Increased levels of Aβ42 decrease the lifespan of ob/ob mice with dysregulation of microglia and astrocytes

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Abstract
Clinical studies have indicated that obesity and diabetes are associated with Alzheimer’s disease (AD) and neurodegeneration. Although the mechanisms underlying these associations remain elusive, the bidirectional interactions between obesity/diabetes and Alzheimer’s disease (AD) may be involved in them. Both obesity/diabetes and AD significantly reduce life expectancy. We generated AppNL-F/wt knock-in; ob/ob mice by crossing AppNL-F/wt knock-in mice and ob/ob mice to investigate whether amyloid-β (Aβ) affects the lifespan of ob/ob mice. AppNL-F/wt knock-in; ob/ob mice displayed the shortest lifespan compared to wild-type mice, AppNL-F/wt knock-in mice and ob/ob mice to investigate whether amyloid-β (Aβ) affects the lifespan of ob/ob mice. AppNL-F/wt knock-in; ob/ob mice displayed the shortest lifespan compared to wild-type mice, AppNL-F/wt knock-in mice, and ob/ob mice. Notably, the Aβ42 levels were increased at minimum levels before deposition in AppNL-F/wt knock-in mice and AppNL-F/wt knock-in; ob/ob mice at 18 months of age. No differences in the levels of several neuronal markers were observed between mice at this age. However, we observed increased levels of glial fibrillary acidic protein (GFAP), an astrocyte marker, in AppNL-F/wt knock-in; ob/ob mice, while the levels of several microglial markers, including CD11b, TREM2, and DAP12, were decreased in both ob/ob mice and AppNL-F/wt knock-in; ob/ob mice. The increase in GFAP levels was not observed in young AppNL-F/wt knock-in; ob/ob mice.

Abbreviations: Aβ, β-amyloid; AD, Alzheimer’s disease; APP, amyloid precursor protein; CNS, central nervous system; DEA, diethylamine; GFAP, glial fibrillary acidic protein; INSR, insulin receptor; IRS1, insulin receptor substrate-1; IRS2, insulin receptor substrate-2; PDK1, pyruvate dehydrogenase kinase 1; PSD95, post-synaptic density protein 95; SYT1, synaptotagmin 1; NF-H, neurofilament heavy chain.

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Thus, the increased Aβ42 levels may decrease the lifespan of ob/ob mice, which is associated with the dysregulation of microglia and astrocytes in an age-dependent manner. Based on these findings, the imbalance in these neuroinflammatory cells may provide a clue to the mechanisms by which the interaction between obesity/diabetes and early AD reduces life expectancy.

KEYWORDS
Alzheimer's disease, astrocytes, diabetes, lifespan, microglia, obesity

1 | INTRODUCTION

Clinical studies have indicated that obesity and diabetes are associated with Alzheimer's disease (AD) and neurodegeneration, although the mechanisms underlying this association remain elusive. Both obesity and diabetes are associated with a significant reduction in life expectancy. Dementia also shortens the life expectancy; estimates of median survival after the onset of dementia range from 5 to 10 years. However, it has not been clearly determined whether obesity/diabetes and Alzheimer's disease (AD) interact with each other to modulate the lifespan. Moreover, the mechanisms by which diabetes and AD decrease the life expectancy have not been elucidated.

Bidirectional interactions may exist between obesity/diabetes and AD. Obesity/diabetes causes neurodegeneration by inducing dysfunction of vascular systems, glucose metabolism, and insulin signaling, as well as by modifying β-amyloid (Aβ)/tau metabolisms. In turn, AD influences systemic glucose metabolism by inducing behavioral changes, memory disturbances, hypothalamic dysfunction, frailty, and possibly changes in plasma/peri-vascular Aβ levels. Thus, various molecular, cellular, inter-organ, physical, and clinical factors might contribute to the bidirectional interactions between obesity/diabetes and AD.

In the present study, we have investigated whether Aβ affects the lifespan of ob/ob mice by crossing AppNL-F/wt knock-in mice and ob/ob mice. AppNL-F/wt knock-in; ob/ob mice have the shortest lifespan than wild-type mice, AppNL-F/wt knock-in; ob/ob mice, and ob/ob mice. Notably, Aβ42 levels were increased before deposition in AppNL-F/wt knock-in mice and AppNL-F/wt knock-in; ob/ob mice, but their levels were mostly comparable in these mice. Moreover, both AppNL-F/wt knock-in; ob/ob mice and ob/ob mice showed decreased levels of microglial markers and only AppNL-F/wt knock-in; ob/ob mice showed increased levels of an astrocyte marker. Thus, increased levels of Aβ42 may shorten the lifespan of ob/ob mice with dysregulation of microglia and astrocytes.

2 | MATERIALS AND METHODS

2.1 | Animals

All mice were on the same genetic background (C57BL/6J), group housed without enrichment structures in a specific pathogen-free environment in ventilated cages, and used in the experiments according to the Guideline for the Care and Use of Laboratory Animals of our research facilities. Male heterozygous AppNL-F/wt knock-in mice were cross-bred with female heterozygous ob/+ mice to generate AppNL-F/wt knock-in; ob/+ founder mice. These male AppNL-F/wt knock-in; ob/+ founder mice were then cross-bred with female heterozygous ob/+ mice to obtain AppNL-F/wt knock-in; ob/ob, AppNL-F/wt knock-in, ob/ob, and WT littermate mice, and housed in the same cage. We employed two animal cohorts: the young cohort and the old cohort. In the old cohort (n = 169), we recorded date of death from the time when genotypes were determined (~1 month of age), until 18 months of age, when the animals were euthanized for biochemical/histochemical analyses. Notably, 23.7% (40/169) of mice died naturally, while 76.3% (129/169) of mice were euthanized at 18 months. These naturally dead mice were treated as the event, while euthanized mice were censored in the survival analysis. The young cohort (n = 50) was euthanized at 6 months of age for biochemical/histochemical analyses.

2.2 | Tissue collection and sample preparation

Mice were anesthetized with xylazine and ketamine (20 and 100 mg/kg, respectively). Blood was collected by the cardiac puncture. After transcardial perfusion with phosphate buffered saline plus a complete protease inhibitor cocktail (Nacalai Tesque), the brain was removed, divided along the sagittal
plane, and the right hemisphere (cortical and hippocampal areas) was stored at −80°C until the biochemical analysis. The left hemisphere was fixed with 4% of paraformaldehyde overnight and embedded in paraffin for the histological analysis. For the biochemical analysis, tissues were pulverized in a prechilled (on dry ice) BioPulverizer (BioSpec) and homogenized with a polytron homogenizer (KINEMATICA) at a ratio of 20 ml/g of wet weight brain tissue in ice-cold RIPA lysis buffer (Millipore) containing 0.1% SDS or diethylylamine (DEA) (for Aβ extraction) and complete protease inhibitor cocktail (Roche) on ice. After centrifugation at 100,000 g for 1 hour at 4°C, the supernatant was collected and used for the biochemical analyses. The order of sample preparation and following analyses were randomly assigned independent of genotypes.

### 2.3 ELISAs and other biochemical assays

Levels of post-synaptic density 95 (PSD95), glial fibrillary acidic protein (GFAP), and CD11b were determined using ELISAs as previously described. Levels of Aβ x-42 were determined using an ELISA employing a mouse monoclonal BC05 (recognizes C-terminus of Aβ42 [43]) capture antibody and a biotin-conjugated mouse monoclonal BNT77 (recognizes 11-28 aa of Aβ) detection antibody (Wako). A synthetic Aβ 1-42 peptide (Peptide Institute Inc) was used as a standard.

Levels of neurofilament heavy chain (NF-H) were determined with an ELISA using a mouse monoclonal SMI 35 capture antibody (BioLegend) and a rabbit polyclonal NF200 detection antibody (Sigma). Purified bovine NF-H (PROGEN) was used as a standard. Levels of synaptotagmin 1 (SYT1) were determined with an ELISA using a rabbit polyclonal SYT1 cytoplasmic tail capture antibody and a biotin-conjugated mouse monoclonal 41.1 detection antibody (Synaptic Systems). Recombinant human SYT1 proteins (Origene) were used as standards. Levels of phospho-tau were determined with an ELISA using a mouse monoclonal Tau5 capture antibody (BioLegend) and a biotin-conjugated mouse AT270 detection antibody (Thermo Fisher Scientific). Synthetic peptides containing both epitopes of Tau5 and AT270 (GenScript) were used as standards. Colorimetric quantification was performed using an iMark plate reader (Bio-Rad) after incubations with horseradish peroxidase (HRP)-linked Avidin-D (Vector) or anti-rabbit IgG (H + L), HRP conjugate (Promega), and the 3,3′,5,5′-tetramethylbenzidine substrate (Sigma). Levels of Aβ x-40 were determined using Human/Rat Aβ(40) ELISA Kit Wako II according to the manufacturer’s protocol (Wako). Levels of insulin were determined using Ultra Sensitive Mouse Insulin ELISA kit according to the manufacturer’s protocol (Morinaga). Levels of glucose was determined using Glucose Assay Kit-WST according to the manufacture’s protocol (Dojindo).

### 2.4 Quantitative real-time PCR

Total RNA was purified using RNeasy Lipid Tissue Mini kit (QIAGEN), eluted in nucleic-acid free water and stored at −80°C. Reverse transcription was performed using ReverTra Ace qPCR RT Master Mix (TOYOBO). Real-time PCR was conducted with the Luna Universal qPCR Master Mix (NEW ENGLAND BioLabs) using an i1000 thermal cycler (Bio-Rad) to detect levels of the TREM2, DAP12, IRS (insulin receptor substrate)-1, IRS-2, PDK1 (pyruvate dehydrogenase kinase 1), INSR (insulin receptor), and β-actin mRNAs. The following primers were used: IRS-1 forward primer CGATGGCTTCTAGACGTG and reverse primer CAGCCCGCTTGTGGATGTGG; IRS-2 forward primer CTGCCGTCTCTCCTCAAGTGC and reverse primer GGGGTCAATGGCCATGGTGC; Pdk1 forward primer GAGCTTCGGTCAAGTGAATGC and reverse primer TCTGGAGAACTTGTGCGGA; INSR forward primer ATGGGGCTCGGAGAGGAT and reverse primer GAGTGCACATCCAGCTGA; and DAP12 forward primer GTTGACTGTGTGCATTGTGCCT and reverse primer CCCCCGGTCTCCTCCCTGA (Integrated DNA Technologies). The relative expression levels were quantified and analyzed using Bio-Rad iCycler iQ software (Bio-Rad). Relative mRNA levels were calculated using ΔΔCt method with β-actin serving as an internal control for each specific gene amplification reaction.

### 2.5 Immunohistochemical staining

Mouse brains were fixed as described above in the section listing the tissue collection and sample preparation methods. Brain samples were embedded in paraffin wax and cut into sections with a thickness of 10 μm. Mouse samples used for immunohistochemical staining were chosen if their ELISA values were similar to the average value for each group. Brain sections were submerged in EDTA buffer (1 mM, pH 8.0), and autoclaved for antigen retrieval. Sections were immersed in 95% of formic acid for Aβ staining. Endogenous peroxidase activity was quenched by incubating the sections with hydrogen peroxide. Sections were incubated with primary antibodies at 4°C overnight. We used an anti-Aβ (6E10, epitope: 3-8 aa of Aβ) antibody (Merck), anti-GFAP antibody (Agilent Technologies), and anti-Iba1 antibody (Wako). Sections were incubated with an HRP-conjugated secondary antibody and diaminobenzidine reaction mixture (Vector Laboratories). Stained sections were observed under a microscope (BZ-X810, Keyence).
2.6 | Statistical analysis

All statistical analyses were performed using JMP Pro software (version 12, SAS Institute Inc). In the survival analysis, Cox proportional hazard models with sex and genotype were as covariates, the date of birth as the time of origin, and the age at death as the time of the event were employed to address the risk of death. In other analyses, effects of each genotype were addressed through (a) the linear regression models adjusting for sex followed by Tukey’s HSD test when both genders were included, or (b) one-way ANOVA followed by Tukey’s HSD test when one gender was included. *P*-values less than 0.05 were considered significant. Data are presented as adjusted means ± standard errors of the means when both genders were included or means ± standard deviations of the means when one gender was included.

3 | RESULTS

3.1 | The shortest lifespan is observed for AppNL-F/wt knock-in; ob/ob mice, despite the minimal and comparable increase in Aβ42 levels

We previously crossed amyloid precursor protein (APP) 23 mice with ob/ob mice to study the interaction between obesity/diabetes and AD,9 while a new generation of AD mouse models that express APP at comparable levels to controls has been developed.22 Thus, we crossed these AppNL-F/wt knock-in mice, instead of APP23 mice, with ob/ob to study the interaction between obesity/diabetes and AD. Interestingly, AppNL-F/wt knock-in; ob/ob mice displayed the shortest lifespan compared to wild-type mice, AppNL-F/wt knock-in mice, and ob/ob mice (Figure 1A), as supported by the results from the proportional hazard model adjusted for sex (Figure 1B). Such trends were also observed in the analyses stratified by sex (Figure S1). Additionally, ob/ob mice had a shorter lifespan than control mice, consistent with a previous study.24 Lower body weights were recorded for AppNL-F/wt knock-in; ob/ob mice than ob/ob mice after adjusting for sex (Figure 1C), while such trends were significant in only female mice in the analysis stratified by sex (Figure S1). Sex difference is becoming focused in AD research field.25,26 A new cohort with increased samples will help us to understand the effects of sex on outcomes. We could not measure blood glucose levels during the course of our study until 18 months of age, because any stressful or invasive procedure might have impacts on lifespan of our mice. We measured insulin levels and casual glucose levels in serum from the mice when they were killed at 18 months of age and found that the levels of insulin in ob/ob mice and AppNL-F/wt knock-in; ob/ob mice were comparable and higher than wild-type mice and AppNL-F/wt knock-in mice and that there were no significant difference between the groups in casual blood glucose levels (Figure S2).

β-amyloid accumulation in the brains of 18-month-old AppNL-F/wt knock-in mice and AppNL-F/wt knock-in; ob/ob mice was below the detection limit of IHC staining (Figure 2A), consistent with the observation that AppNL-F/wt knock-in mice do not form senile plaques until a very old age exceeding 24 months.22 GuHCl extraction, which requires a greater dilution of samples, did not produce detectable Aβ levels in AppNL-F/wt knock-in mice (data not shown). DEA extraction allows us to measure lower Aβ levels using an ELISA.27 Using DEA extraction, we observed increases in Aβ42 levels in the brains of AppNL-F/wt knock-in mice and ob/ob mice (Figure 1A), as supported by the results from the proportional hazard model adjusted for sex (Figure 1B). Such trends were also observed in the analyses stratified by sex (Figure S1). Additionally, ob/ob mice had a shorter lifespan than control mice, consistent with a previous study.24 Lower body weights were recorded for AppNL-F/wt knock-in; ob/ob mice than ob/ob mice after adjusting for sex (Figure 1C), while such trends were significant in only female mice in the analysis stratified by sex (Figure S1). Sex difference is becoming focused in AD research field.25,26 A new cohort with increased samples will help us to understand the effects of sex on outcomes. We could not measure blood glucose levels during the course of our study until 18 months of age, because any stressful or invasive procedure might have impacts on lifespan of our mice. We measured insulin levels and casual glucose levels in serum from the mice when they were killed at 18 months of age and found that the levels of insulin in ob/ob mice and AppNL-F/wt knock-in; ob/ob mice were comparable and higher than wild-type mice and AppNL-F/wt knock-in mice and that there were no significant difference between the groups in casual blood glucose levels (Figure S2).

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knock-in mice or App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice, compared to wild-type or \textit{ob/ob} mice (Figure 2B). Their levels were mostly comparable between App\textsubscript{NL-F/wt} knock-in mice and App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice (Figure 2B), while App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice showed some reduction of Aβ42 levels in male group, but not female group (Figure S2). There is no difference or some trends of reduction of Aβ40 levels in the brains of App\textsubscript{NL-F/wt} knock-in mice or App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice, especially in male group (Figures 2C and S3). However, there is significant increase in the ratio of Aβ42 to Aβ40 by App\textsubscript{NL-F/wt} knock-in (Figure 2D), consistent with the previous study.\textsuperscript{22}

3.2 | Microglia and astrocytes are dysregulated in App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice

To explore the mechanism by which the lifespan of App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice are decreased, we initially analyzed the brain levels of neuronal markers, NF-H (Figure 3A), SYT1 (Figure 3B), PSD95 (Figure 3C), and phospho-tau (Figure 3D) through ELISAs, where sample values were in the non-saturable range of standard curve (Figure 3E-H), but did not detect any changes in App\textsubscript{NL-F/wt} knock-in mice, \textit{ob/ob} mice, and App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice compared to wild-type mice. Because insulin signalling in the brain is involved in regulating the lifespan,\textsuperscript{28,29} we also assessed the levels of molecules related to insulin signalling. \textit{Irs2} and \textit{Pdk1} expression were significantly reduced in \textit{ob/ob} mice, but not App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice compared to wild-type mice. Because insulin signalling in the brain is involved in regulating the lifespan,\textsuperscript{28,29} we also assessed the levels of molecules related to insulin signalling. \textit{Irs2} and \textit{Pdk1} expression were significantly reduced in \textit{ob/ob} mice, but not App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice compared to WT controls (Figure S4). Thus, the shortest lifespan observed in App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice may not be explained by a reduction in the expression of molecules involved in neuronal markers or insulin signalling in the brain.

However, the protein expression of a microglial marker, CD11b, was reduced in both \textit{ob/ob} and App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice (Figure 4A). Interestingly, mRNA levels of \textit{Trem2} and \textit{Dap12}, genes that are abundantly expressed in microglia and play important roles in AD pathogenesis,\textsuperscript{30} were also reduced in both \textit{ob/ob} and App\textsubscript{NL-F/wt} knock-in; \textit{ob/ob} mice (Figure 4B,C), while the numbers of Iba1-expressing...
microglia assessed by IHC staining were not altered (Figure S5). Moreover, levels of the astrocytic marker GFAP were increased in AppNL-F/wt knock-in; ob/ob mice, compared to ob/ob mice, as determined using an ELISA and IHC staining (Figure 4D,E).

We also analyzed young mice that were harvested at 6 months old. Notably, body weights were comparable between ob/ob mice and AppNL-F/wt knock-in; ob/ob mice, compared to ob/ob mice, as determined using an ELISA and IHC staining (Figure 4D,E).

DISCUSSION

While clinical studies have indicated associations between obesity/diabetes and AD and neurodegeneration, it has been not clearly determined whether these diseases interact with each other to alter lifespan. As shown in the present study using AppNL-F/wt knock-in; ob/ob mice obtained by crossing AppNL-F/wt knock-in mice and ob/ob mice, increased Aβ42 levels affected the lifespan of ob/ob mice through a
mechanism associated with the dysregulation of microglia and astrocytes.

Both dementia and obesity/diabetes are associated with a shorter life expectancy. In the present study, $\text{App}^{\text{NL-Fwt}}$ knock-in; $\text{ob}/\text{ob}$ mice had the shortest lifespan compared to wild-type mice, $\text{App}^{\text{NL-Fwt}}$ knock-in mice, and $\text{ob}/\text{ob}$ mice. Additionally, $\text{ob}/\text{ob}$ mice had a shorter lifespan than control mice, consistent with a previous report. Patients with both Apoe4 and diabetes have a shorter lifespan than individuals with Apoe4 and without diabetes, individuals without Apoe4 and with diabetes, and subjects with neither Apoe4 nor diabetes (Shinohara and Sato, unpublished results). Because Apoe4 is a strong genetic risk factor for AD, the results of the present study using an animal model of AD indicate a possible interaction between obesity/diabetes and AD on altering the lifespan of humans. Further human studies are required to confirm this observation.

There is a significant weight loss in $\text{App}^{\text{NL-Fwt}}$ knock-in; $\text{ob}/\text{ob}$ mice compared to $\text{ob}/\text{ob}$ mice, especially in female mice at old age. Because food intake and/or energy expenditure are reported to be altered in some AD models, measurement of these parameters would be interesting. Frailty is also associated with AD. If this weight loss is linked to frailty, the weight loss may also impact survival. Further studies including food intake, energy expenditure, and frailty index are required to determine the mechanisms of weight loss in $\text{App}^{\text{NL-Fwt}}$ knock-in; $\text{ob}/\text{ob}$ mice.

In the present study, comparable Aβ levels were observed between $\text{App}^{\text{NL-Fwt}}$ knock-in mice and $\text{App}^{\text{NL-Fwt}}$ knock-in; $\text{ob}/\text{ob}$ mice. Firm evidence showing that diabetes increases Aβ deposition in humans is not currently available, although diabetes/hyperglycaemia modify Aβ accumulation in the brains of wild-type animals and animal models of AD. Moreover, several neuropathological studies
revealed less senile plaques in patients with AD who have been diagnosed with diabetes compared with patients without diabetes. Thus, we may be able to consider the observation that obesity/diabetes and AD interact to determine lifespan when attempting to obtain a better understanding of the discrepancies among the results of neuropathological studies.

Obesity and diabetes are associated with the dysregulation of macrophages in the peripheral system in humans and animal models. Indeed, Obesity and diabetes increase the susceptibility to infection and impairs wound healing. Recently, emerging evidence supports the hypothesis that the dysfunction of microglia, the counterpart of macrophages in the brain, plays important roles in the pathogenesis of AD. In the present study, levels of the microglial marker CD11b were reduced in ob/ob and App<sup>NL-F<sub>vec</sub></sup> knock-in; ob/ob mice at both 6 and 18 months. Interestingly, the expression of Trem2 and its partner Dap12 were also reduced in these mice. Based on accumulating evidence from recent studies, a reduction in the activity of the TREM2 signalling pathway promotes neurodegeneration. Because brain Aβ42 levels were comparable in App<sup>NL-F<sub>vec</sub></sup> knock-in mice and App<sup>NL-F<sub>vec</sub></sup> knock-in; ob/ob mice, soluble Aβ42 might not be targeted by microglia. Rather, the dysregulation of microglia may cause susceptibility to Aβ.

We observed increased GFAP levels in 18-month-old App<sup>NL-F<sub>vec</sub></sup> knock-in; ob/ob mice, while its levels decreased in 6-month-old animals. Their age-dependent increases were also observed immunohistochemically (Figure S7). Thus, astrocytes might be activated by obesity/diabetes and AD during aging. The activated astrocytes may be reactive astrocytes. Recent studies indicate that reactive astrocytes obtain neurotoxic properties both through a gain of toxic function and a loss of their neurotrophic effects and affect AD pathogenesis, although astrocyte reactions might also be heterogeneous. Because the level of Aβ was not increased in App<sup>NL-F<sub>vec</sub></sup> knock-in; ob/ob mice compared to App<sup>NL-F<sub>vec</sub></sup> knock-in mice, one possible mechanism is that astrocytes might be more vulnerable to the same level of Aβ in App<sup>NL-F<sub>vec</sub></sup> knock-in; ob/ob mice with dysregulated microglia. Further
studies, including single-cell analyses, would be required to test this hypothesis.

Because insulin signalling in the brain is involved in regulating the lifespan, we assessed the expression of molecules involved in insulin signalling. *Irs2* expression was significantly reduced in *ob/ob* mice, but not *App* 

knock-in; *ob/ob* mice. Thus, the shortest lifespan observed in *App* 

knock-in; *ob/ob* mice may not be explained by the decrease in the expression of molecules involved in insulin signalling. The samples analyzed in the present study were obtained from 18-month-old mice that survived to this age. Because, we were unable to analyze the mice that died before reaching 18 months of age, the precise mechanism underlying the substantially decreased lifespan of *App* 

knock-in; *ob/ob* mice should be analyzed further in future studies.

There are several limitations in our animal models. We chose *App* 

knock-in mice as an AD model and *ob/ob* mice as a model of obesity linked to diabetes, with affected peripheral insulin signalling (Figure S8), in this study. *App* 

knock-in mice do not show any amyloid pathology or neurodegenerative changes as shown in this study, and *ob/ob* mice become obese and mildly diabetic due to leptin deficiency. In the future study, we will study the interaction between obesity/diabetes and AD on lifespan in both humans and other animal models. In this study as shown in the original report of *App* 

knock-in mice, there is no difference in lifespan between *App* 

knock-in mice and wild-type mice until 18 months. Further observational studies to older ages are required to determine whether Aβ42 decrease lifespan in wild-type background, in additions to *ob/ob* background. Alternatively, there is also a possibility that Aβ42 decrease lifespan only in *ob/ob* background, but not in wild-type background, suggesting that Aβ42 is enough toxic to decrease lifespan only in *ob/ob* background, but not in wild-type background. It would be also interesting to see the results of the investigation of *App* 

knock-in; *ob/ob* mice at 24 months of age when *App* 

knock-in mice show the formation of senile plaques in the brain. However, we terminated this study at 18 months of age to analyze the mechanism of the shorter lifespan in *App* 

knock-in; *ob/ob* mice using the tissues of mice which survived until 18 months of age. Further investigation, including the analysis of *App* 

knock-in mice with senile plaques, will allow us to investigate the impact of senile plaque formation on lifespan in *ob/ob* mice.

In summary, increased levels of Aβ42 decrease the lifespan of *ob/ob* mice and were associated with the dysregulation of microglia and astrocytes. These findings may provide evidence of an interaction between obesity/diabetes and AD to regulate the lifespan. An understanding of the precise mechanisms of this interaction will likely provide novel targets for the treatment of obesity/diabetes and AD.

5 | CONCLUSIONS

The increased Aβ42 levels may decrease the lifespan of *ob/ob* mice, which is associated with the dysregulation of microglia and astrocytes. The imbalance in these neuroinflammatory cells may provide a clue to the mechanisms by which the interaction between obesity/diabetes and early AD reduces life expectancy.

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DISCLOSURE STATEMENT

The authors have no conflicts of interest related to the content of this article to declare.

AUTHOR CONTRIBUTIONS

N. Sato and Mi. Shinohara designed research; Mi. Shinohara, Y. Tashiro, Mo. Shinohara, J. Hirokawa, M. Onishi-Takeya, and M. Mukouzono performed research; K. Suzuki, S. Takeda, A. Fukumori, and R. Morishita analyzed data; N. Sato and Mi. Shinohara wrote the paper; T. Saito and T.C. Saida contributed to providing a new AD model mouse.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

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