


Postictal Death Is Associated with Tonic Phase Apnea in a Mouse Model of Sudden Unexpected Death in Epilepsy

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Objective: Sudden unexpected death in epilepsy (SUDEP) is an unpredictable and devastating comorbidity of epilepsy that is believed to be due to cardiorespiratory failure immediately after generalized convulsive seizures.

Methods: We performed cardiorespiratory monitoring of seizure-induced death in mice carrying either a p.Arg1872Trp or p.Asn1768Asp mutation in a single *Scn8a* allele—mutations identified from patients who died from SUDEP—and of seizure-induced death in pentylentetrazole-treated wild-type mice.

Results: The primary cause of seizure-induced death for all mice was apnea, as (1) apnea began during a seizure and continued for tens of minutes until terminal asystole, and (2) death was prevented by mechanical ventilation. Fatal seizures always included a tonic phase that was coincident with apnea. This tonic phase apnea was not sufficient to produce death, as it also occurred during many nonfatal seizures; however, all seizures that were fatal had tonic phase apnea. We also made the novel observation that continuous tonic diaphragm contraction occurred during tonic phase apnea, which likely contributes to apnea by preventing exhalation, and this was only fatal when breathing did not resume after the tonic phase ended. Finally, recorded seizures from a patient with developmental epileptic encephalopathy with a previously undocumented *SCN8A* likely pathogenic variant (p.Leu257Val) revealed similarities to those of the mice, namely, an extended tonic phase that was accompanied by apnea.

Interpretation: We conclude that apnea coincident with the tonic phase of a seizure, and subsequent failure to resume breathing, are the determining events that cause seizure-induced death in *Scn8a* mutant mice.

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Sudden unexpected death in epilepsy (SUDEP) is defined as the sudden, unexpected, nontraumatic, nondrowning death of a person with epilepsy, which is not due to *status epilepticus*, nor does a postmortem examination reveal another cause of death.¹ SUDEP is the most common cause of death associated with epilepsy, accounting for between 8 and 17% of all epilepsy-related deaths.² This number increases to 50% for patients whose seizures are refractory to treatment.^{3,4}

Although numerous mechanisms have been proposed for SUDEP,^{3,5,6} a growing body of evidence supports postictal respiratory dysfunction as the primary cause of death for many cases. Apnea and oxygen desaturation have been reported in a large percentage of patients during and after convulsive and nonconvulsive seizures,^{7–12} and in 9 cases of SUDEP with adequate postictal cardiorespiratory monitoring, terminal apnea occurred prior to terminal asystole.¹³ It is believed that most SUDEP cases occur

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after generalized convulsive seizures^{13–17}; thus, mouse models of SUDEP include those in which death occurs immediately after convulsive seizures. In such models, including *Cacna1a*^{S218L}, *Lmx1b*^{fl/fl}, and *Scn1a*^{R1407X} transgenic mice, apnea has also preceded terminal asystole.^{18–21} Of these, only the *Scn1a*^{R1407X} mice represent a patient population known to be susceptible to SUDEP (ie, Dravet syndrome).^{18,22}

The current study utilizes 3 transgenic mouse lines carrying gain-of-function *Scn8a* mutations, which were identified in several patients with developmental epileptic encephalopathy (DEE) who died of SUDEP^{23,24}; either the germline p.Asn1768Asp (referred to as “D/+ mice”) or conditional p.Arg1872Trp (referred to as “W/+ mice”). In the presence of *Emx1-Cre*, the conditional p.Arg1872Trp mutation is expressed in forebrain excitatory neurons (referred to as “W/+^{*Emx1-Cre*} mice”) and in the presence of *EIIA-Cre* it is expressed globally (referred to as “W/+^{*EIIA-Cre*} mice”).²⁵ Phenotypically, expression of either mutation recapitulates clinical features seen in patients, including spontaneous tonic–clonic seizures and seizure-induced death.^{25–27} Prior to the current study, peri-ictal cardiorespiratory monitoring of these pertinent mouse models had not been performed. We utilized the natural epileptogenesis of the W/+^{*Emx1-Cre*} mice to examine the semiology of spontaneous nonfatal and fatal seizures, the inducible fatal seizures of the W/+^{*EIIA-Cre*} mice to examine a causal relationship between apnea and seizure-induced death, and the inducible nonfatal seizures of the D/+ mice to observe tonic diaphragmatic contraction during the tonic phase.

We demonstrate that (1) sudden death in the *Scn8a* mutant mice and pentylenetetrazole (PTZ)-treated mice occurs after convulsive seizures in which apnea occurred coincident with the tonic phase; (2) death can be prevented by mechanical ventilation; (3) tonic phase apnea is necessary, but not sufficient, to produce seizure-induced death; (4) tonic diaphragm contraction may contribute to the fatal apnea; and (5) seizures recorded from a DEE patient with a novel *SCN8A* mutation also presented with apnea during the tonic phase.

Materials and Methods

Mutant Mice

All mice were housed and cared for in accordance with NIH guidelines in a 12-hour light/dark cycle with food and water ad libitum, and all procedures were carried out at the University of Virginia and approved by the Animal Care and Use Committee. Male mice homozygous for the conditional R1872W *Scn8a* mutant allele²⁵ (Fig 1A) were crossed with female mice homozygous for either *Emx1-Cre* (Jax 005628)²⁸ or *EIIA-Cre* (Jax 003724)²⁹ to produce mice heterozygous for both alleles,

resulting in expression of the R1872W mutation in forebrain excitatory neurons (W/+^{*Emx1-Cre*}) or globally (W/+^{*EIIA-Cre*}); see Fig 1B). Mice with germline N1768D *Scn8a* mutant allele were also used.²⁷ Genotyping was performed to confirm the presence of the R1872W and N1768D mutant alleles as described previously,^{25,27} and for *Emx1-Cre* and *EIIA-Cre* alleles as directed by the Jackson Laboratories website. Male and female W/+^{*Emx1-Cre*} and wild-type (WT) mice were used with no difference in seizure phenotype. Due to the preweaning age of the W/+^{*EIIA-Cre*} mice, sex was not recorded.

Continuous Recording of Electroencephalogram, Electrocardiogram/Electromyogram, and Breathing

As previously reported, the W/+^{*Emx1-Cre*} mice develop spontaneous tonic–clonic seizures at 4 weeks of age, and 75% succumb to seizure-induced death by 2 months.²⁵ D/+ mice have a milder phenotype, with seizure onset at 8 to 16 weeks and only 50% lethality by 6 months.²⁷ Custom headsets (Plastics One, Roanoke, VA) were implanted in 4- to 5-week-old W/+^{*Emx1-Cre*} and 8- to 10-week-old D/+ mice using approved surgical techniques as previously described by our laboratories.^{18,25} Briefly, electroencephalographic (EEG) electrodes were placed over the parietal lobe unilaterally, and electrocardiographic (ECG) electrodes were placed in an approximate lead II configuration. In a subset of D/+ mice, ECG leads were replaced with leads that extended to contact the diaphragm, so as to record diaphragm electromyogram (EMG) as previously described.³⁰ A reference electrode was placed over the cerebellum. Normal saline (8ml/kg, intraperitoneal [i.p.]) and ketoprofen (5mg/kg, i.p.) were given postoperatively. Mice were monitored, and additional ketoprofen was administered every 24 hours as needed. After recovery from surgery (minimum of 4 days), mice underwent continuous monitoring of EEG, ECG/EMG, and breathing in custom-designed plethysmography chambers that comply with guidelines for continuous housing described in the Guide for the Care and Use of Laboratory Animals,³¹ and as described previously.¹⁸ The same techniques were used for recording seizures from W/+^{*EIIA-Cre*} mice at postnatal age 15 days. However, because W/+^{*EIIA-Cre*} mice were used prior to weaning, due to their early lethality, they could only be allowed 24 hours to recover from surgery, because degradation of ECG signals was common due to parental grooming.

Audiogenic seizures were induced in W/+^{*EIIA-Cre*} and D/+ mice with a 15kHz pure tone (–80–100dB), generated using Tone Generator software (NCH Software, Canberra, Australia), amplified with a Kinter K3118 stereo amplifier (Kinter USA, Amazon.com), and converted to sound using a small 3W speaker lowered into the plethysmography chamber. For mechanical ventilation experiments, a sonicator (Branson 200 ultrasonic cleaner) was placed next to the mouse cage and turned on to induce audiogenic seizures (see Video S5).

PTZ-induced seizures were generated by i.p. injection of 90mg/kg PTZ, which resulted in 100% lethality, as previously reported.^{32,33} PTZ (Sigma-Aldrich, St Louis, MO) was dissolved in normal sterile saline on the day of administration.

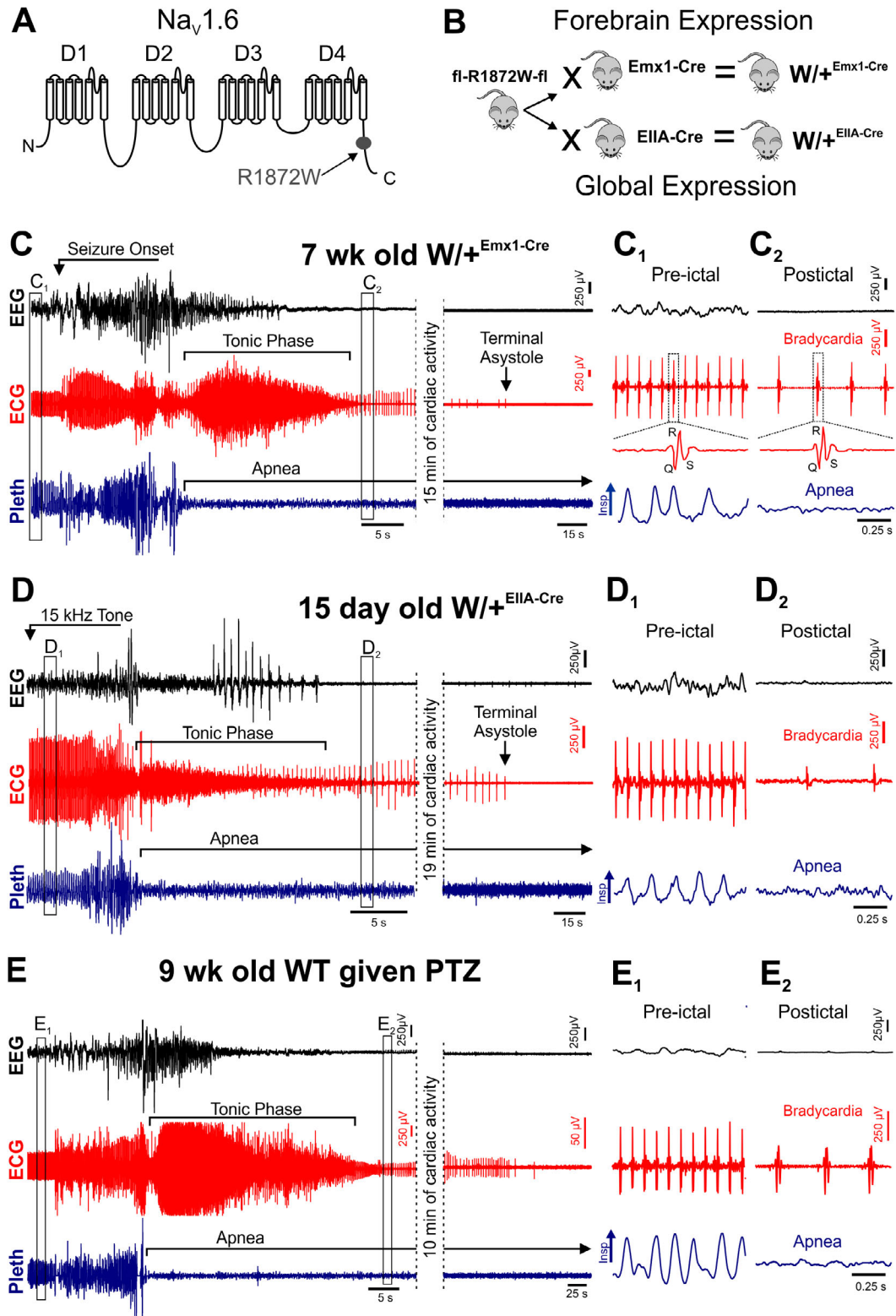


FIGURE 1: Seizure-induced death occurs when breathing does not resume after the tonic phase. (A) Diagram of SCN8A protein structure with location of R1872W mutation depicted. (B) Breeding scheme to generate W⁺/Emx1-Cre (top) and W⁺/EIIA-Cre (bottom) mice. Mice homozygous for the floxed R1872W mutant allele were bred with mice homozygous for Emx1-Cre or EIIA-Cre. The resulting offspring were heterozygous for both alleles. (C) Representative traces showing time from seizure onset to terminal asystole. (C–E) Recording of electroencephalogram (EEG), electrocardiogram (ECG), and plethysmogram (Pleth) during fatal spontaneous seizures of W⁺/Emx1-Cre mice (C), fatal audiogenic seizures of W⁺/EIIA-Cre mice (D), and fatal pentylenetetrazole (PTZ)-induced seizures of wild-type (WT) mice (E). (C₁, C₂) Expanded traces from corresponding boxes in C. (D₁, D₂) Expanded traces from corresponding boxes in D. (E₁, E₂) Expanded traces from corresponding boxes in E. Insp = inspiration.

SCN8A Patient Video-EEG Monitoring and Breathing Analysis

Human studies were approved by the University of Virginia Institutional Review Board and performed under supervision of a faculty neurologist (H.P.G.). A patient with a pathogenic variant of *SCN8A* was admitted to the University of Virginia Medical Center (Charlottesville, VA) for continuous video-EEG scalp recordings (Natus, Pleasanton, CA) with electrodes placed according to the 10–20 system. Breathing movement was visualized using phase-based Eulerian video magnification motion processing with open-source software (<http://people.csail.mit.edu/mrub/vidmag/>) written for MATLAB (MathWorks, Natick, MA) as done previously.¹⁸

Statistical Analysis and Blinding

Because only mice expressing *Scn8a* mutations were used in this study, blinding to genotype was not necessary. For the quantification in Figure 5, measurements of tonic phase and apnea were done independent of one another. Quantification reported in Figure 7 was done with the researcher blinded to seizure outcome (ie, fatal or nonfatal). All statistics were performed using Prism 8 (GraphPad Software, San Diego, CA), presented as mean \pm standard error of the mean, and defined as statistically detectable when $p < 0.05$. Comparison of survival proportions in Figures 2 and 4 was done using a 1-sided Fisher exact test. Comparisons between 2 groups were assessed by unpaired, 2-tailed Student *t*-test when residuals passed the D'Agostino-Pearson Omnibus normality test and by the nonparametric Mann-Whitney test when residuals did not pass the D'Agostino-Pearson Omnibus normality test. Comparisons between more than 2 groups were assessed by 1-way analysis of variance followed by Dunnett multiple comparison test when residuals passed the D'Agostino-Pearson Omnibus normality test and by the nonparametric Kruskal-Wallis test for those that did not pass the D'Agostino-Pearson Omnibus normality test. Details for statistical comparisons that are graphically depicted in figures are described in the figure legends. For comparisons only mentioned in the results section, statistics are reported in line.

Results

Monitoring Seizures and Sudden Death in *Scn8a* Mutant Mice

Eight $W/+^{Emx1-Cre}$ mice were monitored for ECG, cortical EEG, and whole-body plethysmograph, which captured 44 nonfatal and 8 fatal seizures. Only seizures qualifying as R5 or above (ie, R5–R8^{32,33}) on a modified Racine scale were analyzed. These showed rearing and falling, tail-raising, clonic and/or tonic convulsions, and increased ictal EEG amplitude followed by postictal generalized EEG suppression (PGES). Seizures were always separated by at least 1 hour, and never qualified as *status epilepticus*. Three subcategories of R5 or greater seizures were identified in the $W/+^{Emx1-Cre}$ mice: fatal tonic seizures (ie, seizure-induced death; Video S1), nonfatal tonic seizures (Video S2), and nonfatal clonic seizures (Video S3). Fatal tonic seizures were recorded from 8 $W/+^{Emx1-Cre}$ mice. A total of 44 spontaneous nonfatal seizures were recorded from 6 of the 8 $W/+^{Emx1-Cre}$ mice, of which 15 were tonic and 29 were clonic.

Four $W/+^{EIIA-Cre}$ mice were monitored for ECG, EEG, and plethysmography during audiogenic seizure-induced death. We found that high-frequency sound routinely caused seizures in $W/+^{EIIA-Cre}$ mice at 14–18 days of age. In all cases, these seizures resulted in sudden death with the same sequence of events as those experienced by $W/+^{Emx1-Cre}$ mice during spontaneous seizures. All $W/+^{EIIA-Cre}$ mice had convulsive seizures that included wild running followed by a tonic phase and death. The audio stimulus never induced observable seizure behavior in $W/+^{Emx1-Cre}$ or WT mice (data not shown).

To compare results obtained from the *Scn8a* mutant mice to a more widely used model of seizure-induced death, we monitored 7 WT (C57/B6J) mice for ECG, EEG, and plethysmography immediately before and after

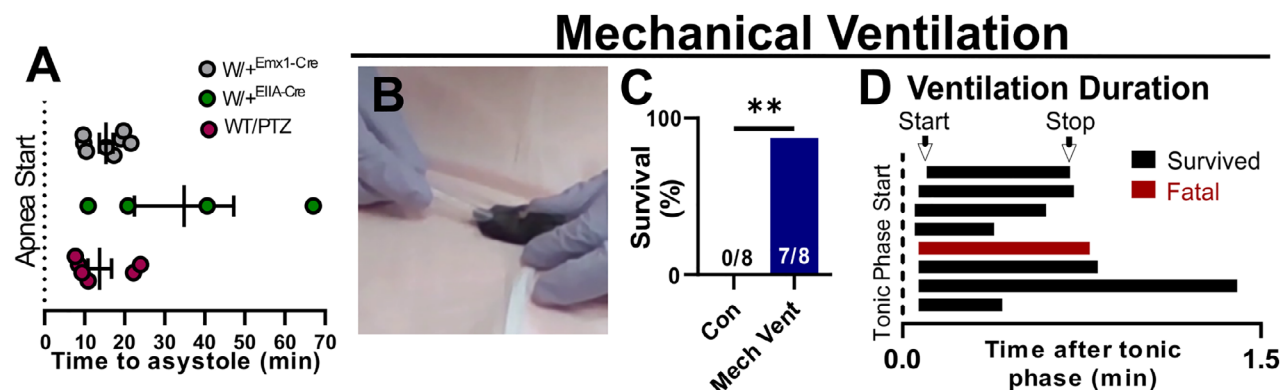


FIGURE 2: Apnea is the determining cause of seizure-induced death. (A) In the event of seizure-induced death, terminal apnea preceded terminal asystole in all mouse models. (B, C) Mechanical ventilation significantly increased survival of audiogenic seizures for $W/+^{EIIA-Cre}$ mice (** $p = 0.0205$, 2-sided chi-squared = 5.367). (D) For each mouse, ventilation was initiated within 6 seconds of the beginning of the tonic phase and was continued for 21 to 80 seconds. Con = control; PTZ = pentylenetetrazole; WT = wild type.

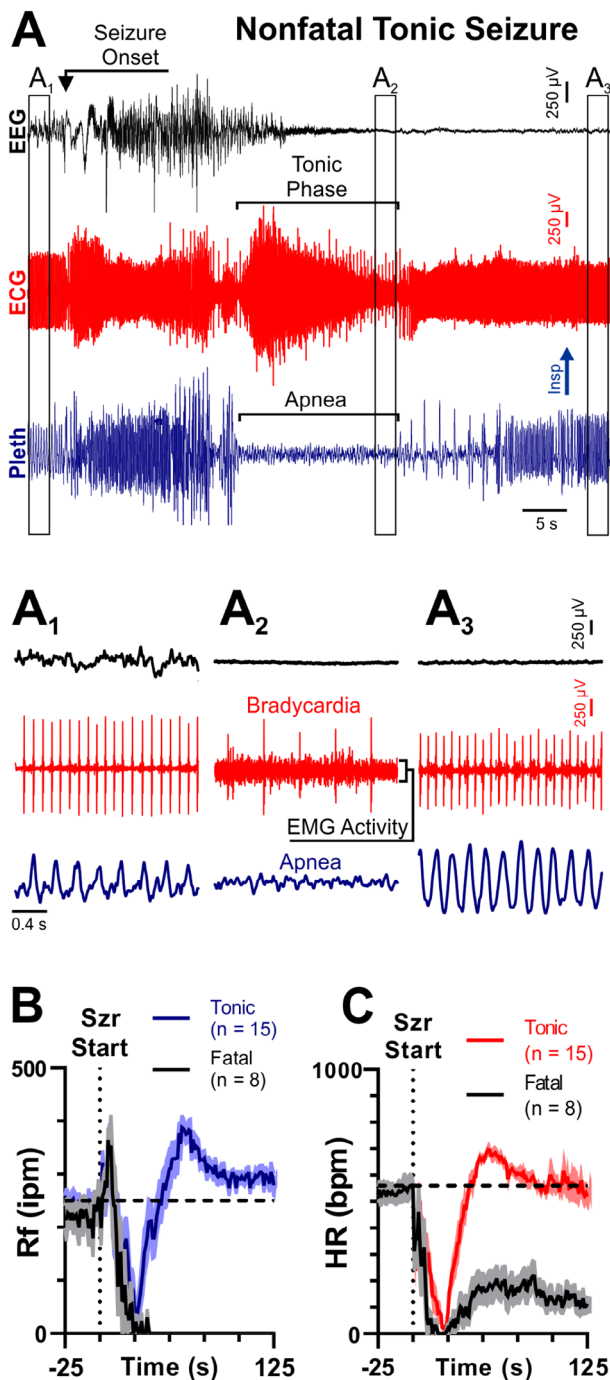


FIGURE 3: Transient apnea and bradycardia also occur during nonfatal tonic seizures. (A) Recording of electroencephalogram (EEG), electrocardiogram (ECG), and plethysmogram (Pleth) during a convulsive seizure in a $W^{+/Emx1-Cre}$ mouse. (A₁–A₃) Expanded views of traces from A, depicting EEG, ECG, and breathing at different time points. In A₂, note the bradycardia, electromyographic (EMG) infiltration of the ECG, and apnea. (B, C) Respiratory frequency (Rf) and heart rate (HR) binned every second from 50 seconds before to 150 seconds after seizure (Szr) initiation for all tonic (red/blue) and fatal (black) seizures. For B and C, solid lines represent mean, and shaded regions represent standard error of the mean. Note that in nonfatal seizures, both breathing and HR increase beyond resting levels immediately after the seizure. bpm = beats per minute; Insp = inspiration; ipm = inspirations per minute.

i.p. injection of 90mg/kg PTZ. All mice died from convulsive seizures with similar semiology to the fatal seizures of the *Scn8a* mutant mice, which included wild running followed by a tonic phase and death (Video S4).

Finally, 3 D/+ mice were monitored for ECG, EEG, and plethysmography during 14 spontaneously occurring nonfatal tonic seizures. Three additional D/+ mice had EEG, plethysmography, and diaphragm EMG measured (as previously described³⁰) during audiogenic nonfatal tonic seizures. As per our previous report, both spontaneous and audiogenic seizures in D/+ mice were characterized by wild running followed by a tonic phase and recovery (manuscript under review).

Apnea Causes Sudden Death in *Scn8a* Mutant Mice

In all cases of sudden death, breathing was absent during the tonic phase of a convulsive seizure and never resumed (see Fig 1C–E). During and after the fatal seizure, cardiac activity continued, albeit at a lower rate (see Fig 1C₂, D₂, and E₂). For $W^{+/Emx1-Cre}$, $W^{+/EIIA-Cre}$, and PTZ-treated WT mice, terminal asystole followed breathing cessation by 15.4 ± 1.7 , 34.4 ± 12.4 , and 13.8 ± 3.0 minutes, respectively (Fig 2A). Thus, the sequence of events did not differ when the *Scn8a* mutation was expressed globally ($W^{+/EIIA-Cre}$ mice) or selectively in forebrain neurons ($W^{+/Emx1-Cre}$ mice), or exogenously induced in otherwise genetically WT mice (ie, PTZ-treated WT mice).

Based on these observations, we hypothesized that apnea was the primary cause of death. To directly test this hypothesis, we induced audiogenic seizures in $W^{+/EIIA-Cre}$ mice that routinely resulted in death (0/8 survived with no intervention; Video S5, left side). When we induced audiogenic seizures in $W^{+/EIIA-Cre}$ mice and performed mechanical ventilation using a transfer pipet adapted to fit snugly around the nose and mouth (see Fig 2B; Video S5, right side), survival markedly improved (7/8 survived when ventilated; see Fig 2C). Ventilation was initiated within 6 seconds after the start of the tonic phase (see Fig 2D) and continued until the mouse began breathing on its own. The one mouse that did not survive was ventilated for 43 seconds before stopping due to lack of recovery. Because another mouse that survived was ventilated for 80 seconds, the single fatality may have resulted from premature termination of mechanical ventilation.

Tonic Phase Apnea Is Not Sufficient for Seizure-Induced Death

Nonfatal tonic seizures in $W^{+/Emx1-Cre}$ mice appeared nearly identical to fatal seizures, except that breathing resumed immediately postictally (compare Fig 3A to Fig 1C, D; Videos S1 and S2). Nonfatal seizures had a

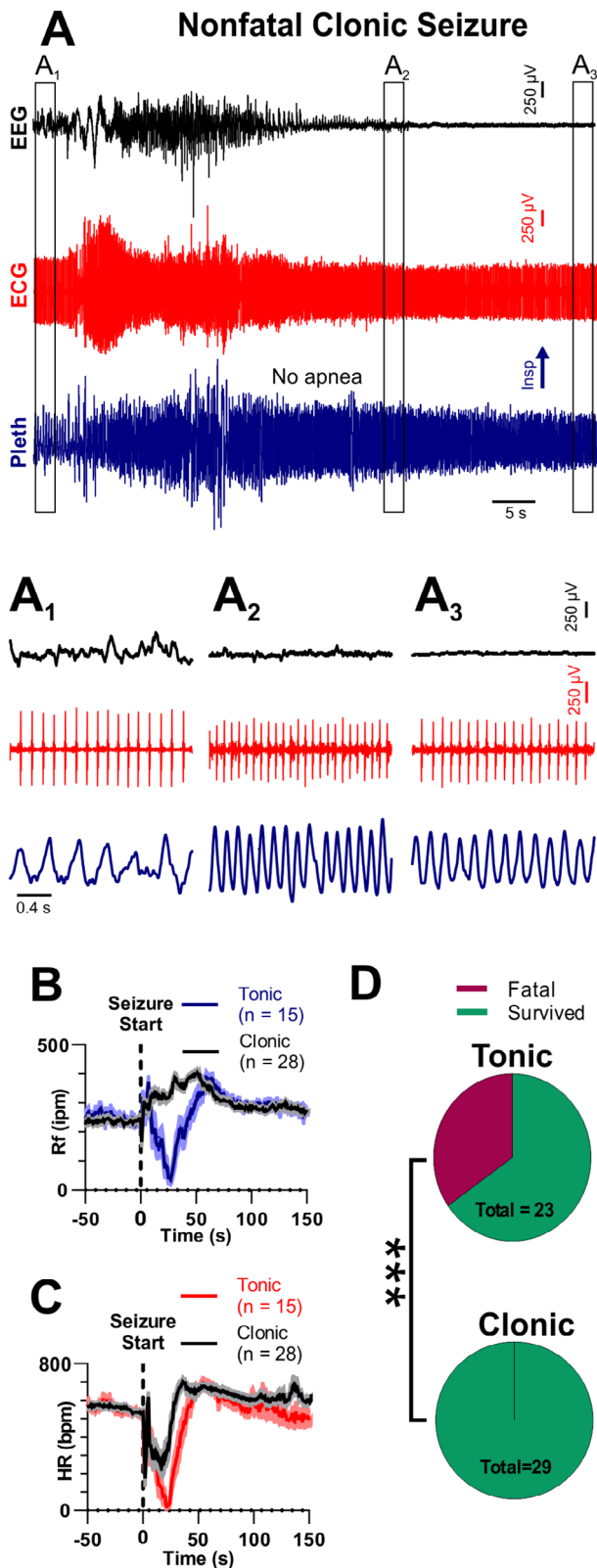


FIGURE 4: Clonic seizures do not produce death or apnea. (A) Recording of electroencephalogram (EEG), electrocardiogram (ECG), and plethysmogram (Pleth) during a clonic seizure in a $W/+^{Emx1-Cre}$ mouse. (A₁–A₃) Expanded views of boxes in A, depicting EEG, ECG, and breathing prior to, during, after the seizure. (B) Respiratory frequency (Figure legend continues on next column.)

pronounced tonic phase, concurrent with apnea and bradycardia (see Fig 3). Following the tonic phase of nonfatal seizures, mice experienced tachycardia and tachypnea, which were different from the bradycardia and apnea observed after fatal tonic seizures. Our analysis likely underestimates heart rate during the tonic phase due to EMG activity obscuring the QRS complexes. However, at the end of the tonic phase, when QRS complexes were distinguishable from the EMG signal, heart rate was clearly reduced. Interestingly, there were no mice that had intermediate responses between complete and permanent respiratory failure, and rapid recovery. The mechanisms for conversion from reversible apnea to terminal apnea are unknown.

Tonic Phase Apnea Is Necessary for Seizure-Induced Death

Clonic seizures were similar to tonic seizures with respect to EEG activity and most behavioral semiology (compare Fig 4A to Fig 3A; Videos S2 and S3). However, instead of a tonic phase denoted by hindlimb extension, clonic seizures had repetitive clonic movements in the upper extremities, visible in the ECG signal as small repetitive bursts of EMG activity (see Fig 4). Instead of apnea, tachypnea occurred during clonic seizures, with the minimum respiratory frequency (Rf) during clonic seizures being detectably higher than for tonic seizures (329 ± 12 and 75 ± 27 inspirations per minute [ipm], respectively; $p < 0.0001$, Mann–Whitney test). Both tonic and clonic seizures presented with bradycardia, which did not differ statistically (70 ± 21 vs 279 ± 60 beats per minute [bpm], respectively; $p = 0.1244$, Mann–Whitney test). Although clonic seizures represented a significant proportion of non-fatal seizures (29/44), no clonic seizure was fatal, and the probability of mortality was higher for tonic compared to clonic seizures.

Tonic Diaphragm Contraction Occurs during Tonic Phase Apnea

In $W/+^{Emx1-Cre}$ mice, ictal apnea is coincident with the tonic phase. Apnea did not occur during clonic seizures. Analysis of the minimum Rf for all nonfatal seizures from $W/+^{Emx1-Cre}$ mice ($n = 43$ seizures) identified 2 distinct groups; those with breathing rates of <100 or >100 ipm,

(Rf) and (C) heart rate (HR) binned every second from 50 seconds before to 150 seconds after seizure initiation for tonic and clonic seizures. (D) Pie charts show tonic seizures are more likely to cause fatality than clonic seizures (***) $p = 0.007$, 2-sided Fisher exact test). For B and C, solid lines represent mean, and shaded regions represent standard error of the mean. bpm = beats per minute; Insp = inspiration; ipm = inspirations per minute.

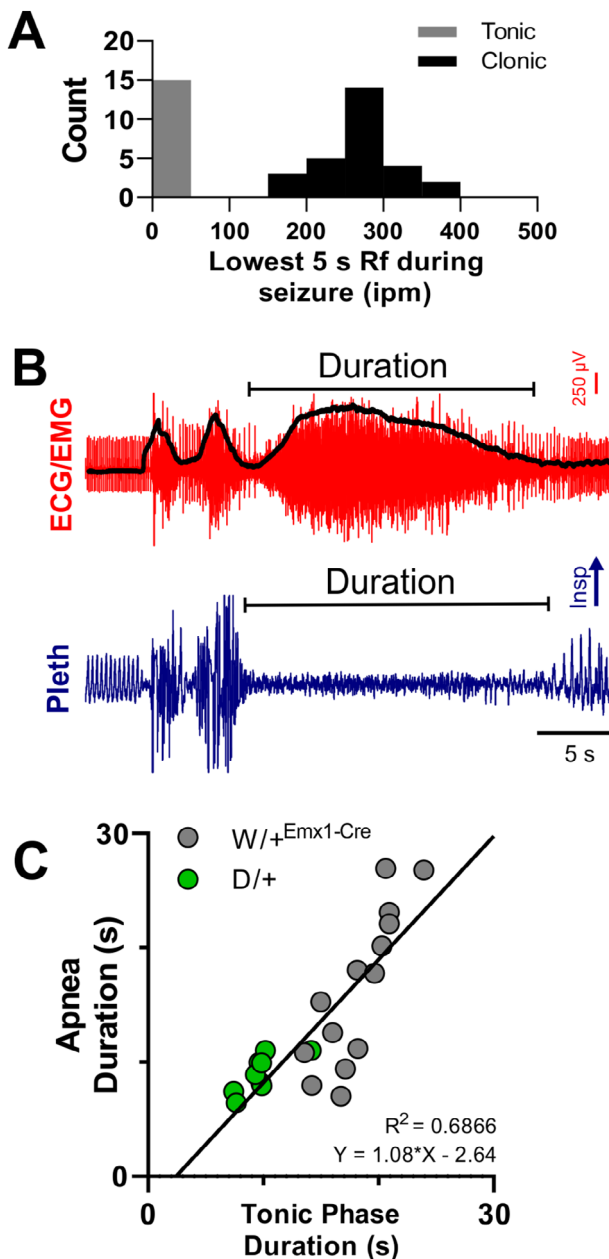


FIGURE 5: The tonic phase is coincident with apnea. (A) Histogram of the nadir of breathing rate for all nonfatal seizures. Breathing rate nadir fell into 2 distinct groups; tonic and clonic, with low (near zero) and high respiratory frequency (Rf), respectively. (B) Depiction of measurement of tonic phase and apnea delay and duration made from all nonfatal tonic seizures. Delay was measured from the seizure onset (ie, the first spike-wave) to apnea or tonic phase start (spike-wave/electroencephalographic [EEG] data not shown in image). The black line superimposed on the electrocardiogram (ECG)/electromyogram (EMG) trace is the same signal after rectification and smoothing. Lower points on the black line were used to determine the start and end of the tonic phase. (C) The duration of apnea is directly proportional to the duration of the tonic phase itself ($R^2 = 0.6886$; $n = 14$ and 9 for $W/+^{Emx1-Cre}$ and $D/+$ mice, respectively). Insp = inspiration; ipm = inspirations per minute; Pleth = plethysmogram.

corresponding to tonic and clonic seizures, respectively (Fig 5A). We hypothesized that tonic phase and apnea share an underlying mechanism, and thus a longer tonic phase should be correlated with longer apnea. We compared duration of the tonic phase and apnea, with data obtained from spontaneous tonic seizures recorded from $D/+$ mice included, as they present with the same behavioral and electrographic signatures, but with shorter apnea than $W/+^{Emx1-Cre}$ mice (9.0 ± 0.5 s and 16.4 ± 1.8 s for $D/+$ and $W/+^{Emx1-Cre}$ mice, respectively; $p = 0.0045$; unpaired t test, $t = 3.182$, $df = 21$). This allowed us to assess the relationship of tonic phase and apnea over a wider range. To estimate tonic phase duration, we rectified and smoothed the ECG/EMG signal and identified the lowest points preceding and following the tonic extension phase (see Fig 5B, black trace on top). Across all apneic seizures of the 2 mouse models, apnea duration was directly proportional to tonic phase duration (see Fig 5C; $R^2 = 0.6866$). It is important to note that wild running occurs immediately prior to the tonic phase. The running produces noise in the plethysmography signal, making it impossible to know whether there is breathing during this short period. For this and other reasons, our determination of the time of onset of breathing cessation is imprecise, and it is possible that apnea starts prior to the tonic phase in some cases.

We hypothesized that tonic muscle contraction prolongs apnea. To test this, we recorded diaphragm muscle activity (diaEMG; as recently described³⁰) during audiogenic seizures in $D/+$ mice. Our diaEMG was reliably synchronized with inspiratory activity recorded by plethysmography (Fig 6A). When we stimulated an audiogenic seizure, we observed a large amount of tonic diaEMG activity during the tonic phase (see Fig 6B). This observation was replicated in a total of 8 audiogenic seizures recorded from 3 $D/+$ mice (see Fig 6C).

Comparison of Fatal and Nonfatal Seizure Dynamics

The ictal duration and amplitude of PGES were compared between clonic, tonic, and fatal seizures from $W/+^{Emx1-Cre}$ mice (Fig 7A). There was no difference in ictal duration among the 3 seizure types (see Fig 7B). PGES, defined as the lowest EEG amplitude within 30 seconds of seizure termination as a percentage of preictal level, was detectably greater for fatal compared to clonic seizures, but was not detectably different between nonfatal tonic and fatal tonic seizures (see Fig 7C).

Duration, peak, and area under the curve (AUC) of EMG tonic activity were analyzed for tonic nonfatal and fatal seizures in $W/+^{Emx1-Cre}$ mice (see Fig 7D). Duration of tonic phase was longer for fatal seizures compared to

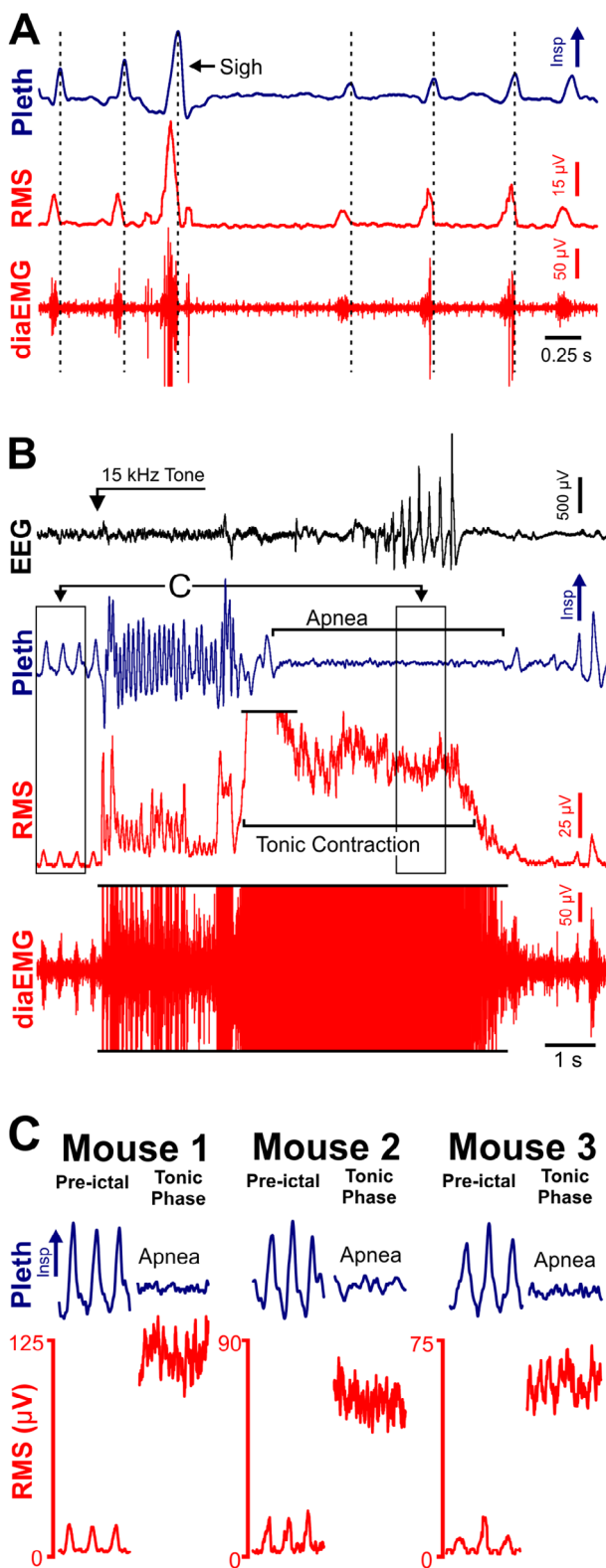


FIGURE 6: Tonic diaphragm activity is coincident with tonic phase apnea. (A) Plethysmogram (Pleth), raw diaphragm electromyogram (diaEMG), and root mean square amplitude of diaEMG (RMS) recorded from a D/+ mouse. (B) Electroencephalogram (EEG), Pleth, diaEMG, and RMS recorded during an audiogenic seizure from the same D/+ mouse as in A. Some of the diaEMG signal is clipped (indicated by black horizontal lines) to better depict lower

nonfatal tonic seizures (see Fig 7E); however, no detectable differences in peak or AUC EMG activity were identified between the seizure types (see Fig 7F, G).

Tonic Phase Apnea in a Patient with SCN8A Epilepsy

An 18-month-old male patient was monitored for seizure characterization in the epilepsy-monitoring unit (EMU) at the University of Virginia Health System. Exome sequencing detected the heterozygous likely pathogenic variant p. Leu257Val (L257V) located in the pore-lining transmembrane segment S5 in domain 1 of Na_v1.6 (Fig 8A). The corresponding leucine residue is conserved in the 3 paralogous human neuronal sodium channels as well as vertebrate and invertebrate homologs (see Fig 8B). Val257 is not present in the gnomAD database of 120,000 control exome sequences lacking early onset neurological disorders. Prediction software rates the L257V substitution as possibly detrimental (PolyPhen-2 score = 0.9) or damaging (Sorting Intolerant from Tolerant score = 0.037), with a high Combined Annotation Dependent Depletion score of 25.4, indicative of deleterious effects.

During EMU monitoring, a 12-minute cluster of 3 electroclinical seizures was recorded, and reviewed by a neurologist (H.P.G.). Each seizure in the cluster had a pronounced tonic phase, during which breathing motions were either absent or markedly diminished. The position of the patient during the first seizure provided clear visibility of the left side of the body, permitting video analysis of breathing movements (see Fig 8C, Video S6). This seizure lasted approximately 80 seconds, of which 60 seconds involved pronounced tonic muscle contraction (see Fig 8D), as indicated by the presence of a large amount of EMG activity in the ECG signal. The patient had tachycardia upon seizure onset followed by bradycardia at the tonic phase onset (see Fig 8D, bottom trace, heart rate).

To assess respiratory frequency and pattern, visualization of respiratory movements was aided by Eulerian video magnification³⁴ as described previously.¹⁸ Dark pixels between the patient's left arm and body were plotted over time to approximate breathing motions (see Fig 8). We were able to reliably analyze breathing with this method during 2 time periods during the seizure when view of the patient was not obstructed by caregivers. During the first period, occurring just after the start of the

amplitude, preictal signal. (C) Left, expanded 1-s traces from preictal and tonic phase time points indicated in B. Note inspiratory motor oscillation occurs during the preictal period and tonic contraction during the tonic phase. In middle and right panels, the same observation was made during seizures in 2 additional D/+ mice. Insp = inspiration.

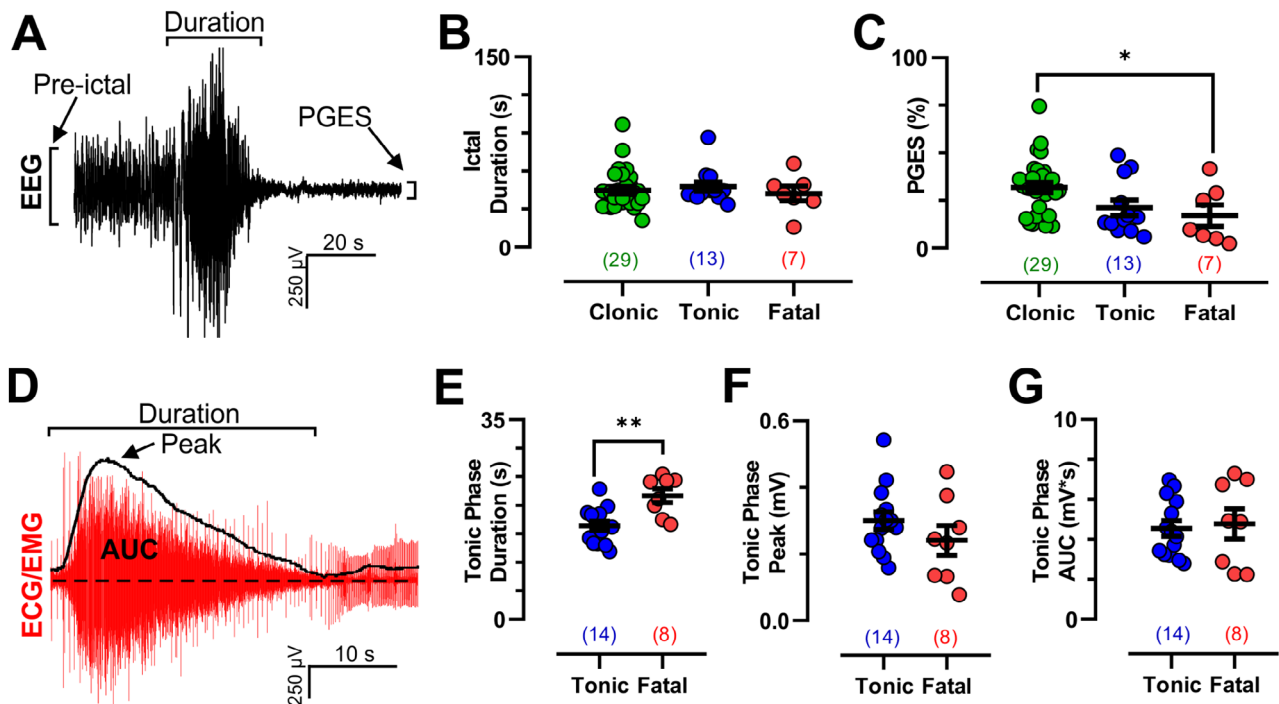


FIGURE 7: Electroencephalographic (EEG) and electromyographic (EMG) dynamics are similar for fatal and nonfatal tonic seizures in $W/+^{Emx1-Cre}$ mice. (A) Depiction of seizure duration, preictal amplitude, and postictal generalized EEG suppression (PGES) amplitude measurements from a peri-ictal EEG trace. (B) Seizure duration was no different across the 3 observed seizure types (Kruskal–Wallis test, $K = 1.166$, $p = 0.5583$). (C) PGES amplitude (as a fraction of preictal amplitude) was smaller for fatal compared to clonic seizures; however, there was no difference between tonic and fatal seizures ($p = 0.0373$ by Dunnett multiple comparison test after significant 1-way analysis of variance, $F_{2,45} = 4.065$, $p = 0.0238$). (D) Depiction of tonic phase duration, peak amplitude, and area under the curve (AUC) measurement from a peri-ictal electrocardiogram (ECG)/EMG trace from a tonic seizure. (E–G) Tonic phase duration was longer for fatal seizures ($p = 0.0013$, $T = 3.729$, $df = 20$, unpaired t test); however, there was no detectable difference in peak ($p = 0.2407$, $T = 1.209$, $df = 20$, unpaired t test) or AUC EMG activity ($p = 0.7583$, $T = 0.3120$, $df = 20$, unpaired t test). * $p < 0.05$, ** $p < 0.01$. Values in parentheses indicate number of seizures analyzed in that dataset.

tonic phase, there were minimal breathing movements, clear tonic muscle contraction, and bradycardia. The second period occurred at the end of the seizure, with minimal breathing motions for >10 seconds prior to clear hyperpnea as breathing resumed (see Fig 8F; Video S6, apnea/hypopnea begins at 0:29 and hyperpnea begins at 1:26). These 2 periods of breathing analysis were different from the eupneic breathing observed several minutes after the seizure when PGES had subsided (see Fig 8G). Fourteen breaths were identified during this 26-second period, resulting in 32 breaths per minute, which is within the normal breathing frequency range for a child of 18 months (29–48 breaths per minute for the 10th and 90th percentiles, respectively).³⁵

Discussion

In the present study, we characterize tonic phase apnea and demonstrate its role in spontaneous and evoked seizure-induced deaths using 3 mouse models of *SCN8A* epilepsy. Recordings of audiogenic seizure-induced deaths from $W/+^{EIIA-Cre}$ mice allowed us to identify breathing

cessation as the primary cause of death. Chronic recording of $W/+^{Emx1-Cre}$ captured spontaneous nonfatal and fatal seizures, which demonstrate that tonic phase apnea is necessary but not sufficient for seizure-induced death. Death only occurred when the animal did not recover from postictal apnea. Recording of diaphragm EMG during audiogenic seizures of D/+ mice allowed us to determine that tonic phase apnea is coincident with tonic respiratory muscle contraction. Finally, data acquired from a patient with a novel *SCN8A* mutation demonstrates that tonic phase apnea can also occur in patients with *SCN8A* epilepsy.

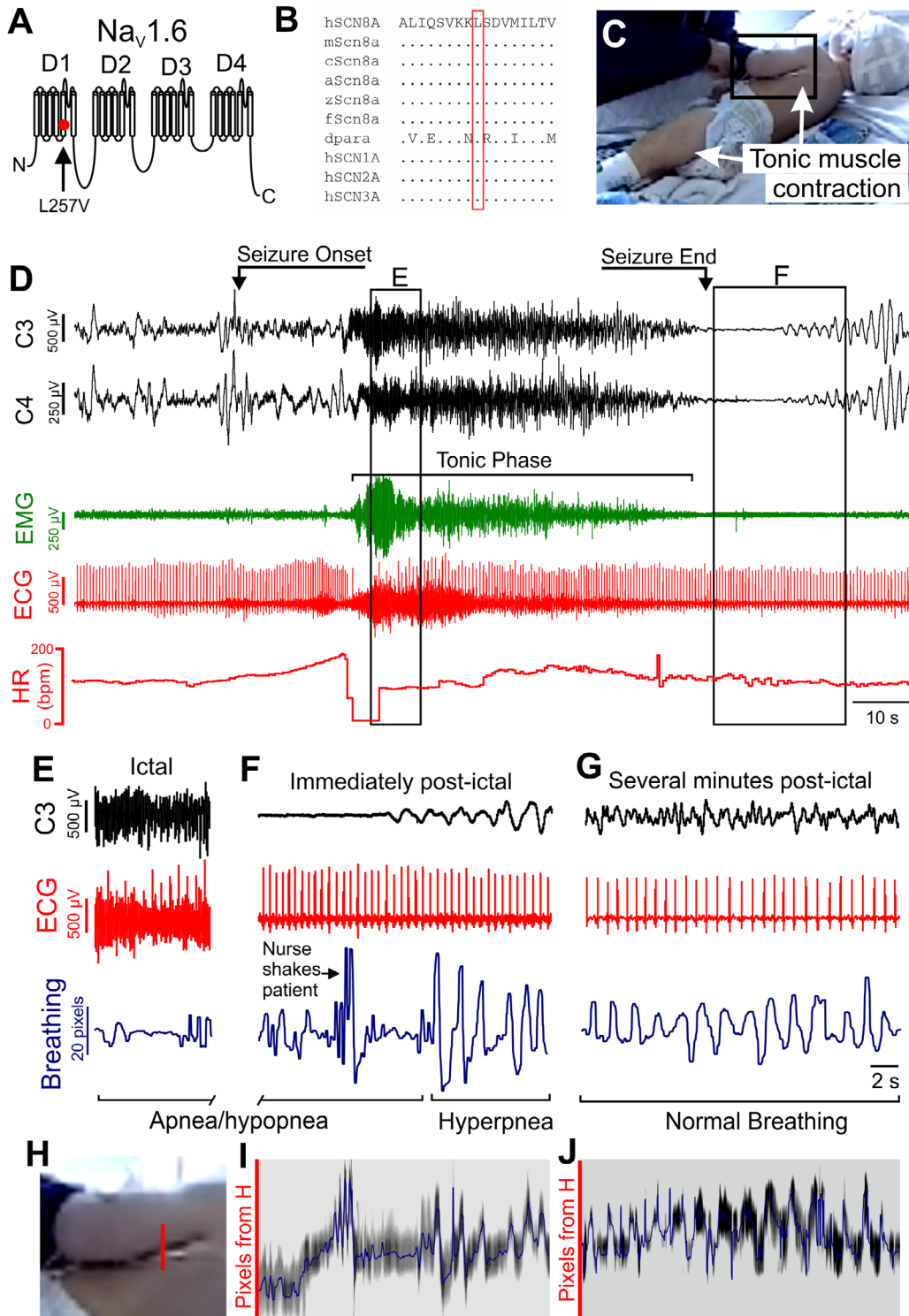
Tonic Phase Apnea

In *Scn8a* mutant and PTZ-treated WT mice, a tonic phase was present in each case in which apnea and death occurred. We further show that tonic phase apnea occurs in a patient with a novel *SCN8A* mutated allele. This is consistent with the limited clinical literature assessing respiratory function during tonic muscle contractions that occur during seizures.³⁶ Tonic phase apnea is likely common to many mouse models of epilepsy. Specifically,

mouse models of Dravet syndrome^{18,37} and *Kcna1*^{-/-38} can demonstrate some apneas coinciding with elevated tonic EMG activity.

Our findings suggest that tonic phase apnea occurs, at least in part, due to tonic contraction of muscles

involved in inspiration (ie, the diaphragm), which prevents the normal rhythmic contraction needed for alveolar ventilation. Although this concept is described in neurology textbooks,³⁹ relevant breathing muscles have not been recorded during tonic seizures in humans or mice.



(Figure legend continues on next page.)

Although tonic diaphragm contraction alone could produce an inspiratory breath hold, effectively an apnea, it should be noted that other respiratory muscles could be affected, including those that could cause airway obstruction, as has been reported for seizures induced with kainic acid in anesthetized rats.⁴⁰

It is important to note that breathing did not resume immediately upon termination of diaphragm contraction for nonfatal seizures. In addition, wild running creates noise in the plethysmography signal immediately before the tonic phase begins, obscuring our ability to determine precise timing of apnea initiation. It is possible, even probable, that apnea is initiated before the tonic phase in some seizures. These periods of apnea outside the tonic phase indicate that mechanisms of central apnea independent of diaphragm contraction must also exist. One such mechanism could be inhibition of brain stem respiratory neural circuitry, as is observed by stimulation of the amygdala.^{41,42} Whatever the mechanism(s) of apnea, our evidence demonstrates a close temporal relationship between the tonic phase and apnea in both the *Scn8a* mutant and PTZ-treated mice; however, the directionality of this relationship is not clear. It is possible that apnea causes tonic contraction or that an unknown third process produces both effects. Because it is currently impossible to selectively suppress the tonic phase, it is not possible to determine the directionality of this relationship.

Apnea Is the Primary Cause of Death for *Scn8a* Mutant Mice

The MORTEMUS study is the most extensive compilation of cardiorespiratory recording during clinical SUDEP events to date. In 9 patients with adequate cardiorespiratory monitoring, terminal apnea occurred before terminal asystole.¹³ This sequence of events was

replicated by both the spontaneous seizure-induced deaths of $W/+^{Emx1-Cre}$ mice and the audiogenic seizure-induced deaths of $W/+^{EIIA-Cre}$ mice; terminal apnea preceded terminal asystole by tens of minutes. Our interpretation is that apnea resulted in anoxia during and after the seizure, which eventually caused cardiac failure. This is supported by our report that WT mice exposed to anoxia succumb to cardiac failure on a similar time scale.¹⁸ In the present study, we were further able to demonstrate a causal relationship between apnea and sudden death by utilizing the audiogenic seizure-induced death of $W/+^{EIIA-Cre}$ mice. These audiogenic seizures were routinely fatal; however, mechanical ventilation greatly increased survival, underscoring the importance of apnea for seizure-induced death in this model.

Tonic Phase Apnea Always Occurs with Seizure-Induced Death in *Scn8a* Mutant Mice and Other Models of SUDEP

All observed instances of seizure-induced death of *Scn8a* mutant mice and PTZ-treated WT mice were preceded by seizures with a tonic phase, denoted by hindlimb extension, and coincident apnea. A number of SUDEP mouse models, including *Scn1a*^{R1407X}, DBA1/2, *Kcna1*^{-/-}, 129/SvTer, and *Cacna1a*^{S218L} mice, experience fatal seizures that are classified as tonic, generally based on the observation of hindlimb extension.^{18–20,38,43–45} However, in many studies of seizure-induced death, seizure-type is not specified, and we have observed a subset of fatal heat-induced seizures of *Scn1a*^{R1407X} mice that were clonic.¹⁸ Thus, the relative predominance of tonic phase apnea during seizure-induced death across other epilepsy models remains to be determined. It is well known that ictal apnea does occur in many patients with non-convulsive seizures,^{8–11} so the same may be true of mouse models.

FIGURE 8: Apnea and bradycardia occur during tonic seizures in a patient with *SCN8A* epilepsy. (A) Diagram of *SCN8A* gene product ($Na_v1.6$) structure, with location of L257V mutation depicted. (B) Evolutionary conservation of residue L257 in multiple species. a = anole; c = chicken; dpara = drosophila "paralytic"; f = fugu; h = human; m = mouse; z = zebrafish. Amino acids are indicated by the single letter code; dots represent identical to the human amino acid. (C) Image of 18-month-old patient during tonic phase of seizure. Note the tonic muscle contraction apparent in the quadriceps and latissimus muscles (arrows). (D) Recording of C3 and C4 electroencephalographic (EEG), electromyographic (EMG), and electrocardiographic (ECG) leads during a convulsive seizure. See Video S6 for full video of the seizure. This seizure has a pronounced tonic phase, as indicated by elevated EMG activity (green signal). bpm = beats per minute; HR = heart rate. (E) At the beginning of tonic phase of the seizure, there was apparent apnea/hypopnea and bradycardia (box E in panel D). Signals shown are EEG, ECG, and breathing assessed with Eulerian video magnification analysis of breathing (blue trace, see below for description of analysis). (F) The same analysis at the end of the seizure showing a period of apnea/hypopnea followed immediately by hyperpnea when breathing recovers (box F in panel D). (G) Breathing analysis several minutes after the seizure, depicting normal, eupneic breathing in addition to normal ictal activity and ECG. (H) Expanded image of region in black box in C depicts column of pixels (red line segment) that were analyzed for breathing movements. (I, J) Grayscale time series of pixels of location depicted in H at the end of apnea (I) and several minutes postictal (J). The blue line corresponds to the midpoint of the band of lowest pixel intensity (binned over 5 pixels). Corresponding signals in F and G were bandpass filtered between 1 and 5 Hz.

Tonic Phase Apnea Is Not Sufficient to Produce Seizure-Induced Death in *Scn8a* Mutant Mice

Although all cases of seizure-induced death in the *Scn8a* mutant mice were preceded by a tonic seizure, there were many occurrences of tonic phase apnea in the $W/+^{Emx1-Cre}$ mice that were not fatal. Thus, tonic phase apnea appears necessary but not sufficient for death to occur in the *Scn8a* and PTZ models used in this study. For the $W/+^{Emx1-Cre}$ mice, recovery from tonic phase apnea was an all or none phenomenon; either breathing rapidly and fully recovered within seconds of tonic phase termination, or it failed completely to recover and death followed. There were no examples of partial or slow recovery. This suggests that a secondary central process responsible for breathing regulation must also fail in order to produce seizure-induced death. It is not unexpected that mechanisms of seizure-induced death and SUDEP will be multifactorial, and different SUDEP mechanisms may exist for different epilepsy etiologies. Other mechanisms that may inhibit recovery from tonic phase apnea include spreading depolarization to the brain stem, impaired homeostatic breathing (ie, central apnea), and cardiovascular dysfunction.^{2,46–48} Cardiac dysfunction appears to vary considerably in different mouse SUDEP models. For example, in $W/+^{Emx1-Cre}$ mice, tonic seizures induce transient asystole that occurs as rapidly as apnea (see Fig 3B, C), whereas in *Scn1a*^{R1407X} mice, fatal seizures are associated with rapid, irreversible apnea, but with much slower development of bradycardia.¹⁸ Thus, future investigation should focus on mechanisms of tonic phase apnea, and uncovering the coexisting factors that impede breathing recovery after the tonic phase.

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Author Contributions

I.C.W., M.K.P., F.A.T., G.B.R., and M.H.M. contributed to the conception and design of the study; E.R.W., P.K.W., J.L.W., P.S.P., P.S., H.P.G., and E.A.B. contributed to the acquisition and analysis of data; I.C.W., M.K.P., F.A.T., G.B.R., and M.H.M. contributed to drafting the text and preparing the figures.

Potential Conflicts of Interest

Nothing to report.

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