Phosphodiesterase 10A PET Radioligand Development Program: From Pig to Human

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Four novel phosphodiesterase 10A (PDE10A) PET tracers have been synthesized, characterized in preclinical studies, and compared with the previously reported 11C-MP-10. Methods: On the basis of in vitro data, IMA102, IMA104, IMA107, and IMA106 were identified as potential PDE10A radioligand candidates and labeled with either 11C via N-methylation or with 18F through an SN2 reaction, in the case of IMA102. These candidates were compared with 11C-MP-10 in pilot in vivo studies in the pig brain. On the basis of these data, 11C-IMA106 and 11C-IMA107 were taken into further evaluation and comparison with 11C-MP-10 in the primate brain. Finally, the most promising radioligand candidate was progressed into human evaluation. Results: All 5 tracers were produced with good radiochemical yield and specific activity. All candidates readily entered the brain and demonstrated a heterogeneous distribution with the known expression of PDE10A. Baseline PET studies in the pig and baboon showed that 11C-IMA107 and 11C-MP-10 displayed the most favorable tissue kinetics and imaging properties. The administration of selective PDE10A inhibitors reduced the binding of 11C-IMA107 and 11C-MP-10 in the PDE10A-rich brain regions, in a dose-dependent manner. In the nonhuman primate brain, the tissue kinetics of 11C-IMA107 and 11C-MP-10 were well described by a 2-tissue-compartment model, allowing robust estimates of the regional total volume of distribution. Blockade with unlabeled MP-10 confirmed the suitability of the cerebellum as a reference tissue and enabled the estimation of regional binding potential as the outcome measure of specific binding. Conclusion: 11C-IMA107 was identified as the ligand with the highest binding potential while still possessing reversible kinetics. The first human administration of 11C-IMA107 has demonstrated the expected regional distribution and suitably fast kinetics, indicating that 11C-IMA107 will be a useful tool for the investigation of PDE10A status in the living human brain.

Key Words: PET; PDE10A; 11C; in vivo; brain

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in the pig brain. The leading 2 molecules were then evaluated more thoroughly in the nonhuman primate. The most favorable candidate, $^{11}$C-IMA107, was progressed into human testing and used to obtain the first human images.

**MATERIALS AND METHODS**

**Chemistry**

All the precursor and reference compounds were prepared by Mitsubishi Tanabe Pharma Corp. according to procedures previously reported (18).

**In Vitro Measurement of PDE10 Inhibitory Activity**

The human enzyme PDE10A (hPDE10A) was isolated from COS-7 cells transfected with plasmid coding for human PDE10A2, according to the methods described by Kotera et al. (19). The PDE assays were performed according to a modified method reported by Kotera et al. (20). A description of the assay can be found in the supplemental material (available at http://jnm.snmjournals.org).

**Radiochemistry**

In vivo pig and baboon experiments were performed at the Aarhus and Yale PET centers, respectively. $^{11}$C-MP-10 radioisynthesis at both centers was previously reported (9). The radiosyntheses of $^{11}$C-IMA104, $^{11}$C-IMA107, and $^{18}$F-IMA106 were all performed under similar conditions. The labeling procedures of the $^{11}$C- and $^{18}$F-pyrazolopyrimidine compounds $^{11}$C-IMA107 and $^{18}$F-IMA102 are reported in the supplemental material.

**Pig PET Studies**

All animal studies were performed in accordance with the Danish Animal Experimentation Act on a license granted by the Danish Ministry of Justice. A description of the subjects' preparation and monitoring, PET data acquisition, image processing, and blocking experiments can be found in the supplemental material.

**Baboon PET Studies**

The study was approved by the local Institutional Animal Care and Use Committee. A description of the subjects' preparation and monitoring, PET data acquisition, image processing, arterial plasma input function generation, tracer kinetic analysis, and blocking experiments can be found in the supplemental material.

**Human Studies**

The data in this manuscript come from the first 2 scans obtained in an ongoing human study at Imanova Ltd. The Research Ethics Committee approved this study, and all subjects signed a written informed consent form. Each subject underwent a PET scan on a Siemens PET/CT TruePoint Biograph 6-camera system, after the administration of $^{11}$C-IMA107 (147–312 MBq) as a 20-s bolus. The specific activity of all of the radiotracers examined, and only tracer level amount of unknown chemical species was observed. The specific activity of hPDE10A; however, other compounds showed similar affinity in our assay for both the bovine and the human PDE10A. The selectivity of IMA104 was not determined; however, the other compounds showed high selectivity for hPDE10A toward other PDEs, with an IC$_{50}$ greater than 1,000 nM, except for IMA102, which showed a PDE4 IC$_{50}$ of 61 nM. The selectivities of IMA102, IMA107, and MP-10 were also tested in vitro against a variety of CNS receptors and enzymes, and no secondary targets of relevance were seen. For IMA107, the highest-affinity secondary target was 5.8 µM, with a 100,000-fold-lower affinity than PDE10 (Supplemental Table 1).

**Radiochemistry**

All $^{11}$C radiotracers were successfully labeled by $N$-alkylation of the secondary amine precursor with $^{11}$C-methyl iodide (Fig. 1). They could be routinely produced in more than a 2-GBq yield, starting from a typical cyclotron bombardment of 55 µA for 40 min, which corresponds to an estimated 30% decay-corrected radiochemical yield from $^{11}$C-carbon dioxide. The total synthesis times were about 40 min including purification and formulation counted from the end of bombardment. The radiosynthesis of $^{18}$F-IMA102 was achieved by nucleophilic substitution of the mesylate precursor IMA101 with $^{18}$F-fluoride with inversion of the configuration (Fig. 1). The radiochemical yield of this reaction was good, with 11 GBq produced from a 60-min irradiation at 40 µA, which corresponds to an estimated 22% non–decay-corrected yield. The radiochemical purity of all tracers was greater than 95%, as determined by analytic high-performance liquid chromatography with radiodetection. No traces of precursor were detected in the injected doses of all of the radiotracers examined, and only tracer level amount of unknown chemical species was observed. The specific activity of MP-10 precursor included in the following figure, the synthesis of a precursor such as IMA105 and the subsequent $^{11}$C-labeling are depicted. The precursor IMA105 was selected over IMA101 because of the high affinity and low selectivity for PDE10A; however, other compounds showed similar affinity in our assay for both the bovine and the human PDE10A. The selectivity of IMA104 was not determined; however, the other compounds showed high selectivity for hPDE10A toward other PDEs, with an IC$_{50}$ greater than 1,000 nM, except for IMA102, which showed a PDE4 IC$_{50}$ of 61 nM. The selectivities of IMA102, IMA107, and MP-10 were also tested in vitro against a variety of CNS receptors and enzymes, and no secondary targets of relevance were seen. For IMA107, the highest-affinity secondary target was 5.8 µM, with a 100,000-fold-lower affinity than PDE10 (Supplemental Table 1).

**RESULTS**

**In Vitro Affinity Determination**

The selected compounds showed subnanomolar affinity for the hPDE10A, with inhibitory concentration of 50% (IC$_{50}$) values of 0.02, 0.02, 0.79, and 0.06 nM for MP-10, IMA106, IMA102, and IMA107, respectively. An affinity assay of IMA104 was conducted using bovine PDE10A; IMA104 exhibited high affinity, with an IC$_{50}$ value of 0.01 nM. IMA104 was not tested at the
11C-MP-10, 11C-IMA106, 11C-IMA104, 11C-IMA107, and 18F-IMA102 at the end of synthesis was 70 ± 92 (n = 7), 214 (n = 1), 87 (n = 1), 121 ± 47 (n = 3), and 250 GBq/μmol (n = 1), respectively, for the pig study. The specific activity of 11C-MP-10, 11C-IMA106, and 11C-IMA107 at the end of synthesis was 285 ± 207 (n = 16), 19 (n = 1), and 68 ± 24 GBq/μmol (n = 4), respectively, for the primate study.

**Pig PET Studies**

All the radiotracers readily entered the brain and showed a heterogeneous CNS distribution in the pig. PET images of summed radioactivity from 0 to 90 min, showing the regional distribution of the tracers at baseline, are depicted in Figure 2. The brain region demonstrating the highest accumulation and retention of radioactivity was the striatum, consistent with reported qualitative immunohistochemical staining indicating the highest expression of PDE10A in this region (4,21) and consistent with previous experience with other PDE10A radioligands. The highest striatum-to-surrounding brain contrast was seen for 11C-IMA104. The regional tissue time–activity curves for each radiotracer are depicted in Figure 3. For all tracers, the highest uptake was seen in the striatum, followed by the diencephalon, with lower uptake in the frontal cortex and cerebellum, which showed similar levels of uptake. 11C-IMA107, 11C-MP-10, and 18F-IMA102 exhibited a fast washout from all regions of interest (ROIs), whereas 11C-IMA104 and 11C-IMA106 showed slower kinetics in the PDE10A-rich striatum. 11C-IMA106 uptake in the striatum peaked by 35 min and plateaued between 40 and 60 min, displaying a slight decrease from then on. For 11C-IMA104, the striatal signal continued increasing during the 90-min scanning session.

Blocking studies of 11C-MP-10 and 11C-IMA107 were performed using a pharmacologic dose of IMA102 (Fig. 3, time–activity curves). After the administration of the blocker, the uptake of both tracers in the striatum was significantly reduced, reaching the levels of other CNS tissues at the highest dose of the blocker (5 mg/kg).

**Baboon PET Studies**

Representative baseline standardized uptake images for 11C-MP-10, 11C-IMA107, and 11C-IMA106 in the baboon brain are given in Figure 4. All 3 tracers readily entered the baboon brain and showed a regional uptake consistent with our findings in the pig. The time–activity curves depicted in Figure 4 revealed that the rank order of uptake was similar for all tracers, with the putamen > caudate = globus pallidus > nucleus accumbens > remaining ROIs. All 3 tracers displayed a comparable initial uptake in the ROIs, in which the PDE10A protein is expressed at a low level, with a standardized uptake value (SUV) around 1.5 at 5 min after injection. 11C-MP-10 and 11C-IMA107 uptake reached peak levels by 50 and 35 min after administration, respectively, and declined after that. In contrast, 11C-IMA106 tissue kinetics were slower, with no decrease observed in the PDE10A-rich regions during the 120-min scanning time.

The in vivo metabolism of all tracers was determined in all scanning sessions and showed that the 3 tracers were metabolized, albeit, to somewhat different extents (Supplemental Fig. 1). The rate of metabolism of the pyrazolopyrimidine tracers, 11C-IMA107 (~50% of parent at 90 min) and 11C-IMA106 (~56% of parent at 90 min), were slower than that of 11C-MP-10 (~30% of parent at 90 min). All the metabolites observed were more polar than their parent compound.

The 2-tissue-compartment analysis with metabolite-corrected plasma input was able to accurately characterize the kinetic behavior of the 3 tracers (time–activity curves and model fits for 11C-IMA107 can be found in Supplemental Fig. 2). The volume of distribution (Vr) for all regions considered and all 3 tracers are presented in Table 1. The Vr values at baseline for all tracers were highest in the putamen and lowest in all ROIs outside the basal ganglia. 11C-IMA107 and 11C-IMA106 with rate constant for transfer from arterial plasma to tissue (Kt) values at baseline of 0.38 ± 0.13 and 0.413, respectively, showed higher delivery than 11C-MP-10 (0.138 ± 0.05).
The slower kinetics of \( ^{11}C\)-IMA106 were reflected in the higher \( V_T \) values for this tracer. The magnitude of the specific binding component of \( ^{11}C\)-IMA107 and \( ^{11}C\)-MP-10 was assessed after blocking experiments with unlabeled MP-10 at doses of 0.6 and 1.8 mg/kg. A dose-dependent decrease in tracer uptake was observed in the PDE10A-rich regions; the time–activity curves obtained for \( ^{11}C\)-IMA107 and \( ^{11}C\)-MP-10 after administration of the highest dose of MP-10 (1.8 mg/kg) are depicted in Figure 5 and showed a reduction of the tracer uptake in the striatal regions to a level close to that of the cerebellum, as previously reported for \( ^{11}C\)-MP-10 (9).

The \( K_1 \) values for \( ^{11}C\)-IMA107 at baseline (0.38 ± 0.13) and after block (0.428 ± 0.09) were consistent, reflecting an unchanged tracer delivery. The quantification of the \( ^{11}C\)-MP-10 and \( ^{11}C\)-IMA107 binding confirmed that the administration of MP-10 induces a profound decrease in the regional \( V_T \) in the striatum and globus pallidus for both tracers (Table 1). These in vivo blocking studies with unlabeled MP-10 demonstrated that the tracers’ binding is specific to binding to the PDE10A enzyme. The \( V_T \) of the cerebellum did not decrease significantly after treatment and confirmed the suitability of the cerebellum as a reference region and allowed for the regional estimation of \( BP_{ND} \). \( BP_{ND} \) estimates obtained under baseline conditions are given in Table 2. \( ^{11}C\)-IMA106 slow tissue kinetics yielded the highest \( BP_{ND} \) estimates in the PDE10A-rich regions with, for example, a value of 12.71 in the putamen.

The occupancy levels obtained after administration of unlabeled MP-10 were consistent using either radioligand, with occupancy values at the dose of 0.6 and 1.8 mg/kg of 70% and 85% for \( ^{11}C\)-MP-10 and 63% and 85% for \( ^{11}C\)-IMA107.

**Human Studies**

There were no adverse or clinically detectable pharmacologic effects in any of the subjects. No significant changes in vital signs or the results of laboratory studies or electrocardiograms were observed after administration of \( ^{11}C\)-IMA107. Representative images from a baseline scan are shown in Figure 6. The regional distribution images of \( ^{11}C\)-IMA107 in the human brain were similar to those of the pig and monkey brain. After intravenous administration of \( ^{11}C\)-IMA107, radioactivity readily entered the brain, with an SUV of up to 3 in the putamen at peak uptake. The distribution was heterogeneous, with highest concentrations in the putamen and globus pallidus. The time–activity curves from a baseline scan are shown in Figure 6. The kinetics of the uptake of \( ^{11}C\)-IMA107 in the brain were reversible, with a measurable washout of the ligand within the time frame of the PET scans, seen in the regional time–activity curves.

**DISCUSSION**

The aim of this study was to develop a PET radioligand for imaging PDE10A in the human brain in vivo. We previously reported the labeling and evaluation of \( ^{11}C\)-MP-10; however, in parallel to the \( ^{11}C\)-MP-10 evaluation, we also investigated the characteristics of pyrazolopyrimidine PDE10A inhibitors. The 4 pyrazolopyrimidine compounds (as well as \( ^{11}C\)-MP-10) were successfully labeled using \( ^{11}C\)-methyl iodide or \( ^{18}F\)-fluoride and produced with high yield, specific activity, and radiochemical purity. Although the radiosynthesis of \( ^{11}C\)-MP-10 leads to a regiosomer, the radiochemical yields of the \( ^{11}C\)-pyrazolopyrimidine were only slightly better than those obtained for \( ^{11}C\)-MP-10. Analysis of the semipreparative high-performance liquid chromatograms showed that the percentage of product in the crude reaction was around 60% for \( ^{11}C\)-IMA107, compared with 49% for \( ^{11}C\)-MP-10. Labeling of IMA107 with \( ^{18}F\) has not been attempted, however, it is
most likely that this could be achieved using the same labeling route as for IMA102. Although the shorter half-life of $^{11}$C is preferred when multiple scans in the same subject are performed on the same day, the $^{18}$F-labeled analog could be attractive to centers without on-site cyclotron or for distribution.

All the radiotracers were initially evaluated in the pig brain in view of selecting the best candidates for further investigation in the nonhuman primate. The uptake for all candidates was heterogeneous, with high uptake localized to the striatum, consistent with the known expression of the enzyme. The tracers’ kinetics and striatum-to-cerebellum ratio at 90 min seen in vivo correlated well with the in vitro affinity. $^{11}$C-IMA107, $^{11}$C-MP-10, and $^{11}$C-IMA106 exhibited suitably fast kinetics in the pig brain. The other candidates were less promising, with a low striatum–to–rest-of-brain ratio in the case of $^{18}$F-IMA102 and slow kinetics that did not appear favorable for quantification in the case of $^{11}$C-IMA104. In vivo blocking studies with unlabeled IMA102 demonstrated that the uptake of $^{11}$C-IMA107 and $^{11}$C-MP-10 in the striatum was significantly reduced after administration of a PDE10A inhibitor, suggesting that both the pyrazolopyrimidine compound series and the MP-10 demonstrate specific binding for PDE10A.

On the basis of these results, $^{11}$C-MP-10, $^{11}$C-IMA107, and $^{11}$C-IMA106 were selected for in vivo evaluation in higher species. In the baboon brain, all 3 tracers displayed slower brain kinetics; this was particularly noticeable for $^{11}$C-IMA106 (Fig. 4C). Although quantification was possible in the baboon with this tracer, the slower kinetics suggested that it was not an ideal candidate for translation to humans. As a result, the binding selectivity of $^{11}$C-IMA106 was not explored. However, the higher affinity of $^{11}$C-IMA106 that led to a higher signal-to-noise ratio may facilitate imaging of extrastriatal regions such as the substantia nigra, in which PDE10A expression is known to be significantly lower. On the other hand, $^{11}$C-IMA107 and $^{11}$C-MP-10 demonstrated good brain penetration, reversible kinetics, and a good specific signal that could be reduced in a dose-dependent manner after competition with unlabeled MP-10. It is worth mentioning that the kinetics of both tracers in the globus pallidus appeared different from those of other basal ganglia regions, displaying a slower washout. We do not believe this phenomenon can be explained by the tracers or a metabolite binding to a different target in this region, because this effect was observed with both tracers, under both baseline and homologous competition conditions.

Quantitative analysis of $^{11}$C-MP-10 and $^{11}$C-IMA107 binding in the baboon brain determined that both ligands possessed good in vivo radioligand characteristics for imaging PDE10A. Analyses of data before and after competition with a PDE10A inhibitor confirmed the specificity of the $^{11}$C-IMA107 and $^{11}$C-MP-10 binding and the suitability of the cerebellum as a reference region for the determination of the regional estimation of $BP_{ND}$, as previously indicated for $^{11}$C-MP-10 (9), this implies that the quantitative analysis of PDE10A PET ligands may not necessitate the accurate measurement of plasma radioactivity and metabolites. The occupancy levels in the striatum calculated for the pharmacologic doses of MP-10 were similar for both radioligands. The high occupancy of 85% measured at the 1.8 mg/kg dose confirmed the specificity of the radioligand binding in this region and indicated that changes in the PDE10A availability in the striatum can be quantitatively measured using either radioligand. The globus pallidus was, however, excluded from the occupancy determination because the occupancy values in this region were similar but lower for the 2 tracers at 68% and 69%, confirming the

<table>
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<tr>
<th>ROI</th>
<th>$^{11}$C-MP-10 Baseline</th>
<th>$^{11}$C-MP-10 1.8 mg/kg</th>
<th>$^{11}$C-IMA107 Baseline</th>
<th>$^{11}$C-IMA107 1.8 mg/kg</th>
<th>$^{11}$C-IMA106, baseline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Putamen</td>
<td>9.62 ± 1.75</td>
<td>3.39</td>
<td>22.48 ± 0.15</td>
<td>7.34</td>
<td>68.13</td>
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<tr>
<td>Caudate</td>
<td>7.03 ± 0.82</td>
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<td>5.70</td>
<td>41.03</td>
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<tr>
<td>Globus pallid</td>
<td>6.38 ± 1.21</td>
<td>3.47</td>
<td>16.66 ± 0.46</td>
<td>8.52</td>
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<tr>
<td>Nucleus accumbens</td>
<td>4.33 ± 0.92</td>
<td>2.57</td>
<td>8.49 ± 0.14</td>
<td>4.45</td>
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<tr>
<td>Insula</td>
<td>3.84 ± 0.38</td>
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<td>5.71 ± 0.03</td>
<td>4.56</td>
<td>6.65</td>
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<tr>
<td>Cingulate</td>
<td>3.46 ± 0.39</td>
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<td>4.31 ± 0.49</td>
<td>4.10</td>
<td>5.39</td>
</tr>
<tr>
<td>Frontal cortex</td>
<td>3.14 ± 0.30</td>
<td>2.27</td>
<td>4.10 ± 0.26</td>
<td>3.84</td>
<td>5.66</td>
</tr>
<tr>
<td>Thalamus</td>
<td>3.04 ± 0.32</td>
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<td>5.42 ± 0.78</td>
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<tr>
<td>Pons</td>
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<td>Brain stem</td>
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<td>Cerebellum</td>
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<td>4.33 ± 0.72</td>
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<td>4.97</td>
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Baseline values are mean and SD of 12 and 2 scans for $^{11}$C-MP-10 and $^{11}$C-IMA107, respectively.
differential behavior of both tracers in this region, compared with the striatum. We cannot exclude regional differences between the nonspecific binding in the globus pallidus and striatal regions.

Qualitative assessment of $^{11}$C-MP-10 uptake in the regions of low or negligible PDE10A expression indicated that the distribution of $^{11}$C-MP-10 appears slightly different from that of the pyrazolopyrimidine tracers, with a higher relative uptake in the insula and cingulate (Fig. 4). The quantitative analysis confirmed the higher uptake of $^{11}$C-MP-10 in these regions, with $BP_{ND}$ values of 0.41 and 0.57 in the cingulate and insula, respectively, compared with those of the pyrazolopyrimidine tracers, with $BP_{ND}$ values in the cingulate and insula of 0.00 and 0.05 for $^{11}$C-IMA107 and 0.08 and 0.34 for $^{11}$C-IMA106, respectively. The uptake of $^{11}$C-MP-10 in these regions was also blocked after administration of MP-10 (1.8 mg/kg), with $BP_{ND}$ values in the cingulate and insula reduced to 0.03 and 0.04, respectively ($P$ values calculated using a 1-tail t test $\leq 0.02$). The magnitude of the signal change in these regions was similar to that observed in the putamen and caudate. However, blocking experiments performed with a structurally different PDE10A inhibitor (data not shown) showed that a dose producing over 95% occupancy in the striatal regions did not significantly reduce the $^{11}$C-MP-10 $BP_{ND}$ values in the insula and cingulate ($P$ values calculated using a 1-tail t test $\geq 0.69$). The $^{11}$C-MP-10 signal reduction in these regions only under homologous pharmacologic challenge may indicate a higher specificity of $^{11}$C-IMA107, which did not show any reduction of signal in the cingulate and insula after treatment with MP-10. In addition, an $^{11}$C-IMA107 effective binding potential, $BP_{ND}$, of 4.26 ± 0.84 in the putamen provides an excellent signal-to-noise ratio, which was superior to that of $^{11}$C-MP-10. For these reasons, it was decided to progress $^{11}$C-IMA107 to humans. After intravenous administration of $^{11}$C-IMA107 in humans, the regional distribution of radioactivity in the brain was consistent with the distribution previously observed in the pig and baboons. The tracer kinetics were also reversible, indicating a good translation of the tracer behavior between all species studied. Work is in progress to fully characterize $^{11}$C-IMA107 in humans. Early indications are that it has good signal to noise, with suitably fast kinetics and consistent good kinetics model fits and can be quantified using a simplified reference tissue model.

**CONCLUSION**

Four new radioligand candidates for the PDE10A were successfully labeled and compared in vivo in the pig and baboon brains with the previously reported radioligand $^{11}$C-MP-10. $^{11}$C-IMA107 proved to be the most suitable for imaging and quantifying PDE10A in preclinical studies. Its initial assessment in humans provides evidence that $^{11}$C-IMA107 is a promising radioligand for probing the PDE10A in vivo by PET.

**DISCLOSURE**

The costs of publication of this article were defrayed in part by the payment of page charges. Therefore, and solely to indicate this fact, this article is hereby marked “advertisement” in accordance with 18 USC section 1734. Part of this research project was performed at the GSK Clinical Imaging Centre before transfer of management to Imanova Ltd. This work was also supported by Mitsubishi Tanabe Pharma Corporation, and the human PET imaging in this project was supported by the National Institute for Health Research (NIHR) Biomedical Research Centre for Mental Health at South London and Maudsley NHS Foundation Trust and the Institute of Psychiatry, King’s College London. This article presents independent research and the views expressed are those of the authors and not necessarily

**TABLE 2**

<table>
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<th>ROI</th>
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<tr>
<td>Brain stem</td>
<td>−0.17</td>
<td>0.04</td>
<td>0.25</td>
<td>0.05</td>
<td>0.20</td>
</tr>
</tbody>
</table>

**FIGURE 6.** Summed PET images in human brain and representative baseline tissue time–activity curves for $^{11}$C-IMA107 for selected ROIs. Slices are displayed as SUV units at level of striatum in transverse, sagittal, and coronal orientations.
those of the NHS, the NIHR, or the Department of Health. No other potential conflict of interest relevant to this article was reported.

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