Conformational changes and CO₂-induced channel gating in connexin26

Graphical abstract

Highlights
- High-resolution cryo-EM structures of connexin26 at varying levels of PCO₂
- CO₂ alone causes conformational changes in the protein under stable pH conditions
- The N-terminal helices regulate the aperture of the pore
- KID syndrome mutations affecting CO₂ sensitivity map to flexion points of structure

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In brief
Brotherton et al. present cryo-EM structures of the connexin26 gap junction at three concentrations of CO₂. They show that at constant pH, the aperture of the pore varies depending on CO₂ concentration. Their results correlate with physiological measurements showing that elevated CO₂ closes gap junctions.

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**Conformational changes and CO2-induced channel gating in connexin26**

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**SUMMARY**

Connexins form large-pore channels that function either as dodecameric gap junctions or hexameric hemichannels to allow the regulated movement of small molecules and ions across cell membranes. Opening or closing of the channels is controlled by a variety of stimuli, and dysregulation leads to multiple diseases. An increase in the partial pressure of carbon dioxide (PCO2) has been shown to cause connexin26 (Cx26) gap junctions to close. Here, we use cryoelectron microscopy (cryo-EM) to determine the structure of human Cx26 gap junctions under increasing levels of PCO2. We show a correlation between the level of PCO2 and the size of the aperture of the pore, governed by the N-terminal helices that line the pore. This indicates that CO2 alone is sufficient to cause conformational changes in the protein. Analysis of the conformational states shows that movements at the N terminus are linked to both subunit rotation and flexing of the transmembrane helices.

**INTRODUCTION**

Twenty connexin genes have been identified in the human genome (Abascal and Zardoya, 2013). Connexins form hexameric plasma-membrane channels, or hemichannels, that can dock together to give dodecameric gap junctions. Gap junctions provide a direct aqueous passageway between cells, with functions independent of hemichannels (Pearson et al., 2005; Stout et al., 2004; Weissman et al., 2004). Connexin mutations underlie many different pathologies affecting all organ systems of the body (Garcia et al., 2016). Mutations of Cx26 constitute a leading cause of congenital deafness (Xu and Nicholson, 2013). Several rare, but dominant, mutations cause a severe syndromic disease: keratitis ichthyosis deafness syndrome (KIDS) (Xu and Nicholson, 2013).

The conductance of gap junctions and hemichannels can be modulated by voltage (Valiunas, 2002; Young and Peracchia, 2004), pH (Bevans and Harris, 1999; Khan et al., 2020; Yu et al., 2007), and, in some instances, by intracellular Ca²⁺ (Peracchia, 2004). We have discovered that a small group of closely related β-connexins can be directly modulated by modest physiologically relevant changes in the partial pressure of CO2 (PCO2) in arterial blood (Huckstepp et al., 2010a; Meigh et al., 2013). The effects of CO2 on Cx26 gap junctions and hemichannels, respectively, are diametrically opposite: in hemichannels, CO2 causes channel opening (de Wolf et al., 2017; Dospinescu et al., 2019; Huckstepp et al., 2010a; Meigh et al., 2013), whereas in gap junctions, CO2 causes channel closure (Nijjar et al., 2021). Physiological and mutational analyses suggest that this CO2 sensitivity is independent of pH (Huckstepp et al., 2010a; Meigh et al., 2013), is intrinsic to the protein (Meigh et al., 2013), and depends on a specific lysine (Dospinescu et al., 2019; Huckstepp et al., 2010a; Meigh et al., 2013, 2015). Binding of CO2 to this specific lysine of Cx26 is physiologically important, as it contributes nearly half of the centrally generated chemosensory regulation of breathing to modest levels of hypercapnia (Huckstepp et al., 2010b; van de Wiel et al., 2020).

Several structures of Cx26 gap junctions have previously been reported, solved by both X-ray crystallography (Bennett et al., 2016; Maeda et al., 2009) and electron microscopy (Khan et al., 2020; Oshima et al., 2007) at low to modest resolution. The hemichannel is composed of 6 connexin subunits arranged around a central funnel. Each subunit consists of 4 transmembrane helices (TM1–4), an N-terminal helix, which lines the funnel entrance, a cytoplasmic loop, two extracellular loops, and a short C-terminal tail. While residues in the extracellular loops, which are involved in hemichannel docking, are generally well ordered among the solved structures, residues on the cytoplasmic side, including the N-terminal helix and the cytoplasmic loop, are much less well defined. Together with structures of other connexins (Flores et al., 2020; Lee et al., 2020; Myers et al., 2018), we are building up a picture of how these proteins are regulated. There remain, however, conflicting ideas of how they open and close in response to various signals. Original ideas, based on low-resolution electron crystallography, posited a mechanism involving subunit rotation (Unwin and Ennis, 1984). However,
the presence of a flexible N-terminal helix in the pore funnel has led to the idea that this may form a movable plug to block and open the channel (Maeda et al., 2009; Oshima et al., 2007, 2011; Yue et al., 2021). Yeager and co-workers suggested that regulation of Cx26 by extracellular Ca²⁺ was conferred by an electrostatic mechanism involving very minor modifications to the protein (Bennett et al., 2016), while in looking at regulation by pH, the same group have proposed a ball-and-chain mechanism involving dramatic conformational changes in which the N terminus completely refolds (Khan et al., 2020).

Here, we report high-resolution structures of Cx26, solved by cryoelectron microscopy (cryo-EM), from protein vitrified in buffers containing three levels of PCO₂ at constant pH. We show that CO₂ by itself is sufficient to change the conformation of the protein. Analyses of the conformational variability among these structures show that movements of the transmembrane helices, in particular, TM2, are correlated with the position of the N-terminal helix within the pore.

RESULTS

Cryo-EM structures of Cx26 under different levels of PCO₂

We first collected data from Cx26 vitrified in a CO₂/bicarbonate (HCO₃⁻) buffer corresponding to a PCO₂ level of 55 mmHg and a pH of 7.4. This resulted in 3D reconstructions with nominal resolutions of 2.5 Å as defined by gold-standard Fourier shell correlations (FSC₆₈) (Rosenthal and Henderson, 2003; Scheres, 2012) without applying symmetry and 2.2 Å when dihedral group 6 (D₆) symmetry, consistent with the dodecameric structure of the gap junction, was used (Figure S1; Table S1). Remarkably, this resolution, limited by the pixel size during data collection, was much higher than similar preparations of protein vitrified in HEPES, where we could only obtain resolutions of approximately 4.5 Å, more similar to those observed by others (Khan et al., 2020).

From this, we inferred that the CO₂/HCO₃⁻ buffer was stabilizing the protein. Encouraged by this, we therefore collected two further datasets at higher (90 mmHg) and lower (20 mmHg) PCO₂ levels. To ensure that differences between the two structures were confined as much as possible to only the PCO₂ levels, the protein was from the same preparation, being divided only in the final buffer-exchange step. The buffers were chosen to ensure that the pH and ionic strength did not differ (see STAR Methods and Figure S2). These datasets, collected with a smaller pixel size, gave reconstructions at even higher resolutions than those of the first data collected (1.9 Å for the 90 mmHg PCO₂ data and 2.1 Å for the 20 mmHg PCO₂ data with D₆ symmetry; 2.1 Å and 2.7 Å, respectively, without symmetry; Figures S3 and S4; Table S1). While it was noted that there was some asymmetry in reconstructions, when symmetry was not applied, it was difficult to draw out any more detailed structural information than was seen in the D₆ maps.

In all datasets, the overall quality of the density in the D₆ reconstructions is very similar. On the extracellularex side of the protein, the density is excellent, with clearly defined side chains, water molecules, and density consistent with the hydrocarbon chains of lipid or detergent (Figures 1 and S5–S7). On the other hand, the cytoplasmic side of the protein is less well defined, and there is no consistent density for the cytoplasmic loop between TM2 and TM3. In all maps, we observe density, which we have been unable to unambiguously assign, that protrudes from near the top of the visible density of TM3 toward the N-terminal helix (Figures S5–S7). In the structure of a related connexin, Cx46/50, more distinctive density in a similar position has been modeled with residues from the loop preceding TM3 (Flores et al., 2020). This corresponds to residues 127 to 130 of Cx26 and would thus be adjacent to the residues implicated in regulation by CO₂ (Dospinescu et al., 2019; Huckstepp et al., 2010a; Meigh et al., 2013, 2015). In structural predictions of Cx26 by both AlphaFold (Jumper et al., 2021) and RosettaFold (Baek et al., 2021), this region is modeled, albeit with a low confidence level, in a similar fashion, rather than by the continuation of the alpha helix that is present in the Cx26 crystal structure (Figure S8). It is possible that the density we observe also corresponds to these residues. We also see density reminiscent of a very short hydrocarbon chain of lipid or detergent lining the pore wall between the N terminus and TM1 and TM3 in all maps (Figures S5–S7). It is possible that the density is due to the dodecyl β-D-maltoside (DDM) that may have inserted into the pore during solubilization and became anchored through the interaction of its hydrocarbon chain with the hydrophobic wall of the funnel.

The major difference in the connectivity of the density associated with the different levels of PCO₂ concerns the density for the N-terminal helix (Figure 2). At the higher levels of PCO₂, the N-terminal helices fold into the pore such that there is a narrow constriction where all of the N-terminal helices from the six subunits meet (Figure 1). The density for these helices is more fragmented for the structure at low PCO₂, indicative of more flexibility of the helices associated with these particles (Figure 2).

Comparison with other Cx26 structures

After refinement of the structures against the respective D₆ averaged maps associated with each level of CO₂, the coordinates look very similar, apart from at the N terminus. The resolution is much higher than has been seen for previous structures of Cx26 (Bennett et al., 2016; Khan et al., 2020; Maeda et al., 2009), enabling the protein to be modeled more accurately. Relative to these structures, the most interesting differences are observed at the N terminus and in TM1 (Figures 3 and S8). TM1 lines the inner surface of the cytoplasmic funnel (Figure 1). In other structures of Cx26, it has been built as a regular helix from residues Lys22 to Glu42 where it kinks as it turns into a 3₁₀ helix (Bennett et al., 2016; Khan et al., 2020; Maeda et al., 2009). This region has been described as the TM1/extracellular loop 1 (TM1/E) interface. In our structures, the conformation of residues Val37 to Glu42 differs from that seen in previously published structures. Given that these residues are located within the pore, are adjacent to the constriction formed by the N termini (Figures 3 and S8), and are critical for function (Bennett et al., 2016; Verselis et al., 1994; Xu and Nicholson, 2013), we investigated whether this was a true conformational variation among the structures or whether it could have arisen from difficulties in map interpretation in the lower-resolution structures. We therefore reanalyzed the data from the previous crystal structures (see STAR Methods). While we could not unambiguously distinguish between these scenarios for the crystal structure from Maeda et al. (2009), the structures published by
Bennett et al. (2016) were consistent with a conformational change (Figure S8).

The only other moderate resolution structure of Cx26 where there is distinct density for the N terminus is the crystal structure solved by Maeda et al. (2009). Comparing this structure with our structure at 90 mmHg PCO2 shows clear differences in this region. Firstly, the trajectory taken by the N terminus into the pore differs slightly in the two structures, giving the constriction at the bottom a slightly different aperture (Figure 3). Secondly, the link between the N terminus and TM1 has been modeled farther from the funnel entrance in the crystal structure, incorporating the density that we observe to link TM3 and the N terminus (Figure 3). It has been shown that this conformation is not stable in molecular dynamics simulations (Myers et al., 2018), and, while unambiguous interpretation of the density in this area in the crystal structure is difficult, inspection of the respective maps shows that there are differences in how the N-terminal helix is situated in the pore with respect to the 90 mmHg PCO2 Cx26 structure.

Structures have also been reported for two other connexins, the Cx46/50 (Flores et al., 2020; Myers et al., 2018) gap junction and the Cx31.3 (Lee et al., 2020) hemichannel. The core regions of these proteins, which correspond to the complete sequence of Cx26, have 50% and 35% sequence identity to human Cx26, respectively, and the profiles of the pores are very similar to Cx26, as seen in Figure 1A. However, whereas in Cx46/50, the N terminus (residues 1–16) is tucked back against the hydrophobic wall of the funnel, giving a much more open structure, in Cx31.3, the N-terminal helix is located closer to the cytoplasmic entrance such that it forms a lid with a narrow constriction of ~13 Å level with the cytoplasmic end of the transmembrane helices (Lee et al., 2020) rather than a plug within the funnel (Figure 3).

**Conformational variability in Cx26 at fixed PCO2**
The cytoplasmic part of Cx26 is clearly flexible as judged, not only from the density and structural comparison but also from elastic network modeling, where considerable flexing of the subunits and transmembrane helices can be observed (Meigh et al., 2013). As cryo-EM datasets can harbor a range of conformations, we analyzed particle subsets in two ways. First, we used variability analysis as implemented in cryoSPARC (Punjani and Fleet, 2021; Punjani et al., 2017) (Videos S1, S2, and S3).
variety of analyses were carried out selecting the hemichannel with the use of a soft-edged mask. Consistently, the most variation was seen at the N termini and in the cytoplasmic sides of the transmembrane helices. In the respective maps derived from data collected at 55 and 90 mmHg PCO2, the N termini oscillated between a conformation where the helices met together in the core of the funnel and a position where the N termini were less defined in the density but pulled back from the pore (Videos S1 and S2). Together with this, the cytoplasmic portions of the transmembrane helices flexed with the greatest movement seen for TM2. In maps derived from data collected from protein vitrified at under 20 mmHg PCO2 conditions, the general trend was the same, but the N termini were never as well defined as those corresponding to higher PCO2, indicative of more mobility of the N termini (Video S3).

This analysis gave some idea of the variability within the datasets. We next sought to use a process of particle expansion, subtraction, and masked, fixed-angle classification in Relion (Scheres, 2012; Zivanov et al., 2018) to investigate whether we could select out particle sets with more consistent conformations in these regions (see STAR Methods). Firstly, we carried out an analysis with C6 symmetry, selecting out the cytoplasmic side of the protein with the use of a mask. The analysis for the data corresponding to the PCO2 of 90 mmHg was the most interesting. Here, out of the eight given classes, the majority of particles divided into three classes, which, in order of particle number, result in maps with nominal resolutions of 2.0, 2.1, and 2.6 Å (Figures S9 and S10). Overall, the maps derived from particles containing 20 and 55 mmHg PCO2 datasets, we only obtained one class in each case with sufficient particles to give high-resolution maps. In each case, the positions of the transmembrane helices in the D6 refined structure resembled most the top class from this classification.

To understand the conformational variation within the 90 mmHg PCO2 data in more detail, structures were built and refined from the maps associated with the top two classes (see Table S3). Comparing the two resulting structures (denoted hereafter as structure Class 1 [C1] and C2) shows that the cytoplasmic part of TM2 flexes at Pro 87 (Figure 4) such that there is an almost rigid body rotation of 14° of the C terminal part of the helix from residues Ala88 to Glu101. This region of TM2 is located between the loop linking the N-terminal helix with TM1 (residues 14 to 18) of the same respective subunit and the top of TM1 of the neighboring subunit. In structure C2, where the N terminus is more defined, TM2 is straighter with interactions between Val95 and His16, though the latter is not well defined in the density. In Structure C1, TM2 is bent toward TM1 of the neighboring subunit so that there is an interaction between Leu90 of TM2 and Trp24 of TM1. Morphing between these structures indicates that as TM2 moves, the N terminus and the following residues, including the first few residues of TM1, also change position (Video S5). The density associated with the side chains in this region is not well defined in either map.
Overall, the conformational change has the aspect of a rotation of the cytoplasmic region of the protein with respect to the extracellular side.

**Effect of PCO2 on conformational distribution of Cx26**

Clearly, the above classification procedure resulted in our being able to discriminate distinct conformations of the protein at a fixed level of PCO2. However, given that the analysis of the 20 and 55 mmHg PCO2 data was hampered by low particle numbers in all but the top classes, the question remained as to how differing levels of PCO2 could alter the conformation of Cx26. Our inference from all the maps we examined, irrespective of particle number, lowpass resolution filter, masking, etc., was that the density associated with the N terminus was more distinct in the data obtained at high PCO2 relative to that with low PCO2. This suggests that elevated CO2 might bias the conformations that are sampled by the N terminus. To examine this more quantitatively, we carried out a further classification without imposing symmetry and with the same numbers of particles from each PCO2 dataset. As the N terminus is located at the interface of two subunits, we created a mask incorporating two subunits and again carried out a process of particle expansion, subtraction, and masked, fixed-angle classification as above, but using four classes (see STAR Methods). The distributions of particles in each of the classes can be seen in Figure 5. With the 90 mmHg PCO2 data, which contained 58% of the particles, the density associated with the N terminus was less defined and indicative of adopting a position nearer to the cytoplasmic side (Figure 5). Altogether, the results from this classification, as seen in Figure 5, support there being a difference between how well the N terminus is defined, with a trend from higher PCO2 to lower PCO2.

**DISCUSSION**

The structures of Cx26 reported here together with the analysis provide insight into the mechanism of these channels. Connexin gap junctions and hemichannels open and close in response to a variety of stimuli with models indicating either blockage through the N-terminal helices or a twisting iris-like movement (reviewed in Oshima, 2014). In our analysis, we see elements of both. Of the structures reported here, the narrowest constriction is observed for the 90 mmHg PCO2 data where the diameter of the pore is approximately 11 Å, slightly narrower than that of the Cx26 crystal structure but much narrower than the pore seen in Cx46/50 (Figure 3). Only slight modifications of the side chains would be required to close the channel completely. The pore diameter is more similar to that of Cx31.3, where the N-terminal helices are closer to the cytoplasm (Figure 3). Whether the conformations seen in either Cx46/50 or Cx31.3 represent possible open and closed conformations that could be adopted in Cx26 is difficult to ascertain. The sequence identity, between the residues of the N-terminal helices in Cx31.3 and Cx26, is only 20%, with little conservation. In Cx46, where the N termini are more conserved
with respect to Cx26 (30% sequence identity), Trp4 of the N-terminal helix slots between Ala39 of TM1 of one subunit and Ile34 and Leu35 of the next (Figure S11). In Cx26, the equivalent residue to Ala39 is replaced with the slightly bulkier residue valine (Val38). Leu35 is also replaced by a methionine (Met 34), for which mutation to alanine or threonine causes a loss of function (Oshima et al., 2003) and reduction in the response to CO2 in hemichannels (de Wolf et al., 2016) and an apparent structural change in the N-terminal helices (Oshima et al., 2007). These conservative substitutions may affect the stability of the position of the N-terminal helix. This interpretation, however, does need to be taken with caution, as this is the site of the presumed detergent that we observe in the binding pocket (Figure S11). Were the detergent not present, we cannot rule out that the N termini would adopt a similar position to that seen in Cx46/50 in the open conformation rather than just becoming more flexible, as we observed at low PCO2. More extensive lipid-like density in this region in Cx31.3 led to speculation that lipids may actually play a role in the mechanism (Lee et al., 2020) by analogy to a similar suggestion for innexins, gap-junction proteins in pre-chordates (Burendei et al., 2020).

The conformation that we observe at the TM1/E boundary in our structures, which includes Val38, discussed above, is more similar to the structures of both Cx46/50 (Flores et al., 2020) and Cx31.3 (Lee et al., 2020) than to the previously reported Cx26 crystal structures (Bennett et al., 2016; Maeda et al., 2009). From an examination of the density associated with each structure, this appears to be a distinct conformational difference rather than differences due to modeling. Why this should be is unclear. In our structures, there does appear to be some
degree of flexibility in this area, and the variability analysis from
cryoSPARC for the structure at low PCO₂ indicates a slight
breathing at this point. This region includes the site of one of
the KIDS mutations, Ala₄₀ to valine (Xu and Nicholson, 2013).
Hemichannels with this mutation are less responsive to extracel-
lar Ca²⁺ (Sanchez et al., 2010), and both gap junctions and
hemichannels are non-responsive to CO₂ (Cook et al., 2019; Nij-
jar et al., 2021). With the change in conformation relative to
the previously published structures, Ala₄₀ moves from a position
that is more accessible to the pore to a more restricted position
juxtaposed to TM₂. This has important consequences for
considering the effects of the mutation, as replacement of the
alanine in this position with the bulkier valine is likely to perturb
the structure and restrict any conformational changes necessary
for activation, explaining why it may have more of an effect than
might be expected from the previous structures (Figure 3). The
region is also next to the Ca²⁺ binding site that is observed in
the crystal structure with bound Ca²⁺, with Glu₄₂ acting as one
of the Ca²⁺ ligands (Bennett et al., 2016). As no large-scale
conformational differences were observed between Ca²⁺-bound
and free structures, an electrostatic mechanism was proposed
to explain how Ca²⁺ was able to prevent the passage of ions
through Cx₂₆. Although this binding site for Ca²⁺ has been ques-
tioned (Lopez et al., 2016), it is also possible that the interaction
of Glu₄₂ with Ca²⁺ leads to the conformational changes we
observe between the Ca²⁺-bound structures and our structures
(Figure S8). It can be speculated that any change in this area will
affect the interaction with the N-terminal helix, as discussed
above. It has previously been shown that Lys₄₁, Glu₄₂, and
the N terminus affect voltage gating (Verselis et al., 1994).

In addition to these conformational differences with respect to
other published structures, all our analyses indicate a great deal
of flexibility in the N terminus and in the cytoplasmic parts of
the transmembrane helices. From the data associated with the
90 mmHg PCO₂, we teased out two distinct conformations of
the protein. In one, the position of the N terminus is relatively
well defined, such that it folds down to a narrow aperture within
the funnel. In the other, the cytoplasmic region of the protein is
rotated relative to the extracellular region, and the N terminus
is less defined, giving the appearance of a more open pore. A
comparison of the two structures also shows that TM₂ twists
around Pro₈₇ and that the density between TM₃ and the N ter-
minus adopts a different trajectory. The rotation that is observed
is, in fact, reminiscent of an early study on the regulation of Cx₂₆
where pore closure was associated with a similar rotation of the
cytoplasmic region (Unwin and Ennis, 1984). The presence of
Pro₈₇, which is highly conserved within connexins, has also
been shown to be important for function (Suchyna et al., 1993),
though whether the flexion of the helix that we observe stems
from the proline itself is difficult to say given the quality of the

Figure 5. Classes observed with a 2-subunit mask at the three levels of PCO₂
From each dataset, 125,000 particles were randomly selected from the final D6 refinement. These were symmetry-expanded in D6, and the same mask was used
for particle subtraction and 3D classification, with appropriate changes in pixel and box size for the 55 mmHg data. The percentage of particles in each class is
shown below each class. Red boxes indicate the full, extended N-terminal helix (NT) position, and blue boxes indicate the truncated, raised NT location. The
remaining two classes in each data set represent intermediates between these two extreme classes. The classes have been arranged in approximate order of the
clarity of the density for the NT in the down position. This is clear for the two extremes but more ambiguous for the intermediate classes.
maps in this region. While we cannot say the range of movements seen represents the full ensemble required for occlusion or opening of the channel, owing to the lack of connectivity for the cytosolic loop and C terminus, there is clearly a series of positions that are intermediates, which together suggest how the protein may transition between open and closed states.

There are nine mutations that give rise to KIDS (Xu and Nicholson, 2013) (Figure 4). Ala40Val has been discussed above, and G45 is located nearby. Four of the mutations occur in the highly flexible regions of the protein near the flexion point of TM2 and in the linker between the N-terminal helix and TM1. These mutations involve substitution of bulkier side chains (Ala88Val and Gly12Arg, Asn14Lys/Tyr and Ser17Phe), which would likely impede the flexions required for channel gating (Video S5). Ala88 is situated next to Pro87 at the flexion point of TM2 and interacts with Val13 on the N-terminal helix between Gly12 and Asn14. Ala40Val, Ala88Val, and Asn14Lys have all been observed to abolish CO2-mediated gap-junction closure (Nijjar et al., 2021).

We have previously reported that in a cellular environment, Cx26 gap junctions and hemichannels can be regulated by changes in PCO2 (de Wolf et al., 2017; Dospinescu et al., 2019; Huckstepp et al., 2010a; Meigh et al., 2013). In this structural study, we clearly see an effect of the CO2/HCO3− buffering system on the resolution of the maps that can be obtained relative to that observed when HEPES is used. This dramatic increase in resolution mirrors the improvement that was seen for Cx46/50 when cryo-EM was carried out with nanodiscs rather than amphipols (Flores et al., 2020; Myers et al., 2018) and suggests that the protein is being stabilized by CO2/HCO3−. In cells, an increase in PCO2 causes gap junctions to close (Nijjar et al., 2021). In our analysis, we observe that the most constricted pore is seen at the highest level of PCO2, which is in broad agreement with the results in cells. It may be that the exact conformation of the N terminus is affected by the detergent in the pore; however, as this is present in all three datasets, we would not expect it to be the cause of the difference. Neither would we expect the difference to be caused by the particle numbers, as reducing the numbers of particles in the high PCO2 datasets gives similar maps. Clearly, multiple conformations are being sampled in all three datasets collected, but it would appear that the more constricted pore is seen only in those associated with high PCO2. The question must be asked why CO2 would have this effect. A comprehensive series of mutational studies has led to the conclusion that the effect of CO2 is mediated by the direct binding of CO2 to Lys125 in a carbamylation reaction (Meigh et al., 2013, 2015; Nijjar et al., 2021; van de Wiel et al., 2020). Lys125 is located in the cytoplasmic loop, close to TM3. While we don’t see Lys 125 in the density, we do see a link between TM3 and the N terminus, which, as described above, may reflect the position of Lys125. Any effect caused by the change in charge of Lys125 caused by carbamylation could therefore be transmitted to the N terminus through this link. Confirmation of this will require further structural studies. It is possible that we have not been able to fully carbamylate the lysine with the preparation of Cx26 in detergent and that we only increase the proportion of the protein carbamylated as we increase PCO2. Regardless of this, we have shown how conformational changes in Cx26 can lead to channel opening and closing by movements of the N-terminal helix and that this effect is modified by buffers containing different levels of CO2 at constant pH. This ties in with previous studies, which propose that the N terminus acts to gate the channel in response to signals such as voltage or pH (Maeda et al., 2009; Oshima et al., 2007, 2011; Yue et al., 2021) and that there is a link between the position of the N-terminal helix, TM2, and the cytoplasmic loop (García et al., 2018; Valdez Capuccino et al., 2019). It is, however, very different from the “ball-and-chain mechanism” recently proposed, which is based on low-resolution structural information (Khan et al., 2020). This model relies on the N-terminal helix unwinding and plugging the central pore. To mediate the pH effect, profound acidification is required to cause the N-terminal helix to unfold and plug the channel, suggesting a function only in serious pathology. Here, we show that modest increases in PCO2 are sufficient to cause conformational changes of the N-terminal helix plug in the absence of a dramatic change in pH.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.str.2022.02.010.

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AUTHOR CONTRIBUTIONS

The project was initiated and supervised by N.D. and A.D.C. Cloning, expression, purification, and grid preparation were carried out by D.H.B. Data collection was performed by C.G.S., T.J.R., and D.H.B. Data processing was done by D.H.B. and A.D.C. with guidance from T.J.R. and C.G.S. D.H.B. and A.D.C. refined the structures, A.D.C., D.H.B., and N.D. wrote the manuscript with contributions from all authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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## STAR★METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and virus strains** | | |
| Sf9 Cells | Fisher Scientific | CAT# 10500343 |
| **Chemicals, peptides, and recombinant proteins** | | |
| Insect Xpress | Scientific laboratory Supplies/Lonza | CAT# LZBELN12-730Q |
| Histidine | Merck/Sigma | CAT# H6034-100G |
| n-Dodecyl (β)-maltoside (DDM-C) | Glycon Biochemicals GMBH | CAT# D97002-C-50g |
| cOmplete™, EDTA-free Protease Inhibitor Cocktail | Merck/Roche | CAT# 4693132001 |
| AEBSF hydrochloride | Fisher Scientific/Acros | CAT# 10742885 |
| Dithiothreitol | Fisher Scientific | CAT# 10592945 |
| DNase I | Merck/Roche | CAT# 10104159001 |
| Econocolumn | Bio-Rad | CAT# 7372532 |
| HisPur Ni-NTA Resin | Thermo Scientific/ Pierce | CAT# 88221 |
| 5/150 Superose 6 column | GE Healthcare Lifescience | CAT# 15383224 |
| Quantifoil 0.6/1 300 mesh Au holey carbon | Quantifoil Micro Tools GMBH | CAT# N1-C11nAu30-50 |
| UltrAuFoil 01.2/1.3 300 mesh Holey gold | Quantifoil Micro Tools GMBH | CAT# N1-A14nAu30-50 |
| UltrAuFoil 0.6/1 300 mesh Holey gold | Quantifoil Micro Tools GMBH | CAT# N1-A11nAu30-50 |
| Formvar/Carbon on copper 300mesh | EM resolutions Ltd. | CAT# FC300Cu100 |
| Vivaspin-20 MWCO 100,000 | Sartorius | CAT# VS2042 |
| Vivaspin-500 MWCO 100,000 | Sartorius | CAT# VS0141 |
| Vivaspin-8 MWCO 100,000 | Sartorius | CAT# VS0641 |
| Slide-a-lyzer, 10 k cutoff | Fisher Scientific/ Pierce | CAT# 11809420 |
| Custom mix 2.5 % CO2/97.5 % N2 | BOC | CAT# 226782-V |
| Custom mix 10 % CO2/90 % N2 | BOC | CAT# 226699-V |
| Custom mix 15 % CO2/85 % N2 | BOC | CAT# 120163-V |
| **Deposited data** | | |
| 20 mmHg PCO2 dodecamer Cryo-EM Volume | This Paper | EMDB: EMD-13940 |
| 20 mmHg PCO2 dodecamer Atomic Coordinates | This Paper | PDB: 7QET |
| 55 mmHg PCO2 dodecamer Cryo-EM Volume | This Paper | EMDB: EMD-13938 |
| 55 mmHg PCO2 dodecamer Atomic Coordinates | This Paper | PDB: 7QER |
| 90 mmHg PCO2 Cryo-EM Volume | This Paper | EMDB: EMD-13937 |
| 90 mmHg PCO2 Atomic Coordinates | This Paper | PDB: 7QEQ |
| Class 1 Hexamer, 90 mmHg Cryo-EM Volume | This Paper | EMDB: EMDB-13944 |
| Class 1 Hexamer, 90 mmHg Atomic Coordinates | This Paper | PDB: 7QEY |
| Class 2 Hexamer, 90 mmHg PCO2 Cryo-EM Volume | This Paper | EMDB: EMDB-13943 |
| Class 2 Hexamer, 90 mmHg PCO2 Atomic Coordinates | This Paper | PDB: 7QEW |
| Cx26 crystal structure | Maeda et al. (2009) | PDB: 2ZW3 |
| Cx26 with Ca2+ crystal structure | Bennett et al. (2016) | PDB: 5ER7 |

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Alexander Cameron (a.cameron@warwick.ac.uk).

Materials availability
This study did not generate unique reagents.

Data and code availability
Cryo-EM density maps have been deposited in the Electron Microscopy DataBank (EMDB) and atomic coordinates have been deposited in the Protein Data Bank. They are publicly available as of the date of publication. Accession numbers are listed in the key resources table. This paper does not report original code. Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Human connexin 26 protein (hCx26) was expressed by infecting Spodoptera frugiperda (Sf9) cells for 72 hours with a baculovirus carrying the hCx26 gene.

METHOD DETAILS

Protein expression and purification
Human connexin 26 protein (hCx26) was expressed with a thrombin-cleavable His(6) affinity tag on the C terminus using baculovirus in Spodoptera frugiperda (Sf9) cells. This construct was a gift from Prof Tomitake Tsukihara. Cells were harvested 72 hours post infection by centrifugation at 2500 x g in a Beckmann JLA 8.1000 rotor, cell pellets were snap frozen in liquid nitrogen, and stored at −80 °C until purification. Cells were thawed in hypotonic lysis buffer, (10 mM sodium phosphate, 10 mM NaCl, 5 mM MgCl2, 1 mM DTT, pH 8.0) DNAse I, cOmplete™ EDTA-free Protease Inhibitor Cocktail (Roche) and AEBSF were added according to manufacturer’s instructions before use. After 30 minutes stirring at 4 °C, the cells were broken using 40 strokes with a dounce homogeniser, and the membranes separated by ultracentrifugation for 1 hour at 4 °C, 158000 x g. Membranes were resuspended in membrane resuspension buffer (25 mM sodium phosphate, 150 mM NaCl, 5 % glycerol, 1 mM DTT, pH 8.0). DNAse I, cOmplete™ EDTA-free Protease Inhibitor Cocktail and AEBSF were added according to manufacturer’s instructions before use. This membrane suspension was diluted (to 400 ml) with solubilisation buffer (10 mM sodium phosphate, 300 mM NaCl, 5 % glycerol, 1 mM DTT, 1%
DDM (Glycon Biochemicals GmbH), pH 8.0) and incubated at 4°C for 3 hours, before a further 1 hour ultracentrifugation at 4°C, 158000 x g to remove insoluble material. The soluble material was batch bound to pre-equilibrated HisPure Ni-NTA resin (Thermo Scientific) overnight, and then poured into an econocolumn for subsequent manual washing and elution steps. Resin was washed with 5x CV wash buffer (10 mM sodium phosphate, 500 mM NaCl, 10 mM histidine, 5 % glycerol, 1 mM DTT, 0.1 % DDM, pH 8.0) before eluting hCx26 with elution buffer (10 mM sodium phosphate, 500 mM NaCl, 200 mM histidine, 5 % glycerol, 1 mM DTT, 0.1 % DDM, pH 8.0). Fractions containing hCx26 were dialysed overnight 4°C against (10 mM sodium phosphate, 500 mM NaCl, 5 % glycerol, 1 mM DTT, 0.03 % DDM, pH 8.0). For the protein used for electron microscopy thrombin (Sigma) 1:1 w/w was added during dialysis to remove the affinity label). hCx26 was then 0.2 μm filtered, concentrated in a vivaspin 100,000 MWCO and loaded onto a Superose 6 Increase 5/150 size exclusion chromatography column (GE Healthcare Lifescience) to remove thrombin and exchange the buffer to either 20 mm, 55 mm or 90 mmHg αCSF buffer (Huckstepp et al., 2010a) (20 mmHg αCSF buffer: 140 mM NaCl, 5 % glycerol, 1 mM DTT, 0.03 % DDM, 10 mM NaHCO₃, 1.25 mM NaH₂PO₄, 3 mM KCl, 1 mM MgSO₄, 2 mM MgCl₂; 55 mmHg αCSF buffer: 100 mM NaCl, 5 % glycerol, 1 mM DTT, 0.03 % DDM, 50 mM NaHCO₃, 1.25 mM NaH₂PO₄, 3 mM KCl, 1 mM MgSO₄, 2 mM MgCl₂; 90 mmHg αCSF buffer: 70 mM NaCl, 5 % glycerol, 1 mM DTT, 0.03 % DDM, 80 mM NaHCO₃, 1.25 mM NaH₂PO₄, 3 mM KCl, 1 mM MgSO₄, 4 mM MgCl₂).

**Cryo-EM sample preparation and data collection**

The hCx26 peak was concentrated to 3.5 mg/ml before being gassed with the correct amount of CO₂ to give a final pH of ~7.4 as described below for each buffer. 25% buffer, 10% and 15% CO₂ in N₂ (BOC) were used for the 20, 55 and 90 mmHg buffers respectively. 2.5%, 10% and 15% CO₂ in N₂ (BOC) were used for the 20, 55 and 90 mmHg buffers respectively. To ensure that the pH was consistent when the protein was vitrified in the relevant buffer, the following controls were applied. Firstly, 10 ml of buffer was equilibrated with the correct percentage of CO₂ for the buffer and the pH checked using a pH meter. Phenol red solution (1:200) was then added to the samples of gassed and ungassed buffers to provide a reference colour for the pH. The volume of the protein to be used for vitrification was noted, and a equal volume of phenol red/buffer was added to two separate Eppendorf tubes, one of which was gassed until the colour matched an equal volume of the pre-gassed 10 ml sample. The time of gassing was noted, and the protein was treated identically (Figure S2). For both the 20 mmHg and 90 mmHg conditions the sample was left overnight dialysing against 60 ml buffer with the correct concentration of CO₂ to give a pH of 7.4. Quantifoil 300 mesh gold grids (either 0.6/1 carbon film, 0.6/1 or 1.2/1.3 UltrAlufoil, Quantifoil Micro Tools GMBH) were glow discharged for 30 seconds prior to use. Vitrification of the protein in liquid ethane/propane at ~180°C was carried out with a Leica GP2 automated plunge freezer with 3 μl protein per grid at 4°C, 95 % humidity, 6 seconds blotting in a CO₂/N₂ atmosphere appropriate for the buffer used. Grids were screened using a Jeol 2100plus microscope, and data were collected on an FEI Titan Krios G3 equipped with a K3 detector and BioQuantum energy filter using a 20 e⁻/Å² slit width. A dose rate of ~10 e⁻/pix/sec on the detector was chosen with a final dose of between 40-45 e⁻/Å² on the specimen. Data collections were prepared and run automatically using EPU2.X and aberration-free image shift (AFIS).

**Cryo-EM data processing**

Data were processed using RELion3.1-beta (Zivanov et al., 2018), using essentially the same protocol for the three data sets, accounting for the different pixel size. Micrographs were motion corrected using the version of MotionCor2 (Zheng et al., 2017) implemented in RELION, and CTFs were estimated using ctffind4 (Rohou and Grigorieff, 2015). Particles were picked using the Laplacian of Gaussian (LoG) picker, and poor, damaged, or junk particles were removed by serial rounds of 2D classifications with particles down-sampled to 4 Å/pixel. 3D classification was carried out in C1 with an initial model generated from a previous Cx26 cryo-EM reconstruction (unpublished data) with a low-pass filter of 30 Å. Multiple rounds of 3D classification in C1 resulted in 4 very similar classes. Exhaustive rounds of refinement, CTF refinement and polishing in RELion with unbinned particles were used to improve the resolution of the Coulomb shells until no further improvement was gained. Trials were made treating the classes individually or pooling them in C1, C2, C3, C6 and D6 symmetry. While variations were noted in the associated maps, it was not possible to identify different structural features. The resolution was estimated based on the gold standard Fourier Shell Coefficient (FSC) criterion (Rosenthal and Henderson, 2003; Scheres, 2012) with a soft solvent mask. Local Resolution estimation was carried out in ResMap (Kucukelbir et al., 2014).

**Variability analysis in cryoSPARC**

The refined particles associated with each data set were downsampled to 1.586 Å/pixel and imported into cryoSPARC (Punjani et al., 2017). An ab-initio reconstruction was made in C1 and the particles refined in cryoSPARC using D6 symmetry to give maps with an estimated resolution of 3.24 Å. Particle expansion was carried out with D6 symmetry. These were then subjected to variability analysis in cryoSPARC with a filter resolution of 4.5 Å and a mask over the six subunits of one of the two docked hemichannels that form the gap junction. The results were displayed as a simple movie of 20 frames shown in Chimera (Goddard et al., 2007; Pettersen et al., 2004). Analysis was also carried out with masks covering single or neighbouring subunits, but these did not appear to give any advantage over the hemichannel mask.
Particle subtraction and masked classification in relion

Hemichannel classification with imposed C6 symmetry

A mask was created in Chimera (Goddard et al., 2007; Pettersen et al., 2004) based on the best-defined region of the D6-refined dodecameric model, and a soft edge added in relion_mask_create. D6-refined particle sets for each dataset were further refined with this mask. These particles were then symmetry expanded in D1. A mask was created in Chimera (Goddard et al., 2007; Pettersen et al., 2004) based on the least-defined region of the D6-refined dodecameric model, and a soft edge added in relion_mask_create. This mask was then used in particle subtraction and 3D classification without image alignment. The top 3 classes from each dataset were selected, and the original particles recovered. These were refined with C6 symmetry imposed, using a hemichannel mask and limited initial angular sampling. The resolution of the maps was estimated by gold standard Fourier Shell Correlations.

Two-subunit classification without imposed symmetry

For consistency among the three data sets, which contain different numbers of particles, 125,000 particles were randomly selected from each of the 55 mmHg and 90 mmHg PCO2 D6 refined particle sets to match the particle numbers in the 20 mmHg PCO2 data set. These were then symmetry expanded in D6. A mask was created in Chimera (Goddard et al., 2007; Pettersen et al., 2004) based on two neighbouring subunits of the refined model and a soft edge added in relion_mask_create. This was then used in particle subtraction and 3D classification without image alignment. Reconstructions were created with half subsets of each class and the resolution was estimate with relion_postprocess.

Model building

Initial model building was carried out in Coot (Emsley and Cowtan, 2004) with the D6 reconstructions of the 55 mmHg CO2 data with an initial model based on the crystal structure of Cx26 (Maeda et al., 2009). Residues at the C-terminus and the cytoplasmic loop between 102 and 128 could not be observed and were not included in the model. While the density clearly showed the position of the N-terminal helix the side chains were not well defined and modelling of the first residues is ambiguous. For this reason, we have omitted the first three residues in the final PDB file. Real space refinement in Phenix (Adams et al., 2010) was carried out with NCS constraints. Water molecules were also added during refinement with further curation of the water molecules added in Coot. Lipids or detergents were clearly visible in the density, though the exact nature of the head groups is ambiguous. The head-groups have been omitted in the final structure. Refinement of the 20 mmHg and 90 mmHg structures was carried out starting from a partially refined structure of the 55 mmHg CO2 structure using a similar protocol. For the structure derived from the second class of the C6 classification of the 90 mmHg PCO2 structure residues were tentatively modelled for the N-terminal helix starting from Asp2.

Structural analysis

All structural images shown in this paper were generated in Chimera (Goddard et al., 2007; Pettersen et al., 2004) or Chimera X (Goddard et al., 2018) except for Figure 1D, which was created in PyMol (Delano, 2002). Superpositions were carried out in Chimera such that only matching Cα pairs within 2 Å after superposition were included in the matrix calculation.

Revaluation of crystallographic data from previous publications

The structure derived from the 90 mmHg data set was refined against the data sets deriving associated with the structures from Maeda et al. (PDB:2ZW3) (Maeda et al., 2009) and Bennet et al. (Bennett et al., 2016) (PDB: 5er7, PDB 5era) using both Refmac5 (Murshudov et al., 2011) and Phenix.refine (Afonine et al., 2012). The resulting maps were examined in Coot (Emsley and Cowtan, 2004).

QUANTIFICATION AND STATISTICAL ANALYSIS

Cryo-EM reconstructions were performed as described in the method details using cryoSPARC (v2.14.2) and Relion 3.1. Phenix (1.18.2) was used for refinement. Statistics of data collection and refinement are given in Tables S1–S3.