Ultrastructure of the Odontocete Organ of Corti: Scanning and Transmission Electron Microscopy

Maria Morell,1,2 Marc Lenoir,3 Robert E. Shadwick,2 Thierry Jauniaux,4 Willy Dabin,5 Lineke Begeman,6 Marisa Ferreira,7 Ianruz Maestre,8 Eduard Degollada,9 Gema Hernandez-Milian,10 Chantal Cazevieille,11 José-Manuel Fortuño,12 Wayne Vogl,13 Jean-Luc Puel,3 and Michel Andrè1*

1Laboratory of Applied Bio-Acoustics, Technological Center of Vilanova i la Geltrú, Technical University of Catalonia-Barcelona Tech, 08800 Vilanova i la Geltrú, Barcelona, Spain
2Zoology Department, The University of British Columbia, V6T 1Z4 Vancouver, Canada
3National Institute of Health and Medical Research U1051, Institute Neurosciences Montpellier, Saint Eloi Hospital, 34091 Montpellier cedex 5, France
4Department of Veterinary Pathology, University of Liège, 4000 Liège, Belgium
5Center for Marine Mammal Research, University of La Rochelle, 17000 La Rochelle, France
6Department of Pathobiology, Faculty of Veterinary Medicine, Utrecht University, 3508 TD Utrecht, The Netherlands
7Portuguese Wildlife Society/Center of Molecular and Environmental Biology, Department of Biology, University of Minho, Gualtar Campus, 4710-057 Braga, Portugal
8AMBAR, Society for the Study and Conservation of Marine Fauna, 48014 Bilbao, Spain
9EDMAKTUB Association for the Study and Disclosure of Aquatic Environment, 08034 Barcelona, Spain
10School of Biological, Earth and Environmental Sciences, University College Cork, Cork, Ireland
11Regional Center for Cellular Imaging Resources, 34091 Montpellier cedex 5, France
12Institute of Marine Sciences, Spanish National Research Council, 08003 Barcelona, Spain
13Department of Cellular and Physiological Sciences, The University of British Columbia, V6T 1Z4 Vancouver, Canada

ABSTRACT

The morphological study of the Odontocete organ of Corti, together with possible alterations associated with damage from sound exposure, represents a key conservation approach to assess the effects of acoustic pollution on marine ecosystems. By collaborating with stranding networks from several European countries, 150 ears from 13 species of Odontocetes were collected and analyzed by scanning (SEM) and transmission (TEM) electron microscopy. Based on our analyses, we first describe and compare Odontocete cochlear structures and then propose a diagnostic method to identify inner ear alterations in stranded individuals. The two species analyzed by TEM (Phocoena phocoena and Stenella coeruleoalba) showed morphological characteristics in the lower basal turn of high-frequency hearing species. Among other striking features, outer hair cell bodies were extremely small and were strongly attached to Deiters cells. Such morphological characteristics, shared with horseshoe bats, suggest that there has been convergent evolution of sound reception mechanisms among echolocating species. Despite possible autolytic artifacts due to technical and experimental constraints, the SEM analysis allowed us to detect the presence of scarring processes resulting from the disappearance of outer hair cells from the epithelium. In addition, in contrast to the rapid decomposition process of the sensory epithelium after death (especially of the inner hair cells), the tectorial membrane appeared to be more resistant to postmortem autolysis effects. Analysis of the stereocilia imprint pattern at the undersurface of the tectorial membrane may provide a way to detect possible ultrastructural alterations of the hair cell stereocilia by mirroring them on the tectorial membrane. J. Comp. Neurol. 000:000–000, 2014.

© 2014 Wiley Periodicals, Inc.

INDEXING TERMS: inner ear; cochlea; cetacean; morphology; acoustic trauma

Grant sponsor: the Spanish Ministry of the Environment; Grant number: 083/SDGTB/2007; Grant sponsor: Agency for Administration of University and Research Grants (AGAUR; Generalitat of Catalonia) fellowships; Grant sponsor: Canadian Natural Sciences and Engineering Research Council Discovery and Accelerator grants.

*CORRESPONDENCE TO: Michel André, Avda. Rambla exposició s/n 08800-Vilanova i la Geltrú, Barcelona, Spain. E-mail: michel.andre@upc.edu

© Wiley Periodicals, Inc.
The ecological and physiological significance of noise effects on toothed whales (order Cetacea, suborder Odontoceti) has recently been recognized (Richardson et al., 1995) because of their vital dependence on acoustic information and their role as apex marine predators. Although some of the physiological damage caused by sound has been observed in organs not directly related to the acoustic pathways (Jepson et al., 2003), most lesions are expected to affect hearing, particularly the organ of Corti and its associated hair cells (Lurie et al., 1944).

In previous studies on terrestrial mammals, structural alterations of the organ of Corti have been found as a consequence of sound exposure. These consist of: 1) changes in the hair cell stereocilia (mainly fusion, size reduction, buckling, formation of “giant” hairs, and disappearance of hairs; Bredberg et al., 1972; Engstrom et al., 1984); 2) degeneration of sensory cells (karyorrhexis, karyopyknosis, and necroptosis; Hu et al., 2000); 3) increased numbers of synaptic vesicles in medial efferent nerve endings and swelling of the afferent nerve endings on the inner hair cells (IHCs) with incipient retrograde nerve degeneration (Spoendlin, 1971); and in more severe cases, 4) complete degeneration of the organ of Corti (Bredberg et al., 1972). At the level of the tectorial membrane, morphological changes on the outer hair cell (OHC) stereocilia imprints have also been shown (Morisaki et al., 1991) after acoustic overstimulation. These changes are coincident with a transformation from circular to oval or irregular shape, fusion of adjacent concavities to form a larger concavity, and occasionally the appearance of filamentous material. Such modification of the imprints has remained for as long as a year, after sensory hairs have disappeared.

Cetaceans include the suborder odontocetes, or toothed whales, comprising approximately 80 different species. Their adaptation to the marine environment has led to the evolution of an echolocation system (Au, 1993; Thomson and Richardson, 1995; Ketten, 2000) that operates at very high frequencies (20–180 kHz), which they use for orientation, foraging, and probably communication among members of the same social group.

Previous morphological studies on the odontocete inner ear by light microscopy can be found in the literature. A detailed description of cochlear morphology was conducted for the bottlenose dolphin (Wever et al., 1971a–c) and the Pacific white-sided dolphin (Wever et al., 1972). Also, cochlear fibers in diverse species (Gao and Zhou, 1992), as well as features of the basilar membrane and osseous spiral lamina in different odontocete species have been compared and related to their hearing capa-

bilities (Ketten and Wartzok, 1990; Ketten, 1992, 1994, 2000). However, ultrastructural descriptions of hearing structures in odontocete species using electron microscopy are still lacking, primarily due to the difficulty of obtaining suitable quality material and reliable protocols for analysis.

Detailed characterizations of the organ of Corti in echolocating horseshoe bats (Rhinolophus rouxi) using scanning (SEM) and transmission (TEM) electron microscopy have been published (Vater and Lenoir, 1992; Vater et al., 1992; Vater and Kossel, 2011). Special features of the very high frequency hearing species were described, such as the very robust attachment between the OHC cuticular plates and the Deiters cells, and the very well developed cup formation of the Deiters cell bodies that house the bottom of the OHC. Because of the similar functionality of the biosonar in bats, we focused on the same hearing structures in odontocetes to investigate possible morphological similarities among these echolocating species, using both SEM and TEM.

The objectives of this study were 1) to describe structural features of the organ of Corti in several odontocete species that can be further correlated with their respective hearing sensitivities; and 2) to conduct a morphological analysis of the cochlea to investigate possible structural alterations as a consequence of sound exposure.

MATERIALS AND METHODS

Extraction and fixation

A total of 150 ears from 13 odontocete species (Pho-

coena phocoena, n = 71; Stenella coeruleoalba, n = 30; Stenella frontalis, n = 12; Tursiops truncatus n = 9; Delphinus delphis, n = 11; Kogia simus, n = 2; Kogia brevi-

ceps, n = 3; Globicephala macrorhynchus, n = 1; Globicephala melas, n = 3; Steno bredanensis; n = 2; Lag

enodelphis hosei, n = 1; Hyperoodon ampullatus, n = 2; and Ziphius cavirostris, n = 2) that stranded in the Mediterranean Sea, Spanish North Atlantic, and North Sea were extracted following the protocol presented at the European Cetacean Society Conference in Istanbul (Morell and André, 2009), and adopted at the Necropsy Workshop, Liège, 2009 (see Table 1 for more details).

After extraction, the samples were fixed with 10% buffered formalin or 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 7.3–7.4). Thirty-six were perfused through the oval and round window replacing the perilymph with the fixative solution.

Decalcification

Because cetacean cochleas are surrounded by very dense periosteal bone, it is necessary to decalcify the
samples to gain access to inner ear structures. Although different solutions were tested (Table 1), most samples were decalcified by using the commercial agent RDO® (Apex Engineering Products, Aurora, IL) as described in Morell et al. (2009). RDO is a rapid decalciﬁer based on hydrochloric acid. Depending on the tympanic–periotic complex volume (Morell et al., 2007), it can take from a few hours to a few days to decalcify the bone tissue. Better control of the decalcification endpoint was obtained by using 50% RDO (diluted in distilled water) and changing to 25% RDO after 24 hours (Morell et al., 2009). Other samples were decalcified by using 14% ethylenediaminetetraacetic acid (EDTA) tetrasodium salt at pH 7.4 and 7.6 either at room temperature (changing the solution once per week; Callis and Sterchi, 1998) or in a microwave oven; SEM, scanning electron microscope; TEM, transmission electron microscope.

Transmission electron microscopy

Only the freshest samples that were perfused with 2.5% glutaraldehyde with 0.1 M phosphate buffer were used for morphological description by TEM. These were completely decalcified, and the endpoint was determined by using the weight loss/weight gain procedure (Mawhinney et al., 1984; Hornbeck et al., 1986; Sanderson et al., 1995) and X-ray imaging. Speciﬁcally, the TEM samples presented here were taken from: 1) an adult striped dolphin (S. coeruleoalbica) ear recovered in the North of Spain, ﬁxed for 5 hours and 30 minutes postmortem and perfused with the same ﬁxative solution 9 hours postmortem 2) a juvenile harbor porpoise (P. phocoena) stranded on the Belgian coast, perfused with ﬁxative 22 hours and 10 minutes postmortem 3) an adult harbor porpoise perfused for at least 3 hours and 15 minutes postmortem 4) a neonate harbor porpoise perfused for less than 4 hours postmortem and 5) a juvenile harbor porpoise perfused for less than 12 hours postmortem (Table 1). Individuals 3, 4, and 5 were stranded on The Netherlands coast.

The samples from individuals 1–3 were postﬁxed for 1 hour with buffered 2% osmium tetroxide, dehydrated with increasing concentrations of ethanol, and embedded in EPON resin by using an automatic microwave tissue processor (Leica AM AMW) for electron microscopy. Semithin sections were examined under a light microscope and ultrathin sections under the Hitachi H-7100 TEM (at CRIC, Montpellier, France).

The samples from individuals 4 and 5 were postﬁxed with 2% osmium tetroxide, dehydrated with increasing concentrations of ethanol, and then embedded in SPURR resin using a Pelco (Ted Pella, Redding, CA) microwave. Semithin and ultrathin sections were cut, and the latter were examined by using the Hitachi H7600 TEM (University of British Columbia Bioimaging Facility, Vancouver, BC, Canada) and the FEI Tecnai G2 Spirit (Department of Cellular and Physiological Sciences, University of British Columbia).

Scanning electron microscopy

The decalcification of the periotic bone was stopped when the vestibular scalae and the stria vascularis of the cochlea were uncovered. A total of 64 cochleas and their tectorial membranes were dissected, dehydrated through ethanol, critical point dried with CO₂,
and then coated with gold–palladium (Table 1). The samples were observed by SEM for morphological description and for possible acoustic trauma assessment. Imaging was done on a Hitachi S-3500N (at the Institute of Marine Sciences, Spanish National Research Council, Barcelona, Spain), a Hitachi S-4000 (at CRIC, Montpellier, France), a Jeol JSM 5600 (at the Technical Center of Vilanova i la Geltrú, Spain), a Hitachi S-570 (at the Autonomous University of Barcelona, Spain), and a Hitachi S-4700 (University of British Columbia Bioimaging Facility).

**Image processing**

Brightness and contrast of images were adjusted in Adobe (San Jose, CA) Photoshop CS3. Several dimensions in the cochlear structures were measured (see F1 Fig. 1 for details), either by using specific tools in the electron microscope software or in ImageJ.

### RESULTS

This section presents the TEM and SEM analysis of the general morphology of the organ of Corti. This structure was typically formed by one row of IHCs and three rows of OHCs. In a harbor porpoise, some additional OHCs were observed forming a fourth row in the lower turn.

**Transmission electron microscopy**

The average cochlear dimensions based on TEM and light microscopy analysis are presented in Tables 2 and 3. Figure 2 shows reconstructions of the spiral course of the basilar membrane performed with Adobe Illustrator from SEM pictures, and indicates the positions that were analyzed by TEM. The most interesting features of each structure are presented below.

#### Basilar membrane

The basilar membrane (Fig. 1A,B) increased in width (i.e., the distance between the foramen of the habenula perforata and the spiral ligament) and decreased in thickness toward the apex, being especially narrow and thick in the lower basal turn. In striped dolphin, the basilar membrane ranged from an average of 112 to 377 μm in width and 14 to 5 μm in thickness, while in harbor porpoise the average values ranged from about 83 to 394 μm in width and 12 to 7 μm in thickness (Table 2). Contrary to this general trend, in the harbor porpoise the basilar membrane width ranged from 412 μm in the apex to 53 μm at approximately 3 mm from the base, but increased again to 118 μm the first 2 mm. There was no information about the most basal region of the cochlea in the striped dolphin (Fig. 2B).

#### Spiral ligament

The spiral ligament was large in the lower basal turn (maximum of 359 μm in striped dolphin and 488 μm in harbor porpoise; Fig.1A). Interestingly, it persisted up to the apical turn in both species.

#### Sensory cells: outer hair cells.

The ultrastructure of the OHCs is illustrated in Figure 3. Measurements of cell body length in over 60 OHCs showed that OHCs of the lower basal turn (Fig. 3A) were very small (striped dolphin: 9.96 μm; harbor

---

**TABLE 2.**

Mean Measurements (in μm) Using Transmission Electron Microscopy and Light Microscopy of the Outer Hair Cells (OHCs), Deiters Cells, Basilar Membrane, and Spiral Ligament of Different Bassoapical Locations

<table>
<thead>
<tr>
<th></th>
<th>OHCs max. length</th>
<th>OHCs max. diameter</th>
<th>OHC cuticular plate Arms</th>
<th>Thickness</th>
<th>First row of the OHC stereocilia Length</th>
<th>Diameter</th>
<th>Rootlet Width</th>
<th>Basilar membrane Thickness</th>
<th>Deiters cells Arm Thickness</th>
<th>Deiters cells Cup Length</th>
<th>Cup length</th>
</tr>
</thead>
<tbody>
<tr>
<td>Striped dolphin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower basal turn</td>
<td>9.97</td>
<td>5.17</td>
<td>2.47</td>
<td>0.93</td>
<td>117.86/112.06</td>
<td>13.95/12.98</td>
<td>336.84/253.45</td>
<td>2.21</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upper basal turn</td>
<td>5.42</td>
<td></td>
<td></td>
<td></td>
<td>199.48/148.16</td>
<td>11.91/11.56</td>
<td>285.71/27.57</td>
<td>3.13</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower apical turn</td>
<td>14.90</td>
<td>6.25</td>
<td>2.65</td>
<td>1.48</td>
<td>0.97/0.17</td>
<td>0.65</td>
<td>220.05/205.80</td>
<td>2.17</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apical end</td>
<td>22.10</td>
<td>13.06</td>
<td>1.77</td>
<td>2.71</td>
<td>1.78/0.29</td>
<td>0.66</td>
<td>377.03/374.35</td>
<td>8.63/2.47</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Harbor porpoise</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower basal turn</td>
<td>10.36</td>
<td>4.10</td>
<td>3.40</td>
<td>1.36</td>
<td>0.77/0.21</td>
<td>0.87</td>
<td>83.79/83.49</td>
<td>11.69/11.56</td>
<td>374.35/27.57</td>
<td>4.00/1.2</td>
<td></td>
</tr>
<tr>
<td>Upper basal turn</td>
<td>10.19</td>
<td>4.60</td>
<td>2.64</td>
<td>1.26</td>
<td>0.46/0.15</td>
<td>0.78</td>
<td>134.25/116.19</td>
<td>8.41/8.90</td>
<td>223.16/32.35</td>
<td>2.47/0.86</td>
<td></td>
</tr>
<tr>
<td>Lower apical turn</td>
<td>14.79</td>
<td>6.90</td>
<td>2.00</td>
<td>1.68</td>
<td>0.21/0.75</td>
<td>0.75</td>
<td>209.49/198.72</td>
<td>7.91/1.74</td>
<td>174.33/37.71</td>
<td>3.9/1.2</td>
<td></td>
</tr>
<tr>
<td>Upper apical turn</td>
<td>18.48</td>
<td>7.07</td>
<td>1.52</td>
<td>1.28</td>
<td>1.85/0.22</td>
<td>0.44</td>
<td>393.95/337.99</td>
<td>7.19/9.2</td>
<td>60.27/12.07</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Max. SD</td>
<td>1.89</td>
<td>2.73</td>
<td>1.38</td>
<td>0.38</td>
<td>0.30/0.07</td>
<td>0.24</td>
<td>23.70/22.75</td>
<td>1.55/2.63</td>
<td>54.8/3.50</td>
<td>3.50/1.08</td>
<td></td>
</tr>
</tbody>
</table>

1See Figure 1A–C for more details on the measurements.
2Measurements performed with light microscopy.
3Measurement that presents the maximum standard deviation (max. SD).

---

M. Morell et al. | The Journal of Comparative Neurology | Research in Systems Neuroscience

Figure 1. Scheme of the cochlear measurements performed using light microscopy (A,B), transmission electron microscopy (C), and scanning electron microscopy (D). These measurements are detailed in Table 2 (a–l) and Table 4 (n–z). A,B: Transverse section of the upper basal turn of the cochlea of a harbor porpoise stained with osmium tetroxide. C: Schematic representation of an OHC at the base of the cochlea and the apical part of the Deiters cell body. D: Schematic representation of the reticular lamina of the organ of Corti epithelium. IHC, inner hair cell; OHC, outer hair cell; IPC, inner pillar cell; OPC, outer pillar cell; PC, phalangeal cell. Scale bars: A) 200 µm, B) 100 µm.
porpoise: 10.36 μm, with extreme values of 7.8 μm). They reached a maximal length of 22.1 μm in the apical turn of the striped dolphin and 18.47 in the upper apical turn of the harbor porpoise. The OHCs were also very narrow, increasing in diameter while approaching the apex of the cochlea (from 5.2 to 13.1 μm in striped dolphin and from 4 to 7.1 μm in the lower apical turn in harbor porpoise).

Stereocilia and cuticular plates
Stereocilia were not very well preserved, and only partial data could be extracted. They were generally of very short length, being longer in the apical cochlear end (1.8 μm in striped dolphin and harbor porpoise) than in the lower basal turn (0.8 μm in harbor porpoise). OHCs had three rows of stereocilia. Cuticular plates of OHCs established tight junctions with the outer pillar cells and phalangeal processes of the three rows of Deiters cells. These attachment zones were extremely long in the OHCs of the basal turn, forming two “arms” (asterisks in Fig. 3A,C).

Subsurface cisternae
OHCs presented a single layer of subsurface cisternae (white arrows in Fig. 3F), which started just below the cuticular plate and continued down to the beginning of the synaptic region, below the nucleus.

Sensory cells: inner hair cells
IHCs were very poorly preserved, and were only observed in the lower apical turn in the striped dolphin and in the lower basal turn in the harbor porpoise. They presented the typical pear shape (Fig. 3E) with a cuticular plate thickness of 1.57 μm. The tallest stereocilia of these cells were 2.97 μm in length and 308 nm in diameter.

Supporting cells
Pillar and Deiters cells, especially at the base of the cochlea, presented an extremely well-developed microtubular cytoskeleton (Fig. 3A,G,H). The phalangeal process of Deiters cells and outer pillar cells had exaggerated long contact zones with the OHCs. This feature was especially evident on the basal turn of odontocete cochlea (asterisks in Fig. 3A,C).

The lower parts of the OHC bodies were completely surrounded by a large cup formed by the Deiters cells. The ultrastructure of the Deiters cup attachment is shown for the lower basal turn in harbor porpoises (black arrows in Fig. 3A). This feature was not observed in apical locations as is the case in other mammalian species like rats (Fig. 3B, courtesy of Marc Lenoir). In

### TABLE 3.

<table>
<thead>
<tr>
<th>Neuron Type</th>
<th>Max. SD</th>
<th>Density (cells/10,000 μm²)</th>
<th>Max. length (μm)</th>
<th>Min. length (μm)</th>
<th>Thickness of myelin layers (nm)</th>
<th>Diameter of afferent fibers before habenula (nm)</th>
<th>Diameter of nerve fibers of the spiral lamina (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spiral ganglion cells</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Striped dolphin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower basal turn</td>
<td>4.56</td>
<td>4.60</td>
<td>54.97</td>
<td>23.21</td>
<td>22.48</td>
<td>601</td>
<td>39.00</td>
</tr>
<tr>
<td>Upper basal turn</td>
<td>4.43</td>
<td>4.01</td>
<td>46.18</td>
<td>29.75</td>
<td>22.23</td>
<td>403</td>
<td>24.00</td>
</tr>
<tr>
<td>Lower apical turn</td>
<td>5.77</td>
<td>6.90</td>
<td>44.99</td>
<td>38.38</td>
<td>25.14</td>
<td>439</td>
<td>34.81</td>
</tr>
<tr>
<td>Apical end</td>
<td>5.13</td>
<td>6.80</td>
<td>37.83</td>
<td>42.23</td>
<td>23.69</td>
<td>443</td>
<td>39.99</td>
</tr>
<tr>
<td>Harbor porpoise</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower basal turn</td>
<td>7.59</td>
<td>8.08</td>
<td>45.97</td>
<td>32.95</td>
<td>29.75</td>
<td>380</td>
<td>30.00</td>
</tr>
<tr>
<td>Upper basal turn</td>
<td>7.13</td>
<td>7.59</td>
<td>44.99</td>
<td>29.75</td>
<td>22.23</td>
<td>439</td>
<td>34.81</td>
</tr>
<tr>
<td>Lower apical turn</td>
<td>7.13</td>
<td>7.59</td>
<td>45.97</td>
<td>32.95</td>
<td>29.75</td>
<td>380</td>
<td>30.00</td>
</tr>
<tr>
<td>Apical end</td>
<td>7.13</td>
<td>7.59</td>
<td>45.97</td>
<td>32.95</td>
<td>29.75</td>
<td>380</td>
<td>30.00</td>
</tr>
</tbody>
</table>

1Measured by light microscopy.
2Measurement that presents the maximum standard deviation (max. SD).
the lower basal turn of the cochlea, this cup reached up to 45% of the total length of the OHCs.

**Innervation**

Along the entire cochlea of both odontocete species that we analyzed, the OHCs were contacted by nerve fibers, which passed between the Deiters cells. These fibers were clearly identified as afferent fibers by the abundant microtubule content of their cytoplasm (Fig. 4C). In contrast, there were no signs of either vesiculated terminals below OHCs or fibers crossing the upper middle region of the tunnel of Corti, belonging to the medial efferent system, in any of the cochlear regions. Additionally, there were no remains of postsynaptic cisterns in the basal pole of the OHCs or medial efferent terminals (Fig. 4A).

The nerve fibers of the inner spiral bundle (below IHCs) showed many postmortem changes (Fig. 3E), with complete loss of nerve terminals. Morphometric information was extracted on the nerve fibers passing in the habenula and spiral lamina, including the diameter of the fibers, the thickness of the myelin sheath, the number of myelin layers, and the number of rows of nerve fibers (Table 3).

The density of spiral ganglion cells (SGCs) was estimated by manually counting the SGCs from many transverse sections of different regions of the cochlear spiral of known area, using light micrographs. The SGC density was expressed as the number of cells per 10,000 μm² in Rosenthal’s canal. The data in Table 3 suggest an increased density toward the apex, but when all data were considered (not shown here) high variability masked any clear tendency. However, the density of SGC in harbor porpoises (from 6.8 to 8.1 SGCs/10,000 μm²) was systematically higher than in striped dolphin (from 4.4 to 5.8 SGCs/10,000 μm²).

The SGCs were well preserved (Fig. 4D) and very large in size in both species. In the striped dolphin, they ranged from 37 to 46 μm at their longest axis, and from 22.4 to 29.7 μm at their shortest axis. By comparison, these dimensions measured 34.8–45 μm and 22.2–25 μm, respectively, in the harbor porpoise.

**Scanning electron microscopy**

Surprisingly, the best preserved regions observed by SEM in all cochleas were the apical end, apical turn, and, in some cases, part of the upper basal turn. Consequently, we present all results from these particular locations. There were no significant differences among the measurements of the apical poles of the sensory and supporting cells within the different measured positions; thus, values in Table 4 are averaged.

**Sensory cells: outer hair cells**

The OHCs had narrow, wing-shaped cuticular plates that ranged in length from 5.0 to 6.2 μm (Fig. 5A,B) and were separated from each other by 500–880 nm within a given row (Table 4). The innermost row (OHC1) and the middle row (OHC2) were distinctly separated by the large hexagonal heads of the outer pillar cells. In...
Figure 3. Ultrastructure of outer hair cells and Deiters cells of the lower basal turn of a harbor porpoise (A,C,E–H), the lower apical turn of a striped dolphin (D), and the apical turn of a rat (B; courtesy of Marc Lenoir) using TEM. Black arrows in A: Deiters cell cup attachment with the lower bodies of OHCs. Asterisks in A and C: large zone of contact between OHCs cuticular plate and supporting cells phalangeal process. Outer hair cell subsurface cisternae (white arrows in F), inner hair cell (E), and the basal body cell of the inner pillar cell (G). BC, border cell; DC, Deiters cell; IHC, inner hair cell; IPC, inner pillar cell; ISB, inner spiral bundle; OHC, outer hair cell. Scale bar = 2 μm in A,G,H; 5 μm in B; 500 nm in C–F.
contrast, there was very little space between the middle (OHC2) and the outermost (OHC3) rows of OHCs, which were significantly different in both harbor porpoise ($P < 0.001$) and striped dolphin ($P < 0.001$).

The OHCs appeared to have three rows of stereocilia (Fig. 5B), but more studies with better preserved samples should be conducted to confirm this result. Lateral links between stereocilia were observed in some cases (Fig. 5C). A few cells forming a fourth row of OHCs in the apical region (Fig. 6B) were only found in one harbor porpoise.

The total number of OHCs was estimated taking into account their average density (175.37 OHCs mm$^{-2}$ of cochlea for striped dolphin and 147.67 OHCs mm$^{-2}$ for harbor porpoise), and extrapolating it to all averaged cochlear lengths (26.878 mm for striped dolphin and 24.813 mm for harbor porpoise). This resulted in a total of 15,012 and 11,405 OHCs, respectively.

The cochlear length was calculated from flat SEM images, performed along the limit of the first row of OHCs with the inner pillar cells. The cochlear length measurements of juvenile individuals were not taken into account for this study.

Even though the samples were not well preserved, it was possible to determine the shape of OHC cuticular plates. It was also possible to determine whether scars were present resulting from the growth of supporting cells after the death of OHCs (Fig. 5D; see a schematic of the scar formation process in Fig. 5F). Moreover, in the case of one of the harbor porpoises, around 420 μm of the apical portion of the cochlea was full of scars (Fig. 5E).

**Sensory cells: inner hair cells**

The IHCs were rarely well preserved in any of the samples analyzed. The IHC cuticular plates were of oval shape (Fig. 6A). The average maximum length in striped dolphin IHC was 8.4 μm, and 4.5 μm was the minimum average length. In the harbor porpoise, the lengths were 8.2 and 5.2 μm, respectively, and the IHCs were found to be separated by 1.15 μm; this latter distance...
was observed to be larger than the separation distance of OHCs in the same row.

The total number of IHCs was estimated following the same procedure as explained for the OHCs, resulting in a total of 3,127 IHCs in harbor porpoise.

**Tectorial membrane**

Despite the poor preservation of the organ of Corti, the tectorial membrane was relatively well preserved. Figure 7 shows the imprints of stereocilia that were found in the tectorial membrane of different species. Even in a sample of one of the bottlenose whales (*Hyperoodon ampullatus*) where there were no remains of the organ of Corti present in the sample, the imprints of the OHC stereocilia were observable in the tectorial membrane (Fig. 7C). By counting the number of imprints, it was possible to estimate the number of the longer stereocilia per OHC, and, assuming that each OHC contained three rows of stereocilia (Fig. 5B), it was possible to estimate the total number of stereocilia. This estimate ranged from 130 to 168, depending on the species (Table 5).

Impressions of the IHC stereocilia were detected in the very basal partition of the tectorial membrane (arrows in Fig. 7G,H) in one of the common dolphins.

---

**TABLE 4.**

Mean Morphometric Measurements and Their Standard Deviations (σ) of the Reticular Lamina 1

<table>
<thead>
<tr>
<th>Striped dolphin</th>
<th>Harbor porpoise</th>
</tr>
</thead>
<tbody>
<tr>
<td>IHC/mm</td>
<td>15.012.02</td>
</tr>
<tr>
<td>Total no. of OHCs</td>
<td>121.46</td>
</tr>
<tr>
<td>IPC/mm</td>
<td>4.41</td>
</tr>
<tr>
<td>OHC/mm</td>
<td>17.14</td>
</tr>
<tr>
<td>Total no. of OHCs</td>
<td>3,126.96</td>
</tr>
<tr>
<td>OPC/mm</td>
<td>7.37</td>
</tr>
<tr>
<td>Distance between OHCs</td>
<td>4.34</td>
</tr>
<tr>
<td>Max. length</td>
<td>10.72</td>
</tr>
<tr>
<td>Min. length</td>
<td>4.41</td>
</tr>
</tbody>
</table>

---

**FIGURE 7.**

The images were obtained by using scanning electron microscopy from the lower and upper apical turn, apical end, and, in some cases, part of the upper basal turn. See Figure 1D for more details on the

---

**FIGURE 5.**

The images were obtained by using scanning electron microscopy from the lower and upper apical turn, apical end, and, in some cases, part of the upper basal turn. See Figure 1D for more details on the
Light microscope measurements revealed that OHC length ranged from 8 to 17 μm in the bottlenose dolphin (Wever et al., 1971a), from 10 to 22 μm in the striped dolphin, and from 8 to 18.5 μm in the harbor porpoise (present study).

A linear relationship exists between frequency and OHC length in terrestrial mammals (Pujol et al., 1991; Dannhof et al., 1991). Indeed, the length of the OHCs increases while the apex is approached. Because the absolute length of OHCs may play a role in an intrinsic

**Figure 5.** Scanning electron microscopic images of the apical pole of outer hair cells (OHCs) from the apical turn of harbor porpoises (A-C-E) and a common dolphin (B), where it is possible to observe three rows of stereocilia. C: Detail of the lateral links (arrows) attaching the outer hair cell stereocilia. D: The rectangle highlights the presence of one scar (asterisk) formed with two expansions of outer pillar cells after the disappearance of one OHC in the first row. E: All OHCs have disappeared and a large scar is formed by many supporting cells including outer pillar cells and Deiters cells. F: Schematic diagram showing the process of a scar formation (source: Leonova and Raphael, 1997, with permission of the editor). Sample B was fixed by SOCPVS before 18 hours postmortem. A, D, and E were fixed by the University of Utrecht, at 16 hours (A and E), at less than 7 hours (C), and for at least 3 hours postmortem (D). IHC, inner hair cell; OHC, outer hair cell; IPC, inner pillar cell; OPC, outer pillar cell; DC, Deiters cell. Scale bar = 2 μm in A; 3 μm in B; 500 nm in C: 50 μm in D; 20 μm in E.
tuning mechanism (Pujol et al., 1991), the small size of odontocete OHCs is consistent with a morphological adaptation to high-frequency hearing. These results confirm that odontocetes, together with bicolored roundleaf bats, present the shortest OHC ever described in mammals and reflect the higher frequency hearing capabilities of these animals. Harbor porpoises, for example, can hear up to 180 kHz (Kastelein et al., 2002) and roundleaf bats up to 200 kHz (Neuweiler et al., 1984), whereas most echolocating bats hear up to 150 kHz (Grinnell, 1995). Interestingly, a comparison of OHC length in the odontocetes we studied and the codified frequency of these particular cells in other species shows that these odontocetes do not follow the relationship described by Pujol et al. (1991) and Dannhof et al. (1991); in fact, we found that OHCs were much shorter than expected all along the cochlear spiral. The reason for this discrepancy is not yet clear.

In addition to their short length, the OHCs also had relatively small diameters. Unlike terrestrial mammals, in which OHC diameter is approximately constant at 7 μm, in the species in this study OHC diameter ranged from 3.9 to 5.4 μm. The smaller OHC dimensions in these odontocetes might coincide with a decrease in the OHC lateral membrane surface and a decrease in the membrane cell capacitance, thus increasing conductance and sensitivity to high frequency (Housley and Ashmore, 1992). The total numbers of IHCs (2,842) and OHCs (10,367), estimated here for harbor porpoise, and OHCs (13,405) for striped dolphin were consistent with previous results for other odontocete species. For example, 3,272 IHCs and 12,899 OHCs were calculated for the Pacific white-sided dolphin (Wever et al., 1972), and 3,451 IHCs and 13,933 OHCs for the bottlenose dolphin (Wever et al., 1971a). The magnitude of these hair cell estimations is similar to what was reported by Retzius (1884) for the human ear (3,475 IHCs and 11,500 OHCs), which has about the same cochlear length.

**Stereocilia**

The number of stereocilia on the outermost row per OHC was high and seemed to be constant along the cochlear length. If the cells have the same number of stereocilia in all rows, the number on each cell would range from 130 in harbor porpoise to 168 in striped dolphin (Table 5), but further research is needed to confirm these results. In some terrestrial mammals the number of stereocilia changes along the basoapical axis and between the OHC rows. For example, in humans the number of OHC stereocilia varies from 50 to 120 between the apical and basal turns (Kimura et al., 1964); in chinchillas the number of OHC1 and OHC2 stereocilia remains quite constant (100–110 stereocilia in the basal turn and 90–100 in the apex), whereas in the OHC3 the number of stereocilia varies from 80 in the basal turn to 18–40 in the apical turn (Lim, 1986). The large number of stereocilia in cetaceans may have at least two consequences: 1) to improve the functional relationship between the tectorial membrane and the organ of Corti; 2) to increase the number of ion channels that would facilitate depolarization of the cell and enhance its sensitivity (Assad et al., 1991).

**Scarring process**

In the case of a harbor porpoise in which we observed extensive scar formation in the OHC region of a large portion of the upper apical turn (Fig. 5E), scarring damage would affect the hearing in the lower frequency range. Once the cochlear frequency map for a species...
is resolved, it will be possible to estimate the acoustic characteristics of a source that may have caused damage by acoustic overstimulation. This is possible because “the greatest hearing-loss occurs at a frequency about half an octave above the exposure tone” (Davis et al., 1950). The frequency range of the suspect source could correspond to several anthropogenic marine activities, for example, shipping, pile driving, or

**Figure 7.** Scanning electron microscopy images of the outer hair cell stereocilia imprints on the undersurface of the tectorial membrane of a striped dolphin (A, arrows, and B), northern bottlenose whale (C), harbor porpoise (D), and common dolphin (E). F: Enlarged imprints in the tectorial membrane of a guinea pig cochlea after a loud sound exposure (reproduced from Morisaki et al., 1991, with permission of the editor). G,H: Inner hair cell stereocilia imprints (arrows) on the Hensen stripe of the most basal portion of the tectorial membrane of a common dolphin. The samples were provided by the Generalitat de Catalunya-Fundació CRAM (A,B), the University of La Rochelle (C), the University of Liège (D), and SOCPVS (E,G,H). Hs, Hensen stripe. Scale bar = 10 μm in A; 3 μm in B,C; 5 μm in D,G; 2 μm in H; 1.5 μm in E.

**TABLE 5.** Analysis of the Outer Hair Cell (OHC) Stereocilia Imprints on the Undersurface of the Tectorial Membrane

<table>
<thead>
<tr>
<th>Tectorial membrane</th>
<th>No. of imprints</th>
<th>No. of stereocilia/OHC</th>
<th>No. of animals</th>
<th>No. of cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>Striped dolphin</td>
<td>56</td>
<td>168</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>Harbor porpoise</td>
<td>43.4</td>
<td>130.2</td>
<td>3</td>
<td>22</td>
</tr>
<tr>
<td>Northern bottlenose whale</td>
<td>49.6</td>
<td>148.8</td>
<td>1</td>
<td>15</td>
</tr>
<tr>
<td>Common dolphin</td>
<td>52.75</td>
<td>158.25</td>
<td>2</td>
<td>8</td>
</tr>
</tbody>
</table>
seismic operations. However, other causes cannot be ruled out. For example, this particular animal was relatively old, and studies of age-related hearing loss in other mammals have shown that the base of the cochlea is consistently affected, as well as, in some cases, a small region of the apex (Johnsson and Hawkins, 1972; Spongr et al., 1997). Nevertheless, although the basal turn of the cochlea was not well preserved, thus preventing determination of the possible influence of aging in the analysis, we could clearly distinguish between disappearance of hair cells prior to death and damage to hair cells due to autolysis. In the first case, the stereociliary bundle was missing and a scar was made by neighboring epithelial cells. In the second case, the stereocilia were generally missing, but the cuticular plate of the hair cells remained and the epithelial mosaic was retained.

Subsurface cisternae

In the horseshoe bat (Vater et al., 1992), Mexican free-tailed bat (Vater and Siefer, 1995), rats, mice, and human (Arnold and Anniko, 1990) there is only one layer of subsurface cisternae. In contrast, guinea pig OHCs display multiple layers of subsurface cisternae, especially in the apical region (Saito, 1983). Subsurface cisternae are believed to play an important role in OHC motility by releasing Ca$^{2+}$ (Schulte, 1993; Tolomeo et al., 1996; Oghalai et al., 1998; Koyama et al., 1999). The present study showed that the harbor porpoise and the striped dolphin present the same pattern as the horseshoe bat (Vater et al., 1992).

Deiters and Pillar cells

In the basal turn of the cochlea in odontocetes as well as echolocating bats, the exaggerated thickness of the reticular lamina, made of both Deiters cell phalanges and heads of outer pillar cells, may provide an advantage for motion at high frequencies (Vater et al., 1992). The Deiters cells, Deiters cups, and pillar cells in the basal cochlear turn of the species evaluated in our study showed robust cytoskeletal structures (Fig. 3G,H), suggesting that the mechanical anchorage of the OHCs is reinforced and extremely stiff compared with the arrangement in the apical turn or throughout most of the cochlea of other mammals. Significantly, the disposition of Deiters cups found in odontocetes is also found in the organ of Corti of the basal turn in echolocating bats (Henson and Henson, 1979; Dannhof and Bruns, 1991; Vater et al., 1992; Vater and Siefer, 1995). This feature may therefore be relevant for high-frequency processing (Reysenbach de Haan, 1957; Wever et al., 1971c, 1972; Vater et al., 1992).

Innervation

Medial efferent innervations

Nerve endings were found connecting with the base of the OHCs (Fig. 4A). We interpret these nerve terminals as afferents (type II), because the OHCs were lacking postsynaptic cisternae and we did not observe thick upper crossing fibers in the tunnel of Corti. Studies in humans have shown that the efferent nerve endings and postsynaptic cistern are very resistant to anoxia, and usually remain present, even when the OHCs are in a very advanced decomposition stage (Bruns and Schmieszek, 1980; Lavigne-Rebillard and Pujol, 1988, 1990). Thus, we could not find evidence that these species receive medial efferent innervation onto the OHCs.

A lack of medial efferent innervation was also shown in echolocating horseshoe bats, in bicolor and Indian roundleaf bats (Bruns and Schmieszek, 1980; Aschoff and Ostwald, 1988; Bishop and Henson, 1988; Dannhof and Bruns, 1991; Vater et al., 1992), and in the mole rat Spalax ehrenbergi (Raphael et al., 1991), a species specialized for low-frequency hearing (Bruns et al., 1988). This evidence supports the notion that the medial efferent innervation of the OHCs may be a regulating system for the mid-frequencies, and may not be needed at extremely high and low frequencies (Vater et al., 1992). However, other studies on the echolocating species Mexican free-tailed bats (Tadaria brasiliensis) and Parnell’s mustached bats (Pteronotus parnellii) showed the presence of medial efferent innervation onto the OHCs (Bishop and Henson, 1988; Vater and Siefer, 1995). Thus, there is no straightforward functional explanation for the absence of medial efferent innervation in some echolocating species (Vater et al., 1992).

Spiral ganglion cells

The density of SGCs, as measured by the number of cells per cross-sectional area, in Rosenthal’s canal (4.4–8 SGCs/10,000 μm$^2$) in the odontocetes studied here is similar to that in guinea pigs (8 SGC/10,000 μm$^2$; Jin et al., 2006), but much lower than in Mongolian gerbils (28.1 SGC/10,000 μm$^2$; Fujita et al., 2007) or mice (between 36.3 in the apex and 39.6 SGC/10,000 μm$^2$ in the base; Sato et al., 2006). Considering that the bottlenose dolphin has three times, and the Pacific white-sided dolphin almost twice as many SGCs as the human (Wever et al., 1971a, 1972), we expected a higher density of SGCs in the odontocetes in our study. However, the SGCs of the harbor porpoise and striped dolphin were much larger in size (maximum 45 × 25 μm and 46 × 30 μm, respectively, Table 3) than those found in rodents (for example, 13–18 μm in rats;
Bichler, 1984), and even larger than those in bottlenose dolphins, in which basal turn SGCs were $38.4 \times 25.2$ $\mu$m and middle turn SGCs were $34.4 \times 21.1$ $\mu$m (Wever et al., 1971a). In fact, odontocetes increase the number of SGCs by increasing the space occupied in Rosenthal’s canal. There are some differences between species. For example, the number of SGCs in a transverse section of Rosenthal’s canal ranged from 128 to 152 in the striped dolphin, and from 122 to 182 in the harbor porpoise (not illustrated here), but was only 81–88 and 57–67 in gerbils (Ruttiger et al., 2007). Our results are consistent with the large range of SGCs to hair cells found by Wever and colleagues (1971a), suggesting “unusual capabilities in the utilization of auditory information.” Considering that the peripheral process of each neuron contacts one IHC, it could be inferred that, in the cochlea of the odontocetes analyzed in this study, IHCs have many more afferent synapses than IHCs in cochleas of terrestrial mammals. In turn, this rich afferent innervation should provide functional advantages to these odontocetes, possibly increasing sensitivity and/or dynamic range.

**Basilar membrane**

The width and thickness of the basilar membrane have been measured in several mammalian species (Nadol, 1988; Roth and Bruns, 1992; Sato et al., 1999; Keller and Richter, 2001). A clear gradient in width and thickness has been reported in most species as well as in the odontocetes analyzed in this study: narrower, thicker, and relatively stiff in the base, and broader and thinner in the apex.

The basilar membrane was wider and thinner in approximately the first 2 mm of the cochlea. A decrease in basilar membrane thickness in the basal hook was previously described in Parnell’s mustached bat and the greater horseshoe bat (Bruns, 1976; Kössl and Vater, 1985), and an increase in width in bicolored roundleaf bat (Dannhof and Bruns, 1991). The functional role of the basilar membrane is not very well understood. In the case of the bottlenose dolphin, the width of the basilar membrane increased by about 14 times from the base (25 $\mu$m) to the apex (350 $\mu$m; Wever et al., 1971b), whereas the thickness diminished by five times (from 25 to 5 $\mu$m). However, the basilar membrane in humans showed an increase in width of only about four times (from 125 to 500 $\mu$m) and a threefold decrease in thickness from the base (7 $\mu$m) to the apex (2 $\mu$m) (Schuknecht, 1993; Ketten, 1998).

The values obtained for the harbor porpoise and the striped dolphin (Table 2) did not show an increasing trend, as in the bottlenose dolphin, even though the striped dolphin and the bottlenose dolphin have similar hearing sensitivities. It is possible that this difference in dimensions could result from a lack of information on the extreme locations of the cochlea (first and last millimeters; Fig. 2), or because the criterion to establish the limits of the basilar membrane was different. Further comparative research should be conducted to investigate differences between these species. In odontocetes, the basilar membrane is very stiff at the base of the cochlea, fixed to both sides by a very well-developed inner and outer osseous spiral laminae that confers an adaptation for very high-frequency hearing (Wever et al., 1971b; Ketten and Wartzok, 1990; Ketten, 1992, 1994).

**Tectorial membrane**

Despite the very fast autolysis processes after death, the tectorial membrane was less affected than other structures, perhaps because it is composed of an extracellular matrix (Slepecky et al., 1992). The high resistance of this structure to anoxia would allow the diagnosis of possible alterations in the OHC stereocilia due to acoustic trauma (Morisaki et al., 1991), even if the cochlea was fixed after more than 20 hours post-mortem. In all our samples, the imprints of stereocilia on the tectorial membrane were regular. Because the shape of these imprints mirrors the initial organization of the stereocilia in the OHCs, there was no evidence of acoustic trauma. That is not to say that acoustic trauma did not occur in some other specimens, as suggested by the presence of epithelial scars seen in SEM (Fig. 5E). We detected the presence of IHC stereocilia impressions in the very basal region of a common dolphin tectorial membrane (Fig. 7G,H) for the first time in odontocetes; this feature has been previously described in rats (Lenoir et al., 1987) and bats (Vater and Lenoir, 1992). However, Matsumura (2001) was unable to demonstrate a direct physical attachment between IHCs and the tectorial membrane in most mammalian species. We believe that this characteristic was difficult to observe in all our samples because of the degraded state of the organ of Corti.

**GENERAL CONCLUSIONS**

This study has, for the first time, described the ultrastructure of the organ of Corti of odontocetes. The two species of odontocetes analyzed by TEM had morphological characteristics that support high-frequency hearing, including: 1) a short length of OHCs; 2) a thick cuticular plate in OHCs, Deiters cells, and outer pillar cells; 3) robust cup formation of the Deiters cell body; 4) a highly developed cytoskeleton in Deiters and pillar cells; and 5) a high thickness of the basilar membrane.
All these features, in addition to a common molecular design of prestin (Li et al., 2010; Liu et al., 2010), an OHC motor protein (Zheng et al., 2000), and a parallel evolution of other auditory genes related to hair cell bundle motility, hair cell activity, and nerve transmission (cadherin 23, protocadherin 15, otoferlin, transmembrane cochlear-expressed gen1, and pejvakin; Davies et al., 2012; Shen et al., 2012), are also shared by echolocating bats (for review, see Vater and Kössl, 2011), suggesting a convergent evolution among echolocating species.

An additional goal of this study was to use morphological features of the inner ear to detect structural alterations that might result from sound overexposure. This proved to be difficult, because delays in tissue fixation from stranded animals resulted in cochlear decomposition that prevented us from diagnosing acoustic trauma from changes in hair cell stereocilia (Bredberg et al., 1972; Engstrom et al., 1984), or other alterations at the sensory cell (Hu et al., 2000) or innervation levels (Spoendlin, 1971). However, the presence of scars (Fig. 5D,E), which indicates premortem hair cell disappearance, could be clearly distinguished from autolysis artifacts. In fact, in one case (harbor porpoise; Fig. 5E), the lesions would be consistent with an acoustic trauma.

The tectorial membrane appeared to be more resistant to postmortem autolysis and remained in acceptable condition for analysis when the cochlea was fixed more than 20 hours postmortem. The analysis of the stereocilia imprints on the tectorial membrane has provided insights into odontocete hair cell stereocilia organization. In addition, because of the difficulties of obtaining and analyzing fresh samples of cetacean ears immediately after stranding, our results suggest an alternative approach using image analysis of the tectorial membrane, a procedure not yet common, as a diagnosis to detect possible lesions due to noise overexposure.

The results presented in this study show that the combination of the two complementary imaging techniques is necessary to document whether an odontocete has suffered from an acoustic trauma: first an overview by SEM to localize possible lesions along the cochlear axis, focusing on the presence of scars and irregularities of the imprints of the tectorial membrane; and second, a more detailed analysis by TEM to characterize the lesion on a transverse view that contains information on all the cochlear cells involved.

Acknowledgments

We thank the auditory team researchers at the Institut des Neurosciences de Montpellier (INM), Jérôme Bourien, Jing Wang, Remy Pujol, Benjamin Delprat, and Michel Eybalin, for their help and advice in interpreting the results, and Jean-Louis Pasquier for help in iconography. We are grateful for the technical assistance provided by Sabine Ladrech and Florence François (INM), Cécile Sanchez (CRIC), Lluisa Matas and Imma Arrom (Research Technical Service of Girona University), and Derrick Horne (Bioimaging Facility, University of British Columbia). We also thank our colleagues and the stranding organizations who helped us in collecting the ears, especially Lidewij Wiersma, Lonneke IJsseldijk, and Sjoukje Hiemstra (University of Utrecht), Pablo Cermeno (AMBAR), Paco Toldano (PROMAR), Ángela Llavona, Josep M. Alonso, Alfredo López, and María Lliarena (CEMMA), Beatriz González (Fundación CRAM), Encarna Gómez (University of Barcelona), Denik Uqini (Universiteit i Skodres), Mardik Leopold (IMARES), and Kees C.J. Camphuysen (Royal NIOZ).

Role of Authors

All authors had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Study concept and design: Morell, André, and Lenoir. Acquisition of data: Morell, Jauniaux, Dabin, Begeman, Ferreira, Maestre, Degollada, and Hernandez-Milian. Analysis and interpretation of data: Morell, Lenoir, Puel, and André. Drafting of the manuscript: Morell, André, and Lenoir. Critical revision of the manuscript for important intellectual content: Jauniaux, Dabin, Begeman, Ferreira, Maestre, Degollada, Hernandez-Milian, Cazevieille, Fortune, Shadwick, Vogl, and Puel. Administrative, technical, and material support: Cazevieille, Fortune, Shadwick, Vogl, and Puel. Study supervision: André, Lenoir, and Shadwick.

Conflict of Interest Statement

The authors have no conflicts of interest of any nature.

Literature Cited


We present scanning (SEM) and transmission (TEM) electron microscopy micrographs of the ultrastructure of the organ of Corti of odontocete species. The species of odontocetes analyzed by TEM presented morphological characteristics, shared with horseshoe bats, suggesting that there has been convergent evolution of sound reception mechanisms among echolocating species.