INTRODUCTION

The skin is the largest organ of the human body and is largely responsible for preserving homeostasis and shielding the body from the harmful effects of the outside world. The skin not only performs essential tasks but also plays a significant element in the immune system’s fight against infections. Skin-resident cells such as Langerhans cells, keratinocytes, melanocytes, mast cells, and macrophages emit tiny, hormone-like signal peptides known as cytokines that serve as local immunity modulators or draw in more immune cells in response to internal or external inputs. Specific membrane receptors that are present in the majority of cells are required for their action. Even though a number of skin conditions, such as psoriasis, atopic dermatitis, seborrheic dermatitis, and contact dermatitis, are characterized by inflammation, their cellular immune responses and cytokine profiles differ from one another. Some in vivo as well as in vitro models replicate the skin’s inflammatory response. Neutrophils and monocytes move quickly into the damaged skin during the inflammatory phase, which primarily involves activation of the innate immune system. Hemostasis occurs concurrently with this phase, which is referred to as the first stage of wound healing. There is several studies that explained the mechanism of inflammation in the skin, the free radical theory is the one of mechanism that is popular in the sciences. According to the free radical hypothesis, improperly produced reactive oxidative species (ROS) can cause DNA damage as well as oxidative proteins, nucleoid acids, and lipids. On the other hand second theory is the programmed cellular senescence theory. Keratinocytes, macrophages, dendritic cells, and mast cells are among the local skin cells that are exposed to danger signals as a result of injury. These warning signals may be roughly divided into two categories: PAMPs are pathogen-specific compounds, such as polynucleotides and essential polysaccharides produced by bacteria, that are not present in the host. Stressed cells going through necrosis emit chemicals known as damage-associated molecular patterns (DAMPs). Bacterial metabolites, notably short-chain fatty acids generated by anaerobic bacteria, impaired the function of white blood cells. More tissue damage is caused by higher levels of cytotoxic enzymes and ROS generation. Exotoxins from bacteria attack different cell types and cause tissue necrosis. This situation is made worse by local hypoxemia brought on by vascular blockage. However, the histological of skin damage after sepsis in the skin is still unclear. This study examined the skin damage after LPS induction in a short time.
METHODS

Study Design
This study used an experimental design with a control group that was randomized solely for the post-test. In this investigation, male mice *Mus musculus* aged 8 weeks were employed. The vivarium chamber was kept at a regulated temperature of 22.5°C and was kept on a 12-hour light/dark cycle (lights on at 9 am, lights out at 9 pm). Before starting the experimental treatment, all rats were confined for 7 days to acclimate. The three groups of mice were arbitrarily divided into the Control group (Ctrl group, n=4), LPS+4h (LPS+4h, n=4), and LPS+8h (LPS+8h, n=4). Sigma-Aldrich Company (Sigma-Aldrich Co., St. Louis, MO, USA) acquired LPS (Escherichia coli; O127:B8). LPS was injected intraperitoneally at a dose of 25 mg/Kg.

Data collection
The abdomen skin was immersed in paraffin, fixed by immersion in 10% buffered formaldehyde overnight, and then cut into coronal slices that were 5 mm thick. Hematoxylin and eosin was used to stain brain slices after deparaffinization for standard histological analysis.

Data analysis
The mean and SEM of the data are displayed. The SPSS ver. 25 statistical analysis program was used. One-Way ANOVA was used to compare more than three sets of data, and Tukey’s multiple comparison test was used to assess the results. When the p-value was 0.05 or below, the difference was deemed significant.

RESULTS

Leukocyte concentration and body weight changes following LPS-induced sepsis
White blood cells were examined to assess how an LPS injection caused sepsis. Figure 1A demonstrated that, when compared to a control group, the 4 h and 8 h groups of LPS injection considerably (p-value<0.05) increased the blood’s leukocyte concentration. On the other hand, LPS-induced sepsis substantially reduced bodyweight (p-value<0.05). (Figure 1)

Skin Histopathologic Alterations Following LPS Induced Sepsis
The morphology of skin thickness in the control group was normal, according to the results of the histopathologic study of the slice stained with hematoxylin and eosin. The thickness did, however, significantly diminish in mice after 4 hours and 8 injections of LPS (p-value<0.05).

DISCUSSION
Infection combined with systemic symptoms of infection is known as sepsis. As a result, it is more than just the local organ pathological damage and suggests that one or more other crucial organs may not be functioning properly. Our study’s major objective was to look at skin thickness in LPS-induced sepsis. In previous investigations, it was discovered that sepsis disturbs the tissue that connects wounds. However, another study found that septic patients had delayed epidermal wound healing. This finding suggested that sepsis had an enhanced blood flow response, which may have resulted from a...
high level of systemic inflammation.\textsuperscript{13} According to a previous study, in this study showed that LPS-induced sepsis in mice model was decreased the skin thickness. This funding is related to mechanism of inflammation in the sepsis mechanism. Recent research has clarified the signaling cascade downstream of interleukin and TNF receptors, two receptors linked to sepsis. In addition to LPS-induced inflammatory systems, this model can be used to monitor and analyze several aspects of cutaneous disorders, oxidative stress, skin irritation, and functional tests.\textsuperscript{14} Based on study by Sommer et al., 2013 stated that the serum levels of IL-6 and TNF-a in septic and non-septic mice to confirm sepsis following the CLP procedure. The weight of the spleen was assessed for persistent inflammatory reaction. After sepsis was induced, septic animals had significantly heavier spleens than non-septic mice which were both statistically significant.\textsuperscript{12} Weight of the spleen was also assessed for persistent inflammatory reaction. After sepsis was induced, septic animals had significantly heavier spleens on days 10 and 16 (0.55 g and 0.35 g, respectively) than non-septic mice (0.21 g and 0.17 g, respectively), which were both statistically significant (p,0.001 for days 10 and 16).

The secreted cytokines have an important role as a modulator of the innate immune system and maintain the homeostasis and function of the various types of cells that make up the skin. Where this is related to the signaling pathway mediated by certain receptors through the activation of JAK-STAT and NF-kB transduction signals in inflammatory reactions in the skin.\textsuperscript{15}

We assessed the inflammatory response to LPS stimulation in the current investigation. The quantity of leukocytes in the blood was observed to have decreased. Leukocyte production has increased, indicating the presence of an infectious process that led to the induction of sepsis by LPS. Similar to this outcome, the earlier study looked at the significance of septic pathophysiology’s temporal responses at various stages.\textsuperscript{16} After 3 hours of LPS treatment in rats, a related investigation discovered that the WBC count in whole blood drastically dropped.\textsuperscript{17} There are still many limitations to this study, including the lack of any design- or analysis-based adjustments and any further compounding variables that might affect the outcomes.

CONCLUSION
In conclusion, we demonstrated that LPS-induced septic mice cause damage to the skin, that changes in skin thickness due to the inflammatory process due to sepsis. Further studies are needed to validate and re-evaluate these findings so that these finding can be used as a base of further studies and treatment.

FUNDING
The authors declare no funding in this study.

CONFlict OF INTEREST
The authors declare no conflict of interest in this study.

ETHICAL STATEMENT
The Use Committee of Hang Tuah University gave its approval to all techniques and processes (I/032/UHT. KEPK.03/VI/2020). In accordance with the Handbook for the Care and Use of Laboratory Animals, experimental techniques and processes (I/032/UHT. KEPK.03/VI/2020) in this study.

AUTHOR CONTRIBUTION
All authors contributed equally to this study.

REFERENCES