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Isolated Amyloid-β(1–42) Protod fibrils, But Not Isolated Fibrils, Are Robust Stimulators of Microglia

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ABSTRACT: Senile plaques composed of amyloid-β protein (Aβ) are an unshakable feature of the Alzheimer’s disease (AD) brain. Although there is significant debate on the role of the plaques in AD progression, there is little disagreement on their role in stimulating a robust inflammatory response within the context of the disease. Significant inflammatory markers such as activated microglia and cytokines are observed almost exclusively surrounding the plaques. However, recent evidence suggests that the plaque exterior may contain a measurable level of soluble Aβ aggregates. The observations that microglia activation in vivo is selectively stimulated by distinct Aβ deposits led us to examine what specific form of Aβ is the most effective proinflammatory mediator in vitro. We report here that soluble prefibrillar species of Aβ(1–42) were better than fibrils at inducing microglial tumor necrosis factor α (TNFα) production in either BV-2 and primary murine microglia. Reconstitution of Aβ(1–42) in NaOH followed by dilution into F-12 media and isolation with size exclusion chromatography (SEC) revealed classic curvilinear β-sheet protofibrils 100 nm in length. The protofibrils, but not monomers, markedly activated BV-2 microglia. Comparisons were also made between freshly isolated protofibrils and Aβ(1–42) fibrils prepared from SEC-purified monomer. Surprisingly, while isolated fibrils had a much higher level of thioflavin T fluorescence per mole, they were not effective at stimulating either primary or BV-2 murine microglia compared to protofibrils. Furthermore, SEC-isolated Aβ(1–40) protofibrils exhibited significantly less activity than concentration-matched Aβ(1–42). This report is the first to demonstrate microglial activation by SEC-purified protofibrils, and the overall findings indicate that small, soluble Aβ(1–42) protofibrils induce much greater microglial activation than mature insoluble fibrils.

KEYWORDS: Alzheimer’s disease, inflammation, amyloid-beta protein, tumor necrosis factor alpha, protofibrils, fibrils

Alzheimer’s disease (AD) is a complex neurodegenerative disease characterized by the accumulation of protein deposits in the affected brain and progressive dementia. The two classic forms of deposits are neurofibrillary tangles composed of tau protein and dense core neuritic plaques composed of amyloid-β protein (Aβ). Both lesions are believed to contribute to disease onset and progression although the initial event appears to be Aβ accumulation. Aβ is commonly produced as an unstructured 40- or 42-residue peptide fragment by proteolytic cleavage of the amyloid-β precursor protein. The monomeric form of Aβ circulates ubiquitously in plasma and cerebrospinal fluid yet an aggregated fibrillar form comprises the characteristic Aβ plaques. The mechanistic complexity of AD is increased by the growing number of reports demonstrating a variety of Aβ structures morphologically distinct from plaques that possess greater solubility and neuronal toxicity.

It has been well documented that inflammatory markers such as activated microglia stained with proinflammatory cytokines have been observed surrounding the neuritic Aβ plaques in the human AD brain. In fact, studies in an AD transgenic mouse model have shown rapid microglial accumulation around newly formed plaques. A chronic inflammatory state induced by accumulated Aβ has been suggested as one of the underlying mechanisms of progressive neurodegeneration in AD and may in fact exacerbate Aβ deposition. Multiple studies suggest that small Aβ(1–42) oligomers may cause early and significant alterations in synaptic function and then as fibrillar structures are produced, concomitant inflammatory responses appear (reviewed in ref 4).

In vitro Aβ aggregation studies have contributed significantly to the understanding of fibrillogenesis mechanisms and the structural properties of monomers, soluble intermediates, and mature fibrils. These studies have identified a continuum of Aβ species in the assembly process which vary in their size, length, solubility, and morphology. In solution, Aβ monomers will undergo noncovalent self-assembly to form soluble oligomers, protofibrils that are enriched in β-sheet structure, and insoluble fibrils. The types of intermediates formed during fibrillogenesis are dependent on the solution conditions.

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In addition to significant Aβ polymorphism exhibited in the AD brain, the structural diversity may even extend to the Aβ senile plaques based on recent observations indicating a significant level of oligomeric Aβ surrounding the plaques in an AD mouse model.18–21 In concert with in vitro aggregation studies, cellular studies have shown that monomeric, oligomeric, protofibrillar, fibrillar, and amorphous Aβ species possess distinct toxic and biological activities and potencies.16,19–22 This is also the case with Aβ as a proinflammatory stimulus. It is generally felt that soluble Aβ oligomers and protofibrils contribute to early dendritic and synaptic injury in AD models (reviewed in ref 23) while fibrillar Aβ acts as a proinflammatory stimulus. However, many groups have reported induction of proinflammatory cytokines and signaling pathways by soluble prefibrillar species in glial cells. In some cases, the type and extent of proinflammatory response differed between Aβ fibrils and soluble oligomeric species.24,25

The prevailing view is still that while small soluble Aβ species display greater cellular toxicity, fibrillar structures are the primary mediators of inflammation. Our own studies have indicated that soluble Aβ(1–42) fibrillar precursors were much more effective than fibrils at stimulating tumor necrosis factor α (TNFα) production in human monocytes.26 In many reports, Aβ solutions were used that likely contained a high degree of polydispersity with respect to size and aggregation state. In this study, multiple aggregation conditions were evaluated in conjunction with separation and rigorous characterization of the isolates in order to identify the optimal Aβ species for microglia activation.

RESULTS AND DISCUSSION

Microglial TNFα Production Induced by Aβ(1–42) Prepared in Oligomer-Forming and Fibril-Forming Conditions. LaDu and colleagues previously compared the inflammatory response in rat astrocyte cultures induced by Aβ incubated in oligomer- or fibril-forming conditions.24 We utilized their preparation methods and conditions to prepare Aβ(1–42) solutions that contain species of distinct morphologies for evaluation of the inflammatory response in isolated murine microglia. The preparation of the Aβ species entailed reconstitution in DMSO followed by dilution into either cold F12 media (oligomer-forming) or 10 mM HCl (fibril-forming). A 24 h incubation of the Aβ(1–42) solutions at either 4 °C (oligomer-forming) or 37 °C (fibril-forming) produced the anticipated morphologies for both conditions as assessed by atomic force microscopy (AFM) (Figure 1A). The images were similar to those of Stine et al.14 Small punctate globular species were observed in the oligomer preparations (heights 3.1 ± 1.2 nm for n = 100 measurements) while long flexible fibers (heights 2.7 ± 0.9 nm for n = 100 measurements) were observed in the fibril preparations. ThT binding and fluorescence levels were significantly higher in the solutions containing fibrils compared to those containing oligomers (Figure 1B). This was consistent with the general understanding of fibrils as they are expected to have increased β-sheet character and a higher number of ThT binding sites. However, the observation that the oligomer preparations exhibited a considerable level of ThT fluorescence was somewhat surprising and suggested a significant level of β-structure.

The Aβ(1–42) oligomer and fibril preparations were evaluated for their ability to stimulate TNFα production by microglia. TNFα is an important product of the MyD88-dependent proinflammatory innate immune response35 and is measurably increased in post-mortem AD brain sections,10 and microvessels36 and cerebrospinal fluid37 of clinically diagnosed AD patients. Although most studies show similar responses between BV-2 and primary microglia, concerns have been raised that in some cases immortalized BV-2 microglia may not fully model primary microglia38 and caution should be exercised when interpreting BV-2 results until direct comparison with primary cells can be made.39 Therefore, it has become commonplace to utilize both microglial cell types for experimental testing which was the case in these studies. A 6 h incubation of 15 μM Aβ(1–42) oligomers or fibrils with murine microglia revealed significant differences in TNFα production between the two Aβ aggregation states (Figure 2). Similar trends of inflammatory activity were observed in both primary (Figure 2A) and BV-2 (Figure 2B) microglia. Despite the lesser ThT fluorescence, the Aβ(1–42) solutions prepared and incubated in oligomer-forming conditions were markedly better at inducing TNFα from murine microglia consistent with previous findings in rat astrocytes.24

Characterization of SEC-Isolated Aβ(1–42) Prototifibrils Prepared and Eluted in Supplemented F12. Regardless of the conditions used to study Aβ(1–42) aggregation there is always a degree of polydispersity present in the solutions which likely include monomers, soluble aggregates of many sizes, and some fibrillar structures. In order to better define and characterize the soluble Aβ(1–42) aggregation species, SEC was used for separation and purification. SEC has been used in numerous studies to separate prototifibrils from monomeric Aβ.11,40 In this

Figure 1. Aβ(1–42) structures formed in oligomer- or fibril-forming conditions are morphologically distinct. Aliquots of lyophyllized Aβ(1–42) were reconstituted to 100 μM as described in the Methods to generate oligomers or fibrils. After 24 h incubation of the given temperatures, aliquots of the solutions were diluted to 10 μM, applied to mica, and imaged by AFM. Representative images are shown for oligomers (panel A, 5 μm × 5 μm) and fibrils (panel B, 10 μm × 10 μm). ThT fluorescence measurements (n = 8 for oligomers and n = 10 for fibrils) were obtained for each preparation at a final concentration of 10 μM Aβ(1–42) (Panel C).
set of experiments, a Superdex 75 column was used for separation. Using a modified version of the NaOH method, lyophilized Aβ(1–42) was reconstituted in NaOH followed by dilution in antibiotic-supplemented F-12 to prevent microbial growth. The use of F-12 media was retained from the oligomer-forming conditions in Figure 1 due to the physiological nature of the solution and the compatibility with microglial cell treatment. The Aβ(1–42) solution (250 μM) was centrifuged at 18 000g and the supernatant was eluted in supplemented F-12 media as described in the Methods. The Aβ(1–42) elution profile was monitored by UV absorbance (280 nm) and exhibited characteristic protofibril (void) and monomer (included) peaks (Figure 3).

The significant Aβ(1–42) void peak was observed without prolonged incubation of the reconstitution solution confirming that Aβ(1–42) forms protofibrils rapidly under these conditions. SEC-purified monomer fractions did not exhibit ThT fluorescence while protofibril fractions at equivalent concentrations showed significant fluorescence (Figure 4A). Although F-12 media is not a common Superdex 75 elution buffer such as Tris or PBS, transmission electron microscopy (TEM) imaging of the F-12-eluted void peak showed classic short curvilinear protofibrils < 100 nm in length (Figure 4B). Dynamic light scattering (DLS) measurements of the protofibril peak from seven separate SEC purifications produced an average R_h value of 21 nm with a standard deviation (std dev) of 6 nm (data not shown). Deconvolution of the average R_h values into histograms by data regularization revealed two predominant peaks of 4.5 ± 0.9 nm and 20.6 ± 6.5 nm. Peak 1 with the smaller R_h value was observed in 4 of 7 protofibril isolations, while peak 2 was always observed. Even within isolated Aβ(1–42) protofibrils, a degree of polydispersity was present as the peak 2 histogram widths varied from narrow (e.g., 15–20 nm) to broad (e.g., 8–41 nm) in different experiments.

Microglial TNFα Production by SEC-Isoalted Aβ(1–42) Protofibrils, But Not Monomer. BV-2 murine microglia were treated with fresh SEC-isolated protofibril and monomer fractions and examined for their ability to stimulate an inflammatory response. Fraction concentrations were calculated based on UV absorbance and were used to determine final cellular treatment concentrations. Protofibrils (15 μM) induced a substantial level of secreted TNFα while monomers (15 μM) were much less effective (Figure 5). The presence of contaminating lipopolysaccharide (LPS) in SEC fractions was assessed in two ways, (1) a cell-free XTT assay in which the presence of any bacteria in the Aβ/ samples would be expected to catalyze reduction of XTT and (2) TNFα production was monitored in the absence or presence of polymyxin B (PMX-B), a neutralizer of LPS signaling. The former method revealed no XTT reduction in the presence of Aβ/ samples and in the latter method it was observed that PMX-B (100 ng/mL) had no effect on the Aβ/ response yet blocked >99% of the LPS (3 ng/mL) response (data not shown). Thus, neither method showed any indication of bacterial or LPS presence. The findings demonstrated that isolated Aβ(1–42) protofibrils were significant stimulators of BV-2 microglia whereas little stimulation was observed with purified monomeric Aβ(1–42). The response to Aβ(1–42) protofibrils was also tested in primary microglia isolated from newborn (3–4 day old) C57BL/6 mouse pups. Freshly isolated protofibrils were tested at multiple concentrations (5, 10, 15, and 20 μM) and exhibited a dose-dependent ability to induce TNFα production in both primary and BV-2 microglia (Figure 6). Typically primary microglia were more responsive to Aβ(1–42)

Figure 3. Dual peaks are observed following SEC elution of Aβ(1–42) reconstituted in NaOH/F-12. Lyophilization Aβ(1–42) (1 mg) was brought into solution with NaOH followed by supplemented F-12 media to a final concentration of 200 μM. The supernatant after centrifugation was eluted from a Superdex 75 column, and 0.5 mL fractions were collected. UV absorbance at 280 nm was monitored during the elution (solid line).

Figure 4. Structure and morphology of Aβ(1–42) protofibrils. (A) Freshly isolated Aβ(1–42) protofibrils and monomers after elution from Superdex 75 in supplemented F-12 were diluted to 5 μM respectively in supplemented F-12 containing 5 μM ThT, and fluorescence emission was measured as described in the Methods. (B) Protofibrils were diluted to 20 μM, applied to a copper formvar grid, and imaged by TEM at a magnification of 43 000x. The scale bar represents 100 nm.
isolated by centrifugation, supernatant removal, and resuspension subjected to gentle agitation for approximately 72 h. Fibrils were stimulated by Aβ.

The current report demonstrates the substantial activation of microglia by soluble Aβ(1–42) aggregates. We observed of the pellet as described in the Methods. In all cases, >98% of the ThT fluorescence of the total solution was removed from the supernatant after centrifugation. ThT fluorescence comparisons of equivalent concentrations of Aβ(1–42) fibrils and SEC-purified Aβ(1–42) protofibrils and monomers immediately following elution showed fibrils with by far the best ThT binding ability (Figure 7A). Protofibrils displayed significantly lower ThT fluorescence and monomers were at background levels. TEM images of the isolated Aβ(1–42) fibrils revealed long fibers with lengths exceeding 1 μm and typical widths of 5–10 nm (Figure 7B). The neutral pH and higher ionic strength of the F-12 media encouraged a considerable degree of lateral association between the fibrils. Surprisingly, fibrils were very poor in their ability to stimulate TNFα production in primary (Figure 7C) and BV-2 (data not shown) microglia. Aβ(1–42) protofibrils invoked a dramatically higher microglial response compared to fibrils and monomers (Figure 7C). A cell-free XTT cell proliferation assay again showed no evidence of contamination in any of the preparations (data not shown). Our typical BV-2 and primary microglial preparations contained cells with both round and ramified morphologies with defined boundaries. After exposure to Aβ, the cells became clustered with rough boundaries and a less ramified morphology. These observations are consistent with those made by Garcia et al in rat microglia.

Primary microglia underwent more pronounced alterations in morphology compared to BV-2 cells. Even with these exterior changes to the cells after treatment with Aβ, no significant toxicity was observed using an XTT cell viability assay. Exposure of the microglia to Aβ(1–42) fibrils, protofibrils, or monomer for 6 h did not inhibit mitochondrial-mediated reduction of XTT in either BV-2 or primary microglia (data not shown). The preparation of Aβ protofibrils, fibrils, and monomers in the supplemented F-12, which is similar in pH and ionic strength to the microglial cell culture medium, likely helped prevent dramatic structural changes in the isolated Aβ species when introduced to the microglia cells. Although this was not verified, very different microglial responses were clearly able to be observed between the distinct Aβ species.

Figure 5. Protofibrils are significant stimulators of BV-2 microglia. SEC-isolated Aβ(1–42) protofibrils and monomers in supplemented F-12 were incubated with BV-2 microglia at a final concentration of 15 μM for 6 h in medium containing 2% FBS. Secreted TNFα was measured by ELISA in the conditioned medium. Data bars represent the average ± std error of n = 6 trials. Control treatments with an equal volume of supplemented F-12 media produced 23 pg/mL TNFα and were subtracted from Aβ-stimulated samples.

Figure 6. Protofibrils display a dose-dependent effect in their ability to induce TNFα production in microglia. SEC-isolated Aβ(1–42) protofibrils in supplemented F-12 were incubated with primary microglia (panel A) or BV-2 microglia (panel B) for 6 h at final concentrations of 5, 10, 15, and 20 μM. Secreted TNFα was measured by ELISA in the conditioned medium. Data bars represent the average ± std error of n = 4–6 trials at each concentration for primary microglia and n = 6 trials at each concentration for BV-2 microglia. Control treatments with supplemented F-12 media produced 6 and 17 pg/mL TNFα respectively for primary and BV-2 microglia and were subtracted from Aβ-stimulated samples. Primary and BV-2 microglia were stimulated by Aβ in serum-free conditions.

Comparison of Microglial TNFα Production between Aβ(1–42) Protofibrils and Aβ(1–40) Protofibrils. In order to compare Aβ(1–40) protofibrils with those formed from Aβ(1–42), a longer incubation was needed after NaOH reconstitution and dilution into supplemented F-12 medium. While Aβ(1–42) protofibrils formed rapidly within minutes, Aβ(1–40) required a 24 h incubation at 25 °C before significant amounts of protofibrils were generated (Figure 8). The Aβ(1–40) protofibril fraction was assessed by TEM which revealed similar structures as observed for Aβ(1–42) although the lengths appeared somewhat smaller (Figure 9A). Manual measurements of the images confirmed that the Aβ(1–40) and Aβ(1–42) protofibril fractions had statistically different (p < 0.001) lengths as Aβ(1–40) protofibrils averaged 27 ± 11 nm while Aβ(1–42) protofibrils averaged 50 ± 16 nm for n = 50 measurements. The length analysis only included unambiguous protofibril structures and excluded the smaller species (<15 nm).

Comparison of Microglial TNFα Production between Aβ(1–42) Protofibrils and Aβ(1–40) Protofibrils. SEC-purified Aβ(1–42) monomer in supplemented F-12 was moved to 25 °C and subjected to gentle agitation for approximately 72 h. Fibrils were isolated by centrifugation, supernatant removal, and resuspension of the pellet as described in the Methods. In all cases, >98% of the ThT fluorescence of the total solution was removed from the supernatant after centrifugation. ThT fluorescence comparisons of equivalent concentrations of Aβ(1–42) fibrils and SEC-purified Aβ(1–42) protofibrils and monomers immediately following elution showed fibrils with by far the best ThT binding ability (Figure 7A). Protofibrils displayed significantly lower ThT fluorescence and monomers were at background levels. TEM images of the isolated Aβ(1–42) fibrils revealed long fibers with lengths exceeding 1 μm and typical widths of 5–10 nm (Figure 7B). The neutral pH and higher ionic strength of the F-12 media encouraged a considerable degree of lateral association between the fibrils. Surprisingly, fibrils were very poor in their ability to stimulate TNFα production in primary (Figure 7C) and BV-2 (data not shown) microglia. Aβ(1–42) protofibrils invoked a dramatically higher microglial response compared to fibrils and monomers (Figure 7C). A cell-free XTT cell proliferation assay again showed no evidence of contamination in any of the preparations (data not shown).

Our typical BV-2 and primary microglial preparations contained cells with both round and ramified morphologies with defined boundaries. After exposure to Aβ, the cells became clustered with rough boundaries and a less ramified morphology. These observations are consistent with those made by Garcia et al in rat microglia. Primary microglia underwent more pronounced alterations in morphology compared to BV-2 cells. Even with these exterior changes to the cells after treatment with Aβ, no significant toxicity was observed using an XTT cell viability assay. Exposure of the microglia to Aβ(1–42) fibrils, protofibrils, or monomer for 6 h did not inhibit mitochondrial-mediated reduction of XTT in either BV-2 or primary microglia (data not shown). The preparation of Aβ protofibrils, fibrils, and monomers in the supplemented F-12, which is similar in pH and ionic strength to the microglial cell culture medium, likely helped prevent dramatic structural changes in the isolated Aβ species when introduced to the microglia cells. Although this was not verified, very different microglial responses were clearly able to be observed between the distinct Aβ species.

Comparison of Microglial TNFα Production between Aβ(1–42) Protofibrils and Aβ(1–40) Protofibrils. In order to compare Aβ(1–40) protofibrils with those formed from Aβ(1–42), a longer incubation was needed after NaOH reconstitution and dilution into supplemented F-12 medium. While Aβ(1–42) protofibrils formed rapidly within minutes, Aβ(1–40) required a 24 h incubation at 25 °C before significant amounts of protofibrils were generated (Figure 8). The Aβ(1–40) protofibril fraction was assessed by TEM which revealed similar structures as observed for Aβ(1–42) although the lengths appeared somewhat smaller (Figure 9A). Manual measurements of the images confirmed that the Aβ(1–40) and Aβ(1–42) protofibril fractions had statistically different (p < 0.001) lengths as Aβ(1–40) protofibrils averaged 27 ± 11 nm while Aβ(1–42) protofibrils averaged 50 ± 16 nm for n = 50 measurements. The length analysis only included unambiguous protofibril structures and excluded the smaller species (<15 nm).

Despite morphological similarities, cellular studies demonstrated that Aβ(1–40) protofibrils were much less effective in stimulating TNFα in primary microglia compared to Aβ(1–42) protofibrils (Figure 9B). The concentration used in the microglial treatment for both Aβ(1–40) and Aβ(1–42) protofibrils was 15 μM.

The current report demonstrates the substantial activation of murine microglia by soluble Aβ(1–42) aggregates. We observed
from Aβ final concentration of 15 μM to 20 μM containing 5 μM ThT and fluorescence emission was measured and plotted as described in the Methods. Fibers were isolated as described after gentle agitation for 72 h at 25 °C. One fluorescence measurement was obtained from each solution. (B) TEM images of isolated Aβ(1–42) fibril pellets (74 μM) at a magnification of 25,000. (C) Aliquots of the solutions described in panel (A) were incubated with primary murine microglia for 6 h at a final concentration of 15 μM Aβ(1–42). Secreted TNFα was measured by ELISA in the conditioned medium. Error bars represent the average ± std error of n = 6. Control treatments with supplemented F-12 media produced 15 pg/mL TNFα for primary microglia and were subtracted from Aβ-stimulated samples.

Figure 7. Protofibrils display less ThT fluorescence but stimulate microglia more effectively than fibrils. (A) Aliquots of SEC-purified Aβ(1–42) protofibrils (PF), monomer (M), and Aβ(1–42) fibrils formed from purified monomer (F) were diluted to 5 μM in 50 mM Tris-HCl pH 8.0 containing 5 μM ThT and fluorescence emission was measured and plotted as described in the Methods. Fibers were isolated as described after gentle agitation for 72 h at 25 °C. One fluorescence measurement was obtained from each solution. (B) TEM images of isolated Aβ(1–42) fibril pellets (74 μM) at a magnification of 25,000. (C) Aliquots of the solutions described in panel (A) were incubated with primary murine microglia for 6 h at a final concentration of 15 μM Aβ(1–42). Secreted TNFα was measured by ELISA in the conditioned medium. Error bars represent the average ± std error of n = 6. Control treatments with supplemented F-12 media produced 15 pg/mL TNFα for primary microglia and were subtracted from Aβ-stimulated samples.

Figure 8. Aβ(1–40) protofibrils require longer incubation for formation. Aβ(1–40) was reconstituted and prepared for SEC as described in the Methods. Aβ(1–40) solutions were eluted on Superdex 75 immediately (dashed line) or after a 24 h incubation at 25 °C (solid line). Fractions containing protofibrils and monomers were immediately placed on ice for further characterization. Concentrations were determined by UV absorbance.

Figure 9. Aβ(1–40) protofibrils do not stimulate microglia as well as Aβ(1–42) protofibrils. (A) TEM image of a sample taken from the Aβ(1–40) protofibril peak in Figure 8 (solid line). Sample was diluted to 20 μM Aβ, applied to a copper formwar grid as described in the Methods, and imaged at a magnification of 43,000×. The scale bar represents 100 nm. (B) Primary microglia were incubated with Aβ(1–40) or Aβ(1–42) protofibrils isolated by SEC in supplemented F-12 at a final concentration of 15 μM each for 6 h. Secreted TNFα was measured by ELISA in the conditioned medium. Data bars represent the average ± std error of n = 12 (Aβ(40)) and n = 9 (Aβ(42)) trials over two separate protofibril preparations for each protofibril type. Protofibrils were directly compared within each cell treatment experiment. Control treatments with an equal volume of supplemented F-12 media produced 31 pg/mL TNFα and were subtracted from Aβ-stimulated samples.

that an enriched fraction of SEC-isolated protofibrils elicited the greatest production of the proinflammatory cytokine TNFα compared to monomers and fibrils. The current results have similarities to our previous findings in THP-1 human monocytes in which soluble fibrillar precursors were optimal for inducing a proinflammatory response.26 While many studies have shown that monocytes and microglia have similar responses to Aβ, microglial cells provide a more relevant model system for investigating Aβ neuroinflammatory mechanisms. Furthermore, this report is the first to provide a direct comparison between SEC-purified Aβ(1–42) protofibrils and isolated Aβ(1–42) fibrils prepared from SEC-purified monomer in their ability to induce microglial activation.

Figure 10. Aβ(1–40) protofibrils do not stimulate microglia as well as Aβ(1–42) protofibrils. (A) TEM image of a sample taken from the Aβ(1–40) protofibril peak in Figure 8 (solid line). Sample was diluted to 20 μM Aβ, applied to a copper formwar grid as described in the Methods, and imaged at a magnification of 43,000×. The scale bar represents 100 nm. (B) Primary microglia were incubated with Aβ(1–40) or Aβ(1–42) protofibrils isolated by SEC in supplemented F-12 at a final concentration of 15 μM each for 6 h. Secreted TNFα was measured by ELISA in the conditioned medium. Data bars represent the average ± std error of n = 12 (Aβ(40)) and n = 9 (Aβ(42)) trials over two separate protofibril preparations for each protofibril type. Protofibrils were directly compared within each cell treatment experiment. Control treatments with an equal volume of supplemented F-12 media produced 31 pg/mL TNFα and were subtracted from Aβ-stimulated samples.
regularization histograms suggests that further separation and characterization may be possible. Structurally, protofibrils have similarities to fibrils based on thioflavin T binding, circular dichroism, and hydrogen exchange, but have not yet developed the full stability of fibrillar Aβ. Protop fibril diameters are typically smaller than fibrils and range from 4 to 6 nm although the proto fibril to fibril transition can occur without a change in diameter. The transition to fibrils can occur via mechanisms that incorporate monomer deposition on protofibril ends, end-to-end annealing of protofibrils, and lateral association of protofibrils. Protofibrils display toxicity to neurons, disrupt ion channels, inhibit hippocampal long-term potentiation, and are likely to possess other detrimental biological activities. Their solubility and diffusible nature quite possibly render them more effective in cellular interactions and engaging microglial receptors compared to mature insoluble fibrils.

The mechanism by which Aβ fibrils evoke a proinflammatory response appears to involve multiple cell-surface receptors. A multireceptor complex comprising the SR-B receptor CD36, αβ-integrin, and the integrin-associated protein CD47 has been shown to mediate fibrillar Aβ initiation of murine microglial activation. Furthermore, receptor components of the innate immune system including CD14, toll-like receptor (TLR) 2, and TLR 4 have also been identified as proinflammatory-linked receptors in microglial cells for Aβ fibrils. Moreover, a TLR4-TLR6-CD36 complex was recently shown to mediate cellular inflammatory responses to Aβ(1–42) fibrils. A cell-surface receptor that is selective for Aβ protofibrils has not been identified although the known fibril receptors may recognize similar structural elements between the two species.

For many years, it has been accepted that fibrillar Aβ is the primary trigger for inducing a glial inflammatory response in AD and in cell culture models. Many studies established and utilized solution conditions that were optimal for Aβ fibril formation but may have contained other Aβ species. Nevertheless, it was generally interpreted as microglial activation by Aβ fibrils. Numerous studies have demonstrated cytokine production by fibrillar Aβ interaction of fibrillar Aβ with proinflammatory receptors, and initiation of proinflammatory signaling pathways by fibrillar Aβ. More recently, studies have shown that a fibrillar state for Aβ may not be required for microglial activation. These findings have been based on a lack of correlation between ThT binding and cell stimulation or modulation of aggregation solution conditions to encourage formation of oligomers that display significant proinflammatory activity in microglia. A recent study altered the Aβ aggregation solution conditions to encourage formation of smaller or larger oligomers and fibrils. Stimulation of primary mouse microglia with these solutions revealed that, while polydispersity in Aβ species, those solutions containing smaller oligomers were able to induce more TNFα mRNA than those containing larger oligomers and fibrils. Less has been published regarding Aβ protofibrils and their role in AD-linked inflammation. In one study, Parvathy et al. found that Aβ protofibrils were not effective at inducing IL-1α mRNA expression in primary murine microglia compared to oligomers and fibrils. This finding indicates that perhaps not all microglial protofibrillar products are upregulated to the same extent by Aβ protofibrils. Interestingly, a new report suggests that soluble Aβ aggregates may be one of the targets of the central nervous system glial cell response which includes both microglia and astrocytes. Da Rocha-Souto et al. demonstrated in double-transgenic APPsw-tau mice that the correlation between oligomeric Aβ burden in the brain (based on NAB61 antibody binding) and the number of reactive astrocytes was better than the correlation between total Aβ plaque burden and reactive astrocytes.

Given the polydispersity observed in Aβ aggregation, there is a good possibility that many preparations are not homogeneous. In fact, many studies focusing on a particular biological activity of Aβ utilize Aβ solutions that have not been rigorously characterized. These solutions likely contain mixtures of Aβ aggregated species including oligomers, protofibrils, and fibrils. The conditions can dictate which species is more highly populated, but the mixture prevents the identification of the most active species. In fact, the Aβ(1–42) solutions prepared in fibril-forming conditions (Figures 1 and 2) may have contained a small population of protofibrils which might help to explain why a moderate amount of microglial TNFα production was observed. The formation of these fibrils at very low ionic strength precluded the ability to utilize centrifugation to separate the fibrils from any contaminating protofibrils. However, the Aβ(1–42) fibrils prepared from SEC-purified monomer at physiological pH and ionic strength (Figure 7) were very poor at stimulating TNFα production from microglia. This observation was possible due to the additional isolation step whereby fibrils were separated from the remaining Aβ solution by centrifugation and resuspended before application to the cells. This procedure likely removed any soluble aggregated species that may have been present in the initial solution thus allowing isolated fibrils to be evaluated. Looking back on previous studies, the persistent use of Aβ solutions containing a polydispersity population of aggregated species when evaluating Aβ biological activity may help to explain significant fluctuations in cell response and irreproducibility between experiments. Furthermore, the Aβ(1–42) fibrils described in Figures 1 and 2 were prepared under nonphysiological solution conditions (e.g., 10 mM HCl), and while the acidic conditions may have enhanced fibril formation, the internal structure may be quite different from the fibrils formed under more physiological conditions in Figure 7. Petkova et al. found that parallel β-sheet Aβ (1–40) fibrils prepared under different conditions had distinct hydrogen-bond registries and marked differences in neuronal toxicity. The same phenomenon has been observed for soluble aggregates. Chiti and co-workers demonstrated that oligomers grown under different solution conditions can have identical morphologies yet show different internal structures and toxicities. There remain potential concerns with in vitro preparation of aggregated Aβ species for cellular studies as their structure may have important differences from those formed in vivo. In the current study, the preparation, isolation, and characterization of Aβ(1–42) protofibrils under close to physiological solution conditions may help avoid some of the concerns. The significant and consistent microglial response to an enriched SEC fraction of Aβ(1–42) protofibrils demonstrates that these soluble fibrillar precursors are potent proinflammatory mediators. The much weaker microglial response to an equal concentration of Aβ(1–40) protofibrils suggests that even though the two protofibril species appear morphologically similar, there are distinct properties between the two types of protofibrils that influence their inflammatory activity.

**METHODS**

**Cell Culture and Primary Microglia Isolation.** BV-2 cells are primary mouse microglial cells immortalized by stable transfection with the c-myc oncogene and are functionally identical to native primary microglia. BV-2 cells were provided by Dr. Gary Landreth,
Case Western Reserve University, and were maintained in Dulbecco’s modified Eagle’s medium (DMEM, 4.5 g/L glucose) (HyClone) containing 50 U/mL penicillin, 50 μg/mL streptomycin, 50 μM β-mercaptoethanol, and 5% fetal bovine serum (FBS, HyClone). BV-2 cells were used for their ease of culture, initial observations, optimization of microglia/Aβ interactions, and to reduce the usage of mouse for primary cells. Primary murine microglia were harvested from newborn C57BL/6 mouse pups as previously described.32 Care and breeding of the C57BL/6 parent mice (Harlan Laboratories) was done at the University of Missouri—St. Louis Animal Facility. Briefly, 3–4 day old mouse pups were euthanized with an overdose of inhaled isoflurane (Fisher Scientific). The brains were isolated and meninges were removed under sterile conditions. Brain tissue was minced using sharp edged forceps, resuspended in 0.5% trypsin (Hyclone), and incubated at 37 °C for 20 min to allow further dissociation of the tissue. Subsequently, tissue was resuspended in complete DMEM containing 10% fetal bovine serum, 4 mM L-glutamine, 100 U/mL penicillin, 0.1 mg/mL streptomycin, and 0.25 μg/mL amphotericin-B (Fisher Scientific), OPI medium supplement (oxalocetate, pyruvate, insulin, Sigma-Aldrich), and 0.5 mg/mL recombinant mouse granulocyte-macrophage colony-stimulating factor (GM-CSF) (Invitrogen). The cell suspension was further triturated using a pipet and filtered through a 70 μm cell strainer to remove debris. The resulting cell suspension was centrifuged at 200g for 5 min at 25 °C, resuspended in complete medium, and seeded into 150 cm² flasks (Corning). Cells were cultured at 37 °C in 5% CO₂ until confluent (1–2 weeks), and microglia were selectively harvested from the adherent astrocyte layer by overnight shaking of the flask at 37 °C in 5% CO₂ and collection of the medium. The flask were replenished with fresh medium and then incubated further to obtain additional microglia. Typically, this procedure was repeated 3–4 times for one flask without removal of the astrocyte layer.

**Cell Stimulation Assay.** For cellular studies, BV-2 microglia were removed from culture flasks with 0.25% trypsin and seeded in a sterile 96-well cell culture plate overnight at a density of 5 × 10⁵ cells/mL in growth medium described above. Prior to cell treatment, medium was replaced with fresh medium containing either 0 or 2% FBS followed by Aβ stimulation at a final concentration of 15 μM. The inclusion of 2% FBS initially was adapted from an established THP-1 monocyte protocol.30 Treatment of BV-2 without serum has been reported elsewhere31 and the absence of serum during BV-2 cell stimulation did not dramatically alter the results. The cells were incubated at 37 °C for 6 h in 5% CO₂ and the medium was collected and stored at −20 °C for subsequent analysis by enzyme-linked immunosorobent assay (ELISA). Primary microglia were collected after shaking flasks overnight and then centrifuged at 200g for 10 min at 25 °C. The cells were resuspended in complete microglial medium without GM-CSF or FBS and plated at a density of 5 × 10⁵ cells/mL in a sterile 96-well cell culture plate for either 2 h or overnight. The longer preincubation, which has been used previously32 and was done to allow greater adherence of the microglia, had no deleterious effects on the cells. Prior to treatment, the medium was replaced again and lipopolysaccharide (3 ng/mL) or Aβ (15 μM) was added to the cells. The medium was collected after 6 h and stored at −20 °C for subsequent determination of secreted TNFα levels by ELISA. The background cellular response was assessed using the particular buffer vehicle for the Aβ.

**ELISA.** Measurement of secreted TNFα in the supernatants was determined by ELISA as previously detailed.32 Briefly, 96-well plates were coated overnight with monoclonal antinmouse TNFα capture antibody, washed with phosphate-buffered saline (PBS) containing 0.05% Tween-20, and blocked with PBS containing 1% BSA, 5% sucrose, and 0.05% NaN₃ following by a wash step. Successive treatments with washing in between were done with samples or standards, biotinylated polyclonal antinmouse TNFα detection antibody in 20 mM Tris with 150 mM NaCl and 0.1% BSA, streptavidin-horseradish peroxidase (HRP) conjugate, and equal volumes of HRP substrates 3,3′,5,5′-tetramethylbenzidine and hydrogen peroxide. The reaction was stopped by the addition of 1% H₂SO₄ solution. The optical density of each sample was analyzed at 450 nm with a reference reading at 630 nm using a SpectraMax 340 absorbance plate reader (Molecular Devices, Union City, CA). The concentration of TNFα in the experimental samples was calculated from a mouse TNFα standard curve of 15–2000 pg/mL. When necessary, samples were diluted to fall within the standard curve. TNFα concentrations for absorbance values below the lowest 15 pg/mL standard were determined by extrapolation of the standard curve regression line.

**Preparation of Aβ Peptides.** Aβ(1–42) was obtained from W.M. Keene in the Abdominal Research Laboratory (Yale School of Medicine, New Haven, CT) in lyophilized form and stored at −20 °C. Aβ(1–40) was obtained from rPeptide (Bogarth, GA) or prepared by solid phase synthesis in the Structural Biology Core at the University of Missouri—Columbia as described previously.33 Aβ peptides were typically dissolved in 100% hexafluoroisopropanol (HFIP) (Sigma-Aldrich, St. Louis, MO) at 1 mM, aliquoted into sterile microcentrifuge tubes, and evaporated uncovered at room temperature overnight in a fume hood. The following day, the aliquots were vacuum-centrifuged to remove any residual HFIP and stored in desiccant at −20 °C. Some Aβ peptides were treated with 100% trifluoroacetic acid and vacuum-centrifuged prior to HFIP treatment. Aβ oligomers and fibrils obtained directly from lyophilized aliquots were prepared as previously described.14 Briefly, lyophilized Aβ42 aliquots were resuspended in sterile anhydrous dimethyl sulfoxide (DMSO) (Sigma-Aldrich, St. Louis, MO) at 5 mM. For oligomer preparation the sample was diluted to 100 μM in sterile ice-cold Ham’s F-12 cell culture medium with 1-glutamine but without phenol red (F-12, BioWorld, Dublin, OH) and incubated for 24 h at 4 °C. For fibril preparation, the sample was diluted to 100 μM in 10 mM HCl and incubated for 24 h at 37 °C. Aβ concentrations in these preparations were based on dry peptide weight.

**Size Exclusion Chromatography.** An amount of 1–1.5 mg of lyophilized Aβ peptide was dissolved in 50 mM NaOH to yield a 2.5 mM Aβ solution. The solution was then diluted to 250 μM Aβ in sterile prefiltered (0.22 μm) Ham’s F-12 cell culture medium (supplemented above) with additional supplementation of 100 U/mL penicillin, 0.1 mg/mL streptomycin, and 0.25 μg/mL amphotericin-B (heretofore referred to as supplemented F-12). Aβ(1–42) solutions were centrifuged immediately at 18 000g for 10 min with a Beckman-Coulter Microfuge18 instrument while Aβ(1–40) solutions were incubated for 24 h at 25 °C prior to size-exclusion chromatography (SEC). The centrifugation supernatant was eluted from a Superdex 75 HR 10/30 column (GE Healthcare) in supplemented F-12 medium. Prior to injection of Aβ, Superdex 75 column was coated with 2 mg bovine serum albumin (BSA, Sigma) to prevent any nonspecific binding of Aβ to the column matrix. Following a 1 mL loading of the sample, Aβ was eluted at 0.5 mL/min and 0.5 mL fractions were collected and immediately placed on ice. Fractions 13, 14 eluting in the void volume were pooled together and designated as protofibrils while fractions 24–25 eluting in the included peak were pooled and designated as monomer. UV absorbance at 280 nm was monitored continuously in milliabsorbance units (mAU) during the elution, and concentrations of both protofibrils and monomers were determined directly from the absorbance trace using an extinction coefficient of 1450 cm⁻¹ M⁻¹ at 280 nm. The delay volume between the absorbance detector and fraction collector was minimal, and this was verified in initial experiments by measuring Aβ concentrations in the fractions on a Cary Bio 50 UV absorbance spectrophotometer and comparing those concentration values to those determined from the chromatography absorbance trace. Aβ(1–42) fibrils were prepared from SEC-purified Aβ(1–42) monomer in supplemented F-12 by incubation at room temperature under gentle agitation for ~72 h. The aggregated Aβ(1–42) solution was then centrifuged at 180 000g for 10 min, supernatant was removed, and the pellet was resuspended in the same volume of supplemented F-12. The concentration of fibrils was assumed to match the original starting concentration of the Aβ(1–42) monomer. Thioflavin-T (ThT) fluorescence measurements were done on the total Aβ(1–42) solution, the 18 000 g supernatant, and the resuspended pellet in order to monitor the conversion of Aβ monomer into fibrils.

**Thioflavin T Fluorescence Measurements.** Aβ(1–42) monomer, protofibril and fibril solutions were assessed by ThT fluorescence as described previously.34 Aβ aliquots were removed and diluted 10-fold into 50 mM Tris-HCl pH 8.0 containing 5 μM ThT. Fluorescence
emission scans (460–520 nm) were acquired on a Cary Eclipse fluorescence spectrophotometer using an excitation wavelength of 450 nm and integrated from 470 to 500 nm to obtain ThT relative fluorescence values. F-12 medium, supplemented or not, did not show any significant ThT fluorescence in the absence of Aβ. All ThT fluorescence numbers are reported in relative fluorescence units denoted arbitrary units (a.u.) in the figures.

**Atomic Force Microscopy.** Aβ(1–42) aggregation solutions (100 μM) were diluted to 10 μM in water. Grade VI mica (Ted Pella, Inc., Redding, CA) was cut into 11 mm circles and affixed to 12 mm metal discs. Aliquots (50 μL) were applied to freshly cleaved mica, allowed to adsorb for 15 min, washed twice with water, air-dried, and stored in a container with desiccant. Images were obtained with a NanoScope III multimode atomic force microscope (Digital Instruments, Santa Barbara, CA) in TappingMode. Height analysis was performed using NanoScope III software on flattened height mode images.

**Transmission Electron Microscopy.** SEC-purified Aβ(1–42) protofibril and aggregation solutions were diluted to 20 μmol/L in water unless otherwise stated and applied (10 μL) to a 200-mesh Formvar-coated copper grid (Ted Pella, Inc.). Samples were allowed to adsorb for 10 min at 25 °C, followed by removal of excess sample solution. Grids were washed three times by placing the sample side down on a droplet of water. Heavy metal staining was done by incubation of the grid on a droplet of 2% uranyl acetate (Electron Microscopy Sciences, Hatfield, PA) for 5 min, removal of excess solution, and air drying. Affixed samples were visualized with a JEOL JEM-2000 FX transmission electron microscope operated at 200 kV.

**Dynamic Light Scattering.** Hydrodynamic radius (R_H) measurements were made at room temperature with a DynaPro Titan instrument (Wyatt Technology, Santa Barbara, CA). Samples (30 μL) were placed directly into a quartz cuvette and light scattering intensity was collected at a 90° angle using a 10 s acquisition time. Particle diffusion coefficients were calculated from autocorrelated light intensity data and converted to R_H with the Stokes–Einstein equation. Average R_H values were obtained with Dynamics software (version 6.7.1). Histograms of percent intensity vs R_H were generated by Dynamics software where data regularization and intensity-weighted mean R_H values were derived from the regularized histograms.

**XTT Cell Viability and Proliferation Assay.** Viability of Aβ-treated microglia was determined by using an XTT [2, 3-bis(2-methoxy-4-nitro-5-sulfophenyl)-2H-tetrazolium-5-carboxanilide] assay. Cellular metabolic activity was monitored by mitochondria-mediated reduction of XTT (Sigma). XTT assay was also used in a cell-free manner to rule out the possibility of any bacterial contaminant of Aβ(1–42) and Aβ(1–40) samples. In cellular XTT assay, cells exposed to Aβ for 6 h were further incubated with 0.33 mg/mL XTT and 8.3 μM phenazine methosulfate (PMS) (Acros, Morris Plains, NJ) for 2 h at 37 °C. The extent of XTT reduction was measured by absorbance of reduced form of XTT at 467 nm. The cell-free XTT assay was done in parallel to the cell stimulation studies, in a similar manner as above except the Aβ sample was incubated with XTT and PMS without any cells at the same final concentration of 15 μM for 6 h.

**Statistical Analysis.** Cellular responses and protofibril length measurements were evaluated for statistical differences by paired t test analysis in SigmaPlot 10.0 software.

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**Author Contributions**

M.N. conceived the project. G.P. performed the following: preparation and purification of protofibril, monomer, and fibril samples, cell culture and primary microglia isolation, microglial activation studies, toxicity assays, ELISA, and fluorescence measurements. L.G. did the AFM imaging and analysis. D.O. did the TEM imaging. M.N. performed the DLS studies. M.N. and G.P. wrote the paper.

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**ABBREVIATIONS**

AD, Alzheimer’s disease; Aβ, amyloid-β protein; HFIP, hexafluoroisopropanol; SEC, size exclusion chromatography; ThT, thioflavin T; TEM, transmission electron microscopy; AFM, atomic force microscopy; DLS, dynamic light scattering

**REFERENCES**


