Cochlear Cell Death Induced Via Cisplatin or Gentamycin in Combination with Furosemide in Rodents

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Abstract

Otototoxic models of animal represent an elemental tool in basic otological research. In the present study, using guinea pigs we compared cochlear lesions mediated via cisplatin applied in terms of two regimens: consecutive application alone and in combination with furosemide. The influences of furosemide alone were also assessed; it was observed to result in temporary hearing loss and reversible damage to the stria vascularis. Consecutive administration of cisplatin alone tended to be disadvantageous because it bred progressive body weight loss and higher mortality compared with the combined regimen, which utilized a smaller cisplatin dose. The combined regimen brought about remarkable hair cell loss without corresponding lesion of spiral ganglion neurons. This difference suggests that the co-administered regimen did not mimic the damage to cochlear neuronal innervation caused by clinical application of cisplatin. In the meantime, we spread this method to the ototoxic model build in mice. Co-administration of gentamicin and furosemide caused marked hair cell loss as well as less mortality compared with consecutive application of gentamicin in mice. Due to different pharmacokinetics between cisplatin in guinea pigs and gentamicin in mice, the injection regimens of co-administration of furosemide - cisplatin in guinea pigs and gentamicin – furosemide in mice were varied. Overall, the methods of co-administration of cisplatin/gentamicin and furosemide in rodents facilitate ototoxic model build.

Keywords: Cisplatin; Gentamicin; Furosemide; Auditory brain-stem response; Hair cell; Spiral ganglion neuron

Introduction

Cisplatin (Cis) is a first-generation antineoplastic drug that is still frequently used in the clinic. Major side effects of this drug include ototoxicity, neurotoxicity and nephrotoxicity [1-3]. Cis has been demonstrated to injure cells and tissues by inducing apoptosis [4,5]. Great efforts have been made to clarify the mechanisms of Cis ototoxicity, but exactly how Cis damages cochlear tissue remains unclear. Specifically, why terminally differentiated cochlear cells (cells that have exited the cell cycle) are killed by Cis, which is designed to kill mitotic cells, remains unknown, and why hair cells are more sensitive to Cis than other cell types is also unclear. Thus, further studies are needed to explore these issues, and such studies will likely be done in animal models, because of ethical considerations.

Rodents are commonly used in such studies due to their availability, low cost and simpler genetic backgrounds. However, a major disadvantage of rodents is the high resistance of their cochlea to Cis damage. Several methods have been used to establish Cis ototoxicity in these animals, and the administration regimen is of great importance in efforts to effectively induce model cochlear lesions. A single high-dose Cis injection produces little hair cell damage, even at a dose that causes 50% mortality [6]. Consecutive injections over several days have been used in many reports as a way to induce cochlear lesions [7,8]. However, this regimen may also result in rapid deterioration in the animals’ health and high mortality before any evident ototoxic effect is established [9-11].

To reduce mortality following Cis administration when establishing cochlear lesions, it is desirable to use an agent that can selectively enhance the effect of Cis in the inner ear [12]. Loop diuretics, including Ethacrynic Acid (EA) and furosemide, have synergistic interactions with Cis in the cochlea [13-15]. These drugs cause temporary hearing loss, which manifests as limited, reversible threshold shifts up when administered alone [16,17]. However, even a small amount of Cis can cause a significant hair cells lesion if it is combined with EA or furosemide [14,18].

Previously, EA was often the loop diuretic of choice for this purpose [14,18]. However, this drug was recently discontinued, because the FDA withdrew its approval for use in humans. Additionally, EA must be administered intravenously, which is difficult in some animal species, such as guinea pigs. Furosemide is a diuretic that can replace EA in establishing ototoxicity in animal models. However, furosemide may not produce exactly the same outcome as EA. Thus, a comprehensive evaluation is necessary to validate the use of furosemide. There was no such report until recently, when a study in mice was published [19]. While the mouse study is important due to the obvious advantages of mouse models in hearing research, this evaluation should be duplicated in other species, such as guinea pigs, that are commonly used in ototoxicity studies. This species is favored by many researchers due to ease of handling and their relatively large cochlea, facilitating morphological evaluations.

While the co-administration of Cis with a diuretic drug facilitates the establishment of Cis-induced cochlear lesions, the possibility exists that the cochlear damage induced via different approaches may differ. In previous studies, even with the use of EA, such potential differences were not fully addressed.

In the present study, using guinea pigs we compared Cis-induced...
The percentage value was calculated as the result of lost OHCs divided by the total OHCs in the corresponding region. Furosemide was given at 200 mg/kg for all subgroups. n = sample size (survived animals). Interval is the time delay between furosemide and Cis administration.

Table 1: Percentage OHC loss by group and anatomical location in guinea pigs.

<table>
<thead>
<tr>
<th>Group</th>
<th>Cis dose</th>
<th>Interval</th>
<th>n</th>
<th>1st turn</th>
<th>2nd turn</th>
<th>3rd turn</th>
<th>4th turn</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 mg/kg × 5 d</td>
<td>N/A</td>
<td>11 (7)</td>
<td>35.4 ± 4%</td>
<td>11.8 ± 1.5%</td>
<td>2.9 ± 1.3%</td>
<td>1.9 ± 0.1%</td>
<td>18 ± 2%</td>
<td></td>
</tr>
<tr>
<td>0.1 mg/kg</td>
<td>30 min</td>
<td>7 (6)</td>
<td>18.9 ± 1%</td>
<td>10.9 ± 1%</td>
<td>3.7 ± 1%</td>
<td>1.1%</td>
<td>11.6 ± 3%</td>
<td></td>
</tr>
<tr>
<td>0.2 mg/kg</td>
<td>0 min</td>
<td>5 (4)</td>
<td>2%</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>30 min</td>
<td>7 (6)</td>
<td>90.8 ± 5%</td>
<td>28.2 ± 4%</td>
<td>9.2 ± 1%</td>
<td>3.9 ± 1%</td>
<td>45.6 ± 2.4%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>60 min</td>
<td>6 (6)</td>
<td>89.9 ± 6%</td>
<td>34.1 ± 4%</td>
<td>8.6 ± 1%</td>
<td>2.8%</td>
<td>48.6 ± 3%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>90 min</td>
<td>8 (6)</td>
<td>98.2 ± 2%</td>
<td>45.5 ± 7%</td>
<td>17.4 ± 3%</td>
<td>5.3%</td>
<td>56.2 ± 1%</td>
<td></td>
</tr>
<tr>
<td>2.0 mg/kg</td>
<td>30 min</td>
<td>8 (6)</td>
<td>100%</td>
<td>98.7 ± 1.2%</td>
<td>78.8 ± 3.6%</td>
<td>97.7 ± 2.8%</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Materials and Methods

Subjects and groupings

In total, 73 Dunkin-Hartley guinea pigs, with body weights from 350 to 450 g, were obtained from the Experimental Animal Service of Shanghai Jiaotong University Medical School. This study was conducted in strict accordance with Declaration of Helsinki principles and the protocol was approved by the Committee on Experimental Animal Services, Shanghai, China (permit number: SYXX (H2011-0128)).

The animals, which were maintained on a 12/12-h light-dark cycle during the experiment, were randomly assigned to four groups: group 1 (n=15) received furosemide alone (fur-alone group), group 2 (n=11) received consecutive injections of Cis alone (Cis-alone group) and group 3 (n=41) received a single injection of Cis in combination with furosemide (co-administration group); group 4 (n=6) were normal controls. The animals in group 3 were further divided into subgroups according to the Cis dose and the time interval between the administration of Cis and furosemide (Table 1). Hearing status was monitored using a frequency-specific Auditory Brain-Stem Response (ABR) test before and at 0.5 hours, 1 hour, 2 hours, 4 hours and 12 hours after drug administration in group 1 and at 7 days post drug application in group 2, 3 and 4. Immediately after the final functional test, the animals were sacrificed and the cochleae were harvested for morphological observation. In each animal, one ear was used to generate a stretched preparation of the basilar membrane to observe the lesion in the organ of Corti, and cross sections were made of the stria vascularis.

As for the ototoxic effect of co-administration of furosemide and gentamycin in mice, seventeen C57 BL/6J mice were recruited, and gentamicin in mice, seventeen C57 BL/6J mice were recruited, which were divided into 2 groups. Group 1 (n=9), a single injection of gentamycin (40 mg/kg) in combination with furosemide (400 mg/kg), and group 2 (n=8), a single injection of gentamycin (60 mg/kg) in combination with furosemide (400 mg/kg). The subjects received hearing function tests before and 7 days post the ototoxic drugs treatment. After the final ABR test, the animals were sacrificed and the cochleae were harvested for morphological observation, only by surface preparation in mice.

Drug administration

For guinea pigs, consecutive Cis administration in the Cis-alone group (group 2) was conducted while the animals were awake. Cis (P4394; Sigma, St. Louis, MO, USA) was prepared in a 0.9% saline solution. A total dose of 15 mg/kg (i.m.) was given to each animal, divided evenly across 5 consecutive days. To administer furosemide (200 mg/kg i.m., cat. #5913020911; Webster Veterinary, Devens, MA, USA) in groups 1 and 3, the animals were anesthetized with 8 mg/kg ketamine (i.p.) followed by 33 mg/kg pentobarbital sodium (P3761; Sigma, St. Louis, MO, USA; i.p.) before the furosemide injection. This regimen kept the animals in deep anesthesia for 2.5-3.5 h. The animals were laid on a thermostatic heating pad to maintain the body temperature at 37°C until they had recovered fully from the anesthesia. In group 3, different Cis doses were given (i.m.) at different time intervals after furosemide administration (Table 1).

For mice, gentamicin (G1914; Sigma-Aldrich; St. Louis, MO, USA) was injected via i.H. to each animal, 40 minutes later, furosemide was applied. The animals were not under anesthesia when ototoxic drugs treated.
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**Stretched preparation**

Animals were sacrificed with a pentobarbital sodium overdose (200 mg/kg, i.p.). In each guinea pig, one ear was randomly selected for a stretched preparation, while the other was used for semi-thin cross sections. In mice, all of the cochleae were used for stretched preparation. The temporal bones were harvested and each bulla was opened to expose the cochlea. The oval window and round window were opened. A small hole was made at the apex of the cochlea, through which fixative (4% paraformaldehyde in 0.1 M PBS) was perfused. Then, the cochleae were submerged in the fixative for at least 4 h. The samples of mice were decalcified in 10% EDTA on a swing bed for 7 days at room temperature before dissection of cochlea. The basilar membrane was dissected out under a dissecting microscope and the tectorial membrane was then removed. To stain the organ of Corti with F-actin, the preparation was treated with a rhodamine phalloidin solution (P1951; Sigma) in the dark for 20 min at room temperature. The specimens were then rinsed three times with 0.1 M PBS (pH 7.4). Fluorescent hair cells were counted under an epifluorescence light microscope. The percentage hair cell loss in each 0.24 mm segment was calculated, and a cochleogram was made for hair cell loss as a function of cochlear length.

**Semi-thin cross sections**

Cochleae were fixed in 2.5% glutaric dialdehyde for 4 h at room temperature. Following several rinses with 0.1 M sodium cacodylate buffer (pH 7.4), the modiolus was decalcified in 10% EDTA on a swing bed for 5 days at room temperature. Next, the specimens were post-fixed in 1% OsO₄ in 0.1 M sodium cacodylate buffer (pH 7.4) for 2 h at 4°C. Dehydration was performed through a graded ethanol series, propylene oxide and a series of ethoxyl ine resins. The modiolus were then embedded in the final resin overnight at 70°C. Semi-thin (1 μm) sections in the mid-modular plane were cut with diamond knives, collected on glass slides, stained with 1% methylene blue, and then rinsed three times with distilled water. The sections were subsequently examined and photographed under a light microscope (Nikon, Tokyo, Japan) for both qualitative and quantitative analysis of SGNs in Rosenthal’s canal from the first to the third turn. In each turn, SGNs were counted in 10 slices that were collected across a ~800 μm distance to get an average SGN number in each cross view of Rosenthal’s canal.

To observe the impact of furosemide on the cochlear lateral wall, the spiral ligament and affiliated Stria Vascularis (SV) were dissected and decalcified for 1 day; the rest of the process was similar to that described above for SGN lesion observation. The samples in experimental groups (Groups 1-3) would be compared against those in normal control group (Group 4).

**Figure 1:** Comparison of the body weight changes caused by consecutive administration of Cis versus one injection of Cis in combination with furosemide.

**Figure 2:** ABR threshold shifts.
(a) Shift of the frequency average after a single furosemide injection. The shift was largest (approximately 53 dB) 30 min after furosemide application (the earliest time tested) and recovered at 12 h.
(b) Threshold shift audiogram 7 days after consecutive cisplatin injections (3 mg/kg × 5d). The shift was biased to high frequencies (up to 50 dB) and was lower at lower frequencies (around 30 dB).
(c) Threshold shift audiogram 7 days after co-administration of 200 mg/kg furosemide and various Cis doses. Cis was administered 30 min after furosemide.
(d) Threshold shift audiogram 7 days after co-administration of 0.2 mg/kg Cis and 200 mg/kg furosemide at various intervals. The threshold shift was nearly zero at a zero interval and higher at longer intervals.
Statistics

The SPSS software (ver. 18.0; SPSS Inc., Chicago, IL, USA) was used for all analyses. The threshold shifts and cell numbers across groups and subgroups were analyzed using one-way or two-way analysis of variance (ANOVA) plus post-hoc tests. A difference was considered statistically significant if \( p < 0.05 \). All data are presented as means ± SEM.

Results

Guinea pig

Body weight changes: The animals’ body weight recovered in the co-administration group, but decreased progressively during the experiment in the Cis-alone group (Figure 1). For example, at the end of the experiment, the body weight loss for the co-administration subgroup (furosemide 200 mg/kg, Cis 0.2 mg/kg, 30 min interval) was only 21.6 ± 6 g, while the corresponding value for cis-alone group was 118.6 ± 5.0 g (\( t_{0.05, 11} = 13.0, p < 0.01 \)).

Effects of a single furosemide injection: Of 15 guinea pigs that received single-injection furosemide, 12 survived, six of which were sacrificed at 0.5 and 1.5 h (three at each time point) after the furosemide injection; the rest were sacrificed 12 h after the injection when the threshold had recovered fully. Diuretic administration caused only a temporary ABR threshold shift, which was greatest 0.5-1.5 h after the injection. The threshold shift was flat across the frequencies tested. Thus, we calculated the frequency average of threshold shifts at various times after furosemide administration (Figure 2a). The largest shift, 53.3 ± 2.5 dB, was seen at 0.5 h after the furosemide injection, the earliest time at which the ABR was recorded after furosemide. The shift decreased gradually thereafter and had disappeared 12 h after the furosemide injection.

Morphological observation of the effects of furosemide focused on the lateral wall of the cochlea, and was demonstrated through comparisons with four vehicle control guinea pigs (Figure 3). Changes in morphology were more evident in the SV of the basal turn. Compared with the normal control (Figure 3d), samples obtained 30 min after furosemide injection showed clearly swollen Marginal Cells (MCs) that bulged into the scala media, so that this surface of the SV became rough (Figure 3a, arrows). At this time point, many vacuoles were seen. (ANOVA) plus post-hoc tests. A difference was considered statistically significant if \( p < 0.05 \). All data are presented as means ± SEM.

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across the three cellular layers of SV into spiral ligament and the SV in the cross-section view was totally broken. The damage appeared to be reduced in samples acquired 90 min after furosemide administration (Figure 3b): the vacuoles were smaller and the swollen MCs decreased in number. Samples taken 12 h after administration showed no obvious pathology compared with the control (Figure 3c), suggesting a full recovery of the structure after furosemide administration. No morphological abnormality was detected by optical microscopy in the basilar membrane or the organ of Corti from any animal in this group.

**Cochlear lesions in the Cis-alone group:** In this group of 11 guinea pigs, Cis was given (i.m.) for 5 consecutive days at a dose of 3 mg/kg per day. Seven animals survived 7 days after the last injection (Table 1), resulting in a mortality rate of 36%. Figure 2b shows that the ABR threshold shifted from the baseline across frequencies 7 days after the last Cis injection. Significant threshold shifts were evident graphically and statistically in a one-way ANOVA on the frequency averaged threshold ($F_{2,21}=226.6, p<0.01$), with a frequency averaged shift of 43 ± 5 dB and the largest shift of 50 ± 7 dB at 16 kHz (Figure 2b).

Hair cell loss corresponded roughly to functional deterioration. The overall Out Hair Cell (OHC) loss was ~18% for the whole cochlea and followed the classic pattern: more in the basal turn and less in the apical turn (Figure 4a). No inner hair cell loss was apparent in this group.

Consecutive Cis application also caused damage to SGNs. Figure 5 compares SGN lesions across all groups. As shown in Figure 5, the survived SGNs were counted under an optical microscope and compared against the normal control across all treatment groups. SGNs that showed clear lesions were not counted as survived (Figure 6a). Since there was obviously no difference across groups above basal turn, our statistical analysis was done only in basal turn. One way ANOVA shows a clear effect of treatment ($F_{3,31}=6.1, p<0.01$). However, post-hoc tests show that only in the Cis-alone group, the SGN count was significantly lower than that of control (Tukey Test, $p<0.01$). Myelin sheath detachment from the perikarya was also found in a few SGNs in the basal turn (Figure 6a) in this group. However, the number of lost SGNs appeared to be fewer than the number of OHC lost.

**Cochlear lesions in the co-administration group:** Out of totally 41 guinea pigs to which furosemide and Cis were co-administered, 34 survived for 7 days after the Cis injection. While mortality in this group was lower than that in the group that received consecutive Cis injections. For example, the mortality in the subgroup that received 0.2 mg/kg Cis 30 min after furosemide was only 14%, compared with 36% in the Cis-alone group. Six subgroups were classified according to the Cis doses and time intervals between the furosemide and Cis injections. Figure 2c shows the threshold shifts in animals 7 days post the drugs administration which received different Cis doses with the time interval fixed at 30 min between furosemide and Cis. Cis administered at 0.1 mg/kg caused slight hearing loss only at very high frequencies in this subgroup (32 kHz, $t_{30.1}=8.215, p<0.01$), while 0.2 mg/kg Cis (30 min interval) caused much greater hearing loss across the whole frequency range with a frequency average of 46.3 + 2.5 dB, similar to that in the Cis-alone group (43.0 + 5.4 dB) in which a total dose of 15 mg/kg Cis was given ($t_{30.1}=0.512, p=0.62$), 75 times as high as that used in this Cis-fur subgroup. A further 10x increase in Cis dose did not lead to a proportional elevation in thresholds. The subgroup that received the highest Cis dose (2 mg/kg) had the largest threshold shifts (52.5 ± 4.4 dB in frequency average). A one-way ANOVA showed a significant effect of dose ($F_{2,30}=170.0, p<0.01$). Post hoc tests on total OHC loss count within the factor of dose showed that the OHC loss was significantly less in 0.1 mg/kg subgroup than in 0.2 mg/kg and 2 mg/kg subgroups, with the latter showing almost 100% loss (Tukey Test, $p<0.01$ for both 0.1 versus 0.2 and 0.1 vs 2 mg/kg) (Table 1). In every subgroup, OHC loss was biased to the basal end (Figure 4b).

Figure 2d shows the effect of time interval between furosemide and Cis administration on ABR threshold shifts when the Cis dose was fixed at 0.2 mg/kg. Unexpectedly, simultaneous co-administration of the two drugs failed to cause any hearing loss (near zero threshold shifts). A two-way ANOVA was performed across the four subgroups of 0, 30,


The present study demonstrated that by the recruitment of the method of co-administration of furosemide and Cis, the survival rate and physical conditions of subjects were boosted. Cis and furosemide co-administration was effective in establishing cochlear lesions. Using this regimen, 0.2 mg/kg Cis was found to cause more hearing loss and OHC lesions, no significant SGN lesion or loss or myelin sheath damage was seen in any subgroup (Figure 5, 6b-d), as evaluated 7 days after co-administration.

**Mice**

**Hearing function:** Of 17 mice receiving gentamicin and furosemide treatments, 16 subjects survived (Table 2). Significant threshold shifts were observed in both gentamicin 40 mg/kg and 60 mg/kg applications. When furosemide was combined with 40 mg/kg gentamicin, significant threshold shift only occurred in 32 kHz, while it was threshold shift across frequencies when gentamicin was up to 60 mg/kg (Figure 7).

**Hair cell lesion:** Hair cell lesion corresponded roughly to functional deterioration in mice. With gentamicin dose increasing, OHC lesion showed a deterioration tendency. No inner hair cell loss was apparent in each group. In the group that accepted gentamicin 40 mg/kg application, OHC loss was 31%. In the group receiving gentamicin 60 mg/kg, OHC loss was 75% (Figure 8).

**Discussion**

The present study demonstrated that by the recruitment of the method of co-administration of furosemide and Cis, the survival rate and physical conditions of subjects were boosted. Cis and furosemide co-administration was effective in establishing cochlear lesions. Using this regimen, 0.2 mg/kg Cis was found to cause more hearing loss and OHC loss than that produced by 3 mg/kg cis alone for 5 consecutive days. This reduction in cis usage was largely responsible for the reduced mortality. Animals were healthier, as illustrated by the difference in body weight changes among the groups. More importantly, the two approaches impacted the SGNs differently: ~10% of the SGN in the basal turn was affected in the group receiving Cis alone, while no such lesion was seen in any of the co-administration subgroups, while the OHC loss was higher.

**Mortality and weight changes**

Use of the various regimens resulted in marked differences in mortality rates. The lower mortality rate, 17%, of co-administration represents a great advantage of this approach. This reduction in mortality is likely due to the lower dose of Cis used. Cis-induced mortality is known to be dose-dependent. In one study in guinea pigs, a single dose of Cis caused no evident gastrointestinal toxicity or nephrotoxicity and thus no body weight loss or mortality, unless the dose was >6 mg/kg [9]. The dose of Cis used in our co-administration regimen was far lower than this critical point. Thus, the mortality

<table>
<thead>
<tr>
<th>GM</th>
<th>Adjudin</th>
<th>n</th>
<th>Frequency average of shifts (dB)</th>
<th>OHC loss % in basal turn</th>
<th>OHC loss % in apical turn</th>
</tr>
</thead>
<tbody>
<tr>
<td>40 mg/kg</td>
<td>Negative</td>
<td>9(8)</td>
<td>8 ± 0.4</td>
<td>31 ± 0.3</td>
<td>10 ± 0.5</td>
</tr>
<tr>
<td>60 mg/kg</td>
<td>Negative</td>
<td>9(9)</td>
<td>33 ± 2</td>
<td>75 ± 1</td>
<td>52 ± 1</td>
</tr>
</tbody>
</table>

The frequency average of shifts was calculated by an average the threshold shift in each tested frequency, the percentage value of OHC loss was calculated as the result of lost OHCs divided by the total OHCs in the corresponding area. n = sample size (survived animals). Apical turn is the region of basilar membrane from 0 to 2.4 mm (begin at apex); basal turn is the region of basilar membrane from 2.4 to 5.2 mm.

Table 2: Threshold shifts of ABR tests and percentage OHC loss by regimen and observation stage in mice.
observed in the co-administration subgroup was not likely caused by Cis; rather, it may be due to the impact of deep anesthesia.

**Impact of furosemide on the blood-labyrinth barrier**

The main effect of these diuretics on biochemistry is to depress the Na^+^-K^-2Cl^- symporter (co-transporter) in the thick ascending limb of the loop of Henle, inhibiting sodium and chloride reabsorption. Because a similar Na^+^-K^-2Cl^- co-transporter is expressed in cochlear MCs [20], which are responsible for the generation and maintenance of Endocochlear Potential (EP), the application of these drugs decreases EP and thus cochlear transduction [21]. However, depression of this transporter seems unlikely to be the reason for the cellular damage in SV. Previous research showed that SV damage appears to be initiated by ischemia observed shortly after diuretic administration [22]. The lack of blood supply to the SV likely causes damage to various cells and the tight junctions among the cells. As the tight junctions bordering the SV and scala media exist among MCs and those bordering the SV and spiral ligament among the basal cells, damage to MCs and basal cells would be expected to create leakage of the drugs between the plasma and cochlear fluid [23]. Although we did not intend to explore the time course of SV pathological changes in the present study, the observed changes in the morphology and function suggested increased permeability of SV to drugs for at least 90 min. This should provide a window of time for furosemide to facilitate Cis entry into the cochlea. The cells in the SV appear to have a notable self-repair ability, and leakage in this region is sealed within 12 h after furosemide application.

**Mechanism of the cis and furosemide synergism in the inner ear**

Enhancement of Cis-induced ototoxicity by diuretic co-administration has been reported previously [14,15,19,24]. This synergistic effect has largely been attributed to two mechanisms. First, as described above, the SV damage by diuretics may increase cochlear drug quantities [25]. Although the Cis concentration with the co-administration of furosemide has not been determined, EA has been demonstrated to significantly increase the concentration of gentamicin in perilymph if they are applied within an appropriate time frame [26]. Second, temporary interruption of blood flow through the SV may result in localized ischemia-reperfusion injury, followed by a burst of oxygen radical species in the endolymph [27]. The increased oxidative stress may interact with the direct effect of Cis on apoptosis in the organ of Corti.

Timing is an important issue for the synergistic interaction between the two drugs. In the present study, we failed to find cochlear lesions in the subgroup in which Cis and furosemide were administered simultaneously.

However, the cochlear damage appeared to increase with increasing interval from 0 to 90-min between furosemide and cis. This change in the synergistic effect may be explained by the pharmacokinetics of the two drugs. Furosemide given as an intramuscular injection likely takes 30 min to damage the SV to an observable level. To our knowledge, however, there has been no systematic observation on the time line of SV damage by furosemide. If the Cis is largely eliminated from plasma before this time, no synergistic effect would be seen. The reported half-life of Cis injected i.v. can be divided into two phases: the fast phase has been reported as 20-30 min [28] or less than 1 h in other studies [29,30], while the half-life in the slow phase lasts several days [29,30]. We assume that only the Cis in the fast-phase would be responsible for the potential cochlear lesion in our case because in the co-administration with zero interval regimen, the Cis dose was low (0.2 mg/kg) and the residual drug in the plasma during the slow phase would likely not be sufficient to cause damage, even if it could freely enter the cochlea. Because the fast phase half-life is short, the Cis likely was largely eliminated when the SV damage peaked, and thus the amount of Cis that entered the cochlea would be insufficient to have any effect. This may be the case even if we take into consideration the fact that an intramuscular injection, not i.v., was used. On the other hand, the fact that Cis injection 60 min and 90 min post-furosemide caused a larger loss of OHC than that at 30 min post-furosemide may be due to the coincidence of increased SV permeability and Cis application, as well as the reperfusion that may have occurred 60 and 90 min after furosemide administration. One possibility is that SV permeability had partially recovered 60 and 90 min after furosemide administration; the increased blood flow at this time may bring more Cis to the SV and thus to the organ of Corti. Another possibility is that free radical levels continued to increase after the onset of ischemia and remained high for at least 40-80 min post-reperfusion [31]. OHC pathological changes, such as swelling and rupture, were found to be induced during this period, presumably via the increased stress of the hydroxyl radicals caused by ischemia-reperfusion [32]. Although the amount of Cis that enters the cochlea may be lower when it is applied 60 and 90 min post-furosemide injection versus if it was applied earlier, the synergistic interaction between Cis and free radicals may cause more hair cell loss. These ideas should be assessed in future studies.

**Differences between co-administration and cis alone**

While OHC death was comparable between the two approaches, the co-administration approach failed to create significant SGN lesions in the present study. To our knowledge, no report addresses the potential difference in SGN lesion induction between the two approaches. The mechanism of this difference is unclear, although we believe that it is likely due to the selective effect of furosemide on SV blood circulation. The lateral wall of the cochlea is supplied by radiating arterioles lateralis, while the modiolus and spiral lamina are supplied by radiating arterioles mediales and the cochlear artery. The blood flow in the vessels supplying the lateral wall of the cochlea diminished rapidly after injection of diuretics. In contrast, the radiating arterioles mediales to the modiolus and the spiral lamina in the cochlea retained a normal appearance at all times [14,25,33]. The mechanism(s) by which the diuretics selectively target lateral wall vessels is unknown. Probably, lateral wall vessels likely have an ultrastructure similar to that of the thick ascending limb of the loop of Henle in the kidney, the basis of why loop diuretics target these two organs. As the dose of Cis was much lower in the co-administration regimens, the amount of Cis that entered the cochlea via other branches of the cochlear artery was also likely low if furosemide did not change the permeability of those vessels. Thus, the Cis leakage through the blood vessels to the SGNs was insufficient to cause SGN lesions.

**Co-administration of gentamicin and furosemide in mice**

Gentamicin-induced ototoxic model was also commonly utilized in otological researches. The ototoxic animal model of gentamicin was also obsessed by the mortality of consecutive injections of gentamicin of mice [34]. The regimen of co-administration of gentamicin and furosemide in mice was extension of that in guinea pig. This regimen established an effective ototoxic model, and performed less animal mortality (6%) compared with 50% in early report. However, the sequence of gentamicin or Cis with furosemide was different due to the different half life of gentamicin and Cis. The
half-life of gentamicin (approximately 1.6 h) was much longer than that of Cis (0.5 h) [35].

In the preliminary experiment of mice, it was difficult to implement the anesthesia by ketamine and pentobarbital, since all of the mice could not survive under this anesthetic regimen. We turned to another approach: increase the dose of furosemide to enhance the permeability of SV.

Conclusions

In summary, in this study we demonstrated the advantages of the co-administration of Cis and furosemide in establishing Cis ototoxicity in terms of a more rapid process and reduced mortality. However, the disadvantage of this approach is the lack of SGN lesions as seen in the Cis-alone regimen. Unless the purpose of a study is an OHC lesion, the disadvantage of this approach is the lack of SGN lesions as seen in the co-administration of Cis and furosemide in establishing Cis ototoxicity.

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Citation:


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