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PIB is a non-specific imaging marker of amyloid-beta (A β) peptide-related cerebral amyloidosis

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The *in vivo* imaging probe [11C]-PIB (Pittsburgh Compound B, N-methyl[11C]2-(4'-methylaminophenyl)-6-hydroxybenzothiazole) is under evaluation as a key imaging tool in Alzheimer's disease (AD) and to date has been assumed to bind with high affinity and specificity to the amyloid structures associated with classical plaques (CPs), one of the pathological hallmarks of the disease. However, no studies have systematically investigated PIB binding to human neuropathological brain specimens at the tracer concentrations achieved during *in vivo* imaging scans. Using a combination of autoradiography and histochemical techniques, we demonstrate that PIB, in addition to binding CPs clearly delineates diffuse plaques and cerebrovascular amyloid angiopathy (CAA). The interaction of PIB with CAA was not fully displaceable and this may be linked to the apolipoprotein E- ϵ 4 allele. PIB was also found to label neurofibrillary tangles, although the overall intensity of this binding was markedly lower than that associated with the amyloid-beta (A β) pathology. The data provide a molecular explanation for PIB's limited specificity in diagnosing and monitoring disease progression in AD and instead indicate that the ligand is primarily a non-specific marker of A β -peptide related cerebral amyloidosis.

Keywords: PIB; amyloid; Alzheimer's disease; positron emission tomography; imaging

Abbreviations: AD = Alzheimer's disease; ApoE = apolipoprotein E; A β = amyloid-beta peptide; BTA-1 = 2-(4'-methylaminophenyl)benzothiazole; CAA = cerebrovascular amyloid angiopathy; CAA-NDB = CAA-associated non-displaceable binding; CPs = classical plaques; DLB = dementia with Lewy bodies; DPs = diffuse plaques; ECX = entorhinal cortex; FDDNP = 2-(1-{6-[(2-fluoroethyl(methyl)amino]-2-naphthyl)ethylidene]malononitrile}; FTD = frontotemporal dementia; MFG = mid-frontal gyrus; NFTs = neurofibrillary tangles; NDB = non-displaceable binding; OCL = occipital lobe; PET = positron emission tomography; PIB = Pittsburgh Compound-B (2-[4'-(methylamino)phenyl]-6-hydroxybenzothiazole); rCMRglc = regional cerebral glucose metabolism rate; SBI3 = 4-N-methylamino-4'-hydroxystilbene; SPL = superior parietal lobe; SPs = senile plaques

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Introduction

Amyloid-beta (A β) peptide deposition in senile plaques (SPs, for plaque terminology see 'Methods' section), and the formation of neurofibrillary tangles (NFTs) from tau are the key histopathological features for the diagnosis of Alzheimer's disease (AD) (Mott and Hulette, 2005). Great efforts are currently underway to define reliable biomarkers to establish the diagnosis of AD *intra vitam* and in early disease phases, a prerequisite for the development of disease-modifying therapies.

Amyloid is a generic term used to describe polypeptide filaments that are folded into a cross β -sheet structure and as a consequence bind dyes such as Congo Red and the Thioflavins S and T (Lockhart, 2006). Both A β , which forms the fibrils variably associated with SPs (and cerebrovascular amyloid), and tau, which forms the paired helical and straight filaments associated with NFTs, are types of amyloid (Serpell, 2000; Berriman *et al.*, 2003; Barghorn *et al.*, 2004). [11C]-PIB is one of a number of positron emission tomography (PET) imaging probes that were derived from

Thioflavin T and the radiotracer is thought to selectively bind to the highly fibrillar (i.e. amyloid rich) SP subtype known as classical plaques (CPs) (Mathis *et al.*, 2004).

In support of this premise a series of reports have demonstrated the increased, region specific, retention of [11C]-PIB in clinically diagnosed AD patients compared with non-demented control groups (Klunk *et al.*, 2004; Verhoeff *et al.*, 2004; Archer *et al.*, 2006; Engler *et al.*, 2006; Fagan *et al.*, 2006; Edison *et al.*, 2007) in a pattern that broadly matches CP distribution (Nordberg, 2004). Additionally, an inverse correlation of PIB retention with the pattern of decreased regional cerebral glucose metabolism rate (rCMRglc) has been observed (Klunk *et al.*, 2004; Buckner *et al.*, 2005; Engler *et al.*, 2006; Edison *et al.*, 2007). Finally, a 2-year-follow-up study on a cohort of AD patients indicated that the amyloid load, as measured by [11C]-PIB, remained stable irrespective of clinical progression determined by both cognitive measures and rCMRglc (Engler *et al.*, 2006). These latter two findings in particular suggest that amyloid deposition, as detected by PIB, is an early event in the pathogenesis of AD that may (i) precede a decline in cognitive function and (ii) reach an early plateau which then remains stable.

However, reconciling these global, *in vivo* observations with data from both the large number of antecedent post-mortem studies detailing the regional, microscopic spread of AD pathology (Braak and Braak, 1991a; Berg *et al.*, 1998; Price and Morris, 1999) and the relatively sparse literature characterizing the interaction of imaging probes with amyloid lesions (Lockhart, 2006) highlights a number of important discrepancies. First, the high retention of [11C]-PIB in brain regions not normally associated with a high CP load such as the striatum and frontal cortex (Klunk *et al.*, 2004; Nordberg, 2004; Buckner *et al.*, 2005; Engler *et al.*, 2006; Edison *et al.*, 2007) cannot readily be explained by the reported specificity of PIB for CPs. For example studies have demonstrated that the vast majority of plaques in the striatum are of a diffuse plaque (DP) type (Wisniewski *et al.*, 1989; Suenaga *et al.*, 1990) and that CPs are not a dominant pathological feature of the frontal cortex during the early stages of AD (Mochizuki *et al.*, 1996; Berg *et al.*, 1998; Price and Morris, 1999). However, the findings associated with this latter observation are not universal (Braak and Braak, 1991a; Markesbery *et al.*, 2006). Secondly, the dichotomy of the ligand retention pattern into essentially 'PIB-positive' or 'PIB-negative' is at odds both with the central role of the CP in disease progression (Selkoe and Schenk, 2003). A number of studies have also demonstrated that CPs begin to form near the time of clinical presentation and then increase with disease severity (Braak and Braak, 1991a; Berg *et al.*, 1998; Price and Morris, 1999). Thirdly, a 'pathologic' or 'PIB-positive' retention pattern has also been identified in 22 to 40% of cognitively normal individuals (Fagan *et al.*, 2006; Mintun *et al.*, 2006; Rowe *et al.*, 2007) and this represents a significant false positive rate. As no longitudinal PIB-PET data is available from these cognitively

normal subjects, it is currently unclear whether these findings reflect the molecular development of prodromal AD (Barnes *et al.*, 2006) or are consistent normal ageing. Finally, only one imaging study to date has considered the potential influence of the apolipoprotein E (ApoE)- ϵ 4 allele on the pattern of PIB retention (Rowe *et al.*, 2007), as this variant is linked with an earlier disease onset, a more rapid cognitive decline, and a more pronounced accumulation of both SPs and cerebrovascular amyloid angiopathy (CAA) (Berg *et al.*, 1998; Attems, 2005). These issues are further exemplified by the application of [11C]-PIB in non-AD patient populations such as dementia with Lewy bodies (DLB) and frontotemporal dementia (FTD) (Bacsikai *et al.*, 2007; Rowe *et al.*, 2007; Rabinovici *et al.*, 2007).

In order to address these important knowledge gaps an autoradiography study was undertaken using a tritium labelled preparation of the PIB ([3H]-PIB) and fresh-frozen brain sections from cases containing AD-associated neuropathological lesions (SPs, CAA and NFTs). Importantly the study utilized subjects of known ApoE genotype, a tracer concentration of radioligand relevant to imaging studies (Mathis *et al.*, 2004; Rowe *et al.*, 2007) and histological examination of adjacent sections, so that radiolabelled features could be positively linked to specific lesions.

Methods

Compounds

Tritiated Pittsburgh Compound-B ([3H]-PIB, *N*-methyl-[3H]-2-(4'-methylaminophenyl)-6-hydroxybenzothiazole) was prepared as a custom synthesis by GE Healthcare UK Ltd (Little Chalfont, UK) at a specific activity 71 Ci/mmol (2.63 TBq/mmol). BTA-1, 2-(4'-methylaminophenyl)benzothiazole (Sigma, Poole, UK) was prepared as a 10 mM stock in dimethylsulphoxide and stored at -20°C .

Preparation of cryosections

Fresh, frozen brain tissues were obtained from the Brain Donation Program at Sun Health Research Institute (Sun City, AZ, USA). Four consecutive cryosections (8–10 μm thickness) were prepared from each brain specimen and mounted onto statically charged ('plus') glass slides prior to airtight storage with desiccants at -80°C . The first and third sections in each series were used for histochemical analysis. The second and the fourth sections were used for radioligand binding studies defining the total ([3H]-PIB) and non-displaceable binding (NDB) ([3H]-PIB plus BTA-1) signals, respectively.

Radiolabelling of sections

Slides were thawed for 10 min at room temperature (25°C) prior to incubation in PBS/E buffer [phosphate buffer saline (PBS) containing 10% v/v ethanol, pH 7.2] for 15 min at room temperature. Slides were then incubated with 0.5 nM [3H]-PIB (total binding) or 0.5 nM [3H]-PIB plus 10 μM BTA-1 (NDB) for 60 min at room temperature and then terminated by consecutive 30 s incubations in PBS/E ($\times 2$) and distilled water (all at 4°C). Slides were dried under a cold air stream and then apposed with Hyperfilm-3H (GE/Amersham, RPN535B; early availability batch).

Table 1 Summary of histological and immunohistological staining techniques used to identify neuropathological lesions

	A β immunostain (6E10)	Thioflavin S	Gallyas
CPs	++	++	N/A
DPs	+, +/-	+/-	N/A
CAA	++	++	N/A
NFTs	N/A	++	++

Note: As noted in the 'Methods' section both the morphology and differential staining intensity (++ strong staining, + moderate staining, +/- weak staining, N/A not applicable) were used to distinguish between lesion types.

Film development and analysis

Films were developed after an 8-week exposure using GBX developer solutions (Kodak, New Haven, USA). Images of the autoradiographs were captured using the MCID digital densitometry system (Interfocus, Linton, UK). No manipulations other than optimization of lighting conditions (for image capture), image cropping and re-sizing were performed.

Nomenclature and classification of pathological features

The identification of SPs, CAA and NFTs was based on data from two separate staining techniques (Table 1) as well as morphological appearance. The terminology for SP phenotypes follows that of Berg *et al.* (1998). Plaques with generally spherical shapes displaying strong staining with both Thioflavin S and the anti-A β monoclonal antibody 6E10 were defined as classical plaques (CPs). CPs were not further differentiated into neuritic and cored plaques. Plaques with spherical or irregular shapes displaying weak staining for Thioflavin S and weak-to-moderate staining with 6E10 were defined as diffuse plaques (DPs). Senile plaques (SPs) are used as a generic term to encompass both types of plaque lesions (i.e. CPs and DPs). CAA was identified on the basis of strong staining with 6E10 and Thioflavin S, and was differentiated from CPs on the basis of morphological appearance. NFTs were identified by Gallyas and Thioflavin S staining.

Histochemical procedures

All procedures were performed at room temperature unless otherwise noted.

Thioflavin S staining

Thioflavin S is a stain used to detect lesions containing high amounts of amyloid structures and will primarily label CPs, CAA and NFTs (Dickson, 2005). DPs contain low amounts of amyloid structure and only stain weakly with this dye (Dickson, 2005). Sections were removed from the freezer and immediately immersed for 30 min in 4% formaldehyde, freshly prepared from paraformaldehyde crystals and buffered to pH 7.4. Following washes in PBS and partial drying at room temperature, the sections were defatted in 1:1 chloroform/ethanol for 1 h, washed briefly twice in 100% ethanol and then hydrated through graded alcohols to water. Sections were then stained in 0.1% Thioflavine

S for 10 min, differentiated rapidly in 80% ethanol and cover-slipped with Apathy's medium.

Immunohistochemical staining for A β peptide using 6E10

The anti-A β monoclonal antibody 6E10 is considered a sensitive marker for CPs and CAA, but may stain diffuse deposits less intensely (Dickson, 2005). Washes were performed between all steps with PBS containing 0.1% Triton X-100. Sections were removed from the freezer and immediately immersed in 70% ethanol for 2 min. Endogenous peroxidase was then suppressed by immersion in 1% hydrogen peroxide for 30 min. After additional washes, sections were incubated overnight in 6E10 (CHEMICON International, Temecula, USA) diluted 1:1000 and detected using VECTASTAIN Elite ABC Kit (Vector Laboratories, Burlingame, USA), dehydrated and mounted in Permount (Fisher Scientific, Fair Lawn, USA).

Gallyas staining

The Gallyas method is a silver-staining technique that in this form is considered specific for NFTs (Braak and Braak, 1991b; Dickson, 2005). Sections were removed from the freezer and fixed as described earlier for Thioflavin S staining. The staining protocol was as described (Braak and Braak, 1991b) with the addition of two pre-treatment steps to minimize non-specific red blood cell staining. These consisted of immersion in 0.25% potassium permanganate for 4 min followed by incubation in 5% oxalic acid for 5 min.

Results

The neuropathological lesions under investigation were studied using four different brain regions: Category A, SP-predominant pathology used mid-frontal gyrus (MFG); Category B, Mixed pathology (DPs, CPs, CAA and NFTs) used superior parietal lobe (SPL); Category C, CAA-predominant pathology used occipital lobe (OCL); and Category D, NFT-predominant pathology used entorhinal cortex (ECX). The neuropathological sections under investigation were pre-selected for the presence of the desired neuropathological lesions and were as far as possible derived from cases with pathologically confirmed AD. However, given the considerable intermingling of lesion types in AD cases and a desire to sample each lesion type in a relatively pure form it was necessary to use additional neuropathological disease groups. A total of 16 cases (four per category) were analysed in the study and representative data from two cases from each category A–D are reported here in detail (Table 2). A full listing of all 16 cases is provided in the supplementary data.

Category A sections (SP-predominant pathology)

The interaction of [3H]-PIB (0.5 nM) with relatively pure populations of SPs (i.e. a mixture of DPs and CPs) was investigated using sections from the MFG of cases with

Table 2 Neuropathological characteristics of sections

Case	Brain region	Pathology scores of sections used in study				ApoE status	CAA-NDB	Neuropathological diagnosis
		DP	CP	NFT	CAA			
A1	MFG	+++	++	–	–	4/4	N.A.*	Probable AD
A2	MFG	+++	+	–	+	3/4	Yes	Probable AD
B1	SPL	+++	+++	+	+++	4/4	Yes	Definite AD
B2	SPL	+++	+++	+	++	4/4	Yes	Definite AD
C1	OCL	++	++	–	++	2/3	No	Possible AD
C2	OCL	–	–	–	+++	4/4	Yes	Probable AD
D1	ECX	–	–	+++	–	2/3	N.A.*	Not AD
D2	ECX	–	–	+	–	–/–	N.A.*	Not AD

Note: Pathology scores: +++, frequent. ++, moderate. +, sparse. –, absent. Note that the neuropathological diagnosis for each case was based on a full pathological examination according to Consortium to Establish a Registry for AD criteria (Mirra et al., 1991). Note that no ApoE genotyping data was available for case D2. N.A.*, not applicable as CAA absent on these sections. An expanded Table listing all 16 cases can be found in the Supplementary data.

probable AD (Table 2). Both cases demonstrated an intense punctate labelling with [3H]-PIB (Fig. 1A and D) that was strongly associated with the cortical region, although there was also a sparse labelling extending into the immediate subcortical white matter (Fig. 1D, case A2). As demonstrated by the labelling reactions performed in the presence of cold competitor ligand (BTA-1, 10 µM) the vast majority of the binding signal in both the grey and white matter was fully displaceable (Fig. 1B and E).

Alignment of the autoradiography images with the 6E10 (Aβ) immunohistochemistry of the adjacent sections (Fig. 1C and F) indicated a substantial overlap with the punctate staining pattern observed with [3H]-PIB and that this extended across both the white and grey matter lesions. This low-magnification correlation was also confirmed by the higher magnification images (Fig. 1G, case A2). However, it is also clear from these images that there are some areas which although intensely labelled with [3H]-PIB display reduced 6E10 immunoreactivity and this may in part reflect the enhanced sensitivity of the autoradiography method.

Consistent with its known poor sensitivity for DPs (Dickson, 2005) Thioflavin S only strongly labelled the sparse CPs (and CAA) associated with this case (Fig. 1G). Nevertheless there was good correspondence between the radiotracer and all forms of Aβ pathology examined. The ability of [3H]-PIB to label DPs more intensely than Thioflavin S probably reflects the >100-fold higher affinity of radioligand for Aβ fibrils relative to the dye (Lockhart et al., 2004). This dataset demonstrates that at typical tracer concentrations [3H]-PIB can bind not only CPs but also DPs.

Category B sections [mixed pathology (DPs, CPs, CAA, NFTs)]

Sections from the SPL were pre-selected for the presence of all lesion types under investigation. The low-magnification images of sections labelling with [3H]-PIB were again very similar to those obtained with the MFG. A dense, punctate cortical staining pattern was observed (Fig. 2A and D) that

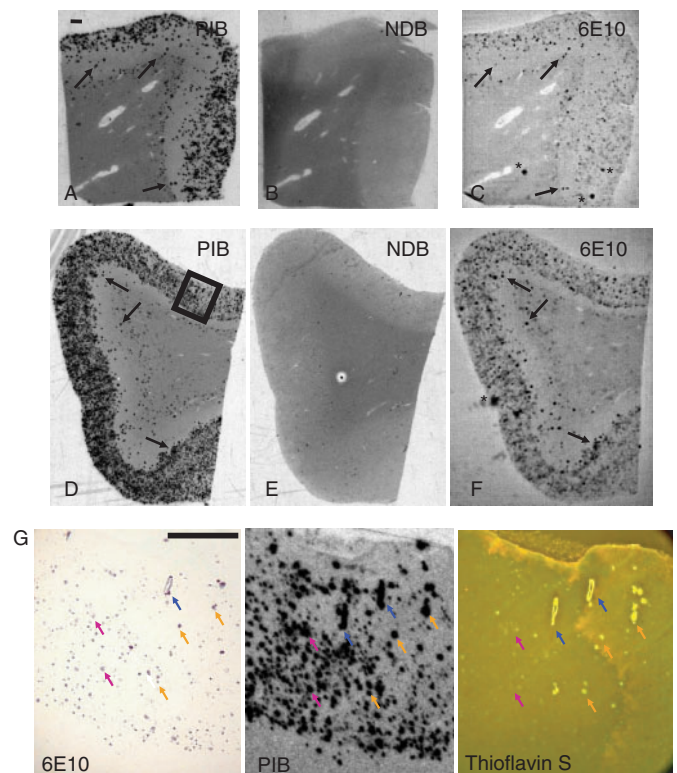


Fig. 1 Images of adjacent MFG sections from two Category A pathology cases (A1 in A–C; A2 in D–G), showing correspondence of [3H]-PIB binding (A and D) with 6E10-immunoreactive Aβ deposits (C and F). NDB is demonstrated in sections treated with the non-radioactive competitive ligand BTA-1 (B and E). Black arrows demonstrate a number of specific corresponding features. Asterisks mark staining artefacts on 6E10 sections. Higher power representative images of boxed area from case A2 arranged in consecutive cryostat order (G). Selected corresponding pathological features are highlighted in pink (DPs), orange (CPs) and blue (CAA). Scale bar 1 mm.

was, in most instances, fully displaceable and which largely overlapped with the 6E10 labelled sections (Fig. 2C and F). Two features however differentiated these sections from the Category A cases (Fig. 1).

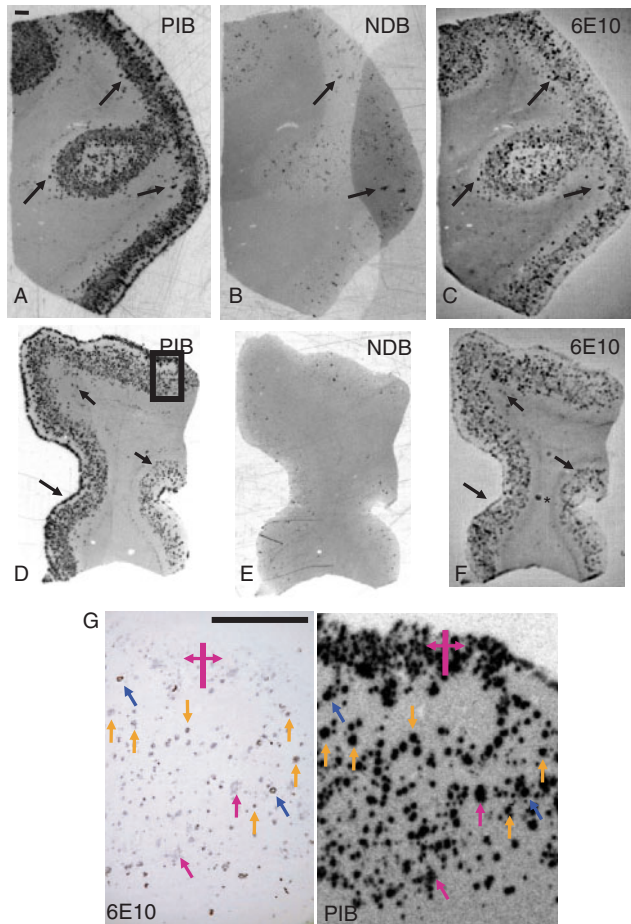


Fig. 2 Images of adjacent SPL sections from two Category B pathology cases (B1 in **A–C**; B2 in **D–G**), showing correspondence of PIB binding (**A** and **D**) with 6E10-immunoreactive A β deposits (**C** and **F**). NDB is demonstrated in sections treated with the non-radioactive competitive ligand BTA-1 (**B** and **E**). Arrows demonstrate a number of specific corresponding features, including subpial diffuse amyloid deposits (black arrows on left of **D** and **F**). Note that on both sections a considerable number of PIB-labelled areas are not completely displaced by the cold BTA-1 compound and that these additionally coincide with 6E10-labelled foci (black arrows on **B**). Higher power representative images of boxed area from case B2 arranged in consecutive cryostat order (**G**). Selected corresponding pathological features are highlighted in pink (DPs), orange (CPs) and blue (CAA). The pink bar and arrows demonstrate corresponding staining of a subpial band of diffuse plaques. Scale bar 1 mm.

First, a number of radiolabelled foci were more obviously retained on the sections incubated with cold competitor ligand (Fig. 2B and E), which as discussed in the Category C cases coincided with CAA. Secondly, the intense labelling observed with [3H]-PIB along the cortical surface of case B2 (Fig. 2D) was associated with a band of weakly 6E10 immunoreactive material (Fig. 2F). Higher power images of the 6E10 sections indicated the presence of subpial ‘band’ or ‘ribbon-like’ A β diffuse deposits associated with the outer portion of the cortical molecular layer (layer 1)

(Fig. 2G). The A β deposits in subpial lesions have been demonstrated to be diffuse, with only small amounts of amyloid fibrils present (Yamaguchi *et al.*, 1991). This finding reinforces our previous contention that PIB will also be sensitive *in vivo* to ‘diffuse’ deposits.

Although Gallyas staining revealed the presence of NFTs in both cases (Table 2) it was not possible to clearly assign any of the [3H]-PIB labelling with this pathological feature due to the high density, close proximity and greater size of the A β lesions. However, as demonstrated for the Category D cases [3H]-PIB clearly labels NFTs [see ‘Category D sections (NFT-predominant pathology) section’].

Category C sections (CAA-predominant pathology)

The OCL is frequently and severely affected by cerebrovascular amyloid deposition in normal ageing and a range of neurodegenerative diseases such as AD (Attems, 2005). Sections labelled with [3H]-PIB from each of the cases displayed strong, punctate staining that at low magnification was broadly similar to the cortical patterns associated with the Category A and B cases (Fig. 3A and D). There was, however, a marked difference in the CAA labelling pattern in the presence of cold competitor ligand. While there was almost complete displacement of the radiolabel in case C1 (Fig. 3B), case C2 retained significant areas of radiolabel, although the overall intensity of the labelled structures was visibly reduced (Fig. 3E). Additional staining of adjacent sections with both 6E10 and Thioflavin S confirmed the presence of A β positive, amyloidotic blood vessels and that the foci of radiolabel retained after treatment with the cold competitor compound remained associated with these structures (Fig. 3G). These features are termed the CAA-associated non-displaceable binding (CAA-NDB).

Although the origins of the CAA-NDB cannot be fully addressed in this study, given the relatively small number of cases examined, one obvious difference between these two cases was their ApoE status. Cases C1 (CAA-NDB negative) and C2 (CAA-NDB positive) were $\epsilon 2/\epsilon 3$ and $\epsilon 4/4$, respectively. Both the remaining two category C cases were negative for CAA-NDB and possessed $\epsilon 3/\epsilon 3$ genotypes (see supplementary data Table). Across the remaining categories CAA pathology was present only in case A2 and cases B1–4. All of these CAA cases were positive for CAA-NDB and contained at least one $\epsilon 4$ allele. Although provocative, further detailed analysis is clearly required to understand the origins of the CAA-NDB.

Category D sections (NFT-predominant pathology)

Hyperphosphorylated tau proteins, the major components of NFTs, are folded into an amyloid structure and thus have the potential to be labelled with [3H]-PIB. The ECX

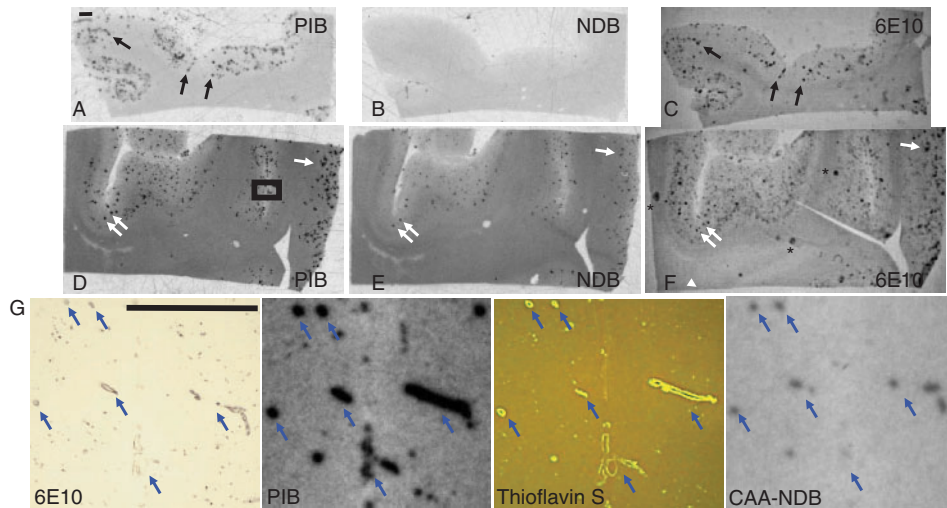


Fig. 3 Images of adjacent OCL sections from two Category C pathology cases (C1 in **A–C**; C2 in **D–G**), showing correspondence of PIB binding (**A** and **D**) with 6E10-immunoreactive A β deposits (**C** and **F**). NDB is demonstrated in sections treated with the non-radioactive competitive ligand BTA-1 (**B** and **E**). Black arrows demonstrate a number of specific corresponding features. Asterisks mark staining artefacts on 6E10 sections. Note that associated with case C2 (**D–F**) the majority of the PIB-labelled areas are not completely displaced by the cold BTA-1 compound and that these additionally coincide with 6E10-labelled foci (black arrows on **E** and **F**). Higher power images of boxed area from case C2 arranged in consecutive cryostat order showing correspondence of PIB binding with 6E10-immunoreactive A β deposits, Thioflavin S-stained amyloid deposits and CAA-NDB (**G**). Blue arrows demonstrate a selection of corresponding amyloidotic blood vessels. Scale bar 1 mm.

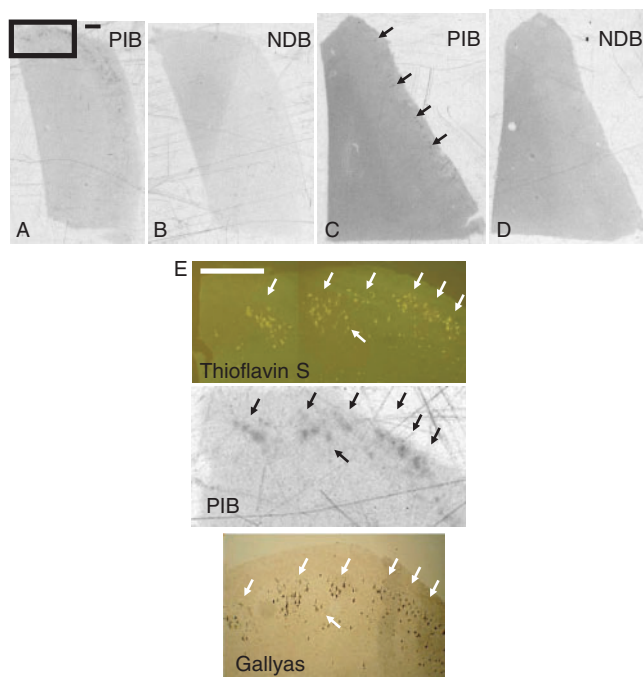


Fig. 4 Representative images of adjacent ECX sections from Category D pathology cases (D1 **A–B** and D2 **C–D**) demonstrating specific binding of [3H]-PIB (**A** and **C**). NDB is demonstrated in sections treated with the non-radioactive competitive ligand BTA-1 (**B** and **D**). Higher power representative images of boxed area from case D1 arranged in consecutive cryostat order showing correspondence of PIB binding with Thioflavin S and Gallyas-positive NFT formations (**E**). White and black arrows demonstrate a selection of corresponding features. Scale bar 1 mm.

appears to be particularly susceptible to the formation of NFTs (Braak and Braak, 1992) and the cases associated with this category were pre-selected for their absence of A β pathology. Associated with case D1 a diffuse, interrupted band of PIB labelling was observed running along the superficial cortical layers (Fig. 4A). Incubation with cold competitor ligand demonstrated that the binding of [3H]-PIB to this lesion was fully displaceable (Fig. 4B). Consistent with its sparse pathological scoring for NFTs (Table 2), case D2 displayed a more discrete punctate labelling with [3H]-PIB that was again associated with the superficial cortical layers and fully displaceable (Fig. 4C and D).

Higher magnification images of the adjacent sections from case D1, labelled with Thioflavin S and Gallyas stains, confirmed the presence of an amyloid structure and NFTs, respectively, that were intimately associated with [3H]-PIB (Fig. 4E). The observed labelling pattern is consistent with localization of the radioligand to the ‘stellate cell islands’ found in the pre-alpha layer (layer II) of the ECX and that as this layer sweeps laterally the ‘islands’ disappears and are replaced by a non-interrupted neuronal band that plunges diagonally through the transentorhinal cortex (Braak and Braak, 1992). Excellent co-registration of individual NFTs with radiolabelled features was also observed in the high-power images of case D2 (see supplementary data Figure). These data indicate that [3H]-PIB can, at a very low concentration, label pathologic human amyloid deposits other than those containing fibrillar A β .

Discussion

The data demonstrate for the first time that at tracer concentrations the neuroimaging agent PIB is not specific for CPs, one of the defining pathological features of AD, but that it additionally binds a range of A β containing lesions including DPs and CAA. The study also established that PIB decorates the tau containing amyloid structures associated with NFTs. However, the contribution of NFTs to the overall AD-associated PIB retention pattern is likely to be minor due to the much greater binding associated with A β lesions. Taken together these findings strongly suggest that *in vivo* [11C]-PIB cortical uptake primarily reflects A β -related cerebral amyloidosis (i.e. is a marker of all fibrillar A β containing pathology: DPs, CPs and CAA).

It is worth emphasising that the experimental protocols used in this study differ significantly from the previous studies investigating the interaction of PIB with human brain sections (Klunk *et al.*, 2003; Mathis *et al.*, 2003, 2004; Bacskai *et al.*, 2007) and that these may explain the novel findings associated with the current study. First, fresh, frozen brain sections were utilized in order to maintain the integrity of both the lesions and the surrounding tissue as close to *in vivo* as possible. Secondly, the use of autoradiography rather than fluorescence techniques permitted the use of concentrations (i.e. low nM range) relevant to *in vivo* imaging studies (Mathis *et al.*, 2004; Rowe *et al.*, 2007), with a detection method insensitive to potential tissue quenching and auto-fluorescence artefacts. Thirdly, detailed supporting histological staining was performed on adjacent sections allowing a direct correlation with the microscopic lesions.

Relevance of findings to *in vivo* imaging studies using PIB

Translation of these findings provides a framework through which the disparities in PIB cortical uptake and neuropathology patterns can be re-evaluated. Plaque type appears to have only a minor influence on PIB binding, as both DPs and CPs are bound with high avidity. PIB uptake in areas with low CP density (e.g. the striatum) and the lack of increase in PIB binding associated with clinical disease progression may both be a consequence of PIB binding to DPs. Histopathological studies indicate that plaque load, a measure of both DPs and CPs, may be high in non-demented individuals (Bouras *et al.*, 1994). While CPs may increase with further overt disease progression, degradation and loss of CPs may also occur as late clinical stages are reached (Brun *et al.*, 1981, 1985; Wegiel *et al.*, 2001). Cortical uptake of PIB in PET is thus a faithful representation of the pathological data, i.e. widespread and dense SP deposition as an early, even preclinical, event in the pathogenesis of AD.

The presence of the CAA-NDB, in the subgroup of $\epsilon 4$ positive cases, highlights the importance of understanding

the interaction of PIB with CAA, due to this fundamental difference in behaviour compared with SPs. Autopsy analysis of a 'PIB-positive' DLB case indicated that CAA was the major pathological source of amyloid and suggested that this lesion type can by itself be a major source of [11C]-PIB cortical uptake (Bacskai *et al.*, 2007). Furthermore, our findings are congruent with those from a study in which the presence of the $\epsilon 4$ allele was linked with a higher cortical uptake of [11C]-PIB, irrespective of clinical diagnosis (Rowe *et al.*, 2007). CAA is biochemically different to parenchymal SPs in that it is formed primarily from A β 1-40 fibrils, although this varies with neuroanatomical location (Attems, 2005). SPs are generally enriched in A β 1-42 species and are additionally subjected to a wide range of post-translational modifications, such as oxidation, isomerization and N-terminal truncation (Lockhart, 2006). Although, *in vitro* data indicates that the interaction of PIB with fibrils formed from either A β 1-40 or A β 1-42 is more or less identical (Klunk *et al.*, 2005) this does not rule the possibility that post-translation modifications may selectively alter the interaction of PIB with one pool. Additionally, the observation that the CAA-NDB was not restricted to the OCL but that it was also observed in the SPL and MFG regions suggests that it is a generic feature. ApoE- $\epsilon 4$ is linked both with CAA severity (Alonzo *et al.*, 1998; Chalmers *et al.*, 2003) and conformation changes to amyloid structure (Sanan *et al.*, 1994), and these factors combined with biochemical differences in the A β pools may account for the occurrence of the CAA-NDB. The data strongly suggests that all individuals undergoing imaging scans with all classes of amyloid imaging probes should be genotyped for ApoE4 and stratified for potential differences in the patterns or kinetics of ligand retention.

The deposition of A β lesions is a common neuropathological finding in a number of conditions other than AD such as vascular dementia (Alonzo *et al.*, 1998) and DLB (Tsuboi *et al.*, 2005; Ballard *et al.*, 2006). The current study clearly supports the continued application of amyloid imaging as another clinical tool to identify and stratify dementias based on their cortical A β load (Rowe *et al.*, 2007; Rabinovici *et al.*, 2007). Although, the current study does not directly address the ability of PIB to act as a pharmacodynamic marker, for drugs targeting A β metabolism (Pangalos *et al.*, 2005), the data suggests that any change in the cortical uptake of the ligand cannot be ascribed to the appearance or disappearance of a particular A β lesion type.

It is reasonable to assume that the interaction between PIB and NFTs is due to the presence of high affinity binding sites associated with the amyloid fold of the tau, given that these structures can be labelled by the chemically related compounds Thioflavin S and T (Dickson, 2005). The weak labelling of the NFTs, relative to that observed with the A β lesions, is consistent with a much lower density of binding sites associated with this form of amyloid and that tangles, in contrast to A β lesions, are therefore unlikely

to be a significant source of tracer uptake *in vivo*. This contention is supported by data from a study using the PIB-related compound, [3H]-BTA-1, which demonstrated that the levels of radioligand associated with ECX homogenates from an AD case were similar to control case levels and around 10-fold less than SP-containing frontal cortex homogenates (Klunk *et al.*, 2003). The higher binding of [3H]-BTA-1 to the frontal cortex homogenates clearly mirrors the intense labelling of SPs and CAA relative to the weakly stained NFTs observed in the current autoradiography study.

Conclusion

In summary, present data supports the application of [11C]-PIB as an important tool for determining the cerebral A β -related amyloid load associated with AD and potentially other types of dementia. However, these findings also provide a molecular explanation for the radiotracer's limited (i) diagnostic specificity and (ii) dynamic range for monitoring disease progression in AD. Alternative amyloid targeting ligands such as [11C]-SB13 (Verhoeff *et al.*, 2004) and [18F]-FDDNP (Shoghi-Jadid *et al.*, 2002; Small *et al.*, 2006) are currently undergoing evaluation studies in dementia populations; understanding their molecular binding profile using analogous experimental procedures will be critical in evaluating their potential as imaging biomarkers.

Supplementary material

Supplementary material is available at *Brain* online.

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