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The incidence and significance of periictal apnea in epileptic seizures


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Summary
Objective: The aim of this study was to investigate periictal central apnea as a seizure semiological feature, its localizing value, and possible relationship with sudden unexpected death in epilepsy (SUDEP) pathomechanisms.

Methods: We prospectively studied polygraphic physiological responses, including inductance plethysmography, peripheral capillary oxygen saturation (SpO2), electrocardiography, and video electroencephalography (VEEG) in 473 patients in a multicenter study of SUDEP. Seizures were classified according to the International League Against Epilepsy (ILAE) 2017 seizure classification based on the most prominent clinical signs during VEEG. The putative epileptogenic zone was defined based on clinical history, seizure semiology, neuroimaging, and EEG.

Results: Complete datasets were available in 126 patients in 312 seizures. Ictal central apnea (ICA) occurred exclusively in focal epilepsy (51/109 patients [47%] and 103/312 seizures [36.5%]) (P < .001). ICA was the only clinical manifestation in 16/103 (16.5%) seizures, and preceded EEG seizure onset by 8 ± 4.9 s, in 56/103 (54.3%) seizures. ICA ≥60 s was associated with severe hypoxemia (SpO2 <75%). Focal onset impaired awareness (FOIA) motor onset with automatisms and FOA nonmotor onset semiologies were associated with ICA presence (P < .001), ICA duration (P = .002), and moderate/severe hypoxemia (P = .04). Temporal lobe epilepsy was highly associated with ICA in comparison to extratemporal epilepsy (P = .001) and frontal lobe epilepsy (P = .001). Isolated postictal central apnea was not seen; in 3/103 seizures (3%), ICA persisted into the postictal period.

Significance: ICA is a frequent, self-limiting semiological feature of focal epilepsy, often starting before surface EEG onset, and may be the only clinical manifestation of focal seizures. However, prolonged ICA (≥60 s) is associated with severe hypoxemia and may be a potential SUDEP biomarker. ICA is more frequently seen in temporal than extratemporal seizures, and in typical temporal seizure semiologies. ICA rarely persists after seizure end. ICA agnosia is typical,
1 | INTRODUCTION

Hypoventilation and hypoxemia are typically seen in focal to bilateral tonic–clonic seizures (FBTCS) and generalized tonic–clonic seizures (GTCS),1–3 and severe alteration of breathing patterns after such seizures has been suggested as a possible mechanism of sudden unexpected death in epilepsy (SUDEP).1 However, oxygen desaturations are also found in 30-60% of focal seizures without generalized convulsions.2 Desaturations are more commonly seen with temporal lobe than with extratemporal seizures.2,4 Electrical stimulation of mesial temporal structures consistently elicits central apnea, potentially explaining this observation.5,6 Ictal apnea has also been noted in 44-48% of nongeneralizing focal seizures,2,4,7,8 and has been reported as the main manifestation of focal seizures in a few case reports.9,10 Ictal and postictal central apnea has been suggested as a potential mechanism in some SUDEP11 and near-SUDEP12 cases. However, the role of ictal and postictal central apnea in SUDEP remains to be definitively demonstrated. We set out to examine the phenomenology, localizing value, and impact of ictal and postictal central apnea in patients with intractable epilepsy, in the epilepsy monitoring unit setting.

2 | METHODS

2.1 | Patients and clinical settings

All patients were prospectively consented and recruited participants in the NINDS Center for SUDEP Research’s Autonomic and Imaging Biomarkers of SUDEP project (U01-NS090407). Patients with epilepsy aged ≥16 years who were undergoing video-electroencephalography (VEEG) evaluation were studied in the epilepsy monitoring units (EMUs) of University Hospitals Cleveland Medical Center, University of Iowa, Northwestern University, New York University, Thomas Jefferson University, University of California at Los Angeles, University College London, and Columbia University. Inclusion criteria were patients in whom inductance plethysmography (abdominal and/or thoracic belts) and VEEG recording were carried out during the evaluation in the epilepsy unit. Exclusion criteria were movement or electrical artifacts obscuring plethysmographic signal, or obstructed or unavailable video.

Key points

- Ictal central apnea (ICA) is frequent in focal seizures and may be their first clinical sign
- Patients are apnea agnostic, and hence polygraphic monitoring of respiration during seizures is necessary for diagnosis
- ICA is 10 times more often seen in temporal than extratemporal seizures
- Prolonged ICA (>60 s) is associated with severe hypoxemia

2.2 | Cardiorespiratory monitoring and VEEG monitoring

All patients had prolonged surface VEEG monitoring using the 10-20 International Electrode System. EEG and electrocardiography (ECG) were acquired using the Nihon Kohden (Tokyo, Japan), Micromed (Modigliani Veneto, Italy), and Xltek (Natus, Pleasanton, CA, USA) acquisition platforms. Peripheral capillary oxygen saturation ($\text{SpO}_2$) and heart rate were monitored using pulse oximetry (Nellcor OxiMax N-600x [Convidien, Minneapolis, MN, USA], Masimo Radical-7 [Masimo, Irvine, CA, USA] and SenTec Digital Monitoring System [Therwil, Switzerland]). Chest and abdominal excursions were recorded using inductance plethysmography (Ambu, Ballerup, Denmark] Sleepmate and Perfect Fit 2 [Dymedix, St. Paul, MN, USA]). Oxygen desaturations were classified as mild ($\text{SpO}_2$ of 90-94%), moderate (75-89%) and severe (<75%). We defined central apnea as cessation of breathing movements lasting for ≥10 s in the absence of generalized tonic or clonic movements, since such movements invariably produced movement artifact in breathing channels. Tachycardia and bradycardia were defined as heart rate >100 beats per minute and <60 beats per minute respectively, or a >20% deviation from baseline. Seizures were classified according to the International League Against Epilepsy (ILAE) 2017 seizure classification based on the most prominent clinical signs: focal onset impaired awareness (FOIA) motor onset with automatisms, FOIA nonmotor onset (dialepsis), FOIA motor onset with hyperkinesis, focal onset aware (FOA)
motor onset tonic and/or clonic, focal onset to bilateral tonic–clonic seizures (FBTCS), and FOA nonmotor onset (auras). Cognitive seizures, where the main clinical manifestation was aphasia, were further classified as aphasic seizures. Electrographic seizures were defined as seizures where the sole clinical manifestation was ictal central apnea. Generalized onset non-motor typical seizures were classified as absence seizures, and generalized onset tonic–clonic seizures of primary generalized epilepsy were classified as GTCS (or GTCS, diagnosis based on history and EEG findings). The putative epileptogenic zone was defined based on clinical history, seizure semiology, neuroimaging, and scalp EEG.

2.3 | Statistical analysis

Statistical analysis was performed using the Statistical Package for Social Science (SPSS, version 24; IBM Corp, Armonk, NY, USA). Summary statistics were reported as mean ± standard deviation (SD; median, range). Chi-square test and binary logistic regressions were used to assess the association between dichotomous variable apnea (yes/no), with other variables and combinations. Because the 103 apneic seizures were not normally distributed, nonparametric testing (Kruskal-Wallis test) was used to assess apnea duration with other variables.

3 | RESULTS

A total of 473 patients underwent polygraphic study of seizures. Reliable inductance plethysmography recordings and unobstructed seizure videos for the assessment of breathing responses were available in 312 seizures in 126 patients (77 female). Mean age was 40.09 ± 14.71 years (median 38.5; range 16–77). One hundred nine patients had focal epilepsy and 17 patients had primary generalized epilepsy. Mean epilepsy duration was 17.8 ± 13.5 years (median 17; range 0–52).

3.1 | A) Ictal central apnea (ICA) incidence and duration

Ictal central apnea (ICA) was found in 103/312 (36.5%) seizures in 51/126 (40.5%) patients (29 female). Mean ICA duration was 28 ± 18.8 (median 22; range 10–97) seconds (s). Oxygen saturation data were available for 227/312 seizures overall, and in 79/103 seizures with ICA. In the remaining seizures, data were rendered unreliable because of dislodged sensors and movement. Prolonged ICA (≥60 s) occurred in 8 patients and was associated with severe hypoxemia (SpO2 <75%) in 6. Seizure and epilepsy details of patients with and without ICA are shown in Table 1.

3.2 | Influence of type and duration of epilepsy, age, and gender on apnea

ICA was seen exclusively in focal epilepsy (36.5% of all partial seizures; P < .001); none of the 17 primary generalized epilepsy patients (15 with primary GTCS and 7 with absence seizures), had ICA. None of the 15 GTCS were preceded by central apnea. In the focal epilepsy group, 10/22 FBTCS had preceding ICA; ICA was the sole clinical manifestation in all 10 before the onset of tonic–clonic activity. We found that older age was significantly associated with ICA presence (P < .001), but not with ICA duration or hypoxemia severity (P = .6). There were no gender differences in ICA incidence (P = .2). Although longer duration of epilepsy was not significantly associated with the presence of ICA (P = .3), it was associated with longer ICA duration (P < .001). Examples of presence or absence of ICA are shown in Figures 1 and 2.

3.3 | Epileptogenic and ictal-onset zones and seizure semiology

Temporal lobe epilepsy was highly associated with ICA presence in comparison with extratemporal epilepsy (odds ratio [OR] 10.1, 95% confidence interval [CI] 5.5–18.5; P = .001) and to frontal lobe epilepsy (OR 8.3, 95% CI 4–17.3; P = .001) (Figure 3). Temporal lobe ictal-onset zone was accordingly significantly associated with ICA (P < .001). We then assessed whether the first ictal discharges at or after ICA onset involved one or both hemispheres. The ictal discharge was unilateral in 85/103 apneic seizures (87%), nonlateralizable in 7/103 (7%) and obscured by artifact at ICA onset in 11/103 (11%). ICA was significantly more likely to be associated with unilateral (left or right) ictal discharge at apnea onset compared to nonlateralizable EEG seizure onset (P < .001). Temporal lobe epilepsy (P < .001) and unilateral temporal lobe EEG ictal onset (P = .001) were both significantly associated with longer ICA duration. There was a higher incidence of ICA in FOIA motor onset with automatisms (71.4%) and FOIA nonmotor onset (dialepsis) (55.9%) compared to other seizure types (Figure 3). Both were highly associated with ICA presence (P < .001), as well as ICA duration (P = .002) and severe hypoxemia (P = .04).

3.4 | Awake and sleep states

Mean duration of ICA in seizures during the awake state was 27.87 ± 19.38 s and 28.13 ± 18.43 s during the non-REM (rapid-eye movement) sleep state; the awake/sleep states at seizure onset did not significantly impact either ICA presence or ICA duration (P = .6). No seizures arose during REM sleep.
### 3.5 | Apnea-induced oxygen desaturation of hemoglobin (SpO2)

Ictal hypoxemia was present in 56/79 (70.8%); desaturation was mild in 26/56 (46%) of seizures (mean 92.5 ± 1.2 [93; 90-94]), moderate in 22/56 (39%) (mean 81.5 ± 4.0 [82.5; 75-89]) and severe in 8/56 (14%) (mean 64.7 ± 9.3 [69.5; 46-72]). Mean oxygen desaturation nadir was 87.7 ± 10.3 (92; 46-98). Duration of ICA was significantly negatively correlated with SpO2 nadir ($r = -0.89; P < .001$) (Figure 4). Fifty-three percent of FOIA motor onset with automatism seizures had moderate or severe hypoxemia compared to all other semiologies combined ($P = .04$). Temporal lobe seizures were more likely to have moderate or severe ictal hypoxemia compared to other epileptogenic zones ($P = .03$).

### 3.6 | Ictal apnea characteristics and relationship with EEG onset/clinical onset

In 16/103 (16.5%) of seizures, ICA was the only clinical manifestation during the entirety of the seizure. In 56/103 (54.3%) seizures, ICA occurred before unambiguous EEG seizure onset (mean 8 ± 4.9 [7.7; 1-29] seconds). In 15/103 (14.5%), EEG onset and ICA onset were simultaneous, and in 32/103 (31%) EEG seizure onset preceded ICA (mean 7.7 ± 9.9 [7.7; 1-28] seconds). In 61/103 seizures (68.5%), ICA onset occurred before clinical onset (mean 12.3 ± 9.7 [10; 1-50] seconds). These ICA onsets were simultaneous in 15/103 (16.8%), and in 13 seizures (14.6%), clinical onset preceded ICA onset (mean 13.6 ± 9.6 [12; 1-33] seconds). ICA always occurred in the expiratory phase and all patients were agnostic to their apneas, confirmed by questioning.
FIGURE 1  A left temporal lobe seizure is shown in 3 consecutive 30 s pages, in polygraphic detail. In A, the patient is awake before seizure onset. Cessation of breathing movements was noted 6 s before epileptiform discharges began. In B, during a 50-s apnea period, complete absence of breathing movement is seen, along with oxygen desaturation, with only pulse artifacts identifiable in the plethysmography signal. In C, the patient restarts breathing 15 s before seizure end, when he is interviewed by nurses. The patient was apnea agnostic.
3.7 | Apnea and bradycardia

During ICA periods, heart rate increased in all 103 seizures; it was always seen at or after EEG seizure onset, rather than with ICA onset. Periictal bradycardia was not observed in any seizure.

3.8 | B) Postictal central apnea

Spontaneous restoration of breathing before seizure end was seen in 100/103 (97%). In 3 FOIA motor onset with automatism seizures (3%), in 2/126 patients (2%), apnea persisted into the postictal period for 16-22 s (total

**FIGURE 2** Differences in polygraphy studies are represented in a typical (A), generalized seizure with 3 Hz spike and wave’s discharges where no apnea is observed and (B), focal epilepsy and right temporal lobe seizure where central apnea is clearly seen.
periictal [ictal plus postictal] apnea periods were 46-97 s). None had apnea beginning exclusively in the postictal period.

4 | DISCUSSION

This study suggests that ICA is a semiological feature exclusive to focal epilepsy, that it most commonly starts before unambiguous surface EEG onset, and can be the only clinical manifestation of focal seizures. Temporal lobe epilepsy and frontal lobe epilepsy accounted for most focal epilepsies associated with apnea; temporal lobe epilepsy not only had an 8-fold greater association with apnea than frontal lobe epilepsy, it was also significantly more likely to be associated with longer apneas and more severe hypoxemia. FOIA motor onset with automatisms and FOIA non-motor (dialepsis)
nonmotor onset seizure semiologies, typical of temporal lobe epilepsy, were similarly much more likely to produce ICA, and longer ICA durations. Thus ICA presence and ICA duration may not only help distinguish focal epilepsies, they may also enhance localization to the temporal lobe. Temporal lobe symptomatogenicity for ICA is in concordance with direct electrical cortical stimulation studies in humans that point to highly reproducible apneic responses with low intensity, unilateral (left or right) amygdalar and hippocampal stimulation. Although seizure spread to bilateral temporal structures has been considered necessary to produce ICA, these unilateral stimulation experiments, and the focal, unilateral ictal discharges at the time of ICA in many of our patients, suggest that such spread is not always the case.

It is likely that seizure discharges impair involuntary suprapontine (amygdalohippocampal) breathing control, resulting in ICA. Because ICA occurred after expiration in all our patients, it is likely that inspiration is immediately inhibited by seizure discharge, whereas expiration is mostly passive and allowed to occur to completion. The most likely downstream driver for ICA is seizure-induced inhibition or disruption of brainstem inspiratory neuronal function. Descending amygdala projections to the parabrachial structures, which exert critical roles in phase switching from expiration to inspiration have been described in cats single pulse amygdala stimulation triggers inspiratory onset. Hippocampal activity increases before apnea termination in cats and some hippocampal and amygdalar neurons phase-lock with the respiratory cycle in humans, suggesting that these structures are intimately involved in breathing regulation. Thus the relatively frequent occurrence of ICA in patients with temporal lobe epilepsy is unsurprising. Whether ICA in patients with extratemporal epilepsy reflects spread to amygdalohippocampal structures (anterior cingulate, orbitofrontal, and anterior insular regions are extratemporal sites with amygdalar connections that have also been implicated in cortical breathing control), or whether this implies involvement of symptomatogenic extratemporal breathing control structures is uncertain; shorter duration of apnea in these patients may indicate the involvement of breathing network nodes that are distinct from those involved in temporal lobe seizures.

The apnea agnosia described in stimulation studies appeared to be true of ICA in our patients, and cessation of apnea in partial seizures was not followed by breathing distress, air hunger, or dyspnea despite significant oxygen desaturations. This may explain why ICA has largely gone unrecognized, aside from a few case series and case reports. An additional explanation is that plethysmography is not commonly used in EMUs. Breathing resumption in ICA patients, prior to seizure end, was the rule (97%) with few exceptions (3%). Lack of ICA awareness may be dangerous only in prolonged apnea. We observed that ICA cessation did not appear to be driven by hypoxemia. Although hypercarbia cannot be commented upon here, since carbon dioxide was not measured, ictal central apneas are not reversed by augmentations in ventilatory drive from increasing carbon dioxide in previous human and animal studies, and similar observations have been made in stimulation experiments. ICA durations were highly varied; therefore, changes in seizure discharge intensity in breathing control structures are a more likely explanation.

In our patients, the complete absence of ictal bradycardia with ICA, reported in a minority of seizures in one series, is surprising because bradycardia is a normal response to hypoventilation. Asphyxia in animal models results in heart rate decline and cardiac arrest in approximately 5 min. Consistent with the literature, even in the rare, prolonged ICA epochs (up to 97 s), no bradycardia was observed in our study. Seizure-driven tachycardia is common, and may have overcome any physiological tendency to bradycardia in these patients. Combined perictal apnea and bradycardia, therefore, appears rare, but when it does occur, may comprise a potentially deleterious, high vagal tone phenotype in seizure patients in the SUDEP context; the observed tachycardia apnea combination in this study may reflect a more benign, self-limiting seizure manifestation.

Is ictal apnea a SUDEP biomarker? Most patients had a brief, self-limiting apnea with mild or moderate hypoxemia, suggesting that ICA poses no danger in most cases. Mean and median ICA durations in this cohort were 28 and 22 s, respectively; the sheer frequency of ICA in this cohort, their short durations, and cessation before seizure end, suggest that in most cases ICA is self-limiting and unlikely to be a SUDEP concern. However, prolonged ICA (≥60 s) was associated with severe hypoxemia (SpO2 <75%) (Figure 4), and hence this combination may prove to be a biomarker of SUDEP that deserves prospective study. Indeed, 2 nonfatal ICA durations of 57 and 58 s, with SpO2 of 68% and 62%, respectively, were recorded in a previously reported patient who subsequently died of SUDEP at home. Longer duration of epilepsy is a known SUDEP risk factor. Our observation of a positive correlation between duration of epilepsy and ICA duration is intriguing, and suggests potential plasticity in respiratory circuitry that may render relatively benign, short duration ICA into potentially lethal, longer duration ICA that may predispose to SUDEP. For example, functional neuroplasticity in the nucleus of the tractus solitarius, as evidenced by long-term changes in glutamate release and γ-aminobutyric acid (GABA)ergic neuronal activity, has been shown to occur with epileptogenesis in mice with acquired temporal lobe epilepsy. Postictal apnea appears to be a rare phenomenon. Only 3% of ICA persisted (for 16-22 s) beyond electroclinical
seizure end in this study. None had isolated postictal central apnea beginning exclusively after seizure end. Duration of apnea continuum beyond seizure end was short (16–22 s) with total periictal apnea periods between 46 and 97 s. Postictal apneic bradycardia, frequently reported in the Postictal, agonal phases of MORTEMUS SUDEP cases, and near-SUDEP cases, after a partial seizure\textsuperscript{11} or GTCS,\textsuperscript{28} did not occur in any of our patients. The persistent apnea observed into the postictal period may not have been truly postictal, as epileptiform discharges can persist in deep regions, such as amygdala or hippocampus, and not be seen on scalp EEG. However, persistent apnea could also represent a phenomenon like Todd’s paralysis or postictal aphasia, due to dysfunction or “exhaustion” in the major breathing control sites in the human brainstem.

Some limitations of our study need to be considered. Our conclusions are based on a relatively small number of seizures in the primary generalized epilepsy group. In addition, by considering GTCS or FBTCS onset as ICA end, we may have underestimated ICA duration, since central apnea may conceivably commence in or continue into the tonic–clonic phase. The invariable loss of plethysmographic breathing signal due to movement artifact and the contribution of respiratory muscle spasm to hypoxia render comment on ICA difficult. The exclusion of data that are contaminated by artifact, may also underestimate ICA in extratemporal seizures, since these may be more likely to induce vigorous movements (for example, hypermotor movements) than temporal seizures.

5 CONCLUSION

ICA is a frequent, self-limiting semiological feature in focal epilepsy and can be its only clinical manifestation. However, prolonged ICA and severe hypoxemia together may comprise a potential biomarker of SUDEP. ICA is seen 10 times more frequently at the beginning of temporal than extratemporal seizures, and in typical temporal (FOIA motor onset with automatisms and FOIA nonmotor onset) seizure semiologies. The apnea is frequently seen before unambiguous scalp EEG or clinical seizure onset. ICA rarely persists after seizure end. Without polygraphic monitoring, including pulse oximetry and breathing plethysmography during VEEG, ICA may go unrecognized, since patients are agnostic to the apnea.

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DISCLOSURE OF CONFLICT OF INTEREST

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