Sudden Hearing Loss: Ten Years’ Treatment Experience

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SUMMARY

Objective: Determine the follow-up between patients sudden sensorineural hearing loss (SSHL) treated in five different groups according to duration of disease and the drugs used for treatment.

Study design: Retrospective cohort study.

Method: Review of patients records included in a standardized protocol of treatment of the SSHL Outpatient unit between January 1996 and January 2006. Inclusion criteria were SSHL occurring within 72 hours and 30 dB or more at three serial frequencies. Patients were included in five treatment groups and had received Dextran®, dexamethasone, acyclovir, nicotinic acid, papaverine chloridrate and or vitamin A. Audiometric assessment was done before and after treatment (30, 90, 120 and 180 days). Outcome was determined by subtracting baseline PTA (first day of the medical assessment) from final PTA (after 180 days).

Results: Complete data were collected for 139 subjects, mean age were 45,4 years, 52,5% were female and 47,5% were male, 92,8% had unilateral SSHL and 7,2% had bilateral one. PTA mean before treatment were similar between groups. Treatment significantly improved PTA mean after 180 days. Nevertheless, no significant difference could be noted in audiometric assessment between treatment groups. We observed a significant linear correlation between time from SSHL onset to initial visit and audiometric recovery.

Conclusion: Dextran® did not provide more benefit than steroid and acyclovir in the treatment of SSHL. As soon as treatment begins better will be the outcome prognosis for SSHL.

Key words: sudden hearing loss, sensorineural hearing loss, treatment, steroid, acyclovir, dextran.
INTRODUCTION

The sudden sensorineural hearing loss (SSHL) is defined as auditory loss (AL) neuro-sensorial, equal or higher than 30 decibel (dB), in 3 or more consecutive frequencies, installed in a period of up to 3 days (1). Most of the times it is severe, not floating, unilateral and idiopathic (1, 2). In about one third of the cases can occur associated complaints as tinnitus, vertigo or dizziness and aural fullness (2, 3). Its incidence is from 5 the 20 cases per 100.000 inhabitants/year in the United States and new 15000 cases/year in the world, what represents 1% of all approximately the cases of SSHL (1, 4).

Since it was described for the first time, 62 years ago (5), innumerable etiologies have been suggested: infectious (6), traumatic, neoplasia, immunological, ototoxic, vascular or ischemic, neurological, amongst others. Amongst the factors which are associated we have the metabolic disorders that are more frequent between cochlea –vestibulopath patients than in the general population (7). However, in only 10 to 15% of the cases the cause can be determined (1).

The treatment is controversial and can include, among others, antiinflammatory (steroids), vasodilatives, anti-viral, volumetric or hemodilutor, diuretic, antagonists of the calcium canals, hyperbaric chamber, carbogenic, anticoagulative expanders and recently corticosteroid intra-tympanic (8-13).

Approximately one third of the patients presents spontaneous improvement, most of them in first the two weeks of evolution of the disease (1, 3). The risk factors associated to a worse prognostic include: time of evolution (delay for beginning of the treatment), extremes of age, degree of the initial SSHL (severe losses), presence of initial associate vestibular symptoms, type of the tonal audiometric curve (descendent) (1, 4, 14).

The objective of this work is to observe the evolution of the SSHL in patients enclosed in different groups of treatment, according to the time of evolution of the illness and with the contraindications for use of medications.

METHOD

The design of the study configures one retrospect group of the handbook analysis of patients taken care in the Clinic of SS and enclosed in protocol of standardized treatment from January 1996 to January 2006. The effective ethical norms in the Institution, as determination of the Committee of Ethics and Research, have all been followed.

Our sample was constituted by 139 patients, with average age of 45.4 + 15.8 years (minimum = 13 years; maximum = 82 years), being 73 (52.5%) female and 66 (47.5%) of the male.

To be included in the study, the individual would have to present equal or bigger neuro-sensorial SSHL than 30 dB, in at least three consecutive frequencies, with sudden installation or in the maximum in 72 hours.

The patients were included in five groups of treatment, according to the time of beginning of the SSHL and the clinical conditions that qualified him/her, or not, to receive proposed medications. The following criteria have been used:

Group I: patients with installed history of SSHL for 5 days at most. They were submitted to the hospitalization and they received plasmatic expander (Dextran® 40000 UI, EV, every 12 h, up to 10 days), dexamethasone (8 mg, VO, once a day, 10 days, with project of gradual follow-up withdrawal), to acyclovir (200 mg, VO, 8/8 h, 15 days), acid nicotinic acid and hydrocloride of papaverine (30 mg and 100 mg, VO, every 12 h, 30 days), vitamin (50000 UI, VO, every 12 h, 30 days).

Group II: patients with history of SSHL installed from 0 to 15 days. Those who had clinical contraindication to the use of the plasmatic expander (systemic arterial hypertension, cardiopathy, coagulation disorder and renal insufficiency) were enclosed in this group. The therapeutical design of group I or III was prescribed, in the dependence of the time of beginning of the SSHL, however without the plasmatic expander.

Group III: patients with history of SSHL installed from 6 up to 15 days. It received the therapeutical design from group I, however without acyclovir.

Group IV: patients with history of SSHL installed from 16 up to 30 days. It received the therapeutical design from group I, however without plasmatic expander and acyclovir.

Group V: patients with installed history of SSHL for more than 30 days. It received nicotinic acid and papaverine hydrochloride (30 mg and 100 mg, VO, every 12 h, 30 days) and vitamin (50000 UI, VO, every 12 h, 30 days).

Audiometric serial examinations were carried through in all patients, before (initial) and after the beginning of the therapy (30, 90, 120 and 180 days).

The patients of groups I and III were hospitalized, as protocol above described, in order to receive plasmatic...
expander through endovenous way. In these cases, the tonal audiometry threshold was carried through on alternated days, the suspension of medication and the hospital discharge being programmed in the absence of improvement of the auditory thresholds after three days. The recovery of at least 10dB in 3 consecutive frequencies or of 15dB in two consecutive frequencies or 20dB in an isolated frequency in the tonal audiometry or still 15% in the speech recognition rate (SRR) were considered criteria of audiometric improvement. In the cases of audiometric improvement, the therapeutical design was kept by up to 10 days.

For documentation of the audiometric variation the PTA was used, pure tone average or pure tonal threshold.

The PTA for low pitch frequencies was determined by the average of the tonal thresholds of the frequencies of 250, 500 and 1000 Hertz (Hz) and the PTA for high pitch frequencies from the average of the 2000 frequencies, 4000 and 8000 Hz. The evolution or audiometric response of the patients among the several groups of treatment was determined by subtracting the initial PTA (first day of medical evaluation) from the end (after 180 days). For analysis statistics of the results, test t of Student and the analysis of variance were used (ANOVA). The considered level of significance was of \( p<0.05 \) in bicaudal tests.

**RESULTS**

In 129 (92.8%) patients the SSHL was unilateral and in only 10 (7.2%) it was bilateral. In terms of ethnics 77 (55.4%) were white, 35 (25.2%) were black and 4 (2.9%) were of the yellow race. In 23 (16.5%) cases this data were not computed in handbooks. The time of evolution of the SSHL until the first examination was in average 17.2 + 24.6 days (minimum = 0 day; maximum = 120 days).

The groups of treatment revealed similar in terms of the averages of the initial PTA, both for low pitch frequencies and for high pitch ones (Table 1). Considering all the assisted cases, a significant reduction in the average of the PTA was observed after the treatment when compared with the initial values, both for low pitch frequencies and for high pitch ones (Table 2).

With relation to the comparison it among the groups in terms of the evolution or audiometric response to the treatment, determined by the difference between final and initial PTA, both for low pitch frequencies and for high pitch ones, there was no statistical significance (Table 3).

We observe that a significant inverse linear correlation exists between the time passed until the first examination and the evolution or audiometric response (initial PTA - final PTA) of the patient to the treatment. Therefore, the bigger the time until the beginning of the treatment, the lesser the audiometric improvement. Thus, for the low pitch frequencies we observe a coefficient of correlation of Spearman \( r = -0.45 \) (\( p<0.001; IC 95\% = -0.58 \) to \( -0.29 \)) (Graph 1) and for high pitch frequencies \( r = -0.42 \) (\( p<0.001; IC 95\% = -0.55 \) to \( -0.25 \)) (Graph 2).

**Table 1.** Mean and standard deviation of initial PTA among the groups.

<table>
<thead>
<tr>
<th>Initial PTA (dB) (mean ± standard deviation)</th>
<th>G-I</th>
<th>G-II</th>
<th>G-III</th>
<th>G-IV</th>
<th>G-V</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grave</td>
<td>79.9 ± 30.5</td>
<td>76.1 ± 26.1</td>
<td>78.8 ± 28.3</td>
<td>75.9 ± 26.3</td>
<td>78.4 ± 27</td>
<td>0.945</td>
</tr>
<tr>
<td>Acute</td>
<td>88.8 ± 31.2</td>
<td>80 ± 33.5</td>
<td>79.1 ± 25.1</td>
<td>79 ± 36</td>
<td>86.9 ± 28</td>
<td>0.275</td>
</tr>
</tbody>
</table>

PTA: pure tone average ; dB: deciBell; G: treatment group; \( p \): significance level (ANOVA).

**Table 2.** Comparison between mean and standard deviation of initial and final PTA, considering the total sample.

<table>
<thead>
<tr>
<th>Frequencies</th>
<th>PTA (dB)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Initial mean ± standard deviation</td>
</tr>
<tr>
<td>Grave</td>
<td>77.4 ± 27.1</td>
</tr>
<tr>
<td>Acute</td>
<td>82.4 ± 30.3</td>
</tr>
</tbody>
</table>

PTA: pure tone average ; dB: deciBell; \( p \): significance level (ANOVA).

**Table 3.** Difference of final and initial PTA means among the groups.

<table>
<thead>
<tr>
<th>Initial PTA - final PTA (dB) (mean ± standard deviation)</th>
<th>G-I</th>
<th>G-II</th>
<th>G-III</th>
<th>G-IV</th>
<th>G-V</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accute</td>
<td>18.9 ± 20.6</td>
<td>29.6 ± 28.3</td>
<td>19.1 ± 19.9</td>
<td>11.8 ± 14.1</td>
<td>13 ± 18.6</td>
<td>0.069</td>
</tr>
<tr>
<td>Grave</td>
<td>25.7 ± 4.5</td>
<td>30.6 ± 31.2</td>
<td>23.1 ± 22.3</td>
<td>14.8 ± 14.9</td>
<td>20.4 ± 20.8</td>
<td>0.237</td>
</tr>
</tbody>
</table>

PTA: pure tone average ; dB: deciBell; G: treatment group; \( p \): significance level (ANOVA).
DISCUSSION

Literature shows that the average age of the patients with SSHL varies from 43 to 53 years, that the distribution for sex is equivalent and that the onset in more 95% of the cases is unilateral, similar to what we have observed in this survey (2, 4, 13, 15, 16). The only epidemiologist survey of SSHL published until the moment was made in Japan in three distinct years, in different decades and shows that the time between the beginning of the SSHL and the first examination varied from 11.6 ± 12.1 days in 1974, 9.1 ± 9.8 days in 1987 to 8.1 ± 9.1 days in 1993 (15). We cannot directly compare our casuistry with the one by Nakashima et al (15), for the type of design and being different populations, however we observe delay until the first examination what can be attributed in part to the indifference of the patients in relation to the symptom or the difficulty of access to the doctor.

Based on homogeneous groups in terms of the initial averages of the PTA and considering all the assisted cases, there was significant reduction in the average of the PTA. This average of improvement (initial PTA - final) was of 21.6dB for low pitch frequencies and 16.7dB for the high pitch. Although there was no control group for comparison, literature shows that the spontaneous improvement (without treatment) occurs in 32% of the SSHL cases and is in average of 15dB of PTA (3, 8), therefore lower than the one observed in this study. The estimated audiometric recovery with the treatment is of approximately 25dB of PTA (3), similar to the one that we observe for low pitch frequencies.

In terms of the evolution of the patients (initial PTA - PTA final) in the five groups of treatment, the observed difference was not significant, both for low pitch frequencies and for high pitch ones. Although we have not observed statistical significance, the improvement of the PTA was lower in the patients of groups IV and V, who initiated the treatment 15 days after the beginning of the SSHL. Groups I, II and III initiated the treatment in first the 15 days of SS, however the best PTA was observed in group II, the only one which did not receive plasmatic expander for clinical contraindication. Although some base illnesses are capable to intervene with the microcirculation of the internal ear making the clinical improvement of the SSHL difficult, the comorbidities presented for the patients of group II, did not apparently mean factors of worse prognostic. In our survey, the plasmatic expander did not contribute to potentiate the effect of the corticosteroid and of acyclovir in the treatment of the SSHL, once the group that better answered to the therapy did not use it.

We observe a significant inverse linear correlation between the passed time until the beginning of the treatment and the audiometric response presented by the patient, both for low pitch frequencies and for high pitch. We agree, therefore, with reports by other authors who show that the prognostic is better as much as precociously the treatment is started (1, 4, 17).

CONCLUSION

After our survey, we can infer that:

- The plasmatic expander did not improve the efficiency of the association between corticostroid and acyclovir for the SSHL treatment;
- The precocious beginning of the treatment determines the best disease evolution.

REFERENCES

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