PRODUCTION OF PLANT PROTECTION AGENTS IN MEDIUM CONTAINING WASTE GLYCEROL BY STREPTOMYCES HYGROSCOPICUS: BIOPROCESS ANALYSIS

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The surplus of waste glycerol, by-product of the biodiesel production process, is available at the global market. Some species of the genera Streptomyces have the ability to assimilate glycerol and convert it into valuable metabolic products. In the present study, the ability of Streptomyces hygroscopicus to assimilate waste glycerol and convert it into metabolic compounds with antifungal activity against four phytopathogenic fungi obtained from apple fruit samples expressing rot symptoms, was investigated. Production of antifungal metabolites by S. hygroscopicus was carried out in 3 l stirred tank bioreactor through 7 days. Fermentation was carried out at 27 °C with aeration rate of 1.5 vvm and agitation rate of 100 r.p.m. The aim of this work was to analyse bioprocess parameters and to determine at which stage of bioprocess the production of antifungal metabolites occurs. Activity of the cultivation liquid on two isolates of Alternaria alternata and two isolates of Fusarium avenaceum were determined every 12 h using in vitro well diffusion method. It was found that the maximum production of antifungal metabolites occurred at 108 hour of cultivation. Formed inhibition zones have shown that the produced antifungal metabolites have high efficacy on tested phytopathogenic fungi (inhibition zone diameter higher than 35 mm for all test organisms).

Keywords: Streptomyces hygroscopicus, bioprocess, waste glycerol, postharvest apple disease, biocontrol

Biodiesel is an alternative to fossil fuels. However, during production of biodiesel and its by-products, waste glycerol is also produced (Roume et al., 2016). As the waste glycerol purification process is unprofitable, after biodiesel production it stays behind as waste. The production of biodiesel could generate 10% of waste glycerol, and an increase in biodiesel production would raise the need for finding efficient treatment or utilisation of waste glycerol (Ondul & Dizge, 2014). However, some microorganisms have the ability to assimilate waste glycerol and convert it into value-added metabolic products (Chatzifragkou et al., 2011). Streptomycetes are known to have a great potential for the use of glycerol, and some scientists have shown that in addition to pure glycerol, this genus is able to grow and produce active metabolites in medium containing waste glycerol as well (Čirić et al., 2012; Dodd et al., 2018).

Usage of microorganisms in biological control of plant pathogens is considered an alternative to chemical-based treatments, with minimal impact on the environment (Jiang et al., 2019). Certainly, the decreasing effectiveness of fungicides, as well as the risks of their use, indicates the need for more effective and safe control measures (Muthukumar et al., 2016; Sharma et al., 2016). However, production of biofungicide is an important step to its commercial use, but one of the biggest problems of production and application of biological products is their high price. Therefore, considerable interest has been shown in usage of agricultural wastes and by-products from food industry as cultivation medium, in order to

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reduce upstream costs (Nunes, 2012). Regarding this fact, waste glycerol seems to be a promising component in cultivation media used for production of active agents by Streptomyces spp.

In this paper, the ability of Streptomyces hygroscopicus to produce storage apple pathogen antagonist substances in medium with waste glycerol was examined. Production of antifungal metabolites by S. hygroscopicus was carried out in 3 l stirred tank bioreactor at 27 °C through 7 days with aeration rate of 1.5 vvm and agitation rate of 100 r.p.m. The aim of this work was to analyse bioprocess parameters and to determine at which stage of bioprocess the production of antifungal metabolites occurs. Activity of the cultivation liquid on two isolates of Alternaria alternata and two isolates of Fusarium avenaceum obtained from rotten apple fruit were determined every 12 h using in vitro well diffusion method.

1. Materials and methods

1.1. Antagonistic microorganism

In this study, soilborne Streptomyces hygroscopicus was used as antagonistic microorganism, isolated from soil sample in the territory of Novi Sad, Serbia and stored in the Microbial Culture Collection of the Faculty of Technology in Novi Sad, Serbia. This microorganism was deposited under GenBank Accession number KT026467.

The growing medium had the following composition (g l⁻¹): glucose (15.0), soybean meal (10.0), CaCO₃, (3.0), NaCl, (3.0), MgSO₄, (0.5), (NH₄)₂HPO₄, (0.5), and K₂HPO₄, (1.0). The pH of the medium was adjusted to 7.2±0.1 (Consort C863, Turnhout, Belgium) prior to autoclaving.

1.2. Fungal pathogens for bioassay

Two Alternaria alternata isolates (KA10 and T1Jg3) and two Fusarium avenaceum isolates (KA12 and KA13) were used in the study as test microorganisms. The isolates were obtained from apple fruit expressing rot symptoms from Ultra Low Oxygen storages in Vojvodina Province, Serbia and kept as monosporial culture on Potato Dextrose Agar (PDA) slants at 4 °C. The isolates were identified by conventional phytopathological and molecular methods (Polymerase Chain Reaction (PCR) using species-specific primers).

All isolates were initially grown on PDA plates for seven days. A small amount of mycelium of each isolate was added to flasks containing 50 ml of Potato Dextrose Broth (PDB). The flasks were incubated for 48 h on a rotary shaker (150 r.p.m.) at 25 °C. Before activity studies, the growth medium was filtered through double layer of sterile cheesecloth (grade 40).

1.3. Bioreactor studies

Production of bioagents effective against Alternaria and Fusarium phytopathogenic isolates were performed in a bioreactor (Biostat Aplus, Sartorius AG, Germany) with a total volume of 3 l and a working volume of 2 l with internal mixing using two parallel Rushton turbines and without internal baffles. Process parameters were controlled automatically. The ratio of vessel to stirrer diameter (Dv/D) was 2.6. Production of antagonists to apple storage pathogens was performed for 168 h under aerobic conditions with aeration rate of 1.5 vvm and agitation rate of 100 r.p.m.
The medium used for the biosynthesis of antifungal metabolites contained waste glycerol enriched with (NH₄)₂SO₄ (0.25 g l⁻¹) as nitrogen source. Since the waste glycerol used in this study contained 60% of glycerol, the medium for the biosynthesis of active bioagents was formulated to contain 20 g l⁻¹ of total glycerol.

Two litres of fermentation medium was inoculated at 10% (v/v) by a preculture after 72 h growth on a rotary shaker (IKA KS 4000i Control Incubating Shaker) at 150 r.p.m.

1.4. Analytical methods

Cultivation liquid was tested every 12 h of cultivation. The sample of the cultivation medium was centrifuged at 10 000 g (Eppendorf Centrifuge 5804th) for 10 min and the supernatant of the cultivation medium was used for further analysis.

Total nitrogen content was determined in the liquid phase of the cultivation medium using Kjeldahl method (EPA Manual). Standard method was used to determine phosphorus content (GALES et al., 1966).

The residual glycerol content was determined by HPLC measurement. Samples were filtered through 0.45 μm nylon filter (Agilent Technologies, Germany) before analysis. The HPLC instrument was equipped with pump HPG-3200SD/RS, autosampler WPS-3000(T) SL (10-ll injection loop), column ZORBAX NH2 (250 mm × 4.6 mm, 5 μm), and detector Refracto- Max520. As eluent, 75% (v/v) acetonitrile was used with flow rate of 1.2 ml min⁻¹ and elution time of 20 min at column temperature of 30 °C.

Samples (10 ml) were centrifuged at 10 000 g for 10 min at 20 °C. The supernatants were discarded and the cell pellet was re-suspended in an equal volume of distilled water and re-centrifuged as above. The cell biomass was dried at 105 °C overnight and weighed. All determinations were performed in duplicate (MEANWELL & SHAMA, 2008).

1.5. In vitro assay

In vitro antagonistic activity assay was performed in 85 mm Petri dishes using wells technique (TADIAN et al., 2016). In short, two layers of PDA medium were poured. The first layer consisted of 2% PDA medium. After solidification, the second layer composed of 1.2% PDA and filtered fungal culture liquid (35%) was added. Three wells per plate with a diameter of 10 mm were made, two plates represented one treatment. For each treatment, 100 μl of Streptomyces hygroscopicus supernatant was added to each well. After 72 h of incubation at 25 °C, diameter (mm) of mycelia growth inhibition zone around the wells was measured.

1.6. Data analysis

Inhibition zone diameter data obtained by testing the supernatant of S. hygroscopicus through 7 days of cultivation on four test isolates were processed by factorial ANOVA using Software STATISTICA 13.

2. Results and discussions

2.1. Time-course of substrate consumption and cell growth during cultivation

In order to monitor the ongoing processes in the laboratory bioreactor, contents of the most important nutrients, i.e. sources of carbon, nitrogen, and phosphate, during the cultivation
were measured. Also, changes in the biomass of the producer microorganism were monitored as dry weight of cells. The results are shown in Figure 1.

Fig. 1. Time course of residual glycerol content (A), residual nitrogen content (B), cell biomass (C), and residual phosphates (D) during 7 days of \textit{S. hygroscopicus} cultivation in 3 l bioreactor.

Based on the data obtained (Fig. 1A), it can be observed that consumption of glycerol is more intense at the start of biosynthesis up to 108 h. This indicates that \textit{Streptomyces hygroscopicus} uses this nutrient during growth phase. Glycerol content was reduced to 9.29 g l$^{-1}$ by the end of the bioprocess, but the residual glycerol can be used as a binder of the final product to plants, which would significantly affect the economy of production (Grahovac et al., 2014). Also, this is an indication that \textit{S. hygroscopicus} uses glycerol obtained as a by-product of biodiesel production very well (Čirić et al., 2012; Dodo et al., 2018). On the other hand, nitrogen and phosphorus are required by microorganisms both for growth and multiplication, and very often for the production of desired metabolites. Certainly, phosphorus affects the control of the biosynthesis of a large number of secondary metabolites (Martin, 2004). Nitrogen (Fig. 1B) and phosphorus (Fig. 1D) consumption is most intense during the exponential phase, and their values at the end of the bioprocess are 0.238 g l$^{-1}$ and 0.171 g l$^{-1}$, respectively.

Results of biomass growth show that exponential phase lasts until the fourth day, when stationary phase of the bioprocess, in which the consumption of nutrients is significantly reduced, begins (Fig. 1C). It can be assumed that the applied conditions did not favour biomass production. Also, it can be concluded that consumption of glycerol, nitrogen, and phosphorus is connected to biomass formation.
2.2. Effect of cultivation time on the production of active agents

Applied conditions were selected according to results from the scientific literature, showing that lower agitation speed and aeration rate of 1.5 vvm represent optimal conditions for the production of desired metabolites using streptomycetes (Techapun et al., 2003; Yen & Li, 2014). As the most favourable form for metabolite production by streptomycetes is small pellets (Ilić et al., 2008), high agitation speed can destroy the desired pellet form, thus adversely affecting the productivity of streptomycetes. The effect of Rushton turbine on the morphology of streptomycetes depends on the applied mixing rate. Also, with increased aeration rate, stationary phase, in which the desired antagonist agents are produced, occurs earlier (Gottschalk et al, 2003). The statistical analysis was performed to determine whether the duration of cultivation affected the production of antifungal metabolites effective against four isolates tested.

### Table 1. Results of factorial analysis of variance: sources of variation of inhibition zone diameter during 7 days of cultivation

| Source of variation | SS     | Degree of freedom | MS     | F-value | P-value |
|---------------------|--------|-------------------|--------|---------|---------|
| Time                | 12658.2| 14                | 904.2  | 1251.9  | 0.00    |
| Test fungi          | 1602.2 | 3                 | 534.1  | 739.5   | 0.00    |
| Time×Fungi          | 1543.2 | 42                | 36.7   | 50.9    | 0.00    |
| Error               | 86.7   | 120               | 0.7    |         |         |

As expected, cultivation time had the most significant effect on inhibition zone diameter. Also, statistically significant differences (P<0.05) in the inhibition zone diameter were observed between the fungal isolates. Table 1 shows that interaction between these two factors (cultivation time and test fungi) also significantly affected inhibition zone diameter.

![Fig. 2. Mean values of inhibition zone diameter (mm) of test phytopathogenic fungi caused by supernatant filtrate of S. hygroscopicus cultivation during 7 days of cultivation](image-url)

Fig. 2. Mean values of inhibition zone diameter (mm) of test phytopathogenic fungi caused by supernatant filtrate of S. hygroscopicus cultivation during 7 days of cultivation

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Figure 2 presents the mean values of inhibition zone diameter of two A. alternate and two F. avenaceum isolates caused by S. hygroscopicus supernatant during 7 days of cultivation on medium containing waste glycerol. Obtained results show that impressive antifungal activity was achieved between 4 and 5 days of S. hygroscopicus cultivation under defined conditions. According to Tónlé and co-workers (2016), inhibition zone radius over 11 mm indicated that the applied antifungal agent is highly efficient. In our study, these inhibition zone diameters were formed after 60 hours of cultivation. This is important information, because it allows the bioprocess to be shortened, increasing its cost-effectiveness.

However, the largest inhibition zone diameters were formed at 108 h of cultivation for all phytopathogenic test isolates. Analysing results presented in Figure 2, it can be concluded that A. alternate T1Jg3 is the most sensitive against produced bioagents (average inhibition zone 44 mm), followed by A. alternata KA10 (41.33 mm). On the other hand, Fusarium isolates show less sensitivity to the produced bioagents, forming inhibition zone diameters of 38.67 mm for KA13 isolate and 35.67 mm for KA12 isolate. Since the tested isolates belong to different genera, their sensitivity to antifungal components could be different. Control of these types of phytopathogenic fungi is important, because they cause infections of apple fruit during storage, but also they are known as producers of mycotoxins harmful to human health (Koncz et al., 2009; Elhairy et al., 2016).

3. Conclusions

The results obtained from these studies indicate the high potential of Streptomyces hygroscopicus to produce antifungal metabolites in medium containing waste glycerol in laboratory stirrer tank bioreactor at aeration rate of 1.5vvm and agitation of 100 r.p.m. Under these conditions, pellets responsible for the productivity of streptomycetes are formed.
Results also showed that cultivation time has a statistically significant effect on production of antifungal agents by \textit{Streptomyces hygroscopicus}. Anyhow, extending the bioprocess duration over 7 days would be considered economically unprofitable. Optimisation of the most important medium nutrients was not performed in this study, which is one of the reasons why not all nutrients of the medium were consumed at the end of the bioprocess. Also, use of different carbon sources as well as different mixing and aeration rates significantly affects the utilisation of the medium nutrients and the trend of production of antifungal metabolites using \textit{S. hygroscopicus}. The in vitro study confirmed that the produced agents can successfully be used for biological control of stored apple fruit infected with phytopathogenic isolates \textit{A. alternate} KA10 and T1Jg3 and \textit{F. avenaceum} KA12 and KA13.

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