Chitosan/Sodium Alginate/Velvet Antler Blood Peptides Hydrogel Promotes Diabetic Wound Healing via Regulating Angiogenesis, Inflammatory Response and Skin Flora

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Background: Diabetic ulcer remains a clinical challenge due to impaired angiogenesis and persistent inflammation, requiring new alternative therapies to promote tissue regeneration.

Purpose: In this study, chitosan/sodium alginate/velvet antler blood peptides (CS/SA/VBPs) hydrogel (CA VBPH) was fabricated and used in the treatment of skin wounds in type 2 diabetes mellitus (T2D) for the first time.

Methods: VBPs were prepared by hydrolysis and ultrafiltration, and their sequences were identified using LC-MS/MS. The CA VBPH was further fabricated and characterized. A mouse model of T2D was induced by a high-sugar and high-fat diet (HSFD) and streptozotocin (STZ) injection. CA VBPH was applied topically to T2D wounds, and its effects on skin repair and potential biological mechanisms were analyzed by appearance observation, histopathological staining, bioinformatics analysis, Western blot, and 16S rRNA sequencing.

Results: VBPs had numerous short-chain active peptides, excellent antioxidant activity, and a low hemolysis rate. CA VBPH exhibited desirable biochemical properties and participated in the diabetic wound healing process by promoting cell proliferation (PCNA and α-SMA) and angiogenesis (capillaries and CD31) and alleviating inflammation (CD68). Mechanistically, the therapeutic effect of CA VBPH on chronic wounds might rely on activating the PI3K/AKT/mTOR/HIF-1α/VEGFA pathway and reversing the expression of inflammatory cytokines TNF-α and IL-1β. The results of 16S rRNA sequencing indicated that T2D significantly altered the diversity and structure of skin flora at the wound site. CA VBPH treatment elevated the relative abundance of beneficial microbes (e.g., Corynebacterium_1 and Lactobacillus) and reversed the structural imbalance of skin microbiota.

Conclusion: These results indicate that CA VBPH is a promising wound dressing, and its repair effect on diabetic wounds by regulating angiogenesis, inflammatory response, and skin flora may depend on the rich small peptides in VBPs.

Keywords: velvet antler blood peptides, diabetic wound, hydrogel, bioinformatics, skin flora

Introduction

Diabetes mellitus is a common chronic metabolic disease, typically manifested as metabolic disorders and hyperglycemia, and its incidence is expected to increase to 592 million patients by 2035.1,2 15–25% of diabetic patients have non-healing skin wounds, such as foot ulcers (DFU), which are usually in a stage of chronic inflammation rather than a series of repairing events, and further pose gangrene, amputation and even death.3 Because traditional therapies and the exploration of underlying mechanisms are usually inadequate, more effective clinical treatment strategies to promote diabetic wound healing are desired.4–7
Uncovered diabetic wounds often display persistent hyperglycemia and immune dysregulation, and are susceptible to bacterial invasion. Predictably, these aforementioned conditions further aggravate the inflammatory response, skin ulcers, and healing impairments.8 Therefore, the treatment of diabetic wounds requires dressings with multiple functions such as good hygroscopicity, drug delivery ability, biodegradability, antibacterial effect, anti-inflammatory activity, and antioxidant capability to overcome the limitations of traditional therapy.9 Hydrogels are becoming an attractive option in creating environments for promoting wound healing due to their skin-mimicking structure and excellent biochemical characteristics. As natural polysaccharides, chitosan (CS) and sodium alginate (SA) are widely applied in the field of tissue engineering, especially in the development of wound dressings.10–12 CS/β-glycerophosphate sodium (β-GP) hydrogel systems have good temperature sensitivity and biocompatibility, which are beneficial to drug release, cell proliferation, and tissue repair and have been utilized in ocular treatment, periodontal repair, and skin trauma at present.13–15 Alginate biomaterial can also enhance granulation tissue formation and wound healing.12 Hence, the combination of biopolymers with healing potential may provide an opportunity to synthesize wound dressings that stimulate skin repair.

The previously reported CS-based oyster peptide microsphere thermosensitive hydrogel promoted skin wound repair in mice via up-regulating the expression of Ki-67 and VEGF and suppressing inflammatory cytokines.16 Wei et al fabricated a CS/alginate hydrogel loaded with FGF/VE-cadherin and evaluated its potential as a wound dressing for repairing defective skin.17 In addition, hydroxypropyl cellulose/SA hydrogel loaded with antibacterial peptide NZ2114 was confirmed to accelerate wound healing by promoting the expression of VEGF, EGF, and CD31 as well as reducing inflammatory response.18 These studies demonstrate that hydrogels encapsulating bioactive peptides/proteins seem to be a promising research direction for wound dressings. However, it is currently insufficient to deeply explore the biological mechanism by which hydrogels accelerate wound repair.

In recent years, animal-derived small peptides, including velvet antler peptides, oyster peptides, chum salmon peptides, and tilapia peptides, have been found to have not only antibacterial, anti-inflammatory, and antioxidant properties but also skin care and tissue repair functions.16,19–21 Velvet antler blood (VB) is the blood extracted from freshly collected velvet antler and has been recorded as a traditional Chinese medicine for restoring and tonifying in Chinese medical classics Shen Nong Ben Cao Jing 2000 years ago.22 VB is rich in protein, providing the material basis for the growth and restoration of velvet antler. It was previously found that velvet antler blood peptides (VBPs), the major components of VB, have significant anti-fatigue and antioxidant effects, yet, it is still unknown whether VBPs are a crucial factor for tissue repair.23,24 Based on the above characteristics, in this study, we fabricated a CS/SA thermosensitive hydrogel loaded with VBPs (CAVBPH) and further explored its effects on diabetic wounds and skin flora. To the best of our knowledge, this is the first comprehensive exploration of the underlying biological mechanisms of CAVBPH for diabetic wound healing.

Materials and Methods

Materials
Freeze-dried VB powder was obtained from Jinlu Pharmaceutical Co., Ltd. (Changchun, China). SA (molecular weight of 32–250 kDa) and CS (deacetylation degree of 95%, viscosity of 200–400 mPa·s) were purchased from Meryer Chemical Technology Co., Ltd. (Shanghai, China). β-GP was purchased from Yuanye Bio-Technology Co., Ltd. (Shanghai, China). Streptozocin (STZ) and chloral hydrate were, respectively, from Sigma-Aldrich Co., Ltd. (Shanghai, China) and Fuchen Chemical Reagent Factory (Tianjin, China). BCA protein assay kit (P0010) and RIPA lysis buffer (P0013B) were obtained from Beyotime (Shanghai, China). HaCat cells were from Cellcook Biotech Co., Ltd. (Guangzhou, China). All other agents were of analytical grade.

Preparation and Characterization of VBPs
VBPs were prepared using in vitro simulated gastrointestinal digestion and ultrafiltration technique as previously reported.25 Peptide sequences were identified and analyzed by utilizing the Ultimate 3000 high-performance liquid chromatography system (HPLC, Thermo Fisher Scientific, USA) coupled with a Q Exactive™ Hybrid Quadrupole-Orbitrap™ Mass Spectrometer (Thermo Fisher Scientific, USA) with an ESI nanospray source.
Preparation of CAVBPH
First, 2% w/v CS solution was prepared in the acetic acid solution (0.1 M, 10 mL), and 2.5 mL of 56% w/v β-GP solution was dripped into the cold CS solution and fully mixed. Then, 0.1 g SA was added to the CS/β-GP solution, and 0.2 mL of 0.5% w/v CaCl$_2$ solution was added dropwise to the mixture to prepare CS/β-GP/SA/Ca$^{2+}$ hydrogel (CAH). Subsequently, VBPs were introduced into CAH scaffold and CAVBPH with peptides concentration of 1 mg/mL was fabricated. The gelation behavior of CAH and CAVBPH was evaluated via the glass vial dumping test at 37 °C. 5 mL of CAH solution or CAVBPH solution transitioned from sol to gel within 5 min at this controlled temperature, indicating that hydrogels possessed good temperature sensitivity. The solutions mentioned above were freshly prepared and kept at 4 °C before use.

In vitro Characterization of Hydrogels
The microstructure of freeze-dried hydrogel was analyzed by scanning electron microscope (SEM; SS-550, Shimadzu, Japan). Infrared (IR) spectra were detected using an FTIR spectrometer (FTIR-650, Gangdong, China). Subsequently, the swelling behaviors and in vitro biodegradation profiles of samples were, respectively, investigated according to previous reports. The release profile of VBPs was further plotted. Briefly, lyophilized CAVBPH ($W_0$) was immersed in 10 mL of PBS with constant shaking. At regular intervals, 1 mL of the co-incubated solution was replaced by the same volume of freshly prepared PBS. The release of VBPs was analyzed utilizing BCA protein assay kit according to the manufacturer’s instructions.

Antioxidant and Biosafety Assays
Hydrogels were soaked in PBS for 24 h to prepare the extraction solutions (1 mg/mL), and GSH and VBPs were used as the positive control. Antioxidant ability of CAH and CAVBPH was evaluated via DPPH and ABTS radical scavenging assays according to a previous study. Meanwhile, 1% w/v TritonX-100 solution and PBS were, respectively, prepared as the positive control and negative control, and the hemolysis rate was determined according to the reference. HaCaT cells were selected to test the cytotoxicity of hydrogel samples. UV-sterilized CAH or CAVBPH was immersed in the culture medium for 24 h to prepare the hydrogel extract medium. And HaCaT cells culture and cell viability assay (MTT method) were performed as previously described.

Bioinformatics Analysis
The dataset of diabetic skin wounds (GSE147890) based on the GPL571 platform was downloaded from the Gene Expression Omnibus database (https://www.ncbi.nlm.nih.gov/geo/). Dataset GSE147890 contains 5 intact normal skin samples (C0h), 7 intact diabetic skin samples (D0h), 6 acute wounds samples 24 h after operation (C24h), and 6 diabetic wounds samples 24 h after operation (D24h). The DEGs of acute wound (C24h vs C0h) and diabetic wound (D24h vs D0h) were identified with the threshold of $|\log_{2}\text{FC}| > 1$ and $p < 0.05$ by utilizing the limma package within the R language environment (version 3.6.2). The overlapping DEGs of acute wounds and diabetic wounds and the specific DEGs of diabetic wounds were identified utilizing the Jvenn plugin (http://www.bioinformatics.com.cn/static/others/jvenn/). Subsequently, the roles of overlapping DEGs and specific DEGs were evaluated on the DAVID Bioinformatics Resources platform (https://david.ncifcrf.gov/).

Animal Experiment for Wound Healing
ICR male mice (18–22 g) came from Yisi Experimental Animal Technology Co., Ltd. (Certificate No. SCXK-2020-0001, Changchun, China) and were fed in a standard laboratory environment (22 ± 2 °C, 60 ± 5% humidity, and a 12 h light/12 h dark cycle). All experimental procedures were approved by the Animal Care and Use Committee of Jilin Agricultural University and strictly followed the NIH Guide for the Care and Use of Laboratory Animals. Type 2 diabetes mellitus (T2D) in mice was induced as previously reported. After one week of adaptive feeding, the normal control (NC) group was fed a normal diet, while the other mice were fed a high-sugar and high-fat diet (HSFD; basic food 49.5%, sucrose 25%, lard oil 15%, milk powder 5%, yolk powder 5%, and sodium cholate 0.5%) for 31 days (Figure S1A). HSFD mice were fasted for 12 h before intraperitoneal injection (i.p.) with STZ in citric acid buffer (0.1 M, pH 4.3) daily for 3 days (80, 70, and 60 mg/kg, successively), whereas NC mice were injected with vehicle solution. Mice with fasting blood glucose (FBG) ≥ 11.1 mM and typical clinical polydipsia, polyphagia, polyuria, and weight loss after one week were considered T2D.
On the day of surgery, all mice were treated with chloral hydrate (400 mg/kg, i.p.) and a full-thickness circular wound with a radius of 0.5 cm was induced when hair was removed from the dorsal region. The T2D animals with wounds were randomly divided into 3 groups (n = 12/group): the T2D group, the CAH group, and the CAVBPH group. The NC and T2D groups were treated with 0.2 mL of PBS, while the CAH and CAVBPH groups received the same volume of CAH and CAVBPH treatment, respectively. Wound size was observed on days 0, 3, 9, and 15 post-surgery and the wound healing rate was determined using the Image J software.

**Histopathological Analysis**

The wound along with the surrounding new skin was fixed in 4% paraformaldehyde for 24 h and further processed for hematoxylin and eosin (H&E) staining, Masson’s trichrome (Masson) staining, and immunohistochemical (IHC) staining. In this study, anti-CD31 (ab182981, Abcam), anti-CD68 (BA3638, BOSTER), anti-PCNA (BM0104, BOSTER), and anti-α-SMA (BM0002, BOSTER) reagents were used for IHC staining of tissue sections as previously described.16 Images were obtained using a Nikon upright microscope (E100, Tokyo, Japan). The collagen content was analyzed using the Image J software and the integrated optical density (IOD) values of different proteins were identified by the Image-Pro Plus 6 software.

**Western Blot**

Skin wound tissues were harvested and homogenized in cold RIPA lysis buffer, and the supernatant of the sample was collected after centrifugation at 4 °C. BCA protein kit was used to measure protein concentration in different groups. Western blot analysis was performed for antibodies p-PI3K (#4228, Cell Signaling, China), PI3K (67071-1-lg, Proteintech, China), p-AKT (ARG51559, Arigobio, China), AKT (60203-2-lg, Proteintech, China), p-mTOR (ARG40666, Arigobio, China), mTOR (10745-1-AP, Proteintech, China), HIF-1α (20960-1-AP, Proteintech, China), VEGFA (19003-1-AP, Proteintech, China), IL-1β (26048-1-AP, Proteintech, China), TNF-α (17590-1-AP, Proteintech, China) and β-actin (66009-1-lg, Proteintech, China) as previously reported.25 β-actin was used as an internal standard. The protein blots were visualized by enhanced chemiluminescent assay and quantified using the Image J software.

**16S rRNA Sequencing of Skin Flora**

Before the mice were euthanized on day 15, the skin microbial samples of each group (n = 10) were collected by gently swabbing the wound along with the surrounding new skin with sterile cotton swabs soaked in normal saline. 16S rRNA sequencing was performed by Shanghai Personal Biotechnology Co., Ltd. The outcomes were analyzed on the Gene cloud platform (https://www.genescloud.cn/home).

**Statistical Analysis**

Triplicate experiments were performed unless stated otherwise. Data are presented as mean ± standard deviation (SD). IBM SPSS Statistics 23 software was applied to compare the differences between groups by one-way analysis of variance with the LSD test. Spearman correlation coefficients between the composition of T2D wound microflora and the expression of active proteins were evaluated by utilizing the OmicStudio platform (https://www.omicstudio.cn/tool). p < 0.05 was considered statistically significant.

**Results**

**VBPs Contained Abundant Short-Chain Peptides**

LC-MS/MS results showed that 89.78% of VBPs had a molecular weight of less than 1 kDa. VBPs had 372 sequences (LKECCDKPV, VGYP, LFP, EHF, FPH, etc.), 96.5% of them were short-chain peptides (no more than 10 amino acids) (Table S1). Only very few sequences have been found with anti-ACE, antibacterial, and antioxidant activities in the BIOPEP database (https://biochemia.uwm.edu.pl/biopep/start_biopep.php), and the bioactivities of most peptides remain unknown.
Hydrogel Had Excellent in vitro Physicochemical Properties

The preparation process of CA VBPH was shown in Figure 1A. Hydrogel maintained liquid at room temperature and transformed into gel when the temperature increased to human body temperature. The interaction between CS and β-GP mainly involved intermolecular hydrogen bonds and electrostatic effect, and the eggshell-like cross-linked structure between SA and Ca²⁺ also improved the mechanical properties of hydrogel. All hydrogels had a highly porous structure at the microscopic level and the introduction of VBPs did not make a difference in the structure of porosity (Figure 1B), which not only built an environment for cell attachment and migration but also satisfied gas exchange of the wound bed.

The IR spectra were presented in Figure 1C. In the IR spectra of CAH and CA VBPH, the N-H vibration absorption peak (1568 cm⁻¹) of -NH₃⁺ in CS molecule and the asymmetric and symmetric stretching vibration peaks (1082 and 973 cm⁻¹) of PO₄³⁻ in β-GP molecule were obviously weakened, showing that electrostatic effect occurred between -NH₃⁺ and PO₄³⁻. The stretching vibration of C=O (1633 and 1420 cm⁻¹) and C-O (1032 cm⁻¹) and the deformation vibration of O-H (1320 cm⁻¹) associated with SA in hydrogel were weakened when cross-linked with Ca²⁺. For VBPs, the absorption peaks at 1630 and 1582 cm⁻¹ corresponded to the characteristic parameters of amide I and II, respectively; the bands at 1455, 1404, and 1111 cm⁻¹ were, respectively, related to C-H bending vibration, O-H bending vibration, and C-O stretching vibration. The IR spectrum of CA VBPH displayed obvious absorption bands of VBPs (1582 cm⁻¹, 1404 cm⁻¹, 1111 cm⁻¹, etc.), which suggested that VBPs have been loaded into the hydrogel.

Figure 1D showed that CA VBPH and CAH quickly absorbed water and achieved swelling ratios of over 200% within 1 h due to their hydrophilicity and interconnected porous structure, which was important to cover the wound and obtain a sustained drug delivery. Moreover, with the increase of incubation time, the mass residue of hydrogels gradually decreased (Figure 1E). On day 14, the residual weights of CA VBPH and CAH dropped from 100% to 59.26% and 56.85%, respectively. The appropriate anti-degradation performance of hydrogels also demonstrated their long-time covering and protective effects on wounds.

The three-dimensional porous structure of the dressing facilitates the rapid delivery of drugs or cells to the affected tissue. As presented in Figure 1F, CA VBPH showed a burst release with a release amount of 44.80% within 12 h, indicating the low molecular weight of short-chain peptides and the high porosity of CA VBPH might increase the diffusion coefficient of VBPs. VBPs in CA VBPH exhibited a nearly linear continuous release (74.79%) within 12–48 h. After 48 h, the release speed slowed down but this process was still maintained and reached 82.24% after 240 h, demonstrating that CA VBPH had desirable sustained release behavior.

Hydrogel Exerted Antioxidant Activity and Biosafety in vitro

An increasing number of reports have shown that skin dressings with antioxidant activity could improve the therapeutic effect for tissue repair. In this study, CAH showed almost no antioxidant activity, DPPH and ABTS radicals scavenging abilities of CA VBPH were 10.97% and 45.81%, respectively, while the antioxidant activities of VBPs and GSH were superior to those of CAH and CA VBPH (p < 0.001), indicating that the antioxidant capacity of CA VBPH was related to the addition of VBPs (Figure 2A).

Wound dressings are usually indirect or direct contact with wounds and blood, so biocompatibility is regarded as a prerequisite for the clinical application of dressings. Dressings are generally considered clinically safe with a hemolysis rate of less than 5%. The results showed that obvious hemolysis appeared in the positive control, but the hemolysis rates of VBPs, CA VBPH, and CAH were not more than 1.5% (Figure 2B). Cytotoxicity is an important index for the practical application of hydrogel dressings. The cell viability maintained almost unchanged with the increase of hydrogel concentration from 50 μg/mL to 500 μg/mL as presented in Figure 2C. The cell vitality of the CAH and CA VBPH groups was 81.29% and 86.74%, respectively, when the concentration was 5000 μg/mL. Samples with more than 75% cell viability are generally regarded as non-cytotoxic referring to ISO 10993–5: 1999 (GB/T 16886.5–2003). Hence, CAH and CA VBPH are considered to possess negligible toxicity and good biosafety.

Bioinformatics Analysis of Acute Wound and Diabetic Wound

A total of 245 up- and 190 down-regulated DEGs were identified in acute skin wounds, and 609 DEGs (299 up- and 310 down-regulated) were found from diabetic skin wounds (Figure 3A and B). Besides, 339 overlapping DEGs between
Figure 1 Preparation and in vitro performance evaluation of hydrogels. (A) The preparation process of hydrogels. (B) SEM micrograph (scale bar = 50 μm). (C) Infrared (IR) spectroscopy. (D) Swelling test. (E) Biodegradation test. (F) In vitro release profile of VBPs from CAVBPH.
acute skin wounds and diabetic skin wounds (199 up- and 140 down-regulated) and 270 specific DEGs of diabetic skin wounds (100 up- and 170 down-regulated) were obtained through Venn diagram analysis (Figure 3C).

As shown in Figure 3D, Gene Ontology-biological processes (GO-BP) of overlapping DEGs were mainly enriched in cell adhesion, positive regulation of cell proliferation, neutrophil degranulation, and inflammatory response. The terms of overlapping DEGs in Kyoto Encyclopedia of Genes and Genomes (KEGG) were predominantly enriched in the PI3K-AKT signaling pathway, IL-17 signaling pathway, TNF signaling pathway, and NF-κB signaling pathway (Figure 3E). The BP category of specific DEGs in diabetic wounds was significantly related to positive regulation of transcription, negative regulation of apoptotic process, and positive regulation of gene expression (Figure 3F). In addition, the specific DEGs of diabetic wounds were predominantly enriched into the PI3K-AKT signaling pathway, cAMP signaling pathway, HIF-1 signaling pathway, and JAK-STAT signaling pathway (Figure 3G).

**CAVBPH Accelerated T2D Wound Healing**

During the in vivo experiment, T2D mice still maintained a higher FBG level than 11.1 mM and the typical symptoms of clinical polydipsia, polyphagia, polyuria, and weight loss after injection of STZ. As shown in Figure S1B, the pancreatic islets of NC mice showed normal architecture. In contrast, considerable degenerative changes in pancreatic islets were detected in T2D mice, including the size and number of islets decreased and islet β-cells damaged. All these results demonstrated that the stable T2D model has been successfully established.

As shown in Figure 4A, the wound size in all groups decreased gradually over time. Compared with the NC group, the T2D group displayed a slower wound shrinkage due to the influence of diabetes, and exudate was still found on the wound surface on day 9. Topical application of CAH and CAVBPH significantly accelerated diabetic wound recovery compared to the T2D group. Quantitative analysis of wound healing confirmed the above results (Figure 4B). The healing rate in the T2D group was less optimistic than that in the NC group ($p < 0.01$ or $p < 0.001$). Compared to the T2D group, the CAH group showed a better healing effect on days 9 and 15 ($p < 0.001$), while the CAVBPH group had the best...
wound shrinkage than the T2D and CAH groups on all observed days ($p < 0.01$ or $p < 0.001$). At day 15, the wounds of the CAVBPH group (96.55% of healing rate) and the NC group (91.77% of healing rate) were basically healed and covered with hair, while the healing rates of the CAH and T2D groups were 84.23% and 63.81%, respectively.

CAVBPH Promoted Remodeling of the Microstructure of Wound Tissue

Concerning microstructure, T2D wound showed necrosis, inflammatory cell infiltration, and no epidermis on day 15 (Figure 5A). Moreover, the T2D group had lower levels of granulation tissue and new capillaries than NC mice. As compared with the T2D group, the CAVBPH and CAH groups presented more regenerated epidermis, granulation tissue, and new capillaries but negligible inflammatory cell infiltration. We also found that the CAVBPH group had more mature proliferating cells and skin appendages than the CAH group. Masson staining results showed that the treatment of CAH and...
CA VBPH increased collagen formation compared to the T2D group (Figure 5A). Notably, the collagen arrangement of the CA VBPH group was denser, ordered, and more mature than that of the CAH group. Quantitative analysis displayed that hydrogel treatment significantly increased collagen content in diabetic wounds ($p < 0.01$ or $p < 0.001$; Figure 5B). Moreover, the CA VBPH intervention recovered the collagen fibers in the diabetic wound tissues to the NC level, which was superior to
the CAH treatment ($p < 0.01$). These results provided evidence that CAVBPH treatment accelerates diabetic wound healing by regulating inflammation and promoting re-epithelialization, angiogenesis, and collagen deposition.

CAVBPH Up-Regulated CD31, PCNA, and α-SMA Proteins but Down-Regulated CD68 Expression in Wound

CD31, a marker of vascular endothelial cells, was used to assess new capillaries during wound healing. PCNA is one of the most prominent markers for cell proliferation, while α-SMA is considered to be directly linked to the production of extracellular matrix (ECM) proteins. In the current study, the expression of CD31, PCNA, and α-SMA in the T2D group was poorer than that in the NC group ($p < 0.001$; Figure 6). Conversely, these proteins were significantly up-regulated in the CAVBPH group compared with the T2D group ($p < 0.01$ or $p < 0.001$). CD68 is a glycoprotein highly expressed on monocytes and macrophages, especially during the inflammatory phase of skin wounds. Treatment with CAH or CAVBPH dramatically suppressed the abnormal expression of CD68 protein in T2D wound tissue ($p < 0.001$). These results suggested that hydrogel treatment, especially CAVBPH, could up-regulate CD31, PCNA, and α-SMA proteins, while down-regulating CD68 expression, i.e., promote angiogenesis, cell proliferation, and ECM secretion as well as alleviate persistent inflammation during T2D wound healing.

CAVBPH Activated the PI3K/AKT/mTOR/HIF-1α/VEGFA Signaling Pathway

According to the results of bioinformatics analysis (Figure 3), we speculated that the PI3K/AKT and HIF-1 pathways might have a great potential effect on the healing of T2D wounds, thus the relative expression of active factors related to the PI3K/AKT/mTOR/HIF-1α/VEGFA pathway in wound tissue was further detected. As displayed in Figure 7A–D, the phosphorylation levels
Figure 6 IHC staining of CD31, PCNA, α-SMA, and CD68 proteins. (A) Representative images for CD31, PCNA, α-SMA, and CD68 (scale bar = 200 μm). (B–E) Quantification analysis of CD31, PCNA, α-SMA, and CD68 proteins expression in wound tissues, respectively. *p < 0.05, **p < 0.01, and ***p < 0.001.
of PI3K, AKT, and mTOR proteins in the T2D group were noticeably lower than in the NC group ($p < 0.01$ or $p < 0.001$). Conversely, the CAVBPH treatment elevated the relative expression of p-PI3K/PI3K, p-AKT/AKT, and p-mTOR/mTOR in diabetic wounds ($p < 0.05$, $p < 0.01$, or $p < 0.001$), demonstrating that the CAVBPH activated the PI3K/AKT/mTOR pathway. We also found that T2D hindered the expression of downstream molecules HIF-1α and VEGFA in skin wound tissues ($p < 0.001$), whereas the CAVBPH intervention elevated the relative expression levels of these ($p < 0.001$; Figure 7A, E, and F). Unfortunately, we did not identify significant differences in the expression of the aforementioned proteins between the CAH group and the T2D group, except for VEGFA ($p < 0.05$). The above data proved that the CAVBPH activated the PI3K/AKT/mTOR/HIF-1α/VEGFA pathway in vivo, which contributed to the rapid healing of diabetic wounds.

**CAVBPH Inhibited the Overexpression of Inflammatory Factors**

To investigate the inflammatory response of T2D wounds, we detected the protein expression of inflammatory cytokines IL-1β and TNF-α by Western blot based on the results of bioinformatics analysis. As presented in Figure 8, the relative expression of TNF-α and IL-1β in the T2D group was dramatically risen as compared to the NC group ($p < 0.001$). However, CAH and CAVBPH treatments significantly reduced the expression levels of IL-1β and TNF-α in T2D wound tissues ($p < 0.05$, $p < 0.01$, or $p < 0.001$). In addition, the CAVBPH intervention remarkably down-regulated the expression of IL-1β and TNF-α when compared with the CAH group ($p < 0.01$). This was consistent with the CD68 staining results, that is, CS/SA hydrogel loaded with VBPs was beneficial for reducing the inflammation level at T2D wound sites.

**CAVBPH Restored the Diversity and Structure of Skin Microbiota at T2D Wounds**

The rarefaction curve and species accumulation curve showed that the increase of species numbers tended to flatten with the increase of sequencing depth and samples, respectively, indicating that the data quality and sample number were sufficient (Figure S2). The number of OTUs is regarded as an index reflecting the abundance of microbial species in the samples. As presented in Figure 9A, 3989, 3627, 3110, and 3879 OTUs were in the NC, T2D, CAH, and CAVBPH groups, respectively. 350 OTUs were co-overlapping in the above four groups. α-diversity, which reflects the richness
and diversity of each sample species, was assessed by the Observed species index and the Simpson index (Figure 9B).

The results showed that T2D caused a decrease in the α-diversity of skin microbiota, whereas the CA VBPH intervention increased the Observed species ($p = 0.065$) and Simpson ($p = 0.0041$) indices when compared with the T2D group.

Principal coordinate analysis represents the community differences between groups and reflects the β-diversity of skin flora in mice. (A) Venn diagrams of operational taxonomic units (OTUs) in the NC, T2D, CAH (T2D+CAH), and CAVBPH (T2D+CAVBPH) groups. (B) α-diversity analysis. (C) β-diversity analysis. (D) Relative abundance of community at the phylum level. (E) Percent of community abundance at the genus level. ***$p < 0.01$. 

**Figure 8** Western blot analysis of inflammatory factors TNF-α and IL-1β proteins. (A) Representative protein bands for TNF-α, IL-1β, and β-actin. (B and C) Quantification analysis of TNF-α/β-actin and IL-1β/β-actin. *$p < 0.05$, **$p < 0.01$, and ***$p < 0.001$.

**Figure 9** 16S rRNA sequencing of skin flora in mice. (A) Venn diagrams of operational taxonomic units (OTUs) in the NC, T2D, CAH (T2D+CAH), and CAVBPH (T2D+CAVBPH) groups. (B) α-diversity analysis. (C) β-diversity analysis. (D) Relative abundance of community at the phylum level. (E) Percent of community abundance at the genus level. ***$p < 0.01$. 

The results showed that T2D caused a decrease in the α-diversity of skin microbiota, whereas the CA VBPH intervention increased the Observed species ($p = 0.065$) and Simpson ($p = 0.0041$) indices when compared with the T2D group.
microflora. As shown in Figure 9C, the sample clustering of the T2D group was significantly far away from the NC group. Community differences began to appear between the CAH group and the T2D group, and the CA VBPH group was clearly separated from the T2D group and had the smallest differences with NC mice.

The dominant bacteria among the four groups, at the phylum level, were Firmicutes, Proteobacteria, Actinobacteria, and Bacteroidetes. The relative abundance of Firmicutes, Proteobacteria, Actinobacteria, and Bacteroidetes in the NC group was 34.05%, 62.63%, 1.46%, and 0.85%, respectively. Compared to the NC group, the relative abundance of Firmicutes (80.22%) was increased, while the relative levels of Proteobacteria (16.13%), Actinobacteria (0.90%), and Bacteroidetes (0.09%) were down-regulated in the T2D group. The relative abundance of Proteobacteria (28.21%) and Bacteroidetes (0.11%) was increased, but Firmicutes (70.52%) was decreased in the CAH group when compared with the T2D group. In the CA VBPH group, the relative abundance of Firmicutes (46.96%) was lower while the relative contents of Proteobacteria (48.79%), Actinobacteria (3.47%), and Bacteroidetes (0.30%) were higher than the T2D group. Firmicutes are normally seen as harmful microorganisms that disrupt lipid and glucose metabolism, and a high ratio of Firmicutes to Bacteroidetes (F: B ratio) is thought to be linked to an increased risk of T2D.\textsuperscript{40} Meanwhile, CA VBPH treatment significantly reduced the F: B ratio of T2D mice from 891.3 to 156.5. At the genus level, the relative abundance of Staphylococcus (69.08%) and Weissella (4.33%) in the T2D group displayed a significant elevation (1.72% and 0.27% in the NC group, respectively), but the relative abundance of Vulcanibacterium (7.59%), Cupriavidus (4.19%), Pseudomonas (1.01%), Lactobacillus (1.48%), Anoxybacillus (1.53%), Chelatococcus (1.03%), Aerococcus (0.73%), and Corynebacterium\_1 (0.68%) was decreased (18.78%, 11.61%, 22.74%, 9.58%, 4.53%, 2.92%, 0.12%, and 0.22% in the NC group, respectively). CA VBPH treatment remarkably reversed T2D-induced skin flora dysregulation at the genus level, except for the up-regulation of Weissella in relative abundance. These changes in skin flora may be important pathways by which diabetic wounds maintain long-time ulcers and CA VBPH promotes skin tissue repair.

**Spearman Correlation Between Skin Microbiota and Active Proteins in Diabetic Wounds**

Spearman heatmap showed significant correlations between the dominant skin microbes around diabetic wounds and factors related to angiogenesis and inflammation in the tissue during topical application of CA VBPH (Figure 10). At the phylum level, Firmicutes were negatively correlated with p-PI3K/PI3K, p-AKT/AKT, p-mTOR/mTOR, HIF-1α, and

![Spearman heatmap](https://doi.org/10.2147/JIR.S376692)  
*Figure 10* Spearman correlation analysis between skin microbiota and active proteins related to angiogenesis and inflammation in diabetic wounds. (A) Correlation heatmap of skin microbes and active proteins at the phylum level. (B) Correlation heatmap of skin microbes and active proteins at the genus level. *p < 0.05 and **p < 0.01.
VEGFA, but positively related to IL-1β and TNF-α ($p < 0.05$ or $p < 0.01$); conversely, *Proteobacteria, Bacteroidetes,* and *Patescibacteria* were positively associated with the PI3K/AKT/mTOR/HIF-1α/VEGFA pathway-related factors, but negatively correlated with inflammatory cytokines IL-1β and TNF-α ($p < 0.05$ or $p < 0.01$; Figure 10A). To obtain more information, we further calculated the spearman coefficient at the genus level (Figure 10B). Specifically, *Staphylococcus* was negatively correlated with the expression of related proteins of the PI3K/AKT/mTOR/HIF-1α/VEGFA pathway, while positively related to IL-1β and TNF-α ($p < 0.05$ or $p < 0.01$). *Lactobacillus* was observed to have an opposite correlation trend to *Staphylococcus*. In addition, *Vulcaniibacterium, Cupriavidus, Pseudomonas, Anoxybacillus,* and *Chelatococcus* were also strongly associated with the expression of angiogenesis and inflammation-related factors. Thus, these skin microbes were indirectly or directly linked to the therapeutic ability of CAVBPH on T2D wounds by the skin flora-active proteins axis.

**Discussion**

Skin wound is a common chronic complication of diabetes and one of the leading factors that threaten the physical and mental health of diabetic patients and cause serious economic and social burdens.1–3 Therefore, it is a pressing need to investigate the pathogenesis and effective strategies for delayed wound healing. Studies have reported that bioactive peptides-loaded hydrogels are a promising option for repairing tissue damage.16,20,21,26 Velvet antler has regenerative property, and both velvet antler and velvet antler peptides have been proved to have wound repair activity.19,41 VB derived from velvet antler is rich in proteins and peptides, and TCM believes that its activity is similar to that of velvet antler. In this study, we encapsulated short-chain VBPs with antioxidant activity in a CS/SA hydrogel scaffold and explored its efficacy on T2D skin wound repair for the first time.

Previously, CS/SA polymer was shown to be more efficient in promoting wound repair than either CS or SA alone, and CS/SA composite hydrogel possessed excellent physicochemical properties to support it as a promising wound dressing scaffold.25,42 The results of in vitro evaluation showed that CAVBPH possessed good porous breathability, swelling, biodegradability, antioxidant property, biocompatibility, biosafety, and release behavior of VBPs (Figures 1 and 2). Subsequently, we topically applied CAVBPH to the wound surface, on the one hand, to avoid direct contact of the wound with the outside, and on the other hand to continuously provide VBPs for wound healing. Compared to the T2D and CAH groups, the CAVBPH group had the best curative effect, with a 96.55% wound healing rate on day 15 (Figure 4). Pathological staining revealed the positive influence of CAVBPH on angiogenesis, re-epithelialization, fibroblast proliferation, collagen deposition, and inflammation regulation (Figures 5 and 6). We also found that the therapeutic effect of CAVBPH on T2D skin wounds was better than recently reported CS-based hydrogels, decellularized skin scaffolds, and herbal ointments based on a literature review.16,17,31,43 However, the specific mechanism by which CAVBPH exerts its efficacy remains unknown.

Bioinformatics studies can comprehensively analyze genome-wide profiles of clinical samples and have been extensively applied to explore the pathogenesis and promising therapeutic targets in various diseases.44,45 In this study, bioinformatics analysis found that diabetic wound-specific DEGs were significantly correlated with the PI3K/AKT and HIF-1 signaling pathways (Figure 3). HIF-1α is a vital molecule regulating angiogenesis and is a downstream node of the PI3K/AKT/mTOR pathway. Excessive glucose levels interfere with the activity and stability of HIF-1α, resulting in the failure of VEGFA up-regulation in diabetic wounds, which in turn causes impaired angiogenesis and wound healing.46 CS-based wound dressings have been shown to exert therapeutic effects by activating the PI3K/AKT pathway.25,47 Interestingly, this work confirmed that CAVBPH more prominently exerted a pro-angiogenic effect than CAH via activating the PI3K/AKT/mTOR/HIF-1α/VEGFA pathway (Figure 7).

The development of DFU is accompanied by persistent chronic inflammation. Accumulating evidence suggests that inflammatory responses mediated by IL-1β and TNF-α damage vascular endothelial cells and weaken insulin sensitivity, causing defects in wound healing. On the contrary, silencing TNF-α or blocking IL-1β can effectively improve the healing of DFU by inducing macrophages to polarize into healing-related phenotypes and inhibiting macrophage infiltration.48,49 In this study, CAVBPH intervention significantly down-regulated the levels of IL-1β, TNF-α, and CD68, indicating that CAVBPH had a good therapeutic effect on chronic inflammation of T2D wounds (Figures 6 and 8).
Skin flora is considered critical for protecting humans from pathogens and maintaining the balance of the immune system. Currently, whether topical application of skin dressings can improve skin ulcers by affecting the microflora around diabetic wounds requires further investigation. It is reported that low microflora diversity was associated with some metabolic disorders, while higher overall diversity meant better health. Our results showed that CA VBPH therapy reversed the poor diversity and structure of skin flora induced by T2D and made them tend to a healthy level (Figure 9). Skin symbiotic bacteria Cutibacterium and Corynebacterium contained genes related to porphyrin metabolism, which could theoretically reduce infections caused by Staphylococcus aureus. In addition, a variety of Lactobacillus species have been reported as beneficial bacteria for the prevention and treatment of gastrointestinal and genitourinary system infections. Live Lactobacillus reuteri DSM 17938 was found to have an inhibitory effect on skin pathogenic bacteria (Pseudomonas aeruginosa, Cutibacterium acnes AS12, Streptococcus pyogenes M1, and Staphylococcus aureus). CAVBPH dramatically increased the relative abundance of Corynebacterium_1 and Lactobacillus at the genus level, suggesting the effectiveness of CAVBPH in chronic wounds and the correlation between the regulation of skin microbiota and the improvement of skin ulcers. Moreover, we believe that the purposeful regulation and improvement of skin microbiota will facilitate the clinical treatment of diabetic wounds. Interestingly, we also noticed that changes in the relative abundance of some dominant microorganisms were significantly associated with the efficacy of CAVBPH in enhancing angiogenesis and alleviating persistent inflammation. However, further studies must be performed to analyze the exact biological mechanism by which CAVBPH improves diabetic wound healing through the skin flora-active protein axis.

**Conclusion**

This is the first study to widely analyze the therapeutic effect of VBPs delivered to T2D skin wounds using a CS/SA hydrogel scaffold, which provides novel insights into the modern application of VBPs and the treatment of diabetic ulcer. The results suggest that CAVBPH, as a promising alternative therapy for diabetic wounds, could promote angiogenesis and cell proliferation and alleviate inflammation via activating the PI3K/AKT/mTOR/HIF-1α/VEGFA pathway and reversing the expression of TNF-α and IL-1β. Also, CAVBPH restored the diversity and structure of T2D skin flora. The excellent therapeutic efficacy of CAVBPH on chronic cutaneous trauma may be related to numerous short-chain VBPs, which needs further study.

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**Disclosure**

The authors report no conflicts of interest in this work.

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