Glitter in the Darkness? Nonfibrillar β-Amyloid Plaque Components Significantly Impact the β-Amyloid PET Signal in Mouse Models of Alzheimer Disease

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β-amyloid (Aβ) PET is an important tool for quantification of amyloidosis in the brain of suspected Alzheimer disease (AD) patients and transgenic AD mouse models. Despite the excellent correlation of Aβ PET with gold standard immunohistochemical assessments, the relative contributions of fibrillar and nonfibrillar Aβ components to the in vivo Aβ PET signal remain unclear. Thus, we obtained 2 murine cerebral amyloidosis models that present with distinct Aβ plaque compositions and performed regression analysis between immunohistochemistry and Aβ PET to determine the biochemical contributions to Aβ PET signal in vivo. Methods: We investigated groups of AppNL-G-F and APPPS1 mice at 3, 6, and 12 mo of age by longitudinal 18F-florbetaben Aβ PET and with immunohistochemical analysis of the fibrillar and total Aβ burdens. We then applied group-level intermodality regression models using age- and genotype-matched sets of fibrillar and nonfibrillar Aβ data (predictors) and Aβ PET results (outcome) for both Aβ mouse models. An independent group of double-hit APPPS1 mice with dysfunctional microglia due to knockout of triggering receptor expression on myeloid cells 2 (Trem2−/−) served for validation and evaluation of translational impact. Results: Neither fibrillar nor nonfibrillar Aβ content alone sufficed to explain the Aβ PET findings in either AD model. However, a regression model compiling fibrillar and nonfibrillar Aβ together with the estimate of individual heterogeneity and age at scanning could explain a 93% of variance of the Aβ PET signal (P < 0.001). Fibrillar Aβ burden had a 16-fold higher contribution to the Aβ PET signal than nonfibrillar Aβ. However, given the relatively greater abundance of nonfibrillar Aβ, we estimate that nonfibrillar Aβ produced 79% ± 25% of the net in vivo Aβ PET signal in AppNL-G-F mice and 25% ± 12% in APPPS1 mice. Corresponding results in separate groups of APPPS1/Trem2−/− and APPPS1/Trem2−/− mice validated the calculated regression factors and revealed that the altered fibrillarity due to Trem2 knockout impacts the Aβ PET signal. Conclusion: Taken together, the in vivo Aβ PET signal derives from the composite of fibrillar and nonfibrillar Aβ plaque components. Although fibrillar Aβ has inherently higher PET tracer binding, the greater abundance of nonfibrillar Aβ plaque in AD-model mice contributes importantly to the PET signal.

Key Words: amyloid; fibrillar; nonfibrillar; PET signal; mouse

J Nucl Med 2022; 63:117–124
DOI: 10.2967/jnumed.120.261858

PET for β-amyloid (Aβ) is now widely used for identification and quantification of amyloidosis in the brain of suspected Alzheimer disease (AD) patients (1) and has been incorporated into the current research framework for diagnostic recommendations in AD (2). Here, the Aβ status (A) identified by PET serves for diagnosis, together with biomarkers for tau and neuronal injury (2). Furthermore, Aβ PET is used as an inclusion criterion in antiamyloid immunotherapy clinical trials (3) and as a progression biomarker for therapy evaluation in these trials (4). In the preclinical setting, Aβ PET has also become a useful tool for the dynamic assessment of neuropathology in transgenic Aβ mouse models (5,6). Despite the excellent correlation of Aβ PET with immunohistochemical gold standard assessments of amyloidosis in patients (7,8) and mouse models of AD (6,9), there has remained an uncertainty about the relative contributions of fibrillar and nonfibrillar Aβ components in plaques to the Aβ PET signal in vivo. This research gap needs to be closed because the 2 forms have differing neurotoxicity, and there is evidence that alterations in AD-related genes such as TREM2 (triggering receptor expressed on myeloid cells 2) and APOE (apolipoprotein E) alter the net Aβ plaque fibrillarity, which would consecutively bias the relationship between plaque density and Aβ PET binding in vivo (5).
A human autopsy–validated 18F-flurbetaben PET study showed preliminary evidence that diffuse plaques may make only a minor contribution to the net Aβ PET signal (10). However, autopsy-controlled data with 18F-flutemetamol in vivo (11) and comprehensive in vitro data (12) indicated that the binding of that structurally distinct tracer to diffuse plaques also contributes to the net PET signal. Furthermore, our recent preclinical study revealed a discernible Aβ PET signal in AppNL-G-F mice (13), although this model displays only very limited fibrillar Aβ pathology (14). Therefore, we aimed to quantify the contributions of fibrillar and nonfibrillar plaque components to the Aβ PET signal in vivo in AD-model mice.

We recently demonstrated that the AppNL-G-F and APPPS1 mouse models exhibit differences in Aβ plaque fibrillarity (14), such that a comparative study of these mice could help to determine the effect of fibrillarity on Aβ PET signal in vivo. Thus, we combined a standardized preclinical 18F-flurbetaben PET study with immunohistochemical characterization of fibrillar versus nonfibrillar Aβ in AppNL-G-F and APPPS1 mice examined at different pathologic stages. We then developed a regression model for immunohistochemistry and Aβ PET to establish the relative proportions of fibrillar and nonfibrillar sources in the Aβ PET signal in vivo. Furthermore, we validated the calculated regression factors in an independent cohort of APPPS1/Trem2−/− and APPPS1/Trem2+/− mice and tested a hypothesis that the nonfibrillar Aβ pool contributes more to the Aβ PET signal in APPPS1/Trem2−/− mice than in APPPS1/Trem2+/− mice.

MATERIALS AND METHODS

Experimental Design

All experiments were performed in compliance with the National Guidelines for Animal Protection, Germany, and with the approval of the regional animal committee (Regierung Oberbayern) and were overseen by a veterinarian. Animals were housed in a temperature- and humidity-controlled environment with a 12-h light–dark cycle and with free access to food (Sniff) and water. We conducted longitudinal 18F-flurbetaben Aβ PET imaging in cohorts of female AppNL-G-F (n = 18) and APPPS1 (n = 14) mice at 3, 6, and 12 mo of age, together with an age- and sex-matched group of wild-type (n = 8) mice. Of the Aβ mouse models, 56% had their baseline examination at 3 mo of age and the remaining 44% were imaged from 6 to 12 mo of age. All mice of each model originated from the same breeding colony. To exclude batch effects within each modality, we used separate cohorts of mice (14) for immunohistochemistry analyses of fibrillar and nonfibrillar Aβ plaque components in wild-type and AD-model mice (n = 3−4) at 3, 6, and 12 mo of age. We then applied intermodality regression models to separate the relative contributions of fibrillar and nonfibrillar Aβ plaque components to Aβ PET signals in the 2 strains.

Animal Models

APP/PS1 (APPPS1-21) mice show extensive fibrillar Aβ plaque pathology, first evident at 6–8 wk of age (15). In contrast, AppNL-G-F (AppNL-G-F/APP) is a murine model with relatively limited fibrillar Aβ plaque pathology but showing Aβ PET signal from 8 wk of age in homozygous mice (16,17). Wild-type controls were C57BL/6 mice.

PET Imaging

PET Data Acquisition, Reconstruction, and Postprocessing.

For all PET procedures, radiochemistry, data acquisition, and image preprocessing were conducted according to an established, standardized protocol (6). In brief, we obtained 18F-flurbetaben Aβ PET recordings (average dose, 12.1 ± 1.8 MBq) with an emission window of 30–60 min after injection.

PET Image Analysis. We performed all analyses using PMOD (version 3.5; PMOD Technologies). Normalization of attenuation-corrected emission images to SUV ratio images was performed using previously validated periaqueductal gray matter (18) and white matter reference regions for the AppNL-G-F and APPPS1 mouse models, respectively (5). We analyzed the wild-type mice separately with both reference regions to serve as controls for the Aβ mouse models. Bilateral necortical volumes of interest (15 mm³) matching the region of interest in the immunohistochemistry analysis were applied for calculation of forebrain–to–white matter SUV ratio or forebrain–to–periaqueductal gray matter SUV ratio.

Immunohistochemical Analysis

Groups of APPPS1 and AppNL-G-F mice at an age of 3 mo (n = 4), 6 mo (n = 3), and 12 mo (n = 4) were transcardially perfused with ice-cold phosphate-buffered saline (0.1 M) followed by 4% paraformaldehyde, after being cryopreserved in 30% sucrose. The mouse tissue used for immunohistochemical analysis included some of the APPPS1 and AppNL-G-F mouse brains used in our previous publication (14). All stainings and analyses were performed newly for the purpose of the present study. We collected 30-μm-thick coronal sections for free-floating immunostaining. We used the 3552 antibody (1:5,000 (19)) to label total Aβ, and we used thiazine red (2 μM; Sigma) to stain the fibrillar Aβ. Twenty-four images were acquired in 4 coronal sections (6 images per section) in regions matching PET using a confocal microscope (×20 dry objective, TCS SP5; Leica). Given the prominent differences in the levels of fibrillar Aβ between the APPPS1 and AppNL-G-F mice, the confocal settings were optimized for each model mouse to acquire the thiazine red signal. For the 3- and 6-mo-old AppNL-G-F mice, the averaging and accumulation confocal functions were set to 2, to better detect the thiazine red signal. An in-house–programmed macro from ImageJ (NIH) was used to analyze the total and fibrillar Aβ coverage.

As a validation analysis, we reanalyzed data from a previous study that included immunohistochemistry markers for fibrillar (x-34) and total (3552) Aβ components of Aβ plaques (5). Immunohistochemistry was obtained from APPPS1/Trem2−/− and APPPS1/Trem2+/− mice (3 and 6 mo, n = 4; 12 mo, n = 8). Aβ PET data were analyzed by the processing pipeline described above and at the same time points for both genotypes (APPPS1/Trem2−/−: three 3-mo-old, ten 6-mo-old, and ten 12-mo-old mice; APPPS1/Trem2+/−: seven 3-mo-old, nine 6-mo-old, and seven 12-mo-old mice). Furthermore, for validation purpose we obtained Aβ coverage for fibrillar (methoxy-x04 or x-34) and total (3552) Aβ components at 13 mo of age in the PET cohorts. In all datasets, nonfibrillar Aβ was calculated by subtraction of fibrillar Aβ from total Aβ (percentage nonfibrillar area = percentage total area − percentage fibrillar area).

Statistics

 Prism (version 8.43; GraphPad Software, LCC) was used for all statistical tests. A P value of less than 0.05 was considered to be significant for rejection of the null hypothesis.

Group-Level Analysis. Nonfibrillar Aβ, fibrillar Aβ, and the Aβ PET z score were compared between AppNL-G-F and APPPS1 mice at different ages by an unpaired Student t test. Mean values of each of the 3 readouts from the AppNL-G-F and APPPS1 groups at different ages were subject to a linear regression analysis. The area between the regression plots served as an index of the potential bias in the estimates of Aβ pathology by Aβ PET.

Individual-Level Analysis. We applied regression models using the Aβ PET z score of all investigated mice in both models as an outcome variable. Fibrillar Aβ and nonfibrillar Aβ estimates deriving
from all age- and genotype-matched mouse groups were used as predictors, and heterogeneity of individual mice with respect to PET results and age were used as additional covariates. We defined heterogeneity as the deviation of individual mice in each genotype from their group mean at each time point. The regression coefficients for fibrillar Aβ and nonfibrillar Aβ were calculated to extract their relative contributions to the Aβ PET signal. Bootstrapping was performed with 1,000 random samples.

**Validation Analysis.** The derived regression coefficients were applied to immunohistochemistry analysis of independent samples of APPPS1/Trem2+/+ and APPPS1/Trem2−/− mice. The predicted Aβ PET z scores were compared with the actual Aβ PET z scores in vivo, and the deviation between the predicted and actual Aβ PET z scores was compared with separate consideration of both plaque components and sole consideration of fibrillar Aβ. The bias resulting from consideration of only fibrillar Aβ was calculated as a function of longitudinal changes in the Aβ PET signal in the comparison of APPPS1/Trem2+/+ and APPPS1/Trem2−/− mice.

**RESULTS**

**Separate Quantification of Fibrillar or Nonfibrillar Aβ Plaque Deposition Fails to Explain the Aβ PET Signal**

First, we performed a direct standardized comparison of nonfibrillar and fibrillar Aβ estimates by immunohistochemistry and Aβ PET between AppNL-G-F and APPPS1 mouse models at different ages. Nonfibrillar Aβ area coverage of AppNL-G-F mice exceeded that of APPPS1 mice at 3 and 6 mo of age, whereas APPPS1 mice had higher nonfibrillar Aβ area coverage at 12 mo of age (Figs. 1A and 2). Fibrillar Aβ area coverage was significantly higher in APPPS1 mice than in AppNL-G-F mice at all ages studied (Figs. 1B and 2). Immunohistochemically assessed area coverage values did not differ between the immunohistochemistry cohorts and the PET cohorts at 12/13 mo of age (all P > 0.05, Supplemental Fig. 1; supplemental materials are available at http://jnm.snmjournals.org). Aβ PET z scores of AppNL-G-F and APPPS1 mice are provided and illustrated in Figures 1C and 1D. There were no interindividual SUV ratio differences between mice imaged 3 times, at 3, 6, and 12 mo, and mice imaged only twice, at 6 and 12 mo (all P > 0.05, Supplemental Fig. 2). Aβ PET showed significantly higher standardized differences in APPPS1 mice than in AppNL-G-F mice at 6 and 12 mo, whereas there were no significant differences at 3 mo of age (Fig. 1C). Plotting of Aβ PET results as a linear function of nonfibrillar or fibrillar Aβ at different ages indicated a mismatch between the 2 mouse models (Fig. 1E). Plotting of fibrillar Aβ as a linear function of nonfibrillar Aβ coverage underpinned that APPPS1 mice had a higher proportion of fibrillar Aβ than did AppNL-G-F mice (Fig. 1E). The comparison of the linear functions of both mouse models (the area transected by the regression lines) indicated that Aβ PET underestimated the proportion of nonfibrillar Aβ in AppNL-G-F mice (−2.08 z score units) but overestimated the proportion of fibrillar Aβ in AppNL-G-F mice (+2.36 z score units). Thus, neither fibrillar nor nonfibrillar Aβ alone could explain the combined Aβ PET findings.

**Nonfibrillar Aβ Contributes Significantly to the Aβ PET Signal**

Next, we hypothesized that a combined model of nonfibrillar and fibrillar Aβ components could improve the explanation of variance in the Aβ PET signals. To test this hypothesis, we established a multiple-regression model using all available combinations of age- and genotype-matched Aβ PET–immunohistochemistry or histochemistry estimates with inclusion of all AppNL-G-F and APPPS1 mice.

Simplified regression models with either fibrillar or nonfibrillar Aβ as predictors of the Aβ PET z score explained 50% and 32% of the variance in Aβ PET, respectively (both P < 0.001, Table 1). Combined consideration of fibrillar Aβ and nonfibrillar Aβ as predictors of the Aβ PET z score increased the explained variance to 57% (P < 0.001, Table 1; Fig. 3A), and fibrillar (β = 0.563, P =
1.11e \(-27\) and nonfibrillar (\(\beta = 0.309, P = 9.38e^{-11}\)) A\(\beta\) were both strong and independent predictors of the A\(\beta\) PET \(z\) score. A model including fibrillar and nonfibrillar A\(\beta\) components with the estimate of individual heterogeneity yielded 85% explanation of variance of the A\(\beta\) PET signal (Fig. 3B), and further inclusion of age further increased the explanation of variance of the A\(\beta\) PET signal (93%, Fig. 3C). Thus, age-related factors influence, importantly, immunochemistry/histochemistry and PET signals (i.e., age-

**TABLE 1**

| Model                             | \(R^2\)  | Corrected \(R^2\) | B Fibrillar       | B Nonfibrillar  | \(P\)       |
|-----------------------------------|----------|--------------------|-------------------|----------------|------------|
| Fibrillar                         | 0.499    | 0.497              | 4.414 (3.915–4.926) |                | <0.001     |
| Nonfibrillar                      | 0.324    | 0.322              | 0.477 (0.378–0.569) |                | <0.001     |
| Fibrillar \times nonfibrillar     | 0.574    | 0.571              | 3.521 (2.954–4.041) | 0.259 (0.178–0.341) | <0.001     |
| Fibrillar \times nonfibrillar \times heterogeneity | 0.854    | 0.852              | 3.521 (2.954–4.041) | 0.259 (0.178–0.341) | <0.001     |
| Fibrillar \times nonfibrillar \times heterogeneity \times age | 0.928    | 0.927              | 2.810 (2.620–3.032) | 0.146 (0.103–0.193) | <0.001     |

\(R^2 = \) coefficient of determination; \(B = \) regression coefficient.
Numbers in parentheses represent 95% CIs as assessed by bootstrapping with 1,000 random samples.

**FIGURE 2.** Representative images of immunochemistry and histochemistry. Total A\(\beta\) was assessed by 3552 staining, and fibrillar A\(\beta\) was assessed by thiazine red. Hoechst (HOE, blue) was used for nuclear visualization (scale bars, 500 \(\mu m\)).
dependent perfusion or partial-volume effects). We considered “fibrillar × nonfibrillar × heterogeneity” and “fibrillar × nonfibrillar × heterogeneity × age” to be the most accurate models, and we calculated the mean regression coefficients from these 2 models to obtain the contributions of fibrillar Aβ and nonfibrillar Aβ to the PET signal. One percent area covered by fibrillar Aβ explained 3.17 PET z score units, and 1% area covered by nonfibrillar Aβ explained 0.20 PET z score units, thus indicating a 16-fold higher contribution of fibrillar than of nonfibrillar Aβ. The opposite edges of the 95% CIs, as assessed by bootstrapping, indicated a possible range between 11-fold and 26-fold for the relationship between fibrillar and nonfibrillar contributions to the Aβ PET signal. Application of this multiplicative factor to the direct comparison of group-averaged immunochemistry or histochemistry and PET endpoints using all available combinations with matched age and genotype. (D) Application of average regression factors for fibrillar (B = 3.17) and nonfibrillar (B = 0.20) Aβ on combined immunochemistry and histochemistry data for both models (group means per age).

Although the resulting Aβ PET tracer signal is inherently 16-fold higher when comparing equal amounts of fibrillar and nonfibrillar Aβ, the larger proportions of nonfibrillar plaque components counterbalance the net contribution. We validated our regression model in an independent cohort of APPPS1 mouse models in vivo. Our data clearly show that nonfibrillar plaque fractions have a significant impact on the net 18F-flurbetaben binding to Aβ plaques of Aβ mouse models in vivo.

**DISCUSSION**

We provide the first—to our knowledge—in vivo analysis to elucidate the contributions of fibrillar and nonfibrillar plaque components to the Aβ PET signal. Application of the regression factors to immunochemistry/histochemistry data indicated an excellent prediction of the actual Aβ PET signal in independent cohorts of APPPS1/Trem2+/− and APPPS1/Trem2+/+ mice (Fig. 4A). APPPS1 mice with Trem2 loss of function showed a higher contribution of nonfibrillar plaque components to the Aβ PET signal (30% at 3 mo, 26% at 6 mo, and 24% at 12 mo) than did APPPS1 mice with intact Trem2 (4% at 3 mo, 15% at 6 mo, and 21% at 12 mo; Fig. 4A). A combined consideration of fibrillar Aβ and nonfibrillar Aβ predicted the actual PET signal more precisely that did sole consideration of fibrillar Aβ (Fig. 4B). Previously calculated increases in Aβ PET signal with age in these mice indicated a considerable bias when considering only the fibrillar Aβ component (Supplemental Fig. 3). In summary, microglial dysfunction altered the relative proportions of fibrillar and nonfibrillar Aβ, thus directly influencing the Aβ PET signal as a function of mouse age.

**Fibrillar and Nonfibrillar Plaque Components Have an Impact on Mice with Dysfunctional Microglia**

Finally, we validated our results in independent cohorts of APPPS1 mice and made an additional investigation of the impact of Trem2 deficiency on the Aβ PET signal in these mice, given that Trem2 is a known driver of changes in the plaque fibrillarity.
examination to elucidate the separate contributions of fibrillar and nonfibrillar Aβ sources to the in vivo Aβ PET signal. We performed PET acquisitions in both Aβ models and wild-type mice with identical housing conditions and using the same tomograph and image reconstruction parameters, thus minimizing the potential methodologic bias. Nonetheless, we acknowledge that scanning of mice on different days of the week, along with social hierarchy factors and technical factors due to different cage positions, might still impact the detection of plaque pathology by Aβ PET. We had to choose between conducting a longitudinal PET examination with immunohistochemistry in a separate cohort and conducting a cross-sectional study in which PET examination directly preceded immunohistochemistry/histochemistry in the same mice. Since animal-batch effects may introduce a bias into cross-sectional PET quantifications between different ages of a given mouse model, we elected to conduct longitudinal PET imaging together with cross-sectional immunohistochemistry and histochemistry to exclude batch effects, at least for PET. To account further for the heterogeneity (22) and asymmetry (13) of amyloidosis in individual mice of a lineage, we used a bilateral target in a regression model including each available combination of PET–immunohistochemistry/histochemistry results for each model at each of 3 ages, controlled for the individual heterogeneity. Regression coefficients for the proportions of fibrillar and nonfibrillar Aβ in different models were robust and revealed that fibrillar Aβ makes an intrinsically 16-fold higher contribution to the Aβ PET signal than nonfibrillar Aβ in the studied Aβ mouse models. Our preclinical in vivo results concur with the postmortem validation of human 18F-flutemetamol PET data, where ligand binding to diffuse plaques was the most likely explanation for positive in vivo signals in patients who later proved to have only sparse neuritic plaques at autopsy (11,23). The same research group recently validated the contributions of diffuse and neuritic plaques to the 18F-flutemetamol and 11C-Pittsburgh compound B autoradiography signals in an in vitro study (12). However, our study was the first translation of such findings into the in vivo setting and enables the arithmetic conversion of Aβ PET signals into fibrillar and nonfibrillar Aβ components. We note that the structure of the stilbene 18F-florbetaben is different from the structures of the benzothiazoles 18F-flutemetamol and 11C-Pittsburgh compound B and that this difference could result in different proportions of fibrillar and nonfibrillar binding capacities in vivo. As usual, the limited resolution of small-animal PET systems in relation to the mouse brain size and resulting partial-volume effects present a limitation for the transfer of the present findings into the human context, and we want to emphasize that detailed regression factors cannot be transferred directly. Yet, the demonstration of an inherently 16-fold higher contribution of fibrillar plaque to the PET signal concurs with a biophysical chemistry study investigating binding mechanisms of Aβ ligands by molecular docking, molecular dynamics, and generalized Born-based free-energy calculations (24). Here, core sites of Aβ fibrils, which are more abundant in fibrillar components of the plaque, dominated over surface sites in producing the Aβ PET signal (24).

Our findings could be of translational relevance since Aβ immunotherapy and other treatment strategies against AD may change the proportions of fibrillar and nonfibrillar plaque

FIGURE 4. (A) z scores of measured Aβ PET signal and predicted fibrillar and nonfibrillar (blue) source components in independent cohort of APPPS1 mice with dysfunctional (Trem2<sup>−/−</sup>) and intact (Trem2<sup>+/+</sup>) microglia. Pie charts show fibrillar and nonfibrillar contributions to measured Aβ PET signals predicted by immunohistochemistry. Representative double staining of APPPS1/Trem2<sup>−/−</sup> mouse shows more nonfibrillar Aβ (3552-positive) surrounding core (x-34-positive) than in APPPS1/Trem2<sup>+/+</sup> mouse, both at 3 mo of age. (B) Bias of predicted z scores when only fibrillar or combination of fibrillar and nonfibrillar plaque contributions was considered. Analysis displays deviation of immuno/histochemically predicted Aβ scores when only fibrillar PET signal. We performed PET acquisitions in both Aβ models and wild-type mice with identical housing conditions and using the same tomograph and image reconstruction parameters, thus minimizing the potential methodologic bias. Nonetheless, we acknowledge that scanning of mice on different days of the week, along with social hierarchy factors and technical factors due to different cage positions, might still impact the detection of plaque pathology by Aβ PET. We had to choose between conducting a longitudinal PET examination with immunohistochemistry in a separate cohort and conducting a cross-sectional study in which PET examination directly preceded immunohistochemistry/histochemistry in the same mice. Since animal-batch effects may introduce a bias into cross-sectional PET quantifications between different ages of a given mouse model, we elected to conduct longitudinal PET imaging together with cross-sectional immunohistochemistry and histochemistry to exclude batch effects, at least for PET. To account further for the heterogeneity (22) and asymmetry (13) of amyloidosis in individual mice of a lineage, we used a bilateral target in a regression model including each available combination of PET–immunohistochemistry/histochemistry results for each model at each of 3 ages, controlled for the individual heterogeneity. Regression coefficients for the proportions of fibrillar and nonfibrillar Aβ in different models were robust and revealed that fibrillar Aβ makes an intrinsically 16-fold higher contribution to the Aβ PET signal than nonfibrillar Aβ in the studied Aβ mouse models. Our preclinical in vivo results concur with the postmortem validation of human 18F-flutemetamol PET data, where ligand binding to diffuse plaques was the most likely explanation for positive in vivo signals in patients who later proved to have only sparse neuritic plaques at autopsy (11,23). The same research group recently validated the contributions of diffuse and neuritic plaques to the 18F-flutemetamol and 11C-Pittsburgh compound B autoradiography signals in an in vitro study (12). However, our study was the first translation of such findings into the in vivo setting and enables the arithmetic conversion of Aβ PET signals into fibrillar and nonfibrillar Aβ components. We note that the structure of the stilbene 18F-florbetaben is different from the structures of the benzothiazoles 18F-flutemetamol and 11C-Pittsburgh compound B and that this difference could result in different proportions of fibrillar and nonfibrillar binding capacities in vivo. As usual, the limited resolution of small-animal PET systems in relation to the mouse brain size and resulting partial-volume effects present a limitation for the transfer of the present findings into the human context, and we want to emphasize that detailed regression factors cannot be transferred directly. Yet, the demonstration of an inherently 16-fold higher contribution of fibrillar plaque to the PET signal concurs with a biophysical chemistry study investigating binding mechanisms of Aβ ligands by molecular docking, molecular dynamics, and generalized Born-based free-energy calculations (24). Here, core sites of Aβ fibrils, which are more abundant in fibrillar components of the plaque, dominated over surface sites in producing the Aβ PET signal (24).

Our findings could be of translational relevance since Aβ immunotherapy and other treatment strategies against AD may change the proportions of fibrillar and nonfibrillar plaque
components and thus bias the Aβ PET outcome. Furthermore, alterations of microglial genes are associated with changes in plaque morphology, which consequently influence the Aβ PET signal (5). We can correlate the more diffuse amyloid plaque morphology in App<sup>NL-G-F</sup> mice with differences in plaque morphology observed in AD mice deficient in TREM2 or APOE (3,25). Although loss-of-function mutations of Trem2 are rare in humans, microglia genes seem in general (26) to have a high impact on AD pathology, and modulation of microglial function is being intensively studied as a therapeutic strategy for AD (27). Another limitation for the direct translation of our findings toward human AD consists in different binding site densities of Aβ plaques in Aβ mouse models when compared with sporadic AD (28). Thus, comparisons of Aβ PET signal intensity between rodents and humans need to be considered with caution. However, the regression model generated in this study should increase awareness of the impact of nonfibrillar Aβ on the Aβ PET signal in both species. Thus, a potential shift in the plaque proportions needs to be considered when designing future Aβ PET monitored studies that target microglia. Assessment of fibrillar and nonfibrillar plaque components and the respective Aβ PET tracer-binding properties in autopsy cases after or during disease-modifying treatment studies of AD could serve to test the impact of our findings on the human situation.

**CONCLUSION**

The Aβ PET signal with <sup>18</sup>F-flurbolten in vivo arises from a combination of fibrillar and nonfibrillar plaque components. Fibrillar Aβ has inherently higher tracer binding, but the greater proportion of nonfibrillar Aβ relative to fibrillar Aβ in most plaques means that the nonfibrillar signal source is a relevant component of the total signal. Since experimental AD therapy regimens can shift the proportion of fibrillar versus nonfibrillar Aβ, any longitudinal changes in Aβ PET signal as a readout of therapy monitoring must be interpreted with caution; a detailed understanding of the biochemical basis of Aβ PET signal is critical for the correct use of PET for monitoring novel AD therapies.

**DISCLOSURE**

This work was supported by an Alzheimer’s Association grant through the AD Strategic Fund (ADSF-21-831226-C) and by the German Research Foundation within the framework of the Munich Cluster for Systems Neurology (EXC 2145 SyNergy/ID 390857198). Sabina Tahirovic was supported by the Alzheimer Forschungs-Initiative e.V (grant 18014). Christian Haass collaborates with Denali Therapeutics, participated in one advisory board meeting of Biogen, and received a speaker honoraria from Novartis and Roche. Christian Haass is chief advisor of ISAR Bioscience. Peter Bartenstein, Axel Rominger, and Matthias Brendel received speaking honoraria from Life Molecular Imaging and GE Healthcare. Matthias Brendel is an advisor of Life Molecular Imaging. No other potential conflict of interest relevant to this article was reported.

**ACKNOWLEDGMENTS**

We thank Mathias Jucker for providing the APPPS1 mice and Takashi Saito and Takaomi C. Saido for providing the App<sup>NL-G-F</sup> mice.

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**QUESTION:** Does nonfibrillar Aβ contribute to the in vivo Aβ PET signal?

**PERTINENT FINDINGS:** Fibrillar Aβ has 16-fold higher tracer binding than does nonfibrillar Aβ, but the greater proportion of nonfibrillar Aβ relative to fibrillar Aβ in most plaques means that the nonfibrillar signal source is a relevant component of the total Aβ PET signal.

**IMPLICATIONS FOR PATIENT CARE:** Shifts of the proportion of fibrillar versus nonfibrillar Aβ need to be considered when interpreting the longitudinal Aβ PET signal for monitoring of therapeutic effects.
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