Amperometric Measurement of Glutamate Release Modulation by Gabapentin and Pregabalin in Rat Neocortical Slices: Role of Voltage-Sensitive Ca^{2+} \alpha_2\delta-1 Subunit

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ABSTRACT

Gabapentin (GBP; Neurontin) and pregabalin (PGB; Lyrica, S-(+)-3-isobutyrlgaba) are used clinically to treat several disorders associated with excessive or inappropriate excitability, including epilepsy; pain from diabetic neuropathy, postherpetic neuralgia, and fibromyalgia; and generalized anxiety disorder. The molecular basis for these drugs’ therapeutic effects are believed to involve the interaction with the auxiliary \alpha_2\delta subunit of voltage-sensitive Ca^{2+} channel (VSCC) translating into a modulation of pathological neurotransmitter release. Glutamate as the primary excitatory neurotransmitter in the mammalian central nervous system contributes, under conditions of excessive glutamate release, to neurological and psychiatric disorders. This study used enzyme-based microelectrode arrays to directly measure extracellular glutamate release in rat neocortical slices and determine the modulation of this release by GBP and PGB. Both drugs attenuated K^{+}-evoked glutamate release without affecting basal glutamate levels. PGB (0.1–100 \mu M) exhibited concentration-dependent inhibition of K^{+}-evoked glutamate release with an IC_{50} value of 5.3 \mu M. R-(−)-3-Isobutyrylgaba, the enantiomer of PGB, did not significantly reduce K^{+}-evoked glutamate release. The decrease of K^{+}-evoked glutamate release by PGB was blocked by the L-amino acid L-isoleucine, a potential endogenous ligand of the \alpha_2\delta subunit. In neocortical slices from transgenic mice having a point mutation (i.e., R217A) of the \alpha_2\delta-1 (subtype) subunit of VSCC, PGB did not affect K^{+}-evoked glutamate release yet inhibited this release in wild-type mice. The results show that GBP and PGB attenuated stimulus-evoked glutamate release in rodent neocortical slices and that the \alpha_2\delta-1 subunit of VSCC appears to mediate this effect.

Introduction

Several neurological and psychiatric disorders characterized by excessive or dysfunctional neurotransmitter release are routinely treated with gabapentin [GBP; Neurontin, 1-(aminomethyl)cyclohexanecarboxylic acid] and pregabalin [PGB; Lyrica, S-(+)-3-isobutyrlgaba, S-(+)-4-amino-3-(2-methylpropyl)butanoic acid] (Dooley et al., 2007). Although multiple mechanisms of action have been proposed historically to account for the preclinical and clinical profiles of these drugs, there is increasing evidence for a significant role of the auxiliary \alpha_2\delta subunit of voltage-sensitive Ca^{2+} channel (VSCC) (Taylor et al., 1998, 2007). The binding of these ligands to the \alpha_2\delta subunit is believed to be the source of their efficacy in treating epilepsy; pain from diabetic neuropathy, postherpetic neuralgia, and fibromyalgia; and generalized anxiety disorder. With the recent availability of transgenic mice with point mutations of the \alpha_2\delta subunit (i.e., \alpha_2\delta-1 and \alpha_2\delta-2 subtypes) (Bian et al., 2006, 2008), preclinical experiments can be designed to test for altered neurochemical and behavioral effects of GBP and PGB (Field et al., 2006).

In the present study, we used enzyme-based microelectrode arrays (MEAs) that have micrometer-sized platinum recording sites with sampling rates of >1 Hz. These MEAs were developed in response to the limitations of other techniques or devices used to measure neurotransmitters (e.g., microdialysis/perfusate sampling coupled to high-performance liquid chromatography) (Barnes et al., 1988; Shinohara et al., 1998; Burmeister et al., 2000). A drawback of microdialysis or perfusate sampling techniques is that they sample from a large area (Borland et al., 2005) and at relatively slow (minutes) sampling rates.
rates. Given the rapid nature of neurotransmission of chemical messengers such as glutamate, faster sampling rates and smaller sampling areas should be beneficial. Enzyme-coated MEAs have been characterized extensively in the brains of anesthetized and behaving animals to measure glutamate (Burmeister et al., 2002; Binns et al., 2005; Day et al., 2006; Nickell et al., 2006; Rutherford et al., 2007; Parikh et al., 2010) but not in brain slices.

We used these MEAs to directly measure K⁺-evoked (extracellular) glutamate release in rat neocortical slices and to determine the modulation of this release by GBP and PGB. As the primary excitatory neurotransmitter in the mammalian central nervous system, glutamate often has been associated with a variety of pathological conditions (Meldrum, 2000), several of which are responsive to α₂δ ligands such as PGB. A reduction of excessive glutamate release by α₂δ ligands conceivably translates into clinically relevant therapeutic effects, especially considering the experimental evidence supporting a relationship between α₂δ-subunit binding and the modulation of processes subserving neurotransmitter release (Dooley et al., 2007).

An additional aspect of this study was to assess the effects of PGB on K⁺-evoked (extracellular) glutamate release in neocortical slices from wild-type and α₂δ-1 mutant mice. The α₂δ-1 transgenic mice have a point mutation (namely, R217A) that markedly reduces [³²H]GBP and [³²H]PGB binding in central nervous system regions (e.g., neocortex) known to preferentially express the α₂δ-1 protein (Bian et al., 2006; Field et al., 2006).

Materials and Methods

Animals. Male rats [Sprague-Dawley, 2–8 weeks old; Harlan, Indianapolis, IN] and male mice [wild-type and mutant α₂δ-1 R217A (Bian et al., 2006; Field et al., 2006), 2–5 months old; Charles River Laboratories, Inc., Wilmington, MA] were housed in an Association for Assessment and Accreditation of Laboratory Animal Care-accredited facility according to the standards outlined in the Guide for the Care and Use of Laboratory Animals (Institute of Laboratory Animal Resources, 1996).

Animals were under a 12-h light/dark cycle, had ad libitum access to food and water, and were maintained for a minimum of 5 days before euthanasia by decapitation. The brains were removed by blunt dissection and placed in ice-cold buffer until slice preparation. The glutamate recordings occurred during the light phase of the light/dark cycle. All of the experimental protocols were approved by the Animal Care and Use Committee of the University of Kentucky.

Glutamate Release Measurements. Neocortical slices from rats and mice were prepared using standard protocols (Hascup et al., 2002). In brief, coronal slices (0.35–0.4 mm in thickness), including the frontal and parietal areas exhibiting relatively high [³²H]GBP and [³²H]PGB binding (Bian et al., 2006), were maintained for at least 1 h at room temperature in artificial cerebrospinal fluid (aCSF; 124 mM NaCl, 5 mM KCl, 2.5 mM CaCl₂, 1.5 mM MgCl₂, 26 mM NaHCO₃, 1.4 mM NaH₂PO₄, and 10 mM d-glucose, saturated with 95% O₂/5% CO₂, pH 7.2–7.4) before the start of an experiment. The slices were transferred to immersion-style chambers (i.e., one slice per chamber) and superfused at a rate of 1.5 to 2.0 ml/min with aCSF (31–33°C). Each chamber was fitted with an Ag/AgCl reference electrode.

Ceramic-based MEAs (four platinum sites in a row, 50 μm each) were assembled, coated with Nafion, and subsequently coated with three layers of a 1% glutamate oxidase (Associates of Cape Cod, East Falmouth, MA) 1% bovine serum albumin/0.125% glutaraldehyde enzyme solution. Coated MEAs were allowed to cure for a minimum of 2 days before use. Enzyme-based MEAs measure glutamate through the enzymatic breakdown of glutamate to yield a reporter molecule of hydrogen peroxide that is subsequently oxidized on the platinum recording surface to yield an oxidation current. The MEAs then were calibrated (in vitro) with glutamate in a phosphate-buffered solution (pH 7.4) at 31–34°C to 1) generate a standard response curve (sensitivity > 2 pA/μM); 2) determine the limit of detection (≥3 times the signal-to-noise ratio; < 2.0 μM); and 3) assess the selectivity for glutamate relative to an endogenous electroactive compound, ascorbic acid (>30:1). A MEA or MEA/micropipette assembly was lowered into the neocortical slice, and extracellular glutamate levels were measured once basal glutamate levels stabilized for at least 10 min.

Test substances were delivered through the superfusion system for a minimum of 15 min before slice stimulation unless stated otherwise. The slices were stimulated twice (S₁, S₂) with high K⁺ by one of two methods to evoke glutamate release: 1) direct, local application of 70 mM K⁺ solution [70 mM KCl, 79 mM NaCl, and 2.5 mM CaCl₂; pH 7.0–7.4] to depolarize the local gluta
maticergic network via pressure ejection or 2) superfusion of 70 mM K⁺ [i.e., an increase of KCl in aCSF with a corresponding decrease of NaCl (59 mM) to maintain iso-osmolarity] to depolarize the whole slice. Slices were allowed to recover for a minimum of 20 min between stimulations. For local application, glass micropipettes (inside tip diameter of 10–15 μm) were formed from stock (1 mm o.d., 0.58 mm i.d.; A-M Systems, Everett, WA) and attached, and the tip was centered over the MEA recording site at a tip-to-tip distance of 70 to 110 μm. The 70 mM K⁺ solution was applied at 1-min intervals until at least two to five reproducible glutamate responses were recorded. Delivery of solution volumes (i.e., 12.5–400 nl over 0.1–3.0 s) was controlled by a pressure-ejection system (2–12 psi; Picospritzer II, Parker Hannifin, Mayfield Heights, OH) and monitored using a stereo microscope fitted with a reticule (Gerhardt and Palmer, 1987). Extracellular glutamate levels were measured at 1 Hz using constant potential amperometry (+0.7 V versus Ag/AgCl reference) controlled by a FAST16 electrochemical recording system (Quanteon, LLC, Nicholasville, KY) and analyzed offline by customized Excel-based software.

Calculations and Statistics. Glutamate release amplitudes were calculated from the difference between maximal K⁺-evoked glutamate release values and basal values. Values given are X ± S.E. (n ≥ 6). In one set of experiments with PGB, a concentration-effect curve with the corresponding IC₅₀ value was calculated by nonlinear regression (Prism 4.0; GraphPad Software Inc., San Diego, CA). If appropriate, the results were analyzed using the t statistic for group means or one- or two-way analysis of variance followed by post hoc comparisons using Dunnett’s or Bonferroni multiple comparison statistic (InStat 3.0; GraphPad Software Inc.). The minimal level of significance was p ≤ 0.05 (two-tail criterion).

Materials. Substances were either commercially available (Sigma-Aldrich, St. Louis, MO) or donated [i.e., GBP, PGB, and R-(-)-3-isobutylglycine (R-IBG) (Pfizer, Inc., New York, NY)]. Test compounds were dissolved directly in aCSF.

Results

The effects of GBP and PGB on resting glutamate levels were evaluated in initial experiments. Neither drug (0.1–100 μM) altered basal glutamate levels in neocortical slices (data not shown).

In the brains of anesthetized animals, delivery of high K⁺ to stimulate glutamate release has been performed with local, pressure-ejected administration (Burmeister et al., 2002; Day et al., 2006; Quintero et al., 2007; Stephens et al., 2009). Meanwhile, studies in brain slices permit the use of two methods of delivering high K⁺ solutions to activate neural networks: local delivery and superfusion. Local stimulation, using pressure delivery, produces a comparable stimulus to...
those used in previous studies with anesthetized animals. An additional benefit of brain slice recordings is the flexibility to also use superfusion of high K⁺ to evoke release. This sustained depolarization resembles prolonged or excessive excitability—a condition that characterizes some neurological disorders such as anxiety. We used both stimulation methods here to characterize the effects of αδ subunit ligands and to compare glutamate measurements with MEAs to those of previous studies.

The repeated pressure-ejection delivery of 70 mM K⁺ solution yielded similar-sized glutamate signals with a mean amplitude of 3.9 ± 0.8 μM (Fig. 1). In the presence of GBP (100 μM), the mean amplitude was decreased significantly by 46% to 2.1 ± 0.7 μM (Fig. 1).

Because GBP was confirmed to modulate K⁺-evoked glutamate release, the more recently developed αδ ligand PGB was chosen for testing in additional experiments. PGB (100 μM) attenuated pressure-ejection delivery of 70 mM K⁺ solution (5.7 ± 1.5 μM glutamate, pre-PGB versus 1.7 ± 1.2 μM glutamate, post-PGB; t(4) = 3.59, p = 0.023). We then transitioned to a paradigm of using repeated stimulation with superfused 70 mM K⁺ (S₃, S₄). The S₃/S₁ ratio of control glutamate signals in rat neocortical slices was 0.97 (Figs. 2A and 3A); this ratio was markedly reduced by 78% to 0.21 by GBP (100 μM) (Figs. 2B and 3A). Other S₃/S₁ ratios for PGB include 0.80 (nonsignificant, 14% inhibition) at 0.1 μM, 0.60 (nonsignificant, 38% inhibition) at 1 μM, and 0.44 (54% inhibition) at 10 μM (Fig. 3A); an IC₅₀ value of 5.3 μM was determined from the concentration-effect relationship (Fig. 3B). The enantiomer of PGB, R-IBG (100 μM), gave an S₃/S₁ ratio of 0.70 (nonsignificant, 28% inhibition), contrasting sharply with the effect (78% inhibition) of an equimolar concentration of PGB (Fig. 3A).

PGB and the endogenous amino acid L-isoleucine are sub-

![Fig. 1. Effect of GBP (100 μM) on K⁺-evoked glutamate release from rat neocortical slices. A, glutamate release evoked by repeated pressure-ejection delivery of 70 mM K⁺ solution (arrowheads on abscissa) in the absence (first three traces) and presence of GBP (last three traces). B, amplitude of K⁺-evoked glutamate release (as derived from A) was decreased by GBP. Values given are X ± S.E. (n = 6). The paired t statistic gave t(5) = 2.930 (p = 0.0326). A significant difference from the control value is indicated by an asterisk (*, p < 0.05).](image)

![Fig. 2. Effect of PGB (100 μM) on K⁺-evoked glutamate release in rat neocortical slices. A, detection of glutamate release by MEAs after repeat superfusion with 70 mM K⁺ (arrowheads; S₁, S₂) for 50 s. B, PGB (closed bar), present 15 min before S₂, attenuated glutamate release.](image)

strates for the system L-amino acid transporter, and both have a similar nanomolar affinity for the αδ ligand binding site on the αδ subunit. The S₂/S₁ ratio associated with the K⁺-evoked glutamate signals in the presence of L-isoleucine (100 μM) was 0.84 (nonsignificant, 13% inhibition), yet this compound reduced the effect (78% inhibition) of PGB (100 μM) as indicated by the S₂/S₁ ratio of 0.59 (nonsignificant, 39% inhibition) (Fig. 3A).

An action of GBP and PGB at the αδ-1 subtype rather than the αδ-2 subtype has been proposed to account for the therapeutic effects of these drugs (Bian et al., 2006, 2008; Field et al., 2006). With neocortical slices from the wild-type and αδ-1 transgenic mice (Bian et al., 2006; Field et al., 2006), PGB (100 μM) significantly decreased K⁺-evoked glutamate signals (Fig. 4, A and B) in wild-type mice by 36% (S₂/S₁ ratio of 0.92 (control) versus 0.59 (PGB); n = 13 and 14, respectively) (Fig. 5); in the αδ-1 transgenic mice, the glutamate signals were unchanged by this drug (S₂/S₁ = 0.74 (control) versus 0.63 (PGB); n = 9 each) (Fig. 4, C and D).

### Discussion

Aberrant glutamate neurotransmission is linked to a variety of neurological and psychological disorders. Thus, identifying mechanisms that could modulate abnormal glutamate release may provide an avenue for developing new therapeutics for modulating glutamate signaling. Here, we used enzyme-based MEAs to directly measure extracelluar glutamate and observed that both GBP and PGB attenuated the K⁺-evoked glutamate release.

GBP has been used as an antiepileptic, but its precise mechanism of action is unknown (Taylor et al., 2007). To help address this, we stimulated the neural network and measured synaptic spillover of glutamate using a technique that...
we have previously examined in anesthetized animals of locally delivering high K+ solution to evoke depolarization and produce a release of glutamate (Burmeister et al., 2002; Day et al., 2006). We observed in these brain slices an attenuation of glutamate release by GBP after locally delivering high K+,

an effect on neurotransmitter release that is repeatedly observed after stimulus delivery (Dooley et al., 2000a). We have observed previously with L-isoleucine and GBP in rat neocortical slices (Cunningham et al., 2004). A more complex role in neurotransmission and GBP and PGB effectiveness may be the case for these α amino acids where these endogenous ligands may act as “positive modulators required for full functionality of the αβ subunit” (Hendrich et al., 2008).

In a previous study, Wang et al. (1999) showed that the arginine at position 217 in the αβ region of the VSCC is involved in GBP and PGB binding to neocortical membranes was greatly reduced in R217A mice compared with that in wild-type mice (Bian et al., 2006; Field et al., 2006). Accordingly, in slices from R217A mice, we concluded that a functional αβ subunit is necessary for GBP to attenuate K+-evoked glutamate release. Although the mean S2/S1 ratio (0.74) in slices from the R217A mice was lower than the S2/S1 ratio in slices from the wild-type mice (0.92), the means were not significantly different. However, we cannot rule out the possibility of a change in excitability properties of the neurons and synapses in these animals given that the αβ subunit is critical for normal synapse formation or function (Eroglu et al., 2009).

In summary, we showed that GBP and PGB can modulate excitatory neurotransmitter release in rat neocortical brain slices and that the αβ subunit of VSCC is involved in the inhibitory effects of these ligands. This ability to modulate excitatory neurotransmitter release may explain, in

neurotransmission (Dooley et al., 2000a,b; Brown and Randall, 2005). One apparent factor that may influence the effectiveness of these compounds is the type of stimulus, such that GBP and PGB may exert an effect on the prolonged, depolarization-induced neurotransmitter release that more closely resembles hypereexcitability as found in pathological states rather than normal physiological neurotransmission (Dooley et al., 2000a, 2007; Maneuf et al., 2001). In addition, some of the GBP or PGB effects on stimulus-evoked neurotransmitter release may have been diluted in studies on slices or synaptic endings where the whole chamber perfusate is sampled (Dooley et al., 2000a; Fink et al., 2000, 2002) versus the limited focal area sampled by the 50 × 150 μm size of these MEAs in slices. For example, with the high-resolution technique of whole-cell patch clamp in cortical slices, GBP and PGB show an effect as high as ~80% on parameters related to glutamate neurotransmission (Cunningham et al., 2004). Nonetheless, the effect of GBP and PGB on neurotransmitter release remains controversial given reports that GBP and PGB have no effect on K+-evoked glutamate release from human synaptosomes (Brawek et al., 2009) or that the GBP and PGB effect may be linked to the trafficking of calcium channels to the cell surface (Hendrich et al., 2008; Mich and Horne, 2008; Bauer et al., 2009; Thorpe and Offord, 2010). The full effect of these ligands may result from both altering calcium channel trafficking and more rapid modulation of synaptic function (Taylor, 2009).

Meanwhile, the pharmacological effects of PGB are stereoselective, and this was born out by the nonsignificant changes to the S2/S1 ratio with the enantiomer R-IBG. The α amino acids L-isoleucine and L-leucine have been proposed as potential endogenous ligands for the αβ subunit (Thurlow et al., 1993). Although L-isoleucine did not produce a significant change to K+-evoked glutamate release, L-isoleucine did inhibit the PGB attenuation of K+-evoked glutamate release similar to what had been described previously with L-isoleucine and GBP in cortical brain slices (Cunningham et al., 2004). A more complex role in neurotransmission and GBP and PGB effectiveness may be the case for these α amino acids where these endogenous ligands may act as “positive modulators required for full functionality of the αβ subunit” (Hendrich et al., 2008).

Fig. 3. Effects of PGB (0.1–100 μM), R(-)-3-isobutylgaba (100 μM), and L-isoleucine (100 μM) to inhibit K+–evoked glutamate release in rat neocortical slices. A, concentration–effect relationship of PGB and inac-
vative activity of R(-)-3-isobutylgaba, L-isoleucine, and PGB (100 μM) and L-isoleucine (100 μM) combination after repeat superfusion with 70 mM K+ (S1, S2) for 50 s. Substances were present 15 min before S2. Values given are X ± S.E. (n = 7). Analysis of variance of S2/S1 values for control and PGB concentrations gave F(4,30) = 5.97 (p = 0.003). A significant difference from the control value is indicated by an asterisk (*, p ≤ 0.05 and ***, p ≤ 0.001). The S2/S1 ratios obtained for the other substances, including the PGB and L-isoleucine combination, were not significantly different from the control value. B, the transformed data from A depict inhibition (%) by PGB relative to the mean control S2/S1 ratio of 0.97 normalized to 1.0; the corresponding IC50 value was 5.3 μM.
part, the efficacy of these molecules in the clinic. The application of the slice recording methodology coupled to the MEA recording technology establishes a new means to better assess drugs’ mechanisms of action by the direct measurement of neurotransmitter release.

Authorship Contributions

Participated in research design: Quintero, Dooley, Pomerleau, Huettl, and Gerhardt.

Conducted experiments: Quintero.

Contributed new reagents or analytic tools: Gerhardt.

Performed data analysis: Quintero, Dooley, Pomerleau, and Gerhardt.

Wrote or contributed to the writing of the manuscript: Quintero, Dooley, Pomerleau, Huettl, and Gerhardt.

Other: Gerhardt directed research efforts.

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