Reserpine Improves Working Memory

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Abstract

Despite exhaustive search, no drug is in sight for AD. Earlier, we reported that reserpine, an antihypertensive and antipsychotic drug, ameliorates Amyloid beta (Aβ-AD causing peptide) toxicity and confers several positive enhancements in the C. elegans model system. Here, we evaluate whether reserpine can provide protection against working memory and against AD in the mouse model. Reserpine (0.08 mg) was administered orally on alternate days to the non-Tg and accelerated Aβ deposition (at 2 months of age) and cognitive deficit (4 months of age) developing 5XFAD AD Tg mouse model expressing mutant human APP (3 familial mutations) and human Presenilin1 (2 familial mutations) in the neurons, and follow their working memory for 2 months using the spontaneous Y-maze alteration behavioral paradigm. Reserpine enhanced working memory in non-Tg mice and improved the cognitive deficit in the 5XFAD AD Tg mice. Hence, reserpine can be considered for a detailed evaluation in the 3X Tg AD mouse model and a pilot study in AD patients.

Keywords

Alzheimer’s Disease, Amyloid-β-Aβ, Reserpine, Cognitive Deficits, Transgenic Mice, Working Memory

1. Introduction

Alzheimer’s disease, the devastating neurodegenerative disease occurring at epidemic proportions (1 in 85 people) is predicted to reach ~115.4 million people in 2050 [1]. Despite screening 1000 s of drugs to treat AD [2] [3], any worthwhile drug is yet to come. Rather, many drugs, immunization with Aβ [4] for its clearance and Aβ generating gamma and beta-secretase inhibitors hardly showed any protection in clinical trials. So a potential treatment for AD is an urgent necessity.

Aβ neuritic plaques [5] and phospho-tau neurofibrillary tangles [6] [7], loss of synapses and neurons in the brain leading to cognitive deficits-progressive dementia and eventually death were observed in AD patients [8].

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Cholinergic nervous system is impaired and lost as the disease progresses. The glutamate NMDA receptor is involved in learning and memory by long term potentiation and synaptic plasticity. A modest protection against moderate and severe AD is brought about by the (NMDA) inverse antagonist memantine ([2] and references within). Acetylcholine esterase inhibitors are the more common treatment which provides minimal delay of the symptoms, but extremely expensive [2].

Majority of AD cases are sporadic. But accelerated familial AD development is noticed in patients with mutations in two genes, namely, Amyloid Precursor Protein (APP) and Presenilin 1 ([6] and references within). Several AD transgenic mice models [9] have been developed with these mutations to understand the mechanism of AD and for the development/identification of drugs, with limited success. Earlier models lacked neuronal loss found in AD patients. To fill in this gap, Oakley et al. [10] have developed a model, which expresses 5 different mutations of human APP and Presenilin 1 namely, three familial mutations (K670N/M671L (Swedish), I716V (Florida), and V717I (London)) in APP and two mutations (PS1 M146L and L286V) in Presenilin 1 (PS1). These 5XFAD AD Tg mice start showing Aβ aggregates from 2 months of age [10]. They display progressive neuronal loss and behavioral deficits [10], the two most important features in AD patients. Hence, we chose this model for our studies.

Antihypertensive drugs provide protection against Dementia. A screen of many antihypertensive drugs to reduce Aβ aggregates in a cell culture model led to the identification of Valsaratran which was later proved to provide protection in AD model mice [11], which is currently in clinical trials [11]. The Cache County study suggests that patients taking antihypertensive drugs are less likely to develop Alzheimer’s disease [12].

Reserpine is a FDA approved antihypertensive drug from the roots of *Rauwolfia serpentina*, known as *Surapaganda* being used in ayurvedic medicine in India for 1000s of years [13] to treat insanity, snake bites and as a tranquilizer. Independently, we identified that reserpine could provide high quality lifespan extension in *C. elegans* [14]. In addition, reserpine ameliorates Aβ-induced toxicity manifested as progressive paralysis, conferred stress tolerance and enhanced locomotion till late age in the *C. elegans* model [15]. The AD mouse model, Tg2576, expresses human mutant APP, which causes Aβ aggregates formation and cognitive deficits at ~11 months of age [16]. In these Tg2576 mice, reserpine reduced Aβ 42 deposits and levels in hippocampus and serum [16]. But reserpine’s effect on the major problem of AD, namely, cognitive deficit—progressive dementia or memory loss was not addressed.

Here, we address whether reserpine: 1) provides protection against Aβ toxicity induced cognitive deficits in the 5XFAD AD Tg model and 2) improves cognition in normal mice.

2. Materials and Methods

2.1. Animals and Maintenance

The mice were maintained in standard animal house facility with ad libitum access to food and water. All animal protocols were approved by the Institutional Animal Ethics Committee and in accordance with the guidelines of CPCSEA. The 5X FAD AD Tg mice [10] (obtained from Jackson laboratory, Ann Arbor, USA) were bred with wild-type non-transgenic mice to yield transgenic and non-transgenic littermates and genotyped using the standard tail DNA genomic PCR with relevant primer sets sets (APP: APP-FP (AGGACTGACCACTCGACCAG) & APP-RP (CGGGGGTCTAGTCTGCAT); Presenilin1: PS1-FP (AATAGAGAACGGCAGGCA) & PS1-RP (GCCATGAGGGCACTAATCAT)).

2.2. Reserpine Treatment

Reserpine (Sigma) (0.08 mg), dissolved in fruit vinegar and diluted with dextrose-saline was orally administered to human *app*<sup>−/−</sup>*psen*<sup>1</sup> (5XFAD)/non-transgenic mice on alternate days from 2 months of age till 5 months. This dosage and duration was arrived at based on the specific concentration at which reserpine had protective effects in *C. elegans* [14] [15] and the prescription used for hypertension [13]. For control, the vehicle, fruit vinegar, diluted with saline dextrose was administered. All the mice were age and sex matched. The treatment was done as three independent sets, with 4 or 5 mice per group with approximately equal number of males and females.

2.3. Behavioral Analyses

After one month of drug regimen, spatial working memory was tested with minor modifications [17]. This
learning task assesses hippocampus dependent spatial learning, working memory and exploratory activity. Mice were individually placed in the center of the symmetrical Y-maze consisting of three arms separated by 120 degrees (arm size: 40 cm long; 13 cm high and 10 cm wide). Mice were allowed to explore freely through the Y-maze during a 5 min session. The sequence of arms entered and total number of arms entered were recorded. Number of arm entry was considered when all the four paws were placed inside the arm. This behavior was evaluated on alternate days starting from 3 months up to 5 months of age, since the 5XFAD Tg mice shows poor performance of spontaneous alternation from the age of ~4 months due to cognitive decline as described previously [10]. The number of triads was calculated as 3 sequential entries into three different arms (i.e. 1, 2, 3 or 2, 3, 1).

### 2.4. Statistical Analysis

The data obtained was analyzed using SigmaPlot version 10.0 and the statistical significance was performed by student’s t test. All results were expressed as mean ± standard deviation. P-value < 0.01 and P-value < 0.001 were considered to be statistically significant.

### 3. Results

Reserpine, an antihypertensive drug was protective in the Aβ toxicity model in *C. elegans*. Further, reserpine reduced Aβ levels in the Tg2576 AD model mice. Here, we evaluated the crux of the disease, improvement from the cognitive deficit impairment, measured using the behavioral paradigm—spontaneous alternation in the Y maze for working memory in the 5XFAD AD Tg mouse model. They develop cerebral Aβ plagues (aggregates) rapidly at two months of age and show memory impairment in Y maze by 4 months of age. Both APP and PS1 segregate together [10] and the mice with these genes were identified.

#### 3.1. Reserpine Delays Cognitive Deficits in the 5XFAD AD Tg Mice

In order to determine reserpine’s ability to alleviate Aβ toxicity in the mouse model, in which the actual behavioral deficit in cognition could be determined, we orally administered the 5XFAD AD Tg mice with ~0.08 mg of reserpine on alternate days from two months of age up to 5 months of age. We followed the improvement in the cognitive paradigm, spontaneous alteration behavior task in the Y-maze on alternate days, usually on the next day after reserpine oral administration. The behavioral paradigm evaluation was started after one month of initiation of drug regimen and monitored for 2 months which is up to 5 months of age.

In the Y-maze paradigm, overall total movement and spontaneous arm alteration counted as triad (entry into all the three arms in a sequential manner) were measured in the 5X FAD AD Tg mice. Each trial lasted for 5 minutes. Reserpine increases the overall movement and the maximum activity is observed after one month of initiation of behavioral paradigm. The actual measure of cognitive enhancement/protection measured as increase in the number of triads entered also peaks after 20 days of start of the Y-maze behavioral paradigm. Of these, ~8 days (118 - 126 days of age) (*Figure 1*) showed the maximum protection which was statistically significant (P < 0.005/P < 0.05).

#### 3.2. Reserpine Treatment Enhances Cognition in the Normal Mice

When normal non-Tg mice were treated with the same dose of reserpine as the human mutant *app*′*psen*1+ mice they showed improvement in cognition as number of triads entered (*Figure 2*). Moreover, the enhancement was statistically significant (**—P < 0.005 and *—P < 0.05**) for a long duration of almost one month (*Figure 2*). Thus, reserpine was able to enhance normal cognition as well.

### 4. Discussion

The 5XFAD AD Tg mouse model showed enhanced spatial working memory measured by increase in the number of triads entered when they were chronically administered with low dosage (0.08 mg) reserpine orally on alternate days for a period of three months. More importantly, reserpine improved the cognition of the 5xFAD AD Tg mice at the age (*Figure 1*) where these mice are reported to show cognitive deficits. While our study was ongoing, Go *et al.* [16] reported that reserpine can help relieve AD pathogenesis in Tg2576 mice through down
regulation of $A\beta_{42}$ deposition, NGF secretion [16], but had not assessed cognitive function. Since decline in cognition is the most devastating feature of AD, here, we show that reserpine improves spatial working memory and delays cognitive deficits in the 5XFAD AD Tg mice at the low dosage of 0.08 mg that too administered only on alternate days which is in line with the low dosage of reserpine currently in use. Reserpine is effective as an antihypertensive drug at as low a dosage as 0.05 mg [13] [18]. In addition, reserpine enhances cognition in the normal mice (Figure 2). This suggests that reserpine can be considered for detailed evaluation in the 3X Tg (human mutant APP, mutant Presenilin 1 and mutant tau) AD mouse model and a pilot screen in AD patients.

In addition to dementia, due to the progressive neuronal loss, other complications like anxiety, paranoia, aggression, hallucinations, delusions, and depression and sleep disturbances make management of AD patients extremely difficult. As reserpine modulates biogenic amines neurotransmitters, the secondary problems mentioned above also could be alleviated by it. Go et al. [16], report that reserpine can significantly reduce the aggressive behavior in the Tg2576 AD mice. In Ayurvedic medicine, reserpine has been used as an antipsychotic and tranquilizer for more than a millennium.

Reserpine has a strong potential to be a candidate drug for AD, because: 1) first and foremost, it improves working memory (Figure 1 and Figure 2); 2) it decreases $A\beta$ 42-deposits/levels in brain and serum [16]; 3) it
increases NGF secretion and its action through TrkA signaling\cite{16}; 4) it induces expression of antiapoptotic protein, BCL-2 which can protect against neuronal loss \cite{16}; 5) it does not inhibit or drastically downregulate the gamma-secretase complex \cite{16} which is crucial for notch signaling, lack of which leads to cancer \cite{3} \cite{4}; 6) it can cross the blood brain barrier; 7) the major caveat about reserpine is the development of depression, and at a low dosage of 1 mg/kg reserpine does not cause depression in mice and the manic depression pathological marker mkb-1 is not induced \cite{19}, but at the dosage of 2 mg/kg both these effects are noticed \cite{19}; 8) it is widely in use, especially as an antihypertensive drug, even as a community based prophylactic at the low dosage of 0.05 mg against hypertension with minimal side effects for several years \cite{13} \cite{20}; 9) it reduces mortality when used as an antihypertensive \cite{17}.

Thus, Reserpine ameliorates Aβ induced cognitive deficit in the 5XFAD Tg mice at a low dosage of 0.08 mg orally administered on alternate days. Hence, reserpine’s efficacy can be evaluated in detail in the 3xTG AD model mice. In addition, reserpine analogs could be synthesized and evaluated for protection against AD. More importantly, given the limitations of animal models, exceptionally fruitful approach will be to carry out a pilot screen in AD patients given the severity, prevalence, social and financial burden on the society and the urgent need of a drug for AD.

5. Conclusion

Reserpine ameliorates Aβ induced cognitive deficit in the 5XFAD Tg mice at a low dosage of 0.08 mg orally administered on alternate days. This makes reserpine a potential drug to be evaluated in AD patients. As reserpine is originally identified to be protective in the Aβ toxicity \textit{C. elegans} model, the \textit{C. elegans} system is a worthwhile model for preclinical drug discovery.

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References

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