ABSTRACT: Mutations in the gene encoding activin receptor-like kinase 2 (ALK2) are implicated in the pathophysiology of a pediatric brainstem cancer, diffuse intrinsic pontine glioma (DIPG). Inhibitors of ALK2 that cross the blood–brain barrier have been proposed as a method of treatment for DIPG. As part of an open science approach to radiopharmaceutical and drug discovery, we developed 11C-labeled radiotracers from potent and selective lead ALK2 inhibitors to investigate their brain permeability through positron emission tomography (PET) neuroimaging. Four radiotracers were synthesized by 11C-methylation and assessed by dynamic PET imaging in healthy Sprague–Dawley rats. One of the compounds, [11C]M4K2127, showed high initial brain uptake (SUV ∼ 2), including in the region of interest (pons). This data supports the use of this chemotype as a brain penetrant ALK2 inhibitor that permeates evenly into the pons with potential application for the treatment of DIPG.

KEYWORDS: Carbon-11, Positron emission tomography, ALK2, DIPG, Blood-brain barrier

Activin receptor-like kinase 2 (ALK2) is a kinase receptor encoded by the ACVR1 gene. Mutations in ACVR1 causing a gain-of-function in ALK2 signaling have been reported in approximately 25–30% of children with diffuse intrinsic pontine glioma (DIPG), a rare pediatric brainstem cancer with a poor prognosis.1–3 DIPG tumors are aggressive and arise from the brain’s glial tissue in the lowest stem-like portion of the brain, the pons. Although the pathology of the relationship is not yet fully known, ALK2 has emerged as a potential therapeutic target for the treatment of DIPG. Kinase inhibitors have notoriously poor selectivity, often showing off-target affinity to other kinases within a family because of the high degree of structural homology in ATP binding domains. In the case of ALK2, off-target binding to the closely related TGFβ pathway receptor ALK5 is of specific concern due to potential cardiotoxicity.4 In addition, the discovery of kinase inhibitors that are sufficiently able to cross the blood–brain barrier (BBB) has been lacking5 and there is no report of work aiming to measure the degree of penetration of DIPG-directed therapeutics into the pons.6–8 We recently reported on leveraging an open science drug discovery model9 to synthesize a series of 3,5-diphenylpyridine ALK2-selective inhibitors that cross the BBB for the treatment of DIPG.10,11 The lead compounds from this effort show brain penetration in rodent models with total brain to plasma (B/P) ratios of 0.8–1.4, although distribution into the therapeutic region of interest in the brain (pons) was unknown. Central nervous system (CNS) drug discovery can be facilitated with positron emission tomography (PET) imaging by measuring the BBB permeability of isotopically labeled drug candidates. In addition, PET imaging can be used to visualize distribution of the radiotracer into specific regions of the brain such as the pons. In addition to our recent work on ALK2 drug discovery, the goal of the present study was to apply an open science approach for PET radiotracer discovery, to label four lead potent and ALK2-selective inhibitors with carbon-11 (half-life = 20.4 min), and to assess the BBB permeability of these radiotracers by PET imaging in rodent models. Assessment of the brain permeability of these compounds as PET radiotracers identified a candidate radioligand that can be used for further drug development of ALK2 inhibitors of this chemotype, facilitating receptor occupancy, target engagement, and dosing regimen studies.

Four lead compounds with IC50 values for ALK2 between 9–19 nM were selected for radiolabeling: M4K2163,
M4K2009, M4K2117, and M4K2127. The structural position of the $^{11}$C radionuclide was chosen based on availability of the desmethyl precursors and ease of synthesis (Figure 1). While there is no consensus for the ideal physiochemical properties of small molecules that are BBB penetrant, CNS Multiparameter Optimization (MPO) scores offer a guideline for compound prioritization. Prior to radiolabeling, CNS MPO desirability scores were calculated for all of the lead compounds, giving scores of 4.34–5.04 (maximum score is 6.0), suggesting that all compounds are rational leads as brain-penetrant radioligands. Calculated logD$_7.4$ values were also within the range for most brain-penetrant PET radioligands (2.03–3.31), with limited off-target binding. As such, none of the initially identified compounds were excluded due to physiochemical characteristics that would suggest low brain permeability (Table 1).

All radiosyntheses were performed on a commercial automated carbon-11 synthesis unit for methylation reactions (GE FX2 C). [11C]M4K2163 and [11C]M4K2009 were synthesized by [11C]-O-methylation with the "loop method" using [11C]CH$_3$I and the respective phenolic precursors (M4K2297 and M4K2298; see the Supporting Information for their preparation) with <1 equiv of potassium tert-butoxide (Scheme 1A). The radioligands were synthesized, isolated, and formulated in radiochemical yields of 6%–10% ([11C]M4K2163, n = 3) and 4%–9% ([11C]M4K2009, n = 2) relative to [11C]CO$_2$. Both radiotracers were obtained in >95% radiochemical purities and high molar activities ([11C]M4K2163: 270–350 GBq/μmol), ([11C]M4K2009: 304 GBq/μmol). Small-scale vial-based radiosyntheses of [11C]M4K2117 and [11C]M4K2127 using [11C]CH$_3$I and various bases (KO$_2$Bu, NaOH, or TBAOH) resulted in low radiochemical purity and high molar activities ([11C]M4K2117 and [11C]M4K2127: 132–644 GBq/μmol). All radiotracers were prepared within 30–40 min from end of bombardment.

Shake-flask logD$_{7.4}$ values were experimentally determined using established procedures. Aliquots of the formulated radiotracers were partitioned between octanol/phosphate buffer, and the radioactivity in each layer was measured on a γ counter. Experimental logD$_{7.4}$ values were slightly higher than the calculated values and were in the range of 2.45–3.37 (Table 1), consistent with most CNS PET tracers.

The brain permeability of the radiotracers was then assessed in healthy Sprague–Dawley rats (Figure 2). Rats were injected (i.v.; tail-vein) with 16–23 MBq of each radiotracer, and dynamic PET imaging was acquired over 90 min. Time-activity histograms were determined for the brain and blood pools, and the ratio of brain to blood (B/P) was calculated. The portrait of the brain-to-blood time-activity is shown in Figure 2B. The results are consistent with the calculatedlogD$_{7.4}$ values.

Table 1. Physiochemical Properties of the Lead Compounds

| ALK2 | IC$_{50}$ (nM) | B/P | pK$_a$ | clogP | clogD$_{7.4}$ | logD$_{7.4}$ | CNS MPO |
|------|--------------|-----|-------|-------|-------------|------------|---------|
| M4K2009 | 13 | 0.97 | 8.5 | 3.20 | 2.03 | 2.76 ± 0.13 | 5.04 |
| M4K2117 | 7 | 0.82 | 9.5 | 4.56 | 2.46 | 2.64 ± 0.05 | 4.34 |
| M4K2127 | 9 | 7.6 | 3.74 | 3.31 | 3.37 ± 0.12 | 4.34 |
| M4K2163 | 19 | 1.41 | 8.5 | 4.05 | 2.88 | 2.45 ± 0.04 | 4.34 |

*Cellular IC$_{50}$ is the concentration of compound that inhibits 50% of the ALK2 activity in male SD-1 mice (2 h) @ 10 mg/kg PO. Properties retrieved from ACD/I-Lab 2.0. *Determined experimentally by shake-flask method with [11C]-labeled compounds. *Scored via an algorithm using a weighted scoring function for clogP, clogD, molecular weight, topological polar surface area, HBDs, and pK$_a$.

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Curves (TACs) were extracted for whole brain and for the pons as a region of interest using the atlas of Schwarz et al. 2006 implemented in VivoQuant (4.0patch1, Invicro). Three tracers, \([11C]M4K2163\) \((n = 2)\), \([11C]M4K2117\) \((n = 1)\), and \([11C]M4K2009\) \((n = 1)\) showed relatively low brain uptake, with peak SUV values of 1.1, 0.9, and 0.6, respectively, within 1 min after injection. Fortunately, \([11C]M4K2127\) \((n = 3)\) showed good permeability into the brain, with a peak SUV of 1.8 ± 0.2 for initial brain uptake within 1 min after injection, followed by a moderate rate of wash out over 30 min. Importantly, we found that \([11C]M4K2127\) has high uptake in the pons (peak SUV = 2.0 ± 0.3). Looking at the similar TACs for \([11C]M4K2127\) in the pons and the whole brain, it is evident the compound permeates homogeneously into the pons as compared to the rest of the brain. This, importantly, constitutes the first example of an ALK2 PET radiotracer and provides evidence that compounds of this chemotype can access the pons, which is the specific region of interest for the treatment of DIPG tumors.

To investigate the properties that could be affecting the BBB permeability of these four compounds, we reviewed their Caco-2 permeability parameters as an indication of membrane permeability as well as to consider the compounds as potential substrates of efflux transporters. As anticipated from the PET imaging results, \(M4K2127\) had the highest Caco-2 permeability, three times greater than that of \(M4K2009\) \((P_{app\ AB} = 16.6 \times 10^{-6}\ \text{cm/s})\) where \(P_{app} > 3\) indicates high permeability. Additionally, the efflux ratio for \(M4K2127\) was found to be almost 2–3-fold better than that of \(M4K2009\) and \(M4K2163\) (efflux ratio \(BA/AB = 0.9, 1.5,\) and 2.7, respectively; see the Supporting Information, Table S1). This could be attributed to the lower \(pK_a\) of the \(N\)-methyl piperazine moiety in \(M4K2127\) (Table 1) along with the molecule having a reduced number of hydrogen bond donors (HBDs), a parameter often associated with a decrease in permeability glycoprotein (P-gp)-mediated efflux. Among the four lead compounds, \(M4K2127\) showed the lowest microsomal stability (see the Supporting Information, Table S1), which is a potential hurdle for further development as a drug candidate. However, when administered via i.v. injection as in our rodent PET imaging study, \(M4K2127\) initially avoids major first-pass metabolism by the liver which increases the initial Cmax. These in vitro ADME results align well with the PET imaging data demonstrating that \([11C]M4K2127\), having the best permeability in vitro, efficiently crosses the BBB.

Herein, we present \([11C]M4K2127\) as an ALK2-targeting radioligand that satisfactorily and homogeneously permeates the brain. By employing an open science approach to this work, we were able to rapidly radiolabel isotopologs of our lead ALK2 inhibitors and apply PET imaging for the assessment of brain penetration. The present study sheds light on our previous work to establish \(B/P\) ratios in this new class of ALK2 inhibitors, with implications both for the drug development process and direct analysis of DIPG-targeting drug candidates in a clinical setting. To our knowledge, this report constitutes the first disclosure of ALK2 PET radiotracers and efforts aimed at assessing the degree of penetration into the brain (including the pons) of a compound being pursued as a potential treatment for DIPG.

While numerous compounds have entered the clinic for DIPG, no studies on their degree of penetration into the pons (using PET imaging, or any other method) have been.
reported. [11C]M4K2127 will be used as a biomarker to further assist the drug development of 3,5-diphenylpyridine ALK2 inhibitors. Blocking studies with [11C]M4K2127 to confirm specific binding in the brain of rodents and higher species, evaluation of radiometabolism, and effect of anesthesia are planned. [11C]M4K2127 may also be used as a tool in receptor occupancy, target engagement, and dosing regimens for other ALK2 inhibitors as they progress toward the clinic.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsmedchemlett.1c00127.

Details for the synthesis and characterization of M4K2127, M4K2117, M4K2297, and M4K2298; details for the radiochemical synthesis and characterization of [11C]M4K2009, [11C]M4K2117, [11C]M4K2127, and [11C]M4K2163; protocols for logD7.4 measurement and in vivo animal PET/MR imaging ADME parameters for ALK2 inhibitors and ADME assay protocols (DOCX).

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Author Contributions

The manuscript was written by E.M. and edited with D.S., I.D.G.W., and N.V., including contributions by all authors. Radiochemical tracers were synthesized by E.M. with precursors synthesized by D.S. (M4K2127, M4K2117, and M4K2297) and I.D.G.W. (M4K2298). Radiochemical tracer compounds were designed, and their syntheses were devised by E.M. and N.V. The precursor compounds were designed, and their syntheses were devised by D.S. and I.D.G.W. Caco-2 HRMS and microsomal stability studies for M4K2127 were conducted by T.K. Animal and imaging studies were performed by J.T including image analysis. E.M., I.D.G.W., and N.V. conceived and initiated the project. All authors have given approval to the final version of the manuscript.

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Notes

The authors declare no competing financial interest.

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ABBREVIATIONS

ADME, absorption distribution metabolism excretion; ALK2, activin-like receptor 2; BBB, blood–brain barrier; B/P, total brain-to-plasma ratio; CNS, central nervous system; DIPG, diffuse intrinsic pontine glioma; HBD, hydrogen bond donor; MPO, multiparameter optimization; PET, positron emission tomography; rt, room temperature; SUV, standard uptake value; TAC, time–activity curve

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