Landiolol Hydrochloride Ameliorates Liver Injury in a Rat Sepsis Model by Down Regulating Hepatic TNF-α

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Abstract

Aims: The effects of a beta blocker, especially an ultra-short acting selective beta blocker, such as landiolol hydrochloride on the organ protection in sepsis are unclear. The present study aimed to investigate whether acute (early hours) liver injury in a rat model of sepsis induced by lipopolysaccharide (LPS) administration: a) can be corrected by administering landiolol and b) whether landiolol’s effects on liver injury is accomplished by diminishing the elevated expression of inflammatory cytokine, such as tumor necrosis factor (TNF-α) and a vasoconstrictor peptide, such as endothelin (ET)-1.

Methods: Eight (8)-week-old male Wistar rats were administered for three hours with either LPS (n=12), or continuously with LPS plus landiolol (n=11). Control rats were treated with saline only in a similar manner as the treatment group during the relevant time points (n=13).

Results: Following LPS administration, blood gas and hemodynamic parameters were significantly altered compared to control rats at 3 h. Also, At 3 h after LPS administration, circulatory levels of ALT, AST, TNF-α and ET-1 were significantly increased. In addition, at 3 h after LPS administration significant features of hepatic injuries at morphological levels were also evident. Co-treatment of rats with LPS and landiolol ameliorated hepatic injury at 3 h post-treatment, as well as reversed elevated circulatory levels of factors associated with liver injury back to normal levels, such as AST and ALT, and local hepatic levels of TNF-α.

Conclusion: Based on the current findings, it can be stated that landiolol may exert protective effects on liver injury in septic rats by normalizing local expression levels of inflammatory cytokine, such as TNF-α.

Keywords: Landiolol hydrochloride; Liver injury; TNF-α; Endothelin-1; Sepsis; Rat model

Introduction

Sepsis, a critical medical emergency, is associated with tissue hypoperfusion and metabolic impairment, which may contribute to the subsequent development of multiple organ failure normally associated with this disorder [1]. The liver is one of the organs that are normally damaged during the pathogenesis of sepsis and septic shock. Therefore, protecting the liver [2-4] is an important target in sepsis treatment and management. Although numerous studies on infection- or sepsis-induced liver injury have been conducted [5,6], to date no effective treatment has been reported on this disorder. We know that during sepsis, liver functions are altered by the activation of inflammatory processes [6,7] to date no effective treatment has been reported on this disorder. We know that during sepsis, liver functions are altered by the activation of inflammatory processes. For instance, both ex-vivo and in-vitro studies have demonstrated that tumor necrosis factor-α (TNF-α) is released in response to lipopolysaccharide (LPS), primarily by Kupffer cells [8,9]. Specifically, LPS stimulates Kupffer cells to secrete TNF-α, which, subsequently, contributes to the pathogenesis of LPS-induced liver injury by a direct or an indirect polymorphonuclear leucocyte-dependent mechanism [9,10].

Endothelin-1 (ET-1), a potent vasoconstrictor with vasoproliferative activity, is believed to participate in the pathogenesis of sepsis, and its plasma (ET-1) levels significantly increase [11] in sepsis. The possible involvement of the ET system in human septic shock is further supported by a clear correlation between endothelin plasma levels and morbidity and mortality in septic patients [12,13]. Infusion of ET-1 in human causes cardiovascular changes, in part resembling those seen during sepsis i.e. decreased cardiac output and vasoconstriction in the pulmonary, renal and splanchnic circulation [13]. In our previous study, we clearly demonstrated that ET-1 is upregulated in liver during sepsis in a time-dependent manner [14].

Landiolol, an ultra-short-acting and highly cardio-selective β-1 blocker, has become useful for various medical problems in recent days, as evidenced from both clinical and animal studies. Recent studies have demonstrated that co-treatment of LPS with landiolol protects against acute lung injury and cardiac dysfunction in a rat model of LPS-induced systemic inflammation, which was also associated with a significant reduction in serum levels of the inflammation mediator HMGB-1 and histological lung damage [15]. More recently, our group has demonstrated that landiolol treatment significantly normalized various components of altered cardiac ET-1 signaling system in septic rat [16]. In addition, Ogura et al. reported the reno-protective effects of landiolol hydrochloride during sepsis by normalizing the altered expression of ET-1 and HIF-1 alpha levels [17]. However, no study has
yet investigated whether landiolol has protective effects on liver injury during sepsis.

In the current study, we investigated whether landiolol hydrochloride plays an important role in ameliorating liver injury during sepsis and whether such blockage involves attenuation of hepatic TNF-α and ET-1 expression.

Materials and Methods

Animal preparation

In the present study, we used male Wistar rats (200–250 g, 8 weeks old). Sepsis was induced by intra-peritoneal (IP) administration of bacterial LPS from Escherichia coli 055: B5 (Sigma Aldrich, Saint Louis, USA) (15 mg/kg), dissolved in sterile saline. The rats were (n = 38) randomized into group 1 (control, n = 13), group 2 (LPS, n = 12) and group 3 (LPS + landiolol hydrochloride, n = 11). Group 1 received an equal volume of vehicle (sterile saline; 2 ml/body), without LPS. LPS (15 mg/kg, intraperitoneal) was administered at time point 0 h in groups 2 and 3, and then the rats were killed after 3 h post-treatment. However, for group 3, 15 min before LPS or vehicle administration, landiolol hydrochloride was administered continuously intravenously (100 μg/kg/min). This dosage and frequency of landiolol (100 mg/kg/ min) from our pilot studies was found to be the minimal effective dose of landiolol required to normalize the LPS-induced hyperdynamic state in the early stages or hours of sepsis. All rats were killed by Nembutal (sodium pentobarbital, IP, 80 mg/kg body weight) at 3 h after LPS or vehicle only. The blood samples were collected from a polypropylene tube catheter inserted into the left carotid artery for blood gas analysis, and hepatic tissues were harvested gently, snap-frozen in liquid nitrogen, and stored at −80°C. All animals received humane care and the experimental procedures were approved by the Animal Care and Use Committee of University of Tsukuba.

Measurements of hemodynamic parameters

The rats were anesthetized with isoflurane inhalation (1.5%, 1 L/min) and a microtip pressure transducer catheter (SPC-320, Millar Instruments, Houston, TX, USA) was inserted into the left carotid artery [14,16,17]. Then arterial blood pressure and heart rate were monitored with a pressure transducer (model SC-590, Gould, Ohio, USA) and recorded with the use of a polygraph system (amplifier, AP-601G, Nihon Kohden, Tokyo, Japan; tachometer, AT-601G, Nihon Kohden; and thermal-pen recorder, WT-687G, Nihon Kohden) [14,16,17].

Measurements of liver injury markers

Serum aspartate aminotransferase (AST) and alanine aminotransferase (ALT) were measured using kits from Wako Pure Chemical Industries, LTD (Osaka, Japan).

Histopathologic examination

For histopathology, the hepatic tissue specimens were fixed in 4% buffered formalin solution, dehydrated through an ethanol series, embedded in paraffin, and sliced into 5-μm-thick sections. After deparaffinization, sections were stained with hematoxylin and eosin (H&E) using the standard method. The slides were analyzed by two pathologists in double-blinded fashion. These pathologists have specialization on liver morphology and are affiliated with University of Tsukuba Hospital, Tsukuba, Japan. Histomorphometry was done using light microscopy. The points of acute liver injury are hepatic necrosis, degenerative change, inflammatory cell infiltration, Kupffer cell, hemorrhage, councilman body, sinusoid dilation. These points of acute liver injury score were determined 0 (none: -), 1 (mild: ±), 2 (moderate: +), 3 (severe: ++), 4 (very severe: ++++) by a pathologist.

Enzyme immunoassay for TNF-α and ET-1

The concentration of each respective protein/peptide of interest in plasma/serum and liver tissue extracts was determined using the following kits: serum/plasma and liver levels of TNF-α and ET-1 (R & D Systems, Minneapolis, MN), according to the manufacturer’s protocol.

Statistical Analysis

The results were expressed as mean ± SE, and the means were compared by a one-way factorial analysis of variance, followed by Scheffe’s test for multiple comparisons. Differences were considered significant at p<0.05.

Results

Table 1 demonstrates hemodynamic, echocardiogram, and blood gas analysis parameters in experimental animals. Both the systolic and diastolic blood pressure levels were significantly lower at 3 h after LPS administration and were unaffected following treatment of rats with landiolol for 3 h (Table 1). Heart rate was significantly increased in LPS group compared to the control group and significantly decreased in LPS-administered rats treated with landiolol (Table 1). Arterial PaO2 was found to be significantly reduced in LPS-administered rats. However, following landiolol administration, there was a significant reversal effect on arterial PaO2 (Table 1). Blood lactate concentrations were increased dramatically after LPS was given and partly normalized with the treatment of landiolol (Table 1).

Representative images from HE staining with the quantitation of liver injury severity by injury score are shown in Figure 1 from the experimental groups studied in the present study. Control group showed normal liver morphology. The morphology of liver from LPS-administered rats at 3 h showed marked morphological disruptions, such as necrosis (±~ +), degenerative change (±~ ++++), inflammatory cell infiltration (±~ +), Kupffer cell hyperplasia (±~ +), councilman body (±~ +) and sinusoid dilation (±~ +). These histopathological changes were significant in LPS-administered rats compared to those of the control liver group. After three hours treatment with landiolol of LPS-administered rats, there were less pronounced histomorphological disruptions, such as degenerative change (±~ +), inflammatory cell infiltration (±~ +), and hemorrhage (±~ +) in liver. In addition, there was no remarkable necrosis in liver.

|                         | Control          | LPS                           | LPS + Landiolol               |
|-------------------------|------------------|-------------------------------|-------------------------------|
| pH                      | 7.412 ± 0.003    | 7.420 ± 0.020                 | 7.400 ± 0.015                 |
| PaO2 (torr)             | 102.6 ± 2.3      | 86.5 ± 2.6*                   | 98.3 ± 2.6*                   |
| PaCO2 (torr)            | 39.8 ± 1.2       | 34.1 ± 1.4                    | 32.1 ± 2.6                    |
| BE (mmol/l)             | 1.0 ± 0.0        | -3.2 ± 0.5*                   | -5.2 ± 1.0*                   |
| Lac (mmol/l)            | 0.8 ± 0.2        | 3.1 ± 0.4*                    | 1.6 ± 0.1*                    |
| HCO3⁻ (mmol/l)          | 24.8 ± 0.8       | 21.5 ± 0.6                    | 20.0 ± 1.2*                   |
| Heart Rate (bpm)        | 462.0 ± 7.7      | 497.0 ± 4.9*                  | 465.0 ± 8.8*                  |
| Systolic BP (mmHg)      | 118.0 ± 0.5      | 102.0 ± 4.4*                  | 101.0 ± 4.5*                  |
| Diastolic BP (mmHg)     | 77.0 ± 2.0       | 66.0 ± 1.2*                   | 65.0 ± 1.1*                   |

LPS, lipopolysaccharide; BE, base excess; Lac, lactate; BP, blood pressure.

* p<0.05 vs control, #p<0.05 vs LPS

Table 1: Hemodynamic and blood gas analysis parameters in experimental animals.
tissue in septic rats after landiolol treatment. As shown in Figure 2B, the enhanced injury score in LPS-administered rats were significantly ameliorated following treatment of rats with landiolol for 3 h.

Levels of serum AST and ALT (Figure 2A and 2B), increased significantly at 3 h after LPS administration, when compared to the control rats and the rise biochemical markers of interest were significantly ameliorated following the treatment of LPS-administered rats with landiolol. The serum levels of TNF-α were elevated after administration of LPS (Figure 2C). However, landiolol treatment failed to normalize the elevated serum levels of TNF-α in sepsis rats. Consistent with our previous report [16,17], the present data show elevated levels of plasma ET-1 in sepsis rats (Figure 2D), which was unaffected by
Landiolol at 3 h post-treatment. Protein expression levels of TNF-α were elevated in liver tissue after LPS administration, compared to control group, and landiolol treatment significantly normalized the elevated hepatic levels of TNF-α (Figure 3A). In liver tissue, ET-1 peptide levels were significantly higher in LPS-administered group compared to the control group. However, landiolol treatment did not alter levels of enhanced ET-1 (Figure 3B) in septic rat liver tissues.

Discussion

The key findings of the present study are that: 1) treatment of septic rats with landiolol 3 h-post administration was effective in ameliorating the histopathological and biochemical (AST and ALT) parameters of liver injury, respectively; 2) this improvement in the condition of liver injury in LPS-administered rats was accompanied by significant normalization of hepatic TNF-α levels; 3) finally the amelioration of liver injury by landiolol treatment, as observed in the present study, may not involve ET-1 levels in liver tissue in septic rats.

Landiolol hydrochloride is an intravenously administered, ultra short-acting β1-blocker with an elimination half-life of 3–4 min and ≈8-fold greater cardioselectivity than esmolol in vitro [18-21]. It is approved in Japan for the treatment of intraoperative and postoperative tachyarrhythmias, but is also used in clinical practice to prevent postoperative tachyarrhythmias, such as atrial fibrillation after coronary artery bypass grafting [18-21]. As an ultra short-acting β1-blocker with a rapid onset of action and readily titratable and rapidly reversible effects, landiolol represents an important agent for the management of intraoperative and postoperative tachyarrhythmias [18-21]. Very recently, landiolol has been reported to confer organ protection in sepsis animal models through various pathways [15-17]. Recent studies have demonstrated that co-treatment of landiolol in LPS-induced systemic inflammation protects against acute lung injury and cardiac dysfunction in a rat inflammation models [15,16] through the suppression of inflammatory markers. Consistent with these previous findings, the current study also found that landiolol treatment confers hepatic protection both at histomorphological as well as biochemical levels during the early hours of sepsis. Proinflammatory mediators, such as LPS and TNF-α, can directly lead to hepatocellular damage [22]. It has been shown in a previous study that LPS and TNF-α may play a central role in the development of acute hepatic failure after severe trauma and sepsis by directly or indirectly inducing hepatocyte necrosis rather than apoptosis [22]. In the current study, we found increased levels of TNF-α both at circulatory and hepatic tissue level after sepsis induction through LPS administration. This upregulation can directly lead to hepatocellular damage.

Amelioration of liver injury by landiolol may involve the inhibitory action of landiolol on local elevated levels of TNF-α in liver tissue of LPS-induced sepsis in rats. In the present study, we found attenuation of histomorphological hepatic changes, such as degeneration, inflammatory infiltration and hemorrhage following co-treatment of LPS with landiolol by three hours. Interestingly, while landiolol significantly normalized the local elevated levels of TNF-α in the livers of LPS-induced sepsis in rats, it (landiolol) failed to reverse increased levels of TNF-α at the circulatory level. For the moment, we do not have any specific explanation underlying the differential effects of landiolol at 3 h post-treatment.

Figure 3: Protein expression levels of TNF-α (A) and ET-1 (B) peptide levels in liver tissues of the control group, LPS-administered rats (3 h-post treatment), and landiolol-treated LPS-administered rats (3 h-post treatment). These data were generated by ELIZA. Values are mean ± SE *p<0.05 vs. control; #p<0.05 vs. LPS.
landiolol on TNF-α expression in liver versus circulation in septic rats. Further studies are required to clarify these issues. ET-1, a potent vasoconstrictor, has been implicated in the pathogenesis of sepsis. Plasma levels of ET-1 have been shown to be significantly higher in septic patients [11] and their levels (ET-1) have a clear correlation with morbidity and mortality in septic patients, suggesting an involvement of ET in the pathogenesis of septic shock in humans [12]. Further, it has been suggested that ET contributes to the dysfunction of several vital organ systems during septic shock. In our previous study, we have shown that ET-1 is highly upregulated in hepatic tissues in septic rats [14]. In fact, ET-1 is a powerful vasoactive peptide and is secreted by various cells in liver including vascular and sinusoidal endothelial cells and hepatic stellate cells [23,24]. ET-1 also causes strong vasoconstrictive action after binding to its receptor [25]. Thus, a growing body of evidence suggests a potential role of ET-1 in the pathogenesis of liver injury. The present findings show that the amelioration of liver injury in sepsis by landiolol does not involve ET-1 pathway, contrary to our recent studies that demonstrated the significant reversal of landiolol on elevated cardiac and renal tissues ET-1 signaling system in sepsis. In contrast, in the current study, landiolol treatment did not normalize upregulated ET-1 levels in liver tissues of septic rats. Thus, it can be concluded that landiolol has organ-specific (reversal) effects on altered ET-1 levels during sepsis. The present study has several limitations. We cannot rule out from the present findings that what types of liver cells are significantly impacted by the landiolol treatment in sepsis. In vitro studies using various cell types of liver should be conducted in future. In addition, other inflammatory cytokines should be investigated in current study design in depth. Longer duration treatment with landiolol in sepsis models can be done in future in current experimental setting.

Conclusion

The present study demonstrates that landiolol hydrochloride, a selective ultra-short acting beta blocker, ameliorates liver injury in a rat model of sepsis possibly through the suppression of elevated hepatic TNF-α levels.

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