

Title

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The protective effect of dietary eicosapentaenoic acid against impairment of spatial cognition learning ability in rats infused with amyloid $\beta_{(1-40)}$

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Abstract

10 **Background:** Amyloid β (A β) peptide (1–40) can cause cognitive impairment.

Experimental design: We investigated whether dietary preadministration of eicosapentaenoic acid (EPA) is conducive to cognition learning 11 12 ability and whether it protects against the impairment of learning ability in rats infused with A β peptide (1-40) into the cerebral ventricle. **Results:** Dietary EPA administered to rats for 12 weeks before the infusion of $A\beta$ into the rat brain significantly decreased the number of 13 reference memory errors (RMEs) and working memory errors (WMEs), suggesting that chronic administration of EPA improves cognition 14 15leaning ability in rats. EPA preadministered to the A_β-infused rats significantly reduced the increase in the number of RMEs and WMEs, 16 with concurrent proportional increases in the levels of corticohippocampal EPA and docosahexaenoic acid (DHA) and in the DHA/ 17arachidonic acid molar ratio. Decrease in oxidative stress in these tissues was evaluated by determining the reactive oxygen species and lipid peroxide levels. cDNA microarray analysis revealed that altered genes included those that control synaptic signal transduction, cell 18 communication, membrane-related vesicular transport functions, and enzymes and several other proteins. 19

20 **Conclusion:** The present study suggests that EPA, by acting as a precursor for DHA, ameliorates learning deficits associated with 21 Alzheimer's disease and that these effects are modulated by the expression of proteins involved in neuronal plasticity.

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24 Keywords: Alzheimer's disease; Hippocampus; Spatial memory; Fatty acid; Rat; Amyloid β

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26 **1. Introduction**

Fish oil provides a host of health benefits because of its 27major polyunsaturated fatty acid (PUFA) components: 28eicosapentaenoic acid [EPA; C20:5(n-3)] and docosahex-29aenoic acid [DHA; C22:6(n-3)]. The beneficial effects of 30 fish oil on brain functions, however, have largely focused on 3132 and highlighted only DHA, the elongated/desaturated product of EPA. That is probably due to the fact that 33 DHA, but not EPA, constitutes the major PUFA of brain 34lipids: DHA alone constitutes >17% of the total fatty acids in 35 the rat brain [1], while EPA, as a precursor for DHA, 36 constitutes only a tiny percentage of the total fatty acids in 37

the brain, thus the extensive studies on the beneficial effects 38 of DHA on cognition learning ability. 39

Alzheimer's disease (AD) is a primary degenerative 40 disease of the central nervous system, and the histopatho-41 logical hallmark of AD is the presence of neurofibrillar 42 tangles and amyloid plaques of insoluble amyloid peptide 43 aggregates, which ultimately leads to dementia and 44 behavioral and cognitive impairments [2]. Epidemiological 45 studies show that intake of fish oil is associated with a 46 reduced risk of AD [3,4]. Chronic administration of DHA 47 improves spatial learning ability by increasing the level of 48 DHA in the hippocampus and cerebral cortex of young 49 and aged rats [5,6]. DHA administration also protects 50 against [7] and ameliorates [8] memory deficits in amyloid 51 β (A β)-peptide-induced AD model rats. DHA protects 52against behavior deficits and dendritic pathology in the AD 53mouse model [9]. 54

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Fig. 1. Schema and schedule of experimental rat groups.

There are few reports on the effects of pure EPA on 55cognition learning ability. Chronic administration of pure 56EPA attenuates interleukin $1\beta_k$ but does not significantly 57enhance memory in control rats, although whether their 58 normal laboratory chow diet contained any n-3 PUFAs is 59unclear [10]. n-3 PUFAs can induce gene expression with 60 61 concomitant effects on synaptic transmission [11] and related signal transduction. In a parallel set of experiments, 62we also investigated whether dietary EPA could induce 63 gene expressions related to its beneficial effects on learning-64related ability. EPA is probably taken into brain tissue, since 65 the presence of Δ^4 -desaturation enzymes in the brain is still 66 unclear [12]. Nonetheless, ¹⁴C-labeled EPA detected in the 67 rat brain 1 h after its oral administration to rats decreases 68 time dependently, while [14C]DHA, a metabolite of EPA, 69 increases time dependently [13]. It is also speculated that in 70the de novo system, each PUFA that is metabolized after 71 being taken into cerebral endothelial cells and astrocytes 72(constituent cells of the blood-brain barrier) is released 73 from those cells, and that DHA is taken into neurons as 74metabolite [14]. Here, we estimated the effects of the 75 chronic administration of pure EPA on spatial learning 76 ability in rats and examined whether EPA can protect 77 against the impairment of learning ability in $A\beta_{(1-40)}$ -78induced AD model rats. 79

80 2. Materials and methods

81 2.1. Animals and diet

The experimental schedule is shown in Fig. 1. Rats were 82 handled and killed in accordance with the procedures 83 outlined in the Guidelines for Animal Experimentation of 84 Shimane Medical University (Shimane, Japan), as compiled 85 from the Guidelines for Animal Experimentation of the 86 Japanese Association for Laboratory of Animal Science. 87 Wistar rats (first generation) (Jcl: Wistar; Clea Japan Co., 88 Osaka, Japan) were housed in a room under conditions of 89

controlled temperature $(23\pm2^{\circ}C)$, relative humidity 90 (50±10%) and light-dark cycles (light: 0800-2000 h; 91 dark: 2000-0800 h), and were provided fish-oil-deficient 92pellet diet (F-1; Funabashi Farm, Funabashi, Japan) and 93 water ad libitum. The inbred third-generation male rats 94(n=48; 5weeks old) were divided into two groups: an EPA 95group (n=24) administrated EPA-95E (300 mg/kg/day; 96 Mochida Pharmaceutical Co., Tokyo, Japan) dissolved in 97 5% gum arabic solution by gavage for 7 weeks initially, and 98 a control group (n=21) given 5% gum arabic solution only. 99

2.2. Preparation of Aβ-infused rats

The surgical techniques for preparing $A\beta$ -infused rats 101 were essentially the same as those described previously 102[7,8]. Briefly, each rat was anesthetized with sodium 103 pentobarbital (50 mg/kg body weight ip), and its skull was 104 exposed and drilled with two holes (right and left, relative to 105 bregma; 0.8 mm posterior, 1.4 mm lateral) in accordance 106 with the atlas of Paxinos and Watson using a stereotaxic 107 frame (Narishige, Tokyo, Japan). A solvent of 35% (vol/vol) 108 acetonitrile plus 0.1% (vol/vol) trifluoroacetic acid (pH 2.0) 109was used as vehicle for A β peptide (1–40) (Peptide Inst., 110Osaka, Japan). A miniosmotic pump (Alzet 2002; Durect 111 Co., Cupertino, CA, USA) containing either A β peptide (1– 112 40) solution (234±13.9 µl) or vehicle alone was quickly 113 implanted into the back of the rats. The outlet of the pump 114 was inserted 3.5 mm into the left ventricle and attached to the 115 skull with screws and dental cement. The infusion rate was 116 0.56μ l/h, and the total amount infused was approximately 117 4.9–5.5 nmol/L AB. Spontaneous infusion for 2 weeks thus 118 brought about completion of the volume used in the 119 miniosmotic pump. 120

2.3. Radial maze learning ability

Seven weeks after the start of EPA administration, the 122learning-related behavior of the rats was assessed by their 123completing a task in an eight-arm radial maze as previously 124described [5,8]: four reward pellets were placed randomly on 125four arms of the maze, and the number of total selections in 126obtaining the four pellets was counted. Two parameters of 127 memory function were examined: reference memory error 128(RME), which was determined by the number of entries into 129unbaited arms, and working memory error (WME), which 130was estimated by the number of repeated entries into arms 131that had already been visited within a trial. Lower numbers of 132RMEs and WMEs implied better spatial learning ability in 133 the rats. Performance was calculated from memory-related 134 behavior. Each rat was given two daily trials, 6 days/week for 135a total of 3 weeks. After completing the behavior test, each of 136the two groups of rats was subdivided into two groups 137 (allowing for the number of errors made by each rat in the 138last six trials of the preliminary behavior test) and infused 139with either $A\beta$ or the vehicle as follows: a control group was 140 divided into an AB-solvent-infused group [control (vehicle) 141 group; n=12)] and an A β -infused group (A β group; n=9); 142

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an EPA group was divided into a vehicle-infused EPA group 143 144 (EPA+vehicle group; n=12) and an A β -infused EPA group (EPA+A β group; *n*=12). The four groups of rats were again 145 behaviorally tested 3 weeks after the implantation of the 146miniosmotic pump to assess the effect of EPA preadminis-147 tration on the impairment of learning ability in AB-infused 148 rats. This testing lasted for a total of 3 weeks. The same 149protocol used for the preliminary behavior test was followed 150in the final behavior test, except for the adaptation periods. 151

152 2.4. Preparation of sample

After undergoing the behavioral tests for 3_1 days, the rats were anesthetized with sodium pentobarbital (65 mg/kg body weight, ip), blood was drawn for plasma analysis, and the hippocampus and cerebral cortex were separated as described previously [7]. The tissues were stored at -80° C by flash-freezing in liquid N₂ until use.

159 2.5. Measurement of fatty acid profile and oxidative status

The brain samples were immediately homogenized on ice 160 in 1.0 ml of ice-cold 0.32 mol/L sucrose buffer (pH 7.4) 161162containing 2 mmol/L EDTA, 0.5 mg/L leupeptin, 0.5 mg/L pepstatin, 0.5 mg/L aprotinin and 0.2 mmol/L phenylmethyl-163sulfonyl fluoride, using a Polytron homogenizer (PCU-2-164 110; Kinematica GmbH, Steinhofhalde, Switzerland), and 165the residual tissues were stored at -80 °C by flash-freezing in 166 liquid N₂ until use. The homogenates were immediately 167 subjected to the assays described below or stored at -80 °C 168 after liquid N₂ flash-freezing and bathing until use. 169

Lipid peroxide (LPO) concentration was assessed by the thiobarbituric-acid-reactive substance assay of Ohkawa et al. [15] as described by Hashimoto et al. [7,8], and its levels were measured in nanomoles of malondialdehyde per milligram of protein. Malondialdehyde levels were calculated relative to a standard preparation of 1,1,3,3tetraethoxypropane.

The levels of reactive oxygen species (ROS) were 177 determined as described previously [7,8]. Briefly, 50 µl of 178 freshly prepared tissue homogenate was mixed with 4.85 ml 179of 0.1 mol/L potassium phosphate buffer (pH 7.4) and 180 incubated with 2',7'-dichlorofluorescin diacetate (Molecular 181 Probes, Eugene, OR, USA) in methanol at a final 182 concentration of 5 µmol/L for 15 min at 37°C. The dye-183loaded samples were centrifuged at 12,500×g for 10 min at 184 4°C. The pellet was mixed on a vortex at 0°C in 5 ml of 1850.1 mol/L potassium phosphate buffer (pH 7.4) and 186 incubated for 60 min at 37°C. Fluorescence was measured 187 with a Hitachi 850 spectrofluorometer (Hitachi, Tokyo, 188 Japan) at excitation and emission wavelengths of 488 and 189525 nm, respectively. A cuvette holder was maintained at 190 37°C. ROS was quantified from a dichlorofluorescin 191 standard curve in methanol. 192

The fatty acid compositions of plasma and brain tissues were determined by gas chromatography as described previously [7]. Protein concentration was estimated by the method of 196 Lowry et al. [16]. 197

2.6. Gene expression analysis

Gene expression analysis was carried out with the 199GeneChip system (Affymetrix) in accordance with the 200manufacturer's protocol [17]. Briefly, double-stranded DNA 201 was synthesized from 5 µg of total RNA, and the cDNA 202 obtained was used as a template for in vitro transcription. 203Fragmented in vitro transcripts were hybridized overnight 204 with Rat Expression Array 230A (Affymetrix), stained, 205washed and scanned with an Affymetrix GeneArray 206 scanner, where the intensity of the fluorescence for each 207feature was measured. The expression value (average 208 difference) for each gene was determined by calculating 209 the average of differences in intensity (perfect match 210 intensity minus mismatch intensity) between its probe 211 pairs. The image files obtained were analyzed with the 212 Affymetrix data suite system Microarray Suite 5.0 (MAS 2135.0). The expression analysis file created from each sample 214 (chip) was imported into GeneSpring 5.1 (Agilent Tech-215nologies, Inc., Palo Alto, CA) for further data characteriza-216 tion. Briefly, a new experiment was generated after 217importing data from the same organ in which data were 218 normalized by array to the 50th percentile of all measure-219ments on that array. Data filtration based on flags present or 220marginal in at least one of the samples was first performed, 221 and a corresponding gene list based on those flags was 222generated. Lists of same-phenotype genes that were either 223induced or suppressed were created by filtration-on-fold 224 function. Gene Ontology (GO) category analyses were 225performed using the GeneSpring GO browser, which 226calculates hypergeometric P-values to measure statistical 227significance for a specific GO category. 228

2.7. Statistical analysis

Results are expressed as mean±S.E. Behavioral data were 230analyzed by a two-factor (Group and Block) randomized 231 block factorial analysis of variance (ANOVA), and all other 232parameters were analyzed for intergroup differences by one-233 way ANOVA. ANOVA was followed by Bonferroni post hoc 234 comparisons. Correlation was determined by simple regres-235sion analysis. The statistical programs used were GB-STAT 2366.5.4 (Dynamic Microsystems, Inc., SilverSpring, MD, 237USA) and StatView 4.01 (MindVision Software; Abacus 238Concepts, Inc., Berkeley, CA, USA). P<.05 was considered 239statistically significant. 240

3. Results

3.1. Body weight

The final body weights did not differ among the groups 243 (vehicle group: 393 ± 14 ; A β group: 401 ± 12 ; EPA+vehicle 244 group: 385 ± 36 ; EPA+A β group: 404 ± 35 g). The brain slices 245

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Fig. 2. Effects of the chronic administration of EPA (left) and the infusion of A β peptide (1–40) into the rat cerebral ventricle (right) on the number of WMEs evaluated by the radial maze task (see Materials and Methods for details). Left: Control rats (5% gum-arabic-administered rats; *n*=21) and EPA rats (*n*=24). After completing the initial behavior test, each of the two groups (control and EPA groups) was subdivided into two groups and infused with either (right) A β (A β group; *n*=9) or A β solvent (control group; *n*=12): the EPA group was divided into a vehicle-infused EPA group (EPA group; *n*=12) and an A β -infused EPA group (EPA+A β group; *n*=12). The four groups of rats were again behaviorally tested (with six trials) after the implantation of the miniosmotic pump. Each value represents the number of WMEs, presented as mean±S.E.M. in each block of six trials. The statistical significance of differences between the groups was determined by randomized two-factor (Block and Group) ANOVA followed by Bonferroni post hoc test.

prepared 16–17 days after infusion of the A β peptides clearly indicated the deposition of the infused A $\beta_{(1-40)}$ in the corticohippocampal regions (data not shown).



Fig. 3. Effects of the chronic administration of EPA (left) and the infusion of A β peptide (1–40) into the rat cerebral ventricle (right) on the number of RMEs evaluated by the radial maze task (see Materials and Methods for details). Left: Control rats (5% gum-arabic-administered rats; n=21) and EPA rats (n=24). After completing the initial behavior test, each of the two groups (control and EPA groups) was subdivided into two groups and infused with either (right) A β (A β group; n=9) or A β solvent (control group; n=12): the EPA group was divided into a vehicle-infused EPA group (EPA group; n=12) and an A β -infused EPA group (EPA+A β group; n=12). The four groups of rats were again behaviorally tested (with six trials) after the implantation of the miniosmotic pump. Each value represents the number of RMEs, presented as mean±S.E.M. in each block of six trials. The statistical significance of differences between the groups was determined by randomized two-factor (Block and Group) ANOVA followed by followed by Bonferroni post hoc test. Groups without a common letter are significantly different at P<.05. The mole percentages of the unsaturated fatty acids times the number of double bonds in each fatty are presented.

3.2. Effect of EPA on radial maze learning ability

The effect of the chronic administration of EPA on 250working-memory- and reference-memory-related learning 251ability is presented as the mean number of WMEs and RMEs 252for each group, with data averaged over blocks of six trials 253[Figs. 2 (left) and 3 (left), respectively]. Randomized two-254factor (Block and Group) ANOVA revealed a significant 255main effect of both blocks of trials (P=.0005) and groups 256(P<.0001), with a significant Block×Group interaction 257(P=.0018), on the number of WMEs (Fig. 2, left). Similarly, 258ANOVA revealed a significant main effect of both blocks of 259trials (P < .0001) and groups (P < .0001), with a significant 260 Block×Group interaction (P=.0484), on the number of 261RMEs (Fig. 3, left). These results indicate that EPA 262administration improves working-memory- and reference-263memory-related learning ability in young rats. 264

The effect of EPA preadministered to the vehicle and Aβ-265infused groups on working-memory- and reference-memory-266 related learning ability is presented as the mean number of 267WMEs and RMEs for each group, with data averaged over 268six trials [Figs. 2 (right) and 3 (right), respectively]. The 269number of WMEs was significantly higher in the A β group 270(P=.0011) than in the vehicle group (Fig. 2, right), 271 suggesting learning impairment - a well-known character-272istic of AD. The number of WMEs and RMEs was 273significantly lower in the EPA+AB group (WMEs: 274 $P \le 0001$; RMEs: P = .0451) than in the AB group [Figs. 2 275(right) and 3 (right), respectively], indicating that preadmi-276nistration of EPA prevents cognitive deficits caused by the 277infusion of $A\beta$ into the cerebral ventricle of rats. 278

Table 1

Major fatty acid composition of plasma, cerebral cortex and hippocampus in control, EPA, $A\beta$ and EPA+ $A\beta$ rats

	Control	EPA	Αβ	EPA+Aβ
Plasma (%)				
EPA	0.36±0.01°	4.53±0.29 ^a	$0.48{\pm}0.04^{c}$	$3.80{\pm}0.21^{b}$
AA	26.7 ± 0.86^{a}	21.5 ± 0.68^{b}	25.2 ± 0.46^{a}	22.0±1.17 ^b
DHA	$2.47{\pm}0.06^{b}$	$3.53{\pm}0.09^{a}$	$2.46{\pm}0.07^{b}$	$3.50{\pm}0.14^{a}$
DHA/AA	$0.09{\pm}0.00^{ m b}$	$0.17{\pm}0.01^{a}$	$0.10{\pm}0.00^{b}$	$0.16{\pm}0.01^{a}$
USI	185±2.35 ^b	199±1.64 ^a	184±1.34 ^b	198 ± 2.69^{a}
Cortex (%)				
AA	12.0±0.23 ^{a,c}	11.1±0.11 ^b	12.4±0.41 ^a	11.4±0.18 ^{b,c}
EPA	$0.10{\pm}0.00^{\rm b}$	$0.13{\pm}0.01^{a}$	$0.09{\pm}0.00^{\rm b}$	$0.13{\pm}0.01^{a}$
DHA	16.8±0.29 ^{b,c}	$18.1{\pm}0.17^{a}$	16.1±0.44 ^b	17.6±0.29 ^{a,c}
DHA/AA	1.41 ± 0.05^{b}	$1.63{\pm}0.03^{a}$	$1.32{\pm}0.08^{b}$	$1.56{\pm}0.05^{a}$
USI	167±1.15 ^b	173±0.86 ^a	164 ± 1.48^{b}	171 ± 1.20^{a}
Hippocampu	s (%)			
AA	13.4±0.21 ^a	12.5±0.17 ^b	13.3±0.11 ^a	12.7±0.16 ^b
EPA	$0.10{\pm}0.00^{b}$	$0.14{\pm}0.01^{a}$	$0.10{\pm}0.00^{b}$	$0.14{\pm}0.00^{a}$
DHA	$14.5 \pm 0.12^{\circ}$	$15.6{\pm}0.16^{a}$	$15.0{\pm}0.13^{b}$	15.8 ± 0.11^{a}
DHA/AA	$1.09{\pm}0.01^{\circ}$	$1.25{\pm}0.02^{a}$	$1.13{\pm}0.01^{b}$	$1.25{\pm}0.02^{a}$
USI	161±0.91°	$166{\pm}0.97^{a}$	163±0.65 ^b	167 ± 0.69^{a}

Values are expressed as mean \pm S.E.M. and as mole percent of the total fatty acids (n=9-12). Means in a row with superscripts without a common letter differ (P<.05). USL was calculated as a function of the sum of the mole percentages of unsaturated fatty acids times the number of double bonds in each fatty acid.

t1.1

t2.1	Table 2								
	Correlation	between	the	mole	percentages	of	plasma	EPA	and
2.2	corticohippoc	campal EP/	A, Dł	HA and	USI				

t2.3	Plasma	X							
t2.4	EPA (y)	Cortex			Hippocar	npus			
t2.5	()	EPA	DHA	USI	EPA	DHA	USI		
t2.6		0.81 (P<.05)	0.57 (P<.05)	0.62 (P<.05)	0.80 (P<.05)	0.67 (<i>P</i> <.05)	0.60 (P<.05)		
		(1 .05)	(1 .05)	(1 .05)	(4 3.05)	(1 .05)	(1 \.05)		

Results are evaluated with simple regression analysis. *P* values are t2.7 expressed inside the parentheses.

279 3.3. Fatty acid profiles of plasma and brain

The major plasma fatty acid composition in the rat 280plasma, cortex and hippocampus is shown in Table 1. In the 281plasma, the proportion of EPA was significantly higher ----282 and that of arachidonic acid [AA; 20:4(n-6)] was signifi-283 cantly lower ($P \le 05$) — in both EPA and EPA+A β rats than 284in the vehicle and $A\beta$ rats, respectively. The proportion of 285DHA was higher in both EPA and EPA+AB rats than in the 286 vehicle rats. EPA administration brought about a significant 287increase in the plasma DHA/AA molar ratio and USI value in 288 both EPA and EPA+AB rats. 289

290 Chronic administration of EPA significantly enhanced 291 the EPA proportion in both the cortex and the hippocam-292 pus of the EPA and EPA+A β rats. In the hippocampus, the 293 proportion of DHA increased, whereas that of AA 294 decreased significantly, effecting a significant increase in



Fig. 4. The effects of EPA on the LPO (upper; A and B) and ROS (lower; A and B) levels of the cortex (A) and hippocampus (B) of the $A\beta$ rats. The correlation between corticohippocampal DHA/AA molar ratio and LPO levels (C) is presented.

the DHA/AA ratio in both the hippocampus and the 295 cortex. EPA administration brought about a significant 296 increase in the corticohippocampal USI values in both 297 these groups of rats. 298

Highly significant positive correlations were observed 299 between the percent compositions of plasma EPA and EPA, 300 DHA or USI values of both the cortex and the hippocampus 301 (Table 2), indicating that dietary administration of EPA 302 accumulates EPA and DHA in brain tissues. 303

Table 3		t3.1
Effects of chronic administration of EPA on hippocampal gene expre	ession	t3.2
Up-regulated genes		t3.3
1. Signal transduction		t3.4
Cystatin C	2.80	t3.5
GABA _B receptor1	2.04	t3.6
mGluR8	1.97	t3.7
Pyruvate dehydrogenase receptor 8	1.84	t3.8
Regulator of G-protein signaling 4	1.79	t3.9
2. Cell communication		t3.1
Syntaxin 1a	1.79	t3.1
PLD1	1.65	t3.1
Suppressor of K ⁺ transport defect 3	1.73	t3.1
Nucleic acid binding		t3.1
Forkhead box E1 (thyroid transcription factor 2)	2.38	t3.1
Basis helix-loop-helix domain containing class B2	2.02	t3.1
v-maf musculoaponeurotic fibrosarcoma (avian)	1.72	t3.1
oncogene family, protein G		
3. Microtubular dynamics		t3.1
VAPs B and C	1.67	t3.1
4. Enzyme		t3.2
VCP	2.07	t3.2
PLD1	1.65	t3.2
Protein arginine N-methyltransferase 3-like 3	1.60	t3.2
Pyridoxine 5-phosphate oxidase	1.59	t3.2
		t3.2
Down-regulated genes		t3.2
1. Signal transduction		t3.2
Insulin-like growth factor binding protein 2	2.09	t3.2
ATP-binding cassette, subfamily G (WHITE), member 5 (sterolin 1)	1.69	t3.2
GABA- α receptor γ 3 subunit	1.63	t3.3
Hyperpolarization-activated cyclic nucleotide-gate K ⁺ channel 2	1.61	t3.3
2. Cell communication		t3.3
TTR	3.10	t3.3
Integrin α1	2.66	t3.3
Aquaporin 1	1.70	t3.3
Solute carrier family 9, member 1	1.63	t3.3
Transferrin receptor	1.61	t3.3
3. Nucleic acid binding		t3.3
Telomeric repeat binding factor 2	2.17	t3.3
ETS domain transcription factor Pet-1	1.80	t3.4
Myogenin	1.73	t3.4
Gonadotropin-inducible ovarian transcription factor 2	1.56	t3.4
4. Membrane component		t3.4
Ectonucleotide pyrophosphatase/phosphodiesterase 2	2.01	t3.4
5. Enzyme		t3.4
Coagulation factor 5	2.83	t3.4
Glycerol kinase	2.00	t3.4
Triadin 1	1.82	t3.4
Metalloprotease/disintegrin	1.78	t3.4
-	1.77	t3.5

Results are the average of triplicate determination, expressed as the ratio of EPA-administered rats to control rats.

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t3.51

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The levels of both LPO and ROS were significantly 338 337 higher in the cerebral cortex of the A β rats (Fig. 4A) than in the cerebral cortex of the vehicle controls and EPA rats. In 338 the cortex of the EPA+A β rats, both LPO and ROS levels 339 decreased, but decreased significantly only in the latter. In 340 the hippocampus of the A β rats, the level of LPO was 341 significantly higher than that in vehicle rats, but not 342 significantly higher than that in EPA rats (Fig. 4B). The 343 LPO levels were significantly lower in EPA+AB rats than in 344A β rats. The ROS level in the hippocampus was also 345 significantly high in $A\beta$ rats and significantly low in 346EPA+A β rats. Finally, the change in oxidative stress in 347 these two tissues demonstrated a significant negative 348 correlation between LPO levels and corticohippocampal 349 DHA/AA ratios (Fig. 4C). 350

351 *3.4. Effects of EPA administration on hippocampal* 352 *gene expression*

353 EPA administration triggered a substantial change in the expression of both up-regulated and down-regulated genes 354355 (Table 3). The up-regulated genes strongly affected by EPA were signal transduction proteins (cystatin C and GABA_B 356 receptor 1), cell communication protein (syntaxin 1a), 357nucleic acid binding protein (thyroid transcription factor 2) 358and microtubular dynamics [vesicle-associated membrane 359 proteins (VAPs) B and C]. The down-regulated genes 360 strongly affected by EPA were signal transduction protein 361 (ATP-binding cassette protein), cell communication proteins 362 [transthyretin (TTR) and integrin α 1], nucleic acid binding 363 protein (telomeric repeat binding factor 2) and membrane-364 associated ectonucleotidase. 365

366 4. Discussion

The present study provides evidence that not only the oral 367 administration DHA but also the oral administration of EPA 368 protects against memory impairment in the AD model rats 369 infused with amyloid peptide. The protective effect was 370 accompanied by corticohippocampal increases in EPA and 371 DHA, DHA/AA molar ratio and USI values. The 10- to 372 12-fold increase in the proportion of plasma EPA in the EPA 373 and EPA+A β rats (Table 1), as compared with that in the 374 controls, clearly suggests effective intestinal absorption of 375this PUFA after oral administration. Moreover, the correla-376tion between plasma EPA and corticohippocampal DHA or 377 the DHA/AA ratio was highly positive, suggesting that 378 plasma EPA effectively deposits DHA in brain tissues after 379 crossing the blood-brain barrier. 380

We speculate that EPA increases the DHA (and DHA/AA ratio) of the corticohippocampal tissues and, in doing so, exerts beneficial effects on memory formation/protection in EPA or EPA+A β rats. The basis of the speculation is that the level of radiolabeled EPA ([¹⁴C]EPA) detected in the rat brain 1 h after its oral administration to rats decreases time dependently with a concomitant increase in the levels of

[¹⁴C]docosapentaenoic acid and [¹⁴C]DHA, metabolites of 388 EPA [13]. It is also inferred that each PUFA is metabolized 389 after being taken into cerebral endothelial cells and 390 astrocytes (constituent cells of the blood-brain barrier), 391 that it is released from both cells and that DHA is taken into 392neurons as metabolite. This speculation is consistent with the 393 fact that DHA is taken into neurons from the extracellular 394medium after its release from glial cells or the capillary 395 endothelium [14]. Indeed, increased DHA and EPA levels in 396 the plasma raise the fatty acid unsaturation index in rat 397 caudal arteries [18,19]. Moreover, lysophosphatidylcholine 398 may be a transporter of DHA in the blood-brain barrier [20]. 399 Remaining to be explored, however, is whether the 400 transformation of [¹⁴C]DHA from [¹⁴C]EPA occurs in the 401 liver and is then transported to the brain, or whether the 402 conversion occurs solely and/or partly in brain tissues. 403Regardless of where EPA is transformed into DHA, it is 404 evident from our study that EPA was converted into DHA 405and exerted its effects on memory functions. 406

The brain has an intrinsic capacity to retain its DHA; 407 however, if DHA is depleted, huge differences in brain 408 functions occur [21]. Thus, a change in brain DHA level 409might be related to behavioral impairments [22]. We 410 reported that a small increase (in mol%) in DHA content 411 contributed significantly to limiting memory deficits in 412 DHA-deficient rats [8]. Thus, a small but highly significant 413increase in corticohippocampal DHA composition (8-414 9 mol%) in the EPA/EPA+AB rats after EPA administration 415is consistent with our previous reports [8]. EPA-mediated 416 actions directly or indirectly could involve effects on 417 antioxidative status, amyloid processing, apoptosis, expres-418 sion of a host of proteins, membrane lipid (disorder) 419fluidity and exocytosis. An increased DHA/AA ratio is 420 associated with increased memory-related learning ability 421 in young [5], aged normal [6] and AD model rats [7,8], 422 with a concurrent decrease in brain LPO levels. Consistent 423 with our previous investigations [7,8], the DHA/AA ratios 424 correlated negatively with the repression of lipid peroxida-425tion (Fig. 4C). The mechanism by which this correlation 426affects memory enhancement and amyloid burden is not 427 clear yet. Free-radical theory of AD pathology involves 428 amyloid-induced oxidative stress [23]. Increasing levels of 429DHA in the cortex of aged rats significantly increase 430 antioxidative enzymes, including catalase, glutathione 431 peroxidase and reduced glutathione [24]. We have 432 hypothesized that the DHA/AA ratio acts as an indicator 433 of antioxidants indirectly by inhibiting the level of AA in 434 the neuronal plasma membrane [7]. An increase in the 435 DHA/AA ratio thus, at least partially, protects the 436 corticohippocampal regions from oxidative insult and 437 provides protection against the impairment of memory 438 in A β -infused rats. DHA inhibits the accretion of A β 439peptide (1–40) in detergent insoluble neuronal membrane 440 domains of the cerebral cortex [25] and of AB-induced 441 apoptosis-like neuronal cell death [7]. Thus, the finding of 442 EPA-administration-induced protection against memory 443

impairment, with concurrent DHA accretion in the brain, is 444 445in line with our studies [7,8] and those of others [10,26]. An alternative mechanism of EPA-induced amelioration 446 of memory may be as follows: we have previously reported 447 448 449450451452453454455456457458

that DHA increases the expression of the Fos protein, the immediate early gene *c-fos* (which acts as a transcription factor and as a functional marker of neuronal activity) of the rat CA1 hippocampus [27]. n-3 PUFAs induce the expression of a host of genes that control synaptic plasticity and underlying signal transduction mechanism(s) [11]. Using microarray analysis, we also examined whether the administration of EPA induces the expression of genes, and whether their function is correlated with EPA-induced memory protection in the AD model rats. The gene chip data are correlative and appear to be the only data from EPAversus-control rats in the present experimental scenario. 459However, if such data had been obtained also from AB and 460 461 EPA+AB rats, then gene expression would have been conferred to the neuroprotective actions of EPA in the 462 EPA+AB rats. Consistent with this possibility, we found that 463 EPA up-regulated 16 genes and down-regulated 25 genes 464 (Table 3). The γ -aminobutyric acid (GABA) receptor 1 465466 (GABA_B) and the metabotropic glutamate receptor (mGluR) 1 were expressed twice, as compared with those of control 467rats. The GABA_B receptor interacts with mGluR1-mediated 468 excitatory transmission [28], suggesting that EPA adminis-469 tration might contribute to a mechanism of regulatory 470synaptic plasticity. Several other genes that participate in 471 cell communication were also overexpressed. For example, 472 the expression of syntaxin 1a and VAPs B and C increased 473concurrently (Table 3). Syntaxin 1a forms a complex with 474 VAP [29] in order to release neurotransmitters and regulates 475their transporter [30]. These activities further support the 476 implication of the n-3 PUFA-induced increase in neuro-477 transmitter release and synaptic plasticity. In addition, the 478expression of the *cystatin* C gene, a lysosomal cysteine 479protease inhibitor [31], increases upon EPA administration. 480 Lysosomal proteases (including cathepsins B and D) are up-481 regulated in the AD brain [32], and cystatin C relates to 482neurogenesis in neural stem cells [33]. In this regard, we 483 have recently shown that DHA significantly enhances 484 neurogenesis both in vivo and in vitro [34]. Phospholipase 485D1 (PLD1) regulates both the release of secretory vesicles 486 from the trans-Golgi network (TGN) [35] and exocytosis 487 [36]. Amyloid proteolysis and subsequent trafficking from 488 the TGN to the cell surface are impaired in AD. Up-489regulation of PLD1 in AD rescues impaired APP trafficking 490 from TGN to the membrane surface [37]. In our study, 491 expression of the valosin-containing protein (VCP) 492 increased concomitantly in EPA-fed rats. VCP senses and 493 pulls abnormal protein accumulation in the cell, and pulls 494 from the endoplasmic reticulum and nucleus [38]. DHA-495induced clearance of AB from neuronal membranes is 496 suggested to be mediated by exocytosis — a process that 497 may be facilitated by the DHA-induced increase in 498membrane fluidity [24,39,40]. Consistently, EPA-induced 499

memory augmentation is attributed to neurotransmitter 500 (noradrenaline) release [10], as facilitated by an increase in 501 membrane disorder (fluidity) [41]. A DHA-induced increase 502in memory-related performance is accompanied by increased 503levels of acetylcholine [42]. Therefore, all these effects are 504speculated to be in concert with the EPA-induced increase in 505memory protection of the EPA+AB rats. Our microarray data 506 also suggest that EPA suppressed the genes involved in 507signal transduction, cell communication and/or membrane-508bound components to a remarkable extent (Table 3). The 509contributions of EPA to the repression of these genes have 510remained largely unrecognized. For example, TTR is a 511thyroid hormone transporter that is secreted by the liver in 512plasma and by choroid epithelial plexus in cerebrospinal 513fluid. The TTR gene was suppressed in our feeding 514paradigm. To date, the effects of n-3 PUFAs on the 515 expression of TTR are inconsistent and conflicting. Short-516term administration of n-3 fatty acids from fish oil for 517 1 month increases TTR expression in 2-year-old rats [43]. In 518contrast, TTR gene expression is unchanged in mice fed a 519 high-DHA diet [44]. Our result is, however, consistent with 520the report of Tanabe et al. [27] wherein the rats were fed n-3521PUFA containing 8% fish oil diet. In our study, the young 522rats were fed 99% purified EPA at a dose of 300 mg/kg body 523weight for 12 weeks. The discrepancies may thus relate to 524the age of the rats, the feeding duration, the fatty acid 525composition of the supplemented oil and the chow diet itself. 526Also, the debate on whether TTR is expressed in the 527hippocampus and/or whether the level found in brain tissues 528is just an effect of contamination from the plexus epithelium 529must be resolved [45]. Integrin $\alpha 1$ expression decreased 530consistently on EPA feeding. Inhibition of this cell-531communicating protein by echistatin or antibodies protects 532against AB-induced degeneration of neurons in vitro [46]. 533 Because the gene chip data are unsubstantiated, with no 534validation at either the mRNA level (by RT-PCR) or the 535protein level, the correlation of EPA-induced memory 536amelioration with other altered genes has remained without 537further predication. The induction of gene expression thus 538 needs to be confirmed by real-time RT-PCR. 539

To summarize, EPA protects against AB-peptide-induced 540memory deficit in AD model rats after its transformation into 541 DHA. This is accompanied by the accumulation of DHA 542and/or an increase in the DHA/AA ratio in the corticohippo-543campal tissues, with a corresponding decrease in oxidative 544stress and an increase in the expression of synaptic-545plasticity-related proteins. Nonetheless, further studies are 546needed for additional data on EPA. 547

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