INTRODUCTION

Resident macrophages in the cochlea have been intensively investigated using advanced immunohistochemistry and laser scanning microscopy techniques. Many studies have reported that cochleae harbor resident macrophages expressing CD45, F4/80, ionized calcium-binding adapter molecule 1, CD11b, and C-X3-C motif chemokine receptor 1 (CX3CR1) [1-4]. These macrophages play important functions in inflammation in response to various insults, including noise exposure, ototoxicity, and infection. Interestingly, some studies have shown that increased numbers of macrophages in response to these insults stem from migrating monocytes, not from pre-existing macrophages [2,5].

According to previous studies, recruited macrophages are distributed widely throughout the cochlea, and resident macrophages are thought to be mainly located in the spiral ligament and stria vascularis [2,3,6]. A few macrophages have been found in Reissner’s membrane and in the basement membrane. Importantly, recruited macrophages can increase significantly in number in the spiral ligament, Reissner’s membrane, and basement membrane and spread to the wall of the scala tympani, scala vestibuli, and spiral limbus. The inferior part of the spiral ligament is believed to be the extracellular migration site of immune cells, as immunohistochemical findings have highlighted a high concentration of macrophages in this area pre- and post-inflammation [2,5].

The spiral ligament is divided into five areas according to the type of fibrocyte that it is composed of. Each type of fibrocyte expresses different cell markers suggestive of different roles. For example, the NKCC1 channel is expressed in type II, IV, and V...
fibrocytes, and the aquaporin 1 channel is expressed in type III fibrocytes. Na/K-ATPase, which is also an important ion channel for generating the potassium gradient, is highly expressed in type II and V fibrocytes [7,8].

In this study, we sought to document macrophage distribution patterns in the spiral ligament in CX3CR1−/GFP mice, in which monocytes and macrophages in the cochlea can be visualized without immunostaining according to previous studies [9,10]. As conventional cochlear sectioning methods have limitations in terms of generalizing the location of cochlear macrophages due to the thin depth of the section, we applied a novel three-dimensional imaging technique without the need for any tissue preparation. Three-dimensional cochlear imaging was conducted using two-photon microscopy, which has a higher penetration depth and poses less damage to the tissue than confocal microscopy [11,12].

MATERIALS AND METHODS

Animals

CX3CR1−/GFP C57/BL6 mice were purchased from the Jackson Laboratory (Bar Harbor, ME, USA) via Orient Bio (Seongnam, Korea). All mice were maintained in a specific pathogen free environment in the animal facility at Avison Biomedical Research Center at Yonsei University College of Medicine (Seoul, Korea). Temperature, humidity, and light/dark cycle were properly controlled, and food and water were accessible ad libitum. Less than five animals were maintained in a single cage. CX3CR1−/GFP mice were produced by crossing wild-type C57/BL6 mice (Orient Bio) with CX3CR1−/GFP mice. Randomly selected male and female mice (aged 4-8 weeks and weight 15–25 g) were used in the experiments. All animals were carefully maintained in cages with area of 25,000 µm². The concentration and amount of injected LPS were based on previous studies [13,14]. The experimental group mice were sacrificed at 1, 4, 7, and 10 days after LPS inoculation. Control group mice were sacrificed without any injection or sacriﬁed 4 days after 5 µl of PBS inoculation in the middle ear. The right side ear was used in all experimental animals. All experiments were performed during the daytime in the animal laboratory. A heating pad (37°C) was used to maintain the body temperature of the animal during general anesthesia. The animal experiments were approved by the Institutional Animal Care and Use Committees at the Yonsei University College of Medicine (Project No. 2019-0182).

Cochlear preparation

After sacrificing the mice in a CO₂ chamber, the bilateral temporal bones of the mice were obtained. The mouse temporal bones were fixed in 4% paraformaldehyde for 24 hours in a 4°C room. After fixation, samples were incubated in 1 mL of 0.25 M ethylenediaminetetraacetic acid (EDTA) solution for 24 hours at 4°C for decalcification. Decalcified cochlea were cleared using Binaree Clearing Kits (Binaree, Daegu, Korea). Cleared cochlea were mounted on a silicone bond in the petri dish, with spiral ligaments were positioned at the top. Eighty percent of glycerol was then dropped on the sample, and a cover slide was positioned on it. Distilled water was filled between cover slide and water-immersing lens of the two-photon microscopy.

Imaging

Two-photon microscopy (LSM7MP; Carl Zeiss, Oberkochen, Germany) was used for imaging data generation. Zen software (Carl Zeiss) was used for image acquisition and basic image analysis. To create three-dimensional images and videos, we used IMARIS software (Bitplane, Zurich, Switzerland). The Mai-Tai HP Ti:Sapphire Deep See laser system (Spectra-Physics, Mountain View, CA, USA) was used to generate excitation lasers with a wavelength of 810 nm. Images were acquired at a resolution of 512 x 512 pixels using band-pass filters with 420–480 nm (blue), 500–550 nm (green), and 575–610 nm (red). A 20 × objective lens was used in all experiments.

Quantification and statistical analysis

Quantification was performed using Zen software (Carl Zeiss), and the type III fibrocyte area, which is parallel to X-Y plane, was investigated. Regions of interest (ROIs) were drawn as a circle with area of 25,000 µm² lateral to the stria vascularis. The number of green cells were counted manually between cortical bone and the stria vascularis in each ROI. Kruskal-Wallis test with post-hoc was used to evaluate the significance (P<0.05) of differences between groups. Statistical analysis was performed by IBM SPSS ver. 25.0 (IBM Corp., Armonk, NY, USA).

RESULTS

Three-dimensional imaging of intact cochlea

We obtained three-dimensional images of intact cochleae from
CX3CR1+GFP mice using two-photon microscopy (Fig. 1, Supplementary Video 1). The cochlear bony capsule was colored blue, and capillaries were colored white because of endogenous fluorescence. Spiral ligaments appeared as dim endogenous fluorescence, whereas the stria vascularis showed brighter fluorescence. Capillaries in the spiral ligament, including the stria vascularis vessels, were clearly seen with their characteristic branching pattern. Resident macrophages appeared in bright green color. Even though the intensity of fluorescence grew dimmer as the depth increased, macrophages in the spiral ligament were clearly distinguishable.

Macrophage distribution in untreated cochlea

To reduce selection bias, we captured extended images of 100 µm in length along the cochlear turn (images were projected to the sagittal plane). The apical and basal turns showed similar patterns of macrophage distribution (Fig. 2A and B). Notably, the spiral ligament macrophages were not evenly distributed in the sagittal plane. The lateral edge of the spiral ligaments showed dense distributions of macrophages. Perivascular macrophages contacting stria vascularis vessels were also found, and several macrophages were seen at Reissner’s membrane. However, resident macrophages were scarce in the area adjacent to the stria vascularis. In
particular, the wide area from the crista basilaris to beneath the stria vascularis showed a lack of macrophages, even though this area was thought to be the primary site where macrophages reside [2].

Macrophage distribution in inflamed cochleae

Upon inflammation, macrophages and monocytes are known to migrate robustly into the cochlea. We induced cochlear inflammation using LPS, which was inoculated in the middle ear through the tympanic membrane. At 1, 4, 7, and 10 days after LPS injection, GFP-positive cells (macrophage and monocytes) were present in higher numbers in the cochlea than in the PBS injection group (Fig. 3). Interestingly, the distribution patterns in the spiral ligament were similar to those in the untreated cochlea, even though macrophage density had significantly increased (Fig. 2). In addition, migration of green cells was identified in the basilar membrane and spiral limbus, a finding which is consistent with previous studies [2,5].

DISCUSSION

It is generally thought that macrophages are distributed in all areas of the spiral ligament and are more abundant in inferior regions represented by type II and IV fibrocyte zones [2,5,15,16], whereas the organ of Corti lacks macrophages even in inflamed conditions, which seems to be a defense mechanism against collateral damage caused by inflammatory cells [3,17]. In this study, interestingly, we noted consistently low populations of macrophages near the stria vascularis including type II fibrocyte zones (Figs. 2 and 4) in both the untreated and inflamed conditions (Fig. 2).

Even though we meticulously reviewed previously published studies, we were unable to uncover any images that showed macrophages in the areas that lacked macrophages in this study [2,3,5,16,17]. These areas seem to be protected from the intrusion of macrophages, given the importance of the stria vascularis and its adjacent area in electro-homeostasis of the cochlea, especially potassium recycling. Furthermore, these areas are associated with type I, II, and V fibrocyte zones, and Na/K-ATPase (a critical ion channel for maintaining endocochlear potential) is highly expressed in type II and V fibrocyte zones [8,18,19].

The mechanism protecting these areas from macrophage intrusion is not clear. However, it seems to be associated with intercellular junctions, which are known to form a blood-labyrinth barrier. It does not seem that fibrocytes themselves play a key role in this protection because the areas lacking macrophages did not precisely match with the known zones of each type of fibrocyte. For instance, only the narrow medial area of the type I fibrocyte zone was lacking in macrophages. In 2017, Liu et al. [7] reported that claudin-11, which is a tight junction protein, acts as an ion barrier in the spiral ligament. They showed that claudin-11 was expressed in areas adjacent to the stria vascularis, with a similar pattern to the areas lacking macrophages in our study. However, the wide area beneath the stria vascularis did not match with this protein marker, and no protein marker corresponding to the areas lacking macrophages in this study is yet known to the best of our knowledge.

Three-dimensional images of intact cochleae were used in this study to investigate the exact locations of macrophages, as con-
CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was reported.

ACKNOWLEDGMENTS

This research was supported by a grant from the National Research Foundation of Korea (NRF-2019R1A2C1084033 to Jinsei Jung) and a faculty research grant of Yonsei University College of Medicine (6-2020-0153 to Seong Hoon Bae).

ORCID

Seong Hoon Bae https://orcid.org/0000-0001-9243-9392
Sang Hyun Kwak https://orcid.org/0000-0002-2827-8456
Jee Eun Yoo https://orcid.org/0000-0001-9372-999X
Kyu Min Kim https://orcid.org/0000-0003-2187-8034
Young-Min Hyun https://orcid.org/0000-0002-0567-2039
Jae Young Choi https://orcid.org/0000-0001-9493-3458
Jinsei Jung https://orcid.org/0000-0003-1906-6969

AUTHOR CONTRIBUTIONS

Conceptualization: SHB. Data curation: SHB, SHK, KMK, JEY. Formal analysis: SHB. Resources: YMH, JYC. Funding acquisition: JJ, SHB. Methodology: SHK, KMK. Supervision: YMH, JYC, JJS. Writing—original draft: SHB. Writing—review & editing: JJ.

SUPPLEMENTARY MATERIALS

Supplementary materials can be available at https://doi.org/10.21053/ceo.2020.00395.

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