ICTAL ECG CHANGES IN TEMPORAL LOBE EPILEPSY

L.M. LI*, J. ROCHE**, J.W.A.S. SANDER***

SUMMARY - Changes in cardiac rhythm may occur during epileptic seizures and this has been suggested as a possible mechanism for sudden unexpected death amongst patients with chronic epilepsy (SUDEP). We have studied ECG changes during 61 complex partial seizures of temporal lobe origin in 20 patients. Tachycardia was observed in 24/61 (39%) and bradycardia in 3/61 (5%). The mean and median tachycardia rate was 139 and 140 beats/min (range 120-180). The longest R-R interval observed was 9 seconds. No difference was found in regard to the lateralisation of seizures and cardiac arrhythmia. One of the patients with bradycardia was fitted with a demand cardiac pacemaker, which appeared to decrease the number of his falls. In conclusion, ictal cardiac changes which may be seen in temporal lobe epilepsy (TLE) are sinus tachycardia and occasionally sinus bradycardia. Patients presenting vague complains suggestive of either TLE or cardiac dysrhythmia, simultaneous monitoring with EEG/ECG is required, and if the episodes are frequent, video-EEG should be considered. Further studies on this subject are warranted as this may shed some light on possible mechanisms for SUDEP.

KEY WORDS: epilepsy, cardiac arrhythmia, sudden death.

Epilepsy is a common neurological condition with a prevalence of 5-10 cases per 1000 people. Chronic epilepsy has been associated with an increased risk of sudden unexpected death.
(SUDEP) when compared to the general population. An annual mortality rate of 1 SUDEP for 200-370 people with chronic uncontrolled epilepsy has been suggested. The mechanism of SUDEP is not clear, and autonomic changes affecting the cardiac rhythm, deleterious action of antiepileptic drugs, and the release of endogenous opiates affecting the respiratory centre have been advanced as possible mechanisms. Autonomic changes may occur as manifestations of epileptic seizures. In daily clinical practice, however, these changes are not commonly recognised or documented, especially with regard to cardiac rhythms. With the advent of simultaneous EEG and ECG recording, different types of ictal cardiac dysrhythmia have been reported, and this has given some insight about a possible mechanism for SUDEP. Simple or complex partial seizures show no concomitant EEG changes in approximately 20-40% of cases, and this may potentially give a misleading interpretation of simultaneous EEG/ECG recording if the correct timing of clinical onset is unclear.

The aim of this study was to ascertain ictal ECG changes in a selected group of patients with chronic temporal lobe epilepsy (TLE) recorded during pre surgical assessment at the video EEG monitoring Unit of the National Hospital for Neurology and Neurosurgery.

PATIENTS AND METHODS

Twenty patients (11 females, 9 males) were randomly selected from a group of 100 consecutive patients with refractory TLE referred for pre surgical assessment. None of them had a history of cardiac dysfunction, and cardiac examination and routine ECG were normal in all cases. The median age was 28 years (range 16-47).

Simultaneous EEG and ECG were recorded digitally using a computer based system. All patients had a diagnosis of unilateral hippocampal sclerosis, which was based on clinical grounds, EEG and volumetric MRI (1.5 Tesla) except for one patient who had a white matter lesion of uncertain nature in the temporal lobe. Seizures were right sided in 12 patients and left sided in 8. Tachycardia and bradycardia were defined as more than 120 and less than 50 beats/min respectively. Increase in heart rate was defined as an increase of more than 10% of baseline heart rate just before the clinical seizure onset.

![Graph showing EEG and ECG recordings](image)

*Fig 1. EEG shows rhythmical activity over left hemisphere, maximum in the antero-mesial temporal region. ECG (lead 1) shows sinus tachycardia at 168 beats/min.*
Fig 2. ECG shows a progressive sinus bradycardia with a pause of up to 9 seconds between R-R intervals.

Fig 3. The patient's habitual attack precedes the progressive sinus bradycardia. The concomitant EEG shows equivocal underlying semi rhythmic slow activity over the left temporal region at the clinical onset. Towards the end of the attack, the EEG shows widespread anterior predominant delta activity, associated with clinical hypotonia, probably reflecting cerebral hypoperfusion. Afterwards, the ECG shows a progressive recovery.
RESULTS

Sixty one clinical attacks were recorded. The ictal EEGs and ECG (lead I) traces were reviewed. The clinical features were those of complex partial seizures of temporal lobe, and one patient had a secondary generalised tonic clonic seizure.

Increase of heart rate were observed in 55 (90%) attacks. Significant ECG changes were observed in 27/61 seizures: tachycardia was seen in 24/61 (39%) and bradycardia in 3/61 (5%). The mean and median tachycardia rate was 139 and 140 beats/min (range 120-180) (Fig 1). The longest R-R interval was 9 seconds (Fig 2).

No difference was seen in regard with seizures lateralization and type of cardiac dysrhythmia albeit the numbers were small.

One of the ictal ECG was of interest (Fig 3). Clinically, the patient stopped his conversation, became vague and detached. His wife recognised the event as his typical attack. After the clinical onset, the EEG showed no definite changes, but the ECG showed progressive bradycardia. Then, the patient became hypotonic and the EEG showed widespread anterior predominant rhythmical delta waves probably reflecting cerebral hypoperfusion, and the ECG showed a progressive recovery. The patient was not considered a surgical candidate for other reasons. In view of the frequency of attacks and the concomitant ictal bradycardia, a demand pacemaker was fitted. This lead to a significant decrease in the number of his falls (> 50%), within 6 months of follow up.

COMMENTS

Changes in cardiac rhythm have been implicated as cause and the consequence of seizures disorders. Cerebral hypoperfusion due to cardiac arrest lasting more than 4 seconds, bradycardia (< 40 beats/min), or tachycardia (> 150 beats/min), my lead to variety of cerebral symptoms. It has been reported that patients referred to specialised centres for assessment of chronic epilepsy turn out to have underlying cardiac dysfunction in approximately 20-30%.

Experimental and clinical data have shown that cardiac rhythm may change during epileptic attacks, and this is dependent on the type of seizures. The ictal ECG changes observed in this study were tachycardia and less frequently bradycardia, which is similar to previous studies. Mechanisms of cardiac arrhythmia suggested in temporal lobe epilepsy include the spread of epileptic activity to autonomic centre, and emotional autonomic reaction in response to an aura. The ECG changes, however, may occur either in subclinical or clinical attacks or even in cases without aura in which case the latter mechanism seems unlikely as a primary cause, but in some cases may play an adjuvant role in ictal tachyarrhythmia.

Differentiating between TLE and cardiac dysfunction may not be easy in some cases, as clinical features of TLE may not be florid and may overlap with symptoms and signs of cardiac dysfunction. Assessment of these cases just based on ECG (Holter) without concomitant EEG may be misleading, as some cases of TLE with cardiac involvement may be considered primarily as a cardiac dysfunction instead of secondary to seizures. In a study carried out in a cardiac pacemaker clinic, patients referred were monitorized using simultaneous EEG and ECG, showed that 83% of patients with complex partial seizures might have been incorrectly diagnosed on referral. This suggests that patients with a vague history that hints either of an epileptic or a cardiac nature, simultaneous EEG/ECG monitoring is required. If the patient's seizures are frequent, video-EEG is therefore the choice, as in some cases of partial complex seizures scalp EEG may show no interictal changes.

The mechanism of SUDEP is uncertain, and secondary cardiac dysrhythmia appears to be an attractive explanation, however, experimental studies in animal models have shown that changes in
cardiac rhythm secondary to epileptic discharges involving subcortical nucleus were unable to lead to death in the experimental animal unless other concomitant factors were present, such as metabolic disturbances. AEDs are known to cause changes in cardiac rhythm. However, as far as SUDEP is concerned, it seems that they are unrelated, as SUDEP has been well described before the advent of any of the modern AEDs.

In conclusion, evaluation of patients with a long history of vague symptoms suggesting either an epileptic or a cardiac cause, who have been carefully investigated, concomitant ECG/EEG or video-EEG is worthwhile. Patients with well documented ictal bradycardia, and who are not surgical candidates, may be considered for a demand pacemaker. Further studies on this area are urgently warranted as this may shed light on possible mechanism for SUDEP.

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REFERENCES