CO₂ directly modulates connxin 26 by formation of carbamate bridges between subunits

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Abstract Homeostatic regulation of the partial pressure of CO₂ (PCO₂) is vital for life. Sensing of pH has been proposed as a sufficient proxy for determination of PCO₂ and direct CO₂-sensing largely discounted. Here we show that connxin 26 (Cx26) hemichannels, causally linked to respiratory chemosensitivity, are directly modulated by CO₂. A ‘carbamylation motif’, present in CO₂-sensitive connexins (Cx26, Cx30, Cx32) but absent from a CO₂-insensitive connexin (Cx31), comprises Lys125 and four further amino acids that orient Lys125 towards Arg104 of the adjacent subunit of the connexin hexamer. Introducing the carbamylation motif into Cx31 created a mutant hemichannel (mCx31) that was opened by increases in PCO₂ and four further amino acids that orient Lys125 towards Arg104 of the adjacent subunit of the connexin hexamer. Introducing the carbamylation motif into Cx31 created a mutant hemichannel (mCx31) that was opened by increases in PCO₂. Mutation of the carbamylation motif in Cx26 and mCx31 destroyed CO₂ sensitivity. Course-grained computational modelling of Cx26 demonstrated that the proposed carbamate bridge between Lys125 and Arg104 biases the hemichannel to the open state. Carbamylation of Cx26 introduces a new transduction principle for physiological sensing of CO₂.

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Introduction

CO₂ is the unavoidable by-product of cellular metabolism. Humans produce approximately 20 moles of CO₂ per day (Marshall and Bangert, 2008). The dissolved CO₂ can readily combine with water, aided by carbonic anhydrase, to form H₂CO₃, which dissociates rapidly to H⁺ and HCO₃⁻. In any physiological solution therefore, the partial pressure of CO₂ (PCO₂) will be in equilibrium with, and inescapably related to, the pH and the concentration of HCO₃⁻ of that solution. Regulation of PCO₂ is thus a vital homeostatic function that is linked to acid-base balance.

As might be expected, chemosensory reflexes regulate the frequency and depth of breathing to ensure homeostatic control of blood gases. The field of respiratory chemosensitivity has been dominated by ‘reaction theory’ which posits that pH is a sufficient signal for detection of changes in PCO₂ (Loeschcke, 1982). Many investigators therefore equate pH-sensing with CO₂-sensing. There are several areas of the medulla oblongata which contain neurons that respond to changes in pH/CO₂, especially near the highly vascularised ventral surface. For example a population of neurons highly sensitive to pH/CO₂ have been described in the retrotapezoid nucleus (RTN) (Mulkey et al., 2004, 2006; Guyenet et al., 2008) and the medullary raphé nucleus (Richerson, 2004; Ray et al., 2011). Despite the acceptance of pH-sensing as the predominant mechanism by which PCO₂ is measured, there is substantial evidence for an additional and independent effect of molecular CO₂ (Eldridge et al., 1985; Shams, 1985; Huckstepp and Dale, 2011). For example, if pH is carefully controlled at the medullary surface, an increase in PCO₂ at constant pH will still enhance breathing by as much as a pH change at constant PCO₂ (Shams, 1985). We have recently shown that connxin 26 (Cx26)
hemichannels, open in response to increases in PCO$_2$ at constant extracellular pH and are an important conduit for the CO$_2$-dependent, as opposed to pH-dependent, release of ATP (Huckstepp et al., 2010a). Cx26 hemichannels contribute to the chemosensory control of breathing (Huckstepp et al., 2010b; Wenker et al., 2012). Hemichannels of two closely related connexins, Cx30 and Cx32, also exhibited CO$_2$-sensitive opening (Huckstepp et al., 2010a). Despite this evidence, widespread acceptance of direct sensing of CO$_2$ requires a detailed molecular explanation of any putative transduction system.

Recent research into structures called connexin hemichannels has challenged this view. Researchers found that when pH levels were held constant, increasing the level of CO$_2$ caused the structures to open up, suggesting that CO$_2$ could be directly detected by the hemichannels. Each hemichannel contains six connexin subunits, but the details of how the CO$_2$ molecules interact with the individual connexin subunits to open up the hemichannels remained mysterious.

Now Meigh et al. show that CO$_2$ molecules bind to a specific amino acid (lysine) at a particular place (residue 125) in one of the connexin subunits to form a carbamate group. This group then interacts with the amino acid (arginine) at residue 104 in a neighbouring connexin subunit to form a carbamate bridge between the two subunits. This leads to structural changes that cause the gap junction hemichannels to open and release signals that can activate other cells. Since connexin hemichannels are found throughout the human body, these results suggest that CO$_2$ might act as a signalling molecule in processes as diverse as the control of blood flow, breathing, hearing and reproduction.

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Results
We have previously demonstrated that Cx26, and two related β connexins, Cx30 and Cx32, open when exposed to modest increases in PCO$_2$ at constant pH (Huckstepp et al., 2010a). This previous study demonstrated, in inside-out and outside-out excised membrane patches at a transmembrane potential of −40 mV, that Cx26 hemichannel gating respectively increased and decreased in response to increases and decreases of PCO$_2$. To reconfirm our previous findings that Cx26, and not some other hemichannel senses CO$_2$ (Huckstepp et al., 2010a), we demonstrated that the CO$_2$-dependent dye loading of HeLa cells expressing Cx26 was blocked by 100 µM carbeneoxide, but unaffected by 1 mM probenecid, a blocker of pannexin-1, (Silverman et al., 2008), and 20 µM ruthenium red, a blocker of calhm1 (Taruno et al., 2013), (Figure 1—figure supplement 1). Parental HeLa cells do not exhibit CO$_2$-dependent dye loading demonstrating that the heterologous expression of Cx26 is essential for this function (Huckstepp et al., 2010a) (Figure 3—figure supplement 1).
The extent of CO₂ chemosensitivity within the β connexins

To document the extent to which this sensitivity to CO₂ is limited within the β connexin family (Figure 1F), and to form the basis of a bioinformatic comparison to identify possible CO₂ binding motifs, we investigated whether another β connexin, Cx31, might also be sensitive to CO₂. We expressed, in HeLa cells, either rat Cx31 or rat Cx26 tagged at the C-terminal with mCherry and used a previously described dye loading assay (Huckstepp et al., 2010a) to test whether the cells could load with carboxyfluorescein (CBF) in a CO₂-dependent manner. As expected from our previous work, HeLa cells expressing the Cx26 readily loaded with CBF when exposed to this dye in the presence of elevated PCO₂ (55 mmHg, at pH 7.5, Figure 1A,B). However, HeLa cells expressing Cx31 failed to dye load in a CO₂-dependent manner (Figure 1A,B). As the connexins were tagged with mCherry, we could verify the presence of fluorescent puncta in both the Cx26 and Cx31 expressing HeLa cells (Figure 1—figure supplement 2). To check for the existence of functional hemichannels in the Cx31-expressing HeLa cells, we removed extracellular Ca²⁺ as a positive control. This manipulation will open all types of connexin hemichannel. Parental HeLa cells do not load with dye when Ca²⁺ is removed from the medium (Figure 3—figure supplement 1); they therefore possess virtually no endogenous hemichannels. The removal of extracellular Ca²⁺ readily caused loading of CBF into the Cx31-expressing HeLa cells (Figure 1A, inset), demonstrating the presence of functional Cx31 hemichannels.

Identification of a carbamylation motif in CO₂ sensitive β connexins

The CO₂-sensitivity in the β connexins therefore appears to be limited to the three closely related connexins, Cx26, Cx30 and Cx32, and Cx31 has no sensitivity to increases in PCO₂ (Figure 1F). We hypothesized that CO₂ carbamylated a lysine residue in Cx26 to induce conformational change and hence opening of the hemichannel. We therefore compared the sequences of the three CO₂-sensitive connexins to Cx31 to identify a lysine present in all three CO₂ sensitive connexins but absent from Cx31 (Figure 1C). This analysis revealed K125 and four further amino acids as forming a motif that was absent from Cx31. The existing crystal structure for Cx26 (Maeda et al., 2009), shows that K125 is at the end of an alpha helix and that the sequence KVREI (‘carbamylation motif’) orients K125 towards the end of an alpha helix and that the sequence KVREI (‘carbamylation motif’) orients K125 towards.

Insertion of the carbamylation motif into Cx31 creates a CO₂-sensitive mutant hemichannel

Our analysis predicts that if we were to introduce the putative carbamylation-motif of Cx26 into Cx31, the resulting mutant Cx31 (mCx31) should be sensitive to CO₂ as the lysine introduced into the sequence should be able to form a salt bridge with the native residue K104 in mCx31 once carbamylated (Figure 1C–E and 2A). We therefore made mCx31 in which the motif TQKVREI was introduced in place of K123H124. Unlike lysine, the arginine side chain cannot be carbamylated as its pKa (12.5) is much higher than that of lysine (10.5), therefore this variant should have no sensitivity to CO₂. mCx31K125R did indeed lack sensitivity to CO₂ (Figure 3A–C, Figure 3—source data 1). This was not because the mutant channel failed to assemble correctly, as the positive control of zero Ca²⁺ dye loading demonstrated the presence of functional hemichannels (Figure 3A, Figure 3—figure supplement 1). Next we investigated the relevant residues in Cx26 itself. The carbamate bridge that we propose must involve K125 (being carbamylated) and R104 (forming the salt bridge with the carbamylated motif). To demonstrate that K125 is the key residue for interaction with CO₂, we made mCx31K125R, by inserting TQRVREI into Cx31 in place of K123H124. Unlike lysine, the arginine side chain cannot be carbamylated by CO₂ as its pKa (12.5) is much higher than that of lysine (10.5), therefore this variant should have no sensitivity to CO₂. mCx31K125R did indeed lack sensitivity to CO₂ (Figure 3A–C, Figure 3—source data 1).
Figure 1. Identification of the motif in Cx26 that imparts CO$_2$ sensitivity. (A) Dye loading assay demonstrates CO$_2$-dependent loading of carboxyfluorescein into HeLa cells expressing Cx26, but not into those expressing Cx31. The inset in Cx31 shows that these hemichannels are expressed and functional in the membrane by utilizing a zero Ca$^{2+}$ stimulus to open them and allow dye loading. Scale bars 40 µm. (B) Cumulative probability plots of pixel intensity in the control and following exposure to PCO$_2$ of 55 mmHg. Each curve is comprised of at least 40 cells. (C) Sequences (from mouse) for Cx26, 30, 32 and 31 to show K125 and four following amino acids that are present in Cx26, Cx30 and Cx32, but absent from Cx31. R104 in Cx26 and 30, and K104 in Cx32 and Cx31 are also highlighted. Accession numbers: Cx26, NP_032151; Cx30, AAH13811; Cx32, AAH26833; and Cx31, NP_001153484. (D) The structure of Cx26 drawn from the 2zw3 PDB file, cytoplasmic face of the channel upwards. On each subunit K125 and R104 are drawn. (E) Detail from the structure of Cx26 (dashed square) showing the orientation of K125 (red) towards R104 (dark grey). The short distance between the two amino acid side chains suggests that this gap could be bridged by carbamylation by CO$_2$ of K125 and a subsequent salt bridge with R104. (F) Phylogenetic tree showing relationship between Cx26 and other β connexins. The three CO$_2$ sensitive connexins are very closely related to each other while Cx31 is more distant.

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Figure 1. Continued on next page
lysine). We therefore made mutations that individually disrupted both of these functions: K125R to prevent carbamylation, and R104A to disrupt formation of the salt bridge. Neither Cx26K125R nor Cx26R104A exhibited sensitivity to CO$_2$ sensitivity. Nevertheless the positive controls demonstrated the presence of functional mutant hemichannels in the expressing HeLa cells (Figure 3A, Figure 3—figure supplement 1).

**Engineering an analogue of the carbamylated lysine into Cx26 makes it constitutively open**

To test further our prediction that the carbamylated K125 forms a salt bridge with R104, we made the mutation K125E in Cx26. The insertion of glutamate in place of the lysine has the potential to act as an analogue of the carbamylated K125. If our hypothesis is correct, this mutated channel should be constitutively open, as the carboxyl group of the E125 should be able to form a salt bridge with R104. We found that HeLa cells expressing Cx26K125E readily loaded with dye under control conditions and exhibited no sensitivity to CO$_2$ (Figure 4). The Cx26K125E-expressing HeLa cells showed much greater loading under control conditions than parental HeLa cells (Figure 4, Figure 4—source data 1). To further confirm that the constitutive dye loading occurred via the misexpressed connexin, we
demonstrated that carbenoxolone (100 µM) completely blocked CO₂-dependent dye loading in HeLa cells expressing Cx26<sub>K125E</sub> (Figure 4, Figure 4—source data 1).

Reasoning that if bridge formation between subunits was key to opening the hemichannel, we also made the further mutation R104E. In the mutated channel E104 has the potential to form a salt bridge in the reverse direction with K125 and we predicted that if this were to happen such a mutant hemichannel should also be constitutively open. We found that HeLa cells expressing Cx26<sub>R104E</sub> did indeed load with dye under control conditions and that this enhanced dye loading was blocked with carbenoxolone (Figure 5, Figure 5—source data 1).

**Elastic network model of Cx26 shows that carbamylation leads to hemichannel opening**

Although our experimental data point to the importance of carbamylation of K125 and the formation of a salt bridge to R104 in the adjacent subunit, it is not clear how this would lead to opening of the
Figure 4. Engineering an analogue of the carbamylated lysine residue, Cx26\(_{K125E}\), creates a constitutively open hemichannel that no longer responds to CO\(_2\). (A) HeLa cells expressing Cx26\(_{K125E}\) readily load with dye under control conditions. Increasing the PCO\(_2\) does not give a further increase in dye loading. This dye loading was blocked by 100 µM carbenoxolone (Carb), indicating that it occurred through the heterologously expressed connexin. Scale bar 40 µm. (B) Cumulative probability plots comparing the median pixel intensities of at least 40 cells per experiment and five independent repetitions for the control, hypercapnic and carbenoxolone treatments with that of parental HeLa cells (four independent repetitions). (C) Summary data showing the median of the median pixel intensity for the three conditions for Cx26\(_{K125E}\) and the background loading for parental HeLa cells. Pairwise comparisons by the Mann-Whitney U-test; KW Kruskall-Wallis Anova. Neither the difference between control and CO\(_2\) nor the difference between Cx26\(_{K125E}\) treated with carbenoxolone and parental HeLa cells is significant.

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The following source data are available for figure 4:

Source data 1. Median pixel intensity values for histogram in Figure 4C and statistical analysis: Kruskal-Wallis anova and pairwise Mann-Whitney tests.

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Addition of springs representing CO₂-binding to the ENM restricted the closing motions of the hemichannel and thus connexin 26 was maintained in the open state (Video 2). We examined the overlap in the ordering of the modes in the unbound and CO₂ bound states to gain insight into how this occurs. A significant reordering of the lowest frequency modes to higher frequencies was observed in the presence of CO₂ rather than the removal of any modes from the total protein motion (Figure 6B). Mode 1, the lowest frequency mode that represents the opening/closing mode, represented about 40% of the total protein motion in the absence of CO₂. In the presence of CO₂ this closing mode is reordered through a change in its frequency as mode 9, which accounts for only about 2% of the total motion (Figure 6B,C). CO₂ therefore opens Cx26 through a reordering of the normal modes of global protein motion such that the normal closing motion of Cx26 no longer significantly contributes to the total motion of the hemichannel. We infer from this that the carbamate bridge formed between Cx26 monomers represents a constraining force that hinders hemichannel closure.

**Discussion**

Evidence from six different experimental tests supports our hypothesis that CO₂ forms a carbamate bridge between K125 and R104 on the adjacent subunit to open the Cx26 hemichannel. Firstly, we demonstrated the sufficiency of the carbamylation motif to confer CO₂ sensitivity by inserting it into a CO₂-insensitive...
connexin, Cx31. Secondly, we showed that K125 of the carbamylation motif was essential for this motif to confer CO₂ sensitivity on Cx31. Thirdly and fourthly, we demonstrated that the mutations K125R and R104A in Cx26 (to prevent carbamate bridging at either end of the bridge) destroyed the CO₂ sensitivity of this connexin. Fifthly, by exploiting glutamate as an analogue of the carbamylated K125 (in Cx26^{K125E}), we demonstrated a gain of function—Cx26^{K125E} was constitutively open, yet had lost sensitivity to CO₂. Finally, we further tested the bridging concept by demonstrating that the bridge is in effect bidirectional: the mutated hemichannel Cx26^{R104E}, in which E104 can bridge to K125 in the reverse direction, was also constitutively open, but had no sensitivity to CO₂.

Although we have not directly demonstrated CO₂ binding to Cx26, our extensive testing of this hypothesis through selective mutations leads us to conclude that CO₂ interacts with Cx26 directly and that no other protein is required for CO₂ sensitivity. This interaction is most probably via carbamylation of K125. Interestingly, the mutations Cx26^{K125E} and Cx26^{K125R} can be considered respectively as open-state and closed-state analogues of the wild type channel. Collectively, our data strongly suggests that CO₂ binds to the intracellular surface of Cx26 and must therefore cross the membrane to reach this site. This could occur either direct diffusion through the membrane bilayer, potentially via Cx26 itself, or via other CO₂ permeable channels (Boron et al., 2011). Amongst its many other functions, Cx26 can therefore be regarded as a receptor for CO₂. Interestingly, this mechanism of modulation applies to both Cx30 and Cx32, which can both potentially form a carbamate bridge at equivalent residues to Cx26. In the case of Cx32 this would involve bridging to K104 rather than R104 (in Cx26 and 30). Cx26 can co-assemble with both Cx30 and Cx32 to form heteromeric hemichannels (Forge et al., 2003; Yum et al., 2007). Our structural studies predict that, as Cx30 and Cx32 have K125 and either R104 or K104, carbamate bridges could form in such heteromeric hemichannels and that they should also therefore be CO₂-sensitive.

Carbamylation involves formation of a labile covalent bond between the carbon of CO₂ and a primary amine. For this to occur the amine must be in a restricted hydration space so that it is not protonated. Some examples of physiologically significant carbamylation are known. The carbamylation of the N-terminal amines of haemoglobin contributes to the Bohr effect (Kilmartin and Rossi-Bernardi, 1971), whereby the affinity of haemoglobin for O₂ is reduced in the presence of elevated CO₂. However in mammalian systems

Figure 6. Elastic network model (ENM) of Cx26 demonstrating that CO₂ binding restricts the motion of the hemichannel and biases it to the open state. (A) Left, diagram of Cx26 from the 2zw3 structure, indicating the ENM (black lines) superimposed on the tertiary structure of Cx26 and showing the position of the hookean springs (white lines) introduced to simulate binding of CO₂ to K125 and bridging to R104. Right, ENM of Cx26 seen end on from the cytoplasmic side of the channel showing the six springs (white lines) that represent CO₂ binding. (B) Frequency modes of channel motion plotted for CO₂ bound against those without CO₂ bound. The grey scale on the right indicates the similarity of the modes between the CO₂ bound and unbound states. Note that there is a high degree of similarity between most modes in the bound and unbound state, indicating that binding of CO₂ reorders the modes of motion. In the graph, the modes that fall on the dotted line (x = y) have not changed between the two states. Mode 1 without CO₂ bound (closing of hemichannel) moves to Mode 9 with CO₂ bound (dashed upward arrow) indicating that it contributes much less to the total channel motions when CO₂ is bound. Most of the other modes fall below the dotted line, indicating that they become more frequent when CO₂ is bound. (C) Vectors indicating the Mode 1 motions of the α helices without CO₂ bound (left) and with CO₂ bound (right). The binding of CO₂ and creation of the carbamylation bridge between subunits greatly restricts hemichannel motion.

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no other examples of carbamylation by CO\(_2\) have been described. In C3 photosynthetic plants, the enzyme RuBisCo, that participates in the Calvin cycle and carbon fixation is activated by carbamylation of a specific lysine residue (Lundqvist and Schneider, 1991). Several microbial enzymes are also carbamylated (Maveyraud et al., 2000; Golemi et al., 2001; Young et al., 2008).

Despite this precedent, the functional significance of CO\(_2\)-carbamylation and its potential as a transduction principle for the measurement of CO\(_2\) has been almost completely overlooked in vertebrate physiology. The mechanism of formation of a salt bridge between a carbamylated lysine and an appropriately oriented arginine on the neighbouring subunit is a unique mechanism for modulation of an ion channel and establishes carbamylation as a mechanistic basis for the direct signalling of PCO\(_2\) in mammalian physiology. This carbamylation of a lysine to transduce the concentration of CO\(_2\) into a biological signal is somewhat equivalent to the nitrosylation of a cysteine residue by NO/nitrite. It establishes a CO\(_2\)-dependent signalling paradigm in which the concentration of CO\(_2\) is signalled by ATP release via Cx26 from the chemosensory cell and consequent activation of neighbouring cells, or potentially by a Ca\(^{2+}\) influx through the Cx26 hemichannel (Fiori et al., 2012) to initiate a Ca\(^{2+}\) wave within the chemosensory cell itself and further Ca\(^{2+}\)-dependent signalling processes.

Materials and methods

Hemichannel expression and mutagenesis

All connexin genes except Cx26\(^{R104A}\), Cx26\(^{K125E}\) and Cx26\(^{R104E}\) site directed mutagenesis was performed using Quikchange II site directed mutagenesis kit. All wild type and mutant genes were sequenced to verify that the correct sequence was present. Hela cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) (Sigma-Aldrich Company Ltd, Gillingham, UK), 10% FCS (Biosera Europe, Labtech International Ltd, Uckfield, UK), 1:1000 pen/strep and supplemented with 3 mM CaCl\(_2\). Cells were grown in a humidified 5% CO\(_2\) incubator at 37°C. The connexin proteins were expressed via transient transfection. Cells were plated in six-well plates at 1 x 10\(^5\) cells per well for Cx26 and its mutants and 5 x 10\(^4\) cells per well for Cx31 and its mutants. Following the GeneJuice transfection reagent (Merck-Millipore UK, Merck KGaA, Darmstadt, Germany) user protocol, cells were transfected with 1 µg of the appropriate DNA. Experiments were performed when the connexin proteins had reached the cell membrane. This was found to be approximately 2 days for Cx26 and its mutants and approximately 3 days for Cx31 and its mutants.

Solutions used

Standard artificial cerebrospinal fluid (aCSF, normocapnic)

124 mM NaCl, 3 mM KCl, 2 mM CaCl\(_2\), 26 mM NaHCO\(_3\), 1.25 mM NaH\(_2\)PO\(_4\), 1 mM MgSO\(_4\), 10 mM D-glucose saturated with 95% O\(_2\)/5% CO\(_2\), pH 7.5, PCO\(_2\) 35 mmHg.
50 mM HCO$_3^-$ aCSF (isohydric hypercapnic)
100 mM NaCl, 3 mM KCl, 2 mM CaCl$_2$, 50 mM NaHCO$_3$, 1.25 mM NaH$_2$PO$_4$, 1 mM MgSO$_4$, 10 mM D-glucose, saturated with 9% CO$_2$ (with the balance being O$_2$) to give a pH of 7.5 and a PCO$_2$ of 55 mmHg respectively.

**Dye loading assay and image analysis**

Connexin expressing HeLa cells were plated on cover slips. A coverslip was placed in a small flow chamber and the cells were exposed to either: control aCSF with 200 µM carboxyfluorescein for 10 min; isohydric hypercapnic aCSF with 200 µM carboxyfluorescein for 10 min; or zero Ca$^{2+}$, 1 mM EGTA-containing aCSF plus 200 µM carboxyfluorescein for 10 min. This was followed by control aCSF plus 200 µM carboxyfluorescein for 5 min and then thorough washing for 30 min with control aCSF. These protocols are summarized in *Figure 7*.

The cells were then imaged by epifluorescence (Scientifica Slice Scope (Scientifica Ltd, Uckfield, UK), Cairn Research OptoLED illumination (Cairn Research Limited, Faversham, UK), 60x water Olympus immersion objective, NA 1.0 (Scientifica), Hamamatsu ImageEM EMCCD camera (Hamamatsu Photonics K.K., Japan), Metafluor software (Cairn Research)). Using ImageJ, the extent of dye loading was measured by drawing a region of interest (ROI) around individual cells and calculating the mean pixel intensity for the ROI. The mean pixel intensity of the background fluorescence was also measured in a representative ROI, and this value was subtracted from the measures obtained from the cells. All of the images displayed in the figures reflect this procedure in that the mean intensity of the pixels in a representative background ROI has been subtracted from every pixel of the image. At least 40 cells were measured in each condition, and the mean pixel intensities plotted as cumulative probability distributions.

For the dye loading experiments, the median pixel intensities of the control and CO$_2$ dye loading conditions (minimum of five independent repetitions) were compared by a Kruskal Wallace ANOVA and pairwise comparisons by the Mann-Whitney test. The false discovery rate procedure (Curran-Everett, 2000) was used to determine whether the multiple pairwise comparisons remained significant.

**Patch clamp recordings**

Cover slips containing non-confluent cells were placed into a perfusion chamber at 28°C in sterile filtered standard aCSF. Standard patch clamp techniques were used to make whole-cell recordings. The intracellular fluid in the patch pipette contained: K-gluconate 120 mM, CsCl 10 mM, TEACl 10 mM, EGTA 10 mM, ATP 3 mM, MgCl$_2$ 1 mM, CaCl$_2$ 1 mM, sterile filtered, pH adjusted to 7.2 with KOH. All whole-cell recordings were performed at a holding potential of −40 mV with regular steps of 5 s to −50 mV to assess whole-cell conductance.
Elastic network model–course-grained simulations

Elastic network model (ENM) simulations were performed based on its regular implementation using pdb file 2ZW3, where all the Cα atoms in the protein within a given cut-off radius are joined with simple Hookean spring (Tirion, 1996; Rodgers et al., 2013a). The spring constants were set to a constant value of 1 kcal mol$^{-1}$ Å$^{-2}$ with a cut-off radius of 8 Å. The presence of CO$_2$ molecules were represented in the ENM by the inclusion of an additional Hookean spring between residues K125 and R104 of each set of neighbouring monomers (Rodgers et al., 2013b). The first six modes, that is the lowest frequency modes, represent the solid body translational and rotational motions of the protein and are thus ignored from the analysis.

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Additional files

Supplementary files

• Supplementary file 1. Conductance changes source data. Raw values for whole cell conductance changes (nS) in response to an isohydric CO$_2$ challenge (PCO$_2$ 55 mmHg) in Cx31 and mCx31 expressing HeLa cells.

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Major dataset

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| Author(s) | Year | Dataset title | Dataset ID and/or URL | Database, license, and accessibility information |
|-----------|------|---------------|-----------------------|-----------------------------------------------|
| Maeda S, Nakagawa S, Suga M, Yamashita E, Oshima A, Fujiyoshi Y, Tsukihara T | 2009 | Structure of the connexin 26 gap junction channel at 3.5 Å resolution | 2ZW3; http://www.rcsb.org/pdb/explore/explore.do?structureId=2ZW3 | Publicly available at RCSB Protein Data Bank (http://www.rcsb.org). |

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