A Network Pharmacology Approach to Explore the Pharmacological Mechanism of Xiaoyao Powder on Anovulatory Infertility

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Aim. To explore the pharmacological mechanism of Xiaoya powder (XYP) on anovulatory infertility by a network pharmacology approach. Method. Collect XYP’s active compounds by traditional Chinese medicine (TCM) databases, and input them into PharmMapper to get their targets. Then note these targets by Kyoto Encyclopedia of Genes and Genomes (KEGG) and filter out targets that can be noted by human signal pathway. Get the information of modern pharmacology of active compounds and recipe’s traditional effects through databases. Acquire infertility targets by Therapeutic Target Database (TTD). Collect the interactions of all the targets and other human proteins via String and INACT. Put all the targets into the Database for Annotation, Visualization, and Integrated Discovery (DAVID) to do GO enrichment analysis. Finally, draw the network by Cytoscape by the information above. Result. Six network pictures and two GO enrichment analysis pictures are visualized. Conclusion. According to this network pharmacology approach some signal pathway of XYP acting on infertility are found for the first time. Some biological processes can also be identified as XYP’s effects on anovulatory infertility. We believe that evaluating the efficacy of TCM recipes and uncovering the pharmacological mechanism on a systematic level will be a significant method for future studies.

1. Introduction

Abnormal ovulation is the most common cause of female infertility, whose mechanism is unable to produce fertilizable oocytes. The most common one is depletion of the oocyte pool, which manifests itself as anovulation, less ovulation, and follicle aging. Anovulation is related to amenorrhea and severe oligomenorrhea; less ovulation is connected with oligomenorrhea (Menstrual Cycle longer than 35 days) [1, 2]. The most common causes of anovulation in adult women are hypothalamic dysfunction (35%), pituitary disease (15%), and ovarian dysfunction (50%) [3, 4]. In the ovary, it is ovarian failure (depletion of the oocyte pool) and ovarian hyperandrogenism (e.g., polycystic ovary syndrome, PCOS) [5–7]. Though its etiology is still not clear, it is the result of the interaction between the genetic and environmental factors(s). Its treatment is mainly focused on promoting follicular maturation. The pharmacological options include clomiphene citrate, clomiphene with hormones, gonadotropin (Gn), gonadotropin-releasing hormone (GnRH), bromocriptine, and glucocorticoid [8]. This can promote ovulation and relieve endocrine disorders related symptoms. However, long-term application will lead to ovarian tumor or other reproductive tumor and ovarian hyperstimulation syndrome [2].

As an important part of the complementary and alternative medical system, traditional Chinese medicine (TCM) has been widely utilized in the treating of infertility for centuries and has been proven efficient in regulating endocrine and promoting ovulation. In the concept of TCM, anovulatory infertility can be classified as “amenorrhea,” “Zhengjia,” “sterility,” “metrorrhagia and metrostaxis,” “depleted blood,” and so on [9]. However, application of TCM has been blocked by the absence of scientific comprehension regarding its mechanism. Therefore, it is important to explore and reveal the TCM mechanism.

The use of Xiaoya powder (XYP) was first recorded in Taiping Huimin Heji Ju Fang, which is regarded as the first monograph about TCM. This formula is for female
menstruation and is composed of *Radix Bupleuri* (Chai Hu), *Angelicae Sinensis Radix* (Dang Gui), *Poria Cocos* (Schw.) *Wolf.* (Fu Ling), *Paeoniae Radix Alba* (Bai Shao), *Atractylodes Macrocephala Koidz.* (Bai Zhu), and *Licorice* (Gan Cao). Based on TCM theory, multiple herbs in one formula should operate cooperatively. In XYP, *Radix Bupleuri* is the main force of dispersing stagnated liver Qi for relieving Qi stagnation; *Angelicae Sinensis Radix* and *Paeoniae Radix Alba* are used for nourishing blood; *Atractylodes Macrocephala Koidz.* and *Poria Cocos* (Schw.) *Wolf.* are able to invigorate the spleen; *Licorice* reconciles the various drugs. These herbs synergistically treat anovulatory infertility through dispersing stagnated liver Qi for relieving Qi stagnation, nourishing blood, and invigorating the spleen. The latest study shows that XYP has a therapeutic effect on ovarian failure and hyperandrogenism [10, 11]. Thus, XYP might be a novel therapeutic strategy for anovulatory infertility. However, its pharmacological mechanism has not been clarified completely.

Chinese herbal formulae are multitarget and multicomponent recipes that achieve their particular therapeutic efficacy through regulation of the molecular network of body systems utilizing its active components [12]. Therefore, new methods and new tactics are required to explore and explain the mechanism of Chinese herbal formulae systematically and comprehensively. Zhang et al. have put forward the concept of network pharmacology [13] to probe the influence or intervention of drugs and to reveal the synergism law of multicomponent drugs to seek high efficacy and low toxicity of multiple target medications. At the same time, the herbal formula is considered as multitarget, multichannel, multicomponent, and multidirectional therapeutic which meet the requirement of curing complicated illnesses in an integrated manner. Thus, we utilize the network pharmacological methods from the perspective of multitarget to combine drugs, targets, and diseases, aiming to provide new ways and new tactics for new medicine research and development [14]. Hence, we select a comprehensive network pharmacology method to uncover the pharmacological mechanism of XYP on anovulatory infertility, which supplies a precious chance for a thorough comprehension of the mechanism for reversing this illness-associated imbalanced network.

### 2. Materials and Methods

#### 2.1. Data Preparation

**2.1.1. Composite Compounds of Each Herb in XYP.** To collect the compounds of XYP, we used the TCM Database@Taiwan [24] (http://tc.mcm.edu.tw/zh-tw/, updated in March 2014), which is the most comprehensive TCM database in the world, and the Traditional Chinese Medicine Systems Pharmacology Database [25] (TcmSP, http://lsp.nwsuaf.edu.cn, updated on May 31, 2014), a unique system pharmacology platform designed for Chinese herbal medicines. Nine hundred and fifty-eight compounds were found, 348 in *Radix Bupleuri*, 175 in *Angelicae Sinensis Radix*, 92 in *Paeoniae Radix Alba*, 52 in *Poria Cocos* (Schw.) *Wolf.*, 63 in *Atractylodes Macrocephala Koidz.*, and 318 in *Licorice*. According to research [26, 27], we filtered these compounds and get 11 representative compounds (active compounds); they are saikosaponin, longispinogenin, ferulic acid, ligustilide, total glucosides of peony (TGP), atractylol, atractylenolide I, atractylenolide III, pachymy, pachymic acid, and glycyrrhizin. The details are described in Table S1 (see Supplementary Material available online at http://dx.doi.org/10.1155/2016/2960372).

**2.1.2. Modern Pharmacology and Traditional Effects of XYP.** We used China National Knowledge Infrastructure (CNKI), Pubmed, and Embase to obtain the information of modern pharmacology of XYP’s active compounds and mechanism of its traditional effects. Then, we found that expect longispinogenin and atractylol, all of the active compounds have various pharmacological effects. Also, we found that XYP can relieve the syndrome of stagnation of liver Qi, blood deficiency, and spleen weakness through dispersing stagnated liver Qi for relieving Qi stagnation, nourishing blood, and invigorating the spleen.

**2.1.3. Compound Target for Each Herb in XYP.** Input all the active compounds into SciFinder (http://scifinder.cas.org), a database of chemical and bibliographic information attached to the Chemical Abstracts Service; get the molecular structure of each active compound. Draw them in ChemBioDraw and save as “mol2” file format. Import them into PharmMapper (http://silab.ecust.edu.cn/pharmmapper/, updated in September 2012), which is a web server for potential drug target identification using pharmacophore mapping approach [28]. Because of the nonstandard naming, we used UniProtKB (http://www.uniprot.org/), which is the central hub for the collection of functional information on proteins, with accurate, consistent, and rich annotation. Input the protein names with the species limited to “Homo sapiens” and we could receive their official symbol. After these operations, protein information of active compounds was obtained. Finally, we utilized Kyoto Encyclopedia of Genes and Genomes (KEGG) database (http://www.genome.jp/kegg/, updated in May 2016) for noting pathway and filtering out protein targets that can be noted by human signal pathway. We used saikosaponin a and saikosaponin d instead of saikosaponin and used peoniflorin instead of TGP to obtain targets because of their high activity. The details are described in Table S2.

**2.1.4. Infertility Targets.** We collected infertility targets in Therapeutic Target Database [29] (http://database.idrb.cqu.edu.cn/TTD/, updated on Sep 10, 2015), which offers information about nucleic acid targets and therapeutic. Nine targets about infertility were acquired. The details are described in Table S3.

**2.1.5. Protein-Protein Interaction Data.** The data of protein-protein interaction (PPI) come from String [30] (http://string-db.org/, ver. 10) with the species limited to “Homo sapiens” and a confidence score >0.4 and InAct [31] (http://www.ebi.ac.uk/intact/, ver. 4.2.3.2).

String is a database of known and forecasted protein-protein interactions and InAct provides an open source database and analysis tools for molecular interaction data.
2.2. Network Construction

2.2.1. Network Construction Method. Network construction was performed as follows: (1) relationship between modern pharmacology and traditional efficacies of XYP; (2) active compound-active compound target network of XYP; (3) herb-active compound target-infertility target network of XYP; (4) active compound target-infertility target-other human proteins’ PPI network.

All the networks can be created via utilizing the network visualization software Cytoscape [32] (http://cytoscape.org/, ver. 3.2.1). It is the software that applies to visualizing biological pathways, intermolecular interaction networks, and many more. Furthermore, it supplies a basic set of features for data integration, analysis, and visualization for complicated network analysis.

2.2.2. Network Topological Feature Set Definition. Every node in a network is evaluated by three indices: degree, node betweenness, and closeness. Degree stands for the number of edges between a node and other nodes in a network [33]. Node betweenness evaluates the participation of a node in the shortest parts of a network and reflects the capability of nodes to manage the rate of information flow in the network [34]. Closeness is the inverse of the sum of the distance from node to other nodes [35]. The higher these three indices are, the more important the node is in the network.

2.3. Gene Ontology Enrichment Analysis. The Database for Annotation, Visualization and Integrated Discovery [36] (DAVID, https://david.ncifcrf.gov/home.jsp, ver. 6.7) was applied for Gene Ontology (GO) enrichment analysis.

3. Results and Discussion

3.1. Relationship between Modern Pharmacology and Traditional Efficacies of XYP

3.1.1. Syndrome of Stagnation of Liver Qi, Blood Deficiency, and Spleen Weakness. In TCM theory, XYP is used for dispersing stagnated liver Qi for relieving Qi stagnation, nourishing blood, and invigorating the spleen. Therefore, XYP can relieve syndrome of stagnation of liver Qi, blood deficiency, and spleen weakness. Based on current evidence, we summarize its mechanism.

According to Xiaolong [37] and Yanyan et al. [38], patients with syndrome of stagnation of liver Qi are immunosuppressed; for example, their immunoglobulin M (IgM), interleukin-1 (IL-1), IL-6, and IL-2 are reduced. Hepatic peroxidation was enhanced. The levels of reactive oxygen species and malondialdehyde (MDA) in liver were increased. Lipid peroxidation and lipid peroxidation (LPO) levels increased. This causes hepatic peroxidation damage. Meanwhile, plasma viscosity and the erythrocyte aggregation index increased significantly; the balance of thromboxane A2 (TXA2)/prostaglandin I2 (PGI2) was broken and the plasma glucose level was increased.

In Yi et al.'s study [39], blood deficiency is related to the reduction of erythrocytes quantity and quality and superoxide dismutase (SOD) activity, imbalance of helper T cell (TH)/suppressor T cell (TS), and lack of glutathione peroxidase (GSH-PX).

“Spleen weakness” has a vague definition in TCM theory. According to XYP’s main effects, we consider it as a syndrome of deficiency of spleen Qi and spleen failing to manage blood. Studies [40, 41] show that patient with the syndrome of spleen weakness has a weak digestive system, which manifests itself by low activity of salivary amylase and dysfunction of gastrointestinal absorption. In addition, reduction of erythrocytes quantity and quality and SOD activity, low total plasma protein (TP), abnormal coagulation and fibrinolysis system, accelerated heart rate, lack of GSH-PXPX, and decrease of thromboxane B2 (TXB2)/6-keto-PGF1α ratio also belong to it.

3.1.2. Relationship between Modern Pharmacology and Traditional Efficacies. We can find the relationship between active compounds’ modern pharmacology and XYP’s traditional efficacies when combined with collected data [15–23] (Table 1).

3.1.3. Complex Network Construction and Analysis. According to the data in Table 1, network is constructed. Based on available evidences, we can find that XYP's traditional effects have something to do with modern pharmacology. XYP may relieve syndrome of stagnation of liver Qi, blood deficiency, and spleen weakness. For example, pachymann can scavenge free radical, reduce LPO and MAD, lower blood sugar, and increase SOD activity. And this may be the mechanism of treating infertility. However, this cannot explain its molecular mechanism. Thus, we need to acquire the predictive targets, in order to explore it (Figure 1).

3.2. Compound-Compound Target Network Analysis. This network contains 244 nodes (233 compound target nodes and 11 active compound nodes) and 1075 edges. In this network, nodes close to the center show more interactions with compounds than peripheral nodes. This indicates that many targets are hit by multiple compounds, but some can be modulated by only one compound (peripheral nodes, such as MAOA, LYZ, and IMPA). AR, BACE1, CA2, GSTA1, and so on can be controlled by all 11 compounds, which may be the key targets in XYP. Atractylenolide I, saikosaponin a, and saikosaponin d can synergistically regulate ERBB4, INSR, and so on. Thus, we can have a rough observation on the relationships between active compounds and targets from compound-compound target network (Figure 2).

This suggests that XYP’s compounds may act on these targets synergistically and thus play a pharmacological role in other diseases besides infertility, which invisibly shows herbal formulae’s feature of multicomponent-multitarget-multidisease. Its potential effects may be found by this network.

3.3. Herb-Compound Target-Infertility Target Network Analysis. This network is set up to clear the relationship between six herbs, compound targets, and infertility targets. It is
### Table 1: Relationship between modern pharmacology and traditional efficacies of XYP.

| Active compounds  | Modern pharmacology of traditional efficacies                                                                 |
|-------------------|---------------------------------------------------------------------------------------------------------------|
| **Saikosaponin**  | Dispersing stagnated liver Qi for relieving Qi stagnation                                                    |
|                   | Nourishing the blood and invigorating the spleen                                                             |
|                   | Keeping a balance of TH/TS [15], reducing LPO [16]                                                          |
| **Ferulic acid**  | Increasing IL-2 and IgM level [15], reducing LPO and MAD, balancing TXA₂/PGI₂, reducing blood viscosity [17] |
|                   | Increasing SOD activity, enhancing hematopoietic function, keeping the balance of TXB₂/6-keto-PGF₁₂ [17]     |
| **Ligustilide**   | Decreasing MAD, balancing TXA₂/PGI₂, reducing blood viscosity [18, 19]                                      |
|                   | Increasing activity of SOD and GSH-PX [18, 19]                                                               |
| **TGP**           | Decreasing MAD, reducing blood viscosity, balancing TXA₂/PGI₂, increasing IL-2 [19, 20]                     |
|                   | Keeping a balance of TH/TS, increasing activity of SOD and GSH-PX, enhancing hematopoietic function, improving the quality of erythrocytes [19, 20] |
| **Atractylenolide I** | —                                                                 |
| **Atractylenolide III** | —                                                                 |
| **Pachyman**      | Scavenging free radical, reducing LPO and MAD, lowering blood sugar [22]                                    |
|                   | Increasing SOD activity [22]                                                                                  |
| **Pachymic**      | —                                                                                                           |
| **Glycyrrhizin**  | Reducing MAD, increasing IL-1β, IL-6, IL-10, scavenging free radical [23]                                    |
|                   | Increasing SOD activity, increasing the plasma TP [23]                                                       |

**Figure 1:** Relationship between modern pharmacology and traditional efficacies of XYP (pink circles, green octagons, fuchsia circles, and orange triangle stand for herbs, active compounds, modern pharmacology, and traditional effects, resp., and green lines, fuchsia lines, and orange lines stand for relationship between herbs and active compounds, active compounds and modern pharmacology, and modern pharmacology and traditional effects, resp.).
composed of 245 nodes (6 herbs, 231 compound targets, 6 infertility targets, and 2 compound-infertility targets) and 841 edges (Figure 3).

In Figure 3, we find that compound targets are also regulated by drug targets (infertility targets). This indicates that drugs may act on drug targets to regulate disease-related proteins indirectly, whereas XYP can act on these proteins directly. Alternatively, XYP may indirectly act on drug targets by regulating associated protein (compound targets) so as to achieve the effects similar to drug treatment.

In Figure 4, according to GO enrichment analysis, these targets are significantly associated with response to steroid hormone stimulus (GO ID: 48545; Fold Enrichment = 7.1; \(P < 0.001\)), steroid hormone receptor signaling pathway (GO ID: 30518; Fold Enrichment = 2.3; \(P = 0.0015\)), ovarian follicle development (GO ID: 1541; Fold Enrichment = 5.8; \(P = 0.001\)), response to insulin stimulus (GO ID: 32868; Fold Enrichment = 11.3; \(P < 0.001\)), and insulin receptor signaling pathway (GO ID: 8286; Fold Enrichment = 9.6; \(P = 0.0057\)). The details are described in Table S4.

In addition, in Figure 4, the number of targets of Angelicae Sinensis Radix and Atractylodes Macrocephala Koidz, is the largest (36 and 37), which indicates the two herbs play a major role in the treatment. This means the two herbs may be the major herbs in XYP.

In this network (Figure 4), we find a lot of infertility-related biological processes. Response to estrogen stimulus (GO: 43627), response to steroid hormone stimulus (GO: 48545), steroid hormone receptor signaling pathway (GO: 30518), response to insulin stimulus (GO: 32868), and insulin receptor signaling pathway (GO: 8286) are considered to
be the possible mechanism of treatment of anovulatory infertility, wherein the most important thing is the steroid hormone imbalance (imbalance of GO: 48545 and GO: 30518). Anovulatory infertility is mainly characterized by the nervous system-hypothalamus-pituitary dysfunction, that is, the balance of steroid hormone imbalance leading to follicular failure and no dominant follicle. And some steroid hormones (Gn, follicle-stimulating hormone [FSH], luteinizing hormone [LH], estrogen, and progesterone) are also used for drug treatment of anovulatory infertility.

Follicles growth benefits from the precise regulation of various steroid hormones. When primordial follicle grows, granulosa cells would express the receptors of FSH, GnRH, glucocorticoids, estrogen, and many more, which make themselves become primary follicles that can respond to hormone stimuli. Then, under the stimulation of hormone such as FSH and estrogen, granulosa cells proliferate and differentiate, and preantral follicle develops into secondary and tertiary follicle. And in the process of dominant follicular formation, estradiol (E_2) plays an important role. FSH promotes granulosa cells to synthesize and secrete estrogen; and estrogen leads them to express more receptors. Still, FSH causes preantral follicular granulosa cells to proliferate and differentiate and secrete follicle fluid, which promote granulosa cells to synthesize and secrete insulin-like growth factor (IGF), IGF receptor, inhibin, activin, and synergies with them. This can influence dominating follicular selection and nondominant follicular degeneration. During the late stages, FSH with estrogen pushes granulosa cells to express the LH receptor, which promote oocyte maturation and make preparations for ovulation and luteinization [42–44].

Secreton of FSH and LH of the pituitary is controlled by hypothalamic GnRH and ovarian estrogen. GnRH can promote pituitary synthesis and secretion of FSH and LH. GnRH with estrogen induces the expression of GnRH receptor so as to improve the pituitary sensitivity to GnRH. Estrogen has
a double effect on the synthesis and secretion of FSH and LH; when the concentration of E2 is low, the effect is negative feedback, and vice versa [45].

Latest research shows that ovulation disorders or ovarian dysfunction (unable to produce fertilizable oocytes) which is caused by endocrine disorders is the most common cause of female infertility [46-48]. The major reasons are serum steroid hormones (glucocorticoids, estrogen, progesterone, and androgen) disorder and ovarian lesions (ovarian dysfunction and follicle dysplasia) caused by abnormal expression of steroid hormone receptors in follicles [49-51]. The classical mode of action of steroids is that the hormone enters the cell, binds to the same receptor, and activates or inhibits transcription of the target gene [52]. However, it is worth noting that nonclassical pathways that are dependent on steroid hormones can also cause changes in gene transcription [53]. In PCOS, follicular development disorder is caused by low levels of estrogen and hyperandrogenism. Moreover, abnormal glucose metabolism is highly related to reproductive dysfunction; abnormal glucose metabolism is caused by insulin resistance and hyperinsulinemia.

Another major factor is neuroendocrine disorders caused by dysfunction of the hypothalamic-pituitary system [54-56]; related hormones include GnRH, FSH, LH, prolactin, GH-(growth hormone) IGF-1, adrenocorticotropin hormone, and thyroid hormone [57]. For instance, hyperandrogenism in PCOS is caused by dysfunction of hypothalamic-pituitary-ovarian axis and adrenal gland. In PCOS patients, the LH/FSH ratio is increasing and LH continues at a higher level. Meanwhile, the relationship between GnRH pulse frequency and Gn (FSH and LH) reactivity may be the key to abnormal secretion of gonadal hormones [58].

3.4. Compound Target-Infertility Target-Other Human Proteins’ PPI Network Analysis. The network contains 1499 nodes (231 compound targets, 7 infertility targets, 2 compound-infertility targets, and 1259 other human proteins) and 54998 edges (Figure 5). In this network, nodes whose three indices are higher than their average (degree ≥ 61.37, node betweenness ≥ 0.000838, and closeness ≥ 0.4485) will be regarded as main nodes. Finally, 235 main nodes are selected. The details are described in Table S5.

Furthermore, according to GO enrichment analysis, a direct interaction network between the main nodes is established, which contains 112 nodes (31 compound targets, 1 infertility target, 1 compound-infertility target, and 79 other human proteins). As shown in Figure 6, the main nodes can be divided into four functional modules, including response to steroid hormone stimulus (GO ID: 30518; Fold Enrichment = 11.1; P < 0.001), response to progesterone stimulus (GO ID: 35020; Fold Enrichment = 11; P < 0.001), response to estrogen stimulus (GO ID: 43627; Fold Enrichment = 11; P < 0.001), response to growth hormone stimulus (GO ID: 32570; Fold Enrichment = 7; P < 0.001), response to insulin stimulus (GO ID: 32868; Fold Enrichment = 13.4; P < 0.001), response to progesterone stimulus (GO ID: 35020; Fold Enrichment = 3.4; P = 0.014), growth hormone receptor signaling pathway (GO ID: 60396; Fold Enrichment = 21.9; P = 0.037), response to growth hormone stimulus (GO ID: 60416; Fold Enrichment = 19.4; P = 0.046), ovulation cycle process (GO ID: 22602; Fold Enrichment = 10.3; P < 0.001), reproductive process in a multicellular organism (GO ID: 48609; Fold Enrichment = 4; P < 0.001), ovarian follicle development (GO ID: 1541; Fold Enrichment = 7.1; P = 0.0051), response to insulin stimulus (GO ID: 32868; Fold Enrichment = 13.4; P < 0.001), insulin receptor signaling pathway (GO ID: 8286;
Figure 5: Active compound target-infertility target-other human proteins' PPI network (blue circle, pink circle, green circle, and red circle stand for another human protein, compound targets, compound-infertility targets, and infertility targets, resp.).
Fold Enrichment = 17.3; \( P < 0.001 \), apoptosis (GO ID: 6915; Fold Enrichment = 4; \( P < 0.001 \)), and antiapoptosis (GO ID: 43066; Fold Enrichment = 7.9; \( P < 0.001 \)). The details are described in Table S6.

By two GO enrichment analyses, one interesting phenomenon is observed; that is, XYP may have similar effects of procreation endocrine regulation to western medicines by target clinically used therapeutic targets such as estrogen receptor alpha (ESR1), estrogen receptor beta (ESR2), progesterone receptor (PGR), insulin receptor (INSR), androgen receptor (AR) follicle-stimulating hormone receptor (FSHR), and lutropin-choriogonadotropic hormone receptor (LHCGR). AR has the highest number of compound target interactions and is likely to play a key role in the treatment of anovulatory infertility. ESR1 and ESR2 are associated with E2 disorder of anovulatory infertility. PGR is related to its progesterone disorder. INSR and AR are linked to insulin resistance and hyperandrogenism, respectively; FSH and LHCGR are associated with neuroendocrine disorders. Through Figure 2, we find that ESR2 is hit by atracylol, pachymic acid, and ligustilide; PGR is regulated by atracylenolide I, atracylenolide III, atracylol, saikosaponin a, saikosaponin d, pachymic acid, ligustilide, paeoniflorin, and glycyrrhizin; INSR is linked to atracylenolide I, atracylenolide III, saikosaponin a, saikosaponin d, paeoniflorin, and glycyrrhizin; AR is connected with all 11 compounds. We also find that the compounds act synergistically on these 5 key targets, which demonstrates herbal formulae’s feature of multicomponent-multitarget and synergistic effects.

In addition, with two GO enrichment analyses, we find that key targets affiliate to GO: 43627, GO: 48545, GO: 30518, GO: 32868, GO: 8286, GO: 30518, GO: 32570, and GO: 30520. This indicates that XYP acts on key targets with its compounds to regulate disease-related biological processes and through regulating disease-related biological processes it intervenes multifactor anovulatory infertility and therefore exerts a therapeutic effect.

Research has shown that steroids and hormones secreted by hypothalamic-pituitary-ovarian axis are able to regulate follicular development and maturation. The steroid metabolic disorder usually results in reproductive system diseases, such as PCOS. Its pathophysiological mechanism consists of four aspects including dysfunction of hypothalamic-pituitary-ovarian axis, insulin resistance and hyperinsulinemia, dysfunction of adrenal endocrine, and multisystem multiorgan abnormalities [59]. Under the circumstance, due to the low level of FSH, aromatase cannot transform androgens sufficiently, and then androgen accumulates in follicles, hindering follicular maturation and ovulation. Meanwhile, the high level of androgen will sensitize hypothalamic-pituitary axis so as to promote LH secretion. This will break normal physiological follicular growth [60–62]. Finally, due to the excessive follicular recruitment, follicular selection and dominance of PCOS patients will stop and then lead to anovulatory infertility.

The dysfunction of hypothalamic-pituitary-ovarian axis in PCOS patients results in Gn imbalance, which leads to LH higher than FSH and LH/FSH ratio increases. The lower level of FSH will result in lower levels of E2, which makes ovulation difficult and makes follicular and oocytes growth slow or stagnant. Meanwhile, because of pituitary sensitization, more FSH and LH are secreted, which causes new follicles continuing to grow but cannot reach maturity. Then hyperplastic theca cells of these small follicles show luteinization stimulated by high LH levels [63]. When using ovulation induction treatment, GH’s application can improve ovarian responsiveness to GnRH. PCOS patients often present low basal GH; after applying levodopa, GH rising magnitude is reduced, which shows that PCOS patients have not only growth hormone deficiency, but also low activity of hypothalamic dopamine [64, 65].

**Figure 6: Active compound target-infertility target-other human proteins’ PPI network: according to the associated biological processes or pathways, these nodes can be categorized into four parts (green hexagon, pink circle, yellow circle, red circle, and purple circle stand for herb, compound target, another human protein, infertility target, and compound-infertility target, resp.; gray lines stand for the relation of herb and black lines stand for the relation of infertility targets and compound targets and compound-infertility targets).**
In clinical setting, cure rate of anovulatory infertility is the highest [66]. The main treatment is symptomatic treatment based on the cause of anovulation, such as clomiphene citrate, clomiphene with hormones, Gn, GnRH, bromocriptine, and glucocorticoid. However, long-term application may induce reproductive system’s tumor, ovarian hyperstimulation syndrome, and so on. In this aspect, current studies show that XYP can regulate steroids and their receptors and control hypothalamic-pituitary-ovarian axis to promote follicular maturation and dominant follicular generation, so that those oocytes can be fertilized [10, 11].

In adult patients, steroid disorders, abnormal expression of steroid receptors, and hypothalamic-pituitary dysfunction can lead to follicular growth stagnation or atresia and depletion of oocytes pool and ultimately result in no dominant follicle or mature follicle. This is the negative effect of the relative imbalance of steroid hormones and ovarian hormones on follicular development. Meantime, this can lead to abnormal proliferation or excessive apoptosis of follicular cells such as the high LH dependent theca cellular proliferation [67] and follicular atresia induced granulosa cell inactivation [68–70] in PCOS and depletion of oocytes pool in premature ovarian failure [71]. Therefore, based on our findings, XYP for anovulatory infertility in two aspects, steroid hormones and their receptors and neuroendocrine, will be the focus of further exploration and research.

Apoptosis, especially in the process of depletion of oocytes pool, oocytes and granulosa cells apoptosis have been shown to have a close relationship with follicular atresia. Moreover, both theca cellular hyperproliferation and follicular atresia induced granulosa cell inactivation in hyperandrogenism (such as PCOS) mediate follicular abnormal development. As the vital ovarian cells, granulosa cells change their form, function, and many more in each stage of follicular development [72], while the oocytes have guided the proliferation and differentiation of them. Still, oocytes maturation is affected by granulosa cells. This procedure is complex and involves multiple pathways and signaling molecules; any link abnormalities may lead to follicular development abnormalities [73]. Thus, inhibiting oocytes and granulosa cells apoptosis and maintaining theca cell growth are the potential strategies to promote the formation of dominant follicle or mature follicle. The current study has shown that saikosaponin d has estrogen-like effects [74] which increase serum E2 and P concentrations in ovariectomized rats. Glycyrrhizin can also increase serum E2 and P concentrations [75]. Jing< found that ferulic acid can reduce the ovary granulosa cell apoptosis rate of PCOS rats and promote luteal formation; still, it can lower serum E2 concentration and increase serum P concentration [76]. Thus, we could assume that XYP can regulate steroid hormone synthesis and secretion, thus promoting follicular development and formation of the dominant follicle, and reduce follicle depletion, so as to protect oocytes and granulosa cells from apoptosis.

4. Conclusions

Currently, as to anovulatory infertility, perfect treatment has not been discovered. Western medicine’s therapeutic strategy is symptomatic treatment. But the side effects of long-term application cannot be ignored. TCM recipe has effect on some incurable diseases such as infertility and is more systematic and holistic. However, many studies are still applying the traditional research idea, “one-drug-one target-one illness,” which ignores the multitarget and multicomponent characteristic of TCM recipes. Inspired by Tang et al’s research [77], we decided to solve this problem by network pharmacology. In this study, a number of network-based computational methods and algorithm-based approaches to predict targets and construct networks are combined to illuminate the molecular synergy of XYP for infertility. This method provides clues to the researcher who explores TCM’s various synergies. It also supplies reference information to researchers who want to explore XYP’s therapeutic (such as treating infertility) mechanism. The most important thing is that we initially identify XYP’s molecular mechanism for treating anovulatory infertility. Our study has successfully found the potential infertility-related targets and biological processes in XYP and uncovered the rationality of herb combinations of XYP. Therefore, such a network pharmacology strategy and platform are expected to make the systematic study of herbal formulae for disease (e.g., XYP for anovulatory infertility) achievable and make the TCM drug discovery predictable.

Competing Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

Authors’ Contributions

Liuting Zeng and Kailin Yang contributed to this work. Liuting Zeng, Kailin Yang, and Huiping Liu are responsible for the study concept, design, and literature searching; Liuting Zeng and Kailin Yang are responsible for data analysis and interpretation; Liuting and Kailin Yang and drafted the paper; Huiping Liu supervised the study; Huiping Liu and Guomin Zhang carried out extensive revision of the manuscript; all authors participated in the analysis and interpretation of data and approved the final paper. Liu Huiping, Liuting Zeng, and Kailin Yang contributed equally to this work; they are co-first authors.

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