Characteristics and Spontaneous Recovery of Tinnitus Related to Idiopathic Sudden Sensorineural Hearing Loss

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Objective: To evaluate the characteristics and spontaneous recovery of tinnitus related to idiopathic sudden sensorineural hearing loss (ISSNHL).

Study Design: Retrospective analysis from two randomized placebo-controlled clinical trials for treatment of ISSNHL within 48 hours from onset (Study A), or of tinnitus related to ISSNHL within 3 months from onset (Study B).

Setting: Forty-eight European sites (academic tertiary referral centers, private ENT practices).

Patients: One hundred thirteen adult patients of which 65 with hearing loss ≥30 dB (Study A) and 48 with persistent acute tinnitus (Study B) at baseline.

Interventions: Intratympanic (i.t.) injection of placebo gel in single dose or in triple dose during 3 consecutive days.

Main Outcome Measures: Frequency of tinnitus, subjective tinnitus loudness, rates of complete tinnitus remission, and complete hearing recovery during 3 months follow-up.

Results: In acute ISSNHL, tinnitus loudness decreased rapidly in cases of mild-moderate hearing loss, and tinnitus had completely resolved in two-thirds of patients after 3 months. Hearing recovery preceded tinnitus resolution. When associated with severe-profound hearing loss, tinnitus improved significantly less. Complete hearing recovery and full tinnitus remission were both about three times more frequent in mild-moderate hearing loss patients than in severe-profound cases. Improvement in tinnitus loudness over time can be approximated by a negative exponential function.

Conclusions: Prognosis for ISSNHL-related tinnitus is relatively poor in case of severe-profound hearing loss and the longer it has persisted. Alleviation or management of tinnitus should be a key therapeutic objective especially in pronounced ISSNHL cases. Key Words: Hearing recovery—Intratympanic—ISSNHL—Spontaneous recovery—Tinnitus. Otol Neurotol 37:634–641, 2016.
initial ISSNHL severity and correlate with hearing recovery rates. Studies enrolling ISSNHL patients with mild to moderate hearing loss already show after 7 to 10 days not only substantial hearing recovery, but also a large decrease in tinnitus incidence from 50 to 70% to 30 to 40% (2,10). On the other extreme, studies enrolling ISSNHL patients with more severe hearing loss show only small to moderate hearing recovery and small to moderate reductions in tinnitus incidence, if at all (11,12); for example, 12 months after onset tinnitus was still present in 60% of patients with severe ISSNHL (11). Recovery in hearing in general seems to occur more rapidly than a decrease in tinnitus loudness or tinnitus remission (13).

Given the scarcity of information on ISSNHL-tinnitus, we sought to obtain more detailed data and gain further insights from two recent prospective double-blind, randomized, placebo-controlled studies that involved ISSNHL patients. One of the studies enrolled ISSNHL patients within 48 hours from onset and followed them for 3 months, with a primary focus on hearing loss (14). The other study enrolled ISSNHL-tinnitus patients within 3 months from onset and followed them for 3 months, with a primary focus on tinnitus (15). Thanks to the inclusion of placebo control groups, the two studies allow for a detailed analysis of the acute and post-acute time course and spontaneous recovery of ISSNHL-related tinnitus.

MATERIALS AND METHODS

Study Design and Participants

For the purposes of the present study we retrospectively analyzed data from a total of 113 placebo-treated ISSNHL patients participating in one of the two randomized double-blind clinical trials AM-111-CL-08-01 (“Study A”; 65 patients) or AM-101-CL-08-01 (“Study B”; 48 patients). Details of the study design and procedures have been described elsewhere (14,15).

Study A enrolled patients (18–60 years of age) suffering from unilateral ISSNHL with hearing loss of at least 30 dB at the average of the three worst affected contiguous audiometric test frequencies (pure tone average [PTA]), determined against the contralateral ear (or a pre-existing audiogram or ISO norm values in case of history of asymmetric hearing loss). They were treated within 48 hours from onset. The presence of tinnitus was not a requirement for study participation. Study B enrolled patients (18–65 years of age) with persistent uni- or bilateral tinnitus following ISSNHL within 3 months from onset. Persistent tinnitus was considered given when patients indicated that they could always hear their tinnitus when they were thinking of it in the past 2 weeks and that their tinnitus during that period occurred either all of the time, most of the time, or a good bit of the time. The presence of hearing loss was not a requirement for study participation.

Randomization and Masking

At baseline (Day 0), study participants were randomized to receive the investigational drug, AM-111 (Study A) or AM-101 (Study B), respectively, or placebo (a viscous phosphate-buffered gel formulation based on 0.7% sodium hyaluronate) at a 2:1 ratio. The study drug formulation was identical in appearance for active and placebo doses and revealed no differences during administration. Study patients and investigators remained blinded throughout the entire study.

Procedures

Participants in Study A received on Day 0 a single dose intratympanic (i.t.) injection, whereas participants in Study B received 3 i.t. injections on Days 0, 1, and 2 under local anesthesia through a small myringotomy with the patient’s head placed in a position tilted 45 degrees towards the untreated ear. In Study B, only the worse affected ear was treated if bilaterally affected (<14% of patients). Patients remained in their supine position for approximately 30 minutes. Patients were followed for 90 days; interim follow-up visits were scheduled at Days 3 (only Study A), 7 and 30. Tinnitus loudness was recorded on a 0 to 10 (Study A) respectively, 0 to 100 (Study B) point numerical rating scale with a recall period of “right now” (0 = “no tinnitus” and 10 = “extremely loud” [Study A], respectively “very loud” [Study B]). For comparison with Study A, values from Study B were converted to a 0 to 10 scale. Hearing loss in Study A was classified by baseline audiogram type (16), initial frequency range (low, medium, or high), and baseline severity (mild to moderate: PTA < 60 dB, severe to profound: PTA ≥ 60 dB; [17]). Complete tinnitus remission was assumed when tinnitus loudness was rated as 0. Hearing was considered as fully recovered if PTA recovered to within 10 dB of the baseline value. In Study A, study participants with insufficient hearing recovery (defined as <10 dB from baseline to Day 7 at the average of the three worst affected contiguous audiometric test frequencies) were given the option to receive oral prednisolone 50 mg b.i.d. for 5 days as a rescue therapy.

Statistical Methods

Statistical analyses were performed with SAS (SAS Institute, Cary, NC, version 9.3). Tests and graphs were based on analysis sets that included all placebo-treated patients for whom data were available both at baseline and Day 90. Mean values of tinnitus loudness at treatment visits and change from baseline were compared using the t test, assuming equal variances for the compared subgroups. A non-linear time function of tinnitus loudness was estimated using the SAS “proc nlm” function. The frequency of complete tinnitus remission and complete hearing recovery was compared between subgroups using Fish-er’s exact test. For pairwise correlation of PTA and tinnitus loudness at baseline and Day 90, Spearman’s correlation coefficient r_s was calculated. A contingency table with complete hearing recovery and complete tinnitus remission (yes/no) was generated to evaluate their interrelations within subjects; in addition a χ² test for independence was performed.

RESULTS

Tinnitus in the Acute ISSNHL Patient

Out of a total of 65 ISSNHL patients enrolled within 48 hours from acute hearing loss onset, 72.3% reported the concurrent onset of tinnitus. 7.7% of patients had pre-existing tinnitus and 20.0% reported no tinnitus at all. The incidence of novel tinnitus (47 patients in total) was 75.9% in cases of severe-profound hearing loss (n = 22) and 70.3% in mild-moderate cases (n = 25; Table 1). In the subgroup of patients with novel-onset tinnitus, cases
with severe-profound hearing loss showed a higher proportion of predominantly high-frequency hearing loss and of audiogram types C or E (descending slope or total or subtotal anacusis) than those with mild-moderate hearing loss. Mean subjective tinnitus loudness in the 47 patients with novel-onset tinnitus was 5.7 points on the 0 to 10 point numerical rating scale; it was significantly lower in the mild-moderate subgroup (5.2 points) than in the severe-profound subgroup (6.4 points; \( p < 0.05 \)).

Seventeen out of the 47 patients with fresh-onset tinnitus received prednisolone reserve therapy starting after the Day 7 visit because of unsatisfactory hearing recovery up to that point; 14 of them suffered from severe-profound hearing loss at baseline. Since active treatment might conflict with our aim of determining spontaneous recovery after Day 7, we compared the change in mean PTA and tinnitus loudness. While those patients who went on to receive reserve therapy on Day 7 had initially improved less than those who did not receive steroids—hence the former’s eligibility for it—the changes thereafter were essentially in parallel. The differences in mean changes in tinnitus loudness and PTA from Day 7 to Day 30 and from Day 7 to Day 90 were statistically not significant (\( p = 0.62 \) and \( p = 0.29 \) and \( p = 0.54 \) and \( p = 0.06 \), respectively). As the reserve therapy could not influence data points up to Day 7 and had no significant impact thereafter and to avoid selection bias, we performed all analyses in the full data set, that is, including steroid treated patients.

Overall, tinnitus loudness dropped rapidly as tinnitus in many cases resolved completely or at least became less intense. Indeed at the follow-up visit on Day 7 mean tinnitus loudness had already more than halved to 2.7 points, and 15.6% of patients experienced complete tinnitus remission. Additional substantial improvement was observed to Day 30, as mean tinnitus loudness decreased further to 2.0 points, and a cumulated 35.6% of patients had complete tinnitus remission. Between Days 30 and 90 the rate of incremental improvement decreased substantially; at the last follow-up visit, the mean tinnitus loudness was down to 1.3 points, and the complete tinnitus remission rate was 44.4%.

The level of tinnitus loudness and rate of complete tinnitus remission differed substantially between the

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**TABLE 1. Baseline characteristics of acute idiopathic sudden sensorineural hearing loss patients**

| Hearing Loss | Mild to Moderate | Severe to Profound | Total |
|--------------|------------------|--------------------|-------|
| Age, years   | 39.6 (11.3)      | 46.6 (10.9)        | 42.8 (11.6) |
| Range        | 19–60            | 19–61              | 19–61 |
| Time from onset, hours | 25.9 (11.8) | 31.9 (12.1) | 28.6 (12.3) |
| Range        | 3–47             | 6–48               | 3–48 |
| Presence of tinnitus, number (%) patients | 36 (100.0) | 29 (100.0) | 65 (100.0) |
| Tinnitus present, concurrent onset | 25 (70.3) | 22 (75.9) | 47 (72.3) |
| Tinnitus pre-existing | 4 (13.5) | 1 (3.4) | 5 (7.7) |
| No tinnitus present | 7 (16.2) | 6 (20.7) | 13 (20.0) |
| Novel-onset tinnitus only: | | | |
| Tinnitus loudness (0–10) | Mean (SD) | 5.2 (1.7) | 6.4 (1.9) | 5.7 (1.9) |
| Range        | 2–10             | 3–10               | 2–10 |
| Initial frequency range, number (%) patients | 25 (100.0) | 22 (100.0) | 47 (100.0) |
| Low frequency hearing loss | 11 (44.0) | 5 (22.7) | 16 (34.0) |
| Medium frequency hearing loss | 5 (20.0) | 4 (18.2) | 9 (19.1) |
| High frequency hearing loss | 9 (36.0) | 13 (59.1) | 22 (46.8) |
| Audiogram type, number (%) patients | 25 (100.0) | 22 (100.0) | 47 (100.0) |
| Type A (ascending) | 4 (16.0) | 0 (0.0) | 4 (8.5) |
| Type B (flat) | 5 (20.0) | 3 (13.6) | 8 (17.0) |
| Type C (descending) | 9 (36.0) | 8 (36.4) | 17 (36.2) |
| Type D (U or V shaped) | 6 (24.0) | 1 (4.5) | 7 (14.9) |
| Type E (total or subtotal anacusis) | 0 (0.0) | 8 (36.4) | 8 (17.0) |
| Not assignable | 1 (4.0) | 2 (9.1) | 3 (6.4) |
| Hearing threshold (pure tone average), dB | Mean (SD) | 47.7 (5.7) | 85.7 (21.0) | 65.5 (24.2) |
| Range | 31.7–58.3 | 60.0–120.0 | 31.7–120.0 |
| Hearing loss, dB | Mean (SD) | 36.1 (6.2) | 72.0 (22.4) | 52.9 (24.0) |
| Range | 20.0–48.3 | 41.0–116.7 | 20.0–116.7 |

PTA indicates pure tone average (mean hearing threshold at three most affected contiguous audiometric test frequencies); SD, standard deviation. Placebo-treated patients in clinical trial AM-111-CL-08-01 (Study A; \( n = 65 \)).

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subgroups of mild-moderate and severe-profound acute hearing loss (Fig. 1). Although the incidence of fresh-onset tinnitus was not much lower at baseline in the mild-moderate subgroup, tinnitus loudness was significantly lower not only at baseline, but even more so at all follow-up visits ($p < 0.01$); mean loudness decreased from 5.3 to 0.7 points at Day 90 (mean decrease 87.6%) versus a decrease from 6.4 to 2.0 points (mean decrease 69.3%) in the severe-profound subgroup. Initially, tinnitus loudness dropped much more quickly than in the severe-profound subgroup; the difference in absolute tinnitus loudness reduction was statistically significant for the change from baseline to Day 7 ($p < 0.05$). Complete tinnitus remission in the mild-moderate subgroup set in much earlier and benefited a significantly higher share of patients (Fig. 2; Table 2). 30.4% of patients in the mild-moderate hearing loss subgroup experienced already at Day 7 complete tinnitus remission, whereas in the severe-profound hearing loss subgroup such complete resolution was observed for the first time only by Day 30. By Day 90, the incidence of complete tinnitus remission was almost three times higher in the mild-moderate subgroup than in the severe-profound subgroup (65.2 versus 22.7%; $p < 0.01$).

The time course of the mean tinnitus loudness was modeled as a negative exponential function

$$f(t) = b_0 + b_1 \cdot \exp(-b_2 \times t)$$

whereas parameters for the hearing loss severity subgroups differed substantially (Fig. 1). For both subgroups, the function appeared to provide a good fit for early time points, but underestimated the tinnitus level at Day 30 and overestimated it at Day 90. While the actual curve suggests the potential for further, albeit only small improvement beyond Day 90, the function yields no or hardly any additional reduction in tinnitus loudness beyond Day 30. Patterns for complete hearing recovery were similar to those observed for complete tinnitus remissions in both hearing loss severity subgroups (Table 2). The difference was statistically significant for complete hearing recovery at all follow-up visits ($p < 0.001$) and for complete tinnitus remission beginning at Day 7 ($p < 0.01$). Whereas 73.9% of patients in the mild-moderate hearing loss subgroup enjoyed complete hearing recovery already at Day 7, this was the case for only 4.5% of patients in the severe-profound hearing loss subgroup. At Day 90, the incidence of complete hearing recovery was 3.7 times higher in the mild-moderate hearing loss subgroup (87.0 versus 23.8%; $p < 0.001$).

The level of tinnitus loudness at Day 90 was statistically significantly correlated with the level of PTA both at baseline ($r_s = 0.47, p = 0.001$) and at Day 90 ($r_s = 0.48, p = 0.001$); the correlation between the PTA at baseline and Day 90, however, was even stronger ($r_s = 0.82; p < 0.001$). A $\chi^2$ test showed that occurrence of complete

*FIG. 1.* Time course of subjective tinnitus loudness from baseline to Day 90 rated on a 0 to 10 scale by placebo-treated patients suffering from idiopathic sudden sensorineural hearing loss and concurrent onset of tinnitus within the past 48 hours in clinical trial AM-111-CL-08-01 (Study A). A. Subgroup of patients with mild to moderate hearing loss at baseline ($n = 23$); B. subgroup of patients with severe to profound hearing loss ($n = 22$) at baseline. Shown are actual mean values as well as estimated mean values based on a negative exponential function: $f(t) = 0.8737 + 4.4327 \times \exp(-0.2477 \times t)$ for mild-moderate cases and $f(t) = 2.3253 + 3.9713 \times \exp(-0.1144 \times t)$ for severe-profound cases.

| Subgroup | Baseline | Day 3 | Day 7 | Day 30 | Day 90 |
|----------|----------|-------|-------|--------|--------|
| Tinnitus remission | | | | | |
| Mild-moderate ($n = 23$) | 0 (0.0%) | 2 (8.7%) | 7 (30.4%) | 13 (56.5%) | 15 (65.2%) |
| Severe-profound ($n = 22$) | 0 (0.0%) | 0 (0.0%) | 0 (0.0%) | 3 (13.6%) | 5 (22.7%) |
| $p$ value | 0.489 | 0.009 | 0.005 | 0.007 | |
| Hearing recovery | | | | | |
| Mild-moderate ($n = 23$) | 0 (0.0%) | 10 (43.5%) | 17 (73.9%) | 19 (82.6%) | 20 (87.0%) |
| Severe-profound ($n = 22$; 21 at Day 90) | 0 (0.0%) | 0 (0.0%) | 1 (4.5%) | 3 (13.6%) | 5 (23.8%) |
| $p$ value | <0.001 | <0.001 | <0.001 | <0.001 | <0.001 |

Acute idiopathic sudden sensorineural hearing loss patients reporting onset of tinnitus concurrently with onset of acute hearing loss who were treated with placebo in clinical trial AM-111-CL-08-01 and had a baseline tinnitus loudness rating (Study A; $n = 45$).

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hearing recovery and complete tinnitus remission were not independent of each other (\( p < 0.01 \)); Cramer’s \( V \) was 0.427 and thus confirmed the association between these two discrete variables. 36.4% of patients experienced both complete hearing recovery and complete tinnitus remission, and 34.1% neither of the two; complete tinnitus remission without complete hearing recovery was the least frequent combination (Table 3).

Complete tinnitus remission lagged complete hearing recovery in both subgroups (Fig. 2, A and B). In the mild-moderate hearing loss subgroup the majority of the observed hearing improvement was achieved already at Day 7, whereas the majority of complete tinnitus remission occurred only between Day 7 and Day 30. In the severe-profound hearing loss subgroup, the frequency of complete tinnitus remission and hearing recovery moved essentially in parallel at relatively low levels.

**Tinnitus Course after Initial Stage**

For the post-acute ISSNHL stage, we focused on those tinnitus patients who had enrolled in Study B between 30 and 90 days from onset—Study A had showed only little further tinnitus reduction beyond the first month. Here, mean tinnitus loudness decreased over the following 90 days from 5.3 to 3.1 points (corresponding to approximately 5 months from tinnitus onset); 17.1% of patients experienced complete tinnitus remission (Table 4). Post-acute stage improvement was smaller than in the acute ISSNHL stage and its magnitude even below that for acute severe-profound hearing loss patients. The rate of tinnitus improvement was again time-dependent: tinnitus loudness dropped more, and spontaneous tinnitus remission occurred more frequently among the patients enrolling in the second month from onset compared with those presenting only in the third month (Fig. 3). The two subgroups showed a similar reduction in the first 7 days from baseline. After 90 days follow-up (corresponding to about 4.5 and 5.5 months from onset), tinnitus loudness dropped to 2.3 points in the earlier and 3.9 points in the later subgroup (Table 4).

**DISCUSSION**

ISSNHL has been the object of a growing number of clinical research studies in otology, yet there is still only limited information available on the natural history. This observation relates to hearing loss and hearing recovery,

### Table 3. Interrelation between complete tinnitus remission and complete hearing recovery

|                | Complete Tinnitus Remission |
|----------------|----------------------------|
|                | No  | Yes | Total |
| Complete hearing recovery |     |     |       |
| No              | 15  | 4   | 19    |
| % overall       | 34.1% | 9.1% | 43.2% |
| % row           | 78.9% | 21.1% | 100.0% |
| Yes             | 9   | 16  | 25    |
| % overall       | 20.4% | 36.4% | 56.8% |
| % row           | 36.0% | 64.0% | 100.0% |
| Total           | 24  | 20  | 44    |
| % overall       | 54.5% | 45.5% | 100.0% |

Acute idiopathic sudden sensorineural hearing loss patients reporting onset of tinnitus concurrently with onset of acute hearing loss who were treated with placebo in clinical trial AM-111-CL-08-01 and had a tinnitus loudness rating both at baseline and Day 90 (Study A).
and even more so to its most frequent comorbidity, tinnitus. Frequently, for hearing recovery reference is made to a ‘‘one third rule’’ (about one-third of patients recover completely, one-third recover partially, while one-third show a remaining hearing loss). This rule draws upon findings from some of the few studies that have assessed spontaneous recovery in non- or placebo-treated patients (18,19). However, these data have to be interpreted and compared across studies with caution given important differences in outcome measures (e.g., tested audiometric frequencies; [20]) or patient characteristics (e.g., time to diagnosis, affected hearing frequencies).

Similar reservations have to be made with regards to the scarce tinnitus data that are available from ISSNHL studies. To our knowledge, the present study represents the first attempt to assess in detail the characteristics of ISSNHL-tinnitus and its spontaneous recovery based on data from well-defined patient populations.

The outcomes from the present study confirm the high incidence of tinnitus that accompanies ISSNHL at the time of onset that has been reported previously. They further show how closely related the two symptoms hearing loss and tinnitus are in terms of severity and recovery patterns. At baseline, tinnitus was not much more frequent in patients with severe-profound acute hearing loss than in those with mild-moderate hearing loss, but its subjective loudness was significantly higher. Over the following 3 months, there was no significant difference between the subgroups in absolute reduction in tinnitus loudness; however, mild-moderate patients enjoyed a significantly higher rate of complete remission. In that, tinnitus recovery followed a similar pattern as in hearing recovery—mild-moderate acute hearing loss recovered more rapidly and to a larger extent than severe-profound hearing loss. Ninety days after onset, complete hearing recovery and complete tinnitus remission were both about three times more frequent in the mild-moderate acute hearing loss subgroup than in the severe-profound hearing loss subgroup.

In spite of important recovery in the average ISSNHL patient, 90 days following the originating incident, there were still 34.8 and 77.3% of patients in the mild-moderate and severe-profound hearing loss subgroups, respectively, who had tinnitus. Data that was available from the second trial from patients enrolling 30 to 90 days after the tinnitus triggering event showed that improvement in tinnitus is still possible beyond the first 90 days, albeit at a lower rate and with complete remission rates of <20%. Whether these patients initially had severe-profound hearing loss at the time of onset or—regardless of hearing loss severity—simply did not improve at all or only slightly is not known. The negative exponential function determined in Study A for severe-profound hearing loss cases would suggest lower tinnitus loudness values than those actually observed post-acutely in Stratum B; accordingly, it appears most likely that both patterns were represented.

With regards to the long-term impact of ISSNHL-tinnitus, a retrospective Swedish study showed that annoying tinnitus (defined as being present always or often) together with remaining vertigo was the strongest predictor of negative impact on the QoL of ISSNHL-patients (7). The authors found significant

### Table 4. Tinnitus evolution in ISSNHL patients presenting within 2 to 3 months from onset

| Days to Enrollment | Total | 30–59 | 60–90 |
|--------------------|-------|-------|-------|
| N                  | 41    | 20    | 21    |
| Mean tinnitus loudness (SD) |       |       |       |
| Baseline           | 5.3 (2.1) | 5.2 (2.2) | 5.4 (2.1) |
| Day 90             | 3.1 (2.6) | 2.3 (1.6) | 3.9 (3.0) |
| Mean change to Day 90 Points (SD) |       |       |       |
| % of baseline      | −44.0% | −57.4% | −31.3% |
| % patients with complete tinnitus remission | 17.1% | 20.0% | 14.3% |

SD indicates standard deviation. Idiopathic sudden sensorineural hearing loss patients with tinnitus who were treated 30 to 90 days from onset with placebo in clinical trial AM-111-CL-08-01 (Study B).
correlations with the QoL measures Hospital Anxiety and Depression Scale, EuroQol-5D, and a problems impact rating scale; in addition, ISSNHL-patients suffering from annoying tinnitus were significantly more often on sick leave directly after onset or in the years thereafter. 22 to 24% of patients in their studied ISSNHL population, which included also cases of less than severe hearing loss, reported the long-term presence of annoying tinnitus.

The present study confirmed earlier findings (13) wherein the pace of tinnitus improvement lags hearing recovery. Active treatment with AM-111 in Study A had also showed earlier effects on hearing recovery than on complete tinnitus remission in patients with severe-profound hearing loss (14). While both hearing loss and tinnitus are symptoms of a dysfunction within the auditory system, there seems to be a difference between the capacity for recovery of afferent input for hearing and for resolution/suppression of the tinnitus perception. Presuming that tinnitus is the result of maladaptive plasticity within the auditory system (for a review see [21–24]), reversing such plasticity changes may take longer than intrinsic repair mechanisms inside the cochlea and may actually depend to some extent on them. In case of insufficient hearing recovery, auditory rehabilitation seems to be warranted to manage or alleviate the negative impact of tinnitus on the well-being and QoL of ISSNHL-tinnitus patients (6,7).

In conclusion, ISSNHL-related tinnitus overall has a good prognosis for rapid improvement up to complete remission in patients suffering from mild-moderate acute hearing loss at onset, whereas recovery is led by improvement in hearing. Watchful monitoring and waiting appear to be appropriate management options, as recommended e.g., in the United States treatment guideline (4). In severe-profound hearing loss patients, both hearing recovery and tinnitus remission are much rarer and prognosis is much poorer. In the absence of effective treatments for the acute hearing loss and/or tinnitus, patient education and auditory rehabilitation should be performed proactively. Tinnitus should be recognized as a patient burden that in case of pronounced ISSNHL may be even more important and bothersome than persisting hearing loss.

Although all patients in the present study received active gel injections, which may have evoked some type of placebo response, outcomes can be considered to depict the natural history of tinnitus rather closely. Importantly, in Study A the primary focus of patients and investigators was on hearing loss rather than tinnitus with both sensations being very recent, which may have mitigated any placebo effect on the tinnitus. That some patients took a course of oral corticosteroids as active treatment 7 days after enrollment does not invalidate their inclusion as the reserve therapy did not have any apparent effect on the course of tinnitus, which is in line with other studies’ outcomes (25,26). In Study B the primary focus was on tinnitus, and there was a reduction of tinnitus loudness of 16 to 18% within the first week of trial participation that seemed atypical of spontaneous recovery at this stage; however, any placebo response may have been mitigated by previous unsuccessful treatment attempts.

Given the limited size of the database for the present study, further analyses in a larger number of patients seem to be warranted. It would especially be worthwhile to compare ISSNHL cases with and without tinnitus as comorbidity; however, ISSNHL without tinnitus represents a clear minority. In addition, given the acute nature of the condition, it may be challenging or impossible to obtain pure observational or placebo data at all.

Acknowledgments: The authors wish to thank Drs. Manfred Wargenau and Frauke Friedrichs for their biostatistical contributions, Prof. Marlies Knipper for her critical review of the manuscript, and the two anonymous reviewers for their thoughtful suggestions.

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