Network Pharmacology Prediction and Molecular Docking-Based Strategy to Discover the Potential Pharmacological Mechanism of Wen-yu-jin Against Pulmonary Fibrosis in Mouse Model

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Abstract

Background

Pulmonary fibrosis (PF) is a devastating lung disease. The two drugs approved by the FDA, pirfenidone and nintedanib, can only delay the progression of the disease but cannot cure the disease. These drugs also present adverse effects. Wen-yu-jin (WYJ) obtained from steamed roots of Curcuma wenyujin showed a variety of pharmacological activities. In this study we investigated whether WYJ present anti-lung fibrosis effects.

Methods

Ultra-high pressure liquid chromatography combined with linear ion trap-orbital tandem mass spectrometry (UHPLC-LTQ-orbital trap) was used to identify chemical composition of WYJ. PF-related and WYJ-related targets were obtained from public databases. Network pharmacological was performed to acquire potential targets and major signaling pathways. The binding activity of composition with core targets was predicted by molecular docking. Based on the predicted results, the anti-lung fibrosis effect of WYJ was verified in vivo and in vitro experiments.

Results

23 major compositions of WYJ were identified based on UHPLC-LTQ-Orbitrap. According to the results of network pharmacology, MAPK signaling pathway might play an important role in WYJ against lung fibrosis and STAT3 also could be the potential therapeutic targets. Molecular docking results indicated that most of the compositions have good binding activities with core targets. In vivo and in vitro experiments showed that WYJ alleviated process of fibrosis by inhibiting MAPK signaling pathway and the levels of phosphorylated STAT3 (p-STAT3).

Conclusion

According to the results of network pharmacology and molecular docking, in vivo and in vitro experiments further verified potential targets and molecular mechanism of WYJ against lung fibrosis. Our study provided a novel approach to explain the pharmacological basis of other herbs.

Introduction

PF characterized by bilateral pulmonary interstitial infiltration, restriction on pulmonary function testing, and progressive dyspnea with respiratory failure[1–3], is an irreversible lung disease[4]. Idiopathic pulmonary fibrosis (IPF) is the most common type of pulmonary fibrosis. PF susceptibility is closely related to aging, which leads to telomeres shortening and mitochondrial dysfunction[5]. Accumulating exposures to numerous risk factors[6] such as smoking, occupational dust, drug stimulation, bacterial and virus infection, also result in PF. Pirfenidone and nintedanib[7, 8] approved by FDA are the only two effective drugs for clinical PF medical therapies. However, these drugs only can delay the progression of the disease and maintain lung function but cannot cure the disease[9, 10]. And these drugs can cause adverse effects[10–12], resulting in treatment discontinuation and adverse gastrointestinal effects.

Multiple active ingredients and targets of Traditional Chinese Medicines (TCMs) arouse the attention of pharmacologists. Previous studies showed that TCMs or active ingredients from TCMs have pulmonic protective benefits[13], which represent an attractive source of drug discovery for treating PF. In TCMs, there are many theories according to the pathogenesis of PF, such as qi stagnation and blood stasis (Qi-Zhi-Xue-Yu in Chinese) and binding of phlegm and qi (Tan-Qi-Hu-Jie in Chinese). Wen-yu-jin (WYJ, Curcuma Radix, [III]), derived from the steamed root of Curcuma wenyujin, has been used for at least 1500 years in China and was mainly dependent on move qi and activate blood to intervene diseases.

In our study, a novel approach combining TCMs with network pharmacology was established to explore the molecular mechanism of WYJ against PF. Molecular docking was performed to predict the affinity strength between the active Compounds and the important targets. Finally, the anti-fibrosis mechanism of WYJ was further verified via in vitro and in vivo experiments based on the predicted results.

Materials And Methods

Preparation of Chinese Medicine

WYJ was purchased from Tong Ren Tang Co., Ltd. (Beijing, China). The herb was boiled in purified water twice for 1h each time. Lyophilized powder was obtained by condensation and freeze-dry of liquids. Lyophilized powder was stored at -80°C and used for a series of experiments.

LC/MS Analysis

The coupling of UHPLC System (Thermo Fisher Scientific) equipped with a binary pump, an autosampler, a column thermostat and DAD detector and LTQ-Orbitrap XL (Thermo Fisher Scientific) equipped with an electrospray ionization source (ESI) was used For LC/MS experiments. Data was controlled and processed by Xcalibur software (Thermo Fisher Scientific). The column used in the LC analysis was Thermo Scientific Hypersil BDS C18
Histopathological examination

The level of hydroxyproline in the lung tissue was measured at the absorbance of 550nm and the results were expressed with µg/ lung. The lung tissue was processed according to the instructions of the Hydroxyproline Kit (Nanjing Jiancheng Institute of Bioengineering, Nanjing, China).

Hydroxyproline determination (HYP)

The body weight (BW) and the whole lung weight (PW) were recorded after mice were sacriced. Pulmonary indexes (PI) were calculated using the equation PI = PW/BW*100.

Animals and experimental design

75 male mice (18-22g, 8 weeks old) were purchased from Beijing Vital River Laboratory Animal Technology Co., Ltd. (Beijing, China). The mice were anesthetized to reduce pain in the procedures. Mice of the Control group were given normal saline via oropharynx, and other mice were given bleomycin (BLM) via oropharynx at a dose of 5 mg/kg to establish PF model. BLM-induced PF model were randomly divided into four groups: BLM treatment group (Model); BLM with WYJ high concentration treatment group (WYJ-H); BLM with WYJ medium concentration treatment group (WYJ-M); and BLM with WYJ low concentration treatment group (WYJ-L).

Control group and Model group were treated with saline. According to the previous reports and the results of pre-experiments, WYJ-H, WYJ-M and WYJ-L groups were given WYJ at doses of 1.2, 0.6, and 0.3g/kg respectively. All the treatments were initiated 7 days after the establishment of PF model with bleomycin, once a day for 14 days. All the mice were sacriced on day 22. Lung tissue was stored at -80°C.

Network Pharmacology

According to the Compounds identified by UHPLC-LTQ-Orbitrap, WYJ-related targets were obtained from SwissTargetPrediction database[14] (http://www.swisstargetprediction.ch/) and Traditional Chinese Medicine Systems Pharmacology Database and Analysis Platform (TCMSP, https://tcmsp-e.com/). The SDF format file of each Compound structure was obtained from PubChem database (https://pubchem.ncbi.nlm.nih.gov/) and loaded into the SwissTargetPrediction database to predict the targets. Compounds-related protein names obtained from TCMSP need converting to the gene names using UniProt (https://www.uniprot.org/). Combined with the results of the two databases, the targets of WYJ were obtained.

PF-related targets were obtained from GeneCards database v5.0[15] (https://www.genecards.org/, Relevance score ≥ median values) and OMIM database[16] (https://omim.org/) using "pulmonary fibrosis (PF)", "idiopathic pulmonary fibrosis (IPF)" or "interstitial lung disease (ILD)" as the keywords. After combining the targets obtained from different databases using the same keywords, the intersection of the targets from different keywords were the targets of PF.

WYJ-related targets and PF-related targets were intersected to obtain gene symbols. Using STRING database (https://www.string-db.org/)[17], the protein-protein interaction (PPI) network was constructed to explore the potential interactions of the intersected gene symbols. Cytoscape v3.6.0 visualized PPI network. Critical nodes calculated by median values of three topological features (degree, closeness centrality, and betweenness centrality) were identified as the key targets of WYJ.

To determine biological meaning behind key targets, pathway enrichment was analyzed by Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) databases in DAVID v6.8 (https://david.ncifcrf.gov/)[18, 19].The threshold value of confident gene enrichment was set at P≤0.05. Compound–target-pathway network explored the connections of compounds of WYJ, key targets, and important signaling pathways and was visualized using Cytoscape.

Molecular Docking

Based on results of LC/MS analysis and network pharmacology, we downloaded the 2D structures of ligands from PubChem database and the structures of target proteins from the PDB (https://www1.rcsb.org/). MGLTool-1.5.6 was used for generating necessary pdbqt file format of proteins and ligands along with the respective grids and dock files. AutoGrid 4.2 and Auto vina[20] were adopted to generate grids and execute docking respectively. 10 different conformations of each protein and compound were obtained and analyzed. PyMOLs visualization property[21] was used for the image construction of the docked ligand protein complexes. The binding affinities of the complexes were recorded in the log files of the docks generated by Auto vina.

Experimental Verification

Animals and experimental design

The body weight (BW) and the whole lung weight (PW) were recorded after mice were sacrificed. Pulmonary indexes (PI) were calculated using the equation PI = PW/BW*100.

Hydroxyproline determination (HYP)

The lung tissue was processed according to the instructions of the Hydroxyproline Kit (Nanjing Jiance Institute of Bioengineering, Nanjing, China). The level of hydroxyproline in the lung tissue was measured at the absorbance of 550nm and the results were expressed with µg/ lung.

Histopathological examination
Lung tissues fixed with 10% formalin were embedded in paraffin and sectioned. Sections were stained with hematoxylin-eosin (H&E) and Masson. Experienced pathologists performed sections pathological evaluation under light microscopy.

**Culture of human pulmonary fibroblast (HPF)**

HPF, a normal human fibroblast cell line from the National Infrastructure of Cell Line Resource (Beijing, China). HPF were cultured in medium (DMEM) with 10% FBS and 1% PS and maintained at the cell incubator with 5% CO2 and 37°C. Cells were passaged at a confluence of 80%-90% by standard trypsinization techniques. HPF stimulated with TGF-β1 (10ng/mL) were compared with cells that grown in the same conditions, but without stimulation of TGF-β1.

**Cell viability analysis**

The viability of HPF treated with WYJ was assessed using CCK-8 (CK001, Lablead). Firstly, HPF were seeded into 96-well plate in the density of 2×104 cells/well and treated with or without WYJ (0, 0.1, 1, 10, 20 or 50µg/mL) after 12-24h. Secondly, each well was added 200µL medium with 10% CCK-8 solution after 48h and then incubated at 37°C for 1-2h. Finally, the absorbance of well was measured at 490 nm using a microplate reader (Molecular Devices, Sunnyvale, CA, USA).

**Quantitative real-time polymerase chain reaction (qPT-PCR)**

Table 1: Gene Primer Sequences

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer(5’-3’)</th>
<th>Accession no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mus-GAPDH</td>
<td>Forward AGTTGTGCTCCCTGGAGCTTCA</td>
<td>NM_008084.3</td>
</tr>
<tr>
<td></td>
<td>Reverse TGGTCAGGGTTCTTACTCC</td>
<td></td>
</tr>
<tr>
<td>Mus-Col1a1</td>
<td>Forward CCAAGAAGACATCCCTGAGTCA</td>
<td>NM_007742.4</td>
</tr>
<tr>
<td></td>
<td>Reverse TGACGTCATGCACACA</td>
<td></td>
</tr>
<tr>
<td>Mus-TGF-β1</td>
<td>Forward ATGACATGAACCGGACCTTTT</td>
<td>NM_011577.2</td>
</tr>
<tr>
<td></td>
<td>Reverse AGTTGTATCCAGGGCTTCTCC</td>
<td></td>
</tr>
</tbody>
</table>

Proteins obtained from lung tissue or cells were further analyzed with Bicinchoninic Acid (Applygen, Beijing, China) for protein concentration. Protein at 30–50 µg/lane was loaded and separated by SDS-PAGE gel electrophoresis. After SDS-PAGE, proteins were transferred to the Polyvinylidene Fluoride (PVDF) membranes (Millipore, Massachusetts, USA). Membranes were blocked in 5% fat-free milk dissolved with Tris-buffered saline with Tween 20 (TBST) for 120 min at room temperature. Rinsed with TBST, these membranes incubated with primary antibodies at 4°C overnight. On the second day, the blots were rinsed with TBST and then incubated with secondary antibody for 1h. Blots were visualized by gel imager. GAPDH was used loading positive control and the results were analysed using image J.

**Antibodies:** Anti-α-SMA (ab5694, abcam, Cambridge, UK), Anti-Collagen 1 (14695, Proteintech), anti-p-MAPK (ab201015, abcam), anti-MAPK (ab184699, abcam), anti-p-STAT3 (ab76315, abcam), anti-STAT3 (ab68153, abcam), anti-GAPDH (60004, Proteintech, Chicago, USA), anti-rabbit IgG (ab205718, abcam) and anti-mouse IgG (ab6728, abcam).

**Data and statistical analysis**

One-way ANOVA were performed using GraphPad Prism software (version 8.0.2). P values < 0.05 are adopted as statistically significant.

**Results**

**Identified Ingredients of WYJ**

Using UHPLC-LTQ-Orbitrap to detect and identify chemical Compounds of WYJ (Fig. 1), a total of 23 compounds were confirmed based on comprising with the published data. Information about the identified ingredients was summarized in Tables 2 and 3.
Based on methods above, 436 WYJ-related targets were obtained from SwissTargetPrediction and TCMSP after removing duplicate items. Information about WYJ-related targets was provided in Supplementary Table S1. Combining the search results of the same keyword from GeneCards and OMIM database, a total of 2,867 targets for PF, 1,287 targets for IPF and 1,613 targets for ILD were collected. As shown in Fig. 2A, 846 overlapping targets were considered to be important targets about PF. The details were shown in Supplementary Table S2. Combining 463 WYJ-related targets with 846 PF-related targets, 124 targets could be the potential targets for the WYJ treatment on PF (Fig. 2B). Detailed information about 124 potential targets was provided in Supplementary Table S3.

STRING databases were used to explore the underlying interactions of 124 potential targets. There were 124 nodes and 1,716 edges in the PPI network visualized by Cytoscape (Fig. 3). According to the conditions of three topological (degree >22, betweenness centrality >0.00208, and closeness centrality > 0.5371179), 48 nodes were identified as the key targets of WYJ against PF (Supplementary Table S4).
The 48 key targets obtained from PPI were selected to investigate the biological processes and mechanisms of WYJ on PF treatment. Results indicated that biological processes involve in PF treatment include positive regulation of transcription from RNA polymerase II promoter, signal transduction, etc. Pathway enrichment analyses using the KEGG database revealed that most of the therapeutic targets are associated with signal transduction (e.g., MAPK signaling pathway and PI3K-protein kinase B (Akt) signaling pathway), inflammation (e.g., TNF signaling pathway and HIF-1 signaling pathway) and immune response (e.g., Toll-like receptor signaling pathway and Jak-STAT signaling pathway). The disease is caused by basic biological dysfunction, so the KEGG pathway about human disease were deleted. 20 significant BP terms and KEGG pathway terms were shown in Fig. 4.

Compound–target-pathway network explored the connections of important compounds of WYJ, key targets, and top 20 related pathways. The integrated compound–target-pathway network visualized by Cytoscape consisted of 69 nodes and 258 edges (Fig. 5). STAT3, SRC, IL6, MAPK1, AKT1, EGFR, MAPK8, MAPK14 and IL1B were the nine most important nodes in the compound–target-pathway network based on comparing betweenness centrality, closeness centrality and degree (Supplementary Table S5). These nodes also interacted closely with MAPK signaling pathway(MAPK8, IL1B, MAPK1, AKT1, PRKCA, MAPK14, RELA, EGFR)in the KEGG results. These results indicated that WYJ relieves PF mainly via the MAPK signaling pathway. Interestingly, we also found that STAT3 is the top node in the compound–target-pathway network and involved in multiple pathways. Above, these results demonstrated that WYJ worked in PF treatment through overall cooperation with multiple pathways and multiple targets.

Molecular Docking

In this paper, molecular docking using AutoDock Vina was applied to verify the reliability of the results of network pharmacology. The association between 13 active compounds in WYJ and target genes was visualized by compound–target-pathway network. Considering the results of KEGG and compound–target-pathway network. Therefore, we explored the interactions between 13 key compounds and nine important targets (STAT3, SRC, IL6, MAPK1, AKT1, EGFR, MAPK8, MAPK14, IL1B). Docking and binding affinity of ligands into the binding site of receptors are important information for prediction of pharmacology[22]. Results showed that most of the ligands have good binding activity with receptors (Vina scores less than −5 kcal/mol). The Vina scores were shown in Supplementary Table S6. The predicted target-compound with the highest affinity was visualized using PyMOL (Fig. 6).

Experimental verification of WYJ effects on the idiopathic pulmonary fibrosis

WYJ alleviated BLM-induced pulmonary fibrosis in mice

21 days after BLM injection, significant decreases in survival rate, severe weight loss and increase lung/body weight ratio were shown in BLM-induced PF model compared with the Control group. The WYJ treatment increased the survival rate, attenuated weight loss, and reduced the lung/body weight ratio (Fig. 7A, B, C). Hydroxyproline level, an important fibrotic indicator of collagen deposition, also decreased by the WYJ treatment (Fig. 7D). TGF-β1 is the important fibrotic factor in the development of pulmonary fibrosis[23, 24]. In our study, WYJ treatment down-regulated increased lung tissue TGF-β1 mRNA in PF mice model (Fig. 7E). BLM-induced PF mice model showed typical features of lung fibrosis, and WYJ prevented the development of lung fibrosis, as revealed by Masson staining (Fig. 7F) and Histopathological examination of lung sections (Fig. 7G). Above results confirmed that WYJ exerted a protective effect on lung fibrosis.

Excessive deposition of extracellular matrix (ECM), such as collagen[25], is the feature of fibrosis. The WYJ groups inhibited mRNA expression of Collagen-1 compared with the Model group (Fig. 8A). Fibroblasts lose their marker E-cadherin and transform to myofibroblasts characterized with α-SMA[24], occurs in the PF. As shown in Fig. 8B, bolts indicated the decreased α-SMA expression after the WYJ treatment compared with the Model group, indicating that the fibrosis process was inhibited by the WYJ treatment.

According to the network pharmacology analysis results, we assumed that MAPK signaling pathway could be the pharmacological mechanism of the therapeutic effect of WYJ against PF. Phosphorylation levels of MAPK1 (p-MAPK1) was increased in the Model group compared with the Control group (Fig. 8C). The WYJ treatment reduced the upregulation of p-MAPK1 expression in BLM-induced PF model.

Network pharmacology analysis showed that STAT3 is the top node in the compound–target-pathway network and involved in multiple pathways. Studies showed STAT3 is the common downstream target of fibrotic factors. Levels of phosphorylated STAT3 (p-STAT3) was elevated in the Model group. We detected that the WYJ treatment remarkably reduced the increased p-STAT3 (Fig. 8D). Pulmonary fibrosis is the result of multiple cytokines or signal transduction pathways. WYJ reduced pulmonary fibrosis by down-regulating p-STAT3, which may be a new perspective to explain anti-lung fibrosis of WYJ.

WYJ inhibited fibroblast activation in vitro experiments

In vitro experiments further verified the anti-lung fibrosis effect of WYJ. CCK-8 assay was used to determine the effect of WYJ on HPF activity. As shown in Fig. 9A, WYJ shows no effect on the viability of normal cells at 1–20 ng/ml. Activated lung fibroblasts, the primary roles in the progress of fibrogenesis, express excessive α-SMA and Collagen-1. To determine whether WYJ inhibits fibroblast activation, we used TGF-β1 to promote fibroblast activation[24] and evaluated the expression of α-SMA and collagen-1. The results demonstrated that HPF stimulated with TGF-β1 significantly increased the production of α-SMA (Fig. 9B) and Collagen-1 (Fig. 9C) protein. HPF pretreated with WYJ markedly inhibited expression of α-SMA and
Collagen-1 protein. And results also showed that WYJ prevented the activation of fibroblasts by down-regulating p-MAPK and p-STAT3 in vitro (Fig. 9D, E).

Discussion

TCMs has accumulated valuable information alleviating lung diseases. WYJ, from steamed roots of Curcuma wenyujin, was included in the 2020 edition of the Pharmacopoeia of People’s Republic of China. Modern pharmacological studies have demonstrated that the ingredients or extracts from WYJ have antibacterial, antitumor anti-inflammatory and antioxidant effects. There are studies reported that curcumin, an essential ingredient of WYJ, have anti-pulmonary fibrosis in murine model.[27, 28]. Hui’s study showed potential anti-fibrotic effects of WYJ in liver fibrosis.[29] In our present study, we investigated the effect of WYJ against PF and the related molecular mechanism.

In this study, we established PF mice model by administering (via the trachea) bleomycin at a dose of 5 mg/kg. This model could cause a pulmonary pathological change similar to that of patients with lung fibrosis.[30, 31] We explored the effect of WYJ on PF from the perspectives of the survival rate, HYP, and histopathological examination. In our study, three dosage groups of WYJ were evaluated. The results showed that all dosage groups hindered lung fibrosis, and the WYJ-H and WYJ-M groups demonstrated the best therapeutic effect.

We identified 23 major compounds from WYJ using UHPLC-LTQ-Orbitrap. Further the network pharmacology results revealed that 128 targets could be potential targets on the treatment of PF, and PPI network showed 48 of them were key targets. KEGG analysis results and the compound-target-pathway network suggested that STAT3, SRC, IL6, MAPK1, AKT1, EGFR, MAPK8, MAPK14 and IL1B are important targets. MAPK8, MAPK1, AKT1, MAPK14, EGFR and IL1B were the important targets in MAPK signaling pathway. STAT3 was the top node in the compound-target-pathway network and involved in multiple pathways. The prediction results indicated that WYJ relieved pulmonary fibrosis mainly through inhibiting the MAPK signaling pathway and down-regulating the expression of p-STAT3.

MAPKs signaling cascades participate in the regulation of processes including cell cycle progression, cell migration, cell survival and differentiation.[32, 33]. Katerina M and colleagues[34] found BRAF major signaling intermediate protein that regulates the MAPK/ERK pathway increased in lung tissue of patients with IPF compared with normal. Another study found activated MAPKs are significantly increased in lung parenchyma of patients with IPF compared that of normal[35]. Bach1 ameliorated fibrosis and inflammation via blocking MAPK pathway in pulmonary fibrosis in recent study.[36]. Another study proved that the blockade of ERK by PD98059 attenuated the increase of collagen mRNA expression induced by TGF-β1.[37]. In our study, the WYJ treatment remarkably reduced the increased α-SMA, Collagen-1, and levels of p-MAPK1 in BLM-induced PF model and TGF-β1 induced HPF cells. The inhibitory effect of WYJ on PF may function mainly through modulating the MAPK signaling pathway.

Compound–target-pathway network revealed that STAT3 is the top node in PF treatment with WYJ. STAT3 regulates cell growth, proliferation, differentiation, and migration.[38]. Important fibrotic factors including TGF-β1, PDGF, FGF-2, and IL-13 can activate STAT3[39–43] and further lead to pulmonary fibrosis. Levels of phosphorylated STAT3 have been shown to be elevated in patients with IPF and bleomycin model.[44–47]. C-188-9, a synthetic small molecule inhibitor of STAT3 phosphorylation, inhibited STAT3 activation and decreased the development of pulmonary fibrosis in BLM-induced PF model. Accordingly, targeting common downstream mediators of fibrogenic mediators, STAT3, PF can be treated by several of the cellular pathways. In our study, elevated levels of p-STAT3 were observed in BLM-induced PF model and TGF-β1 induced HPF cells. The WYJ treatment remarkably reduced the increased p-STAT3. WYJ relieves pulmonary fibrosis by down-regulating the expression of P-STAT3.

We also explored the effect of WYJ on PF from another perspective using molecular docking. The results proved that compounds of WYJ have good binding activity with key targets (MAPK8, MAPK1, AKT1, MAPK14, EGFR, IL1B and STAT3), further illustrating that WYJ could hinder lung fibrosis via inhibiting the MAPK signaling pathway and down-regulating the expression of p-STAT3.

This study has some drawbacks. Compounds which have the most effective therapeutic effects require further definite and research. The network pharmacology had limitations. A total of 23 Compounds of WYJ were confirmed based on UHPLC-LTQ-Orbitrap. However, only targets of 15 Compounds were obtained from databases and the network pharmacology only involved targets of 13 Compounds. Whether the remaining Compounds play an anti-pulmonary fibrosis effect needs further research. The pharmacological mechanism by which WYJ treated PF need further exploration.

Conclusions

In conclusion, our study identified that WYJ alleviated pulmonary fibrosis (Fig. 10). According to the results of network pharmacology and molecular docking, the pharmacological mechanism by which WYJ-treated PF was further validated by experiment. The inhibitory effect of WYJ on PF may function mainly through modulating the MAPK signaling pathway. Interestingly, WYJ reduced pulmonary fibrosis by down-regulating p-STAT3, which may provide new insights for the treatment of WYJ on pulmonary fibrosis. The method of exploring the pharmacological mechanism of WYJ in our study provides a novel approach to explain the pharmacological basis of other herbs. The limitations of network pharmacology indicate that more experiments are needed to verify the reliability anti-fibrotic effect of WYJ.

Abbreviations
Declarations

Ethics approval and consent to participate

Animal procedures were performed according to the protocols of Animal Care and Use Committee of Beijing University of Chinese Medicine.

Consent to publish

All authors have provided consent for publication in the Journal of Chinese Medicine.

Availability of data and materials

The datasets used and analyzed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.

Funding

Not applicable.

Authors' Contributions

LW and WX-Z made substantial contributions to the conception and design of the study, performed the experiments and wrote the paper. RS, BB-Z and QH-M assisted in the performance of experiments. JL participated in data analysis. YY-S modified the experimental protocol, revised the manuscript, and finally approved the version to be published. All authors read and approved the final manuscript.

Acknowledgements

Not applicable.

References


Figures
Figure 1

Total ion chromatography

Figure 2
(A) Venn diagram of IPF-related targets. (B) Venn diagram of potential targets for WYJ treatment on PF.

Figure 3
PPI network of potential targets.

Figure 4
Bubble diagram of functional analysis. (A) Enrichment analysis of biological process (B) KEGG pathway enrichment analysis.
Figure 5

The integrated compound–target-pathway network visualized by Cytoscape.

Figure 6

The binding modes of ligand and receptor protein. (A) Schematic diagram of the interplay between 1,2-Dihydrocurcumin and EGFR (PDB ID chimeric 5YU9). (B) Schematic diagram of the interplay between ar-turmerone and STAT3 (PDB ID chimeric 6TLC). (C) Schematic diagram of the interplay between bisdemethoxycurcumin and SRC (PDB ID chimeric 2BDF). (D) Schematic diagram of the interplay between Curcumolactone A and MAPK14 (PDB ID chimeric 1SFO). (E) Schematic diagram of the interplay between turmeronol A and MAPK1 (PDB ID chimeric 1TVO). (F) Schematic diagram of the interplay between turmeronol A and MAPK8 (PDB ID chimeric 3ELJ).

Figure 7

WYJ alleviated BLM-induced pulmonary fibrosis in mice. (A) Survival rate. (B) The body weight. (C) Lung/Body weight ratio. (D) Hydroxyproline. (E) Lung tissue TGF-β1 mRNA. (F) The representative images of Masson staining. (G) The representative images of H&E staining. C: The Control group. M: The Model group. The data were presented as mean ± SEM (n≥3). *P< 0.05, **P< 0.01 and ***P< 0.001 versus the Model group; #P< 0.05, ##P< 0.01 and ###P< 0.001 versus the control group.

Figure 8

The effect of WYJ on BLM-induced pulmonary fibrosis mice. (A) mRNA expressions of Collagen-1 in lung tissues were measured by qRT-PCR. (B–D) The relative protein levels of α-SMA, p-MAPK1 and p-STAT3. C: The Control group. M: The Model group. The data were presented as mean ± SEM (n≥3). *P< 0.05, **P< 0.01 and ***P< 0.001 versus the Model group; #P< 0.05, ##P< 0.01 and ###P< 0.001 versus the control group.

Figure 9

(A) The effect of WYJ on HPF activity. (B–E) The relative protein levels of α-SMA, Collagen-1, p-MAPK1 and p-STAT3. C: The Control group. M: The Model group. The data were presented as mean ± SEM (n≥3). *P< 0.05, **P< 0.01 and ***P< 0.001 versus the Model group; #P< 0.05, ##P< 0.01 and ###P< 0.001 versus the control group.
Figure 10

The underlying mechanisms involved in the anti-lung fibrotic activity of WYJ.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- SupplementaryTable.xlsx