Rhamnolipid and Surfactin production from olive oil mill waste as sole carbon source

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**Abstract[[1]](#footnote-1)**

Olive mill waste (OMW) creates a major environmental problem due to the difficulty of further waste processing. In this work we present an approach to give OMW added value by using it for the production of biosurfactants. Two bacterial species, *Pseudomonas aeruginosa* and *Bacillus subtilis,* were grown with OMW as the sole carbon source. Glycerol and waste frying oil were used as comparative carbon sources. *B. subtilis* produced surfactin (a lipopeptide) at a maximum concentration of 3.12 mg/L with 2% w/v of OMW in the medium, dropping to 0.57 mg/L with 10% w/v of OMW. In contrast, *P. aeruginosa* produced 8.78 mg/L of rhamnolipid with 2% w/v OMW increasing to 191.46 mg/L with 10% w/v OMW. The use of solvent-extracted OMW reduced the biosurfactant production by 70.8 % and 88.3 % for *B. subtilis* and *P. aeruginosa* respectively. These results confirm that OMW is a potential substrate for biosurfactant production.

**Keywords:** Biosurfactants, Agroindustrial wastes, Alperujo, Surfactin, Rhamnolipids, Olive mill waste

# Introduction

Biosurfactants (BS) are surface active molecules produced mainly through fermentation employing microorganisms. Biosurfactants can be classified according to their chemical structure into glycolipids, lipopeptides, fatty acids, phospholipids, neutral lipids and polymeric BS (Pacwa-Płociniczak et al., 2011). These compounds have the same properties as their synthetic counterparts i.e., emulsification, detergency, wetting, foaming, etc. (Rosen and Kunjappu, 2012). Furthermore, BS are usually more biocompatible and biodegradable than the synthetic ones due to the fact that they are produced by a biological process using renewable resources, avoiding chemical synthesis (Banat et al., 2010). Additionally, they have important surfactant characteristics like stability at extreme pH, salinity and temperature conditions (Lotfabad et al., 2009). For this reason BS are a real alternative to surfactants produced through chemical processes, and are already being used as replacements for them (Marchant and Banat, 2012a), in addition they can be employed in completely new applications like soil remediation, recovery of heavy metals, food or medicine (Banat et al., 2014a; Campos et al., 2013; Diaz et al., 2014; Gudiña et al., 2013).

The main challenge for BS to become a real option in industry are the production costs. Biosynthesis and downstream steps are still expensive, and cheaper and more efficient production methods need to be developed (Marchant and Banat, 2012b). Although fermentation substrates form only a small part of the total production costs some industrial and agroindustrial by-products and wastes could play an important role since their use would provide added value for these wastes that otherwise would be simply discarded, causing a harmful impact on the environment (Henkel et al., 2012; Makkar et al., 2011).

There are a number of different waste products like fatty acids, waste oils or soap stock, which may be used in BS production (Banat et al., 2014b). For example, glycerol is the main co-product from biodiesel production and has low value due to the presence of impurities. De Sousa et al. (2011) used this waste in the production of rhamnolipids with a yield of 1.9 g/L. Soap stock, produced during oil refinery and mainly composed of fatty acids, is another suitable industrial waste for BS production (Helmy et al., 2011). For example, Nitschke et al. (2005) obtained 11.7 g/L of rhamnolipids using soybean soap stock. Vedaraman and Venkatesh (2011) obtained surfactin with a *Bacillus subtilis* strain using waste frying oils (sunflower and rice bran) with a yield up to 650 mg/L of crude extract. Finally other authors proposed the use of residual glycerol and okara (soy pulp) as carbon source in the production of lipopeptides (Slivinski et al., 2012; Sousa et al., 2012). It is necessary to sound a note of caution when considering claims for yields of BS from any oily substrate since the methods of quantification commonly used are extremely unreliable (Marchant and Banat, 2014). Rudden et al.(2015) have recently provided the first fully validated UPLC-MS/MS method for the quantification of the range of rhamnolipid congeners produced by *P. aeruginosa* and have shown that other methods often give significant overestimations of yield.

This paper deals with “alperujo” or olive oil mill waste (OMW), an agroindustrial waste very common in the Mediterranean area, produced after extraction of olive oil. Nowadays, the most frequently used extraction method for olive oil is the two-stage-process, which yields olive oil and the solid waste, “alperujo.” Alperujo is a paste with a high content of lignocellulosic material together with salts and residual oil (Dermeche et al., 2013; Tortosa et al., 2012). Furthermore, this waste is also rich in phenols and polyphenols, substances with phytotoxic and microbial inhibitory properties (McNamara et al., 2008), which can hinder its further processing or recycling. At the present time, alperujo is being re-used in composting, animal feed, or it is simply burned (Tortosa et al., 2012; Vitolo et al., 1999). However, since OMW contains residual oil not extracted in the primary processing, it is worth considering its use in the production of BS, an approach which has hardly been explored, with the exception of Mercadé et al. (1993), which obtained rhamnolipids using the liquid fraction of a three stage olive oil extraction as sole carbon source, and our first work (Maass et al., 2015), where we explored the availability of producing surfactin from OMW.

Therefore, the objective of our work is to explore the suitability of using OMW as a sole carbon source in the production of two biosurfactants: surfactin and rhamnolipid. For this purpose we used batch fermentations of two bacterial species: *Bacilus subtillis* and *Pseudomonas aeruginosa*. We have examined the major parameters that can be optimized to achieve an efficient production process, such as OMW concentration, OMW oil content and kinetics. Two additional carbon sources, i.e. waste cooking oil and glycerol, have been used for comparative purposes.

# Materials and Methods

## Chemicals

Salts for culture media were purchased from BHD (England). Culture medium salts, nutrient agar, protease peptone and glucose were purchased from Oxoid (England). Glycerol, ethyl acetate, methanol, chloroform, hexane and rhamnolipid and surfactin standards for UPLC-MS analysis were purchased from Sigma (Germany). Waste oil (WO) was generously provided by two local restaurants in Granada (Spain). OMW was kindly provided by Cooperativa LA UNIÓN (Montilla, Spain). Its composition is detailed in Table 1, similar to those reported by Niaounakis and Halvadakis (2006)

## Microorganisms

The identities of the microorganisms used were confirmed as *Bacillus subtilis* through sequencing the 16S rRNA gene and *Pseudomonas aeruginosa* PAO1 available at University of Ulster’s laboratory.

## Culture conditions

The bacteria were first inoculated onto nutrient-agar and incubated at 37 oC for 24 h. Two seed cultures were prepared before starting the batch culture. Seed culture 1 was a PPGAS medium with Tris-HCl (19 g/L), protease peptone (10 g/L), glucose (5 g/L), KCl (1.5 g/L), NH4Cl (1 g/L) and MgSO4 (0.4 g/L) in distilled water. Twenty ml of this medium was inoculated with cells from the plate culture and maintained at 37oC for 24 h at 160 rpm. Seed culture 2 was a mineral salt medium (MSM) composed of glucose (20 g/L), NaNO3 (2 g/L), Na2HPO4 (0.9 g/L), KH2PO4 (0.7 g/L), MgSO4· 7H2O (0.4 g/L), CaCl2·2 H2O (0.1 g/L), FeSO4· 7H2O (0.001 g/L) and the following trace elements ZnSO4· 7H2O (0.7 mg/L), CuSO4· 5H2O (0.5 mg/L), MnSO4· H2O (0.5 mg/L), H3BO3 (0.26 mg/L) and Na2MoO4·2 H2O (0.06 mg/L). Seed culture 2 was inoculated with 5 % v/v of seed culture 1, and was grown again at 37oC and 160 rpm for 24 h, for adaptation of cells in MSM.

For the batch fermentation experiments, 100 mL of medium containing the selected concentration of carbon source (2, 5 or 10 % w/v) was added to a 1 L Erlenmeyer flask. The medium was the MSM (see above for seed culture 2) without glucose. Glycerol, WO and/or OMW were also tested as carbon sources. Cultures were inoculated with 5 %v/v of seed culture 2 and maintained at 37 oC and 160 rpm. All experiments were carried out in triplicate.

## Analytical assays

For OMW characterization, evaporation at 105 oC to constant weight was used to determine the moisture content (AOAC, 1999) and the Kjeldahl method (ASTM E258) for total nitrogen. Extraction of the residual oil fraction in OMW was carried out with hexane. The ratio waste/solvent was 1/5 (w/v), and the process was repeated three times. Afterwards oil free OMW was dried, and finally the moisture content was reset again to the initial value.

## Culture media processing and measurements

Cell growth was measured by dry weight (DW) of pellets obtained from 1 mL of culture, centrifuged at 105xg for 15 min and dried at 80oC to constant weight (Maass et al., 2015) . This measurements were accomplished by triplicate for each sample. But obtained results were not accurate enough, due to the solid fraction of OMW, and therefore they were only used as illustrative results.

The culture medium was centrifuged at 105xg and 4oC for 15 min and the supernatant was carefully collected. Surface tension (ST) of the clarified medium was measured using the Wilhelmy method in a Kruss K 10 ST tensiometer (Hamburg, Germany). Afterwards BS (rhamnolipids or surfactins) were extracted. For this purpose 50 mL of supernatant was gently mixed with the same volume of ethyl acetate. Organic and aqueous phases were separated in a funnel. These two last steps were repeated three times. The three organic fractions obtained were combined and rotary evaporated. The crude extract was dissolved in a small amount of methanol in order to be transferred to a pre-weighted vial, then dried and weighed to give the yield of crude extract.

## Surfactant quantification

The BS produced was identified and quantified by UPLC-MS. The measurements were conducted in an UPLC Waters Acquity H-Class chromatograph (Waters Corporation, Milford-MA, USA) equipped with a Waters UPLC BEH C-18 column and coupled to a mass spectrometer (Waters Xevo-TG-S). For surfactin analysis the mobile phase consisted of a mixture of 20 %(v/v) of water and 80%(v/v) of acetonitrile, both with a 0.1% w/w of formic acid. The volumetric flow rate of the mobile phase and the sample injection volume were 0.5 mL/min and 3.0 µL, respectively. Mass spectrometry was conducted in positive mode (ES+), under the following conditions: capillary voltage 3 kV, ion source temperature 150oC, desolvation temperature 500oC, and desolvation gas flow rate 800 L/h. For rhamnolipids analysis the mobile phases was 70 %(v/v) water and 30 % (v/v) acetonitrile and the mass spectrometry was carried out in negative mode (ES-), all the other conditions were as described above.

# Results and discussion

## The effect of Carbon source

Initially, the effect of the carbon source on the cell dry weight (DW), surface tension (ST) of the supernatant and BS concentration after 7 days of culture was determined.

Both the type of carbon source and its concentration affected the DW and ST of the cultures. With the exception of 2 % w/v *P. aeruginosa* glycerol yielded the highest values of DW (Figure 1). The DW values for waste oil and OMW were lower, and similar for these two carbon sources. Higher concentrations of carbon source yielded higher values of DW in all cases, although the increase with concentration was higher for the glycerol cultures. However, in the case of *B. subtilis* the DW increased with the concentration of glycerol and OMW, while waste oil as carbon source yielded less biomass (Figure 1). Remarkably, the values of DW obtained for glycerol and OMW when using *Bacillus* were very similar at concentrations of 5 and 10 % w/v.

In the case of ST, the lowest values were achieved in the cultures with glycerol for both microorganisms, slightly decreasing with increasing substrate concentration for *P. aeruginosa* and the reverse for *B. subtilis*, and ranging between 29.2 and 33.4 mN/m. The same effect was observed for WO as carbon source. Of special interest is that for OMW, ST decreased with concentration from 45.1 to 33.7 mN/m in *Pseudomonas* cultures, whereas an increase from 39.3 to 46.6 mN/m was obtained with *Bacillus*.

These results suggest that both microorganisms are able to grow in OMW and produce molecules with interfacial properties. *P. aeruginosa* seems better able to grow and produce surfactant with OMW in the medium, especially at higher OMW concentrations. The increase in final ST with OMW concentration in *B. subtilis* cultures suggests that this waste inhibits the ability of this microorganism to produce substances with interfacial activity (Figure 1). This effect could be due to the antimicrobial activity of phenols found in OMW, which is well known and described in the literature (Obied et al., 2005; Perez et al., 1992). For instance, Moreno et al. (1990) reported that this antimicrobial effect is higher for Gram-positive than for Gram-negative bacteria. Therefore, Gram-negative species, such as *P. aeruginosa,* could be more suitable to grow on OMW as carbon source.

UPLC-MS analysis of the extracts from the culture media confirmed that *P. aeruginosa* had produced rhamnolipids and *B. subtilis* surfactins. This result supports the contention that it is feasible to produce BS from OMW.

Furthermore, the direct quantification of the produced BS confirmed that the highest BS amount was produced with glycerol for both species (see Table 2), while WO and OMW yielded less BS. With regard to the effect of the carbon source concentration, the amount of rhamnolipid produced remained relatively constant with increasing glycerol concentration, while surfactin decreased almost to a half as the glycerol concentration was increased from 2% to 10 % w/v. Additionally, both strains yielded less BS with increasing WO concentration, with the reduction in surfactin more marked. Under the best conditions using OMW in these experiments 191.46 mg/L of rhamnolipids and 3.12 mg/L of surfactin were obtained, these conditions corresponded to 10 % and 2 % w/v of OMW for *P. aeruginosa* and *B. subtilis* respectively. Again it is interesting to note the effect of OMW concentration in both strains: while *P. aeruginosa* grew better with OMW as carbon source, increasing its productivity at higher concentration, *B. subtilis* had reduced production of surfactin as the concentration of OMW was increased from 2% to 10 % w/v. Furthermore, in terms of efficiency, rhamnolipid YP/S was considerably higher than surfactin at their respective optimum condition for OMW as carbon source: 19.15 mg rhamnolipids/g OMW at 10 % w/v OMW versus 1.56 mg surfactin/g OMW at 2% w/v OMW.

The higher BS yields obtained with OMW for rhamnolipid and surfactin were respectively 12 % and 4 % of the obtained yield with glycerol at 2% w/v. In spite of this, OMW would be a suitable raw material in the BS production process due to its low cost. Although carbon substrate cost is not a critical element in the overall production costs of BS the opportunity to add considerable value to the OMW is potentially important. The OMW used in these experiments contained little residual oil, however, if OMW was used prior to any secondary extraction of oil using solvents the levels of BS could be increased thereby improving the economics of downstream processing. An additional advantage of using OMW is the purity of the extract, which has intermediate values between glycerol and WO. Actually, the experiments carried out with WO presented the lowest purity; particularly for *B. subtilis* cultures (see Table 3). In these cultures unconsumed oil is quite difficult to separate from the rest of the medium, while when using OMW, this problem does not exist, because solid particles are easily separated. This is a clear advantage of OMW over WO for industrial BS production, because it will simplify downstream processing, which is a major factor to take into account, in addition to the price of raw materials (Smyth et al., 2010).

## Kinetics

Once it had been demonstrated that it is possible to produce BS with OMW as sole carbon source the next steps were oriented towards process optimisation. With this aim, a kinetic study was carried out to examine the course of BS production during the culture using this waste as sole carbon source. DW and BS concentration were measured at different times. Both parameters increased rapidly in the exponential growth phase, and remained almost constant in the log phase for both microorganisms. Figure 2 shows how biomass concentration in the culture medium reaches a maximum after 6 days for *P. aeruginosa* and 4 days for *B. subtilis*. In the case of BS maximum concentrations were reached after 8 days for *P. aeruginosa* and 6 days for *B. subtilis*. The production of surfactin by *B. subtilis* appears to be closely associated with the growth of the organism since surfactin is detected in the medium very early. In contrast rhamnolipid production by *P. aeruginosa* follows the pattern described by other workers where production takes place as the cells enter stationary phase (Perfumo et al., 2013).

In order to gain knowledge of the BS produced with OMW, we studied the different congeners produced by using UPLC/MS. Table 4 presents the different congeners detected and their average relative abundance, which remained almost constant during the whole kinetic study for both microorganisms. For *P. aeruginosa* the most abundant congener was the dirhamnolipid with 10 carbon in each fatty acid chain (2 Rha-C10-C10). Whereas *B. subtilis* mainly produced surfactins with 14 and 15 carbon atoms in the β-hydroxy-fatty acid tail.

## Combined carbon source study; OMW toxicity.

With the purpose of achieving a better understanding of the problems produced by using OMW, we combined it with glycerol as an additional carbon source. Glycerol concentration was kept constant at 2% w/v, an amount that was completely consumed after the 7 days of culture (data not shown). OMW was added at 2% and 10 % w/v, as shown in Figure 3.

For both microorganisms combined carbon sources yielded less BS than experiments carried out only with glycerol, and more than when only OMW was used (Figure 3). Compared with experiments when only glycerol was used, rhamnolipid productivity decreased by 47.6 % and 75.9 % in cultures with 2% and 10 % w/v of OMW respectively. For surfactin the reduction was of 5.6 % and a 92.2 % for the same conditions. This indicates that some component(s) in OMW have an inhibitory effect on the production of BS. Previously we noted the inhibition of surfactin production by *B. subtilis* at high concentrations of OMW. Additionally these results show how this waste could have hindered the ability of *P. aeruginosa* to produce rhamnolipids, although this effect was not fully obvious when OMW was the sole carbon source. As discussed previously, this reduction could be due to the presence of phenols in OMW. Therefore in order to develop an optimal BS production process the use of more phenol-resistant strains or phenol removal should be contemplated.

## Relevance of residual oil in OMW

Our first hypothesis when using the waste from olive oil extraction was that bacteria could use the residual oil in this waste (1.4 % w/w of the wet product) to produce BS, as reported in many studies where waste oil is used as carbon source. With the purpose of confirming if bacteria could process this residual oil, we carried out fermentations with oil-free OMW (OF-OMW).

Figure 4 shows how for both bacteria the biomass and the BS production were considerably lower with oil-free OMW, which yielded a 88.5 % and 70.7 % less BS, compared to fresh OMW, for *P. aeruginosa* and *B. subtilis* respectively. This confirms the importance of residual oil in the BS production, despite its low concentration in OMW.

Surprisingly, both strains still produced some BS with oil-free OMW without oil. This suggests that there is something else besides oil that bacteria are able to use as carbon source. Probably, hemicellulosic material, and more specifically, free sugars could be an available and easy carbon source to incorporate (Jain et al., 2013; Makkar et al., 2011). However this hypothesis needs further investigation.

# Conclusions

In this study we demonstrate that olive oil mill waste (OMW) is a potential carbon source for biosurfactant production. OMW is somewhat inhibitory to BS production but can be used at appropriate levels. The residual oil in OMW provides a major source of carbon but other nutrients in the waste also contribute. OMW is an abundant by-product in Mediterranean countries which could be used for biosurfactant production achieving two important objectives: (i) recycling of a waste that is difficult to process, and (ii) production of high added value products. However, questions remain before an economically viable process can be developed.

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# Figure captions

Figure 1. Influence of carbon source (glycerol, WO and OMW) and its concentration (2, 5 and 10% w/v) on cell dry weight (coloured columns-left) and ST (uncoloured columns-right) in culture supernatant after 7 days of culture. (a) *P*. *aeruginosa* (b) *B*. *subtilis*. Error bars correspond to the standard deviation of three replicates.

Figure 2. Time course of DW (▲) and BS (●) production for *P. aeruginosa* (continuous line) and *B. subtilis* (broken line) cultures. OMW was the sole carbon source at concentrations of 10 % w/v for *Pseudomonas* and 2 % w/v for *Bacillus*.

Figure 3. Production of rhamnolipids and surfactin by *P. aeruginosa* and *B. subtilis* respectively using glycerol, OMW and glycerol mixed with OMW, after 7 days of culture.

Figure 4. Effect of oil extraction of OMW on the biomass (DW, coloured columns) and biosurfactant production (BS, uncoloured columns). OMW concentration of fresh and oil-free OMW were fixed as 10 % for *P. aeruginosa* cultures and 2% for *B. subtilis.* The duration of cultures was of 7 days.

# Figures and Tables

Figure 1.



Figure 2.



Figure 3.



Figure 4.



Table 1. Oil mill waste composition

|  |  |  |
| --- | --- | --- |
| Water content (%) | | 64.37 |
| Lipid content (%) | | 3.89 |
| Protein content (%) | | 7.10 |
| Free sugars (%) | | 9.50 |
| Elemental analysis (%) | C | 48.15 |
| N | 1.18 |
| H | 7.14 |

Percentages referred to dry OMW

Table 2. Influence of carbon source (glycerol, waste oil and oil mill waste) and its concentration (2, 5 and 10 % w/v) on rhamnolipids (Rha, mg/L) and surfactin (Surf, mg/L) concentration in the culture medium for *P. aeruginosa* and *B. subtilis* cultures.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | **Glycerol** | | **WO** | | **OMW** | |
|  | Rha (mg/L) | Surf. (mg/L) | Rha (mg/L) | Surf. (mg/L) | Rha (mg/L) | Surf. (mg/L) |
| **2 %** | 1564.70 ± 106.70 | 21.72 ± 2.61 | 899.94 ± 74.04 | 1.01 ± 0.15 | 8.78 ± 0.49 | 3.12 ± 0.13 |
| **5 %** | 1295.05 ± 82.62 | 20.76 ± 2.49 | 496.23 ± 139.23 | 0.08 ± 0.02 | 27.07 ± 1.71 | 0.93 ± 0.21 |
| **10 %** | 1906.83 ± 121.66 | 13.51 ± 1.63 | 315.59 ± 8.87 | n.d. | 191.46 ± 18.76 | 0.57 ± 0.10 |

n.d. = not detected

Table 3. Purity (in %) of rhamnolipids (Rha) and surfactin (Surf) in crude extracts from culture medium, at three different concentration of glycerol, waste oil (WO) and olive oil mill waste (OMW).

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | Glycerol | | WO | | OMW | |
| Concentration | Rha | Surf | Rha | Surf | Rha | Surf |
| 2 % | 69.09 | 6.38 | 20.23 | 0.05 | 6.62 | 1.24 |
| 5 % | 57.97 | 1.06 | 10.52 | 0.01 | 6.86 | 0.13 |
| 10 % | 62.93 | 0.52 | 5.11 | n.d. | 20.43 | 0.05 |

n.d. = not detected

Table 4. Rhamnolipid and surfactin relative abundance of different congeners during kinetic study with 10% and 2% w/v of OMW as carbon source respectively. The relative abundance remained almost constant throughout each experiment.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **Rhamnolipid congeners** | | | | |
| Congener | Rha-C8-C10 | Rha-C10-C10 | | Rha-C10-C12:1 | 2 Rha-C8-C10 |
| M.W. ion-H (m/z) | 475 | 503 | | 529 | 621 |
| Relative abundance (%) | 4.98 | 27.49 | | 0.61 | 4.10 |
| Congener | 2 Rha-C8-C12:1 | 2 Rha-C10-C10 | | 2 Rha-C10-C12:1 | 2 Rha-C10-C12 |
| M.W. ion-H (m/z) | 647 | 649 | | 675 | 677 |
| Relative abundance (%) | 0.13 | 58.12 | | 3.72 | 0.85 |
|  |  |  |  |  |  |
|  | **Surfactin congeners** | | | | |
| Congener | C12 | C13 | C14 | C15 | C16 |
| M.W. ion+H (m/z) | 994.6 | 1008.6 | 1022.6 | 1036.6 | 1050.6 |
| Relative abundance (%) | 1.01 | 9.07 | 33.55 | 46.06 | 10.31 |

Data are average of duplicate determinations

1. Abbreviations: BS (biosurfactant); WO (waste cooking oil); OMW (Olive oil mill waste); OF-OMW( Oil-free olive oil mill waste); DW (dry weight); ST (surface tension) [↑](#footnote-ref-1)