Abstract

Objective. This report describes the main clinical features associated with specific reading disability (RD) in a group of 778 school-age children studied in a Neuropsychological Clinic in Mexico City. Material and Methods. The study was performed retrospectively, using data abstracted from clinical records of subjects seen in 1995-1996. Children were mainly from low and middle economic strata and aged between 6 to 12 years. The following data were collected: age, gender, diagnosis, school grade, food intake, maternal complications during pregnancy, perinatal and postnatal neurological risk factors, and neurological signs and handedness. Results. Subjects with RD had a mean age of 102.9 months, were predominantly male (male:female ratio, 2:1). Among the study group, 49.1% of the children were diagnosed with RD of a visuo-sensory-motor type, and 75.1% were from early school years (1st to 3rd grades). 27.6% showed evidence of malnutrition. A previous history of language disorders (49.2%), and a high frequency of perinatal risk factors and neurological soft signs were also found. Conclusions. This study shows that variables such as gender, food intake, and genetic and neurological risk factors, were associated with reading disabilities in school children.

Key words: reading disability; language disorders; Mexico

Resumen

Objetivo. Describir las características clínicas asociadas con los problemas específicos de la lectura en un grupo de 778 escolares estudiados en una Clínica de Neuropsicología Infantil de la Ciudad de México. Material y métodos. El estudio fue realizado retrospectivamente en los años de 1995 a 1996, con los expedientes clínicos de pacientes de estratos socioeconómicos medio y bajo, valorando las siguientes variables: edad, sexo, diagnóstico, grado escolar, alimentación, complicaciones maternas durante el embarazo, factores de riesgo pre-peri y posnatales para daño neurológico, enfermedades neurológicas y lateralidad manual de los pacientes. Resultados. Los pacientes con problemas de la lectura tuvieron un promedio de edad de 102.9 meses, fueron predominantemente del sexo masculino (2:1). Entre el grupo de estudio, 49.1% de los niños fueron diagnosticados con RD de tipo visuo-sensorial-motor, y 75.1% cursaba los primeros grados de escolaridad; 27.6% mostró evidencia de malnutrición. Un historial previo de trastornos del lenguaje (49.2%), y una alta frecuencia de factores perinatales y signos neurológicos blandos también fueron encontrados. Conclusiones. Este estudio sugiere que variables como sexo, alimentación, factores genéticos y diversos factores de riesgo para daño neurológico, se asociaron con la presentación de problemas específicos del aprendizaje de la lectura.

Palabras clave: problemas específicos de la lectura; problemas del lenguaje; México

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Received on: November 30, 2000 • Accepted on: April 3, 2002

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Childhood Reading Disability (RD) is a generic term that refers to an abnormal difficulty in learning to read and write, presumably as a result of neuropsychological abnormalities in otherwise healthy children. Some authors consider that reading and writing have not exerted evolutionary pressure on humans and therefore no brain specific biological adaptations have occurred for these skills. Effective treatment depends on early recognition of the underlying disease.

Population-based studies suggest that clinical characteristics of RD in children include: a preponderance of males (male: female ratio of 3.5:1), substandard performance in spelling, a frequent history of delayed language milestones, and common familial occurrence. These features suggest a biological substrate that some authors speculate might be related to cerebral lateralization. In developing countries like Mexico, the clinical characteristics of learning disability (LD) in children may be influenced by malnutrition and socio-cultural disadvantages, as has been suggested by Cravioto et al. The DSM-III and the DSM-IV definitions require that children with LD must have normal intelligence and show evident failure in reading and writing abilities.

The epidemiology of RD in underdeveloped countries presents a challenge to the health system, due to the lack of specialists and clinics for treatment of children with RD. In Mexico, only a small proportion of school children have the opportunity to receive care at the few neuropsychological clinics in place. Despite their small numbers, these clinics provide excellent care—although not population-based—information on learning disabilities. On the other hand, it is important to remind the health authorities of the need for systematic treatment and clinical care of children with development disorders.

The RD literature includes many proposals for classification and subtyping of this clinical entity. However, none of these proposals has gained general acceptance among researchers. Azcoaga et al. have proposed subtyping focused on developmental neuropsychological performance, such as language-based (expression and comprehensive delay) and non-language-based RD (underperformance in visuo-sensory-motor abilities). The first subtype is probably linguistic and the second one corresponds to the perceptual subtype described by Bakker. No previous studies have been performed in Mexico to identify the prevalence of both subtypes and additional clinical characteristics of RD children.

The goal of this paper is to describe the main clinical characteristics of school age children with RD, evaluated by the Children’s Neuropsychology Clinic at the Institute for Communication Disorders in Mexico City.

Material and Methods

The clinical records of 778 patients were reviewed; these subjects had been evaluated and diagnosed with specific reading disability during 1995-1996 at the Children’s Neuropsychology Clinic of the Institute for Communication Disorders in Mexico City. Patients were referred mainly by their school teachers or school administrators from neighborhoods surrounding the Institute, which is located in one of the most populated areas of Mexico City. The patients were residents of Mexico City aged between 6 and 12 years, from lower and middle-economic strata (86.2% and 13.7%, respectively, according to socioeconomic evaluation by social workers). A diagnosis of reading disability was based on reading age within two or more standard deviations below the mean for children of similar age and intelligence, as established with appropriate testing and a battery to assess writing and reading ability in Spanish. Diagnoses were made according to Azcoaga’s classification, with subtyping according to differential deficits. Three main groups were identified: expression delay, comprehensive delay, and underperformance in visuo-sensory-motor abilities. These tests have been administered for more than ten years in Mexico and are well-established at the Institute, which has now accumulated a vast experience in applying and interpreting them RD diagnosis.

Clinical histories were performed by three specialist in child communication disorders with a percent of concordance of 99% in neuropsychological diagnoses. Neuropsychologic and neuropsychologic tests were also performed on each patient but results from those tests are not included in the present report. Data from clinical records included: (a) age; (b) gender; (c) clinical diagnosis; (d) school grade level; (e) food intake noted in the clinic history using a questionnaire; (f) perinatal and postnatal neurological risk factors and maternal complications during pregnancy; (g) neurological examination, and (h) handedness. Descriptive data of the sample are presented as percentages and cumulative frequencies of the whole study population.

Results

A total of 778 subjects aged 6-12 years (average 102.9 months or 8.6 years) were included in the study. The male/female ratio was 2:1. Three hundred and eighty
two patients (49.1%) were diagnosed with a visuo-sensory-motor type of RD, three hundred and forty four (44.2%) showed a language plus non-verbal deficit alteration type, and fifty two (6.6%) were in the expression delay group.

Of the 778 children studied, 75.1% were in the first school grades (1, 2, 3) and the remainder (24.8%) in the higher grades of elementary school (Table I). Food intake was clinically qualified as a lower caloric/protein intake in 27.6% of the group (215 patients).

One or more maternal complications during pregnancy were noted in 21.4% of subjects. The main complications were: transvaginal bleeding (n=113, 67.7%), preclampsia (n=21, 12.6%), pregnancy after 35 years (n=21, 12.6%), and hypertension (n=12, 7.2%). The main perinatal risk factors found were: hyperbilirubinemia (n=121, 15.5%, including 3 patients who required transfusions, and 47 who received phototherapy), asphyxia (n=88, 11.3%), and preterm birth (n=55 cases, 7%).

Motor development was delayed in 60 patients (7.7%), and language development delay was detected in 383 patients (49.2%) prior to the diagnosis of RD. A postnatal neurological history disclosed mild head traumatism without loss of consciousness in 162 patients (20.8%), spells of apnea in 110 patients (14.1%), attention deficit-hyperactivity disorder in 35 patients (4.5%), and febrile seizures in 30 patients (3.8%).

The more common soft neurological signs were dysgraphystesias in 185 patients (23.7%), dysterognosias in 54 patients (6.9%), hand dyspraxias in 43 patients (5.5%), dysdiadochokinesia in 40 patients (5.1%), and dysmetry in 39 patients. The investigation of handedness showed that 98.4% (766 patients) were right-handed and only 1.5% (12 patients) were left-handed.

Discussion

The present work included only children treated in the neuropsychological clinic, and not the general elementary school population. Our results are thus limited only to the urban population with access to specialized health care services. Despite these methodological limitations, to our knowledge no previous studies have been performed in Mexico to describe the main clinical characteristics of RD children. No association between RD children and risk factors were explored and no statistical analyses were conducted to measure correlations between variables, thus in the discussion we only comment on possible relationships among variables that cannot be considered as strong conclusions.

It is now recognized that children with RD may present some neuropsychological alterations. Our sample shows that diagnosis were performed mainly from the first to the third grade of elementary school (75.1%, see Table I), which indicates the interest of parents and teachers of children with these conditions. Delayed diagnosis may result in mild disabilities. However, the average age in our study population (8.6 years) shows a delayed diagnosis, which may be due to the lack of information of school teachers leading to delayed identification and referral of RD children to the neuropsychological services.

Characteristics of RD in children from other studies include a higher prevalence in males, consistent with the male/female 2:1 ratio, similar to that found in a study conducted in the Isle of Wight. However, methodological and population differences between studies suggest caution in making comparisons. In underdeveloped countries such as Mexico, the parents have the idea that males will be the economic support of the family, and thus the female difficulties at school do not receive the same attention. Recently, some investigators have shown that this bias is important and suggest that boys and girls present similar frequency of RD. In the future this hypothesis deserves more attention.

The frequency of RD alterations in underdeveloped countries is unknown. In our study visuo-sensory-motor disabilities (49.1%) and mixed group (44.2%) were the most frequent types of alterations. Pure expressive language deficit were the less frequent (6.6%).

Some researchers believe that learning disabilities occur at higher rates in families with low nutrition levels, which may occur with greater frequency in underdeveloped countries. A relationship between poverty, malnutrition and academic failure is well established.

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Table I

<table>
<thead>
<tr>
<th>Grade</th>
<th>n</th>
<th>%</th>
<th>af</th>
</tr>
</thead>
<tbody>
<tr>
<td>1º</td>
<td>145</td>
<td>18.6</td>
<td>18.6</td>
</tr>
<tr>
<td>2º</td>
<td>260</td>
<td>33.4</td>
<td>52.0</td>
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<tr>
<td>3º</td>
<td>180</td>
<td>23.1</td>
<td>75.1</td>
</tr>
<tr>
<td>4º</td>
<td>112</td>
<td>14.4</td>
<td>89.5</td>
</tr>
<tr>
<td>5º</td>
<td>61</td>
<td>7.8</td>
<td>97.3</td>
</tr>
<tr>
<td>6º</td>
<td>20</td>
<td>2.6</td>
<td>99.9</td>
</tr>
</tbody>
</table>

% = percent
af = cumulative frequency
In our study we found that almost one third of the patients may have suffered some degree of malnutrition. Some of the main alterations in mental development on school performance correlated with infant malnutrition were: Delayed vocabulary and learning achievement, impaired visual perception and impaired visual-kinesthetic and auditory-visual intersensory integration. These conditions may help to explain poor school performance.

Our study results also seem to suggest a possible role of obstetric and perinatal events in the development of RD. Studies about risk factors for neurodevelopmental disabilities can prove this observation. The relationships between obstetric and neurological conditions of the infant were not clear in our sample. For example, transvaginal bleeding is a possibly damaging obstetrical risk factor that may increase neurological damage and secondarily RD, but its influence could not be ascertained; the only significant relationships found in the literature refer to asphyxia and RD. At present, the relationship between perinatal risk factors and learning disabilities is still under study. The majority of prospective studies deals with low-birth weight infants and indicates an increased frequency for learning disabilities, but the literature on this issue topic is not conclusive. In our study, the most common perinatal events and maternal complications were neonatal jaundice, asphyxia, and transvaginal bleeding.

Some investigators suggest that RD is related to language disorders. In our study we found that almost half our patients (49.2%) showed delayed language milestones. On the other hand, the parents of children diagnosed with specific language impairments showed a higher frequency of learning problems. Many studies suggest that there exists a familial factor that may reflect a genetic component. The Mendelian mode of transmission remains unclear, but recent works suggested that specific learning disabilities are associated with chromosomes 15 and 6. The soft signs more frequently found are those involving fine movements, asymmetries, dyskinesia, visual-motor coordination, sensory integration, laterality, right-left discrimination, and clumsiness. In our study dysgraphystesia was the most common sign.

An association between learning disabilities and left-handedness and immune diseases has been proposed previously, but it was not clearly observed in our study. Despite a direct search for immune disorders in the hospital records of our patients, only a few cases with auto-immune alterations were detected and thus are not shown in the tables. No surveys of learning disabilities in left-handed children in the general school population support this hypothesis, however some studies show that poor readers tend to have weak left hand skills. Our work was not designed to research this issue.

In conclusion, our data suggest that in a specialized Neuropsychological Clinic from Mexico City, reading disabilities may be associated to many variables such as: gender, language development, food intake, frequency of perinatal risk factors, and neurological soft signs. On the other hand, probabilistic population studies and school screenings must be undertaken in the future, to assess the true risk factors of reading disability behavior in Mexico.

Acknowledgments

We are grateful to Patricia Muñiz, MD for her collaboration, and to Robert Burns, PhD, for his help in correcting the manuscript. Thanks to the anonymous reviewer for recommending on-line information on Learning Disabilities in the website www.ldonline.org.

References

Reading disability in Mexican children