

Assessment of microbial pools by an innovative microbiological technique during the co-composting of olive mill by-products

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ABSTRACT

Different mixtures of olive pomace (OP), olive mill wastewater (OMWW) and olive pruning residues (OPR) were aerobically co-composted under natural conditions. Compost temperature showed a sharp increase in the first 40 - 60 days, followed by a stabilization at 60°C and a decline after 150 days, whereas compost water content ranged from 50% - 55% to 25% - 30%. Total and selective microbial counts were followed throughout the experiment by means of innovative (IMT) and conventional (CMT) microbiological techniques. *Pseudomonas* spp., anaerobic bacteria, actinomycetes, and fungi reached levels of 8, 7, 5 and 6 log CFU g⁻¹ compost, respectively, with a slight depression after 30-80 days. Total and fecal coliforms strongly decreased during the composting process. The use of IMT allowed to detect a higher and more stable growth of microorganisms if compared to CMT. IMT was demonstrated to be an appropriate and reliable method for monitoring the microbial pools during the co-composting process.

Keywords: Composting; Olive Mill Wastewater; Olive Pomace; Pruning Residues

1. INTRODUCTION

The olive pomace (OP), also called '*sansa vergine*' in Italian or '*orujo*' in Spanish, is defined as the residue that remains after the first oil extraction from olives (crude olive cake). The OP is a dry material with a 8% - 10% moisture and is composed of ground olive stones and pulp, with a high lignin, cellulose, and hemicelluloses content, and a 3% - 5% oil content, depending on the olive mill typology (pressure or centrifugation) [1].

This by-product is usually used for residual oil extraction by solvents, heating, animal feed supplement, or as an organic amendant for olive grove or other crops soils [2]. In terms of agronomic value, OP watered with olive mill wastewater (OMWW), another product of olive milling, or with other organic material lead to a product that supplies nutrients to plants and is an efficient method for the disposal of olive mill residuals [3,4]. According to the Italian law 574/1996, it is possible to use not-composted OMWW and OP for agronomic purposes, as they are considered simple plant amendants, with no limitations on the amounts of OP to be applied to the soil, but the CEE Regulation 91/156 indicates that composting is one of the methods to recycle and recovery organic wastes.

Olive-mill wastewater (OMWW) is composed by the own water of the olives (vegetation water) and the water used in the different stages of oil elaboration [1]. From an environmental point of view, OMWW is an environmental emergency as it has a considerable polluting organic load, with a maximum biological oxygen demand and chemical oxygen demand of about 100 and 220 kg·m⁻³, respectively, an average concentrations of volatile solids and inorganic matter of 15% and 2%, respectively, and organic matter fraction that includes sugars, tannins, polyphenols, polyalcohols, pectins and lipids [5]. Therefore, a series of studies focused on the degradation of OMWW and its chemical components [5-8], and many authors used specific microorganisms for OMWW treatment [9-13]. Microbiological and physicochemical parameters were used as indicators to study the kinetic of OMWW biodegradation, such as chemical oxygen demand (COD), dissolved organic carbon, counts of heterotrophs, filamentous fungi, and yeasts, and content of K, P and N [8-14]. As OMWW does not generally contain sufficient N and P for an adequate aerobic purification process, OMWW degradation may be performed

by co-composting, anaerobic digestion or enzymatic treatment [8-15]. Some authors obtained satisfactory results, in terms of OMWW degradation and amelioration of soil physicochemical properties, by adding this liquid waste to agro-industrial and urban wastes and monitoring the physicochemical parameters during the composting process of these matrices [16-18]. Angelidaki and Ahring [19] studied a combined anaerobic digestion of OMWW together with manure, household waste or sewage sludge, and with this method they managed to degrade OMWW without previous dilution, without addition of external alkalinity and without addition of external nitrogen source. Other authors [3] efficiently monitored over four months a compost made of OP, OMWW and poultry manure, by following temperature, pH, humidity and C/N ratio, in order to ascertain its maturity, and tested its effectiveness in increasing potato agronomic production. Furthermore, the co-composting of exhausted olive cake with poultry manure and sesame shells was recently investigated, and this process was followed by studying some physicochemical parameters [4].

Generally, the study on composting process of olive mill by-products was focused on their physicochemical aspects. On this basis, the present study was performed to evaluate if mixtures of olive mill pomace, olive mill wastewater and olive pruning residues (OPR), without the adding of any other additive external to olive grove, can be efficiently composted under "in farm", non-industrial procedures conditions, based on spontaneous aerobic degradation by autochthonous microorganisms. This methods of compost production needs limited resources, low energetic inputs, and uses machinery and equipment often already present in the farm. As information on selective media for the microorganism responsible for the spontaneous aerobic degradation of compost deriving from different olive material, and in particular from OP, is lacking, we have tested an innovative microbiological technique (IMT) based on microorganisms cultivation using a broth extracted from the matrix to be composted. This method could be used to monitor the biomass degradation process during OP co-composting.

2. MATERIALS AND METHODS

2.1. Compost Production

The composting trials were carried out in a three-year period (2006-2008) under "in farm" and open field environmental conditions located in San Demetrio Corone (Southern Italy, Calabria Region, 39°34' N, 16°21' E). The matrix for compost production included: olive pomace (OP) and olive mill wastewater (OMWW) deriving from a three-phases olive mill, and rain water, added with grinded olive branches and leaves deriving from

pruning residues (OPR). OPR were added in order to give a better structure of the composting matrices.

Depending on the by-products availability in the experimental field, the matrix composition in the three years was, respectively:

year 2006 (from 17 March to 4 October): 200 q OP + 8 m³ OMWW + 20 q OPR;

year 2007 (from 14 March to 28 September): 300 q OP + 80 q OPR;

year 2008 (from 21 March to 30 September): 400 q OP + 5 m³ OMWW + 250 q OPR.

The matrix, uniformly mixed to form a trapezoidal parallelepiped pile (volume = 52, 60 and 129 m³ in 2006, 2007 and 2008, respectively) placed outdoor in open field, resulted to have a semi-solid consistency. It was blended with a mechanical shovel every 7 days, in order to ensure the aeration and control the warmth of the biomass, and was subjected to a spontaneous degradation by autochthonous microorganisms. The aeration allowed to reduce anaerobic fermentation and the consequent high production of putrefactive, toxic compounds for aerobic microorganisms.

Compost maturation was followed by weekly samplings to monitor compost temperature (CT) and compost water content (CWC). The values of CT were measured by a digital thermometer (model 46908; TR snc, Italy). The values of CWC were determined from the weight differences of compost samples before and after drying at 105°C for 24 h, and expressed as percentages of water on dry matter. The mean chemical parameters measured at the end of the composting process in the three years were the following: humidity = 27.5 ± 1.48% (SD), pH = 7.05 ± 0.35, C/N = 24.15 ± 1.06, total organic carbon = 31.4 ± 0.99%, organic matter = 54 ± 1.70%, N = 1.3 ± 0.10%, P₂O₅ = 0.7 ± 0.03%, K₂O = 1.15 ± 0.10%.

Approximately every 25 days, compost samples (5 kg) were randomly collected in different parts and depths of the pile during reversal procedures using sterile gloves, placed in sterile plastic bags, and stored in a refrigerated box at 4°C for 1 h. Before the microbiological analyses, compost samples were sieved at 4 mm and successively at 2 mm, to eliminate the roughest fractions.

2.2. Microbiological Analysis

The screening of microbic pools has been carried out at different steps of composting process. Two methods were utilized: an innovative microbiological technique (IMT) here applied for the first time, and a conventional microbiological technique (CMT) by methods usually adopted in soil analysis [20].

For CMT, a 10 g aliquot of compost was mixed to 90 ml of sterile peptoned water (triptone 10 g·L⁻¹ and sodium chloride 5 g·L⁻¹; pH 7.2) and diluted up to 10⁸ in a

physiological solution. Microbial community has been assessed through plate-counter and expressed as colony-forming units (CFU) per mL. Serial dilutions, using peptoned water as diluent, were prepared to obtain CFU counts in the range of 30 - 300 per plate and 100 μ l of the corresponding decimal dilutions were plated in triplicate the following specific nutritive media: Plate Count Agar (PCA; Oxoid Lim., Hampshire, UK) for total cultural bacteria; Yeast Dextrose Agar (YPD: 1% yeast extract, 2% peptone, 2% glucose) supplemented with 150 ppm chloramphenicol (Sigma-Aldrich, MO, USA) for moulds and yeast; *Pseudomonas* Agar Base medium (Oxoid) with the addition of *Pseudomonas* C-N supplement (Oxoid) for *Pseudomonas* spp.; PCA under anaerobic conditions for anaerobic bacteria; Violet Red Bile Agar (VRBA; Merck and Co. Inc., NJ, USA) for total coliforms; and Slanetz and Bartley medium (Merck) for fecal coliforms.

All plates were incubated at 25°C, as this temperature is adequate for mesophilic microorganisms [20], with the exception of fecal coliforms, that were incubated at 44°C. Microbial counting took place after 24, 36 and 48 h.

For IMT, a 10-g aliquot of compost was mixed to 90 ml of sterilized olive matrix broth (OMB), extracted from the matrix. The OMB was obtained by melting 1 kg of compost matrix in 3 L of sterile de-mineralized water. The 1:3 ratio was chosen on the basis of various trials carried out in our laboratory. The solution was filtered with a cheesecloth, and successively sterilized at 121°C for 20 min, in order to obtain the final OMB. For the microbiological analyses of each compost sample, the OMB used was obtained from that specific sample. The values of pH in the OMBs used ranged from 6.7 to 7.2, with a mean of 7.0 ± 0.2 (SD). For IMT, OMB was utilized as a diluents both for scalar dilutions and substrate preparation. The serial dilutions procedures and the growth media used were the same of CMT. Each measurement was replicated three times and the mean values (\pm SD) were calculated.

2.3. Statistical Analysis

Data were treated by analysis of variance (ANOVA) using the SAS software (SAS Institute, NC, USA) in order to detect significant differences (PROC GLM).

3. RESULTS AND DISCUSSION

A new approach in fruit orchard management is imposed by environmental emergencies, such as soil degradation, water shortage and greenhouse effect [21]. In particular, in olive groves a positive influence of sustainable orchard management, including pruning residues re-use and compost application, on soil biochemical characteristics and soil microbial diversity was recently ob-

served [22,23]. Among the agronomic sustainable practices, the input of soil organic matter as compost is one of the most important factor affecting soil fertility, in terms of enhancement of soil permeability and water retention, better endowment and availability of nutrients for plants, higher CO₂ uptake and carbon fixation, and reduction of soil erosion [23-26].

In all the three years of the experiment, the CT values of the matrices showed a similar trend, with a sharp increase in the first 40 - 60 days followed by a stabilization (**Figure 1**). This period, characterized by high values of CT, corresponds to the so called active composting time (ACT), the phase of bio-oxidation, during which the degradation processes of labile organic components facilitated by pile aeration occurs, with high heat release and oxygen consumption [2,15].

During the ACT, CT remained quite stable for about 40-50 d, reaching maximum values of about 60°C (**Figure 1**). These high temperatures allows compost hygienization during the ACT and thus are an important requisite for the following utilization of the compost. The bio-oxidative phase was considered finished after 140, 160 and 155 days in 2006, 2007, 2008, respectively, when CT started to decrease and reheating did not occur (**Figure**

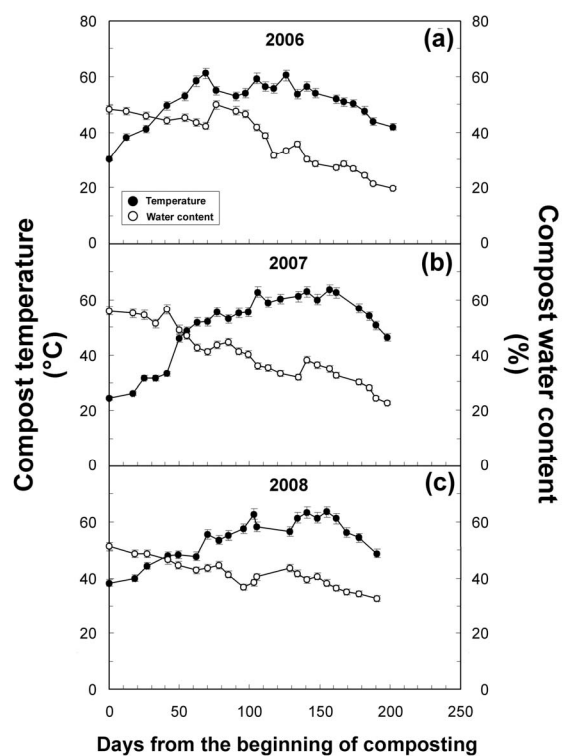


Figure 1. Compost temperature (black circles) and compost water content (white circles) during the year (a) 2006, (b) 2007 and (c) 2008. The values represent the averages (\pm SD) of 10 independent replicates.

re 1). In fact, during the following maturation time (curing), degradation processes and heat production finish, whereas the biosynthesis of humic compounds occurs by fungi [24]. As for CT, the values of CWC showed similar trends in the three years, starting from values of 50-55% and reaching values of about 25% - 30% at the end of curing, with a progressive and gradual decrease throughout the degradation process (Figure 1).

While some authors used OMWW as a substrate for the production of enzymes and organic compounds by microorganisms [27,28], the microbiological application of sterilized compost extracts were never studied. The use of IMT allowed to detect a higher growth of microorganisms if compared to CMT (Figure 2), likely due to the fact that OMB is richer of nutrients if compared to peptoned water, and its compositions is similar to that usually experienced by the microorganisms living in the compost. Another

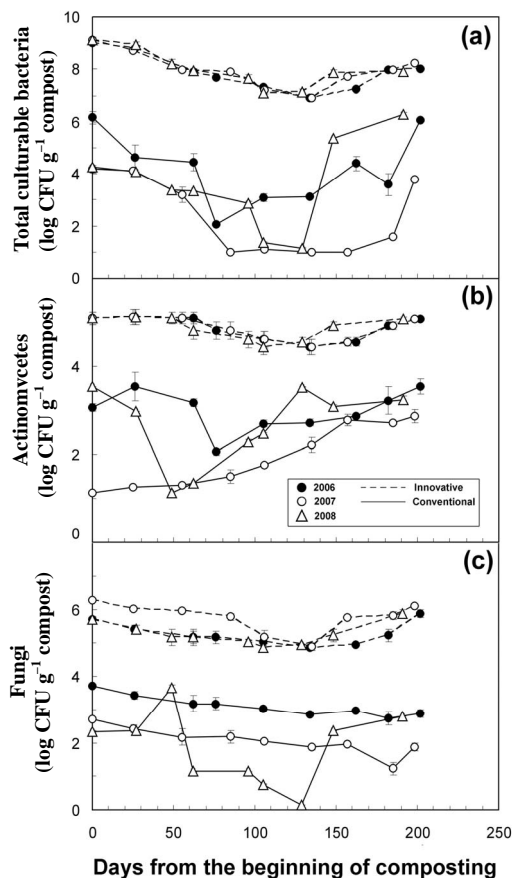


Figure 2. (a) Total cultural bacteria, (b) Actinomycetes, and (c) yeasts and moulds counts (read after a 36-h incubation) using the innovative microbiological technique (IMT; dashed lines) or the conventional microbiological technique (CMT, continuous lines). Experimental period: 2006 (black circles), 2007 (white circles) and 2008 (triangles). The values represent the averages (\pm SD) of three independent replicates. All the IMT and the respective CMT resulted to be significantly different ($P \leq 0.05$).

advantage is that the cultures based on OMB showed more stable and comparable values both throughout the experimental period and between the three years (Figure 2), a basic requisite for their use as bio-indicators of the composting degradation process. The trends of microbial number detected by IMT (Figures 2 and 3), and in particular the kinetics of total and fecal coliforms (Figure 3(c,d)), indicate that this technique could be also adopted for olive grove soils, and urban and industrial sludge, two

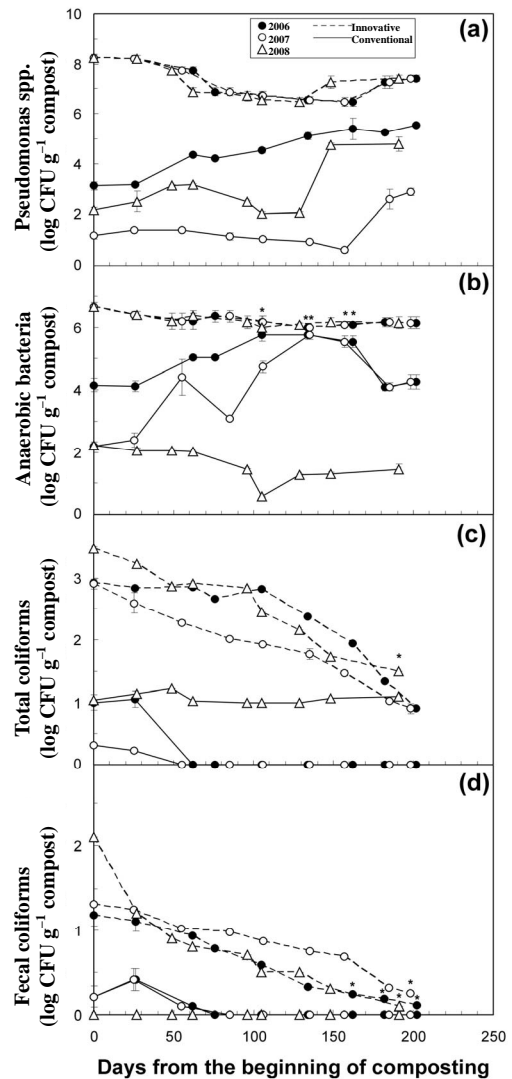


Figure 3. (a) *Pseudomonas* spp., (b) anaerobic bacteria, (c) total coliforms, and (d) fecal coliforms counts (read after a 36-h incubation) using the innovative microbiological technique (IMT; dashed lines) or the conventional microbiological technique (CMT, continuous lines). Experimental period: 2006 (black circles), 2007 (white circles) and 2008 (triangles). The values represent the averages (\pm SD) of three independent replicates. All the IMT values and the respective CMT values resulted to be significantly different ($P \leq 0.05$), with the exception of those with the asterisk on the top.

matrices that host microorganisms living in conditions similar to those studied in this paper [3,8].

Total cultural bacterial counts determined by IMT did not significantly differ during the three years (**Figure 2(a)**), so highlighting that bacterial growth is not inhibited by the different percentages of OMWW, a by-product with a very high concentration of polyphenols [1,4]. Actinomycetes (e.g., *Streptomyces*), fungi produce a number of enzymes that help degrade organic plant material, such as lignin and chitin, and so are abundant in soils rich of organic matter [20]. Furthermore, actinomycetes are able to use root exudates as carbon source, supply roots with easily assimilable nitrates, and are involved in the suppressive action of the soil [23]. Actinomycetes (**Figure 2(b)**), and fungi (**Figure 2(c)**), counted by IMT, reached levels of about 5 and 6 log CFU g⁻¹ compost, respectively.

The use of specific cultural media allowed the isolation of important physiological groups of microorganisms related to compost maturation (**Figure 3**). *Pseudomonas* spp. and anaerobic bacteria (e.g., *Bacillus* spp.) are the main decomposers of complex polymers, such as lignocelluloses and chitin, but they also have proteolytic enzymes and are of key-importance important for the production of assimilable nitrogen in soils [20]. The observed trends of microbial counts using IMT, show that *Pseudomonas* spp. (**Figure 3(a)**) and anaerobic bacteria (**Figure 3(b)**) reached levels of about 8 and 7 log CFU g⁻¹ compost, respectively. The slight decrease of the microbial numbers of these two groups during the first 30-40 days (**Figure 3(a,b)**) was likely due to the increase of temperature [1]. Moreover, this depression started earlier if compared to Actinomycetes, and yeasts and moulds, that declined during the first 70-80 days (**Figure 2(b,c)**). The high number of *Pseudomonas* spp. observed throughout the composting process (**Figure 3(a)**) indicates that these microorganisms are affected by the organic content of the matrix and are able to initiate their oxidative biodegradation without being particularly inhibited by polyphenols. The presence of anaerobic bacteria (**Figure 3b**) indicates that, considering the size of the compost pile and the high microbial activity, with the consequent strong oxygen demand, mechanical aeration did not allow the complete removal of anaerobic degradation processes.

On the other hand, the use of compost in agriculture is often associated with health risks because of the possible presence of human pathogens, enteric in origin, such as bacteria, viruses, protozoa and helminthes [25]. The number of total and fecal coliforms strongly decreased during compost maturation (**Figure 3(c,d)**), so attesting the efficiency of degradation process and the good hygienic conditions of the matrix. In fact, coliform bacteria are of key importance because the low number or the absence

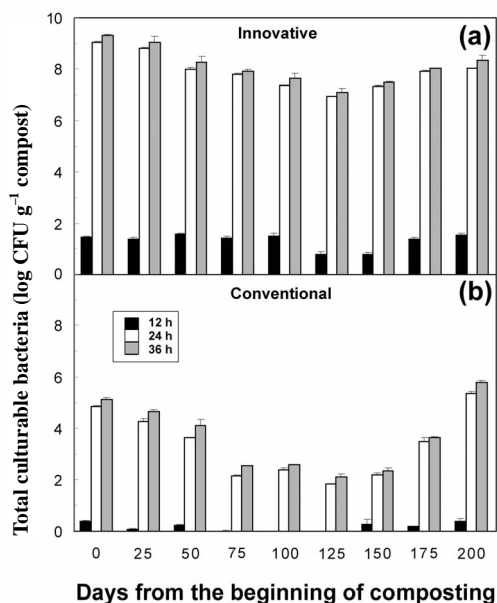


Figure 4. Total cultural bacteria counts read after an incubation time of 12 h (black columns), 24 h (white columns) and 36 h (grey columns) by using (a) the innovative microbiological technique (IMT) or (b) the conventional microbiological technique (CMT). The values represent the three-years averages (\pm SD) of three independent replicates. All the values obtained by 12-h incubation time were significantly different ($P \leq 0.05$) from those of 24-h and 36-h incubation times, whereas no significant differences ($P \leq 0.05$) were found between the values of 24-h and 36-h incubation times.

of this group of micro-organisms indicates compost healthiness and safety [25]. Notwithstanding the widespread use as amendants of olive mill residues and by-product in Mediterranean countries, legislative limits for compost deriving from these matrices are lacking. The only microbiological threshold values for compost application in agriculture, present in the EC Regulation 1774/2002, are 1000 CFU g⁻¹ compost for both *Escherichia coli* and *Enterococcus* spp., and 0 CFU g⁻¹ compost for *Salmonella* spp. Thus, we suggest that the use of total and fecal coliforms could be efficacious bio-indicators, more specific and less variable than the microbial groups listed in EC Regulation 1774/2002.

Our investigation also focused on the more appropriate incubation time before colonies counting. In **Figure 4** we reported the averages total cultural bacterial counts calculated from the values of the three years, determined by IMT (**Figure 4(a)**) and CMT (**Figure 4(b)**). For both CMT and IMT, a counting time of 36 h was the most reliable, as total and specific microbial counts ranged between 4 and 9 log CFU g⁻¹ compost. In fact, the range of microbial number after 24 h was lower (between 0 and 2 log CFU g⁻¹ compost), whereas after 48 h it did not vary significantly from the that obtained after a 36-h incubation.

4. CONCLUSIONS

Our experiment demonstrated that sterilized olive matrix broth can represent an efficacious diluent to be used for monitoring the microbial pools during the co-composting of olive mill by-products. The results also show that the aerobic stabilization of olive pomace, suitably co-composted with olive mill wastewater and/or other crop by-products, is a real sustainable agronomical practice, that allows to obtain a low-cost stabilized organic amendament, in line with the legislative parameters of mixed composted amendaments (Italian law 748/1984). The application of this product could recovery, maintain or increase soil fertility, with environmental benefits and positive influence on crop yields.

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