Original Article

Adipose-derived mesenchymal stem cell-derived exosomes markedly protected the brain against sepsis syndrome induced injury in rat

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Abstract: This study tested the hypothesis that sepsis syndrome (SS-induced by cecal-ligation and puncture (CLP))-induced systemic inflammation and brain damage in rats were effectively suppressed by allogenic adipose-derived mesenchymal stem cell-derived exosome (AMSC EXO). SD rats (n = 72) were divided into group 1 [sham-control (SC)], group 2 (SS only) and group 3 (SS + AMSC EXO) and equally euthanized at 6/24/48/72 h after SS induction, respectively. By 6/16/24/72 h, flow cytometric analyses demonstrated the numbers of inflammatory cells (Ly6G+/CD11b+/c), immune (CD3+/CD4+ cells/CD3+/CD8+ cells) and early (AN-V+/PI-)/late (AN-V+/PI+) apoptotic cells in circulation were significantly increased in group 2 than in groups 1 and 3, and significantly increased in group 3 than in group 1, whereas the number of T-reg+ cells was significantly progressively increased from groups 1 to 3 (all P < 0.0001). At 6/16/24/72 h, the numbers of (CD3+/CD4+ cells/CD3+/CD8+ cells/T-reg+ cells) in spleen exhibited an identical pattern of circulation among the three groups (all P < 0.0001). ELISA showed inflammatory mediators (IL-6/TNF-α) in circulating/cerebrospinal fluid at 6/24/72 h displayed an identical trend as the immune cells among the three groups (all P < 0.0001). ELISA showed inflammatory mediators (IL-6/TNF-α) in circulating/cerebrospinal fluid at 6/24/72 h displayed an identical trend as the immune cells among the three groups (all P < 0.0001). Microscopic findings demonstrated that the cellular expressions of inflammatory (F4/80+/MMP-9+/CD14+/GFPA+) and brain-damaged (AQP4+/γ-H2AX+) biomarkers at 24/72 h exhibited an identical pattern of immune cells among the three groups (all P < 0.0001). The protein expressions of inflammatory (IL-1β/MMP-9/TNF-α/NF-κB/TLR2/TLR-4/MyD88/HMGB1), apoptotic (cleaved-caspase3/PARP/mitochondrial-Bax) and oxidative-stress (NOX-1/NOX-2/oxidized protein) biomarkers displayed an identical pattern as the immune cells among the three groups (all P < 0.0001). In conclusion, SS elicited vigorously inflammatory reaction not only in circulation but also in spleen/brain, resulting in serious brain damage.

Keywords: Sepsis syndrome, inflammation, oxidative stress, apoptosis, brain damage

Introduction

Sepsis syndrome (SS), characterized by systemic inflammatory reaction caused by infection, is a very common public health issue with an extremely high economic burden through healthcare costs worldwide [1-3]. Abundant data have shown that SS is still the major leading cause of death in ICU, ranging from 20% in sepsis to more than 60% in SS complicated by septic shock [1-8]. Although utilization of advanced antibiotics is the golden standard method for treatment of SS [1, 2, 9, 10], the in-hospital morbidity and mortality rates of SS remain unacceptably high, highlighting that there is an urgent need for innovative and efficacious therapies for treatment.

Undoubtedly, severe sepsis frequently causes multiple organ failure [8, 10-12]. Of these major organs, brain is not only commonly involved but also one of the most vulnerable organs subject-
ed to serious damage via metabolic or inflammatory disturbances. In fact, neurologic complications, such as sepsis-associated encephalopathy, frequently occur early in septic patients and are associated with even higher mortality and long-term complications such as cognitive dysfunction [13-18]. Studies have further demonstrated that septic encephalopathy and polyneuropathy occur in 70% of septic patients [13]. Interestingly, despite the incidence of SS-associated central nervous system (CNS) complication and relevant prognosis are clear; however, the underlying mechanisms remain a heatedly debated issue [14-18].

Previous studies have exhibited that host’s defense mechanisms, including uncontrolled overwhelming inflammatory response and hyperactivity of immune system in response to infectious stimulation along with complement cascade which not only fight against bacterial infection but also attack and damage the host’s organs, are the major contributors of high mortality in SS patients [9, 18-22]. Interestingly, some studies have revealed that systemic inflammation appears to play a key role in both altering the blood-brain barrier permeability and amplifying the inflammatory response [15]. These aforementioned issues [9, 15, 18-22] raised the hypothesis that early targeting on the inflammatory signaling might have a potentially therapeutic impact on protecting the brain organ failure against SS.

Intriguingly, plentiful studies have shown that the mesenchymal stem cell (MSC)-derived exosomes which have capacities of angiogenesis, immunomodulation, anti-inflammation and the paracrine effect protect the organ function following injury in preclinical studies [23-28]. Our previous studies [15, 24, 29, 30] have shown that adipose-derived MSC (i.e., ADMSC)-derived exosomes effectively protected the organs from sepsis or ischemia-reperfusion damage mainly through regulating the inflammatory-oxidative signaling axis. Accordingly, we proposed that ADMSC-derived exosomes might have therapeutic benefit on protecting the brain organ from SS induced injury.

Materials and methods

Ethics

All animal procedures were approved by the Institute of Animal Care and Use Committee at Kaohsiung Chang Gung Memorial Hospital (Affidavit of Approval of Animal Use Protocol No. 2016032201) and performed in accordance with the Guide for the Care and Use of Laboratory Animals.

Animals were housed in an Association for Assessment and Accreditation of Laboratory Animal Care International (AAALAC; Frederick, MD, USA)-approved animal facility in our hospital with controlled temperature and light cycles (24°C and 12/12 light cycle).

Procedure for SS induction by cecal ligation and puncture (CLP)

The procedure and protocol of SS induction have been described in our previous reports [11, 12, 31]. In detail, Pathogen-free, adult male Sprague-Dawley (SD) rats weighing 320-350 g (Charles River Technology, BioLASCO, Taiwan) were utilized in the present study. Rats were anesthetized with inhalational 2.0% isoflurane and placed supine on a warming pad at 37°C with the abdomen shaved. Under sterile conditions, the abdominal skin and muscle were opened and the cecum exposed in all groups. In the sham-control animals, the abdomen was then closed and the animal was allowed to recover from anesthesia. In the experimental CLP groups, the cecum was ligated with prolene suture over its distal portion (i.e., distal ligation) and the cecum distal to the ligature was punctured twice with an 18# needle to allow the cecal contents to be expressed intraperitoneally, as previously described [11, 12, 31]. The wound was closed and the animal was allowed to recover from anesthesia.

Isolation of ADMSCs

For preparation of allogenic ADMSCs, additional 20 SD rats were utilized in the present study. Adipose tissue surrounding the epididymis was dissected, excised and prepared based on our recent reports [11, 12, 31]. Briefly, the tissue was carefully cut into pieces < 1 mm³ using a pair of sharp, sterile surgical scissors. Sterile saline (37°C) was added to the homogenized adipose tissue in a ratio of 3:1 (saline: adipose tissue), followed by the addition of stock collagenase solution to a final concentration of 0.5 units/mL. The centrifuge tubes with the contents were placed and secured on a Thermaline shaker and incubated with constant agitation...
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for 60 minutes at 37°C. After 40-45 minutes of incubation, the contents were triturated with a 25-mL pipette for 3 minutes. The contents of the flask were transferred to 50 mL tubes after digestion, followed by centrifugation at 600 g for 5 minutes at room temperature. The cell pellet was resuspended in 40 mL saline and then centrifuged again at 600 g for 5 minutes at room temperature. After being resuspended again in 5-mL saline, the cell suspension was filtered through a 100 μm filter into a 50-mL conical tube to which 2 mL of saline was added to rinse the remaining cells through the filter. The flow-through was pipetted into a new 50-mL conical tube through a 40-μm filter. The tubes were centrifuged for the third time at 600 g for 5 minutes at room temperature. The cells were resuspended in saline. An aliquot of cell suspension was then removed for cell culture in Dulbecco’s modified Eagle’s medium (DMEM)-low glucose medium containing 10% FBS for 14 days. Approximately 2.0-3.0 × 10⁶ ADMSCs were collected from each rat.

**Exosomes preparation for SS treatment**

The preparation, isolation and dosage of ADMSC-derived exosomes have been described in detail in our previous report [26]. Briefly, the exosomes were isolated from the culture medium of ADMSCs and pooled for protein separation and characterization, as well as western blot analysis. The proteins in Dulbecco’s modified Eagle’s medium (DMEM)-low glucose medium containing 10% FBS for 14 days. Approximately 2.0-3.0 × 10⁶ ADMSCs were collected from each rat.

**Animal grouping, therapeutic strategy, blood sampling and time intervals of euthanized animals for individual study**

The adult male SD rats (n = 72) were equally categorized into sham-operated control (SC, i.e., only opened the abdomen, then closed the muscle and skin layers), SS + 500 μ normal saline and SS + allogenic ADMSC-derived exosomes (AMSC<sup>EXP</sup>) (i.e., 100 μg intravenous administration 3 h after CLP) groups. The exosome dosage administered was described in our previous report [24].

Blood samples were drawn at time intervals of 6, 16, 24 and 72 h for assessment of circulating levels of immune cells, cellular apoptosis and inflammatory biomarkers. Additionally, 6 animals in each group at these four time points were euthanized and brain tissues were harvested for individual study.

**Isolation of splenic mononuclear cells for assessment of immune cells**

Mononuclear cells in the spleen were isolated by homogenization of the spleen using a Tenbroeck tissue grinder followed by passage through a 0.4-mm-pore-size cell strainer to obtain a single cell suspension. These mononuclear cells were then suspended in RPMI and separated by Ficoll-paque Plus (GE Healthcare) for identification of immune cells.

**Flow cytometric quantifications of inflammatory and apoptotic cells in circulation, and immune cells in circulation and spleen**

The procedure and protocol of flow cytometric analysis which was performed to identify and characterize immune cells, apoptotic and inflammatory cell surface markers have been described in our previous reports [12, 24]. Briefly, peripheral blood mononuclear cells and splenocytes (1.0 × 10⁶ cells for each population) were triple-stained with FITC-anti-CD3 (BioLegend), PE-anti-CD8α (BD Bioscience), and PE-Cy™5 anti-CD4 (BD Bioscience). To identify CD4+CD25+Foxp3+ Tregs, peripheral blood mononuclear cells and splenocytes were triple-stained with Alexa Fluor 488-anti-CD25 (BioLegend), PE-anti-Foxp3 (BioLegend), and PE-Cy5 anti-CD4 (BD Bioscience) using the Foxp3 Fix/Perm buffer set according to the manufacturer’s protocol. The numbers of CD3+/CD4+ helper T cells, CD3+/CD8+ cytotoxic T cells, and CD4+CD25+Foxp3+ Tregs were analyzed using flow cytometry (FC500, Beckman Coulter).

The percentages of viable and apoptotic cells were determined by flow cytometry using double staining of annexin V and propidium iodide.
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(Pi). This is a simple and popular method for the identification of apoptotic cells (i.e., early [annexin V+/Pi-] and late [annnexin V+/Pi+] phases of apoptosis).

Additionally, the numbers of inflammatory cells in circulation [i.e., CD11b/c, LyG6] were assessed using the flow cytometric method.

**Analysis of circulating and cerebrospinal fluid (CSF) levels of inflammatory cytokines**

Blood samples and CSF were drawn from rats in each group (n = 6/each time interval) at time points of 6, 24 and 72 h after SS induction and stored at -80°C until analyses of tumor necrosis factor (TNF)-α and interleukin (IL)-6 were performed in batches at the end of the experiment. Serum TNF-α and IL-6 concentrations were assessed in duplicate with a commercially available ELISA kit (R&D Systems, Minneapolis, MN, USA). Intra-individual variabilities in TNF-α and IL-6 levels were assessed in each group with the mean intra-assay coefficient of variance < 1.8%.

**Immunohistochemical (IHC) and immunofluorescent (IF) staining**

The procedure and protocol for IHC and IF staining have previously been reported in detail [12, 31, 32]. For IHC and IF staining, rehydrated paraffin sections were first treated with 3% H2O2 for 30 minutes and incubated with ImmunoBlock reagent (BioSB, Santa Barbara, CA, USA) for 30 minutes at room temperature. Sections were then incubated with primary antibodies specifically against CD14 (1:200, Proteintech), F4/80 (1:100, Santa Cruz), glial fibrillary acidic protein (GFAP) (1:500, Dako) and MMP-9 (1:200, Themo Fisher Scientific) while sections incubated with irrelevant antibodies served as controls. Three kidney sections from each rat were analyzed. For quantification, three randomly selected HPFs (200× for IHC; 400× for IF) were analyzed in each section. The mean number of positively-stained cells per HPF for each animal was then determined by summation of all numbers divided by 9.

**Western blot analysis**

The procedure and protocol for Western blot analysis have been described in our previous reports [12, 31, 32]. Briefly, equal amounts (50 μg) of protein extract were loaded and separated by SDS-PAGE using acrylamide gradients. After electrophoresis, the separated proteins were transferred electrophoretically to a polyvinylidene difluoride (PVDF) membrane (Amer sham Biosciences, Amersham, UK). Nonspecific sites were blocked by incubation of the membrane in blocking buffer [5% nonfat dry milk in T-TBS (TBS containing 0.05% Tween 20)] overnight. The membranes were incubated with the indicated primary antibodies [caspase 3 (1:1000, Cell Signaling, Danvers, MA, USA), Poly (ADP-ribose) polymerase (PARP) (1:1000, Cell Signaling, Danvers, MA, USA), high mobility group box protein 1 (HMGB1) (1:1000, Cell Signaling, Danvers, MA, USA), toll-like receptor 2 (TLR-2) (1:1000, Abcam, Cambridge, MA, USA), TLR-4 (1:1000, Novus Biologicals, Littleton, MA, USA), myeloid differentiation primary response 88 (MyD88) (1:1000, Abcam, Cambridge, MA, USA), NADPH oxidase (NOX)-1 (1:1500, Sigma, St. Louis, MO, USA), NOX-2 (1:750, Sigma, St. Louis, MO, USA), interleukin (IL)-1β (1:1000, Cell Signaling, Danvers, MA, USA), matrix metalloproteinase (MMP)-9 (1:1000, Abcam, Cambridge, MA, USA) and actin (1: 10000, Chemicon, Billerica, MA, USA)] for 1 hour at room temperature. Horseradish peroxidase-conjugated anti-rabbit immunoglobulin IgG (1:2000, Cell Signaling, Danvers, MA, USA) was used as a secondary antibody for one-hour incubation at room temperature. The washing procedure was repeated eight times within one hour. Immunoreactive bands were visualized by enhanced chemiluminescence (ECL; Amersham Biosciences, Amersham, UK) and exposed to Biomax L film (Kodak, Rochester, NY, USA). For quantification, ECL signals were digitized using Labwork software (UVP, Waltham, MA, USA). The full gel images of western blot were available in the Figure S1.

**Assessment of oxidative stress**

The procedure and protocol for assessing the protein expression of oxidative stress have been detailed in our previous reports [12, 31, 32]. The Oxyblot Oxidized Protein Detection Kit was purchased from Chemicon, Billerica, MA, USA (S7150). DNPH derivatization was carried out on 6 μg of protein for 15 minutes according to the manufacturer's instructions. One-dimen-
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Measurement of circulating and CSF levels of inflammatory cytokines by ELISA at time intervals of 6, 24, and 72 h after SS induction (Figure 4)

The circulating and CSF levels of TNF-α and IL-6, two indicators of acute inflammatory cytokines, were significantly increased in group 2 than in groups 1 and 3, and significantly increased in group 3 than in group 1 at time intervals of 6, 24 and 72 h.

Inflammatory cell infiltration in brain tissues by time intervals of 6 and 72 h after SS induction (Figures 5 and 6)

The cellular expressions of F4/80 and CD14, two indicators of inflammation, at time points of 6 and 72 h after SS induction, were significantly higher in group 3 than in group 1 (Figure 5). Additionally, cellular expressions of MMP-9 and GFAP, another two indicators of inflammation, exhibited an identical pattern of CD14 among the three groups at these two time points (Figure 6).

Cellular expressions of brain-damaged biomarkers at time intervals of 6 and 72 h after SS induction (Figure 7)

The cellular expressions of AQP4 (i.e., brain edema index) and γ-H2AX (DNA-damaged indicator) were significantly increased in group 2 as compared to those of groups 1 and 3 and significantly higher in group 3 than in group 1.

Protein levels of apoptotic and brain damaged released inflammatory biomarkers at time intervals of 6 and 72 h after SS induction (Figure 8)

The protein expressions of cleaved caspase 3 and cleaved PARP, two indicators of apoptotic

Statistical analysis

Quantitative data are expressed as means ± SD. Statistical analysis was adequately performed by ANOVA followed by Bonferroni multiple-comparison post hoc test. Statistical analysis was performed using SPSS statistical software for Windows version 22 (SPSS for Windows, version 22; SPSS, IL, USA). A value of $P < 0.05$ was considered as statistically significant.

Results

Circulating levels of inflammatory and apoptotic cells at time points of 6, 16, 24 and 72 h after SS induction (Figure 1)

The circulating numbers of Ly6G+ and CD11b/c+ cells, two indicators of inflammation, were significantly higher in group 2 (SS) than in group 1 (SC) and group 3 (SS + AMSC<sub>EXO</sub>), and significantly higher in group 3 than in group 1 at time intervals of 6, 16, 24 and 72 h. Additionally, the circulating levels of early (i.e., annexin V+/PI-) and late [i.e., annexin V+/PI+] cellular apoptosis also exhibited an identical pattern of inflammatory cells at time intervals of 6, 16, 24 and 72 h.

Serial changes of immune cells in circulation and spleen (Figures 2 and 3)

The circulating and splenic levels of CD3+/CD4+ cells and CD3+/CD8+ cells, two indices of immune reaction, were significantly increased in group 2 than in groups 1 and 3, and significantly increased in group 3 than in group 1 at time points of 6, 16, 24, and 72 h (Figure 2). On the other hand, the circulating and splenic levels of T-reg+ cells, an indicator of immunomodulation, were progressively increased from groups 1 to 3, highlighting an intrinsic response to inflammatory reaction in SS than the corresponding MSC<sub>EXO</sub>-enhanced treatment for purpose of attenuating the overwhelming immune response in SS (Figure 3).
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Biomarkers, were significantly higher in group 2 than in groups 1 and 3, and significantly higher in group 3 than in group 1 at time intervals of 6 and 72 h after SS induction.

Additionally, protein expression of high mobility group box protein 1 (HMGB1), a component of damage-associated molecular patterns (DAMPs), displayed an identical pattern of apoptosis among the three groups at these two-time intervals.

The protein expressions of inflammatory biomarkers at time intervals of 6 and 72 h after SS induction (Figures 9 and 10)

The protein expressions of TLR-2, TLR-4, and MyD88, three indices of inflammation, were
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significantly higher in group 2 than in groups 1 and 3, and significantly higher in group 3 than in group 1 at time intervals of 6 and 72 h after SS induction (Figure 9). Furthermore, the protein expressions of IL-1β, MMP-9, TNF-α and NF-κB, another four indicators of inflammatory biomarkers, displayed an identical pattern of MyD88 among the three groups at these two-time intervals (Figure 10).

The protein expressions of oxidative-stress biomarkers at time intervals of 6 and 72 h after SS induction (Figure 11)

The protein expressions of NOX-1, NOX-2 and oxidized protein, three indicators of oxidative stress, were significantly increased in group 2 than in groups 1 and 3, and significantly increased in group 3 than in group 1 at time intervals of 6 and 72 h after SS induction.
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Figure 3. Circulatory and splenic levels of T-reg+ cells at time intervals of 6, 16, 24 and 72 h after SS induction. (A, B) Number of T-reg+ cells in circulation (A) and spleen (B) ay 6 h, *vs. other groups with different symbols (†, ‡), P < 0.001. (C, D) Number of T-reg+ cells in circulation (C) and spleen (D) ay 16 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. (E, F) Number of T-reg+ cells in circulation (E) and spleen (F) ay 24 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. (G, H) Number of T-reg+ cells in circulation (G) and spleen (H) ay 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. All statistical analyses were performed by one-way ANOVA, followed by Bonferroni multiple comparison post hoc test (n = 6 for each group). Symbols (*, †, ‡) indicate significance (at 0.05 level). SC = sham-operated control; SS = sepsis syndrome; AMSC^EXO = allogenic adipose-derived mesenchymal stem cells-derived exosomes.

Figure 4. Measurement of circulating and CSF levels of inflammatory cytokines by ELISA at time intervals of 6, 24, and 72 h after SS induction. (A, B) Level of tumor necrosis factor (TNF)-α in circulation (A) and Cerebrospinal fluid
(CSF) (B) at 6 h, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). (C, D) Level of TNF-\(\alpha\) in circulation (C) and CSF (D) at 24 h, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). (E, F) Level of TNF-\(\alpha\) in circulation (E) and CSF (F) at 72 h, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). (G, H) Level of interleukin (IL)-6 in circulation (G) and CSF (H) at 6 h, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). (I, J) Level of IL-6 in circulation (I) and CSF (J) at 24 h, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). (K, L) Level of IL-6 in circulation (K) and CSF (L) at 72 h, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). All statistical analyses were performed by one-way ANOVA, followed by Bonferroni multiple comparison post hoc test (\(n = 6\) for each group). Symbols (*, †, ‡) indicate significance (at 0.05 level). SC = sham-operated control; SS = sepsis syndrome; AMSC\(^{EXO}\) = allogenic adipose-derived mesenchymal stem cells-derived exosomes.

**Figure 5.** Inflammatory cell infiltration in brain tissues by time intervals of 6 and 72 h after SS induction. A-C. Illustrating the immunofluorescent (IF) microscopic finding (400×) for identification of cellular expression of F4/80 at 6 h (green color). D. Analytical result of number of F4/80\(^+\) cells, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). E-G. Illustrating the IF microscopic finding (400×) for identification of cellular expression of CD14 at 6 h (green color). H. Analytical result of number of CD14\(^+\) cells, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). I-K. Illustrating the IF microscopic finding (400×) for identification of cellular expression of F4/80 at 72 h (green color). L. Analytical result of number of F4/80\(^+\) cells, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). M-O. Illustrating the IF microscopic finding (400×) for identification of cellular expression of CD14 at 72 h (green color). P. Analytical result of number of CD14\(^+\) cells, \(^*\) vs. other groups with different symbols (†, ‡), \(P < 0.0001\). Blue color indicated the nuclei stained by DAPI. Scale bars in lower right corner represent 20 \(\mu m\). All statistical analyses were performed by one-way ANOVA, followed by Bonferroni multiple comparison post hoc test (\(n = 6\) for each group). Symbols (*, †, ‡) indicate significance (at 0.05 level). SC = sham-operated control; SS = sepsis syndrome; AMSC\(^{EXO}\) = allogenic adipose-derived mesenchymal stem cells-derived exosomes.

**Discussion**

This study which investigated the therapeutic impact of ADMSC-derived exosomes on protecting the brain from SS-induced brain damage yielded several striking implications. First, the results of this current study showed that original source of abdominal sepsis elicited not
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Figure 6. Inflammatory cell biomarkers in brain tissues by time intervals of 6 and 72 h after SS induction. A-C. Illustrating the immunofluorescent (IF) microscopic finding (400×) for identification of cellular expression of glial fibrillary acidic protein (GFAP) at 6 h (red color). D. Analytical result of number of GFAP+ cells, *vs. other groups with different symbols (†, ‡), P < 0.0001. Scale bars in lower right corner represent 20 µm. E-G. Illustrating the immunohistochemical (IHC) stain for identifying the cellular expression of matrix metalloproteinase (MMP)-9 (gray color). H. Analytical result of number of MMP-9+ cells, *vs. other groups with different symbols (†, ‡), P < 0.0001. Scale bars in lower right corner represent 50 µm. I-K. Illustrating the IF microscopic finding (400×) for identification of cellular expression of GFAP at 72 h (red color). L. Analytical result of number of GFAP+ cells, *vs. other groups with different symbols (†, ‡), P < 0.0001. Scale bars in lower right corner represent 20 µm. M-O. Illustrating the IHC stain for identifying the cellular expression of MMP-9 (gray color). P. Analytical result of number of MMP-9+ cells, *vs. other groups with different symbols (†, ‡), P < 0.0001. HPF = high-power field. Blue color indicated nuclei stained by DAPI. Scale bars in lower right corner represent 50 µm. All statistical analyses were performed by one-way ANOVA, followed by Bonferroni multiple comparison post hoc test (n = 6 for each group). Symbols (*, †, ‡) indicate significance (at 0.05 level). SC = sham-operated control; SS = sepsis syndrome; AMScEXO = allogenic adipose-derived mesenchymal stem cells-derived exosomes.

only circulatory inflammatory reaction but also brain organ inflammatory response (i.e., a “sepsis victim” as likely as other organs), highlighting that SS affected systemically and involved multiple innocent organs. Second, the time points of measuring inflammatory biomarkers demonstrated that the vigorous immune-inflammatory reactions emerged at a peak level as early as a few hours (i.e., ≤ 6 h) after SS induction that were not only in circulatory but also extended to brain organ and CSF. Third, prompt administration of ADMSC-derived exosomes effectively suppressed the systemic immune-inflammatory responses and protected the brain organ from SS-induced damage.

Whatever the original source of infection is, vast previous studies have shown that sepsis can spread into circulation causing SS which ultimately leads to multiple organ failure and unacceptably high mortality [1-5, 9, 10]. A principal finding in the present study was that the
immune-inflammatory reactions were identified not only in the circulation, but also in spleen as well as in the brain tissue and CSF, indicating that an abdominal origin of SS could disseminate throughout the human body. Our results using an experimental study support the findings from previous clinical trials and observational studies [1-5, 9, 10].

Intriguingly, even though the sepsis-associated encephalopathy and the incidence of SS-associated metabolic disturbance and CNS complication as well as its unfavorable prognostic outcome have been earnestly clarified, yet regrettably the underlying mechanisms remain a heatedly debated topic [14-18]. An essential finding in the present study was that not only the cytokine level but also molecular-cellular and protein levels of inflammatory biomarkers were identified to markedly increase in CNS (i.e., expressed in brain tissues and CSF) in SS animals. Our findings, could, at least in part, explain the phenomena of CNS complications after sepsis that have been revealed by previ-
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Our recent study has demonstrated that DAMP inflammatory axis was markedly upregulated and participated in brain damage in brain-dead animals [33]. The most important finding in the present study was that the levels of a DAMP component (i.e., HMGB1) and downstream inflammatory biomarkers, i.e., TLR-2, TLR-4, MYD88, IL-1β, TNF-α, NFκB and MMP-9, were substantially higher in brain tissue from SS animals than the SC. In this way, our findings, in addition to being consistent with the results of our recent study [33], could powerfully explain why the brain is frequently as an innocent victim seriously damaged by SS, highlighting that not only the brain is an organ failing in sepsis but also sepsis overwhelmingly...

Figure 8. Protein levels of apoptotic and brain-damaged released biomarkers at time intervals of 6 and 72 h after SS induction. A. Protein expression of cleaved caspase 3 (c-Casp-3) at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. B. Protein expression of cleaved Poly (ADP-ribose) polymerase (c-PARP) at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. C. Protein expression of high mobility group box protein 1 (HMGB1) at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. D. Protein expression of c-Casp-3 at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. E. Protein expression of c-PARP at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. F. Protein expression of HMGB1 at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. All statistical analyses were performed by one-way ANOVA, followed by Bonferroni multiple comparison post hoc test (n = 6 for each group). Symbols (*, †, ‡) indicate significance (at 0.05 level). SC = sham-operated control; SS = sepsis syndrome; ADMSC = allogenic adipose-derived mesenchymal stem cells-derived exosomes.
influences all inflammatory processes on a variety of pathophysiologic levels, thus contributes to the initiation and propagation of septic processes [17]. Of particular importance was that these molecular-cellular perturbations in circulatory and splenic levels and in brain tissue as well as in CSF were remarkably suppressed by ADMSC-derived exosomes. Intriguingly, ours [24, 26, 29, 30] and other studies [15, 23, 25, 27, 28] have previously shown that MSC-derived exosomes protected the organs from ischemia/ischemia-reperfusion injury or from sepsis-induced damage mainly through immunomodulation and suppression of the inflammatory reaction and oxidative stress. In this way, the results of the current study, in addition to being consistent with the findings of the previous studies [15, 23-30], highlight that these preclinical findings convey a significant clinical information that ADMSC-derived exosomes may have a potential therapeutic impact on the setting of SS complicated with CNS involvement, especially in patients who are refractory to conventional treatment.

Study limitations

This study has limitations. First, the neurological function was not examined in the present

Figure 9. Protein levels of upstream inflammatory biomarkers at time intervals of 6 and 72 h after SS induction. A. Protein expression of toll-like receptor (TLR)-2 at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. B. Protein expression of TLR-4 at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. C. Protein expression of myeloid differentiation primary response 88 (MyD88) at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. D. Protein expression of TLR-2 at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. E. Protein expression of TLR-4 at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. F. Protein expression of MyD88 at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. All statistical analyses were performed by one-way ANOVA, followed by Bonferroni multiple comparison post hoc test (n = 6 for each group). Symbols (*, †, ‡) indicate significance (at 0.05 level). SC = sham-operated control; SS = sepsis syndrome; AMSC\textsuperscript{EXO} = allogenic adipose-derived mesenchymal stem cells-derived exosomes.
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study. Thus, we did not provide the information regarding the impairment of neurological function in setting of SS and the recovery of the neurological status after ADMSC-derived exosome therapy. Second, the study period was relatively short. As a consequence, the long-term outcome of ADMSC-derived exosome therapy on protecting the brain against SS injury remains to be clarified.

In conclusion, allogenic ADMSC-derived exosome therapy significantly protected the brain from SS-induced damage.

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Disclosure of conflict of interest

None.

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Figure 10. The protein expressions of downstream inflammatory biomarkers at time intervals of 6 and 72 h after SS induction. A. Protein expression of nuclear factor (NF)-κB at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. B. Protein expression of matrix metalloproteinase (MMP)-9 at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. C. Protein expression interleukin (IL)-1β at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. D. Protein expression of tumor necrosis factor (TNF)-α at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. E. Protein expression of NF-κB at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. F. Protein expression of MMP-9 at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. G. Protein expression IL-1β at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. H. Protein expression of TNF-α at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. All statistical analyses were performed by one-way ANOVA, followed by Bonferroni multiple comparison post hoc test (n = 6 for each group). Symbols (*, †, ‡) indicate significance (at 0.05 level). SC = sham-operated control; SS = sepsis syndrome; AMSCEXO = allogenic adipose-derived mesenchymal stem cells-derived exosomes.
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Figure 11. Protein expressions of oxidative-stress biomarkers at time intervals of 6 and 72 h after SS induction. A. Protein expression of NOX-1 at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. B. Protein expression of NOX-2 at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. C. Oxidized protein expression at 6 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. (Note: left and right lanes shown on the upper panel represent protein molecular weight marker and control oxidized molecular protein standard, respectively). MW = molecular weight; DNP = 1,3 dinitrophenylhydrazone. D. Protein expression of NOX-1 at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. E. Protein expression of NOX-2 at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. F. Oxidized protein expression at 72 h, *vs. other groups with different symbols (†, ‡), P < 0.0001. All statistical analyses were performed by one-way ANOVA, followed by Bonferroni multiple comparison post hoc test (n = 6 for each group). Symbols (*, †, ‡) indicate significance (at 0.05 level). SC = sham-operated control; SS = sepsis syndrome; AMSCEXO = allogenic adipose-derived mesenchymal stem cells-derived exosomes.

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Figure S1. A-D. Illustrating the raw materials of Western blotting (i.e., including the whole films/membranes).