

ORIGINAL RESEARCH

Exploration in the Therapeutic and Multi-Target Mechanism of Ketamine on Cerebral Ischemia Based on Network Pharmacology and Molecular Docking

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Background: Ketamine is famous for its dissociative anesthetic properties. It is also analgesic, anti-inflammatory and anti-depressant, and even has a cerebral protective effect. We searched the evidence of the correlation between ketamine target and clinical efficacy and utilized network pharmacology to gather information about the multi-target mechanism of ketamine against cerebral ischemia (CI). We found that ketamine's clinical significance may be more extensive than previously thought.

Methods: The drug target of ketamine and CI-related genes were predicted by SwissTargetPrediction, DrugBank, PubChem, GeneCards and DisGeNET databases. The intersection of ketamine's drug-targets and CI-related genes was analyzed by using GO and KEGG. We predicted the molecular docking between the potential target and ketamine.

Results: The results indicated that the effect of ketamine on CI was primarily associated with the target of α-synuclein (SNCA), muscarinic acetylcholine receptor M1 (CHRM1) and nitric oxide synthase 1 (NOS1). It principally regulates the signal pathways of circadian transmission, calcium signaling pathway, dopaminergic synapse, cholinergic synapse and glutamatergic synapse. Molecular docking analysis exhibited that hydrogen bond and Pi-Pi interaction were the predominant modes of interaction.

Conclusion: There are protein targets affected by ketamine in the treatment of CI. Three pivotal targets involving 298 proteins, SNCA, CHRM1 and NOS1, have emerged as multi-target mechanisms for ketamine in CI therapy. Similarly, this study also provides a new idea for introducing network pharmacology into the evaluation of multi-targeted drugs for CI and cerebral protection.

Keywords: ketamine, cerebral ischemia, treatment target, network pharmacology

Introduction

Cerebral ischemia (CI) is a widespread acute cerebrovascular disease, which is the second leading cause of death in the world, and China has the highest burden of stroke in the world. Ischemic stroke is characterized by high incidence rate, high disability rate and high mortality rate. The critical aspect of treatment is to reduce or prevent the nerve injury caused by CI. The neuroprotective effects of CI have been studied for more than half a century, especially in the past 30 years. Nevertheless, the really effective method is still under exploration. Therefore, it is imperative to seek new treatment methods.

Ketamine is a classic intravenous anesthetic. Its anesthetic and analgesic properties are typically attributed to its direct inhibition of N-methyl-D-aspartate receptor (NMDAR).⁹ Not only that but it also has affinity for many other receptors, including γ-aminobutyric acid, dopamine, serotonin, opioid and cholinergic receptors, as well as voltage-gated sodium and hyperpolarized cyclic nucleotide gated channels. Hence, its clinical significance may be broader than previously thought.^{10,11} In recent years, ketamine has made significant breakthroughs as an antidepressant.^{12,13} In the treatment of CI, although previous studies have demonstrated that ketamine can ameliorate the behavioral and

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neuroprotective effects on animal models.¹⁴ It may be achieved by affecting the NF-κB signaling pathway, reducing TNF-α, IL-6 and increasing the content of IL-10.¹⁵ It may also activate mTOR signal transduction pathway through p-mTOR and down-regulate the expression of key autophagic proteins (such as LC3 and beclin-1), while attenuating secondary brain injury and reducing inflammatory response.¹⁶ Nevertheless, the precise mechanism of ketamine for CI is not known.

Along with the rapid development of bioinformatics, systems biology and pharmacology, web-based drug discovery and evaluation databases are considered to be an effective and cost-effective method for drug development. The combination of online pharmacology and bioinformatics tools has broadened the understanding of potential drug targets and their intersection with disease-specific key genes. Experimental data on gene, protein and metabolic interactions are combined with clinical knowledge of disease and pharmacology to explore new therapeutic approaches, which provides a macroscopic way to understand and treat complex diseases. Understanding pharmacokinetic and pharmacodynamic models will accelerate the discovery of new drugs with better therapeutic outcomes by combining them with pathway and network analysis. These approaches may mitigate the risk of single-target and symptom-based drug discovery and treatment failure in the future. Along the systems of the single-target and symptom-based drug discovery and treatment failure in the future.

In this paper, we applied a network pharmacology approach to integrate the multi-target mechanism of ketamine against CI, and further explore how ketamine can be effective against CI and facilitate new drug development using evidence of the relevance of ketamine targets to CI disease targets. In general, the interpretation of Ketamine's network pharmacology will hopefully lead to the development of new drugs suitable for the clinical treatment of CI disease.

Materials and Methods

Statement of Human Data

In this study, the databases containing human data include SwissTargetPrediction, ^{25,26} DrugBank, ²⁷ PubChem, ²⁸ PDB²⁹ ZINC³⁰ GeneCards and DisGeNET. ^{31,32} This study made use of data from public datasets and was approved by the human Experiment Ethics Committee of the First Affiliated Hospital of Kunming Medical University.

Identification of Ketamine Target Genes

In order to avoid missing all the potential target proteins of Ketamine as far as possible, we searched for "ketamine" in three databases: SwissTargetPrediction, ^{25,26} DrugBank²⁷ and PubChem. ²⁸ All the results were limited to "*Homo sapiens*", and 39 human target genes that might be targeted by ketamine were obtained. The 39 target genes were uploaded to the string database for retrieval, and the species was defined as "*Homo sapiens*", and the protein–protein interaction (PPI) network map was obtained. The node1, node2 and combined score information from the file exported from string database were imported into the software of Cytoscape 3.8.0^{33,34} for visual analysis, and the network analysis results were obtained. Further, set the node size and color reflection (degree) value, the thickness of the edge reflects the combined score size, and finally get PPI network.

Identification of CI-Related Genes

GeneCards and DisGenNET databases^{31,32} were used to identify CI-related genes, and a total of 2929 results were identified in GeneCards database and 358 results were identified in DisGeNET database. After screening and deleting repetitive target genes, 2930 CI-related genes were obtained.

Identification of Related Targets of Ketamine and CI

CI-related genes (2930) and Ketamine targets (39) were uploaded to Venny 2.1.0 software to obtain the intersection map of Ketamine target and CI target, so as to obtain the potential target of Ketamine for CI treatment.

Screening of Key Target Proteins and GO and KEGG Analysis

The BisoGenet application in Cytoscape was used to build a PPI network of mutual targets between CI and ketamine. The key targets were screened by the scores of "degree", "intermediate" and "intimacy" calculated by double genome. GO and KEGG enrichment analysis was performed in the Metascape database to identify the molecular function and systematic involvement of the target gene. The common targets of ketamine and CI were imported into Metascape database. Biological function (BP), molecular function (MF), cell composition (CC) and KEGG were checked respectively. Data with P < 0.05 were selected from BP, CC, MF and KEGG values, and then input into R software to draw GO analysis diagram and KEGG pathway enrichment analysis bubble diagram, so as to further explore the potential mechanism of ketamine in CI treatment. 35,36

Molecular Docking

Three-dimensional (3D) structure of α-synuclein (SNCA; PDB ID: 3Q29), nitric oxide synthase 1 (NOS1; PDB ID: 6PNA) and muscarinic acetylcholine receptor M1 (CHRM1; PDB ID: 6WJC) was retrieved from RCSB PDB database.²⁹ Using Autodock Vina software,³⁷ water molecules were removed, proteins were separated, non-polar hydrogen was added, the Gasteiger charge of the structure was calculated, and saved as a PDBQT file. The two-dimensional (2D) and 3D structures of Ketamine (PubChem CID: 3821) were downloaded using the PubChem database.²⁸ The 2D structures were converted to PDB format by Chem3D processing and saved as docking ligands in Autodock Vina software in PDBQT format. According to visually inspected and docking score, SNCA, NOS1 and CHRM1 were used as receptors, and ketamine was used as ligand. The conformation with the best affinity was selected as the final docking conformation, and PyMOL was used to analyze the docking results. The higher the absolute value of the binding free energy, the greater the stability of a protein–ligand complex.

Results

The Expected Target of Ketamine and Its Network

The 2D and 3D structures of Ketamine (PubChem CID: 3821) are shown in Figure 1A. For the purpose of investigating the target proteins of ketamine, we used SwissTargetPrediction, DrugBank and PubChem databases for prediction. Then, 39 target proteins were accessed by taking the intersection of the targets from these databases (Figure 1B). The three predominant types of these targets are ligand gated ion channels (28%), G-group protein coupled receptors (28%), and electrochemical transporters (12%) (Figure 1C). According to PPI network (Figure 2A), the critical potential target proteins with more interaction (binding fraction) with other targets were screened by using Cytoscape software. The target proteins located in the center of PPI network are 5-hydroxytryptamine receptor 1A (HTR1A), dopamine D2 receptor (DRD2) and 5-hydroxytryptamine receptor 3A (HTR3A) (Figure 2B). The top 15 target genes were mapped in R software according to the interactive data of PPI (Figure 2C).

Analysis of Gene Topological Networks Related to CI and Ketamine

To obtain the critical genes in CI, we performed prediction using GeneCards and DisGeNET databases. The result indicated that a total of 2930 CI-related human genes were identified from the database. Using the Wayne map, there are 30 intersecting genes between ketamine targets and CI-related genes (Figure 3A). The information of 30 intersecting genes was shown in Figure 3B. In a PPI network, the degree center (DC) of a node is simply the number of edges it has. Betweenness centrality (BC) captures the position of a given node in relation to each other. The combination of DC and BC has been shown to be effective for screening reliable and important proteins. Subsequently, the PPI network of 30 target proteins was carried out by Cytoscape software and its BisoGenet application program. An extended network of 296 proteins associated with 30 target proteins was obtained. As shown in Figure 4A, the extended network has a total of 296 nodes (targets) and 1871 edges (interactions). In order to identify the most important nodes in these 296 proteins, the DC > 18 (twofold median) was used as the primary screening criteria, and 64 nodes and 520 edges were obtained (Figure 4B). Then, BC range of 0.548–38.573 was further screened. Finally, 24 related proteins and 149 correlations were obtained (Figure 4C), which may play an important role in the treatment of CI by ketamine. In addition, the first 20 of them were screened (Table 1).

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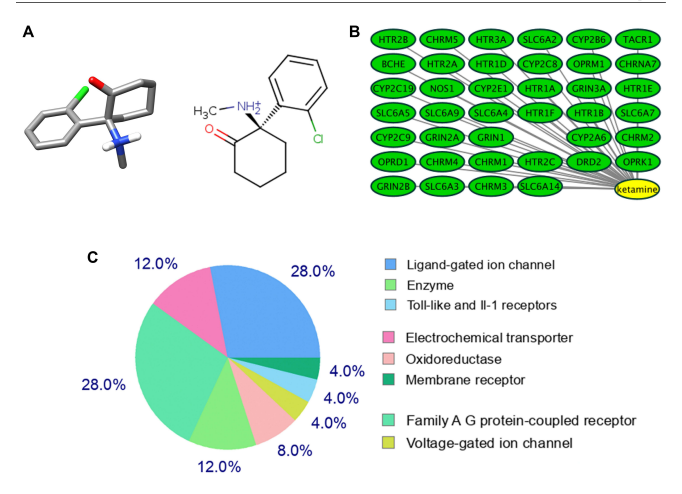


Figure 1 Target proteins predicted by the database. (A) Three-dimensional (left) and chemical structure (right) of ketamine (PubChem CID: 3821). (B) SwissTargetPrediction, DrugBank and PubChem databases predicted 39 co-potential targets for ketamine. (C) Categories of top 25 targets for ketamine.

GO and KEGG Enrichment Analysis of 298 Related Genes

To explore the functions and related pathways of 298 target genes, we used GO and KEGG for macroscopic evaluation. According to CC analysis of GO, these proteins are mainly located in glutamatergic synapses, postsynaptic density, asymmetric synapses, synaptic membranes, and postsynaptic specialization (Figure 5A); in terms of MF, they mainly focus on ion channel binding, protein C-terminal binding, glutamate receptor binding, neurotransmitter receptor activity, and G protein coupled receptor binding (Figure 5B) Their BP mainly focus on divalent inorganic cation transport, divalent metal ion transport, calcium ion transport, regulation of trans-synaptic signal and regulation of chemical synaptic transmission (Figure 5C). KEGG pathway analysis further displayed that these proteins were mainly involved in daynight transmission, calcium signaling pathway, dopaminergic synapses, cholinergic synapses and glutamatergic synapses (Figure 5D and Table 2).

Molecular Docking Analysis of Ketamine and Its Potential Protein Targets

Molecular docking is a technique that simulates the interaction between small ligand molecules and receptor protein macromolecules, allowing the binding energy between the two counterparts to be calculated to predict their affinity. The analysis of the top 20 ketamine's potential targets in Table 1 and the above key target proteins using molecular docking provides an intuitive explanation for the interaction between Ketamine and its CI-related potential target proteins. The top 10 targets for ketamine based on docking scores are shown in Table 3. Docking scores below 0 indicate spontaneous binding of two molecules, with smaller docking score leading to more stable conformations. According to visually

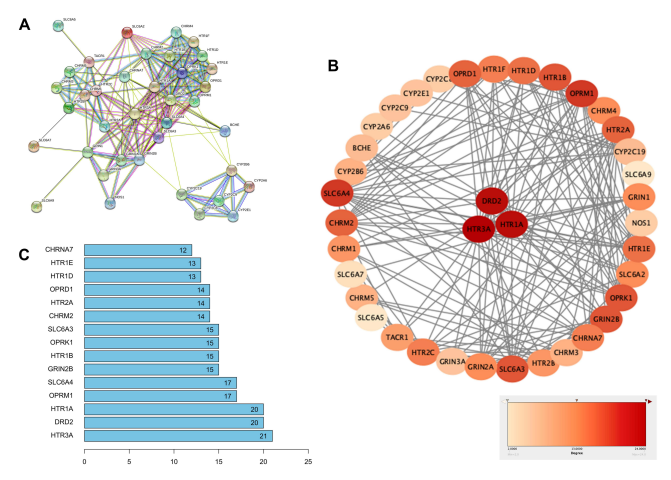


Figure 2 Establishment of PPI network for ketamine's targets. (A) PPI network of 39 potential targets was constructed via Cytoscape software. (B) Critical potential target proteins were selected according to binding fraction. (C) Top 15 critical target proteins.

inspected and docking score, we selected SNCA (PDB ID: 3Q29), NOS1 (PDB ID: 6PNA) and CHRM1 (PDB ID: 6WJC) with docking score of -9.9, -8.9 and -7.9 kcal/mol, respectively, for molecular docking with ketamine. The result confirmed that SNCA, NOS1 and CHRM1 are the most likely targets of ketamine, which have spatial adaptability, conventional hydrogen bound, Pi-alkyl interaction and Pi-Pi interaction are the main forms of interaction. As shown in Figure 6, ketamine interacted with amino acid residues TRP⁶³, TRP³⁴¹, TYR¹⁵⁶ and GLU¹⁵⁴ of SNCA, and formed hydrogen bonds with GLU¹⁵⁴, Pi-alkyl interaction with TRP³⁴¹, and Pi-Pi interaction with TRP⁶³ and TYR¹⁵⁶. As seen in Figure 7, ketamine formed hydrogen bonds with amino acid residues CYS⁴²⁰ and PRO⁵⁷⁰ of NOS1, and formed a Pi-Pi interaction with TRP⁴¹⁴. In addition, amino acid residues TYR⁴⁰⁴ and TYR⁴⁰⁸, TYR³⁸¹ and ASP¹⁰⁵ of CHRM1 formed hydrogen bonds and Pi-Pi interaction with ketamine, respectively (Figure 8). The above results indicated that ketamine has good binding activity with SNCA, NOS1 and CHRM1.

Discussion

Ketamine has been used in medicine for more than 50 years, but its mechanism is still not well understood. It has a wide range of functions and complex pharmacological effects. Clinically, ketamine is applied as an analgesic, antidepressant, anti-inflammatory drug, dissociative anesthetic, local anesthetic, neuroprotective agent, psychotic, and even neurotoxins.¹⁰ In the past, it was mainly used in anesthesia, analgesia and antidepressant therapy. Now, ketamine has been shown to have improved behavioral and neuroprotective effects in animal models of CI.³⁹ According to our network pharmacology results, SNCA, NOS1 and CHRM1 may play a key role in the treatment of Ci by ketamine.

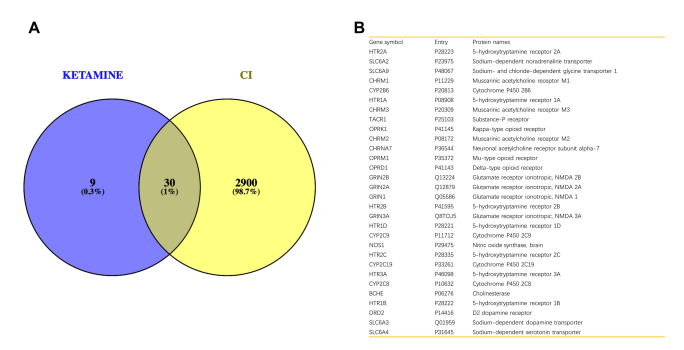


Figure 3 Identification of genes related to CI and ketamine. (A) The intersection result of ketamine targets and CI-related genes. (B) Information on gene symbol, entry and protein names of 30 intersecting genes.

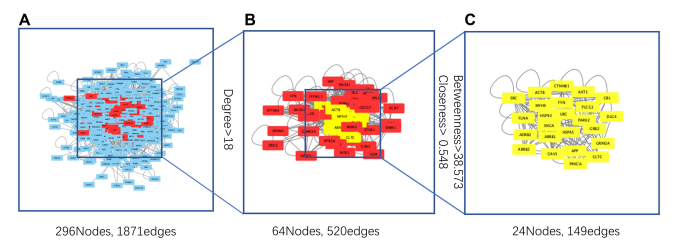


Figure 4 Analysis of gene topological networks related to CI and ketamine. (A) PPI network of intersecting target proteins. (B) PPI network based on degree center (DC) > 18 (twofold median) for the first screening. (C) PPI network based on betweenness centrality (BC) range of 0.548–38.573 for the second screening.

Target SNCA

SNCA is a neuronal protein that plays multiple roles in synaptic activity, such as regulating synaptic vesicle transport and subsequent neurotransmitter release. ⁴⁰ Mechanistically, SNCA regulates dopamine transmission by increasing the release of Ca2+ in the middle of the microregion, which is also a major component of the Louis body. The aggregation of the protein in the cell represents a histological marker of Parkinson's disease. ⁴¹ In rodents suffering from focal ischemia, *SNCA* gene knockdown significantly diminished the induction of Drp1, 3-nitrotyrosine phosphate, caspase-3 and LC-3II/I after ischemia. These results indicated that it can induce mitochondrial fragmentation, oxidative stress, and autophagy, thus promoting neuronal cell death. Therefore, it plays a key role in ischemic brain injury and is a potential therapeutic target for reducing brain injury after stroke. ⁴² If the expression of SNCA protein is stopped in the early stage after CI, brain injury can be minimized and a better functional outcome can be guided. ⁴³ Inhibition of *SNCA* gene expression in

Table I Top 20 Potential Targets Associated with CI in Ketamine Treatment

Gene Symbol	Protein Names	Degree	Betweenness
Зуппоот			
APP	Amyloid-beta precursor protein	71	111
SRC	Proto-oncogene tyrosine-protein kinase Src	61	350
HSP90AA I	Heat shock protein HSP 90-alpha	56	240
GRB2	Growth factor receptor-bound protein 2	54	98
UBC	Polyubiquitin-C [Cleaved into: Ubiquitin]	45	101
DLG4	Disks large homolog 4	45	92
ADRB2	Beta-2 adrenergic receptor	44	52
FYN	Tyrosine-protein kinase Fyn	43	112
PARK2	E3 ubiquitin-protein ligase parkin	39	134
SNCA	α-Synuclein	38	88
CTNNBI	Catenin beta-I	37	51
AKTI	RAC-alpha serine/threonine-protein kinase	36	67
CBL	E3 ubiquitin-protein ligase CBL	34	67
ACTB	Actin, cytoplasmic I	34	76
ARRBI	Beta-arrestin-I	34	49
CAVI	Caveolin-I	33	85
PRKCA	Protein kinase C alpha type	33	72
HSPA5	Endoplasmic reticulum chaperone BiP	32	60
GRIN2A	Glutamate receptor ionotropic, NMDA 2A	32	70
PLCGI	I-Phosphatidylinositol 4,5-bisphosphate	31	60
	phosphodiesterase gamma-1		

the animal model of CI can significantly decrease lesion volume, enhance exercise and facilitate cognitive function recovery.⁴⁴ During the whole CI, the ablation of *SNCA* gene can increase the formation of prostaglandins in the brain, which have a protective effect against CI. Hence, reducing the level of SNCA can effectively reduce the post-ischemic brain injury.⁴⁵ Notably, no studies have reported the progression of ketamine involvement in CI through regulation of SNCA. In this study, the network pharmacology studies we have done suggests that ketamine can bind tightly to SNCA firmly, inhibit protein function, reduce neuronal apoptosis, protect post-ischemic brain cells, and facilitate recovery of brain function.

Target NOSI

It has been demonstrated that activation of c-Jun N-terminal kinase 1/2 (JNK1/2) is a signaling pathway for early neuronal death in CI. This is because the expression of neuronal nitric oxide synthase (nNOS) is increased and more NO is produced. NO, as a free radical, can directly destroy protein structure, disrupt mitochondrial function, and inspire apoptosis. Previous studies disclosed that transient focal ischemia increases endothelial-type NOS1 in the cerebral vasculature. Threefore, inhibition of NOS1 (neuronal NOS) activity can alleviate cell death in CI. According to previous studies, intrathecal ketamine can significantly suppress the expression of NOS1 in the dorsal horn of the rat's spinal cord. Ketamine has been shown to be involved in the development of disorders such as schizophrenia and depression by regulating the expression of NOS1. Our molecular docking simulation results indicated that ketamine has strong docking ability with NOS1. It can effectively inhibit the activity of NOS1, which in turn reduces the production of NO, thus avoiding the damage to neurons by oxidative stress. Moreover, CI-reperfusion injury is associated to the calcium influx mediated by NMDA receptor, which activates nNOS and thus induces NO production. Ketamine can simultaneously inactivate NMDA receptor and NOS1, which should be one of the mechanisms by which ketamine can prevent CI injury.

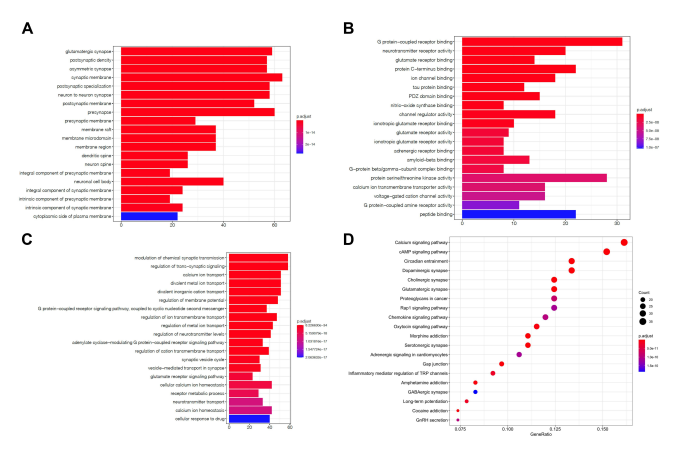


Figure 5 GO and KEGG enrichment analysis of 298 related genes. The top 20 terms of (A) CC, (B) MF and (C) BP obtained by GO enrichment analysis of 298 target genes. (D) KEGG pathway enrichment analysis results in the top 20 pathway related to 298 target genes.

Abbreviations: CC, cell composition; MF, molecular function; BP, biological process.

Target CHRMI

Ketamine can significantly inhibit CHRM1 and muscarinic signal. This effect may explain some anticholinergic effects of ketamine, including central (effect on memory and consciousness) and peripheral (significant enhancement of sympathetic excitability, bronchiectasis, mydriasis, etc). Frevious studies have demonstrated that the absence of CHRM in microglia affects CI injury. T-type calcium channel enhancer SAK3 inhibits neuronal death after transient CI via CHRM1 stimulation. Previous studies have revealed that ketamine can reduce the need for variable force drug support in patients with septic shock, and its effect may be related to the inhibition of CHRM1.

Table 2 P Values of 10 Pathways and Pathways Related to Ketamine Involvement in Treatment of CI

KEGG ID	Term	P value	Count
hsa04713	Circadian entrainment	3.86E-23	29
hsa04020	Calcium signaling pathway	2.58E-19	35
hsa04728	Dopaminergic synapse	5.02E-19	29
hsa04725	Cholinergic synapse	8.74E-19	27
hsa04724	Glutamatergic synapse	1.12E-18	27
hsa05032	Morphine addiction	6.95E-18	24
hsa04024	cAMP signaling pathway	1.82E-16	33
hsa04726	Serotonergic synapse	2.31E-15	24
hsa04540	Gap junction	8.04E-15	21
hsa05030	Cocaine addiction	6.51E-14	16

Table 3 Ketamine's Top 10 Targets Based on Docking Scores

Gene Symbol	Protein Names	Docking Score (kcal/ mol)
SNCA	α-Synuclein	-9.9
NOSI	Nitric oxide synthase	-8.9
CHRMI	Muscarinic acetylcholine receptor MI	-7.8
SRC	Proto-oncogene tyrosine-protein kinase Src	-7.7
HSPA5	Endoplasmic reticulum chaperone BiP	-7.6
HTR2A	5-Hydroxytryptamine receptor 2A	-7.4
FYN	Tyrosine-protein kinase Fyn	-6.9
HSP90AA I	Heat shock protein HSP 90-alpha	-6.8
ADRB2	Beta-2 adrenergic receptor	-6.8
ARRB I	Beta-arrestin-I	-6.7

intravenous anesthetic that can stimulate brain function and leads to a 50% increase in cerebral blood flow. At the same time, it can reduce catecholamine reuptake, increase mean arterial pressure, and make hemodynamics more stable. It can also increase heart rate and myocardial contractility by stimulating sympathetic nerve, and its cardiovascular excitation response can improve cerebral perfusion and has neuroprotective properties. Based on this, ketamine is helpful for the recovery of blood perfusion of tissue cells after CI by regulated CHRM1.

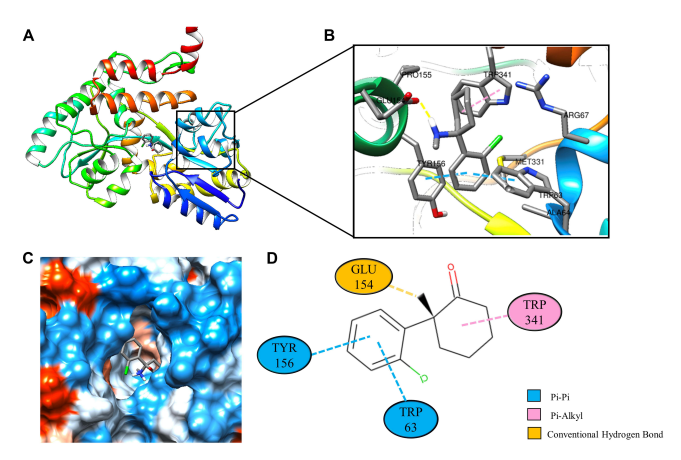


Figure 6 Molecular docking analysis of ketamine and SNCA protein. (A) Crystal structure of the SNCA (PDB ID: 3Q29) with ketamine as the ligand in the active binding site of SNCA. (B) 3D docking pattern and molecular interactions of SNCA and KET. Interactive keys are represented by dashed lines. (C) Pocket view of KET binding to SNCA. (D) Two-dimensional docking pattern of KET with amino acids TRP,⁶³ TRP,³⁴¹ TYR¹⁵⁶ and GLU. ¹⁵⁴

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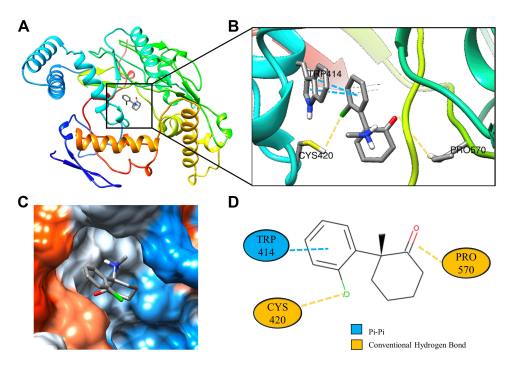


Figure 7 Molecular docking analysis of ketamine and NOS1 protein. (A) Crystal structure of the NOS1 (PDB ID:6PNA) active site-binding ligand KET. (B) 3D docking and molecular interaction of NOS1 and KET, the interaction bonds are represented by dashed lines. (C) Pocket view of KET binding to NOS1. (D) Two-dimensional docking patterns of KET and 6PNA amino acids CYS, 420 TRP414 and PRO.570

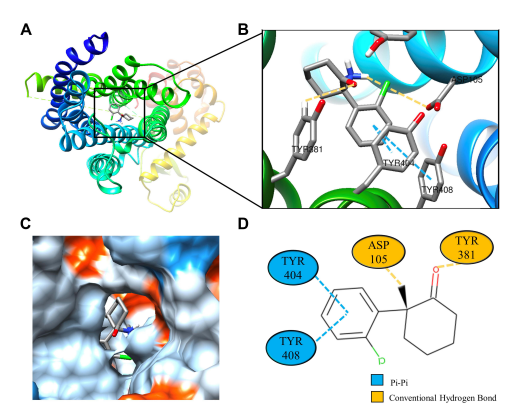


Figure 8 Molecular docking analysis of ketamine and CHRMI protein. (A) Crystal structure of the active site-binding ligand KET of CHRMI (PDB ID: 6WJC). (B) 3D docking and molecular interactions of CHRMI and KET, the interaction bonds are represented by dashed lines. (**C**) Pocket view of KET bound to CHRMI. (**D**) Two-dimensional docking pattern of KET with amino acids TYR, ⁴⁰⁴ TYR, ⁴⁰⁸ ASP¹⁰⁵ of CHRMI.

More Relevant Mechanisms

We conclude through the network pharmacology that SNCA, NOS1, and CHRM1 are the key targets of ketamine treatment CI. Molecular docking simulation has the best and most stable binding force, which is consistent with the previous research results. Ketamine even has anti-inflammatory properties, which can skew macrophages to an M2-like phenotype. Furthermore, ketamine can inhibit the activation of leukocytes in inflammatory response and reduce the production of inflammatory cytokines TNF-α and interleukin-6, thereby reducing the neurological damage caused by the inflammatory response. Previous studies have confirmed that subanesthetic doses of ketamine can activate autophagy activity and protect neurons in a mouse model of Parkinson's disease. The expression of c-Jun protein in the hippocampus of mice can be regulated to protect the mice from CI and reperfusion injury. This is in line with the results we have analyzed from KEGG and GO. According to CC analysis of GO, 296 related target genes involved in ketamine treatment CI were mainly located in glutamate synapses, postsynaptic density, asymmetric synapses, synaptic membrane and postsynaptic specialization. The analysis of KEGG pathway exhibited that these proteins mainly involved in the day and night transmission, calcium signaling pathway, dopamine synapse, cholinergic synapse and glutamate synapse, and almost all involved in the information transmission of nervous system.

Limitations and Future Directions

Ketamine has unique pharmacological advantages and multi-target characteristics that distinguish it from other traditional anesthetics. It has a wide range of clinical prospects and has already exceeded the scope of general anesthetics. Sub-anaesthetic dose of ketamine is now more widely used in the treatment of chronic pain, Alzheimer's disease, depression and CI. Nevertheless, it is not limited to this. In the future, there are more application directions to be explored. However, there is still lack of large sample, long-term and repeated clinical evidence of ketamine in the prevention of nerve injury after CI. The neurotoxic effect of ketamine is also a major obstacle limiting its clinical application, and the problem of addiction must also be taken seriously. Further research is needed to determine the selection criteria, optimal scheme and dose selection, and minimize adverse reactions to improve its clinical efficacy and safety.

Conclusion

In summary, we explored the multi-target mechanism of ketamine to treat CI through network pharmacology. The findings indicated SNCA, NOS1 and CHRM1 are the most likely targets for ketamine on treating CI. This article uses network pharmacology and molecular docking simulation methods to explore the possible mechanism of ketamine for CI treatment, which will help facilitate its clinical application in more directions.

Data Sharing Statement

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

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Disclosure

The authors report no conflicts of interest in this work.

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