Voltage Regulation of Connexin Channel Conductance

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Voltage is an important parameter that regulates the conductance of both intercellular and plasma membrane channels (undocked hemichannels) formed by the 21 members of the mammalian connexin gene family. Connexin channels display two forms of voltage-dependence, rectification of ionic currents and voltage-dependent gating. Ionic rectification results either from asymmetries in the distribution of fixed charges due to heterotypic pairing of different hemichannels, or by channel block, arising from differences in the concentrations of divalent cations on opposite sides of the junctional plaque. This rectification likely underpins the electrical rectification observed in some electrical synapses. Both intercellular and undocked hemichannels also display two distinct forms of voltage-dependent gating, termed V₁ (fast)-gating and loop (slow)-gating. This review summarizes our current understanding of the molecular determinants and mechanisms underlying these conformational changes derived from experimental, molecular-genetic, structural, and computational approaches.

Key Words: Connexin, voltage dependence, rectification, gating

INTRODUCTION

Connexins are tetraspan membrane proteins with intracellular N- and C-termini (Fig. 1) that in mammals are encoded by a 21 member gene family.¹ Six connexin protein subunits assemble to form a hemichannel or connexon. In most cases, hemichannels are assemblages of six identical subunits, termed homomeric hemichannels. Heteromeric hemichannels comprised of different protein subunits can form in cells that express multiple connexin genes, if the assembly and trafficking cellular machinery allows their mixing. An intercellular channel is formed by the head to head docking of two hemichannels. These are termed homotypic channels, when the subunit composition of the docked hemichannels is identical, and heterotypic channels, when the subunit composition of the docked hemichannels is different. In principle, all combinations are possible (e.g., homomeric homotypic, heteromeric homotypic, etc.). However, not all homomeric hemichannels can pair to form intercellular channels. Pairing compatibility between any two given connexons has been mapped to the second extracellular domain (E2), although both extracellular loops are involved in intercellular channel formation.²
Intercellular channels aggregate to form gap junction plaques with characteristic morphology in EM micrographs (Fig. 1). Interestingly, gap junctions were first observed in invertebrates, however, these are encoded by the innexin gene family.\(^3,4\) Innexins share no substantial protein sequence homology with vertebrate connexins; however, they do share structural homology, which imparts functional similarity (both form large pore, voltage-gated intercellular ion channels). Innexins are not found in the chordate genome, nor are connexins found in the invertebrate genome, suggesting that the two families may have evolved from a common but distant ancestral protein. The pannexin gene family,\(^5\) which encodes membrane channels but not intercellular channels apparently, is found in both chordates and invertebrates and is a candidate ancestral gene. Despite lacking sequence homology, another gene family that encodes calcium homeostasis modulator (CALMH1) plasma membrane channels\(^6\) also shares structural homology to connexins, innexins, and pannexins. Interestingly, CALHM1 channels, like connexin channels, are gated by voltage and calcium ions and play an important role in regulating neuronal excitability in response to extracellular Ca\(^{2+}\). Undocked hemichannels have the potential to regulate membrane potential (and hence excitability) in response to changes in calcium, as do CALMH channels.\(^7\) Reductions in extracellular calcium would increase hemichannel open probability, resulting in membrane depolarization that would move membrane potential closer to the threshold voltage required for firing action potentials. Notably, only a small fraction of intercellular channels appear to be active in a junctional plaque.\(^8\) The molecular bases for this property are unknown. Undocked hemichannels also may be organized into plaques, with only a small proportion of channels active; however, this possibility has not been examined to date.

For the most part, connexins form large pore ion channels that span the plasma membrane of single channels. These plasma membrane channels have been termed undocked or unapposed hemichannels. Intercellular channels play a vital physiological function in the heart and brain, where gap junctions form electrical synapses, and in non-excitable tissues, such as the liver and pancreas, where they couple groups of cells to synchronize cellular activity and secretion. Undocked hemichannels play an important role in paracrine signaling.\(^8\) The open probability of undocked hemichannels is low at the resting potentials of most cells and further reduced by the presence of extracellular Ca\(^{2+}\). Undocked hemichannels have the potential to regulate membrane potential (and hence excitability) in response to changes in calcium, as do CALMH channels.\(^7\) Reductions in extracellular calcium would increase hemichannel open probability, resulting in membrane depolarization that would move membrane potential closer to the threshold voltage required for firing action potentials. Notably, only a small fraction of intercellular channels appear to be active in a junctional plaque.\(^8\) The molecular bases for this property are unknown. Undocked hemichannels also may be organized into plaques, with only a small proportion of channels active; however, this possibility has not been examined to date.

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**Fig. 1.** Membrane topology of connexin subunits and channel composition. A connexin subunit is composed of four transmembrane domains (TM1, TM2, TM3, and TM4), two extracellular loops (E1 and E2), amino and carboxyl termini (NT and CT, respectively), and a cytoplasmic loop (CL). Six connexin subunits assemble to form a hemichannel or connexon, which is termed homomeric if all subunits are identical and heteromeric if comprised of different connexin subunits. Head to head docking of two connexons forms an intercellular channel, aggregates of which are termed gap junctions and visible as plaques in freeze fracture and transmission electron microscopy. Homotypic intercellular channels are formed by docking two hemichannels of identical subunit composition, be it homomeric or heteromeric. Heterotypic intercellular channels are formed by docking hemichannels that differ in subunit composition.
Connexin Voltage Dependence

Membrane potential is an important parameter that regulates the conductance of both intercellular channels and undocked hemichannels. Strictly speaking, connexin channels display two forms of voltage dependence: 1) Instantaneous change in conductance that arises as a consequence of current rectification through single channels (i.e., the rate of ion permeation is a non-linear function of voltage). This is a property of the open channel and channel substates\(^{19,20}\) that occur without conformational changes. 2) Voltage-dependent changes in channel open probability that arise as a consequence of changes in protein conformation (i.e., allosteric transitions between open and closed states that are driven by voltage).

**Rectification of ionic currents**

Although considerably less steep, the rectification of initial currents and single channel currents in symmetric salt, observed in the heterotypic Cx32/Cx26 junction (Fig. 2), qualitatively resembles that of a rectifying electrical synapse.\(^{21,22}\) Furshpan and Potter\(^{21}\) hypothesized that the junctional membrane behaved as an “electrical rectifier” or diode rather than a simple electrical resistor. The separation of fixed positive and negative charges across the junctional membrane would form a diode (p-n junction). P-n junctions can result in steeply rectifying current-voltage (I-V) relations, with steepness increasing markedly as greater amounts of positive and negative are segregated. Thus, they provide an attractive model for the molecular underpinning of steeply rectifying electrical synapses.

We\(^{19}\) examined the diode hypothesis by determining the molecular basis of electrical rectification displayed by Cx32/Cx26 heterotypic junctions. The results of the molecular-genetic studies supported the hypothesis by demonstrating that charged amino acid residues at the amino terminus and first extracellular loop are major determinants of the rectification of the heterotypic Cx32/Cx26 junction. The rectifying I-V relation of the heterotypic junction results from an asymmetry in the position and number of fixed charges present in the Cx32 and Cx26 hemichannels; indeed, the heterotypic channel forms a p-n junction with Cx26 side contributing the negative and Cx32 contributing the positive elements.

The significant concept arising from this study was that rectification of ionic currents requires some form of asymmetry. The asymmetry could be produced by differences in the distribution of fixed charges in the pore\(^{14,23,24}\) or by differential post-translational modification (e.g., phosphorylation or acetylation) of one element in a homotypic intercellular channel that forms the rectifying synaptic junction. Rectification of ionic current can also result from channel block, as exemplified by inwardly rectifying potassium channels, where outward currents are reduced by high-affinity channel block by endogenous polyamines and/or magnesium ions. In this case, the asymmetry reflects a difference in the concentrations of the blocking particles on opposite

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(most with pore diameters of about 15 Å); remarkably, however, there is no strict correlation between single channel conductance, charge selectivity, and permeability to larger molecules.\(^{10}\) In general, connexin channels are permeable to ions, second messengers, (IP\(_3\), cAMP, and Ca\(^{2+}\)),\(^{11}\) and large molecules (in some cases –1 kDa), probably including small regulatory RNA molecules;\(^{12}\) nevertheless, these properties differ according to the subunit composition of the channel.\(^{13}\) Single channel conductance of intercellular channels varies widely (from ~5 to 300 pS), and charge selectivity ranges from slight anion selectivity to moderate cation selectivity.\(^{14,15}\) Notably, connexin channels do not discriminate among small metallic cations or anions. In most cases, the permeability ratios of metallic ions (e.g., Pd/Pb) can be predicted from their mobility in bulk solution. This suggests that ions are fully hydrated when they traverse the connexin channel pore.

Connexins are expressed in virtually all organ systems and not surprisingly, mutations in connexin genes underlie several human diseases. At present, mutations in 10 connexin genes are known to underlie the etiology of at least 15 human hereditary diseases.\(^{16,17}\) These include myelin related diseases, Charcot-Marie-Tooth (Cx32) and Pelizaeus-Merzbacher-Like disease (Cx47); both non-syndromic deafness and syndromic deafness associated with lethal skin disease, such as keratitis-ichthyosis-deafness syndrome (Cx26, Cx30, and Cx31); skin disease not associated with deafness, erythrokeratoderm (Cx30.3); cataract formation (Cx46 and Cx50); cardiovascular disease (Cx40 and 43); and oculodentodigital dysplasia (Cx43). Disease causing mutations can be grouped as loss of function or gain of function. A large category of loss of function mutations result from large shifts in voltage-dependence, resulting in channel closure at physiological conditions where connexin channels would normally be fully open.\(^{15,18}\)
Recent studies have shown that most mammalian electrical synapses are formed by homotypic pairings of Cx36. While most mammalian electrical synapses do not rectify (i.e., they display linear current-voltage relations), the molecular basis for rectification, when observed, results from asymmetries in the concentrations of intracellular magnesium and ATP on the post and presynaptic sides of the synapse. Mg$^{2+}$ and other divalent cations have been shown to interact with a charged residue, D47, located in the channel pore to block the channel, and this is likely to underlie rectification of some electrical synapses formed by Cx36. However, the electrical synapse found between auditory af-

Voltage-dependent gating

Voltage-dependent gating of ion channels is an allosteric process in which channels undergo structural changes in response to voltage to allow or prevent ion flux. Voltage (potential energy), does work on the molecule by changing the position (kinetic energy) of charged amino acids that in turn are coupled to the permeation barrier that blocks ionic flux when the channel resides in the closed state.

As intercellular channels span the plasma membrane of

![Fig. 2. Representative current traces and conductance-voltage plots of initial and steady state junctional currents obtained from pairs of Xenopus oocytes. (A) Cx32 homotypic channels. (B) Cx26 homotypic channels. (C) Cx32/Cx26 heterotypic channels. Initial conductance (▼) and steady-state conductance (▼) are plotted as the function of the applied V relative to the cytoplasm of the right-side hemichannel. Taken with permission from Oh, et al. J Gen Physiol 1999;114:339-64.]

![Graph A](image)

![Graph B](image)

![Graph C](image)
Connexin Voltage Dependence

two adjacent cells, they can in principle be sensitive to two different electric fields: 1) that created by application of transjunctional voltage \( (V_j) \), which is defined as the difference in electric potential between the interior of two coupled cells, and 2) that created by the inside-outside voltage (termed \( V_{i-o} \) or \( V_m \)), which is defined as the difference in electric potential of the cytoplasm and the extracellular space (Fig. 3). In the case of undocked hemichannels, \( V_j \) is equivalent to changes in membrane potential in that a component of the electric field falls across the channel pore. These features of connexin voltage gating have been detailed by Bargiello and Brink.\(^{30}\)

The conceptual framework underlying our understanding of voltage-dependent regulation of gap junction open probability is based on two papers from the Bennett lab,\(^{31,32}\) that described the equilibrium and kinetic properties of gap junction channels found in amphibian blastomeres. The major findings of their studies that are applicable to the majority of vertebrate connexins include the following:

1) Most vertebrate connexins are sensitive only to \( V_j \); Cx26 gap junctions appear to be an exception to this generalization, as the initial conductance of these channels displays weak sensitivity to \( V_{i-o} \),\(^{33}\) at least in macroscopic recordings of junctional currents. Interestingly, single-channel current-voltage elicited by voltage ramps do not rectify.\(^{19}\) In contrast to vertebrate gap junctions, steady-state currents of \( Drosophila \) salivary gland gap junctions formed by innexin protein are markedly sensitive to both \( V_{i-o} \) and \( V_j \).\(^{34}\)

2) Voltage-gating is an intrinsic hemichannel property. Voltage sensors and gates are present in each hemichannel. Application of a given polarity of \( V_j \) favors closure of gates in one hemichannel and opening of gates in the apposed hemichannel. This is a consequence of the head to head docking and serial arrangement of the two hemichannels. This geometry also results in contingent gating, where closed gates in one hemichannel must open before gates can close in the apposed hemichannel when polarity of \( V_j \) is reversed.\(^{30,31}\)

3) As the open probability of homotypic gap junction channels is maximal at \( V_j = 0 \) and decreases with non-zero \( V_j \), voltage must act to destabilize the open state and most likely stabilizes the closed state.

A major advance toward establishing the structure-function relations of voltage-gating in connexin channels was the discovery that connexin hemichannels could open in the plasma membrane of \( Xenopus \) oocytes,\(^{35}\) when unapposed by another hemichannel (undocked hemichannels). This property allowed direct observation of the operation of voltage gates and removed the complication of inferring the op-

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**Fig. 3.** Diagram of a gap-junction channel showing presumed isopotential lines resulting from the application of \( V_{i-o} \) (A) and \( V_j \) (B). \( V_{i-o} \) was established by simultaneously holding the membrane potential of both cells at -50 mV. \( V_j \) does not exist in this case. \( V_j \) was established by voltage clamping the left cell to a holding potential of -100 mV, while the right cell was voltage clamped to 0 mV. Note that the generation of \( V_j \) also generates \( V_{i-o} \) in regions of the channel, but \( V_j \) varies with the voltage paradigm. Holding the left cell at -50 mV and the right cell at -100 mV generates the same \( V_j \) as the previous example (i.e., the voltage difference across the channel is -100 mV relative to the left cell). Note that \( V_j \) would differ in this case, because the absolute membrane potentials used to generate \( V_j \) differ in the two cases. Taken with permission from Bargiello and Brink.\(^{30}\)
eration of serially arranged voltage-gates in intercellular channels.

Remarkably, single channel records of undocked Cx46 hemichannels displayed two distinct forms of voltage-gating, Vj- and loop-gating (also respectively known as fast- and slow-gating). Since this transition passes through a number of meta-stable intermediate conductance states, it has a measurable duration, and consequently, it appears to be a slow-gating event. The duration of the full transition is usually in the range of tens of milliseconds. The term loop-gating was coined by the Verselis lab that postulated that this gating resulted from conformational changes in extracellular loops.

Notably, both gating mechanisms are evident in single channel records of intercellular channels. This indicates that neither process is impaired by constraints imposed on the conformation of extracellular domains when intercellular channels are formed.

Molecular determinants and mechanism of Vj-gating

The molecular determinants of Vj-gating appear to differ among connexins. Two different mechanisms have been proposed to account for channel entry into substates through fast transitions: 1) conformational changes resulting from movement of a voltage sensor located in the N-terminus, which is based on studies of Cx26 and Cx32, and 2) conduction of serially arranged voltage-gates in intercellular channels.

In single channel recordings, Vj-gating is characterized by transitions between the open and one or more sub-conductance states. These transitions cannot be resolved within the time resolution of the voltage clamp; hence the term fast-gating. In contrast, loop-gating is characterized by transitions between the open and a fully closed state; however, since this transition passes through a number of meta-stable intermediate conductance states, it has a measurable duration, and consequently, it appears to be a slow-gating event. The duration of the full transition is usually in the range of tens of milliseconds. The term loop-gating was coined by the Verselis lab that postulated that this gating resulted from conformational changes in extracellular loops.

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a particle-receptor mechanism (ball and chain model) where
the carboxyl terminus (CT) acts as a gating particle by bind-
ing to a receptor located in the cytoplasmic loop (CL),45-48
which is based on studies of Cx43 and Cx40.

Role of the N-terminus in gating polarity of Cx32 and Cx26
The role of the N-terminus in voltage gating arose from stud-
ies that sought to explain why the conductance-voltage rela-
tion of heterotypic Cx32/Cx26 junctions could not be pre-
dicted from the current-voltage relations of the correspond-
ing homotypic channels (Fig. 2).35,49 We43 demonstrated that the asymmetry in the relaxation of steady-state junctional currents was a consequence of the opposite polarity of Vj-
gating in hemichannels formed by the two connexins. Clo-
sure of the Cx26 hemichannel is favored when that side of
the junction is relatively positive, whereas closure of the Cx32
hemichannel is favored when the Cx32 side of the junction is
relatively negative. Thus, application of positive Vj to the
Cx26 side of the junction favors closure of the Vj-gates in
both hemichannels, while application of negative Vj to the
Cx26 side favors opening of Vj-gates in both hemichannels.

Extensive analyses of gating polarity in a series of Cx26/
Cx32 chimeras mapped the differences in gating polarity of
Cx26 and Cx32 hemichannels to a difference in the charge of
the second amino acid. Substitution of the negatively charged
aspartic acid (D2) found in Cx26 for the neutral asparagine
residue (N2) in Cx32 reversed the polarity of Vj-gating of
Cx32 hemichannels from closure favored on relative nega-
tivity to closure favored on relative positivity. Substitution of
neutral and positively charged residues at the second posi-
tion in Cx32 conserved the negative gating polarity of Cx32
junctions, whereas substitution of the negatively charged
aspartic acid with neutral or positively charged residues re-
versed the Vj-gating polarity of Cx26 hemichannel from
closure favored at relatively positive potentials to closure
favored at relatively negative potentials.

Interestingly, charge neutralization of D3 appears to re-
verse the polarity of Vj-gating in Cx46 and Cx50 hemi-
channels.50,51 The Cx50 study51 was based on the asymme-
ty of the conductance-voltage relation of the heterotypic
junction Cx50/Cx50D3N, with closure observed for a sin-
gle polarity of Vj. However, single channel recordings are
needed to confirm the results, because it is known that Cx50
hemichannels display bi-polar fast gating events in single
channel recordings.50

Single channel studies of negative charge and positive
charge substitutions of the second residue in the Cx32*43E1
chimera confirmed that the polarity of Vj-gating was deter-
mimed by the sign of the amino acid residing at the second
position.42,43,52 Subsequent studies demonstrated that nega-
tive charge substitutions up to but not including the 12th resi-
due could reverse the polarity of Vj-gating.42,52 These data
were interpreted to indicate that the first 12 amino acids of
the N-terminus reside in the channel pore, for only in this
position would the domain be able to sense changes in just
transjunctional voltage in intercellular channels. This con-
clusion was supported by structural determinations of Cx26
and Cx32 N-terminal peptides by NMR. These studies dem-
onstrated the presence of a flexible open turn initiated by
the 12th residue (glycine in both Cx26 and Cx32) that would
allow the n-terminal half of the N-terminus to be positioned
within the channel pore.53-55 Demonstration that cysteine sub-
stitutions of the 4th, 5th, and 8th position can be modified by
a MTS reagents strongly support the proposed structure14 and
Oh and Bargiello (unpublished observations). Significantly,
the NMR derived structure agrees well with that of the N-ter-
minus in the crystal structure of Cx26 gap junctions.86

The perfect electrostatic correlation between gating polari-
ty and sign of charges in the N-terminus was interpreted to
indicate that charges in the N-terminus formed at least part of
the voltage-sensor for Vj-gating,43 although movement of the
N-terminus in response to voltage has not been shown direct-
ly. Others58-61 have suggested that the N-terminus forms a
gating particle to block the channel pore in Vj-gating, howev-
er, this possibility needs to be investigated further.37

Oh, et al.44 established that Vj-gating polarity reversal re-
quired only a negative charge at the second position in one
of the six subunits of a connexin hemichannel. Interesting-
ly, these heteromeric hemichannels displayed bi-polar Vj-
gating (i.e., the channel open probability decreases at both
inside positive and negative voltages and is maximal at 0
mV). Presumably, closure at inside positive potentials was
initiated by movement of negatively charged subunits to-
toward the cytoplasmic entrance to the channel pore, while
closure at inside negative potentials was initiated by move-
ment of positively charged subunits toward the cytoplasmic
entrance. The data were interpreted to indicate that Vj-gat-
ing results from conformational changes in individual sub-
units rather than by a concerted movement of all six chan-
el subunits. If gating were concerted (i.e., required the
simultaneous repositioning of all six subunits), then closure
of hemichannels containing negatively charged subunits in
response to inside positive potentials would be opposed by the
tendency of positively charged subunits to move in the
opposite direction and immediately open. Hence, one would expect that the residency in a closed state in these heteromeric hemichannels would be exceedingly brief; the channels would flicker rapidly between open and closed states with both membrane depolarization and hyperpolarization. This was not observed.

Remarkably, several 43E1 homomeric hemichannels containing charge substitutions in the N-terminus also display bi-polar $V_j$-gating. These include, T8D, N2E+G5K, and N2R+G5D. While the mechanism underlying the basis of bi-polar channels has not been established, it can be explained by modeling the $V_j$-voltage sensor as a center open toggle switch. In this scenario channel closure would initiated by either the inward or outward movement of the voltage sensor, such that the voltage sensor crosses one energy barrier for one voltage polarity and the other barrier for the opposite polarity. Interestingly, the charges at the second position appear to have a larger role in determining gating polarity, as voltage sensitivity of bipolar channels correlates with the sign of the charge at this position. However, this class of model cannot adequately explain the bipolarity of heteromeric channels, whose explanation tacitly assumes that the direction of movement of the voltage sensor is identical for all mutations (always toward the cytoplasm). Resolution of the role of the N-terminus requires direct observation of movement of the N-terminus in response to voltage.

In other ion channels, fluorescence quenching, luminescence resonance energy transfer, and fluorescence resonance energy transfer have been used to demonstrate voltage sensor movements. While feasible, application of these methods to undocked connexin hemichannels is technically challenging, as hemichannels appear to aggregate into well dispersed plaques. Furthermore, the results may prove difficult to interpret because only a small fraction of connexin channels contained in plaques appear to be active. Thus, signals reporting movements would probably be very small.

Finally, it is important to understand that the gating model tacitly assumes that the difference in gating polarity of Cx26 and Cx32 hemichannels reflects the oppositely charged voltage sensors, negative in Cx26 and positive in Cx32. Originally, we suggested that the positive charge was contributed by the N-terminal methionine residue. However, characterization of the N-termini of Cx26 and Cx32 (Bargiello, unpublished) by tandem mass spectroscopy have demonstrated that the positive charge of the N-terminal methionine residue is neutralized by acetylation. It is conceivable, that not all methionine residues are acetylated, thus at least one sub-unit carries a positive charge. Alternatively, it is possible that positive charge results from the dipole associated with the $\alpha$-helical N-terminus, or that that residues carry partial positive charges. For example, in the Chemistry at HARvard Macromolecular Mechanics (CHARMM) topology file, atoms HD21 and HD22 in the asparagine side chain together carry a positive charge (0.62) that is balanced by the negative charge of atom ND2 (-0.62). We also cannot rule out the possibility that the N-terminus does not contain the voltage sensor and that charge substitutions in this domain alter the electrostatic profile of the channel and reverse the direction of the voltage drop across conserved pore lining charges that form the voltage sensor. Knowledge of the structure of the $V_j$-closed state would be of great value in clarifying the gating mechanism and the role of the N-terminus.

Role of the CL and CT in fast gating

The ball and chain model (also termed the particle-receptor model) was originally proposed to explain pH gating of Cx43 channels by Delmar and co-workers. The model proposes that changes in pH cause a conformational change in the CL of connexin channels that exposes a receptor domain (L2 region) to which binding of a region of the CT either closes or stabilizes the closed state. The model was later extended to provide a mechanism to explain $V_j$ (fast)-voltage-dependent gating, following demonstration that deletions of the CT of Cx43 and Cx40, or addition of green fluorescent protein to the C-terminus appear to abolish $V_j$-gating.

Revilla, et al. concluded that the CT of Cx32, like that of Cx43, was critical to the expression of $V_j$-gating following truncation of the Cx32 CT to residue 220. However, this was based on their interpretation of kinetic data, which equated a fast component in current relaxations observed in macroscopic recordings of Cx32 intercellular channels to $V_j$-gating. Loss of $V_j$ gates and persistence loop (slow) gates were not confirmed by single-channel recordings. Recently, Kwon, et al. demonstrated that both $V_j$ and loop gates are operational in a truncation mutation that removes all but the first four CT residues (ACAR219) of the Cx32*43E1 hemichannel. Thus, it does not appear that the CT is required for expression of $V_j$-gating in Cx32.

Given the difference in molecular determinants reported for Cx32, Cx26, Cx43, and Cx40 hemichannels, the fundamental questions become whether there a mechanistic relation between the role of the NT and that of the CL-CT interaction in $V_j$-gating and whether fast-gating arises by the
same mechanism in different connexin channels. One possibility, consistent with the view that three domains participate in the same process, is that voltage acts upon charges in the NT to induce conformational changes that include exposure of the L2 receptor in the CL, and that this conformation is stabilized by low-affinity binding of specific sequence contained in the CT. If this mechanism is shared by all connexin hemichannels, then the operation of the \( V_j \) (fast)-gate in the CT truncations of Cx32*43E1 can be explained if the four remaining CT residues (ACAR) are sufficient to interact with a region of the CL near the TM3 border to stabilize the closed conformation. Alternatively, it is possible that the conformation of the Cx32 \( V_j \)-closed state is stable in the absence of interactions with the CT, and thus, the CT would not be required for Cx26 and Cx32, but may be required to stabilize the closed state of Cx43 and Cx40 hemichannels.

**Molecular determinants and mechanism of loop-gating**

Voltage-dependent loop-gating is the mechanism that in conjunction with divalent cations reduces the open probability of undocked hemichannels in the plasma membrane. It has been proposed that the voltage dependent-loop gate is the same as the one closed by chemical gating agents such acidification and extracellular \( Ca^{2+} \), however this is not established and is currently being investigated by both structural and experimental approaches.

In all connexins examined to date, loop-gate closure in undocked hemichannels is favored at inside negative potentials (membrane hyperpolarization). In intercellular channels, loop gate closure is favored on the relatively negative side of the junction. Unlike, \( V_j \)-gating, there are no known mutations or combinations of mutations that reverse loop-gating polarity. In many intercellular channels, the voltage sensitivity of loop gates is less than that of \( V_j \)-gates, such that over intermediate ranges of transjunctional voltages, the plateau observed in conductance-voltage relations of steady-state currents (e.g., Cx32 homotypic junctions) (Fig. 2), is due to entry of a hemichannel into a \( V_j \)-substate, and there is little if any closure of loop-gates at these potentials. At larger \( V_j \)'s, the conductance-voltage relations can decrease to 0 as a consequence of loop-gate closure (see for example, the conductance-voltage relation of Cx32*KE intercellular channels in\(^{68}\)).

Physiologic concentrations of \( Ca^{2+} \) and \( Mg^{2+} \) have been shown to stabilize the loop-gate closed state of Cx46 hemichannels\(^{69}\) and endogenous *Xenopus* Cx38 hemichannels (see Supplemental information in\(^{68}\)). \( Ca^{2+} \) has also been reported to gate connexin channels without application of voltage. This may result by destabilization of the open state as reported by Contreras and co-workers\(^{67,68}\) for Cx26 hemichannels. Barrio and colleagues showed that mutations of Asp169 and Asp178 at the extracellular vestibule significantly reduce the ability of extracellular \( Ca^{2+} \) to close Cx32 hemichannels; however, it is unclear whether this is due to changes in interaction(s) that are involved in stabilizing voltage-dependent loop-gates. Atomic force microscopy studies of connexin hemichannels have indicated a narrowing of extracellular channel entrance with 2 mM \( Ca^{2+} \) (but not \( Mg^{2+} \)) for Cx26\(^{70}\) in the absence of membrane voltage. Divalent cations have been shown to block Cx36 hemichannels by interactions with the conserved pore lining residue D47. Based on the apparent multiplicity in reported actions of extracellular \( Ca^{2+} \), it is likely that there are multiple mechanisms underlying \( Ca^{2+} \) regulation of connexin hemichannels and that different connexin channels may have multiple regulatory binding sites. The stabilization of loop-gates is likely to be just one of several different mechanisms.

The distinctive feature of loop-gating is the slow time constant of the full gating transition from fully open to fully closed states (tens of milliseconds) (Fig. 4). The time course is explained by passage through a series of meta-stable states with occupancy in the fully closed state strongly favored at large negative potentials. A macroscopic recording of Cx32*43E1 N2E ΔCT hemichannels expressed in *Xenopus* oocytes is shown in Fig. 5. Current relaxations at inside positive potentials (red traces) correspond to closure of \( V_j \)-gates, while those at inside negative potentials result from loop-gate closure. For Cx32*43E1 N2E ΔCT the gating charge for loop-gating calculated from limiting slope shown in Fig. 5B is ~0.5. The gating charge of Cx46 hemichannels is ~–2.2 (limiting slope from \( P_{open} \) data of\(^{68}\)). The difference in gating charge could mean that more charge moves through the same electric field in Cx46, or that the electric field differs in the two hemichannels, or both. Knowledge of open and closed structures allows these alternatives to be distinguished.

For Cx32*43E1 N2E ΔCT, the macroscopic current relaxation at -70 mV is well fitted by two exponentials with time constants ~6.46 and 0.83 seconds (0 Mg\(^{2+}\)), 4.19 and 0.55 (1.8 mM Mg\(^{2+}\)), and 4.07 and 0.52 (5 mM Mg\(^{2+}\)). Progressive time constant lengthening as a function of (Mg\(^{2+}\)) is consistent with closed state stabilization. The time constants of Cx46 are shorter, ~1.2 and 0.23 seconds at -70 mV in 1.8
the complexity of the maze increases, the average time taken to successfully transit the maze increases.

Our current understanding of the molecular mechanism underlying loop-gating derives from three sources: 1) experimental studies that defined the structure of the loop-gate closed state; 66,74,75 2) the atomic structure of the Cx26 channel (PDB:2ZW3) 56 and its refinement by all atom Molecular Dynamics (MD) simulation and validation by Grand Canonical Monte Carlo Brownian Dynamics simulation (GCMD/BD); 23 3) identification and computational analyses of van der Waals and electrostatic interactions that stabilize the permeability barrier in the open state.

Mapping the loop-gate closed conformation
Solution of the atomic structure of a native ion channel in a voltage-gated closed state is a challenge common to all voltage-gated channels. One cannot impose a stable, defined voltage gradient to an isolated channel that is currently compatible with structural determination by NMR, electron paramagnetic resonance, or X-ray crystallography. A powerful alternative method is to use biochemical/functional assays to define the distance between substituted cysteine residues along the length of the channel pore. Methods such as state dependent formation of Cd²⁺-thiolate metal bridges, bi-functional chemical cross-linkers, and disulfide bonds serve as “molecular rulers” that define the distance between substituted cysteines when the channel resides in a conformation of interest. The experimental rationale and interpretation of results of Cd²⁺-thiolate metal bridge formation in determining conformational changes has been discussed previously. 37,66,76

Mg²⁺. The time constant of Cx26 is longer, exceeding 10 seconds at -70 mV in 1.8 Mg²⁺. The longer of each pair of time constants most likely corresponds to full channel closures and the shorter time constants to partial closures (“failed transitions”).

According to classical transition state theory (TST), the longer time constant suggests that the energy barrier separating fully open and fully closed states is in the range of 15–17 kcal/mol. Remarkably, undocked Cx46 single channel records from the Verselis lab show loop-gating transitions in response to small membrane depolarizations, as well as membrane hyperpolarization. 36 This indicates that transition between open and closed states can take place in the absence of voltage [i.e., with thermal energy alone (~0.62 kcal/mol at 310 K)], despite the extremely high energy barrier predicted by TST. This suggests that approaches based on TST, which relate rate constants exclusively to the heights of energy barriers, may not provide a suitable conceptual framework to elucidate the mechanisms of conformational change of loop-gating. An alternate conceptual framework is provided by transition path theory (TPT). 72,73 TPT is founded in statistical mechanics and considers the transition-path ensemble in terms of the free energy landscape of different conformational states. It takes into account the likelihood that only a subset of the transient conformational fluctuations of open or closed states has access to paths that lead to successful transitions. TPT interprets rate constants of conformational transitions to reflect not only barrier heights but also the complexity and “ruggedness” of the free energy landscapes. The simplest analogy is to picture a maze: as the complexity of the maze increases, the average time taken to successfully transit the maze increases.

Our current understanding of the molecular mechanism underlying loop-gating derives from three sources: 1) experimental studies that defined the structure of the loop-gate closed state; 66,74,75 2) the atomic structure of the Cx26 channel (PDB:2ZW3) 56 and its refinement by all atom Molecular Dynamics (MD) simulation and validation by Grand Canonical Monte Carlo Brownian Dynamics simulation (GCMD/BD); 23 3) identification and computational analyses of van der Waals and electrostatic interactions that stabilize the permeability barrier in the open state.
The fundamental idea is to trap or “lock” the channel in a given conformation by formation of metal bridges.

Fig. 6 summarizes the results of metal-bridge formation following investigation of cysteine substitutions at 25 loci that span the length of the Cx32*43E1 channel pore. The largest conformational changes are in the most stable region of the channel pore, the 310 or parahelix, formed by amino acids 41–50, where pore diameter is reduced from 15 to 20 Å to less than 4 Å. This reduction in pore diameter is sufficient to form a permeability barrier, given that the diameter of hydrated K+ and Cl− ions is 6.62 Å and 6.64 Å, respectively.77 The data, in conjunction with the atomic structure models of Cx2623,56 and Cx32*43E1 channels,64 show a marked reorganization of the parahelix, with A43C moving into the channel pore with loop-gate closure. Similarly, A40C only coordinates Cd2+ in the loop-gate closed state, a feature which requires straightening of the TM1/E1 bend angle. Straightening of this angle predicts that the intracellular channel formed by residues at the border of TM2 and CL and NT will move closer together. This prediction was confirmed by formation of metal bridges at E109C and L108C66 and at 3 residues in the N-terminus (Kwon, Oh, Bargiello, Dowd, and Bargiello, submitted). The diameter of the channel pore at the intracellular entrance would be reduced to about 10 Å. The extracellular entrance, defined by residue Q56, does not appear to undergo large conformational changes with voltage-gating. We interpret these results to indicate that the loop-gate permeability is essentially focal; in that, conformational changes in the parahelix, but not the channel entrances, are sufficient to prevent ion flux.

Insights into loop-gating from MD simulation

Surprisingly, the largest conformational change in loop-gating occurs in the most stable region of the Cx26 hemichannel pore,78 the parahelix formed by amino acids 42–51 in Cx26. As voltage effects channel closure by destabilization of the open state, elucidation of the atomic interactions that stabilize the permeability barrier formed by the parahelix should provide insights into the mechanism of loop-gating at the atomic level.

The picture of loop-gating that emerges from MD simulations23,78 is that an electrostatic network (emanating from the parahelix) that interconnects all six subunits forms the nucleus of the loop-gate voltage sensor. In Cx26, this includes residues E42, D46, E47, R75, R184, E187, and K188. This charge ring meets the requirement that a connexin voltage sensor must reside in the channel pore where it can sense transjunctional voltage in both open and closed states.
conformation. The MD simulations show how reorganization of the electrostatic network could be directly coupled to the structure of the permeability barrier formed by reorganization of the parahelix and straightening of the TM1/E1 bend angle. The intersubunit connection provided by the electrostatic network suggests a high degree of positive cooperativity, such that changes in conformation in any given subunit will be transmitted to the two adjacent subunits. This suggests that loop-gating most likely arises from the simultaneous movement of all six subunits (i.e., a concerted gating mechanism). We favor a mechanism in which the intermediate states observed in single channel records correspond to the linked electrostatic networks stepping together through a series of intermediate structures, each of which increases pore occlusion.

Finally, Kwon, et al. also described an extensive intrasubunit van der Waals network that stabilizes both the parahelix and TM1/E1 bend and modulates the dynamics of electrostatic interactions among residues that are believed to form the loop-gate voltage sensor. We proposed that because the open state is stabilized by an additive effect of multiple weak vdW and electrostatic interactions, the transition to the closed state most likely proceeds by the sequential destabilization of open-state interactions and formation of new interactions that stabilize the intermediate and fully closed states. This provides a mechanism to surmount the large energy barrier predicted by TST and one that is compatible with application of TPT.

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