Network pharmacology-based analysis of the role of traditional Chinese herbal medicines in the treatment of COVID-19

Shengwei Yu, Junwu Wang, Haitao Shen

Department of Emergency Medicine, Shengjing Hospital of China Medical University, Shenyang 110004, China

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Correspondence to: Haitao Shen. Department of Emergency Medicine, Shengjing Hospital of China Medical University, 36 Sanhao Street, Shenyang 110004, China. Email: shenht@sj-hospital.org.

Background: The novel coronavirus named COVID-19, which originated in Wuhan, China, has spread to many countries around the world. Currently, no effective medical treatment exists to combat this disease. Traditional Chinese herbal medicines (CHM) have unique roles in the treatment of viral infections. In this article we analyzed the effectiveness and possible molecular mechanisms of CHM formulas for the prevention of COVID-19.

Methods: The active ingredients and action targets of CHM formulas were obtained from the TCMSP database. Genes related to severe acute respiratory syndromes (SARS) and Middle East respiratory syndrome (MERS) were queried on the GeneCards database. The action mechanisms of these genes were predicted using a Gene Ontology (GO)-based functional enrichment and annotation tool and the Kyoto Encyclopedia of Genes and Genomes (KEGG).

Results: CHM formulas played a positive role in preventing COVID-19 and warrant further application.

Conclusions: Our research provides new evidence to support the possible value of CHM formulas for the prevention of COVID-19. However, further clinical studies with large sample sizes are required to verify their effectiveness.

Keywords: Novel coronavirus; COVID-19; traditional Chinese herbal medicines (traditional CHM); network pharmacology

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Introduction

As 2019 drew to a close, several cases of pneumonia of unknown etiology were reported in Wuhan, the capital of Hubei province in China (1). Up to now, the outbreak of COVID-19, caused by a new coronavirus, Genes related to severe acute respiratory syndromes (SARS) and Middle East respiratory syndrome (MERS) were queried in the GeneCards database (2). As of February 20, 2020, based on data released by China CDC (http://2019ncov.chinacdc.cn/nCoV/), the total number of confirmed cases was 74,680, with 2,122 of these cases resulting in death. At present, no drug exists to effectively treat or prevent COVID-19 (3). In addition, immune response to the new coronavirus differs according to the individual, leading to a variety of clinical symptoms, disease course, and treatment response to drugs and vaccines (4). Based on the National Health Commission of China Diagnosis and Treatment Protocol for COVID-19 Infection (Sixth Edition) as well as the experience of managing the virus in Hubei province (especially in Wuhan), the Hubei Province Diagnosis and Treatment Protocol for COVID-19 has been developed and released. In our current study, we analyzed data relevant to exploring the
effectiveness and possible action mechanism of traditional Chinese herbal medicine (CHM) formulas for the prevention and treatment of the novel coronavirus.

**Methods**

**Collection of active ingredients**

Two CHM formulas were obtained from the *Hubei Province Diagnosis and Treatment Protocol for COVID-19*: Formula A: Rhizoma Atractyloidis, Flos Lonicerae, Pericarpium Citri Reticulatae, Rhizoma Phragmitis, Foliun Mori, and Radix Astragali seu Hedysari; and Formula B: Radix Astragali seu Hedysari, Rhizoma Atractylodis Macrocephalae, Radix Saposhnikoviae, Cyrtomium fortunei J. Sm., Flos Lonicerae, Eupatorium fortunei Turcz., and Pericarpium Citri Reticulatae.

**Screening for active ingredients and target genes**

Oral bioavailability (OB) reflects the rate of absorption of an orally administered drug that enters the circulation via the liver after absorption into the gastrointestinal tract. Drug-likeness (DL) refers to the structural similarity of herbal ingredients to a known drug. There is a positive correlation between these two descriptors. With the help of the TCMSP data platform (5) (http://tcmspw.com/tcmsp.php), the chemical constituents in the compounds in Formulas A and B that had OB ≥30% and DL ≥0.18 were retrieved as the active ingredients in the study. A high OB value often reflects better DL (6).

**Screening of disease targets**

No data on COVID-19-related genes were available in the GeneCards (https://www.genecards.org/) database at the time of study. Since the new coronavirus is highly similar to SARS-CoV and MERS-CoV (7,8), especially the bat SARS-like coronavirus (Genbank accession number MG772933), “Severe Acute Respiratory Syndromes” (SARS) and “Middle East Respiratory Syndrome” (MERS) were used in our study to search for genes that may be associated with the new coronavirus.

**Screening of COVID-19-related genes acted by the active CHM ingredients**

A Venn diagram was created to visualize the amount of overlap between the genes related to the active CHM ingredients and COVID-19-related genes.

**Construction of protein-protein interaction (PPI) networks and screening of its core network**

The protein-protein interaction core network (PPICN) refers to the correlation between compounds and disease-related protein molecules, taking into account biochemistry, signal transduction, and genetic networks. The obtained intersection genes were uploaded onto STRING11.0 (http://string-db.org/cgi/input.pl) to obtain the relationships of PPIs.

**Comparison and analysis of target pathways**

We analyzed the core genes screened out from the two formulas, as well as their roles in signaling pathways, to explore their functions. To achieve this, a Gene Ontology (GO)-based functional enrichment and annotation tool and the Kyoto Encyclopedia of Genes and Genomes (KEGG) were used. Data were obtained from the website (http://bioconductor.org/biocLite.R) and the results were visualized with Rstudio.

**Results**

**Active compounds**

Some of the compounds (i.e., those with the 10 highest OB values) in these two formulas were retrieved through the Traditional Chinese Medicine Systems Pharmacology Database and Analysis Platform (TCMSP). OB ≥30% and DL ≥0.18 served as the criteria (*Tables 1,2*).

**Core action genes in the CHM formulas**

Aided by the TCMSP database, the genes linked to the effects of the active ingredients in these two CHM formulas were obtained. Genes related to COVID-19 were retrieved from the GeneCard database using the keywords “SARS” and “MERS”. Gene intersections were generated based on the intersection results from these two groups (*Figures 1,2*).

**Construction and screening of PPICN between CHM formulas and COVID-19**

The 217 and 203 intersected genes were analyzed using the
STRING software, with a coefficient of 0.990 indicating correlation. The PPICN (PPI correlation network) of Formula A contained 217 nodes and 99 connections, with the average degree of nodes being 0.912. Formula B contained 203 nodes and 98 connections, with the average degree of nodes being 0.314 (Figures 3,4).

Target pathways
GO-based functional enrichment and annotation of Formula A yielded 192 GO entries (P<0.05). The 20 pathways with the highest GO enrichment are shown in Figures 5 and 6.
GO-based functional enrichment and annotation of

| Table 1 Basic information of some active compounds in Formula A |
|---|---|---|---|
| No. | Chemical composition | Relative molecular weight | OB% |
| MOL000379 | 9,10-dimethoxypterocarpan-3-O-b-D-glucoside | 462.49 | 0.92 |
| MOL000729 | Oxysanguinarine | 347.34 | 0.87 |
| MOL000211 | Mairin | 456.78 | 0.78 |
| MOL000033 | (3S,8S,9S,10R,13R,14S,17R)-10,13-dimethyl-17-[(2R,5S)-5-propan-2-yloctan-2-yl]-2,3,4,7,8,9,11,12,14,15,16,17-dodecahydro-1H-cyclopenta[a]phenanthren-3-ol | 428.82 | 0.78 |
| MOL000184 | NSC63551 | 412.77 | 0.76 |
| MOL000186 | Stigmasterol 3-O-beta-D-glucopyranoside | 412.77 | 0.76 |
| MOL000092 | Daucosterin | 414.79 | 0.76 |
| MOL000094 | Daucosterol | 414.79 | 0.76 |
| MOL003036 | ZINC03978781 | 412.77 | 0.76 |
| MOL000449 | Stigmasterol | 412.77 | 0.76 |

OB, oral bioavailability.

| Table 2 Basic information of some active compounds in Formula B |
|---|---|---|---|
| No. | Chemical composition | Relative molecular weight | OB% |
| MOL000379 | 9,10-dimethoxypterocarpan-3-O-b-D-glucoside | 462.49 | 0.92 |
| MOL000592 | Dammaraadienyl acetate | 468.84 | 0.82 |
| MOL000211 | Mairin | 456.78 | 0.78 |
| MOL000033 | (3S,8S,9S,10R,13R,14S,17R)-10,13-dimethyl-17-[(2R,5S)-5-propan-2-yloctan-2-yl]-2,3,4,7,8,9,11,12,14,15,16,17-dodecahydro-1H-cyclopenta[a]phenanthren-3-ol | 428.82 | 0.78 |
| MOL000033 | (3S,8S,9S,10R,13R,14S,17R)-10,13-dimethyl-17-[(2R,5S)-5-propan-2-yloctan-2-yl]-2,3,4,7,8,9,11,12,14,15,16,17-dodecahydro-1H-cyclopenta[a]phenanthren-3-ol | 428.82 | 0.78 |
| MOL000028 | β-Amyrins | 426.8 | 0.76 |
| MOL003036 | ZINC03978781 | 412.77 | 0.76 |
| MOL000449 | Stigmasterol | 412.77 | 0.76 |
| MOL000449 | Stigmasterol | 412.77 | 0.76 |
| MOL000296 | Hederagenin | 414.79 | 0.75 |

OB, oral bioavailability.
Figure 1 The intersection generated between genes related to the effects of the active ingredients of Formula A and COVID-19-associated genes.

Figure 2 The intersection generated between genes related to the effects of the active ingredients of Formula B and COVID-19-associated genes.

Figure 3 The PPICN between Formula A and COVID-19. Each node represents a protein and the connections represent the interaction between two proteins. A thicker connection represents higher correlation. PPICN, protein-protein interaction core network.
**Figure 4** The PPICN between Formula B and COVID-19. Each node represents a protein and the connections represent the interaction between two proteins. A thicker connection represents higher correlation. PPICN, protein-protein interaction core network.

**Figure 5** The 20 most enriched GO pathways in Formula A (ranking by P value). GO, Gene Ontology.
Formula A yielded 192 GO entries (P<0.05). Twenty pathways with highest GO enrichment are shown in Figures 7 and 8.

The KEGG pathways of Formulas A and B were enriched and screened to obtain the 20 most enriched signal pathways (P<0.05), including the hepatitis B pathway, Kaposi sarcoma-associated herpesvirus (KSHV) infection-related pathways, and human cytomegalovirus infection-related pathways (Figures 9-12).

**Discussion**

In the absence of drugs to effectively treat COVID-19 infection, CHM offers unique antiviral benefits. For instance, astragalus polysaccharide at a non-cytotoxic concentration (30 μg/mL) significantly suppresses the expressions of two early viral proteins (Zta and Rta) in the Epstein-Barr virus lytic cycle to exert an antiviral effect (9). CHM can directly inhibit respiratory pathogens or coordinate immune system activity to prevent or alleviate
A combination of traditional Chinese and Western medicine was capable of promoting the absorption of pulmonary infiltrates in SARS patients (11). In addition to their antiviral benefits, CHMs can also be applied at the rehabilitative stage (12). Therefore, CHM may be effective for the treatment of COVID-19 (13).

By using the technology offered by network pharmacology, we analyzed the molecular mechanisms of two CHM formulas for the prevention of COVID-19. Although different herbal drugs were used in these two formulas, the results of the PPI networks, GO-based enrichment analysis, and KEGG enrichment analysis were fairly similar, which is not coincidental.

The internal regulation of the body involves a complex regulatory network rather than a single signaling pathway. Signal transduction exists among different signaling pathways, and the interaction of different signaling pathways produces diverse biological effects. Therefore, in order to target a single signaling pathway, we need to first identify the key proteins in this signaling pathway and then target these proteins. However, this approach may not be effective in treating COVID-19, which may involve multiple signaling pathways. One possible explanation for the similarity of the results of the two CHM formulas is that the molecular mechanisms of these formulas may involve multiple signaling pathways, which may be coincidental.

Figure 8 The 20 most enriched GO pathways in Formula B (ranking by number of enriched genes). GO, Gene Ontology.

Figure 9 The 20 most enriched KEGG pathways in Formula A (ranking by P value). KEGG, Kyoto Encyclopedia of Genes and Genomes.
pathways and targets; therefore, the therapeutic effects of drugs are not just a result of direct targeting. Instead, they more commonly directly regulate the target while indirectly regulating other targets. Data on PPIs facilitate our understanding of the regulatory roles of targets. Akt, as the core target of CTHM formulas, plays an important role in containing coronavirus infections. In SARS patients, the phosphorylation level of the cell survival protein Akt is downregulated in cells expressing M protein. Meanwhile, the overexpression of 3-phosphoinositide-dependent protein kinase-1 (PDK1), an upstream kinase for Akt, inhibits M protein-induced apoptosis, indicating that M-protein perturbs the PDK1 and PKB/Akt cell survival signaling pathway (14). Through Akt activation,
which inhibits apoptosis, CHM formulas can suppress viral replication (15), reduce apoptosis, and thereby repair damage to the body. Based on PPI networks, we can further investigate the targets via which CHM formulas affect the occurrence and development of COVID-19, map the PPI target networks, and thus pave the way for further network analysis and investigation of their underlying mechanisms.

In the GO-based enrichment analyses, the most enriched biological processes of these two CHM formulas included nuclear receptor activity, transcription factor activity, and direct ligand regulated sequence-specific DNA binding pathways. Nuclear receptors (NRs) are a superfamily of ligand-dependent transcription factors that regulate a variety of biological processes including growth and development, metabolism, and inflammation (16,17). After coronavirus infection, some of the NR-encoded proteins (e.g., nsp1) can inhibit the translation of the host without seriously affecting the expressions of viral genes (18). Fortunately, the targets of CHM formulas are enriched in NR activity, which, to a certain extent, can alleviate the damage caused by coronavirus infection. However, whether viral replication can be suppressed via the activation of coronavirus membrane fusion, which takes place through a receptor-driven ratcheting mechanism, remains unclear and further verification is required (19).

The KEGG analysis showed that a vast majority of the most enriched pathways in these two formulas were associated with viral infections. These pathways included the cytomegalovirus infection, hepatitis B virus infection, and PI3K/Akt signaling pathways. Notably, the PI3K/Akt pathway serves as a central regulator of many important processes that control translation, metabolism, and apoptosis. Active PI3K/Akt signals can meet the needs of replication for many viruses. When the "proviral" kinase is activated, it is also involved in the host’s response to viral infection and ultimately inhibits viral replication (20).

**Limitations and prospects**

CHMs have been widely recognized not only for their multiple biologically active ingredients and multiple pharmacological activities, but also their production of other biologically active or inactive metabolites when delivered in vivo. Therefore, it is difficult to determine whether the antiviral activity of a CHM is a consequence of a single and precise action mechanism or a result of a synergistic therapeutic effect. Furthermore, the medicinal properties or toxicities of CHM may be influenced by many other factors, including the methods of processing, combining, and frying involved with some medicines (21,22).

In summary, a network pharmacology method was used in our current study to investigate the effectiveness of CHM for COVID-19. Although the results were promising, more clinical trials are warranted for our findings to be confirmed.
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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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