Type 1 cannabinoid receptor mapping with $[^{18}F]$MK-9470 PET in the rat brain after quinolinic acid lesion: a comparison to dopamine receptors and glucose metabolism

Cindy Casteels · Emili Martinez · Guy Bormans · Lluïsa Camon · Nuría de Vera · Veerle Baekelandt · Anna M. Planas · Koen Van Laere

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Abstract

Purpose Several lines of evidence imply early alterations in metabolic, dopaminergic and endocannabinoid neurotransmission in Huntington’s disease (HD). Using $[^{18}F]$MK-9470 and small animal PET, we investigated cerebral changes in type 1 cannabinoid (CB$_1$) receptor binding in the quinolinic acid (QA) rat model of HD in relation to glucose metabolism, dopamine D$_2$ receptor availability and amphetamine-induced turning behaviour.

Methods Twenty-one Wistar rats (11 QA and 10 shams) were investigated. Small animal PET acquisitions were conducted on a Focus 220 with approximately 18 MBq of $[^{18}F]$MK-9470, $[^{18}F]$FDG and $[^{11}C]$raclopride. Relative glucose metabolism and parametric CB$_1$ receptor and D$_2$ binding images were anatomically standardized to Paxinos space and analysed voxel-wise using Statistical Parametric Mapping (SPM2).

Results In the QA model, $[^{18}F]$MK-9470 uptake, glucose metabolism and D$_2$ receptor binding were reduced in the ipsilateral caudate-putamen by 7, 35 and 77%, respectively ($p<2 \times 10^{-5}$), while an increase for these markers was observed on the contralateral side (>5%, all $p<7 \times 10^{-4}$). $[^{18}F]$MK-9470 binding was also increased in the cerebellum ($p=2 \times 10^{-5}$), where it was inversely correlated to the number of ipsiversive turnings ($p=7 \times 10^{-6}$), suggesting that CB$_1$ receptor upregulation in the cerebellum is related to a better functional outcome. Additionally, glucose metabolism was relatively increased in the contralateral hippocampus, thalamus and sensorimotor cortex ($p=1 \times 10^{-6}$).

Conclusion These data point to in vivo changes in endocannabinoid transmission, specifically for CB$_1$ receptors in the QA model, with involvement of the caudate-putamen, but also distant regions of the motor circuitry, including the cerebellum. These data also indicate the occurrence of functional plasticity on metabolism, D$_2$ and CB$_1$ neurotransmission in the contralateral hemisphere.

Keywords Type 1 cannabinoid receptor · Small animal PET · QA · Huntington’s disease · $[^{18}F]$MK-9470

Introduction

Huntington’s disease (HD) is a devastating genetic neurodegenerative disorder, clinically characterized by involuntary movements, emotional disturbances and cognitive impairments. HD is caused by a CAG repeat expansion within exon 1 of the HD gene on chromosome 4 [1]. The mutated HD gene encodes an extended polyglutamine...
stretch in the N-terminal domain of the huntingtin (htt) protein, which results in widespread neuronal degeneration preferentially within the striatum [2]. Despite the discovery of the genetic mutation associated with HD, the function of the abnormal gene product and the pathogenic mechanisms of the disease are still unknown.

Dysregulation of cannabinoid-mediated control of striatal function might play a critical role in the development of HD symptoms [3]. One of the earliest neurochemical alterations observed in HD patients is the loss of type 1 cannabinoid (CB₁) receptor binding in the basal ganglia, an alteration that significantly precedes the development of identifiable striatal neuropathology [4]. Likewise, CB₁ receptor mRNA levels were decreased in the absence of neuronal loss in the lateral striatum, cortex and hippocampus of transgenic mouse models of HD [5-9]. In the HD94 transgenic mice also decreases in the number of basal ganglia-specific binding sites and the activation of guanosine triphosphate (GTP)-binding proteins by CB₁ receptor agonists were noticed [5]. Loss of CB₁ receptors in the basal gangia not only occurred in transgenic mice HD models, but also in rats after local intrastriatal application of 3-nitropropionic acid (3-NP), a toxin that reproduces the mitochondrial complex II deficiency characteristic of HD patients [10]. Delaying the onset of HD symptoms by enriched environments has been shown to selectively slow down the loss of CB₁ receptors in experimental HD [11].

In normal conditions, CB₁ receptors are abundantly distributed in different structures of the brain, controlling motor, cognition, emotional and sensory functions [12]. CB₁ receptors encompass together with a family of plant-derived, synthetic or endogenous compounds the endocannabinoid system (ECS). The ECS is a modulatory system that interacts with and regulates functioning of other neurotransmitter systems such as glutamate and γ-aminobutyric acid (GABA) [13].

So far, only in vitro and ex vivo data on the ECS exist in experimental HD. Studies of CB₁ receptor changes in experimental HD have focused on the basal ganglia, hippocampus and motor cortex, but have not assessed other brain regions [7, 9]. Also unknown is whether CB₁ receptor levels are changed in vivo. In vivo imaging of CB₁ receptors in the rat brain has recently become feasible due to the development of a CB₁-selective radioligand, [¹⁸F]MK-9470 [14, 15].

A well-characterized and frequently used rat model of HD relies on the use of quinolinic acid (QA). QA is an endogenous metabolite in the brain that results in the degeneration of GABAergic striatonigral and striatopallidal projection neurons [16] with the relative [10] sparing of interneurons [17] after intrastratial injection, a cellular pathology remarkably similar to that in patients with HD.

In this study, we have characterized the QA lesion rat model of HD in vivo with respect to CB₁ receptor binding using [¹⁸F]MK-9470 and small animal positron emission tomography (PET) imaging. Also, dopamine D₂ receptor availability and glucose metabolism, all shown to be altered (early) in patients and animal models of HD [18, 19], were investigated in the same animals and correlated to the regional CB₁ receptor status.

Material and methods

Animals

Experiments were conducted on 21 female Wistar rats (Charles River, France) weighing 160–170 g at the start of the experiment. All animals were housed three to a cage, at an average room temperature of 22°C and a 12-h light/dark cycle. Food and water were given ad libitum. The research protocol was approved by the local Animal Ethics Committees of the Universities of Leuven and Barcelona and was according to European Ethics Committee guidelines (decree 86/609/EEC).

Striatal quinolinic lesion

All surgical procedures were performed under 1.5% halothane anaesthesia (4% during the induction period), at a rate of 1.5 ml/min. All rats were placed in a stereotactic head frame (David Kopf Instruments, Tujunga, CA, USA) [20], and a single unilateral hole was drilled in the skull over the left striatum using the bregma as reference. The neurotoxic and sham lesions in the left striatum were made by injecting 1 μl of QA (n=11, 240 nmol) or phosphate-buffered saline (PBS) (n=10, pH 7.4), respectively, at the following coordinates: anteroposterior +0.2 mm, lateral +2.8 mm and dorsoventral −4.5 mm [21]. The rate of infusion was 0.5 μl/min. After the injection, the needle was left in place for an additional 5 min before being slowly withdrawn from the brain.

Behavioural testing and weight

To assess toxin efficacy, the presence of QA-specific stereotypes was assessed in QA-lesioned rats after recovering using digital recording (Videotrack 2000, View Point, Lyon, France) [22]. Briefly, each animal (immediately after recovering from anaesthesia) was introduced into an individual cage (33.3×33.3 cm), in a soundproof room, and the behavioural activity was digitally recorded for a 3-h period. Digital records were analysed by a well-trained researcher evaluating the presence of QA-related stereotypes (head nodding, circling and rolling behaviour). All QA-treated rats
developed these stereotypes. Four weeks post-lesioning amphetamine-induced asymmetric rotational behaviour was monitored [23]. Amphetamine sulphate, supplied by the Ministry of Health of Spain, was injected intraperitoneally (i.p.) at a dose of 4 mg/kg free base. For each test, 5 min after amphetamine administration, the total number of complete turns clockwise and anticlockwise was counted over 30 min. The direction ipsilateral to the lesion is considered as positive.

The analysis of amphetamine tests was based on net ipsilateral turns (defined as anticlockwise turning in cases of a left-sided injection). Additionally, changes in body weight were measured before lesioning, 24 h and 4 weeks after.

Radiotracer synthesis

CB1 receptor imaging was performed in all animals (n=21) using the radioligand \[18F\]MK-9470 \( (N\)-[2-(3-cyano-phenyl)-3-(4-[18F]fluorethoxy)phenyl]-1-methyl[2-propyl]-2-(5-methyl-2-pyridyloxy)-2-methylproponamide), which is characterized by high specificity and high affinity for the CB1 receptor (rat IC50 0.9 nM) [14]. The precursor for tracer synthesis was obtained from Merck Research Labs (West Point, PA, USA) and labelling was performed using an \[18F\]ethylbromide procedure as described previously [14]. The final product was obtained after high-performance liquid chromatography separation and had a radiochemical purity >95%. Specific activity was on average 219 GBq/\( \mu \)mol (specific activity range: 53–606 GBq/\( \mu \)mol). The tracer was administered in a sterile solution of 5 mM sodium acetate buffer pH 5.5 containing 6% of ethanol [24].

Functional images of the striatal dopamine system were obtained from each surviving rat using the D2 receptor radioligand \[11C\]raclopride (n=20) [25], while glucose metabolism was assessed using \[18F\]FDG (n=20). \[11C\] Raclopride was obtained by methylating the corresponding nor-precursor with \[11C\]methyl triflate, whereas \[18F\]FDG was prepared by using an MRC Ion Beam Applications \[18F\]FDG synthesis module. Approximately 18 MBq (500 \( \mu \)Ci) of each radioligand (specific activity range: 53–760 GBq/\( \mu \)mol; injection volume: 500 \( \mu \)l) were injected into the tail vein using an infusion needle set.

Data acquisition

Small animal PET imaging was performed using a lutetium oxyorthosilicate detector-based tomograph (microPET Focus 220, Siemens Medical Solutions USA, Inc., Malvern, PA, USA), which has a transaxial resolution of 1.35 mm in full-width at half-maximum. Data were acquired in a \( 128 \times 128 \times 95 \) matrix with a pixel width of 0.475 mm and a slice thickness of 0.796 mm. The coincidence window width was set at 6 ns. Before imaging, the rats were anaesthetized with an i.p. injection of 50 mg of sodium pentobarbital (Nembutal, Ceva Sante Animale, Brussels, Belgium) per kilogram of body weight.

Imaging studies were performed on age-matched animals within 11–13 weeks post-lesioning for \[18F\] MK-9470, 16–18 weeks post-lesioning for \[18F\]FDG and 15–25 weeks post-lesioning for \[11C\]raclopride. Dynamic 60-min \[18F\]MK-9470 and \[11C\]raclopride acquisitions were started immediately after tracer injection (frame duration: 4×15 s, 4×60 s, 5×180 s, 8×300 s), while \[18F\]FDG measurements were obtained during 40 min starting 1 h after injection. Animals were scanned after overnight fasting for \[18F\]MK-9470 and \[18F\]FDG. The acquisition timing rationale and kinetics of the radioligands in rats have been described previously [24, 26].

Sinograms were reconstructed using filtered backprojection (FBP). No corrections were made for attenuation or scatter.

Small animal PET data processing and Statistical Parametric Mapping (SPM)

Parametric images based on standardized uptake values (SUV) [activity concentration (MBq/ml) × body mass/\( \text{ng} \)/injected dose (MBq)] were generated as a measure of absolute CB1 receptor binding [14, 27]. No significant differences in body weight or injected activity were present between QA- and PBS-infused rats. We investigated absolute as well as relative uptake. Relative \[18F\]MK-9470 uptake was expressed as SUV values normalized on whole-brain SUV. Parametric D2 binding index (BI) images, representing D2 receptor availability, were constructed by reference to the cerebellum using the Ichise multilinear reference tissue model 2 (MRTM2) module in PMOD [28]. Relative regional glucose metabolism was determined by count normalizing \[18F\]FDG data to the whole-brain uptake. The within-subject test-retest variability of our imaging procedures is in a magnitude consistent with human data, i.e. \( <15\% \) [15, 24, 29].

To obtain maximal use of image information without a priori knowledge, images were analysed on a voxel-by-voxel basis using SPM2. The procedure of spatial normalization and its validation have been described previously [24]. This methodology allows reporting results in coordinates directly corresponding to the Paxinos coordinate system for the rat brain.

SPM analysis was carried out using a categorical subject design (conditions: disease vs controls) on parametric \[18F\] MK-9470 and \[11C\]raclopride images, and on \[18F\]FDG data of QA-lesioned rats. Spatially normalized images were masked to remove extracerebral signals that would disrupt the global normalization. All images were smoothed with an isotropic Gaussian kernel of 1.6 mm. For analysis of absolute \[18F\]MK-9470 receptor binding, no proportional scaling was used and an analysis threshold of 0.8 of the mean
image intensity was applied. To study regional $^{[18F]}$MK-9470 uptake, proportional scaling was used. SPM analysis of absolute parametric D$_2$ binding data was performed without global normalization and with an absolute analysis threshold of $-\infty$ (i.e. whole-brain analysis), while $^{[18F]}$FDG data were proportionally scaled with an analysis threshold of 80% to exclude white matter and ventricular activity.

To minimize false-positive findings, T-map data were interrogated at a peak level of $p<0.005$ (uncorrected) and extent threshold $k_E >200$ voxels (1.6 mm$^3$), as described previously [30]. Only those clusters that were significant at the $p<0.05$ (corrected) level were considered.

In addition, a voxel-based correlation analysis between relative $^{[18F]}$MK-9470 uptake/$^{[18F]}$FDG data and the following covariates was performed within the QA-lesioned group: (1) striatal $^{[11C]}$raclopride binding to characterize D$_2$ impairment and (2) the number of net ipsiversive turnings. A correlation analysis between relative $^{[18F]}$MK-9470 uptake and $^{[18F]}$FDG was carried out voxel-by-voxel wise using the BPM toolbox for SPM2 [31]. D$_2$ values were determined by a predefined volume of interest (VOI) analysis based on definition on the Paxinos atlas [24]. For D$_2$ impairment, the striatal affected to non-affected side BI ratio was obtained. D$_2$ impairment, FDG data and relative $^{[18F]}$MK-9470 uptake were expressed as percentage of normal values.

General statistics

Conventional statistics were carried out using STATISTICA v8.0 (StatSoft, Tulsa, OK, USA). Behavioural outcomes and body weight gain were analysed using unpaired $t$ tests. Significance was defined at the 95% probability level. Data are mean ± SD.

Results

Subjects and behavioural data

PBS- and QA-infused rats showed the same pattern of body weight gain during the experiment (body weight gain after 24 h: $-2\pm1$ vs $-14\pm2$ g, $p<0.05$, and after 4 weeks: $62\pm20$ vs $65\pm1$ g, NS). No significant differences in the time of recovery from anaesthesia (5.6±1.8 vs 5.5±1.1 min) were found between PBS- and QA-infused rats. After surgery, QA-lesioned rats displayed, among others, head nodding, rolling and circling behaviour, with the net ipsiversive turnings after amphetamine administration significantly different from sham controls 4 weeks post-lesioning ($151\pm23$ vs $8\pm6$, $p<0.05$).

SPM analysis

Absolute $^{[18F]}$MK-9470 binding values were not significantly different between QA-lesioned rats and control animals. Average cross-sectional small animal PET images of absolute $^{[18F]}$MK-9470 binding in controls and HD rats are shown in Fig. 1. As can be seen from this figure, the pattern of $^{[18F]}$MK-9470 uptake observed in the rat control brain is consistent with that previously reported ex vivo

![Fig. 1 Average cross-sectional small animal PET images, coregistered to MRI (a), of $^{[18F]}$MK-9470 binding (b), glucose metabolism (c) and $^{[11C]}$raclopride binding in the rat brain of sham-operated animals (left) and QA-lesioned rats (right). Colour bars indicate SUV values for $^{[18F]}$MK-9470, relative intensities for $^{[18F]}$FDG and binding indices for the dopamine D$_2$ receptor. Intersection points of 3 planes have been set to the mid-striatal level of the left hemisphere [i.e. (x, y, z) = (3.4, −0.2, −6.0, Paxinos coordinates)]. Images are oriented in neurological convention](image-url)
[12], i.e. a fairly homogeneous and high uptake in the cortex, cerebellum and caudate-putamen.

When looking at regional changes by relative scaling to the global mean tracer binding, [18F]MK-9470 values were decreased in the ipsilateral caudate-putamen by 7% (p<2.10^{-5}). Also, glucose metabolism and D_2 receptor binding were reduced in this region by 35 and 77%, respectively (all p<2.10^{-11}). An increase for each of these markers was observed on the contralateral side (>5%, all p<7.10^{-4}). The contralateral increase in relative metabolism

![Fig. 2](image-url) Coronal brain sections showing overlays on the regions where a statistically significant increase (red) and decrease (blue) in relative [18F]MK-9470 binding (a), D_2 receptor binding (b) and relative glucose metabolism (c) were observed in QA-lesioned rats (figure given at p_{\text{height}}<0.005 uncorrected) as compared to controls. Significance is shown with a T statistic colour scale, which corresponds to the level of significance at the voxel level. The distance between the sections is 1.00 mm with the position relative to the bregma (positive values for sections anterior to the bregma) on top of the sections in the left column. Images are oriented in neurological convention. The D_2 receptor deficit is located mainly in the anterior part of the caudate-putamen, which is in correspondence to the site of injection. The contralateral cluster of relative glucose metabolism encompasses the caudate-putamen, hippocampus, thalamus and sensorimotor cortex.

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extended towards the contralateral hippocampus, thalamus and sensorimotor cortex (+5.8±1.9%, p=1.10^{-6}).

Statistical parametric maps of [18F]MK-9470, [11C] raclopride and [18F]FDG analysis are shown in Fig. 2. Average cross-sectional small animal PET images of glucose metabolism and [11C]raclopride BI in controls and QA-lesioned rats are shown in Fig. 1.

Relative [18F]MK-9470 binding was also increased in the cerebellum (+6.9±3.3%, p=2.10^{-5}), where it was inversely correlated to the number of ipsiversive turnings (r=−0.94, p=7.10^{-6}; Fig. 3a, b). No homologous regional correlations of [18F]FDG and [11C]raclopride with relative [18F]MK-9470 uptake were observed. Relative [18F]FDG values of the ipsilateral caudate-putamen were positively correlated with D2 impairment (r=0.92, p=9.10^{-5}; Fig. 3c, d). Detailed cluster peak locations and p values of SPM analyses are shown in Table 1.

**Discussion**

Neurodegenerative diseases such as HD are characterized by gradually evolving selective neuronal death. Animal models using neurotoxins that produce similar patterns of neuronal degeneration may yield important information to elucidate its aetiology and may be used to assess neuroprotective/neuromodulatory approaches. In HD, much interest has recently been focused on the ECS as alterations in CB1 receptor-mediated brain signalling might be related to hyperkinesia, typical of the earliest phase of the disease, or might be involved in the process of pathogenesis itself [32].

In this study, we have characterized CB1 receptor alterations for the first time in vivo in QA-lesioned rats of HD using [18F]MK-9470 and small animal PET. A whole-brain characterization of CB1 receptors in this model is valuable and necessary prior to starting drug treatment [9].

![Fig. 3 a, c Coronal, transverse and sagittal brain sections with overlay of the cluster with decreased relative [18F]MK-9470 binding (a) and decreased relative [18F]FDG (c), expressed as percentage of controls, in relation to rotational behaviour (net ipsiversive turnings) and D2 impairment for the QA group, respectively. Images are oriented in neurological convention. b Scatter plot of relative [18F]MK-9470 uptake at the maximal peak location of the cerebellum in relation to ipsiversive turning behaviour of QA-lesioned rats. d Scatter plot of relative [18F]FDG uptake at the maximal peak location of the lesioned caudate-putamen in relation to D2 impairment of QA-lesioned rats, expressed as striatal affected to non-affected side BI ratio.](image-url)
Compared with sham controls, QA-infused rats showed an in vivo decrease in CB1 receptor binding in the lesioned caudate-putamen. Reductions in CB1 receptor levels of this region are in line with ex vivo samples of HD models at early and late stages of the disease. Lastres-Becker et al. demonstrated, in the absence of neuronal loss, a 30% reduction in the striatal CB1 receptor density of HD94 transgenic mice using \[^{3}H\]CP55,940 autoradiography [5]. This reduction became more severe in 3-NP-treated rats when striatal degeneration was evident, presumably as a mere side effect caused by the progressive destruction of medium spiny GABAergic neurons [10]. We reported in the present study merely modest CB1 receptor loss in the QA model, despite also evidence of prominent degeneration in these animals, as reflected by D2 receptor imaging [33]. It appears from our findings as if CB1 receptors try to restore basal levels, but remain insufficient as mentioned before. In the normal caudate-putamen, CB1 receptors are expressed on presynaptic glutamatergic terminals originating in the cortex [34, 35]. In experimental and human HD, corticostriatal glutamatergic transmission is severely impaired (for review see [36]), and QA lesions result in a secondary dying back of these terminals [37]. Relative upregulation of this CB1 receptor population could conceivably represent a compensatory response, damping the excessive corticostriatal glutamatergic drive that results from QA infusion [38] and reducing the excitotoxic mechanisms underlying HD [39]. However, within the caudate-putamen, CB1 receptors are also expressed on striatal GABAergic interneurons that are labelled with parvalbumin and a few interneurons of the cholinergic population [40], from which their contribution to the observed effect cannot be excluded. Other contributing mechanisms may be changes in receptor affinity or in conformational state.

No correlation of this striatal CB1 receptor deficiency with disease severity, measured by \[^{18}F\]FDG and \[^{11}C\]raclopride, was observed. In patients and animals with HD, metabolic abnormalities and reductions in dopamine binding are well described in striatal and extrastriatal regions, progressively decreasing upon symptom severity [18, 19, 41–45]. Here, \[^{18}F\]FDG values of the caudate-putamen were higher than, but positively correlated to D2 impairment. Generally, \[^{18}F\]FDG uptake is a correlate of neuronal activity [46], but cellular metabolic decreases may also in part be compensated by \[^{18}F\] FDG uptake in reactive microglia, as has previously been shown in this model [47, 48].

Table 1  Peak locations for the clusters in the group comparison and correlation analysis (at \(p_{\text{uncorr}} \leq 0.005\) uncorrected, \(k_{E} > 200\))

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<tr>
<th>Cluster level</th>
<th>Voxel level</th>
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<td>2523</td>
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<td>Correlation analysis: negative correlation with net ipsiversive turnings</td>
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sphere with regard to D₂/CB₁ receptors and glucose metabolism has not been demonstrated before in unilateral HD models by histology and imaging techniques. Previous animal experiments made use of between-hemisphere comparisons or only focused on the lesioned side in comparison to control data [19, 48]. Notably, studies with unilateral intranigral 6-hydroxydopamine (6-OHDA) application to model Parkinson’s disease (PD) found similar findings of increased D₂ receptor densities in the contralateral caudate-putamen [49, 50]. There is also histologic evidence for interhemispheric projections from the striatum to the motor cortex [51], and upon lesioning, evidence of increased sprouting from intact to damaged hemispheres [52].

From our observations, it also appears as if reactive compensation is not limited to striatal structures only. In the QA model, we also found increased [¹⁸F]MK-9470 tracer activity in the cerebellum, which inversely correlates to the number of ipsiversive turnings, suggesting that animals with the highest binding were related to a better functional outcome. In agreement with this finding, reduced CB₁ receptor levels in the cerebellum have been shown to be associated with locomotor hyperactivity in congenitally hypothyroid rats [53]. Motor hyperactivity is also a well-characterized feature observed in QA-treated animals, which appears already 2 weeks post-lesioning and lasts up to 6 months [54]. Here, the amphetamine-induced asymmetry test has been performed 4 weeks following the administration of QA intrastrially, which is a time interval of 7–9 weeks from the [¹⁸F]MK-9470 PET imaging. Putative alterations of the lesions occurring within that time interval are unlikely to have affected the behavioural outcome. It has clearly been shown that the lesions induced by QA result in stable ipsiversive turnings over time (range: 2–9 weeks) [55, 56]. Also, the striatal D₂ receptor asymmetry, underlying this behavioural effect, has been shown to reach a steady state, starting 16 days post-administration [19]. The striatal affected to non-affected side BI ratio of this study was also nearly constant among subjects between 15 and 25 weeks (y = 0.0042x + 0.1875).

In the design of this study, we have explicitly set the aim to investigate CB₁ receptor network alterations by using both absolute and relative [¹⁸F]MK-9470 measurements. It is known that there is a higher physiologic interindividual variability of absolute regional [¹⁸F]MK-9470 determinations in the human brain, in the order of several tens of percent [14, 57]. Relative measurements have the advantage of being much more sensitive, allowing changes of 5–10% to be measured using SPM [58], as proven here.

Although QA neurotoxicity provokes loss of medium-sized spiny neurons in the injected striatal region, associated with motor hyperactivity [54] and spatial learning deficits [59], comparable to HD, it does not replicate the genetic hallmark underlying this disorder. This finding, together with the suggestion that (modest) species-specific differences exist in the brain CB₁ receptor distribution [14], reinforce the importance of translational research. Therefore, careful comparison to the human condition is needed to assess the in vivo validity and functional significance of our current findings, especially on the ECS which may open perspectives for neurochemical modulation of this currently untreatable neurodegenerative disorder.

In the present study, we have used QA rats because of the smaller brain size in transgenic mice which decreases sensitivity due to spatial resolution limits of small animal PET. In comparison to human studies, these QA rats received a relatively high dose of radiotracer. As the average weight of humans vs rats shows a ratio of approximately 300, care needs to be taken not to induce pharmacological effects. For the specific activity administered, the calculated % receptor occupancy of [¹¹C]raclopride was found to be in the range of 0.8–11%. This is still unlikely to induce a major pharmacological effect, such as dyskinesia, observed at an occupancy of over 50% for dopamine D₂ receptors. Also for [¹⁸F]MK-9470, occupancies that are well below the pharmacological threshold were obtained (1–10 %) [60].

In conclusion, this in vivo study suggests modest regional changes in endocannabinoid transmission specific for CB₁ receptors in the QA rodent model of HD and points towards a compensatory role of the cerebellum. Our results additionally demonstrate the occurrence of functional plasticity in the QA lesion model on metabolism, D₂ and CB₁ neurotransmission, which implicates the need for carefulness when using the contralateral side as control condition.

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Conflicts of interest None.

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