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The Potential Therapeutic Effects of THC on Alzheimer's Disease

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Abstract. The purpose of this study was to investigate the potential therapeutic qualities of Δ^9 -tetrahydrocannabinol (THC) 10 with respect to slowing or halting the hallmark characteristics of Alzheimer's disease. N2a-variant amyloid- β protein precursor 11 $(A\beta PP)$ cells were incubated with THC and assayed for amyloid- β $(A\beta)$ levels at the 6-, 24-, and 48-hour time marks. THC was 12 also tested for synergy with caffeine, in respect to the reduction of the A β level in N2a/A β PPswe cells. THC was also tested 13 to determine if multiple treatments were beneficial. The MTT assay was performed to test the toxicity of THC. Thioflavin T 14 assays and western blots were performed to test the direct anti-AB aggregation significance of THC. Lastly, THC was tested 15 to determine its effects on glycogen synthase kinase-3 β (GSK-3 β) and related signaling pathways. From the results, we have 16 discovered THC to be effective at lowering AB levels in N2a/ABPPswe cells at extremely low concentrations in a dose-dependent 17 18 manner. However, no additive effect was found by combining caffeine and THC together. We did discover that THC directly interacts with A peptide, thereby inhibiting aggregation. Furthermore, THC was effective at lowering both total GSK-3 plevels 19 and phosphorylated GSK-3ß in a dose-dependent manner at low concentrations. At the treatment concentrations, no toxicity was 20 observed and the CB1 receptor was not significantly upregulated. Additionally, low doses of THC can enhance mitochondria 21 function and does not inhibit melatonin's enhancement of mitochondria function. These sets of data strongly suggest that THC 22 could be a potential therapeutic treatment option for Alzheimer's disease through multiple functions and pathways. 23

Keywords: Alzheimer's disease, amyloid-β peptide, cannabinoid, CB1 receptor, CB2 receptor, delta(9)-tetrahydrocannabinol, 24 neurodegeneration 25

INTRODUCTION 26

In 2011 alone, 15 million family members have pro-27 vided more than 17.4 billion hours of care to diagnosed 28 Alzheimer's disease (AD) patients. That care translates 29 into more than \$210 billion of AD-related services [1]. 30 This disease translates into an enormous burden on 31 32

caregivers, as well as the health care system, both med-

ically and economically. To date, there have been no effective treatments developed to cure or delay the progression of AD [2, 3]. By 2050, an estimated 11 to 16 million Americans will be living with the disease [1, 4].

AD pathology can be divided into two cate-38 gories, familial inherited AD and sporadic AD. The 39 histopathologies of early onset familial AD and late 40 onset sporadic AD are indistinguishable. Both forms of 41 AD are characterized by extracellular amyloid- β (A β) 42 peptide, and by amyloid plaques and tau-containing 43 neurofibrillary tangles [3]. The misfolded structure of 44 the A β peptides generates a characteristic tendency 45 for their aggregation [5]. It has long been believed 46

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that $A\beta_{1-40}$ (A β_{40}) and $A\beta_{1-42}$ (A β_{42}) aggregates are 47 the constituents of the insoluble plaques that are char-48 acteristic of AD. This disease is also associated with 49 neuroinflammation, excitotoxicity, and oxidative stress 50 [6, 7]. However, the continuous aggregation of $A\beta$ 51 peptides along with hyperphosphorylation of the tau 52 protein inside the cell, causing neurofibrillary tangle 53 formation, are generally accepted as the major etiolog-54 ical factors of the neuronal cell death associated with 55 the progression of AD [8–10]. 56

Recent studies have also suggested that glycogen 57 synthase kinase 3 (GSK-3) has a key role in the patho-58 genesis of both sporadic and familial AD [11, 12]. It 59 has been reported that GSK-3ß induces hyperphos-60 phorylation of tau [13–17]. Moreover, overexpression 61 of GSK-3 in Tet/GSK-3ß mice reveal pathological 62 symptoms that correspond to AD pathology with 63 respect to spatial learning deficits, reactive astrocyto-64 sis, increased AB production, and plaque associated 65 inflammation, as well as tau hyperphosphorylation 66 resulting in Aβ-mediated neuronal death [18]. Addi-67 tionally, chronic lithium (GSK-3 inhibitor) treatment 68 in double transgenic mice overexpressing GSK-3B and 69 tau has shown to prevent tau hyperphosphorylation 70 and neurofibrillary tangle formation [19]. Some reports 71 have also indicated that GSK-3α plays a role in regu-72 lating amyloid- β protein precursor (A β PPP) cleavage, 73 resulting in increased AB production [20, 21]. It has 74 also been shown that the $A\beta$ load in mouse brain can 75 be robustly ameliorated by the inhibition of GSK-3β 76 [22]. 77

Along with past research suggesting an involve-78 ment of GSK-3 in the pathogenesis of AD, there 79 have also been recent studies suggesting the intricate 80 involvement of the cannabinoid system in AD. It was 81 reported that the cannabinoid system can limit the neu-82 rodegenerative processes that drive the progression of 83 the disease, and may provide a new avenue for dis-84 ease control [23]. Currently the complete pathway and 85 mechanism of action of the cannabinoid system are 86 unknown, however, studies have been conducted to 87 determine the involvement of the cannabinoid $1 (CB_1)$ 88 and cannabinoid $2(CB_2)$ receptors in AD brain [6]. The 89 CB1 receptor is abundant in the brain and contributes to 90 learning, memory, and cognitive processes which are 91 interrupted early in the course of AD [24]. To the con-92 trary, CB₂ receptor expression is more limited and has 93 been anatomically found in neurons within the brain-94 95 stem [25], cerebellum [26], and microglia [27]. Recent research has also investigated the propensity of endo-96 cannabinoid receptor sub-types 1 (CB₁) and 2 (CB₂) 97 to elicit a neuroprotective and anti-inflammatory effect 98

on the brain when stimulated by endocannabinoids [28]. Postmortem studies of AD brains have detected increased expression of CB1 and CB2 receptors on microglia within the plaque, while CB₁ expression is reduced in neurons more remote from the plaque [29]. It is also noted that the endocannabinoid metabolizing enzyme, fatty acid amide hydrolase, is upregulated in the plaque [30]. There is also an increase in expression of anandamide metabolites, such as arachidonic acid, in the vicinity of the plaque [30]. These findings may indirectly suggest that the increase in CB₁ and CB₂ receptors may be to offset the lack of activity with their ligands due to increased metabolic activity of fatty acid amide hydrolase. These alterations in the cannabinoid system suggest an involvement of endogenous cannabinoids in the pathogenesis of AD or that this system may be altered by the pathophysiology of the disease [6]. Understanding that microglial activation is reserved in all cases of AD, it is important to identify that endogenous cannabinoids prevent ABinduced microglial activation both in vitro and in vivo [31]. These receptors are known to experience time dependent and brain region specific alterations during neurodegenerative and neuroinflammatory disorders to attempt to counteract excitotoxicity and inflammation [32].

Endocannabinoid receptors, CB₁ and CB₂, have been reported to interact with the endocannabinoid molecules: 2-arachidonoyl glycerol and anandamide. However, it has also been reported that CB₁ and CB₂ also react interact with Δ^9 -tetrahydrocannabinol (THC) isolated from the *Cannabis sativa* plant [33]. Furthermore, early reports indicate that Dronabinol, an oil-based solution of Δ^9 -THC, improves the disturbed behavior and stimulates appetite in AD patients [34], and alleviates nocturnal agitation in severely demented patients [35]. Accumulated evidence also suggests antioxidants having anti-inflammatory and neuroprotective roles [23].

It has also been shown that THC can decrease the level of A β -induced increases in reactive oxygen species, decreases in mitochondrial membrane potential, and caspase (a protein that is intimately involved in the regulation of apoptosis) activation, as well as protect human neurons from oligomeric A β -induced toxicity [36]. While it is understood that cannabinoids are active against inflammation, our research investigated the neuroprotective properties of THC, the active component of marijuana. Here we evaluated: 1) the effects of THC against A β expression in N2a/A β PPswe cells against the effects of caffeine, a reported A β expression suppressor [37]; 2) the direct

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effects of THC against A β aggregation, one pathological marker of AD; 3) the mechanism behind the anti-pathological properties of THC on AD; 4) the toxicity of THC and caffeine individually; and 5) the effects of THC on GSK-3 β and other related signal pathways in N2a/A β PPswe cells.

157 MATERIALS AND METHODS

158 Drugs used in this study

THC solution was purchased from Sigma (T4764-1ML Sigma Aldrich); caffeine was purchased from Sigma (C0750-100G, Sigma Aldrich); melatonin was purchased from Sigma (M5250-5G, Sigma Aldrich).

163 ELISA for detection of total $A\beta$ in protein samples

50 µl of goat anti-PWT1-42 antibody solution was 164 added to the sample and incubated overnight, followed 165 by a 1-hour incubation with 0.1% I-block buffer. The 166 tissue culture supernatant was diluted 1:10 with dilu-167 ent buffer containing a protease inhibitor. Standards 168 (1000, 500, 250, 125, 62.5, 31.25 pg/ml) were prepared 169 by serial dilution. The plate was washed and 50 µl of 170 sample or standard was added with triplication. $50 \,\mu l$ 171 of both Biosource 40/42 (HS) (primary antibody) A β 172 and a standard solution was added to each well and 173 incubated for 3 hours followed by $5 \times$ wash with PBST. 174 100 µl prepared secondary antibody (1:350 anti-rabbit 175 HRP) was added and incubated at 37°C for 45 minutes 176 on a shaker. The plate was washed; TMB substrate 177 was added (100 µl) and incubated for 10-30 minutes 178 in the dark. The reaction was halted by adding $100 \,\mu$ l 179 stop solution for detection at 450 nm. A 4 parameter 180 regression was used for the standard. 181

182 *Cell culture and drug treatment*

N2a/ABPPswe cells, N2a cells stably expressing 183 human ABPP carrying the K670N/M671L Swedish 184 mutation (ABPPswe), were grown in Dulbecco's modi-185 fied Eagle medium containing 10% fetal bovine serum, 186 100 U/mL penicillin, 100 µg/mL streptomycin, and 187 400 µg/mL G418 (Invitrogen), at 37°C in the pres-188 ence of 5% CO2. N2a/ABPPswe cells were diluted 189 with medium to a concentration of 2×10^{5} /ml, and 190 plated into the each well in 3 ml. 2 ml of trypsin was 191 incubated at room temperature, or 37°C. When most of 192 the cells began to float, trypsin was decanted and 5 ml 193 of fresh pre-warmed medium was added. Pipetting was 194 performed more than 30 times to ensure cells were sep-195

arated into individual cells. One drop of medium was 196 put into 1.5 ml tubes for counting; 10 µl of trypan blue 197 and 10 µl of medium of cells were added and applied to 198 cytometer for counting. The rule was total number of 199 cells of all for diagonal blocks/4 X 2 X 10000 = number 200 of cells/ml. The proper amount of cell medium and 201 fresh medium was added into new flasks according 202 to the ratio of dilution. Pipetting was performed 10 203 times to homogenize cells. 3 ml of cells were seeded 204 into medium into each 6 well plate. When one pipette 205 was used up, the cells were mixed in the flask before 206 using them for the next pipette. Compounds for screen-207 ing were resolved in DMSO, at 1000 fold to the final 208 concentration in the well. Pipetting of 10 µl solution, 209 then addition into 990 µl medium was performed; mix-210 ing followed. 12 hours after cells were plated, 400 µl 211 of compounds were added into 3.6 ml medium. The 212 medium was then removed from the six-wells. 3 ml of 213 medium with 1% DMSO was added to well 1; in well 214 2, 3 ml melatonin solution was added. In well 3, 4, 5, 215 and 6, compound solutions of 3 ml were added. 216

MTT assay

Cells were plated in 96-well tissue culture plate 218 at 10,000 cells/well, 100 µl/well. 100 µl THC solu-219 tion was added at $2 \times$ concentrations in each well. 220 Control groups are: 1) cells without THC treatment, 221 cells and fresh medium only; and 2) blank, wells with 222 medium without cells. All wells were replicated. Wells 223 were incubated for 36 hours. Cell proliferation kit 224 (Roche 11465007001) was then applied for toxicity 225 assay according to the standard protocol. 10 µl of MTT 226 reagent was first added to each well and incubated at 227 37°C for 4 hours. Then 100 µl of solubilization solu-228 tion was added to each well. These were incubated 229 overnight and optical density (OD) values were read at 230 575 nm. The percentage of cell viability was calculated 231 as: Cell viability % = (OD - OD blank) / (OD control - OD blank) / (232 OD blank) 233

Western blot for anti-aggregation assay

HFIP pretreated $A\beta_{1-40}$ peptide were obtained 235 from Biomer Technology, California. AB1-40 pep-236 tide solution was prepared in Ham's F-12 solution to 237 concentrations of 200 µM as stock. In the 15 µl aggre-238 gation system: 1) THC at final concentration of 25 239 nM, 2.5 nM, or 0.25 nM; and 2) 1.5 µl peptide stock 240 solution was added. Then 15 µl with F-12 medium 241 was made. Aggregation was allowed for 48 hours at 242 $37^{\circ}C$. After incubation, isomers of A β peptide were 243

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separated by 12% Tris-Tricine gel electrophoresis at 244 100 V for 180–210 minutes, a temperatures under 4°C. 245 The protein was transferred to PVDF membrane with 246 semi-dry transfer at 200 mA for 70 minutes. For west-247 ern blot detection, the membrane was blocked with 248 1.5% BSA solution in PBST solution (0.5%), then incubated in 1st Antibody: 6E10 (Signet) 1 mg/ml, 250 diluted by 1:1000 dilution in blocking buffer. It was 251 then washed 3 times with $1 \times PBST$ solution, and 252 then incubated with second antibody (anti-mouse IgG-253 HRP sigma A9044. 1:5000 diluted in blocking buffer). 254 After membrane was developed, film with bands were 255 scanned, followed by analysis of gel-quantification 256 software (QuantityOne, from Bio-rad). 257

258 Thioflavin T fluorescence assay

HFIP pretreated A β_{1-40} peptide was obtained from 259 Biomer Technology, California. In thioflavin T (ThT) 260 solutions (1.6 µg/ml dissolved in 20 mM Tris-HCL), 261 THC solution was prepared at concentration of 250, 262 25, 2.5, and 0.25 nM. THC solution was added contain-263 ing ThT buffer into black 96 well plates. Unaggregated 264 Aβ peptide solution was thawed, diluted, and imme-265 diately added to wells, making the final concentration 266 of A β_{1-40} at 1 μ M. Control groups were setup as: 1) 267 aggregation control; 2) control with ThT buffer only; 268 and 3) Tris-HCl buffer only. Plate was mixed and flu-269 orescence was read at 482 nm with excitation 440 nm 270 with Biotek All-in-One plate reader. Fluorescence was 271 screened for 2 hours with 5-minute intervals. 272

Western blot for total and phosphorylated GSK-3β, total tau and phosphorylated tau, and β-actin

Followed by THC treatment in tissue culture, 275 N2a/AβPPswe cell lysate were collected, quantified, 276 and aliquoted. Using 12% Tris-Glycine gel system 277 (Biorad), protein were separated by electrophoresis 278 and semi-dry transferred to PVDF membrane. GSK-3β 279 and β -actin antibodies were used as primary antibody. 280 After adding secondary antibody, the membranes were 281 exposed using ECL substrate (Pierce). After mem-282 brane was developed, film with bands were scanned, 283 followed by analysis of gel-quantification software 284 (QuantityOne, from Biorad). 285

Mitochondria isolation and respiratory measurements

The respiratory function of isolated mitochondria was measured using a miniature Clark type oxygen electrode (Strathkelvin Instruments, MT200A chamber, Glasgow, UK). Detail method is published in Dragicevic et al. [38].

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Statistical analysis and graphs

All data were analyzed with one-way ANOVA and *post hoc* analysis was conducted with Turkey's group analysis and p < 0.05 was considered as statistical significance (GraphPad 6.0). All graphs were graphed with GraphPad 6.0 software.

RESULTS

THC can decrease Aβ level in N2a/AβPPswe

ELISA assay was performed for $A\beta_{40}$ levels in N2a/ABPPswe cells 6 hours after cells were treated at different concentrations individually with THC, and caffeine—a reported compound to lower serum A β_{40} levels in a mouse model [39]—showed a significant reduction in AB40 levels of THC and caffeine versus the control (Fig. 1A). However, 24 hours after treatment of N2a/A β PPswe cells, A β_{40} concentrations were measured again in the THC treated cells versus the control. An increasing difference in $A\beta_{40}$ concentrations were noted in both THC treated cells and caffeine treated cells in a dose-dependent manner (Fig. 1B). The assay was performed again, 48 hours after treatment of N2a/ABPPswe cells with THC versus the control at each concentration of the drugs originally used. THC-treated N2a/ABPPswe cells significantly differed more in $A\beta_{40}$ concentrations versus the control then at the 6- and 24-hour time point. The significant difference was conserved and greater over each increasing dose of THC and caffeine administered versus the control (Fig. 1C). These data suggest THC's and caffeine's inherent anti-A β_{40} properties are time and dose dependent in N2a/ABPPswe cell models. This data also reveals that THC may delay of halt the progression of AD by inhibiting the production of $A\beta_{40}$ peptide in the central nervous system.

Synergy between THC and caffeine on $A\beta_{40}$ concentration in N2a/A β PPswe cells

THC and caffeine were assayed for a synergistic effect on $A\beta_{40}$ concentration in N2a/A β PPswe cells (Fig. 2). However, no synergistic properties of THC and caffeine are seen as there is no significant difference in the concentration of $A\beta_{40}$ in N2a/A β PPswe cells solely treated with THC as compared to cells



Fig. 1. (A) $A\beta_{40}$ (pg/ml) *in vitro* measured 6 hours from incubation in N2a/AβPPswe cells. Three groups of cells were assayed: 1) those that were not treated with THC; 2) those that were treated with THC; and 3) those that were treated with caffeine. Treatment in both the THC group and in the caffeine group resulted in a dose-dependent decrease in A β_{40} concentration after 6 hours. There are no significant differences among all groups (p > 0.05). The concentrations of THC from A to F are 0 nM, 0.25 nM, 2.5 nM, 25 nM, 250 nM, and 2500 nM respectively, and concentrations of caffeine from A to F are 0 μ M, 0.625 μ M, 1.25 μ M, 2.5 μ M, 5 μ M, and 10 μ M, respectively, (B) A β_{40} (pg/ml) *in vitro* measured 24 hours from incubation in N2a/AβPPswe cells. A dose-dependent decrease in concentration of A β_{40} was still observed. THC: A, B, C, versus F are p < 0.05 and all other groups in comparison are p > 0.05. Caffeine: A versus B, B versus E, and all other groups versus F are p < 0.05. The concentrations of THC from A to F are 0, 0.25 nM, 25 nM, 250 nM, and 2500 nM, respectively, and concentrations of raffeine from A to F are 0, 0.25 mM, 2.5 nM, 25 nM, 250 nM, and 2500 nM, respectively, and concentrations of caffeine from A to F are 0, 0.25 nM, 2.5 nM, 25 nM, 250 nM, and 2500 nM, respectively, and concentrations of caffeine from A to F are 0, 0.625 μ M, 1.25 μ M, 5 μ M, and 10 μ M, respectively, (C) A β_{40} (pg/ml) *in vitro* measured 48 hours from incubation in N2a/A β PPswe cells. A dose-dependent decrease in concentrations of THC from A to F are 0, 0.55 nG are 0.05. The concentrations of THC from A to F are 0, 0.25 nM, 2.5 μ M, 30 nM, respectively, (C) A β_{40} (pg/ml) *in vitro* measured 48 hours from incubation in N2a/A β PPswe cells. A dose-dependent decrease in A β_{40} (pg/ml) in conserved. THC groups: p > 0.05 for A versus B, and all other groups are p < 0.05. Caffeine groups: p < 0.05 for B versus D, and all other comparisons between groups are p > 0.05. The

treated with $2.5 \,\mu$ M caffeine and THC at various concentrations.

Repeated treatment can continuously decrease Aβ production

Our data also illustrates N2a/ABPPswe cells treated 339 with THC twice, 24 hours apart from each treatment, 340 showed a significant decrease in AB40 concentra-341 tion compared to cells treated once (Fig. 3A). While 342 the decrease in $A\beta_{40}$ expression is not observed 343 at concentration close to $10 \,\mu$ M, they are seen at 344 $25 \,\mu M$ and greater suggesting multiple treatments 345 may be efficacious in reducing $A\beta_{40}$ concentration in 346 N2a/ABPPswe cells and animal models. 347

Cell toxicity detection of THC on N2a/AβPPswe cells

THC was also measured for toxicity versus the caffeine and the untreated N2a/AβPPswe cells, which served as the control. The MTT assay showed no significant difference from the control for toxicity as compared to each concentration of THC and caffeine administer suggesting THC and caffeine lack toxicity to the cells at each concentration assayed (Fig. 3B).

THC can inhibit $A\beta_{40}$ aggregation as shown by ThT assay and western blot

The ThT assay was to exhibit the direct interaction THC has with $A\beta$ demonstrates that as the concen-



Fig. 2. $A\beta_{40}$ (pg/ml) concentration in N2a/A β PPswe cells at various drug concentrations among groups. Treatment with both THC and caffeine resulted in a dose-dependent decrease in $A\beta_{40}$ concentration. However, no synergistic effect was observed.

tration of THC added to the assay was increased, the 361 intensity of fluorescence in AB decreased. This data 362 suggests that $A\beta$ peptide directly binds to THC and 363 prevents the uptake of fluorescence (Fig. 4A). More-364 over, our lab performed an additional ELISA assay to 365 confirm that the interaction of the AB peptide with THC 366 did not shield amino acids 1-10, the major B-cell epi-367 tope [40] (Fig. 4B). There is no significant difference in 368 absorbance at each concentration of THC, indicating 369 that at each concentration of THC the $A\beta$ antibodies 370 were able to bind with equal distribution and affinity. 371



Fig. 3. (A) $A\beta_{40}$ (pg/ml) concentration N2a/A β PPswe cells treated with THC, as well as the $A\beta_{40}$ (pg/ml) concentration of N2a/A β PPswe cells treated with THC twice, 24 hours apart. The number of treatments has shown to decrease the concentration of $A\beta_{40}$ (pg/ml), (B) This shows the data obtained from the reduction of MTT at different concentrations of THC versus the different concentration of caffeine. Untreated N2a/A β PPswe cells were also assayed to compare with the MTT reduction of N2a/A β PPswe cells treated with THC and caffeine at different concentrations.



Fig. 4. (A) ThT assay measuring the fluorescence of Thioflavin T which binds to β -sheet structure of A β aggregation. With addition of THC, dose-dependent decreases in intensity of fluorescence indicates THC directly interferes with the binding of ThT to A β peptide. (B) THC incubated with A β peptide to determine the occurrence of THC interference with the major B cell epitope. No identified interference was observed at each increasing concentration of THC.



Fig. 5. (A) Polyacrylamide gel from a western blot indicating the concentration of aggregated A β peptide with and without the treatment of THC at various concentrations. Groups: 1: Aggregation control; 2: THC 100 nM; 3. THC 10 nM; 4: THC 1 nM, (B) THC anti-aggregation assay gel quantification indicating the relative percent monomeric A β .



Fig. 6. ELISA assay elucidating a possible mechanism through which THC functions to decrease the synthesis of AB in N2a/ABPPswe cells. AB level increases at 36 hours and reaches its peak level at 48 hours. Follow this mark; it then starts decreasing at 60 hours. The drug treatment benefit time is seen at 36 hours and last to 48 hours (the best window time). THC can significantly lower AB and this function can be partially blocked by CB1 antagonist Rimon at 10^{-4} M. However, inhibition function is lost at 10^{-2}

Therefore, we can postulate that THC's direct interac-372 tion with the A β peptide will not dampen an immune 373 response to clear the A β peptide. 374

Further analysis with western blot was performed 375 measuring the anti-aggregation properties of THC with 376 A β peptide. At each increasing concentration of THC, 377 a higher relative % of A β monomer was observed 378 correlating with a lower intensity of aggregated AB 379 peptide. This data suggests the direct interaction of 380 THC with $A\beta$ peptide and its ability to bind to the 381 peptide and inhibit aggregation (Fig. 5A,B). 382

CB1 receptor antagonist can partially rescue $A\beta$ 383 level inhibited by THC 384

An ELISA was performed to determine the mech-385 anism of THC in supporting the reduction of $A\beta$ in 386 N2a/ABPPswe cells. A known inhibitor of the CB1 387 receptor, rimonabant, was mixed with THC at differ-388 ent concentrations. Untreated N2a/ABPPswe cell AB 389 concentrations were used as a control. It was noted that 390 a dose dependent increase in $A\beta$ was observed as the 391 concentration of the inhibitor was increased. A time 392 dependent effect of the inhibitor was also witnessed as 393 the assay was repeated at the 12-, 36-, 48-, 60-, and 394 72-hour mark (Fig. 6). Due to increasing AB concen-395 trations as the inhibitor concentration is increased, this 396 suggests that THC partially functions through the CB1 397 receptor to mediate the synthesis of AB. The RT-PCR 398 results for CB1 receptor expression level showed that 399

there is no significant upregulation by THC to CB1 receptor (data not shown).

THC can inhibit total GSK-3β and phosphorylated $GSK-3\beta$ (pGSK-3 β) production

The western blot assay performed to examine the 404 effect of THC on GSK-3ß exhibits a dose-dependent 405 decrease in GSK-3β. β-actin, a housekeeping gene, 406 was used as a control to indicate that GSK-3 β was 407 expressed at a constant rate and that the changes in 408 intensity are not related to the change in expression 409 amount. As shown in Fig. 7A-D, this data suggests that 410 THC is efficacious in modulating and ameliorating the 411 expression of GSK-3ß and could decrease neuronal 412 apoptosis by down regulating GSK-3β. 413

THC can inhibit phosphorylated (pTau) production, but not affect $A\beta PP$ production

We detected pTau and ABPP levels among different 416 treatment conditions. THC can lower pTau expression 417 level with dose-dependent administration, but we did 418 not see the differences in ABPP levels detected with 419 6E10 antibody (Fig. 8A-F). 420

THC can enhance mitochondrial function but will not interfere with melatonin's enhancement of the mitochondria

Isolated mitochondria from N2a/ABPPswe cells showed higher oxygen utilization when treated with THC. When combined with melatonin, the function of the mitochondria is not altered (Fig. 9A, B).

DISCUSSION

Advances in therapeutics to prevent AD, or delay the 429 progression, are currently being made. Recent research 430 has shown caffeine and coffee are effective in limiting cognitive impairment and AD pathology in the trans-432 genic mouse model by lowering brain AB levels, which 433 are thought to be central to the pathogenesis of AD [41]. 434 Similarly, the current study shows the in vitro anti-AB activity of caffeine, and of another naturally occurring compound, THC. 437

N2a/ABPPswe cells were incubated separately with 438 various concentrations of caffeine, melatonin, and 439 THC. The relative anti-A β effect of THC was observed 440 to increase in a time dependent manner. A dose-441 dependent decrease in AB concentration was noticed 442 at lower concentrations of THC, as compared to caf-443

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Fig. 7. (A) A western blot performed to determine the effects of THC on GSK-3 β in N2a/A β PPswe. β -actin was used as a control to indicate that the expression rate was constant. The left indicator is molecular weight. Lane 1, 2, and 3 are β -actin level and lane 4, 5, and 6 are GSK-3 β expression. 1 and 4 are cell controls, 2 and 5 are cells treated with 2.5 nM THC, and lane 3 and 6 are cells treated with 0.25 nM THC. THC can inhibit GSK-3 β level at 2.4 nM concentration, (B) Graph representing the expression decrease in GSK-3 β in a dose-dependent manner by using β -actin to obtain a value for the ratio of expressed GSK-3 β . As shown in the bar graph, the total GSK-3 β decrease after using β -actin standardized protein loading, (C) GSK-3 β expression in N2a/A β PPswe treated with different drugs: Cells were plated in 6 well plate for overnight and then drugs were added into each designated wells in duplicate. Cells were lysed after 36 hours incubation. Proteins were loaded onto SDS-page gel and then blotted with each antibody after transfer onto PVDF membrane. Groups are: CTRL, Control; M1T2, 10⁻⁵ M Melatonin + 2.5 nM; T2, THC 2.5 nM; T3, THC 0.25 nM, (D) Expression of pGSK-3 β following melatonin and THC treatment in N2a/A β PPswe cells. *The same batch protein samples were used in this test as in Fig. 7C. Bands were quantified. One-way ANOVA was applied to the data. *p* < 0.05 when compared with control group. ***p* < 0.01 when compared with control groups are: Ctrl, Control; M1T2, 10⁻⁵ M Melatonin + 2.5 nM THC; M2T2, 10⁻⁶ M Melatonin + 2.5 nM; T2, THC 2.5 nM; T3, THC 0.25 nM THC; T1, THC 25 nM; T2, THC 2.5 nM; T3, THC 0.25 nM THC; T1, THC 25 nM; T2, THC 2.5 nM; T3, THC 0.25 nM THC; T1, THC 25 nM; T2, THC 2.5 nM; T3, THC 0.25 nM, T0] with control group. The control; M1T2, 10⁻⁵ M Melatonin + 2.5 nM THC; M2T2, 10⁻⁶ M Melatonin + 2.5 nM; T2, THC 2.5 nM; T3, THC 0.25 nM THC; T1, THC 25 nM; T2, THC 2.5 nM; T3, THC 0.25 nM.

feine. Further evidence shows that N2a/A β PPswe cells, treated twice with THC, show an even greater reduction in A β levels at slightly higher concentrations. Although it might have been predicted that caffeine and THC may function in a synergistic effect to reduce the A β load in N2a/A β PPswe cells, no synergy was observed. The MTT assay confirmed that cells treated at efficacious concentration of THC showed no toxicity, suggesting such a treatment to be safe and effective for further experimentation in the AD animal model. However, valid arguments have transpired in recent times regarding the concern for acute and long-term memory impairment with the use of THC. It has been



Fig. 8. (A) AβPP expression in N2a/AβPPswe treated with different drugs. The sample protein samples as in Fig. 7C were used for western blotting assay. 6E10 anti-Aβ antibody was used to detect AβPP and β-Actin was detected as protein loading control. Groups are: CTRL, Control; M1T2, 10^{-5} M Melatonin + 2.5 nM THC; M2T2, 10^{-6} M Melatonin + 2.5 nM THC; T1, THC 25 nM; T3, THC 0.25 nM, (B) Quantification result of AβPP in western blotting: We used quantification method to further compare the differences among drug treatment to AβPP level. There are no statistical significant differences among all treatment (*p* > 0.05). This data indicates that THC did not change AβPP expression level. Groups are: Ctrl, Control; M1T2, 10^{-5} M Melatonin + 2.5 nM THC; M2T2, 10^{-6} M Melatonin + 2.5 nM THC; T1, THC 25 nM; T3, THC 0.25 nM; (C) Tau expression in N2a/AβPPswe treated with different drugs. The sample protein samples as in Fig. 7C were used for western blotting assay. Anti-Tau and pTau antibodies were used to detect AβPP and β-Actin was detected as protein loading control. Groups are: CTRL, Control; M1T2, 10^{-5} M Melatonin + 2.5 nM THC; M2T2, 10^{-6} M Melatonin + 2.5 nM THC; M2T2, 10^{-6} M Melatonin + 2.5 nM THC; M2T2, 10^{-6} M Melatonin + 2.5 nM THC; T1, THC 25 nM; T3, THC 0.25 nM; T3, THC 0.25 nM; T3, THC 0.25 nM, (D) No significant difference of pTau expression shown among six groups in N2a/AβPPswe cells. THC treatment has no function to pTau expression. Groups are: Ctrl, Control; M1T2, 10^{-5} M Melatonin + 2.5 nM THC; M2T2, 10^{-6} M Melatonin + 2.5 nM THC; T1, THC 25 nM; T3, THC 0.25 nM, (E) Expression of pTau/Tau following melatonin and THC treatment in N2a/AβPPswe cells. +p < 0.05 when compared with THC 25 nM and THC 0.25 nM groups. *p < 0.01 when compared with THC 25 nM, T2, THC 2.5 nM; T3, THC 0.25 nM; T3, THC 0.25 nM; T2, THC 2.5 nM; T3, THC 0.25 nM; T2, THC 2.5 nM; T3, THC 0.25 nM; T2, THC 2.5 nM; T3, THC 0.



Fig. 9. (A) The enhancement of mitochondria function to cells treated with different: N2a/A β PPswe cells were cultured in 10 cm tissue culture plate and then treated with drugs for 36 hours and mitochondria were harvested and tested for their ability of using oxygen utilization. Ctrl, Control; M1T2, 10⁻⁵ M Melatonin + 2.5 nM THC; M2T2, 10⁻⁶ M Melatonin + 2.5 nM THC; T2, THC 2.5 nM; M1, 10⁻⁵ M Melatonin, (B) The enhancement of mitochondria function to cells treated with different: N2a/A β PPswe cells were cultured in 10 cm tissue culture plate and then treated with drugs for 36 hours and mitochondria were harvested and tested for their ability of using oxygen utilization. Ctrl, Control; M1T2, 10⁻⁵ M Melatonin + 2.5 nM THC; M2T2, 10⁻⁶ M Melatonin + 2.5 nM THC; T2, THC 2.5 nM; M1, 10⁻⁵ M Melatonin, (B) The enhancement of mitochondria function to cells treated with different: N2a/A β PPswe cells were cultured in 10 cm tissue culture plate and then treated with drugs for 36 hours and mitochondria were harvested and tested for their ability of using oxygen utilization. Ctrl, Control; M1T2, 10⁻⁵ M Melatonin + 2.5 nM THC; M2T2, 10⁻⁶ M Melatonin + 2.5 nM THC; T2, THC 2.5 nM; M1, 10⁻⁵ M Melatonin.

Table 1 Difference and percent decrease of $A\beta_{40}$ (pg/ml) in THC treated cells at 2.5 µg/ml compared with the control at different time points

Time Point	6 h	24 h	48 h
Control	1064.025	5303	5935.525
THC 2.5 µg/ml	965.827	3648.975	2894.175
Percentage of decreased $A\beta_{40}$	9.23%	31.19%	51.24%

shown that memory impairment was identified in rats treated with THC [42]. It should be clear, however, that the memory impairment observed occurred at con-centrations more than a thousand times higher than what is presented here as a beneficial treatment in AD model N2a/ABPPswe cells. The concentrations used in the study are considered to be extremely low, as the concentrations that we focused on in the study were from 2.5 nM of THC down to 0.25 nM of THC. Although some studies with ultra-low doses of THC have indicated neurotoxic roles [42], newer research shows a neuroprotective role and actually promotes elevation of phosphorylated cAMP response element-binding protein (pCREB) by increasing the levels of brain-derived neurotrophic factor in the frontal cortex [43]. Furthermore, the dosing used in our study is a lower concentration than that in the aforementioned research. Therefore, we believe that THC has a thera-peutic value, and that at low enough doses, the potential benefits strongly prevail over the risks associated with THC and memory impairment.

In addition to the Aβ concentration suppression,
benefits of THC, analyzed with a western blot and
ThT assay, confirmed anti-Aβ aggregate properties
by a dose-dependent decrease in fluorescence uptake,

and a decrease in intensity of aggregated A β in a dose-dependent manner. The positive results suggest possible intermolecular force interactions, preventing the molecular aggregation of A β peptides. The conducted ELISA, to ensure the intermolecular interaction of THC with A β did not block the major B-cell epitope, showed no interference with antibody binding, which indicated that regardless of the molecular interaction of THC with A β , an immune response should not be inhibited.

One pathway in which THC function was shown through the cannabinoid receptor inhibition with rimonabant. The dose- and time-dependent increase of $A\beta$ with respect to CB₁ inhibition was noted. It is likely that the time deference was observed due to the slow interaction of rimonabant with the CB₁ receptor. However, the difference in A β concentration becomes more evident at the later time points. Lastly, we showed a dose-dependent decrease in GSK-3 β expression influenced by THC.

To date, no $A\beta$ specific therapeutic options for AD have been approved. While progression is being made in this field, rigorous efforts focus on developing compounds that can address or possess the inhibition of A β synthesis and anti-A β aggregation properties or characteristics that down regulate GSK-3 β and pGSK-3 β . Our results demonstrate that THC possesses all of the above mentioned properties. All of these areas address major etiological characteristics of AD. GSK-3 β , pGSK-3 β , and A β -plaque brain concentrations, the hallmark of AD, are major targets for current AD research. Furthermore, we have shown that THC functions are pathway dependent of the endoge-

nous cannabinoid CB1 receptor recently discovered 516 to possibly function in AD disease modulation by 517 suppressing microglial activation upon receptor inter-518 action. Notwithstanding, it should also be noted that 519 low doses of THC are used to address the above men-520 tioned targets, thus avoiding risks induced by THC 521 associated with memory impairment and risks asso-522 ciated with toxicity. In addition, we also discovered 523 that low doses of THC can also enhance mitochon-524 dria function and has no negative drug interactions to 525 melatonin, a potential therapeutic for AD. 526

Here we have presented a promising compound that 527 addresses many major targets for AD therapeutics cur-528 rently being research. We have shown THC, at an 529 extremely low dose level (2.5 nM), has the proclivity to 530 slow or halt AD progression by dampening the synthe-531 sis of the major pathological marker of AD, AB. Also, 532 our lab has elucidated a potential mechanism respon-533 sible for the anti-pathological properties of THC with 534 respect to AD. Furthermore, we have clearly exhib-535 ited lack of toxicity at low concentrations of both THC 536 and caffeine individually. In conclusion, we believe the 537 multifaceted functions of THC will ultimately decrease 538 downstream tau hyperphosphorylation and neuronal 539 death thereby halting or slowing the progression of 540 this devastating disease. 541

542 DISCLOSURE STATEMENT

Authors' disclosures available online (http://www.jalz.com/disclosures/view.php?id=2309).

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