The effects of hyperventilation on seizure length and cerebral oxygenation during electroconvulsive therapy

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ABSTRACT

OBJECTIVE: Previous studies have reported that hyperventilation prolongs seizure length. However, there is no clear consensus in clinical guidelines on how to perform hyperventilation during Electroconvulsive Therapy (ECT). The present study aims to investigate the effects of hyperventilation on seizure length and cerebral oxygenation.

METHODS: Forty patients aged 18–65 and classified as ASA I-II, who would have their first ECT course were included in the study. Ethics committee approval was obtained and all patients' consent was taken. The consecutive patients were randomized into two groups as follows: group H (20 patients; target etCO₂: 25–30 mmHg) and group N (20 patients; target etCO₂ 35–40 mmHg). All patients were ventilated with a facial mask for two minutes and later were ventilated by a laryngeal mask (LMA) for one minute. Vital signs, peripheral oxygen saturation (SpO₂), and regional oxygen saturation (rSO₂) were measured before general anesthesia induction, on the 3rd minute of ventilation with an LMA (LMA₃), on the 1st minute postictal (PI₁), on the 5th (PI₅), and 10th (PI₁₀) minutes. The motor seizure duration, Richmond sedation-agitation scale, and the time needed to reach Aldrete Score 9 were also recorded.

RESULTS: There was a significant difference between the groups when they were compared concerning seizure length and recovery time. However, when we compared the rSO₂ values that were measured at different times in group H, the difference between the measurements was statistically significant. When rSO₂ values in group H were compared in doubles, there were significant differences between measurements between the basal and LMA₃, basal and PI₁, and the basal and PI₅. When Richmond agitation scores in both groups are compared, there were no significant differences between the groups.

CONCLUSION: This study found that seizure length was longer, and the recovery time was shorter in group H. There was a contribution of hyperventilation on cerebral oxygenation that was measured on the same person at different times, but cerebral oxygenation was not statistically different from patients that were normoventilated. More studies are required to form a consensus regarding how hyperventilation applies to ECT.

Keywords: Cerebral oxygenation; electroconvulsive therapy; hyperventilation; seizure length.

Electroconvulsive shock is applied to one or both cerebral hemispheres to initiate a seizure. Variables of this shock include stimulus pattern, amplitude and duration. The aim of electroconvulsive therapy (ECT) is to create a therapeutic generalized seizure for 25–60 seconds [1]. Routine use of general anesthesia to induce unconsciousness and to prevent the damage by neuromuscular blockade raised interest in ECT.

Seizure activity is characteristically associated with prolonged sympathetic discharge following an initial parasympathetic discharge. The initial phase is characterized by bradycardia and increased secretion. Sometimes,
severe bradycardia and even temporary asystole are seen. Subsequent hypertension and tachycardia usually last a few minutes. Transient autonomic imbalance can make arrhythmias and T-wave changes in electrocardiography. Cerebral blood flow and intracranial pressure, intragastric and intraocular pressures temporarily increase [2, 3].

Unconsciousness is only required for a short time (1–5 min) from the delivery of a neuromuscular blocker to the start of a successful therapeutic seizure. The seizure itself usually produces anterograde amnesia, somnolence and often confusion for a short time. As a result, only a short-acting induction agent is required. Since many induction agents (barbiturates, etomidate, benzodiazepines and propofol) have anticonvulsive effects, small doses of them should be used [4]. With all these agents, the seizure threshold increases, and seizure duration is shortened. Due to anesthetic drugs that shorten the seizure duration, many studies have been carried out by anesthesiologists to increase seizure duration without increasing the electrical stimulus, and in some studies, results have shown that hyperventilation prolongs seizure duration [1].

Hyperventilation was first performed according to the number of insufflation. However, the number of insufflation is insufficient to show whether hyperventilation has occurred. Since the number of insufflation can be arbitrarily determined, it has been replaced by end-tidal carbon dioxide (etCO₂), which is a more objective parameter in hyperventilation [5]. There is no consensus on the duration of ventilation. According to a current review about hyperventilation in ECT sessions, ventilation duration was determined as 1 minute, some for two minutes, and in others for three minutes in various studies [6–10].

This study aims to investigate the effects of hyperventilation on seizure length and to research the effects of hyperventilation on the brain metabolic rate, which increases during the electrical current by monitoring cerebral oxygenation.

**MATERIALS AND METHODS**

After obtaining the approval of the local ethics committee (2017-12/01), 40 patients were classified as American Society of Anesthesiologists (ASA) I-II, aged between 18–65 years who would have electroconvulsive therapy for their first time under anesthesia, were included in this study. The written consents of the participants were obtained. Patients with uncontrolled cardiovascular, respiratory, metabolic disease, patients that had any intracranial pathology and difficult airway criteria, and patients with a preoperative hemoglobin (hb) value of less than 10 g/dl or above 17 g/dl were excluded from this study. In addition, patients that were taking lithium, sodium valproate and benzodiazepines were included in this study but these patients discontinued lithium for two weeks, sodium valproate for five days and benzodiazepines for two days before the ECT session.

The age, weight, gender, hemoglobin levels and psychiatric diagnoses of the patients were evaluated preoperatively and were recorded. Patients who underwent electroconvulsive therapy for the first time under anesthesia without premedication due to their effect on the convulsion threshold, were taken to the operating table. Before anesthesia induction, systolic blood pressure (SBP), diastolic blood pressure (DBP), mean arterial pressure (MAP), heart rate (HR), peripheral oxygen saturation (SpO₂), and regional oxygen saturation (rSO₂) values were recorded. These values were taken as basal values. Following induction of general anesthesia, etCO₂ values were recorded at the third minute of the ventilation and at the first minute of convulsion of patients with their airway controlled by a laryngeal mask (LMA). Following general anesthesia induction, hyperventilation with an LMA (target etCO₂ 25–30 mmHg) to one group of 20 patients (group H), normoventilation (target etCO₂ 35–40 mmHg) with a laryngeal mask to the other group of 20 patients (group N) were applied with O₂ 100% 6 lt/min for three minutes. The consecutive patients were randomized to group H or group N. The preoxygenation was completed after general anesthesia induction with a facial mask (FM) for two minutes until reaching the maximum muscle relaxation. Subsequently, the preoxygenation laryngeal mask was applied to the patients. The ventilation maneuvers were stopped when the electrical stimulus started. Ventilation maneuvers were applied according to targeted etCO₂ intervals.

All procedures performed under general anesthesia may cause impaired brain perfusion, especially in elderly patients [11]. However, in routine practice, the brain is rarely monitored. Many studies have shown that monitoring cerebral oxygen saturation in the intraoperative period is a convenient and useful method. Near-infrared spectroscopy (NIRS) monitoring is a noninvasive analysis method that demonstrates regional cerebral oxygenation that presents venous values with brain oxyhemoglobin and deoxyhemoglobin tissue concentration [12]. Our device (Somanetics INVOS oximeter) used in this study to measure cerebral oxygenation works with
NIRS technology. The cerebral oximeter was applied unilaterally to the patients’ left temporal region.

The ECT electrodes were applied to frontotemporal regions of the patients bilaterally. The protocol for seizure threshold titration starts with the administration of an initial stimulus of 48 millicoulombs (mC) for women receiving bilateral ECT, and 60 mC for men receiving bilateral ECT. The maximum stimulus charge was 504 mC (THYMATRON™ DGx Somatics, INC).

Before the electrical current and postictal 1st, 5th, 10th minutes, convulsion times were recorded with SBP, DBP, MAP, HR, SpO2 and rSO2 values. The efficacy of the given electrical current was determined by observing convulsive movements in the patient’s extremities. When the electrical stimulus was ended, and the convulsive movements in the extremities were started, a chronometer was started, and it was stopped when the movements were ended. Then, this duration was recorded as the motor seizure duration.

After the patients were awakened, the time required for the modified Aldrete recovery score to be 9 and agitation scores using the Richmond agitation-sedation scale (RASS) were determined and recorded (Table 1). The RASS score was determined at the 10th minute after ECT. In the induction of general anesthesia, propofol (Fresenius Kabi 2%) was used at a dose of 2.5 mg/kg and rocuronium as a muscle relaxant at a dose of 0.6 mg/kg. Sugammadex (Schering-plough 100 mg/ml) was administered to patients at a dose of 4 mg/kg after convulsion at the adequate time and quality.

### Table 1. Modified Aldrete and Richmond Sedation-Agitation Scales

| Modified Aldrete Scale | Richmond Sedation-Agitation Scale |
|------------------------|----------------------------------|
| Consciousness          |                                  |
| Totally awake and oriented (name, place, time) (2). | +4 Combative/Violent/Staff is in danger |
| Response to voice (1).  | +3 Very agitated/Pulls or removes tubes or catheters/Aggressive |
| No response (0).       | +2 Agitated/Non-purposeful movements/Fights ventilator |
| Activity               | +1 Restless/Anxious but movements are not aggressively vigorous |
|                        | 0 Alert and calm |
| Breathing              |                                  |
| Can breathe deeply and cough (2). | -1 Drowsy/Not fully alert but has sustained awakening/ eye-opening to voice >10 sn. |
| Dyspnea, limited breathing or tachypnea (1). | -2 Light sedation/Briefly awakening to voice with eye contact (<10 sn). |
| Apneic or with mechanical ventilation support (0). | -3 Moderate sedation/Movement or eye-opening to voice but no eye contact |
| Circulation            |                                  |
| Blood pressure is ±20% of the pre-anesthetic value (2). | -4 Deep sedation/No response to voice but movement response to physical stimulation |
| Blood pressure is ±20–49% of the pre-anesthetic value (1). |                                  |
| Blood pressure is ±50% of the pre-anesthetic value (0). | -5 Unarousable/No response to voice or physical stimulation |
| Oxygen saturation      |                                  |
| Room-air SpO2 >92% (2). |                                  |
| Need O2 support for SpO2 >90% (1). |                                  |
| Although O2 support, SpO2<90% (0). |                                  |

### Statistical Methods

The data obtained from our study were loaded into the SPSS (ver: 22.0) program. Parametric test assumptions were performed (Kolmogorov-Smirnov) in the evaluation of the data, and variance analysis was used in repetitive measurements in comparing the measurements obtained at different times or situations in the same individuals. The Bonferroni test was used to find the measurement, or measurement groups, which that differ when the decision was made as significant as a result of the analysis. In the comparison of the measurements of
the individuals in both groups at different times, the significance test of the difference between two means and the significance test of the difference between two peers were used. The Chi-square test and correlation analysis with the Pearson test were used to evaluate the data obtained by counting. The data were included in tables with their arithmetic mean, standard deviation, number, and the percentage of individuals. The margin of error was taken as 0.05. When the parametric test assumptions were not fulfilled, the Man-Whitney U test was applied, and the margin of error was taken as 0.05.

**RESULTS**

The difference between the groups concerning gender was insignificant in the individuals included in this study ($\chi^2=1.61; \ p=0.204; \ p>0.05$). No significant difference was found between preoperative hemoglobin levels with the comparison with the groups ($p>0.05$).

The mean age of the patients who underwent hyperventilation was $38.5\pm10.70$, and the mean age of the patients who underwent normoventilation was $37.5\pm8.90$. The difference between the groups concerning age was insignificant ($t=0.32; \ p=0.750; \ p>0.05$).

When patients in both groups were compared for the duration of convulsion and recovery time, the difference between the groups was significant ($p<0.05$) (Fig. 1, 2). When the Richmond Agitation scores of the two groups were compared, the difference was insignificant ($p>0.05$).

Comparing the $\text{rSO}_2$ values measured at different times the group H, the difference between the measurements was significant ($p<0.05$). When $\text{rSO}_2$ values in

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**Table 2.** Mean arterial pressure, heart rate, and peripheral oxygen saturation values at different times between the groups*

|                      | Group H Mean±SD | Group N Mean±SD | p    |
|----------------------|-----------------|-----------------|------|
| Mean arterial pressure (mmHg) |                 |                 |      |
| Basal                | 92.85±11.62     | 90.65±8.42      | 0.49*|
| LMA 3rd minute       | 92.35±11.62     | 89.90±9.51      | 0.47*|
| Postictal 1st minute | 96.25±15.11     | 95.40±9.70      | 0.83*|
| Postictal 5th minute | 93.50±10.95     | 92.70±7.22      | 0.78*|
| Postictal 10th minute| 91.45±10.34     | 91.05±7.22      | 0.88*|
| Heart rate (beat/minute) |                |                 |      |
| Basal                | 80.30±9.81      | 76.15±11.33     | 0.22*|
| LMA 3rd minute       | 84.45±10.10     | 79.30±10.07     | 0.11*|
| Postictal 1st minute | 88.40±13.92     | 88.75±12.15     | 0.93*|
| Postictal 5th minute | 87.60±12.15     | 84.80±10.77     | 0.44*|
| Postictal 10th minute| 86.20±12.94     | 82.15±10.00     | 0.27*|
| Peripheral oxygen saturation ($\text{SpO}_2$) (%) | | | |
| Basal                | 97.50±2.21      | 97.60±1.72      | 0.87*|
| LMA 3rd minute       | 99.70±0.80      | 99.65±0.58      | 0.82*|
| Postictal 1st minute | 99.45±0.99      | 99.55±0.75      | 0.72*|
| Postictal 5th minute | 98.30±2.20      | 98.00±2.24      | 0.67*|
| Postictal 10th minute| 98.15±1.98      | 98.10±1.68      | 0.93*|

SD: Standard deviation; LMA: Laryngeal mask airway; *: p>0.05 insignificant; Mann-Whitney U test was used for the comparison.

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**Table 3.** The difference between the groups concerning etCO$_2$*

| Time of measurement method (n=20) | Ventilation method | Mean±SD | Result t | p   |
|-----------------------------------|--------------------|---------|----------|-----|
| Basal                             | LMA 3rd minute     | 26.75±1.29 | t=14.01  | 0.001* |
| etCO$_2$                          | Hyperventilation   |         |          |     |
| Normoventilation                  | 36.75±2.91         |          |          |     |
| PI 1st minute                     | LMA 3rd minute     | 27.45±1.43 | t=14.94  | 0.001* |
| etCO$_2$                          | Hyperventilation   |         |          |     |
| Normoventilation                  | 37.50±2.64         |          |          |     |

SD: Standard deviation; LMA: Laryngeal mask airway; etCO$_2$: End-tidal carbon dioxide; PI: Postictal; *p<0.05 significant; t= t-test results between groups; n= Number of patients.

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**Figure 1.** The difference between the groups concerning convulsion length.
group H were compared in doubles, there was a significant difference in measurements between the basal and LMA3, basal and PI1, basal and PI5 and the difference between the other measurements was found to be insignificant. Comparison of the rSO2 values measured at different times in group N showed the difference between the measurements to be insignificant (p>0.05) (Table 4).

In group H, there was no significant correlation between hemoglobin values and rSO2 measurements that were measured at different times. The same directional positive correlations were found between hemoglobin-rSO2 in group H, but these correlations were statistically insignificant.

In group N, the same directional correlation coefficients were found between the hgb-basal rSO2 \( (r=0.551) \), the hgb-LMA 3rd rSO2 \( (r=0.654) \), hgb-PI 1st rSO2 \( (r=0.657) \), hgb-PI 5th rSO2 \( (r=0.557) \) and hgb-PI 10th rSO2 \( (r=0.525) \). These correlation coefficients were statistically significant. Accordingly, when the hgb value increased, rSO2 values of the basal, LMA 3rd, PI 1st, PI 5th, and PI 10th minutes increased (Table 5).

Furthermore, a negative correlation was found in both groups between the values of MAP measured at different times and rSO2 values also measured at different times. However, these correlation coefficients were statistically insignificant.

### Table 4. Distribution of the cerebral oxygenation (rSO2) values within the groups

| Technic      | Time of measurement | Mean±SD | Result | p   |
|--------------|---------------------|---------|--------|-----|
| Hyperventilation | Basal rSO2          | 74.25±7.85 |        |     |
|               | LMA 3rd minute rSO2 | 78.60±9.05 |        |     |
|               | PI 1st minute rSO2  | 80.90±9.26 | F=18.16| 0.001*|
|               | PI 5th minute rSO2  | 79.75±9.04 |        |     |
|               | PI 10th minute rSO2 | 78.30±9.17 |        |     |
| Normoventilation | Basal rSO2          | 74.00±7.26 |        |     |
|               | LMA 3rd minute rSO2 | 76.05±9.16 |        |     |
|               | PI 1st minute rSO2  | 76.45±8.61 | F=3.89 | 0.06 |
|               | PI 5th minute rSO2  | 75.65±9.37 |        |     |
|               | PI 10th minute rSO2 | 74.95±9.47 |        |     |

SD: Standard deviation; rSO2: Regional cerebral oxygenation saturation; LMA: Laryngeal mask airway; PI: Postictal; ****p<0.05 significant in Hyperventilation group; F=Friedman analysis of variance test results between groups.

### Table 5. The correlations between cerebral oxygenation (rSO2) and preoperative hemoglobin (hgb) values and in group N

|                   | Pearson correlation coefficient | p   |
|-------------------|--------------------------------|-----|
| Basal rSO2-preop hgb | 0.55                          | 0.01*|
| LMA 3rd minute rSO2-preop hgb | 0.65                          | 0.002*|
| PI 1st minute rSO2-preop hgb | 0.60                          | 0.005*|
| PI 5th minute rSO2-preop hgb | 0.55                          | 0.01*|
| PI 10th minute rSO2-preop hgb | 0.52                          | 0.01*|

rSO2: Regional cerebral oxygenation saturation; preop hgb: Preoperative hemoglobin value; LMA: Laryngeal mask airway; PI: Postictal; *p<0.05 significant.
In only one patient, hemorrhagic secretion was observed due to the LMA application after the ECT course. No further side effects were observed due to LMA and ventilation maneuvers.

**DISCUSSION**

Ventilation has a role in increasing the effectiveness of ECT by prolonging seizure duration [13]. The electrical dose is usually increased to pass the seizure threshold, which may lead to a transient weakening of cognitive functions. Therefore, an optimal dose adjustment is required in ECT.

Many pro-convulsant strategies have been attempted to avoid increasing the electrical dose. These include methylxanthine drugs, such as caffeine and theophylline, opioids (remifentanil), flumazenil, etomidate, anesthetic drugs, such as ketamine, sleep deprivation and hyperventilation alone [1, 14-18]. All of these applications reduce the seizure threshold and the required dose of electrical dose for convulsions [19].

Today, clinical guidelines do not have a clear view of how ventilation should be in ECT. Each patient is routinely pre-oxygenated and is intended to achieve a reliable duration of apnea during the seizure by filling intrapulmonary oxygen reserves. In some sources, hyperventilation with pre-oxygenation is recommended, but the optimal hyperventilation maneuvers are not indicated [8]. The role of hyperventilation for prolonging the seizure duration has been investigated for many years, and these studies were collected in a current review of Gómez-Arnau et al. [6]. We also took this review as a guide for determining our methods in this study.

In our study, we placed a laryngeal mask after induction of general anesthesia to the oropharynx of the individuals and provided 3-minute ventilation before the electrical current was delivered. During this procedure, we performed end-tidal CO₂ monitoring with a capnograph in the breathing circuit that was connected to the LMA. etCO₂ reflects the maximum CO₂ concentration at the end of each exhalation and is close to the arterial CO₂ value in people without a pulmonary disease [7]. Although etCO₂ monitoring in ECT is a useful monitor for controlling hemodynamic parameters, there is no effective monitoring due to mask leaks in patients who are ventilated with a facial mask (FM). Thus, we used an LMA instead of a facial mask for effective etCO₂ monitoring. Because of minimal leaks, better airway control, and better ventilation features, LMA could be a first-line option to manage airway and ventilation in ECT in the future, although nowadays, facial mask ventilation is the most frequently used method. In addition, the most optimal way to determine the level of CO₂ would be to intubate the patient and follow etCO₂, and to look for PaCO₂ in the patient’s arterial blood gas. However, we did not implement this technique because it would be a much more invasive attempt. Due to its invasiveness, intubation is not a common practice in ECT [7].

The facial mask is the most widely used ventilation device for ECT ventilation. However, the facial mask is criticized by many authors since it cannot provide airway security effectively. Instead, a minimally invasive laryngeal mask is recommended [10, 13]. In the study performed in patients with difficult ventilation, the LMA was compared with the FM and patients with an LMA having lower PaCO₂ and longer seizures [10]. In other studies supporting this study, LMA-related positive results were obtained and many authors noted that a more tolerable side effect profile of ECT has been established by reducing the electrical dose that is needed for stimulus in ECT patients ventilated with LMA [13]. However, routine use of the LMA in ECT is not currently recommended in clinical guidelines [8].

We assumed that three minutes of hyperventilation is enough to reach the target values of etCO₂. The long duration of ventilation may affect the depth of anesthesia and may cause excessive hypocapnia because of a long period between general anesthesia induction and electrical stimulation in ECT. Bergsholm et al. [7] made severe hypocapnia, and the target etCO₂ value was 15 mmHg. Many other authors have determined the target etCO₂ as 30 mmHg [5, 8, 9, 13].

Pande et al. [20] evaluated hyperventilation in three stages; etCO₂ values were determined as 20, 25 and 30 mmHg in three different groups, and he did not find a significant difference in seizure durations.

In another study, there was no difference between the 18 mmHg targeted etCO₂ and 25mmHg targeted etCO₂ groups concerning seizure duration [21]. According to the results of these studies, hyperventilation due to the target etCO₂ of less than 30 mmHg may not show any additional benefits in ECT. On the other hand, there is still no consensus on what the target etCO₂ should be.

In addition to the seizure duration, quality parameters, such as ictal amplitude and postictal suppression index, were also used to assess seizure adequacy. In fact,
these parameters are used to evaluate the effects of ECT on the patient’s clinical situation [22]. Although there is evidence of the relationship between hyperventilation and seizure quality, some authors have used these seizure quality parameters [23]. In our study, we did not assess seizure quality with these parameters. Only seizure duration was determined by observing the extremity movements of the patients.

Haeck et al. [13] found a significantly higher ictal intensity in the hyperventilated patient group with an LMA. Di Pauli et al. [21] reported a higher level of postictal suppression index and amplitude in the group of patients hyperventilated with LMA by keeping the target etCO₂ around 28 mmHg. When cerebral blood flow was altered by excessive maneuvers, the efficacy of NIRS to detect sudden changes in cerebral oxygenation was determined by Levy et al. [24].

Aydin et al. [25] randomly divided their 40 patients into two groups in their study referred to as “brain oxygenation during electroconvulsive therapy”. Following general anesthesia induction, one group (group O) was given oxygen at 6 l/min with the nasal cannula and not applied to the other group (group K). There were no statistically significant differences between the groups after 10 minutes of measuring cerebral oxygenation with NIRS. In addition, five patients had desaturation during seizures, and four of them were in group K, with no oxygen. Three of these five patients were accompanied by cerebral desaturation. In addition, postictal etCO₂ values were significantly higher in group K. This is probably due to the lack of adequate filling of the pulmonary reservoirs with preoxygenation and alveolar hypoventilation after the seizure.

Saito et al. [26] investigated the change in cerebral oxygenation using NIRS in electroconvulsive therapy and found a correlation between rSO₂ values and mean arterial pressure that decreased at the time of the seizure and then increased and came back to basal levels. However, in this 43 patient study of Saito et al.’s, patients were premedicated with atropine sulfate (0.01 mg/kg IM). It should be noted that this may affect the cardiac functions of the patient. In our study, no statistically significant correlation was found between mean arterial pressure and cerebral oxygenation in patients without premedication. This is possibly because of the patient’s unstable arterial blood pressure values due to cardiac arrhythmia caused by electrical currents applied to the patient. etCO₂ values were kept between 30 and 35 mmHg in all patients in this study, where we cannot reach satisfactory information about how ventilation is done. Therefore, it is difficult to say whether hyperventilation in ECT contributes to rSO₂ due to Saito et al.’s study.

We found that the effects of ECT on cerebral oxygenation were examined in our literature search, but the studies about the relationship between hyperventilation/hypocapnia and cerebral oxygenation were insufficient. In this sense, our study is important.

Some authors have defined cerebral tissue desaturation as an absolute rSO₂ below 50%, or more than a 20% reduction from the baseline. Other researchers have used a decline of over 20% of the baseline value as the cut-off value for cerebral ischemia-hypoxia. Monitor based cerebral NIRS on continuous-wave technology is a trend monitor. Consequently, clinical decision-making should be based on the change from baseline together with the clinical situation instead of the absolute value observed [27]. In our study, we consider a decline of over 20% of the baseline value as the cut-off value and none of our patients’ rSO₂ values decreased below this level.

The condition limiting our study is that we cannot detect basal etCO₂ in patients because we use a laryngeal mask after general anesthesia induction. Detecting how much the etCO₂ values decrease from their basal values could be an effective parameter in determining the depth of hyperventilation. In addition, data analysis could not be performed according to the diagnosis because the patients were not distributed equally according to their diagnoses. Seizure duration alone should not serve as a marker of the adequacy of treatment. Some research strongly suggests that there are complex relations between stimulus dosing and seizure duration, as when the electrical stimulus is increased further above the seizure threshold, the seizure duration may actually decrease particularly early in the treatment course [28]. This makes the seizure duration a confusing and inadequate outcome measure, despite this, due to its ease, it is still one of the most used. The study did not report the effects on seizure quality, which is a factor to guide clinical decisions during the ECT, together with seizure length and patient’s clinical response. Not monitoring the electrical seizure with EEG was another limiting condition of this study.

In conclusion, our study discussed a controversial issue such as hyperventilation in ECT, which could not reach a definite consensus in the world. Seizure duration is determined as longer in the group of patients, which were hyperventilated. According to the analysis of the
obtained rSO2 data, hyperventilation was found to have a significant contribution to cerebral oxygenation measured at different times in the same individual. However, there was no statistically significant difference from group N.

It was also found that the duration of recovery in group H was shorter than the group N. We contend that because of the rSO2 values is higher in group H than group N, the recovery time was found to be shorter. In hyperventilated patients, cerebral oxygenation is more protected and this situation can be a reason to have a shorter recovery time. The recovery time in hyperventilation has also been found to be shorter in a study of Mayur et al. [29] and in another study of De Arriba-Arnau et al. [30].

To make a definitive judgment on how to perform hyperventilation in ECT and to investigate hyperventilation effect controlling for other potential confounder variables, future controlled studies are needed. The entire ECT course should be considered to understand the longitudinal effects of hyperventilation during the sessions and its effects on clinical outcomes or seizure threshold.

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