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ORIGINAL RESEARCH





Synthesis and evaluation in rats of homologous series of [18 F]-labeled dopamine D_{2/3} receptor agonists based on the 2-aminomethylchroman scaffold as potential PET tracers

Vladimir Shalgunov¹, Jan-Peter van Wieringen², Henk M. Janssen³, P. Michel Fransen³, Rudi A.J.O. Dierckx¹, Martin C. Michel⁴, Jan Booij² and Philip H. Elsinga^{1*}

Abstract

Background: Agonist positron emission tomography (PET) tracers for dopamine $D_{2/3}$ receptors ($D_{2/3}$ Rs) offer greater sensitivity to changes in endogenous dopamine levels than $D_{2/3}$ R antagonist tracers. $D_{2/3}$ R agonist tracers currently available for clinical research are labeled with the short-lived isotope carbon-11, which limits their use. We aimed to develop high-affinity D_2 R agonists amenable for labeling with the longer-living fluorine-18. Here, we report the evaluation as potential PET tracers of two homologous series of [¹⁸F]fluorinated tracers based on the 2-aminomethylchroman-7-ol (AMC) scaffold: (R)-2-((4-(2-fluoroalkoxy)benzylamino)methyl)chroman-7-ols (AMC13 homologues) and (R)-2-((2-(4-(4-(fluoroalkoxy)phenyl)piperazin-1-yl)ethylamino)methyl)chroman-7-ols (AMC15 homologues). We varied the length of the ¹⁸F-fluoroalkyl chain in these structures to balance brain penetration and non-specific binding of the radioligands by adjusting their lipophilicity.

Methods: The tracers were evaluated in brain slices of Sprague-Dawley rats by in vitro autoradiography and in living rats by microPET imaging and ex vivo autoradiography. PET data were analyzed with one- and two-tissue compartmental models (1TCM/2TCM), simplified reference tissue model (SRTM), and Logan graphical analysis. Specificity of binding was tested by blocking D_{2/3}R with raclopride.

Results: Homologues with a shorter fluoroalkyl chain consistently showed greater $D_{2/3}R$ -specific-to-total binding ratios in the striatum than those with longer chains. The fluoroethoxy homologue of AMC13 ([¹⁸F]FEt-AMC13) demonstrated the highest degree of $D_{2/3}R$ -specific binding among the evaluated tracers: mean striatum-to-cerebellum uptake ratio reached 4.4 in vitro and 2.1/2.8 in vivo/ex vivo (PET/autoradiography). Striatal binding potential (BP_{ND}) relative to cerebellum was 0.51–0.63 depending on the estimation method. Radiometabolites of [¹⁸F]FEt-AMC13 did not enter the brain. In vitro, application of 10 µmol/L raclopride reduced $D_{2/3}R$ -specific binding of [¹⁸F]FEt-AMC13 in the striatum by 81 %. In vivo, pre-treatment with 1 mg/kg (2.9 µmol/kg) raclopride led to 17–39 % decrease in $D_{2/3}R$ -specific binding in the striatum.

Conclusions: Varying the length of the [18 F]fluoroalkyl chain helped improve the characteristics of the original candidate tracers. Further modifications of the current lead [18 F]FEt-AMC13 can provide an agonist radiopharmaceutical suitable for D_{2/3}R imaging by PET.

Keywords: Agonist tracer; Dopamine receptor; PET; Fluorine-18

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Background

Dysregulation of dopamine signaling through dopamine D_2 and D_3 receptors ($D_{2/3}$ Rs) is implicated in many neuropsychiatric disorders [1–4], making imaging of $D_{2/3}$ Rs by positron emission tomography (PET) highly relevant.

Majority of PET tracers developed for $D_{2/3}R$ imaging are antagonists [5], but in the last two decades, agonists have attracted attention as potential PET ligands for G protein-coupled neurotransmitter receptors (GPCRs). In in vitro competition studies, agonists consistently show higher affinity for receptor molecules bound to G proteins ("high-affinity state") than for free receptors ("low-affinity state") [6–8]. Agonists are therefore expected to selectively recognize the "high-affinity" subset of receptor population in the imaging experiments, while antagonists bind to all receptors disregarding "affinity states" [9].

Though the existence of a separate "high-affinity state" subpopulation of $D_{2/3}R$ in the brain is still not conclusively demonstrated in vivo [10], radiolabeled $D_{2/3}R$ agonists indeed turned out to be more sensitive than the antagonist [¹¹C]raclopride to amphetamine-induced release of dopamine (itself also an agonist) in rodents, cats, non-human primates, and humans [11–17]. Alteration of the relative abundance of "high-affinity state" $D_{2/3}Rs$ ($D_{2/3}R$ -high), detected by in vitro methods, is implied in dopamine supersensitivity, a state relevant to psychosis, Parkinsonism, and drug addiction [18]. Changes in this in vitro-based parameter are likely to be translated into effects observable in vivo, so agonist tracers might provide new insights for the research into, and (early) diagnosis of, these common neuropsychiatric disorders.

Numerous scaffolds have been investigated in the development of D_{2/3}R agonist PET tracers [19], but representatives of only two classes, namely the apomorphines [¹¹C](-)NPA and [¹¹C]MNPA and the naphtoxazine [¹¹C](+)PHNO, have hitherto been used in human PET studies. These tracers can only be used in hospitals with an on-site cyclotron due to the short half-life of carbon-11. Fluorine-18-labeled tracers (half-life of 109.8 min) have greater potential of widespread application in the clinic and are also more suitable for long-duration experiments in the research set-up. [¹⁸F]5-OH-FPPAT, an aminotetraline tracer, showed promising results in rats and monkeys [20], but no evaluation in humans has been published so far. A [¹⁸F]fluorinated analog of [¹¹C](+)PHNO failed to demonstrate $D_{2/3}$ R-specific binding in rats [21], while the recently described [¹⁸F]MCL-524, structurally related to [¹¹C]MNPA, showed very promising results in nonhuman primates [22].

We aimed to increase the availability of fluorine-18labeled $D_{2/3}R$ agonist tracers for PET. For this purpose, we selected the 2-aminomethylchroman-7-ol (AMC) scaffold. Structure-activity relationships of this class of high-affinity D_2R -agonists were previously investigated by Mewshaw and co-workers [23].

Recently, we have described the preparation of two [¹⁸F]labeled AMCs (Fig. 1): (R)-2-[(4-(4-[¹⁸F]fluorobutoxy)benzylamino)methyl]chroman-7-ol ([18F]FBu-AMC13) and (*R*) $1-(4-(2-[^{18}F]fluoroethoxy)phenyl)-4-(4-(7-hydroxychro$ man-2-yl)-3-azabutyl)-piperazine ([¹⁸F]FEt-AMC15). Both compounds, when tested in their non-radioactive $[^{19}F]$ form, showed nearly full agonism at the long isoform of human D₂R and low-nanomolar affinities towards the high-affinity state of these receptors: on average 5.6 and 6.7 nM for FBu-AMC13 and FEt-AMC15, respectively. The affinities of these two compounds towards human D₂R and D₃R were comparable to each other, with no detectable preference for any of the two receptor subtypes [24]. Both [¹⁸F]FBu-AMC13 and [¹⁸F]FEt-AMC15 also demonstrated specific binding to the D_{2/3}R-high in rat striata. We wanted to further evaluate these radioligands in vivo and optimize their brain penetration and nonspecific binding by adjusting their lipophilicity. We reasoned that the latter goal could be achieved by varying the length of the [¹⁸F]fluoroalkyl chain in the structures of the tracers, because these chains are situated outside the main pharmacophore of the AMCs [24], so shortening or elongating them will likely have no influence on the affinity of the resulting ligands towards $D_{2/3}R$.

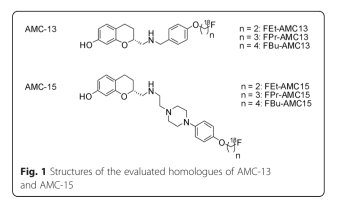
This study reports on the synthesis of [¹⁸F]-labeled AMC13 and AMC15 homologues with varied alkyl chain lengths, and on the in vitro and in vivo evaluation of these compounds.

Methods

Chemistry

Reagents, chemicals, materials, and solvents were obtained from commercial sources and were used as received: Biosolve, Merck for solvents, Cambridge Isotope Laboratories for deuterated solvents, and Aldrich, Acros, ABCR, Merck, and Fluka for chemicals, materials, and reagents. All solvents were of analytical grade quality.

¹H-NMR and ¹³C-NMR spectra were recorded on Varian MR (300 or 400 MHz for ¹H-NMR, 100 MHz for



 13 C-NMR) spectrometers at an ambient temperature. Chemical shifts are reported in ppm, applying deuterated chloroform (CDCl₃) or other deuterated solvents as internal reference. Abbreviations used for splitting patterns are s = singlet, t = triplet, q = quartet, m = multiplet, and dd = double doublet.

LC-ESI-MS/MS analyses were performed on a Shimadzu LC-20AD series LC coupled to an API3000 mass spectrometer (Applied Biosystems/MDS SCIEX, Toronto, Canada), a triple quadrupole mass spectrometer supplied with an atmospheric pressure ionization source, and an ion-spray interface. Analyses were executed at 298 K using an Alltech Alltima C18 5 μ 150 × 2.1 mm column using an injection volume of 10 μ L, a flow rate of 0.4 mL/min, and a gradient of MeCN/propan-2-ol in H₂O (from 5 to 95 % organic component, where both MeCN/propan-2-ol and H₂O contain 0.1 % formic acid). MS/MS analysis was performed in the multiple reaction monitoring mode.

Analytical thin layer chromatography (TLC) was performed on Kieselgel F-254 precoated silica plates. Normalphase column chromatography was carried out on regular silica gel 60 (0.040–0.063, Merck).

1,2-Ethanediol ditosylate, 1,3-propanediol ditosylate, 1,4butanediol ditosylate, 2-fluoroethyl tosylate, 3-fluoropropyl tosylate, and 4-fluorobutyl tosylate were prepared according to procedures described in the literature [24, 25]. Analytical data (¹H-NMR, ¹³C-NMR) were according to expectations.

The preparation and characterization of phenol precursors for ¹⁸F-fluoroalkylation, (R) N-[7-(methoxymethoxy)-chroman-2-yl]methyl 4-hydroxybenzyl amine, and (R) 1-(4-hydroxyphenyl)-4-(4-(7-hydroxychroman-2-yl)-3-azabutyl)-piperazine were described earlier by van Wieringen et al. [24].

[¹⁹F]-references for FBu-AMC13 and FEt-AMC15 were prepared and characterized as described previously by van Wieringen et al. [24]. Of the new homologous ligands, [¹⁹F]-reference was only prepared for FEt-AMC13, the homologue that showed the most promising results in vivo. Given the use of the very similar radiolabeling methods for all ligands, and the close structural relatedness of the new homologues to the originally developed FBu-AMC13 and FEt-AMC15, we decided not to prepare [¹⁹F]-references for other homologues.

(*R*)-2-[(4-(4-Fluoroethoxy)benzylamino)methyl]chroman-7-ol ([¹⁹F]-reference of FEt-AMC13)

A solution of (*R*) N-[7-(methoxymethoxy)chroman-2yl]methyl 4-hydroxybenzyl amine (28.4 mg, 0.11 mmol) in 1.5 ml dry DMF and 5 mg NaH were stirred in an oil bath at 80 °C. To this mixture 2-fluoroethyl tosylate (22.8 mg, 0.11 mmol) dissolved in 1.2 ml dry DMF was added in three portions of 0.4 ml over the period of 1 hour. The mixture was stirred at 80 °C for 24 h. After that, the reaction mixture was cooled down and N₂ was bubbled through the solution to create an inert atmosphere. Then, 2 mL of 4 M HCl solution in dioxane was added to the mixture, and it was stirred for 5 h at room temperature in the atmosphere of N₂. N₂ was further bubbled through the solution to remove volatile reaction products from the mixture. Thereafter, the reaction mixture was diluted with 50 mL water and the pH was brought to 8–8.5 by addition of a Na₂CO₃-solution. The mixture was extracted by CH₂Cl₂ (3 × 10 mL), organic extract was washed with water (3 × 10 mL), dried with Na₂SO₄, and evaporated in vacuo. The residue was purified using silica column chromatography (ethyl acetate/ methanol/triethylamine 100/5/1), giving 8.5 mg of the product (24% yield), as a slightly yellowish solid.

¹H NMR (CD₃OD/CDCl₃): δ = 7.37 (d, 2H), 6.98 (d, 2H), 6.87 (d, 1H), 6.39 (dd, 1H), 6.21 (d, 1H), 4.83 (t, 1H), 4.71 (t, 1H), 4.29 (t, 1H), 4.22 (t, 1H), 4.12 (m, 1H), 4.01 (bs 2H), 3.01 (m, 2H), 2.9–2.5 (multiple signals, 2H), 2.05–1.65 (multiple signals, 2H). ¹³C NMR (CD₃OD/CDCl₃): δ = 162.7, 160.0, 158.2, 134.5, 133.9, 133.0, 118.8, 116.6, 112.5, 107.0, 86.6, 84.8, 76.6, 71.3, 71.1, 55.6, 55.2, 35.7, 33.4, 29.5, 27.3.

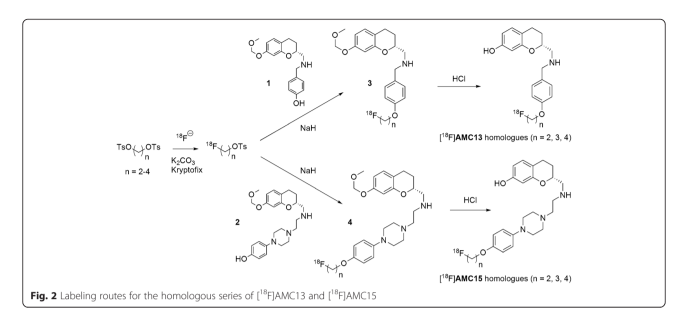
Radiochemistry

[¹⁸F]-fluoride was produced in the Scanditronix MC-17F cyclotron (GE Healthcare) and delivered from the target into the hotcell with helium overpressure. Fig. 2 gives an overview of the applied radiochemistry.

Synthetic procedures were performed manually. Radiochemical conversions and purities were determined by radio-HPLC using beta-sensitive detectors and dosecalibrators (Veenstra Instruments) and by radio-TLC using Cyclone phosphor storage screens (multisensitive, Packard) and OptiQuant software.

Preparation of [¹⁸F]fluoroalkyl tosylates

[¹⁸F]-Fluoride in enriched H₂¹⁸O water was adsorbed on an activated anion-exchange column (QMA) and later eluted into the reaction vial with 1-1.5 mL of 1.5 mg/mL K_2CO_3 , 5 mg/mL Kryptofix solution in 9:1 ν/ν mixture of acetonitrile and water, respectively. After the elution of QMA, acetonitrile and water were evaporated at 130 °C under nitrogen flow. For further water removal, 1 mL anhydrous acetonitrile was added into the reaction vial and evaporated. Then, 5 mg of alkyl ditosylate precursor (ethyl, propyl, or butyl) in 1 mL anhydrous acetonitrile was added into the vial, the vial was sealed, and reaction was carried out for 4 min at 130 °C followed by cooling 5 min. To the cooled reaction mixture, 1 mL of water was added, and the resulting mixture was loaded into the HPLC loop. [¹⁸F]fluoroalkyl tosylates were purified by reversephase HPLC using Alltima C18 5 μ 250 × 10 mm column eluted with acetonitrile/water at 5 mL/min. Acetonitrile content in the mobile phase was 65 % for [¹⁸F]fluorobutyl



tosylate and 50 % for [¹⁸F]fluoropropyl and [¹⁸F]fluoroethyl tosylate. Under these conditions, retention times of all thee [¹⁸F]fluoroalkyl tosylates were in the range of 6–8 min. Collected radioactive fraction was diluted two- to three-fold with water and 4-[¹⁸F]fluoroalkyl tosylate was adsorbed on tC18 light Sep-Pak cartridge. The cartridge was rinsed with 5 mL pure water and dried for 5 min in strong nitrogen flow.

Preparation of [¹⁸F]FEt-AMC13, [¹⁸F]FPr-AMC13, and [¹⁸F]FBu-AMC13

(R) N-[7-(Methoxymethoxy)chroman-2-yl]methyl 4-hydroxybenzyl amine (1, 0.6 mg) in 30 µL dry DMF and 2-3 mg NaH were put into a V-vial. Thereafter, the corresponding [18F]fluoroalkyltosylate was eluted into the vial from the tC18 light Sep-Pak cartridge with 0.6-0.7 mL dry DMF. The reaction vial was sealed and the alkylation reaction to acquire compounds 3 were conducted for 10 min at 120 °C. Then, 1 mL 0.4 N HCl was added into the vial with a syringe and deprotection of the MOM-group followed that also lasted 10 min at 120 °C. The product was purified by reverse-phase HPLC using Alltima C18 5µ 250×10 mm column eluted with acetonitrile/10 mM H₃PO₄ at 5 mL/min. Acetonitrile content in the mobile phase was 30 % for [18F]FEt-AMC13 and 35 % for [¹⁸F]FPr-AMC13 and [¹⁸F]FBu-AMC13. Retention times were 10.6 min for [18F]FEt-AMC13, 9.2 min for [18F]FPr-AMC13, and 10.8 min for [¹⁸F]FBu-AMC13. Collected HPLC fraction was diluted with 2-3 volumes of water and the radioactivity was adsorbed on Oasis HLB SPE cartridge. The cartridge was rinsed with 5 mL pure water and radioactivity was then eluted with 1-2 mL absolute ethanol. Typical recovery was 20-40 % for pure ethanol and 80–90 % for 100/1 ν/ν mixture of ethanol with acetic or ortho-phosphoric acid; in synthesis runs for in vivo experiments, only pure ethanol was used. After partial or complete evaporation of ethanol in nitrogen flow (with or without application of vacuum), the tracer was formulated in physiological saline (ethanol content in the formulated tracer was kept below 10 %).

Preparation of [¹⁸F]FEt-AMC15, [¹⁸F]FPr-AMC15, and [¹⁸F]FBu-AMC15

The procedure described above for [¹⁸F]AMC13 homologues was also used for the preparation of [¹⁸F]AMC15 homologues, with the following modifications:

Instead of (R) N-[7-(methoxymethoxy)chroman-2-yl]methyl 4-hydroxybenzyl amine (1), (R) 1-(4-hydroxyphenyl)-4-(4-(7-hydroxychroman-2-yl)-3-azabutyl)-piperazine (2) was used as the precursor. MOM-protected molecules 4 were the intermediates for the $[^{18}F]AMC15$ homologues (Fig. 2). At the end of the deprotection step, the reaction mixture was diluted with 10 mL water, titrated with 1 M NaOH to pH of 4-7 and then put through the Oasis HLB SPE cartridge. Adsorbed activity was then eluted from the cartridge with 1 mL acetonitrile/ DMF mixture $(1/1 \nu/\nu)$. Eluate was further diluted with water to a total volume of 1.5-2 mL, and the crude product was purified by HPLC using Alltima C18 5µ 250×10 mm column. For [¹⁸F]FEt-AMC15 purification, the column was eluted with a gradient of acetonitrile in 10 mM acetate buffer with pH 4.0 at 5 mL/ min (45 % acetonitrile during the first 2 min, then a linear decrease from 45 % at 2 min to 30 % acetonitrile at 15 min). Retention time of [18F]FEt-AMC15 was 8.4 min. [¹⁸F]FPr-AMC15 and [¹⁸F]FBu-AMC15 purification the column was eluted with 25 % acetonitrile in 10 mM H₃PO₄ at 5 mL/min. Retention times were 9.2 min for [¹⁸F]FPr-AMC15 and 11.0 min for [¹⁸F]FBu-AMC15.

Non-carrier-added [¹⁸F]FEt-AMC15 was found to decompose on the silica plate, so before TLC runs with [¹⁸F]FEt-AMC15 and its longer-chain homologues (including the analysis of plasma and tissue extracts), the start positions of the lanes on the TLC plate were pretreated with AMC15 oxalate standard aqueous solution (2 µg in 2 µL) to hamper the on-plate decomposition.

Characterization and identity confirmation of tracers

All tracers were characterized by analytical radio-HPLC and radio-TLC. Conditions and results of the characterization are presented in Additional file 1: Table S1.

The presence of the corresponding "carrier" (i.e., tracer molecules that contain a [¹⁹F]-atom) in the formulated tracer solutions was confirmed by high performance liquid chromatography-electron spray tandem mass spectrometry (LC-ESI-MS/MS) analysis with the monitoring of fragmentation reactions of the tracer molecules. For all radioligands, ion current peaks with expected M/z ratios giving expected fragmentation products were observed (Additional file 1: Table S2). The identity of [¹⁸F]FEt-AMC13, [¹⁸F]FBu-AMC15, and [¹⁸F]FEt-AMC15 was also confirmed by co-elution with reference products.

LogD measurement

LogD of the tracers was measured using the radioactivity remaining on the HLB SPE cartridges after their elution with ethanol during the tracer formulation procedure (see above). The cartridge was dried with nitrogen for 5 min and eluted with at least 1.5 mL n-octanol. A 500 μ L portion of octanol eluate was mixed with 500 μ L of 1 M phosphate buffer (pH 7.4) and vigorously vortexed for 5 min. The mixture was then centrifuged at 17,000g for 10 min and $3 \times 100 \ \mu$ L aliquots of organic and aqueous phase were loaded into a gamma-counter for radioactivity measurement. To correct for possible presence of radioactive impurities in the tracer, aliquots of organic and aqueous phases were later analyzed by radio-TLC, and the counts obtained from the gamma-counter were corrected according to the percentage of impurities found in aqueous and organic phases. LogD was calculated with the following formula:

$$\text{LogD} = \text{lg} \left(\frac{\text{CPM}_{\text{org}} \times X_{\text{org}}^{\text{tracer}}}{\text{CPM}_{\text{aq}} \times X_{\text{aq}}^{\text{tracer}}} \right)$$

where CPM_{org} is the mean radioactivity of the organic aliquots, $X_{\text{org}}^{\text{tracer}}$ is the mean percentage of radioactivity accounting for intact tracer in organic aliquots (as determined from radio-TLC), and CPM_{aq} and $X_{\text{aq}}^{\text{tracer}}$ are the corresponding values for the aqueous phase.

For each tracer batch used for logD determination, the experiment was performed in duplicate. LogD values are means calculated from at least two independent experiments.

In vitro autoradiography

In vitro autoradiography assay was performed as previously described [24]. Frozen brains of young (10–12 weeks of age; 300–350 g body weight) male Sprague-Dawley rats (Harlan, Netherlands) were cut at -12 °C into sagittal slices 20 µm thick using a Leica microtome, and the slices were thaw-mounted on Superfrost (70 × 22 mm, Fischer) adhesive slides. Only slices containing both striatal and cerebellar regions were used. The slides with slices mounted them were allowed to dry, then put into storage boxes with slica gel bags and stored at -80 °C.

On the day of the experiment (within 1 week from the preparation of the slices), the slides were taken out of storage and allowed to come to room temperature for 5-10 min. Then, 1-1.2 mL of incubation buffer (50 mM Tris-HCl, 5 mM KCl, 2 mM CaCl₂, 2 mM MgCl₂, 120 mM NaCl, pH 7.4 25 °C) was applied per slide, and the slides were pre-incubated for 15 min at room temperature. The buffer was then removed, and the slides were placed into staining jars containing radioligand solution (concentration range of 0.1-10 nM) in the incubation buffer with or without 10 μ M raclopride (D_{2/3} antagonist) or 100 μ M guanosine-5'-triphosphate sodium salt (GTP, stimulator of G-protein uncoupling from the receptors). After 35 min of incubation at 37 °C, slides were washed once with ice-cold incubation buffer (3.5 min) and dipped for 30 s into icecold distilled water to remove buffer salts.

After drying the slides in a stream of room-temperature air, they were exposed on phosphor storage screens for 6–10 h. The storage screens were read by Cyclone Storage Phosphor System (Packard Instruments Co.). Quantification of plate readings was done with OptiQuant software (version 3.00, Packard Instruments Co.).

MicroPET study in rats

Three radioligands were evaluated in vivo: the originally developed [¹⁸F]FBu-AMC13 and [¹⁸F]FEt-AMC15, and [¹⁸F]FEt-AMC13, the homologue that showed the best results in the in vitro autoradiography experiments (see "Results" section).

Animal experiments were performed by licensed investigators in compliance with the Law on Animal Experiments of The Netherlands. The protocol was approved by the Committee on Animal Ethics of the University of Groningen. Young male (10–12 weeks of age, 300–350 g body weight) Sprague-Dawley rats (Harlan, the Netherlands) were used for all experiments. The rats were maintained at a 12-h light/12-h dark regime and were fed standard laboratory chow ad libitum. Distribution of the radiotracers was studied in rats pretreated, at random, with physiological saline (controls) or raclopride tartrate (1 mg/kg calculated per raclopride base, equivalent to 2.9 μ mol/kg). Saline and raclopride were administered intravenously (1 mL/kg injected volume).

An additional group of saline pre-treated rats, injected with $[^{11}C]$ raclopride, served as positive controls for $D_{2/3}R$ -specific brain uptake.

Two rats were scanned simultaneously. Before all manipulations, animals were anesthetized with a mixture of isoflurane/air (inhalation anesthesia, 5 % ratio during induction, 2 % at maintenance). Cannulae were placed into the rats' left femoral arteries and veins; the operation took 45–50 min. The rats were positioned supine inside the camera (Focus 220 microPET, Siemens-Concorde), one above the other with their heads in the camera's field of view. A 515-s transmission scan with a Co-57 point source was performed. Tracer was injected through the venous cannula as a slow bolus (1 mL volume, 60 s long) using an infusion pump. The second (upper) animal was injected 16 min after the first (lower). Injected activities and doses are given in Table 1.

PET acquisition was started during the injection of radioactivity in the first rat. A 106-min long (76 min long in the case of $[^{11}C]$ raclopride) list-mode acquisition protocol was used for the scan.

Throughout the scan, arterial blood samples (15–19 samples; 0.10–0.15 mL each) were withdrawn from each rat through the arterial cannula. Time intervals between samples gradually increased from 5–10 s right after injection to 30 min at the end of the scan. From each sample, 25 μ L aliquots of whole blood were withdrawn. The rest was centrifuged at 3500 g for 5 min; 25 μ L supernatant (plasma) aliquots were withdrawn and deproteinated with 75 μ L of ice-cold acetonitrile. From the resulting 100 μ L, 2–8 μ L portions (i.e., 2–8 % of volume) were taken for thin layer chromatography analysis (radio-TLC) to assess radiometabolite content.

The radioactivity of the plasma and whole blood samples was measured using a well-type gamma-counter (LKB-1282-Compugamma, LKB Wallac).

Radio-TLC was performed on silica plates. Samples of the formulated tracers (diluted) were run along with deproteinated plasma samples to confirm the identity of the parent compound in plasma. Eluent systems used and TLC R_f values recorded are presented in Additional file 1: Table S1.

After the end of the PET scans (i.e., 98 ± 8 min after tracer injection), anesthetized animals were sacrificed by heart extirpation and dissected. Samples of brain and peripheral tissues were taken. All samples were weighed, and their radioactivity was measured in a gamma-counter.

MicroPET image reconstruction and data analysis

List-mode data from the 106-min-long and 76-min-long scans were reframed into, respectively, 90-min-long and 60-min-long dynamic sequences of 6×10 , 4×30 , 2×60 , 1×120 , 1×180 , 4×300 , 3×600 , and (for 90-min-long scans) 2×900 s frames. The data were reconstructed per time frame using an iterative reconstruction algorithm (attenuation-weighted 2-dimensional ordered-subset expectation maximization, provided by Siemens; 4 iterations, 16 subsets; zoom factor 2). Datasets were fully corrected for random coincidences, scatter, and attenuation. Data from the transmission scan were used for attenuation correction. The final datasets consisted of 95 slices, with a slice thickness of 0.8 mm and an in-plane image matrix size of 128×128 and pixel size of 0.47×0.47 mm.

Reconstructed images were analyzed with Inveon 2.0 software (Siemens Medical Solutions, USA, Inc). Regions of interest (ROIs) were drawn manually on a T_2 -weighted MRI template of rat brain around the striatum, brainstem, cortex, hippocampus, hypothalamus, thalamus, olfactory bulbs, and cerebellum around the whole brain and around the pituitary gland. The MRI template was co-registered with the PET scan by image fusion. The time-activity curves (TAC) per ROI were determined in Bq/cm³ units and converted into standardized uptake values (SUVs).

Kinetic analysis is described using the nomenclature proposed by Innis et al. [26]. ROI TACs of [¹⁸F]FEt-AMC13 were analyzed with one-tissue and two-tissue compartmental models of reversible binding (1TCM and 2TCM, respectively) to obtain individual rate constants. From these, the regional distribution volumes ($V_{\rm T}$) were calculated and compared to those obtained with Logan graphical analysis. Binding potentials (BP_{ND}) were calculated from the distribution ratios as $V_{\rm T}$ (target) / $V_{\rm T}$ (cerebellum) – 1, or estimated with the simplified reference

Table 1 Injection data for the PET study

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Tracer	Injected activity, MBq/rat	Injected dose, nmol/kg	Rats per treatment group (saline/raclopride)	Pre-treatment time interval, min		
[¹⁸ F]FEt-AMC13	9.4 ± 1.2	0.55 ± 0.14	4/4	47 ± 27		
[¹⁸ F]FBu-AMC13	20.8 ± 10.6	0.45 ± 0.26 , 10.4, 11.0 ^a	3/3	52 ± 19		
[¹⁸ F]FEt-AMC15	26.6 ± 15.2	1.06 ± 0.65	3/3	48 ± 10		
[¹¹ C]raclopride	16.9 ± 5.3	1.26 ± 0.16	4 (saline only)	36 ± 3		

One rat in the saline-treated group and one rat in the raclopride-treated group were injected, respectively, with 10.4 and 11.0 nmol/kg tracer. As all outcome measures (time-activity curves, uptake ratios) obtained from these animals were very close to the results from other animals in the corresponding groups, these results were not omitted or separated, but combined with the rest of the data

tissue model (SRTM) using the cerebellum as reference region.

Metabolite-corrected plasma-derived arterial input function and whole blood time-activity curve were used for Logan, 1TCM, and 2TCM analyses. Fractional cerebral blood volume was set to 3.6 % [27]. For one rat in the raclopride-treated group, no input curves could be obtained; therefore, its PET data were only analyzed with SRTM.

Ex vivo autoradiography study in rats

To further assess the specific binding of $[^{18}F]$ FEt-AMC13 to the striatal $D_{2/3}R$, two groups of rats were anesthetized, pre-treated with saline (n = 3) or raclopride (n = 3), and 26 ± 3 min later injected with a bolus of $[^{18}F]$ FEt-AMC13 (4.2 ± 1.4 MBq, 0.14 ± 0.04 nmol/kg) into the penile vein. Rats were sacrificed by heart extirpation 35 min later. Brains were quickly extracted and separated into two halves along the sagittal symmetry plane or along the coronal plane spanning the thalamus.

One half of the brain (rostral or randomly taken left/right) was quickly frozen over liquid nitrogen and prepared for cutting on the microtome in the same manner as brains used for in vitro autoradiography [24]. Slices 40 μ m thick were cut, mounted on Superfrost glass slides, permitted to dry, and then directly applied to the phosphor storage screens.

Another half of the brain was homogenized in 3 mL of ice-cold acetonitrile using Heidolph IDAX600 homogenizer at maximum speed. Homogenate was centrifuged at 3500g for 5 min, and the supernatant (containing >95 % total radioactivity) was analyzed by radio-TLC in the same way as described above for the plasma.

D_{2/3}R-specific binding quantification

Degree of $D_{2/3}R$ -specific binding of the tracers was characterized by BP_{ND} (as given above) or the specific binding ratio (SBR) given as target region activity/cerebellar activity - 1. Both were calculated under control conditions and in response to the raclopride challenge.

Statistics

All data are presented as means \pm standard deviations. Comparison of means was done using 2-sided unpaired Welch *t* test. *p* values below 0.05 were considered significant. No multiple comparison corrections were performed.

Results

Preparation of radioligands

The labeling scheme used for the preparation of the tracers is given in Fig. 2. Decay-corrected yields, radiochemical purities, specific activities, and logD values of the prepared tracers are listed in Table 2. Radio-HPLC and radio-TLC characterization data of the prepared are described in Additional file 1: Table S2. For both [¹⁸F]AMC13 and [¹⁸F]AMC15 homologues, reverse-phase radio-HPLC retention times, normal-phase TLC retention factors, and logD values increased with the length of the [¹⁸F]fluoroalkyl chain.

compounds in the decayed samples of purified tracers

In vitro autoradiography

All [¹⁸F]fluoroethyl and [¹⁸F]fluoropropyl compounds showed preferential uptake in the $D_{2/3}$ R-rich striatum (Fig. 3). [¹⁸F]FEt-AMC13 had the highest striatal SBR, followed by [¹⁸F]FPr-AMC13 and [¹⁸F]FEt-AMC15 (Table 3). [¹⁸F]FBu-AMC13 only showed discernible specific binding at the lowest concentration tested.

In the presence of 10 μ M of raclopride, SBR values of all AMC13 homologues decreased by 79–88 %, and those of all AMC15 homologues by 60–64 % compared to control conditions. GTP decreased the SBR values to the same extent, implying that the striatal binding of all tracers was predominantly to the high-affinity subset of the D_{2/3}Rs (Table 3).

In vivo and ex vivo brain uptake of the tracers under control conditions

 $[^{18}F]$ FEt-AMC13 and $[^{18}F]$ FBu-AMC13 showed good blood-brain barrier (BBB) penetration (Fig. 4). Brain uptake peaked at 0.7 % injected dose (ID) and 0.9 % ID, respectively, 3 min after radiotracer injection.

For both tracers, the brain region with the lowest uptake was the cerebellum. [¹⁸F]FBu-AMC13 had the highest uptake in the brainstem, followed by the striatum; striatum-to-cerebellum uptake ratio peaked at 1.41 \pm 0.08 (Additional file 1: Figure S1. [¹⁸F]FEt-AMC13 and [¹¹C]raclopride had the highest uptake in the striatum (Fig. 5). Striatum-to-cerebellum uptake ratio of [¹⁸F]FEt-AMC13 peaked 45 min post-injection at 2.08 \pm 0.25 (Fig. 5b); the same ratio measured by ex vivo autoradiography was 2.81 \pm 0.29 (35 min post-injection). The striatum-to-cerebellum ratio of [¹¹C]raclopride was 7.45 \pm 1.60 (45 min post-injection; Fig. 5d).

[¹⁸F]FEt-AMC15 essentially did not penetrate the BBB. Its in vivo brain uptake peaked at 0.2 % ID 3 min postinjection and fell to 0.04 % ID 60 min post-injection.

Ex vivo measurements of the uptake of the tracers in the brain and peripheral tissues performed after the PET scan are given, respectively, in Table 4 and Additional file 1: Table S3.

Tracer metabolism

[¹⁸F]FEt-AMC13 and [¹⁸F]FBu-AMC13 were quickly metabolized in the plasma, while [¹⁸F]FEt-AMC15 was more stable (Fig. 6). Tracers were eliminated from the

Compound	Isolated radiochemical yield, %	Radiochemical purity, %	End-of-synthesis specific activity, GBg/µmol	logD
[¹⁸ F]FEt-AMC13	12±6	>95	102±30	1.67 ± 0.07
[¹⁸ F]FPr-AMC13	7±6	>95	29 ± 27	1.98 ± 0.08
[¹⁸ F]FBu-AMC13	8±7	93±2	85 ± 83	2.50 ± 0.06
[¹⁸ F]FEt-AMC15	5 ± 3	93±2	25 ± 15	1.48 ± 0.002
[¹⁸ F]FPr-AMC15	3 ± 1	93 ± 5	9 ± 7	2.39 ± 0.001
[¹⁸ F]FBu-AMC15	6 ± 0.3	93±6	10±3	2.71 ± 0.07

 Table 2 Synthetic and analytical data for [¹⁸F]AMC13 and [¹⁸F]AMC15 homologues

Values are means \pm SD of \geq 3 determinations from \geq 2 independent experiments

plasma with half-lives of 3 to 12 min. Radiometabolites of all tracers in the plasma and in the brain tissue extract were considerably more hydrophilic than their corresponding parent compounds, as they all had TLC R_f values of 0, compared to R_f of 0.16–0.56 for the tracers themselves (Additional file 1: Table S1). Intact [¹⁸F]FEt-AMC13 constituted 92 % of radioactivity from brain tissue extract 35 min post-injection (Additional file 1: Figure S2).

Uptake in the parietal bone was lower than in the brain tissue for all $[^{18}F]AMC13$ homologues (Table 4), suggesting low levels of in vivo defluorination.

Kinetic modeling for [¹⁸F]FEt-AMC13

2TCM and SRTM fits of ROI TACs were consistently better than 1TCM fits (Additional file 1: Figure S3) as assessed by Akaike's information criterion, but the estimates of individual rate constants of 2TCM were highly variable and often physiologically implausible (Additional file 1: Table S4).

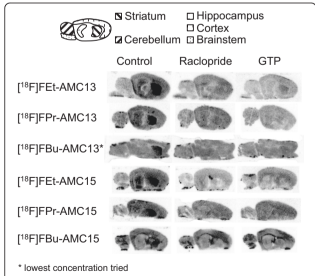


Fig. 3 Representative in vitro autoradiography images of rat brain slices for the assayed radioligands. *Left column*—slides incubated with radioligand only. *Middle column*—radioligand and 10 μ M raclopride. *Right column*—radioligand and 100 μ M GTP

The most robust ROI $V_{\rm T}$ estimates were obtained from the Logan analysis (Additional file 1: Table S5). Logan $V_{\rm T}$ values correlated better with 2TCM $V_{\rm T}$ (r = 0.97) than with 1TCM $V_{\rm T}$ values (r = 0.89; Additional file 1: Figure S4). No robust BP_{ND} estimates could be obtained from the k₃/k₄ rate constant ratio of 2TCM (data not shown), but BP_{ND} estimates calculated from 1TCM, 2TCM, and Logan $V_{\rm T}$ data and obtained by SRTM correlated well with each other for all regions except the pituitary and olfactory bulbs (Additional file 1: Figure S5).

The highest BP_{ND} values in control rats were found in the striatum (1TCM 0.63 ± 0.08; 2TCM 0.56 ± 0.05, Logan 0.52 ± 0.01; SRTM 0.51 ± 0.02).

Blocking experiments

Pre-treatment with 1 mg/kg (2.9 µmol/kg) raclopride decreased peak PET-based striatal SBR of [¹⁸F]FEt-AMC13 by 17 % (from 1.08 ± 0.25 to 0.90 ± 0.11), though the SBR change was not statistically significant (p = 0.25). Striatal SBR based on after-scan biodistribution measurements decreased by 31 % (from 1.07 ± 0.10 to 0.74 ± 0.27), but was also not statistically significant (p = 0.17). Ex vivo autoradiography-based SBR fell by 39 % (from 1.81 ± 0.29 to 1.06 ± 0.17), which was statistically significant (p < 0.01), and the blocking of D_{2/3}R specific striatal binding was clearly visible on the autoradiography images (Additional file 1: Figure S6).

Striatal BP_{ND} fell by 18–27 % (p < 0.05 for 2TCM, Logan and SRTM; see Table 5 and Additional file 1: Table S6) depending on the kinetic model used. Modified Lassen plot analysis [28] estimated the raclopride-induced D_{2/3}R occupancy at 20–25 % (p < 0.05 for 1TCM and 2TCM), while 41–55 % of striatal $V_{\rm T}$ and 11–26 % of cerebellar $V_{\rm T}$ were estimated to be potentially displaceable, i.e., represent D_{2/3}R specific binding (Additional file 1: Figure S7).

For [¹⁸F]FBu-AMC13 pre-treatment with raclopride, decreased absolute uptake in all brain regions, and striatum-to-cerebellum uptake ratios were higher in raclopride-treated group than in the saline-treated group (Additional file 1: Figure S1c).

Tracer	Tracer	Striatal SBR ^b (% change relative to control condition)				
	concentration, nM ^a	Control	Raclopride	GTP		
[¹⁸ F]FEt-AMC13	0.9 ± 0.7 (n = 3)	3.40 ± 1.25	0.63 ± 0.53 (-81%)*	0.66 ± 0.52 (-81%)*		
[¹⁸ F]FPr-AMC13	1.4 ± 1.6 (n = 3)	2.07 ± 1.06	0.41 ± 0.23 (-80%)*	0.42 ± 0.23 (-80%)*		
[¹⁸ F]FBu-AMC13 ^c	0.2 (<i>n</i> = 1)	0.85 ± 0.23	0.10 ± 0.19 (-88%)*	0.20 ± 0.16 (-76%)*		
	4.5 ± 3.6 (n = 3)	0.19 ± 0.18	0.04 ± 0.22 (-79%)	-		
[¹⁸ F]FEt-AMC15 ^c	5.8 ± 1.6 (n = 4)	1.64 ± 1.04	0.65 ± 0.37 (-60%)*	0.67 ± 0.28 (-59%)*		
[¹⁸ F]FPr-AMC15	5.0 ± 7.2 (n = 3)	1.01 ± 0.46	0.39 ± 0.22 (-61%)*	0.37 ± 0.24 (-63%)*		
[¹⁸ F]FBu-AMC15	2.3 ± 1.8 (n = 3)	0.69 ± 0.30	0.25 ± 0.23 (-64%)*	0.23 ± 0.23 (-67%)*		

 Table 3 Quantitative results of in vitro autoradiography

*p < 0.05 relative to control values, 2-sided Welch test

^aConcentrations are means \pm SD from *n* independent experiments

^bSpecific binding ratio (SBR) values are means ± SD from the total number of slides (3–28 per condition) assayed in all experiments

^cPublished earlier [24]

For [¹⁸F]FEt-AMC15, pre-treatment with raclopride caused no detectable alterations of radioactivity distribution in the brain (data not shown).

Discussion

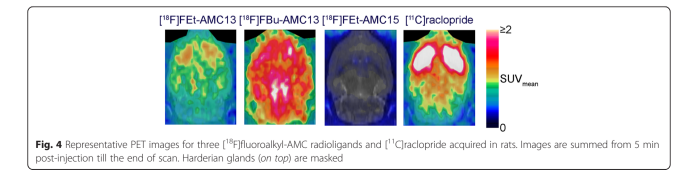
We have previously described the design, synthesis, and in vitro evaluation of a series of high-affinity $D_{2/3}$ agonists based on the AMC scaffold. Two compounds from that series, FBu-AMC13 and FEt-AMC15, were confirmed to have high-affinity towards $D_{2/3}R$ and act as agonists at D_2R [24]. They were [¹⁸F]-labeled and showed favorable results in the in vitro autoradiography experiments. In this study, we have evaluated these compounds in vivo and further optimized their characteristics by shortening or lengthening the [¹⁸F]fluoroalkyl chains in their structures. We did not directly measure the affinities and intrinsic activities of the shorter-chain homologues of FBu-AMC13 and longerchain homologues of FEt-AMC15 towards D_{2/3}R, assuming that changing the fluoroalkyl chain would have negligible influence on these parameters, because this fluoroalkyl chain is situated outside the main AMC pharmacophore [24].

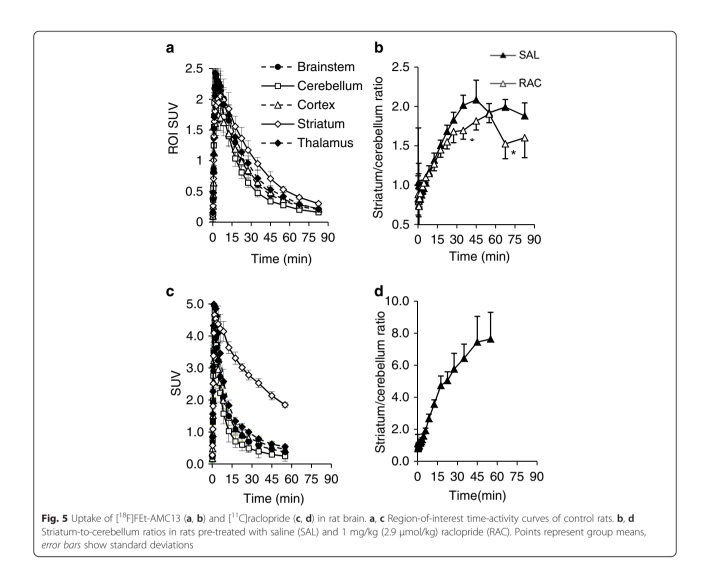
Both raclopride and GTP decreased striatal SBRs of all tracers significantly and to a similar extent, which was consistent across the homologous series (Table 3). This indicates that all our tracers specifically bind to striatal dopamine $D_{2/3}$ receptors in their G-protein-coupled "high-affinity state." Tracers with a shorter alkyl chain showed higher in vitro autoradiography-based striatal SBR values (Et > Pr > Bu). These findings are in agreement with our hypothesis that varying the $[^{18}\mathrm{F}]$ fluoroalkyl chain length would influence non-specific binding, but not the $D_{2/3}R$ -affinity or agonism of the tracers.

In vivo, [¹⁸F]FEt-AMC15 showed very low levels of BBB penetration, which may be the consequence of the tracer's low lipophilicity and/or may imply that it is a substrate for one of the drug efflux pump proteins such as P-glycoprotein. Meanwhile, [¹⁸F]FEt-AMC13, which is the short-alkyl-chain homologue of [¹⁸F]FBu-AMC13, showed greatly improved specific-to-non-specific binding ratios compared to the "original" radioligand, while retaining good brain penetration and favorable metabolic profile.

Our approach has proven to be a useful and relatively simple tactic for the adjustment of the radioligands' lipophilicity with the goal of optimizing their non-specific binding and brain penetration.

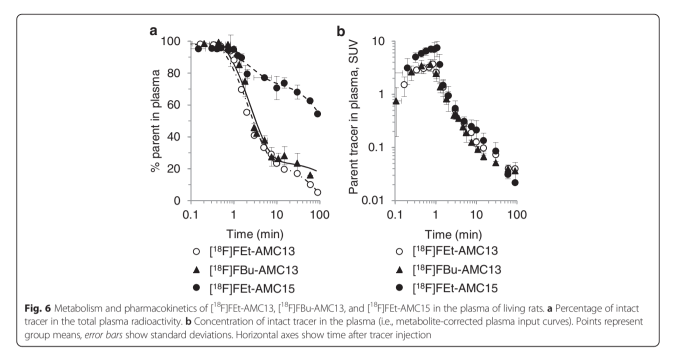
The time-activity curves of [¹⁸F]FEt-AMC13 were better approximated with the 2TCM model than with 1TCM model, and 2TCM-derived $V_{\rm T}$ and BP_{ND} estimates correlated better than 1TCM-derived $V_{\rm T}$ and BP_{ND} estimates with the same data obtained from Logan analysis that makes no assumptions regarding the





Tracer	[¹⁸ F]FEt-AMC13		[¹⁸ F]FBu-AMC13		[¹⁸ F]FEt-AMC15	
Treatment	Saline	Raclopride $(n = 4)$	Saline (<i>n</i> = 3)	Raclopride	Saline	Raclopride
Region	(<i>n</i> = 4)			(n = 3)	(n = 3)	(<i>n</i> = 3)
Striatum	0.28 ± 0.03 (1)	0.23 ± 0.08	1.60 ± 0.33 (5)	1.21 ± 0.14	0.02 ± 0.02	0.03 ± 0.01
Hippocampus	0.22 ± 0.02 (2)	0.23 ± 0.08	1.50 ± 0.25 (6)	1.21 ± 0.06	0.03 ± 0.02	0.03 ± 0.01
Thalamus + hypothalamus	0.19 ± 0.003 (4)	0.19 ± 0.05	2.07 ± 0.70 (1)	1.07 ± 0.10	0.03 ± 0.01	0.03 ± 0.01
Cortex	0.18±0.01 (5)	0.16 ± 0.04	1.50 ± 0.32 (7)	1.01 ± 0.11	0.03 ± 0.003	0.03 ± 0.01
Brainstem	0.18±0.02 (6)	0.16 ± 0.04	1.97 ± 0.42 (2)	1.25 ± 0.14	0.03 ± 0.004	0.03 ± 0.01
Olfactory bulbs	0.17 ± 0.01 (7)	0.15 ± 0.04	1.85 ± 0.64 (3)	0.86 ± 0.09	0.03 ± 0.001	0.03 ± 0.02
Rest of brain	0.20±0.02 (3)	0.18 ± 0.03	1.60 ± 0.25 (4)	1.10 ± 0.17	0.02 ± 0.02	0.03 ± 0.02
Cerebellum	0.13 ± 0.01 (8)	0.13 ± 0.02	1.11 ± 0.29 (8)	0.73 ± 0.08	0.03 ± 0.01	0.03 ± 0.01
Mean brain uptake	0.18 ± 0.01	0.16 ± 0.04	1.56 ± 0.33	1.03 ± 0.11	0.03 ± 0.01	0.03 ± 0.01
Pituitary	0.76 ± 0.08	0.59 ± 0.20	3.19 ± 0.29	2.72 ± 0.77	1.36 ± 0.30	1.08 ± 0.27
Parietal bone	0.10 ± 0.01	0.10 ± 0.01	0.61 ± 0.17	0.70 ± 0.18	0.09 ± 0.03	0.08 ± 0.03

Data are presented as means \pm standard deviations of standard uptake values. The ranking of the uptake magnitude (highest to lowest) among brain regions within the BBB is shown in parentheses (only for [¹⁸F]AMC13 homologues)



number of kinetic compartments. However, the 2TCM model did not always provide reliable estimates of the individual rate constants of [¹⁸F]FEt-AMC13 uptake in the brain. This may have been caused by the quality of input data: insufficient counts or imprecise estimates of the plasma input can make the optimization routine converge at a local minimum. Fortunately, the estimates of $V_{\rm T}$ and BP_{ND} were more robust and consistent, so these were used for the evaluation of [¹⁸F]FEt-AMC13.

Estimation of the magnitude of the $D_{2/3}R$ blockade induced by 1 mg/kg (2.9 μ mol/kg) raclopride in rats' striata in vivo performed with the use of different approaches produced consistent results: changes in striatal BP_{ND} and (PET and autoradiography-derived) SBR were similar to the modified Lassen plot-based estimate of $D_{2/3}R$ occupancy by raclopride. Depending on the estimate, raclopride pre-

treatment blocked 17–39 % of D_{2/3}R-specific signal in the striatum. Though a bolus dose injected in an in vivo experiment cannot be directly compared with ligand concentration applied in an in vitro experiment, our in vivo findings are also arguably consistent with our in vitro findings, where a higher concentration of raclopride (10 μ mol/L) used in the in vitro autoradiography study induced a greater decrease (81 %) in striatal SBR in rat brain slices compared to control conditions.

Our in vivo findings, however, do not agree with the published studies which reported >90 % displacement of apparent D_{2/3}R-specific striatal uptake of [¹¹C]MNPA and [¹¹C]PHNO in Sprague-Dawley rats with 1–2 mg/kg (2.9–5.8 µmol/kg) raclopride [29–31]. Even a fivefold lower dose (0.2 mg/kg, 0.58 µmol/kg) displaced over 80 % of apparent D_{2/3}R-specific [¹¹C](+)PHNO uptake [30]. It is

Table 5 2TCM and SRTM-derived BP	D Of	¹⁸ F1FFt-AMC13 r	per brain regio	n in control	and raclopride-treated rats

Region of	2TCM BP _{ND} (V _T -base	ed)	SRTM BP _{ND}		
interest	Control	Raclopride	Control	Raclopride	
Striatum	0.56 ± 0.05	0.41 ± 0.05 (-27%)*	0.51 ± 0.02	0.38 ± 0.05 (-26%)**	
Hippocampus	0.30 ± 0.07	0.26 ± 0.06 (-13%)	0.26 ± 0.04	0.23 ± 0.05 (-13%)	
Thalamus	0.33 ± 0.02	0.30 ± 0.01 (-10%)*	0.28 ± 0.06	0.26 ± 0.02 (-7%)	
Hypothalamus	0.22 ± 0.03	0.18±0.06 (-19%)	0.21 ± 0.03	0.16±0.04 (-23%)	
Cortex	0.13 ± 0.04	0.10 ± 0.02 (-27%)	0.11 ± 0.03	0.08 ± 0.02 (-24%)	
Brainstem	0.22 ± 0.02	0.10 ± 0.02 (-53%)**	0.18 ± 0.04	0.07 ± 0.03 (-60%)**	
Olfactory bulbs	0.08 ± 0.13	0.01 ± 0.04 (-88%)	0.10 ± 0.13	0.11 ± 0.05 (+10%)	
Pituitary	0.36 ± 0.07	0.13±0.04 (-23%)	0.71 ± 0.32	0.55 ± 0.53 (-22%)	

Data are presented as means \pm SD, n = 4 for the control group and n = 3 for the raclopride-treated group (except for the SRTM BP_{ND} values, where n = 4). Next to the BP_{ND} values of the raclopride-treated group, percentage of change relative to the control group is shown

*p < 0.05; **p < 0.01, 2-sided Welch test

not clear why the apparent displacement of [18F]FEt-AMC13 from the striatum by 1 mg/kg raclopride is lower than what is reported for other D_{2/3}R tracers in the literature. It is unlikely that we violated the so-called tracer conditions because crude estimation of the D_{2/3}R occupancy by [¹⁸F]FEt-AMC13 itself using the approach described by Skinbjerg et al. [32] produced the values of no more than 3 % in all our experiments. It may be that the signal-to-noise ratio in our data was insufficient, which made it difficult to precisely estimate the contribution of specific binding to total tracer binding in various ROIs. The presence of specific binding of [¹⁸F]FEt-AMC13 to sites other than $D_{2/3}R$, which has a comparatively high density in the striatum, may also be implied. We have previously demonstrated that FBu-AMC13, a longer-chain homologue of FEt-AMC13, has low affinity towards D_1 receptors (7 µM on average) [24], while (R)-2-[(benzylamino)methyl]chroman-7-ol and 2-[(4-hydroxybenzylamino) methyl]chroman-7-ol, compounds structurally closely related to FEt-AMC13, were shown by Mewshaw and coworkers to be selective for D_{2/3}R against serotonin-1A and adrenergic α_1 receptors [23]. In the in vitro autoradiography study, presence of 10 µmol/L raclopride decreased striatal SBR of [¹⁸F]FEt-AMC13 by 81 %, which also does not support the hypothesis of the existence of significant non- $D_{2/}$ ₃R-specific binding of [¹⁸F]FEt-AMC13 in the striatum. Still, a separate study of pharmacological selectivity of FEt-AMC13, including direct determination of the affinities of FEt-AMC13 towards high- and low-affinity states of D_{2/3}R, is necessary to clarify this issue.

Apart from the striatum, the olfactory bulbs and pituitary gland are known to contain high densities of $D_{2/}$ ₃R available to agonist radioligands [33]. However, no blocking effect of raclopride could be observed in these regions (Tables 4 and 5), and the relatively high tracer uptake in the pituitary is more likely to be explained by the pituitary being situated outside the BBB than by the presence of $D_{2/3}$ R-specific binding of [¹⁸F]FEt-AMC13 there.

A striatum-to-cerebellum ratio of 2.08 is on par with the value of 2 reported in rats for [¹⁸F]5-OH-FPPAT [20] and with the value of 1.97 obtained for our other separately evaluated AMC derivative [¹⁸F]AMC20 [34]. However, the striatal BP_{ND} values of 0.51–0.63 obtained for [¹⁸F]FEt-AMC13 are lower than the values of at least 0.8–1.0 reported in rodents for existing [¹¹C]-labeled D_{2/3} agonists [19, 29, 31]. Non-specific binding of the tracer in the tissue can slow down the association rate of the tracer-receptor binding by leaving less free tracer in the tissue. The launch of the canonical signaling cascade induced by agonist binding eventually leads to the uncoupling of the G-protein from the receptor and the relaxation of the receptor into its low-affinity state [18, 35], which may increase the effective dissociation rate of an agonist

tracer from the receptors. To find out which factors limit the signal-to-noise ratio of [¹⁸F]FEt-AMC13, a more thorough study of its binding kinetics in a more controlled environment (e.g., in vitro) is necessary.

Conclusions

We have evaluated two homologous series of agonist radiopharmaceuticals based on the AMC scaffold that bind with high affinity and selectivity to $D_{2/3}R$ in vitro. Varying the length of the [¹⁸F]fluoroalkyl group allowed us to significantly improve the specific-to-non-specific binding ratio of one of our original structures. The resulting ligand, [¹⁸F]FEt-AMC13, demonstrated specific binding to the striatal $D_{2/3}R$ in vitro as well as in vivo. In vitro findings also confirmed the tracer's binding to the high-affinity state of $D_{2/3}R$. Further investigation of the binding characteristics of FEt-AMC13 and optimization of its molecular structure can lead to an improved [¹⁸F]-labeled radioligand suitable for clinical $D_{2/3}R$ imaging by PET.

Additional file

Additional file 1: Supplementary material. Supplemental tables and supplemental figures referred to in this article.

Competing interests

MCM is an employee of Boehringer Ingelheim, a company marketing the dopamine D_2R agonist pramipexole. JB is consultant at GE Healthcare, a company marketing the $D_{2/3}R$ radiopharmaceutical $[^{123}]JBZM$. The other authors declare that they have no competing interest.

Authors' contributions

VS synthesized the [¹⁸F]-labeled aminomethylchroman compounds, performed the in vitro and in vivo experiments, analyzed the data, and drafted the manuscript. JPvW helped with the design of the in vitro experiments and with the drafting of the manuscript. HMJ and MF synthesized the "cold" aminomethylchroman compounds and precursors for radiolabeling and provided critical feedback on the manuscript. RAJOD helped with the general design of the study and provided critical feedback on the manuscript. MCM helped with the design of the in vitro experiments and provided critical feedback on the manuscript. MCM helped with the design of the invitro experiments and provided critical feedback on the manuscript. JB helped with the interpretation of the results and critically revised the manuscript. PHE helped with the design of the radiosynthetic protocols and interpretation of the results and critically revised the manuscript. All authors read and approved the final manuscript.

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References

- 1. Howes OD, Kapur S. The dopamine hypothesis of schizophrenia: version III—the final common pathway. Schizophr Bull. 2009;35(3):549–62.
- Booij J, Tissingh G, Winogrodzka A, van Royen EA. Imaging of the dopaminergic neurotransmission system using single-photon emission tomography and positron emission tomography in patients with parkinsonism. Eur J Nucl Med. 1999;26(2):171–82.
- Volkow ND, Fowler JS, Wang G-J, Swanson JM. Dopamine in drug abuse and addiction: results from imaging studies and treatment implications. Mol Psychiatry. 2004;9(6):557–69.
- Volkow ND, Fowler JS, Wang GJ, Baler R, Telang F. Imaging dopamine's role in drug abuse and addiction. Neuropharmacology. 2009;56 Suppl 1:3–8.
- 5. Elsinga PH, Hatano K, Ishiwata K. PET tracers for imaging of the dopaminergic system. Curr Med Chem. 2006;13(18):2139–53.
- Chio CL, Lajiness ME, Huff RM. Activation of heterologously expressed D3 dopamine receptors: comparison with D2 dopamine receptors. Mol Pharmacol. 1994;45(1):51–60.
- Sibley DR, De Lean A, Creese I. Anterior pituitary dopamine receptors. Demonstration of interconvertible high and low affinity states of the D-2 dopamine receptor. J Biol Chem. 1982;257(11):6351–61.
- Liu IS, George SR, Seeman P. The human dopamine D2(Longer) receptor has a high-affinity state and inhibits adenylyl cyclase. Brain Res Mol Brain Res. 2000;77(2):281–4.
- George SR, Watanabe M, Seeman P. Dopamine D 2 receptors in the anterior pituitary: a single population without reciprocal antagonist/agonist states. J Neurochem. 1985;44(4):1168–77.
- Skinbjerg M, Sibley DR, Javitch JA, Abi-Dargham A. Imaging the high-affinity state of the dopamine D 2 receptor in vivo: fact or fiction? Biochem Pharmacol. 2012;83(2):193–8.
- Cumming P, Wong DF, Gillings N, Hilton J, Scheffel U, Gjedde A. Specific binding of [11C]raclopride and N-[3H]propyl-norapomorphine to dopamine receptors in living mouse striatum: occupancy by endogenous dopamine and guanosine triphosphate-free G protein. J Cereb Blood Flow Metab. 2002;22:596–604.
- McCormick PN, Ginovart N, Wilson AA. Isoflurane anaesthesia differentially affects the amphetamine sensitivity of agonist and antagonist D2/D3 positron emission tomography radiotracers: implications for in vivo imaging of dopamine release. Mol Imaging Biol. 2011;13:737–46.
- Ginovart N, Galineau L, Willeit M, Mizrahi R, Bloomfield PM, Seeman P, et al. Binding characteristics and sensitivity to endogenous dopamine of [11C]-(+)-PHNO, a new agonist radiotracer for imaging the high-affinity state of D2 receptors in vivo using positron emission tomography. J Neurochem. 2006;97:1089–103.
- Narendran R, Hwang DR, Slifstein M, Talbot PS, Erritzoe D, Huang Y, et al. In vivo vulnerability to competition by endogenous dopamine: comparison of the D2 receptor agonist radiotracer (–)-N-[11C]propyl-norapomorphine ([11C]NPA) with the D2 receptor antagonist radiotracer [11C]-raclopride. Synapse. 2004;52:188–208.
- Seneca N, Finnema SJ, Farde L, Gulyas B, Wikstrom HV, Halldin C, et al. Effect of amphetamine on dopamine D2 receptor binding in nonhuman primate brain: a comparison of the agonist radioligand [11C]MNPA and antagonist [11C]raclopride. Synapse. 2006;59:260–9.
- Narendran R, Mason NS, Laymon CM, Lopresti BJ, Velasquez ND, May MA, et al. A comparative evaluation of the dopamine D(2/3) agonist radiotracer [11C](-)-N-propyl-norapomorphine and antagonist [11C]raclopride to measure amphetamine-induced dopamine release in the human striatum. J Pharmacol Exp Ther. 2010;333(2):533–9.
- Shotbolt P, Tziortzi AC, Searle GE, Colasanti A, van der Aart J, Abanades S, et al. Within-subject comparison of [(11)C]-(+)-PHNO and [(11)C]raclopride sensitivity to acute amphetamine challenge in healthy humans. J Cereb Blood Flow Metab. 2012;32(1):127–36.
- Van Wieringen J-P, Booij J, Shalgunov V, Elsinga P, Michel MC. Agonist high- and low-affinity states of dopamine D2 receptors: methods of detection and clinical implications. Naunyn Schmiedebergs Arch Pharmacol. 2013;386(2):135–54.

- Finnema SJ, Bang-Andersen B, Wikström HV, Halldin C. Current state of agonist radioligands for imaging of brain dopamine D2/D3 receptors in vivo with positron emission tomography. Curr Top Med Chem. 2010;10(15):1477–98.
- Shi B, Narayanan TK, Christian BT, Chattopadhyay S, Mukherjee J. Synthesis and biological evaluation of the binding of dopamine D2/D3 receptor agonist, (R, S)-5-hydroxy-2-(N-propyl-N-(5'-(18F)fluoropentyl) aminotetralin ((18)F-5-OH-FPPAT) in rodents and nonhuman primates. Nucl Med Biol. 2004;31(3):303–11.
- Vasdev N, Seeman P, Garcia A, Stableford WT, Nobrega JN, Houle S, et al. Syntheses and in vitro evaluation of fluorinated naphthoxazines as dopamine D2/D3 receptor agonists: radiosynthesis, ex vivo biodistribution and autoradiography of [18F]F-PHNO. Nucl Med Biol. 2007;34(2):195–203.
- Finnema SJ, Stepanov V, Nakao R, Sromek AW, Zhang T, Neumeyer JL, et al. 18F-MCL-524, an 18F-labeled dopamine D2 and D3 receptor agonist sensitive to dopamine: a preliminary PET study. J Nucl Med. 2014;55(7):1164–70.
- Mewshaw RE, Kavanagh J, Stack G, Marquis KL, Shi X, Kagan MZ, et al. New generation dopaminergic agents. 1. Discovery of a novel scaffold which embraces the D2 agonist pharmacophore. Structure-activity relationships of a series of 2-(aminomethyl)chromans. J Med Chem. 1997;40(26):4235–56.
- Van Wieringen J-P, Shalgunov V, Janssen HM, Fransen PM, Janssen AMG, Michel MC, et al. Synthesis and characterization of a novel series of agonist compounds as potential radiopharmaceuticals for imaging dopamine D2/3 receptors in their high-affinity state. J Med Chem. 2014;57(2):391–410.
- Kämäräinen EL, Kyllönen T, Airaksinen A, Lundkvist C, Yu M, Någren K, et al. Preparation of [¹⁸F]β-CFT-FP and [11C]β-CFT-FP, selective radioligands for visualisation of the dopamine transporter using positron emission tomography (PET). J Labelled Compd Radiopharm. 2000;43(12):1235–44.
- Innis RB, Cunningham VJ, Delforge J, Fujita M, Gjedde A, Gunn RN, et al. Consensus nomenclature for in vivo imaging of reversibly binding radioligands. J Cereb Blood Flow Metab. 2007;27(9):1533–9.
- Julien-Dolbec C, Tropres I, Montigon O, Reutenauer H, Ziegler A, Decorps M, et al. Regional response of cerebral blood volume to graded hypoxic hypoxia in rat brain. Br J Anaesth. 2002;89:287–93.
- Cunningham VJ, Rabiner EA, Slifstein M, Laruelle M, Gunn RN. Measuring drug occupancy in the absence of a reference region: the Lassen plot re-visited. J Cereb Blood Flow & Metab. 2009;30:46–50.
- Seneca N, Zoghbi SS, Skinbjerg M, Liow JS, Hong J, Sibley DR, et al. Occupancy of dopamine D2/3 receptors in rat brain by endogenous dopamine measured with the agonist positron emission tomography radioligand [11C]MNPA. Synapse. 2008;62(10):756–63.
- Wilson AA, McCormick P, Kapur S, Willeit M, Garcia A, Hussey D, et al. Radiosynthesis and evaluation of [11C]-(+)-4-propyl-3,4,4a,5,6,10bhexahydro-2H-naphtho[1,2-b][1,4]oxazin-9-ol as a potential radiotracer for in vivo imaging of the dopamine D2 high-affinity state with positron emission tomography. J Med Chem. 2005;48(12):4153–60.
- Egerton A, Hirani E, Ahmad R, Turton DR, Brickute D, Rosso L, et al. Further evaluation of the carbon11-labeled D(2/3) agonist PET radiotracer PHNO: reproducibility in tracer characteristics and characterization of extrastriatal binding. Synapse. 2010;64(4):301–12.
- 32. Skinbjerg M, Seneca N, Liow J-S, Hong J, Weinshenker D, Pike VW, et al. Dopamine beta-hydroxylase-deficient mice have normal densities of D(2) dopamine receptors in the high-affinity state based on in vivo PET imaging and in vitro radioligand binding. Synapse. 2010;64(9):699–703.
- Levant B, Grigoriadis DE, DeSouza EB. [3H]Quinpirole binding to putative D2 and D3 dopamine receptors in rat brain and pituitary gland: a quantitative autoradiographic study. J Pharmacol Exp Ther. 1993;264:991–1001.
- Shalgunov V, van Wieringen JP, Janssen HM, Fransen PM, Dierckx RAJO, Michel MC, et al. Synthesis and evaluation in rats of the dopamine D2/3 receptor agonist 18F-AMC20 as a potential radioligand for PET. J Nucl Med. 2015;56(1):133–9.
- van Wieringen JP, Michel MC, Janssen HM, Janssen AG, Elsinga PH, Booij J. Agonist signalling properties of radiotracers used for imaging of dopamine D2/3 receptors. EJNMMI Res. 2014;doi:10.1186/s13550-014-0053-3.