INTRODUCTION

Mammalian brains consume a highly disproportionate amount of energy given their relatively small size, thus requiring a system capable of balancing cellular oxygen metabolism and the control of cerebral blood flow to maintain brain oxygen levels (partial oxygen pressure [pO₂]) within the normoxic zone. Impaired cerebral blood flow typically leads to hypoxia, which can have profound and opposing effects on brain tissue oxygenation that may serve as a biomarker for ongoing pathological activity in the brain.
leads to severely hypoxic conditions, insufficient production of adenosine triphosphate, and a failure to meet the high energetic demands of synaptic plasticity and results in behavioral dysfunction. In contrast, when oxygen delivery to the brain becomes too great, hyperoxia typically occurs. Hyperoxia favors the production of reactive oxygen species, which can also lead to neuronal damage and behavioral dysfunction. Seizures have previously been shown to move oxygen levels out of the normoxic range, which raises the possibility that seizure-driven changes in brain pO₂ could influence brain function during and after seizures.

Induced self-terminating seizures in rodent models result in postictal vasoconstriction, hypoperfusion, and severe hypoxia that is responsible for acute behavioral deficits and cognitive dysfunction. Likewise, postictal hypoperfusion is observed in people with self-generated recurrent seizures. In contrast, following exposure to a long-acting convulsant agent, neuronal hyperexcitability correlates with increased cerebral blood flow and hyperoxia in rats. However, the effect of both induced status epilepticus (SE) and post-SE self-generated epileptiform activity on local tissue pO₂ is unknown. Moreover, whereas oscillations in spectral oxygen signals have been shown to be predictive of absence seizures, their role as a biomarker in models of focal epilepsy is also unknown.

In the following study, we determined how local dorsal hippocampal pO₂ responds in both the intrahippocampal kainic acid (KA) and perforant pathway (PP) electrical stimulation models. We measured pO₂ during SE and for several subsequent weeks in relation to self-generated epileptiform activity. We observed SE-induced long-lasting hyperoxia in the dorsal hippocampus in both models. Self-generated spiking activity emerged within days following the termination of SE and was associated with robustly altered oxygen spectra, indicating that oxygen dynamics can be a useful biomarker for active nonconvulsive focal epilepsy. Post-SE epileptiform activity increased in frequency, self-organized into electrographic seizure activity that drove progressively more severe behavioral seizures (endogenous kindling), and was associated with postictal hypoxia. These results provide evidence that epilepsy is not only an electrical disorder, but a vascular one as well.

2 MATERIALS AND METHODS

2.1 Animals and housing conditions

The subjects were 27 male Sprague Dawley rats (Charles River), weighing 200-273 g at the time of surgery, housed individually in transparent plastic cages with food and water available ad libitum. The colony room was maintained on a 12:12-hour light-dark cycle with lights on at 07:00 hours, and all testing took place during the light cycle. Rats were handled and maintained according to the Canadian Council for Animal Care guidelines, and all procedures were approved by the Health Sciences Animal Care Committee of the University of Calgary (AC16-0272).

2.2 Surgery

Bipolar stimulating electrodes were made from 31-gauge twisted Teflon-coated stainless-steel wires (A-M Systems) with gold-plated Amphenol pins attached to the ends. Recording electrodes for both PP-stimulated rats and KA-infused rats were composed of a two-channel twisted stainless-steel wire electrode (Plastics One). Rats were anesthetized using isoflurane (5% induction and 1.5%-2.0% maintenance) mixed with 100% oxygen, and lidocaine (2%) was administered subcutaneously at the incision site as a local anesthetic. The skull was secured in a stereotaxic apparatus (David Kopf Instruments), and burr holes were drilled in the skull at the stereotaxic coordinates. For PP-stimulated rats, two bipolar stimulating electrodes were implanted bilaterally into the perforant paths (anterior/posterior [A/P], −7.3 mm; medial/lateral [M/L], ±5.0 mm; dorsal/ventral [D/V], −6.0 mm) and a unilateral recording electrode into the right dorsal hippocampus (A/P, −3.0 mm; M/L, +2.0 mm; D/V, −3.5 mm). A pO₂ optode (Oxford Optronix) was implanted into the right dorsal hippocampus as well (A/P, −4.5 mm; M/L, +4.5 mm; D/V, −4.0 mm) to closely record the tissue oxygenation changes in the hippocampus. For KA-infused rats, a 22-gauge guide cannula (Plastics One) was implanted into the right ventral hippocampus (A/P, −6.0 mm; M/L, +5.0 mm; D/V, −6.0 mm), a unilateral recording electrode in the right dorsal hippocampus, and a pO₂ optode in the right dorsal hippocampus (A/P, −3.5 mm; M/L, +2.5 mm; D/V, −3.5 mm). Stainless steel screws and dental cement were used to secure the implanted devices to the skull, with one screw serving as a ground reference. All animals were given buprenorphine in a flavored gelatin preparation for 3 days following surgery and allowed at least 5 days rest for recovery prior to experimentation.
2.3 | Electroencephalographic and oxygen recordings

Electroencephalographic (EEG) signals were amplified \( \times 1000-2000 \) (Grass Neurodata Acquisition System) and digitized at 100 Hz with DATAQ software (DATAQ Instruments). The \( \text{pO}_2 \) optode was 250 µm in diameter and produces no detectible damage to brain tissue over time. Tissue oxygenation was monitored using an Oxylite Pro (Oxford Optronix) within a local area of approximately 500-1000 µm\(^3\). Unless otherwise stated, oxygen measurements were sampled at 0.333 Hz.

2.4 | KA infusions

Following a 1-week postsurgical recovery period and prior to SE induction, rats were administered with a sedating dose of urethane (Sigma), which has previously been shown to limit electrical activity during SE to the temporal lobe, thereby preventing extrahippocampal damage and mortality.\(^{22,23}\) Urethane was dissolved in saline at 0.833 g/mL and injected subcutaneously (0.833 g/kg) at 3-4 injection sites along the back of the rat. Following this, rats were placed into the recording cage to record baseline oxygen and local field potentials, which in the context of seizure monitoring will subsequently be referred to as EEG. Sixty minutes following urethane injection, the rat was unplugged, and KA (4.0 mg/mL) was infused unilaterally into the right ventral hippocampus at 0.1 µL/min for 4 minutes through a 1.0-µL Hamilton syringe (Hamilton Robotics) with a microsyringe pump (Harvard Apparatus, model 55-2222). A ventral hippocampal location was chosen, as it has been shown to produce a temporal lobe epilepsy model with a seizure phenotype similar to the standard dorsal model,\(^{24}\) and is located at an appropriate distance such that the dorsal optode is not within the area of KA-induced damage. The microsyringe needle was kept in the ventral hippocampus for 5 additional minutes before removing it and returning the animal to the recording cage. Oxygen measurements and EEG were recorded for a minimum of 90 minutes, or until SE ended, and the rat was removed, and transferred to colony housing for further oxygen measurements.

2.5 | PP stimulation

Following the postsurgical recovery period and prior to SE induction, rats were administered a sedating dose of urethane (Sigma) as described above. Following this, rats were placed into the recording cage to record baseline oxygen and EEG. Sixty minutes following urethane injection, rats were stimulated bilaterally in the PP for 24 hours (2-Hz paired-pulse stimuli 40 milliseconds apart at 20 V and a 10-second 20-Hz train once per minute at 20 V) in accordance with previous work.\(^{20}\) Oxygen was recorded at a sample rate of 0.333 Hz with an on/off cycle of 1.5 hour/2.5 hour to preserve oxygen probe life. Rats were administered additional subcutaneous saline, heating, and eye lubrication throughout the 24-hour stimulation. At the 24-hour mark, the stimulator was turned off and the rats could briefly return to their home cage before immediately being transferred to a new cage for 24/7 video-EEG recording.

2.6 | 24/7 Video-EEG recording

Video-EEG was monitored continuously over the course of 2 to 8 weeks. Electrode cables (Plastics One) were attached to the recording electrodes and a two-channel commutator. EEG was at a sample rate of 250 Hz. Video was recorded concurrently with EEG using day/night infrared cameras. Following the 8 weeks of video-EEG recording, remaining rats were removed, and transferred to colony housing for further oxygen measurements.

2.7 | EEG scoring for epileptiform activity

EEG was scored by two independent reviewers blinded to animal condition. EEG was viewed in 2-minute segments and monitored for large-amplitude self-generated events compared to baseline. When a candidate event was observed, it was compared to the corresponding video to confirm animal behavior and potential sources of artifact. Self-generated epileptiform events were placed into four categories (Figure 1): spiking, low-frequency discharges without behavior, high-frequency focal seizures (FSs), and high-frequency generalized seizures (GSs). Spiking (Figure 1A) was defined as low-frequency (0.5-1.0 Hz), high-amplitude individual spikes that lasted for 1 to 20 minutes. Low-frequency discharges without seizure behavior (Figure 1B) were defined as low-frequency (1-3 Hz) events of brief duration (typically <30 seconds). The amplitude of these events was spindlelike; they would begin at low amplitude, gradually increase in amplitude, and peak before decreasing in amplitude toward the end.

High-frequency FSs (Figure 1C) were high-frequency (∼5-10 Hz), high-amplitude events of moderate duration (typically between 30 and 90 seconds). These seizures are associated with Racine stage 1 and 2 behaviors;\(^{25}\) this includes facial automatisms, twitching, and eye-blinking
stage 1), and chewing and clonic head movements (stage 2). Seizures with these behavioral manifestations were categorized as “focal” because seizure activity is likely discretely localized to a specific brain region or hemisphere. Seizures with unilateral forelimb clonus (stage 3) as the most severe behavioral manifestation were not observed in this study. High-frequency GSs (Figure 1D) were high-frequency (~5-10 Hz), high-amplitude events of even longer duration (typically >60 seconds), and were initially associated with Racine stages 1, 2, and 3 with progression to bilateral forelimb clonus (stage 4) and rearing, loss of balance, and hindlimb clonus (stage 5) within the same electrographic bout. Stage 4 and 5 seizures were categorized as “generalized” because seizure activity engages bilaterally distributed networks.

The first occurrence of a self-generated spike discharge or low-frequency discharge without behavior (Figure 1A,B) was recorded as the first “epileptiform event.” However, subsequent events of this type were not tracked due to their highly frequent occurrence. Each high-frequency FS and GS in each rat was tracked and analyzed for seizure duration.

2.8 | Oxygen dynamics experimentation and analysis

In a separate group of rats (n = 6), KA infusion experiments were performed in the same manner as described above. Following recovery from induced SE, oxygen and EEG were concurrently recorded in the dorsal hippocampus at three time points (48 hours, 1 week, and 2 weeks post-SE). Oxygen measurements were sampled at 1 Hz to perform additional analysis. Python 3.6 was used for oxygen dynamics analysis. To isolate the relative changes and remove the absolute component of the data, a centered sliding average of 60 seconds was subtracted from the data. One minute of data was removed on both ends. This manipulation allowed us to do a power spectral density analysis (Matplotlib.org) and made the data more comparable to techniques only capable of measuring relative changes, which are most commonly used. Data were filtered for low-, medium-, and high-frequency components using a bandpass Butterworth filter (SciPy.org). Data were isolated at the following frequency ranges: 0-0.02 Hz, 0.02-0.04 Hz, 0.04-0.06 Hz, 0.06-0.08 Hz, 0.08-0.10 Hz, and 0.10-0.12 Hz.
2.9 | Statistical analysis

Statistical analysis was performed using Prism (GraphPad) version 6.01. Student t tests (within subjects) were used for experiments with only two groups, or where an a priori hypothesis was tested in experiments with more than two groups. Repeated measures analyses of variance (ANOVAs) were also used when more than two groups were compared. Bonferroni post hoc tests were used where appropriate. Linear regressions were used to analyze the relationship between seizure duration and time post-SE.

3 | RESULTS

3.1 | Hippocampal oxygen levels during KA-induced SE

Local pO2 was measured in the dorsal hippocampus of rats during KA-induced SE (n = 13; Figure 2). Pre-KA baseline hippocampal pO2 was 27.86 ± 2.29 mm Hg. Induction of KA-induced SE significantly increased local hippocampal pO2 (F9, 120 = 4.76, P < .0001). Mean hippocampal pO2 slowly increased 0.8 mm Hg every 10 minutes and was significantly elevated 50-60 minutes after KA infusion (33.09 ± 2.29 mm Hg) compared to baseline. Hippocampal oxygen remained significantly elevated 80-90 minutes after KA infusion (35.41 ± 1.98 mm Hg). Although mean hippocampal oxygen levels remained within the normoxic range for the first 50 minutes, graphs of pO2 for individual rats show highly variable changes in local oxygen levels (Figure 3). Sharp transient increases in pO2 outside of the normoxic range coincided with electrographic seizures during SE. As depicted in Figure 3A, bursts of electrical activity during SE precede brief increases in hippocampal pO2. During KA-induced SE, for this rat, a brief seizure (30-40 seconds in duration) caused an immediate 4- to 10-mm Hg increase in pO2, which peaked at seizure termination before declining. In other rats (Figure 3B,C), these oxygen “spikes” were of larger amplitude (11-29 mm Hg) and occurred more frequently during SE.

3.2 | Hippocampal oxygen levels during perforant path stimulation-induced SE

Hippocampal pO2 was continuously recorded during 24-hour PP stimulation (n = 8; Figure 4). Prestimulation baseline hippocampal mean pO2 was 19.82 ± 2.37 mm Hg, but sharply increased to well above the normoxic range (40.39 ± 2.62 mm Hg) 1.5 hours after stimulation was initiated and remained significantly (F6, 49 = 32.71, P < .0001).

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**FIGURE 2** Mean hippocampal partial oxygen pressure (pO2) in rats receiving intrahippocampal kainic acid infusions. A, Mean hippocampal oxygen trace (n = 13) before and after kainic acid infusion. Pre–status epilepticus (SE) oxygen levels are within the normoxic range (blue horizontal bar), but following kainic acid infusion (red vertical bar) pO2 rises and becomes hyperoxic. B, Quantification of mean hippocampal pO2. Within-subject one-way analysis of variance revealed a significant effect of time (F9, 120 = 4.76, P < .0001). Bonferroni posttests demonstrate oxygen is significantly elevated 50-60 minutes (33.09 ± 2.44 mm Hg), 60-70 minutes (33.95 ± 2.30 mm Hg), 70-80 minutes (34.53 ± 2.20 mm Hg), and 80-90 minutes (35.41 ± 1.98 mm Hg) after induction of SE. Significantly different at *P < .05, **P < .01, ***P < .001.
elevated above baseline for the duration of stimulation. A distinct oscillatory pattern in the pO2 trace was observed in each rat related to the stimulus parameters; every minute, a 10-second train would induce an electrographic seizure (Figure 5A; dark blue) and subsequent increase in pO2. pO2 would rise and then peak shortly after seizure termination and begin to decline, at which point another 10-second train would begin, repeating the pattern (Figure 5A). Although the amplitude of this oscillatory oxygen activity varied at different timepoints during SE and between different rats (Figure 5B,C), the temporal relationship between oxygen changes and seizure activity remained consistent.

3.3 Endogenous kindling of self-generated epileptiform events

Twenty-four hours after the induction of SE, rats from both groups were continuously monitored for 60 days to determine the occurrence and progression of self-generated epileptiform activity within the hippocampus. The first 24-hours following SE termination were not included in the analysis to ensure that the observed epileptiform events were not due to residual SE activity.

In both models, 100% of rats developed and displayed a self-generated epileptiform event in the form of spiking or low-frequency electrographic seizures without seizure behavior. This was observed within 4.88 ± 1.70 days in KA-infused rats and 2.75 ± 0.49 days in PP-stimulated rats (Figure 6A,B). Endogenous kindling progressed in the following manner: spiking and low-frequency electrographic seizures without seizure behavior always preceded the onset of FSs, which in turn always preceded the onset of the first GS in all rats. In KA-infused rats, five of eight had at least one high-frequency FS with a mean latency of 18.80 ± 5.73 days, and three of eight had at least one high-frequency GS with a mean latency of 33.00 ± 10.02 days. In PP-stimulated rats, seven of eight had at least one high-frequency FS with a mean latency of 22.00 ± 8.31 days (Figure 6A,B), and five of eight had at least one high-frequency GS with a mean latency of 33.00 ± 7.17 days. Combining KA-infused and PP-stimulated rats, 63% displayed self-generated epileptiform activity within the second day post-SE, 75% within the first 3 days, and 88% within the first week.
KA-infused rats had 27 FSs or a mean of 3.38 FSs per rat and 17 GSs or a mean of 2.13 GSs per rat, with FSs significantly ($t_{42} = 13.68, P < .0001$) shorter in duration (29.67 ± 2.96 seconds) than GSs (93.71 ± 3.55 seconds; Figure 6C). PP-stimulated rats had 33 FSs or 4.13 FSs per rat and 20 GSs or 2.50 GSs per rat. The PP-stimulated rats also displayed FSs that were significantly ($t_{51} = 4.73, P < .0001$) shorter (54.40 ± 5.54 seconds) in duration than GSs (102.20 ± 9.24 seconds; Figure 6D).

To precisely determine how seizure durations and phenotypes developed over time, individual FSs and GSs from both models were compared against the number of days following the induction of SE (Figure 6E,F). In KA-infused rats, FSs were the first to emerge after the termination of SE and persisted for the duration of their recordings without significantly increasing in duration ($R^2 = 0.08, P = .21$). GSs began later after SE and also did not increase in duration over time ($R^2 = 0.002, P = .88$). In PP-stimulated rats, FSs were also the first to emerge after the termination of SE and did not significantly increase in duration ($R^2 = 0.003, P = .76$). GSs began later after FSs and significantly decreased in duration over time ($R^2 = 0.41, P = .002$).

3.4 | Self-generated seizures and postictal hypoxia

Hippocampal pO₂ was also recorded in rats during self-generated seizures. Oxygen tracings from self-generated GSs in both KA-infused and PP-stimulated rats are seen in Figure 7A,B. As depicted in Figure 7A, a 94-second seizure induced a mild increase in hippocampal pO₂ from 20 to 23 mm Hg, which rapidly declined following seizure termination to between 1 and 2 mm Hg. The hypoxia was sustained for 60 minutes before returning to baseline by minute 80. As depicted in Figure 7B, a 63-second seizure caused an immediate increase in hippocampal pO₂ from 30 to 39 mm Hg. Following seizure termination, oxygen levels dropped and varied between 4 and 12 mm Hg for 40 minutes before slowly rising back into the normoxic range.

3.5 | Interictal oxygen dynamics in rats with KA-induced epilepsy

Hippocampal oxygen measurements were taken every week post-SE to determine whether oxygen dynamics...
were altered as a result of self-generated spiking activity. Following the termination of KA-induced SE, the absolute pO$_2$ in the hippocampus was significantly elevated ($F_{3, 20} = 7.58$, $P = .003$) compared to baseline (Figure 8A,B). Baseline hippocampal pO$_2$ (24.48 ± 1.88 mm Hg) fluctuated within the normoxic range but was significantly increased at 48 hours post-SE (35.53 ± 3.94 mm Hg) and 1 week post-SE (35.20 ± 3.13 mm Hg). There was no difference between baseline hippocampal pO$_2$ and 2 weeks post-SE (31.08 ± 1.68 mm Hg).

Interictal oxygen dynamics were determined by calculating the power of the hippocampal oxygen signal before, during, and after KA-induced SE. Previous research has determined that vascular dynamics in the brain fluctuate at frequencies of <0.1 Hz, thus the oxygen signal was analyzed at various bandwidths between 0 to 0.12 Hz. ANOVA revealed a significant effect of time after SE on the power of the oxygen signal ($F_{5, 30} = 25.96$, $P < .0001$). Specifically, at 1 week post-SE, there was a significant increase in the power of the oxygen signal ($F_{1, 35} = 96.37$, $P < .0001$) compared to baseline (Figure 8C,D). Posttests revealed a significant increase in power at 0.02-0.04 Hz. The changes observed at 1 week post-SE were consistent with data at 48 hours and 2 weeks post-SE (Figure S1).

To further understand changes in oxygen dynamics, the standard deviation (SD) of the filtered oxygen signal was measured across all frequencies. ANOVA revealed a significant alteration in the SD of the oxygen signal ($F_{5, 30} = 17.03$, $P < .0001$). Specifically, at 1 week post-SE, there was an increase in the SD of the hippocampal oxygen signal ($F_{1, 35} = 27.25$, $P < .0001$) compared to baseline (Figure 8E). Posttests revealed a significant increase in oxygen signal SD at 0.02-0.04 Hz. The changes observed at 1 week post-SE were consistent with data at 48 hours and 2 weeks post-SE (Figure S2).

To demonstrate that the observed increases in both oxygen power and oxygen SD were due to self-generated epileptiform activity, we isolated segments of the oxygen signal that corresponded to spiking (Figure 1A) on the EEG. EEG traces were analyzed, and four 15- to 20-minute segments of self-generated spiking from post-KA rats were used in the analysis. ANOVA revealed a significant effect of spiking on the power of the oxygen signal ($F_{1, 41} = 105.7$, $P < .0001$). Posttests demonstrate a significant increase in the power of the oxygen signal between 0.02 and 0.12 Hz (Figure 9A,B). There was also a main effect of spiking on the SD of the oxygen signal ($F_{1, 41} = 21.93$, $P < .0001$), with posttests indicating a significant increase at 0.02-0.06 Hz compared to pre-KA baseline recordings (Figure S3). Next, post-KA EEG traces were analyzed, and four 15- to 20-minute segments, which contained no self-generated epileptiform activity, were isolated and the corresponding oxygen signal was analyzed. ANOVA revealed a significant alteration in the power of the oxygen signal ($F_{1, 41} = 10.03$, $P = .004$), but posttests demonstrated no significant changes in the power of the oxygen signal compared to pre-KA baseline (Figure 9A,B). There was also a significant effect on the SD of the oxygen signal ($F_{1, 41} = 5.43$, $P = .03$).
WOLFF et al.

Using two established models of SE-induced temporal lobe epilepsy, the relationship of local dorsal hippocampal oxygen levels with SE and self-generated epileptiform activity was determined. Temporal lobe SE was associated with increases in local hippocampal pO2 to hyperoxic levels. Following termination of KA-induced SE, the hippocampus remained hyperoxic for at least 2 weeks. During the first 2 weeks following SE termination, when self-generated spiking was present, oxygen dynamics were significantly altered relative to baseline and can serve as a biomarker for the presence of epileptiform activity for focal nonconvulsive seizures. Self-generated epileptiform activity progressively developed into focal and then later generalized behavioral seizures (endogenous kindling). These self-generated seizures led to postictal severe hypoxia similar to previous descriptions following induced seizures in nonepileptic rodents. Thus, SE and self-generated behavioral seizures led to local oxygen levels outside of the normoxic zone but in opposite directions.

The initial observation of SE-induced hyperoxia is consistent with previous work, which has reported hyperoxia in the neocortex in response to soman-induced generalized SE. SE increases the cerebral metabolic rate of oxygen and neuronal glucose metabolism, leading to increases in cerebral blood flow and local pO2 levels. Rodent imaging studies have observed increases in cerebral blood flow in the cortex during pilocarpine-induced generalized SE; however, this study is the first characterization of oxygen levels during focal SE. This sustained hyperoxia may also contribute to brain damage, as the brain is extremely vulnerable to oxidative damage, and KA-induced SE leads to the production of reactive oxygen species. The damage to neurons during this period is sufficient to permanently alter neuronal excitability and cause self-generated recurrent seizures, but the contribution of SE-induced hyperoxia remains to be determined.

The observation that KA-induced SE resulted in hyperoxia in the hippocampus that was sustained for 2 weeks is consistent with previous work that reported hyperperfusion of the rat amygdala between 2 and 14 days following...
pilocarpine-induced SE.\textsuperscript{35} Interictal hyperperfusion has also been measured in clinical populations and correlates with interictal spike rate.\textsuperscript{36} In animal models, bicuculline injections into the cortex lead to measurable increases in cerebral blood flow\textsuperscript{37} and local capillary perfusion,\textsuperscript{38} which align with interictal spikes. SE may drive vasodilation in response to heightened metabolic demand, with post-SE interictal spiking responsible for the maintenance of dilated vessels and subsequent hyperoxia. Metabolic rate may also be chronically reduced following SE. Hypometabolism of glucose and mitochondrial metabolic dysfunction in the hippocampus following SE have been reported.\textsuperscript{39,40} Compromised mitochondria that are consuming less oxygen for aerobic cellular respiration could contribute to hyperoxia. Neurons also become more dependent on glycolysis following KA-induced SE,\textsuperscript{41} and consume less oxygen. The balance between vascular and metabolic factors that contribute to sustained hyperoxia in the hippocampus following SE requires further study.

This study also provides evidence that oxygen dynamics are altered following SE. Previous work demonstrated an increase in the spectral power of the oxygen signal at and above 0.08 Hz in the Genetic Absence Epilepsy Rat From Strasbourg model.\textsuperscript{16} In the KA-induced focal epilepsy model, we observed increases in the power of the spectral oxygen signal at frequencies between 0 and 0.12 Hz, as well as an increase in the SD of the signal at 0.02-0.06 Hz. These changes occur in the absence of clinically obvious (behavioral) seizure activity. Subclinical epileptiform activity, occurring post-SE, likely drives changes in blood flow to the focal region without the sustained vasoconstriction and hypoperfusion typically observed after a clinically apparent seizure.\textsuperscript{13,14} Interictal spikes are known to display blood oxygenation level–dependent (BOLD) changes,\textsuperscript{42,43} and thus are a likely candidate to affect oxygen dynamics. These findings support the use of this technique in clinical populations to locate the seizure onset zone (SOZ), or to differentiate between epileptic and nonepileptic tissue. Human studies using simultaneous intracranial EEG–functional magnetic resonance imaging have shown local BOLD responses to focal interictal discharges, which are colocalized with the SOZ.\textsuperscript{44,45} Furthermore, surgical studies have used optical mapping during nonictal periods to separate epileptogenic cortex from healthy tissue. The successful use of low-frequency hemodynamic oscillations to map the cortex in pediatric patients with epilepsy,\textsuperscript{46} along with similar findings in animal studies, further suggests that these analytical tools can be applied to other noninvasive blood flow imaging modalities.

At longer times after SE, self-generated recurrent focal and generalized behavioral seizures are associated with long-lasting and severe postictal hypoxia (<10 mm Hg). Oxygen levels in this range are associated with cell damage and worse outcomes.\textsuperscript{47,48} Antagonism of the COX-2 enzyme and L-type calcium channels has been shown to effectively prevent seizure-induced hypoxia in model

\textbf{FIGURE 7} Hippocampal partial oxygen pressure (pO$_2$) during self-generating seizure events. A, Hippocampal pO$_2$ before, during, and after a self-generated generalized seizure in a rat that previously received kainic acid–induced status epilepticus. B, Hippocampal pO$_2$ before, during, and after a self-generated generalized seizure in a rat that previously received perforant pathway–induced status epilepticus. Red bars indicate the severely hypoxic range (0-10 mm Hg), whereas blue bars indicate the normoxic range (18-30 mm Hg). Dark blue traces indicate corresponding ictal electroencephalogram.
**FIGURE 8** Altered interictal oxygen dynamics in rats with kainic acid (KA)-induced epilepsy. A, Mean hippocampal partial oxygen pressure (pO2) in rats prior to and after the termination of KA-induced status epilepticus (SE). Horizontal blue bar indicates the normoxic range (18-30 mm Hg). B, Quantification of A indicates significantly elevated hippocampal pO2 at 48 hours and 1 week post-SE. C, Power spectral density analysis of the hippocampal oxygen signal 1 week post-SE plotted on a logarithmic dB scale. D, Quantification of C at binned frequencies. There is an increase in the power of the spectral oxygen signal at 0.02-0.04 Hz and 0.04-0.12 Hz. E, Standard deviation (SD) of the filtered oxygen signal at binned frequencies. There is an increase in the SD of the oxygen signal at 1-week post-SE at the lowest frequency range of 0.02-0.04 Hz compared to pre-KA baseline. Significantly different at *P < .05, **P < .01, ***P < .001, ****P < .0001

**FIGURE 9** Altered oxygen dynamics is due to the presence of self-generated epileptiform activity. A, Power spectral density analysis of the hippocampal oxygen signal from segments where interictal self-generated epileptiform activity was observed (green; post–kainic acid [KA] with spiking) and from post-KA segments that lacked self-generated epileptiform activity (blue; post-KA no spiking) compared to pre-KA baseline (black). B, Quantification of A at binned frequencies. During post-KA periods with interictal spiking, there is an increase in the power of the spectral oxygen signal between 0.02 and 0.12 Hz. There is no change in the power of the oxygen signal at any frequency range during post-KA periods without interictal spiking. This indicates that self-generated interictal activity drives the observed changes in oxygen dynamics following KA infusion. Significantly different at *P < .05, **P < .01, ***P < .001, ****P < .0001
systems, and clinical trials in Canada and the Netherlands are currently underway.

This study demonstrates that oxygen levels become hyperoxic during SE but hypoxic following self-generated seizures and that during self-generated focal nonconvulsive spiking, changes in oxygen dynamics can serve as a biomarker. Thus, epilepsy must also be understood as a vascular disorder, and oxygen levels may contribute to seizure-induced behavior and anatomical anomalies.

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CONFLICT OF INTEREST
None of the authors has any conflict of interest to disclose.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.