Defective claudin-10 causes a novel variation of HELIX syndrome through compromised tight junction strand assembly

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Abstract Formation of claudin-10 based tight junctions (TJs) is paramount to paracellular Na⁺ transport in multiple epithelia. Sequence variants in CLDN10 have been linked to HELIX syndrome, a salt-losing tubulopathy with altered handling of divalent cations accompanied by dysfunctional salivary, sweat, and lacrimal glands. Here, we investigate molecular basis and phenotypic consequences of a newly identified homozygous CLDN10 variant that translates into a single amino acid substitution within the fourth transmembrane helix of claudin-10. In addition to hypohidrosis (H), electrolyte (E) imbalance with impaired urine concentrating ability, and hypolacremia (L), phenotypic findings include altered salivary electrolyte composition and amelogenesis imperfecta but neither ichthyosis (I) nor xerostomia (X). Employing cellular
TJ reconstitution assays, we demonstrate perturbation of cis- and trans-interactions between mutant claudin-10 proteins. Ultrastructures of reconstituted TJ strands show disturbed continuity and reduced abundance in the mutant case. Throughout, both major isoforms, claudin-10a and claudin-10b, are differentially affected with claudin-10b showing more severe molecular alterations. However, expression of the mutant in renal epithelial cells with endogenous TJs results in wild-type-like ion selectivity and conductivity, indicating that aberrant claudin-10 is generally capable of forming functional paracellular channels. Thus, mutant proteins prove pathogenic by compromising claudin-10 TJ strand assembly. Additional ex vivo investigations indicate their insertion into TJs to occur in a tissue-specific manner.

Introduction

Recently, sequence variants in CLDN10 have been linked to a non-Bartter, non-Gitelman salt-losing tubulopathy accompanied by altered handling of divergent cations accompanied by dysfunctional salivary, sweat, and lacrimal glands. The acronym HELIX has been coined for the associated clinical syndrome encompassing hypohidrosis, electrolyte imbalance, lacrimal gland dysfunction, ichthyosis, and xerostomia. CLDN10 encodes the tight junction (TJ) protein claudin-10 which comes in two major isoforms, claudin-10a and claudin-10b. While the expression of claudin-10a is restricted to kidney and uterus, claudin-10b has been found in many epithelia. In the kidney, claudin-10 is expressed by tubular epithelial cells from the proximal convoluted tubule to the cortical thick ascending limb of Henle’s loop (TAL) with claudin-10a found in the proximal tubule and claudin-10b expressed along TAL segments.

Claudin-10 belongs to the family of claudins that are integral components of TJs and largely determine their sealing and permeability properties. These proteins share a common topology of four transmembrane helices from which two extracellular loops protrude into the paracellular cleft, forming an extracellular domain. Claudins assemble both in cis (within one membrane) and in trans (between adjacent cells) to form intramembranous strands and networks thereof that vary in complexity, continuity, and depth. The extracellular domains of claudins critically contribute to strand formation and mediate molecular barrier or selective channel formation, thereby directing the paracellular transport of ions, unchanged solutes, and water.

The differential expression of claudin-10 isoforms along the kidney tubule has major implications for the ion selectivity of the paracellular pathway. While claudin-10a is thought to mediate paracellular Cl⁻ reabsorption in the proximal tubule, claudin-10b complements transepithelial Na⁺ reabsorption in the TAL by allowing for its paracellular passage due to a lumen-positive transepithelial potential. In this way, in the TAL, half of the total amount of Na⁺ is reabsorbed via the paracellular route. In glandular physiology, claudin-10b appears to play an even more critical role in saliva, sweat, and tear secretion as it is thought to mediate all of the transepithelial Na⁺ transport.

Here, we investigate phenotypic features, molecular consequences, and renal pathophysiology associated with a novel homozygous CLDN10 missense variant. Besides an expansion of the HELIX phenotype we provide evidence of defective TJ strand assembly accounting for altered paracellular ion transport.

Methods

For comprehensive methods, see Supplementary Methods.

Study approval

Patient data and material were used following approval (#402/16-ek) of the institutional review board (#IRB00001750) at University of Leipzig Medical School. Written informed consent was obtained from the patient involved in this study, including the consent to publish patient photographs. From healthy controls for diuretics responsiveness testing (4 males and 1 female, aged 30.8 ± 4.6 years, mean ± SD), written informed consent to participate was obtained prior to the study.

Genetic analysis

Genomic DNA was extracted from whole blood, analyzed with next generation sequencing technology using Illumina sequencing-by-synthesis technology and bioinformatically interpreted as described before. Details are provided in Supplementary Methods. The reported homozygous CLDN10 variant was confirmed by Sanger sequencing. Copy number variation analysis did not suggest hemizygosity.

Clinical testing

Diuretics responsiveness tests were based upon a previously published protocol. Water deprivation and hypertonic saline infusion tests were essentially performed as previously described. Details and further clinical tests are described in Supplementary Methods. We remark that the patient was 18 or 20 years of age when clinical tests were performed.
Immunofluorescence and immunohistochemistry

Immunofluorescence and immunohistochemical labeling of TJ components was performed as detailed in Supplementary Methods.

Structural bioinformatics and molecular modeling

Homology models were created utilizing MODELLER \(^{19}\) and Schrödinger’s BioLuminate/Maestro (Biologics Suite, Release 2019-2, Schrödinger, Germany). For the claudin-10b protomer model, the claudin-15 crystal structure \(^{20}\) (PDB ID 4P79) was used as template. For the claudin-10a protomer model, the claudin-10b model was used as template. Images were created with PyMOL 2.3 (Schrödinger, Germany).

**CLDN10 expression constructs, cell culture, and transfection**

CLDN10 expression constructs were generated and HEK293 and MDCK-C7 cells were cultured and transfected as detailed in Supplementary Methods.

**Förster resonance energy transfer (FRET) assay**

FRET was assessed between CFP- and YFP-claudin fusion proteins in transiently transfected HEK293 cells essentially as previously reported. \(^{21}\) A description can be found in Supplementary Methods.

**Freeze-fracture electron microscopy**

Stably transfected HEK293 cells were grown in 6-well plates until confluency and processed for freeze-fracture electron microscopy as previously reported. \(^{22}\) For quantification of strand breaks, electron micrographs were analyzed as detailed in Supplementary Methods.

**Electrophysiology**

Dilution and biionic potential measurements in stably transfected MDCK-C7 cell layers were essentially carried out as previously described. \(^{7}\) Liquid junction potentials were corrected for as previously reported. \(^{23}\) Relative permeabilities were computed as detailed in Supplementary Methods. Pore diameter was estimated by plotting the square root of \(P_x/P_{Na} \) (\(X, NH_4^+\) and various organic cations) against cation diameters and extrapolating the linear part of the relationship. \(^{23}\)

**Results**

**Clinicals**

A 16-year-old male of consanguineous Afghan descent presented with nocturnal enuresis, polyuria, and polydipsia since early childhood. Upon clinical examination, arterial hypotension and bradycardia were noted. Beard growth and axillary hair were markedly reduced while ichthyosis was not evident on dermal inspection. Laboratory tests revealed marked hypochloremic alkalosis with hypokalemia as well as high serum renin and aldosterone levels (Table S1). Additionally, hypermagnesemia and hypocalciuria were evident. Serum creatinine was normal and there were no morphological abnormalities on renal ultrasound, especially no signs of nephrocalcinosis or nephrolithiasis.

**Genetic testing**

Genetic testing by means of a tubulopathy specific gene panel revealed a novel homozygous missense variant in CLDN10 (c.494G>C; NM_006984.4). Consequences at the protein level are p.(Gly163Ala) for claudin-10a and p.(Gly165Ala) for claudin-10b, henceforth referred to as G163A and G165A, respectively. G163/G165 is located within the fourth transmembrane helix of claudin-10a/b and is highly conserved across species and among claudin family members (Fig. S1). Absence of the variant in both mutation (HGMD, ClinVar) and population databases (gnomAD, ExAc) as well as recent reports of claudin-10 associated disease \(^{1–4}\) suggested pathogenicity. Unfortunately, additional family members were unavailable for segregation analysis.

**Assessment of renal tubular pathophysiology**

To further characterize underlying mechanisms of enuresis, polyuria, and polydipsia, we performed diuretics responsiveness testing. Blocking the Na\(^+\)-Cl\(^-\) cotransporter in the proximal part of the distal convoluted tubule by hydrochlorothiazide led to an exaggerated response of fractional Na\(^+\) and Cl\(^-\) excretions (\(\Delta FE_{Na}\) 8.40% and \(\Delta FE_{Cl}\) 13.65%, respectively, with \(\Delta\) signifying maximal increase over mean basal level), resembling that observed in Bartter syndrome type I. \(^{17}\) Loop diuretic responsiveness testing revealed no overt differences in increases of fractional Na\(^+\) and Cl\(^-\) excretions between patient and controls (Fig. 1A, S2). Assuming complete blocking of TAL paracellular Na\(^+\) transport, this is compatible with partial compensation by the transcellular pathway. Additionally, a profound effect on renal Mg\(^{2+}\) and K\(^+\) handling was evident. While an exaggerated response of fractional Mg\(^{2+}\) excretion in the patient is suggestive of enhanced TAL paracellular Mg\(^{2+}\) reabsorption, the loop diuretic effect on fractional K\(^+\) excretion is compatible with adaptive changes in the distal nephron promoting enhanced K\(^+\) secretion with an increase in NaCl delivery.

Urine concentrating ability was assessed during water deprivation and turned out to be markedly reduced with urine osmolalities remaining below 600 mosmol/kg (Fig. 1B). Administration of desmopressin, an analogue of arginine vasopressin (AVP), resulted in an increase of urine osmolality of less than 10%. Determination of hypertonic saline-stimulated serum copeptin (25.3 pmol/L) indicated no defect in central AVP release, \(^{18}\) as did basal copeptin levels (12.5 ± 3.5 pmol/L, mean ± SD, \(n = 3\)). \(^{24}\) In the absence of a compensatory increase in endogenous AVP release, \(^{24,25}\) both compromised urine concentrating capacity and limited renal response to desmopressin most likely reflect a reduction in the buildup of medullary...
interstitial osmolality. In order to assess the impact of short-term increases in serum K$^+$ on maximal urine concentration while bypassing any gut-mediated kaliuretic effects, water deprivation testing was repeated with continuous intravenous infusion of KCl. Raising serum K$^+$ by about 1 mmol/L did neither impact urine osmolality nor desmopressin responsiveness (Fig. 1B). Yet, activity of the distal nephron K$^+$ secretory process as determined by transtubular K$^+$ concentration gradients rose with serum K$^+$ levels (Fig. 1C).

Assessment of extrarenal manifestations

Given the vast bodily distribution of claudin-10b and its significance for proper functioning of several exocrine glands, we sought to define the extrarenal phenotype.

Stimulated saliva flow rates did not differ between patient and controls (Fig. 2A). Yet, measurements of electrolytes in salivary fluid revealed pronounced ion imbalances in the patient (Fig. 2B): Na$^+$ and Cl$^-$ levels were markedly lower than in controls, compatible with the critical role of paracellular Na$^+$ transport for NaCl secretion in glandular epithelia. On the contrary, K$^+$ levels were exceedingly high, while Ca$^{2+}$ contents were unaltered. Immunofluorescence labeling of claudin-10 in submucosal salivary glands from the labial mucosa revealed a complete absence of the protein in comparison with control tissue that showed clear staining (Fig. 2C, S3).

Notably, despite the absence of xerostomia, severe enamel wear was found clinically and radiographically (Fig. 2D, E), suggesting impaired amelogenesis.

Schirmer’s test was used to assess tear production and revealed bilateral hypolacrimia (Fig. 3A). The starch-iodine test was used to qualitatively assess sweat secretion. Results were compatible with generalized hypohidrosis with the axillae more severely affected than palmar and plantar aspects of hands and feet, respectively (Fig. 3B). Quantitative assessment of sweat secretion indicated reduced basal and stimulated axillary sweat secretion rates (Fig. 3C). Immunohistochemical staining of axillary skin biopsy specimens showed the presence of claudin-10 in secretory tubules of eccrine sweat glands from both patient and control samples (Fig. 3D, E). Similarly, immunofluorescence labeling of claudin-10 in eccrine sweat glands demonstrated its presence in canaliculi of secretory portions (Fig. 3F).

Figure 1  Assessment of renal phenotype. (A) Furosemide responsiveness testing in the patient and in 5 control subjects. Shown are maximal absolute initial changes during loop diuretic exposition over mean basal levels. The transtubular K$^+$ concentration gradient (TTKG) was defined as urinary K$^+$/serum K$^+$ whenever serum osmolality exceeded urinary osmolality and else was calculated from the standard formula. Overt differences between the patient and controls are observed only for fractional excretions of Mg$^{2+}$ and K$^+$ as well as for the TTKG. The loop diuretic effect on fractional Mg$^{2+}$ excretion signifies that paracellular Mg$^{2+}$ reabsorption in the thick ascending limb of Henle’s loop is enhanced in the patient and may be involved in the development of hypermagnesemia. Loop diuretic effects on fractional K$^+$ excretion can be explained by adaptive changes in the distal nephron promoting enhanced K$^+$ secretion upon an increase in NaCl delivery. Median and IQR. (B) Development of urine osmolality in the patient over time during water deprivation with and without administration of KCl. Results are similar for both tests and indicate a markedly reduced urine concentrating ability with a blunted response to desmopressin, irrespective of an increase in serum K$^+$ level. (C) In the patient, the increase in serum K$^+$ parallels that of TTKG during water deprivation with concomitant intravenous KCl substitution. i.v., intravenously.
Figure 2  Assessment of dental phenotype and saliva secretion. (A) Stimulated saliva flow rates in the patient and in 4 control subjects. The amount of saliva produced did not differ between the patient and controls. Mean. (B) Assessment of Na\(^+\), Cl\(^-\), K\(^-\), and Ca\(^{2+}\) concentrations in salivary fluid from the patient and from 4 control subjects. While Na\(^+\) and Cl\(^-\) concentrations were markedly lower in the patient, K\(^-\) levels were exceedingly high. Ca\(^{2+}\) concentrations turned out to be comparable between the patient and control subjects. Median. (C) Immunofluorescence labeling of claudin-10 and ZO-1 in submucosal salivary glands from the labial mucosa. When compared with control tissue, a complete absence of claudin-10 and a weaker ZO-1 signal are evident in the patient. Nuclei are stained with DAPI (blue). Scale bars, 5 μm. (D,E) Dental findings in the patient. Both clinical view and radiographs demonstrate severe enamel wear of the permanent teeth especially in the anterior region (arrowheads). Note the anteriorly open bite which is frequently associated with amelogenesis imperfecta\(^41\) and makes enamel wear unlikely to result from bruxism. On radiographs, enamel is still evident on molar teeth (arrows).
Figure 3  Assessment of tear production and sweat secretion. (A) Tear production as assessed by Schirmer’s test in the patient and in 4 control subjects. Results demonstrate bilateral hypolacrinia in the patient. Median and IQR. (B) Exemplary images from starch-iodine test in the patient. The starch-iodine reaction does only occur in the presence of water, indicated by dark discoloration. Axillae are more severely affected from hypohidrosis than palmar aspects of hands. (C) Quantitative assessment of sweat secretion before and after exercise in the patient and in 5 control subjects, indicating reduced basal and stimulated sweat secretion rates in both patient axillae. Median and IQR. (D,E) Immunohistochemical staining of claudin-10 in axillary skin biopsy specimens. In patient and control samples, claudin-10 (pink) is well-detectable and localizes to secretory tubules of eccrine sweat glands but is not found in excretory ducts (nuclei in dark blue). (D) Scale bars, 50 μm (left column) or 20 μm (right column). (E) demonstrates the loss of claudin-10 expression as a secretory tubule merges into an excretory duct (arrow) in a control
Homology modeling of claudin-10 3D structures and oligomerization

To assess the variant’s impact on tertiary protein structures, we employed homology modeling of human claudin-10 based upon claudin-15 crystal structure and previously reported claudin-10b models. In the resulting claudin-10a/-10b models, G163/G165 points to the center of the transmembrane four-helix bundle (Fig. 4A–C, E). This glycine residue is strongly conserved among claudins (Fig. S1), indicating that its small size and great torsional angle space are important for tight conformational packing of the four-helix bundle which is challenged by the insertion of a methyl group in the G163A/G165A variant (Fig. 4D, F). The models further indicate that the substitution G165A in claudin-10b is less well tolerated than G163A in claudin-10a because of the larger sidechain of T24 in claudin-10b which corresponds to A22 in claudin-10a. Additionally, data available for the region surrounding G163/G165 support the idea that G163A/G165A directly affects claudin-10 conformation (Fig. 4B).

As G165 does not line an interface between claudin protomers (Fig. 4G, H), we propose G165A (and, to a lesser extent, G163A in claudin-10a) to indirectly affect claudin assembly by perturbing the transmembrane four-helix bundle. Underlying mechanisms might involve alterations of the cis-interface close by, slight repositioning of W30 and the extracellular β-sheet involved in cis- and trans-interactions, or alterations of the membrane environment between claudin protomers (Fig. 4G, H).

Assessment of trans-interaction capability of wild-type and mutant claudin-10

To analyze claudin oligomerization, a TJ strand reconstitution assay was performed. YFP tagged claudin-10 proteins were transiently overexpressed in HEK293 cells (Fig. 5A). As these cells do not contain endogenous TJs, alterations in de novo formation of claudin-10 TJ strands could be probed in this way. Both YFP-claudin-10a wild-type (WT) and G163A showed similar enrichment at cell-cell contacts. In addition, they prominently localized to non-acidic intracellular granules in a similar fashion (Fig. 5A). Also, YFP-claudin-10b WT localized to the cell membrane with strong enrichment at cell-cell contacts, indicating trans-interactions and TJ strand formation. In contrast, YFP-claudin-10b G165A appeared to be more evenly distributed across the surface of neighboring cells, even though enrichment at cell-cell contacts was found to some extent. The lack of prominent enrichment indicates perturbed trans-interactions between claudin-10b G165A proteins. These data were supported by quantification of claudin-10 enrichment at individual cell-cell contacts, showing significantly lower enrichment factors for claudin-10b G165A when compared with the corresponding wild-type (Fig. 5B). We conclude that trans-interaction properties of both claudin-10 isoforms are differentially affected by the mutation.

Assessment of cis-interaction properties of wild-type and mutant claudin-10

For analysis of claudin cis-interaction, a FRET based cellular assay was employed. To probe the spatial cis-proximity between claudins, we assessed FRET efficiencies at cell-cell contacts between HEK293 cells coexpressing different CFP- and YFP-claudin fusion proteins that functioned as FRET-donor/-acceptor pairs (Fig. 5C). In the case of claudin-10a, FRET efficiencies were reduced only when assessing the interaction between claudin-10a G163A proteins. Coexpression of claudin-10a WT led to a complete rescue of proper cis-interactions. For claudin-10b, results indicate perturbed cis-interactions between claudin-10b G165A and both claudin-10b WT and G165A with a partial rescue by coexpression of the wild-type, indicating that the mutation’s impact on claudin-10 function is more severe for claudin-10b. Thus, G163A/G165A perturbs cis-interactions between claudin-10a/b proteins in reconstituted TJ strands.

Subcellular distribution of exogenous wild-type and mutant claudin-10 in MDCK-C7 and HEK293 cells

Differentiated renal epithelial cells (MDCK-C7) were stably transfected with wild-type and mutant claudin-10 constructs. Note that endogenously, MDCK-C7 cells express claudin-1,-3, -4, -5, -7, and -8 but no claudins that are known to form paracellular channels such as claudin-10a or claudin-10b. Immunofluorescence labeling of claudin-10 showed membrane localization and varying degrees of intracellular accumulation (Fig. 5S). While mutant proteins exhibited clear staining in a mottled perinuclear pattern, these effects were markedly less pronounced in claudin-10a/b WT expressing cells. For claudin-10b, these findings were corroborated by the fact that relative amounts of tight junctional claudin-10 were reduced in the mutant case (Fig. 5S). In contrast, when transiently overexpressed in HEK293 cells, both claudin-10a WT and G163A localized in the cell interior to non-acidic granules while claudin-10b WT and G165A showed no notable intracellular accumulation (Fig. 5A, 5S). These differential findings may reflect tissue-specificity of the cellular processing of aberrant claudin-10.

Ultrastructures of wild-type and mutant claudin-10 TJ strands

Freeze fracture electron microscopy was employed to analyze the ultrastructure of TJ strands reconstituted by stable expression of CLDN10 constructs in HEK293 cells. In cells expressing either claudin-10a or claudin-10b WT, regular meshworks of continuous TJ strands were detected on the proteo- (P-) and exoplasmic (E-) faces of the plasma membrane (Fig. 6A, D, E), as was shown previously. In the case of claudin-10a G163A, TJ strands were frequently found to be discontinuous, consisting of beaded particles on the P-face (Fig. 6B, C), which indicates altered TJ strand formation of claudin-10.
assembly. Claudin-10b G165A strands were altered even more drastically with highly discontinuous and fragmented TJ strands on both P- and E-faces (Fig. 6F, G). Quantitatively, the proportion of gaps along strands was significantly higher in the mutant cases when compared with the respective wild-types (Fig. 6H). This held also true for claudin-10b G165A strands compared with claudin-10a G163A strands. Additionally, strand abundance appeared to be reduced in both mutant isoforms, an effect more pronounced for claudin-10b G165A. Thus, for both mutants, TJ strand formation is compromised with claudin-10b strands being more severely affected by the mutation.

Dilution and bionic potential measurements

In order to determine whether the mutation affects paracellular channel selectivity, we investigated NaCl dilution potentials and bionic potentials for monovalent cations in MDCK-C7 cell layers stably transfected with wild-type and mutant claudin-10 constructs. Given their endogenous claudin expression pattern, MDCK-C7 monolayers are characterized by high transepithelial resistance and the lack of charge preferences of the paracellular pathway, hence constituting a standard expression system for functional analyses of channel forming claudins.30 Permeability ratios of Cl−/C0 and Na+ turned out to be comparable (PCl/PNa > 1) between claudin-10a WT and G163A expressing cell layers (Fig. 7A). Similarly, mutant claudin-10b retained the wild-type preference for Na+ over Cl− (PNa/PCl > 1) (Fig. 7B). Thus, the mutation does not affect charge preferences of claudin-10 channels. The absence of overt differences in transepithelial resistance between claudin-10a/b WT and G163A/G165A expressing cell layers with comparable permeability ratios indicated no substantial alteration of...
Figure 5  The mutation G163A/G165A differentially affects trans- and cis-interaction properties of claudin-10 isoforms in tight junction (TJ) strands reconstituted in HEK293 cells. (A, B) Cells were transiently transfected with YFP tagged claudin-10 constructs. (A) YFP-claudin-10a WT and G163A show similar enrichment at cell-cell contacts between claudin-expressing cells (arrows) and in intracellular granules. Similarly, YFP-claudin-10b WT shows strong contact enrichment (arrow), indicating trans-interactions and TJ strand formation. In contrast, YFP-claudin-10b G165A frequently distributes more evenly across the membranes of neighboring cells (arrowheads), indicating inhibition of trans-interactions. Nuclei are stained with DAPI (blue). Representative images from 3 biological replicates; scale bars, 5 μm. (B) Enrichment factors computed for individual cell-cell contacts between claudin-expressing cells. Claudin-10b G165A expressing cells show significantly less contact enrichment than cells expressing the corresponding wild-type. In contrast, claudin-10a WT and G163A expressing cells exhibit no significantly different contact enrichment. The dotted line indicates enrichment factor = 1. Enrichment factors >1 are considered to indicate trans-interactions. Different colors represent data from different biological replicates (n = 2–20 per replicate per condition). Welch’s one-way ANOVA with Dunnett’s T3 multiple comparisons test; ns, not significant; ***, P < 0.001. Mean ± SEM. (C) Cis-interaction assay using Förster resonance energy transfer (FRET) between CFP- and YFP-claudin-10 fusion proteins transiently coexpressed in HEK293 cells. Mean FRET efficiencies are significantly reduced for both YFP-/CFP-claudin-10a G163A and YFP-/CFP-claudin-10b G165A when compared with the corresponding wild-types. Complete (claudin-10a) or partial (claudin-10b) restoration of FRET efficiencies can be achieved by coexpression of the respective wild-type isoform. Thus, G163A/G165A perturbs cis-interactions between claudin-10a/b proteins and coexpression of the respective wild-type rescues cis-interactions more efficiently in the case of claudin-10a. One-way ANOVAs followed by two-tailed t-tests with Bonferroni-Holm correction; ns, not significant; **, P < 0.01; *, P < 0.05. Mean ± SEM (n = 11–47). WT, wild-type.
Figure 6  Freeze fracture electron micrograph analysis of HEK293 cells stably expressing claudin-10 constructs. (A) Claudin-10a WT forms meshworks of continuous tight junction (TJ) strands (white arrows) detected on the protoplasmic fracture face (PF) of the plasma membrane. On the exoplasmic fracture face (EF), complementary continuous grooves are evident (white arrowheads). (B) For claudin-10a G163A, similar continuous TJ strands (white arrows) on the PF and grooves (white arrowheads) on the EF are detected to some extent. (C) In most instances, however, discontinuous claudin-10a G163A TJ strands (black arrows) are detected in the form of beaded particles on the PF. The abundance of strands appears to be lower than for claudin-10a WT. (D, E) As is the case with claudin-10a WT, claudin-10b WT forms meshworks of continuous TJ strands (white arrows) on the PF and complementary continuous grooves on the EF (white arrowheads). (F) In contrast, for claudin-10b G165A, fewer and highly discontinuous, fragmented TJ strands (black arrows) can be detected as beaded particles on the PF. On the EF, irregular grooves, pits, and particles are evident (black arrowheads). (G) For claudin-10b G165A, only in few cases, short continuous TJ strands (white arrows) are observed on the PF. Representative images; scale bars, 200 nm. (H) Quantification of strand breaks. Percentages of strand breaks are higher in the mutant cases. In addition, claudin-10b G165A strands exhibit higher break percentages than do claudin-10a G163A strands. Along strands, the particle-free distances between particles or strand segments were determined. The percentage of strand breaks was calculated from the quotient total particle-free length/total strand length. For each sample, $n = 10$ electron micrographs were analyzed. Kruskal-Wallis test followed by post-hoc two-tailed Mann Whitney tests with Holm–Šidák multiple comparisons test; *, $P < 0.05$; ***, $P < 0.001$. Mean ± SEM. WT, wild-type.
Figure 7  Electrophysiological properties do not differ between claudin-10a/b WT and G163A/G165A expressing MDCK-C7 cell layers. (A, B) Permeability quotients as computed from NaCl dilution potentials demonstrate a preservation of charge selectivity in both mutant cases. Paracellular channels reconstituted from claudin-10a G163A retain a preference for Cl\(^-\) over Na\(^+\) (P\(_{Cl}\)/P\(_{Na}\) > 1). Paracellular channels reconstituted from claudin-10b G165A favor the permeation of Na\(^+\) over Cl\(^-\) (P\(_{Na}\)/P\(_{Cl}\) > 1) as do claudin-10b WT channels. Different symbols represent data from different stably transfected clones. Dashed lines indicate P\(_{Na}\) = P\(_{Cl}\).

(C, D) Transepithelial resistance is not overtly different between claudin-10a/b WT and G163A/G165A expressing MDCK-C7 cell layers. Permeability quotients as computed from NaCl dilution potentials are plotted against transepithelial resistance. Different symbols represent data from different stably transfected clones (blue: claudin-10a/b WT, red: claudin-10a/b G163A/G165A, black: vector control). Dashed lines indicate P\(_{Na}\) = P\(_{Cl}\). (E) For both claudin-10b WT and G165A, monovalent cation permeability sequences as computed from bionic potentials correspond to Eisenman sequence X (Na\(^+\) > Li\(^+\) > K\(^+\) > Rb\(^+\) > Cs\(^+\)). For both claudin-10b WT and G165A, data from three different stably transfected clones (#) are shown. Mean ± SEM (n = 6–13). (F) Estimation of paracellular pore diameter from bionic potentials for NH\(_4\)\(^+\) and various organic cations. Pore diameters do not differ between claudin-10b WT and G165A (maximally 5 Å). For both claudin-10b WT (blue) and G165A (red), data from two stably transfected clones are shown as indicated by differing symbols (per clone n = 6–11 for NH\(_4\)\(^+\), n = 6–8 for MA, n = 6–8 for DMA, n = 3–6 for TriMA, n = 2–6 for TeMA). Two-tailed t-tests; ns, not significant. Square root of mean ± SEM. WT, wild-type; MA, methylammonium; DMA, dimethylammonium; TriMA, trimethylammonium; TeMA, tetramethylammonium.
single channel ion conductivity (Fig. 7C, D). Monovalent cation permeability sequences for claudin-10b WT and G165A expressing cell layers both corresponded to Eisenman sequence X (Na\(^+\) > Li\(^+\) > K\(^+\) > Rb\(^+\) > Cs\(^+\)), indicating that channel selectivity was unaltered (Fig. 7E). Pore diameter was estimated from bionic potentials for the monovalent cations NH\(_4\)^+\), methylammonium, dimethylammonium, trimethylammonium, and tetramethylammonium to be identical for both claudin-10b WT and G165A (maximally 5 Å) (Fig. 7F).

**Discussion**

In this study, we demonstrate pathogenicity of a novel missense variant in CLDN10 by investigating associated molecular alterations and phenotypic traits. Homology modeling predicted destabilization of the claudin transmembrane four-helix bundle, leading to perturbed oligomerization processes.

Indeed, in HEK293 cells which do not form TJs by themselves, reconstituted mutant claudin-10 TJ strands exhibited defective architecture and compromised protein interaction properties. Ultrastructural analysis of mutant TJ strands showed fragmentation, discontinuity, and reduced abundance. *Cis*-interactions between claudin-10 proteins as probed by the assessment of FRET efficiencies were perturbed in both mutant isoforms. Throughout, claudin-10b G165A displayed more severe alterations than claudin-10a G163A, in line with structural implications from homology modeling. Fluorescence microscopy suggested compromised claudin-10 *trans*-interactions only for claudin-10b G165A. Of note, however, a distinction between a primary *trans*-interaction defect and a merely indirect effect induced by the inhibition of *cis*-interactions cannot be made at this point. Together, the observed defects in claudin-10 interaction properties do well explain the perturbation of TJ architecture evident on electron micrographs by an impairment of TJ strand assembly.

Our electrophysiological findings in MDCK-C7 cells demonstrate that mutant claudin-10 is generally capable of forming paracellular channels with preserved selectivity and conductivity if inserted into pre-existing TJ structures. In line with results from homology modeling, G163A/G165A hence does not affect the pore lining and in turn does not alter the interaction with permeating ions. Instead, the mutation impedes claudin oligomerization and TJ strand assembly which manifests in various ways, depending on cell type or tissue: (i) In HEK293 cells, reconstituted TJ strands are fragmented; (ii) in MDCK-C7 cells and in epithelial cells from secretory tubules of eccrine sweat glands, TJ strands contain functional claudin-10 channels even if absolute claudin-10 abundance at the TJ may be reduced; and (iii) small salivary glands lack claudin-10 altogether.

We conclude that mutant proteins assemble to form functional paracellular channels in a tissue-specific manner which may depend on TJ composition and regulatory factors. In regards to our data we speculate that the specific claudin environment promotes stabilization of mutant claudin-10 in secretory tubules of eccrine sweat glands, yet is unfavorable in those of small salivary glands. Phenotypically, sweat secretion rates were preserved to some extent, whereas the rate of saliva secretion was unaffected but ionic saliva composition showed marked abnormalities. These data indicate diverging roles of claudin-10 in glandular processes of fluid secretion. Given the mosaic pattern of claudin expression in the kidney's TAL with mutual exclusion of claudin-3/-16/-19 and claudin-10 TJs,\(^3\) mutant claudin-10 is not expected to be inserted into TAL TJs. For our patient, this remains elusive in the absence of a kidney biopsy. However, claudin-10 was found to be absent from the TAL of two other HELIX patients.\(^1,5\)

Of note, our data contrast with previous reports of xerostomia in HELIX patients.\(^2,5,28\) However, quantitative assessment of saliva secretion has previously been performed solely by Hadj-Rafia et al.\(^2\) in three patients. Our dental findings indicate impaired enamel formation to be a novel phenotype of HELIX syndrome. Indeed, amelogenesis imperfecta has previously been causally related to defects in claudin-16, -19, and -3.\(^34,36\) Moreover, claudin-10 expression has been found in mature murine ameloblasts.\(^37\)

We remark that all pathogenic CLDN10 variants reported to date affect claudin-10b with the associated phenotype matching its bodily distribution. An impairment of claudin-10a function may thus not become clinically evident.

Regarding TAL ion transport in the absence of functional claudin-10b TJs, paracellular Na\(^+\) reabsorption is expected to be markedly reduced while paracellular Ca\(^2+\) and Mg\(^2+\) transport via claudin-3/-16/-19 TJs should be enhanced.\(^33,38\) These TJs are spatially separated from those comprised of claudin-10b and expand in its absence.\(^3\) Our results from loop diuretic responsiveness testing suggest transcellular Na\(^+\) transport to partially compensate for the defect in paracellular reabsorption. In addition, hydrochlorothiazide test results and secondary hyperaldosteronism suggest adaptive changes in the distal nephron, accounting for larger Na\(^+\) transport capacities with concomitant K\(^+\) secretion upon increased NaCl delivery. These mechanisms are well in line with limited salt-wasting on baseline. Pathophysiologically, defective TAL NaCl reabsorption has two major implications. First, it leads to renal NaCl wasting despite compensatory efforts, and second, it reduces the buildup of interstitial osmolality, accounting for hyposthenuria. The ensuing NaCl deficit in the extracellular fluid compartment induces metabolic alkalosis by volume contraction. Activation of the renin-angiotensin-aldosterone system following the reduction of effective arterial blood volume is implicated in maintaining the alkalotic state\(^39\) and leads to the release of aldosterone which promotes a secondary KCl deficit. Hypokalemia in turn aggravates metabolic alkalosis.\(^40\) Thus, in HELIX syndrome, metabolic alkalosis is caused by net deficits of NaCl and KCl.

Reduced buildup of interstitial osmolality in the renal medulla is consistent with the preserved renal aquaporin-2 response to desmopressin in another HELIX patient.\(^1\) Prolonged thirsting unmasked a defective urine concentration mechanism in our patient, similar to previous results.\(^1\) The defective urine concentration ability likely accounts for nocturnal enuresis in our patient. In particular, polyuria is unlikely to result from xerostomia which has been suggested previously.\(^3\)

Given the above pathophysiological considerations, the question of suitable treatment options arises. Major goals...
of medical therapy should be to ameliorate polyuria and to meet the imminent danger of acute-on-chronic hypokalemia. With defective claudin-10, long-term K\(^+\) homeostasis appears to be achieved when plasma K\(^+\) is low enough to limit urinary and exocrine K\(^+\) losses to an amount that matches its input. Our data indicate that aggressive KCl substitution, therefore, is not expected to be of sustained benefit. Rather, mineralocorticoid receptor antagonists or blockers of the epithelial Na\(^+\) channel (ENaC) should be employed.

Clearly, one drawback of our study is its confinement to the assessment of a single patient. However, this is only the sixth reported instance of a homozygous **CLDN10** variant associated with human disease and the fourth such variant affecting both protein isoforms.\(^1\)\(^{–}\)\(^5\) In conclusion, we demonstrated that claudin-10a/b G163A/G165A causes a variation of HELIX syndrome based on defective claudin-10 TJ strand assembly. Thereby, claudin-10b is more severely affected than claudin-10a. Phenotypic traits result from the tissue-specific deficiency in insertion of aberrant claudin-10 into TJs, a process that may depend on the presence of other TJ components. Although both mutant isoforms are generally capable of forming paracellular channels with preserved electrophysiological properties if inserted into pre-existing TJs, their absolute TJ abundance may be reduced and thereby determine the degree of paracellular route dysfunction. Whether structural interference with claudin oligomerization constitutes a more general mechanism of pathogenicity in HELIX syndrome remains to be determined in future studies.

**Author contributions**

SS, DG, and JH conceived the study. SS, JP, and DG performed experiments. SS, JP, RS, and DG analyzed experimental data. SS performed clinical testing and analyzed the data. JP performed homology modeling. SS, SG, and AR contributed clinical data. SN and CB performed genetic analyses. SS wrote the manuscript, with contributions and editions from JP, DG, and JH. All authors approved the final version of the manuscript.

**Conflict of interests**

The authors declare no conflict of interests.

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**Appendix A. Supplementary Data**

Supplementary Data to this article can be found online at https://doi.org/10.1016/j.gendis.2021.06.006.

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