A Preliminary Inquiry Into the Potential Mechanism of Huang-Lian-Jie-Du Decoction in Treating Rheumatoid Arthritis via Network Pharmacology and Molecular Docking

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Huang-Lian-Jie-Du decoction (HLJDD) has been widely applied to treat inflammation-associated diseases for thousands of years in China. However, the concrete molecular mechanism of HLJDD in the treatment of rheumatoid arthritis (RA) remains unclear. In this work, network pharmacology and molecular docking were applied to preliminarily analyze the potential active ingredients, drug targets, and related pathways of HLJDD on treating RA. A total of 102 active compounds with corresponding 189 targets were identified from HLJDD, and 41 common targets were further identified by intersecting with RA-related targets. Functional enrichment analysis was performed to screen the biological pathways associated with RA. Ten hub targets were further identified through constructing the protein–protein interaction (PPI) network of common targets, which were mainly enriched in the interleukin-17 (IL-17) signaling pathway, tumor necrosis factor (TNF) signaling pathway, and Toll-like receptor signaling pathway. Furthermore, a complex botanical drugs-ingredients-hub-targets-disease network was successfully constructed. The molecular docking results exhibited that these vital ingredients of HLJDD had a stable binding to the hub targets. Among these ingredients, quercetin (MOL000098) was the most common molecule with stable binding to all the targets, and PTGS2 was considered the most important target with multiple regulations by the most active ingredients. In vitro, we successfully validated the inhibitory role of quercetin in the cellular proliferation of human RA fibroblast-like synoviocyte cell line (MH7A cells). These findings indicated that the potential mechanisms of HLJDD for RA treatment might be attributed to inhibiting the immune-inflammatory response, reducing the release of chemokines, and alleviating the destruction of extracellular matrix (ECM) in the synovial compartment.

Keywords: Huang-Lian-Jie-Du decoction, rheumatoid arthritis, functional enrichment analysis, network pharmacology, molecular docking
INTRODUCTION

Rheumatoid arthritis (RA) is a chronic inflammatory autoimmune disease characterized by synovitis, pannus formation, and cartilage erosion, eventually leading to progressive joint dysfunction, deformity, and increased mortality risk (Smolen et al., 2016). The prevalence rate of RA is approximately 1% in the world’s population, with higher morbidity among elderly women (Abbasi et al., 2019). So far, the treatment of RA was mainly dependent on the appropriate combination of physical, pharmacological, and surgical approaches, of which nonsteroidal anti-inflammatory drugs (NSAIDs) and disease-modifying antirheumatic drugs (DMARDs) were widely used to lighten inflammation and improve joint function (Wilsdon and Hill, 2017; Conigliaro et al., 2019). However, these drugs may result in some adverse effects, including infection, gastrointestinal reaction, skin rashes, bone marrow suppression, and liver and kidney toxicity (Chen et al., 2019).

Recent advancements in pharmacy have attracted increasing attention on the use of traditional Chinese medicine (TCM) for the treatment of RA due to its significant therapeutic efficacy and mild side effects (Huang et al., 2016). As a well-known classic TCM formula for clearing heat and detoxification, Huang-Lian-Jie-Du decoction (HLJDD) is composed of Scutellaria baicalensis Georgi (Huangqin, HQ), Phellodendron amurense Rupr. (Huangbai, HB), Coptis chinensis Franch. (Huanglian, HL), and Gardenia jasminoides J.Ellis (Zhizi, ZZ) with a ratio of 2:2:3:3 (Supplementary Table S1). HLJDD has been widely applied to the treatment of inflammation-associated diseases including hepatitis (Wei et al., 2016), pneumonia (Li et al., 2021), Alzheimer’s disease (AD) (Gu et al., 2018), inflammatory bowel disease (IBD) (Yuan et al., 2019), and RA (Hu et al., 2013). Furthermore, numerous studies have shown that HLJDD could significantly reduce the levels of inflammatory cytokines such as TNF-α and inflammatory mediators such as prostaglandin E2 (PGE2) to realize the anti-inflammatory efficacy (Fang et al., 2004; Lu et al., 2011). In vitro, Sung et al. (2012) demonstrated that quercetin could significantly inhibit unstimulated and IL-1β induced proliferation of rheumatoid synovial fibroblasts (RASFs) and the messenger ribonucleic acid and protein expression of MMP-1, 3, COX-2. In animal models, Hu et al. also successfully demonstrated the treatments of HLJDT on collagen-induced arthritis in rats (Hu et al., 2013). During the period of COVID-19, HLJDD was also found to play a therapeutic role in COVID-19 through regulating multiple signaling pathways based on targeting genes such as IL6, IL10, MMP9, NOS2, VEGF, and TGFβ1 (Liu et al., 2021). Nevertheless, the concrete molecular mechanism of HLJDD in the treatment of RA was still unclear.

As a novel, promising, and cost-effective drug research approach based on bioinformatics and pharmacology, network pharmacology has been widely used in studies on the molecular mechanism of drugs with molecular docking. In previous studies, Yang et al. (2013) identified berberine, baicalin, and geniposide as three major ingredients of HLJDD and further demonstrated their anti-inflammatory effects through inhibiting NF-κB and MAPK signaling pathways in a dose-dependent manner. In addition, using network pharmacology and metabolomics analysis, Qu et al. found the antidepressant effects of HLJDD through candicine, berberine, tetrahydroberberine, and baicalin to regulate SLC6A4 and MAOA in tryptophan metabolism (Qu et al., 2021). In Li’s study, the network pharmacology and molecular docking method was also well conducted to decode the potential mechanism of HLJDD in treating pneumonia and construct a complex botanical drugs-ingredients-targets-disease network (Li et al., 2021). These studies provided a convincing possibility that network pharmacology and molecular docking analysis can be effectively applied to investigate potential therapeutic targets of HLJDD and facilitate a deep understanding of its underlying mechanism in RA.

In the present study, we investigated the potential targets and pathways of HLJDD in treating RA and constructed the botanical drugs-ingredients-targets-disease network through network pharmacology analysis and molecular docking validation. We were convinced that our results would help in illuminating HLJDD’s possible mechanism of the treatment in RA, thus improving the curative effect and prognosis for RA. The workflow of the whole process of our study is shown in Figure 1.

MATERIALS AND METHODS

Screening Active Ingredients and Potential Targets in HLJDD

In this study, the Traditional Chinese Medicine Systems Pharmacology (TCMSP) (Ru et al., 2014) database was utilized to screen active ingredients of HLJDD by using “Huangbai,” “Huanglian,” “Huangqin,” and “Zhizí” as keywords with the filtration criteria of bioavailability (OB) ≥30% and drug-likeness (DL) ≥0.18, respectively (Song et al., 2018). The corresponding molecular structure, structural information, and PubChem CID of those active ingredients were further obtained from the “PubChem” online tool (Wang et al., 2017). Subsequently, using the above structural information of ingredients, we utilized the “BATMEN-TCM” online tools to identify potential targets in HLJDD with a score ≥20 and adjusted p ≤ 0.05 (Liu et al., 2016).

Identification of Target Genes in Rheumatoid Arthritis

The RA-associated target genes were obtained from the OMIM database (Rigden and Fernandez, 2020), PharmGKB database (Whirl-Carrillo et al., 2012), DrugBank database (Wishart et al., 2018), and TTD database (Li Y. H. et al., 2018), respectively. By searching “rheumatoid arthritis” as the keyword, we identified 12, 13, 299, and 157 targets related to RA from these databases, respectively. The UniProt database was further employed to standardize their protein structures in unique ways (UniProt Consortium, 2019).

Construction of PPI Network and Cluster Analysis

The PPI data of the potential target interactions were downloaded from the STRING database (version 11.0) (Szklarczyk et al., 2019) with setting the Organism as “Homo sapiens” and confidence
score $\geq 0.4$. Then, the PPI network was constructed and visualized by the Cytoscape software (Shannon et al., 2003), and hub targets were detected according to levers of degree value (the number of interactions for each node) in the PPI network. In the further cluster analysis, the plug-in MCODE (Deng et al., 2019) was used to identify significant modules using the filter conditions: $k\text{ core }$
2, maximum depth = 100, node score cutoff = 0.2, and degree cutoff = 2.

GO and KEGG Functional Enrichment Analysis

To investigate the biological function of potential targets in RA, we performed gene ontology (GO) analysis, including biological processes (BP), molecular functions (MF), and cellular components (CC), and Kyoto Encyclopedia of Genes and Genomes (KEGG) functional enrichment analysis. These analyses were conducted through the online tool “g:Profiler” (Raudvere et al., 2019) with Bonferroni correction and adjusted $p \leq 0.05$. Moreover, the ClueGo (Bindea et al., 2009) and CluePedia (Bindea et al., 2013) plug-in of Cytoscape software were further applied to visualize the network of significant pathways and their corresponding target genes.

Identification of Hub Targets and Critical Network of HLJDD on RA

The common targets between HLJDD and RA were identified through taking the intersection, and the PPI network of these targets was also constructed and visualized in the Cytoscape software. Subsequently, we used plug-in cytoHubba (Chin et al., 2014) to explore important nodes in the above PPI network by 12 topological algorithms including MCC, MNC, DMNC, BottleNeck, Eccentricity, EPC, Radiality, Betweenness, Closeness, Clustering Coefficient, Degree, and Stress. The top 10 nodes identified by each topological algorithm were chosen to identify the shared genes more than six ways as the most pivotal hub genes in the network (Luo et al., 2020). Finally, a complex botanical drugs-ingredients-hub-targets-disease network was further constructed and visualized in the Cytoscape.

Visualization and Validation of Molecular Docking

To validate the binding of hub targets and active ingredients in HLJDD, the 3D molecular conformations of ingredients were retrieved from the PubChem Compound database. The crystal structures of target proteins were obtained from the RCSB Protein Data Bank (PDB) database (Burley et al., 2021). AutoDock 4.2.6 tool was used to remove the redundant structure, ligands, and water molecules and add polar hydrogen atoms and partial charges into protein crystal structures before running molecular docking (Li H. et al., 2018). The visualization of molecular docking was exhibited by PyMol software (Seeliger and de Groot, 2010). The transcriptomic dataset of peripheral blood mononuclear cells (PBMCs) (GSE17755, including 112 RA and 45 controls) was downloaded from Gene Expression. Omnibus (GEO) database to validate the normalized expression of RA-related target genes. Values were presented as the mean ± standard deviation (SD), and the comparison between groups was performed using the Wilcoxon test by the “ggpubr” R package. $p < 0.05$ was considered statistically significant.

Cell Culture and Treatment

Human RA fibroblast-like synoviocyte cell line (MH7A cells) was purchased from the Riken cell bank (Ibaraki, Japan), and cells were maintained in DMEM supplemented with 10% fetal bovine serum (FBS) and 1% penicillin-streptomycin solution (HyClone, Shanghai). Subsequently, the MH7A cells were plated into a six-well plate ($5 \times 10^3$ cells/well) and treated with different concentrations of quercetin (0, 25, 50, 75, and 100 μM; Sigma, St Louis, MO, United States) in 37°C incubator with 5% CO$_2$ atmosphere for 24 h. Meanwhile, the culture medium with 10% DMEM and 0.05% DMSO but without MH7A cells was set as control groups and also plated in the same condition for 24 h.

Cell Proliferation Assay

Equal numbers of the above MH7A cells were plated in a 96-well plate ($5 \times 10^3$ cells/well), and Cell Counting Kit-8 (CCK-8) (Dojindo, Tokyo, Japan) (10 μL/well) was applied to measure the cell proliferation at 24 h. Finally, the imaging of cell proliferation was recorded, and the absorbance value (OD value; A$_{450nm}$) of MH7A cells was further detected using a Microplate Reader (Thermo Fisher, MA). Based on the OD value of different quercetin concentrations, we successfully drew the drug-concentration curve to identify the half-maximal inhibitory concentration (IC50) value of quercetin in RA.

Statistical Analysis

All statistical analyses were performed in R software (version 4.0.1, https://www.r-project.org/). Continuous variables were presented as mean ± standard deviation (SD), and Wilcoxon signed-rank test was used to compare continuous variables. The two-tailed $P$ value $<0.05$ was considered statistically significant.

RESULT

Identification of Potential Targets-Active Ingredients Network

Based on the criteria of OB $\geq$ 30% and DL $\geq$ 0.18, a total of 102 active ingredients were predicted in HLJDD through the TCMSP database, including 37 from HB, 14 from HL, 36 from HQ, and 15 from ZZ (Supplementary Figure 1A). Moreover, we also screened a total of 1008 corresponding targets of HLJDD using the TCMSP and BATMEN-TCM, of which 271 were from HB, 216 from HL, 226 from HQ, and 295 from ZZ (Supplementary Table S3). Combined with the 102 active ingredients, 189 targets of these ingredients were further identified, and the complex targets-active ingredients network is constructed in Figure 2A. Notably, by setting the degree value $\geq$ 45, the top 10 active ingredients were identified in the network, including quercetin (MOL000098), beta-sitosterol (MOL000358), stigmastanol (MOL000449), kaempferol (MOL000422), wogonin (MOL000173), baicalein (MOL002714), palmatine (MOL00785), berberine (MOL001454), coptisine
(MOL001458), and 5-hydroxy-7-methoxy-2-(3,4,5-trimethoxyphenyl) chromone (MOL003095). The detailed characteristics of these active ingredients and targets are shown in Supplementary Table S2 and Supplementary Table S3. To further identify the changes of gene expression in the progress of RA, four databases, namely, OMIM, GenBank, PharmGKB, and TTD, were screened to obtain RA-related target genes, and a total of 440 targets of RA were obtained (Supplementary Figure 1B, Supplementary Table S4). Subsequently, comparing the target genes of HLJDD and RA,
we found that RA shared 41 common targets with that of active ingredients (Figure 2A).

### PPI Network and Cluster Analysis of Disease Targets in RA

To identify central target genes of RA in the physical interaction network and provide potential clues for the possible pathogenic mechanisms of RA, we successfully constructed an interconnected PPI network of 440 RA-related targets with 394 nodes and 9391 edges (Figure 2C) based on the STRING database. Using the MCODE plug-in in Cytoscape with MCODE score greater than 5, we further identified five clusters with close interconnection (Supplementary Figure 1C, Table 1). The blue nodes represented the significant targets with high degree values in the network, which might play essential roles in the progression of RA.

### Functional Enrichment Analysis of Disease Targets in RA

In order to further interpret biological processes associated with the RA-related targets, GO and KEGG enrichment were performed based on the 440 target genes. It turned out that these target genes were enriched in 1237 BPs, 81 MFs, 67 CCs, and 76 KEGG pathways (Supplementary Table S5). The results showed that the biological processes of RA were mainly related to the immune response process, such as “immune system process,” “response to organic substance,” “cellular response to cytokine stimulus,” and “inflammatory response.” The molecular functions of RA might be related to “signaling receptor binding,” “molecular transducer activity,” and “signaling receptor activity.” The cellular components were principally associated with “cell periphery,” “extracellular region,” and “plasma membrane” (Figure 3A). The KEGG pathway analysis also revealed these targets were significantly enriched in immune-activated associated diseases and pathways, including “rheumatoid arthritis,” “inflammatory bowel disease,” “cytokine-cytokine receptor interaction,” “TNF signaling pathway,” “NF-kappa B signaling pathway,” and “Th17 cell differentiation” (Figure 3B). In addition, the concrete network of the KEGG pathways indicated that massive inflammatory cytokines participated in the progression of RA (Figure 3C).

### Identification of Hub Genes and Critical Network of HLJDD on RA

In order to identify the hub genes of HLJDD in treating RA, we also imported the 41 common targets into the STRING database and subsequently constructed their PPI network (Figure 2D). The cytoHubba plug-in was further applied to calculate the significance of targets with 12 methods, and we found that there were 10 hub genes shared with more than 6 topological analyses (MMP9, IL1β, IFNG, IL10, ICAM1, CCL2, PTGS2, IL6, CXCL8, and JUN) by selecting top 10 genes of each method (Table 2). The PPI network of the hub genes included 10 nodes and 45 edges, with an average node degree of 9 and P value of 3.22e-14 (Figure 2D).

Furthermore, by selecting the hub gene-related active ingredients in HLJDD, we successfully constructed the botanical drugs-ingredients-targets-disease network (Figure 3D), containing 1 disease, 4 botanical drugs, 54 ingredients, and 10 hub genes. It could be seen that quercetin (MOL000098), beta-sitosterol (MOL000358), wogonin (MOL000173), kaempferol (MOL000422), coptisine (MOL000458), and stigmasterol (MOL000449) were pivotal ingredients that were identified with high degree value, suggesting these ingredients might be the material basis for HLJDD in treating RA.

### Selecting and Analyzing Critical GO and KEGG of Hub Genes

In order to further interpret the mechanism of HLJDD in the treatment of RA, we used the common 41 targets to perform GO and KEGG functional enrichment analysis. A total of 631 BPs, 5 CCs, 19 MFs, and 78 KEGG pathways were identified, and the top 10 biological processes are exhibited in Figure 4. GO enrichment analysis demonstrated that the common targets were mainly enriched in the inflammatory response and cytokine-associated processes such as “inflammatory response,” “cellular response to cytokine stimulus,” “cytokine-mediated signaling pathway,” and “response to cytokine.” The main cellular components were
FIGURE 3 | Functional enrichment analysis of total RA-related targets. (A) Bubble diagram showing the results of GO enrichment analysis, including top 10 terms in BP, MF, and CC, respectively. (B) Top 10 pathways of KEGG enrichment analysis were associated with immune activation. (C) The network showing the detailed genes involved in the top 10 pathways and the hub targets were enlarged. (D) The network showing the KEGG pathways enrichment of 10 hub targets.
located in the extracellular region, and the molecular functions were mainly enriched in "signaling receptor binding," "cytokine receptor binding," and "cytokine activity" (Figure 4A). As to KEGG pathways analysis, the most significant pathways of the common targets were "IL-17 signaling pathway," "TNF signaling pathway," "AGE–RAGE signaling pathway in diabetic complications," and "rheumatoid arthritis" (Figure 4B), and the interaction network also demonstrated that the ten hub genes participated in these significant pathways (Figure 4C). Moreover, KEGG analysis indicated that the hub targets were significantly enriched in biological processes associated with immune activation, including IL-17 signaling pathway, Toll-like receptor signaling pathway, TNF signaling pathway, fluid shear stress, atherosclerosis, and rheumatoid arthritis pathways (Figure 4D).

Validation of Hub Genes by Molecular Docking and mRNA Expression

To further evaluate and validate the effective binding of targets and active ingredients, ten hub genes and nine pivotal ingredients were used to perform molecular docking as receptors and ligands, respectively. The stability and strength of the binding were based on the binding energy. The lower the binding energy value was, the more stable the binding was. Setting the binding energy value as the threshold, we successfully demonstrated the ligands binding well to the target receptor with the lowest energy (Table 3). Among these ingredients, quercetin (MOL000098) was the most significant molecule with stable binding to all the targets, and PTGS2 was considered the most important target with multiple regulations by the most active ingredients (Figure 5). Normalized expression of 10 hub targets is also validated in GEO datasets, and it revealed that IL6, IL1β, JUN, CXCL8, MMP9, PTGS2, ICAM1, and CCL2 were significantly upregulated. In contrast, IFNG and IL10 were significantly downregulated in the PBMC of RA patients compared to controls (Figures 6A–J).

Quercetin Inhibited Cell Proliferation in MH7A Cells

Initially, we estimated the proliferation of MH7A cells after the treatment of series quercetin concentrations (0, 25, 50, 75, and 100 μM) and found the inhibitory effectiveness was generally increased with the enhancement of quercetin (Figure 6K). However, the growth did not increase speedily when the concentration was more than 50 μM (Figure 6L), and the IC50 value of quercetin was identified as 38.02 μM according to the drug-concentration curve (Figure 6M).

DISCUSSION

As one of the most common systematic autoimmune disorders, RA has a high proportion of incidence and joint deformity with high disease activity. HLJDD is a classical formula from TCM and has been widely used to treat inflammation-related diseases for thousands of years, and recent preclinical studies also demonstrated its effective treatment in RA (Hu et al., 2013). However, its concrete mechanism of the clinical effects remains unclear. Therefore, we investigated the potential mechanisms of the therapeutic effect of HLJDD in RA employing network pharmacology and molecular docking analysis.

In our study, the ingredients-target network of HLJDD was constructed using 102 compounds, and their 189 targets and complicated network demonstrated that most ingredients of the HLJDD could regulate multiple targets. Combined with the PPI network analysis, nine pivotal ingredients were ultimately identified, including quercetin, coptisine, stigmasterol, baicalein, oroxylin A, beta-sitosterol, kaempferol, wogonin, and rutaecarpine. Interestingly, using High Performance Liquid Chromatography (HPLC) analysis, Yuan et al. (2019) detected the contents of 13 ingredients and indirectly supported our findings, including phellodendrine (1), chlorogenic acid (2), magnoflorine (3), geniposide (4), coptisine (5), epiberberine (6), jatrorrhizine (7), berberine (8), palmatine (9), baicalin (10), wogonoside (11), wogonin (12), and oroxylin A (13). Quercetin was a class of flavonol that has unique therapeutic properties, including anti-inflammatory, antioxidant, anticarcinogenic, and antiviral activities (Li et al., 2016).

Wogonin was a natural flavonoid derived from the root extract of Huangqin and had demonstrated potential anti-inflammatory and chondroprotective effects in osteoarthritis through activating the ROS/ERK/Nrf2 signaling pathway (Khan et al., 2017). Baicalein was considered an...
anticarcinogenic compound to inhibit the deterioration of multiple tumors, such as hepatocellular carcinoma and breast cancer (Bie et al., 2017; Yan et al., 2018). Kaempferol has been reported as a potent immunosuppressant that significantly reduced the hypernomic immune response, including autoimmune and chronic inflammation responses (Lin et al., 2011). In this study, the above compounds regulated most RA-related targets and exhibited anti-inflammatory effects.

**FIGURE 4** Functional enrichment analysis of 41 common and 10 hub targets. (A) Top 10 terms of BP, MF, and CC using GO enrichment analysis of 41 common targets. (B) Top 10 KEGG pathways of 41 common targets. (C) The network included detailed targets of these KEGG pathways. (D) The complex botanical drugs-ingredients-hub-targets-disease network of HLJDD in the treatment of RA. The blue nodes represent botanical drugs. The red nodes represent botanical drugs activation ingredients. The orange nodes represent hub targets. The green node represents RA disease.
Furthermore, the molecular docking results exhibited appropriate docking between the crystal structures of target proteins and activated ingredients. Increasing evidence indicated quercetin could inhibit cell apoptosis and reduce the expression levels of inflammatory factors such as IL6 and TNF-α (Wang et al., 2019). In addition, quercetin significantly inhibited MPP9’s expression through NF-kappa B signaling pathway to attenuate cellular proliferation and invasion (Lu et al., 2018). Interestingly, Sung et al. (2012) also demonstrated that quercetin could significantly inhibit unstimulated and IL-1β induced proliferation of rheumatoid synovial fibroblasts (RASFs) and the messenger ribonucleic acid and protein expression of MMP-1, 3, COX-2. Consistent with previous literature, through cell experiments, we successfully validated the inhibitory role of quercetin in the cellular proliferation of MH7A cells in vitro. These results suggested that the therapeutic mechanism of activated ingredients of HLJDD focused on anti-inflammatory and anticancerous biological effects, interpreting the efficacious treatment in RA.

Moreover, 10 hub targets (MPP9, IL1β, IFNG, IL10, ICAM1, CCL2, PTGS2, IL6, CXCL8, and JUN) of HLJDD in treating RA were identified through constructing the PPI network of the 41 common targets from HLJDD and RA. These targets were significantly enriched in the biological process of the cellular inflammatory response to different substances such as organic substance and chemical and cytokine stimulus and associated with IL-17 signaling pathway (CCL2, CXCL10, CXCL8, IFN-γ, IL1β, IL6, JUN, MPP9, PTGS2), TNF signaling pathway (CCL2, CXCL10, ICAM1, IL1β, IL6, JUN, MPP9, PTGS2), rheumatoid arthritis pathway (CCL2, CXCL8, ICAM1, IFNG, IL1β, IL6, JUN), and Toll-like receptor signaling pathway (CXCL10, CXCL8, IL1β, IL6, JUN). Accumulating studies showed that IL-17 and IL-17-producing T helper (Th17) cells played a critical role during the development and progression of RA (Kim et al., 2017) and Lee’s study exhibited that IL-17 upregulated the expression of TLR3 by STAT3 pathways in fibroblast-like synoviocytes of RA (Lee et al., 2014). Tumor necrosis factor (TNF) signaling pathway has been reported to mediate leukocyte recruitment and inflammatory responses, and various anti-TNF inhibitors also acquired prominent potent effects on RA (Yang et al., 2018). In addition, Toll-like receptor (TLRs) and its corresponding downstream signaling pathways, such as Wnt, NF-κB, and MAPK signaling pathways, have been demonstrated in the biological process of synovial inflammation and bone remodeling of RA (Andreakos et al., 2005). Interestingly, low IL-10 and IFN-γ levels were found in RA, and this contradictory finding might be explained by published studies. Seung et al. (Lee et al., 2017) found IFN-γ regulated inflammatory cell death by targeting necroptosis, and IFN-γ deficiency increased the number of Th17 cells and upregulated the expression levels of IL-17 and TNF-α using experimental mice of collagen-induced arthritis (CIA). In addition, an authoritative study declared that IL-10 was also considered a highly promising treatment target for RA attributing to its unique capacity to inhibit cellular immunity and deactivate macrophages through downregulating the production of multiple proinflammatory cytokines, including IL1 and TNF-α (St Clair, 1999). Overall, the results suggested that complicated immune-associated signaling pathways were activated in the progression of RA, and inhibiting associated targets would alleviate the disease activity and prognosis for RA.

### Table 3

| Target genes | PDB ID | Active ingredients | Binding energy (kcal/mol) |
|--------------|--------|---------------------|---------------------------|
| PTGS2        | 5F19   | Quercetin           | -9.6                      |
| PTGS2        | 5F19   | Baicalin            | -9.3                      |
| PTGS2        | 5F19   | Beta-sitosterol     | -9.7                      |
| PTGS2        | 5F19   | Coptisine           | -9.6                      |
| PTGS2        | 5F19   | Stigmasterol        | -9.0                      |
| IL6          | 1IL6   | Quercetin           | -7.4                      |
| IL6          | 1IL6   | Wogonin             | -7.2                      |
| IL6          | 1IL6   | Oroxolin A          | -7.0                      |
| MMP9         | 4H3X   | Quercetin           | -10.7                     |
| MMP9         | 4H3X   | Baicalin            | -10.1                     |
| MMP9         | 4H3X   | Rutacearpine        | -8.6                      |
| CXCL8        | 4DX    | Quercetin           | -6.8                      |
| CXCL8        | 4DX    | Wogonin             | -6.7                      |
| JUN          | 1JUN   | Quercetin           | -5.5                      |
| JUN          | 1JUN   | Beta-sitosterol     | -5.5                      |
| ICAM1        | 1D3L   | Quercetin           | -6.2                      |
| ICAM1        | 1D3L   | Kaempferol          | -5.9                      |
| IFNG         | 1HIg   | Quercetin           | -7.7                      |
| IL1β         | 31Bl   | Quercetin           | -7.5                      |
| IL10         | 1Lk    | Quercetin           | -6.7                      |
| CCL2         | 1DOM   | Quercetin           | -6.1                      |

### Table 3 | Molecular docking results between ligands and core target receptors.
FIGURE 5 | The molecular docking results between active ingredients and hub targets. (A) PTGS2, (B) MMP9, (C) IL6, (D) JUN, (E) ICAM1, (F) CXCL8, (G) CCL2, (H) IL1β, (I) IL10, (J) IFNG. Different molecular rings with different colors represent different active ingredients.
FIGURE 6 | Normalized expression of 10 hub genes in PBMC of RA and control and cell proliferation assay. (A–J) The results showed that the expression levels of IL6, IL1β, CXCL8, MMP9, PTGS2, ICAM1, and CCL2 were increased in RA while the expression levels of IFNG and IL10 were decreased in RA compared with the control. (K,L) The histogram and cellular imaging showed the quercetin’s inhibitory effectiveness was generally increased with the enhancement of quercetin. (M) The drug-concentration curve displayed the IC50 value of quercetin was identified as 38.02 μM in the treatment of RA.
patients (Alwan and Ghali, 2021). Prostaglandin-endoperoxide synthase 2 (PTGS2), also called COX-2, has been recognized as a significant proinflammatory target of RA, and COX-2 inhibitor (NSAIDs) was the preferred anti-inflammatory drug to control pain and stiffness (Crofford, 2013). Notably, Nicole et al. found that the AP-1 Transcription Factor (JUN) could directly activate proinflammatory factor COX-2 in macrophages to promote arthritis in rat models (Hannemann et al., 2017). Summarily, the potential mechanisms of HLJDD in the treatment of RA might be attributed to the following aspects: inhibiting the immune inflammatory response, reducing the release of chemokines, and alleviating the destruction of ECM in the synovial compartment.

However, there still are some limitations in our study. On the one hand, although network pharmacology provides a comprehensive approach based on the combination of pharmacology, biochemics, bioinformatics, and network biology, the data of active ingredients and targets were collectively obtained from multi-public databases, and still selection bias exists. On the other hand, in this study, we just validated querectin’s therapeutic role in inhibiting the proliferation of RA cells in vitro, but the ingredients-targets network and concrete molecular mechanism of HLJDD in the treatment of RA remain to be verified by in-depth in vivo and in vitro studies. Finally, the curative effect of querectin might be non-specific due to artificial synthesis in this experiment and need more validation using extractive compounds in the future.

CONCLUSION

In this study, we successfully screened and obtained the pivotal ingredients of HLJDD in treating RA, and 10 hub targets were identified to construct the botanical drugs-ingredients-targets-disease network. Functional enrichment analysis suggested HLJDD could regulate related targets through associated pathways, including TNF signaling pathway, Toll-like receptor signaling pathway, and IL-17 signaling pathway. Furthermore, based on the molecular docking analysis, important compounds such as querectin, beta-sitosterol, wogonin, kaempferol, coptisine, and stigmasterol exhibited good binding ability with these core proteins. In conclusion, our study systematically expounded the mechanisms and molecular targets of HLJDD in the treatment of RA through the network pharmacology and molecular docking approach. The therapeutic action of HLJDD in RA was mainly by inhibiting the immune inflammatory response, reducing the release of chemokines, and alleviating the destruction of ECM in the synovial compartment.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

CL and JP contributed to the data analysis and drafting of the manuscript. CX contributed to data acquisition, figures presentation, and revision of the manuscript. ZJ and XC contributed to the design of the study. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcell.2021.740266/full#supplementary-material

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