Saturated fatty acids activate microglia via Toll-like receptor 4/NF-κB signalling

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(Received 13 January 2011 – Revised 24 March 2011 – Accepted 30 March 2011 – First published online 29 June 2011)

Abstract

Diets rich in SFA have been implicated in Alzheimer’s disease (AD). There is strong evidence to suggest that microglial activation augments the progression of AD. However, it remains uncertain whether SFA can initiate microglial activation and whether this response can cause neuronal death. Using the BV-2 microglial cell line and primary microglial culture, we showed that palmitic acid (PA) and stearic acid (SA) could activate microglia, as assessed by reactive morphological changes and significantly increased secretion of pro-inflammatory cytokines, NO and reactive oxygen species, which trigger primary neuronal death. In addition, the mRNA level of these pro-inflammatory mediators determined by RT-PCR was also increased by PA and SA. We further investigated the intracellular signalling mechanism underlying the release of pro-inflammatory mediators from PA-activated microglial cells. The present results showed that PA activated the phosphorylation and nuclear translocation of the p65 subunit of NF-κB. Furthermore, pyrrolidine dithiocarbamate, a NF-κB inhibitor, attenuated the production of pro-inflammatory mediators except for IL-6 in PA-stimulated microglia. Administration of anti-Toll-like receptor (TLR)4-neutralising antibody repressed PA-induced NF-κB activation and pro-inflammatory mediator production. In conclusion, the present in vitro study demonstrates that SFA could activate microglia and stimulate the TLR4/NF-κB pathway to trigger the production of pro-inflammatory mediators, which may contribute to neuronal death.

Key words: Microglia; SFA; NF-κB; Toll-like receptor 4

Epidemiological data suggest that a diet rich in SFA is considered an increased risk factor for the development of Alzheimer’s disease (AD). For example, in a 21-year follow-up study, it was found that abundant SFA intake from milk products and spreads at midlife was associated with poorer global cognitive function and prospective memory(1). Other studies have demonstrated that a greater intake of saturated fat increased the risk for impaired cognitive function in middle-aged or aged populations(2–4). This notion has been supported by animal studies. In this connection, it has been reported that rodents fed high levels of SFA also show impaired learning and memory performance and develop AD-like pathophysiological changes in their brains(5,6). Fatty acids are free to cross the blood–brain barrier(7). Therefore, brain fatty acid homeostasis may be dependent on their levels in the periphery. It is therefore conceivable that diets rich in SFA may increase brain uptake of intact NEFA from the plasma through the blood–brain barrier(8). In addition, the fatty acid profile of neurofibrillary tangles in the AD brain is rich in palmitic acid (PA) and stearic acid (SA)(9), and the white matter in the AD brain is characterised by high total fatty acid contents(10).

PA and SA were reported to increase hyperphosphorylation of tau, and up-regulate β-secretase, the rate-limiting enzyme in the production of amyloid β peptides in primary rat cortical neurons(11,12). These actions were mediated by the above-mentioned two SFA on astrocytes, possibly through enhanced astrocytic synthesis of ceramide(13). Several fatty acids have been reported to stimulate the aggregation of tau protein and amyloid β in vitro(14). Despite the accumulating data, the basic mechanism behind the causal relationship...
between SFA and the pathogenesis of AD has not been well established.

It is well documented that microglia, the resident macrophages in the brain, play a central role in mediating chronic inflammatory conditions in AD\[^{15}\]. In the ramified state, microglia actively survey the microenvironment and ensure normal central nervous system (CNS) activity by secreting neurotrophic factors such as neuronal growth factor. They are activated in response to specific stimuli and produce a host of pro-inflammatory cytokines, chemokines and reactive oxygen species (ROS). Although microglial activation plays an important role in phagocytosis of dead cells in the CNS, microglia cause inflammatory responses leading to neuronal death and brain injury when they are over-activated and dysregulated\[^{16}\]. Therefore, identification of the regulators involved in the initiation and maintenance of microglial activation may lead to a better understanding of inflammatory processes leading to AD. However, as far as can be ascertained, there is a total lack of information relating to the modulation of microglial activation by SFA. We report here that SFA could activate microglia to a pro-inflammatory state as evidenced by reactive morphological changes and significantly increased secretion of pro-inflammatory cytokines including TNF-\(\alpha\), IL-1\(\beta\) and IL-6, as well as NO and ROS via Toll-like receptor (TLR)-4/NF-\(\kappa\)B signalling in the microglial cell line, BV-2 cells, and primary microglia.

Materials and methods

Animals

BALB/c mice were used. All animals were obtained from the Laboratory Animal Centre, Shandong University. All animals were kept under controlled 12 h light–12 h dark conditions, temperature (23\(^\circ\)C) and humidity (60%). In the handling and care of all animals, the International Guiding Principles for Animal Research, as stipulated by the WHO (1985) and as adopted by the Laboratory Animal Centre, Shandong University, were followed. During the study, the number of animals used and their suffering were minimised.

Microglial cell culture

BV-2 cells were maintained in Dulbecco's modified Eagle's medium (DMEM; Hyclone Co., Logan, UT, USA) with 10% fetal bovine serum (Hyclone Co.), 2 mg/l-glutamine, penicillin (100 U/ml) and streptomycin (100 \(\mu\)g/ml) (Sigma-Aldrich, St Louis, MO, USA) in a 5\% CO\(_2\) incubator. For all experiments, BV-2 cells were used at 75 to 80\% confluency. Before the experiment, plated cells were incubated with serum-free DMEM for 1 h. After this, the medium was replaced with serum-free DMEM containing appropriate fatty acid–albumin complexes for 12 h. Controls received BSA and vehicle only.

Preparation of fatty acid–albumin complexes

PA or SA was solubilised in ethanol at 70\%. Then PA or SA was combined with fatty acid-free and low-endotoxin bovine serum albumin (BSA) at a molar ratio of 10:1 (fatty acid:albumin) in serum-free medium at 50\°C for 6 h for a final PA or SA concentration of 25–200 \(\mu\)M as described previously\[^{18}\]. The fatty acid–albumin complex solution was freshly prepared before each experiment. The final concentration of ethanol was \(< 0.5\%\). In most of the experiments, BV-2 cells or primary microglial cells were treated with individual SFA at 25–200 \(\mu\)M concentration, while the controls received BSA and vehicle only.

To evaluate the possible contamination of PA or SA with LPS, the endotoxin content was determined by the chromogenic Limulus amebocyte lysate test, following the manufacturer's instructions (Cambrex Bio Science, Walkersville, MD, USA). The endotoxin content in the 100 \(\mu\)M-PA and 100 \(\mu\)M-SA solution was \(\leq 3.45 \times 10^{-3}\) pg/ml, which is far below the concentration required to induce microglial activation under our assay conditions.

Conditioned medium

To assess bystander neuronal death by factors released by microglial cells following PA treatment, BV-2 cells were seeded in 60 mm culture plates at a density of 3 \(\times\) 10\(^5\) cells/plate. After the cells became confluent, they were incubated with serum-free DMEM for 1 h. After this, the medium was replaced by serum-free DMEM containing appropriate fatty acid–albumin complexes for 12 h. Controls received BSA and vehicle only. After 12 h, the medium was changed with fresh serum-free DMEM for 12 h, and then the supernatant fractions were collected, filtered and added onto primary neurons cultured in poly-D-lysine-coated twelve-well plates.

Neuronal culture and apoptosis analysis

Primary cultures of mouse cortical neurons were prepared as previously described\[^{19}\]. Briefly, cortical neurons were harvested from mice, aged 1–2 d, using the serum-free Neurobasal medium with B27 supplement system (Invitrogen Corp., Carlsbad, CA, USA). Cortical cells were plated at a density of 1.5 \(\times\) 10\(^5\) cells per well in poly-D-lysine-coated twelve-well plates and allowed to differentiate for 7 d. At day 7, the mouse neuronal medium was removed and replaced by a conditioned medium from microglial cells. Neurons were incubated with microglia-conditioned medium for 2 d, and...
then analysed by Hoechst 33342 nuclei staining for the
detection of morphological features of apoptotic cell death.
Hoechst was added to the culture medium at a final
centration of 10 µg/ml, which was then incubated in the dark at
100% humidity for 10 min at 37°C. The cells were then exam-
ined under a fluorescence microscope. Undamaged cell nuclei
were large and diffusely stained whereas apoptotic nuclei
showed chromatin that was condensed and fragmented.

**ELISA**

BV-2 cells (3 × 10^5 cells per well in a twelve-well plate) were
pretreated with different concentrations of PA (0, 25, 50 or
100 µM), LPS or inhibitors at the indicated times. The supernat-
ate fraction of the culture medium from the various
treatments was then collected. The levels of cytokines, TNF-
α, IL-1β and IL-6 in the culture medium were measured
using commercially available ELISA kits (R&D Systems Inc.,
Minneapolis, MN, USA) according to the manufacturer’s
instructions. Briefly, serial dilutions of protein standards and
samples were added to ninety-six-well ELISA plates, followed
by biotinylated anti-TNF-α, IL-1β or IL-6 Ab. After rinsing with
wash buffer, the prepared solution of avidin, horseradish
peroxidase-conjugated complex was added followed by the
addition of substrate solution. The reaction was terminated
by the stopping solution. The optical density was detected
at 450 nm in a microplate reader (Bio-Rad Laboratories,
Hercules, CA, USA). Each sample concentration was calculated
from the linear equation derived from the standard curve of
known concentrations of the cytokine.

**Assay of NO production**

BV-2 cells (3 × 10^5 cells per well in a twelve-well plate), were
pretreated with different concentrations of PA (0, 25, 50 or
100 µM), LPS or inhibitors at the indicated times. The superna-
atant fraction of the culture medium from the various treat-
ments was collected. NO production was assessed by
measuring the accumulation of nitrite in the culture medium
by the Griess reaction. The culture medium was mixed with
an equal volume of Griess reagent (0.1% N-(1-naphthy-
lethylenediamine dihydrochloride and 1% sulfanilamide in 5% phos-
phoric acid; Sigma-Aldrich) in a ninety-six-well plate and
incubated at room temperature for 10 min. Absorbance was
measured at 550 nm in a microplate reader (Bio-Rad Labora-
tories). Sodium nitrite, diluted in culture medium at concen-
trations ranging from 10 to 100 µM, was used to prepare a
standard curve.

**Reactive oxygen species assay**

Intracellular ROS levels were measured by 2',7'-dichlorodihy-
drofluorescein diacetate (H2DCFDA) and dihydroethidium
(DHE) assays. H2DCFDA or DHE is a membrane-permeable
dye that is oxidized by intracellular ROS to the fluorescent pro-
duct DCF or ethidium, respectively. Briefly, BV-2 cells were
stimulated with or without PA, or LPS for 12 or 24 h, and
the culture medium was first removed. The cells were washed
three times with PBS, and were incubated with either 10 µM-
H2DCFDA or 2 µM-DHE (Molecular Probes, Eugene, OR,
USA) for 20 min at 37°C. The cells were then washed three
times with PBS and examined with a Nikon TE2000U micro-
scope (Nikon, Tokyo, Japan). The fluorescence was measured
at 485 nm for excitation and 530 nm for emission with a
fluorescence plate reader (Fluroskan Ascent II; Labsystems,
Helsinki, Finland). The increased value compared with control
was considered as the increase of intracellular ROS.

**Cell viability assay**

Cell viability was determined using the 3-[4, 5-dimethylthiazol-
2-yll-2, 5-diphenyltetrazolium bromide (MTT) assay. BV-2 cells
were plated into ninety-six-well culture plates at a density of
1 × 10^4 cells/well with 200 µl culture medium per well in
triplicate. When reaching 75% confluency, the cells were
incubated in the absence or presence of PA (25, 50, 100,
200 µM), respectively, for 48 h. Then, 20 µl MTT solution
(5 mg/ml; Sigma-Aldrich) were added to each well and
incubated at 37°C for 4 h. The culture medium was aspirated
and followed by the addition of 200 µl dimethyl sulfoxide.
The absorbance value was measured in a microplate reader
(Bio-Rad Laboratories) at 490 nm. Values were expressed as
a percentage relative to those obtained in controls.

**Immunocytochemistry**

Microglial cells were seeded onto glass coverslips. Following
 treatment with or without PA, LPS or inhibitors at the indicated
times, cells were fixed in 4% paraformaldehyde for 10 min,
and blocked with 10% goat serum in PBS. Slides were incu-
bated overnight in a humid chamber at 4°C with the primary
Ab (mouse monoclonal anti-CD11b Ab (1:100 dilution);
rabbit polyclonal anti-inducible NO synthase (iNOS) Ab
(1:100 dilution); rabbit monoclonal anti-IFN-γ Ab p65 Ab (1:100
dilution); all Cell Signaling Technology, Beverly, MA, USA).
After primary Ab incubation, samples were washed again
and incubated in the appropriate fluorescent-conjugated
secondary Ab (goat anti-mouse/rabbit IgG (1:100 dilution);
Sigma-Aldrich) for 1 h. The cells were counterstained by
4',6-diamidino-2-phenylindole (DAPI). Images were captured
with a Nikon TE2000U microscope.

**Reverse transcription-PCR**

Total RNA was extracted from induced cell cultures using the
Trizol reagent (Gibco, Invitrogen Corp.) according to the manufac-
turer’s instructions. RNA concentration was determined
by a spectrophotometer (Bio-Rad Laboratories) at 260 nm.
Identical amounts of RNA (1 µg) were reverse transcribed
into cDNA by using a commercial RT-PCR kit (Fermentas,
Vilnius, Lithuania) according to the manufacturer’s instruc-
tions. cDNA was subsequently amplified by PCR with specific
primers (Table 1). PCR amplification of the resulting cDNA
template was conducted by using the following conditions
for thirty-two (TNF-α, IL-6, iNOS and β-actin) and thirty-five
(IL-1β) cycles; denaturation at 94°C for 30 s, annealing at
Western blot analysis

Cell-associated proteins were washed with cold PBS and lysed in ice-cold radioimmunoprecipitation assay (RIPA) buffer containing protein inhibitors. Cell lysates were incubated at 4°C for 20 min. The sample was centrifuged at 12 000 rpm for 10 min at 4°C, the supernatant fraction was then collected and protein concentration was assayed colormetrically. A quantity of 30 μg total proteins was loaded onto a 4–20% gradient polyacrylamide gel, electrophoretically transferred to a polyvinylidene difluoride membrane and probed with primary antibodies (rabbit polyclonal anti-phospho-NF-kB p65 (S536) Ab (1:500 dilution), Bioworld Technology, Inc., Minneapolis, MN, USA; rabbit monoclonal anti-NF-kB p65 Ab (1:1000 dilution), Cell Signaling Technology; rabbit polyclonal anti-iNOS Ab (1:1000 dilution), Cell Signaling Technology; rabbit polyclonal anti-β-actin (1:2000 dilution; Sigma-Aldrich) was used as an internal control. Secondary antibodies were horseradish peroxidase conjugated to goat/mouse anti-rabbit IgG (1:5000 dilution; Sigma-Aldrich). The membranes were developed using an enhanced chemiluminescence detection system (Pierce, Rockford, IL, USA).

Transient transfection and luciferase reporter gene assay

The NF-κB reporter plasmid contained three copies of the NF-κB-binding sequence fused to the firefly luciferase gene (Clontech Laboratories, Inc., Mountain View, CA, USA). The cells were cultured in a twenty-four-well plate until they reached 75–80% confluency. Transfection of the NF-κB reporter gene into the cells was performed using Lipofectamine 2000 (Invitrogen Corp.) according to the manufacturer’s instructions. Cells were transfected with 0·8 μg NF-κB reporter plasmid, 0·04 μg pRL-TK vector (Promega Corp., Madison, WI, USA) mixed with Lipofectamine 2000. After 48 h, cells were harvested and a luciferase assay was performed. To determine SFA-induced NF-κB activity, cells were incubated with or without PA or LPS for 24 h before harvesting cells for the luciferase assay. Luciferase assays were performed using the Dual-Luciferase reporter assay system (Promega Corp.) according to the manufacturer’s instructions. Luciferase activity was measured using a Monolight 2010 luminometer (Analytical Luminescence Laboratory, San Diego, CA, USA). Renilla luciferase activity was used as an internal control. The relative luciferase activity was then calculated by normalising firefly luciferase activity to Renilla luciferase activity.

Statistical analysis

Quantitative data are presented as the mean values and standard deviations of at least three independent experiments. Statistical analysis of data was done by Student’s t test or by one-way ANOVA using Dunnett’s test in multiple comparisons of means. Differences were considered statistically significant if the P value was <0·05.

Results

Treatment with SFA led to activation of microglial cells

In order to confirm that incubation with SFA would not induce microglia death, cell viability was assessed at 48 h after PA treatment by MTT. BV-2 cell viability following treatment with PA at 25 μM (94·78 (SD 4·34) %), 50 μM (101·09 (SD 14·61) %) and 100 μM (89·49 (SD 6·51) %) was not significantly different from the control (100·08 (SD 5) %). However, exposure of BV-2 cells to PA at 200 μM resulted in significantly fewer viable cells (66·82 (SD 4·91) %) as compared with cells in the control condition (Fig. 1(a)). In view of this and because PA is common in the diet and constitutes a large proportion of circulating NEFA, we used PA (25, 50 and 100 μM) as a representative SFA in most of the subsequent experiments.

The MTT assay showed that PA did not have any cytotoxicity at the concentrations of 25, 50 and 100 μM for at least 48 h on primary microglia (Fig. 1(a)); it was clearly toxic to few viable cells (66·82 (SD 4·91) %) as compared with cells in the control condition (Fig. 1(a)). In view of this and because PA is common in the diet and constitutes a large proportion of circulating NEFA, we used PA (25, 50 and 100 μM) as a representative SFA in most of the subsequent experiments.

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Table 1. PCR primers used in the present study

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward</th>
<th>Reverse</th>
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<tbody>
<tr>
<td>TNF-α</td>
<td>5′-CGTCAGCCGCTTGTCTCT-3′</td>
<td>5′-CGGAATCTCCGAAATCTAAG-3′</td>
</tr>
<tr>
<td>IL-1β</td>
<td>5′-AAAGATGAAAGGCTGTTTCCAACC-3′</td>
<td>5′-ATACTGGCTGCTGACGGTTGT-3′</td>
</tr>
<tr>
<td>IL-6</td>
<td>5′-CACTTCCAAGTCGAGGCTT-3′</td>
<td>5′-CCAGTTATCGTTAGGAGA-3′</td>
</tr>
<tr>
<td>iNOS</td>
<td>5′-CCTCTCCTCACAACCTACGAAT-3′</td>
<td>5′-CACACAAAGTGCTTGATCGTA-3′</td>
</tr>
<tr>
<td>β-Actin</td>
<td>5′-GTGGGGGCCGCCAGGCAACA-3′</td>
<td>5′-CTCCCTTAAATGTCCAGCACGATTTC-3′</td>
</tr>
</tbody>
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iNOS, inducible NO synthase.
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We next examined whether SFA treatment could affect intracellular ROS levels in BV-2 cells. The cells were treated with PA (100 µM) for 12 and 24 h, or LPS (500 ng/ml) for 24 h, following the addition of the ROS fluorescent probes H$_2$DCFDA and DHE to detect H$_2$O$_2$ and superoxide (O$_2^−$) production, respectively (Fig. 4(a)). It was observed that both PA

SFA caused increased NO release and intracellular inducible NO synthase levels

We first tested the effect of PA on NO release by measuring nitrite quantities in the supernatant fractions of BV-2 cell culture. It was found that upon treatment with different concentrations of PA (25, 50 and 100 µM) for 12, 24 and 48 h or LPS (500 ng/ml) for 24 h, NO release was significantly increased from the cells in all three treatment groups in a dose-dependent and time-dependent manner compared with the control (Fig. 3(a)). The cells exposed to different concentrations of PA (25, 50 and 100 µM) for 4 h also exhibited an increase in iNOS mRNA expression in a dose-dependent manner (Fig. 3(b) and (c)).

In conjunction with nitrite quantification, iNOS expression responsible for NO production was evaluated by Western blot and immunocytochemistry. Western blot results showed that PA dose-dependently increased iNOS expression in BV-2 cells (Fig. 3(c)). Moreover, in the PA (100 µM)-treated group, iNOS expression was higher than that in LPS-treated cells. By immunofluorescence, iNOS expression was induced after 6 h of treatment with PA (100 µM) in primary microglia cells; at 24 h after treatment, the expression was visibly more intense (Fig. 3(d)). In order to investigate whether the above effects are PA specific, we then tested another SFA – SA, which is present in the serum, accounting for close to 13% of the total fatty acids. We found that treatment with 100 µM-SA (a non-toxic concentration) for 4 h also induced a marked morphological change (data not shown) and increased expression of pro-inflammatory cytokine mRNA and iNOS mRNA in BV-2 cells (Fig. 3(e)). These results suggest that SFA play a rather general role in microglial activation.

SFA caused elevation of reactive oxygen species production

We next examined whether SFA treatment could affect intracellular ROS levels in BV-2 cells. The cells were treated with PA (100 µM) for 12 and 24 h, or LPS (500 ng/ml) for 24 h, following the addition of the ROS fluorescent probes H$_2$DCFDA and DHE to detect H$_2$O$_2$ and superoxide (O$_2^−$) production, respectively (Fig. 4(a)). It was observed that both PA

SFA induced expression and secretion of pro-inflammatory cytokines

RT-PCR analysis showed that in BV-2 cells exposed to different concentrations of PA (25, 50 and 100 µM) or LPS (500 ng/ml) for 4 h, the levels of TNF-α, IL-1β and IL-6 mRNA expression were significantly increased compared with the control (Fig. 2(a) and (b)). By ELISA, we then determined the production of TNF-α, IL-1β and IL-6 in the medium of BV-2 cells treated with PA at different concentrations (25, 50 and 100 µM) for 12, 24 and 48 h. As a positive control, BV-2 cells were stimulated with LPS (500 ng/ml) for 24 h. As shown in Fig. 2(c), both LPS and PA, at either low or high concentrations, stimulated microglia to produce increased amounts of cytokines. In addition, PA stimulated the release of TNF-α, IL-1β and IL-6 in BV-2 cells from 12 h onward at all the concentrations (25, 50 and 100 µM); the maximum production was observed at 24 h. Remarkably, IL-1β secretion of the PA-treated microglia (409.47 (SD 54.54) pg/ml) was higher than that in LPS-treated cells (306.93 (SD 45.10) pg/ml) (Fig. 2(c)) at 24 h. Nonetheless, the levels of TNF-α and IL-6 were about 1- to 2-fold higher in LPS (TNF-α, 451.14 (SD 40.95) pg/ml; IL-6, 209.88 (SD 22.60) pg/ml) than those in the PA-treated microglia (TNF-α, 344.06 (SD 18.38) pg/ml; IL-6, 65.19 (SD 8.49) pg/ml) at 24 h.

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and LPS markedly increased H₂O₂ and O₂⁻ production in BV-2 cells compared with the control (Fig. 4(b)).

**Activation of microglia by palmitic acid treatment leads to bystander neuronal death**

Activated microglia are known to produce an array of cytokines and other inflammatory mediators that are in turn deleterious for surrounding neurons in the CNS. BV-2 cells were incubated in the absence or presence of PA (25–100 μM) for 12 h and the medium was changed with fresh serum-free DMEM. After 12 h, supernatant fractions were collected and filtered. To check whether PA-induced microglia activation causes bystander neuronal death, we treated primary neurons with the medium mentioned above. The control comprised culture supernatant fractions from BSA- and vehicle-treated BV-2 cells. Primary cortical neurons were incubated with microglia-conditioned medium for 2 d and then neuronal apoptosis was measured by morphological analysis. The results in Fig. 5 demonstrated a significant induction of apoptosis in the neurons. These results indicate that microglia produced inflammatory mediators in response to PA and that the mediators accumulated in the medium were capable of inducing neuronal death.

**SFA-activated NF-κB signalling**

NF-κB is an essential transcription factor for the expression of cytokine and iNOS expression in microglia. We therefore investigated the potential nuclear translocation of NF-κB following the stimulation of microglia with PA. For these experiments, BV-2 cells were treated with or without PA (25, 50 and 100 μM) for 12 h, and the p65 subunit of NF-κB in the nuclear fraction was assessed by using immunofluorescence. It was observed that either PA or LPS was capable of activating NF-κB, as demonstrated by the increased levels of the NF-κB subunit, p65, in the nucleus, whereas p65 was localised primarily in the cytosol during the resting state (Fig. 6(a)).

It has been demonstrated that phosphorylation of serine residues 529 and 536 of the RelA/p65 subunit leads to a transactivation of NF-κB. We next investigated whether PA regulates the phosphorylation of p65. The proteins harvested from the cells after 1 h treatment with or without PA (25, 50 and 100 μM) were processed for Western blot to detect intracellular levels of phospho-p65 (ser536). As shown in Fig. 6(b), phospho-p65 levels were significantly elevated in all three treatment groups.

In order to observe the effect of PA on the transcriptional activity of NF-κB, cells were transfected with a plasmid construct containing 3 × NF-κB binding sites associated with the
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NF-κB inhibitor suppressed SFA-induced pro-inflammatory cytokines and NO production

The role of NF-κB in PA-induced pro-inflammatory cytokines and NO production was examined using the specific NF-κB pathway inhibitor PDTC. In BV-2 cells treated with PDTC (100 μM), a non-toxic concentration, for 4 h, PA-induced gene expression of iNOS and pro-inflammatory cytokines was significantly suppressed (Fig. 7(a) and (b)). In addition, PDTC reduced PA-induced NO, TNF-α and IL-1β secretion (Fig. 7(c)). However, PDTC did not exert a significant effect on PA-induced IL-6 production (Fig. 7(c)). Taken together with the result shown in Fig. 6, these findings indicate that SFA are capable of inducing a rapid response of NF-κB in microglia, triggering the expression of cytokines (for example, TNF-α and IL-1β) and inflammatory mediators such as NO.

Anti-Toll-like receptor 4 antibody inhibited SFA-induced NF-κB activation and pro-inflammatory mediator production

We next investigated whether PA-induced activation of NF-κB was regulated via TLR4. Fig. 8(a) shows that at 1 h of PA treatment, the p65 translocation of NF-κB was significantly increased. Incubating the cells with anti-TLR4 Ab (10 μg/ml, a non-toxic concentration), however, prevented the PA-induced activation of NF-κB, suggesting an involvement of TLR4 in the activation of the transcriptional factors. Moreover, treatment of BV-2 cells with anti-TLR4 Ab inhibited PA-induced production of pro-inflammatory mediators (Fig. 8(b)).
The present results have shown that SFA can induce microglial activation as manifested by its actions on BV-2 cells and primary microglial cells. We have shown that SFA treatment induced microglial activation, as shown by changes in cell morphology consistent with a reactive phenotype, and caused significantly higher production of ROS, NO, and pro-inflammatory cytokines including TNF-α, IL-1β and IL-6 in microglia, resulting in bystander neuronal death. Moreover, PA treatment induced a marked expression of IL-1β and iNOS comparable with that with LPS. Additionally, we have shown that PA treatment activated NF-κB. It is striking that inhibition of NF-κB activation, with its inhibitor PDTC resulted in inhibition of iNOS, TNF-α, IL-1β and IL-6 mRNA expression, and production of TNF-α, IL-1β and NO except for IL-6. Another major finding was that in cells treated with anti-TLR4 Ab, PA-induced NF-κB activation and pro-inflammatory mediator production were repressed. These results suggest that SFA could activate microglia and stimulate the TLR4–NF-κB pathway to trigger the production of pro-inflammatory mediators, which may contribute to neuronal death.

Microglia adapt to different CNS environments and exhibit diverse morphological types and functional specialisations. Our findings have demonstrated that BV-2 cells and primary microglial cells exposed to SFA assumed an amoeboid morphology and increased CD11b expression, which are indicative of its activated state.

Discussion

It has been reported that the pro-inflammatory cytokines TNF-α, IL-1β and IL-6, which are important factors in the regulation of inflammatory processes, are overexpressed in the brain of AD patients, indicating the possible involvement of these cytokines in the pathology of the disease. Indeed, the present results have shown that PA and SFA increased TNF-α, IL-1β and IL-6 mRNA expression and cytokine secretion. Along with the pro-inflammatory cytokines, NO is an important contributor to neuronal damage in AD development. Following PA stimulation, there was a marked increase in NO production by BV-2 cells. Moreover, iNOS expression was also enhanced after PA treatment for 6 and 24 h. It is suggested that SFA treatment had induced iNOS up-regulation in microglia resulting in increased production of NO.

Like LPS, PA enhanced TNF-α, IL-6 and IL-1β production in microglia. However, it is remarkable that PA treatment had resulted in more vigorous IL-1β up-regulation. IL-1β is a critical inflammatory cytokine in AD, and found in activated microglia localised to amyloid plaques. It induces amyloid β deposition through directly up-regulating expression and processing of β-amyloid precursor proteins. It increases tau phosphorylation by mitogen-activated protein kinase p38. It activates astrocytes to overexpress S100 β, which stimulates neurite growth and increases Ca flux (a deadly event) in neurons. Moreover, it also stimulates astrocytes to produce additional pro-inflammatory cytokines such as IL-6. IL-1β has been reported to promote the
activation activity of the enzyme acetylcholinesterase, thus down-regulating the cholinergic system (36). Finally, IL-1 directly promotes microglial proliferation (37) and increases microglial expression of IL-1β and IL-6 (38). All these IL-1β-regulated processes might result in neuronal stress or injury, which in turn further enhances microglial activation and IL-1β overexpression. Thus IL-1β plays a pivotal role in the pathogenesis of AD. In addition, IL-1β is known to be involved in the expression and activation of iNOS (39). Interestingly, we have found that PA treatment enhanced a marked iNOS expression at a level comparable with that induced by LPS. It is suggested that PA directly activates microglia, and

Fig. 5. Bystander neuronal death caused by SFA-treated microglia. (a) BV-2 cells were incubated in the absence or presence of palmitic acid (PA; 25–100 µM) for 12 h. The medium was changed with fresh serum-free Dulbecco’s modified Eagle’s medium (DMEM) for 12 h. The supernatant fractions were collected, filtered and stored at −20°C. Primary cortical neurons were treated with these culture supernatant fractions for 48 h and stained with Hoechst 33342. **, Representative apoptotic nuclei (scale bar = 10 µm). Images are representative of triplicate sets. (b) Apoptotic nuclei were quantified in ten random fields for each experimental condition. Values are the means of three independent experiments, with standard deviations represented by vertical bars. Mean value was significantly different from that of the control: ** P < 0.01, *** P < 0.001.

Fig. 6. SFA induce NF-κB activation. (a) BV-2 cells were incubated in the absence or presence of palmitic acid (PA; 100 µM) or lipopolysaccharide (LPS; 500 ng/ml) for 1 h and stained for NF-κB p65 and counterstained with 4,6-diamidino-2-phenylindole (DAPI). Then images were captured by a fluorescence microscope. Scale bar = 20 µm. (b) BV-2 cells were incubated in the absence or presence of PA (25–100 µM) or LPS (500 ng/ml) for 1 h and 30 µg total protein were subjected to Western blot analysis. A phospho-specific antibody that recognises the phosphorylation of the serine 536 residue on the p65 (p-p65) determined the relative activation state of NF-κB. A non-phospho-specific antibody to NF-κB p65 (p65) served as protein loading controls. Images are representative of triplicate sets. (c) BV-2 cells were incubated in the absence or presence of PA (25–100 µM) or LPS (500 ng/ml) for 24 h. The effect of PA on NF-κB promoter activity was evaluated by luciferase assay as described in Materials and methods. Values are the means of three independent experiments, with standard deviations represented by vertical bars. Mean value was significantly different from that of the control: * P < 0.05, ** P < 0.01, *** P < 0.001.
increases IL-1β production, which may lead to enhanced iNOS expression. The above results indicate that IL-1β may play a key and detrimental role in regulating the SFA-induced inflammatory response.

Analysis of the regulation of inducible transcription factors has claimed a major role for the NF-κB system in the activation of microglial cells in neurodegenerative diseases. Activation of NF-κB has been linked to the up-regulation of potentially inflammation-related genes, including iNOS, cyclo-oxygenase-2, TNF-α, IL-1β and IL-6. Previous studies have demonstrated that SFA can induce NF-κB activation in macrophages (40). The present results have shown that PA enhanced p65 nuclear translocation, phosphorylation and NF-κB promoter activity. Furthermore, PDTC, a potent NF-κB inhibitor, suppressed PA-induced NO, TNF-α and IL-1β production in BV-2 cells, while PA-induced IL-6 production was not affected. It is therefore suggested that NO, TNF-α and IL-1β are downstream gene products of the NF-κB pathway induced by PA, whereas the effect of PA on IL-6 is independent of the NF-κB pathway. It is possible that other transcriptional factors may be involved in IL-6 production, for example, activating protein-1 (41), cAMP-induced transcription factors such as cAMP-responsive element binding protein and CCAAT-enhancer box binding protein (42,43), or signal transducer and activator of transcription (44). PA may be able to activate one of these transcriptional factors, increasing IL-6 production.

Several previous results have demonstrated that the stimulation of TLR4 by SFA can trigger transcription factor activation, leading to the production of pro-inflammatory mediators in monocytes and macrophages in vitro (45). However, recent studies have called into question the ability of
TLR to directly bind SFA as a ligand and suggest that SFA might modulate TLR activity via lipid raft changes or that SFA may induce inflammation via TLR-independent mechanisms. In the present study anti-TLR4 Ab reversed the PA-induced NF-κB p65 translocation and pro-inflammatory mediator production. Thus, it is hypothesised that TLR4 may mediate NF-κB activation in PA-induced microglia activation. On the other hand, the mechanism by which PA can activate TLR4 remains to be clarified. It is evident that the above-mentioned effects of SFA on BV-2 cells were not due to cytotoxicity, because no significant cell death was observed in the concentration ranges examined.

Against the above background, we then determined whether culture supernatant fractions derived from microglia treated with PA could actually affect neuronal survivability. Indeed, on addition of the supernatant fractions into primary neuronal culture, there was a significant reduction of neuronal viability. Thus, the combination of ROS, NO and pro-inflammatory cytokines seems to act in a synergistic way to cause bystander death to neurons.
In conclusion, the present study has shown that SFA can cause activation of microglial cells, resulting in the generation of pro-inflammatory cytokines, NO and ROS, which could, in turn, induce neuronal dysfunctions. PA treatment markedly increases the expression of IL-1β and iNOS to a level comparable with that with LPS. PA induced the TLR4-mediated activation of NF-κB, which is responsible for TNF-α, IL-1β and NO production. The possibility of PA activation of other pathways should be considered. This takes into consideration the fact that inhibition of NF-κB did not alter IL-6 production. The present novel findings suggest the potential mechanisms in SFA-induced microglial activation and, to this end, nutrition rich in SFA may be linked to some inflammatory diseases of the CNS.

Acknowledgements

The present study was supported by funding from the National Basic Research Program of China (973 Program, no. 2007CB512001, 2011CB966201); National Natural Science Foundation of China (no. 30771142, 81071057); Natural Science Foundation of Shandong Province (no. Z2007C11, J2008S2, ZR2010HQ022).

Z. W. and D. L. contributed equally to the present study. The authors’ contributions were: A. H. was involved in study design, data interpretation and manuscript editing; Z. W. and D. L. performed the majority of the laboratory work and contributed to the analysis of data and writing of the manuscript; F. W. and S. L. were responsible for the cell culture; S. Z. and E.-A. L. were involved in manuscript editing.

The authors have no conflict of interest to declare.

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