BIODEGRADATION OF OLIVE MILL WASTEWATER
BY TRICHOSPORON CUTANEUM AND GEOTRICHUM CANDIDUM

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Olive oil production generates large volumes of wastewater. These wastewaters are characterised by high chemical oxygen demand (COD), high content of microbial growth-inhibiting compounds such as phenolic compounds and tannins, and dark colour. The aim of this study was to investigate biodegradation of olive mill wastewater (OMW) by yeasts Trichosporon cutaneum and Geotrichum candidum. The yeast Trichosporon cutaneum was used because it has a high potential to biodegrade phenolic compounds and a wide range of toxic compounds. The yeast Geotrichum candidum was used to see how successful it is in biodegrading compounds that give the dark colour to the wastewater. Under aerobic conditions, Trichosporon cutaneum removed 88 % of COD and 64 % of phenolic compounds, while the dark colour remained. Geotrichum candidum grown in static conditions reduced COD and colour further by 77 % and 47 %, respectively. This investigation has shown that Trichosporon cutaneum under aerobic conditions and Geotrichum candidum under facultative anaerobic conditions could be used successfully in a two-step biodegradation process. Further investigation of OMW treatment by selected yeasts should contribute to better understanding of biodegradation and decolourisation and should include ecotoxicological evaluation of the treated OMW.

KEY WORDS: decolourisation, detoxification, phenolic compounds, yeasts
compounds, free fatty acids, low pH, low dissolved oxygen (DO), high total suspended solids, and tannins (7-9). Some claim that OMW toxicity does not depend on extraction procedure (8), while others report that OMW from the continuous process is less toxic that from the traditional (10). The concentration of phenolic compounds in OMW may reach 5 g L\(^{-1}\) to 10 g L\(^{-1}\) due to higher solubility in the water phase than oil (10-12). The types and concentrations of phenolic compounds in OMW vary tremendously with region, type of used process, conditions of everyday use, and local operational procedure (13). More than 30 different phenolic compounds have been identified in OMW: monocyclic aromatic molecules such as hydroxytyrosol, tyrosol, catechol, methylcatechol, caffeic acid, and compounds with greater molecule mass obtained by their polymerisation (12, 14, 15).

OMW is an important environmental problem in the Mediterranean countries where it is generated in huge quantities over short periods of time. Discharge of OMW directly into soil may affect its physical and chemical properties such as porosity and pH, but the main problem, due to the direct use of OMW for irrigation, is the presence of phenolic compounds which are phytotoxic and can inhibit seed germination (16).

In the past two decades, many processes have been investigated in order to reduce the toxicity of OMW and to use it as food or raw material in various biotechnological processes (1, 13, 17, 18). Treatment and reuse of OMW presents significant challenges both due to the nature of olive oil production and due to the characteristics of the wastewater (high COD, high phenolic content, and dark colour) (13). A number of different microorganisms and processes have been tested to treat OMW (13). Biological processes investigated for OMW treatment are aerobic and anaerobic (6-11). The major problem with anaerobic processes is the inhibition of anaerobic microorganisms by phenolic compounds and long-chain fatty acids present in OMW and their persistence in effluent (13). A build up of recalcitrant phenolics such as condensed tannins is also reported (19). Aerobic treatment with activated sludge is only efficient for diluted wastewater, because phenolic compounds present in the wastewater inhibit the activity of the activated sludge microorganisms (20).

Literature reports about degradation of phenol (21-24) and phenolic compounds extracted from OMW (24) by the yeast *Trichosporon cutaneum*. Other yeasts of the genus *Geotrichum* (12, 13, 15, 25-27), *Candida* (7, 10, 28) and *Yarrowia* (28), moulds (13, 25) and white-rot fungi (13, 25) have also been investigated in studies of OMW biodegradation.

Phenol removal is an important step in OMW biodegradation and environmental protection. The other significant component of environmental pollution is the dark colour that is to be removed during OMW treatment before discharge.

The purpose of this study was to investigate OMW biodegradation by *Trichosporon cutaneum* and *Geotrichum candidum*. *T. cutaneum* was used because of its potential to degrade phenolic compounds in wastewaters of different origin (21). *G. candidum* was isolated from the investigated OMW and used in decolourisation experiments, because of its known enzymatic potential for OMW decolourisation (27). The main goal was to remove phenol compounds by *T. cutaneum* to reduce the OMW toxicity and the second goal was to remove the dark colour in so treated OMW using *G. candidum*.

**MATERIALS AND METHODS**

**Wastewater**

Olive mill wastewater used in this study was obtained from a local olive oil manufacturer (Croatia). OMW was centrifuged at 5000 x g for 10 min (5 °C) (Beckman J-21B, USA) to remove suspended solids, and stored at -20 °C. The main chemical and physical characteristics of raw OMW, expressed as mean values ± standard deviation (n=3), were: pH (5.5±0.1), COD (54±0.6) g L\(^{-1}\), total suspended solids (42±1) g L\(^{-1}\), total phenols (9.2±0.2) g L\(^{-1}\), and total nitrogen (0.9±0.1) g L\(^{-1}\). pH was measured using a pH meter (WTW 320, Germany). COD is used as a measure of the oxygen equivalent of the organic matter content of a sample that is susceptible to oxidation by a strong chemical oxidant. It was determined using the APHA standard closed reflux method (5520 C) (29). Total suspended solids (TSS) were determined by filtering known volume of sample through a previously weighed and dried filter paper (overnight), then dried overnight to the constant weight (at 103 °C to 105 °C) according to the APHA standard method 2540 D (29), and expressed in g L\(^{-1}\). Total phenols were determined using the Folin-Ciocalteu’s phenol reagent (Fluka, Switzerland) (30) and total nitrogen (N-tot) was analysed photometrically (Merck)
Spectroquant® Vega 400) with standard kits (Merck-Spectroquant, Germany). Prior to biological treatment, pH was adjusted to 6 with 2 mol L\(^{-1}\) NaOH (Gram-mol, Croatia). For aerobic experiments, OMW was diluted with tap water to the initial value of the COD and phenols of (11.8±0.3) g L\(^{-1}\) and (2±0.13) g L\(^{-1}\), respectively. Aerobically treated OMW, with addition of glucose (Gram-mol, Croatia) as carbon source (1 g L\(^{-1}\)) was used for further decolourisation with \(G.\) candidum. UV-spectrophotometric measurements of colour intensity were performed with a UV-VIS spectrophotometer (Pye Unicam, Helios beta, UK) at 390 nm.

**Yeasts**

Yeasts \(T.\) cutaneum and \(G.\) candidum used in this research belong to the collection of microorganisms of the Laboratory for Biological Wastewater Treatment, Faculty of Food Technology and Biotechnology, University of Zagreb. \(T.\) cutaneum was isolated from activated sludge used to treat oil refinery wastewater (21) while \(G.\) candidum was isolated from the investigated OMW.

The strains were identified using the API Candida System (bioMérieux, France), maintained on a yeast growth medium (YGM) solidified with agar (14 g L\(^{-1}\), Biolife, Italy) and on diluted OMW (40 %, v/v) solidified with agar (14 g L\(^{-1}\), Biolife, Italy). They were replicated every month. The composition of YGM was: glucose (20 g L\(^{-1}\), Gram-mol, Croatia), bacto peptone (10 g L\(^{-1}\), Difco, USA), and yeast extract (5 g L\(^{-1}\), Biolife, Italy).

For experiments with phenol removal, the biomass of \(T.\) cutaneum grown on YGM agar plates (at 28 °C for three days) was transferred in 500 mL Erlenmeyer flasks with 100 mL of YGM and with diluted and sterilised (at 121 °C for 30 min) OMW in a volume portion of 30 % to 50 %, and incubated on a rotary shaker (Certomat IS, Sartorius, Germany) at 150 rpm and 28 °C. After three days of cultivation, the biomass was centrifuged at 5000 x g and 5 °C for 10 min, washed with sterile saline solution, and the suspension of so prepared biomass used as inoculum in the OMW biodegradation experiment.

For the OMW decolourisation experiment, \(G.\) candidum grown on YGM agar plates (at 30 °C for three days) was transferred in 500 mL Erlenmeyer flasks containing 100 mL YGM with aerobically treated OMW (in volume portion of 50 % to 60 %) and incubated at 30 °C. After three days of cultivation, the biomass was centrifuged at 5000 x g and at 5 °C for 10 min, washed with sterile saline solution and used as inoculum.

All cultures were grown in triplicates.

**Experimental methods**

The experiments were carried out in 2 L glass serum bottles with working volume of 1.5 L and at ambient temperature. Serum bottles were equipped with an aeration system that included an air pump (SPP-40GJ-L, Techno Takatsuki Co., LTD, Japan), a silicone tube (Tecno Plast, Germany), a sterile filter unit (pore size 0,45 μm, Wathman, Germany), and a sterile serological pipette (10 mL, TPP, EU). Constant air flow ensured sufficient oxygenation of the mixture by bubbling.

OMW was biodegraded by \(T.\) cutaneum under aerobic conditions (dissolved oxygen 2.5 mg L\(^{-1}\) to 3.5 mg L\(^{-1}\)) and at pH 6.0 to 6.5. The dark colour of so treated OMW persisted. At the end of the experiment with \(T.\) cutaneum aeration was stopped, the biomass separated by centrifugation at 5000 x g and 5 °C for 10 min, and the supernatant was then treated with \(G.\) candidum.

Decolourisation experiment with \(G.\) candidum was performed under static conditions (dissolved oxygen 0.5 mg L\(^{-1}\) to 0.8 mg L\(^{-1}\)) because static conditions improve decolourisation (15).

All experiments were performed in triplicate. Control experiments were done without inoculation with biomass.

**Statistical analysis**

Results presented are mean values ± standard deviation. Statistical analysis was performed using Student’s \(t\)-test and Pearson’s correlation coefficient to test the differences between variables. Statistical significance was set at \(P<0.05\).

**RESULTS AND DISCUSSION**

Efficient treatment of olive oil industry wastewaters is a difficult task, as they contain many phenolic compounds and other toxic and biomass growth inhibiting compounds (11).

OMW is recalcitrant to conventional wastewater treatment, and its management and disposal present a serious environmental problem (8). The removal of phenolic compounds is therefore the first step in OMW biodegradation and detoxification (24). For
that purpose, many types of yeast were successfully used (7, 10, 13, 28). In this study we used *T. cutaneum* because it has the greatest phenolic biodegradation potential for wastewaters of different origin among the yeasts of the *Trichosporon* genus (21-24).

*T. cutaneum* used in this study was isolated from activated sludge originating from a system for biological treatment of oil refinery wastewater (21). It has been reported for degradation of phenol and other toxic compounds simultaneously as model compounds or present in wastewater (21-23) and phenolic fraction from OMW (24).

In our research the ability of *T. cutaneum* for phenolic compounds removal, and the effect of OMW composition on their growth and activity for OMW degradation was investigated in real OMW. Because of inhibitory effects of phenolic compounds on microorganisms in this research was used OMW diluted with tap water (11, 15, 26, 27).

Figure 1 shows removal of total phenols from OMW with *T. cutaneum*. Degradation of phenolic components was accompanied by biomass growth, COD reduction and pH decrease. In the first 18 h, pH decreased to 5.85 and then increased to 6.16 by the end of the experiment. There was no colour reduction. Under aerobic conditions, the reduction of COD and phenols over 54 h was 88 % and 64 %, respectively. Also, the increase in biomass concentration (Figure 1) confirmed that *T. cutaneum* utilised OMW compounds for its growth. In control experiments, there were no significant changes in COD (Figure 1).

A previous study (21) reported that *T. cutaneum* decomposed 1.5 g L\(^{-1}\) of phenol, as a sole source of carbon and energy over 50 h. Similarly, 2 g L\(^{-1}\) of phenol was degraded after 48 h (24) and with lag phase (18 h). However, Chtourou et al. (24) observed that phenol concentrations above 2.5 g L\(^{-1}\) were inhibitory. In our investigation biodegradation of (1.96±0.03) g L\(^{-1}\) total phenols in OMW was achieved in 54 h without the lag phase (Figure 1). This result confirms high enzymatic activity of *T. cutaneum* for phenol reduction.

The literature data (24) reported that *T. cutaneum* was able to grow on phenolic compounds extracted in ethyl acetate from OMW. The removal of phenols was 90.6 % (from initial 2 g L\(^{-1}\)) as well as COD removal (from initial 19 g L\(^{-1}\)) during 8 days. By HPLC analysis it was confirmed the degradation and transformation of phenolic compounds and formation of new compounds that persisted degradation.

However, this first-pass removal of phenolic compounds and COD reduction under aerobic conditions did not remove the OMW’s dark colour. This colour is owed to polyphenols, phenolic compounds with high molecular weights that are not easily biodegraded (1, 26, 27). This is why we used *G. candidum* in the second pass, as it is known to depolymerise polyphenols and produce enzymes that decolour OMW (11-13, 25-27). The results of decolourisation of aerobically treated OMW with *G. candidum* are shown in Figure 2. Over the six days of the experiment, the treatment with *G. candidum*
removal of COD and reduction in color by 47%. The biomass concentration increased insignificantly. A decrease in pH (data not presented) suggests that the remaining phenolic compounds were biodegraded to products which lower pH. Similar was reported elsewhere for *G. candidum* (11, 26, 27). All changes observed during aerobic treatment of OMW under facultative anaerobic conditions indicate high enzymatic activity of *G. candidum* in biodegradation of less readily biodegradable compounds, especially those that give color. A number of studies investigated different conditions of *G. candidum* growth and OMW decolourisation (11, 15). Some have pointed out the importance of the addition of an easily biodegradable carbon source for lignin peroxidase production and the biomass growth (11, 12, 27). In our decolourisation experiments with *G. candidum* glucose was used as carbon source. It was also noted that *G. candidum* possessed the ability to decolourise fresh OMW, but when it was incubated with stored black OMW as the sole carbon source, there was no decolourisation. Therefore, glucose seems to play an important role in the *Geotrichum* decolourisation process (26).

CONCLUSIONS

Because of OMW’s composition and environmental issues involved, it is necessary to control its disposal and develop new technologies to reduce pollution. However, this may present a considerable technical and economic challenge, not only because of the complex composition of OMW, but also because of the seasonal nature of olive oil production and wide geographical dispersion of mills. Effective biotreatment of OMW that significantly reduces COD, phenolic compounds, and color allows for safe and economical disposal of OMW into soil or into surface waters. Further investigation of OMW treatment by selected yeasts should contribute to a better understanding of the processes of biodegradation and decolourisation (production of enzymes, definition of process factors) and should include ecotoxicological evaluation of the treated OMW.

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REFERENCES


**Sažetak**

BIORAZGRADNJA OTPADNE VODE IZ PROCESA PROIZVODNJE MASLINJAVA ULJA S POMOĆU *TRICHOSPORON CUTANEUM* I *GEOTRICHUM CANDIDUM*

Tijekom proizvodnje maslinova ulja nastaju velike količine otpadne vode. Karakterizira je visoka kemijska potrošnja kisika (KPK), visoka koncentracija sastojaka koji inhibiraju rast mikroorganizama, poput fenolnih spojeva i tanina i tamna boja. Cilj ovog rada je bio istražiti biorazgradnju otpadne vode iz procesa proizvodnje maslinova ulja s pomoću kvasaca *Trichosporon cutaneum* i *Geotrichum candidum*. Kvasac *Trichosporon cutaneum* u ovom je istraživanju uporabljen zbog svog visokog potencijala za biorazgradnju fenolnih i drugih toksičnih spojeva. Kvasac *Geotrichum candidum* uporabljen je kako bi se istražilo koliko je uspješan u biorazgradnji spojeva koji otpadnoj vodi daju tamnu boju. Pri aerobnim uvjetima *Trichosporon cutaneum* uklonio je 88 % KPK i 64 % fenolnih spojeva, dok tamna boja otpadne vode preostaje. *Geotrichum candidum* u statičkim je uvjetima smanjio KPK i boju za 77 %, odnosno 47 %. Ovo je istraživanje pokazalo da *Trichosporon cutaneum* pri aerobnim uvjetima i *Geotrichum candidum* pri fakultativno anaerobnim uvjetima mogu biti uspješno uporabljeni u dvostupanjskom procesu biorazgradnje. Daljnje istraživanje obrade otpadne vode iz procesa proizvodnje maslinova ulja s pomoću odabranih kvasaca trebalo bi pridonijeti boljem razumijevanju biorazgradnje i uklanjanju boje i trebalo bi biti provedeno zajedno sa ekotoksikološkim vrednovanjem obradene otpadne vode.

**KLJUČNE RIJEČI:** detoksifikacija, fenolni spojevi, kvasci, obezbojenje

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