Brivaracetam attenuates pain behaviors in a murine model of neuropathic pain

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Abstract

Background: The antiseizure racetams may provide novel molecular insights into neuropathic pain due to their unique mechanism involving synaptic vesicle glycoprotein 2A. Anti-allodynic effects of levetiracetam have been shown in animal models of neuropathic pain. Here, we studied the effect of brivaracetam, which binds to synaptic vesicle glycoprotein 2A with 20-fold greater affinity, and has fewer off-target effects.

Methods: Mice underwent unilateral sciatic nerve cuffing and were evaluated for mechanical sensitivity using von Frey filaments. Pain behaviors were assessed with prophylactic treatment using levetiracetam (100 or 10 mg/kg) or brivaracetam (10 or 1 mg/kg) beginning after surgery and continuing for 21 days, or with therapeutic treatment using brivaracetam (10 or 1 mg/kg) beginning on day 14, after allodynia was established, and continuing for 28 or 63 days. Spinal cord tissues from the prophylaxis experiment with 10 mg/kg brivaracetam were examined for neuroinflammation (Iba1 and tumor necrosis factor), T-lymphocyte (CD3) infiltration, and synaptic vesicle glycoprotein 2A expression.

Results: When used prophylactically, levetiracetam, 100 mg/kg, and brivaracetam, 10 mg/kg, prevented the development of allodynia, with lower doses of each being less effective. When used therapeutically, brivaracetam extinguished allodynia, requiring 10 days with 10 mg/kg, and six weeks with 1 mg/kg. Brivaracetam was associated with reduced neuroinflammation and reduced T-lymphocyte infiltration in the dorsal horn. After sciatic nerve cuffing, synaptic vesicle glycoprotein 2A expression was identified in neurons, activated astrocytes, microglia/macrophages, and T lymphocytes in the dorsal horn. Conclusion: Synaptic vesicle glycoprotein 2A may represent a novel target for neuropathic pain. Brivaracetam may warrant study in humans with neuropathic pain due to peripheral nerve injury.

Keywords
Brivaracetam, levetiracetam, neuropathic pain, astrocyte, microglia, T lymphocyte, sciatic nerve cuff model, mouse

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Introduction

Neuropathic pain, which is characterized by dysesthesia, hyperalgesia, and allodynia, encompasses a wide range of heterogeneous conditions caused by lesions or diseases of the somatosensory nervous system, either at the periphery or centrally.1 Neuropathic pain that occurs after peripheral nerve injury arguably may be the most troubling type, since it is often refractory to treatment and thus is especially burdensome clinically.2 Numerous classes of drugs have been utilized for the treatment of neuropathic pain, including many anticonvulsants.3 Among anticonvulsants, the racetams used for seizure disorders may be of particular interest, due to their unique mechanism of action involving synaptic vesicle glycoprotein 2A (SV2A), a membrane glycoprotein localized to secretory vesicles in neurons, endocrine, and other cells.4–6

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Antihyperalgesic and anti-allodynic effects of the antiseizure racetam, levetiracetam (LEV), have been demonstrated in two animal models of chronic neuropathic pain: sciatic nerve (n.) constriction and streptozotocin-induced diabetes. Notably, LEV was more potent in the diabetic pain model (first active dose was 17 mg/kg) than in the sciatic n. constriction model (first active dose was 540 mg/kg). LEV has also been found to exert analgesic effects in nonneuropathic pain models such as postsurgical pain and inflammatory pain.

In humans, LEV was found to have analgesic effects in the electrical sural n. stimulation pain model and has shown efficacy in treating patients with trigeminal neuralgia, multiple sclerosis, and migraine. Notably, a recent Cochrane review failed to find the evidence of clinical efficacy for LEV based on six studies of 344 patients with six different types of neuropathic pain. However, the failures in these trials may have been due to the fact that numerous patients treated with LEV experienced adverse events that caused them to withdraw from the studies, precluding proper assessment of LEV's effects on neuropathic pain.

Another anticonvulsant of the racetam group, brivaracetam (BRV; UCB 34714), has gained recognition due to its greater antiseizure potency and reduced off-target effects, compared to LEV. BRV was first reported in 2004 as a structural derivative of LEV, differing from the parent compound by a single propyl group (Figure 1). BRV was identified during a drug-discovery program based on SV2A binding. Like LEV, BRV binds to SV2A but with 20-fold greater affinity. BRV, similar to LEV, enters into recycling synaptic vesicles and produces a frequency-dependent decrement in synaptic transmission at 100-fold lower concentrations than LEV. BRV is reported to be 10 times more potent as an antiseizure medication compared to LEV.

The effects of BRV have been reported in two rat models of neuropathic pain, albeit only as a meeting abstract. In both the streptozotocin-induced diabetes model and the sciatic n. constriction model, BRV (21 mg/kg) significantly increased the vocalization thresholds and completely reversed hyperalgesia. BRV (200 and 400 mg/day) was also tested in a clinical trial involving 152 subjects with postherpetic neuropathic pain, with the negative outcome of this study recently being made public.

Here, we sought to expand on the available literature regarding BRV in neuropathic pain, focusing on pain induced by peripheral nerve injury, because this type is relatively refractory to treatment compared to diabetic pain. We used a murine model of sustained neuropathic pain induced by sciatic n. cuffing to examine the effects of BRV when given either prophylactically or therapeutically, and we evaluated not only pain behaviors but also neuroinflammation in the spinal cord, an important factor in the initiation and maintenance of pain hypersensitivity.

Methods

Ethics statement

We certify that all applicable institutional and governmental regulations concerning the ethical use of animals were followed during the course of this research. Animal experiments were performed under a protocol approved by the Institutional Animal Care and Use Committee of the University of Maryland, Baltimore and in accordance with the relevant guidelines and regulations as stipulated in the National Research Council Publication, Guide for the Care and Use of Laboratory Animals. All efforts were made to minimize the number of animals used and their suffering.

Subjects and surgical procedure

Adult male C57BL/6 mice, approximately 22 to 25 g were obtained from Charles River (Frederick, MD, USA). Mice were given free access to food and water, except during behavioral testing. They were housed in plastic cages in specially constructed rooms with controlled humidity, exchange of air, and controlled lighting (12/12 h light/dark cycle).
For surgery, mice were anesthetized (100 mg/kg ketamine plus 10 mg/kg xylazine, intraperitoneal (IP)) and breathed room air spontaneously. Core temperature was maintained at 37°C using a heating pad (Deltaphase® Isothermal Pad, Braintree Scientific, Braintree, MA, USA). Hair was clipped from the right proximal lateral thigh, and the surgical site was prepared using iodine and alcohol. A sterile environment was maintained throughout the procedure. Lidocaine solution (2%) was injected prior to making an incision.

The procedure for sciatic n. cuffing was as previously described, with only minor modification. Using a surgical microscope, the common branch of the right sciatic n. was exposed by separating the muscles and the nerve by blunt dissection. After isolation, the nerve was gently stretched for 15 min by placing a 5-mm diameter plastic rod beneath it, which caused the nerve to blanch (see Figure 1 of Benbouzid et al.). A 2-mm long section of PE20 tubing, presplit and gas sterilized, was placed around the nerve. After the surgical procedure, mice were nursed on a heating pad to maintain temperature approximately 37°C until they emerged from anesthesia.

Exclusions
No mice became infected, required early euthanasia or died. No mice in the vehicle-treated groups failed to develop stable mechanical allodynia. There were no exclusions.

Treatments
LEV was obtained from West-Ward Pharmaceutical Corp. (now Hikma Pharmaceuticals USA Inc., Eatontown, NJ, USA). For doses of 100 and 10 mg/kg LEV, solutions of 500 or 50 mg, respectively, in 5 mL normal saline (NS) were administered via IP injection in a volume of 25 μL once daily. BRV was obtained from Toronto Research Chemicals (Ontario, Canada). For doses of 10 and 1 mg/kg BRV, solutions of 25 mg in 10 or 100 mL NS were administered via IP injection in a volume of 100 μL once daily. IP injections were performed using a 27-gauge needle with the depth of the injection limited to 3 mm by a sleeve of PE20 tubing placed over the needle.

Sample size calculation
For mechanical allodynia, we based our sample size calculation on a previous study that used the same model of neuropathic pain but tested a different drug. Values derived from Figure 2 of Yalcin et al. suggested an effect size (Cohen’s d) of ≈2, where $d = (M_1 - M_2)/SD_{\text{pooled}}$, $M_1$ and $M_2$ are the means, and $SD_{\text{pooled}} = [(SD_1^2 + SD_2^2)/2]^{1/2}$. Using the following assumptions:

two-tailed hypothesis, $\alpha$, 0.05; desired power, 80%; and $d$, 2.1, sample size calculation indicated a minimum sample size of five mice per group. This group size is similar to other reports using this model.

Experimental series
In series 1, 15 mice were randomly divided into three groups and were administered LEV (100 or 10 mg/kg) or vehicle (NS), with treatments beginning shortly after surgery, on postop day 0 (pod-0) and continuing daily until pod-21. In series 2, 15 mice were randomly divided into three groups and were administered BRV...
(10 or 1 mg/kg) or NS, with treatments beginning shortly after surgery, on postop day 0 (pod-0), and continuing daily until pod-21; tissues from these mice (BRV, 10 mg/kg and NS) were used for immunohistochemistry. In series 3, 10 mice were randomly divided into two groups and were administered BRV (10 mg/kg) or NS, with treatments beginning on pod-14 and continuing daily until pod-28. In series 4, 12 mice were randomly divided into two groups and were administered BRV (1 mg/kg) or NS, with treatments beginning on pod-14 and continuing daily until pod-63. In all cases, the non-injured hind paw served as a control.

**Mechanical allodynia**

Outcomes were assessed by investigators blinded to treatment group. Mice were handled two to three times weekly for acclimatization to handlers. Sensory testing was performed before surgery and at three- to five-day intervals thereafter. Mice were placed in elevated Perspex cages with a wire mesh floor (15 × 10 × 10 cm) (ITCC Life Science, Woodland Hill, CA, USA) and were acclimatized for 30 min prior to testing. The paw withdrawal threshold to mechanical stimulation of both the ipsilateral and contralateral hind paws was measured using a series of von Frey filaments, which exerted forces ranging from 0.16 to 4 g (North Coast Medical, San Jose, CA, USA). The von Frey filaments were pressed perpendicularly onto the plantar surface of the hind paw for 2 s (four times for each filament), and a positive response was noted if there was a sharp flinching of the hind paw. The “up-down method” was used to determine the withdrawal threshold.

**Thermal hyperalgesia**

Outcomes were assessed by investigators blinded to treatment group. Thermal sensitivity was assessed in mice from series 2 at 14 days after sciatic n. cuffing using the Hargreaves method and a Hargreaves-type apparatus (Plantar Test Analgesia Meter, ITCC Life Science). An unrestrained mouse was placed in a Perspex enclosure on the top of a glass pane. An infrared generator placed below glass pane was aimed at the plantar surface of the hind paw, and the time to withdrawal was recorded automatically via an optical sensor. Paw withdrawal latency was calculated as the mean of three to five different measurements taken at 15-min intervals.

**Immunoblot for validation of anti-SV2A antibody**

To validate the rabbit anti-SV2A antibody used for immunohistochemistry (cat#TA322365; Origene, Rockville, MD, USA), we performed an immunoblot of lysate from mouse brain, from an immortalized rat astrocyte cell line (DI TNC1; ATCC, Gaithersburg, MD, USA) and from an immortalized human T-lymphocyte cell line (TIB-152; ATCC), using standard methods as we described.

**Immunohistochemistry and quantification of specific labeling**

Under deep anesthesia, mice were euthanized, underwent trans-cardiac perfusion with NS (15 mL) followed by 10% neutral buffered formalin (15 mL). Spinal cord tissues at spine segments L1 to L5 were harvested and postfixed. Tissues were cryoprotected with 30% sucrose, frozen in optimal cutting temperature (OCT), and cryosectioned (10 μm).

Immunohistochemistry was performed as described. Sections were incubated at 4°C overnight with primary antibodies, including rabbit anti-Iba1 (1:200; cat#019–19741; Wako, Osaka, Japan), goat antitumor necrosis factor (TNF) (1:200; cat#sc1350 (N-19); Santa Cruz Biotechnology, Sant Cruz, CA, USA), mouse antiglial fibrillary acidic protein (GFAP) (1:300; cat#C9205; Sigma, St. Louis, MO, USA), rabbit anti-CD3 (1:100; cat# AB5690; Abcam, Cambridge, UK), mouse anti-CD45 (1:50; cat# 05–1410; EMD Millipore, Temecula, CA, USA), and rabbit anti-SV2A (1:50; cat#TA322365; Origene).

After several rinses in phosphate-buffered saline, sections were incubated with species-appropriate fluorescent secondary antibodies (Alexa Fluor 488 and 555, Molecular Probes; Invitrogen, Carlsbad, CA, USA) for 1 h at room temperature. Controls included the omission of primary antibodies.

Unbiased measurements of specific labeling within regions of interest (ROI) were obtained using NIS-Elements AR software (Nikon Instruments, Melville, NY, USA) from sections immunolabeled as a single batch. All images for a given signal were captured using uniform parameters of magnification, area, exposure, and gain. Segmentation analysis was performed by computing a histogram of pixel intensity for a particular ROI, and pixels were classified as having specific labeling based on signal intensity greater than two times that of background. The area occupied by pixels with specific labeling was used to determine the percentage area in the ROI with specific labeling (% ROI). For Iba1 and TNF, the ROI was a rectangle, 500 × 400 μm, positioned at the dorsal edge of the dorsal horn. For CD3, individual CD3+ cells were counted manually as described in two distinct areas: the dorsal horn and the remainder of the gray matter, both ipsilateral and contralateral.

**Statistics**

Nominal data are presented as mean ± standard error. Nominal data were analyzed using a t test or analysis of
variance with post hoc Bonferroni correction, as appropriate. Statistical tests were performed using Origin Pro (V8; OriginLab, North Hampton, MA, USA). Significance was assumed if $P < 0.05$.

**Results**

**LEV and BRV prophylaxis**

LEV (540 mg/kg) was tested previously in a rat model with chronic constriction of the sciatic n. Here, we tested LEV (100 or 10 mg/kg) in a mouse model with sciatic n. cuffing. LEV or vehicle was administered beginning shortly after sciatic n. cuffing and daily thereafter. Von Frey filaments were used to assess ipsilateral and contralateral hindpaw withdrawal thresholds at several day intervals up to 21 days. In mice that were administered vehicle, sciatic n. cuffing gave rise to mechanical allodynia involving the ipsilateral hindpaw that developed over the course of 7 days and that persisted for the remainder of the 21-day duration of the experiment (Figure 2(a)). Mice that were administered LEV, 10 mg/kg, exhibit allodynia that was significantly less severe than vehicle-treated animals. However, in mice that were administered LEV, 100 mg/kg, hypersensitivity of the ipsilateral hindpaw to mechanical stimuli failed to develop, and withdrawal thresholds were not different from those of the contralateral (uninjured) hindpaw (Figure 2(a)).

We repeated this experiment, except that LEV was replaced by BRV (10 or 1 mg/kg daily). Again, in control mice that were administered vehicle, sciatic n. cuffing gave rise to mechanical allodynia involving the ipsilateral hindpaw that persisted (Figure 2(b)). Mice that were administered BRV, 1 mg/kg, exhibit allodynia that was significantly less severe than vehicle-treated animals. However, in mice that were administered BRV (10 mg/kg daily) beginning at the time of sciatic n. cuffing, hypersensitivity of the ipsilateral hindpaw to mechanical stimuli failed to develop, and withdrawal thresholds were not different from those of the contralateral (uninjured) hindpaw (Figure 2(b)).

We also tested thermal sensitivity in the mice that had received BRV (10 mg/kg daily) versus vehicle. Thermal sensitivity was assessed on day 14 using Hargreaves test. Compared to the contralateral hindpaw, the ipsilateral hindpaw of vehicle-treated mice exhibited significant thermal hyperalgesia (Figure 2(c)). By contrast, BRV-treated mice exhibited ipsilateral thermal sensitivity that was not different from the contralateral (uninjured) hindpaw (Figure 2(c)).

**Neuroinflammation and T-cell infiltration**

Neuroinflammation with microglial activation is a key component of neuropathic pain and is reportedly ameliorated by LEV. We studied tissues from mice in the foregoing experiment administered BRV (10 mg/kg daily) versus vehicle. The involved spinal cord segment was immunolabeled for Iba1 and colabeled for TNF. In the dorsal horn of uninjured controls, Iba1+ microglia were sparsely distributed and exhibited a ramified, nonreactive morphology, consistent with a quiescent phenotype; minimal immunoreactivity for TNF was evident (Figure 3(a) and (d)). After sciatic n. cuffing, numerous Iba1+ cells with a plump, activated morphology were evident in the ipsilateral dorsal horn and to a lesser extent in the contralateral dorsal horn, and immunoreactivity for TNF was greatly increased (Figure 3(b) and (e)). In mice with sciatic n. cuffing treated with BRV, Iba1+ cells were less prominent, and these cells exhibited less-plump morphologies; TNF immunoreactivity was also less prominent (Figure 3(c) and (f)). Quantification of Iba1 and TNF immunoreactivity confirmed a significant increases in vehicle-treated mice with sciatic n. cuffing compared to uninjured controls and...
significant decreases in mice with sciatic n. cuffing treated with BRV, both ipsilateral and contralateral (Figure 3(g)).

T lymphocytes (T cells) comprise an important component of the inflammatory response to nerve injury.42,44–46 Immunolabeling for CD3 showed small clusters of cells at the involved segment, located predominantly in the ipsilateral dorsal horn, often at the lateral gray-white junction (DH-LGWJ) (Figure 4(a) to (c)), as well as near the central canal (Figure 4(d)).42 Occasionally, a few CD3+ cells could also be identified in the mirror location contralaterally. Counts of CD3+ cells showed a significant increase in the dorsal horn as well as outside the dorsal horn of mice with sciatic n. cuffing treated with vehicle, compared to uninjured controls (Figure 4(e) and (f)). Treatment with BRV resulted in significant reductions in CD3+ cells in all regions (Figure 4(e) and (f)).

**SV2A expression in the affected spinal cord**

A likely site of action of LEV and BRV in neuropathic pain due to sciatic n. cuffing is the spinal cord, where SV2A expression was reported previously using autoradiography and immunoblot.27 Here, we sought to determine which cell type(s) in the spinal cord express SV2A. Spinal cord tissues were immunolabeled using an anti-SV2A antibody that we independently validated (Figure 5). In uninjured controls, SV2A was identified almost exclusively in gray matter neuropil and neurons (not shown), as reported.27 In mice with sciatic n. cuffing, coronal sections of the involved segment showed robust labeling in gray matter (Figure 5(a) and (b)), with occasional, sporadic immunoreactivity in white matter subpial astrocytes. Most of the SV2A immunoreactivity in gray matter was in neuropil and neurons, the latter easily identified in the ventral horn by their large distinct perikarya (Figure 5(a) and (b)). Double immunolabeling for SV2A and GFAP showed that activated astrocytes in the ipsilateral dorsal horn also expressed SV2A (Figure 5(c)). In addition, double immunolabeling for SV2A and Iba1 or CD45 showed that small numbers of microglia/macrophages as well as leukocytes located in the ipsilateral dorsal horn, including the DH-LGWJ, also expressed SV2A (Figure 5(d) and (e)). The CD45+ leukocytes that expressed SV2A were the same cells that were CD3+, consistent with previous reports of SV2A expression by T lymphocytes.47

**BRV treatment**

To examine a clinically relevant scenario, we studied the effect of BRV when administered beginning after allodynia was fully established. Mice underwent sciatic n. cuffing and were allowed to develop mechanical allodynia over the course of two weeks. They then were randomly assigned to one of the four groups: two vehicle control groups, or BRV at two doses, 10 mg/kg or 1 mg/kg daily. Over the course of the next 10 days, mice receiving 10 mg/kg daily gradually reverted to mechanical sensitivities equivalent to the contralateral hindpaw, whereas those receiving vehicle exhibited persistent allodynia (Figure 6(a)). Similarly, mice receiving 1 mg/kg exhibited a slow recovery to normal sensitivities, although at this dose, the recovery required six weeks (Figure 6(b)). Note that vehicle-treated mice in the last experiment showed no extinction of symptoms, continuing to exhibit mechanical allodynia for the full nine-weeks of the experiment.

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**Figure 4.** Treatment with BRV reduces spinal cord infiltration of CD3+ cells in the murine sciatic n. cuff model. Low-magnification (a) and high-magnification (b and c) views of the ipsilateral dorsal horn of two mice with sciatic n. cuffing-administered vehicle, immunolabeled for CD3; note several small CD3+ cells in the dorsal most part of the dorsal horn (red arrows), and several larger CD3+ cells at the dorsal horn lateral gray-white junction (red asterisk), the latter shown at high magnification in (c). (d) High-magnification view of the central canal, identified by the circle of 4’,6-diamidino-2-phenylindole (blue) positive ependymal cells, with nearby clusters of CD3+ cells (green); bars: 250 μm in (a), and 100 μm in (b) to (d). (e and f) Quantification of CD3+ cells in ipsilateral (I) and contralateral (C) areas, the DH and the remainder of the gray matter (non-DH), of mice with sciatic n. cuffing-administered vehicle (Veh) or BRV (10 mg/kg); LWM: lateral white matter; five mice per group; *P < 0.05; **P < 0.01. BRV: brivaracetam; DH: dorsal horn.
Discussion

The principal findings of the present study are that (i) both BRV and LEV are effective in reducing neuropathic pain behaviors in the murine sciatic n. cuff model; (ii) the salutary effects of BRV are observed with a dose that is 10 times less than that of LEV, similar to findings in seizure models, and consistent with a mechanism of action involving SV2A; (iii) the salutary effects of BRV on neuropathic pain behaviors correlate with reduced neuroinflammation in the spinal cord; (iv) BRV exhibits beneficial effects when administered both prophylactically, at the time of sciatic n. cuffing, before the onset of neuropathic pain, and later, after symptoms have fully developed, underscoring its potential for translation to the pain clinic. An interesting, but unexplained observation is the length of time required for BRV to reverse allodynia after it is established—two and six weeks with 10 and 1 mg/kg, respectively.

Neuronal hyperexcitability is a critical element in neuropathic pain, and for this reason, most current drug treatments for neuropathic pain are directed toward decreasing neuronal excitability. However, neuroinflammation linked to glial cell activation is also recognized to play a prominent role in the initiation and maintenance of pain hypersensitivity. Nonneuronal cells such as immune cells (macrophages and lymphocytes) and glial cells (Schwann cells and satellite cells in the peripheral...
nervous system (PNS) and astrocytes and microglia in the central nervous system (CNS) play an important role in the induction and maintenance of neuropathic pain. Injury-induced inflammation at the site of the damaged or affected nerve(s) precedes microglial activation in the dorsal horns of the spinal cord. Under chronic pain conditions, neuroinflammation is characterized by infiltration of immune cells in the dorsal root ganglion and sequential activation of microglia and astrocytes in the spinal cord and brain, with subsequent release of numerous pro-inflammatory cytokines and chemokines.

Our data showing a reduced neuroinflammatory response within the dorsal horn with BRV is in keeping with the concept of a key role for neuroinflammation in neuropathic pain. We showed that BRV significantly reduced microglial activation, TNF expression, and leukocyte infiltration into the dorsal horn, in conjunction with reduced neuropathic pain behaviors. Our observations here with BRV accord with a previous report on LEV, which is also effective in preclinical models of neuropathic pain and also reduces neuroinflammation within the spinal cord.19

Considerable work indicates that LEV and BRV act, at least in part, via a specific mechanism involving SV2A.51 However, apart from SV2A, LEV has additional mechanisms of action, including inhibition of voltage-gated K+ and Ca2+ channels, inhibition of 1,4,5-triphosphate-mediated release of intracellular Ca2+, and interactions with multiple excitatory and inhibitory ligand-gated ion channels.52–58 By contrast, BRV may be somewhat more selective. A single report suggests that BRV inhibits Na+ current in cultured rat cortical neurons,59 but other studies indicate that BRV does not modulate high- and low-voltage-activated Ca2+ currents, voltage-gated delayed rectifier K+ currents, and persistent voltage-gated Na+ currents, and that BRV is devoid of any direct effect on currents gated by γ-aminobutyric acidergic type A, glycine, kainate, N-methyl-D-aspartate, and γ-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid.60–62

Given that LEV and BRV overlap strongly in terms of SV2A binding, but not in terms of actions on ion channels or receptors, it seems plausible that the mechanism for the similar salutary effects of LEV and BRV on pain behaviors involves SV2A. The 10-fold higher potency that we observed here with respect to antiallodynic activity, which is similar to its relative efficacy with respect to antiseizure activity,24 is in keeping with 20-fold greater binding affinity for SV2A by BRV28 and is in keeping with the novel hypothesis that SV2A is the relevant target of the anticonvulsant racetams in neuropathic pain.

Although SV2A may be a target of the drug, the specific role of SV2A in neuropathic pain is enigmatic. SV2A is an integral constituent of synaptic vesicle membranes in presynaptic terminals and has been implicated in mechanisms that control the efficiency of vesicle release as well as upstream vesicle-trafficking mechanisms, accounting for the ability of the anticonvulsant racetams to reduce neuronal excitation.5,6 SV2A is an essential protein for maintaining Ca2+-regulated exocytosis not only at the synapse but in neuroendocrine and other cells as well.

Of the three known SV2 paralogs, SV2A is the only member ubiquitously expressed in the adult brain.5,6,63 SV2A is also expressed in spinal cord, predominantly in gray matter, as previously reported27 and as shown here. SV2A is also found in neuroendocrine cells and at neuromuscular junctions.48,64,65 However, the expression of SV2A by other CNS or PNS cells has not been well studied. In culture, astrocytes reportedly upregulate SV2A,66 and in cultured astrocytes, LEV inhibits the release of glutamate67 and exhibits anti-inflammatory properties.68,69 Reports indicate that acutely isolated astrocytes as well as astrocytes in situ do not express SV2A,66 whereas here, we found that activated astrocytes in the involved dorsal horn, as well as an astrocyte cell line, express SV2A. Our immunolabeling experiments also revealed the expression of SV2A by Iba1+ microglia/macrophages. Thus, regarding involvement of CNS cell types, neurons, astrocytes, and microglia in the spinal cord may be plausible targets of the racetams in neuropathic pain.

Apart from the nervous system, SV2A mRNA and protein have been identified in T lymphocytes47,70 and in antral mucosal cells,64 and in these cells, like in neurons, Ca2+-regulated exocytosis (in this case, degranulation) is inhibited by LEV.47,71 Here, we found that CD3+ T lymphocytes in situ, as well as a human T-lymphocyte cell line, express SV2A, with counts of T lymphocytes in situ significantly reduced by BRV. T lymphocytes are known to contribute to neuropathic pain induced by peripheral nerve injury. CD4+ and CD8+ T lymphocytes infiltrate the injured sciatic n. after both a local inflammatory insult72 and chronic constriction injury (CCI),34,46 the latter similar to the sciatic n. cuff model used here. Following CCI of the sciatic n., congenitally athymic nude rats,34 as well as RAG-1 knockout mice,46 both of which lack functional T lymphocytes, develop reduced mechanical allodynia and thermal hyperalgesia. In murine nerve transection-induced neuropathic pain models, T lymphocytes infiltrate into the dorsal horn of the spinal cord, and in these models as well, T lymphocyte deficiency (RAG-1 or CD4 knockout) is partially protective.42,73 Thus, both peripheral and central T lymphocyte may be plausible targets of the racetams in neuropathic pain.

A possible peripheral action of the racetams is further supported by other lines of evidence. Ipsilateral but not contralateral intraplantar injection of LEV in a model of
localized inflammation (intraplantar carrageenan) produces local peripheral antihyperalgesic and antiedematous effects. Botulinum neurotoxins, which enter neurons by binding to SV2A, are injected peripherally as a third-line treatment for neuropathic pain in humans. In a model of diabetic neuropathy, untreated mice with allodynia showed degeneration and vacuolization in the sciatic n. along with central neuroinflammation, whereas in mice treated with LEV, allodynia was reduced, nerves showed minimal histopathological changes, and spinal cord microgliosis and astrocytosis were reduced. Thus, the anticonvulsant racetams, LEV and BRV, may be acting synergistically via SV2A both centrally in neurons, astrocytes, microglia, and T lymphocytes and peripherally in T lymphocytes and sensory neurons.

Finally, the mechanism for the salutary effects of LEV and BRV on neuropathic pain may extend beyond SV2A. The racetams are a broad class of drugs that share a 2-pyrrolidinone nucleus but may otherwise differ greatly in structure and biological actions. Notably, other racetams, including piracetam, nefiracetam, and dimiracetam, have been found to be effective in neuropathic pain models, generally at doses comparable to those used with LEV. In some cases (excluding piracetam), their chemical structures are quite different from those of LEV and BRV (Figure 1). LEV, BRV, and piracetam are known to bind to SV2A, but to our knowledge, SV2A binding has not been demonstrated for dimiracetam and nefiracetam, although the latter is reported to have antiseizure properties. At present, a unifying molecular mechanism is lacking to explain how such structurally diverse molecules as the racetams depicted in Figure 1 can be effective in neuropathic pain.

This study has limitations. Among them, we studied only a single model of pain in one sex of a single species, and most of our efficacy evaluations focused on mechanical allodynia. Future studies will be needed to broaden our experimental approach, to include other models of neuropathic pain, nonreflexive pain behaviors, both sexes, and other species. Foremost, we implicated SV2A in the neurobiology of neuropathic pain by relying on pharmacological rather than molecular experiments. Although BRV is highly selective for SV2A, molecular approaches may now be within reach to address this shortcoming.

**Conclusion**

Neuropathic pain remains a major public health problem that is magnified by the prevalence of opioid use disorder. Its pathogenesis remains incompletely understood, and major challenges remain to discover novel drugs with therapeutic efficacy that will be well tolerated, have minimal side effects, and be devoid of addictive potential. Here, we report that BRV is highly effective in reducing neuropathic pain behaviors in a murine sciatic n. injury model when administered both prophylactically, before the onset of neuropathic pain, and later, after symptoms have fully developed. Attenuation of pain behaviors by BRV was found to correlate with reduced neuroinflammation in the spinal cord. When used therapeutically, especially at low doses, daily administration of BRV for several weeks was required to achieve an optimal benefit. Compared to LEV, BRV’s greater potency, fewer off-target effects, and more benign side-effect profile suggest that it may be an attractive candidate drug for the treatment of some forms neuropathic pain due to peripheral nerve injury.

**Author’s Contributions**

Study conceptualization, ST; study design and oversight, OT, VG, and JMS; surgeries, OT; data collection for neurofunctional tests, ST, MS, CG, and OT; immunohistochemistry, SI and VG; funding procurement, CS and JMS; first draft of the manuscript, JMS; critical contributions to the manuscript, ST, CS, VG, and JMS. All authors have read and approved the final manuscript.

**Declaration of Conflicting Interests**

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