Nanoscale segregation of channel and barrier claudins enables paracellular ion flux

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The paracellular passage of ions and small molecules across epithelia is controlled by tight junctions, complex meshworks of claudin polymers that form tight seals between neighboring cells. How the nanoscale architecture of tight junction meshworks enables paracellular passage of specific ions or small molecules without compromising barrier function is unknown. Here we combine super-resolution stimulated emission depletion microscopy in live and fixed cells and tissues, multivariate classification of super-resolution images and fluorescence resonance energy transfer to reveal the nanoscale organization of tight junctions formed by mammalian claudins. We show that only a subset of claudins can assemble into characteristic homotypic meshworks, whereas tight junctions formed by multiple claudins display nanoscale organization principles of intermixing, integration, induction, segregation, and exclusion of strand assemblies. Interestingly, channel-forming claudins are spatially segregated from barrier-forming claudins via determinants mainly encoded in their extracellular domains also known to harbor mutations leading to human diseases. Electrophysiological analysis of claudins in epithelial cells suggests that nanoscale segregation of distinct channel-forming claudins enables barrier function combined with specific paracellular ion flux across tight junctions.

During development and homeostasis, tissues not only have to tightly control passage of small molecules and ions through transepithelial transport mechanisms but also via paracellularly located adhesion complexes including tight junctions (TJs)1. TJs form apical cell–cell contacts in epithelia2 and restrict the passage of pathogens, small molecules, ions, and water by very close paracellular membrane contacts2. Claudins are transmembrane proteins that form ~10 nm thick strands interwoven into TJ meshworks3 and associate intracellularly with numerous scaffolding proteins and the cytoskeleton4. These strands can act as paracellular diffusion barriers against small and large solutes, e.g., when composed of barrier claudins like claudin-1 (Cldn1)5,6 or Cldn37. In addition, size and charge selective ion/water channels are formed within strands by Cldn28–10, Cldn10a/b11,12, Cldn1513–15, and Cldn1616,17.

Early electrophysiological measurements, mathematical modeling18, as well as structural and functional data on channel and barrier claudins19,20 led to a model in which paracellular pores formed by selected claudins integrate into TJ strands. How TJ meshworks composed of single or multiple claudins are organized at the nanoscale to integrate paracellular barrier and channel functions is, however,
unknown. Besides amino acids that enable paracellular charge selectivity, comparison of the primary protein sequences of barrier- vs. channel-forming claudins has not uncovered distinctive channel features that could explain their distinct organization. Moreover, freeze fracture electron microscopy (FFEM) images of endogenous or reconstituted claudin strands failed to reveal specific ultrastructural features of barrier and channel claudins, potentially due to structural effects of strong chemical fixation required for this analysis. Super-resolution fluorescence microscopy now enables molecular imaging of living cells and fixed tissues with nanometer resolution and has been applied to visualize selected claudins.

Here we use super-resolution stimulated emission depletion (STED) microscopy in live and fixed cells and tissues to reveal the nanoscale TJ organization of all 26 mammalian claudins. We find that only half of the individually expressed claudins form TJ strands or meshworks on their own, whereas TJs formed by multiple claudins display nanoscale organization principles of intermixing, integration, induction, segregation, and exclusion of strand assemblies. The notable nanoscale segregation of channel- and barrier-forming claudins within individual strands is mediated, at least in part, by claudin extracellular domains, while claudin association with ZO-1 adaptor proteins is dispensable. We hypothesize that segregation of channel and barrier claudins is a key mechanism for proper claudin channel function, that is compromised in certain patients with claudin missense mutations (e.g., N48K). Functional ion permeability measurements of claudins re-expressed in a TJ-strand free epithelial cell line indicated that segregation enables the parallel flow of oppositely charged ions and increases the ion specificity. Our data show that paracellular ion transport is mediated by segregated TJ meshworks. These findings may have wide implications for cell physiology and our understanding of tissue barrier function.

**Results**

**STED microscopy reveals the nanoscale organization of mammalian claudins in living cells and tissue.**

Claudins, the key constituent proteins of TJs, are tetraspan transmembrane proteins with two extracellular loops, encoded by 26 genes in mice and 25 genes in humans. They determine the paracellular barrier and flux properties of TJs by forming up to one micrometer-sized meshworks at the most apical part of the lateral membrane of epithelial cell-to-cell contacts (Fig. 1a). Super-resolution STED microscopy with >50 nm lateral (XY) resolution enabled us to visualize endogenous Cldn3 within TJs of mouse duodenum labeled with specific antibodies (Fig. 1c). While single strands and close strand assemblies below 50 nm distance could not be resolved within the intestinal TJ meshwork, large individual meshes and the overall TJ thickness of ~500 nm became apparent, consistent with earlier observations from FFEM. To characterize the nanoscale organization of individual claudins within TJs, and to overcome limitations of antibody labeling, we reconstituted TJ meshworks formed by individual claudins (and
their isoforms) in COS-7 cells, e.i., non-polarized fibroblast-like cells isolated from African green monkey kidney (Fig. 1b). These non-polarized cells lack endogenous tight junctional transmembrane proteins including claudins, but can still form TJ meshworks that colocalize with ZO-1, an important scaffolding TJ protein when claudins are exogenously expressed. YFP-tagged Cldn3 visualized in the flat overlapping areas of neighboring COS-7 cells formed large meshworks, comprised of multiple individual claudin strands and small meshes (Fig. 1d), greatly resembling the organization of endogenous Cldn3 in intestinal tissue (Fig. 1c). Similarly organized meshworks were formed by untagged Cldn3 labeled with anti-Cldn3 antibodies and by mouse Cldn3 tagged at either its N- or C-terminal end with an YFP (Supplementary Fig. 1d). Moreover, TJ-like meshworks formed by Cldn3 or related claudins (i.e., Cldn1, Cldn2, Cldn10a) colocalized generally with ZO-1 endogenously expressed in COS-7 cells but only peripherally with actin (Supplementary Fig. 1a–c). Since meshwork morphology might be affected by chemical fixation (Supplementary Fig. 1e) we analyzed the nanoscale organization of SNAP-Cldn3 in living COS-7 cells. Live STED imaging revealed strand dynamics, the formation of large Cldn3 meshes in living cells and a similar Cldn3 meshwork organization as in fixed cells with 4% PFA (Fig. 1e; Supplementary Fig. 1d; Supplementary Movie 1). Individual claudin strands of ~10 nm thickness can form intricate meshworks of various morphology. To characterize individual TJ strands quantitatively, we measured isolated SNAP-Cldn3 strands in fixed and living COS-7 cells and found characteristic widths of 59 ± 11 nm and 69 ± 14 nm, respectively (Fig. 1f). The slightly larger value in living cells could be due to strand mobility and/or a lower signal-to-noise ratio in cell culture medium compared to fixed cell mounting medium. Of note STED images generally show a lower signal-to-noise ratio than imaging techniques that were previously used to image claudin strands including widefield, confocal, and SIM. We note that single-molecule localization microscopy of YFP-Cldn3 in fixed HEK cells yielded similar strand widths. Collectively, these results show that STED microscopy can resolve meshes of single strands in living or fixed cells. Therefore, STED is capable of revealing the nanoscale molecular architecture of TJ assemblies in cells and tissue.

### Mammalian claudins show differences in strand formation and meshwork organization

Based on these results and taking advantage of the excellent structural preservation of chemically fixed strand morphologies we analyzed all 26 mammalian claudins, tagged at their N-termini with YFP and stained with Atto647N-labeled anti-GFP nanobodies in fixed COS-7 cells (Supplementary Fig. 2). Since half of the analyzed claudins formed no clear meshworks we wanted to objectively classify TJ strand formations based on STED intensity texture features that are spatial intensity variations across a scale of 20–200 nm, as explained in the method section. Indeed, hierarchical clustering of basic intensity texture features largely confirmed our separation between meshwork-forming from non-meshwork forming claudins (Supplementary Figs. 2 and 3). Most claudins previously classified as classic claudins were able to polymerize into TJ strands; exceptions were Cldn4, Cldn8 and Cldn17. We confirmed the mostly uniform and unstructured distribution of SNAP-Cldn4 or SNAP-Cldn8 by STED imaging of living COS-7 cells (Supplementary Fig. 5i). In contrast, all claudins previously attributed as non-classic, except for Cldn11 and Cldn20, did not form TJ strands on their own. Instead, they appeared either as uniformly distributed, small punctae, irregular clusters or small strands. Cldn22 and Cldn27 localized partly to ER (Supplementary Fig. 2 and 3b, c). Non-meshwork forming claudins might require co-assembly with classic claudins or other TJ proteins to polymerize into TJ assemblies. We did not observe any correlation between the ability to form strands with claudins either acting as a barrier or a channel (Supplementary Fig. 2 and 3). Claudins previously implicated in bivalent cation transport such as Cldn2, Cldn3 or Cldn6 were unable to form detectable strands and meshworks under these conditions (Supplementary Fig. 2 and 3).

To further classify TJ meshwork formation by individual claudins, STED images of meshwork-forming claudins were segmented and features of meshwork morphology such as average mesh size, branch length, and the number of branches were extracted and combined with intensity texture features (Fig. 2a). Using hierarchical clustering and principal component analysis, we identified three classes of meshworks: Class A claudins, comprising Cldn7, Cldn10a (anion channel), Cldn19a/b, and Cldn20, formed large mesh sizes. Claudins of class B included cation channel-forming Cldn2, Cldn10b, Cldn15, as well as the barrier-forming Cldn3, Cldn5, and Cldn14, and formed meshes of much smaller sizes. Finally, class C claudins, such as Cldn1, Cldn6, Cldn9, and Cldn11 assembled into very dense meshwork structures with predominantly parallel strands (Fig. 2b; Supplementary Fig. 4). On rare occasions, single strands of Cldn11 were observed and measured to have a similar width as Cldn3 strands (Supplementary Fig. 3d; Fig. 1f). Interestingly class C claudins are known to form particularly tight seals in epithelia, a feature that appears to be recapitulated upon exogenous expression in non-polarized COS-7 cells.

Taken together, we found that out of half of all mammalian claudins are able to form meshworks and this ability seems to correlate with protein sequence homology, but not with the far known barrier vs. channel function. This suggests that meshwork formation may be an intrinsic feature of claudin 3D-structure and polymer assembly and barrier as well as channel claudins can assemble into similarly organized meshworks that may facilitate their co-assembly.

### Claudin combinations display nanoscale organization principles of intermixing, integration, induction, segregation and exclusion of strand assemblies

TJ meshworks are mostly composed of multiple barrier- and/or channel-forming claudins. Copolymers of these claudins may form at different length scales ranging from mixed strands to small separated clusters or larger segments within meshworks thereby creating different local ion permeabilities as suggested previously (Fig. 3a). To investigate the nanoscale organization of claudin copolymers we co-expressed Cldn3 or Cldn4, two typical barrier-forming claudins found in the respiratory, urinary and gastrointestinal tracts, with other claudins in COS-7 cells and analyzed overlapping areas by two-color STED microscopy (Fig. 3b; Supplementary Fig. 5a; Supplementary Fig. 6a). Strikingly, five different types of behaviors of co-expressed claudins were observed: (i) Intermixing of barrier-forming Cldn1, Cldn5, Cldn6, Cldn7, Cldn9, Cldn14, and Cldn19b with Cldn3. (ii) Integration of Cldn4, a claudin incapable of forming TJ strands on its own, into Cldn3 strands. (iii) Induction of co-assembled strands and meshworks from two non-meshwork forming claudins, Cldn4 and Cldn8. (iv) Segregation of channel-forming Cldn2, Cldn10a, Cldn10b and Cldn15 from Cldn3 within individual TJ strands and meshworks resulting in alternating areas of channel-forming and barrier-forming TJs. Finally, (v) exclusion of Cldn3 and Cldn11 from each other by formation of independent TJ meshworks. These five different types of nanoscale organization of claudin copolymers were also observed in the mouse fibroblast cell line 3T3 and in COS-7 cells using a GFP-nanobody labeled with a more hydrophilic fluorophore (Supplementary Fig. 7e, f).

The intermixing, segregation, and exclusion behavior of the various claudins with respect to Cldn3 was confirmed by quantitative colocalisation analysis based on Pearson correlation of pixel intensity, where positive values indicate intermixing and negative values indicate segregation or exclusion (Fig. 3c; Supplementary Fig. 5b–d). This analysis revealed that Cldn3 intermixing with other barrier-forming claudins was indistinguishable from colocalization of Cldn3 with itself.
In contrast, channel-forming claudins such as Cldn2, Cldn10a, Cldn10b, and Cldn15 were nearly completely segregated from Cldn3 within the TJ-like meshwork, as indicated by a significantly lower and sometimes even negative colocalization index (Fig. 3c). A similarly negative colocalization index was seen for Cldn3 and Cldn11 that formed mutually exclusive TJ meshworks (Supplementary Fig. 5d). To test whether these behaviors reflect a general organizational principle of TJ formation by claudins we analyzed the barrier-forming Cldn1. We found Cldn1 to intermix with Cldn3, but segregate from channel-forming Cldn2, Cldn10a, Cldn10b, and Cldn15 (Supplementary Fig. 5f).

To challenge these findings by an alternative approach, we analyzed claudin interactions in living cells on a molecular scale of ~10 nm.

Fig. 2 | Automated analysis of TJ meshwork enables the overall classification of all meshwork-forming claudins. a Hierarchical clustering of all meshwork forming mammalian claudins expressed in COS-7 cells into three different classes using an automated analysis based on Haralick texture features and image segmentation. All claudins were N-terminally tagged with YFP or GFP and boosted with α-GFP-NB-Atto647N. Columns are centered and unit variance scaling was applied to columns. Correlation distance with average linkage was used for clustering of rows and columns. For the Haralick texture features one pixel (px) equals 20 nm. Three classes are labeled in cyan (class A), yellow (class B), and magenta (class C). Claudins were color-coded by function, as barrier (black), cation channel (green), anion channel (red), cation and water channel (blue), and heteromeric ion channel formed by two different claudins (orange). Color code of heatmap represents unit variance scaling and represents number of standard deviations. Based on a dataset of 15 claudins with 202 total images. Source data are provided as a Source data file. b Representative STED images of TJ-like meshworks from three identified meshwork classes (A, B, C) in (a). ROIs were taken from cell–cell overlaps of two transfected cells. All claudins were N-terminally tagged with YFP and boosted with α-GFP-NB-Atto647N. Scale bar, 200 nm (b).
Fig. 3 | Claudins form characteristic copolymers including segregation of channel-forming Claudins from Cldn3. a. Scheme is illustrating predicted claudin-claudin organization patterns. Two claudins (yellow and magenta) that are compatible with each other tightly colocalize in the TJ (pink). Incompatible claudins separate into different strands or even larger claudin-specific parts. This separation might facilitate permeability of specific ions (cyan) over the TJ meshwork.

b. Representative STED images of TJ-like meshwork of the observed five organization patterns formed by the indicated co-expressed claudins (SNAP-tagged (yellow; BG-Atto590) and YFP-tagged (magenta; α-GFP-NB-Atto647N)) in fixed COS-7 cells are shown. Tight colocalization: intermixing of mesh-forming claudins, integration of non-mesh-forming with a mesh-forming claudin or induction (de novo mesh-forming of co-expressed non-mesh-forming claudins). Separated claudins: segregation and exclusion.

c. Pearson correlation analysis of Cldn3 co-expressed with barrier-forming (gray) or channel-forming (magenta) Claudins. Co-expression of Cldn3 with Cldn3 (yellow) served as positive control. Data represent the mean ± SD. Every n represents the Pearson of one TJ-like meshwork. n(Cldn3 + Cldn3) = 55; n(Cldn3 + Cldn1) = 54; n(Cldn3 + Cldn5) = 20; n(Cldn3 + Cldn6) = 20; n(Cldn3 + Cldn7) = 17; n(Cldn3 + Cldn9) = 15; n(Cldn3 + Cldn14) = 19; n(Cldn3 + Cldn19b) = 25; n(Cldn3 + Cldn2) = 36; n(Cldn3 + Cldn10a) = 30; n(Cldn3 + Cldn10b) = 24;

n(Cldn3 + Cldn15) = 42; from 3–6 independent experiments; one-way ANOVA with Dunnett’s multiple comparison test; **P ≤ 0.001, ns (not significant).

d. Spectral FRET analysis of Trq2-Cldn3 alone (blank; negative control) or co-expressed with indicated YFP-tagged barrier Cldn3 (yellow; positive control) and channel-forming Cldn2 or Cldn15 (magenta) in HEK293 cells. Data represent the mean ± SD. Every n represents one cell–cell contact. n(negative control) = 57, from one experiment; n(Cldn3 + Cldn3) = 312; n(Cldn3 + Cldn2) = 162; n(Cldn3 + Cldn15) = 154; all from 4 to 5 independent experiments; one-way ANOVA with Dunnett’s multiple comparison test; **P ≤ 0.001, ns (not significant).

e. Representative STED image of segregating SNAP-Cldn3 (yellow; BG-JF646) and Halo-Cldn15 (magenta; CA-Atto590) in living COS-7 cells. Image noise was removed for visualization using noise2Void®.

f. Representative STED images of intermixing and segregation of SNAP-Cldn3 (yellow; BG-Atto990) with YFP-tagged Cldn1 and Cldn15 (both magenta; α-GFP-NB-Atto647N), respectively expressed in Caco-2 cells. g. Representative STED images of the endogenous formed TJ in cryo-sections from mouse duodenum immunostained for Cldn3 (yellow; 2nd-Atto647N) and Cldn15 (magenta; 2nd-AF594). White arrows indicate regions in the TJ that shows the segregation of Cldn3 and Cldn15 strands. A Gaussian blur with a sigma of 20 nm was applied. All representative images derive from 3 independent experiments. Scale bars, 500 nm (g), 200 nm (b, e, f, magnification in g). Source data are provided as a Source data file.
Channel-forming claudins form segregated meshworks in cells and tissue

Channel-forming claudins such as Cldn2, Cldn10a, Cldn10b, and Cldn16 in combination with Cldn19 are highly abundant in the kidney and facilitate Na\(^+\), Cl\(^-\), and Mg\(^{2+}\) resorption\(^{10,16,43}\). We thus wanted to know whether channel claudins would not only segregate from barrier Cldn3 but also from each other to preserve their unique channel properties or create localized ion permeabilities. When co-expressed Cldn2, i.e., a paracellular Na\(^+\) channel, with other channel-forming claudins such as the Cl\(^-\)-selective Cldn10a or the Na\(^+\)-selective claudins Cldn10b and Cldn15, or with barrier Cldn1 in COS-7 cells, we observed segregated TJ-like meshworks as evidenced by their low colocalization index in STED analyses (Fig. 4a, b; Supplementary Fig. 6b). FRET measurements in HEK293 and COS-7 cells confirmed the observed lack of interaction between Cldn2 and Cldn10a on a scale below 10 nm in living cells (Fig. 4c; Supplementary Fig. 7c). Segregation of channel-forming Cldn2 and Cldn10a was also found in living COS-7 cells visualized by live STED imaging (Fig. 4d) and in chemically fixed not fully-polarized kidney epithelial cells (Fig. 4e, f).

In the thick ascending limb of Henle's loop of the nephron paracellular Na\(^+\) and Mg\(^{2+}\) flux is mediated by three claudins Cldn10b/16/19 that displayed a low degree of colocalisation (Fig. 4g, h; Supplementary Fig. 7d). Apart from their segregation behavior in proximal tubules, Cldn2 and Cldn10a displayed considerable heterogeneity with respect to strand length between different tubules and even within the same tubule without any indication of a claudin or location-specific strand length (Fig. 4i).

We conclude that TJ meshworks formed by different claudins not only are segregated between barrier- and channel-forming claudins but also between distinct channel-forming claudin isoforms, possibly to spatially restrict ion flux on the nanoscale within cells and tissues.

Claudin protein levels and extracellular loop sequences regulate claudin segregation

Next, we set out to determine the molecular basis of the observed segregation of distinct channel claudins. As the ability to form segregated claudin meshworks may depend on relative protein copy numbers, we first tested the effect of different protein expression ratios. Channel-forming Cldn2 and Cldn10a displayed segregation with low colocalisation at all expression ratios tested, whereas strand lengths positively correlated to expression levels (Fig. 5a).

While these data demonstrate that TJ strand length is determined by claudin copy numbers and polymerization properties, they fail to provide insight into the mechanism underlying channel claudin segregation. In principle, two main mechanisms could mediate the segregation of distinct channel claudins: (i) claudin segregation might be a consequence of their differential association with other factors, e.g., membrane lipids such as cholesterol, which was shown to be required for TJ formation\(^{44}\) and to associate with claudins in distinct membrane domains\(^{45}\), or TJ-associated intracellular scaffold proteins, most notably the ZO-1 that has been proposed to promote TJ assembly via multiple mechanisms\(^{46-48}\) including liquid-liquid phase separation\(^{49}\). (ii) Alternatively, claudin segregation might reflect an intrinsic property of claudin proteins themselves that could either relate to their channel activity or the ability to assemble via their extracellular and transmembrane domains.

We tested these mechanisms by combined chemical and genetic approaches. Treatment of COS-7 cells with the HMG-CoA reductase blocker mevastatin or with the cholesterol-sequestering drug Methyl-\(\beta\)-cyclodextrin (MβCD) markedly reduced plasma membrane cholesterol levels (Supplementary Fig. 8a) but did not impair claudin segregation (Fig. 5b). Deletion of the C-terminal PDZ binding motif (required for binding to ZO-1) or even almost the entire C-terminal cytoplasmic domain (Fig. 5c) in either Cldn2 and Cldn10a or Cldn3 and Cldn15 resulted in smaller and less frequent meshworks, but did not affect the ability of claudins to segregate along TJ strands (Fig. 5d, e; Supplementary Fig. 8d, e). Hence, claudin segregation does not appear to be a consequence of their association with extrinsic factors such as cholesterol, proteins that bind to phosphorylated residues within the C-terminal tail, or PDZ domain proteins such as ZO-1.

We therefore pursued the alternative hypothesis that claudin segregation is a protein-intrinsic property built into their 3D architecture. The fact that distinct channel-forming claudins segregated from each other as well as from barrier-forming claudins indicates that segregation might be a consequence of ion flux. Potentially, this stabilizes claudin channel polymers. We tested this possibility by...
Channel-forming claudins form segregated meshworks in non-polarized cells, epithelial cells and the murine proximal tubule.

**a** Representative STED image of TJ-like meshwork formed by SNAP-Cldn2 (yellow; BG-Atto590) co-expressed with YFP-tagged Cldn2, Cldn1, Cldn10a, Cldn10b, and Cldn15 (all magenta; α-GFP-NB-Atto647N) in COS-7 cells.

**b** Pearson correlation analysis of Cldn2 co-expressed with Cldn2 (yellow), Cldn1 (gray) and Cldn10a, Cldn10b, Cldn15 (magenta). Data represent the mean ± SD. Every n represents the Pearson of one TJ-like meshwork. n(Cldn2 + Cldn2) = 21; n(Cldn2 + Cldn1) = 29; n(Cldn2 + Cldn10a) = 35; n(Cldn2 + Cldn10b) = 28; n(Cldn2 + Cldn15) = 22; from 3 to 5 independent experiments; one-way ANOVA with Dunnett’s multiple comparison test; ***P ≤ 0.001, ns (not significant).

**c** Spectral FRET analysis of Trq2-Cldn2 (blank; negative control) or co-expressed with YFP-tagged Cldn2 (yellow; positive control) and Cldn10a (magenta) in HEK cells. Data represent the mean ± SD. Every n represents one cell–cell contact. n(negative control) = 126, from one experiment; n(Cldn2 + Cldn2) = 405; n(Cldn2 + Cldn10a) = 115; from 4 to 10 independent experiments; one-way ANOVA with Dunnett’s multiple comparison test; ***P ≤ 0.001, ns (not significant).

**d** Representative STED image of segregating SNAP-Cldn2 (yellow; BG-JF646) and Halo-Cldn10a (magenta; CA-Atto590) in living COS-7 cells. Image noise was removed for visualization using noise2Void.

**e** Representative STED image of segregation of SNAP-Cldn2 (yellow; BG-Atto590) and YFP-Cldn10a (magenta; α-GFP-NB-Atto647N) localized in an overlap region of two transfected MDCKC7 cells. F Pearson correlation analysis of Cldn2 co-expressed with Cldn2 (yellow) and Cldn10a (magenta) in MDCKC7. Data represent the mean ± SD. Every n represents the Pearson of one TJ-like meshwork. n(Cldn2 + Cldn2) = 17; n(Cldn2 + Cldn10a) = 16; from 3 independent experiments; Mann-Whitney test, two-tailed; ***P ≤ 0.0001.

**f** Representative STED image of the TJ in mouse proximal tubule immunostained for Cldn2 (yellow; 2nd-Atto647N) and Cldn10a (magenta; 2nd-AF594). The white rectangle indicates the area of the magnification.

**g** Pearson correlation analysis of Cldn2 and Cldn10a in three mouse proximal tubules. Data represent the mean ± SD. Every n represents the Pearson of one STED image of the tubular TJ. n(Tubule I) = 9; n(Tubule II) = 12; n(Tubule III) = 11; one-way ANOVA Tukey’s multiple comparison test; ns (non-significant).

**h** Strand length analysis of Cldn2 and Cldn10a strands in three proximal tubules (Tub I-III). Data represent the mean ± SD; n = 4. Mean of the measured strands for each tubule; strands Tubule I: 541 (Cldn2)/664 (Cldn10a), strands Tubule II: 747 (Cldn2)/791 (Cldn10a), strands Tubule III: 693 (Cldn2)/828 (Cldn10a); one-way ANOVA with Tukey’s multiple comparison test; ***P ≤ 0.001, ns (non-significant). All representative images derive from 3 independent experiments. Scale bars, 500 nm (g), 200 nm (a, d, e, magnification in g). Source data are provided as a Source data file.
mutating I66 in the pore center of Cldn2 to Cysteine to allow pharmacological blockade of ion permeation by application of the cysteine-reactive reagent MTSET. MTSET treatment of mutant Cldn2I66C expressed in COS-7 cells did not affect its ability to form a TJ-like meshwork. When we expressed Cldn2I66C with Cldn10a we observed segregation in both control (DMSO) and MTSET conditions (Fig. 5f). Hence, channel activity is dispensable for channel-forming claudin segregation.

A pathogenic mutation (N48K) in the first extracellular loop of Cldn10b causes HELIX (hypohidrosis, electrolyte imbalance, lacrimal gland dysfunction, ichthyosis, and xerostomia) syndrome. Cldn10b dysfunction alters the paracellular cation permeability in HELIX syndrome patients, resulting in anhidrosis and kidney damage. Consistently, we found that Cldn10bN48K fails to form Cldn10b WT-like meshworks (Supplementary Fig. 8b) and is not integrated into TJ meshworks formed by Cldn3 and Cldn10b (Fig. 5g; Supplementary Article...
Fig. 5 | Claudin segregation is conserved in the extracellular loops and does not depend on cholesterol or the binding to ZO-1. a, Representative STED images, immunoblot analysis, strand length, and Pearson correlation analysis of TJ-like meshwork formed by SNAP-Cldn2 (yellow; BG-Atto590) and YFP-Cldn10a (magenta; α-GFP-NB-Atto647N) with different expression ratios (3:1, 3:3) in COS-7 cells. Blotting was performed against SNAP tag and YFP tag. HSP70 (70 kDa) served as loading control. For the strand and Pearson analysis the same TJ-meshworks were used. Data represent the mean ± SD. For the strand length analysis every n represents the mean of 40 strands of Cldn2 and Cldn10a per meshwork from 12 TJ-like meshworks (n = 12); one-way ANOVA Tukey’s multiple comparison test; **P < 0.0001, ***P < 0.01, ns (not significant); from 3 independent experiments; for the Pearson analysis every n represents one TJ-like meshwork (n = 12); Mann-Whitney test, two-tailed; ns (not significant) (P = 0.8623) from 3 independent experiments. b, Representative STED images of control and cholesterol-depleted COS-7 cells expressing SNAP-Cldn2 (yellow; BG-Atto590) and YFP-Cldn10a (magenta; α-GFP-NB-Atto647N). White arrows point at claudin-containing vesicles that were increasingly formed under cholesterol depletion conditions. c, Schematic representation of a claudin in the plasma membrane. The sites for the ECL exchange of Cldn2ECL10a and the PDE- and C-term deletion mutants (ΔPDE and ΔCCT) are pointed out with red lines. d, Representative STED images of COS-7 cells co-expressing SNAP-Cldn2ECL10a/Cldn3ECL10a (yellow; BG-JF646) with YFP-Cldn10aΔPDZ/-Cldn3ECL10a (magenta; α-GFP 2nd-AF594) or SNAP-Cldn2ECL10a/Cldn3ECL10a (yellow; BG-JF646) with YFP-Cldn10aΔPDZ/-Cldn3ECL10a (magenta; α-GFP 2nd-AF594) in COS-7 cells. Pearson analysis of co-expressed combinations from (b). Data represent the mean ± SD. Every n represents the Pearson of one TJ-like meshwork. n(Cldn2ΔPDZ + Cldn10aΔPDZ) = 15; n(Cldn2ΔPDZ + Cldn10aΔCT) = 15; n(Cldn2ΔPDZ + Cldn10aΔCCT) = 15; n(Cldn2ΔPDZ + Cldn10aΔCCT + Cldn3ΔPDZ) = 16; n(Cldn2ΔPDZ + Cldn10aΔCCT + Cldn3ΔPDZ + Cldn15ΔPDZ) = 16. Data represent the mean ± SD. Every n represents the Pearson of one TJ-like meshwork. n(Cldn2ΔPDZ + Cldn10aΔPDZ) = 15; n(Cldn2ΔPDZ + Cldn10aΔCT) = 15; n(Cldn2ΔPDZ + Cldn10aΔCCT) = 15; n(Cldn2ΔPDZ + Cldn10aΔCCT + Cldn3ΔPDZ) = 16; n(Cldn2ΔPDZ + Cldn10aΔCCT + Cldn3ΔPDZ + Cldn15ΔPDZ) = 16. Data represent the mean ± SD. Every n represents the Pearson of one TJ-like meshwork. n(Cldn2ΔPDZ + Cldn10aΔPDZ) = 15; n(Cldn2ΔPDZ + Cldn10aΔCT) = 15; n(Cldn2ΔPDZ + Cldn10aΔCCT) = 15; n(Cldn2ΔPDZ + Cldn10aΔCCT + Cldn3ΔPDZ) = 16; n(Cldn2ΔPDZ + Cldn10aΔCCT + Cldn3ΔPDZ + Cldn15ΔPDZ) = 16.
Fig. 6 Segregation enables paracellular ion flux over the TJ meshwork.

a Representative confocal images of MDCKII QKO cells stably expressing (sT) FLAG-Cldn2, FLAG-Cldn10a, and FLAG-Cldn2+FLAG-Cldn10a. Single claudin expressing cells were immunostained with anti-Cldn2 (magenta; 2nd-Atto647N) or anti-Cldn10 (magenta; 2nd-Atto647N) and anti-ZO-1 (yellow; 2nd-AF594) antibody. Double claudin expressing cells were immunostained with anti-Cldn2 (magenta; 2nd-Atto647N) and anti-Cldn10 (yellow; 2nd-AF594). The white rectangle indicates the region of interest for the magnification in (b).

b Magnification of FLAG-Cldn2+FLAG-Cldn10a expressing cells from (a). White arrows point out differently sized TJ parts that contain only Cldn10a.

c Representative confocal images of MDCKII QKO sT FLAG-Cldn3, FLAG-Cldn15 and FLAG-Cldn3+FLAG-Cldn15. Single and double claudin expressing cells were stained with anti-Cldn3 (magenta; 2nd-Atto647N) or anti-Cldn15 (magenta; 2nd-Atto647N) and anti-ZO-1 (yellow; 2nd-AF594) antibody.

d Summary of the TER (ohm·cm²), fluorescein flux (10⁻⁶ cm/s) and PNa/PCl ratio from the measured cell lines: MDCKII, MDCKII QKO and MDCKII QKO sT FLAG-tagged Cldn2, Cldn3, Cldn10a, Cldn15 and MDCKII QKO sT FLAG-tagged Cldn2 + Cldn10a and FLAG-tagged Cldn3 + Cldn15. Data represent the mean ± SD from 3 to 5 independent experiments. Single data points are shown in Supplementary Fig. 12.

e Absolute permeability for sodium (PNa; magenta bars) and chloride (PCl; cyan bars) of MDCKII, MDCKII QKO and MDCKII QKO sT FLAG-tagged Cldn2, Cldn3, Cldn10a, Cldn15, and MDCKII QKO sT FLAG-tagged Cldn2 + Cldn10a and FLAG-tagged Cldn3 + Cldn15. Data represent the mean ± SD. For the electrophysiological measurements every n represents one transwell filter: n(MDCKII) = 14; n(MDCKII QKO) = 14; n(FLAG-Cldn2) = 9; n(FLAG-Cldn10a) = 11; n(FLAG-Cldn2+FLAG-Cldn10a) = 8; n(FLAG-Cldn3) = 9; n(FLAG-Cldn15) = 12; n(FLAG-Cldn3+FLAG-Cldn15) = 9; from 3–5 independent experiments; one-way ANOVA with Kruskal-Wallis test; **P ≤ 0.01, ***P ≤ 0.001, ns (not significant). The total claudin expression level was determined by immunoblotting for Cldn2, Cldn3, Cldn10, and Cldn15 (20–25 kDa). β-actin (40 kDa) served as control. All representative images derive from 3 independent experiments. Scale bars, 10 μm (a, c), 2 μm (b). Source data are provided as a Source data file.
pathogens and their toxins. How TJs and their proteins coordinate this interplay at the nanoscale level has remained unknown since their initial discovery.12,24,29,60 Using super-resolution STED microscopy, we overcame the diffraction limit of fluorescence microscopy and the limitations of electron microscopy for the visualization of the molecular nanoscale composition of TJ in living or fixed cells and in different tissues (Figs. 1–3). Here we systematically show which of the mammalian claudins are able to form TJ-like meshworks (Supplementary Fig. 2, 3). The ability observed to form TJ strands grossly (but not strictly) parallels the previous classification into classic versus non-classic claudins based on sequence homology.21 TJ-like meshworks were formed by 11 out of 14 classic claudins but only two (Cldn11 and Cldn20) of the non-classic claudins formed a TJ-like meshwork. Notably, multiple claudins previously proposed to act as paracellular ion channels, e.g., Cldn461, 84, 16, 16.27, and 177 were unable to form strands and may require co-assembly with meshwork- or other non-meshwork-forming claudins, as observed for Cldn16 and Cldn197 or Cldn4 and Cldn8.7. Albeit a uniform distribution of Cldn4 and Cldn8 was observed here in fixed and living cells further analyses are required to determine the precise nanoscale architecture and organization of non-meshwork forming claudins in living cells. In the future, classification of nanoscale intensity features of non-meshwork forming claudins in living cells could reveal new polymerization principles, the interplay of cis/trans interactions and general structure-function relationships for these incompletely characterized claudin family members.

A different clustering analysis approach based on clustering of intensity texture as well as meshwork morphology features identified three distinct groups of meshwork forming claudins (Fig. 2). Class A and B showed distinct mesh sizes and contained both channel and barrier claudins. Class C comprised exclusively strongly sealing claudins (Cldn1, Cldn6, Cldn9, and Cldn11) and leads us to propose that dense TJ meshworks with parallel strands, as found solely in this class, perform sealing functions. Overall, our analysis of claudin meshworks suggests that there is no strict correlation between meshwork organization and channel or barrier function of individual claudins, confirming earlier work using FFEM on a limited set of claudins.7 However, it is important to note that unsupervised machine learning can only produce classification hypotheses that then need to be tested by further experimental analyses, e.g., by super-resolution imaging, genetic manipulations or paracellular flux analysis.

Since TJs are formed by multiple claudins, we combined multi-color STED microscopy with colocalisation analysis and FRET to investigate claudin interactions on a nanoscale. We found that barrier-forming claudins can intermix and that channel-forming claudins segregate from barrier-forming claudins and other channel-forming claudins when overexpressed in non-polarized cells, epithelial cell lines, and at endogenous level in responsive tissues (Figs. 3, 4). Individually and co-expressed barrier claudins Cldn1 and Cldn3 appeared as homogenous strands in FFEM, produce high colocalisation signals in STED images and high FRET values consistent with a highly intermixed organization of this and maybe other barrier claudin combinations. Conversely, very small clusters were found in FFEM images of recombinant Cldn2 combined with Cldn1 or Cldn3 and endogenous Cldn2 but their molecular identity was so far unknown. Cluster from Cldn2 FFEM images could reflect differential association of Cldn2 with the protoplasmic and extracellular FFEM surfaces or alternatively a segregation pattern of Cldn2 from Cldn1 or Cldn3. Trans interactions reported between Cldn2 and Cldn3, but not between Cldn2 and Cldn113 are mirrored by higher STED colocalisation of Cldn2/3 compared to Cldn2/1. We speculate that stable trans interactions between Cldn2 and Cldn3 might lead to very short Cldn2 strands that cannot be resolved by STED microscopy. Such short strands may underlie the high degree of colocalisation between Cldn3 and Cldn2 compared to that of Cldn3 with other channel-forming claudins, e.g., Cldn10a, Cldn10b, or Cldn15. Therefore the balance of cis and trans interactions can fine-tune the segregation properties of claudins as also observed by exchanging extracellular loops of Cldn2 and Cldn10a (Fig. 5h, i).

Neither Cldn2 channel inactivation by MTSET, cholesterol depletion, nor the deletion of the entire claudin C-terminus or the PDZ motif required for phosphorylation or ZO-1 binding respectively affected the nanoscale segregation. Unexpectedly, we observed smaller, unstructured, and less frequent meshworks formed by combinations of C-terminal or PDZ deletion mutants of Cldn2/10a in COS-7 cells. Conversely, FFEM and live imaging data of C-terminal or PDZ deletion mutants of Cldn1, Cldn2, or Cldn3 expressed in non-polarized fibroblasts were reported to lead to unperturbed strand and meshwork morphologies, albeit showing higher meshwork dynamics.23,24.7 We therefore speculate that segregated meshworks could be more dynamic and unstable, especially at segregation break points with possibly imperfect interaction interfaces. Here ZO-1 interactions could be required to stabilize segregated meshworks. Future work based on live-cell super-resolution microscopy could characterize nanoscale strand dynamics, strand breaks, and molecular factors affecting the barrier properties of claudin homo- and heteropolymers in TJs.

The extracellular loops of channel claudins form not only the paracellular ion pore through structural organization, but together with transmembrane helices and other interacting surfaces contribute to strand formation and segregation. Molecular modeling based on the crystal structure of Cldn157.68 and experimental data7 indicate that multiple claudin molecules interacting in cis and trans are implicated in pore formation7.23,24.66. The segregation mechanism described here could ensure that these pores are not disturbed by residues of ECLs from nearby barrier- or channel-forming claudins. The maximum length of a segregated strand on the other hand must be smaller than the total length of a typical mesh to increase the probability of integration of a selective ion pore that spans all meshwork strands (Supplementary Fig. 13). Finally, discontinuities in the barrier that lead to non-specific leaks must be prevented when TJ strands consisting of claudins with different permeability properties are formed. These non-specific leaks could be sealed by close membrane contacts formed at strand boundaries or by a so far undefined transmembrane protein enriched at TJs.

The integration of non-meshwork forming Cldn16 into Cldn19 strands or the induction of strands by copolymers of the two non-meshwork forming Cldn4 and Cldn8 were not distinguishable from colocalisation values of intermixed barrier strands by our STED microscope (Fig. 3; Supplementary Figs. 5 and 7), so could enable the formation of ion pores at so far unresolved interfaces that are different from the paracellular pore structures proposed for Cldn15 polymers7.68.64. The combination of integration, segregation, and exclusion observed for Cldn10b, Cldn16, and Cldn19 is in accord with differential interactions of these claudins7 and could lead to the observed mosaic spatial separation of paracellular Na+/ Mg2+ (Cldn3/16/19) transport found in cell–cell-contact areas of the thick ascending limb of Henle’s loop of the nephron7.

Because nanoscale segregation of barrier and channel claudins could ensure specific claudin polymerization and meshwork incorporation, as described above, and was observed in epithelial cells and in two resorbing tissues, it may be important for paracellular transport processes. To measure the barrier and channel properties of the segregated claudin combinations (Cldn2/Cldn10a and Cldn3/Cldn15) and their individual claudin components (Cldn2, Cldn3, Cldn10a, and Cldn15), claudins were reconstituted in MDCKII QKO cells, an epithelial cell line lacking five major claudins and thus the TJ meshwork. Of note, very low levels of non-meshwork forming Cldn12 and Cldn16 were expressed in MDCKII QKO (Supplementary Fig. 9a), but these failed to colocalize with occludin or ZO-1 at the plasma membrane (Supplementary Fig. 9b) and did not copolymerize with Cldn2, Cldn3,
Cldn10a, and Cldn15 (Supplementary Fig. 10). The apparent absence of both claudins from the TJ could be explained by their inability to form strands and/or the absence of copolymerizing claudins such as Cldn19, as previously observed for Cldn16.

Reconstitution of Cldn3 in QKO showed that a single barrier claudin can form an efficient paracellular seal. When Cldn2 or Cldn15 were reconstituted in QKO cells, a paracellular barrier containing Na+/Cl−selective channels was formed with a high Na+/Cl− permeability that closely match predictions form molecular models of Cldn15 homopolymers. Conversely, reconstitution of Cldn10a created barrier containing a Cl−selective channels consistent with the results from former studies. Albeit based on a limited set of claudins we noted that barrier claudins show significantly lower permeability to the small molecule fluorescein than all channel-forming claudins tested. This could be a consequence of more complex meshworks and/or fewer strand breaks occurring in restituted barrier claudins and will be subject of further investigation.

Since single barrier or channel claudins can form tight or ion selective TJ strands in QKO cells, we conclude that segregation interfaces between different claudin segments do not form specific ion pores (Fig. 6; Supplementary Fig. 13). Coexpression of segregating Cldn2 and Cldn10a produces parallel Na+/Cl− system in MDCKI cells from Curry et al. in 2020 and ion absorption consistent with the results of an established Cldn2/Cldn10a Tet On/Off consistent with the predicted Na+/Cl− fluxes (Fig. 6; Supplementary Fig. 13). Coexpression of segregating barrier claudins show significantly lower permeability to the small molecule fluorescein than all channel-forming claudins tested. This could be a consequence of more complex meshworks and/or fewer strand breaks occurring in restituted barrier claudins and will be subject of further investigation. We propose that segregation enables balanced, interspaced, and alternating incorporation of claudins with different permeability properties into common TJ meshworks. This can produce parallel fluxes of different ions or increase the charge selectivity of paracellular ion channels at the TJ. Thereby, the segregation/intermixing pattern of the claudins expressed critically contributes to regulation of paracellular permeability in a given tissue. Our model (Supplementary Fig. 13) that is based on predictions from Claude et al. and Weber et al. assures that sufficiently large pore stretches enable a balanced flux of oppositely charged ions, by having minimal occurrence of dead ends for ions, while ensuring paracellular barrier integrity.

Methods

Plasmid cloning

All used constructs are listed in Supplementary Dataset 1. For all standard cloning and sub-cloning approaches self-made chemically competent TOP10 were used. Commercial NEB 5-alpha chemically competent E. coli (New England Biolabs Inc., #C2987H) were used for site-directed mutagenesis and HiFi-assembly cloning. HB101 (recA) (Promega GmbH, #L201) bacteria were used for cloning and amplifying pLIB plasmid DNA. Standard PCR for amplification was performed using the Phusion High-Fidelity DNA Polymerase (Thermo Fisher Scientific, #F530S) according to the manufacturer’s instructions. All used primers in this study were synthesized by BioTeZ Berlin. All used constructs are listed in Supplementary Dataset 1. All human claudin (Cldn) constructs are labeled with "hu" (huCldn). All murine Cldn constructs are labeled with an "m" (mCldn). For the following constructs pYFP-Cldns, pTrq2-Cldns and pGFP-Cldns an identical linker ("SLVSSPD" (8 AA)) between fluorescent tag and Cldn sequence was used. The linker for pSNAP-Cldns and pHalo-Cldns contains three additional AA ("LYKSLVSSPD" (11 AA)).

The pYFP-Cldn constructs (pYFP-huCldn1, -2, -3, -4, -8, -10a, -10b, -11, -12, -15, -16, -19a, -19b, 20, 22, 23), pmCldn3-YFP, phuCldn3-YFP, phuCldn1-YFP as well as pTOPO-huCldn6 and pTOPO-huCldn9 were provided by Susanne Krug, Dorothée Günzel and Jörg Piontek (Clinical Medicine, Charité – Universitätsmedizin Berlin).

The pYFP-Cldn constructs (pYFP-huCldn5, -7, -12, -20, -24, -25 and pYFP-mCldn13, -14, -18.1, -18.2) and pCFP-huCldn17 were provided by Lorena Suarez from the Mertins Lab (Max-Delbrück-Center for Molecular Medicine (MDC), Berlin).

pHuCldn3-C1 was generated by sub-cloning Cldn3 from pYFP-huCldn3 via PCR and restriction enzymes HindIII and Sall into p-C1 (generated from a pGFP-C1).

pYFP-huCldn17 was generated by sub-cloning Cldn17 from pCFP-huCldn17 via restriction enzymes BsrGI and Sall into pYFP-huCldn15.

pYFP-huCldn6 and pYFP-huCldn9 were generated by sub-cloning Cldn6 and Cldn9 from pTOPO-huCldn6 and pTOPO-huCldn9 via PCR and restriction enzymes BsrGI and EcoRV into pYFP-huCldn15.

pGFPCldn-C1 constructs (pGFPCldn2ΔmCC and pGFPCldn2ΔEC and pGFPCldn26–27) were de novo synthesized and obtained from General Biosystems (Chuzhou, Anhui, China).

pHaloCldn constructs (pHalo-huCldn10a, -15, -16, -19a, -19b, -20, -22) were generated by sub-cloning Cldns from pYFP-Cldns via PCR and restriction enzymes BshTI and BsrGI into pmCherry-C1.

pSNAP-Cldn constructs (pSNAP-huCldn1, -2, -3, -4, -8, -10a, -11, -15, -16, -19a, -22) were generated by sub-cloning Cldns from pYFP-Cldns via PCR and the restriction enzymes BshTI and BsrGI into pmCherry-C1.

pSNAP-huCldn10a, -15, -16, -19a, -19b, -20, -22 via PCR and the restriction enzymes BshTI and BsrGI into pSNAP-C1.

pSNAP-huCldn12 construct was de novo synthesized and obtained from General Biosystems (Chuzhou, Anhui, China).

pHaloCldn constructs (pHalo-huCldn10a, -15, -16) were generated by sub-cloning Cldns from pYFP-Cldns via PCR and the restriction enzymes BsrGI and Sall into pHalo-C1.

pTrq2-Cldn constructs (pTrq2-huCldn2, -3) were generated by sub-cloning huCldn2 and huCldn3 from pSNAP-huCldn2 and pSNAP-huCldn3 via restriction enzymes BsrGI and Sall into Trq2-C1.

pSNAP-Cldn PDZ binding motif deletion mutants (pSNAP-huCldn2ΔCT, -10a ΔPDZ, -15 ΔCT) were generated by site-directed mutagenesis PCR and following phosphorylation of the PCR product with a T4 Polynucleotide Kinase (Thermo Scientific, #EK0032) using pSNAP-huCldn2, -3, -10a, -15. For all constructs the last three amino acids of the Cldn sequence were deleted: pSNAP-huCldn2ΔCT for...
In the first approach pSNAP-huCldn2ECL2C10a was generated by using pSNAP-huCldn2 as backbone and pSNAP-Cldn10a for generating ECL1C10a with Cldn2 homology arms. The ECL2 “WNLHGILRDFYSPLVPDSMKFEI” (amino acids 138–162) of huCldn2 was used as backbone plasmid and pSNAP-huCldn10a for generating ECL2C10a with huCldn2 homology arms to sub-clone pSNAP-huCldn2ECL2C10a. The ECL2 “WNLHGILRDFYSPLVPDSMKFEI” (amino acids 138–162) of huCldn2 was substituted by the ECL2 “LYANKITTEFFDPLFVEQKYELGA” (amino acids 135–158) of Cldn10a. pTrq2-huCldn2ECL2C10a was generated via sub-cloning of huCldn2ECL2C10a from pSNAP-huCldn2ECL2C10a using restriction enzymes BsrGI and Sall into pTrq2-CL.

pLJ-CMV-FLAG-huCldn2-10a-15-Puro constructs were generated by sub-cloning Cldns from pSNAP-Cldn constructs (pSNAP-huCldn2-10a, 15-Puro) via PCR and the restriction enzymes BshTI and NotI into pLJ-CMV-GFP-N1-Puro. 

Retroviral generation of stable cell lines
HEK293T cells were stably transfected with a lentiviral construct carrying a puromycin resistance cassette under the control of a CMV promoter. The lentiviral construct was generated by co-transfecting HEK293T cells with the pLJ-CMV-GFP-N1 construct and a packaging plasmid pMD2.G, along with a VSV-G pseudotype and the Rev expression plasmid psiPAX2. After 24 h, supernatants containing concentrated virus were collected and stored at –80°C. To infect the cells, a 1:1 mix of concentrated virus and a 10:1 mix of DMEM was used to infect the cells. GFP-N1 transducing virus was used to infect the cells.

Mammalian cell culture
The following cell lines were used in the study. COS-7 (ATCC CRL 1651), HEK293 (ATCC CRL 1573), HEK293T (ATCC CRL 3216), 3T3 fibroblasts (DSMZ, no.: ACC 173), MDCKII (ECACC 00062170), and Caco-2 cells (ATCC HTB 37) were taken from own lab cell culture stocks. MDCKC7 cells were kindly provided by Lorena Suarez Artiles from the Mertins Lab (Max-Delbrück-Center for Molecular Medicine (MDC), Berlin). MDCKII quintuple claudin knock-out (QKO) cells were kindly provided by the lab of Miklo Furuse. MDCKII QKO cells stably expressing claudins were generated in this study. COS-7, HEK293, HEK293T, 3T3 fibroblasts, MDCKII, MDCKII QKO, and MDCKC7 were cultured in DMEM with high glucose (Gibco, Thermo Fisher Scientific, #11965084) supplemented with 10% FBS (Gibco, Thermo Fisher Scientific, #10082147) and 50 μg/ml of penicillin-streptomycin (Pen-Strep) (Gibco, Thermo Fisher Scientific, #15140122) at 37°C and 5% CO2. MDCKQII QKO cells stably expressing claudins were cultured as described above and under additional selection pressure of 2–5 μg/ml of Puromycin (Invivogen, #ant-pr-1) and 300–500 μg/ml of G418 (Genetix, #G418) for 2 weeks. Caco-2 were cultured in MEM (Gibco, Thermo Fisher Scientific, #11095080) containing 15% FBS and 50 μg/ml Pen-Strep at 37°C and 5% CO2. PBS (Gibco, Thermo Fisher Scientific, #14190144) was used for all washing steps. For transfection, cells were seeded on 18 mm or 25 mm glass coverslips (#1.5H) (Thermo Fisher Scientific) or in µm-Slide 8-well glass bottom dishes (Ibidi, #80827) coated with 2% tritratigel in DMEM in a confluent of 70%. After 24 h the transfection was performed with Lipofectamine 2000 Transfection Reagent (Thermo Fisher Scientific, #11668019) according to the manufacturer’s instructions. Snap-tag or Halo-tag labeling was performed 24 h after transfection. The transfected cells were incubated for 1 h at 37°C and 5% CO2 with 2 μM BG-Atto590 (SNAP-tag ligand) or BG-JF646 (SNAP-tag ligand) and 1 μM CA-Atto590 (Halo-tag ligand) or CA-JF646 (Halo-tag ligand) in DMEM, intensively washed and post-incubated in DMEM for another 30 min at 37°C and 5% CO2. For electrophysiological measurements, MDCKII, MDCKII QKO, and MDCKQII QKO cells stably expressing claudins were seeded in a cell density of 1.5 × 10^5 cells on 0.6 cm² PCF transwell coverslips (#1.5H) (Fisher, Quality One) for 7 days at 37°C and 5% CO2.

Retrovirus production via calcium phosphate transfection of HEK293T cells
HEK293T cells were seeded in 6-well plates or in 10 cm petri dishes and were transfected with plasmid mix of packaging plasmid DNA pCIG3.NB, lentiviral envelope plasmid pMD2.G, and genomic plasmid DNA (pLJ-CMV-FLAG-Cldn2-10a-15-Puro) according to the manufacturer’s instructions. Snap-tag or Halo-tag labeling was performed 24 h after transfection. The transfected cells were incubated for 1 h at 37°C and 5% CO2 with 2 μM BG-Atto590 (SNAP-tag ligand) or BG-JF646 (SNAP-tag ligand) and 1 μM CA-Atto590 (Halo-tag ligand) or CA-JF646 (Halo-tag ligand) in DMEM, intensively washed and post-incubated in DMEM for another 30 min at 37°C and 5% CO2. For electrophysiological measurements, MDCKII, MDCKII QKO, and MDCKQII QKO cells stably expressing claudins were seeded in a cell density of 1.5 × 10^5 cells on 0.6 cm² PCF transwell filter with a pore size of 0.4 μm (Fisher Scientific, Millipore Millicell inserts, #10126240) and cultivated for 5–7 days at 37°C and 5% CO2.
selection of successfully infected cells and non-infected cells, up to 10 µg/ml Puromycin and 800 µg/ml G418 were used. The growth medium was changed to DMEM with 10% FBS but without Pen/Strep when G418 was used. The cells were routinely checked for viability and expression level for about one week. After one week, the selection pressure was reduced to 2–5 µg/ml Puromycin and 300–500 µg/ml G418. The selection process was continuously checked and the expression of the claudins was controlled by immunofluorescent staining and immunoblotting.

Antibodies

All used antibodies and their dilutions are listed in Supplementary Dataset 1.

Antibody fluorophore conjugation

Donkey anti-rabbit Atto542, donkey anti-mouse Atto542, and rabbit anti-Cldn3 Atto590 were produced by incubating 100 µl donkey anti-rabbit (AffiniPure Donkey Anti-Mouse IgG (H + L) from Jackson Immuno Research Ltd., #711-005-152) and 100 µl donkey anti-mouse (AffiniPure Donkey Anti-Mouse IgG (H + L) from Jackson Immuno Research Ltd., #715-005-151) with a 5–10x excess of Atto542-NHS (AttoTEC, #AD542-31) and rabbit anti-Cldn3 (Thermo Fisher Scientific, #34-1700) with a 5–10x excess of Atto590-NHS (Sigma-Aldrich, #79636-SMG-F) for 1 h at RT under constant agitation. The labeled antibody was purified by using Zeba™ Spin Desalting Columns (7 K MWCO, 0.5 ml) (Thermo Fisher Scientific, #89883) according to the manufacturer’s instructions. The degree of antibody labeling was determined by absorbance measurements via NanoDrop (NanoDrop ND-1000, Thermo Fisher Scientific).

Immunocytochemistry

On glass coverslips (#1.5) seeded and transfected COS-7, 3T3 fibroblasts, MDCKC7 and Caco-2 were washed with PBS supplemented with 0.5 mM magnesium and 1 mM calcium (PBS), fixed with 37 °C pre-warmed 4% PFA/sucrose for 10 min, permeabilized with 0.2% Triton X-100 in PBS for 5 min and blocked in blocking solution containing 10% NGS, 1% BSA, 0.05% Tween-20 dissolved in PBS. For YFP-booster, cells were incubated with anti-GFP Atto647N nanobody (α-GFP-NB-Atto647N; Chromotek, #gba647n100, 1:200), anti-GFP Atto594 nanobody (α-GFP-NB-Atto594; Chromotek, #gba594100, 1:200) or anti-GFP Atto488 nanobody (α-GFP-NB-Atto488; Chromotek, #gba488100, 1:200) for 1 h at room temperature or with a mouse anti-GFP antibody (Thermo Fisher Scientific, #A-11201, 1:500) in blocking solution overnight at 4 °C. MDCKII, MDCKII QKO and MDCKII QKO stably expressing FLAG-tagged claudins were seeded on transwell filters were cultivated for 5–7 days, washed with PBS, fixed with ice-cold ethanol at −20 °C for 20 mins and blocked in blocking solution containing 10% NGS, 1% BSA, 0.05% Tween-20 dissolved in PBS. All used primary antibodies, rabbit anti-ZO-1 (Thermo Fisher Scientific, #61-7300), mouse anti-ZO-1 (Thermo Fisher Scientific, #33-9100), mouse anti-Occludin (Thermo Fisher Scientific, #33-1500), rabbit anti-Calreticulin (Abcam, #ab9526), mouse anti-Cldn2 (Thermo Fisher Scientific, #32-5600), rabbit anti-Cldn3 (Thermo Fisher Scientific, #34-1700), rabbit anti-Cldn10 (antibodies-online.de, ABLIN389395), rabbit anti-Cldn12 (BBI America, #118801), rabbit anti-Cldn15 (Thermo Fisher Scientific, #38-9200), mouse anti-Cldn16 (gift from Prof. Henrik Dimke) were incubated in blocking solution overnight at 4 °C. For secondary antibody labeling the following antibodies were incubated in blocking solution for 1 h at room temperature: donkey anti-mouse Atto542 (Atto542-NHS from AttoTEC, #AD542-31 and AffiniPure Donkey Anti-Mouse IgG (H + L) from Jackson Immuno Research Ltd., #715-005-151, 1:200), goat anti-mouse Alexa Fluor Plus 594 (Thermo Fisher Scientific, #A32740, 1:200) for 1 h at room temperature or with a mouse anti-Cldn16 (Thermo Fisher Scientific, #38-9200, 1:200) overnight at 4 °C. For secondary labeling goat anti-rabbit Alexa Fluor Plus 594 (Thermo Fisher Scientific, #A32740, 1:200) for 1 h at room temperature followed by intensively washing and an over blocking step using donkey anti-rabbit (Jackson Immuno Research Ltd., #711-005-152, 1:200) in a 10x excess for 2 h at room temperature. The samples were washed intensively with PBS. As control the over blocked slices were incubated with a goat anti-rabbit Alexa Fluor 488 (Thermo Fisher Scientific, #A-11008, 1:200) for 1 h at room temperature. The samples were washed intensively with PBS and incubated with rabbit anti-Cldn15 (Thermo Fisher Scientific, #38-9200, 1:100) overnight at 4 °C. For secondary labeling goat anti-rabbit Atto647N (Active Motif, #15048, 1:200) for 1 h at room temperature used. The tissue was washed and mounted using Pro Long Gold Antifade (Thermo Fisher Scientific, #P36934) on object slides and incubated for 24 h at room temperature. For every sample, the control staining with Alexa Fluor 488 was imaged to verify the quality of the over blocking step.

Immunohistochemistry of single murine tubules from proximal tubule

C57BL/6j mice were sacrificed by cervical dislocation. The Landesamt für Gesundheit und Soziales (LAGeSo) Berlin with their permission under the license T-025/16 approved the use of animals for this study. The whole duodenum was taken out, washed with a Ringer solution (7.2 g NaCl, 0.17 g CaCl2, 0.37 g KCl) and dissolved in reagent-grade H2O, cut in lateral direction into multiple 1–2 cm parts, embedded in Tissue-Tek (Sakura Finetek, Fisher Scientific, #12351753) and immediately frozen in liquid nitrogen. 10 µm thick longitudinal slices were cut with a cryostat (Microm HM 560 Cryostat, Thermo Fisher Scientific) and placed on acid cleaned and organosilan coated glass coverslips (#1.5H). The tissue sections were fixed with −20 °C cold ethanol for 20 min, washed with PBS, blocked with PBS containing 6% NGS, 1% BSA and 0.05% Tween-20 for 1 h at room temperature and stained with rabbit anti-Cldn3 (Thermo Fisher Scientific, #34-1700, 1:100) overnight at 4 °C. After 24 h the slices were incubated with goat anti-rabbit Alexa Fluor Plus 594 (Thermo Fisher Scientific, #A32740, 1:200) for 1 h at room temperature followed by intensively washing and an over blocking step using donkey anti-rabbit (Jackson Immuno Research Ltd., #711-005-152, 1:200) in a 10x excess for 2 h at room temperature. The samples were washed intensively with PBS. As control the over blocked slices were incubated with a goat anti-rabbit Alexa Fluor 488 (Thermo Fisher Scientific, #A-11008, 1:200) for 1 h at room temperature. The samples were washed intensively with PBS and incubated with rabbit anti-Cldn15 (Thermo Fisher Scientific, #38-9200, 1:100) overnight at 4 °C. For secondary labeling goat anti-rabbit Atto647N (Active Motif, #15048, 1:200) for 1 h at room temperature used. The tissue was washed and mounted using Pro Long Gold Antifade (Thermo Fisher Scientific, #P36934) on object slides and incubated for 24 h at room temperature. For every sample, the control staining with Alexa Fluor 488 was imaged to verify the quality of the over blocking step.
subsequently vigorously removed from the slide under visual control and washing with PBS containing 0.3% Triton X-100 (PBS-T100). Primary antibodies (either mouse anti-Clnd2 (Thermo Fisher Scientific, #32-560), rabbit anti-Clnd10 (Antibodies-online, #ABIN383935), or rabbit anti-Clnd2 (Thermo Fisher Scientific, #51-6100) and mouse anti-Clnd10 (Thermo Fisher Scientific, #41-S100), 50 µl/slide) were incubated at 4 °C overnight. After extensive washing under visual control (PBS-T100), secondary antibodies goat anti-mouse Alexa Fluor Plus 594 (Thermo Fisher Scientific, #A32744, 1:200) and goat anti-rabbit Attos647N (AttoTEC, #15048, 1:200) were incubated for 1–2 h at room temperature. After final vigorous washing with PBS-T100, embedding was performed with Pro Long Gold Antifade (Thermo Fisher Scientific, #P36934) together with #1 LSH coverslips.

Cell lysates
COS-7 were seeded in 6-well plates or 10 cm petri dishes and after 24 h incubation scraped with 100–500 µl lysis buffer (1% Triton X-100, 20 mM HEPES, pH 7.4, 130 mM NaCl, 10 mM NaF, 0.03% PIC) on ice. MDCKC7, Caco-2, MDCKII QKO, and MDCKII QKO FLAG-tagged or untagged Cldns expressing cells were seeded in 6-well plates and after 5–7 days scraped with 100 µl lysis buffer (1% Triton X-100, 20 mM HEPES, pH 7.4, 130 mM NaCl, 10 mM NaF, 0.03% PIC) on ice. The supernatants were transferred into pre-cooled 1.5 ml tubes and incubated for 30 min on ice. The protein-containing supernatant was isolated via centrifugation at 17,000 g for 20 min at 4 °C. 1–10 µl of the supernatant was incubated with 490–499 µl H2O and 500 µl 2x Bradford reagent for 5 min and the protein concentration was determined by measuring the OD595 with a photometer (BioPhotometer plus, Eppendorf). The protein lysates were denaturized with 6x Laemmli buffer (0.375 M Tris pH 6.8, 12% SDS, 60% glycerol, 0.6 M DTT, 0.06% bromophenol blue) for 5 min at 95 °C and stored at −20 °C.

Immunoblot-based analysis
For protein separation via SDS-PAGE (sodium dodecylsulfate polyacrylamide gel electrophoresis) 15–30 µg of lysate was loaded on a 10–12% polyacrylamide gel or 4–15% MINI-PROTEAN TGX Prestain Gels (Bio-Rad, #4561083/86). The PageRuler Plus Prestained (Thermo Fisher Scientific, #26620) or PageRuler Prestained (Thermo Fisher Scientific, #26616) were used as protein ladder. The gel was run in SDS-running buffer (25 mM Tris base, glycine 0.192 M, SDS 0.1%) at 120 Volt for 60–90 min using the Mini-PROTEAN Tetra Vertical Electrophoresis Cell (Bio-Rad, #1668004). The protein transfer on a nitrocellulose membrane was done by Wet Blot in a 20% methanol containing 15–20 °C. The protein-containing supernatant was incubated with Alexa Fluor Plus 594 (Ex.: 561 nm; EmF.: 660–630 nm), Atto647N (Ex.: 633 nm; EmF.: 636–740 nm). Images were acquired with a PL APO DIC M27 63×/1.40 NA oil objective (Carl Zeiss Microscopy).

For the detection in the LSM780, a photomultiplier was used. The LSM780 was controlled by the Zeiss ZEN2010 software (Carl Zeiss Microscopy). Single- and multi-color confocal imaging of fixed samples was performed in sequential mode with the following fluorophore-specific excitation (Ex.) and emission filter (EmF.) settings: Alexa Fluor Plus 594 (Ex.: 561 nm; EmF.: 660–630 nm), Atto647N (Ex.: 633 nm; EmF.: 636–740 nm). Images were acquired with a HC PL APO CS2 100×/1.40 NA objective (Leica Microsystems).

Time-gated single- and multi-color STED imaging
STED images were taken with a Leica SP8 TCS STED microscope (Leica Microsystems) equipped with a pulsed white-light excitation laser (WLL; ~80 ps pulse width, 80 MHz repetition rate (NKT Photonics)) and two STED laser for depletion at 592 nm and 775 nm. The microscope was housed in a heatable incubation chamber (ILS Life Imaging Services). The system was controlled by the Leica LAS X software. Single- and multi-color STED imaging of fixed samples was performed in sequential mode with the following fluorophore-specific excitation (Ex.) and emission filter (EmF.) settings: Alexa Fluor 488 (Ex.: 488 nm; EmF.: 519–586 nm), YFP (Ex.: 514 nm; EmF.: 524–568 nm), Alexa Fluor Plus 594 (Ex.: 590 nm; EmF.: 600–640 nm), Atto647N (Ex.: 640/650 nm; EmF.: 650/670–700 nm). Images were acquired with a HC PL APO CS2 100×/1.40 NA oil objective (Leica Microsystems).

One- and two-color live STED imaging
Live STED imaging was performed with COS-7 seeded on 25 µm glass coverslips (#1.5H) using an Attofluor cell chamber (Thermo Fisher Scientific, #A7816) or in µ-Slide 8-well glass bottom dishes (Ibidi, #80827). The glass surface was coated with 2% matrigel. The cells were transfected with SNAP-tag or Halo-tag constructs. A HEPES buffered live imaging solution (Thermo Fisher Scientific,
#A14291DJ) was used. 24 h prior imaging, the heatable incubation chamber was set to 37 °C to provide focus stable imaging. For single-color live STED imaging JF646 (Ex.: 640 nm; Em.: 650–700 nm) was used. The imaging settings were the same as described above (see the “Time-gated single- and multi-color STED imaging” section) except for the reduced scanning format of $12 \times 12$, increased 16–32x line averaging and a 12x optical zoom yielding in a pixel size of $18.9 \times 18.9$ nm. The acquisition time per frame for serial imaging was set to 10 sec/frame.

Two-color Live STED imaging was performed with the JF646 (Ex.: 640 nm; Em.: 650–700 nm) as SNAP-ligand, Atto590 (Ex.: 590 nm; Em.: 600–640 nm) as Halo ligand or YFP. For the combination of $p$SNAP-Cldn3 with $p$YFP-Cldn4 and $p$SNAP-Cldn8 with $p$YFP-Cldn4 the imaging settings were the same as described above (see the “Time-gated single- and multi-color STED imaging” section). For the combination $p$SNAP-Cldn2 and $p$Halo-Cldn10a a reduced scanning format of $12 \times 12$, 16x line averaging and a 12x optical zoom yielding in a pixel size of $18.9 \times 18.9$ nm was used. Imaging was performed in an acquisition speed of 10 sec/frame. For imaging of the combination of $p$SNAP-Cldn3 and $p$Halo-Cldn15 a resonance scanner at 8000 Hz was used. Single live images were taken with a scanning format $12 \times 12$, 32x line averaging and a 12x optical zoom yielding in a pixel size of $18.9 \times 18.9$ nm. Imaging was performed in an acquisition speed of 19.5 sec/frame. For further imaging processing we used nothing. We applied a deep learning-based restoration method to remove noise from images for visualization as specified in the figure legends. We trained the noise model in Fiji on a GPU with 150 epochs and 200 steps per epoch on a large set of training data with a batch size of 100 and a dimension of 180 × 60 px. The neighborhood radius was set to 5. After training the best model was chosen for predicting the image with filtered noise.

**FWHM measurements**

Full-width half-maximum (FWHM) measurements were performed with $p$SNAP-huCldn3 or $p$SNAP-huCldn11 transfected COS-7 cells labeled with BG-JF646. The same imaging settings, except for temperature and imaging solution differences (fixed: PBS at room temperature; live: live imaging solution at 37 °C), were used for live and fixed cell imaging. For Cldn3 line profiles (straight line 0.3 μm length, 10 px = 189 nm width) of 20 single strands per TJ-like meshwork (8 TJ-like meshworks in total) were analyzed for FWHM and average fluorescence intensity by Gaussian Fitting using Fiji ImageJ. For Cldn11 only line profiles of 20 strands in one TJ-like meshwork were measured. The FWHM was determined by multiplication of sigma with the factor 2.35.

**Automated TJ-like meshwork analysis**

**Cluster analysis of claudins.** A dataset of 29 claudins with in total 168 images with an average number of 5 images per claudin was used for this analysis. Analysis for texture and meshwork analysis was performed within a single 200 × 200 px crop per image. Crops were defined by an expert annotator when possible in the center of a structure, avoiding bright potential artefacts and outside edges. Images were filtered for the texture analysis using a 3 px Gaussian filter kernel and a sigma of 1. Gray level co-occurrence matrices were computed at distances 1, 3, 5, and 10 px and Haralick texture features were extracted from 4 angles and averaged using the python mahotas library. Three texture features (sum average, sum variance, and sum of squares: variance) were selected based on literature. For segmenting the meshwork, the images were filtered using Multiscale Oriented-Flux Tubularity filter implemented in the Fiji plugin, simple neurite tracer. A fixed threshold value was then applied to the filtered image and the largest connected region comprising the meshwork was kept. Within the ROI the percent of segmented area was measured. A skeleton analysis was performed to measure within the ROI the number of branches, average and maximum branch length, number of total junctions as well as triple and quadruple junctions. The binary mask was inverted to measure the mesh size within the ROI, excluding objects touching the edge. The number of meshes, average mesh size, and variance of the mesh size was measured.

For clustering the meshwork forming claudins, both the meshwork analysis and texture analysis features were used. ROIs with a segmented area of less than 10% or more than 90% were excluded as well as meshworks with less than 10 branches. The features were then averaged for each ROI. ClustVis server was used for visualizing the clustering with ln(x)-transformation applied to the values, the rows were centered, and unit variance scaling was applied. Clustering was computed using correlation distance and average linkage. Code and example images provided at [https://doi.org/10.5281/zenodo.7009994](https://doi.org/10.5281/zenodo.7009994).

**Pearson correlation analysis**

COS-7, Caco-2, or MDCK-C7 cells were seeded on with 2% matrigel-covered coverslips and transfected with equal amounts of two or three plasmids coding YFP-, SNAP- or Halo-tagged claudins. For reproducible Pearson analysis over several different experiments, the imaging parameter were set with the following positive controls set as reference: $p$SNAP-Cldn2/pYFP-Cldn2, $p$SNAP-Cldn3/pYFP-Cldn3, $p$SNAP-Cldn19a/pYFP-Cldn19a and then kept constant for the rest of an experiment. In general, the cells were labeled with the SNAP-ligand BGGatto590 and GFP-booster α-GFP-NB-Atto647N. For the $p$SNAP-Cldn2/pYFP-Cldn2, $p$SNAP-Cldn19a/pYFP-Cldn19a and $p$SNAP-Cldn15/pYFP-Cldn15 Pearson analysis the combination of SNAP-ligand BGGatto646 and mouse anti-GFP with anti-mouse Alexa Fluor Plus 594 was used. For the triple claudin expression experiments the combination of Halo-ligand CA-JF646, SNAP-ligand BGGatto590, and mouse anti-GFP with anti-mouse 2nd-Atto542 was used. For every condition at least 5 meshworks over 3 independent experiments with the same settings (1024 × 1024, 16x line averaging, pixel size of 18.9 × 18.9 nm).
18.9 × 18.9 nm) were imaged. From every meshwork a ROI with a representative signal of the transfected claudins was picked and the Pearson above threshold was measured with a Coloc2-based script in Fiji ImageJ with the PSF set to 2 px = 38.8 nm. At least >30 STED images from three different isolated proximal tubules stained with anti-Cldn2 mouse, anti-mouse Alexa Fluor Plus 594 and anti-Cldn10 rabbit, anti-rabbit Atto647N (see Immunofluorescence of single murine tubules from proximal tubule) were taken and imaged with the same settings (1024 × 1024, 16x line averaging, pixel size of 18.9 × 18.9 nm). Out of every meshwork, a ROI with a representative signal of the stained claudins was picked and the Pearson above threshold was measured with a Coloc2-based script in Fiji ImageJ with the PSF set to 2 px = 38.8 nm.

Fluorescence resonance energy transfer (FRET) measurement HEK293 cells were transfected with pTrq2- and pYFP-Cldn constructs. A plasmid DNA ratio of 1:1 was used for all pTrq2-Cldn3 approaches. For pTrq2-Cldn2 or pTrq2-Cldn2Stx the ratio was changed in all approaches to 1:5. Transfected HEK293 cells were visualized on a LSM510-NLO inverted microscope (Carl Zeiss Microscopy GmbH, Jena, Germany) using a 40x/1.3 oil objective. Trq2 fluorescence signals were recorded (IR laser, λ exc = 810 nm, two-photon technique, META detector, spectral range 436–650 nm and split using a MBS KP 700). Channel pictures were taken prior to the recording of spectra in order to estimate expression of the Trq2 and YFP-tagged constructs (Trq2: IR laser λ exc 810 nm, two-photon technique, λ em = 430–505 nm, META detector; YFP: argon laser, λ exc = 514 nm, λ em = 560 nm long pass filter). FRET data analysis was performed using the software ZEN2010 (Carl Zeiss Microscopy, Jena, Germany), and Excel2013 (Microsoft Office), respectively. For the analysis, ROIs were set on the contact side of two neighboring HEK293 cells that were expressing both claudins. The fluorescence spectra of Trq2-tagged constructs in presence and absence of the YFP-tagged constructs receptor were measured. To calculate the FRET-based fluorescence, a λ-stack with a linear spectral unmixing mode was used to correct any YFP fluorescence crosstalk into the FRET channel (523–532 nm). The λ-stack is an integral part of the confocal laser system software. YFP correction was carried out to correct for direct excitation of the acceptor during donor excitation. The effect was expressed by changes in the FRET ratio, which is calculated by dividing the acceptor emission (YFP, λem = 532 nm) by the donor emission (Trq2, λem = 478 nm). For pTrq2-Cldn2 or pTrq2-Cldn2Stx the ratio was changed in all approaches to 1:5. Transfected HEK293 cells were visualized on a LSM510-NLO inverted microscope (Carl Zeiss Microscopy GmbH, Jena, Germany) using a 40x/1.3 oil objective. Trq2 fluorescence signals were recorded (IR laser, λ exc = 810 nm, two-photon technique, META detector, spectral range 436–650 nm and split using a MBS KP 700). Channel pictures were taken prior to the recording of spectra in order to estimate expression of the Trq2 and YFP-tagged constructs (Trq2: IR laser λ exc 810 nm, two-photon technique, λ em = 430–505 nm, META detector; YFP: argon laser, λ exc = 514 nm, λ em = 560 nm long pass filter). FRET data analysis was performed using the software ZEN2010 (Carl Zeiss Microscopy, Jena, Germany), and Excel2013 (Microsoft Office), respectively. For the analysis, ROIs were set on the contact side of two neighboring HEK293 cells that were expressing both claudins. The fluorescence spectra of Trq2-tagged constructs in presence and absence of the YFP-tagged constructs receptor were measured. To calculate the FRET-based fluorescence, a λ-stack with a linear spectral unmixing mode was used to correct any YFP fluorescence crosstalk into the FRET channel (523–532 nm). The λ-stack is an integral part of the confocal laser system software. YFP correction was carried out to correct for direct excitation of the acceptor during donor excitation. The effect was expressed by changes in the FRET ratio, which is calculated by dividing the acceptor emission (YFP, λem = 532 nm) by the donor emission (Trq2, λem = 478 nm).

Strand lengths measurement COS-7 cells were transfected with pSNAP-Cldn2 and pYFP-Cldn10a in three different ratios (Cldn2/Cldn10a: 3:1, 1:3) with a total plasmid conc. of 1.6 µg/ml. In addition, phalo-C1 was used to normalize the plasmid concentration over the different ratios. For the detection and protein amount determination via Western Blot (see: Immunoblot-based analysis) the antibodies rabbit anti-SNAP (New England Biolabs Inc., #P93105) (1:1000) and mouse anti-GFP (Thermo Fisher Scientific, #A1120) (1:2000) as well as a mouse anti-HSP70 (Thermo Scientific Fisher, #MA3006) (1:5000) were used. For the strand length analysis via IF (see: Immunofluorescence of overexpressed claudins), the cells were labeled with BG-Atto590 and immunofluorescent stained with α-GFP-NB-Atto647N (1:200) and imaged in two-color STED (see the “Time-gated single- and multi-color STED imaging” section). For the measurement of each condition, 40 individual strands of 3–4 different meshworks were analyzed using the segmented line function of Fiji ImageJ.

Four STED images of three different isolated proximal tubules stained with mouse anti-Cldn2, donkey anti-mouse Atto647N and rabbit anti-Cldn10 rabbit, donkey anti-rabbit AF594 (see: Immunofluorescence of single murine tubules from proximal tubule) were taken and imaged with the same settings (1024 × 1024, 16x line averaging, pixel size of 18.9 × 18.9 nm). For the measurement ≥132 individual claudin strands per image were analyzed using the segmented line function of Fiji ImageJ.

Cholesterol depletion assay COS-7 cells were transfected with pSNAP-Cldn2 and pYFP-Cldn10a with a total plasmid conc. of 1.6 µg/ml and treated with 10 μM Mevas- tatin (Sigma-Aldrich, #567022-SMC) for 24 h in DMEM with FBS or 10 mM Methyl-β-cyclo-dextrin (MβCD) (Sigma Aldrich, #C4555-1G) for 1 h in DMEM without FBS. As control in an additional approach the equal amount of H2O was used. The transfected cells were labeled with BG-Atto590 and immunofluorescent stained with α-GFP-NB-Atto647N (1:200) and imaged in two-color STED.

For Filipin III staining, the cells were fixed with 4% PFA/Sucrose for 15 min, quenched with 0.1 M glycine in PBS for 30 min and incubated for 2 h with a freshly prepared Filipin III solution (final concentration: 0.05 µg/ml) (Sigma-Aldrich, #F4767-IMG) under light protection. The cells were washed with PBS and imaged with a Leica SP8 TCS microscope (Leica Microsystems). Filipin III staining was imaged with the following settings: excitation, a UV-laser was used (405 nm) and the emission filter was set to 415–470 nm. The fluorescence signal was detected by a photomultiplier tube (PMT). Images were acquired with a HC PL APO CS2 100 ×/1.4 NA oil objective (Leica Microsystems), a scanning format of 1024 × 1024 pixels, 8-bit sampling, 6x line averaging and 1x optical zoom, yielding a pixel dimension of 113 × 113 nm.

Pore blocking assay COS-7 cells were transfected with pSNAP-Cldn2Stx and pYFP-Cldn10a with a total plasmid conc. of 1.6 µg/ml. Cells were either treated with 1 mM MTSET (Biotium, #567022-5MG) for 24 h in DMEM with FBS or 10 mM Methyl-β-cyclo-dextrin (MβCD) (Sigma Aldrich, #C4555-1G) for 1 h in DMEM without FBS. As control in an additional approach the equal amount of H2O was used. The transfected cells were labeled with BG-Atto590 and immunofluorescent stained with α-GFP-NB-Atto647N (1:200) and imaged in two-color STED.

Co-culture assay COS-7 cells were seeded in 6-well plates and transfected with pSNAP-Cldn2Stx, pYFP-Cldn10a, or pYFP-Cldn2. After 24 h the cells were detached and intensively washed with PBS. After resuspension in DMEM pSNAP-Cldn2Stx and pYFP-Cldn10a transfected cells were mixed with pYFP-Cldn2 or pYFP-Cldn10a transfected cells in a ratio of 1:1 and seeded on 25 mm glass coverslips (#1L3H) and incubated for 24 h at 37 °C and 5% CO2. For SNAP-Tag labeling the BG-Atto590 was used (see: Cell culture). And the YFP-signal was boosted with the α-GFP-NB-Atto647N
Electrophysiological of monovalent ion permeabilities and fluorescein flux measurements

For the electrophysiological and flux measurement cells (MDCKII, MDCKII QKO, MDCKII QKO FLAG-Cldn2/3/10a/15, MDCKII QKO FLAG-Cldn2 + FLAG-Cldn10a, MDCKII QKO FLAG-Cldn3 + FLAG-Cldn15) were seeded on 0.6 cm² and 0.4 µm pore sized transwell filter (see: cell culture). In addition, for all measurements, cells were seeded on filters or in 6-well plates for an additional immunofluorescence labeling and immunoblotting approach to control the claudin expression. The measurements for the ion permeabilities were performed with an Ussing-Chamber as described in detail in Günzel et al. 2009. The measurements were performed in circulating Ringer’s solution (21 mM NaHCO₃, 119 mM NaCl, 5.4 mM KCl, 1 mM MgSO₄, 1.2 mM CaCl₂, 3 mM HEPES, 10 mM glucose; pH of 7.4) with a total concentration of 140 mM Na⁺ and 128.8 mM Cl⁻. The solution was gazed and mixed using a bubble lift (95% O₂ and 5% CO₂) and warmed up to 37 °C. For dilution potential measurements, each side of the Ussing-Chamber was filled with 5 ml Ringer’s solution. After acclimatization of the cells, 5 ml of a modified Ringer’s solution containing 238 mM mannitol and only 80.5 mM Na⁺ and 69.3 mM Cl⁻ were added to the apical bathing solution. At the same time an equal amount of Ringer’s solution was added to the basolateral bathing solution. The trans-epithelial resistance (TER) and voltage were recorded (every 10 s) during the whole experiment. The relative permeability ratios for Na⁺ and Cl⁻ (PNa/PCl) were calculated from the relative permeabilities and voltages according to the Goldman-Hodgkin-Katz equation. Details are described in Günzel et al. 2009. The absolute permeabilities (PNa, PCl) were calculated from the relative permeabilities and trans-epithelial resistances as described in Günzel et al. 2009. For the flux measurement of fluorescein (332.31 Da), the bathing solution was changed back on both sides of the chamber to a Ringer’s solution (21 mM NaHCO₃, 119 mM NaCl, 5.4 mM KCl, 1 mM MgSO₄, 1.2 mM CaCl₂, 3 mM HEPES, 10 mM glucose; pH of 7.4) with a total concentration of 140 mM Na⁺ and 128.8 mM Cl⁻. The solution was gazed and mixed using a bubble lift (95% O₂ and 5% CO₂) and warmed up to 37 °C. For dilution potential experiments, each side of the Ussing-Chamber was filled with 5 ml Ringer’s solution. After acclimatization of the cells, 5 ml of a modified Ringer’s solution containing 238 mM mannitol and only 80.5 mM Na⁺ and 69.3 mM Cl⁻ were added to the apical bathing solution. At the same time an equal amount of Ringer’s solution was added to the basolateral bathing solution. The trans-epithelial resistance (TER) and voltage were recorded (every 10 s) during the whole experiment. The relative permeability ratios for Na⁺ and Cl⁻ (PNa/PCl) were calculated from the relative permeabilities and voltages according to the Goldman-Hodgkin-Katz equation. Details are described in Günzel et al. 2009. The absolute permeabilities (PNa, PCl) were calculated from the relative permeabilities and trans-epithelial resistances as described in Günzel et al. 2009. For the flux measurement of fluorescein (332.31 Da), the bathing solution was changed back on both sides of the chamber to a Ringer’s solution. A voltage clamp was applied to the system and voltage was recorded (every 10 s) during the whole experiment. The relative permeability ratios for Na⁺ and Cl⁻ (PNa/PCl) were calculated from the relative permeabilities and voltages according to the Goldman-Hodgkin-Katz equation. Details are described in Günzel et al. 2009. The absolute permeabilities (PNa, PCl) were calculated from the relative permeabilities and trans-epithelial resistances as described in Günzel et al. 2009. For the flux measurement of fluorescein (332.31 Da), the bathing solution was changed back on both sides of the chamber to a Ringer’s solution. A voltage clamp was applied to the system and voltage was recorded (every 10 s) during the whole experiment. The relative permeability ratios for Na⁺ and Cl⁻ (PNa/PCl) were calculated from the relative permeabilities and voltages according to the Goldman-Hodgkin-Katz equation.

Statistics and reproducibility

All data were derived from at least three independent experiments and are presented as means ± standard deviation (SD) or median and interquartile range unless differently noted in the figure legend. Comparisons among groups were performed using a one-way ANOVA and additional Dunnett’s multiple comparison test or Tukey’s multiple comparison test or Kruskal–Wallis test. Comparisons between two independent groups, statistical significance was analyzed with a two-tailed non-parametric Mann-Whitney test in GraphPad Prism Version 5.04. The level of significance is indicated in the figures by asterisks (*P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001, ****P ≤ 0.0001). No statistical method was used to predetermine sample sizes as sample sizes were not chosen based on a prespecified effect size. Instead, multiple independent experiments were carried out using several sample replicates as detailed in the figure legends.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

All relevant data supporting the key findings of this study are available within the article and its Supplementary Information files or from the corresponding author upon reasonable request. Source data are provided with this paper.

Code availability

Code and example data to reproduce data from Fig. 2 are available here: https://doi.org/10.5281/zenodo.7009994.

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Author contributions

H.G. performed experiments and analyzed data, C.S. wrote code and performed image and data analysis with help from R.E.v.d.V., J.E. and H.G. acquired and analyzed FRET data, H.G. and D.G. acquired and analyzed ion flux data, R.E.v.d.V. acquired & quantified STED images of selected Claudins with ER, N.H. and M.B. prepared & stained mouse kidney tubules and found initial segregation phenotype in kidney; D.G., J.P., M.B. and MF provided reagents and analyzed data, H.G., V.H. and M.L. designed the study, analyzed data, and wrote the manuscript with input from all authors.

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Competing interests

The authors declare no competing interests.

Additional information

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