Assessment of pulmonary infectious disease treatment with Mongolian medicine formulae based on data mining, network pharmacology and molecular docking

Baochang Zhou a, Zhanhong Qian a, Qinyu Li b, Yuan Gao c,*, Minhui Li a,b,d,e,⇑

a College of Traditional Chinese Medicine, Inner Mongolia Medical University, Hohhot 010110, China
b Department of Pharmacy, Baotou Medical College, Baotou 014040, China
c Inner Mongolia Autonomous Region Hospital of Traditional Chinese Medicine (Inner Mongolia Medical University of Clinical College of Traditional Chinese Medicine), Hohhot 010020, China
d Inner Mongolia Institute of Traditional Chinese and Mongolian Medicine, Hohhot 010010, China
e Inner Mongolia Key Laboratory of Characteristic Geoherbs Resources Protection and Utilization, Baotou Medical College, Baotou 014040, China

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Objective: Pulmonary infectious diseases (PID) include viral pneumonia (VP) and pulmonary tuberculosis (PT). Mongolian medicine (MM) is an effective treatment option in China, however, the core group medicines (CGMs) in the treatment of PID and their underlying therapeutic mechanisms remain unclear. In this study, through the method of data mining, the CGMs of MM for the treatment of PID were excavated, and the possible mechanism of action of the CGMs in the treatment of PID was explored by using network pharmacology.

Methods: First, 89 MM formulae for the treatment of pulmonary infectious diseases collected from Gan Lu Si Bu, Meng Yi Jin Kui, People’s Republic of China Ministry of Health Drug Standards (Mongolian Medicine Volume), Standard of Mongolian Medicine Preparations in Inner Mongolia (2007 Edition), and Standard of Mongolian Medicine Preparations in Inner Mongolia (2014 Edition). The CGMs of MM for PID were excavated through association rule analysis and cluster analysis. Then, the active ingredients and potential targets of the CGMs were obtained from TCMSP, TCMIP, BATMAN-TCM databases. PID targets information was collected from OMIM, GeneCards, and DrugBank databases. The possible targets of CGMs treatment for PID were obtained by intersection. The PPI network was constructed through the STRING database, and the topology analysis of the network was performed. Through the enrichment analysis of the intersection targets by R language, the main action pathways and related target proteins of CGMs in the treatment of PID were screened out. The results were verified by molecular docking.

Results: A total of 89 formulae were included, involving 164 MM herbs. The efficacy of the drugs was mainly cough-suppressing and panting-calming herbs, and heat-clearing herbs. The nature and flavor were mainly bitter and cold. The CGMs of MM to treatment of PID was excavated as the classic famous formula Sanzi Decoction (Toosendan Fructus-Chebulae Fructus-Gardeniae Fructus). A total of 28 candidate components and 237 predicted targets of CGMs were collected, and 61 common targets with PID were obtained, including key compounds such as quercetin, kaempferol, β-sitosterol and stigmasterol and key targets such as VEGFA, IL6, TP53, AKT1. KEGG enrichment analysis yielded AGE-RAGE signaling pathways, IL-17 signaling pathways, and TNF signaling pathways. Molecular docking results showed that the key targets were well matched with the potential active ingredients of CGMs.

Conclusion: This study found that MM commonly used cough-suppressing and panting-calming herbs in combination with heat-clearing herbs to treat PID, and the CGMs for the treatment of PID is “Toosendan Fructus-Chebulae Fructus-Gardeniae Fructus”. CGMs mainly play a role in the treatment of PID by acting on VEGFA, IL6, TP53, AKT1 and other targets, regulating AGE-RAGE signaling pathways, IL-17 signaling pathways, and TNF signaling pathways.

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1. Introduction

Pulmonary infectious diseases (PID) include viral pneumonia (VP) and pulmonary tuberculosis (PT). According to estimates by the World Health Organization (WHO), approximately 450 million patients are diagnosed with pneumonia worldwide each year, of whom approximately 4 million (7% of pneumonia deaths) die from VP (Mathers et al., 2003). Viral infections account for 25%–50% of non-bacterial pneumonia and are its main cause of death (Kao et al., 2017; Burk et al., 2016). Primary symptoms of VP include fever, cough with or without phlegm, shortness of breath, and pharyngalgia. Despite its significance, there is a lack of relevant research data on the pathogenesis of VP, which limits its clinical treatment. The U. S. Food and Drug Administration (FDA) has approved the use of antiviral neuraminidase inhibitors such as oseltamivir and peramivir (Dandachi & Rodriguez-Barradas, 2018) in the treatment of VP. However, antiviral drug efficacy is negatively affected by the high degree of respiratory virus variability, emergence of drug-resistant strains, and adverse drug reactions. According to the “Global Pulmonary Tuberculosis Report 2019” released by the WHO, tuberculosis is one of the top 10 causes of death globally and is responsible for the greatest number of deaths caused by a single infectious pathogen. Currently, antibiotics remain the primary treatment for patients with PT. However, with the widespread application of anti-tuberculosis drugs, the global incidence of drug-resistant PT is gradually increasing. As a result, the multidrug resistance rate is as high as 82%, which has made the treatment of PT significantly more difficult (Liu et al., 2021). Thus, the search for highly effective antiviral drugs and anti-tubercle bacilli drugs are gaining an increasing level of research interest.

The curtain rose on the application of traditional medicine millennia ago, and a >2800-year history of traditional Mongolian medicine (MM) has continued thereafter (Robesan, 1922). Throughout history, the use of MM has developed distinctive ethnic, regional, and clinical characteristics, and it is characterized by the incorporation of theories and advanced clinical experiences of various ethnic groups in Han, Tibet, and other regions in China (Zhang et al., 2015). MM has a long history of treating PID. MM believes that “VP” belongs to the category of “Pulmonary fever”, which is due to the fact that “Badagan” (Badagan is feminine, belongs to water, and its attribute is cold) is prevalent, and “Xila” (Xila is masculine, is a fire source, and its attribute is heat) fever is flourishing in the pulmonary. Treatment should focus on clearing heat; PT belongs to the category of “pulmonary carbuncle” is due to the fact that “Badagan” is too prosperous, and “Xi la” fever is blazing in the pulmonary, so this disease occurs. Therefore, “VP” and “PT” have a common cause. MM has distinctly affected the treatment of PID, and has had a therapeutic impact on the disease treatment via multiple targets and pathways deregulated in PID. However, the underlying mechanisms by which MM therapies affect PID treatment remain unclear, which restricts the internationalization and standardization of MM use in the treatment of PID.

Data mining technology has the potential to improve the assessment of the medication mode of the clinical upper agent, identify potential compatibility rules from existing medication methods, and provide references for subsequent clinical applications (Tang, Shen, & Yang, 2019; Wu et al., 2021; Wu et al., 2021). However, these studies do have limitations. Effects of MM occur via multicomponent, multitarget, and multipathway mechanisms, it makes MM uniquely effective in the treatment of complex diseases. However, it also brings out difficulty with regard to clarifying the therapeutic mechanisms of MM. The emergence of network pharmacology has helped clarify the therapeutic effects of MM (Cao et al., 2020). Network pharmacology is a new method for studying the biologically active components, potential drug targets, and molecular mechanisms of MM (Wan et al., 2021). In addition, it can be used to elucidate the pharmacological mechanisms of candidate drugs from an overall perspective, which is consistent with the overall concept of MM theory (Hopkins, 2011). Research has shown that network pharmacology is a new and feasible method for identifying drug candidates in the early stages of drug screening (Deb, 2019). In addition, molecular docking is based on the docking of small molecules to target proteins and is used in basic research studies (Lohning, Levonis, Williams, & Schweiker, 2017).

In this study, representative classes of MM therapies from the Qing Dynasty to the modern period (CE 1751–2014) were selected, which included therapies described in Gan Lu Si Bu (Qing dynasty, CE 1751–1785, One of the three classic works of Mongolian medicine), Meng Yi Jin Kui (Qing dynasty, CE 1829, Mongolian medical prescription monograph, one of the three classic works of Mongolian medicine), People's Republic of China Ministry of Health Drug Standards (Mongolian Medicine Volume), and The Standard of Mongolian Medicine Preparations In Inner Mongolia (Technical standards for the preparation, use, inspection and supervision and management of Mongolian medicinal preparations in medical institutions in Inner Mongolia). Gan Lu Si Bu is the one of the three classic works of Mongolian medicine, which laid a solid theoretical foundation for the development of MM. It was written in 1751–1785. Meng Yi Jin Kui also known as Fang Hai, is particularly important since it was the first systematic and complete book of prescriptions in the history of MM. The People's Republic of China Ministry of Health Drug Standards (Mongolian Medicine Volume) was the first, highly scientific Mongolian medicine standard, which was issued by the Ministry of Health in 1998. Moreover, the Standard of Mongolian Medicine Preparations In Inner Mongolia was the first book to include prescription specifications that was issued by Inner Mongolia, and included prescriptions that basically reflect the clinical use of MM therapeutics at the time. In our study, core group medicines (CGMs) in PID treatment were determined by mining PID prescription data from classic MM texts. Then, network pharmacology and molecular docking were used to explore CGMs therapeutic mechanisms in the treatment of PID to provide data needed for further research (Sun, Sun, Yan, Li, & Xin, 2020; Sun, Jiang, Wang, & Liu, 2020; Yu et al., 2021).

2. Materials and methods

2.1. Data source and normalization

In this study, 89 MM formulae for the treatment of PID and 164 MMMs were collected from Gan Lu Si Bu (Qing dynasty, CE 1751–1785), Meng Yi Jin Kui (Qing dynasty, CE 1829), People's Republic of China Ministry of Health Drug Standards (Mongolian Medicine Volume) (1998), Standard of Mongolian Medicine Preparations In Inner Mongolia (2007 Edition) (2007), and Standard of Mongolian Medicine Preparations In Inner Mongolia (2014 Edition) (2014) texts. Data were used to establish a database of MMss and perform subsequent analyses. Moreover, the name of each prescription used in MM was standardized based on conventions detailed in the Pharmacopeia of the People's Republic of China (2020 Edition) (Chinese Pharmacopoeia Commission, 2020) and Nei Meng Gu Meng Yao Cai Biao Zhun (1987 Edition) (Mongolian herbal medicine standards promulgated by Inner Mongolia Autonomous Region) (Inner Mongolia Department of Health, 1987).

2.2. Data processing and analysis

A database of MM forms for the treatment of PID was established using Excel 2013 and was converted into a format compatible with the data mining software. Based on MM medication rules,
categories, properties, and tastes associated with single drugs included in MM prescriptions, Studio software was used for implementing a hierarchical clustering algorithm to categorize high-frequency drugs. Further, the Apriori algorithm was used to assess associations between high-frequency drugs.

2.3. CGMs active compound collection

Compounds comprising all herb were identified using the following three traditional Chinese medicine databases: Traditional Chinese Medicine Systems Pharmacology Database and Analysis Platform (TCMSP) (Ru et al., 2014), Integrative Pharmacology-based Research Platform of TCM (TCMIP) (Xu et al., 2017), and Bioinformatics Analysis Tool for Molecular Mechanisms of Traditional Chinese Medicine Database (BATMAN-TCM) (Liu et al., 2016). Screening criteria for bioactive compounds were set based on the TCMSP database and included oral bioavailability (OB) ≥ 30% and drug-likeness (DL) ≥ 0.18. Putative targets of bioactive compounds were collected from the three previously mentioned TCM databases, and the gene name of each target protein was searched using the UniProt database. Conditions associated with data were reviewed; duplicate, non-human targets were removed; the corresponding gene abbreviation of targets were identified to facilitate follow-up research. A simultaneous herb-compound-target network was constructed to explore active compounds and their potential targets. Core compounds were identified using an herb-compound-target network.

2.4. Therapeutic targets of CGMs in treatment of PID

PID-related therapeutic targets were obtained from OMIM (Amberger & Hamosh, 2017), GeneCards (Safran et al., 2010), and DrugBank (Wishart et al., 2018) databases; the search terms used were “Viral Pneumonia” (including Corona Virus Disease 2019), and “Pulmonary Tuberculosis”, which was defined as the key word. Results were restricted to human genes and proteins. Targets with a relevance score above the median value were selected from the GeneCards database. CGMs and PID targets that overlapped were identified using a Venny 2.1. Overlapping CGMs and PID targets identified were considered the best potential targets of CGMs in the treatment of PID.

2.5. Network construction and analysis

First, using the Cytoscape software, an open-source software platform for visualizing complex networks, a “disease chemical composition target” network of CGMs in the treatment of VP and PT was constructed. Second, using the same software, a common mechanism network of CGMs in the treatment of VP and PT was constructed. Third, to obtain protein–protein interaction (PPI) data, candidate targets (common targets) were used as inputs in STRING with a minimum interaction score of >0.7, and the search was limited to “Homo sapiens” visual network graphs were created using Cytoscape. Target interaction network parameters were calculated using NetworkAnalyzer, and core genes of the network were predicted.

2.6. Protein functional enrichment analysis

To elucidate the therapeutic mechanisms of the CGMs on PID, we used the R language Bioconductor biology data package to perform gene GO function and KEGG pathway enrichment analysis on common targets in the PPI network. The three functional categories cell component, molecular function, and biological process were separately assessed to identify target protein functions. According to the statistical method, set P to 0.05 and q to 0.05 to perform GO (gene ontology) biological enrichment analysis and KEGG pathway enrichment analysis, which were visualized using the “ggplot2” package in R.

2.7. Molecular docking

Molecular docking was conducted to validate whether CGMs were capable of binding targets. Two-dimensional structures of the top 10 core compounds (top 10 degrees in the herb-compound-target network) were downloaded from TCMSP and PubChem databases. Structures were charged and displayed rotatable keys using AutoDock Tools. The protein crystal structures corresponding to the core target genes (top 10 degrees in the PPI network) were downloaded from the Protein Data Bank (PDB) (Burley et al., 2017). The screening conditions were set as follows: 1) the protein structure was obtained using X-ray diffraction; 2) the protein crystal resolution was <3 Å; 3) the species was Homo sapiens. Water and other hetero molecules of proteins were removed using AutoDock Tools, and hydrogen atoms and charge operations of proteins were added. The 3D Grid box for molecular docking simulation was also obtained using AutoDock tools and displayed using AutoDock Vina (Trott & Olson, 2010). Pymol software was used to visualize docking results.

3. Results

3.1. Herb frequency and analysis

The 89 formulae in the analysis included 164 MM herbs, of which 43 were used more than five times, and that with the highest frequency was Glehniae Radix, which occurred 45 times at a frequency of 50.56%. Next, Bambusae Concretio Silicea was identified 44 times (49.43%), and Aucklandiae Radix was identified 41 times (46.06%). There were 43 MM that were used >5 times. Details regarding MM herb identification are included in Table 1.

3.2. Frequency and analysis of Mongolian medicinal categories

According to the classification standard of Neimenggu Mengyao Biao Zhun (Mongolian herbal medicine standards promulgated by Inner Mongolia Autonomous Region), we analyzed the efficacy of high-frequency drugs (use frequency ≥5), and sorted 43 MMs with eight functions used 659 times. Among them, the top three most frequently used herbs had cough-suppressing and anti-hyperventilation (245 times, 37.17%), heat-clearing (199 times, 30.19%), and Qu “Badagan” (78 times, 11.83%) effects. The results of the analyses are shown in Table 2.

3.3. Analysis of MM properties

According to the classification system of Neimenggu Mengyao Biao Zhun, properties and tastes associated with high-frequency drugs (use frequency ≥5) were statistically analyzed. If multiple properties and tastes of MM were identified, they were then counted. Among the 43 MMs, 43 different “nature” and 74 “flavor” were identified; the principal properties of MMs were either cool (observed 18 times, 41.86%) or warm (observed 11 times, 25.58%). The taste most frequently associated with MMs was bitter (observed 25 times, 58.88%) The properties and tastes associated with MMs are shown in Fig. 1.

3.4. Results of association rules analysis

In this study, the R data mining software was used for modeling, while the Apriori algorithm was used to analyze association rules.
of core herbs. The following parameters were applied: support degree $\geq 10\%$ and confidence level $\geq 85\%$ (Xia et al., 2020). A total of 61 herb pairs and herb suit association rules were obtained (Table 3); correlation rules are shown in Fig. 2. The herb pair identified with the highest degree of confidence and strong support was Chebulae Fructus - Toosendan Fructus - Gardeniae Fructus, with a confidence of 100% and a support of 19.31%. According to the Apriori algorithm in association rule, it is considered that the two drug pairs with the highest support are the two drug pairs with the highest probability of simultaneous occurrence in prescription. Therefore, we believe that Chebulae Fructus - Toosendan Fructus - Gardeniae Fructus are the core group medicines of Mongolian medicine for the treatment of PID.

3.5. Cluster analysis results

The R hierarchical clustering algorithm was used to cluster 43 high-frequency drugs (use frequency $\geq 5$), using the R Studio software. The drugs were clustered into six categories, and a cluster dendrogram was generated. The cluster analysis is shown in Fig. 3. We found that the core group medicines screened by association rule analysis were clustered into one group (the fourth group), further illustrating the importance of core group medicine and the accuracy of association rule analysis. Therefore, we took Chebulae Fructus - Toosendan Fructus - Gardeniae Fructus as the core group medicine to further explore its mechanism of action in the treatment of PID.

3.6. Screening active ingredients of CGMs

Active ingredients of each herb included in the three TCM databases (TCMSP, TCMP, and BATMAN-TCM databases) were combined. Toosendan Fructus (Chuanlianai in Chinese, number CLZ) and Chebulae Fructus (Hezi in Chinese, number HZ) have seven

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**Table 1**

Frequency analysis of main Mongolian medicines formulae for treatment of pulmonary infectious diseases.

| Mongolian medicines         | Frequency | Rate (%) |
|----------------------------|-----------|----------|
| Glehniae Radix             | 45        | 50.56    |
| Bambusae Concretio Silicea | 44        | 49.43    |
| Aucklandiae Radix          | 41        | 46.06    |
| Glycyrrhizae Radix et Rhizom | 39   | 43.82    |
| Chebulae Fructus           | 36        | 40.44    |
| Carthami Flrs              | 32        | 35.95    |
| Hippophae Fructus          | 31        | 33.83    |
| Santali Albi Lignum        | 29        | 32.58    |
| Gardeniae Fructus          | 28        | 31.46    |
| Bisorvae Rhizoma           | 23        | 25.84    |
| Biskeiae Scopariae Herba    | 20        | 22.47    |
| Lanceae Herba              | 20        | 22.47    |
| Carophylli Fls             | 19        | 21.34    |
| Pipersi Longi Fructus      | 19        | 21.34    |
| Toosendan Fructus          | 17        | 19.10    |
| Bovis Calculus             | 15        | 16.85    |
| Lignum Pterocarpi          | 13        | 14.60    |
| Alpine Gentian             | 13        | 14.60    |
| Acomitius Kusnezoffii Foli | 11        | 12.35    |
| Lida                     | 11        | 12.35    |
| Rubiae Radix et Rhizoma    | 9         | 12.35    |
| Myris Ticiae Semen         | 9         | 12.35    |

**Table 2**

Frequency of Mongolian medicinal categories.

| Herb category                                | Frequency | Rate (%) |
|----------------------------------------------|-----------|----------|
| Cough-suppressing and panting-calming herbs  | 245       | 37.17    |
| Heat-clearing herbs                          | 199       | 28.37    |
| Qi “Badagian” herbs                          | 78        | 11.83    |
| Zhen “He yi” herbs                          | 33        | 5.46     |
| Qi ta herbs (Chebulae Fructus)               | 24        | 3.64     |
| “Sha nian” herbs                             | 24        | 3.64     |

Qu “Badagian” herbs: Anti-cold herbs; Zhen “He yi” herbs: Regulating Qi movement herbs; Zao “Xie ri wu su” herbs: Dryness and dampness herbs; “Sha nian” herbs: Insecticidal and antifungal herbs.

Fig. 1. Medicinal natures (A) and flavors (B) of 43 MMs.
chemical components, respectively, while *Gardeniae Fructus* (Zhizi in Chinese, number ZZ) has 14. In addition, the assessment revealed that quercetin (number Q) is found in both *Toosendan Fructus* and *Gardeniae Fructus*. Results of CGMs ingredient screening are shown in Table 4.

### 3.7. Prediction of the potential targets of active ingredients

Twenty-nine active ingredients and 702 predicted targets of CGMs were identified; after eliminating overlapping targets, 237 remained. To further clarify interactions between compound and targets, a compound-target network was constructed. As shown in Fig. 4, the MM-compound-target network included a total of 267 nodes (three drugs, 28 compounds, and 237 target nodes) and 686 edges.

### 3.8. Disease prediction

Using OMIM, GeneCards, and DrugBank databases, 653 and 638 disease targets related to VP and PT, were respectively identified.
Targets of both diseases were paired with CGMs component targets, and 79 and 72 CGMs for treating VP and PT, respectively, were identified. Further, 61 targets of CGMs, VP, and PT were shared (Fig. 5).

3.9. Common target multi-level network construction

Cytoscape software was used to construct a compound-target network of active ingredients of CGMs associated with VP, PT, and common targets (Figs. 6 and 7). Key targets are arranged according to degree value in Table 5. By including common targets of CGMs in the treatment of VP and PT in the network, it was possible to obtain a network diagram of the common mechanisms of VP and PT treatment (Fig. 7). The network diagram revealed that CGMs co-treatment effects on VP and PD mainly involved 13 chemical components, including quercetin, kaempferol, and ellipticine, and 61 targets, including PTGS2, PPARG, and EGFR.

Fig. 8 showed a PPI network of 61 targets. We initially entered common targets into the STRING database and imported nodes 1 and 2, and the binding score information from result files into Cytoscape 3.6.1. The resultant network contained 61 nodes and 1,091 edges, which denoted proteins and the interrelations among proteins, respectively. The larger and darker the node, the greater the value indicated. Similarly, the thicker the edge, the greater the combined score of the connected targets. Using a Network Analyzer topology attribute analysis, we found that nodes with higher betweenness and closeness values tended to have a greater degree of connection than those with lower values. Core targets with betweenness of >0.00432403, closeness of >0.73170732, and degree values of >38 were screened. For example, those with values greater than their respective median values were selected. As a result of this analysis, 35 key proteins were identified, and proteins with the top five scores were vascular endothelial growth factor A (VEGFA), interleukin-6 (IL-6), cellular tumor antigen p53 (TP53), and serine/threonine-protein kinase akt-1 (AKT).

3.10. Module and functional enrichment analysis

GO and KEGG enrichment analyses of common targets were performed ($P < 0.05$). The mechanism of drug pair action in the treatment of the disease was also explored. GO function analysis of CGMs in the treatment of VP produced a total of 2,540 GO entries ($P < 0.05$), and the top 10 biological processes (BP), cellular components (CC), and molecular functions (MF) were selected for visualization (Fig. 9A). In the histogram shown, the redder the color, the higher the degree of enrichment and the greater the potential for the component to be used as a drug target. The results of the BP assessment showed that the active components of modified drug pairs in the human body mainly include the response to lipopolysaccharide (GO:0032496), response to the molecule of bacterial origin (GO:0002237), cellular response to chemical stress (GO:0062197), extrinsic apoptotic signaling pathway (GO:0097919), and response to oxidative stress (GO:0006979). Enriched CC mainly included membrane rafts (GO:0045121),...
Table 4
CGMs active ingredient list.

| No. | Compounds       | OB   | DL   | Chemical structures |
|-----|----------------|------|------|---------------------|
| CLZ1| Melianone      | 40.73| 0.81 | ![Chemical structure](image) |
| CLZ2| Stigmasterol   | 43.41| 0.76 | ![Chemical structure](image) |
| CLZ3| Balanophonin   | 54.74| 0.40 | ![Chemical structure](image) |
| CLZ4| Ethyl linolenate| 46.10| 0.20 | ![Chemical structure](image) |
| CLZ5| Medioresil     | 57.20| 0.62 | ![Chemical structure](image) |
| CLZ6| Nimbolidin B   | 30.22| 0.61 | ![Chemical structure](image) |
| HZ1 | Ellagic acid   | 43.06| 0.43 | ![Chemical structure](image) |
| HZ2 | 7-Dehydrosigmasterol | 37.42| 0.75 | ![Chemical structure](image) |
| No. | Compounds                                      | OB (%) | DL (%) | Chemical structures                                      |
|-----|-----------------------------------------------|--------|--------|----------------------------------------------------------|
| HZ3 | Chebulic acid                                 | 72.00  | 0.32   | ![Chemical structure of Chebulic acid](image)             |
| HZ4 | Ellipticine                                   | 30.82  | 0.28   | ![Chemical structure of Ellipticine](image)              |
| HZ5 | Peraksine                                     | 82.58  | 0.78   | ![Chemical structure of Peraksine](image)                |
| HZ6 | (R)-(6-methoxy-4-quinolyl)-[(2R,4R,5S)-5-vinylquinuclidin-2-yl]methanol | 55.88  | 0.40   | ![Chemical structure of (R)-(6-methoxy-4-quinolyl)-[(2R,4R,5S)-5-vinylquinuclidin-2-yl]methanol](image) |
| HZ7 | Cheilanthifoline                              | 46.51  | 0.72   | ![Chemical structure of Cheilanthifoline](image)         |
| ZZ1 | Crocetin                                      | 35.30  | 0.26   | ![Chemical structure of Crocetin](image)                 |
| ZZ2 | Ammidin                                       | 34.55  | 0.22   | ![Chemical structure of Ammidin](image)                  |
| ZZ3 | Sudan III                                     | 84.07  | 0.59   | ![Chemical structure of Sudan III](image)                |
| ZZ4 | Beta-sitosterol                                | 36.91  | 0.75   | ![Chemical structure of Beta-sitosterol](image)          |

(continued on next page)
### Table 4 (continued)

| No. | Compounds                                             | OB   | DL  | Chemical structures          |
|-----|-------------------------------------------------------|------|-----|-----------------------------|
| ZZ5 | Kaempferol                                            | 41.88| 0.24|                             |
| ZZ6 | Stigmasterol                                          | 43.83| 0.76|                             |
| ZZ7 | Mandenol                                              | 42.00| 0.19|                             |
| ZZ8 | Isoimperatorin                                        | 45.46| 0.23|                             |
| ZZ9 | Ethyl oleate (NF)                                     | 32.40| 0.19|                             |
| ZZ10| 5-hydroxy-7-methoxy-2-(3,4,5-trimethoxyphenyl)chormone| 51.96| 0.41|                             |
| ZZ11| 3-Methylkempferol                                     | 60.16| 0.26|                             |
| ZZ12| Artemisetin                                           | 49.55| 0.48|                             |
| ZZ13| Sitosterol                                            | 36.91| 0.75|                             |
membrane microdomains (GO:0098857), membrane regions (GO:0098589), transferase complexes, and the transfer of phosphorus-containing groups (GO:0061695). Enriched MF mainly included cytokine receptor binding (GO:0005126), cytokine activity (GO:0005125), receptor ligand activity (GO:0048018), signaling receptor activator activity (GO:0030546), and protein phosphatase binding (GO:0019903).

Through a KEGG pathway enrichment analysis, a total of 172 related signaling pathways affected by CGMs in the treatment of VP were identified \((P < 0.05)\), and the top 30 items were listed for visual analysis (Fig. 9B). In the bubble diagram shown, the absissa indicates the gene ratio, the color corresponds to the degree of enrichment, and the bubble size indicates the number of included genes. Enriched pathways identified mainly included the AGE-RAGE signaling in diabetic complications, lipid and atherosclerosis, fluid shear stress and atherosclerosis, Kaposi sarcoma-associated herpesvirus infection TNF signaling, and IL-17 signaling.

A GO function analysis of drug pairs used to treat PT produced a total of 2,435 GO entries \((P < 0.05)\); the top 10 BP, CC, and MF are visualized in Fig. 9C. BP affected by drug pairs included the response to lipopolysaccharide (GO:0032496), cytokine activity (GO:0005125), receptor ligand activity (GO:0048018), signaling receptor activator activity (GO:0030546), and protein phosphatase binding (GO:0019903).

Table 4 (continued)

| No. | Compounds | OB   | DL   | Chemical structures |
|-----|-----------|------|------|---------------------|
| Q   | Quercetin | 46.43| 0.28 | ![Chemical Structure](image) |

Note: CLZs (1–6) represent active compounds in Toosendan Fructus, HZs (1–7) represent the active compounds of Chebulae Fructus, and ZZs (1–13) represent the active compounds of Gardeniae Fructus. Qi is Toosendan Fructus and Gardeniae Fructus common active compounds.
Through a KEGG pathway enrichment analysis, a total of 162 related signaling pathways of drug pairs in the treatment of PT were obtained \((P < 0.05)\); the top 30 items were listed for visual analysis (Fig. 9D). Pathways determined to be enriched mainly included AGE-RAGE signaling in diabetic complications, fluid shear stress and atherosclerosis, lipid and atherosclerosis, TNF signaling, and IL-17 signaling.

A GO function analysis of common targets of drug pairs produced 2,391 GO entries \((P < 0.05)\), and the top 10 BP, CC, and MF were visualized (Fig. 9E). The BP analysis revealed that active components of modified drug pairs in the human body mainly affect the response to lipopolysaccharides (GO:0032496), response to molecules of bacterial origin (GO:0002237), response to chemical stress (GO:0062197), cellular response to biotic stimulus (GO:0071216), and metabolism of reactive oxygen species (GO:0072593). These results suggest that CGMs may affect the treatment of PID by regulating the biological processes identified.

CC identified mainly included membrane rafts (GO:0045121), membrane microdomains (GO:0098857), membrane regions (GO:0098589), and the vesicle lumen (GO:0031983). Further, MF identified mainly included cytokine receptor binding (GO:0005126), cytokine activity (GO:0005125), receptor ligand activity (GO:0048018), and signaling receptor activator activity (GO:0030546). These MF may play important roles in the occurrence and development of PID.

Through KEGG pathway enrichment analysis, a total of 159 related signaling pathways of drug pairs of common targets were obtained \((P < 0.05)\), and the top 30 items were listed for visual analysis (Fig. 9F). Enriched pathways mainly included the AGE-RAGE signaling pathway in diabetic complications, lipid and atherosclerosis, TNF signaling, fluid shear stress, atherosclerosis, and IL-17 signaling. The IL-17 signaling pathway, which plays crucial roles in both acute and chronic inflammatory responses, was selected as an example (Fig. 10). Red labeled nodes indicate the targets of CGMs, which shows that CGMs play a key role in modulating the IL-17 signaling pathway by regulating marker targets.

### 3.11. Molecular docking

Among the core targets of the PPI network (Fig. 8), we selected those with the top 10 scores for subsequent molecular docking analyses. To analyze the docking results, findings were depicted using a heat map, as shown in Fig. 11. The minimum binding energy needed for binding of each compound to a key protein was determined (Fig. 12). As a result, a semiflexible docking model was adopted. For example, a rigid configuration of the protein versus a flexible configuration of small-molecules was considered. Grid energy was calculated using AutoGrid, and the docking operation was based on a Lamarck genetic algorithm. Docking results showed that the binding energy required for the binding of a compound and a protein was \(< -5 \text{ kcal/mol}\), indicating that each compound and protein was capable of binding tightly.

### 4. Discussion

#### 4.1. Three core medicines we mined are Sanzi Decoction, a famous classical formula in Mongolian medicine

In this study, we found that Toosendan Fructus-Chebulae Fructus-Gardeniae Fructus was the core group medicine of Mongolian medicine for the treatment of PID. “Toosendan Fructus-Chebulae Fructus-
Fig. 7. CGMs treatment of PID "disease-compound-target" network. Common target mechanism network; 13 active compounds are shown as blue circles and 61 common target genes are shown as purple circles.

Table 5
CGMs’s targets for treatment of PT, VP and PID for three diseases (Top ten degree).

| Diseases | Number of targets | Target name (Top 10 degree) |
|----------|-------------------|-----------------------------|
| PT       | 79                | PTGS2, ADRB2, PIK3CG, BAX, BCL2, CASP3, NOS2, GSTM1, CYP1A1, MAPK14 |
| VP       | 72                | PTGS2, DPP4, PIK3CG, ESR1, AR, BAX, BCL2, CASP3, NOS3, NOS2 |
| PID      | 61                | VEGFA, IL6, TP53, AKT1, TNF, CXCL8, IL1B, PTGS2, MMP9, JUN |

Note: TNF, tumor necrosis factor; CXCL8, interleukin-8; IL1B, interleukin-1 beta; PTGS2, prostaglandin G/H synthase 2; MMP9, matrix metalloproteinase-9; JUN, transcription factor AP-1.

Fig. 8. PPI network of common targets by STRING database (A) and Cytoscape software (B).
Fig. 9. GO enrichment analysis (A, C, E) and KEGG analysis (B, D, F) of therapeutic targets of CGMs for VP (A, B), PT (C, D) and common targets (E, F).

Fig. 10. IL-17 signaling pathway. CGMs may further regulate IL-17 signaling pathway by affecting IL6, PTGS2, TNF, JUN, CXCL8, MMP9 and other targets to achieve anti-inflammatory purposes.
Gardeniae Fructus is the composition of the famous classical formulas of Mongolian medicine Sanzi Decoction (it is composed of three herbs of Toosendan Fructus, Chebulae Fructus, Gardeniae Fructus) (Sa & Siqin, 2019), which was first published in the Four Medical Tantras (Wu, Du, & Jin, 2015).

Gardeniside is one of the active and toxic components of Chebulae Fructus-Gardeniae Fructus in Sanzi Decoction. The combination of the three medicines can increase the curative effect and reduce the toxicity of gardeniaside. (Wang, Bao, & Wang, 2015). Sanzi Decoction has the functions of clearing away heat and detoxification and cooling blood, and can be used to treat plague. The Mongolian Medicine Jingui also records that the plague can be treated with Sanzi Decoction for 4–6 d, with heat-clearing and detoxifying (Li & Tu, 2020). Pharmacological studies have found that Sanzi Decoction has the effects of improving immunity, anti-inflammatory and antibacterial (Luo, Liu, Zhou, & Yun, 2001). It has significant antibacterial effect on Staphylococcus aureus, Pseudomonas aeruginosa and Streptococcus (Zhang, Wang, & Bao, 2009). Therefore, it is the traditional basis for analysing the therapeutic mechanism of “Toosendan Fructus-Chebulae Fructus-Gardeniae Fructus” in the treatment of pulmonary infectious diseases by network pharmacology.

4.2. CGMS treat PID mainly through anti-inflammatory and antibacterial effects

4.2.1. Quercetin, kaempferol, β-sitosterol and stigmasterol might be responsible compounds

In this study, the potential mechanism of “Toosendan Fructus-Chebulae Fructus-Gardeniae Fructus” in treating PID was explored by using network pharmacology method. Among the active ingredients screened, quercetin has a wide range of biological activities, including those that are antiviral, anti-inflammatory, immunomodulatory (Li et al., 2016), and antioxidant (Xu, Hu, Wang, & Cui, 2019). It can reduce inflammatory damage in lipopolysaccharide-caused WI-38 lung fibroblasts by inhibiting NF-κB and JNK signaling pathways, and also inhibit early-stage influenza A virus infection and viral invasion (Wu et al., 2015). Kaempferol has anti-inflammatory, antibacterial, antioxidative stress activity, and anti-tumoral effects (Zhang et al., 2017). It can inhibit the expression of inflammatory cytokines TNF-α, IL-6, IL-1β and PGE2, improve pulmonary edema, and also inhibit the phosphorylation of NFκB and MAP kinase, exerting anti-inflammatory effects in vitro and in vivo (Sun et al., 2018). Studies showed that β-sitosterol has immunomodulatory, anti-inflammatory, antioxidant and anti-tumor effects (Liu, Ji, & Huang, 2019; Zhi et al., 2020). It can play an anti-inflammatory effect by inhibiting the synthesis of NO and the activity of interleukin-6 (IL-6), and reducing the secretion of inflammatory factors such as IL-1 and TNF-α (Choi et al., 2012). Stigmasterol can exert anti-inflammatory, immune regulation, and increase blood oxygen saturation by acting on CASP3, PTGS2, NOS2, NOS3 and other targets (Lin et al., 2006). Stigmasterol and β-sitosterol also have certain anti-mycobacterial activity (Dai, 2009). At the same time, we found that quercetin, kaempferol, β-sitosterol and stigmasterol are also key compounds of Qingfei Paifu Decoction (Ephedrae Herba, Glycyrrhizae Radix et Rhizoma Preparata Cum Melle, Armeniacae Semen Amarum, Gypsum Fibrosum, Cinnamonomum Ramulus, Alismatis Rhizoma, Polyporus, Atractylodis Macrocephalae Rhizoma, Poria, Bupleuri Radix, Scutellariae Radix, Pinelliae Rhizoma Preparatum cum Zingibere et Alumine, Zingiberis Rhizoma Recens, Asteris Radix et Rhizoma, Farfarae Flos, Belamcandae Rhizoma, Asari Radix et Rhizoma, Dioscoreae Rhizoma, Aurantii Fructus Immaturus, Citri Reticulatae Pericarpium, Pogostemonis Herba) in the treatment of COVID-19 (Ren, Yang, Zhang, &

Fig. 11. Heat map of binding energy of active ingredient and core target molecule docking. The horizontal is the active ingredient, and the vertical is the core target. Each number represents the binding energy between the active ingredient and the core target. The darker the color, the lower the binding energy and the more stable the binding.
It shows that these compounds may have the effect of inhibiting virus replication and inhibiting the binding of virus spike protein to ACE2, which can prevent SARS-CoV-2 from infecting cells.

4.2.2. VEGFA, IL6, TP53, AKT1, and TNF might be key targets for treatment of PID

The core targets of VEGFA, IL6, TP53, AKT1 and TNF were obtained by protein–protein interaction network analysis, suggesting that these targets play a key role in the treatment of PID with “Toosendan Fructus-Chebulae Fructus-Gardeniae Fructus”. VEGFA is one of the main factors that stimulate angiogenesis, and plays an important role in activating signaling pathways and inducing endothelial cell proliferation and migration (Claesson & Wslsh, 2013). Studies showed that elevated VEGFA levels are likely to be associated with resolution of lung inflammation (Strouvalis et al., 2018). As a pro-inflammatory cytokine, IL-6 can stimulate the inflammatory process in various diseases (Tanaka, Narazaki, & Kishimoto, 2014). Cytokine storm is an important part of the severe response of COVID-19 patients, and IL6 plays a key role in cytokine release syndrome, which has been used as a clinically important indicator of patient deterioration (Zhang, Wu, Li, Zhao, & Wang, 2020). At the same time, WHO studies have confirmed that IL-6 inhibitors can reduce the risk of death in hospitalized patients with COVID-19 (Tharaux, 2021). TP53 can regulate cell cycle arrest and apoptosis. The experimental study of mycobacterium tuberculosis

![Diagram of compound and key target molecule. The combination with the lowest binding energy to the core target for each active ingredient.](image)

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**Fig. 12.** Docking modl diagram of compound and key target molecule. The combination with the lowest binding energy to the core target for each active ingredient.
infection of alveolar epithelial cells showed that, p53 can resist the invasion of Mycobacterium tuberculosis by inhibiting the activation of TLR-4 and NF-κB and reducing the secretion of inflammatory factors (Wang et al., 2019). AKT1 can regulate the proliferation and growth of cells. By acting on JNK, it further promotes cell apoptosis and reduces the viability of mycobacterium tuberculosis (Zhang et al., 2021). TNF can mediate cell apoptosis, inflammatory response and other processes, and can activate other downstream inflammatory factors, attracting inflammatory cells to the respiratory tract (Tang et al., 2002). At the same time, it was found that TNF is closely related to lung injury and pulmonary fibrosis (Liang, Yuan, Wang, Lei, & Ouyang, 2020).

4.2.3. AGE-RAGE signaling pathways, IL-17 signaling pathways, and TNF signaling pathways might be regulated by CGMs

The results of the KEGG pathway enrichment analysis of “Toosendan Fructus-Chebulae Fructus-Cardiaceae Fructus” in the treatment of PID showed that it was mainly enriched in AGE-RAGE signaling pathways, IL-17 signaling pathways, and TNF signaling pathways. The AGE-RAGE signaling pathway can enhance the expression of VEGF, TGF-β, and MCP-1, and promote the release of inflammatory factors, thereby accelerating the inflammatory response, cell apoptosis, and tissue damage (Wendt et al., 2003). Relevant studies showed that the overexpression of IL-17 can cause a large number of neutrophils to aggregate and secrete inflammatory factors, activate the inflammatory response, and cause alveolar endothelial damage and increased tissue permeability (Liu & Wen, 2021). In addition, IL-17 in the peripheral blood of patients with severe pneumonia was significantly increased, and the level of IL-17 was closely related to the development and severity of the patient's disease, suggesting that IL-17 may be a cytokine that plays an important role in the inflammatory response system (Cai et al., 2017). TNF signaling pathway plays an important role in regulating inflammation and immune response in vivo. Relevant studies have shown that the level of TNF-α and the incidence of ARDS in severe patients with COVID-19 are higher than those in non-severe patients (Li et al., 2020). At the same time, the levels of TNF-α and TNF-β in the serum of pulmonary tuberculosis patients were significantly increased, which made Mycobacterium tuberculosis in a relatively static and stable state by promoting tissue damage and further activation of macrophages and phagocytosis of bacteria (Li et al., 2016).

4.3. Limitations of this study

First, due to the small number of included prescriptions, the reliability of the results was affected to a certain extent. Therefore, it is expected that more prescriptions can be collected in the future to improve the accuracy of the results. Second, network pharmacology is a research technology based on network data and computer simulation analysis, and the database used is not updated timely enough. Therefore, the obtained results still need further experimental verification.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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