INTRODUCTION

Starting with the early reports of antibiotic resistance towards penicillin and sulphonamides just after the World War II, it then involved many other cases of antibiotic resistance occurred including the establishment of MRSA, a pathogenic strain of Staphylococcus aureus that resistant to methicillin [1]. It is one of the human pathogens that involved in nosocomial acquired infection and caused serious infectious diseases with high rate in morbidity and mortality [2]. Ocimum sanctum Linn is a medicinal herb that is known locally as basil or 'selasih hitam' in Malaysia can be found commonly planted in home gardens due to its traditional uses and medicinal values [3]. Traditionally, the plant is used to relieve common cold, bronchitis, cough and digestive problems [4]. Besides that, researchers have proven that various part of this plant exhibited antibacterial [5], antifungal [6], and antioxidant [7] and insecticides properties [8]. Microorganisms such as endophytes have been recognised as a potential source of novel bioactive compounds with promising antimicrobial activity [9]. Endophytic fungi can be defined as highly diverse microorganisms that live within plant tissues of their host but remain asymptomatic [10]. High diversity of endophytic fungi associated with medicinal herbs is observed and their distributions, as well as composition, are affected by tissues of their host but remain asymptomatic [10]. High diversity of endophytic fungi associated with medicinal herbs is observed and their distributions, as well as composition, are affected by tissues of their host but remain asymptomatic [10]. High diversity of endophytic fungi associated with medicinal herbs is observed and their distributions, as well as composition, are affected by tissues of their host but remain asymptomatic [10]. High diversity of endophytic fungi associated with medicinal herbs is observed and their distributions, as well as composition, are affected by tissues of their host but remain asymptomatic [10]. High diversity of endophytic fungi associated with medicinal herbs is observed and their distributions, as well as composition, are affected by tissues of their host but remain asymptomatic [10]. High diversity of endophytic fungi associated with medicinal herbs is observed and their distributions, as well as composition, are affected by tissues of their host but remain asymptomatic [10].
Cultivation and extraction

The inoculum was prepared by introducing two mycelial agar plugs approximately 1.0 cm in diameter and 4.0 mm thickness, which were excised from the periphery of 7 d old endophytic fungal cultures into 250 ml Erlenmeyer flasks containing 100 ml of YES broth medium. The cultures were cultivated at 30 °C with an agitation speed of 120 rpm for 20 d in the dark. After the incubation period, the fermented broth and fungal biomass were separated out using filter papers (Whatman, No. 1). The filtered broth was extracted thrice with an equal volume of ethyl acetate (1:1, v/v). The upper organic phase was collected and subsequently was concentrated to dryness under reduced pressure in a rotary evaporator to obtain the ethyl acetate crude extract paste. Due to the antibacterial activity exhibited by *O. sanctum*, a control was included by extracting the sterile medium following exactly the same procedure as that for the endophytic cultures.

Test microorganisms

The Methicillin-resistant *Staphylococcus aureus* ATCC 33591 culture was provided by the Industrial Biotechnology Research Laboratory, Universiti Sains Malaysia, Penang, Malaysia. The bacterial culture was grown on nutrient agar (NA) (Merck, Germany) and incubated at 37 °C. The inoculum suspension was prepared by picking up five single isolated colonies from 24 h old culture and transferred them into 5.0 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).

MIC and MBC determinations

The minimal inhibitory concentration (MIC) of the fungal ethyl acetate crude extract was determined by broth microdilution assay as described by Jorgensen and Ferraro [20] with some modifications. The broth microdilution was conducted in a sterile, 96-well, U-shaped microtiter plate (TPP, Switzerland). A single-fold dilution of the fungal crude extract was prepared in sterile Muller Hinton Broth (MHB) (Merck, Germany) medium and 100 µl of the extract was dispensed into each wells of a microtiter plate. On the other hand, 100 µl of bacterial inoculum at approximately 1 x 10^7 CFU/ml was added into each of the wells for a final volume of 200 µl and the final concentration of bacterial in each wells was 1 x 10^7 CFU/ml. Chloramphenicol (Sigma-Aldrich, USA) was used as a reference drug and a control with 5% methanol and bacterial inoculum was included. After 24 h of incubation period at 37 °C, 40 µl of 0.2 mg/ml p-iodonitrotetrazolium violet (INT) (Sigma-Aldrich, USA) dissolved in 99.5% ethanol solution was loaded to each wells as a growth indicator. The MIC value was determined and recorded as the lowest concentration of the ethyl acetate crude extract that capable to inhibit the visible growth of test bacteria after 24 h of incubation period [21].

Minimal bactericidal concentration (MBC) of the fungal ethyl acetate crude extract was subsequently determined upon reading of minimal inhibitory concentration (MIC) value. The viable cell was enumerated on Mueller Hinton agar (MHA) (Merck; Germany) by a standard viable plate count after 24 h of incubation period at 37 °C for overnight. The MBC was observed and recorded as the lowest concentration of fungal ethyl acetate crude extract that resulted in a reduction of 99.9% bacterial growth relative to the growth control.

Time-kill study

In the time-kill study, the ethyl acetate crude extract was tested at concentrations of 1/2MIC, MIC, and 2MIC. Prior to that, the minimal inhibitory concentration (MIC) value of MRSA was 125.0 µg/ml. A volume of 0.1 ml of bacterial suspension (approximately 1 x 10^8 CFU/ml) was transferred into a 25 ml Erlenmeyer flask containing 19.9 ml of MHB with different concentrations of extract yielded the initial bacterial suspension of approximately 5 x 10^7 CFU/ml. For comparison, the inoculum consisted of bacterial culture and 1% Dimethyl sulfoxide (DMSO; v/v) in MHB was set as a control. The cultures were subsequently incubated in a rotary orbital shaker (Lab-Companion, Korea) at 37 °C with agitation rate of 150 rpm for 48 h. A volume of 0.1 ml of the aliquot was withdrawn for viable cell count at every 4 h during the time intervals of 0 to 48 h. The samples were diluted and spread onto fresh nutrient agar plates followed by incubating at 37 °C for 24 h and the viable bacterial colonies were then counted. To determine the colony unit per millilitre (CFU/ml), only the plates with the number of colonies ranging from 30–300 were counted. A time-kill curve (log CFU/ml vs. time) was plotted for each extract concentrations and control. At the meantime, the growth reduction in which the time to achieve 50%, 90%, and 99.9% of bacterial cells reduction was calculated according to equation 1. The experiment was carried out in triplicates on separate occasions. The result was expressed as the mean ± SD. P values<0.05 were considered statistically significant.

\[
\text{Growth reduction} = \frac{V_t - V_c}{V_c} \times 100 \quad (1)
\]

Whereby *V_i* is the initial viable cell count and *V_v* is the viable cell count at time *t*.

**Scanning electron microscopy (SEM)**

For sample preparation, 50 µl of bacterial inoculum (approximately 1 x 10^8 CFU/ml) was transferred into a 25 ml Erlenmeyer flask containing 945 µl of MHB and incubated in an orbital shaker at 37 °C, 150 rpm for 18–20 h. A volume of 0.5 ml of ethyl acetate crude extract (2.5 mg/ml) was added into the bacterial culture after incubation period to yield a volume of 10 ml mixture with extract concentration of 0.25 mg/ml. At the meantime, 0.5 ml of 20% DMSO (v/v) was added into the bacterial culture as a replacement to the extract and served as control. The mixture was then incubated at 37 °C, 150 rpm for 36 h. The cultures were harvested at 0, 12, 24 and 36 h of the incubation period and proceed for SEM works as described previously by Borges [22]. The prepared samples were then viewed under SEM (Leica Cambridge, S-360, UK).

**Statistical analysis**

The data obtained in this study were analyzed by Student t-test for comparing the effect of ethyl acetate extract on test microorganism cells against control using SPSS Version 12.0. All tests were independently performed in triplicates. The results obtained were analyzed using a one-way ANOVA test and reported as the mean±SD. P values<0.05 were considered statistically significant.

**RESULTS**

**MIC and MBC determinations**

The susceptibility test result for ethyl acetate crude extract is presented in table 1. The result revealed that MBC value (250.0 µg/ml) was significantly higher than the MIC (125.0 µg/ml), indicating that the concentration of the fungal extract would have to be significantly increased to kill bacterial cells, instead of inhibiting their growth. The result also showed that the MBC/MIC ratio was 2 (which was less than 4). Hence, the fungal ethyl acetate crude extract possesses bactericidal effect towards MRSA.

**Table 1: MIC and MBC values of the fungal ethyl acetate crude extract towards MRSA**

| Microorganisms            | MIC (µg/ml) | MBC (µg/ml) | Ratio |
|---------------------------|-------------|-------------|-------|
| Methicillin-resistant S. aureus (MRSA ATCC 33591) | 125.0 | 250.0 | 2 |

**Time-kill study**

Time-kill study of the ethyl acetate crude extract of *L. pseudotheobromae* IBRL OS-64 against MRSA ATCC 33591 at different extract concentrations viz. 1/2MIC (62.5 µg/ml), MIC (125.0 µg/ml), 2MIC (250.0 µg/ml) as well as a control are shown in fig. 1. At the extract concentration of 1/2MIC, the MRSA cell was constantly growing at the lag phase mirroring the growth control.
However, the viable cell counts slightly decreased after 4 until 8 h of incubation. Then, the bacterial growth was observed to be increased after 12 h of incubation and afterwards which reached the viable cell counts almost mirroring the control. The emergence of resistance cells revealed that the extract had not exerted a bactericidal effect on MRSA ATCC 33591 at low concentration such as at 1/2MIC. Besides that, there was a slight decline in a number of cell growth in the culture treated with the MIC concentration of extract from 0 to 8th h. The decrement of viable cell counts was observed throughout the incubation period until all of the bacterial cells were killed after exposure to the extract for 48 h.

Similarly, the viable cell counts of MRSA ATCC 33591 after treated with concentration extract of 2MIC showed the same pattern as at MIC. As for 2MIC, the bacterial growth started decreasing after 4 h of exposure to the extract and declining throughout the incubation period until it reached 36 h whereby all bacterial cells were killed. The time for killing activity of the extract on MRSA ATCC 33591 was shorter at higher concentration of extract (2MIC) as compared to lower concentration (MIC and 1/2MIC). The results revealed that the ethyl acetate crude extract of *L. pseudotheobromae* IBRL OS-64 possessed strong bactericidal effect against MRSA ATCC 33591 at higher concentration. The results also reflected that the bactericidal effect of the extract was concentration and exposure time dependent.

The time-kill study of control showed that the bacterial growth occurred exponentially until reaching its maximal bacterial concentration at the stationary phase. As for the lowest extract concentration (1/2MIC), the reduction of bacterial cell was observed in the first 4 h of exposure and the re-growth was then occurred.

**Fig. 1:** Time-kill study of the ethyl acetate crude extract of *L. pseudotheobromae* IBRL OS-64 against MRSA ATCC 33591 at different extract concentrations [Triplicate readings were recorded, mean ± SD]

Table 2 shows the time to achieve 25, 50, 90, 99, and 99.9% growth reduction in initial inoculum of MRSA ATCC 33591. The result reflects that the longer incubation time was required to obtain a higher percentage of bacterial cell reduction. On the other hand, results also revealed that higher concentration of extract was required to exert in inhibiting or killing effects of MRSA cells.

Reduction in cell viability was only observed in the extract twice the concentration of the MIC. As shown in table 2, higher concentrations of the extract are capable to kill bacterial cells up to 99.9%. In fact, at the highest concentration of extract (2MIC) shorter time was needed to reduce and killed the MRSA cells (32-36 h) as compared to the MIC value with incubation period in the range of 44-48 h.

**Table 2:** The time to achieve 25, 50, 90, 99, and 99.9% growth reduction in initial inoculum of MRSA ATCC 33591

| Percentage of reduction (%) | Control | ½ MIC | MIC | 2 MIC |
|----------------------------|---------|-------|-----|-------|
|                            | NR      | NR    | 4-8 | 0-4   |
| 50                         | NR      | NR    | 4-8 | 4-8   |
| 90                         | NR      | NR    | 16-20 | 4-8 |
| 99                         | NR      | NR    | 20-24 | 16-20 |
| 99.9                      | NR      | NR    |       |       |

Key: NR = not reached

**Structural degeneration and morphological changes of the extract treated cells**

The effects of structural degeneration of MRSA ATCC 33591 cells after exposure to the ethyl acetate crude extract of *L. pseudotheobromae* IBRL OS-64 were studied via SEM observations. Based on the comparison among the SEM micrographs, the ethyl acetate crude extract was found to affect significantly on the morphological changes and the degeneration of MRSA cell structures (fig. 2) of the treated and untreated (control) cells. Fig. 2a demonstrates the untreated or control cells with the presence of regular Gram-positive bacterial cells, undamaged and smooth surface with the intact spherical shape. The cells were also observed growing actively which indicated by the cells underwent binary fission (red arrows). After 12 h exposed to the extract (fig. 2b), the formation of small dimples and dents (red arrows) or cavities were observed which indicated the primary effect of the extract. Fig. 2c shows the bacterial cells after exposed to fungal extract for 24 h. Results revealed that some of the bacterial cells become irregular in shape and bigger in size (black arrow). Despite of irregular shape, some of the bacterial cells were shrunken and crumpled. Fig. 2d illustrates the effect of fungal crude extract towards bacterial cells after 36 h of exposure time.

Severe damaged of the cell morphologies were observed as the cells envelope (cell wall) were broken (indicated by black arrow) and resulted in leakage of cytoplasm. The lysed cell walls were extremely devoid of cytoplasmic contents which led to completely collapsed and left only the homologous cell masses (or cells debris) which was indicated by red arrow. Eventually, the cell death completely occurred.
The development of resistance by existing pathogenic bacteria to commercial drugs is a relevant problem faced by health services and has become a serious concern around the world [23]. Several factors have favoured this scenario, such as the extensive and often inappropriate use of antibiotics, poor hygienic conditions, continuous movement of travellers, increased numbers of immunocompromised patients, and delay in diagnosis of infections [24]. As a result, an intensive search for new, effective antimicrobial agents is necessary, which is facilitated by exploring new niches and habitats [25, 26].

In the search for new anti-MRSA compounds, _L. pseudotheobromae_ IBRL OS-64, an endophytic fungus residing in the leaf of _O. sanctum_, one of the medicinal plants that have many usages to treat various illnesses was isolated. Medicinal plants have been proven to accumulate a wealth of bioactive compounds and are thus chemically highly defended [27]. Any endophyte that is able to colonize these plants needs to cope with their chemical weapons that will restrict any successful microbial invasion [28]. Again, these pronounced stress conditions are expected to favour unusual endophytes that will give rise to likewise unusual metabolites [29].

The _in vitro_ antibacterial activity of drugs is usually assessed by determining of the MIC and MBC levels after overnight aerobic incubation following standard protocols [20]. The MIC is defined as the minimal concentration of antibiotic that prevents a clear suspension of 10^5 colony-forming units (CFUs) of bacteria/ml from becoming turbid after overnight incubation. Turbidity usually connotes to growth at least a 10-fold increase in bacterial density. Because clear bacterial suspensions may have bacterial densities that are 10^6 CFU/ml or less, the MIC may actually be bactericidal to some extent. The minimal bactericidal concentration (MBC) on the other hand is defined as the minimal concentration of an antibiotic that is bactericidal. It is determined by subculturing broth dilutions that inhibit the growth of test bacterial cells (i.e., those at or above the MIC level). The broth dilutions are streaked onto agar and incubated for 24 to 48 h. The MBC is the lowest broth dilution of drug that prevents the growth of the test bacteria on the agar plate. Failure of the bacteria to grow on the plate implies that only nonviable organisms are present.

From the data obtained in this study, _L. pseudotheobromae_ IBRL OS-64 ethyl acetate crude extract on MRSA cells was found to be bactericidal. This was due to the low ratio of MBC/MIC values (equivalent to 2) where the MBC value was two-fold higher than MIC. Levison [30] stated for bacterial drugs the MBC values are usually not more than four-fold higher than their MIC values.

Time-kill test has been used to investigate a wide range of antimicrobial agents and it is the most appropriate method for determining the bactericidal effect. It is a flexible alternative approach that provides more dynamic information regarding on relationship effect of concentration over the time of antimicrobial agents [31]. The time-kill test reveals a time-dependent or a concentration-dependent antimicrobial effect. This approach was frequently used as the basis in the investigation of pharmacodynamics drug interactions and it provides descriptive information on the pharmacodynamics of antimicrobial drugs [32]. The antimicrobial effect of various extract concentrations levels can be directly compared over a wide range of concentrations and times through this approach [33]. The result was consistent with the previous study in the bacterial kill-time curve of tedizolid [31]. At low extract concentration, the post-antibiotic effect might happen whereby the remaining bacteria will begin to resume their growth [34]. Biphasic killing curve with a decline in killing rate over the time was observed when bacterial cells were exposed to a lower concentration of the extract, indicating the presence of persister which less susceptible to antimicrobial drugs [35]. The regrowth phenomenon was attributed to two distinct subpopulations with different susceptibility in which the selective growing of resistant sub-population take over the preferential killing of the susceptible sub-population at a specified time of interaction [32].

The Time Kill Test is used in microbiology to assess a test object's _in vitro_ antimicrobial activity in relation to time. Generally, in a Time-Kill Test, a 3 Log_{10} reduction is considered the minimum level that would indicate a product has significant killing activity against a particular test microorganism, while, as in the minimal bactericidal (MBC) test, bactericidal activity is defined as a 99.9% or greater killing efficacy at a specified time. In conclusion, the results of the present study suggest that time-kill methodologies offer an _in vitro_ approach to the initial selection of new agents or combinations of agents for therapy.

Usually the crude extracts of natural products, whether plants or microbial extracts would affect the cell wall biosynthesis and also the cell membrane permeability. As revealed by the results from SEM micrographs obtained from this study, the effects of the extract on MRSA cells were concentration and exposure time dependants. Meaning at higher concentrations (more than the MIC value) the extract causes the formation of cavities and small dents. The sizes and the depth of those cavities and dents increased as the

**DISCUSSION**

**Fig. 2: SEM micrographs of the MRSA ATCC 33591 cells treated with 0.25 mg/ml of _L. pseudotheobromae_ IBRL OS-64 ethyl acetate crude extract at different exposure times, (a) 0 h [control] (b) 12 h (c) 24 h (d) 36 h. Scale bars: 200 nm**
concentration increased and the exposure time prolonged. The cell wall and cell membrane were destructed or disintegrated, possibly indicated the rupture of cell membrane as well. Hartmann [36] suggested the formation of holes, craters and dents on the surface of bacteria cells indicated a failure or a mechanism rupture of the cell wall and membrane. Besides that, the bacterial cells were observed to compactly resemble and clump together and this may led to stress response to the extract. According to Bible [37], cell-to-cell clumping is a special metabolical scavenging strategy by bacterial cells in order to prepare their cells for further stress conditions. Basri [2], found the MRSA cell treated with acetone extract which combined with vancomycin produced the bigger size of cells and the cell treated with solely acetone extract were in inhomogeneous and irregular shape. Watanabe [38] postulated that the use of arbekacin promoted the formation of thick cell walls of MRSA which could be due to the rapid continuation of cell wall biosynthesis whereby the cell division was inhibited. Some antibiotics including aminoglycosides able to inhibit the initial step in DNA replication and thus inhibited bacterial cell division, but they are not able to completely block the protein synthesis of the bacteria [39, 40]. Besides that, the MRSA cells were observed to be in crumpled and shrunken stages. Ibrahim [41], stated that the reason of the irregular shape of bacterial cell treated with fungal extract (such as cell surface crumpled and shrinkage of the cells) could be due to extensive loss of cell organelles and also the leakage of cytoplasm, supported by the damage of cell envelope.

This condition also may be due to the initiation of cell's autolytic mechanisms and removal of crucial ions and molecules [42]. The collapsed cells with shrunken cell residues may represent the leakage or loss of cytoplasmic contents. Another interesting point is that MRSA is a Gram positive bacteria and is more susceptible to any antibacterial agents compared to Gram negative bacteria.

CONCLUSION
The result of this study revealed that the ethyl acetate crude extract of L. pseudotheobromae IBRL OS-64 exhibited bactericidal activity towards MRSA ATCC 33591 with time and concentration dependents. The fungal extract also showed antibacterial activity by disrupting the cell wall and cell membrane of MRSA which cause cell lysis and ultimate death.

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AUTHORS CONTRIBUTIONS
Taufiq MMJ did all the experiments including leaf sampling, isolation of endophytic fungi, antibacterial activity, time-kill study and structural degeneration (SEM). Darah I was the principle investigator and she supervised the whole works including sample preparation, isolation of endophytes, time-kill study, the antibacterial activity of extract as well as structural degeneration through SEM. All the works, except SEM was performed in her laboratory. She, along with Taufiq MMJ was prepared and corrected the manuscript.

CONFLICT OF INTERESTS
Authors have declared that no competing interests exist.

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