Candidate Chinese Herbal Medicine Alleviates Methamphetamine Addiction via Regulating Dopaminergic and Serotonergic Pathways

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Methamphetamine (METH) addiction and its induced mental disorders have become a severe worldwide problem. A candidate Chinese herbal medicine (CCHM) in our lab had therapeutic effects on METH-induced locomotor sensitization, however, its chemical and pharmacological profiles remain to be elucidated. The current study aimed to investigate the effect of CCHM on conditioned place preference (CPP) induced by METH and screen the main active ingredients and key targets by using network pharmacology and molecular docking methods. Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment, Gene ontology (GO) analysis and protein-protein interaction (PPI) network were performed to discover the potential mechanisms. Results showed that CCHM could significantly inhibit METH-induced CPP behaviors in mice. A total of 123 components and 43 targets were screened. According to the network pharmacology analysis, ten hub targets including D(2) dopamine receptor (DRD2) and 5-hydroxytryptamine receptor 3A (HTR3A) were screened. GO analysis and KEGG enrichment indicated that mechanisms of CCHM treatment of METH addiction were related to multiple pathways such as dopaminergic synapse and serotoninergic synapse. Western blot results showed that the protein expressions of DRD2 in nucleus accumbens and prefrontal cortex were significantly decreased in METH group, while the protein expressions of HTR3A were significantly increased. These changes caused by METH could be prevented by CCHM pretreatment. The results of molecular docking displayed that the five active ingredients such as (S)-Scoulerine, Hyndarin, and Beta-Sitosterol had good affinities with DRD2 and HTR3A. In conclusion, this study constructed the CCHM’s pharmacologic network for treating METH addiction based on the method of network analysis and experimental verification, and analyzed its major active ingredients and potential targets, indicating a new direction for further revealing its mechanisms of effect on METH addiction.

Keywords: Chinese herbal medicine, methamphetamine, addiction, network pharmacology, dopamine, serotonin
**INTRODUCTION**

Drug addiction is a severe worldwide social problem, which has attracted great attention. Compared with drugs extracted from plants such as heroin and cocaine, the proportion of people who abuse amphetamine-type synthetic drugs is increasing year by year. According to reports released by United Nations Office on Drugs and Crime and the Ministry of Public Security of China, methamphetamine (METH) has become the most commonly abused amphetamine-type stimulant in Southeast Asia and North America. METH can easily enter the central nervous system through the blood-brain barrier, so its effect on brain is more evident than its peripheral effect (Paulus and Stewart, 2020). Studies have shown that METH abuse could seriously damage the structure and function of neurons and gliocyte in the prefrontal cortex, nucleus accumbens, hippocampus, and other brain regions of addicts, and cause a variety of central nervous system diseases, such as cognitive impairment, depression, and mental disorders (Zhang et al., 2018; Nie et al., 2020). The brain structure and function of most addicts cannot recover even after a prolonged period of abstinence, and the persistent mental disorders and negative emotions could also trigger the continuous drug-seeking behaviors, which led to relapse (Glasner-Edwards et al., 2009). METH addiction not only seriously affected the physical and mental health of abusers, but also led to crime, which has become a social problem endangering
public security. Although antipsychotic drugs have been used to control the psychiatric symptoms caused by addiction, there is still a lack of drugs with definite efficacy on METH addiction in clinic. Therefore, the research and development of drugs for the prevention and treatment of METH addiction with better efficacy and fewer side effects has become the focus in drug abuse field.

The mechanism of METH addiction remains unclear, and multiple brain regions such as the prefrontal cortex and nucleus accumbens are involved, therefore, traditional Chinese medicine has become the focus of development of anti-addiction drugs due to its multi-system and multi-target mechanism (Chen et al., 2021). Our previous work have shown that a candidate Chinese herbal medicine (CCHM), which was composed of 8 traditional Chinese medicines including Corydalis yanhusuo and Codonopsis, could significantly improve the learning and memory impairment induced by METH, and inhibit the acquisition and expression of METH-induced locomotor sensitization (Can et al., 2019; Jiaming et al., 2019). The results of acute and long-term toxicity tests showed that CCHM had no detectable toxic effects (Can, 2019). However, as a Chinese herbal compound, the composition of CCHM is complex. A large number of experiments are still needed to analyze its main active ingredients and possible mechanisms. Based on the theory of systems biology, network pharmacology has the characteristics of systematic and holistic, and it is similar to the mechanism of traditional Chinese medicine in treating diseases. Network pharmacology could reveal the complex biological network relationships between drugs, targets, and diseases, analyze and predict the pharmacological mechanism of candidate drugs (Guo et al., 2019). Therefore, this study aimed to analyze the main active ingredients and critical targets of CCHM in the treatment of METH addiction based on the network pharmacology and molecular docking, and to provide the evidence for the clinical application of CCHM.

**MATERIALS AND METHODS**

**Animals**

Male C57BL/6 mice (2 months old, weighing 18–22 g) were purchased from Beijing Charles River Laboratories (Beijing, China). Mice were housed in a temperature-controlled room with a humidity of 45–55% and subjected to a 12-h cycle of alternating light and dark. All mice had free access to food and water, and were pre-adapted to the environment for at least 3 days before the experiment. All experiment procedures were approved by the Ethics Committee of Jianghan University. As shown in Figure 1A, the mice were randomly divided into four groups: the control group, the METH group, CCHM low group, and CCHM high group. METH was offered by Hubei Public Security Bureau and was dissolved in saline, and CCHM was decocted and condensed into extract before use. Mice were intraperitoneally injected with METH (2 mg/kg) or saline following 1 h intragastric administration of CCHM (14.12, 56.48 g/kg) or saline. The dosage of CCHM was selected based on our published articles (Can et al., 2019).

**The Conditioned Place Preference Apparatus and Procedure**

The conditioned place preference (CPP) facility was brought from Shanghai Xinruan Information Technology Co., Ltd. (Shanghai, China) and comprised three compartments. One chamber had a grid floor and black walls (L × W × H: 35 cm × 35 cm × 35 cm). Another same-sized chamber had a mesh floor and white walls. The middle chamber had a flat, gray floor and gray walls (L × W × H: 15 cm × 35 cm × 35 cm) and is connected to the other two compartments by removable doors. The white and black chamber had a ceiling light and camera, respectively. The general activity in the facility was monitored by video, and the time spent in each chamber was recorded using XR-XT401 software.

The CPP procedure used in this experiment was performed as previously described (Yang G. et al., 2020). The pretreatment period lasted from day 1 to day 3. Mice were placed in the middle chamber and then moved freely between the black and white chambers for 15 min per day. The time spent in each chamber on the third day was used as pre-processing data. The most visited chamber (black chamber) was designated as the preferred compartment and the other compartment (white chamber) was defined as drug compartment. Each animal received individual CPP training. The CPP training period lasted for 8 days (day 4–day 11) and included four METH sessions and four saline sessions. The saline- and drug-paired compartments were separated by closed doors. On day 4, 6, 8, and 10, mice in METH or CCHM group were pretreated with saline or CCHM for 1 h and then confined to a drug-paired chamber (white chamber) for 1 h after intraperitoneally injected with 2 mg/kg METH. On days 5, 7, 9, and 11, the mice in METH or CCHM group were given saline or CCHM respectively and confined to the saline paired chamber (black chamber) for 1 h after saline injection. After each session, the floor and walls of each chamber were wiped with 75% ethanol. On day 12, the post-adaptation test was conducted for 15 min, and each mouse was placed in the middle chamber and then moved freely between the black and white chambers. Preference (CPP score) was calculated by subtracting the time spent in the white chamber on day 3 from the time spent in the white chamber on day 12 (Yang C. et al., 2020). CPP score reflects the difference in the time spent in the post-conditioning and pre-conditioning phases of METH.

**Identification of Candidate Components in Candidate Chinese Herbal Medicine**

Candidate Chinese herbal medicine was composed of eight Chinese medicinal herbs, including C. yanhusuo, Poria cocos, Codonopsis pilosula, Licorice, Atractylodes macrocephala Koidiz, Astragalus membranaceus, Angelica sinensis, and American ginseng. The components of CCHM were retrieved from Traditional Chinese medicine system pharmacology (TCMSP) database. Oral Chinese medicine must overcome the obstacles of absorption, distribution, metabolism, and excretion (ADME) process in order to be effective, so oral bioavailability (OB) is one...
Ru et al. Herbal Medicine Alleviates Methamphetamine Addiction

FIGURE 1 | Effect of candidate Chinese herbal medicine (CCHM) on Methamphetamine (METH)-induced conditioned place preference (CPP) in mice. (A) The experiment procedure of CPP test. Mice received METH (2 mg/kg, i.p) or saline (10 ml/kg, i.p) injections following 1 h pretreatments of CCHM (14.12, 56.48 g/kg, i.g) or saline (10 ml/kg, i.g) treatments. (B) Time spent in the white chamber was collected before and after METH abuse. (C) CPP score was calculated by subtracting the time spent in the drug-paired compartment in the pre-conditioning test from that after METH exposure. Data were presented as mean ± SE, **p < 0.01, compared with control group; #p < 0.05, compared with METH group, n = 10.

of the most important pharmacokinetic parameters in ADME. High OB is usually an important index for determining drug similarity (DL) of active substances (Guo et al., 2019). Substances with OB no less than 30% are considered to have a higher OB. DL index, as a qualitative concept used to estimate the medicinal properties of molecules in drug design, can be used for rapid screening of active substances (Tao et al., 2013; He et al., 2019). In the Drugbank database, the average DL index is 0.18. Substances with a DL index no less than 0.18 have higher medicinal properties. Therefore, compounds with OB ≥ 30% and DL index ≥ 0.18 and blood brain barrier (BBB) ≥ −0.3 in CCHM were selected as active substances in this study. The possible targets of the active substances of CCHM were searched from TCMSP database. The Uniprot database3 and the bioDBnet website4 were used to verify the gene symbols corresponding to the possible targets.

Screening of Possible Targets of Methamphetamine Addiction

Potential targets involved in METH addiction were collected from OMIM,5 Drugbank,6 and Genecards7 databases. Potential targets from different databases were combined and the duplicates has been deleted.

3http://www.Unitprot.org/
4https://biodbnet-abcc.ncifcrf.gov/
5https://www.omim.org/
6https://www.drugbank.ca/
7https://www.genecards.org/
Network Analysis of Active Compounds and Potential Targets

The potential targets of active compounds in CCHM were mapped with METH addiction-related targets on the Bioinformatics and Evolutionary Genomics website. Cytoscape 3.8.0. software was used to construct the “herb-compound-target” network (Han et al., 2020). In these graphical networks, the compounds and proteins were expressed as nodes, whereas the compound-target interactions were expressed as edges.

Network Analysis of Protein-Protein Interaction

Potential targets of CCHM on METH addiction were analyzed by online STRING 11.0 to construct protein-protein interaction (PPI) network. The network was visualized with Cytoscape (3.8.0) and CytoHubba, a plug-in in Cytoscape, to filter the modules from the PPI network and to obtain the most important hub genes based on the degree score.

Gene Ontology and Kyoto Encyclopedia of Genes and Genomes Enrichment of Hub Genes

To further characterize the molecular mechanism of CCHM on METH addiction, the DAVID website was used to conduct the GO analysis and KEGG enrichment of hub targets. Top terms of GO analysis and KEGG enrichment with thresholds of $p$-values $\leq 0.05$ were chosen in functional annotation clustering.

Literature Verification

Literature retrieval was used to excavate and analyze the active components and possible mechanisms of CCHM in the treatment of METH addiction. This may provide literature verification for prediction results of the network pharmacology.

Western Blotting Analysis

Mice were sacrificed by cervical dislocation under anesthesia after behavioral testing. Protein extracts in nucleus accumbens and prefrontal cortex were isolated, lysed, and the supernatant was collected after centrifugation. BCA Protein Assay Kit was used to detect the concentration of protein extracts. Protein samples were separated and electroblotted onto the PVDF membrane. After blocking, membranes were incubated with primary antibodies followed by a horseradish peroxidase conjugated secondary antibody. Enhanced chemiluminescence was used to visualize the band, and the intensity of each sample was determined quantitatively using Image J. Actin was used as loading control, and the results were normalized to the control group.

Molecule Docking of Main Active Compounds of Candidate Chinese Herbal Medicine With Hub Targets

The amino acid sequence of the two hub targets D(2) dopamine receptor (DRD2) and 5-hydroxytryptamine receptor 3A (HTR3A) were downloaded from the PDB protein structure database. Furthermore, AutoDock v.4.2.6 was used to add the hydrogens, calculate charges, and merge the non-polar hydrogens of protein structure. Next, the protein file was saved in PDBQT format. The three-dimensional (3D) structures of main active compounds were downloaded from the PubChem structure database. The SDF format of main active compounds was converted to PDB format using OpenBabel 2.4.1. AutoDock v.4.2.6 was used to convert the PDB format of resveratrol to PDBQT format and for molecular docking of main active compounds with hub targets. PyMol software was used to conduct a visual analysis of docking results.

Statistical Analysis

Statistical analysis was processed with SPSS 23.0 and Prism 8 software. Data were shown as the mean ± SE and analyzed using one-way ANOVA and LSD post-hoc test. Differences between groups were considered to be statistically significant if values of $p < 0.05$.

RESULTS

Candidate Chinese Herbal Medicine Inhibited Methamphetamine-Induced Conditioned Place Preference Behavior in Mice

Conditioned place preference, a classic model widely used to assess drug-seeking behaviors, was used in this study to assess the preference for visual and tactile cues associated with METH after being administered with varying doses of CCHM. As shown in Figure 1B, mice in the control group did not produce motivational effects before or after place conditioning, which set the baseline measurement for the CPP test. METH administration at 2 mg/kg significantly increased preference of white chamber compared to the controls $(p < 0.01)$. As shown in Figure 1C, compared to the METH group, pretreatment of CCHM reduced METH-induced motivational effects $(p < 0.05)$, and CPP scores decreased with the increase of CCHM dose. Taken together, the results suggested that 2 mg/kg of METH was sufficient to induce CPP behaviors, and the co-administration of CCHM attenuated the acquisition of METH-induced CPP in mice.

1. http://bioinformatics.psb.ugent.be/webtools/Venn/
2. http://string-db.org/
3. https://david.ncifcrf.gov/
4. http://www.rcsb.org/
5. https://pubchem.ncbi.nlm.nih.gov/
6. 10.1007/s10934-018-1183-5
7. https://www.frontiersin.org/articles/10.3389/fnins.2018.01183/full
8. 10.3389/fnmol.2020.00042
9. https://bioinformatics.psb.ugent.be/webtools/Venn/
Identification of Bioactive Compounds and Possible Targets in Candidate Chinese Herbal Medicine

A total of 163 compounds in CCHM obtained from TCMSP database met the requirement of OB ≥ 30%, DL index ≥ 0.18 and BBB ≥ −0.3, and 48 of which belong to C. yanhusuo, six to P. cocos, 16 to C. pilosula, 69 to Licorice, four to A. macrocephala Koidz, 12 to A. membranaceus, two to A. sinensis, and six to American ginseng. After eliminating the overlaps, 151 compounds were chosen as candidate bioactive compounds for further analyses, and the detailed information was shown in Table 1 and Supplementary Material 1. Among the 151 candidate bioactive components, 2,492 possible protein targets were retrieved from the TCMSP database. The detailed information was shown in Supplementary Material 2. After eliminating the overlaps, 138 protein targets were obtained for further analyses.

Targets Identification of Candidate Chinese Herbal Medicine on Methamphetamine Addiction

Sixty METH addiction-related targets were collected from OMIM, 356 targets were collected from the Genecards database, and 11 targets were obtained from Drugbank database. After eliminating the overlaps, 392 targets were obtained for further analyses. The detailed information was shown in Supplementary Material 3. Then, these protein targets of CCHM were mapped with METH addiction-related targets using the Bioinformatics and Evolutionary Genomics website. As a result, 43 proteins were selected, corresponding to 123 active ingredients of CCHM and the detailed information of 43 proteins was shown in Supplementary Material 4.

Construction of “Herb-Compound-Target” Network

Cytoscape software was used to construct the of “herb-compound-target” network, as shown in Figure 2. Among these bioactive compounds, the top 10 compounds were 7-O-Methylisoumorculatol, 7-Methoxy-2-Methylisoflavone, Formononetin, Beta-Sitosterol, S-Scoulerine, Isocorypalmine, Leonticine, Medicarpin, Licochalcone, and Hyndarin. The detailed information was shown in Table 2. These bioactive components may be the main active compounds of CCHM in treating METH addiction.

Gene Ontology and Pathway Enrichment Analysis

To identify the biological characteristics of putative targets of CCHM on METH addiction in detail, the GO and KEGG enrichment of involved targets were conducted via the functional annotation tool of DAVID Bioinformatics Resources 6.8. There were 116 terms of biological process (BP), 20 terms of cellular component (CC), and 31 terms of molecular function (MF) in total, which met the requirements of Count ≥ 2 and p-value ≤ 0.05. The detailed information was shown in Supplementary Material 5. The top 10 significantly enriched terms in BP, CC, and MF categories were shown in Figure 3A, which indicated that CCHM may regulate neuron functions via adrenergic receptor signaling pathway, dopamine neurotransmitter receptor activity, dopamine binding to exert its therapeutic effects on METH addiction. To explore the underlying involved pathways of CCHM on METH addiction, KEGG pathway analysis of involved targets was conducted. The detailed pathway information of CCHM on METH addiction was shown in Supplementary Material 6. The top 15 significantly enriched pathways were shown in Figure 3B. The neuroactive ligand-receptor interaction pathways exhibited the largest number of involved targets (21 counts).

Screen of Hub Genes

The PPI network of possible targets involved in the treatment of CCHM on METH addiction was constructed using STRING (Figure 4A). There were 43 nodes and 200 edges, and the average node degree was 9.3. The average local clustering coefficient was 0.637, and the p-value of PPI enrichment was less than 1.0e-16. To obtain the hub genes in the PPI network, these node pairs were entered into the Cytoscape software. The scores of nodes were calculated by cytoHubba, and the top 10 hub genes were shown in Figure 4B.

The hub gene symbols, full names, and functions were shown in Table 3. These 10 key targets corresponded to 75 major active components of CCHM. The network diagram of “herb-major active component-hub target” constructed by Cytoscape software was shown in Figure 5.

Gene Ontology and Pathway Enrichment Analysis of Hub Genes

The GO analyses and KEGG enrichment of hub genes were conducted via DAVID 6.8. There were respectively 51 BP, 5 CC, and 10 MF terms in total, which met the requirements of Count ≥ 2 and p-value ≤ 0.05. The detailed information was shown in Supplementary Material 7. The top enriched terms in BP, CC, and MF categories were shown in Figure 6. Among them, the results of BP (Figure 6A) showed that the treatment of METH addiction by CCHM was mainly related to adenylate cyclase-activating adrenergic receptor signaling pathway, response to cocaine and dopamine catalytic process. The results of CC
(Figure 6B) showed that its treatment was mainly related to plasma membrane, axon, axon terminus, and so on. The results of MF (Figure 6C) showed that items including dopamine binding, protein homodimerization activity, and dopamine neurotransmitter receptor activity had close relationship with the treatment of METH addiction by CCHM, and the genes involved were shown in Table 4. There were 11 KEGG pathways met the requirements of Count ≥ 2 and p-values ≤ 0.05. The detailed pathway information of hub genes was shown in Supplementary Material 8 and Figure 6D, which indicated that CCHM may...
FIGURE 3 | Clustering analysis of possible targets of candidate Chinese herbal medicine (CCHM) in the treatment of methamphetamine (METH) addiction. (A) Gene ontology (GO) analysis of the possible targets. (B) Kyoto encyclopedia of genes and genomes (KEGG) enrichment analysis of the possible targets. Counts referred to the number of the enriched targets.

FIGURE 4 | Protein-protein interaction (PPI) network and Hub gene screening map of targets. (A) PPI network exported from STRING database. (B) Key subnetwork of top 10 nodes analysed by CytoHubba.
regulate METH addiction via dopaminergic synapse, cocaine addiction, neuroactive ligand-receptor interaction, amphetamine addiction, and serotonergic synapse.

**Literature Verification**
It was found that Scoulerine could significantly inhibit the CPP behavior and alleviate the anxiety-like behavior caused by METH (Mi et al., 2016). Hyndarin, also known as L-tetrahydropalmatine, had significant inhibitory effects on locomotor sensitization, CPP behavior, self-administration behavior, and relapse behavior induced by METH (Zhao et al., 2014; Gong et al., 2016; Su et al., 2020), and hyndarin could also improve METH-induced learning and memory impairment (Cao et al., 2018). Studies have shown that hyndarin could improve METH-induced locomotor sensitization by regulating the activity of 5-HT neurons and the expression of dopamine D3 receptor (Yun, 2014b). These literature reports further validated the network pharmacological prediction results. The relationship between the other eight compounds and METH addiction has not been reported.

**Protein Expression Validation of Predicted Target Genes**
To further verify the effects of CCHM on the expressions of the potential targets identified via network pharmacology, the expression of DRD2 and HTR3A in nucleus accumbens and prefrontal cortex, which play important roles in the reward system, were examined using Western blotting. As shown in Figure 7, compared with the control group, the protein level of DRD2 in nucleus accumbens was significantly decreased in the METH group, while the protein expression of HTR3A was greatly increased. Furthermore, pretreatment with CCHM could prevent METH-induced changes of DRD2 and HTR3A protein expressions in nucleus accumbens. Similar results were also found in the prefrontal cortex (Figures 7B, D). These results validated that CCHM may inhibit the METH induced abuse mainly through dopaminergic synapse and serotonergic synapse pathways.

**Molecular Docking Main Active Compounds and Hub Targets**
To screen the main active compounds in CCHM, the docking results of 10 major active components of CCHM acting on DRD2 and HTR3A were shown in Table 5 and Figure 8. Results of molecular docking showed that eight of the 10 major active components had good binding abilities with the potential targets of methamphetamine treatment (DRD2 and HTR3A), and β-Sitosterol, scoulerine, isocorypalmine, medicarpin, and hyndarin had better binding activities with the two target proteins, and the molecular conformational binding was stable, which may affect the downstream signaling pathway to participate in the treatment of METH addiction by binding with the two targets.

**DISCUSSION**
Methamphetamine is widely abused worldwide, and effective clinical treatment strategy for METH abuse is still urgently needed. CCHM is a traditional Chinese medicine developed for the treatment of METH addiction, and our previous results showed that CCHM could significantly inhibit the acquisition and expression of METH-induced behavioral sensitization (Can et al., 2019), indicating that CCHM had the potential to ameliorate METH addiction. In this study, we found that pre-treatments of CCHM could significantly inhibit METH-induced conditional place preference, which further verified that CCHM had the potential to be developed as a candidate drug for the treatment of METH addiction. However, as a multi-component Chinese herbal compound, the pharmacodynamics basis and mechanism of CCHM are still unclear. In this study, GO and KEGG analysis revealed that CCHM could regulate dopaminergic synapse, serotonergic synapse, and neuroactive...
ligand-receptor interaction pathways. Among 10 hub targets out of 43 genes by PPI analyses, we focused on two most significant genes DRD2 and HTR3A, and performed molecular docking to verify interactions between active compounds of CCHM and DRD2 (or HTR3A). Our results demonstrated the effectiveness of CCHM in the treatment of METH addiction from bioinformatics and experimental perspectives, and provided more evidence for its clinical application.

Long-term abuse of METH would damage the nervous system, resulting in rapid cognitive decline, anxiety, mental disorders, hyperalgesia, depression, etc. (Prakash et al., 2017). At the same time, long-term use of METH also inhibited immune function and increased susceptibility to infection (Harms et al., 2012; Peerzada et al., 2013). CCHM consists of eight Chinese medicinal herbs. C. yanhusuo had the effect of promoting blood circulation and relieving pain (Ingram, 2014; Zhou et al., 2019). A. macrocephala Koidz, P. cocos, American ginseng, Licorice, and C. pilosula could improve the body immunity and reduce the damage of METH to the central nervous system (Fu et al., 2016; Tian et al., 2019; Bai et al., 2020). A. sinensis and A. membranaceus had good sedation and tranquilizing effects (Chen et al., 2004; Park et al., 2009). Therefore, it is reasonable to speculate that different herbs in CCHM may play an effective synergistic role in regulating METH addiction and its various complications, and the cooperation among different components could be achieved from the distinct mode of actions to achieve a complementary pharmacological synergy.
In the current study, the components in CCHM with OB ≥ 30%, DL index ≥ 0.18 and BBB ≥ −0.3 were considered pharmacokinetically active as they are possibly absorbed and distributed in human brain. 123 components and 43 targets of CCHM for the treatment of METH addiction were screened by databases. Compounds with high-degree may account for the major therapeutic effects of CCHM on METH addiction. Ten major active components including Beta-Sitosterol, S-Scoulerine, Isocorypalmine, Leonticine, Hyndarin, and 10 hub targets including DRD2, HTR3A, and MAOA were screened out. Results of hub targets GO and KEGG enrichment analysis showed that dopaminergic synapses, cocaine addiction, neuroactive ligand-receptor interaction, alcoholism, and serotonergic synapse were the main possible pathways involved in the CCHM treatment of METH addiction.

The neural circuits that project from dopamine neurons in the ventral tegmental area to the nucleus accumbens and frontal cortex are called the mesolimbic dopamine system (MLDS). Numerous studies have identified the MLDS as a common reward circuit and main neuroanatomical basis for the rewarding or positive reinforcement of addictive drugs, involving in drug-induced reinforcement, pathological memory, craving, and relapse (Peters et al., 2021). Addictive substances affect the release of monoamine neurotransmitters such as dopamine, adrenaline and 5-hydroxytryptamine by acting on the reward system, and regulate the excitability of neurons. Under physiological conditions, the presynaptic release and reuptake of dopamine in dopaminergic neurons are in equilibrium (Volkow et al., 2017; Limanaqi et al., 2018). METH enters neurons through dopamine transporter, spurs presynaptic dopamine release and increases the concentration of dopamine in the synaptic cleft. Moreover, it could replace endogenous dopamine vesicles via combination of monoamine transporter in neurons, inhibit dopamine uptake, and promote the interaction of dopamine and receptors to produce neuroadaptive and addictive behaviors (Krasnova and Cadet, 2009; Shin et al., 2017). Michael et al. found that intravenous METH injection could significantly increase the levels of dopamine in the nucleus accumbens, and the locomotor
Herbal Medicine Alleviates Methamphetamine Addiction

FIGURE 7 | Candidate Chinese herbal medicine (CCHM) administration inhibited methamphetamine (METH)-induced expression changes of D(2) dopamine receptor (DRD2) and 5-hydroxytryptamine receptor 3A (HTR3A) in the nucleus accumbens and prefrontal cortex. (A) Representative bands of DRD2 and HTR3A in the nucleus accumbens. (B) Representative bands of DRD2 and HTR3A in the prefrontal cortex. (C) METH significantly decreased expression of DRD2 and increased expression of HTR3A, while CCHM pretreatment could greatly upregulate DRD2 expression and downregulate HTR3A expression. (D) The expression of DRD2 notably decreased and the level of HTR3A increased in METH group, which could be reverse by CCHM pretreatment. n = 3 for per group, *p < 0.05, **p < 0.01 compared to the control group, #p < 0.05, ##p < 0.01 compared to the METH group.

activities of rats was consistent with the change of dopamine levels in the nucleus accumbens (Baumann et al., 2011), and reducing dopamine levels in the nucleus accumbens could reduce the locomotor activities of addicted mice (Yun, 2014a). These studies suggested that dopaminergic system played important roles in METH-induced addictive behaviors. Among the 10 hub targets screened in this study, SLC6A3 is a dopamine transporter, which regulates the reuptake of dopamine, and plays a role in maintaining the homeostasis of dopamine in the synaptic cleft. The expression of SLC6A3 in the striatum of METH abusers was significantly decreased, accompanied with reduced learning and memory ability and motor ability (Volkow et al., 2001b,c). After 4 weeks of drug withdrawal, the expression of dopamine transporter in the striatum of abusers gradually recovered, but it was still significantly lower than that of healthy subjects (Yuan et al., 2014). Dopamine transporter inhibitors such as JHW007 and tetrabenazine significantly reduced relapse and locomotor sensitization in METH-addicted rats (Meyer et al., 2011; Ferragud et al., 2014). DRD2 and DRD3 belong to dopamine receptors. DRD3 antagonists could significantly inhibit the reinforcement of METH and relapse (Chen et al., 2014; Sun et al., 2016). METH abuse has been associated with DRD2 and DRD3 deficits in the caudate nucleus and nucleus accumbens, and DRD2 receptor availability was lower in the METH abusers and may mediate impulsive temperament and thereby influence addiction (Volkow et al., 2001a; Lee et al., 2009; Groman et al., 2012; Wang et al., 2012; Okita et al., 2018). Animal experiment also confirmed that mRNA expression of DRD2 was significantly decreased after METH administration (Landa et al., 2012), suggesting poor DRD2 function may be a biomarker that predicts a greater likelihood for relapse. Therefore, we focused on the expression of DRD2 in mesolimbic dopamine system to further verify the results of bioinformatics analysis. Consistent with previous literatures, our results also proved that DRD2 in nucleus accumbens and prefrontal cortex was significantly decreased in METH group, while CCHM treatment could increase the levels of DRD2. Our findings combined with previous reports further verified the vital role of dopamine system in the prevention and treatment of METH addiction.

In addition to the dopamine system, 5-hydroxytryptamine, adrenaline system and their related metabolic enzymes are also the hub targets of CCHM in the prevention and treatment of METH addiction. MAOA and MAOB specifically

| Molecule Name          | Binding energy with DRD2/KJ mol⁻¹ | Binding energy with HTR3A/KJ mol⁻¹ |
|------------------------|-----------------------------------|-----------------------------------|
| 7-O-Methyllisomucronulatol | −2.77                             | −2.92                             |
| Formononetin           | −1.43                             | −3.52                             |
| Beta-Sitosterol        | −2.94                             | −4.07                             |
| (S)-Scoulerine         | −3.3                              | −3.96                             |
| Isocorypalmine         | −3.42                             | −3.6                              |
| Leonticine             | −1.87                             | −2.84                             |
| Medicarpin             | −3.09                             | −4.28                             |
| Hyndarin               | −3.54                             | −3.91                             |
oxidize and inactivate monoamine transmitters, and MAOA gene polymorphisms were associated with susceptibility to METH-induced mental disorders (Nakamura et al., 2009). ADRA2A, ADRA2C, and ADRA1B are adrenergic receptors. METH could increase the expression of ADRA2A in the hippocampus (Nishio et al., 2002), and downregulation of ADRA2A inhibited METH-induced hyperactivity (Nishio et al., 2003), and knockdown of ADRA1B had a protective effect on METH-induced neurotoxicity (Battaglia et al., 2003). It was reported that karbalatin, an inhibitor of the acetylcholinesterase AChE could also reduce METH self-administration behavior (De La Garza et al., 2008), suggesting that acetylcholine may be involved in METH addiction. HTR3A is one of the 5-hydroxytryptamine receptors. Electrophysiological results showed 0.1 µM METH increased 5-HT-induced inward peak current ($I_{5-HT}$) in oocytes expressing HTR3A receptor, while high dose METH inhibited $I_{5-HT}$, and HTR3A receptor antagonist MDL72222 had an inhibitory effect on the acquisition and expression of METH locomotor sensitization (Yoo et al., 2006). HTR3A receptor antagonist ondansetron had a protective effect on the neurotoxicity caused by METH (LaFuente et al., 2018). Among all the hub genes, HTR3A has the highest degree score in the cytoHubba analysis, therefore, the expression of HTR3A in mesolimbic dopamine system was also detected. Our results demonstrated that METH addiction significantly increased HTR3A expression in nucleus accumbens and prefrontal cortex, while CCHM treatment decreased the levels of HTR3A while inhibiting METH addiction, which were consistent with results in previous literatures. These findings further validated that CCHM may inhibit the METH induced abuse mainly through regulating serotonergic system.

In this study, 10 major active compounds were molecularly docked with DRD2 and HTR3A, two receptors with higher degrees in the hub targets. The results showed that most active compounds could effectively bind to DRD2 and HTR3A receptors. Among them, β-Sitosterol, Scoulerine, Isocorypalmine, Medicarpin, and Hyndarin had lower binding energy with DRD2 and HTR3A. These results were consistent with previous studies which showed Scoulerine and Hyndarin had the potential to inhibit METH addiction, and Isocorypalmine could reduce the behavioral sensitization and reward of cocaine in mice by acting on dopamine receptors (Xu et al., 2013). To the best of our knowledge, there are no related literature reports on β-Sitosterol and Medicarpin and drug addiction, and our results suggested they might be novel potential molecules for the treatment of drug addiction. Based on literature verification and results of molecular docking, β-Sitosterol, Scoulerine, Isocorypalmine, Medicarpin, and Hyndarin may be potential active components for the treatment of CCHM on METH addiction, and the underlying mechanism of action may be through DRD2 and HTR3A signaling pathways.

**CONCLUSION**

To sum up, the pharmacological mechanism by which CCHM inhibited METH addiction was investigated based on the network pharmacology analysis, literature, and experimental validation. The results of the current study have shed light on further research on the pharmacodynamics and molecular mechanism of CCHM in the treatment of METH addiction and related mental disorders. In the future, in vivo and in vitro experiments are still needed to systematically verify the main active components and possible key targets in detail. The potential clinical therapeutic effects of CCHM may benefit from further studies.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

**ETHICS STATEMENT**

The animal study was reviewed and approved by the Ethics Committee of Jianghan University.
AUTHOR CONTRIBUTIONS
QR, LC, and YW designed the study, analyzed the data, and prepared the manuscript together. QR, XT, CL, and QX performed the experiments. All authors have read and approved the final version of the manuscript.

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SUPPLEMENTARY MATERIAL
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