# Positron Emission Tomography and Single-Photon Emission Computed Tomography in Central Nervous System Drug Development

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**Summary:** In this review, the value of functional imaging [positron emission tomography (PET)/single-photon emission computed tomography (SPECT)] in drug development is considered. Radionuclide imaging can help establish the diagnosis of neurodegenerative disorders where this is in doubt and provides a potential biomarker for following drug effects on disease progression. PET and SPECT can help understand mechanisms of disease and determine the functional effects of therapeutic approaches on neurotransmission and metabolism. Synthesizing radiotracer analogs of novel drugs can provide proof of principle that these agents reach their enzyme or receptor targets and delineate their regional brain distribution. If such radiotracers do not prove to have ideal properties for

### INTRODUCTION

Positron emission tomography (PET) and single-photon emission computed tomography (SPECT) provide a means of examining regional cerebral blood flow, metabolism, and pharmacology *in vivo* under both resting and activating conditions. These molecular imaging techniques rely on radiolabeled molecules (tracers) that bind to enzyme sites or surface receptors. PET utilizes shortlived positron emitting isotopes (<sup>15</sup>O, <sup>11</sup>C, <sup>18</sup>F, <sup>76</sup>Br), whereas SPECT uses lower energy  $\gamma$ -emitting isotopes (<sup>123</sup>I, <sup>99m</sup>Tc). Both techniques can detect nanomoles of tracer, but PET has the advantage that it is more sensitive and versatile and allows scatter correction to be performed. SPECT is, however, cheaper and more widely available as it does not rely on a local cyclotron for production of isotopes. A list of some of the neurotransimaging, the concept of microdosing potentially allows multiple other drug analogs to be tested with less stringent regulatory requirements than for novel medicinals. Finally, PET tracers can provide receptor and enzyme active site dose occupancy profiles, thereby guiding dosage selection for phase 1 and phase 2 trials. The eventual hope is that radiotracer imaging will provide a surrogate marker for drug efficacy, although this has yet to be realized, and progress the concept of personalized medicine where receptor/enzyme binding profiles help predict therapeutic outcome. **Key Words:** Positron emission tomography/PET, SPECT, drug development, Alzheimer/Parkinson, neuroprotection, dose occupancy.

mitter systems now amenable to study and the more commonly employed radiotracers is detailed in Table 1.

To derive information about the integrity of neurotransmitter systems from the time course of regional cerebral tracer uptake and arterial plasma activity, it is necessary to use a kinetic model describing the tracer distribution between different physical and metabolic compartments. PET detects 511 keV photons and so cannot distinguish native tracer activity from that of a metabolite. Fortunately, most of the ligands used as PET tracers are not metabolized significantly by brain tissue. The rate of plasma tracer metabolite formation can be determined using serial HPLC measurements enabling a metabolite-corrected input function to be generated.

# KINETIC MODELING

Traditionally, brain uptake of PET tracers has been described using compartmental modeling. The simplest approaches assume that brain regions contain a nonspecific freely exchanging compartment and a compartment where tracer is bound specifically; see Figure 1. A brain

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Biological Application	Tracers
Blood flow	H <sub>2</sub> <sup>15</sup> O, <sup>15</sup> O-butanol. <sup>99m</sup> Tc-HMPAO, <sup>133</sup> Xe
Oxygen metabolism	$^{15}O_{2}$
Glucose metabolism	<sup>18</sup> F-2-fluoro-2-deoxyglucose (FDG)
Dopamine storage/DDC activity	$^{18}$ F-6-fluorodopa (F-dopa), $\beta$ - $^{11}$ C-dopa
Monoamine vesicle transporters	<sup>11</sup> C-dihydrotetrabenazine (DTBZ)
Dopamine transporters (DAT)	<sup>11</sup> C-CFT, <sup>11</sup> C-RTI 32, <sup>18</sup> F-CFT, <sup>123</sup> I-β-CIT
	<sup>123</sup> I-FP-CIT, <sup>123</sup> I-IPT, <sup>123</sup> I-altropane
Dopamine D1 type sites	<sup>11</sup> C-SCH23390
Dopamine D2 type sites	<sup>11</sup> C-raclopride, <sup>11</sup> C-FLB456, <sup>11</sup> C-methylspiperone, <sup>18</sup> F-spiperone,
	<sup>18</sup> F-fluorethylspiperone, <sup>76</sup> Br-bromospiperone, <sup>123</sup> I-epidepride,
	<sup>123</sup> I-iodobenzamide (IBZM)
Noradrenaline transporters	<sup>11</sup> C-BATA
Noradrenaline $\alpha 2$	<sup>18</sup> F-2-fluorethoxyidazoxan
Serotonin storage	<sup>11</sup> C-methyltryptophan
Serotonin transporters	<sup>11</sup> C-DASB, <sup>123</sup> I- $\hat{\beta}$ -CIT
Serotonin HT <sub>1a</sub>	<sup>11</sup> C-WAY100635
Serotonin HT <sub>2a</sub>	<sup>11</sup> C-MDL100907, <sup>18</sup> F-altanserin, <sup>18</sup> F-setoperone
Acetylcholinesterase activity	<sup>11</sup> C-MP4A, <sup>11</sup> C-physostigmine
Cholinergic vesicle transporters	<sup>18</sup> F-fluoroethoxybenzovesamicol, <sup>11</sup> C-vesamicol, <sup>123</sup> I-benzovesamicol
Muscarinic M1 sites	<sup>11</sup> C-tropanylbenzylate, <sup>11</sup> C-NMPB, <sup>18</sup> F-FP-TZTP, <sup>123</sup> I-QNB
Nicotinic sites	<sup>11</sup> C-MPA, <sup>11</sup> C-A-85380, <sup>18</sup> F-A-85380, <sup>123</sup> I-A-85380
Histamine H1 sites	<sup>11</sup> C-dothiepin
Opioid $\mu$ sites	<sup>11</sup> C-carfentenil, <sup>18</sup> F-cyclofoxy
Opioid $\mu$ , $\kappa$ , and $\delta$ sites	<sup>11</sup> C-diprenorphine
Central benzodiazepine sites	<sup>11</sup> C-flumazenil
Central benzodiazepine sites ( $\alpha$ 5 subunits)	<sup>11</sup> C-R015-4513
Peripheral benzodiazepine sites	<sup>11</sup> C-PK11195, <sup>18</sup> F-PK11195, <sup>123</sup> I-PK11195
Substance P/NK1 sites	<sup>18</sup> F-SPARQ, <sup>11</sup> C-GR205171
Adenosine $A_{2A}$ sites	<sup>11</sup> C-SCH 442416
NMDA voltage channels	<sup>11</sup> C-CNS 5161
Amyloid	<sup>1</sup> °F-FDDNP, <sup>1</sup> ′C-PIB, <sup>1</sup> ′C-SB13 <sup>123</sup> I-IMPY
Phosphoglycoprotein activity	<sup>11</sup> C-carfentenil

**TABLE 1.** Radiotracers Available for Studying Neuropharmacology

reference area devoid of the target receptors may provide a measure of nonspecific binding in certain instances. More complex kinetic models include separate brain compartments representing nonspecific ligand binding, freely exchanging pools, ligand metabolism, and ligand diffusion to binding sites.

At tracer doses, a series of first order rate constants can be used to describe the kinetic behavior of a ligand. Rate constants are obtained by solving the linear simultaneous differential equations describing the kinetics of tracer binding using nonlinear regression analysis. The ratio of the two rate constants describing ligand passage from free to bound brain compartments,  $k_3/k_4$ , is known as the binding potential (BP) and, at tracer doses of ligands, reflects  $f_2.B_{max}/K_d$  where  $f_2$  is the tissue free tracer fraction, B<sub>max</sub> the receptor availability, and K<sub>d</sub> the receptor dissociation constant.<sup>1</sup> Measurement of the BP in the presence of different levels of cold ligand along with conventional Scatchard analysis allows B<sub>max</sub> and K<sub>d</sub> to be independently estimated. As in most neurodegenerative conditions K<sub>d</sub> appears to remain unchanged, measurements of BP provide a reflection of changes in B<sub>max</sub> without necessitating displacement studies.



If  $B_{max} >> B$ 

$$k_3/k_4 = BP = f_2 x B_{max} / K_d$$

FIG. 1. A two-tissue compartment model describing radiotracer uptake by the brain. All figures are courtesy of the Medical Research Council Clinical Sciences Centre, UK.

Compartmental modeling requires an input function and regional brain uptake time activity curves (TACs). The input function is generally a metabolite-corrected arterial plasma curve, but, if a nonspecific brain reference area is available (e.g., cerebellum for dopaminergic,  $HT_{1A}$ , and NK1 tracers or occipital cortex for opioid tracers), then a brain reference uptake curve can often be substituted for arterial plasma activity as an input function avoiding the need for arterial cannulation.<sup>2</sup>

A problem with conventional compartmental analysis is that it makes a priori assumptions about the number of pharmacological brain compartments present. A way around this problem is to use spectral analysis that estimates the minimum number of kinetic components required to describe regional brain data sets without making any assumptions about their physiological nature.<sup>3</sup> In essence, the brain tissue function is modeled as a convolution of the measured plasma input function with a sum of k exponential terms where k is the number of frequency components required to describe brain uptake. This exponential function is known as an impulse function and, when integrated, yields parametric images of tracer total volume of distributions (V<sub>d</sub>) which can then be sampled using a conventional region of interest (ROI) approach or subjected to statistical parametric mapping (SPM)-see examples of the use of SPM. Specific V<sub>d</sub> for a region can be estimated by subtracting the reference area  $V_d$  from the total  $V_d$ .

Finally, graphical approaches that effectively linearize brain regional TACs when nonspecific uptake has reached a secular equilibrium with plasma can be used. The Patlak approach assumes an irreversibly bound tracer in the specific compartment and derives an influx constant Ki, which reflects the product of the freely exchangeable ligand  $V_d$  and the association rate constant  $k_3$ .<sup>4</sup> The Logan plot assumes totally reversible ligand binding and derives the reversible specific  $V_d$ .<sup>5</sup> These graphical approaches have the advantage that they are simpler computationally but the linearization process ignores early time points, tends to crowd time points to one end of the plot, and the fitted gradients can be critically dependent on the end points being accurately measured.

# ROI ANALYSIS VERSUS STATISTICAL PARAMETRIC MAPPING

ROI analysis is conventionally used to sample regional brain levels of tracer uptake and to compare these in normal and pathological situations or before and after blockade with cold ligand. There are, however, a number of problems that arise when the ROI approach is employed: First, *a priori* assumptions have to be made about the shape and size of ROIs used to sample brain activity. This can, in part, be overcome by coaligning all functional images to the subject's magnetic resonance imaging (MRI) and using the latter to define ROIs; however, brain shape and orientation vary from subject to subject. Second, an *a priori* selection of the locations of ROIs relevant to the analysis has to be made. One way around this problem is to transform functional images into standard stereotaxic space and apply a predefined template of ROIs from a probabilistic atlas,<sup>6</sup> although this approach inevitably results in some loss of functional resolution.

An alternative to the use of ROIs for analyzing changes in ligand binding associated with pathological situations or the presence of competitive agents is employment of SPM.<sup>7</sup> Here, the whole three-dimensional (3D) brain voxel set is considered without any requirement for a priori definition of ROIs. Parametric images of ligand uptake-either influx constants (Ki) or specific volumes of distribution (V<sub>d</sub>)-are transformed into standard stereotaxic space for each individual; see Figure 2. This can be done using an MRI or brain blood flow template in Montreal Neurological Institute space as provided by the SPM software. Alternatively, a spatial template for the particular radioligand derived from scans of normal subjects can be employed. This last approach has the advantage that, where ligand uptake is concentrated in particular regions, such as the striatum, the accuracy of the spatial normalization is weighted to these areas rather than the cortical rim.

After spatial normalization, group mean images with associated SDs are then generated for each voxel. Tracer binding can subsequently be compared either between different groups of subjects or between an individual subject and a control group. Alternatively, the effects of blocking doses of cold ligands or activating behavioral paradigms on regional tracer binding can be studied. SPM localizes foci of significantly altered Ki or V<sub>d</sub> at preassigned thresholds (generally P < 0.05 with a correction for the number of independent multiple comparisons performed) and the magnitude of changes at these foci can then be measured. The SPM approach has the clear advantage that it is exploratory, and so changes in regional ligand binding can be detected that might not have been otherwise predicted. Its disadvantage is the requirement for stereotaxic transformation of 3D data sets with a consequent loss of functional resolution and a danger of detection of false positives due to the large number of comparisons being performed. The latter can be reduced by the use of masking or small volume correction when a target region is being considered.

#### DIAGNOSIS OF DEGENERATIVE DISORDERS

In a longitudinal survey, serial clinical assessments over several years of 134 patients initially thought to have early Alzheimer's disease (AD) were reported to provide a diagnostic sensitivity of 83% but a specificity



**FIG. 2.** Spatial transformation of <sup>18</sup>F-dopa PET images of a PD patient into standard space using an <sup>18</sup>F-dopa template created from 12 normal subjects. Groups of spatially normalized early PD cases and healthy volunteers are then interrogated with statistical parametric mapping. The SPM shows voxels where <sup>18</sup>F-dopa uptake is significantly reduced in early PD patients clinically affected on only one side. It can be seen that bilateral disease activity is present.

of only 50%.8 Early AD cases generally show little atrophy on conventional MRI, and formal measurements of hippocampal volume show reductions in only 50%.<sup>9</sup> <sup>18</sup>F-2-fluro-2-deoxyglucose (FDG) PET can demonstrate a characteristic Alzheimer pattern of reduced resting regional cerebral glucose metabolism (rCMRGlc) targeting association temporal and parietal cortex. Adding baseline FDG PET measurements to the assessment of 284 early AD cases (138 with subsequent autopsy diagnosis) was shown to improve diagnostic sensitivity to 93% and specificity to 73%.<sup>10</sup> Such an AD pattern can also be detected when present in amnestic mild cognitive impairment (MCI) cases, so aiding the diagnosis.<sup>11</sup> Additionally, asymptomatic adult apo- $\epsilon$ 4 gene carriers,<sup>12</sup> identical AD co-twins,<sup>13</sup> and subjects with mutations of the amyloid precursor protein or presenilin genes may show subclinical metabolic derangement.14,15

Currently there are a number of agents under development designed to prevent amyloid formation (e.g.,  $\beta$  and  $\gamma$  secretase inhibitors)<sup>16</sup> or to remove plaques (active and passive immunization strategies) in Alzheimer's disease.<sup>17</sup> To follow the effects of these agents, it would be beneficial to be able to reliably identify the earliest cases of this disorder. Over half of amnestic MCI patients will progress to develop AD on Diagnostic and Statistical Manual of Mental Disorders, fourth edition criteria over 5 years.<sup>18</sup> The use of FDG PET to screen amnestic MCI for the presence of the characteristic AD pattern of reduced parieto-temporal rCMRGlc has been shown to aid in identifying that minority of cases least likely to progress.<sup>19</sup>

Parkinson's disease (PD) patients show no consistent structural changes on conventional MRI though inversion recovery sequences designed to suppress gray or white matter signals may demonstrate altered nigral structure.<sup>20,21</sup> In contrast, there is a characteristic asymmetrical pattern of loss of striatal dopamine terminal function in PD, the putamen being more affected than the caudate.<sup>22 18</sup>F-dopa PET can detect a subclinical parkinsonian pattern of dopaminergic dysfunction in 50% of asymptomatic adult identical twins<sup>23</sup> of apparently sporadic PD cases, one quarter of relatives of patients with familial PD<sup>24</sup> and, interestingly, in occasional subjects with olfactory dysfunction<sup>25</sup> and rapid eye movement sleep behavior disorder.<sup>26</sup>

There have been two reported trials involving imaging where de novo cases of PD were recruited with a view to assessing the effects of medication on disease progression. In the REAL PET study, 21 of 183 recruited cases (11%) thought to have PD by referring clinicians proved to have normal <sup>18</sup>F-dopa PET.<sup>27</sup> Similarly, in the ELL-DOPA trial, 21 out of 135 cases (16%) felt to have PD had normal  ${}^{123}$ I- $\beta$  SPECT.<sup>28</sup> These subjects have now been followed for up to 6 years, and both their clinical syndromes and imaging findings are unchanged. Clearly, if one is interested in testing putative neuroprotective agents in de novo PD, these subjects with normal striatal dopaminergic function would be best excluded given their failure to show progression. Either one has to screen out these cases by imaging dopaminergic function at baseline or allow for a 10-15% discordance between clinical impression and PET or SPECT assessments of striatal dopaminergic function when powering trials to show a given effect size.

# DEMONSTRATING MECHANISMS OF DISEASE

# Measuring amyloid load in AD

Definitive diagnosis of Alzheimer's disease relies on the demonstration of amyloid plaques and neurofibrillary tangles at autopsy. Amyloid plaques are composed of 40to 42-amino acid amyloid- $\beta$  (A $\beta$ ) peptides. Neurofibrillary tangles contain a hyper-phosphorylated form of the microtubule-associated protein, tau. The time course of amyloid deposition in AD is uncertain, but post-mortem studies of Down syndrome, a congenital disorder associated with trisomy 21, A $\beta$  amyloid deposition, and dementia, suggests that amyloid deposition is likely to start a decade before the clinical symptoms of dementia.

The relationship between deposition of  $A\beta$  plaques and cognitive dysfunction in AD remains uncertain, but a strong link is suggested by the finding that several mutations in the  $A\beta$  precursor protein (APP) gene on chromosome 21 cause early onset AD.<sup>29</sup> Although there are a very small number of families affected with this form of AD, this earlier onset disease is phenotypically indistinguishable from the more common later sporadic form of AD. Further genetic support for the amyloid cascade hypothesis comes from the finding that mutations in the presenilin-1 gene on chromosome 14, which codes for a protein that is strongly implicated to be the  $\gamma$ -secretase enzyme responsible for C-terminal cleavage of  $A\beta$  from its precursor, APP, is associated with early onset AD.<sup>30</sup>

The metabolism of  $A\beta$  has become an important therapeutic target in AD research. Antiamyloid immunization agents<sup>17</sup> and secretase inhibitors<sup>16</sup> are being developed and there are now *in vivo* tools available to directly assess the success of these agents in delaying or reversing amyloid deposition in humans. A demonstration that clinical treatments effectively reduce plaque load while improving cognition will provide valuable proof of concept and strengthen the "amyloid hypothesis" of AD.

There are now three PET imaging agents available that bind to nonphysiological sites in  $\beta$ -sheeted neuritic amyloid plaques. The first is a nonpolar carbon-11-labeled derivative of the fluorescent dye thioflavin-T. This hydroxybenzothiazole, <sup>11</sup>C-6-hydroxybenzothiazole-1, also known as <sup>11</sup>C-PIB (Pittsburgh interesting compound B), showed a significant difference between the retention pattern in 16 AD cases and 9 healthy controls<sup>31</sup>; see Figure 3. Thirteen of the 16 AD cases showed retention of <sup>11</sup>C-PIB in areas of brain association cortex known to contain large amounts of amyloid deposits, whereas retention was minimal in the cerebellum. Three of the clinically diagnosed AD cases, albeit with normal mini-



Normal subject

Alzheimer's disease

**FIG. 3.** <sup>11</sup>C-PIB PET scans obtained from a normal subject and patient with early Alzheimer's disease. The AD case shows wide-spread amyloid deposition in cortical areas and the striatum.

mental state examination (MMSE) scores, showed no abnormal <sup>11</sup>C-PIB retention.

The second agent is a naphthol 2-(1-{6-[(2-[<sup>18</sup>F]fluoroethyl)(methyl)amino]-2-naphthyl}-ethylidene) malononitrile (<sup>18</sup>F-FDDNP) that, in contrast to <sup>11</sup>C-PIB, binds to intracellular neurofibrillary tangles as well as extracellular amyloid plaques.<sup>32</sup> FDDNP appears to compete with certain nonsteroidal anti-inflammatory agents, such as ibuprofen, for amyloid binding, although this is not true of thioflavins. Temporal lobe <sup>18</sup>F-FDDNP transit times have been shown to be prolonged in AD and to correlate with MMSE, verbal, and visual recall scores.

Finally, stilbenes also bind to amyloid plaques. It has been reported that <sup>11</sup>C-4-N-methylamino-4'-hydroxystilbene, may be useful as an amyloid PET imaging agent.<sup>33</sup>

#### Detecting activated microglia

The isoquinoline ligand PK11195 (1-[2-chlorophenyl]-N-methyl-N-[1-methyl-propyl]-3-isoquinolone carboxamide) binds specifically to the peripheral benzodiazepine binding site (PBBS). The PBBS is expressed by mitochondrial outer membranes in cells of the mononuclear phagocyte lineage. Within the CNS, the PBBS is highly expressed by activated, though not resting, microglia-the brain's intrinsic population of tissue macrophages.<sup>34</sup> In combination with volumetric MRI to provide detailed structural information, the entantiomeric PET ligand  $[^{11}C](R)$ -PK11195 has been used to measure microglial activation in acute and chronic inflammatory, and noninflammatory brain diseases including multiple sclerosis,<sup>35</sup> herpes encephalitis,<sup>36</sup> and Rasmussen's encephalitis,<sup>37</sup> Alzheimer's disease<sup>38</sup> and frontal dementias,<sup>39</sup> parkinsonian syndromes,<sup>40</sup> and motor neurone disease.41

The neuronal loss in Alzheimer's disease (AD) is associated with activation of microglial cells whose distribution is closely associated with neuritic amyloid plaques.<sup>42</sup> It may be that microglia play an active role of in mediating amyloid toxicity by causing secondary tissue damage via release of cytokines such as tumor ne-



SN

# Parkinson's

# Alzheimer's

**FIG. 4.** <sup>11</sup>C-PK11195 PET scans obtained from Parkinson's and Alzheimer's disease cases. Increased microglial activation can be seen in the midbrain and basal ganglia of the PD (SN = substantia nigra; GP = globus pallidus) and the cortex (TC = temporal cortex) and brainstem of the AD case.

crosis factor (TNF) $\alpha$  and IL1 $\beta$ . They could also mediate adaptive mechanisms via synaptic stripping and remodeling. A recent <sup>11</sup>C-PK11195 PET study reported two-fold raised levels of microglial activation in AD temporal cortex; see Figure 4.<sup>38</sup>

Microglial activation has also been detected in the motor cortex and brainstem of patients with amyotrophic lateral sclerosis<sup>41</sup> and the basal ganglia of patients with both typical and atypical parkinsonian syndromes,<sup>40,43</sup> being more extensive in the latter. A recent study has reported in abstract that, despite progression of Parkinson's disease, both clinically and as evidenced by <sup>18</sup>F-dopa PET, levels of microglial activation remained relatively unchanged over 2 years.<sup>44</sup>

It has been suggested that suppression of the brain's inflammatory and immune reaction to neuronal degeneration may slow the progression of degenerative disease. Agents such as the peroxisome proliferator-activated receptor  $\gamma$  agonist minocycline, which are known to suppress microglial activation in cell culture and animal models of disease, are currently in trial. <sup>11</sup>C-PK11195 PET potentially provides a biomarker for examining the anti-inflammatory properties of putative neuroprotective agents in neurodegenerative and neuroinflammatory disorders.

#### FOLLOWING DISEASE PROGRESSION

There are a number of difficulties when trying to assess the rate of Parkinson's disease progression clinically. First, timed tests such as finger tapping and hand pronation/supination appear relatively insensitive to progression of early disease though the standing-walkingsitting test is of value.<sup>45</sup> Second, clinical rating scales such as the Unified Parkinson's Disease Rating Scale (UPDRS) are semiquantitative, subjective, and nonlinear and tend to emphasize particular aspects of the disorder, such as bradykinesia. Third, symptomatic effects of therapies can mask disease progression. PD patients generally require symptomatic medication within months of diagnosis and the clinical relief obtained can be profound. Attempts to fully washout such symptomatic effects are poorly tolerated in practice. Additionally, putative neuroprotective agents often turn out to have symptomatic effects-even when these are unsuspected.46 A mild symptomatic effect that improves the UP-DRS rating by two to three points may be enough to delay the need for dopaminergic drugs by several months. Fourth, when assessing neuroprotective efficacy, it is by no means clear what the best primary endpoint should be. Time to requiring medication or a change of medication can be confounded by drug symptomatic effects as can a predefined absolute or percentage change in rating scales. Time to treatment complications such as dyskinesias and motor fluctuations may reflect dopaminergic drug administration schedules as much as disease progression.47

Given these difficulties, there has been an interest in developing imaging biomarkers to follow PD progression. A number of studies have reported that the annual rate of loss of putamen <sup>18</sup>F-dopa and <sup>123</sup>I-β-CIT uptake ranges from 7-12% in early L-dopa-treated cases. 45,48-50 Trials have now examined relative rates of PD progression in patients randomized to either levodopa or a dopamine agonist. Both the REAL PET (<sup>18</sup>F-dopa)<sup>27</sup> and the CALM-PD  $(^{123}I-\beta-CIT)^{51}$  studies showed a one-third relative slowing of loss of dopamine terminal function in the agonist-treated groups. In contrast, the levodopatreated cohorts showed better clinical outcome rated with the UPDRS, although this finding could have been confounded by symptomatic drug effects given that no washout was performed in the REAL PET study and only an overnight washout in the CALM-PD trial. In contrast, the prevalence of dyskinesias was significantly lower in both the agonist-treated groups.

The imaging findings of the above trials have been criticized on the grounds that the medications concerned may have had differential direct effects on the scanning modalities.<sup>52</sup> Putamen <sup>18</sup>F-dopa uptake reflects dopa decarboxylase activity, which is down-regulated by high doses of dopaminergic agonists and upregulated by antagonists in rodents.<sup>53–55</sup> Putamen <sup>123</sup>I-β-CIT uptake measures dopamine transporter binding, which becomes reduced following reserpine induced dopamine depletion<sup>56</sup> and can be increased following levodopa exposure in rats.<sup>57–59</sup> There are relatively limited data concerning the direct effects of medications on brain <sup>18</sup>F-dopa and  $^{123}$ I- $\beta$ -CIT uptake. Findings in animal studies have been inconsistent.<sup>52,60</sup> Human series (albeit small) have failed to show direct drug effects on striatal <sup>18</sup>F-dopa and <sup>123</sup>I- $\beta$ -CIT uptake in both normal and PD subjects but these were underpowered to detect subtle changes. Clearly, if imaging is to be used as a biomarker of disease progression, trials will need to build in an assessment of direct medication effects on the modality, possibly by using a randomized delayed wash-in strategy or by recruiting a separate subject group specifically to address this point.

Other degenerative disorders where PET and SPECT may be of potential value in assessing disease progression include Alzheimer's and Huntington's diseases. MRI shows 3–8% annual changes in hippocampal volume in  $AD^{61}$ , whereas FDG PET shows similar annual rates of reduction in temporal and frontal cortex glucose metabolism (rCMRGlc).<sup>62,63</sup> The annual rate of change of amyloid load in AD has yet to be determined with PET but clearly will be relevant when powering trials to determine the efficacy of antiamyloid agents such as secretase inhibitors. Given the reports of microglial activation in  $AD^{17}$  and its murine models,<sup>64,65</sup> the annual rate of change of levels of microglial activation measured with <sup>11</sup>C-PK11195 PET may also be of interest.

Huntington's disease is a degenerative disorder currently without any effective therapy and so a fertile area for testing neuroprotective agents. Clinical ratings of severity in this condition are less likely than in PD to be confounded by symptomatic drug effects and the Unified Huntington's Disease Rating Scale (UHDRS) has now been validated. In Huntington's disease, annual reductions in striatal glucose metabolism and dopamine D2 receptor binding have been shown to average 3% and 6%, respectively.<sup>66–69</sup> This makes these functional measures a more sensitive marker of HD progression than loss of striatal volume.

#### **DOSE-OCCUPANCY PROFILES**

PET and SPECT can help determine the dose-occupancy profiles of novel ligands for pharma and so help guide dose selection for phase 1 and 2 trials. There are, however, a few caveats: First, there is an inherent assumption that the receptor or enzyme being targeted is relevant to the therapeutic action of the drug. That this may not be the case is illustrated by the use of antidepressants or antipsychotics where symptomatic relief only develops after weeks of exposure to the agent despite high levels of serotonin transporter or dopamine D2 receptor blockade, respectively. Second, this approach only works well for antagonists and partial agonists as full agonists can be therapeutically effective at low levels of receptor occupancy.<sup>70</sup> Third, it is still not clear what level of receptor or enzyme binding is required to achieve the minimum clinically effective therapeutic dose. Based on experience with classical neuroleptics an occupancy of over 70% is generally recommended for antagonists,<sup>71</sup> and this would appear to hold true for other agents such as NK172 and monoamine oxidase B inhibitors.<sup>73</sup> It has to be said, however, that atypical neuroleptics, such as clozapine, are clinically effective when only occupying 50% of D2 receptors.<sup>74</sup>





**FIG. 5.** Blockade of striatal <sup>11</sup>C-raclopride binding by the D2 inhibitor zaprasidone. It can be seen that a 40-mg oral dose of zaprasidone achieves 80% receptor occupancy.<sup>89</sup>

The next question concerns selection of the radiotracer for evaluating dose-receptor occupancy. Options are to use an established radioligand for the receptor type or subtype under investigation or to radiolabel the novel drug under consideration. The advantage of the first approach is that one is ready to run assuming the radiotracer concerned is freely available for use; see Figure 5. The disadvantage is that the specificity of the novel drug being examined and the radiotracer for the receptor concerned may differ. Radiolabeling a novel drug with <sup>11</sup>C or <sup>18</sup>F to produce a PET tracer is generally possible provided a suitable precursor (preferably prepared under GMP conditions) is available. It may, however, take up to 6 months to prepare and validate the new tracer, and not all radiotracers turn out to be useful PET ligands due to their low brain extraction or high nonspecific background signals.

One way of making radiotracer design and selection more efficient involves the concept of microdosing where multiple candidates are prepared as radiotracers and tested simultaneously.75 Because subpharmacological doses are being administered (<20  $\mu$ g) on only one or two occasions, the toxicology requirements for administration of these novel radiotracers in man are less stringent than for medicinals. PET can then assess the plasma, regional brain, and other organ pharmacokinetics in man ahead of therapeutic trials. An example of a toxicology package currently provided for a novel radiotracer being administered two to three times in man would be: six rats given 15  $\mu$ g/kg (100 times human dose) with 3 killed after 48 h and 3 on day 15. Clinical signs, body weight, pathology on limited list of tissues and genotoxicity (Ames test, mammalian cell mutation assay) are performed.

Pharmaceutical companies often ask for nonhuman primate studies with a novel radioligand before going

into man, but these are often not necessary and are expensive in practice. Generally, *ex vivo* studies in the rat give a good indication of whether a radioligand will make a good tracer, but there are some receptors, such as the NK1 site, which show species sensitivity. Additionally, rodent and primate tracer plasma metabolite profiles can differ. In these situations, monkey studies become valuable. Having said that, microPET is capable of examining *in vivo* brain binding in the guinea pig and marmoset, which can avoid the need for higher primate studies.

The next question concerns the optimum design of studies to determine dose-occupancy profiles. A typical approach is to start with an open single dose protocol and choose a time to scan after oral drug administration when brain levels are expected to be peaking. When an oral dose has been identified that results in high receptor occupancy the doses can then be successively reduced to establish a dose-occupancy profile at that time point. Subsequently, occupancy at trough plasma levels can be measured by delaying PET to later times following oral dosing while using a dose that gives high occupancy at peak.<sup>76</sup> Different individuals can be used to establish each baseline and occupancy point on the dose-occupancy profile, or one can use a design combining unoccupied baseline and occupied receptor binding in the same individuals. Generally, establishing baseline receptor availability requires four to eight individuals and each dose occupancy point at least two individuals depending on the interindividual variability.

An alternative, more rigorous approach than the above open design is to blind the study and randomize groups of subjects to different drug doses and placebo. This design, however, is far less flexible than an open design and assumes that one already knows the dose-occupancy profile before starting the study.

As an alternative to a single-dose occupancy design, or following establishment of a peak single dose occupancy profile, a multiple dose paradigm may be employed. Here, the subject is dosed for 1-2 weeks to establish a steady state, and then a trough dose occupancy profile is generated.<sup>77</sup>

# **EFFECTS OF DRUG ACTION**

An advantage of PET and SPECT is that they can examine the downstream effects of novel agents on brain metabolism and pharmacology. An example of this are the studies with <sup>18</sup>F-dopa PET in normal subjects and Parkinson's disease patients before and after administration of the catechol-*O*-methyl transferase inhibitor entacapone with the peripheral dopa decarboxylase inhibitor carbidopa. Four hundred micrograms of entacapone given orally 1 h before PET boosted levels of striatal <sup>18</sup>F-dopa uptake by 50% in normal and early PD subjects



placebo metamphetamine



by blocking peripheral metabolism of the tracer.<sup>78</sup> In advanced PD, the effect was attenuated due to the reduced ability of these patients to store levodopa in the striatum.<sup>79</sup>

An exciting new development of radioisotope imaging is the potential to detect changes in neurotransmitter fluxes. Neurotransmitters that bind to G-coupled receptors, such as dopamine, internalize these receptors transiently after activating them. Radioligands, such as the benzamides <sup>11</sup>C-raclopride, <sup>11</sup>C-FLB, and <sup>123</sup>I-IBZM, compete with dopamine for binding to D2 receptors but can only attach to surface and not internalized sites. As a consequence, the difference in binding of <sup>11</sup>C-raclopride, <sup>11</sup>C-FLB, and <sup>123</sup>I-IBZM before and after a drug or behavioral challenge gives an indirect measure of changes in synaptic dopamine levels.<sup>80</sup> This approach has been used to demonstrate that amphetamine<sup>81</sup> (see FIG. 6), methylphenidate,<sup>82</sup> and scopolamine,<sup>83</sup> all increase levels of synaptic dopamine. Fluxes of opioid transmission can be followed using <sup>11</sup>C-diprenorphine PET.<sup>84</sup> In the future, it may become possible to examine fluxes of other neurotransmitters acting at G-coupled sites, including glutamate.

An indirect approach for examining drug effects on brain function is to examine changes in patterns of resting or activation-induced blood flow and metabolic changes. FDG PET provides a measure of regional cerebral glucose metabolism (rCMRGlc). Patterns of change in rCMRGlc associated with drug administration give a measure of drug effect and may aid dose selection for clinical trials.<sup>85</sup> The advantage of this metabolic approach is that it is equally applicable to agonists and antagonists though its utility still remains to be validated. Along similar lines, changes in activation-induced regional cerebral blood flow (rCBF) patterns measured with either  $H_2^{15}O$  PET or functional MRI using the blood oxygen level-dependent (BOLD) effect can be studied before and after a drug challenge. Buspirone has been shown to augment verbal memory induced increases in prefrontal blood flow while these are attenuated by low doses of apomorphine.<sup>86</sup> Ketamine has been shown to abolish the limbic blood flow response to viewing faces exhibiting a fearful expression.<sup>87</sup> The HT2c agonist, m-chlorophenylpipermazine (mCPP), increased BOLD signal in the hypothalamus, caudate, pallidum, amygdala and pyriform cortex, anterior cingulate gyrus and choroid plexus.<sup>88</sup> In a subsequent Go/NoGo task, mCPP enhanced activation in right lateral orbitofrontal cortex. These findings suggest that pharmacologial challenges with H<sub>2</sub><sup>15</sup>O PET and fMRI are potentially powerful tools for investigating neurotransmitter function in humans.

#### CONCLUSIONS

PET and SPECT imaging can aid drug development by:

- Demonstrating drugs reach their receptor and enzyme targets so providing proof of concept.
- Defining dose occupancy curves for phase 1 and phase 2 studies and so rationalizing choice of dosing in clinical trials.
- Providing biomarkers for monitoring disease progression alongside clinical ratings.
- Monitoring disease mechanisms and measuring inflammation/glial activation in brain disorders.
- Elucidating the downstream effects of drug action.

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