Compositions for use in methods for treating hearing loss
Zusammensetzungen zur Verwendung in Verfahren zur Behandlung von Gehörverlust
Compositions pour l’utilisation dans des méthodes de traitement contre la surdité

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FIELD OF THE INVENTION

[0001] The present invention relates to compositions for treating and preventing hearing loss induced by noise.

BACKGROUND OF THE INVENTION

[0002] A major cause of acquired hearing loss is loud noise. Exposure to harmful noise levels is common in the workplace. The National Institute for Occupational Safety and Health estimates that about 30 million workers in the United States encounter hazardous levels of noise. (Franks et al. (1996) Preventing Occupational Hearing Loss - A Practical Guide, DHHA (NIOSH) Publication No. 96-110, p.1). These levels are encountered in, for example, construction, mining, agriculture, manufacturing and utilities, transportation, and in the military. The incidence of noise associated hearing loss continues to increase in spite of efforts to regulate job related noise exposure, and to improve the use of hearing protective devices such as ear muffs and ear plugs.

[0003] Another cause of hearing loss is exposure to ototoxic drugs such as cisplatin and aminoglycoside antibiotics. Accordingly, there is a need for methods and compositions to prevent or treat hearing loss.


[0005] US6177434 discloses the use of agents that augment the inner ear antioxidant defenses such as adenosine agonists or up-regulating agents and/or agents which increase inner ear glutathione levels to prevent or reverse hearing loss induced by noise or toxins.

SUMMARY OF THE INVENTION

[0006] In one aspect the present invention provides compositions for use in methods for ameliorating hearing loss induced by noise, the methods each comprising the step of administering to a subject an amount of an otoprotective composition that is effective to ameliorate hearing loss. The otoprotective composition includes at least ebselen. In some embodiments, the otoprotective composition comprises a pharmaceutically effective amount of at least one glutathione peroxidase mimic which is ebselen. In some embodiments, the otoprotective composition comprises a pharmaceutically effective amount of (a) at least one glutathione peroxidase mimic which is ebselen and (b) at least one xanthine oxidase inhibitor (e.g., a composition comprising ebselen and allopurinol). In some embodiments, the otoprotective composition comprises a pharmaceutically effective amount of (a) at least one glutathione peroxidase mimic which is ebselen and (b) at least one glutathione or glutathione precursor (e.g., a composition comprising ebselen and N-acetyl-cysteine).

[0007] In some embodiments, the otoprotective composition comprises a pharmaceutically effective amount of (a) at least one glutathione peroxidase mimic which is ebselen, (b) at least one xanthine oxidase inhibitor, and (c) at least one glutathione or glutathione precursor (e.g., a composition comprising ebselen, allopurinol, and N-acetyl-cysteine).

[0008] Another aspect of the present invention provides otoprotective compositions useful for ameliorating hearing loss induced by noise. The otoprotective compositions include at least ebselen. In some embodiments, the otoprotective compositions comprise at least one glutathione peroxidase mimic which is ebselen. In some embodiments, the otoprotective compositions comprise at least one glutathione peroxidase mimic which is ebselen and at least one glutathione or glutathione precursor.

[0009] In some embodiments, the otoprotective compositions comprise at least one glutathione peroxidase mimic which is ebselen, at least one xanthine oxidase inhibitor, and at least one glutathione or glutathione precursor.

BRIEF DESCRIPTION OF THE DRAWINGS

[0010] The foregoing aspects and many of the attendant advantages of this invention will become more readily appreciated as the same become better understood by reference to the following detailed description, when taken in conjunction with the accompanying drawings, wherein:

FIGURE 1 shows a graph comparing the threshold shifts of the auditory evoked brainstem response (ABR) tested with a 4-16 kHz click stimulus 1 day, 1 week, 2 weeks, and 3 weeks after exposing control rats (A), N-acetyl-cysteine-treated rats (B), and ebselen-treated rats (C) to 115 dB, 4-16 kHz noise for 4 hours. The difference in threshold shifts between control and ebselen-treated rats was highly significant at 1 day, 1 week, and 2 weeks post noise exposure (p <0.01) and significant at 3 weeks post noise exposure (p <0.05). The difference in threshold shifts...
between control and N-acetyl-cysteine-treated rats was significant at all time points post noise exposure (p <0.05). Threshold shifts from baseline were measured in 5 dB intervals.

FIGURE 2 shows a graph comparing the threshold shifts of the ABR tested with a 4-16 kHz click stimulus, and stimuli at frequencies of 4 kHz, 8 kHz, 12 kHz, and 16 kHz, 3 weeks after exposing control rats (A), 16 mg/kg ebselen-treated rats (B), 16 mg/kg allopurinol-treated rats (C), and 8 mg/kg ebselen/allopurinol-treated rats (D) to 115 dB, 4-16 kHz noise for 4 hours. The difference in threshold shifts between control (A) and ebselen-treated rats (B) was highly significant for the click and 4 kHz stimuli (p<0.01). The difference in threshold shifts between control (A) and ebselen/allopurinol-treated rats (C) was highly significant for the click, 4 kHz, and 16 kHz stimuli (p <0.01) and significant for the 12 kHz stimulus (p <0.05). Threshold shifts from baseline were measured in 5 dB intervals.

FIGURE 3 shows a graph comparing the threshold shifts of the ABR tested with stimuli at frequencies of 4 kHz, 8 kHz, 12 kHz, and 16 kHz, 3 weeks after exposing control rats (A) and ebselen-treated rats (B) two times to 110 dB, 4-16 kHz noise for 4 hours, separated by three weeks. The difference in threshold shifts between control and ebselen-treated rats was highly significant for the 8 kHz stimulus (p <0.01), and significant for the 16 kHz stimulus (p <0.05). Threshold shifts from baseline were measured in 5 dB intervals.

FIGURE 4A-D shows graphs comparing the percentage of outer hair cell loss as a function of distance from the apex of the cochlea, after exposing control rats (A and C) and 16 mg/kg ebselen treated-rats (B and D) at two times, separated by three weeks, to 110 dB, 4-16 kHz noise for 4 hours. The number of outer hair cells lost in rats A and C was 401 and 246, respectively, the number of outer hair cells lost in rats B and D was 90 and 56, respectively.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

[0011] As used herein, the term "otoprotectant" refers to a chemical substance that is capable of ameliorating hearing loss.

[0012] As used herein, the term "ameliorating hearing loss" includes: (a) reducing the magnitude and/or duration of hearing loss; and/or (b) slowing the progression of hearing loss; and/or (c) preventing the onset of hearing loss that would occur without administration of an otoprotectant composition of the invention.

[0013] As used herein, the term "ototoxic agent" means an agent that is likely to impair the function of any component of the ear involved in hearing and, therefore, is likely to induce temporary or permanent hearing loss. Examples of ototoxic agents are ototoxic drugs and ototoxic noise.

[0014] As used herein, the term "exposure to an ototoxic agent" includes single or multiple exposures to an ototoxic agent that is recognized in the art as being likely to cause temporary or permanent hearing loss. For example, the Occupational Safety and Health Administration (OSHA) considers exposures to noise greater than or equal to 85 decibels (dB) to be hazardous to hearing. Thus, OSHA mandates that workers not be exposed to greater than or equal to 85 dB of noise over a continuous eight hour period based on a time weighted average, unless noise reduction measures (i.e., ear muffs) are employed.

[0015] As used herein, the term "otoprotectant composition" refers to a composition that includes at least one otoprotectant, and may include more than one otoprotectant. Otoprotectant compositions may also include, in addition to one or more otoprotectant(s), pharmaceutically acceptable carriers that facilitate administration of an otoprotectant composition to a mammalian subject.

[0016] In one aspect the present invention provides compositions for use in methods for ameliorating hearing loss induced by noise, the methods each comprising the step of administering to a subject an amount of an otoprotectant composition that is effective to ameliorate hearing loss. The compositions of the invention are applicable to any mammalian subject, such as a human subject. The otoprotectant composition may be administered before, during or after exposure to an ototoxic agent such as noise.

[0017] The otoprotectant compositions can include one or more than one otoprotectant. Unless stated otherwise, any isomeric or tautomeric form of any of the otoprotectants disclosed herein can be used in the invention. Some otoprotectants that can be included in otoprotectant compositions of the invention include glutathione and glutathione precursors.

Representative examples of otoprotectants in this category are: methionine; N-acetyl-DL-methionine; S-adenosylmethionine; cysteine; homocysteine; N-acetylcysteine; glutathione; glutathione ethylester; glutathione diethylester; glutathione triethylester; cysteamine; cystathione; N,N'-diacetyl-L-cystine (DiNAC); 2(R,S)-D-ribo-(1',2',3',4'-tetrahydroxybutyl)-thiazolidine-4-carboxylic acid (RibCys); 2-alkythiazolidine 2(R,S)-D-ribo-(1',2',3',4'-tetrahydroxybutyl)thiazolidine-4-carboxylic acid (OTCA).

[0018] Xanthine oxidase inhibitors, for example allopurinol (C₄H₇N₄O) and its tautomers, are useful as otoprotectants in the practice of the invention. The following representative allopurinol derivatives are useful as otoprotectants in the practice of the invention: 1-methylallopurinol; 2-methylallopurinol; 5-methylallopurinol; 7-methylallopurinol; 1,5-dimethylallopurinol; 2,5-dimethylallopurinol; 1,7-dimethylallopurinol; 2,7-dimethylallopurinol; 5,7-dimethylallopurinol; 2,5,7-trimethylallopurinol; 1-ethoxycarbonylallopurinol; and 1-ethoxycarbonyl-5-methylallopurinol.

[0019] Glutathione peroxidase mimics which is ebselen are useful as otoprotectants in the practice of the invention.
Other examples of glutathione peroxidase mimics: 6A,6B-diseleninic acid-6A',6B'-selenium bridged \( \beta \)-cyclodextrin (6-diSeCD); and 2,2'-diseleno-bis-Beta-cyclodextrin (2-diSeCD).

Table 1 sets forth representative effective dosage ranges for some of the otoprotectants described herein. The otoprotectants set forth in Table 1 are preferably administered orally or intravenously. The otoprotectants set forth in Table 1 can be administered to a mammalian subject before, during or after exposure to an ototoxic agent, such as ototoxic noise. Typically, a mammalian subject receives at least one dose of at least ebselen before and after each exposure to ototoxic noise. In some embodiments, a mammalian subject receives one dose of at least ebselen before exposure to ototoxic noise and at least one dose of at least ebselen after exposure to ototoxic noise. In some embodiments, a mammalian subject receives at least two daily doses of at least ebselen for a single exposure to an ototoxic agent, such as an exposure to ototoxic noise lasting for about 1 to about 6 hours. In some embodiments, a mammalian subject receives at least three daily doses of at least ebselen for repeated exposures to ototoxic noise or prolonged exposures to an ototoxic agent, such as exposures to ototoxic noise lasting longer than about 6 hours.

In some embodiments of the invention, an otoprotectant composition comprising one or more otoprotectants is administered to a mammalian subject at one or more times during a period extending from 18 hours before exposure of the mammalian subject to ototoxic noise, to 18 hours after exposure of the mammalian subject to ototoxic noise. In some embodiments of the invention, an otoprotectant composition comprising one or more otoprotectants is administered to a mammalian subject at one or more times during a period extending from one hour before exposure of the mammalian subject to ototoxic noise, to one hour after exposure of the mammalian subject to ototoxic noise. In some embodiments of the invention, an otoprotectant composition comprising one or more otoprotectants is administered to a mammalian subject at one or more times during a period extending from 30 minutes before exposure of the mammalian subject to ototoxic noise, to 30 minutes after exposure of the mammalian subject to ototoxic noise. In some embodiments of the invention, an otoprotectant composition comprising one or more otoprotectants is administered to a mammalian subject at one or more times during a period extending from 10 minutes before exposure of the mammalian subject to ototoxic noise, to ten minutes after exposure of the mammalian subject to ototoxic noise. In some embodiments of the invention, an otoprotectant composition comprising one or more otoprotectants is administered to a mammalian subject concurrently with exposure of the mammalian subject to an ototoxic agent, such as ototoxic noise.

The abbreviation "mg" means milligrams.
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<th>Compound(s)</th>
<th>Chemical name</th>
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<th>Presently more preferred range</th>
<th>Presently most preferred range</th>
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<td>500-1000mg/day</td>
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<td>5-5000mg/day</td>
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<td>2-diSeCD</td>
<td>2,2’-diseleno-bis-Beta-cyclodextrin</td>
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<td>500-1000mg/day</td>
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<td>Presently more preferred range</td>
<td>Presently most preferred range</td>
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<tr>
<td>-----------------------------</td>
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<td>---------------------------</td>
<td>-------------------------------</td>
<td>--------------------------------</td>
</tr>
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<td>5,7-dimethylallopurinol</td>
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</table>
The otoprotectant compositions can include one, or more than one, otoprotectant(s). Otoprotectant compositions of the invention are defined in the appended claims. In some embodiments of the otoprotectant compositions that include more than one otoprotectant, the otoprotectant compositions are formulated to provide an effective dosage of the individual constituent otoprotectants as set forth in Table 1. In some embodiments, the combination of otoprotectants may act synergistically, as described in EXAMPLE 2.

Also disclosed are otoprotectant compositions that each comprise at least two (e.g., two, three, four, five, six, seven, eight, nine, or ten) of the individual otoprotectants. For example, some otoprotectant compositions include at least one otoprotectant selected from Group A, at least one otoprotectant selected from Group B, and at least one otoprotectant selected from Group C, wherein Groups A, B and C include the following otoprotectants:

- **Group A (glutathione or a glutathione precursor):** methionine; N-acetyl-DL-methionine; S-adenosylmethionine; cysteine; N-acetylcysteine; glutathione; glutathione ethylester; glutathione diethylester; glutathione triethylester; DIPC; RbCys; homocysteine; cystathione; cysteamine; OTCA and RibCyst.
- **Group B (xanthine oxidase inhibitors):** allopurinol; 1-methylallopurinol; 2-methylallopurinol; 5-methylallopurinol; 7-methylallopurinol; 1,5-dimethylallopurinol; 2,5-dimethylallopurinol; 1,7-dimethylallopurinol; 2,7-dimethylallopurinol; 5,7-dimethylallopurinol; 2,5,7-trimethylallopurinol; 1-ethoxyxycarbonylallopurinol; and 1-ethoxyxycarbonyl-5-methylallopurinol.
- **Group C (glutathione peroxidase mimics):** Ebelsen; 2-diSeCD; and 6-diSeCD.

The otoprotectant compositions of the invention are useful for ameliorating hearing loss induced by exposure to ototoxic noise.

The otoprotectant compositions of the invention can be formulated to provide a dosage that is effective to ameliorate hearing loss in a subject exposed to an ototoxic agent. For example, in some embodiments the otoprotectant compositions are formulated to provide an effective dosage of the individual otoprotectants as set forth in Table 1.

Administration of the otoprotectant compositions of the invention is accomplished by any effective route, e.g., orally or parenterally, as described in EXAMPLES 1-3. Methods of parenteral delivery include topical, intra-arterial, subcutaneous, intramedullary, intravenous, or intranasal administration. In addition to one or more otoprotectants, the otoprotectant compositions may contain suitable pharmaceutically acceptable carriers comprising excipients and other compounds that facilitate administration of the otoprotectant compositions to a mammalian subject. Further details on techniques for formulation and administration may be found in the latest edition of “Remington’s Pharmaceutical Sciences” (Maack Publishing Co, Easton PA).

Otoprotectant compositions for oral administration can be formulated using pharmaceutically acceptable carriers well known in the art, in dosages suitable for oral administration, as described in EXAMPLES 1-3. Such carriers enable the otoprotectant compositions to be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions, etc., suitable for ingestion by a subject.

Otoprotectant compositions for oral use can be obtained, for example, through combination of one or more otoprotectants with solid excipient, optionally grinding the resulting mixture, and processing the mixture of granules, after adding suitable additional compounds, if desired, to obtain tablets or dragee cores. Suitable excipients include carbohydrate or protein fillers. These include, but are not limited to, sugars, including lactose, sucrose, mannitol, or sorbitol, starch from corn, wheat, rice, potato, or other plants; cellulose such as methyl cellulose, hydroxypropylmethylcellulose, or sodium carboxymethylcellulose; and gums including arabic and tragacanth; as well as proteins, such as gelatin and collagen. If desired, disintegrating or solubilising agents may be added, such as the cross-linked polyvinyl pyrrolidone, agar, alginic acid, or a salt thereof, such as sodium alginate.

Dragee cores are provided with suitable coatings such as concentrated sugar solutions, which may also contain gum arabic, talc, polyvinylpyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures. Dyestuffs or pigments may be added to the tablets or dragee coatings for product identification or to characterise the quantity of active compound (i.e., dosage).

Otoprotectant compositions, which can be used orally, can be formulated, for example, as push-fit capsules made of gelatin, as well as soft, sealed capsules made of gelatin and a coating such as glycerol or sorbitol. Push-fit capsules can contain otoprotectants mixed with filler or binders such as lactose or starches, lubricants such as talc or magnesium stearate, and, optionally, stabilizers. In soft capsules, the otoprotectant(s) may be dissolved or suspended in suitable liquids, such as fatty oils, liquid paraffin, or liquid polyethylene glycol with or without stabilizers.

Otoprotectant compositions for parenteral administration include aqueous solutions of one or more otoprotectants, as described in EXAMPLES 1-3. For injection, the otoprotectant compositions of the invention may be formulated in aqueous solutions, preferably in physiologically compatible buffers such as Hank’s solution, Ringer’s solution, or physiologically buffered saline. Aqueous injection suspensions may contain substances, which increase the viscosity of the suspension, such as sodium carboxymethyl cellulose, sorbitol, or dextran. Additionally, suspensions of otoprotectants may be prepared as appropriate oily injection suspensions. Suitable lipophilic solvents or vehicles include fatty oils such as sesame oil, or synthetic fatty acid esters, such as ethyl oleate or triglycerides, or liposomes. Optionally, the suspension may also contain suitable stabilizers or agents, which increase the solubility of the compounds to allow for the preparation of the suspension.
For topical or nasal administration, penetrants appropriate to the particular barrier to be permeated are typically used in the formulation. Such penetrants are generally known in the art.

The otoprotectant compositions of the present invention may be manufactured in a manner similar to that known in the art (e.g., by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping or lyophilising processes). The otoprotectant compositions may also be modified to provide appropriate release characteristics, e.g., sustained release or targeted release, by conventional means (e.g., coating).

The otoprotectant compositions may be provided as a salt and can be formed with many acids, including but not limited to hydrochloric, sulfuric, acetic, lactic, tartaric, malic, succinic, etc. Salts tend to be more soluble in aqueous or other protonic solvents than are the corresponding free base forms.

After such otoprotectant compositions formulated in an acceptable carrier have been prepared, they can be placed in an appropriate container and labeled for use.

The amount actually administered will be dependent upon the individual to which treatment is to be applied, and will preferably be an optimized amount such that the desired effect is achieved without significant side-effects. The determination of an effective dose is well within the capability of those skilled in the art. Of course, the skilled person will realize that divided and partial doses are also within the scope of the invention.

For any otoprotectant composition, the effective dose can be estimated initially either in cell culture assays or in any appropriate animal model (e.g., primate, rats and guinea pigs and other small laboratory animals), as described in EXAMPLES 1-3. The animal model is also typically used to achieve a desirable concentration range and route of administration. Such information can then be used to determine useful doses and routes for administration in humans or other mammals.

Therapeutic efficacy and possible toxicity of otoprotectant compositions can be determined by standard pharmaceutical procedures, in cell cultures or experimental animals (e.g., ED50, the dose therapeutically effective in 50% of the population; and LD50, the dose lethal to 50% of the population). The dose ratio between therapeutic and toxic effects is the therapeutic index, and it can be expressed as the ratio ED50/LD50. Otoprotectant compositions which exhibit large therapeutic indices are preferred. The data obtained from cell culture assays and animal studies is used in formulating a range of dosage for use in humans or other mammals. The dosage of such compounds lies preferably within a range of circulating concentrations that include the ED50 with little or no toxicity. The dosage typically varies within this range depending upon the dosage form employed, sensitivity of the patient, and the route of administration.

EXAMPLE 1

This example shows that ebselen administration protects against hearing threshold shifts and cochlear hair cell loss in rats exposed to noise.

1. Methods

Noise exposure paradigm: 8-week old female Fisher-344 rats were exposed to 110-115 dB noise at 4-16 kHz for 4 hours.

Physiologic analyses: The auditory evoked brainstem response (ABR) was used to assess hearing in each ear in each animal before and after exposure to noise using equipment from Intelligent Hearings System. ABR generated with a click stimulus (broad spectrum stimulus, 4-16 kHz) was measured in 5 dB intervals. Animals were reevaluated at 1 day, 1 week, 2 weeks, and 3 weeks post noise exposure. ABR changes measured 1 day post noise exposure are considered to represent a temporary threshold shift (TTS); ABR changes measured 3 weeks post noise exposure are considered to represent a permanent threshold shift (PTS).

Morphologic analyses: Hair cell counts in cochlea from test animals following noise exposure and physiologic evaluation using ABR were determined by carefully dissecting cochlea to obtain inner ear sensory epithelia. The tissues were stained with fluorescein-labeled phalloidin (specific for actin filaments, which are abundant in hair cells) and 4,6-diamidino-2-phenylindole (DAPI; specific for the DNA in the nucleus in cells). Samples were then mounted on microscope slides and viewed with epifluorescence to determine the extent of hair cell loss and retention. The correlation between physiologic data (ABR) and morphological data (hair cell counts) allows for confirmation of the protective effects of otoprotectants of the invention.

Dosing of Otoprotectants: Ebselen was dissolved at 4 mg/mL in 10% DMSO and administered to rats at 16 mg/kg. N-acetyl-cysteine was dissolved at 100 mg/mL in saline and was administered to rats at 325 mg/kg. About 0.5 ml of ebselen solution or N-acetyl-cysteine solution was injected intra-peritoneally or delivered by oral gavage twice daily the day prior to, the day of, and the day following exposure to noise. Control animals were dosed on an identical schedule with vehicle (10% DMSO) only.

Statistical analysis: All experiments were performed with four rats per study group, and measuring each ear
independently before and after noise exposure. Data was collected in blinded studies. Statistical analyses were performed by analysis of variance (ANOVA) between study groups.

2. Results

[0049] As shown in FIGURE 1, ABR data generated with a 4-16 kHz click stimulus measured in 5 dB intervals show a highly significant reduction in TTS in animals administered ebselen at one day after noise exposure, and a significant reduction in PTS at three weeks after noise exposure, compared with controls. The reduction in TTS and PTS in animals administered ebselen was at least two-fold, compared to control animals. Similarly, the ABR data show a significant reduction in TTS in animals administered N-acetyl-cysteine at all time points after noise exposure.

[0050] The physiologic data correlates with morphologic data from the same test animals showing less hair cell loss in cochlea from animals treated with otoprotectants compared to the control group. For example, in an ebselen-treated animal with 0 dB PTS and a significantly reduced TTS, preservation of most outer hair cells was observed, whereas there was almost complete loss of outer hair cells in a control animal with a 15 dB PTS. Similar preservation of outer hair cells was observed in N-acetyl-cysteine-treated rats. Therefore, the administration of ebselen or N-acetyl-cysteine results in physiologic protection from noise-induced hearing loss and hair cell loss.

[0051] Similarly, the administration of allopurinol was found to provide protection from noise-induced hearing loss and hair cell loss (see EXAMPLE 2).

EXAMPLE 2

[0052] This example shows that combined administration of ebselen and allopurinol provides greater protection from hearing loss than each compound administered alone.

1. Methods

[0053] Noise exposure paradigm: 8-10 week old female Fisher-344 rats were exposed to 110-115 dB noise at 4-16 kHz for 4 hours.

[0054] Physiologic analyses: ABR generated with a click stimulus (broad spectrum stimulus, 4-16 kHz), and with tone stimuli at frequencies of 4 kHz, 8 kHz, 12 kHz, and 16 kHz, was measured in 5 dB intervals, as described in EXAMPLE 1. Animals were tested before and at 3 weeks post noise exposure to assess PTS.

[0055] Dosing of compounds: Ebselen was dissolved at 4 mg/ml in 10% DMSO and allopurinol was dissolved at 4 mg/ml in water. Ebselen and allopurinol were administered to rats at an individual dose of 16 mg/kg or at a combined dose of 8 mg/kg each. About 0.5 ml of ebselen solution, allopurinol solution, or ebselen/allopurinol solution was injected intra-peritoneally or delivered by oral gavage twice daily the day prior to, the day of, and the day following exposure to noise. Control animals were dosed on an identical schedule with vehicle only.

[0056] Statistical analysis: All experiments were performed with four rats per study group, and measuring each ear independently before and after noise exposure. Data was collected in blinded studies. Statistical analyses were performed by ANOVA between study groups.

2. Results

[0057] Physiologic data as measured by ABR indicates that ebselen and allopurinol, alone or in combination, afford significant protection from both temporary and permanent thresholds shifts in rats exposed to noise. FIGURE 2 shows that ebselen and allopurinol delivered by intra-peritoneal injection at 16 mg/kg individually result in significant reductions in PTS at three weeks after noise exposure, compared with controls. At half the individual dose (i.e., 8 mg/kg each), the combination of ebselen and allopurinol provides a level of protection that is greater than additive (synergy).

EXAMPLE 3

[0058] This example shows that ebselen administration protects against hearing threshold shifts and cochlear hair cell loss in rats repeatedly exposed to noise.

1. Methods

[0059] Noise exposure paradigm: 8-week old female Fisher-344 rats were exposed to 110 dB noise at 4-16 kHz for 4 hours two times three weeks apart.

[0060] Physiologic analyses: ABR generated with tone stimuli at frequencies of 4 kHz, 8 kHz, 12 kHz, and 16 kHz,
was measured in 5 dB intervals, as described in EXAMPLE 1. Animals were tested before and 3 weeks following the repeated noise exposure to evaluate permanent threshold shift (PTS).

**[0061] Morphologic analyses:** Hair cell counts in cochlea from test animals following noise exposure and physiologic evaluation using ABR were determined as described in EXAMPLE 1.

**[0062] Dosing of Ebselen:** Ebselen was dissolved at 4 mg/mL in 10% DMSO and administered to rats at a dose of 16 mg/kg. About 0.5 ml of ebselen solution was injected intra-peritoneally or delivered by oral gavage twice daily the day prior to, the day of, and the day following each exposure to noise. Control animals were dosed on an identical schedule with vehicle only.

**[0063] Statistical analysis:** All experiments were performed with four rats per study group, and measuring each ear independently before and after noise exposure. Data was collected in blinded studies. Statistical analyses were performed by ANOVA between study groups.

2. Results

**[0064]** Physiologic data measured by ABR indicates that ebselen provides significant protection from both temporary and permanent threshold shifts in rats from repeated exposure to noise. The administration of ebselen results in a significant reduction in TTS at 1 day after repeated noise exposures, compared with controls. In addition, the PTS at three weeks after repeated noise exposures was significantly reduced in these animals compared to controls, as shown in FIGURE 3. The physiologic data correlates with morphologic data from the same test animals showing less hair cell loss in cochlea from animals treated with ebselen compared to the control group, as shown in FIGURE 4.

Claims

1. An otoprotectant composition for use in ameliorating noise-induced hearing loss in a mammalian subject, comprising a pharmaceutically effective amount of 2-phenyl-1,2-benzoisoselenazol-3(2H)-one (ebselen) or a pharmaceutically acceptable salt thereof.

2. The composition for use of Claim 1 comprising:

   (a) a pharmaceutically effective amount of 2-phenyl-1, 2-benzoisoselenazol-3 (2H)-one (ebselen) or a pharmaceutically acceptable salt thereof; and

   (b) a pharmaceutically effective amount of at least one otoprotectant selected from the group consisting of allopurinol, 1-methylallopurinol, 2-methylallopurinol, 5-methylallopurinol, 7-methylallopurinol, 1,5-dimethylallopurinol, 2,5-dimethylallopurinol, 1,7-dimethylallopurinol, 2,7-dimethylallopurinol, 5,7-dimethylallopurinol, 2,5,7-trimethylallopurinol, 1-ethoxycarbonylallopurinol, 1-ethoxycarbonyl-5-methylallopurinol, and pharmaceutically acceptable salts thereof.

3. The composition for use of Claim 1 or claim 2 comprising:

   (a) a pharmaceutically effective amount of 2-phenyl-1, 2-benzoisoselenazol-3 (2H)-one (ebselen) or a pharmaceutically acceptable salt thereof; and

   (b) a pharmaceutically effective amount of at least one otoprotectant selected from the group consisting of allopurinol, 1-methylallopurinol, 2-methylallopurinol, 5-methylallopurinol, 7-methylallopurinol, 1,5-dimethylallopurinol, 2,5-dimethylallopurinol, 1,7-dimethylallopurinol, 2,7-dimethylallopurinol, 5,7-dimethylallopurinol, 1-ethoxycarbonylallopurinol, 1-ethoxycarbonyl-5-methylallopurinol, and pharmaceutically acceptable salts thereof;

   wherein ebselen is present in an amount of 5-5000 mg/day and each of the otoprotectants in group (b) is present in an amount of 10-2400 mg/day.

4. The composition for use of Claim 2 or 3, wherein the otoprotectant composition comprises ebselen and allopurinol, or a pharmaceutically acceptable salt thereof.

5. The composition for use of Claim 2 or 3, further comprising a pharmaceutically effective amount of N-acetylcysteine or a pharmaceutically acceptable salt thereof.

6. The composition for use of Claim 5, comprising ebselen, allopurinol, and N-acetylcysteine, or pharmaceutically acceptable salts thereof.
7. The composition for use of Claim 1, wherein ebselen is present in an amount of 5-5000 mg/day.

8. The composition for use of Claim 1, wherein ebselen is present in an amount of 50-2000 mg/day.

9. The composition for use of Claim 1, wherein ebselen is present in an amount of 500-1000 mg/day.

10. The composition for use of Claim 4, wherein ebselen is present in an amount of 5-5000 mg/day and allopurinol is present in an amount of 10-2400 mg/day.

11. The composition for use of Claim 4, wherein ebselen is present in an amount of 500-1000 mg/day and allopurinol is present in an amount of 100-800 mg/day.

12. The composition for use of Claim 6, wherein each of the ebselen and N-acetylcysteine is present in an amount of 5-5000 mg/day and allopurinol is present in an amount of 10-2400 mg/day.

13. The composition for use of Claim 6, wherein each of the ebselen and N-acetylcysteine is present in an amount of 500-1000 mg/day and allopurinol is present in an amount of 100-800 mg/day.

14. The composition for use of any one of the preceding Claims, further comprising at least one pharmaceutically acceptable carrier suitable for oral administration to a mammalian subject.

15. The composition for use of any one of the preceding Claims, further comprising a solid excipient comprising at least one of a carbohydrate or protein filler.

16. The use of a composition as defined in any one of the preceding Claims in the manufacture of a medicament for use in ameliorating noise-induced hearing loss in a mammalian subject.

17. The use of Claim 16, wherein the mammalian subject is a human subject.

Patentansprüche

1. Otoprotektive Zusammensetzung zur Anwendung in der Verbesserung von lärminduzierten Hörverlust bei einem Säugetiersubjekt, die eine pharmazeutisch wirksame Menge an 2-Phenyl-1,2-benzisoselenazol-3(2H)-on (EBSELEN) oder einem pharmazeutisch annehmbaren Salz davon, umfasst.

2. Zusammensetzung zur Anwendung nach Anspruch 1, umfassend:

   (a) eine pharmazeutisch wirksame Menge an 2-Phenyl-1,2-benzisoselenazol-3(2H)-on (EBSELEN) oder einem pharmazeutisch annehmbaren Salz davon; und

   (b) eine pharmazeutisch wirksame Menge an mindestens einem otoprotektiven Präparat, ausgewählt aus der Gruppe, die aus Allopurinol, 1-Methylallopurinol, 2-Methylallopurinol, 5-Methylallopurinol, 7-Methylallopurinol, 1,5-Dimethylallopurinol, 2,5-Dimethylallopurinol, 1,7-Dimethylallopurinol, 2,7-Dimethylallopurinol, 5,7-Dimethylallopurinol, 2,5,7 Trimethylallopurinol, 1-Ethoxycarbonylallopurinol, 1-Ethoxycarbonyl-5-Methylallopurinol und pharmazeutisch annehmbaren Salzen besteht.

3. Zusammensetzung zur Anwendung nach Anspruch 1 oder Anspruch 2 umfassend:

   (a) eine pharmazeutisch wirksame Menge an 2-Phenyl-1,2-benzisoselenazol-3(2H)-on (EBSELEN) oder einem pharmazeutisch annehmbaren Salz davon; und

   (b) eine pharmazeutisch wirksame Menge an mindestens einem otoprotektiven Präparat, ausgewählt aus der Gruppe, die aus Allopurinol, 1-Methylallopurinol, 2-Methylallopurinol, 5-Methylallopurinol, 7-Methylallopurinol, 1,5-Dimethylallopurinol, 2,5-Dimethylallopurinol, 1,7-Dimethylallopurinol, 2,7-Dimethylallopurinol, 5,7-Dimethylallopurinol, 1-Ethoxycarbonylallopurinol, 1-Ethoxycarbonyl-5-Methylallopurinol und pharmazeutisch annehmbaren Salzen davon besteht,

   wobei EBSELEN in einer Menge von 5-5000 mg/Tag vorliegt und jedes sich in Gruppe (b) befindliche otoprotective Präparat (b) in einer Menge von 10-2400 mg/Tag vorliegt.
4. Zusammensetzung zur Anwendung nach Anspruch 2 oder 3 wobei die otoprotektive Zusammensetzung EBSELEN und Allopurin umfasst, oder ein pharmazeutisch annehmbares Salz davon.

5. Zusammensetzung zur Anwendung nach Anspruch 2 oder 3, die ferner eine pharmazeutisch wirksame Menge an N-Acetylcystein, oder einem pharmazeutisch annehmbaren Salz davon, umfasst.


7. Zusammensetzung zur Anwendung nach Anspruch 1, wobei EBSELEN in einer Menge von 5-5000 mg/Tag vorliegt.

8. Zusammensetzung zur Anwendung nach Anspruch 1, wobei EBSELEN in einer Menge von 50-2000 mg/Tag vorliegt.

9. Zusammensetzung zur Anwendung nach Anspruch 1, wobei EBSELEN in einer Menge von 500-1000 mg/Tag vorliegt.

10. Zusammensetzung zur Anwendung nach Anspruch 4, wobei EBSELEN in einer Menge von 5-5000 mg/Tag vorliegt und Allopurinol in einer Menge von 10-2400 mg/Tag vorliegt.

11. Zusammensetzung zur Anwendung nach Anspruch 4, wobei EBSELEN in einer Menge von 500-1000 mg/Tag vorliegen und Allopurinol in einer Menge von 100-800 mg/Tag vorliegt.

12. Zusammensetzung zur Anwendung nach Anspruch 6, wobei EBSELEN und N-Acetylcystein jeweils in einer Menge von 5-5000 mg/Tag vorliegen und Allopurinol in einer Menge von 10-2400 mg/Tag vorliegt.

13. Zusammensetzung zur Anwendung nach Anspruch 6, wobei EBSELEN und N-Acetylcystein jeweils in einer Menge von 500-1000 mg/Tag vorliegen und Allopurinol in einer Menge von 100-800 mg/Tag vorliegt.

14. Zusammensetzung zur Anwendung nach jedem der vorangehenden Ansprüche, die ferner mindestens einen pharmazeutisch annehmbaren Träger, der zur oralen Verabreichung an ein Säugetiersubjekt geeignet ist, umfasst.


17. Verwendung von Anspruch 16, wobei es sich bei dem Säugetiersubjekt um ein menschliches Subjekt handelt.

Revendications

1. Composition de protection auditive destinée à être utilisée pour l’amélioration de l’état de surdité induite par du bruit chez un sujet mammifère, comprenant une quantité efficace sur le plan pharmaceutique de 2-phenyl-1,2-benzoiselenazonol-3(2H)-one (ebselect) ou d’un sel de celui-ci acceptable sur le plan pharmaceutique.

2. Composition destinée à être utilisée selon la revendication 1, comprenant :
   (a) une quantité efficace sur le plan pharmaceutique de 2-phenyl-1,2-benzoiselenazonol-3(2H)-one (ebselect) ou d’un sel de celui-ci acceptable sur le plan pharmaceutique ; et
   (b) une quantité efficace sur le plan pharmaceutique d’au moins un protecteur auditif sélectionné dans le groupe se composant d’allopurinol, 1-methylallopurinol, 2-methylallopurinol, 5-methylallopurinol, 7-methylallopurinol, 1,5-dimethylallopurinol, 2,5-dimethylallopurinol, 1,7-dimethylallopurinol, 2,7-dimethylallopurinol, 5,7-dimethylallopurinol, 2,5,7-trimethylallopurinol, 1-ethoxycarbonylallopurinol, 1-ethoxycarbonyl-5-methylallopurinol et des sels de ceux-ci acceptables sur le plan pharmaceutique.

3. Composition destinée à être utilisée selon la revendication 1 ou la revendication 2, comprenant :

13.
(a) une quantité efficace sur le plan pharmaceutique de 2-phenyl-1,2-benzoiselenazol-3(2H)-one (ebselen) ou d’un sel de celui-ci acceptable sur le plan pharmaceutique ; et
(b) une quantité efficace sur le plan pharmaceutique d’au moins un protecteur auditif sélectionné dans le groupe se composant d’allopurinol, 1-methylallopurinol, 2-methylallopurinol, 5-methylallopurinol, 7-methylallopurinol, 1,5-dimethylallopurinol, 2,5-dimethylallopurinol, 1,7-dimethylallopurinol, 2,7-dimethylallopurinol, 5,7-dimethylallopurinol, 1-ethoxycarbonylallopurinol, 1-ethoxycarbonyl-5-methylallopurinol et des sels de ceux-ci acceptables sur le plan pharmaceutique ; dans laquelle ebselen est présent dans une quantité de 5 à 5000 mg/jour et chacun des protecteurs auditifs dans le groupe (b) est présent dans une quantité de 10 à 2400 mg/jour.

4. Composition destinée à être utilisée selon la revendication 2 ou 3, dans laquelle la composition de protection auditive comprend ebselen et allopurinol, ou un sel de ceux-ci acceptable sur le plan pharmaceutique.

5. Composition destinée à être utilisée selon la revendication 2 ou 3, comprenant en outre une quantité efficace sur le plan pharmaceutique de N-acetylcysteine ou d’un sel de celui-ci acceptable sur le plan pharmaceutique.

6. Composition destinée à être utilisée selon la revendication 5, comprenant ebselen, allopurinol et N-acetylcysteine, ou des sels de ceux-ci acceptables sur le plan pharmaceutique.

7. Composition destinée à être utilisée selon la revendication 1, dans laquelle ebselen est présent dans une quantité de 5 à 5000 mg/jour.

8. Composition destinée à être utilisée selon la revendication 1, dans laquelle ebselen est présent dans une quantité de 50 à 2000 mg/jour.

9. Composition destinée à être utilisée selon la revendication 1, dans laquelle ebselen est présent dans une quantité de 500 à 1000 mg/jour.

10. Composition destinée à être utilisée selon la revendication 4, dans laquelle ebselen est présent dans une quantité de 5 à 5000 mg/jour et allopurinol est présent dans une quantité de 10 à 2400 mg/jour.

11. Composition destinée à être utilisée selon la revendication 4, dans laquelle ebselen est présent dans une quantité de 500 à 1000 mg/jour et allopurinol est présent dans une quantité de 100 à 800 mg/jour.

12. Composition destinée à être utilisée selon la revendication 6, dans laquelle chacun d’ebselen et N-acetylcysteine est présent dans une quantité de 5 à 5000 mg/jour et allopurinol est présent dans une quantité de 10 à 2400 mg/jour.

13. Composition destinée à être utilisée selon la revendication 6, dans laquelle chacun d’ebselen et N-acetylcysteine est présent dans une quantité de 500 à 1000 mg/jour et allopurinol est présent dans une quantité de 100 à 800 mg/jour.

14. Composition destinée à être utilisée selon l’une quelconque des revendications précédentes, comprenant en outre au moins un porteur acceptable sur le plan pharmaceutique adapté à une administration orale à un sujet mammifère.

15. Composition destinée à être utilisée selon l’une quelconque des revendications précédentes, comprenant en outre un excipient solide comprenant au moins l’un d’un carbohydrate ou d’une charge de protéine.


17. Utilisation selon la revendication 16, dans laquelle le sujet mammifère est un sujet humain.
TIME POST NOISE EXPOSURE

Fig. 1.
Fig. 3.
**Fig. 4A.**

**Fig. 4B.**
**Fig. 4C.**

**Fig. 4D.**
REFERENCES CITED IN THE DESCRIPTION

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