The Microbiome of the Medicinal Plants *Achillea millefolium* L. and *Hamamelis virginiana* L.

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In the recent past many studies investigated the microbiome of plants including several medicinal plants (MP). Microbial communities of the associated soil, rhizosphere and the above-ground organs were included, but there is still limited information on their seasonal development, and in particular simultaneous investigations of different plant organs are lacking. Many studies predominantly addressed either the prokaryotic or fungal microbiome. A distinction of epi- and endophytic communities of above-ground plant organs has rarely been made. Therefore, we conducted a comprehensive investigation of the bacterial and fungal microbiome of the MP *Achillea millefolium* and studied the epi- and endophytic microbial communities of leaves, flower buds and flowers between spring and summer together with the microbiome of the associated soil at one location. Further, we assessed the core microbiome of *Achillea* from four different locations at distances up to 250 km in southern Germany and Switzerland. In addition, the bacterial and fungal epi- and endophytic leaf microbiome of the arborescent shrub *Hamamelis virginiana* and the associated soil was investigated at one location. The results show a generally decreasing diversity of both microbial communities from soil to flower of *Achillea*. The diversity of the bacterial and fungal endophytic leaf communities of *Achillea* increased from April to July, whereas that of the epiphytic leaf communities decreased. In contrast, the diversity of the fungal communities of both leaf compartments and that of epiphytic bacteria of *Hamamelis* increased over time indicating plant-specific differences in the temporal development of microbial communities. Both MPs exhibited distinct microbial communities with plant-specific but also common taxa. The core taxa of *Achillea* constituted a lower fraction of the total number of taxa than of the total abundance of taxa. The results of our study provide a basis to link interactions of the microbiome with their host plant in relation to the production of bioactive compounds.

**Keywords:** fungi, bacteria, epiphytic, endophytic, diversity, phyllosphere, season, core-community

**INTRODUCTION**

Epi- and endophytic microorganisms, including prokaryotes and fungi, colonize plants above ground (phyllosphere) and below ground (rhizosphere) and exhibit different ecological interactions with their host from beneficial and commensal symbiosis to pathogenic relationships (Hardoim et al., 2015; Jia et al., 2016; Brader et al., 2017). Uniquely shaped microbiomes were found in a
variety of plants and their different compartments such as rhizosphere, leaf, flower, fruit, and seed (Copeland et al., 2015; Grudzinska-Sterno et al., 2016; Hamonts et al., 2018; Wei and Ashman, 2018; Grady et al., 2019; Comeau et al., 2020; Massoni et al., 2020). Embedded in a wide range of environmental constraints, the structure of the plant microbiome is far from random; its formation and temporal development is a continuous process, expressed by specific co-occurrence patterns and microbial interactions (Cardinale et al., 2015; Berg et al., 2016; Grady et al., 2019). Variables affecting the microbiome and its temporal development in the different compartments include environmental factors such as wind, rain, temperature and soil chemistry, leading to horizontal transmission of microbes and structuring mainly the epiphytic microbial communities (Müller et al., 2016). Some microbes can also establish in the phyllosphere by vertical transmission from seeds and plant compartments and affect mainly the endophytic microbial communities (Müller et al., 2016).

Past research on plant microbiomes has demonstrated growth-promoting effects (Naik et al., 2019), the enhancement of plant resistance against pathogens (Ansary et al., 2018; Vurukonda et al., 2018) and positive effects on overall plant fitness (Rodriguez et al., 2009; Uroz et al., 2019). Further, implications of the microbiome on agricultural management (Toju et al., 2018; Jones et al., 2019; Pacifico et al., 2019; Lyu et al., 2020), crop quality and postharvest processes (Rillig et al., 2018) have been addressed. Several studies investigated the microbiome of medicinal plants (MPs), e.g., *Origanum vulgare* (Pontonio et al., 2018; Castronovo et al., 2020), *Matricaria chamomilla*, and *Calendula officinalis* (Köberl et al., 2019) as well as *Salvia miltiorrhiza* (Chen et al., 2018; Huang et al., 2018). MPs are characterized by a high variety of unique secondary metabolites and many endophytes producing pharmacologically active substances (Köberl et al., 2013; Golinska et al., 2015; Ivanova et al., 2016; Rahman et al., 2017; Caruso et al., 2020; Gupta et al., 2020; Teimoori-Boghsani et al., 2020). Thus, beside the potential impact of the plant microbiome on pathogen resistance and general fitness evidence has accumulated that the microbiome has a significant impact on the production of secondary metabolites of MPs (Köberl et al., 2013; Schmidt et al., 2014; Jia et al., 2016; Huang et al., 2018; Pacifico et al., 2019). This has recently been specified for *Echinacea purpurea* (Maggini et al., 2017; Haron et al., 2019), *Camellia sinensis* (Sun et al., 2019), and *Cannabis sativa* (Taghinasab and Jabaji, 2020). Further work is still required to illuminate mutual interactions of MPs and their microbiome and its potential role in producing beneficial bioactive compounds.

A very important but almost unexplored aspect in this context is the seasonality of the microbiome. To obtain a high-quality herbal drug, the MP has to be harvested at the optimum developmental stage of the desired plant organ (Chen et al., 2012; Duckstein et al., 2012b; Duda et al., 2015). It is completely unclear how this optimum stage is reflected in the plant microbiome and how it establishes over time. As a first step toward this aim, Comeau et al. (2020) identified spatio-temporal variations in the microbiome of *C. sativa* at different developmental stages. Further, endophytic *Actinobacteria*, known for their high potential of producing bioactive compounds, vary seasonally in tropical MPs (Barman and Dkhar, 2020). Therefore, knowledge about the seasonal development and dynamics of the microbiome should be part of a comprehensive understanding of the complex interactions between MP metabolites and the plant’s microbiome. This may also have an impact on the pharmacological potential and further processing such as fermentation of MPs (Duckstein et al., 2012b; Hussain et al., 2016; Köberl et al., 2019).

To elucidate differences in microbial communities and their temporal dynamics of two MPs with different habitus and lifestyle we investigated the microbiome of *Achillea millefolium* (in the following *Achillea*) and *Hamamelis virginiana* (in the following *Hamamelis*). *Achillea*, commonly referred to as yarrow, is a perennial plant native to the Northern hemisphere of Asia, Europe and America. Pharmaceutical preparations of this MP are used to treat loss of appetite, gut complaints, wounds, and menstrual spasms (EMA/HMPC, 2018), to reduce inflammation (Benedek et al., 2007) and even show in vitro potential against cancer (Königl et al., 2017; Pereira et al., 2018). *Hamamelis*, with the common name witch hazel, is a deciduous arborescent shrub or small tree native to eastern North America and is used in the treatment of inflammations of skin and mucous membranes, varicose veins, and hemorrhoids. Furthermore, *Hamamelis* is frequently applied in cosmetics, i.e., skin lotions, nourishing creams, pre-, and after-sheaves (EMEA, 2009). The endophytic bacterial community of *Achillea* on hydrocarbon-contaminated and natural soil (Lumactud et al., 2016; Lumactud and Fulthorpe, 2018) and endophytic fungi of this plant have been investigated recently (Satari et al., 2016; Hatamzadeh et al., 2018, 2020). We extended the investigations of the microbiome of *Achillea* by including epi- and endophytic bacterial and fungal communities of the associated soil, leaf, flower bud and flower, including their temporal dynamics and a comparison of four different locations. We further investigated for the first time the epi- and endophytic microbiome of *Hamamelis*. We addressed the following questions: (i) how do the endo- and epiphytic bacterial and fungal communities of *Achillea* change from the early leaf development stage in spring to the flowering stage in summer and the respective microbial leaf communities of *Hamamelis* from spring to summer? (ii) How does the composition of the epi- and endophytic bacterial and fungal communities differ between each other and both plants? (iii) *Does Achillea* harbor a distinct core community of bacteria and fungi? Illumina Miseq sequencing was applied to generate amplicon datasets of the 16S rRNA gene of the bacterial and the intergenic transcribed spacer (ITS) for the fungal community. The results show that *Achillea* and *Hamamelis* exhibit distinct microbiomes above ground both in their bacterial and fungal communities, which also undergo seasonal changes and that *Achillea* harbors a specific core community when comparing different locations up to 250 km apart.

**MATERIALS AND METHODS**

**Sampling**

Plant and soil samples were collected in 2016 during the growing period of both MPs in the biodynamic garden of WALA Heilmittel GmbH (location 1a) from April until July for *Achillea*
and until September for Hamamelis, respectively, (Table 1). Both plants have been growing at this location in close proximity for more than a decade. To evaluate the impact of different locations, we sampled Achillea again in July 2018 at the same place (location 1b) in comparison with three other locations, i.e., wild plants beside a meadow with cows near Göppingen (Germany; location 2), a meadow close to the river Thur near Kleinandelfingen (Switzerland; location 3) and in a biodynamic garden in the Swiss Alps (1,096 m above sea level; location 4).

To ensure sufficient coverage of the phyllosphere, each biological sample contained leaves, flower buds or flowers of three individual plants growing in close proximity. Sampling was done in triplicate and at least 48 h after rainfall. Leaves of Hamamelis were collected at 1.2 to 2.0 m above ground. The herbaceous plant Achillea develops different leaves during the growing period. Thus, we first sampled leaves of the rosette in April and May and switched to the upper leaves of the stem in June and July (because rosette leaves decay until summer). The flower buds of Achillea in June were still closed and sampled together with their small surrounding young leaves (Figure 1). At each sampling also bulk soil was sampled in triplicate (top 20 to 70 mm) next to the plant below the canopy. Samples were stored at −80°C until analysis. The whole sampling procedure was carried out with gloves, DNA-free tweezers and DNA-free bags. Samples were transported in cooling bags at 4°C to the adjacent laboratory for further sample preparation.

### Differentiation Between Epi- and Endophytic Microorganisms

To obtain epiphytic communities all plant samples were run through a washing procedure by shaking (vortex) for 3 min in sterile 1x PBS-Silwet buffer; pH 7.4 (Silwet L77, final conc. 0.02%), followed by ultrasonication (640 W, Sonorex Digipuls, DL 510 H, Bandelin, Berlin, Germany) for 3 min. The sonication power was reduced to 0.2 µm filter and stored at −80°C until DNA extraction. The endophytic plant samples, after ultrasonication treatment, were transferred in 50 ml Falcon tubes, rinsed additionally by shaking in fresh sterile PBS buffer to remove remaining epiphytes and stored at −80°C until extraction. For each extraction, PBS-Silwet buffer served as control.

### DNA-Extraction

To investigate the closely related and specialized endophytic community, all washed plant compartments were ground in liquid nitrogen just before DNA extraction. We used the DNeasy PowerSoil Kit (Qiagen, Hilden, Germany) following the manual to extract DNA. The final extracts were quantified using a Qubit 2.0 fluorometer (Invitrogen/ThermoFisher, Waltham, United States) and diluted with DNA-free water or concentrated by using Co-Precipitant Pink (Bioline, Memphis, TN, United States) following the manual if necessary. All samples were adjusted to a final concentration of 1–10 µg/µl DNA.

### Sequencing

Samples were sequenced by LGC Genomics GmbH (Berlin, Germany) on an Illumina MiSeq platform (Illumina, San Diego, United States). Fungi were amplified with primer ITS1F (5′-CTTGGTCATTGAGGAAGTAA-3′)/ITS2 (5′-GCTGCGTTCTTCATCGATGC-3′) targeting the ITS 1 region. Bacteria were amplified with primer pair 799f/1115r to reduce chloroplast-DNA signals. Controls were visually checked through an agarose gel.

### Table 1

| Species          | Sampling date | Time series | Spatial comparison | Location               | Location ID | Elevation (m aSL) | pH Coordinates | Coordinates | Type       |
|------------------|---------------|-------------|-------------------|------------------------|-------------|------------------|----------------|-------------|------------|
| Hamamelis       | 06 Apr 2016   | X           |                   | D- WALA garden         | 1a          | 444              | n/a            | 46―3745"      | 9735′13"   | Leaf, soil |
| Hamamelis       | 18 Apr 2016   | X           |                   | D- WALA garden         | 1a          | 444              | n/a            | 46―3745"      | 9735′13"   | Leaf, soil |
| Hamamelis       | 04 May 2016   | X           |                   | D- WALA garden         | 1a          | 444              | n/a            | 46―3745"      | 9735′13"   | Leaf, soil |
| Hamamelis       | 08 May 2016   | X           |                   | D- WALA garden         | 1a          | 444              | n/a            | 46―3745"      | 9735′13"   | Leaf, soil |
| Hamamelis       | 04 Jul 2016   | X           |                   | D- WALA garden         | 1a          | 444              | 7.2            | 46―3745"      | 9735′13"   | Leaf, soil |
| Hamamelis       | 20 Sep 2016   | X           |                   | D- WALA garden         | 1a          | 444              | n/a            | 46―3745"      | 9735′13"   | Leaf, soil |
| Achillea        | 11 Apr 2016   | X           |                   | D- WALA garden         | 1a          | 444              | n/a            | 46―3745"      | 9735′14"   | Leaf, soil |
| Achillea        | 04 May 2016   | X           |                   | D- WALA garden         | 1a          | 444              | n/a            | 46―3745"      | 9735′14"   | Leaf, soil |
| Achillea        | 10 Jun 2016   | X           |                   | D- WALA garden         | 1a          | 444              | n/a            | 46―3745"      | 9735′14"   | Leaf, soil |
| Achillea        | 23 Jul 2016   | X           |                   | D- WALA garden         | 1a          | 444              | 7.7            | 46―3745"      | 9735′14"   | Leaf, bud, soil |
| Achillea        | 19 Jul 2018   | X           |                   | D- WALA garden         | 1b          | 444              | 7.7            | 46―3745"      | 9735′14"   | Leaf, flower, soil |
| Achillea        | 12 Jul 2018   | X           |                   | CH- Alp-garden         | 4           | 1096             | 7.6            | 46―4341"      | 80′1′15"   | Leaf, flower, soil |
| Achillea        | 12 Jul 2018   | X           |                   | CH- Kleinandelfingen, meadow | 3          | 578              | 7.8            | 47―3554"      | 84′1′25"   | Leaf, flower, soil |
| Achillea        | 19 Jul 2018   | X           |                   | D- Goepppingen, meadow | 2           | 354              | 7.8            | 48―4057"      | 9838′08"   | Leaf, flower, soil |

Type indicates plant compartment and soil sampled. Time series and spatial comparison highlight the samples used for the seasonal or spatial comparison. Samples of both plants in July 2016 were used for direct comparison and are marked with *.
Bioinformatics

Datasets of the bacterial 16S rRNA gene and fungal ITS sequences were processed as follows: Trimmomatic 0.36 (Bolger et al., 2014) was used to truncate low quality read ends if the average quality dropped below 15. Primer sequences were removed from amplicon sequences using bbduk. All datasets were subsequently processed with USEARCH v.10.0.240 (Edgar, 2010). Sequences were merged and low quality sequences were discarded [shorter than 300 base pairs (bp), accumulative sequencing error rate ≥ 1%] resulting in 9,652,175 and 9,128,731 high quality (HQ) amplicon sequences for the 799f-1115r primer set and the fungal ITS, respectively. HQ sequences were pooled according to the used primer set. In addition, sequences from the 16S rDNA dataset were truncated to equal length of 300 bp. Due to high variance in ITS sequence length this step was skipped for fungal ITS datasets. Subsequently, all sequences were dereplicated and sorted by abundance. Chimeric sequences were removed and remaining sequences clustered into zero-radius taxonomic units (ZOTUs) using the unoise3 algorithm with a minimum unique sequence abundance of 8 across all samples. In total 20,133 (16S rDNA dataset) and 5,932 (ITS) ZOTUs were generated. All ZOTUs were taxonomically classified by alignment employing the USEARCH algorithm against the GTDB SSU database (Parks et al., 2018; release 89) for 16S rRNA gene sequences and the UNITE ITS database 8.0 (Abarenkov et al., 2020) with an e-value cut-off of 1e-10 and minimum sequence identity of 90% and maxaccepts/maxrejects option disabled. An abundance table was created by mapping HQ sequences of each sample to the ZOTUs.

Statistical Analysis

Statistical analyses were performed using R v3.6.0 (R Core Team, 2018) with the packages vegan (Oksanen et al., 2017), ape (Paradis and Schliep, 2018), drc (Ritz et al., 2015), ade4, and picante (Kembel et al., 2010). Only samples with more than 1,000 mapped reads were considered for further analysis.

To account for varying sequencing depth, count tables of bacterial 16S rDNA and fungal ITS datasets were repeatedly rarefied to 1,000 sequences per sample (99 times). Subsequently, richness and Shannon entropy as well as species coverage were calculated for each iteration and the mean value was used for further analysis. Effective Number of taxa (EN) was calculated according to Jost (2006). Linear model fitting was used to determine a relationship between richness and EN of the associated leaf community to the growth period of Achillea and Hamamelis. P-values < 0.05 were considered significant. Differences between richness and diversity of the microbiome of different plant compartments were tested using analysis of variance (ANOVA) and Tukey’s honest significant difference test after checking for normal distribution (Shapiro–Wilk test) and homoscedasticity (Flinger test).

Prior to further analysis unrarefied samples were converted to relative abundances by dividing individual ZOTU counts by the total number of reads per sample. To determine core and
specific plant species/tissue community, only bacterial and fungal taxa that occurred in at least two triplicates or in more than 50% of the replicates, if more than three replicates were used, were considered. Bray–Curtis distances (Bray and Curtis, 1957) of bacterial and fungi abundances at each location were visualized by non-metric multidimensional scaling (NMDS; $k = 2$; 999 permutations). A permutational multivariate analysis of variance (PERMANOVA) with 9,999 permutations was performed to determine the dataset variance explained by sample type, plant species, tissue, and soil pH. The core microbiome of the four locations was determined only for the July samples of Achillea. To exclude rare taxa in the core microbiome analysis, all bacterial and fungal genera, which occurred with at least 0.01% of the total community in more than two thirds of a particular sample type (leaf epi, leaf endo, flower epi, or flower endo) at one location, were considered to be present. Mantel tests (9,999 permutations) using Bray–Curtis distances were used to determine correlation between bacterial and fungal datasets.

**Soil Analysis**
Wavelength-Dispersive X-Ray Fluorescence (XRF; Axios FAST, Malvern Panalytical, Malvern, United Kingdom) analyses was used to determine the concentrations of selected major (Si, Al, Fe, Mg, Ca, N, K, P) and trace (Mn, As, Co, Cr, Cu, Mo, V, U, Zn, Zr) elements of soil samples. For each analysis 0.7 g of dried and sieved (2 mm) sample were used and measured according to Atar et al. (2019). Major and trace elements are reported as weight% and ppm, respectively.

Soil pH was measured in an aqueous suspension (soil:deionized water, 1:2.5, v:v) using a InLab Micro (Mettler Toledo, Giessen, Germany).

**RESULTS**
We assessed the microbial community of 45 soil and 120 plant samples from above ground, comprising epiphytic and endophytic communities of Achillea (24 leaves, 3 flower buds, 15 flowers) and Hamamelis (18 leaves). The soil communities showed the highest richness and EN diversity of bacterial species and fungal genera compared to plant samples, the latter exhibiting a decreasing diversity from epiphytic leaf to endophytic flower communities. Richness of each of the epiphytic communities was significantly higher than that of their corresponding endophytic compartment, except for fungi on flowers (ANOVA and post hoc test, $p < 0.05$). The EN differed only significantly between epi- and endophytic bacterial communities on leaves ($p < 0.05$; Figure 2; Supplementary Table 1). As shown by rarefaction analysis, sequencing efforts

![FIGURE 2](FIGURE 2 | Box-Whisker plots of Richness and Effective Number diversity of bacterial and fungal epi- and endophytic communities of the flower and leaf compartments as well as soil of Achillea and leaf and soil of Hamamelis. Given are mean values, quantiles 2 and 3 (box) and 1 and 4 (lines) and outliers (<5% and >95% of the mean). In parenthesis: number of samples. Samples were subsampled (repeatedly rarefied to 1,000 sequences) for comparison. For statistical comparisons of these data see Supplementary Table 1.)
of most of the plant samples and also of the highly diverse soil samples reached almost saturation (Supplementary Figure 1).

The Microbiome of Achillea and Hamamelis

In July, Achillea starts to bloom and Hamamelis has fully developed leaves. Therefore, we assessed the microbiome of both plants in the WALA garden (location 1a) at this time. Hamamelis blooms in winter, consequently, flowers of this plant were not included in this survey. As described for the overall assessment (see above) both plants at this location and time also showed a decreasing number of total detected bacterial and fungal taxa from soil to epi- and endophytic leaf and further for Achillea to the epi- and endophytic floral communities (Figures 3A–D).

Besides this generally decreasing number of total bacterial and fungal taxa, proportions of compartment-overlapping taxa of Achillea decreased from soil to leaf and flower as indicated by the numbers in intersecting areas of Figures 3A,B. Forty-eight percent of all 878 plant-associated bacterial species were exclusive to plant tissue (28% to leaf, 10% to flower, and 10% to both) and not detected in soil. Respective proportions of fungi were even higher as 58% of all 102 plant-associated fungal genera were exclusive to plant tissue (15% to leaf, 18% to flower, and 25% to both). Only 7% of all bacterial species and 31% of all fungal genera associated with plant tissue were shared among all plant compartments (Figures 3A,B). For Hamamelis, 57% of all 418 leaf-associated bacterial species and 80% of all 137 leaf-associated fungal genera were exclusive to the plant compartments and not detected in soil. Thus, Hamamelis harbored a higher relative proportion of leaf-exclusive bacteria and in particular of fungi than Achillea (Figures 3C,D).

Hamamelis and Achillea grew in close proximity (max. 10 m distance) in the WALA garden. The composition of the soil communities of both plants showed high similarity, the bacterial communities by more than 79% and the fungal communities by more than 68% (Figures 3E,F). Pronounced differences, however, were detected in the leaf microbiome of both plants. Leaves of

FIGURE 3 | Venn-diagrams of the partitioning of the bacterial species and fungal genera of Achillea (A,B) among the epi- and endophytic leaf, flower and the soil compartments, and of Hamamelis (C,D) among the epi- and endophytic leaf and the soil compartments in July in the WALA garden in 2016 (location 1a). Partitioning of the bacterial species (E) and fungal genera (F) of the endo- and epiphytic leaf and soil communities between Achillea (Ach) and Hamamelis (Ham) is also shown. Intersection without number: detected taxa = 0. In parenthesis: number of compartment-specific taxa.
*Achillea* exhibited a much higher total number of bacterial species and endophytic fungal genera than *Hamamelis*, whereas the total number of epiphytic fungal genera was higher in leaves of the latter (Figures 3E,F).

**Seasonal Leaf Diversity**
To examine whether leaf development over the season also affected the associated microbial communities, we analyzed time series of the microbial leaf communities of *Achillea* and *Hamamelis* collected in the WALA garden. At the first sampling point in April, the endophytic bacterial and fungal communities exhibited a lower richness and EN than the epiphytic communities (Figure 4). The epiphytic bacterial community of *Achillea* revealed some temporal fluctuations but no consistent temporal pattern. However, the richness and EN of the endophytic bacterial community increased consistently over time (Figure 4). In contrast, the fungal leaf community of *Achillea* decreased over time except in richness of the endophytic

![Graphs showing microbial diversity over time](image-url)
community (Figure 4). At the last sampling point in July, richness and EN of the epi- and endophytic communities of bacterial species and fungal genera were similar (Figure 4).

For Hamamelis, different patterns were observed. Richness and EN of the endophytic bacterial leaf community remained similar over time, whereas both variables of the epiphytic community increased significantly, finally yielding higher values than the endophytic community (Figure 4). The fungal epi- and endophytic leaf communities of Hamamelis increased over time as well, but temporal fluctuations were rather high (Figure 4).

An NMDS analysis confirmed the temporal patterns and changes of both microbial communities of both MPs (Figure 5). This analysis was extended to the microbial communities of the soils of both MPs and to the flower buds and flowers of Achillea. It revealed distinct and temporally stable soil communities of both MPs. The epi- and endophytic leaf communities of Hamamelis were slightly different with some temporal variations. For Achillea the NMDS analysis revealed distinct leaf-, bud- and flower-associated bacterial communities. The epiphytic fungal community was slightly but consistently different from the respective endophytic community. A PERMANOVA considering only plant-associated samples revealed the plant species as the major determinant explaining the differences in the bacterial and fungal communities, 34% for bacteria and 40% for fungi. The tissue type (leaf or flower) covered 23% and 17% of the variation of bacteria and fungi, respectively. The distinction in epi- and endophytic communities was not significant and explained less than 3% of the community variance (Supplementary Figure 2).

**Community Composition and Seasonal Dynamics**

Characteristics and temporal dynamics of the microbial communities of Achillea, Hamamelis and the associated soil between April and September were clearly reflected in the 50 most abundant bacterial and fungal taxa (Figures 6, 7 and Supplementary Figures 8, 9). As indicated above for the entire communities also the most abundant taxa exhibited a much greater diversity of the soil-associated microbial communities than the above-ground plant-associated communities.

Bacterial communities of both MPs were composed of Alphaproteobacteria, Gammaproteobacteria, Actinobacteria, Firmicutes, Deinococci, and Bacteriodota (Figure 6). Alphaproteobacteria were most abundant with 40 to over 60% of total abundance on leaves and higher proportions on Hamamelis than on Achillea. Sphingomonadales and Rhizobiales constituted major alphaproteobacterial fractions and increased in abundance over the progressing season. Furthermore, Actinobacteria were detected on leaves of Hamamelis and constituted up to 12% of total abundance. Gammaproteobacteria were the second most abundant group on leaves of both MPs and on Achillea even dominated initially on leaves, flower buds and flowers to more than 70%. On leaves of Hamamelis, Gammaproteobacteria decreased in proportion and finally constituted less than 15% in the epiphytic and below 5% in the endophytic communities. Burkholderiales were the most abundant order of this subclass on leaves of both plants and on flower buds of Achillea. Pseudomonadales and Enterobacterales also constituted substantial proportions, in particular in spring on leaves and the latter on flowers constituting 35% of the epiphytic and 70% of the endophytic community. Further, Actinobacteria with several orders, Firmicutes, mainly their orders Clostridiales, Lactobacillales and Phaenibacillales, Bacteriodota, mainly their order Cytophagales, and Deinococci with their order Deinococcales were consistently present on leaves of both MPs. Actinobacteria constituted between 5 and 18% and the other groups not more than 7% each (Figure 6).

Genera such as Sphingomonas, Methylobacterium, Massilia, Herbaspirillum, Kineococcus, and Deinococcus were shared by both MPs but leaves of each plant exhibited specific patterns, which differed in relative proportions of these genera.

**Figures 6**
FIGURE 6 | Relative proportions of the 50 most abundant bacterial orders of the epi- and endophytic leaf, bud, flower and soil communities of Achillea (Ach) over a period of 103 days (April to July) and of the epi- and endophytic leaf and soil communities of Hamamelis (Ham) over periods of 89 and 167 days (April to July and September), respectively.

(Supplementary Figure 8). Plant-specific seasonal dynamics were also recorded. On Achillea leaves, relative proportions of Methylobacterium, Kinneococcus and Spiroforma increased over time and those of Pseudomonas, Janthinobacterium and FW-11 decreased. Few genera like Gilliamella, Conexibacter, and Microvirga exhibited higher proportions in the epiphytic relative to the endophytic leaf community of Achillea in April and compared to later time points (Supplementary Figure 8). On Hamamelis leaves, Deinococcus and Microvirga increased their relative proportions over time and those of Herbaspirillum, Burkholderia and Massilia decreased. Gilliamella exhibited a higher proportion of the epiphytic leaf community of Hamamelis relative to Achillea (Supplementary Figure 8).

The dominant bacteria of the associated soil of both plants were almost identical and showed only minor temporal variations (Figure 6). Some typical plant taxa were not detected or were exceedingly rare in soil such as Methylobacterium, Kinneococcus, Lactobacillus, and Deinococcus (Figure 6 and Supplementary Figure 8).

Fungal communities of both MPs and the associated soils were clearly different (Figure 7 and Supplementary Figure 9). On leaves of Hamamelis, taxa of the phylum Ascomycota dominated by over 90% of total fungal abundance, in particular the order Capnodiales. On Achillea, taxa of the phylum Basidiomycota and various orders were also prominent and constituted proportions of roughly 20 to about 60% (Figure 7). Highest proportions of taxa of this phylum were detected on leaves at the second sampling date (May) when new rosette leaves had developed, and the first stem growth appeared as well as later on flower buds. Respective proportions of Ascomycota on these dates and compartments were below 40%. On flowers, unidentified fungi constituted ∼35% of total abundance whereas they constituted <10% on buds and leaves (Figure 7). Fungi also showed a clear plant-specific community, but several dominant genera also overlapped between both plants such as Didymella, Mycosphaerella and Cladosporium but often with a plant-specific dominance (Supplementary Figure 9). The fungal communities of Hamamelis were dominated by the genus Microcyclospora, an unidentified Capnodiales, Uwebraunia and Xenosonderhena, whereas Achillea harbored other dominant fungi, such as Leucosporidium, Septoria, Filobasidium, Tetracladium, and Stragonosporopsis.
The fungal leaf communities of *Achillea* underwent seasonal variations. Basidiomycotal genera such as *Leucosporidium*, *Filobasidium* and *Dioszegia* relatively decreased in both epi- and endophytic leaf communities during the growing season (Supplementary Figure 9). In contrast, ascomycotal genera such as *Didymella*, *Boeremia* and *Ramularia* increased, together with the basidiomycotal genus *Thanatephorus* (Supplementary Figure 9). Only the latter was enriched in the endophytic leaf community at the final sampling point in July, together with an unidentified member of the *Plectosphaerellaceae*. The composition of the fungal leaf communities of *Hamamelis* remained rather stable seasonally. However, *Capnodiales* relatively decreased during summer in the endophytic leaf community and exhibited reduced numbers in the epiphytic as well as at the last sampling date in September. At this date, *Dothideomycetes* with its lineages *Didymella*, *Mycosphaerella* and *Epicoccum* increased concomitantly (Supplementary Figure 9).

Fungi detected in soil were mostly absent or present in low abundances in the above-ground plant compartments. The fungal soil communities were dominated by 50 to 80% of total abundance by various orders of *Ascomycota* (Figure 7). *Basidiomycota* constituted <30% and in most cases <15%. Further, *Mortierellales*, not at all detected in the MPs above ground, constituted ~8 to 23% in the soil of both plants with higher proportions in that of *Hamamelis*. Other and unidentified fungi of generally minor proportions revealed some differences between the soils of both MPs (Figure 7). A few dominant genera were only detected in the soil of *Achillea*, such as *Didymella* and *Tetracladium*, whereas three genera of the *Clavariaceae* (*Clavaria*, *Clavulinopsis*, one unidentified) dominated in *Hamamelis* soil (Figure 7 and Supplementary Figure 9).

On *Achillea*, the number of genera among the most abundant 50 bacterial and fungal taxa decreased from leaves to flower buds and flowers (Supplementary Figure 8). Two enterobacterial genera, *Nissabacter* and *Pectobacterium*, and one fungal genus, *Botryosphaeria*, were detected exclusively on flowers and even as prominent community members (Supplementary Figures 8, 9).

To explore the connection between fungal and bacterial community we computed Mantel correlations of Bray-Curtis distances between both communities for all soil, *Achillea* and *Hamamelis* samples. Communities present in *Achillea* (Mantel’s $r = 0.59$, $p < 0.001$) and soil (Mantel’s $r = 0.49$, $p < 0.001$)
were tightly linked whereas communities from Hamamelis (Mantel’s $r = 0.21$, $p = 0.002$) showed a weaker but still significant correlation.

**Effect of Location on Composition of Microbial Communities of Achillea**

To investigate the geographic impact on the microbial communities of Achillea, soil, leaves and flowers were collected in July 2018 at four locations in southern Germany and Switzerland, up to 250 km apart. The sites differed by altitude, soil pH (Table 1 and Figure 8D) and inorganic soil elements (Supplementary Figure 3). Locations 2 and 3 exhibited pronounced differences in their relative proportions of Ca and Si as compared to locations 1b and 4, which are biodynamically cultivated. A PERMANOVA revealed that location explains 18% of the variance of bacterial and fungal plant communities, closely followed by plant compartment (leaf or flower), explaining 18% and 14% of the variance of bacterial and fungal communities, respectively. Differences in the epiphytic and endophytic sub-communities were only significant for the fungal communities and explained 10% of the variance (Figure 8C). Soil pH explained only 5% of the plant-community variance.

The composition of the soil-associated microbial communities was location-specific (Supplementary Figure 4). These patterns of the microbial soil communities were not reflected in the respective plant communities as illustrated by an NMDS-analysis (Figures 8A,B). Clustering of the leaf- and flower-associated microbial communities was mostly compartment-specific and only the fungal and endophytic bacterial floral communities at location 4 were distinctly different from those of the other locations (Figure 8 and Supplementary Figure 7).

**Core Community and Unique Taxa of Plant Compartments of Achillea and Soil**

To elucidate general and location-specific aspects of the microbiome of Achillea, we assessed the microbial core community and unique taxa of a given location. We defined the core community as a subset of taxa occurring at all locations and unique taxa as those found only at one location and more specific in a plant compartment or soil. The analysis is based on
all detected taxa that occurred in at least two of the triplicates. Numbers of core taxa decreased from soil to flower and from epiphytic to endophytic compartments (Table 2) and reflect the general patterns described above. Numbers of total and unique taxa followed the same general pattern, except for the epiphytic bacterial floral communities which exhibited the highest number of taxa (Table 2) possibly reflecting that the flower is affected greatest by environmental impacts and possible transmission by microbes including pollinating insects. The proportion of bacterial core taxa as percent of total bacterial abundance decreased less from soil to flower than their percentage as number of total bacterial taxa. Respective percentages of fungi even increased (Table 2). The fraction of bacterial core taxa as percent of total abundance in the plant compartments ranged between 54 and 95% not exceeding 29% as fraction of the total number of bacterial taxa. Percentages in the endophytic compartments were consistently higher than in the epiphytic compartments. In the endophytic bacterial floral community with the lowest number of taxa, the 28 core genera comprised only 17% of the number of total bacterial taxa but almost 76% of total bacterial abundance. Accordingly, the 19 core genera of the fungal endophytic floral community constituted 18% of the number of total genera but 95% of the total abundance (Table 2).

As the pure "core" number does not reflect whether a core taxon is specific to one compartment or included in the core community of another one, we also assessed the compartment-specific core taxa, comprising all taxa solely detected in a specific compartment. Soil samples revealed the largest compartment-specific core community, both in total number and in relative abundance (Supplementary Table 2). Compartment-specific bacterial epiphytic core communities revealed more taxa than endophytic core communities, and no bacterial genus was exclusively detected in the endophytic core community of leaves. Five genera were exclusive to the endophytic floral community of which three belonged to Enterobacteriaceae (Supplementary Tables 2, 3). Regarding fungi, nine and five genera occurred exclusively in the epiphytic and endophytic core community of leaves, respectively. Only three basidiomycotal genera occurred exclusively as core genera of the epiphytic flower community (Supplementary Tables 2, 4).

An even more rigorous analysis excluding any detection in soil revealed a plant-exclusive endophytic core community in leaf and flower of two and three bacterial genera and eight and six bacterial species, respectively, (Supplementary Table 2, plant core never in soil). Similarly, four and one of the endophytic fungal core genera were never detected in soil (Supplementary Table 2, plant core never in soil). These results show that the core microbiome of Achillea harbors soil-independent plant compartment-specific core taxa in both its endophytic and epiphytic compartments.

### DISCUSSION

Since the advent of high throughput sequencing our knowledge and understanding of the composition and diversity of bacterial and fungal communities of the various plant compartments above ground, the rhizosphere and the associated soil has greatly improved. The general pattern emerged that the diversity decreases from below ground to stem, twig, leaf and flower with the strongest decrease from below to above ground (Ottesen et al., 2013; Gomes et al., 2018; Hamonts et al., 2018; Wei and Ashman, 2018; Grady et al., 2019; Comeau et al., 2020; Massoni et al., 2020), even though different trends for leaf and flower were also reported, e.g., for the endemic Hawaiian tree Metrosideros polymorpha (Junker and Keller, 2015) and wild

| Compartment       | Total taxa | Sum of unique taxa of one location | Unique taxa at one location | Core taxa, absolute | Core taxa,% of total taxa | Core taxa,% of total abundance |
|-------------------|------------|----------------------------------|---------------------------|--------------------|--------------------------|--------------------------------|
| **Bacteria species** |            |                                  |                           |                    |                          |                                 |
| Soil              | 1985       | 465                              | 0.7                       | 836                | 42.1                     | 93.7                           |
| Leaf epiphytic    | 913        | 334                              | 2.2                       | 230                | 25.2                     | 80.5                           |
| Leaf endophytic   | 659        | 307                              | 2.4                       | 113                | 17.1                     | 90.1                           |
| Flower epiphytic  | 1002       | 612                              | 9.0                       | 80                 | 8.0                      | 54.2                           |
| Flower endophytic | 329        | 163                              | 1.9                       | 34                 | 10.3                     | 64.9                           |
| **Bacteria genera** |            |                                  |                           |                    |                          |                                 |
| Soil              | 826        | 150                              | 0.2                       | 438                | 53.0                     | 97.2                           |
| Leaf epiphytic    | 409        | 133                              | 0.3                       | 120                | 29.3                     | 90.1                           |
| Leaf endophytic   | 263        | 103                              | 1.9                       | 59                 | 22.4                     | 94.6                           |
| Flower epiphytic  | 451        | 236                              | 1.8                       | 73                 | 16.2                     | 73.7                           |
| Flower endophytic | 170        | 77                               | 0.8                       | 28                 | 16.5                     | 76.1                           |
| **Fungi genera**  |            |                                  |                           |                    |                          |                                 |
| Soil              | 292        | 111                              | 4.4                       | 64                 | 21.9                     | 65.4                           |
| Leaf epiphytic    | 160        | 52                               | 1.8                       | 40                 | 25.0                     | 91.5                           |
| Leaf endophytic   | 163        | 64                               | 5.4                       | 42                 | 25.8                     | 88.5                           |
| Flower epiphytic  | 213        | 112                              | 1.0                       | 30                 | 14.1                     | 86.5                           |
| Flower endophytic | 104        | 56                               | 1.6                       | 19                 | 18.3                     | 94.6                           |

*Shown are numbers of total taxa, unique taxa (occurring at only one location) and their relative abundance (% of the total community), core taxa and their percentages of the total number of taxa and the total abundance of the community. A taxon was considered as core when it was detected in at least 66% of the replicates.*
strawberries (Wei and Ashman, 2018). In quite a few studies only one or two plant compartments and soil were investigated and even fewer studies investigated prokaryotes and fungi simultaneously (e.g., Hamonts et al., 2018; Comeau et al., 2020). A distinction between epi- and endophytic communities was made even more rarely (e.g., Gomes et al., 2018; Yao et al., 2019) and not yet for prokaryotes and fungi of the same plant. Further, there is still limited information available on the temporal development of plant-associated microbial communities. For a better understanding of the colonization patterns and how they evolve and establish it is important to study these features from below ground to leaf and flower at one plant species.

Therefore, we investigated the epiphytic and endophytic bacterial and fungal communities of leaves, flower buds and flowers of Achillea and the respective epiphytic and endophytic microbial communities of leaves of Hamamelis including the associated soil communities. Flower buds and flowers of Hamamelis were not considered because this deciduous shrub flowers in winter. Overall richness and diversity of bacterial and fungal taxa decreased from soil to leaves of both MPs and further to flowers of Achillea, ending in a specialized flower-associated microbial community of a few dominant members reflecting presumably the shorter life time of flowers as compared to leaves. Such a reduction in richness from leaf to flower has also been reported for bacterial communities in a very recent study on several wild flowers (Massoni et al., 2020) and of tomatoes which further showed that richness of top leaves was similar to that of flowers (Ottesen et al., 2013). In wild strawberries, however, significant differences of the richness of bacterial communities between leaves and flowers were not found (Wei and Ashman, 2018). Similarly, fungal communities associated with leaf and flower of C. sativa neither exhibited differences (Comeau et al., 2020). Studying the endemic Hawaiian tree M. polymorpha Junker and Keller (2015) found a higher bacterial diversity on flower organs than on leaves but a higher Shannon index on the latter. These findings indicate that despite the general decrease in diversity from below to above ground different plants exhibit distinct trends, obviously reflecting intrinsic features such as their specific habitus, e.g., tree, shrub, herbaceous, perennial or annual plant, size, and developmental time. Environmental and climatic factors may further contribute to shaping the plant microbiome.

**Seasonal Patterns of Microbial Leaf Communities**

The seasonal patterns of the epi- and endophytic leaf communities of bacteria and fungi of both MPs exhibited pronounced differences, obviously reflecting the distinct habitus and host-microbe interactions of each MP. In spring, Achillea first develops basal rosette leaves in close contact to the soil; further leaves follow along the growing stem until flower buds are produced. All leaves are relatively short-lived and in contrast to the arborescent shrub Hamamelis do not bud out of the twigs well above the soil. Leaves of Hamamelis are generally bigger, sturdier and persist throughout the season from April to November. Richness and EN of Achillea’s epiphytic bacterial community was highest on the rosette leaves at the first sampling and dropped strongly at the second sampling. Presumably, most of these epiphytic bacteria were transferred from soil, which can be considered as key reservoir or seedbank of leaf microbes (Copeland et al., 2015; Grady et al., 2019). Our NMDS analysis supports this assumption, as the composition of the early epiphytic bacterial community was most similar to that of soil (Figure 5). The strong decline in richness and diversity of this bacterial community at the second sampling of young leaves on the stem seems to reflect that these leaves develop newly without direct contact to the soil and rather little environmental exposition. The slight increase in richness thereafter may reflect that these leaves accumulated bacteria transmitted from the environment. The endophytic bacterial leaf community of Achillea continuously increased from April to July, indicating that the endophytic leaf compartment provided niches for the successful establishment of new taxa in the progressing season. In contrast to bacteria, endo- and epiphytic fungal leaf communities of Achillea decreased in EN diversity and the epiphytic fungal community also in richness from April to July, indicating that a substantial fraction of the fungi associated with the young leaves early in the season did not establish a long term relationship with the MP. As discussed for epiphytic bacteria most of the fungi were presumably transferred from soil to the leaf-rosette, persisted in early leaf stages but did not establish.

For Hamamelis the seasonal development of the bacterial and fungal leaf community was quite different. Whereas the richness and EN diversity of the epiphytic bacterial leaf community increased slightly from April to July, these indices remained almost constant for the endophytic communities. As also the temporal colonization patterns of the bacterial community changed only slightly, these findings indicate that only few new bacteria were able to settle and dwell on leaves after their initial development after budding (Figures 4–6). It appears that bacterial leaf communities of Hamamelis reached a stable composition shortly after early leaf development possibly promoted by certain bacterial taxa transferred from adjacent bark and branch communities. It has been reported from other woody plants and trees that bacterial bark and branch communities are more diverse than the leaf communities (Leff et al., 2015; Harrison and Griffin, 2020). Richness and EN of the fungal leaf communities increased from April to July and that of the epiphytic community further until September, reflected also in the taxonomic colonization pattern (Figures 4, 5, 7). Settling and establishing of new bacterial taxa was of minor significance in Hamamelis, whereas the fungal leaf community responded positively to the leaf development over time by increasing its diversity.

Reasons for these differences in the temporal colonization patterns of the bacterial and fungal leaf communities of Achillea and Hamamelis presumably include differences between the size, development and physiology of a herbaceous perennial plant and a deciduous arborescent shrub. Both MPs provide their respective microbiome with distinct substrates and/or growth-promoting and growth-inhibiting compounds which may also lead to distinct interactions of their bacterial and fungal components (Da Silva et al., 2000; Candan et al., 2003; Saeidnia et al., 2011; Saénchez-Tena et al., 2012; Duckstein et al., 2012a;
Achillea, suggesting that the Nitrogen demand, supply and metabolism of Hamamelis was different from that of Achillea. The leaves of Hamamelis presumably provide a favorable environment for these bacteria. In particular, Acetobacter is able to grow in an acidic environment with a pH below 5.5 (Kersters et al., 2006) and the leaf cytoplasm of Hamamelis exhibits a pH of ~4 (S. Sauer, unpubl. data). In addition to the prominent phylogenetic groups we further detected consistently other groups of lower abundances in the plant compartments in particular of Achillea, such as Deinococcales, Firmicutes, and their order Lactobacillales. This group has also been reported

Microbial Taxa of the Plant and Soil Communities

Our results show that each MP harbors a specific bacterial and fungal microbiome. For Achillea we further showed that the leaf-, bud- and flower-associated microbial communities were composed of a different array of taxa, even though some were also present in adjacent compartments. Other comparative studies have shown that the microbiome of different plants is distinct, reflecting the specific feature of the given plant and its environment (Copeland et al., 2015; Lumactud and Fulthorpe, 2018; Massoni et al., 2020). The various microbial taxa include commensals, potential pathogens but also probiotic and host growth-promoting partners interacting with other microbes and the host plant in various mutualistic and antagonistic and still little understood ways (Hardoim et al., 2015; Berg et al., 2016; Brader et al., 2017; Castronovo et al., 2020).

The bacterial taxa we detected in soil and the plant compartments of both MPs are rather typical as similar phylogenetic groups have also been detected on other plants. Members of Sphingomonadales and Rhizobiales of Alphaproteobacteria are typical leaf-associated bacterial taxa (Copeland et al., 2015; Hardoim et al., 2015; Hamonts et al., 2018; Pontonio et al., 2018; Wei and Ashman, 2018; Grady et al., 2019; Comeau et al., 2020; Massoni et al., 2020). A continuous temporal relative increase of Actinobacteria in the endophytic leaf community, as we observed for Achillea, has already been reported for bean, soybean and canola (Copeland et al., 2015). Endophytic Actinobacteria may be of particular importance in MPs as they produce a diverse range of bioactive compounds (Golinska et al., 2015). Gammaproteobacteria and in particular their orders Pseudomonadales and Burkholderales are further prominent members of the leaf and soil communities and may exhibit plant growth promoting features (Hardoim et al., 2015; Castronovo et al., 2020; Comeau et al., 2020; Massoni et al., 2020). They were abundant members of the leaf communities of both MPs, but differences on the genus level were monitored, indicating that each plant selects for its specific bacterial community. As our and data from other studies show, bacteria can also be prominent members of the flower communities, in particular Enterobacteriaceae (Shade et al., 2013; Aleklett et al., 2014; Junker and Keller, 2015; Wei and Ashman, 2018; Castronovo et al., 2020). In fact, Enterobacteriaceae appear to be specifically associated to the style and stamen of the flower as reported for the tree M. polymorpha (Junker and Keller, 2015). Interestingly, the nitrogen-fixing Acetobacter and further putative N₂-fixing Rhizobiales such as Devosia, Bosea, and Roseomonas were prominent on leaves of Hamamelis and very rare on leaves of Achillea, suggesting that the Nitrogen demand, supply and metabolism of Hamamelis was different from that of Achillea. The leaves of Hamamelis presumably provide a favorable environment for these bacteria. In particular, Acetobacter is able to grow in an acidic environment with a pH below 5.5 (Kersters et al., 2006) and the leaf cytoplasm of Hamamelis exhibits a pH of ~4 (S. Sauer, unpubl. data). In addition to the prominent phylogenetic groups we further detected consistently other groups of lower abundances in the plant compartments in particular of Achillea, such as Deinococcales, Firmicutes, and their order Lactobacillales. This group has also been reported
in other studies (Shade et al., 2013; Copeland et al., 2015; Hardoim et al., 2015; Pontonio et al., 2018; Massoni et al., 2020). *Lactobacillales* may have important and so far only little understood beneficial functions for plant health by producing secondary metabolites promoting plant growth and/or inhibiting growth of pathogens (Lamont et al., 2017; Pontonio et al., 2018; Daranas et al., 2019).

In contrast to bacteria, the fungal communities of both MPs exhibited pronounced differences. Whereas *Basidiomycota* dominated on leaves of *Achillea* during the early developmental stages and on flower buds and constituted 20–25% of total relative abundance on flowers, they comprised only minor proportions on leaves of *Hamamelis*. Interestingly, on flowers fungal taxa, which could not be further affiliated taxonomically, constituted >35%. This indicates that *Achillea* flowers harbor still unknown and obviously prominent fungi, which still need to be identified. Leaves of *Hamamelis* were greatly dominated by *Ascomycota*, as were the later leaf stages of *Achillea*. In soil, also *Ascomycota* were abundant but different orders than on *Hamamelis* leaves. The ascomycotal taxa detected on *Hamamelis* and *Achillea*, such as *Didymella*, *Microsphaerella*, *Cladosporium*, *Microcyclospora*, and *Uwebraunia*, are widespread among many different plants like *Cannabis*, wheat, sugar cane and olive tree (Hardoim et al., 2015; Grudzinska-Sterno et al., 2016; Brader et al., 2017; Gomes et al., 2018; Hamonts et al., 2018). These fungi include saprophytes but also a few potential pathogens (Persoh, 2015; Jia et al., 2016; Thapa and Prasanna, 2018). The *Basidiomycota* in the *Achillea* microbiome consisted predominantly of yeasts such as *Leucosporidium* and *Filobasidium*, which were present on leaves during the earlier stages and later on flower buds and flowers. Basidiomycotal yeasts can reduce infection and sporulation of fungal pathogens (Elad et al., 1994) and thus may act as protectants for *Achillea*. Yeasts of both fungal phyla on flowers appear to be protective and beneficial for floral health by suppressing growth of pathogenic fungi and bacteria and by producing volatile organic compounds attracting pollinators, but are little studied so far (Klaps et al., 2020). The predominance of ascomycotal genera such as *Didymella* and *Septoria*, already reported for *Achillea* (Hatamzadeh et al., 2018, 2020), on the later leaf stages may, in fact, indicate a stronger impact of these pathogens on the aging leaves. *Basidiomycota* seem to be less widespread among plants and constitute lower fractions of fungal phyllosphere communities than *Ascomycota* (Jin et al., 2015; Grudzinska-Sterno et al., 2016; Cregger et al., 2018; Hamonts et al., 2018). Thus, there is a need to intensify studies on *Basidiomycota*, which appear to be more important interaction partners with other plant-associated microbes and the host plant than assumed previously (Martin et al., 2015; Sidorova and Voronina, 2019).

**Core Taxa of Achillea**

The identification of core microbial communities of plants is critical to better understand the significance of the core taxa for the health of the host plant and its compartments and maybe of the functional roles and mutual interactions of these core taxa with the respective host plant (Shade and Handelsman, 2011).

Microbial core communities of plants so far have been rarely investigated and their epi- and endophytic communities have never been distinguished. Hamonts et al. (2018) studied the core bacterial and fungal communities in the leaf, stalk and rhizosphere of sugar cane of different regions in Queensland, Australia. The bacterial core communities of switchgrass and miscanthus of different plots in an experimental area of biofuel crops in Michigan, United States, were studied by Grady et al. (2019) and bacterial core communities of apple flowers of six trees of the same cultivar in an agricultural research station in Wisconsin, United States (Shade et al., 2013). The bacterial core community of the phyllosphere of the Manuka tree (*Leptospermum scoparium*) from five different regions in New Zealand was studied by Noble et al. (2020).

We assessed the bacterial and fungal communities of *Achillea* at four different locations in summer when the plant was fully developed. Hence, we were able to identify the bacterial and fungal epi- and endophytic core communities. Our findings revealed that the epiphytic bacterial and fungal core communities were more diverse than the endophytic communities on leaves and flowers, except for fungi on leaves, even though there was a considerable overlap in the taxa of both communities. The core communities constituted a systematically much higher proportion, i.e., >75% of the total abundance as compared to the total number of taxa, and this difference was consistently higher for the endo- than for the epiphytic communities. A similar finding was reported for the bacterial core community of Manuka trees (Noble et al., 2020), for bacterial and fungal core communities of sugar cane (Hamonts et al., 2018), and switchgrass and miscanthus (Grady et al., 2019), thus, emphasizing the general significance of the microbial core communities for the plants. Our findings further imply, in agreement with previous work on mangrove ecosystems (Yao et al., 2019), that the highly abundant endophytic microbial core communities have a generally greater impact on the interactions with the host plant than the epiphytic core communities. The latter reveal a higher diversity, assuming that they are more affected by local effects such as transmission of location-specific taxa, weather and microclimatic conditions. The differences between the epi- and endophytic microbial core communities also indicate that our methodological separation of both communities was effective. However, we cannot rule out that it did not always remove all attached bacteria from the surfaces during the washing procedure and some overlap between both fractions might still occur.

The bacterial and fungal core communities comprised quite a few rare taxa, which were not among the most abundant ones. This suggests that these members of the rare phyllosphere do play a significant role in the mutual interactions among the microbes and with the host plant. This has been hypothesized for rare microbial taxa in a habitat context in general (Jousset et al., 2017) and emphasizes that these taxa should not be neglected in such investigations. Consequently, future work needs to elucidate their specific functions within the phyllosphere.

The bacterial core community comprised genera, which occurred on leaf and flower in the epi- as well as endophytic
primarily Lactobacillales these habitats. The flower-exclusive core community comprised indicates that some taxa were distinctly adapted to one of these habitats. The flower-exclusive core community comprised primarily Lactobacillales and Enterobacteriaceae and some of them were exclusive to the epi- or endophytic compartment indicating their very specific adaptation to these habitats rich in easily degradable carbohydrates. One mode of transmission of the flower-associated taxa certainly was insect pollination as indicated by the flower-specific epiphytic Gilliamella, well known from the gut of bees, hornets and bumble bees (Moran, 2015; Graystock et al., 2017; Suenami et al., 2019; Zhang et al., 2020). Interestingly, we also detected Gilliamella on leaves of Hamamelis and on leaves of Achillea in spring, assuming a transmission by insect feces or resting insects, as both plants did not flower at this time.

Several fungal core taxa occurred both in the epi- and endophytic leaf and flower compartments. They affiliated to Mycosphaerellaceae, Didymellaceae and other ascomycotal families but also to basidiomycotal families such as Sporidiobolaceae and Bulleraceae. Quite a few fungal core genera were exclusive to leaves and four to the flower including the highly abundant phytopathogenic ascomycotal genus Botryosphaeria and three basidiomycotal genera, exclusive in the epiphytic flower. Only few of these core genera were reported in a comparable study by Hamonts et al. (2018) on the core fungal community of sugar cane, such as Bullera, even though most core taxa affiliated to orders also found in our study. Several fungi of the core community have been reported to inhibit growth of pathogenic microbes and to produce volatile organic compounds attracting pollinators such as Aureobasidium and Sporobolomyces (Golubev and Nakase, 1997; Caruso et al., 2020; Klaps et al., 2020).

As our study on the fungal and bacterial core communities of Achillea represents pioneering work, it is too early to draw conclusions on a more general significance of these core communities. Taxa of the core communities certainly are most important for the health of the host plant, in our study the MP Achillea, presumably including growth-promoting and protecting features but also by controlling potentially pathogenic microbes of the core communities. Further, our results show that it is worth distinguishing between epi- and endophytic communities as we found distinct differences between both communities.

The Microbiome of Achillea and Hamamelis Under Medicinal Aspects and Outlook

Both MPs we studied are well known for various applications, and a variety of bioactive compounds have been identified in Achillea (Candan et al., 2003; Saeidnia et al., 2011; Dias et al., 2013; Hatamzadeh et al., 2020; Apel et al., 2021) and Hamamelis (Da Silva et al., 2000; Sænchez-Tena et al., 2012; Duckstein et al., 2012a; Rocasalbas et al., 2013). Most of these studies focused on the bioactive compounds and their potential application, whereas only one study assessed the chemical transformation by the natural microflora upon spontaneous fermentation (Duckstein et al., 2012a). In the present investigation we comprehensively characterized the epi- and endophytic bacterial and fungal microbiome of above-ground compartments of Achillea and of the leaves of Hamamelis. The results show distinct microbiomes of each MP and seasonal variations. Among the bacteria and fungi in particular of the endophytic compartments taxa were identified of which certain species are known to produce bioactive compounds including ones with antimicrobial and antioxidant activity, which appear to be important in establishing and maintaining the health of the host plant. These microbes interact with each other and the host plant, and it has been shown that such interactions may be instrumental for the production of bioactive compounds (Caruso et al., 2020). Hence, it is highly probable that the microbiome of both MPs investigated affects or may even be instrumental in shaping the bouquet of the bioactive compounds the MP produced and which are key to the medical treatment. The microbiome and/or single microbial taxa of both MPs presumably are also important in further processing these MP and their extracts, e.g., by fermentation as already shown for other MPs and their microbiomes (Schwarzerberger et al., 2012; Lorenz et al., 2013; Duckstein and Stintzing, 2015; Hussain et al., 2016). Thus, future research needs to elucidate the role of the microbiome in the production and conversion of host plant-specific bioactive compounds.

DATA AVAILABILITY STATEMENT

The bacterial 16S rRNA gene and fungal ITS amplicon sequencing data have been deposited in the European Nucleotide Archive (ENA) at EMBL-EBI under accession number PRJEB40947 (https://www.ebi.ac.uk/ena/browser/view/PRJEB40947).

AUTHOR CONTRIBUTIONS

FS, DK, SS, and MS designed the study. SS carried out the practical work, sample processing, and data analyses. LD carried out the bioinformatics and statistical analyses. SS and MS wrote the manuscript. All authors reviewed the manuscript and approved it for publication.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.696398/full#supplementary-material

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Conflict of Interest: SS, DK, and FS were employed by company WALA Heilmittel GmbH.

The remaining 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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