Investigating the Mechanism of Guizhi Fuling Formula in the Treatment of Primary Dysmenorrhea based on Network Pharmacology and Molecular Docking

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Research

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Abstract

Background: Primary dysmenorrhea (PD) is the most common gynecologic disorder. Despite the prevalence is high, it is often underdiagnosed, undertreated and normalized even by patients themselves. Guizhi Fuling Formula (GFF) is experientially used for the treatment of PD in a long time. Therefore, the efficiency and potential mechanism are waiting to identify.

Methods: We adopted network pharmacology integrated molecular docking strategy in this study. Based on published literatures, the relative compounds of GFF were selected preliminarily. Secondly, the putative targets of PD were obtained by wide-searching DisGeNET, OMIM, Drugbank and GeneCards databases. With protein-protein interaction (PPI) analysis, GO and KEGG pathway enrichment analysis and molecular docking, we systematically evaluated the relationship of herb ingredients and disease targets.

Results: The results showed that 30 ingredients of GFF and 43 hub targets made a difference. Under the further analysis, 8 targets (EGFR, AKT1, PTGS2, TNF, ESR1, AHR, CTNNB1, CXCL8) were recognized as key therapeutic targets with excellent binding. The enrichment analyses indicated that the GFF had the potential to influence varieties of biological pathways, especially the pathways in cancer and steroid hormone biosynthesis, which play an important part in the pathogenesis of primary dysmenorrhea.

Conclusion: GFF influenced primary dysmenorrhea through the synergistic effect of multiple components, multiple targets, and multiple pathways. This study predicted the potential mechanism, hope that could make contribution for clinical application and scientific research.

Introduction

Primary dysmenorrhea (PD) is the most common gynecological disorder, whose definition was epitomized a painful menstruation without the pelvic pathology[1, 2].

There are many studies have demonstrated that PD affect women's academic performance or work management, even the quality of life (QoL), this phenomenon not only bring troubles to people but also have an impact on social and economy[3, 4].

The latest PD guideline systematically reviewed the intervention of primary dysmenorrhea. Medical therapy is first-recommended, then the complementary and alternative therapy is also recommended, surgical management should be considered only the above-mentioned therapies has not been satisfactory[5]. As the first-line drug, NSAIDs were found to be consistently more effective than placebo, although adverse effects were significantly more common. Besides, some RCTs confirmed that cyclic oral contraceptives (COC) was superior to placebo in the treatment of primary dysmenorrhea[6–7]. Recently, Yutaka O.et al. verified the favorable tolerability and efficiency of the long-term use of Dienogest.
for patients with PD, which as a progestin and probably caused irregular uterine bleeding and associated anemia[8].

Compared with those interventions, Traditional Chinese medicine may be able to offer more cost-effective and safe alternative treatments against PD. Guizhi Fuling Formula was initially seen in Synopsis of Golden Chamber(Fig. 1), which is mainly composed of Cinnamomi Ramulus(Guizhi), Poria(Fuling), Moutan Cortex(Danpi), Persicae Semen(Taoren), Peony(Baishao). In the treatment of many gynecological diseases such as hysteromyoma, dysmenorrhea, ovarian cyst etc, it showed ideal curative effect. Evidences on literatures and animal-model trials verified the efficiency of GFF treating PD[9–10].

However, researches about systematic network analysis are deficient comparatively and contribution of the special compatibility of these five herbs on ameliorating PD is not corroborated. Thus, we conducted network pharmacology integrated molecular docking analysis based on the interaction of drugs, diseases, genes, proteins and targets in the biological network to investigate the bioactive constituents and potential mechanism of PD therapy.

**Materials And Methods**

**Screening of bioactive compounds of GFF**


**Prediction of putative targets of GFF**

We utilized the SwissTargetPrediction [15] (http://www.swisstargetprediction.ch/) to predict potential targets of GFF compounds. Besides, the UniProt [16] (http://www.uniprot.org/) was applied to convert the protein name of the GFF bioactive compounds to the gene names. We filtered items by "Reviewed" and "Human", then used the result to standardize targets and deleted data duplicates.

**Identification of PD related targets**

With a keyword searching of “Primary Dysmenorrhea” among four database: (1) the GeneCards database [17] (https://www.genecards.org/); (2) the Online Mendelian Inheritance in Man database [18] (OMIM, http://omim.org/); (3) the DisGeNET database[19](http://www.tdisgenet.org/search), (4) the DrugBank
database [20] (https://www.drugbank.ca/), we gathered related PD genes and made intersection with bioactive compound genes for further analysis.

**Protein-protein interaction (PPI) network construction**

Most of the proteins appear to activate their functionalities through their interaction. By evolving core regions with surface residues and hotspot residues at the interface, it could modulate cellular processes such as signal transduction, which provided evidences for drug designing or discovering. [21-22] Thus, we submitted the PD targets and the intersection separately to STRING 11.0 database [23] (https://string-db.org/) for PPI analysis, which set filter as “Homo sapiens” organism, to identify important targets in anti-PD system.

**Network construction**

To characterize the mechanisms of GFF in treating PD from a network target perspective, we employed the Cytoscape 3.8.2 [24] (https://cytoscape.org/) to construct visualization networks as follows: (1) Compound-target network; (2) PPI network of PD targets; (3) PPI network of compound PD targets; (4) Compound-target-pathway network. When analyzed the topological properties of the PPI network by the Cytoscape, there were three indices recommended to assess nodes and edges. Firstly and commonly, we used “Degree” to evaluate the interaction between nodes [25]. Secondly, “Betweenness Centrality” was applied to determine the amount of information passing between nodes [26]. Thirdly, “Closeness Centrality” reflects the closeness of a node to other nodes [27]. The higher values of this three parameters are, the more important the nodes are. Thus, the hub genes for molecular docking were selected systematically with the help of above parameters.

**Functional enrichment analysis**

As same as the choosing principle of bioactive compounds, Functional Annotation tool of Database for Annotation, Visualization and Integrated Discovery (DAVID) 6.8 [28] (https://david.ncifcrf.gov/) was not the best candidate, whose speed of updating is too sluggish. What’s more, Metascape [29] (https://metascape.org/) are recommended to conducted GO functional analysis and KEGG pathways analysis recently. We set parameters as follow: Overlap>2, P value>0.01 and Enrichment>1.5.

**Molecular docking simulation**

This study selected the " component target-pathway " network moderate value (degree) top 8 target protein, and these eight potential targets with nine corresponding compounds were simulated by molecular docking. Initially, downloading the PDB format of the 3D structure of the target protein from the Protein Data Bank database [30] (https://www.rcsb.org/). The screening conditions were set as follows: (1) the organism comes from Homo sapiens; (2) the protein structure is obtained by X-crystal diffraction; (3) the crystal resolution of the protein is less than 2 Å; (4) preferential selection of protein structures reported in the literature of molecular docking. And the mol2 format of the 3D structure of the compounds...
were downloaded in the ZINC database[31] (https://zinc.docking.org/). Secondly, the Pymol 2.4[32] was used to remove water molecules and raw ligands. After, utilizing AutoDockTools1.5.6[33] (http://mgltools.scripps.edu/documentation/links/autodock) to hydrogenate and charge, and then the pdb format of compounds and core protein were converted into pdbqt format and active pocket was found. Eventually, Running AutoDock Vina1.1.2[34] with CMD command characters for molecular docking and calculating binding energy. Visualization of results using PyMOL 2.4 software.

Results

Compound-target network

Based on the Pharmacokinetics and Druglikeness parameters in SwissADME[14], we selected a total of 30 candidate bioactive compounds. (Complementary Table 1). Then, every compound was submitted on Swiss Prediction for identifying related targets. There were 374 compound-predicted targets of GFF made up the network, which consist of 409 nodes and 907 edges (Fig. 3). We found that a single target can be co-regulated by a variety of compounds to trigger the biological effects, which may play a vital role in treating PD.

PPI network of PD targets

Integrated targets which retrieved from GeneCards, DisGeNet, OMIM and DrugBank database, then we uploaded them to the STRING database and constructed a PPI network with 319 nodes and 4270 edges. For better visualization, the Cytoscape was applied for network mapping. Based on analyzing network, we used degree (value > 2) cutting low-related targets. From yellow to blue, the degree was increasing. (Fig. 4) The bigger the node was, the higher the degree was.

PPI network of compound PD targets

We made a intersection between GFF bioactive targets and PD related targets, 43 targets were identified as the potential targets for treating PD (Fig. 5a). Similarly, we conducted a PPI network to explore the mechanism of the drugs in disease therapy and discover the interactive effects. Then employed Cytoscape to analyze the topological feature of the PPI network and cut low-related targets, 39 genes arranged in a concentric circle according to the degree. (Fig. 5b) According to average degree (value > 17), Betweenness Centrality (>0.03) and Closeness Centrality (>0.06), we selected a total of 8 hub genes (Table 1), including EGFR, AKT1, PTGS2, TNF, ESR1, AHR, CTNNB1, CXCL8. These hub genes are of significance in the network, which would be more suitable for further molecular docking analysis.

Table 1. Topological information of 8 hub genes
GO and KEGG enrichment analysis

The GO items and KEGG pathways are selected based on counts of hit genes and P values (Fig. 6). For biological processes, it can be found that the targets were mainly enriched in cellular response to organic cyclic compound (GO:0071407) or nitrogen compound (GO:1901699), steroid metabolic process (GO:0008202), membrane raft (GO:0045121) and oxidoreductase activity (GO:0016705).

For pathway analysis, most involved in pathways in cancer (hsa05200), proteoglycans in cancer (ko05205), human cytomegalovirus infection (hsa05163) and steroid hormone biosynthesis (ko00140).

Molecular docking analysis

As far as current research is concerned, we choose 8 hub genes and 9 core bioactive compounds (Table 2) to test the potential interaction activity by molecular docking verification. A total of 72 pairs were used for molecular docking simulation. The smaller the value of docking affinity, the more easily the active ingredient binds to the receptor. According to the results of docking, the great majority of binding complexes possessed high binding affinity with an average of -7.5 kcal/mol. Molecular docking affinity values <-7 accounted for 69.44%, indicating better binding activity. The first eight binding complexes with affinity less than -9 were selected and displayed in Table 3, including EGFR-ellagic acid docking (-9.4 kcal/mol), AHR-taxifolin docking (-10.3 kcal/mol), AHR-(+)-catechin docking (-10.2 kcal/mol), AHR-baicalin (-10.2 kcal/mol), CTNNB1-Stigmasterol docking (-10.3 kcal/mol), CTNNB1-ellagic acid docking (-9.9 kcal/mol), CTNNB1-baicalin docking (-9.4 kcal/mol), CTNNB1-(+)-catechin docking (-9.3 kcal/mol).

Table 2. Topological information of 9 core bioactive compounds

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Protein name</th>
<th>BetweennessCentrality</th>
<th>ClosenessCentrality</th>
<th>Degree</th>
</tr>
</thead>
<tbody>
<tr>
<td>EGFR</td>
<td>Epidermal growth factor receptor</td>
<td>0.078120983</td>
<td>0.701754386</td>
<td>23</td>
</tr>
<tr>
<td>APT1</td>
<td>Rac-alpha serine/threonine-protein kinase</td>
<td>0.077030349</td>
<td>0.701754386</td>
<td>23</td>
</tr>
<tr>
<td>PTGS2</td>
<td>Prostaglandin G/H synthase 2</td>
<td>0.076204982</td>
<td>0.68965172</td>
<td>23</td>
</tr>
<tr>
<td>TNF</td>
<td>Tumor necrosis factor</td>
<td>0.10958786</td>
<td>0.68965172</td>
<td>23</td>
</tr>
<tr>
<td>ESR1</td>
<td>Estrogen receptor</td>
<td>0.062011048</td>
<td>0.67766102</td>
<td>23</td>
</tr>
<tr>
<td>AHR</td>
<td>Aryl hydrocarbon receptor</td>
<td>0.042161256</td>
<td>0.644516129</td>
<td>19</td>
</tr>
<tr>
<td>CTNNB1</td>
<td>Catenin beta-1</td>
<td>0.033031049</td>
<td>0.625</td>
<td>18</td>
</tr>
<tr>
<td>CXCR3</td>
<td>Interleukin-8</td>
<td>0.032675776</td>
<td>0.63492635</td>
<td>18</td>
</tr>
</tbody>
</table>
Table 3. Analysis of molecular docking results

<table>
<thead>
<tr>
<th>Molecule name</th>
<th>MOLID</th>
<th>BetweennessCentrality</th>
<th>ClosenessCentrality</th>
<th>Degree</th>
</tr>
</thead>
<tbody>
<tr>
<td>quercetin</td>
<td>MOL000098</td>
<td>0.660565978</td>
<td>0.57208238</td>
<td>145</td>
</tr>
<tr>
<td>kaempferol</td>
<td>MOL000422</td>
<td>0.14134038</td>
<td>0.411861614</td>
<td>56</td>
</tr>
<tr>
<td>beta-sitosterol</td>
<td>MOL000358</td>
<td>0.111475745</td>
<td>0.390035601</td>
<td>38</td>
</tr>
<tr>
<td>baicalein</td>
<td>MOL002714</td>
<td>0.096768005</td>
<td>0.374812594</td>
<td>34</td>
</tr>
<tr>
<td>Stigmasterol</td>
<td>MOL000449</td>
<td>0.090225666</td>
<td>0.382848399</td>
<td>30</td>
</tr>
<tr>
<td>hederagenin</td>
<td>MOL000296</td>
<td>0.055000783</td>
<td>0.372578241</td>
<td>24</td>
</tr>
<tr>
<td>ellagic acid</td>
<td>MOL01002</td>
<td>0.031733698</td>
<td>0.354609929</td>
<td>18</td>
</tr>
<tr>
<td>taxifolin</td>
<td>MOL004576</td>
<td>0.026248081</td>
<td>0.341997264</td>
<td>11</td>
</tr>
<tr>
<td>(+)-catechin</td>
<td>MOL000492</td>
<td>0.016160162</td>
<td>0.353606789</td>
<td>11</td>
</tr>
</tbody>
</table>

For further explanation, Give an example of AHR-taxifolin docking, small molecule ligand taxifolin could possibly fit into the active interface pocket formed by interaction amino acid residues in protein (Fig.7). It displayed that a hydrogen bond formation between ligand and residues involved in TRP-
Consequently, taxifolin stably bound to AHR protein bonds through interaction forces. On account of this experiment, the results of molecular docking showed that the core compounds and key targets screened by network pharmacology had fairly binding activity.

**Compound-target-pathway network**

For further exploring the interaction among pathway, gene and compound, Cytoscape was applied again to construct a compound-target-pathway network (Fig. 8a).

Based on KEGG enrichment analysis, we obtained 14 pathways which related different targets. Thanks to compound-target network above, the new network was responsibly easy to conducted. As shown in Fig. 8a, there were a total of 72 nodes and 187 edges. 14 pathways were interacted with 31 targets and 27 compounds. Subsequently, degree was adopted as the threshold values to identify the significant compounds, targets and pathways. Six compounds, 20 targets and 8 pathways were selected for a more detailed explanation of the associations described above, which displayed in the sub-network (Fig. 8b).

**Discussion**

From past to now, the difference between Traditional Chinese Medicine (TCM) and Modern Medicine is the concept of whole and part. The principle of choosing Chinese herbs was based on four properties, five flavors and channel tropism [35]. Appropriate compatibility made prognosis of disease better, it was hard to explain concretely the interplay among those herbs for people who never learned about TCM. Thus, we need a evidence-based and illustrative methodology to explore the mechanism.

According to a series of analysis above, we gained six significant compounds, keampferol concentrated in Baishao (Peony), quercetin and isorhamnetin were enriched in Danpi (Moutan Cortex), cerevisterol and 3,5-Dihydroxyergosta-7,22-dien-80-one were found in Fuling (Poria), glycosmisisic acid existed in Guizhi (Cinnamomi Ramulus).

Keampferol and quercetin always appeared at the same time due to the familiar effectiveness in anti-inflammatory, antioxidation and proapoptosis [36-39]. In a network study of TCM treating cancer-related pain [40], quercetin and kaempferol also worked as very important compounds, which could regulate the phosphoinositide 3-kinase (PI3K)/protein kinase B (AKT) pathways to achieve the purpose of antitumor and alleviating pain. It might provide evidences for the effect of quercetin and kaempferol ameliorating the pain of menstruation. Cerevisterol and 3,5-Dihydroxyergosta-7,22-dien-80-one were usually isolated and identified from the mycelia, which could treat inflammatory diseases by targeting an MAPK, NF-κB, AP-1, and Nrf2-mediated H0-1 signaling cascade [41].

Epidermal growth factor receptor (EGFR) belongs to a family of four homologous tyrosine kinase receptors (TKRs). This family are one of the major drivers of cancer and is involved in several of the most frequent malignancies such as non-small cell lung cancer, breast cancer, colorectal cancer and ovarian cancer [42]. Although EGFR related to many diseases, there were not published trials confirmed the association...
between EGFR and PD. RAC-alpha serine/threonine-protein kinase AKT1 had a close relationship with gynecologic disease, such as endometriosis [43] and endometrial carcinoma [44]. In our study, it presented a significant relevance to PD, which could provide evidences for further studies. Prostaglandin G/H synthase 2 PTGS2 is an inducible prostaglandin enzyme whose derivative (PGE2 and PGI2) are the major mediators in decidualization [45]. Immunohistochemical tests, western blotting (WB) assays and polymerase chain reaction (PCR) show that PTGS2 expression in the uterine myometrium and glands is directly related to organ inflammation intensity [46]. One pathological mechanism of PD is the increase of PGE2 and PGF2α [47], verifying the great effect of PTGS2 on PD.

The results of pathway analysis suggested that Pathways in cancer, Proteoglycans in cancer, Human cytomegalovirus infection and Steroid hormone biosynthesis showed significant association with PD. It is worth thinking about its close relationship with cancer-related pathways.

Steroid hormone biosynthesis (Complementary Fig. 1) contributed to regulate metabolism and stress response, initiate and maintain sexual differentiation and reproduction [48]. Follicle-stimulating hormone (FSH) controls the progesterone synthesis from granulosa cells via G protein/adenylate cyclase (cAMP)/protein kinase A (PKA)/MAPK pathway, then progesterone transforms to estrogen under the regulation of enzymes in the steroid hormone metabolism pathway [49]. Thus, it indirectly confirmed the effectiveness of drugs or herbs therapy via hormone pathway. Ji et al. revealed the enhanced steroid hormone biosynthesis during decidualization, which was influenced by metabolic alteration [50]. Serotonergic synapse is widely associated with mood disorders, especially in women [51]. Thus, PD caused by stress, depression or other mood disorders could treat in this train of thoughts.

Studies about the Adipocytokine signaling pathway and PPAR signaling pathway are limited. Hu et al. corroborated that PPAR signaling pathway participated in the regulation of inflammatory response [52]. Dotun et al. indicated that the downregulation of several key genes in the KEGG Adipocytokine Pathway was linked to insulin resistance or inflammation [53].

Overall, it was speculated that the main compositions of GFF may play a significant role in the treatment of PD through hub targets in these top-ranking signaling pathways. However, some limitations of our study should be considered. For instance, the results are only based on screening already known chemical constituents of GFF, related targets, and signaling pathways from literatures and existing databases. Consequently, more in-depth researches are required for characterization of the underlying mechanisms.

**Conclusion**

In the present study, there were four networks that we derived to reveal the significant compounds, targets and pathways of GFF in PD treatment, including compound-target network, PPI network of PD targets, PPI
network of compound-PD targets and compound-target-pathway network. After sieving and consolidating data from available databases, we got 30 bioactive compounds of GFF and 43 potential targets for treating PD. According to the PPI network, 8 hub genes were highly connected and could be defined as vital targets in the process of treating PD with GFF. In GO and KEGG pathway enrichment analysis, we found that these hub targets ameliorated PD by participating in signal transduction, hormone-metabolism system and inflammatory response. In short, the above results preliminarily predicted the potential mechanism of GFF against PD, visualizing the characteristics of Chinese herbs interaction, which providing more possibility of PD therapy. Actually, further extensive researches such as animal experiments and clinical investigations will be needed, which is necessary to provide more valuable evidences for the application of GFF in treating PD.

Declarations

Availability of data and materials

All data in the manuscript are available and they are exhibited in figures and tables.

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Contributions

SL, WZN and XL: study design; acquisition of data; analysis of data; drafting of the manuscript. LJ and XXO: revision of the manuscript and study supervision. All the author(s) read and approved the final manuscript.

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Correspondence to Xiao-ou Xue and Lu Sun.
Ethics declarations

Ethics approval and consent to participate

No applicable.

Competing interests

Authors declare no conflict of interest.

References


Figures

Figure 1

Synopsis of Golden Chamber and Guizhi Fuling Formula
Figure 2

Flowchart of investigating the mechanism of GFF in PD treatment

Figure 3
Compound-target network. Colourful octagen nodes represent bioactive compounds of GFF and blue diamond nodes represent corresponding targets.

**Figure 4**

PPI network of PD targets. The sizes and colors of the nodes are illustrated from big to small and blue to yellow in descending order of degree values.

(a) [Diagram showing Venn diagram of intersecting targets of GFF and PD]

(b) [Diagram of PPI network of compound-PD targets]

**Figure 5**

Venn diagram and PPI network of compound-PD targets. a. Venn diagram of intersecting targets of GFF and PD. b. PPI network of compound-PD targets. The sizes and colors of the nodes are illustrated from big to small and blue to yellow in descending order of degree values.
Figure 6

GO & KEGG enrichment analysis. a GO enrichment analysis. BP: biological processes, CC: cell component, MF: molecular function. b KEGG pathway analysis. The sizes of the bubbles are illustrated from big to small in descending order of the number of the potential targets involved in the pathways.
Figure 7

Analysis of target-compound docking simulation

**Supplementary Files**

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

- CTable11.png
- CTable12.png
- CTable13.png
- CTable14.png
- C.Fig.1.png