PET amyloid imaging as a tool for early diagnosis and identifying patients at risk for progression to Alzheimer’s disease

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Introduction

The development in 1984 of consensus criteria [1] for diagnosis of Alzheimer’s disease (AD) capped a period of evolving knowledge that AD could be differentiated not only from normal aging but also from other causes of neurodegenerative dementias. On average, clinical diagnosis using these consensus criteria is approximately 81% sensitive and 70% specific compared to the gold standard, pathology at autopsy [2], a performance that equals or exceeds the performance of proposed diagnostic criteria for many other neurodegenerative diseases [2,3].

Nevertheless, there remains both room and a need for improvement in diagnostic accuracy. Up to 20% of subjects clinically diagnosed with AD do not have AD pathology at autopsy [4-6], a percentage that is essentially unchanged from the estimate in the 1984 consensus publication [1]. In addition, under-diagnosis in the community setting is significant. Approximately 10% of community-dwelling elderly have undiagnosed dementia [7,8] and community physicians may fail to diagnose up to 33% of individuals with mild dementia [8].

Perhaps the biggest limitation in current practice is a reliance on the presentation and progression of symptoms to identify an AD phenotype. This inherently leads to delays in diagnosis as physicians must wait for symptoms to appear and must track progressive decline over time. However, the past 25 years have seen dramatic improvements in technology and understanding of biomarkers that offer potential to improve this diagnostic algorithm. As a result, new draft criteria [9,10] have proposed that diagnosis can be enhanced by use of biomarkers to increase certainty, and, in early stages, to identify prodromal AD. This approach has the potential to allow earlier and more specific diagnosis and will possibly identify patients with AD before the point where irreversible damage precludes effective treatment [11].

A number of different biomarkers, including atrophy on magnetic resonance imaging (MRI), regional metabolism as assessed by 18F-fluorodeoxyglucose positron emission tomography (PET), and cerebrospinal fluid (CSF) concentrations of tau and β-amyloid (Aβ) are potentially useful [11,12], but molecular imaging with amyloid targeted PET ligands is a particularly attractive approach. Rate of atrophy on volumetric MRI and pattern of metabolic deficits on 18F-fluorodeoxyglucose PET can provide useful information on stage of deterioration and functional status, but may lack specificity, since multiple types of neurologic disorders can cause the same type of changes [13-17]. CSF markers provide information (albeit indirect) more relevant to the underlying molecular pathology, including both Aβ and tau, but require a relatively invasive procedure (lumbar puncture) and may

Abstract

Current theory suggests that β-amyloid accumulation may be an early step in the cascade that leads to cognitive impairment in Alzheimer’s disease. β-Amyloid targeted positron emission tomography (PET) imaging potentially provides a direct, relatively noninvasive estimate of brain β-amyloid burden. This has recently been supported by demonstration that amyloid plaque binding on PET was strongly correlated with brain β-amyloid burden at autopsy. Additionally, there is growing consensus that PET imaging can identify subjects with elevated β-amyloid burden, even at early stages of disease. Finally, preliminary evidence suggests that abnormal β-amyloid accumulation, as evidenced by PET imaging, has implications for both present and future cognitive performance. Although large longitudinal studies like the ongoing ADNI trial will be required for definitive evaluation, present data suggest that PET amyloid imaging has the potential to promote earlier and more specific diagnosis of dementia.

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not be entirely specific for AD [18]. In contrast, Aβ imaging potentially provides a direct, relatively non-invasive estimate of brain Aβ burden, which together with tau and a progressive pattern of neuronal loss is a defining pathology and an import link in the pathogenesis of AD [19,20].

The first, and to date most widely studied, ligand for PET imaging of Aβ aggregates (subsequently referred to as amyloid PET or amyloid imaging) is the 11C-labeled agent known as Pittsburgh compound B (PIB) [21-23]. Although 11C-PIB has been a highly valuable tool in the research setting, the short (20-minute) half-life of the 11C label limits the utility of 11C-PIB in routine clinical application. Thus, there has been a push to develop a longer lived 18F-labeled amyloid PET agent. Three compounds are currently in the late stages of development. One of these, florbetapir F 18 [24-26] has now completed phase III trials [27], while florbetaben [28] and flutemetamol [29,30] are currently enrolling to phase III trials.

The utility of PET amyloid imaging as an aid in early diagnosis rests on three major assumptions: first, that PET imaging accurately reflects Aβ burden in the brain; second, that PET imaging can detect brain Aβ at an early stage of disease, that is, prior to the onset of dementia; and finally, that the presence of β-amyloid, as detected by PET imaging, has consequences for current and future cognitive performance. We will examine the available evidence for each of these assumptions in turn.

Relationship between PET amyloid imaging and brain Aβ burden by histopathology

In vitro studies have shown that PET imaging ligands such as 11C-PIB [21,31], florbetaben [32] and florbetapir F 18 [24] bind to Aβ and co-localize with plaques stained by thioflavin and other amyloid labeling agents. However, a definitive demonstration of the relationship requires a comparison between in vivo imaging and brain pathology, for example, at autopsy.

Five single subject/single center PET to pathology comparison studies with 11C-PIB have produced mixed results. Two studies described patients with clinical diagnosis and autopsy confirmation of dementia with Lewy bodies (DLB) who had amyloid-positive 11C-PIB PET scans in life, and borderline Aβ pathology at autopsy. Bacskai and colleagues [33] reported a visually positive 11C-PIB PET scan from a 76 year old with DLB and severe cerebral amyloid angiopathy. Regional quantification of the PET image, expressed as distribution volume ratio (DVR), revealed low to moderately elevated tracer levels (DVR = 1.3 to 1.5), which was consistent with the autopsy findings of low to moderate levels of diffuse plaques and infrequent cored plaques (intermediate probability of AD by National Institute of Aging - Reagan Institute (NIA-Reagan) [34] criteria). However, there was no relationship across brain regions between regional DVR and regional levels of Aβ42 in autopsy tissue as assessed by ELISA. Kantarci and colleagues [35] reported a positive 11C-PIB PET scan from a 77 year old with DLB. At autopsy neuritic plaques were moderately common in some brain regions, including mid-frontal gyrus, amygdala and superior parietal lobe, but sparse in the areas used for pathological diagnosis, resulting in an NIA-Reagan classification of low likelihood AD. In contrast to the previous study, there was a strong correlation between regional quantification of the PET image and regional Aβ density by immunohistochemistry at autopsy. Two other reports studied subjects with a clinical diagnosis of AD. Ikonomovic and colleagues [31] reported an amyloid positive 11C-PIB PET scan in a 64 year old with severe AD. Strong correlations (0.7 to 0.8) were seen between regional 11C-PIB PET tracer uptake (DVR) and various postmortem measures of Aβ burden, including immunohistochemistry, histopathology and Aβ levels by ELISA. Cairns and colleagues [36] reported on a 91 year old with clinical diagnosis of early AD with a negative 11C-PIB PET scan but reduced CSF Aβ. The autopsy revealed numerous diffuse plaques, but sparse cored plaques and isolated neurofibrillary tangles (NFT). The neuropathologic diagnosis in this subject was borderline: low probability of AD by NIA-Reagan criteria, and possible AD by CERAD (Consortium to Establish a Registry for Alzheimer’s Disease) criteria [37]. Additionally, the 11C-PIB PET scan was taken more than 2 years prior to autopsy. Thus, it is difficult to determine whether this case should be considered a failure of the 11C-PIB PET scan to detect an early stage of AD, or a successful rejection of a case that lacked convincing AD pathology. Finally, Leinonen and colleagues [38] reported that five of ten subjects who had a tissue removed for a shunt for normal pressure hydrocephalis had significant numbers of Aβ aggregates by immunohistochemistry at biopsy. Four of these subjects had abnormal 11C-PIB PET scans (elevated cortex to cerebellum standard uptake volume ratio (SUVR)). The overall correlation between SUVR and number of amyloid aggregates across the ten subjects was 0.85.

Clark and colleagues [27] recently reported the first prospective multicenter phase III study to evaluate the correlation between the level of cortical amyloid burden on PET scan and true Aβ burden assessed by postmortem histopathology. In this study, 152 subjects with cognitive status ranging from cognitively normal to mild cognitive impairment (MCI) to AD or other dementing disorders agreed to both florbetapir-PET scan and subsequent autopsy. As specified by the protocol, the first six subjects to come to autopsy were considered front runners and were used to confirm the experimental methods, and the next 29 subjects to come to autopsy were considered the
primary efficacy population. Cortical amyloid burden on florbetapir-PET scans was visually assessed on a 0-to-4 scale (no-to-high cortical tracer uptake) by three independent raters, blinded to clinical information, with the median rater score as the primary outcome variable, and by a semi-automated quantification of the SUVR in six cortical target areas (frontal, temporal and parietal cortex, precuneus, anterior and posterior cingulate) relative to the cerebellum reference region. Amyloid burden at autopsy was assessed by quantitative immunohistochemistry (primary outcome variable) and by a modified CERAD scoring (silver stain) in the six cortical target regions. The results showed a strong, statistically significant correlation between the level of cortical tracer uptake in the PET image, whether assessed by median visual read or SUVR, and true Aβ burden, whether assessed postmortem by quantitative immunohistochemistry or silver stain (ρ = 0.71 to 0.78, P < 0.0001). Similar results were obtained in the primary efficacy set (n = 29) and in the entire autopsy data set (n = 35, including the front runners). There was qualitative agreement between florbetapir-PET and postmortem results in 97% of the autopsied subjects. Of 19 subjects that met pathologic criteria (CERAD and NIA-Reagan) for AD, 18 were rated visually positive for amyloid by median read, and all 19 had SUVR above a predefined cutpoint. Conversely, all 16 subjects that did not meet pathologic criteria (amyloid free) at autopsy were amyloid free by both visual and quantitative analysis of the PET scan.

Although the data with 11C-PIB are somewhat limited, the results with florbetapir F 18 provide a strong preliminary indication that PET amyloid imaging can provide an accurate reflection of underlying Aβ burden. However, further studies are required to understand how early in the disease course the amyloid pathology can be detected. In both the 11C-PIB [36,38] and florbetapir F 18 [27] studies there were some subjects with measurable but low levels of amyloid pathology at autopsy that were not associated with amyloid-positive PET scans. In most cases, the level of pathology in these patients at autopsy was below the threshold for neuropathological diagnosis of AD (that is, rated low likelihood or no AD). Thus, the threshold for detection of amyloid on the PET scan appears close to the levels of neuropathology typical for a diagnosis of AD. It is presently unclear whether levels of Aβ burden at autopsy that are insufficient to be thought of as AD actually represent an early stage of disease [35,36], or whether they represent variants of amyloid deposition, including normal aging [39]. Longitudinal studies, with periodic repeat scans and cognitive testing, would be useful to determine how much or for how long a negative scan in a cognitively normal individual reduces risk of future amyloid accumulation and cognitive impairment. Such studies are now starting as part of the second phase Alzheimer’s Disease Neuroimaging Initiative (ADNI; for example, ADNI-2) protocol [40].

On the other hand, across both the 11C-PIB and the florbetapir F 18 image-autopsy studies there were no cases in which a positive amyloid PET scan was obtained in a subject found to be cognitively normal and amyloid free at autopsy. These results suggest that there is a high probability of underlying brain Aβ pathology in subjects with positive amyloid PET scans. This kind of high specificity and positive predictive value, compared to the autopsy gold standard, is a prerequisite for a biomarker to be used as an aid to early diagnosis of dementia.

**Early detection of amyloid by PET imaging in MCI and cognitively normal subjects**

Current theories of AD pathophysiology hold that Aβ deposition may be a precipitating event that begins years in advance of the onset of dementia [41-43]. Evidence in support of the hypothesis includes the finding that 15% or more of cognitively normal subjects coming to autopsy may have plaque burden sufficient to support a diagnosis of AD [44-46] and 33 to 62% of subjects with MCI have significant accumulation of Aβ plaques [47,48]. Corresponding changes in biomarkers have also been reported in non-demented individuals. Notably, studies of CSF biomarkers have consistently shown decreases in CSF Aβ in 30 to 40% of cognitively normal subjects [49,50]. Changes in CSF tau, MRI volume and cerebral metabolism may occur slightly later than changes in CSF Aβ [41,49,51].

Amyloid PET imaging studies have yielded results similar to those from autopsy and CSF studies. Studies using 11C-PIB have reported amyloid-positive scans in 14 to 47% of cognitively normal elderly volunteers [40,43, 52-55], and 55 to 72% of subjects with MCI [51,54-57]. Where data from both 11C-PIB PET scans and CSF Aβ have been available, strong correlations between these measures have generally been reported [49,57]. Results with 18F-labeled imaging agents are similar to those for 11C-PIB. The proportion of Aβ-positive scans in cognitively normal subjects has ranged from 7% and 12% with flutemetamol [29,30], to 13% with florbetapir [26], and 20% with florbetaben [28]. In MCI subjects the proportion of positive scans was about 50% for flutemetamol [30] and florbetaben [58] and about 38% in the studies with florbetapir [59].

The differences across PET studies, which are similar to the differences in the pathological studies of cognitively normal controls and MCI, could easily be related to differences in subject age and inclusion criteria rather than differences in sensitivity of the different tracers. Consistent with findings in the autopsy literature [45,60], the proportion of cognitively healthy control subjects that are Aβ-positive by PET scan increases with age
The mean age of cognitively healthy subjects varied by more than 10 years across the studies above [29,55]. Additionally, the florbetapir trial [58] was designed to evaluate early stage MCI patients, diagnosed within the past year. These subjects may be more difficult to diagnose and thus more heterogeneous, leading to inclusion of a greater number of subjects with non-amyloid/AD-related impairments.

Jagust and colleagues [40], reporting on $^{11}$C-PIB subjects from the ADNI study, further evaluated quantitative values (cortical to cerebellar SUVR) for the Aβ-positive and Aβ-negative subjects by diagnostic presentation group (cognitively healthy, MCI, and AD). Interestingly, there was no apparent difference in SUVR between Aβ-positive MCI and Aβ-positive AD, but SUVR in Aβ-positive MCI and AD both appeared greater than SUVR in Aβ-positive healthy controls. These results are consistent with histopathology findings [47], indicating that the relative proportion of patients with high versus moderate levels of Aβ pathology at autopsy (definite versus probable AD by CERAD criteria) does not increase from MCI to AD patients, and suggests that Aβ accumulation reaches asymptote at early stages of disease.

Together with the image-autopsy results described above [27], these results suggest that PET imaging can detect the presence of Aβ aggregates sufficient to support a pathological diagnosis of AD in upwards of 15% of cognitively healthy elderly subjects (prevalence increasing with age) as well as in 40 to 70% of subjects with MCI. Thus, the results are consistent with the hypothesis [41] that PET amyloid imaging can detect Aβ accumulation well in advance of the onset of dementia. The next section will consider the available literature regarding consequences of a positive amyloid scan for present and future cognitive performance in cognitively healthy and MCI subjects.

**Relationship between amyloid PET imaging and cognitive performance/progression**

The most obvious prediction from the model of Jack and colleagues [41] is that compared to subjects who have a negative amyloid PET scan, cognitively healthy control and MCI subjects who have positive amyloid PET scans will, as a group, show greater deterioration in cognitive performance, and will be more likely to progress to an advanced stage of disease (for example, from MCI to AD). A significant number of studies have looked at the relationship between PET amyloid binding and concurrent cognitive performance. Multiple studies have reported no correlation between amyloid binding and degree of cognitive deficits in AD patients [55,61,62]. This is consistent with the hypothesis that amyloid is an early initiating event in a pathological cascade, that Aβ accumulation approaches asymptote by the time that symptoms appear, and that other pathological processes (tau phosphorylation, inflammation, synaptic degeneration) are more closely linked to expression of cognitive impairment in AD patients [41].

Results are more mixed for MCI subjects. Pike and colleagues [55] found a good correlation ($r = 0.61$) between $^{11}$C-PIB SUVR and a working memory composite score. Others have found no consistent differences in cognition as a function of PET amyloid imaging [43,54]. However, it is likely that correlational studies in MCI subjects are particularly sensitive to the diagnostic algorithms used to select and define MCI subjects. Overlap between the diagnostic algorithm and cognitive outcome variables can reduce the chances of finding a relationship between an independent variable and cognitive performance; for example, if all subjects must have objectively demonstrated memory deficits for inclusion in the study cohort, it becomes difficult to demonstrate a relationship between amyloid burden and memory performance within the cohort. Additionally, as noted above, amyloid levels may approach asymptote by the MCI stage, and differences in brain amyloid burden beyond that point may have as much to do with modulating factors influencing the individual subject’s asymptotic level as they do with disease stage.

In cognitively healthy elderly subjects, Mintun and colleagues [52], Storandt and colleagues [62] and Jack and colleagues [54] reported no relationship between concurrent cognitive performance and $^{11}$C-PIB amyloid binding. Other studies have found mixed results. Mormimo and colleagues [63] reported a relationship between $^{11}$C-PIB amyloid binding and episodic memory for one population of normal elderly, but not for a second population. Rowe and colleagues [43] reported that subjects with high PIB amyloid binding had significantly reduced memory scores relative to subjects with low amyloid binding, but the correlation between binding and memory was not significant. In contrast, several studies [55,64,65] have now reported correlations between $^{11}$C-PIB amyloid binding and memory scores. Similarly, Rosenberg and colleagues [61] examined cognitive performance in the cohort of subjects described by Wong and colleagues [26] and found a significant correlation between florbetapir F 18 binding and ADAS-cog (Alzheimer’s Disease Assessment Scale Cognitive Sub-scale) performance by normal elderly controls. Park and colleagues [66] have also recently reported a relationship between florbetapir PET amyloid binding and working memory performance in cognitively normal aging subjects.

It is not surprising that the strength of correlation between PET result and cognitive performance, and/or the magnitude of the difference in cognitive performance between cognitively normal subjects with Aβ-positive and Aβ-negative PET scans, was modest and sometimes...
variable. At least three factors work to limit the magnitude of effect that can be obtained in cognitively normal subjects. First, the range of cognitive performance in cognitively normal subjects is constrained by the criteria used to separate cognitively impaired subjects from cognitively normal. The earlier and more aggressively the diagnosis of impairment is made, the less potential for variance within the normal group as a function of amyloid level, as subjects with greater amyloid burden, and more advanced impairment, may be classified as cognitively impaired. Second, the outcome may depend on the difficulty of the cognitive tests used. More difficult tests are more likely to uncover deficits that may otherwise go unnoticed [64]. Finally, the relationship between amyloid binding and cognitive performance can be modified by the subject’s education/cognitive reserve [64,65]. Subjects with high education/high cognitive reserve may appear to maintain cognitive function in the normal range for a longer period or in the face of greater PET amyloid binding than subjects with lower cognitive reserve.

The Pike and colleagues [55] and the Rentz and colleagues [64] reports above both include scatterplots of cognitive performance as a function of amyloid binding (SUVR). Rather than a preferential distribution of abnormally low memory scores in association with high amyloid binding, the scatterplots are notable for the relative absence of high memory scores in the high amyloid group. It is tempting to speculate that this kind of distribution is the result of the limiting factors discussed above. In the amyloid-positive cohort, subjects with low cognitive reserve cannot sustain performance and become classified as MCI, whereas subjects with high cognitive reserve, who otherwise would have been above average memory performers, have deteriorated but are still performing near the middle of the normal range. However, this kind of hypothesis can only be addressed by longitudinal studies.

The relationship between amyloid burden as assessed by PET imaging and longitudinal change in cognitive function in cognitively normal and MCI populations is currently under examination in multiple trials, including the US ADNI study [40] (11C-PiB, phase 1, and flortebaip F 18, phase 2), the Australian Imaging, Biomarkers and Lifestyle Initiative (AIBL) initiative [43] (11C-PiB) and several ongoing longitudinal trials of aging [62,67], as well as in several trials with 18F-labeled agents that are either still ongoing (flumetamol, NCT01028053; flortebaiben, NCT01138111; ClinicalTrials.gov) or recently completed (flortebaip) [59]. First results, now coming into the literature, strongly suggest a relationship between amyloid burden and AD progression.

Four published studies have examined the potential of 11C-PiB PET amyloid imaging to predict progression from MCI to AD. Forsberg and colleagues [57] imaged 27 MCI subjects and reported that 7 who subsequently converted to AD had higher PiB retention than non-converting subjects. Okello and colleagues [56] studied 31 MCI subjects, 17 (55%) of whom were considered amyloid-positive on an 11C-PiB PET scan. Of these 17 subjects, 14 (82%) converted from MCI to AD in the follow-up period (up to 3 years). Only 1 of 14 (7%) amyloid-negative subjects converted in the same time period. A comparison of fast (<1 year) versus slower converters suggested that fast converters (within one year of scan; 8 of 17 amyloid-positive subjects) had higher 11C-PiB PET cortical to cerebellar uptake ratios than the slower converters, despite a similar mean age. Notably, all fast converters for whom genotype was available carried an apolipoprotein E ε4 allele, whereas only two of six slow converters with genotype information carried an αpolipoprotein E ε4 allele. Thus, the ε4 allele may have contributed to both the elevated amyloid burden (increased SUVR) and the more rapid conversion. Wolk and colleagues [68] similarly reported a higher rate of conversion in subjects classified as amyloid-positive (5 of 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET. Finally, Jack and colleagues [69] recently published 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET. Finally, Jack and colleagues [69] recently published a higher rate of conversion in subjects classified as amyloid-positive (5 of 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET. Finally, Jack and colleagues [69] recently published 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET. Finally, Jack and colleagues [69] recently published a higher rate of conversion in subjects classified as amyloid-positive (5 of 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET. Finally, Jack and colleagues [69] recently published a higher rate of conversion in subjects classified as amyloid-positive (5 of 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET. Finally, Jack and colleagues [69] recently published a higher rate of conversion in subjects classified as amyloid-positive (5 of 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET. Finally, Jack and colleagues [69] recently published a higher rate of conversion in subjects classified as amyloid-positive (5 of 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET. Finally, Jack and colleagues [69] recently published a higher rate of conversion in subjects classified as amyloid-positive (5 of 13, 38%) versus amyloid-negative (zero of 10) by 11C-PiB PET.
the relationship between PET amyloid binding and continuous measures, that is, change in objectively measured cognitive performance. Storandt and colleagues [62], working with essentially the same subject population as Morris and colleagues [70], found that concurrent cognitive performance was unrelated to $^{11}$C-PIB binding, but the estimated annual rate of cognitive deterioration, as evidenced by change in visuospatial and working memory performance composite scores, was significantly greater in subjects with an amyloid-positive $^{11}$C-PIB PET scan than in subjects with an amyloid-negative $^{11}$C-PIB scan. High amyloid binding on $^{11}$C-PIB scans was also associated with reduced regional brain volume on MRI, further suggesting that even in cognitively normal subjects (CDR 0) amyloid accumulation is not benign. Villemagne and colleagues [71] imaged 34 elderly subjects that had been previously followed longitudinally for 6 to 10 years. On average, subjects with memory decline over the observation period had higher $^{11}$C-PIB retention; 7 of 11 subjects with elevated $^{11}$C-PIB retention showed memory decline, versus 4 of 23 subjects with normal $^{11}$C-PIB retention. Finally, Resnick and colleagues [67] obtained $^{11}$C-PIB PET images on 57 subjects who had been followed for an average of 10.8 years as part of the Baltimore Longitudinal Study of Aging and found a significant correlation between $^{11}$C-PIB binding (DVR) and Mini Mental State Exam and verbal memory (California Verbal Learning Test).

One weakness of the Storandt and colleagues [62], Villemagne and colleagues [71] and Resnick and colleagues [67] studies is that they rely primarily on retrospective analysis of cognitive decline. Although several groups have now reported that change in $^{11}$C-PIB binding is relatively slow, particularly in amyloid-positive subjects [40,51], it is difficult to judge from a retrospective analysis how early the $^{11}$C-PIB PET could have predicted subjects likely to show cognitive decline. Of course, these groups and others (for example, ADNI) are now following subjects prospectively from the point of imaging. One recent preliminary report [59] was consistent with the results above showing a relationship between florbetapir PET amyloid binding and prospectively measured cognitive decline.

In summary, the data to date are limited, but taken together provide evidence that abnormal accumulation of Aβ as evidenced by PET amyloid imaging is associated with increased risk of both concurrent cognitive deficits and subsequent progression of cognitive impairment, and thus may be pathological even in apparently cognitively normal subjects.

**Conclusion**

Emerging consensus regarding diagnostic algorithms and criteria suggests that diagnosis of AD can be enhanced by use of biomarkers to increase certainty, and, in early stages, to identify the group of patients at risk for progression to AD. The data reviewed above suggest that PET amyloid imaging may be well suited to both tasks. Amyloid binding on PET has been shown to be strongly correlated with brain Aβ burden at autopsy, and PET imaging identified amyloid-positive subjects with a high sensitivity and specificity in relationship to postmortem histopathological criteria for AD. Additionally, there is consistent evidence that PET imaging can identify subjects with elevated Aβ burden, even at early stages of disease, and preliminary evidence suggests that excess Aβ accumulation, as evidenced by PET imaging, has implications for both present and future cognitive performance.

Current theory suggests that Aβ accumulation may be a critical early step in a cascade of events, including phosphoryl tau and inflammation-mediated synaptic damage and neuronal loss, that leads to cognitive impairment in AD. Early identification of subjects with Aβ accumulation may be critical to the development of potential disease-modifying therapies because amyloid targeted therapies may not be effective once later stages of the cascade have begun.

There is an opportunity to identify patients earlier than occurs in current clinical practice. Typical patients in clinical trials, who are generally well educated and well integrated into the medical system, report delays of approximately 2 years between symptom onset and diagnosis. Delays may be even greater in a community setting where physicians are known to overlook diagnoses in a substantial proportion of patients. However, improved diagnostic aids, such as amyloid-targeted PET scans, alone may not be sufficient to overcome this problem. Diagnostic delays may be partly a matter of patient education (recognition and acceptance of AD symptoms, readiness to seek treatment) and physician practice. In particular, some physicians may be unwilling to commit to diagnosis in the absence of viable treatments. On the other hand, tools that provide evidence of the underlying pathology might improve physician’s confidence, and lead to an earlier diagnosis, by reducing the need for longitudinal follow-up and progression to a more advanced stage of symptoms. Equally important, the evidence indicates that PET amyloid scans can identify patients with early cognitive impairments who do not have pathological levels of brain Aβ at autopsy. Since Aβ pathology is required for a diagnosis of AD, the early demonstration of the absence of Aβ may lead instead to further evaluation of potentially treatable causes of impairment (for example, depression) in these patients.

When, and in what population of patients, should amyloid PET imaging be used? It is easy enough to identify and rule out the extremes. On the one extreme, a
well characterized patient whose disease has progressed beyond the point where a scan would influence medical management would likely derive little benefit from a PET scan. On the other extreme the evidence to date is not sufficient to support routine use in screening cognitively normal subjects, even in the presence of risk factors. Although the results discussed above (for example, [55,61]) suggest that subjects who are amyloid-positive on PET scan may perform worse on cognitive tests, the results have not been entirely consistent across trials, and the effects are subtle and of uncertain clinical relevance. Most important, too few amyloid-positive subjects have been identified and followed longitudinally to give guidance to the patient regarding likelihood and time course of future cognitive deterioration. Current estimates of 10 years or more between the first signs of excess Aβ accumulation and onset of dementia suggest that many amyloid-positive elderly patients might pass on before experiencing significant cognitive decline.

In between these extremes lie a large number of patients that could potentially benefit from PET amyloid scans. With three 18F-labeled amyloid targeted ligands having entered or already completed phase III trials, it is likely that amyloid PET scans will be broadly available within the next few years. Additional studies and consensus evaluations are needed to determine the best use for these agents. Despite the positive results described above, it is clear that an amyloid PET scan is not sufficient to confer a diagnosis of AD. Aβ can be present in association with other disease conditions, including DLB, Parkinson's disease and cerebrovascular disease. It remains unclear whether this reflects the coincidence of two or more disease entities (for example, AD independently in addition to DLB) or whether Aβ (and tau) pathology can be found independently in multiple disease entities.

In either case, the advent of PET amyloid imaging techniques does not obviate the need for clinical/cognitive evaluation. Moreover, the information obtained from amyloid PET imaging may be enhanced by additional biomarker studies, including, for example, functional imaging [72], or molecular imaging aimed at dopamine systems [73-75]. Additional studies are required to identify which patients most benefit from PET amyloid imaging and which additional diagnostic assessments are most useful in developing a practice parameter to optimize the potential for early evaluation of cognitive impairment.

**Abbreviations**
Aβ, β-amyloid; AD, Alzheimer's disease; ADNI, Alzheimer's Disease Neuroimaging Initiative; CDR, Clinical Dementia Rating; CERAD, Consortium to Establish a Registry for Alzheimer's Disease; CSF, cerebral spinal fluid; DLB, dementia with Lewy bodies; DVR, distribution volume ratio; ELISA, enzyme-linked immunosorbent assay; FDG, 18F-fluorodeoxyglucose; MCI, mild cognitive impairment; MRL, magnetic resonance imaging; NFT, neurofibrillary tangles; NIA-Reagan, National Institute of Aging - Reagan Institute; PET, positron emission tomography; PIB, Pittsburgh compound B; SUV, standard uptake volume ratio.

**Competing interests**
MP and MM are employees and stockholders in Avid Radiopharmaceuticals. Avid owns the patent license for florbetapir F 18, one of the PET amyloid imaging compounds discussed in this review.

**Authors' contributions**
Both MP and MM contributed to the analysis and views expressed in this review. MP drafted the manuscript with critical revision by MM.

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