Long-term precision editing of neural circuits using engineered gap junction hemichannels

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Summary

The coordination of activity between brain cells is a key determinant of neural circuit function; nevertheless, approaches that selectively regulate communication between two distinct cellular components of a circuit, while leaving the activity of the presynaptic brain cell undisturbed remain sparse. To address this gap, we developed a novel class of electrical synapses by selectively engineering two connexin proteins found in *Morone americana* (white perch fish): connexin34.7 (Cx34.7) and connexin35 (Cx35). By iteratively exploiting protein mutagenesis, a novel *in vitro* assay of connexin docking, and computational modeling of connexin hemichannel interactions, we uncovered the pattern of structural motifs that broadly determine connexin hemichannel docking. We then utilized this knowledge to design Cx34.7 and Cx35 hemichannels that dock with each other, but not with themselves nor other major connexins expressed in the human central nervous system. We validated these hemichannels *in vivo*, demonstrating that they facilitate communication between two neurons in *Caenorhabditis elegans* and recode a learned behavioral preference. This system can be applied to edit circuits composed by pairs of genetically defined brain cell types across multiple species. Thus, we establish a potentially translational approach, ‘Long-term integration of Circuits using connexins’ (LinCx), for context-precise circuit-editing with unprecedented spatiotemporal specificity.

Keywords: neural circuits; editing; connexins; docking; behavior; neuromodulation; electrical synapses
Introduction

Many molecular events at synapses regulate the communication between cells, and studies have linked changes in synchrony across brain circuits to cognitive and emotional behaviors. For example, electrical oscillations in hippocampus and prefrontal cortex synchronize during spatial memory in rats [1], and between amygdala and hippocampus during fear memory retrieval in mice [2]. Moreover, altered long range synchrony has been observed in preclinical rodent models of schizophrenia [3], depression [4], and autism [5], and manipulating synchrony between infralimbic cortex and thalamus has been shown to enhance resilience to acute stress [6]. Though these studies highlight the therapeutic potential for modulating synchrony to drive long lasting changes in behavior, it remains a major challenge to manipulate the activity of precise brain circuits with sufficient precision to elicit synchrony. Both spatial (i.e., the interaction of specific cells in space) and temporal constraints of endogenous neural circuits (i.e., the ongoing integrated activity across multiple circuits) make this goal especially challenging. For example, classic pharmacological therapeutics targeting ion receptors or neuromodulators pathways impact many brain circuits, and emerging brain stimulation-based approaches directly impact the activity of many neuronal populations (i.e., the rate code) and the timing of activity between distinct cells that comprise neural circuits (i.e., the timing code as reflected by synchrony). As such, tools that selectively regulate synchrony within specific circuits in animals have been scarce.

Gap junctions (electrical synapses) enable direct flow of ions and small molecules between two cells and play a prominent role in broadly synchronizing electrical activity in many organs such as the heart and the brain [7]. To achieve such synchrony, gap junction consists of two docked segments called hemichannels, embedded in the membranes of two adjoining cells. Each hemichannel, in turn, is an oligomer consisting of six monomeric proteins called connexins (Cx), of which there are 21 isoforms in humans [8, 9]. Many Cxs can form single-isoform hemichannels that dock with themselves to create homotypic gap junctions (Fig. 1A, left). This common feature limits the potential utility of Cxs as tools to
selectively regulate brain circuit activity in complex organisms. Expression of ectopic Cx proteins that form homotypic synapses would establish connections between individual neurons of the same cell type, though they may operate in distinct neural circuits (Fig. 1A). In such a scenario, information flow would flow orthogonally across otherwise independent circuits. On the other hand, several Cx hemichannels are capable of recognizing hemichannels composed of other Cx protein isoforms and can generate heterotypic, gap junctions (Fig 1A, right) [10, 11]. Thus, we reasoned that an exclusively heterotypic hemichannel pairing would provide substantial control over the direction of information flow between distinct cell types, and that mechanistic characterization of homotypic and heterotypic hemichannel docking may support the rational design of such a pair (Fig. 1A). We also reasoned that knowledge of these mechanisms could be utilized to engineer connexin proteins that were docking incompetent to the major connexins endogenous to the mammalian CNS, a feature that would be essential for any tools developed for precise circuit editing in mammalian species.

*Morone americana* (white perch fish) express two homologs of mammalian neuronal connexin36 (Cx36) – Cx34.7 and Cx35 – that create heterotypic electrical synapses between auditory afferents and Mauthner cells in the CNS [12]. Critically, the Cx34.7/Cx35 heterotypic gap junction exhibits rectification in the Cx34.7 to Cx35 direction [12], with an intrinsic conductance that increases above the threshold (i.e., -20mV) for action potential firing in the brains of mammals [13]. Since this former feature of the Cx34.7/Cx35 heterotypic gap junction enables a preferred directional modulation that mirrors the general operation of chemical synapses, we chose Cx34.7 and Cx35 as our platform to engineer the synthetic electrical synapse. We systematically altered the residues responsible for Cx34.7 and Cx35 docking to identify two mutant hemichannels that exhibit heterotypic but not homotypic interactions, and do not dock with other connexin proteins endogenous to the mammalian CNS. We then deployed this synthetic synapse in C. elegans to demonstrate its functionality and specificity as a neuronal circuit editing tool.
Results

**In vitro analysis of connexin protein docking**

Though the precise interactions that guide hemichannel docking are incompletely characterized for the majority of connexins, structure-function and sequence analyses indicate that the second extracellular loop (EL2) plays the greatest role in hemichannel docking specificity [14]. We developed an *in vitro* approach for rapid evaluation of connexin hemichannel docking [15]. Here, we reasoned that this approach could be used to rapidly screen a library of mutations for their ability to disrupt homotypic docking for the two mutants. Subsequently, a subset of non-homotypic docking Cx34.7 and Cx35 mutants could be screened against each other to identify putative heterotypic docking pairs.

Our rapid screening approach, Flow Enabled Tracking of Connexosomes in HEK cells (FETCH), enables rapid assessment of docking for many combinations of connexin proteins using flow cytometry. Specifically, FETCH utilizes the presence of connexosomes (a by-product of gap junction formation) as an indicator of docking compatibility. Connexosomes are double-bilayer, vesicular structures composed of fully-docked gap junctions that are internalized into each cell contributing to gap junctions, as part of normal turnover (Fig. 1B)[16, 17]. Thus, docking compatibility of different fluorescently-labeled Cxs can be characterized as a function of the fluorescence exchange between cells, mediated by connexosomes [15]. In our assay, individual Cxs are expressed as either GFP- or RFP670-fluorescent fusion proteins in HEK cells. Following an initial expression, HEK cells that are expressing intended Cx counterparts are then co-plated and incubated. This allows apposed Cxs to potentially interact, and fluorescence exchange is evaluated using flow cytometry. Docking is characterized as the proportion of dual-labeled fluorescent cells that develop in a co-plated sample.

To identify potential mutations that destabilize Cx34.7 and Cx35 homotypic hemichannel docking, we rationally introduced seventy and sixty-seven mutations at sixteen positions on both extracellular loops
ELs) of Cx34.7 and Cx35 respectively (see methods for mutant library design). We then quantified homotypic docking interactions of these mutant proteins using FETCH screening (Fig. 1C). Here, our goal was to identify residues that completely disrupted docking; thus, we benchmarked the mutant scores against a heterotypic pairing of human Cx36 and Cx45 proteins that has previously been shown not to yield functional gap junctions [18]. Critically, for this screening, we were agnostic as to whether these mutations directly disrupted docking interactions or indirectly disrupted docking via interference with an upstream process such as folding or trafficking. Homotypic FETCH screening revealed that most mutants retained their docking character; however, several non-homotypic docking mutant proteins were identified. Non-homotypic Cx34.7 mutants included Y78S, Y78T, Y78V, E225K, E225R, L238Y and K222Q, and non-homotypic Cx35 mutants included N56E, Y78V, Y78S, Y78T, E224H, E224K, E224R, and L237Y (Fig. 1C-E)

Next, to identify mutant protein pairs that exhibit exclusively heterotypic docking, we screened the Cx34.7 and Cx35 non-homotypic mutants against each other using FETCH. Here, a successful docking interaction verified that upstream process such as folding or trafficking remained intact for both Cx mutants. Strikingly, we discovered three connexin mutant pairs whose FETCH scores were higher than the FETCH scores observed for wild type Cx34.7\textsubscript{WT}/Cx35\textsubscript{WT} gap junctions. These results provided clear evidence that it was indeed feasible to engineer a connexin hemichannel pairs that demonstrated heterotypic, but not homotypic, docking. Interacting mutant pairs were comprised of Cx34.7\textsubscript{K222Q} with either Cx35\textsubscript{E224H}, Cx35\textsubscript{E224K}, or Cx35\textsubscript{E224R} as counterparts (Fig. 1F).

Since our long-term objective was to develop a modulation approach that would be amenable for use in the mammalian nervous system, we also probed whether the four mutant isoforms docked with endogenous connexin isoforms– specifically Cx36 and connexin43 (Cx43), the major connexins in mammalian neurons and astrocytes, respectively [19, 20]. Using heterotypic FETCH analysis, we observed that none of the mutant proteins interacted with human Cx43 (FETCH=1.3±0.1%; T\textsubscript{96}=0.29;
P=0.61 for Cx34.7K222Q/Cx43 compared to established non-docking pair replicates using a one tailed t-test, with a Bonferroni correction for 20 comparisons; FETCH=0.4±0.1%, 0.5±0.1%, and 0.5±0.1%; T96=1.41, 1.36, and 1.28; P=0.92, 0.91 and 0.90 for Cx35E224H/Cx43, Cx35E224K/Cx43, and Cx35E224R/Cx43, respectively; N=6 replicates for all experimental connexin pairs; FETCH=1.5±0.2%, N=92 for established non-docking pairs; see methods); however, Cx34.7K222Q and Cx35E224H interacted with human Cx36 (FETCH=22.8±1.9%; T91=−24.4; P=3.4×10⁻⁴³ for Cx34.7K222Q/Cx36; FETCH=5.9±1.1%, 0.8±0.1%, and 0.6±0.1%; T96=−5.50, 0.89, and 1.24; P=1.6×10⁻⁷, 0.81, and 0.89 for Cx35E224H/Cx36, Cx35E224K/Cx36, Cx35E224R/Cx36, respectively).

Rational design of exclusively heterotypic and isoform specific Cx34.7 and Cx35 pair using in silico modeling

To address the unintended docking of Cx34.7K222Q with Cx36 determined via FETCH, we utilized homology modeling to rationally design a heterotypically exclusive connexin pairs that does not interact with endogenous Cx43 and Cx36. Briefly, first we developed computational models of the Cx34.7 and Cx35 wild type and mutant, homotypic and heterotypic gap junctions that we had tested in the initial FETCH screen. We then validated the computational model by comparing the key residues predicted to underly hemichannel docking against the initial docking characteristics of the mutants we determined using FETCH. Next, we modeled the docking interactions between Cx34.7K222Q, Cx35E224H, Cx35E224K, and Cx35E224R mutant proteins and Cx36. Finally, we utilized the insights of residue-wise interaction models to design Cx34.7 and Cx35 hemichannels that would exhibit exclusively heterotypic docking and would not dock with Cx36. We then validated the docking characteristics of our designer Cx34.7 and Cx35 proteins in vitro using FETCH (Fig. 2).

Specifically, to model docking interactions between Cx34.7 and Cx35 gap junctions, we ran molecular dynamics simulations of modeled homotypic and heterotypic pairs of wild type and mutant proteins
Our modeling revealed large negative interaction energies involving residues E214, K222, E223, and E225 in wild type Cx34.7 and residues E213, K221, D222, E224 in wild type Cx35 for both the homotypic and heterotypic docking simulations. These large negative interaction energies were suggestive of salt bridges that stabilize both homotypic and heterotypic docking interactions, since results from our initial homotypic FETCH screens showed that mutations of Cx34.7-K222 and Cx35-E224 disrupted docking. Providing further support for this observation, residues in EL2 that had the lowest interaction energies in the wild type pairs showed much higher energy in mutants for which the charge was switched (e.g., Cx35E224R, Cx35D222H, Cx35D222R, and Cx34.7E223K), or for which a smaller residue of the same charge was introduced (e.g., Cx34.7K222H). Integrating these results, we observed a common interaction motif for both Cx34.7 and Cx35 consisting of three negative residues (E214/E213, E223/D222, and E225/E224 for Cx34.7 and Cx35, respectively), and a positive residue (K222/K221 for Cx34.7 and Cx35, respectively, see Fig. 3). Thus, we determined that the four targeted residues indeed play a prominent role in docking interactions for wild type homotypic and heterotypic Cx34.7 and Cx35 gap junctions.

Next, we introduced Cx36 into our computational modeling pipeline to characterize its homotypic and heterotypic interaction principles. Upon simulating the homotypic Cx36 gap junction, we determined the same pattern of interactions observed for Cx34.7 and Cx35, with residues E230, K238, E239, and E241 serving as key interaction sites. Given that the predicted Cx36 interaction motif is identical to that of Cx34.7, it is unsurprising that Cx36 was additionally modeled to interact with Cx34.7 and Cx35 heterotypically. After verifying these heterotypic docking interactions via FETCH (FETCH=11.9±1.2%; T96=-12.93; P=4.7×10^{-23} for Cx34.7/Cx36; FETCH=18.0±2.0%; T96=-18.69; P=4.7×10^{-34} for Cx35/Cx36), we introduced the four Cx mutants identified in our initial FETCH analysis (Cx34.7K222Q, Cx35E224H, Cx35E224K, Cx35E224R) into our computational framework and modelled their modified heterotypic interactions with Cx36.
Mutating K222Q in Cx34.7 disrupted the large negative interaction energies we observed in the homotypic wild type Cx34.7 model, consistent with our findings from FETCH analysis that this mutation impaired homotypic docking (FETCH=0.4±0.1%; T96=1.43; P=0.92 homotypic Cx34.7K222Q). On the other hand, the three negative residues in Cx34.7K222Q continued to show large negative interaction energies with the positive residue of Cx36, explaining the preserved heterotypic docking between Cx34.7K222Q and Cx36 observed via FETCH. Screened mutant Cx35 proteins (Cx35E224H, Cx35E224K, and Cx35E224R) maintained the positive K221 residue that formed strong interactions with the negative residues of Cx36; however, the Cx35E224K, and Cx35E224R mutations induced strong repulsion with the positive K238 residue of Cx36, explaining why these two mutants failed to heterotypically dock with Cx36 in our FETCH analyses. Interestingly, introducing a smaller positive charge at the same position, Cx35E224H restored the interaction with Cx36 in the computational model, and docking in our heterotypic FETCH analyses.

Having determined putative interaction principles underlying the docking of the three connexin homologs (Cx34.7, Cx35, and Cx36), we set out to rationally design a Cx34.7/Cx35 pair that would exhibit isoform-specific, exclusively heterotypic docking. Our initial strategy was to mutate residues at the four positions of our identified docking motif such that one connexin isoform contained all negative charge interactors, and the counterpart connexin contributed all positive interactors. Cx35K221E (all negative charges at motif residues) showed strong repulsions in our homotypic model, and it did not exhibit homotypic docking in FETCH analysis (FETCH=1.2±0.4%; T96=0.35; P=0.64). Additionally, this mutant protein failed to dock with Cx36 and Cx43 (FETCH=1.5±0.1%; T91=0.02, P=0.51 and 1.7±0.2, T96=−0.32, P=0.37 for Cx35K221E/Cx36 and Cx35K221E/Cx43, respectively). Similarly, Cx34.7E214K/E223K/E225K (all positive charges at motif residues) showed strong repulsions in our homotypic computation model and it did not exhibit homotypic docking in FETCH analysis (FETCH=0.2±0.0%; T96=1.76; P=0.96). However, when Cx35K221E was paired with Cx34.7E214K, E223K, E225K for heterotypic FETCH analysis, the two mutant proteins did not dock (FETCH=1.2±0.3; T96=0.37; P=0.64), despite our model predicting strong interaction. Follow-
up confocal imaging analysis of HEK 293FT cells expressing the constructs revealed that Cx34.7\textsubscript{E214K,E223K}K failed to properly localize to the cell membrane (compare Fig. 4D and 4E). We did not test this failed mutant this against Cx36 or Cx43; rather we evaluated an intermediate Cx34 mutant protein that exhibited strong positive charges at only three of the residues, Cx34.7\textsubscript{E214K/E223K}. This mutant showed strong attractive interactions with Cx35\textsubscript{K221E} in our computational model, and it localized to the cell membrane where it docked with Cx35\textsubscript{K221E} as confirmed by FETCH analysis and confocal microscopy (FETCH=35.7±4.1%; \(T_96=-28.11\); \(P=2.0\times10^{-48}\), see Fig. 4F for confocal image). Critically, Cx34.7\textsubscript{E214K/E223K} did not show homotypic docking in our FETCH analysis (FETCH=1.1±0.2%, \(T_96=0.46, P=0.68\)), nor did it dock with Cx36 or Cx43 (FETCH=1.0±0.2, \(T_96=0.58, P=0.72\) and FETCH=0.9±0.1%, \(T_96=0.73, P=0.77\) for Cx34.7\textsubscript{E214K/E223K/Cx36} and Cx34.7\textsubscript{E214K/E223K/Cx43}, respectively). From here on we refer to this rationally designed, exclusively heterotypic and isoform-specific connexin pair, Cx34.7\textsubscript{E214K/E223K} and Cx35\textsubscript{K221E}, as Cx34.7\textsubscript{M1}/Cx35\textsubscript{M1} (designer Cxs version 1.0, from \textit{Morone americana}).

**Synthetic Cx34.7\textsubscript{M1}/Cx35\textsubscript{M1} electrical synapses regulate behavior**

Using our novel designer Cxs, we set out to determine whether Cx34.7\textsubscript{M1}/Cx35\textsubscript{M1} could establish functional gap junctions with electrical properties sufficient to synchronize the activity of distinct neurons that compose a circuit. We capitalized on \textit{C. elegans} for this analysis because the nematode nervous system is composed of well-characterized circuits of individual cells that regulate behavior.

\textit{C. elegans} do not have an innate temperature preference and can thrive in a broad range of temperatures [21]. However, \textit{C. elegans} trained at a temperature in the presence of food will migrate towards that temperature when they are subsequently placed on a temperature gradient without food [21]. This learned behavioral preference is in part mediated by plasticity of the chemical synapse occurring between a thermosensory neuron (called AFD, presynaptic) and an interneuron (called AIY,
postsynaptic) [22]. Critically, plasticity in the thermosensory neuron AFD can be genetically manipulated to affect transmission to AIY, and to predictably code the otherwise learned behavioral preference [23].

As with other invertebrates, *C. elegans* do not express connexins. Thus, ectopic expression of vertebrate connexins result in formation of electrical synapses that are inert to endogenous gap junction proteins. We have previously shown that ectopic expression of Cx36 could be used to edit the thermotaxis circuit, by bypass the presynaptic plasticity mechanisms between thermosensory neuron AFD and interneuron AIY that contribute to the learned temperature preference [23]. As such, these circuit-edited animals show a persistent preference for warmer temperatures (Fig 5A). We therefore used the thermotaxis circuit in *C. elegans* to validate *in vivo* the utility of our engineered gap junction proteins via *de novo* formation of electrical synapses (assessed by calcium imaging) and to recode behavior (assessed by quantitative thermotaxis behavior testing).

We first expressed Cx34.7M1/Cx35M1 cell-specifically in the AFD/AIY pair and examined whether these proteins were capable of reconstituting functional electrical synapses (see supplemental fig. S1 and supplemental table S2). Like Cx36/Cx36, expression of Cx34.7M1/Cx35M1 between the AFD/AIY pair resulted in functional coupling between AFD and AIY, as assessed via calcium imaging (Fig. 5B left and 4C; P<0.0005 using Fisher exact test with an FDR correction). These *C. elegans* constitutively moved towards warmer temperatures when placed on a thermal gradient, again mirroring the animals expressing ectopic Cx36/Cx36 (F7,17.91= 84.99; P<0.0001 using Welch one-way ANOVA followed by Dunnett’s T3 multiple comparisons; p<0.005; Fig. 5B right and 5D). Expression of Cx34.7M1 or Cx35M1 in both the AFD and AIY neurons (i.e., Cx34.7M1/Cx34.7M1 or Cx35M1/Cx35M1) failed to reconstitute an electrical synapse, though homotypic configuration of both wild type connexins (i.e., Cx34.7WT/Cx34.7WT and Cx35WT/Cx35WT) synchronized the two cells facilitated neurotransmission and modulated behavior (Fig 5B-D, Supplementary Figure SX; P<0.0005). Taken together, these findings confirmed that our designer heterotypic gap junction, Cx34.7M1/Cx35M1, exhibited electrical properties sufficient to
synchronize the activity of specific neurons \textit{in vivo}, and rationally recode the behavior. Moreover, our findings confirmed the docking properties we predicted for a range of connexin proteins using our \textit{in vitro} screen and \textit{in silico} studies, since both Cx34.7\textsubscript{M1} and Cx35\textsubscript{M1} failed to exhibit homotypic docking.

**Discussion**

We developed a novel approach we term ‘Long-term integration of Circuits using Connexins’ (LinCx) that can be used to rationally edit brain circuits. This approach employs an engineered connexin hemichannel pair capable of heterotypic, but not homotypic, docking. When expressed in two adjacent cells, these hemichannels compose a heterotypic electrical synapse, facilitating the transfer of activity between them. We chose to engineer LinCx using two connexin proteins found in \textit{Morone americana} (white perch fish), connexin34.7 (Cx34.7) and connexin35 (Cx35). These connexins are expressed in the CNS at pre- and post-synaptic terminals in \textit{Morone americana} and show rectification in the Cx34.7 to Cx35 direction [12]. Our approach is analogous to the strategy we previously deployed in \textit{C elegans}, using Cx36 [23]; however, here our engineering proteins have a preferential direction for facilitating the transfer of electrical activity, and they do not dock homotypically. This latter feature ensures that even when a hemichannel is expressed across a cell type, those cells will not form electrical synapses between themselves (Fig. 1A). Thus, our LinCx system can still be deployed in high order animals, which have many cells per cell type, to target distinct neural circuits.

We employed a two-part strategy to engineer a Cx34.7 and Cx35 pair that solely exhibit heterotypic docking. First, we created a library of Cx34.7 and Cx35 mutants based on sites previously shown to conferring docking specificity, as well as those implicated in docking based on homology modeling from the structures of Cx26[24, 25]. We then subjected these mutants to our novel rapid screening approach to identify the mutations that disrupted homotypic docking for each Cx protein. Next, we combinatorially screened non-homotypic Cx34.7 and Cx35 mutant proteins against each other, to
identify pairs which retained their heterotypic docking character. Second, we developed a computational model of Cx protein docking. We then employed this computational model to probe the docking properties of our mutants identified in our initial FETCH screen and to design new Cx mutants, \textit{in silico}. At each step of our \textit{in silico} design processes, we validated the homotypic and heterotypic docking properties of our designed mutants by directly assay them using our \textit{in vitro} docking screen. Finally, we integrated all of the mechanistic understanding of Cx docking interactions gained through this two-pronged approach to engineer Cx34.7 and Cx35 mutants that show heterotypic, but not homotypic docking. To our knowledge, this exclusively heterotypic docking profile has never been observed for pairs of connexin hemichannels. The functionality of our Cx34.7 and Cx35 mutant pair and their unique docking configuration was validated \textit{in vivo}, in \textit{C. elegans}, where we demonstrated that their expression in two distinct cells can edit the circuit between them and modulate behavior.

The integrated engineering approach we utilized to develop LinCx (see Fig. 2) can likely be deployed to develop a toolbox of connexin protein pairs that exhibit selective docking properties. Future work may also yield novel hemichannel pairs with customized conductance properties, mirroring approaches applied to modify the conductance of invertebrate electrical synapses [26]. Thus, we believe that it may be possible to deploy multiple LinCx pairs in the same animal to simultaneously edit multiple circuits and ultimately regulate brain function. We also optimized the translational potential for LinCx by engineering Cx34.7 and Cx35 to not only disrupt their homotypic docking, but also to disrupt their heterotypic hemichannel docking with other connexin proteins endogenous to the human brain (e.g., Cx36, Cx43, and Cx45). Given the sequence homology/identity of ELs across mammals (see supplemental Figure S2), we believe that our tool can be broadly applicable to other preclinical model organisms including rodents and non-human primates. Critically, LinCx can also be deployed alongside other well established preclinical modulation approaches including optogenetics and DREADDs enabling broad manipulation of brain networks across multiple scales of spatial, temporal, and context resolution concurrently.
Within brain circuits, space is operationalized as the physical boundaries of individual cells, and time is operationalized as the sub-millisecond level electrical changes of those cells. Though many classic electrical stimulation approaches exhibit high temporal precision in their targeting, these techniques often stimulate volumes of brain tissue that include many brain cell-types and local axonal fibers of passage (see Fig. 6A; electrical stimulation). Preclinical approaches such as optogenetics provide a substantial improvement with regards to spatial targeting by enabling the selective stimulation of specific cell bodies (based on their genetic identities). Moreover, optogenetics modulates cells via temporally precise light pulses, maintaining the temporal precision of electrical stimulation.

Nevertheless, both electrical and optogenetics stimulation bear substantial potential to override circuit computations, which integrate space and time across precise cell types, since these approaches are typically utilized under conditions that modulate the activity of many neurons concurrently, and outside of the activity context of their inputs (see Fig. 6A; optogenetics - soma stimulation).

Multiple studies have sought to address this limitation by delivering stimulation within a closed loop framework. In a closed loop framework, neural activity is modulated based the ongoing activity in the brain. For example, in our prior work we modulated the activity of neurons in medial dorsal thalamus while mice were engaged in a classic task used to model the action of antidepressants [6]. When we activated neurons based on ongoing oscillatory activity in infralimbic cortex (an input region to medial dorsal thalamus), we observed an antidepressant-like behavioral effect. Conversely, when we stimulated these cells using two distinct temporal patterns that were untimed to activity in infralimbic cortex, we observed a pro depression-like behavioral effect, or no behavioral affect at all. These observations provided evidence of the importance of modulating neural activity with ‘context precision’ – stimulation time to the appropriate context of brain activity – to regulate behavior. Similarly, studies have shown that stimulating hippocampus at the peak vs. trough of the endogenous theta oscillatory cycle differentially impacts spatial memory [27]. These and other closed loop studies demonstrate that
context precision serves as a critical axis of circuit operation in the brain that is orthogonal to both spatial and temporal precision.

Over the last decade, several strategies have been employed across myriad studies to enhance the context precision of circuit targeting. One such strategy is based on selectively modulating projection neurons, whereby the inputs at a targeted brain site are selectively activated (See Fig. 6A; optogenetics - projection stimulation) [28]. By directly modulating presynaptic nerve fibers, investigators can regulate the context (presynaptic neurotransmitter release) that drives the target cellular response.

Nevertheless, this approach has the potential to change physiological variables that define the context of the presynaptic neuron (and thus the circuit). For example, terminal stimulation can activate axonal collaterals of the presynaptic neuron thereby decreasing the spatial precision of circuit targeting, or it can induce retrograde activation of the presynaptic cell in a manner that disrupts the activation context of that neuron relative to its own inputs. This approach can also drive the activation of non-target circuits since inputs from a brain region can synapse onto multiple distinct cell types.

Other strategies utilized to enhance the context precision of circuit targeting include the stable step function opsins (SSFOs) [29] and designer receptors exclusively activated by designed drugs (DREADDs) [30], which function to increase the resting membrane potential of target cells. SSFOs and DREADDs maintain the cell type specific spatial precision characteristic of initial optogenetic targeting approaches. Moreover, because cells are rendered more likely to fire in response to their input signals under optimal conditions, these approaches provide improved temporal and context precision. Nevertheless, SSFOs and DREADDs can render target neurons more responsive to all their excitatory inputs (including those from non-targeted circuits), raising the potential of circuit-level off-target effects (see Fig. 6A; DREADDs). Thus, there is still demand for neuromodulation approaches that function within the spatial, temporal, and context constraints that together define brain circuit operation.
Like established protein-based modulation tools such as optogenetics and DREADDs, LinCx can target to precise cell types. However, LinCx builds upon these technologies by enabling each hemichannel to be expressed in a different cell type. The hemichannels expressed by these two distinct cell types then integrate in vivo to form an electrical synapse. As such, LinCx offers unprecedented spatial precision compared to optogenetics and DREADDs in that it enables targeting of one of the specific spatial features that constrains circuits (e.g., the structural integration of two distinct cell types). Importantly, no electrical synapses are constituted between distinct cells of the same cell type since we disrupted both hemichannels’ abilities to dock homotypically (Fig. 1A). LinCx is also designed to optimize the context precision of neuromodulation. Because this electrical synapse rectifies and it only forms between the target pre- and post-synaptic neuron, LinCx constrains the modulation of each individual post-synaptic neuron by the endogenous activity of its genetically defined pre-synaptic partner. Moreover, this feature also yields a level of temporal precision that mirrors the precision of endogenous brain activity. Finally, unlike established modulation approaches, LinCx does not require an exogenous actuator such as light, electricity, or an inert pharmacological compound. Rather, LinCx utilizes endogenous brain activity to modulate target neurons, yielding a tool for precise circuit editing.

Direct stimulation of the brain is a well-established treatment for neurological and psychiatric disorders. For example, electrical convulsive therapy (ECT), which delivers energy to the whole brain, has remained the most effective treatment for major depressive disorder for nearly a century [31]. Deep brain stimulation (DBS) of the sub thalamic nucleus is a widely utilized therapeutic for Parkinson’s disease [32]. Nevertheless, both these modalities have important spatiotemporal constraints. ECT can induce short term cognitive dysfunction, likely due the spatially untargeted nature of energy delivery to the brain, ultimately limiting its clinical use. On the other hand, DBS of the subthalamic nucleus has limited impact on the devastating emotional and cognitive symptoms that accompany Parkinson’s disease, likely due to the brain region targeting selectivity [32]. While non-invasive techniques such as transcranial
magnetic and focused ultrasound hold future promise for expanding the therapeutic landscape for brain disorders due to their increased clinical accessibility (these approaches do not require anesthesia/brain surgery), there continues to be demand for novel therapeutic approaches that modulate brain activity within both the spatial and temporal constrains of brain circuit activity.

We believe there is great potential for LinCx as a tool to probe the causal relationship between brain circuit function and behavior, and as a potential therapeutic approach to ameliorate human neuropsychiatric disorders. Moreover, there are many potential applications for LinCx technology that extend beyond the central nervous system. For example, future studies could optimize and deploy LinCx to the neuromuscular junction as a potentially treatment for Myasthenia Gravis, the cardiac nervous system as a potential treatment for arrhythmias, to the splenic nerve to regulate inflammation, the autonomic nervous system as a potential treatment for a range of disorders associated with gastrointestinal dysfunction, or to the integumentary system to ameliorate chronic pain.

**Figure Legends**

**Figure 1: Screening to identify mutant connexin hemichannel pair that exhibits exclusively heterotypic docking.**

A) Schematic outlining the limitation of introducing ectopic wild type connexin hemichannels (pink rectangles) as a method for selectively modulating precise neural circuits composed by brown and light green neurons (left). Note that connexin hemichannels yield off-target electrical synapses between pre-synaptic neurons, and thus off-target modulation of other circuits. Putative strategy for deploying exclusively heterotypic docking hemichannels (green and red rectangles) to selectively modulate precise neural circuits (right). Note the rectification of this putative gap junction. B) Counterpart fluorescently labeled connexin proteins (i.e., C-terminally fused mEmerald or RFP670) are expressed in different cell populations. Cells are then combined and accessed for opposing hemichannel internalization using flow
cytometry (left). Representative flow cytometry plot for wild type Cx34.7/Cx35 pair is shown on top, and representative flow cytometry plot for Cx36/Cx45 pair is shown on bottom (right). Pink square highlights portion of individual cells labeled by two distinct fluorescent proteins. 

C) Schematic of Morone Americana Cx34.7 and Cx35 extracellular loop mutations used to screen for novel, heterotypic exclusive hemichannels. Positions and mutations unique to Cx34.7 and Cx35 are shown in green and red, respectively. Positions and mutations common to both proteins are shown in black. 

D-E) Circular plot showing homotypic FETCH results for d) Cx34.7 and e) Cx35 mutations. Bar graphs show the effect size (portion of dual labeled cells) of homotypic mutant combinations relative to the heterotypic pairing of human Cx36 and Cx45 which fails to dock. Mutations that disrupted docking are highlighted by black arrows. Purple circle corresponds to zero effect size. Black line corresponds to scale bar for effect size. 

F) Heterotypic FETCH results for Cx34.7 and Cx35 mutant protein combinations. Bar graphs show the effect size of homotypic mutant combinations relative to the wild type Cx34.7 and Cx35 pair.

**Figure 2: Integrated approach used to model connexin hemichannel docking.** Our approach consists of seven integrated components: 1) homology model generation, 2) *in silico* protein mutagenesis, 3) embedding of proteins in a lipid bilayer and aqueous solution, 4) system minimization, equilibration, and molecular dynamics simulation, 5) residue-wise energy calculation, 6) *in vitro* protein mutagenesis, and 7) FETCH validation.

**Figure 3: Model of extracellular loop -2 (EL2) residues mediating homotypic and heterotypic hemichannel connexin docking.** A-C) EL2-to-EL2 interactions predicted between wild type Cx34.7 and Cx35 using homology modeling. Residues predicted to form strong attractive/repulsive interactions are highlighted in blue/red respectively (top). Contact plots for EL2-to-EL2 interactions produced by molecular dynamics simulation (middle), and summary of interactions predicted to stabilize hemichannels pairs (bottom). Plots are shown for A) homotypic Cx34.7, B) homotypic Cx35, and C) heterotypic Cx34.7 and Cx35 interactions.
Figure 4: Engineering Cx34.7 and Cx35 mutants to show heterotypic, but not homotypic, hemichannel docking. A-C) Homology models predicting EL2-to-EL2 residue interactions for Cx34.7 and Cx35 mutant hemichannels. Residues predicted to form strong attractive/repulsive interactions are highlighted in blue/red respectively (top). Contact plots for EL2-to-EL2 interactions produced by molecular dynamics simulation (middle), and summary of interactions predicted to stabilize/destabilize hemichannels pairs (bottom). Plots are shown for A) homotypic Cx34.7_E214K, E223K, B) homotypic Cx35_K221E, and C) heterotypic Cx34.7_E214K, E223K and Cx35_K221E interactions (Cx34.7 residues are shown along the y-axis and Cx35 residues are shown along the x axis). D-F) Confocal images of heterotypic connexin pairs D) Cx34.7_WT/Cx35_WT, E) Cx34.7_E214K, E223K/Cx35_K221E, and F) Cx34.7_E214K, E223K/Cx35_E221E expressed in HEK 293FT cells. All Cx34.7 and Cx35 proteins are expressed as mEmerald and RFP670 fusion proteins, respectively. White arrows highlight dual fluorescent labeled vesicles. Note the cytoplasmic localization of Cx34.7_E214K, E223K, E225K in panel E.

Figure 5. Ectopic connexin hemichannels couple C. elegans neurons and recode thermal preference. A) Schematic of the AFD→AIY synaptic communication and expressed temperature preference. The AFD thermosensory neuron has a robust calcium response to warming stimuli. C. elegans raised in the presence of food at 15°C, or animals with a Protein Kinase C (PKC)-1 gain-of-function mutation, move towards cooler temperatures when placed on a thermal gradient (top). Ectopic expression of connexin hemichannels between AFD and AIY results in synchronization of the signal to AIY and promotes warm-seeking behavior (bottom). B) Calcium traces of neurons expressing ectopic connexin hemichannel pairs (left). Baseline AFD and AIY responses are also shown. Each panel depicts the average trace a group (top, data shown as mean±SEM), heatmaps of individual animals (middle), and the temperature stimulus (bottom). Behavioral traces for each group are shown on the right. Traces are shown for C. elegans homotypically expressing wild type connexin hemichannels (pink highlight), heterotypically expressing the mutant pair (tan highlight), and homotypically expressing mutant connexin hemichannels (cyan...
highlight). **C** Portion of animals showing neuronal calcium responses based on the traces shown in B; ***p<0.0005 using Fisher’s exact test for penetrance. Error bars denote 95% C.I. **D** Thermotaxis indices corresponding to experimental groups. Each data point represents the thermotaxis preference index of a separate assay (12-15 animals/assay), with the median for each group plotted denoted by a black horizontal. **p<0.005; ***p<0.0005; Error bars denote 95% C.I.

**Figure 6. Complementary utility of LINCx compared to widely adopted neural modulation approaches in selectively targeting a spatially and cell-type pair defined circuit.** **A** Electrical stimulation activates many cell types and pass-through fibers within a tissue volume. Stimulation also modulates cellular activity independent of the context of input fibers, and potentially drives retrograde activation of cellular inputs. Open-loop optogenetic stimulation of neuronal soma modulates cell-type specific activity independent of the context of input fibers. Optogenetic terminal stimulation enables modulation of connections between brain regions, but potentially activates multiple circuits defined by distinct post-synaptic cell types, and potentially drives retrograde activation of target axons. DREADDs modulate target cells, but potentially modulate the response of target cells to all of their input fibers. **B** LinCx enables selective targeting based on a cell-type defined pair and selective modulation of the post-synaptic cell-type based on the activity context of the pre-synaptic cell.

**Materials and Methods**

**Design of Cx34.7 and Cx35 mutant library**

A semi-rational design approach was used to design the mutant library. Sequence alignments between the *Morone americana* connexins and the connexins for which the most structure-function data existed (Cx26, Cx32, Cx36, Cx40, and Cx43) were performed in ClustalW. Sites identified by previous studies as
conferring specificity for docking were used as well as those identified by homology modeling from the structures of Cx26 [11]. Specifically, we primarily focused on the extracellular loops and four residues at the interface in loop two, KEVE/KDVE (M. americana Cx34.7/Cx35) and one residue of E1. The homologous residues in other connexins had been demonstrated to be highly tolerant to mutation and critical for docking specificity [33]. Mutations were modeled in Swiss PDB Viewer using homology models of Cx34.7 and Cx35 from a Cx26 and Cx32 interface structure so as not to create mutations with obvious steric hindrance. A wide range of substitutions was made for these five residues of interest, including both those intended to introduce compatible electrostatic interactions as well as less likely candidates. Mutations were also created targeting other residues nearby and/or adjacent to these five for which there was some evidence in the literature that they contributed to docking specificity. However, our semi-rational approach was such that not as many variants were tried for these more distal site mutations and the mutations that were made in those sites were more conservative with regard to the steric and electrostatic properties of the change.

**Construct cloning and preparation**

*Morone Americana* Cx34.7 and Cx35 cDNA constructs we initially acquired failed to efficiently express in HEK 293FT cells. Thus, Connexin gene information was procured from the National Center for Biotechnology Information (NCBI, ncbi.nlm.nih.gov) and the Ensembl genome browser (ensembl.org). The human codon-optimized genes were ordered from Integrated DNA Technology as gBlocks Gene Fragments (IDT, idtdna.com). To generate constructs for transient transfection of HEK 293 FT cells, genes were subcloned into Emerald-N1 (addgene:53976) and piRFP670-N1 (addgene: 45457) vectors using In-Fusion cloning (takarabio.com), resulting in connexin fluorescent fusion proteins, specifically with the fluorescent proteins being adjoined to the connexin carboxy-terminus. Mutant constructs were
generated by employing overlapping primers within standard Phusion polymerase PCR reactions to facilitate site-directed mutagenesis.

The Gateway recombination (Invitrogen) system was used to generate all Connexin 36, Cx34.7, Cx35, wild type and mutant protein *C. elegans* expression plasmids. For PCR-based cloning and subcloning of components into the Gateway system, either Phusion or Q5 High-Fidelity DNA-polymerase (NEB) was used for amplification. All components were sequenced within the respective Gateway entry vector prior to combining components into expression plasmids via a four-component Gateway system [34].

The different connexins versions were introduced into pDONR221a using a similar PCR-based strategy from plasmid sources [35-37]. Cell-specific promoters were introduced using the pENTR 50 -TOPO vector (Invitrogen) after amplification from genomic DNA or precursor plasmids. Transgenic lines were created by microinjection into the distal gonad syncytium [38] and selected based on expression of one or more co-injection markers: Punc-122::GFP, Pelt-7::mCherry::NLS.

**Cell Culture**

HEK 293FT cells were purchased from Thermo Fisher Scientific (cat# R70007) and were maintained according to manufacturer instructions. Briefly, cultures were grown in 10 cm tissue culture treated dishes in high-glucose DMEM (Sigma Aldrich, D5796) supplemented with 6mM L-glutamine, 0.1 mM MEM non-essential amino acids and 1 mM MEM sodium pyruvate in a 5% CO₂, 37°C incubator. Cells were passaged via trypsinization every 2-3 days or until 60-80% confluency was reached.

**FETCH data analysis**

Complete FETCH methodology is outlined in Ransey et. al. 2021 [15]. Briefly, replica multi-well plates with HEK293FT cells were transfected with either of the two connexin proteins being evaluated for docking. Following a transfection incubation period, experimental counterparts were combined, replated, and further incubated for ~20-24hrs, allowing cells to make contacts and potentially generate
connexosomes. Co-plated samples were trypsinized, fixed in suspension and analyzed via flow cytometry. Flow cytometry data was processed with two selection gates prior to their fluorescence evaluation. First, putative HEK cells were identified by evaluating sample forward vs. side scatter area. Next, single cells were identified as the putative cells that maintained a linear correlation of forward scatter height to forward scatter area. Finally, the dual fluorescence (mEmerald vs. RFP670) profiles of each sample was generated, and the FETCH score was defined as the proportion of dual-labeled fluorescent cells that develop in a co-plated sample.

For screening analysis, five FETCH replicates were obtained for each condition (mutation). These scores were benchmarked against scores for Cx36 and Cx45 (FETCH=1.2±0.1%, N=54 replicates). To quantitatively determine whether a connexin pair docked, we determined FETCH scores for the dual fluorescence of cells under conditions where docking was not anticipated. These conditions included pairs of connexins previously established to not show docking: Cx36 and Cx45 (FETCH=0.7±0.0%, N=59 replicates), homotypic Cx23 (FETCH=0.9±0.4%, N=6 previously collected replicates[15]), Cx36 and Cx43 (FETCH=1.2±0.2%, N=10 previously collected replicates[15]), and under conditions for which cells were transfected with cytoplasmic fluorophores rather than tagged connexins (FETCH=4.4±0.6%, N=17 previously collected replicates[15]). These 92 FETCH scores were used as the ‘known-negative’ distribution. FETCH scores from each experimental condition were then compared against the established negative score distribution using a one-tailed t-test, with a Bonferroni correction for the total number of experimental conditions tested. These FETCH replicates were independent of the replicates utilized for our screening analysis. Stats are reported as mean±s.e.m, and only uncorrected P values are reported through the text.

Confocal Imaging Analysis of Gap Junction Partners
For imaging of putative gap junction partners, different populations of HEK 293FT cells were transfected with counterpart connexin proteins, incubated and combined as described for FETCH analysis. Combined samples of HEK 293FT cells were co-plated onto 10 ug/ml Fibronectin coated 35 mm, glass-bottom Mattek dishes (cat# P35GC-1.5-14-C). Cells were imaged at ~20 hrs post co-plating. Images were acquired on a Leica SP5 laser point scanning inverted confocal microscope using Argon/2, HeNe 594nm and HeNe633nm lasers, conventional fluorescence filters and a 63X, HCX PL APO W Corr CS, NA: 1.2, Water, DIC, WD: 0.22mm, objective. Images were taken with 1024 x 1024 pixel resolution at 200Hz frame rate.

For assessing Cx34.7M1::Cx35M1 expression in vivo in C. elegans, we imaged strain DCR8669 olaEx5214 [Pgcy-8::CX34.7(E214K, E223K)::GFP; Pttx-3::CX35(K221E)::mCherry; Punc-122::GFP]. L4 animals were mounted in 2% agarose in M9 buffer pads and anaesthetized with 10mM levamisol (Sigma). Confocal images were acquired with dual Hamamatsu ORCA-FUSIONBT SCMOS cameras on a Nikon Ti2-E Inverted Microscope using a confocal spinning disk CSU-W1 System, 488nm and 561nm laser lines and a CFI SR HP PLAN APO LAMBDA S 100xC SIL objective. Images were captured using the NIS-ELEMENTS software, with 2048px x 2048px, 16-bit depth, 300nm step size, 200ms of exposure time and enough sections to cover the whole worm depth.

Protein modeling pipeline

Our protein modeling pipeline is based on previously published methodology [39] and integrates five components: 1) homology model generation, 2) embedding of proteins in a lipid bilayer and aqueous solution, 3) protein mutagenesis, 4) system minimization, equilibration, and molecular dynamics simulation production run, and 5) residue-wise energy calculation.

Homology Modeling
We initially tested five homology modeling software suites: Robetta, SwissModel, Molecular Operating Environment (MOE; Chemical Computing Group ULC, Montreal, QC, Canada, H3A 2R7, 2021), I-Tasser, Phyre2 [40-48]. A quality assessment suite, — MOLProbity [49-51] revealed that Robetta models outperformed the rest, based on a set of standard metrics (Ramachandran plot outliers, clashscore, poor rotamers, bad bonds/angles, etc).

Since our aim was to model the extracellular loops responsible for connexin hemichannel docking, we picked all the resolved connexin structures that possessed a high degree of extracellular loop homology to our connexins of interest as the inputs for Robetta. The top homolog hits were generally the same for the three Cxs of interest: Connexin-26 Bound to Calcium (5er7.1) [25], Human Connexin-26 (Calcium-free) (5era) [25], Structure of connexin-46 intercellular gap junction channel at 3.4 angstrom resolution by cryoEM (6mhq) [52], Structure of connexin-50 intercellular gap junction channel at 3.4 angstrom resolution by cryoEM (6mhy) [52], Structure of the connexin-26 gap junction channel at 3.5 angstrom resolution (2zw3) [24]. Cx34.7 and Cx35 wild type sequences had the greatest homology degree with 6mhq, while Cx36 was most homologous to 5er7.1. We generated three wild type hemichannels for Cx34.7, Cx35, and Cx36.

System Assembly

Next, we assembled hemichannels into homotypic and heterotypic gap junctions, embedded them in two double bilayers, dissolved them in water, and added appropriate ion concentrations for the extracellular and two intracellular compartments. The primary software suite used for this modeling step was VMD [53, 54]. We also utilized CHARMM GUI to generate the naturalistic model of a region of a double bilayer [55-59]. Membrane components were then selected in appropriate proportions to resemble experimentally-derived data from a neuronal axonal membrane.
Specifically, since Robetta was unable to model the full gap junction, we merged hemichannels into full homotypic/heterotypic gap junctions in a semi-automated way. First, to make homotypic gap junctions, we loaded the two-homology models for a hemichannel. We then aligned them using the center of mass of the extracellular loops. A slight rotation along the z axis was implemented for several pairs to optimize their fit. To make heterotypic gap junctions, we created a homotypic gap junction for each hemichannel, aligned the extracellular loops for the two homotypic gap junctions, then removed an opposing hemichannel from each homotypic gap junction (leaving the two different hemichannels aligned). Next, using the constructed gap junction, we aligned two pre-made membrane bilayers with the center of mass assigned as each embedded hemichannel. We then removed membrane molecules that overlapped with the hemichannel or the hemichannel pore. Next, we dissolved the system in water, and removed water that overlapped with the lipid bilayer. Extracellular water was then separated to a new file, where Na+, K+, Cl−, and Ca2+ ions were added to yield concentrations mirroring the extracellular environment of mammalian neurons [60]. Finally, Na+, K+, Cl−, and Ca2+ ions were added to the intracellular space to mirror the intracellular environment of mammalian neurons, and the files containing the embedded Cx hemichannels and extracellular water were merged. Notably, these stages were automated yielding a streamlined progression from a protein-only hemichannel model to a fully embedded gap junction model ready for subsequent simulation and/or mutagenesis.

**Mutagenesis**

We developed a python command-line tool that utilizes VMD to generate mutation configuration files for subsequent MD simulation. Here, we simply specify the connexin hemichannels of interest and the position at which a specific mutation should be introduced.

**System Minimization, Equilibration, and MD Simulation Production Run**
Next, we minimized atomic energies, equilibrated the system, and ran the stable system in a production simulation run. Specifically, MD simulation was performed using NAMD [61] and is divided into five steps:

1) **Melt lipid tails while keeping remaining atoms fixed (simulate for 0.5 ns)**
2) **Minimize the system, then allow the bilayers and solutions to take natural conformation while keeping gap junction fixed (split in two stages to accommodate reduction in volume of relaxing system; simulate 0.5 ns total)**
3) **Release the gap junction and equilibrate the whole system (simulate 0.5 ns)**
4) **Run minimized and equilibrated system in a production run (simulate 0.5 ns)**

Though MD simulation (step 4) is highly reliant on the input file provided by the System Assembly process, these steps render the simulation much more robust to modeling imperfections. For example, the membrane model developed though System Assembly is very rigid and has the potential to behave like a solid rather than like a liquid. Thus, melting the lipid tails encourages the model to embody a liquid. Similarly, many atoms in the input file may have unnatural initial energies, such that if they are all released at once, they would start moving at high velocities and the simulation would fail. Therefore, bringing the system to a local energy minimum increases stability. Removing constraints on the water and lipids enables them to surround the gap junction in a naturalistic form. Finally, releasing the constraints on the gap junction enables it to take the most energetically stable conformation given the environment.

**Energy Calculation**

To predict the residues that play a prominent role in docking, we quantified all non-bonding interactions between the two connexin hemichannels at key residues on the extracellular loops. Output from the MD simulation was loaded into the VMD “NAMD Energy” plugin. We then calculated nonbonding energies
for all residues on each hemichannel that were within 12 angstroms of at least one residue on the other hemichannel. For each residue pair we then averaged energies across the 250 simulation frames.

**C. elegans strains and genetics**

Nematodes were cultivated at 20°C on nematode growth medium seeded with a lawn of Escherichia coli strain OP50 using standard methods [62]. One-day-old adult hermaphrodites were used for all experiments. The strains used in this study are listed in the Strain Table (supplemental table S2). All thermotaxis behavioral assays, molecular biology, transgenic lines, and calcium imaging were performed following the methods of Hawk et al. 2018 [23], with minor modifications as outlined in the following subsections.

**Thermotaxis Behavioral Assay**

Animals were reared at 20°C for all experiments with shifts to the 15°C training temperature 4 hours prior to testing. High-throughput behavioral analyses were performed as described [23, 63]. Briefly, synchronized 1-day-old adult populations were washed in M9 buffer [64], then transferred by pipette to the 20°C isotherm of the behavioral test plate (22cm x 22cm plates with a 18°C to 22°C thermal gradient). Each behavioral test plate was split in half along the temperature gradient using a thin and clear plastic divider. This allowed for wild type controls to be assayed on one half of the arena and connexin-expressing animals on the other half. Migration was monitored for 60 min at 2fps using a MightEx camera (BCE-B050-U). Trajectories were analyzed using an adaptation of the MagatAnalyzer software package as previously described [23, 63, 65].

**Calcium Imaging**

For imaging, worms were mounted on a thin pad of 5% agarose in M9 buffer between two standard microscope coverslips. Worms were immobilized with 7.5mM levamisol (Sigma). All solutions were pre-equilibrated to the holding temperature prior to sample preparation. Prepared coverslip assemblies
were placed at TH on the peltier surface of the thermoelectric control system. A thermal probe (SRTD-2, Omega) mounted onto the surface of the peltier was used for feedback control via a commercially available proportional-integral-derivative (PID) controller (FTC200, Accuthermo). Target temperatures were supplied to the PID controller by a custom computer interface written in LabView (National Instruments), which subsequently gated current flow from a 12V power supply to the peltier via an H-bridge amplifier (FTX700D, Accuthermo). Excess heat was removed from the peltier with a water-cooling system. Precise temperature control was initially confirmed with an independent T-type thermal probe (IT-24P, Physitemp) attached to a hand-held thermometer (HH66U, Omega) and routinely compared to incubator temperatures with an infrared temperature probe (Actron). After mounting worms and placing them on the thermal control stage, fluorescence time-lapse imaging was begun immediately prior to implementing the temperature protocol in LabView, and temperature readings were recorded continuously while imaging. Fluorescence time-lapse imaging (250ms exposure) was performed using a Leica DM5500 microscope with a 10X/0.40 HC PL APO air objective and a Hamamatsu ORCA-Flash4.0 LT camera. Image acquisition was performed using MicroManager [66]. Segmentation into regions of interest and downstream data processing was performed using FIJI [67], and custom scripts written in MATLAB (MathWorks), including alignment of fluorescence intensity values to the temperature stimulus, calcium response detection, and initial figure production. Temperature readings were assigned to image frames in MATLAB based on CPU timestamps on images and temperature readings. Heatmaps were displayed using the MATLAB ‘imagesc’ function, which scales data to the full color range available. The same scaling was applied to all comparable data within a figure. Unless otherwise specified, the scaling was based upon the highest intensity dataset to minimize saturation. For analyses of AFD calcium signals, we measured an ROI of one AFD soma per animal. For analyses of AIY, we quantified signal intensity in the synaptic Zone 2. AIY is a unipolar neuron with both inputs, including those from AFD, in zone 2 and outputs to postsynaptic partners. We also quantified a reference region outside of AIY.
Responses were scored as the initial rise of the AFD or AIY calcium signal as determined by a blind human observer. Consistent with previous reports [68], we observed clear and qualitatively similar responses to thermal stimuli in the synaptic Zone 2 region. The genetic background for the AFD and AIY calcium imaging lines used in this study (control and experimental) contained olals23, a caPKC-1 GOF mutation. This was done to match prior work [23] in which Connexin 36 was demonstrated to evoke AFD-locked responses in AIY compared to caPKC-1 animals without Connexin 36.

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Author contributions


Declaration of Interests

The authors declare no competing interests


Amino Acid Sequences

Wild-type  ...PGIFECRDYPCLKEVCRYVSRPTEK...
Mutants    ...PGIFECRDYPCLKKVEVCRYVSRPTEK...

In vitro mutagenesis

Homology Modeling
In silico mutagenesis

High-Scoring Single Connexon Models

Robetta
MOL Probity

CHARMM-GUI

VMD Toolbox
Custom Scripts

Membrane Bilayer Model

Embedding Proteins

NAMD Toolbox
Custom Scripts

Molecular Dynamic Simulation

Residue-Wise Energy Readings

Interaction Energy (kcal/mol)

-600
-500
-400
-300
-200
-100
0
100

Cx Residues

226 225 224 223 222 221 220 219 218 217 216 215 214 213
A

Trained at 15°C or PKC-1 GOF

AFD
Sensory Neuron

Cal²⁺ Signal

18°C 22°C

18°C 22°C

Endogenous Chemical Synapse

Cool Seeking Behavior

AIY
Interneuron

Ca²⁺ Signal

18°C 22°C

Warm Seeking Behavior

Exogenous Connexin Gap Junction

AFD::AIY Connexin

+ Trained at 15°C or PKC-1 GOF

AFD
Sensory Neuron

Ca²⁺ Signal

18°C 22°C

Endogenous Chemical Synapse

AIY
Interneuron

B

Sensory Neuron (AFD)

GCaMP A/F

1.5

1

0.5

0

120%

60%

40%

20%

0%

animals (n=60)

18°C

22°C

time (s)

10 20 30 40 50

Interneuron (AIY)

GCaMP A/F

0.5

0.2

0.1

0.05

0.01

60%

40%

20%

0%

animals (n=50)

18°C

22°C

time (s)

10 20 30 40 50

AIY [Cx36/Cx36]

GCaMP A/F

0.5

0.2

0.1

0.05

0.01

60%

40%

20%

0%

animals (n=50)

18°C

22°C

time (s)

10 20 30 40 50

AIY [Cx34.7M1/Cx35M1]

GCaMP A/F

0.5

0.2

0.1

0.05

0.01

60%

40%

20%

0%

animals (n=50)

18°C

22°C

time (s)

10 20 30 40 50

C

Response Penetrance (% Animals Responding)

*** n.s. ***

n.s.

n.s.

n.s.

10 20 30 40 50

18°C 22°C

Cx Configuration (AFD::AIY)

D

TTX Index

n.s.

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A
Electrical stimulation

Optogenetics (terminal stimulation)

Optogenetics (soma stimulation)

DREADDs

*External Actuator/Ligand

B
Long-term integration of
Circuits using Connexins (LinCx)

*No External Actuator/Ligand

Legend:
- Target cell/synapse
- Stimulated region
- Off-target signal propagation
- Desired natural inputs
- Undesired input uncoupling
- Off-target synaptic effect

Target Circuit

Soma
Axon
Synapse
### Supplementary Table S1, Detailed Author Contributions

<table>
<thead>
<tr>
<th>Author</th>
<th>Contributions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elizabeth Ramsey</td>
<td>Conceived FETCH methodology as approach for screening connexin mutant docking; coordinated and performed connexin DNA construct cloning; performed all FETCH experiments and subsequent microscopy; conceived strategy to employ computational models for rational design; coordinated design of computationally guided mutations; secured funding and resources; prepared figures; wrote original draft of the introduction, <em>in vitro</em> study results and associated materials and methods, and discussion; and revised the paper.</td>
</tr>
<tr>
<td>Kirill Chesnov</td>
<td>Conceived and developed the computational modeling pipeline to enable rational design; performed all computational modeling analysis, conceived methodology for quantification of FETCH results, analyzed FETCH results; discovered motif interaction schema that guided Cx34.7, Cx35, and Cx36 docking; prepared figures; wrote original draft of <em>in silico</em> study results and associated materials and methods, and discussion; and revised the paper.</td>
</tr>
<tr>
<td>Elias Wisdom</td>
<td>Conceived <em>C. elegans</em> neurophysiological and behavioral experiments with AAP; Generated connexin expression vectors for <em>C. elegans</em>; performed all <em>C. elegans</em> experiments and analyzed results; prepared figures; wrote original draft of <em>in vivo</em> study results and associated methods with A.A.P. and D.C.; and revised the paper.</td>
</tr>
<tr>
<td>Ryan Bowman</td>
<td>Researched gap junction structural/functional properties across animal species, systematically organized findings, selected candidate proteins as substrates for LinCx, and presented recommendations (to RH and KD) with EA, GET, and HS; Cloned connexin DNA constructs; Tested in vitro methods for evaluating gap function.</td>
</tr>
<tr>
<td>Tatiana Rodriguez</td>
<td>Analyzed FETCH results; Cloned connexin DNA constructs; prepared figures; revised the paper.</td>
</tr>
<tr>
<td>Elise Adamson</td>
<td>Researched gap junction structural/functional properties across animal species, systematically organized findings, selected candidate proteins as substrates for LinCx, and presented recommendations (to RH and KD) with GET, HS, and RB; Cloned connexin DNA constructs;</td>
</tr>
<tr>
<td>Gwenaëlle Thomas</td>
<td>Initiated research on gap junction structural/functional properties across animal species, including target residues to alter docking properties; systematically organized findings, selected candidate proteins as substrates for LinCx, and presented recommendations (to RH and KD) with EA, RB, and HS.</td>
</tr>
<tr>
<td>Agustin Almoril-Porras</td>
<td>Conceived <em>C. elegans</em> neurophysiological and behavioral experiments with EW; generated <em>C. elegans strains</em>; assisted in calcium imaging analyses; coordinated all <em>C. elegans</em> experiments; prepared figures; wrote original draft of <em>in vivo</em> study results and associated methods with E.W. and D.C.R.; and revised the paper.</td>
</tr>
<tr>
<td>Hannah Schwennesen</td>
<td>Researched gap junction structural/functional properties across animal species, systematically organized findings, selected candidate proteins as substrates for LinCx, and presented recommendations (to RH and KD) with EA, GET, and RB;</td>
</tr>
<tr>
<td>Daniel Colón-Ramos</td>
<td>Conceived <em>C. elegans</em> based strategy to assess designer gap junction functional properties; supervised all <em>C. elegans</em> experiments; wrote original draft of <em>in vivo</em> study results and associated methods with E.W. and A.A.P.; and revised the paper.</td>
</tr>
<tr>
<td>Rainbo Hultman</td>
<td>Coordinated research of gap junction structural/functional properties across animal species; co-selected fish Cx34.7/Cx35 pair for subsequent protein</td>
</tr>
<tr>
<td><strong>Engineering</strong></td>
<td><strong>Nenad Bursac</strong></td>
</tr>
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<tr>
<td></td>
<td><strong>Kafui Dzirasa</strong></td>
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<tr>
<td><strong>Attribution Process</strong></td>
<td>Each team member outlined their individual contributions across a standard set of domains (conceptualization and methodology, formal analysis, investigation, resources, writing -original draft, writing -review &amp; editing, visualization, Supervision, and Project Administration and Funding Acquisition), and subsequently had the opportunity to edit a summarized attribution description to their satisfaction. Contribution summaries were then shared across all team members. Each team member had the opportunity to raise concerns with regards to any other team member’s outlined contributions, and issues that remained unaddressed after additional revisions were subjected to a mediation process led by the lead principal investigator. The assigned authorships and these detailed author contribution descriptions reflect the outcome of this process.</td>
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<td>Genotype</td>
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<td><code>olaEx5215 [Pgcy-8::CX34.7(E214K, E223K)::GFP; Pttx-3::CX35(K221E)::mCh; Pelt-7::NLS::mCh (All 25ng/ul)]; oalaIs17 [Pmod-1::GCaMP6s (25ng/ul) PPttx-3::mCherry (25ng/ul) Punc-122::dsRed (40ng/ul)]; oalaIs23 [Pgcy-8(800)::caPKC-1B (30ng/ul), Pgcy-8(800)::tagRFP (10ng/ul), Punc-122::RFP (30ng/ul)]</code></td>
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</table>
Supplementary Figure S1: Confocal maximum intensity projections of *C. elegans* expressing the Cx34.7<sub>M1</sub>::Cx35<sub>M1</sub> pair, related to Fig. 5. GFP-tagged Cx34.7<sub>M1</sub> is expressed in the AFD neuron, with puncta along its axon (left). mCherry-tagged Cx35<sub>M1</sub> is expressed in the AIY neuron, with puncta along its neurite (middle). Composite image showing the colocalizing GFP/mCherry puncta (right). Scale bar is 10um. White arrows highlight puncta.
Supplementary Figure S2: Sequence alignment of several connexin proteins' predicted extracellular loop 2. Predicted EL2 regions of Cx36 (GJD2), Cx43 (GJA1), and Cx45 (GJC1) for humans and several species broadly utilized in neuroscience research, related to Fig. 6. Identical residues are shown in black, residues that are variable are highlighted with red text. Residues of Cx36 that align to the interaction motif of Cx34.7 and Cx35 are indicated by blue underline. Note zebrafish (Danio rerio) do not have a Cx36 gene, thus the closest homolog, Cx34.7, was used for comparison.