

## Biosurfactant Production by *Pseudomonas Aeruginosa* Strains on 2ml of Inoculum Size

SHOAIB ASIF, HAROON HABIB, MUDASER HUSSAIN ABBASI, RANA MUHAMMAD AKHTAR, MIAN ABDUR RASHEED, MUHAMMAD WAQAS, RANA M. SHAHID ZAFAR, HEENA AWAIS,

### ABSTRACT

**Objective:** To produce biosurfactants from *Pseudomonas aeruginosa* using agricultural resource and to produce Biosurfactants using low cost materials.

**Study design:** Descriptive study

**Place and duration of study:** Study was conducted at Institute of molecular biology and biotechnology in university of Lahore. Duration of the study was two years.

**Methods:** The volume of sample taken are 2ml, of innoculum from growing culture of *Pseudomonas aeruginosa* was isolated from contaminated soil collected from industrial area of District Kasoor and flasks were than placed into an orbital shaker at speed of 120rpm. The samples were collected in sterile screw capped bottle, 4-5 cm deep from the soil surface aseptically. The samples were stored at 4°C till further use. After every 24h the culture broth from each flask was taken to estimate bacterial cell mass.

**Results:** Surface tension was 65.8, 52.6, 47.2 and 33.8 mN/m at time 24, 48, 72 and 96 hours respectively at constant temperature of 37°C and molasses used 0.25g with 2ml inoculum size. The rhamnolipid production was 0.38, 0.97, 1.92 and 2.21 g/L respectively. Similarly the bacterial cell mass was 0.3, 0.65, 0.75 and 0.8 g/L respectively

**Conclusion:** After optimizing various growth and environmental factors a production of rhamnolipid was achieved .

**Keywords:** Biosurfactants, Molases, *Pseudomonas aeruginosa*

---

### INTRODUCTION

In recent years greater emphasis has been placed on the environmental impacts of chemical surfactants and new surfactants for use in the pharmaceutical and biomedical. For example, a range of new nonionic gemini aldonamide-type surfactants consisting of two hydrophobic chains and two aldonamide polar head groups fused with a linker region have been developed that have low critical micelle concentration values ( $3.8 \times 10^{-6}$  to  $1.3 \times 10^{-4}$  M)<sup>1</sup>. Biosurfactants are amphiphilic biological compounds produced extracellularly or as part of the cell membrane by a variety of yeast, bacteria and filamentous fungi from various substances including sugars, oils and wastes. However, carbohydrates and vegetable oils are among the most widely used substrates for research on biosurfactant production by *Pseudomonas*

*aeruginosa* strains<sup>2</sup>. Bacteria of the genus *Pseudomonas* are known to produce glycolipid surfactant containing rhamnose and 3-hydroxy fatty acids<sup>3</sup>. Rhamnolipids produced by *Pseudomonas aeruginosa* have been widely studied and reported as a mixture of homologous species RL1 (RhC<sub>10</sub>C<sub>10</sub>), RL2 (RhC<sub>10</sub>), RL3 (Rh<sub>2</sub>C<sub>10</sub>C<sub>10</sub>) and RL4 (Rh<sub>2</sub>C<sub>10</sub>)<sup>4</sup>. From a combined application/cost perspective rhamnolipid, produced by *P. aeruginosa*, represents the leading commercial microbial biosurfactant and hence this brief discourse on industrial biosurfactant production will be confined to this product/host system. Extensive investigations have been implemented at both the molecular and cell culture level aimed at understanding factors influencing rhamnolipid biosurfactant biosynthesis by *P. aeruginosa* with a view to optimising the fermentation process<sup>4</sup>. Molasses: Molasses is a co-product of sugar production, both from sugar cane as well as from sugar beet. It is defined as the runoff syrup from the final stage of crystallization, in which further crystallization of sugar is uneconomical. Molasses generally consists of 48-56% total sugar (mainly sucrose), 9-12% non-sugar organic matter, 2-4% protein (N×6.25), 1.5-5% potassium, 0.4-0.8% calcium, 0.06% magnesium, 0.6-2.0% phosphorus,

1.Biochemist Scholar 2.Lecturer Biochemistry in Avicenna Medical College, Lahore 3.Associate Prof. Forensic Medicine & Toxicology Avicenna Medical College Lahore 4. Prof. Community Medicine, Avicenna Medical College Lahore 5.Prof.Forensic Medicine& Toxicology,Shaheed Benazir Bhutto Medical college,Azad Kashmir 6.Biochemist Scholar 7.M-phil student 8.Physiotherapist  
Correspondence to Mr. Shoaib Asif, Biochemist Email: biofilm@yahoo.com

1.0-3.0 mg/kg biotin, 15-55mg/kg pantothenic acid, 2500-6000 mg/kg inositol and 1.8mg/kg thiamine (4). Different kinds of bacteria have been employed by many researchers in producing biosurfactant using culture media. Most of such bacteria used are isolated from contaminated sites usually containing petroleum hydrocarbons by-products and/or industrial wastes<sup>5</sup>.

There is shortage of this type of studies so this study was designed to produce rhamnolipid (A glycol-lipid biosurfactant composed of one rhamnose unit and a lipid tail) by *Pseudomonas aeruginosa* using agricultural resources i.e., molasses.

## MATERIALS AND METHODS

It was designed to optimize the inoculum size for the production of rhamnolipid. The volume of sample taken are 2ml, of inoculum from growing culture of *Pseudomonas aeruginosa* was isolated from contaminated soil collected from industrial area of District Kasoor and flasks were then placed into an orbital shaker at speed of 120rpm. The samples were collected in sterile screw capped bottle, 4-5 cm deep from the soil surface aseptically. The samples were stored at 4 °C till further use (7). After every 24h the culture broth from each flask was taken to estimate bacterial cell mass. All the chemicals including L-rhamnose, Orcinol reagent, Diethyl ether, Molasses, Na<sub>2</sub>HPO<sub>4</sub>, K<sub>2</sub>HPO, MgSO<sub>4</sub>, NaH<sub>2</sub>PO<sub>4</sub>, FeSO<sub>4</sub>, Peptone were purchased from Sigma Aldrich from their local distributor in Lahore, Pakistan. The bacterial strains were isolated from the industrial contaminated soil by using soil enrichment technique. Briefly; 1g soil from sample, in 100ml sterile mineral salt media with 1g of molasses was incubated for 96 hours at 37 °C on an orbital shaker at 100 revolutions per minute. After enrichment, 2ml cell suspension was taken from the flask and spread over nutrient agar plate and was incubated at 30 °C for 48 hours. Colonies that appear on nutrient agar plates were selected randomly and sub-cultured to obtain pure isolates.(7). An organic nitrogen medium, with phosphate was prepared. The composition of the medium was (gL<sup>-1</sup>): NaH<sub>2</sub>PO<sub>4</sub> .H<sub>2</sub>O, 4.0, Na<sub>2</sub>HPO<sub>4</sub>

.H<sub>2</sub>O, 1.0, MgSO<sub>4</sub> .7H<sub>2</sub>O, 1.0, CaCl<sub>2</sub> .2H<sub>2</sub>O, 0.005, Peptone, 1.38, 25ml of glycerol was used as source of carbon substrate<sup>8</sup>. A total of 2.5litres of distilled water was used, hence the above measured weights and volume respectively was calculated based on that. The pH of the medium was adjusted to 7 using 211 Microprocessor pH meter with 1.0M NaOH. Sixteen Erlenmeyer flasks (250ml) were used during the experiment. 150ml of the prepared medium was measured into each flask using a 200ml measuring cylinder. Each flask was clogged using cushion foam and covered with Aluminium foil. The prepared medium was autoclaved for 3 days before being inoculated. Nutrient broth media (100ml) was inoculated with bacterial strain and growth was monitored at 37°C in shaking incubator at 100 rpm for 72 hours<sup>9</sup>.

## RESULTS

It was designed to optimize the inoculum size for the production of rhamnolipid. Various volumes of inoculum were taken and added into the fermentation media. The experiment was monitored for 96hours and temperature was set at 37°C and pH was set at 7. The volumes taken 2ml of inoculum from growing culture of *Pseudomonas aeruginosa* and flasks were then placed into an orbital shaker at speed of 120rpm. After every 24h the culture broth from each flask was taken to estimate bacterial cell mass, rhamnolipid estimation and surface tension reduction. (Tables 1). Biosurfactant Production is a growth associated production, parallel relationships exist between production, substrate utilization and biosurfactant production. The production of rhamnolipid by *Pseudomonas* species is an example of growth associated biosurfactant production (10). The results of the present study(table-1 and fig-1&2) revealed that surface tension was 65.8, 52.6, 47.2 and 33.8 mN/m at time 24, 48, 72 and 96 hours respectively at constant temperature of 37°C and molasses used 0.25g with 2ml inoculum size. The rhamnolipid production was 0.38, 0.97, 1.92 and 2.21 g/L respectively. Similarly the bacterial cell mass was 0.3, 0.65, 0.75 and 0.8 g/L respectively.

Table 1: Results with 2ml inoculum size

Time Hours	Inoculum size ml	Temp. °C	Molases g	Surface Tension mN/m	Rhamnolipids g/L	Bacterial cell mass g/L
24	2	37	0.25	65.8	0.38	0.3
48	2	37	0.25	52.6	0.97	0.65
72	2	37	0.25	47.2	1.92	0.75
96	2	37	0.25	33.8	2.21	0.8

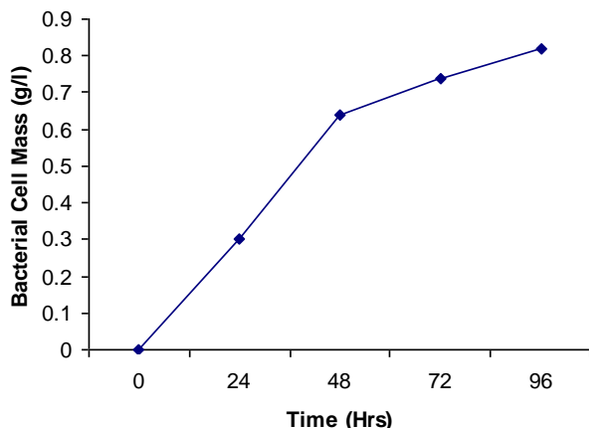
**Bacterial Cell Mass:**

Fig. 1: Estimation of Bacterial cell mass using 2 ml inoculum size

The fig. 1 represents that bacterial cell mass (g/L) increased with the passage of time as revealed in the fig that at zero time the bacterial cell mass was zero and it increased to 0.8g/L bacterial cell mass when the time passage was 96 hours.

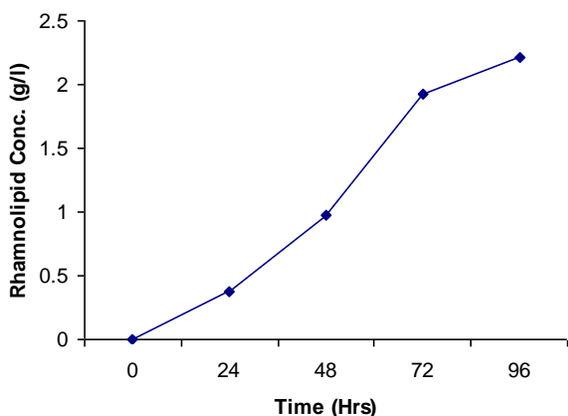


Fig. 2: Estimation of Rhamnolipid production using 2 ml inoculum size

The fig. 2 represents that rhamnolipid concentration (g/L) increased with the passage of time as revealed in the fig that at zero time the rhamnolipid concentration was zero and it increased to 2.21g/L rhamnolipid concentration when the time passage was 96 hours.

**DISCUSSION**

Inoculum size is one of the most important parameter for the production of microbial metabolites. Microorganisms required a certain cell number in a

particular media to start their rapid growth and metabolite production (log phase) so it is important to determine the exact initial bacterial size to start an experiment leading to the successful end. As rhamnolipid is a growth associated process<sup>10,11</sup>, the optimization of the inoculum size in proposed media was the most important parameter to be optimized. it was found that 1ml inoculum size was best for rhamnolipid production during this research. it was shown by that biosurfactant production was growth associated so increase in inoculum size will increase the nutritional demand by microorganisms<sup>10,11</sup> so it was very essential for the experiment to maintain a balance between the inoculum size and the volume of the media component as it effected the biosurfactant production shown by the results. Biosurfactant Production is a growth associated production, parallel relationships exist between growth, substrate utilization and biosurfactant production. The production of rhamnolipid by *Pseudomonas* species is an example of growth associated biosurfactant production<sup>10,11</sup>.

**CONCLUSIONS**

After optimizing various growth and environmental factors a production of rhamnolipid was achieved

**Suggestions<sup>11</sup>**

1. Various inoculum sizes such as 3ml and 4ml/100ml of broth should be tested
2. At the end rhamnolipid production, surface tension and bacterial cell mass was should be estimated .
3. Such product can be used for numerous industrial, therapeutic, biomedical, and environmental applications.

**Acknowledgments:** the authors highly acknowledge the honourable dean dr Saghir Ahmad Jafri, TI, AF , Dean, Faculty of Sciences, Director, IMBB, Mr. Asif Jamal, IMBB The University of Lahore.

**REFERENCES**

1. Syldatk C and F Wagner (1987). Production of biosurfactants. Editors, Biosurfactants and Biotechnology, Surfactant Science Series vol. 25: Marcel Dekker, New York , 89–120.
2. Mata SJC, J Karns and A Torrents (2000). Effects of rhamnolipids produced by *Pseudomonas aeruginosa* UG2 on the solubilization of pesticides. Environmental Science and Technology , 34: 4923–4930.
3. Lang and Wagner, 1987 Lang S and F Wagner (1987). Structure and properties of biosurfactants Editors, Biosurfactants and Biotechnology, Marcel Dekker: New York , 21–45.
4. Maier RM and G Soberon-Chavez (2000). *Pseudomonas aeruginosa* rhamnolipids: biosynthesis and potential applications. Applied Microbiology and Biotechnology , 54: 625–633

5. (Makkar and Cameotra, 1997). Makkar RS and SS Cameotra (1997). Utilization of molasses for biosurfactant production by two *Bacillus* strains at thermophilic conditions. Journal of the American Oil Chemists' Society , 74: 887–889.
6. (Burd and Ward, 1996). Burd G and OP Ward (1996). Bacterial degradation of polycyclic aromatic hydrocarbons on agar plates the role of biosurfactants . Biotechnology Techniques, 10: 371–374.
7. Anna *et al*, 2002 Anna LMS, GV Sebastian, EP Menezes, TLM Alves, AS Santos, N Pereira and DMG Freire (2002). Production of biosurfactants from *Pseudomonas aeruginosa* PA1 isolated in oil environments. Brazilian journal of Chemical Engineering, 56: 159-166.
8. Patel and Desai, 1996 Patel RM and AJ Desai (1996). Biosurfactant production by *Pseudomonas aeruginosa* GS3 from molasses. Letters in Applied Microbiology , 25: 91–94.
9. (Wei *et al*, 2005). Wei Y, L Wang and J Chang (2005). Optimizing iron supplement strategies for enhanced surfacting production with *Bacillus subtilis*. Biotechnol Prog , 20: 979–983.
10. Desai JD and IM Banat (1997). Microbial production of surfactants and their commercial potential. Microbiological Molecular Reviews , 61” 47–64.
11. Shoaib Asif, Haroon Habib, Mudaser Hussain Abbasi (2013). Biosurfactant production by *Pseudomonas aeruginosa* strains on 1 ml of inoculum size , Pakistan journal of Medical & Health sciences, Vol.7, Apr-Jun.2013, 421.