Culture-Dependent and Culture-Independent Characterization of the Olive Xylem Microbiota: Effect of Sap Extraction Methods

Manuel Anguita-Maeso¹, Concepción Olivares-García¹, Carmen Haro¹, Juan Imperial², Juan A. Navas-Cortés¹ and Blanca B. Landa¹*

¹ Institute for Sustainable Agriculture, Spanish National Research Council (CSIC), Córdoba, Spain, ² Institute of Agricultural Sciences, Spanish National Research Council (CSIC), Madrid, Spain

Microbial endophytes are well known to protect host plants against pathogens, thus representing a promising strategy for the control of xylem-colonizing pathogens. To date, the vast majority of microbial communities inhabiting the olive xylem are unknown; therefore, this work pursues the characterization of the xylem-limited microbiome and determines whether the culture isolation medium, olive genotype, and the plant material used to analyze it can have an effect on the bacterial populations retrieved. Macerated xylem tissue and xylem sap extracted with the Scholander chamber from olive branches obtained from two cultivated and a wild olive genotypes were analyzed using culture-dependent and -independent approaches. In the culture-dependent approach using four solid culture media, a total of 261 bacterial isolates were identified after performing Sanger sequencing of 16S rRNA. Culturable bacteria clustered into 34 genera, with some effect of culture media for bacterial isolation. The cultivated bacteria belonged to four phyla and the most abundant genera included Frigoribacterium (18.8%), Methylobacterium (16.4%), and Sphingomonas (14.6%). On the other hand, in the culture-independent approach conducted using Illumina MiSeq 16S rRNA amplicon sequencing [next-generation sequencing (NGS)] of the xylem extracts, we identified a total of 48 operational taxonomic units (OTUs) belonging to five phyla, being Sphingomonas (30.1%), Hymenobacter (24.1%) and Methylobacterium (22.4%) the most representative genera (>76% of reads). In addition, the results indicated significant differences in the bacterial communities detected in the xylem sap depending on the genotype of the olive tree studied and, to a minor extent, on the type of sap extraction method used. Among the total genera identified using NGS, 14 (41.2%) were recovered in the culture collection, whereas 20 (58.8%) in the culture collection were not captured by the NGS approach. Some of the xylem-inhabiting bacteria isolated are known biocontrol agents of plant pathogens, whereas for others little information is known and are first reported for olive. Consequently, the potential role of these bacteria in conferring olive tree protection against xylem pathogens should be explored in future research.

Keywords: microbiome, xylem, culture, next-generation sequencing, vascular pathogens
INTRODUCTION

Olive tree (*Olea europaea* L.) is one of the oldest cultivated trees and has been part of traditional Mediterranean agriculture ever since Roman times. Totaling over 2,000 cultivars (Lavee, 1990), the culture spread from Asia along Syria, Iran, and Palestine to the rest of the Mediterranean basin ca. 6,000 years ago. Since then, olive oil has been a food staple of Mediterranean countries (Müller et al., 2015). Olive oil is considered a high-quality food with multiple beneficial effects for human health, mainly due to its high content in polyphenols, which has induced an increase in olive oil demand and trade worldwide (Landa et al., 2019). Nowadays, world olive cultivation is estimated to cover 10.8 million ha, which results in an olive oil production of about 3.05 million tons, of which ca. 9.5 million ha of olives are grown in the Mediterranean region, accounting for 95% of the cultivated olive area worldwide (i.e., about 98% of the olive oil and 80% of table olive production are from Mediterranean countries) (IOC; www.internationaloliveoil.org; FAOSTATS, http://www.fao.org/faostat/). Olive crop is of great value within the Mediterranean Basin, being critical for the sustainment and maintenance of Mediterranean ecosystems as an integral part of their landscape. Moreover, olive crop is particularly well adapted to cultivation in less accessible areas, including mountain slopes and hillsides, where it helps control soil erosion by reducing the surface runoff of soil and increases soil fertility and nutrient retention (Gómez et al., 2014). Finally, as a major landscape player, it contributes to the establishment of ecological niches for different organisms, thus helping in the maintenance of biodiversity (Rey, 2011). Consequently, since olive is a “high natural value” agricultural system with relevant economic and environmental roles, representing an important part of the heritage and sociocultural life across the Mediterranean, its cultivation should be maintained and preserved.

Nowadays, the health of the olive groves is being seriously threatened, as consequence of a notable increase, both in extent and in severity, of diseases caused by various pathogens, which are capable of adversely affect their growth and production. Among olive diseases, those caused by the vascular plant pathogenic bacterium *Xylella fastidiosa* (specifically from subspecies *multiplex* and *pauca*) and the soilborne vascular fungus *Verticillium dahliae* are, without a doubt, global threats for olive production worldwide (Jiménez-Díaz et al., 2011; Saponari et al., 2018; Almeida et al., 2019; Landa et al., 2019). Xylem vessels are considered ideal niches for microbial endophytes (bacteria or fungi; both beneficial or pathogenic) by providing an effective internal pathway for dispersion throughout the plant and a continuous source of nutrients (McCully, 2001). However, despite significant progress in research on plant microbiota over past years, only a few number of publications have revealed the nature and role of the xylem microbiome, and its relationship to plant health and crop productivity (e.g., Deyett et al., 2017; Fausto et al., 2018; Deyett and Rolshausen, 2019). This fact may be due to microbiome complexity, technical difficulties in the isolation of the xylem-inhabiting microorganisms, as well as to the myriad of biotic and abiotic factors that may affect and determine the microbial community composition and their interaction within the host plant (Vandenhoornhuyse et al., 2015).

With olive, most microbiome studies have focused on determining the microbial composition of the rhizosphere (Mercado-Blanco et al., 2004; Berg and Hallmann, 2006; Mendes et al., 2007; Aranda et al., 2011; Prieto et al., 2011; Montes-Borrego et al., 2013; Caliz et al., 2015; Gómez-Lama Cabanás et al., 2018), whereas only a few studies have focused on the endosphere or xylem microbiomes (Müller et al., 2015; Fausto et al., 2018; Sofo et al., 2019). Unraveling the microbiome of the olive xylem sap should provide a better understanding of the microbes that systemically move throughout the plant or are horizontally transmitted from plant to plant during vegetative propagation. Combined use of culture-dependent and culture-independent approaches such as next-generation sequencing (NGS) technologies may provide a better characterization of the xylem microbiome composition (Turner et al., 2013; Berg et al., 2014) than that obtained when using each approach independently. An essential, parallel step would be to determine the proportion of the microbiome that can be easily isolated and cultured, in order to be subsequently tested and potentially exploited as biocontrol agents of the xylem-inhabiting plant pathogens, such as *X. fastidiosa* or *V. dahliae*. These resident microorganisms could act through direct inhibition or through niche displacement of those pathogens, as commensals and symbiotic organisms that can support the olive immune system, and/or as plant growth promoting agents.

Our research was designed to address this gap in the knowledge by characterizing the bacterial taxa that shape the olive xylem sap microbial communities by using culture-dependent and culture-independent approaches. We also determined in which extent the culture isolation media and the type of plant extract or olive genotype used to isolate the xylem sap bacterial microbiome have an effect on the characterization of its composition, diversity, and structure. Knowledge of the effect of those factors may be essential to identify, isolate, and establish a bacterial collection that might be used subsequently as biocontrol agents against xylem-inhabiting plant pathogens.

MATERIAL AND METHODS

Sampling of Olive Trees

Branches (ca. 35 cm long, 2 years old) from 8-year-old cultivated olive (*O. europaea var. europaea*) trees of “Picual” and “Arbequina” cultivars were sampled from an experimental rainfed field located at the Institute for Sustainable Agriculture from the Spanish National Research Council (IAS-CSIC) in Córdoba (Southern Spain). Trees of both varieties have been submitted to the same agricultural practices throughout the years. Cv. Picual has been found to be highly susceptible to Defoliating (D) and susceptible to Nondefoliating (ND) *V. dahliae* isolates, respectively, while cv. Arbequina has been shown to be susceptible to D and moderately resistant to ND *V. dahliae* (Calderon et al., 2014). In addition, clones of a wild olive tree of *O. europaea var. sylvestris* “Acebuche” known to be moderately resistant to *D. dahliae* infection from a collection of wild-olive genotypes at IAS-CSIC was also sampled. Three 35-cm-long terminal branches (one per tree) from independent
“Picual” and “Arbequina” cultivated olive trees and from clone trees of “Acebuché” were sampled in December 2018 to perform xylem sap extraction with the Scholander chamber. Xylem sap was extracted at the end of Autumn, a season of the year when the olive stem water potential is more stable (Iniesta et al., 2009) and lower than 40 bar of pressure, the maximum pressure allowed by the Scholander chamber device that was used to extract the xylem sap (see below). Likewise, for wood chips collection, a similar number of branches (one branch per tree, with three trees in total per olive genotype) were sampled, and 6-cm-long stem portions were selected from each branch (three pieces per branch). All pruned branches were placed in sterile plastic bags, sprayed with distilled water and kept in a cold room at 4°C to avoid desiccation until later processing in the same day.

**Microbiome Extraction From Xylem**
The three 6-cm-long pieces of branches from each sample were debarked with a sterile scalpel. Bark tissue or xylem chips were obtained by scraping the debarked woody pieces with a sterile scalpel. A total of 0.5 g of xylem chips was weighted by mixing the chips obtained from all the samples from same branch and tree and placed in a Bioreba bag containing 5 ml of sterile phosphate-buffered saline (PBS). Bioreba bags were closed with a thermal sealer and the content was homogenized with a hand homogenizer (BIOREBA, Reinach, Switzerland). Extracts were stored at 4°C until plating onto culture media, and then at -80°C, until DNA extraction. A total of three replicates per olive tree within each genotype was processed. All the processes described above took place under sterile conditions within a flow hood chamber.

Xylem sap extraction from olive branches was performed with a Scholander pressure chamber connected to a nitrogen cylinder following the Bollard process described by Alexou and Peuke (2013). An external port allowed switching from the internal Scholander chamber to an external 60-cm-long super chamber admitting a maximum of 40 bar of pressure. After inserting the branch in the super chamber, 5 cm of the main stem protruded to the exterior of the lid. At this point, it was important to preserve the branch with no cuts or loose leaves that could compromise the pressure within the chamber. To avoid microbial contamination of the xylem sap from bark and phloem, 2 cm of the main stem was debarked and disinfested. The edge between bark and xylem tissue was covered with parafilm to avoid leaks, and the pressure was increased gradually until xylem sap drops were observed to a maximum of 35 bars. The first drops of xylem sap were discarded to avoid external contamination. Xylem sap was collected within a 15 ml sterile falcon tube placed on ice. An average of 10 ml of xylem sap per genotype and branch was obtained and was preserved at 4°C until plating onto culture media, and then kept at -80°C until DNA extraction. All the processes described above took place under sterile conditions within a flow hood chamber.

**Culture-Dependent Characterization of Xylem Bacteria**
For the culture-dependent approach, four solid culture media were evaluated: BCYE (Wells et al., 1981), PD2 (Davis et al., 1981), R2A (Reasoner and Geldreich, 1985), and Nutrient agar (NA) (CONDALAB, Madrid, Spain). R2A and NA are general culture media of low- or high-nutrient contents, respectively. BCYE and PD2 are media specifically designed for isolation of the fastidious bacterium X. fastidiosa from xylem tissues.

Aliquots or 1/10 dilutions (100 µl each) of xylem extracts obtained by each of the two techniques indicated above were plated directly onto three plates of each medium and incubated at 28°C in the dark for 2 weeks. After incubation, all bacterial colonies were counted and a representative number of colonies from each medium and genotype was selected based on abundance and colony feature morphological criteria. Selected colonies were purified by triple serial colony isolation in the same medium. Purified isolates were grown at 28°C in the dark for 1 to 2 weeks depending on their rate of growth prior to DNA extraction.

DNA was extracted from a total of 261 bacterial isolates using the DNeasy kit (QIAGEN, Madrid, Spain). The near-complete 16S rDNA gene was amplified using primers 8f (5’-AGAGTTTGATCCTGGRCTG-3’) and 1492r (5’-ACGGTACCTTGTTACGACTC-3’) (Weisburg et al., 1991) as described in Aranda et al. (2011). Amplicons were purified with ExoSAP-IT (Thermo Fisher Scientific, Madrid, Spain), and directly sequenced in both directions using primers 8f and 1492r (STABVIDA, Caparica, Portugal). Sequences were assembled and manually corrected using DNASTAR software version 15.3.0.66 (Madison, WI, USA). Isolates were identified to genus/species level by the nearest neighbor in the GenBank “nt” database after alignment with reference 16S rRNA gene sequences using the BLAST algorithm according to Altschul et al. (1997).

**Culture-Independent Characterization of Xylem Bacteria**
Aliquots of xylem sap samples (0.5 ml) obtained from macerated xylem chips were placed in PowerBead tubes (DNeasy PowerSoil Kit, QIAGEN) and homogenized 7 min at 50 pulses s⁻¹ with the Tissuelyser LT (QIAGEN). Sap extracts were incubated in the lysis buffer for 1 h at 60°C to increase cell lysis, and then processed following the DNeasy PowerSoil Kit manufacturer's instructions.

Aliquots (8 ml) of xylem sap samples extracted with the Scholander chamber were filtered through a 0.22-µm pore MF-Millipore™ filter (Merck Millipore, Madrid, Spain). Then, filters were placed into 1.5-ml Eppendorf tubes and the filtrate resuspended by vortexing for 5 min in the DNeasy PowerSoil Kit lysis buffer, incubated 1 h at 60°C and processed as described before. DNA obtained was quantified and outsourced to the Integrated Microbiome Resource (IMR) at Dalhousie University (Canada) to perform V5–V6 amplicon library sequencing with primers 799F (5’-AACMGATTAGATACCCKG-3’) and 1115R (5’-AGGTTGCGCTGTGGT-3’) and paired-end sequenced by using Illumina MiSeq sequencing platform (V3; PE 2x 300 bp). The ZymoBIOMICS microbial standard (Zymo Research Corp., Irvine, CA, USA) and water (no template DNA) were used as internal positive and negative controls, respectively, for library
construction and sequencing. Raw sequence data were deposited in the Sequence Read Archive (SRA) database at the NCBI under BioProject accession number PRJNA574439.

Statistical and Bioinformatics Analysis

To determine the effects of the olive genotype, type of xylem sap extraction procedure, and culture media, data of culturable bacterial populations obtained in the culture-dependent approach were subjected to analysis of variance (ANOVA) using the GLM (General Linear Models) procedure in Statistical Analysis System v. 9.4 (SAS Institute Inc.). Data of culturable bacterial population were log-transformed to fulfill ANOVA assumption. The experiment had a completely randomized design, with olive genotype, type of xylem sap, and culture media as factors with three replications (plates) per experimental unit. Data fulfilled the assumptions for ANOVA according to proper statistics. Orthogonal single-degree-of-freedom contrasts were computed to test the effect of selected experimental treatment combinations.

The 16S rRNA sequences obtained were analyzed using the Quantitative Insights Into Microbial Ecology bioinformatics pipeline, QIIME2 (version 2018.11; https://view.qiime2.org/) (Caporaso et al., 2010; Bolyen et al., 2018) with default parameters unless otherwise noted. DADA2 pipeline was used for denoising raw fastq paired-end sequences and filtering chimeras. Operational taxonomic units (OTUs) were obtained at 1% of dissimilarity and were taxonomically classified using RDP Bayesian classifier (Wang et al., 2007) against Silva SSU v.132 reference database. Singletones were discarded for taxonomy and statistical analyses.

Differences among bacterial communities derived from the culture-independent approach were calculated in QIIME2 using rarefaction curves of alpha-diversity indexes (including Shannon, Simpson, Faith_PD, and Richness) at the genus level. Alpha and beta diversity as well as alpha rarefaction curves were conducted rarefying all samples to the minimum number of reads found. The Kruskal-Wallis test ($P<0.05$) with FDR correction (Benjamini and Hochberg, 1995) was used to find differences in alpha diversity indexes among the studied factors. Venn diagrams were generated using the “Venn diagram” online tool (http://bioinformatics.psb.ugent.be/webtools/Venn/) and were used to identify shared (core microbiome) or unique taxa for different studies. Venn diagrams derived using the “Venn diagram” online tool (http://bioinformatics.psb.ugent.be/webtools/Venn/) were used to identify shared (core microbiome) or unique taxa for different studies.

RESULTS

Bacterial Abundance and Alpha Diversity Measures

In the culture-dependent approach, bacterial population densities in xylem sap extracted from woody chips ranged from 40 to 1,920 colony forming units (cfu/ml), whereas those obtained from xylem sap samples obtained with the Scholander pressure chamber ranged from 10 to 610 cfu/ml. Only two factors in the study were found to be significant [olive genotype ($F=12.6, P<0.0001$) and type of xylem sap extraction method ($F=22.4, P<0.0001$), representing 16.6% and 29.7% of the mean square error (MSE) in the model, respectively]. Only the interaction type of xylem sap extraction method by olive genotype was significant ($F=35.67, P<0.0001$) representing 47.2% of the MSE in the model.

A significantly higher ($F>32.09, P<0.0001$) bacterial population density was estimated on xylem samples extracted from woody chips compared to xylem sap extracted with the Scholander chamber for “Acebuche” and “Picual.” while the opposite occurred for “Arbequina” ($F=17.13, P=0.0001$) for both xylem extraction procedures (Figure 1). “Arbequina” showed a significantly higher ($F>10.45, P<0.0021$) bacterial population than “Picual” and “Acebuche” when xylem sap was obtained with the Scholander chamber, whereas “Acebuche” showed a significantly higher ($F>9.24, P<0.0037$) bacterial population than “Arbequina” and “Picual” when the xylem sap was extracted from woody chips macerates (Figure 1).

Maximal recoveries of bacterial population were observed with R2A medium (478.9 ± 131.9 cfu/ml), and decreased with BCYE (352.2 ± 101.3 cfu/ml), PD2 (335.0 ± 97.6 cfu/ml), and NA (270.0 ± 82.0 cfu/ml), in that order, although no significant differences were found among culture media ($F=1.98, P=0.1298$) (data not shown).

A total of 261 bacterial isolated were selected for further study from xylem sap and xylem chips extracts after cultivation in NA, PD2, R2A, or BCYE culture media (Figure 1, Table S1). Out of those, 137 bacterial isolates were from xylem sap extracted with the Scholander chamber while 124 were selected from xylem chips extracts. From the 261 isolates, 51 and 117 bacterial isolates were recovered from cultivated olives “Picual” and “Arbequina,” respectively, and 93 bacterial isolates from the wild olive genotype “Acebuche” (Figure 1).

In the culture-independent approach, Illumina MiSeq sequencing analysis resulted in a total of 21,411 good quality reads with an average of 334 bp after removal of chimeras, unassigned or mitochondrial reads. No chloroplasts reads were detected in our samples. A total of 58 OTUs were identified for all treatments, out of which 48 OTUs were retained after construction and sequencing. Raw sequence data were deposited in the Sequence Read Archive (SRA) database at the NCBI under BioProject accession number PRJNA574439.
samples (Maximum values of Good’s coverage of 1.0 were obtained for all corresponded normally to single isolates (and AN, and BCYE, respectively. Those unique genera seven, four and four, and one genera were unique to R2A, PD2 and AN, and BCYE, respectively. Those unique genera corresponded normally to single isolates (Figure S2; Table S1).

Regarding olive cultivars, 13 genera were identified in “Picual,” 22 in “Arbequina,” and 21 in “Acebuche.” Only four bacterial genera (11.8% of the total identified) were shared among the three olive genotypes and were present in more than 50% of all samples (Core bacterial genera: Bacillus, Frigoribacterium, Methylobacterium, Sphingomonas), six were shared between the two cultivated olive genotypes (Brevibacillus, Dermacoccus, Kineococcus, Marmoricola, Micrococcus, Staphylococcus), seven were shared between the wild olive genotype “Acebuche” and “Arbequina” (Annabacterium, Curtobacterium, Frondihabitans, Microbacterium, Modestobacter, Nocardoides, Paenibacillus), and only one (Variovorax) between “Acebuche” and “Picual” (Figure 3). “Acebuche” showed the highest number of unique genera (nine in total). Finally, regarding xylem sap extraction method, 13 (38.2%) of the culturable bacterial genera (Bacillus, Brevibacillus, Curtobacterium, Dermacoccus, Frigoribacterium, Methylobacterium, Micrococcus, Modestobacter, Nocardoides, Paenibacillus, Sphingomonas, Staphylococcus and Variovorax) were isolated by both extraction methods, whereas 11 (32.4%) and 10 (29.4%) of bacterial genera were unique to the Scholander chamber extraction method (Acinetobacter, Amnibacterium, Clavibacter, Corynebacterium, Frondihabitans, Kocuria, Marmoricola, Novosphingobium, Patulibacter, Quadrisphaera, Roseomonas) or the woody chips extraction method (Brachybacterium, Cellulomonas, Enterococcus, Friedmanniella, Kineococcus, Microbacterium, Rhodococcus, Risingbinella, Sporosarcina, Terrabacter) respectively (Figure 4).

When using the culture-independent approach, a total of 5 phyla, 8 classes, 17 orders, 23 families, and 31 genera were identified (Figure 2). Twenty-one of the genera were identified in “Picual,” 14 in “Arbequina,” and 16 in “Acebuche” (Figure 3). Only seven bacterial genera (22.6% of the total identified) were shared among the three olive genotypes (Core bacterial genera: Curtobacterium, Friedmanniella, Geodermatophilus, Hymenobacter, Kineococcus, Marmoricola, Methylobacterium, Sphingomonas), two were shared between the two cultivated olive genotypes (Cutibacterium and Pseudomonas), two were shared between the wild olive genotype “Acebuche” and “Arbequina” (Annabacterium and Mucilaginibacter), and two (Exiguobacterium and Novosphingobium) between “Acebuche” and “Picual.” “Picual” showed the highest number of unique genera (10 in total) (Figure 3). Regarding xylem sap extraction method 14 (45.2%) of the bacterial genera identified were isolated by both extraction methods (Acidibacter, Curtobacterium, Cutibacterium, Exiguobacterium, Friedmanniella, Geodermatophilus, Hymenobacter, Kineococcus, Marmoricola, Methylobacterium, Pseudomonas, Roseomonas, Sphingomonas, Sporosarcina), whereas 14 (45.2%) and 3 (0.10%) of bacterial genera were unique to the Scholander chamber extraction method (1174-901-12, Acidiphilium, Amnibacterium, Arthrobacter, Brevibacillus, Deinococcus, Frigoribacterium, Mucilagibacter, Nocardoides, Novosphingobium, Quadrisphaera, Rathayibacter, Streptococcus, Variovorax) or the woody chips extraction method (Bradyrhizobium, Pectobacterium, Rubellimicrobium), respectively (Figure 4).

Four of the phyla, Actinobacteria, Bacteroidetes, Firmicutes, and Proteobacteria, were obtained both by the culture-dependent and culture-independent approaches. Only the phylum Deinococcus-Thermus emerged exclusively when using the culture-independent
approach (Figure 2). At the Class level Actinobacteria, Bacilli, Alphaproteobacteria, Betaproteobacteria, and Gammaproteobacteria were identified following both approaches. Thermophilia class was detected only in the culture-dependent approach while Sphingobacteria and Deinococci classes were identified only when using the NGS procedure (Figure 2). At the genus level, a total of 51 genera were identified combining both culture-dependent and culture-independent approaches; 20 and 17 genera were exclusive of the culture-dependent and culture-independent approaches, respectively, and 14 bacterial genera were shared by both methodologies (Figure 2).

**Bacterial Abundance Distribution**

Actinobacteria phylum presented the highest relative abundance in the culture-dependent approach (46.37%) followed by Proteobacteria (34.09%), Firmicutes (19.15%), and Bacteroidetes (0.38%) (Figure S3). Differently, Proteobacteria showed more than half of the total bacterial genera in the culture-independent approach (56.25%), followed by Bacteroidetes (24.76%), Actinobacteria (13.44%), Firmicutes (0.77%), and Deinococcus-Thermus (0.21%) (Figure S4). At the family level, the most abundant bacterial families were Microbacteriaceae (34.48%), Methylobacteriaceae (16.86%), Sphingomonadaceae (31.44%), and Bacillaceae (13.03%) when using the culture-dependent approach, whereas Sphingomonadaceae (31.44%), Hymenobacteraceae (24.11%), and Methyllobacteriaceae (22.36%) were the most abundant when using the culture-independent approach. Focusing on the culture-dependent methodology, the most abundant genera were Frigoribacterium (18.77%), Methyllobacterium (16.36%), Sphingomonas (14.56%),
FIGURE 3 | Prevalence Venn diagram showing the unique and shared bacterial genera obtained using culture-dependent approaches (upper panel) or culture-independent approach (lower panel) in olive xylem sap samples when compared by olive genotype (“Acebuche,” “Arbequina,” and “Picual”).
**FIGURE 4**| Prevalence Venn diagram showing the unique and shared bacterial genera obtained using culture-dependent approaches (upper panel) or culture-independent approach (lower panel) in olive xylem sap samples when compared by xylem sap extraction method [Scholander chamber (SCh) or wood chips (WC) maceration].
Bacillus (12.64%), and Curtobacterium (11.88%), while when we used the NGS approach, the most abundant genera identified were Sphingomonas (30.10%), Hymenobacter (24.11%), and Methylobacterium (22.36%) (Figures S3 and S4).

**Bacterial Community Structure**
Hierarchical clustering analysis and PCA using OTU frequencies at the genus level differentiated xylem bacterial communities according to the sap extraction method or the olive genotype irrespective of the culture approach used (Figure 5; Figure S5). However, these differences were more noticeable when using the culture-independent approach, for which there was a clear trend to group the bacterial communities first by olive genotype and then by extraction method with only one exception, and with "Acebuche" showing the most distant bacterial communities as compared to the cultivated olive genotypes.

**DISCUSSION**
This study resulted in an initial characterization of the bacterial microbiome of the xylem sap from two cultivated and one wild olive genotypes following culture-dependent and -independent approaches, thus providing new insights into the olive microbiome profile, especially in a vascular system where some of the most destructive olive pathogens thrive. Our results suggest that the characterization of the xylem bacterial endophyte community composition is strongly dependent on the use of culture-dependent or -independent approaches and on the xylem sap extraction method used (Scholander chamber vs. woody chips). This initial characterization of olive xylem bacterial microbiome represents a first step making inroads into a potentially promising strategy for the identification of prospective biological control agents well adapted to the ecological niche in which these organisms would be potentially applied.

We found that the population of culturable xylem-associated bacteria ranged from $10^3$ to $2 \times 10^3$ cfu/ml, depending on the olive genotype and the extraction method. These population ranges are in agreement with those obtained for other woody crops such as citrus and grapes (Bell et al., 1995; Gardner et al., 1982). Although the estimates obtained per gram of homogenized vessel xylem tissue are not strictly comparable to data obtained for the vacuum Scholander chamber, mainly due to differences in the amount of plant tissue sampled, the population levels obtained when using the former method appeared to be much higher in general, even when the amount of tissue sampled was smaller. Microscopic analysis of xylem samples from grapes and citrus (Bell et al., 1995; Gardner et al., 1982) revealed that many bacteria remain attached to xylem vessel walls with fibrillar material. These observations might also explain our differences in olive, not only quantitatively but also qualitatively, i.e., the differences noticed in bacterial genera retrieved by both extraction methods. It may be possible that certain groups of xylem-inhabiting bacteria show, or certain environmental conditions induce, a preference for planktonic growth as compared to a static-biofilm associated growth on xylem tissue. Since biofilm formation in plants, and specifically on xylem tissues, is associated with both symbiotic (beneficial) and pathogenic responses, these potential differences in growth styles or changes due to environmental conditions within the xylem vessels should be taken into consideration in future studies (Cruz et al., 2012; Bogino et al., 2013).

![Figure 5](image)

**FIGURE 5** | Principal component analysis of the relative abundance of bacterial genera obtained using culture-dependent approaches (left panel) or culture-independent approach (right panel) in olive xylem sap samples when compared by olive genotype (Ace: "Acebuche," Arb: "Arbequina," and Pi: "Picual") or by xylem sap extraction method [Scholander chamber (SCh) or wood chips (WC) maceration].
Olive is one of the most ancient domesticated plants, widely distributed in Roman times and its culture documented since ca. 6,000 years ago (Lavee, 1990). A key feature of cultivated plants is represented by the process of domestication and breeding, with a net reduction of genetic diversity and the promotion of growth with external inputs, which may interfere with the establishment of microbiome assemblages (Doebley et al., 2006). We found differences in the microbiome community composition associated to xylem-sap depending of the olive genotype, indicating a more different bacterial composition of wild olive as compared to both cultivated olive genotypes. Several studies with domesticated crops have indicated differences in microbiome community composition between wild accessions and modern cultivated varieties, mainly studied in herbaceous species (e.g., barley, bean maize, and rice; Peiffer et al., 2013; Bulgarelli et al., 2015; Edwards et al., 2015; Pérez-Jaramillo et al., 2017), less so in woody crops (e.g., Bullington and Larkin, 2015; Deyett et al., 2017; Yang et al., 2017), with only a few studies in olive (Aranda et al., 2011; Montes-Borrego et al., 2013; Montes-Borrego et al., 2014; Müller et al., 2015). Although this study shows some evidence on the existence of significant differences in population densities and community composition of the xylem-limited microbiome according to the olive genotype, this observation should be further explored including a higher number of olive cultivars growing under different environmental conditions.

Our NGS results with olive xylem indicated the presence of endophytic bacteria from the phyla Actinobacteria, Firmicutes, Bacteroidetes, Proteobacteria, and Deinococcus-Thermus. However, while Actinobacteria and Proteobacteria bacteria were present in all three olive genotypes, only “Picual” contained Firmicutes and Bacteroidetes, and only “Acebuch” contained bacteria from the Deinococcus-Thermus phylum. These results are in accordance with those obtained previously following similar approaches from olive leaves and xylem sap (Müller et al., 2015; Fausto et al., 2018; Sofo et al., 2019) as well as from other woody plants, like grapes, oak, poplar, and Pinus (Uroz et al., 2010; Bonito et al., 2014; Cregger et al., 2018; Deyett and Rolshausen, 2019). Interestingly, our results were more in agreement with those obtained by Deyett and Rolshausen (2019) that found similar proportions for Proteobacteria and Actinobacteria on grapevine xylem sap than those on olives obtained by Fausto et al. (2018), when using a similar xylem sap extraction method with a Scholander pressure chamber. Some differences were also found among those studies at the genus level. Thus, Deyett and Rolshausen (2019) found bacteria in the Enterobacteriaceae family, and belonging to Streptococcus, Bacteroides, Bacillus, Acinetobacter, and Pseudomonas genera as the most abundant on grapevine sap, whereas Fausto et al. (2018) identified some bacterial genera in the olive sap common to our study (e.g., Curtobacterium, Hymenobacter, Methylobacterium, and Sphingomonas), although they did not provide their relative abundance. Those differences may be due to the fact that different primers targeting different variable regions of the 16 rRNA were used for amplification in each study, or to the differences on environmental growing conditions and the plant genotypes evaluated.

In our study the bacterial genera detected in the highest proportions were, in this order, Sphingomonas, Hymenobacter, and Methylobacterium. Interestingly, despite the abundance of Hymenobacter determined by NGS, we did not isolate it in culture, although this bacterium has been readily isolated from stem and leaves of other plant species (Izhaki et al., 2013; Ding and Melcher, 2016; Durand et al., 2018; Ginnan et al., 2018). Other abundant genera identified through NGS were Kineococcus and Friedmanniella, both in xylem sap and woody chips. However, they were only isolated in culture from woody chips, which might indicate the presence of different species of cultivable or non-cultivable bacteria of Kineococcus and Friedmanniella in the olive xylem microbiota or a preference of these genera for a stationary growth within the xylem. Cultivated members of these genera had been previously isolated from plant stems (Qin et al., 2009; Tuo et al., 2016) and, as members of the Actinobacteria phylum, they have been proposed to increase agricultural productivity through plant-growth promotion and to have the potential to be used as an alternative to chemical fertilizers (Palaniyandi et al., 2013; Hamedi and Mohammadipanah, 2015).

Traditional approaches for studying the diversity of plant microbial communities have relied on cultured-dependent approximations. Although the advances in -omic and NGS technologies over the last decade have enabled the culture-independent study of plant microbiomes, we believe that both approaches should be used in parallel to provide deeper knowledge and exploitation of the plant-associated microbiome, especially for perennial crops (Cagliardi et al., 2001; Aranda et al., 2011; Mendes et al., 2011; Jackson et al., 2013; Dissanayake et al., 2018). With olive, our results using the culture-dependent approach indicated that endophytic bacteria from phyla Actinobacteria, Firmicutes, Bacteroidetes, and Proteobacteria can be readily isolated from the different plant genotypes. Considering the unique microbial communities found for each olive genotype, however, no Bacteroidetes could be isolated from “Arbequina,” while for “Picual,” unique bacteria belonging to Actinobacteria phylum were isolated. These results are in line with the gross taxonomic bacterial distribution of isolates already identified in the olive rhizosphere and phyllosphere (Ercolani, 1991; Aranda et al., 2011), as well as with the composition of bacterial endophytes identified in other woody plants, such as citrus, grapevine, or poplar (Bell et al., 1995; Compant et al., 2011; Azevedo et al., 2016; Durand et al., 2018). At the genus level, Sphingomonas, Methylobacterium, Curtobacterium, Frigoribacterium, and Bacillus were among the most abundant cultivable genera. Frigoribacterium are endophytic bacteria isolated from poplar trees (Ulrich et al., 2008) and from potato, where they have an antagonistic capacity against phytopathogenic fungi (Berg and Hallmann, 2006). Bacillus bacteria are well-known rhizosphere and endosphere colonizers with diverse antagonistic activities against plant pathogens (Kumar et al., 2012). The widespread occurrence of the Sphingomonas and Methylobacterium genera has been described as crucial in plants due to a diverse range of functions, including degradation of certain contaminants and facilitation of soil nutrient cycling and plant growth (White et al., 1996; Chen et al., 2016). This may explain their ample distribution in the
plant kingdom, from legume species (Khan et al., 2014) to woody plants (Van Aken et al., 2004) and throughout different plant organs, such as seed, leaf, and flower tissues (Kim et al., 1998). Interestingly, *Methylobacterium* isolates have been shown to exhibit antagonistic properties that induce resistance against the attack of diverse fungi (Rajendran et al., 2009; Ardanov et al., 2012) or that, more importantly, can modify the response of the xylem-inhabiting bacterial pathogen *X. fastidiosa* in citrus, resulting in a reduction of Citrus Variegated Chlorosis (CVC) symptoms (Azevedo et al., 2017). Similarly, *Curtobacterium* has been also described in citrus plants as an endophyte interacting with *X. fastidiosa* and reducing the severity of the CVC symptoms (Lacava et al., 2007). All these characteristics make several of the bacterial isolates from xylem sap of olive obtained in this study good candidates for testing their potential to confer protection against the xylem-inhabiting pathogens *X. fastidiosa* and *V. dahliae* in future research.

Although several studies on woody plant species have revealed the power of using culture-independent approaches for revealing previously unappreciated microbial diversity on xylem samples (Müller et al., 2015; Deyett et al., 2017; Fausto et al., 2018; Deyett and Rolshaumen, 2019; Sofo et al., 2019), our study is unique in that culture-dependent and -independent methods were compared directly and simultaneously (i.e., applied to the same samples) in different genotypes of the host plant using two different extraction methods, to perform a quantitative and qualitative assessment of the xylem bacterial microbiome. Although knowing the specific limitations of each of the culture-dependent and culture-independent approaches, and the short number of olive genotypes and samples evaluated, our study is relevant in that it paves the way for future studies on the identification and characterization of olive xylem sap endophytic bacteria that could be involved in plant defense by acting as biocontrol agents against diverse xylem-inhabiting pathogens and by promoting plant health and growth promotion of olive plants. This information, in turn, will be generally useful for innovative microbiome research based both on culture-dependent and culture-independent approaches wherever vascular plant pathogens and nutrient requirements are involved.

**DATA AVAILABILITY STATEMENT**

The raw sequence data have been deposited in the Sequence Read Archive (SRA) database at the NCBI under BioProject accession number PRJNA574439.

**AUTHOR CONTRIBUTIONS**

MA-M and BL: conceived research, performed statistical and bioinformatics analyses, interpreted results, and wrote the manuscript. CO-G, MA-M, and CH: prepared materials and equipment and performed the experiments. JN-C and JI: contributed to reviewing the manuscript and interpreting results. All authors viewed and approved the manuscript.

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**SUPPLEMENTARY MATERIALS**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2019.01708/full#supplementary-material

**FIGURE S1** | Comparison of rarefaction curves of observed OTUs present in the xylem sap of ‘Picual’ (Pic), ‘Arbequina’ (Arb), and ‘Acebuche’ (Ace) olive genotypes extracted with the Scholander chamber (SCh) or from woody chips macerates (WC). Error bars represent standard derivation of three independent tree replicates. Data were rarified to 1,234 sequences.

**FIGURE S2** | Prevalence Venn diagram and table of cases showing the unique and shared bacterial genera obtained using culture-dependent approaches when compared by culture isolation media.

**FIGURE S3** | Bacterial taxonomic abundances (%) from phylum to genus level derived from culture-dependent approach.

**FIGURE S4** | Bacterial taxonomic abundances (%) from phylum to genus level derived from culture-independent approach.

**FIGURE S5** | Hierarchical cluster analysis and heat-map of the relative abundance of bacterial genera obtained using culture-dependent approaches (left panel) or culture-independent approach (right panel) in olive xylem sap samples when compared by olive genotype (Ace: ‘Acebuche’, Arb: ‘Arbequina’ and Pic: ‘Picual’) or by xylem sap extraction method (Scholander chamber (SCh) or wood chips (WC) maceration). The relative abundance of the bacteria are scaled by a color gradient bar.

**TABLE S1** | Bacterial isolates obtained from xylem sap of olive in the present study, indicating the sap extraction method, culture media of isolation and their identification at genus level based on partial sequencing of 16S rRNA and BLAST analysis.
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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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