HEART RATE ANALYSIS DIFFERENTIATES DIALEPTIC COMPLEX PARTIAL TEMPORAL LOBE SEIZURES FROM AURAS AND NON-EPILEPTIC SEIZURES

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ABSTRACT - The distinction of non-epileptic from epileptic events is difficult even for experienced neurologists. We retrospectively evaluated 59 dialeptic events from 27 patients admitted for video EEG monitoring to check whether heart rate (HR) analysis could help in differentiating dialeptic complex partial temporal lobe seizures (TLS) from dialeptic simple partial TLS, and non-epileptic dialeptic events. Baseline HR was increased in the simple partial TLS in comparison to complex partial TLS and non-epileptic groups (p<0.05). HR increase accompanied each individual dialeptic complex partial TLS (100% of the events, p<0.05) but HR returned to baseline in the post-ictal phase. Ictal HR was not altered in the non-epileptic or simple partial TLS groups. Our findings suggest that ictal centrally mediated tachycardia is characteristic of dialeptic TLS (both tachycardia and bradycardia have been reported during TLS). This finding may be used as a criterion to distinguish dialeptic complex partial TLS from simple partial and non-epileptic dialeptic events.

KEY WORDS: heart rate, dialeptic seizures, temporal lobe epilepsy, epileptic auras.

A análise da frequência cardíaca diferencia crises dialépticas parciais complexas de auras e crises não epilépticas

RESUMO - A distinção entre eventos não epilépticos de epilépticos é difícil mesmo para neurologistas experientes. Analisamos 59 eventos dialéticos de 27 pacientes internados para monitorização por vídeo-EEG para checar se a análise da frequência cardíaca (FC) poderia auxiliar na diferenciação de crises dialépticas parciais complexas de crises dialépticas parciais simples e eventos dialépticos não epilépticos. A frequência cardíaca basal estava aumentada nos pacientes com crises parciais simples em comparação com o período basal dos grupos parcial complexa e não epiléptico (p<0,05). Houve aumento da frequência cardíaca em cada crise dialéptica parcial complexa (100% dos eventos, p<0,05), mas a FC retornou aos níveis basais na fase pós-ictal. A FC ictal não foi alterada nos grupos de crises não epiléticas e nos pacientes com crises parciais simples. Nosso achado sugere que a taquicardia ictal com mediação central é característica de crises parciais complexas dialépticas (tanto taquicardia quanto bradicardia têm sido relatadas durante crises temporais parciais complexas). Tal achado poderá ser utilizado como critério para diferenciar crises dialépticas parciais complexas de crises dialépticas parciais simples e eventos dialépticos não epilépticos.

PALAVRAS-CHAVE: frequência cardíaca, crises dialépticas, epilepsy do lobo temporal, auras epilépticas.

Autonomic nervous system changes are common in epileptic and non-epileptic seizures and may cause sudden death in epileptic patients¹⁵. These autonomic changes are not fully understood, but may be the result of increased motor activity, emotional distress or modulation of central autonomic circuitry.

Several studies have demonstrated that heart rate (HR) commonly increases during seizures¹⁷, but bradycardia²⁶ and even cardiac asystole³ can occasionally occur during temporal lobe seizures. Opherk et al.⁷ demonstrated that ictal HR analysis could aid in differentiating non-convulsive epileptic seizures from non-epileptic events. They proposed that in “quiet spells”, if ictal HR was increased by more than 30% of baseline, there was a 97% chance that the spell was epileptic. However, no adequate control for movement was performed and HR was evaluated only during the first 10 seconds of the ictal period and postictally only in the first 1-2 minutes.

In this study, we evaluated the HR changes in epileptic and non-epileptic events with minimal amount of motor activity and staring to establish whether

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patterns of HR progression could differentiate them. For this purpose, we compared temporal lobe seizures with particular clinical manifestations, i.e. alteration of consciousness, staring, and loss of or minimal motor activity (therefore fulfilling the criteria of dialeptic seizures using the classification of ictal semiology proposed by Lüders) with similar non-epileptic and partial temporal lobe seizure (TLS) counterparts. Part of this study has been reported in abstract form elsewhere.

**METHOD**

Patients – We performed a retrospective analysis of 59 dialeptic events defined as seizures characterized by alteration of consciousness, staring, and minimal motor activity from 27 patients admitted for video EEG monitoring (11 men, 16 women; maximum of 3 events/patient to avoid selection bias) at the Saint Louis University Epilepsy Unit. This study was approved by the Institutional Review Board from Saint Louis University.

Each dialeptic event - epileptic or non-epileptic - was classified according to the presence of ictal scalp EEG changes in: 1) "Dialeptic complex partial TLS": well-formed, high amplitude, ictal rhythmic theta and delta frequencies (N=28 events); 2) Non-epileptic dialeptic seizures: spells in patients with a final clinical diagnosis of non-epileptic seizures, without background changes on scalp EEG (N=20 events); 3) Dialeptic aura (simple partial TLS): dialeptic events in patients with known temporal lobe epilepsy, not associated with background changes on scalp EEG (N=11 events).

Data acquisition – We analyzed the HR, as well as behavioral changes and EEG patterns during, and 1 h prior to and 1 h after each event. HR was measured by counting the QRS complex in 10 s epochs throughout 3 different periods: baseline (one 10 s epoch every 1 min for 10 min starting 1 h prior to each seizure), ictal (one 10 s epoch every 1 min until 60 min), and post-ictal (one 10 s epoch every 30 s in the first minute, every 1 min during 10 min (from +1 to +10 min), and thereafter every 10 min until 60 min post event (from +10 to +60 min). The onset of the electrographic TLS events was established by the presence of scalp EEG changes and was considered as the initial initial period to start ictal HR measurements in the dialeptic complex partial TLS. The onset of a non-epileptic dialeptic event was defined by the observation of a behavioral change and patient’s subsequent report or family’s assessment (in each event the patient or relative pushed the button to indicate the onset of the event). Patients with non-epileptic events were exhaustively investigated, being admitted for continuous inpatient monitoring during several days in at least one occasion. Most of them had more than one hospital admission for inpatient monitoring. The aura (partial TLS) onset was defined by a behavioral change during monitoring, which was not considered by the patient to be his/her usual seizure event. None of these events progressed to full-blown complex TLS with scalp EEG changes.

Statistical analysis – We used ANOVA for repeated measures and Bonferroni’s test to evaluate HR changes during the event progression (baseline versus ictal versus post-ictal periods) for each individual seizure, within each group and between the different groups. Mean values from the first 30 min, last 10, 5 and 1 min of baseline monitoring and the first 10 and last 30 min of the post ictal period were also compared with the baseline HR values to evaluate the progression of HR changes over time. Differences were considered significant if p<0.05.

**RESULTS**

Mean age was 34.9±2 years. There was no significant age difference (p>0.05) in the 3 groups: 31.2±2.9 (complex partial TLS), 40.7±2.4 (non-epileptic) and 34.2±2.8 (simple partial TLS). In addition, there was no difference in gender distribution, history of smoking or obesity between the 3 groups (p>0.05). No patient had history of coronary artery disease or cardiac arrhythmia. All complex partial TLS events had duration of at least 30 s, and most of them had duration of at least 1 minute. Most of the simple partial TLS (auras) had similar duration, except for few events lasting less than a minute.

Figure shows that the baseline HR was similar (p>0.05) in the non-epileptic (75±1.4 beats/min) and complex partial TLS groups (73±2.5 beats/min) but was increased in the simple partial TLS group: 86±2.6 beats/min (p<0.05). However, as can be seen in Table, motor activity was not increased during the baseline period in the simple partial TLS. Therefore, this increase in baseline HR observed in the simple partial TLS group was not due to increased activity.

HR increase occurred in each individual complex partial TLS, and as a group HR was increased to 109±3.2 beats/min, p<0.05 (Figure), with return to the baseline HR in the post ictal phase: 82±2.4 beats/min (p>0.05). Bradycardia or unchanged HR was not observed in any complex partial TLS. Sixty-one % of the complex partial TLS events started on the right and 39% on the left, with 100% contralateral spreading. In the non-epileptic and simple partial TLS groups, the HR did not increase in the ictal and post-ictal phases (p>0.05): in the simple partial TLS 91±2.6 beats/min (ictal) and 88±2.3 beats/min (post-ictal) and in the non-epileptic events, 78±1.9 beats/min (ictal) and 73±2.5 beats/min (post-ictal).

Despite the increased baseline HR in simple partial TLS, the HR in the last 10, 5 and 1 min of the baseline simple partial TLS period was not significantly different from the HR values in the first 10, 5
and 1 min of the baseline: 86±2.6 beats/min (60 min mean baseline) versus 90±3.7, 91±3.8 and 89±4 beats/min (last 10, 5 and 1 min).

Similarly, the HR in the last 10, 5 and 1 min of the baseline simple partial TLS period was not significantly different from the HR values in the first 10, 5 and 1 min of the baseline, in patients with complex partial TLS or non-epileptic seizures: 73±2.5 beats/min (60 min mean baseline) versus 76±2.8, 77±2.9 and 78±3 beats/min, respectively for the last 10 min, last 5 min and last 1 min of the baseline period (complex partial TLS) and 75±1.4 beats/min (mean 60 min baseline) versus 75±1.8, 76±1.8 and 76±1.9 beats/min, respectively for the last 10 min, last 5 min and last 1 min of the baseline period (non-epileptic seizures).

**DISCUSSION**

HR changes during seizures are common and may increase the risk of sudden death. Sinus tachycardia is by far the most commonly reported autonomic change during seizures. However, bradycardia and many other types of arrhythmias have been described during focal or generalized epileptic events. Its mechanisms are not entirely understood, but a recent study has demonstrated altered baroreflex function, decreased total autonomic variability and relative increase in baseline sympathetic tone, which may further explain the propensity to cardiac arrhythmias and sudden death in temporal lobe epilepsy patients. A recent study from India also reported autonomic dysfunction in 56.3% of the epileptic patients. The existence of unilateral parasympathetic cardiomotor representation in the left hemisphere has been reported but recently questioned. Although one study has correlated the amount of dissemination of the epileptiform activity throughout the cortex with the tachycardia magnitude, simple partial seizures can also be associated with HR changes. Experimental data have also proved that selective stimulation of areas that belong to the central autonomic network, such as the insula, can trigger major autonomic changes. Other important factors which can determine autonomic changes include motor activity and emotional distress.

Our study revealed that dialeptic complex partial TLS (which is not associated with significant motor changes) is characteristically accompanied by tachycardia (100% of the events). Since increased motor activity could not be the culprit for the tachycardia, it is likely that it was due to the activation of components of the central autonomic network. Our interpretation is further supported by our observation that tachycardia started with focal EEG changes and was not necessarily related to spreading of the epileptiform activity. Alternatively, one can argue that the tachycardia was emotionally driven, due to self-perception of the seizure. Although we cannot completely rule out this hypothesis, in most of our patients fear was not reported after the event.

Our study also reveals that dialeptic complex partial TLS can be differentiated from dialeptic simple partial TLS (auras) and dialeptic non-epileptic seizures by HR analysis, since no dialeptic non-epileptic seizure or dialeptic partial TLS was associated with tachycardia, whereas 100% of the complex partial dialeptic TLS caused tachycardia. These findings are also in agreement with Opherk et al., who demonstrated that ictal HR analysis could aid in differentiating non-convulsive epileptic seizures from non-epileptic events, since in “quiet spells”, if ictal HR was

**Table. Motor activity during the baseline period in patients with non-epileptic dialeptic seizures, simple partial dialeptic temporal lobe seizures (TLS) and complex partial dialeptic TLS. Values are expressed as % of the total patients.**

<table>
<thead>
<tr>
<th>Activity</th>
<th>Non-epileptic patients (%)</th>
<th>Simple partial TLS (%)</th>
<th>Complex partial TLS (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any sleep</td>
<td>17.7</td>
<td>20</td>
<td>29</td>
</tr>
<tr>
<td>Standing</td>
<td>82.3</td>
<td>50</td>
<td>8.7</td>
</tr>
<tr>
<td>Seating</td>
<td>44.4</td>
<td>70</td>
<td>50</td>
</tr>
<tr>
<td>Walking</td>
<td>76.9</td>
<td>40</td>
<td>8.3</td>
</tr>
</tbody>
</table>
increased by more than 30% of baseline, there was a 97% chance that the spell was epileptic.

A baseline state of enhanced HR was common prior to the dialeptic partial TLS (auras) and was not due to increased motor activity during the baseline period (Table). This finding could be the result of the self-perception (body awareness) of an imminent complex partial TLS, which was not fully developed subsequently, since ictal tachycardia was present in each individual (all) dialeptic complex partial TLS.

In summary, our findings indicate that in TLS associated with a dialeptic state, tachycardia is invariably present. These findings raise the possibility that even without EEG monitoring, one can distinguish complex partial TLS from simple partial TLS and nonepileptic events by looking at conventional HR monitoring.

REFERENCES