

Identification of Active Compounds and Mechanism of Huangtu Decoction for the Treatment of Ulcerative Colitis by Network Pharmacology Combined with Experimental Verification

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
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Introduction: Huangtu decoction (HTD) has been widely used in the treatment of gastrointestinal bleeding, ulcerative colitis (UC) and gastrointestinal tumors in China, but its active compounds and mechanism are still not clear yet. The present research aimed to identify the active compounds and mechanism of HTD for the treatment of UC.

Methods: Firstly, the chemical compounds of HTD were qualitatively identified based on Q Exactive Orbitrap LC-MS/MS, and their potential targets were predicted through SwissTargetPrediction. Secondly, the differential expressed genes (DEGs) in colon tissues of UC patients and normal controls were retrieved from the GEO database. Thirdly, the overlapping targets of DEGs and the predicted targets were obtained and subjected to GO and KEGG analysis. Finally, the key targets in the most significantly enriched pathway were verified by in vivo experiment, and the protein and mRNA expressions of matrix metalloproteinase-1 (MMP1), MMP3, MMP7, MMP9 and MMP12 were determined by immunohistochemistry (IHC), Western blotting (WB) and quantitative real-time-PCR (qRT-PCR).

Results: A total of 47 compounds were identified and 29 overlapping targets were obtained from HTD extract. The most significantly enriched pathway of overlapping targets involved was MMP. HTD improved the pathological damage in colon tissues of DSS-induced UC model and significantly decreased the serum levels of IL-1 β and IL-6. The protein and mRNA expressions of MMP1, MMP3 and MMP9 in colon tissues were significantly decreased after HTD treatment.

Conclusion: HTD treatment can alleviate the colonic inflammation via inhibiting MMPs including MMP1, MMP3 and MMP9.

Keywords: bioinformatic analysis, ulcerative colitis, Huangtu decoction, matrix metalloproteinases

Introduction

Ulcerative colitis (UC) is a kind of chronic colitis involving colorectal mucosa and submucosa, which is mainly caused by chronic inflammation and ulceration. The incidence of UC is increasing worldwide,^{1,2} and its clinical manifestations are primarily diarrhea, mucous purulent and bloody stool with abdominal pain and acute severe symptoms. It is estimated that the total annual economic burden of UC ranges from

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Received: 8 July 2021
Accepted: 17 September 2021
Published: 29 September 2021



\$8.1 to \$14.9 billion in the United States and from €12.5 to €29.1 billion in Europe.³ The duration and extensive feature of disease are considered as important factors associated with an increased risk of colorectal cancer in patients with UC.⁴⁻⁶ Intense efforts have been made to elucidate the pathogeny of UC, but its molecular mechanisms are still poorly understood. Drug treatment is the first choice for UC, and the available drugs include mesalazine, corticosteroids, immunosuppressive drugs, and monoclonal antibodies to TNF- α currently in Western world.⁶ Nevertheless, the limitations of these drugs are also obvious, such as poor therapeutic effects, high price and bothersome side effects.⁷

In recent years, Traditional Chinese Medicine (TCM) has been gradually recognized as an effective method for treating various diseases not only in China and Southeast Asia, but also in many Western countries.^{8,9} TCM also plays an important role in drug development, providing abundant novel compound lead sources. Huangtu Decoction (HTD) was first recorded in the Treatise on “JingGuiYaoLue (Synopsis of Golden Chamber)” written around 205 A.D by Zhang Zhongjing, a famous medical scientist in the Eastern Han Dynasty. Its prescription is composed of 7 medicinal herbs, including *Terra Flava Usta*, *Rehmanniae Radix*, *Scutellariae Radix*, *Aconiti Lateralis Radix Praeparata*, *Asini Corii Colla*, *Atractylodis Macrocephalae Rhizoma* and *Glycyrrhizae Radix et Rhizoma*. HTD has been mainly used in the treatment of bleeding symptoms, such as blood under stool, hematemesis, bleeding, metrorrhagia and dull blood color, and highly respected by doctors for thousands of years in China.^{10,11} In

modern times, HTD is often used to treat gastrointestinal bleeding, chronic UC, gastrointestinal tumor and so on.^{12,13} However, the active compounds and mechanism of HTD for the treatment of UC are still not clear.

During the last decades, bioinformatics analysis was widely used to screen genetic alterations at the genome level,^{14,15} so as to identify the differentially expressed genes (DEGs). Besides, network pharmacology has been successfully introduced to reveal the mechanisms of TCM, which coincides with the connotation of multi-component, multi-targets, and multi-pathways holistic strategy.^{16,17}

In the present study, network pharmacology, bioinformatics analysis and in vivo experiments were combined to uncover the possible targets and mechanisms of HTD for treating UC, and the detailed flowchart of the study design is shown in Figure 1.

Materials and Methods

Reagents and Materials

Terra Flava Usta (Batch No: 2006002) was purchased from AnguoJuyaotang Pharmaceutical Co., Ltd. *Rehmanniae Radix* (Batch No: Y2105018), *Scutellariae Radix* (Batch No: Y2007053), *Aconiti Lateralis Radix Praeparata* (Batch No: Y062011002), *Asini Corii Colla* (Batch No: Y052105035), *Atractylodis Macrocephalae Rhizoma* (Batch No: Y2006043) and *Glycyrrhizae Radix et Rhizoma* (Batch No: Y2008055) were purchased from Sichuan Neautus Traditional Chinese Medicine Co., Ltd.

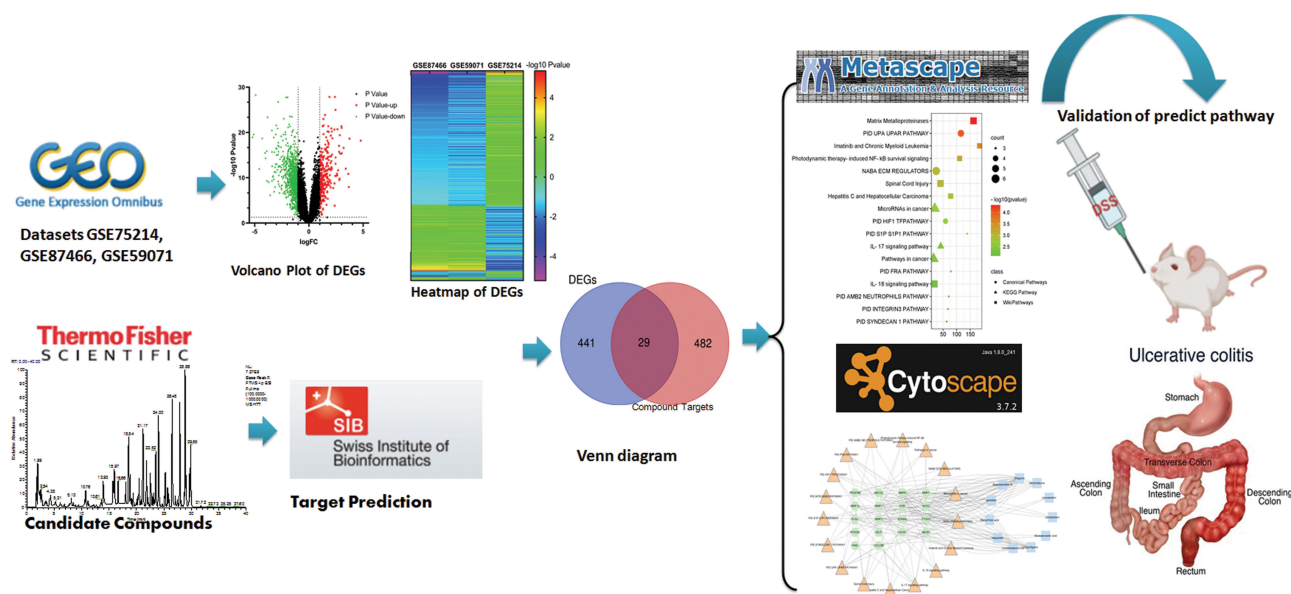


Figure 1 Flowchart of the analysis strategy in the study.

HPLC grade methanol was provided by Fisher Scientific (USA). Other chemicals were of analytical grade.

Animals and Diets

The C57BL/6 male mice were provided by the SPF (Beijing) Biotechnology Co., Ltd. (DCXK (Beijing) 2019–0010). The mice (6–8 weeks old, n=24) were randomly divided into 3 groups, and acclimated in independent ventilation cages (IVC) for 6 days with free access to standard mouse chow and tap water in the SPF facility under controlled temperature ($22 \pm 2^\circ\text{C}$), humidity ($55\% \pm 10\%$), and 12 light-dark cycle. Animal experiments were performed according to the Recommended Guidelines for the Care and Use of Laboratory Animals issued by the Chinese Council on Animal Research and approved by the Ethics Committee of Chengdu University of TCM.

Preparation of the HTD Extracts

The HTD extracts were prepared according to the recorded method in “JingGuiYaoLue”. Briefly, the seven herbs *Terra Flava Usta* (40 g), *Rehmanniae Radix* (15 g), *Scutellariae Radix* (15 g), *Aconiti Lateralis Radix Praeparata* (15 g), *Asini Corii Colla* (15 g), *Atractylodis Macrocephalae Rhizoma* (15 g) and *Glycyrrhizae Radix et Rhizoma* (6 g) were mixed with purified water, and *Aconiti Lateralis Radix Praeparata* were soaked for 12 hours, while other herbs were soaked for half an hour, then they were boiled for an hour. The filtrate were collected and centrifuged at 3000 rpm for 5 min. The supernatant was harvested and steamed dry, and the residue was dissolved to 5 mL with methanol. The sample was filtered through a 0.22 μm microporous membrane for subsequent testing.

Identification of Active Compounds in HTD

The active compounds of HTD extracts were qualitatively identified by Q Exactive Orbitrap LC-MS/MS (Thermo Fisher Scientific, Waltham, MA, USA). The XBridgeBEHC18 (2.1 \times 100 mm, i.d.; 2.5 μm , Waters, MA, USA) was applied for sample separation at 35 $^\circ\text{C}$. Methanol (A) and 0.1% formic acid-water (B) (95:5) were used as mobile phase. Detection wavelength was set at 280 nm. The flow rate and injection volume were set as 0.2 mL/min and 1.0 μL , respectively. Nitrogen was applied as auxiliary gas and sheath gas at a flow rate of 12 L/min. The mass determination was carried out based on positive and negative scanning mode with the m/z of 100–1000.

Acquisition of Corresponding Targets of Active Compounds

All the chemical structures of the identified active compounds were imported into SwissTargetPrediction (<http://www.swisstargetprediction.ch/>) to predict the targets.¹⁸ Targets of these compounds were collected through high-throughput screening and reverse docking.

Acquisition of UC Targets

Gene Expression Omnibus (GEO) (<https://www.ncbi.nlm.nih.gov/geo/>), a public functional genomics data repository, was used to establish UC targets library.¹⁹ Keywords such as “ulcerative colitis” or “ulcer colonitis” were used to search for the known targets related to the pathogenesis of UC from GEO database, so as to find out the difference in gene expression datasets between intestinal tissue samples from patients with UC and normal intestinal tissue samples. Finally, 3 datasets of UC were downloaded from the GEO database: GSE75214 on the (HuGene-1_0-st) Affymetrix Human Gene 1.0 ST Array (GPL6244 platform), including colon tissue samples from 97 UC patients and 11 healthy people; GSE87466 on the (HT_HG-U133_Plus_PM) Affymetrix HT HG-U133+ PM Array Plate (GPL13158 platform), including colon tissue samples from 87 UC patients and 21 healthy people; GSE59071 on the (HuGene-1_0-st) Affymetrix Human Gene 1.0 ST Array (GPL6244 platform), including colon tissue samples from 97 UC patients and 11 healthy people.

The datasets downloaded from the GEO database were preprocessed by background correction, quantile normalization and base logarithm conversions. Finally, the difference in gene expression between normal human and UC patients was clarified. In this study, log FC (fold change) ≥ 1 and adjusted p-value < 0.01 considered that the difference was statistically significant, and the corresponding genes were identified as DEGs, listed in the UC target library.

GO and Pathway Enrichment Analysis

The GO and pathway enrichment analysis was performed through Metascape (<https://metascape.org/>).²⁰ GO functionally annotates key genes into 3 main terms, ie cellular components (CCs), molecular functions (MFs), and biological processes (BPs). Besides, CytoscapeClueGO plugin was used to further analyze the enrichment of BPs. Pathway enrichment analysis unveiled the possible BPs with key targets. In addition, the bubble chart of GO and

pathway enrichment analysis were performed on the bioinformatics platform (<http://www.bioinformatics.com.cn/>).

Construction of Compound-Target-Pathway Network

To characterize the therapeutic mechanisms of HTD for UC, the compound-target-pathway network was constructed using Cytoscape version 3.7.2.²¹ In the network, the nodes with different colors and shapes represented the drug, compounds, disease, target genes, or disease-related pathways, respectively, and an “edge” was an association between the nodes.

Experimental Validation

Dextran Sodium Sulfate (DSS)-Induced Acute UC Model

Following the previous methods of Mariano et al and Meurer et al^{22,23} after adaptive feeding for 6 days, the mice in control group (n=8) were fed with water, while those in model group (n=8) and DSS + HTD group (n=8) freely drank 3% DSS (36,000–50,000 Da) solution (prepared by dissolving 3 g of DSS in 100 mL of deionized water). The drinking amount in each group was recorded, and DSS solution was updated every day. Furthermore, the mice in DSS + HTD group were administrated with 2.5 g/kg HTD extracts once a day according to their weight. The drinking amount, body weight, stool properties and blood in stool of mice in each group were recorded every day, and the colitis scores were given by weight loss and stool conditions shown in Table 1. After investigations for one

week, the mice in the 3 groups were euthanized, and the serum and colon tissues were collected for further analysis.

Enzyme-Linked Immunosorbent Assay (ELISA)

The levels of inflammatory factors were detected by ELISA kits. The mouse IL-1 β ELISA kit (Multi Sciences (Lianke) Biotech, CO., LTD., Hangzhou, China), and mouse IL-6 ELISA kit (Multi Sciences (Lianke) Biotech, CO., LTD., Hangzhou, China) were utilized in accordance with the directions of the manufacture.

Histopathology

The colon samples were cut into small pieces, fixed in 4% paraformaldehyde, embedded in paraffin, and stained with hematoxylin and eosin (H&E). Then histopathological changes were examined under a microscope.

Western Blotting (WB)

Proteins were extracted from the colon tissues using RIPA Lysis Buffer (Wuhan Servicebio Technology Co., Ltd, China) and the protein concentration was determined by bicinchoninic acid (BAC) method on the enzyme labelling instrument (Thermo Fisher Scientific, USA). The SDS-PAGE was used to separate the protein, and subsequently transferred to polyvinylidene fluoride (PVDF) membranes. The membranes were first blocked for 1h with 5% BSA at room temperature, and then probed with primary antibodies of MMP1 (Affinity, dilution of 1:1000), MMP3 (Abcam, dilution of 1:5000), MMP7 (Abcam, dilution of 1:1000), MMP9 (Abcam, dilution of 1:1000), MMP12 (Proteintech, dilution of 1:1000) and β -Actin (Affinity, dilution of 1:5000) at 4°C for overnight and then incubated with HPR-conjugated antibody at 37°C for one hour. Finally, the protein bands were recorded by ELC (Thermo Fish Scientific, USA) and quantified by Image J software to evaluate the protein relative expressions compare with β -actin.

Quantitative Real-Time-PCR (qRT-PCR)

The mRNA expression of UC was detected by qRT-PCR. The colon tissues in each group of mice were pulverized in liquid nitrogen before RNA extraction. The total RNA was isolated and reversely transcribed into cDNA using the RT-qPCR EasyTM (One Step)-SYBR Green I kit (Foregene; Chengdu, China), followed by synthesis via reverse transcription using the Direct RT Mix in the kit, and amplification using Direct qPCR Mix-SYBR of kit

Table 1 Scoring Standards of Ulcer Colitis

Score	Weight Loss (%)	Stool Consistency	Blood in Stool
0	None	Normal	None
1	None	Normal	Occult+
2	1–5	Normal	None
3	1–5	Soft	None
4	1–5	Soft	Occult+
5	1–5	Diarrhea	Occult+/Gross
6	5–10	Soft	Occult+/Gross
7	5–10	Diarrhea	Occult+/Gross
8	10–20	Soft	Occult+/Gross
9	10–20	Diarrhea	Occult+/Gross
10	>20	Diarrhea	Occult+/Gross

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according to the manufacturer's instructions. Then qPCR was carried out on a Bio-Rad real-time PCR system (CFXConnect, USA) for 40 cycles. The expression levels of target genes were normalized to the control house-keeping gene GAPDH. The gene expression was analyzed by the $2^{-\Delta\Delta Ct}$ method. Primer sequences are shown in Table 2.

Immunohistochemistry and Staining Intensity

Immunolabeling for matrix metalloproteinase-1 (MMP1), MMP3, MMP7, MMP9 and MMP12 was performed on fixed and paraffin-embedded sections. Briefly, following deparaffinization and rehydration, slides were subjected to antigen retrieval in citrate buffer (pH 6.0) for 20 min in a steamer, rinsed in PBS (0.01 mol/L phosphate buffer containing 0.15 mol/L NaCl, pH7.4), and incubated for 10 min in 3% H₂O₂ to block endogenous peroxidases. After that, the sections were incubated with anti-MMP1 (1:200; Bioss, China), anti-MMP3 (1:200; Bioss, China), anti-MMP7 (1:200; Bioss, China), anti-MMP9 (1:200; Bioss, China), and anti-MMP12 (1:200; Bioss, China) primary antibodies at 4°C overnight in a wet box, and then with HRP-labeled anti-rabbit secondary antibodies for 30 min, followed by counterstaining with hematoxylin, rehydration and mounting. Staining intensity was quantified by Image-Pro Plus 6.0 (Media Cybernetics) and expressed as mean using the formula "mean density = integrated optical density/area of interest".

Statistical Analysis

Each experiment was repeated 3 times, and the results were expressed as mean \pm standard deviation (SD). Significant differences between means were compared using the Bonferroni and Dunnett's T3 test in the case of homogeneity of variance with the p-value <0.05.

Table 2 Primer Sequences for qRT-PCR

Gene	Forward Primer (5'-3')	Reverse Primer (5'-3')
MMP1	ACTGCCAAATGGGCTTGAAG	TTCCCTTTGAAAACCGGACTT
MMP3	GAGGCATCCACACCCTAGGTT	TCAGAAATGGCTGCATCGATT
MMP7	GAGTGAGCTACAGTGGGAACA	CTATGACGCGGGAGTTTAACAT
MMP9	CCCTTGTGCTCTTCCCTGGA	TCTGCCACCCGAGTGTAAACC
MMP12	TTGGATTATTGGAATGCTGC	GCACATTTTGATGAGGCAGA
GAPDH	AGCCACATCGCTCAGACAC	GCCCAATACGCCAAATCC

Results

Compounds of HTD Extract and Corresponding Targets

The main compounds of HTD extracts were analyzed qualitatively based on Q Exactive Orbitrap LC-MS/MS. The total ion chromatogram (TIC) is shown under positive-ion polarity mode (A) and negative-ion polarity mode (B) in Figure 2.

More than 60 major compounds were eluted by the Q Exactive Orbitrap LC-MS/MS system from the HTD extracts within 30 min. Among them, a total of 47 compounds were finally identified by comparing with the known chemical constituents in the reported literature data, and deduced according to their mass spectrometry and fragment ion characteristics, as shown in Table 3 and Figure 3.

A total of 511 corresponding targets of compounds were acquired from Swiss Target Prediction.

Prediction of Targets of HTD for UC

After data preprocessing, a total of 470 DEGs were extracted from 3 datasets (GSE87466, GSE59071, and GSE75214) downloaded from the GEO database, as shown in the volcano map and hot map (Figure 4A and B). Finally, 154 upregulated and 316 downregulated genes were identified in UC tissues compared with normal colon tissues.

Finally, a total of 29 overlapping genes (OGs) of compound targets and UC-related targets, namely HSD17B2, KDR, PLA2, ABCG2, MMP7, NOS2, MMP3, FADS1, CXCR1, AKR1B10, CDC25B, SELL, CD38, MAOA, EDNRA, LRRK2, CA2, ABCB1, STS, PDGFRB, SELP, PTGS2, PFKFB3, PIM2, MMP1, IGFBP5, EPHX2, MMP12 and MMP9, were regarded as potential targets of HTD for the treatment of UC (Figure 4C).

GO and Enrichment Analysis

As shown in Figure 5A and Table 4, the top terms (P<0.01) in the BPs, CCs, and MFs were selected. The enrichment of BPs

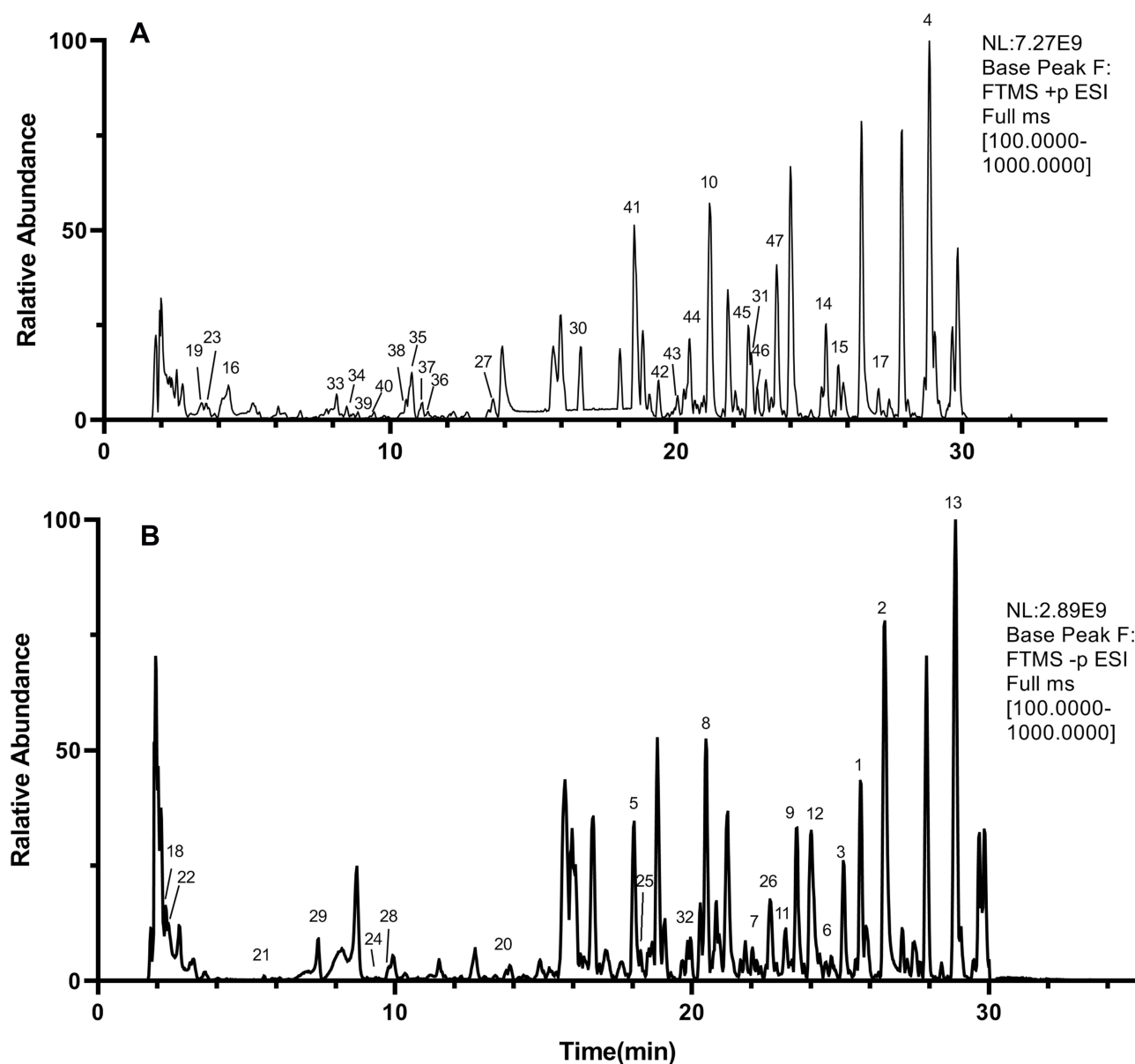


Figure 2 Chromatograph of the extracts of HTD by Q Exactive Orbitrap LC-MS/MS on positive-ion polarity mode (A) and negative-ion polarity mode (B).

involved the collagen catabolic process, collagen metabolic process, extracellular matrix disassembly, regulation of neuroinflammatory response, extracellular matrix organization, extracellular structure organization, external encapsulating structure organization, response to oxidative stress, reactive oxygen species metabolic process, and response to inorganic substance. Besides, further analysis of the network was performed by Cytoscape ClueGO plugin, and it was found that the collagen catabolic process, positive regulation of smooth muscle cell proliferation, long-chain fatty acid biosynthetic process, and positive regulation of phospholipase activity were mainly involved in BPs (Figure 5B). The enrichment

of CCs mainly involved secretory granule membrane, and other cell components. The enrichment of MFs involved the metalloendopeptidase activity and metallopeptidase activity.

The signal transduction pathways are shown in Table 5 and Figure 6 ($P < 0.01$). The result of pathway enrichment analysis of the HTD for UC revealed that the OGs were significantly associated with MMPs, imatinib, chronic myeloid leukemia, and photodynamic therapy-induced NF- κ B survival signaling, in WikiPathways, with IL-4 and IL-13 signaling and collagen degradation in Reactome Gene Sets, and also with MicroRNAs in cancer and IL-17 signaling pathway in KEGG.

Table 3 Compositions and Their Product Ions in Huangtu Decoction

No	Name	RT [min]	Formula	Ion Source Model	Observed MS ₁ (m/z)	Theoretical MS ₁ (m/z)	Error (ppm)	MS ₂ (m/z)	Source	Reference
1	LicoricesaponinG2	25.97	C ₄₂ H ₆₂ O ₁₇	[M-H] ⁻	837.39465	837.39142	3.23	661.36273,643.34851,485.32883,351.05743	E	[25,26]
2	Glycyrrhizic Acid	26.63	C ₄₂ H ₆₂ O ₁₆	[M-H] ⁻	821.39917	821.39651	2.66	645.36481,403.87167,351.05740	E	[25]
3	Liquiritigenin	25.68	C ₁₅ H ₁₂ O ₄	[M-H] ⁻	255.06674	255.06628	0.46	251.82465,153.01883,135.00815	E	[25]
4	Wogonin	28.86	C ₁₆ H ₁₂ O ₅	[M+H] ⁺	285.07596	285.07575	0.21	270.05237,269.04456,250.27339,242.05725	B/E	[27-30]
5	Eriodictyol	18.15	C ₁₅ H ₁₂ O ₆	[M-H] ⁻	287.05682	287.05611	0.71	269.04544,161.02400,151.00317,125.02377	E	[27]
6	Luteolin	24.70	C ₁₅ H ₁₀ O ₆	[M-H] ⁻	285.04126	285.04046	0.80	267.03111,241.05080,213.05574,199.04033,151.00322	E	[27]
7	Quercetin	22.46	C ₁₅ H ₁₀ O ₇	[M-H] ⁻	301.03635	301.03538	0.97	273.04129,257.04614,229.05078,178.99835	E	[25-27,31]
8	Genistein	20.21	C ₁₅ H ₁₀ O ₅	[M-H] ⁻	269.04630	269.04555	0.75	225.05536,201.05576,153.01889,133.02890	B/E	[27-29]
9	Pinobanksin	23.41	C ₁₅ H ₁₂ O ₅	[M+H] ⁺	271.06195	271.06120	0.75	253.05077,197.06038,153.01884,135.04456	B/E	[27-29]
10	Hesperetin	21.65	C ₁₆ H ₁₂ O ₆	[M+H] ⁺	301.07111	301.07066	0.45	286.04730,253.05286,167.03400,153.01833	B/E	[27]
11	Isorhamnetin	23.27	C ₁₆ H ₁₂ O ₇	[M-H] ⁻	315.05194	315.05103	0.91	300.02805,283.02634,255.02972	E	[27]
12	Chrysin	24.44	C ₁₅ H ₁₀ O ₄	[M-H] ⁻	253.05135	253.05063	0.72	232.98445,209.06047,197.05319,153.01881	B/E	[27-29]
13	Pinocembrin	28.78	C ₁₅ H ₁₂ O ₄	[M-H] ⁻	255.06676	255.06628	0.48	213.05573,211.0762,171.04478,151.00317	B/E	[27-29]
14	Calycosin	24.72	C ₁₆ H ₁₂ O ₅	[M+H] ⁺	285.07611	285.07575	0.36	270.05237,253.04919,242.05699,170.02107	B/E	[27-29]
15	Retrochalcone	25.84	C ₁₆ H ₁₄ O ₄	[M+H] ⁺	271.09680	271.09649	0.31	229.08594,170.02122,167.03412	E	[27]
16	5-Hydroxymethyl-2-Furaldehyde	4.34	C ₆ H ₆ O ₃	[M+H] ⁺	127.03929	127.03897	0.32	109.02880,99.04453,81.0341	A/D	[32-34]
17	Atractylenolide III	27.4	C ₁₅ H ₂₀ O ₃	[M+H] ⁺	249.14871	249.14852	0.19	231.13779,213.12749,175.07549	D	[32,33]
18	Uridine	2.62	C ₉ H ₁₂ N ₂ O ₆	[M+H] ⁺	243.06285	243.06226	0.59	188.05617,170.04547,144.06602,128.03465	A	[34]
19	Adenosine	3.56	C ₁₀ H ₁₃ N ₅ O ₄	[M+H] ⁺	268.10428	268.10403	0.25	170.26382,136.06189	A	[34]
20	Phlinoside A	13.77	C ₃₅ H ₄₆ O ₂₀	[M-H] ⁻	785.25299	785.25097	2.02	623.22131,477.16083,431.97238	A	[34]
21	Rehmannioside D	5.59	C ₂₇ H ₄₀ O ₂₀	[M-H] ⁻	685.22125	685.21910	-2.15	505.15607,341.10971,263.07773,179.05580	A	[34]
22	Citric Acid	2.7	C ₆ H ₈ O ₇	[M-H] ⁻	191.01965	191.01973	-0.08	173.00873,129.01881,111.0080	A	[34]
23	Guanosine	3.87	C ₁₀ H ₁₃ N ₅ O ₅	[M-H] ⁺	284.09921	284.09895	0.27	152.05684	A	[34]

(Continued)

Table 3 (Continued).

No	Name	RT [min]	Formula	Ion Source Model	Observed MS ₁ (m/z)	Theoretical MS ₁ (m/z)	Error (ppm)	MS ₂ (m/z)	Source	Reference
24	Mussaenosidic Acid	9.34	C ₁₆ H ₂₄ O ₁₀	[M-H] ⁻	375.13083	375.12967	1.16	331.13101, 252.75468, 213.07686, 169.08653	A	[34]
25	Verbascoside	18.27	C ₂₉ H ₃₆ O ₁₅	[M-H] ⁻	623.20050	623.19814	2.36	461.16702, 415.10403, 295.06149, 161.02397	A	[34]
26	Martynoside	22.62	C ₃₁ H ₄₀ O ₁₅	[M-H] ⁻	651.23041	651.22944	0.97	475.18237, 193.05034, 175.03967	A	[34]
27	p-Coumaric Acid	13.87	C ₉ H ₈ O ₃	[M+H] ⁺	163.03970	163.04007	-0.37	119.0495, 93.03382, 91.05441	B	[28]
28	Vanillic Acid	9.56	C ₈ H ₈ O ₄	[M-H] ⁻	167.03477	167.03498	-0.21	152.0110, 139.0031, 124.01594, 123.04449, 108.02094	B	[28]
29	4-Hydroxybenzoic Acid	7.77	C ₇ H ₆ O ₃	[M-H] ⁻	137.02400	137.02442	-0.42	136.01599, 109.02878, 108.89879, 93.03375	B	[28,35]
30	Scutellarin	17.12	C ₂₁ H ₁₈ O ₁₂	[M+H] ⁺	463.08777	463.08710	0.67	316.05765, 287.05515, 269.04486	B	[28-30,35]
31	Apigenin-7-O-Glucuronide	22.63	C ₂₁ H ₁₈ O ₁₁	[M+H] ⁺	447.09201	447.09219	-0.18	293.49832, 271.06015	B	[28,29,35,36]
32	2,5,6,7-Tetrahydroxyflavanone	19.81	C ₁₅ H ₁₂ O ₆	[M-H] ⁻	287.05692	287.05611	0.81	201.05571, 161.02400, 133.02881, 125.02381	B	[28,30,35]
33	Karakoline	8.02	C ₂₂ H ₃₅ NO ₄	[M+H] ⁺	378.26443	378.26389	0.54	360.25357, 332.22223, 328.22644, 310.21783	C	[37]
34	Isotalatizidine	8.51	C ₂₃ H ₃₇ NO ₅	[M+H] ⁺	408.27481	408.27445	0.36	390.26410, 376.25168, 372.25351, 358.23663	C	[38]
35	Fuziline	10.64	C ₂₄ H ₃₉ NO ₇	[M+H] ⁺	454.28012	454.27993	0.19	436.26950, 422.25400, 404.24393, 386.23343	C	[37-40]
36	Talatizamine	11.15	C ₂₄ H ₃₉ NO ₅	[M+H] ⁺	422.29053	422.29010	0.43	404.27063, 390.26404, 372.25360, 358.23761	C	[37-39]
37	Neoline	11.11	C ₂₄ H ₃₉ NO ₆	[M+H] ⁺	438.28530	438.28501	0.29	420.27451, 388.24826, 370.23965, 356.22174	C	[37-40]
38	Songorine	10.48	C ₂₂ H ₃₁ NO ₃	[M+H] ⁺	358.23807	358.23767	0.40	340.22723, 322.21643, 315.74133	C	[37,38]
39	Aconine	9.26	C ₂₅ H ₄₁ NO ₉	[M+H] ⁺	500.28632	500.28541	0.91	482.27646, 468.25943, 450.24875, 436.23288	C	[37,38]
40	Hetisine	11.15	C ₂₀ H ₂₇ NO ₃	[M+H] ⁺	330.20676	330.20637	0.39	312.19818, 294.18567, 258.60516	C	[37,38]
41	Benzoylmesaconine	18.55	C ₃₁ H ₄₃ NO ₁₀	[M+H] ⁺	590.29550	590.29597	-0.47	572.28363, 558.27057, 540.26007, 526.24512	C	[37-40]
42	l4-Benzoylaconine	19.39	C ₃₂ H ₄₅ NO ₁₀	[M+H] ⁺	604.31100	604.31162	-0.62	572.28558, 554.27521, 540.26080, 522.24957	C	[38-40]
43	Benzoylhyapaconine	20.05	C ₃₁ H ₄₃ NO ₉	[M+H] ⁺	574.30130	574.30106	0.24	542.27521, 510.24875, 482.21729	C	[40]
44	Mesaconitine	21.32	C ₃₃ H ₄₅ NO ₁₁	[M+H] ⁺	632.30682	632.30654	0.28	600.28052, 572.28595, 540.25922, 512.26617	C	[37,38,40]
45	Hypaconitine	22.53	C ₃₃ H ₄₅ NO ₁₀	[M+H] ⁺	616.31146	616.31162	-0.16	584.28546, 556.29077, 524.26483, 496.27551, 492.23889	C	[37,38,40]
46	Aconitine	22.84	C ₃₄ H ₄₇ NO ₁₁	[M+H] ⁺	646.32230	646.32219	0.11	586.30212, 554.27502, 526.28052	C	[37,38,40]
47	Deoxyaconitine	23.34	C ₃₄ H ₄₇ NO ₁₀	[M+H] ⁺	630.32780	630.32727	0.53	598.29962, 570.30670, 538.28058, 510.28748	C	[38,40]

Notes: A: *Rehmanniae Radix*; B: *Scutellariae Radix*; C: *Aconiti Lateralis Radix Praeparata*; D: *Atracylodes Macrocephalae Rhizoma*; E: *Glycyrrhizae radix et rhizoma*.

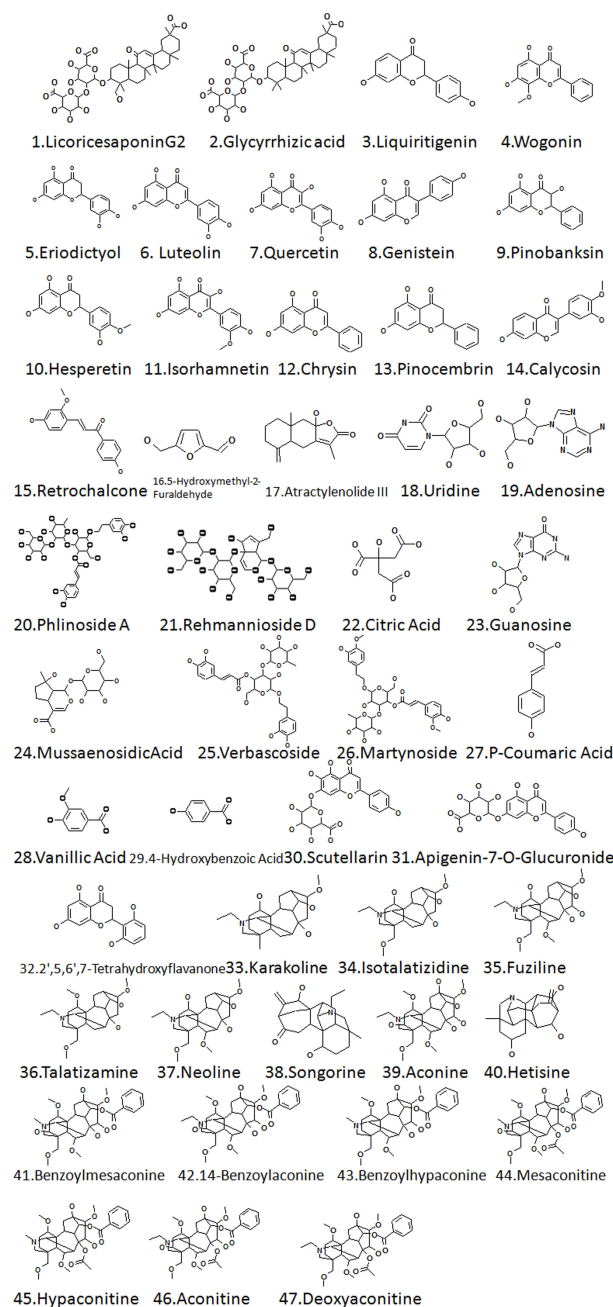


Figure 3 Compounds detected in the extracts of HTD by Q Exactive Orbitrap LC-MS/MS.

Compound-Target- Pathway Network

As shown in Figure 7, the compound-target-pathway network included 46 nodes (11 compounds, 18 targets, and 17 pathways) and 158 edges. According to the analysis of the network, each active compound acted on at least 2 target genes, each gene was regulated by at least 2 active compounds, and at least 18 genes were potentially involved in each pathway related to UC, indicating the action characteristics of multiple active compounds and targets of HTD in the treatment of UC.

Experimental Validation

Effect of HTD on Histopathological Changes

As shown in Figure 8A and B, the mice in model group and DSS+HTD group manifested varying degrees of loose stool and weight loss compared with those in control group. The pathological conditions of colon tissues of the mice treated with DSS+HTD were improved compared with those in model group. The histopathological changes in each group are shown in Figure 8C. The results of H&E staining showed that DSS induced massive inflammatory infiltration, mucosal edema, and hyperplasia in colonic tissues of model group, but less necrosis was observed in DSS+HTD group.

Effect of HTD on Serum Levels of IL-1 β and IL-6

As shown in Figure 8F and G, the serum levels of IL-1 β and IL-6 were significantly increased in model group compared with those in control group ($P < 0.01$). Compared to the model group, serum IL-1 β and IL-6 levels decreased greatly in DSS+HTD group ($P < 0.01$).

Effect of HTD on MMP Expressions

The expression levels of MMP1, MMP3, MMP7, MMP9 and MMP12 in colon tissues were investigated via immunohistochemical staining (Figure 9) and WB (Figure 10A and B). In contrast with model group, the expressions of MMP1, MMP3 and MMP9 in colon tissues in DSS+HTD group were significantly lower, and the expressions of MMP7 and MMP12 declined with no significant difference. In addition, qRT-PCR was applied to explore the role of HTD in regulating the expression of MMPs in colon tissues. As shown in Figure 10C, DSS significantly up-regulated the expressions of MMP1, MMP3, MMP7, MMP9 and MMP12 in colon tissues, but after HTD treatment, the upregulation of MMP1, MMP3 and MMP9 was reversed, while MMP7 and MMP12 had no significant changes.

Discussion

UC is a chronic disease with an increasing incidence worldwide over recent decade, especially in developing nations.² HTD is a very effective prescription for the clinical treatment of hematochezia in TCM, and has been reported as non-surgical treatment of UC with minimal side effects. However, there is a lack of modern animal experimental evidence and no literature report on its active compounds and mechanism. In this study, the UC model was established to evaluate the therapeutic effects of HTD on UC, and the possible mechanism of this TCM formula

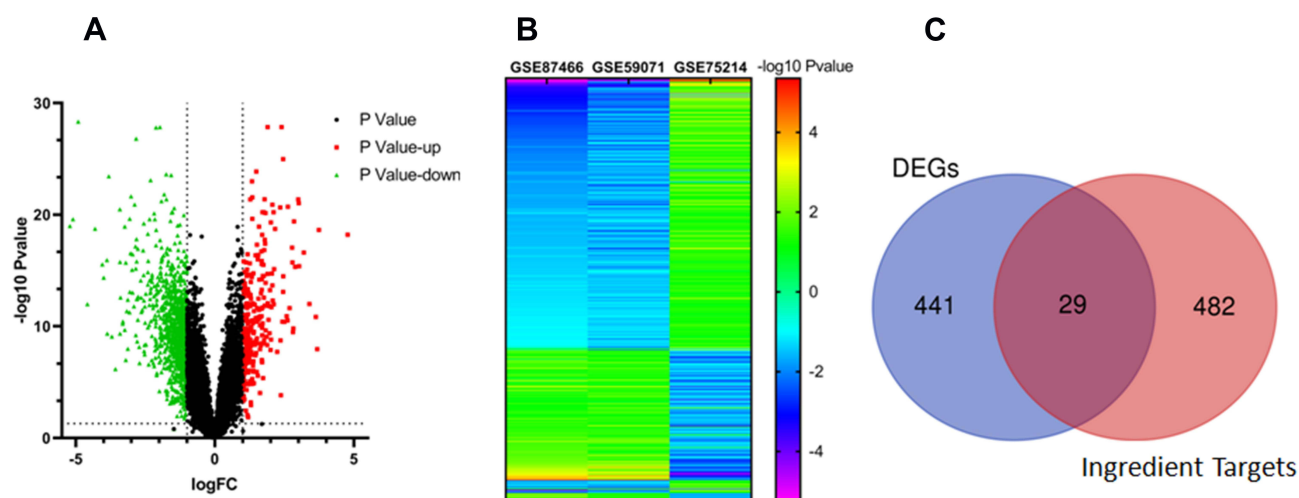


Figure 4 The overlapped targets screening strategy. **(A)** The volcano plot of the differentially expressed genes (DEGs) in datasets of GSE87466, GSE59071, and GSE75214. The red nodes represent up-regulated DEGs, while green nodes represent down-regulated DEGs; the horizontal dashed lines show the P value is less than 0.05 and the vertical dashed lines indicate the value of \log_2 fold change (FC) is more than 1. **(B)** Heatmap of DEGs. The vertical axis represents the genes. The horizontal axis shows the change of expression levels of DEGs in datasets of GSE87466, GSE59071, and GSE75214 by $-\log_{10}$ Pvalue. **(C)** Overlap genes of DEGs and predicted compound targets.

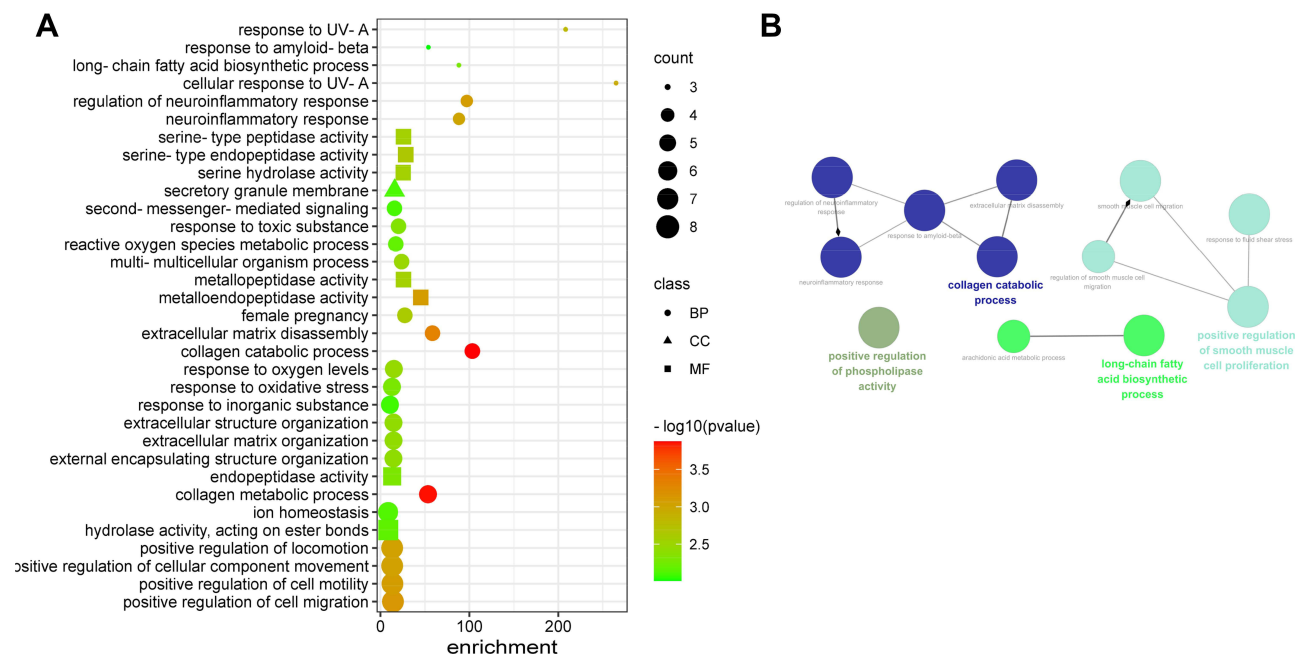


Figure 5 Bubble diagram of GO functional enrichment analysis of overlapped genes **(A)** The bubble chart. **(B)** The visualization analysis of BPs.

for treating UC was also explored based on network pharmacology.

HTD contains 7 medicinal herbs, including *Rehmanniae Radix*, *Scutellariae Radix*, *Aconiti Lateralis Radix Praeparata*, *Atractylodis Macrocephalae Rhizoma* and *Glycyrrhizae Radix et Rhizoma*, *Terra Flava Usta* and *Asini Corii Colla*. In the present study, HPLC-MS/MS was used to analyze the chemical compositions of HTD, and a total of 47

compounds were identified from it, mostly flavones and alkaloids. Furthermore, network pharmacology was used to explore its possible mechanism. The compound-target-pathway network showed that wogonin with the highest degree was regarded as the most effective compound, followed by LicoricesaponinG2, Liquiritigenin, Retrochalcone, Genistein, Atractylenolide III, Glycyrrhizic acid, Pinobanksin, Pinocembrin, Mussaenosidic acid and Hesperetin. These

Table 4 GO Functional Enrichment Analysis Results of Overlapping Genes (OGs)

GO	ID	Description	Gene ID	Count
BP	GO:0030574	Collagen catabolic process	MMP1/MMP3/MMP7/MMP9/MMP12	5
BP	GO:0032963	Collagen metabolic process	MMP1/MMP3/MMP7/MMP9/MMP12/PDGFRB	6
BP	GO:0022617	Extracellular matrix disassembly	MMP1/MMP3/MMP7/MMP9/MMP12	5
BP	GO:0150077	Regulation of neuroinflammatory response	MMP3/MMP9/PTGS2/LRRK2	4
BP	GO:0150076	Neuroinflammatory response	MMP3/MMP9/PTGS2/LRRK2	4
BP	GO:0071492	Cellular response to UV-A	MMP1/MMP3/MMP9	3
BP	GO:0070141	Response to UV-A	MMP1/MMP3/MMP9	3
BP	GO:0030198	Extracellular matrix organization	KDR/MMP1/MMP3/MMP7/MMP9/MMP12	6
BP	GO:0043062	Extracellular structure organization	KDR/MMP1/MMP3/MMP7/MMP9/MMP12	6
BP	GO:0045229	External encapsulating structure organization	KDR/MMP1/MMP3/MMP7/MMP9/MMP12	6
BP	GO:0006979	Response to oxidative stress	CD38/MMP3/MMP9/PDGFRB/PTGS2/LRRK2	6
BP	GO:0072593	Reactive oxygen species metabolic process	MMP3/NOS2/PDGFRB/PTGS2/LRRK2	5
BP	GO:0010035	Response to inorganic substance	CA2/KDR/MMP3/MMP9/PDGFRB/LRRK2	6
BP	GO:1904645	Response to amyloid-beta	MMP3/MMP9/MMP12	3
BP	GO:0030335	Positive regulation of cell migration	IGFBP5/KDR/MMP7/MMP9/PDGFRB/PLAU/PTGS2/SELP	8
BP	GO:2000147	Positive regulation of cell motility	IGFBP5/KDR/MMP7/MMP9/PDGFRB/PLAU/PTGS2/SELP	8
BP	GO:0051272	Positive regulation of cellular component movement	IGFBP5/KDR/MMP7/MMP9/PDGFRB/PLAU/PTGS2/SELP	8
BP	GO:0040017	Positive regulation of locomotion	IGFBP5/KDR/MMP7/MMP9/PDGFRB/PLAU/PTGS2/SELP	8
BP	GO:0007565	Female pregnancy	STS/CD38/IGFBP5/MMP7/MMP9	5
BP	GO:0044706	Multi-multicellular organism process	STS/CD38/IGFBP5/MMP7/MMP9	5
BP	GO:0070482	Response to oxygen levels	CD38/EDNRA/NOS2/PDGFRB/PLAU/PTGS2	6
BP	GO:0009636	Response to toxic substance	EPHX2/PDGFRB/PTGS2/ABCG2/AKR1B10	5
BP	GO:0042759	Long-chain fatty acid biosynthetic process	EPHX2/FADS1/PTGS2	3
BP	GO:0050801	Ion homeostasis	CA2/CD38/EDNRA/CXCR1/KDR/ABCG2/LRRK2	7
BP	GO:0019932	Second-messenger-mediated signaling	CXCR1/KDR/NOS2/SELP/LRRK2	5
CC	GO:0030667	Secretory granule membrane	CD38/CXCR1/PLAU/SELL/SELP	5
MF	GO:0004222	Metalloendopeptidase activity	MMP1/MMP3/MMP7/MMP9/MMP12	5
MF	GO:0004252	Serine-type endopeptidase activity	MMP1/MMP3/MMP7/MMP9/PLAU	5
MF	GO:0008236	Serine-type peptidase activity	MMP1/MMP3/MMP7/MMP9/PLAU	5
MF	GO:0008237	Metallopeptidase activity	MMP1/MMP3/MMP7/MMP9/MMP12	5
MF	GO:0017171	Serine hydrolase activity	MMP1/MMP3/MMP7/MMP9/PLAU	5
MF	GO:0004175	Endopeptidase activity	MMP1/MMP3/MMP7/MMP9/MMP12/PLAU	6
MF	GO:0016788	Hydrolase activity, acting on ester bonds	STS/CA2/CDC25B/EDNRA/EPHX2/PFKFB3/LRRK2	7

Table 5 Pathway Enrichment Analysis Results of Overlapping Genes (OGs)

Class	Description	Gene ID	Count
WikiPathways	Matrix Metalloproteinases	MMP1/MMP3/MMP7/MMP9/MMP12	5
WikiPathways	Imatinib and Chronic Myeloid Leukemia	PDGFRB/ABCB1/ABCG2/PIM2	4
WikiPathways	Photodynamic therapy-induced NF-kB survival signaling	MMP1/MMP3/MMP9/PTGS2	4
WikiPathways	Spinal Cord Injury	MMP9/MMP12/NOS2/PTGS2/SELP	5
WikiPathways	Hepatitis C and Hepatocellular Carcinoma	CXCR1/MMP1/NOS2/PTGS2	4
WikiPathways	IL-18 signaling pathway	MMP1/MMP3/MMP9/NOS2/PTGS2	5
Reactome Gene Sets	Interleukin-4 and Interleukin-13 signaling	MAOA/MMP1/MMP3/MMP9/NOS2/PTGS2	6
Reactome Gene Sets	Collagen degradation	MMP1/MMP3/MMP7/MMP9/MMP12	5
Reactome Gene Sets	Activation of Matrix Metalloproteinases	MMP1/MMP3/MMP7/MMP9	4
Reactome Gene Sets	Degradation of the extracellular matrix	MMP1/MMP3/MMP7/MMP9/MMP12	5
Reactome Gene Sets	Extracellular matrix organization	KDR/MMP1/MMP3/MMP7/MMP9/MMP12	6
Reactome Gene Sets	Signaling by Interleukins	MAOA/MMP1/MMP3/MMP9/NOS2/PTGS2	6
KEGG Pathway	MicroRNAs in cancer	CDC25B/MMP9/PDGFRB/ABCB1/PLAU/PTGS2	6
KEGG Pathway	MicroRNAs in cancer	CDC25B/MMP9/PDGFRB/ABCB1/PLAU/PTGS2	6
KEGG Pathway	IL-17 signaling pathway	MMP1/MMP3/MMP9/PTGS2	4
KEGG Pathway	Pathways in cancer	EDNRA/MMP1/MMP9/NOS2/PDGFRB/PTGS2	6
KEGG Pathway	IL-17 signaling pathway	MMP1/MMP3/MMP9/PTGS2	4

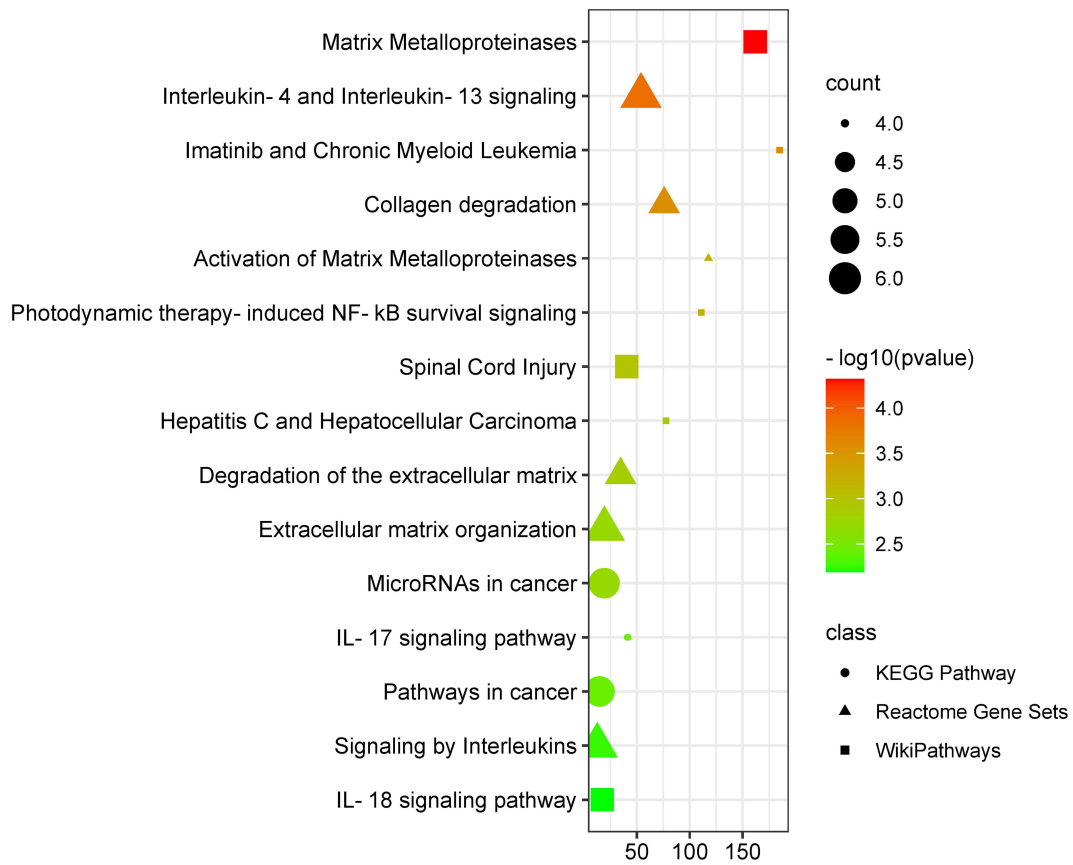


Figure 6 Enriched pathway analysis of potential targets of HTD compounds against UC.

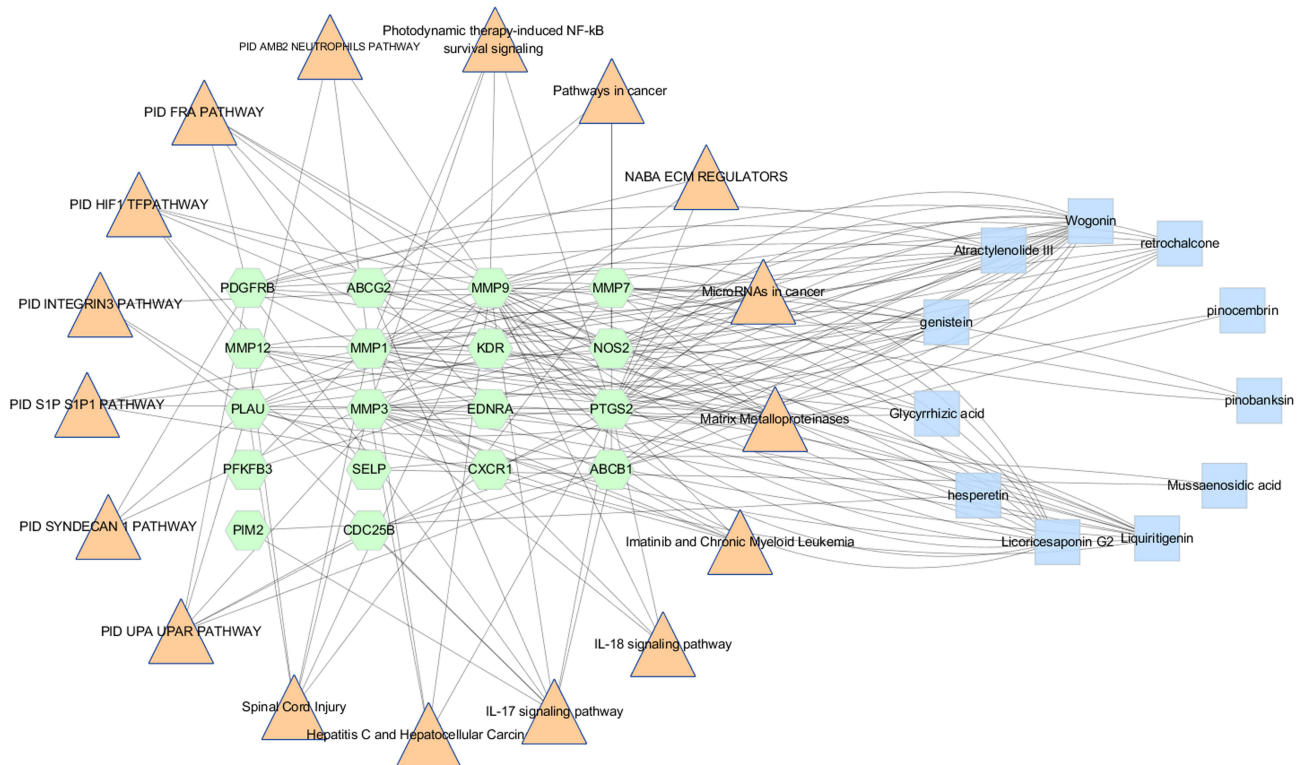


Figure 7 Compound-Target-Pathway Network, the pink triangle node are the signal transduction pathways; green hexagon nodes are OGs, blue rectangle nodes are the compounds of HTD extract, lines represent the interactions between them.

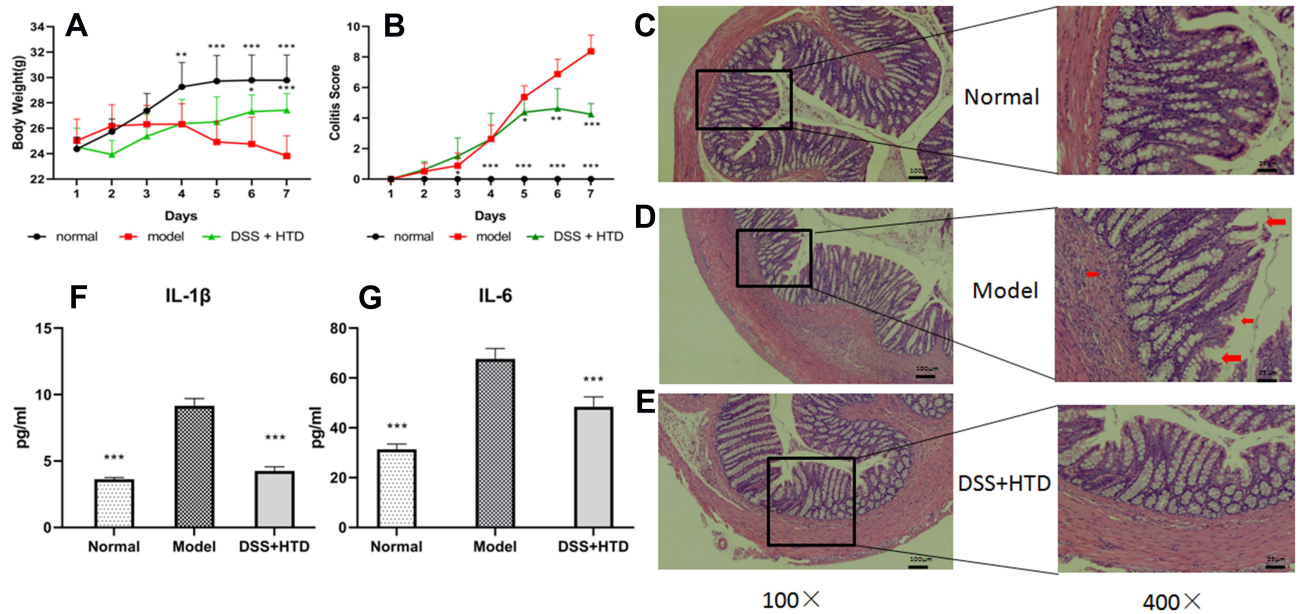


Figure 8 HTD ameliorates DSS-induced colitis. (A) Curve of daily Colitis scores. (B) Curve of daily body weights. (C) Colonic mucosa of normal group mice. (D) Colon of a DSS-treated mouse, severe inflammatory infiltration, edema, loss of crypts and ulcerations are seen. (E) HTD decreased DSS-induced colonic inflammation. (F and G) Effect of HTD on serum level of IL-1β and IL-6. Data are expressed as mean±SD (n = 8). *P < 0.05, **P < 0.01 and ***P < 0.001 versus model group.

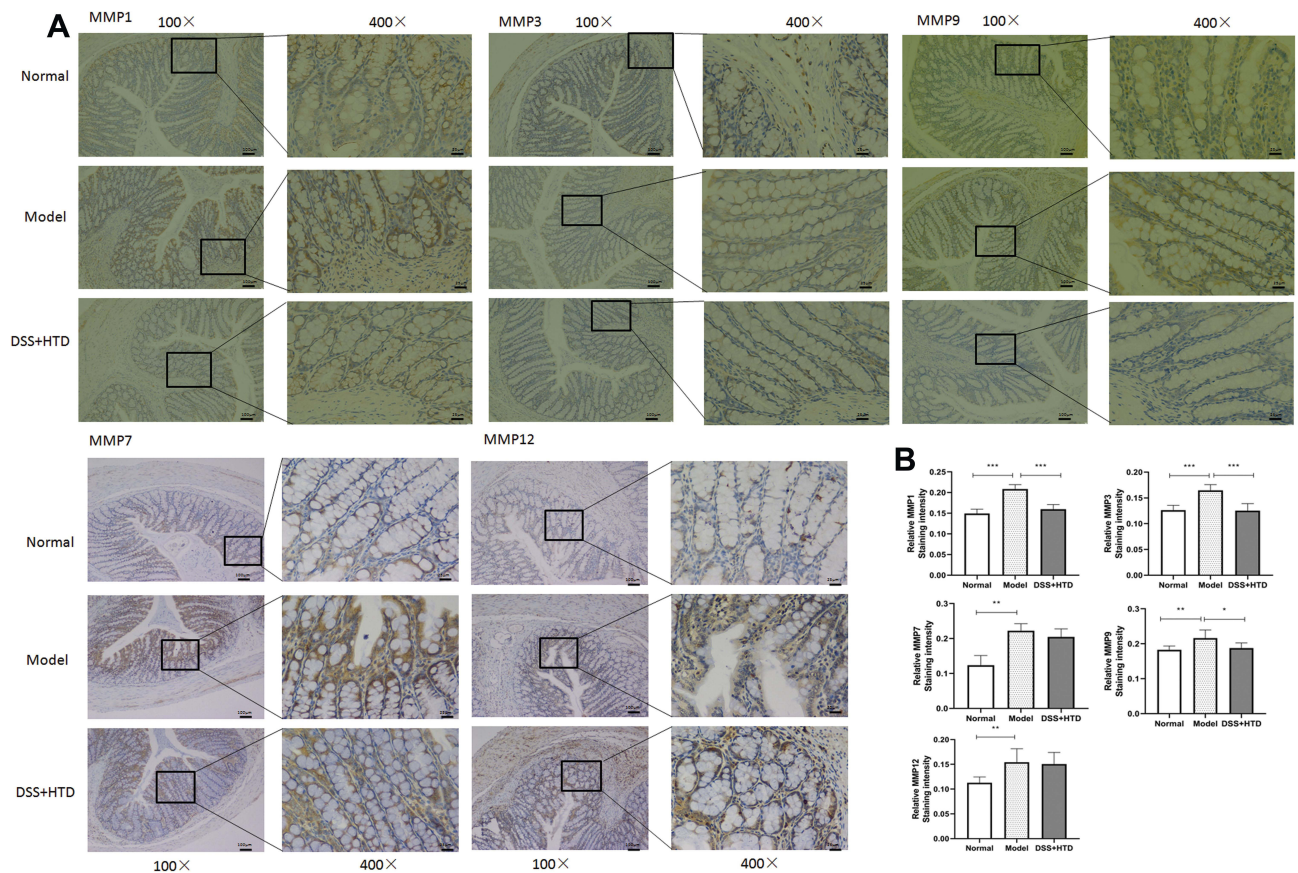


Figure 9 Representative immunohistochemical staining with MMP1, MMP3, MMP7, MMP9 and MMP12 in colon tissues in mice treated with DSS. (A) Immunohistochemical staining for MMP1, MMP3, MMP7, MMP9 and MMP12 in normal, model and DSS+HTD group. (B) Statistical analysis of relative immunohistochemical staining intensity for MMP1, MMP3, MMP7, MMP9 and MMP12 in normal, model and DSS+HTD group. *P < 0.05, **P < 0.01 and ***P < 0.001 versus model group.

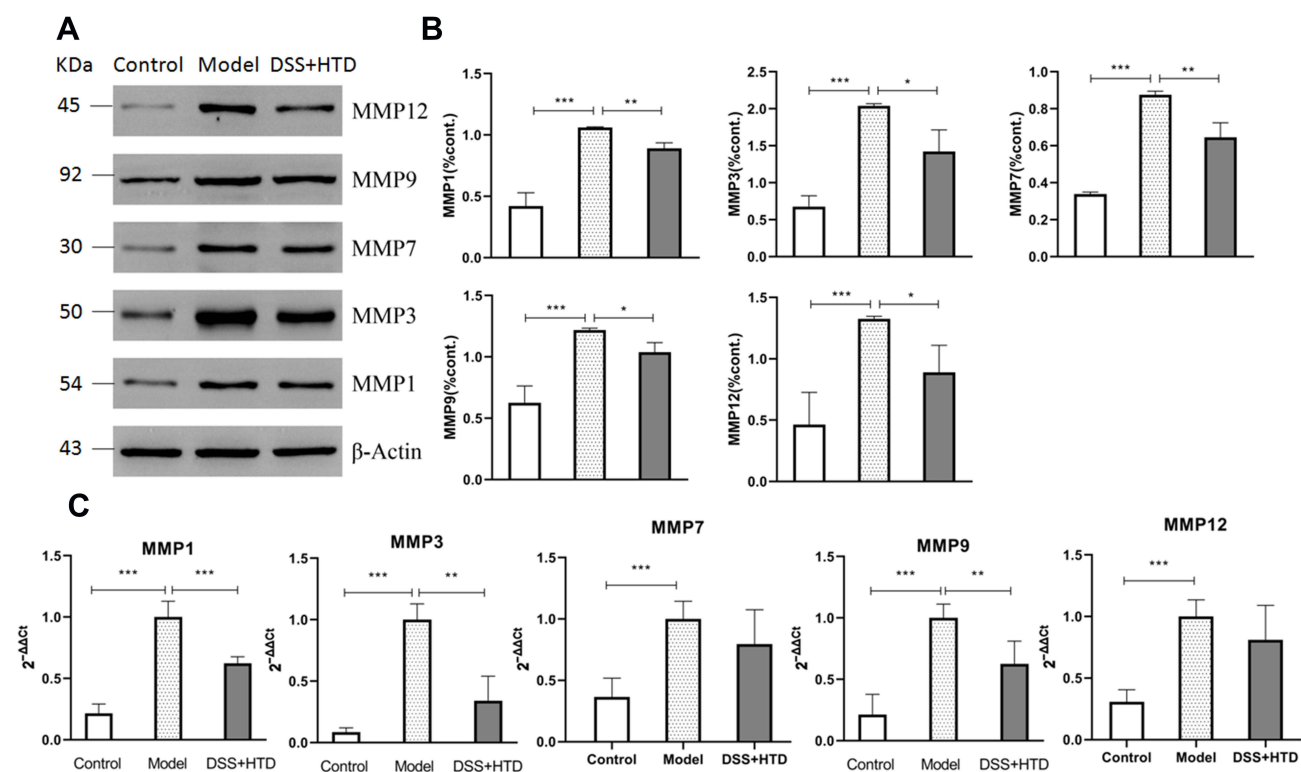


Figure 10 Effects of HTD on the expression of MMPs. **(A)** Western blot of MMP1, MMP3, MMP7, MMP9 and MMP12 in colon tissues. **(B)** Quantification of MMP1, MMP3, MMP7, MMP9 and MMP12. **(C)** Decreased protein expressions of MMP1, MMP3, MMP7, MMP9 and MMP12 in the colon obtained from mice treated with DSS. * $P < 0.05$, ** $P < 0.01$ and *** $P < 0.001$ versus model group.

compounds identified in this formula may provide fundamentals for establishing Q-markers for further reliable quality control of HTD. In addition, the possible molecular mechanism and signaling pathway for HTD were also predicted based on GO and KEGG enrichment analysis. The results revealed that the enrichment of BPs of HTD involved collagen catabolic process, collagen metabolic process, and extracellular matrix disassembly. The pathway enrichment analysis results showed that the pathways covered MMPs, MicroRNAs in cancer, IL-17 signaling pathway and so on. Therefore, the prediction results were further verified by in vivo animal experiments, and a similar result to network pharmacology was obtained. HTD was effective in the treatment of UC through decreasing the protein and mRNA expressions of MMP1, MMP3 and MMP9. MMPs are a family of proteolytic enzymes characterized by the ability to degrade extracellular matrix (ECM) proteins such as collagen, proteoglycan, fibronectin, elastin, and laminin.^{41,42} Chronic inflammation and aberrant tissue remodeling with excessive accumulation or degradation of ECM components are markers in UC. Hence, MMPs are thought to be predominant proteases involved in the

pathogenesis of UC.^{41–43} MMPs have been recognized as noninvasive diagnostic tools for UC in many studies.^{41,42} Specific synthetic MMP inhibitors and some natural products or therapeutic interventions have been described to ameliorate intestinal inflammation.⁴³

Conclusion

In conclusion, HTD can alleviate the colonic inflammation via inhibiting MMPs including MMP1, MMP3 and MMP9. The present study will benefit the further development of this classic TCM formula.

Acknowledgments

The authors wish to thank the editor and the anonymous reviewers whose constructive comments are very helpful in strengthening the presentation of this paper. The research was supported by Chengdu High-level Key Clinical Specialty Construction Project.

Disclosure

The authors report no conflicts of interest in this work.

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