### Nanostructure of mouse otoconia

2	Dimitra Athanasiadou <sup>a</sup> , Wenge Jiang <sup>a,1</sup> , Natalie Reznikov <sup>b</sup> , Alejandro B. Rodríguez-Navarro <sup>c</sup> ,
3	Roland Kröger <sup>d</sup> , Matthew Bilton <sup>e</sup> , Alicia González-Segura <sup>f</sup> , Yongfeng Hu <sup>g</sup> , Valentin Nelea <sup>a</sup> , Marc
4	D. McKee <sup>a,h*</sup>
5	<sup>a</sup> Faculty of Dentistry, McGill University, Montreal, QC, Canada H3A 0C7
6	<sup>b</sup> Object Research Systems Inc., Montreal, QC, Canada H3C 1M4
7	<sup>c</sup> Departamento de Mineralogía y Petrología, Universidad de Granada, Granada, Spain 18002
8	<sup>d</sup> Department of Physics, University of York, York, UK YO10 5DD
9	<sup>e</sup> Imaging Centre at Liverpool, University of Liverpool, Liverpool, UK L69 3GL
10	<sup>f</sup> Centro de Instrumentación Científica, Universidad de Granada, Granada, Spain 18002
11	<sup>g</sup> Canadian Light Source, University of Saskatchewan, Saskatoon, SK, Canada S7N 2V3
12	<sup>h</sup> Department of Anatomy and Cell Biology, McGill University, Montreal, QC, Canada H3A 0C7
13	
14	*Corresponding author
15	Email address: Prof. Marc D. McKee marc.mckee@mcgill.ca
16	
17	<sup>1</sup> Present address: Department of Chemistry, Tianjin Key Laboratory of Molecular Optoelectronic
18	Sciences, and Tianjin Collaborative Innovation Center of Chemical Science & Engineering,
19	Tianjin University, Tianjin, P. R. China, 300072.
20	
24	
21	

#### 23 Abstract

Mammalian otoconia of the inner-ear vestibular apparatus are calcium carbonate (calcite)-24 containing mineralized structures critical in maintaining balance and in detecting linear 25 acceleration. The mineral phase of otoconia coherently diffracts X-rays much like single-crystal 26 calcite. Among mineral-associated proteins, otoconia contain osteopontin (OPN) - a highly 27 phosphorylated mineral-binding protein influencing mineralization processes in bones, teeth and 28 29 avian eggshells. Here, we investigate mineral structure and OPN distribution in mouse otoconia at the ultrastructural level. Scanning electron and atomic force microscopy of intact mouse otoconia 30 and otoconia cleaved open using a microtome revealed an internal nanostructure (averaging 50-31 32 nm Feret diameters). Further investigation by transmission electron microscopy and electron tomography of focused ion beam-prepared sections of otoconia confirmed this mineral 33 nanostructure, and identified even smaller (approximately 10 nm) nanograin dimensions. X-ray 34 35 diffraction identified only calcite mineral in the otoconia, and Raman and X-ray absorption spectroscopy – both methods being sensitive to the detection of crystalline and amorphous forms 36 in the sample – showed no evidence of amorphous calcium carbonate in the otoconia. Scanning 37 and transmission electron microscopy combined with colloidal-gold immunolabeling for OPN 38 39 revealed that this protein was located at the surface of the otoconia, correlating with a site where 40 surface nanostructure was observed. OPN addition to calcite growing in vitro produced similar surface nanostructure. Finally, these findings provide details on the composition and nanostructure 41 of mammalian otoconia, and suggest that while OPN may influence surface rounding and surface 42 43 nanostructure in otoconia, other incorporated proteins (also possibly including OPN) likely participate in creating internal nanostructure. 44

45 Keywords: Biomineralization; otoconia; osteopontin; calcite; nanostructure; tomography

#### 46 **1. Introduction**

Biomineralization processes have been broadly studied for many mineralized tissues and 47 structures (Lowenstam and Weiner, 1989). Otoconia (and otoliths in fish) are composites of 48 calcium carbonate ( $CaCO_3$ ) mineral and proteins found in the utricle and the saccule of the 49 vestibular apparatus of vertebrate mammals, and in the lagena of nonmammalian vertebrates 50 (Hiatt, 2014; Lundberg et al., 2015). The vestibular system of the inner ear is responsible for head 51 52 movement detection, for spatial orientation, and for body balance (Hiatt, 2014; Marieb, 1991). In mammals, the vestibular system has three fluid-filled semicircular canals which respond to 53 rotational acceleration, and two receptor organs – the utricle and the saccule – which sense linear 54 55 acceleration and gravity (Hughes et al., 2006; Lundberg, 2012; Lundberg et al., 2015).

Otoconia are formed in mammals during the late embryonic stages, and they are considered 56 stable until mid-life where signs of their degeneration can be detected in humans (Anniko et al., 57 58 1984). In mice – which share many similar otoconial features to humans – the initial seeding of otoconia is detected at embryonic day 14, with the highest rate of mineralization occurring at 59 embryonic days 15 and 16 (Ornitz et al., 1998). Murine otoconia acquire their final size (a few 60 micrometers) by postnatal day 7, and from there, in health, they generally are preserved throughout 61 62 life with minimal changes/transitions (House and Honrubia, 2003; Ornitz et al., 1998). Otoconia, 63 surrounded by a low-calcium solution termed the endolymph, are embedded in a membranous structure called the otoconial membrane, which rests on the extremities of hair cell kinocilia and 64 stereocilia in the utricular and saccular sensory epithelium, regions known as the macula 65 66 (Lundberg, 2012; Lundberg et al., 2015). With each head movement, the otoconia are displaced 67 by their moments of inertia relative to the underlying cells, leading to bending of the sensory hair bundles (Lundberg, 2012). This mechanical impetus is transformed to electrical signals that are 68

transmitted by the hair cells to the central nervous system of the brain, providing information onthe spatial position of the head (Lundberg, 2012; Marieb, 1991).

Morphologically, mammalian otoconia are barrel-shaped calcitic structures bounded by 71 well-defined rhombohedral facets at both ends and which diffract as single crystals (Lundberg, 72 2012). They are unique in being the only calcium carbonate-containing biomineral found in normal 73 74 healthy mammals; the main mineral phase of bones and teeth is a calcium phosphate (apatite). However, in some species, calcium phosphate otoconia can be found such as in the Agnatha 75 species (jawless fish) where there is negligible or absent crystalline structure (Carlstrom, 1963), 76 77 or in malformed human inner ears where a mixture of apatite and calcite has been reported (Johnsson et al., 1982). Otoconia can occur as any of the  $CaCO_3$  polymorphs depending upon the 78 organism within which they reside (Ross and Pote, 1984). In mammals and birds, the polymorph 79 for otoconia is typically calcite, the most thermodynamically stable calcium carbonate polymorph 80 existing under ambient conditions (Lins et al., 2000). In amphibians and fish, the predominant 81 polymorph is aragonite (Carlstrom, 1963; Ross and Pote, 1984). Vaterite is found in primitive 82 jawfish or in pathologic conditions in the human inner ear (Addadi et al., 2003; Johnsson et al., 83 1982; Ross and Pote, 1984; Wright et al., 1982). Specific matrix proteins have been shown to 84 85 promote the calcium carbonate polymorph selection of otoconia (Pote and Ross, 1991).

As in other biominerals, and important for the CaCO<sub>3</sub> formation in the inner ear, is the presence of an organic matrix consisting mainly of glycoproteins and proteoglycans (Fermin et al., 1995; Lundberg, 2012; Lundberg et al., 2015; Pote and Ross, 1993; Tachibana and Morioka, 1992). The presence of otoconial proteins is necessary for the formation of otoconia where they sequester and concentrate calcium ions from the endolymph fluid (Lundberg, 2012), likely amongst other important functions in guiding otoconial growth. Prominent proteins identified in

mammalian otoconia are otoconin 90 (OC90), otolin-1 (or otolin), osteopontin (OPN), fetuin-A, 92 SPARC-like protein 1 (SC1), secreted protein acidic and rich in cysteine (SPARC), dentin matrix 93 protein 1 (DMP1) (Lundberg, 2012; Lundberg et al., 2015; Thalmann et al., 2006) and  $\alpha$ -tectorin 94 (Xu et al., 2010). OC90 – the main soluble matrix protein of otoconia – modulates the form of 95 calcite crystals in vitro (Lu et al., 2010), whereas in vivo in mice, in the absence of this protein, 96 97 there is either an absence of otoconia, or only a few massive abnormal otoconia are formed, with overall calcite amounts being decreased roughly by half (Andrade et al., 2012; Xu et al., 2010). 98 Otolin – identified in bony fish (Murayama et al., 2005) – is a member of the collagen X family, 99 100 and is found in both otoconia and the surrounding otoconial matrix where it might serve as a scaffold protein for biomineralization (Moreland et al., 2014; Yang et al., 2011). Other proteins 101 such as fetuin-A, SPARC, OPN and DMP1, are considered as minor otoconins because of there 102 being a negligible phenotype effect on otoconial formation and vestibular function in transgenic 103 mice (Xu et al., 2010; Zhao et al., 2007; Zhao et al., 2008). Other candidates for mediating 104 otoconial mineralization are the keratan sulfate proteoglycans (KSPGs) (Xu et al., 2010), these 105 being extended biomolecules having strong negative charges for attracting calcium ions and 106 appearing to interact with OC90 and otolin proteins (Lundberg, 2012). 107

OPN is a highly phosphorylated mineral-binding protein, having multiple roles in cell adhesion and protein binding in the extracellular matrix of bone and teeth, where the phosphoserine residues and the overall negative charge of this molecule elicited from Asp and Glu residues appear to be important for regulating mineralization processes (Fisher et al., 2001; Sodek et al., 2000; Sorensen et al., 1995). OPN is expressed by the sensory hair cells, the nonsensory dark cells, and the cells of the endolymph sac (Zhao et al., 2008). In recent work pertaining to OPN and biomineralization in another calcium carbonate structure – the avian eggshell – we have shown an association of OPN with the nanostructured texture of the calcitic shell, and we have reproduced
similar internal nanostructure in calcite crystals grown *in vitro* in the presence of OPN
(Athanasiadou et al. 2018).

Structural studies of mammalian otoconia are sparse, and those that exist mainly describe 118 pathologic circumstances and developmental abnormalities (Johnsson et al., 1982; Wright et al., 119 120 1982). In terms of their ultrastructure, human otoconia are known to be composed of three sectors/branches at each end that extend outwards from the central so-called belly region, a site 121 seemingly more susceptible to otoconial degradation (Walther et al., 2014). Despite having both 122 123 rounded and faceted external morphology at the microscale suggesting perhaps nonclassical crystallization pathways (De Yoreo et al., 2015; Rodriguez-Navarro et al., 2016; Wolf et al., 2016), 124 it remains possible that an internal polycrystalline, coherent nanostructure forms during their 125 126 formation in the presence of mineral and calcium-binding proteins, as has been shown in numerous calcium-carbonate mineralizing systems (Athanasiadou et al., 2018; Jean-Pierre Cuif, 2010; Tseng 127 et al., 2014). 128

Morphological and compositional alterations of otoconia are frequently produced by head 129 trauma, ototoxic drugs, aging, and environmental and genetic factors, and these can lead to 130 131 balance-related disorders (Lundberg et al., 2015). Benign paroxysmal positional vertigo (BPPV) is a serious disease in which patients suffer from intense nausea and loss of balance (Salvinelli et 132 al., 2004). BPPV occurs when otoconia are dislodged from their initial position and migrate into 133 134 the semicircular canals (canalithiasis), or when otoconia exist in larger numbers than the active surface area of the utricular cells (utriculithiasis) (Oas, 2001). Aging-related otoconial 135 136 degeneration is a high-risk factor for creating free otoconial debris that results in loss of balance

in elderly people, consequently leading to falls which frequently cause bone fractures and evendeath (Agrawal, 2009).

Because of the important physiologic functions effected by otoconia, and because otoconial 139 pathologies lead to BPPV, accurate and detailed knowledge of the formation of otoconia and their 140 fine structure is fundamental to understanding the function of the vestibular system. Given the 141 142 limited understanding of otoconial formation and structure, and to better consider ways to influence their regrowth/regeneration to potentially treat their abnormalities, we describe here the 143 internal structure of mouse otoconia at the nanoscale level. We also describe the localization of 144 145 OPN at the ultrastructural level in these otoconia, and combine this with an examination of the *in* vitro effects of OPN on growing calcite morphology. 146

147

#### 148 2. Materials and Methods

#### 149 2.1 Harvesting and embedding of otoconia

150 Otoconia were processed by manual dissection under a stereomicroscope from C57BL/6 normal (wild-type) and *Opn<sup>-/-</sup>* (knockout) mice obtained from the Jackson Laboratory (Bar Harbor, 151 152 ME, USA). All mice used were 8 days old (except for those used for the 2D XRD and EBSD 153 analyses which were 3 months old, Sections 2.3 and 2.9 below). Animals were kept at 24°C in pathogen-free conditions using alternating 12-hour light and 12-hour dark cycles. Mice were fed 154 normal mouse chow (2920X, Teklad global soy protein-free extruded rodent diet, Envigo, 155 156 Huntingdon, UK), and had access to water ad libitum. Mice were sacrificed under isoflurane anesthesia by decapitation, and mouse heads were either fixed with aldehyde and then dissected to 157 isolate the inner ear and dehydrated and embedded in Epon or LR White resin, or the heads were 158 159 transferred unfixed immediately into 100% ethanol to prevent any potential dissolution of otoconia 160 from exposure to aqueous solutions. From the latter, otoconia were carefully retrieved through 161 dissection under a stereoscope, and stored in 100% ethanol. Embedded samples were sectioned for 162 histology and mounted on either glass slides for atomic force microscopy or onto grids for scanning 163 electron microscopy (that were also used additionally for transmission electron microscopy). 164 Animal procedures were evaluated and approved by the McGill University Institutional Animal 165 Care and Use Committee following the guidelines of the Canadian Council on Animal Care.

#### 166 **2.2 Light microscopy of the vestibular inner ear system**

For the histology of the inner ear vestibular system and observation of otoconia *in situ*, 1µm-thick sections from wild-type mouse otoconia were cut from polymerized blocks using a Leica
Ultracut E Ultramicrotome (Leica, Wetzlar, Germany). These sections were von Kossa-stained for
mineral (black reaction) followed by counter-staining with toluidine blue, with visualization and
image recording using an optical microscope (model DMRBE, Leica) equipped with a 3-CCD
Sony DXC-950 camera (Sony, Tokyo, Japan).

### 173 2.3 X-ray diffraction (XRD), Raman spectroscopy and X-ray absorption (XAS) of the 174 mineral phase of mouse otoconia

The mineral phase of wild-type mouse otoconia was analyzed by powder and single-crystal
X-ray diffraction (XRD), Raman spectroscopy and X-ray absorption spectroscopy (XAS). Powder
XRD was run on a Bruker D8 Discover diffractometer equipped with a Cu X-ray tube (wavelength
0.154056 nm) and an area detector (GADDS 2D XRD).

For XRD, measurements were performed in coupled θ-θ scan mode (500 µm beam spot
 size) with samples of otoconia spotted as aggregates onto a quartz sample substrate. Single-crystal
 XRD analyses were performed using a Bruker D8 Venture diffractometer equipped with a photon

area detector and a Cu X-ray microsource (200  $\mu$ m beam size). Measurements were taken in the transmission mode on aggregates of otoconia (dissected from 3-month-old wild-type mice). Frames were recorded while the sample was rotating in  $\varphi$  angle within a 10° angular range using 0.3° steps and an integration time of 60 seconds.

Raman spectroscopy was performed using a Renishaw inVia Raman microscope 186 187 (Renishaw, Gloucestershire, UK) equipped with a holographic spectrometer and a Leica DM2500 M optical microscope (Leica Microsystems GmbH, Wetzlar, Germany). The excitation source was 188 a 514.5 nm Ar laser with a ca. 2 µm laser spot size and a 25 mW excitation power. The laser was 189 190 focused through a  $50 \times$  objective (numerical aperture 0.75) on a dense array of otoconia. For each measurement, Raman spectra were acquired for 10 seconds where 3 scans were accumulated for 191 minimizing any noise effects. Spectral reproducibility was confirmed by taking several spot 192 analyses. Measurements were done on 1-µm-thick sections of utricle cut with an ultramicrotome 193 from Epon blocks containing sectioned otoconia and placed on glass coverslips. 194

195 XAS synchrotron data collection was performed at the Canadian Light Source (CLS) using 196 the SXRMB beamline. Sections of resin-embedded otoconia cut at 1- $\mu$ m thickness were placed 197 onto Si wafers. The electron beam was run at 2.9 GeV with a beam current of approximately 200 198 mA. A set of 5 scans was averaged from three samples to achieve the best signal-to noise ratio. 199 All spectra were obtained at the calcium K-edge and were normalized in reference to the positions 200 of the intrinsic monochromator glitches and of the calcite edge spectrum.

201

#### 2.4 Scanning electron microscopy (SEM)

To analyze by SEM both the outer surface of otoconia and the internal structure of microtome (diamond knife)-cleaved otoconia, isolated intact otoconia and resin-embedded otoconia were prepared. For external morphology examination, intact isolated whole otoconia were placed on aluminum SEM stubs, coated with a 2-nm thick Pt layer, and examined by SEM at an accelerating voltage of either 2 or 5 kV using an Inspect F-50 FE-SEM (FEI Company). Without sputter coating, ultramicrotome-cut 1- $\mu$ m-thick microtome sections of otoconia were placed on conductive grids (also for future TEM use) where likewise imaged by SEM under the same conditions.

#### 210 **2.5 Atomic force microscopy (AFM)**

211 To further investigate possible internal nanogranular structure of otoconia, and to correlate 212 this with the nanoscale distribution of organic matrix in the same sample, atomic force microscopy (AFM) was conducted on 1 µm-thick ultramicrotome cut sections from wild-type mice. AFM 213 214 height and amplitude images were taken using a Multimode Nanoscope IIIa atomic force 215 microscope (Veeco, Santa Barbara, CA, USA) operating in the tapping mode in air at room 216 temperature using a nonvertical engage E-scanner and NanoScope version 5.30 software (Veeco/Bruker-AXS Inc., Madison, WI, USA). In the AFM experiments, V-shaped tapping mode 217 probes (typical tip apex radius of approximately 7 nm) with Si cantilevers having a spring constant 218 219 k = 42 N/m (Bruker-AXS Inc.) were used. The tip force exerted on the surface was optimized by 220 the amplitude set-point being as high as possible to reduce imaging artefacts. The Feret diameters of the units comprising the observed nanostructure were calculated using ImageJ software (1.x 221 222 version) (Schneider et al., 2012). At least 200 Feret diameters of the otoconial nanostructure 223 domains were calculated from the AFM images (obtained in amplitude mode) after performing the software's high-pass processing to enhance domain boundaries. 224

# 225 2.6 Focused-ion beam (FIB) sample preparation and transmission electron microscopy 226 (TEM)

227 The TEM investigation was performed on a thin FIB section of an otoconium prepared by a dual-beam FIB system (FEI Helios 600 NanoLab, FEI, Hillsboro, OR, USA) equipped with a 228 gallium ion source. For this, a nickel TEM grid with otoconia was placed on a flat aluminium SEM 229 stub. A single otoconium was then identified and covered with a protective 2 µm-thick Pt layer 230 from which a 2 µm-thick slab cut from the otoconium was milled/thinned with the FIB. Afterward 231 this initial thinning, the section was transferred onto a Cu TEM half-grid using an EasyLift 232 nanomanipulator for the final thinning at 5 kV and 9.4 nA, from which an 80-nm-thick section was 233 obtained. Bright-field TEM images and selected-area electron diffraction (SAED) patterns were 234 235 obtained using a Tecnai TF-20 (FEI) microscope operating at 200 kV.

#### 236 **2.7 Electron tomography**

237 The 80-nm-thick FIB-cut section of an otoconium collected on a TEM Cu grid was used to 238 collect a series of single-axis tilt images at an accelerating voltage of 200 kV using a Tecnai G2 239 F20 cryo-S/TEM (FEI) equipped with a Gatan Ultrascan 4000  $4k \times 4k$  digital CCD camera system (model 895). Images were taken at a magnification of 62,000X over a tilt range from  $-40^{\circ}$  to  $+50^{\circ}$ 240 241 for the samples (2° increments in both low tilts and high tilts). The resulting images had pixel sizes of 0.19 nm. For electron tomography in a scanning transmission electron microscope (STEM) 242 (shown in Supplementary Material), images were recorded using a ThermoFisher 300 kV Titan<sup>3</sup> 243 244 Themis X-FEG S/TEM at a magnification of 62,000 over a tilt range from  $-50^{\circ}$  to  $+70^{\circ}$  (2° increments in low tilts and 1° high tilt step on the 80-nm-thick sections). The images from the tilt 245 series were aligned, filtered and reconstructed into a tomogram using the IMOD software package 246 (Kremer et al., 1996). The movies for the raw tilt series and reconstruction were done in IMOD, 247 whereas the movies with 3D volume with solid and surface rendering were generated using UCSF 248 Chimera (version 1.10.1) software. 249

#### 250 **2.8 3D imaging and reconstruction**

### Three-dimensional imaging of intact utricular otoconia embedded in an LR White resin block was performed using a Zeiss Xradia 520 Versa (Carl Zeiss Canada Ltd). A series of X-ray frames (totalling 1989 projections) was collected over 360-degrees of rotation at 60 V with a pixel resolution of 0.5 $\mu$ m. A reconstructed movie of the rotation series was generated using Dragonfly<sup>TM</sup> v3.6 software (Object Research Systems Inc., Montreal).

#### 256 **2.9 Electron backscatter diffraction (EBSD) and pole figures**

Electron backscattering diffraction (EBSD) maps were collected for 20 h from FIB-cut sections of otoconia in the transmission mode using an Auriga Zeiss SEM instrument and a 0.3 μm-step size resolution. All EBSD data were collected and analyzed with AZtec 2.1 software (Oxford Instruments). Data was visualized as crystallographic orientation maps using pseudocolors to represent crystal orientations or pole figures.

#### 262 2.10 Calcite crystal growth in the presence of OPN

Synthetic calcite crystals were grown *in vitro* in a 10 mM CaCl<sub>2</sub> solution with (or without) 263 added full-length phosphorylated bovine milk OPN (0.15, 0.3, 0.45 and 0.9 µM) as provided by 264 265 Arla Foods and prepared according to the procedure described by Sørensen and Petersen (Sorensen and Petersen, 1993). Calcite crystallization took place for 2 hours on round glass coverslips in 266 267 small wells placed within a sealed desiccator previously charged with 1 g of (NH<sub>4</sub>)<sub>2</sub>CO<sub>3</sub> powder. Glass coverslips with calcite crystals were gently removed from solution, and lightly rinsed with 268 distilled water and ethanol, and then, air-dried and placed in a desiccator for storage until further 269 270 characterization. Triplicates were used for all the crystallization experiments.

#### 271 2.11 Immunogold labeling for OPN

272 For immunogold OPN labeling of otoconia in situ in normal wild-type mice and OPNdeficient mice (negative control), TEM grid-mounted microtome-cut sections 80 nm in thickness, 273 as well as the respective polymerized block faces themselves from which the sections were cut, 274 were used to localize OPN at the EM level. Otoconia-containing thin sections and block faces were 275 incubated with anti-mouse OPN antibody (R&D Systems Oakville, Canada) for 1 hr followed by 276 277 washing and incubation with protein A-colloidal gold complex (14-nm gold particles, from G. Posthuma, University of Utrecht) for 1 hr to detect the immunolabeling reactions. For the sections, 278 conventional staining with uranyl acetate and lead citrate was performed after the immunolabeling 279 280 protocol. The microtome-cut sections were visualised by TEM as described above, and the block faces were visualized by SEM. 281

282 **3.1 Results** 

# 3.2 Optical microscopy and scanning electron microscopy (SEM) – external morphology of otoconia

Figure 1a shows by light microscopy an overview of the location of otoconia in the mouse 285 286 inner ear from a von Kossa and toluidine blue-stained section of the utricle from wild-type mice. Otoconia stained black for mineral (arrow) rest upon a gelatinous mass - the otoconial membrane 287 - overlying the hair cells of the utricular macula. Figures 1b and 1c show by SEM the external 288 morphology of wild-type mouse otoconia at low and high magnification, respectively, while 289 290 Movie S1 (Supp. Material) displays the 3D organization of otoconia in the utricle as determined from microcomputed X-ray tomography. The size of the majority of the otoconia generally ranged 291 between 5 to 8 µm, with some smaller otoconia of approximately 2 µm in length being present 292

(Fig. 1b). Higher magnification of the otoconial surface reveals an external nanogranular structure
on the barrel-shaped body (Fig. 1c inset), whereas the triplanar faceted surfaces appear smooth
(Fig. 1c).



296

Figure 1. External morphology of wild-type mouse otoconia. (a) Histological staining of a
 section of the mouse utricle observed by light microscopy after von Kossa staining for mineralized
 otoconia (black, arrows). (b) Low-magnification SEM image of otoconia. (c) High-magnification
 SEM image showing details of otoconia having surface nanostructure (inset).

## 302 3.3 XRD, Raman spectroscopy and XAS – identification of CaCO<sub>3</sub> polymorphs and 303 crystallinity of otoconia

304 As has been observed previously (Pote and Ross, 1986), mouse otoconia are crystallized 305 in the form of calcite. XRD patterns from a cluster of many agglomerated otoconia revealed

characteristic peaks corresponding only to calcite (Fig. 2a). Similar to the XRD results, Raman 306 spectra revealed characteristic peaks in the region between 100-1200 cm<sup>-1</sup> attributable to vibrations 307 arising from the calcite polymorph (Fig. 2b). Particularly demonstrating this, the bands 281 cm<sup>-1</sup> 308 and 155 cm<sup>-1</sup> are characteristically attributable to the vibrations of the complete unit cell, referred 309 to as lattice modes. Other bands above 400 cm<sup>-1</sup> are attributable to the internal modes of the 310 carbonate ions, the symmetric stretching at 1085 cm<sup>-1</sup> and the in-plane bending at 711 cm<sup>-1</sup> 311 (Wehrmeister et al., 2011). No evidence of amorphous calcium carbonate was observed from 8 312 day-old mice otoconia. The pre-edge peak at 4035.8 eV, corresponding to the electronic transition 313 1s-3d being characteristic for ACC samples (Politi et al., 2006), was absent from the Ca K-edge 314 XAS spectra of the otoconia (Fig. 2c), and only crystalline forms of Ca (calcite) was present. 315



Figure 2. Calcitic mouse otoconia. (a) XRD pattern showing calcite as the mineral phase in mouse otoconia indicating the predominant crystallographic plane reflections. (b) Raman spectrum obtained from a single otoconium. The Raman spectrum was compared with spectra obtained from a Raman database (<u>http://rruff.info/</u>). (c) XAS spectra of wild-type mouse otoconia showing peaks characteristic for calcite. The spectrum was compared with XAS spectra of geological calcite and ACC-Mg obtained under the same experimental imaging conditions.

#### 323 **3.4 Internal structure of otoconia**

324 Figure 3a shows an SEM image of an uncoated, ultramicrotome-cut single otoconium 325 revealing ubiquitous nanostructure throughout its interior. To further analyze and confirm this nanogranular structure, ultramicrotome-cut sections were also examined by AFM. AFM height 326 327 (Fig. 3b) and amplitude (Fig. 3c) analysis revealed an aligned (inset) otoconial nanogranular structure consisting of about 50  $\pm$  14 nm (SD) in Feret diameter. Phase-mode AFM images, as 328 shown in Figure 3d, revealed delineating margins of different composition at the boundaries of the 329 330 nanostructure (arrows). Since there was no evidence of ACC in our samples according to the XRD, Raman and XAS spectroscopy data, at this mouse age (8-day-old), these phase variations are 331 332 attributed to the presence of organic content delineating the nanostructure.



Figure 3. Interior nanostructure of wild-type mouse otoconia. (a) SEM image of the interior of an otoconia from an uncoated otoconium. AFM image in (b) height and (c) amplitude mode showing internal otoconial nanostructure. (d) AFM phase-mode image showing linear arrays of inorganic calcitic nanogranules (yellow, white arrows) surrounded by organic material (red, black arrows). Otoconia were cut/fractured open using a microtome-mounted diamond knife.

340 As examined by TEM, an even smaller nanostructure size of approximately 10 nm in diameter was detected as seen by higher magnification lattice imaging (Fig. 4a). SAED of these 341 regions showed predominantly single-crystal alignment (Fig. 4a, inset), demonstrating highly 342 coherent alignment of the nanostructured subunits. Electron tomography 3D reconstructions of 343 mouse otoconia (Figs. 4b-d, and Supp. Material Movies S2, S3) showed abundant and 344 homogeneously dispersed 10-nm-sized nanodomains together with the larger nanostructure 345 initially observed and measured by AFM (Figs. 3b-d). STEM bright and high-angle annular dark-346 field imaging (HAADF) confirmed the nanostructure of otoconia as shown in Figures S1a,b. 347



Figure 4. Nanostructure of otoconia by TEM after FIB sectioning. (a) High-resolution TEM
lattice imaging of nanodomains, with SAED indicating predominantly single crystal alignment.
(b) Bright-field TEM image from a tilt series of a nanostructured branch region. (c and d) Threedimensional tomographic reconstructions of the nanodomain branch region indicated by the box
in (b) [solid and surface rendering in (c) and (d), respectively].

354 355

Single-crystal 2D XRD of otoconia showed calcitic diffraction rings with rounded spots 356 357 that were similar to those produced by ground geological calcite crystal (optical-quality Iceland spar) which indicate that aggregates of otoconia are generally randomly oriented with respect to 358 359 each other, and that the crystalline phase of individual otoconia behaves like high-quality calcite single crystals (Figs. 5a,b). Despite this single crystal behaviour in diffraction, we observed by 360 SEM, AFM and electron tomography an internal granular nanostructure similar to that observed 361 362 in other calcium carbonate biomineralization systems. Thus, the nanograins forming the otoconia calcite crystals must have a nearly perfect coherent orientation as seen in the SAED patterns (Fig. 363 364 4a, inset). These nanograin results are supported by the data in the 001 pole figures and their 365 contour plots obtained after whole-otoconia EBSD data analysis where the angular spread values for geological calcite and otoconia are quite low and quite similar (0.58 deg and 0.47 degrees, 366 respectively) (Figs. 5c-f). 367



368

**Figure 5. 2D XRD and EBSD analysis of otoconia.** (a) 2D X-ray diffraction pattern produced by otoconia showing single-crystal diffraction spots (white box). (b) Intensity profile along the 104 calcite ring as a function of the  $\gamma$  angle. (c and d) 001 pole figures showing almost identical scattering of the *c*-axis within geological calcite and otoconia, respectively. (e and f) Contour plots of the same data shown in c and d, respectively.

374

### 375 **3.5 Immunogold labeling for OPN in mouse otoconia**

The colloidal-gold immunolabeling imaging approach was used to provide a means for

377 high-resolution localization and mapping of proteins in mineralized tissues by electron microscopy

(McKee and Nanci, 1995). Furthermore, if samples can be cut with microtome, postembedding
immunogold labeling allows protein localization within the interior of biomineralized specimens.
Here, we were able to detect OPN in otoconia by both TEM and SEM in wild-type and *Opn<sup>-/-</sup>* mice,
the latter being used as a negative control where the transgenic mice have been designed to lack
entirely OPN in all their tissues, induced by intentional gene ablation approach.

383 OPN at the surface of the otoconia was readily detected, with Figures 6a-d showing TEM and SEM immunogold labeling for OPN as indicated in wild-type mice. As expected, no labeling 384 was observed in negative-control mice lacking OPN (*Opn*<sup>-/-</sup>) (Figs. 6e-h). A limiting difficulty with 385 386 this approach is that the aqueous procedures necessary to do antibody labeling and other protocol steps particularly for relatively soluble calcite-containing samples, produces essentially a total 387 decalcification of the otoconia and release of mineral-bound proteins into the solution (indeed, this 388 389 is how otoconial proteins can be extracted for biochemical analyses). In the present study, this aqueous dissolution unfortunately resulted in voids in the prepared samples (all panels of Fig. 6) 390 which precluded potential labeling of the interior of the otoconia, but allowed OPN localization at 391 the surface of the otoconia where OPN molecules are trapped locally at the surface by the 392 infiltrated embedding resin and are not affected by the dissolution. 393



394

Figure 6. Immunogold labeling of OPN in otoconia. (a and c) In wild-type mice, otoconial voids 395 396 shown by TEM and SEM, respectively, after immunolabeling for OPN (aqueous procedures dissolve the calcitic otoconia). Inset: Intact wild-type mouse otoconia (without aqueous exposure) 397 398 shown by SEM. By TEM (b) and SEM (d), immunogold labeling for OPN shows gold particles (arrows) at the surface of otoconial voids. (e and g) In OPN-deficient mice, otoconial voids shown 399 by TEM and SEM, respectively, after immunolabeling for OPN. Inset: Intact otoconia from OPN-400 401 deficient mice (without aqueous exposure) shown by SEM. By TEM (f) and SEM (h), as expected in this negative control, immunogold labeling for OPN was absent. 402

#### 403 **3.6 Effect of OPN on calcite crystal growth** *in vitro*

404 To investigate a possible contribution of OPN protein to the production of nanostructure at the surface of otoconia, we examined the effects of OPN on calcite crystal growth in vitro as shown 405 in Figure 6. At a low concentration of OPN (0.15  $\mu$ M), calcite crystals retained their {104} 406 rhombohedral morphology (Fig. 7a), but after increasing the OPN concentration to 0.3 µM and 407 0.45 µM, morphological changes became more evident to produce altered rounded growth step 408 edges (Figs. 7b,c, respectively). At the highest OPN concentration used (0.9 µM), calcite crystals 409 start developing aggregates of {104} rhombohedra (Fig. 6d), all with rough surfaces and apparent 410 surface nanostructure similar to that seen at the surface of otoconia (Fig. 1c). 411



Figure 7. Effect of full-length purified OPN protein on calcite growth *in vitro*. Increasing concentration of OPN (a)  $0.15 \mu$ M, (b)  $0.3 \mu$ M, (c)  $0.45 \mu$ M and (d)  $0.9 \mu$ M gradually alters the external morphology of synthetically grown calcite crystals, producing evident surface nanostructure (inset).

#### 417 **4. Discussion**

The structural and functional relationship between the organic matrix and the mineral 418 component of various biomineralized tissues and structures – including otoconia – is a topic of 419 420 considerable interest in the field of biomineralization. In recent years, reports on amorphous mineral phase precursors and mineral nanostructure have increasingly provided new insights into 421 422 mechanistic processes for biomineralization at the molecular and atomic level. In biomineralized 423 tissues and structures, mineral nanostructure has emerged as a common theme shared by many organisms where nanostructured morphology and mechanical properties can be attributed to the 424 presence of incorporated organic matrix biomolecules (Wolf et al., 2016). 425

For calcitic mammalian otoconia – where in humans there exist major recognized debilitating health consequences linked to otoconial abnormalities and displacement – little is known about the structure of otoconia at the nanoscale. Using high-resolution, nanostructuredetermining 2D and 3D characterization techniques, here we describe that there is an internal calcitic nanostructure in mouse otoconia consisting of domains approximately 50 nm in size with interwoven organic material surrounding individual inorganic calcitic nanostructures.

AFM in the tapping mode and using the phase function can discriminate between inorganic 432 433 and organic phases within complex biomineral structures, being highly sensitive to sample 434 inhomogeneity and compositional variations (Mass et al., 2014). Using this approach to study the interior of otoconia exposed by various means, we observed that the mineral phase consisted of 435 closely packed and aligned nanogranules throughout the otoconial interior, a structural 436 437 arrangement consistent with other descriptions of similar features in numerous calcareous 438 biominerals such as, for example, the sea urchin spicule (Seto et al., 2012), mature tablets of nacre (Hovden et al., 2015), and chicken and guinea fowl eggshells (Athanasiadou et al., 2018; Perez-439

440 Huerta and Dauphin, 2016). Commonly it seems in these cases, the nanogranules are aligned and surrounded by a fine organic matrix, or by a combination of organics and amorphous calcium 441 carbonate (ACC) (Jean-Pierre Cuif, 2010). Notably, most observations on such biomineralized 442 tissues/structures have revealed a nanostructure similar to what we report here for mammalian 443 otoconia, this being roughly spheroidal fields of nanogranules ranging in size from between 50 to 444 445 100 nm (Perez-Huerta and Dauphin (Athanasiadou et al., 2018; Perez-Huerta and Dauphin, 2016). Such results are consistent with there being a nonclassical crystallization pathway to develop this 446 nanostructure that deviates from the classical crystallization pathway (Rodriguez-Navarro et al., 447 448 2016; Wolf et al., 2016), but additional work under cryo-conditions is needed to confirm this. Previous model studies have indicated that organic additives, e.g. acidic biomacromolecules, could 449 favor a nonclassical aggregation-based crystal growth mechanism by stabilizing ACC particles 450 451 (Cölfen, 2008; Wolf et al., 2016). In the present study in 8-day-old mice, ACC was not detected in the otoconia from the Ca-K edge XAS and Raman spectroscopy, but this could be attributed to 452 its loss during sample preparation, or to the relatively mature state of the examined otoconia, where 453 ACC could in fact be present at earlier prenatal or perinatal developmental stages of mouse 454 otoconial development. 455

The smallest nanodomain size that we observed in the otoconia was approximately 10 nm, as observed by HRTEM. Differences in these dimensions could be related to regional variations in organic content (likely type or amount of proteins) acting as inhibitors/regulators of mineralization as has been proposed for the calcitic nanostructure of chicken eggshell *G. gallus domesticus* (Athanasiadou et al., 2018). Previously, *in vitro* crystal growth work by others has shown that OC90, fetuin-A and OPN act as inhibitors of calcite crystal growth, primarily attributable to their having high-binding affinity to calcium (Hong et al., 2015). These results are 463 in agreement with our own calcite crystal growth experiments in which the presence of OPN modifies the typical rhombohedral shape of calcite in a concentration-dependent manner. With 464 increasing OPN concentration, the {104} crystal faces of calcite have abundant growth islands and 465 rounded step edges, and evident surface nanostructure, disclosing changes in the growth 466 mechanism of these crystal faces attributable to the added OPN. Even at higher OPN 467 468 concentrations, the synthetic calcite crystals tend to create aggregates with rough surfaces of slightly mismatched {104} rhombohedra. Although this reflects surface activity, we previously 469 have shown that beyond the surface, additional incorporation of OPN occurs into the crystal 470 471 interior to generate internal nanostructure (Athanasiadou et al., 2018).

In previous *in vitro* calcite growth experiments by others, it was shown that the addition of 472 473 either OC90 or otolin protein alone could not produce the external shape of mature otoconia, but when both proteins were added in combination, they could to some degree produce calcitic 474 structures that resembled otoconial morphology (Moreland et al., 2014). Similar to this, calcite-475 gelatin composites grown by a double-diffusion method in gelatin gel matrices revealed similar 476 morphological features to mammalian otoconia (having branch and belly regions), but these were 477 much larger in size (Huang et al., 2008). Focused-ion beam-cut sections of these artificial 478 479 composites demonstrated a dense crystalline branch region and a poorly crystalline belly region (Simon et al., 2011). 480

Occluded OPN into synthetic calcite crystals can induce a nanostructure similar to the nanostructure observed in calcitic chicken eggshell, where higher OPN concentration creates smaller nanostructure and increases hardness (Athanasiadou et al., 2018). Our finding showing that OPN is concentrated at the surface of the otoconia as detected by high-resolution, postembedding immunogold labeling is novel at the ultrastructural level, and is consistent with 486 previous studies using conventional immunohistochemistry and light microscopy (Sakagami, 2000; Takemura et al., 1994). However, because otoconial calcite is readily soluble under aqueous 487 conditions, dissolution of the mineral rapidly occurs during the required aqueous immunolabeling 488 incubation and washing steps, and we were not able to associate OPN with interior structure (only 489 with the surface region where the embedding medium retained resident components). Thus, while 490 491 likely affecting surface mineralization of the otoconia, it remains possible that OPN is also present within the interior of otoconia and contributes to the nanostructure that we have observed here, 492 and which can be reproduced *in vitro* using OPN. Indeed, otoconins such as OC90 could likewise 493 494 participate in this nanostructuring process. OPN is co-expressed locally in the inner ear with OC90 by the vestibular dark cells and in the endolymphatic sac (Ignatova et al., 2004; Sakagami, 2000; 495 Verpy et al., 1999), meaning that OC90 and OPN could have similar functions in some regard, 496 even partially compensating for each other during otoconial formation. In particular, OC90 has a 497 strong effect on otoconial formation; OC90-knockout mice have rodlike large calcitic aggregates 498 that are susceptible to dissolution (Zhao et al., 2007). In contrast, in OPN-null mice where Oc90 499 levels are presumably not altered, there is normal balance behavior and normal otoconial 500 morphology (Zhao et al., 2008). Since OPN has now been localized to the very periphery of 501 502 otoconia in mice (our work) and in rats (Takemura et al., 1994), and given its role in creating mineral nanotexture as we have recently shown for calcite crystals in the chicken eggshell, OPN 503 could indeed be a protein involved in controlling surface growth and nanostructure in otoconia. 504 505 However, the interplay between OPN and OC90, if any, along with interactions amongst other otoconial proteins in this biomineralization process, remains to be determined. 506

507

#### 509 **5. Conclusions**

510 In summary, the internal structure of mouse otoconia at the nanoscale has been determined. Within the interior of otoconia, using a combination of advanced imaging techniques, we show 511 that otoconia consist of densely packed calcite nanodomains surrounded by organic material, with 512 513 OPN being present at the very surface of the otoconia. The surface localization of OPN suggests 514 involvement in the growth process and/or termination of otoconial growth via the mineralizationinhibiting activity of this protein on crystal growth. This study provides detailed findings on the 515 516 internal and external structure of otoconia, information that may lead to a better understanding of 517 what goes wrong with otoconia in diseases such as vertigo.

518

#### 519 Acknowledgments

520 We are grateful to Ms. Lydia Malynowsky and Dr. Betty Hoac for otoconia dissections. We also thank Dr. Kelly Sears, Dr. David Liu, Ms. Weawkamol Leelapornpisit, Dr. Kaustuv Basu 521 522 and Prof. Khanh Huy Bui for assistance with FIB, TEM and tomography work, and Dr. Rui Tahara for assistance with Xradia 520 Versa imaging. We would like also to thank the Soft X-ray 523 Microcharacterization Beamline (SXRMB) 06B1-1 at the Canadian Light Source (CLS) for their 524 525 technical support, and the Canadian Light Source (CLS) for financial travel support. We also gratefully thanks Dr. Nicole Hondow and Dr. Zabeada Aslam of the Leeds Electron Microscopy 526 and Spectroscopy Centre. This work was supported by Canadian Institutes of Health Research 527 528 (MOP-142330) and the Natural Sciences and Engineering Research Council of Canada (RGPIN-2016-05031) to MDM. MDM is a member of the FRQ-S Network for Oral and Bone Health 529 530 Research. This research was undertaken, in part, thanks to funding from the Canada Research 531 Chairs program.

#### 533 6. Supplementary Material



- **Fig. S1** Otoconia nanostructure observed by STEM. (a) Bright-field and (b) HAADF TEM
- 536 images from a STEM tilt series where nanostructure is clearly evident.
- 537 Movie S1. Three-dimensional reconstruction of utricular otoconia (otoconia in grey, and bone in
   538 orange) from an X-ray computed tomography series.
- 539 Movie S2. Three-dimensional reconstruction from the branch region of an 8-day-old mouse
   540 utricular otoconium from a TEM tomography tilt series.
- 541 Movie S3. Three-dimensional reconstruction by solid- and surface-rendering of a nanostructured
  542 area from the branch region of an 8-day-old mouse utricular otoconium from a TEM tomography
  543 tilt series.

#### 553 **References**

- Addadi, L., et al. 2003. Taking advantage of disorder: amorphous calcium carbonate and its roles
   in biomineralization. Adv. Mater. 15, 959-970.
- Agrawal, Y., et al. 2009. Disorders of balance and vestibular function in us adults: data from the
   national health and nutrition examination survey, 2001-2004. Arch. Intern. Med. 169,
   1419-1419.
- Andrade, L.R., et al. 2012. Immunogold TEM of otoconin 90 and otolin relevance to
   mineralization of otoconia, and pathogenesis of benign positional vertigo. Hear. Res. 292,
   14-25.
- 562 Anniko, M., et al. 1984. Microprobe analysis of human otoconia. Acta Oto-Laryngol. 97, 283-289.
- Athanasiadou, D., et al. 2018. Nanostructure, osteopontin, and mechanical properties of calcitic
  avian eggshell. Sci. Adv. 4, eaar3219.
- 565 Carlstrom, D.D., 1963. Crystallographic study of vertebrate otoliths. Biol. Bull. 125, 441-463.
- Cölfen, H., Antonietti, M., 2008. Mesocrystals and nonclassical crystallization. John Wiley &
   Sons, Ltd.
- De Yoreo, J.J., et al. 2015. Crystal growth. Crystallization by particle attachment in synthetic,
   biogenic, and geologic environments. Science 349, aaa6760.
- Fermin, C.D., et al. 1995. The glycan keratan sulfate in inner ear crystals. Cell Mol. Biol. 41, 577591.
- Fisher, L.W., et al. 2001. Flexible structures of SIBLING proteins, bone sialoprotein, and osteopontin. Biochem. Bioph. Res. Comm. 280, 460-465.
- Hiatt, G.A., 2014. Color atlas and text of histology. Wolters Kluwer.
- Hong, M.N., et al. 2015. Effect of otoconial proteins fetuin a, osteopontin, and otoconin 90 on the
  nucleation and growth of calcite. Cryst. Growth Des. 15, 129-136.
- House, M.G., Honrubia, V., 2003. Theoretical models for the mechanisms of benign paroxysmal
  positional vertigo. Audiol. Neurotol. 8, 91-99.
- Hovden, R., et al. 2015. Nanoscale assembly processes revealed in the nacroprismatic transition
  zone of Pinna nobilis mollusc shells. Nat. Commun. 6, 1-7.
- Huang, Y.X., et al. 2008. Shape development and structure of a complex (otoconia-like?) calcitegelatine composite. Angew. Chem. Int. 47, 8280-8284.
- Hughes, I., et al. 2006. Mixing model systems: using zebrafish and mouse inner ear mutants and
  other organ systems to unravel the mystery of otoconial development. Brain Res. 1091, 5874.
- Ignatova, E.G., et al. 2004. Molecular mechanisms underlying ectopic otoconia-like particles in
   the endolymphatic sac of embryonic mice. Hear. Res. 194, 65-72.
- 588 Cuif, J.-P., et al. 2010. Biominerals and fossils through time. Cambridge university press,
   589 Cambridge.
- Johnsson, L.G., et al. 1982. Pathology of neuroepithelial suprastructures of the human inner-ear.
   Am. J. Otolaryngol. 3, 77-90.
- Kremer, J.R., et al. 1996. Computer visualization of three-dimensional image data using IMOD. J.
   Struct. Biol. 116, 71-76.
- Lins, U., et al. 2000. The otoconia of the guinea pig utricle: internal structure, surface exposure,
  and interactions with the filament matrix. J. Struct. Biol. 131, 67-78.
- 596 Lowenstam, H.A., Weiner, S., 1989. On biomineralization. Oxford university press, New York.
- Lu, W., et al. 2010. In vitro effects of recombinant otoconin 90 upon calcite crystal growth.
  Significance of tertiary structure. Hear. Res. 268, 172-183.

- Lundberg, Y.W., Xu, Y., 2012. Proteins involved in otoconia formation and maintenance, in:
  Gendeh, B.S., (Ed.), Otolaryngology, InTech, pp. 1-22.
- Lundberg, Y.W., et al. 2015. Mechanisms of otoconia and otolith development. Dev. Dynam. 244,
   239-253.
- Marieb, E.N., 1991. Essentials of human anatomy and physiology, Essentials of human anatomy
   and physiology.
- Mass, T., et al. 2014. Immunolocalization of skeletal matrix proteins in tissue and mineral of the
   coral Stylophora pistillata. Proc. Natl. Acad. Sci. USA 111, 12728-12733.
- Moreland, K.T., et al. 2014. In vitro calcite crystal morphology is modulated by otoconial proteins
   otolin-1 and otoconin-90. Plos One 9, e95333.
- Murayama, E., et al. 2005. Otolith matrix proteins OMP-1 and Otolin-1 are necessary for normal
   otolith growth and their correct anchoring onto the sensory maculae. Mech. Develop. 122,
   791-803.
- Oas, J.G., 2001. Benign paroxysmal positional vertigo: a clinician's perspective. Ann. N. Y. Acad.
   Sci. 942, 201-209.
- Ornitz, D.M., et al. 1998. Otoconial agenesis in tilted mutant mice. Hear. Res. 122, 60-70.
- Perez-Huerta, A., Dauphin, Y., 2016. Comparison of the structure, crystallography and composition of eggshells of the guinea fowl and graylag goose. Zoology. 119, 52-63.
- Politi, Y., et al. 2006. Structural characterization of the transient amorphous calcium carbonate
   precursor phase in sea urchin embryos. Adv. Funct. Mater. 16, 1289-1298.
- Pote, K.G., Ross, M.D., 1986. Ultrastructural morphology and protein-content of the internal
   organic material of rat otoconia. J. Ultra. Mol. Struct. Res. 95, 61-70.
- Pote, K.G., Ross, M.D., 1991. Each otoconia polymorph has a protein unique to that polymorph.
  Comp. Biochem. Phys. B 98, 287-295.
- Pote, K.G., Ross, M.D., 1993. Utricular otoconia of some amphibians have calcitic morphology.
  Hear. Res. 67, 189-197.
- Rodriguez-Navarro, C., et al. 2016. Nonclassical crystallization in vivo et in vitro (II):
   nanogranular features in biomimetic minerals disclose a general colloid-mediated crystal
   growth mechanism. J. Struct. Biol. 196, 260-287.
- Ross, M.D., Pote, K.G., 1984. Some properties of otoconia. Philos. T. Roy. Soc. B 304, 445-452.
- Sakagami, M., 2000. Role of osteopontin in the rodent inner ear as revealed by in situ
   hybridization. Med. Electron Microsc. 33, 3-10.
- Salvinelli, F., et al. 2004. Benign paroxysmal positional vertigo: diagnosis and treatment. Clin.
   Ter. 155, 395-400.
- Seto, J., et al. 2012. Structure-property relationships of a biological mesocrystal in the adult sea
  urchin spine. Proc. Natl. Acad. Sci. USA 109, 7126-7132.
- Simon, P., et al. 2011. Structural relationship between calcite-gelatine composites and biogenic
   (human) otoconia. Eur. J. Inorg. Chem. 5370-5377.
- 637 Sodek, J., et al. 2000. Osteopontin. Crit. Rev. Oral Biol. M 11, 279-303.
- Sorensen, E.S., Petersen, T.E., 1993. Purification and characterization of 3 proteins isolated from
   the proteose peptone fraction of bovine-milk. J. Dairy Res. 60, 189-197.
- Sorensen, E.S., et al. 1995. Posttranslational modifications of bovine osteopontin: identification of
   twenty-eight phosphorylation and three O-glycosylation sites. Protein Sci. 4, 2040-2049.
- Tachibana, M., Morioka, H., 1992. Glucuronic acid-containing glycosaminoglycans occur in
   otoconia: cytochemical evidence by hyaluronidase-gold labeling. Hear. Res. 62, 11-15.

- Takemura, T., et al. 1994. Localization of osteopontin in the otoconial organs of adult rats. Hear.
   Res. 79, 99-104.
- Thalmann, I., et al. 2006. Microscale analysis of proteins in inner ear tissues and fluids with
  emphasis on endolymphatic sac, otoconia, and organ of Corti. Electrophoresis 27, 15981608.
- Tseng, Y.H., et al. 2014. CaCO<sub>3</sub> nanostructured crystals induced by nacreous organic extracts.
   Cryst. Eng. Comm. 16, 561-569.
- Verpy, E., et al. 1999. Characterization of otoconin-95, the major protein of murine otoconia,
   provides insights into the formation of these inner ear biominerals. Proc. Natl. Acad. Sci.
   USA 96, 529-534.
- Vibert, D., et al. 2003. Benign paroxysmal positional vertigo in older women may be related to
   osteoporosis and osteopenia. Ann. Oto. Rhinol. Laryn. 112, 885-889.
- Walther, L.E., et al. 2014. The inner structure of human otoconia. Otol. Neurotol. 35, 686-694.
- Wehrmeister, U., et al. 2011. Amorphous, nanocrystalline and crystalline calcium carbonates in
   biological materials. J. Raman Spectrosc. 42, 926-935.
- Wolf, S.E., et al. 2016. Nonclassical crystallization in vivo et in vitro (I): process-structure property relationships of nanogranular biominerals. J. Struct. Biol. 196, 244-259.
- Wright, C.G., et al. 1982. A calcareous concretion in the posterior semicircular duct of a human
  labyrinth. Am. J. Otolarynogol. 3, 196-201.
- Ku, Y., et al. 2010. Expression, functional, and structural analysis of proteins critical for otoconia
   development. Dev. Dyn. 239, 2659-2673.
- Yang, H., et al. 2011. Matrix recruitment and calcium sequestration for spatial specific otoconia
   development. Plos One 6, e20498.
- Zhao, X., et al. 2007. Gene targeting reveals the role of Oc90 as the essential organizer of the
   otoconial organic matrix. Dev. Biol. 304, 508-524.
- Zhao, X., et al. 2008. Osteopontin is not critical for otoconia formation or balance function. J.
   Assoc. Res. Otolaryngol. 9, 191-201.