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PET Imaging Demonstrates Histone Deacetylase Target Engagement and Clarifies Brain Penetrance of Known and Novel Small Molecule Inhibitors in Rat

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Supporting Information

ABSTRACT: Histone deacetylase (HDAC) enzymes have been demonstrated as critical components in maintaining chromatin homeostasis, CNS development, and normal brain function. Evidence in mouse models links HDAC expression to learning, memory, and mood-related behaviors; small molecule HDAC inhibitor tool compounds have been used to demonstrate the importance of specific HDAC subtypes in modulating CNS-disease-related behaviors in rodents. So far, no direct evidence exists to understand the quantitative changes in HDAC target engagement that are necessary to alter biochemistry and behavior in a living animal. Understanding the relationship between target engagement and *in*



vivo effect is essential in refining new ways to alleviate disease. We describe here, using positron emission tomography (PET) imaging of rat brain, the *in vivo* target engagement of a subset of class I/IIb HDAC enzymes implicated in CNS-disease (HDAC subtypes 1, 2, 3, and 6). We found marked differences in the brain penetrance of tool compounds from the hydroxamate and benzamide HDAC inhibitor classes and resolved a novel, highly brain penetrant benzamide, CN147, chronic treatment with which resulted in an antidepressant-like effect in a rat behavioral test. Our work highlights a new translational path for understanding the molecular and behavioral consequences of HDAC target engagement.

KEYWORDS: Neuroimaging epigenetics, preclinical, translational, rodent, radiotracer

H istone deacetylase (HDAC) enzymes, particularly isoforms from class I (HDAC 1, 2, 3, and 8), class IIa (HDAC 4, 5, 7, and 9), and class IIb (HDAC 6 and 10), have been demonstrated as critical components in maintaining chromatin homeostasis, CNS development, and normal brain function. Altered HDAC expression has been linked to CNS diseases including Alzheimer's disease, bipolar disorder, schizophrenia, and major depressive disorder via post-mortem human brain research.¹⁻⁴ Modulating HDAC expression in mice has been shown to impact learning, memory, and moodrelated behaviors.⁵⁻⁷ Further, small molecule HDAC inhibitor tool compounds have been used to demonstrate the importance of specific HDAC subtypes in modulating CNSdisease-related behaviors in rodents.^{5,8,9}

However, only indirect evidence, obtained via invasive, destructive methods, exists to demonstrate the engagement of HDAC targets in brain. Inhibitor-induced changes in acetylation of histone proteins and altered gene expression from dissected brain tissue are related to HDAC function but do not provide information about the quantitative changes in HDAC target engagement necessary to alter biochemistry or behavior in a living animal.

We describe here, using positron emission tomography (PET) imaging of rat brain, the *in vivo* target engagement of a subset of class I/IIb HDAC enzymes implicated in regulating CNS-disease-related behaviors. Using a [11 C]labeled probe that permits direct visualization of HDAC subtypes 1, 2, 3, and 6,¹⁰ we demonstrate evidence of the mechanism of HDAC inhibitors in brain with known and novel HDAC inhibitors from the hydroxamate and *ortho*-aminoanilide chemical classes.

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IC₅₀ (nM) В. Α. Assav Martinostat SAHA SAHA HDAC1 0.3 4.0 HDAC2 0.8 11 ОН HDAC3 0.6 3.0 [¹¹C]Martinostat Givinostat HDAC4 1970 >30000 HN-OH HDAC5 352 8750 EC₅₀ (nM) PNC HDAC6 4.1 2.0 Assay Martinostat SAHA HDAC7 >20000 >30000 H3K9ac 100 3400 HDAC8 >15000 1020 PK5: X = S H4K12ac 100 **CN54** 1900 HDAC9 >15000 >30000 PK6: X = S(O)₂ C CI-994 Cpd60 RGFP966 **CN147** Valproic acid

Figure 1. [¹¹C]Martinostat and HDAC inhibitor tool compounds. (A) [¹¹C]-labeled martinostat is a versatile HDAC imaging probe characterized by (i) potent induction of histone acetylation in cells ($EC_{50} = 100$ nM for histone H3 lysine 9 acetylation and histone H4 lysine 12 acetylation compared with >1900 for SAHA) and (ii) robust selectivity for HDAC subtypes 1, 2, 3, and 6 ($IC_{50} = 0.3-4.0$ nM) in a recombinant human enzyme assay. (B) Hydroxamate HDAC inhibitor tool compounds. (C) *ortho*-Aminoanilide tool compounds and the short-chain fatty acid, valproic acid.

Understanding the relationship between target engagement and *in vivo* effect is essential in refining new ways to alleviate disease. This concept is established for dopamine D2-receptor drug studies, where binding measured by the D2/D3 probe raclopride (among many others) has facilitated dose-finding for human trials and provided insight into the phamacodynamic origins of clinical dopaminergic drug effects.^{11–13} Here, our work highlights a new translational path for understanding the molecular and behavioral consequences of HDAC target engagement.

RESULTS

The target engagement of HDAC subtypes 1, 2, 3, and 6 in living rodent brain has not yet been evaluated, impeding the interpretation of treatment effect relationships for myriad HDAC inhibitors in preclinical research. Therefore, we used $\begin{bmatrix} 11 \\ C \end{bmatrix}$ martinostat, a radiotracer that we have recently described¹⁰ with robust potency and selectivity for a subset of HDACs that make it an excellent in vivo PET probe (Figure 1 A and synthesis Scheme S1, Supporting Information). We first measured "baseline" tracer uptake and averaged binding in whole brain in vehicle-treated rats (n = 9). Dynamic uptake is represented in the averaged time-activity curve (Figure 2B), with raw data scaled to 100% whole brain uptake at time = 600 s post-[¹¹C]martinostat administration. Time-activity curve profiles were consistent across all scans, with percent radiotracer bound declining from a scaled mean of 100% (t =600 s) to $95\% \pm 2.2\%$ (*t* = 3600 s, mean \pm SEM). The average variance for this time range was 12.4%.

Specific target binding was demonstrated with self-blocking experiments. Radioprobe bound in whole brain was decreased following intravenous (iv) pretreatment with a dose range of unlabeled ($[^{12}C]$)martinostat, 5 min prior to radiotracer administration (Figure 2C). Blocked binding of the radiotracer by the nonradioactive martinostat was quantified by measuring

the change in the radioactivity accumulated from time 600 s to time 3600 s, and demonstrated tracer binding was reduced dose-dependently from 86.7% to 36.5% of baseline levels by pretreatment with martinostat at 0.001-2 mg/kg. (Figure 2D). We measured the blocked binding with replicate experiments at the 2 mg/kg pretreatment dose of martinostat and resolved good reproducibility from three independent experiments $(36.5\% \pm 1.75\%, \text{ mean } \pm \text{ SEM}, \text{ Figure 2D})$, with an average variance over the time range of 10.7%, consistent with the baseline group. Formal comparison by Student's t test revealed a significant difference in radiotracer binding between rats pretreated with 2 mg/kg martinostat and baseline controls (Figure 2D, $*p = 2.0 \times 10^{-8}$). Given the similarity in variance at low (baseline) and high (2 mg/kg self-blocking) measures of HDAC target engagement, we performed power analysis calculations (two-sample z test) using a representative standard deviation of 6.2% (Supportive Information, Table S1). These results showed that for blocked binding measurements $\leq 85\%$ of baseline, statistical significance could be achieved with ≤ 3 replicates, a feasible group size for preclinical imaging throughput. Moreover, we calculated that blocked binding measurements ≤92.5% of baseline could resolve statistically significant differences using a higher, but achievable 15 animals/group. Therefore, we hereafter considered blocked binding experiments demonstrating ≤92.5% of baseline tracer uptake in brain as positive initial evidence for measurable target engagement.

Quantification of absolute radiotracer uptake via total distribution volume $(V_{\rm T})$ analysis requires normalization to full arterial input function from blood sampling and was not feasible in rats. However, $V_{\rm T}$ quantification was achieved at baseline and in a range of self-blocking doses in analogous experiments conducted in nonhuman primates (NHPs) (described in detail in a forthcoming manuscript). We found that the dose-dependent decreases observed in NHP $V_{\rm T}$ were



Figure 2. Uptake of [¹¹C] martinostat in rat brain. (A) Timeline schematic of blocking dose administration and PET/CT imaging. All animals were stabilized on anesthesia at least 20 min prior to [¹¹C]martinostat administration. (B) Dynamic tracer uptake in whole brain was evaluated using ROI analysis on dynamic imaging data sets from n = 9 "baseline" rats blocked with vehicle (10% DMSO, 10% Tween 80, 80% saline) 5 min prior to tracer administration. Data are expressed as percent uptake in whole brain relative to uptake at time = 600 s (mean ± standard deviation). (C) Self-blocking of [¹¹C]martinostat tracer binding (change in % of whole brain [¹¹C] uptake) is demonstrated via pretreatment (5 min, iv) with a dose range (0.001–1 mg/kg, n = 1/group or 2.0 mg/kg, n = 3/group) of unlabeled martinostat and compared with baseline controls, described in panel B. (D) Quantification of [¹¹C] activity in whole brain at time 3600 s measured via trend in accumulated radioactivity from time 10 min to time 60 min for baseline (white bar, 0 mg/kg) and each blocking condition (grayscale). (E) Spearman correlation with directional *t* test identified significant dose–response relationships for relative [¹¹C] blockade in whole brain in rat (gray triangles, r = 0.89; p = 0.006) and whole-brain volume of distribution (V_T) from analogous PET experiments in nonhuman primate (NHP; white squares, r = 0.90; p = 0.042); * indicates blocking doses scaled for NHP equivalent dose (mg/kg). (F) Pretreatment with unlabled martinostat (1 mg/kg) 5 min, 4 h, or 24 h prior to tracer administration demonstrated time-dependent effects of target engagement, calculated as difference from baseline (100% – % blocked binding).

significantly correlated (Figure 2E, white squares, r = 0.90; p = 0.042). We also observed a significant correlation by the same test using scaled whole brain uptake levels of total [¹¹C] activity in rat (Figure 2E, gray triangles, r = 0.89; p = 0.006). This comparison gave confidence that in rat, in the absence of a blood input function, relative HDAC target binding could be resolved using the time–activity curve slope, given the unique tracer kinetics of [¹¹C]martinostat, which show a relatively nondynamic time–activity curve within the scan duration.

Using this method, we then varied the pretreatment time for nonradioactive martinostat (1 mg/kg) from 5 min to 4 h and 24 h before radiotracer administration. Target binding was blocked most (40% reduction) after a short, 5 min, blocking dose pretreatment (Figure 2F) with a lesser blockade observed 4 h after pretreatment and no appreciable change observed 24 h after nonradioactive martinostat pretreatment (n = 1 rat/blocking condition; 5 min time point same data as in Figure 2C,D.) These data demonstrate a relative peak of binding of the nonradioactive martinostat at the 5 min pretreatment time point and are consistent with the established, fast-binding kinetics of hydroxamate HDAC inhibitors like martinostat.

The hydroxamic acid class of HDAC inhibitors has been reported to modulate behavioral response in CNS-disease-

related paradigms in rodents; however the extent of target engagement in the brain has not yet been demonstrated. Therefore, we next applied the metric of $\left[{}^{11}\mathrm{C}\right]\!martinostat$ target binding in whole rat brain to investigate the relative HDAC1-3 and 6 engagement of structurally distinct hydroxamate HDAC inhibitors (structures shown in Figure 1B). Anesthetized rats were pretreated with known and novel hydroxamate HDAC inhibitors as indicated (iv pretreatement, 3-10 min prior to tracer administration). Quantification of dynamic imaging data revealed that the hydroxamates SAHA and givinostat both resulted in only subtle blockade of radiotracer binding in whole brain (Figure 3A), consistent with recent reports demonstrating poor brain penetrance of the prototypical hydroxamic acid HDAC inhibitor, SAHA.14,15 Poor brain penetrance was overcome in the development of martinostat by the addition of an adamantyl group, a chemical moiety frequently used developing in CNS-penetrant compounds.¹⁰ Therefore, we tested PK5 and PK6, two hydroxamate HDAC inhibitor compounds containing heterocyclic capping groups often found in CNS drugs.¹⁶ We found that pretreatment resulted in a modest ~20% decrease in whole brain target binding of $[^{11}C]$ martinostat (n = 2 replicates/ condition, mean \pm SEM). Further, we quantified a 40%



Figure 3. HDAC inhibitor pretreatment blocks [¹¹C]martinostat uptake in brain. (A) Pretreatment with hydroxamic acids (iv) resulted in a 10–14% blockade of tracer uptake in whole brain for the known hydroxamic acids, SAHA and givinostat (n = 1/group), with greater, 18–22%, blockade for the two novel compounds, PK5 and PK6 (n = 2/group; mean \pm SEM). Imaging experiments revealed robust 40% blockade of whole brain [¹¹C] levels by the hydroxamic acid, CN54 (n = 1/group). (B) Pretreatment with the protypical orthoaminoanilide benzamide, CI-994, via ip injection revealed modest blockade (9–14%) with limited impact of dose (15–60 mg/kg) acute pretreatment blocking time (2.5–4 h) or duration of treatment (7 days treatment with final treatment 4 h prior to tracer administration). (C) Pretreatment (ip) with the HDAC subtype-selective benzamides Cpd60 and RGFP966 also resulted in limited, 7–10%, blockade of [¹¹C]martinostat binding in whole brain. A 500 mg/kg ip dose of sodium valproate, administered 30 min prior to tracer administration blocked uptake by 16%. (D) Brain penetrance of novel benzamides was evaluated via blocked tracer binding for the tool compound CN147. CN147 demonstrated robust 23% tracer blockade at 5 mg/kg (ip, 2.5 h pretreatment). At 1 mg/kg (ip), maximal blocking effects were observed 4 h after pretreatment and were maintained at a lower dose of 0.1 mg/kg. Consistent with the slow-binding kinetics of benzamide HDAC inhibitors, no appreciable blockade was observed following a short, 5 min, pretreatment time with CN147, 5 mg/kg, ip (data not shown).

reduction in whole brain target binding of $[^{11}C]$ martinostat following pretreatment with CN54, a heterocycle-capped hydroxamate HDAC inhibitor identified from the patent literature base¹⁷ that showed a robust 4.5:1 ratio of brain/ plasma partitioning after iv treatment. Taken together, these data support that $[^{11}C]$ martinostat binding can quantify target engagement of structurally distinct, brain-penetrant hydroxamate HDAC inhibitors in living rat brain.

HDAC inhibitor tool compounds from the ortho-aminoanilide (OAA) class (Figure 1C) have been demonstrated to selectively inhibit a subset of class I HDAC isoforms and as a result have emerged as a way to evaluate selective inhibition of specific HDAC isoforms in the CNS.^{5,8,9,18,19} We investigated the impact of benzamide HDAC inhibitor treatment on [¹¹C]martinostat target binding in whole brain. Given the slow kinetics described for members of the OAA class of HDAC inhibitors,^{9,18} we administered compounds via intraperitoneal (ip) injection 2.5-7.5 h prior to tracer administration. Treatment with the prototypical benzamide-based OAA HDAC inhibitor, CI-994, a selective inhibitor for HDAC subtypes 1, 2, and 3 recently shown to enhance memory reconsolidation in mice,⁵ revealed that radiotracer binding was reduced by <14% (Figure 3B). The reduction in target binding was not altered by CI-994 dose (15, 30, or 60 mg/kg) or by protracted treatment (Figure 3B, striped bar; 15 mg/kg, 7 days

of daily ip treatment with an additional treatment on the day of PET imaging, 4 h before tracer administration).

Radiotracer target binding in whole brain was also blocked, at 7-10%, by Cpd60 and RGFP966, two selective OAA HDAC inhibitors previously shown to modulate behavior in rodents.^{8,9} These two compounds differ in their reported selectivity for recombinant HDACs with Cpd60 demonstrated as a selective nanomolar inhibitor of HDAC 1 and 2⁹ and RGFP966 reported as selective for HDAC3.8 We observed a more robust 16% reduction in basal HDAC target engagement following pretreatment with 500 mg/kg of sodium valproate, a shortchain fatty acid containing anticonvulsant and mood stabilizer with selective class I HDAC inhibitory properties (Figure 3C).^{20,21} These data are consistent with prior findings indicating limitations for brain penetrance of OAA as a compound class.^{9,22,23} They also support the notion that valproic acid may function as an HDAC inhibitor in vivo within the dose ranges that result in behavioral effects.^{24–26}

To test whether benzamide brain penetrance could be improved using a strategy of adamantane incorporation, we synthesized the novel compound CN147 (MW 429.3, Figure 1C and synthesis in Scheme S2, Supporting Information). In HDAC biochemical assays with recombinant human enzymes, this compound demonstrated HDAC selectivity (HDAC 1, 2, 3) and potency (IC₅₀ 150–600 nM) similar to CI-994, a close



Figure 4. Pharmacokinetics in rat brain and plasma. Acute treatment following single systemic treatment with (A) CN147 (1 mg/kg) or (B) CI-994 (10 mg/kg) was used to evaluate the concentration of each compound by LC-MS/MS in plasma (red) or brain tissue (blue) 5 min to 24 h after administration using n = 3 rats per compound per time point. (C) Brain/plasma ratios for CN147 and CI-994 demonstrate dramatic differences in CNS vs systemic exposure.



Figure 5. Initial behavioral impact of the novel brain penetrant HDAC inhibitor, CN147. (A) Chronic (7 days) treatment with CN147 at 0.1 mg/kg resulted in a 17% reduction in forced-swim test immobility compared with vehicle-treated controls as evaluated by a trained scorer blinded to treatment groups. Treatment with CN147 at a lower dose (0.025 mg/kg) had no appreciable effect on FST immobility score compared with controls, and analysis by Spearman correlation revealed that CN147 dose and FST immobility times were significantly correlated (Spearman r = -0.0671, two-tailed p = 0.0192). (B) Locomotor effects of CN147 treatment were evaluated after 7 days treatment with 0.5 mg/kg or 2.0 mg/kg compared with vehicle-treated controls; distance traveled (cm) was lower in CN147-treated rats than in control animals and was not considered a confound to interpretation of decreased immobility observed in the forced-swim test. No treatment-group differences were observed in rat weight gain (data not shown).

structural homologue. We then evaluated target engagement in brain by $[^{11}C]$ martinostat PET imaging. We identified a 25% blockade of tracer uptake that varied by dose (5, 1, or 0.1 m/kg) and pretreatment time (ip 2.5, 4, or 7.5 h prior to tracer administration; Figure 3D, solid grayscale bars).

Given the robust HDAC engagement indicated for CN147 by our PET data, we wanted to confirm presence in brain using an independent method. Therefore, we obtained pharmacokinetic data for CN147 (1 mg/kg) in rat brain tissue and in blood using established (invasive and destructive) LC-MS/MS methodology (Figure 4A).²⁷ The brain pharmacokinetic profile for CN147 revealed a C_{max} of 315 nM, a half-life of 6.8 h, and that the compound was present in the brain 8-24 h after ip or iv administration, a result similar to that reported for other benzamides.^{9,28} The brain to plasma exposure profile for CN147 was in marked contrast to that obtained for CI-994 (brain/plasma ratio 1:10), as measured by the same method, following a 10 mg/kg ip treatment (Figure 4B). The robust 20:1 brain partitioning of CN147 compared with CI-994 was further illustrated by coplotting brain/plasma ratios for (Figure 4C).

A growing number of reports have described the beneficial impact of OAA-based HDAC inhibitors in CNS-relevant rodent behavioral tests.^{5,8,9} On the basis of the promising brain

penetration data and evidence for HDAC target engagement from our PET imaging, we prioritized CN147 for evaluating the impact of HDAC inhibition on behavior. As a first step to provide evidence of behavioral changes induced by CN147, we chose the modified forced swim test because another OAA, Cpd60 was recently demonstrated to be ameliorative in the mouse FST after 7–10 days systemic treatment at 45 mg/kg.⁹ Given that the brain/plasma exposure of Cpd60 was 0.5 (1:20) in mice after a single 45 mg/kg ip dose, we rationalized (equilibrating for species differences in body surface area)²⁹ that a dose of CN147 in rat of 0.025–1 mg/kg would result in brain exposure close to that achieved by Cpd60 in mice.

We observed an antidepressant-like effect after 7 days ip treatment with 0.1 mg/kg CN147 treatment compared with vehicle-treated controls (two-tailed Student's *t* test *p* = 0.16, *n* = 4/treatment group; Figure 5A). Rats treated with 0.1 mg/kg CN147 showed a 17% reduction in FST immobility time compared with controls, albeit not significant by Student's *t* test (*p* = 0.17). Notably, comparison of the reduced FST immobility times (0.1 mg/kg) with rats treated at a dose of 0.025 mg/kg (*n* = 4/group) revealed a significant difference (*p* = 0.035). Further, regression analysis by Spearman correlation indicated immobility time and CN147 dose (0, 0.025, or 0.1 mg/kg) were significantly correlated (*r* = -0.6072; two-tailed *p* = 0.0196), supporting the observed antidepressant-like effects by chronic 0.1 mg/kg CN147 treatment (Figure 5A). We confirmed our ability to detect reduced FST immobility in an independent cohort of rats using an established subchronic administration of the antidepressant imipramine, which significantly reduced immobility times (104 ± 3.2 s, mean \pm SEM) compared with rats injected with vehicle (185.7 ± 19.8 s; n = 3 rats/treatment group; Student's *t* test p = 0.014).

Locomotor activity in a novel open field was evaluated in the same vehicle/CN147-treated rats, following a subsequent oneweek treatment paradigm with vehicle or a 20-fold increased dose of CN147 (treatment group assignments were increased in dose, but not scrambled). Results from a 10 min session showed that rats treated with CN147 had a lower distance traveled in a novel open field compared with vehicle-treated controls. The locomotor activity that we observed in vehicletreated rats during the 10 min session was consistent with previously reported independent results for the same strain.³⁰ This result clarifies that CN147 treatment did not induce locomotor stimulatory effects and, thus, did not confound the observed antidepressant-like effect in the FST (Figure 5B). Furthermore, it suggests that CN147, and by inference class I HDACs, may impact neurocircuits involved in both mood and anxiety-related behavior.³¹

DISCUSSION

We demonstrate here the immediate preclinical utility of investigating the brain uptake and *in vivo* target engagement of known and novel small molecule HDAC inhibitors. Increasing reports indicate the therapeutic benefit of HDAC inhibitor compounds in preclinical models, including behavioral testing of wild-type rodents,^{9,32,33} in ameliorating behavioral deficits in a genetic^{5,6} or etiological (stress) model of brain disease.³⁴ Until now, no tool has been available to profile the density or engagement of HDAC targets in such models. This, in part, has impeded translational studies of novel HDAC inhibitor drugs in humans with CNS dysfunction, despite the fact that HDAC density changes have been reported in post-mortem human brain.^{1–4} We begin here to understand the relationship between HDAC density and occupancy and disease and treatment related behaviors *in a living animal.*

The expression of HDACs in the brain is robust and widespread, 35,36 and the optimal reference region to evaluate nonspecific binding of [11 C]martinostat has not yet been resolved due to the recent development of this tool. We have shown that relative measures of binding can be made in the absence of a reference region or blood sampling, providing a readily applicable HDAC imaging tool for rodents.

Identifying lead HDAC inhibitor molecules in preclinical and translational research can be catalyzed using PET imaging with [¹¹C]martinostat, which demonstrated a high proportion of specific binding, with more than 60% of probe binding in brain prevented in homologous ([¹²C]martinostat) and heterologous (CN54) *in vivo* blocking experiments.

Our imaging results highlight a novel, highly brain penetrant HDAC inhibitor, CN147, which demonstrated an antidepressant-like effect in the forced swim test after chronic treatment. OAA compounds with similar features have been previously shown to result in beneficial changes in preclinical behavioral responses relevant to mood and memory disorders, including Cpd60, CI-994, and RGFP966.^{5,8,9} Investigating the impact of CN147 in additional behavioral paradigms and disease models will further clarify the role of HDAC targets in the brain.

Additionally, it will be important to investigate the behavioral impact of other novel HDAC inhibitors to clarify whether a low level of HDAC target occupancy in CNS is sufficient to induce meaningful molecular and behavioral changes, which is relevant for current efforts to determine whether HDACs can be targeted for beneficial therapeutic effects in a range of CNS disorders.^{37–40}

One potential explanation for a mechanism by which low engagement of HDAC targets could differentially result in such profound changes at the level of behavior is inhibitor binding selectivity at the level of HDAC multiprotein complexes. HDAC subtypes, particularly HDAC 1, 2, and 3, are members of distinct protein complexes. Efficient engagement of HDAC inhibitors to HDAC targets in vivo likely depends in part on other components of HDAC protein complexes and, potentially, on activity-dependent conformational changes.⁴¹ Therefore, it is fully possible that in vivo, OAA HDAC inhibitors only bind to class I HDAC targets in a subset of protein complexes. Further work using isolation of endogenous HDAC complexes from specific tissues of interest like the brain along with detailed analysis of selectivity and potency in combination with the use of autoradiography methods³⁶ will likely provide an additional layer of clarification and complexity to the impact of HDAC inhibitors on transcription, neural signaling, and brain output. In doing so, this strategy holds promise to facilitate improvements in HDAC radiotracer development and the application of neuroimaging to novel therapeutic development efforts.

ASSOCIATED CONTENT

S Supporting Information

Materials and methods, chemical synthesis schemes and characterization for martinostat and CN147 and statistical power analyses are included. This material is available free of charge via the Internet at http://pubs.acs.org.

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Author Contributions

Imaging experiments were contributed by F.A.S., C.W., G.C.V., H.Y.W., R.N., W.R.T., S.J.H., and J.M.H. Chemistry and radiochemistry were contributed by C.W., R.N., A.K., E.B.H., S.J.H., and J.M.H. Biochemistry and *in vitro* assays were contributed to by Y.L.Z., J.G., S.A.R., F.A.S., E.B.H., and S.J.H. Behavioral experiments were contributed by F.A.S., G.C.V., S.J.H., and J.M.H. F.A.S. drafted the manuscript with contributions from C.W., G.C.V., H.Y.W., J.G., W.R.T., S.J.H. and J.M.H.

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Notes

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ABBREVIATIONS

HDAC, histone deacetylase; PET, positron emission tomography; OAA, *ortho*-aminioanilide; iv, intravenous; ip, intraperitoneal; SEM, standard error of the mean; NHP, nonhuman primate; V_{T} , total volume of distribution; FST, forced swim test

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