Special Section on The Opioid Crisis

Pharmacological Assessment of Sepiapterin Reductase Inhibition on Tactile Response in the Rat


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ABSTRACT

There is an unmet medical need for nonopioid pain therapies in human populations; several pathways are under investigation for possible therapeutic intervention. Tetrahydrobiopterin (BH4) has received attention recently as a mediator of neuropathic pain. Recent reports have implicated sepiapterin reductase (SPR) in this pain pathway as a regulator of BH4 production. To evaluate the role of SPR inhibition on BH4 reduction, we developed analytical methods to monitor the relationship between the plasma concentration of test article and endogenous pterins and applied these in the rat spinal nerve ligation pain model. Sepiapterin is an endogenous substrate, which accumulates upon inhibition of SPR. In response to a potent inhibitor of SPR, plasma concentrations of sepiapterin increased proportionally with exposure. An indirect-effect pharmacokinetic/pharmacodynamic model was developed to describe the relationship between the plasma pharmacokinetics of test article and plasma sepiapterin levels in the rat, which was used to determine an in vivo SPR IC50 value. SPR inhibition and mechanical allodynia were assessed coordinately with pterin biomarkers in plasma and at the site of neuronal injury (i.e., dorsal root ganglion). Upon daily oral administration for 3 consecutive days, unbound plasma concentrations of test article exceeded the unbound in vivo rat SPR IC50 throughout the dose intervals, leading to a 60% reduction in BH4 in the dorsal root ganglion. Despite evidence for pharmacological modulation of the BH4 pathway, there was no significant effect on the tactile paw withdrawal threshold relative to vehicle-treated controls.

Introduction

Chronic pain affects close to 300 million people in the United States annually. Due to the lack of therapeutic options, one common pharmacological intervention is to prescribe opioids. The mortality rate from opioid overdose exceeds 90 Americans per day (Rudd et al., 2016). Overprescription of opioids (such as fentanyl) leads to misuse and addiction, which has an increasing burden on societal and health care systems (Volkow et al., 2014). In an effort to reduce the production and use of opioid analogs, the pharmaceutical industry is investigating alternative biologic pathways of pain reduction, which has led to the rapid prosecution and validation of new pain targets. Accordingly, several putative targets for analgesia have emerged in recent years; of particular interest are enzymes that regulate tetrahydrobiopterin (BH4) synthesis. BH4 is reported to enhance pain sensitivity in the dorsal root ganglia (DRG) in preclinical models with associated links to human pain (Latremoliere and Costigan, 2011; Latremoliere et al., 2015). Subsequently, drug discovery and development efforts have emerged to reduce BH4 by developing inhibitors of biosynthetic enzymes involved in BH4 biosynthesis (Tebbe et al., 2017).

BH4 is an essential cofactor for several enzymes (nitric oxide synthase and aromatic amino acid hydrolases), and its production occurs through de novo synthesis and enzymatic salvage pathways in vivo (Latremoliere and Costigan, 2011). GTP cyclohydrolase 1 (GCH1) is elevated during chronic pain and performs the rate-limiting step in de novo synthesis of BH4 by converting GTP to 7,8-dihydروbiopterin-triphosphate (Hatakeyama et al., 1991; Tegeder et al., 2006). Sepiapterin reductase (SPR), which is also elevated during chronic pain,
is involved in both de novo biosynthesis and scavenging pathways (Tegeder et al., 2006; Latremoliere and Costigan, 2011) (Fig. 1). Since GCH1 is a master regulator of BH4 production in vivo, inhibition of GCH1 is likely to significantly reduce systemic BH4 levels, which may lead to undesirable biologic consequences. Therefore, the downstream enzyme, SPR, has emerged as a more desirable target to reduce BH4 levels because several SPR-catalyzed steps in BH4 production are also catalyzed by other enzymes (Latremoliere and Costigan, 2011; Latremoliere et al., 2015). Recent efforts have focused on development of SPR inhibitors that would lead to reduction of BH4 but not its complete elimination. As due diligence to assess new targets in the discovery pipeline for neuropathic pain, we recently evaluated candidate inhibitors of SPR. To evaluate our conviction in the target, it was necessary to benchmark internal efforts against those reported in the literature. Therefore, we developed bioanalytical methods to quantify BH4, 6-biopterin (BIO), and test article in both plasma and DRG tissues to interpret measurement of BH4 via oxidative stabilization to BIO. A total of six assays were developed to measure biomarkers and test article in both plasma and DRG matrixes (Fig. 2). These efforts were designed to support an early discovery portfolio with efficient use of resources; thus, the analytical assays were developed as fit-for-purpose without the requirement of analytical method validation (Cummings et al., 2008; Food and Drug Administration, 2013; Leonard et al., 2014; Cowan et al., 2016). These biomarker assays for BH4, BIO, and SEP were successfully used for the quantitative measurement of target engagement (SPR inhibition) in preclinical pharmacokinetic (PK) and pharmacodynamic (PK/PD) studies.

Materials and Methods

Chemicals

Dimethylsulfoxide (DMSO), water for high-performance liquid chromatography (HPLC), methanol for HPLC, isopropanol, acetone for HPLC, formic acid (89%–91%), hydrochloric acid (37%), iodine (99.9%), potassium iodide United States Pharmacopeia (USP) 99%, ascorbic acid, ammonium hydroxide (28%–30%), BH4, 6-biopterin, verapamil-HCl (VER), tolbutamide (TOL), gabapentin, and methyl sulfonic acid were obtained from Sigma-Aldrich (St. Louis, MO). BH4 was obtained from Combi Blocks (San Diego, CA). Rhamnoperin (RHAM [6-([N-(2-(5-hydroxy-2-methyl-1H-indol-3-yl)ethoxy)-2-methoxyacetamide (SPRi3) and Q-1195 was identified from US patent 2017/0096435 and a synthetic route was designed (Tebbe et al., 2017). In brief, selective hydrolysis of 2,4-dichloropyrrolotriazime followed by nucelopeinic displacement with hydrazine afforded the intermediate hydracinylpyrrolotriazine.
The condensation product of methyl acetocetate with dimethylformamide dimethyl acetal was then coupled with the hydrazinylpyrrolo-triazine to afford the methyl pyrazolecarboxylate intermediate, which was saponified to the corresponding carboxylic acid in preparation for amide formation. The amine coupling partner was prepared via reductive amination of Boc-protected diazabicyclo[3.1.1]heptane with 1,1-ethoxy (trimethylsilyl) cyclopropane, followed by Boc deprotection. Amide formation of the acid with the amine afforded Q-1195. The synthesis is further described in the Supplemental Methods.

### Instrumentation and Liquid Chromatography–Mass Spectrometry Conditions

**BIO and SEP Liquid Chromatography–Tandem Mass Spectrometry.** The HPLC system contained the following modules: CBM20A controller, solvent degassing unit, two LC20ADXR pumps, SIL30ACMP temperature-controlled autosampler, and CTO30A column oven (Shimadzu, Kyoto, Japan). Chromatographic separation was achieved on a Phenomenex Synergi 4u Polar reverse-phase (RP) (80A, 50 × 2 mm) column maintained at 40°C. Mobile phase A consisted of water with 0.002% formic acid (FA), and mobile phase B consisted of methanol with 0.002% FA. The gradient profile was as follows: 0.0–0.2 minutes at 0% B; 0.2–1.2 minutes, linear increase to 10% B; 1.2–1.3 minutes, linear increase to 100% B; 1.3–1.95 minutes, hold at 100% B; 1.95–2.0 minutes, return to 0% B; 2.0–2.5 minutes, hold at 0% B. The total run time was 2.5 minutes at a flow rate of 0.5 ml/min. Samples were kept at 15°C in the autosampler. Twenty microliters of sample extract was injected with the column effluent plumbed to an AB Sciex 4000 triple quadrupole mass spectrometer (AB Sciex, Framingham, MA). The mass spectrometer was operated in positive ion mode turbo spray with the following parameters: curtain gas at 20, GS1 and GS2 at 30, gas temperature at 500°C, collision-activated dissociation gas at 12, ion spray voltage at −4500 V, and entrance potential at −10 V. Tuning parameter settings were optimized by postcolumn infusion of 1 μg/ml analyte at 1–5 μl/min into a flow of 0.1 ml/min 50% B. All biomarker analytes and internal standards were scanned in multiple reaction monitoring mode using the parameter settings shown in Supplemental Table 1. Analyst version 1.6.1 software (AB Sciex) was used for HPLC control, data acquisition, and initial data processing.

**Q-1195 Liquid Chromatography–Tandem Mass Spectrometry Conditions.** Chromatographic separation was achieved on a Phenomenex Synergi 4u Polar RP (80A, 50 × 2 mm) column maintained at 40°C. Mobile phase A consisted of water with 0.002% FA, and mobile phase B consisted of methanol with 0.002% FA. The gradient profile was as follows: isocratic hold of 5% B from 0.0 to 0.15 minutes, linear increase to 100% B from 0.15 to 0.75 minutes, isocratic hold at 100% B from 0.75 to 1.25 minutes, linear decrease to 5% B from 1.25 to 1.3 minutes, and isocratic hold (re-equilibration) at 5% B from 1.3 to 1.7 minutes. The total run time was 1.7 minutes at a flow rate of 0.6 ml/min. Samples were kept at 15°C in the autosampler over the course of analysis. Two microliters of sample extract was injected with the column effluent in line with an AB Sciex 4000 triple quadrupole mass spectrometer. The mass spectrometer was operated in positive ion mode turbo spray with the following parameters: curtain gas at 20, GS1 and GS2 at 30, gas temperature at 500°C, collision-activated dissociation gas at 12, ion spray voltage at 4500 V, and entrance potential at 10 V. Tuning parameter settings were optimized by postcolumn infusion of 1 μg/ml analyte at 1–5 μl/min into a flow of 0.1 ml/min 50% B. Analyte and internal standard were scanned in multiple reaction monitoring mode using the parameter settings shown in Supplemental Table 2.

### Compound Stock Solutions

BH4, BIO, SEP, RHAM, Q-1195, VER, and TOL were weighed in duplicate and dissolved in DMSO:methanol (1:1) at 1.0 mg/ml as free base. RHAM required the addition of 20 mM hydrochloric acid (HCl) to solubilize. Prior to use, the duplicate 1.0 mg/ml stock solutions were analyzed with multiple injections (n = 6); the mean area was determined for each stock solution and, if within 20%, then aliquoted at 50 μl per tube and stored at −70°C for up to 4 months.

### Standard and Sample Preparation and Analysis

#### Homogenization of DRG.
Due to the challenges in handling the small DRG tissues (1 to 2 mg per DRG), DRG L4, L5, and L6 from the ipsilateral side were pooled; separately, DRG L4, L5, and L6 from the contralateral side were pooled from each subject on study. The pooled DRGs were weighed and 9 times their weight in water was added, representing a 10-fold dilution of the matrix and providing sufficient volume for subsequent sample preparation steps. Homogenization of the DRGs was accomplished using two 3-mm tungsten carbide beads per sample in snap cap tubes and processing in the TissueLyzer (Qiagen, Hilden, Germany) with a frequency of 15 cycles/s for 5 minutes.

#### Selection of Stock Solution Solvent.
There are varied literature reports of solvents used to prepare stock solutions of BH4, BIO, and SEP. Stock solutions (1.0 mg/ml) were prepared by weighing in duplicate in the following solvents: water, 25 mM HCl in water, DMSO, and DMSO:methanol (1:1) (Fukushima and Nixon, 1980a; Zhao et al., 2009; Fismen et al., 2012). DMSO:methanol (1:1) showed a near 10-fold increase in mass spectrometry (MS) signal. Subsequently, all stock solutions were prepared in DMSO:methanol (1:1). Stock solution aliquots (50 μl) in individual tubes were capped and stored at −70°C. Stock solution stability was established up to 4 months under these conditions.

#### Rat Plasma.
Endogenous levels of BIO and SEP were evaluated in six individual lots of blank K2EDTA rat plasma using this method. BIO levels were below quantitation limit (BQL) (<2.5 ng/ml) in four of six lots; the remaining two lots contained 11 and 55 ng/ml of BIO. The lots with less than 2.5 ng/ml were reserved for use as a blank matrix. SEP levels were BQL (<1 ng/ml) in all six individual lots of K2EDTA rat plasma. These results are similar to previous literature reports (Fukushima and Nixon, 1980a; Schmidt et al., 2006; Latremoliere et al., 2015).

#### Preparation of Standard Curves.
To facilitate high-throughput analysis of Q-1195, BIO, and SEP, both standards and samples were prepared in 96-well deep block microtiter plates (Microliter Analytical, Suwanee, GA). Standard curve concentrations at 1.0, 2.5, 5, 10, 25, 50, 100, and 250 ng/ml were prepared in control Sprague-Dawley rat K2EDTA plasma. The Tcan D300e was used to dispense analyte from 100 μg/ml stock solution in DMSO:methanol (1:1) into wells containing 20 μl of control plasma. Standard curve concentrations at 10, 15, 25, 50, 100, 250, 500, and 1000 ng/g in DRG homogenate (1.0 diluted in water prior to homogenization) were prepared using the Tcan D300e to dispense analyte from 100 μg/ml stock solution in DMSO:methanol (1:1) into 20 μl of control DRG homogenate.

#### Bioanalysis of Test Article, BIO, and SEP.
For determination of Q-1195, BIO, and SEP in both plasma and DRG matrices, standards and samples were processed as follows: to a 20-μl sample, 150 μl of internal standard was added (VER and TOL at 25 ng/ml in acetonitrile), vortexed for 30 seconds, and then centrifuged for 5 minutes at 3200g in an Allegra 25R centrifuge (Beckman-Coulter, Danvers, MA). A Tomtec Quadra 4 liquid handling system was used to transfer 125 μl of supernatant to a fresh 96-well deep block microtiter plate. The supernatant was taken to dryness under 15 psi N2 gas at 60°C. The dry-down procedure required 50 minutes in an UltraVap (Porvair Sciences, Wrexham, UK). Standards and samples were reconstituted in 100 μl of mobile phase A, sealed with a preslit 96-round-well plate. Supernatant was vortexed for 15 seconds in a plate shaker (VX-2500; VWR, Manasquan, NJ), and placed in the autosampler.

#### Bioanalysis of BH4.
Autooxidation of BH4 to BIO can complicate direct assessment of BH4 in biologic matrices. Stabilization of BH4 in its reduced form immediately upon sample collection has been reported (Fukushima and Nixon, 1980a; Zhao et al., 2009), but due to limited sample volumes and sizes, sample processing in this manner...
was determined to be impractical for routine analysis. Other well established reports oxidized BH4 all the way to BIO and indirectly determined BH4 levels by quantitation of BIO (Schmidt et al., 2006); this method was adopted in our analysis. Correspondingly, a separate analysis arm that implemented solid-phase extraction (SPE) was incorporated into our procedure to remove these oxidizing reagents from downstream workflow so as not to interfere with the stability of other analytes. Plasma and DRG standards and samples were processed as follows: a 20-μl sample was combined with 100 μl of RHAM (25 ng/ml in 0.1 M HCl) then vortexed for 30 seconds. Fifty microliters of oxidizing reagent (10 mg/ml iodine (I) in 20 mg/ml potassium iodide (KI) in water) was added to this mixture, vortexed for 30 seconds, and incubated for 30 minutes at room temperature in the dark. To stop the reaction, 100 μl of ascorbic acid at 50 mg/ml in water was added, and the plate(s) was vortexed for 30 seconds. Note that in addition of ascorbic acid, the standards and samples went colorless. Both the ascorbic acid and iodine solutions were made fresh immediately before use.

**SPE Method for BH4 Bioanalysis.** Oasys MCX 96-well 30-μm/30-ng plates (Waters, Wexford, Ireland) were prepared as follows: 1.0 ml of methanol was added and washed through each well with positive pressure (~1.0 psi). This was followed by 1.0 ml of water that was washed through each well, being careful to ensure that a small amount of water remained to cover the column head. The oxidized BH4 standards and samples were transferred to the plate and loaded under ~1.0-psi positive pressure. The plates were washed with 1.0 ml of 0.1 M HCl followed by 1.0 ml of methanol. BIO, from oxidized BH4, was eluted with two 250-μl additions of 8% ammonium hydroxide (NH2OH) in 25% methanol in water. The eluates were collected into a fresh 96-well plate and taken to dryness under N2 gas (15 psi) at 60°C for 1 hour. Standards and samples were reconstituted in 100 μl of mobile phase A, sealed with a preslit 96-round-well sealing mat (Phenomenex), vortexed for 1 minute on a plate shaker (VX-2500; VWR), and placed in the autosampler. Standard curve concentrations at 10, 15, 25, 50, 100, 250, 500, and 1000 ng/ml in Sprague-Dawley rat K2EDTA plasma were prepared using the Tecan D300e dispenser, and BH4 was dispensed from 100 μg/ml stock solution in DMSO:methanol (1:1) into wells containing 20 μl of blank plasma. Standard curve concentrations at 10, 15, 25, 50, 100, 250, 500, and 1000 ng/g in DRG homogenate (1:10 diluted in water prior to homogenization) were prepared using the Tecan D300e dispenser, and BH4 was dispensed from 100 μg/ml stock solution in DMSO:methanol (1:1) into 20 μl of control homogenate. Standard curves were oxidized and applied to the SPE for wash and elution as described earlier for the sample analysis.

**Standard Curve Parameters.** Standard curves were acquired at the beginning and end of each analytical run. Standard curve linear regression fitting was performed in Watson version 7.4.2 (ThermoFisher Scientific, Waltham, MA) using 1X2 weighting. The interassay precision [(S.D./mean)×100] and accuracy [(measured/true)×100] of the standard calibrators across multiple analytical runs are reported in Supplemental Table 3. Supplemental Table 4 summarizes the standard curve performance in both plasma and DRG matrices.

**Fractional Binding.** Pooled plasma and brain homogenates from male Sprague-Dawley rats were obtained from BioIVT (Hicksville, NY). Plasma was treated with lithium heparin, and each homogenate was prepared as 1 g of tissue per 4 ml of phosphate-buffered saline. Plasma or homogenate (1 ml) was spiked with test article to a final concentration of 5 μM. Triplicate 200-μl aliquots were transferred to 225 μl of ultracentrifuge tubes and spun at 37°C and 190,000g for 4.5 hours in a Type 42.2 rotor (Beckman Coulter). Twenty-five-microliter aliquots of spiked matrix were transferred to a sample plate and kept on ice to represent total drug. The remainder of spiked plasma was incubated at 37°C for the duration of the centrifugation period and used as a stability control. After centrifugation, 25-μl aliquots of the unbound fractions were transferred to the sample plate. Twenty-five-microliter aliquots of the stability control plasma were also transferred to the sample plate. All samples were matched with an equal volume of blank matrix. Samples were extracted by adding 3 volumes of acetonitrile containing internal standard (1 μM TOL) and 0.1% formic acid. Samples were vortexed, centrifuged at 4°C for 15 minutes at 3400g, and analyzed by liquid chromatography (LC)–MS using a Q-Exactive (ThermoFisher Scientific, Waltham, MA) in positive ion mode.

Fraction unbound (fu) was calculated from the ratio of test article detected in the water layer after centrifugation relative to the total concentration in the original matrix. The fu in tissue homogenates was corrected for dilution according to the equation described by Kalvass et al. (2007):

\[
fu = \frac{f_{u2} - \left( \frac{1}{\text{plate}} - 1 \right) + \left( \frac{1}{\text{matrix}} \right)}{D}
\]

where fu2 is the adjusted fraction unbound in tissue, fu1 is the estimated fraction unbound assuming no dilution, and D is the fold-dilution of the tissue.

**Formulation.** Q-1195 was formulated as a solution of 2 mg/ml in DMSO (Sigma-Aldrich) for intravenous bolus administration in rats. A platform by-mouth (p.o.) formulation vehicle of 2% (w/v) HPMC (50 cps grade; Spectrum Chemicals) and 1% (w/v) Tween 80 (Crodina) in water was used, and Q-1195 was formulated as an in situ salt in the vehicle by adjusting to pH 2 with methyl sulfonic acid (Sigma-Aldrich) for oral administration in rats.

**In Vivo Experiments.** Animals were cared for in accordance with the Guide for the Care and Use of Laboratory Animals, 11th Edition [National Research Council (U.S.) Committee for the Update of the Guide for the Care and Use of Laboratory Animals, 2011]. Animals were individually or group housed at the Association for Assessment and Accreditation of Laboratory Animal Care, an internationally accredited facility, in static or ventilated caging on corn cob bedding. All research protocols were approved by the Institutional Animal Care and Use Committee. Animals had ad libitum access to pelleted feed (2020X; Harlan, Indianapolis, IN) and reverse osmosis–purified water via water bottles or an automatic watering system. Animals were maintained on a 12:12-hour light:dark cycle in rooms at 18–26°C and a 30%–70% humidity range and had access to enrichment opportunities. All animals were determined free of specific pathogens (http://www.cverr.com/files/pdfs/rms/hmsummary.aspx). Animals were allowed at least 1 week acclimation to the facility prior to any procedures.

**Pharmacokinetic In Vivo Experiments.** A total of 18 surgically modified male Sprague-Dawley rats (8–12 weeks old) were obtained from Charles River Laboratories (Hollister, CA). Rats that were to receive intravenous administration had a femoral vein catheter for dosing and a jugular vein catheter for blood sampling. Rats that were to receive p.o. administration via gavage had only a jugular vein catheter for blood sampling. Rats were arbitrarily assigned to treatment groups (n = 3 rats/group) and received either a single 1-mg/kg i.v. bolus dose of Q-1195 administered at a dose volume of 0.5 ml/kg in 100% DMSO or a single 5-, 15-, or 45-mg/kg p.o. dose of Q-1195 administered at a dose volume of 10 ml/kg in 2% HPMC, 1% Tween 80, pH 2 adjusted with methyl sulfonic acid. Q-1195 was also administered orally once daily at 15 mg/kg over 3 subsequent days to a group of rats. For single-dose studies, blood was collected at 0.083 (intravenous treatment group only), 0.25, 0.5, 1, 2, 4, 6, 8, and 24 hours postdose. For multiple-dose studies, blood was collected...
PK/PD Modeling of Q-1195 Exposure to SPR Inhibition

Phoenix WinNonlin/NLME software (Certara USA, Princeton, NJ) was used for the PK and PK/PD modeling. Parameter imprecision was estimated using 95% confidence intervals (CIs) around the mean estimates. The plasma PK after single-administration i.v. (1 mg/kg) and p.o. (5, 15, and 45 mg/kg) dosing was consistent with a linear extravascular one-compartmental model with first-order elimination. Because lower than dose-proportional plasma PK was observed at the 45-mg/kg dose, bioavailabilities were estimated for each dose, and the doses were adjusted for bioavailability when fitting the PK/PD model. Sequential PK/PD model fitting was performed for animals with plasma SEP data. A standard indirect-effect model (inhibition of dissipation) (Mager et al., 2003) was used to describe the effect of Q-1195 on the time course of SEP in plasma with Imax fixed at 1. This model is consistent with the anticipated mechanism of Q-1195 inhibiting the conversion of SEP to BH4 by SPR, with the plasma levels of SEP as the pharmacodynamic (PD) biomarker. The PK/PD model and best-fit parameters for the 5-, 15-, and 45-mg/kg data sets were used to project the plasma concentrations of Q-1195 and SEP following the 60-mg/kg p.o. dose administered during the open-field activity and spinal nerve ligation (SNL) studies (described later).

Rat Open-Field Activity

Male Sprague-Dawley rats were habituated in a reversed light cycle room in home cages for at least 1 week and acclimated to the testing room for 1 hour prior to dosing. Animals were counterbalanced to n = 8 per treatment groups. Q-1195 (15 and 60 mg/kg) or vehicle (2% HPMC, 1% Tween 80, pH 2.2, with methyl sulfoxide acid) was administered 3 hours before placing an animal in the open-field apparatus. Open-field activity was measured using a system that counts interruptions of a set of photobeams for the course of 30 minutes (Kinder Scientific, Poway, CA). To begin a session, animals were removed from the home cage and placed individually into an independent Plexiglas box [41 x 41 x 38 cm (length x width x height)] surrounded by a frame consisting of 32 photocells (16Y and 16X) that track the movement of the animal. Photobeam breaks were used as an indication of activity and were measured as the following parameters per minute: basic movements (counts) and total rearing (counts). Following completion of behavioral measurements, animals were euthanized with carbon dioxide followed by immediate blood collection via cardiac puncture for pharmacokinetic analysis (approximately 3.5 hours postdose). Blood collections were mixed in SARSTEDT Microvette K3EDTA plasma separation tubes and centrifuged at 4°C at 14,000g for 5 minutes to generate plasma. All plasma and DRG specimens were maintained at −70°C (±10°C) prior to PK/PD analysis. All behavioral data were scored by an automated device.

SNL-Induced Tactile Alloodynia

Adult male Sprague-Dawley rats weighing 120–150 g were obtained from Harlan (Livermore, CA) and kept in the reversed light cycle for the entire study. SNL surgeries were performed under a stereomicroscope using standard aseptic surgical techniques as described previously (Kim and Chung, 1992). Spinal nerve injury was caused by ligating the left L5 and L6 (ipsilateral) spinal nerves, with special care taken to avoid damage to the L4 spinal nerve. In brief, under gaseous anesthesia with a mixture of O2 and isoflurane (4% for induction and 2% for maintenance), skin was excised and longissimus lumborum muscle and the fascia above L6 spinal nerves were carefully removed. These procedures provided a clean and spacious working area to enable complete resection of the L6 transverse process and to separate the L5 from L4 spinal nerves. The L5 and L6 spinal nerves were each tightly ligated with 6-0 silk suture. The entire surgical procedure, beginning with skin incision and ending with wound clipping of the skin, lasted 7 minutes or less.

SNL Behavioral Testing

Two weeks postoperation, mechanical sensitivity was measured by determining the median 50% ipsilateral hind paw withdrawal threshold for von Frey filaments using the up-down method. Rats were placed under a plastic cover (9 x 9 x 20 cm) on a metal mesh floor (Chaplan et al., 1994). von Frey filaments (Semmes-Weinstein monofilaments; Stoelting, Wood Dale, IL) were applied to the middle glabrous area between the footpads of the plantar surface of the injured hind paw. This plantar area was touched with a series of nine calibrated von Frey filaments with approximately exponentially incremental bending forces (von Frey filament numbers: 3.61, 3.8, 4.0, 4.2, 4.41, 4.6, 4.8, 5.0, and 5.2; equivalent to 0.41, 0.63, 1.0, 1.58, 2.51, 4.07, 6.31, 10, 15.8 g). The von Frey filament was presented perpendicular to the plantar surface with sufficient force to cause slight bending and held for approximately 3 to 4 seconds. To avoid possible reflex responses, only abrupt withdrawal of the foot accompanied by pain-indicative behaviors (namely, paw flinching, shaking, or licking for more than 2 seconds) was recorded as a response. Any postsurgery rat that displayed a mechanical threshold of more than 3.16 g or less than 0.7 g was eliminated from the study.

After measuring basal threshold, animals were randomized into four groups (a minimum of 10 animals per group) and treated orally with vehicle (single dosed), Q-1195 (60 mg/kg, single dosed or multidosed once per day for 3 days), or gabapentin (100 mg/kg, single dosed). Measurement of the tactile threshold was assessed at 3 hours postdose for Q-1195–treated and vehicle groups and at 1 and 2 hours postdose for gabapentin-dosed animals. Following completion of behavioral measurements, animals were euthanized with carbon dioxide followed by immediate blood collection via cardiac puncture for pharmacokinetic analysis (approximately 3.5 hours postdose). Lumbar DRGs (L4, L5, and L6) from the ipsilateral side of individual animals were collected and pooled in 1.5-ml microcentrifuge tubes; separately, DRG L4, L5, and L6 from the contralateral side were pooled from each subject into microcentrifuge tubes. Blood collections were mixed in SARSTEDT Microvette K3EDTA plasma separation tubes and centrifuged at 4°C at 14,000g for 5 minutes to generate plasma. All plasma and DRG specimens were maintained at −70°C (±10°C) prior to LC-MS/MS analysis.

Human and Rat SPR Protein Production

SPR Cloning and Expression. Human (NM_003124.5) and rat (NM_019181.1) sepiapterin reductase genes were synthesized by Integrated DNA Technologies and cloned into pET28 vector (Millipore Sigma, Burlington, MA) containing an N-terminal FLAG tag followed by thrombin cleavage site. The sequence confirmed expression constructs were transformed into BL21(DE3), and subsequently, a single colony of each construct was inoculated into 60 ml of Terrific Broth medium (MilliporeSigma, Burlington, MA) supplemented with 50 μg/ml kanamycin. The overnight cultures were used to inoculate 6 l of Terrific Broth/kanamycin at a 1:100 (v/v) ratio. The cultures were grown at 37°C by shaking at 250 rpm until optical density 600 (OD600) of ~1.0; protein expression was then induced by the addition of 0.5 mM isopropyl-β-D-thiogalactopyranoside for 3 hours. Cells were pelleted by centrifugation, washed once with Tris-buffered saline
Burlington, MA) with 10-kDa molecular mass cutoff.

**SPR Protein Purification.** Frozen cells were resuspended in TBS with protease inhibitor cocktail (Roche) and lysed by a microfluidizer. Cell debris was removed by centrifugation at 40,000 rpm at 4°C using a Beckman Coulter Ti75 rotor. The cleared cell lysates were incubated with 3 ml of anti-FLAG M2 resin (Sigma-Aldrich) overnight at 4°C with gentle nutation. Resin was packed into a gravity column and washed with 20 column volumes of TBS, and proteins were then eluted with 20 ml of TBS supplemented with 0.15 mg/ml 2X FLAG peptide (Sigma-Aldrich). SPR proteins were further purified by size-exclusion chromatography using a HiLoad 16/60 Superdex-200 column (GE Healthcare, Chicago, IL) equilibrated in buffer B (20 mM 4-morpholineethanesulfonic acid (MES) (pH 6.5), 100 mM NaCl). SPR-containing fractions were pooled and concentrated to ~0.5 mg/ml using Amicon Ultra-15 filters (MilliporeSigma, Burlington, MA) with 10-kDa molecular mass cutoff.

**Biochemical Assessment of SPR Binding.**
A low-volume, high-throughput NADP/NADPH-Glo Assay (Promega, Madison, WI) was developed to measure the NADPH-dependent SPR reduction of SEP to 7,8-dihydrobiopterin. The conversion of NADPH to NADP is proportional to the conversion of SEP to 7,8-dihydrobiopterin by SPR. NADP/NADPH-Glo technology generates a luminescent signal upon reaction with reductase, reducing the proluciferin substrate to luciferin. Luciferin is then detected with Ultra-Glo Luciferase (Promega), where the light generated is proportional to the NADPH in the sample. Inhibition of SPR is assessed by the inhibition of luciferin signal in response to increasing concentrations of inhibitor.

Assay plates were prepared by dispensing either human FLAG-SPR or rat FLAG-SPR to final concentrations of 10 and 3 nM, respectively, in assay buffer (25 mM HEPES (pH 7.5), 150 mM NaCl, 0.01% Tween-20) containing 10 nM SEP into a Corning 384-well white, low-volume plate with a Thermo Multidrop Combi. Inhibitors were dissolved in 100% DMSO to a top concentration of 5 mM and then serially diluted (1:2 stepwise) with 100% DMSO to generate a 22-point curve in a standard 384-well polypropylene plate. Assay buffer supplemented with 10 µM NADPH was added to the inhibitor titration plate, and the reaction was initiated by transferring from the inhibitor titration plate to the assay plate using a Thermo VPrep liquid dispenser. The final top concentration of inhibitor was 50 µM in the assay. The reaction was incubated for 40 minutes at room temperature and stopped by the addition of sulfasalazine (to a final concentration of 10 µM). The detection buffer NADP/NADPH-Glo was prepared and added to the stopped reaction as recommended by the manufacturer (omitting the use of NADP cycling enzyme). After 45 minutes of incubation, the plate was read on an Envision 2104 Multimode Plate Reader (PerkinElmer) using the ultrasensitive luminescence-detection method.

**Statistical Analyses.**
Since the von Frey filament set was calibrated on a logarithmic scale by the vendor (Stoelting), our selection of nine filaments for the up-down method was also based on nearly equal logarithmic intervals, and because it is our experience that variability noticeably increases with threshold value (Dixon, 1980), data were analyzed following logit transformation prior to statistical analysis. Actual gram values were plotted on a gram scale y-axis of the figures for convenience. Behavioral data are expressed as the S.E.M. Results were analyzed using one-way analysis of variance (ANOVA) with Bonferroni’s multiple-comparisons post hoc test for significance relative to vehicle. Statistical calculations and graphs were made using GraphPad Prism 7.01 (GraphPad Software Inc., San Diego, CA).

**Results.**

**Chemical Synthesis and Biochemical Inhibition of SPR Inhibitors.** Literature evaluation of chemical matter revealed several molecules that have demonstrated biochemical inhibition of SPR as well as reduction in BH4 in vivo. Synthesis of these molecules was initiated; concurrently, we developed a biochemical assay to assess in vitro potency of inhibition. The SPR NADPH-Glo assay IC50 determinations were consistent with reported values in Table 1 (Zhao et al., 2009; Latremoliere et al., 2015; Tebbe et al., 2017). Q-1195 was determined to be the most potent inhibitor of those tested. Thus, Q-1195 was moved forward into in vivo characterization to determine its suitability as a tool molecule to probe for behavioral and biochemical changes associated with SPR inhibition in vivo.

**Bioanalytical Method.** Accurate and reliable bioanalytical methods were required to monitor the pterin biomarkers associated with SPR inhibition. A coordinated series of method development steps were undertaken in a fit-for-purpose matter during bioanalytical method development. Several bioanalytical methods provided the basis for internal development efforts (Fukushima and Nixon, 1980b; Schmidt et al., 2006). Details of these investigations may be found in the Materials and Methods section. In brief, stock solutions were prepared in 1:1 DMSO:methanol, which resulted in a near 10-fold increase in MS response compared with aqueous and DMSO-alone solutions. A shallow LC gradient with methanol and water (containing 0.002% formic acid) was selected with separations performed on a Phenomenex Polar RP column. Stabilization of BH4 to BIO was achieved by oxidation of matrix (plasma or DRG) using KI in HCl solutions. Biologic samples (plasma and DRG) were split and processed by two routes: 1) protein precipitation with Q-1195 detection in positive ion mode with threshold value (Dixon, 1980), data were analyzed following logit transformation prior to statistical analysis. Actual gram values were plotted on a gram scale y-axis of the figures for convenience. Behavioral data are expressed as the S.E.M. Results were analyzed using one-way analysis of variance (ANOVA) with Dunnett’s multiple-comparisons post hoc test for significance relative to vehicle. Statistical calculations and graphs were made using GraphPad Prism 7.01 (GraphPad Software Inc., San Diego, CA).

**TABLE 1**

<table>
<thead>
<tr>
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<tr>
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<tr>
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SPR3, N-[2-(5-hydroxy-2-methyl-1H-indol-3-yl)ethyl]-2-methoxyacetamide.

**PK/PD Relationship with Sepiapterin Reductase Inhibition**

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SPR3, N-[2-(5-hydroxy-2-methyl-1H-indol-3-yl)ethyl]-2-methoxyacetamide.
corresponding unbound in vivo SPR IC50 well with measured values (Phoenix WinNonlin/NLME and adequately described the PK/PD-derived in vivo SPR IC50 for inhibition of the conver-
mination (5 mg/kg) demonstrated good bioavailability (86.7%) with a Cmax of 26.3 \mu M at 2 hours postdose. Based on these initial findings, Q-1195 was deemed suitable for subsequent p.o. dose-escalation studies. P.o. dose escalation at 15 and 45 mg/kg indicated less than dose-proportional increases in area under the curve (AUC) (Table 2). Additionally, repeat dosing of Q-1195 at 15 mg/kg for 3 days demonstrated minimal dose accumulation (day 1 AUCinf = 434 \mu M*h; day 2 AUCinf = 509 \mu M*h). A PK model was developed using Phoenix WinNonlin/NLME and adequately described the PK of Q-1195 (Fig. 3); predicted concentrations correlated well with measured values (R^2 = 0.91). The plasma PK was 80% of dose-proportional between 5 and 15 mg/kg and 46% of dose-proportional between 5 and 45 mg/kg, suggesting decreased bioavailability. The model estimations of bioavailability were consistent with this trend, fitting to 100% (CI: 80%–120%) at 5 and 15 mg/kg and 58% (CI: 53%–63%) at 45 mg/kg. PK parameter values and CI estimated with the model are shown in Table 3.

**PK/PD Results.** Concomitant with the assessment of plasma concentrations of Q-1195, SEP was measured in plasma for each dose group (intravenous and p.o.). The appearance of SEP in plasma increased in response to increased p.o. exposure of Q-1195, indicating Q-1195 was engaged with its target (SPR) and that SPR inhibition was measurable in the plasma compartment (Fig. 3). An indirect PK/PD model adequately described the relationship between plasma PK of Q-1195 and the plasma SEP levels (Fig. 3). Predicted SEP levels correlated with measured values (R^2 = 0.86). The PK/PD-derived in vivo SPR IC50 for inhibition of the conversion of SEP to BH4 by SPR was 2.2 \mu M (Table 3), with the corresponding unbound in vivo SPR IC50 = 0.44 \mu M. The PD response was 80% dose proportional between 5 and 15 mg/kg and 74% dose proportional between 5 and 45 mg/kg, demonstrating that increased Q-1195 exposure led to an increase in SEP concentration. We then used the indirect PK/PD model to project a dose that would cover the in vivo SPR IC90 at Cmin for use in subsequent pain studies. Results from the model projection indicated that a Q-1195 dose of 60 mg/kg dose yields plasma concentrations at Cmin exceeding 2.2 \mu M.

To determine the extent of SPR inhibition at the presumed site of action (DRG), SEP was measured in excised DRG from intact rats. After p.o. dosing at 15 mg/kg, SEP levels in DRG increased from undetectable (<15 ng/g) in sham-dosed animals to 26.0 ng/g on day 1 and 59.0 ng/g on day 3. The corresponding plasma concentrations of SEP were 20.8 and 19.1 ng/ml on day 1 and day 3, respectively. Q-1195 was measured in the same DRG samples and found to be 9.1 nmol/g (9.1 \mu M) on day 1 and 11.8 nmol/g (11.8 \mu M) on day 3. The corresponding plasma concentrations of Q-1195 were 29.6 and 30.7 \mu M on day 1 and day 3, respectively.

**Q-1195 Did Not Affect Open-Field Activity.** Potential analgesic effects of a new chemical entity can be confounded by behavioral changes that produce sedative or motor side effects and may lead to a false-positive response. Therefore, prior to testing in a pain behavioral model, we evaluated Q-1195 in an open-field assessment to observe general behavior changes after exposure to test article. Following 15 and 60-mg/kg p.o. administration of Q-1195, the mean plasma concentrations at 3.5 hours postdose were 79.8 and 193 \mu M, respectively, while the SEP plasma concentrations were 41.1 and 59.8 ng/ml, respectively. The C_{plasma,plasma} was 38.6 \mu M at the 60-mg/kg dose, indicating that an 8.6-fold coverage of the unbound in vivo SPR IC90 was achieved near Cmax. The 60-mg/kg projection from the PK/PD model matched reasonably well to the data, with both the Q-1195 and SEP falling within 2-fold of the prediction (Fig. 3). The measured Q-1195 plasma concentrations were 1.5-fold higher than projected (130 \mu M), and the measured plasma SEP concentrations were 1.6-fold higher than projected (37.9 \mu M). At 3 hours postdose, basic movement counts (5532 ± 165.7) were not significantly different relative to the vehicle-treated group (5406 ± 386.2), and total rearing counts (184.4 ± 14.8) were not significantly different relative to the vehicle-treated group (173.3 ± 15.1) (Fig. 4, A and B). Similarly, at the lower dosing group (15 mg/kg),

### TABLE 2

<table>
<thead>
<tr>
<th>Route</th>
<th>Dose</th>
<th>AUCinf</th>
<th>CL</th>
<th>Vss</th>
<th>t1/2</th>
<th>%F</th>
<th>AUCt</th>
<th>AUCt/dose</th>
<th>AUCt/AUCi</th>
<th>low dose</th>
<th>% of linear</th>
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<tbody>
<tr>
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<td>1.0</td>
<td>42.3</td>
<td>0.063</td>
<td>0.4</td>
<td>4.5</td>
<td>87</td>
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<td></td>
</tr>
<tr>
<td>p.o.</td>
<td>5.0</td>
<td>183</td>
<td>5.5</td>
<td>87</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>p.o.</td>
<td>15</td>
<td>434</td>
<td>9.1</td>
<td>100</td>
<td>39</td>
<td>1.0</td>
<td>100</td>
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</tr>
<tr>
<td>p.o.</td>
<td>15</td>
<td>509</td>
<td>9.0</td>
<td>15</td>
<td>31</td>
<td>2.4</td>
<td>80</td>
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<tr>
<td>p.o.</td>
<td>45</td>
<td>850</td>
<td>7.8</td>
<td>50</td>
<td>21</td>
<td>4.1</td>
<td>46</td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

AUC_{inf}, area under the curve to infinity; AUCt, area under the curve to last time point; CL, clearance; %F, bioavailability; t1/2, half-life; Vss, volume of distribution at steady state.

*Pharmacokinetic parameters determined by noncompartment analysis.

*Pharmacokinetic parameters determined by noncompartment analysis.

*Day 1.

*Day 2.
no significant effect on basic movements or total rearing counts was observed. Results from the open-field assessment indicate that any behavioral change in the SNL assessment would likely be attributable to a pharmacological effect rather than an overt behavioral change.

**Q-1195 Did Not Reverse SNL-Induced Tactile Allodynia.** Following no observable effect in the open-field assessment, Q-1195 was assessed for its ability to reverse tactile allodynia in the SNL rat pain model. Q-1195 was administered at 60 mg/kg p.o. for 3 consecutive days to SNL rats (n = 10); testing was assessed on day 1 and reassessed on day 3 to evaluate the pharmacological response from acute and repeat administration, respectively. The typical baseline, presurgery threshold for tactile response was 13.5 g. Two weeks after SNL surgery, sham-treated rats exhibited tactile thresholds of 2.53 ± 1.45 g (3 hours post–single dosing of vehicle) and 2.30 ± 1.71 g (3 hours post-multidosing of vehicle on day 3), indicating SNL surgery produced the expected end-point of reduced pain threshold. Gabapentin (100 mg/kg p.o.) was administered as a positive control to separate cohorts of SNL rats on day 1 or day 3 of the experiment. Previous open-field assessment of gabapentin at this dose level was determined to have no impact on basic movements or rearing counts (negative in the open-field assessment). Gabapentin significantly reversed SNL-induced tactile allodynia with an increase in the tactile threshold to 8.94 ± 3.95 g (day 1) and 10.12 ± 4.45 g (day 3). Q-1195 at 60 mg/kg PO did not increase the tactile threshold relative to vehicle on either day 1 or day 3 of this multidose study (P > 0.05 by Dunnett’s multiple comparisons test), with observed mean tactile thresholds of 2.61 ± 1.45 and 2.94 ± 1.71 g on day 1 and day 3, respectively (Fig. 4C).

**Biochemical Assessment of SPR Inhibition in the SNL Rat.** In addition to the von Frey behavioral response assessment of tactile threshold, biochemical endpoints of SPR inhibition (BH4 and SEP) were also evaluated. Plasma samples were collected after behavioral assessment (3.5 hours postdose) on day 1 and day 3 of the study, and DRG were

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>%CV</th>
<th>Mean</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>kₐ</td>
<td>Adsorption rate</td>
<td>20</td>
<td>2.5 l/h</td>
<td>1.5–3.6 l/h</td>
</tr>
<tr>
<td>V</td>
<td>Volume of distribution</td>
<td>8.9</td>
<td>0.92 l/kg</td>
<td>0.76–1.1 l/kg</td>
</tr>
<tr>
<td>CL</td>
<td>Clearance</td>
<td>2.9</td>
<td>0.075 l/h/kg</td>
<td>0.071–0.079 l/h/kg</td>
</tr>
<tr>
<td>kₛₑ</td>
<td>SEP synthesis rate</td>
<td>9.6</td>
<td>18 μg/l/h</td>
<td>14–21 μg/l/h</td>
</tr>
<tr>
<td>kₑₑ</td>
<td>Baseline rate of SEP conversion to BH4</td>
<td>26</td>
<td>14 l/h</td>
<td>7.0–22 l/h</td>
</tr>
<tr>
<td>IC₅₀</td>
<td>Concentration at half-maximal inhibition</td>
<td>29</td>
<td>2.2 μM</td>
<td>0.89–3.5 μM</td>
</tr>
</tbody>
</table>

CL, clearance; %CV, percent coefficient of variation.
Vehicle-dosed animals were run as the negative control. Analysis of plasma samples (3.5 hours postdose) from day 1 to day 3 demonstrated Q-1195 concentrations of 241 and 231 \( \mu M \), respectively (Fig. 4D). These plasma levels were consistent with (and slightly higher than) the measured plasma concentrations from the 60-mg/kg p.o. dose in the open-field activity assessment. The unbound in vivo IC\(_{90}\) coverage was determined to be 11-fold on both day 1 and day 3. Correspondingly, an increase in plasma SEP levels was observed in Q-1195–treated rats relative to the vehicle-dosed controls. SEP plasma concentrations increased from undetectable (lower limit of quantitation 1 ng/ml) to mean plasma levels of 46.1 and 55.6 ng/ml on day 1 and day 3, respectively, following administration of 60 mg/kg Q-1195. Vehicle-treated SNL rats exhibited tactile thresholds of 2.53 ± 1.45 and 2.30 ± 1.71 g on day 1 and day 3, respectively. SNL rats treated with 60 mg/kg Q-1195 displayed tactile thresholds of 2.81 ± 1.45 and 2.94 ± 1.71 g on day 1 and day 3, respectively, which was indistinguishable from vehicle-treated controls. SNL rats treated with 100 mg/kg gabapentin displayed tactile thresholds of 8.94 ± 3.95 and 10.12 ± 4.45 g on day 1 and day 3, respectively, which was statistically different from vehicle-treated controls. (D and E) LC-MS/MS analysis of plasma samples from Q-1195–treated SNL rats demonstrated exposure to Q-1195 (241 and 231 \( \mu M \) on day 1 and day 3, respectively) and elevations in SEP (46.1 and 55.6 ng/ml on day 1 and day 3, respectively) relative to vehicle-treated controls. (F and G) Q-1195 was quantified in DRG on day 3 and determined to be 26.8 \( \mu M \); SEP levels in the ipsilateral and contralateral DRG of SNL rats were 156.8 and 57.6 ng/ml, respectively, upon exposure to Q-1195 at 60 mg/kg p.o. for 3 consecutive days. (H) LC-MS/MS analysis of DRG samples from Q-1195–treated SNL rats demonstrated a 60% reduction in BH4 from 124.7 to 51.5 ng/ml relative to vehicle control in the ipsilateral DRG \((P = 0.008)\). In the vehicle-treated group, BH4 was elevated in the ipsilateral DRG but not the contralateral DRG \((P < 0.0001)\). Contra, contralateral; Ipsil, ipsilateral.
an internal control to monitor changes in BH4. Quantitation of BH4 in the ipsilateral DRG of vehicle-dosed animals demonstrated a 6.7-fold increase relative to the contralateral DRG (from 18.5 to 124.7 ng/g) (Fig. 4H), which indicated SNL surgery led to elevated BH4 levels, as previously reported (Latremoliere et al., 2015). This is also in agreement with the decrease in tactile threshold observed during von Frey testing (described earlier). Analysis of the ipsilateral DRG from Q-1195–dosed animals demonstrated a 59% reduction in BH4 levels relative to sham-dosed controls (from 124.7 to 51.5 ng/g). Despite this decrease in BH4 in Q-1195–exposed rats, there was no significant increase in tactile threshold observed during von Frey testing. In good agreement with the decrease in BH4 in the ipsilateral DRG following exposure to Q-1195, levels of SEP were significantly increased over sham-decrease in BH4 in the ipsilateral DRG following exposure to Q-1195

6 Revealed the presence of Q-1195 at levels of 26.8 ng/g (from 18.5 to 124.7 ng/g) (Fig. 4H), which indicated SNL

9 projections, the unbound plasma concentrations from a 60-mg/kg PO dose in the SNL experiment were designed to exceed the in vivo SPR IC50 at Cmin over the dose interval. Plasma quantitation of Q-1195 in the SNL experiment demonstrated the unbound plasma concentrations exceeded the in vivo unbound IC50 at Cmin as well and were consistent with the projections.

The opportunity to assess target engagement (SPR inhibition) through SEP appearance in a typical p.o. PK experiment was very attractive. The PK/PD model produced a reliable prediction of SPR inhibition and likely would have been a useful tool in a drug-discovery campaign, provided a relationship between PD and efficacy was established. This approach would have minimized the reliance on SNL studies, which are time- and resource-intensive. Prioritizing compounds based on both PK and PD early in the discovery flow scheme likely would have provided an efficient use of resources, whereby the SNL studies would have been reserved for true lead candidates, and greater confidence in the outcomes would be expected.

There was a large difference in potency observed between the calculated in vitro rat SPR IC50 (0.004 μM) and the in vivo SPR IC50 (2.2 μM) and unbound in vivo SPR IC50 = 0.44 μM) of approximately 110-fold. Prior studies demonstrated a strong right shift in potency from biochemical to cell for human SPR with an SPRi3 of approximately 70-fold (not adjusted for fractional binding in the cell-based assay) (Latremoliere et al., 2015). Large shifts in potency from biochemical to cell-based assays are common; further, shifts from cell-based potency to in vivo potency are frequently right shifted. The biochemical, recombinant protein-based assay is largely a measure of potency and may not fully account for mode of action and intracellular compartment effects that would be increasingly prevalent from cell culture to in vivo. In the absence of a reliable cell-based potency assessment, we used the calculated in vivo assessment of SPR inhibition (measured by the increase in SEP in plasma) to benchmark the target coverages for the determination of efficacy in the SNL investigation.

Despite biochemical evidence for inhibition of SPR in both plasma and DRG, there was no observable increase in tactile paw withdrawal for SNL rats. The reason for this lack of behavioral change is unclear, but several explanations are possible. A possible disconnect between the unbound plasma coverage and behavior effect may result from unanticipated differences in unbound concentrations of test article in the DRG. SEP, BH4, and Q-1195 were all quantified in DRG; increased levels of SEP were observed as well as a decrease

Discussion

To support the early pipeline and evaluate novel nonopioid targets for analgesia, we developed a fit-for-purpose analysis suite to quantify both test article and relevant target-specific biomarkers in a pharmacological model of pain sensitivity. The requirement to generate data for multiple analytes from low-volume rodent samples presented several analytical challenges. The methods reported here accommodated the analysis of BIO and SEP in plasma using the same negative ion acquisition method. Q-1195, along with other test articles, in plasma was successfully analyzed by immediate reinsertion of the sample plates using a positive ion acquisition method.

The methods reported here allowed for the rapid turnaround of bioanalytical results for BH4, BIO, SEP, and Q-1195 in both plasma and DRG samples. The turnaround time for the complete analysis of a behavioral (SNL) study including sample preparation, data analysis, and report generation was approximately 4 days.

The DRG matrix presented unique sample-handling challenges. Homogenates were prepared for both matrix spiking of analytical standard curves and the analysis of in vivo samples. Homogenates of DRG were typically prepared as 2:1 mixtures of PBS:tissue. Gradually, these mixtures would seize and become unworkable to any subsequent liquid transfers, and it became impossible to determine fractional binding in DRG by ultracentrifugation because a total recovery sample could not be reproducibly transferred due to these severe matrix effects. Homogenates diluted 10:1 in PBS:tissue produced similar results. However, when total recovery was desired, precipitation and extraction with organic solvent (acetone) was successful as evidenced by the linearity of spiked standard curves used for bioanalysis. Liu et al. (2018) recently reported a strong correlation in fractional binding among neural tissues (brain, spinal cord, sciatic nerve, and DRG) by using rapid equilibrium dialysis. Dialysis was not attempted in our analysis, but it's possible that congealing events observed in our studies may be overcome using dialysis, provided the solidified DRG matrix permitted small molecule equilibrium to be achieved between dialysis solution reservoirs.

Application of these bioanalytical methods to preclinical studies enabled reliable determinations of test article and pterin metabolites in both plasma and DRG. Collectively, this provided a quantitative basis to derive PK/PD relationships and assess pharmacological inhibition of SPR in a rat model of neuropathic pain. Using the PK analysis and PK/PD projections, the unbound plasma concentrations from a 60-mg/kg PO dose in the SNL experiment were designed to exceed the in vivo SPR IC50 at Cmin over the dose interval. Plasma quantitation of Q-1195 in the SNL experiment demonstrated the unbound plasma concentrations exceeded the in vivo unbound IC50 at Cmin as well and were consistent with the projections.

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PK/PD Relationship with Sepiapterin Reductase Inhibition

485

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in BH4, which suggested that SPR inhibition was achieved in DRG. Total concentrations of Q-1195 were measured in DRG, but the unbound fraction estimates were determined from brain matrix studies ($f_{u,brain}$). Even though there appears to be a strong relationship between fractional binding in brain relative to DRG (Liu et al., 2018), it is possible that some chemotypes will deviate from this relationship. If the actual $f_{u,DRG}$ was 10-fold lower than $f_{u,plasma}$ then it is conceivable that in DRG only the in vivo SPR IC50 was exceeded at Cmin. However, fractional binding was also evaluated in spinal cord and nerve homogenates with Q-1195, and these values were within 2-fold of the $f_{u,brain}$ value (data not shown), which is consistent with prior reports (Liu et al., 2018). Penetration into the tissue could also impact the local concentration; however, in vitro permeability assessment of Q-1195 in parental MDCK cell monolayers indicated favorable passive permeability ($P_{app} = 15 \mu m/s$, data not shown), which should not be limiting for tissue distribution. BH4 levels in healthy DRG were generally in the 20–30 ng/g range. SNL surgery caused an increase in BH4 that was greater than 100 ng/g. There was often a tight S.D. around the measured levels but a large range of values within a given experiment. This heterogeneity in response may also have impacted our ability to effectively reduce BH4. A 60% reduction in BH4 was observed following exposure to Q-1195, and it is possible that a larger magnitude of BH4 reduction is required to return animals to their tactile threshold baseline. It is worth noting that even for the positive control, gabapentin, there was a large variation in tactile threshold response (ranging from 2.32 to 15.0 g). A large sample number ($n = 10$ rats) for all SNL groups was used to gain a better estimate of the sample mean. Although not evaluated in this study, the selection of rodent strains and vendors can impact physiologic and behavioral endpoints in preclinical studies. Additionally, several neuropathic pain models are used in preclinical research; it is possible that varied results could be observed with SNL, spared nerve injury, or chronic constriction injury models. Behavioral models often exhibit large variability in response (pain models included) that present additional obstacles in the drug-discovery pipeline.

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