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Abstract. Sound processing in the inner ear involves separation of the constituent frequencies along the length of the cochlea. Frequencies relevant to human speech (100 to 500 Hz) are processed in the apex region. Among mammals, the guinea pig cochlear apex processes similar frequencies and is thus relevant for the study of speech processing in the cochlea. However, the requirement for extensive surgery has challenged the optical accessibility of this area to investigate cochlear processing of signals without significant intrusion. A simple method is developed to provide optical access to the guinea pig cochlear apex in two directions with minimal surgery. Furthermore, all prior vibration measurements in the guinea pig apex involved opening an observation hole in the otic capsule, which has been questioned on the basis of the resulting changes to cochlear hydrodynamics. Here, this limitation is overcome by measuring the vibrations through the unopened otic capsule using phase-sensitive Fourier domain optical coherence tomography. The optically and surgically advanced method described here lays the foundation to perform minimally invasive investigation of speech-related signal processing in the cochlea. © The Authors. Published by SPIE under a Creative Commons Attribution 3.0 Unported License. Distribution or reproduction of this work in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.JBO.21.2.025003]

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1 Introduction

Sound entering the ear stimulates the cochlea and launches a fluid-structure traveling wave along the structures in the inner ear. The constituent sound frequencies are separated along the length of the cochlea with high frequencies near the base, which is closer to the middle ear, and low frequencies near the apex. The sound-induced vibrations inside the cochlea are highly complex in a living mammal and are vulnerable to damage. These vibrations could, therefore, provide important information on the mechanisms of normal and impaired hearing. Most previous measurements of inner ear vibrations were limited to the first surface in the optical view, which is the basilar membrane in the basal turn and the reticular lamina or the tectorial membrane in the apical turn, upon opening the bony otic capsule.1 Furthermore, reflective beads were introduced to enhance the signal-to-noise ratio, as inner ear tissues have poor reflectivity.

Phase-sensitive Fourier domain optical coherence tomography (PhS-FDOCT) is ideal to measure inner ear vibrations because it can detect vibrations of samples with very low native reflectance without the need to introduce reflective objects. Vibrations on the order of a fraction of a nanometer can be measured across tissue and over a wide frequency range. Additionally, it provides the capability to simultaneously image the structure to identify where the vibrations are detected. This technique has recently been used to measure inner ear vibrations without opening the cochlea in the hook region of the guinea pig cochlea,2 hook and next turn in the mouse cochlea,3 and in the mouse apical turn.4

The fundamental frequency of the human voice ranges from 100 to 300 Hz, and a large part of the frequency range used in music has fundamental frequencies and at least some harmonics at frequencies below 1000 Hz. The ability to hear a low stimulus frequency is therefore fundamental for speech and music perception, and, indeed, oral communication becomes difficult if low-frequency hearing is disturbed. It is believed that the cochlear processing of low frequencies in the apex is different from the processing of high frequencies in the basal region.5 The different behavior in the apex could be concomitant with the fact that the auditory nerve phase-locks to sound stimulus at low frequencies (below ~4000 Hz), whereas it is not phase-locked at higher frequencies.

While much information from physiologically intact preparations is available from the high-frequency or basal regions of the cochlea,1,3,5 this is not the case for the apical regions. One problem is that previous techniques required opening the cochlea, which is doable at high frequencies (because acoustic
wavelengths are smaller than the cochlear dimensions), but the response of the apical turn may be altered by such observation windows in the bone (e.g., Ref. 7). The problem is compounded by the anatomical inaccessibility of the guinea pig cochlear apex, which has meant that previous studies were either performed in vitro, or in preparations where an extensive and potentially damaging surgical approach was necessary. The history of cochlear imaging and interferometry is discussed in detail in Ref. 8.

We have developed measurement techniques and surgical approaches that improve this situation. The method described here enables us, for the first time, to measure the cochlear mechanical response to frequencies <1000 Hz in the absence of artificial reflectors or observation windows through the bone. The method is based on two technical advancements: (1) optical, to allow optical penetration into the cochlea, in particular to the organ of Corti and tectorial membrane, without opening the otic capsule and (2) anatomical, to allow measurements apically along the cochlear axial direction with reduced surgery compared with that used thus far. The optical development is based on PhS-FDOCT and the anatomical development on the novel idea of using a mirror to direct the beam axially, thus eliminating the need for ventral surgery. Since the current surgery is less traumatic and the fluid boundary conditions are unaltered, these two developments together enable gathering of important, less artifact-prone vibration data for understanding sound processing in the low-frequency region of the cochlea.

2 Materials and Methods

2.1 Ethical Approval

Measurements reported in this article were made in vivo in normal-hearing albino guinea pigs of either sex (250 to 350 g). Albino animals are expected to have the same normal cochlear mechanics as pigmented, and the lack of pigmentation reduces light scatter in the cochlea. All procedures in this study were approved by the Institutional Animal Care and Use Committee of Oregon Health and Science University.

2.2 Experimental Setup

The schematic of the PhS-FDOCT system setup used in this study is shown in Fig. 1. The optical source is a broadband superluminescent diode (SLD, SuperLum) with a central wavelength of 840 nm and a spectral bandwidth of 45 nm. The light from the SLD is coupled into a fiber-based Michelson interferometer via an optical circulator with a coupling ratio of 90:10. In the reference arm, the light is delivered onto a stationary mirror. The light in the sample arm is focused onto the sample via an objective lens with a focal length of 30 mm. The sample is scanned along the x-direction by a galvo mirror (GVSM002, Thor Labs) to acquire reflectance image using M-B mode scanning.

A 2×2 optical fiber coupler recombines the light backscattered from the sample and reflected from the reference mirror. This combined beam is routed to a purpose-built, high-speed spectrometer via the optical circulator. The spectrometer consists of a 30-mm focal length collimator, a 1200-lines/mm transmission grating, an achromatic lens with 100 mm focal length, and a 12-bit, 4096-pixel CMOS line-scan camera (Basler Sprint spL4096-140 km), of which only 2048 pixels are used, to reduce data-acquisition time. To improve the signal-to-noise ratio, we use vertical binning as well as horizontal binning, both over 2 pixels, which effectively yields 1024 pixels. The maximum line-scan rate of the camera is 140 kHz. We use a reduced sample rate of 10 kHz to improve the signal-to-noise ratio, as the maximum frequency of interest is <1 kHz in this study.

The coherence length and central wavelength of the light source determine the axial resolution, which, in the current instrument, is ~5 μm in tissue, an advantage of the 840-nm source. The axial resolution is calculated using the formula

\[
l_c = \frac{4 \ln(2) \lambda_0^2}{\pi n \Delta \lambda} = 6 \ \mu m
\]

in air and 4.3 in tissue for \( n = 1.4 \). In PhS-FDOCT, the entire penetration depth of the beam is used at once to image along the optical axis of the system. To generate images of the cochlear structures, a computer-controlled galvo mirror moves the beam across the preparation. Once the organ of Corti is identified, the phase and amplitude of sound-evoked motion is determined for every image pixel. Due to the large penetration depth of infrared light, the technique makes it possible to image and measure the vibration of all cellular and noncellular structures of the organ of Corti in the apical turn. The good penetration of the beam makes opening of the cochlea unnecessary. Vibrations are measured over a wide frequency range, and structures are simultaneously imaged, which allows us to determine the sound-evoked vibration of all cells within the organ of Corti, not only those at the surface.

2.3 Surgical Preparation

Young guinea pigs weighing 250 to 350 g were anesthetized by an intraperitoneal injection of a combination of ketamine (40 mg/kg) and xylazine (10 mg/kg). Supplemental 1/2 doses were given every hour or when need was indicated by paw withdrawal to toe pinch. The guinea pig was mounted on a heated head holder. A tracheotomy and placement of a short section of ventilation tubing allowed free breathing. No artificial respiration was needed, and motion from respiration or cardiovascular vibrations did not significantly affect the measurement noise floor. Surgical operation was performed to expose the left
bulla, which was then incised to open for access to the cochlea.\textsuperscript{10} The surgical dissection to expose the auditory bulla was from a ventral direction, and the bulla was widely opened to expose the cochlea but preserve the tympanic membrane. With a widely opened bulla, the apex of the cochlea becomes visible. During the surgery as well as during the experiment, the core temperature of the animal was maintained at 37 to 38 °C by a heating blanket and a rectal thermometer, controlled by a servo temperature controller (FHC, Bowdoinham, Maine).

### 2.4 Measurement in Axial and Radial Directions

For apex vibration measurements in the axial direction, it is not possible to simply adjust the head orientation in a guinea pig. The reason is evident from the location of the bulla and the cochlea inside the skull shown in Fig. 2. Because the axial viewing direction is occluded by the jaw and associated neck tissue, extensive surgery is needed to remove part of the jaw and musculature to expose the apex in the axial direction. However, such extensive surgery takes significant time and adds physiological stress and blood loss, which can affect the viability of the guinea pig for long-duration studies while under anesthesia. In addition, the hearing sensitivity could be affected by extensive surgery, and we lack easy-to-use and reliable electrophysiological indicators of cochlear function at these low frequencies (unlike compound action potential and cochlear microphonics at high frequencies for basal turn studies;\textsuperscript{10} see also Ref. 11).

To overcome this issue and gain access to the axial direction, we use a mirror device that has two right-angled faces with large openings and a reflective front surface mirror is attached to a 45-deg slanted face [Fig. 3(c)]. This device is placed inside the live animal’s bulla facing the cochlea, as shown in Fig. 3(d) and schematically in Fig. 3(a). Because the light is deflected by 90 deg inside the bulla, surgical removal of the jaw and neck tissue is not necessary. This makes it possible to measure vibration using a standard surgical approach developed for the basal turn, a very significant advantage, since extensive dissection is avoided. The placement of the mirror device inside the bulla needs to be done carefully in order to get correct orientation to access the tissues in the cochlear apex. We placed the device inside the bulla by viewing under a dissection microscope and oriented the device such that the top plastic surface was horizontal, as shown in Fig. 3(a).

To measure the vibrations in the radial direction, the animal’s head orientation was adjusted and the light from the sample arm was directed at the apex as shown in Fig. 3(b) without the use of the mirror device. The depth of the inner ear structures to be viewed in this direction is larger compared to the depth in the axial direction. Therefore, the otic capsule is thinned, but not opened, in older animals to improve the signal-to-noise ratio. In younger animals (less than five weeks of age), the native bone is already sufficiently thin even for the radial vibration measurement.

### 2.5 Sound Stimulation and Data Acquisition

All sound stimuli were delivered to the ear through a custom speculum fitted tightly into the ear canal to form a closed sound field. An earbud speaker was used to generate the sound. The pressure at the ear canal was precalibrated using a microphone (Bruel and Kjaer type 4138). The data acquisition was done using custom software codes written in Lab View software (National Instruments), which generates the voltage signal to drive the speaker as well as recording the data acquired by the line-scan camera using a camera link. The input voltage to the speaker was adjusted to deliver a constant sound level to the ear. The data recorded by the camera were a function of the optical wavenumber and contained information about the reflectance and vibration of the sample. The postprocessing method described in Ref. 12 was used to derive vibration and reflectance image from the recorded data. The lowest detectable vibration depends on several factors such as the reflectivity of the tissue, animal-to-animal variability, camera sampling rate used, number of averages, etc. For the parameters used in this study (100 to 500 Hz, 40 averages, 10-kHz camera sampling rate, or 4 s of averaging time), and for the guinea pig apex organ of Corti through the otic capsule, the lowest detectable vibration amplitudes were around 1 nm. Only displacements with signal higher than the mean noise floor by at least five standard deviations were considered for analysis.

The displacement amplitude is given by

\[
A_D = \frac{A_0}{\Phi} \frac{\Phi}{4\pi n}
\]

where \(A_D\) is the displacement amplitude, \(A_0\) is the center wavelength of the light source, \(\Phi\) is the phase of the depth-resolved optical signal, and \(n\) is the sample refractive index. The largest displacement that can be measured without phase wrapping is equal to a quarter wavelength of the optical source for \(n = 1\). In order to reduce optical noise, as well as to improve this maximum displacement limit, velocity is derived via time differentiation of the raw phase data as discussed in Ref. 12. The time differentiation is done numerically by deriving the phase difference between adjacent A-scans. The velocity magnitude is then given by

\[
A_V = \frac{A_0}{\Phi} \frac{\Phi}{4\pi n} S
\]

where \(S\) is the camera sampling rate. The maximum velocity measurable without phase wrapping is therefore given by

\[
A_{V_{max}} = \frac{A_0}{4\pi n S}
\]
For the detector, we used the Basler Sprint spL4096-140 km CMOS line-scan camera with a maximum sampling rate of 140 kHz. However, because this study was done at low frequencies $<1$ kHz, we operated the camera at a reduced rate of 10 kHz to take advantage of the larger signal-to-noise ratio. Reducing the camera rate further would limit the maximum vibration that can be measured without phase wrapping. In this study, $\lambda_0 = 840$ nm. This gives a maximum velocity limit of 0.21, 2.1, and 21 mm/s for camera sampling rates of 1, 10, and 100 kHz, respectively. Figure 4 shows the corresponding maximum displacements that can be measured using PhS-FDOCT with center wavelength of 840 nm and for camera sampling rates $S$ indicated in the legend. Note that a quarter wavelength of the optical source is 210 nm. The sound-induced displacements in the guinea pig apex can approach several hundred nanometers for 90 dB SPL stimulus level, as shown in (b). In the sample axial frequency response shown in (b), the maximum displacement at 94 dB SPL (or 1 Pascal) is 300 nm at 300 Hz. This value can be higher in some animals.
displacement limit versus stimulus frequency for the three camera sampling rates. For the 90 dB SPL sound stimulus used in this study, the vibrations in the guinea pig apex can be as large as 250 to 500 nm below 300 Hz and lower at higher frequencies, as shown by a typical recording in Fig. 4(b). Therefore, we chose to use a 10 kHz camera sampling rate in this study.

3 Results and Discussion

3.1 Vibrations in the Axial Direction

As described in Sec. 2, a mirror device placed inside the live guinea pig bulla is used to measure the inner ear vibrations in the guinea pig apex through the otic capsule in the axial direction. The reflectance (structural) image is shown in Fig. 5(a), and a schematic is shown in Fig. 5(c) for comparison. The Reissner’s membrane, basilar membrane, and the organ of Corti can be identified in the reflectance image. The tectorial membrane appears faint or sometimes not easy to discern. This could be improved by thinning the otic capsule. The axial displacement response in nanometers at 90 dB SPL, 200 Hz is shown in Fig. 5(b). The displacement is higher in the organ of Corti compared to the displacement at the basilar membrane. The Reissner’s membrane also has a high displacement response. The response pattern has important physiological implications and is indicative of the status of the inner ear.

For the reflectance image shown in Fig. 5(a), 1000 points are used across the radial direction with 200 time averages. For the vibration pattern shown in Fig. 5(b), 25 points are used across the radial direction with 40 time averages. A smaller number of lateral points is used for the vibration recording to save recording time.

3.2 Vibrations in the Radial Direction

The vibrations in the radial direction are measured without using the mirror device by positioning the animal’s head as shown in Fig. 3(b). Because the depth of the inner ear structures to be viewed in this direction is larger, the otic capsule may need to be thinned (but not opened) in older animals to improve the signal-to-noise ratio. For the data shown in Fig. 6, thinning the

![Fig. 5 PhS-FDOCT measurement of the guinea pig apex in the axial direction.](image-url)
The reflectance (structural) image viewed through the bone in the radial direction is shown in Fig. 6(a), and the schematic oriented similarly is shown in Fig. 6(c). Although the deeper structures appear faint, their vibrations can be detected above noise. The radial displacement in nanometers for a 90 dB SPL, 150 Hz stimulus is shown in Fig. 6(b). These data are from a guinea pig different from that shown in Fig. 5. The radial vibration pattern also contains information on the inner ear physiology.

4 Conclusions and Future Directions

This paper describes the method we have developed to measure in vivo vibrations in the apex of the guinea pig cochlea through the otic capsule in two directions (axial and radial) using PhS-FDOCT. A mirror device is placed inside the cochlear bulla to significantly reduce the surgery needed to access the cochlear apex in the axial direction. This method has enabled, for the first time, the measurement of cochlear response to frequencies <1000 Hz in the absence of artificial reflectors or observation windows through the otic capsule. Although the primary sound-induced excitation for the cochlear structures is in the axial direction, it has been hypothesized\textsuperscript{13} that the tectorial membrane resonates radially in response to sound. Models have shown that this resonance could play an important role in active feedback amplification in the cochlea.\textsuperscript{14–16} The tectorial membrane resonance hypothesis has been supported by in vitro experiments in the apex,\textsuperscript{17} but it has not been tested experimentally in vivo. The method developed in this study will enable us to address this and other questions in unopened cochleae of guinea pigs. The apical turn of the guinea pig cochlea has the best frequencies around 100 to 500 Hz. Investigating this region is important to understand how the cochlea processes speech signals.

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Biographies for the authors are not available.