Serum of limb remote ischemic postconditioning inhibits fMLP-triggered activation and reactive oxygen species releasing of rat neutrophils

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Introduction

Limb remote ischemic postconditioning (LRIP) refers to the repeated limb ischemia after a serious ischemic brain or myocardial injury. LRIP protects against cerebral or myocardial ischemia/reperfusion (I/R) injury [1,2], eventually promoting the recovery in ischemic disease patients. Thus, LRIP has exhibited a promising therapeutic potential. I/R injury is a complicated pathophysiological process that occurs through various pathways including oxidative stress and inflammation. However, the underlying mechanisms remain to be fully elucidated. Recent studies support that the reactive oxygen species (ROS) and proteases released from the neutrophils act as a major role in acute reperfusion injury following ischemic stroke [3].

Our previous study revealed that the limb operated with LRIP can ‘attract’ the activated neutrophils, thereby reducing the neutrophils migration to the cerebral ischemic area [3]. Meanwhile, LRIP in rats inhibits the NADPH oxidase activation in neutrophils, a key enzyme which is dedicated to ROS production [4], through TLR4-MyD88-TRAF6 signaling pathway. A recent report presented that the serum extracted from humans who were subjected to remote ischemic preconditioning (RIPC) has a protective effect on human endothelial and intestinal cells against hypoxia-induced damage [5,6]. However, whether the serum of LRIP mediates the protective effect through inhibiting neutrophils activation or migration.

We hypothesized that the LRIP serum from rats could directly reduce the ROS release from neutrophils through...
inhibiting NADPH oxidase. The associated pathways were examined. The finding contributes to the elucidation of the L RIP protective mechanism.

Materials and methods

Animals

Adult male Sprague–Dawley rats (Qinglongshan, Nanjing, China) weighing 250–280 g were used in this study. All animal experiment protocols were approved by the Animal Ethics Committee of China Pharmaceutical University (No. 81503284) and were conducted according to the guidelines of the Institutional Animal Care and Use Committee at China Pharmaceutical University. Animals were housed in a temperature and light-controlled room (23 ± 1°C; 12 h light/12 h dark cycle) and allowed to freely access to food and water.

Reagents

Hanks’ Balanced Salt Solution (HBSS), Trizol (R0016) and SB203580 (SB, S1863) were obtained from Beyotime Biotechnology Co. Ltd. RPMI 1640 and Fetal bovine serum (FBS, 26140-079) were obtained from Wisent Biotechnology Co., Ltd. Formyl-methionyl-leucyl-phenylalanine (fMLP, C-1711) and Apocynin (APO, HY-N0088) were obtained from Sigma-Aldrich and MedChemExpress.

Collection and preparation of LRIP serum of rat

The rats were randomly divided into Serum Sham (SSham), Serum LRIP0 (S LRIP0) and Serum LRIP1 (S LRIP1) groups (n = 5). S LRIP0 and S LRIP1 rats were anesthetized with 10% chloral hydrate and placed on the surgical plate. The femoral artery of the left hindlimb of rats was clamped with a modified blood pressure cuff for 5 min (ischemia) and blood flow was restored for 5 min (reperfusion), which was regarded as a cycle and repeated three cycles [3]. Interruption of the blood supply to the hindlimb was confirmed by cyanosis of the skin on the limb. The blood was collected from the abdominal aorta of rats immediately (SLRIP0) or at 1 h after the completion of LRIP (SLRIP1) or without LRIP operation (SSham) [6]. After clotted for 10 min at room temperature, all of the blood samples were then centrifuged at 5000 rpm for 10 min. The serum was collected and stored at −80°C for further use.

Isolation and culture of rat bone marrow neutrophils

The neutrophils were isolated from rat bone marrow and purified following the manufacturer’s instructions of Rat Bone Marrow Isolation Kit (TBD2013NR, Tianjin Haoyang Biological Manufacture CO., Ltd., Tianjin, China). Rats were sacrificed by cervical dislocation under anesthesia. The femur and tibia of rat were isolated and the soft tissue attached were carefully removed. F solution in Rat Bone Marrow Isolation Kit was forced through the bone with a 10 mL syringe. After dispersing cell clumps, the cell suspension was centrifuged (500 g, 10 min, room temperature) and resuspended in HBSS (Ca2+,Mg2+) solution. Cells were then overlay two-layer solution A gradient of 100% and 80% solution A (diluted in diluent with 100% solution A), and centrifuged (750 g, 25 min, room temperature). Neutrophils from the upper phases were harvested and the remaining red cells were eliminated by red blood cell lysis. After washed twice with cell detergent, the purity of neutrophils was identified by using Wright-Gemssa staining (purity > 90%). The isolated neutrophils were adjusted to 2.5×106/mL with RPMI 1640 complete medium, then subjected to the following treatments and measurements.

Detection of neutrophil viability

To detect the neutrophil viability, cells were separated into 2 groups. In group 1, cells were cultured with RPMI 1640 containing 10% FBS [7]. In group 2, cells were cultured with RPMI 1640 containing 10% S Sham at 37°C under 95% O2, 5% CO2. The neutrophil viability was detected at 0, 3, 6, 12, 24 and 48 h by using the Cell Titer-Lumi™ Cell Viability Assay Kit (C00685, Beyotime Biotechnology Co., Ltd., Shanghai, China) according to the manufacturer’s instructions.

Release of ROS in neutrophils

To measure the release of ROS and the mRNA/protein levels of associated markers, the neutrophils were randomly distributed into 6 groups: S Sham, S Sham + ROS inducer fMLP group, S LRIP0 group, S LRIP1 group, S Sham+ p38 MAPK inhibitor SB group, S Sham+ NADPH oxidase inhibitor APO group [8,9], S LRIP0 and S LRIP1 group cultured with RPMI 1640 containing 10% S LRIP0 or 10% S LRIP1, respectively, the other four groups cultured with RPMI 1640 containing 10% S Sham. 200 μL cell suspension was inoculated into the 96-well culture plate coated with 0.1% gelatin, then cultured at 37°C under 95% O2 and 5% CO2. 24 h later, p38 MAPK inhibitor SB (10 μM, DMSO) [10] and NADPH oxidase inhibitor APO (100 μM, DMSO) [11] were added into SB and APO groups, respectively, then SB group was cultured for 30 min and APO group for 15 min, followed by stimulation with fMLP (10 μM, DMSO) [12] for 10 min. After carefully transferred into 96-well plate and gently washed with PBS, the culture medium was added with 100 μL of 2, 7 dichlorofluorescein diacetate (DCFH-DA) diluted (1:1000) to PBS, then incubated at 37°C for 25 min. Fluorescence value was measured at the excitation at 488 nm and emission at 525 nm with an automated microplate reader (Varioskan Flash, ThermoFisher Scientific, Waltham, MA, USA).

qPCR

Total RNA of neutrophils was extracted by Trizol similar as previously described [3]. Then 1 μL cDNA product and 2 μL primer were added into qPCR super mixture with a final volume of 10 μL [3]. qPCR was performed with 1 cycle for 30 s at 94°C, followed by 5 s at 94°C, 40 cycles (1 cycle for 30 s) at 60°C using a Real-time quantitative PCR instrument (Bio-Rad, USA). The following primer sequences were used (5’ to 3’): p47phox (AC008033.9), forward, ATTTCCTTGGTGATTGATG, reverse, GAGGGATGTTACTACTG; p22phox (AC008080.1) forward, CAGCCACCGAGATTGACCA, reverse, AGG-CAGGGACAGCAGTAA; GAPDH (LC627775.1), forward, TCCTGACCACCAACTGCTTAG, reverse, AGTGCGCATGATGCCATGGACT.
Western blot

Western blot was performed as described previously [3]. The total protein was extracted with cold RIPA lysis buffer from neutrophils and protein concentration was measured using BCA Protein Assay Kit (P0012S, Beyotime Biotechnology Co., Ltd., Shanghai, China), then 40 μg protein from each group were used for protein gel electrophoresis. After the electrophoresis, the protein band was transferred onto the PVDF membrane and blocked with 3% BSA. Followed by incubation with the first antibody overnight at 4°C, (antibodies as below, p47phox (sc-17845, Santa Cruz Biotechnology, Texas, USA), p-p47phox Ser 345 (SA4301368, Sigma-Aldrich, St. Louis, MO, USA), p-p47phox Ser 304 (P14598, Bioworld, Beijing, China), TRAF6 (ab33915, Abcam, Cambridge, MA, USA), MyD88 (ab133739, Abcam, Cambridge, MA, USA), p38 MAPK (86905, CST, Danvers, Massachusetts, USA), p-p38 MAPK (4511S, CST, Danvers, Massachusetts, USA), P13K (17366S, CST, Danvers, Massachusetts, USA), Akt (46855, CST, Danvers, Massachusetts, USA), p-AKT (46855, CST, Danvers, Massachusetts, USA), ERK (46955, CST, Danvers, Massachusetts, USA), p-ERK (43705, CST, Danvers, Massachusetts, USA), JNK (23055, CST, Danvers, Massachusetts, USA) and p-JNK (92555, CST, Danvers, Massachusetts, USA), the membranes were washed and incubated with the horseradish labeled second antibody for 2 h at room temperature. The protein band was analyzed with the gel-imaging system (Bio-Rad, Hercules, CA, USA). The values were calculated after normalization to the amount of GAPDH.

Statistical analysis

All data was statistically analyzed by using GraphPad Prism 5 software (GraphPad Software, Inc., La Jolla, CA, USA). Results were expressed as mean ± SEM. Student’s t-test was used for the comparison between the two groups. One-way ANOVA analysis was used for the comparison of three or more groups, followed by using Dunnett’s post-test, compared with SSham group, SSham + fMLP group significantly increased the excessive of ROS (P < .001). SLRIP0, SLRIP1, SB203580 and APO all could significantly inhibit ROS production (Figure 2).

LRIP serum inhibits fMLP-induced neutrophil ROS release

Compared with SSham group, SSham + fMLP group significantly increased the excessive of ROS (P < .001). SLRIP0, SLRIP1, SB203580 and APO all could significantly inhibit ROS production (Figure 2).

LRIP serum inhibits the activation of neutrophil NADPH oxidase induced by fMLP

The mRNA expression of p47phox and p22phox, two key regulators of NADPH oxidase, were significantly upregulated in the SSham + fMLP group (P < .05). Similar to the SSham group, the phosphorylation of p47phox Ser 304 and p47phox Ser 345 were also significantly upregulated (Figure 3). p47phox mRNA of SLRIP1 + fMLP, SB and APO groups, as well as p22phox mRNA of SLRIP0, SLRIP1 and SB groups were significantly downregulated (P < .05). SLRIP0, SLRIP1, SB and APO all could significantly inhibited the phosphorylation of p47phox Ser 304 and Ser 345. These results indicated that LRIP serum could inhibit the activation of neutrophil NADPH oxidase induced by fMLP.

LRIP serum inhibits the activation of neutrophil NADPH oxidase induced by fMLP

To determine whether LRIP serum inhibits the activation of neutrophil NADPH oxidase induced by fMLP, neutrophils were treated with SSham, SLRIP0, SLRIP1, SB and APO before being activated by fMLP. Compared with SSham group, the mRNA expression of p47phox and p22phox as well as the phosphorylation of p47phox Ser 304 and p47phox Ser 345 was significantly increased in the SSham + fMLP group (P < .05, Figure 3). SLRIP1, SB and APO all could downregulated the expression of p47phox mRNA (P < .05). SLRIP0, SLRIP1, SB and APO all could significantly inhibit the phosphorylation of p47phox Ser 304 and Ser 345. SLRIP0, SLRIP1, SB also could downregulate the expression of p22phox mRNA (P < .05). These results indicated that LRIP serum could inhibit the activation of neutrophil NADPH oxidase induced by fMLP.

LRIP serum inhibits the activation neutrophil via MyD88-TRAF6-MAPKs signaling pathway

Compared with SSham group, the protein expression of MyD88, TRAF6, p-ERK, p-JNK and p-p38 MAPK of SSham + fMLP groups were significantly upregulated. SLRIP0, SLRIP1, SB and APO all could downregulated the protein expression of MyD88, TRAF6, p-ERK, p-JNK and p-p38 MAPK. These results proved that the LRIP serum could inhibit the activation of neutrophil NADPH oxidase induced by fMLP via the MyD88-TRAF6-MAPKs signaling pathway (Figure 4).

LRIP serum inhibits the activation neutrophil induced by fMLP independently of PI3 K/ AKT signaling pathway

Continuously, to determine whether PI3K/Akt pathway is involved. As shown in Figure 5, compared with the SSham group, the expression of PI3K of the SSham + fMLP group was significantly upregulated but the p-Akt expression levels were downregulated by fMLP. SLRIP0, SLRIP1, SB and
APO all could downregulate the expression of PI3K. These results revealed that LRIP serum inhibits the activation of neutrophil induced by fMLP, the PI3K may play a role in it but independently of p-AKT.

Discussion

In this study, we wanted to test the effect and mechanism of LRIP serum on a neutrophil respiratory burst model in vitro. The lifespan of neutrophils in culture with FBS or rat serum was tested. Recent studies have confirmed that the lifespan of neutrophils in the blood circulation of mice is 18 h [13], and in humans, neutrophils can survive in the blood for 5.4 days [14]. Previous research has proved that there are a great amount of functionally competent neutrophils in mice bone marrow, the half-life of neutrophils from bone marrow is longer than the neutrophils from blood [15]. This study showed that the activity of neutrophils cultured with FBS decreased significantly after 3 h, this is similar to previous studies [15]. It is very interesting to find that neutrophils cultured with SSham remained more than 90% viability after a 24 h culture (Figure 1). This proved that there should be some nutritional factors in SSham which satisfied the rat neutrophil survival needs. Based on this result, we chose neutrophils extracted from the bone marrow and the rat serum as the cell culture media, 24 h as the culture time in this study. In addition, SLRIP0, SLRIP1, SB and APO were all found to inhibit the ROS production from neutrophils activated by fMLP (Figure 2). This result proved the inhibiting effect of SLRIP0, SLRIP1 on the ROS release from neutrophils under the stimulated I/R condition and also proved LRIP can exert protective effects through the serum route. Furthermore, results of qPCR and western blot showed that SLRIP1 could downregulate the mRNA expression of p47phox. Both SLRIP0 and SLRIP1 could downregulate the mRNA and protein expression of p22phox, p47phox ser 304 and p47phox ser 345 (Figure 3).

ROS was thought to be the major cause of I/R injury [16–18] and the activated neutrophil NADPH oxidase was thought to be the main source of ROS [19–21]. The reperfusion injury would occur [22,23] after the thrombolytic therapy was used on ischemic disease patients to restore blood flow into the ischemic regions [24,25]. LRIP is effective in alleviating I/R injury of brain, heart, kidney and other organs [3,26,27]. The related mechanisms include reversing the eNOS uncoupling [28], upregulating the expression of Nrfr2 along with heme oxygenase-1 (HO-1) [29], quinone oxidoreductase 1 (NQO-1) [30], inhibiting the inflammation [31–33]. Our previous work has proved that LRIP can inhibit

Figure 1. Combine LRIP serum collecting and neutrophil viability cultured in vitro. (A) Blood was collected before LRIP operation, or 0, 1 h after LRIP operation, serum was named SSham, SLRIP0, SLRIP1 separately. The viability of bone marrow neutrophils was assessed at 0, 3, 6, 12, 24, 48 h under different serum conditions; (B) Cultured with FBS; (C) Cultured with SSham. Data are expressed as mean ± SEM, n = 4–5, ***P < .001 vs. 0 h, *P < .05 vs. 0 h.

Figure 2. The inhibiting effects of LRIP serum on intracellular superoxide generation from neutrophils. Neutrophils were cultured with RPMI 1640 containing SSham, SLRIP0 or SLRIP1 serum for 24 h, followed by incubating with DMSO, Aponycin (100 μM) for 15 min or incubated with SB203580 (10 μM) for 30 min, then activated with fMLP (10 μM) for 10 min. Data are expressed as mean ± SEM, n = 5, ***P < .001 vs. SSham, *** P < .0001 vs. SSham + F.
the activation of neutrophil NADPH oxidase through the TLR4-MyD88-TRAF6 pathway, working to reduce the cerebral I/R injury [3]. But till now, no researches have been done to clarify the effect and mechanism of LRIP serum in the activation of neutrophil NADPH oxidase in vitro. In this study, we wanted to check the protective effect of LRIP serum and clarify the related mechanisms.

![Figure 3](image)

**Figure 3.** The inhibiting effects of LRIP serum on activation of neutrophil NADPH oxidase. qPCR results showed the mRNA expression of p47<sup>phox</sup> (A) and (B) p22<sup>phox</sup>, GAPDH was used as the internal control, (n = 3); (C-D) Western bolt analysis of p47<sup>phox</sup> phosphorylation at Ser 304 and Ser 345 of neutrophils (n = 5). Data are expressed as mean ± SEM, ###P < .001 vs. SSham, #P < .05 vs. SSham, ***P < .001 vs. SSham + F, **P < .01 vs. SSham + F, *P < .05 vs. SSham + F.

The activation of neutrophil NADPH oxidase induced by fMLP will produce a great deal of ROS [34], which can simulate the respiratory burst of neutrophils under I/R injury. In our study, 10 μM fMLP was used to stimulate neutrophils [35]. The catalytic core of NADPH oxidase in neutrophils is gp91phox (Nox2). Activation of Nox2 needs the active p47<sup>phox</sup> (the adaptor protein) to bind onto p22<sup>phox</sup> [36] and

![Figure 4](image)

**Figure 4.** The inhibiting effects of LRIP serum on fMLP-induced neutrophil activation via MyD88/TRAF6/p38MAPK pathway. Representative images of Western bolt assessments and quantitative analysis of the ratio of MyD88 (A); TRAF6 (B); p-ERK (C); p-JNK (D); and p-p38 MAPK (E). Data are expressed as mean ± SEM, n = 5, ###P < .001 vs. SSham, #P < .05 vs. SSham, ***P < .001 vs. SSham + F, **P < .01 vs. SSham + F, *P < .05 vs. SSham + F.
p67phox onto Nox2. The function of p47phox is governed by serine phosphorylation, which occurs during activation of the NADPH oxidase. Phosphorylation of Ser-304 [37] and Ser-345 [38] on p47phox is a necessary event in the priming of neutrophils. Results in this study proved fMLP could upregulate the mRNA expression of p22phox, p47phox in neutrophils and upregulate the phosphorylation of ERK, JNK, P38, p47phox ser 304, ser 345, also upregulate the expression of MyD88, TRAF6 in neutrophils. These proved the activation of neutrophil NADPH oxidase simulated by fMLP was based on the MyD88/TRAF6/p38 MAPK pathway (Figure 6). The results also showed that this kind of activation can be inhibited by SLRIP0 and SLRIP1. This demonstrates that SLRIP0 and SLRIP1 inhibited the production of ROS from activated neutrophils by inhibiting the activation of neutrophil NADPH oxidase.

This study still has some limitations. For example, the kind of serum factor/factors which mediated the protective effect of the LRIP serum and how PI3K regulated the protective effect of LRIP serum, both remain to be identified.

**Conclusion**

In conclusion, the rat serum is more adept to keeping viability of the bone marrow neutrophils. LRIP operation can induce some serum factors to inhibit the activation of neutrophil NADPH oxidase via the MyD88/TRAF6/p38 MAPK pathway, which then alleviate the I/R injury. This study might be meaningful for clarifying the protection mechanisms of LRIP and allowing for further research and development of biological drugs related to I/R injury.
Disclosure statement
No potential conflict of interest was reported by the author(s).

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