Combined therapy of photobiomodulation and adipose-derived stem cells synergistically improve healing in an ischemic, infected and delayed healing wound model in rats with type 1 diabetes mellitus

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ABSTRACT

Objective We assessed the therapeutic effects of photobiomodulation (PBM) and adipose-derived stem cell (ADS) treatments individually and together on the maturation step of repairing of a delayed healing wound model in rats with type 1 diabetes mellitus (DM1).

Research design and methods We randomly assigned 24 rats with DM1 to four groups (n=6 per group). Group 1 was the control (placebo) group. In group 2, allograft human ADSs were transplanted. Group 3 was subjected to PBM (wavelength: 890 nm, peak power output: 80 W, pulse frequency: 80 Hz, pulsed duration: 180 ns, duration of exposure for each point: 200 s, power density: 0.001 W/cm², energy density: 0.2 J/cm²) immediately after surgery, which continued for 6 days per week for 16 days. Group 4 received both the human ADS and PBM. In addition, we inflected an ischemic, delayed healing, and infected wound simulation in all of the rats. The wounds were infected with methicillin-resistant Staphylococcus aureus (MRSA).

Results All three treatment regimens significantly decreased the amount of microbial flora, significantly increased wound strength and significantly modulated inflammatory response and significantly increased angiogenesis on day 16. Microbiological analysis showed that PBM+ADS was significantly better than PBM and ADS alone. In terms of wound closure rate and angiogenesis, PBM+ADS was significantly better than the PBM, ADS and control groups.

Conclusions Combination therapy of PBM+ADS is more effective that either PBM or ADS in stimulating skin injury repair, and modulating inflammatory response in an MRSA-infected wound model of rats with DM1.

INTRODUCTION

Type 1 diabetes mellitus (DM1) is a chronic illness that results from inadequate insulin production by the pancreas. In 2017, there were 451 million individuals worldwide who suffered from DM. The incidence of DM1 is estimated to increase to nearly 700 million
by 2045. Approximately 50% of all persons living with DM are unidentified.2

People who suffer from DM are at greater risk for diabetic foot ulcers (DFUs). DFUs are the result of concurrent activities of many causative sources. Persistent hyperglycemia, peripheral neuropathy, ischemia from peripheral vasculopathy, immunosuppression and infection are the major pathological elements that lead to foot deformities and DFUs.3 Of note, any DFU is assumed to have blood vessel deficiencies, even without direct verification.4 In addition to metabolic aberrations, DM causes overproduction of mitochondrial superoxide and reactive oxygen species (ROS) in endothelial cells, which further leads to reduced angiogenesis and ischemia.5

Lower limb amputation (LLA) occurs in 5%-24% of patients with DFUs within 180-545 days after the initial diagnosis.6 Worldwide, DFUs are the most common complications that lead to huge financial problems for the patients, their relatives and the public.7 The total clinical cost for managing diabetic lower limb complications in the USA is approximately US$10.5 billion in addition to the cost of DM management.8

Approximately 50% of LLA occur in patients with DM, and most are from microbial DFU.9 The prevalence of methicillin-resistant Staphylococcus aureus (MRSA) in infected DFU is 15%-30%.10 Among microbes S. aureus is the most prevalent. Overuse of antibiotics to combat infected diabetic ulcers have led to an explosion of antibiotic resistance with delayed macrophage response, which poses a serious threat to global public health.11

DFU is the main medical threat due a lack of knowledge about its pathophysiology.12 Weakened wound repair in DM is identified by reduced angiogenesis, decreased endothelial progenitor cell recruitment, reduced fibroblast and keratinocyte proliferation and migration and delayed macrophage response.12-15 The capability to quickly repair the wound is demanding and is the crucial aim for a cure for DFU.14 Both protective and curative measures consist of the use of bactericidal mediators, wet absorbent and bioengineered gauzes, vacuum-assisted closure, Regranex PDGF gel and unloading to stimulate the repair process.15 However, none of these measures show any distinct advantage over the others; the cure of chronic wounds and DFU remains problematic and prolonged.14 15 Thus, new approaches are urgently needed to combat infection and stimulate tissue repair in diabetic wounds. Concentrated efforts on the effectiveness of prognostic laboratory experiments, technologies for new cures and selective and focused approaches are required.4

Mesenchymal stem cells (MSCs) can effectively enhance repair of wound. The ability of MSCs to differentiate into various cell lines and their low immunogenicity make them highly attractive for therapeutic use in DFUs. MSCs stimulate cell migration, new blood vessel formation, re-epithelialization and new wound bed formation and maturation. In addition, they reduce the inflammatory response, enhance wound contraction, and can improve healing.16

Adipose-derived stem cells (ADs) are great replacements of bone marrow (BM) MSCs because they can be simply harvested from adipose tissue and can grow in a culture system. Published investigational probes have indicated that ADs could improve wound repair via boosting re-epithelialization and new wound bed formation. They modulate the inflammatory reaction, have an anti-apoptotic impact and secrete angiogenic growth factors.17

Probes using animal simulation18 and studies in humans19 have been performed to display the probable favorable impacts of ADs in wounds and ischemic organs in animals and in patients who suffer from DM. Despite the current achievements, several chief obstacles remain before they can be effectively used to benefit diabetic ADs in wound repair. First, the aberrant in vivo milieu of people with DM adversely affects the therapeutic biological action of the ADs on wound repair.20 Second, the systemic deficiency of the healing potential of diabetic ADs in vivo.21 Third, ADs is physiologically damaged.22 Finally, ADs cannot provide a blood vessel complex. Consequently, these diabetic ADs are weak in stimulating new blood vessel formation and soft tissue repair.23

Photobiomodulation (PBM) decreases pain, and inflammation, and enhances injury repair, and prevents cell and tissue damage.24 We believe that PBM could overcome these ADs deficiencies and synergize its positive effects in stimulating tissue repair. PBM has long been used as a means to increase circulation and improve tissue repair by enhancing new blood vessel formation through increases in vascular endothelial growth factor release and hypoxia-induced factor-1α expression.25 PBM has been shown to improve the angiogenic impact of ADs by increasing its viability and encouraging the release of cytokines in the cutaneous flap,26 ischemic limbs27 and injured skin.28

More than 100 recognized factors are involved in the insufficient skin injury repair observed in people with DM.29 The therapeutic use of a combination of beneficial mediators and biomodulators appears to be of benefit in non-healing wounds; they are likely to show a synergistic effect and can improve the success of a cure for different wound simulations. In recent experiments, Bayat group have reported the capability of the combination of PBM and conditioned media (CM) from human bone marrow mesenchymal stem cells (hBM-MSCs) to stimulate repair of weakened wounds in a rat model of streptozotocin (STZ)-induced DM1. The outcomes of these experiments showed that PBM+CM prompted anti-inflammatory and angiogenic actions, and accelerated skin injury repair in a DM1 simulation of an MRSA-infected skin injury.30 The combination of CM and PBM revealed a synergistic influence.31 Here, we intend to assess the impact of PBM and ADs treatments, alone and in combination, on the maturation step of a repair course of ischemic, delayed healing, and infected wound simulations in rats with DM1. The combined application of PBM and ADs could advance the repairing course and assist with healing severe cases of DFUs in people with DM.
Figure 1  A photo of the wound, photobiomodulation (PBM) target points, and adipose-derived stem cells (ADS) injection points.

MATERIALS AND METHODS

Animals and study design

We randomly assigned 24 male Wistar rats aged 3 months into four groups (n=6 per group). Group 1 was the control (placebo) group. In group 2, allograft human ADS were transplanted. Group 3 received PBM treatment. The fourth group received an allograft of human ADS and PBM treatment. DM1 was induced in all rats. In addition, an ischemic, delayed healing and infected wound simulation was inflicted on all of the rats. We performed microbiological examinations, wound closure rate, wound strength and histological and stereological tests on days 0, 4, 8, 12 and 16. We considered day 4 to be the inflammation step, day 8 to be the proliferation step and days 12 and 16 were the early and late maturation steps for the skin injury repair.

Isolation, expansion and immunophenotyping of human hypodermis ADS

Human hypodermis adipose tissue was donated by a healthy adult donor (aged 35–40 years) who underwent cosmetic mammoplasty surgery. The patient provided informed consent for the use of her tissue. Approximately 5 cc of adipose tissue was manually minced, washed with phosphate buffered saline (PBS) supplemented with 100 U/mL penicillin+100 µg/mL streptomycin and subjected to digestion with 0.1% collagenase type I solution. Then, the cell pellets were suspended in Dulbecco’s Modified Eagle’s medium (DMEM)+20% fetal bovine serum (FBS) and then seeded in T-75 flasks with DMEM+20% FBS, and the above-mentioned antibiotics. We performed flow cytometry analysis to characterize the ADS cells for MSC markers, as reported previously.32,33

ADS transplantation

At 24 hours after surgery, we mixed 1×10^6 passage-4 ADS cells34 in 300 µL PBS.31 The mixture was immediately injected intradermally by an insulin syringe into eight sites around each wound at a 4–5 mm distance from the wound’s margin (figure 1).

Induction of DM1

DM1 was induced through injection of STZ (40 mg/kg, intraperitoneal).31 DM1 was validated as a blood sugar level higher than 250 mg/dL. All rats with diabetes were kept for 1 month to ensure that DM1 was established in the animals.31

Clinical examinations

We monitored the rats’ body weights and blood sugar levels throughout the experiment.

Surgery

The rats were anesthetized by intramuscular injections of ketamine (50 mg/kg) and xylazine (5 mg/kg). A dorsal, bipedicle skin flap (10×3.5 cm) was generated deep into the skin muscle. A 12 mm full-thickness excisional round wound that included the skin muscle was produced in the midpoint region of the flap by a biopsy punch. A donut-shaped silicone skin holder was fixed around each skin defect with a 04-silk suture (figure 1). Before surgery, all rats were treated with 20 mg/kg ibuprofen every 8–12 hours, which was continued until 5 days after surgery.

Inoculation of MRSA into the wounds and microbiological examination

We used the MRSA strain of S. aureus (ATCC 25923) according to a previously described procedure.31 In summary, a colony of MRSA was prepared for a final concentration of 2×10^8 at 1 cc and a 100 µL aliquot that contained 2×10^7 MRSA was administered topically onto each wound immediately after surgery. Microbiological samples for routine microbiological analyses were taken from the wounds on days 8 and 16. The numbers of bacterial per animal were counted as colony-forming units (CFUs).31

Photobiomodulation

Wounds from the rats in groups 3 and 4 were subjected to PBM immediately after surgery, which continued for 6 days per week for 16 days. The rats were sedated with the anesthetizing drugs (figure 1).31 Since the PBM target area (wound area and adjacent unwounded skin) was larger than laser probe beam area, we radiated nine
Table 1  Complete specifications for the photobiomodulation

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Dose and unit</th>
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<tbody>
<tr>
<td>Peak power output</td>
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<tr>
<td>Average power</td>
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<tr>
<td>Power density</td>
<td>0.001 W/cm²</td>
</tr>
<tr>
<td>Wavelength</td>
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</tr>
<tr>
<td>Wavelength range of the device</td>
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<tr>
<td>Pulse frequency</td>
<td>80 Hz</td>
</tr>
<tr>
<td>Spot size</td>
<td>1 cm²</td>
</tr>
<tr>
<td>Diameter</td>
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</tr>
<tr>
<td>Pulsed duration</td>
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<td>Duration of exposure for each point</td>
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</tr>
<tr>
<td>Energy density</td>
<td>0.2 J/cm²</td>
</tr>
<tr>
<td>Number of laser shootings in each session</td>
<td>9</td>
</tr>
<tr>
<td>Total energy densities at one session, and total sessions</td>
<td>1.8 and 25.2 J/cm²</td>
</tr>
<tr>
<td>Probe</td>
<td>L07</td>
</tr>
<tr>
<td>Company</td>
<td>MUSTANG 2000, Technica, Russia</td>
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</table>

shootings of the laser probe in each session of PBM therapy in order to cover all target area (a 33 mm×33 mm quadrangular area including wound in the center and adjacent unwounded skin around wound), and to make sure that all the PBM target area could receive almost the same energy density of the PBM. Complete information of PBM protocol is reported in table 1.

Wound closure rate
We photographed the wounds on days 0, 4, 8, 12 and 16, and calculated and compared the wound closure rate in the study groups and at each time point.

Wound closure rate=((original wound area−open area on X day)/original wound area)×100%.35

Wound strength examination
We harvested one 5×50 mm standard sample from the wounds of all the euthanized rats at day 16 after they were euthanized. The samples were placed in a material testing machine. The deformation ratio was 0.166 mm/s. From the load-deformation curve, we calculated the tensiometric properties including bending stiffness (MPa), maximum force (N), stress high load (N/cm²) and energy absorption (J) of the samples.31

Histological and stereological analyses
The sample from euthanized rats were excised, prepared for light histological study and serially sectioned into 5 μm sections. We stained 10 sections by H&E and 5 sections by Mallory’s trichrome staining methods.

Estimation of the cell numbers
The physical dissector method was used to determine the numerical density (Nv) of the neutrophils, macrophages, and fibroblasts, as follows:

\[
Nv = \frac{\sum Q}{(h \times a/f \times \sum p)}
\]

where Nv is numerical density; ΣQ is number of nuclei; h is height of the dissector; a/f is counting frame area; Σp is number of counting frames in all fields.

\[
N \times V = Nv \times \Sigma Q / \Sigma p 
\]

where Nv is numerical density; V is final total volume.35

Estimation of vascular length=2ΣQ/(Σp×a/f)

where 2ΣQ (total number of the vessel profiles counted per rat skin)/ΣP (number of counting frames in all fields (a/f)).36 Vascular length was considered as a biomarker for angiogenesis.

Statistical analysis
Data are shown as mean±SD. We used the t-test, one-way analysis of variance, repeated measurement analysis and the least significant difference (LSD) tests for statistical analyses. A p value of <0.05 was considered to be statistically significant.

RESULTS
Marker expressions
Flow cytometry analysis showed that the ADS cells slightly expressed CD45 (0.8%) and CD11b (0.33%). In addition, the ADS cells completely expressed CD44H (100%) and CD105 (100%) as seen in online supplementary material.

Clinical observations
All p values were related to the LSD test. All of the rats developed clinical evidence of DM1 with a significant increase in blood sugar levels and a decrease in body weight after STZ injection. On day 4, rats in the PBM and PBM+ADS groups had a significant decrease in weight compared with the ADS group (LSD, p=0.006 and 0.016) and the control group (LSD, p=0.020 and 0.046), respectively. Although this downtrend continued for the PBM group until day 16, the PBM+ADS group experienced a weight gain. The combination of PBM+ADS significantly increased body weight compared with PBM alone. Complete information of body weights and glucose levels are shown in table 2, and by a graph in online supplementary material.

Microbial findings
Day 8
PBM+ADS treatment significantly decreased CFU in the wounds compared with ADS treatment (p=0.008) (figure 2).

Day 16
There were significant decreases in CFU in the PBM+ADS, PBM and ADS groups compared with the control group (all p=0.000). Treatment with PBM+ADS was significantly
Table 2  Comparison of body weights and blood sugar levels in the studied groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>Control</th>
<th>PBM</th>
<th>ADS</th>
<th>PBM+ADS</th>
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<tr>
<td>Initial blood sugar (mg/dL)</td>
<td>445±44.96</td>
<td>366.0±62.39</td>
<td>332.86±72.89</td>
<td>403.87±67.47</td>
</tr>
<tr>
<td>Final blood sugar (mg/dL)</td>
<td>460±21.03</td>
<td>425.3±74.85</td>
<td>361.6±58.50</td>
<td>354.14±60.93</td>
</tr>
<tr>
<td>Initial body weight (g)</td>
<td>310.4±12.6</td>
<td>300.75±19.14</td>
<td>301.57±12.2</td>
<td>304.75±9.58</td>
</tr>
<tr>
<td>Final body weight (g)</td>
<td>265.75±15.95*</td>
<td>246.5±14.26*</td>
<td>295.80±14.17</td>
<td>294±16.35</td>
</tr>
</tbody>
</table>

*P<0.05, **p<0.01; Student’s t-test.

Better than the individual PBM (p=0.006) and ADS (p=0.003) treatments (figure 2).

Wound closure rate

Day 4

There were no significant differences in the wound closure rate among the studied groups (figure 3).

Day 8

PBM+ADS (p=0.000), PBM (p=0.000), and ADS (p=0.001) treatments significantly increased wound closure rate compared with the control group. Treatment with PBM+ADS and PBM were significantly better than ADS treatment (both p=0.000). At the same time, PBM+ADS was significantly better than PBM treatment (p=0.000) (figure 3).

Day 12

Treatment with PBM+ADS and PBM significantly increased wound closure rate compared with the control and ADS groups (all, p=0.000) (figure 3).

Day 16

The PBM+ADS and PBM groups had significantly increased wound closure rate compared with the control and ADS groups (all, p=0.000). PBM+ADS treatment was significantly better than the individual PBM treatment (p=0.007) (figure 3).

Wound strength, bending stiffness

PBM+ADS (p=0.000), PBM (p=0.001) and ADS (p=0.003) treatments significantly increased bending stiffness compared with the control group (figure 4A).

Maximum force

PBM+ADS (p=0.000), PBM (p=0.004) and ADS (p=0.035) treatments significantly increased maximum force compared with the control group. PBM+ADS treatment was significantly better than ADS treatment (p=0.020) (figure 4B).

Stress high load

PBM+ADS (p=0.000), PBM (p=0.004) and ADS (p=0.035) treatments significantly increased stress high load compared with the control group. The PBM+ADS treatment was significantly better than the ADS treatment (p=0.020) (figure 4C).

Energy absorption

All treatments significantly increased energy absorption compared with the control group (all, p=0.000). Concurrently, ADS treatment was significantly better than treatment with PBM+ADS and PBM (both, p=0.000) (figure 4D).

Figure 2  Comparison of colony-forming units (CFUs) of methicillin-resistant *Staphylococcus aureus* (MRSA) wounds in the groups by the least significant difference test. **P<0.01; ***p<0.001. ADS, adipose-derived stem cell; PBM, photobiomodulation.

Figure 3  Comparison of wound closure rate from all of the groups by the least significant difference test. **P<0.01; ***p<0.001. ADS, adipose-derived stem cells; PBM, photobiomodulation.
Clinical Care/Education/Nutrition

Figure 4  Comparison of bending stiffness (A), maximum force (B), stress high load (C), and energy absorption (D), of the wounds in the studied groups according to the least significant difference test. *P<0.05; **p<0.01; ***p<0.001. ADS, adipose-derived stem cells; PBM, photobiomodulation.

Stereological findings
In the cases of inflammatory cells (neutrophils and macrophages), all treatments regimes significantly decreased neutrophils and macrophages counts individually and totally (figure 5, panels A, B and C, all p=0.000). For neutrophil count, the results of PBM+ADS and PBM were better than ADS (p=0.001, p=0.01). For macrophage count, the results of PBM+ADS and PBM were better than ADS (p=0.017, p=0.032). For inflammatory cells count, the results of PBM+ADS and PBM were better than ADS (p=0.003, p=0.01). For fibroblast number, PBM+ADS significantly increased fibroblast number compared with PBM, control and ADS groups (p=0.000, p=0.012, p=0.027) (figure 5, panel D). ADS significantly increased fibroblast number compared with PBM (p=0.04). All treatments regimes significantly increased vascular length compared with control group. In case of angiogenesis, we observed significantly more vascular length in PBM+ADS compared with ADS and PBM (p=0.007, p=0.029) (figure 5, panel E).

Histological micrographs of four studied groups which were stained with H&E staining method are shown in online supplementary materials.

Findings of Mallory's trichrome staining
Photos of Mallory's trichrome staining are shown in online supplementary materials. Collagen fibers orientation in control group were mixed, while in treatment
groups were mostly horizontal. Pattern of collagen fibers in control group were mostly reticular and thin, whereas in treatment groups they were mostly fascicular and they seem thicker than collagen fibers of the control group.

**DISCUSSION**

In the current study, we evaluated the impact of PBM and ADS administration alone and together on the maturation step of the repair process in an ischemic, delayed healing and infected wound simulation in rats with DM1.

The complications in treatment of DFUs include an aberrant regional milieu, excessive inflammation and angiogenesis maladies. The results from this study indicated that all three treatment regimens significantly modulated inflammatory response, and significantly increased angiogenesis on day 16.

A previous study from one of the authors of the current study have shown that because of ADSs immunomodulatory impressions, they might be considered an appropriate treatment of some inflammatory diseases in healthy individuals and patients with DM. The most recent tissue engineering methods in stem cell therapy have shown encouraging choices to repair the weakened, injured skin, including DFUs. Stem cell therapy has arisen as a unique healing method for many illnesses, comprising injury repair and tissue restoration. Many diverse stem cell types have been considered in both animal and human studies and included BM-MSCs and ADS cells for wound repair. Fat tissue provides a plentiful source for MSCs and has shown promising results in wound repair experiments. ADS cells have the capability to change into different cell lines and release paracrine factors that start the tissue restoration process. The plentiful source of adipose tissue, simplicity of extraction, huge proliferative abilities in vitro and their capacity to release pro-angiogenic cytokines make them a perfect cell type as treatment for chronic wounds and wounds. In addition, ADS cells can be very valuable for medical use as they prevent immunological reactions and do not require a chief histocompatibility equivalent for allogeenic engraftment. Engraftment of ADSs encourages new blood vessel formation and increases blood flow to the ischemic organ in mice. It has also been revealed that growth factors released by ADS cells stimulate angiogenesis in ischemic tissue by paracrine signaling. To our knowledge, the combination of PBM+ADS on the repair of infected wounds in patients or animals with DM1 has not been documented. This study aimed to assess the impacts of PBM+ADS on microbial flora, wound area, wound strength and histological parameters in rats with DM1. We found that the use of PBM or ADS alone, or together significantly increased skin injury repair of the MRSA-infected wounds in rats with DM1. Additionally, PBM+ADS showed a synergistic impact. Therefore, PBM+ADS treatment was significantly better than individual treatment by PBM or ADS. This was probably the result of the enhanced survival of ADSs by inhibition of apoptosis, along with increased paracrine signaling of ADS under the influence of PBM.

Our data demonstrated that an anti-inflammatory agent as well as a mitogenic agent, PBM, displayed a synergistic impression on ADS-promoted skin injury repair in rats (figure 5). The wound closure rate was increased in the group that received PBM+ADS. Interestingly, the initiation of bactericidal impact by ADSs was specially improved by PBM, accompanied by the elevated wound strength. These outcomes suggested that administration of PBM at the right protocol might assist with the healing effects of ADS in a DM1 simulation of an MRSA-infected ischemic delayed healing wound. Taken together, our findings recommend that PBM, an anti-inflammatory and mitogen agent, along with a cost-effective modality could be a beneficial enhancement for ADS-based therapy in treating DFUs.

Guo and DiPietro in their review article emphasized on important of wound strength in different medical situations. Our study showed that treatment of wounds with ADS significantly increased wound strength compared with control and PBM groups. This result showed importance of ADS in treating non healing wounds such as diabetic wounds. However, combined administration of ADS and PBM demonstrated a synergistic effect. In current study, the results of fibroblast count and Mallory’s trichrome stain were in the line of wound strength results.

In the current study, stereological analysis supported our hypothesis that the treatments, particularly PBM+ADS modulated the inflammatory response, simultaneously increased wound strength and wound closure rate, and significantly decreased the CFU.

It should also be noted that our unique outcomes reveal the synergistic impact of PBM+ADS on skin injury repair in comparison with other related experiments that only investigated the impression of a solo cure with either PBM or ADS.

Bacterial contamination is a main contributing issue in the deterioration of wound healing. MRSA is a common staphylococcal strain accountable for numerous regional and general contaminations that quickly progresses to resistance to regularly recommended antibiotics. Therefore, the observed occurrence and adaptation of MRSA in DFUs has stemmed from a growing range of non-curable staphylococcal contaminations. Novel approaches to treat MRSA contamination and counter the growing complication of MRSA are mandatory. Specifically, our experiment aimed to focus on the bactericidal effects of PBM and ADS because both have been individually reported to have bactericidal effects in animal models. To date, there is no probe that has examined the effects of PBM on immunomodulation of ADS.

Our outcomes (figure 2) show that PBM treatment could further promote the immunomodulatory efficacy of ADS and might be medically applicable to cell therapies, while better healing impact are required as DFUs, or scalds. In a study of a culture system probe with PBM (415 nm, 100 mW and 30, 60, 120 J/cm²), Lipovesski et al presented an inhibitory impact on S. aureus growth.
by stimulation of ROS synthesis. Consistent with this finding, we hypothesized that the bactericidal effect of the PBM-treated groups in this probe might be due to the induction of ROS by PBM.

Several studies have suggested that stem cells, specifically ADS cells, can speed up repair of a wound, and ADS cells can employ many actions to stimulate skin injury repair. In theory, ADS cells might immediately contribute to tissue restoration by renewing impaired cells via epidermal differentiation, although the weak efficiency of ADS transplantation and viability at the injury site has restricted this direct involvement. Recent investigations have shown significant decreases in stem cells quantity in some experimental simulations of fractures and wounds within the first 14 days of engraftment. Wu et al reported a considerable increase in stem cell survival 7 days after creating the skin injury and a fast decrease in surviving cells 14 days after the infliction of this skin injury. Levi et al studied the effect of human ADS implantation on the repair of a non-healing mice skull defect. They observed significant repair in the implanted skull defect. In their study, at 28 and 56 days after the human ADS implantation, many of the fluorescence in situ hybridization positive cells were not recognized. The outcomes from two probes demonstrated that some implanted ADS cells expressed endothelial markers and were absorbed into vascular networks in ischemic organs, but the rate of contribution was small because many cells expired inside the hypoxic milieu of the ischemic organ. Hence, to improve positive stem cell therapies, it is essential to improve stem cell function, so they can stay alive in ischemic tissue and be able to differentiate into vascular cells, as well.

PBM has been successfully applied for enhancement of local blood circulation and improve repair of an injury by triggering new blood vessel formation in some non-diabetic animal simulations of ischemic tissues. Park et al reported that the release of cytokines was greater in the ADS+PBM group compared with the ADS-only group in a random skin flap (RSF) in mice. ADS improved tissue repair by endothelial cell differentiation and release of angiogenic growth factors. The ADS+PBM group showed better treatment efficiency in comparison with ADS alone. Park et al found that transplanting ADSs to ischemic RSF enhanced healing efficiency for ischemia as an outcome of improved ADS viability and paracrine impact. Moreover, Park et al concluded that PBM improved the viability of ADS and encouraged the release of growth factors in RSF.

The impacts of PBM+CM of hBM-MSC on the infected wounds in rats with diabetes were studied by Bayat group. In the first study, Kouhkheil et al reported the positive impacts of PBM (890 nm, 80 Hz, 0.2 J/cm²) and CM from hBM-MSCs (four injections) together and alone on the CFU and wound strength of an infected wound simulation with MRSA in rats with DM1. In the second study, Fridoni et al demonstrated the positive impact of PBM (890 nm, 80 Hz, 0.2 J/cm²) and CM from hBM-MSC (four injections) alone or together on the stereological parameters of an infected wound simulation with MRSA in rats with DM1. Fridoni et al determined that the use of PBM+CM prompted anti-inflammatory and neovascular actions, and hastened skin injury repair in an MRSA-infected injury in rats with DM1.

In current study, stereological analysis supports our hypothesis regarding our treatments and especially PBM+ADS modulates inflammatory response and at the same time increases wound strength, and significantly increased wound closure rate and CFU. Bayat and Chien have reported the positive effects of combined application of ADS and PBM in some ischemic tissues. They stressed that these results might support new healing attitudes for a cure and resolution of delayed healing of DFU in patients.

CONCLUSION

PBM and ADS alone or together significantly hasten the skin injury repair in an ischemic delayed healing and MRSA-infected wound simulation in rats with DM1. Furthermore, combined use of PBM and ADS have confirmed a synergistic impact. We suggest that PBM+ADS treatment should be used for infected skin infections and wounds in translational studies, and in patients to reduce inflammation, and infection, and promote repair. Additional details of cellular and molecular mechanisms regarding the effect of PBM+ADS on the repair of ischemic delayed healing and MRSA-infected wounds in rats with DM1 should be elucidated by further research.

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Correction notice This article has been corrected since it was published. Dr. Abdollah Amini is added as co-corresponding author.

Contributors MB is the guarantor of this work, and researched data and wrote the manuscript. SC, AA, and SMH contributed to the conception and design. RE-M, F2, AM, SD, ND, MR, MH, and HA performed the experiments and collection of data and SB primary edited the manuscript. SKG contributed to the data analysis and interpretation. SC and SS edited scientifically the manuscript.

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