IN SILICO AND IN VIVO PHARMACOLOGICAL STUDIES OF CLOZAPINE AND D-AMINO ACID OXIDASE INHIBITOR FOR COGNITIVE ENHANCEMENT

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ABSTRACT

Objective: D-amino acid oxidase inhibitors (DAAOIs) are of particular focus for cognition study. Atypical antipsychotics are known DAAO inhibitors. The present examination was done to check out the binding affinity of atypical antipsychotics by docking toward the DAAO protein; in conclusion, the picked antipsychotic drug was checked for their cognition enhancing activity with scopolamine-induced amnesia.

Methods: The crystal structure of DAAO was obtained from Protein Data Bank, the energy minimization was performed with CHARMM program, then active site prediction was made out using Ramachandran plot, and finally, docking examination was finished using AutoDock 4.2 tool. For in vivo study, the mice were divided into three groups. Group I - vehicle (Saline) treated, Group II – saline +scopolamine (1 mg/kg, intraperitoneal [i.p]) treated, and Group III - clozapine (20 mg/kg, i.p) + scopolamine (1 mg/kg, i.p).

Results: The Autodock examination shows significant binding affinity of - 5.22 for brexpiprazole and least or positive binding affinity of +1 for loperidone. Clozapine with binding energy of - 2.87 was decided for completing the in vivo cognition study. The in vivo shows up that clozapine (20 mg/kg, i.p) exhibits a change in the impairment of spatial memory.

Conclusion: The results recommend that the clozapine produces cognitive enhancement through both DAAOI and antipsychotic action. Clozapine has cognitive improvement potential, favoring its usage in reducing toxic impacts of scopolamine

Keywords: D-amino acid oxidase, Antipsychotics, Clozapine, Cognition.

INTRODUCTION

D-amino acid oxidase (DAAO) is a flavoenzyme that degrades the D-amino acids through the system of oxidative deamination. DAAO control of keeping up D-amino acid levels has been connected with a couple of physiological techniques running from hormone secretion to synaptic transmission and cognition [1]. Preclinical figures suggest that second-generation antipsychotics could diminish cognitive impairment [2]. Second-generation antipsychotics have regularly outflanked their original partners in switching or weakening phencyclidine- or methamphetamine-instigated shortfalls in inversion learning, restrictive learning, consideration, spatial memory, spatial learning, and long-haul potentiating effect in rodents and nonhuman primates [3].

In this work, we predict the three-dimensional (3D) structure and the protein-ligand and protein-protein docking of DAAO using various Bioinformatics methods. The essential purpose of our investigation was to expect the 3D structure and docking. The objective of the present examination was to clear up the participations of DAAO proteins with ligands and distinctive proteins and to perceive the relationship of DAAO to insight. Protein-protein docking and association reproduction reveal hydrogen and ionic bonds.

Scopolamine, a non-selective muscarinic receptor antagonist, causes debilitation in learning and memory by decreasing cholinergic action. On intraperitoneal (i.p.) administration of scopolamine, the cholinergic neurotransmission was blocked, provoking cholinergic brokenness and obstructed cognitive impairment in mice [4]. Starting late, it has been represented that memory impairment induced by scopolamine in mice is connected with changed cerebral amino acid levels by the protein DAAO. Along these lines, mice with scopolamine-induced memory deficiencies were used as an animal model for the screening of cognitive enhancement agents.

Here, we investigate whether the neuroprotective effect of clozapine reduced learning and memory impairment induced by a muscarinic antagonist scopolamine in mice. We assessed the effect of clozapine on scopolamine-induced learning and memory impairment with Morris water maze tests.

MATERIALS AND METHODS

In silico method

Retrieval of crystal structure and target preparation

The high-resolution crystal structure of human DAAO was retrieved from the Protein Data Bank. FASTA sequence of the retrieved structure revealed the presence of glycine at position. The target was pre-processed by standard methods before binding analysis [5].

Energy minimization [6]

CHARMM is a general and flexible program for macromolecular energy minimization and dynamic calculations that utilize both classical and quantum mechanical energy functions for molecular systems. Energy minimization of target enzyme was carried out under CHARMM27 force field. Gradient was set to 0.05.

Structural assessment of proteins [7]

3D structure and conformational stability of the protein were analyzed by means of Ramachandran plot.
Active site prediction [8]
The ligand-binding domain of the human DAAO was predicted using the Site Finder module of the molecular operating environment.

Molecular docking analysis [9]
To understand the binding conformation and affinity, antipsychotics were individually docked to the active site of human DAAO. Autodock 4.2 tool was used for molecular docking analysis. Both the receptor and ligands were prepared by the addition of hydrogen and Gasteiger charges. A grid defining the active site was constructed before running the docking simulation. Genetic algorithm was adopted for conformer search while docking.

In vivo methods

Animals
Male Swiss Albino mice (25–30 g; 24 mice) were obtained from the Animal house, TANUVAS, Chennai, and housed into four groups of six mice each. The mice were housed in cages at a distorted temperature of 25°C±2°C with a 12 h light/dark cycle. The animals had free access to standard pellet eating regimen and drinking water. Behavioral studies were performed in a silent room between 9.00 am and 11.00 am to stay away from circadian variation. The investigation was endorsed by the Institutional Animal Ethical Committee, and work was completed according to the CPCSEA Guidelines, New Delhi. XIX/VELS/PCOL/05/2000/CPCSEA/IAEC/03.10.2016

Drugs and chemicals
Scopolamine (Sigma-Aldrich), Clomaph (La Pharmaceuticals, New Delhi), and sodium chloride (Sigma-Aldrich) were utilized. Every other substance and reagents used are of analytical grade.

Experimental groups
- Group I - Saline treated (10 mL/Kg, i.p.) (Vehicle)
- Group II - Saline (10 mL/Kg, i.p.)+scopolamine (1 mg/Kg, i.p) treated,
- Group III - Clozapine (20 mg/Kg, i.p)+scopolamine (1 mg/Kg, i.p).

Learning and memory evaluation: Morris maze test [11]
Spatial learning and memory were assessed by the Morris water maze. The technique included two stages. The initial step was the place navigation test from day 1 to 4, in which the escape latency (EL) (the time required to escape onto the hidden platform) was utilized to assess learning and memory function. Mice that found the platform were permitted to stay on the platform for 20 s and were then come back to the home cage. On the off chance that mice did not achieve the platform inside 120 s, it was tenderly guided to the platform by the experimenter, where it stayed for 20 s. The last trial was viewed as the probe test. The second step was the spatial test on day 5 after evacuation of the platform and after the probe test, which was performed to test the capacity of mice to locate the expelled platform by memory

RESULTS

In silico
Energy minimization of DAAO was carried out. The initial and post-minimization potential energies are listed in Table 1.

The conformation and stability of the enzyme were analyzed in terms of its dihedral angles phi and psi using Ramachandran plot. A typical Ramachandran plot consists of a favored, allowed, and disallowed region. The Ramachandran plot of human DAAO is shown in Fig. 1.

Table 2 shows the effect of clozapine (20 mg/Kg, i.p) on (a) escape latency and (b) time spent, in the Morris water maze test. Clozapine (20 mg/Kg, i.p) was administered to the mice. The mice were then treated with scopolamine (1 mg/kg) and tested in the Morris water maze test. Probe trial sessions were performed for 60 s. The data represent mean ± standard error of mean **p<0.001 compared to scopolamine-treated mice

In vivo
The result of clozapine on scopolamine-induced memory deficit was evaluated using Morris water maze (Fig. 2). Morris water maze evaluates spatial, working, and reference memory. During this study, the escape latency time in animals administered by scopolamine was considerably reduced by clozapine-treated group (20.35±5.7**) compared with the group received vehicle only (32.21±2.69) as shown in Table 2.

**Table 1: Energy minimization of human DAAO**

| S. No | Type of variant | Pre-minimization potential energy | Post-minimization potential energy |
|-------|-----------------|----------------------------------|-----------------------------------|
| 1     | HS-DAO          | 5150.8530 kcal/mol               | −744.0501 kcal/mol                |

*Force field: CHARMM27, Gradient: 0.05, H and LP adjusted. DAAO: D-amino acid oxidase

**Table 2: Effect of clozapine on scopolamine-induced amnesia**

| S. No | Groups  | Treatment | Escape latency (sec) | Time spent in platform quadrant (sec) |
|-------|---------|-----------|----------------------|---------------------------------------|
| 1     | Group I | Saline treated (10 mL/Kg, i.p.) (Vehicle) | 22.43±2.5** | 22.43±2.5** |
| 2     | Group II| Saline (10 mL/Kg, i.p.)+Scopolamine (1 mg/Kg, i.p) | 46.39±3.8 | 46.39±3.8 |
| 3     | Group III| Clozapine (20 mg/Kg, i.p)+Scopolamine (1 mg/Kg, i.p) | 29.37±3.57 | 29.37±3.57 |
To gain better insight for the interactions between antipsychotics and DAAO, molecular docking studies were carried out. The interactions of the ligands with the active site residues (Table 3) of the target are analyzed in terms of the following parameters: Binding energy, number of hydrogen bonds established by the ligand with residues of the active site, π-π interactions, conformation oriented by the ligand within the active site, and root mean square deviation (RMSD) of the active site residues. The dock score of Autodock is reported in kcal/mol. Autodock uses the following empirical formula to calculate the free energy of binding:

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\text{Binding energy (ΔG) = Intermolecular energy + van der Waals hydrogen bond desolvation energy + electrostatic energy + total internal energy + torsional energy – unbound energy of the system.}
\]

Desolvation energy is a prime parameter that decides a molecule’s interaction with its pharmacodynamic target [5]. In the biological environment, all drug binding pockets of a target protein remain solvated, and hence, a ligand cannot as such occupy the active site unless it dislodges the water molecules. The similarity of docked structures is measured by computing the RMSD, and clusters are created based on the comparison of conformations and estimated RMSD values [12,13-16]. The docking score of selected ligands with DAO is shown in Table 4, and Fig. 3 shows the 3D docked confirmations of atypical antipsychotic agents.

Impairment of memory and learning and the foremost characteristic manifestation of psychological feature dysfunction are caused by scopolamine, a cholinergic antagonist which interferes with neurotransmitter transmission within the central nervous system [17,18]. The Fig. 2 shows the experimental study, escape latencies in Table 2 it indicates a major decrease for clozapine treated group., whereas the scopolamine-treated group exhibited a characteristic swimming behavior consisting of circling around the pool and longer escape latency indicating impairment of their spatial memory. These results suggested the cognitive enhancement of clozapine.

**CONCLUSION**

Clozapine has cognition improvement potential, approving its utilization in alleviating toxic effects of scopolamine. Clozapine being a DAAO inhibitor is an added benefit for the therapy, and it is a better insight for future cognitive studies.

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**AUTHOR’S CONTRIBUTION**

Both the authors have contributed equally to the research work.

**CONFLICTS OF INTEREST**

The authors do not have conflicts of interest.

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**Table 3: Active site analysis of DAAO**

| S. No | Type of variant | Active site residues |
|-------|----------------|----------------------|
| 1     | DAAO           | GLN37 GLN38 LYS39 LYS42 SER43 PRO44 LYS45 LEU47 VAL58 PRO59 ARG61 PHE62 ILE75 ALA81 ASP82 TYR86 |

**Table 4: Molecular docking analysis of antipsychotics with DAAO**

| Drug       | ΔG (kcal/mol) | kI (mM) | Intermolecular energy | VHBDE   | Electrostatic Energy | Total internal Energy | Torsional energy |
|------------|---------------|---------|-----------------------|---------|----------------------|-----------------------|------------------|
| Amisulpride| −3.97         | 1.23    | −6.35                 | −6.62   | 0.26                 | −1.17                 | 2.39             |
| Amoxapine  | −4.13         | 0.94    | −7.41                 | −7.83   | 0.42                 | −0.73                 | 3.28             |
| Aripiprazole| −4.25        | 0.76    | −6.34                 | −6.53   | 0.19                 | −0.23                 | 2.09             |
| Blonanserin| −3.67         | 2.05    | −5.75                 | −5.9    | 0.15                 | −0.01                 | 2.09             |
| Brexpiprazole| −5.22       | 0.14    | −7.31                 | −7.56   | 0.26                 | −0.13                 | 2.09             |
| Clozapine  | −2.72         | 10.13   | −6.0                  | −6.55   | 0.55                 | −0.95                 | 3.28             |
| Clorectine | −3.46         | 2.91    | −5.55                 | −5.62   | 0.07                 | −0.07                 | 2.09             |
| Clozapine  | −3.04         | 5.88    | −5.13                 | −5.76   | 0.63                 | −0.49                 | 2.09             |
| Clozapine  | −2.87         | 7.93    | −4.95                 | −5.55   | 0.6                  | −0.66                 | 2.09             |
| Iloperidone| 1.0           | N/A     | −4.67                 | −4.85   | 0.18                 | 0.09                  | 5.67             |

DAAO: D-amino acid oxidase, VHBDE: Van der Waals hydrogen bond desolvation energy
Fig. 3: Three-dimensional docked confirmations of atypical antipsychotic agents. (a) Amisulpride with D-amino acids oxidase (DAAO). (b) Amoxapine with DAAO. (c) Aripiprazole with DAAO. (d) Blonanserin with DAAO. (e) Brexpiprazole with DAAO. (f) Cariprazine with DAAO. (g) Caripipramine with DAAO. (h) Clozapramine with Arg16 variant. (i) Clotepine with DAAO. (j) Clotiapine with DAAO. (k) Clozapine with DAAO. (l) Iloperidone with DAAO

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