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Original paper

Actions of mesenchymal stem cell secretome on microglia amyloid-β uptake

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Abstract

In the brains of elderly dementia patients, important proteins (*i.e.*, amyloid β [A β], tau, and α -synuclein) undergo posttranslational modification, polymerization, aggregation, and deposition in tissues. Glycative stress, a non-physiological factor inducing these phenomena, is a state of excess aldehydes, and is induced by hyperglycemia, a high-fat diet, and alcohol consumption. In the lipid-rich brain, concurrently with carbohydrate-derived aldehydes, FA-derived aldehydes generated by the oxidation of FAs modify brain proteins like a double punch. Our previous reports have shown that glycated A β can become resistant to microglia phagocytosis and may interfere with clearance. In this study, we created an A β phagocytosis evaluation model using microglia-derived BV2 cells and evaluated the effects of mesenchymal stem cell secretome (SCST). SCST contains various growth factors and extracelluar vesicles (EVs), and it has been reported that nasal administration improves symptoms in elderly dementia patients. Present results showed that SCST promotes A β phagocytosis in a dosedependent manner. These findings suggest that SCST may activate microglia and contribute to the homeostasis of A β clearance. We speculate that it may also be important from the perspective of gliaprotection.

KEY WORDS: microglia, amyloid β , phagocytosis, dementia, stem cell secretome, gliaprotection

Introduction

Various important proteins (*i.e.*, amyloid β , tau, and α synuclein) in the brains of elderly dementia patients undergo post-translational modification, polymerization, aggregation, and tissue deposition. Although the physiological role of amyloid beta (A β) is unknown, it plays an important role in the pathogenesis and progression of Alzheimer's disease (AD). It is said that A β polymerization increases neurotoxicity, is difficult to decompose, and deposits in the brain, reducing A β clearance. We focus on glycative stress (GS) as a risk factor for AD progression. GS is a state of aldehyde excess, caused by hyperglycemia, a high-fat diet, and excessive alcohol intake. In the lipid-rich brain, concurrently with carbohydrate-derived aldehydes, fatty acid (FA)-derived aldehydes generated by the oxidation of FAs induce a double punch of glycative modification of brain proteins.

In patients with diabetes, a typical disease with severe GS, various types of aldehydes including methylglyoxal (MGO) and acrolein (Acro) increase. Our previous studies have focused on A β phagocytosis by microglia, which is involved in A β clearance, and have examined the effects of GS. Glycated A β is generated by MGO or Acro treatment. Fluorescence-labeled A β (TAMRA-A β) and primary cultured mouse-derived microglia cells (Cosmo Bio Co., Ltd., Tokyo, Japan) were used in the experiments. Experiments using these cells have shown that microglia phagocytose A β , while in contrast phagocytosis of glycated A β is markedly reduced¹). When $A\beta$ was added to primary cultured microglia cells, there was a tendency for activation, proliferation, and natural death to decrease compared to when $A\beta$ was not added, so the possibility of neurotrophic gliaprotection should be seriously considered. Similar comments have been made in

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other papers²⁾. In light of the above, it is considered important in preventive medicine to "improve A β clearance and prevent the progression of AD" by lifestyle habits such as preventing glycation modification of A β through GS care and improving "quality of sleep" rather than completely eliminating A β .

Given this background, our laboratory, as stated in the Grant-in-Aid for Scientific Research application, began searching for components that promote or inhibit microglia $A\beta$ phagocytosis³⁾. As a candidate substance, we are focusing on extracellular vesicles (EVs) derived from mutualistic bacteria in the gut microbiota.

During this period, Morita et al. reported that nasal administration of mesenchymal stem cell-derived secretome (SCST) improves the Hasegawa's dementia score in elderly patients with dementia⁴⁾. There have been other similar reports^{5.7)}. We hypothesized that one of the mechanisms underlying the improvement of dementia symptoms is that SCST promotes microglia A β phagocytosis. To verify this hypothesis, we examined the effect of SCST on phagocytosis in patients with dementia in an A β phagocytosis, A β clearance may improve. We hope that if factors that improve A β clearance can be used, we will be able to obtain a useful means for preventing the progression and exacerbation of AD.

Methods

From primary cultured microglia cells to cell lines

The established cell lines used in microglia experiments to date have retained macrophage-like properties, and are said to differ significantly from in vivo microglia in terms of their response to inflammation and neuroprotection. Primary cultured microglia are prepared from the brain immediately after birth, and are grown in a mixed culture system that mainly contains astrocytes and neurons in addition to microglia, thereby maintaining the original functions of microglia⁸⁾. In previous studies, a rat primary microglia culture kit (Cosmo Bio) was used to verify the function of microglia, which is to recognize and eliminate Aβ⁹. Primary cultured cells are not suitable for an experimental model in which the efficacy of a relatively large number of samples is repeatedly evaluated due to large lot-to-lot differences. Therefore, the mouse microglia cell line BV2 (Code No.: 305156-Academic, Cell Lines Service GmbH, Germany) was used in this study.

Microglia cell line BV2

BV2 cells are a type of microglia cell line derived from C57BL/6 mice that is widely used in animal experiments^{10,11}. The cells were immortalized and established using a J2 retrovirus carrying the v-raf and v-myc oncogenes. BV2 cells express env gp70 antigen on their surface, which is one of the important features of BV2 cells, along with expression of nuclear v-myc and cytoplasmic v-RAF oncogenes. The env gp70 antigen contributes to the cell function in immune responses and inflammation in the brain.

BV2 cells were cultured in RPMI 1640 Medium (Thermo Fisher Scientific Inc.) supplemented with 10% fatal bovine serum (FBS) and Penicillin-Streptomycin (Fujifilm Wako Pure Chemicals) at 100 IU/mL, 0.1 mg/mL. For fluorescence observation, phenol red-free medium was RPMI 1640 medium, no phenol red (Thermo Fisher Scientific Inc.) supplemented with 10% of FBS, penicillin-streptomycin (Fujifilm Wako Pure Chemicals) at 100 IU/mL, 0.1 mg/mL.

Synthesis of fluorescence-labeled $A\beta$ (TAMRA- $A\beta$)

The TAMRA-A β used in the previous report¹) was fluorescently labeled TAMRA-A β (Cosmo Bio). This reagent consists of A β -(1-42) labeled with red fluorescent carboxytetramethylrhodamine (TAMRA) via a PEG spacer²).

The A β -(1-42) peptide is a difficult reagent to handle. Its amphipathic sequence and strong self-aggregation tendency complicate characterization of both structure and function. Large lot-to-lot variations affect both aggregation behavior and biological activity. Dahlgren KN et al. recently took steps to minimize this variation by removing what they call "structural history"²). Structural history refers to secondary, tertiary, or quaternary structures that can act as templates or seeds and direct a large fraction of a peptide solution into a specific aggregation pathway. These structural seeds are not detected in synthetic products using traditional quality control methods that focus primarily on chemical purity and not on structural heterogeneity.

However, this A β -(1-42) peptide, which has the property of good stability, was discontinued in 2022, and the stock was depleted. After that, we tried to use various commercially available fluorescence-labeled A β , but we were unable to obtain labeled A β suitable for the experiment due to unstable properties, poor solubility, and poor compatibility with primary microglia. Therefore, we decided to synthesize new A β . The TAMRA-labeled mouse A β used in this experiment was synthesized by the Sapporo Division of Cosmo Bio. The sequence was 5-TAMRA-XDAEFGHDSGFEVRHQKL VFFAEDVGSNKGAIIGLMVGGVVIA X:AEEAc (PEG), molecular weight 4975.57.

Verification of $A\beta$ phagocytosis

Aβ pretreatment

After dissolving mouse A β in trifluoroacetic acid (TFA) (Nacalai Tesque) to 1 mg/mL, 100 µL/tube of the dissolved solution was divided into two microtubes, diluted 10-fold with distilled water, and dried in a centrifuge evaporator. To loosen the aggregation of A β , 100 µL/tube of 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP) (Kanto Chemical Co.,Inc.) was added to A β after drying in a centrifugal evaporator, and the solution was re-dissolved. The cells were again dried in a centrifugal evaporator and stored at -20 °C until use. The day before cell addition, the dried A β was dissolved in 40 µL of dimethyl sulfoxide (DMSO) (Merck KGaA) (2.5 mg/mL) and further diluted 25-fold by adding 960µL of medium (final A β concentration: 100 µg/mL = 20 µM, DMSO concentration: 4%). A β diluted in medium was incubated at 4°C for 24 hours to oligomerize A β .

Experimental procedure: Verification of Aß phagocytosis

BV2 cells, which had been cultured until sufficient volume was reached by passaging, were seeded in 96-well plates at a density of 5×10^3 cells/well. The day after seeding, supplemental medium was prepared at the following concentrations and added at 100 µL/well.

Negative control	Aβ: None	DMSO: 0.2%					
Positive control	Αβ: 1μΜ	DMSO: 0.2%					
Yeast EV							
×100	Αβ: 1μΜ	DMSO: 0.2%	EV: 1%				
×1,000	Αβ: 1μΜ	DMSO: 0.2%	EV: 0.1%				
×10,000	Αβ: 1μΜ	DMSO: 0.2%	EV: 0.01%				
SCST-A							
$\times 2$	Αβ: 1μΜ	DMSO: 0.2%	HGF: 50 ng/mL				
×20	Αβ: 1μΜ	DMSO: 0.2%	HGF: 5 ng/mL				
×200	Αβ: 1μΜ	DMSO: 0.2%	HGF: 0.5 ng/mL				
SCST-B			-				
$\times 2$	Αβ: 1μΜ	DMSO: 0.2%	HGF: 5 ng/mL				
× 5	Αβ: 1μΜ	DMSO: 0.2%	HGF: 2 ng/mL				
×20	Αβ: 1μΜ	DMSO: 0.2%	HGF: 0.5 ng/mL				
(Note that HGF is an estimated concentration							

At 24 hours after addition, the supernatant was removed, washed once with HBSS (–), and 100 μ L of phenol red-free medium was added. The cells were then observed with an all-in-one microscope (BZX-710, Keyence). From the TAMRA-A β fluorescence and phase contrast images, image analysis was performed to calculate the TAMRA fluorescence intensity per image and the total area value of cells.

Formula:

TAMRA fluorescence intensity / total area value of cells

The TAMRA fluorescence intensity was corrected for the total area value of the cells, and the control ratio of each group to the negative control group (% NC) and the control ratio of each group to the A β -supplemented group (positive control) (% A β) were calculated and graphed.

Stem cell-derived secretome (SCST)

Two types of cell supernatants were used in this study. The first was the same SCST used in the previous report ⁴), and freeze-dried mesenchymal SCST (SCST-A) was used (Nature Bionics, Inc., Tokyo). When dissolved in 1 mL, one vial contains approximately 1×10^8 exosomes and 1.2×10^5 pg/mL of hepatocellular growth factor (HGF), one of the main growth factors, as well as many other types of growth factors, nutritional factors, cytokines, etc. In addition to HGF, the product is manufactured to meet the respective index amounts for several representative components, so that the active ingredients are almost the same in amount and there is little difference between lots.

The donors of the stem cells were healthy Japanese women in their 20s and 30s, and a completely serum-free medium was used without insulin or antibiotics and without animal or human serum to avoid the risk of viral infection. Unnecessary components such as ammonia, which are produced in large quantities during the cultivation process, are removed to below the detection limit, and active components such as various growth factors and exosomes are concentrated.

The second SCST sample (SCST B) was provided by a medical institution in Japan. They say it contains an estimated HGF concentration of 5 ng/mL. It also contains exosomes and growth factors, but the amount contained is unknown.

Yeast-derived extracellular vesicles (EV) were used for comparison. Yeast-derived EV (Yeast EV) used was YSEV-R3 (Cosmo Bio).

Verification of gene expression: Microarray method

BV2 cells were passaged to reach sufficient volume and seeded into 100-mm dishes at a density of about 1.6×10^6 cells per dish. The day after seeding, medium for addition was prepared to the following concentrations of A β and DMSO and added at 10 mL/dish by changing the medium. 24 hours after addition, the supernatant was removed, the cells were lysed, and total RNA was extracted. After extraction, microarray analysis was performed by Macrogen Japan (Tokyo).

Negative control	Aβ: None	DMSO: 0.3%
Positive control	Aβ: 1 μM	DMSO: 0.3%

Results

Microglia $A\beta$ phagocytosis

Figures 1 and 2 show the culture status of microgliaderived BV2 cells as fluorescence and phase contrast microscopic images.

TAMRA fluorescence was not detectable in the A β non-added group (negative control), whereas fluorescence was detectable in the A β added group (positive control). It was difficult to evaluate by image observation alone whether the addition of yeast EV or SCST changed the microglia A β phagocytosis.

In *Tables 1* and 2, the fluorescence intensity of each was measured and used as an index of microglia $A\beta$ phagocytosis. Each group consisted of n = 4. Since differences in measurement conditions caused differences in the measured values of the positive control, negative and positive controls were established for each plate.

Figure 3 shows the ratio of fluorescence intensity to the positive control. Only the mesenchymal SCST-A used in the previous clinical study⁴⁾ enhanced microglia A β phagocytosis in a dose-dependent manner.

Microarray analysis

To verify whether $A\beta$ is actually taken up by microglia and exerts its effects at the molecular level, we analyzed the changes in gene expression using microarrays. The results of the Gene Ontology (GO) analysis are classified into three categories: biological process (*Fig. 4-a*), cellular component (*Fig. 4-b*), and molecular function (*Fig. 4-c*). In the biological process, gene expression changes were observed in the regulation of cellular processes, regulation of biological processes, biological regulation, cellular response to stimuli, and signal transduction. In the cellular component, gene expression changes were observed in the anatomical entity of cells. In the molecular function, gene expression changes were observed in signal receptor activity, signal transduction system, transmembrane receptor activity, and G proteincoupled receptor activity.



EV, extracellular vesicle.



EV, extracellular vesicle. SCST, mesenchymal stem cell-derived secretome.

Group (Plate 1)	FI (Integrated)	Area (Integrated)	FI/Area (Corrected)	Average	SEM	% NC	% PC
	5.662	5,886,304	9.62.E-04	2.57.E-04	2.03.E-04	100%	-
Negative control $(NC) \land \beta(.)$	108	5,173,551	2.09.E-05				
Positive control (PC) $A\beta(+)$	135	6,504,722	2.08.E-05				
	162	6,253,515	2.59.E-05				
	604,653	4,816,245	1.26.E-01	1.51.E-01		58,529%	100%
	720,560	5,875,886	1.23.E-01		1.33.E-02		
	1,163,465	6,7042,90	1.74.E-01				
Positive control (PC) $A\beta$ (+)	1,332,503	7,369,924	1.81.E-01				
······································	978,860	6,210,947	1.58.E-01				
	756,885	6,543,283	1.16.E-01				
	871,041	6,453,243	1.35.E-01				
	395,331	4,434,738	8.91.E-02	1.20.E-01 1		46,600%	80%
	746,704	6,398,303	1.17.E-01		1.03.E-02		
Yeast EV x100 A β (+)	820,308	6,409,137	1.28.E-01				
	915,831	6,278,260	1.46.E-01				
Yeast EV×1,000 A β (+)	295,519	4,741,174	6.23.E-02	1.03.E-01 1.37.E-02		39,831%	68%
	471,174	3,434,813	1.37.E-01		1.37.E-02		
	494,521	4,304,150	1.15.E-01				
	422,725	4,420,153	9.56.E-02				

Table 1. Results of fluorescence intensity: Aβ phagocytosis index (Plate 1).

FI, fluorescence intensity; SEM, standard error mean; EV, extracellular vesicle.

Table 2. Results	of fluorescence	intensity: Aß	phagocytosis	index (Plate 2)
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Group (Plate 2)	FI (Integrated)	Area (Integrated)	FI/Area (Corrected)	Average	SEM	% NC	% PC
	5	5,825,113	8.58.E-07				
Nagative control $AB()$	28	7,062,462	3.96.E-06	2 20 E 04	1.83.E-04	% NC 100% 27,333% 16,198% 131,519% 70,917% 49,771% 16,178% 26,215%	-
Negative control Ap (-)	132	6,300,474	2.10.E-05	2.20.E-04			
	5,839	6,845,136	8.53.E-04				
	264,613	6,015,738	4.40.E-02			% NC 04 100% 02 27,333% 03 16,198% 03 16,198% 03 70,917% 03 49,771% 03 16,178% 03 26,215% 02 50,232%	
Positive control $\Delta\beta(\pm)$	264,740	5,136,753	5.15.E-02	6 00 E 02	1.01.E-02		100%
r ositive control Ap(+)	259,160	5,192,431	4.99.E-02	0.00.E-02			
	554,611	5,852,807	9.48.E-02				
	163,736	6,176,329	2.65.E-02			% NC 04 100% 02 27,333% 03 16,198% 02 131,519% 03 70,917% 03 49,771% 03 16,178% 03 26,215% 02 50,232%	
	196,163	5,547,236	3.54.E-02				59%
Yeast EV ×10,000 A β (+)	168,509	5,372,446	3.14.E-02				
	321,880	6,555,047	4.91.E-02	3.56.E-02	4.21.E-03		
	1,103,828	5,097,519	2.17.E-01				
	875,578	4,886,988	1.79.E-01				
	1,525,473	5,143,644	86,988 1.79.E-01 43,644 2.97.E-01 81,199 4.59.E-01 44,514 2.37.E-01 01,226 1.99.E-01 29,043 2.61.E-01 60,483 1.56.E-01				
	1,277,428	2,781,199	4.59.E-01		5.04.E-02	131,519%	481%
	1,054,059	4,444,514	2.37.E-01	2.89.E-01			
SCS1-A $\times 2$ A β (+)	913,478	4,601,226	1.99.E-01				
SCST-A ×2 A β (+)	1,285,373	4,929,043	2.61.E-01				
SCST-A ×20 A β (+)	647,831	4,160,483	1.56.E-01	156 5 01	3.EM % NC 1.83.E-04 100% 1.01.E-02 27,333% 4.21.E-03 16,198% 5.04.E-02 131,519% 3.61.E-03 70,917% 1.66.E-03 49,771% 5.40.E-03 16,178% 1.27.E-03 26,215% 1.40.E-02 50,232%	70,917%	259%
	854,001	5,840,659	1.46.E-01				
	974,431	5,850,179	1.67.E-01	1.30.E-01			
	895,619	5,788,828	1.55.E-01				
	491,261	4,412,781	1.11.E-01	4 2 6.00.E-02 1.01.E-02 27,333% 2 3.56.E-02 4.21.E-03 16,198% 1 2.89.E-01 5.04.E-02 131,519% 1 1.56.E-01 3.61.E-03 70,917% 1 1.09.E-01 1.66.E-03 49,771% 1 1.09.E-01 1.66.E-03 16,178% 2 3.55.E-02 5.40.E-03 16,178% 2 5.76.E-02 1.27.E-03 26,215% 2 1.10.E-01 1.40.E-02 50,232%			
SCST A V200 AR(1)	620,828	5,920,794	1.05.E-01		1.66.E-03	49,771%	182%
$SCST-A \times 200$ Ap (+)	696,461	6,465,712	1.08.E-01	1.09.E-01			
	721,674	6,359,165	1.13.E-01				
	137,547	4,530,323	3.04.E-02		5.40.E-03	16,178%	59%
$SCST_B \times 2 A\beta(+)$	164,859	5,775,365	2.85.E-02	3.55.E-02			
$5C51-D^2$ Ap (1)	161,843	5,569,193	2.91.E-02				
	281,414	5,191,789	5.42.E-02				
	262,819	4,586,097	5.73.E-02	5.76.E-02 1.2		26,215%	96%
SOST DUS AQ(1)	332,899	6,202,677	5.37.E-02		1.27.E-03		
$SCSI-B \times 5 A\beta(+)$	371,320	6,127,799	6.06.E-02				
$3C31-B \times 5$ Ap (+)	381,500	6,488,118	5.88.E-02				
	448,949	5,450,273	8.24.E-02			70,917% 49,771% 16,178% 26,215% 50,232%	184%
	327,562	3,844,944	8.52.E-02	1 10 E 01	1 40 E 02		
$SUSI-B \times 20$ AB (+)	565,440	3,778,175	1.50.E-01	1.10.E-01	1.40.E-02		
	417,598	3,362,082	1.24.E-01				

FI, fluorescence intensity; SEM, standard error mean; EV, extracellular vesicle; SCST, mesenchymal stem cell-derived secretome.

Fig. 3. Microglia Aβ phagocytosis potential: Comparison with positive control. Results are expresses as % change of fluorescence intensity derived from TMRA-Aβ uptaken by microglia VB2 cell (data from Table 1 & 2). EV, extracellular vesicle; SCST,

mesenchymal stem cell-derived secretome.

Fig. 4-a.

Fig. 4. Microarray GO functional analysis.

a) Biological process. b) Celluar component. c) Molecular function. Results are expressed as Gene Ratio (intersection-size/query-size.). GO, gene ontology.

Discussion

Glial cells (astrocytes, oligodendrocytes, and microglia) play important roles in supporting the activity of the central nervous system (CNS), maintaining the tissue environment, providing nutrition, and clearing A_β. Damage to glial cells also has a negative effect on the CNS¹²). Factors that cause damage to glial cells include inflammation, oxidative stress (OS), and GS¹³⁾. Although the term "metabolic stress" has often been used, in this paper we refer to it as GS, as excess aldehydes are involved in most cases. Reducing these factors and aiming for gliaprotection is extremely important for maintaining homeostasis of the CNS.

Many AB removal therapies have been attempted to date, however, there have been many adverse events and no successful cases. The current situation is that therapeutic drugs are being developed with the main pharmacological effects being the inhibition of A β production mechanism by controlling secretase activity, the inhibition of amyloid formation by inhibiting $A\beta$ aggregation, and the inhibition of amyloid deposition to promote $A\beta$ removal. It has been shown that the A β clearance rate is significantly decreased in patients with sporadic AD14, and the complete picture of the metabolic pathway of $A\beta$ degradation is awaited.

According to Dahlgren KN et al., early in vitro studies suggested that A\beta-induced neurotoxicity required the peptide to adopt a fibrillar aggregated state, and that low doses of non-aggregated peptides were indeed neurotrophic²⁾. Their validation showed that oligometric A β -(1-42) significantly reduced neuronal viability, whereas non-aggregated peptides had less effect on viability. Non-aggregated AB showed a biphasic response, exerting neurotrophic effects at low concentrations (1-100 µM) and neurotoxicity at higher concentrations (1–15 μ M). Non-aggregated A β -(1-40) had no significant effect on neuronal viability even at 20 µM, and at concentrations below 1 µM appeared to be as neurotrophic as

non-aggregated $A\beta$ -(1-42) preparations. Supporting the CNS by maintaining the health of microglia's $A\beta$ phagocytosis, restoring its reduced function, and promoting its activity may also be effective in preventing the progression of dementia (especially AD), and new treatment strategies are expected.¹⁵⁻²¹). Metformin^{16,22}, curcumin^{23,24}, and resveratrol^{23,25} not only have anti-oxidation functions, but also have anti-glycation effects that inhibit the formation of advanced glycation products (AGEs). These findings suggest that GS care is important in gliaprotection as well.

On the other hand, oxidation and lipid loading induce a decline in microglia function²⁶. Restoration of mitochondrial function also restores A β phagocytosis²⁷. Oxidation and lipid loading induce the oxidation of FAs, enhancing the formation of FA-derived aldehydes, leading to an excess state. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) and aldehyde dehydrogenase (ALDH) consume NAD (nicotinamide adenine dinucleotide) in the process of metabolizing aldehydes, and NAD is also consumed in the process of β -oxidation enhanced by lipid loading. As a result, NAD becomes insufficient in the TCA cycle in mitochondria, subsequently ATP production decreases, and ultimately the cellular function of microglia is weakened.

Deposition of aldehyde-modified proteins such as MGO and Acro has been observed in senile plaques (amyloid plaques) by immunohistochemistry²⁸⁻³⁸, the presence of glycated A β is strongly suspected. It has been pointed out that urinary aldehyde-modified proteins may be an indicator of dementia³⁹.

In the lipid-rich brain, important proteins are vulnerable to a double whammy attack from not only carbohydrate-derived aldehydes (*i.e.*, GO [glyoxal], MGO, 3DG [3-deoxyglucosone], GA [glyceraldehyde]) but also FA-derived aldehydes (*i.e.*, MGO, Acro, MDA [malondialdehyde], etc.)⁴⁰⁻⁴²) produced by the oxidation of FAs¹³.

In our previous study, we reported that MGO and Acromodified A β are difficult to phagocytose by microglia¹). This result suggests that aldehyde modification of A β due to GS may be the cause of reduced A β clearance. In the subsequent clinical study⁴), we reported that nasal administration of SCST may lead to improvement of symptoms in dementia patients. In order to explore the mechanism of improvement, this study examined the effect on microglia A β phagocytosis, which is one of the indicators of A β clearance.

The SCST contains various growth factors and EVs. Among them, HGF is a physiologically active protein that is responsible for regeneration of the liver and kidney and protection of neural tissue. It is known as the most powerful growth factor for hepatocytes and is also produced in organs other than the liver ⁴³. HGF strongly promotes the proliferation of primary cultured mature rat hepatocytes at concentrations of 1 ng/mL or more⁴⁴⁾. The HGF concentration in human serum is high in patients with fulminant hepatitis, averaging approximately 10 ng/mL. In patients with acute hepatitis, liver cirrhosis, and liver cancer, the levels are 2-3 times higher than in healthy adults (approximately 0.2 ng/mL), but rarely exceed 1 ng/mL. The estimated HGF concentration in the culture medium of SCST-A×200 (HGF: 0.5 ng/mL) is 2 to 3 times that of healthy adults. The results of this experiment showed that the SCST administration increases microglia Aß phagocytosis in a dose-dependent manner. Nasal SCST administration may have a similar effect and contribute to

improving $A\beta$ clearance or maintaining homeostasis.

Of course, it is expected that the growth factors contained in SCST contribute to the protection and activation of neurons and glial cells. Since neurons in the hippocampal dentate gyrus, which are deeply involved in memory function, maintain their division and proliferation functions even in the elderly, it is expected that activation of these cell functions will greatly contribute to the improvement of dementia symptoms. This test did not clarify to what extent the active ingredient is transferred into the brain parenchyma by nasal administration, that can be a future topic of research.

Molecular impacts of $A\beta$ on microglia

To verify whether $A\beta$ is actually taken up by microglia and has an impact at the molecular level, we analyzed the changes in gene expression using microarrays. Genes with significant changes in expression were extracted in the categories of biological process, cellular component, and molecular function. It was shown that the $A\beta$ administration had an impact on microglia gene expression. In this analysis, $A\beta$ oligomers were used, and different results may be obtained with non-aggregated $A\beta$. Later, we plan to analyze the changes in the expression state of individual genes over time to see whether $A\beta$ is involved in gliaprotection. It is an extremely interesting question which genes are affected by SCST.

Microglia $A\beta$ phagocytosis model

In this study, we have established an evaluation model for microglia $A\beta$ phagocytosis. We placed emphasis on reproducibility so that the efficacy of various test substances can be compared and evaluated also at other facilities.

When preparing the model, we considered whether to use primary cultured cells or lineage cultured cell lines as microglia. The former is assumed to retain the characteristics of microglia at a high level, but they need to be collected from animals for each experiment, and there is a possibility of individual differences and differences due to techniques. The latter may have lost some of the microglia characteristics, but it expectedly has higher reproducibility. The latter was selected as a model that requires multiple experiments over a long period of time and analyzes multiple comparative factors.

We selected the $A\beta$ to be used. This was because TAMRA- $A\beta$, which was used in the previous report ^D, was discontinued. We examined the addition of fluorescent labels, changes in properties due to aldehyde modification and the creation of oligo- and fibril bodies, and the compatibility and cytotoxicity between primary cultured microglia and culture strains. As a result, we used newly synthesized $A\beta$. The details of this research will be reported separately.

Next, Polaric (Cosmo Bio) and TAMRA were compared as fluorescent dyes for labeling $A\beta$. After fluorescent labeling of $A\beta$, oligomeric and fibril bodies were prepared as previously reported²⁾ and the fluorescence intensity was evaluated. TAMRA- $A\beta$ was selected, which showed little change in fluorescence intensity after phagocytosis.

Regarding measurement methods, a method using a confocal laser microscope and an all-in-one fluorescence microscope were compared. The former is suitable for taking beautiful photographs, but requires separate image analysis using a separate device, making the process complicated.

With the latter, the process of taking photographs under fixed conditions, saving the photo files, and analyzing the images can all be done with a single device, and the procedure is less complicated than the former. This is the reason for selecting the latter for the model.

Conclusion

We evaluated microglia A β phagocytosis using SCST, which was reported to improve the symptoms of dementia patients in a previous study⁴). The results showed that the SCST addition may increase phagocytosis activity in a dose-dependent manner. Microarray analysis showed that various gene expressions were induced during A β phagocytosis. In the future, we plan to verify whether the SCST addition to microglia is effective for gliaprotection and analyze what changes occur in gene expression.

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Conflict of Interest Declaration

None in particular.

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