Vigabatrin-Induced Retinal Functional Alterations and Second-Order Neuron Plasticity in C57BL/6J Mice

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PURPOSE. Vigabatrin (VGB) is an effective antiepileptic that increases concentrations of inhibitory \( \gamma \)-aminobutyric acid (GABA) by inhibiting GABA transaminase. Reports of VGB-associated visual field loss limit its clinical usefulness, and retinal toxicity studies in laboratory animals have yielded conflicting results.

METHODS. We examined the functional and morphologic effects of VGB in C57BL/6J mice that received either VGB or saline IP from 10 to 18 weeks of age. Retinal structure and function were assessed in vivo by optical coherence tomography (OCT), ERG, and optomotor response. After euthanasia, retinas were processed for immunohistochemistry, and retinal GABA, and VGB quantified by mass spectrometry.

RESULTS. No significant differences in visual acuity or total retinal thickness were identified between groups by optomotor response or optical coherence tomography, respectively. After 4 weeks of VGB treatment, ERG b-wave amplitude was enhanced, and amplitudes of oscillatory potentials were reduced. Dramatic rod and cone bipolar and horizontal cell remodeling, with extension of dendrites into the outer nuclear layer, was observed in retinas of VGB-treated mice. VGB treatment resulted in a mean 3.3-fold increase in retinal GABA concentration relative to controls and retinal VGB concentrations that were 20-fold greater than brain.

CONCLUSIONS. No evidence of significant retinal thinning or ERG a- or b-wave deficits were apparent, although we describe significant alterations in ERG b-wave and oscillatory potentials and in retinal cell morphology in VGB-treated C57BL/6J mice. The dramatic concentration of VGB in retina relative to the target tissue (brain), with a corresponding increase in retinal GABA, offers insight into the pathophysiology of VGB-associated visual field loss.

Keywords: vigabatrin, mouse, GABA

The antiepileptic drug, vigabatrin (VGB; \( \gamma \)-vinyl-\( \gamma \)-aminobutyric acid [GABA]) is indicated as first-line, monotherapy treatment for infantile spasms and as adjuvant therapy for refractory complex partial seizures; it is also used as empirical treatment of a rare metabolic disorder, succinic semialdehyde dehydrogenase deficiency.1,2 VGB is a selective, irreversible inhibitor of GABA transaminase and increases intracellular concentrations of GABA, a major neurotransmitter in inhibitory pathways of the central nervous system.3 VGB is rapidly acting and highly effective in controlling epilepsy, with complete cessation of seizures reported in 35% to 75% of infantile spasm cases across multiple studies.2 However, visual field loss has been recognized as an adverse effect of VGB administration that limits the prescription and duration of use of VGB, with the risk of visual field loss increasing from 9% after less than 1 year of treatment to 63% with treatment of more than 2 years in duration.4 In a review of 1678 patients receiving VGB, the random effects estimate of VGB-associated visual field loss (VAVFL) was 34% in children and 52% in adults.5 The frequency and severity of VAVFL as an adverse effect has limited the prescription of VGB to only those seizure disorders where the benefits outweigh the risks and, even then, for a minimal length of time.6 Changes associated with and used as markers for VAVFL in infants and other patients where the visual field cannot
be directly assessed, include attenuation of 30 Hz flicker ERG responses, decreased scotopic and photopic b-wave amplitude, and a marked increase in d-wave latency.7–11 However, there have been conflicting reports of which of these markers are significantly associated with VAVFL in the clinical setting.12–16 Similarly, with VGB administration in laboratory models, reports of VGB’s effects in the retina have been inconsistent. Both enhancement and attenuation of the photopic b-wave amplitude have been reported along with significantly increased, decreased, and insignificantly altered scotopic b-wave amplitudes.12–16 However, there is a paucity of detailed reports on morphologic retinal changes and of VGB and GABA retinal concentrations in pigmented mouse models.

In ongoing efforts to elucidate and mitigate the toxic effects of VGB on the retina, administering VGB to mice and monitoring alterations in retinal structure and function in vivo, we were unable to replicate a previously described model of VGB-associated retinal toxicity in pigmented mice.16 We report striking findings, including marked second-order neuron remodeling, enhancement of scotopic b-wave amplitude with attenuation of later oscillatory potential (OP) wavelets and significant concentration of VGB and GABA in the retina of the VGB-treated pigmented mouse. These findings provide insight into potential causes of VAVFL and suggest specific avenues for future exploration to reduce the frequency and severity of VAVFL in humans.

**METHODS**

**Animals**

Ten-week-old male C57BL/6j (n = 18 per group) were purchased from Jackson Laboratories (Bar Harbor, ME) and housed under relatively bright light intensity conditions (approximately 120 lux; 12-hour light/dark cycle). Mice were maintained on Teklad 8604 rodent diet (Envigo, Madison, WI). All procedures were approved by the University of Wisconsin–Madison Institutional Animal Care and Use Committee and performed in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

**VGB Treatment**

Mice were randomized to receive either VGB 140 mg/kg/d intraperitoneally or vehicle only (0.9% NaCl) at the same volume, duration, and route, as above for 4 weeks (n = 6 per treatment group) or 8 weeks (n = 12 per treatment group). The mice were injected 5 days per week with 2 consecutive days without injection. On the last day of the 5 consecutive days of intraperitoneal injection, a 280-mg/kg dose was delivered. This VGB dose of 140 mg/kg/d was based on human clinical doses.2 VGB was synthesized as described in Walters et al., 2019 for the first cohort of 12 C57BL/6j mice (n = 6 per treatment group) and purchased from Sigma-Aldrich (St. Louis, MO) for the second and third cohorts of mice (n = 12 per treatment group).

ERG was performed between 5 AM and 1 PM on Tuesday and Wednesday of each week of testing, with alternation between treatment groups, in mice anesthetized with ketamine (100 mg/kg, Akorn, Lake Forest, IL) and xylazine (10 mg/kg, Akorn) and pupils dilated with 0.5% tropicamide (Akorn). Silver thread electrodes (OcuScience, Henderson, NV) attached to 2.5-mm contact lenses were placed on the corneas with 2.5% methylcellulose gel (Akorn), and subdermal reference and ground needle electrodes were placed on the nose between the eyes, and at the base of the tail, respectively, under dim red light. The red light was extinguished for at least 2 minutes before obtaining recordings. Body temperature was maintained around 37°C with a heating pad. An Espion c system with a ColorDome Ganzfeld stimulator (Diagnosys LLC, Lowell, MA) was used to elicit and record full-field flash and flicker ERG responses under both scotopic (after a >10-hour overnight dark adaptation) and subsequently photopic conditions (after 10 minutes of 30 cd/m² light adaptation) with corresponding waveforms acquired from both eyes. Scotopic flash intensity ranged in 8 steps from 0.001 to 10.000 cd/s/m² with a 2- to 5-second interstimulus interval. Photopic flash intensity ranged in six steps from 0.78 to 20.00 cd/s/m² with a 2- to 10-second interstimulus interval in the presence of a 6 cd/m² rod-suppressing background. Flicker responses were obtained at 8 Hz and 16 Hz at the maximum intensity for photopic and scotopic conditions, respectively. The c-wave was measured in response to 2.5 and 25 cd/s/m² stimuli immediately before light adaptation. The a-wave was measured from baseline to the trough of the first corneal negative deflection. The b-wave was measured from the trough of the a-wave to the peak of the succeeding corneal positive deflection, with care taken to avoid selecting OPs. The a-, b-, and c-wave amplitudes and implicit times were measured with Espion V6 software (Diagnosys LLC). The OP root mean square, amplitude, and implicit time of individual wavelets and c-wave amplitude were measured with an in-house R script after a bandpass filter was applied from 100 to 300Hz by the Espion V6 software. The OP root mean square was calculated from 10 to 60 milliseconds after the stimulus; OP1 was measured from baseline to the first positive peak; each successive OP (OP2–OP4) was measured from the preceding negative deflection to the immediately following positive peak. Analysis of the ERG data was performed with open-source software R (version 3.5.2; The R Foundation, Vienna, Austria) for summary statistics of the a- and b-wave and OPs. Naka-Rushhton functions were fitted to b-wave data with an in-house R script following the equations described by Naka and Rushton.18,19

**Optomotor Response (OMR) Testing**

The qOMR system (Phenosys GmbH, Berlin, Germany) was used to determine the visual acuity threshold (maximum spatial frequency at which an OMR is elicited to a consistent, high-contrast grating) in awake, unrestrained mice. Sinusoidal gratings ranging from 0.05 to 0.50 cycles per degree and at maximal contrast were displayed in a randomized order for 1 minute with a solid gray pattern displayed for 10 seconds in between gratings. Data were averaged from six sessions per time point for each subject, using integrated software for detection of head tracking movements and projection of a virtual cylinder of revolving gratings of different spatial frequencies. Mice were tested for no more than 30 minutes at one time with at least 2 hours between testing sessions. The visual acuity threshold was determined by fitting a sigmoid curve as described by Frund et al., where the curve reaches 25% of the maximum response.20–22

Optical coherence tomography (OCT) was performed in ketamine/xylazine anesthetized mice using Spectralis HRA+OCT (Heidelberg Engineering Inc., Heidelberg, Germany). Pupils were dilated with 0.5% tropicamide, and
Corneal desiccation prevented by the application of a corneal contact lens. OCT volume scans (20 × 20 degrees, 120 μm width) were obtained centered on the optic nerve head and in consistent locations and orientation between testing sessions by use of proprietary tracking software. Total retinal thicknesses were calculated by the manufacturer’s proprietary Eye Explorer algorithm, with manual correction of segmentation as needed, and averaged for each region.24

**Immunohistochemistry**

After euthanasia by cervical dislocation, freshly enucleated eyecups were placed in artificial cerebrospinal fluid (pH 7.4, containing in millimole: 119 NaCl, 2.5 KCl, 2.5 CaCl2, 1.3 MgCl2, 1 NaH2PO4, 11 glucose, and 20 HEPES) with the retinas dissected within 30 minutes and fixed for 30 minutes at room temperature in 4% (w/v) paraformaldehyde prepared in artificial cerebrospinal fluid. Thereafter the retinas were rinsed in 1 × PBS (pH 7.4) and embedded in 4% agarose for vibratome sectioning (120 μm thickness). Retina slices were incubated with primary antibodies in PBS containing 0.5% Triton X-100 and 5% normal donkey serum for 24 hours at 4°C. The primary antibodies used were directed against protein kinase C alpha (1:1000; Sigma, St. Louis, MO), glutamic acid decarboxylase 67 (1:1000; Millipore, Burlington, MA), glycine transporter 1 (1:1000; Synaptic Systems, Goettingen, Germany), Ribeye (1:1000, Synaptic Systems), synaptotagmin 2 (1:1000; Zebrafish International Resource Center, Eugene, OR), glial fibrillary acidic protein (1:1000; Novus Biologicals, Centennial, CO, and Invitrogen, Carlsbad, CA), cellular retinaldehyde-binding protein (1:1000; Cambridge, UK), and calbindin (1:500; Swant, Marly, Switzerland). After rinses in PBS, the samples were routinely labeled with appropriate Alexa Fluor-conjugated secondary antibodies (1:1000) for 3 hours at room temperature. Fluorescein-conjugated peanut agglutinin (1:200; Vector Laboratories, Burlingame, CA) was included within the secondary antibody incubation step. The samples were then rinsed, mounted on glass slides with Vectashield (Vector Laboratories) and imaged with a Leica TCS SP8 confocal microscope (Leica Microsystems, Wetzlar, Germany). Images were acquired with a 20× (Numerical Aperture (NA) = 0.75) or 60× (NA = 1.4) objective at a resolution of 0.1 μm × 0.1 μm × 0.3 μm. Cell numbers were quantified by manual counting in FIJI.24

**Mass Spectrometry**

The brain, liver, and retina were rapidly dissected within 5 minutes after cervical dislocation and flash frozen in liquid nitrogen. The tissues were stored at –80°C and shipped on dry ice for extraction and quantification of VGB and GABA concentrations in tissue samples, performed as described by Walters et al.17

**Statistical Analyses**

Statistical comparisons were performed with Prism 8 (GraphPad, San Diego, CA, USA) by t-tests with Holm-Sidak correction for multiple comparisons, and mixed-effects models with Tukey post hoc tests. Results were considered significant at a P value of less than .05.

**ERG**

Statistically significant enhancement of the scotopic b-wave amplitude in response to mesopic light stimuli was identified in VGB-treated mice between 2 and 4 weeks of VGB treatment, relative to vehicle-treated mice at the same time points (Fig. 1). The photopic b-wave amplitude tended to be higher in VGB-treated mice than control between 2 and 4 weeks as well, but differences were not statistically significant (Fig. 2). Although the photopic and scotopic b-wave amplitudes were lower than baseline values after 8 weeks of treatment in the vehicle-treated group, this age-related decrease was expected.25

Implicit times of the scotopic b-wave were shorter in VGB-treated mice compared with age-matched vehicle-treated controls in response to 0.05, 5.00 and 10.00 cd*s/m² stimuli between 2 and 8 weeks of treatment (Fig. 3). In contrast, implicit times of the photopic b-wave were prolonged in response to all tested stimuli in mice treated with VGB for between 2 and 4 weeks, relative to age-matched vehicle-treated controls (Fig. 3). Root mean square values of scotopic (Fig. 4) and photopic (Fig. 4) OPs were significantly lower in VGB-treated mice compared with controls between 2 and 8 weeks of VGB treatment. Specifically, there was a significant reduction in the scotopic OP3 and OP4 waveform components with 1 cd/s/m² stimulus (Fig. 5) and a significant reduction of the photopic OP3 with 10 cd/s/m² stimulus (Fig. 6). Implicit times of the photopic OP2 and OP3 were significantly prolonged in VGB-treated mice compared with age-matched controls between 2 and 6 weeks of VGB treatment (Fig. 6). In contrast with the b-wave, after 8 weeks of treatment, the scotopic a-wave amplitude of the ERG was significantly decreased in VGB-treated mice compared with age-matched vehicle-treated mice (Fig. 7). No significant changes were identified in the photopic and scotopic a-wave implicit time, photopic a-wave amplitude, scotopic and photopic flicker responses or c-wave amplitude (not shown). (For all ERG results presented, from baseline to 4-week timepoints: VGB-treated n = 18, control n = 17; at 6 week timepoint: VGB-treated n = 12, control n = 11; and at 8 weeks: VGB-treated n = 10, control n = 11. The control group is smaller than the VGB-treated group because initial ERG indicated no response to light in a single mouse assigned to that group. Six mice per group were euthanized at the 4-week timepoint for histology. A further two mice were lost from the VGB group when they died between weeks 6 and 8 of the study (the cause of death was not established in either mouse).

**OMR**

No statistically significant differences in visual acuity were identified between groups by qOMR testing over 8 weeks of VGB treatment (t-test) (Fig. 8A).
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**Optical Coherence Tomography**

No alteration in retinal morphology were identified with OCT imaging over 8 weeks of VGB treatment (Fig. 8B). There were no significant differences in retinal thickness between control and VGB-treated mice throughout the study (t-test) (Fig. 8C).

**Immunohistochemistry**

In mice treated with VGB for 8 weeks, antibodies directed against protein kinase C alpha consistently identified striking extension of rod bipolar cell dendrites into the outer nuclear layer of the retina. Protein kinase C alpha–labeled rod bipolar cell dendritic extensions were closely apposed to the ribbon synapse marker Ribeye, enriched at photoreceptor terminals (Fig. 9). Additionally, extension of synap-
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**FIGURE 2.** The photopic b-wave (A) amplitude of VGB-treated mice (orange) is qualitatively increased between 2 and 4 weeks of VGB treatment compared with controls (blue) at those timepoints (B) as well as baseline values (C). LA, light adapted.

totagmin 2-positive (type 2 OFF and type 6 ON bipolar) cone bipolar cell dendrites into the outer nuclear layer could be observed in 8-week VGB-treated retinas (Fig. 10). Dendritic sprouting of calbindin-positive horizontal cell processes into the outer nuclear layer could also be observed in VGB-treated retina (Fig. 11), providing further evidence of retinal neuronal plasticity upon VGB treatment. These findings were not identified in retinas of mice treated with VGB for 4 weeks or in vehicle-treated mice at any time point (Figs. 9, 10, 11). Initial studies provided
FIGURE 3. The scotopic b-wave implicit time (A) of VGB-treated mice (open circles) is selectively decreased compared with age-matched controls (filled circles) between 2 and 8 weeks of VGB treatment. The photopic b-wave implicit time (B) of VGB-treated mice is globally increased compared with controls between 2 and 4 weeks of VGB treatment with a trend toward increased implicit time at 6 and 8 weeks as well.

no evidence suggestive of cone photoreceptor loss in VGB treated retinas as determined by labeling with peanut agglutinin (Fig. 11, but with only one mouse per treatment group), or of gliosis in response to VGB treatment when labeled with antibodies against the glial fibrillary acidic protein (Fig. 10). No significant differences in either rod bipolar cell, glycinergic, or GABAergic amacrine cell numbers were identified between vehicle-treated and VGB-treated retinas (Fig. 12).

Mass Spectrometry
The active $S^+$ enantiomer of VGB was concentrated in the retinas of VGB-treated mice by a mean of 21-fold compared with the brain of VGB-treated mice ($n = 6$; $P < .0001$, Welch’s $t$-test) (Fig. 13A). No appreciable amounts of VGB were detected in tissues of vehicle-treated mice. In VGB-treated mice compared with saline-treated controls, GABA concentrations in the retina were increased 3.3-fold ($n = 6$; $P < .0001$, $t$-test) (Fig. 13B).

DISCUSSION
In this study of pigmented adult mice, we identified subtle changes in retinal function: a transitory increase in the ERG b-wave amplitude that was maximal after 4 weeks of VGB treatment and persistent depression of OP amplitudes. Enhanced b-wave amplitudes have been described previously both in VGB-treated albino and pigmented rats. In contrast, there are also multiple reports of ERG deficits
Figure 4. VGB attenuates the OP RMS in mice. The scotopic OP RMS (A) is decreased in VGB-treated mice (orange) compared with controls (blue) at the 2-, 4-, and 8-week timepoints. The photopic OP RMS (B) is decreased in VGB-treated mice at 2 weeks and remains qualitatively decreased from 4 to 8 weeks of VGB treatment. RMS, root mean square. *P < .05; **P < .01; ***P < .001.

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of varying magnitude in laboratory animals treated with VGB, including rabbits, pigmented mice, and albino and pigmented rats. In laboratory studies of VGB, OPs have been reported infrequently. However, VGB treatment has been found in two separate studies to have either no effect on OP amplitudes or to decrease OP amplitudes in pigmented rats. These discrepancies between studies could be reflective of differences in VGB dose, route of administration, dietary taurine and other amino acid concentrations, and environmental light conditions, as well as species and strain of the animals. In our studies, we used a pigmented mouse strain, C57BL/6J, and confirmed the absence of a common confounding mutation associated with retinal pathology in mice of this general genetic background. Care was taken to ensure that all mice were subject to consistent light levels because light exposure has been reported to exacerbate severity of VGB toxicity. Our findings were reproduced in multiple cohorts of mice using two sources of VGB. Retinal VGB concentrations were confirmed, as was elevated retinal GABA, indicating a pharmacologic effect of VGB on the physiology of the retina.

Accumulation of GABA within the retina was pronounced, as previously reported, and we extend these findings to confirm that there was marked increase in VGB concentration in the retina of VGB-treated mice. It is likely that supraphysiologic retinal GABA concentrations impact GABAergic signaling in retinal neurons and could result in abnormal bipolar cell-driven responses (principally ERG b-wave) and amacrine cell-associated ERG responses (OPs), as we identified in this study. However, the precise underlying mechanism(s) for effects of accumulation of endogenous retinal GABA on the ERG remains unclear. In both in vivo and ex vivo studies, administration of exogenous GABA has been found to enhance, reduce, or have no significant effect on b-wave amplitude. In C57BL/6 mice, intravitreal injection of GABA has been found to decrease the b-wave amplitude of responses to dim light stimuli, but increase the b-wave amplitude of responses to brighter stimuli. Inhibitory inputs in the retina are largely mediated by GABA_A, GABA_C, and glycine receptors, which are expressed in different ratios on various retinal cell types. Studies conducted to examine differential effects of GABA_A and GABA_C receptors on
FIGURE 5. VGB attenuates the amplitude of later scotopic OP wavelets in mice. In VGB-treated mice (open circles), although OP1 and OP2 amplitudes were not significantly different (A, B), the DA 1 cd/s/m² OP3 amplitude was decreased in VGB-treated mice (filled circles) compared with time-matched controls at 4 weeks of VGB treatment (C). The OP4 amplitude was attenuated in VGB-treated mice at 2, 4, and 8 weeks of VGB treatment (D). Implicit times of OP wavelets were not significantly different between groups (A–D). Representative waveforms from both right and left eyes shown for one control and one VGB-treated mouse (E). DA, dark adapted. *P < .05; **P < .01; ***P < .001.
FIGURE 6. VGB decreases the amplitude of later photopic OP wavelets in mice while increasing the implicit time. In VGB-treated mice (open circles), although OP1 amplitudes (A) were not significantly different between groups, the LA 10 cd*s/m² OP2 and the OP3 implicit time (B, C) was increased compared with age-matched controls (filled circles) between 2 and 6 weeks of VGB treatment. Additionally, the OP3 amplitude was decreased in VGB-treated mice relative to age-matched controls between 2 and 8 weeks of VGB treatment (C). Representative waveforms from both right and left eyes shown for one control and one VGB-treated mouse (D). LA, light adapted. *P < .05; **P < .01; ***P < .001.
the b-wave by pharmacologic blockade have also yielded conflicting results between and within mammalian species. GABA<sub>B</sub> receptor blockade resulted in an increase in b-wave amplitude in the rat, but abolished the b-wave in the rabbit. Effects of GABA<sub>C</sub> receptor blockade have been similarly variable, with reports of increase, decrease, or no effect on the b-wave amplitude of the ERG. These inconsistencies could be reflective of differences in species, experimental techniques, and concentration and choice of pharmacologic agents used, but leaves no clear consensus on the effect of GABA and its receptors on the b-wave.

Our finding of attenuation of the later OP waveform components (OP3 and OP4) in VGB-treated mice is likely due to supraphysiologic concentrations of GABA in the retina, as supported by previously published literature. It has been shown that intravitreal injection of GABA abolishes OPs in C57BL/6 mice, with similar results reported in the rat, including complete OP removal and selective abolishment of later wavelet components. This finding is consistent with reports that later OPs are primarily generated or at least modulated by third-order neurons in the ON pathway, particularly GABAergic amacrine cells. Further, it is clear that GABA receptors in bipolar cells and third-order neurons, likely GABAergic amacrine cells, play an important role in shaping the ERG b-wave, as well as the OPs. There is a compelling body of work demonstrating that pharmacologic blockade of GABA<sub>B</sub> receptors can decrease b-wave amplitude while enhancing OP amplitude, GABA receptors respond differentially to tonic versus phasic stimuli and GABA<sub>C</sub> receptors do not desensitize and are more sensitive to GABA than GABA<sub>B</sub> receptors. Importantly, differential uptake and activity of VGB have been shown in various retinal cell types, with a preference toward GABAergic amacrine cells. Thus, our findings underscore an apparent association between supraphysiologic concentrations of GABA within the VGB-treated retina, b-wave amplitude enhancement, and depression of the later OPs. These effects on retinal electrophysiology may be attributable to preferential stimulation of GABA<sub>B</sub> receptors and/or GABA-modulated disinhibition of retinal inhibitory circuits at the bipolar axon terminal. However, the observed enhancement of the b-wave was not sustained throughout the entire duration of the 8-week study period. The transitory nature of this effect may reflect homeostatic processes regulating GABA receptors and changes in other contributions to the b-wave. Unfortunately, within the scope of the present study, it was not possible to interrogate the precise cellular and molecular mechanisms responsible for the observed functional abnormalities.

In addition to these functional changes, we identified morphologic evidence of pronounced retinal plasticity of second-order neurons. This neuronal plasticity was, however, not directly associated with our finding of b-wave amplitude enhancement, although it affected bipolar cells, because neuronal plasticity was not identified at the 4-week time point when the b-wave enhancement was most evident. Thus, neuronal plasticity may instead reflect changes associated with more chronic VGB administration. Broad extension of rod bipolar dendrites upon VGB treatment has been previously described in albino rats and mice, along with horizontal cell dendritic outgrowth. Here we extend these findings to pigmented mice and additionally provide evidence for dendritic sprouting of cone bipolar cells, which was reported to not occur in albino mice. An outgrowth of bipolar cell dendrites may be observed under pathologic conditions such as progressive photoreceptor degeneration and retinal detachment and in mutant knockouts such as the nob2 mouse with a calcium channel Cac1.4 null mutation and Bsn mice lacking functional Bassoon protein, as well as in aged wild-type C57BL/6 mice.

Concurrent horizontal cell plasticity has also been identified under most of those conditions outlined. In contrast with the presynaptic changes described, bipolar cell dendrite outgrowth has not been associated with...
FIGURE 9. After 8 weeks of treatment, extension of rod bipolar cells dendrites (arrowheads) into the ONL is evident (anti-PKCα, green) in VGB-treated mice (A, B) and is not seen in sham-treated mice (C). This finding was consistent in four retinas. There is colocalization of synaptic ribbons (anti-Ribeye, magenta) at the sites of PKCα labeling. Bipolar cell plasticity was not identified at 4 weeks of VGB treatment (D). Scale bar = 25 μm. GCL, ganglion cell layer; INL, inner nuclear layer; IPL, inner plexiform layer; ONL, outer nuclear layer; OPL, outer plexiform layer; PKCα, protein kinase C alpha.

FIGURE 10. After 8 weeks of treatment, extension of cone bipolar cell dendrites (arrowheads) into the ONL is evident (antisynaptotagmin 2, cyan) in VGB-treated mice (A, B) and is not seen in sham-treated mice (C). No apparent changes in GFAP expression (anti-GFAP, yellow) or glycineric amacrine cells (anti-GlyT1, red) can be observed upon VGB treatment. Scale bar = 25 μm. GCL, ganglion cell layer; GFAP, glial fibrillary acidic protein; GlyT1, glycine transporter 1; INL, inner nuclear layer; IPL, inner plexiform layer; ONL, outer nuclear layer; OPL, outer plexiform layer.

postsynaptic changes in the photoreceptor–bipolar cell synapse.57–59 In light of this evidence, it is reasonable to suggest that photoreceptor presynaptic transmission is impaired by VGB, such that the photoreceptor terminals retract, and bipolar and horizontal cell dendritic extend to maintain the synaptic connection, although the initiating event is unclear and could not be elucidated in our study design. Evidence for significant cone photoreceptor damage or gliosis was absent in our mice, contrary to previously published work in albino rodents,13,14 perhaps reflective of the protective nature of ocular pigmentation.90 Light-induced retinal toxicity has been well-documented in albino rodents, including photoreceptor damage and retinal gliosis, and could have been possible even at standard vivarium illumination levels, and particularly in conjunction with VGB-induced retinal toxicity.61–64

In addition to light exposure, deficiency of the amino acid taurine has been shown to influence the severity of VGB toxicity in albino rats.65 Administration of VGB has been reported to interfere with taurine transport and result in taurine plasma deficiency below reference levels in VGB-treated animals and human infants.65,66 In VGB-treated neonatal and adult albino rats, bipolar cell plasticity and cone photoreceptor and retinal ganglion cell damage have been demonstrated with concurrent taurine deficiency,27,65 with taurine supplementation partially rescuing these changes in VGB toxicity.27,65 Taurine depletion in rodent models with guanidoethane sulfonate also results in bipolar cell plasticity, as well as cone photoreceptor and retinal ganglion cell damage.67 In these rodent models, animals were fed a low protein, low nutrient, and vegetarian diet without taurine, whereas mice in the current study were maintained on a nonvegetarian diet with animal protein, including a source of taurine.15 Although quantification of dietary or tissue taurine levels was outside the scope of our study, the manifestation of enhanced ERG responses,
FIGURE 11. After 8 weeks of treatment, there was no identifiable change in cone photoreceptors (PNA, green) in VGB-treated retina (A) compared with sham treated (B). Extension of horizontal cell dendrites (arrowheads) into the ONL is evident (anticalbindin, red) in VGB-treated mice (C, D) and is not seen in sham-treated mice (E). Horizontal cell plasticity was not identified at 4 weeks of VGB treatment (F). Scale bar = 25 μm. GCL, ganglion cell layer; INL, inner nuclear layer; IPL, inner plexiform layer; ONL, outer nuclear layer; OPL, outer plexiform layer; OS/IS, outer and inner segments; PNA, peanut agglutinin.

or even complete rescue from morphologic effects of VGB toxicity, has not been demonstrated with taurine supplementation. In one study of pigmented rats, taurine was not found to have a protective effect against retinal VGB toxicity. Further, some subsets of human patients receiving VGB have not shown significantly decreased taurine concentrations compared with reference levels. However, a recent report of just two cases suggested that taurine supplementation may help to mitigate retinal VGB toxicity in pediatric patients with the rare metabolic disorder, succinate semialdehyde dehydrogenase deficiency. Then, there are additional factors involved in VGB-associated retinal morphologic and functional changes described, because the effects of VGB on amino acid metabolism and synthesis within neural tissues, including the retina, is complex and taurine deficiency models alone do not fully address the observed effects.

In human patients, there have been only very limited histologic reports of retinal morphologic changes associated with VAVFL and, as with animal models, functional changes attributed to VGB administration have been inconsistent. The pathophysiology of VGB resulting in visual field loss remains uncertain with a majority of the literature describing electrophysiological markers, testing and/or prevalence of VAVFL. Despite significant retinal remodeling and ERG perturbation in our mice, the lack of significant alteration in the visual acuity threshold between VGB-treated and sham-treated mice suggests that these morphologic and functional retinal changes identified at 8 weeks of VGB treatment do not severely impact vision. In this respect, our finding is consistent with the generally asymptomatic nature of VAVFL in people and highlights the importance of developing sensitive and specific tests for VAVFL for patients where perimetry is impossible. With regard to the electrophysiologic findings described in humans, it is important to separate unassociated, reversible physiologic effects, and pathologic findings associated with VAVFL. A reduction in the International Society for Clinical Electrophysiology of Vision light-adapted 30-Hz flicker has been generally accepted as a reliable marker for VAVFL, although not without some controversy. Abnormalities in the 30-Hz flicker is considered reflective of (largely postreceptor) dysfunction in the cone pathway and is consistent with our findings of cone bipolar and horizontal cell plasticity. Although photopic ERG b-wave amplitudes were comparable in our study, in contrast with previous findings in human patients, this finding may reflect differences in the duration of treatment, because an initial increase preceding a decrease in the photopic b-wave amplitude was also described in humans. Consistent with our findings in
mice, a reduction in OP amplitudes was reported in human patients, but with a greater effect on the early OP wavelets.\(^8^2\) These findings in human subjects are broadly similar to the results of this study, although we found later OP wavelets to be significantly affected rather than early OP wavelets. The reason for this apparent species difference is unclear. Because the OP deficits demonstrate dramatic improvement upon discontinuation of VGB in human patients, it has been postulated that OP alterations represent a physiologic response that is not directly associated with the pathology of VAVFL.\(^7^9\) Bipolar cell abnormalities with presynaptic photoreceptor disruption could explain ERG perturbations described commonly in humans, such as attenuation of the light-adapted 30-Hz flicker, and warrant further study in our model.

In summary, we document significant alterations in retinal function and morphology secondary to VGB administration in C57BL/6j mice, primarily enhancement of the b-wave amplitude with concurrent depression of OPs and second-order neuron plasticity. Within the scope of the current study, we were unable to establish the precise mechanisms responsible for these functional and morphologic effects of VGB administration. However, we confirmed the dramatic preferential accumulation of VGB in the mouse retina over the brain, the target organ for VGB's antiepileptic activity. Irrespective of the precise mechanisms responsible for VAVFL, it is evident that our results underscore the association between the intense concentration of VGB and GABA in the retina and VGB-associated retinal toxicity. Our findings also highlight the current knowledge gaps that are important to address in ongoing efforts to understand the pathophysiology of VGB-associated retinal toxicity, namely, (1) mechanisms responsible for the preferential accumulation of VGB in the retina, (2) the effects of chronic supraphysiologic concentrations of GABA on retinal circuitry, and (3) the role of photoreceptors, horizontal, and amacrine cells in the observed changes in retinal bipolar cells. In turn, the answers to these questions are likely to advance efforts to mitigate VAVFL in human patients by informing the design of pharmacologic approaches to avoid VGB concentration in off-target tissues, or of drugs in this class that do not share VGB's adverse effects.

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VGB-Induced Retinal Neuron Plasticity in Mice

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