Xenopus connexins: how frogs bridge the gap

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Abstract Animal species use specialized cell-to-cell channels, called gap junctions, to allow for a direct exchange of ions and small metabolites between their cells' cytoplasm. In invertebrates, gap junctions are formed by innexins, while vertebrates use connexin (Cx) proteins as gap-junction-building blocks. Recently, innexin homologs have been found in vertebrates and named pannexins. From progress in the different genome projects, it has become evident that every class of vertebrates uses their own unique set of Cxs to build their gap junctions. Here, we review all known Xenopus Cxs with respect to their expression, regulation, and function. We compare Xenopus Cxs with those of zebrafish and mouse, and provide evidence for the existence of several additional, non-identified, amphibian Cxs. Finally, we identify two new Xenopus pannexins by screening EST libraries.

Key words Xenopus · connexin · pannexin · gap junction · development · mouse · zebrafish

Introduction

Many of today’s genes find their origins in ancient prototypes. They first appeared in pro- and uni-cellulars, and function in basic cell metabolism or division. The evolutionary leap into integrated multicellular life forms demanded the exchange of information between the co-operating cells, for instance, providing positional information to generate the body plan of the organism. Many of the patterning genes are thought to have found their origin at that time. Cells also started communication by many different secreted hormones and transmembrane receptor molecules, whereby old genes adapted to new functions and new genes were added to the so-called metazoan “genetic toolkit”. Furthermore, direct cell-to-cell communication was established by means of gap junctions in animal species. Gap junctions are found in all vertebrates and invertebrates.

Upon the discovery of the vertebrate gap-junction proteins in the mid-1980s, the so-called connexin (Cx) molecules, it became obvious that sequence-related invertebrate Cxs were absent. More recently, it became clear from Drosophila and Caenorhabditis elegans genome projects that proteins structurally related to Cxs are present in these invertebrates, and are now known as innexins (Phelan and Starich, 2001). It appears that innexins predate Cxs that have evolved independently (Stout et al., 2004). The finding of Cx molecules in invertebrate chordates demonstrates that Cxs have evolved earlier than was initially thought (Sasakura et al., 2003; White et al., 2004), but still after the appearance of innexins. As a result of an in-depth search for innexin sequences present in the vertebrate genomes, several orthologs were identified in mouse, rat, and man, and subsequently named pannexins (from the Latin pan, all, and nexus, connection) (reviewed by Panchin, 2005). Functional analyses demonstrated the ability of the pannexins to form functional gap junctions (Bruzzone et al., 2003). Furthermore, it is argued that innexins, pannexins, and Cxs are members of three different gene families, of which the first one is not co-expressed with the latter two in the same organism (White et al., 2004). Most peculiarly, it became clear that even between the different vertebrate classes, many differences in Cx molecules exist, and for many mammalian Cxs, no genuine ortholog can be easily nominated in birds, fish, amphibians, or tunicates on the basis of sequence characteristics only. It has become clear that every class of vertebrates uses its own unique set of Cxs to build their gap junctions.
This review focuses on amphibian Cxs, in particular, *Xenopus*. For excellent recent reviews on Cxs in human disease and mammalian development, we refer to Wei et al. (2004) and Houghton (2005). Here, we review *Xenopus* Cx expression patterns, regulation, and functional properties and compare these with Cxs from other species. Finally, as the genome sequencing projects of *Xenopus* progress rapidly, we will explore the generated databases for new *Xenopus* Cx and pan-nexin molecules.

**Xenopus Cxs**

While genome sequencing reveals that mouse and human express at least 20 Cxs (Willecke et al., 2002; Söhl and Willecke, 2003), only seven *Xenopus* Cx isoforms have been described. These are, named by their predicted molecular weight, Cx30, Cx31, Cx38, Cx40.4, Cx41, Cx43, and Cx43.4 (Gimlich et al., 1988, 1990; Ebihara et al., 1989; Yoshizaki and Patiño, 1995; Landesman et al., 2003; De Boer et al., 2005). Based on their nucleotide sequence only, they can be separated into an α-group (Cx38, Cx40.4, Cx41, Cx43), a β-group (Cx30, Cx31), and an unspecific group (Cx43.4) (De Boer et al., 2005).

Expression data of the *Xenopus* Cxs, i.e. temporal, developmental pattern, and adult tissue distribution, are summarized in Table 1. Cx31, Cx38, Cx43, and Cx43.4 are maternally expressed and are therefore possibly involved in pre-midblastula transition (MBT) development. Of these, Cx31 and Cx38 have very high expression levels, while pre-MBT expression levels of Cx43 are very low, and detectable by RT-PCR (Landesman et al., 2003), but not by whole-mount *in situ* hybridization (Van der Heyden et al., 2001a). Cx30 and Cx40.4 are only expressed post-MBT. From the summation in Table 1, it becomes evident that analyses of the embryonic expression patterns and adult tissue distribution are far from complete for many of the Cxs. The best studied are Cx30, Cx40.4, and Cx43 for which the temporal expression, developmental pattern, and tissue distribution are well established.

**Transcriptional and translational control**

Gap-junctional communication is regulated at different levels: transcription, translation, intracellular trafficking, oligomerization, docking, and gating (Segretain and Falk, 2004; Teunissen and Bierhuizen, 2004). Virtually nothing has been described for the latter four mechanisms with respect to *Xenopus* Cxs.

In recent years, enormous technical progress has been made in producing transgenic *Xenopus* embryos. This now enables *in vivo* promoter analysis during development (Amaya et al., 1998; Hirsch et al., 2002). Of the known *Xenopus* Cxs, only the promoter of Cx43

**Table 1 Expression data**

<table>
<thead>
<tr>
<th>Stage</th>
<th>Development</th>
<th>Adult</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cx30</td>
<td>11, 14, 15, 16, 19, 20,</td>
<td>Hatching gland anus,</td>
<td>Lung, liver, intestines</td>
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<tr>
<td></td>
<td>25, 30, 35, 38, 41</td>
<td>endoderm, cement gland,</td>
<td>stomach, kidney</td>
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<tr>
<td>Cx31</td>
<td>VI oocyte</td>
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<td>Ovary</td>
</tr>
<tr>
<td></td>
<td>Oocyte</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2, 15, 30</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cx38</td>
<td>1–IV oocyte</td>
<td>Not determined</td>
<td>Ovary</td>
</tr>
<tr>
<td></td>
<td>IV–VI oocyte</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Oocyte</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1, 2, 8, 9, 10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cx40.4</td>
<td>10, 18, 22, 26, 30, 35, 41</td>
<td>Presomitic mesoderm</td>
<td>Brain, heart, stomach</td>
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<tr>
<td>Cx41</td>
<td>Oocyte</td>
<td>somites</td>
<td>leg muscle, kidney</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Ovary, heart</td>
</tr>
<tr>
<td>Cx43</td>
<td>I–IV oocyte</td>
<td>Lens, limb bud,</td>
<td>Lung, liver, intestines</td>
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<tr>
<td></td>
<td>IV–VI oocyte</td>
<td>Notochord, Branchial</td>
<td>Stomach, spleen</td>
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<td></td>
<td>Oocyte</td>
<td>arches, Cement gland,</td>
<td>Body wall muscle,</td>
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<td>1, 7, 38, 41</td>
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<td>Neural folds, eye vesicle</td>
<td>Ovary, testis, eye</td>
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<td>Oocyte</td>
<td>Head, spinal cord, tail,</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2, 6, 8.5, 10, 15, 17, 24, 30, 33, 35</td>
<td>Branchial arch, brain</td>
<td></td>
</tr>
</tbody>
</table>

*No connexin43 expression was found in the working myocardium of the heart. Traces of connexin43 expression were found in whole-heart homogenates.*
has been cloned (Van der Heyden et al., 2001b). The immediate upstream regions, with respect to the transcription initiation site, display strong homology to mammalian Cx43 promoters. This region has been characterized as the proximal promoter in rat Cx43 (Teunissen et al., 2003). Further upstream, the sequence homology is very limited (own unpublished observations). The proximal promoter region displays developmentally regulated promoter activity that correlates with low Cx43 levels in neurula-stage embryos (st 18/19) and an increased promoter activity in early and late tadpole stages (st 28/29, st 37/38) (Van der Heyden et al., 2001b).

Translational control provides an excellent mechanism for fast adaptation to rapidly changing demands, as the mRNA is already present. The mechanism of translational derepression may play a role in different stages of development or at different locations within the rapidly developing embryo. A few studies investigated translational control of Xenopus Cxs mediated by their 5' and 3' untranslated regions. In two elaborate studies, it was demonstrated that the translation efficiency of Cx41 was strongly affected by the presence of three short upstream open reading frames (uORFs) (Meijer et al., 2000a; Meijer and Thomas, 2003). The first, uORF1, generated the largest inhibition, although translation initiation at this uORF was relatively infrequent. However, the use of a rare leucine codon made the translation elongation of uORF1 fairly slow. Furthermore, the termination codon of uORF1 caused another delay. The combinatorial effect most likely results in upstream ribosome stalling, leading to inhibited mRNA scanning and therefore reduced translation initiation at the downstream Cx41 coding ORF. Only very limited translational control was observed for Cx30 (Meijer et al., 2000b). A similar mechanism has now been suggested for mouse Cx43, which also contains multiple uORFs (Pfleifer et al., 2004).

Some Cxs have been detected as molecular targets of upstream signaling pathways. In vitro stimulation of ovarian fragments with human chorionic gonadotropin hormone resulted in decreased Cx41 and Cx43 levels within 4 hr, while Cx38 expression remained unaffected (Yoshizaki and Patiño, 1995). Subtractive hybridization approaches using animal caps of Wnt3a-injected embryos revealed Cx30 as a target gene for Wnt3a-induced signaling (McGrew et al., 1999). In mammals, Cx43 has been determined as a genuine Wnt target (Van der Heyden et al., 1998; Ai et al., 2000).

### Role of Xenopus Cxs in development

Patterns of gap-junctional communication in Xenopus

An important role for gap-junctional communication in embryonic development was proposed in the 1960s (Potter et al., 1966; reviewed in Furshpan and Potter, 1968). In those initial days, the developmentally powerful Xenopus model system also became involved into this rapidly expanding research field. In 1969, halothane-inhibitory electrical coupling between Xenopus blastomeres was demonstrated (Palmer and Slack, 1969). In contrast, metabolic coupling by fluorescent injection in animal pole cells could not be detected (Slack and Palmer, 1969). Subsequently, electrical coupling and its underlying gap junctions were investigated in detail during the 1970s in cleavage stage embryos, isolated blastomeres, oocyte–follicle interaction systems, neural induction, and other systems (Warner, 1973; DiCaprio et al., 1975, 1976; Sanders and DiCaprio, 1976; Turin and Warner, 1977; Browne et al., 1979; Spray et al., 1979). In the 1980s and 1990s, differences in communication capacity within the developing embryo and regulation of this by upstream signaling pathways and cell cycle transition were reported (Rink et al., 1980; Bluemink et al., 1983; Guthrie, 1984; Van den Hoeft et al., 1984; Guthrie et al., 1988; Nagajski et al., 1989; Su et al., 1990; Olson et al., 1991; Olson and Moon, 1992; Kruftka et al., 1998; Levin and Mercola, 1998). Many of these studies partly rely on fluorescent dye injection and subsequent whole-mount analysis. Although dye injections in Xenopus blastomeres are relatively easy to perform, their interpretations are rather cumbersome and are prone to technical artifacts, as convincingly shown by Landesman et al. (2000). They repeated many of the previous experiments and injected embryos with different fluorescent and non-fluorescent dyes in combination with non-permeable tracers. While whole-mount analysis by conventional fluorescence microscopy of lucifer yellow-injected embryos indicated communication in many examples, subsequent embryo sectioning revealed this to be an optical artifact. The earlier observed differences in communication when comparing the ventral with the dorsal side of the embryo also appeared to be confused by artifacts caused by a combination of several factors such as differential pigment distribution, cell size, yolk distribution, and the so-called lens artifacts. This led to the conclusion that fluorescent dye injection as a tool for studying gap-junctional communication should be used very carefully. However, injection of the non-fluorescent dye neurobiotin revealed extensive gap-junctional communication between the different blastomeres. In these experiments, no differences in the amount of dye coupling was observed within the different regions of the embryo. Nevertheless, conclusions of earlier studies concerning the existence of communication within the developing embryo and selectivity for different dyes were validated. Using more restricted dyes, regional differences in gap-junctional communication within the developing Xenopus embryo might still be revealed in the end. When using the appropriate methodology, Xenopus will be a valuable model to pursue this goal.
Manipulating gap junctions during *Xenopus* development

An early study on the involvement of gap-junctional communication in early *Xenopus* development demonstrated that blocking of communication by halothane led to delayed cleavage rates and inhibited gastrulation in some embryos (Palmer and Slack, 1969). Warner et al. (1984) injected anti-gap-junction antibodies (raised against rat Cx32) into the right anterodorsal cell of an eight-cell embryo, resulting in a disruption of electrical coupling. The resulting tadpoles bore a characteristic set of abnormalities, like absence or size reduction of the right eye and underdevelopment of the brain at this site. Antibody injection just before first cleavage or in both cells of the two-cell embryo resulted in death before gastrulation. A Cx32–Cx43 chimeric Cx acts as a dominant-negative Cx on maternally expressed Cx38 (Paul et al., 1995). Injection of this chimera into the right anterodorsal cell of the eight-cell embryo resulted in communication deficiency in this cell’s progeny from stage 7 onward (Paul et al., 1995). This resulted in delamination and extrusion of the affected cells from stage 8 onward. Rescue was seen with co-expression of Cx37, which is not subject to dominant-negative inhibition by the Cx32–Cx43 chimera. The authors did not observe developmental defects in ventrally injected embryos. The developmental defects in the tadpole stages were similar to those reported for the antibody injection study. However, in the antibody inhibition study, no extrusion or delamination was reported. Although the effects of gap-junctional inhibition on development are clear from these examples, they cannot be easily attributed to a specific Cx isoform.

A strong correlation has been found between Cx30 expression and hatching of embryos from the vitellin membrane (Levin and Mercola, 2000). Hatching depends on the secretion of hatching-associated enzymes, like metalloproteases. Many secretion processes have been shown to depend on functional gap junctions (reviewed by Serre-Beimier et al., 2002). In the developing embryo, Cx30 is strongly expressed in the hatching gland. The application of three different general gap-junction blockers independently to batches of embryos prior to hatching resulted in strongly inhibited hatching efficiency. Furthermore, targeted expression of the dominant-negative Cx32–Cx43 chimera to the hatching gland also resulted in decreased hatching efficiencies. These data might indicate an important function of Cx30 in hatching. However, it cannot be excluded that other, still unknown, Cx isoforms function in the hatching gland.

The *Xenopus* developmental system has been instrumental in establishing a role for gap-junctional communication in generating left–right asymmetry. A comprehensive review on left–right asymmetry incorporating the *Xenopus* results appeared recently (Levin, 2005). Using a variety of pharmacologic gap-junction inhibitors, Levin and Mercola (1998, 1999) were able to induce complete and partial heterotaxia in a large percentage of embryos. The strongest sensitivity was found to occur between stages 5 and 12, while enhanced heterotaxia could be demonstrated until stage 21. Furthermore, either inhibition of dorsal gap-junctional communication by injection of the dominant-negative Cx32–Cx43 chimera or increasing ventral gap-junctional communication by bilateral Cx26 injection caused significant increases in heterotaxia. In a follow-up paper, it was demonstrated that gap-junctional involvement in heterotaxia was upstream of asymmetric Sonic hedgehog and nodal expression (Levin and Mercola, 1999). The first left–right asymmetric distributed gene product in *Xenopus* appears to be the H+/K+-ATPase, which is depleted from the left ventral blastomere at the four-cell stage (Levin et al., 2002). In chick embryos, membrane potential at the left side of the node is more depolarized than on the right side. Given the asymmetric H+/K+-ATPase distribution, a similar difference in membrane potential seems obvious in *Xenopus* too. It is hypothesized that the gap junctions play a role in the propagation of a calcium wave in the vicinity of the node (Mercola, 2003). Whether or not the wave itself displays an asymmetric orientation is not clear, but voltage-dependent gap-junction gating (Turin and Warner, 1977; Spray et al., 1979) could display left–right differences as a result of asymmetrical H+/K+-ATPase function.

Like in mouse knock-out studies, Cx redundancy in *Xenopus* has hampered the ability to attribute certain Cxs to specific roles in development, as described already for Cx30. Another example came from a study on maternal Cx38, which, at the time of the study, was the only known maternally expressed Cx. Using antisense technology, expression of Cx38 could be severely inhibited without causing developmental abnormalities (Landesman et al., 2003). This suggested the existence of additional maternal Cxs, which were promptly identified in the same study.

In conclusion, some of the above-mentioned examples display a clear role for Cxs in *Xenopus* development. Because of Cx redundancy, however, no one-to-one correlation could be made between phenotype and Cx isoforms. Solving this problem will require further identification of additional *Xenopus* Cxs in combination with forward genetic techniques like RNAi.

**Interspecies comparisons**

Cxs have evolved relatively late in evolution (Stout et al., 2004), and Cx molecules are restricted to chordata only. Even between the different vertebrate classes, sev-
eral Cxs appear to have limited sequence homology, while others have high levels of amino acid identity (Van der Heyden et al., 2004). Therefore, it is questionable as to whether every Cx isoform has a genuine ortholog in other vertebrate classes.

In Fig. 1, a comprehensive amino acid-based sequence comparison between *Xenopus* and mouse Cxs is depicted. As can be seen, some *Xenopus* Cxs display a high level of identity to murine Cx isoforms, i.e. Cx30, Cx31, and, in particular, Cx43. Other *Xenopus* Cxs, however, display less or even limited sequence homology, i.e. Cx43.3, Cx38, and Cx40.4, and thereby it is difficult to indicate a mammalian ortholog, if any, for these.

Similar observations have been made in the studies of zebrafish Cxs (i.e., Dermietzel et al., 2000). There too, some of the zebrafish Cxs display a high degree of amino acid identity, while several others show only limited amounts of amino acid identity to mammals, or higher vertebrates. As stated earlier, by others for zebrafish, “this observation raises the question why evolution does conserve some Cxs . . . but not others?” (Zoidl et al., 2004). It seems this issue can now be extended into the *Xenopus* field. A definitive answer to this intriguing question awaits further research. Triggered by these observations, we questioned whether a comparison of *Xenopus* and zebrafish Cxs would yield higher levels of amino acid identity than those with mouse. As seen in Fig. 2, this is certainly true for some Cxs (i.e., XICx40.4 display a 63% amino acid identity with DrCx39.9). However, other Cx pairs yield even lower identity levels (i.e., XICx43 displays 73.6% identity with DrCx43, but 85.6% with human Cx43) when compared with mammalian Cxs. Therefore, *Xenopus* Cxs might be considered as intermediates between mammals and fish. Of the seven known *Xenopus* Cxs, six of them have orthologs in either mammals, fish, or both. Only Cx38 appears as a *Xenopus*-specific Cx isoform. In conclusion, the evolutionary conservation among Cxs appears very complex and seems to result from an interplay between evolutionary history, functional convergence, pleiotropy, and maybe other, unknown mechanisms. Further elucidation of the intriguing evolutionary history for all Cxs will require massive input of comprehensive sequence analyses, developmental expression patterns, and functional data.

*Xenopus* Cx43 displays a very high level of amino acid identity to mammalian Cx43 (Van der Heyden et al., 2004). Detailed expression pattern analysis also highlights many similarities in expression during development and in adults. For instance, Cx43 is expressed in the lens epithelium in frogs and mammals, and it has almost identical expression patterns during limb outgrowth in both classes (Van der Heyden et al., 2001a and references therein). It is therefore most remarkable that other important sites of mammalian Cx43 expression are not reflected in *Xenopus*. For instance, no expression is found in the developing somites (Van der Heyden et al., 2001a) or in the working myocardium (Becker et al., 1998; Van der Heyden et al., 2001a). In the developing *Xenopus* somites, Cx40.4 is expressed instead (De Boer et al., 2005), while in the working myocardium, expression of a β-variant has been suggested (Becker et al., 1998). Until the evolutionary patterns of Cx distribution have been elucidated in more detail, it remains speculative whether Cx43 in the lens epithelium, developing limb buds, and notochord are the ancient sites of expression, while myocardial and somitic functions of Cx43 in mammals have been acquired more recently.

Analysis of the electrophysiological properties of the cloned *Xenopus* Cxs has been limited to three isoforms. Several studies were performed on Cx38, while functional descriptions of Cx43.3 and Cx40.4 were only covered in one study (see Table 2 for references). Although not unexpected and while the analyses are far from complete, it appears that *Xenopus* Cxs behave in a manner similar to their mammalian counterparts.

### Additional *Xenopus* Cxs?

When compared with humans, mouse, and zebrafish, relatively few Cxs have been described in *Xenopus*. At least two observations suggest the existence of additional amphibian Cxs. First, the cumulative expression pattern of all seven known *Xenopus* Cxs does not cover every structure in the developing frog, leaving “white areas” to be filled in by additional Cxs. Second, from the genome sequencing projects of *Xenopus*, it has become evident that additional Cx-like sequences are present in the genomic and EST databases. Screening of the mouse and human genome for Cxs resulted in the identification of 20 and 21 Cx isoforms in the fully sequenced mouse and human genomes (Willecke et al., 2002, Söhl and Willecke, 2003). Exploiting the rapid progress in the *Xenopus* genome sequencing projects, we screened the *Xenopus* ESTs databases for additional Cxs. This resulted in the identification of full-length sequences for Cx25 (BC055269) and Cx26 (BC043797) and a partial sequence for Cx50 (BQ737050). Cx25 is most similar to human Cx25 (64.4% amino acid identity), Cx26 displays the strongest identity with mouse Cx26 (65.5%), while the partial Cx50 sequence has a more than 65% identity to mouse Cx50 within the overlapping region. Given that at the time of screening the EST database contained over 400,000 ESTs (compared with over 4,000,000 for mouse), it can be expected that even more Cxs are hidden within the *Xenopus* genome.

When screening the databases for other amphibian Cxs, we obtained two partial Cx sequences each in the Axolotl (*Ambystoma mexicanum*) and the Japanese fire-
Fig. 1 Comparison of Xenopus and mouse connexins (Cxs). Percentage of amino acid-based identity (x-axis) of a Xenopus Cx (indicated at the top right in each graph) is plotted against 21 mouse Cxs (left axis). Identity percentages are displayed at the right of each bar.
Fig. 2 Comparison of *Xenopus* and zebrafish connexins (Cxs). Percentage of amino acid-based identity (x-axis) of a *Xenopus* Cx (indicated at the top right in each graph) is plotted against 15 zebrafish Cxs (left axis). Identity percentages are displayed at the right of each bar.
and mouse PANX1 mRNAs are ubiquitously expressed, while PANX2 is brain specific (Baranova et al., 2004).

To look for the presence of pannexins in Xenopus, we screened X. tropicalis and X. laevis EST databases with a region of mouse PANX1 (NP_062355), PANX2 (NP_001002005), and PANX3 (NP_766042) encompassing the highly conserved first two transmembrane regions and their intervening extracellular stretch of amino acids (Baranova et al., 2004). This resulted in one positive EST for X. laevis (BU912264), most similar to mouse PANX1, and two different positive ESTs for X. tropicalis (CX978500 and CX799160), most similar to mouse PANX1 and PANX2, respectively. No ortholog was identified for mouse PANX3. Amino acid identity of XiPANX1 (stretch of 153 amino acids) is 54.9% with mouse PANX1 and 45.5% with zebrafish PANX1. The amino acid identity of XtPANX1 (N-terminal 217 amino acids) with mouse and zebrafish PANX1 is 75.6% and 61.9%, respectively. XiPANX2 identity with mouse PANX2 is 89.9% over the N-terminal 129 amino acids. Alignment of the Xenopus pannexins with those of mouse and zebrafish is depicted in Fig. 3. Amino acids shaded in green represent conserved sites between invertebrate innexins and vertebrate pannexins (Baranova et al., 2004). These data demonstrate the presence of genuine expressed pannexins within the Xenopus genomes.

XtPANX1 was identified in an oocyte EST library. XiPANX1 was found in EST databases from oocyte, gastrula, neurula, tailbud, intestine, testes, and brain EST libraries. Finally, XiPANX2 was present in an adult brain EST library. These provisional expression data are in accordance with the tissue distribution of mouse and human PANX1 and PANX2.

As for other vertebrates, ensued by our identification of pannexins in Xenopus, we can now regard these proteins as an alternative source of gap-junction proteins in the frog too.

### Table 2 Electrophysiological properties

<table>
<thead>
<tr>
<th>Single channel conductance (pS)</th>
<th>Vₒ (mV)</th>
<th>Gₘᵢₙ (%)</th>
<th>Hemi-channels</th>
<th>References</th>
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</thead>
<tbody>
<tr>
<td>Cx38 –</td>
<td>~ 25</td>
<td>~ 35</td>
<td>Yes</td>
<td>Ebihara et al. (1989, 1996)</td>
</tr>
<tr>
<td>Cx40.4 90–120</td>
<td>~ 55</td>
<td>~ 40</td>
<td>ND</td>
<td>De Boer et al. (2005)</td>
</tr>
<tr>
<td>Cx43.4 35</td>
<td>~ 20</td>
<td>5</td>
<td>ND</td>
<td>Landesman et al. (2003)</td>
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</table>

### Identification of Xenopus pannexins

Cxs have long been considered as the only gap-junction-forming proteins in vertebrates, while invertebrate gap junctions are formed by an unrelated protein family named as innexins. Recently, however, three orthologs of innexins were identified in the genomes of several vertebrates like mouse, rat, and man (reviewed by Panchin, 2005). These so-called pannexins (PANX) are able to confer electrical coupling in a classical paired Xenopus oocyte assay (Bruzone et al., 2003). Human belly newt (Cynops pyrrhogaster) databases. AmCx45 is likely the ortholog of XICx43.4 (81.4% identity over 44 amino acids of available sequence), and AmCx31 might be the XICx30 ortholog (52.6% identity over 134 amino acids). CpCx43 is the ortholog of XICx43 (82.3% identity over 304 amino acids), while CpCx35 has no obvious ortholog in Xenopus (<30% identity over 258 amino acids), and is likely another unknown amphibian Cx.

![Fig. 3 Amino acid alignment of Xenopus, mouse, and zebrafish pannexins.](attachment:alignment.png)
Conclusions and future directions

Although once a very popular model system in gap-junction research, *Xenopus* is currently merely used as a producer of oocytes for the study of ectopically expressed xeno-Cxs. The molecular identification of *Xenopus* Cxs has become less of a priority than mammalian and zebrafish Cxs in recent decades. Nevertheless, Cx research in *Xenopus* has clearly elucidated some important aspects of Cx regulation at the translational level. Furthermore, it has played an important role in understanding the developmental roles of Cxs, for instance, in establishing the left–right axis, valuable knowledge that reaches beyond the amphibians.

Two developments within the *Xenopus* field will boost molecular approaches in this model system with respect to Cxs. Firstly, the genome project has already elucidated several additional Cxs in this species. The number of identified *Xenopus* Cxs will increase further in the coming years, and we might well end up in the same numbers as in mammals. Secondly, the transgenic and RNAi approaches in this model system will be very useful in pinpointing the role of specific Cxs in developmental processes. Furthermore, upon completion of the genome projects, a comprehensive comparison of the complete Cx sets from mouse, chicken, zebrafish, *Xenopus*, and *Ciona* will likely shed more light on the enigmatic evolution of Cxs. Finally, the presence of pannexins in *Xenopus* hints at an alternative source of gap-junction proteins in the frog.

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References


