Clinical and neuropsychological correlation in patients with rolandic epilepsy

Correlação entre achados clínicos e neuropsicológicos em pacientes com epilepsia rolândica

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ABSTRACT

Objectives: To evaluate the presence of neurological soft signs (NSS) and to correlate them with the Wechsler Intelligence Scale for Children (WISC III) in patients with rolandic epilepsy (RE). Methods: Forty children and adolescents aged between 9 and 15 years were studied. They were divided into two groups: G1 – patients with RE (n=20) – and G2 – healthy controls without epilepsy (n=20). They were assessed with the Quick Neurological Screening Test (QNST II) – clinical trial to search for NSS –, and the WISC III – neuropsychological test. Results: No statistical difference between groups was found in WISC III and QNST II. However, children with poorer motor skills had worse performance in the QNST II and also in the execution intelligence quotient – IQ (p=0.001) and in total IQ (p=0.004), thus showing a positive correlation between them. Conclusions: The QNST II is a good screening tool for the neurologist to detect abnormalities in fine motor skills.

Key words: epilepsy, rolandic epilepsy, neurological soft signs.

RESUMO

Objetivos: Avaliar a presença de sinais neurológicos menores (SNM) e correlacioná-los com a Escala de Inteligência de Wechsler para Crianças (WISC III) em pacientes com epilepsia rolândica (ER). Métodos: Foram estudados 40 crianças ou adolescentes entre 9 e 15 anos, divididos em dois grupos: G1 – pacientes com ER (n=20); G2 – controles saudáveis sem epilepsia (n=20). Foram avaliados por meio do QNST II – teste clínico que pesquisa SNM – e do WISC III – teste neuropsicológico. Resultados: Não foi encontrada nenhuma diferença estatística entre os grupos no WISC III e QNST II. Entretanto, crianças com habilidades motoras pobres tiveram pior desempenho no QNST II, assim como no quociente de inteligência (QI) de execução (p=0,001) e no QI total (p=0,004), mostrando, portanto, correlação positiva entre os dois instrumentos. Conclusões: O QNST II é uma boa ferramenta de rastreamento para o neurologista detectar anormalidades nas habilidades motoras finas.

Palavras-Chave: epilepsia, epilepsia rolândica, sinais neurológicos menores.

Focal epilepsy of childhood with centrotemporal spikes, or rolandic epilepsy (RE), is the most common form of childhood epilepsy, occurring in children between three and 13 years-old with normal psychomotor development, according to the International Classification of Epilepsies and Epileptic Syndromes¹. Patients may present with simple or complex partial seizures, most commonly involving the face, with or without involvement of the ipsilateral upper limb and, more rarely, evolving to generalized seizure.

RE is usually considered ‘benign’ because there is seizure remission during adolescence without future major impairments²⁻⁴. However, subtle neuropsychological deficits, behavioral disturbances, and difficulties in school performance, particularly in relation to reading and writing skills, may occur. It is still not clear if these abnormalities can be observed only in the active phase of epilepsy⁵⁻⁶, that is, if the transitory cognitive impairments occur exclusively during subclinical epileptiform discharges⁶⁻⁸. On the other hand, evidence suggest that such abnormalities are observed both before the active phase and during the postactive phase of epilepsy⁹.

The presence of neurological soft signs in RE patients has not been addressed. Our study aimed at investigating and correlating clinical features, through the search of neurological soft signs, with the neuropsychological deficits in patients with RE.

METHODS

Forty children and adolescents aged between 9 and 15 years old were studied. RE patients were selected from the outpatient Clinic of Childhood Epilepsy of our university hospital, and the Control Group consisted of healthy children and adolescents from the Department of Neurology, Faculdade de Ciências Médicas (FCM), Universidade Estadual de Campinas (Unicamp), Campinas SP, Brazil.

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Children and adolescents were divided into two groups: G1 (n=20) consisted of patients with RE according to the International League Against Epilepsy (ILAE) criteria, and G2 (n=20), the Control Group, consisted of healthy children without epilepsy. Most G1 patients (85%) had their epilepsy controlled (17 out of 20 patients).

All children and adolescents were assessed with the Quick Neurological Screening Test (QNST II), which is a clinical trial to search for neurological soft signs, and the Wechsler Intelligence Scale for Children (WISC III). The neuropsychological tests were applied by a qualified professional.

Data analysis was performed using the Statistical Package for Social Sciences (SPSS), version 17.0, and the Mann-Whitney’s test to verify the difference between the two groups.

RESULTS

Table 1 compares the QNST II data, considering its subdivisions and the chronological age of both groups. There was no statistically significant difference between them.

Table 2 shows the comparison between the final scores of the QNST II and neuropsychological assessments (WISC III) in patients who underwent both tests. There was no statistical difference between the groups. However, there was agreement between the QNST II (when searching for neurological soft signs) and the WISC III (assessment of intelligence quotient – IQ) with statistically significant results between total IQ and QNST II (p=0.004) and execution IQ and QNST II (p=0.001). Therefore, higher IQ correlated with better performance in the QNST II.

DISCUSSION

Epilepsy is often associated with cognitive impairment, particularly in children. The relationship between epilepsy and cognitive functions is multifactorial and involves age of seizure onset, seizure frequency, presence or absence of structural damage, use of antiepileptic drugs (mono- or polytherapy), and so on. RE is one of the most common forms of childhood epilepsy, accounting for 14 to 17% of children with epilepsy under 16 years of age. The clinical presentation is relatively homogeneous and there is no structural damage of the brain, which makes RE a good model to investigate neuropsychological, behavioral, and educational comorbidities. Neurological soft signs have not been investigated yet. To our knowledge, this is the first study to address this issue.

Although neuropsychological and behavioral changes in idiopathic benign childhood epilepsy have already elicited interest and subtle impairments, a wide variety of higher cortical functions (memory, visuomotor skills and visuospatial, spatial perception, memory spatial, coordination, different areas of language, content discrimination, and auditory-verbal dyspraxia) have been documented. Even after the active phase of seizures, little is known about neurological soft signs in such group of patients.

In our results, the QNST II showed no statistically significant difference between controls and children with RE. However, some considerations can be given. Firstly, neurological soft signs might be present in the active phase of epilepsy, and most of our patients (85%) were out of the active phase of seizures. Secondly, data correlation from the WISC III and QNST II, particularly the execution IQ with QNST II, should be emphasized. The QNST II is designed to detect soft neurological signs related to motor activity, coordination, and balance. Motor skills are necessary to carry out the tasks of both QNST II and execution IQ. Our findings showed that children with poorer motor skills have worse performance in QNST II as well as in execution IQ, thus showing a positive correlation between the two tests.

We have gotten to the conclusion that the QNST II is a good screening tool for the neurologist to detect abnormalities in fine motor skills, since there was correlation of the findings with the WISC III.
References


