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Fluvoxamine maleate alleviates amyloid-beta load and neuroinflammation in 5XFAD mice to ameliorate Alzheimer disease pathology

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Introduction: Alzheimer pathology (AD) is characterized by the deposition of amyloid beta (A β) and chronic neuroinflammation, with the NLRP3 inflammasome playing a significant role. This study demonstrated that the OCD drug fluvoxamine maleate (FXN) can potently ameliorate AD pathology in 5XFAD mice by promoting autophagy-mediated clearance of A β and inhibiting the NLRP3 inflammasome.

Methods: We used mice primary astrocytes to establish the mechanism of action of FXN against NLRP3 inflammasome by using various techniques like ELISA, Western blotting, confocal microscopy, Immunofluorescence, etc. The anti-AD activity of FXN was validated in transgenic 5XFAD mice following two months of treatment. This was followed by behavior analysis, examination of inflammatory and autophagy proteins and immunohistochemistry analysis for A β load in the hippocampi.

Results: Our data showed that FXN, at a low concentration of 78 nM, induces autophagy to inhibit NF-κB and the NLRP3 inflammasome, apart from directly inhibiting NLRP3 inflammasome in primary astrocytes. FXN activated the PRKAA2 pathway through CAMKK2 signaling, leading to autophagy induction. It inhibited the ATP-mediated NLRP3 inflammasome activation by promoting the autophagic degradation of NF-κB, resulting in the downregulation of pro-IL-1β and NLRP3. The anti-NLRP3 inflammasome effect of FXN was reversed when autophagy was inhibited by either genetic knockdown of the PRKAA2 pathway or pharmacological inhibition with bafilomycin A1. Furthermore, FXN treatment led to improved AD pathology in 5XFAD mice, resulting in significant improvements in various behavioral parameters such as working memory and neuromuscular coordination, making their behavior more similar to that of wild-type animals. FXN improved behavior in 5XFAD mice by clearing the Aβ deposits from the hippocampi and significantly reducing multiple inflammatory proteins, including NF-κB, GFAP, IBA1, IL-1β, TNF-α, and IL-6, which are associated with NF-κB and

NLRP3 inflammasome in the brain. Moreover, these changes were accompanied by increased expression of autophagic proteins.

Discussion: Our data suggest that FXN ameliorates AD pathology, by simultaneously targeting two key pathological features: $A\beta$ deposits and neuroinflammation. As an already approved drug, FXN holds potential as a candidate for human studies against AD.

KEYWORDS

fluvoxamine maleate, Alzheimer disease, NLRP3 inflammasome, autophagy, amyloid beta, neuroinflammation, 5XFAD

Highlights

- Fluvoxamine maleate (FXN) potently inhibited NLRP3 inflammasome through induction of autophagy in primary astrocytes.
- Two months of treatment with FXN led to a significant reduction in amyloid beta load and inflammatory proteins in the hippocampi of 5XFAD mice.
- Mice treated with FXN displayed significantly improved memory behavior and neuromuscular coordination.

1 Introduction

Deposition of amyloid beta $(A\beta)$ and neuroinflammation are essential pathological features of Alzheimer disease (AD). AB contributes to neuroinflammation through multiple mechanisms. It activates NLRP3 inflammasome either directly by acting as a DAMP for sensor NLRP3 and indirectly by activating TLR4 mediated induction of NF-kB pathway (1-6). It can also activate NLRP3 inflammasome by inhibiting PRKAA2 and autophagy (7). Inflammation, in turn, also promotes the A β pathology through various pathways. NLRP3-mediated activation of IL-1ß inhibits the microglial cells from engulfing the A β plaques and thus contributes to its enhanced deposition (8-10). ASC specks secreted by immune cells directly bind to the A β and augment the accumulation of A β aggregates (11). NF-kB can also increase the activity of BACE1 to enhance the production of A β (12). Therefore, it's a kind of vicious cycle which helps in the progression of AD. Many studies have shown that autophagy is a common link between A β deposition and activation of NLRP3 inflammasome (13, 14). Autophagy, by virtue of its ability to recycle the cellular contents, can help clear protein aggregates and limit the activation of NLRP3 inflammasome by reducing the availability of proteins linked with inflammasome. Therefore, autophagy has emerged as one of the most important mechanisms that regulate NLRP3 inflammasome (15, 16). Some recent studies have shown that induction of autophagy by pharmacological or non-pharmacological means can help mitigate NLRP3 inflammasome mediated or other types of inflammation (17–20). Therefore, targeting NLRP3 inflammasome and autophagy together can be a good strategy, particularly when most of the monotargeted therapies have not shown any significant clinical advantage for treating AD.

In this study, as a part of our program related to the repurposing of drugs, we identified fluvoxamine (FXN) as a potent inhibitor of NLRP3 inflammasome and NF- κ B pathway. We show here that the anti-inflammatory properties of FXN are regulated by PRKAA2mediated autophagy. We attempted to utilize the ability of FXN to simultaneously target inflammation and autophagy in mitigating AD pathology in transgenic 5XFAD mice. We found that FXN can clear A β from the hippocampus of mice at a minimal dose 5 mg/kg along with a significant decline in neuroinflammation. These molecular changes in the brain helped improve working memory and overall brain health in 5XFAD mice. Our data projects FXN as a lead drug that should be explored for its possible development for the treatment of Alzheimer disease.

2 Materials and methods

2.1 Chemicals and antibodies

Chemicals- RPMI-1640 (Sigma-Aldrich-R6504), Dulbecco's modified eagle medium (Sigma-Aldrich-D1152), Penicillin (Sigma-Aldrich-P3032), Streptomycin (Sigma-Aldrich-S6501), Sodium bicarbonate (Sigma-Aldrich-S5761), Fetal Bovine Serum (FBS) (GIBCO-10270106), Phosphate buffer saline (PBS) (Sigma-Aldrich-D5652), EDTA (Invitrogen-15575038), Lipopolysaccharide (LPS) (Sigma-Aldrich-L3129), Adenosine 5'-triphosphate (ATP) (Sigma-Aldrich-A6419), Acrylamide (MP Biomedical-193982), Glycine (MP Biomedical-194825),Triton X-100 (Sigma-Aldrich-T8787), Albumin Bovine Fraction V (MP Biomedical-160069), Phenylmethylsulfonyl fluoride (PMSF) (MP Biomedical-195381), Skimmed milk (HimediaGRM1254), Strataclean resin (Agilent-400714-61), Bafilomycin A1 (Sigma-Aldrich-B1793), Rapamycin (Sigma-Aldrich-R8781),DAPI (Sigma-Aldrich-D9542), glycerol (Sigma-Aldrich-G5516), Tween 20 (Sigma-Aldrich-P7949), HEPES (Sigma-Aldrich-H3375), Paraformaldehyde (Sigma-Aldrich-P6148), MTT (Sigma-Aldrich-M5655), Trizma (Sigma-Aldrich-P6148), MTT (Sigma-Aldrich-L3771), Uric acid sodium salt (Sigma-Aldrich-U2875), Suberic acid (Sigma-Aldrich-S1885), OPTI-MEM media (Gibco-11058-021), β-Amyloid (Anaspec-AS-60479).

Antibodies- NLRP3 (CST-15101S), ASC (Santa Cruz Biotechnology-SC-22514), CASP1 (Santa Cruz Biotechnology-SC-56036), anti-mIL-1 β (R & D biotechnology-AF-401-NA), pCAMKK2 (CST-12818S), PRKAA2 (CST-2532S), pPRKAA2 (CST-2535S), pULK1 (CST-14202S), ULK1 (Santa Cruz Biotechnology-SC-33182), FIP200 (CST-12436S), BECN1 (Santa Cruz Biotechnology-SC-48341), ATG5 (Santa Cruz Biotechnology-SC-133158), ATG7 (Santa Cruz Biotechnology-SC-33211), ATG13 (CST- D4P1K), NF-KB (Santa Cruz Biotechnology-SC-372), pNF-κB (Santa Cruz Biotechnology-SC-136548), MTOR (CST-2972S), pMTOR (CST-5536S), LAMP1 (Santa Cruz Biotechnology-SC-20011), HRP-linked anti-goat IgG (Santa Cruz Biotechnology-SC-2354), siRNA PRKAA2 (CST-6620S), GFAP (CST-80788S), IBA1 (CST-17198S), HRP-linked anti-rabbit IgG (CST-7074S), HRP-linked anti-mouse IgG (CST-7076S), anti-mouse IgG Alexa flour 488 (CST-4408S), anti-mouse IgG Alexa flour 555 (CST-4409S), anti-rabbit IgG Alexa flour 488 (CST-4412S), anti-rabbit IgG Alexa flour 555 (CST-4413S), anti-ACTB (Sigma-Aldrich-A3854), ANTI-LC3B-II (Sigma-Aldrich-L7543), Anti-p62/SQSTM1 (Sigma-Aldrich-P0067).

Kits and reagents- PVDF Membrane (Millipore-ISEQ00010), Precision plus protein markers (Bio-Rad-161-0375), ECL-kit (Millipore-WBKLS0500), Bradford reagent (Bio-Rad-500-0006), FuGENE HD (Promega-E2313), mouse TNF- α ELISA kit (Invitrogen-88-7324-88), mouse IL-6 ELISA kit (Invitrogen-88-7064-88), mouse IL- 1 β ELISA kit (Invitrogen-88-7013-88), mouse IL- 18 ELISA kit (Invitrogen-88-50618-88), Human A β 42 ELISA kit (Invitrogen-KHB3441).

2.2 Synthesis of fluvoxamine maleate

The synthesis of fluvoxamine maleate was done by a wellestablished method (Figure 1A) (WO2014178064, US 6,433,225 B1). The scheme for the synthesis of fluvoxamine maleate (Scheme 1) and data related to its characterization are given in the Supplementary Material File.

2.3 Cell culture

2.3.1 Primary astrocyte culture

The brain was isolated from four to five days old pups of C57BL/6 mice and cortical hemisphere was treated with trypsin. Following centrifugation, the cells were passed through cell strainer and cultured in low glucose DMEM media and supplemented with 10% FBS. After every 2 days, fresh media was added to the cells.

After one week, the shaking was carried out using rocker to separate out the astrocytes from the top layer of the microglial cells. Thereafter, astrocytes were trypsinized and cultured for future experiments under 95% humidity and 5% $\rm CO_2$ at 37°C in an incubator chamber.

2.3.2 Cell line maintenance

The N9 microglial cells were cultured in RPMI-1640 media supplemented with 10% Fetal bovine serum (FBS), penicillin (70 mg/L), streptomycin (100 mg/L) and NaHCO₃ (3.7 g/L), under constant supply of 5% CO₂ and 95% humidity at 37°C.

2.4 NLRP3 inflammasome activation in primary astrocytes

In order to activate the NLRP3 inflammasome in primary astrocytes, N9 microglial cells were seeded in 24 well plates and allowed to grow for 24 h. The cells were then treated with LPS (1 μ g/ml) and incubated for next 24 h. The LPS activated microglial conditioning media was collected and primary astrocytes were pretreated with the conditioned medium for 24 h. Next day, primary astrocytes were primed with LPS (1 μ g/ml) for 4 h and treated with different concentrations of FXN and standard MCC 950 (100 nM) for 1 h in serum free media, followed by ATP (5 mM) stimulation for 30 min.

2.5 Enzyme-linked immunosorbent assay

After activation of NLRP3 inflammasome and drug treatments (as mentioned in the previous section), the supernatants were collected and analyzed to assess the levels of IL-1 β and IL-18 using ELISA following the manufacturer's instructions (Invitrogen). For detection of LPS induced pro inflammatory cytokines, TNF- α and IL-6, primary astrocytes were seeded in 24 well plate and pretreated with N9 conditioning media. Then the astrocytes were incubated with LPS for 24 h and the levels of proinflammatory cytokines were examined in the supernatants. The levels of cytokines were normalized by dividing the values with total protein content present in the sample. For in vivo studies, brain cortex was homogenized in RIPA buffer and centrifuged at 12000 rpm for 20 min at 4°C. The supernatants were collected and analyzed for IL-1 β , TNF- α and IL-6 cytokine levels. After completion of 2 months oral dosing, blood was collected from retro-orbital plexus and the $A\beta_{42}$ levels in plasma of 5X FAD mice were analyzed using ELISA following the manufacturer's guidelines (Invitrogen).

2.6 ASC oligomerization

After drug treatments under NLRP3 inflammasome activation condition, primary astrocytes were lysed in the cold buffer consisting of 1% cocktail, sodium orthovanadate (1 mM), KCL (150 mM), PMSF (0.1 mM), HEPES-KOH (20 mM) and 1% of NP-



Effect of FXN on the activation of the NLRP3 inflammasome in primary astrocytes. (A) Structure of Fluvoxamine maleate. (B, C) Graphs representing IC_{50} of FXN for IL-1 β and IL-18, respectively. The IC50 values were calculated in primary astrocytes after treatment with FXN in the serum free media for 1 h and ATP (5 mM) for 30 minutes. (D) Immunoblots depicting the expression of NLRP3 inflammasome complex proteins including IL-1 β , CASP1 and NLRP3. Densitometric analysis of the western blots is provided in the Supplementary Data (Supplementary Figure S1D). (E) Representative confocal images indicating the ASC speck formation in primary astrocytes, which generally form after NLRP3 inflammasome activation, and the graph showing average number of specks is provided in the Supplementary Data (Supplementary Figure S1C). (F) Western blots showing the ASC oligomerization. Densitometric analysis of immunoblots is given in Supplementary Data (Supplementary Figure S1F). (G) Representative images showing co-localization of NLRP3 (green) and CASP1 (red) in FXN (78 nM) treated primary astrocytes, which were stimulated with LPS and ATP. The mean fluorescence intensities of NLRP3 and CASP1 is provided in the Supplementary Data (Supplementary Figure S1G). The data provided are Mean±SD of three independent experiments and the statistical analysis was measured using one-way ANOVA, followed by *post-hoc* Bonferroni test.

40. The lysed cells were centrifuged at 330g for 10 min at 4°C and the supernatant was collected to analyze the protein expression by western blotting. The cell pellets were washed with PBS and 500 μ L of cold PBS was added to the cells. Suberic acid (2 mM) was added and incubated with the pellets for 30 min at 37°C for cross-linking of ASC proteins. Afterwards the pellets were centrifuged at 330g for 10 min at 4°C and 2X Laemmli buffer was added to the cell pellets. Cell lysates were heated at 95°C for 5 min and proteins were analyzed by western blotting.

2.7 Western blotting

For in vitro protein analysis by western blotting, primary astrocytes were seeded in 60 mm petri dishes and treated with FXN and standard MCC 950 under NLRP3 inflammasome activation condition. Cytokines secreted in the media were concentrated using Strataclean resin according to the instructions provided by manufacturer and cells were lysed with RIPA buffer [PMSF (2mM), Na3OV4 (0.5mM), NaF (50mM) and 1% cocktail] and centrifuged at 12,000 rpm for 20 min at 4°C. For in vivo studies, hippocampi from 5X FAD mice brain were homogenized and the lysed samples were centrifuged. Supernatants were collected and the protein content was estimated using Bradford's reagent. For in vitro studies 60 µg protein and for in vivo studies 20 µg protein was loaded for separation using SDS PAGE and proteins were transferred to PVDF membrane for 2 h at 4°C. The blots were blocked using 5% BSA or 5% skimmed milk for 1 h at RT and incubated overnight with primary antibody at 4°C. Afterward blots were incubated with HRP-conjugated secondary antibody for 2 h at RT. Chemiluminescent HRP substrate (Millipore) was used for detection of protein bands and visualized by Chemidoc system (Syngene G:BOX chemi XT 4).

2.8 Autophagy flux measurement

Autophagy flux was measured by treating primary astrocytes with different concentrations of FXN in the presence and absence of bafilomycin A1 for 24 h. The protein expression of LC3B-II and SQSTM1 were analyzed by western blotting.

2.9 Confocal microscopy

For confocal microscopic examination of the proteins, primary astrocytes were seeded in 6-well plate over coverslip. After different treatments, cells were washed thrice with PBS and fixed with 4% paraformaldehyde for 15 min. The cells were permeabilized with 0.2% Triton X-100 for 7 min and blocked using blocking buffer (2% BSA and 0.2% Triton X-100) for 1 h. The cells were incubated overnight with primary antibody at 4°C. Next day, the cells were incubated with secondary antibody (Alexa flour 555 or Alexa flour 488) for 1 h at RT and washed thrice with PBS. After counterstaining the nuclei with DAPI, the coverslips were mounted using the mounting media and the images were acquired using Yokogawa CQ1 Benchtop High-Content Analysis System at 40x or 60x.

2.10 A β clearance assay

 $A\beta_{42}$ -HiLyte flour488 peptide was prepared following the manufacturer's protocol (Anaspec Inc.). Primary astrocytes seeded on coverslips were treated with FXN or rapamycin. After 12 h, 2 µg/ml fluorochrome tagged (Hilyte flour488) $A\beta_{42}$ protein was added to cells for another 12 h. Bafilomycin A1 was given 3 h prior to experiment termination. Cells were washed with PBS and fixed with 4% PFA, afterwards cells were permeabilized with Triton-X 100, and the nuclei were counterstained with DAPI. Slides were prepared and images were taken and analyzed in the Yokogawa CQ1 Benchtop High-Content Analysis System at 40x or 60x.

2.11 Transfection of primary astrocytes with PRKAA2 siRNA

Primary astrocytes were grown in 6-well plate for confocal microscopic analysis and 60-mm dishes for western blotting. For transfection of PRKAA2, cells were incubated with OPTI-MEM media and PRKAA2 siRNA in FuGENE HD was added for 24 hours. Transfected primary astrocytes were incubated with microglial conditioning media for 24 h, followed by FXN (78 nM) treatment under NLRP3 inflammasome activation condition and the samples were analyzed by western blotting and confocal microscopy at 60x (Yokogawa CQ1 Benchtop High-Content Analysis System).

2.12 Drug formulation

For *in vivo* studies, FXN was formulated in 5% of DMSO, 30% PEG400, 20% PEG200 and 45% distilled water. LPS and ATP were dissolved in PBS.

2.13 Animals and ethical clearance

Six months old 5XFAD transgenic mice were used in the study and randomly divided into three groups (n = 5). C57BL6/J mice were used as Control wild type (WT). They were housed under a 12hour light/dark cycle in a temperature (65–75°F; ~18-23°C) and humidity-controlled (40–60%) environment, supplied with free access to the food and water (ad libitum). Prior to initiation of study all the animals were acclimatized for one week under standard laboratory conditions, animals were drug naive with no prior procedures performed. All testing were performed from 1 to 4 p.m. Mice were randomized in groups based on their body weights for all behavioral assays and testing were performed by an experimenter blinded to the treatment groups. Total study duration was 2 months. All experiment protocols were approved by the Institutional Animal Ethics Committee (IAEC) (IAEC

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approval no.- 321/82/2/2023), and followed the Committee for Control and Supervision of Experiments on Animals (CCSEA; Ministry of Environment and Forest, Government of India) guidelines for animal care.

2.14 Open-field test

The open field arena box (60 cm x 45 cm x 25 cm) made up of white colored non-reactive plastic was used for all the assessments. The mice were acclimatized to the testing room 30 minutes prior to experiment. Individual mice were placed in the center of the open field arena and allowed to freely explore it for 5 minutes. The arena box was cleaned with a 70% ethanol after every trial. The locomotor activity and exploratory behavior of mice were recorded using a video camera connected to AnyMaze software. Mice were assessed in the open field and the following parameters were recorded automatically: Total distance travelled (cm), average speed (m/s), time spent in the center area (sec), and time spent in the corners (sec).

2.15 Radial arm maze test

The eight-armed radial arm maze (UGO Basile) was used to assess spatial memory behavior. The mice were habituated to the maze for three days before the experiment. Each mouse was trained to reach to the baited arm having a butter cookie as food reward and flag as a visual cue. Mice were trained for five days to locate the baited arm. Memory retention trials include recording of several parameters for three consecutive days. The movement of mice was tracked with the help of automated AnyMaze software connected with a video tracking camera.

2.16 Neuromuscular coordination test

The neuromuscular coordination of mice was recorded by rotarod. For this assessment mice were trained for three consecutive days with three trials per day. The maximum time allowed on the rotarod during the experiment was 300 seconds. The rotarod was initially set at a speed of 4 rpm at the start of the test and accelerated to 40 rpm over 300 seconds. The latency to fall from the rotarod was recorded for each mouse, and the mean latency to fall for each group was calculated.

2.17 Immunohistochemical analysis of 5XFAD mice brains

After the treatment completion, the hippocampi were isolated from the mice brains and the 10 μ m thick tissue-sections were prepared. For IHC, the slides were deparaffinized and histochemical staining was carried out. The sections were incubated with primary antibody (overnight at 4°C) under humid conditions after blocking using 2% BSA solution in TBS. For analysis of total A β , sections were incubated with anti-A β antibody (1-42) D3E10 followed by secondary antibody (Alexa fluor 488) and for localizing reactive astrocytes, anti-GFAP was used followed by secondary antibody (Alexa fluor 555). The alteration in autophagy levels were assessed by examination of LC3B levels in the tissue sections of mice hippocampi. The sections were immuno-stained with LC3B followed by secondary antibody (Alexa fluor 488). The nuclei were counterstained using DAPI. The images were acquired and quantified using Confocal Quantitative Image Cytometer at 40X (Yokogawa CQ1).

2.18 MTT assay

To assess cytotoxicity, primary astrocytes were cultured in 96well plates and exposed to varying concentrations of FXN for 24 and 48 hours. Four hours before termination, MTT dye was added to each well. Termination involved the addition of DMSO to dissolve the formazan crystals, followed by measuring the absorbance at 570 nm.

2.19 Statistical analysis

Statistical analyses were analyzed using GraphPad Prism 9 software. The data shown here are Mean±SD of three independent experiments. The response of independent mice from the group was noted for animal experiments and the mean was calculated. The statistical analysis of data was calculated using one-way ANOVA, followed by *post-hoc* Bonferroni test. The p-value<0.05 was considered to be statistically significant with values assigned as ****p < 0.0001, ***p < 0.001, **p< 0.01, *p< 0.05 and ns= not significant.

3 Results

3.1 FXN potently inhibited the release of IL- 1β , indicating the inhibition of NLRP3 inflammasome activity in primary astrocytes

NLRP3 inflammasome is an important player in neuroinflammation, so we treated the LPS-primed primary astrocytes with FXN before induction with ATP to analyze the anti-NLRP3 inflammasome activity of FXN. The ELISA results showed that FXN could inhibit NLRP3 inflammasome even at very low concentrations of nine nanomolar, which was evidenced by the measurement of released IL-1 β and IL-18 as inflammasome activation products. FXN inhibited the release of IL-1 β and IL-18 with IC50 values of 16 nM & 25.96 nM, respectively (Figures 1B, C). We also analyzed the inhibitory effect of FXN on NLRP3 inflammasome by using different types of second stimulus in three different types of cells. In nigericin-induced microglial N9 cells, FXN inhibited the release of IL-1 β with an IC50 value of 7.02 nM, in ATP induced macrophage J774A.1 cells, it showed IC50 value of 18.40 nM, and in hemin induced THP-1 cells, FXN

displayed the IC50 value 17.24 nM (Supplementary Figures S1A-C). Based on the concentration-dependent inhibitory activity of FXN; we chose 78 nM concentration for further detailed studies in astrocytes. We also confirmed the inhibition of release of cleaved IL-1β and CASP1 through western blotting of the proteins collected from the supernatant of primary astrocytes treated with FXN (Figure 1D). The analysis of whole cell lysates revealed reduced levels of pro-IL-1ß and NLRP3, while pro-CASP1 was unchanged (Figure 1D; Supplementary Figure S1D). Further, the oligomerization of ASC is an important step in the activation of NLRP3 inflammasome. Therefore, we analyzed the effect of FXN on ASC by using two methods, including confocal microscopy and western blotting. Both experiments showed similar results with the potent inhibitory effect of FXN on ASC oligomerization (Figures 1E, F; Supplementary Figures S1E, F). We also analyzed the colocalization of CASP1 and NLRP3 in the presence of FXN. Data revealed that FXN and MCC950 (used as a standard) had a similar inhibitory effect on the co-localization of CASP1 and NLRP3 (Figure 1G; Supplementary Figure S1G).

3.2 FXN inhibited the NLRP3 inflammasome through NF-κB pathway

In the initial experiments, we treated the LPS-primed astrocytes with FXN to know whether it could inhibit the assembly of the NLRP3 inflammasome complex. We also wanted to know if it can inhibit the NLRP3 inflammasome at the transcriptional level by inhibiting the NF- κ B. Therefore, we treated the cells with FXN before priming them with LPS. We found that FXN could significantly inhibit NLRP3 inflammasome at a concentration of 78 nM contrary to nine nanomolar, observed during FXN treatment post-priming. The ELISA measurement of IL-1ß showed an IC50 value of 131.1 nM (Figure 2A). We confirmed the inhibitory effect of FXN on NF- κ B by direct and indirect methods. The dwindling levels of NF- κB dependent proinflammatory cytokines TNF- α and IL-6 after treatment with FXN at 78 nM indicated the inhibition of NF- κ B in astrocytes (Figures 2B, C). Direct evidence for inhibition of NF-kB pathway came from western blot analysis of NF-kB in the cytosolic and nuclear fraction of astrocytes treated with FXN. The data showed that the movement of NF-KB, p65 from the cytoplasm to the nucleus is hindered in the cells treated with FXN in the presence of LPS and ATP stimuli, wherein the levels of NF-KB in the cytoplasm were significantly higher than in the untreated control. On the contrary, its level in the nucleus was significantly reduced (Figures 2D-F).

3.3 FXN induced the autophagy under the inflammatory conditions in primary astrocytes

After confirming the inhibitory effect of FXN on NLRP3 inflammasome, we wanted to know its inhibition mechanism. Therefore, we focused on autophagy as a major mechanism regulating inflammasome activity. We found that FXN could induce autophagy at a low concentration of 39 nM; however, its effect was enhanced with the increasing concentration (Figure 3A; Supplementary Figure S2A). Further, to confirm the completion of autophagy, we calculated the autophagy flux by using LC3B-II and SQSTM1 expression after treatment of astrocytes with FXN (78 nM) in the presence and absence of end-stage autophagy inhibitory bafilomycin A1. We found an autophagy flux at all the tested time points through 24 h of astrocyte treatment with FXN (Figure 3B; Supplementary Figure S2B). Induction of autophagy by FXN was also confirmed by confocal microscopy analysis of co-localization of LC3B-II and LAMP1 in primary astrocytes. The cells treated with FXN and rapamycin showed a significant amount of co-localization of these proteins, while in the presence of bafilomycin A1, FXNtreated cells did not show any observable co-localization of LC3B-II and LAMP1 (Figures 3C, D). After confirming the induction of autophagy by FXN, we checked if it can induce autophagy in primary astrocytes under inflammatory conditions where the cells have been challenged with LPS and ATP. We found that the treatment of astrocytes with FXN before ATP induced the expression of LC3B-II, while SQSTM1 was downregulated, thus confirming the induction of autophagy (Figure 3E; Supplementary Figure S2C). We further analyzed the expression of various proteins involved in the induction of autophagy. We found that FXN induced autophagy through CAMKK2 mediated induction of PRKAA2 pathway leading to activation of various proteins including pPRKAA2 (Thr 172), pULK (Ser 317), ATG13, FIP 200, BECN1, ATG5 and ATG7 and downregulation of pMTOR (Figure 3F; Supplementary Figures S2C). However, induction of autophagy can also play a detrimental role in cell survival. Therefore, to rule out any toxicity due to induction of autophagy by FXN, we treated the primary astrocytes with various concentrations of FXN for up to 48 h and checked their viability by MTT assay. However, we did not find any significant toxicity of FXN even at 320 times higher concentration (25 µM) of FXN (Supplementary Figure S2D).

3.4 FXN inhibited NLRP3 inflammasome by inducing autophagy

For the confirmation of the involvement of autophagy in FXNmediated inhibition of NLRP3 inflammasome, we inhibited the autophagy through genetic and pharmacological methods. We pretreated the primary astrocytes with siPRKAA2 and analyzed the expression of autophagic and NLRP3 inflammasome-related proteins. The inhibition of PRKAA2 led to the reversal of the autophagic effect of FXN, which was reflected in the form of a marked decrease in the expression of LC3B-II and BECN1, while SQSTM1 expression was significantly increased (Figure 4A; Supplementary Figure S3A). Similarly, the inhibition of NLRP3 inflammasome by FXN was also reversed in the cells pretreated with siPRKAA2. The cleaved IL-1 β levels were significantly increased, whereas the reduction in expression of NLRP3 was also reversed in the cells treated with FXN in the presence of *siPRKAA2* (Figure 4A; Supplementary Figure S3A). We also confirmed these findings by using confocal microscopy, where the co-localization of CASP1 and



Effect of FXN on the NF-κB signaling pathway. The effect of FXN on NF-κB pathway was analyzed by treating the primary astrocytes with FXN, prior to LPS priming and ATP activation of NLRP3 inflammasome. (A) Graph representing IC50 value of FXN for IL-1β. Primary astrocytes treated with FXN (78 nM) and the standard dexamethasone in the presence of LPS for 24 h release NF-κB mediated pro-inflammatory cytokines TNF-α and IL-6 (B, C) Graphs representing the TNF-α and IL-6 levels. (D) The impact of FXN on nuclear translocation of pNF-κB (p65) in the presence of LPS and ATP was analyzed by western blotting. LMNA was used as a nuclear marker and ACTB was used as cytosolic marker. (E, F) showing densitometric analysis of nuclear and cytosolic fractions of pNF-KB (p65), respectively. The data shown here are Mean±SD of three independent experiments and the statistical analysis was calculated using one-way ANOVA, followed by post-hoc Bonferroni test. The p-value<0.05 was considered to be statistically significant with values assigned as ****p < 0.0001, ***p < 0.001, **p < 0.001, **p < 0.01.



Impact of FXN treatment on autophagic processes in primary astrocytes under inflammatory conditions. (A) Immunoblots showing concentration dependent effect of FXN and rapamycin on the expression of LC3B-II and SQSTM1 after 24 h treatment in primary astrocytes. Densitometric analysis is given in Supplementary Data (Supplementary Figure S2A). (B) Immunoblots showing the time dependent effect of FXN (78 nM) on autophagic flux in the presence and absence of bafilomycin A1 in primary astrocytes. Densitometric analysis is provided in Supplementary (Supplementary Figure S2B). (C) Representative images showing the effect of FXN (78 nM) on co-localization of LAMP1 (green) and LC3B (red) in primary astrocytes. (D) Graph representing the mean fluorescence intensities of LAMP1 and LC3B. (E) Immunoblots showing the effect of FXN (78 nM) on LC3B-II and SQSTM1 under NLRP3 inflammasome activation conditions in primary astrocytes. Densitometry of immunoblots is provided in Supplementary Data (Supplementary Figure S2C). (F) Immunoblots showing the effect of FXN on pMTOR (Ser 2448) and other autophagic proteins involved in the initiation, elongation, maturation and fusion of autophagosome [pCAMKK2 (Ser 511), pPRKAA2 (Thr 172), pULK1 (Ser 317), ATG 13, FIP 200, BECN1, ATG5 and ATG7] in primary astrocytes. The data provided are Mean±SD of three independent experiments and the statistical analysis was performed using one-way ANOVA, followed by *post-hoc* Bonferroni test. The p-value<0.05 was considered to be statistically significant with values assigned as ****p < 0.0001, ***p < 0.001, ***p < 0.01 and ns, not significant.

NLRP3 was restored to the level present in cells primed with LPS and ATP when they were treated with FXN along with knockdown of PRKAA2 by using siRNA (Figure 4B; Supplementary Figure S3B). The knockdown of PRKAA2 was confirmed by using western blotting, where the expression of PRKAA2 and pPRKAA2 was found to be significantly reduced (Figure 4C). We found that a similar kind of reversal of the inhibitory activity of FXN against NLRP3 inflammasome could be achieved if the autophagy is inhibited by using the end-stage pharmacological inhibitor bafilomycin A1. The cleavage of IL-1 β was almost completely inhibited, while the expression of NLRP3, NF- κ B (p65) along with autophagy markers LC3B-II, SQSTM1 was restored when the cells were treated with FXN in the presence of bafilomycin A1 (Figure 4D; Supplementary Figure S3C).

3.5 FXN cleared $A\beta_{42}$ in primary astrocytes by inducing autophagy

To confirm whether the autophagy induced by FXN can help in the clearance of $A\beta_{42}$, we treated the cells for a total of 24 h with FXN, including 12 h treatment with hilyte fluor 488 tagged $A\beta_{42}$. The confocal analysis revealed that the cells treated with FXN and rapamycin were almost completely devoid of green fluorescence indicating clearance of $A\beta_{42}$ in comparison to control cells. Further, 3 h treatment of cells with autophagy inhibitor bafilomycin A1 prior to termination of the experiment completely repressed the clearance of $A\beta_{42}$, as indicated by the same level of green fluorescence as that of control cells (Figures 5A, C). The involvement of autophagy in the clearance of $A\beta_{42}$ by FXN was further confirmed when we inhibited the autophagy by using *siRNA* against PRKAA2. The clearance of $A\beta_{42}$ by FXN in the presence of *siPRKAA2* was similarly stopped as we observed with bafilomycin. A1 (Figures 5B, D).

3.6 FXN improved working memory, exploratory behavior and neuromuscular coordination in 5XFAD mice

After confirming the mechanism of inhibition of NLRP3 inflammasome by FXN, we wanted to know if its dual pharmacological activity can improve Alzheimer disease pathology. Therefore, we treated the 6-month-old 5XFAD mice for two months and analyzed multiple behavior parameters. In the radial arm maze test, the untreated 5XFAD mice displayed significant deterioration of neuronal functions in comparison to wild-type control (Figure 6A). However, the groups treated with FXN (5 and 10 mg/kg) took significantly less time and traveled a reduced distance in finding the baited arm and spent more time in the baited arm apart from fewer entries into non-baited arms in comparison to untreated 5XFAD mice, indicating improvement in the spatial memory (Figures 6B-E). In the open field test, untreated 5XFAD mice spent significantly more time in the center than in corners and showed little interest in exploring the surroundings (Figure 6F). However, the mice treated with FXN covered a significantly higher distance at greater speed and showed normal behavior by spending more time in the corners than in the center (Figures 6G–K). We also observed a significant improvement in the neuromuscular coordination in the mice treated with FXN, which showed delayed latency to fall than the untreated 5XFAD mice (Figure 6L). In all the behavior parameters studied, the mice treated with FXN behaved more like wild-type animals than the untreated 5XFAD control group.

3.7 FXN reduced the A β_{42} in the hippocampus and plasma of 5XFAD mice

Improvement in neuronal health of 5XFAD mice after treatment with FXN led us to investigate the underlying cause for such change. Therefore, through immunohistochemistry (IHC), we checked the levels of $A\beta_{42}$ in the hippocampus after two months of treatment with FXN. We found a significant decline in the levels of $A\beta_{42}$ in the hippocampus of both groups of mice treated at 5 and 10 mg/kg, respectively compared to the control group, as indicated by reduced green fluorescence (Figure 7A). The average number of $A\beta_{42}$ plaques, skeletal length of $A\beta_{42}$ plaques, branch count and circumference of $A\beta_{42}$ plaques were significantly reduced in FXN treated groups as compared to 5XFAD control group (Figures 7B-E). A β_{42} is strongly related to activation of astrocytes, we therefore checked if reduced levels of $A\beta_{42}$ had any impact on the reactive phenotype of astrocytes. The analysis of GFAP in the vicinity of $A\beta_{42}$ revealed that as the levels of $A\beta_{42}$ were depleted after treatment with FXN, the expression of GFAP reduced proportionately, as indicated by reduced red color fluorescence (Figures 7A, F). Further, the reduction of $A\beta_{42}$ levels in the brain also led to reduced circulatory levels of $A\beta_{42}$ in the blood plasma of 5XFAD mice treated with FXN in comparison to untreated control mice, as observed through ELISA (Figure 7G).

3.8 FXN ameliorated the AD pathology in 5XFAD mice through autophagy-mediated reduction in amyloid beta levels and neuroinflammation

Based on the effect of FXN on *in vitro* and *in vivo* clearance of $A\beta_{42}$, we hypothesized that autophagy induced by FXN is responsible for the reduced level of $A\beta_{42}$ in the hippocampi of 5XFAD mice brains. Therefore, to test the hypothesis, we analyzed the expression of LC3B-II in the hippocampi and found a highly significant increase in its levels in the mice treated with FXN (Figure 8A). Further, analysis of expression of various autophagic proteins including pCAMKK2 (Ser 511), pPRKAA2 (Thr 172), pMTOR (Ser 2448), LC3-IIB, pULK1 (Ser 317), BECN1, and ATG5 revealed a significant upregulation of these proteins in the hippocampi of mice treated with FXN in comparison to untreated mice, while the effect of FXN at 10 mg/kg was more pronounced as compared to the dose of 5 mg/kg (Figure 8B; Supplementary Figures S4A).

The analysis of pro-inflammatory proteins in the hippocampi also supported the anti-inflammatory effect of FXN. We found that



Role of autophagy in FXN-mediated inhibition of the NLRP3 inflammasome. Autophagy was genetically and pharmacologically inhibited by *siPRKAA2* and bafilomycin. A1 respectively in primary astrocytes under inflammasome activation conditions. **(A)** Western blots indicating the effect of PRKAA2 knockdown on the expression levels of pPRKAA2 (Thr 172), BECN1, LC3B-II, SQSTM1, NF- κ B (p65), IL-1 β and NLRP3. Densitometric analysis is given in **Supplementary Data** (Supplementary Figure S3A). **(B)** Representative images indicating the colocalization of NLRP3 and CASP1 under aforementioned conditions (genetic knockdown of PRKAA2 by *siPRKAA2*) after FXN treatment. The mean fluorescence intensities of NLRP3 and CASP1 are provided in the **Supplementary Data** (Supplementary Data (Supplementary Data (Supplementary Data) (C) Immunoblots depicting the PRKAA2 expression in primary astrocytes following treatment with the *siPRKAA2*, in comparison to mock *siRNA* and their densitometric analysis. **(D)** Immunoblots showing the effect of FXN on expression levels of IL-1 β , NLRP3, NF- κ B (p65), LC3B-II and SQSTM1 after pharmacological inhibition of autophagy using bafilomycin A1. Densitometric analysis is given in **Supplementary Data** (Supplementary Figure S3C). The data represent Mean±SD of three independent experiments and the statistical analysis of data was performed using one-way ANOVA, followed by *post-hoc* Bonferroni test. The p-value<0.05 was considered to be statistically significant with values assigned as ****p < 0.0001.



Impact of FXN-induced autophagy on $A\beta_{42}$ clearance in primary astrocytes. (A) Representative images showing the effect of FXN, in the presence Baf. A1, on the intracellular deposition of A β in primary astrocytes treated with fluorescently tagged (HiLyte 488) A β peptide. Rapamycin was used as a standard autophagy inducer. (B) Representative images showing the effect of FXN on intracellular deposition of A β in primary astrocytes transfected with *siPRKAA2*. (C) Graph representing the mean fluorescence intensity of images shown in panel (A). (D) Graphical representation of mean fluorescence intensity of images shown in panel (B). The data shown here are Mean±SD of three independent experiments and the statistical analysis of data was calculated using one-way ANOVA, followed by *post-hoc* Bonferroni test. The p-value<0.05 was considered to be statistically significant with values assigned as ****p < 0.0001, ***p < 0.001, **p< 0.01 and ns, not significant.



Analysis of behavioral parameters in 5XFAD mice following FXN treatment. The effect of FXN (5 and 10 mg/kg) on cognitive behavior was assessed after two months of oral dosing in Alzheimer (5XFAD) mice model (n=5 per group). Vehicle 5XFAD control mice were compared with wild-type C57BL/6J mice of same age. Radial Arm Maze Test: (A) Representative images showing track plots of radial arm maze post five days training. The data values represents average of three independent memory retention trials. The different parameters were assessed such as (B) Latency to first entry to baited arm, (C) Time spent in baited arm, (D) Entries to non-baited arms and (E) Distance travelled by mice. (F) Images depicting the track plots obtained from open field test. Various parameters were investigated such as (G) Distance travelled by mice, (H) Mobility time, (I) Time spent in the Centre zone, (J) Corner zone time and (K) Maximum speed. The latency to fall time of mice from rotarod was assessed by using rotarod test. (L) Graph representing the latency to fall time. Automated camera was used to capture the movement of mice in radial arm maze and open field test and all the activities were measured by using AnyMaze software. The statistical analysis of data was measured using one-way ANOVA, followed by *post-hoc* Bonferroni test. The p-value <0.05 was considered to be statistically significant with values assigned as **** p < 0.0001, *** p < 0.001, **p < 0.05.



Effect of FXN treatment on the $A\beta_{42}$ deposition in the hippocampus and plasma of 5XFAD mice. Immunostaining post FXN treatment (5 and 10 mg/ kg for 2 months), the formalin-fixed paraffin embedded (FFPE) tissues of brain hippocampi were immuno-stained with D3E10 for detection of total A β (green), GFAP for activated astrocytes (red) and the nuclei were counterstained with DAPI (blue). (A) Representative images showing co-localized A β plaques and reactive astrocytes in hippocampi of FXN treated mice (n=5 per group). Graphs showing (B) Average number of A β plaques (C) Average skeletal length of A β plaques (D) Average branch count of A β plaques (E) Average circumference of A β plaques (F) Fold change in the GFAP fluorescence (G) The plasma levels of A β -42 in mice measured using ELISA. Confocal microscopic images were examined using cell pathfinder software. The statistical analysis was performed using one-way ANOVA, followed by *post-hoc* Bonferroni test. The p-value <0.05 was considered to be statistically significant with values assigned as ****p < 0.0001, ***p < 0.001.

treatment of mice with FXN significantly reduced the levels of both NF-κB, p50 and p65 subunits confirming its effect on the most important inflammatory pathway. Further analysis of NLRP3 inflammasome proteins showed significant down-regulation of NLRP3, which significantly reduced ASC oligomerization and cleaved CASP1 and IL-1β in the mice treated with FXN (Figures 8C, D; Supplementary Figures S4B). The reduction of inflammation in the hippocampi was indicated by reduced levels of microglial marker protein IBA1 (Figure 8C; Supplementary Figures S4B). We also observed a significant decline in the levels of pro-inflammatory cytokines TNF- α , IL-6 and IL-1β, which were measured in the cortex region of the brain by using ELISA (Figures 8E–G). These data emphasized a potent effect of FXN on the pathology of AD in 5XFAD mice.

4 Discussion

Alzheimer disease progresses with a complex pathology. However, most cases of AD are accompanied by the presence of $A\beta_{42}$ plaques and neuroinflammation where NLRP3 inflammasome plays a significant role (2, 21-23). There are multiple types of damage associated molecular patterns (DAMPs) that can upregulate NLRP3 inflammasome in the brain, including $A\beta$, TAU NFTs, and ATP (24-27). All these DAMPs contribute to chronic inflammation and AD pathology. We initiated this study focusing on finding NLRP3 inflammasome inhibitors that can be used to mitigate AD pathology. Our preliminary studies found that fluvoxamine maleate (FXN) is a potent inhibitor of NLRP3 inflammasome with an inhibitory concentration in the nanomolar range in various types of cells that were induced by using diverse type of second stimulus including ATP, nigericin, and hemin. US FDA had approved FXN for the treatment of obsessive-compulsive disorder (OCD) in 1994. Its proposed mechanism against OCD is through the inhibition of serotonin reuptake transporter (5-HT) (28-30). However, our studies revealed new targets of FXN that can be effectively exploited against AD. Additionally, being an approved drug for CNS disease, FXN gave us an additional advantage to be used for AD studies without being worried about its safety and crossing of the blood-brain barrier (BBB). For detailed studies on the effect of FXN on neuroinflammation, we continued working on primary astrocytes because of their emerging role in AD and other neurodegenerative diseases. We found that FXN downregulated the expression of NLRP3 and pro-IL1 β along with reduced oligomerization of ASC leading to poor assembly of NLRP3 inflammasome and thus cleavage of CASP1. These data indicated that FXN works on both the steps of NLRP3 inflammasome activation to inhibit its assembly. This assumption was confirmed when we observed a significant reduction in the nuclear translocation of NF-KB, p65 and inhibition of pro-inflammatory cytokines TNF- α and IL-6, which depend on it. These data emphasized that FXN has a wider anti-inflammatory spectrum that targets multiple inflammation-related pathways. These findings are important as inflammation is the key driver in neuronal cell death during AD (31, 32).

After confirming the anti-inflammatory activity of FXN, we intended to know how the upstream mechanisms of two important molecular pathways were inhibited by FXN. Therefore, we explored the possible involvement of autophagy in the FXN mediated downregulation of inflammation. Autophagy can regulate inflammation through multiple ways including recycling of proteins, and organelles like mitochondria, clearance of protein aggregates and other DAMPs, invading microorganisms and PAMPs associated with them, etc. (33-35). We found that FXN is a very potent inducer of autophagy, which works in the same concentration range, which was effective against inflammation. We further observed that it can also induce autophagy under the inflammatory conditions generated by using LPS and ATP. The treatment of astrocyte with FXN under inflammatory conditions showed a significant decline in the expression of NF-κB and other proteins associated with activation of NLRP3 inflammasome. This was an interesting observation. However, we wanted to be sure about the involvement of autophagy in the anti-inflammatory effects of FXN. Therefore, we inhibited the autophagy by both pharmacological and genetic means and found that all the antiinflammatory effects of FXN were reversed, which was evidenced by re-instatement of expression level of inflammatory proteins to the pre FXN treatment levels.

Apart from being an important regulator of inflammation, the key physiological role of autophagy is to remove protein aggregates (35). The failure of autophagy to clear proteins aggregates is strongly linked to number of pathological conditions including AD, where it helps to maintain the physiological levels of A β along with other clearance mechanisms (36). We found that FXN could also help in the clearance of A β by inducing autophagy in primary astrocytes. This was further confirmed when the clearance of A β was completely reversed in the presence of *siPRKAA2*.

These findings encouraged us to validate the efficacy of FXN to ameliorate the AD pathology in transgenic 5XFAD mice. Therefore, we treated the 5XFAD mice for two months with FXN at two different doses. At the end of treatment, the analysis of different behavior parameters revealed that the mice treated with both doses of FXN had significantly improved brain health. The data from different parameters studied during the radial arm maze test showed a marked improvement in the working memory of mice. The hippocampus is the most important part of the brain that processes working memory and is severely affected during AD, as was observed in untreated control mice. Further, the exploratory behavior of FXN-treated mice was also significantly improved and it appeared more like non-demented animals. They spent more time in corners, traveled more distances at higher speeds, etc. The neuromuscular coordination was also found to be similarly improved in mice treated with FXN. All these parameters clearly indicated the improvement of brain health after treatment with FXN. These data made us to explore molecular mechanisms behind this improvement in brain health, particularly memory behavior. Therefore, we isolated the hippocampi of these mice and analyzed the expression of various proteins related to autophagy and inflammation. We found that the mice treated with FXN had a significantly increased expression of proteins involved in autophagy



(p50), NF- κ B (p65), NLRP3, ASC oligomer, ASC dimer and ASC monomer, CASP1, IL-1 β and IBA1. (**D**) Densitometric quantification of ASC oligomer, dimer and monomer forms. The densitometric analysis of other western blots is provided in the Supplementary Data (Supplementary Figure S4B). Graphs depicting the levels of pro-inflammatory cytokines in brain cortex of FXN treated 5XFAD mice (**E**) TNF- α , (**F**) IL-6 and (**G**) IL-1 β . The samples were compared by using one-way ANOVA, followed by *post-hoc* Bonferroni test. The p-value <0.05 was considered to be statistically significant with values assigned as ****p < 0.0001, ***p < 0.001, **p < 0.05.

induction including pCAMKK2 (Ser 511) and pPRKAA2 (Thr 172), and downregulation of pMTOR (Ser 2448). On the similar lines the expression of autophagy initiation and execution proteins pULK1 (Ser 317), BECN1, LC3B-II, and ATG5 was found to be significantly increased. The most important impact of autophagy induction by FXN in the hippocampus was the clearance of A β_{42} . We found only traces of A β_{42} in the hippocampus of FXN treated mice in comparison to untreated control. Interestingly, reduced A β_{42}

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levels were directly related to low-reactive phenotype of astrocytes in the hippocampi as evidenced by reduced expression of GFAP. Furthermore, the expression of microglial marker IBA1 was also found to be significantly down regulated in mice treated with FXN. These data indicate that FXN through autophagy not only cleared $A\beta_{42}$ but it also helped mitigation of overall inflammatory microenvironment in the brain. These data were further supported by down regulation of NF- κ B and other proteins associated with the activation of NLRP3 inflammasome in the hippocampus. Moreover, these effects of the FXN were not localized to hippocampi as the analysis of proinflammatory cytokines TNF- α , IL-6, and IL-1 β in the cortex region of the brain clearly established the strong effect of FXN against neuroinflammation. These data directly correspond to the improvement of memory behavior.

5 Conclusion

The analysis of whole data indicates that FXN exerts a potent autophagic effect through activation of PRKAA2 pathway, which helps in the clearance of $A\beta_{42}$ and inhibition of neuroinflammation. In AD, deposition of $A\beta_{42}$ and neuroinflammation together inflict severe damage to neuronal tissue in the brain. Interestingly, FXN exerted these anti-Alzheimer effects at the minimal dose of 5 mg/kg, which corresponds to approximately 25 mg/kg of human equivalent dose, which is a minimum prescribed dose of FXN for OCD. Therefore, FXN through its ability to work against both these key pathological hallmarks of AD at a minimal dose can raise a new hope and warrants its further development against AD.

Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

Ethics statement

The animal study was approved by Institutional Animals Ethics Committee of CSIR-IIIM, Jammu. The study was conducted in accordance with the local legislation and institutional requirements.

Author contributions

SK: Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Software, Validation, Visualization, Writing – original draft. KS: Data curation, Investigation, Methodology, Writing – original draft. AS: Conceptualization, Investigation, Methodology, Validation, Visualization, Writing – original draft. KKS: Data curation, Formal analysis, Investigation, Methodology, Visualization, Writing – original draft. SA: Investigation, Software, Validation, Visualization, Writing – original draft. RA: Formal analysis, Investigation, Methodology, Visualization, Writing – original draft. PR: Formal analysis, Investigation, Methodology, Resources, Validation, Visualization, Writing – original draft. PS: Data curation, Formal analysis, Investigation, Methodology, Validation, Visualization, Writing – original draft. ZA: Visualization, Funding acquisition, Supervision, Writing – original draft, Resources, Validation. AK: Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation, Methodology, Resources, Supervision, Validation, Visualization, Writing – review & editing.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2024. 1418422/full#supplementary-material

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Glossary

ADAlzheirer's diseaeFRAM2Fordin kinase AMP-activated catalytic subunit apha 2ATGAndyag relatedAND4Andysis of varianceATGAlonsine triphosphateATGAlonsine triphosphateATGAlorisinasciatical specific line triphosphateATGAlorisinatical specific line triphosphateATGAlorisinatical specific line triphosphateATGAlorisinatical specific line triphosphateATGAlorisinatical calorATGAlorisinatical calorATGAlorisinational calorATGAlorisinat	Αβ	Amyloid-beta
PRKAA2Protein kinase AMP-activated catalytic subunit alpha 2ATGAudopagy relatedANOVAAnalysis of varianceATTAdenosine triphosphateAGCApoptosis-associated speck-like protein containing a CaspaseBECNIBeclin 1BACNBovine serum albuminCAMKACalcium/calmodulin-dependent protein kinase kinaseCASPICalspase-1CASPICalspase-1DAMKACalcium/calmodulin-dependent protein kinase kinaseDAMKACalcium/calmodulin-dependent protein kinase kinaseDAMFADamage-associated molecular patternsDAMFACalsace-1DAMFACalsace-1DAMADubecco's modified agle mediumDAMAEthlenediamine tetra actic acidDTAEthlenediamine tetra actic acidEDAFederal drug administrationFDAFederal drug administrationFDAGlaidifiel proteinFMAJousenal-associated molecule 1GAMAJousenal-associated molecule 1GAMAJousenal-associated molecule 1FDAFederal drug administrationFDAGlaidifiel proteinFDAJousenal-associated molecule 1GAMAJousenal-associated molecule 1GAMA <td< td=""><td>AD</td><td>Alzheimer's disease</td></td<>	AD	Alzheimer's disease
NTGAutophagy relatedANOVAAlayisa or varianceATUAlaosina trachapaseATUAlaosina scalada opechika protein containing a CaspaseASCBoin a containa containing a CaspaseBACMBoin a contain and anno anno	PRKAA2	Protein kinase AMP-activated catalytic subunit alpha 2
NANOVAAnalysis of varianceATPAdenosine triphosphateATPAdenosine triphosphateASCAportosis-associated speck-like protein containing a CaspaseERCN1Belin 1BACM2Borine serum albuminCAM4KAlfomycin A1CAM5K1Calcun/calmodulin-dependent protein kinase kinaseCAM5K2Caspase-1CAM5K3Alora factor kappa B subunitDAM5Danage-associated molecular patternsDAM5Jounder-DependiendienDAM6Obieco's modified eagle mediumDAM6Aloraterido eagle mediumDAM5Aloraterido eagle mediumDAM6Disco's modified eagle mediumDAM7Bielend right eagle mediumDAM8Disco's modified eagle mediumDAM8Disco's modified eagle mediumDAM9Disco's modified eagle mediumDAM9Disco's modified eagle mediumDAM9Disco's modified eagle mediumDAM9Disco's modified pagle mediumDAM9Disco's modified pagle mediumDAM9Disco's modified pagle	ATG	Autophagy related
ATPAdenosine triphosphateASCApptosis-associated speck-like protein containing a Caspase cruitment domainBECN1Belin 1BSABoine serum albuminBAABaflomycin A1CAMK8Caspase-1CAMK9Caspase-1DAMP0Danage-associated molecular patternsDAMP1Gadianizio-2-phenylindoleDMAMDisco's modified eagle mediumDMAMDisco's modified eagle mediumDMAMDisco's modified eagle mediumDMAMDisco's modified eagle mediumDMAMEdubarice trat accit acidDMAMDisco's modified eagle mediumDMAMDisco's modified eagle mediumDMAMEagle discover medica eagleDMAMEagle discover medica eagle	ANOVA	Analysis of variance
ASCApoptosis-associated speck-like protein containing a CaspaseBECN1Beclin 1BAABerlin and the protein and the protein kinaseBAABafiomycin A1CAMKKCalciun/calmodulin-dependent protein kinase kinaseCASP1Caspase-1ArAMKNuclear factor kappa B subunitDAMP3Jamage-associated molecular patternsDAMP4Jouneco's modified agle mediumDMAMDimethyl SulfoxideEGUEnhanced chemiluminescenceEGUAEndyeneliamine trat acetic acidEGNAEndyeneliamine tratactic acidFBAFedral drug administrationFBAGial forlilary acidic proteinFBAJourentyl SulfoxideFBAGial forlilary acidic proteinFBAFedral activationFBAJourentaine maleateFGAJourentainescenceFIAJourentainescenceFBAJourentaine maleateFGAGial forlilary acidic proteinFBAJourentaine maleateFIAJourentainescenceFIAJourentainescenceFIAJourentainescenceFIAJourentainescenceFIAJourentainescenceFIAGial forlilary acidic proteinFIAJourentainescenceFIAJourentainescenceFIAJourentainescenceFIAJourentainescenceFIAJourentainescenceFIAJourentainescenceFIAJourentainescenceFIAJourentainescence	ATP	Adenosine triphosphate
BECN1Beclin 1BSABovine serum albuminBAABaflomycin A1CAMKKCalcum/calmodulin-dependent protein kinase kinaseCAMKDCaspase-1CAMPBanage-associated molecular patternsDAPMJonage-associated molecular patternsDAMMJonage-associated molecular patternsDMAMJolbecco's modified eagle mediumDMAMDimethyl SulfoxidePMSODimethyl SulfoxideFDAEthanced chemiluminescenceFDAEthalbovine serumFDAFederal drug administrationFDAFolderal drug administrationFDAGlaid fortilary acidic proteinFDAIonized calcum-binding adapter molecula 1FDAJonized calcum-binding adapter molecula 1FDAJonage-associated membrane protein 1FDAJongolysaccharide	ASC	Apoptosis-associated speck-like protein containing a Caspase recruitment domain
BSABovine serum albuminBaf. A1Bafilomycin A1CAMKKCalciun/calmodulin-dependent protein kinase kinaseCAMKNCaspase-1CASP1Sapase-1DAMPaDanage-associated molecular patternsDAMPaJandarg-associated molecular patternsDAMMJublecco's modified eagle mediumDMSUDinethyl SulfoxideFDABinhaned chemiluminescenceEDTABindynel serumEDTAFeal bovine serumFDAFeal dovine serumFDAFeal dovine serumFDASildi forillary acidic proteinFDAJoizecalischemistrationFDAAiceradischemistrationFDAGainericularisdi adapter molecule 1FDAJoizecalischemistryFDAJoizecalischemistryFDAJoizecalischemistryFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDAJoinelylascharideFDA <td>BECN1</td> <td>Beclin 1</td>	BECN1	Beclin 1
Baf. A1Bafilomycin A1CAMKKCakium/calmodulin-dependent protein kinase kinaseCASPICakapae-1CASPINuclear factor kappa B subunitDAMPaJomage-associated molecular patternsDAMPaJomage-associated molecular patternsDMFMJoubecco's modified eagle mediumDMSODimethyl SulfoxideENTAEhhanced chemiluminescenceEDTAEtal bovine serumFDAFedral drug administrationFDAFedral drug administrationFDAGildi fibrillary acidic proteinFMAJonzeradish peroxidaseFMAJonzeradish peroxidaseFMAJonzeradish peroxidaseFMAJonzendish peroxidaseFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJouroinformangenFMAJouroinformangenFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJouroinformangenFMAJouroinformangenFMAJouroinformangenFMAJouroinformangenFMAJopolysaccharideFMAJopolysaccharideFMAJopolysaccharideFMAJouroinformangenFMAJouroinformangenFMA <td>BSA</td> <td>Bovine serum albumin</td>	BSA	Bovine serum albumin
CAMKKCalcium/calmodulin-dependent protein kinase kinaseCASP1Caspase-1NF-RBNuclear factor kappa B subunitDAMPsDamage-associated molecular patternsDAMPaGubacco's modified eagle mediumDMS0Dubecco's modified eagle mediumDMS0Dimethyl SulfoxideECLEnhanced chemiluminescenceEDTAEthylenediamine tetra acetic acidFBSFetal bovine serumFDAFederal drug administrationFDAFoloreani munosorbent assayFBAGilaf fibrillary acidic proteinFMPJoizee calcium-binding adapter molecule 1FIRPIonized calcium-binding adapter molecule 1FIRPLight chain 3LAMP-1Light chain 3FIRPLight chain 3FIRPSosomal-associated membrane protein 1FIRPNeurofibrillary tanglesFIRPNeurofibrillary tanglesFIRPNeurofibrillary tanglesFIRPSosomal-associated membrane protein 1FIRPJight chain 3FIRPNeurofibrillary tanglesFIRPNeurofibrillary tangles	Baf. A1	Bafilomycin A1
CASP1Capase-1NF-RBNuclear factor kappa B subunitDAMPsJamage-associated molecular patternsDAMPa4.6-diaminido-2-phenylindoleDMEMJubecco's modified eagle mediumDMSOJuinethyl SulfoxideENAROEhhanced chemiluminescenceEDTAEhtylenediamine tetra acetic acidEDTAEhtalovine serumFDSFetal bovine serumFDAFoldovine farta acetic acidFDAFoldovine maleateGFAPGial fibrillary acidic proteinFDAJoized calcium-binding adapter molecula 1FDAJoiseonal-associated membrane protein 1FIAPJoisoanal-associated membrane protein 1FIAPLight chain 3FIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoinal-astociated membrane protein 1FIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoinal-astociated reporterin 1FIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAPJoipolysaccharideFIAP <td< td=""><td>САМКК</td><td>Calcium/calmodulin-dependent protein kinase kinase</td></td<>	САМКК	Calcium/calmodulin-dependent protein kinase kinase
NF-8BNuclear factor kappa B subunitDAMPsJomage-associated molecular patternsDAPI4.6-diamindo-2-phenylindoleDMEMJolbecc's modified eagle mediumDMSOJomethyl SulfoxideDMSOJomethyl SulfoxideFDABhanced chemiluminescenceEDTABidylenediamine tetra acetic acidEDTAFotal bovine serumFDAFetal bovine serumFDAFetal odi minutosorbent assayFDAFotal odi administrationFDAGidai fibrillary acidic proteinFMPGidai fibrillary acidic proteinFMPJoreardish peroxidaseFMAInized calcium-binding adapter molecule 1FMAJosomal-associated membrane protein 1FMAJosomal-associated protein 2FMAJosomal-associated protein 2<	CASP1	Caspase-1
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DMSO0Dimethyl SulfoxideFCLEnhanced chemiluminescenceFDTAEthylenediamine tetra acetic acidFDTAEthylenediamine tetra acetic acidFISAForzme-linked immunosorbent assayFBSFetal bovine serumFDAFederal drug administrationFXNFluvoxamine maleateGFAPGlial fibrillary acidic proteinHEPES4/2-hydroxyethyl)-1-piperazineethanesulfonic acidHRPHorseradish peroxidaseIRAInized calcium-binding adapter molecule 1ILAMP-1Lysosomal-associated membrane protein 1LS3Light chain 3ILTGMarmalian target of rapamycinNTRNoD-like receptorNLRNDL-like receptorSURPJustafamily pyrin domain containing 3OCDObessive-compulsive DisorderPEGPolyethylene glycol	DMEM	Dulbecco's modified eagle medium
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OCD Obsessive-compulsive Disorder PBS Phosphate buffer saline PEG Polyethylene glycol	NLRP3	NLR family pyrin domain containing 3
PBS Phosphate buffer saline PEG Polyethylene glycol	OCD	Obsessive-compulsive Disorder
PEG Polyethylene glycol	PBS	Phosphate buffer saline
	PEG	Polyethylene glycol

(Continued)

PFA	Paraformaldehyde
PMSF	Phenylmethylsulfonyl fluoride
PVDF	Polyvinylidene difluoride
PYD	Pyrin domain
Rap	Rapamycin
RIPA	Radioimmunoprecipitation assay
SDS	Sodium dodecyl sulfate
SDS- PAGE	SDS poly acrylamide gel electrophoresis
SQSTM1	Sequestosome 1
ULK-1	unc-51-like kinase 1

Continued