Original Research Article

ENUMERATION OF POLYHYDROXYALKANOATE (PHA) PRODUCING BACTERIA FROM DAIRY SEWAGE SAMPLES

ABSTRACT:

Aims: The synthetic polymer plastics have become an integral part of contemporary life. Excess use of plastics and indiscriminate dumping of it in soil and water is polluting the environment. In order to overcome this problem, the production and applications of eco-friendly biodegradable products from microbes are becoming inevitable from the last decade and also are the good alternatives for synthetic polymers.

Methods and Results: Polyhydroxyalkanoate producing bacterial strains were confirmed by serial dilution of sewage samples from dairies and pour plating using modified nutrient agar medium with 2% glucose and 0.3% sudan black. Commercial dairy sewage sample from III Dairy showed highest count of PHA producers (3.80 log₁₀cfu/ml) followed by II Dairy (3.68 log₁₀cfu/ml) and I Dairy (3.35 log₁₀cfu/ml). On an average, 70 per cent were PHA producers among TBC of sewage samples.

Conclusion: Dairy sewage sample from III Dairy showed highest count of PHA producers (3.80log₁₀cfu/ml)

Significance and Impact of the Study: This study provides importance of polyhydroxyalkanoates and their role against synthetic plastic by enumerating the polyhydroxyalkanoate (PHA) producing bacteria from Dairy sewage samples that can be effectively utilized for the synthesis of bioplastics.

KEYWORDS: Polyhydroxyalkanoate, synthetic plastic, Thermoplastic, Biodegradable.

1. INTRODUCTION:

"Plastics" are polymers stemming from petro-chemistry. Based on favourable material features such as low density, high resistance, and well-optimized manufacturing processes, plastics are by far the fastest emerging group of materials used for manufacturing of customized items. In this context, plastics are needed for packaging of diverse goods, agriculture, electronics, construction industry, transportation, health care, or the sport and leisure sector. However, based on the limitation of petrochemical resources and the recalcitrance of plastics towards biodegradation, there is a growing global concern related to traditional plastics of petrochemical origin. Reliable estimates speak about a pile of 8 to 9 x 10⁹ tonnes of plastics that have been made globally in recent decades urgently require proper disposal (Koller and Braunegg, 2018).

Biodegradable polymers offer an alternative, which, due to their technical and economic viability, present a great potential for expansion. Biodegradable polymers are defined as materials that undergo breakage of their chains by the action of microorganisms, resulting in decomposition under specific conditions of pH, humidity, oxygenation, and the presence of catalytically active metals. Biodegradable polymers can be produced from derivatives of

fossil sources and renewable natural resources, as well as can be synthesized by bacteria (Ray and Bousmina, 2005). Among the biodegradable polymers, Polyhydroxyalkanoates (PHAs) are biodegradable polymer family that are produced by diverse microorganisms as Polyhydroxybutyrate (PHB) such as *Pseudomonas putida, Ralstonia eutropha* and *Bacillus megaterium* under nutrient-starved conditions (Jeon *et al.*, 2014). In 1926, French scientist Maurice Lemoigne of Pasteur Institute discovered intracellular accumulation of 3-hydroxybutyric acid polymers in *Bacillus megaterium*, and this was the first report of PHB accumulation in bacteria. These biopolymers are accumulated in the cell to store their utilized carbon sources in depletion of nutrients. PHA has been explored over the years because of its relevant thermal and mechanical properties; furthermore, it can be obtained from renewable resources, degrading enzymatically in different ecosystems such as water, soil, and sludge, among others (Siracusa *et al.*, 2017).

Scanty literature is available regarding enumeration of PHA producing bacteria directly as many scientists have done after colonies are formed by staining. The current study was aimed at enumeration of the PHA producing bacteria from various dairy sewage samples directly.

2.0 MATERIALS AND METHODS:

2.1 Collection of dairy sewage samples

Three Dairy sewage samples were collected in sterile bottles one from student experimental dairy (I Dairy), Dairy Science college, KVAFSU, Hebbal,

Bangalore and other two samples from commercial dairies of Bengaluru (II & III Dairies).

2.2 Characterization of dairy sewage samples

The collected dairy sewage samples were subjected to determination of pH, BOD and COD using standard procedure of BIS (IS 3025 (Part 44): 2003)(IS 3025 (Part 58):2006)

2.3 Enumeration of PHA producing bacteria

The dairy sewage samples were serially diluted and pour plated for enumeration of PHA producers. Dairy sewage samples of 11 ml was pipetted and transferred to the sterile 99 ml flask containing physiological saline to make 1st dilution. Further required dilutions were prepared serially using 1st dilution. Serially diluted samples were transferred to labelled sterile petri plates for the enumeration of PHA producing bacteria using sterile pipettes. Sterile molten modified nutrient agar with 2% glucose and 0.3% sudan black (Paul *et al.*, 2017) maintained at 55°C water bath was poured to labelled plates containing 1 ml of dilution and mixed thoroughly without spilling the medium. Later the poured agar plates were allowed to solidify. All the poured plates were incubated at 37° C/48 h by inverting the plates. After the completion of the incubation period, the colonies with dark black coloured colonies were considered as PHA produced and were counted in countable plates ranging between 30-300 and average count was expressed as log₁₀ cfu/ml of the sewage sample.

3. RESULTS AND DISCUSSION

All the three dairy sewage samples were neutral with COD on an average of 2944 and BOD of 1400 ppm (Table 1). All the dairy sewage samples from commercial dairies such as I, II and III were subjected to enumeration of PHA bacteria using modified Nutrient agar enriched with 2% glucose and 0.3% sudan black, the plates were incubated at 37° C/48 hrs (Table 2). I dairy sewage sample showed 3.35 log₁₀cfu/ml of PHA producer followed by II dairy sewage of 3.68 log₁₀cfu/ml and III dairy sewage sample of 3.80 log₁₀cfu/ml. Nehra *et al.* (2015) revealed that the colonies obtained on sterile modified nutrient agar medium when flood with sudan black observed black coloured colonies indicating the presence of PHA producers. Many literature identified the PHA producers on nutrient agar after colony formation. But in present research work an attempt was made to use modified nutrient agar with 2% glucose and sudan black (0.3%) and found to be better medium for direct selection of PHA producers.

Table 1: Chemical characteristics of collected Dairy Sewage Samples

Sample Code	рН	COD	BOD
I Dairy	6.8 ^b	2100 ^b	800 ^b
II Dairy	7.2 ^a	2400 ^{ab}	1500 ^{ab}

III Dairy	7.2 ^a	3200 ^a	1900 ^a
CD (P=.05)	0.10	431.79	307.46

Note:

- Values are average of three trials
- Lower case alphabets as superscript indicate significant difference
- CD Critical Difference
- Significantly there was difference in COD and BOD values

Table 2: Enumeration of PHA Producers of collected Dairy Sewage

Samples

Sample Code	ТВС	PHA	
	(log ₁₀ cfu/ml)		
Dairy I	5.00 ^a	3.35 ^a	
	(100 %)	(67%)	
Dairy II	5.07 ^a	3.68 ^a	
	(100 %)	(72.6 %)	
Dairy III	5.32 ^a	3.80 ^a	
	(100 %)	(71.4%)	
CD (<i>P</i> =.05)	0.43	0.54	

Note: Values are average of three trials

- ➤ Lower case alphabets as superscript indicate significant difference
- > TBC Pour plated using standard plate count agar and incubated at 37° C/72 h
- ▶ PHA Pour plated using Modified Nutrient Agar enriched with 2% glucose and 0.3 % Sudan Black and incubated at 37° C/ 72 h.
- > Colonies with black colour were counted as PHA producer

4. CONCLUSION:

In the present research work, dairy sewage samples were collected, determined for pH, BOD and COD and further serially diluted and pour plated using 2% nutrient agar with sudan black for the enumeration of PHA producing *Bacillus* spp. Many of the literature revealed that *Bacillus* spp. resulted as potential PHA producing organism by showing positive with Sudan black B staining.

5. REFERENCES

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