Delayed Auditory Tone Perception in Multiple Sclerosis

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SUMMARY: Delays of auditory perception at three frequencies were measured in 30 multiple sclerosis patients using a psychophysical technique. Nineteen patients had abnormal delays at one or more tone frequencies, though 15 had normal audiograms at those frequencies. In addition, auditory acuity for left-right asynchrony was abnormally poor in 13 patients, 9 of whom had normal audiograms. Such delays of auditory perception within a restricted frequency band may provide a partial explanation for degraded speech comprehension in some multiple sclerosis patients.

RÉSUMÉ: Chez 30 patients souffrant de sclérose en plaques nous avons mesuré la perception auditive à trois fréquences en employant une technique psychophysique. Dix-neuf patients avaient des réponses retardées pour une ou l’autre des fréquences tonales, même si 15 de ces patients avaient des audiogrammes normaux à ces fréquences. Chez 13 patients il y avait aussi une anomalie de l’asynchronie droite-gauche; 9 de ces patients avaient des audiogrammes normaux. De tels retards dans la perception auditive pour certaines fréquences restreintes pourraient expliquer partiellement la compréhension diminuée de la parole notée chez certains patients atteints de sclérose en plaques.


The rationale for this study is based on the finding that, in some multiple sclerosis (MS) patients, visual perception is markedly delayed in one region of the visual field though perception is delayed much less or not at all in nearby regions only a few degrees away (Heron, Regan, and Milner, 1974; Regan, Milner, and Heron, 1976). Among possible explanations for such delays of visual perception, the most likely is demyelination of central nervous system (CNS) axonal fibres (Koles and Rasminsksy, 1972; McDonald, 1974).

Authors who compare the visual and the auditory pathways commonly point out the analogy between the frequency of an auditory tone and location in the visual field or spatial frequency of a visual stimulus (Julesz, 1980). In particular, since position in the visual field has a topological representation in the optic radiations on visual cortex, it is easy to imagine how a local plaque of demyelination might slow or disrupt signal transmission from only a local region of the visual field.

We surmised that, in analogous manner, a local plaque of demyelination and/or neural damage in the auditory pathway might slow transmission of auditory signals originating in a local region of the basilar membrane. We were already aware that MS could produce a general slowing of auditory signals, since it has been reported that evoked potentials originating in auditory brainstem were delayed in some MS patients (Robinson and Rudge, 1975).

The present study was undertaken to determine whether MS patients experience delayed auditory perception and, if so, whether such delays can be restricted to a limited range of tone frequencies.

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The data reported below were obtained using the method of ascending and descending limits. The duration of the stimulus tone was 1250 msec.

Tone onset was linear over 10 msec and tone intensity was 50 dB SL. In the first run the right ear’s tone started between 200 and 250 msec before the left ear’s. Both tones switched off together. The subject pressed one of two buttons, depending on whether he judged that the left ear’s tone started before the right ear’s or vice versa. Then the two tones were presented again with stimulus onset asynchrony (SOA) reduced by 30 msec. The process was repeated, reducing asynchrony in steps of 30 msec until the subject judged the tone to start in the left ear twice in succession. A second run was then carried out with the left ear’s tone switching on 200-250 msec before the right ear’s, and SOA being again reduced in 30-msec steps. The starting point was randomized at SOA between 200 and 250 msec. Four runs were carried out for each of the five different tone combinations, except for 1000 Hz versus 1000 Hz which was run eight times — four runs at the start and four at the end of the session. This 1000Hz/1000Hz combination was used to assess reproducibility. Every session started with several practice runs.

Apparatus

All psychophysical sessions were controlled by a microcomputer (Commodore PET, 32K). Stimulus tones were generated by circuits of our own construction in which outputs were attenuated (Hewlett-Packard model 350D) and amplified (Citation 12 DeLuxe) before driving electrostatic earphones (Stax model SRX Mk 3). Auditory calibration was carried out by coupling the earphones to a Bruel and Kjaer model 2215 Sound Level Meter via an artificial ear (Bruel and Kjaer model 4152). Subjects sat in a soundproof chamber (Industrial Acoustics model 403-A). Audiograms were measured for each patient and control subject by means of a Grason-Stdler model 1704 audiometer. We regarded an audiogram as abnormal when the hearing loss exceeded 20 dB at any of the test frequencies (500, 1000 or 2000 Hz).

Patients and control subjects

Patients were diagnosed according to the criteria set out by Schumacher et al. (1965) and classified into “clinically definite”, “probable”, and “possible” groups (Rose et al., 1976). In addition, each patient was rated on the 10-point disability scale of Kurtzke (1965). A total of 30 patients with multiple sclerosis were tested, comprising 7 males and 23 females with ages ranging from 19 to 66 years. Clinical data are summarized in Table I. We did not require that the patients should have prior evidence of auditory involvement to be included in this study.

Eleven naive subjects were tested, comprising 4 males and 7 females, ranging in age from 18 to 60 years.

RESULTS

Interaural delay

Interaural delays were measured for 30 patients and 11 control subjects. Interaural delays in control subjects ranged from -95 msec to +83 msec with a mean of 18 msec and SD of 14 msec (Figure 1A). (Negative values correspond to left ear leading while positive values correspond to right ear leading.) For patients, interaural delays ranged from -167 msec to +126 msec with a mean of 27 msec and SD of 27 msec (Figure 1B). MS patients showed significantly longer interaural delays than did the control subjects [Mann-Whitney U test for large samples: z = 3.13, p = 0.0009 (Siegel, 1956)]. In the five experimental conditions (tone combinations), only 4 of 66 control tests (6%) exceeded the 2.5 standard deviation (< -17 msec or > +53 msec), while 45 out of 180 tests (25%) on patients exceeded this criterion.

Of the 30 patients tested, 19 showed abnormal interaural delay at one or more tone frequencies. One, two, or all three test frequencies could show an abnormal delay. Ten patients had a delay in both ears. In 15 of these 19 patients, the frequency-selective delay was associated with a clinically normal audiogram (i.e. less than 20 dB loss) in the affected part of the auditory spectrum. To emphasize this point further, we note that in the parts of the auditory spectrum where interaural delay was abnormal, 13 patients had pure tone thresholds for the left and right ears that differed by only 5 dB or less, 4 patients had a 10-dB difference, 1 differed by 15 dB, and 1 differed by 20 dB.

Although test-to-test repeatability was poorer than we could have wished in quantitative terms, patients with abnormally large delays generally gave delays outside normal limits on the second and subsequent retests, while patients whose delays were inside control limits generally remained within control limits on the second and subsequent retests.
Acuity for binaural synchrony

For each tone combination, we computed the difference between the mean interaural delay for the two runs in which the stimulus began in the left ear and the mean interaural delay for the two runs in which the stimulus began in the right ear. We took this difference as a measure of acuity for binaural asynchrony. For control subjects the range of values was 0 msec to 195 msec with a mean of 62 msec and SD of 45 msec (Fig. 2A), while for MS patients the values ranged from 0 msec to 427 msec with a mean of 99 msec and SD of 72 msec (Fig. 2B). MS patients showed significantly greater ranges than the control subjects did (Mann-Whitney U test for large samples: z = 3.84, p < 0.00007 (Siegel, 1956)).

The 2.5 standard deviation criterion (175 msec) was exceeded in only 1 of the 66 control tests (1.5%), while it was exceeded in 25 of 180 tests on patients (14%).

Of 30 patients tested, 13 had abnormally low acuity at one or more test frequencies. Of these 13 patients, 9 had normal audiograms in the relevant parts of the auditory spectrum. To emphasize this point further we note that, in the parts of the auditory spectrum where acuity was abnormal, 8 patients had pure tone thresholds for the left and right ears that differed by 5 dB or less, 4 patients had a 10 dB difference, and 1 differed by 15 dB (Table I).

**DISCUSSION**

**Delays of auditory perception that varied with tone frequency**

Our chief conclusion is that auditory perception can be delayed over a restricted frequency band, even though perception is not delayed for tones of other frequencies. In particular, significant variations of delay can exist between tones as close as 500 and 1000 Hz, or 1000 and 2000 Hz. These frequency-specific delay can occur in the absence of any abnormality in pure tone intensity thresholds.

The frequency-dependent delays might be explained in terms of local plaques of demyelination and/or neural damage if we can assume that such pathology occurs at a neuroanatomical site where signals evoked by different frequency tones are spatially separated.
Acuity for binaural asynchrony

The range of interaural delays over which the tones were judged to be simultaneous was abnormally broad in a substantial number of patients (13/30). We take this range as a measure of acuity for binaural asynchrony.

In studies of temporal order discrimination, Jerger et al. (1969) reported control subjects to have an interaural resolution of 20-40 msec in a paradigm similar to ours. Our slightly poorer thresholds (62 msec in controls) can probably be attributed to less experienced subjects. Like Jerger et al. (1969) we used stimuli of random phase. This enabled us to compare temporal resolution of different frequencies.

Hasler and Levine (1980), found interaural time discrimination was degraded in 13 of 29 patients, though their technique differed in that the same (noise) waveform was presented to each ear. Their phase-locked stimuli enabled subjects to use phase cues which our stimuli did not. (Phase cues permit lateralization of stimuli with only a fraction of a millisecond discrepancy.)

Relation to SBMPL test results

Our psychophysical findings may relate to Noffsinger et al.'s (1972) report that 12/60 patients with MS gave abnormal results on the simultaneous binaural median plane location (SBMPL) test. In this test the same pure tone was presented to left and right ears, and the intensity of the stimulus to one ear was varied until the patient perceived the sound source to be located at the midline. Of 60 patients tested, Noffsinger et al. reported that 12 either could not experience a midline image or did so only when different intensities were presented to the two ears. They underlined their finding that, of these 12, only 5 had difficulties in performing the task, contrasting this with the well-known observation that patients with brainstem lesions often find the SBMPL task difficult since they do not perceive a single fused image or cannot shift the image.

Frequency-dependent delays and speech comprehension

We find that auditory perception can be delayed even when the audiogram is normal. In other words, the intensity threshold for pure tones may be unaffected even though perception is delayed. Nevertheless, it is possible that delayed auditory signals might degrade everyday hearing, especially when delays are different for different pure tones. In particular, it seems possible that speech comprehension would be affected by distorting the relative timing of auditory signals originating in different parts of the basilar membrane. Thus, frequency-dependent delays might degrade speech comprehension in MS patients, even though the disorder might not be detected by pure tone audiometry. Frequency-dependent delays might be a factor in the unexplained poor performance on speech sound recognition tests of some patients with MS who have normal audiograms (Citron et al., 1963; Noffsinger et al., 1972; Sidtis et al., 1979).

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REFERENCES


