Daily Isoflurane Exposure Increases Barbiturate Insensitivity in Medullary Respiratory and Cortical Neurons via Expression of ε-Subunit Containing GABA_{A}Rs

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Abstract

The parameters governing GABA_{A} receptor subtype expression patterns are not well understood, although significant shifts in subunit expression may support key physiological events. For example, the respiratory control network in pregnant rats becomes relatively insensitive to barbiturates due to increased expression of ε-subunit-containing GABA_{A}Rs in the ventral respiratory column. We hypothesized that this plasticity may be a compensatory response to a chronic increase in inhibitory tone caused by increased central neurosteroid levels. Thus, we tested whether increased inhibitory tone was sufficient to induce ε-subunit upregulation on respiratory and cortical neurons in adult rats. Chronic intermittent increases in inhibitory tone in male and female rats was induced via daily 5-min exposures to 3% isoflurane. After 7d of treatment, phrenic burst frequency was less sensitive to barbiturate in isoflurane-treated male and female rats in vivo. Neurons in the ventral respiratory group and cortex were less sensitive to pentobarbital in vitro following 7d and 30d of intermittent isoflurane-exposure in both male and female rats. The pentobarbital insensitivity in 7d isoflurane-treated rats was reversible after another 7d. We hypothesize that increased inhibitory tone in the respiratory control network and cortex causes a compensatory increase in ε-subunit-containing GABA_{A}Rs.

Introduction

One of the features defining functional and regional subgroups of neurons in the CNS is the local expression of different patterns of GABA_{A}R subunits [1]. Tonic GABA_{A}Rs, which regulate network excitability, are principal targets of allosteric positive GABA_{A}R modulators [2,3], which include ethanol [4], many anesthetics [5,6], some drugs of abuse such as barbiturates...
[7], and neurosteroids that are increased during pregnancy [8,9]. The expression patterns of these GABA_{A}R subtypes are regulated in a compensatory manner (for review, see [10]), such that increases in allosteric modulators downregulate hippocampal and cerebellar GABA_{A}Rs involved in tonic current generation [11,12]. Disruption in GABA_{A}R regulation is associated with a variety of affective disorders (for review, see [13,14]), and during pregnancy, hippocampal networks are less stable and more easily rendered epileptic than in non-pregnant animals [15].

Regulation of GABA_{A}R subunit expression in the CNS is complex and poorly understood, especially during pregnancy when both increases [16] and decreases [15] in hippocampal δ-subunit expression are observed. We previously described a compensatory plasticity during pregnancy in which ε subunit-containing GABA_{A}Rs, which conduct a tonic current and are insensitive to many allosteric modulators [17,18, 19], are upregulated on respiratory rhythm-generating medullary neurons [20]. We hypothesized that a subunit-specific form of GABA_{A}R plasticity promotes stable respiratory output by decreasing neuronal sensitivity to circulating inhibitory neurosteroids. Despite recent interest in the complex patterns of GABA_{A}R subunit plasticity, it remains unclear what stimuli are required to engage these mechanisms. One possibility is that neurosteroid receptor activation, which is a potent activator of gene transcription [21], may result in a transcriptional feedback control over GABA_{A}R subunit composition. Alternatively, neurosteroids can activate the PKC-dependent phosphorylation of residues on specific subunits leading to increased membrane insertion of receptor complexes [22]. Finally, neurosteroids may not be required at all. Chronic changes in activity are sufficient to induce homeostatic regulation of neuronal activity in vivo [23, 24], and tonic GABA_{A}Rs can be recruited by cortical neurons to stabilize perturbations in channel expression [25].

While seeking an answer to these questions, we serendipitously observed that the respiratory plasticity observed during pregnancy can be induced in virgin animals: female rats exposed daily to a brief dose of isoflurane (for estrous cycle tracking) developed a phenotype strikingly similar to that of pregnant animals. Combined with data suggesting that chronic ethanol administration and pregnancy have similar effects on cerebellar and hippocampal GABA_{A}Rs [15, 26], we hypothesized that GABA_{A}R plasticity is stimulated in the respiratory system via repetitive manipulation of inhibitory tone. It is important to note that isoflurane acts on a variety of systems. In the nucleus ambiguus, which is adjacent to the medullary respiratory regions investigated here, isoflurane potentiates both tonic and phasic GABA_{A}R inhibition [27]. While a primary target of isoflurane in medullary and spinal neurons is tonic/phasic GABAergic inhibition, others targets include glycine receptors [28] as well as excitatory synaptic currents [29].

Previously [20], we reported that ε subunit-containing GABA_{A}Rs may be under activity-dependent transcriptional control because the 5’ flanking region of the gene encoding the ε subunit has conserved binding sites for CREB and SRF which are both implicated in activity-dependent neuronal gene expression and plasticity [30, 31, 32]. Similarly, GABA_{A}R subunit expression patterns are partly regulated by activity dependent transcriptional control in cortical cultures [33]. Thus, we predicted that respiratory rhythm-generating neurons would increase expression of ε subunit-containing GABA_{A}Rs in a predictable, compensatory manner when challenged with isoflurane.

Accordingly, we employed a chronic intermittent anesthetic exposure paradigm to determine whether increased ε subunit expression: 1) can be experimentally induced in the respiratory control network and decrease pentobarbital sensitivity both in vivo and in medullary slices in vitro, 2) can be induced in non-respiratory neurons, such as cortical neurons, 3) is restricted to only female animals. Here we demonstrate that 7d isoflurane exposures reversibly increase GABA_{A}R ε subunit expression on medullary respiratory neurons, especially neurons in the...
PreBötzinger Complex (preBötC) which is hypothesized to be the inspiratory rhythm generator in the mammalian brain [34, 35]. Furthermore, we show that after 30 days of treatment, GABA\(_{A}\)R \(\varepsilon\) subunit expression is also increased on cortical neurons.

Methods

Ethical Approval

All experimental procedures were conducted in accordance with NIH guidelines and approved by the University of Wisconsin-Madison Institutional Animal Care and Use Committee (protocol V00936). A total of 95 rats (51 females and 44 males) were used in this study. All efforts were made to minimize discomfort and suffering of animals. For sacrifice, animals were either deeply anesthetized prior to decapitation.

Isoflurane Treatment

Rats were exposed to 3% isoflurane (balance O\(_2\)) for 5 min per day for 7d or 30d. Breathing frequency unambiguously decreased during the 5-min isoflurane exposures, although this was not systematically quantified. One group of rats was allowed to recover for 7d following a 7d treatment. Control rats were exposed to 5 min of 100% O\(_2\), the vehicle, or room air in the anesthetic chamber. No differences were observed between these two control treatments, thus the data were merged.

In Vivo Phrenic Nerve Recordings

Three groups of rats (Sprague Dawley, Harlan) were studied: adult male and female untreated control rats (3–4 mo; n = 12), 30d isoflurane-treated female rats (3–4 mo, n = 4), and 7d isoflurane-treated male and female rats (4 mo n = 8). The methods for phrenic nerve recordings and pentobarbital dose response were described previously [20]. Briefly, rats were anesthetized initially with isoflurane (3.0–3.5%, 50% O\(_2\)) for approximately 1 h, and then slowly converted to urethane anesthesia (1.6 mg/kg, i.v.). Rats were paralyzed (pancuronium bromide, 2.5 mg/kg, i.v.), bilaterally vagotomized and ventilated with a rodent respirator (Small Animal Ventilator, Model 683, Harvard Apparatus Inc., Holliston, MA, USA). Blood samples (60 \(\mu\)l) were drawn to determine arterial blood gases (PaO\(_2\) and PaCO\(_2\)), pH and base excess (ABL 810, Radiometer, Copenhagen, Denmark). Body temperature was maintained at approximately 37°C using a heated table. End-tidal CO\(_2\) was measured with a flow-through capnograph (Capnogard, Novametrix, Wallingford, CT, USA). The right phrenic nerve was isolated via a dorsal approach, cut distally, desheathed, submerged in mineral oil and placed on bipolar, silver wire electrodes. Nerve activity was amplified (10,000x), band pass filtered (100 Hz to 10 kHz) (Model 1700, A-M Systems, Inc., Carlsborg, WA, USA) and integrated (time constant = 50 ms, Model MA-821RSP, CWE Inc., Ardmore, PA, USA).

Recordings began approximately 60 min post-surgery. The nerve was allowed to stabilize under baseline conditions of hyperoxia (PaO\(_2\) = 150 mmHg) and hypercapnia (PaCO\(_2\) = 60 mmHg). Hypercapnia was maintained throughout an experiment to ensure a chemical drive to breathe. Ten pentobarbital injections were administered (10 mg/kg/injection i.v.), each separated by 5 min. Following the final pentobarbital injection, 5 min of hypercapnic hypoxia was administered (PET CO\(_2\) = 80 mmHg, 13% inspired O\(_2\)) to estimate the scope of phrenic nerve activity.
In Vitro Recordings

A total of 54 rats were used for in vitro electrophysiology studies. Ten groups of rats were studied: adult males (3–4 months; n = 6), adult virgin females (3–4 months; n = 6), adult male and female oxygen controls (3–4 months; n = 4), adult male and female time controls (3–4 months; n = 5), adult male and female bicuculline controls (3–4 months; n = 4), 7d isoflurane-treated adult males (3–4 months, n = 6), 7d isoflurane-treated adult females (3–4 months, n = 6), 30d isoflurane-treated adult males (4 months, n = 6), 30d isoflurane-treated adult females (4 months, n = 6), 7d isoflurane-treated, 7d recovery adult males and females (3–4 months, n = 7).

Methods for in vitro multielectrode array recordings and analysis were described previously [20]. Briefly, brains were removed and coronal medullary and cortical slices were cut in cold (0°C) 3 mM KCl artificial cerebrospinal fluid (aCSF) with a vibrating microtome (Campden Instruments, Layfayette, IN, USA). The aCSF composition was (in mM): 120 NaCl, 26 NaHCO₃, 20 glucose, 2 MgSO₄, 1.0 CaCl₂, 1.25 Na₂HPO₄, 7 KCl. Cortical slices (375 μm thick) contained primary motor and primary somatosensory areas. To remove the medulla, transverse cuts were made at caudally at spinal segment C1 and rostrally at the pontomedullary junction. A series of slices (375 μm thick) were made through the medulla from the pontomedullary junction to the obex. The first slice used for recording contained the rostral VRC [36]. The next two adjacent medullary slices used for recording contained the preBötC as identified using tissue landmarks (i.e., hypoglossal nuclei were separated at the midline and the caudal extremity of the subcompact nucleus ambiguus was visible). Slices were immediately placed into an interface recording chamber (Warner Instruments, Hamden, CT, USA) and subfused with aCSF (37°C) at a rate of 8 ml/min. Slices were maintained at 37°C by an automated temperature controller (Harvard Apparatus, Holliston, MA, USA). Three 16-channel extracellular electrodes arrays (model a4x4-3μM100–177, Neuronexus, Ann Arbor, MI, USA) were placed ventrolateral to the subcompact nucleus ambiguus in VRC slices. Arrays were inserted into medullary slices at a 45° angle such that the top of the array touched the ventral border of the subcompact nucleus ambiguus. The array spanned the entire VRC and extended into tissue immediately adjacent. One array was inserted perpendicular to the cortical layers, centered on layer 3. Using this approach, multiple neurons (up to 25 neurons) were recorded from each of the three medullary slices and the cortical slice obtained from each animal in each condition. Slices were allowed to equilibrate in 7 μM KCl aCSF at 37°C with electrodes inserted for 60 min prior to recording. The following drugs were applied in our experiments: 300 μM pentobarbital (barbiturate, Fort Dodge Animal Health, Fort Dodge, Iowa, USA), 100 μM bicuculline (GABAAR antagonist, Tocris Bioscience, Ellisville, MO, USA), and 20 μM muscimol (GABAAR agonist, Tocris).

Colocalization Immunohistochemistry

The methods for immunofluorescence and image acquisition were described previously [20]. Brain slices (375 μm) used for in vitro recordings were immersion-fixed in 4.0% formaldehyde, cryoprotected with a 30% sucrose solution and sectioned coronally (30 μm). Sections were first placed in blocking solution for 1h (10% NDS in 0.01 M PBS) and then reacted with antibodies against NK1-R (1:30,000; S8305; Sigma-Aldrich, St Louis, MO, USA) and GABAAR ε subunit (1:1000; Abcam, Cambridge, MA, USA) applied overnight at room temperature in blocking solution and 0.3% Triton X-100. Sections were exposed to a fluorescent secondary antibody (1:300; Invitrogen, Carlsbad, CA, USA), mounted and coverslipped. Images of the entire slice were acquired at 4x and 10x. Images of the medullary ventrolateral quadrant were acquired bilaterally at 40x. There were no labeled cells in negative control slices from all conditions. A tyramine signal amplification kit (Perkin Elmer, Waltham, MA, USA) was used to diminish
false-positive label between antibodies. Sections were treated according to manufacturer’s recommendations. A tyramine amplified primary antibody against NK1-R (1:30,000; Sigma-Aldrich, S8305) was applied. A biotin-SP-conjugated secondary was applied (1:200; Jackson ImmunoResearch, West Grove, PA, USA) prior to the tyramine reagent (Perkin Elmer, Waltham, MA, USA). Sections were then washed and blocked again, before the application of a primary antibody against GABA<sub>ε</sub> subunit as described above (1:1000; Abcam 35971, produced against a synthetic peptide immunogen corresponding to human GABA<sub>ε</sub> subunit intracellular amino acids 237–286). Negative controls included sections reacted without primary antibodies, sections reacted without secondary antibodies, and sections reacted without tyramine signal amplification of the NK1-R primary antibody. Negative controls demonstrated specificity of antibody labeling. There were no labeled cells in control sections lacking primary antibodies.

**Image Acquisition and Analysis**

Images were acquired during the same session using an Olympus Fluoview 500 laser-scanning confocal system (Tokyo, Japan) mounted on an AX-70 upright microscope. Images were analyzed using Image J software (W. Rasband, National Institutes of Health, Bethesda, MD, USA). Images were scanned with different wavelengths sequentially to prevent bleed-through across channels. User-defined thresholds were applied uniformly to all images to measure the average pixel intensity, the number of particles, and the area of particles. Total and average fluorescence was measured on cell-by-cell basis. Total intensity was calculated as the mean cell intensity times the number of cells. All positive cells more than 200 µm from the ventral edge of the slice were quantified. Background fluorescence of negative control sections was subtracted from positive label. Data were normalized to immunofluorescence levels in virgin female rats. For figures, images were uniformly processed in Adobe Photoshop (Adobe Systems Incorporated, San Jose, CA). Images (350x350 µm) were centered in the VRC based on anatomical landmarks (i.e., the subcompact nucleus ambiguous was visible under brightfield conditions and demonstrated GABA<sub>ε</sub> subunit immunoreactivity). The viewfinder was centered ventral to the subcompact nucleus ambiguous, one third of the distance between the ventral edge of the slice and the subcompact nucleus ambiguous. Consistent microscope and laser settings were applied for the collection of each image.

**Statistical Analyses**

For statistical analyses, R was used with function “lmer” in package “lme4” (R foundation for statistical computing, Vienna, Austria). Data were analyzed with a mixed effect linear model using experimental condition as a fixed effect. Overall group differences were tested with an F test. Subsequent post hoc comparisons were performed with a Wald T-test. Differences were considered significant if p<0.05. Data are reported as means +/- S.E.M. Categorical data (hypercapnic hypoxic challenge) were analyzed with a Pearson’s Chi-Square test (R).

**Quantitative Polymerase Chain Reaction (qPCR)**

A total of 17 rats were used for qPCR experiments, including 30d isoflurane- treated rats (n = 5, 3 male, 2 female) and untreated rats (n = 12, 6 male and 6 female). Cortical and whole medullary samples contained the regions described in in vitro recording methods. Tissue was homogenized in Tri- Reagent (Sigma-Aldrich, St. Louis, MO, USA), and total RNA was collected according to the manufacturer’s protocol. Reverse transcription PCR (RT-PCR) was conducted using 1 µg of total RNA as a template for the reverse transcription reaction with a combination of oligo dT and random hexamer primers and ImProm-II Reverse Transcriptase (Promega, New York, NY).
Quantitative RT-PCR was conducted by monitoring increases in fluorescence of SYBR-GREEN dye in real-time using the TaqMan 7300 Sequence Detection System (Applied Biosystems, Carlsbad, CA, USA). Using the comparative delta delta CT method, the relative quantities of each gene transcript were measured [37]. CT values were normalized to the levels of 18s in each sample. The following primer sequences (rattus norvegicus) were used in this study:

18s F: 5′-AAC GAG ACT CTC GGC ATG CTA A-3′
18s R: 5′-CCG GAC ATC TAA GGG CAT CA-3′
Epsilon F: 5′-TGG AGC CTC AGC CTA GTG GAA AGA-3′
Epsilon R: 5′-GGC GCA GTT TAT GGT CGT AGT TGC-3′

After the final amplification cycle, a dissociation curve was generated to ensure that a single gene product was amplified. Delta CT values were used for statistical analyses. Data were analyzed in R with a linear model with experimental condition as a fixed effect. For analysis of variance, function “lme” in package “nlme” was used (R foundation for statistical computing, Vienna, Austria). Post hoc analyses were conducted with a Tukey’s HSD test. Data are graphed as fold-change relative to respective untreated regional controls.

Results
Isoflurane Treatment Increased Resistance to Pentobarbital-Dependent Frequency Depression

To test whether chronic intermittent exposure to isoflurane was sufficient to induce a compensatory increase in ε subunit-containing GABAARs in respiratory neurons, rats were pre-treated with 5 min of 3% isoflurane per day for 7 or 30 days before testing the effects of pentobarbital on respiratory motor output. Four groups of rats were used to examine changes in phrenic nerve sensitivity to pentobarbital: 1) untreated male and female rats, 2) male rats treated for 7d with intermittent isoflurane, 3) virgin female rats treated for 7d with intermittent isoflurane, and 4) virgin female rats treated for 30d with intermittent isoflurane. Phrenic nerve burst insensitivity to pentobarbital was maximally elicited at 7d in both males and females and showed no statistical differences between 7d and 30d treatments, therefore we grouped phrenic nerve recordings into treated and untreated categories for statistical comparisons.

A dose response analysis to pentobarbital was performed using untreated (n = 6 male and 6 female) and treated (n = 4 male and 8 female) rats (Fig. 1). For untreated control male and female rats, phrenic burst frequency and amplitude decreased during sequential pentobarbital injections (top two traces in Fig. 1A) with all control rats not producing phrenic bursts by the sixth injection (60 mg/kg administered). The Hill slope of the burst frequency IC50 curve was −6.0 ± 1.7 (Fig. 1B), and the IC50 was 39.9 mg/kg (95% CI 36.0 to 44.2 mg/kg).

Following isoflurane treatment, rats continued to produce measurable phrenic motor bursts under higher doses of pentobarbital than control animals. One animal continued to produce phrenic output at the maximal dose (100 mg/kg; lowest trace in Fig. 1A). The mean dose to silence phrenic output in isoflurane-treated rats was 53 ± 7 mg/kg versus 38 ± 2 mg/kg in untreated control rats (p < 0.05). The Hill slope increased to −11.4 ± 9.1 (Fig. 1B) and the burst frequency IC50 significantly increased to 60.3 mg/kg (95% CI 51.9 to 70.0 mg/kg; p < 0.01). To normalize the lethal dose of pentobarbital across rats, individual approximate Hill slopes were calculated for burst frequency and averaged for treated and untreated groups. This revealed that the slope in isoflurane-treated rats (−218.4 ± 71.1) was significantly steeper than in untreated control rats (−57.6 ± 30.4, p < 0.05).

In contrast to the pentobarbital-dependent decrease in phrenic burst frequency, the amplitude of phrenic nerve output was highly variable following pentobarbital application. The
parameters defining the best-fit models for the phrenic nerve amplitude dose response curve did not differ by condition (p = 0.35; Fig. 1C), thus one IC50 (19.1 mg/kg; 95% CI 17.3 to 21.1 mg/kg) and one Hill slope (−1.68 ± 0.15) described the entire amplitude dataset. These data indicate that following isoflurane treatment, pentobarbital was still inhibitory on neurons contributing to phrenic nerve amplitude but not burst frequency.

To assess whether respiratory networks were truly silenced by 100 mg/kg pentobarbital, we sought to provide maximal stimulus for breathing to prompt the system to surmount barbiturate-induced depression. There was notable variability in the dose of pentobarbital required to silence the phrenic nerve. While ≤60 mg/kg of pentobarbital was sufficient to depress phrenic nerve activity in all untreated control rats, not all isoflurane-treated rats were depressed by the maximal dose of 100 mg/kg. To assess whether the degree of inhibition differed between groups, we challenged rats with a strong stimulus (hypercapnic-hypoxia challenge) at the end of each experiment (Fig. 1A). Six of 12 isoflurane-treated rats mounted a detectable phrenic response to the challenge, while only two of 12 untreated control rats responded (p = 0.08).
Pentobarbital Sensitivity of Spontaneous Neuronal Activity in Medulla and Cortex

During sequential pentobarbital injections, respiratory phrenic burst frequency was sustained, thereby suggesting that respiratory rhythm-generating neurons in medulla express pentobarbital-insensitive GABA\(_A\)Rs. In addition, it is not clear whether isoflurane exposures increase the expression of pentobarbital-insensitive GABA\(_A\)Rs in other parts of the CNS. To answer these questions, pentobarbital sensitivity was examined in the neural regions responsible for rhythm generation (VRC, preBötc) as well as in cortical neurons, which exhibit a GABA\(_A\)R mediated tonic current \[38\]. Four groups of rats were tested: 1) untreated male and female rats, 2) 7d isoflurane-treated male and female rats, 3) 30d isoflurane-treated male and female rats, and 4) male and female rats that were allowed 7d of recovery following 7d isoflurane treatment (data from the untreated groups are reported in \[20\]).

Consistent with prior data \[39, 40, 20\], in slices from untreated control male (n = 6) and female rats (n = 6), 60 min exposure to bath-applied 300\(\mu\)M pentobarbital inhibited spontaneous activity in VRC neurons to 44 ± 15% of baseline (n = 103 neurons, p < 0.05) and cortical (CTX) neurons to 24 ± 8% of baseline (n = 84 neurons, p < 0.05) compared to time controls (n = 5 animals, 56 VRC neurons and 48 CTX neurons, 136 ± 19% and 123 ± 19% of baseline, respectively) (Fig. 2A, B). There were no significant differences between untreated male and female animals.

The 7d intermittent isoflurane treatment (n = 12 rats) reduced the inhibitory effects of 1h pentobarbital on VRC (n = 102 neurons, 91 ± 21% of baseline, p < 0.05) and CTX neurons (n = 84 neurons, 49 ± 10% of baseline, p < 0.005) compared to untreated control rats (Fig. 2C, D). There were no significant differences between male and female animals. The effects of 7d isoflurane treatment were reversible in the VRC, as 7d recovery significantly shifted the effect of 1h pentobarbital towards the response of untreated control rats (VRC, n = 27 neurons, 44 ± 14% of baseline, p < 0.01) (Fig. 2E). There was no significant difference between spontaneous activity after pentobarbital application in CTX neurons from 7d isoflurane-treated rats and 7d recovery rats.

The 30d of intermittent isoflurane treatment rendered VRC neurons insensitive to pentobarbital application (n = 163 neurons, 104% of baseline, p > 0.05 compared to time control) (Fig. 2F, G). The 30d intermittent isoflurane treatment differentially affected CTX neurons in males and females (p < 0.05). In males, CTX neurons maintained 83 ± 18% of baseline activity (n = 33 neurons) (Fig. 2F) while in females, CTX neurons maintained 48 ± 9% of baseline activity (Fig. 2G). In both males and females, CTX neurons were significantly less inhibited by pentobarbital after 30d of intermittent isoflurane than untreated animals (female, p < 0.05; male, p < 0.005).

The persistence of spontaneous neuronal activity following pentobarbital application to the CTX and VRC of 30d treated animals could reflect either a lack of GABA\(_A\)Rs on neurons, or the presence of pentobarbital-insensitive GABA\(_A\)Rs. To address this question, 20\(\mu\)M muscimol (GABA\(_A\)R agonist) was bath-applied to from slices from isoflurane-treated rats after 1h of 300\(\mu\)M pentobarbital (Fig. 2F, G). After 1h of exposure to 300\(\mu\)M pentobarbital, CTX and VRC neurons from isoflurane-treated rats were rapidly (<5 min) inhibited by muscimol, confirming the presence of functional, pentobarbital-insensitive GABA\(_A\)Rs. The response of CTX and VRC neurons to muscimol did not differ between isoflurane-treated rats and time control rats (p > 0.05), indicating the presence of pentobarbital-insensitive GABA\(_A\)Rs in CTX and VRC after 30d of isoflurane.

The inhibitory effects of pentobarbital in CTX and VRC neurons were mediated by GABA\(_A\)Rs, as bath application of 100\(\mu\)M bicuculline (GABA\(_A\)R antagonist) to slices (n = 4...
Fig 2. Spontaneous neuronal activity resistance to bath-applied pentobarbital in medullary and cortical slices from 7d isoflurane-treated rats. Spontaneous activity of neurons in the region anatomically consistent with the VRC and PreBötC (all labeled as VRC neurons) are insensitive to bath-applied pentobarbital (300 μM) following 7d isoflurane treatment. A, Mean normalized firing rate of VRC (n = 62, open circles) and CTX (n = 37, filled circles) neurons from male rats in response to 300 μM pentobarbital (applied at vertical bar, t = 30 min). B, Pentobarbital had equivalent depressive effects on VRC (n = 37) and CTX (n = 47) neurons from virgin female rats. C, Following pentobarbital treatment, spontaneous activity of VRC neurons (n = 62) from male rats treated with intermittent isoflurane for 7d were significantly more active than VRC neurons from untreated control rats. The response of cortical neurons (n = 49) to pentobarbital was non-
rats; 2 male, 2 female) prior to co-application of 100 μM bicuculline and 300 μM pentobarbital prevented the inhibition of spontaneous activity of both CTX and VRC neurons by pentobarbital observed in virgin female and male slices (data not shown).

A summary of the steady-state pentobarbital responses in VRC and CTX neurons in slices from male and female rats shows that isoflurane exposures increased pentobarbital resistance after 7d and 30d (Fig. 2H, 2I). The response in female rats was greater compared to male rats. Together, these data suggest that intermittent isoflurane treatment is sufficient to increase the expression of a pentobarbital-insensitive GABAAR subtype in VRC and CTX neurons.

Immunohistochemical Localization of ε Subunit Expression in Medulla and Cortex

To confirm the presence or absence of GABAAR ε subunit expression at the in vitro recording sites, medullary slices from recording experiments were treated with an antibody against the GABAAR ε subunit. Medullary sections were co-labeled for GABAAR ε subunit and for NK1-R (Substance P receptor) that is used as a marker for putative respiratory neurons in the VRC and preBöTc [41, 42]. NK1-R and the GABAAR ε subunit exhibited >90% colocalization, both in the nucleus ambiguus (subcompact and compact; Lin et al., 2008), which was used as a landmark for electrode placement, and in the VRC (Fig. 3A; [36]). Neurons in the VRC displayed enriched somatic ε subunit staining as well as in primary neuronal processes (Fig. 3A). The pentobarbital-insensitive GABAARs identified in vitro were functionally consistent with immunohistochemically identified ε subunit-containing GABAARs in the VRC. To our surprise, the GABAAR ε subunit was detectable in CTX neurons in all groups as well. Layer 5 pyramidal cells demonstrated the most robust expression, and staining appeared to be largely restricted to the soma (Fig. 3B).

ε Subunit mRNA Expression in Medulla and Cortex

To quantify whether intermittent isoflurane treatment altered the transcriptional regulation of the GABAAR ε subunit, cortical and medullary samples from 30d isoflurane-treated rats (n = 5) and untreated control rats (n = 12) were harvested for GABAAR ε subunit mRNA analysis (Fig. 4). The 30d isoflurane treatment increased GABAAR ε subunit mRNA in medullary samples compared to untreated samples (3.9 ± 1.39-fold increase, p<0.05). Cortical samples from 30d isoflurane-treated rats also demonstrated increased GABAAR ε subunit mRNA levels compared to controls (4.2 ± 1.35-fold increase, p<0.05, Wilcoxon Rank Sum). These data demonstrate a correlation between the emergence of barbiturate-insensitive GABAARs in VRC and cortex and the transcriptional upregulation of GABAAR ε subunit mRNA.
Multiple forms of neuronal plasticity must be carefully coordinated to produce a stable and functional nervous system. The mechanisms that are actively expressed are regulated both developmentally and spatially, and depend upon external and internal environments. This study shows that repeated exposure to brief episodes of an inhaled anesthetic is sufficient to functionally reorganize medullary and cortical neuronal circuitry. Following 7 days of 5 min/day exposure to isoflurane, both in vivo and in vitro measures of respiratory neuronal output revealed a

**Discussion**

Multiple forms of neuronal plasticity must be carefully coordinated to produce a stable and functional nervous system. The mechanisms that are actively expressed are regulated both developmentally and spatially, and depend upon external and internal environments. This study shows that repeated exposure to brief episodes of an inhaled anesthetic is sufficient to functionally reorganize medullary and cortical neuronal circuitry. Following 7 days of 5 min/day exposure to isoflurane, both in vivo and in vitro measures of respiratory neuronal output revealed a
significantly increased barbiturate tolerance. After 30 days of treatment, cortical networks were similarly affected.

This study initially sought to test whether non-pregnant animals could be induced to increase $\epsilon$ subunit expression and barbiturate tolerance in VRC neurons. Our findings make it clear that gestational hormones are unnecessary for this effect, as both virgin female and male rats have the capacity to express this form of plasticity. We cannot rule out the effects of neurosteroids, as there is evidence that GABAergic modulators may increase neurosteroid levels in in vitro hippocampal preparations [43]. Surprisingly, our data also reveal that this phenomenon is not restricted to medullary neurons, as cortical neurons displayed a similar pharmacological profile following longer treatment periods. Whether this is a global feature of the cortex is unclear, but these data suggest that the incorporation of pharmacologically distinct GABA$_{A}$R configurations is a general mechanism of neuronal plasticity.

If this plasticity is not gated by the interaction of hormones and nuclear receptors (one of our initial hypotheses), how might this process be triggered in neurons? Long duration (minutes to hours) application of GABA to cultured neurons leads to an increase in receptor

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**Fig 4. Increased GABA$_{A}$R $\epsilon$ subunit mRNA levels in medulla and cortex in 30d isoflurane-treated rats.**

Medullary GABA$_{A}$R $\epsilon$ mRNA levels were more than 3-fold greater in 30d isoflurane-treated rats compared to untreated control rats. Likewise, cortical GABA$_{A}$R $\epsilon$ mRNA levels were more than 4-fold increased in 30d isoflurane-treated rats. Statistics were conducted on delta CT values. Asterisk denotes $p<0.05$. Error bars indicate SEM.

doi:10.1371/journal.pone.0119351.g004
internalization, an uncoupling of the allosteric modulatory site from the GABA binding site, and a decrease in the expression of specific subunits [33]. The mechanism linking increased allosteric modulatory pressure and the regulation of the ε subunit is unclear. This will be a critical question to answer in future studies.

The demonstration that the frequency of phrenic nerve output in vivo is highly resistant to sequential pentobarbital injections in isoflurane-treated rats strongly suggests that rhythm-generating neurons in the brainstem increase ε subunit expression in GABA\(_\alpha\)Rs. Since it is hypothesized that inspiratory activity produced mainly by preBötC neurons in the VRC [44, 34, 35], pentobarbital sensitivity of spontaneous neuronal activity was tested in medullary slices from control and isoflurane-treated rats. None of the recorded neurons in the medullary slice experiments could be defined as being respiratory-related because adult medullary slices do not produce spontaneous respiratory rhythm-generating motor activity similar to that produced in neonatal rodent medullary slice [44]. However, it is likely most of the recorded neurons were respiratory-related preBötC neurons because the preBötC region has distinctive surrounding neural landmarks that define the region and are easily visualized in medullary slices in vitro. Furthermore, ε subunit expression was increased in NK1-R-positive neurons within the preBötC region in isoflurane-treated rats. Since the NK1-R is one of several markers for preBötC neurons [45], this finding is consistent with the hypothesis that ε subunit expression increased in preBötC neurons to compensate for the isoflurane-dependent decrease in respiratory motor activity. This study does not address the question as to whether intermittent isoflurane exposures altered ε subunit expression in other respiratory-related brainstem regions controlling expiration, postinspiration, or central chemosensation. Furthermore, modulatory neurons (e.g., serotonergic, noradrenergic, dopaminergic) projecting to the preBötC may also have increased expression of ε subunits in GABA\(_\alpha\)Rs, and thereby contributed to the altered responsiveness of the respiratory control network to pentobarbital.

The in vivo and slice physiology data presented here provide measures of network activity as opposed to cell autonomous responses. Our data demonstrate the presence of pentobarbital insensitive GABA\(_\alpha\)Rs in the cortex and the brainstem, but it is difficult to predict specific expression patterns from these experiments. For example, the transient increases in cortical neuronal activity of 30d isoflurane treated male rats (Fig. 2F) likely represent pentobarbital-mediated inhibition of a subclass of inhibitory interneuron, resulting in disinhibition of excitatory cells. Subunit distribution rules will be important to understand in the future.

ε subunit-containing GABA\(_\alpha\)Rs are insensitive to most allosteric positive modulators [18, 19]; both the human and rat isoform of the ε subunit confers anesthetic insensitivity to neurons [46], and GABA\(_\alpha\)Rs containing the ε subunit are upregulated on VRC neurons during pregnancy when pentobarbital sensitivity decreases [20]. Our data fall short of establishing a mechanistic link between the induced expression of ε subunit-containing GABA\(_\alpha\)Rs and barbiturate insensitivity. We do, however, demonstrate the conditional presence of fully functional, barbiturate-insensitive GABA\(_\alpha\)Rs in brain regions that are immunohistologically enriched with the ε subunit and show increased ε subunit mRNA transcription following treatment. Furthermore, these receptors are rapidly inhibited by muscimol, and the inhibitory actions of pentobarbital are blocked by bicuculline (indicating the involvement of GABA\(_\alpha\)Rs). These data are difficult to understand without the specific pharmacological properties imbued by the ε subunit, thus this is the most likely explanation for our findings. However, because the connection between the ε subunit and the respiratory plasticity described here is correlative, it will take further investigation to unequivocally confirm a molecular mechanism.

Cortical networks have the capacity to homeostatically regulate gain control via plasticity of intrinsic excitability and changes in synaptic strength such that network activity is maintained following perturbations in drive [24, 47, 48]. Similarly, precise balancing of respiratory
neuronal input/output ratio is necessary for survival, and gain modulation of respiratory rhythm generating neurons is provided by a tonic GABA_A current [49]. Neurosteroids, like barbiturates, act on these currents and suppress activity in medullary slices from naïve animals [50]. Thus, these data suggest a compensatory mechanism of gain modulation that counteracts the hyperpolarizing properties of chronic exposure to allosteric positive GABA_A receptors, such as neurosteroids, ethanol, and in some cases, anesthetics. Without a compensatory response to allosteric modulators, respiratory pathologies would arise as a result of pregnancy, regular alcohol consumption, or multiple exposures to anesthetics. To our knowledge, these are the first data to demonstrate reorganization of the neural control of respiration by repeated anesthetic exposure. Prior data demonstrate the ability of anesthetics to alter GABA_A receptor expression patterns. Sekine et al. (2006) described changes in forebrain mRNA for the GABA_A receptor α_4 subunit during anesthesia with propofol or isoflurane [51]. Similar to our findings, chronic ethanol exposure is associated with a cross-tolerance to many GABAergic anesthetics [26, 52, 53]. It is likely that cross-tolerance is a feature of tonic GABA_A receptor plasticity, as we have previously described VRC insensitivity to both ETOH and barbiturates in hibernating ground squirrels [40].

**Significance**

This study presents a novel, clinically relevant, inducible model of neuroplasticity that was clearly demonstrated in an intact animal with measurable, functional spinal respiratory motor output. We provide evidence that this isoflurane-induced neuroplasticity likely involves the expression of a relatively unknown, yet highly conserved, GABA_A receptor subunit that confers insensitivity to allosteric modulation by neurosteroids, anesthetics, ethanol, barbiturates, and benzodiazepines [18, 19, 39, 40, 20]. Here we demonstrate that this plasticity is restricted neither to specific classes of animals (i.e. pregnant or hibernating), nor respiratory neurons. These data raise the possibility that this type of respiratory neuroplasticity, which mirrors some of the changes associated with hibernation and pregnancy, is inducible and may be harnessed to treat pathological conditions.

Many long-term in vivo preparations in basic neurobiology research rely on regular administration of isoflurane and other GABAergic anesthetics in order to prepare an animal for data collection. The findings presented here suggest that these protocols may be inadvertently manipulating the very networks being studied. The impact of these methods needs further study.

**Author Contributions**

Conceived and designed the experiments: KBH SMJ GSM MB. Performed the experiments: KBH NRN KMS SMS. Analyzed the data: KBH NRN SMS. Contributed reagents/materials/analysis tools: JJW SMJ MB GSM. Wrote the paper: KBH.

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