

# Molecular analysis of voltage dependence of heterotypic gap junctions formed by connexins 26 and 32

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**ABSTRACT** Heterotypic gap junctions formed by pairing *Xenopus* oocytes expressing hemichannels formed of Cx32 with those expressing hemichannels formed of Cx26 displayed novel transjunctional voltage ( $V_j$ ) dependence not predicted by the behavior of these connexins in homotypic configurations. Rectification of initial and steady-state currents was observed. Relative positivity and negativity on the Cx26 side of the junction resulted in increased and decreased initial conductance ( $g_{j0}$ ), respectively. Only relative positivity on the Cx26 decreased steady-state conductance ( $g_{j\infty}$ ). This behavior suggested that interactions between hemichannels influences gap junction gating. The role of the first extracellular loop (E1) in these interactions was examined by pairing Cx32 and Cx26 with a chimeric connexin in which Cx32 E1 was replaced with Cx26 E1 (Cx32\*26E1). Both junctions rectified with  $g_{j0}/V_j$  relations that were less steep than that observed for Cx32/Cx26. Decreases in  $g_{j\infty}$  occurred for either polarity  $V_j$  in the Cx32/Cx32\*26E1 junction. Mutation of two amino acids in Cx26 E1 increased the steepness of both the  $g_{j0}/V_j$  and  $g_{j\infty}/V_j$  relations. These data demonstrate that fast rectification can arise from mismatched E1 domains and that E1 may contribute to the voltage sensing mechanisms underlying both fast and slow  $V_j$ -dependent processes.

## INTRODUCTION

Molecular genetic studies of ion channels have generated new insights into the mechanisms of ion channel gating and permeability. Through the identification of domains whose activities are critical determinants of protein function, the molecular basis of these functions becomes addressable. Some properties of proteins appear to be determined by discrete domains that confer these characteristics even when part of chimeric constructions (1–3). Others (4) and those discussed here, seem to arise from the interactions of multiple domains.

Gap junction channels comprise a unique family of ion channels, some members of which have proven to be voltage dependent (5). They show no sequence homology with the superfamily of sodium, potassium, and calcium channels or ligand-gated channels (6). A gap junction channel forms by the association of two hemichannels, one contributed by each of two coupled cells. Each hemichannel is a hexamer of molecules termed connexins. The connexins are a multigene family with at least 11 mammalian members (for three of these amphibian homologues are known; Bennett, M. V. L. manuscript in preparation). Multiple connexins can be expressed by a single cell type and as demonstrated for Cx26 and Cx32 in hepatocytes, more than one connexin can be localized to the same gap junction plaque (7). The topology of connexins, predicted from hydropathy plots (8) and in the cases of Cx32 and Cx43, verified by

protease and antibody studies (9–11), includes four transmembrane domains (M1–M4), a cytoplasmic loop, cytoplasmic amino and carboxy termini and two extracellular loops (E1 and E2). Presumably the association between hemichannels occurs by contact between extracellular loops which creates a continuous, insulated channel between the coupled cells (Fig. 1A).

By virtue of spanning two cell membranes and the intervening gap between them, gap junctions can be subjected to two different kinds of voltage stimuli (Fig. 1B). A difference between the potentials in the interiors of two coupled cells constitutes a transjunctional potential ( $V_j$ ) and a difference between the potential in the interior of cells and the extracellular space is an inside-outside potential ( $V_{i-o}$ ).  $V_{i-o}$  will be developed largely across the channel wall because the access resistance through the intercellular gap is small compared to the leakage resistance of the wall (12).  $V_{i-o}$  can be changed without changing  $V_j$  by simultaneously applying equal polarizations to the coupled cells. When  $V_j$  is changed by applying a voltage step to one of two coupled cells,  $V_{i-o}$  is changed in the stepped cell as well as along the channel. Many gap junctions are sensitive to  $V_j$  with no sensitivity to  $V_{i-o}$ . A few have been found that are sensitive to both (13, 14). The responsiveness of gap junctions to applied voltage is most likely the result of electrical work done in changing the conformation of the channel protein rather than due to blockade by ions at sites within the channel. Single channel recordings show transitions between open and closed states that are abrupt, with no evidence

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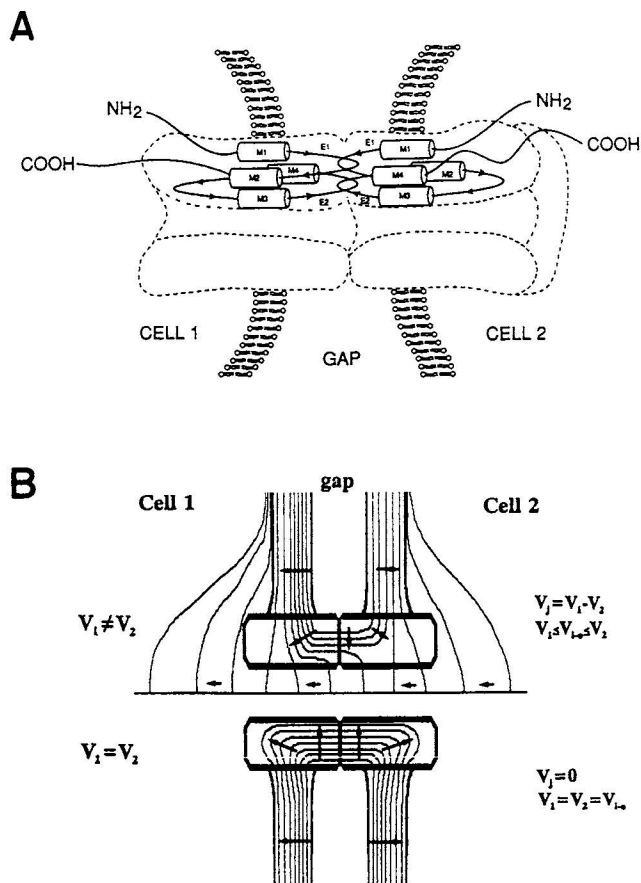


FIGURE 1 (A) Schematic representation of a gap junction channel between two cells. The dotted lines indicate the channel boundaries and the superimposed solid lines diagram the position of single connexin molecules in each of the paired hemichannels. The transmembrane domains are labeled M1–M4, the extracellular loop between M1 and M2 is indicated as E1 and the extracellular loop between M3 and M4 is labeled E2; the amino and carboxy termini are labeled NH<sub>2</sub> and COOH, respectively. (B) Diagram (from reference 13) illustrating the presumed isopotential lines in the presence of a  $V_j$  (top half) and a  $V_{i-o}$  in the absence of a  $V_j$  (bottom half). Arrows indicate field direction and suggest relative magnitude.

of flicker (6). Opening rate constants, derived from macroscopic currents, are relatively insensitive to voltage (15) and the size of the channel and its permeability to large organic ions (16) all argue against blockade as the basis for voltage dependent decreases in gap junctional conductance. In addition, voltage dependent gating has been observed in perfused and reconstituted preparations without impermeant ions in the bathing media (17).

Studies of voltage dependence of gap junctions formed by cloned connexins have been conducted primarily on homotypic junctions, those formed by the association of identical hemichannels (18–22). With the exception of

Cx26, these junctions have been shown to be dependent only on  $V_j$  with changes in conductance that are symmetric about  $V_j = 0$  mV, i.e., identical for hyperpolarization or depolarization of either cell. Junctions formed by Cx26 possess a small degree of sensitivity to  $V_{i-o}$  in addition to sensitivity to  $V_j$ . The presence of  $V_{i-o}$  and  $V_j$ -dependence results in an asymmetry in the  $G_j/V_j$  relationship of Cx26 about  $V_j = 0$  mV (18).

In the nervous system, gap junctions serve as electrical synapses. Some of these synapses are not significantly voltage dependent (5, 23), whereas others, such as the giant motor synapse of crayfish nerve cord (24, 25), hatchetfish medulla (26), and lamprey spinal cord (27) display asymmetric  $V_j$ -dependence in which depolarization on the presynaptic side rapidly increases junctional conductance. The time course of these changes in conductance is in the submillisecond range and thus are much more rapid than the changes in conductance in response to  $V_j$  observed for homotypic gap junctions. The connexins that form these fast rectifying electrical junctions have not been identified but insight into how electrical asymmetry in gap junctions may arise has recently been offered by Barrio et al. (18). They found, by expression of exogenous RNAs in *Xenopus* oocytes, that the heterotypic junctions formed between an oocyte that expresses only Cx26 and another that expresses only Cx32, display a fast  $V_j$ -dependence not present for these connexins in their homotypic configurations. These results imply that gating by hemichannels can differ depending on the identity of the hemichannels to which they are joined. In contrast, the Cx38/Cx43 heterotypic junction described by Swenson et al. (28) behaved as a composite of Cx38 and Cx43 homotypic junction behavior. These hemichannels appeared to operate identically regardless of whether they were paired with Cx38 or Cx43; and the hemichannel interactions that occurred in Cx38/Cx38, Cx43/Cx43, and Cx38/Cx43 appeared to be functionally equivalent.

We developed a novel technique for the creation of gene chimeras and used it to explore the basis for the difference in voltage dependence in homotypic and heterotypic junctions of Cx26 and Cx32 (28a). Because the interaction between hemichannels is likely to be mediated by contacts between the extracellular loops, we examined the possibility that the unpredicted electrical asymmetry of the heterotypic junctions arose from mismatching of the extracellular loops. Our initial approach was to make a chimera in which the first extracellular loop of Cx32 was replaced with the first extracellular loop of Cx26, and to pair it with Cx26 and Cx32. We found that mismatching the first extracellular loops in a heterotypic junction could create rectifying behavior but that it did not fully account for all of the properties of the Cx32/Cx26 junction.

## MATERIALS AND METHODS

### Construction of chimeric connexins

The first extracellular loop (E1) of Cx32 encoded by nucleotides 152–256 (8) was replaced by the corresponding sequence, nucleotides 121–225 of Cx26 (29) by the procedure described by Rubin et al. (28a) (Fig. 2). Briefly, the first extracellular loop of Cx26 was amplified by the polymerase chain reaction (PCR) using two bifunctional oligonucleotide primers: 5'-CTG GTG GTG GCT GCA AAG GAG GTG TGG GGA-3' and 5'-TTG CAG GGA CCA CAG CCG GAT GTG AGA GAT-3'. The underlined portions of these oligonucleotides were complementary to sequences of Cx26 at 5' and 3' boundaries of E1 and served as "forward" and "reverse" primers for the amplification of the intervening Cx26 DNA by PCR. The remaining sequences at the 5' ends of both primers were complementary to the regions of Cx32 that were on either side of the borders of the E1 domain. PCR product was phosphorylated and served as a primer for the in vitro mutagenesis of single stranded Cx32 in standard methods of mutagenesis using a kit supplied by Amersham, (Arlington Heights, Illinois) with modifications as described (28a). The sequence of these clones was confirmed by dsDNA sequencing using Sequenase (United States Biochemical Corp., Cleveland, Ohio). The first two amino acid residues, Lys 41 and Glu 42, of Cx26 E1 were replaced with the amino acids Glu 41 and Ser 42 of Cx32 using a 30 base oligonucleotide primer and the standard in vitro mutagenesis kit.

### Preparation of RNA

RNA was transcribed from linearized plasmids using T7 RNA polymerase. Briefly, synthesis was performed in the presence of the cap

analogue m<sup>7</sup>G(5')ppp(5')G (Boehringer-Mannheim, Indianapolis, Indiana) at a 10:1 ratio to added rGTP for 2 h at 37°C under standard reaction conditions followed by an additional 5 min synthesis in the presence of equimolar rGTP to ensure full length transcription of initiated capped transcripts. Integrity of the synthesized RNA was determined by electrophoresis through a 1% agarose gel.

### Expression of junctional currents in pairs of *Xenopus* oocytes

Adult female *Xenopus laevis* frogs were purchased from Xenopus I, (Ann Arbor, Michigan) and maintained at 18°C in a 12 h L/D cycle. Defolliculated oocytes were placed in ND96 medium containing 1.8 mM CaCl<sub>2</sub>, allowed to recover overnight and then coinjected with 50 nl of an aqueous solution containing approximately 1 µg/µl RNA and 0.25 µg/µl of two antisense oligonucleotides 5'-G CTT TAG TAA TTC CCA TCC TGC CAT GTT TC-3' and 5'-TTC CTA AGG CAC TCC AGT CAC CCA TGC TCA-3' that are complementary to endogenous *Xenopus* Cx38 (commencing at nt -5 in the sequence reported in reference 19) and Cx43 (commencing at nt 190 in the sequence reported in reference 30) mRNA, respectively. These antisense oligonucleotides block all endogenous junctional communication within 48 h of pairing (28a; see also reference 18). 24 h post-injection, oocytes were devitellinized manually in hypertonic medium (200 mM K-aspartate, 20 mM KCl, 1 mM MgCl<sub>2</sub>, 10 mM Hepes, pH 7.6, see reference 31) and paired in ND96 medium containing calcium. Junctional currents produced by exogenous RNA were evident 6–8 h after pairing at room temperature.

Junctional conductance was measured with a dual voltage clamp as described by Verselis et al. (13). Data were collected with a PC-AT computer using pCLAMP software (Axon Instruments Inc., Foster

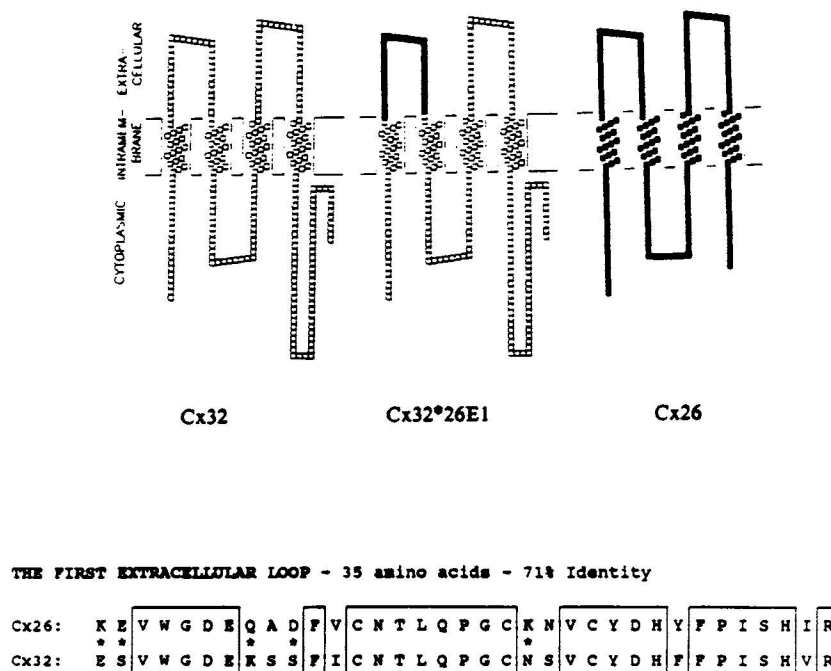


FIGURE 2 Topology of Cx26, Cx32, and the chimera Cx32\*26E1 and comparison of the amino acid sequences of the first extracellular loops of Cx26 and Cx32. Cx32 residues are represented by open boxes and Cx26 residues by solid boxes. Sequences involved in the domain replacement which correspond to amino acids 41–75 of Cx32 and Cx26 are shown below. Changes in charged residues that occurred as a result of the replacement are indicated by asterisks. The domain replacement results in no net change in the overall charge of this domain.

City, California) and a LABMASTER/TL-1 interface combination. Junctional currents were filtered at 1–2 kHz with an eight-pole Bessel filter (AP Circuit Corp., New York, NY). Initial and steady-state  $g_j$  ( $g_{j0}$  and  $g_{j\infty}$ ), were obtained by extrapolating exponential fits of responses to step changes in  $V_j$ . The steady-state conductance,  $g_{j\infty}$  was fit to the Boltzmann relation of the form

$$g_{j\infty} = \{g_{j\max} - g_{j\min}\} / (1 + \exp[A(V_j - V_0)]) + g_{j\min}, \quad (1)$$

where  $g_{j\max}$  is the maximal conductance,  $g_{j\min}$  is the residual conductance approached at large values of  $V_j$ ,  $V_0$  is the voltage at which  $g_{j\infty} = (g_{j\max} + g_{j\min})/2$ , and  $A = zq/kT$  is a constant expressing voltage sensitivity in terms of the number of equivalent gating charges,  $z$ , moving through the entire applied field, where  $q$  is the electron charge, and  $k$  and  $T$  have their usual meanings. The heterotypic junctions display an initial fast  $V_j$ -dependence of  $g_j$ , the time course of which was not resolved by the voltage clamp. These initial values of  $g_j$ ,  $g_{j0}$  were fit by the Boltzmann relation of Eq. 1, but with different parameters. For analysis of the subsequent slow changes in  $g_j$ , it was assumed that the fast and slow processes operated independently and in series. Thus, Eq. 1 was fit to the ratio  $g_{j\infty}/g_{j0}$ . Both  $g_{j0}$  and  $g_{j\infty}$  changed with  $V_j$  in the heterotypic pairs. The results are plotted as  $G_{j0}$  and  $G_{j\infty}$ , values of  $g_j$  normalized to  $g_j$  at  $V_j = 0$  to allow comparison of the conductance-voltage relations of different junctions.

## RESULTS

### Voltage dependence of junctions formed with Cx26 and Cx32

The steady-state conductance-voltage ( $G_{j\infty}/V_j$ ) relations for homotypic and heterotypic junctions of Cx32 and Cx26, as determined in the *Xenopus* oocyte expression system, have been reported (18, 28a) and are presented for reference in Fig. 3, *A* and *B*. The smooth curves are fits to the Boltzmann relation described by Eq. 1 and are based on data from Table 1. Also shown, in Fig. 3 *C*, is a hypothetical  $G_{j\infty}/V_j$  relation of heterotypic junctions formed by Cx32 and Cx26. Homotypic junctions formed from Cx32 (Fig. 3 *A*) display symmetric reduction in steady-state junctional conductance about  $V_j = 0$  mV. Homotypic junctions formed by Cx26 are characterized by a more complex voltage dependence (Fig. 3 *B*). In addition to slow decreases in  $G_{j\infty}$  apparent for large values of  $V_j$ , of either sign, initial currents display a small degree of fast rectification whose time course cannot be resolved. This fast rectification increases  $G_{j0}$  for depolarization and decreases  $G_{j0}$  for hyperpolarization of either cell and thus depends on  $V_{i-o}$ . For polarizations of  $\pm 100$  mV this fast  $V_{i-o}$ -dependence results in conductance changes of  $\sim \pm 10\%$  of the value for  $G_j$  at  $V_j = 0$  mV (18). The asymmetry of the  $G_{j\infty}/V_j$  relation also indicates a small degree of  $V_{i-o}$  dependence shown in Fig. 3 *B*. The hypothetical curve shown in Fig. 3 *C* was generated with the assumption that the component hemichannels would operate independently and would retain characteristics determined for each connexin in homotypic

junctions. The small degree of  $V_{i-o}$  dependence of the Cx26 hemichannel was omitted. It was further assumed that, like Cx38 (28, 32, 33), Cx26, and Cx32 hemichannels would close for relative positivity imposed on their side of the junctions.

Injection into *Xenopus* oocytes of antisense oligonucleotides directed against the endogenous connexins, Cx38 and Cx43, blocked the formation of endogenous gap junctions between paired oocytes and permitted the unambiguous characterization of macroscopic currents produced by injection of exogenous RNAs (see also reference 18). Heterotypic junctions formed by pairing an oocyte injected with Cx26 RNA with one injected with Cx32 RNA exhibited voltage dependent behavior (Fig. 4 *A*) that was markedly different from that of homotypic junctions composed of these connexins and the predicted behavior illustrated in Fig. 3 *C*.  $G_{j0}$  decreased substantially on hyperpolarization and increased on depolarization of the cell expressing Cx26 producing a somewhat sigmoidal  $G_{j0}/V_j$  relation. The same relation could be obtained by equal and opposite polarizations of the cell expressing Cx32 indicating dependence only on  $V_j$ . We term this process fast  $V_j$ -dependent rectification. The absence of  $V_{i-o}$  dependence was confirmed by equal simultaneous polarizations of both cells (data not shown). The  $G_{j0}/V_j$  relation was fit by the Boltzmann relation (Eq. 1) although in another paper (18) a linear fit was satisfactory. Slow changes in  $G_{j\infty}$  in response to  $V_j$  were also present and were also asymmetric. The effective gating charge of the  $G_{j\infty}/V_j$  relation for Cx32/Cx26 junctions with  $V_j$  relatively positive on the Cx26 side was close to that of homotypic Cx26/Cx26 junctions, but the  $V_0$  was decreased by 25 mV (Table 1). We observed no slow decrease in  $G_j$  for voltages that made the Cx32 side relatively positive. The asymmetry in the  $G_{j\infty}/V_j$  relation of heterotypic junctions results from the apparent loss of slow  $V_j$ -dependence in one of the opposed hemichannels. If Cx32 and Cx26 hemichannels are closed by relative positivity on their side as are Cx38 hemichannels, it would be the slow gating mechanism of the Cx32 hemichannel that was not seen and that of the Cx26 hemichannel that was preserved. The maximum slope of the  $G_{j0}/V_j$  relation was less than that of the  $G_{j\infty}/V_j$  relation for relative positivity on the Cx26 side implying a smaller gating charge. However, because of uncertainty about the asymptotes of the  $G_{j0}/V_j$  relation, the values of  $z$  and  $V_0$  for this process could not be accurately determined. The gating charge estimated for the fast process is in the range of 0.6 equivalent charges. These data are in general agreement with those presented recently by Barrio et al. (18).



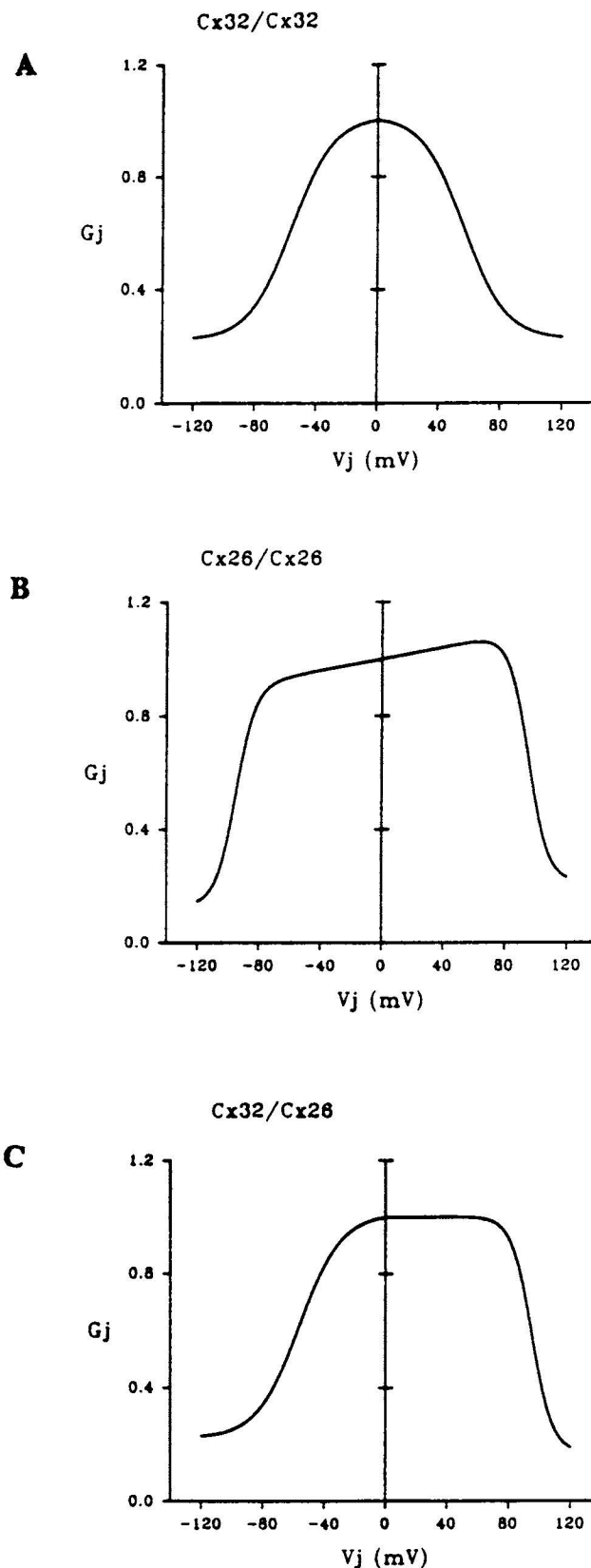


FIGURE 3 Conductance-voltage relations for (A) homotypic Cx32, (B) homotypic Cx26 junctions, and (C) the predicted conductance-voltage relation for heterotypic Cx32/Cx26 junctions. Smooth curves are fitted Boltzmann relations of the form described in the materials and methods. Values for the parameters are those calculated (28a), and are presented in Table 1. The theoretical curve for the conductance-voltage relationship of heterotypic junctions formed by Cx32 and Cx26, C is the relation that would obtain from independent hemichannel gating if, as was shown for Cx38, Cx26, and Cx32 hemichannels close in response to relative positivity on their side of the junction. The small contribution to the conductance-voltage relation that could be made by the inside-outside voltage dependence of the single Cx26 hemichannel has been omitted.

TABLE 1 Boltzmann parameters for homotypic and heterotypic junctions

Connexins	$z$	$A$	$-V_0$	$G_{\min}$	$z$	$A$	$+V_0$	$G_{\min}$
Cx32/Cx32 <sup>‡</sup>	1.8	$0.073 \pm 0.018$	$53.8 \pm 6.4$	$0.22 \pm 0.06$	1.9	$0.075 \pm 0.020$	$60.3 \pm 7.6$	$0.22 \pm 0.07$
Cx26/Cx26 <sup>‡</sup>	4.0	$0.160 \pm 0.029$	$93.1 \pm 6.1$	$0.15 \pm 0.04$	3.8	$0.150 \pm 0.270$	$98.8 \pm 4.5$	$0.19 \pm 0.03$
Cx32/Cx26		No slow $V_j$			3.3	$0.132 \pm 0.009$	$75.7 \pm 3.3$	$0.16 \pm 0.02$
Cx32*26E1/Cx32*26E1 <sup>§</sup>	$\geq 1.5$	$\geq 0.061$	$\leq 96.6$	$\geq 0$	$\geq 1.4$	$\geq 0.056$	$\leq 101.5$	$\geq 0$
Cx32/Cx32*26E1 <sup>§</sup>	$\geq 0.9$	$\geq 0.037$	$\leq 113.8$	$\geq 0$	1.9	$0.076 \pm 0.009$	$88.8 \pm 3.3$	$0.19 \pm 0.02$
Cx32*26E1/Cx26		No slow $V_j$			2.8	$0.112 \pm 0.011$	$76.8 \pm 2.8$	$0.13 \pm 0.05$
Cx32/Cx26*32ES		No slow $V_j$			5.6	$0.222 \pm 0.002$	$66.1 \pm 4.0$	$0.20 \pm 0.04$

Values for  $V_0$  are absolute values for polarity indicated; <sup>‡</sup>Values determined (28a); <sup>§</sup> $G_{j\infty} - V_j$  relation did not approach a asymptote  $G_{\min}$  for  $V_j$  as large as  $\pm 120$  mV. Values are presented as lower limits for  $z$  and upper limits for  $V_0$  with  $G_{\min} = 0$ .

## Voltage dependence of junctions formed with chimeric connexins

The electrical asymmetry observed in heterotypic junctions is likely to arise from interactions of the extracellular loops as these are the only regions of the connexins that are expected to be in contact. We examined the contribution of the first extracellular loop to the voltage dependence of gap junctions by producing a chimeric connexin in which the first extracellular loop of Cx32 was replaced with the corresponding domain of Cx26. This chimeric connexin is designated Cx32\*26E1 or precisely Cx32 (amino acids 1–40)/Cx26 (41–75)/Cx32 (76–283).

Heterotypic pairing of Cx32\*26E1 with either Cx26 or Cx32 resulted in electrically asymmetric junctions with both fast and slow processes and  $G_j/V_j$  relations similar in form to those of Cx32/Cx26 junctions. (We write these junctions as Cx32/Cx32\*26E1 and Cx32\*26E1/Cx26 where for clarity, the connexin on the right hand side of the pairing designation is relatively positive for positive  $V_j$  in the  $G_j/V_j$  plots). The fast  $V_j$ -dependence of the heterotypic Cx32/Cx32\*26E1 junctions increased  $G_{j0}$  for depolarization on the Cx32\*26E1 side (or hyperpolarization on the Cx32 side), but resulted in a  $G_{j0}/V_j$  relation that was less steep than those of all other heterotypic junctions examined (Fig. 4B). The slow  $V_j$ -dependence of these junctions decreased  $G_{j\infty}$  markedly with depolarization of the Cx32\*26E1 side, but unlike the other heterotypic junctions also decreased  $G_{j\infty}$  upon hyperpolarization of the cell expressing Cx32\*26E1. The gating charge and  $V_0$  of the slow  $V_j$ -dependence for relative positivity on the Cx32\*26E1 side were close to those reported for homotypic Cx32\*26E1/Cx32\*26E1 junctions (Table 1). The gating charge and  $V_0$  of the slow  $V_j$ -dependence for relative positivity on the Cx32 side could not be precisely calculated as  $G_{\min}$  was not achieved for this polarity within the 120 mV range of applied transjunctional voltages (Table 1). The values given in Table 1 for this polarity are limits determined by assuming  $G_{\min} = 0$ . Regardless of the uncertainty in  $G_{\min}$ ,  $V_0$  for slow  $V_j$ -dependence in response to relative positiv-

ity on the Cx32 side of these junctions was increased by at least 35 mV compared to that reported for the Cx32 homotypic junction (Table 1). With  $G_{\min} = 0$ , the gating charge of the slow  $V_j$ -dependence in response to relative positivity on the Cx32 side would be 0.9. This would represent a significant reduction compared to the gating charge of either component hemichannel in homotypic junctions. A  $G_{\min}$  of 0.25, the conductance measured at 120 mV for this polarity of  $V_j$ , would correspond to a gating charge of 1.9. If 0.25 was close to the true  $G_{\min}$ , there was no change in gating charge, compared to the homotypic values.

In heterotypic Cx32\*26E1/Cx26 junctions, fast  $V_j$ -dependent increases in  $G_{j0}$  and slow  $V_j$ -dependent decreases in  $G_{j\infty}$  were observed upon depolarization of the cell expressing Cx26 (or hyperpolarization of the cell expressing Cx32\*26E1) (Fig. 4C). The fast  $V_j$ -dependent rectification was less steep than for Cx32/Cx26 junctions but steeper than for Cx32/Cx32\*26E1 junctions (compare Fig. 4, A–C). The fit of the slow  $V_j$ -dependence to the Boltzmann equation was close to that obtained for Cx32/Cx26 junctions (Table 1). As with Cx32/Cx26 junctions, no slow  $V_j$ -dependence was observed when the cell expressing Cx26 was hyperpolarized by as much as 120 mV.

These results indicate that asymmetrical E1 pairing can cause electrical asymmetry, as the  $G_j/V_j$  relation for Cx32/Cx32\*26E1 junctions were similar in form to that of Cx32/Cx26 junctions. However, similar electrical asymmetry was seen in Cx32\*26E1/Cx26 junctions in which only the E1 domain was the same in both hemichannels. The slow decrease in  $G_j$  was attenuated when  $V_j$  was opposite to that causing the fast increase in all pairings, but the reduction was only partial with the Cx32/Cx32\*26E1 junctions.

Additional evidence supporting a role for the first extracellular loop in asymmetry of  $V_j$ -dependence was provided by the behavior of heterotypic junctions comprised of Cx32 and a mutant of Cx26 termed Cx26\*32ES in which Lys 41 (K) and Glu 42 (E) that are contained

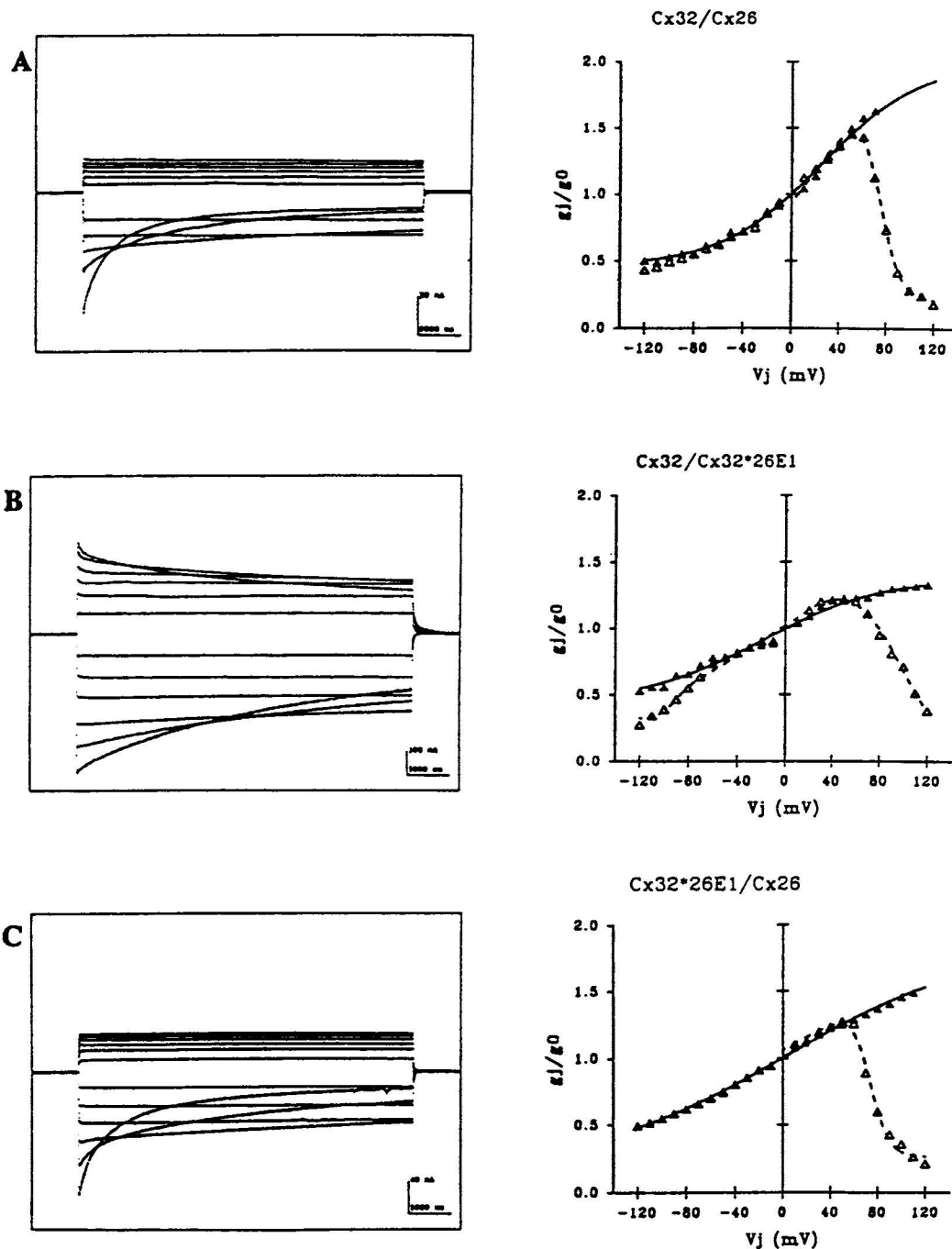


FIGURE 4 Representative junctional currents and initial and steady-state conductance voltage relations for heterotypic junctions formed by (A) Cx32/Cx26, (B) Cx32/Cx32\*26E1, and (C) Cx32\*26E1/Cx26. Junctional currents shown were elicited by  $\pm 20$ , 40, 60, 80, 100, and 120 mV voltage steps applied to the oocyte expressing (A) Cx26, (B) Cx32\*26E1, and (C) Cx26, the connexins indicated on the right side of the pairing designations. Boltzmann parameters are given in Table 1. Calibration bars; (A) 30 nA, 1 s; (B) 100 nA, 1 s; (C) 40 nA, 1 s.

within the first extracellular loop of Cx26 were replaced by the Glu (E) and Ser (S) residues present at these positions in Cx32 and most other sequenced connexins (Fig. 5). The  $G_j/V_j$  relations were again similar in shape

to those of Cx32/Cx26 junctions, but there was a marked increase in the slope of the fast  $V_j$ -dependent rectification and the slope of the slow decrease for relative positivity on the Cx26\*32ES side (for  $G_{jss}$ ,  $z = 5.6$ , Table

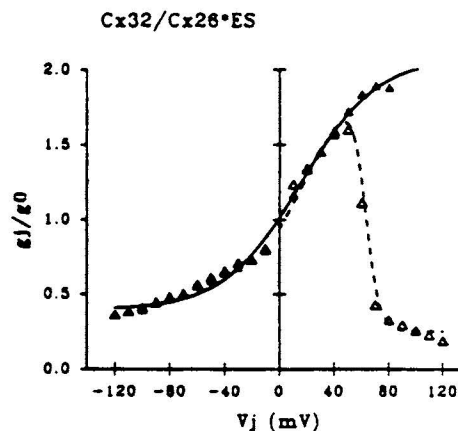
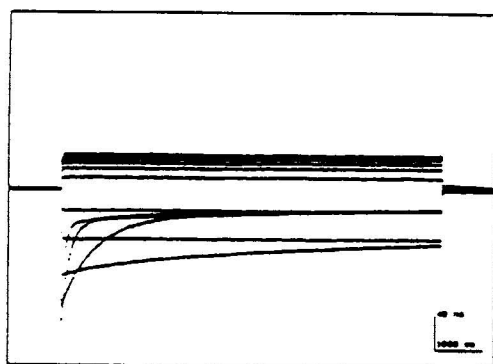


FIGURE 5 Representative junctional currents and initial and steady-state conductance voltage relations for heterotypic Cx32/Cx26\*32ES junctions. Junctional currents shown were elicited by voltage steps, as in Fig. 4, applied to the cell expressing Cx26\*32ES. Normalization and fits are as in Fig. 4. The Boltzmann parameters are given in Table 1. Calibration bar: 40 nA, 1 s.

1). In these junctions, the difference in primary sequence between the E1 domains was smaller than in Cx32/Cx26 junctions, but the asymmetry of the  $G_j/V_j$  relations was greater.

## DISCUSSION

The voltage dependence of Cx32/Cx26 heterotypic junctions (reference 18 and Fig. 4A) differed qualitatively from that of all characterized homotypic junctions and heterotypic junctions formed by pairing *Xenopus* Cx38 with rat Cx43 (28). The voltage dependence of heterotypic Cx38/Cx43 junctions was consistent with the hemichannel properties inferred from homotypic Cx38/Cx38 and Cx43/Cx43 junctions. Each hemichannel appeared to act independently in that there was no indication that either hemichannel influenced the gating properties of the other. In contrast, heterotypic junctions formed by pairing Cx32 with Cx26 exhibit novel voltage dependent properties that would not have been predicted from the behavior of Cx26 and Cx32 homotypic junctions. Heterotypic Cx32/Cx26 junctions exhibited a fast  $V_j$ -dependent rectification that was not seen in either homotypic pair. Although a small degree of fast voltage dependence was described in Cx26/Cx26 junctions, it was sensitive to  $V_{i-o}$ . The slow  $V_j$ -dependence of Cx32/Cx26 junctions was markedly asymmetric, and was present only when the cell expressing the Cx26 hemichannel was made relatively positive. There was a conspicuous absence of slow  $V_j$ -dependence in response to transjunctional voltages that would be expected to close the Cx32 hemichannel if, as demonstrated for Cx38, it were to close in response to relative positivity on its side

of the junction. The slow  $V_j$ -dependence present when the Cx26 hemichannels were made relatively positive resembled that observed for Cx26 homotypic junctions. These data suggest that the slow gating process in only the Cx26 hemichannel was operational within the range of applied transjunctional voltages in these heterotypic junctions. Unlike heterotypic Cx38/Cx43 junctions, protein interactions between the hemichannels of heterotypic Cx32/Cx26 junctions appear to alter their gating properties.

Consideration of the inferred membrane topology of gap junctions suggests that contact between the extracellular loops in the intercellular gap would mediate any interactions between the hemichannels. We used domain replacement to examine the interactions between the first extracellular loops of paired hemichannels. Heterotypic Cx32/Cx32\*26E1 junctions differ from homotypic Cx32/Cx32 junctions by the presence of the first extracellular loop of Cx26 in one of the component hemichannels. The appearance of fast  $V_j$ -dependent rectification in these junctions suggests that mismatch of E1 domains contributes to the observed electrical asymmetry of Cx32/Cx26 heterotypic junctions. The steepness of the fast process in Cx32/Cx32\*26E1 junctions was less than that observed for Cx32/Cx26 junctions and, as is discussed below, slow  $V_j$ -dependence was observed for both polarities of  $V_j$  indicating that mismatch of the first extracellular loops is not the sole cause of the novel behavior of Cx32/Cx26 junctions. If mismatching of E1 alone had led to complete replication of the Cx32/Cx26 properties, it would have been expected that Cx32\*26E1/Cx26 junctions, in which E1 was matched, would not rectify. Such complimentary results would have implied that E1 mismatch made a modular



contribution to rectification, a contribution that was not dependent upon interactions with other domains and was identical whether the E1's were part of Cx26 or Cx32. Mismatching of E1 alone, however, did not result in fully rectifying junctions and conversely, junctions formed with matched E1's, Cx32\*26E1/Cx26, did have properties similar to Cx32/Cx26 junctions. Thus, it would appear that mismatching of other domains, in addition to E1, is necessary for the full expression of Cx32/Cx26 rectification. The requirement for multiple domains could signify that contributions to voltage dependence will be made by more than one domain acting additively or through interdomain interactions that are necessary to generate the appropriate protein conformations for rectification. If mismatch of E1 were responsible for the generation of rectification, then the tertiary structure of the E1 domain must have been altered by interdomain interactions when it was part of Cx32 in order to account for the fact that Cx32/Cx32\*26E1 junctions were not electrically identical to Cx32/Cx26 junctions. If distortion of the Cx26 E1 structure was induced by interaction with other domains, then the rectification observed in Cx32\*26E1/Cx26 junctions could have arisen by E1 mismatch.

The distinction between modular contributions to rectification and dependence upon interdomain interactions will be distinguished in future experiments in which chimeras composed of substitutions of other domains as individual units and in combination will be paired with Cx32, Cx26, and each other. Recent results with a chimera in which E2 of Cx32 has been replaced with E2 of Cx26 and another chimera in which both of the extracellular loops of Cx32 have been replaced with the extracellular loops of Cx26 have indicated that Cx32/Cx32\*26E2 heterotypic junctions do not rectify and that Cx32/Cx32\*26(E1 + E2) heterotypic junctions behave identically to Cx32/Cx32\*26E1 junctions. Thus, rectification like that observed for Cx32/Cx26 cannot be generated by mismatching only the extracellular loops (data not shown). This result suggests that the Cx26 E1 domain can function in the same manner in the presence of Cx32 E2 or Cx26 E2 and that nonextracellular loop domains are important in rectification.

The steepness of the fast process was greater in Cx32/Cx26\*32ES junctions than in the other heterotypic junctions studied. The increased steepness of the fast  $V_j$ -dependent process in these heterotypic junctions, which differs from these Cx32/Cx26 junctions by only two amino acids in the E1 domain of Cx26, further implicates this domain's involvement in the observed rectification. The mutation made the primary amino acid sequence of the E1 domain of Cx26 more like that of Cx32 but rather than reducing the degree of electrical asymmetry, the asymmetry was markedly enhanced. It

remains to be determined how the mutation altered the nature of hemichannel interactions.

The conductance-voltage relationships of these gap junctions were fit to a Boltzmann relation to generate values for the parameters  $V_0$ ,  $z$ , and  $G_{\min}$ . These parameters can be used to compare the voltage dependent behavior of different connexins regardless of the complexity of the actual transitions between open and closed states. In both Cx32/Cx26 and Cx32\*26E1/Cx26 heterotypic junctions, the gating charge of the slow  $V_j$ -dependence observed when the cell expressing Cx26 was made relatively positive was very similar to that observed in Cx26/Cx26 homotypic junctions (Table 1), but in both heterotypic junctions,  $V_0$  was less by  $\sim 25$  mV. When the cell expressing Cx32\*26E1 in the pairs forming Cx32/Cx32\*26E1 junctions was made relatively positive, the gating charge of the slow  $V_j$ -dependence was also similar to that seen in homotypic Cx32\*26E1/Cx32\*26E1 pairs, but unlike the two preceding cases,  $V_0$  was also about the same. However, when the cell expressing Cx32 in Cx32/Cx32\*26E1 junctions was made relatively positive (by convention, negative  $V_j$ ) the  $V_0$  of the slow decrease in  $G_{j\infty}$  was increased by as much as 50–60 mV relative to the value characteristic of Cx32 homotypic junctions. These changes suggest that interactions between hemichannels can significantly shift the  $G_{j\infty}/V_j$  relation along the voltage axis. One interpretation of this shift is that the interactions altered the chemical free energy difference between open and closed states without significantly affecting the gating charge. The large shifts in  $V_0$  caused by heterotypic pairings could explain the electrical asymmetry of slow  $V_j$ -dependence in heterotypic junctions, if the  $V_0$  for the Cx32 side positive was larger than the test pulses applied.

The fast  $V_j$ -dependence could arise from either voltage dependent gating or changes in single channel conductance. These possibilities should be distinguishable by single channel studies of the rectifying junctions. In the present study the fast changes in conductance have been treated as voltage dependent gating. The conservation of the gating charge of the unattenuated slow  $V_j$ -dependent process and the apparent differences in the steepness of the fast rectification in these junctions suggests that these two  $V_j$ -dependent processes arise by separate mechanisms. We recently demonstrated that the differences in the calculated gating charges for slow  $V_j$ -dependence in Cx26 and Cx32 cannot be accounted for by differences in the sequences of their first extracellular loops (28a). Thus, the generation of fast  $V_j$  sensitivity by first extracellular loop interactions may occur independently of their role in slow  $V_j$ -dependent gating. Interactions between the two distinct  $V_j$ -dependent processes may be reflected in the correlation between the degree of attenuation of the

slow  $V_j$ -dependent process for negative  $V_j$  and the steepness of the fast rectification. The heterotypic Cx32/Cx26\*26E1 junctions were the only heterotypic junction in which slow  $V_j$ -dependent gating was observed in response to  $V_j$  of either polarity, and they also exhibited the shallowest slope for the fast rectification. A correlation between the fast and slow processes is also suggested by the behavior of Cx32/Cx26\*32ES junctions in which the steepness of both the fast and slow  $G_j/V_j$  relations was increased by the mutation.

We have used the domain replacement procedure to construct a chimeric connexin in which the first extracellular loop of Cx26 was substituted for the corresponding region in Cx32. The properties of junctions formed by heterotypic pairings of this chimera demonstrated that the E1 domain contributes to the fast  $V_j$ -dependent rectification of heterotypic junctions, but that other domains are involved. Substitution of two charged amino acids in Cx26 with two different residues present in other sequenced connexins had a marked effect on the steepness of the slope of both the fast and slow  $V_j$ -dependent processes in heterotypic junctions with Cx32. This result suggests that E1 may contribute to the formation or operation of the voltage sensing mechanisms underlying both fast and slow  $V_j$ -dependent processes.

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## DISCUSSION

*Session Chairman:* Alan Finkelstein *Scribes:* Han-qing Xie and Marc J. Glucksman

OLAF ANDERSEN: What is the basis for the asymmetry of the Cx26 gating curve for the symmetrical channel? That is, why does the inside-out voltage-dependent process produce asymmetry?

JOSHUA RUBIN: There are two independent processes interacting to produce the asymmetry. The steady-state curves reflect the presence of a fast  $V_{i-o}$  dependence and a slower  $V_j$  dependence.

Asymmetry is a result of a process that is dependent on the membrane potential. One oocyte is kept at a fixed potential while the other oocyte is stepped to various potentials. The current flow across the membrane is measured and indicates that there is a fast change in conductance as a result of changes in the membrane potential. Depolarization of the cell causes a fast increase in conductance and hyperpolarization causes a fast decrease in conductance. On top of this there are slower changes in conductance arising from the  $V_j$  dependence.

ANDREW HARRIS: If you include all of the data from stepping one cell then stepping the other and plot against to  $V_j$ , would you get a symmetrically shaped curve?

RUBIN: Yes.

ALAN FINKELSTEIN: Let me clarify some terms:  $V_j$  refers to the potential difference between the two cells. If for example there is a 20-mV difference between cell 1 and 2, the absolute voltage does not matter.

There is another voltage dependent factor  $V_{i-o}$ , the potential difference between the cell and the external medium. There is no potential difference between the cells, and you change the potential difference between the inside and outside of the cells. In some gap junctions the change in coupling between the cells is a consequence of this transmembrane potential.

ANDERSEN: Getting back to the issue of hemichannels. One cannot “tear” the gap junctions apart, we are told that they can dissociate by incubation in hypertonic solutions, which may be analogous to the two different ways you can open a zippered jacket.

(a) If you pull in a direction perpendicular to the zipper, you will tear the fabric.

(b) If you unzip from the end you will have two hemizippers.

DAVID SPRAY: We looked for that “zipper” for a long time. Many people would agree that there is no stoichiometric result in hemichannels in gap junction preparations.  $V_{i-o}$  dependence is an uncommon property found only in some vertebrate connexins. So what is being measured? Is it the field across one or another extracellular loops?

RUBIN: We have exchanged both extracellular loops and have not changed the  $V_{i-o}$  voltage dependence. The ES mutant also has inside-out voltage dependence. The extracellular loops may have a role in  $V_{i-o}$  dependence but they certainly cannot by themselves confer this kind of voltage dependence.

JOE MINDELL: If you normalize out the fast  $V_{i-o}$  in the symmetric Cx26 junction, is the slow process becoming symmetrical?

JOSHUA RUBIN: Yes.

GERHARD DAHL: In connexins, there is no equivalent to the S4 segment of other voltage-gated ion channels. In fact, all connexins known today have the same set of a few charged amino acids in their transmembrane segments, while voltage sensitivity of various connexons differs considerably. Do you dare to speculate where the voltage gate could be located?

RUBIN: In our studies we are trying to identify regions of the molecule involved in voltage gating. There are biophysical data generated by Andrew Harris and David Spray that suggests that a component of the voltage sensor may reside along the channel lining sequences. They studied the time course of changes in conductance in

amphibian blastomeres when they flipped the polarity of  $V_j$  from +100 to -100 mV. It appeared that the open hemichannels could not close until the closed hemichannels first opened, suggesting that with one gate closed the other hemichannel voltage sensor could not see the change in the  $V_j$  field. Another interpretation could be allosteric interaction preventing the open gate from closing so there is no double closure.

We replaced all the predicted pore lining sequences of Cx32 with the sequences from Cx26 and observed no change in the gating charge.

Recently, we have observed the behavior of a mutant in the first extracellular loop, ES mutants (Fig. 2). As Cx26 comes out of the first transmembrane domain, it has a Lys, Glu at the border; all the other connexins have Glu, Ser. We changed these amino acids and now we see a large increase in the gating charge. This suggests that we are in a region of the molecule involved in voltage sensation; no other charges have done this.

**HARVEY POLLARD:** This discussion about the voltage sensor could be broadened to include more than a few of the residues. Indeed, the global ensemble of charges does result in a dipole, which can be acted on by a voltage pulse. A reorientation of the protein could ensue. This could explain how in the absence of a discrete charge site, voltage dependence can still be found.

**RUBIN:** We approached the voltage sensor as if there were discrete functional domains. We have exchanged various regions and there was no difference. There are 1.8 equivalent gating charges for Cx32 and 3.8 for Cx26, this means that only a fraction of charges move with each subunit. The mutant Cx32 closes for relative negativity. Since Cx26 closes for relative positivity, it would seem that somewhere there must be localized charges conferring polarity.

**POLLARD:** If the structural model were correct and you looked at the distribution of charges in the membrane, would there be any asymmetry of charge? Could a vector of some sort be modified by the mutant.

**RUBIN:** The transmembrane charges for Cx26/Cx32 are conserved.

**FINKELSTEIN:** Can you tell us how much of Fig. 2 is based on data?

**RUBIN:** The original evidence is hydropathy plots. Additional evidence of topology is from membrane protection studies and antibody binding studies. These have been done with Cx32 and Cx43 and these two connexins represent two branches of the connexin family tree.

**FINKELSTEIN:** What about evidence for  $\alpha$ -helix structure within the membrane?

**RUBIN:** There was x-ray diffraction data that suggested the presence of a  $\beta$ -sheet, but that may have been due to the tilt of the  $\alpha$ -helices.

**B. VEERAPANDIAN:** What x-ray structures are you referring to?

**RUBIN:** Those of Makowski et al. from two-dimensional (2-D) crystals. Current 2-D x-ray diffraction data has a resolution of 18 Å.

**HARRIS:** For the mechanism of contingent gating proposed by David Spray and myself, it is not necessary for the voltage sensor charges to be inside the pore but only for the field across those charges to be affected by the conductance state of the pore. My question is, what other aspects of the channel behavior were altered in these mutants?

**RUBIN:** In homotypic junctions, those composed of identical hemichannels, the only changes we observed were shifts in  $V_o$ . These

occur with replacement of the first extracellular loop or the cytoplasmic loop. The second loop shows no change in voltage dependence of the chimera.

**HARRIS:** More specifically, in mutants that change voltage sensors, were there also changes in kinetics?

**RUBIN:** The Cx32\*26 KE mutant has a greatly increased gating charge but there were obvious changes in the kinetics.

**DAHL:** Does mutation of the extracellular loops, including domain swapping, affect the efficiency of channel formation? How do the macroscopic junctional conductances compare between wild-type connexins and mutants?

**RUBIN:** Not for the most part. The ES mutant does not appear to make homotypic junctions but will make heterotypic ones with Cx26 or Cx32.

**MARC GLUCKSMAN:** I believe there are more than five connexins that have been cloned. Are there significant homologies beyond the first loop that may play the "other" role in changing the conformation of the channel.

**RUBIN:** There are regions of tremendous conservation. The two extracellular loops are conserved between all vertebrate Cxs as well as the transmembrane domains.

The amino terminus diverges as well as the cytoplasmic domain and carboxyl terminus.

**GLUCKSMAN:** Are there any features of the models of Nigel Unwin (with electron microscopy), or of Lee Makowski (combining EM and x-ray diffraction) to support or to conflict with the heterotypic constructs, that would work at the resolution of those structural models?

**RUBIN:** Unwin predicted a clockwise rotation in the hemichannel as part of the gating mechanism. It occurred to us that asymmetry in slow  $V_j$  dependence could be explained by steric interactions.

**GLUCKSMAN:** Have you tried any single mutations instead of the double mutation of Lys, Glu to Glu, Ser in Cx26?

**RUBIN:** Not yet.

**MARCO COLOMBINI:** I have a comment about the location of the sensor. Any portion of the protein that moves out of the field, whether neutral or not, once you introduce a charge, will act as a sensor. Maybe the protein moves and you have voltage dependence, or maybe somewhere a charge moves.

**RUBIN:** That is right. These charges could be serving as a probe for regions of the molecule that are moving.

**MICHAEL GREEN:** Why is the gating charge so small (is it possible that there are charges of opposite sign, or charges of the same sign moving in opposite directions)?

**RUBIN:** That is possible. There are positive and negative charges conserved in different transmembrane domains. We made mutants that have not been characterized that may address whether multiple ion movements yield a small change in net charge distribution.

**GREEN:** Is this (above) connected to the effect of changing two



charges by changing two amino acids on Cx26 in the chimera with Cx32 (at the end of your paper)?

RUBIN: The opposite occurs too.

GREEN: Can these charges be titrated?

RUBIN: We never tried. Shifts in voltage dependence as a result of a change in the pH have been observed.

THADDEUS BARGIELLO: In many of the chimeras we have made for examining the cytoplasmic loop, we have simultaneously changed the position and number of charges in the molecule without changing the gating charge.

RUBIN: That brings up a good point. The first extracellular loop exchange involves changes in five charged positions but doesn't produce any change in the gating charge. Mutation of two charges within the loop creates a large change in gating charge. Perhaps the whole domain transfer preserved intradomain interactions and these same interactions were disrupted by the two amino acid mutations. This may give us a clue as to the nature of the intradomain interactions that are important in structure and function.

GLUCKSMAN: You mention at several points in your paper that not all of the properties of rectification could be accounted for in the heterotypic junction, could you speculate what other than the first loop may be responsible for rectification?

RUBIN: In Cx32 and Cx26 there are only slow  $V_j$  changes when the Cx26 side is relatively positive. The slow changes have Boltzmann parameters consistent with Cx26 closure. Cx38 has been demonstrated to close for relative positivity so we thought Cx32 was inactivated in this junction. Recent results indicate that Cx32 may close for relative

negativity and so the asymmetry in slow  $V_j$  may arise from simultaneous closure of Cx32 and Cx26 hemichannel slow  $V_j$  gates.

DAHL: Could you comment on the effects of the environment on the voltage gate? When Cx32 is expressed in transfected cells, the voltage gating appears different compared with hepatocytes and oocytes.

RUBIN: The gating charge is identical in all cases, the  $V_o$  has shifted.  $V_o$  in oocyte is 55 mV, in hepatocyte 40 mV, and in transfected cells 25 mV. The  $G_{min}$  may also change.

Oocytes may influence Cx38 gating, which is different in oocytes and blastomeres. If there is an environmental change, it can shift the  $\Delta G_{chem}$ .

VALERIE HU: My question concerns the irreversibility of gap junction formation. How do you explain the decrease in gap junction permeability along the progression of the cell cycle towards G2/M phase? We have observed this and permeability may be metabolically regulated.

DAHL: Turnover of gap junction protein is very fast with a  $t_{1/2}$  of 2–3 h in hepatocytes.

HU: So uncoupling observed as cells go through mitosis is degradation without resynthesis at that time.

DAHL: Cells have sufficient time for a cycle of synthesis and degradation.

HU: Once gap junction channels form irreversibly, are they subject to degradation?

RUBIN: Degradation could occur on the whole channel. In cells, double membrane structures have been observed. You may be degrading full channels and not hemichannels.