Rett syndrome associated with continuous spikes and waves during sleep

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Abstract

Major speech impairment is a cardinal feature of Rett syndrome. Epilepsy, of variable presentation, is also frequently described. We report a girl who presented rapid speech regression while EEG showed continuous spikes and waves during slow sleep. The clinical picture was consistent with Rett syndrome, confirmed by MECP2 mutation R133C. We hypothesized that speech regression was partially related to nocturnal epileptic activity. Several antiepileptic drugs were used unsuccessfully but valproic acid was accompanied by improvement of verbal fluency, social interaction and manual abilities as measured by the Quebec Scale of Adaptive Behaviors and the Rett syndrome adapted Kerr scale. Continuous spikes and waves during slow sleep are unexpected in the early stages of Rett syndrome. This report suggests that it might contribute to the clinical impairment, in particular communicative capabilities, and that adequate antiepileptic treatment may be beneficial.

Key words: Rett syndrome; MECP2; Continuous spikes and waves during sleep; Speech delay.

Introduction

Rett syndrome is a neurodevelopmental disorder affecting girls almost exclusively, characterized by early neurological regression with severe motor, cognitive and communication impairment. Diagnosis is primarily based on clinical criteria including deceleration of head growth from the first years of life, regression with hand stereotypies, reduced manual and communication skills, scoliosis, abnormal gait and sleep disturbances (1, 2). Rett syndrome is caused by MECP2 mutations (3, 4), with some genotype-phenotype correlations (5, 6). Epilepsy, which is a frequent clinical problem, may have an atypical presentation, posing differential diagnosis difficulties (7). It is most common in stages II and III of the disease, but is rarely problematic beyond 10 years of age.

Seizures may display large variations in their typology: tonic, generalized tonic-clonic, complex partial, myoclonic, atonic, etc. (8). The electroencephalogram is almost invariably abnormal, showing loss of expected developmental features, rhythmic slow (theta) patterns primarily in the frontal central regions and diffuse or generalized spikes or spikes and waves (9). We describe a girl whose speech regression was contemporaneous with continuous spikes and waves during sleep, actually signalling the diagnosis of Rett syndrome.

Case report

A 3½ year-old girl presented with language regression from the age of three. Previous medical and family history was unremarkable with a normal birth weight and head circumference. Until 2½ years, milestones were achieved in the normal range, including communication skills. She then developed manual beat/friction stereotypies with progressive loss of manual function. From around 3 years, she lost previously reported openness toward social interaction. She had reduced eye contact and joint attention, seemed inattentive and did not respond to her name or to complex orders. Speech was limited to a few single words, and word repetition was not observed. She did not perform symbolic activities such as gesture imitation, setting in scene or drawing. She never presented any clinical seizures.

On clinical examination, head circumference was at the 25th percentile. Nonverbal communication was extremely poor with an inconstant visual contact. She showed hypermotor behavior, prominent stereotyped movements of both hands, severe bimanual dyspraxia and walking with marked lateral swinging. Her development was evaluated using the Quebec Scale of Adaptive Behaviours, Rett syndrome adapted Kerr scale and N-EEL scale for language (11, 12) (Table 1).
Biological work-up showed normal standard parameters. Brain MRI and auditory evoked potentials were normal. Heterozygous mutation R133C (g.66248 C > T) within the MECP2 confirmed the diagnosis of Rett syndrome.

Standard EEG showed occasional left centro-temporal discharges without any clinical correlation. 24-hour EEG showed continuous spike-and-waves during slow-wave sleep (CSWS) with a spike-wave index of 75% (Fig. 1).

Clobazam (5 mg bid) and ethosuximide (25 mg/kg/day; from age 3 years 8 months to 3 years 10 months), topiramate (4 mg/kg/day; from age 3 years 10 months to 4 years), leviteracetam (50 mg/kg/day; from age 4 years to 4 years 4 months) and prednisolone (2 mg /kg/d; 2 months; from age 4 years 4 months to 4 years 6 months) were not accompanied by clinical or EEG improvement. Valproic acid given from the age of 4 years 6 months on was associated with a moderate but significant improvement of verbal fluency (mostly in repetition but also in denomination and lexical comprehension), social interaction, manual prehension abilities (see Table 1) and a reduction of 37% in the spike-wave index after 8 months of treatment.

**Discussion**

This girl with Rett syndrome presented with electrical status epilepticus during slow-wave sleep.
Fig. 1. — Initial 24 hour EEG (3, 5 year-old). A) transition from wake to non REM sleep (see arrow); B) Non REM sleep; recording of CSWS with a spike-wave index of 75%. This index is defined as the total number of minutes of spikes and slow-wave abnormalities multiplied by 100 and divided by the total number of NREM sleep minutes. 30 seconds recording; vertical bar = 200 μV on EEG recording.
coinciding with speech and behavioral regression. This electrographic phenomenon has been well documented in continuous spikes and waves during slow sleep (CSWS) and Landau-Kleffner syndrome (LKS), where it contributes to progressive neurological dysfunction and behavioral alterations (13, 14, 15). CSWS typically starts in 4-8 year-old children (14, 15, 16), two thirds of whom have normal previous neurocognitive development (17). CSWS was also reported in children with various early brain lesions (13, 15, 18). EEG often shows focal spike-waves during wakefulness. During NREM sleep, spike-waves spread and become continuous, occupying 50-85% or more of the time of sleep (19). In LKS, CSWS is associated with acquired aphasia, usually developing over a few months and affecting both receptive and expressive speech. Antiepileptic medication commonly improves communication skills in children with LKS (13, 15, 18).

Our patient presented at 3 years with speech and general communication skills regression. CSWS was identified but the clinical picture was evocative of Rett syndrome in stage II of the disease, confirmed by the demonstration of a R133C mutation in the MECP2 gene. In stage II of the disease, sleep EEG shows frequent focal (central) or multifocal epileptiform abnormalities (9). In stage III, multifocal discharges and generalized slow spike waves patterns during NREM sleep are observed, often reaching the criteria for CSWS in stage IV (9, 10). To our knowledge, this has not been described previously at the early regression stage II, when the clinical diagnosis of Rett syndrome is commonly made.

Although CSWS and its implications on cognition have been poorly described in Rett syndrome, we find it reasonable to question its possible contribution to deteriorating some symptoms observed in our patient, such as speech and communication regression.

Patients harbouring the R133C mutation tend to show a milder phenotype, with better ambulation and hand use, and greater likelihood of using speech (12). Indeed, our patient showed normal development of motor and communications skills until the age of 2½.

In terms of treatment, only valproic acid induced improvement in verbal fluency, social interaction, manual stereotypies and spike-wave index. The observation period was however too short to consider long-term effectiveness of antiepileptic medications on the CSWS and its clinical repercussion.

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