Comparison of two newborn hearing screening protocols with distinct reference criteria of distinct pass and failure

Comparação de dois protocolos de triagem auditiva neonatal com critérios de referência de passa e falha distintos

Priscila de Araújo Lucas Rodrigues(1)
Taina Maiza Bilinski Nardez(1)
Mariano Martinez Espindola(2)
Keyla Cristina Costa Gomes(1)
Bruna Luana da Silva(1)

ABSTRACT

Purpose: to compare two newborn hearing screening protocols with benchmarks of passes and failure distinct.

Methods: a retrospective study. It was evaluated the records of 312 infants (s) and babies of both sexes and up to 90 days after birth in September 2013 to September 2014 in the Neonatal Hearing Screening Service of Clinical Speech and Hearing Division School of the origin institution. TEOAE were analyzed in two different benchmarks and they were compared.

Results: of the 312 patients evaluated, there was a greater number of male patients (53.85%), the majority of patients aged 30 days or less (65.06%) and only 6.09% had more than 60 days. The proportion of observed agreement was 43.91%, that is, both methods coincide in 43.91% of the results and the Kappa index was 0.0628, with a confidence interval of 95% (0.03 ; 0.0942) and 0.001 statistical significance (p = 0.001). It is observed that the value of Kappa is very low, considering that the perfect agreement is 1.00.

Conclusion: there was no statistical correlation between the protocols analyzed, there were more failures in the NHS by step benchmark protocol 1.

Keywords: Screening; Optoacoustic Emissions; Newborn

RESUMO

Objetivo: comparar dois protocolos de triagem auditiva neonatal com critérios de referência de passa e falha distintos.

Métodos: estudo retrospectivo. Foram avaliados os prontuários de 312 indivíduos, entre eles, RN(s) e bebês de ambos os sexos com até 90 dias de nascidos, sendo de baixo e alto risco para a deficiência auditiva, no período de setembro de 2013 a setembro de 2014 no Serviço de Triagem Auditiva Neonatal da Clínica Escola da instituição de origem. Os prontuários consultados referem-se a RN(s) ou bebês submetidos à triagem auditiva neonatal por meio das Emissões Otoacústicas Evocadas Transientes (EOAT). As EOAT foram analisadas considerando tanto o critério de referência proposto por FINITZO (1998) (PROTOCOLO 1) quanto os critérios de referência vindos de fábrica no equipamento (PROTOCOLO 2), e os mesmos foram comparados. Foi realizada uma análise descritiva para caracterização da amostra.

Resultados: dos 312 indivíduos avaliados, observou-se maior número do sexo masculino (53.85%), a maioria tinha 30 dias ou menos (65.06%) e somente 6.09% tinham mais de 60 dias. A proporção de concordância observada foi de 43,91% e o índice de Kappa foi de 0.0628, com o intervalo de confiança de 95% de (0.03 ; 0.0942) e uma significância estatística de 0.001 (p=0.001). Observa-se que o valor do Kappa é bem baixo, considerando que a concordância perfeita é de 1.00.

Conclusão: não houve concordância estatística entre os protocolos analisados, houve mais falhas na etapa da TAN pelo critério de referência do protocolo 1.

Descritores: Triagem; Emissões Otoacústicas; Neonato
INTRODUCTION

Through hearing the individual develops his capacity of speech and language, which, in turn, guarantees the development of his understanding and expression. Sensory deprivation affects the communication, the language, the literacy and his social and emotional development.

The Neonatal Hearing Screening (NHS) enables the early diagnosis of child hearing impairment, and should be done soon after birth or in the first month of the newborn’s life (NB) so as to have a diagnosis in the first three months and any intervention in up to the first six months, reducing the impairments caused by hearing loss, and making possible the development of language very close to that of a child that can hear1. The NHS is feasible due to its low cost, its facility and speed2.

The Joint Committee on Infant Hearing (JCIH)3 and the Multi-professional Committee on Auditory Health1 both suggest that the implementation of the Neonatal Hearing Screening use electroacoustic and electrophysiological measures, such as the Otoacoustic Emissions Evoked by Transient Stimulus (OAET) and the Auditory Brainstem Response (ABR).

The OAET is an objective test which evaluates the cochlear function without depending on neurological conditions of the newborn. It is a fast, noninvasive technique of easy interpretation, high specificity and sensitivity, performed when the baby is sleeping or quiet. An OAET is generated from a nonlinear click which stimulates the cochlea as a whole, from the basis to the top, in the frequencies from 300 to 5000 Hz, resulting in an acoustic energy, generated by contractility of the outer hair cells during the active mechanism of the cochlear function that propagates to the middle ear and external ear canal, where it can be sensed4.

This test does not quantify hearing loss, but detects the presence of answers in individuals that have auditory thresholds of up to 30dB.

The ABR assesses the neuro-electrical activity in the auditory pathway, from the auditory nerve to the brain cortex, in answer to an acoustic stimulus. The reception of the auditory evoked potential is done using electrodes that are fixed on the scalp, forehead, ear lobes or mastoids. The captured neuro-electric answers are submitted to a filtering and amplifying process and, after being separated from the devices and added, can be observed as waves in the computer5.

There is no obligatory protocol defined as pass or fail criteria for the OAET, although there are some suggestions in literature1,6-8. The equipment that analyzes the otoacoustic emissions comes with defined pass or fail criteria, however these do not correspond to those mentioned in the literature.

The study of sensitivity and specificity of screening protocols used in NHS services are of great importance, as well as the search for definition of a universal protocol aiming at an increase of reliability of the results obtained by services which exist worldwide. Also, a standardization of pass and failure reference criterion of these OAET, mainly for Hearing Screening, would make the comparison of the results in scientific research easier and would increase the evidence of effectiveness of NHS services.

This research aims at comparing two different protocols of neonatal hearing screening with benchmark reference criteria of passes and failure.

METHODS

This study was initially sent for ethical consideration to Plataforma Brasil, being approved under the protocol number 38226314.3.0000.5164.

A retrospective study through the analysis of medical records was carried out, in which records were analyzed considering specifically hearing screening which occurred from September 2013 to September 2014 at the Neonatal Hearing Screening Service of the section of Speech Therapy at the School Clinic, belonging to the researcher’s institution of origin in Mato Grosso State, Brazil.

The results of the OAET of 312 individuals of both genders and ages of up to 90 days were analyzed.

The inclusion criteria were newborn(s) or babies of up to 90 days old, of both low and high risk. The exclusion criteria were: infants evaluated any time after completing 90 days of age, or who presented inconclusive results due to lack of occlusion of the external auditory canal, or extreme noise at the moment of evaluation and non-attendance for retesting. Babies of up to three months of age were included, since, despite the national and international organs stating that the NHS must be performed before the end of the first month of life, several infants’ parents search NHS services after this period.

The medical records checked refer to newborn(s) or infants submitted to the neonatal auditory screening through reception of the transient evoked otoacoustic emissions. The equipment used was a model Otoread made by Inter-acoustics. The stimulus presented was a non-linear click a 60dBpNPS (± 5). This equipment...
analyzes the functionality of the outer hair cells and discards the possibility of hearing loss.

Two protocols were used for the analyses of the results. Protocol 1 follows the international reference criterion, \(^6\) that considers the newborns’ response which occurs from 3 to 6 dB above the noise (3dB in the frequency ranges of 1500Hz and 6 dB in the frequency ranges of 2000Hz, 3000Hz and 4000Hz).

Protocol 2 follows the reference criteria from factories that produce Otoread equipment, where 6 frequency ranges are tested (1,500Hz, 2,000Hz, 3,000Hz, 3,500Hz and 4,000Hz) and to pass the screening, there must be at least 3 frequencies with a sign-noise relation (S/R) of 4dB.

A comparison of the results obtained by the NB(s) and babies evaluated was done, considering both the reference criterion proposed by PROTOCOL 1 and the reference criteria which came from factories of the Otoread equipment (PROTOCOL 2).

Finally, a descriptive analysis was carried out for the description of the sample and inferential in order to compare the results obtained by both reference criteria applied. For such a comparison the non-parametric Kappa test was used, where two groups with different qualitative results are compared to verify their concordance.

**RESULTS**

In Table 1, the characterization of the 312 individuals evaluated is presented according to the variables gender and group age in days (corrected age), at the moment of the evaluation. In this table it is observed that there is a larger number of male patients (53,85%), most of the patients were 30 days old or less (65,06%) and only 6,09% were more than 60 days old.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Frequency</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gender NB</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>168</td>
<td>53,85</td>
</tr>
<tr>
<td>Female</td>
<td>144</td>
<td>46,15</td>
</tr>
<tr>
<td><strong>Age group in days</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 to 15 days</td>
<td>80</td>
<td>25,64</td>
</tr>
<tr>
<td>16 to 30 days</td>
<td>123</td>
<td>39,42</td>
</tr>
<tr>
<td>31 to 45 days</td>
<td>36</td>
<td>11,54</td>
</tr>
<tr>
<td>46 to 60 days</td>
<td>54</td>
<td>17,31</td>
</tr>
<tr>
<td>More than 60 days</td>
<td>19</td>
<td>6,09</td>
</tr>
</tbody>
</table>

Legend: NB = Newborn

In Table 2 the comparison between the result obtained according to protocol 1 and 2 is presented. The values presented are given by the number of patients and respective percentage in each classification.

In Table 2, the observed proportion of concordance was 43,91%, that is, both methods coincide on 43,91% of the results and the Kappa ratio was 0,0628, with the confidence interval of 95% of (0,03; 0,0942) and a statistic significance of 0,001 (p=0,001). It can be observed that the Kappa value is very low, considering that the perfect concordance is 1,00.

<table>
<thead>
<tr>
<th>Result Protocol 2</th>
<th>Result Protocol 1</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Failed</td>
<td>Failed</td>
<td>15 (4,81)</td>
</tr>
<tr>
<td>Passed</td>
<td>Passed</td>
<td>175 (56,09)</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td>190 (60,90)</td>
</tr>
</tbody>
</table>

Table 1. Sample characterization according to gender and age group variables

Table 2. Comparison of the results according to protocols 1 and 2
DISCUSSION

Of the 312 participants in this research, 53.85% were males, and 46.15% females (Table 1). According to other authors, the percentage of male newborns or babies that seek NHS service is higher, and the majority are under 30 days old\(^1\),\(^2\). The age of the newborns that did the NHS varied from 0 to 90 days old, and the majority was less than 30 days old. These numbers are in accordance with what is recommended by national and international organs that indicate that neonatal hearing screening should be done in the first month after birth\(^1\),\(^3\).

The literature describes some national and international studies using different pass and fail criteria, indicating that there is not yet an agreement about which would be the best criteria a speech therapist should choose, using the transient evoked otoacoustic emissions in neonatal auditory screening\(^8\),\(^11\).

In this study, the number of newborn(s) or babies that failed in protocol 1 was higher than that obtained by protocol 2, and both have low concordance, as can be seen on Table 2. A possible explanation is that, as protocol 1 is more thorough, it can detect a higher number of auditory losses, avoiding false negatives. However, this same rigor might lead to an excessive number of false positive results and generate problems in the quality of the program, generating unnecessary anxiety in parents as well as increasing referral for repetition of tests of fails of OAET and creating an overload in the clinics responsible for the auditory diagnostic\(^12\),\(^13\).

Another relevant result is that the number of fails in protocol 1 is higher than the literature considers acceptable (4%) – Table 2. Maybe the protocol has low sensibility, thus generating many false-positive results. Another hypothesis for this finding is that the 1.5 KHz frequency evaluated by protocol 1 can negatively influence the results of NHS service, for it is highly liable of fail as shown in a study\(^11\) where it was confirmed that the OAE register in lower frequency ranges can be disguised by noisy environments and physiological noises.

It was not possible to verify the sensibility and specificity of the protocols in this study as the research focused only on the tests results and neonatal auditory screening. It is suggested that future studies seek to assess the same newborn(s) through ABR to confirm or discard auditory loss. Therefore, it was not possible to confirm that the babies that failed at NHS actually had a confirmed hearing loss in the diagnostic stage.

Thus, the great importance of introducing a universal neonatal auditory screening should be emphasized. One that is sensitive and specific enough to avoid false positives and false negatives and that will make worldwide scientific studies easier.

CONCLUSION

- There was no statistical concordance between the analyzed protocols.
- There were more fails in the NHS stage by the reference criteria of protocol 1.

REFERENCES


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REFERENCES


**ERRATUM**

In this article, “Comparison of two newborn hearing screening protocols with distinct reference criteria of distinct pass and failure”, with DOI number: 10.1590/1982-021620161842816, published in the journal Revista Cefac, 18(4):876-880, on page 876:

Where it was:
Taina Maiza Bilinsky Nardez

Read:
Taina Maiza Bilinsky Nardez