Toward in vivo proof of binding of $^{18}$F-labeled inhibitor $[^{18}$F$]^{\text{TRACK}}$ to peripheral tropomyosin receptor kinases

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Abstract

**Background:** Tropomyosin receptor kinases (TrkA, TrkB, TrkC) are a family of tyrosine kinases primarily expressed in neuronal cells of the brain. Identification of oncogenic alterations in Trk expression as a driver in multiple tumor types has increased interest in their role in human cancers. Recently, first- and second-generation $^{11}$C and $^{18}$F-labeled Trk inhibitors, e.g., $[^{18}$F$]^{\text{TRACK}}$, have been developed. The goal of the present study was to analyze the direct interaction of $[^{18}$F$]^{\text{TRACK}}$ with peripheral Trk receptors in vivo to prove its specificity for use as a functional imaging probe.

**Methods:** In vitro uptake and competition experiments were carried out using the colorectal cancer cell line KM12. Dynamic PET experiments were performed with $[^{18}$F$]^{\text{TRACK}}$, either alone or in the presence of amitriptyline, an activator of Trk, entrectinib, a Trk inhibitor, or unlabeled reference compound TRACK in KM12 tumor-bearing athymic nude mice as well as B6;129S2-Ntrk2$^{tm1Bbd}$/J mice. Western blot and immunohistochemistry experiments were done with KM12 tumors, brown adipose tissue (BAT), and brain tissue samples.

**Results:** Uptake of $[^{18}$F$]^{\text{TRACK}}$ was increasing over time reaching $208 \pm 72\%$ radioactivity per mg protein ($n = 6/2$) after 60 min incubation time. Entrectinib and TRACK competitively blocked $[^{18}$F$]^{\text{TRACK}}$ uptake in vitro (IC$_{50}$ 30.9 ± 3.6 and 29.4 ± 9.4 nM; both $n = 6/2$). $[^{18}$F$]^{\text{TRACK}}$ showed uptake into KM12 tumors (SUV$_{\text{mean,60 min}}$ 0.43 ± 0.03; $n = 6$). Tumor-to-muscle ratio reached 0.9 (60 min) and 1.2 (120 min). In TrkB expressing BAT, $[^{18}$F$]^{\text{TRACK}}$ uptake reached SUV$_{\text{mean,60 min}}$ 1.32 ± 0.08 ($n = 7$). Activation of Trk through amitriptyline resulted in a significant radioactivity increase of 21% in KM12 tumor (SUV$_{\text{mean,60 min}}$ from 0.53 ± 0.01 to 0.43 ± 0.03; $n = 6$; $p < 0.05$) and of 21% in BAT (SUV$_{\text{mean,60 min}}$ from 1.32 ± 0.08; $n = 5$ to 1.59 ± 0.07; $n = 6$; $p < 0.05$) respectively. Immunohistochemistry showed TrkB > TrkA expression on BAT fat cells, but TrkA > TrkB in whole brain. WB analysis showed sevenfold higher TrkB expression in BAT versus KM12 tumor tissue.

**Conclusion:** The present data show that radiotracer $[^{18}$F$]^{\text{TRACK}}$ can target peripheral Trk receptors in human KM12 colon cancer as well as brown adipose tissue as confirmed through in vitro and in vivo blocking experiments. Higher TrkB versus TrkA protein expression was detected in brown adipose tissue of mice confirming a peripheral functional role of brain-derived neurotrophic factor in adipose tissue.

**Keywords:** TrkA, TrkB, PET, Molecular imaging, Radiotracer, $[^{18}$F$]^{\text{TRACK}}$

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transmembrane region and an intracellular section containing the tyrosine kinase domain [1, 2]. Extracellular regions of each Trk subtype bind a preferred neurotransmitter: nerve growth factor (NGF) on TrkA, brain-derived neurotrophic factor (BDNF), and neurotrophin-4 on TrkB and neurotrophin-3 on TrkC [3–9]. The intracellular domain endogenously binds ATP [10, 11].

Trk receptors play an important role during prenatal development of central and peripheral nervous system and normal brain function [10, 12]. Neurotrophin activation of Trk has an impact on neuronal events such as neuronal cell differentiation and survival, cell proliferation, synaptic formation and plasticity, membrane trafficking, and axon and dendrite formation [2, 13, 14]. Neurodegenerative diseases such as Alzheimer's and Parkinson's induce changes in Trk expression and signaling [15–19].

Identification of oncogenic alterations in peripheral Trk expression has stimulated interest in their role in human cancers [1]. Unrestrained activation of Trk-dependent pathways resulting from Trk fusion proteins leads to cell transformation, growth, and proliferation [1, 2, 20, 21]. However, Trk fusions are rare and found in only <1% of solid tumors [21]. A recent immunohistochemical patient analysis found Trk fusions in only 31 (0.3%) out of 11,502 patients [22]. Other studies detected TrkA fusions in 3.3% lung cancers, 0.5–2% colorectal cancers, <12% papillary thyroid carcinomas, TrkA/B/C fusions in 7% pediatric gliomas, and TrkC fusions in 92–100% secretory mammary tumors [23, 24]. Targeting Trk fusions represents one of the first examples of a 'personalized medicine' approach [25]. Two pan-Trk inhibitors have been recently approved for clinical treatment, larotrectinib [26] and entrectinib [27]. For a patient's inclusion into a Trk inhibitor therapy, noninvasive tools are needed to determine their Trk fusion expression status. Fluorescence in situ hybridization (FISH), next sequencing generation (NSG), standard RT-PCR, or immunohistochemistry analysis using tumor biopsy specimen collected invasively are known [28–36].

There is a need to develop noninvasive methods to reduce the invasiveness of diagnostic procedures. Radiolabeled probes for positron emission tomography (PET) represent a sensitive noninvasive approach. Over the past years, several 11C- and 18F-labeled kinase domain-binding Trk inhibitors derived from pan-Trk and TrkA subtype selective lead structures were developed [37]. A first-generation radioligand based on the 4-aza-2-oxindole structure resulted in 11C-labeled GW441756 [38]. Due to its isomeric mixture, and observed metabolism and pulmonary retention, it was not further developed [37]. Subsequent radioligands in the first generation had linear elongated structures for better cell membrane penetration. 18F-labeled GW2580 and QM1CF were kinome selective pan-Trk inhibitors [39, 40]. A second radiotracer generation was developed based on imidazo[1,2-b]pyridazine- and pyrazolo[1,5-a]pyrimidine-containing structures resulting in 13C-labeled IPMICF16 [41]. Compared to the first-generation compounds pure enantiomer (R)-IPMICF16 resulted in higher potencies (IC50) against Trk: 0.2 nM (TrkB), 0.1 nM (TrkC), 4.0 nM (TrkA) [42]. In vivo evaluation in mice, rats, non-human primates and ‘first-in-human’ showed moderate brain uptake. [11C]-(R)-IPMICF16 is a substrate for active P-glycoprotein 1 efflux [42]. Non-human and human PET data revealed no efflux from the brain and enriched accumulation in thalamus > cerebellum and cortex.

To overcome disadvantages of 11C (half-life 20 min), 18F-labeled pan-Trk inhibitor TRACK was developed representing an 18F-labeled analogue of (R)-IPMICF16 [43]. In vitro potency toward all three Trk subtypes was similar to [11C]-(R)-IPMICF16. Non-human and ‘first-in-human’ PET imaging revealed slightly higher uptake in the cerebellum, thalamus, and cortex [44].

Based on the complex transmembrane protein structure of Trk, its activation and dimerization status as NTRK fusion protein, and the localization of the intracellular ligand-binding sites, it remains challenging to demonstrate ‘proof-of-target’ binding for these novel radiolabeled Trk inhibitors in vivo. The goal of the present study was to analyze direct interaction of [18F] TRACK with peripheral Trk proteins in vivo to prove its specificity for its use as a functional PET imaging probe.

Methods

General

All chemicals, reagents, and solvents for synthesis and analysis were analytical grade. If not stated otherwise, all chemicals and reagents were obtained from Sigma-Aldrich (Sigma-Aldrich, Oakville, ON, Canada).

Radiosynthesis

The radiosynthesis of [18F]TRACK was performed similarly as previously reported [43, 44] utilizing alcohol-enhanced copper-mediated radiofluorination [45]. In brief, no carrier-added aqueous [18F]fluoride was produced by an 18O(p,n)18F nuclear reaction through the bombardment of an 18O(H2O target. Aqueous [18F]fluoride was passed through the female end of a QMA Sep Pak Light Carbonate (46 mg) cartridge (1.6–1.9 GBq in 1 mL H2O). The trapped [18F]fluoride was pre-dried by passing n-butanol (3 mL) through the cartridge, followed by air (10 mL). [18F]fluoride was eluted from the cartridge directly into a glass V-vial containing the boronic ester precursor (4.8 mg. 9.1 μmol) by passing a solution
of Et$_2$NHCO$_3$ (3.5–4.0 mg) in n-butanol (0.4 mL) through
the male end of the cartridge, with a routine elution effi-
ciency of 75–80%. The reaction vial, with no vial cap,
was placed in a 110 °C oil bath for 20 min. The vial was
removed from the oil bath and the reaction solution
diluted with HPLC eluent (0.5 mL) and drawn up in
a syringe. The reaction solution was injected on HPLC (elu-
ent (isocratic): 55% MeCN in H$_2$O; flow rate: 3 mL/min,
injection loop: 1 mL, column: Phenomenex Luna 3 μm
PFP(2) 100 Å, 250 × 10 mm). The peak eluting at ~ 21 min
was collected and diluted with H$_2$O (15 mL) and passed
through a C18 Sep Pak Light cartridge, followed by H$_2$O
(5 mL) and air (10 mL). [$^{18}$F]TRACK was eluted from
the cartridge with ethanol (10 drops). The volume was
reduced to ~ 20 μL under an N$_2$ stream at 90 °C to afford
112–140 MBq of [$^{18}$F]TRACK for injection.

Cell culture
Human colon adenocarcinoma KM12 cells were obtained
from Charles River Laboratories, National Cancer Insti-
tute, Frederick National Laboratory Cancer Research
(Fort Detrick, Frederick, MD, USA.), and cultivated in
RPMI 1640 cell growth medium supplemented with 10%
fetal bovine serum (FBS) (Gibco) and 2 mM L-glutamine
(Sigma-Aldrich). For the in vitro cell experiments, cells
were seeded in 12-well plates in their medium and grown
for 24 h.

In vitro cell uptake experiments
Right before the experiment, the media was removed and
the cells were washed two times with phosphate-buffered
saline solution (PBS). Next, 200 μL Krebs–Ringer buffer
(120 mM NaCl, 4 mM KCl, 1.2 mM KH$_2$PO$_4$, 2.5 mM
MgSO$_4$, 25 mM NaHCO$_3$, 70 μM CaCl$_2$, 5 mM glucose
pH 7.4) was added to each well. Next 300 μL Krebs–
Ringer buffer with 0.1–0.5 MBq [$^{18}$F]TRACK radiotracer
was added to each well, and the plates were incubated at
37 °C for specific periods of time (1, 5, 10, 15, 30, 45, and
60 min). Radiotracer uptake was stopped with 1 mL ice-
cold PBS, and the cells were washed twice with PBS and
lysed in 0.4 mL radioimmunoprecipitation assay buffer
(RIPA buffer). Radioactivity in the cell lysates was mea-
sured as counts per minute [CPM] using a WIZARD2
Automatic gamma counter (PerkinElmer; Waltham, MA,
USA) and converted to the radioactivity dose SI unit
Becquerel [Bq]. Total protein concentration in the sam-
ple was determined by the bicinchoninic acid method
(BCA; Pierce, Thermo Scientific 23,227) using bovine
serum albumin (BSA) as protein standard. Data were
normalized as percent of measured radioactivity per mg
protein (%radioactivity / mg protein). Graphs displaying
cell uptake over time were constructed using GraphPad
Prism 5.0 (GraphPad Software, San Diego, Ca, USA).

Competition binding experiments were carried out in
the presence of increasing concentrations of reference
compound TRACK ($10^{-8}$ – $10^{-7}$ M) or pan-Trk inhibitor
entrectinib ($10^{-8}$ – $10^{-6}$ M; Tocris Biosciences, Bristol,
UK). Blocking compounds were added to the cells imme-
diately before radiotracer [$^{18}$F]TRACK was applied and
incubated for 60 min. After incubation cells were rinsed
with ice-cold PBS, lysed, and counted as described above
to obtain CPM values. Data then were normalized as %
maximum of total added radioactivity from control wells
not receiving any blocker compound. Competition bind-
ing curves were also generated using GraphPad Prism 5.0
(GraphPad Software, San Diego, Ca, USA).

Animal experiments
Female athymic nude mice (10–12 weeks old) were
received from Charles River Laboratories (Saint-Con-
stant, QC, Canada) and injected subcutaneously with
3 × 10$^6$ KM12 cells in 100 μL PBS/Matrigel (50:50) into
the upper left flank. After about 3 to 4 weeks, subcuta-
eous KM12 tumors reached sizes of ~400–500 mm$^3$
suitable for PET imaging experiments. Female B6129SF2/J
(wild-type control) and female B6129S2-Ntrk2$^{tm1Bbd}$/J
(TrkB 50% knockout) mice were obtained from Jackson
Laboratories (JAX™ Bar Harbor, ME, USA) at the age of
10–12 weeks and used over the next 3–5 months.

Dynamic PET imaging
General anesthesia of wild-type control or tumor-bearing
mice was induced with inhalation of isoflurane in 40%
oxygen/60% nitrogen (gas flow = 1 mL/min), and mice
were subsequently fixed in prone position. The body tem-
perature was kept constant at 37 °C for the entire exper-
iment. Mice were positioned in a prone position into the
center of the field of view. A transmission scan for atten-
uation correction was not acquired. Radioactivity present
in the injection solution (0.5 mL syringe) was determined
using dose calibrator (AtomlabTM 300, Biodex Medi-
cal Systems, New York, USA). After emission scan was
started, radioactivity (3–8 MBq in 100–150μL saline) was
injected with a delay of ~ 15 s through a tail vein catheter.
For blocking experiments, 5 mg/kg entrectinib (pan-Trk
inhibitor) was pre-injected 15 min before the radiotracer
or 0.1–0.5 mg/kg unlabeled reference compound TRACK
was co-injected i.v. together with the radiotracer. Pre-
dosing with 15 mg/kg amitriptyline was done i.p. 4 h
before the radiotracer was injected. PET data acqui-
sition was performed in 3D list mode for 60 min. Dynamic
list mode data were sorted into sinograms with 54 time
frames (10 × 2 s, 8 × 5 s, 6 × 10 s, 6 × 20 s, 8 × 60 s, 10 × 120 s, 5 × 300 s). Frames were reconstructed using
ordered subset expectation maximization (OSEM) or
maximum a posteriori (MAP) reconstruction mode. No
correction for partial volume effects was performed. Image files were further processed using the ROVER v2.0.51 software (ABX GmbH, Radeberg, Germany). Masks defining 3D regions of interest (ROI) were set and defined by 50% thresholding. Mean standardized uptake values \[\text{SUV}_{\text{mean}} = \frac{\text{activity/mL tissue}}{\text{injected activity/body weight}}, \text{mL/kg}\] were calculated for each ROI. Time–activity curves (TAC) were generated from the dynamic scans.

Western blotting
Tissue samples were homogenized on ice using a sonicator using RIPA buffer with Triton-X-100 (Biopower, Markham, ON, Canada). The protein concentrations were determined using Pierce™ BCA Protein Assay Kit (Thermo Scientific, Waltham, MA, USA). Samples were loaded onto a 4–12% precast polyacrylamide gel (Bio-Rad, Hercules, CA, USA), separated by SDS-PAGE, and transferred to PVDF membranes. The membranes were first air-dried for 1 h and re-hydrated with 100% methanol. The membranes were blocked with 5% non-fat powdered milk and then incubated with a combination of either rabbit monoclonal anti-TrkA antibody (ab76291; Abcam, Cambridge, UK; 1:1000) and mouse monoclonal anti-TrkB [MM0586-7Y6] (ab89925; Abcam, Cambridge, UK; 1:1000) or rabbit polyclonal anti-TrkB (ab18987; Abcam, Cambridge, UK; 1:10,000) and mouse monoclonal anti-TrkA antibody (ab86474; Abcam, Cambridge, UK; 1:1000) overnight at 4 °C. The membranes were incubated with 3% H2O2 in water for 15 min, slides were incubated with DakoCytomation Envision + anti-mouse- or anti-rabbit-labeled Polymer HRP (DakoCytomation, Glostrup, Denmark) for 1 h, developed, using Dako Liquid DAB+Substrate Chromagen System and 1% copper sulfate and counterstained with hematoxylin. Slides were dehydrated by reversing re-hydration and cover slipped. Quantification analysis of protein expression was conducted with ImageJ 1.x (NIH and LOCI, University of Wisconsin; USA) and IHC profiler [46].

Data analysis
All data are expressed as means± standard error of the mean (SEM) from three or more experiments. All graphs were constructed using GraphPad Prism 5.0 (GraphPad Software, San Diego, CA, USA). IC50 values were calculated using a nonlinear regression (curve fit) analysis (GraphPad Prism 5.0). Statistical differences were considered significant if p < 0.05 and were evaluated using the Student’s t test.

Results
Figure 1 shows the chemical structure of [18F]TRACK in comparison with the [11C]-labeled derivative. Radiolabeling occurred at the same molecule position.

In vitro cell uptake of [18F]TRACK was analyzed in human KM12 colon cancer cells. These cells express TrkA, specifically tropomyosin 3-TrkA fusion protein (Vaishnavi et al. 2013; Tatematsu et al. 2014). Uptake of [18F]TRACK was increasing over time reaching 208 ± 72% radioactivity per mg protein (n = 6/2) after 60 min incubation time (Fig. 2). Blocking of TrkA was measured through competitive binding using increasing concentrations of pan-Trk inhibitor entrectinib and compared to the effects of the non-radioactive reference compound TRACK. Similar half-maximum
Inhibition concentrations (IC50) values were determined: 29.4 ± 9.4 nM for TRACK and 30.9 ± 3.6 nM for entrectinib (both n = 6/2), respectively, confirming that radiotracer [18F]TRACK was indeed binding to TrkA in vitro and that both compounds, TRACK and entrectinib, functionally blocked TrkA in the analyzed KM12 cells.

The radiolabeled [18F]TRACK was also analyzed in vivo in KM12 tumor-bearing mice. Original PET images as well as semiquantitative radioactivity uptake into tumor and muscle tissue are depicted in Fig. 3 (brain uptake in WT B6129SF2/J and B6129S2-Ntrk2tm1Bbd/J TrkB 50% knockout mice in Additional file 1: Fig. S1). The images show brain uptake and washout over time as well as increasing signal in BAT. The tumor area is visible at 60 and 120 min post-injection, while clearance mainly occurs through the hepatobiliary system as the liver and intestines show high radioactivity levels (Fig. 3A). Tumor uptake was continuously increasing over time from a standardized uptake value (SUV) of 0.22 ± 0.03 at 10 min to 0.43 ± 0.03 (n = 6) after 60 min post-injection (Fig. 3B). In contrast, radioactivity in muscle tissue uptake was initially systematically higher with a SUV of 0.30 ± 0.03 at 10 min increasing to 0.46 ± 0.03 (n = 6) at 40 min post-injection with no further change up to 60 min post-injection indicative of a beginning washout of the radioactivity. To analyze this further, two KM12 tumor-bearing mice were measured dynamically over 120 min post-injection. While the SUV in the tumor tissue reached 0.53 (n = 2) at 120 min in these mice, SUV in muscle tissue amounted to 0.45 after that time frame (n = 2; Fig. 3D). Taken together, radioactivity washout from muscle started after ~90 min post-injection, while tumor uptake was still increasing. This observation led to the conclusion that accumulating continuous tumor uptake was only occurring after later time points, while uptake into muscle tissue later also resulted in a slow washout process. After 120 min post-injection, tumor-to-muscle ratio (T/M) reached 1.13, while at 60 min it amounted to only 0.9 (Fig. 3C).

Based on the relatively low radioactivity uptake in the peripheral tumors, the next experimental step involved the protein analysis of TrkA and TrkB in brain, BAT, and KM12 tumor tissue of wild type (WT), TrkB 50% KO (B6129SF2/J), and KM12 tumor-bearing athymic mice (Fig. 4). In the whole brain, both TrkA and TrkB proteins were found to be sufficiently expressed with TrkB about twofold–threefold higher than TrkA (Fig. 4B). Expression of TrkB was similar in KM12 tumor tissue and brown adipose tissue (BAT), and 50% knockout of TrkB confirmed reduced protein expression in whole brain and BAT (Fig. 4C). A similar pattern was observed in white adipose tissue (AT) as well (Fig. 4D).

Immunohistochemistry analysis also revealed TrkB protein expression in the membrane of fat cells in brown adipose tissue which was higher than TrkA expression (Fig. 5) as quantified protein positive staining showed: 11.2 ± 1.1% for TrkA versus 20.0 ± 4.7% for TrkB (n = 3; p = 0.0974; Fig. 5D), respectively. Using the same antibody, no TrkB protein expression was detectable in KM12 tumor tissue as well as TrkA. In the latter sample, only some inclusions into the tumor tissue detected some protein levels (Fig. 5A). For comparison,
protein expression of TrkA and TrkB in the brain of KM12 tumor-bearing athymic mice and B6129SF2/J wild-type mice is shown in Additional file 1: Fig. S2.

Analysis of dynamic PET data after injection of $[^{18}F]$TRACK resulted in substantially higher radioactivity uptake into brown adipose tissue compared to KM12 tumor tissue before. In KM12 tumor-bearing athymic mice, uptake into BAT resulted in $\text{SUV}_{60\text{min}}$ $1.32 \pm 0.08$ ($n=7$), while in WT mice uptake amounted to $1.00 \pm 0.03$ ($n=3$), respectively (Fig. 6). From the dynamic time–activity curves, it was also visible that the uptake levels reached the maximum earlier after $\sim 60$ min compared to the KM12 tumor uptake ($>2$ h). Interestingly, 50% knockout of TrkB in B6129S2-NTkr²tm1Bbd/J mice did not lead
Fig. 4  Protein expression of TrkA and TrkB. A Original gel of selected samples from KM12 tumor tissue, brown adipose tissue (BAT), white adipose tissue (AT), and the whole brain from KM12 tumor-bearing mice, control wild type (WT), and TrkB 50% KO mice from combined red (700 nm) and green (800 nm) light wavelength analysis showing using antibodies for TrkA and TrkB and β-actin as housekeeping gene. For the full-length gel please, see Additional file 1: Fig. S5. B Semiquantitative comparison of TrkA and TrkB protein levels in brain samples from KM12 tumor-bearing mice, control wild type (WT), and TrkB 50% KO mice. C Semiquantitative comparison of TrkB protein levels in BAT and KM12 tumor tissue from KM12 tumor-bearing mice, control wild type (WT), and TrkB 50% KO mice. D Semiquantitative comparison of TrkB protein levels in white adipose tissue (AT) from KM12 tumor-bearing mice, control wild type (WT), and TrkB 50% KO mice. B–D All data are shown as normalized ratios (Trk / β-actin) and as mean ± SEM from 5 different tissue samples. * p < 0.05; n.s.—not significant
Fig. 5  Protein expression of TrkA and TrkB. Immunohistochemical staining of TrkA and TrkB in KM12 tumor tissue samples (A) and in BAT of wild-type (WT) control mice (B, C). Pictures were taken using a 20 x objective. D Quantification of positive TrkA and TrkB immunohistochemical staining in BAT. Data as mean ± SEM from n = 3 slices.
to a reduced uptake of $^{[18F]}$TRACK in BAT (Additional file 1: Fig. S3).

Specificity of $^{[18F]}$TRACK and its functional binding to Trk in vivo were tested with blocking experiments. Using the non-radioactive reference compound TRACK with doses of 0.1 mg/kg and 0.5 mg/kg co-injected with the radiotracer resulted in a concentration-dependent reduction of the uptake curve in BAT with the highest effects observed at ~30 min with a reduction of SUV from $1.00 \pm 0.05$ ($n = 3$) to $0.77$ ($n = 2$) amounting to ~23%. Interestingly, this blocking effect was eliminated again at 60 min post-injection (Fig. 7B). In addition to the self-blocking experiment, pan-Trk inhibitor entrectinib was used. A dose of 5 mg/kg pre-injected i.p. 15 min before the radiotracer resulted in a similar blocking pattern: a blocking effect at ~30 min post-injection resulting in an SUV of $0.82 \pm 0.11$ ($n = 3$) corresponding to ~18% which was also completely abolished again after 60 min. This was very much like the blocking with TRACK indicating the same functional pattern.

To further prove functional involvement of Trk, the classic tricyclic antidepressant compound amitriptyline was chosen for further in vivo experiments as it was reported to act as a TrkA and TrkB agonist in the literature [47]. Pre-dosing of KM12 tumor-bearing mice with 15 mg/kg amitriptyline 4 h i.p. before radiotracer injection resulted in significant increases in absolute $^{[18F]}$TRACK uptake in KM12 tumor tissue by 23% ($SUV_{60\min}$ from $0.53 \pm 0.01$ to $0.43 \pm 0.03$; $n = 6$; $p < 0.05$) and also in brown adipose tissue by 21% ($SUV_{60\min}$ from $1.32 \pm 0.08$ ($n = 5$) to $1.59 \pm 0.07$; $n = 6$; $p < 0.05$), respectively (Fig. 8B, C). Effects in the brain were somewhat smaller amounting to an increase of 13% only which was not significant ($SUV_{60\min}$ from $0.52 \pm 0.07$ ($n = 5$) to $0.60 \pm 0.06$; $n = 6$; $p > 0.05$; Fig. 8C). Taken together, amitriptyline did indeed increase uptake of $^{[18F]}$TRACK in selected peripheral tissue, however, not significantly in the brain. In addition, $^{[18F]}$TRACK uptake into muscle tissue was also increased in the presence of amitriptyline by 15% ($SUV_{30\min}$ from $0.45 \pm 0.03$ to $0.52 \pm 0.02$; $n = 5$; $p = 0.057$; Additional file 1: Fig. S4). However, at 60 min this difference was diminished which contrasted with the profiles observed in brown adipose tissue and KM12 tumor tissue.

In a final experimental setup, effects of the pan-Trk inhibitor entrectinib were also tested in KM12 tumor-bearing mice pre-treated with 15 mg/kg amitriptyline before to analyze if there would be still a functional blocking effect in peripheral tissue. Indeed, dynamic PET experiments revealed that in KM12 tissue, the addition
of 5 mg/kg of entrectinib reduced $[^{18}\text{F}]$TRACK uptake slightly by ~8% from SUV$_{60\text{min}}$ from 0.53 ± 0.01 ($n = 5$) to 0.48 ± 0.05 ($n = 3$; $p > 0.05$; Fig. 9A) and in brown adipose tissue by ~14% from SUV$_{60\text{min}}$ from 1.56 ± 0.08 ($n = 5$) to 1.37 ± 0.12 ($n = 3$; $p > 0.05$; Fig. 9B), respectively. Although these effects were small and did not satisfy the criteria for statistical significance, they fit the observable pattern.

**Discussion**

The present study revealed the following main results: (i) Trk inhibitor $[^{18}\text{F}]$TRACK is taken up into TrkA expressing human KM12 colon cancer cells, and this uptake can be inhibited by pan-Trk inhibitor entrectinib; (ii) $[^{18}\text{F}]$TRACK is also taken up into KM12 tumor tissue as well as brown adipose tissue (BAT) in vivo which can also be blocked by entrectinib; and (iii) amitriptyline acts as a functional Trk agonist and increases uptake of $[^{18}\text{F}]$TRACK in peripheral target tissue.
Trk expression and function have been mainly studied in the neuronal system as well as for neurological disorders and neurodegenerative diseases [15–19]. In addition, Trk fusion protein expression and activation promoting cell proliferation have been found in many cancers [1]. However, based on their rarity, variety and clinical sample detection limits to date, functional effects, and importance in cancers are not fully understood yet [48]. In 2018 and 2019, first-generation Trk inhibitors, larotrectinib and entrectinib, were introduced into the clinic for targeted therapy effectively blocking the kinase domain activity [49]. Next-generation Trk inhibitors, selitrectinib and repotrectinib, are currently undergoing clinical development to overcome therapy resistance [50]. Clinical data have shown that specific therapy with first-generation Trk inhibitors achieves high response rates independent of tumor histology, age, or NTRK fusion type [51]. Patient selection for Trk therapy may become a key step for personalized medicine and treatment decisions. As current clinical testing for NTRKs is still based on invasive biopsy sampling or costly novel bioassays such as NGS [48], access to noninvasive imaging methods such as PET would tremendously facilitate therapy decision-making processes in the clinical setting. Even with the limitation of Trk protein expression in muscle tissue [52] which would lead to a certain increased background uptake, overexpression of Trk fusion proteins in certain cancer types could contribute to valuable diagnostic PET information for a specific patient population to support the therapeutic decision and to select only eligible patients for a Trk inhibitor therapy.

After numerous imaging probe developments [37], [18F]TRACK has been identified as an optimized 18F-labeled Trk targeting PET radiotracer [43, 44]. So far, [18F]TRACK has been mainly evaluated for its binding to Trk in the brain. While it has shown substantial brain uptake in preclinical studies in murine, monkey, and human brain [43, 44], in vivo proof of target specificity remains challenging.

In the present study, inhibition of [18F]TRACK cellular uptake into KM12 cells using pan-Trk inhibitor entrectinib was successful. KM12 represents a human colon cancer cell line expressing TrkA and more specifically tropomyosin 3-TrkA (TPM3-NTRK1) fusion protein [14, 53], since non-radiolabeled TRACK and entrectinib resulted in similar IC50 values in the nanomolar range when competing with [18F]TRACK binding in KM12 cells. These results serve as a strong functional proof of target in this colon cancer model in vitro. In enzymatic assays, entrectinib has shown IC50 values of 1–5 nM against isolated TrkA/B/C [54]. TRACK resulted in an IC50 of 0.3–4.2 nM IC50 for human TrkA/B/C measured in an ATP-based enzymatic assay [43]. However, uptake into KM12 tumors in vivo was like muscle tissue (SUV60min 0.43 vs. 0.46). This could be attributed to the muscle expression of TRACK specifically TrkB as known from the literature [52, 55]. Interestingly, uptake profiles over time resulted in a later slight washout from muscle tissue, while uptake into tumor tissue continued to rise even after 2 h. resulting in a T/M ratio of 1.1.

PET imaging experiments also revealed uptake into brown adipose tissue (BAT) in mice reaching significantly higher uptake levels (SUV60min > 1) versus KM12 tumor tissue. In the literature, adipose tissue has been shown to express TrkB mRNA [56] and TrkB protein in bovine adipose tissue of lactating cows [57]. The neuronal BDNF/TrkB axis and its function in adipose tissue play an important role for metabolic regulation and its influence on food intake and obesity. Therefore, it seems not surprising to detect uptake of [18F]TRACK in murine BAT.

TrkB protein expression in brown and white adipose tissue as well as KM12 tumor samples was confirmed using Western blotting and immunohistochemistry and compared to TrkB expression in the whole mouse brain. Experimental analysis revealed that TrkB expression in the brain is high and ubiquitous. Immunohistochemistry also showed somewhat higher expression of TrkA over TrkB in the mouse brain. In contrast, expression of TrkB was stronger than TrkA on the membrane surface of fat cells analyzed from murine BAT samples confirming previous observations in lactating cows [57]. However, present analysis of KM12 tissue with immunohistochemistry did not reveal substantial expression of both TrkA and TrkB. TrkA was only found in small inclusions in the tumor tissue confirming only a somewhat weak expression although KM12 cells are known to strongly express TrkA [58] although as the fusion protein TPM3-NTRK1
Blocking of $[^{18}\text{F}]$TRACK uptake after stimulation with amitriptyline

**A) KM12 tumors**

- + 15 mg/kg amitriptyline i.p.
- + 15 mg/kg amitriptyline i.p. + 5 mg/kg entrectinib i.p.

**B) Brown Adipose Tissue (BAT)**

- $[^{18}\text{F}]$TRACK
- $[^{18}\text{F}]$TRACK + amitriptyline
- $[^{18}\text{F}]$TRACK + amitriptyline + entrectinib

**Fig. 9** (See legend on previous page.)
It is possible that the antibody used for western blot and immunohistochemistry experiments in the present study was not able to detect the fusion protein.

The present functional in vivo analysis using radiotracer $^{18}$FTRACK supports the fact of lower protein expression and functional availability in KM12 tumors versus higher expression in BAT. To confirm specific targeting of $^{18}$FTRACK to Trk (mainly TrkB) in BAT, in vivo blocking studies were performed with non-radioactive TRACK reference compound for self-blocking and pan-Trk inhibitor entrectinib. With both compounds, similar blocking effects on the radiotracer uptake were observed at 30 min p.i. (minus 23% with TRACK and 18% with entrectinib). It must be added that intracellular targets are often difficult to engage experimentally with in vivo blocking PET experiments. This is further based on the necessary higher lipophilicity of the PET tracer compound to reach its intracellular target. We have made similar observations before when analyzing specific binding of radiotracers to intracellular COX-2. While inhibition with selective COX-2 inhibitor, celecoxib, revealed only 24% blocking effect from PET experiments using $^{18}$FPyricoxib, ex vivo biodistribution showed a 50% inhibition [60]. Based on that, blocking effects with TRACK and entrectinib were observed within the same order of magnitude. However, this effect was not persistent and at 60 min p.i., no blocking was detectible indicating reversible effects over time. Based on the reversibility of the observed effects, it is difficult to conclude that these in vivo experiments using standard Trk inhibitors and therefore antagonists binding to the intracellular kinase domain would serve as proof of target in vivo although they show clear blocking effects in KM12 cells in vitro.

Consequently, an alternative approach using an agonist on Trk receptors might provide a way to prove specific targeting of TrkB via radiotracer $^{18}$FTRACK in vivo. A basic analysis from 2009 revealed that the classic tricyclic antidepressant amitriptyline directly binds to the extracellular domain of both TrkA and TrkB triggering their dimerization and activation [47]. It was found that amitriptyline is in fact acting as an agonist on both TrkA and B in neuronal cells and that inhibition of TrkB eliminated its neuroprotective effect without impairing its antidepressant activity. Subsequent follow-up studies found that amitriptyline induces neurite outgrowth in rat primary cortical neurons through Trk receptor and MAP kinase activation [61] and that similarly to neurotrophins amitriptyline also acts as a Trk receptor agonist on the regeneration of afferent cochlear synapses in the inner ear [62]. Using amitriptyline in the present study, overall uptake levels of $^{18}$FTRACK in KM12 tumor tissue and in BAT could be significantly increased by 21 and 23% indicating that ligand activation of TrkA and TrkB in both tissues resulted in increased binding of $^{18}$FTRACK at the intracellular kinase binding site as well. For comparison, radiotracer uptake in the whole brain was increased by only 13%, which did not reach statistical significance, though the trend confirmed the observations in the peripheral tumor and BAT tissue. Interestingly, $^{18}$FTRACK uptake into muscle tissue was also elevated by 15%, however, only after 30 min and diminished after 60 min which is different from the effect in tumor and BAT tissue attributing maybe to different intracellular ATP levels to compete for binding with $^{18}$FTRACK or different dimerization forms of the oncogene Trk versus the muscular Trk. It is known that muscle tissue expresses Trk [55] and specifically TrkB and that BDNF mimetics such as 7,8-di-hydroxyflavone [52] or other antidepressant drugs such as fluoxetine can activate muscular TrkB by direct binding to the receptor [63]. Taken together, tricyclic antidepressant amitriptyline also acts as a Trk agonist in peripheral target tissue by activating Trk receptors resulting in an uptake increase in Trk targeting radiotracer $^{18}$FTRACK. In addition, this in vivo experiment also serves as a pharmacodynamic proof-of-target experiment.

Finally, experiments combining agonistic activation of Trk with amitriptyline (binding at the extracellular domain) and Trk inhibition with entrectinib (binding the intracellular kinase domain) revealed that there was a small but also systematic blocking effect detectable at 60 min post-injection of radiotracer $^{18}$FTRACK in both KM12 tumor tissue (Δ8%) and BAT (Δ14%). This effect was persistent in the equilibrium phase of the radiotracer’s in vivo distribution which contrasted with the observed reversible effects detected without amitriptyline. This combination experiment may serve as the best functional proof of target for selective target binding of $^{18}$FTRACK in vivo and in combination with the in vitro results.

**Conclusion**

The present study demonstrated that novel Trk targeting PET radiotracer $^{18}$FTRACK binds to TrkA and TrkB in peripheral tissue such as colon KM12 cancer and adipose tissue but also muscle tissue. Besides its use for noninvasive functional brain PET imaging to detect changes in neurodegenerative diseases as analyzed before, the present data also support the potential application as an additional tool for multiple diagnostic tumor imaging to detect the right patients with Trk protein expression in cancers and to select them for specific Trk inhibitor therapy. Additionally, visualization of TrkB in adipose tissue of obese patients may open a new field for imaging Trk involvement in in metabolic disease. While first-in-human PET has been
recently performed with $[^{18}F]$TRACK for brain PET imaging and current clinical studies are undergoing in Alzheimer disease patients, the present dataset may also support a clinical translation for cancer and metabolic disease as well.

Abbreviations

BAT: Brown adipose tissue; BDNF: Brain-derived neurotrophic factor; FISH: Fluorescence in situ hybridization; MAP: Maximum a posteriori; NGF: Nerve growth factor; NGS: Next-generation sequencing; OSEM: Ordered subset expectation maximization; PET: Positron emission tomography; ROI: Region of interest; SUV$_{max}$: Mean standardized uptake value; TAC: Time–activity curve; Trk: Tropomyosin receptor kinase; T/M: Tumor-to-muscle ratio; WB: Western Blot.

Supplementary Information

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Additional file 1: Fig. S1: Brain uptake of $[^{18}F]$TRACK in control and 50% TrkB knockout mice from dynamic PET experiments. Representative images at 30 min post injection and time-activity curves over 120 min time course. Fig. S2 TrkA and TrkB protein expression in mouse brain. Immunohistochemical staining in slices from a KM12 tumor bearing athymic mouse and a control wild-type B6129SF2/J mouse. Fig. S3: Uptake of $[^{18}F]$TRACK into brown adipose tissue (BAT). Time activity curves for BAT tissue in control wild-type B6129SF2/J and B6129SF2-Ntrk2tm1Bbd/J 50% TrkB knockout mice over 120 min time course. Fig. S4: Muscle uptake of $[^{18}F]$TRACK in KM12 tumor bearing mice. Effect of 15 mg/kg amitriptyline on the time-activity curve over 60 min time course. Fig. S5: Original full gel for the protein expression of TrkA and TrkB. Data are shown in samples from KM12 tumor tissue, brown adipose tissue (BAT), white adipose tissue (AT) and the whole brain from KM12-tumor bearing mice, control wild-type B6129SF2/J mice and B6129SF2-Ntrk2tm1Bbd/J 50% TrkB knockout mice.

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Author contributions

MW, JJB, and RS contributed to conception and design; MW and RS were involved in development of methodology; MW, JJB, JD, DG, and VO contributed to acquisition of data; MW, JJB, DG, THJ, JMB, and RS were involved in analysis and interpretation of data; and MW, JJB, THJ, JMB, and RS contributed to writing, review, and/or revision of the manuscript. All authors have read and approved the final manuscript.

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Availability of data and materials

Experimental data are available from the corresponding authors on reasonable request.

Declarations

Ethics approval

All animal experiments were carried out in accordance with the ARRIVE guidelines as well as the guidelines of the Canadian Council on Animal Care and approved by the local animal care committee (Cross Cancer Institute, University of Alberta, protocol number AC14214).

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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