



The pharmacological mechanism of Huashi Baidu Formula for the treatment of COVID-19 by combined network pharmacology and molecular docking

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Background: Huashi Baidu Formula (HSBDF) is a traditional Chinese medicine formula consisting of fourteen parts, which has been proven effective for treating coronavirus disease 2019 (COVID-19) clinically. However, the therapeutic mechanism of the effect of HSBDF on COVID-19 remains unclear.

Methods: The components and action targets of HSBDF were searched in the TCMSP, YaTCM, PubChem, and TargetNet databases. Disease targets related to *ACE2* were screened in single-cell sequence data of colon epithelial cells from other reports. The therapeutic targets of HSBDF for COVID-19 were obtained by integrated analysis, and the protein-protein interaction was analyzed using the STRING database. The Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) processes were analyzed using the OmicsBean and Metascape databases. The communication between networks [component-target (C-T) network, component-target-pathway (C-T-P) network, herb-target (H-T) network, target-pathway (T-P) network, and meridian-tropism (M-T) network] was constructed by Cytoscape software. The Cloud computing molecular docking platform was used to verify the molecular docking.

Results: The obtained 223 active ingredients and 358 targets of HSBDF. The 5,555 COVID-19 disease targets related to *ACE2* were extracted, and 84 compound-disease common targets were found, of which the principal targets included *ACE*, *ESR1*, *ADRA1A*, and *HDAC1*. A total of 3,946 items were seized by GO enrichment analysis, mainly related to metabolism, protein binding, cellular response to the stimulus, and receptor activity. The enriched KEGG pathways screened 46 signaling pathways, including the renin-angiotensin system, the renin secretion, NF-kappa B pathway, the arachidonic acid metabolism, and the AMPK signaling pathway. The molecular docking results showed that the bioactive components of HSBDF have an excellent binding ability with main proteins related to severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2).

Conclusions: HSBDF might act on SARS-CoV-2 through multiple components, targets, and pathways. Here we reveal preliminary results of the mechanism of action of HSBDF on SARS-CoV-2, providing a theoretical basis for future clinical applications.

Keywords: Coronavirus disease 2019 (COVID-19); severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2); Huashi Baidu Formula (HSBDF); network pharmacology; signaling pathway; molecular docking

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Introduction

The 2019 new coronavirus [severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2)] epidemic was first reported in Wuhan, China, in December 2019, and has been declared by the World Health Organization as an urgent public health event of international concern, which evolved into a large-scale epidemic-related pandemic (1). Coronavirus disease 2019 (COVID-19) is caused by SARS-CoV-2, a virus that belongs to the coronavirus of the genus β . The virus has a diameter of 60–140 nm, and the particles are round or oval; the shape is often polymorphic. SARS-CoV-2 cell invasion depends on *TMPRSS2*, *ACE2*, and other main proteins, and its invasion mechanism is similar to that of SARS-CoV (2,3). After the virus invades, the body dysregulates inflammatory factors, resulting in excessive inflammation, which can accelerate the disease progress and produce clinical manifestations such as multiple organ failure (4). Patients with mild disease have symptoms such as fever, fatigue, and dry cough, and severe cases may have dyspnea, respiratory distress syndrome, or septic shock, and there are currently no specific antiviral drugs available in the clinic (1). As of Nov 29, 2020, more than 62.34 million people have been diagnosed with COVID-19, and the number of deaths is caused by the disease is close to 1,460,000 (5).

When the immune system is overactivated due to infection, disease, drugs, and other factors, immune cells will produce large amounts of cytokines, intensifying the inflammation. Many secreted pro-inflammatory factors will cause a positive feedback loop to break through a certain threshold, eventually forming a cytokine storm (6). Patients infected with SARS-CoV-2 with severe COVID-19 have a significant increase in pro-inflammatory cytokines such as IL-6, IL-7, TNF- α , IFN- γ , and their symptoms are characterized by a cytokine storm (7-9). The body produces inflammatory factors after the virus invades, causing an inflammatory overreaction, accelerating the disease and clinical manifestations such as single or multiple organ failure (10,11). In addition to the antiviral treatment and supportive care, it is necessary to monitor for cytokine storm in patients, as early treatment positively affects the reduction of critical conversion rate and reduces the mortality rate (12).

In the treatment of COVID-19, Traditional Chinese medicine (TCM) and humanized monoclonal antibodies play a significant role in reducing the cytokine storm syndrome in COVID-19, including tocilizumab, Lianhuaqingwen, other antibodies targeting inflammatory cytokines, and other TCM prescriptions (13-15).

TCM has achieved good clinical results in preventing and treating new infectious diseases, especially in the prevention and treatment of COVID-19 (16-18). “Three Chinese Patent Medicines, Three Traditional Chinese Medicine Compounds” has been proven effective in the treatment of COVID-19. Huashi Baidu Formula (HSBDF) has been recommended for treating epidemic toxin blocking in the lung, a pattern seen in severe cases of COVID-19. HSBDF consists of 14 herbs, including *Ephedra Herba*, *Amygdalus Communis Vas*, *Gypsum Fibrosum*, *licorice*, *Pogostemon Cablin (Blanco) Benth.*, *Magnolia Officinalis Rehd Et Wils*, *Atractylodes Lancea (Thunb.)Dc.*, *Amomum Tsao-Ko Crevostet*, *Pinelliae rhizoma preparata*, *Poria Cocos(Schw.) Wolf*, *Radix Rhei Et Rhizome*, *Hedysarum Multijugum Maxim.*, *Lepidii Semen Descurainiae Semen*, and *Radix Paeoniae Rubra* (19).

In 75 severe cases treated in Wuhan Jinyintan Hospital, the CT diagnosis showed significant improvement after using HSBDF in lung inflammation and clinical symptoms, and nucleic acid-negative time conversion and hospital stay time were shortened by an average of 3 days (20). In another study, including 124 moderate cases in Street Health Center at Jiangjun Road and 894 mild and moderate cases (452 in the TCM group) in Dongxihu Makeshift Cabin Hospital, showed that there were significant differences in the time to achieve symptoms relief and nucleic acid negative conversion after using HSBDF (21).

Although numerous studies have been carried out on the clinical efficacy, not many studies exist which initially reveal the potential mechanism HSBDF use may have on COVID-19, and the pharmacological mechanism remains unclear. Network pharmacology is a novel technology, and it improves the efficiency of drug discovery based on the component-target-pathway multi-network level. It has been widely used to screen active components, the mechanism of drug action, and the pathogenesis of diseases (22,23). It would be an essential tool for researching complex systems of TCM with the continuous accumulation of disease and

drug-related targets, and the constant mining of chemical components of TCM (24). Molecular docking is a theoretical simulation method to predict its binding mode and affinity based on the characteristics of the receptor and the interaction between the receptor and drug molecules (25). This study aims to investigate the potential therapeutic mechanism of HSBDF on COVID-19 through network pharmacology combined with molecular docking. The flow chart of the research is shown in *Figure 1*.

We present the following article in accordance with the MDAR checklist (available at <http://dx.doi.org/10.21037/apm-20-1759>).

Methods

The Component Database establishment

The components of HSBDF were initially searched in the TCMSP (<http://tcmsp.com/tcmsp.php>) and YaTCM (<http://cadd.pharmacy.nankai.edu.cn/yatcm/home>) databases. Then, using PubChem (<https://pubchem.ncbi.nlm.nih.gov/>), the SMILES structures were acquired, resulting in the Mol2 structure of all compositions, establishing the component database (26,27). The TCMSP and YaTCM databases are TCM databases with information on various herbs, chemicals, and targets, which are beneficial to study TCM scientifically and precisely, and the PubChem database is the central database of chemical research (28).

The component database was established according to the compendial flow chart (*Figure 2*). We initially collected the herbal medicine information, including 14 crude drug medicinal materials. The TCMSP and YaTCM databases were used to obtain the component information of the 14 crude drug medicinal materials. Pharmacokinetic and drug screening models were used to screen the active ingredients from the chemical compositions, including the Obioavail 1.1 system, the Tanimoto coefficient, and Lipinski Rule of Five, and the parameters included $DL \geq 0.18$, $OB \geq 30\%$, Lipinski Rule of Five ≥ 0.75 (29-32). The $T_{(a,b)}$ index was defined as follows:

$$T_{(a,b)} = \frac{a \cdot b}{\|a\|^2 + \|b\|^2 + a \cdot b} \quad [1]$$

a represents the descriptor vector of the tested compound, and b represents the database descriptor vector of drug or drug-like compound.

After collecting the active ingredients, the chemical

structural formula (the Mol2 2D structure, the Mol2 3D structure, the SMILES structure) was obtained from the TCMSP, YaTCM, and PubChem databases, as well as the properties of the active ingredients.

The active components acquirement and ADME screening

The absorption, distribution, metabolism, and excretion (ADME) screening, including the oral bioavailability (OB) and drug-like (DL) properties, was performed in all components to acquire active components from crude drug medicinal materials. The condition parameters were $OB \geq 30\%$ and $DL \geq 0.18$, Lipinski Rule of Five ≥ 0.75 , which effectively screened the active components. The information was normalized using the settings of the OBioavail 1.1 system and the Tanimoto coefficient, which are efficient in collecting the active components (29,31).

Potential targets prediction and crucial targets screening

The prediction of the potential target was carried out using the TargetNet database (<http://targetnet.scbdd.com/>) for netting or predicting potential targets. It constructs QSAR models based on current chemogenomics data to make future predictions (33). We included models with $AUC \geq 0.7$, and the result score of ≥ 0.5 was used for precise and efficient prediction. After gaining the active targets of HSBDF, disease targets of COVID-19 were obtained from other reports, which identified 5,555 genes co-expressed with *ACE2* (34), which has been identified as the crucial cellular receptor for SARS-CoV and SARS-CoV-2. Then we used a Venn diagram to merge and intersect component target genes and COVID-19 disease genes to retrieve crucial target genes of HSBDF on SARS-CoV-2. Finally, the essential target genes database was established, and the genes were annotated and normalized using the Uniprot database (<https://www.uniprot.org/>) (35).

Protein-Protein Interaction (PPI) network construction

The PPI network was constructed through the STRING database (<https://string-db.org/cgi/input.pl>), a database of protein-protein interactions that covers 24,584,628 proteins from 5,090 organisms (36-37). We then visualized the data with the Cytoscape software (38). The Molecular Complex Detection (MCODE) plug-in for critical targets analysis was used with the following setting parameters: Degree Cutoff: 2, Node Score Cutoff: 0.2, K-Core: 2, and Max Depth: 100 (39).

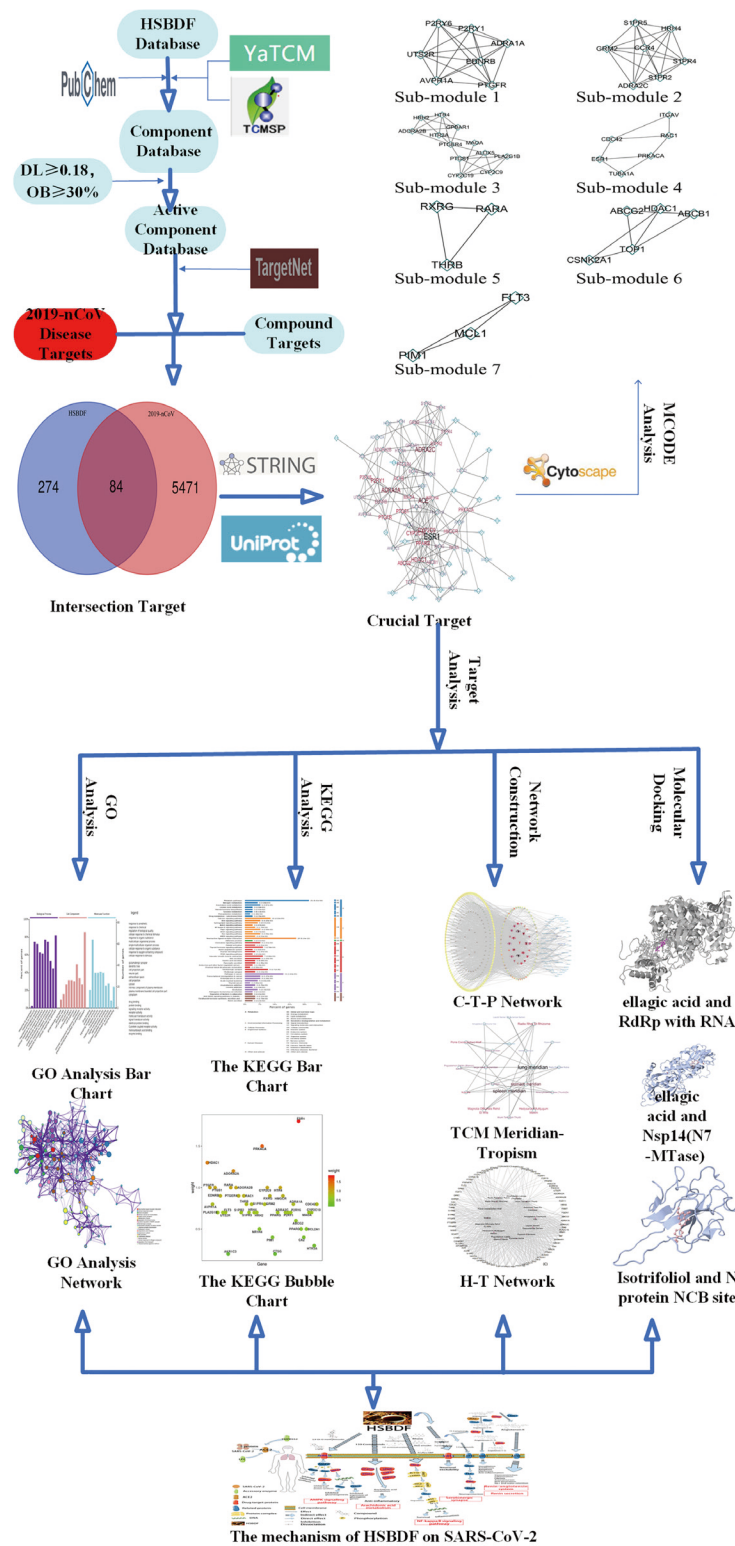


Figure 1 The flowchart of the study process. The flowchart includes the acquisition of key targets, protein protein interaction (PPI) analysis, Molecular Complex Detection (MCODE) analysis, Gene Ontology (GO) analysis, Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis, network construction, molecular docking, and the pharmacological mechanism assessment.

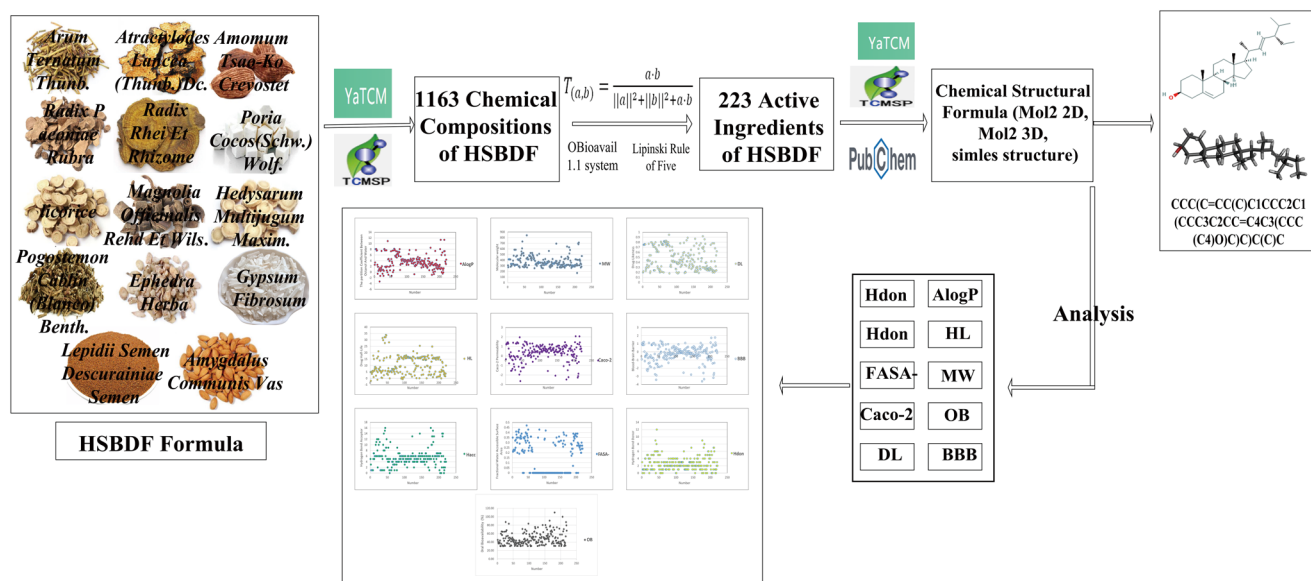


Figure 2 The flow chart of the database establishment. The flowchart explains the collection of crude drug medicinal materials, the acquisition of all ingredients, active ingredients, and the analysis of active compounds properties.

The Gene Ontology (GO) analysis

The key targets were analyzed using the Metascape (<http://metascape.org/gp/index.html>) and OmicsBean (<http://www.omicsbean.cn/>) databases (40). Metascape uses the Benjamini-Hochberg P value correction algorithm and well-adopted hypergeometric test. In Metascape and OmicsBean databases, the investigation was based on fundamental and routine parameters that had a statistically significant difference (P value <0.05). Moreover, Cytoscape and other graphic software were used to draw related charts based on the GO analysis (41).

The Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analysis

A KEGG pathway enrichment was carried out using the Metascape database to find essential targets involved in physiological regulation (42). Differential expression gene and protein lists were used to design the KEGG analysis experiment, to allow for the detection of essential target proteins enriched in associated biological pathways.

Network construction

Networks were constructed using the Cytoscape software, including a component-target (C-T) network, component-

target-pathway (C-T-P) network, herb-target (H-T) network, target-pathway (T-P) network, and meridian-tropism (M-T) network. A topology analysis was performed in each network, including information of the node, edge, degree, betweenness, and closeness centrality, which aids in elucidating the nature of the networks (38).

Molecular docking

Molecular docking was executed using the COVID-19 Docking Server (<https://ncov.schanglab.org.cn/index.php>), which aims to test the correlation between active compositions and essential proteins related to COVID-19. The molecules were acquired by topology analysis from the C-T-P network, and the target proteins that are essential for the treatment of COVID-19 are listed in *Tables 3,4* (43). There were three groups of drugs used for the molecular docking: a group of HSBDF compounds, a control group of chemical drugs, and a negative control group. We included five molecules for the group of HSBDF compounds: baicalein, ellagic acid, genkwanin, isotrifoliol, and rhein. We included six compounds for the control group of chemical drugs: Lopinavir, Ritonavir, Remdesivir, darunavir, Arbidol, and Chloroquine, which might have a potential effect on COVID-19 (44). Water was used to verify the correctness of the database in the negative control group. The D3Targets-2019-nCoV database (<https://www.d3pharma>.

com/D3Targets-2019-nCoV/index.php) was also used to verify the binding energy between bioactive components and other main proteins related to SARS-CoV-2; detailed information is shown in the results table (45).

The study conformed to the provision of the Declaration of Helsinki (as revised in 2013).

Statistical analysis

The quantitative data were analyzed with Student's *t*-test using SPSS (25.0 version, SPSS Inc., Chicago, IL, USA). *P* value <0.05 was considered a statistically significant value.

Results

Identification compositions of HSBDF

The composition of HSBDF was acquired from the TCMSP and YaTCM databases, which included 223

bioactive compositions, and the basic information is listed in *Table 1* and *Figure 3*. All medicinal chemistry information is listed in the supplementary information table (<https://cdn.amegroups.cn/static/public/atm-20-1759-1.xlsx>). We calculated the average of each chemical information (MW: 374.61, AlogP: 3.15, Hdon: 3.86, Hacc: 5.95, OB: 45.37%, Caco-2: 0.08, BBB: -0.50, DL: 0.49, FASA-: 0.29, HL: 10.48). Various compositions included glycoside compounds, ketones, steroid acid compounds, and other compounds rich in herbs, representing the diversity of compositions, revealing multi-compositions of HSBDF. Among those compositions, isoflavones and cynotoxin had a high OB score; (-)-medicocarpin and 9,10-dimethoxypterocarpan-3-O-β-D-glucoside had a high level of DL score.

Crucial targets of HSBDF on COVID-19

The 358 potential target genes were acquired using the TargetNet database, and the disease target genes related

Table 1 The compendious compositions of Huashi Baidu Formula (HSBDF)

No.	Name	No.	Name	No.	Name
M001	2-hydroxyisoxypyl-3-hydroxy-7-isopentene-2,3-dihydrobenzofuran-5-carboxylic	M076	Poricoic acid C	M151	Quercetin der.
M002	3β-acetoxyatractylone	M077	Trametenolic acid	M152	Semilicoisoflavone B
M003	Beta-daucosterol_qt	M078	Hederagenin	M153	Shinpterocarpin
M004	Beta-sitosterol 3-O-glucoside_qt	M079	(-)-medicocarpin	M154	Sigmoidin-B
M005	Daucosterin_qt	M080	(2R)-7-hydroxy-2-(4-hydroxyphenyl)chroman-4-one	M155	Vestitol
M006	Daucosterol_qt	M081	(2S)-2-[4-hydroxy-3-(3-methylbut-2-enyl)phenyl]-8,8-dimethyl-2,3-dihydropyrano[2,3-f]chromen-4-one	M156	Xambioona
M007	NSC63551	M082	(2S)-6-(2,4-dihydroxyphenyl)-2-(2-hydroxypropan-2-yl)-4-methoxy-2,3-dihydrofuro[3,2-g]chromen-7-one	M157	Calycosin
M008	Stigmasterol 3-O-beta-D-glucopyranoside_qt	M083	(2S)-7-hydroxy-2-(4-hydroxyphenyl)-8-(3-methylbut-2-enyl)chroman-4-one	M158	Formononetin
M009	Wogonin	M084	(E)-1-(2,4-dihydroxyphenyl)-3-(2,2-dimethylchromen-6-yl)prop-2-en-1-one	M159	Jaranol
M010	(1S,2S,4R)-trans-2-hydroxy-1,8-cineole-B-D-glucopyranoside	M085	(E)-3-[3,4-dihydroxy-5-(3-methylbut-2-enyl)phenyl]-1-(2,4-dihydroxyphenyl)prop-2-en-1-one	M160	Mairin
M011	(2R,3R)-4-methoxyl-distylin	M086	1,3-dihydroxy-8,9-dimethoxy-6-benzofurano[3,2-c]chromenone	M161	Naringenin

Table 1 (continued)

Table 1 (continued)

No.	Name	No.	Name	No.	Name
M012	1-o-beta-d-glucopyranosyl-8-o-benzoylpaeonisuffrone_qt	M087	1,3-dihydroxy-9-methoxy-6-benzofurano[3,2-c]chromenone	M162	Isorhamnetin
M013	1-o-beta-d-glucopyranosylpaeonisuffrone_qt	M088	18 α -hydroxyglycyrrhetic acid	M163	Quercetin
M014	4-ethyl-paeoniflorin_qt	M089	1-Methoxyphaseollidin	M164	Kaempferol
M015	4-o-methyl-paeoniflorin_qt	M090	2-(3,4-dihydroxyphenyl)-5,7-dihydroxy-6-(3-methylbut-2-enyl)chromone	M165	Glabridin
M016	8-debenzoylpaeonidanin	M091	2-[(3R)-8,8-dimethyl-3,4-dihydro-2H-pyrano[6,5-f]chromen-3-yl]-5-methoxyphenol	M166	Glycyrol
M017	9-ethyl-neo-paeoniaflorin A_qt	M092	3-(2,4-dihydroxyphenyl)-8-(1,1-dimethylprop-2-enyl)-7-hydroxy-5-methoxy-coumarin	M167	Licochalcone B
M018	Albiflorin	M093	3-(3,4-dihydroxyphenyl)-5,7-dihydroxy-8-(3-methylbut-2-enyl)chromone	M168	Phaseol
M019	Albiflorin_qt	M094	3,22-Dihydroxy-11-oxo-delta(12)-oleanene-27-alpha-methoxycarbonyl-29-oic acid	M169	Liquiritin
M020	Benzoyl paeoniflorin	M095	3'-Hydroxy-4'-O-Methylglabridin	M170	Eucalyptol
M021	Ellagic acid	M096	3'-Methoxyglabridin	M171	Neohesperidin
M022	Ethyl oleate (NF)	M097	5,7-dihydroxy-3-(4-methoxyphenyl)-8-(3-methylbut-2-enyl)chromone	M172	(3R)-3-(2-hydroxy-3,4-dimethoxyphenyl)chroman-7-ol
M023	Evofolinb	M098	6-prenylated eriodictyol	M173	(3S,8S,9S,10R,13R,14S,17R)-10,13-dimethyl-17-[(2R,5S)-5-propan-2-yloctan-2-yl]-2,3,4,7,8,9,11,12,14,15,16,17-dodecahydro-1H-cyclopenta[a]phenanthren-3-ol
M024	Isobenzoylpaeoniflorin	M099	7,2',4'-trihydroxy-5-methoxy-3-arylcoumarin	M174	(6aR,11aR)-9,10-dimethoxy-6a,11a-dihydro-6H-benzofurano[3,2-c]chromen-3-ol
M025	Lactiflorin	M100	7-Acetoxy-2-methylisoflavone	M175	1,7-Dihydroxy-3,9-dimethoxy pterocarpene
M026	Paeoniflorigenone	M101	7-Methoxy-2-methyl isoflavone	M176	3,9-di-O-methylnissolin
M027	Paeoniflorigenone	M102	8-(6-hydroxy-2-benzofuranyl)-2,2-dimethyl-5-chromenol	M177	5'-hydroxyisomuronulatol-2',5'-di-O-glucoside
M028	Paeoniflorin	M103	8-prenylated eriodictyol	M178	7-O-methylisomucronulatol
M029	Paeoniflorin_qt	M104	dehydroglyasperins C	M179	9,10-dimethoxypterocarpan-3-O- β -D-glucoside
M030	Stigmast-7-en-3-ol	M105	DFV	M180	Bifendate
M031	Sitosterol	M106	euchrenone	M181	FA

Table 1 (continued)

Table 1 (continued)

No.	Name	No.	Name	No.	Name
M032	Campest-5-en-3beta-ol	M107	Eurycarpin A	M182	Isoflavanone
M033	(+)-catechin	M108	gadelaicid acid	M183	Isomucronulatol-7,2'-di-O-glucosiole
M034	Spinasterol	M109	Gancaonin A	M184	3,23-dihydroxy-12-oleanen-28-oic acid
M035	(-)-catechin	M110	Gancaonin B	M185	5-hydroxy-7,4'-dimethoxyflavanon
M036	Aloe-emodin	M111	Gancaonin G	M186	Acanthoside B
M037	Daucosterol_qt	M112	Gancaonin H	M187	Diop
M038	Emodin-1-O-beta-D-glucopyranoside	M113	Glabranin	M188	Irisolidone
M039	EUPATIN	M114	Glabrene	M189	Pachypodol
M040	Gallic acid-3-O-(6'-O-galloyl)-glucoside	M115	Glabrone	M190	Patchoulan 1,12-diol
M041	Mutatochrome	M116	Glepidotin A	M191	Phenanthrone
M042	Palmidin A	M117	Glepidotin B	M192	Quercetin 7-O-β-D-glucoside
M043	Physciondiglucoside	M118	Glyasperin B	M193	Genkwanin
M044	Procyanidin B-5,3'-O-gallate	M119	Glyasperin C	M194	(+)-Leucocyanidin
M045	Rhein	M120	Glyasperin F	M195	Delphinidin
M046	Sennoside D_qt	M121	Glyasperins M	M196	Diosmetin
M047	Sennoside E_qt	M122	Glycyrin	M197	Eriodictyol
M048	Torachrysone-8-O-beta-D-(6'-oxayl)-glucoside	M123	Glycyroside	M198	Herbacetin
M049	Toralactone	M124	Glycyrrhiza flavonol A	M199	Leucopelargonidin
M050	(3S,6S)-3-(benzyl)-6-(4-hydroxybenzyl)piperazine-2,5-quinone	M125	Glypallichalcone	M200	Luteolin
M051	10,13-eicosadienoic	M126	Glyzaglabrin	M201	Pectolarigenin
M052	12,13-epoxy-9-hydroxynonadeca-7,10-dienoic acid	M127	HMO	M202	Poriferast-5-en-3beta-ol
M053	beta-D-ribofuranoside, xanthine-9	M128	Icos-5-enoic acid	M203	Resivit
M054	Cavidine	M129	Inermine	M204	Supraene
M055	Coniferin	M130	Inflacoumarin A	M205	Taxifolin
M056	Cycloartenol	M131	Isoglycyrol	M206	Truflex OBP
M057	Glycyrrhizic acid ammonium salt	M132	Isolicoflavonol	M207	Calcium sulfate
M058	Baicalein	M133	Isotrifoliol	M208	Cynotoxin
M059	Baicalin	M134	Kanzonol F	M209	Dihomolinolenic acid
M060	Stigmasterol	M135	Kanzonols W	M210	Erysimoside
M061	24-ethylcholest-4-en-3-one	M136	Licoagrocarpin	M211	Evobioside
M062	Beta-sitosterol	M137	Licoagroisoflavone	M212	K-strophanthoside
M063	Gondoic acid	M138	licoalcone a	M213	K-strophanthoside_qt

Table 1 (continued)

Table 1 (continued)

No.	Name	No.	Name	No.	Name
M064	(2R)-2-[(3S,5R,10S,13R,14R,16R,17R)-3,16-dihydroxy-4,4,10,13,14-pentamethyl-2,3,5,6,12,15,16,17-octahydro-1H-cyclopenta[a]phenanthren-17-yl]-5-isopropylhex-5-enoic acid	M139	Licochalcone G	M214	11,14-eicosadienoic acid
M065	(2R)-2-[(3S,5R,10S,13R,14R,16R,17R)-3,16-dihydroxy-4,4,10,13,14-pentamethyl-2,3,5,6,12,15,16,17-octahydro-1H-cyclopenta[a]phenanthren-17-yl]-6-methylhept-5-enoic acid	M140	Licocoumarone	M215	(6Z,10E,14E,18E)-2,6,10,15,19,23-hexamethyltetracos-2,6,10,14,18,22-hexaene
M066	(2R)-2-[(5R,10S,13R,14R,16R,17R)-16-hydroxy-3-keto-4,4,10,13,14-pentamethyl-1,2,5,6,12,15,16,17-octahydrocyclopenta[a]phenanthren-17-yl]-5-isopropylhex-5-enoic acid	M141	Licoisoflavanone	M216	CLR
M067	3beta-Hydroxy-24-methylene-8-lanostene-21-oic acid	M142	Licoisoflavone	M217	Diisooctyl succinate
M068	7,9(11)-dehydropachymic acid	M143	Licoisoflavone B	M218	Estrone
M069	Cerevisterol	M144	Licopyranocoumarin	M219	I-SPD
M070	Dehydroeburicoic acid	M145	Licorice glycoside E	M220	Machiline
M071	Ergosta-7,22E-dien-3beta-ol	M146	Licoricone	M221	Ziziphin_qt
M072	Ergosterol peroxide	M147	Lupiwighteone	M222	Ent-Epicatechin
M073	Pachymic acid	M148	Medicarpin	M223	(4E,6E)-1,7-bis(4-hydroxyphenyl)hepta-4,6-dien-3-one
M074	Poricoic acid A	M149	Odoratin		
M075	Poricoic acid B	M150	Phaseolinisoflavan		

to *ACE2* included 5,555 target genes built from the attachment in published reports. Finally, 84 crossed target genes were obtained (Figure 4), representing potentially crucial target genes for HSBDF on SARS-CoV-2. The essential target genes database was established after the annotation and normalization of the Unprior database (Table 2), and most proteins found in the intersection targets were related to metabolism, sex hormones, and immunological stress.

PPI network of crucial target proteins

The 84 crucial target proteins were analyzed in the STRING database, and the PPI network was constructed using Cytoscape software (Figure 5). The network included 75

nodes, 211 edges, and 5,550 shortest paths. The essential proteins in the PPI network were *ESR1*, *ADRA1A*, *ADRA2C*, *HDAC1*, and *ACE*, which were considered the vital proteins for the treatment of HSBDF on COVID-19. The network was divided into seven submodules by the MCODE analysis (Figure 6), which revealed the particular interactions.

The GO biological analysis

The GO results were acquired after 84 essential proteins were analyzed by the Metascape and OmicsBean databases, and the overall histogram of GO biological enrichment was obtained using the OmicsBean database (Figure 7). For the biological processes, significant roles included the cellular response to the stimulus, the multicellular organismal process, which



Figure 3 The scatter diagram of component properties, including oral bioavailability (OB), molecular weight (MW), hydrogen bond donor (Hdon), hydrogen bond acceptor (Hacc), blood-brain barrier (BBB), drug-likeness (DL), drug half-life (HL), Caco-2 permeability (Caco-2), fractional water accessible surface area (FASA-), and the partition coefficient between octanol and water (AlogP). Each note represents the compound properties.

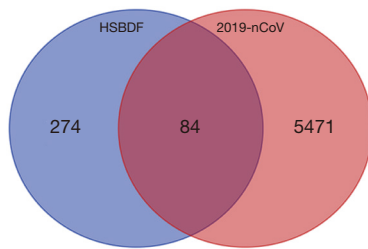


Figure 4 The Venn diagram comparing the composition and disease target genes. The blue section represents 358 targets of Huashi Baidu Formula (HSBDF), the red section represents 5,555 targets of the disease, and the intersection part represents 84 crucial target genes.

indicates the blocking of SARS-CoV-2 and the diversity action of the HSBDF components. The considerable role of cell components included the cytoplasm and cytosol, which could potentially be the therapeutic mechanism of HSBDF on COVID-19 since it might target the synthetic metabolism, primarily related to the protein, fat, and carbohydrate metabolism. For molecular functions, a significant role was found for protein binding, reflecting the mechanisms related to proteins consistent with the biological processes and the cell components involved. The specific bubble diagram and gene weight diagram are shown in *Figure 8*.

Based on the Metascape database, the interaction network of the GO biological process was acquired

Table 2 The special information on crucial target proteins

Number	Target name	Gene symbol	Uniprot ID	Degree	Betweenness	Closeness centrality
1	AdenosinATP-dependent translocase ABCB1	<i>ABCB1</i>	P08183	7	0.01986246	0.37755102
2	Broad substrate specificity ATP-binding cassette transporter ABCG2	<i>ABCG2</i>	Q9UNQ0	9	0.03675141	0.38743455
3	Angiotensin-converting enzyme	<i>ACE</i>	P12821	13	0.18839999	0.48051948
4	Acetylcholinesterase	<i>ACHE</i>	P22303	8	0.06378896	0.4180791
5	Adenosine deaminase	<i>ADA</i>	P00813	5	0.03303039	0.32888889
6	Adenosine receptor A2a	<i>ADORA2A</i>	P29274	4	0.01066671	0.30204082
7	Adenosine receptor A2b	<i>ADORA2B</i>	P29275	7	0.01580731	0.31759657
8	Alpha-1A adrenergic receptor	<i>ADRA1A</i>	P35348	12	0.04248098	0.39361702
9	Alpha-2C adrenergic receptor	<i>ADRA2C</i>	P18825	13	0.11221669	0.40883978
10	Aldo-keto reductase family 1 member C3	<i>AKR1C3</i>	P42330	5	0.00871493	0.37373737
11	Arachidonate 5-lipoxygenase	<i>ALOX5</i>	P09917	6	0.01039743	0.3490566
12	Aminopeptidase N	<i>ANPEP</i>	P15144	4	0.02742846	0.35576923
13	DNA dC->dU-editing enzyme APOBEC-3A	<i>APOBEC3A</i>	P31941	0	0	0
14	Vasopressin V1a receptor	<i>AVPR1A</i>	P37288	6	0	0.33035714
15	Cholinesterase	<i>BCHE</i>	P06276	6	0.01551649	0.39572193
16	Bcl-2-related protein A1	<i>BCL2A1</i>	Q16548	2	0.00037	0.26714801
17	Carbonic anhydrase 1	<i>CA1</i>	P00915	0	0	0
18	Carbonic anhydrase 13	<i>CA13</i>	Q8N1Q1	0	0	0
19	Carbonic anhydrase 2	<i>CA2</i>	P00918	0	0	0
20	Carbonic anhydrase 4	<i>CA4</i>	P22748	2	0.00432285	0.34101382
21	Carbonic anhydrase 6	<i>CA6</i>	P23280	0	0	0
22	C-C chemokine receptor type 4	<i>CCR4</i>	P51679	6	0	0.31355932
23	Cell division control protein 42 homolog	<i>CDC42</i>	P60953	5	0.062072	0.3627451

Table 2 (continued)

Table 2 (continued)

Number	Target name	Gene symbol	Uniprot ID	Degree	Betweenness	Closeness centrality
24	Cocaine esterase	<i>CES2</i>	O00748	5	0.00554243	0.33183857
25	Casein kinase II subunit alpha	<i>CSNK2A1</i>	P68400	2	0	0.30081301
26	Cathepsin G	<i>CTSG</i>	P08311	2	0.00969708	0.32888889
27	Cytochrome P450 2C19	<i>CYP2C19</i>	P33261	10	0.03088508	0.39784946
28	Cytochrome P450 2C9	<i>CYP2C9</i>	P11712	11	0.03338479	0.41111111
29	Dipeptidyl peptidase 2	<i>DPP7</i>	Q9UHL4	2	0.000527	0.26523297
30	Endothelin receptor type B	<i>EDNRB</i>	P24530	7	0.01108262	0.37
31	Estrogen receptor	<i>ESR1</i>	P03372	18	0.26203358	0.48366013
32	Squalene synthase	<i>FDFT1</i>	P37268	2	0	0.296
33	Receptor-type tyrosine-protein kinase FLT3	<i>FLT3</i>	P36888	4	0.0043992	0.30962343
34	Glutamate carboxypeptidase 2	<i>FOLH1</i>	Q04609	0	0	0
35	G-protein coupled bile acid receptor 1	<i>GPBAR1</i>	Q8TDU6	6	0.03345795	0.35576923
36	Metabotropic glutamate receptor 2	<i>GRM2</i>	Q14416	7	0.01265554	0.32599119
37	Histone deacetylase 1	<i>HDAC1</i>	Q13547	11	0.06573624	0.41340782
38	3-hydroxy-3-methylglutaryl-coenzyme A reductase	<i>HMGCR</i>	P04035	8	0.10118612	0.41340782
39	Hepatocyte nuclear factor 4-alpha	<i>HNF4A</i>	P41235	5	0.00461644	0.38341969
40	Histamine H2 receptor	<i>HRH2</i>	P25021	4	0	0.296
41	Histamine H4 receptor	<i>HRH4</i>	Q9H3N8	6	0	0.31355932
42	Corticosteroid 11-beta-dehydrogenase isozyme 1	<i>HSD11B1</i>	P28845	1	0	0.29482072
43	Estradiol 17-beta-dehydrogenase 2	<i>HSD17B2</i>	P37059	2	0	0.33333333
44	5-hydroxytryptamine receptor 3A	<i>HTR3A</i>	P46098	6	0.01348139	0.3627451
45	5-hydroxytryptamine receptor 4	<i>HTR4</i>	Q13639	5	0.00524066	0.31623932
46	Integrin alpha-V	<i>ITGAV</i>	P06756	2	0	0.27819549
47	Potassium voltage-gated channel subfamily A member 3	<i>KCNA3</i>	P22001	1	0	0.22981366
48	Amine oxidase [flavin-containing] A	<i>MAOA</i>	P21397	8	0.041508	0.41111111
49	Induced myeloid leukemia cell differentiation protein Mcl-1	<i>MCL1</i>	Q07820	5	0.01972533	0.34101382
50	Monoglyceride lipase	<i>MGLL</i>	Q99685	2	0.000444	0.30081301
51	Macrophage migration inhibitory factor	<i>MIF</i>	P14174	0	0	0
52	Bile acid receptor	<i>NR1H4</i>	Q96R11	8	0.074357	0.42528736
53	P2Y purinoceptor 1	<i>P2RY1</i>	P47900	10	0.03601648	0.37
54	P2Y purinoceptor 6	<i>P2RY6</i>	Q15077	8	0.0131273	0.3490566
55	Poly [ADP-ribose] polymerase 1	<i>PARP1</i>	P09874	4	0.00226999	0.35238095

Table 2 (continued)

Table 2 (continued)

Number	Target name	Gene symbol	Uniprot ID	Degree	Betweenness	Closeness centrality
56	cAMP-specific 3',5'-cyclic phosphodiesterase 4B	<i>PDE4B</i>	Q07343	1	0	0.28682171
57	Phosphatidylinositol 4,5-bisphosphate 3-kinase catalytic subunit gamma isoform	<i>PIK3CG</i>	P48736	2	0.00277659	0.27715356
58	Serine/threonine-protein kinase pim-1	<i>PIM1</i>	P11309	4	0.00564785	0.29959514
59	Phospholipase A2	<i>PLA2G1B</i>	P04054	5	0.00237198	0.36453202
60	Phenylethanolamine N-methyltransferase	<i>PNMT</i>	P11086	4	0.01301743	0.38541667
61	Peroxisome proliferator-activated receptor delta	<i>PPARD</i>	Q03181	2	0	0.30081301
62	Peroxisome proliferator-activated receptor gamma	<i>PPARG</i>	P37231	9	0.07798613	0.41573034
63	cAMP-dependent protein kinase catalytic subunit alpha	<i>PRKACA</i>	P17612	8	0.10792044	0.4
64	Protein kinase C beta type	<i>PRKCB</i>	P05771	2	0.00739673	0.29718876
65	Presenilin-1	<i>PSEN1</i>	P49768	5	0.05491323	0.3681592
66	Gamma-secretase subunit PEN-2	<i>PSENE1</i>	Q9NZ42	1	0	0.27007299
67	Prostaglandin E2 receptor EP4 subtype	<i>PTGER4</i>	P35408	7	0.03122439	0.34418605
68	Prostaglandin F2-alpha receptor	<i>PTGFR</i>	P43088	9	0.06060018	0.4
69	Prostaglandin G/H synthase 1	<i>PTGS1</i>	P23219	9	0.04383119	0.38541667
70	Tyrosine-protein phosphatase non-receptor type 22	<i>PTPN22</i>	Q9Y2R2	0	0	0
71	Tyrosine-protein phosphatase non-receptor type 7	<i>PTPN7</i>	P35236	1	0	0.28682171
72	Ras-related C3 botulinum toxin substrate 1	<i>RAC1</i>	P63000	3	0.00918002	0.31223629
73	Retinoic acid receptor alpha	<i>RARA</i>	P10276	7	0.02999583	0.37755102
74	Retinoic acid receptor RXR-gamma	<i>RXRG</i>	P48443	6	0.01528434	0.36453202
75	Sphingosine 1-phosphate receptor 2	<i>S1PR2</i>	O95136	8	0.03432741	0.3681592
76	Sphingosine 1-phosphate receptor 4	<i>S1PR4</i>	O95977	7	0.01772187	0.34418605
77	Sphingosine 1-phosphate receptor 5	<i>S1PR5</i>	Q9H228	6	0	0.31355932
78	Sigma non-opioid intracellular receptor 1	<i>SIGMAR1</i>	Q99720	3	0.02702703	0.29718876
79	Sodium/glucose cotransporter 1	<i>SLC5A1</i>	P13866	0	0	0
80	Thyroid hormone receptor beta	<i>THRB</i>	P10828	4	0.00100465	0.3507109
81	DNA topoisomerase 1	<i>TOP1</i>	P11387	7	0.0148477	0.3681592
82	Tubulin alpha-1A chain	<i>TUBA1A</i>	Q71U36	3	0.01416577	0.34741784
83	Urotensin-2 receptor	<i>UTS2R</i>	Q9UKP6	6	0	0.33035714
84	Xanthine dehydrogenase/oxidase	<i>XDH</i>	P47989	1	0	0.24832215

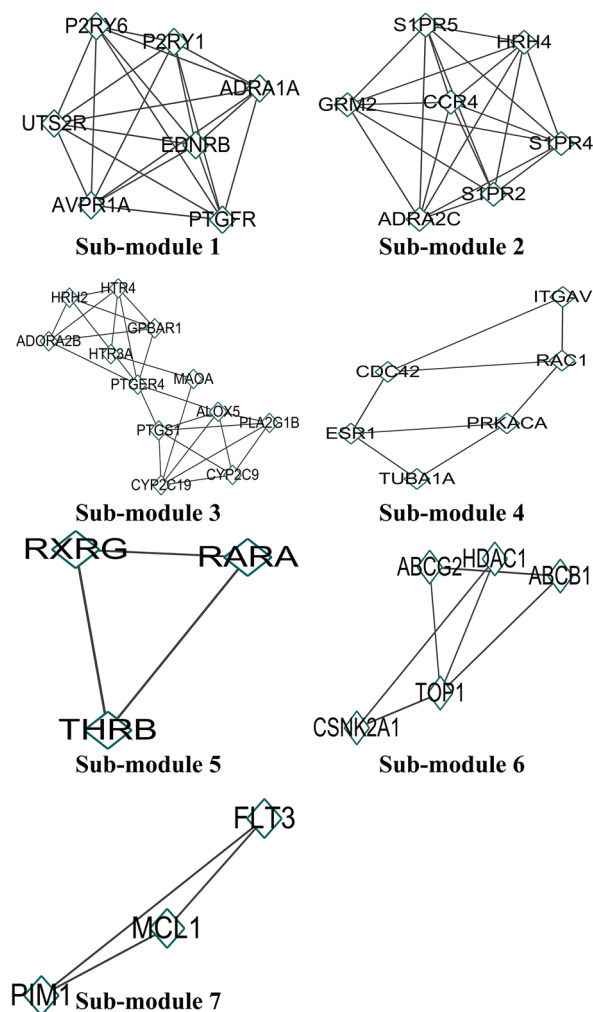


Figure 6 The submodule of the protein-protein interaction (PPI) network by using the Molecular Complex Detection (MCODE). Each part reflects a submodule, each node represents proteins, and the edge represents the interaction of the specific interaction. The submodule 1–7 consists of 7, 7, 12, 6, 3, 5, and 3 nodes, respectively.

Network construction

The constructed networks are shown in *Figure 14*. The C-T-P network (*Figure 14A*) reflects the relationship between components, targets, and pathways. It includes 355 nodes, 1,575 edges, and 125,670 shortest paths, suggesting the crucial pathways in the use of HSBDF to affect SARS-CoV-2. The C-T network (*Figure 14B*) included 298 nodes, 1,575 edges, and 62,250 shortest paths, showing the relationship between components and targets without pathways and explains the mechanism from different perspectives. The H-T network (*Figure 14C*) describes the targets of 14 parts, which is beneficial for studying the

treatment mechanism, and 93 nodes, 517 edges, and 8,556 shortest paths were involved in this network. The T-P network (*Figure 14D*) showed the relationship between targets and pathways. We found that *PRKCB*, *PRKACA*, and *CDC42* are crucial targets, and the pathways associated with cancer, metabolic pathways, and neuroactive ligand-receptor interactions are essential pathways for the T-P network. Moreover, the T-P network included 125 nodes, 248 edges, and 15,550 shortest paths. The meridian-tropism network (*Figure 14E*) was significant for the TCM study, we found that the lung meridian, the spleen meridian, and the stomach meridian were potential target sites of HSBDF for the treatment of COVID-19, and there were 20 nodes,

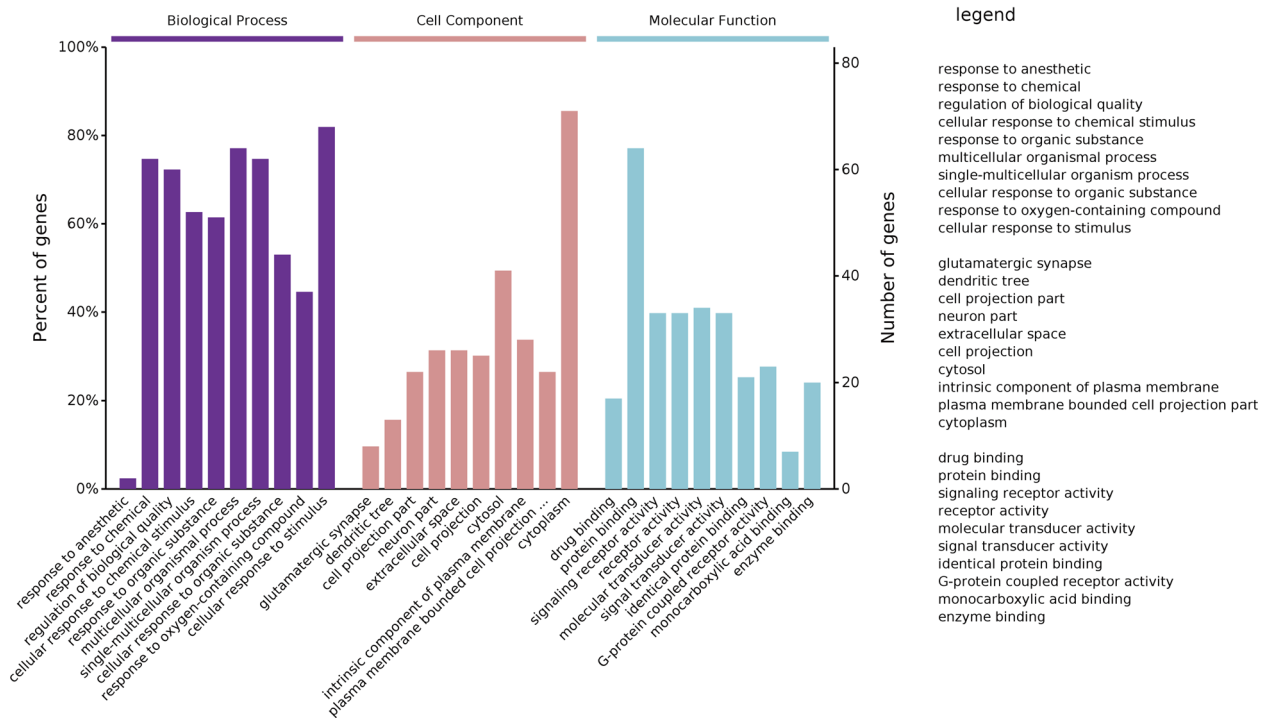


Figure 7 The overall histogram of the Gene Ontology (GO) biological analysis. The color represents different GO processes: the purple section describes the biological process, the pink section describes the cell component, and the blue section describes the molecular function. The length of the bar chart represents the percentage of genes in the corresponding section, P value <0.05.

38 edges, and 380 shortest paths covered in this meridian-tropism (M-T) network.

Discussion

TCM plays a significant role in the treatment of COVID-19, as most of the more than 50,000 discharged patients in China were treated with TCM as of Mar 15. The proportion of TCM applications in Wuhan city, Hubei province, and China was 89.10%, 91.64%, and 92.41%, respectively (19,21,46,47). HSBDF was gradually implemented in the treatment process of the first batch of Chinese medicine medical teams in Jinyintan Hospital and Dongxihu square cabin hospital (21,48). The use of HSBDF for the treatment of COVID-19 was based on multiple components, targets, and pathways that might significantly shorten the duration of the nucleic acid transfer, the average time of hospitalization, improve clinical symptoms, and improve findings at the physical and chemical examination and lung CT improvement (21,48). The experimental results showed that the viral load of lung tissue could be reduced by 30%, and in mice infected with SARS-CoV-2,

the inflammation of the lung can be improved significantly with this drug (20).

Evidence-based medicine and systematic reviews show that the combination of TCM and standard care can significantly improve the treatment outcome in COVID-19 cases, including the overall treatment efficacy, the rate of symptom disappearance, and the improvement of other clinical symptoms (20,49). TCM has been found to have a positive effect on symptoms in a study among 732 adults in seven study groups, and no significant adverse events were recorded in the analysis (50). Recent evidence suggests that the remarkable effect of the combination of TCM and standard care reveals the clinical effect of Chinese medicine in the treatment of COVID-19 in a study including 855 patients in seven study groups (51). Existing research recognizes the critical role TCM played in treating COVID-19 in a study including 2,275 enrolled patients in 18 study groups, and improvements in clinical parameters, such as clinical cure rate, lung CT, inflammatory biomarkers, and other parameters were found (52). Recently, investigators have examined the effects of TCM on COVID-19 in a meta-analysis including 1,474 patients in

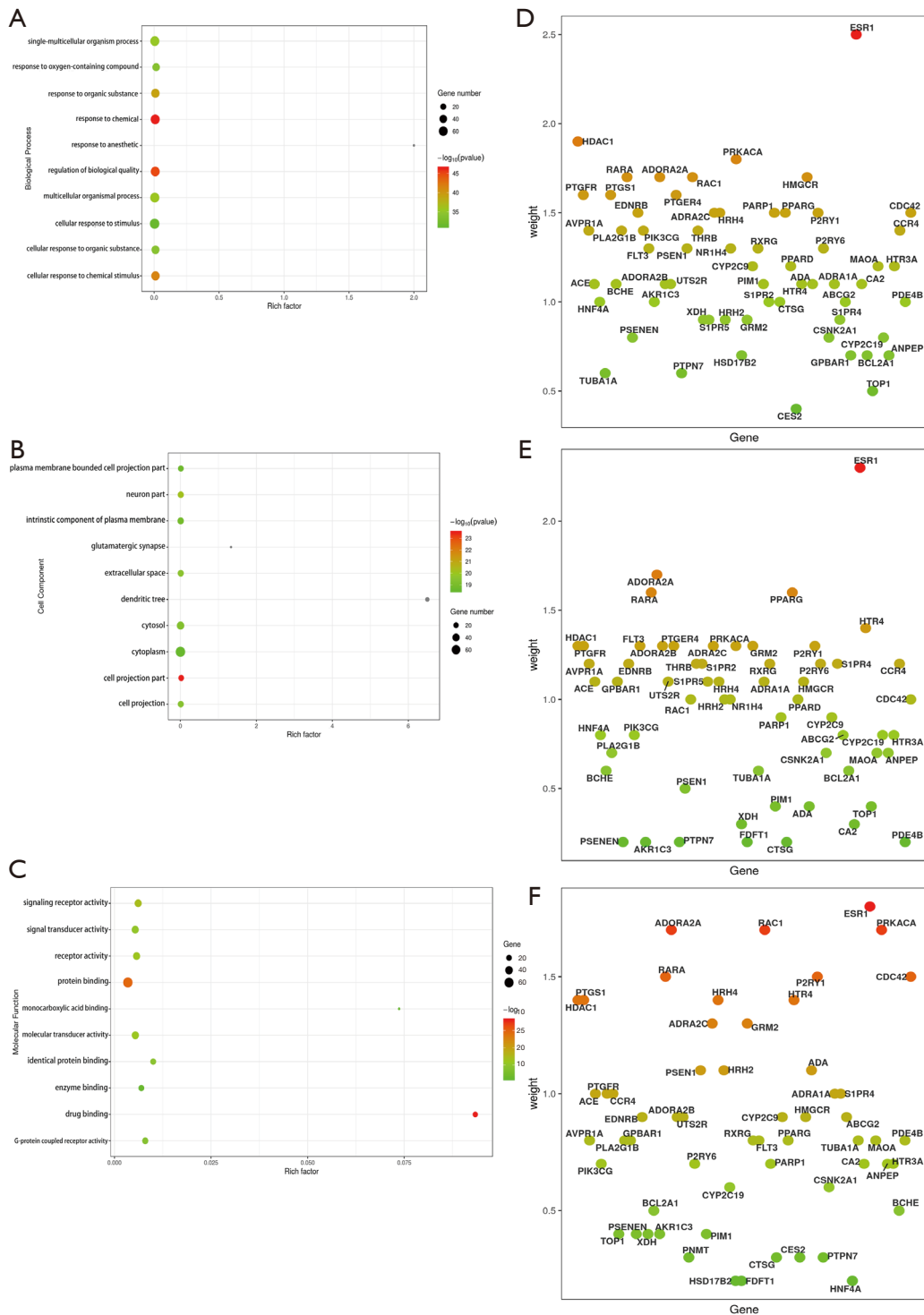


Figure 8 The bubble chart and gene weight chart of the Gene Ontology (GO) enrichment analysis. The bubble chart of the biological process (A), cell component (B), and molecular function (C). The gene weight chart of the biological process (D), cell component (E), and molecular function (F). (A,B,C) The different enrichment of differentially expressed genes in the corresponding section; (D,E,F) the weight of differentially expressed genes in the corresponding section.

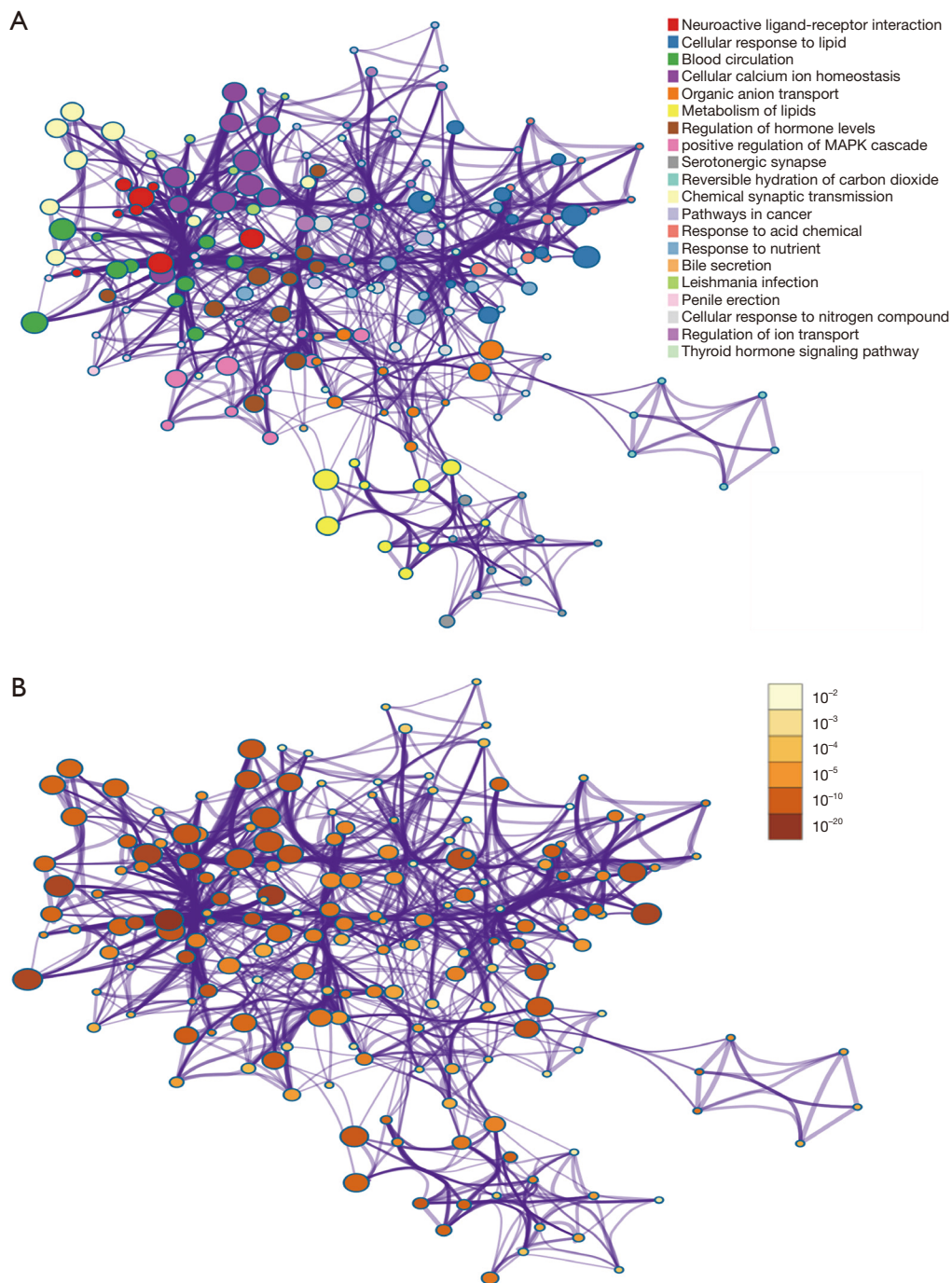


Figure 9 The interaction network of the Gene Ontology (GO) biological enrichment analysis of differentially expressed genes. (A) The network by the criticality. Each node represents the GO processes and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways. The edge represents the relationship of the GO processes and KEGG pathways. Different colors represent different biological processes, the darker the node color, the more critical the biological process. The network includes 180 nodes, 2,549 edges, and 32,220 shortest paths. (B) The network by statistical properties. The depth of the color represents the P value. Different shades of color represent different statistical properties, the darker the node color, the higher the statistical significance of the biological process. The network includes 180 nodes, 2,549 edges, and 32,220 shortest paths.

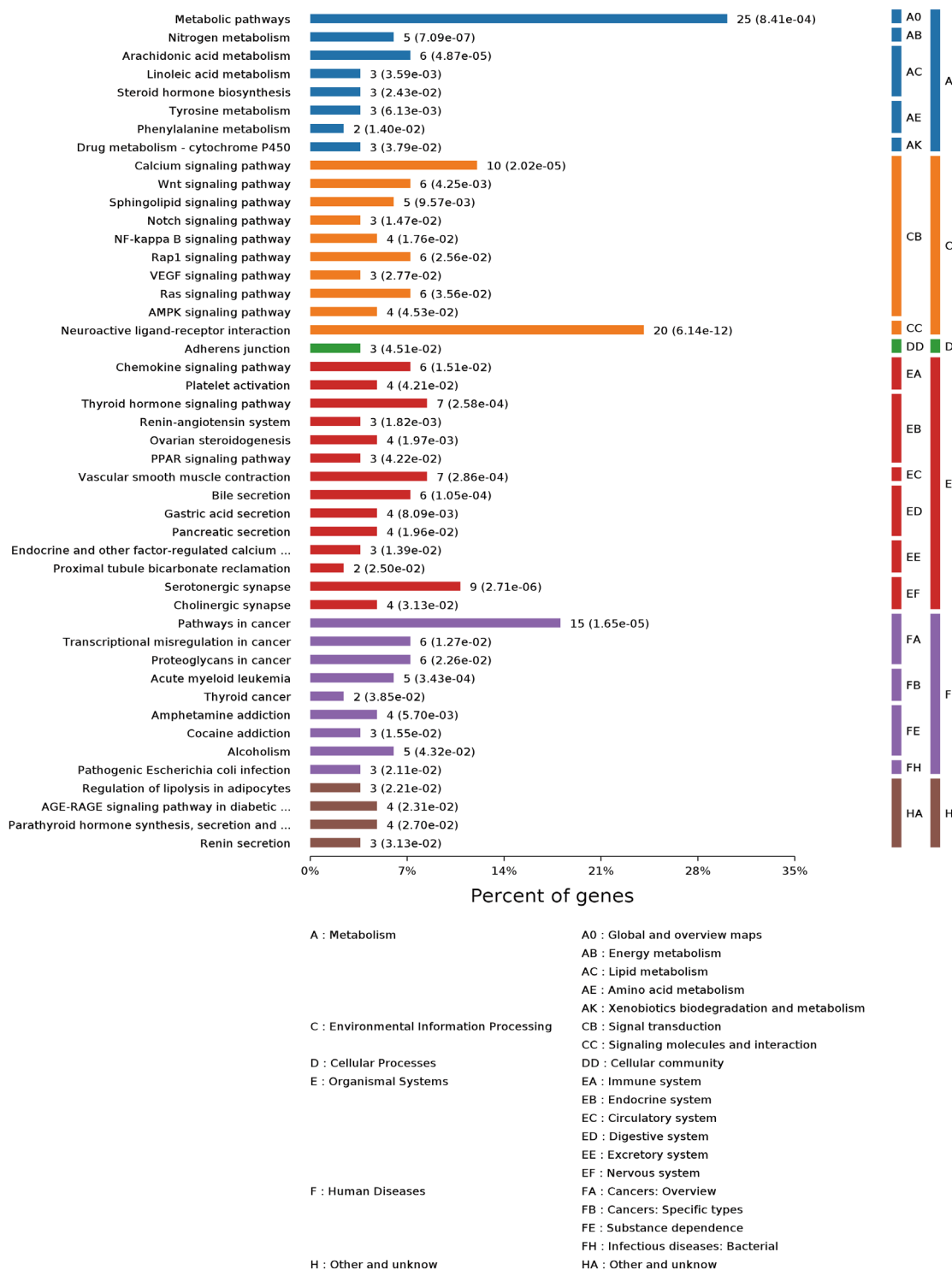


Figure 10 Histogram of the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways associated with the effect of Huashi Baidu Formula (HSBDF) on severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). Different colors represent different pathway types, the bar reflects the specific biological pathways, and the length of the bar represents the percentage of differential genes in the pathway, and the content in brackets represents the statistical significance of pathway enrichment for each pathway, P value <0.05.

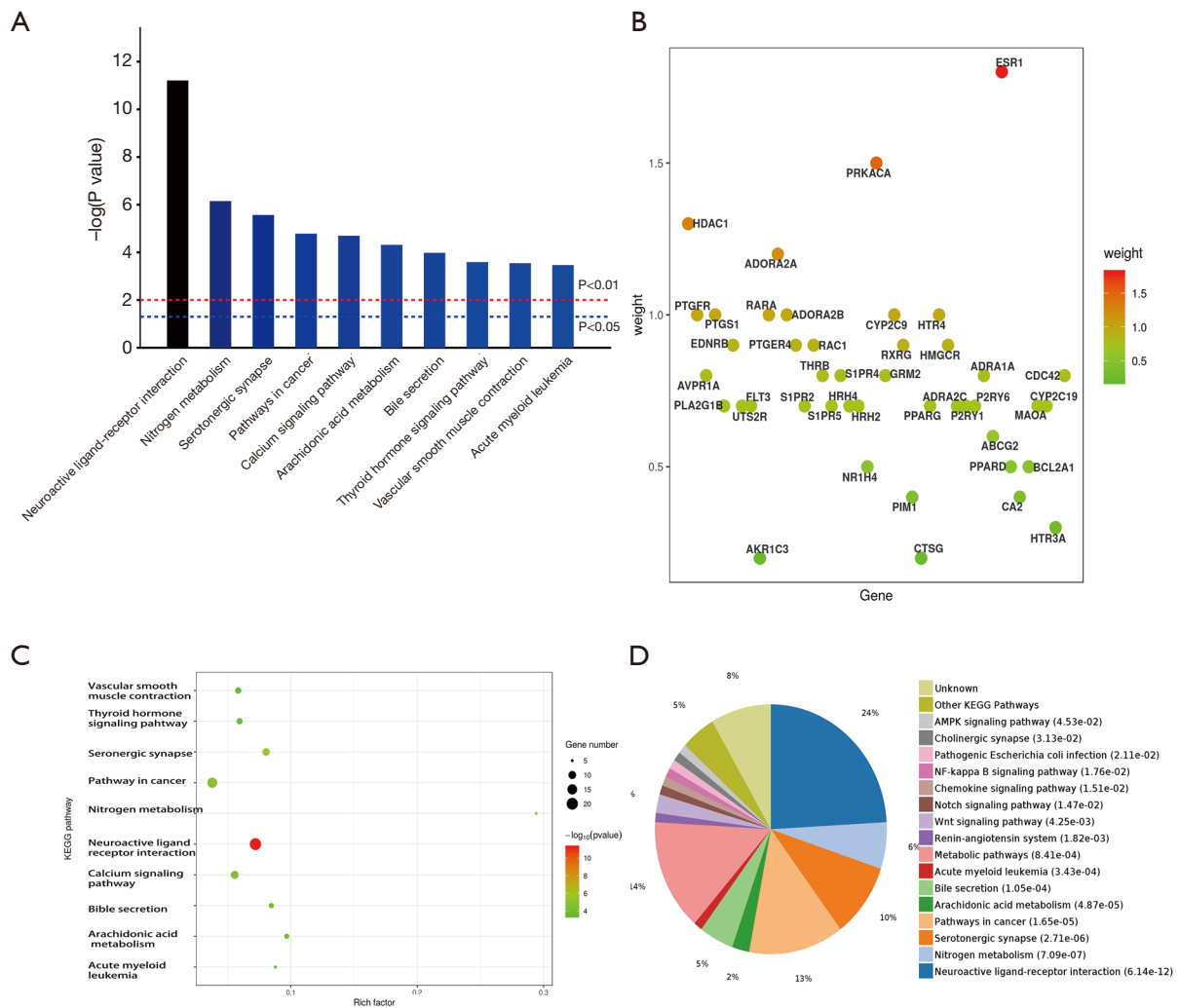


Figure 11 The related graphs of the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analysis. (A) The sector diagram depicting the $-\log(P\text{-value})$ of crucial pathways, the length of the bar reflects the statistical significance of specific KEGG pathways, the higher the length, the more convincing the statistical significance; (B) the gene weight of the KEGG pathway enrichment, different color shades represent the weight of differentially expressed genes; (C) bubble chart of crucial pathways, the color and size of the node represent the number of genes in its specific pathway and its statistical significance; (D) pie chart depicting the P value level and criticality of biological pathways, the higher proportion degree, the more critical the pathway.

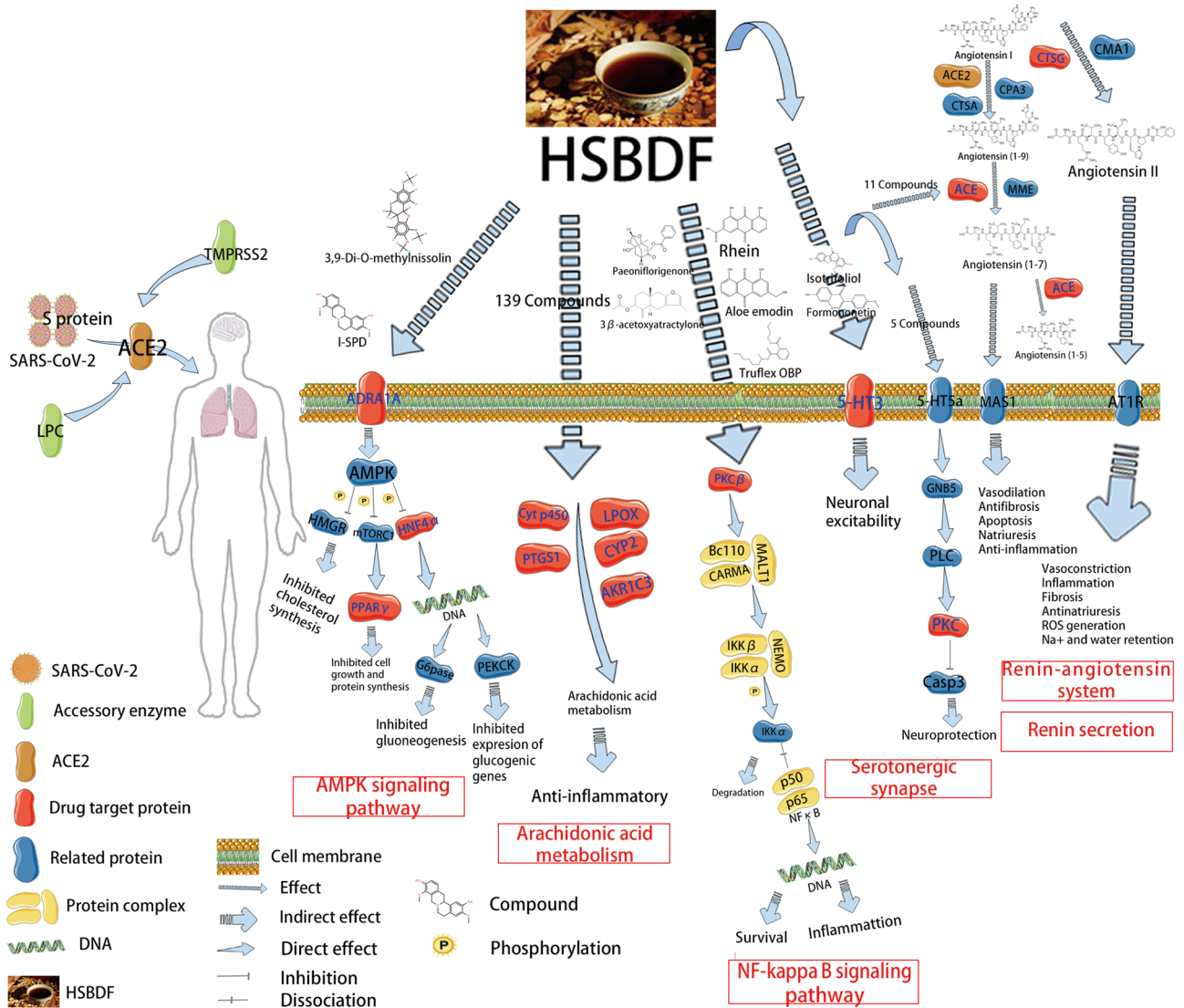


Figure 12 The potential mechanism diagram of the effect of Huashi Baidu Formula (HSBDF) on severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), involving potential therapeutic pathways of HSBDF: adenosine 5'-monophosphate-activated protein kinase (AMPK) signaling pathway, arachidonic acid metabolism, nuclear factor kappa-B (NF-kappa B) signaling pathway, serotonergic synapse, renin-angiotensin system, and renin secretion. The annotation is shown in the figure.

Table 3 The molecule docking results of molecules and proteins (kcal/mol)

Name	Molecular formula	MW	Main Protease	Papain-like protease	RdRp with RNA	RdRp without RNA	Helicase NCB site	Nsp14 (ExoN)	Nsp14 (N7-MTase)	Nsp15	Nsp16	N protein NCB site	E protein (ion channel)
Baicalein	C15H10O5	270.24	-7.80	-8.80	-8.20	-7.10	-6.60	-7.50	-8.70	-6.30	-7.90	-8.20	-7.30
Ellagic acid	C14H6O8	302.19	-7.40	-8.50	-9.60	-8.30	-7.00	-7.60	-9.10	-6.50	-7.80	-8.60	-6.80
Genkwanin	C16H12O5	284.26	-7.10	-8.30	-8.20	-7.20	-6.20	-7.50	-8.30	-6.40	-7.90	-7.90	-7.20
Isotrifoliol	C16H10O6	298.25	-7.90	-8.70	-8.60	-7.90	-6.70	-7.80	-8.80	-6.20	-7.60	-8.90	-8.90
Rhein	C15H8O6	284.22	-7.50	-8.80	-8.70	-7.50	-6.80	-8.00	-9.10	-6.30	-8.00	-8.50	-7.30
Lopinavir	C37H48N4O5	628.80	-7.70	-10.80	-8.50	-8.00	-7.40	-8.30	-9.80	-7.20	-8.80	-7.50	-9.50
Ritonavir	C37H48N6O5S2	720.90	-8.80	-11.70	-11.00	-10.10	-6.90	-9.50	-9.20	-7.10	-8.30	-8.30	-9.80
Remdesivir	C27H35N6O8P	602.60	-7.70	-10.20	-9.50	-7.50	-7.20	-7.40	-9.10	-7.10	-8.70	-7.00	-7.40
Darunavir	C27H37N3O7S	547.70	-8.10	-9.20	-8.60	-7.40	-6.90	-8.00	-9.50	-6.60	-7.90	-7.20	-7.70
Arbidol	C22H25BN2O3S	477.40	-6.60	-8.80	-7.90	-6.10	-6.10	-6.30	-7.60	-5.10	-6.60	-5.80	-6.90
Chloroquine	C18H26ClN3	319.90	-6.20	-7.10	-7.00	-5.60	-5.70	-6.20	-7.70	-5.30	-6.60	-6.00	-6.50
Water	H ₂ O	18.02	-1.70	-1.60	-1.90	-1.70	-1.60	-1.70	-1.50	-1.90	-1.90	-1.50	-1.10

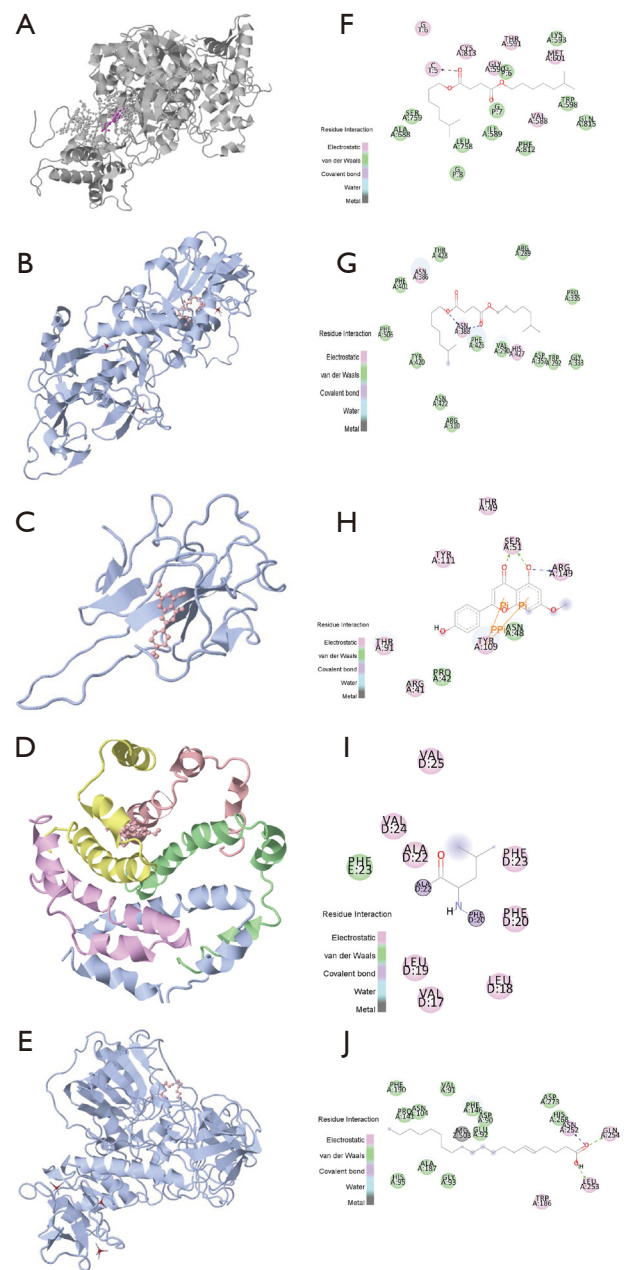
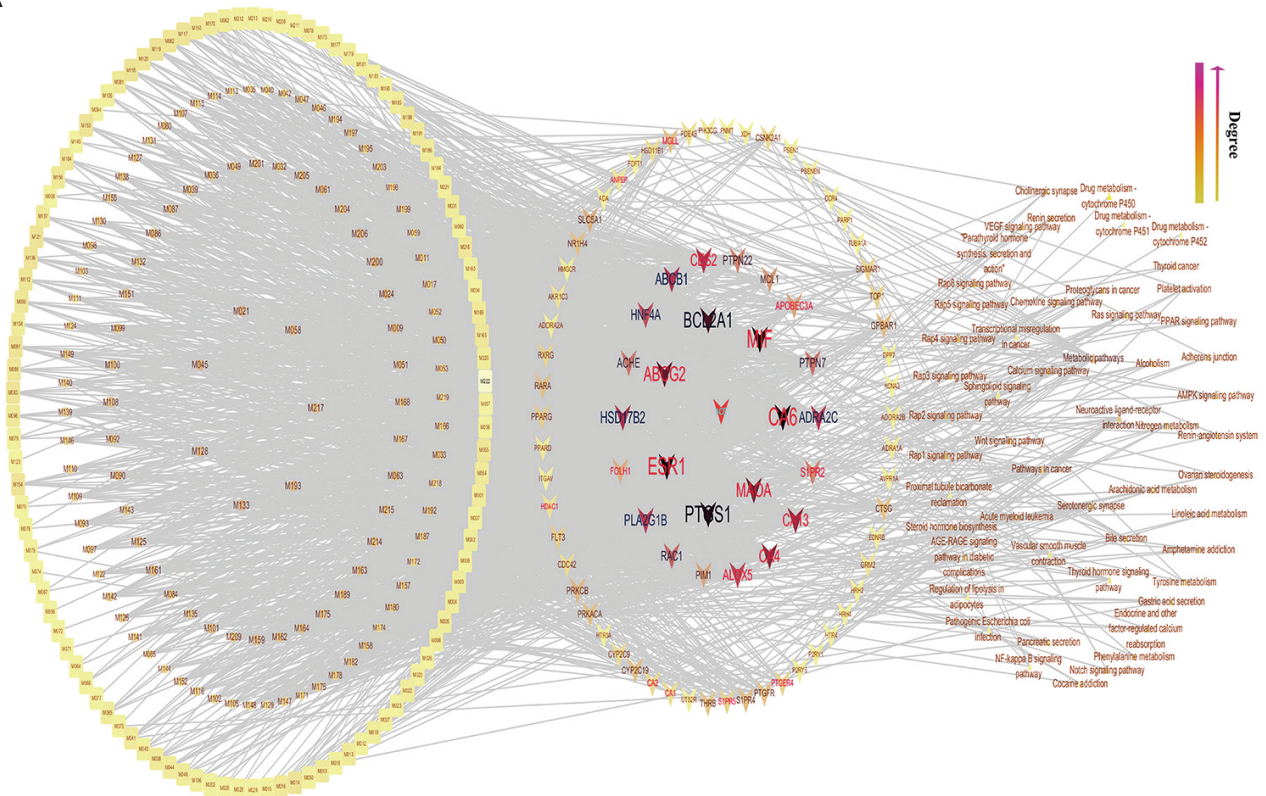
**Figure 13** The visualization of the molecular docking, different parts represents different molecular docking. (A,F) Ellagic acid and RNA-dependent RNA polymerase (RdRp with RNA); (B,G) ellagic acid and Guanine-N7 methyltransferase (Nsp14); (C,H) Isotrifoliol and nucleocapsid protein NCB cite (N protein NCB site); (D,I) Isotrifoliol and envelope proteins (E protein); (E,J) rhein and Guanine-N7 methyltransferase (Nsp14). (A,B,C,D,E) represent the 3D binding diagrams of active compounds and crucial proteins, (F,G,H,I,J) represent the 2D binding diagrams of active compounds and crucial proteins, the circular patterns represent the amino acid residue, and a different color represents variable intermolecular interactions.

Table 4 The molecule docking results of molecules and proteins (kcal/mol)

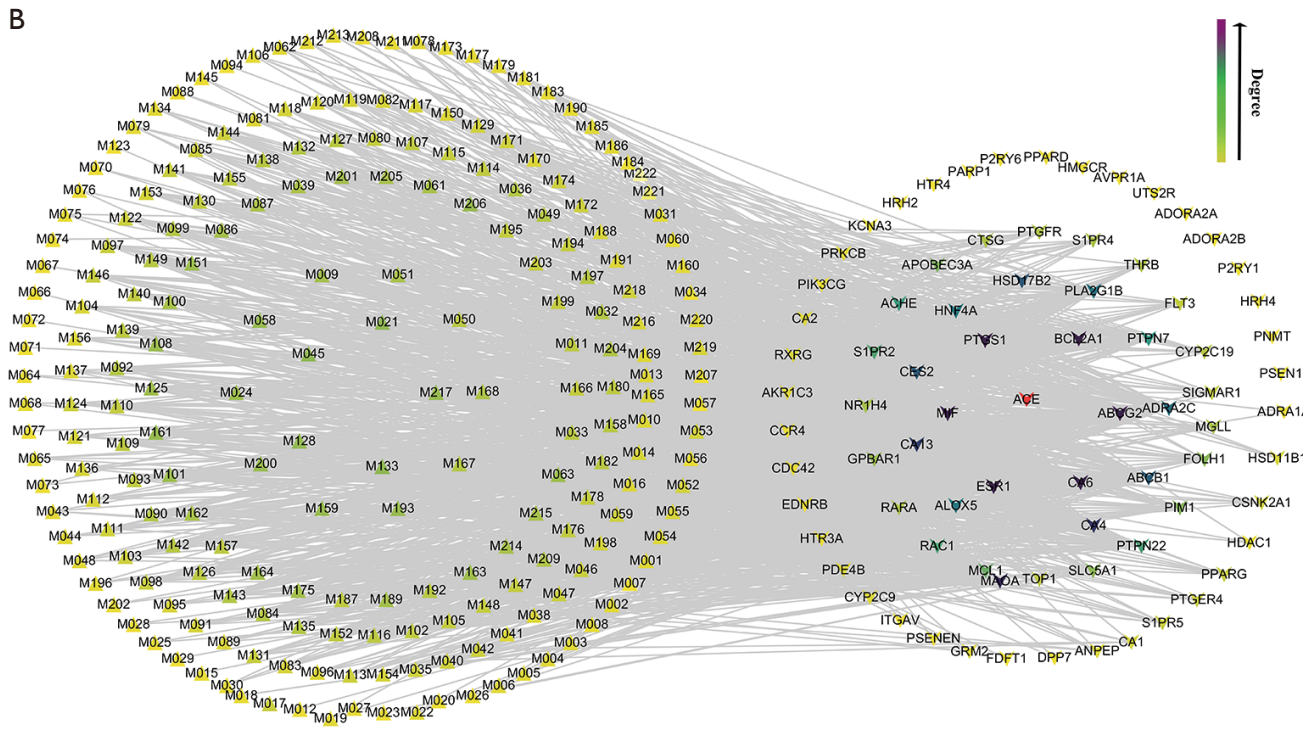
Name	Molecular formula	MW	ACE2	CD209L	CTSL	DHODH	ADAM17 (+Zn)	ADAM17	SARS-CoV-2	TMPRSS2	Mpro (Dimer)	3CLpro/3CLpro/ Mpro (monomer)	S protein (open)	S protein (close)	S protein-S2 subunit	Spike protein-heptad repeat 1
Baicalein	C15H10O5	270.25	-8.41	-7.38	-8.39	-10.97	-8.61	-8.57	-8.67	-8.91	-8.60	-8.36	-9.49	-9.39	-8.12	-7.77
Beta-sitosterol	C29H50O	414.79	-8.15	-6.14	-7.80	-9.23	-7.65	-7.54	-8.31	-8.43	-8.47	-7.61	-8.27	-8.08	-7.66	-7.50
Calycosin	C16H12O5	284.28	-7.51	-6.15	-7.37	-10.83	-8.91	-8.82	-7.41	-8.05	-8.62	-8.00	-8.53	-8.88	-7.83	-6.86
Catechin	C15H14O6	290.29	-9.06	-7.59	-8.54	-10.44	-9.73	-9.66	-8.88	-8.92	-8.44	-9.14	-9.76	-10.02	-8.74	-7.56
Ellagic acid	C14H6O8	302.20	-8.31	-6.94	-7.74	-11.34	-8.23	-8.17	-8.91	-9.11	-9.57	-8.03	-10.70	-10.15	-8.55	-7.32
Formononetin	C16H12O4	268.28	-7.09	-5.88	-7.51	-10.38	-8.31	-8.29	-7.42	-7.40	-8.05	-7.74	-8.18	-8.52	-7.61	-6.59
Genkwanin	C16H12O5	284.26	-7.88	-6.50	-7.97	-10.44	-8.70	-8.59	-8.55	-8.13	-8.31	-7.47	-8.65	-9.05	-7.98	-7.70
Glycyrol	C21H18O6	366.37	-8.92	-6.85	-8.47	-11.01	-7.81	-7.71	-9.26	-8.79	-9.59	-7.69	-10.48	-9.11	-8.40	-7.60
Isorhamnetin	C16H12O7	316.26	-8.58	-7.58	-8.71	-10.52	-8.68	-8.56	-8.15	-8.43	-8.60	-8.16	-9.53	-9.58	-8.38	-7.40
Isotrifolol	C16H10O6	298.26	-8.50	-6.40	-8.19	-11.15	-7.71	-7.53	-8.18	-8.25	-8.84	-7.61	-9.58	-9.38	-7.79	-6.82
Kaempferol	C15H10O6	286.25	-9.02	-7.76	-8.30	-11.00	-8.98	-8.91	-8.29	-8.58	-8.77	-7.95	-9.63	-9.48	-8.21	-8.03
Licochalcone A	C21H22O4	338.40	-8.32	-6.55	-7.52	-9.54	-8.23	-8.15	-7.88	-7.53	-8.44	-7.47	-8.56	-8.31	-7.68	-6.88
Luteolin	C15H10O6	286.23	-8.91	-7.44	-8.57	-11.53	-9.42	-9.35	-8.84	-8.65	-8.86	-8.54	-9.96	-9.69	-8.57	-7.56
Naringenin	C15H12O5	272.25	-8.35	-7.09	-8.05	-10.95	-8.91	-8.82	-8.52	-8.12	-8.34	-7.89	-9.28	-9.33	-8.52	-7.53
Quercetin	C15H10O7	302.24	-8.65	-7.80	-8.62	-11.23	-9.74	-9.71	-8.95	-9.03	-8.88	-8.38	-10.26	-10.16	-8.68	-7.80
Rhein	C15H8O6	284.23	-8.41	-6.45	-8.32	-11.21	-7.85	-7.66	-8.30	-8.49	-8.83	-7.59	-9.53	-9.48	-8.11	-7.20
Stigmasterol	C29H48O	412.69	-8.75	-6.89	-6.69	-9.46	-7.98	-7.94	-8.57	-8.76	-8.55	-7.65	-8.83	-8.62	-8.29	-7.46
Wogonin	C16H12O5	284.26	-7.56	-6.16	-7.57	-10.65	-8.51	-8.25	-7.42	-8.06	-7.78	-7.39	-9.13	-9.18	-7.75	-8.00
Arbidol	C22H25BrN2O3S	477.40	-6.98	-5.55	-6.89	-7.45	-6.83	-6.31	-7.00	-7.75	-7.27	-6.62	-7.32	-7.09	-6.58	-7.48
Chloroquine	C18H26ClN3	319.90	-6.12	-5.17	-6.15	-8.40	-5.81	-6.09	-6.23	-5.91	-6.55	-5.81	-6.80	-6.68	-5.43	-5.68
Darunavir	C27H37N3O7S	547.70	-8.24	-6.24	-6.98	-10.01	-7.79	-8.49	-7.38	-8.63	-9.00	-7.42	-9.52	-8.47	-7.71	-7.54
Lopinavir	C37H48N4O5	628.80	-9.05	-7.12	-7.34	-8.77	-8.95	-8.80	-9.30	-8.60	-9.39	-7.65	-9.87	-9.08	-7.64	-7.54
Remdesivir	C27H35N6O8P	602.60	-8.44	-7.29	-7.73	-9.08	-7.63	-7.78	-7.65	-8.43	-9.49	-7.42	-8.99	-8.83	-8.42	-7.16
Ritonavir	C37H48N6O5S2	720.90	-9.48	-7.25	-7.02	-9.82	-9.77	-8.92	-9.39	-8.30	-10.09	-7.04	-9.42	-9.07	-7.48	-7.65

ACE2, angiotensin converting enzyme 2; CD209L, C type lectin domain family 4 member M; CTSL, Cathepsin L; DHODH, dihydroorotate dehydrogenase; ADAM17 (+Zn), disintegrin and metalloproteinase domain-containing protein 17 +Zn state; ADAM17, disintegrin and metalloproteinase domain-containing protein 17; SARS-CoV-2 chimeric RBD+ACE2, SARS-CoV-2 chimeric receptor-binding domain+Angiotensin converting enzyme 2; TMPRSS2, transmembrane protease serine 2; 3CLpro/Mpro(Dimer), 3C-like proteinase Dimer State; 3CLpro/Mpro(monomer), 3C-like proteinase Monomer State; S protein (Open), Spike protein Open State; S protein (Close), Spike protein Close State; S protein-S2 subunit, Spike protein S2 subunit; S-HR1, Spike protein-heptad repeat 1.

A



B



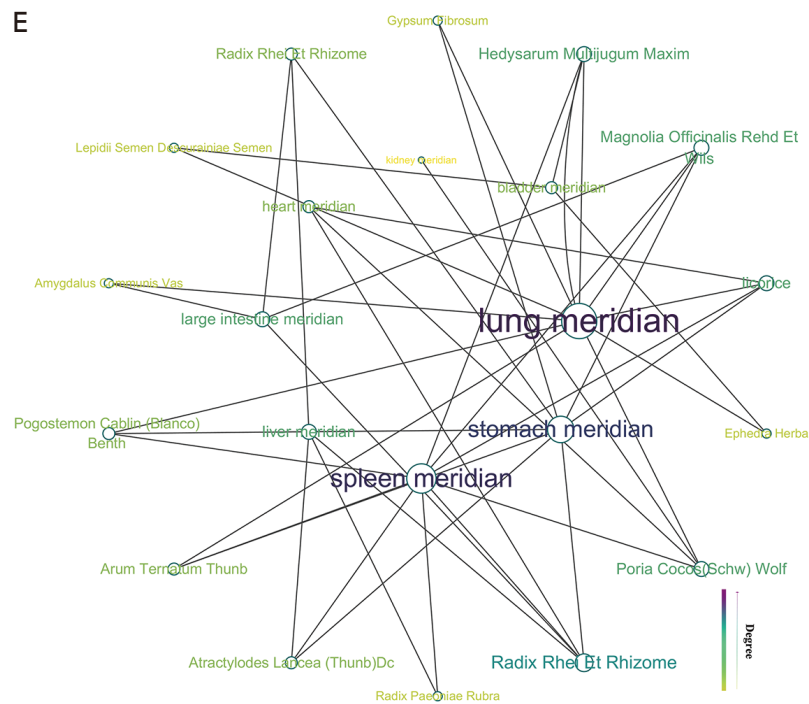


Figure 14 The network information. The component-target-pathway (C-T-P) network (A), rectangle: active ingredients, arrow: crucial targets, triangle: biological pathways; the component-target (C-T) network (B), triangle: active ingredients, arrow: crucial targets; the herb-target (H-T) network (C), triangle: crude drug medicinal materials, circle: crucial targets; the target-pathway (T-P) network (D), triangle: crucial targets, biological pathways, the darker the node, the more critical it is; the meridian-tropism (M-T) network (E), circle: crude drug medicinal materials, organ meridian. The depth and size of the node color reflect the criticality of the node, the darker the color and the larger the size of the node, the greater the criticality of the node.

19 study groups. Combining Chinese and Western medicine to treat COVID-19 may effectively control symptoms and reduce disease progression, which is seen reflected by the improvement of symptoms, reduction of hospital stay, and conversion rate of critical illness of TCM (53). These findings have significant implications for the understanding of the clinical effect of HSBDF on COVID-19.

In this study, the mechanism behind the efficacy of HSBDF for the treatment of COVID-19 was studied based on network pharmacology and molecular docking. Databases and networks were applied to gain the herbs, components, targets, pathways, and the relationships between them. Using network pharmacology and molecular docking tools, the therapeutic mechanism of HSBDF on COVID-19 was preliminarily explained.

Natural products and TCM with high safety, high convenience, low toxicity, and broad-spectrum properties are widely used to prevent and treat infectious diseases, especially in the SARS and SARS-CoV-2 outbreaks (17,54).

Many Chinese herbal extracts have antiviral abilities, as well as anti-inflammatory and immune regulatory functions. Some of the 223 biological components in HSBDF have such effects; for example, wogonin, baicalein, and baicalin have antiviral activity (55). Genkwanin can inhibit proinflammatory transmission to prevent the cytokine storm caused by SARS-CoV-2 infection via targeting the miR-101/MKP-1/MAPK pathway (56). Albiflorin may inhibit the inflammatory response through the LOX-1/NF- κ B pathway (57). Ellagic acid has anti-inflammatory activity and inhibits HIV-1 infection *in vitro*, and has been shown to have antiviral activity against the Ebola virus, Human Papillomavirus (HPV), and human rhinoviruses (58-61). By the suppression of TLR/NF- κ B and TLR/MAPK signaling pathways, isotrifoliol can inhibit pro-inflammatory mediators *in vitro*, which might inhibit the formation of a cytokine storm caused by SARS-CoV-2 infection (62). Besides, rhein attenuates inflammation by inhibiting NF- κ B and NALP3 inflammasome, both *in vivo* and *in vitro* (63).

For 223 components of HSBDF, the compositions play a significant direct role in the antiviral capability, the regulation of metabolism and synthesis, as well as the inflammatory immune regulation.

The mechanism of action of HSBDF on COVID-19 includes seven KEGG pathway components: (I) vascular function regulation pathways; (II) pathways associated with the immune response and inflammation; (III) metabolism pathways; (IV) sex hormone pathways; (V) microbial infection pathways; (VI) nerve-related pathways; and (VII) other pathways. The specific pathways mechanisms were analyzed as follows:

The vascular function regulation pathway includes vascular smooth muscle contraction, the VEGF signaling pathway, the renin-angiotensin system, and renin secretion. Its mechanism might be based on the indirect regulation of *ACE2* both in the renin-angiotensin system and renin secretion (Figure 12), and HSBDF influences the regulation of renin and the biotransformation of angiotensin (64). The genetic deletion of *ACE2* induces vascular dysfunction in mice, suggesting a relationship between *ACE2* and vascular function, reflecting the association of COVID-19 and vascular function (65-68). Moreover, COVID-19 is closely related to cardiovascular diseases, and the regulation of cardiovascular diseases is beneficial for treatment, especially for critically ill patients (68,69). Because of the association of *ACE2* with cardiovascular diseases, there is a potential for treatment with HSBDF by targeting the vascular function regulation pathway.

For pathways associated with the immune response and inflammation, we found significant roles for arachidonic acid metabolism, platelet activation, NF-kappa B signaling, chemokine signaling, and PPAR signaling (70-75). Notably, arachidonic acid is essential for prostaglandin, and is crucial for the prevention of a cytokine storm and is also the crucial target for cytochrome P450 (73). Therefore, arachidonic acid metabolism has the pharmacological effect of preventing a cytokine storm. For platelet activation, when the virus is invading the body, inflammation occurs. The immune defense mechanism is based on recruiting patrolling monocytes, and this recruitment process is based on the *CCN1* protein generated from platelets and vascular endothelium, thus reflecting the significance of platelet activation (74). Moreover, through the NF-kappa B signaling pathway, HSBDF directly activates PKC β protein and indirectly affects a series of complex proteins, affecting survival and inflammation. Through the immune response and inflammation pathways, HSBDF plays a

pharmacological role in the treatment of COVID-19 by stopping the "cytokine storm".

According to the single-cell RNA sequencing research of patient tissues, the digestive system is a vital target organ for SARS-CoV-2 infection, and metabolism and endocrine regulation is closely related to COVID-19 (76). Therefore, bile secretion, metabolic pathways, the AMPK signaling pathway, and other metabolic pathways are related to the treatment. The metabolism pathways intervene in the endocrinology and metabolism processes to protect the body, which reduces the number of cases from mild to severe symptoms, especially for patients with diabetes, cardiovascular disease, and metabolic disease. Due to the target organ infected by the virus, impairment of the intestinal barrier function would lead to the translocation of enteric bacteria and endotoxin, thus aggravating respiratory diseases of distal organs through intestinal lymphatic circulation. HSBDF could improve a patients' loss of appetite after infection with SARS-CoV-2, avoid deterioration of the disease, and affect the synthesis of cholesterol and protein to speed up the healing process (77).

A gender difference was found for infections with SARS-CoV-2, and the incidence and mortality in men are higher than that in women. A total of 41,510 out of 77,932 patients (53.3%) were males, and the mortality rates of male and female patients were 2.8% and 1.7%, respectively (78). Research elucidated the biological factors involved apart from differences in living habits: firstly, *ACE2* is commonly expressed in prostate and testicles; secondly, *ACE2* is positively correlated with the expression of androgen receptor (AR) and *TMPRSS2*, which is the crucial molecule that promotes the development of male prostate cancer and can be positively regulated by androgens in the body; thirdly, *CCL14*, *CCL23*, *IL7*, *IL16*, and *IL1* which are pro-inflammatory cytokines and chemokines are also preferentially expressed in males, and the chemokines *CCL2*, *CCL3*, and *CCL4* that could play a protective role are higher expressed in females; besides, T cells in the lungs of men were found to be less virus-killing than those found in women (2,78). For the effect of HSBDF on COVID-19, the regulation of sex hormone pathways consists of ovarian steroidogenesis and steroid hormone biosynthesis, and the virus infection process is intervened by the regulation of steroid hormone pathways, especially the sex steroid hormones.

Critically ill patients have neurological symptoms, which are manifested as acute cerebrovascular disease, unconsciousness, and skeletal myopathy (79), and HSBDF

can be used for those patients. Moreover, through the neuropathic pathway, including neuroactive ligand-receptor interaction, serotonergic synapse, and the sphingolipid signaling pathway, HSBDF could potentially protect critically ill patients from neurological symptoms. Besides, some other pathways, such as pathogenic *Escherichia coli* infection, elaborate on the treatment mechanism of HSBDF on SARS-CoV-2. The antiviral and antimicrobial effects of HSBDF could be revealed from the view of microbial infections.

The therapeutic mechanism of HSBDF on SARS-CoV-2 can further be elucidated by molecular docking. We found that ellagic acid could dock well with RdRp and with RNA, in which the Nsp12 part has the function of RNA polymerase and plays a vital role in the replication and transcription of viral genomes (80). Ellagic acid could also dock well with Nsp14 (N7-MTase), which has a double enzyme activity, forms a complex with Nsp10 to function, and is responsible for capping mRNA (81). Isotrifoliol has a great docking ability with the N protein NCB site of Helicase (Nsp13), the ideal target for wide-spectrum antiviral drugs (82). Isotrifoliol could dock well with E protein, which forms pentapolymer and has ion channel functions, and HSBDF could cause interference with virus-related ion channels and block the toxic expression of the virus (83). Rhein could dock well with Nsp14 (ExoN), proofreading RNA replication and transcription, which reveals HSBDF might prevent the replication and transcription of RNA. Furthermore, the bioactive components of HSBDF also have an excellent binding ability with the proteins directly related to SARS-CoV-2. The discussion above explains the docking status between molecules of HSBDF and crucial proteins and indicates the therapeutic mechanism behind the effects of HSBDF for the treatment of COVID-19, which is mainly based on multiple effects of TCM.

While preliminary, this finding suggests that the pharmacological mechanism of HSBDF on COVID-19 is based on direct and indirect antiviral effects. The binding activity of active ingredients and crucial proteins related to SARS-CoV-2 plays direct antiviral effects, and the key biological pathway mechanisms of HSBDF on the body play indirect antiviral effects. This study has identified the active ingredients, such as isotrifoliol and ellagic acid, which have an excellent binding ability with crucial proteins directly related to COVID-19, including ACE2, ADAM17, and 3CLpro. Thus, it achieves a pharmacological effect through the combination of molecules and then directly

achieves the therapeutic effect. Likewise, the indirect pharmacological mechanisms of HSBDF on COVID-19 are based on biological pathways, especially the crucial biological signaling pathways, including the AMPK signaling pathway, arachidonic acid metabolism, NF-kappa B signaling pathway, serotonergic synapse, renin secretion, and the renin-angiotensin system. The indirect pharmacological effects of HSBDF on COVID-19 are based on multiple target proteins of HSBDF on the body, such as *AMPK*, *PPAR* γ , arachidonic acid metabolism-related proteins, *5-HT3*, and *ACE*, thus having mainly indirect pharmacological effects on metabolism, inflammation, and nerves. The contribution of this study has been to preliminarily confirm the potential pharmacological mechanisms of HSBDF on COVID-19, which provides a theoretical basis and a reasonable preliminary explanation for the clinical effects, including severe conversion (hospitalization) rate, the increase in total effective treatment rate, and the improvement in lung imaging signs and major related symptoms (50).

Above all, TCM plays an essential part in the antiviral process by targeting multiple components, targets, and pathways based on the network pharmacology and molecular docking, and different networks reflect the various properties of the mechanisms. Moreover, this research needs to be further improved and analyzed through experiments, and the molecular mechanism of HSBDF on COVID-19 should be deepened. Western-Blot (WB) and Polymerase Chain Reaction (PCR) experiments could be used for mechanism verification.

Conclusions

In summary, 223 components, 84 crucial target proteins, and 46 potential pathways were acquired by network pharmacology, molecular docking, and TCM theory, which preliminarily elaborated the therapeutic mechanism of HSBDF on COVID-19 at the molecular level. HSBDF may affect SARS-CoV-2 infection through multiple components, targets, and pathways.

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Footnote

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Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at <http://dx.doi.org/10.21037/apm-20-1759>). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

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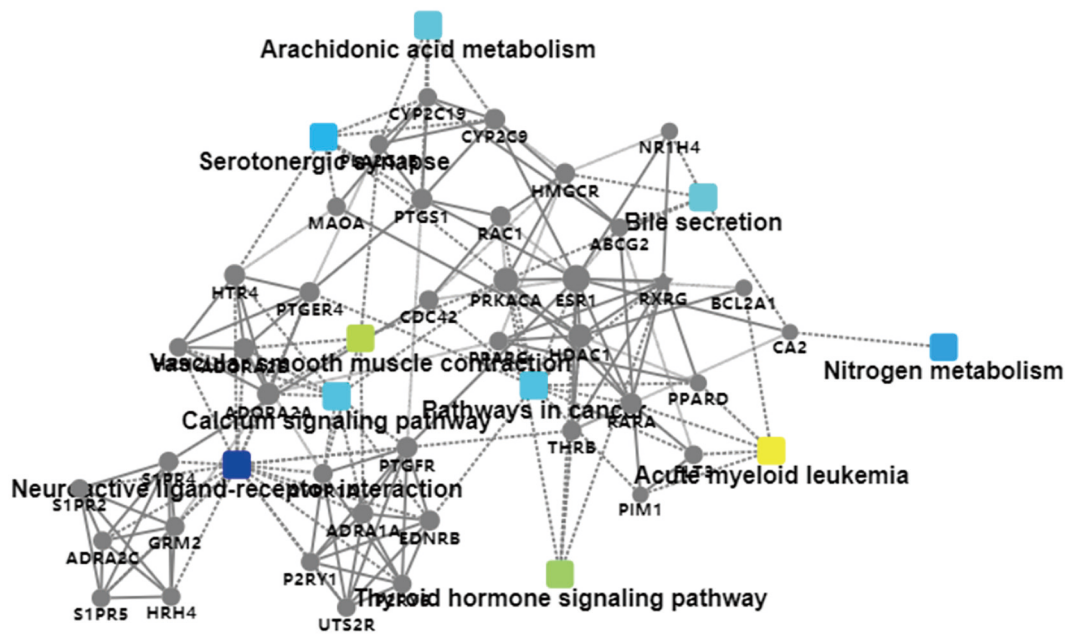


Figure S1 The essential part of the Protein-Protein Interaction (PPI) and target-pathway (T-P) network.

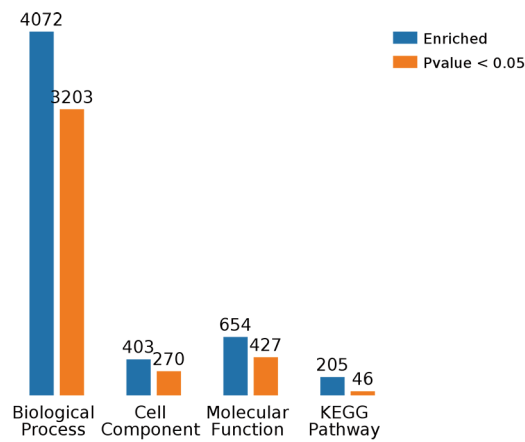


Figure S2 The summary of the Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis.

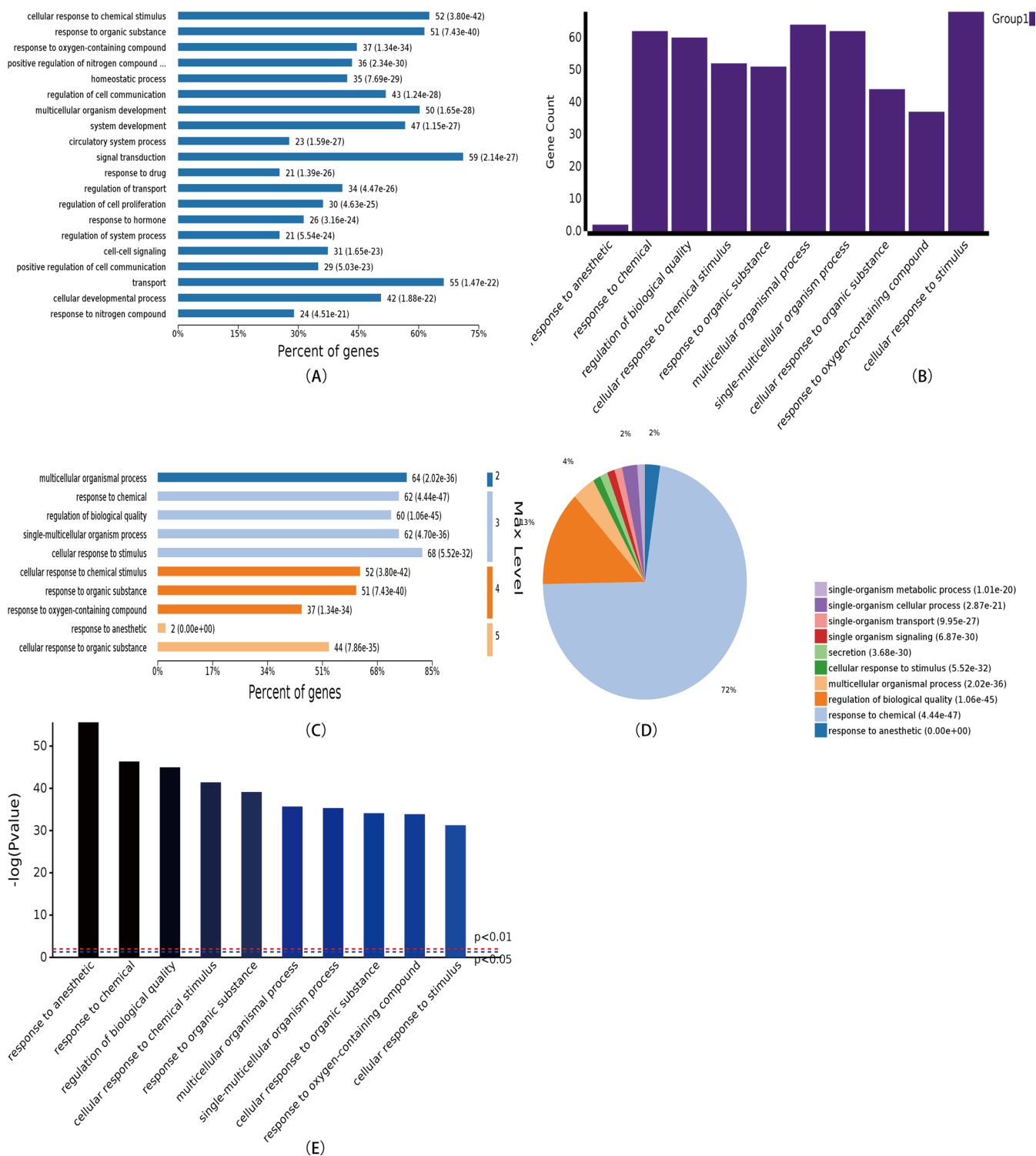


Figure S3 (A) Bar chart of the biological process (percent of genes); (B) bar chart of the biological process group (gene count); (C) bar chart of the biological process group (percent of genes); (D) pie chart of the biological process level; (E) bar chart of statistically significant biological process.

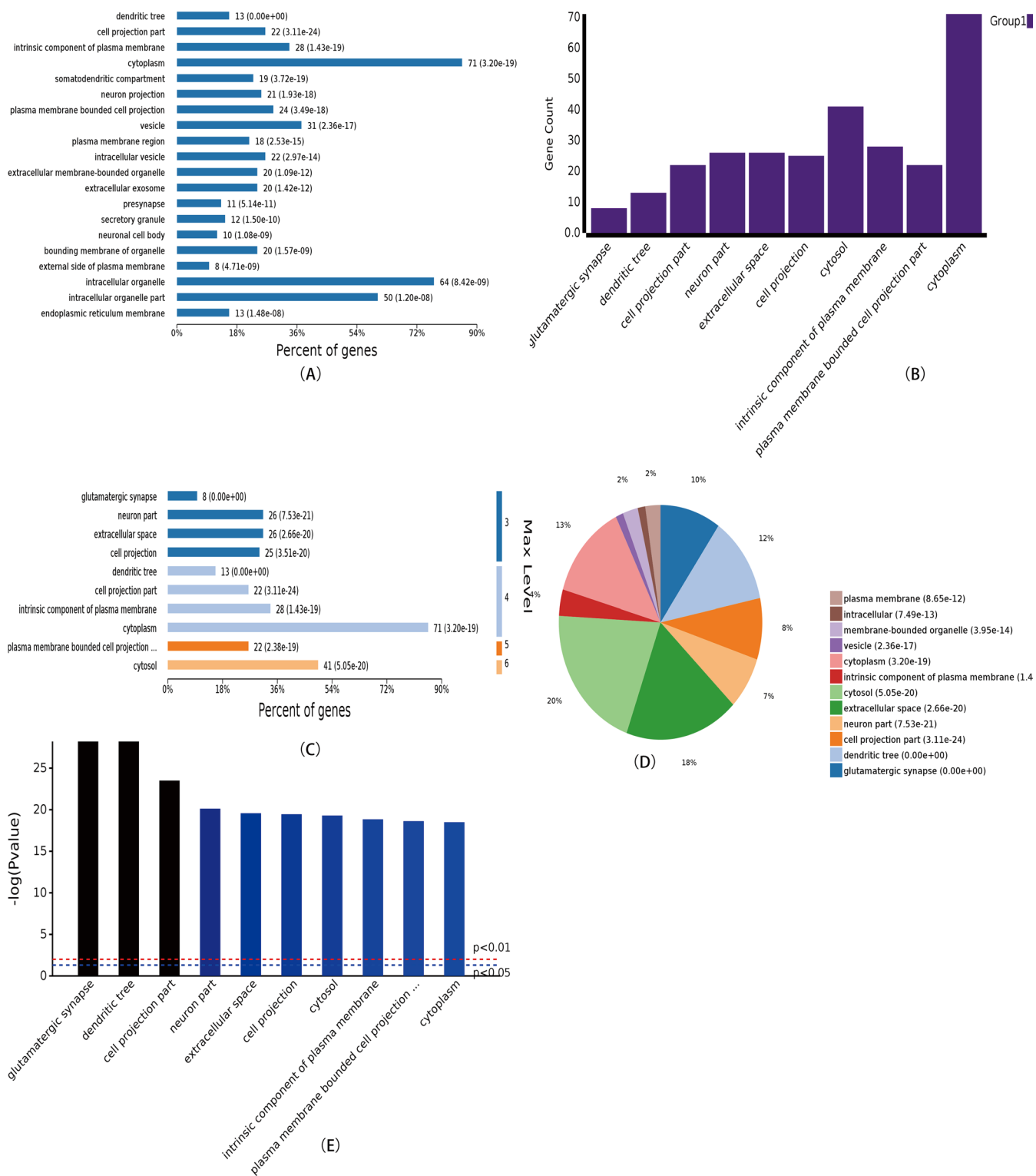


Figure S4 (A) Bar chart of the cell component (percent of genes); (B) bar chart of the cell component group (gene count); (C) bar chart of the cell component level (percent of genes); (D) pie chart of the cell component; (E) bar chart of statistically significant cell component [-log(Pvalue)].

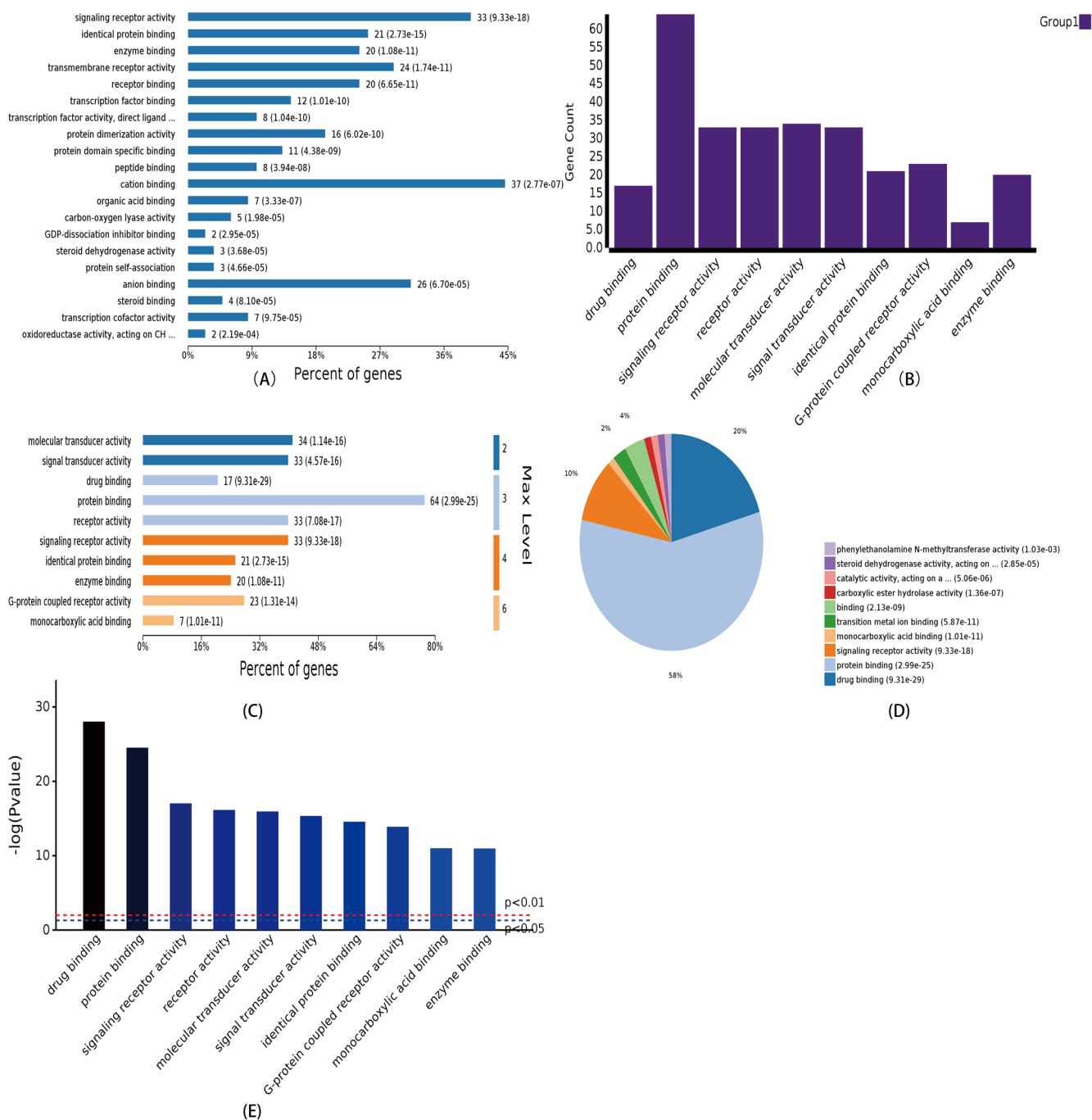


Figure S5 (A) Bar chart of the molecular function (percent of genes); (B) bar chart of the molecular function group (gene count); (C) bar chart of the molecular function level (percent of genes); (D) pie chart of the molecular function; (E) bar chart of statistically significant molecular function [-log(Pvalue)].