

SUPERINTENDENCIA DE INDUSTRIA Y COMERCIO
DIRECCIÓN DE NUEVAS CREACIONES

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Bogotá, D.C.,

Abogado
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SUPERINTENDENCIA DE INDUSTRIA Y COMERCIO



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Act. 519 TRASLADOINFORME Folios: 15

ASUNTO	Radicación	09-22307
	Trámite	011
	Evento	379
	Actuación	519
	Oficio	<u>16674</u>
	Folios	015

Patente de Invención	☒	Patente de Modelo de Utilidad	☐
Vía Nacional	☒	Cap. I	☐
Vía PCT (fase nacional)	☒	Cap. II	☒

No. Solicitud Internacional PCT/EP2007/006148
Fecha de Presentación Internacional 06 de julio de 2007

Como resultado del Examen de Fondo realizado, con fundamento en el Artículo 45 de la Decisión 486 de la Comisión de la Comunidad Andina, se comunica:

1. Documentos estudiados

Descripción:	Folios 74-137	04 de marzo de 2009
Reivindicaciones:	Folios 138-153	04 de marzo de 2009
Prioridad:	EP 080184455.3	4 de septiembre 2008

2. Análisis de la Prioridad

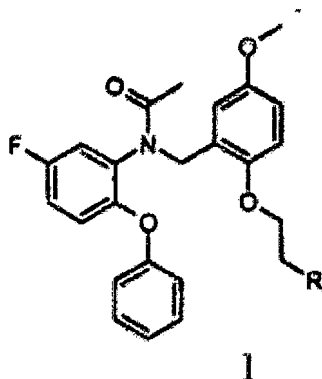
Se acepta la reivindicación de Prioridad porque esta solicitud fue presentada dentro del plazo establecido (Art. 9 D 486) y su contenido técnico corresponde con el de la prioridad reivindicada.

3. Objeto de la invención

Título de la solicitud: "derivados de la feniloxiamina"

El problema planteado en esta solicitud consiste en la necesidad de proporcionar compuestos con actividad sobre los receptores periféricos de las benzodíacepinas

Para resolver este problema técnico la solicitud en estudio proporciona un compuesto de la fórmula (I) y composiciones que lo contienen.



El objeto de la presente Solicitud se relaciona con la Clasificación Internacional de Patentes (CIP): C07C 309/66; C07C 233/88; C07F 7/18

4. Reivindicaciones relativas a un uso o a un segundo uso (Art 14 y 21 D 486)

El uso de las reivindicaciones 31-33 no es patentable, de acuerdo con el Art 14.

5. De la Solicitud de Patente

Reivindicaciones (Art 30 D 486, Art 22 PCT)

Las reivindicaciones 1-7 no son concisas porque los términos:

- Alquilo, perhaloalquilo, di-alquil, tri-alquil, alcoxi, alilo: porque no se define el número de carbonos del grupo.
- Aralquilo, alquilfenilo, cicloalquilo: porque no define el número de carbonos del anillo.
- Heteroarilo: no define el número de miembros del anillo, y el número y tipo de heteroátomos que conforman el anillo.
- C₁₋₆ alquilfenilo, 2,4,6-C₁₋₆ trialquil, C₁₋₅ alquilo porque dan lugar a un alto número de alternativas que no encuentran soporte en el capítulo descriptivo.

No permiten entonces, limitar el alcance de la materia que se busca proteger ni definirla adecuadamente, por lo tanto se requiere al solicitante que restrinja la solicitud a una razonable generalización del tipo de compuestos que han sido sintetizados o probados.

Las reivindicaciones 21 al 24 definen un proceso para la preparación de un compuesto de fórmula uno, pero no definen el proceso por sus características esenciales (catalizadores, condiciones de reacción, solventes, etapas, esta última es la única que se encuentra en la definición del proceso), se sugiere definir el proceso de manera adecuada, y retirar de éstas reivindicaciones las definiciones de los radicales, las cuales pueden ir referidas en las reivindicaciones dependientes que definieron el compuesto de fórmula uno.

La reivindicación 31 está definida de manera inadecuada puesto que establece como característica un uso, lo cual no es patentable, por lo tanto se sugiere que sea retirada del capítulo reivindicatorio.

6. Determinación del Estado de la Técnica

El Estado de la Técnica se determinó con base en la fecha de Presentación de la Solicitud de Prioridad: 04 de septiembre de 2006

Y se encontraron los siguientes documentos que anticipan el objeto de esta solicitud:

Item	Documento	Título	Fecha de Publicación*
D1	Journal Of Medicinal Chemistry Volume 47, Page 2228-2235 DOI: 10.1021/jm0304919	Development of a New Radioligand, N-(5-Fluoro-2-phenoxyphenyl)-N-(2-[¹⁸ F]fluoroethyl-5-methoxybenzyl)acetamide, for PET Imaging of Peripheral Benzodiazepine Receptor in Primate Brain	March 26, 2004 (Article)

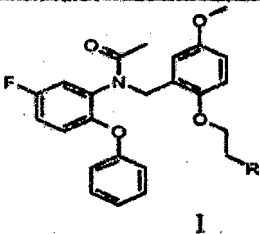
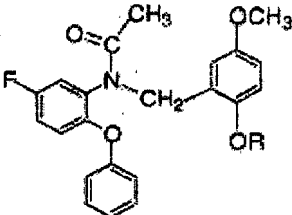
D1
Journal Of Medicinal Chemistry Chemistry, Volume 47, (folio 198-205, página 2228-2235)
Divulga el compuesto 5 que corresponde a un compuesto de fórmula (I) en el que R = ¹⁸F el cual es afín con el Receptor periférico de las benzodiazepinas y puede ser utilizado como radio ligando (página 2228, columna 2, Esquema 1)

7. Examen de Patentabilidad (Art 14 D486)

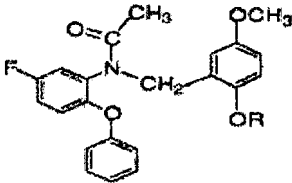
Nivel Inventivo (Art18 D486)

La reivindicación 1 consiste en un compuesto de fórmula (I)

Comparación de las características técnicas **esenciales**, con las del Estado de la Técnica:

Características esenciales	D1
Un compuesto de fórmula (I)	Un compuesto de fórmula (I)
 I	
R es seleccionado de Cloro, Bromo, Iodo	Un compuesto 5 donde R es -CH ₂ CH ₂ F, un compuesto 6 donde R es -CH ₂ I
R puede ser un grupo -OS(O) ₂ -L	Un compuesto 7 donde R es tosiloxietyl

El documento D1 se considera el estado de la técnica cercano a la invención definida en la reivindicación 1 porque se refiere al desarrollo de nuevos radioligandos para los



receptores periféricos de benzodiazepinas (PBR) de fórmula (I) donde R es H (compuesto 3), FCH₂ (compuesto 4), FCH₂CH₂ (compuesto 5), ICH₂ (compuesto 6), tosiloxietyl (compuesto 11), trfluorometanosulfoniloxyetyl bromuro (compuesto 12), (folios 198-201, columna de la derecha, estructura y tabla 1)

La diferencia entre la invención, definida en la reivindicación 1 y los compuestos de D1 consiste en que la actual reivindicación presenta análogos 2-cloroetil-5-

metoxibencil, 2-bromoetil-5-metoxibencil y 2-iodoetil-5-metoxibencil, de los compuestos de fórmula I.

El efecto técnico que logra la diferencia, consiste en que se obtienen compuestos de fórmula I que incluyen halógenos diferentes al flúor en la estructura de los compuestos de D1, ya que el capítulo descriptivo no revela que los compuestos reivindicados tengan mayor afinidad por los receptores PBR, por lo tanto no se puede concluir que los compuestos de la actual solicitud tengan una mejora técnica inesperada sobre los compuestos revelados en D1.

Por lo tanto, el Problema Técnico Objetivo que pretende resolver esta invención se puede formular así: "cómo proporcionar compuestos alternativos a los conocidos en D1 y que sean útiles como ligandos de los PBR".

Sin embargo, la utilización de la introducción de radicales etil-halógeno, o un grupo alquilo unido a un grupo sulfonilo saliente ya se encuentra divulgada en D1 (folio 200 compuestos 5, 7) los cuales se refieren a la sustitución por 2 fluoroetil-5-metoxifenil y tosiloxietyl análogos respectivamente.

En consecuencia, la persona del oficio normalmente versada en la materia enfrentada al problema de presentar compuestos alternativos derivados de la feniloxianilina, intercambiaría el flúor del grupo 2-halo-etil-5-metoxifenil por cloro, bromo o iodo, intercambiaría el grupo halo por un grupo sulfonilo saliente como en el compuesto 7 llegando así al objeto de la presente solicitud. Por lo cual se considera obvia.

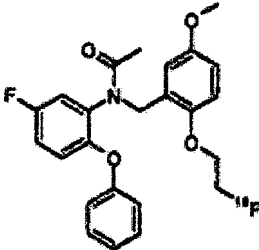
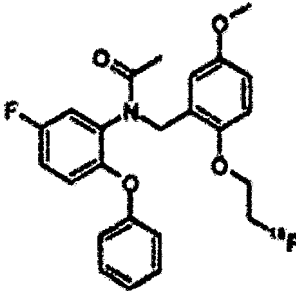
Conclusión:
La reivindicación 1 cumple el requisito de novedad (Art. 16), pero no cumple con el requisito de nivel inventivo (Art. 18).

Puesto que las reivindicaciones dependientes 2-6 no mencionan características inventivas adicionales, por lo que tampoco se pueden considerar inventivas.

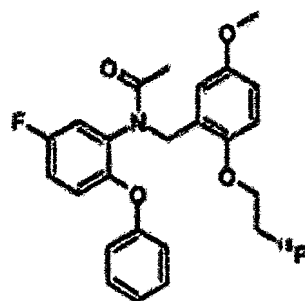
Conclusión:
Las reivindicaciones 1-20 y 29-30 cumplen el requisito de novedad (Art. 16), pero no cumplen con el requisito de nivel inventivo (Art. 18).

La reivindicación 25 consiste en un proceso para formar el compuesto de fórmula 3

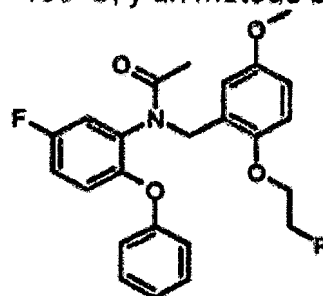
Comparación de las características técnicas **esenciales**, con las del Estado de la Técnica:

Características esenciales	D1
Un proceso para preparar un compuesto de fórmula 3	Un proceso para preparar un compuesto de fórmula 5
 (3)	
Literal a)	Metodo 2

La diferencia entre la invención, definida en la reivindicación 25 y los procesos de D1 consiste en que la actual reivindicación presenta un método a) donde se hace

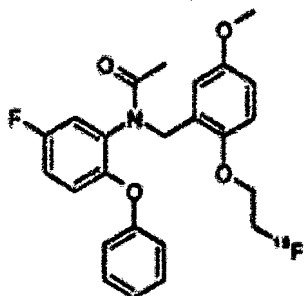


reaccionar un compuesto análogo tosiloimetil de fórmula 7 con aniones ^{18}F a unas condiciones de temperatura entre 121°C - 160°C ; y un método b)



donde se hace reaccionar un compuesto de fórmula I con aniones ^{18}F a unas condiciones de temperatura entre 50°C - 160°C .

El efecto técnico que logra la diferencia, consiste en que se obtiene un compuesto de



fórmula 3

partiendo de intermediarios alternativos.

Por lo tanto, el Problema Técnico Objetivo que pretende resolver esta invención se puede formular así: "cómo presentar un proceso alternativo para obtener el compuesto de fórmula (5) conocido en D1 partiendo de intermediarios alternativos"

Sin embargo, la preparación de un compuesto de fórmula 5 a partir de un compuesto de tosiloxi sustituido o a partir de un compuesto halo alquil sustituido saliente ya se encuentra divulgado en D1 (folio 199, página 2229 columna derecha último párrafo) el cual se refiere a un proceso donde se obtiene un compuesto de fórmula 5 a partir de precursores tosiloxi sustituidos (literal a)) o haloalquil sustituidos (literal b)).

En consecuencia, la persona del oficio normalmente versada en la materia enfrentada al problema de presentar procesos alternativos para la formación del compuesto de fórmula 5, tomaría las enseñanzas de D1 haciendo reaccionar aniones ^{18}F con análogos tosiloxi sustituidos (como el tosiloimetil) o haloetil sustituido, para obtener el compuesto 5. Por otro lado, una persona normalmente versada como resultado de la experimentación encontraría la temperatura óptima. De esta manera no puede atribuirse nivel inventivo al objeto de la reivindicación 25.

La reivindicación 25 cumple el requisito de novedad (Art. 16), pero no cumple con el requisito de nivel inventivo (Art. 18).

Puesto que las reivindicaciones dependientes 26-28 no mencionan características inventivas adicionales, por lo que tampoco se pueden considerar inventivas.

Conclusión:

Las reivindicaciones 1-6, 25-28 el requisito de novedad (Art. 16), pero no cumplen con el requisito de nivel inventivo (Art. 18).

8. Conclusión

Los requisitos de Patentabilidad de la presente Solicitud están afectados así:

Item	Documento	Reivindicaciones afectadas	Requisito que afecta
D1	Journal Of Medicinal Chemistry Volume 47, Page 2228-2235 DOI: 10.1021/jm0304919	1-6, 25-28	Nivel Inventivo

En cumplimiento del artículo 45 de la Decisión 486 de la Comisión de la Comunidad Andina, se le indica que debe dar respuesta dentro del término de sesenta (60) días contados a partir de la fecha de notificación de la presente comunicación. En caso de no dar respuesta al presente requerimiento, o si a pesar de la respuesta subsistieran los impedimentos para la concesión, se denegará la patente.

Notifíquese el presente oficio conforme a lo dispuesto en el numeral 5.2, literal e), del capítulo quinto del título primero de la Circular Única No. 10 de 2001.


JOSÉ LUIS SALAZAR LÓPEZ
Director de Nuevas Creaciones (C)

El anterior requerimiento fue notificado por fijación en lista, de fecha
125 SET. 2012

Elaborado por: Harvey A. Jaramillo
Revisado Por: Jacqueline Alfonso *OKU*

Development of a New Radioligand,

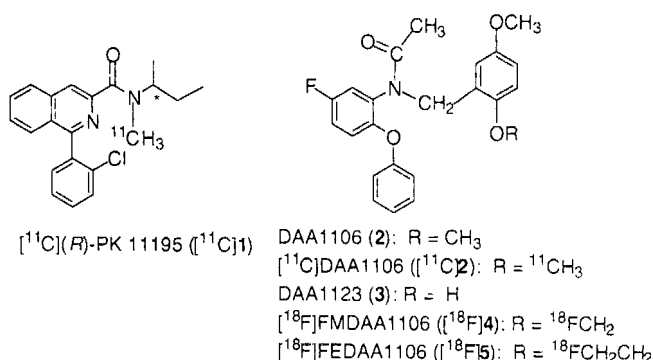
N-(5-Fluoro-2-phenoxyphenyl)-N-(2-[¹⁸F]fluoroethyl-5-methoxybenzyl)acetamide, for PET Imaging of Peripheral Benzodiazepine Receptor in Primate BrainMing-Rong Zhang,^{1,2,*} Jun Maeda,^{2,§} Masanao Ogawa,^{1,§} Junko Noguchi,^{1,§} Takchito Ito,^{1,§} Yuichiro Yoshida,^{1,§} Kazutoshi Okajima,^{2,§} Shigeru Ohayashi,² Tetsuya Suhara,² and Kazutoshi Suzuki¹¹Department of Medical Imaging, National Institute of Radiological Sciences, 4-9-1 Anagawa, Inage-ku,Chiba 263-8525, Japan; ²Japan Brain Imaging Project, National Institute of Radiological Sciences, 4-9-1 Anagawa, Inage-ku,

Chiba 263-8525, Japan, and SHI Accelerator Service Co. Ltd., 5-9-11 Kitashinagawa, Shinagawa-ku, Tokyo 141-8686, Japan

To develop a positron emission tomography (PET) ligand for imaging the peripheral benzodiazepine receptor (PBR) in brain and elucidating the relationship between PBR and brain diseases, four analogues (**4**–**7**) of *N*-(2,5-dimethoxybenzyl)-*N*-(5-fluoro-2-phenoxyphenyl)-acetamide (**2**) were synthesized and evaluated as ligands for PBR. Of these compounds, fluoromethyl (**4**) and fluoroethyl (**5**) analogues had similar or higher affinities for PBR than the parent compound **2** ($K_i = 0.16$ nM for PBR in rat brain sections). Iodomethyl analogue **6** displayed a moderate affinity, whereas tosyloxyethyl analogue **7** had weak affinity. Radiolabeling was performed for the fluoroalkyl analogues **4** and **5** using fluorine-18 (¹⁸F, β^+ ; 96.7%, $T_{1/2} = 109.8$ min). Ligands [¹⁸F]**4** and [¹⁸F]**5** were respectively synthesized by the alkylation of desmethyl precursor **3** with [¹⁸F]fluoromethyl iodide ([¹⁸F]**8**) and 2-[¹⁸F]fluoroethyl bromide ([¹⁸F]**9**). The distribution patterns of [¹⁸F]**4** and [¹⁸F]**5** in mice were consistent with the known distribution of PBR. However, compared with [¹⁸F]**5**, [¹⁸F]**4** displayed a high uptake in the bone of mice. The PET image of [¹⁸F]**4** for monkey brain also showed significant radioactivity in the bone, suggesting that this ligand was unstable for in vivo defluorination and was not a useful PET ligand. Ligand [¹⁸F]**5** displayed a high uptake in monkey brain especially in the cerebral cortex, a region with richer PBR than the other regions in the brain. The radioactivity level of [¹⁸F]**5** in monkey brain was 1.5 times higher than that of [¹¹C]**2**, and 6 times higher than that of (*R*)-(1-(2-chlorophenyl)-*N*-[¹¹C]methyl-*N*-(1-methylpropyl)isoquinoline ([¹¹C]**1**). Moreover, the in vivo binding of [¹⁸F]**5** was significantly inhibited by PBR-selective **2** or **1**, indicating that the binding of [¹⁸F]**5** in the monkey brain was mainly due to PBR. Metabolite analysis revealed that [¹⁸F]**4** was rapidly metabolized by defluorination to [¹⁸F]F[−] in the plasma and brain of mice, whereas [¹⁸F]**5** was metabolized by debenzoylation to a polar product [¹⁸F]**13** only in the plasma. No radioactive metabolite of [¹⁸F]**5** was detected in the mouse brain. The biological data indicate that [¹⁸F]**5** is a useful PET ligand for PBR and is currently used for imaging PBR in human brain.

Introduction

The peripheral benzodiazepine receptor (PBR), which has been identified in the peripheral tissues, is located on the mitochondrial outer membrane in several organs including the kidney, nasal epithelium, lung, heart, and endocrine organs such as the adrenal, testis, and pituitary gland.^{1–3} Later studies further demonstrated the presence of PBR in the central nervous system.^{4–12} The PBR density in the brain was increased in injured brain, and this increase has been used as an indicator of neuronal damage⁷ and several neurodegenerative disorders, such as Alzheimer's disease,⁸ Huntington's disease,⁹ Wernicke's encephalopathy,¹⁰ multiple sclerosis,¹¹ and stroke.¹² In addition, a relationship between alterations of PBR and epileptic foci have been proposed.^{13–15} These findings have resulted in the de-

Scheme 1. Chemical Structures of [¹¹C]**1** and [¹¹C]**2** Analogues

velopment of a radioligand labeled by a positron-emitter, which has made possible to visualize the distribution of PBR in the brain.^{15–19}

(*R*)-(1-(2-Chlorophenyl)-*N*-[¹¹C]methyl-*N*-(1-methylpropyl)isoquinoline ([¹¹C](*R*)-PK 11195, [¹¹C]**1**, Scheme 1) is a clinically useful positron emission tomography (PET) ligand for imaging PBR in cases of several brain

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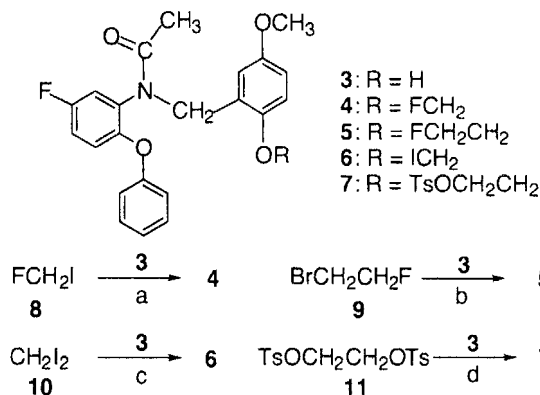
cases.^{23–25} However, the signal of [¹¹C]**1** in the monkey brain was not high enough for quantitative analysis using PET because of its low uptake into the brain.²³ Therefore, the precise characterization of PBR in monkey brain awaits development of new PET ligands with improvements over [¹¹C]**1**. In 1998, *N*-(2,5-dimethoxybenzyl)-*N*-(5-fluoro-2-phenoxyphenyl)acetamide (DAA1196, **2**, Scheme 1) was reported as a potent and selective ligand for PBR by the Taisho Pharmaceutical's group.^{23–27} Compound **2** had a high affinity for PBR in mitochondrial fractions of rat ($K_i = 0.043$ nM) and monkey ($K_i = 0.188$ nM) brains.²⁵ Moreover, **2** had weak or negligible affinities ($IC_{50} > 10$ μ M) for other receptors, ion channels, uptake/transporters, and second messengers.^{23–27} Thus, we previously synthesized [¹¹C]**2**²⁸ as a PET ligand for PBR by reacting the desmethyl precursor (*N*-(5-fluoro-2-phenoxyphenyl)-*N*-(2-hydroxy-5-methoxybenzyl)acetamide²⁴) (**3**) with [¹¹C]-methyl iodide in vitro and in vivo evaluation using mouse. In a monkey demonstration that [¹¹C]**2** had high affinity binding with PBR in the brains and about a 3-fold higher uptake of radioactivity in the monkey brain than [¹¹C]**1**.²⁸ Now [¹¹C]**2** is being used to investigate PBR in the human brain in our facility.

In general, the short half-life of a ¹¹C-labeled ligand ($t_{1/2} = 20.4$ min) often limits its usefulness in a dynamic PET experiment has a turnover time longer than 100 min. Since ¹⁸F has advantages over ¹¹C, with a longer half-life (110 min vs. 20 min) and a lower positron energy (650 keV vs 960 keV), an ¹⁸F-labeled ligand produces higher quality images with a higher spatial resolution in PET measurements. Moreover, ¹⁸F is more convenient for time storage and long distance transportation to other facilities. To develop ¹⁸F-labeled compounds with improved properties compared to [¹¹C]**2**, we previously prepared two [¹⁸F]fluoroalkyl analogues (**4**, **2**-(*N*-(5-fluoro-2-phenoxyphenyl)-*N*-(2-(¹⁸F)fluoromethyl-5-methoxybenzyl)acetamide ([¹⁸F]FM-DAA1196, [¹⁸F]**4**) and *N*-(5-fluoro-2-phenoxyphenyl)-*N*-(2-(¹⁸F)fluoroethyl-5-methoxybenzyl)acetamide ([¹⁸F]EE-DAA1196, [¹⁸F]**5**) as putative radioligands for PBR (Scheme 1). Preliminary studies revealed that analogues **4** and **5** have similar or higher inhibitory activity (IC_{50}) on PET in the rat brain than **2**.²⁴ The ex vivo autoradiograms of [¹⁸F]**4** and [¹⁸F]**5** binding sites in rat brain showed significantly high radioactivity in the cerebral cortex and cerebellum, the richer regions of PBR compared with other brain regions.²⁴

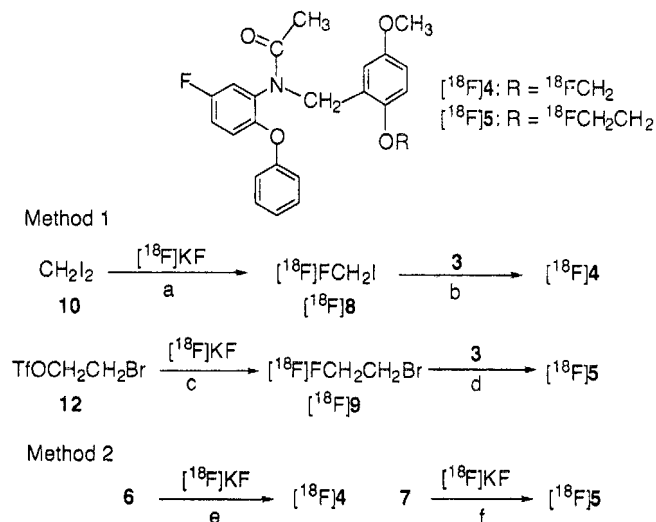
In this study, we synthesized four analogues (**4**–**7**, Scheme 2), including two novel compounds: (*N*-(5-fluoro-2-phenoxyphenyl)-*N*-(2-iodomethyl-5-methoxybenzyl)acetamide (**6**) and (*N*-(5-fluoro-2-phenoxyphenyl)-*N*-(5-tosyloxyethylbenzyl)acetamide (**7**). Using autoradiography technique, we measured the binding affinities (K_i) of **4**–**7** to PBR and the central benzodiazepine receptor (CBR). We labeled the fluoroalkyl analogues **4** and **5** using ¹⁸F and prepared the radioactive products in reproducible radiochemical yields. We examined the regional distribution of [¹⁸F]**4** and [¹⁸F]**5** in mice and measured the percentages of unmetabolized [¹⁸F]**4** and [¹⁸F]**5** in the plasma and brain. Using PET, we determined the uptake of [¹⁸F]**4** and [¹⁸F]**5** and elucidated the specific binding of [¹⁸F]**5** to PBR in the monkey brain.

Scheme 2. Chemical Synthesis of Compounds **4**–**7**.

199



^a Reagents and conditions: (a) NaH, DMF, 0 °C, 5 h, 79%; (b) NaH, DMF, 80 °C, 8 h, 65%; (c) NaH, DMF, 25 °C, 3 h, 34%; (d) NaH, DMF, 0 °C, 5 h, 46%.

Scheme 3. Radiosynthesis of [¹⁸F]**4** and [¹⁸F]**5** by Two Methods^a

^a Reagents and conditions: (a) 110 °C, 10 min, 8%; (b) NaH, DMF, −15 °C, 3 min, 90%; (c) *o*-dichlorobenzene, 130 °C, 5 min, 55%; (d) NaH, DMF, −15 °C, 10 min and then 120 °C, 10 min, 82%; (e) DMF, 30–120 °C, 1–15 min, 2–37%; (f) DMF, 30–120 °C, 1–15 min, 2–60%.

Results and Discussion

Chemistry. The four analogues **4**–**7** were synthesized according to the reaction sequences linearized in Scheme 2. Reaction of the desmethyl precursor **3**^{24–27} with excess fluoromethyl iodide³¹ (FCH₂I, **8**) in the presence of NaH at 0 °C for 5 h gave the fluoromethyl analogue **4**. Treatment of **3** with 2-fluoroethyl bromide (FCH₂CH₂Br, **9**) and NaH at 80 °C gave the fluoroethyl analogue **5**. The iodomethyl analogue **6** was prepared by the reaction of **3** with diiodomethane (CH₂I₂, **10**) and NaH at 25 °C for 3 h. The tosyloxyethyl analogue **7** was obtained by treating **3** with excess ethylene ditosylate (**11**) and NaH at 0 °C.

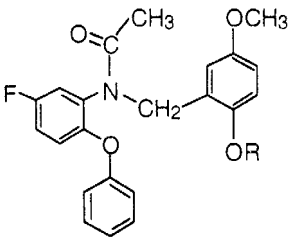
Radiosynthesis. Ligands [¹⁸F]**4** and [¹⁸F]**5** were prepared by two methods according to Scheme 3. The first method was a two step reaction sequence which involved the preparation of the radioactive intermediate [¹⁸F]fluoromethyl iodide ([¹⁸F]FCH₂I, [¹⁸F]**8**) or 2-[¹⁸F]fluoroethyl bromide ([¹⁸F]FCH₂CH₂Br, [¹⁸F]**9**), followed by the alkylation of **3** with [¹⁸F]**8** or [¹⁸F]**9**. The second method was direct nucleophilic replacement of the iodomethyl analogue **6** or tosyloxyethyl analogue **7**

with [¹⁸F]F₂. As for the two-step method, the alkylating intermediate [¹⁸F]8 or [¹⁸F]9 was prepared by the reaction of [¹⁸F]F₂ with 10 or 2-trifluoromethanesulfonyloxethyl bromide (BrCH₂CH₂OTf, 12) by using a newly developed automated system.^{32,33} After the fluorination reaction finished, [¹⁸F]8 or [¹⁸F]9 was purified by distillation and directly trapped in a solution of DMF containing 3 and NaH at -15 °C. The purification of [¹⁸F]8 (bp: 53–54 °C) or [¹⁸F]9 (bp: 71.5 °C) by distillation was effective, because this procedure can leave behind all nonvolatile impurities such as metal ions from the cyclotron target, the unreacted [¹⁸F]F₂, and the nucleophilic displacement reagent Kryptofix 222 K⁺CO₃. After the nucleophilic [¹⁸F]8 or [¹⁸F]9 trapping ended, the subsequent benzomethylation finished perfectly, while the subsequent demethylation required a further 10 min at 120 °C. As for the direct method, heating 6 and 7 with [¹⁸F]F₂ at 30–120 °C gave [¹⁸F]4 and [¹⁸F]5. However, the radiochemical yields were not reproducible (2%–40%), and purifying [¹⁸F]4 and [¹⁸F]5 from the reaction mixtures was often difficult since the many impurities resulting from the target and reaction greatly reduced the efficiency. After comparing the reaction conditions and radiochemical yields, we synthesized [¹⁸F]4 and [¹⁸F]5 by the two-step reaction method (Method 1) via the intermediates [¹⁸F]8 and [¹⁸F]9, respectively.

Reversed phase semipreparative HPLC purification of the reaction mixtures gave [¹⁸F]4 and [¹⁸F]5 in 45 ± 3% and 40 ± 4% and 12 ± 4% (*n* = 4) radiochemical yields (radiochemical yield corrected for the physical decay in a sample's time of 63 ± 4 min and 50 ± 3 min). The structure of the desired product was confirmed by coinjection with the corresponding nonradioactive 4 or 5 on reversed phase analytic HPLC. In the final product solutions, the radiochemical purity of [¹⁸F]4 or [¹⁸F]5 was higher than 98%, and the specific activity was 40–65 GBq/μmol for [¹⁸F]4 and 110–145 GBq/μmol for [¹⁸F]5, determined from the mass measured by HPLC-UV analysis. No significant peak corresponding to 3 was observed in the HPLC for the final products. Moreover, the radiochemical purities of [¹⁸F]4 and [¹⁸F]5 remained >95% after 180 min at 25 °C, and they were stable as PET ligands while performing the evaluation.

In Vitro Binding Assays. The in vitro binding affinities (*K*_i) of the four analogues 4–7 for PBR were determined from competition for the [¹¹C]2 binding to PBR using quantitative autoradiography of rat brain sections (Table 1). Among these compounds, the fluoromethyl analogue 5 was the most active toward PBR. The *K*_i value (0.078 nM) of 5 was 2-fold higher than that of 2 and 10-fold higher than that of 1. This result suggested that substituting the OCH₃ group with OCH₂F group was favorable for augmenting its binding affinity for PBR. The fluoromethyl analogue 4 displayed a similar affinity to 2, suggesting that substitution with a fluoromethyl group did not obviously affect the affinity. This may be due to the molecular similarity and bioisosteric property of the OCH₂F and OCH₃ groups. The iodomethyl analogue 6 had 30 times lower affinity than 2, and the tosylloxethyl analogue 7 was 110 times lower than 2. These results suggested that the relative positions of this series were not favorable for the binding of PBR. However, although it was moderately active (*K*_i ≈ 18 nM) for PBR, the iodomethyl analogue

Table 1. In Vitro Binding Affinity (*K*_i) for PBR and CBR, and Octanol/Phosphate Buffer Distribution Coefficient (log *D*)



ligand	R	<i>K</i> _i (nM) ^a		log <i>D</i> ^d
		PBR ^b	CBR ^c	
4	FCH ₂	0.17 ± 0.02	>1000	3.70
5	FCH ₂ CH ₂	0.078 ± 0.01	>1000	3.81
6	ICH ₂	4.8 ± 0.68	>1000	4.02
7	TsOCH ₂ CH ₂	18.1 ± 1.30	>1000	4.65
2	CH ₃	0.16 ± 0.02	>1000	3.65
1		0.83 ± 0.24	>1000	2.78

^a Values represent the mean obtained from 9 concentrations of compound using at least 8 slices of rat brain (*n* = 3). ^b [¹¹C]2 was incubated in the presence of the compounds examined. ^c [¹¹C]Flumazenil was incubated in the presence of the compounds examined. ^d The log*D* values were determined in the phosphate buffer (pH = 7.4)/octanol system using the shaking flask method. All results were presented as mean values (*n* = 3) with a maximum range of ±5%.

6 may become a suitable candidate of a SPECT ligand for imaging PBR. Similarly, the binding affinities of 4–7 for CBR were measured using CBR-selective [¹¹C]flumazenil. As shown in Table 1, 4–7 did not display significant inhibitory effects (*K*_i > 1 μM) on [¹¹C]flumazenil binding in the rat brain. The negligible affinities of these analogues for CBR may be due to a difference from the typical benzodiazepine structure, which resulted in high selectivity for PBR.

Biodistribution on Mice. The radioactivity distribution of [¹⁸F]4 and [¹⁸F]5 was measured in mice over 120 min (the data are shown in Supporting Information). These results revealed that [¹⁸F]4 and [¹⁸F]5 showed a similar distribution pattern of radioactivity in the regions examined except bone. As can be seen, high radioactivity levels (>5% ID/g) were found in tissues such as the lung, heart, kidney, and adrenal gland, organs known to be rich in PBR. The distribution patterns of uptake were in agreement with that of PBR in the peripheral system, which was consistent with the distribution of [¹¹C]2 or [¹¹C]1 in the mouse. The highest accumulations of [¹⁸F]4 and [¹⁸F]5 were found in the lung and these high levels were probably due to the high mitochondrial contents containing PBR in the organ. On the other hand, high radioactivities (2.6–4.4% ID/g for [¹⁸F]4 and 2.2–4.9% ID/g for [¹⁸F]5) were also found in the brain, the target tissue in this experiment. The values were about 1.3–1.6-fold higher than that of [¹¹C]2, and 2–3-fold higher than that of [¹¹C]1, which was consistent with their higher partition coefficients as shown in Table 1. These findings suggested that [¹⁸F]4 and [¹⁸F]5 do pass across the brain–blood barrier (BBB), which is a prerequisite for a promising PET ligand.

In the bone of mouse, a significant difference in radioactivity was observed between [¹⁸F]4 and [¹⁸F]5. The accumulation of [¹⁸F]5 in the bone was initially low and showed no increase from 30 min (0.31% dose/g) to 120 min (0.09% dose/g), which demonstrated the ex-

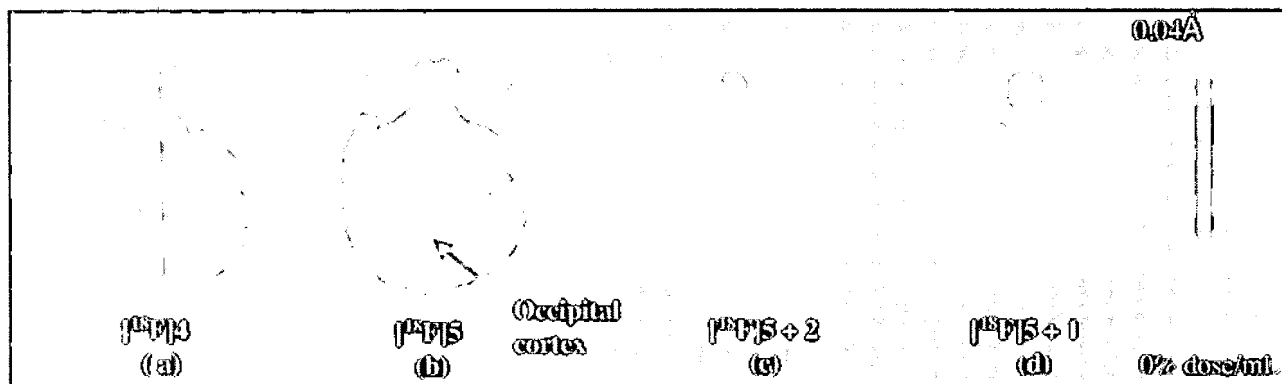


Figure 1. PET summation images of the monkey brain acquired between 30 and 180 min after [^{18}F]ligand injection. The images were obtained from the same subject. (a) [^{18}F]4; (b) [^{18}F]5; (c) [^{18}F]5 after treatment with **2** (1 mg/kg); (d) [^{18}F]5 after treatment with **1** (5 mg/kg). Significant radioactivity was observed in the bone after injection of [^{18}F]4 (a). High radioactivity was observed in the occipital cortex after injection of [^{18}F]5 (b), which was significantly inhibited by **2** (c) and **1** (d).

and stability of the fluoroethyl group against *in vivo* metabolism. In contrast, after injection of [^{18}F]4, the radioactivity began to accumulate in the bone and the brain at 120 min, which was 50 times higher than that of [^{18}F]5 in the bone. On consideration of the characteristic of [^{18}F]F $^-$ to readily accumulate in bone, [^{18}F]4 may be decomposed to [^{18}F]F $^-$ by *in vivo* defluorination. Our previous findings also demonstrated that [^{18}F]fluoroethyl compounds are relatively unstable in comparison with the corresponding [^{18}F]fluoroethyl and [^{11}C]methyl compounds.³³ Therefore, [^{18}F]4 may not be a useful PET ligand, even if it entered the brain and had some binding sites with PBR.

Monkey PET. The uptake of [^{18}F]4 and [^{18}F]5 in monkey brain was examined using PET. In the previous *in vitro* study for postmortem human brain, the highest density of PBR was observed in the occipital cortex.⁴⁶ The region of interest (ROI) in this PET experiment was placed on the occipital cortex. Figure 1 shows the PET summation images of the monkey brain acquired from 30 to 180 min after [^{18}F]4 (a) and [^{18}F]5 (b) injection (80–85 MBq/1.5 mL). The two images displayed a high accumulation of radioactivity in the brain, especially in the occipital cortex. However, in comparison with [^{18}F]5, a marked accumulation of [^{18}F]4 was observed in the bone. This image is visual evidence that [^{18}F]4 was decomposed to [^{18}F]F $^-$ *in vivo* as reflected by the high accumulation of [^{18}F]4 in the mouse bone. Therefore, no further evaluation of [^{18}F]4 was carried out using PET in monkey brain.

Figure 2 (circles) shows the time activity curve (TAC) of [^{18}F]5 in the occipital cortex of monkey brain after intravenous (iv) injection. At 2 min after injection, a high level of radioactivity was observed in the occipital cortex, which then remained almost same level during the measurement (180 min). The radioactivity of [^{18}F]5 was 1.5 times higher than that of [^{11}C]2,²⁹ and 6 times higher than that of [^{11}C]1¹⁹ at 30 min after injection. Pretreatment with nonradioactive **2** (1.0 mg/kg) gave a marked reduction of uptake for the image (Figure 1c) and TAC (Figure 2, triangles) of [^{18}F]5 as compared to the control experiment which was obtained under the same conditions. As shown in Figure 2, pretreatment with **1** enhanced the initial maximal uptake of [^{18}F]5. This increase may be derived from [^{18}F]5 dispossessed by mass nonradioactive **2** from lung abundant in PBR. Similar cases have been reported in the PET studies of

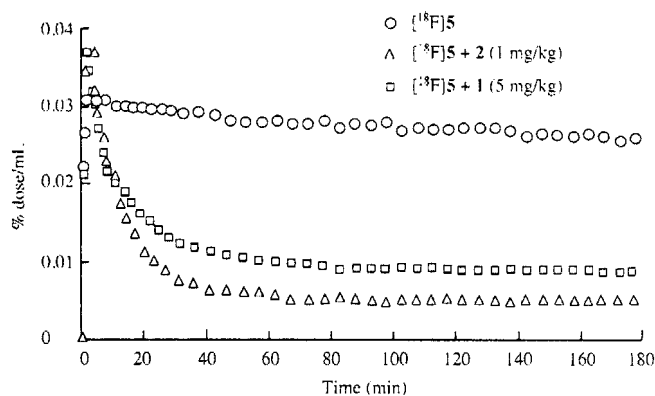


Figure 2. Time-activity curves (TACs) of [^{18}F]5 in the occipital cortex of monkey brain. The radioactivity of the control (circles) was strongly inhibited by pretreatment with **2** (triangles) and **1** (squares).

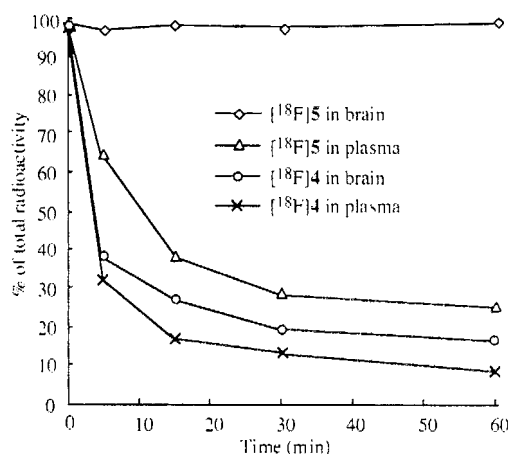
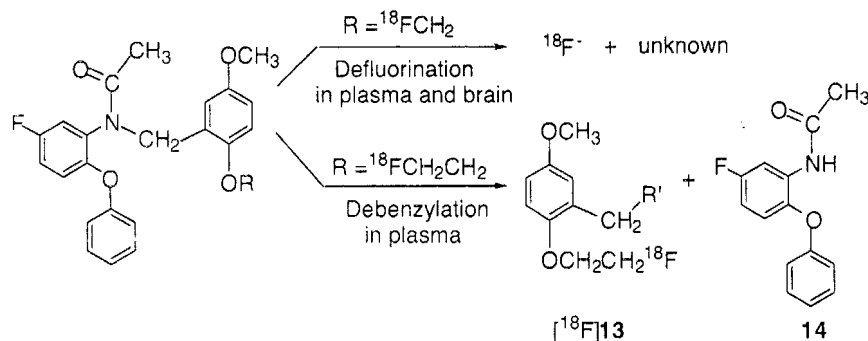


Figure 3. Percent conversion of [^{18}F]4 or [^{18}F]5 to metabolite in the mouse plasma and brain at several time points after iv injection of ligand (5–10 MBq) into the mice ($n = 3$). The unchanged tracers and metabolites were analyzed by HPLC for CH_3CN extracts from the plasma and brain homogenate prepared as described in the Experimental Section.

[^{11}C]1³⁶ and [^{11}C]2.²⁹ From 30 min after injection to the end (180 min) of the PET scan, pretreatment with **2** reduced the level of radioactivity to about 20% of the control. This result suggested high specific binding of [^{18}F]5 present in the occipital cortex. Pretreatment with the PBR-selective **1** (5 mg/kg) also produced a significant reduction of radioactivity in the occipital cortex, as shown in the image (Figure 1d) and TAC (Figure 2; squares). The radioactivity was reduced to about 30%

Scheme 4. Metabolism of [^{18}F]4 and [^{18}F]5 in the Plasma and Brain

control from 30 min after injection to the end of the PET scan. The reduction percentage on the uptake of [^{18}F]5 by **1** was slightly lower than that by **2**, which was probably due to the weaker affinity of **1** for PBR and its lower penetration²⁷ into the brain than that of **2**. These findings confirmed that [^{18}F]5 may have high specific binding for PBR in the monkey brain.

Metabolite Analysis. In the imaging brain study, the presence of metabolites in the plasma of the subject may preclude the evaluation of a PET ligand if the metabolites enter the brain and are retained/bound at the target sites. Therefore, metabolite analyses were performed for [^{18}F]5 in the plasma and brain of mouse and the plasma of monkey. For comparison, similar analyses were also performed for [^{18}F]4.

Figure 3 shows the percentages of unchanged [^{18}F]4 and [^{18}F]5 in the plasma and brain homogenate of mice determined by HPLC. After injection into the mouse, the amount of [^{18}F]4 corresponding to the unmetabolized [^{18}F]4 in the plasma rapidly decreased to 32% or 35% at 5 min and 22% or 23% at 30 min. A major radioactive metabolite with high polarity was observed in the HPLC. On strong ion exchange chromatography, the metabolite was assigned to [^{18}F]F. Since [^{18}F]F is hardly not brain permeable, [^{18}F]4 may be metabolized in the plasma and brain, respectively (Scheme 4).

After its injection into the mouse, the amount of [^{18}F]5 in the plasma continued to decrease during the entire experiment. The fraction corresponding to the unchanged [^{18}F]5 in the plasma was 64% at 5 min, 29% at 30 min, and 25% of the total radioactivity at 60 min after injection. No [^{18}F]F⁻ but another radioactive metabolite was observed in the HPLC. As estimated by its retention time ($t_R \approx 1.9$ min), the metabolite was much more polar than [^{18}F]5 ($t_R = 11.8$ min). On the other hand, only [^{18}F]5 was detected in the brain homogenate with no evidence (<5%) of any radioactive metabolites even at 60 min after injection.

The metabolite analysis was also performed using monkey plasma. After its injection into the monkey, [^{18}F]5 in plasma decreased to 77% at 5 min, 56% at 30 min, and 18% at 90 min. From then, the amount of unchanged [^{18}F]5 continued to decrease, while that of radioactive metabolite increased to 90% by the end of 180 min of the PET experiment. The radioactive metabolite in the monkey plasma was the same metabolite determined in the mouse plasma.

As reported previously, debenzylation of **2** and [^{11}C]2 was a main route of metabolism.^{25,26} Since the fluorinated analogue [^{18}F]5 had molecular similarity to and the same property of [^{11}C]2, it may be metabolized

with a similar profile to [^{11}C]2 (Scheme 1). A radioactive metabolite ([^{18}F]13) with a benzyl moiety and nonradioactive metabolite (*N*-(5-fluoro-2-phenoxyphenyl)acetamide (**14**) may be the putative metabolite of [^{18}F]5. The debenzylated compound **14** had no affinity for PBR and CBR ($\text{IC}_{50} > 10 \mu\text{M}$).²⁵ Therefore, the presence of the nonradioactive **14** could not interfere with the specific binding of [^{18}F]5 to PBR in the brain, even if it passed the BBB and entered the brain. Since no radioactive metabolites including [^{18}F]13 were detected in the mouse brain homogenates, the chemical structure of [^{18}F]13 was not further identified. These findings could reveal that although [^{18}F]5 was extensively metabolized in the plasma, all of the specific binding determined in the monkey brain may be due to this ligand itself and not influenced by its radioactive or nonradioactive metabolite.

Conclusion

To develop PET ligands for imaging PBR in the primate brain, four analogues **4**–**7** for PBR were synthesized and evaluated in this time. The two fluoroalkyl analogues **4** and **5** had higher or same affinities for PBR than the parent compound **2** and had no potency for CBR. The iodomethyl analogue **6** had a moderate affinity for PBR and may become a candidate for a SPECT ligand. The PET ligands [^{18}F]4 and [^{18}F]5 were respectively synthesized by the alkylation of the desmethyl precursor **3** with [^{18}F]8 and [^{18}F]9 in reproducible radiochemical yields. The distribution patterns of [^{18}F]4 and [^{18}F]5 in mice were consistent with the distribution of PBR. However, compared with [^{18}F]5, [^{18}F]4 exhibited a high accumulation of radioactivity in the bone. The PET image of [^{18}F]4 in the monkey brain also showed significant radioactivity in the bone, indicating that it was not a useful PET ligand because of its remarkable in vivo defluorination. The uptake of [^{18}F]5 in the occipital cortex of the monkey brain was 1.5 times higher than that of [^{11}C]2, and 6 times (maximum value) higher than that of [^{11}C]2. Pretreatment with **1** or **2** significantly reduced the radioactivity levels of [^{18}F]5 to 20–30% of the control in the brain, suggesting a high specific binding of [^{18}F]5 to PBR in the monkey brain. No radioactive metabolite of [^{18}F]5 was detected in the brain although it was metabolized in the plasma by debenzylation. This study has succeeded in developing a potential PET ligand [^{18}F]5 which has high specific binding for PBR in the monkey brain.

Experimental Section

^1H NMR spectra were recorded on a JNM-GX-270 spectrometer (JEOL, Tokyo) with tetramethylsilane as an internal

standard. All chemical shifts (δ) were reported in parts per million (ppm) downfield from the standard. FAB/MS was obtained on a JEOL NMS-SX102 spectrometer (JEOL, Tokyo). Column chromatography was performed using Merck spherigel gel 60 F₂₅₄ (70–230 mesh). 18 Fluorine (^{18}F) was produced by the ^{18}O p, n ^{18}F nuclear reaction using a CYPRIS 18 cyclotron (Sumitomo Heavy Industry, Tokyo). If not otherwise stated, radioactivity was determined with an ICC countermeeter (Aloka, Tokyo). HPLC was performed using a JASCO HPLC system (JASCO, Tokyo); effluent radioactivity was monitored using a NaI (TI) scintillation detector system. Samples **2** and **3** were kindly provided by Dr. A. Nakazoto (Mitsubishi Chemical). All chemical reagents with the highest grade commercially available were purchased from Wako Pure Chem. (Milwaukee, WI) and Wako Pure Chem. Ind. (Saka). The animal experiments were carried out according to the guidelines of the committee for the care and use of animals of the National Institute of Radiological Sciences (NIRS).

Chemistry. *N*-(5-Fluoro-2-phenoxyphenyl)-*N*-(2-fluoromethyl-5-methoxybenzyl)acetamide (**4**). A mixture of **3** (19 mg, 0.1 mmol), FCH_2I (**4**, about 20 mg in 1 mL of CH_3CN), and NaH (4 mg, 0.1 mmol) in DMF (2 mL) was stirred at 0 °C for 5 h. The reaction mixture was quenched with AcOEt and washed with water and a saturated NaCl solution. After the organic layer was dried over Na_2SO_4 , the solvent was removed to give a residue. Column chromatography of the residue on silica gel with CHCl_3 /hexane (1/9) gave **4** (16 mg, 79%) as a white powder: mp: 71–72 °C; ^1H NMR (300 MHz, CDCl_3) δ : 7.83 (d, 2H, m), 6.71–7.13 (8H, m), 6.28–6.44 (1H, m), 5.30 (dd, $J = 4.9$ Hz), 4.80 (2H, dd, $J = 2, 12$ Hz), 3.71 (3H, s), 3.41 (3H, s). FAB/MS (m/z): 414.2 ($\text{M}^+ - 1$); Anal. ($\text{C}_{21}\text{H}_{21}\text{F}_2\text{NO}_2$) $\text{C}, \text{H}, \text{N}$.

N-(5-Fluoro-2-phenoxyphenyl)-*N*-(2-fluoroethyl-5-methoxybenzyl)acetamide (**5**). A mixture of **3** (38 mg, 0.1 mmol), $\text{FCH}_2\text{CH}_2\text{I}$ (**5**, 1.0 mmol) and NaH (8 mg, 0.2 mmol) in DMF (5 mL) was heated at 80 °C for 8 h. The reaction mixture was quenched with AcOEt, and washed with water and a saturated NaCl solution. After the organic layer was dried over Na_2SO_4 , the solvent was removed to give a residue. Column chromatography of the residue on silica gel with CHCl_3 /hexane (1/13) gave **5** (27 mg, 65%) as a white powder: mp: 74–75 °C; ^1H NMR (300 MHz, CDCl_3) δ : 7.26–7.59 (2H, m), 6.22–7.19 (9H, m), 4.88 (2H, dt, $J = 4, 41$ Hz), 4.65 (2H, dt, $J = 2, 12$ Hz), 4.12 (2H, dt, $J = 4, 27$ Hz), 3.80 (3H, s), 3.41 (3H, s). FAB/MS (m/z): 428.2 ($\text{M}^+ - 1$); Anal. ($\text{C}_{22}\text{H}_{23}\text{F}_2\text{NO}_2$) $\text{C}, \text{H}, \text{N}$.

N-(5-Fluoro-2-phenoxyphenyl)-*N*-(2-iodomethyl-5-methoxybenzyl)acetamide (**6**). A mixture of **3** (19 mg, 0.05 mmol), FCH_2I (**6**, 0.15 mmol) in DMF (5 mL) was stirred at 0 °C for 5 h. The reaction mixture was quenched with AcOEt and washed with water and a saturated NaCl solution. After the organic layer was dried over Na_2SO_4 , the solvent was removed to give a residue. Column chromatography of the residue on silica gel with CHCl_3 /hexane (1/15) gave **6** (8.9 mg, 46%) as a colorless oil: ^1H NMR (300 MHz, CDCl_3) δ : 7.18–7.60 (2H, m), 7.33 (9H, m), 4.81 (dd, $J = 2, 12$ Hz), 4.70 (s), 4.78 (1H, s), 2.20 (3H, s). FAB/MS (m/z): 524.1 ($\text{M}^+ - 1$); Anal. ($\text{C}_{21}\text{H}_{21}\text{F}_2\text{INO}_2$) $\text{C}, \text{H}, \text{N}$.

N-(5-Fluoro-2-phenoxyphenyl)-*N*-(5-methoxy-2-tosyloxyethylbenzyl)acetamide (**7**). A mixture of **3** (19 mg, 0.05 mmol), $\text{TsOCH}_2\text{CH}_2\text{OTf}$ (**7**, 37 mg, 0.1 mmol) and NaH (8 mg, 0.2 mmol) in DMF (5 mL) was stirred at 0 °C for 5 h. The reaction mixture was quenched with AcOEt and washed with water and a saturated NaCl solution. After the organic layer was dried over Na_2SO_4 , the solvent was removed to give a residue. Column chromatography of the residue on silica gel with CHCl_3 /hexane (1/20) gave **7** (11 mg, 46%) as a colorless oil: ^1H NMR (300 MHz, CDCl_3) δ : 6.52–7.48 (15H, m), 4.62 (dd, $J = 2, 12$ Hz), 4.12–4.28 (2H, m), 4.12–3.92 (2H, m), 3.80 (3H, s), 3.41 (3H, s). FAB/MS (m/z): 580.3 ($\text{M}^+ - 1$); Anal. ($\text{C}_{27}\text{H}_{29}\text{FNO}_6$) $\text{C}, \text{H}, \text{N}$.

Radiosynthesis. [^{18}F]Fluoride ($^{18}\text{F}^-$). [^{18}F]F $^-$ was produced by the ^{18}O p, n ^{18}F reaction on 10–20 at % H_2^{18}O

using 18 MeV protons (14.2 MeV on target) from the cyclotron and separated from [^{18}O]H $_2\text{O}$ using Dowex 1 X8 anion-exchange resin in an irradiating room. The [^{18}F]F $^-$ was eluted from the resin with aqueous K_2CO_3 (3.3 mg/0.3 mL) into a vial containing CH_3CN (1.5 mL)/4,7,13,16,21,24 hexaoxa 1,10-diazabicyclo[8.8.8]hexacosane (Kryptofix 222, 25 mg) and transferred into a reaction vessel in a hot cell.

N-(5-Fluoro-2-phenoxyphenyl)-*N*-(2-[^{18}F]fluoromethyl-5-methoxybenzyl)acetamide ([^{18}F]4). [^{18}F]F $^-$ from the irradiating room was transported to a Pyrex glass vessel (5 mL) containing 100 μL of *o*-dichlorobenzene (*o*-DCB), and the [^{18}F]F $^-$ was dried to remove H_2O and CH_3CN at 120 °C for 15 min. After CH_2I_2 (**10**, 100 μL) was added to the radioactive mixture by a helium flow (50 mL/min) at 120 °C, [^{18}F]FCH $_2\text{I}$ ([^{18}F]8) resulted in this vessel was distilled at once under helium for 3 min and bubbled into another Pyrex glass vessel containing the demethyl precursor (**3**, 1.5 mg) and NaH (10 μL , 1.5 g/20 mL DMF) in anhydrous DMF (300 μL) at –15 °C. After maximum radioactivity was bubbled into the solution, the reaction was terminated by adding $\text{CH}_3\text{CN}/\text{H}_2\text{O}$ (6/4, 500 μL). The radioactive mixture was applied to a semipreparative HPLC column. HPLC semipreparative purification was completed on YMC Jsphere ODS-H80 column (10 mm ID $\times$ 250 mm) using a mobile phase of $\text{CH}_3\text{CN}/\text{H}_2\text{O}$ (60/40) at a flow rate of 6.0 mL/min. The retention time (t_R) for [^{18}F]4 was 11.2 min, whereas that for **3** was 6.7 min. The radioactive fraction corresponding to [^{18}F]4 was collected in a sterile flask containing polysorbate (80) (75 μL) and ethanol (150 μL), evaporated to dryness under vacuum, redissolved in 7 mL of sterile normal saline, and passed through a 0.22 μm Millipore filter to obtain the final product. At the end of synthesis (EOS), 180–300 MBq of [^{18}F]4 was obtained as an iv injectable solution at a beam current of 10–15 μA and 20–25 min proton bombardment.

N-(5-Fluoro-2-phenoxyphenyl)-*N*-(2-[^{18}F]fluoroethyl-5-methoxybenzyl)acetamide ([^{18}F]5). After the [^{18}F]F $^-$ was dried, $\text{BrCH}_2\text{CH}_2\text{OTf}$ (**12**, 8 μL) in *o*-dichlorobenzene (400 μL) was added to the radioactive mixture. The [^{18}F]FCH $_2\text{CH}_2\text{Br}$ ([^{18}F]9) in this vessel was distilled under a helium flow (90–100 mL/min) at 130 °C for 5 min and bubbled into another vessel containing **3** (1.5 mg) and NaH (10 μL , 1.5 g/20 mL DMF) in anhydrous DMF (300 μL) at –15 °C, and the reaction mixture was heated and kept at 120 °C for 10 min. HPLC semipreparative purification was performed on a YMC Jsphere ODS-H80 column (10 mm ID $\times$ 250 mm) using a mobile phase of $\text{CH}_3\text{CN}/\text{H}_2\text{O}$ (55/45) at a flow rate of 6.0 mL/min. The t_R for [^{18}F]5 was 14.2 min. At EOS, 570–780 MBq of [^{18}F]5 was obtained as an iv injectable solution at a beam current of 10–15 μA and 20–25 min proton bombardment.

Radiochemical Purity and Specific Activity Determinations. Radiochemical purity was assayed by analytical HPLC (column: CAPCELL PAK C $_{18}$, 4.6 mm ID $\times$ 250 mm, UV at 254 nm, mobile phase: $\text{CH}_3\text{CN}/\text{H}_2\text{O}$ = 6/4). The t_R for [^{18}F]4 and [^{18}F]5 was 6.1 and 6.3 min at a flow rate of 2.0 mL/min. The specific activity of [^{18}F]4 and [^{18}F]5 was determined by comparison of the assayed radioactivity to the mass associated with the carrier UV peak at 254 nm.

In Vitro Binding Assays. Male Sprague–Dawley rats ($n = 4$) weighing 220–250 g were killed by decapitation under ether anesthesia, and their brains were quickly removed and frozen on powdered dry ice. Brain sagittal sections (20 μm) were cut on a cryostat microtome (HM560, Carl Zeiss Co., Germany) and thaw-mounted on glass slides (Matsunami Glass Ind., Tokyo), which were then dried at room temperature and stored at –18 °C until use for experiments. The brain sections were preincubated at 25 °C for 20 min in 50 mM Tris-HCl (pH 7.4) buffer. After the preincubation, these sections were incubated at 37 °C for 30 min in the assay buffer containing [^{14}C]2 (1 nM, specific activity: 110 GBq/ μmol) or [^{14}C]flumazenil (1 nM, specific activity: 220 GBq/ μmol). To determine the IC_{50} values of **1**, **2**, **4**, **7** for the [^{14}C]2 (for PBR) or [^{14}C]flumazenil (for CBR) binding, the brain sections were incubated with [^{14}C]2 or [^{14}C]flumazenil in the presence of increasing concentrations of the corresponding **1**, **2**, **4**–**7** (0.1–1000 nM). Nonradioactive **2** or flumazenil (10 μM) of was used

to determine the nonspecific binding for PBR or CBR. After autoradiation the sections were washed three times for 2 min each time with the cold assay buffer, dipped into cold distilled water, and dried with a stream of warm air (about 50 °C). These sections were then placed in contact with imaging plates (BAS-SR-224; Fuji Photo Film Co. Ltd., Tokyo, Japan) for 60 min. The distribution of their radioactivity with a BAS-EX BAS-3000 bioimaging analyzer (Fuji). The region of interest (ROI) on the sections was placed on the cerebellum. The data corresponding to the radioactivity on the cerebellum were used to correct the absence of the displacement. 1, 2, 4–7 were used to assess the specific binding for PBR or CBR was defined as the binding minus binding in the presence of 2 or flumazenil (100 nM). The PBR data corresponding to the specific binding of each compound concentration was calculated as a percent of the control specific binding, and which were converted to pIC₅₀ values to determine the IC₅₀ of each compound. The IC₅₀ value was further converted to K_i according to the Cheng–Prusoff equation.⁴⁶

Biodistribution on Mice. A saline solution of ¹⁸F-ligand (specific activity of 8 MBq/200 μ L) was injected into ddv mice (30–40 g) in the back, raised through the tail vein. Four mice for each compound were sacrificed by cervical dislocation at 5, 15, 30, 60, and 180 min after injection. Whole brain, liver, lung, heart, kidney, adrenal, bone, and blood samples were quickly removed and weighed. The radioactivity present in the various tissues was measured in a Packard autogamma scintillation counter and expressed as a percentage of the injected dose per gram of wet tissue ($\pm$ SD g). All radioactivity measurements were corrected for decay.

PET on Monkeys. The PET scan was performed using a small-animal PET camera (Hamamatsu Photonics, Hamamatsu, Japan) designed for laboratory animals, which acquires 16 transaxial slices 3.6 mm (center-to-center) apart and a 33.1-cm field of view. A male rhesus monkey (*Macaca mulatta*) weighing about 5 kg was repeatedly anesthetized with ketamine (ketalar, 10 mg/kg/h, im) every hour for 24 h. After anesthesia, transmission scans for attenuation correction were performed for 1 h using a 74 MBq ⁶⁷Ge Tl Co source. A dynamic emission scan in the 3D acquisition mode was performed for 20 min (2 min $\times$ 5 scans, 4 min $\times$ 10 scans, 4 min $\times$ 5 scans). All emission scan images were reconstructed with a 3-mm slice filter of 1 mm, and circular regions of interest (ROIs) with a 5-mm diameter were placed over the occipital cortex using an image analysis software.^{47,48} A solution of ¹⁸F-ligand 4 or 5 (80–85 MBq) was injected iv into the monkey. After sequential tomographic scanning was performed on a transverse section of the brain for 180 min. In the following experiments, nonradioactive 2 (1 mg/kg) or 1 (5 mg/kg) was injected at 2 min before the [¹⁸F]5 injection. The time-activity curves (TACs) in the occipital cortex were obtained for each scan of the brain.

Metabolite Assay for Mouse Plasma and Brain Tissue. After iv injection of [¹⁸F]4 or [¹⁸F]5 (5–10 MBq/100 μ L) into ddv mice (n = 3), these mice were sacrificed by cervical dislocation at 5, 15, 30, or 60 min. Blood (0.7–1.0 mL) and whole brain samples were removed quickly. The blood sample was centrifuged at 15 000 rpm for 1 min at 4 °C to separate plasma (250 μ L), which was collected in a test tube containing 100 μ L of CH₃CN (10:1), and a solution of the authentic unlabeled 4 (100 ng) in 50 μ L of CH₃CN (10:1). After the tube was vortexed for 15 s and centrifuged at 15 000 rpm for 2 min for deproteinization, the supernatant was collected. The extraction efficiency of radioactivity into the CH₃CN supernatant ranged from 73% to 94% of the total radioactivity in the plasma. On the other hand, the whole brain including the cerebellum, etc. was dissected from the mouse brain and homogenized in 1 mL of ice-cold CH₃CN/H₂O (1:1, 1.0 mL/mL). The homogenate was centrifuged at 15 000 rpm for 1 min at 4 °C, and supernatant was collected. The recovery of radioactivity into the supernatant was 68–87% based on the total radioactivity in the brain homogenate.

An aliquot of the supernatant (100–300 μ L) obtained from the plasma or brain homogenate was injected into the HPLC

system for radioactivity and analyzed under the same conditions described above except that the mobile phase was CH₃CN/H₂O with a ratio of 1/1. The percent ratio of [¹⁸F]ligand (t_R = 10.6 min for [¹⁸F]4 and 11.8 min for [¹⁸F]5) to total radioactivity (corrected for decay) on the HPLC chromatogram was calculated as % = (peak area for [¹⁸F]ligand/total radioactive peak area) $\times$ 100.

Metabolite Analysis for Monkey Plasma. After iv injection of [¹⁸F]5 (80 MBq) into the monkey, arterial blood samples (1 mL) were collected at 2, 10, 30, 60, 90, 120, and 180 min. All samples were centrifuged at 15 000 rpm for 1 min at 4 °C to separate plasma (250 μ L), which was collected in a test tube containing CH₃CN (0.5 mL). The tube was vortexed for 15 s and centrifuged at 15 000 rpm for 1 min for deproteinization. The extraction efficiency of radioactivity into the CH₃CN ranged from 73% to 94% of the total radioactivity in the plasma. The radioactive fractions in these samples were determined using HPLC.

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Supporting Information Available: Radioactivity distribution of [¹⁸F]4 and [¹⁸F]5. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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