Claudin expression during early postnatal development of the murine cochlea

Takayuki Kudo, Philine Wangemann and Daniel C. Marcus*

Abstract

Background: Claudins are major components of tight junctions, which form the paracellular barrier between the cochlear luminal and abluminal fluid compartments that supports the large transepithelial voltage difference and the large concentration differences of $K^+$, $Na^+$ and $Ca^{2+}$ needed for normal cochlear function. Claudins are a family of more than 20 subtypes, but our knowledge about expression and localization of each subtype in the cochlea is limited.

Results: We examined by quantitative RT-PCR the expression of the mRNA of 24 claudin isoforms in mouse cochlea during postnatal development and localized the expression in separated fractions of the cochlea. Transcripts of 21 claudin isoforms were detected at all ages, while 3 isoforms ($Cldn-16$, −17 and −18) were not detected. Claudins that increased expression during development include $Cldn-9$, −13, −14, −15, and −19v2, while $Cldn-6$ decreased. Those that do not change expression level during postnatal development include $Cldn-1$, −2, −3, −4, −5, −7, −8, −10v1, −10v2, −11, −12, −19v1, −20, −22, and −23. Our investigation revealed unique localization of some claudins. In particular, $Cldn-13$ expression rapidly increases during early development and is mainly expressed in bone but only minimally in the lateral wall (including stria vascularis) and in the medial region (including the organ of Corti). No statistically significant changes in expression of $Cldn-11$, −13, or −14 were found in the cochlea of $Slc26a4^{−/−}$ mice compared to $Slc26a4^{+/−}$ mice.

Conclusions: We demonstrated developmental patterns of claudin isoform transcript expression in the murine cochlea. Most of the claudins were associated with stria vascularis and organ of Corti, tissue fractions rich in tight junctions. However, this study suggests a novel function of $Cldn-13$ in the cochlea, which may be linked to cochlear bone marrow maturation.

Keywords: Tight junctions, Inner ear, Pendrin, SLC26A4, Mouse

Background

Tight junctions are structures consisting of proteins that join epithelial and endothelial cells to form continuous sheets and tubules which separate two liquid compartments. They consist of claudins [1], occludins [2] and other proteins that form a band-like network known as tight junction strands. These junctions are known to perform several functions (barrier, pore and fence), and are composed of several types of proteins: transmembrane (e.g., claudins and occludin), cytoplasmic, signaling and adapter links to the cytoskeleton [1, 3]. Barrier function refers to the restriction of paracellular movement of fluid constituents between the two fluid compartments that are separated by the cell layer. Pore function refers to the selective permeability of the paracellular barrier to those solutes that can pass between the fluid compartments. The fence function refers to the restriction of lateral movement of membrane proteins and lipids within the face of the plasma membrane, which retains the separate physiological functions of the luminal and abluminal cell membranes that are necessary to carry out vectorial transport of solutes and water.

Claudins are a family of more than 20 subtypes [1]. The specific isoforms of claudin included in a tight junction are the primary determinant of paracellular permeability [3]. Common structures of the claudin family include four transmembrane domains and two extracellular loops (Fig. 1). It is thought that charged amino acids in the first extracellular loop define the
of stria vascularis, mutation of Cldn-14 knockout mice [6]. In contrast to this pathology, chlear potential and the resulting deafness in adult basal cells, as confirmed by the reduction of endocochlear potential across the basal cell layer is supported by the barrier function of the highly dense tight junctions between the cell membranes [8]. This voltage is generated within the multi-layered stria vascularis in the cochlear lateral wall and originates as a potential difference across the basal cell layer of the stria between the cochlear lateral wall and the fibrous spiral ligament [8]. This potential difference is known to be critical for normal hearing [6, 7]. A major driving force for the ionic currents underlying the cellular transduction of sound into corresponding electrical signals to hearing centers in the brain is the endocochlear potential, the transepithelial voltage across the inner ear epithelium [8]. This voltage is generated within the multi-layered stria vascularis in the cochlear lateral wall and originates as a potential difference across the basal cell layer of the stria between the intrastrial fluid space and the perilymph pervading the fibrous spiral ligament [8]. This potential difference across the basal cell layer is supported by the barrier function of the highly dense tight junctions between the basal cells, as confirmed by the reduction of endocochlear potential and the resulting deafness in adult Cldn-11 knockout mice [6]. In contrast to this pathology of stria vascularis, mutation of Cldn-14 led to degeneration of a different cochlear structure, the sensory organ of Corti, and was associated with the human hereditary deafness DFNB29 [9, 10]. It is to be expected that mutations of other claudin isoforms in the cochlea could lead to impaired hearing. In addition to these examples of claudin isoform localization and expression, three other groups of investigators have reported expression of claudins in the cochlea [11–13]. Kitajiri and colleagues [12] examined Cldn-1 to Cldn-18 using immunohistochemistry, but their study was limited by a lack of antibodies for some claudins. We localized in the current study transcript expression of most of the claudin isoforms in multiple tissue fractions of the cochlea, including the outer layer of cochlear bone.

One of the most common hereditary deafness genes is pendrin (SLC26A4), which has been surprisingly shown to exert its strongest effects on cochlear function by expression in the endolymphatic sac during early development [14]. Lack of pendrin expression was found to be accompanied by delays in cochlear bone development and in expression of other genes due to an apparent local hypothyroidism [15]. It therefore was of interest to determine whether the expression of Cldn-13, observed in the present study to be predominantly expressed in the outer bone fraction, would be altered by deletion of the Slc26a4 gene in the mouse model.

The aim in the present study was to determine a) expression of claudin transcripts during early development, b) localization of the claudin isoforms among the cochlear regions and c) the potential effects of Slc26a4 knockout on claudin isoform expression.

**Methods**

Slc26a4+/− and Slc26a4−/− mice were obtained from a colony at Kansas State university and the heterozygous mice served as controls. Animals were deeply anesthetized with sodium pentobarbital (100 mg/kg i.p.). Temporal bones were removed from both male and female mice and whole cochleae were collected from age-sex matched littermates of Slc26a4+/− and Slc26a4−/−. Expression of claudins was determined on RNA isolated from a) whole cochlea, b) cochlear lateral wall tissues, c) cochlear medial fraction tissues, and d) outer cochlear bone. Lateral wall tissues were further microdissected into spiral ligament and stria vascularis fractions, while the medial fraction was further microdissected into organ of Corti and modiolus fractions. All procedures involving animals were approved by the Institutional Animal Care and Use Committee of Kansas State University (protocol 2925).

Total RNA was isolated from these tissues using the RNeasy Micro Kit (Qiagen, Valencia, CA; Cat #7400) and care was taken that RNA was extracted from all cell types. Recombinant bovine DNase I, Grade 1 (Roche
Diagnosics Corp, Indianapolis, IN; catalog # 04536282001) was used to remove residual DNA. The quality and quantity of 18S rRNA were determined by using the RNA 6000 Nano Kit (Agilent Technologies, Santa Clara, CA; catalog # 5067–1511) with a Bioanalyzer (Agilent Technologies; Model 2100) and a spectrophotometer (Thermo Scientific, Wilmington, DE; NanoDrop 8000). The amount of RNA in each sample was calculated as the average of the results of the Bioanalyzer and NanoDrop assays.

Primers pairs for Cldn-1 to − 23 (excluding Cldn-21) were designed using software Primer3 (http://primer3.sourceforge.net/). The sequences of primers are documented in Table 1 and were validated with RNA from positive control tissues (tibia, liver, lung, intestine, kidney, stomach, skin, brain). mRNA expression was measured by quantitative RT-PCR using a Bio-Rad iCycler iQ thermocycler and QuantiTect SYBR Green RT-PCR Kit (Qiagen; catalog # 204243). Claudin mRNA was normalized against 18S. The calculation method has been described previously [16]. Melting curve measurements were made with the Bio-Rad thermocycler and the PCR product size was measured by using a DNA assay (Agilent DNA 1000 Kit; catalog #5067–1504) on the Agilent Bioanalyzer to exclude the detection of non-specific PCR products. This method yields quantitative measures of claudin isoform transcript expression that can be compared within each isoform; however, comparisons between and among isoforms are not quantitative in these experiments due to undetermined efficiencies in the RT step that vary for each primer pair [16].

Data are given as means ± standard deviation (SD) or ± standard error of the mean (SEM), as reported in Results. N values refer to the number of cochleae (Figs. 2 and 4) or to the number of isolated tissues (Fig. 3a and b) which are the same as the numbers of RT-PCR reactions analyzed. A one-way analysis of variance (ANOVA) with Holm-Sidak method post-test (Figs. 2 and 3) or a two-way ANOVA which tested for statistical significance of interaction between age and genotype. Since no statistically significant interaction was found for the 3 transcripts tested in the experiments shown in Fig. 4, individual paired differences were not assessed. P values of < 0.05 were considered as significant differences; analyses were performed with SigmaStat for Windows Version 4.0 software.

Results and discussion

We first determined the transcript expression level of 24 claudin isoforms in the whole cochlea of normal (Slc26a4−/−) mice at three different ages after birth: P2 and P6 before the onset of endocochlear potential generation and hearing in mice, and at P15, after acquisition of hearing. Transcripts of 21 claudin isoforms were detected at all ages, while 3 isoforms (Cldn-16, − 17 and − 18) were not detected (Fig. 2). The permeability properties of several isoforms have been unambiguously determined [17, 18] and are shown in (Fig. 2), as described in the figure legend. Cldn-10 [19] and Cldn-19 were each determined for two splice variants, v1 and v2 (Table 1).

Six cochlear claudin isoforms increase with development at P6 and/or P15: Cldn-9, − 13, − 14, − 15, and -19v2. By contrast, Cldn-6 expression decreases with development. Cochlear claudins that do not change significantly with development include: Cldn-1, − 2, − 3, − 4, − 5, − 7, − 8, − 10v1, − 10v2, − 11, − 12, − 19v1, − 20, − 22, and − 23. As described above, endocochlear potential normally develops between P6 and P15. So genes that change their expression in this period might be involved in establishment of the special properties of the paracellular barrier of the epithelial cells that border the endolymph, and thereby provide the resistive barrier that supports the large endocochlear potential. Cldn-19v2 appeared to increase expression only transiently during this period. The post-natal changes in expression of multiple claudin isoforms are consistent with the likely presence of factors that regulate claudin expression during development. Most striking of all, Cldn13 shows a remarkably large increase in cochlear expression compared to the others. Previously, Abuazza et al. [20] reported maturational decrease of Cldn-6, − 9 and − 13 transcripts and of paracellular protein in several segments of the mouse kidney. They suggested these changes may contribute to developmental changes in the paracellular permeability of kidney tubules. In our study of the cochlea, Cldn-6 undergoes developmental decrease in transcript expression from P2 to P6, and further from P6 to P15, as in the kidney (Fig. 2). By contrast to the kidney, Cldn-9 and -13 transcripts increased from P2 to P6, and further from P6 to P15 (Fig. 2).

The cochlear tissues expressing these claudins were resolved in two subsequent experimental series. In the first series (Fig. 3a), cochlea of adult (P18-P32, mean P22.6) Slc26a4−/− mice were subdivided into three fractions: 1) lateral wall (exclusive of outer bone), 2) medial region, and 3) outer bone. These fractions were assayed for 12 claudin isoforms: Cldn-5, − 6, − 7, − 10v1, − 10v2, − 13, − 15, − 19v1, − 19v2, − 20, − 22, and − 23. All of these claudins were detected both in the lateral wall and the medial region fraction. Cldn-19v1 and Cldn-19v2 were expressed most strongly in the medial region.

Interestingly, Cldn−13 was expressed virtually exclusively in the outer bone fraction, in spite of the statistically significant difference in the minimal expression in the two soft tissues. Wongdee et al. examined claudin expression in skull and tibia bone [21] and determined localization of Cldn-5, − 11, − 14, − 15 and − 16. The expression was limited to the cells lining the bone
| Template (v: splice variant) | Primers | Sequences | Product Length | GenBank Accession Number |
|----------------------------|---------|-----------|----------------|-------------------------|
| 18S | 18S_L | gaggttccgaagacgatcaga | 316 | X00686 |
| 18S_R | tcgtccaccaactaaagac | | |
| Claudin-1 | cldn1L | cgaccttcttgtgaactctga | 390 | NM_016674 |
| | cldn1R | cgtgggttggtgaggaagt | |
| Claudin-2 | cldn2L | ggtggtctcttgagggacat | 333 | NM_016675 |
| | cldn2R | ctttcctctgccctttgtg | |
| Claudin-3 | cldn3L | cgggagtgctttctctgtt | 344 | NM_009902 |
| | cldn3R | tgctgtcactggtggactcgta | |
| Claudin-4 | cldn4L3 | cgccgactttctaaacactta | 326 | NM_009903 |
| | cldn4R3 | gttccacagtaagcctgtg | |
| Claudin-5 | cldn5L2 | gacgccgttgtggtgag | 307 | NM_013805 |
| | cldn5R2 | gccccttcagttgacagt | |
| Claudin-6 | cldn6L1 | ctactagggctggagaggtg | 363 | NM_018777 |
| | cldn6R1 | tttgtgtgagcagggaactgt | |
| Claudin-7 | cldn7L1 | caactgtggccttttcaat | 329 | NM_016887 |
| | cldn7R1 | gctctccccctctggctgcttctc | |
| Claudin-8 | cldn8L4 | aagcggaaatcactctctctca | 399 | NM_016878 |
| | cldn8R4 | caggttggttcctcca | |
| Claudin-9 | cldn9L2 | tctactcctctctcgttc | 331 | NM_020293 |
| | cldn9R2 | ctaggtcagttccttgacag | |
| Claudin-10v1 | cldn10v1L2 | gggatatttctgtgctattt | 378 | NM_023878 |
| | cldn10v1R2 | tctcttctgtggctgata | |
| Claudin-10v2 | cldn10v2L | ttttctgtgctatcttgc | 375 | NM_021386 |
| | cldn10v2R | atctcctctagccctgt | |
| Claudin-11 | cldn11L2 | gcggaaaaatgcgaacggt | 315 | NM_008770 |
| | cldn11R2 | ggccgacatacagggaacaccag | |
| Claudin-12 | cldn12L3 | cagatgtgctctggtcagct | 304 | NM_022890 |
| | cldn12R3 | cccgtgtaaatcgtcagggt | |
| Claudin-13 | cldn13L2 | tgcggagaaacagggtgatac | 385 | NM_020504 |
| | cldn13R2 | gttgacacagacagagagtgc | |
| Claudin-14 | cldn14L3 | cttggtccatctctctcact | 332 | NM_019500 |
| | cldn14R3 | aagacgactctctctctgt | |
| Claudin-15 | cldn15L2 | aagacgacagcagaaatctg | 305 | NM_021719 |
| | cldn15R2 | caaagatgtggttggtggtg | |
| Claudin-16 | cldn16L1 | gcagggcaccaccattctcatt | 389 | NM_053241 |
| | cldn16R1 | taaccggcaccaggacagcag | |
| Claudin-17 | cldn17L11 | ggtgtaacaggtaggcccaag | 314 | NM_0181490 |
| | cldn17R11 | tgtgagcaacgacagaaagga | |
| Claudin-18 | cldn18L4 | gaacctttcccccaagaagag | 355 | NM_019815 |
| | | caagcgtgaaatagaggagg | |
| Claudin-19v1 | cldn19v1L | gaaggtctgtggtgatcttg | 321 | NM_001038590 |
| | cldn19v1R | aggatgtcgtggtggtgagtga | |
| Claudin-19v2 | cldn19v2L2 | tgctgtctcatctttcttgtg | 306 | NM_0153105 |
| | cldn19v2R2 | gacagttaaatgggtgtgct | |
(periostium), suggesting a function of claudin other than tight junction formation. They, however, did not test bone for the presence of Cldn-13. Johnson et al. reported Cldn-13 expression in G1E cells, a proerythroblastic cell line [22] and Cldn-13 was identified in a stress induced erythropoiesis pathway that is mainly expressed in tissues associated with haematopoietic function [23]. It is therefore likely that expression of Cldn-13 in cochlear outer bone might originate from the associated bone marrow, which develops during the

| Template (v: splice variant) | primers sequences product length | GenBank Accession Number |
|-----------------------------|----------------------------------|--------------------------|
| Claudin-20                  | cldn20L2 cagctcctgtctctcatct   | 356 NM_001101560         |
|                             | cldn20R2 aagcagactcctcagcaaa  |                          |
| Claudin-22                  | cldn22L2 gccttgagagacacaggag   | 342 NM_029383            |
|                             | cldn22R2 tttctggattggcttgcttc  |                          |
| Claudin-23                  | cldn23L2 tactacagcagagacacgca  | 320 NM_027998            |
|                             | cldn23R2 cagttagaggaagggcagca  |                          |

Fig. 2 Developmental expression levels of 21 cochlear transcripts for claudin isoforms at P2, P6 and P15 in Slc26a4+/− mice. Bars for each isoform are in chronological order, left to right; Top, middle and bottom panels are numerically-increasing isoforms. Cldn-16, −17 and −18 did not show detectable specific amplification. Claudins associated with established permeability properties are designated in the second row of the labels: B, barrier; P, permeable pore; +, cation-selective pore; −, anion-selective pore [17]. Asterisks indicate significant difference (P < 0.05) between bars embraced by brackets using one-way ANOVA and the Holm-Sidak post-test. The absence of brackets and asterisks indicates differences are not significant. Error bars, Standard Deviation. The individual descriptive statistics are derived from n cochleae, as indicated on the graph.
early postnatal period [15, 24]. In support of this proposition, it was found that Slc26a4−/− mice exhibit delayed bone marrow maturation between P6 and P15 [24]. Mouse Cldn-13 does not have a human homolog [3].

In the second series, cochleae of adult (P19-P28, mean P22.0) Slc26a4+/− mice were subdivided further into four micro-dissected fractions: 1) the stria vascularis and 2) spiral ligament fractions were separated from the lateral wall; 3) the organ of Corti and the 4) modiolus were separated from the medial structures. Claudins in the lateral wall and medial fractions that were highly expressed (claudin mRNA/18S > 4.5; dashed line) in this experimental series were analyzed in the more-finely separated tissues of the following series. The lateral and medial fractions were each subdivided into two smaller fractions in order to obtain finer resolution of location (n = 6). Lateral wall: stria vascularis and spiral ligament; Medial region: organ of Corti and modiolus. Ages of samples are between P19 and P28 (days), mean: 22.0. Asterisks indicate significant difference (P < 0.05) using the two-way analysis of variance as indicated by brackets; *, significant. Non-significant comparisons are not shown and comparisons other than stria vascularis – spiral ligament and organ of Corti – modiolus are given in the Additional file 1. Non-significant comparisons are not shown in Fig. 3b and comparisons other than stria vascularis – spiral ligament and organ of Corti – modiolus are given in the Additional file 1.

Three claudins were selected to investigate the possible effect of Slc26a4 gene deletion on inner ear developmental expression of claudins. Developmental expression of the three isoforms demonstrated a dramatic postnatal increase in Cldn-13 that was not characteristic of the other two claudins, consistent with the notion that Cldn-13 is not regulated by a mechanism common to the claudins highly expressed in the epithelial tissues. We examined RNA from whole cochleae from age- and sex-matched littermates of Slc26a4+/− and Slc26a4−/− and analyzed by two-way ANOVA 1) Cldn-11, which is expressed in basal cells of stria vascularis [12], and whose deletion in mice causes hearing loss, 2) Cldn-13, which is expressed in cochlear outer bone (this report), and 3) Cldn-14, which is expressed in organ of Corti and is responsible for human hereditary deafness DFNB29. The results of analysis (Fig. 4) showed no statistically significant interaction between age and genotype in all three genes and no further comparisons of individual paired genotypes were made.
“Cldn-21” was not included in this study. The nomenclature has varied and developed since 2001 and was not identified in mouse at the time of this study [25]. The mouse gene currently accepted as Cldn-21 [25, 26] has been heterologously expressed in MDCK epithelial cell cultures, immunolocalized to sites that also express the tight-junction protein occludin, and was shown to participate in a Na⁺-selective paracellular transport pathway [26].

Some of our data differ from previous observations: expression of Cldn-5, −6 and −15 was not detected by Kitajiri et al. [12], but were observed in our experiments. In kidney, Cldn-5 and -15 are expressed in endothelial cells, not epithelial cells [27]. By contrast, Kitajiri et al. [12] reported that there had been no expression of Cldn-5 and Cldn-15 in stria vascularis nor spiral ligament, both highly vascularized tissues. We found Cldn-6 expression, but it gradually decreased during early development. Consistent with our observation, Kitajiri et al. [12] did not see any expression of Cldn-6 in the adult cochlea.

Conclusions
We analyzed 24 claudins in structures of the inner ear. Previous studies did not show the presence and localization of Cldn-7, Cldn-13, Cldn-19 to −23 in the cochlea, but the results of our study showed regional localization of transcripts of these genes in the cochlea and developmental changes in two of them. We observed that Cldn-13 is expressed in bone and that its expression increased rapidly during early postnatal development. Most of the claudins were expressed in stria vascularis and organ of Corti, tissue fractions rich in tight junctions. However, this study suggests a novel function of Cldn-13 in the cochlea, which may be linked to cochlear bone marrow maturation.

Additional file

Additional file 1: “Claudin expression raw data”. This file contains the data points collected and analyzed in the text and figures. (XLSX 86 kb)

Abbreviations
Slc26a4+/−: Slc26a4 gene knockout mouse; Slc26a4+/−: Slc26a4 heterogeneous mouse

Acknowledgments
We thank Dr. Kalidou Ndaiye for his contributions to RT-PCR primer design and other assistance as Manager of the Molecular Biology Core of the College of Veterinary Medicine. We also thank Donald G. Harbridge and Joel D. Sanneman for their excellent technical support.

Funding
This work was supported by NIH grants R01-DC00212 (DCM), P20-RR017686 (DCM) and R01-DC01098 (PW) and by the College of Veterinary Medicine, Kansas State University. Publication of this article was funded in part by the Kansas State University Open Access Publishing Fund.
Availability of data and materials
The raw data from which Figs. 2, 3 and 4 are based and on which conclusions are made are contained in Additional file 1.

Authors’ contributions
TK contributed to the design and analysis of experiments, collected the experimental data and contributed to the writing of the manuscript. PJW and DM contributed to the design and analysis of experiments and writing of the manuscript. All authors read and approved the final manuscript.

Authors’ information
The current address for TK is Dept. of Otorhinolaryngology, South Miyagi Medical Center, Japan.

Ethics approval and consent to participate
Experiments were conducted according to an ethics protocol approved by the Kansas State University Institutional Animal Care and Use Committee (protocol #2925).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Received: 10 April 2017 Accepted: 15 January 2018
Published online: 25 January 2018

References
1. Muto S. Physiological roles of claudins in kidney tubule paracellular transport. Am J Physiol Renal Physiol. 2017;312:F9–F24.
2. Furuse M, Hirase T, Itoh M, Nagafuchi A, Yonemura S, Tsukita S, Tsukita S. Occludin: a novel integral membrane protein localizing at tight junctions. J Cell Biol. 1998;132:1777–88.
3. Günzé D, Yu AS. Claudins and the modulation of tight junction permeability. Physiol Rev. 2013;93:525–69.
4. Krause G, Winkler L, Mueller SL, Haseloff RF, Piontek J, Blasig IE. Structure and function of claudins. Biochim Biophys Acta. 2008;1778:631–45.
5. Furuse M, Sasai H, Tsukita S. Manner of interaction of heterogeneous claudin species within and between tight junction strands. J Cell Biol. 1999;147:891–903.
6. Gow A, Davies C, Southwood CM, Frolenkov G, Chrustowski M, Ng L, Yamauchi D, Marcus DC, Kachar B. Deafness in Claudin 11-null mice reveals the critical contribution of basal cell tight junctions to stria vascularis function. J Neurosci. 2004;24:7051–62.
7. Nakano Y, Kim SH, Kim HM, Sanneman JD, Zhang Y, Smith RJ, Marcus DC, Wangemann P, Nessler RA, Banfi B. A claudin-9-based ion permeability barrier is essential for hearing. PLoS Genet. 2009;5:e1000610.
8. Marcus DC. Acoustic transduction. In: Cell Physiology Source Book Essentials of Membrane Biophysics Edited by Sperelakis N San Diego: Academic Press; 2012:649–668.
9. Wilcox ER, Burton QL, Naz S, Riazuddin S, Smith TN, Ploplis B, Belyantseva I, Ben Yosef T, Liburd LA, Moore RJ, Kachar B, Wu DK, Griffith AJ, Riazuddin S, Friedman TB. Mutations in the gene encoding tight junction claudin-14 cause autosomal recessive deafness DFNB29. Am J Hum Genet. 2009;84:736–46.
10. Ben-Yosef T, Belyantseva IA, Saunders TL, Hughes ED, Kavamoto K, Van Itallie CM, Beyer LA, Halsey K, Gardner DJ, Wilcox ER, Rasmussen J, Anderson JM, Dolan DF, Forge A, Raphael Y, Camara SA, Friedman TB. Claudin 14 knockout mice, a model for autosomal recessive deafness DFNB29, are deaf due to cochlear hair cell degeneration. Hum Mol Genet. 2003;12:2049–61.
11. Florian P, Arnaheh S, Lessidensky M, Trott T, Bloedow A, Ernst A, Fromm M, Gitter A. Claudins in the tight junctions of stria vascularis marginal cells. Biochim Biophys Commun. 2003;304:5–10.
12. Kitajji SJ, Furuse M, Morita K, Saishin-Kiuchi Y, Kido H, Ito J, Tsukita S. Expression patterns of claudins, tight junction adhesion molecules, in the inner ear. Hear Res. 2004;187:25–34.
13. Elkouby-Naor L, Abassi Z, Lagziel A, Gow A, Ben-Yosef T, Double gene deletion reveals lack of cooperation between claudin 11 and claudin 14 tight junction proteins. Cell Tissue Res. 2008;333:427–38.
14. Li X, Sanneman JD, Harbridge DG, Zhou F, Ito T, Nelson P, Picard N, Chambery R, Eladari D, Miesner T, Griffith AJ, Marcus DC, Wangemann P. SLC26A4 targeted to the endolymphatic sac rescues hearing and balance in Slc26a4 mutant mice. PLoS Genet. 2013;9:e1003641.
15. Wangemann P, Kim HM, Billings S, Nakaya K, Li X, Singh R, Sharlin DS, Forrest D, Marcus DC, Fong P. Developmental delays consistent with cochlear hydropsyndrome contribute to failure to develop hearing in mice lacking Slc26a4/pendrin expression. Am J Physiol Renal Physiol. 2009;297:F1435–47.
16. Wangemann P, Iza EM, Albrecht B, Wu T, Jabbva SV, Maganti RJ, Lee JH, Everett LA, Wall SM, Rouday E, Green ED, Marcus DC. Loss of KCNJ10 protein expression abolishes endocochlear potential and causes deafness in Pendred syndrome mouse model. BMC Med. 2004;2:30.
17. Günzél D, Claudins: vital partners in transcellular and paracellular transport coupling. Pflugers Arch. 2017;469:35–44.
18. Günzél D, Fromm M, Claudins and other tight junction proteins. Compr Physiol. 2012;2:1819–52.
19. Van Itallie CM, Rogan S, Yu A, Vital LS, Holmes J, Anderson JM. Two splice variants of claudin-10 in the kidney create paracellular pores with different ion selectivities. Am J Physiol Renal Physiol. 2006;291:F1288–99.
20. Abugaza G, Becker A, Williams SS, Chakravarty S, Truong HT, Lin F, Baam M, Claudins 6, 9, and 13 are developmentally expressed renal tight junction proteins. Am J Physiol Renal Physiol. 2006;291:F1312–F1411.
21. Wongdee K, Pandaranandaka J, Teerapornpuntak J, Tudpor K, Thongbuncho J, Thongon N, Jantarajit W, Krishnamra N, Charoenphandhu N. Osteoblasts express Claudins and tight junction-associated proteins. Histochem Cell Biol. 2008;130:79–90.
22. Johnson KD, Kim SI, Biessnick EH. Differential sensitivities of transcription factor target genes underlie cell type-specific expression profiles. Proc Natl Acad Sci U S A. 2006;103:15939–44.
23. Thompson PD, Tipney H, Brass A, Noyes H, Kemp S, Naessens J, Tassabehji M, Claudin 13, a member of the claudin family regulated in mouse stress induced hydropsyndrome. PLoS One. 2010;5.
24. Kudo T, Li X, Wangemann P. Bone marrow cell migration in early postnatal cochlea in a mouse model of Slc26a4-related syndromic and non-syndromic deafness (abstract). Assoc Res Otolaryngol. 2010.
25. Mineta K, Yamamoto Y, Yamazaki T, Tanaka H, Tada Y, Saito K, Tamura A, Igarashi M, Endo T, Takeuchi K, Tsukita S. Predicted expansion of the claudin multigene family. FEBS Lett. 2011;585:606–12.
26. Tanaka H, Yamamoto Y, Kashihara H, Yamazaki T, Tani K, Fujiyoshi Y, Mineta K, Takeuchi K, Tamura A, Tsukita S, Claudin-21 has a Paracellular Channel role at tight junctions. Mol Cell Biol. 2016;36:954–64.
27. Kichu-Saishin Y, Gotoh S, Furuse M, Takasuga A, Tano Y, Tsukita S. Differential expression patterns of Claudins, tight junction membrane proteins, in mouse neuron segments. J Am Soc Nephrol. 2002;13:875–86.