P300: Waves Identification with and without Subtraction of Traces

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Abstract

Introduction The P300 test requires well-defined and unique criteria, in addition to training for the examiners, for a uniform analysis of studies and to avoid variations and errors in the interpretation of measurement results.

Objectives The objective of this study is to verify whether there are differences in P300 with and without subtraction of traces of standard and nonstandard stimuli.

Method We conducted this study in collaboration with two research electrophysiology laboratories. From Laboratory 1, we selected 40 tests of subjects between 7–44 years, from Laboratory 2, we selected 83 tests of subjects between 18–44 years. We first performed the identification with the nonstandard stimuli; then, we subtracted the nonstandard stimuli from the standard stimuli. The examiners identified the waves, performing a descriptive and comparative analysis of traces with and without subtraction.

Results After a comparative analysis of the traces with and without subtraction, there was no significant difference when compared with analysis of traces in both laboratories, within the conditions, of right ears (p = 0.13 and 0.28 for differences between latency and amplitude measurements) and left ears (p = 0.15 and 0.09 for differences between latency and amplitude measurements) from Laboratory 1. As for Laboratory 2, when investigating both ears, results did not identify significant differences (p = 0.098 and 0.28 for differences between latency and amplitude measurements).

Conclusion There was no difference verified in traces with and without subtraction. We suggest the identification of this potential performed through nonstandard stimuli.
Introduction

P300 is an auditory-evoked potential denominated endogenous because it reflects the functional use individuals make of the auditory stimulus, being highly dependent on cognitive abilities, including attention and auditory discrimination. It is an objective procedure which depends on the examiner’s experience in detecting the peaks of the waves. A methodology for the identification of traces that facilitates the analysis of presence of response and interpretation of results is relevant in this case.1

The P300 potential provides a window to observe the neurophysiological substrate of processes that occur in the cerebral cortex related to cognition, memory, and auditory attention.2 The feature is the most often reported in research focusing on hearing. However, latency and amplitude measures are variable and must be adjusted to age and to the studied population.3

For the P300 analysis, examiners should take into account possible interference that contribute to its variability, such as the parameters used in the test (intensity, frequency and type of acoustic stimulus evoking, filter, type of task, interstimuli interval, among others), the identification conditions (time of day), the subject (age/maturity, sex, cognitive ability, body temperature), and the examiner’s skills for verifying such a potential.4

According to Schochat,5 P300 is identified as a wave of positive polarity with latency of ~300 milliseconds post-stimulus, obtained after subtracting the trace corresponding to nonstandard stimuli from the trace corresponding to standard stimuli.

Recent research with diverse populations have performed the identification of P300 in nonstandard traces, without performing the subtraction from the standard trace, and have found reliable results.6–8

Although P300 is considered an objective test, a single, well-defined criteria, and training of its examiners is essential for a more uniform analysis of studies, and a verification whether or not subtraction of the traces is necessary to benefit from these potentials.

In this context, our study aims to determine whether there is difference in P300 with and without subtraction.

Material and Method

This study received approval from the Research Ethics Committees of both institutions involved in this research under the Research protocols Number 842/2010 and 12790/2011, respectively.

This is a documental, experimental, and descriptive study. It was a partnership between two Electrophysiology Research Laboratories in Hearing.

The examiners of both services were experienced in Auditory Electrophysiology, specifically in Long Latency Auditory Evoked Potentials.

From Laboratory 1, we selected and analyzed 40 cognitive auditory evoked potential tests of subjects (24 women and 16 men) with normal development, previously performed in participating volunteers from previous research in the same field. The subjects were of both sexes, aged between 7 and 44 years, without cognitive impairment, according to history available in medical records, and audiometric thresholds within the normal range (20 a 30 dBnHL).9

From Laboratory 2, we selected and analyzed 83 cognitive auditory evoked potential tests of subjects (44 women and 30 men) with normal development and audiometric thresholds within the normal range (20 dBnHL), previously performed in participating volunteers from previous research in the same field. The study subjects were men and women aged 18–44 years.

In Laboratory 1, for P300 assessment, we used Biologic Navigator Pro and captured the responses with active electrode in the central region Cz, referring to the electrodes positioned in the right lobe (A2) and left (A1) and the ground electrode at the forehead (Fpz). In Laboratory 2, we used 3A insert phones, the acoustic stimulus was tone-burst at 70 dBnHL, presented randomly by the computer, at a proportion of 20% nonstandard stimuli of a total of 200 stimuli, identified in a 500 milliseconds window, 100.000x amplification, alternating polarity, with bandpass filtering of 0.5–30Hz, monaural stimulation, and stimulation rate of 1.1/second.

For frequency discrimination, we used a standard stimulus at a 750Hz frequency and nonstandard stimulus at 1000Hz frequency with 20ms - rise/fall and 60ms - plateau, both with duration of 100ms. Subjects were instructed to pay attention and identify random nonstandard stimuli within a series of standard stimuli and name them out loud: fine to 1000Hz.

In Laboratory 2, for the P300 evaluation, we used Bio-Logic equipment, version 5.70, model 317 (Bio-Logic Science Instruments S.A.S, Seyssinet-Pariset, France). We captured responses with active electrodes positioned at Cz and Fz and connected to the input of a preamp channels 1 and 2, respectively. We placed the reference electrodes on the earlobe (A1 and A2), interconnected and connected to input 2 of channel 1, and interconnected to channel 2 by the preamplifier jumper. We placed the ground electrode at the forehead (Fpz). We used 3A insert phones, the acoustic stimulus was presented through tone-burst at 75 dBnHL, binaurally, presented randomly at a proportion of 20% stimuli of a total of 240 stimuli, identified in a 500ms window, amplification 100.000x, alternating polarity, with band filtering of 1–30 Hz and 1.1 stimuli per second stimulation rate.

For frequency discrimination, we used a standard stimulus at 1000 Hz frequency and nonstandard stimuli at 2000 Hz frequency with 20ms rise/fall and 60ms plateau, both with duration of 100ms. Subjects were instructed to pay attention and identify the nonstandard stimuli (random) within a series of standard stimuli, counting mentally.

The tests took place with two successive passages to allow good definition and replication. Impedance was maintained at 3 kW or less.

The tests lasted ~50 minutes. As a protocol to ensure quality examination, we suggested to those with myogenic interference that they change position. We also ensured quality by controlling the answers for a maximum of 5% deviation between stimuli data and patient responses and,
when necessary, repeating the test. Also, we asked the subjects to keep their eyes closed to eliminate eye artifacts.

For the identification of P300 waves, we instructed all examiners to use the criterion proposed by Junqueira and Colafêmina adapted as follows: identification of N1-P2-N2 complex—the first three waves that appear in sequence—and present the polarities: negative - positive - negative, respectively, occurring in the replication of traces, standard and nonstandard between 60 and 300ms; P3 identification, the highest positive wave, right after N1-P2-N2 complex, occurring in trace replication to nonstandard stimulus, between 240 and 700ms. We identified latencies at the highest peak, that is, the maximum wave amplitude point. We identified amplitudes at the wave peak to the base line and inter-amplitude in the case of N2-P3 inter-amplitude; when there was duplication of P3 wave in P3a and P3b, we always performed the identification in the second wave, P3b. In addition, the identification of waves always occurred in the identification of the nonstandard stimuli.

We first analyzed and identified the tests for nonstandard stimulus, then performed a second identification by subtracting the nonstandard stimulus from the standard stimulus. Examiners were instructed according to pre-established criteria for identifying the potential cognitive waves.

In this study, we chose to focus on the analysis of P300 with and without subtraction, since this has been the most used analysis in research involving auditory electrophysiological assessments.

As a study that analyzes the intersubject response, we did not consider small variations of the protocols in the services as variables that could affect the results and objectives of this study.

We performed a descriptive analysis of the test results, and then conducted a comparative analysis using Student t-test among the variables of P300 latency and amplitude, comparing these variables identified in the traces with subtraction and without subtraction. We only analyzed the active electrode arrangement positioned in Cz, a position that ensures records with better morphology and greater amplitude.

We describe results as p-value, and the significance level was always 5% or 0.05 (p ≤ 0.05). The statistics software used was Bioestat (Instituto de Desenvolvimento Sustentável Mamirauá, Tefé, Brazil).

Results

Table 1 shows the mean value, standard deviation, and p-value of P300 latency and amplitude, respectively, from Laboratories 1 and 2 after statistical analysis through t-test. We found no significant differences between the measures with and without subtraction.

Figs. 1 and 2 present a representative box plot comparing the measures with and without subtraction from both laboratories.

Discussion

LLAEP is one of the most promising measures in objective auditory assessment as it comprehends from simple listening skills to the most complex ones. According to Magliaro, auditory-evoked potentials have been useful tools for functional diagnostics, allowing one to observe, through an increase in latency or decrease in amplitudes, objective evidence of clinical and sub-clinical problems.

This potential’s analysis, although an objective procedure, is highly subjective, depending on good clinical experience to visually detect the waves. In addition, standardizations for their identification are required to avoid variations of traces and difficulties in the interpretation of the study. One of the procedures for interpreting the results is whether there is variation when traces are analyzed with subtraction or no subtraction of the nonstandard stimulus waveform from the standard stimulus wave.

Table 1 Descriptive statistics and p-values: t test of P300 latency and amplitude of measures with and without subtraction

| Group   | Ear | Analysis | Variable | Mean   | SD    | Min   | 1st Quartile (25%) | 3rd Quartile (75%) | Max  | p Value t Test |
|---------|-----|----------|----------|--------|-------|-------|--------------------|-------------------|------|----------------|
| Laboratory 1 | R   | with sub | lat_P300 | 335.82 | 32.35 | 275   | 308               | 352.50            | 246  | 0.136          |
|         |     |          | lat_P300 | 339.78 | 35.40 | 246   | 319               | 356              | 415  |                |
|         |     | without sub | amp_P300 | 6.25   | 2.77  | 2     | 4                 | 8                | 2    | 0.286          |
|         |     |          | amp_P300 | 6.59   | 2.69  | 13    | 4                 | 8                | 14   |                |
|         | L   | with sub | lat_P300 | 327.17 | 35.29 | 274   | 301.5             | 346.50            | 411  | 0.158          |
|         |     |          | lat_P300 | 330.44 | 41.92 | 246   | 304               | 355.75            | 416  |                |
|         |     | without sub | amp_P300 | 5.69   | 3.01  | –     | 4                 | 7                | 12   | 0.095          |
|         |     |          | amp_P300 | 5.81   | 3.12  | –     | 4                 | 8                | 15   |                |
| Laboratory 2 | BIN | with sub | lat_P300 | 289.90 | 34.02 | 225   | 264               | 315              | 371  | 0.095          |
|         |     |          | lat_P300 | 296.89 | 34.33 | 237   | 271               | 325              | 409  |                |
|         |     | without sub | amp_P300 | 7.15   | 4.10  | –     | 4                 | 9.50             | 21   | 0.280          |
|         |     |          | amp_P300 | 7.46   | 3.82  | 1     | 5                 | 10               | 22   |                |

Abbreviations: R, right; L, left; BIN, bineural; amp, amplitude (µV); SD, standard deviation; lat, latency (ms); sub, subtraction.
In this study, after a comparative analysis of the traces with and without subtraction, we found that none of the variables showed statistically significant results, suggesting no difference in identifications of traces with and without subtraction. Thus, we suggest that identification of this potential be conducted for nonstandard stimulus traces.

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