Isolation and characterization of oleaginous yeasts from dairy waste

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Abstract: Microbial lipids (biodiesel) can be produced by oleaginous yeasts via converting carbohydrates into lipids under certain conditions. Sudan black ‘B’ staining technique was used to screen cellular lipid accumulation. Extraction of the bacterial lipids was carried out by Bligh and Dyer method and fatty acid methyl esters were analysed by GC. Primary screening of sample resulted in 22 isolates, out of which 6 isolates were found to generate lipid. After detection of good oleaginous lipid productivity, one best yielding isolate was preferred for upward spring. Secondary selection of potential isolate was based on the production in addition to parameters like pH and nutritional requirements. Lipid from selected isolate CNK 1 was subjected to determination of saponification value and PUFA screening. The isolate CNK1 was identified as Zygrosaccharomyces rouxii by FAME analysis. The above results were promising hence it has importance for supplementary development of the yeast isolate Zygrosaccharomyces rouxii as another source of lipid for biodiesel production.

Keywords: FAME-GC, PUFA, Oleaginous Yeast; Sudan Black B, Zygrosaccharomyces rouxii,
Materials and Methods

Soil sample collection

Soil sample were collected from district milk collection centre, Nanded (industry waste area).

Enrichment of microorganism

1 gm soil sample was added in to 250 ml flask containing 100 ml sterile enrichment medium (composition (g/L): Glucose-100, Yeast extract-1, NH₄Cl-1, KH₂PO₄-2, MgSO₄-0.75, CaCl₂-0.05, ZnSO₄-0.01, FeCl₃-0.01 and Na₂HPO₄-1). This mixture was cultured at 28 C, 120 rpm for 48 hrs. Serially diluted soil samples were spread on sterile enrichment medium plate. Then plates were kept in incubator at 28 C for 1-4 days. After incubation each separate colonies were picked and stored on same medium agar slant for further study. Screening was done by lipid staining (Burdan K 1946).

PUFA producers screening / H₂O₂ plate assay method

PUFA producers screening was done as per the method mentioned by (Ashwini et al. 2012). YPD medium containing (1mM) NaN₃ were prepared and sterilized and isolated strains were spread on plates. After spreading sterile filter paper disc (5mm) were placed on medium. 10 µl of H₂O₂ solution having different concentration (0.5% and 1%) were added on separate filter paper disc on the plate. Plates were incubated at 28°C for 24- 48 hrs.

FAME analysis

The fatty acids were extracted by a procedure which consists of saponification in sodium hydroxide/methanol solution followed by derivatization with hydrochloric acid/methanol solution to give the respective fatty acid methyl esters (FAMEs). The FAMEs are then extracted from the aqueous phase by the use of an organic solvent and the resulting extract was analysed by GC. FAMEs are more volatile than their respective fatty acids and therefore more suitable to GC analysis. The Sherlock software automates all analytical operations and uses a sophisticated pattern recognition algorithm to match the unknown FAME profile to the stored library entries for identification (Anju et al. 2011).

Production and extraction of lipid

100 ml fermentation medium were prepared and sterilized at 121 C for 15 mins and cultures of positive screened organism were inoculated into fermentation medium. Flasks were incubated at 28 C, 120 rpm for 6 days. These set were used to determine biomass, dry weight and lipid content. Extraction of lipid was done by (Bligh and dyer 1959) method. 100 ml cultured medium were centrifuged at 5000g for 10 mins. Supernatant was discarded, pellet was washed with sterile distilled water, dried and weight was taken. 15 ml of 4M HCl was added and kept at room temperature for 30 mins. This mixture was kept in freezer (-20 C) for 20 minutes and immediately transfered to boiling water bath for 10 min. 30 ml chloroform : methanol mixture was added and centrifuged at 5000 g for 10 min. Lower layer chloroform contained lipid were collected and used for further study.

Estimation of lipid

Estimation of total lipid was carried out by (Barnes and blackstock 1973) method.

Production media optimization study

Effect of pH on biomass lipid production

100ml fermentation medium of different pH (4.5, 5.5, 6.5, 7.5 & 8.5) were prepared. Isolated strains were inoculated into each different pH medium and incubated at 28 C, 120 rpm for 6 days for lipid production. After incubation the biomass and lipid content were checked in each pH medium. The lipid content was determined by lipid estimation. The optimum pH medium suitable for isolated strain was determined.

Effect of carbon source on biomass and lipid production

100ml of fermentation medium of different carbon source (glucose, sucrose, lactose, maltose, and fructose) were prepared. Isolated strains were inoculated into each different carbon source medium and incubated at 28 C, 120 rpm for 6 days for lipid production. Biomass and lipid content were checked in each carbon source medium. The high lipid content was determined by lipid estimation. The optimum carbon source medium suitable for isolated strain was determined.

Effect of nitrogen source on biomass and lipid production

100ml fermentation medium of different nitrogen source (organic nitrogen source: yeast extract, peptone and inorganic nitrogen source: NH₄Cl, KNO₃, (NH₄)₂SO₄, NH₄NO₃, NaNO₃). Isolated strains were inoculated into each different nitrogen source medium and incubated at 28 C, 120 rpm for 6 days for lipid production. After incubation the biomass, lipid content was checked in each nitrogen source medium. The high lipid content was determined by lipid estimation and the optimum nitrogen source medium suitable for isolated strain was determined.

Saponification value

Saponification value was determined by the methods of Association of Official Analytical Chemists (1990). 4-5 gm lipid sample was added in to flask and 50 ml alcoholic KOH was added. Blank was prepared by taking only 50 ml alcoholic KOH. Air condenser was connected to the flask and reaction mixture was boiled for 60 min. 1ml phenolphthalein indicator was added and

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reaction mixture was titrated against 0.1 N HCl up to pink colour disappears.

**Statistical analysis**

All the incubations were performed at triplicates and the data was analysed using MS-Excel and the result were expressed as the mean ± SD.

**Results and Discussion**

Soil sample were collected from district milk collection centre, Nanded (industry waste area) and successfully used for isolation of lipid producers. Total 43 isolates were obtained on nitrogen limited medium. Out of these isolates, 22 isolates were recovered and used for screening of lipid producers. Twenty two isolates preliminary screened trough Sudan black B staining method and six isolates showed lipid producers (Figure 1). Extraction of lipid was done by using advanced Bligh and dyer method. Amount of lipid produced in test sample was calculated by interpolating optical density of test sample on standard lipid curve and the estimated concentration of lipid was found to be 0.098gm/100ml. The Microbial Identification System (MIDI) for fatty acid methyl ester (FAME) analysis is a standard method for identification of microorganisms (Schutter et al. 2000). Whole cell fatty acids are converted to methyl esters and analysed by gas chromatography. The fatty acid composition of the unknown is compared to a library of known organisms in order to find the closest match. The list of the fatty acids composition like straight Chain fatty acids 21.84%, branched chain fatty acid 25.50%, Mono Unsaturated Fatty Acid 12.68% and oleic acid (C18:1w9c) 38.36% was given clearly according to the GC report. The FAME GC analysis data obtained (Table 1) is more descriptive and elaborative. Our experimental data matches and establish the similar result mentioned in the report of MIDI Sherlock software databases and the similarity was matched with organism *Zygosaccharomyces rouxii* with selected ion monitoring (SIM) index 0.49 (Figure 2). Based on morphological characteristics and FAME-GC analysis it was identified as *Zygosaccharomyces rouxii*. Ravikumar et al. (2012) worked on biodiesel production from oleaginous fungi which involves the mixture of fatty acyl methyl/ethyl esters, produced from transesterification of neutral lipids. In similar way, Mrinal et al. (2011) has studied the comparative lipid profiling of endophytic fungi by FAME analysis techniques. Gao et al. (2010) also studied the screening, fermentation and optimisation of microbial lipid producing molds.

**Production media optimization study**

*Effect pH on biomass and lipid production*

Effect of pH on biomass and lipid production were studied, the strains was incubated at varying pH (4.5 to 8.5). Outcomes of the stated reports (Figure 3) shows increase in the biomass as well as lipid yield with respect to the increase in pH value but the relevant increase was maximum at 7.5 and shows minimum at 8.5 pH respectively.

*Effect of carbon source on biomass and lipid production*

Effect of carbon source on biomass and lipid production was studied; different carbon sources were used in the production medium (Figure 4). The result showed lactose, maltose, glucose, sucrose and fructose were suitable carbon source for the growth and lipid accumulation of the isolated strain among which glucose is the most suitable. The lipid production was 0.092 g/100ml.

*Effect of Nitrogen source on biomass and lipid production-

The effect of nitrogen source on biomass and lipid production was shown in (Figure 5). Organic nitrogen source (yeast extract and peptone) whereas inorganic (KNO₃, NH₄SO₄, NH₄NO₃, NH₄Cl and NaN₃) were used. In the organic nitrogen source yeast extract is the suitable and lipid production was 0.098 g/100ml. Similarly, NH₄Cl is the most suitable inorganic nitrogen source; lipid production was found to be 0.098 g/100ml.

**PUFA screening**

As shown in (Figure 6), yeast cells were grown in presence of H₂O₂ to check membrane shielding effect of PUFA. PUFAs are the most vulnerable to oxygen and ROS. To confirm this, growth of yeasts in presence of H₂O₂ was in reality mainly due to presence of PUFA; NaN₃ was added into the media which is a very powerful inhibitor of catalase. If microorganism is producing catalase enzyme, NaN₃ inhibits catalase enzyme. Out of selected 6 strains, strain of *Zygosaccharomyces rouxii* has given false positive results at all H₂O₂ concentrations used during plate assay method.

| RT   | Response | Ar/Ht | RFact | ECL | Peak Name | Percent (%) |
|------|----------|-------|-------|-----|-----------|-------------|
| 1.000| 24951    | 0.100 | 1.000 | 1.000| Straight  | 21.84       |
| 2.000| 3865     | 0.100 | 1.000 | 2.000| Branched  | 25.50       |
| 3.000| 5564     | 0.100 | 1.000 | 3.000| Hydroxy   | 1.62        |
| 5.000| 2485     | 0.100 | 1.000 | 5.000| MUFA      | 12.68       |
| 9.000| 168      | 0.100 | 1.000 | 9.000| 18:1w9c  | 38.36       |
| 12.000| 353    | 0.100 | ———— | 12.000| Other     | ————        |
Saponification value

Saponification value for *Zygrosaccharomyces rouxii* has calculated 1122. Saponification value is one of the chemical properties of biodiesel which contribute to fatty acid profile. Saponification value indicates amount of triacylglycerol present in total lipid. Since, the proportion of neutral lipid is the major component in total lipid content of *Zygrosaccharomyces rouxii* capable of yielding high amount of lipid biomass. The basic mechanism of lipid accumulation in microorganisms has been well studied. It was observed that when the culture medium lacks the nitrogen source, the isocitric dehydrogenase (ICDH) get
suppressed, therefore the tricarboxylic acid (TCA) was blocked. Extra carbon source was transformed to triglyceride (TAG) by a series of enzymes like the citric acid lytic enzyme, the malic acid enzyme, the fatty acid synzyme, thus completed the fat accumulation. When the culture medium contains sugar but low nitrogen, lipid accumulates in order to isolate oleaginous
microorganism, Sudan Black B staining is usually used to determine lipid content. However, this method roughly indicates the presence of microbial lipids.

**Conclusions**

A methodical study to search the soil sources for eukaryotes, producing lipid was performed. The study involved isolation and screening of PUFA (poly unsaturated fatty acid) producing oleaginous yeast, identification of the yeast with fatty acid profile by FAME-GC, physico-chemical parameters, optimization and saponification value from dairy waste soil sample. Primary screening of sample resulted in 22 isolates. Among them 6 isolates were found to generate lipid. Based on lipid productivity, the potent isolate CNK1 was selected and identified as *Zygrosaccharomyces rouxii* by FAME analysis. The lipid yield of *Zygrosaccharomyces rouxii* was found 0.098gm/100ml in optimized media and at optimized condition. *Zygrosaccharomyces rouxii* has shown saponification value 1122. In addition to that *Zygrosaccharomyces rouxii* is good PUFA producing and with high percent oleic acid side chain which are the best characteristics of oleaginous yeast used for biodiesel production. The above results are promising but still there is a need for protein sequencing, gene identification and other bioinformatics parameters study for further establishment of value of the yeast isolate *Zygrosaccharomyces rouxii*.

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