Microbial Community Succession and Organic Pollutants Removal During Olive Mill Waste Sludge and Green Waste Co-composting

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Olive mill wastewater sludge (OMWS) is the main by-product of the olive industry. OMWS is usually dumped in landfills without prior treatment and may cause several eco-environmental hazards due to its high toxicity, which is mainly attributed to polyphenols and lipids. OMWS is rich in valuable biocompounds, which makes it highly desirable for valorization by composting. However, there is a need to understand how microbial communities evolve during OMWS composting with respect to physicochemical changes and the dynamics of pollutant degradation. In this study, we addressed the relationship between microbial community, physicochemical variations and pollutants degradation during the co-composting of OMWS and green wastes using metagenomic- and culture-dependent approaches. The results showed that in raw OMWS, Pichia was the most represented genus with almost 53% of the total identified fungal population. Moreover, the bacteria that dominated were Zymobacter palmae (20%) and Pseudomonas sp. (19%). The addition of green waste to OMWS improved the actinobacterial diversity of the mixture and enhanced the degradation of lipids (81.3%) and polyphenols (84.54%). Correlation analysis revealed that Actinobacteria and fungi (Candida sp., Galactomyces sp., and Pichia manshurica) were the microorganisms that had the greatest influence on the composting process. Overall, these findings provide for the first time some novel insights into the microbial dynamics during OMWS composting and may contribute to the development of tailored inoculum for process optimization.

Keywords: composting, metagenomic, microbial communities, organic pollutants, olive mill wastewater sludge (OMWS)

INTRODUCTION

The world production of olive oil is estimated at 3 million tons annually, 98% of which are produced in the Mediterranean area (Souilem et al., 2017). The cultivation of olive trees is largely dedicated to oil and table olive production and has significant socioeconomic impacts (Uylaşer and Yildiz, 2014). However, the olive transformation industry generates large quantities of olive mill wastewater
OMWS, which is considered a serious environmental concern in Mediterranean countries (Galliou et al., 2018). Due to the complexity of OMWW, valorization strategies are scarcely implemented, and most industries often resort to storing byproducts in dedicated evaporation ponds (Jarboui et al., 2010; Kavvadias et al., 2010; Rajhi et al., 2018). Such a strategy, although more convenient in economic terms, is flawed because it induces the formation of olive mill waste sludge (OMWS), which is more hazardous than OMWW due to the higher load of pollutants (Kavvadias et al., 2017). The number of evaporation ponds throughout the Mediterranean area is continuously increasing, and their implication on the contamination of the soil and groundwater is frequently reported (Barbera et al., 2013; Mekki et al., 2020).

OMWS is a blackish-red solid material with complex and variable physicochemical characteristics. OMWS could be either acidic or basic depending on the olive quality, ripeness degree and extraction process (Rigane et al., 2015; Bouhia et al., 2020). Moreover, the stability of the generated sludge fluctuates significantly, as several aspects may affect the evaporation process, namely, the wind speed, light intensity and humidity of the area where OMWS is often dried for several weeks or even months depending on the weather conditions (Montero et al., 2015). Limited research has focused on the valorization of OMWS in comparison with OMWW. Successful examples include the production of construction materials (Hytiris et al., 2004) and the production of alternative fuels via pyrolysis of OMWS and waste tires (Grioui et al., 2019). With respect to biological methods, composting stands as a promising technology that has attracted the interest of scientific and industrial communities for the valorization of OMWS. Composting is one of the best approaches employed for the management of solid wastes and the production of stable soil amendments, which is perfectly aligned with the circular economy concept, as it fully integrates both economic and environmental aspects (Pergola et al., 2018). Nevertheless, due to the peculiarities of OMWS, composting remains intricate, and the use of classical approaches seems to be ineffectiveness in removing a large part of the existing organic pollutants (Hachichia et al., 2012; Sáez et al., 2021). Consequently, process optimization is required, which could be plausibly achieved by controlling the composting environment and/or microbial dynamics.

During OMWS composting, the variation in microbial community succession is dependent on the composting phase. The duration of each phase is dependent on the initial composition of organic material, moisture content, quantity, and composition of the microbial community. During the mesophilic phase, the initial decomposition of organic matter is ensured by mesophilic populations such as *Lactobacillus, Acetobacter, Penicillium, Aspergillus,* and *Streptomyces* by using readily available carbon sources at the beginning of the process (Chandna et al., 2013; Hefnawy and Nagdi, 2014; Li et al., 2019; Palaniveloo et al., 2020). As a result of mesophilic microorganism metabolic activity, the resulting heat leads to the transition to the thermophilic phase, where a high rate of biodegradation is recorded. In this phase, the growth of thermophilic organisms such as *Aspergillus,* *Bacillus,* *Actinobacteria,* *Thermoactinomyces,* *Talaromyces,* and *Pseudomonas* is observed (Chandna et al., 2013; Hefnawy and Nagdi, 2014; Li et al., 2019; Palaniveloo et al., 2020). However, it has been found that the most efficient composting process is achieved by both bacterial and fungal communities. The cooling phase, in which mesophilic microorganisms proliferate again in the compost, degrades the remaining organic matter. Many bacterial communities are associated with this phase, such as *Anthrobacter,* *Bacillus,* *Enterobacter,* and *Micrococcus,* as well as fungal genera such as *Alternaria,* *Aspergillus,* *Cladosporium,* *Trichoderma,* and *Verticillium* (Palaniveloo et al., 2020). However, until today, there is a big lack of literature regarding OMWS microbiological structural community that are likely to play a major role during composting hygienization process. The objective of this investigation was to study microbial dynamics during OMWS composting by combining metagenomic and culture-dependent methodologies to pinpoint key microbial species and classes that may be involved in pollutant degradation to optimize the process and improve the quality of the final product.

**MATERIALS AND METHODS**

**Composting Assay**

The co-composting assay was conducted in a bioreactor under controlled conditions. The feedstock was prepared by mixing equivalent quantities (20 kg) of OMWS and green waste (GW) consisting of grass and follicular parts of gardening work (Table 1).

The mixture was homogenized, its moisture level was adjusted between 50 and 60%, and then the mixture was introduced into a 100 L, cylindrical bioreactor in stainless steel, with a superficial layer of insulating polyurethane equipped with a brewing system and air input and output to ensure uniform aeration during the experiment. At each stage of the composting process, 1 kg of the mixture was sampled and stored at -20°C, a difference in temperature is observed (Chandna et al., 2013). The pH of the OMWS was measured at 5.32 ± 0.02 (pHb 5.2); EC was measured at 3.72 ± 0.73 (ECb 3.7); TOC was measured at 50.18 ± 5.1 (TOCb 50.18); TN was measured at 47.64 ± 2.15 (TNC 47.64); and C/N was measured at 57.64 ± 1.2 (CNb 57.64). The co-composting assay was conducted in a bioreactor under controlled conditions. The feedstock was prepared by mixing equivalent quantities (20 kg) of OMWS and green waste (GW) consisting of grass and follicular parts of gardening work (Table 1).

**TABLE 1 | Physicochemical characteristics of OMWS (olive mill waste sludge) and GW (green waste): values are means ± SD of 3 replicates.**

| Parameters                  | OMWS (ppm) | Green waste (ppm) |
|-----------------------------|------------|-------------------|
| Moisture %                  | 53.11 (4.8) | 57.64 (0.7)       |
| pH                          | 4.9 (0.01)  | 7.15 (0.08)       |
| ECb (ms.cm⁻¹)               | 3.72       | 0.73 (0.08)       |
| TOCb %                      | 50.18 (0.14)| 47.38 (0.04)     |
| TNCb %                      | 0.22 (0.01)| 2.15 (0.29)       |
| C/Nb                        | 228        | 22                |
| Total phosphorus %          | 0.126 (0.001) | 0.04 (0.02)     |
| Exchangeable potassium K%   | 1.36 (0.01)| 0.22 (0.02)       |
| Exchangeable calcium %      | 0.58 (0.01)| 0.34 (0.01)       |
| Exchangeable Mg%            | 0.16       | 0.08 (0.01)       |
| Total Cu (ppm)              | 20.58 (0.52)| 2.91 (0.62)      |
| Total Mn (ppm)              | 35.76 (0.36)| 38.49 (0.18)     |
| Total Fe (ppm)              | 1563.25 (38.81)| 800.42 (156.9)  |
| Total Zn (ppm)              | 58.86 (3.37)| 24.24 (0.8)       |

<sup>a</sup> Results expressed per unit weight fresh matter.

<sup>b</sup> Results expressed per unit weight dry matter.
supernatant was recovered and filtered through a 0.45 µm filter. Then, a volume of 50 ml of distilled water was added to 1/10 mixture of compost and distilled water. The total organic carbon (%TOC) was dosed by organic matter oxidation using K₂Cr₂O₇ according to Walkley and Black (1934). The ash content was calculated after calcination at 600°C for 6 h. Total nitrogen (TKN) was assayed with 0.5 g dried samples by using classical Kjeldahl distillation according to AFNOR methods.

Phenols were measured based on the water extracts, successively treated with petroleum ether and ethyl acetate in the presence of ammonium sulfate and phosphoric acid, and the polyphenols were recovered with pure methanol. Quantitative measurements were carried out through colorimetric methods according to Box (1993) using Folin and Ciocalteu’s reagent at 760 nm. Lipid content was directly extracted from mixture samples using a dichloromethane/methanol solution with a 1:1 ratio.

Phytotoxicity was evaluated according to Zucconi et al. (1981) and Paulino et al. (2006) with slight modifications. Four grams of dry material from each composting phase was moistened with water and then treatment for 10 to 15 min by sonication (El Fels et al., 2015; Bouhia et al., 2021). Afterward, the suspensions were serially diluted to 10⁻⁹. The pH of each co-composting sampled mixture was adjusted to be the same.

For Actinobacteria enumeration, actinomycete isolation agar (AIA) (Sigma) was supplemented with 50 µg/ml cycloheximide and 10 µg/ml nalidixic acid to inhibit fungi and gram-negative bacteria without affecting the growth of Actinobacteria. Bacterial community enumeration was performed using standard medium (nutrient agar) with the addition of cycloheximide (Sigma). For estimation of the fungal community, suspension calibration was performed in potato dextrose agar (PDA) (Panreac) by the addition of 5 µg/ml chloramphenicol. Colony forming units (CFU) were counted in triplicate, and the results are expressed as CFU g⁻¹ dry weight. The pH of each medium was adjusted to 5.6; 5.7; 5.9; 6.5; 7.4; and 8, respectively, for 0, 4, 9, 21, 52, and 120 days to be similar to that of each composting phases. Samples were incubated at both mesophilic (35°C) and thermophilic (45°C) temperatures.

Metagenomic Approach: DNA Extraction, Gene Amplification, and Sequencing

To study the variation in the total microbiota (bacteria and fungi) during the co-composting phases (mesophilic, thermophilic and maturation phases), we performed comprehensive metagenomic analysis. DNA from the samples (≈ 1 g of compost from each phase) was extracted by using a DNAeasy PowerSoil Pro Kit (Qiagen Inc.) following the manufacturer’s instructions. Then, the quantity and quality of the extracted DNA were evaluated spectrophotometrically using a NanoDrop 2000c before being processed for 16S and ITS amplicon sequencing. Sequence libraries were generated using the complete primers ITS1 (5′-CTTGGTACTTTAGGAAGTA 3′) and ITS2 (5′-GCTGCGTCTTCCATCGATG 3′) for fungi. For bacteria, the 16S rRNA gene V4 variable region PCR primers 515/806 were used in a 30–35 PCR cycle using HotStarTaq Plus Master Mix Kit (Qiagen Inc.). PCR was performed under the following conditions: 95°C for 5 min, then 30–35 cycles at 95°C for 30 s, followed by 53°C for 40 s and 72°C for 1 min. Finally, a final elongation was performed at 72°C for 10 min. Afterward, samples were multiplexed using distinctive dual indices and equally pooled together based on their molecular weight and DNA concentration. Then, they were purified using calibrated Ampure XP beads and used to prepare an Illumina DNA library. Sequencing was performed with MR DNA (Shallowater, TX, United States)¹ on a MiSeq following the manufacturer’s guidelines, and sequence data were processed using MR DNA analysis pipeline (MR DNA, Shallowater, TX, United States).

Briefly, sequences were joined, and sequences < 150 bp or with ambiguous base calls were removed. Sequences were then quality filtered (error threshold of 1.0) and dereplicated. The dereplicated or unique sequences were denoised, and unique sequences identified with sequencing and/or PCR point errors were removed, followed by chimera removal. thereby providing

¹www.mrdnalab.com

notably at T0 (initial mixture), T4 (after 4 days of composting), T9 (after 9 days of composting), T21 (mixture recovering from the bioreactor, after 21 days), T52 (after 52 days of the composting process; maturation in perforated bags) and T120 (after 120 of the composting process; maturation in perforated bags). The resulting compost product was equal to 14 kg.

Phytotoxicity Analysis

Phytotoxicity was evaluated according to Zucconi et al. (1981) and Paulino et al. (2006) with slight modifications. For estimation of the fungal community, suspension calibration was performed in potato dextrose agar (PDA) (Panreac) by the addition of 5 µg/ml chloramphenicol. Colony forming units (CFU) were counted in triplicate, and the results are expressed as CFU g⁻¹ dry weight. The pH of each medium was adjusted to 5.6; 5.7; 5.9; 6.5; 7.4; and 8, respectively, for 0, 4, 9, 21, 52, and 120 days to be similar to that of each composting phases. Samples were incubated at both mesophilic (35°C) and thermophilic (45°C) temperatures.
a denoised sequence or zOTU. Final zOTUs were taxonomically classified using BLASTn against a curated database from the NCBI.2

Statistical Analysis
The results are the mean of 3 values, and measurement data are shown as the mean ± standard deviation (SD). Statistical analysis was carried out using IBM SPSS Win software version 20. Principal component analysis and Pearson's correlation were executed using Excel Analyze-it 1.56.

RESULTS
Evolution of the Physicochemical Parameters
The temperature evolution during OMWS-GW co-composting showed a classical dynamic represented by the succession of mesophilic, thermophilic and maturation phases. The ambient temperature of the composted mixture on the first day of this experiment was 23°C (Figure 1A), and after only 2 days, the temperature reached 42°C and then 54°C after 7 days. The thermophilic phase was maintained for 17 days. Between days 17 and 26, the temperature steadily decreased until reaching an ambient value of 23°C, which was maintained until the end of the composting process.

The initial pH value was 5.57, which then increased to 6.5 on the 21st day, corresponding to the end of the cooling phase (Table 2). The final pH value was 8, resulting from the high decomposition rate (DR)% of organic matter, which exceeded 62% after 120 days of composting (Table 2). For the EC, the initial value of the mixture was 2.9 mS cm⁻¹, and it then followed the same trend as that of the pH, as it reached 3.8 and 4.2 mS cm⁻¹ after 21 and 120 days, respectively.

The TOC decreased progressively during 120 days of composting (Table 2). The TOC contents decreased from 46.23 to 42.24% after 21 days and then further decreased to 37.7% at the end of the process. The initial nitrogen content (0.41%) significantly increased during the thermophilic phase by 173%. Thereafter, it gradually stabilized at a lower value (0.41%) at the end of the process. The initial nitrogen content (0.41%) was recorded.

Evolution of Toxicity
Co-composting of OMWS-GW resulted in a significant reduction in polyphenols and lipids, which constitute the main toxic organic compounds in OMWS. The water-soluble polyphenol (WSPP) concentration and total extract lipid (TELP) variations during composting followed a zero-order kinetic model (Figure 1B). The curve fitting of the experimental data gave the following equations:

\[
\begin{align*}
\text{WSPP} & = 0.695t + 11.9014 \quad (R^2 = 0.853) \\
\text{TELP} & = 0.5514t + 24.418 \quad (R^2 = 0.763)
\end{align*}
\]

For both fractions, a steady decrease was registered throughout composting. Two distinct phases were observed: a fast degradation dynamic during the thermophilic phase, which resulted in the elimination of nearly 50% WSPP and 60% TELP. Then, a slow degradation dynamic until the end of the experiment resulted in a maximum decrease of 81.30 and 84.54% for TELP and WSPP, respectively. Evaluation of the germination index (GI) showed a positive correlation with TELP and WSPP degradation dynamics. Indeed, after the 4th day (Figures 1C,D), the GI values were 8.13 and 0.6 for Cress and Turnip, respectively (highly phytotoxic GI < 50%), showing the high toxicity of the compost during the initial composting phase, which was maintained even after the thermophilic phases for the two-studied species, with values of 47 and 4.6% for Cress and Turnip, respectively. However, at the end of the maturation phase, the phytotoxic effects were completely alleviated as the GI reached 110 and 82% for Cress and Turnip, respectively.

Change in Microbial Diversity During Olive Mill Wastewater Sludge-Green Waste Co-composting
Change in the Microbial Communities Through a Culture-Dependent Approach
The evolution of the total mesophilic and thermophilic microbiota during the composting process (Figure 2) was correlated with temperature variation. The total mesophilic microbiota showed a maximum increase at day 4 from 3.53 × 10^7 to 26.61 × 10^7. Then, a gradual decrease was observed, and a value of 2.77 × 10^7 was reached after 4 months of co-composting. For the thermophilic microbiota, the optimal increase in enumerated microorganisms (from a value of 6.54 to 29.7 × 10^6) was determined at day 9, corresponding to the thermophilic phase. Later, a decrease to 3.9 × 10^6 by the end of the co-composting process was recorded.

During the biodegradation phase (Figure 2A), the mesophilic microbiota was initially present. Once the temperature surpassed 40°C, thermophilic microorganisms became increasingly dynamic and progressively replaced the mesophilic microorganisms. From days 0 to 9 of the composting process, mesophilic Actinobacteria showed a decrease in total abundance from 16.56 to 9.15%, while thermophilic Actinobacteria showed an increase from 6.88 to 15.72%). In the bacterial community, the total abundance was less than 7% for both categories, with a successive increase from 4.63 to 6.46% and decrease from 5.5 to 4.47% for mesophilic and thermophilic bacteria. However, the fungal abundance presented the highest values during the entire process of composting, with variations of 78.8 to 84.37% and 87.61 to 79.79% for mesophilic and thermophilic fungi, respectively, after 9 days of composting.

By the end of the co-composting process, the abundance of mesophilic and thermophilic fungi decreased from 78.80 to 79.79% and 87.61 to 60%, respectively. Nevertheless, a decrease in Actinobacteria abundance from 30.69 and 25% to 20.58 and 15% for mesophilic and thermophilic bacteria, respectively, was recorded.
Change in Microbial Communities Through Metagenomic Analysis

Metagenomic analysis of samples representing the key phases of the composting process (mesophilic, thermophilic and maturation) revealed that bacterial taxonomic diversity initially existing in raw OMWS was mostly dominated by Gammaproteobacteria (52.4%), Bacilli (18%) and Actinobacteria (17.8%). At the species level, bacteria were prevalent, namely, Zymobacter palmae (20%), Pseudomonas sp. (19%), and Lactobacillus acidipiscis (6%). Moreover, fecal bacteria were also identified, such as Staphylococcus sciuri (0.003%), Enterococcus faecalis (0.1%), Enterococcus gallinarum (0.1%), and Enterococcus raffinosus (0.003%), which should be considered potential infectious agents. For fungi, diversity analysis revealed that yeasts were the most abundant genus in raw OMWS (Figure 3). These were mostly represented by the Pichia genus, with almost 53% of the total identified population. The remaining genera were Tortispora (16%), Dipodascus (12%), Rhizophydium (11%), Candida (4.4%), Galactomyces (2.4%), and Scedosporium (0.4%).

In the initial mixture (Figure 3C), the fungal microbiome was dominated by four species (> 5%), namely, Dipodascus australiensis (33.18%), Penicillium roqueforti (27.27%), Pichia sp. (8.17%) and Cryptococcus sp. (7%). Similarly, the bacterial microbiome (Figure 3D) was dominated by Psychrobacter aquaticus (30.3%), Corynebacterium variabile (28.7%), Carnobacterium maltaromaticum (11.5%), and Psychrobacter pulmonis (9.7%).

By the end of thermophilic phase, major changes were observed in taxa abundance (Figure 3C), as the compost was dominated by only two fungal and three bacterial species, namely, Candida freyschussii (87.3%), Sporopachydermia lactativora (10.6%), Pseudomonas syringae (47.8%), Actinobacter sp. (30.3%) and Corynebacterium variabile (6.1%). After 120 days of compost maturation (Figure 3D), the fungal community...
was mainly represented by four species, *Penicillium crustosum* (48%), *Nectria mauritiicola* (20.7%), *Acremonium* sp. (6.7%) and *Guehomyces pullulans* (5.9%). The minor representatives (>1%) were dominated by *Penicillium* sp. (2.9%), *Penicillium roqueforti* (2.9%) and *Penicillium hirsutum* (1.6%). On a related note, the bacterial diversity significantly shifted at the end of the maturation phase compared to the initial mixture. Indeed, fecal bacteria were not detected, and the percentage of each bacterial class decreased except for Actinobacteria, for which a threefold increase (60%) was noted compared to raw OMWS. At the species level, the most dominant bacteria were *Promicromonospora sukumoe* (13.9%), *Pseudomonas stutzeri* (7.1%), *Brevibacterium linens* (7%) and *Arthrobacter protophormiae* (7%), followed by *Isoptericola* sp. (4.6%), *Carnobacterium maltaromaticum* (4%), *Psychrobacter aquaticus* (2.3%) and *Promicromonospora vindobonensis* (2%).

Moreover, alpha diversity was determined for important composting phases by calculating the number of OTUs, Shannon index and Faith’s PD (Figure 4). Higher microbial diversity index was recorded for raw OMWS and samples of the compost maturation phase, while the lower diversity was recorded for samples of the thermophilic phase.

**DISCUSSION**

**Physicochemical and Toxicity Changes During Green Waste-Olive Mill Wastewater Sludge Cocomposting**

**Evolution of Temperature, pH, and Nitrogen**

Temperature evolution during the composting process is related to microbial community changes during the biodegradation of composted wastes, which plays an important role during maturation and stabilization (Ma et al., 2018). The initial increase in pH is attributed to the organic matter richness of the mixture, especially the lipid and phenolic compounds that microorganisms metabolize more quickly, in addition to the low molecular weight organic acids that are volatilized due to the higher temperature of the thermophilic phase (Tsai and Chang, 2019). The final pH value was 8, which is a direct result of the high decomposition rate of organic matter, which exceeded 62% after 120 days of composting, and the elimination of large amounts of secondary metabolites produced by microorganisms during biodegradation, such as acetic acid and butyric acid (Wei et al., 2016). Furthermore, the %TOC decrease during the composting process was attributed to microbial activity, which reduced the organic content in the composted mixture. According to Huang et al. (2019), this effect is related to active thermophilic microorganisms with a great ability to degrade organic carbon and eliminate various organic compounds initially present in the mixture. The enhancement in nitrification might explain the decrease in the \(\text{NH}_4^-/\text{NO}_3^-\) ratio to 0.91, indicating that compost reached maturity based on the standard maturation index reported by Barje et al. (2012).

**Evolution of Toxicity**

After 4 months of composting, the total lipids and polyphenols decreased, which is in alignment with the findings of Hachicha et al. (2009), who reported similar results for OMWS/sesame bark cocomposting. Those authors attributed this reduction to the use of those compounds as a carbon source by indigenous microorganisms through the conversion of phenols to simpler structures or quinones through lignolytic enzymes such as laccase and peroxidase, which are supplied during humic substance polymerization (Ait Baddi et al., 2009). Lipid degradation could be explained by the ability of microbes to breakdown several forms of lipids, such as fatty acids and triglycerides.
The use of the germination index (GI) to evaluate compost maturity by studying phytotoxicity risks for plants is acceptable. Compost is exempt from any potential toxicity to plants if the GI values are greater than 80% (Zhang et al., 2020). Several factors have been previously reported to be responsible for inducing phytotoxicity effect, during the OMWS compost
evaluation, this effect has been attributed to the presence of several compounds such as organic acids and phenol compounds, impacting negatively the seed germination (Hachicha et al., 2009; Filippi et al., 2013). According to Piotrowska et al. (2006), many seed species are sensitive to phenol presence such as tomato, which are related to their own genotype. In additions to the low pH value, the high concentrations of lipids in OMWS, can negatively influence the growth and biological activities of seeds, which was negatively correlated with GI reduction (Paulino et al., 2006; Hachicha et al., 2009). In agreement with our results, Said-Pullcino et al. (2007), concluded that a value of GI superior to 60%, the toxicity could be considered as well treated. Our results revealed that phytotoxic effects remained even after the thermophilic phase, as the PP and Lp contents were still relevant. However, at the end of the maturation phases, the phytotoxic effects were completely alleviated, as the GI% reached 110 and 82% for Cress and Turnip, respectively. Hence, the occurrence of plausible biostimulatory effects was demonstrated.

Microbial Community Structure of the Olive Mill Wastewater Sludge

Microbial diversity analysis showed that raw OMWS was mainly dominated by fungi, which represented nearly 54% of all identified kingdoms (including bacteria). Similar results were recently reported by Martínez-Gallardo et al. (2021), who found that in OMWS sampled from several ponds, the fungal community systematically outnumbered its bacterial counterpart. Furthermore, the same authors revealed that the bacterial community structure of OMWS is highly dependent on physicochemical parameters, notably nutrient content, pH and moisture content. Indeed, Proteobacteria were prevalent in OMWS with neutral pH and high moisture. Actinobacteria were dominant in dry/alkaline OMWS, which is in alignment with our results. Notably, physicochemical traits are not the only factors that influence the microbial diversity and functionality of OMWS. According to Tsiamis and Tzagkaraki (2012), the latter are significantly affected by olive cultivars. Those authors investigated the bacterial profile of OMWS generated from various olive varieties and found only 15% similarity in terms of the identified OTUs. Moreover, the cultivation and harvesting practices had a large influence on the microbial community distribution in OMWS. For example, Carlozzi et al. (2015) showed that fermentative bacteria in OMWS resulting from the early collection and harvesting “hand collection” of the Olea europaea variety were restricted to the two families Peptococcaceae and Sporolactobacillaceae (Kavoulakis and Ngougas, 2011). On another note, the very few studies that thoroughly investigated the fungal diversity in OMWS reported very contrasting results. For example, Martínez-Gallardo et al. (2021) revealed that the OMWS fungal community was dominated by genera such as Fusarium, Aspergillus, Scopulariopsis, Trittirachium, Scedosporium, and Microascus, depending on the pond characteristics. Slama et al. (2021) identified four dominant genera, namely, Nakazawaea, Saccharomyces, Lachancea, and Candida. Our findings were very different from those previously reported, as the dominant fungal genus was Pichia, which constituted half of the identified fungal population, followed by Tortispora, Dipodascus, and Candida. The prevalence of Pichia sp. was not surprising, as several studies have reported a strong correlation between the occurrence of Pichia species and the detoxification of OMWS (Morillo et al., 2008; Sinigaglia et al., 2010; Arous et al., 2016). Moreover, to our knowledge, this is the first time that the Diplodocus genus has been identified in OMWS. This genus was represented solely by the cactophilic Dipodascus australiensis, which is not surprising knowing that the pond from where the sample was taken was surrounded by Opuntia spp. Similarly, Tortispora and Pichia are usually associated with decaying cactus tissue (Lachance et al., 1988). These observations suggest that the community structure of OMWS is highly dependent on the faunal and floral specificities of the sampling sites.

Changes in Microbial Community Structure During Olive Mill Wastewater Sludge/Green Waste Cocomposting Depend on Physicochemical Traits and Microbe-Microbe Interactions

Adding green waste to OMWS significantly affected the composition of the microbial communities, as the diversity of Actinobacteria increased twofold compared to raw OMWS, which is plausible due to the actinobacterial richness of green waste. However, members of the Proteobacteria family (primarily represented by Moraxellaceae) were still the most dominant. Similarly, for fungi, the previously dominant Pichiaea were greatly outnumbered by Microascales, which represented more than 70% of the fungal diversity of the initial mixture. At the end of the thermophilic phase, half of the polyphenols and nearly 60% of total lipids were degraded, which was accompanied by a significant change in both fungal and bacterial diversity, suggesting the existence of a significant correlation between microbial changes and pollutant dynamics during the composting process. Principal component and Pearson correlation analyses (Figure 5 and Supplementary Material 1) revealed that overall pH, temperature and EC were the factors that affected most community structures. Moreover, the pace at which those parameters changed during the composting process was paramount, which was demonstrated by the occurrence of fast and slow phases of polyphenol and lipid degradation. Temperature is particularly important, as this is the first factor to undergo significant changes, which combined with PP content, initially shape the compost microbial community. Our study showed that most of the microbial species were negatively affected by temperature increases, thus demonstrating the prevalence of the mesophilic microbiota. For fungi, only three families were positively correlated with temperature, namely, Saccharomycetaceae, Dipodascaceae, and Pichiaceae (Figure 5A).

Surprisingly, those fungal families were not strongly correlated with polyphenol and lipid removal when taking into consideration the whole composting process. However, this does not exclude a plausible involvement in the degradation of those organic compounds at early stages of composting, notably during the fast degradation phase observed in the first several days (Figure 1B). Indeed, the works of Ben Sassi et al. (2006) and Sinigaglia et al. (2010) previously reported the ability of Pichia and Candida species (Candida dildensisiae, Candida
ernobii, Pichia holstii, and Pichia membranifaciens) isolated from Moroccan OMWS to degrade PP, such as p-coumaric, caffeic and vanillic acids. Additionally, many other authors have reported that OMWS detoxification can be directly attributed to species of the Pichia genus (Morillo et al., 2008; Arous et al., 2016). Similar trends were observed for bacteria, as the only bacterial classes (Clostridia and Gammaproteobacteria) that were positively correlated with temperature had limited effects on PP and Lp removal, as shown by PCA. More importantly, even if Actinobacteria were negatively affected by temperature, some thermotolerant representatives (Supplementary Material) were likely to be involved in PP and Lp degradation. For example, at the end of the thermophilic phase, the actinobacterial composition of the mixture was dominated by Corynebacterium sp., which have been reported as potent PP degraders (Pradeep et al., 2015). Moreover, the highly sensitive Pseudomonas syringae constituted more than 50% of the identified bacterial species, which could be explained by the favorable physicochemical conditions, including the decrease in PP content following microbial removal by fungal and actinobacterial species.

FIGURE 5 | Correlation heatmap and Principal Component Analysis (PCA) of the main parameters during composting treatment: TOC, Total organic carbon; K, potassium; Mn, Manganese; Phenols; TD, Decomposition rate; pH; Ca, Calcium with Fungi (A) and Bacteria (B) genera diversity.
In fact, Pseudomonadaceae was strongly correlated with Actinomycetaceae (0.995), which demonstrates the importance of microbial species interactions during the composting process. Regarding fungi, the end of the thermophilic phase was marked by the prevalence of two species, namely, Candida freyschussii, an oleaginous yeast knowing for its ability to produce lipids from glycerol (Amaretti et al., 2012), and Sporopachydermia, which was previously reported to be a lipolytic microorganism (Fickers et al., 2005; Agnolucci et al., 2013). Following the thermophilic phase, pH and EC were the most determinant factors influencing microbial community structure, and similar trends were observed for fungi and bacteria, as phyla (Actinobacteria and Ascomycota) that were favored by pH and EC increases were positively correlated with WSPP and TLEP. On a related note, the results of the culture-dependent analysis revealed that fungi were probably more active than bacteria during the thermophilic phase, as they represented more than 70% of all cultivated species. Usually, bacteria constitute the majority of the microorganisms during composting, with a greater number than Actinobacteria and fungi. Our findings revealed opposite results, which could be attributed to the antimicrobial effect of OMWS (richness in monomeric and polymeric phenolic compounds). Additionally, fungi are largely known for being directly involved in biodegradation through the production of polyphenol oxidase during composting (Carraro et al., 2014; Rigane et al., 2015; Steinmetz et al., 2019). Overall, few studies have investigated shifts in microbial communities during the cocomposting of OMWS, and most of them could not identify relevant causative factors (Milanović et al., 2019). In our case, even if relative abundancy lacks the functionality aspect, both correlation analysis and culture-dependent assays clearly demonstrated that pH, temperature, EC, PP content and microbial competition are the main factors affecting microbial succession.

At the species level, Pichia manshurica seems to specialize in lipid degradation, and Candida sp. and Galactomyces sp. are likely to be strongly involved in phenol biodegradation (Supplementary Material 2), which agrees with several previous works (Pinedo-Rivilla et al., 2009; Alves et al., 2014; Karimi and Hassanshahian, 2016; Oliveira et al., 2018). The identified bacterial species revealed that Lactobacillus acidipiscis was the agent most correlated with phenolic and lipid biodegradation (Supplementary Material 2), which was previously identified in a work recently done by Uylaşer and Yildiz (2014) and has been identified as a halophilic lactic acid bacterium able to tolerate up to 8% NaCl from black and green olive samples, representing almost 28% of the total identified genus. The same authors showed important lipolytic and pectolytic activity up to 1.09 and 5.29 U ml\(^{-1}\), respectively, as well as a positive decarboxylase activity. To the best of our knowledge, this species has never been identified as a direct agent of phenol degradation, despite its high potential, which will be very useful in the detoxification of organic pollutants in olive mill waste. L. acidipiscis, first described by Tanasupawat et al. (2000), was found in isolates from fermented fish. It has also been isolated from soy sauce mash (Tanasupawat et al., 2002).

**CONCLUSION**

These findings highlight the distinct impact of the composition and functional microbiota of the OMWS substrate. Fungi were the most dominant taxon of the cultivable microorganisms and were dominated by the genera Pichia, Candida, and Galactomyces. The genus Lactobacillus was the most efficient OMWS bacterial taxon with respect to pollutant degradation. Microbial community succession with respect to physicochemical changes during OMWS has been scarcely investigated, and most of the studies could not clearly identify relevant causative factors explaining the prevalence of a specific microbial community during a given composting phase. At the end of the co-composting experiment, polyphenol and lipid content was reduced by 84.54 and 81.30%, respectively, which suggest that the process needs to be further improved as the degradation of contaminants was not fully optimal. Our study combined culture-dependent and metagenomic approaches to identify key microbial species/classes that have significant effect on distinct phases of the composting process. These results are valuable, as they demonstrate that 1. OMWS composting should be tailored taking into consideration the specifics of the collecting ponds, as the microbial profile is highly dependent on-site characteristics; and 2. An inoculation methodology taking into account both inoculum composition and time of inoculation may plausibly improve the degradation rate of organic pollutants though providing a competitive advantage for key species during composting.

**DATA AVAILABILITY STATEMENT**

The metagenomics raw data of the bioproject are available at NCBI under accession number PRJNA784753 (ID: 784753).

**AUTHOR CONTRIBUTIONS**

YB, MH, YO, and KL conceptualized the study. YB, MH, YO, LE, and KL designed the experiments. YB, MH, YO, ME, and KL reviewed the manuscript. YB and KL performed the statistical data analysis. YZ contributed to investment. All authors agreed to be accountable for the content of the work.

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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.2021.814553/full#supplementary-material
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