The cytotoxicity, anti-inflammation, anti-nociceptive and oral ulcer healing properties of coconut shell liquid smoke

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Introduction: Coconut shell liquid smoke (CS-LS) is the bio-economy product, consists of light tar or liquid smoke resulting from the pyrolysis of coconut shell, which has components similar to wood (1,2). The benefits of using CS-LS as a natural preservative are safer food preservation, lower cost, and ease of application in the home industry. The use and application of CS-LS itself as an alternative for food product preservative already exist in Indonesia (2). In recent years, CS-LS has also been explored as a natural remedy for treating human ailments. In a diabetic model, the CS-LS has shown the potential to lower fasting blood
glucose and maintain body weight (3). The other potential is in treating diabetic oral ulcers through inhibition of macrophage activity (4), production of the tumor necrosis factor-alpha (1), increasing collagen synthesis (5), and induction of oral ulcer healing (3), by increased fibroblast and growth factor (6).

In traditional medicine in Indonesia, it has been used for generations in the health care system (7). Liquid smoke has long been used by Indonesian ancestors to treat skin diseases caused by fungi, viruses, and bacteria by applying it to the affected area (8). Liquid smoke might also be used as an alternative topical agent for treating burn wounds by reducing pain, increasing fibroblast proliferation and capillary formation (9). The use of natural ingredients as traditional medicine in society is increasingly widespread. Research is needed to evaluate its use in accordance with the rules of health service, and it must be accountable scientifically for the quality, safety, and efficacy of traditional medicine (10).

The chemical changes that occur over time may affect the sensory properties of smoke in the generation and use of liquid smoke, which has potential significance in storage of liquid smoke. The lowest sensory score can be found in after-processed liquid smoke, specifically 5 hours after production. However, they gradually improved over time because the methyl alcohol could react with acetic and formic acids to form esters with softer flavors (11). Additional chemical changes will certainly occur during the storage of liquid smoke, which might result in the loss of undesirable or desirable compounds with the development of other flavor-based compounds (11).

The characteristic of after-process CS-LS obtained from the pyrolysis process and purified through distillation has a yellow color, acidity 2.39, and density 1.0643 g/mL with major component identified as phenol (36.6%), 2-methoxy-phenol (25.2%), and furfural (17.09%) (2). As a natural preservative, the after-processed CS-LS is not a toxic agent (12), possesses anti-nociceptive (13), anti-inflammatory (13), and stimulates the oral ulcer healing by increasing the fibroblast (6), and collagen (5). The production of liquid smoke is not easy and has become a major challenge. During pyrolysis process with a final temperature of 400°C, coconut shell as raw material, only 32.23% of liquid smoke and 58% of charcoal produced (14). During the purification, the amount of liquid smoke produced will be less. Therefore, in addition to the production aspect, storage is also a concern. It is found that the storage of liquid smoke may change its characteristics, cytotoxicity, anti-inflammation, anti-nociceptive, and ulcer healing properties of CS-LS after storage.

Materials and Methods

CS-LS production and storage

The coconut shell of Cocos nucifera L. was collected from the local market in Tembok Dukuh Surabaya (latitude: 7.2520962°S, longitude: 112.7177336°E). The sample was identified by Purwodadi Botanic Garden, Indonesia Institute of Science and identified as species of Cocos nucifera L. The plant name was also verified at http://www.theplantlist.org (accessed November 29, 2016).

CS-LS from coconut shells was obtained through a pyrolysis process. Five kilograms of dried coconut shells were used. The pyrolysis furnace was equipped with a kerosene pump stove and an encircling reactor with a diameter and height of 30 cm and 40 cm, respectively. The furnace was connected to the cooling tubes to condense the fumes and generate liquid smoke. Pyrolysis was carried out at a temperature of 400°C with a heating rate of 3.33°C for 4.5 hours (3). CS-LS was made in Forest Products Research and Development Center Laboratory, Bogor, in 2016 and then stored at a stable temperature at 5°C.

CS-LS characteristics

Acidity was analyzed using a digital pH meter (Mettler Toledo S220, Malaysia). The density was analyzed using a pycnometer (Brand, Germany). The components were determined using gas chromatography–mass spectrometry (GC-MS) model 6890N (Agilent Technologies, Inc., Santa Clara, CA), equipped with a mass spectrometer detector 5975B and DB-5MS UI column (Agilent Technologies, stationary phase; polyethylene glycol, 30 m × 0.25 mm; i.d. 0.25 μm) (1).

CS-LS concentration

CS-LS obtained from the pyrolysis process was considered as CS-LS 100%. CS-LS was prepared in eight different concentrations (1%, 2%, 4%, 6%, 8%, 10%, 12%, and 14%) diluted by sterile water (% v/v).

Cytotoxicity test

Baby hamster kidney (BHK-21) cells were cultivated in a roux bottle and harvested with Trypsin-Versene solution. BHK-21 cells were inserted into 96 wells microplate, added with D-MEM media containing 10% fetal bovine serum albumin and incubated at 37°C for 24 hours.

Eight concentrations of CS-LS (1%, 2%, 4%, 6%, 8%, 10%, 12%, 14%, 100%) were added into 96 microplate wells. Each concentration replicated 8 times. The microplate wells were then incubated for 24 hours, and after 24 hours, each well was added a solution containing MTT reagent in phosphate buffer saline (PBS), then re-incubated for 6 hours. Furthermore, to each well was added dimethyl sulfoxide (DMSO) and shaken with a plate shaker. The
wells would be read at Elisa Reader in 620 wavelengths. The calculation results are said to be non-toxic if ≥ 60% of cells be live. The percentage of the live cells was calculated by the formula (15):

$$\text{% live cell} = \frac{(\text{test group} + \text{test media}) \times 100}{\text{cell} + \text{media}}$$

Information:
% live cells: optical density for each sample
Test group: optical density after each test
Media: optical density on the average of each media control
Cell: optical density on average of cell control

**Anti-inflammatory properties**
The anti-inflammatory properties of the CS-LS were observed by the carrageenan-induced rat paw edema test. Forty healthy male *Rattus norvegicus* rats weighing 110–120 g, 3–4-month-old were used. Animals were divided into 10 groups and each consisted 4 animals. The animals were acclimatized under normal environmental conditions for a few days before starting the experiments (temperature 20–22°C and 12-hour dark/12-hour light cycle).

The first group served as control and received normal saline solution (10 mL/kg, p.o). The second group received indomethacin (10 mg/kg, p.o) suspended as the standard medication in normal saline solution (0.9 % w/v). The other eight groups received the CS-LS 1%, 2%, 4%, 6%, 8%, 10%, 12%, 14%, and 100% (10 mL/kg, p.o).

One hour after oral administration, all animals were injected with 0.1 mL of 1% (v/v) carrageenan solution into saline at the right hind paw sub-plantar region. The paw volume of each rat was measured using a digital caliper before the carrageenan injection and then at 15 minutes intervals up to 1 hour after carrageenan administration. The volume of each rat paw was measured before carrageenan injection using a digital caliper and then at intervals of 15 minutes to 1 hour after carrageenan administration. The percentage of edema was determined for each category as follows:

$$\text{Edema} = \frac{(V_t - V_0)}{V_0} \times 100$$

Information:
$V_0$: the volume before carrageenan injection (mm)
$V_t$: the volume at t minutes after carrageenan injection (mm).

**Anti-nociceptive properties**
A hot-plate method was performed for anti-nociceptive properties. A total of 40 adult male *Mus musculus* mice, weighing 20–30 g were used and divided into 10 groups of 4 animals each. The animals were acclimatized under normal environmental conditions for a few days before starting the experiments (temperature 20–22°C and 12-hour dark/12-hour light cycle).

The first group was controlled and received normal saline solution (10 mL/kg, i.p). The second group was treated orally with tramadol (40 mg/kg, i.p) as normal medication dissolved in saline solution (0.9% w/v). The other eight groups received the CS-LS 1%, 2%, 4%, 6%, 8%, 10%, 12%, 14%, and 100% (10 mL/kg, p.o), respectively.

The rats were accustomed to the hot-plate for 3 consecutive days prior to the experiment by putting them on a plate that was held at room temperature for 15 minutes. To carry out the test, every animal was put on a 52°C hot-plate daily. Latency to exhibit nociceptive responses, such as licking paws, was determined 30, 45, and 60 minutes after the test substances or saline was administered. Prior to the administration of the standard medication and tested substance, the reaction time for thermal pain was measured. A cut-off time for a response to the thermal stimulus was set at 60 seconds to avoid tissue damage of the rat paws.

**Oral ulcer healing**
Twenty-one male *Rattus norvegicus* rats, weighed around 120–160 g were used as animal models. A 10 mm oral ulcer was created using a round stainless blade, in the labial fornix incisive inferior, after anesthetizing the animals, using a combination of xylazine and ketamine. After 24 hours, the oral ulcer appeared as white color surrounded by erythematous area. At this point, the treatment of CS-LS 8% and CS-LS 100% were performed at a dose of 1 μL/g weight once a day on the oral ulcer for three, five, and seven days (3).

**Statistical analysis**
Statistical analysis for all tests was performed using one-way analysis of variance (ANOVA) sequentially followed by LSD post hoc test using SPSS software, version 24.0 for Mac (SPSS, Inc.). Data are represented as the mean and standard deviation. A $P<0.01$ was considered to be significant.

**Results**
**CS-LS characteristic**
CS-LS had a yellow color. The acidity of CS-LS was 2.296 and density was 1.0102 g/mL. After long-term storage, CS-LS was identified as having 14 components (Figure 1). The major component analyzed in GC-MS was phenol (32.75%), 2-methoxy-phenol (guaiacol) (17.45%), and furfural (13.09%). Other compounds such as creosol (6.14%), 2-methyl-phenol (4.23%), 4-ethyl-2-methoxy-phenol (2-EMP) (4.61%), and 16-Hentriacontanone (4.62%) were also be found (Table 1).

**Cytotoxicity test**
The highest living cells were observed in CS-LS 100% (68.08%) and CS-LS 8% (60.64%). The lowest living...
cells of BHK-21 were observed in CS-LS 1% (36.95%). The increased number of living cells was followed by an increase in the CS-LS concentration of 1% until 8%, then decreased until to a concentration of 12% (Figure 2).

**Anti-inflammatory properties**
The effect of CS-LS in carrageenan-induced paw edema is presented in Table 2. The CS-LS in lowest concentration (CS-LS 1%) until highest concentration showed no difference between indomethacin and control in 15 minutes ($P = 0.935$), 45 minutes ($P = 0.984$), and 60 minutes ($P = 1.000$).

**Anti-nociceptive properties**
The anti-nociceptive effect of CS-LS in the hotplate method is presented in Table 3. CS-LS showed a lower thermal pain reaction compared with tramadol. CS-LS 1%, 4%, 6%, and 8% showed lower licking behavior compared to tramadol ($P=0.001$) after 30 minutes administration. The CS-LS 100% had higher thermal pain reaction compared to CS-LS 1%-8% ($P=0.001$).

After 45 minutes administration, the CS-LS 1%, 2%, 4%, and 6% showed a lower thermal pain reaction compared with tramadol ($P=0.001$). The CS-LS 100% had a higher thermal pain reaction compared to CS-LS 1%-8% ($P=0.001$).

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**Table 1.** Component analysis of coconut shell of liquid smoke by gas chromatography mass spectrometry (GC-MS)

<table>
<thead>
<tr>
<th>Name</th>
<th>Area (%)</th>
<th>RT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Furfural</td>
<td>13.09</td>
<td>4,526</td>
</tr>
<tr>
<td>1-(2-furanyl)-ethanone</td>
<td>2.33</td>
<td>6,809</td>
</tr>
<tr>
<td>5-methyl-2-furancarboxaldehyde</td>
<td>1.01</td>
<td>9,392</td>
</tr>
<tr>
<td>Phenol</td>
<td>32.75</td>
<td>10,287</td>
</tr>
<tr>
<td>2-methyl-phenol</td>
<td>4.32</td>
<td>13,921</td>
</tr>
<tr>
<td>p-Cresol</td>
<td>2.85</td>
<td>15,222</td>
</tr>
<tr>
<td>2-methoxy-phenol</td>
<td>17.54</td>
<td>15,829</td>
</tr>
<tr>
<td>2,4-dimethyl-phenol</td>
<td>1.15</td>
<td>19,514</td>
</tr>
<tr>
<td>Creosol</td>
<td>6.14</td>
<td>22,160</td>
</tr>
<tr>
<td>4-ethyl-2-methoxy-phenol</td>
<td>4.61</td>
<td>27,645</td>
</tr>
<tr>
<td>16-Hentriacontane</td>
<td>4.52</td>
<td>61,981</td>
</tr>
<tr>
<td>16-Hentriacontane</td>
<td>2.63</td>
<td>62,694</td>
</tr>
<tr>
<td>16-Hentriacontane</td>
<td>2.68</td>
<td>63,407</td>
</tr>
<tr>
<td>16-Hentriacontane</td>
<td>4.38</td>
<td>64,101</td>
</tr>
</tbody>
</table>

**Table 2.** The anti-inflammatory properties of coconut shell of liquid smoke (CS-LS) in carrageenan-induced paw edema

<table>
<thead>
<tr>
<th>Group</th>
<th>15 minutes</th>
<th>45 minutes</th>
<th>60 minutes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>3.90 ± 1.73</td>
<td>14.08 ± 3.09</td>
<td>23.56 ± 7.51</td>
</tr>
<tr>
<td>Indomethacin</td>
<td>3.24 ± 1.29</td>
<td>11.42 ± 3.97</td>
<td>19.60 ± 3.16</td>
</tr>
<tr>
<td>CS-LS 1%</td>
<td>5.62 ± 0.72</td>
<td>12.43 ± 4.08</td>
<td>21.72 ± 12.19</td>
</tr>
<tr>
<td>CS-LS 2%</td>
<td>5.54 ± 2.27</td>
<td>12.32 ± 1.19</td>
<td>21.58 ± 3.63</td>
</tr>
<tr>
<td>CS-LS 4%</td>
<td>5.61 ± 1.39</td>
<td>11.60 ± 3.67</td>
<td>21.32 ± 8.17</td>
</tr>
<tr>
<td>CS-LS 6%</td>
<td>5.69 ± 2.16</td>
<td>11.36 ± 4.40</td>
<td>21.27 ± 5.64</td>
</tr>
<tr>
<td>CS-LS 8%</td>
<td>5.27 ± 2.48</td>
<td>11.23 ± 2.32</td>
<td>20.82 ± 4.24</td>
</tr>
<tr>
<td>CS-LS 10%</td>
<td>5.08 ± 2.79</td>
<td>11.11 ± 4.88</td>
<td>20.67 ± 8.88</td>
</tr>
<tr>
<td>CS-LS 12%</td>
<td>5.04 ± 1.56</td>
<td>10.97 ± 2.02</td>
<td>20.52 ± 3.77</td>
</tr>
<tr>
<td>CS-LS 14%</td>
<td>4.52 ± 2.36</td>
<td>10.79 ± 1.77</td>
<td>20.40 ± 4.79</td>
</tr>
<tr>
<td>CS-LS 100%</td>
<td>5.71 ± 3.45</td>
<td>11.46 ± 3.42</td>
<td>20.57 ± 2.24</td>
</tr>
</tbody>
</table>

*Each value represents mean ± SD (%) volume of paw edema.*
The properties of coconut shell liquid smoke

Table 3. The anti-nociceptive properties of coconut shell of liquid smoke (CS-LS) in hot-plate method

<table>
<thead>
<tr>
<th>Group</th>
<th>Licking reaction*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>30 minutes</td>
</tr>
<tr>
<td>Control</td>
<td>5.50 ± 0.58 a</td>
</tr>
<tr>
<td>Tramadol</td>
<td>17.50 ± 1.29 b,c,d,e</td>
</tr>
<tr>
<td>CS-LS 1%</td>
<td>7.50 ± 0.58 a</td>
</tr>
<tr>
<td>CS-LS 2%</td>
<td>8.00 ± 0.82 a</td>
</tr>
<tr>
<td>CS-LS 4%</td>
<td>9.25 ± 0.58 a,b</td>
</tr>
<tr>
<td>CS-LS 6%</td>
<td>11.50 ± 0.58 a,b</td>
</tr>
<tr>
<td>CS-LS 8%</td>
<td>15.25 ± 0.50 a,b</td>
</tr>
<tr>
<td>CS-LS 10%</td>
<td>16.00 ± 0.82 a</td>
</tr>
<tr>
<td>CS-LS 12%</td>
<td>16.75 ± 1.50 a</td>
</tr>
<tr>
<td>CS-LS 14%</td>
<td>17.25 ± 1.89 a</td>
</tr>
<tr>
<td>CS-LS 100%</td>
<td>18.75 ± 1.50 a</td>
</tr>
</tbody>
</table>

*Each value represents mean±SD (s) licking behavior as thermal pain reaction.
Same character and symbol in every value is considered as significantly different with ANOVA and LSD test with P value < 0.01.

The CS-LS showed lower thermal pain reaction compared with tramadol after 60 minutes of administration (P = 0.001). The CS-LS 100% has higher thermal pain reaction compared to CS-LS 1%-6% (P = 0.001).

Oral ulcer healing properties

The oral ulcer healing ability was measured based on the number of fibroblasts in the oral ulcer tissue stained with hematoxylin-eosin. The fibroblast was calculated on the edge of the ulcer (blue box). The number of fibroblasts after giving CS-LS topically for three days, five days, and seven days is presented in Figures 3, 4, and 5, respectively.

Fibroblast number in the oral ulcer after treating with CS-LS 100% was higher compared with the control group after treatment for three (P = 0.002), five (P = 0.001) and seven days (P = 0.002). When compared with CS-LS 8%, the fibroblast number was lower after seven days (P = 0.001) (Figure 6).

Discussion

This study was aimed to evaluate the characteristics, cytotoxicity, anti-inflammation, anti-nociceptive, and ulcer healing properties of CS-LS after storage. In a previous study, CS-LS obtained from the pyrolysis process had a yellow color, acidity 2.39, and density of 1.0643 g/mL. After 5 years storage of CS-LS in 5°C, these characteristics were the same (yellow color, acidity 2.29, and density of 1.0102 g/mL). The lower temperature was able to maintain the physical characteristics and physicochemical properties of CS-LS itself. Research conducted by Piccirilli et al confirmed that incorporating the liquid smoke with whey protein concentration was able to maintain the physicochemical properties after stored 28 days in 8°C compared to 25°C (16). This result may also confirm that the lower temperature is better at maintaining the physical characteristic and physicochemical properties of CS-LS itself.
The quality of liquid smoke is determined by phenolic compound and lower acidity because that characteristics could determine the antibacterial effect in food preservation (17). The lower acidity is caused by formic acid, acetic acid, and furfural (18). The phenolic compound, polyaromatic hydrocarbon and lower acidity have allegedly played an important role in liquid smoke toxicity (19). Based on these reasons, the safety of liquid smoke needs to be investigated because of its relevance to human health (20).

The major phenolic compounds identified in the CS-LS were such as phenol (32.75%), 2-methoxy-phenol (guaiacol) (17.45%), and furfural (13.09%). This structure has OH- or hydroxyl structure (21). OH- structure has a high quantitative structure-activity relationship and OH bond dissociation enthalpy. Both of these characteristics are responsible for inducing toxicity, because OH can be deposited into the membrane cells, transported into cells by cellular transport, processing the toxicity (20). The process of toxicity to cells through apoptosis mechanism is by generating excessive intracellular ROS production (22). The excessive intracellular ROS production can induce G1-cell cycle arrest by upregulating p21 expression (23), activating p53 (24), and regulating the nuclear factor kappa B (NF-κB) activity (25). On the other hand, phenolic compounds in the CS-LS act as antioxidants. The antioxidant activity of CS-LS is thought to inhibit or reduce free radicals by transferring hydrogen atoms from their hydroxyl groups. A phenolic compound with peroxyl radical (ROO-) reaction mechanism involves a coordinated shift of the hydrogen cation from the phenol to the radical, creating a transition state of an H-O bond with one electron (26).

In a previous study, the important CS-LS compounds identified were phenol (36.6%), 2-methoxy-phenol (25.2%) and furfural (17.09%) (1). The composition had
The properties of coconut shell liquid smoke

In the present study, CS-LS 100% and CS-LS 8% showed safe concentrations by an in-vitro toxicity test. The CS-LS 10%, 12%, and 14% showed lower living cells of BHK21 less than 60% and considered toxic concentrations. At low concentrations, the phenolic compounds activate the mitogen-activated protein kinase, leading to the expression of genes provoking protective mechanisms leading to gene expressions, which trigger protective mechanisms. Nevertheless, these compounds additionally activated the caspase pathway by increasing the phenolic concentrations, contributing to apoptosis (21). Even though the cytotoxicity data of after-process of CS-LS were not available, its therapeutic potentials have been proven (3). There is no relevant data that revealed in-vivo toxicity tests of after-process of CS-LS. Based on the lack of data, the toxicity of after-process and after-storage CS-LS cannot be compared. But, in the present study, it is confirmed that the after-storage CS-LS showed no toxicity of CS-LS 100% and CS-LS 8%.

The concentrations of CS-LS 1% to CS-LS 14% (except CS-LS 8%) showed the toxicity properties. The toxicity of CS-LS can be related to phenolic compounds and polyaromatic hydrocarbons. The phenolic compound that has a relationship with toxicity is phenol. In a study, 60 mg/kg of phenol exhibited toxicity in an animal model (27). A study confirmed that the liquid smoke from rice hulls had components similar to CS-LS, and had no toxicity at doses of 50, 500, 5000, and 15 000 mg/kg body weight (28). A study showed that the doses of phenol around 10-15 mg/L had toxicity (29). Nevertheless, the CS-LS 1% as the lowest concentration, effectively prevented the growth of bacteria such as Escherichia coli, and Salmonella and prevented the production of histamine (30,31).

A previous study demonstrated the anti-nociceptive properties of CS-LS. The CS-LS 100% had better anti-nociceptive properties compared to the CS-LS 50% and CS-LS 25% in the model of acetic acid-induced writhing reflex (13), the same as this study.

The possible mechanism is that phenolic compounds inhibit tissue cyclooxygenase reducing the synthesis of prostaglandin E2. The reduced number of prostaglandins could disrupt the main transduction mechanisms of the afferent nociceptor, inhibiting pain response (13).

CS-LS has the ability to increase the oral ulcer healing in diabetic (3) by increased macrophages (4), decreased pro-inflammatory cytokine (1), increased the fibroblast, growth factor (6), and collagen synthesis (5). In this study, we showed the properties of CS-LS to stimulate oral ulcer healing after 5 years of storage. The CS-LS 100% and CS-LS 8% were able to stimulate fibroblasts after treatment for three, five and seven days. The possible mechanism is related to the stable physical characteristic of CS-LS (Acidity, density, and phenolic compounds). The phenolic compounds, especially phenol and guaiacol, are able to inhibit NF-κB expression in macrophages (4) and decrease the released pro-inflammatory cytokines such as TNF-α (1). The reduction of inflammation will enhance the release of growth factors such as fibroblast growth factor and vascular endothelial growth factor (6) to increase fibroblast proliferation and collagen synthesis (5). Another mechanism involved in the increased fibroblast proliferation is the inhibition of ROS production by phenol and guaiacol. The inhibition of ROS production will inhibit the activation of the FOXO1 transcription factor to increase caspase-3 that has an important role in fibroblast apoptosis (32). This process will increase the number of fibroblasts during oral ulcer healing.

Based on the present results it possesses anti-nociceptive activity. This research confirms that the right temperature will maintain the properties of CS-LS and proves its anti-nociceptive, non-toxic, and oral ulcer healing properties.

Conclusion
After-storage CS-LS showed a high level of acidity. The CS-LS 100% was the optimum concentration for non-toxicity properties, anti-nociceptive and oral ulcer healing.
healing properties. The CS-LS 100% showed potent anti-nociceptive, but not for the anti-inflammation. The CS-LS is a promising natural herb for various medicinal uses, especially for oral ulcer therapy. The future research is needed to compare after-processed CS-LS and after storage CS-LS in-vivo model in the same time and concentrations to confirm its potential in oral ulcer healing.

Authors’ contributions
MDCS got the funding, wrote the draft of manuscript, data analysis, conceived and performed the experiments; FYM performed the experiments; NFA wrote the draft of manuscript; ABRS revised the manuscript and language correction; DSE revised the manuscript; DM revised the manuscript; IA revised the manuscript; AABN performed the experiments; IPR performed the experiments. All authors read and confirmed publication of the paper.

Conflict of interests
All authors declare no conflict of interest related to this manuscript.

Ethical considerations
This study was conducted in full compliance with the Guide for the Care and Use of Laboratory Animals, in National Health Research and Development Ethics Standard and Guidelines Council (2017), Minister of Health, Republic of Indonesia. The study was approved on February 05, 2020, by the Ethical Committee of Health Research, Faculty of Dental Medicine, Universitas Airlangga (registered number: 047/HRECC.FODM/II/2020).

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