# **Manuscript Details**

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Title	Nanoparticle drug delivery systems for inner ear therapy: an overview
Short title	Nanoparticles for inner ear drug delivery
Article type	Review Article

#### Abstract

Local drug delivery based on nanoparticles (NP) represents a novel strategy to improve inner ear treatments. The intratympanic delivery of NP may be suitable to treat or prevent hearing loss originating from damage to hair cells and spiral ganglion neurons in the cochlea. Numerous experimental studies support in vitro and in vivo the biocompatibility of NP, their physical stability, target specificity, cell/tissue uptake and ability to internalize therapeutic agents. The topical use of NP helps to reduce the amount of drug required and avoid systemic side effects. This review focuses on recent findings and applications of different NP systems locally administered in the inner ear. The perspectives for clinical application of NP in inner ear drug delivery are also discussed.

Keywords	Nanoparticles; inner ear; drug delivery; intratympanic administration; local administration.
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Corresponding Author's Institution	University of Padua
Order of Authors	Filippo Valente, Laura Astolfi, Edi Simoni, Serena Danti, Valeria Franceschini, Milvia Chicca, alessandro martini
Suggested reviewers	Wei Liu, Marcelo N. Rivolta, barbara zavan, Isabel Varela Nieto

# Submission Files Included in this PDF

#### File Name [File Type]

Cover letter 1.docx [Cover Letter]

Answer\_to\_Reviewers1.docx [Response to Reviewers]

MS review Revision Marked.docx [Revised Manuscript with Changes Marked]

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NP\_Figure\_Legends1.docx [Figure]

Figure 1 inner ear barrier.jpg [Figure]

Figure 2 administration routes.jpg [Figure]

Figura 3 nanoparticles.jpg [Figure]

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Dear Professor Siepmann,

thank you very much for sending me the Reviewers' comments on the manuscript "Nanoparticle drug delivery systems for inner ear therapy: an overview" (JDDST\_2017\_23) by Valente, Astolfi and coworkers.

One of the Reviewers mentioned that he/she did not find the Figure captions. Indeed, we had some problems in manuscript uploading: we checked on the journal site and could not find the file of Figure captions. We therefore enclose the missing Figure captions file for you and for the Reviewer.

Here enclosed you will find the detailed answers to the Reviewers. We hope that now our manuscript is suitable for publication on JDDST.

Kind regards,

Laura Astolfi, PhD

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Reviewer 1

The review presented by Valente et al « Nanoparticle drug delivery systems for inner ear therapy: an overview" is well-written and interesting.

I have the following minor comments/suggestions regarding the manuscript.

1. Lines 43-44 "The NP with size between 10 and 100 nm are useful for application in biology and medicine for innovative DD systems". This range is a little bite too restrictive. The range 10 to 200 nm is commonly used.

We thank the Reviewer for his/her comment and corrected the sentence according to the suggestion.

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According to the Reviewer's suggestion, we introduced the references in the sentence.

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- A prolonged residence time at the site of injection as well as in the round window were achieved without any negative effect on the hearing thresholds of the animals.

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#### 22 Abstract

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31

#### 32 Keywords

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#### 34 **1. Introduction**

The treatment of inner ear diseases through drug delivery (DD) faces numerous challenges (1), 35 among which the limited blood flow to the inner ear (2), the presence of physical barriers acting as 36 a selective filter for drug transportation to the inner ear from the circulatory system (3), the small 37 size of the cochlea and its isolated location in the petrous bone. As a result, research in local drug 38 applications and medications has recently attracted interest because it is a more effective and 39 40 preferable treatment than the systemic one. Case studies involving steroids (4) and gentamicin 41 treatment for Meniere's disease (5) have been documented, but these approaches could be improved for clinical protocols by the development of controlled and targeted delivery systems. 42

Nanoparticles (NP) are a possible option to improve existing therapeutic strategies (6). The NP with
size between 10 and 200 nm are useful for application in biology and medicine for innovative DD
systems. NP-based strategy could be more efficient and reduce drug-associated side effects because

of the ability to deliver the therapeutic agent to the target site. Moreover, the controlled release of
compounds conjugated to NP results in a lower dose of drug required to achieve the therapeutic
effects (1, 7).

49 The cochlea is a good model for studying the NP-based DD due to its isolated structure and the

50 perilymph rheology. The intratympanic delivery of NP could be suitable to treat the hearing loss

- and prevent its progression when hair cells and spiral ganglion neurons are damaged (8).
- 52 Several works and reviews have been published in the past decade, focusing on NP type, pathology 53 involved, delivery approach or a combination of these topics (1-4, 6-8). The goal of the present 54 review is to provide an updated general overview of NP-based strategies and their advantages and
- 55 disadvantages for local DD into the inner ear.
- 56

### 57 **2. Ear barriers**

The human inner ear consists of two main parts, the auditory system (the cochlea) and the vestibular 58 system. The cochlea is a bony spiral canal, about 30 mm long and divided in three fluid-filled 59 compartments: the scala tympani, the scala media and the scala vestibuli. The round window 60 membrane (RWM), the blood inner ear barrier (BB) and the oval window are physical barriers that 61 isolate the cochlea from the middle ear and from the circulatory system (Figure 1). The RWM is a 62 63 three-layer semi-permeable membrane, composed of an outer epithelial cell layer, a middle 64 connection layer and an inner connection layer facing the perilymph of the scala tympani (9). In 65 humans, the variable thickness of RWM affects the response of patients to DD treatments. In animal models, its thickness is different among species but its composition is similar (10). 66

Both the RWM and the oval window membranes have been investigated for DD, as connections between the middle ear cavity and the cochlear perilymph. The DD strategies for the inner ear currently rely mostly on RWM (11). The passage of molecules across this membrane is not only influenced by thickness, but also by its morphological integrity, inflammation and weight, concentration, liposolubility and external charge of the therapeutic compound (12). The drugs deposited topically in the middle ear cavity are internalized by pinocytosis and transported to the perilymph through blood vessels or by diffusion. Thus the direct application of drugs in the proximity of RWM is a suitable approach for treatment of inner ear pathologies (13).

The BB is a major barrier in the stria vascularis separating the cochlear tissues from the circulatory 75 system (14). Its role is to maintain the homeostasis of cochlear fluids and protect the inner ear 76 77 integrity. Its main components are principally the endothelial capillaries whose cells are connected by tight junctions, which lay over a basement membrane. However numerous accessory cells have 78 recently been observed in the complex structure of the barrier, such as pervcites and perivascular 79 80 resident macrophage-like (11). The BB has been described to act as a physical and biochemical 81 barrier through an efflux pump, the P-glycoprotein 1 (P-gp) (15). The BB is therefore considered a rate-limiting barrier in the passage of therapeutic agents from the circulatory system to the inner ear. 82 83 However, the current knowledge about drug transportation processes through BB is still limited (16). 84

85

#### 86 **3. Administration routes**

87 The clinical protocols for inner ear therapies mostly rely on systemic and local DD routes. The systemic administration represents a classical route for DD, but in the inner ear only few drugs may 88 reach the target site at therapeutic concentrations. If high doses of systemic drugs are employed, 89 90 often side effects are developed (17, 18). Systemic applications of NP in inner ear have been recently investigated: poly(lactic-co-glycolic acid) NP conjugated with rhodamine B and applied 91 systemically were detected in the liver, but not in the cochlea (19). The limited bioavailability of NP 92 after systemic administration could be due to the rapid clearance from the circulation in liver and 93 spleen (20). 94

Local administration appears more suitable for inner ear DD (19). This approach allows a quick 95 96 distribution of the drug inside the cochlea, improving their delivery to the target site; it also requires lower drug doses, avoiding side effects (21) (Figure 2). Two main routes are presently used for this 97 purpose, the intratympanic (IT) or the intracochlear administration, but the second one is rarely 98 performed because it is highly invasive and limited to surgery cases (22). On the contrary, the IT 99 injection is minimally invasive and relies on passive diffusion of the active molecules through 100 101 RWM to access the inner ear. This review focuses on development of these methods for DD with minimal trauma for the cochlea. However, local delivery trials show a high variability in results 102 (23) because of some key factors: 1) the drug clearance within the middle ear through the 103 104 Eustachian tube; 2) the permeability of RWM; and 3) the residence time of the drug in contact with RWM (24). A method to reduce variability of results and increase the drug concentration in the 105 perilymph could be to better control the residence time of the drug at close range with RWM, using 106 107 specific delivery systems based on NP (25).

108

#### 109 4. Nanoparticle-based systems

The NP (also called nanocarriers or nanovectors) are artificial compounds with size at the 110 111 nanoscale, which aim to compensate for adverse drug properties such as low solubility, degradation and short half-life (26). The NP may also be adapted to target a specific tissue of the inner ear. 112 113 However, when injected in the middle ear as a liquid suspension, NP will undergo clearance 114 through the Eustachian tube (27), thus significantly reducing their residence time near RWM. The NP suitable for DD systems should therefore increase the residence time, together with the ability to 115 cross RWM and their biocompatibility (Figure 3). A detailed description of physico-chemical 116 117 characteristics of NP and their applications is reported.

118 4.1. Lipid Core NP

5

Lipid Core NP (LCN) possess a lipid core matrix (usually triglycerides) with a surrounding shell of 119 120 lecithin, polyethylene glycol or poloxamers as stabilizing agents. The LCN structure can be changed to include different drugs and control the kinetics of drug release (28). It has been shown to 121 be stable up to six months in aerosol dispersion (29). These NPs did not induce toxicological effects 122 in vivo in mice after systemic applications (12 mg/kg intravenously for five days) (30) and their cell 123 uptake and cell viability was *in vitro* verified on fibroblasts by confocal scanner laser microscopy 124 125 (31). In rat animal models LCN were able to cross RWM and reach inner ear targets after middle ear application *in vivo*, while not affecting hearing capacity (32). Their preferred pathway to diffuse 126 inside the cells was also investigated: they followed a "nerve pathway", diffusing from the 127 128 perilymph in the scala tympani to the spiral ganglion, nerve fibres and later approaching the inner and the outer hair cells (33). Their variability in diffusion and ability to cross RWM depends on 129 their lipid composition, size and external charge. The ability to cross the RWM has been shown to 130 131 be size-dependent, because the percentage of particle diffusion was inversely proportional to their size (31). Surface charge may also affect the uptake and biodistribution of LNC. Some NP 132 candidates based on glycerol mono-oleate were studied under different external charges: after an in 133 vivo application to RWM, LCN expressing stronger positive charges were detected in the deeper 134 turns of the cochlea (34). The LCN were also tested as a drug carrier, delivering dexamethasone in 135 136 the inner ear through IT injection and comparing the results with a systemic application of the same LCN. The amount of dexamethasone detected in cochlear fluid after local LCN application was 137 significantly higher compared to the systemic application, also increasing the half-life and the 138 average residence time of the drug in the perilymph by 1.9 folds (35). All these results indicate a 139 great potential for LCN for sustained drug release and targeting of inner ear tissues after local 140 administration. 141

142

#### 143 **4.2. Liposomes**

Liposomes are artificial phospholipid bilayers, similar to those found in the cell membrane, but 144 145 surrounding an aqueous core. They exhibit a wide size range (between 50 nm and 5 µm) and morphology, depending on the phospholipid used and the preparation method (36). Liposomes can 146 encapsulate either hydrophobic molecules in the phospholipid bilayer or hydrophilic molecules in 147 their aqueous core (37). The uptake of these NP in vitro or in vivo usually relies on the passive 148 diffusion inside the cells, but their surface can be modified with polyethylene glycol, antibodies, 149 150 peptides, carbohydrates, hyaluronic acid and folic acid (35). Such modified liposomes successfully targeted cells expressing tropomyosin receptor-B (TrkB) by using 18-mer peptides to promote 151 cellular uptake (38). Liposomes labelled with fluorescent markers applied in vivo to a mouse model 152 153 with a single IT injection were identified in all cochlear turns, with a concentration gradient decreasing from the base to the apex and, to a lesser extent, in the lateral wall and in the organ of 154 Corti. No morphological or functional damages to the inner ear were detected 24 hours after the 155 156 application (8). Disulfiram, a neurotoxic agent, was used as model payload for DD analysis: NP loaded with Disulfiram damaged the spiral ganglion 48 hours after application, with an associated 157 threshold shift reaching 35 dB. No significant effects were observed with a similar application of a 158 pure Disulfiram solution (8). To test the drug delivery efficiency of liposome nanocarriers, NP of 159 different size (95, 130, 240 nm) encapsulating the contrast agent gadolinium-tetra-azacvclo-160 161 dodecane-tetra-acetic acid (Gd-DOTA) were applied in the middle ear and analyzed with MRI: the results showed that the liposome carrier efficiency was inversely proportional to NP size (39, 40). 162

## 163 **4.3. Polymersomes and copolymers**

The polymersomes (also called multifunctional NP) are a wide class of amphiphilic copolymers, consisting of a self-assembled membrane of hydrophobic units, surrounding an aqueous core, and of a hydrophilic corona (41). Structurally they are similar to liposomes, with the advantages that the membrane thickness can be controlled by the molecular weight of the hydrophobic block of copolymer to achieve stronger, thicker and more stable membranes. The hydrophilic corona can be modified to regulate the biodistribution of polymersomes and induce specific cellular uptake (42).
Hydrophilic drugs can be loaded in the core, while hydrophobic ones in the membrane (43).

Different multifunctional polymersomes were studied for inner ear DD targeting specific tissue orconjugated with ferromagnetic materials.

In a mouse model, poly(ethylene glycol)-*b*-poly (ε-caprolactone) NP (PEG-*b*-PCL) labeled with fluorescent markers were detected in the spiral ganglion, in the organ of Corti and in the lateral wall after 24 hours from RWM application *in vivo* (8). Tissue specificity was also investigated: PEG-*b*-PCL were conjugated with a nerve growth factor derived peptide and tested *ex-vivo* on explanted mouse cochleae and *in vitro* on PC12 cells. No significant toxic effect was observed and a specific targeting to spiral ganglion neurons, Schwann cells and nerve fibres was achieved by conjugating the NP with tyrosin kinase and p75 neurotrophin receptors (44).

Poly(2-hydroxyethyl aspartamide) NP (PHEA) were observed to enter in vitro the immortalized 180 mouse organ of Corti cell line (HEI-OC1) and the human middle ear cell line (HMEEC). When 181 applied *in vivo* near the RWM in a mouse model, PHEA were also detected in the inner ear tissue 182 (45). In order to improve NP uptake, PHEA were modified with oligoarginine peptide, a positively 183 charged copolymer, and conjugated with fluorescent Nile red as a hydrophobic model drug (46). In 184 185 these conditions the NP uptake *in vitro* on HEI-OC1 and HMEEC cells was significantly improved 186 after 15 and 24 hours, compared to pure Nile red solution. Modified PHEA were detected after 24 hours from application in the inner hair cells and supporting cells (47). 187

Poly(lactic-co-glycolic acid) (PLGA) NP are copolymers among the novel carrier developed for DD. The Food and Drug Administration (FDA) and the European Medicines Agency (EMA) approved PLGA NP for parenteral administration (47). PLGA NP are interesting because of their hydrophilicity, biocompatibility and easy derivatization by functional groups on the surface or inside the polymer. Their surface may be modified for target specificity by PEGylation, chitosan

absorption and binding of antibodies and oligopeptides (48), and different molecules (proteins, 193 194 steroids, antibiotics and nucleic acids) have been successfully encapsulated and delivered by PLGA **NP** (49). Programmed degradation of the polymer may therefore yield quantitative delivery of 195 drugs, plasmids or other bioactive molecules. The PLGA NP tested in the inner ear were first 196 conjugated with rhodamine B, a red fluorescent dye, and applied via IT injection: they were 197 identified in the scala tympani, showing that PLGA NP are able to cross RWM by diffusion and 198 199 their clearance depends on the perilymph flow rate (19). A quantitative pharmacokinetic study recently showed that **PLGA NP** applied locally *in vivo* in Guinea pigs significantly improved the 200 drug distribution within the inner ear (52). When PLGA NP were loaded with the fluorescent dye 201 202 coumarin-6 and applied through IT injection, the concentration of the compound after 96 hours from treatment was 10.9-fold higher in the perilymph than when administered in pure solution. 203 Similar results were obtained for other therapeutic payloads such as antioxidants and antiapoptotic 204 205 drugs (50). Thus **PLGA NP** are an useful DD system for inner ear because of their high versatility in adaptation to drug properties and tissue targets (51). 206

207

#### 208 4.5. Silica NP

Silica NP are modified colloidal silica particles (52) used to transfect *in vitro* plasmid DNA (53) but also as a DD system (54). A pilot study in mice tested the efficacy of diffusion of Cy3-labeled silica NP administered near the RWM: these NP were found inside the inner hair cells, the vestibular hair cells, the spiral ganglion neurons and the supporting cells, without any hearing impairment. Since the NP also reached the dorsal cochlear nucleus and the superior olivary complex, the authors suggested a retrograde axonal transport and concluded that silica NP could be applied for safe drug deliver in the auditory system (55).

216 4.6. Supermagnetic iron oxide NPs (SPIONs)

Magnetic NP are synthetic  $Fe_3O_4$  (magnetite) particles, with a core diameter around 15 nm, that can 217 218 be widely applied for magnetic targeting of cells (56). Unlike large ferromagnetic materials, the smaller supermagnetic iron oxide NP (SPION) are characterized by the absence of residual 219 magnetic interactions when the magnetic field is not active, thus they are more suitable for 220 biomedical applications (57). The SPION derivatized to increase biocompatibility and cell 221 interactions could be guided by an external magnetic field to a specific biological target, but they 222 223 cannot encapsulate any drug (58). For in vivo applications, to prevent particle aggregation and favour dispersion SPION were coated by organic compounds (59). In inner ear drug delivery, 224 SPION have been encapsulated in PLGA (60), silica (58) and dextran (61) and their 225 226 biocompatibility was tested and verified in vitro and in vivo (59). The mobility of SPION induced by a magnetic field was also quantified and the results of flux density, gradients and NP properties 227 were compared between in vitro and in vivo models (62). The magnetic force required for SPION to 228 229 cross RWM in vivo in Guinea pigs was significantly lower than that of the in vitro RWM model (63). Another study in vivo in Guinea pigs revealed that the concentration of coated SPION inside 230 the cochlea significantly increased (330% above control) when a magnetic field was active (64). 231 Recently, SPION coated with PGLA NP were tested as drug carriers with dexamethasone-acetate 232 233 (Dex-Ac) as a payload: the levels of Dex-Ac detected in the inner ear fluids after 1 hour from 234 treatment were significantly higher compared with those in absence of a magnetic field (65). All these results support the application of SPION for inner ear drug delivery protocols. 235

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#### 237 4.7. Hyperbranched poly-L-lysine NP

Hyperbranched poly-L-lysine (HBPL) are high cationic charged dendrimers widely used for nonviral gene transfer (67, 68). The HBPL were applied *in vivo* in Guinea pig inner ears without any sign of cell toxicity or permanent hearing loss (31): they were detected in the stria vascularis and hair cells (31). Nanoparticles based on HBPL and conjugated with fluorescein isothiocyanate were tested *ex-vivo* on freshly frozen human temporal bones, placing them near the intact RWM: HBPL
were detected in hair cells, nerve fibres and other cochlear tissues (66).

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# 245 5. Key aspects for nanoparticle-based drug delivery in the inner ear

There are several key parameters to consider for NP-based local DD in the inner ear: the RWM permeability; the NP cochlear targeting; their payload ability and the controlled drug release; their biocompatibility and their stability in cochlear fluids and tissues. All these aspects were evaluated with different NP systems *in vitro, ex-vivo* or *in vivo* in animal models. However, studies on their therapeutic efficacy are still in progress (26).

251 The RWM is considered the main access to the inner ear after the administration in the middle ear (67). NP with different composition and size between 10 and 640 nm were able to cross RWM. The 252 size and the surface charge are determinant factors that affect NP diffusion through RWM. The 253 254 number of NP crossing from middle ear to inner ear was inversely proportional to lipid NP size (39) and in the cochlea the positively charged glycerol mono-oleate NP achieved a larger distribution 255 256 than neutral or negatively charged ones (34). The process responsible for this passage was firstly described for lipid NPs as a paracellular pathway (33). Recent studies in rat RWM suggested that 257 the passage of liposome NP may occur either via the paracellular pathway or by endocytotic 258 259 mechanisms based on clathrin and caveolin (8).

In most studies NP were loaded or labelled with a fluorescent dye (Rhodamine B, Carboxycyanine, Nile-red) or a contrast agent (gadolinium) for visualization of particles in cochlear cells or fluids by imaging techniques: however, the reported data mostly detected the presence of NP in inner ear tissues without a quantitative analysis. Liposome NP were detected in RWM until 11 days and in the cochlea until 6 days post-injection (67); lipid nanocapsules were detected in the cochlea until 7 days post-injection (33). However, for treatment of sensorineural hearing loss, the cochlear target cells are hair cells and spiral ganglion neurons: all populations are selectively reached by the
functionalized NP tested (51). For example, the NP functionalized with nerve growth factor-derived
peptides showed specificity for spiral ganglion neurons and nerve fibres (44).

The ability of NP to carry the drugs into the cochlea through the RWM was shown in vivo by 269 several studies. Smaller supermagnetic iron oxide NP (SPION) coated with PGLA and conjugated 270 with dexamethasone enabled the release of the drug in inner ear fluids, resulting in a higher 271 concentration of dexamethasone in the perilymph compared to the pure drug diffusion (10% higher 272 273 after 60 minutes, p<0.01) (65). The PLGA loaded with coumarin-6 enhanced up to 10.9 times the local bioavailability of the dye in the perilymph in comparison to pure drug solution (50). When the 274 neutoxic agent disulfiram was loaded on liposomes and polymersomes, the number of spiral 275 ganglion cells significantly decreased two days after administration (8). However, drug release by 276 NP has not yet been examined by long-term studies. 277

Biocompatibility is one of the major concerns in NP clinical applications. Up to date no hearing 278 impairment, loss of hair cells or histological damages were reported (31, 32, 45, 68), thus NP 279 systems appear reasonably safe. However, SPION tend to aggregate when the magnetic field is 280 removed and the long persistence of these nanoparticles on the inner ear may induce toxicity due to 281 282 accumulation (8). The effects of LNC were evaluated 20 days post-injection and no toxicity was detected (67). Topic applications of liposomes in rats did not affect hearing, but a NP concentration-283 284 dependent toxicity was observed in vitro in primary cochlear cell cultures (32). A possible 285 explanation was that a NP overload occurred in these cells, resulting in cytoplasm condensation and 286 cell function impairment (69). In most of these studies a single intratympanic administration was employed: recently, in order to improve liposome efficiency, a continuous NP release was obtained 287 through a high-performance polymide tubing (HPPT) equipped with an ALZET<sup>©</sup> micro-pump 288 (DURECT Corp, CA; USA). Liposomes loaded with gadolinium-tetra-azacyclo-dodecane-tetra-289 acetic acid were visualized both in vitro by TEM and in vivo by MRI. In vitro, intact NP were free 290

to diffuse in the medium, and *in vivo* were detected in the cochlea without adverse effects within six
days (67). Again, no long-term effects of exposure to NP after multiple applications in the inner ear
have yet been evaluated.

The residence time of the drug within the middle ear cavity may be increased by NP, but this does 294 not guarantee direct contact between the loaded NP and the RWM, because the RWM is the access 295 point for the inner ear in the case of trans-tympanic administration. The NP also undergo middle ear 296 clearance through the Eustachian tube (70). A possible strategy to bypass these limits could be to 297 298 combine different DD systems together, for example using hydrogels. The incorporation of loaded NP into hydrogels could increase their residence time in the middle ear, thus enhancing drug release 299 in the perilymph (26, 71). The hydrogel is applied near the RWM and releases the loaded NP in the 300 perilymph along with its degradation. This approach has been recently reported: a poloxamer 407 301 hydrogel combined with SPION was successfully applied on ex vivo models (human temporal 302 bones and explanted mouse inner ear cultures) (72). More recently, a nanohydrogel based on 303 chitosan polymer incorporating liposomes was tested in vitro and in vivo in the mouse model (73). 304 The *in vitro* results showed that NP persisted without significant degradation for at least two weeks 305 306 and were released in a controlled and continuous way by the nanohydrogel. The *in vivo* results 307 showed that the NP were successfully released by the nanohydrogel across the RWM and were able to reach the perilymph and Organ of Corti cells (73). 308

Although NP research in local DD for inner ear therapy appears promising, there are still many difficulties to overcome, mostly related to the inner ear anatomy, to the complexity of the cochlea and its highly differentiated cell populations, as well as the possibility to cause hearing loss using microsurgical approaches. The *in vivo* analyses of perilymph samples represent a technical challenge (27), because of the small volume of inner ear fluids in animal models (the total volume of perilymph in a Guinea pig amounts to about 10  $\mu$ l) (74), and the possible contamination of samples by cerebrospinal fluid (27). The pharmacokinetics of drugs in the inner ear is therefore still

unclear and there are no reliable quantitative data about local bioavailability, drug distribution and 316 317 RWM permeability (26). Only recently, a computer pharmacokinetic for inner ear fluids and drug distribution has been developed (75). The software (Cochlear Fluids Simulator V3.083) outlines a 318 model based on inner ear anatomy in humans and rodents, pharmacokinetic and solute distribution 319 parameters. This model has been used to simulate the distribution of therapeutic drugs and other 320 compounds in the perilymph (24, 76). However, because of intraspecific variability of animal 321 models in the volume of inner ear fluids and RWM thickness and conditions, it is difficult to 322 compare the current studies using different DD systems and to draw quantitative conclusions about 323 drug pharmacokinetics in the inner ear. 324

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#### 326 6. Conclusions and Future Perspectives

The NP-based systems show a high potential for inner ear delivery of various therapeutic agents. Their use could minimize the side effects of treatments, allow target specificity and provide a sustained release of drugs in inner ear fluids. The type of NP may be adapted to the drug to be carried and different formulations have been tested. The NP could also be combined with other nanomaterials, such as hydrogels, to improve the local application of drugs. However, several problems have yet to be solved and more *in vivo* studies are necessary to verify their bioavailability and effectiveness before a successful clinical application.

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335

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# INNER EAR NANOPARTICLES DELIVERY



1	Nanoparticle drug delivery systems for inner ear therapy: an overview
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#### 22 Abstract

23 Local drug delivery based on nanoparticles (NP) represents a novel strategy to improve inner ear treatments. The intratympanic delivery of NP may be suitable to treat or prevent hearing loss 24 originating from damage to hair cells and spiral ganglion neurons in the cochlea. Numerous 25 experimental studies support *in vitro* and *in vivo* the biocompatibility of NP, their physical stability, 26 target specificity, cell/tissue uptake and ability to internalize therapeutic agents. The topical use of 27 NP helps to reduce the amount of drug required and avoid systemic side effects. This review 28 focuses on recent findings and applications of different NP systems locally delivered to the inner 29 ear. The perspectives for clinical application of NP in inner ear drug delivery are also discussed. 30

31

#### 32 Keywords

33 Nanoparticles, inner ear, drug delivery, intratympanic administration, local administration

#### 34 **1. Introduction**

The treatment of inner ear diseases through drug delivery (DD) faces numerous challenges (1), 35 among which the limited blood flow to the inner ear (2), the presence of physical barriers acting as 36 a selective filter for drug transportation to the inner ear from the circulatory system (3), the small 37 size of the cochlea and its isolated location in the petrous bone. As a result, research in local drug 38 applications and medications has recently attracted interest because it is a more effective and 39 40 preferable treatment than the systemic one. Case studies involving steroids (4) and gentamicin 41 treatment for Meniere's disease (5) have been documented, but these approaches could be improved for clinical protocols by the development of controlled and targeted delivery systems. 42

Nanoparticles (NP) are a possible option to improve existing therapeutic strategies (6). The NP with
size between 10 and 200 nm are useful for application in biology and medicine for innovative DD
systems. NP-based strategy could be more efficient and reduce drug-associated side effects because

of the ability to deliver the therapeutic agent to the target site. Moreover, the controlled release of
compounds conjugated to NP results in a lower dose of drug required to achieve the therapeutic
effects (1, 7).

The cochlea is a good model for studying the NP-based DD due to its isolated structure and the perilymph rheology. The intratympanic delivery of NP could be suitable to treat the hearing loss and prevent its progression when hair cells and spiral ganglion neurons are damaged (8).

52 Several works and reviews have been published in the past decade, focusing on NP type, pathology 53 involved, delivery approach or a combination of these topics (1-4, 6-8). The goal of the present 54 review is to provide an updated general overview of NP-based strategies and their advantages and 55 disadvantages for local DD into the inner ear.

56

#### 57 **2.** Ear barriers

The human inner ear consists of two main parts, the auditory system (the cochlea) and the vestibular 58 59 system. The cochlea is a bony spiral canal, about 30 mm long and divided in three fluid-filled compartments: the scala tympani, the scala media and the scala vestibuli. The round window 60 membrane (RWM), the blood inner ear barrier (BB) and the oval window are physical barriers that 61 isolate the cochlea from the middle ear and from the circulatory system (Figure 1). The RWM is a 62 63 three-layer semi-permeable membrane, composed of an outer epithelial cell layer, a middle 64 connection layer and an inner connection layer facing the perilymph of the scala tympani (9). In 65 humans, the variable thickness of RWM affects the response of patients to DD treatments. In animal models, its thickness is different among species but its composition is similar (10). 66

Both the RWM and the oval window membranes have been investigated for DD, as connections between the middle ear cavity and the cochlear perilymph. The DD strategies for the inner ear currently rely mostly on RWM (11). The passage of molecules across this membrane is not only

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influenced by thickness, but also by its morphological integrity, inflammation and weight, concentration, liposolubility and external charge of the therapeutic compound (12). The drugs deposited topically in the middle ear cavity are internalized by pinocytosis and transported to the perilymph through blood vessels or by diffusion. Thus the direct application of drugs in the proximity of RWM is a suitable approach for treatment of inner ear pathologies (13).

The BB is a major barrier in the stria vascularis separating the cochlear tissues from the circulatory 75 system (14). Its role is to maintain the homeostasis of cochlear fluids and protect the inner ear 76 77 integrity. Its main components are principally the endothelial capillaries whose cells are connected by tight junctions, which lay over a basement membrane. However numerous accessory cells have 78 recently been observed in the complex structure of the barrier, such as pervcites and perivascular 79 80 resident macrophage-like (11). The BB has been described to act as a physical and biochemical 81 barrier through an efflux pump, the P-glycoprotein 1 (P-gp) (15). The BB is therefore considered a rate-limiting barrier in the passage of therapeutic agents from the circulatory system to the inner ear. 82 83 However, the current knowledge about drug transportation processes through BB is still limited (16). 84

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#### 86 **3. Administration routes**

87 The clinical protocols for inner ear therapies mostly rely on systemic and local DD routes. The systemic administration represents a classical route for DD, but in the inner ear only few drugs may 88 reach the target site at therapeutic concentrations. If high doses of systemic drugs are employed, 89 90 often side effects are developed (17, 18). Systemic applications of NP in inner ear have been recently investigated: poly(lactic-co-glycolic acid) NP conjugated with rhodamine B and applied 91 systemically were detected in the liver, but not in the cochlea (19). The limited bioavailability of NP 92 after systemic administration could be due to the rapid clearance from the circulation in liver and 93 spleen (20). 94

Local administration appears more suitable for inner ear DD (19). This approach allows a quick 95 96 distribution of the drug inside the cochlea, improving their delivery to the target site; it also requires lower drug doses, avoiding side effects (21) (Figure 2). Two main routes are presently used for this 97 purpose, the intratympanic (IT) or the intracochlear administration, but the second one is rarely 98 performed because it is highly invasive and limited to surgery cases (22). On the contrary, the IT 99 injection is minimally invasive and relies on passive diffusion of the active molecules through 100 101 RWM to access the inner ear. This review focuses on development of these methods for DD with minimal trauma for the cochlea. However, local delivery trials show a high variability in results 102 (23) because of some key factors: 1) the drug clearance within the middle ear through the 103 104 Eustachian tube; 2) the permeability of RWM; and 3) the residence time of the drug in contact with RWM (24). A method to reduce variability of results and increase the drug concentration in the 105 perilymph could be to better control the residence time of the drug at close range with RWM, using 106 107 specific delivery systems based on NP (25).

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#### 109 4. Nanoparticle-based systems

The NP (also called nanocarriers or nanovectors) are artificial compounds with size at the 110 111 nanoscale, which aim to compensate for adverse drug properties such as low solubility, degradation and short half-life (26). The NP may also be adapted to target a specific tissue of the inner ear. 112 113 However, when injected in the middle ear as a liquid suspension, NP will undergo clearance 114 through the Eustachian tube (27), thus significantly reducing their residence time near RWM. The NP suitable for DD systems should therefore increase the residence time, together with the ability to 115 cross RWM and their biocompatibility (Figure 3). A detailed description of physico-chemical 116 117 characteristics of NP and their applications is reported.

118 4.1. Lipid Core NP

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Lipid Core NP (LCN) possess a lipid core matrix (usually triglycerides) with a surrounding shell of 119 120 lecithin, polyethylene glycol or poloxamers as stabilizing agents. The LCN structure can be changed to include different drugs and control the kinetics of drug release (28). It has been shown to 121 be stable up to six months in aerosol dispersion (29). These NPs did not induce toxicological effects 122 in vivo in mice after systemic applications (12 mg/kg intravenously for five days) (30) and their cell 123 uptake and cell viability was *in vitro* verified on fibroblasts by confocal scanner laser microscopy 124 125 (31). In rat animal models LCN were able to cross RWM and reach inner ear targets after middle ear application *in vivo*, while not affecting hearing capacity (32). Their preferred pathway to diffuse 126 inside the cells was also investigated: they followed a "nerve pathway", diffusing from the 127 128 perilymph in the scala tympani to the spiral ganglion, nerve fibres and later approaching the inner and the outer hair cells (33). Their variability in diffusion and ability to cross RWM depends on 129 their lipid composition, size and external charge. The ability to cross the RWM has been shown to 130 131 be size-dependent, because the percentage of particle diffusion was inversely proportional to their size (31). Surface charge may also affect the uptake and biodistribution of LNC. Some NP 132 candidates based on glycerol mono-oleate were studied under different external charges: after an in 133 vivo application to RWM, LCN expressing stronger positive charges were detected in the deeper 134 turns of the cochlea (34). The LCN were also tested as a drug carrier, delivering dexamethasone in 135 136 the inner ear through IT injection and comparing the results with a systemic application of the same LCN. The amount of dexamethasone detected in cochlear fluid after local LCN application was 137 significantly higher compared to the systemic application, also increasing the half-life and the 138 average residence time of the drug in the perilymph by 1.9 folds (35). All these results indicate a 139 great potential for LCN for sustained drug release and targeting of inner ear tissues after local 140 administration. 141

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#### 143 **4.2. Liposomes**

Liposomes are artificial phospholipid bilayers, similar to those found in the cell membrane, but 144 145 surrounding an aqueous core. They exhibit a wide size range (between 50 nm and 5 µm) and morphology, depending on the phospholipid used and the preparation method (36). Liposomes can 146 encapsulate either hydrophobic molecules in the phospholipid bilayer or hydrophilic molecules in 147 their aqueous core (37). The uptake of these NP in vitro or in vivo usually relies on the passive 148 diffusion inside the cells, but their surface can be modified with polyethylene glycol, antibodies, 149 150 peptides, carbohydrates, hyaluronic acid and folic acid (35). Such modified liposomes successfully targeted cells expressing tropomyosin receptor-B (TrkB) by using 18-mer peptides to promote 151 cellular uptake (38). Liposomes labelled with fluorescent markers applied in vivo to a mouse model 152 153 with a single IT injection were identified in all cochlear turns, with a concentration gradient decreasing from the base to the apex and, to a lesser extent, in the lateral wall and in the organ of 154 Corti. No morphological or functional damages to the inner ear were detected 24 hours after the 155 156 application (8). Disulfiram, a neurotoxic agent, was used as model payload for DD analysis: NP loaded with Disulfiram damaged the spiral ganglion 48 hours after application, with an associated 157 threshold shift reaching 35 dB. No significant effects were observed with a similar application of a 158 pure Disulfiram solution (8). To test the drug delivery efficiency of liposome nanocarriers, NP of 159 different size (95, 130, 240 nm) encapsulating the contrast agent gadolinium-tetra-azacvclo-160 161 dodecane-tetra-acetic acid (Gd-DOTA) were applied in the middle ear and analyzed with MRI: the results showed that the liposome carrier efficiency was inversely proportional to NP size (39, 40). 162

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## 4.3. Polymersomes and copolymers

164 The polymersomes (also called multifunctional NP) are a wide class of amphiphilic copolymers, 165 consisting of a self-assembled membrane of hydrophobic units, surrounding an aqueous core, and of 166 a hydrophilic corona (41). Structurally they are similar to liposomes, with the advantages that the membrane thickness can be controlled by the molecular weight of the hydrophobic block of 167 copolymer to achieve stronger, thicker and more stable membranes. The hydrophilic corona can be 168

modified to regulate the biodistribution of polymersomes and induce specific cellular uptake (42).
Hydrophilic drugs can be loaded in the core, while hydrophobic ones in the membrane (43).

Different multifunctional polymersomes were studied for inner ear DD targeting specific tissue orconjugated with ferromagnetic materials.

In a mouse model, poly(ethylene glycol)-*b*-poly (ε-caprolactone) NP (PEG-*b*-PCL) labeled with fluorescent markers were detected in the spiral ganglion, in the organ of Corti and in the lateral wall after 24 hours from RWM application *in vivo* (8). Tissue specificity was also investigated: PEG-*b*-PCL were conjugated with a nerve growth factor derived peptide and tested *ex-vivo* on explanted mouse cochleae and *in vitro* on PC12 cells. No significant toxic effect was observed and a specific targeting to spiral ganglion neurons, Schwann cells and nerve fibres was achieved by conjugating the NP with tyrosin kinase and p75 neurotrophin receptors (44).

Poly(2-hydroxyethyl aspartamide) NP (PHEA) were observed to enter in vitro the immortalized 180 mouse organ of Corti cell line (HEI-OC1) and the human middle ear cell line (HMEEC). When 181 applied *in vivo* near the RWM in a mouse model, PHEA were also detected in the inner ear tissue 182 (45). In order to improve NP uptake, PHEA were modified with oligoarginine peptide, a positively 183 charged copolymer, and conjugated with fluorescent Nile red as a hydrophobic model drug (46). In 184 185 these conditions the NP uptake *in vitro* on HEI-OC1 and HMEEC cells was significantly improved 186 after 15 and 24 hours, compared to pure Nile red solution. Modified PHEA were detected after 24 hours from application in the inner hair cells and supporting cells (47). 187

Poly(lactic-co-glycolic acid) (PLGA) NP are copolymers among the novel carrier developed for DD. The Food and Drug Administration (FDA) and the European Medicines Agency (EMA) approved PLGA NP for parenteral administration (47). PLGA NP are interesting because of their hydrophilicity, biocompatibility and easy derivatization by functional groups on the surface or inside the polymer. Their surface may be modified for target specificity by PEGylation, chitosan

absorption and binding of antibodies and oligopeptides (48), and different molecules (proteins, 193 194 steroids, antibiotics and nucleic acids) have been successfully encapsulated and delivered by PLGA NP (49). Programmed degradation of the polymer may therefore yield quantitative delivery of 195 drugs, plasmids or other bioactive molecules. The PLGA NP tested in the inner ear were first 196 conjugated with rhodamine B, a red fluorescent dye, and applied via IT injection: they were 197 identified in the scala tympani, showing that PLGA NP are able to cross RWM by diffusion and 198 199 their clearance depends on the perilymph flow rate (19). A quantitative pharmacokinetic study recently showed that PLGA NP applied locally in vivo in Guinea pigs significantly improved the 200 drug distribution within the inner ear (52). When PLGA NP were loaded with the fluorescent dye 201 202 coumarin-6 and applied through IT injection, the concentration of the compound after 96 hours from treatment was 10.9-fold higher in the perilymph than when administered in pure solution. 203 Similar results were obtained for other therapeutic payloads such as antioxidants and antiapoptotic 204 205 drugs (50). Thus PLGA NP are an useful DD system for inner ear because of their high versatility in adaptation to drug properties and tissue targets (51). 206

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#### 208 4.5. Silica NP

Silica NP are modified colloidal silica particles (52) used to transfect *in vitro* plasmid DNA (53) but also as a DD system (54). A pilot study in mice tested the efficacy of diffusion of Cy3-labeled silica NP administered near the RWM: these NP were found inside the inner hair cells, the vestibular hair cells, the spiral ganglion neurons and the supporting cells, without any hearing impairment. Since the NP also reached the dorsal cochlear nucleus and the superior olivary complex, the authors suggested a retrograde axonal transport and concluded that silica NP could be applied for safe drug deliver in the auditory system (55).

216 4.6. Supermagnetic iron oxide NPs (SPIONs)

Magnetic NP are synthetic  $Fe_3O_4$  (magnetite) particles, with a core diameter around 15 nm, that can 217 218 be widely applied for magnetic targeting of cells (56). Unlike large ferromagnetic materials, the smaller supermagnetic iron oxide NP (SPION) are characterized by the absence of residual 219 magnetic interactions when the magnetic field is not active, thus they are more suitable for 220 biomedical applications (57). The SPION derivatized to increase biocompatibility and cell 221 interactions could be guided by an external magnetic field to a specific biological target, but they 222 223 cannot encapsulate any drug (58). For in vivo applications, to prevent particle aggregation and favour dispersion SPION were coated by organic compounds (59). In inner ear drug delivery, 224 SPION have been encapsulated in PLGA (60), silica (58) and dextran (61) and their 225 226 biocompatibility was tested and verified in vitro and in vivo (59). The mobility of SPION induced by a magnetic field was also quantified and the results of flux density, gradients and NP properties 227 were compared between in vitro and in vivo models (62). The magnetic force required for SPION to 228 229 cross RWM in vivo in Guinea pigs was significantly lower than that of the in vitro RWM model (63). Another study in vivo in Guinea pigs revealed that the concentration of coated SPION inside 230 the cochlea significantly increased (330% above control) when a magnetic field was active (64). 231 Recently, SPION coated with PGLA NP were tested as drug carriers with dexamethasone-acetate 232 233 (Dex-Ac) as a payload: the levels of Dex-Ac detected in the inner ear fluids after 1 hour from 234 treatment were significantly higher compared with those in absence of a magnetic field (65). All these results support the application of SPION for inner ear drug delivery protocols. 235

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#### 237 4.7. Hyperbranched poly-L-lysine NP

Hyperbranched poly-L-lysine (HBPL) are high cationic charged dendrimers widely used for nonviral gene transfer (67, 68). The HBPL were applied *in vivo* in Guinea pig inner ears without any sign of cell toxicity or permanent hearing loss (31): they were detected in the stria vascularis and hair cells (31). Nanoparticles based on HBPL and conjugated with fluorescein isothiocyanate were tested *ex-vivo* on freshly frozen human temporal bones, placing them near the intact RWM: HBPL
were detected in hair cells, nerve fibres and other cochlear tissues (66).

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# 245 5. Key aspects for nanoparticle-based drug delivery in the inner ear

There are several key parameters to consider for NP-based local DD in the inner ear: the RWM permeability; the NP cochlear targeting; their payload ability and the controlled drug release; their biocompatibility and their stability in cochlear fluids and tissues. All these aspects were evaluated with different NP systems *in vitro, ex-vivo* or *in vivo* in animal models. However, studies on their therapeutic efficacy are still in progress (26).

251 The RWM is considered the main access to the inner ear after the administration in the middle ear (67). NP with different composition and size between 10 and 640 nm were able to cross RWM. The 252 size and the surface charge are determinant factors that affect NP diffusion through RWM. The 253 254 number of NP crossing from middle ear to inner ear was inversely proportional to lipid NP size (39) and in the cochlea the positively charged glycerol mono-oleate NP achieved a larger distribution 255 256 than neutral or negatively charged ones (34). The process responsible for this passage was firstly described for lipid NPs as a paracellular pathway (33). Recent studies in rat RWM suggested that 257 the passage of liposome NP may occur either via the paracellular pathway or by endocytotic 258 259 mechanisms based on clathrin and caveolin (8).

In most studies NP were loaded or labelled with a fluorescent dye (Rhodamine B, Carboxycyanine, Nile-red) or a contrast agent (gadolinium) for visualization of particles in cochlear cells or fluids by imaging techniques: however, the reported data mostly detected the presence of NP in inner ear tissues without a quantitative analysis. Liposome NP were detected in RWM until 11 days and in the cochlea until 6 days post-injection (67); lipid nanocapsules were detected in the cochlea until 7 days post-injection (33). However, for treatment of sensorineural hearing loss, the cochlear target cells are hair cells and spiral ganglion neurons: all populations are selectively reached by the
functionalized NP tested (51). For example, the NP functionalized with nerve growth factor-derived
peptides showed specificity for spiral ganglion neurons and nerve fibres (44).

The ability of NP to carry the drugs into the cochlea through the RWM was shown in vivo by 269 several studies. Smaller supermagnetic iron oxide NP (SPION) coated with PGLA and conjugated 270 with dexamethasone enabled the release of the drug in inner ear fluids, resulting in a higher 271 concentration of dexamethasone in the perilymph compared to the pure drug diffusion (10% higher 272 273 after 60 minutes, p<0.01) (65). The PLGA loaded with coumarin-6 enhanced up to 10.9 times the local bioavailability of the dye in the perilymph in comparison to pure drug solution (50). When the 274 neutoxic agent disulfiram was loaded on liposomes and polymersomes, the number of spiral 275 ganglion cells significantly decreased two days after administration (8). However, drug release by 276 NP has not yet been examined by long-term studies. 277

Biocompatibility is one of the major concerns in NP clinical applications. Up to date no hearing 278 impairment, loss of hair cells or histological damages were reported (31, 32, 45, 68), thus NP 279 systems appear reasonably safe. However, SPION tend to aggregate when the magnetic field is 280 removed and the long persistence of these nanoparticles on the inner ear may induce toxicity due to 281 282 accumulation (8). The effects of LNC were evaluated 20 days post-injection and no toxicity was detected (67). Topic applications of liposomes in rats did not affect hearing, but a NP concentration-283 284 dependent toxicity was observed in vitro in primary cochlear cell cultures (32). A possible 285 explanation was that a NP overload occurred in these cells, resulting in cytoplasm condensation and 286 cell function impairment (69). In most of these studies a single intratympanic administration was employed: recently, in order to improve liposome efficiency, a continuous NP release was obtained 287 through a high-performance polymide tubing (HPPT) equipped with an ALZET<sup>©</sup> micro-pump 288 (DURECT Corp, CA; USA). Liposomes loaded with gadolinium-tetra-azacyclo-dodecane-tetra-289 acetic acid were visualized both in vitro by TEM and in vivo by MRI. In vitro, intact NP were free 290

to diffuse in the medium, and *in vivo* were detected in the cochlea without adverse effects within six
days (67). Again, no long-term effects of exposure to NP after multiple applications in the inner ear
have yet been evaluated.

The residence time of the drug within the middle ear cavity may be increased by NP, but this does 294 not guarantee direct contact between the loaded NP and the RWM, because the RWM is the access 295 point for the inner ear in the case of trans-tympanic administration. The NP also undergo middle ear 296 clearance through the Eustachian tube (70). A possible strategy to bypass these limits could be to 297 298 combine different DD systems together, for example using hydrogels. The incorporation of loaded NP into hydrogels could increase their residence time in the middle ear, thus enhancing drug release 299 in the perilymph (26, 71). The hydrogel is applied near the RWM and releases the loaded NP in the 300 perilymph along with its degradation. This approach has been recently reported: a poloxamer 407 301 hydrogel combined with SPION was successfully applied on ex vivo models (human temporal 302 bones and explanted mouse inner ear cultures) (72). More recently, a nanohydrogel based on 303 chitosan polymer incorporating liposomes was tested in vitro and in vivo in the mouse model (73). 304 The *in vitro* results showed that NP persisted without significant degradation for at least two weeks 305 306 and were released in a controlled and continuous way by the nanohydrogel. The *in vivo* results 307 showed that the NP were successfully released by the nanohydrogel across the RWM and were able to reach the perilymph and Organ of Corti cells (73). 308

Although NP research in local DD for inner ear therapy appears promising, there are still many difficulties to overcome, mostly related to the inner ear anatomy, to the complexity of the cochlea and its highly differentiated cell populations, as well as the possibility to cause hearing loss using microsurgical approaches. The *in vivo* analyses of perilymph samples represent a technical challenge (27), because of the small volume of inner ear fluids in animal models (the total volume of perilymph in a Guinea pig amounts to about 10  $\mu$ l) (74), and the possible contamination of samples by cerebrospinal fluid (27). The pharmacokinetics of drugs in the inner ear is therefore still

unclear and there are no reliable quantitative data about local bioavailability, drug distribution and 316 317 RWM permeability (26). Only recently, a computer pharmacokinetic for inner ear fluids and drug distribution has been developed (75). The software (Cochlear Fluids Simulator V3.083) outlines a 318 model based on inner ear anatomy in humans and rodents, pharmacokinetic and solute distribution 319 parameters. This model has been used to simulate the distribution of therapeutic drugs and other 320 compounds in the perilymph (24, 76). However, because of intraspecific variability of animal 321 models in the volume of inner ear fluids and RWM thickness and conditions, it is difficult to 322 compare the current studies using different DD systems and to draw quantitative conclusions about 323 drug pharmacokinetics in the inner ear. 324

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#### 326 6. Conclusions and Future Perspectives

The NP-based systems show a high potential for inner ear delivery of various therapeutic agents. Their use could minimize the side effects of treatments, allow target specificity and provide a sustained release of drugs in inner ear fluids. The type of NP may be adapted to the drug to be carried and different formulations have been tested. The NP could also be combined with other nanomaterials, such as hydrogels, to improve the local application of drugs. However, several problems have yet to be solved and more *in vivo* studies are necessary to verify their bioavailability and effectiveness before a successful clinical application.

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# Figure Legends

Fig. 1. Scheme of the cochlea structure, highlighting the cochlear barriers, the round window and the stria vascularis.

Fig. 2. Scheme of nanoparticle administration routes. Arrow: intratympanic route; black arrowhead: intracochlear route by round window; white arrowhead: intracochlear route by cochleostomy.

Fig. 3. Structures of nanoparticles (NP) useful for drug delivery. LCN: lipid core NP; SPION: supermagnetic iron oxide NP; HBPL: hyperbranched poly-L-lysine NP; P: hydrophilic region; NP: hydrophobic region.





