Quantitative assessment of somatosensory-evoked potentials after cardiac arrest in rats: Prognostication of functional outcomes*

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Objective: High incidence of poor neurologic sequelae after resuscitation from cardiac arrest underscores the need for objective electrophysiological markers for assessment and prognosis. This study aims to develop a novel marker based on somatosensory evoked potentials (SSEPs). Normal SSEPs involve thalamocortical circuits suggested to play a role in arousal. Due to the vulnerability of these circuits to hypoxic-ischemic insults, we hypothesize that quantitative SSEP markers may indicate future neurologic status.

Design: Laboratory investigation.

Setting: University Medical School and Animal Research Facility.

Subjects: Sixteen adult male Wistar rats.

Interventions: None.

Measurements and Main Results: SSEPs were recorded during baseline, during the first 4 hrs, and at 24, 48, and 72 hrs postasphyxia from animals subjected to asphyxia-induced cardiac arrest for 7 or 9 mins (n = 8/group). Functional evaluation was performed using the Neurologic Deficit Score (NDS). For quantitative analysis, the phase space representation of the SSEPs—a plot of the signal vs. its slope—was used to compute the phase space area bounded by the waveforms recorded after injury and recovery. Phase space areas during the first 85–190 mins postasphyxia were significantly different between rats with good (72 hr NDS ≥50) and poor (72 hr NDS <50) outcomes (p = .02). Phase space area not only had a high outcome prediction accuracy (80–93%, p < .05) during 85–190 mins postasphyxia but also offered 78% sensitivity to good outcomes without compromising specificity (83–100%). A very early peak of SSEPs that precedes the primary somatosensory response was found to have a modest correlation with the 72 hr NDS subscores for thalamic and brainstem function (p = .066) and not with sensory-motor function (p = .30).

Conclusions: Phase space area, a quantitative measure of the entire SSEP morphology, was shown to robustly track neurologic recovery after cardiac arrest. SSEPs are among the most reliable predictors of poor outcome after cardiac arrest; however, phase space area values early after resuscitation can enhance the ability to prognosticate not only poor but also good long-term neurologic outcomes. (Crit Care Med 2010; 38:1709–1717)

Key Words: cardiac arrest; prognosis; functional outcomes; somatosensory evoked potentials; electrophysiology; quantitative neural monitoring

According to the Resuscitation Outcomes Consortium, 294,851 emergency medical service-treated cardiac arrests (CAs) occurred in the United States in 2008 (1). The overall survival after CA remains low, and only 3–7% of the survivors regain normal neurologic status (2). Among survivors, poor neurologic outcomes are the main causes for morbidity and mortality (3–5). In light of these facts, the American Heart Association has recently redefined traditional cardiopulmonary resuscitation as cardiopulmonary-cerebral resuscitation to emphasize the importance of neuroprotection after CA (6).

CA-induced hypoxic ischemic injury initiates a cascade of molecular events that lead to disruption of membrane potential, cellular depolarization, and swelling, ultimately causing secondary neuronal injury (7). Thalamocortical pathways are shown to be especially vulnerable to hypoxic-ischemic insults (8) with a higher degree of neuronal cell death and injury in regions of the cortex and thalamus observed both in animals (9–12) and in humans (13–16). A loss of scalp potentials in patients with thalamic lesions has also been reported (17). Injury to the cortical and subcortical structures has been suggested as the cause of comatose states after ischemia (18). Furthermore, deep-brain thalamic nuclei, such as the intralaminar (19) and central median (20) nuclei, have been found to play vital roles in arousal pathways. These findings, along with the knowledge of the thalamus as a sensory relay center, have led us to hypothesize that the thalamus may play a major role in recovery after CA-induced coma. The normal conduction of somatosensory evoked potentials (SSEPs) requires both an intact cortex and an intact thalamus and provides an avenue to assess the integrity of the somatosensory pathway, the restoration of normal thalamocortical coupling, and the onset of arousal.

While the prognostic value of SSEPs in comatose patients is well established, their clinical use is limited to studying the presence or absence of characteristic short and long latency peaks (N20 and N70 in humans). While the absence of cortical evoked potentials within 24 hrs after cardiopulmonary resuscitation has been shown to result in poor neurologic outcomes (13), the presence of these peaks does not necessarily correlate with
better recovery. In addition to the lack of reliable quantitative indicators for SSEP monitoring, electrophysiological examination of patients is often done 8–24 hrs after CA. Since neural damage due to ischemia begins soon after CA, we hypothesize that studying neural activity as soon as possible after injury may be of value not only in prognostication but also in guiding the treatment regime by studying in real time the effect of therapeutic interventions.

SSEPs are only a few microvolts in amplitude and are often corrupted by biological (from nonspecific neural activity) and electrical noise. Thus, waveform averaging (typically 100 to 1000 sweeps in intraoperative monitoring) is required to enhance signal quality (21). SSEPs are traditionally evaluated using peak-to-peak amplitudes and their latencies relying on sophisticated peak detection techniques. Several time-frequency measures have been used to evaluate evoked responses including but not limited to peak power, peak frequency, and time-to-peak frequency (22, 23). An inherent drawback of all the above methods is that they ignore other signal characteristics, such as the rate of rise and fall, the width of the crests and troughs, and the overall morphology of the waveform, which might contain information about the functionality of the conduction pathway. Prior research suggests that the negative slope of the scalp evoked potential in humans is generated by thalamocortical radiation, whereas the subsequent positive wave reflects the cortex (24). Further, SSEPs after injury and during recovery are complex low-amplitude signals, often with indistinguishable peaks which motivated us to assess the shape of the evoked response.

We studied SSEPs using a dynamical systems perspective in the phase space. In order to obtain the phase space curve (PSC) for an SSEP waveform, we plotted its first time derivative against its magnitude. We propose to use the area bounded by the PSC as a prognostic tool and as a quantitative aid in clinical decision-making. We evaluated the association of both the PSA and the amplitudes of the obligate potentials of the rat somatosensory response with neurologic outcomes. In addition, we used PSA to study the correlation between different parts of the SSEP with the thalamic, brainstem, and cortical function. Finally, we sought to identify the time frame most suited for electrophysiological examination after CA to optimize the use of PSA as a prognostic tool and as a quantitative aid in clinical decision-making.

MATERIALS AND METHODS

Sixteen adult male Wistar rats (361 ± 12 g; Charles River, Wilmington, MA) were randomly divided into two groups and subjected to a 7-min (n = 8) or a 9-min (n = 8) asphyxia. This experimental model of global cerebral ischemic injury after CA has been previously well established and validated (11, 25). The animal protocol used in this study was approved by the Institutional Animal Care and Use Committee at the Johns Hopkins University School of Medicine.

Electrode Implantation

One week before the experiments, the rats were implanted with five epidural screw electrodes (Plastics One, Roanoke, VA) to record bipolar SSEPs from the left and right cerebral hemispheres. Electrodes were placed over the somatosensory cortex near areas that topographically represent the fore- and hindlimbs. A ground electrode was placed over the paraspinal right frontal lobe. Care was taken to ensure that electrodes did not puncture the dura mater so as to not come into contact with the brain. The electrodes, lead wires, and the exposed portion of the skull were covered with dental cement. Finally, the skin was closed with sutures and the animal was allowed to recover for a week.

Experimental Procedures

On the day of the experiment, the rats were intubated and mechanically ventilated using a pressure-controlled ventilator. Ventilation was adjusted to maintain physiologic pH, pO2, and pCO2. The rats were anesthetized using 1.5% isoflurane in a 1:1 N2/O2 gas mixture. The femoral artery and vein distal to the inguinal ligament were cannulated in order to monitor blood pressure and arterial blood gas and to provide medications. Baseline SSEPs (15 mins) were recorded using the previously implanted electrodes, followed by a 5-min anesthesia washout. During the last 2 mins of anesthesia washout, 2 mg/kg of vecuronium, a muscle paralytic, was administered, and the gas mixture was switched to room air. At the end of washout, mechanical ventilation was stopped by clamping the endotracheal tube, and the time to CA was recorded as the time taken for the mean arterial pressure to fall below 10 mm Hg. The total duration of asphyxia for each rat was predetermined to be either 7 or 9 mins. Cardiopulmonary resuscitation was performed with external chest compressions and mechanical ventilation, along with the use of epinephrine. Sodium bicarbonate was used to counter acidosis. The time to return of spontaneous circulation (ROSC) was indicated by a rise in mean arterial pressure to a value greater than 60 mm Hg.

Isoflurane was restarted at 0.5% 45 mins post-ROSC to maintain animal comfort during SSEP recording. All animals were subjected to an identical and rigid protocol, and the isoflurane was maintained constant at 0.5%. If the rats showed any visible signs of discomfort, the anesthetic was turned up to 1 or 1.5% for short durations. Since all the signals analyzed in this study had similar levels of anesthesia in the baseline and recovery period, the effect of the anesthetic is the same across all animals.

SSEP Recording and Stimulation Protocol

SSEPs were recorded using the TDT System3 data acquisition system (Tucker-Davis Technologies, Alachua, FL). The median nerves were stimulated with subdural needle electrodes placed in the distal forelimbs. The needle electrodes were connected to a stimulus generator and direct current stimulation was applied with 200-μsec-long 0.6-mA pulses. Instead of recording at 3 Hz, the typical stimulation frequency used in clinical studies, we selected a lower stimulus rate of 0.5 Hz to reduce animal discomfort, as minimal anesthetic was used during recovery. Further, since our experimental design required near-continuous stimulation for at least 4 hrs postresuscitation, there was no need for stimulation at higher frequencies, the purpose of which is to obtain a high number of sweeps to average over short bursts of time. The electrodes were connected to an amplifier and SSEP data were sampled at a frequency of 6.1 kHz using the TDT system. Baseline SSEPs were recorded for 15 mins followed by a 5-min anesthesia washout. Following CA, SSEPs were recorded continuously for 1 hr and then at 15-min intervals with a 15-min rest period in between each interval for the next 3 hrs. After the last 15-min interval, the rats were returned to their respective cages and allowed to recover naturally with access to food and water. Finally, SSEPs were recorded for 15 mins at 24, 48, and 72 hrs after injury.

Preprocessing of SSEP Waveforms

The forelimbs of the rats were stimulated in an alternating fashion at 0.5 Hz, and the SSEPs were recorded from the corresponding contralateral hemisphere. Sixty-sweep aver-
Neurologic function in rats was evaluated by an independent trained observer using Neurologic Deficit Score (NDS) at 24, 48, and 72 hrs postresuscitation (26, 27). This scale has been developed and previously validated by our group and seeks to assess arousal, brainstem function, motor and sensory inputs, behavior, and seizures (11, 27); it was adapted from several human and animal scales (28–31). The score ranges from 0 to 80, where 80 is the best possible outcome and 0 the worst (details described in the results). The 72-hr NDS score was the primary outcome measure of the experiment.

Quantitative SSEP Assessment: The PSA

The phase space is the set of all possible states of a system with each state occupying a distinct point. In order to extract morphologic information from the SSEP, we plot the slope against its magnitude. We call this transformation the PSC. Whereas the signal itself dictates the shape of the PSC, the area bounded by the curve is indicative of the power (energy) of the signal. In order to quantify the transformed waveform, we compute the area enclosed by the curve, termed the PSA, using numerical integration by first fitting a convex hull to it using the Quickhull algorithm described by Barber et al (32). Analogous to the manner in which a stretched rubber band takes the shape of the object it is placed around, the convex hull, a simple closed polygon chain, captures the extent of a nonempty set of points in a planar sense.

The PSCs of three different SSEPs and their corresponding PSAs are shown in Figure 1. The PSC stretches further out with larger amplitudes and sharper transitions (steeper slopes). The points selected by the convex hull algorithm in order to capture the extent of the curve include the salient signal peaks and points that capture the slopes of transition. The other features of the signal, such as the smaller peaks, are within the PSC. Upon comparing the normalized peak-to-peak amplitudes with the PSA, it was seen that the PSA was able to better discriminate different morphologies. Whereas the ratio of the N10-P15 peak-to-peak amplitudes (N10-P15 pk-pk) of the SSEPs shown is 3.6:1:1.96 (normalized to the second waveform), the ratio of the PSAs is 6.25:1:2.7 (Fig. 1). This indicates that PSA is more sensitive to differences in SSEP shapes as it incorporates instantaneous slope information while indirectly incorporating information about peak amplitudes and interpeak latencies. A steeper high-amplitude evoked potential encloses a greater area in the phase space domain than does a low-amplitude slowly rising/falling response.

Experimental Stages and PSA Calculation

The experimental monitoring for the evaluation of SSEPs was divided into three stages: immediate injury (0–60 mins), early recovery (60–240 mins), and late recovery (day 1–3), with all time points referenced to the onset of asphyxia. Each stage was further subdivided into smaller time intervals to study the temporal evolution of the SSEPs. Successive 60-sweep averages (2-min averages) were calculated, and the PSA was obtained for each averaged waveform. PSA values were normalized with respect to the baseline PSA for each rat.

Statistical Methods

Statistical analyses were performed using MedCalc for Windows, version 11.2.1.0 (MedCalc Software, Mariakerke, Belgium). Parametric group values (arterial blood gas results, times to CA and ROSC) were reported as mean ± sd. Univariate analysis was performed for parametric data with the use of Student’s t test for continuous variables (two-tailed under the assumption of unequal variances). Nonparametric variables (NDS) were reported as median (25th–75th percentile). The nonparametric Mann-Whitney test was used to test if the NDS in one group was statistically greater than the other by replacing actual data points with a set of ordered ranks.

The Kolmogorov-Smirnov test was used to establish normality of the quantitative SSEP marker (PSA values) and the peak-to-peak amplitudes at each experimental phase. Repeated-measures analysis of variance was used to compare the PSA values between the functional groups across time. Further, to com-
RESULTS

Animal Survival and Functional Outcomes

The rats were a posteriori grouped based on neurologic outcomes recorded at 72 hrs after CA. We prespecified the NDS cutoff into two functional outcome groups: G1, NDS 72 hr <50; and G2, NDS 72 hr ≥50. Rats with NDS <50 were immobile, had sluggish responses, and reacted minimally to environmental stimuli. Rats with NDS ≥50 generally retained some mobility and reacted briskly to stimuli. Table 1 shows the NDS breakdown of a few animals with different functional outcomes. The NDS scores for the groups at 24, 48, and 72 hrs postasphyxia are shown in Figure 24. As can be seen from the Kaplan-Meier curves in Figure 2B, 67% of the rats in G1 did not survive 72 hrs postasphyxia. Since SSEPs were recorded in an alternating fashion from both hemispheres, two bipolar SSEPs were analyzed for each rat, and the average PSA of the two signals at every time was used for the analysis. Data from one rat that did not have characteristic baseline SSEP waveforms, possibly due to loosened electrode connections, was excluded from the shape analysis.

Animal Characteristics and Arterial Blood Gas Monitoring

In order to ensure that the longer duration of CA in half the cohort did not skew those rats to worse neurologic recovery, we compared the time to CA, time to ROSC, and arterial blood gas data between the two groups (Table 2) and found no significant differences in values during baseline and after CA.

Table 1. Representative breakdown of Neurologic Deficit Score of three rats with poor, borderline, and good outcomes, respectively

<table>
<thead>
<tr>
<th>NDS 72 &lt;50</th>
<th>NDS 72 ≥50</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n = 6)</td>
<td>(n = 9)</td>
<td></td>
</tr>
<tr>
<td>Weight (g)</td>
<td>360 ± 12</td>
<td>361 ± 13.4</td>
</tr>
<tr>
<td>Time to CA (sec)</td>
<td>181 ± 36</td>
<td>184 ± 34</td>
</tr>
<tr>
<td>Time to ROSC (sec)</td>
<td>30 ± 0</td>
<td>29 ± 4</td>
</tr>
<tr>
<td>Arterial Blood Gas Data</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>7.47 ± 0.04</td>
<td>7.43 ± 0.02</td>
</tr>
<tr>
<td>pCO₂ (mm Hg)</td>
<td>36 ± 7</td>
<td>42 ± 7</td>
</tr>
<tr>
<td>pO₂ (mm Hg)</td>
<td>185 ± 51</td>
<td>195 ± 42</td>
</tr>
<tr>
<td>HCO₃⁻ (mmol/L)</td>
<td>26 ± 4</td>
<td>28 ± 4</td>
</tr>
<tr>
<td>O₂ Sat (%)</td>
<td>100 ± 0</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>Post-CA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>7.42 ± 0.05</td>
<td>7.40 ± 0.08</td>
</tr>
<tr>
<td>pCO₂ (mm Hg)</td>
<td>42 ± 5</td>
<td>44 ± 11</td>
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<td>pO₂ (mm Hg)</td>
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<tr>
<td>HCO₃⁻ (mmol/L)</td>
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<td>27 ± 4</td>
</tr>
<tr>
<td>O₂ Sat (%)</td>
<td>100 ± 0</td>
<td>100 ± 0</td>
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</tbody>
</table>

CA, Cardiac arrest; ROSC, time to return of spontaneous circulation; Post-CA, timeframe within 1 hour after return of spontaneous circulation; Sat, saturation. All measurements were taken at 37°C.
Temporal Evolution of SSEPs and PSCs After Hypoxic-Ischemic Injury

Representative SSEP waveforms at different experimental stages from a rat in each group are depicted in Figure 3. The onset of ischemia is accompanied by the sudden disappearance of evoked activity. During the first 60 mins, the SSEPs in all rats reappear slowly with the negative peak at around 10 msec (N10) appearing before the recovery of the positive peak at 15 msec (P15). Interestingly, a small negative peak at 7 msec (N7) was also observed in every rat. This peak gets submerged as the N10-P15 peaks recover. It can also be seen that the reappearance of SSEPs follows a pattern with sequential return of peaks: N7, N10, and P15 in that order, giving rise to signals of varying morphologies.

The PSCs corresponding to the SSEPs in Figure 3, A and B, are shown in Figure 3, C and D, respectively. The typical PSC of a normal SSEP is a curve that forms a closed loop that incorporates not only the peak amplitudes but also the slopes, the interpeak latency, and other features of the response. The PSC diminishes in size at the onset of asphyxia in both groups but recovers faster and to a greater extent in G2 (Fig. 3A, C), most prominently during the early recovery phase (85–240 mins).

Recovery of PSAs in the Different Outcome Groups

In all rats combined, the PSA increased significantly as time progressed \( (p = .003) \) highlighting the dynamic nature of SSEP recovery during the first 4 hrs postasphyxia. During the first 60 mins, the PSA was on average higher in G2 than G1, but no statistical difference was observed \( (p = .44) \). The SSEPs in both groups reappeared around the same time. Analysis during the next 3 hrs of recovery—the early recovery duration—demonstrated a significant separation of PSAs between two groups, with G2 showing a greater recovery than G1 \( (85–190 \text{ min}, p = .02; 85–250 \text{ min}; p = .04) \) \( (Fig. 4A) \). G2 also had a significantly greater overall trend of recovery over time compared to G1, as estimated by fitting a linear mixed effects model to the data \( (slopes \text{ of } G2 \text{ and } G1 \text{ are } 0.028 \text{ and } 0.010, \text{ respectively}, p = .05) \).

For the sake of completeness and comparison, peaks were manually inspected, and the N10-P15 pk-pk results for both of the outcome groups were compared with each other. The N10-P15 pk-pk results are in close agreement with the results of PSA, but they offer lesser separation and have a higher degree of variability. Further, the N10-P15 pk-pk trends across time are less statistically significant than PSA \( (Fig. 4B) \).

Comparison of the Predictive Abilities of PSA and N10-P15 pk-pk

The receiver operating characteristic curves were plotted for the early recovery stages to study the costs versus benefits of using PSA as a tool for diagnostic decision making. The binary neurologic outcome classification on the basis of PSAs was found to be 80–93% accurate during the period 85–190 mins after asphyxia. The sensitivity, specificity, accuracy, and corresponding optimal threshold \( (\text{Th}_{\text{optimal}}) \) for the normalized PSAs are summarized in Table 3.

The absence of N20 predicts poor outcome in humans with excellent specificity. However, in our study, N10, the rat correlate of the human N20, was present in all rats at 24 hrs except for one that did not survive till then. Thus, our objective was to gauge eventual outcomes in subjects whose outcomes are uncertain, i.e., in which N10 was present. Since a direct comparison of our marker to the presence/absence of N10 could not be evaluated, we compared the prognostic abilities of normalized N10-P15 pk-pk at different time points (Table 3). Amplitudes were found to have a lower accuracy in classifying rats based on 72-hr functional outcomes than PSA. The sensitivity–specificity tradeoff using PSA was found to be superior. Interestingly, the PSA provided us with consistent thresholds for outcome classification across time \( (\text{greater than } 6–9%) \) recovery with respect to baseline during 85–160 mins and >9–14% recovery during 175–
250 mins), whereas the criteria provided by N10-P15 pk-pk fluctuated randomly between 0% and 20% of recovery during 1–4 hrs after CA.

**The N7-Phase Space Subarea: Correlation With NDS Components at 72 hrs**

The N7 peak appeared before the main N10-P15 signal during early stages of recovery (Fig. 5). In order to study whether this early response associates with the underlying functioning of deep-brain regions or the cortex, we assessed the correlation between PSA of the early (5–9-msec) portion, the phase space subarea, of the SSEP and NDS "subscores" of interest. The N7 signal at the end of 60 mins has a fair but not significant correlation with the functional sub scores corresponding to arousal and brainstem function (Spearman's $\rho = 0.49, p = 0.066$), whereas the sensory-motor subscore showed no significant correlation (Spearman's $\rho = 0.49, p = 0.30$). Groupwise analysis showed that in rats with good outcomes this early signal was more prominent during 25–60 mins postasphyxia (Fig. 5).

**DISCUSSION**

Bedside neurophysiological monitoring using SSEPs is not used extensively in intensive care units (ICUs) for post-CA evaluation except for prognostication of negative outcomes and withdrawal of life support from comatose survivors. In a 407-patient study conducted in 2006 across 32 ICUs, it was observed that SSEP monitoring is superior to clinical tests in predicting neurologic outcomes, but this evaluation was conducted only at 24, 48, and 72 hrs after injury (33). Furthermore, the monitoring relied only on studying the characteristic N20 peak in this study, where 87% of the patients had a poor neurologic outcome, defined as death or persistent unconsciousness 1 month after CA (33). Our study in the rodent model is the first of its kind to evaluate neural injury and recovery using SSEPs during the early stages (>4 hrs) after CA using a novel quantitative tool. Early evaluation is critical for titration of treatments because excitotoxic-ischemic cascades begin immediately after injury, and it is during this time that interventions like hypothermia and drugs might have the most benefit.

We have shown that early SSEP assessments using an objective measure of signal morphology, PSA, does not require sophisticated peak detection and can track brain injury and recovery in addition to differentiating between good and poor neurologic outcomes. There was no discernible difference in PSAs of the two outcome groups during the first 60 mins postasphyxia as signals begin to reappear in a similar fashion after ROSC, but we observed significantly higher PSAs in the good outcome group during the next 3 hrs. We also noted that the early assessment of SSEPs using PSA was able to predict neurologic outcomes at 72 hrs postasphyxia. This is an important result that highlights the need for SSEP monitoring during early stages of recovery after CA. The most common argument against early monitoring is due to the fact that impaired cerebral perfusion and reperfusion damage might affect SSEP recordings. However, we believe that...
Table 3. Summary of the receiver operating characteristic analysis during the early recovery duration

<table>
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<tr>
<th></th>
<th>85–90 mins</th>
<th>115–130 mins</th>
<th>145–160 mins</th>
<th>175–190 mins</th>
<th>205–220 mins</th>
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<tr>
<td>PSA</td>
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<td>0.80</td>
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<tr>
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<tr>
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<td>10.4</td>
<td>11.9</td>
<td>21.0</td>
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AUC. Area under the receiver operating characteristic curve; PSA, phase space area. The AUC was used to evaluate the overall accuracy of the test. A p value of <.05 (italicized) indicates that the AUC is significantly different from 0.5, indicating that the quantitative marker has the ability to distinguish between the two outcome groups. Sensitivity was defined as the proportion of rats with good functional outcomes classified correctly using PSA as the classification criterion. ThOptimal gives the value of the marker (as a percentage of baseline) for best separation between the groups, i.e., lowest false-positive and false-negative rates. Receiver operating characteristic parameters derived using N10-P15 peak-to-peak amplitudes are indicated below the values derived using the PSA marker.

Figure 5. The N7 potential and its recovery. The inset at the top left of the figure shows a typical baseline somatosensory evoked potential (SSEP) waveform showing the characteristic peaks. A very early peak appears about 7 msec poststimulus followed by a biphasic signal characterized by the N10 and P15 potentials. The N7 is the first to recover after ischemic injury and eventually gets submerged as the N10-P15 signal recovers after cardiac arrest. The graph summarizes the evolution of the SSEP N7 phase space subarea during the first hour from the onset of asphyxia. The first 15 min postasphyxia are characterized by suppressed SSEPs, after which the N7 signal begins to recover. As can be seen from the plot, the recovery of N7 stabilizes in all rats by the 60-min mark. Phase space analysis of the N7 peak shows that the recovery of N7 is different between the two outcomes during 35–60 mins postasphyxia (35–60 min, p = .037, one-tailed t test under the assumption of unequal variances under the hypothesis that G2 shows greater recovery than G1). NDS, Neurologic Deficit Score.

monitoring early electrical changes that reflect underlying metabolic states may allow for better prognostication and may be useful in subsequent titration of treatment protocol to maximize the chances of neurologic recovery. It should be noted that SSEPs on days 1–3 in our study offered lower prognostic value for differentiating outcomes.

Another study on the same set of 407 patients mentioned earlier (33) used the presence or absence of short (N20)- and long (N70)-latency peaks to show that, while these peaks give information about the likelihood of poor outcomes, they are not precise enough to base treatment decisions solely on their absence and cannot predict good outcomes. Thus, while traditional SSEP monitoring is shown to reliably prognosticate poor outcomes, our quantitative SSEP assessment using PSA shows 78% sensitivity in identifying good outcomes between 85–220 mins with 83% specificity during 85–160 mins and 100% specificity during 175–190 mins postasphyxia. These findings indicate that a holistic approach to studying SSEPs, taking into account the entire morphology of the waveform, in contrast with the binary assessment based on the presence/absence of characteristic peaks, can provide information about good outcomes with high sensitivity without compromising specificity.

An earlier study evaluated SSEPs for reliable outcome prediction as early as 4 hrs after CA, with subsequent recordings at 12, 24, and 48 hrs after cardiopulmonary resuscitation, and recommended waiting for at least 24 hrs to make a reliable prognosis based on SSEP (34). However, a recent study in dogs revealed that there is a critical window of time immediately after CA during which hypothermia has maximum therapeutic potential (35). The sudden disappearance of the evoked response in all rats was analogous to the isoelectric phase observed in the electroencephalograph immediately after an ischemic insult (25). This can be attributed to the depletion of the energy stores of the brain tissue due to lack of oxygen, which leads to an immediate cessation of electrical activity. It is within this window of time that secondary neuronal injury results from the damaging molecular cascades that are initiated by the lack of oxygen. Therapeutic interventions seek to alter the pathophysiological sequence of events to reduce neuronal injury. Therefore, it is essential to evaluate neural activity as soon as possible after injury and to continue monitoring it during and after the application of hypothermia or other pharmacologic treatments.

Reported mechanistic explanations of coma suggest the involvement of structures like the cortex and the thalamus (18). Injury to the thalamocortical pathway and loss of normal rhythms are speculated to result in coma and the characteristic electroencephalographic bursting patterns through the disinhibition of intrinsic “pacemaker” rhythms (36, 37).
Under anesthesia-induced burst suppression, spontaneous cortical activity is absent during periods of electrical silence, whereas thalamic potentials persist (38). During bursts, coherent potentials are observed in the thalamus and cortex with the thalamus driving the cortex. These observations indicate that burst suppression coma after CA involves abnormal coupling of the thalamic and cortical potentials and that arousal is dependent upon the restoration of the bidirectional thalamocortical coupling. A possible thalamic component of the auditory evoked potential that preceded the cortical response was reported in rats (17). A negative far-field potential that begins around 16 msec preceding N20 in humans is known to be preserved after thalamic lesions and is suggested to originate in the brainstem. The roles of the thalamus and brainstem in arousal and consciousness make them interesting targets to study in order to gain insight into the mechanism of recovery from ischemic coma. We have shown that there is an association of the very early peak (N7) with the brainstem and arousal components of the NDS in contrast to sensory motor assessment. This finding hints at a noninvasive means to evaluate deep-brain regions critical to arousal and consciousness from comatose states with robust implications for clinical translation. Further, these observations set the stage for future studies on the neuroelectrophysiology of the brainstem, thalamus, and cortex using micro-electrode arrays in order to ascertain the origins of the SSEP peaks using spike times.

The absence of histopathological data is a potential limitation of this study. Whereas we have previously demonstrated that histopathological markers for ischemic cell death correlate with NDS (26, 28, 39, 40), we also acknowledge that postmortem histologic markers in rats have been a poor indicator of clinical significance in human trials and have been less predictive than early behavioral assessments (41–43). As such, we put more emphasis on functional outcome and real-time electrophysiologic measures of recovery (PSA) as a means to clinical translation.

The main barrier to clinical translation of these findings is that, in clinical practice, electrophysiological testing is generally delayed for hours to days after resuscitation. As a basic question, the early evolution of SSEP after CA and its prognostic utility remain largely unknown. Our study assesses the early recovery of SSEP recovery using a novel marker that can be automated to provide a real-time monitor that does not require expert interpretation. This study lays the foundation for a more involved study design that may include therapeutic strategies like hypothermia.

A challenge to the implementation of quantitative SSEP monitoring in the ICU is the lack of a baseline signal in CA patients. A possible solution to this problem is to create standardized cortical SSEP waveforms with typical morphologies that can be used for comparison. Another major challenge is the lack of expertise in interpreting SSEPs in the ICUs. Advanced neural monitoring systems with novel quantitative tools such as PSA can offer health personnel the option to select windows of interest to study different components of SSEPs, simultaneously providing a better means to track injury, recovery, and the effect of neuroprotective interventions in real time and a clinically translational means of evaluating the integrity of deep-brain regions.

**CONCLUSION**

The area enclosed by the SSEP in the phase space—a space of all possible configurations of magnitudes and slopes in a signal—was introduced as a quantitative descriptor for SSEPs. We demonstrated that this measure was able to objectively track recovery of evoked responses and differentiate between good and poor neurolologic outcomes with a good predictive ability. Furthermore, we identified a critical window of interest (1–4 hrs postasphyxia) during which the recovery of SSEPs correlates with neurologic outcome, emphasizing the potential benefits of SSEP monitoring in the ICU soon after CA. Finally, we identified a subcomponent of SSEP that many be associated with deep-brain structures.

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