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Estimation of Single Brainstem Auditory Evoked Potential Using Time-Sequenced Adaptive Filtering

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Abstract - Time-sequenced adaptive filtering (TSAF) technique has been successfully used to track variation of brainstem auditory evoked potential (BAEP) in human. 400 ensembles used by TSAF will produce a satisfactory result rivaling to EA using 2000 ensembles. Furthermore, single BAEP signal can be obtained which make it possible for clinicians to analyze the variation of signals across trials.

Keywords: Time-Sequenced Adaptive Filter, Brainstem auditory evoked potential, Ensemble averaging.

INTRODUCTION

Evoked potentials (EPs), which are brain electrical activities resulting from sensory stimulation, are time-locked signals. Ensemble averaging (EA) is the most widely used method for obtaining the EPs. However, this method ignores the fact that the EPs are also time-varying signals and the results of EA may introduce significant distortion and loss of information about the response variability from trial to trial. In this paper, we present an adaptive filtering method based on [1] to track variation of brainstem auditory evoked potential (BAEP) and improve the SNR.

METHOD

Fig. 1 shows a conceptual realization of TSAF. Differing from conventional adaptive signal enhancer, the TSAF uses multiple filters and each filter is adapted for filtering a particular portion of the interval between regeneration times, and is suitable for tracking signals whose statistical properties recur at various points in time. The filters are trained and the filter weights are obtained via an adaptive algorithm.

The TSAF filter has been applied to detect BAEP in humans. The stimulus pulse was of 0.1 msec duration and the click rate was 10 Hz. The 100 dB sound pressure level (SPL) was given to a subject. The response was recorded at a sampling rate of 10 kHz. The first 9 ms of response has been processed for BAEP evaluation. The SNR of the raw data of human is about -20 dB in our case. The regeneration time were set at the instant of stimuli were given. The sequence number was initialized at the regeneration time and then incremented after every point until 90 set of weights have been cycled through. Each set contains 25 weights.

RESULTS AND CONCLUSIONS

Using TSAF we can track the variation across trials after convergence. Fig. 2 shows two results of single BAEP in human estimated by TSAF. The SNR of single trial was be significantly improved and the five peaks are clear. Fig. 3 gave the trace of BAEP of 600 trials. Since the result of EA is still the standard of studying BAEP in clinical application, it is necessary to make a comparison the result of averaged TSAF with that of EA. We use 400 trials to estimate an averaged result of TSAF. First 200 trials are used for training the filters and last 200 trials processed by TSAF are averaged to get an averaged estimate of BAEP. The averaged TSAF is shown in Fig. 4 together the EA using 2000 trials. TSAF and EA are very similar and their correlation coefficient is 0.983, and the measurement time greatly reduced using TSAF. The tracking ability of TSAF makes it possible for the clinician to observe the signal variation trace in every single ensemble which may be helpful for the detail analysis of EPs.

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REFERENCES


Fig. 1 The time-sequences adaptive filter

Fig. 2 Two individual BAEP waveforms estimated by TSAF

Fig. 3 Trace of BAEP of 600 trials. (a) a 3D view of BAEP trace. (b) a gray level 2D view of (a).

Fig. 4 A comparison of averaged TSAF (solid line) and EA (dotted line)