Production of Red Pigment from Fungal Isolate DMMS-1

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ABSTRACT

Colors have a very noteworthy role in one’s life; Colours have been used for therapeutic tenacity since ages. They play a significant role in setting up one’s mood, emotions, feelings and state of mind. Colour is even a medium of communication. Since colours mean so much in one’s life, they have a wide variety of applications; they are used in food industries as colorants, in textile industries as dyes, as antioxidants in pharmaceuticals and cosmetic industries etc. (Kumar et al., 2015). Man puts color on everything he creates in order to make it more eye catching. One wants to be surrounded by a colorful environment. The market is full of industrialized goods, the increased market demand of color has put a lot of pressure on the industries to synthesize high amount of color in order to satisfy its insatiable demand, to do so the industries have been producing the color by the chemical means. Most of the chemical used are of very poor quality. There have been cases reported the chemical dyes causing dermatitis, respiratory disorders, irritation in the upper respiratory tract and mucosal membrane (Bansal and Yadav 2016;
Ozkurt et al., 2012). Some researchers have been successful in proving the carcinogenic nature of these chemicals (Azo dyes to be specific) (Khandare and Govindwar 2015). Though one wants to be surrounded by an environment which is colourful but healthy as well. Due to their compatibility with the livings and the environment the natural pigments are gaining interest as an alternative to these chemical dyes globally. In natural pigments microbial pigments are the most efficient to use as the time and capital investment in case of plant origin pigment is very high and the efficiency is very poor, where as in case of animal origin pigments it’s not only the cost and poor yield but the ethical issues too. Besides this nature consists a plethora of microbes producing pigment which not only imparts color but many additional advantages like anti-proliferative, anti-biotic etc. At one side where the chemical dyes have potential to cause cancer on the other the microbial pigments possess anti-cancerous traits (Heer and Sharma, 2017). Here we have worked on the isolation and purification of a fungal isolate DMMS-1 and the standardization of the conditions for maximum pigment production.

**Materials and Methods**

**Isolation**

A soil sample from the agricultural fields of DAV University campus was taken diluted and was spread on N.A plates incubated at 37°C. A mixed culture was obtained in which the fungal isolate DMMS-1 was found to secrete a red coloured pigment in its periphery.

**Purification**

The fungal isolate was purified on the potato dextrose agar (PDA) and malt extract agar (MEA) medium by streaking. MEA was found to favour the pigment production so MEA was selected for the future use.

**Preparation of seed culture**

For preparation of seed culture an MEA plate streaked with fungal isolate was incubated at 37°C for 3 days. The pH of the medium was adjusted to be 5.5. After 3 days a full plate growth was obtained and that plate served as the seed culture for the liquid culture.

**Production of pigment under liquid culture condition**

Plugs from the 3-day old seed culture were cut with the help of a borer 5mm in diameter. The plugs were then inoculated to the MEA broth and the flasks with inoculated MEA broth were incubated at 37°C for an incubation period of 10 days. After 10 days of incubation a significant amount of pigment was obtained in the flasks. Since the pigment produced by the fungal isolate DMMS-1 is water soluble in nature, it was directly obtained in the broth.

**Pigment Filtration and analysis**

The layer of the fungal biomass formed at the top of the broth containing pigment was filtered using normal filter paper. The absorbance maxima for the obtained pigment were found to be 535 nm and hence the pigment obtained was analysed spectrophotometrically at a wavelength of 535 nm for further analysis. Un-inoculated MEA broth was used as the control/blank.

**Standardization of conditions**

The effect of conditions like inoculum size, pH of the medium, carbon source, nitrogen source, temperature, shaking vs non-shaking conditions were assessed by changing one parameter at a time.
Results and Discussion

A soil sample from the agricultural fields of DAV University campus was taken diluted and was spread on N.A plates incubated at 37°C. A mixed culture was obtained in which the fungal isolate (later named as DMMS-1) was found to secrete a red coloured pigment in its periphery. The fungal isolate was purified on the potato dextrose agar (PDA) and malt extract agar (MEA) medium by streaking. MEA was found to favour the pigment production so MEA was selected for the future use as a pronounced pigment secretion was observed (Fig. 1). The isolate was further cultivated under liquid culture conditions and the λ max of pigment secreted in the broth was determined spectrophotometrically (Fig. 2). The absorbance maxima for the pigment was found to be 535 nm and hence the pigment obtained was analysed spectrophotometrically at a wavelength of 535 nm for further analysis.

The effects of various cultural parameters on the pigment production by the fungal isolate were assessed. Flasks containing MEA broth were inoculated with different number of plugs (5mm in diameter) of seed culture. After the incubation of 10 days flasks with the inoculum size of 3 plugs was found to have synthesized the maximum pigment as compared to the flasks with inoculum size of 1, 2 and 4 plugs (Fig. 3). Similar results were reported by Bhat and Marar (2015) where increasing inoculum percentage first lead to increase and then constant decrease in the pigment production by Salinicoccus sp M KJ99797. Also, Studies conducted by Babitha et al., (2007) also suggested that the increase in inoculum size may increase the biomass but decreases the pigment production.

For assessing the impact of pH, flasks containing MEA broth with pH ranging from 3.5- 8.5 were inoculated with 3 plugs of the seed culture and were incubated for 10 days at 37°C resulting in the flask with pH 7.5 yielding the maximum pigment production (Fig. 4). Similar results were also observed by Babitha et al., (2007) and reported the highest pigment yield by Monascus perpureus was at a pH range of 4.5 to 7.5. Also, Hernández-Rivera, et al., (2008) found that the pH at which the highest production of red pigment produced by Monascus cultures was between 7.0 – 7.5.

For analyzing the impact of temperature on the pigment production, MEA flasks (pH 7.5 & 3 plugs) were incubated for 10 days at 28°C, 37°C and 45°C. Maximum yield of pigment was obtained in the flasks kept at 37°C whereas almost negligible growth in flasks incubated at 45°C. An intermediate amount of pigment was obtained in the flasks kept at 25°C (Fig. 5). While analyzing pigment produced by Penicillium purpurogenum, Patil et al., (2015) found that the temperature of 27°C is the temperature at which the highest pigment production was observed. Ahn et al., (2006) observed that the pigment produced by Monascus cultures at 30°C is ten times lesser than the pigment produced by it at 25°C.

In order to assess the effect of shaking on pigment production of the isolate, two sets of flasks containing MEA (pH 7.5) inoculum size of 3 plugs were incubated at 37°C for 10 days. Set 1 was kept in the non-shaking incubation whereas set 2 was kept under shaking incubation at 110 rpm. It was found that the amount of pigment produced in the flasks kept under shaking condition (110 rpm) was double than that of produced in the non-shaking condition (Fig. 6). Similar results were reported by Usman et al., (2018) where the orange pigment produced by Salinococcus roseus under shaking incubation was approximately three times the pigment...
produced under non-shaking incubation. The nutrients also play a very significant effect on the metabolic activities of all the living organisms, hence the effect of carbon and nitrogen on pigment production was assessed.

Effect of different carbon sources was analysed by replacing the Malt Extract with an equal amount of lactose, fructose and dextrose in the medium/broth.

**Fig.1** (a) Front view of Petri plate with Fungal isolate DMMS-1 growth, (b) Rear view of the Petri plate with pigment secreted by the fungal isolate

**Fig.2** Uninoculated ME Broth used as control (Left) and pigment obtained from the broth inoculated with the fungal isolate DMMS-1 (Right)

**Fig.3** Effect of inoculum size on the pigment production by fungal isolate DMMS-1
**Fig.4** Effect of pH on the pigment production by fungal isolate DMMS-1

![Graph showing the effect of pH on pigment production](image)

**Fig.5** Effect of temperature on the pigment production by fungal isolate DMMS-1

![Graph showing the effect of temperature on pigment production](image)

**Fig.6** Effect of Shaking/non-shaking condition on the pigment production by fungal isolate DMMS-1

![Graph showing the effect of shaking/non-shaking](image)
Fig. 7 Effect of carbon source on the pigment production by fungal isolate DMMS-1

![Graph showing pigment production by different carbon sources]

Carbon Source

Fig. 8 Effect of Nitrogen source on pigment production by fungal isolate DMMS-1

![Graph showing pigment production by different nitrogen sources]

pH of the media was adjusted to 7.5 and flasks were inoculated with 3 plugs each. The order of increasing pigment production was found to be: Lactose < fructose < dextrose < malt extract. Here malt extract supported the maximum pigment production whereas the pigment production in lactose and fructose were disappointingly low (Fig. 7). Similar results were found by da Costa Souza et al., (2016) they found that potato dextrose and Malt extract favoured increased pigment production. Also Subhsree et al., (2011) found fructose as the best suited carbon source for the pigment production from Monascus perpureus. For nitrogen source, flasks containing MEA broth with 3 plugs of seed culture and pH 7.5 were supplemented with different nitrogen sources like
Ammonium sulphate, Ammonium Molybdate, beef extract, peptone and yeast extract. The supplementation of nitrogen to the medium rather led to the decrease in the pigment production. Out of all maximum pigment was obtained in the flasks containing beef extract but was less than the pigment amount obtained without any nitrogen supplementation (Fig. 8). Chen and John (1993) observed that the Monascus perpureus growth along with the pigment production is affected by the nature of the nitrogen, where peptone and ammonium gave higher growth and pigment production than nitrate. Also Subhsree et al., (2011) reported Yeast Extract as the nitrogen source under which the highest pigment production by Monascus perpureus was achieved.

In conclusion, the present investigation was carried out with the aim of characterization of the fungal isolate DMMS-1, with respect to red pigment production. The isolate shows promising results in the preliminary investigations, since it secreted a red colored pigment into the culture broth. Thus, this pigment could be easily harvested without breaking of the fungal biomass. It was observed that the temperature of 37°C, inoculum size of 3 plugs (5 mm diameter), malt extract as carbon source, pH of 7.5 and the shaking incubation (110 rpm) resulted in enhancement of pigment production. So, more research is required to identify the fungus and analyze the characteristics of the red colored pigment in order to ascertain its possible commercial application in various industries.

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