Structural determinants of CO$_2$-sensitivity in the β connexin family suggested by evolutionary analysis

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A subclade of connexins comprising Cx26, Cx30, and Cx32 are directly sensitive to CO$_2$. CO$_2$ binds to a carbamylation motif present in these connexins and causes their hemichannels to open. Cx26 may contribute to CO$_2$-dependent regulation of breathing in mammals. Here, we show that the carbamylation motif occurs in a wide range of non-mammalian vertebrates and was likely present in the ancestor of all gnathostomes. While the carbamylation motif is essential for connexin CO$_2$-sensitivity, it is not sufficient. In Cx26 of amphibia and lungfish, an extended C-terminal tail prevents CO$_2$-evoked hemichannel opening despite the presence of the motif. Although Cx32 has a long C-terminal tail, Cx32 hemichannels open to CO$_2$ because the tail is conformationally restricted by the presence of proline residues. The loss of the C-terminal tail of Cx26 in amniotes was an evolutionary innovation that created a connexin hemichannel with CO$_2$-sensing properties suitable for the regulation of breathing.

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There are 20 connexin genes in the human genome. This large number of variants in the connexin gene family, implies diversity of cellular and physiological function which may depend on the precise properties of the different connexins. Connexins form gap junctions, which comprise two hexameric hemichannels in the membranes of adjacent cells docked together to form a dodecameric complex. Gap junctions are aqueous pores that permit ion flow and transfer of small molecules between the coupled cells. In addition to this canonical function of coupling cells, the hexameric hemichannels can have an independent function by acting as large conductance plasma membrane channels. Hemichannels are a particularly important mechanism for the release of ATP into the extracellular space. We have discovered that the β connexins, Cx26, Cx30 and Cx32 are modulated by CO2. Hemichannels of each of these connexins can be opened by CO2. In the case of Cx26, this direct CO2-gated hemichannel opening, and subsequent release of ATP, mediates an important part of respiratory chemosensitivity. There are other important molecules that may also contribute to respiratory chemosensitivity — these include pH-sensitive channels and receptors such as the TASK channels and GPR41. The physiological significance of the CO2 sensitivity of Cx30 and Cx32 has not yet been elucidated. We have analyzed the structural basis of the CO2 dependent modulation of Cx26 hemichannels in detail, and have discovered that it most likely depends upon the carbamylation of Lys125, and formation of a salt bridge from the carbamylated lysine to Arg104 of the neighbouring subunit (a "carbamate bridge"). This carbamate bridge increases the time that the hemichannel spends in the open configuration. Our structural studies have allowed us to define a "carbamylation motif" that is present in CO2-sensitive connexins, but absent from those that are insensitive to CO2.

Very recently we have discovered that there are two actions of CO2 mediated via the carbamylation motif. Whereas CO2 opens hexameric hemichannels, it has the opposite effect and causes closure of the gap junctions. Mutational analysis shows that this closing effect of CO2 on the Cx26 gap junction is most likely mediated by its binding to the same residues that effect hemichannel opening. Both effects of CO2 are likely therefore to be mediated via the carbamylation motif.

Several authors have examined the evolution of the β connexin family. In this paper we use our insights about the nature of the carbamylation motif to further refine our understanding of the phylogenetic occurrence of this motif, and hence CO2-sensitivity, in the β connexin family. This approach has given us new insight into the structural determinants of the CO2 sensitivity of both gap junctions and hemichannels and has shown that the carbamylation motif was present in the ancestor of all gnathostomes. Interestingly, it is a common feature of the amniote species tested that their Cx26 hemichannels lack an extended C-terminal tail and are consequently sensitive to CO2. This suggests that the common ancestor of all extant amniotes had already evolved CO2-sensitive Cx26 hemichannels.

### Results

**Molecular phylogenetic and microsyntenic analysis.** The amino acid sequences from fifty-three β connexin family members, from 24 vertebrate species, were used for molecular phylogenetic analysis (Fig. 1 and Supplementary Table 1). Additional species that we inspected (and support our conclusions) but did not include in the phylogenetic analysis of Fig. 1 are listed in Supplementary Table 2, and shown in Supplementary Fig. 1. The resulting tree topology exhibited two main clades named A and B, supported by high values of posterior probability. Clade A consisted of the sequence Cx27.5 of the jawless Petromyzon marinus and a cluster comprising the Cx26, Cx30 and Cx30.3 sequences of gnathostomes. These sequences are distributed further into two subgroups: one containing the sequences for Cx26 and Cx30 of reptiles, birds, and mammals; the other containing sequences belonging to elephant shark, actinopterygians, coelacanth, lungfish, and amphibians. The analysis was not able to establish the correct orthology and paralogy relations in the clade A. The microsyntenic analysis of the chromosome region harbouring these genes showed a conserved pattern of flanking genes from lamprey to mammals (Fig. 2a). This indicates that the common craniate ancestor of agnathans and gnathostomes already had this genomic arrangement. Furthermore, this analysis reveals that in amniotes two genes are located between Cyl1 and Gja3, corresponding to Cx30 and Cx26, respectively (Blue box, Fig. 2a). This result, together with the phylogenetic analysis, supports the hypothesis proposed by Abascal and Zaroya that these connexin genes are derived from a duplication event that occurred in the common amniote ancestor of reptiles, birds, and mammals. However, for the clade containing Cx30 and Cx26 sequences of amniotes, our phylogenetic analysis does not allow the orthology and paralogy to be ascertained probably because mechanisms such as gene conversion, common in tandemly arranged genes, may have hidden the real relationships. In the amphibian Xenopus tropicalis, only one gene is located between Cyl1 and Gja3. This gene is annotated as Cx26 but is probably orthologous to the ancestral gene from which Cx26 and Cx30 of amniotes originated. The orthology relation documented for X. tropicalis can be extended to all sequences of non-amniote organisms here analyzed. The presence of more genes in this chromosome region, as for example in coelacanth, is due to lineage-specific duplication. Thus, the sequences of amphibians and lungfish are most probably more closely related evolutionarily to the ancestral gene from which the amniote Cx26 and Cx30 genes arose and are more correctly named Cx26-like. For simplicity, we shall refer to these genes as Cx26.

The clade B included the sequences corresponding to Cx32 of the cartilaginous and actinopterygian fish and of sarcopterygians. Furthermore, the analysis allowed elucidation of the orthology relation between these sequences and those of actinopterygians. This finding is in agreement with the microsyntenic analysis performed between the main vertebrate lineages (Fig. 2b). Indeed, the flanking regions of Cx32 gene shared several genes indicating a common origin. However, the pattern between birds and mammals is more conserved compared to that of actinopterygians probably due to genomic rearrangements. The genes named Cx27.5 and Cx31.7 in teleosts are ohnolog genes thus derived from the lineage-specific genomic duplication event that affected the genome of these organisms.

The carbamylation motif is present in both clades A and B. In clade B, this motif is almost universally present. Interestingly in clade A, there are two substantial branches where it has been lost: non-mammalian, amniote Cx30; and actinopterygian Cx30.3. Interestingly, Cx30.3 may play a similar role in cochlea of fish to that of Cx26 of mammals. The lamprey has a version of the carbamylation motif that is unusual: it possesses Arg104 and Lys125, but there are two prolines in the sequence (Pro124, Pro123). No sequence from any other vertebrate that we have studied possesses this sequence and, given the steric restrictions that the two proline residues would introduce, it is questionable whether Lys125 could be properly oriented to form a salt bridge to Arg104 following carbamylation. The carbamylation motif therefore is definitely present in the ancestor to all gnathostomes, and may have evolved in the agnathans too, albeit in a heavily modified form. Over the two clades, almost all sequences are characterized by a long C-terminal tail. The only notable
Sensitivity of Cx26 hemichannels to CO2. Our sequence comparisons show that the carbamylation motif is present in sarcopterygian fish and tetrapods. As we have already established the CO2-sensitivity of Cx26 hemichannels coded by the mammalian and avian genes, we tested whether the Cx26 hemichannels of reptiles (Chelonia and Gekko), amphibia (Xenopus) and lungfish (Lepidosiren), also exhibit CO2-dependent opening. To evaluate this, we used our well established and validated dye-loading assay to test whether we could detect entry of carboxyfluorescein into HeLa cells expressing these Cx26 genes during a CO2 challenge (Fig. 3). As a positive control to check for functional expression of hemichannels in the membrane we used a zero Ca2+ stimulus which is effective at opening hemichannels by a CO2-independent mechanism, and provides a measure of maximal dye loading to compare to the CO2-dependent dye loading. All four tested Cx26 genes possessed a carbamylation motif, very similar to that of human Cx26 (Fig. 3a). However only HeLa cells expressing the reptilian Cx26 exhibited CO2-dependent dye loading (Fig. 3b, c). Nevertheless HeLa cells expressing all four genes showed dye loading to the zero Ca2+ stimulus demonstrating the presence of functional hemichannels (Fig. 3b, c). We confirmed these results by means of whole-cell patch clamp recordings to demonstrate the presence of a CO2-dependent conductance in HeLa cells expressing Chelonia and Gekko Cx26, but not in HeLa cells expressing Xenopus Cx26 or non-transfected HeLa cells (Fig. 4). Thus, the Cx26 hemichannels from Xenopus and Lepidosiren are not sensitive to CO2. Within the species tested therefore, only Cx26 hemichannels from annmotes possess CO2 sensitivity.

The C-terminal tail controls CO2-sensitivity of Cx26. On inspection of amphibian, coelacanth (Fig. 5a) and lungfish Cx26 amino acid sequences, we noticed that Cx26 of these species possessed a C-terminal tail considerably longer than that of the Cx26 of annmotes. We therefore tested whether removal of this extended C-terminal tail could restore the CO2 sensitivity of Cx26 in these species. We truncated the tail of Xenopus Cx26 and altered the final two residues so they were the same as in mammalian Cx26 to improve trafficking (xtxCx26ΔPV). HeLa cells expressing this truncated Cx26 now demonstrated both CO2-dependent dye loading (Fig. 5b, c) and CO2-dependent conductance changes (Fig. 6). Conversely the addition of the C-terminal tail to human Cx26 (hCx26 + XerCT) effectively abolished the CO2 sensitivity of human Cx26 hemichannels (Fig. 5b, c). Finally, evolution has performed the same manipulation for us: Latimeria (Coelacanth) has 3 different homologues of Cx26 and the C-terminal tail has been truncated to only a few amino acids.

exception to this is Cx26 of annmotes (green box, Fig. 1, see also Supplementary Table 2, Supplementary Fig. 1) in which the C-terminal tail has been truncated to only a few amino acids.

Fig. 1 Molecular phylogenetic tree of Cx26, Cx30 and Cx32 genes. Bayesian Inference on amino acid sequences belonging to the main vertebrate lineages. Numbers close to nodes indicate posterior probability values (>0.95). Amia Calva GEUG01000334.1 (Cx31-like) was used as the outgroup. If the species name is red, the sequence possesses the carbamylation motif. The green box indicates the presence of functional hemichannels. COM
Fig. 2 Microsynteny of the genomic regions harboring Cx26, Cx30 and Cx32 genes in the main vertebrate lineages. a Comparative analysis of microsynteny related to the genomic region harboring Cx26 and Cx30 genes in the main vertebrate lineages. b Comparative analysis of microsynteny related to the genomic regions harboring Cx32 in the main vertebrate lineages. Arrow-heads indicate gene direction and colour help to follow orthology relations. Black arrow-heads indicate genes not showing evident orthology relations. Lines underneath genes indicate syntenic arrangement. Square brackets indicate the presence of multiple genes in that genomic region. The blue box indicates the Cx26 and Cx30 genes present in amniotes.
Fig. 3 Amniote Cx26 hemichannels can be opened by CO₂. a Amniotes, amphibia and lungfish share the CO₂ carbamylation motif (highlighted boxes). b Images of dye loading in response to CO₂ and the zero Ca²⁺ positive control. c Summary data showing that reptile Cx26 hemichannels open to CO₂ but those of *Xenopus* and *Lepidosiren* do not (expressed as the change in pixel intensity from control 35 mmHg PCO₂). Mann Whitney test used in comparisons. Box and whisker plots: box, first and third quartiles; horizontal line, median; whiskers, the furthest point that lies no more than 1.5 times the interquartile range from the median. Each point is the median change in pixel intensity from an independent transfection. Scale bar 20 µm.
human gene (Fig. 5a). We therefore tested whether the truncated Latimeria Cx26 gene encodes CO2-sensitive hemichannels. We found that HeLa cells expressing this truncated gene did indeed exhibit CO2-dependent dye loading (Fig. 5b, c) and CO2-dependent whole-cell conductance changes (Fig. 6). We therefore conclude that the two critical criteria necessary for CO2 sensitivity in Cx26 hemichannels are the lack of an extended C-terminal tail and the presence of the carbamylation motif. This condition is met by the Cx26 of many amniote species.

CO2-sensitivity of Cx32 hemichannels. We have previously shown the Cx32 hemichannels from rat can be opened by CO2, but require higher levels of PCO2 than Cx26. Inspection of the Cx32 amino acid sequence in a variety of actinopterygian and cartilaginous fish revealed the presence of a carbamylation motif very similar to that of human Cx32 (Fig. 7). This implies that this motif was already present in the common ancestor of Chondrichthyes and Osteichthyes. Unlike Cx32, Cx26 in actinopterygian fish does not have the carbamylation motif except in a very few cases for primitive fish (Fig. 1, Supplementary Table 2, Supplementary Fig. 1). Furthermore, Cx32 (like Cx30, which is also CO2-sensitive) possesses a long C-terminal tail, which in Cx26 would abrogate CO2 sensitivity.

We therefore tested whether Danio (Zebrafish) and Rhincodon (whale shark) Cx32 hemichannels were CO2 sensitive (Fig. 7). We found that there was a small amount of CO2 dependent dye loading at a PCO2 of 55 mmHg, and robust dye loading at a PCO2 of 70 mmHg (Fig. 7b, c). Like human Cx32 hemichannels, the fish homologues are sensitive to CO2 but require a substantially higher stimulus than those of amniote Cx26 to open them.

This leads us to the intriguing question of why the extended C-terminal tail in Cx32 does not abrogate the CO2 sensitivity of hemichannels, whereas in Cx26 it does. By inspecting the sequences of Cx32 we noticed that, unlike the amphibian and lungfish Cx26, there were multiple proline residues in the C-terminal tail (Fig. 8a). As prolines will conformationally restrict an unstructured peptide sequence, we hypothesized that the resulting structure could prevent the C-terminal tail of Cx32 interfering with the CO2-dependent opening of the hemichannels.

We therefore mutated all proline residues to glycine (Fig. 8) in the C-terminal tail of human Cx32. This completely removed the sensitivity of Cx32 hemichannels to CO2. We also performed the converse experiment: Cx26 hemichannels of Lepidosiren are not sensitive to CO2. To explore whether introduction of prolines into the C-terminal tail gave a gain of CO2-sensitivity in Lepidosiren Cx26, we changed two glycine residues in the extended C-terminal tail to proline (Fig. 8a). Remarkably, the presence of the prolines in the C-terminal tail conferred CO2-sensitive opening on Lepidosiren Cx26 hemichannels (Fig. 8b, c). We conclude that hemichannels of Cx32, and by extension Cx30, both of which have extended C-terminal tails, are CO2 sensitive because the presence of proline residues prevents the extended tail from interfering with either CO2 binding to the carbamylation motif or the subsequent conformational change that leads to hemichannel opening.

The ancestral function of the carbamylation motif. The carbamylation motif exists in Cx32, including Cx32 of shark, suggesting a very ancient evolutionary origin to at least the ancestor of all gnathostomes. The motif has also been conserved in homologues of Cx26 (amphibian and lungfish) in which the hemichannels are not CO2 sensitive. It is notable that only a single base change is needed to effect the K125R mutation, which destroys CO2 sensitivity. That the motif has been preserved over 400 MY, suggests selection pressure to maintain an important biological function. This in turn suggests that the original function of the motif in Cx26 must have been something other than opening the hemichannel.

We have recently discovered that modestly elevated CO2 has two actions on mammalian Cx26: (1) opening of hemichannels; and (2) to closing of gap junctions. As mutations that remove the ability of the hemichannel to open to CO2 also remove the ability of CO2 to close Cx26 gap junctions, the CO2 dependent closure of Cx26 gap junctions seems to depend on CO2 binding to the same residues that open the hemichannel i.e. the carbamylation motif. To explore whether gap junction closure might be the ancestral function of the carbamylation motif in Cx26, we
**Fig. 5** The extended C-terminal tail of non-amniote sarcopterygians inhibits the CO₂ sensitivity of Cx26 hemichannels. 

**a** Comparison of the C-terminal tail of Cx26 in amniote and non-amniote vertebrates. 

**b** Removal of the C-terminal tail of Xenopus and substitution of PV for final two residues (xtCx26ΔPV) gives good expression of the modified Cx26 and demonstrates gain of CO₂ sensitivity. Addition of the Xenopus C-terminal tail to human Cx26 (hCx26 + XenCT) causes loss of CO₂ sensitivity. In *Latimeria*, a copy of the Cx26 gene lacking the C-terminal tail is CO₂ sensitive. 1. XP_014348762.1, 2. ENSLACG000000007568. 

**c** Summary data showing the increase in dye loading with the 55 mmHg PCO₂ and zero Ca²⁺ stimuli (expressed as the change in pixel intensity from control 35 mmHg PCO₂). Box and whisker plots: box, first and third quartiles; horizontal line, median; whiskers, the furthest point that lies no more than 1.5 times the interquartile range from the median. Each point represents the median change in pixel intensity from the control saline (35 mmHg PCO₂) from an independent transfection. Scale bar 20 µm.
examined whether CO$_2$ could close *Lepidosiren* Cx26 gap junctions.

We tested whether exposure to different levels of PCO$_2$ could affect the movement of a fluorescent tracer from a single cell (loaded via a patch pipette) through gap junctions to coupled cells. A PCO$_2$ stimulus of 55 mmHg prevented dye-spread through *Lepidosiren* gap junctions, and permeation of the dye only occurred once the saline had been changed to a PCO$_2$ of 35 mmHg (Fig. 9a–c). This demonstrates that *Lepidosiren* Cx26 gap junctions are closed by CO$_2$ even though the hemichannels are insensitive (Fig. 3b, c). The extended C-tail therefore does not interfere with binding of CO$_2$ to the carbamylation motif or the conformational changes that this induces in the gap junction to close it. Presumably the C-terminal tail prevents conformational change leading to hemichannel opening. By association we reasoned that the gap junctions of Cx32 may also be sensitive to CO$_2$. To our surprise, we found that permeation of fluorescent tracer occurred through the gap junction very rapidly at all levels of PCO$_2$ tested (Fig. 9d, e). Thus, gap junctions of Cx32, unlike those of Cx26, are not sensitive to CO$_2$ at these doses.

**Discussion**

By studying CO$_2$ sensitivity in the β connexin clade in a number of different phylogenetic groups from shark to mammals, we have unexpectedly revealed new structural requirements that determine the actions of CO$_2$ on these connexins. As we have already described, the carbamylation motif is a necessary requirement for CO$_2$-dependent modulation\(^\text{12}\). The presence of this motif engenders CO$_2$-dependent closure of gap junctions of Cx26. For Cx26 hemichannels, a further structural condition is required to gain CO$_2$-dependent opening: the truncation of the C-terminal tail. When this tail is present, it prevents this opening action of CO$_2$. That the extended tail does not prevent gap junction closure, strongly suggests that the tail does not interfere with the carbamylation event, or indeed the conformational change leading to gap junction closure, but instead prevents the conformational changes required to open the hemichannels once CO$_2$ has bound. A parsimonious explanation might be that the extended C-terminal tail stabilises the closed conformation of both the gap junction and hemichannel when CO$_2$ is bound.

However, our analysis highlights a further essential structural feature of the C-terminal tail for CO$_2$-dependent hemichannel opening. The very long tail of Cx32 still permits CO$_2$-dependent opening of Cx32 hemichannels, albeit at significantly higher levels of PCO$_2$. The presence of prolines in this tail permits hemichannel opening in response to CO$_2$. Changing these prolines to glycine abrogates the CO$_2$ sensitivity, and introducing prolines into the C-terminal tail of the non-sensitive *Lepidosiren* Cx26 gives a gain of function and permits CO$_2$-dependent opening of the hemichannel. Presumably, the proline residues introduce a degree of conformational restriction into the C-terminal tail that prevents the extended tail from interfering with hemichannel opening.

Given that there are two functions of the carbamylation motif -gap junction closing and hemichannel opening, what was the original ancestral function of this motif? Our finding that some Cx26 orthologues (*Xenopus*, lungfish) possess the motif, but do not open to increased CO$_2$, strongly suggests that modulation of gap junction activity might be the original function. This receives further support from our demonstration that the lungfish Cx26 gap junctions can indeed be closed by CO$_2$.

The level of PCO$_2$ tested in this study (55 mmHg) is high compared to the typical levels of PCO$_2$ found in lungfish and amphibia. This dose of PCO$_2$ is near saturating for Cx26, which in mammals and birds is sensitive to changes of PCO$_2$ over the range 20–60 mmHg\(^\text{6,18}\). Ventilation in *Lepidosiren* responds to changes in PCO$_2$ over the range 21–42 mmHg and is controlled by central chemoreceptors that are sensitive to both pH and PCO$_2$\(^\text{22}\). Breathing in *Rana catesbeiana* responds to changes in PCO$_2$ from 6–42 mmHg\(^\text{23}\). While the hemichannels of both these species are insensitive to CO$_2$, the *Lepidosiren* gap junctions were completely closed by a PCO$_2$ of 55 mmHg. It is therefore possible that the CO$_2$-sensitivity of Cx26 gap junctions (i.e. involving partial closure) at lower levels of PCO$_2$ could contribute to the chemosensory control of ventilation in these species. Further experimental data is needed to test this proposition.

Gap junctions of Cx32 are insensitive to the levels of PCO$_2$ used in this study. This suggests that in Cx32 the original function of the carbamylation motif was to open hemichannels. Cx32 hemichannels of fish and humans can be opened by sufficiently high levels of PCO$_2$ (55–70 mmHg). In entirely water breathing vertebrates, such as elasmobranch or actinopterygian fish, systemic PCO$_2$ is only slightly above the ambient\(^\text{24}\). It is therefore very unlikely that systemic PCO$_2$ would reach a range of 55–70 mmHg sufficient to open Cx32 hemichannels. Thus, Cx32 hemichannels in fish are most probably not used as systemic CO$_2$ sensors to regulate breathing in the way that Cx26 hemichannels are used in mammals. The possible functions of the
The CO2-sensitivity of Cx32 remains enigmatic. The preservation of the carbamylation motif in Cx32 over a long evolutionary period, suggests that there is indeed some important physiological function for the CO2-sensitivity of this connexin. A possible hypothesis is that the Cx32 hemichannels are important to detect locally-produced CO2. We speculate that a metabolically active group of cells (such as hepatocytes, which abundantly express Cx32) might produce very high localized concentrations of CO2 that would be sufficient to open Cx32 hemichannels.

We hypothesize (Fig. 10) that the connexin ancestor of the Cx32 and Cx26 clades would have possessed the carbamylation motif and that most likely this motif served to permit CO2 opening of the ancestral hemichannel at high levels of PCO2 (70 mmHg). When the two clades split, the Cx26-like clade gained a new functionality — ability of CO2 to close the gap junction at more modest levels of PCO2 (55 mmHg) but simultaneously lost the old functionality — the ability of CO2 to open the hemichannels. During the evolution of amniotes, when the Cx26-like gene duplicated to give Cx26 and Cx30, a further evolutionary innovation occurred — loss of the C-terminal tail from the amniote subclade of Cx26. This permitted the opening of Cx26 hemichannels at modest levels of PCO2, at a sensitivity range that was appropriate for systemic CO2 sensing, and retained the ability of CO2 to close the gap junction.

Fig. 7 Cx32 hemichannels in fish can be opened by high levels of CO2. a Sequence of the carbamylation motif (pink box) and extended C-terminal tail in Cx32 of human, Danio and Rhincodon. Note the presence of proline residues in the extended C-terminal tails of each species. b Dye loading experiments showing a small amount of dye loading with 55 mmHg CO2, and substantial loading with 70 mmHg PCO2 and zero Ca2+. c Box and whisker plots: box, first and third quartiles; horizontal line, median; whiskers, the furthest point that lies no more than 1.5 times the interquartile range from the median. All points the median change in pixel intensity from the 35 mmHg control saline. Each point represents result from an independent transfection. Scale bar 20 µm.
It is striking that Cx26 hemichannels with the structural features that permit opening by CO₂ have so far only been found in amniotes (Fig. 1, Supplementary Table 2). Equally notable, is that the Cx30 of non-mammalian amniotes lacks the carbamylation motif. Thus, the universal CO₂ sensor in amniotes is the hemichannel of Cx26 rather than that of Cx30. The key additional step to evolve CO₂-sensitive Cx26 hemichannels in amniotes was to truncate the extended C-terminal tail, which permits CO₂ sensitive opening of hemichannels. This is illustrated in Fig. 8a, where sequences of modified human Cx32 C-terminal tail with glycine in place of proline are shown, as well as Lepidosiren Cx26 C-terminal tail, showing the glycines that were changed to proline. Human Cx32 hemichannels can be opened by CO₂. Mutation of prolines in the extended C-terminal tail abolishes CO₂ sensitivity (Human Pro to Gly). Introduction of two prolines into the Lepidosiren C-terminal tail gives a gain of CO₂ sensitivity (compare to Fig. 3b, c).

**Fig. 8** Prolines in the extended C-terminal tail, permit CO₂ sensitive opening of hemichannels. a Sequences of: modified human Cx32 C-terminal tail with glycine in place of proline; and Lepidosiren Cx26 C-terminal tail, showing the glycines that were changed to proline. b Human Cx32 hemichannels can be opened by CO₂. Mutation of prolines in the extended C-terminal tail abolishes CO₂ sensitivity (Human Pro to Gly). Introduction of two prolines into the Lepidosiren C-terminal tail gives a gain of CO₂ sensitivity (compare to Fig. 3b, c). c Summary data showing the change in median pixel intensity compared to the control (PCO₂ 35 mmHg) from five independent replications for each connexin. KW, Kruskal–Wallis ANOVA, pairwise comparisons Mann Whitney. Box and whisker plots: box, first and third quartiles; horizontal line, median; whiskers, the furthest point that lies no more than 1.5 times the interquartile range from the median. Each point represents result from an independent transfection. Scale bar 20 µm.
The ancestral function of the carbamylation motif in Cx26 but not Cx32 is to close gap junctions. a Images showing rapid permeation of NBDG (within 1 min of establishing whole-cell recording) through the Lepidosiren Cx26 gap junction when PCO2 is 35 mmHg. In the images, red shows the distribution of the mCherry-tagged Lepidosiren Cx26, green is NBDG fluorescence, the yellow arrow indicates the gap junction between the cells. The numbers in bottom right hand corner are minutes after establishing whole-cell recording configuration. Scale bar, 20 μm. b Permeation of NBDG through the gap junction is delayed by elevated PCO2. The cells were perfused with hypercapnic saline (PCO2 55 mmHg) for 2 min following breakthrough, and then transferred to control saline (PCO2 35 mmHg). Significant permeation of the dye into the coupled cell is apparent only by 6th minute. c Summary data showing the effect of PCO2 on delaying permeation of dye through the gap junction to the coupled cell. d Cx32 Gap junctions are insensitive to CO2. NBDG permeates rapidly (within seconds after establishing the whole-cell configuration) through the gap junction at all levels of PCO2. e Summary data showing that there is no difference in the time required for dye transfer between coupled cells at different levels of PCO2. N = 6 for each treatment (independent replicates); box and whisker plots show the median and interquartile range (IQR), with the whisker indicating the furthest point that lies no more than 1.5 times the IQR from the median. The time for dye transfer was calculated to be when the acceptor cell had reached 10% of the fluorescence of the donor cell.

Fig. 10 Inferred evolution of CO2-dependent functionality in the Cx32 and Cx26-like clades. The common ancestor of the Cx32 and Cx26-like genes (Pre Cx32) most likely had the carbamylation motif (CM). We postulate that this was originally used to regulate the opening of hemichannels; the CM and this functionality has been preserved in Cx32 to the present day. The emergence of the Cx26-like gene was accompanied by a de novo function for the CM — gain of CO2-dependent gap junction closure, but at the cost of losing CO2-dependent hemichannel opening. In the pre-amniote world, the functions of opening hemichannels and closing of gap junctions were subserved by different gene products. With the evolution of amniotes, the Cx26-like gene was duplicated to give Cx26 and Cx30. Cx30 gained a long C-terminal tail and in many cases lost the carbamylation motif. Cx26 in amniotes lost the C-terminal tail and regained the ability of CO2 to open the hemichannel. (Green box indicates near-universal presence of carbamylation motif, light green box presence of carbamylation motif in some species but not others.)
Extant amniotes can only exchange gases by breathing air — they have no capacity for gas exchange via water. One litre of air contains about 30 times the amount of O₂ as the same volume of water. Consequently, amniotes can have much lower ventilation rates than water breathing animals. As a result of these lower ventilation rates, air breathing vertebrates accumulate much higher levels of CO₂ (compared to water-breathers). For example, mammals typically have a PCO₂ in arterial blood of ~40 mmHg, whereas water breathing fish have a blood PCO₂ of ~5 mmHg. Amniotes have adapted to the high levels of PCO₂ by retaining much higher concentrations of HCO₃⁻, thus regulating their blood pH to the required physiological levels. Nevertheless, for amniotes, the regulated excretion of CO₂ and consequent homeostatic control of acid base balance is a key rate-limiting step critical for life. Amniotes have therefore shifted the primary regulation of breathing from the detection of O₂ to the detection of CO₂ and pH. While pH-sensitive mechanisms of central respiratory chemosensitivity are clearly important, the evolutionary innovation of a CO₂ sensor (hemichannels of Cx26) of other therms, they use basking behaviour to elevate their body temperature. Although reptiles are poikilotherms, they use basking to elevate their body temperature (and metabolic rates). The arterial PCO₂ of reptiles is much higher than that of mammals, and can exceed 40 mmHg in sun-bathing lizards, and can exceed 55 mmHg in the tuatara. The CO₂-sensitivity of reptiles is very temperature dependent, but is usually above 20 mmHg, can be particularly valuable for amniotes.

Methods

Phylogenetic and microsyntenic analyses. For the phylogenetic analysis the Cx26, Cx32, and Cx30 orthologous sequences were collected from Ensembl or NCBI databases. The Proteus anguinus sequence was retrieved from the transcriptome previously published by Biscotti et al., while the sequence of Neoceratodus forsteri was obtained from NCBI (PRJNA317231). Callorhinus milii sequences were collected from http://esharkgenome.imcb.a-star.edu.sg. Moreover, sequences of genes located in the same genomic regions of those of interest were also added to the phylogenetic analysis. Accession numbers of all sequences used are reported in Supplementary Table 1.

The alignment was performed with MUSCLE (https://www.ebi.ac.uk/Tools/msa/muscle/) using default parameters. The phylogenetic analysis was carried out with MrBayes-3.2, and, on the basis of the results of microsyntenic analysis, the sequence of Petromyzon marinus (Cax27.5) was constrained to form a monophyletic clade with those located in the same genomic region. The Jones aa model was identified by the MrBayes program with a posterior probability of 1.00. The connectivity sequence of Amia calva was used as the outgroup (accession number EU016003331). 6,000,000 generations were run and sampling was conducted every 100 generations. Stationarity was defined as the condition where the standard deviation of split frequencies reached 0.0077. The first 15,000 trees were discarded as the burn-in.

The microsyntenic arrangement of the connexin genes here analysed were obtained from Ensembl with the exception of C. milii obtained from the UCSC Genome Browser (http://genome.cse.ucsc.edu/).

Generation of connexin expression constructs. The Cx26 gene sequences from Latimeria chalumnae (XM_014493276.1), Chelonias midas (XM_007051931.1), and Xenopus (XM_015429500.1), Xenopus tropicalis (CR848317.2), Lepidosiren paradoxa (GHE20103311.1) and Cx32 gene sequences from Danio rerio (XM_001921588.7) and Rhinodontens typus (XM_020523441.1) were synthesised by GenScript and subcloned into the pCAG-GS vector prior to mammalian cell transfection. xtcx26A and xtcx26DPV were made by creating PCR fragments with the forward primer shown in Table 1 and the reverse primers XTHLrev2 and Xen244PVrev, respectively (Table 1). The presence of the correct Cx26 was confirmed by DNA sequencing (GATC Biotech).

The Xenopus Tail was added to human Cx26 (to create hcx26 + XenCT of Fig. 5) by Gibson Assembly with the following fragments: the vector fragment was created by PCR from pCAG-humCx26-mCherry using human-Xen reverse and mchnid forward primers (Table 1); the insert fragment was created by PCR from pAG-Xenopus-mCherry using XenTail forward and reverse primers (Table 1).

Mutations to change prolines to glycines in the human Cx26 tail sequence were introduced stepwise by the Quikchange protocol (Agilent) using the primers shown in Table 1 for P228G/P229G, P242G, and P268G. Mutations to change glycines to prolines in the Lepidosiren paradoxa Cx26 tail sequence were introduced stepwise using the primers shown in Table 1 for G222P and G238P.

HeLa cell culture. HeLa DH cells were grown in Dulbecco’s Modified Eagle Medium (DMEM), or HeLa Ohio cells were grown in Eagle’s Minimum Essential Medium, supplemented with 10% fetal bovine serum, 50 μg/mL penicillin/streptomycin and 3 mM CaCl₂. HeLa DH cells were used for patch clamp studies on all Cx26 variants and for the dye loading experiments, the cells were plated onto coverslips at a density of 5 × 10⁴ cells per well, and transiently transfected with the Cx26:3x-mCherry expression constructs following the GeneJuice Transfection Reagent protocol.

Recording solutions. Control (35 mmHg PCO₂): 124 mM NaCl, 26 mM NaHCO₃, 1.25 mM NaH₂PO₄, 3 mM KCl, 10 mM D-glucose, 1 mM MgSO₄, 2 mM CaCl₂. This was bubbled with 95% O₂/5% CO₂ and had a final pH of ~7.4.
Hypercapnic (55 mM Hg PCO2; 100 mM NaCl, 50 mM NaHCO3, 1.25 mM NaH2PO4, 3 mM KCl, 10 mM D-glucose, 1 mM MgSO4, 2 mM CaCl2. This was bubbled with 95%CO2/5%O2 to establish the whole-cell mode, images were collected every 10 s. The time required for dye transfer was calculated to be when the background ROI for each image was subtracted from each cell measurement from the same image analysis via ImageJ.

Patch clamp recordings. Cover slips containing non-confluent HeLa DH cells (prefered over the HeLa Ohio cells for their more rounded morphology) were placed into a chamber and superfused with control saline. An MCl Cleverscope, Photometrics Prime camera and Caim Instruments OptoLED illumination, and an Olymix 60x water immersion objective (NA 1.0) objective were used to visualize the cells under brightfield DIC, and mCherry expression (470 nm). Micromanager software was used to control the illumination and camera settings and to save images for offline analysis via ImageJ.

Standard patch clamp techniques were used to make whole-cell patch clamp recordings from HeLa cells that expressed Cx26 as assessed by mCherry fluorescence. The intracellular fluid in the patch pipette contained: K-gluconate 130 mM, KCl 10 mM, EGTA 10 mM, CaCl2 2 mM, and HEPES 10 mM, pH adjusted to 7.3 with KOH and was adjusted with pure water to a final osmolality of 295 mOsm. All whole-cell recordings were performed at a holding potential of −50 mV with steps to −40 mV, lasting 2.5 s and delivered every 5 s, to assess whole-cell conductance. Imaging of fluorescent tracer movement through gap junctions was achieved by using 2-Dexoy-2-[7-nitro-2,1-benzoxadiazol-4-ylamino]-D-glucose, NBDG, which was included at 200 µM in the patch recording fluid which was either the same as above or had lowered EGTA concentration (5 mM). Following break-through of the patch pipette to establish the whole-cell mode, images were collected every 10 s. The time required for dye transfer was calculated to be when the acceptor cell reached 10% of the fluorescence of the donor cell.

Statistical analysis and reproducibility. Data has been plotted as either cumulative probabilities (showing every data point) or box and whisker plots where the box is interquartile range, bar is median, and whisker extends to most extreme data point that is no more than 1.5 times the interquartile range. All individual data points are superimposed on the plots. For the patch clamp experiments, an individual replicate is a recording from a single cell. For the gap junction permeation studies, dye transfer between a pair of cells is regarded as a single replicate. For the dye loading studies, single replicate is the analysis of CO2 sensitivity from cells resulting from an independent transfection. In this case to avoid pseudoreplication, statistical analysis was performed on the median values arising from each of the independent replicates. Statistical analysis was performed with the R language. All analysis was performed using the Kruskal–Wallis ANOVA for multiple comparisons and the Mann Whitney U test for pairwise comparisons.

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

V.M.D., F.S. and E.d.W. performed the dye loading experiments and data analysis. S.N. performed the patch clamp recordings and gap junction permeation experiments and data analysis. J.C. generated the constructs used to express Cx26, Cx32 and the various mutations thereof. M.A.B. performed the molecular phylogenetic and synteny analysis. M.A.B. and M.G. contributed transcriptomic data for lungfish Cx26 and Cx32. N.D. conceived the study, analyzed the data, and wrote the paper. All authors commented on drafts of the paper.

Additional information

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