Effects of Cultural Conditions in Enhancing the Production of Anti-MRSA Activity of *Lasiodiplodia pseudotheobromae* IBRL OS-64, an Endophytic Fungus Isolated from Leaf of *Ocimum sanctum* L. in Submerged Fermentation System

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Abstract

The effects of cultural conditions on the production of anti-MRSA activity of the endophytic fungal isolate, *Lasiodiplodia pseudotheobromae* IBRL OS-64 isolated from the leaf of a local medicinal herb, *Ocimum sanctum* Linn. was studied. The highest anti-MRSA activity of 50.0±0.1 U/mL and 1.46±0.1 g/L of fungal growth were obtained after incorporating all the improved cultural conditions which consisted of yeast extract sucrose broth supplemented with host plant extract, with initial medium pH of 6, 2 mycelial plugs of 4 days old seed culture, cultivation temperature of 30°C and cultivated for 16 days under static conditions without the presence of light. After improvement of cultural conditions, a significant increment of anti-MRSA activity of about 57.57% (2.4 folds) and a slight increment of fungal growth of about 8.15% (1.1 folds) were obtained compared to the cultural condition before improvement. Indeed the improvement of cultural conditions greatly affected the anti-MRSA activity and growth production by *L. pseudotheobromae* IBRL OS-64 isolate.

Keywords: Endophytic fungus, *Lasiodiplodia pseudotheobromae* IBRL OS-64, *Ocimum sanctum* Linn, anti-MRSA activity, submerged fermentation.

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INTRODUCTION

Endophytic fungi are ubiquitous and largely unexplored sources of novel bioactive compounds with structurally diverse secondary metabolites. They possess highly valuable bioactive compounds to human such as antibiotics and can be exploited as huge repertoire of potent biological activities (Mishra et al. 2012). Generally the secondary metabolites produced by the endophytic fungi are biosynthetically derived from primary metabolites via genetically controlled method and enzymatic catalysed reaction that lead to formation of complex compounds (natural product). The biosynthesis of the microbial metabolites is tightly control by regulatory mechanisms to avoid overproduction. These regulatory mechanisms often limit the yield of metabolites production especially through submerged fermentation system (SmF). SmF is a fermentation process that uses soluble substrate and growth medium by inoculating with microbes in shake flask or bioreactor systems. All the needed nutrients such as nitrogen and carbon sources are supplied in to this system to support microbial growth. This fermentation processes normally are done in shake flask or bioreactor whereby the pH, temperature, agitation speed, mixing frequency are controlled (Favella-Torres et al., 1998; Ruiz-Sanchez et al., 2010). According to Vidyalakshmi et al., (2009), submerged culture has their own advantages such as easy to sterilize and controlling process. To circumvent this problem, an attempt can made to optimize the cultural and physiological conditions in order to enhance the production of secondary metabolites as well as its biomass.

Many reports have shown the effectiveness of the endophytic fungal extracts against MRSA. We have reported the potential of Lasiodiplodia pseudotheobromae IBRL OS-64, an endophytic fungus isolated from leaf of Ocimum sanctum Linn which possess anti-MRSA activity (Taufiq and Darah, 2018a; 2018b). The ethyl acetate extract of the fungus has shown significance effects on the MRSA cells and proven to have bactericidal effect. Thus, the optimization of cultural conditions such as pre-culture seed, initial medium pH, incubation temperature, inoculum size, and agitation speed were performed in order to maximize the anti-MRSA activity and fungal growth of the isolate. The study is very important and significance since this is the first report of L. pseudotheobromae isolated from the leaf of O. sanctum Linn. and many of its characteristics are needed to be established.

MATERIALS AND METHODS

Endophytic fungal culture and maintenance

The endophytic fungus, L. pseudotheobromae IBRL OS-64 used in this study was provided by the Industrial Biotechnology Research Laboratory (IBRL), School of Biological Sciences, Universiti Sains Malaysia, Penang, Malaysia. It was isolated from the old leaf of O. sanctum and was cultured on potato dextrose agar (PDA) supplemented with powdered host plant powder (2 g/L) and incubated at 30°C for 9 days before keeping it at 4°C prior to use. The isolate was subcultured regularly on sterile fresh PDA medium once a month to ensure its purity and viability.

Test microorganism

The culture of Methicillin-resistant Staphylococcus aureus ATCC 33591 which was also provided by the Industrial Biotechnology Research Laboratory, School of Biological Sciences, Universiti Sains Malaysia, Penang, Malaysia was grown on nutrient agar (NA) and incubated at 37°C for 24 hours. The inoculum was prepared by picking-up five single isolated colonies from 24 hours old culture and transferred into 5 mL of 0.85% sterile physiological saline (w/v). The turbidity of the bacterial suspension was adjusted to match 0.5 McFarland standards (approximately 1 x 10^8 CFU/mL). The isolate was subcultured on fresh NA medium once a month to ensure its purity and viability.

Initial profiling of anti-MRSA activity

The anti-MRSA activity and fungal growth profile before optimization were carried out to determine the cultivation day with highest anti-MRSA activity. One plug of the 6 days old endophytic fungus was inoculated into 250 mL Erlenmeyer flasks containing 100 mL of yeast extract sucrose (YES) broth supplemented with powdered host plant powder (Taufiq and Darah, 2018a), initial medium pH was adjusted to pH 6.0. The flasks were then incubated at 30°C with 120 rpm agitation speed in dark condition. The anti-
MRSA activity as well as the fungal growth was determined every two days interval for 30 days. All experiments were carried out in triplicates.

### Optimization of different cultural conditions on the production of anti-MRSA activity and fungal growth

The optimization of physical cultural conditions of the submerged fermentation system for enhancing the production of anti-MRSA activity and fungal growth was carried out on pre-culture seed. The fragment of endophytic fungal culture was inoculated on PDA supplemented with powdered host plant (2 g/L) and incubated at 30°C. At 2, 4, 6, 8, 10 and 12 days of cultivation, the periphery of the colony was punched using a 6.0 mm diameter of a sterile cork borer to get fungal plugs which then be used as inoculum.

The other physical parameters evaluated were the supplementation of growth medium with and without the host plant powder or extract), light intensity (presence or absence), incubation temperature (25°C, 30°C, 35°C, 40°C and 45°C), pre-culture seed (2, 4, 6, 8, 10, 12 days), number of agar plug (1, 2, 3, 4 and 5), agitation speed (static, 50, 100, 120 and 150 rpm) and initial medium pH of the culture medium (4, 5, 6, 7 and 8). All experiments were carried out in triplicates.

### Extraction process

The fermentative broth was extracted using method described by Tong et al., (2011). The fungal biomass was separated from the broth using a sterile Whatman filter paper no.1 and the filtrate was then kept in a sterile glass container for subsequent study.

### Anti-MRSA determination assay

The method described previously by Lorian (1991) with slight modifications was followed to determine the anti-MRSA activity quantitatively. An amount of 2.0 mL of crude extract (fermentative broth) was introduced into a clean aluminium screw cap universal bottle (Favorit) containing 7.9 mL of nutrient broth (NB). Then, 0.1 mL of 24 hour old culture of MRSA (in the form of bacterial suspension) was added to the nutrient broth (NB). For blank, 2.0 mL of fermentative broth was introduced into an aluminium screw cap universal bottle (Favorit) containing 8.0 mL of NB. For control, 0.1 mL of 24 hour old culture of MRSA (in the form of bacterial suspension) was added into the aluminium screw cap universal bottle (Favorit) containing 9.9 mL of NB without crude extract. The samples were then incubated at 37°C for 18 hours and the OD was measured at the wavelength of 560 nm. The antibacterial activity of the extract was defined as one unit (U) of the antibacterial activity which resulted in the reduction or inhibition of the growth of MRSA (Darah et al., 2014). The experiment was done in triplicate.

### Determination of fungal growth

The fungal growth was determined according to method described by Ibrahim et al., (2015) and the fungal cell dry weight was then expressed as g/L.

### Statistical Analysis

One-way analysis of variance (ANOVA) and Duncan multiple range test (DMRT) with PASW (SPSS) Statistics version 12.0 were used to analyse the significant difference of the mean of experimental data. A 5% confidence level or $\alpha = 0.05$ were used to test all the experimental data.

### RESULTS

#### Time course profiles of the anti-MRSA activity and fungal growth before improvement of physical parameters

The time course before the improvement of cultural conditions (yeast extract sucrose [YES] broth nourished with host plant powder, pH 6.0, one mycelial plug, incubation temperature of 37°C and 120 rpm agitation speed) was studied for 30 days and the results are shown in Fig. 1. The results showed a gradual increment in the anti-MRSA activity which started from day 7 of cultivation and reached its maximal activity on day 22 of cultivation with 21.22±0.9 U/mL. The anti-MRSA activity decreased slowly thereafter and achieved about 14.04±0.6 U/mL on day 30 of cultivation. The fungal growth on the other hand started earlier on day 2 of cultivation and reached its maximal growth on day 18 of cultivation with about 1.35±0.1 g/L. However, its growth was stagnant thereafter and achieved about 1.32±0.1 g/L on day 30 of cultivation. The significance of the study was tested using Duncan test, $p < 0.05$. Thus, 22 days of cultivation period was optimal for anti-MRSA activity and retained in the next experiment.
Improvement of cultural conditions for maximal anti-MRSA production

Effects of pre-culture seed

Six different pre-culture seeds, i.e., 2, 4, 6, 8, 10 and 12 days old were tested throughout this experiment and the results are shown in Fig. 2. The 4 days old pre-culture seed produced the highest anti-MRSA activity with the value of 30.58±1.1 U/mL and the fungal growth was 1.39±0.1 g/L. Meanwhile, 10 and 12 days old pre-culture seed produced the least anti-MRSA activity with the value of 7.77±0.3 U/mL and 8.01±0.3 U/mL, respectively. The anti-MRSA activity production of 2, 6 and 8 days old pre-culture seed were 22.06±0.5 U/mL, 15.20±0.9 U/mL and 10.00±0.2 U/mL, respectively. The significance of the study was tested using Duncan test, p < 0.05. The statistical analysis revealed that 4 days old pre-culture seed was significantly promoted the highest anti-MRSA activity and was then used as a pre-culture seed for the subsequent experiment.
Effects of culture medium incorporated with or without host plant powder or extract and light intensity

The effects of culture medium incorporated with host plant, and light intensity on anti-MRSA activity was investigated and the results are shown in Fig. 3. The highest anti-MRSA activity of 38.17±0.5 U/mL was produced in the medium nourished with host plant extract and incubated in dark condition (without the presence of light). Meanwhile, the culture medium without the addition of host plant and cultivated in the presence of light produced the lowest anti-MRSA activity with the value of 22.63±1.5 U/mL. The current findings revealed that the cultivation medium supplemented with host plant extract were significantly enhanced the anti-MRSA activity (37.07±0.6 U/mL) compared to the cultivation supplemented with host plant powder (28.41±0.3 U/mL). On the other hand, the cultivation medium incorporated with host plant either extracts or powder (32.74±0.9 U/mL) significantly increased the antibacterial activity compared to the medium without the addition of plant materials (25.48±0.6 U/mL). Results indicated that the endophytic fungus need its host plant material in order to enhance the anti-MRSA activity. For the effect of light intensity, the culture medium cultivated in the dark condition produced the highest anti-MRSA activity (32.47±0.9 U/mL) compared to the one that cultivated in the presence of light (28.17±0.6 U/mL). However the anti-MRSA activity production was not growth dependant. The significance of the study was tested using Duncan test, p < 0.05. The statistical analysis revealed that the culture medium supplemented with host plant extract in dark condition showed significantly the highest anti-MRSA activity. Therefore, this condition was selected to be applied in the next experiment.

![Fig. 3. Effects of growth medium and light intensity on the production of anti-MRSA activity and fungal growth by L. pseudotheobromae IBRL OS-64. Different letters indicate significant differences between means (Duncan, p<0.05). Standard deviation values of triplicate analyses are shown by error bars. [notes: D-dark condition, WEP-without host plant extract/powder, E-host plant extract, P-host plant powder, L-in the presence of light]](image)

Effects of initial medium pH

The effects of initial culture medium pH on anti-MRSA activity produced by L. pseudotheobromae IBRL OS-64 in a shake flask system were investigated and the results are shown in Fig. 4. The maximal production of anti-MRSA activity of about 37.75±0.7 U/mL was found in pH 6 (Duncan, p < 0.05). Again the anti-MRSA activity production was not growth dependant. Meanwhile, the initial medium pH beyond or lower than the optimal level produced lower amount of anti-MRSA activity. Thus, the initial medium pH of 6 was used in the subsequent studies.

Effects of cultivation temperature

The effects of various cultivation temperatures were carried out and the results are shown in Fig. 5. With the increase in cultivation temperatures, the production of anti-MRSA activity by L. pseudotheobromae IBRL OS-64 increased and reached the maximal production
of 38.56±0.7 U/mL at the temperature of 30°C. However, further increased in the cultivation temperature (35°C) showed an abrupt declining in anti-MRSA activity. Furthermore, it is noteworthy that further increase in cultivation temperature beyond 40°C inhibited the fungal growth as well as the production of anti-MRSA activity. Besides that, cultivation temperatures below than 30°C (optimal cultivation temperature) produced lower anti-MRSA activity. Since cultivation temperature of 30°C produced the highest anti-MRSA activity (Duncan, p < 0.05), thus, it was used in the subsequent experiment.

Fig. 4. Effects of initial medium pH on the production of anti-MRSA activity and fungal growth by L. pseudotheobromae IBRL OS-64. Different letters indicate significant differences between means (Duncan, p<0.05). Standard deviation values of triplicate analyses are shown by error bars.

Fig. 5. Effects of temperature on the production of anti-MRSA activity and fungal growth by L. pseudotheobromae IBRL OS-64. Different letters indicate significant differences between means (Duncan, p<0.05). Standard deviation values of triplicate analyses are shown by error bars.

**Effects of the number of mycelial plug**

The effects of number of the mycelial plug on the production of anti-MRSA activity were studied and the results are shown in Fig. 6. The highest anti-MRSA activity was produced by the 2 mycelial plugs of the age of 4 days old with the activity of 41.51±0.8 U/mL and the fungal growth of 1.42±0.1 g/L (Duncan, p < 0.05). However, further increased in numbers of mycelial plug (3, 4 and 5) gradually decreased the production of
anti-MRSA activity as well as the fungal growth. Thus, the mycelial plug of 2 gave highest anti-MRSA activity and was then used in the next experiment.

**Effects of agitation speed**

Agitation speed is an important condition for the fungal growth and production of anti-MRSA activity in submerged fermentation. Five agitation speeds were studied and the results are tabulated in Fig. 7. The findings revealed that the static condition (0 rpm) produced the highest anti-MRSA activity of about 47.56±0.6 U/mL and the fungal growth of 1.41±0.1 g/L (Duncan, p < 0.05). However, the increased in agitation speeds from 50 to 150 rpm showed a gradual decreased in anti-MRSA activity (42.58±1.3 U/mL to 20.32±1.2 U/mL, respectively). Since static condition showed the highest anti-MRSA activity, it was selected and retained in the next experiment.

**Fig. 6.** Effects of the number of mycelial plugs on the production of anti-MRSA activity and fungal growth by *L. pseudotheobromae* IBRL OS-64. Different letters indicate significant differences between means (Duncan, p<0.05). Standard deviation values of triplicate analyses are shown by error bars.

**Fig. 7.** Effects of agitation speeds on the production of anti-MRSA activity and fungal growth by *L. pseudotheobromae* IBRL OS-64. Different letters indicate significant differences between means (Duncan, p<0.05). Standard deviation values of triplicate analyses are shown by error bars.

**Time course profiles after the improvement of physical parameters of submerged fermentation in a shake flask system for the production of anti-MRSA**

A time course profiles for the production of anti-MRSA activity after the improvement of physical parameters were carried out and the results are shown in Fig. 8. The profiling was
performed under improved culture conditions (culture medium nourished with host plant extract, initial medium pH of 6, cultivation temperature of 30°C, without the presence of light, pre-culture seed of 4 days, inoculum size of 2 mycelial plugs and under static condition) for 30 days in submerged fermentation using a shake flask system. Overall, the results showed a gradual increment in the production of anti-MRSA activity and fungal growth with the cultivation time. The anti-MRSA activity was detected as early as day 5 of cultivation and increased with the increment of cultivation time and reached its maximal production of 50.00±0.1 U/mL on the day 16 of cultivation (Duncan, p<0.05). The anti-MRSA activities were stagnant on days 18 and 20 of cultivation. However, further increased in cultivation time beyond day 20th showed a gradual decreased in anti-MRSA activity and, on day 30th, it reached the lowest activity of about 25.77±1.5 U/mL. In term of fungal growth, the growth started immediately after the cultivation and achieved its maximal growth of about 1.46±0.1 g/L on day 16 of cultivation and then grow slowly until day 22 before declining thereafter. Thus, the findings of present study revealed that the improvement of physical parameters provided substantial effects on the production of anti-MRSA activity by the endophytic fungus, *L. pseudotheobromae* IBRL OS-64.

Table 1 summarizes the production of anti-MRSA activity as well as fungal growth before and after incorporating all the improved cultural conditions. Results revealed that the anti-MRSA activity and fungal growth increased up to 57.57% (2.4 folds) and 8.15% (1.1 folds) under improved conditions, respectively. Throughout this study, it was observed that the pre-culture seed, additional of host plant extract, number of plugs and agitation speeds played major roles in improving and enhancing fungal metabolite production. Besides that, the results also exhibited that the improvement of cultural conditions were not only enhance the production of anti-MRSA activity, but also shorten the cultivation time. Therefore, the shorter cultivation period would definitely reduce the cost for production at a commercial scale.

**DISCUSSION**

As far as culture conditions are concerned, there is usually a dilemma between achieving maximal fungal growth rates and maximal antibiotic yields because conditions that allow fast cell growth could be unfavourable to metabolite production (Miao *et al.*, 2006). Furthermore in submerged fermentation there is always production of metabolites which is not dependent to the fungal growth. The yield of metabolites can sometimes be substantially increased by the improvement of physical parameters such as...
The maturity of the pre-culture seed was studied to determine its effect on fungal growth and production of their secondary metabolites. The current findings revealed that younger seed culture produced the highest anti-MRSA activity as well as fungal growth. Meanwhile, older pre-culture seed resulted in decrement of fungal growth as well as their secondary metabolite production. This condition could be due to the old culture that was in inactive condition which needs an extent cultivation period to reproduce and re-propagate new mycelia, whereas the young active culture just continues their growth phase in a new fresh medium. Brookman et al., (2000), did a comparative evaluation on fungal growth and the results revealed that the active growth of older cultures as inoculum was delayed compared to young cultures. Similarly, Villiers (1973), reported that older seed culture might delay the germination due to repairs and replacements mechanisms in the cell that might affected organelles and enzymes. Besides that, Herker et al., (2004) reported that apoptotic regulation occurred in older microbial cells that releasing a substance into medium to stimulate the survival of new younger and fitter cells. This resulted in delayed biomass production.

Endophytes are microorganisms that symbiotically colonized within plant tissues for all or a part of their life cycle without causing any harmful effect to their host (Hirsch and Braun, 1992; Wilson, 1995). Fungal endophytes benefit and offer several advantages to their host by preventing them from desiccation, promoting plant growth hormone, increasing nutrients acquisition, protecting from insect, pathogenic fungi and environmental harsh conditions (Dutta et al., 2014; Nair and Padmavathy, 2014). Meanwhile, the host plant would provide the endophytes with shelters, space for their colonization and also nutrients acquired by them (Saikkonen et al., 2004; Schardl et al., 2004). It was observed that culture medium supplemented with host plant extract enhanced the production of fungal metabolites and their growth. The current findings were in agreement with Yenn et al., (2012) who reported the anti-candidal activity was produced by Phomopsis sp. ED32 when the culture medium was supplemented with host plant extract. Host plant extract is believed to supply the endophytes with crucial needed nutrients or growth compounds that could be used in biosynthesis of cellular components resulting in increment of fungal secondary metabolite production (Tong et al., 2011; Padhi and Tayung, 2013).

The effects of light on the secretion of fungal bioactive metabolites and their growth

Table 1. Summary of the anti-MRSA activity and fungal growth production by L. pseudotheobromae IBRL OS-64 after improvement of physical parameters in a submerged fermentation using a shake flask system

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Profiling before improvement</th>
<th>Profiling after improvement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-culture seed</td>
<td>6 days old</td>
<td>4 days old</td>
</tr>
<tr>
<td>Effect of host plant light intensity</td>
<td>host plant powder extract</td>
<td>host plant powder extract</td>
</tr>
<tr>
<td>Light intensity</td>
<td>dark</td>
<td>dark</td>
</tr>
<tr>
<td>Temperature</td>
<td>30°C</td>
<td>30°C</td>
</tr>
<tr>
<td>Inoculum size (no. of plugs)</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Agitation speed (rpm)</td>
<td>120</td>
<td>0 (static)</td>
</tr>
<tr>
<td>Initial pH</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Optimum activity (U/mL)</td>
<td>21.22±0.9</td>
<td>50.00±0.1</td>
</tr>
<tr>
<td>Increment (%)</td>
<td>-</td>
<td>57.57</td>
</tr>
<tr>
<td>Fungal growth (g/L)</td>
<td>1.35±0.1</td>
<td>1.46±0.1</td>
</tr>
<tr>
<td>Increment (%)</td>
<td>-</td>
<td>8.15</td>
</tr>
</tbody>
</table>

incubation temperature, initial pH medium value, size of inoculum, agitation speed and light (Calvo et al. 2002; Llorens et al. 2004).
were investigated and the result showed that the presence of light slightly decreased the anti-MRSA activity of *L. pseudotheobromae* IBRL OS-64. Schmidt-Heydt *et al.*, (2011) reported the same observation in which the presence of light reduced the production of ochratoxin. Similarly, Soliman and Raizada, (2018) claimed that the presence of light exhibited inhibitory effect on the production of fungal secondary metabolites. In the Taxol-producing experiment, they observed that the complete loss of taxol production occurred when the endophytic fungus SSM001 exposed to light. Kim *et al.*, (2013a), reported the inhibitory effect of the light might be due to unavailability of LaeA nuclear protein which was a regulator protein that involved in biosynthesis of fungal metabolites and thus leads to the inactivation of secondary metabolites production. Furthermore, Kim *et al.*, (2013b) found that white light induced the carotenoid biosynthesis by *Fusarium graminearum* strains. They suggested this phenomenon might be due to the role of fgWC-1 and fgWC-2 that act as negative and positive regulators, respectively that monitored the light requirement of *F. graminearum* in its biological process. Furthermore, in filamentous fungus, the light-dependent gene was expressed with the light exposure and thus, enhancing secondary metabolite production by activating the light-mediated control of the DNA expression (Wang *et al.*, 2014). However, Velmurugan *et al.*, (2010) stated that the fungi may react differently according to light wavelength. For instance, certain light wavelength might enhance extracellular pigment production and some are not.

The initial medium pH of growth medium is a crucial factor for fungal growth and production of their secondary metabolite since it affect the function of several enzymes in catabolic reaction and also influences on complex physiological phenomena such as cell morphology and their cell membrane permeability (Guimaraes *et al.*, 2004). In the present study, the slightly acidic growth medium (pH 6) was the best for the production of fungal secondary metabolite. The result was in agreement with previous study that reported pH 6 was the best for bioactive metabolite production of *Streptomyces coeruleorubidus* isolated from soil (Bundale *et al.*, 2015). The current findings also revealed that production of fungal secondary metabolite of *L. pseudotheobromae* IBRL OS-64 was decreased with the increment of pH and the similar observation was reported by Keller *et al.*, (1997). Drori *et al.*, (2003) reported the increasing in pH of the growth medium from 4 to 6 was an ambient pH transduction pathway which increases the production of secondary metabolites by *C. gloeosporioides*. Noaman *et al.*, (2004) suggested the pH of growth medium is closely related to permeability of microbial cell wall and membrane and therefore, affected in ions uptake and loss in the growth medium. According to Rizk *et al.*, (2007), the concentration of hydroxyl and hydrogen ion might have direct or indirect effects on the cell by varying dissociation level of substances in the medium. Thus, the change of pH in the medium was crucial for microbial enzyme activity especially for dissociation and solubility of intermediate products. Therefore, the extremely low and high pH of the growth medium is believed to disturb enzyme activity resulted in suppression of their secondary metabolite production.

Another cultural condition that has big effects on fungal secondary metabolite production is cultivation temperature (Rizk *et al.*, 2007). In the present study, cultivation temperature of 30°C produced the highest anti-MRSA activity by *L. pseudotheobromae* IBRL OS-64. The results were in agreement with Bhattacharyya and Jha, (2011) that reported the fungal growth and their bioactive metabolite production increased with the increment of cultivation temperature from 25°C to 30°C. The current finding also revealed that lower cultivation temperature resulted in slightly decrement of bioactive compounds and the temperature higher than 40°C killed the growth of *L. pseudotheobromae* IBRL OS-64. Gogoi *et al.*, (2008) stated that a fungal endophyte, *Hypocrea* spp. NSF-08 which was cultivated at temperature of 28°C produced the highest production of bioactive metabolite, and the fungus could not grow at temperature below 20°C and above 50°C. Agastian *et al.*, (2013), stated the lower temperature might cease the fungal metabolite activity whilst at higher temperature, they might be killed.

**Inoculum sizes are important in fermentation as well as secondary metabolites production by fungi.** Low inoculum size may influence the time extension for cell proliferation...
to utilize substrate and produce the desired product of secondary metabolites (Darah et al., 2015). Moreover, low inoculum sizes might lead to prolongation of the lag phase of fungal growth and resulted in slow growing rate and greater fungal viability (Baert et al., 2008; Garcia et al., 2010). In the present study, this problem could be observed at inoculum size of 1 mycelial plug, where lower anti-MRSA activity was produced. On the other hand, higher inoculum size than 1 mycelial plug increased anti-MRSA activity. Thus, an inoculum size of 2 mycelial plugs was found to be optimal for the production of maximal anti-MRSA activity. The anti-MRSA activity declined slightly with further increment in inoculum sizes. Similar observation also reported by Sidhu et al., (2017) in which two mycelial discs was sufficient to stimulate higher laccase production by *Scytalidium lignicola* under submerged fermentation. Higher inoculum sizes than the optimal may stimulate too much production of microbial biomass which lead to nutrient depletion for their secondary metabolite production (Kumar et al., 2010). According to Ahsan et al., (2017), moderate inoculum size has a significant positive effect on the antibiotic production and growth yield. Limited space and less oxygen supply might occur if higher sizes of inoculum used in cultivation. However, if lower sizes of inoculum applied, the excess materials in the culture may turn toxic and thus lead to unfit fermentation products.

Microbial activity of aerobic cultures is markedly affected by the oxygen supply to the system. There are varying functional aspects of aeration in submerged fermentation including the maintenance of oxygen supply, removing carbon dioxide from the system and heat transfer. Aeration provides good transfer of nutrients and product gases in the submerged fermentation system. Agitation speed and dissolved oxygen are important factors in determining the successful of microbial cultivation whereby the sufficient dissolved oxygen in growth medium is crucial in mass transfer characteristics of microbial cells (Ibrahim et al., 2015). The present study revealed that the static condition produced the highest anti-MRSA activity and the activity was decreased with the increment of agitation speeds. The result obtained was in agreement with previous studies that reported the highest production of bioactive compound was obtained under static condition (Bringmann et al., 2007; Tamminen et al., 2014). The favorable effect of air flow on the secondary metabolite production could be attributed to the enhancement of product formation by microorganism under-forced aeration. The cell differentiation might be occurred under insufficient dissolved oxygen which leads to more formation of aerial hyphae and conidiophores and thus, increases the accumulation of bioactive compounds (Scott and Eaton, 2008; Tang et al., 2015). This statement was supported by Pearce, (1997) who postulated that the fungal are able to store bioactive compounds in their mycelia. According to Ahsan et al., (2017), agitation speed influenced the dissolved oxygen temperatures and oxygen availability and therefore affecting the antibiotic production. Moreover, oxygen availability is one of the crucial factors for cell growth and production of their secondary metabolites during fermentation process (Wang et al., 2010; Dou et al., 2013). Besides that, Song et al., (2012) postulated that proper ventilation should be considered during fermentation process since the culture media containing organic and inorganic compounds that lead to the decrement in dissolved oxygen level. In contrast, Noraziah and Izyani (2012) reported the agitation could increase biomass formation by supplementing more dissolved oxygen. However, the fungal growth and their fungal mycelium morphology is dependent on several factors including the growth medium, hydrodynamic in the bioreactor, fungal strains, and method of initiation of culture such as mycelium dispersion (Lopez et al., 2005). Interestingly, fungal growth of the *L. pseudotheobromae* IBRL OS-64 was not affected either under stationary or agitated condition since the value was almost the same even though the anti-MRSA activity was varies. According to Barborkakova et al., (2012), the bioactive production is not dependent to the quantity of biomass. The agitated and static conditions of cultivation conditions might have their pros and cons, but the adaptation to the harsh environment is depends on the fungal strains and their response mechanisms to stress conditions. Even agitation is believed to provide adequate mixing especially for dissolved oxygen, heat and mass transfers, perhaps it also induce shear forces which resulted in morphological
changes and damage of the cell structures (Darah et al., 2011; Zhu et al., 2012).

Cultivation time is also a crucial parameter in determining the success of fermentation process to gain fungal secondary metabolites. In the present study, the anti-MRSA activity was detected after about a week of cultivation time. The late detection of anti-MRSA activity might be due to the production of anti-MRSA compound as fungal secondary metabolites. According to Bu’Lock (1961), secondary metabolites are secreted during the late stage of microbial growth, at a stationary phase. It was noted in this study that the extended cultivation time (22th day of cultivation time) produced maximal yield of anti-MRSA activity. The result was in agreement with Yamanaka et al., (2008) who reported the prolonged cultivation period could induce secondary metabolite production by Trametes versicolor. They also suggested that the cells are physiologically active at a long period of cultivation. Generally, the extended cultivation time would create unfavorable conditions which due to starvation of carbon or nitrogen, accumulation of self-toxic metabolites, translocation of nutrients, elevation of pH, increment of temperature and degeneration of fungal mycelium. These unfavorable conditions are believed to adversely affect the expression of microbial secondary metabolites (Griffin, 1995). According to Sejiny (1991), during the propagation of antibiotics producers, two phases were observed which were trophophase and idiophase. Trophophase is characterized by rapid growth of microbial cells producing biomass and during the idiophase, slow microbial growth was observed and maximal antibiotic production was obtained. In the present study, all the major physical parameters which were believed to affect the fermentation process for anti-MRSA production were investigated, and the results showed that the 16th day of cultivation time produced maximal production of anti-MRSA activity.

CONCLUSION

The present study suggests that fermentative broth of L. pseudotheobromae IBRL OS-64 exhibited a potent anti-MRSA activity through submerged fermentation in a shake flask system. The optimal cultivation conditions were significantly affected the production of anti-MRSA. Under improved conditions, the production of anti-MRSA activity was about 50±0.1 U/mL, which corresponded to 57.57% of increment compared to the yield before improvement of cultivation conditions.

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CONFLICTS INTEREST

The authors declare that there is no conflict of interest.

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AUTHOR’S CONTRIBUTION

MMJT performed all the experiments. ID supervised the entire work performed in her laboratory and was the principle investigator. ID and MMJT wrote, revised and approved the manuscript for publication.

DATA AVAILABILITY

All datasets obtained throughout this study are incorporated in the manuscript.

ETHICS STATEMENT

The authors declare that this article does not contain any studies with animals or human participants.

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