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The spatial distribution pattern of Connexin26 expression in supporting cells and its role in outer hair cell survival

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Abstract
Mutations in the GJB2 gene (which encodes Connexin26 (Cx26)) account for about a quarter of all cases of nonsyndromic deafness. Previous studies have indicated that knockout (KO) of Gjb2 gene during early postnatal days can cause outer hair cell (OHC) loss in mouse models. However, the postnatal spatial distribution pattern of Cx26 in different types of supporting cells (SCs) and the role of such distributions for the survival of OHCs is still obscure. In this study, the spatial distribution patterns of Cx26 in SCs were observed, and based on these observations different spatial Cx26-null mouse models were established in order to determine the effect of changes in the spatial distribution of Cx26 in SCs on the survival of OHCs. At postnatal day (P)3, unlike the synchronous expression of Cx26 along both longitudinal and radial boundaries of most types of SCs, Cx26 expression was primarily observed along the longitudinal boundaries of rows of Deiter’s cells (DCs). From P5 to P7, radial expression of Cx26 was gradually observed between adjacent rows of DCs. When Gjb2 gene was knocked out at random in different types of SCs, about 40% of the total DCs lost Cx26 expression and these Cx26-null DCs were distributed randomly in all three rows of DCs. The mice in this randomly Cx26-null group showed normal hearing and no significant OHC loss. When using a longitudinal KO pattern to induce knockout of Gjb2 gene specifically in the third row of DCs, about 33% of the total DCs lost Cx26 expression in this specific longitudinally Cx26-null group. The mice in this group showed late-onset hearing loss and significant OHC loss, however, the morphology of corresponding DCs was slightly altered. In both experimental groups, no substantial DC loss was observed. These results indicate that longitudinal Cx26-based channels are predominant in DCs during P3–P5. The Cx26 expression along rows of DCs might play a key role in the survival of OHCs, but this longitudinal KO pattern in DCs has a limited effect on DC survival or on its postnatal development.

Introduction
Mutations in the GJB2 gene (which encodes Connexin26 (Cx26)) account for a quarter of all cases of nonsyndromic deafness in different populations1–3. Cx26 self-assembles to form intercellular channels called gap junctions, and these allow ions, second messengers, microRNAs, and other small molecules to pass through4–6. Several different mouse models have been established to study the mechanism through which GJB2 mutations lead to deafness, but this process is still poorly understood.

In the inner ear of mice and humans, the Cx26 protein is not expressed in auditory sensory hair cells (HCs), but it is expressed in different types of supporting cells (SCs) in the cochlear epithelium7–9. In the mouse cochlea, Cx26 is first detected in a few cells on embryonic day 14.5, and the region of cochlear Cx26 expression gradually expands during postnatal development10. Radial sections show that the Cx26 protein is found in the cells of the greater
epithelial ridge (GER) at postnatal day (P) 2, and pillar cells (PCs), Deiter’s cells (DCs), and Claudius cells (CCs) show punctate Cx26 signals at P3. By P7, Cx26 is expressed in all types of SCs in the cochlear epithelium. These studies have provided the time-sequence expression pattern of cochlear Cx26 in the radial plane. However, the cochlear epithelium is a finely detailed spatial structure, these spatial distribution patterns of cochlear Cx26 during different stages of postnatal development still need further study.

According to previous studies, loss of outer HCs (OHCs) and developmental abnormalities in SCs were the two main pathological changes in different Gjb2 mutation or knockout (KO) mouse models. In these models, cochlear Gjb2 gene could be extensively knocked out in different types of SCs, and rapid loss of SCs and OHCs was seen at P14–P15. The changes in the Gjb2 gene were induced at embryonic or early postnatal days in these models. When cochlear Cx26 was reduced in the mature cochlea (P16 or P18), the mice showed normal hearing with no significant OHC or SC loss over the course of at least 1 month. These results indicated that early postnatal Cx26 expression might play a vital role in the survival of both OHCs and SCs. However, the specific mechanism by which early postnatal Cx26 expression affects the survival of cochlear cells is still obscure.

In this study, flattened cochlear preparations combined with radial sections were used to investigate the spatial distribution of Cx26 in different SCs at different postnatal time points. Furthermore, two distinct spatial Cx26-null mouse models were established to study the effect of the spatial distribution of Cx26 in SCs on the survival of OHCs. Use of the Lgr5 promoter allowed us to establish a specific longitudinally Cx26-null mouse model, and in these mice Gjb2 gene was knocked out in all of the third row DCs (DC3s) and some of the inner PCs (IPCs). The second mouse line carried the Rosa26 promoter, and injection of a small dose of tamoxifen into these mice caused random cells in all three rows of DCs and other SCs to lose their Cx26 expression. Comparisons between these two models in terms of the relationship between the number of Cx26-null DCs in a row and the rate of OHC loss have provided a better understanding of the effect of the early postnatal Cx26 distribution patterns of SCs on the survival of OHCs.

**Results**

**The distinct distribution patterns of Cx26 in different types of SCs during early postnatal development**

In cochlear radial sections, the Cx26 signal (red) was detected in cells of the GER at P2 (Figs. 1a, e). At P4, weak Cx26 signals could be detected in both PCs and DCs (white arrows, Figs. 1b, f). At P7, the Cx26 signals were found in greater numbers of SCs (white arrows and arrowheads, Figs. 1c, g). At P9, Cx26 signals were detected in almost all SCs but not in HCs (Figs. 1d, h). The flattened preparations provided a more detailed view of the spatial distribution of Cx26. At P3, the Cx26 staining (red) was clearly shown in DCs and PCs. Unlike PCs, which had both longitudinal and radial distributions of Cx26, the Cx26 in DCs was mainly expressed longitudinally in rows of DCs. At P3, a very small amount of Cx26 expression was observed at the boundary between two adjacent rows of DCs (radial expression, white arrowheads, Figs. 1i, l).

Compared with the radial expression of Cx26 in DCs, the longitudinal expression of Cx26 in DCs was still predominant at P5 (white arrows, Figs. 1j, m). By P7, however, this longitudinally dominant Cx26 distribution pattern in DCs disappeared. At this time, the Cx26 signal was almost the same along both the longitudinal and radial boundaries of all DCs (Figs. 1k, n). To confirm this unique distribution, the distribution patterns of Cx26 in different SCs were compared at P3. The cell membranes were labeled by phalloidin (green). In inner sulcus cells (ISCs) and CCs, similar Cx26 signals were evenly distributed along all of the cells’ boundaries (Figs. 1o, q). However, a strong Cx26 signal (white arrows, Fig. 1p) was clearly observed longitudinally in the same row of DCs. Occasionally, weak Cx26 signals in DCs (white arrowheads, Fig. 1p) were detected at the boundaries between adjacent rows of DCs.

**Two distinct spatial Cx26-null mouse models were established**

To confirm the spatial KO patterns of transgenic mice upon activation of the Lgr5 promoter, the expression of Cre was visualized by tdTomato. Consistent with a previous report, the tdTomato signals (red) were mainly observed in DC3s, IPCs, and inner phalangeal cells at P7 (Figs. 2a, b). Next, the expression levels of cochlear Cx26 were evaluated at P7. The littermates without Cre were used as controls, therefore two control groups (one for the randomly Cx26-null group (control R) and one for the specific longitudinally Cx26-null group (control L)) were included in the statistics. Western blot quantification showed that cochlear Cx26 expression in the randomly Cx26-null group decreased to 57.2 ± 8.8% (n = 4, P < 0.05). However, there was no statistically significant difference between the control L and specific longitudinally Cx26-null groups (n = 4, P = 0.32). To better understand the residual Cx26 expression in these models, Cx26-null cells were quantified at P7. In the randomly Cx26-null group, the Cx26-null cells were observed in nearly all types of SCs. Cx26-null DCs (asterisks in Figs. 2g–i) were observed in different rows (DC1, DC2, and DC3), and a number of IPCs (white arrows, Figs. 2g–i) and OPCS (arrowheads, Figs. 2g–i) lost Cx26 expression in this.
group. However, in the specific longitudinally Cx26-null group, nearly all DC3s (asterisks, Figs. 2j–l) lost their Cx26 signal, whereas the DC1s and DC2s still had normal Cx26 expression. In this group, all of the OPCs still had Cx26 expression, whereas some IPCs lost Cx26 expression (white arrows, Figs. 2j–l). In the randomly Cx26-null
group \((n = 3)\), quantitative analysis showed that the proportions of Cx26-null SCs (DCs and PCs) were \(45.5 \pm 1.4\%\), \(44.7 \pm 2.8\%\), and \(37.9 \pm 1.4\%\) in the apical, middle, and basal turns of the cochlea, respectively. However, the proportions of Cx26-null SCs (DCs and PCs) in the specific longitu-dinally Cx26-null group \((n = 3)\) were \(36.3 \pm 2.1\%\), \(31.9 \pm 2.5\%\), and \(28.6 \pm 3.1\%\) in the apical, middle, and basal turns, respectively. Additionally, the proportions of total Cx26-null DCs in the randomly Cx26-null group \((n = 3)\) were \(42.5 \pm 3.2\%\), \(40.8 \pm 1.6\%\), and \(36.0 \pm 2.6\%\) in the apical, middle, and basal turns, respectively. However, the proportions of total Cx26-null DCs in the specific longitu-dinally Cx26-null group \((n = 3)\) were \(33.0 \pm 1.8\%\), \(30.5 \pm 0.5\%\), and \(32.0 \pm 0.7\%\) in the different turns, respectively (Fig. 2n). Furthermore, Cx26-null DCs in the different rows were also quantified. In the specific longitu-dinally Cx26-null group, only DC3s lost their Cx26 expression (>95%). However, about 40% of all DC1s, DC2s, and DC3s lost their Cx26 expression in the random-ly Cx26-null group (Fig. 2o).

Late-onset high-frequency hearing loss was seen in the specific longitu-dinally Cx26-null group

The auditory brainstem response (ABR) was tested at P18 and P60. The littermates without Cre were tested as controls; therefore, two control groups (control R and control L) were included in the statistics. At P18, all four groups of mice \((n = 5\) in each group) showed normal hearing. At P60, the hearing thresholds of the specific longitu-dinally Cx26-null group at 8, 16, 32, and 40 kHz were \(42.0 \pm 3.7\), \(28.0 \pm 2.5\), \(39.0 \pm 1.9\), and \(59.0 \pm 1.0\) dB SPL, respectively, and the hearing thresholds of the corre-sponding control group at 8–40 kHz were \(35.0 \pm 3.2\), \(26.0 \pm 1.9\), \(35.0 \pm 1.6\), and \(48.0 \pm 3.7\) dB SPL, respectively. Compared with controls, there was a statistically sig-nificant difference at 40 kHz \((n = 5, P = 0.02)\) (Fig. 3).

OHC loss without corresponding DC loss was observed in the basal turn of the cochlea from the specific longitu-dinally Cx26-null group

Quantification of OHCs and DCs was performed at P18 and P60. The mice from the two control groups showed no differences in the ABR test, so the two control groups were combined into one group for quantification analysis \((n = 4)\). In the randomly Cx26-null group, there was no obvious OHC or DC loss at P18 (Fig. 4c). However, in the specific longitu-dinally Cx26-null group, scattered OHC3 (the third row of OHC) loss was found in the basal turn (asterisks, Fig. 4b), whereas their corresponding DCs were still intact at P18 (white arrows, Fig. 4b). Additionally, a cochleogram showed that 17.7 ± 3.3%–62.9 ± 10.7% of OHC3 loss was seen in the cochlear middle to basal turn of the specific longitu-dinally Cx26-null group at P18 (Fig. 4d). No substantial DC loss was observed in either of the two experimental groups at P18 (Fig. 4e).

At P60, OHC3 loss was increased (white arrowheads, Fig. 4g) in the basal turn of the cochlea of the specific longitu-dinally Cx26-null group. Accordingly, no sub-stantial DC loss was observed in the corresponding region (white arrows, Fig. 4g). Occasionally, a few missing OHCs were observed in the basal turn of the randomly Cx26-null group, whereas no obvious DC loss was found in the corresponding region (Fig. 4h). The cochleogram showed that 53.0 ± 11.4% to 96.1 ± 2.8% OHC3 loss was found from the middle to the basal turn of the cochlea of the specific longitu-dinally Cx26-null group at P60 (Fig. 4i). No substantial DC loss was observed in either of the two experimental groups at P60 (Fig. 4). In two experimental groups, quantification analysis showed that no substantial OHC1 or OHC2 loss was found at P18 or P60 (Fig. S2 A–D).

Two other specific longitu-dinally Cx26-null mouse models were established by varying the dose of tamoxifen injection. In these two models, one-third or one-sixth of the standard tamoxifen dose was given in the same manner as in the specific longitu-dinally Cx26-null mouse lines and are referred to as the longitu-dinally Cx26-null group \((1/3)\) and longitu-dinally Cx26-null group \((1/6)\), respectively. In the longitu-dinally Cx26-null group \((1/3)\) \((n = 3)\), quantitative analysis showed that the proportions of SCs (DCs and PCs) without Cx26 were 16.1 ± 1.3%, 13.8 ± 0.9%, and 12.5 ± 0.6% in the apical, middle, and basal turns, respectively (Fig. 5g), and there were very few Cx26-null IPCs. In the longitu-dinally Cx26-null group \((1/6)\) \((n = 3)\), these proportions were reduced to 11.4 ± 1.0%, 9.7 ± 0.8%, and 9.2 ± 0.8% in the three turns of the
cochlea, respectively (Fig. 5g). The proportions of Cx26-null DC3s were 74.5 ± 3.9%, 71.2 ± 1.4%, and 70.3 ± 3.3% in the different turns from the longitudinally Cx26-null group (1/3) (Fig. 5h), whereas they were 51.1 ± 2.7%, 49.4 ± 1.2%, and 49.9 ± 3.0% in the longitudinally Cx26-null group (1/6) (Fig. 5h). The cochleogram (n = 3 or 4)
showed that the rates of OHC3 loss between the specific longitudinally Cx26-null group and the longitudinally Cx26-null group (1/3) were not significantly different. However, the OHC3 loss was significantly reduced in the longitudinally Cx26-null group (1/6) (Fig. 5o).

**Normal structure of the organ of Corti and no spiral ganglion neuron loss were observed in the specific longitudinally Cx26-null group**

Radial sections showed that the tunnel of Corti was open and Nuel’s space was well formed at P18 in the specific longitudinally Cx26-null group (black arrows and arrowheads, Figs. 6e–g). Consistent with the HC counts, the OHC3s were lost in the basal turn (black arrow, Fig. 6h), whereas the DCs remained intact (Fig. 6h). The pathological change of the longitudinally Cx26-null group at P60 was similar to that at P18 (Figs. 6m–o), and the missing OHC3 can be also observed in the basal turn (black arrow, Fig. 6p). Additionally, there was no substantial spiral ganglion neuron loss in this group at P60 (Fig. 6w).

**The morphology of the phalangeal processes and the ultrastructure of DC3s in the specific longitudinally Cx26-null group**

In the control group, the phalangeal processes of DCs were thin cylindrical prominences (black arrows, Fig. 7a), and the bodies of the DCs (black arrowheads, Fig. 7a) were relatively uniform. In the specific longitudinally Cx26-null group, the phalangeal processes (black arrows, Fig. 7b) of the DC3s can develop into finger-like structures, but the diameters of the phalangeal processes changed a little bit, and most of them became larger. The bodies of the DC3s were not uniform, and some of the DC3 bodies were swollen (black arrowheads, Fig. 7b). The ultrastructural observation showed that DC1s in the specific longitudinally Cx26-null group were not affected, and bundles of microtubules and normal mitochondria (Figs. 7g, h) were observed in the bodies of DC1s. In DC3s, some lysosomes and mitochondria could be found in the perinuclear zone. Except for a few lipid droplets (black arrowhead, Fig. 7j), no signs of karyopyknosis, karyorrhexis, or autophagy were observed (Fig. 7i) in the DC3s from the specific longitudinally Cx26-null group.

**Discussion**

Cx26 protein expression in DCs has both time-dependent and spatial distribution patterns. First, Cx26 is expressed longitudinally to form a dotted line pattern along each row of DCs at P3–P5. However, very little Cx26 is expressed at the boundaries between two adjacent rows of DCs during this period. At P5–P7, Cx26 begins to be expressed at the boundaries between two adjacent rows of DCs. These results suggest that intercellular communication within rows of DCs is preferentially built up at P3, and these longitudinal Cx26-based channels in rows of DCs predominate from P3 to P5. Previous studies showed the time course of Cx26 expression in radial sections, and Cx26 expression was initially observed in the cells of the GER and spread toward the cells in the lesser epithelial ridge (LER) from P1 to P810,11. Thus, it would seem that the radial Cx26-based channels were established from the GER to the LER in turn, and these junctions are likely to be necessary for the radial delivery of small molecules between the GER and the LER. However, according to our observations, this radial intercellular communication between SCs might have limited effects in DCs during P3 to P5. Due to the lack of radial Cx26 expression in DCs, small molecules from the GER might be slowed down or even blocked in different rows of DCs during this period. In contrast, longitudinal delivery of small molecules through Cx26-based channels predominates within rows of DC at this time.

The postnatal expression of Cx26 in the organ of Corti is important for the survival of OHCs, and early postnatal Cx26 expression in rows of DCs might play a key role for the survival of OHCs. In the randomly Cx26-null group,
Fig. 4 OHC and DC counts in the control and experimental groups. a–c Representative images of HCs (Myosin7a, red) and corresponding DCs (Sox2, white) of different turns in control (a), specific longitudinally Cx26-null (b), or randomly Cx26-null groups (c) at P18. Asterisks indicate the missing OHC3s (b), and white arrows indicate the corresponding DC3s (b). d Quantifications of OHC3 loss at specific cochlear locations in the different groups at P18. e Quantifications of DC’s survival at specific cochlear locations in the different groups at P18. f–h Representative images of HCs (Myosin7a, red) and corresponding DCs (Sox2, white) of different turns in control (f), specific longitudinally Cx26-null (g), or randomly Cx26-null groups (h) at P60. White arrowheads indicate the regions of missing OHC3s (g), and white arrows indicate the regions of corresponding DC3s in specific longitudinally Cx26-null group at P60 (g). i Quantifications of OHC3 loss at specific cochlear locations in the different groups at P60. j Quantifications of DC3’s survival at specific cochlear locations in the different groups at P60. The scales in panel a represent 40 µm.
Fig. 5 Cx26 expression patterns and OHC loss patterns in different longitudinally Cx26-null groups. a–c Cx26 immunolabeling (red) in the apical, middle, and basal turns of the longitudinally Cx26-null group (1/3), respectively. d–f Cx26 immunolabeling in the apical, middle, and basal turns of the longitudinally Cx26-null group (1/6), respectively. Asterisks indicate Cx26-null DCs. g Cx26-null cell counts at P7 in different turns of the organ of Corti (n = 3 in each group). h Cx26-null DC3 counts at P7 (n = 3 in each group). * and # indicate significant differences between two groups (P < 0.05). i–k Representative images of HCs in the apical (i), middle (j), and basal turns (k) from the longitudinally Cx26-null group (1/3). l–n Representative images of HCs in the different turns from the longitudinally Cx26-null group (1/6). o Quantifications of OHC3 loss at specific cochlear locations in different groups at P18. Asterisk indicates a significant difference compared with the specific longitudinally Cx26-null group. The hash symbol indicates a significant difference compared with the longitudinally Cx26-null group (1/3). p–s Representative images of the middle-basal parts of flattened cochlear preparations in different groups. White arrowheads indicated the regions of OHC3 loss. The scales in panels a, i, and p represent 30, 40, and 200 µm, respectively.
numerous OPCs, IPCs, CCs, and ISCs had no Cx26 expression (Fig. S1). In the whole cochlea, Cx26 expression was reduced by about 42.8% in the randomly Cx26-null group, whereas there was no reduction in overall cochlear expression of Cx26 in the specific longitudinally Cx26-null group. However, OHC counts showed that no substantial OHC loss occurred in the randomly Cx26-null group. Thus, it would appear that KO of Cx26 in a certain
number of SCs (except DCs) might not cause significant OHC loss. To validate this speculation, ISC-, IPC-, and CC-specific KO mouse models would be needed for further investigations.

The total proportion of Cx26-null DCs in the randomly Cx26-null group was about the same as that in the specific longitudinally Cx26-null group; however, OHC loss was only seen in the latter group. This observation indicated that there was no strong correlation between the total proportion of Cx26-negative DCs and OHC loss. However, the proportion of Cx26-null DC3s in the randomly Cx26-null group was about 40% (ranging from 36.0 ± 2.6% to 42.5 ± 3.2% in the different turns). By using different doses of tamoxifen injection (including 1/6, 1/3, or the full standard dose), these proportions could be manipulated to range from 49.4% to 100% in different longitudinally Cx26-null groups. In accordance with the rates of OHC loss in these groups, a strong correlation between the proportion of Cx26-negative DCs in a row and the corresponding OHC loss.
was seen. Based on our data, there might not be significant OHC loss when <40% of the DCs in a row lose Cx26 expression. OHC loss was observed in the basal turn when the proportion of Cx26-null DCs in a row exceeded 50%, and OHC loss reached its maximum when this proportion exceeded 70%.

Based on the above results, we propose the following hypothesis. From P3 to P5, the cells in rows of DCs are connected together by longitudinal Cx26-based channels. In other words, a row of DCs is connected by Cx26-related channels like a functional syncytium, which has a certain similarity with cardiomyocytes coupled by gap junctions. If >70% of the DCs lose Cx26 expression, this will disrupt the longitudinal intercellular communication in a row of DCs and severely impair the function of that row of DCs. This will ultimately lead to a significant loss of corresponding OHCs. When the proportion of Cx26-null DCs in a row is <40%, these residual Cx26-based channels might be sufficient to maintain the normal function of DCs and prevent the loss of OHCs. Another considerable factor is that we still lack the specific promoter for DC1 (or DC2), the accurate proportion of Cx26-null DC1 (or DC2) to maintain the survival of corresponding OHC1 (or OHC2) still needed further investigation.

Compared with the effect seen on OHCs, the two spatial Cx26-null patterns had no obvious negative effects on the survival of DCs or SGNs. In a previous study using a systemic Cx26-null mouse model, we found that the phalangeal processes of the three rows of DCs did not develop into finger-like structures. Although the morphology of DC3s in the present work changed a little bit, the phalangeal processes of DC3s could still develop into finger-like structures in the specific longitudinally Cx26-null group. Moreover, the ultrastructure of DC3s showed no obvious pathological changes in this group. These results suggest that the longitudinal KO pattern in DCs has limited effect on DC’s survival or DC’s postnatal development.

The late-onset hearing loss at high frequency might be caused by OHC3 death in the basal turn in the specific longitudinally Cx26-null group. According to Muller’s formula, the frequency of 40 kHz corresponds to 75% distance from the apex. At P18, the rate of OHC3 loss at the location near the 75% distance from the apex only ranged from 14.2 ± 3.3% to 33.2 ± 7.8%. If calculated by the total number of OHCs, the rate of OHC loss would not exceed 12%. This slight OHC loss might not cause significant hearing loss that can be detected by the ABR test. However, the rate of OHC3 loss near this location (for 40 kHz) ranged from 53.0 ± 11.2% to 77.2 ± 9.1% at P60, whereas the rate of OHC3 loss near the location responsible for hearing at 32 kHz was 25.0 ± 7.9% at this time point. Thus, it is reasonable that a statistically significant increase in hearing threshold was only found at 40 kHz.

In conclusion, Cx26 expression in DCs has time-dependent and spatial distribution patterns during postnatal development. Cx26 is predominantly expressed longitudinally in rows of DCs during P3–P5 (Figs. 8a, b). In randomly Cx26-null group, about 40% of the total DCs lost Cx26 expression and these Cx26-null DCs were distributed randomly in all three rows of DCs (Fig. 8c). It would not cause significant OHC loss in this group (Fig. 8f). However, when using a longitudinal KO pattern to induce KO of Gjb2 gene specifically in the third row of DCs (Fig. 8e), it can induce a substantial OHC3 loss in the basal turn (Fig. 8d). Our results indicated that the postnatal expression of Cx26 in the organ of Corti is important for the survival of OHCs, and this longitudinal expression of Cx26 in DCs during early postnatal development might play a key role in the survival of OHCs.

**Materials and methods**

**Mouse models**

Cx26loxP/loxP mice and Rosa26CreER mice were provided by Prof. Xi Lin at Emory University. Lgr5EGFP-Ires-CreERT2 (Lgr5CreER) and Rosa26CAG-loxP-stop-loxP-tdTomato (tdTomato) mice were provided from Prof. Renjie Chai at Southeast University. Tamoxifen-inducible Cx26loxP/loxP;Rosa26CreER and Cx26loxP/loxP; Lgr5CreER mice were generated by crossing the Cx26loxP/loxP mice with Rosa26CreER or Lgr5CreER mice, respectively. Mouse genotyping was performed by PCR amplification of tail genomic DNA. Details of the mice are given in our previous papers15,19. The genotyping primers were as follows:

- **Cx26 (F):** 5′-ACAGAATGTGTTGTGAATGG-3′
- **Cx26 (R):** 5′-CTTTCAATGTCTGGTGGAGTG-3′
- **Rosa26Cre (F):** 5′-AGCTAAACATGCTTCATCGTG GTC-3′
- **Rosa26Cre (R):** 5′-TATCCAGGGTACGGATAGTT CATG-3′
- **Lgr5CreER (F):** 5′-CTGTCTCTCTGCTCCGAGCTCT-3′
- **Lgr5CreER (R):** 5′-ATACCCCATCCCTTTTGAG C-3′
- **Lgr5-mutant (R):** 5′-GAACCTTCAGGTTACGTCG C-3′
- **tdTomato-wild-type (F):** 5′-AAGGGAGCTGCAGTG GAGT-3′
- **tdTomato-wild-type (R):** 5′-CTGTTTCGTCAGGGCAT GG-3′

To confirm the activation of the Cre recombinase, we crossed Lgr5CreER mice with tdTomato mice, and
tamoxifen (T5648-1G, Sigma-Aldrich, USA) was injected subcutaneously at P0 and P1 (the total dose was 1.5 mg/10 g body weight, once a day for 2 consecutive days). It was previously reported that CreER activation is tamoxifen dose-dependent; consequently, a smaller dose of tamoxifen (total dose 0.75 mg/10 g for 2 days) was administered to Cx26loxP/loxP; Rosa26CreER mice at P0 and P1, and cochlear Cx26 was partly and randomly knocked out in different types of cells of this group (the randomly Cx26-null group). To obtain the specific longitudinally Cx26-null mouse model, Cx26loxP/loxP; Lgr5CreER mice were given a tamoxifen dose of 1.5 mg/10 g for 2 days. Two reduced doses (total doses of 0.5 mg/10 g or 0.25 mg/10 g for 2 days) were injected into Cx26loxP/loxP; Lgr5CreER mice to establish the specific longitudinally Cx26-null group (1/3) and the specific longitudinally Cx26-null group (1/6), respectively. The littermates without Cre were used as controls.

All mice were raised in the specific-pathogen free Experimental Animal Center of Huazhong University of Science and Technology. All experimental procedures were conducted in accordance with the policies of the Committee on Animal Research of Tongji Medical College, Huazhong University of Science and Technology.

ABR
ABR was performed at P18 and P60. As we previously reported, mice (n = 6–8 in each group) were anesthetized by intraperitoneal injection with a mixture of ketamine (120 mg/kg i.p.) and chlorpromazine (20 mg/kg, i.p.). Body temperature was maintained by placing the anesthetized mice on a heating pad. The recording electrode was placed at the vertex of the skull, and the reference electrode was inserted at the tested ear with a ground electrode at the contralateral ear. Tone bursts of 8, 16, 32, and 40 kHz were generated and responses were recorded using a Tucker-Davis Technologies System (RZ6, Tucker-Davis Tech., USA). The responses were the average from 1024 stimuli and were recorded in decreasing 10 dB steps, then narrowing to 5 dB steps near the threshold. The ABR threshold for each frequency was determined by the lowest sound level that gave a measurable signal.

Protein extraction and western blots
Mice were deeply anesthetized and sacrificed at P7. Mouse cochleae (n = 4 in each group) were carefully dissected from the temporal bone in ice-cold 0.01 M phosphate-buffered saline (PBS), and the total proteins of the membranous labyrinth were extracted using RIPA lysis buffer (P0013B, Beyotime Biotechnology, China).

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Fig. 8 A cartoon showing the different spatial-specific Cx26-null patterns. a Cx26 expression (red) in the radial section of a mature organ of Corti. b Cx26 expression (red) in flattened preparations at P3. c, e Cx26 expression in the specific longitudinally Cx26-null (c) and randomly Cx26-null (e) groups at P7. d, f HC loss pattern in the specific longitudinally Cx26-null (d) and randomly Cx26-null (f) groups at P18.
The protein concentrations were quantified using a bicinchoninic acid protein assay kit (P00125, Beyotime Biotechnology, China). Equal amounts of protein samples were separated by electrophoresis on 12% sodium dodecyl sulfate–polyacrylamide gel electrophoresis gels and then transferred to polyvinylidenedifluoride (PVDF) membranes. The membranes were blocked in Tris-buffered saline (TBST with 0.1% Tween-20) containing 5% milk for 1 h. The Cx26 and β-actin proteins were detected using rabbit polyclonal antibodies against Cx26 (1:1,000 dilution, 512800, Invitrogen, USA) or rabbit polyclonal antibodies against β-actin (1:1000 dilution, 04-1116, Millipore, USA). After washing in TBST, the PVDF membranes were incubated with horseradish peroxidase (HRP)-conjugated goat anti-rabbit secondary antibody for 1 h at room temperature. Bands were visualized with an ECL reaction kit (P0018, Beyotime Biotechnology, China). The protein levels of Cx26 were measured by Quantity One 4.6.2 (Bio-Rad, USA) and normalized to the levels of β-actin (1:200 dilution, 04-1116, Millipore, USA). After washing in TBST, the PVDF membranes were incubated with horseradish peroxidase (HRP)-conjugated goat anti-rabbit secondary antibody for 1 h at room temperature. Bands were visualized with an ECL reaction kit (P0018, Beyotime Biotechnology, China). The protein levels of Cx26 were measured by Quantity One 4.6.2 (Bio-Rad, USA) and normalized to the levels of β-actin in the corresponding lane.

**Cochlear tissue preparation and immunofluorescent labeling**

For Cx26-null cell quantification, mice (n = 3) were sacrificed at P7. For cochlear HC and SC counting, mice (n = 4) were deeply anesthetized and sacrificed at P18 or P60. The cochleae were carefully dissected from the temporal bones and fixed in 4% paraformaldehyde in 0.01 M PBS for 1 h at room temperature. The flattened cochlear preparations were carefully dissected in ice-cold 0.01 M PBS. After blocking in a solution of 10% donkey serum with 1% Triton X-100 for 1 h, the samples were incubated with polyclonal rabbit anti-Cx26 antibodies (1:200 dilution, 512800, Invitrogen, USA), polyclonal rabbit anti-myosin7a antibodies (1:500 dilution, 25-6790, Proteus Bio-Sciences, USA), or polyclonal goat anti-sox2 antibodies (1:100 dilution, sc-17320, Santa Cruz Biotechnology, USA) diluted in 0.01 M PBS with 0.3% Triton X-100 overnight at 4°C. Samples were washed three times in 0.01 M PBS with 0.1% Tween-20 and then stained with Alexa Fluor 647 donkey anti-rabbit IgG or Alexa Fluor 594 donkey anti-goat IgG (1:200 dilution, ANTI032 and ANTI031, Antigene, China) for 1.5 h. 4',6-diamidino-2-phenylindole (DAPI, C1005, Beyotime Biotechnology, China) and phalloidin (0.05 mg/mL, P5282, Sigma, USA) were used for nucleus and F-actin staining, respectively. Images were taken with a laser scanning confocal microscope (Nikon, Japan). To better understand the spatial expression pattern of Cx26 in DCs, images from different levels of DCs were merged together.

**Resin sections and transmission electron microscopy**

Mice (n = 3) were deeply anesthetized and sacrificed at P60. The cochleae were removed and fully fixed with a mixture of 2% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M PBS. The samples were decalcified for 48–72 h in 10% disodium EDTA (pH = 7.2) and post-fixed for 1 h in 1% osmium tetroxide. After dehydration through a graded ethanol series, the samples were embedded in resin. The samples were sectioned (1.5 μm in thickness) and stained with toluidine blue (89640-5G, Sigma-Aldrich, USA) for light microscope observation. The ultrathin sections were stained with uranyl acetate and lead citrate for transmission electron microscopy examination (FEI Tecnai G2 20 TWIN, Thermo Fisher Scientific, USA).

**Scanning electron microscopy**

Mice were sacrificed at P18, and the cochleae were removed and fully fixed with a mixture of 2% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M PBS. After decalcification, the bony capsule and lateral walls were removed from the cochlea. The basal turn was exposed to show the morphology of the DC3s and OHC3s. Samples were subsequently dehydrated through a graded series of ethanol, critical point dried from liquid CO2 (HCP-2, Critical Point Dryer, HITACHI, Japan), mounted on stubs, and sputter-coated with a layer of gold (Eiko Engineering, Japan). Photographs were taken using a scanning electron microscope (VEGA 3 LMU, Tescan, Czechoslovakia).

**Statistical analysis**

All data are presented as means ± s.e.m. and plotted with Sigma Plot (version 12.5, Systat Software, Inc., USA). The t-tests were performed in SPSS software (version 19, IBM SPSS Statistics, USA), and P < 0.05 was considered to be statistically significant.

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**Conflict of interest**

The authors declare that they have no conflict of interest.
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