TALERO-GUTIÉRREZ, CLAUDIA; ROMERO, LILIANA; CARVAJALINO, IRMA; IBÁÑEZ, MILCIADES

Epidemiology of prelingual sensorineural hearing impairment at a children’s center in Bogotá, Colombia between 1997 and 2008


Universidad del Valle
Cali, Colombia

Available in: http://www.redalyc.org/articulo.oa?id=28318450009
Epidemiology of prelingual sensorineural hearing impairment at a children’s center in Bogotá, Colombia between 1997 and 2008

CLAUDIA TALERO-GUTIÉRREZ, MD¹, LILIANA ROMERO, MD², IRMA CARVAJALINO, AUDIÓL³, MILCIADES IBÁÑEZ, MS⁴

SUMMARY

Introduction: Hearing loss is a frequent problem in childhood with an incidence of about one case per 1000 births. Control of deafness should be aimed at prevention and early diagnosis in efforts to provide appropriate treatment and stimulate adequate communication in children affected. The objective of this study was to determine the prevalence of different etiologies among deaf children with a diagnosis of prelingual sensorineural hearing loss referred to the Fundación CINDA in Bogotá, Colombia, between 1997 and 2008.

Materials and methods: The medical records were selected from those with prelingual hearing loss. Information was gathered in a format containing variables related to the risk factors suggested by the Joint Committee of Infant Hearing.

Results: We studied 254 children; boys and girls were equally distributed. The most common etiological diagnosis was «unknown cause», followed by genetic causes (31 cases), and 38 cases from TORCH infections (toxoplasmosis, others – syphilis, rubella, cytomegalovirus, herpes), with rubella as the most common cause.

Conclusions: Review of prenatal, perinatal, and postnatal history often reveals the cause of the deafness in children; therefore, appropriate evaluation of pregnant mothers could result in decreased frequency of deafness in children in our country.

Keywords: Deafness; Sensorineural hearing loss; TORCH; Hyperbilirubinemia; Premature.

Colomb Med. 2011; 42: 199-206

Epidemiología de la pérdida auditiva neurosensorial pre-lingual en un centro de atención a niños de Bogotá, Colombia entre 1997 y 2008

RESUMEN

Introducción: La deficiencia en la capacidad auditiva es la alteración sensorial que con más frecuencia se encuentra en la literatura científica. La incidencia en la infancia es de un niño con pérdida auditiva profunda por cada mil recién nacidos. Los esfuerzos en el manejo de la pérdida auditiva deben estar orientados a la prevención primaria y a la detección temprana para estimular el desarrollo de la comunicación en los niños. El objetivo del presente trabajo fue establecer la prevalencia de los diferentes factores etiológicos en una población de niños sordos que asistieron a la Fundación CINDA, entre 1997 y 2008.

Material y métodos: Se seleccionaron las historias clínicas de aquellos con pérdida auditiva pre-lingual y se recolectó la información en un formato que contenía las variables relacionadas con los factores de riesgo planteados por el Joint Committee of Infant Hearing.

Resultados: Cumplieron el criterio de selección 254 niños (56.7% niños y 43.3% niñas). El diagnóstico etiológico más frecuente (47.2%) fue desconocido, 31 casos se identificaron como genéticos y 38 tuvieron diagnóstico de TORCH (sifilis,
toxoplasmosis, rubeola, citomegalovirus, herpes), entre los que se encontró con mayor frecuencia la rubeola.

**Conclusiones:** Los antecedentes positivos de tipo pre (63.4%), peri (37.8%) y post-natal (75.2%) permiten afirmar que un control adecuado del embarazo probablemente lograría disminuir en forma importante la incidencia de sordera en Colombia.

**Palabras clave:** Sordera; Hipoacusia neurosensorial; STORCH; Hiperbilirrubinemia; Prematuro.

**Colomb Med. 2011; 42: 199-206**

Deficiencia en la capacidad auditiva es la alteración sensorial más frecuentemente descrita en la literatura científica. La incidencia de sordera en la infancia es una de cada mil nacimientos (1-3). La pérdida auditiva presentada en esta etapa, previa a la exposición al lenguaje, es denominada pre-lingual; mientras que la presentada después de haber escuchado se llama post-lingual.

Los estudios epidemiológicos realizados en Europa revelan que más del 50% de los casos diagnosticados corresponden a causas genéticas, mientras que los relacionados con eventos ocurridos durante la gestación o al nacer, infecciosos, traumáticos, o otros procesos post-natales corresponden al resto de los casos (1-3). Las causas de pérdida auditiva han cambiado con las infecciones being the least frequent in developed nations, whereas in developing nations they continue being a frequent risk factor (3-5).

Los estudios epidemiológicos de la región americana publicados por Madriz (6) muestran la falta de una estructura adecuada de la información en la mayoría de los países de la región. Nobrega et al. (7) en el Instituto de Cirugía de la Universidad Federal de São Paulo, realizó un estudio retrospectivo y encontró que la etiología es desconocida en la mayoría de los casos (190 de un total de 442 pacientes). La rubéola congénita sigue en importancia con 67 pacientes, seguida por los genéticos, perinatales y meningíticos. Los datos analizados por Madriz sobre los casos de sordera en Colombia se refieren al año 1993 colombiano. En el país se realizó un censo de población con una prevalencia de 5.2 individuos con pérdida auditiva por cada 1000 nacimientos. In Colombia, the technical reports from the Health and Education ministries and from the Instituto Nacional para Sordos (INSOR) refer to the etiology of hearing impairments as having perinatal origin in 52.5% of the cases, unknown in 40%, and hereditary origin in 7.5%. The consolidated analysis from the data base of the Sistema Nacional de Información (National Information System) on the disability provides a prevalence of 12 per every thousand individuals with hearing loss, of which 18% present total hearing loss. Mencher and Madriz (9), calculated an incidence of 1.5 deaf children per every 1000 live births in Costa Rica.

Según los autores previamente mencionados, entre 38.75% y 50% de los diagnósticos de pérdida auditiva congénita se deben a causas genéticas. Algunas de estas infecciones pueden haber sido infectadas sin detectar el virus de la citomegalovirus (CMV). Este virus ha reemplazado la meningitis como la causa más común de sordera congénita en países desarrollados con incidencias variando entre 0.2% y 2%. Los efectos de las infecciones congénitas pueden variar desde una presentación asintomática hasta el desarrollo de un síndrome congénito que puede causar pérdida auditiva.

Bacterial meningitis is another one of the frequent causes of sensorineural hearing loss, embracing 60-90% of post-lingual deafness. In Colombia, the group led by Tamayo (12) conducted a screening study, between November 2005 and May 2006, in the country’s institutions for individuals with hearing loss. Ophthalmology studies were done on this population to detect “salt and pepper” retinopathy, typical of congenital rubella. A 33.5% positive frequency for rubella was found in a population of 1383 subjects from the main cities in the nation. These results urge this group of researchers state that prevention measures are not being effective to diminish the presence of this type of pathology nor its detection. This group proposes a visual screening as a basic tool in studying the population with hearing loss.

Efforts to manage hearing loss are aimed -in the first place- at early detection to therapeutically intervene and stimulate development of communication in children. Said detection is done through the hearing screening, and monitoring of children from high-risk pregnancies. Screening programs have been evaluated to verify the early-detection feature, confirming better development of communicative abilities in children with early diagnosis (12).

The Joint Committee of Infant Hearing identifies 11 risk factors of congenital permanent hearing loss, late onset or progressive, which are:

1. Concept of the parents or caregivers with respect to
MATERIALS AND METHODS

This is a correlation, descriptive, and retrospective study of children who attended the Fundación CINDA from 1997 to 2008. The inclusion criteria were the diagnosis of prelingual sensorineural hypoacusis; the exclusion criteria involved hearing losses of conductive or mixed type and post-lingual origin. Some 254 clinical charts were sequentially selected from those complying with the selection criteria.

Information was gathered in a format containing variables that evaluated family antecedents, presence of prenatal alterations like infectious processes, taking of ototoxic medications; perinatal alterations like weeks of gestation, hypoxia or suspicion of fetal suffering, and postnatal alterations of infectious and medication type and syndromes associated to hypoacusis and others related to the risk factors suggested by the Joint Committee of Infant Hearing. A previous pilot test was carried out.

The information was systematized on an Excel 2007 data base. This work was carried out by a general physician, a speech therapist, and medical students who received prior training in handling clinical charts and in the search and selection of data to be analyzed.

For data quality control and to avoid bias, we kept in mind the confirmation of the hearing loss secondary to a sensorineural deficiency and not to conductive processes; clinical charts without complete information were excluded and the information was purged with logical frequencies and cross-referencing to identify inconsistencies in the information gathered.

Statistical analysis. Information was processed in SPSS version 16 and STATA 10 statistical packages. In the description of etiological factors in general, in qualitative variables, measurements of absolute and relative frequencies were established expressed in percentages, and in quantitative through measurements of central tendency with the average, median, and dispersion, the range and standard deviation.

For the purpose of evaluating the association between etiological factors and sex and the magnitude of hearing loss, we used Pearson’s Chi square test of association or the likelihood exact tests and Fisher’s (expected values <5). These tests were evaluated at a 5% (p<0.05) significance level.

Ethical aspects. Each patient was assigned an
identification code to maintain the confidentiality of the information. This study was classified without risks according to norms established by resolution 008430 of 1993 by the Ministerio de Salud de Colombia (scientific, technical, and administrative norms for health research). The recommendations of the Helsinki Declaration by the World Medical Association were followed. It was a study of observational design and lacking of direct intervention with the children. Upon admission to the institution, all parents signed an authorization letter, which permits using the information related to their children in teaching and research activities. This was done bearing in mind the functions of the Fundación CINDA, from the care, academic, research, diffusion, and social sensitivity points of view. This study was approved by the Ethics Committee of the School of Medicine and Health Sciences at Universidad del Rosario.

RESULTS

The total population completing the selection criteria was 254 patients of which 144 (56.7%) were boys and 110 were girls (43.3%). The minimum age to enter the study was 2.5 months, and a maximum of 16 years of age with an average of 3.7±2.7 years. The degree of hearing loss was classified according to 1996 ANSI criteria, presented by Ross and Downs14, and bearing in mind the configuration of the audiometric curve by frequency ranges (Table 1).

The audiological diagnosis carried out in the population revealed profound hypoacusis of the right ear in 126 (49.6%) of the children and severe to profound hypoacusis in 53 (20.9%). For the left ear, profound hypoacusis was found in 129 (50.8%) children and severe to profound hypoacusis in 51 (20.1%) children. Hearing loss was severe in the right ear for 32 (12.6%) children and in the left ear for 31 (12.2%) children. In total, 211 children were classified in the highest levels of hearing loss.

The minimum age for the mother at the moment of giving birth was 13 years of age; the maximum was 44 years of age with a mean age of 25.6±6.5 years.

In 141 children (55.5%), some type of positive family antecedent was found (Table 2) among which it is worth remarking hearing loss, delay or other type of language alteration, psychomotor delay, and neurological pathology.

Prenatal antecedents were positive in 161 (63.4%) cases (Table 3). In these, it is important to note the frequency of infection during pregnancy by agents like rubella, syphilis, toxoplasmosis, and cytomegalovirus, which constituted 15.7% of the cases. There were 10 children with other infections not classified at the moment of doing the clinical history, among which any of the aforementioned agents may be found.

Threat of abortion, hypertension, and ingestion of prescribed medications, including antibiotics, and urinary infection were the most frequent adverse events during gestation.

Among the perinatal antecedents, positive in 96 (37.8%) of the population studied, it was found that in 7.4% of the children under 30 weeks of gestation and

<table>
<thead>
<tr>
<th>Degree*</th>
<th>dB HL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal hearing</td>
<td>0 - 10</td>
</tr>
<tr>
<td>Minimum hypoacusis</td>
<td>11 - 25</td>
</tr>
<tr>
<td>Slight hypoacusis</td>
<td>25 - 40</td>
</tr>
<tr>
<td>Moderate hypoacusis</td>
<td>40 - 55</td>
</tr>
<tr>
<td>Moderate to severe hypoacusis</td>
<td>55 - 70</td>
</tr>
<tr>
<td>Severe hypoacusis</td>
<td>70 - 90</td>
</tr>
<tr>
<td>Profound hypoacusis</td>
<td>&gt;90</td>
</tr>
</tbody>
</table>

* Ross and Downs, 2004 - ANSI 1996
25.8% between 31 and 35 weeks, there were 3 newborns who weighed less than 1000 g. In total, 65 children were underweight at birth (Table 4).

Sixty percent of the children measured less than 50 cm at birth. Eleven presented sepsis (4.3%) and there is data in the charts regarding perinatal hypoxia in 74 children (29.1%).

Positive post-natal antecedents were found in 191 children (75.2%) among which the highest frequency was found in oxygen requirement, phototherapy, exanguinotransfusion, upper respiratory infections, and middle-ear infections, as well as meningitis among other causes according to that observed in Table 5.

Etiological diagnosis. For each patient, a causal diagnosis was done considering for each case what was the principal etiological factor for the hearing loss; nevertheless, in some cases there was concurrence of several significant etiological factors that were included in the definite diagnosis (Table 6).

The first issue coming to our attention is that almost half the children (120) corresponding to 47.2% have a diagnosis of unknown etiology. There are 31 cases clearly identified as genetic and among these there is a boy with Waardengurg syndrome and three cases identified with alteration of 26-Conexine, as well as two with Trisomy 21.

Rubella and other prenatal infections were the etiological diagnosis in 38 children (rubella 31, syphilis 4, toxoplasma 1, cytomegalovirus 1, and herpes 1). Events related to perinatal risks: prematurity (19), hyperbilirrbinemia (8), perinatal hypoxia (16), and toxemia (3) (in total 18.1% of the population) were important etiological factors; while hearing loss was...
attributed to meningitis in 8 children.  

**Relationship among variables.** Upon verifying the degree of hearing loss in children in relationship to gender, it was found that 53.6% of the girls had profound hearing loss of the right ear and 57.3% of them had the same degree of hearing loss in the left ear; while the boys revealed the same type of hearing loss in lower percentages for the right ear (46.5%) and for the left ear (47.8%). This does not constitute a statistically significant difference between the degree of hearing loss and gender.

No gender differences are noted with family antecedents related to delayed or altered language. When analyzing the antecedent of hypoacusis, we found that boys have more antecedents for hypoacusis (18.8%) than the girls (11.1%), yielding p: 0.068. In antecedents related to STORCH and other prenatal factors, no differences were found between boys and girls.

Infection of respiratory pathways was significantly more frequent in boys (30.6%) than in girls (19.1%) with p:0.026. The presence of identifiable syndromes like Gregg’s and Down’s syndromes, and others was found most frequent in females tan in males with p: 0.039. There was no significant correlation between the etiological diagnosis and the gender of the children.

**DISCUSSION**

The results obtained reveal an important frequency of positive antecedents in the pre-, peri-, and post-natal stages, which could have been triggering or coadjuvant factors of the hearing loss. Likewise, several of these factors converged on particular cases of children at high-risk because of their gestation conditions, birth, and perinatal period.

Upon reviewing family antecedents, these were positive in 55.5% of the cases. Hypoacusis was found as a factor present in the family, which substantiates the risk of hearing loss in children due to genetic factors, as well as delayed language that can be associated to hearing losses and unspecified neurological pathology.

In 63.4% of the children, prenatal antecedents are positive and among these the greatest relevance is related to STORCH-type infections with rubella being the most frequent (30 children). We found 10 cases of unspecified viruses that could correspond to both undiagnosed rubella and cytomegalovirus. Blood hypertension in the form of preeclampsia or toxemia and the threat of abortion during the first trimester constituted an equally frequent risk factor. Although there are reports of ingestion of medications in an important number of mothers, in general these correspond to vitamins, iron, and broad-spectrum antibiotics like ampicillin, amoxicillin, and others that are not related to deleterious effects in the fetus.

In many cases, there is history of prematurity and underweight, as well as hypoxia and hyperbilirubinemia sufficiently important to have required phototherapy or exanguinotransfusion and support oxygen.

The infectious causes during the early period of the newborn and their complications, along with exposure to ototoxic medications, also constitute risk factors for hearing loss.

Upon conducting a definite diagnosis, we kept in mind the causal factor in each case that weighed more in the history, aside from the clinical and para-clinical exams done at different stages of the pregnancy and the post-natal period. A total of 47.2% of the cases were

<table>
<thead>
<tr>
<th>Table 6</th>
<th>Etiological diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Etiologies*</td>
<td>Nº</td>
</tr>
<tr>
<td>Unknown</td>
<td>120</td>
</tr>
<tr>
<td>STORCH (38)</td>
<td></td>
</tr>
<tr>
<td>Rubella</td>
<td>31</td>
</tr>
<tr>
<td>Toxoplasma</td>
<td>1</td>
</tr>
<tr>
<td>Syphilis</td>
<td>4</td>
</tr>
<tr>
<td>Cytomegalovirus</td>
<td>1</td>
</tr>
<tr>
<td>Herpes</td>
<td>1</td>
</tr>
<tr>
<td>Genetic (1 Wanderburg syndrome, 2 Trisomy 21, and 3 26-Conexine)</td>
<td>31</td>
</tr>
<tr>
<td>Prematurity</td>
<td>19</td>
</tr>
<tr>
<td>Perinatal hypoxia</td>
<td>16</td>
</tr>
<tr>
<td>Hyperbilirubinemia</td>
<td>8</td>
</tr>
<tr>
<td>Meningitis</td>
<td>8</td>
</tr>
<tr>
<td>Feverish episodes</td>
<td>8</td>
</tr>
<tr>
<td>Toxemia</td>
<td>3</td>
</tr>
<tr>
<td>Ototoxic drugs</td>
<td>2</td>
</tr>
<tr>
<td>Upper respiratory infection</td>
<td>1</td>
</tr>
</tbody>
</table>

* Of this group of children, 9 presented double diagnosis; for this reason, the total number of children increases by this proportion

attributed to meningitis in 8 children.

---

The presence of identifiable syndromes
classified as unknown etiology. Only 23 cases were

diagnosed as clearly genetic.

According to the authors mentioned, there is evidence
of growing numbers of cases reported of unknown
etiology, corresponding to non-syndrome genetic cau-
ses. Other cases may be the result of infections like
CMV, considered more and more an important
etiological factor for hearing loss and that may go
unnoticed from a clinical point of view. In many cases,
when not finding relevant data complementary exams
are not performed and there is no confirmation if the
loss is due to a pre or perinatal infection.

If we compare the data herein reported with the
screening studies conducted by Tamayo\textsuperscript{12} in centers for
individuals with hearing deficiencies in the nation, we

can note that in the figures reported in her study there
is prevalence for the diagnosis of congenital Rubella in
30\% of the population and in the group studied at the
Fundación CINDA that percentage, although still
important as an etiological factor, only constitutes
12.2\% of the population. It is also possible that some of
the causes defined as unknown may correspond to
undiagnosed rubella and if the recommended rear-of-
the-eye exam had been performed on these children as
a fundamental part of the exam, more cases would have
been found attributable to this cause.

Vaccination campaigns against rubella, carried out
as part of the primary prevention policy by the Minis-
terio de la Protección Social (Colombian Ministry of
Social Protection) since August 2005, are still not
reflected in the current results.

The 7.5\% of children whose etiology is based on
prematurity and all the risk factors underlying to this
condition can generate false hypoaucus cases that
corresponded to early stages of hearing maturity, as
stated in the study by Talero et al.\textsuperscript{15}

The important percentage of antecedents related to
pre and perinatal risk factors highlight the importance
of improving controls to mothers during gestation.

The definition of the epidemiological characteristics
of a population with a specific pathology, and in this
case in particular with hearing losses, determines the
effects of health policies and supports plans aimed at

carrying out primary prevention.

The etiology of hearing loss permits knowing the
origin of the alteration and determining the consequences
in terms of the functional development of the individual’s

hearing and, hence, of communication. It is clear in the
data reported in this study that there is a high percentage
of infectious causes like hypoxia and hyperbilirubin-
emia, where the compromise is not exclusively in the
hearing, but which is associated with neurological-
type alterations that may affect -to a greater extent- the
child’s communicative development. Etiologies of
genetic or ototoxic type that generate a specific alteration
of hearing permit foreseeing what will be the therapeutic
focus and the prognosis in each patient, bearing in mind
the different tendencies of rehabilitation in hearing
deficiencies.

\textbf{Conflict of interest.} None of the authors has conflicts
of interest related to this study.

\textbf{REFERENCES}

4. Debeba K, Janssens de Varebekeb S, Coxb T, Van de Heyning P. Epidemiology of hearing impairment at three Flemish
7. De Nóbrega M, Weckx LL, Juliano Y. Study of the hearing loss in children and adolescents, comparing the periods of
Bogotá: Vicepresidencia de la República, Consejería Presi-
dencial de Política Social, Ministerio de Educación Nacional,
Pontificia Universidad Javeriana, Corporación Interinsti-
tucional de Rehabilitación; 1995.
278-83.
10. Samilhe N, Ahmad S, Mohammad F, Framarz M, Azarodkht T, Jomeht E. Role of cytomegalovirus in sensorineural hearing