Betulinic acid accelerates diabetic wound healing by modulating hyperglycemia-induced oxidative stress, inflammation and glucose intolerance

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Abstract

Background: Diabetes significantly delays wound healing through oxidative stress, inflammation and impaired re-epithelialization that lead to defective regulation of the healing process, although the related mechanism remains unclear. Here, we aim to investigate the potential role and mechanism for the beneficial effect of betulinic acid (BA) on diabetic wound healing.

Methods: The molecular effect of BA on hyperglycemia-mediated gene expression, oxidative stress, inflammation and glucose uptake was evaluated in endothelial, fibroblast and muscle cells. Burn injury was introduced to streptozotocin-induced diabetic rats and BA administration through either an intraperitoneal (IP) or topical (TOP) technique was used for wound treatment. Glucose tolerance was evaluated in both muscle tissue and fibroblasts, while oxidative stress and inflammation were determined in both the circulatory system and in wound tissues. The effect of BA on the wound healing process was also evaluated.

Results: BA treatment reversed hyperglycemia-induced glucose transporter type 4 (GLUT4) suppression in both muscle and fibroblast cells. This treatment also partly reversed hyperglycemia-mediated suppression of endothelial nitric oxide synthase (eNOS), nuclear factor erythroid 2-related factor 2 (Nrf2) signaling and nuclear factor NFκB p65 subunit (NFκB p65) activation in endothelial cells. An in vivo rat study showed that BA administration ameliorated diabetes-mediated glucose intolerance and partly attenuated diabetes-mediated oxidative stress and inflammation in both the circulatory system and wound tissues. BA administration by both IP and TOP techniques significantly accelerated diabetic wound healing, while BA administration by either IP or TOP methods alone had a significantly lower effect.

Conclusions: BA treatment ameliorates hyperglycemia-mediated glucose intolerance, endothelial dysfunction, oxidative stress and inflammation. Administration of BA by both IP and TOP techniques was found to significantly accelerate diabetic wound healing, indicating that BA could be a potential therapeutic candidate for diabetic wound healing.
Betulinic acid administration by both intraperitoneal and topical methods significantly accelerates diabetic wound healing. Betulinic acid ameliorates hyperglycemia-mediated glucose transporter type 4 suppression and glucose intolerance in muscle cells. Betulinic acid attenuates hyperglycemia-mediated oxidative stress and inflammation and vascular dysfunction in endothelial cells. Betulinic acid administration by both intraperitoneal and topical methods significantly accelerates diabetic wound healing.

Key words: Betulinic acid, Diabetes, Inflammation, Oxidative stress, Wound healing, Nuclear factor κB, Glucose, Hyperglycemia

Highlights

- Betulinic acid ameliorates hyperglycemia-mediated glucose transporter type 4 suppression and glucose intolerance in muscle cells.
- Betulinic acid attenuates hyperglycemia-mediated oxidative stress and inflammation and vascular dysfunction in endothelial cells.
- Betulinic acid administration by both intraperitoneal and topical methods significantly accelerates diabetic wound healing.

Background

Wound healing is a complicated pathophysiological process that involves the steps of hemostasis, inflammation, proliferation, re-epithelialization and remodeling [1] as well as different cell types, including endothelial cells, fibroblasts, blood cells, muscle and stem cells [2–4]. Diabetes significantly delays wound healing, leading to several defects in the regulation of the normal healing process via consistent oxidative stress [5–7], inflammation [8] and impaired re-epithelialization [9,10]. Diabetes is associated with impaired wound healing [11] among other complications, resulting in a heavy social burden, severe health issues and high rates of morbidity and mortality [12,13]. Development of a potential mechanism and therapeutic treatment for diabetic wound healing is still urgently needed [14–16].

Betulinic acid (BA) is a pentacyclic triterpene product that is purified from natural products including pulsatilla chinensis [17]. It possesses many biological properties and impressive anti-tumor effects [18–20], although the related mechanism is still unclear [21]. Furthermore, it has been reported that BA induces endothelial nitric oxide synthase (eNOS) expression with potential vascular protective effects [22], attenuates oxidative stress by activation of the nuclear factor erythroid 2-related factor 2 (Nrf2) signaling pathway [23–25] with subsequent activation of antioxidant genes, such as superoxide dismutase 2 and glutathione peroxidase [23–27], and ameliorates inflammation by inhibition of the nuclear factor κB (NFκB) signaling pathway [28–30]. Interestingly, BA induces the expression of glucose transporter type 4 (GLUT4) [31] with potential anti-diabetic effects [32–35]. Based on the above observations, BA may appear to have a protective effect on diabetes-mediated oxidative stress, inflammation, vascular dysfunction and glucose intolerance, even as the potential mechanism remains largely unclear [36,37]. Thus, we hypothesize that BA may be a candidate for promoting diabetic wound healing [38].

In addition to the significant negative effects of diabetes complications on diabetic wound healing, diabetes also contributes to severe wound infection, which is usually the major reason for amputations in diabetic foot ulcers [39]. *Staphylococcus aureus* (*S. aureus*) has been identified as the dominant pathogen in diabetic infections such as diabetic foot ulcers [40]. Recent literature has shown that BA may be a potential candidate as a therapeutic agent for *S. aureus*-mediated infection [41]; additionally, it has been reported that BA has promising antimicrobial [42,43] and anti-viral effects [17,44]. These reports indicate that BA may be a promising candidate for diabetic wound healing due to its beneficial effects on diabetes-mediated infection and vascular damage, in addition to its anti-diabetic effects [32–35].

In this study, we aimed to investigate the effect of BA on diabetic wound healing. The results of our *in vitro* study showed that BA partly reversed hyperglycemia-mediated GLUT4 suppression in muscle cells and fibroblasts. A similar effect was observed in partly reversing hyperglycemia-mediated eNOS suppression, NFκB signaling activation and Nrf2 suppression in endothelial cells. An *in vivo* study in rats showed that BA administration partly reversed hyperglycemia-mediated oxidative stress and inflammation in the circulatory system and wound tissues. Similar effects were seen on diabetes-mediated glucose intolerance in muscles and fibroblasts. Furthermore, BA administration by both intraperitoneal (IP) and topical (TOP) treatment significantly accelerated cutaneous burn injury healing in diabetic rats. We conclude that BA accelerates diabetic burn injury healing by modulating hyperglycemia-induced oxidative stress, inflammation and glucose intolerance.

Methods

An expanded Materials and Method section is provided in Supplementary Information (see online supplementary material, Data S1) and the primers used in this study are shown in Table S1 (see online supplementary material).

Materials and reagents

Human primary aorta smooth muscle cells (HASMCs, #PCS-100-012) were purchased from ATCC and maintained in vascular cell basal medium supplemented with vascular smooth muscle growth kit. Antibodies for eNOS (sc-376751), GLUT4 (sc-53566) and NFκB p65 (sc-8008) were obtained from Santa Cruz Biotechnology (Shanghai, China). The transcriptional activity of NFκB p65 was determined by the NFκB p65 transcription factor assay kit (#ab133112 from Abcam) according to the manufacturer’s instructions and BA (#B8936) was purchased from Sigma (Shanghai, China).

DNA methylation analysis

Human genomic DNA was extracted and purified from HASMCs, treated by bisulfite modification and then amplified by the following primers: methylated primer: forward 5′- tgt ttt cag ttt tgg ttt ttc -3′, reverse 5′- aac ccc ata aat aaa ttc cta cgt a -3′; unmethylated primer: forward 5′- ttt ttt gag ttt gta ttt tgg ttt gcg -3′; reverse 5′- ttt ttg aac ccc ata aat aaa ttc cta cgt a -3′. Product size: 166 bp (methylated)
and 164 bp (unmethylated); CpG island size: 231 bp; melting temperature (Tm): 63.8°C. The final results for DNA methylation were normalized using unmethylated results as input [45].

**In vivo rat experiments**

The animal protocol conformed to US NIH guidelines (Guide for the Care and Use of Laboratory Animals, No. 85–23, revised 1996) and was reviewed and approved by the Institutional Animal Care and Use Committee.

**Rat protocol 1: generation of diabetic rats** Two-month-old rats were fed a high-fat diet for 1 month followed by injection of 35 mg/kg streptozotocin (STZ) after an 8-h fast. Blood glucose levels were monitored for 4 weeks continuously and rats with blood glucose levels > 11.1 mM were selected as the diabetes positive group. The control (CTL) group received only vehicle (VEH) injection [24,46].

**Rat protocol 2: induction of cutaneous burn injury** The cutaneous burn injury was introduced to rats in protocol 1. The dorsa of the rats were exposed to a hot copper pillar (2-cm diameter) at 75°C for 15 s and the subsequent wound healing process was monitored and evaluated [24,46].

**Rat protocol 3: BA treatment of cutaneous burns in a diabetic rat model** The rats in protocol 2 received either VEH or BA administration. In the procedure for IP administration, BA was dissolved in 0.1% Dimethylsulfoxide (DMSO) (diluted by 0.9% NaCl solution) and administered intraperitoneally every 3 days at a dose of 10 mg/kg body weight for 4 weeks starting from 1 week before the introduction of burn injury. For TOP administration, the dissolved BA was administrated by spraying in a dose of 20 μM each day continuously for 3 weeks starting from the second day after the introduction of burn injury. The experimental rats were randomly separated into four groups as follows: group 1: CTL rats with VEH treatment (CTL/VEH); group 2: STZ-induced diabetic (STZ) rats with VEH treatment (STZ/VEH); group 3: STZ rats with BA TOP administration (STZ/BA-TOP); group 4: STZ rats with BA IP administration (STZ/BA-IP); group 5: STZ rats with BA treatment by both TOP and IP administration (STZ/BA-IP/TOP). Wound status was monitored and evaluated throughout the treatment. After treatment, the animals were subjected to glucose/insulin tolerance tests. Whole blood was then withdrawn by heart puncture and the serum was prepared by centrifugation. Peripheral blood mononuclear cells (PBMCs) were isolated from the blood using Ficoll-Paque Plus lymphocyte separation medium and fibroblast cells were isolated from the underarm area of skin from treated mice for *in vitro* biological assays [47]. The rats were sacrificed and the wound tissues and soleus muscles were collected.

**Analysis of glucose uptake**

Treated cells and the soleus muscles were used for this assay. The soleus muscles were isolated from treated rats and were dissected, weighed, pre-incubated (30 min) and then incubated (60 min) at 37°C in Krebs Ringer–bicarbonate buffer with a composition of 122 mM NaCl, 3 mM KCl, 1.2 mM MgSO4, 1.3 mM CaCl2, 0.4 mM KH2PO4 and 25 mM NaHCO3 and bubbled with O2/CO2 (95:5%, v/v) until pH 7.4. [U-14C]-2-Deoxy-D-glucose ([14C]-DG) (0.1 μCi/mL) was added to each sample during the incubation. After incubation, the muscles were placed in screw-cap tubes containing 1 mL of distilled water and frozen at −20°C in a freezer followed by 10 min of boiling. Aliquots of tissue and external medium (25 μL) were placed in a scintillation counter for radioactivity measurements [48] and the glucose uptake results were expressed as counts per min/ml of incubation medium [31].

**Intraperitoneal glucose tolerance test**

For the glucose tolerance test, pre-treated mice were injected intraperitoneally with 2 g/kg body weight of glucose after a 6-hour fasting period. Blood samples were collected from the tail vein and blood glucose was monitored using a One-Touch Ultra®2 Glucometer at time points of 0, 15, 30, 60, 90 and 120 min after injection. Serum insulin levels were evaluated using the rat insulin enzyme-linked immunosorbent assay (ELISA) kit (#ERINS from Invitrogen) according to manufacturer’s instructions [34,49].

**Measurement of oxidative stress**

The Glutathione/Glutathione disulfide (GSH/GSSG) ratio was determined using the GSH/GSSG ratio detection assay kit (fluorometric - green) (#ab205811 from Abcam) and the transcriptional activity of Nrf2 was determined using the Nrf2 transcription factor assay kit (colorimetric) (#ab207223 from Abcam). Methylglyoxal (MG) formation was determined using the methylglyoxal assay kit (#ab241006 from Abcam) and 3-nitrotyrosine formation was measured using the 3-nitrotyrosine ELISA kit (#ab116691 from Abcam) according to the manufacturer’s instructions.

**Statistical analysis**

One-way Analysis of variance (ANOVA) analysis together with the Bonferroni post hoc test was employed to evaluate statistical significance among the treatments under normal distribution. The experiments were repeated at least four times (n = 4) unless otherwise mentioned. The data are presented as mean ± standard deviation (SD) and SPSS 22 software was employed for the analysis of statistical difference. *P value* < 0.05 indicates a significant difference [24,46].

**Results**

**BA ameliorates hyperglycemia-induced oxidative stress and inflammation in human umbilical vein endothelial cells (HUVECs)**

We determined the potential effect of BA and hyperglycemia on oxidative stress and inflammation. The HUVECs were incubated with 5 mM glucose (low glucose, LG), 25 mM glucose (high glucose, HG) or HG plus 20 μM BA for
4 days, and the cells were harvested for biological assays. Our results showed that HG treatment significantly decreased mRNA (Figure 1a) and protein levels (Figures 1b, c and S1a, see online supplementary material) of eNOS compared to the LG group, but had no effect on Nrf2 or NFκB p65. Additionally, BA treatment (HG/BA) completely reversed the HG-mediated effect. Moreover, HG treatment significantly increased NFκB p65 activity and phospho-Ser139 histone H2AX (γH2AX) formation, but decreased Nrf2 activity compared to the LG group. BA treatment (HG/BA) either partly or completely reversed this effect (Figures 1d, e and S1b, see online supplementary material). We also measured oxidative stress, and the results showed that BA treatment partly reversed hyperglycemia-mediated increased reactive oxygen species (ROS) formation (Figure 1f) and 8-hydroxy-2′-deoxyguanosine (8-OHdG) formation (Figure 1g). We then established dose–response curves for the effect of BA on hyperglycemia-mediated oxidative stress and inflammation, and the results showed that BA (20 μM) treatment significantly ameliorated hyperglycemia (HG)-mediated increased MG formation (Figure 1h) and ROS formation (Figure 1i), as well as hyperglycemia (HG)-mediated decreased Nrf2 activity (Figure 1j) and increased NFκB p65 activity (Figure 1k).

We conclude that BA treatment ameliorates hyperglycemia-mediated oxidative stress and inflammation in HUVECs. BA reverses hyperglycemia-induced GLUT4 suppression by modulation of histone methylation on the GLUT4 promoter

We determined the molecular effect of BA and hyperglycemia on GLUT4 expression in HASMCs and found that 15 μM BA partly, while 20 μM of BA completely, reversed hyperglycemia-mediated decreased expression of GLUT4 mRNA (Figure 2a, b) and protein (Figures 2c, d and S1c, see online supplementary material) as well as hyperglycemia-mediated decreased GLUT4 reporter activity (Figure 2e) and 14C-DG uptake (Figure 2f). We then investigated the possible mechanism for hyperglycemia-mediated GLUT4 suppression. A series of progressive 5′-promoter deletions for GLUT4 reporter constructs was generated, and either GLUT4 full length (pGLUT4–2000) or deletion reporter plasmids were transiently transfected into conditionally
immortalized HASMCs. Reporter activity was then measured after a 3-day treatment with either LG or HG. We found that hyperglycemia-mediated GLUT4 suppression occurred in the GLUT4 deletion constructs of pGLUT4–2000, −1600, −1400, −1200, −800, −600 and −400, while reporter suppression was partly restored in pGLU4–200 and −100 and completely restored in the pGLUT4–0 construct, suggesting that hyperglycemia-mediated transcriptional suppression is located in the range of −300 to −0 on the GLUT4 promoter (Figure 2g). We then evaluated possible epigenetic changes in this area, and results showed that HG treatment significantly increased histone methylation at H3K9me3 and H3K27me3 compared to LG treatment and that BA treatment partly reversed this effect (Figure 2h). On the other hand, there was no difference among treatments in histone methylation at H3K9me2 or H3K27me2 or in DNA methylation on the GLUT4 promoter (Figure 2i).

Additionally, we evaluated possible epigenetic changes on histone 4 methylation (Figure S2a, see online supplementary material) and histone 3 acetylation (Figure S2b, see online supplementary material), and the results showed no effect among the treatments. We conclude that BA treatment partly reverses hyperglycemia-mediated GLUT4 suppression by modulation of histone methylation on the GLUT4 promoter.

BA administration ameliorates diabetes-induced oxidative stress and inflammation in PBMCs

We determined the potential effect of BA and diabetes on oxidative stress in PBMCs. Diabetic rats with burn injury received BA administration by IP, TOP or both treatments (IP/TOP), and then the serum and/or PBMCs were isolated for biological assays. We found that the diabetes (STZ/VEH) group showed significantly reduced Nrf2 transcriptional activity in PBMCs compared to the CTL/VEH group, and that TOP administration of BA (BA-TOP) had little effect, while BA administration by either IP alone or a combination of IP and TOP (IP/TOP) partly reversed this effect (Figure 3a). Additionally, STZ/VEH treatment significantly potentiated ROS generation (Figure 3b), MG formation (Figure 3c) and 8-oxo-dG formation (Figure 3d) in PBMCs compared to the CTL/VEH group. BA-TOP had little effect, while treatment with either BA-IP or BA-IP/TOP significantly reversed this effect. We also evaluated the GSH/GSSG ratio in serum, and the results showed that treatment with either BA-IP or BA-IP/TOP significantly reduced the GSH/GSSG ratio, while the BA-TOP group showed no effect (Figure 3f). We then evaluated the effect of BA on diabetes-mediated inflammation and found that STZ/VEH treatment significantly potentiated NFκB p65 transcriptional activity in PBMCs compared to the CTL/VEH group; BA-TOP treatment showed little effect, while treatment with either BA-IP or BA-IP/TOP partly reversed this effect (Figure 3g). We also determined the mRNA levels of pro-inflammatory cytokines and found that treatment with BA-IP or BA-IP/TOP either partly or completely reversed diabetes (STZ/VEH)-induced increased cytokine expression, while BA-TOP treatment showed little effect (Figure 3h). We also determined the serum levels of the pro-inflammatory cytokines, including interleukin-1β (IL-1β) (Figure 3i), IL-6 (Figure 3j) and monocyte
Figure 3. BA administration ameliorates hyperglycemia-induced oxidative stress and inflammation in PBMCs. Rat models with cutaneous burn injury received either CTL or STZ treatment and then received treatment of either vehicle (VEH) or BA administration through intraperitoneal (IP), topical (TOP) or both methods (IP/TOP); the serum and/or PBMCs were then isolated on day 23 after introduction of burn injury for biological assays. (a) Nrf2 transcriptional activity, n = 5. (b) ROS formation, n = 5. (c) Methylglyoxal formation, n = 5. (d) Quantitation of 8-oxo-dG formation, n = 5. (e) Representative pictures of 8-oxo-dG staining (red) and DAPI staining (blue) for (d). (f) GSH/GSSG ratio in serum, n = 5. (g) NF-κB p65 transcriptional activity, n = 5. (h) mRNA level in PBMC, n = 4. (i) serum IL-1β level, n = 5. (j) serum IL-6 level, n = 5. (k) serum MCP1 level, n = 5. *p < 0.05 vs CTL/VEH group; ¶p < 0.05 vs STZ/VEH group. Data are expressed as mean ± SD. BA: betulinic acid, PBMCs: peripheral blood mononuclear cells, CTL: control, STZ: streptozotocin, DAPI: 4,6-diamidino-2-phenylindole, IL: interleukin, MCP1: monocyte chemoattractant protein-1, SD: standard deviation, OD: optical density.

BA administration ameliorates diabetes-mediated GLUT4 suppression and glucose intolerance in muscles

We determined the effect of BA on diabetes-mediated GLUT4 suppression in muscles and found that STZ/VEH treatment significantly reduced GLUT4 expression of both mRNA (Figure 4a) and protein levels (Figures 4b, c and S1d, see online supplementary material) compared to the CTL/VEH group. BA-TOP treatment showed little effect, while treatment with either BA-IP or BA-IP/TOP partly reversed this effect. Additionally, treatment with either BA-IP or BA-IP/TOP partly reversed diabetes (STZ/VEH)-mediated suppression of 14C-DG uptake, and again BA-TOP treatment had little effect (Figure 4d). We also evaluated the potential effect of BA on fibroblast cells isolated from the skin of treated mice and found that BA treatment completely reversed diabetes (STZ)-mediated GLUT4 suppression (Figure S3a, see online supplementary material), partly reversed diabetes-induced epigenetic changes on the GLUT4 promoter (Figure S3b), and completely reversed diabetes-mediated glucose uptake (Figure S3c). Finally, we determined the possible effect of BA administration on diabetes-mediated glucose intolerance and found that STZ/VEH treatment significantly increased blood glucose levels after glucose injection during the intraperitoneal glucose tolerance test compared to the CTL/VEH group. BA-TOP treatment showed little effect, while treatment with either BA-IP or BA-IP/TOP partly reversed this effect (Figure 4e). We again determined the insulin levels and found that the diabetic (STZ/VEH) group had significantly less insulin secretion during the intraperitoneal glucose tolerance test compared to the CTL/VEH group. BA-TOP treatment showed little effect, while treatment with either BA-IP or BA-IP/TOP partly reversed this effect (Figure 4f). We conclude that BA-IP administration has significant anti-diabetic effects on diabetes-mediated GLUT4 suppression and glucose intolerance.

BA administration ameliorates diabetes-mediated oxidative stress and inflammation

We determined the effect of BA and hyperglycemia on gene expression in wound tissues. We found that STZ/VEH...
Figure 4. BA administration ameliorates diabetes-mediated GLUT4 suppression and glucose intolerance in muscles. Rat models with cutaneous burn injury received CTL or STZ treatment and then received treatment of either vehicle (VEH) or BA administration by intraperitoneal (IP), topical (TOP) or both methods (IP/TOP). The tissues or soleus muscles were isolated and used for analysis on day 23 after burn injury. (a–d) The muscles were used for biological assays. (a) mRNA levels, n=4. (b) Quantitation of protein level by western blot, n=5. (c) Representative western blots for (b). (d) 14C-DG uptake assay, n=5. ∗p < 0.05 vs LG/VEH group; ¶p < 0.05 vs STZ/VEH group. (d,e) Rats were used for the intraperitoneal glucose tolerance test. (e) Blood glucose levels, n=5. (f) Serum insulin levels, n=5. Data are expressed as mean ± SD. BA betulinic acid, GLUT4 glucose transporter type 4, CTL control, STZ streptozotocin, LG low glucose, SD standard deviation.

treatment significantly decreased mRNA (Figure 5a) and protein levels (Figures 5b, c and S1e, see online supplementary material) of eNOS in whole cell lysates compared to the CTL/VEH group; treatment with either BA-TOP or BA-IP partly, while BA-IP/TOP treatment completely, reversed this effect. Additionally, there was no difference in the expression of Nrf2 and NFκB p65 among all the treatments. We then determined the transcriptional activity of Nrf2 and NFκB p65 by measuring related protein levels in nuclear extracts, and the results from western blotting showed that treatments with BA-TOP and BA-IP partly, while BA-IP/TOP treatment completely, reversed diabetes (STZ/VEH)-mediated NFκB p65 activation and Nrf2 suppression (Figures 5d, e and S1f, see online supplementary material). We evaluated the effect of BA and diabetes on oxidative stress, and the results showed that treatment with BA-TOP and BA-IP partly, while BA-IP/TOP treatment almost completely, reversed diabetes (STZ/VEH)-mediated increased γH2AX formation (Figures 5e, f and S1f), 3-nitrotyrosine formation (Figure 5g, h) and 8-oxo-dG formation (Figure 5i, j). We also determined the effect of BA and diabetes on inflammation in wound tissues, and the results showed that diabetes (STZ/VEH) treatment significantly increased mRNA levels of IL1β, IL6 and MCP1; treatment with BA-TOP and BA-IP partly, while BA-IP/TOP treatment completely, reversed this effect (Figure 5k). We also evaluated protein levels from wound tissues for the pro-inflammatory cytokines, including IL1β (Figure 5l), IL6 (Figure 5m) and MCP1 (Figure 5n), and the expression was similar to related mRNA levels. We conclude that BA administration ameliorates hyperglycemia-mediated oxidative stress and inflammation.

BA administration accelerates diabetic wound healing

We determined the effect of BA administration on diabetic wound healing. The CD31 positive cells in wound tissues were identified by immunohistochemistry on days 7, 14 and 21, respectively after burns, and we found that diabetes (STZ/VEH) significantly reduced the generation of CD31 positive cells compared to the CTL/VEH group; treatment with either BA-TOP or BA-IP showed little effect, while this effect was completely reversed in the BA-IP/TOP group (Figure 6a, b). We then determined the deposition of granulation cells by hematoxylin and eosin (H&E) staining and found that treatment with either BA-TOP or BA-IP showed little effect, while diabetes-mediated decreased deposition of granulation cells (Figure 6c, d). Finally, we evaluated the effect of BA administration on the progress of diabetic wound healing. Pictures of
Figure 5. BA administration ameliorates diabetes-mediated oxidative stress and inflammation in wound tissues. Rat models with cutaneous burn injury received CTL or STZ treatment and then received treatment of either vehicle (VEH) or BA administration by intraperitoneal (IP), topical (TOP) or both methods (IP/TOP); the wound tissues were then isolated on day 23 after burns for biological assays. (a) mRNA levels by qPCR, \(n=4\). (b) Protein quantitation by western blot, \(n=5\). (c) Representative western blots for (b). (d) Protein quantitation in nuclear extracts by western blot, \(n=5\). (e) Representative western blots for (d). (f) Quantitation of \(\gamma\)H2AX levels for (e). (g) 3-Nitrotyrosine (3-NT) quantitation, \(n=6\). (h) Representative pictures of 3-NT staining for (g). (i) 8-oxo-dG quantitation, \(n=6\). (j) Representative pictures of 8-oxo-dG staining for (i). (k) mRNA level in wound tissues, \(n=4\). (l) IL-1\(\beta\) in tissues, \(n=5\). (m) IL-6 in tissues, \(n=5\). (n) MCP1 in tissues, \(n=5\). *\(p<0.05\) vs CTL/VEH group; ¶\(p<0.05\) vs STZ/VEH group. Data are expressed as mean \(\pm\) SD. BA betulinic acid, \(\gamma\)H2AX PhosphoSer139 Histone H2AX, CTL control, STZ streptozotocin, MCP1 monocyte chemoattractant protein-1, IL interleukin, SD standard deviation.

Figure 6. BA administration accelerates diabetic wound healing. Rat models with cutaneous burn injury received CTL or STZ treatment and then received treatment of either vehicle (VEH) or BA administration by intraperitoneal (IP), topical (TOP) or both methods (IP/TOP). The wounds were then isolated for analysis and the healing process was evaluated. (a) Representative pictures for evaluation of vascularity (assessed by CD31 immunohistochemistry). (b) Quantitative numbers of CD31 positive per HPF area on day 14 after burns for (a), \(n=8\). (c) H&E staining of granulation cells on day 21 after burns. (d) Granulation cell deposition from (c), \(n=8\). (e) Photographs of representative wounds on days 1, 7, 14 and 21 after burns. (f) Quantitation of burn area on day 21, \(n=8\). (g) Graphical depiction of wound areas on different days after burns, \(n=8\). *\(p<0.05\) vs CTL/VEH group; ¶\(p<0.05\) vs STZ/VEH group. Data are expressed as mean \(\pm\) SD. BA betulinic acid, CTL control, STZ streptozotocin, H&E hematoxylin and eosin, HPF high power field, SD standard deviation.
the wound were taken every day after burns, and the results indicated that treatment with either BA-TOP or BA-IP partly, and BA-IP/TOP treatment completely, reversed diabetes-mediated delayed diabetic wound healing (Figure 6e–g).

**Schematic model for BA-mediated acceleration of diabetic wound healing**

Diabetes or hyperglycemia-mediated overgeneration of superoxide anions and MG formation results in decreased expression of eNOS, epigenetic changes and GLUT4 suppression, as well as suppressed Nrf2 transcriptional activity and increased NFκB p65 activity, subsequently triggering oxidative stress, inflammation, vascular dysfunction and glucose intolerance. On the other hand, BA administration can either partly or completely reverse those diabetes-mediated effects; thus, BA administration could significantly accelerate diabetic wound healing (Figure 7).

**Discussion**

In this study, we demonstrate that BA reverses hyperglycemia-mediated GLUT4 suppression in HASMCs and also partly reverses hyperglycemia-mediated eNOS suppression, Nrf2 suppression and NFκB p65 activation in HUVECs. Additionally, BA administration in rats ameliorates diabetes-mediated glucose intolerance in muscles and fibroblasts, as well as diabetes-induced oxidative stress and inflammation in the circulation and wound tissues. Finally, we show that administration of BA by both IP and TOP methods significantly accelerates diabetic wound healing.

**BA-mediated effect on oxidative stress and inflammation**

It has been reported that BA attenuates oxidative stress by activation of the Nrf2 signaling pathway [23,50]. Our results showed that BA treatment significantly increased Nrf2 transcriptional activity but showed no effect on Nrf2 expression [25], subsequently ameliorating hyperglycemia-mediated ROS generation, 8-oxo-dG formation and MG formation, and decreased the GSH/GSSG ratio in the circulatory system. Additionally, our results showed that BA treatment significantly ameliorated hyperglycemia-mediated NFκB p65 transcriptional activation [51], subsequently decreasing the release of pro-inflammatory cytokines, including IL1β, IL6 and MCP1 [24]. Taken together, BA can ameliorate hyperglycemia-induced oxidative stress and inflammation; this effect may partly contribute to its beneficial effect on diabetic wound healing, although the detailed mechanism remains largely unknown.

**BA-mediated anti-diabetic effect**

It has been reported that BA induces GLUT4 activation in muscle cells [31]. A BA-mediated anti-diabetic effect has been previously reported [32–35], although the related mechanism remains largely unknown. Our results showed that BA treatment increased GLUT4 expression in both muscle tissue and fibroblasts, subsequently ameliorating diabetes-mediated glucose intolerance. In this scenario, BA could be a potential candidate for treatment of diabetes and could subsequently be beneficial for therapeutic treatment of diabetic wound healing by ameliorating related diabetic complications [52,53].

**BA-mediated beneficial effect on diabetic wound healing**

In addition to the beneficial effect of BA on diabetes-mediated oxidative stress, inflammation and glucose intolerance, our results also showed that BA treatment significantly reversed hyperglycemia-mediated eNOS suppression [22], indicating that BA may play a vascular protective role in diabetic complications [54]. In this study, BA was administrated through either IP or TOP methods for the treatment of burn injury in diabetic rats, and the results showed that BA administration by IP significantly attenuated diabetes-mediated oxidative stress and inflammation in PBMCs, while BA administration by TOP showed little effect. On the other hand, IP administration of BA had a significantly lower effect on wound tissues compared to TOP administration; this can be explained as BA has very low gastrointestinal absorption. Finally, our results showed that BA administration by both IP and TOP significantly accelerated diabetic wound healing; this suggests that treatment of diabetic wound healing should not just focus on the wound/injury itself; instead, more attention should be paid to diabetes and diabetes-mediated complications and dysfunction. This is the first time that BA has been used for the treatment of diabetic burn injury with positive results due to its anti-diabetic effect and potential protective role in diabetes-mediated dysfunction, such as oxidative stress and inflammation [55,56].

**Conclusions**

BA treatment ameliorates diabetes-mediated glucose intolerance by activation of GLUT4 expression, attenuates
hyperglycemia-induced oxidative stress and inflammation by Nrf2 activation and NFκB p65 signaling suppression, and plays a vascular protective role by stimulation of eNOS expression. BA administration by both IP and TOP methods significantly accelerates diabetic wound healing. We conclude that BA accelerates diabetic wound healing by modulating hyperglycemia-induced oxidative stress, inflammation and glucose intolerance and that BA may be a promising therapeutic candidate for the treatment of diabetic wound healing.

**Abbreviations**

BA: Betulinic acid; CTL: Control; 14C-DG: [U-14C]-2-Deoxy-D-glucose; eNOS: Endothelial nitric oxide synthase; GLUT4: Glucose transporter type 4; HAMSC: Human primary aorta smooth muscle cells; HG: High glucose; γH2AX: Phospho-Ser139 Histone H2AX; IL1β: Interleukin-1β; IP: Intraperitoneal; LG: Low glucose; MCP1: Monocyte chemoattractant protein-1; MG: Methylglyoxal; NFκB p65: Nuclear factor NFκB p65 subunit; Nrf2: Nuclear factor erythroid 2-related factor 2; 8-OHdG: 8-Hydroxy-2′-deoxyguanosine; PBMC: Peripheral blood mononuclear cells; ROS: Reactive oxygen species; STZ: Streptozotocin; VEH: Vehicle; TOP: topical.

**Supplementary data**

Supplementary material is available at *Burns & Trauma Journal* online.

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**Authors’ contributions**

PY wrote the paper, PY and WX designed, analyzed the data and interpreted the experiments. WH, ZH and XH performed part of the animal experiments, HZ performed part of the gene expression analysis. ML performed statistical analysis. WX performed the remaining experiments. All authors read and approved the final manuscript.

**Ethics approval and consent to participate**

The animal protocol conformed to US NIH guidelines (Guide for the Care and Use of Laboratory Animals, No. 85–23, revised 1996), and was reviewed and approved by the Institutional Animal Care and Use Committee.

**Conflicts of interest**

None declared.

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