Exploration of potential targets and mechanisms of Naringenin in treating autism spectrum disorder via network pharmacology and molecular docking

Jialin Gai, MSca, Jinxiao Xing, MScb, Yangyang Wang, MScb, Junfang Lei, MScb, Chengdong Zhang, MScb, Jinfei Zhang, MScb, Jiqin Tang, MSa-e

Abstract
Naringenin (NR) is a kind of flavonoid which plays a great role in the treatment of autism spectrum disorder (ASD). However, the underlying mechanism of NR in treating ASD still remains unclear. This study used network pharmacology and molecular docking to examine the potential targets and pharmacological mechanism of NR on ASD. Targets related to NR were screened from Traditional Chinese Medicine System Pharmacology Database and Analysis Platform (TCMSP), Encyclopedia of Traditional Chinese Medicine Database (ETCM), Traditional Chinese Medicine Integrated Database (TCMID), PharmaMapper database, and targets related to ASD were screened from Online Mendelian Inheritance In Man (OMIM), Disgenet, GeneCards, Therapeutic Target Database (TTD), Drugbank, and ETCM. Screened of the intersected gene targets. Then, we used the protein–protein interaction (PPI) networks to construct a PPI network and used Network Analyzer plug-in to perform topological analysis to screen out the core target. We used Metascape platform to perform gene ontology (GO) functional enrichment analysis and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analysis, and used Chem draw, Pymol, AutoDock 1.5.6 software for molecular docking verification with core targets. A total of 149 targets of NR and 1594 potential targets of ASD were screened, and 43 intersected targets and 8 key targets were obtained and screened. A total of 176 GO items were obtained by GO enrichment analysis (P < .05), 153 entries on biological process (BP), 12 entries on BP and 11entries on cell composition (CC) were included. A total of 100 signaling pathways were obtained by KEGG pathway enrichment screening (P < .05). The pathways that are closely related to the pathogenesis of ASD are estrogen signaling, thyroid hormone signaling pathway, prolactin signaling pathway, and endocrine resistance pathway. Molecular docking results showed that NR had the best docking activity with the core target CASP3, and had good binding ability with AKT1, ESR1, ACTB and MAPK3. Taken together, our findings support that NR exerts therapeutic effects on ASD with multi-target, and multi-pathway characteristics, which provides a preliminary theoretical basis for clinical trials. The mechanism of anti-oxidative stress response, anti-apoptosis, regulation of cell growth and metabolism, anti-inflammatory, balance hormone levels may be important for the therapeutic effect.

Abbreviations: ASD = autism spectrum disorder, BDNF = brain-derived neurotrophic factor, CC = cellular component, ETCM = Encyclopedia Of Traditional Chinese Medicine Database, GO = gene ontology, KEGG = Kyoto Encyclopedia of Genes and Genomes, MF = molecular function, NR = Naringenin, OB = oral bioavailability, OMIM = Online Mendelian Inheritance In Man, PDB = Protein Data Bank, PPI = protein-protein interaction, ROS = reactive oxygen species, STRING = protein-protein interaction networks functional enrichment analysis, TCM = Traditional Chinese Medicine, TCMID = Traditional Chinese Medicine Integrated Database, TCMSP = Traditional Chinese Medicine System Pharmacology Database and Analysis Platform.

Keywords: autism, autism spectrum disorder, molecular docking, Naringenin, network pharmacology
1. Introduction

Autism Spectrum Disorder (ASD) is a complex chronic neuro-developmental disease that occurs in childhood and is accompanied by persistent deficits such as various social impairments, behavioral impairments, language communication impairments, and repetitive stereotyped behaviors. These include anxiety, intellectual disability, motor abnormalities, epilepsy, attention disturbance, hyperactivity, sleep disturbance, immune system deregulation, and gastrointestinal problems. ASD has become increasingly common in recent years, with a reported global incidence rate of about 1% to 2%.[1] In China, autistic children account for about 36.9% of mentally ill children, which brings a heavy economic burden to society and their families.[2] Clinical treatment with western medicine consists primarily of drug combinations and behavior modification, educational intervention, speech training, and modern rehabilitation training. The target drugs for this disease are also in the exploratory stage, requiring a prolonged disease course to achieve curative effect while growing evidence suggests that traditional NR can be beneficial.[3]

In ancient Chinese medicine books, “dementia,” “late speech,” “fetal weakness,” “coma” and “depression syndrome” were assigned to the autism category. The Traditional Chinese Medicine (TCM) pathogenesis is based on deficiency and excess and is closely related to the brain, heart, kidney, and liver, involving qi deficiency, essence deficiency, blood deficiency, fire (heat), and phlegm-dampness, which can easily lead to brain disorders and insufficiency.[4] Children with ASD often show typical heart and spleen deficiency symptoms and gastrointestinal problems such as food intolerance, food allergy, constipation or diarrhea, partial eclipse, and picky eating.[5] TCM often embodies the overall concept and dialectical thinking of TCM concepts. Therefore, this study sought to obtain the structure and target information of NR, and the molecular docking technology, hoping to provide a scientific theoretical basis for the optimization of drug combinations and behavior modification, educational intervention, speech training, and modern rehabilitation training. The target drugs for this disease are also in the exploratory stage, requiring a prolonged disease course to achieve curative effect while growing evidence suggests that traditional NR can be beneficial.[6]

NR in treating ASD therapeutic potential has been reviewed.[7] but the targets, pathway, and mechanism remain unclear. Hitherto, few specific studies have been conducted. As an emerging frontier science for TCM research, network pharmacology can integrate and systematize drugs, targets, and interactions, and repetitive behaviors.[8] Likewise, it indicates anti-hyperlipidemic, anti-depressant, anti-proliferative effects and its effect on metabolic disorders and cognitive dysfunction which provides a pharmacological and experimental foothold for the prevention and treatment of ASD.[9]

2. Materials and methods

2.1. Obtain NR structure and predict targets information

The TCMSp[10] (http://tcmspw.com/tcmsp.php) was used to obtain the structure and target information of NR, and the

2.2. Screening intersected gene targets related to NR and ASD

2.2.1. Screening of the target genes of NR. The target genes of NR were screened from ETCM[11] (http://www.tcmip.cn/ETCM/), TCMID[12] (http://www.megabionet.org/tcmid/), and PharmaMapper[13] (http://lilab.ecust.edu.cn/pharmmapper/) platform using “naringin” as the keyword, which were merged to remove duplicates, calibrated and compared through the Uniprot database[14] (http://www.uniprot.org/). After unifying the ID name, we finally obtained the corresponding compound target genetic information.

2.2.2. Screening of ASD-related genes. By using “Autism Spectrum Disorder” as the keyword, we searched 6 disease databases of OMIM (http://omim.org/), Disgenet (https://www.disgenet.org/), GeneCards (http://www.genecard.org/), TTD (http://bidd.nus.edu.sg/group/cjtd), Drugbank (http://
www.drugbank.ca), and ETCM to obtain ASD-related genes, and the target names were standardized through the Uniprot database. Then, the target names were combined and deduplicated to obtain ASD-related genes.

2.2.3. Screening of the intersected gene targets. The NR action targets and disease targets retrieved above were imported into Venny 2.1.0 (https://bioinfogp.cnb.csic.es/tools/venny) to obtain the intersected genes, and a Venn plot was created to visualize the potential targets of NR in the treatment of ASD.

2.3. Construction of protein–protein interactions (PPI) network

The construction of PPI is crucial for understanding the indication to what extent 2 proteins interact with each other. We imported the potential targets obtained above into the STRING (https://string-db.org) platform. “Organism” was set to “Homo Sapiens,” free nodes were hidden, and PPI network was constructed to derive node information to prepare for the next step, which involved generating a network graph to explore the interaction between NR-related targets and ASD targets.

2.4. Screening of core targets

In order to obtain core target genes, we used the tool Network Analyzer in Cytoscape 3.9.0 to analyze the network topology parameters. The node size and color depth reflected the degree value. And the key targets were filtered through the plug-in centiscape2.2 by calculating the mean values of betweenness, closeness, and degree. The targets above the 3 thresholds of betweenness, closeness, and degree were selected as “key targets,” which could be considered as the core target genes of NR in the treatment of ASD.

2.5. GO and KEGG pathway enrichment analysis

GO biological function enrichment analysis and KEGG metabolic pathway enrichment analysis were performed on the screened key targets through the Metascape platform. Homo Sapiens (human source) was selected for the species background of Metascape; the P-value was set to <0.01, the Min Overlap value was 3, and the Min Enrichment value was 1.5. GO annotation was conducted on 3 aspects: BP, CC, and molecular function (MF). The microscopic letters (http://www.bioinformatics.com.cn) generated the bubble plots and histogram, and the Cytoscape 3.9.0 platform was used to draw the “component-target-disease-pathway” network diagram.

2.6. Molecular docking

2.6.1. Preparation of target proteins and small molecule structures. We conducted a molecular docking analysis of the key targets screened in 1.4 with the NR component. The structure of the NR compound was obtained from the PubChem database. The structure optimization was performed by Chem3D 17.0 software; the energy of the downloaded compound was minimized and converted into mol2 format. Small molecule compounds were imported into AutoDock Tools-1.5.6 software and saved as Protein Data Bank (PDB) files after adding atomic charges, assigning atomic types, and making all flexible bonds rotatable by default.

The crystal structure of the target protein was obtained from the Uniprot protein database. After using Pymol 2.1 software to delete irrelevant small molecules in the protein molecule, the protein molecule was imported into AutoDock Tool 1.5.6 software to delete water molecules, add hydrogen atoms, set the atom type, and finally saved as PDB files.

2.6.2. Molecular docking process. The treated compound was used as a small molecule ligand, and 8 protein targets were used as receptors. The center position and length, width, and height of the Grid Box were set to 50 × 50 × 50 according to interactions between the small molecule and the target. Finally, molecular docking was performed by AutoDock Tool software to verify their binding efficiency, and a Lamarckian genetic algorithm was used for molecular docking calculation.

3. Result

3.1. Pharmacological and molecular structure information of NR

Through the TCMSP and the PubChem database, we obtained the structure of NR. The 2D and 3D structures of NR are shown in Figure 2A and B. Its molecular ID is MOL004328, molecular formula C15H12O5, molecular weight 272.27, the ratio of lipid water partition coefficient (AlogP) 2.30, OB 59.29, blood-brain barrier (BBB) −0.37, and drug-likeness (DL) 0.21, drug half-life (HL) 16.98.

3.2. Screening of the intersected gene targets related to NR and ASD

Thirty-six potential targets of NR were obtained from the TCMSP database, 29 potential targets were obtained from the Swiss Target Prediction database, and 44, 48, 65 potential targets of NR were obtained from ETCM, TCMID, and PharmaMapper, respectively. After the targets obtained in the above databases were merged and deduplicated, 149 NR-related targets were obtained.

Ninety-one, 1113, 733, and 68 ASD-related targets were obtained from the OMIM, Disgenet, GeneCard, and Drugbank databases, respectively, and combined with the 2 databases of TTD and ETCM to supplement related targets. A total of 1594 ASD disease targets were obtained after merging and deleting duplicate targets.

The screened NR and ASD-related targets were mapped, and the intersection of the 2 was obtained, yielding 43 targets, and a

![Figure 2](https://example.com/figure2.png)

**Figure 2.** (A) The 2D structure of NR, (B) the 3D structure of NR, and (C) NR-ASD Intersection Target VENN Map. Compounds = NR (blue), Disease = ASD (yellow). A total of 43 gene products were identified as common. ASD = autism spectrum disorder, NR = Narirutinin.
“drug-disease” Venn plot was generated in Figure 2C. The potential targets of NR in the treatment of ASD are shown in Table 1.

3.3. Construction of PPI network of NR against ASD

The protein interaction PPI information obtained from the STRING database was imported into Cytoscape 3.9.0 software as a “TSV” file to generate a PPI network map of potential targets involving 40 nodes and 234 edges (Fig. 3). The node size and color depth in the figure represent the node degree value. A larger node and darker color corresponded to a larger node degree value. The thicker and darker the edge, the higher the interconnectivity and the greater its role in the network. The results showed that the average node degree of the network was 11.7, and the clustering coefficient was 0.593.

3.4. The screening results of core targets network

In Cytoscape3.9.0 software, the plug-in centiscape2.2 was used to calculate betweenness, closeness, and degree values. The betweenness, closeness, and degree values were 0.014236605, 33.5, and 11.7. Targets with values above the 3 thresholds were set as key targets (Fig. 4). The details are shown in Table 2. Targets that met the above requirements were AKT1, ACTB, ESR1, CASP3, MAPK3, SRC, CAT, and MAPK1, suggesting they are closely related to other target proteins. Based on this, we hypothesized that they might be involved in the pharmacological effects of NR in the treatment of ASD.

3.5. KEGG pathway analysis and GO enrichment analysis

The screened core targets were imported into the Metascape database for GO gene function annotation and KEGG pathway enrichment analysis. The top 10 most significant GO terms according to the count are shown in Figure 5A–C, and display the bubble chart according to the P value, as shown in Figure 5D. During GO annotation, a total of 176 GO terms were obtained for BP (n = 153), CC (n = 11), and MF (n = 12). Significantly enriched biological processes consisted of response to reactive oxygen species (ROS), chemical stress, oxidative stress and growth factors, and organic cyclic compounds. Significantly enriched CC GO terms consisted of focal adhesion, cell-substrate junction, membrane raft, membrane microdomain, caveola, plasma membrane raft, etc. Finally, the significantly enriched MF GO terms included phosphoprotein binding, protein serine/threonine/tyrosine kinase activity, RNA polymerase II-specific DNA-binding transcription factor binding, protein kinase binding, kinase activity, etc.

The KEGG pathway enrichment analysis revealed 100 enriched metabolic pathways (P < 0.01), the top 20 representative pathways were selected for bubble diagram according to the P-value, as shown in Figure 6A, and the pathways were classified according to KEGG pathway classification, as shown in Figure 6B, and the specific information is provided in Table 3. KEGG network includes the first 20 pathways, of which MAPK1, MAPK3, AKT1, SRC, ACTB and other genes appear the most. It involves prolactin signaling pathway, estrogen signaling, proteoglycans in cancer, thyroid hormone signaling pathway, endocrine resistance, apoptosis, platelet activation, VEGF signaling pathway, Yersinia infection, etc.

3.6. “NR-Target-ASD-pathway” PPI network diagram

The “NR-Target-ASD-pathway” PPI network diagram is shown in Figure 7. The purple diamond node on the left represents the NR compound, the yellow circle icon in the middle represents the key target, the green icon in the middle represents the representative top 20 signaling pathways where the target is located, and the blue rectangle icon on the right represents autism. After further topological analysis by Network Analyzer, we found that NR may play a co-regulatory role on ASD mediated by AKT1, MAPK1, MAPK3 and involve the proteoglycan pathway in cancer, thyroid hormone signaling pathway, ROS pathway, and other pathways.

3.7. The results of molecular docking

To further explore the interaction between NR and key targets, the NR compound and key targets AKT1, ACTB, CAT, ESR1, MAPK1, MAPK3, CASP3, and SRC were molecularly docked. The results of the final molecular docking study are shown in Table 4. It is well-established that binding energy can predict the

### Table 1

| Number | Gene      | Number | Gene |
|--------|-----------|--------|------|
| 1      | ESR1      | 23     | SHBG |
| 2      | PTGS2     | 24     | NLRP3|
| 3      | Pik3CG    | 25     | NF2L2|
| 4      | AKT1      | 26     | MTOR |
| 5      | BCL2      | 27     | MABO |
| 6      | MAPK3     | 28     | ABG2 |
| 7      | MAPK1     | 29     | ADORA2A |
| 8      | CASP3     | 30     | AR   |
| 9      | SOD1      | 31     | TIR  |
| 10     | CAT       | 32     | CTSB |
| 11     | PPARG     | 33     | POE4D|
| 12     | APOB      | 34     | F2   |
| 13     | CYP19A1   | 35     | PPARD|
| 14     | GSTP1     | 36     | FGR1 |
| 15     | UGT1A1    | 37     | PAH  |
| 16     | ADIPQ     | 38     | SRC  |
| 17     | ABAT      | 39     | HK1  |
| 18     | SODAT1    | 40     | POE4B|
| 19     | ACTB      | 41     | AMY1A|
| 20     | CYP1B1    | 42     | AMY1B|
| 21     | ESR2      | 43     | AMY1C|
| 22     | NR112     | 44     |      |

NR = Naringenin.
binding activity between 2 parties. Indeed, the lower the binding energy, the tighter the compound binds to the target protein, and the more stable their conformation is. Binding energy $< -5 \text{ kcal/mol}$ is associated with good binding ability, and binding energy $< -7 \text{ kcal/mol}$ suggests strong activity.\cite{33} As seen in Table 4, the binding energies of all targets and compounds were $< -5 \text{ kcal/mol}$, indicating that NR has a good binding effect with these target proteins and has a high degree of matching. The binding energies of the 5 target proteins CASP3, AKT1, ESR1, ACTB, and MAPK3 were all $< -7 \text{ kcal/mol}$ and exhibited the strongest binding ability to the compound, suggesting that NR plays an essential role in ASD through these targets.

The complex formed by the compound and the protein after docking was visualized with Pymol2.1 software, and the binding mode of the compound and the protein was obtained. According to the binding mode, we can clearly see the amino acid residues that the compound binds to the protein pocket. Molecular docking details of NR with each protein are shown in Figures 8–12. Then we selected the top 3 target proteins with the lowest binding energies for analysis.

We found that NR has the best match with the active site of CASP3 protein; its phenolic hydroxyl group could form multiple hydrogen bonds with SER-120, GLN-161, ARG-64, yielding a strong effect on protein stability. In addition, the benzene ring of this compound can form strong π-π conjugated interactions with the benzene ring of TYR-204, playing an essential role in the stabilization of small molecules. The amino acid residues that interact with the active site of NR and AKT1 protein are THR-291, MET-227, GLU-228, TYR-229, ALA-230, ALA-177, LEU-156, VAL-164, PHE-438, and PHE-44. NR has strong hydrophobicity and can form strong hydrophobic interactions with amino acids in the active site (MET-227, TYR-229, ALA-230, ALA-177, LEU-156, VAL-164, PHE-438, and PHE-442), playing an essential role in stabilizing small molecules in the protein cavity. This compound can also form hydrogen bonds with GLU-228 and ALA-230 amino acids, which contribute to the small molecules in the anchor protein cavity. Moreover, we found that NR and the ESR1 protein site can form hydrogen bonds, hydrophobic interactions, van der Waals forces, and other interactions; for example, NR can interact with LEU-387, PHE-404, LEU-391, PHE-404, LEU-353, MET-343, LEU-346, and other amino acids to form strong hydrophobic interactions, and can form strong hydrogen bond interactions with LEU-387, ARG-394, GLU-353.

In addition, NR can interact with several other targets in various ways, effectively promoting the formation of stable complexes between proteins and small molecules and having strong associations with the targets.

4. Discussion

In recent years, there has been a burgeoning interest in exploring the underlying molecular and cellular mechanisms of ASD. Ample evidence suggests that its pathogenesis includes genetic, environmental, and immune factors. High genetic variability involving disruption of messenger ribonucleic acid (mRNA) transcription or protein function, deletion of gene exons, abnormal gene methylation, single nucleotide polymorphism, and copy number variation represents a significant risk factor for
Other risk factors include increasing parental age, influenza vaccine, drug infection or exposure to valproic acid and folic acid during pregnancy, maternal obesity, maternal smoking, preterm birth, low birth weight, and altered gut microbiota. The current treatment approach for ASD involves modulating neurotrophic factor levels, regulating gene expression, restoring cell synapse levels, restoring protein homeostasis, and stimulating oxidative stress responses.

NR, chemically known as 5,7-dihydroxy-2-(4-hydroxyphenyl)chroman-4-one, is a disaccharide derivative at position 7 via a glycosidic linkage. It is a natural flavonoid glycoside with various pharmacological activities including high antioxidant, anti-inflammatory, anti-apoptotic and neuroprotective properties that can be found in a diversity of citrus fruits, vegetables, nuts and Chinese medicine.

NR as a flavanone flavonoid that can have a protective effect on the gastrointestinal tract and effectively improve neurological diseases.

NR has oral bioavailability (OB) as an important index to evaluate the effectiveness of drugs entering human circulation and the OB value of NR is 0.01620032.

| Number | Core target | Betweenness unDir | Closeness unDir | Degree unDir |
|--------|-------------|-------------------|-----------------|--------------|
| 1      | CAT         | 98.60251676       | 0.016666667     | 30           |
| 2      | MAPK3       | 62.83480866       | 0.017241379     | 27           |
| 3      | SRC         | 59.50302501       | 0.016949153     | 23           |
| 4      | CASP3       | 76.96309488       | 0.01754386      | 23           |
| 5      | ESR1        | 64.03900537       | 0.017241379     | 22           |
| 6      | ACTB        | 141.6436652       | 0.019230769     | 20           |
| 7      | AKT1        | 268.8019609       | 0.020833333     | 19           |
| 8      | MAPK1       | 38.27931872       | 0.018129032     | 17           |

Figure 5. (A) GO function enrichment analysis bubble diagram. The circle size represents the number of enriched genes; color represents the significance of enrichment. BP = biological process, (B) MF = molecular function, (C) CC = cell component, (D) Histogram of GO enrichment analysis of NR in the treatment of ASD. ASD = autism spectrum disorder, GO = gene ontology.

Table 2

Core target information.
Figure 6. (A) Bubble plot of KEGG pathway enrichment analysis of NR – ASD, (B) KEGG pathway classification plot of NR-ASD. The 4 modules from top to bottom are cellular process, environmental information processing, Human Diseases, and Organismal Systems. ASD = autism spectrum disorder, KEGG = Kyoto Encyclopedia of Genes and Genomes, NR = Naringenin.

Table 3

| Pathway                                | Enrichment | P-value   | Count | Gene                      |
|----------------------------------------|------------|-----------|-------|---------------------------|
| Proteoglycans in cancer                | 128.943092 | 4.75E–15  | 7     | ACTB|AKT1|CASP3|ESR1|MAPK1|MAPK3|SRC |
| Thyroid hormone signaling pathway      | 187.2582645| 1.01E–13  | 6     | ACTB|AKT1|CASP3|ESR1|MAPK1|MAPK3|SRC |
| Prolactin signaling pathway            | 269.741074 | 3.22E–12  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Endocrine resistance                   | 192.6721939| 1.80E–11  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Platelet activation                    | 152.2731855| 5.95E–11  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Apoptosis                              | 138.8373162| 9.51E–11  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Yersinia infection                     | 137.8239051| 9.87E–11  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Estrogen signaling pathway             | 136.8251812| 1.02E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Hepatitis B                            | 116.554784 | 2.30E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Influenza A                            | 110.4203216| 3.03E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Tuberculosis                           | 104.8993056| 3.92E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Neutrophil extracellular trap formation| 99.37828947| 5.15E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Kaposi sarcoma-associated herpesvirus infection| 97.3292528| 5.72E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Pathogenic Escherichia coli infection  | 95.84708122| 6.18E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Focal adhesion                         | 93.93967662| 6.83E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Rapt signaling pathway                 | 89.91369048| 8.52E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Chemical carcinogenesis - receptor activation| 89.06544811| 8.93E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| VEGF signaling pathway                 | 256.0254237| 9.13E–10  | 4     | AKT1|ESR1|MAPK1|MAPK3|SRC |
| Lipid and atherosclerosis              | 87.82267442| 9.59E–10  | 5     | AKT1|ESR1|MAPK1|MAPK3|SRC |

KEGG = Kyoto Encyclopedia of Genes and Genomes, NR = Naringenin.

Figure 7. Target-pathway PPI network of NR in the treatment of ASD. ASD = autism spectrum disorder, NR = Naringenin, PPI = protein-protein interaction.
>30 is often used as a standard for compound screening. Our study found that the OB value of NR was 59.29, indicating that NR may be easily absorbed into the blood after oral administration, and it thus can conveniently exert biological activity. Some studies showed NR could improve spatial learning and memory in aging mice by inhibiting Aβ levels and neuroinflammation, phosphorylation, and oxidative stress. It can also downregulate anti-apoptotic genes to improve cognitive dysfunction, dyskinesia, depression, and anxiety-like behavior in hypoxic mice.

In our study, a total of 43 common targets of NR in the treatment of ASD were obtained through TCMSP, OMIM, and other related databases, and 8 key targets were obtained after screening. Topological analysis and molecular docking results showed that CASP3 had the closest binding ability to NR, and other targets such as AKT1, ESR1, ACTB, MAPK3, MAPK1, SRC, CAT, suggest that their binding ability was also good. CASP3 (Caspase 3) is a group of proteases that play a key executive role during apoptosis, capable of eliminating redundant and nonfunctional synapses and removing extra cells during early childhood development, and play an important role in embryonic neuronal development. Experiments have shown that Casp3-deficient mice exhibit abnormal motor behaviors such as impaired social interaction, limited interest, and repetitive stereotypes associated with endoplasmic reticulum stress. Ample evidence substantiates that children with ASD have significantly higher levels of CASP3, which can lead to apoptosis in autistic brain tissue. NR can inhibit CASP3 elevations and alleviate endoplasmic reticulum stress and oxidative stress-induced apoptosis in ASD. Moreover, AKT1 (AKT serine/threonine kinase 1) reportedly regulates cell proliferation, metastasis, and apoptosis. Abnormalities in the PI3K/AKT/mTOR pathway have been associated with abnormal synaptic protein synthesis and the development of ASD. Current evidence suggests that NR can control cell apoptosis and maintain cell activity by regulating AKT1 target proteins. At the same time, NR can significantly inhibit AKT1 phosphorylation, promote gene transcription and protein synthesis, and activate Akt-mTOR signaling to inhibit apoptosis in autism and alleviate abnormal social behavior in mice.

ESR1, a transcription factor of the nuclear receptor family, can mediate estrogen ligands, promote growth and cell survival, and is associated with social behavior and emotional regulation in children with ASD. Interestingly, women with abnormal

| Compounds | Structure | Target ID | PDB ID | Docking Score/kcal/mol | Combination Type |
|-----------|-----------|-----------|--------|------------------------|------------------|
| NR        | C₄₃H₆₂O₅ | CASP3     | 1NMS   | –7.59                  | Hydrogen bonds,  |
|           |           |           |        |                        | Hydrophobic      |
|           |           |           |        |                        | π-stacking       |
| AKT1      | 4GV1      |           |        | –7.54                  |                  |
| ESR1      | 3ERT      |           |        | –7.48                  |                  |
| ACTB      | 6NBW      |           |        | –7.18                  |                  |
| MAPK3     | 5W62      |           |        | –7.16                  |                  |
| MAPK1     | 6SL6      |           |        | –6.87                  |                  |
| SRC       | 6ATE      |           |        | –6.31                  |                  |
| CAT       | 1DGF      |           |        | –5.63                  |                  |

NR = Naringenin, PDB = Protein Data Bank.
antenatal testosterone and sex steroid precursor exposure have higher rates of autistic traits. Abnormal chemical stimuli into cells send signals at the membrane and nuclear levels through the Estrogen receptor, stimulate kinases and phosphatases in the body, change the phosphorylation state of key protein kinases, and reduce estrogen biosynthesis.\textsuperscript{[51,52]} Experiments have demonstrated that the ER\(\alpha/\beta\)-mediated estrogen system of NR regulates oxytocin secretion in rodent hypothalamic neurons during rodent central nervous system development and can effectively activate ER-\(\alpha\) inhibitory activity, reduce cellular stress through the inhibition of ESR1-mRNA expression or the downregulation of estrogen protein involving the corresponding signaling pathway.\textsuperscript{[53]} Moreover, estrogen-induced activation of the estrogen results in a reduction in the levels of nuclear
DNA-binding activity of NF-kB, which in turn regulates the expression of inflammatory genes. Therefore, anti-inflammatory action as a potential mechanism in mediating the neuroprotective effects of estrogen.

GO and KEGG enrichment analysis enabled further analysis of their gene functions. The results showed that the therapeutic effect of NR in the treatment of ASD involves various pathways such as estrogen signaling, thyroid hormone signaling pathway, prolactin signaling pathway, endocrine resistance pathway, platelet activation pathway, and apoptosis pathway while involving different biological processes such as ROS metabolism, chemical stress, oxidative stress, cell growth, and apoptosis.

ROS are widely acknowledged as a major cause of cellular damage, premature cellular aging, and neurological diseases. ROS have been associated with increased activity of mitogen-activated protein kinases and play essential roles in cell growth,
differentiation, development, cell cycle, survival, and cell death by activating the redox reaction of tyrosine kinases while affecting the activity of critical growth and metabolism-related transcription factors, and is sensitive to redox changes.[15,66] Environmental and genetic risk factors may exacerbate ASD patients’ vulnerability to oxidative stress, impair antioxidant defense mechanisms, and lead to cell membrane damage, altered membrane fluidity, and permeability. For example, when a child is exposed to toxic air or environmental pollution in the mother’s body, an excess of free radicals is produced in the child’s body, leading to increased inflammation, toxicity, and oxidative stress. On the other hand, patients with autism exhibit blood-brain barrier leakage and are relatively sensitive to oxidative damage, leading to the manifestations of neuropsychiatric disorders, such as depression, cognitive dysfunction, psychosis, anxiety, etc. [58] Recent studies found that NR is a flavonoid that has an inhibitory effect on signal transduction enzymes and controls cell growth by regulating proteins such as protein tyrosine and protein kinases.[14] Moreover, NR possesses antioxidant properties, which can regulate the levels of ROS, inhibit oxidative stress and neuroinflammation, thereby improving cognition levels.

Interestingly, thyroid hormone signal transduction pathways have been shown to play important roles in metabolic processes such as central nervous system development and anti-apoptosis during early development.[49] NR can reduce hormone levels through the thyroid hormone signaling pathway to restore normal cell growth and metabolism in children with ASD.[60] In this regard, mothers who drink heavily or take antipsychotics for a long time during pregnancy can cause hormone imbalance, reduce oxytocin secretion, and have a higher chance of developing ASD in the baby. Overwhelming evidence substantiates that plasma oxytocin levels in autistic patients are lower than usual, and low levels of oxytocin can affect children’s social recognition behavior.[81–83] It has also been reported that children with ASD have hypoplasia of the nervous system, accompanied by inflammation, oxidative stress and mitochondrial dysfunction, and defective platelet aggregation in endothelial cells.[64] Current evidence suggests that platelets release brain-derived neurotrophic factor (BDNF) upon activation. BDNF is a growth factor and a member of the neurotrophic factor family.[65] After platelets activate BDNF, angiogenesis-related and inflammatory cytokines are released, while NR can regulate platelet activation, aggregation, secretion, and promote the expression of BDNF, angiogenesis-related and inflammatory, balance hormone levels. At the same time, the molecular docking also corresponds to the predicted results, showing good target binding ability, which verifies the feasibility of the predicted components, core targets and their pathways in this study.

This study provides a potential biological basis for the further study of NR in the treatment of ASD, but still has some limitations. We used only modern bioinformatics methods to explore the effects of NR in the treatment of ASD by using network pharmacology and molecular docking. Firstly, data from online databases are based on reviewed and predicted data, and unconfirmed and unrecorded data were not included in our study. Secondly, although CASP3, AKT1, ESR1, ACTB, MAPK3, were identified as core targets, pharmacodynamic and molecular biology experiments need to be considered to further investigate our results. We believe that this topic has great research potential and application value.

5. Conclusion

In summary, the present study found a series of novel targets and pathways for NR treating ASD and the mechanism of action on the human body suggests the involvement of multiple pathways and multiple targets. It was preliminarily predicted that NR could regulate the targets of CASP3, AKT1, ESR1, ACTB, MAPK3 and may modulate various pathways like estrogen signaling, thyroid hormone signaling pathway, prolactin signaling pathway, proteoglycan in cancer pathway, endocrine resistance pathway. Its mechanism of action are involving anti-oxidative stress response, anti-apoptosis, regulation of cell growth and metabolism, anti-inflammatory, balance hormone levels. At the same time, the molecular docking also corresponds to the predicted results, showing good target binding ability, which verifies the feasibility of the predicted components, core targets and their pathways in this study.

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Author contributions

Conceptualization: Jialin Gai.
Data curation: Jinxiao Xing.
Formal analysis: Jialin Gai, Yangyang Wang.
Funding acquisition: Jiqing Tang.
Methodology: Jialin Gai, Junfang Lei, Chengdong Zhang.
Software: Jialin Gai, Junfang Lei.
Supervision: Jiqin Tang, Jinfeng Zhang.
Writing—original draft: Jialin Gai.
Writing—review & editing: Jialin Gai, Jinxiao Xing, Yangyang Wang, Junfang Lei, Chengdong Zhang, Jinfeng Zhang, Jiqin Tang.

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