FOURTEEN AND SIX C/SEC POSITIVE BURSTS IN COMATOSE PATIENTS *

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There have been numerous contributions regarding the clinical and etiologic significance of 14 and 6 c/sec positive bursts, aptly summarized by Gibbs and Gibbs (1952, 1963), Poser and Ziegler (1958), Henry (1963) and Hughes (1965), but none of the cases discussed by these authors were of an acute nature. Silverman (1964) observed 14 and 6 c/sec positive bursts in a patient with a coma due to infectious hepatitis and he stressed the rarity of this finding in an acutely ill, comatose subject. The purpose of this communication is to describe the presence and discuss the possible clinical significance of 14 and 6 c/sec positive bursts in acute coma due to Reye’s syndrome.

Material and method

A total of ten consecutive patients with Reye’s syndrome, age 7–14 years, were treated at the University of Michigan Mott Children’s Hospital during the early part of 1974. From one to five EEGs were obtained from each patient, using an 8-channel Grass electroencephalograph. A total of 30 EEGs were recorded. Scalp electrodes were placed according to the 10–20 International system.

The EEG abnormalities were classified into six grades (Aoki and Lombroso 1973): grade I: predominant theta and less alpha; grade II: predominant theta and less delta; grade III: diffuse high voltage rhythmic or arrhythmic delta activity with no or little theta activity; grade IV: diffuse mostly low voltage delta activity with no or little theta activity; grade V: nearly isoelectric activity; grade VI: electrocerebral silence.

The diagnosis of Reye’s syndrome was made on the basis of the clinical presentation, a course consistent with this syndrome (Reye et al. 1963; Bradford and Latham 1967; Hutttenlocher et al. 1969), and abnormal laboratory findings, specifically the blood ammonia level, serum glutamic oxaloacetic transaminase (SGOT) and serum glutamic pyruvic transaminase (SGPT).

Coma was graded into four stages (Silverman 1963):

Stage I: Arousal and cooperation at times were possible; if unstimulated, the patient was lethargic and tended to lapse into a comatose state.

Stage II: Arousal was difficult and was only possible after repeated and strong stimuli.

Stage III: All communication with the patient was lost except for a protopathic avoidance to noxious stimuli.

Stage IV: No behavioral responses to any external stimuli were observable.

Results

Three cases will be described in detail.
Case 1

A 10-year-old boy who was previously in good health developed a cough, malaise, and a low grade fever 5 days prior to admission. A few days later severe vomiting started and persisted until admission. A day prior to admission, the parents noticed that the patient was confused and irritable. The examination on admission revealed the patient to be delirious with agitated behavior and the patient then rapidly became stuporous. His temperature was 101°F (rectal); pulse rate was 150/min; respiratory rate was 38/min; blood pressure was 120/80 mm Hg. The liver edge was palpable and the spleen was slightly enlarged. His pupils were dilated but of equal size and reactive to light. There was a mild blurring of the disc margins bilaterally by funduscopic examination. He had increased muscle tone, hyperactive reflexes and bilateral Babinski signs.

Laboratory studies on admission revealed a serum ammonia level of 320 pg% (control 30 pg%), SGOT of 110 IU and SGPT of 480 IU. His prothrombin level was 33% of normal. A lumbar puncture was not performed because of suspected increased intracranial pressure. A few hours after admission, the patient became comatose and responded to pain in a decerebrate manner.

The first EEG was obtained 12 h after admission when the patient was in stage III coma with a dilated, non-reactive pupil on the left. The EEG was a grade IV with intermittent, diffuse 0.5–1 c/sec delta waves of less than 100 μV in amplitude (Fig. 1, A). Episodic 14 c/sec positive bursts were noted in the bioccipital region. The patient was then placed on a subdural pressure monitoring system and intracranial pressure was continuously monitored (Kindt et al. 1975). The second EEG was taken the following day when the patient showed some improvement. The EEG was grade III with higher amplitude and shorter wave length delta waves as compared to the first EEG (Fig. 1, B). Again, positive bursts were noted. The patient steadily improved and upon discharge he was neurologically normal except for a partial 3rd cranial nerve paresis on the left side. EEGs were obtained on the 5th, 6th and 12th day of hospitalization which were grades II, I and borderline, respectively. None of the last three EEGs showed positive bursts (Fig. 1, C).

Case 2

A 9-year-old male was well until 7 days prior to admission when he developed “flu-like” symptoms characterized by low-grade fever, cough, myalgia and lethargy. Two days afterwards, the patient began vomiting. On the night prior to admission, the patient was noted to be agitated and confused with occasional visual and auditory hallucinations.

On admission, the patient exhibited alternating agitation and stupor. His blood pressure was 140/95 mm Hg; pulse rate was 80/min; respiratory rate was 24/min. Pupils were slightly dilated but equal and reacted briskly to light. Funduscopic examination was normal. Slight hepatomegaly was noted. He moved all extremities spontaneously. Deep tendon reflexes were slightly increased, and there were bilateral Babinski signs. Laboratory data disclosed a SGOT of 1210 IU, a SGPT of 1430 IU and a serum ammonia of 100 μg% (control 30 μg%). The opening pressure on lumbar puncture was 100 mm water; the spinal fluid was clear and colorless with no cells; cerebrospinal fluid protein was 17 mg%; cerebrospinal fluid glucose was 97%. The EEG taken 6 h after admission when the patient was in stage II coma was a grade III characterized by high amplitude 2–3 c/sec delta waves (Fig. 2). Prominent 14 c/sec positive bursts with some 6 c/sec components were noted in occipital and temporal regions. The patient showed steady improvement and was discharged 2 weeks after admission.

Case 3

A previously healthy 8-year-old male developed fever and a sore throat 5 days prior to admission. Three days later he began vomiting repeatedly. He became progressively delirious.
Fig. 1. A: Grade IV EEG when patient was in stage III coma. Note 14 and 6 c/sec positive bursts (arrow). B: Grade III EEG when patient was in stage II coma on the following day. Note 14 c/sec positive bursts (arrow). C: Improvement of EEG on 12th day of hospitalization. Patient was alert. Note absence of positive bursts (light stages of sleep).
and agitated. On admission, his temperature was 98°F (rectal); pulse rate was 100/min; respiratory rate was 32/min; blood pressure was 124/62 mm Hg.

The patient's behavior was characterized by periods of agitation and delirium alternating with periods of stupor. His pupils were slightly enlarged but of equal size and reactive to light. Funduscopic examination revealed sharp disc margins. He had a mild systolic murmur. The liver and spleen were not palpable. He had increased muscle tone, hyperactive reflexes, sustained ankle clonus and bilateral Babinski signs. Laboratory findings revealed a SGOT of 180 IU, a SGPT of 400 IU, a prothrombin level of 40% of normal and a serum ammonia of 140 μg% (control 40 μg%). Lumbar puncture revealed clear and colorless cerebrospinal fluid with no cells, cerebrospinal fluid protein of 15 mg% and sugar of 95 mg%. The first EEG was taken 6 h after admission when the patient was in stage II coma. Prominent 14 c/sec positive bursts were noted in the right central, temporal and occipital regions (Fig. 3, A). Intermittent sleep spindles were also observed in the central area. At this time strong alerting stimuli abolished both sleep spindles and positive bursts but high voltage rhythmic delta slow waves remained dominant. The EEG was graded III. Second and third EEGs were obtained on the 2nd and 7th days of hospitalization and were graded II and borderline, respectively. Positive bursts were no longer present in either EEG (Fig. 3, B). The patient improved, and he was discharged after 8 days of hospitalization with normal motor and mental function.

**Cases 4 and 5**

These patients were 12- and 7-year-old males who were in stage III coma on admission. Their first EEGs were grade IV, and episodic 14 c/sec positive bursts were noted superimposed on the background of low-voltage delta waves, similar to the EEG of Fig. 1, A. Both patients expired on the third day of hospitalization.

The EEGs of five of ten patients who were in various stages of coma revealed 14 and 6 c/sec positive bursts in a background of diffuse delta waves of variable frequency and ampli-
Fig. 3. A: 14 c/sec positive bursts (arrow) appear when the EEG resembles spindle stage of sleep. Note sleep spindles on the right side of the figure (C₃). Patient was in stage II coma. B: Improvement of the EEG on 7th day of hospitalization. Patient was alert. Positive bursts were not seen (light stage of sleep).
TABLE I

<table>
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<td>V VI</td>
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B: Borderline.

- : EEG with 14 and 6 c/sec positive burst.

All the positive bursts but one (Case 2) were 14 c/sec without 6 c/sec components. Table I shows the gradings of the EEGs and the outcomes of the ten patients. The patients were divided into two groups with group 1 (Cases 1–5) having 14 c/sec positive bursts and group 2 (Cases 6–10) being without 14 c/sec positive bursts.

Out of six EEGs with positive bursts, three were grade IV (Cases 1, 4, 5), three were grade III (Cases 1, 2, 3). Three of the second group had grade V EEGs on admission with subsequent isoelectric EEGs (Cases 7, 8, 10). Only one case (Case 9), which had a similar degree of abnormality to the first group, failed to reveal the positive bursts. As a whole, six of eight EEGs graded from III to IV showed the 14 c/sec positive bursts. When the EEG became grade V or worse, positive bursts were no longer present (Cases 4, 5). Disappearance of positive bursts was also noted upon improvement (Cases 1, 3).

There were three survivors from group 1 and one from group 2.

Discussion

Since the early studies by Berger (1932) and Gibbs et al. (1937), diffuse EEG slowing has been regarded as characteristic of a variety of comatose states. Davis and Davis (1939), Schwab (1951) and Strauss et al. (1952) considered that the delta activity in comatose states was similar to that of deep sleep. Comatose states may be associated with a multitude of abnormal wave forms other than delta waves (Fischgold and Bounes 1946; Mathis et al. 1957; Loeb 1958; Fischgold and Mathis 1959). The presence of various sleep patterns such as sleep spindles, vertex sharp waves and K-complexes have been described in patients with coma of diverse etiologies (Jasper and Van Buren 1953. Chatrian et al. 1963; Silverman 1963). However, the presence of 14 and 6 c/sec positive bursts in comatose patients has been seldom reported in the literature. This is true even though positive bursts are most commonly seen during drowsiness or early sleep and certain workers have regarded the bursts as a distorted form of sleep pattern (Gibbs and Gibbs 1951) or even a type of sleep spindle (Kooi 1971). Silverman (1964) described one case of infectious hepatitis in which the 14 and 6 c/sec positive bursts were present during comatose state. Poser and Ziegler (1958) included a single case of hepatic coma due to Laennec's cirrhosis among 136 patients with various diagnosis who had positive bursts in their EEGs. In our own experience, positive bursts are also extremely uncommon in coma in patients of any age, al-
though the bursts are a common finding in normal adolescence. We have recently seen a single case of 6 c/sec positive bursts in a 40-year-old man with hepatic coma.

In view of the prevalent opinion that 14 and 6 c/sec positive bursts are generally found in normal, young persons (Walter et al. 1960; Lombroso et al. 1966; Eeg-Olofsson 1971), and that the present group of patients range from 7 to 14 years of age, it may be that the positive bursts were preserved in these young comatose patients as a type of sleep pattern. Sleep spindles were noticed in some cases of Reye’s syndrome studied by Aoki and Lombroso (1973) and also in one of our cases (Case 3). In the latter case, positive bursts were noted only when the EEG resembled sleep because of the presence of intermittent sleep spindles.

Although the etiology and pathogenesis of the Reye’s syndrome have yet to be elucidated, several studies have implicated short-chain fatty acids as potential toxins (Bourgeois et al. 1971; David et al. 1972; Trauner et al. 1972) and elevated concentration of serum short-chain fatty acid in Reye’s syndrome was reported (Trauner et al. 1975). Administration of short-chain fatty acids to experimental animals was shown to induce sleep patterns in EEGs and a further increase of dosage to result in irreversible coma (White and Samson 1956). These experimental results further support the observation that some of the EEG features of Reye’s syndrome resemble sleep patterns.

However, the following reasons raise the possibility that the positive bursts may not be simply preserved but are enhanced in some stages of coma in Reye’s syndrome: (1) This is an unusual prominence and frequency of occurrence of positive bursts in the face of actual reduction of other patterns characteristic of light to moderately deep sleep (Fig. 1, A, Cases 4 and 5. (2) Subsidence of positive bursts upon improvement is observed (Fig. 1, C; 3, B). This finding was also noted in the case reported by Silverman. (3) Positive bursts are found in association with EEG patterns resembling deep-level sleep, whereas they are usually seen with lighter levels of sleep in normal youths (Fig. 1, B; 2).

It is of further interest to note that our cases, Silverman’s case, and Poser and Ziegler’s case all had hepatic dysfunction. A causal relationship between hepatic disorder and positive bursts cannot be established with certainty because of the limited number of cases reported to date. The data suggest the need for further observations relative to a possible correlation between positive bursts and etiology of coma.

Summary

Of ten patients with Reye’s syndrome, there were five with stage II or III coma where EEGs revealed 14 c/sec positive bursts in a background of diffuse delta waves. Positive bursts disappeared upon EEG improvement in two survivors and when the EEG became nearly isoelectric in two other patients.

Although 14 and 6 c/sec positive bursts are seen commonly during sleep in normal young persons, their occurrence in association with diffuse delta waves in acutely ill, comatose patients has been rarely reported.

It is not certain whether the present findings should be regarded as selective preservation of a type of sleep pattern or whether there are special factors that enhance positive bursts in stage II or III coma of Reye’s syndrome.

Résumé

Bouffées positives de 14 et 6 c/sec chez les malades dans le coma

Chez 10 malades présentant un syndrome de Reye, 5 cas ont présenté un coma de stade II ou III dans lequel l’EEG a montré des bouffées positives de 14 c/sec sur un fond d’activité delta diffuse. Ces bouffées positives disparaissent lorsque l’EEG s’améliore chez deux survivants et lorsque l’EEG devient pratiquement isœlectrique chez deux autres malades.
Bien que les bouffées positives de 14 et 6 c/sec s'observent habituellement au cours du sommeil chez les sujets jeunes normaux, leur survenu en association avec des ondes delta diffuses chez des malades aigus comateux n'a été qu'exceptionnellement rapportée.

Il n'est pas certain que les données observées ici puissent être considérées comme une conservation élective d'un certain type de pattern de sommeil, ni qu'il existe des facteurs particuliers qui augmentent les bouffées positives au cours de comas aux stades II ou III de syndrome de Reye.

References


