The AMIGO1 adhesion protein activates Kv2.1 voltage sensors

Condensed Title: AMIGO1 activates Kv2.1 voltage sensors

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1 Abstract

Kv2 voltage-gated potassium channels are modulated by AMIGO neuronal adhesion 2 3 proteins. Here, we identify steps in the conductance activation pathway of Kv2.1 channels that are modulated by AMIGO1 using voltage clamp recordings and spectroscopy of heterologously 4 expressed Kv2.1 and AMIGO1 in mammalian cell lines. AMIGO1 speeds early voltage sensor 5 6 movements and shifts the gating charge-voltage relationship to more negative voltages. The gating charge-voltage relationship indicates that AMIGO1 exerts a larger energetic effect on 7 voltage sensor movement than apparent from the midpoint of the conductance-voltage 8 9 relationship. When voltage sensors are detained at rest by voltage sensor toxins, AMIGO1 has a 10 greater impact on the conductance-voltage relationship. Fluorescence measurements from voltage sensor toxins bound to Kv2.1 indicate that with AMIGO1, the voltage sensors enter their 11 earliest resting conformation, yet this conformation is less stable upon voltage stimulation. We 12 conclude that AMIGO1 modulates the Kv2.1 conductance activation pathway by destabilizing 13

14 the earliest resting state of the voltage sensors.

15 Statement of Significance

Kv2 potassium channels activate a potassium conductance that shapes neuronal action potentials. The AMIGO family of adhesion proteins modulate activation of Kv2 conductances, yet, which activation steps are modified is unknown. This study finds that AMIGO1 destabilizes the earliest resting conformation of the Kv2.1 voltage sensors to promote activation of channel conductance.

21 Introduction

Voltage-gated potassium (Kv) channels of the Kv2 family open following membrane 22 23 depolarization and are critical regulators of neuronal electrical excitability. Mammals have two 24 Kv2 pore-forming α subunits, Kv2.1 and Kv2.2, which function as homo- or heterotetramers (1). The molecular architecture of Kv2 channels is similar to Kv1 channels for which atomic 25 26 resolution structures have been solved (2). Each α subunit monomer has six transmembrane helical segments, S1-S6. S1-S4 comprise a voltage sensor domain (VSD) while S5 and S6 27 together form one quarter of the central pore domain. In response to sufficiently positive 28 29 intracellular voltages, gating charges within the VSD translate from an intracellular resting position to a more extracellular activated conformation. This gating charge movement powers 30 31 the conformational changes of voltage sensor activation, which are coupled to subsequent pore opening and K⁺ conduction (3). Kv channels progress through a landscape of conformations 32 leading to opening, all of which define a pathway for the activation of the K^+ conductance. The 33 activation pathway of Kv2 channels is distinct from Kv1 channels, as Kv2.1 channels have a 34 pore opening step which is slower and more weakly voltage-dependent than the VSD movement 35 of Kv1 channels (3–5). The unique kinetics and voltage dependence of Kv2 currents are critical 36 37 to neuronal activity, as they regulate action potential duration and can either support or limit repetitive firing (6–10). 38 Kv2 channels are abundant in most mammalian central neurons (11). Genetic deletion of 39 40 Kv2.1 leads to seizure susceptibility and behavioral hyperexcitability in mice (12), and human

- 41 Kv2.1 mutations result in developmental epileptic encephalopathy (13–15), underscoring the
- 42 importance of these channels to brain function. Homeostatic Kv2.1 regulation maintains
- 43 neuronal excitability (16). Kv2.1 regulation by ischemia (17, 18), glutamate (19),

44 phosphorylation (20) and SUMOylation (21) and AMIGO auxiliary subunits (22, 23) all shift the 45 midpoint of the conductance–voltage relation (G–V). However, it is not known which steps in the 46 conductance activation pathway are modulated by any of these forms of regulation.

47 To identify steps in the Kv2.1 conduction activation pathway that are susceptible to modulation, we studied the impact of an AMIGO auxiliary subunit. The AMIGO (AMphoterin-48 Induced Gene and Open reading frame) family of proteins contains three paralogs in mammals: 49 AMIGO1, AMIGO2, and AMIGO3. AMIGO proteins are single-pass transmembrane proteins 50 51 with an extracellular immunoglobulin domain and several leucine-rich repeats (24). AMIGO1 has been proposed to play a role in schizophrenia biology (25). In vertebrate brain neurons, 52 AMIGO1 is important for cell adhesion (24), neuronal tract development (26), and circuit 53 formation (25–27). AMIGO1 colocalizes with Kv2 in neurons throughout the brains of multiple 54 mammalian species (22, 28). Coimmunoprecipitation of AMIGO1 and Kv2.1 (22, 23, 26) and 55 co-diffusion through cell membranes (22) indicate a robust interaction, consistent with an 56 AMIGO1-Kv2.1 complex being sufficiently stable for intensive biophysical studies. All three 57 AMIGO proteins activate the conductance of both Kv2 channel subtypes, shifting the 58 conductance-voltage relation by -5 to -15 mV (22, 23). While these shifts may seem small in 59 60 excitable cells that can have voltage swings of more than 100 mV, human mutations that shift the conductance-voltage relation of ion channel gating by similar magnitudes are correlated with 61 physiological consequences (13, 29–31). However, it is difficult to determine whether the 62 63 physiological consequences of mutations are caused by the gating shifts themselves. Here we investigate which steps in the Kv2.1 conductance activation pathway are 64 modulated by AMIGO1. In other voltage-gated ion channels, the G-V relation can be shifted to 65 more negative voltages by modulating pore opening (32–34), voltage sensor movement (35, 36), 66 or voltage sensor-pore coupling (37–39). Single-pass transmembrane auxiliary subunits modulate 67 other voltage-gated ion channel α subunits by a variety of mechanisms (32, 38, 40, 41). 68 However, AMIGO1 only shares a limited degree of homology with other single-pass 69 70 transmembrane auxiliary subunits (42), and divergent structural interactions have been observed among single-pass transmembrane auxiliary subunits (43, 44). As there is no consensus binding 71 pose or mechanism of interaction for auxiliary subunits, it is difficult to predict on which step in 72 the conductance activation pathway AMIGO1 acts. A recent study proposed that AMIGO 73 74 proteins shift Kv2.1 conductance by increasing voltage sensor-pore coupling and that AMIGOconferred changes to Kv2 voltage-sensing machinery are unlikely (23). Here we ask whether 75 76 AMIGO1 alters conformational changes associated with pore opening or with voltage sensor movement using a combination of electrophysiological and imaging approaches. We find that 77 78 AMIGO1 modulates voltage sensor movements which occur before pore opening. We find 79 AMIGO1 to have a greater impact on early voltage sensor movements than the conductance-80 voltage relation. We conclude that AMIGO1 destabilizes the earliest resting conformation in the

81 pathway of channel activation.

82 Methods

83 GxTX peptides

A conjugate of a cysteine-modified guangxitoxin-1E and the maleimide of fluorophore Alexa594 (GxTX Ser13Cys(Alexa594)) was used to selectively modulate Kv2.1 channel gating and to fluorescently identify surfaceexpressing Kv2.1 channels (45). Conjugates of propargylglycine (Pra)-modified GxTX and the fluorophore JP-N₃ (GxTX Ser13Pra(JP) and GxTX Lys27Pra(JP)) were used to monitor the chemical environment surrounding GxTX when localized to the channel (46). All modified GxTX-mutants were synthesized by solid phase peptide synthesis as described (46–48). Stock solutions were stored at -80 °C and thawed on ice on the day of experiment.

90

91 Cell culture and transfection

92 The HEK293 cell line subclone TS201A was a gift from Vladimir Yarov-Yarovoy and was maintained in 93 DMEM (Gibco Cat# 11995-065) with 10% Fetal Bovine Serum (HyClone, SH30071.03HI, LotAXM55317) and 1% 94 penicillin/streptomycin (Gibco, 15-140-122) in a humidified incubator at 37°C under 5% CO₂. Chinese Hamster 95 Ovary (CHO) cell lines were a Tetracycline-Regulated Expression (T-REx) variant (Invitrogen, Cat# R71807), and 96 cultured as described previously (47). The Kv2.1-CHO cell subclone (49) was stably transfected with pCDNA4/TO 97 encoding the rat Kv2.1 (rKv2.1) channel. Cell lines were negative for mycoplasma by biochemical test (Lonza, 98 LT07). 1 µg/ml minocycline (Enzo Life Sciences), prepared in 70% ethanol, was added to Kv2.1–CHO cells to 99 induce rKv2.1 channel expression for 1.5 hours to minimize series resistance-induced voltage errors in K⁺ current 100 recordings or for 48 hours to produce sufficient Kv2.1 density necessary for recording gating currents. 5 minutes 101 prior to transfection, cells were plated at 40% confluency in unsupplemented culture media free of antibiotics, 102 selection agents, and serum and allowed to settle at room temperature. For imaging studies (except concentration-103 response), cells were plated in 35 mm No. 1.5 glass-bottom dishes (MatTek, P35G-1.5-20-C). For concentration-104 response time-lapse imaging, cells were plated onto 22 x 22 mm No. 1.5H cover glass (Deckglaser). For 105 electrophysiological studies, cells were plated in 35 mm tissue culture treated polystyrene dishes (Fisher Scientific, 106 12-556-000). Transfections were achieved with Lipofectamine 2000 (Life Technologies, 11668-027). Each 107 transfection included 220 µL Opti-MEM (Life Technologies, 31985062), 1.1 µL Lipofectamine, and the following amount of plasmid DNA. HEK293 cell experiments: 0.1 μ g of mKv2.1 DNA and either 0.1 μ g of pEGFP, 108 109 mAMIGO1-pIRES2-GFP DNA, or hSCN1β-pIRES2-GFP. The pIRES2-GFP vector has an encoded internal 110 ribosome entry site which promotes continuous translation of two genes from a singular mRNA (50) so that GFP 111 fluorescence indicates the presence of AMIGO1 or SCN1 β mRNA. Kv2.1–CHO cell experiments: 1 μ g of either mAMIGO1-pEYFP-N1, pEGFP, rAMIGO2-pEYFP-N1, or rAMIGO3-pEYFP-N1. CHO cell experiments: 1 µg 112 113 of both pCAG-ChroME-mRuby2-ST and mAMIGO1-pEYFP-N1. Cells were incubated in the transfection cocktail 114 and 2 mL of unsupplemented media for 6-8 hours before being returned to regular growth media, and used for 115 experiments 40-48 hours after transfection. pEGFP, mAMIGO1-pEYFP-N1, and pCAG-ChroME-mRuby2-ST (51) 116 plasmids were gifts from James Trimmer. mAMIGO1-pEYFP-N1 uses a VPRARDPPVAT linker to tag the 117 internal C-terminus of wild-type mouse AMIGO1 (NM 001004293.2 or NM 146137.3) with eYFP. pCAG-118 ChroME-mRuby2-ST encodes an mRuby2-tagged channelrhodopsin with a Kv2.1 PRC trafficking sequence (51, 119 52). mKv2.1 (NM 008420) was purchased from OriGene (MG210968). hSCN18-pIRES2-GFP was a gift from Vladimir Yarov-Yarovoy. mAMIGO1 was subcloned into pIRES2-GFP between NheI and BamHI restriction sites. 120 121 rAMIGO2-pEYFP-N1 and rAMIGO3-pEYFP-N1 were generated by subcloning rat AMIGO2 (NM 182816.2) or 122 rat AMIGO3 (NM 178144.1) in place of mAMIGO1 in the mAMIGO1-pEYFP-N1 vector.

123

124 Whole-cell K⁺ ionic currents

125 Voltage clamp was achieved with an Axopatch 200B patch clamp amplifier (Axon Instruments) run by 126 Patchmaster (HEKA). Solutions: HEK293 internal (in mM) 160 KCl, 5 EGTA, 10 HEPES, 1 CaCl₂, 2 MgCl₂, and 127 10 glucose, adjusted to pH 7.3 with KOH, 345 mOsm. HEK293 external (in mM) 5 KCl, 160 NaCl, 10 HEPES, 2 128 CaCl₂, 2 MgCl₂, 10 glucose, pH 7.3 with NaOH, 345 mOsm, 5 µM tetrodotoxin added to recording solution: LJP 3.9 129 mV, E_K: -89.0 mV with HEK293 internal. Kv2.1–CHO internal (in mM) 70 KCl, 5 EGTA, 50 HEPES, 50 KF, and 130 35 KOH, adjusted to pH 7.4 with KOH, 310 mOsm. Kv2.1–CHO external (in mM) 3.5 KCl, 155 NaCl, 10 131 HEPES, 1.5 CaCl₂, 1 MgCl₂, adjusted to pH 7.4 with NaOH, 315 mOsm: LJP 8.5 mV, E_K: -97.4 mV with Kv2.1-CHO cell internal. High Mg²⁺ Kv2.1–CHO external (in mM) 3.5 KCl, 6.5 NaCl, 10 HEPES, 1.5 CaCl₂, 100 MgCl₂, 132 133 adjusted to pH 7.4 with NaOH, 289 mOsm: LJP 13.1 mV, EK: -97.4 mV with Kv2.1-CHO internal. Osmolality 134 measured with a vapor pressure osmometer (Wescor, 5520), 5% difference between batches were tolerated. Liquid

135 junction potential (LJP) values were tabulated using Patcher's Power Tools version 2.15 (Max-Planck), and

136 corrected post hoc, during analysis. Voltage protocols list command voltages, prior to LJP correction. Kv2.1-CHO

137 cells were harvested by scraping in Versene (Gibco, 15040066) or TrypLE (Gibco, 12563011), HEK293 cells were 138 dislodged by scraping. Cells were washed three times in a polypropylene tube in the external solution used in the

recording chamber bath by pelleting at 1,000 x g for 2 min, and rotated at room temperature (22-24 °C) until 139

140 use. Cells were then pipetted into a 50 μ L recording chamber (Warner Instruments, RC-24N) and allowed to settle

141 for 5 or more minutes. After adhering to the bottom of the glass recording chamber, cells were rinsed with external

142 solution using a gravity-driven perfusion system. Cells showing plasma membrane-associated YFP, or intracellular

143 GFP of intermediate intensity, were selected for patching. Thin-wall borosilicate glass recording pipettes (BF150-144 110-7.5HP, Sutter) were pulled with blunt tips, coated with silicone elastomer (Sylgard 184, Dow Corning), heat

145 cured, and tip fire-polished to resistances less than 4 M Ω . Series resistance of 3–9 M Ω was estimated from the

146 whole-cell parameters circuit. Series resistance compensation (of < 90%) was used as needed to constrain voltage

147 error to less than 10 mV, lag was 10 us. Cell capacitances were 4-15 pF. Capacitance and Ohmic leak were

148 subtracted using a P/5 protocol. Output was low-pass filtered at 10 kHz using the amplifier's built-in Bessel and

149 digitized at 100 kHz. Traces were filtered at 2 kHz for presentation. Intersweep interval was 2 s. HEK293 cells with 150 less than 65 pA/pF current at +85 mV were excluded to minimize impact of endogenous K^+ currents (53).). The

151 average current in the final 100 ms at holding potential prior to the voltage step was used to zero-subtract each

- 152 recording. Mean outward current ($I_{avg,step}$) was amplitude between 90-100 ms post depolarization. Mean tail current
- was the current amplitude between 0.2-1.2 ms into the 0 mV step. 100 μ L of 100 nM GxTX-594 was flowed over 153
- cells with membrane resistance greater than 1 G Ω , pulses to 0 mV gauged the time course of binding, and the G-V154
- protocol was run. Data with predicted voltage error, $V_{error} \ge 10 \text{ mV}$ was excluded from analysis. V_{error} was tabulated 155
- 156 using estimated series resistance post compensation $(R_{s,post})$

$$V_{error} = I_{avg,step} * R_{s,post}$$
(Eqn. A)

For
$$G-V$$
 profiles cell membrane voltage (V_{membrane}) was adjusted by V_{error} and LJP .

$$V_{membrane} = V_{command} - V_{error} - LJP$$
 (Eqn. B)

Tail currents were normalized by the mean current from 50 to 80 mV. Fitting was carried out using Igor Pro 158

159 software, version 7 or 8 (Wavemetrics, Lake Oswego, OR) that employs nonlinear least squares curve fitting via the

160 Levenberg-Marquardt algorithm. To represent the four independent and identical voltage sensors that must all

- 161 activate for channels to open, G-V relations were individually fit with a 4th power Boltzmann

$$f(V) = A\left(1 + e^{\frac{-(V-V_{1/2})zF}{RT}}\right)^{-x}$$
 (Eqn. C)

162 where f(V) is normalized conductance (G), A is maximum amplitude, x is the number of independent identical

transitions required to reach full conductance (for a 4th power function, x=4), $V_{1/2}$ is activation midpoint, z is the 163

164 valence in units of elementary charge (e_0) , F is the Faraday constant, R is the ideal gas constant, and T is absolute 165 temperature. The half-maximal voltage (V_{Mid}) for 4th power functions is

$$V_{\rm Mid} = V_{\rm i,1/2} + \frac{42.38}{z_{\rm i}}$$
 (Eqn. D)

166 Reconstructed Boltzmann curves use average z_i and $V_{1/2} \pm SD$. The minimum Gibbs free energy (ΔG_{AMIGO1}) that

AMIGO1 imparts to conductance, was tabulated as 167

$$\Delta G = -\mathbf{R} \times \mathbf{T} \times \ln(K_{eq})$$
(Eqn. E)

168 Here R = 0.00199 kcal/(K•mol) and T = 298K. K_{eq} , or the equilibrium constant of channel opening, was

approximated by $\frac{f_{\text{Kv2.1+AMIGO1}}(V_{i,\text{Mid},\text{Kv2.1}})}{1-f_{\text{Kv2.1+AMIGO1}}(V_{i,\text{Mid},\text{Kv2.1}})} \text{ where } f_{\text{Kv2.1+AMIGO1}}(V_{i,\text{Mid},\text{Kv2.1}}) \text{ is the reconstructed relative conductance}$ 169 170 of Kv2.1 + AMIGO1 at V_{i,Mid} of Kv2.1–control cells (Table 1).

- 171 Activation time constants (τ_{act}) and sigmoidicity values (σ) (54) were derived by fitting 10-90% current rise 172 with

$$I_K = A \left(1 - e^{-\frac{t}{\tau_{\text{act}}}} \right)^{\sigma}$$
(Eqn. F)

173 Where current at end of step, $I_{avg,step}$, was set to 100%. t = 0 was adjusted to 100 μ s after voltage step start to correct

- 174 for filter delay and cell charging. Deactivation time constants (τ_{deact}) were from fitting 1 to 100 ms of current decay
- 175 during 0 mV tail step with an exponential function

$$y = y_0 + Ae^{-\frac{t-t_0}{\tau_{deact}}}$$
(Eqn. G)

 $I_K = y_0 + Ae^{-\tau_{deact}}$ Reported τ_{deact} was the average after steps to +10 mV to +120 mV or +50 mV to +120 mV in GxTX-594. Kv2.1 176

177 deactivation kinetics became progressively slower after establishment of whole-cell mode, similar to Shaker 178 deactivation after patch excision (55). Due to the increased variability of deactivation kinetics expected from this

179 slowing phenomenon, deactivation kinetics were not analyzed further.

180

181 On-cell single channel K⁺ currents

182 Single channel recordings were made from on-cell patches, to avoid Kv2.1 current rundown that occurs 183 after patch excision (56). Methods same as whole-cell K⁺ ionic currents unless noted. While cells selected for recording had AMIGO1-YFP fluorescence apparent at the surface membrane, we cannot be certain each single 184 185 Kv2.1 channel interacted with AMIGO1. Solutions: Kv2.1-CHO single channel internal (in mM) 155 NaCl, 50 186 HEPES, 20 KOH, 2 CaCl₂, 2 MgCl₂, 0.1 EDTA, adjusted to pH 7.3 with HCl, 347 mOsm. Kv2.1–CHO single 187 channel external (in mM) 135 KCl, 50 HEPES, 20 KOH, 20 NaOH, 2 CaCl₂, 2 MgCl₂, 0.1 EDTA, adjusted to pH 7.3 with HCl, 346 mOsm: LJP -3.3 mV with Kv2.1-CHO single channel internal. Thick-wall borosilicate glass 188 189 (BF150-86-7.5HP; Sutter Instruments) was pulled, Sylgard-coated and fire–polished, to resistances $>10 \text{ M}\Omega$. 190 Analysis methods were same as prior (5) unless noted. To subtract capacitive transients, traces without openings 191 were averaged and subtracted from each trace with single-channel openings. Peaks in single channel amplitude 192 histograms were fit to half maximum with a Gaussian function to define single channel opening level for 193 idealization by half-amplitude threshold. Open dwell times were well described by a single exponential component 194 which was used to derived $\tau_{closing}$. Average open dwell times were also described as the geometric mean of all open 195 dwell times. Closed dwell times appeared to have multiple exponential components and were solely described as the 196 geometric mean of all closed dwell times.

197

198 Whole-cell gating current measurements

199 Methods same as whole-cell K⁺ ionic currents unless noted. Solutions: gating current internal (in mM) 90 200 NMDG, 1 NMDG-Cl, 50 HEPES, 5 EGTA, 50 NMDG-F, 0.01 CsCl, adjusted to pH 7.4 with methanesulfonic acid, 201 303 mOsm. Gating current external (in mM) 150 TEA-Cl, 41 HEPES, 1 MgCl₂ · 6 H₂O, 1.5 CaCl₂, adjusted to 202 pH to 7.3 with NMDG, 311 mOsm: LJP -3.3 mV with gating current internal. To avoid KCl contamination of the 203 recording solution from the pH electrode, pH was determined in small aliquots that were discarded. Cells were 204 resuspended in Kv2.1-CHO external and washed in the recording chamber with 10 mL gating current 205 external. Pipettes has resistances of 6-14 M Ω . Series resistances were 14-30 M Ω and compensated 50%. Cell capacitances were 6-10 pF. Verror was negligible (< 1 mV). P/5.9 leak pulses from -133 mV leak holding potential. 206 207 An early component ON gating charge movement was quantified by integrating ON gating currents in a 3.5 ms 208 window (O_{ON,fast}) following the end of fast capacitive artifacts created from the test voltage step (which usually 209 concluded 0.1 ms following the voltage step). The slow tail of the ON charge movement is difficult to accurately 210 integrate in these cells, making the cutoff point arbitrary. This 3.5 ms integration window resulted in a more positive 211 $Q_{\text{ON,fast}} - V$ midpoint than with a 10 ms window (5), and more positive midpoint than the G - V relation. Differences in 212 gating current solutions compared to prior studies may also contribute to the different midpoints reported (4, 5, 57). 213 Currents were baseline-subtracted from 4 to 5 ms into step. Q_{OFF} was determined by integration of OFF charge 214 movement in a 9.95 ms window after capacitive artifacts (usually 0.1 ms). Currents were baseline-subtracted from 215 10 to 20 ms into the step. Gating charge density fC/pF was normalized by cell capacitance. Q-V curves normalized 216 to average from 100-120 mV. Q-V relations were individually fit with a 1st power Boltzmann (Eqn. C., x=1). Time 217 constants (τ_{ON}) were determined from a double-exponential fit function

$$I_{\rm g,ON} = A\left(e^{\frac{-t}{\tau_{\rm ON}}}\right) + B - A_{\rm rise}\left(e^{\frac{-t}{\tau_{\rm rise}}}\right)$$
(Eqn. H)

- 218 τ_{rise} was not used in analyses. $I_{g,OFF}$ was not well fit by Eqn. H and τ_{OFF} was not analyzed. The voltage-dependence of
- 219 the forward voltage sensor activation (α) rate was determined by fitting the average $\pi_{ON}-V$ weighted by the standard
- 220 error

 $\tau_{\rm ON} = \frac{1}{\alpha_{\rm OM} e^{V z_{\alpha} F} /_{RT} + \beta_{\rm OM} e^{V z_{\beta} F} /_{RT}}$ (Eqn. I)

Reverse rates were not analyzed. Energy of AMIGO1 impact on the activation rate of all 4 voltage sensors 221 222 $(\Delta G^{\ddagger}_{AMIGOI})$ was

 $\Delta G = -4 \times R \times T \times \ln\left(\frac{k_{\text{Kv2.1 + AMIGO1}}}{k_{\text{Kv2.1}}}\right)$ where $k = \alpha_{0\text{mV}}$. Estimates of ΔG_{AMIGO1} from Q-V relations were with Eqn. E or (Eqn. J)

223

$$G = V_{1/2} \times Q \times F \tag{Eqn. K}$$

Here $F = 23.06 \text{ kcal/V} \cdot \text{mol} \cdot e_0$. Q was either z_g from fits or 12.5 e_0 as determined from a limiting slope analysis of the Kv2.1 open probability-voltage relation (3). $V_{1/2}$ was either $V_{g,\text{Mid}}$ or a median voltage ($V_{g,\text{Med}}$) as calculated from integration above and below $Q_{\text{OFF}}-V$ relations using a trapezoidal rule (58).

228 Fluorescence imaging

227

Images were obtained with an inverted confocal/airy disk imaging system with a diffraction grating separating 400-700 nm emission into 9.6 nm bins (Zeiss LSM 880, 410900-247-075) run by ZEN black v2.1. Laser lines were 3.2 mW 488 nm, 1.2 mW 514 nm, 0.36 mW 543 nm, 0.60 mW 594 nm. Images were acquired with a 1.4 NA 63x (Zeiss 420782-9900-799), 1.3 NA 40x (Zeiss 420462-9900-000), or 1.15 NA 63x objectives (Zeiss 421887-9970-000). Images were taken in either confocal or airy disk imaging mode. The imaging solution was Kv2.1–CHO external supplemented with 0.1% bovine serum albumin and 10 mM glucose. Temperature inside the microscope housing was 24-28 °C. Representative images had brightness and contrast adjusted linearly.

236 Concentration-effect imaging. Cells plated on coverslips were washed 3x with imaging solution then 237 mounted on an imaging chamber (Warner Instruments, RC-24E) with vacuum grease. 100 µL GxTX-594 dilutions 238 were applied for 10 minutes, then washed-out by flushing 10 mL at a flow rate of ~ 1 mL / 10 sec. 15 minutes after 239 wash-out, the next GxTX-594 concentration was added. Airy disk imaging, 1.4 NA 63x objective (Zeiss 420782-240 9900-799), 0.13 µm pixels, 0.85 µs dwell, 5 sec frame rate. YFP excitation 488 nm 2% power, emission 495-550 241 nm. GxTX-594 excitation 594 nm 2% power, emission 495-620 nm. Intensities extracted using FIJI (59). ROIs 242 drawn around groups of cells \pm YFP fluorescence. Dissociation constant (K_d) fit with fluorescence intensity at 0 nM 243 GxTX-594 set to 0 with

$$f(x) = A \frac{1}{(1 + K_{\rm d}/r)} + B$$
 (Eqn. L)

244 Voltage clamp fluorimetry was conducted as described (45). Briefly, 100 µL 100 nM GxTX-594 in imaging external was applied for 10 minutes then diluted with 1 mL Kv2.1-CHO external for imaging. Airy disk 245 246 imaging, 1.15 NA 63x objective (Zeiss 421887-9970-000), 0.11 µm pixels, 0.85 µs dwell, 2x averaging, 1 sec frame 247 rate. GxTX-594 excitation 594 nm 1% power, emission 605nm long-pass. Cells with obvious GxTX-594 labeling 248 were whole-cell voltage-clamped. Voltage clamp fluorimetry internal (in mM) 70 mM CsCl, 50 mM CsF, 35mM 249 NaCl, 1 mM EGTA, 10 mM HEPES, adjusted to pH 7.4 with CsOH, 310 mOsm: LJP -5.3 mV with Kv2.1-CHO 250 external. Pipettes from thin-wall glass were less than 3.0 MΩ. Cells were held at -100 mV for 30 images and stepped 251 to +35 mV until fluorescence change appeared complete. Intensity data was extracted using Zen Blue from ROIs 252 drawn around apparent surface membrane excluding pipette region. For presentation, fluorescence intensity traces 253 were normalized from minimum to maximum. Rate of GxTX-594 dissociation (k_{AF}) was fit with a monoexponential 254 function (Eqn. G), and K_{eq} for resting vs. activated voltage sensors was calculated as described (45). ΔG_{AMIGO1} from 255 with Eqn. J where $k = K_{eq}$.

256 Environment-sensitive fluorescence imaging with GxTX Ser13Pra(JP) and GxTX Lys27Pra(JP). Cells were 257 incubated in 100 μ L of GxTX(JP) solution for 5-10 minutes then washed with imaging solution. Spectral confocal 258 imaging, 1.4 NA 63x objective, 0.24 um pixels, 8.24 us dwell, 2x averaging. YFP excitation 514 nm. GxTX 259 Ser13Pra(JP) excitation 594 nm. GxTX Lys27Pra(JP) excitation 543 nm. Fluorescence counts extracted in Zen 260 Blue. JP emission spectra were fit with two-component split pseudo-Voigt functions (46) using the curve fitting 261 software Fityk 1.3.1 (https://fityk.nieto.pl/), which employed a Levenberg-Marquardt algorithm. Goodness of fit 262 was determined by root-mean-squared deviation (R^2) values, which are listed in Supplemental Table 2 along with 263 the parameters of each component function. To avoid YFP overlap, fittings for spectra from cells expressing 264 AMIGO1-YFP include emission data points from 613-700 nm for GxTX Ser13Pra(JP) and 582-700 nm for GxTX Lys27Pra(JP). Fittings for JP spectra from cells without AMIGO1-YFP included all data from 550-700 nm. 265

266

267 Experimental Design and Statistical Treatment

Independent replicates (*n*) are individual cells pooled over multiple transfections. The *n* from each transfection for each figure are listed in Supplemental Tables 3 and 4. In each figure panel, control and test cells

were plated side by side from the same suspensions, transfected side by side, and the data was acquired from control

and test cells in an interleaved fashion. Identity of transfected constructs was blinded during analysis. ANOVA

and est cents in an interfeaved fashion. Reputy of transferred constructs was officed during analysis. Areo VA analysis of transfection- or acquisition date-dependent variance of Boltzmann fit parameters and PCC/COV did not

reveal a dependence, and all *n* values were pooled. Statistical tests were conducted with Prism 9 (GraphPad

274 Software, San Diego, CA), details in figure legends.

275 Results

276 AMIGO1 shifts the midpoint for activation of Kv2.1 conductance

Voltage-clamp recordings from cotransfected HEK293 cells indicate that mouse 277 AMIGO1 shifts the G-V relation of mouse Kv2.1 by -5.7 ± 2.3 mV (SEM) (Supplemental Fig. 278 1). This shift was similar to the -6.1 mV \pm 1.6 mV shift reported of rat Kv2.1–GFP by human 279 AMIGO1–mRuby2 (23), and smaller than the -15.3 mV (no error listed) shift of mouse Kv2.1– 280 GFP by mouse AMIGO1 (22). This small effect of AMIGO1 was similar to the cell-to-cell 281 variability in our recordings. We suspected that endogenous voltage-activated conductances of 282 HEK293 cells (53, 60) and variability inherent to transient co-transfection could increase 283 variability. To minimize possible sources of cell-to-cell variability, further experiments were 284 with a Chinese Hamster Ovary K1 cell line with inducible rat Kv2.1 expression (Kv2.1–CHO) 285 transfected with a YFP-tagged mouse AMIGO1. Inducible Kv2.1 expression permits tighter 286 control of current density (49) and fluorescence tagging of AMIGO1 permits visualization of 287 protein expression and localization. Unlike HEK293 cells, CHO cells lack endogenous voltage-288 gated K^+ currents (61). 289

As expression systems can influence auxiliary protein interactions with ion channels (62– 66), we assessed Kv2.1–AMIGO1 association in these CHO cells. We evaluated two hallmarks of Kv2.1 and AMIGO1 association: Kv2.1 reorganization of AMIGO1, and AMIGO1 / Kv2.1 colocalization (22, 23, 28).

In HEK293 cells, heterologously expressed AMIGO1 localization is intracellular and 294 diffuse (23, 28). However, when co-expressed with Kv2.1, AMIGO1 reorganizes into puncta 295 296 with Kv2.1, similar to the expression patterns in central neurons (23, 28). To determine whether Kv2.1 reorganizes AMIGO1 in Kv2.1-CHO cells, the degree of AMIGO1-YFP reorganization 297 was quantified using the Coefficient of Variation (COV), which captures non-uniformity of YFP 298 299 localization (67). COV was quantified following the limited 1.5 h Kv2.1 induction period used in 300 whole-cell and single channel K⁺ current recordings and the prolonged 48 h induction period used for gating current recordings or imaging studies. COVs were compared against an 301 uninduced control (0 h induction) and against an engineered protein, ChroME-mRuby2, which 302 contains the Kv2.1 PRC trafficking sequence, but lacks the Kv2.1 voltage sensing and pore 303 forming domains (51, 52). COVs were evaluated from the glass-adhered, basal membrane where 304 evidence of reorganization is most notable (Fig. 1). Both $COV_{1.5h}$ and COV_{48h} were greater than 305 the COV_{0h} or COV_{ChroME-mRuby} control. This result is consistent with Kv2.1 and AMIGO1 306 association in CHO cells. 307

As an additional measure of whether Kv2.1 reorganizes AMIGO1 in Kv2.1–CHO cells, we assessed AMIGO1–YFP and Kv2.1 colocalization using the Pearson's correlation coefficient

310 (PCC) (68). Surface-expressing Kv2.1 on live cells was labeled with GxTX

311 Ser13Cys(Alexa594), a conjugate of a voltage sensor toxin guangxitoxin-1E derivative with a

fluorophore, abbreviated as GxTX–594 (45). As auxiliary subunits can impede binding of toxins

to voltage-gated ion channels (69), we tested whether AMIGO1 impacted GxTX-594 binding to

314 Kv2.1. Under conditions where AMIGO1 modulates most, if not all, Kv2.1 voltage sensor

movements (Fig. 6, 7), we found no evidence that AMIGO1 impedes GxTX-594 binding to

316 Kv2.1 (Supplemental Fig. 5). Colocalization between AMIGO1–YFP and GxTX–594 was

317 apparent as PCC_{48h}, measured from the glass-adhered basal membrane, was greater than the

negative control, PCC_{ChroME-mRuby2} (Fig. 2B). With a limited 1.5 h induction, GxTX–594 was

difficult to detect at the glass-adhered membrane, so we moved the confocal imaging plane

320 further from the cover glass to image Kv2.1 on apical cell surfaces where GxTX–594 labeling

was more apparent. On these apical surfaces, PCC_{1.5h} and PCC_{48h} were greater than PCC_{0h} (Fig. 321

2A), consistent with some colocalization of AMIGO1-YFP and Kv2.1. The weakly significant 322

increase of the PCC_{1.5h} compared to PCC_{0h} is consistent with some colocalization. 323

- Disproportionate expression can skew PCC values (70), and the limited GxTX-594 signal is 324
- expected to depress the PCC_{1.5h} value. Similarly, the lower PCC_{48h} values were associated with 325 either minimal or exceptionally bright AMIGO1-YFP signal. Overall, we see no sign of Kv2.1
- 326 channels lacking colocalized AMIGO1 in cells with high levels of AMIGO1 expression.
- 327
- Altogether, the reorganization and colocalization indicate that AMIGO1-YFP and Kv2.1 interact 328 in the CHO cells used for K⁺ current recordings and for gating current measurements. 329
- 330

331 AMIGO1 shifts the midpoint of activation of Kv2.1 conductance in CHO cells

To determine whether AMIGO1 affected the macroscopic K⁺ conductance in Kv2.1– 332 CHO cells, we conducted whole-cell voltage clamp recordings. Cells were transfected with GFP 333 (Kv2.1-control cells) or with AMIGO1-YFP (Kv2.1 + AMIGO1 cells) and identified for whole-334 cell voltage clamp based on the presence of cytoplasmic GFP fluorescence or plasma membrane-335 associated YFP fluorescence, respectively (Fig. 3A). Macroscopic ionic current recordings were 336 made in whole-cell voltage-clamp mode and K⁺ conductance was measured from tail currents 337 (Fig. 3B, C). In expectation of small AMIGO1 effects relative to cell-to-cell variation, recordings 338 from control cells and AMIGO1 cells were interleaved during each day of experiments and cell 339 identity was blinded during analysis. G-V relations were fit with a 4th power Boltzmann function 340 (Eqn. C) (Fig. 3D, E, F) and average midpoints of half-maximal conduction ($V_{i,Mid}$) and 341 steepness equivalents (z_i) were determined (Table 1). In Kv2.1–control cells, the average $V_{i,Mid}$ 342 was -1.8 mV (Fig. 3H), consistent with prior reports of $V_{i,Mid}$ ranging from -3 mV to +8 mV in 343 344 CHO cells (4, 23, 47, 71). Cell-to-cell variation in V_{i,Mid} remained notable between Kv2.1–CHO cells, with variation in $V_{i \text{ Mid}}$ on par with other reports (see *Discussion/Limitations*). The range of 345 $V_{i,Mid}$ values of Kv2.1 + AMIGO1 cells overlapped with Kv2.1–control cells (Fig. 3H), yet the 346 average $V_{i \text{ Mid}}$ was negatively shifted by -5.7 ± 2.2 mV (SEM), similar to $\Delta V_{i \text{ Mid}}$ from mouse 347 348 Kv2.1 in HEK293 cells (Table 1). No effect on z_i was observed. We also tested AMIGO2 and AMIGO3 on Kv2.1, and found they colocalize and induce $\Delta V_{i,Mid}$ shifts similar to those reported 349 from HEK293 cells by Maverick and colleagues (23) (Supplemental Fig. 3, 4), indicating that the 350 351 small G-V shifts by the AMIGO proteins are robust across different experimental preparations. To test if AMIGO1 also alters the rate of activation of Kv2.1 conductance, we analyzed 352 activation kinetics. The 10-90% of the rise of Kv2.1 currents following a voltage step (Fig. 3A, 353 354 B) was fit with the power of an exponential function (Eqn. F) for sigmoidicity (σ) which quantifies delay before current rise, and activation time constant (τ_{act}). σ was not significantly 355 altered by AMIGO1 (Fig. 3J, L, N), suggesting that the Kv2.1 activation pathway retains a 356 similar structure with AMIGO1 (5). At a subset of voltages less than +70 mV, AMIGO1 357 expression accelerated activation, decreasing τ_{act} (Fig. 3I, K, M), consistent with results of 358 Maverick and colleagues (23). Following the +10 to +120 mV activating steps, time constants of 359 360 tail current decay at 0 mV were similar to τ_{act} at 0 mV (Fig. 3O, Eqn. G). A prior study found no impact of AMIGO1 on Kv2.1 deactivation kinetics at -40 mV (23), and deactivation is not 361 studied further here. A model of Kv2.1 activation kinetics suggests that voltage sensor dynamics 362

influence τ_{act} below ~+70 mV, and that at more positive voltages a slow pore opening step limits 363

kinetics (5). This analysis suggests that AMIGO1 accelerates activation kinetics only in the 364

voltage range which is sensitive to voltage sensor dynamics. 365

Effects of AMIGO1 on pore opening conformational changes were not apparent in single 366 channel recordings 367

To more directly assess whether the pore opening step of the Kv2.1 activation pathway is 368 modulated by AMIGO1, we analyzed pore openings of single Kv2.1 channels during 1 s long 369 recordings to 0 mV (Fig. 4A, B). At 0 mV we expect >85% of all Kv2.1-control voltage sensors 370 or >95% of all Kv2.1–AMIGO1 voltage sensors (Fig. 6T) to activate in less than 2 ms (Fig. 6N), 371 such that the majority of single channel openings represent stochastic fluctuations between a 372 373 closed and open conformation of the pore. Neither the single channel current amplitude (Fig. 4C, D, E) nor the intra-sweep open probability (Fig. 4F) were significantly impacted by AMIGO1. 374 AMIGO1 did not significantly impact the single channel open or closed dwell times (Fig. 4G-L). 375 376 These results constrain any impact of AMIGO1 on Kv2.1 pore opening to be smaller than the variability in these single channel measurements. 377

378

A voltage sensor toxin enhances modulation of AMIGO1 on the Kv2.1 conductance 379

To test whether AMIGO1 modulation is dependent on voltage sensor dynamics, we 380 altered voltage sensor movement with a voltage sensor toxin. GxTX binds to the voltage sensing 381 domain of Kv2.1 (72), such that exit from the earliest resting conformation limits opening to 382 more positive voltages (5). If AMIGO1 modulates voltage sensors, then GxTX might be 383 expected to amplify the AMIGO1 effect. Alternately, if AMIGO1 acts directly on pore opening, 384 the AMIGO1 impact on the pore opening equilibrium should persist, regardless of voltage sensor 385 modulation. To distinguish between these possibilities, we measured AMIGO1 modulation in the 386 presence of the imaging probe GxTX-594, which modulates Kv2.1 by the same mechanism as 387 GxTX (45) and has a similar affinity for the resting conformation of Kv2.1 with or without 388 AMIGO1 (Supplemental Fig. 5). We applied 100 nM GxTX-594 to cells and activated the 389 Kv2.1 conductance. We note that the 100-ms activating pulses are much shorter than the >2390 second time constants of GxTX–594 dissociation at extreme positive voltages (45) and during 391 392 these short activating pulses we saw no evidence of GxTX-594 dissociation. The AMIGO1 $\Delta V_{i,Mid}$ of -22.1 ± 4.8 (SEM) with GxTX–594 was distinct from the AMIGO1 $\Delta V_{i,Mid}$ of -5.7 ± 393 2.2 mV (SEM) without GxTX-594 (p = 0.00018, unpaired, two-tailed t-test), indicating that 394 395 GxTX–594 amplifies the impact of AMIGO1 on Kv2.1 conductance. We did not observe a significant effect of AMIGO1 on τ_{act} or σ in GxTX–594 (Fig. 5J-N). We calculated the impact 396 of AMIGO1 on a pore opening equilibrium constant (K_{eq}) at the midpoint of the Kv2.1 G-V 397 relation and found a 3.7-fold bias towards a conducting conformation in 100 nM GxTX-594 398 versus a 1.4-fold bias under control conditions ($\Delta G_{AMIGO1} = -0.77$ versus -0.28 kcal/mol 399 400 respectively, Table 1). This result indicates that the impact AMIGO1 has on the Kv2.1 conductance is dependent on the dynamics of the activation path. Further, this result indicates 401 that AMIGO1 opposes the action of GxTX-594, which stabilizes the earliest resting 402 403 conformations of Kv2.1 voltage sensor. We also note that the more dramatic modulation by AMIGO1 with GxTX-594 verifies that most Kv2.1 channels are modulated by AMIGO1 in this 404 cell preparation in which only a small impact on $V_{i,Mid}$ was observed without GxTX-594 (Fig. 3). 405

406

AMIGO1 facilitates the activation of Kv2.1 voltage sensors 407

To determine if AMIGO1 affects voltage sensor movement, we measured gating currents 408 409 (I_g) , which correspond to movement of Kv2.1 voltage sensors across the transmembrane electric field. Kv2.1–CHO cells were patch clamped in whole–cell mode in the absence of K⁺ (Fig. 6A) 410

and given voltage steps to elicit gating currents (Fig. 6B, C). The resolvable ON gating currents ($I_{g,ON}$) represent an early component of gating charge movement, but not all of the total gating charge; the later charge movements, which include any charge associated with the pore opening, move too slowly for us to resolve accurately in ON measurements (4, 5). If AMIGO1 acts solely through the pore we would not expect to detect an impact on early components of ON gating currents which occur before pore opening.

At voltages above 50 mV, the charge density translocated over the first 3.5 ms, Q_{ON,fast}, 417 was not significantly different with AMIGO1 (Fig. 6D, E, F), indicating that AMIGO1 did not 418 alter the total charge translocated during early conformational transitions. However, between -10 419 mV and +50 mV, Kv2.1-control cells did not move as much gating charge as Kv2.1 + AMIGO1 420 421 cells, indicating a shift in gating current activation (Fig. 6F). The shift in voltage dependence was quantified by fitting $Q_{\text{ON,fast}}$ -V with a Boltzmann (Fig. 6G, H, I) yielding $\Delta V_{\text{g,Mid,ON,fast}}$ of -12.8 ± 422 423 3.5 mV (SEM) (Fig. 6K) and a $\Delta z_{g,ON,fast}$ of 0.215 ± 0.058 e_0 (SEM) (Fig. 6J) (Table 2). This result indicates that AMIGO1 modulates the early gating charge movement which occurs before 424

425 pore opening.

To determine whether AMIGO1 modulates the kinetics of early gating charge movement, 426 427 we extracted a time constant (τ_{ON}) from the decay phase of $I_{g,ON}$ that occurs before 10 ms (Fig. 6B top, C top) (Eqn. H) as in (5). In Kv2.1 + AMIGO1 cells, the $\tau_{ON}-V$ relation shifts to more 428 negative voltages compared to control (Fig. 6L, M, N). Above +30 mV, the mean τ_{ON} for Kv2.1 429 + AMIGO1 cells was faster than the mean τ_{ON} from Kv2.1–control cells (Fig. 6N). Fitting the 430 $\tau_{ON}-V$ with rate theory equations indicated AMIGO1 accelerates the forward rate of gating 431 charge movement by 1.7x at neutral voltage and decreases the voltage dependence of this rate by 432 13% (Fig. 6N). This result indicates that voltage sensors activate faster in the presence of 433 AMIGO1, consistent with destabilization of the earliest resting conformation of the voltage 434 sensors by AMIGO1. 435

To measure if AMIGO1 alters the total gating charge movement, we integrated OFF 436 gating currents (IgOFF) at -140 mV after 100 ms voltage steps (Fig. 6B bottom, C bottom, O, P, 437 Q). The density of Q_{OFF} elicited by voltage steps above -10 mV was not significantly different 438 between Kv2.1-control and Kv2.1 + AMIGO1 cells (Fig. 6Q), indicating that AMIGO1 did not 439 alter the density of channels expressed, nor the total gating charge per channel. However, 440 441 between -25 mV and -10 mV, Kv2.1-control cells did not move as much gating charge as Kv2.1 + AMIGO1 cells, indicating a shift in voltage dependence (Fig. 6Q). Boltzmann fits (Fig. 6R, S, 442 T), yielded $\Delta V_{g,Mid,OFF}$ of -10.8 ± 2.4 mV (SEM) (Fig. 6V) and a $\Delta z_{g,OFF}$ of 0.43 ± 0.20 e_0 (SEM) 443 (Fig. 6U) (Table 2), indicating that AMIGO1 shifts total gating charge movement to more 444 negative voltages. Overall, we find that AMIGO1 affects every aspect of gating current we have 445 analyzed to a greater degree than the K⁺ conductance. As both $Q_{ON,fast}-V$ and α_{0mV} measurements 446 report the gating charge movements out of the earliest resting conformation, these results 447 448 indicate that AMIGO1 destabilizes the earliest resting conformation relative to voltage sensor conformations later in the conduction activation pathway of Kv2.1. 449

450

451 AMIGO1 accelerates voltage-stimulated GxTX–594 dissociation

To further test the hypothesis that AMIGO1 specifically destabilizes the earliest resting conformation of Kv2.1 voltage sensors, we probed the stability of this conformation with GxTX– 594 fluorescence. The earliest resting conformation is stabilized by GxTX (5) and when occupancy of this conformation is decreased by voltage activation, the rate of GxTX–594

dissociation accelerates (45). Destabilization of the earliest resting conformation by AMIGO1 is

expected to increase the rate of GxTX-594 dissociation when voltage sensors are partially 457 activated. To test this prediction, we measured the rate of GxTX-594 dissociation at +30 mV, a 458 potential at which about 20% of Kv2.1 gating charge is activated with GxTX bound (5). The rate 459 of GxTX–594 dissociation from Kv2.1 (k_{AF}) accelerated from 0.073 ± 0.010 s⁻¹ (SEM) in control 460 cells to $0.115 \pm 0.015 \text{ s}^{-1}$ (SEM) in cells positive for AMIGO1–YFP fluorescence (Fig. 7). As we 461 see no evidence that AMIGO1 alters GxTX-594 affinity in cells at rest (Supplemental Fig. 5), 462 this 1.6-fold acceleration of $k_{\rm AF}$ is consistent with AMIGO1 destabilizing the earliest resting 463 conformation of voltage sensors. The thermodynamic model developed to interpret the $k_{\Delta F}$ of 464 GxTX-594 dissociation (45) estimates that AMIGO1 decreases the stability of the earliest 465 resting conformation of each voltage sensor by 1.9-fold or a ΔG_{AMIGO1} of -1.5 kcal/mol for Kv2.1 466 tetramers (Eqn. L). This result is consistent with AMIGO1 destabilizing the resting voltage 467 sensor conformation to speed up voltage sensor activation and shift conductance to lower 468 voltages. 469

470

471 An extracellular surface potential mechanism of AMIGO1 was not detected

To differentiate between mechanisms through which AMIGO1 could change voltage 472 sensor activation we probed whether the large AMIGO1 extracellular domain is directly 473 changing the electrostatic environment of Kv2.1's voltage sensors. Per surface charge theory, 474 local extracellular negative charges could attract positive gating charges to activate channels 475 (73). AMIGO1 possesses five extracellular glycosylation sites (74), each potentially decorated 476 with negatively-charged sugar moieties (28). AMIGO1 also has a conserved negatively charged 477 residue predicted to be near the extracellular side of the membrane (24, 74). Similar structural 478 characteristics are found in Nav β auxiliary subunits which, like AMIGO1, are glycosylated, 479 single transmembrane pass protein with an immunoglobulin-domain. Nav β1 has been proposed 480 to interact with Nav1.4 α subunit through surface charge effects (75–77). We tested if AMIGO1 481 likewise affects Kv2.1 activation through electrostatic surface charge interactions. 482

To measure the electrostatics of the environment immediately surrounding the Kv2.1 483 voltage sensor domain complex with and without AMIGO1, we employed far-red polarity-484 485 sensitive fluorescence (78). The polarity-sensitive fluorophore, JP, was localized to the Kv2.1 voltage sensor by conjugating GxTX to JP at either residue Ser13 or Lys27 (46). When GxTX 486 487 binds to the extracellular S3b region of the Kv2.1 channel, Ser13 and Lys27 occupy positions of distinct polarity (46). At resting membrane potentials, GxTX Ser13Pra(JP) has an emission 488 maximum of 644 nm, consistent with the homology-based prediction that Ser13 of GxTX 489 localizes in an aqueous environment branched away from S4. Conversely, GxTX Lys27Pra(JP) 490 491 has an emission maximum of 617 nm, consistent with the prediction that Lys27 sits in the polar region of the membrane adjacent to S4 (46). If AMIGO1 were to alter the electrostatic 492 environment of the resting conformation of the Kv2.1 voltage sensor domain, we would expect 493 either of these environmental point detectors, GxTX Ser13Pra(JP) or GxTX Lys27Pra(JP), to 494 495 exhibit an altered emission maximum.

Full emission spectra of JP fluorescence from Kv2.1–CHO cells transfected with AMIGO1–YFP and treated with GxTX Ser13Pra(JP) or GxTX Lys27Pra(JP) were fitted with 2component split pseudo-Voigt functions (Fig. 8C, F). Fitting shows emission peaks, 644 nm and 617 nm, respectively, are unchanged with or without AMIGO1–YFP, consistent with the local electrostatic environment surrounding the JP probes positioned on resting Kv2.1 voltage sensors not being altered by AMIGO1 expression. Previous work has shown that GxTX Lys27Pra(JP) emission peak wavelength is sensitive to conformational changes among early resting states of

voltage sensors (46). The absence of any AMIGO1-induced change in environment for either of 503 these GxTX sidechains suggests that AMIGO1 does not cause significant changes to the local 504 environment of the GxTX binding site on the S3b segment of Kv2.1, nor the GxTX position in 505 the membrane when bound to the channel. These results are consistent with destabilization of the 506 GxTX binding site by AMIGO1 being indirect, as the binding site itself appears to retain the 507 same conformation and local environment in the presence of AMIGO1. However, it remains 508 possible that AMIGO1 acts extracellularly to modulate Kv2.1 by a mechanism that these 509 GxTX(JP)-based sensors do not detect. 510

511 We also tested whether AMIGO1 acts by a surface charge mechanism with a classical 512 charge screening approach. Surface charge interactions can be revealed by increasing the

513 concentration of Mg^{2+} to screen, or minimize, the impact of fixed negative charges near the

voltage sensors (73, 79). If AMIGO1 alters surface potential, we would expect elevated Mg^{2+} to

shrink $\Delta V_{i,Mid}$. To determine whether surface charge screening suppresses the AMIGO1 effect,

voltage clamp experiments were conducted as in Fig. 3, except external recording solutions

517 contained 100 mM Mg²⁺ (Fig. 9A, B, C). Kv2.1 requires more positive voltage steps to activate

in high Mg^{2+} solutions (Table 1), consistent with sensitivity to surface charge screening (80). In

high Mg²⁺, AMIGO1 effected a $\Delta V_{i,Mid}$ of -7.4 ± 2.4 mV (SEM) (Fig. 10H) but did not change z_i

520 (Fig. 9G) (Table 1). When compared to low Mg^{2+} conditions by Ordinary 2-way ANOVA,

521 $\Delta V_{i,Mid}$ was not significantly different in normal versus 100 mM Mg²⁺ (interaction of p = 0.33).

522 Hence, Mg^{2+} altered Kv2.1 activation in a manner consistent with surface charge screening, yet

523 Mg^{2+} did not detectably abrogate the AMIGO1 effect. However, we cannot rule out the

possibility of a screened site that is inaccessible to Mg^{2+} . While neither extracellular fluorescence

525 measurements nor surface charge screening detected an extracellular impact of AMIGO1, we are

526 not able to rule out the possibility of an extracellular coupling to AMIGO1 that was not detected

527 by these methods.

528 Discussion

We asked whether AMIGO1 modulates Kv2.1 conductance by modulating conformational changes of pore opening or voltage sensor activation. We found that AMIGO1 destabilizes the resting, inward conformation of Kv2.1 voltage sensors, causing channels to activate at more negative voltages. This conclusion is supported by three major results:

533

534 1) AMIGO1 destabilizes the earliest resting conformation of Kv2.1 voltage sensors.

AMIGO1 expression accelerated conductance activation only at a subset of voltages where the activation kinetics are voltage sensitive (Fig. 3M). When voltage sensor movements were measured directly, gating current recordings revealed an acceleration of the forward rate constant (τ_{ON}) of gating charge activation in cells with AMIGO1. Between 0 and 120 mV, pore opening is

539 10-30x slower than $I_{g,ON}$ decay (Fig. 3M, 6N), too slow to influence the first few ms of $I_{g,ON}$.

540 When the change in the forward rate α_{0mV} (Fig. 6N), was used to estimate the amount of energy 541 AMICOL contributes to modulating K_{V2} 1 conformational bias we found that AMICOL

541 AMIGO1 contributes to modulating Kv2.1 conformational bias, we found that AMIGO1 542 imparted -1.3 kcal/mol per channel (Eqn. J) to $\Delta G^{\ddagger}_{AMIGO1}$. From this result we conclude that

AMIGO1 speeds the rate of conformational change between the earliest resting conformation

and its transition state in the activation path. Additionally, the AMIGO1 effect on GxTX–594

44 and its transition state in the activation path. Additionally, the AWIGOT effect on GxTX=594545 dissociation at +30 mV is consistent with AMIGO1 opposing the action of GxTX=594, which 546 stabilizes resting voltage sensors. All available evidence indicates that AMIGO1 destabilizes the

earliest resting conformation of Kv2.1 voltage sensors. We estimate that AMIGO1 destabilizes
the fully resting conformation of Kv2.1 channels by ~3 kcal/mol, relative to the fully active open
state, and that about half of this energy lowers the barrier for the initial exit of voltage sensors

- 550 from their resting conformation (Fig. 10A).
- 551

552 2) AMIGO1 has a greater impact on the voltage sensors than the pore opening.

Free energy estimates indicate more AMIGO1 perturbation of the Q-V than the midpoint of 553 the G–V. The ΔG for AMIGO1's impact on voltage sensor activation ranged from -1.9 kcal/mol 554 to -3.1 kcal/mol depending on the calculation method (Table 2). Yet, the ΔG_{AMIGO1} calculated at 555 the conductance midpoint was only -0.3 kcal/mol (Table 1). This lesser impact on pore opening 556 is consistent with a direct impact of AMIGO1 on voltage sensor movements which are coupled 557 to pore opening. Notably ΔG_{AMIGO1} calculated at the conductance midpoint widens to -0.8 558 kcal/mol when voltage sensor activation is limited with GxTX-594. When we looked at pore 559 opening directly, we saw no evidence suggesting a direct effect of AMIGO1. We saw no change 560 in the slope of the G-V relationship with AMIGO1 (Table 1), nor sigmoidicity (Fig. 3), nor 561 single channel measurements (Fig. 4). While these negative results do not eliminate the 562 possibility that AMIGO1 has a small direct effect on pore opening, these negative results 563 constrain the effect size of AMIGO1 on pore opening equilibria to be smaller than the error 564 associated with our measurements. 565

566

567 3) The AMIGO1 impact on conductance is malleable

In Kv2.1–CHO cells, AMIGO1 shifts the V_{Mid} of conductance by -5.7 ± 2.2 mV (SEM). With GxTX–594, the AMIGO1 *G*–*V* shift widens to -22.3 ± 4.8 (SEM) (Table 1). This remarkable result indicates that the AMIGO1 effect on conductance can change in magnitude. While we

have not completely excluded the possibility that AMIGO1 has a direct interaction with GxTX–

594, we think this unlikely, as we saw no sign of an AMIGO1-dependent environmental change 572 around GxTX–JP conjugates, and GxTX–594 had a similar affinity for resting Kv2.1. We think 573 it is more likely that AMIGO1 and GxTX-594 interact only allosterically, and favor the 574 575 explanation that GxTX makes the $V_{i,Mid}$ of conductance more sensitive to the early voltage sensor transition which AMIGO1 modulates. After its fast voltage sensor movement, Kv2.1 has a slow 576 conductance-activating step that makes the 4th power of the Q-V not predictive of the G-V (3–5, 577 57). GxTX stabilizes the earliest resting conformation of Kv2.1 voltage sensors such that 4th 578 power Boltzmann fits to the G-V are similar to the Q-V(5). This suggests the $V_{i,Mid}$ is more 579 responsive to AMIGO1 in GxTX-594 because the G-V becomes limited by early voltage sensor 580 movement. 581 To test the idea that AMIGO1 modulation of voltage sensors could result in different $\Delta V_{i,Mid}$ 582 of G-Vs, we performed calculations with a voltage sensor shift model composed of simple gating 583 equations. This voltage sensor shift model incorporates distinct $V_{1/2}$ values assigned to 584 independent voltage sensor $(V_{VSD,1/2})$ and pore $(V_{Pore,1/2})$ transitions, all of which must activate to 585 allow channel opening. Calculations incorporating a constant $\Delta V_{\text{VSD},1/2}$ shift with no change in 586 $V_{Pore,1/2}$ demonstrate that the $\Delta V_{i,Mid}$ of G-V can be malleable. In these calculations an AMIGO1 587 shift of $\Delta V_{\text{VSD},1/2}$ = -22.4 mV resulted in $\Delta V_{i,\text{Mid}}$ = -5.0 mV (Fig. 10B), similar to the empirical 588 measurement $\Delta V_{i,Mid}$ = -5.7 mV of Kv2.1 with AMIGO1 (Fig. 3). However, when $V_{VSD,1/2}$ was 589 modified to fit GxTX-594 data, this same AMIGO1 shift of $\Delta V_{\text{VSD},1/2}$ = -22.4 mV yielded a 590 591 larger shift G-V shift, $\Delta V_{i,Mid} = -21.8$ mV (Fig. 10B). While the gating model implied by these calculations is highly simplified and does not recapitulate all of our data, it does demonstrate a 592 mechanism by which a fixed modulation of voltage sensors could result in varying $\Delta V_{i,Mid}$ shifts. 593 As the voltage dependence of Kv2.1 activation is dynamically modulated by many forms of 594 cellular regulation and can vary dramatically (16–20, 81–86), the impact of AMIGO1 might also 595 fluctuate. A malleable impact of AMIGO1 in response to Kv2.1 regulation could perhaps explain 596 597 why a larger G-V shift was originally reported (22), than was observed here or elsewhere (23). The voltage sensor shift mechanism we propose does not require changes in pore opening, or 598 voltage sensor-pore coupling. Maverick and colleagues (23) suggested that the effects of 599 AMIGO proteins on Kv2.1 conductance could be described by increasing the coupling between 600 the voltage sensor and pore opening without a shift in the O-V curve (23), similar to a 601 mechanism by which leucine-rich-repeat-containing protein 26, LRRC26, modulates large-602 conductance Ca^{2+} -activated K⁺ channels (37). As the precise voltage sensor-pore coupling 603 604 mechanisms for Kv2.1 channels have yet to be defined, we cannot rule out the possibility that AMIGO1 also alters coupling. However, we see no reason that AMIGO1 must do anything other 605 than destabilize the earliest resting conformation of voltage sensors to modulate Kv2.1 606 607 conductance.

608

609 Limitations

610 More detailed investigation of the AMIGO1 impact on the Kv2.1 activation pathway was

611 limited by the relatively small magnitude of AMIGO1-dependent effects versus the cell-to-cell

variability, with $\Delta V_{i,Mid}$ as low as 5 mV, and standard deviations for $V_{i,Mid}$ of 4 to 9 mV (Table 1,

excluding GxTX–594). While we compensated for the limited power of the AMIGO1 effect by

614 increasing replicates, a decreased cell-to-cell would enable more precise biophysical

615 investigation. This degree of cell-to-cell variability does not appear to be unique to our

- laboratory. Midpoints reported for rat Kv2.1 activation in HEK293 cells span a 36 mV range,
- from -20.2 mV to 16.4 mV (22, 23, 67, 72, 87–95). When we calculated V_{Mid} standard deviation

values from the standard errors and n-values in these studies, standard deviations ranged from 1

to 17 mV, on par with our own. We suspect these notable V_{Mid} deviations result from the many

different types of regulation to which Kv2.1 channels are susceptible (20, 21). Techniques to

constrain the cell-to-cell variability in Kv2.1 function could allow more precise mechanistic
 studies of AMIGO1 modulation.

Our interpretations assume that the AMIGO1 effect is similar whether Kv2.1 is expressed at low density to measure K⁺ currents or at high density for gating current and imaging experiments. Auxiliary subunit interactions with pore α subunits can be influenced by many factors that can alter their assembly and functional impact on channel currents (96–101). However, if Kv2.1 channels in K⁺ current recording were modulated less by AMIGO1, we would expect a decrease in Boltzmann slope of the fit, a bimodal *G*–*V* relation, or increased cellto-cell variability with AMIGO1. We do not observe any of these with CHO cells. The similar

630 impact of AMIGO1 on Kv2.1 conductance in two cell lines (Table 1) and consistency in effect

magnitudes with an independent report (23), further suggest that AMIGO1 effect is saturating in

632 our K⁺ conductance measurements. Thus, while incomplete complex assembly and other factors

could in theory influence the magnitude of the AMIGO1 impact on Kv2.1 conductance, we do

not see evidence that would negate our biophysical assessment of the mechanism through which

635 AMIGO1 alters Kv2.1 conductance.

The most parsimonious explanation for the effect AMIGO1 has on the Kv2.1 conduction– voltage relation seems to be a direct interaction with Kv2.1 voltage sensors. However, it also seems possible that AMIGO1 proteins could change cellular regulation of which in turn modulates Kv2.1. Even if AMIGO1 acts by an indirect mechanism, our mechanistic conclusions remain valid, as they are not predicated on a direct protein–protein interaction between AMIGO1 and Kv2.1.

641 642

643 Potential physiological consequences of an AMIGO1 gating shift

The impact of AMIGO1 on Kv2.1 voltage sensors suggests that all voltage-dependent 644 Kv2 functions are modulated by AMIGO1. How might the AMIGO1 impact on voltage sensor 645 dynamics affect cellular physiology? As AMIGO1 is colocalized with seemingly all the Kv2 646 647 protein in mammalian brain neurons (22, 28, 102), our results suggest that AMIGO1 could cause Kv2 voltage-dependent functions to occur at more negative potentials in neurons. Consistent 648 with this suggestion, $I_{\rm K}$ currents from hippocampal pyramidal neurons isolated from AMIGO1 649 knockout mice are altered compared to wild type $I_{\rm K}$ currents (25). AMIGO1 knockout mice 650 display schizophrenia-related features (25) and AMIGO1 knockdown zebrafish have deformed 651 neural tracts (26). However, it is unclear whether these deficits are due to effects on channel 652 gating or other functions of AMIGO1, such as extracellular adhesion. In addition to electrical 653 signaling, Kv2 proteins have important nonconducting functions (28, 67, 103–106), which 654 AMIGO1 could potentially impact. Currently, we can only speculate about whether 655 physiological impacts of AMIGO1 are due to alteration of Kv2-mediated signaling. 656 Are the AMIGO1 effects on Kv2.1 conductance activation big enough to meaningfully 657 impact cellular electrophysiology? To begin to address this question, we estimated the impact 658 that AMIGO1 would have on neuronal action potentials. In mammalian neurons, Kv2 659 conductance can speed action potential repolarization (7, 107), dampen the fast 660 afterdepolarization phase (107), deepen trough voltage, and extend after-hyperpolarization (7) to 661 impact repetitive firing (7, 107-110). To estimate the impact AMIGO1 might have on the action 662 potentials, we superimposed the impact of AMIGO1 measured in Kv2.1-CHO cells onto the 663

Kv2 conductance in rat superior cervical ganglion (SCG) neurons, which Liu and Bean (7) found 664 to account for ~55% of outward current during an action potential. We roughly approximated an 665 SCG action potential as a 1.5 ms period at 0 mV, during which the parameters fit by Liu and 666 Bean predict 2.2% of the maximal Kv2 conductance will be activated. If the Kv2 parameters are 667 modified to mimic removal of AMIGO1, SCG neuron Kv2 conductance at the end of the mock 668 action potential decreases by 70% (Table 3). This large effect due to small changes in 669 conductance activation suggests that the AMIGO1 gating shift could have a profound impact on 670 electrical signaling. Furthermore, we think the AMIGO1 impact could be even greater. Liu and 671 Bean found that in SCG neurons, Kv2 activation lacks the slow pore-opening step we see in 672 Kv2.1-CHO cells, and SCG Kv2 kinetics were effectively modeled by a Hodgkin-Huxley n⁴ 673 674 model of activation (111). This suggests that only voltage sensor activation limits conductance activation in the SCG neurons. When the impact of AMIGO1 on Kv2.1-CHO voltage sensors is 675 applied to SCG neuron parameters, Kv2 conductance at the end of the mock action potential 676 decreases by 89% (Table 3). This analysis suggests that removal of the AMIGO1 effect in 677 neurons could be functionally equivalent to blocking the majority of the Kv2 current during an 678 action potential, which would in turn be expected to have impacts on repetitive firing (7, 107– 679

- 110). However, we stress that any predicted impact of AMIGO1 on action potentials is merely
- 681 speculation.

682 Conclusions

- To shift the activation midpoint of Kv2.1 conductance to lower voltages, AMIGO1
- destabilizes the earliest resting conformations of Kv2.1 voltage sensors relative to more activated
- 685 conformations. While we cannot rule out a direct influence on pore dynamics, we saw no
- 686 indication of such. We propose that AMIGO1 shifts the voltage-dependence of Kv2.1
- 687 conduction to more negative voltages by modulating early voltage sensor movements.
- 688 We also propose that because AMIGO1 acts on early voltage sensor movements, modulation of
- Kv2 gating can alter the impact of AMIGO1 on K^+ conductance.

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Competing Interests

We declare no competing interests.

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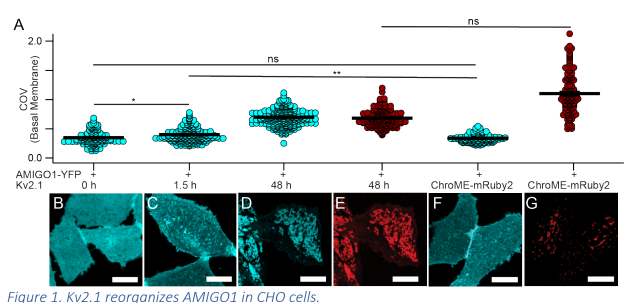
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(A) Coefficient of variation of fluorescence from AMIGO1–YFP (blue circles), GxTX-594 (red circles), or ChroME-mRuby2 (red circles). Bars are mean ± SEM. COV measurements were calculated from confocal images acquired from the glass–adhered basal membrane of the cell (exemplar confocal images in B-G). All cells were transfected with AMIGO1–YFP 48 h prior to imaging. COV from individual cells (*n*) were pooled from 4 separate transfections for each experimental condition. AMIGO1–YFP fluorescence from cells (**B**) not induced for Kv2.1 expression (COV_{0h} = 0.3492 ± 0.0098, *n* = 134), (**C**) induced 1.5 h (COV_{1.5h} = 0.4013 ± 0.0077, *n* = 217), (**D**) induced 48 h (COV_{48h} = 0.6984 ± 0.0083, *n* = 277). (**E**) GxTX–594 labeling from panel D (COV_{48h(GxTX-594)} = 0.6822 ± 0.010, *n* = 197). (**F**) AMIGO1–YFP fluorescence from CHO cells which lack Kv2.1 co-transfected with ChroME-mRuby2 (COV_{lack} = 0.3377 ± 0.0059, *n* = 125). (**G**) ChroME-mRuby2 fluorescence from panel F (COV_(ChroME-mRuby2) = 1.102 ± 0.030, *n* = 128). Scale bars 10 μ m. (**Statistics**) Outliers removed using ROUT, Q = 1%. Ordinary one-way ANOVA with multiple comparisons. P-values: COV_{0h}COV_{1.5h}: p = 0.0467; COV_{0h}COV_{lack}: p = 0.0981; COV_{48h(GxTX-594}) COV_{(ChroME-mRuby2}): p =0.9010. All other p-values ≤ 0.0001.

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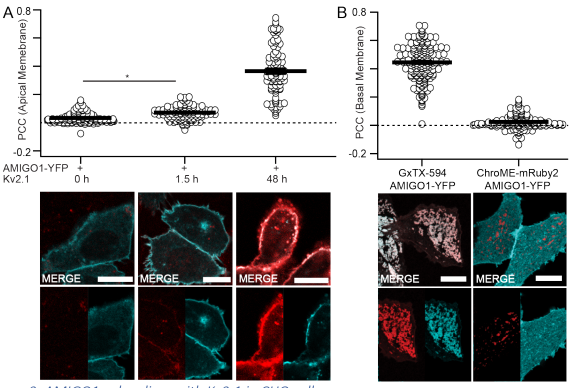


Figure 2. AMIGO1 colocalizes with Kv2.1 in CHO cells.

(A) Costes thresholded, Pearson's colocalization between AMIGO1–YFP and GxTX–594 at cell membrane following, from left to right, 0, 1.5, or 48 h of Kv2.1 induction (exemplar confocal images in B-J below). Mean \pm SEM (one-tailed ≥ 0 t-test): PCC_{0h} = 0.0321 \pm 0.0033, (p < 0.0001), n = 101; PCC_{1.5h} = 0.0718 \pm 0.0042, (p < 0.0001), n = 118; and PCC_{48h} = 0.365 \pm 0.017, (p < 0.0001), n = 101. (B) Costes thresholded, Pearson's colocalization between (left to right) AMIGO1–YFP/GxTX–594 and AMIGO1–YFP/ChroME-mRuby2 at the glass-adhered basal membrane of the cell. Exemplar images are the same as in Fig. 1 D-G. From left to right: PCC_{GxTX-594} = 0.4449 \pm 0.0090, (p < 0.0001), n = 195; PCC_{ChroME-mRuby2} = 0.0242 \pm 0.0045, (p < 0.0001), n = 129. Image panels with merge overlays (white) of GxTX–594 (red) and AMIGO1–YFP (cyan) correspond to conditions above. All scale bars are 10 μ m. (Statistics) Outliers were removed using ROUT, Q = 1%. Ordinary one-way ANOVA with multiple comparisons. P-values: PCC_{0h}PCC_{1.5h}: p = 0.346; PCC_{1.5h}PCC_{ChroME-mRuby2}: p = 0.0025; PCC_{0h5h}PCC_{ChroME-mRuby2}: p = 0.9777. All other p-values were \leq 0.0001.

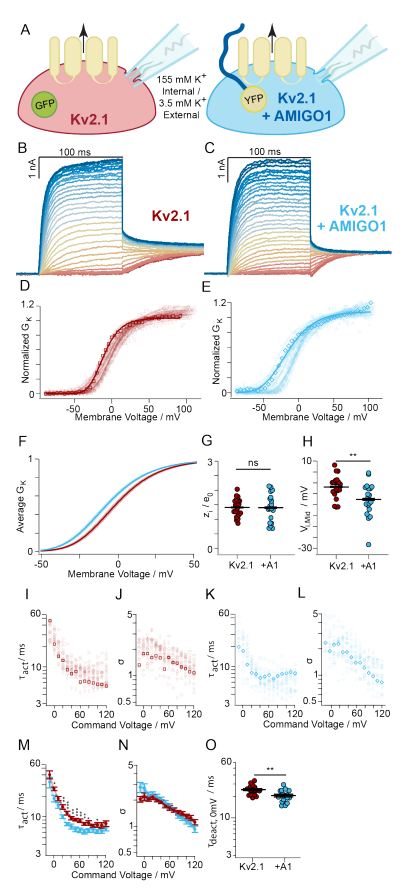


Figure 3. AMIGO1 shifts the midpoint and speeds activation of the Kv2.1 conductance in CHO cells.

(A) Experimental set up: Whole-cell K⁺ currents (arrow) from Kv2.1-CHO transfected with GFP (red) or AMIGO1-YFP (blue). (B, C) Representative Kv2.1control (6.0 pF) or Kv2.1 + AMIGO1 (14.5 pF) cell. 100 ms voltage steps ranging from -80 mV (dark red trace) to +120 mV (dark blue trace) in 5 mV increments and then to 0 mV for tail currents. Holding potential was -100 mV. Data points from representative cells are bolded in analysis panels. (D, E) Normalized tail G-V relationships for Kv2.1-control or Kv2.1 + AMIGO1 cells. Symbols correspond to individual cells. Lines are 4th order Boltzmann fits (Eqn. C). (F) Reconstructed Boltzmann fits from average $V_{i,Mid}$ and z_i (Table 1). Shading $V_{i,Mid} \pm SEM.$ (G) Steepness and (H) midpoint of fits. (I, K) τ_{act} and (J, L) σ from fits of Eqn. F to activation (M) Mean τ_{act} and (N) σ . (O) τ_{deact} fits of Eqn. G to 0 mV tails: Kv2.1–control 24.9 ± 3.6 ms, Kv2.1+AMIGO1 20.6 ± 3.8 ms. Unpaired t-test p > 0.5 between 0 mV τ_{act} and τ_{deact} for Kv2.1-control and Kv2.1 + AMIGO1. All other statistics in Table 1. ***: p = ≤ 0.001 , **: $p = \leq 0.01$, *: $p = \leq 0.05$, ns: not significant. Bars are mean \pm SEM.

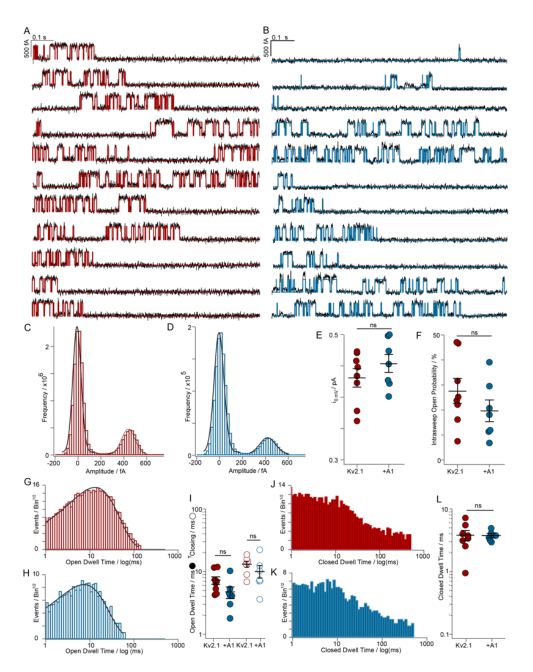


Figure 4. Effects of AMIGO1 on pore opening conformational changes were not apparent in single channel recordings.

(A) Representative single channel currents at 0 mV from Kv2.1–control and (B) Kv2.1 + AMIGO1. Red or blue lines are idealizations. (C,D) Amplitude histograms at 0 mV from the patches in A,B fit with Gaussians. (E) Mean single channel current amplitude: Kv2.1–control 0.43 ± 0.01 pA, Kv2.1 + AMIGO1 0.45 ± 0.02 pA. (F) Open probability from amplitude histograms: Kv2.1–control 28 ± 4.9%, Kv2.1 + AMIGO1 20 ± 4.2%. (G) Open dwell-time distributions and single exponential fits for a Kv2.1–control or (H) Kv2.1 + AMIGO1 patch. (I) Open dwell times from mean (filled circles) or exponential fit (hollow circles). Kv2.1–control: 13.0 ± 1.3 μ s. Kv2.1 + AMIGO1: 9.98 ± 2.3 μ s. (J) Closed dwell-time distributions and single exponential fit for a Kv2.1–control: 3.80 ± 0.67 μ s. Kv2.1 + AMIGO1: 3.73 ± 0.250 μ s. ns = two-tailed t-test p-value > 0.05. Means ± SEM.

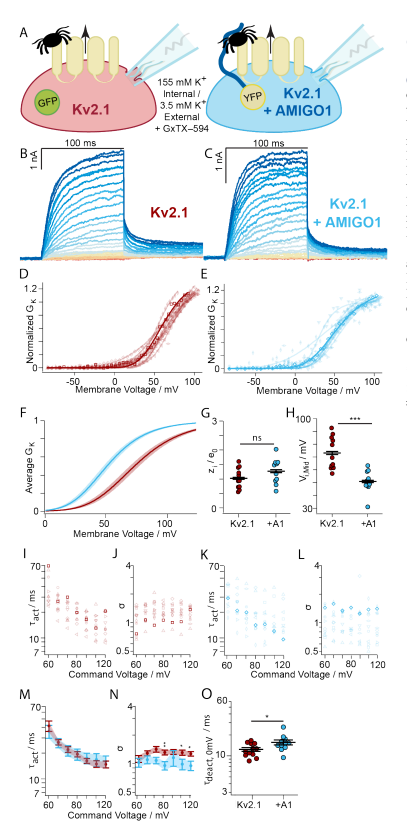
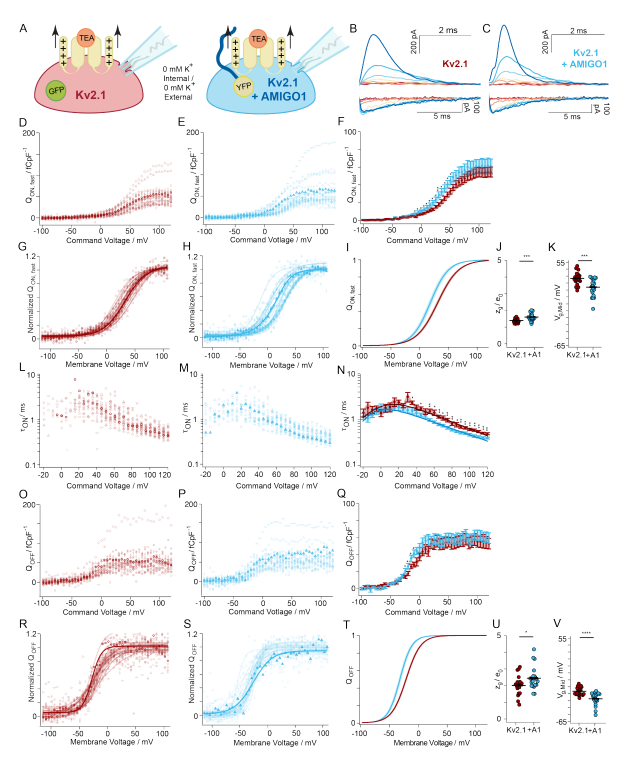


Figure 5. The voltage sensor toxin GxTX–594 enhances AMIGO1

modulation of Kv2.1 conductance. (A) Experimental set up: Whole-cell K⁺ currents (arrow) from Kv2.1-CHO transfected with GFP (red) or AMIGO1-YFP (blue). Cells were treated with 100 nM GxTX-594 (tarantulas). (B, C) Representative Kv2.1-control (6.0 pF) or Kv2.1 + AMIGO1 (14.5 pF) cell. Same voltage protocol and representations as Fig. 3. (**D**, **E**) Normalized G-Vrelationships (F) Reconstructed 4th order Boltzmann fits from $V_{i,Mid}$ and z_i in Table 1. Shading $V_{i,Mid} \pm SEM$. (G) Steepness and (H) midpoint of fits. (I, K) τ_{act} and (J, L) σ from fits of Eqn. F to activation (M) Mean τ_{act} and (N) σ . (O) τ_{deact} fits of Eqn. G to 0 mV tails: Kv2.1 with GxTX-594 = 12.4 ± 2.7 ms. Kv2.1+AMIGO1 with GxTX-594 = 15.7

 \pm 4.2 ms. All other statistics in Table 1. ***: $p = \le 0.001$, **: $p = \le 0.01$, *: $p = \le 0.05$, ns: not significant. Bars are mean \pm SEM. bioRxiv preprint doi: https://doi.org/10.1101/2021.06.20.448455; this version posted October 11, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.





(A) Experimental set up: Gating currents (arrows) from Kv2.1–CHO transfected with GFP (red) or AMIGO1–YFP (blue). K⁺ currents were eliminated removal of K⁺ ions and the external tetraethylammonium, a Kv2 pore-blocker (orange). (**B**, **C**) Top/Bottom: Representative $I_{g,ON}/I_{g,OFF}$ from Kv2.1–control (11.9 pF) or Kv2.1 + AMIGO1 (8.2 pF). Cells were given 100 ms voltage steps ranging from -100 mV (dark red trace) to +120 mV to record $I_{g,OFF}$. The holding potential was -100 mV. Voltage pulses to -100, -50, -25, +0, +25, +50, and +100 mV are presented. Data points from representative cells are bolded in analysis panels.

(**D**, **E**) $Q_{\text{ON,fast}}/pF-V$ relation from individual cells. $Q_{\text{ON,fast}}/pF$ is gating charge integrated over the first 3.5 ms normalized to cell capacitance. (**F**) Mean $Q_{\text{ON,fast}}/pF$ (**G**, **H**) $Q_{\text{ON,fast}}-V$ relations normalized to maximum $Q_{\text{ON,fast}}$ from +50 to +100 mV voltage steps. Solid lines represent Boltzmann fit (Eqn. C). (**I**) Reconstructed Boltzmann fits from average $V_{g,\text{Mid,ON,fast}}$ and $z_{g,\text{ON,fast}}$ (Table 2). Shading $V_{g,\text{Mid,ON,fast}} \pm$ SEM. (**J**) Steepness and (**K**) midpoint of Boltzmann fits. (**L**, **M**) τ_{ON} from individual cells fit with Eqn. I. (**N**) Average $\tau_{\text{ON}}-V$. Solid lines are Eqn. I fit. Fit values \pm SD for Kv2.1–control cells: $\alpha_{0\text{mV}} = 254 \pm 26 \text{ s}^{-1}$, $z_{\alpha} = 0.468 \pm 0.026 e_0$, $\beta_{0\text{mV}} = 261 \pm 50 \text{ s}^{-1}$, $z_{\beta} = -1.31 \pm 0.37 e_0$; for Kv2.1 + AMIGO1 cells: $\alpha_{0\text{mV}} = 443 \pm 26 \text{ ms}^{-1}$, $z_{\alpha} = 0.405 \pm 0.019 e_0$, $\beta_{0\text{mV}} = 157 \pm 52 \text{ ms}^{-1}$, $z_{\beta} = -2.00 \pm 0.55 e_0$. (**O**, **P**) Q_{OFF}/pF relation from individual cells normalized to cell capacitance. (**Q**) $Q_{\text{OFF}}/pF-V$ relation. (**R**, **S**) $Q_{\text{OFF}}-V$ relations normalized to maximum Q_{OFF} from +50 to +100 mV voltage steps. Solid lines are Boltzmann fits (Eqn. C). (**T**) Reconstructed Boltzmann fits using the average $V_{g,\text{Mid,OFF}}$ and $z_{g,\text{OFF}}$ (Table 2). Shading $V_{g,\text{Mid,OFF}} \pm$ SEM (**U**) Steepness and (**V**) midpoint of Boltzmann fits. Mean \pm SEM. Statistics in Table 2. ****: $p = \leq 0.0001$, ***: $p = \leq 0.001$, *: $p = \leq 0.01$, *: $p = \leq 0.05$, ns: not significant. Bars are mean \pm SEM.

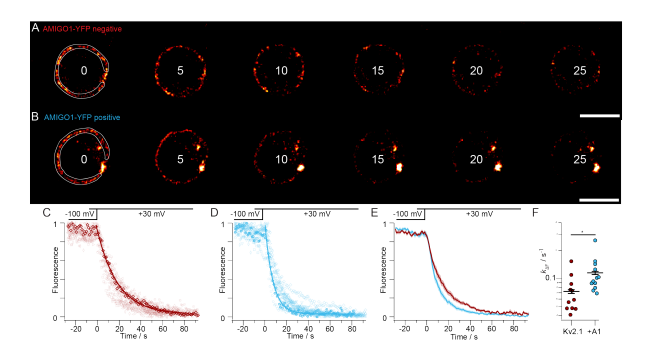


Figure 7. AMIGO1 accelerates voltage-stimulated GxTX–594 dissociation.

(A, B) Fluorescence from the solution-exposed membrane of voltage-clamped Kv2.1–CHO cells \pm AMIGO1–YFP. Kv2.1 expression was achieved through a 48-hour induction period. Cells were held at -100 mV for 30 seconds before being stimulated to +30 mV (time = 0 s) to trigger GxTX–594 dissociation. The time point in seconds of each image is listed. Region of interest for analysis is shown by the white line in left panel, which excludes the point contact with pipette and intracellular regions which have voltage-insensitive fluorescence. 10 μ m scale bar. (C, D) Normalized fluorescence intensity decay plots for Kv2.1–CHO cells without (red) and with (blue) AMIGO1–YFP fluorescence. The bolded traces correspond to exemplar cells in (A) and (B). Solid line is monoexponential fit (Eqn. G). (E) Averaged fluorescence intensity decay for AMIGO1–YFP negative (red), and AMIGO1–YFP positive (blue) cells. SEM is shaded. (F) Rates of fluorescence change ($k_{\Delta F}$) were calculated as 1/ τ from Eqn. G fits. *: p = 0.03 unpaired, two-tailed, t-test

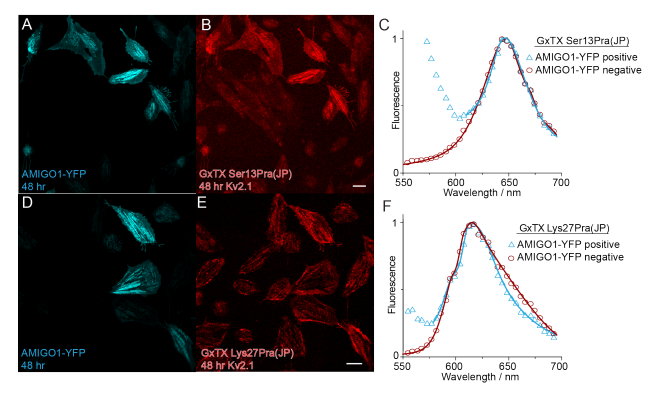


Figure 8. AMIGO1 does not alter the Kv2.1–GxTX interface on resting voltage sensors.

Kv2.1–CHO cells transfected with AMIGO1-YFP were treated with GxTX Ser13Pra(JP) or GxTX Lys27Pra(JP) (**A**, **D**) Confocal image of AMIGO1–YFP fluorescence (blue) and (**B**, **E**) JP fluorescence. (**C**, **F**) Fitted emission spectra of cells positive (blue) and negative (red) for AMIGO1–YFP fluorescence. Data points for all spectra are the mean of normalized emission from AMIGO1–YFP positive cells and AMIGO1–negative cells. Spectra were fit with two–component split pseudo–Voigt functions with shape parameters and root–mean–squared values found in Supplemental Table 1.

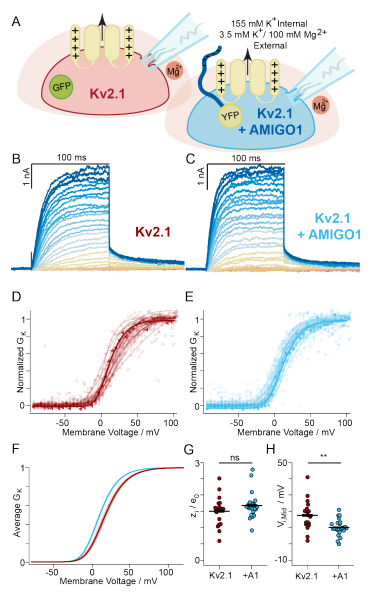


Figure 9. Surface charge screening does not suppress the AMIGO1 effect.

(A) Experimental set up: Whole-cell K⁺ currents (arrow) from Kv2.1–CHO transfected with GFP (red) or AMIGO1–YFP (blue). 100 mM magnesium was used to shield surface charges (peach halo). Same voltage protocol and representations as Fig. 3. (**B**, **C**) Representative Kv2.1–control (10.0 pF) or Kv2.1 + AMIGO1 (6.3 pF) cell. (**D**, **E**) Normalized *G*–*V* relationships. (**F**) Reconstructed 4th order Boltzmann fits from average $V_{i,Mid}$ and z_i (Table 1). Shading $V_{i,Mid} \pm$ SEM. (**G**) Steepness and (**H**) midpoint of 4th order Boltzmann fits. Mean \pm SEM. Statistics in Table 1. **: $p = \leq 0.01$, ns: not significant.

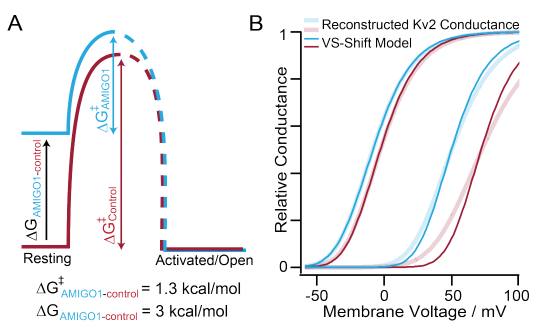


Figure 10. AMIGO1 destabilizes the resting conformation of Kv2.1 voltage sensors.

(A) AMIGO1 raises resting state energy (ΔG) of Kv2.1 voltage sensors and lowers the energy barrier (ΔG^{\ddagger}) of Kv2.1 activation. (B) Voltage sensor shift model of AMIGO1 modulation (dark lines) plotted with reconstructed *G*–*Vs* from Kv2.1–CHO Table 1 values (pale lines). From left to right: Kv2.1+AMIGO1, Kv2.1–Control, Kv2.1+AMIGO1 with GxTX–594, Kv2.1–Control with GxTX–594. Voltage sensor shift model is $f(V) = (1 + e^{-(V-V_{VSD,1/2})(z^2/25.46)})^{-4} \cdot (1 + e^{-(V-V_{POTP,1/2})(z^2/25.46)})^{-1}$, where $z = 1.5 e_0$, $V_{Pore,1/2} = -16$ mV, and $V_{VSD,1/2}$ varies. Kv2.1–Control $V_{VSD,1/2} = -33$ mV and Kv2.1–Control with GxTX–594 $V_{VSD,1/2} = 51$ mV. AMIGO1 $\Delta V_{VSD,1/2} = -22$ mV with or without GxTX–594.

		G - V fit paran	neters		ΔG_{AMIGO1} (kcal/mol)
	$V_{i,1/2}(mV)$	$V_{i,Mid}(mV)$	$z_i(e_0)$	n	(Eqn. E)
HEK293 cells					
mKv2.1 + GFP	-26.8 ± 3.0	-1.7 ± 1.4 ^A	1.79 ± 0.17 ^D	7	-0.31
mKv2.1+ AMIGO1 + GFP	-30.9 ± 0.8	-7.4 ± 1.8 ^B	1.95 ± 0.16 ^E	14	
$mKv2.1 + SCN\beta1 + GFP$	-24.8 ± 1.5	0.2 ± 1.8 ^C	1.720 ± 0.074 F	8	
Kv2.1–CHO cells					
rKv2.1 + GFP	-33.4 ± 1.7	-1.8 ± 1.2 ^G	1.411 ± 0.070 ^I	20	-0.28
rKv2.1+ AMIGO1-YFP	-42.0 ± 3.3	-7.6 ± 1.8 ^H	1.40 ± 0.11 ^J	19	
Kv2.1–CHO cells + Mg ²⁺					
rKv2.1 + GFP	-13.8 ± 1.8	17.6 ± 2.2 ^к	1.51 ± 0.11 ^M	18	-0.37
rKv2.1+ AMIGO1-YFP	-16.3 ± 1.5	10.2 ± 1.0 ^L	1.682 ± 0.082 ^N	23	
Kv2.1–CHO cells + GxTX–594					
rKv2.1 + GFP	26.8 ± 2.9	73.2 ± 3.8 ^o	1.03 ± 0.11 Q	13	-0.77
rKv2.1+ AMIGO1-YFP	12.9 ± 4.4	50.9 ± 2.8 P	1.27 ± 0.14 ^R	12	

Table 1. Fourth order Boltzmann parameters for G–V relationships.

Average $V_{i,1/2}$, $V_{i,Mid}$, and z_i values were derived from a 4th order Boltzmann fits (Eqn. C) of *n* individual cells. All values are given \pm SEM. Brown-Forsythe and Welch (appropriate for differing SD) ANOVA test with a Dunnett's T3 multiple comparisons p-values: AB: 0.046. AC: 0.64. DE: 0.75. DF: 0.91. Unpaired, two-tailed t-test p-values: GH: 0.012. IJ: 0.95. KL: 0.0051. MN: 0.21. OP: 0.00018. QR: 0.19. ΔG_{AMIGO1} from Eqn. E, at $V_{i,Mid}$ for Kv2.1 + GFP.

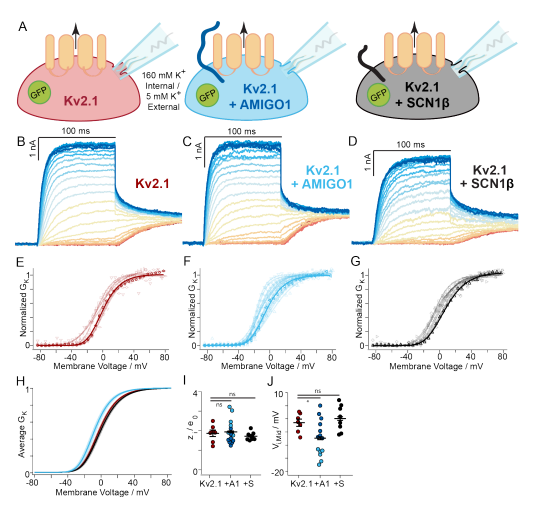
Kv2.1–CHO cells	Q-V fit parameters					ΔG_{AMIGO1} (kcal/mol)			
$Q_{ m ON, fast}$	$V_{\rm g,Mid}$	(mV)	$z_{g}(e_{0})$	п	Eqn. E				
rKv2.1 + GFP	30.6 ±	2.0 ^s	1.38 ± 0.03 ^U	20	-1.92				
rKv2.1+ AMIGO1-YFP	17.8 ±	2.9 ^T	1.61 ± 0.05 V	20	-1.92				
$Q_{ m OFF}$	$V_{\rm g,Mid}(\rm mV)$	$V_{g,Med}(mV)$	$z_{\rm g}(e_0)$	п	Eqn. E	Eqn. K*	Eqn. K*°		
rKv2.1 + GFP	-22.0 \pm 1.3 $^{\rm W}$	-19.5	$2.00\pm0.13\ ^{\rm Y}$	20	-2.45	-3.11 ± 0.69	-2.74		
rKv2.1+ AMIGO1-YFP	-32.8 ± 2.0 ^x	-29.0	2.43 ± 0.15 ^z	20	-2.43	-2.74			

Table 2. Boltzmann parameters and ΔG calculations for voltage sensor movement.

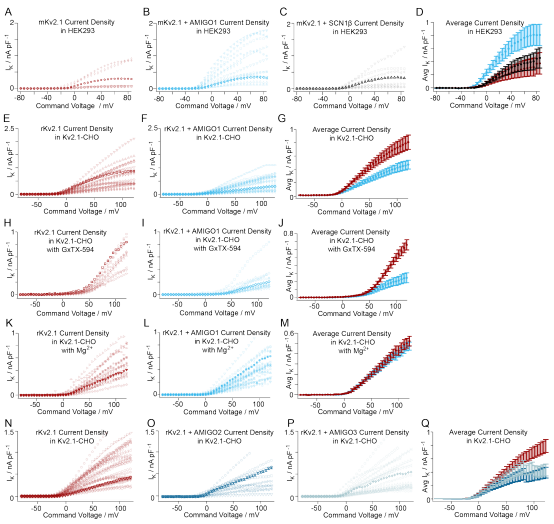
Average $V_{g,Mid}$ and z_g values were derived from 1st order Boltzmann fits of *n* individual cells. Means ± SEM. $V_{g,Mid} = V_{g,1/2}$. $V_{g,Med}$ is median voltage (58). Unpaired, two-tailed t-test p-values: $Q_{ON,fast}$: ST: 0.00093. UV: 0.00084. OFF Gating currents: WX: 7.82x10⁻⁵. YZ: 0.038. * $z = 12.5 e_0$, ° $V_{g,Med}$ was used.

Calculated activation of native Kv2 conductance after 1.5 ms at 0 mV								
Type of AMIGO1 impact	$ au_{0mV}(s)$	$V_{\rm Mid}({ m mV})$	Relative Conductance					
none, values from (7)	0.0029	-13.1	0.022					
from conductance data	0.0040^{\dagger}	-7.1	0.0067					
from voltage sensor data	0.0050^{\dagger}	-2.3	0.0024					

Table 3. Prediction of AMIGO1 impacts on Kv2 conductance in superior cervical ganglion neurons. Liu and Bean fit Kv2 kinetics with $(1 - e^{-t/\tau_{0}mv})^4$ and the G-V with $(1 + e^{-(V-V_{Mid})/k})^{-1}$, and these equations are used to calculate relative conductance here τ_{0mv} and ΔV_{Mid} adjusted for the impact of loss of AMIGO1 from Kv2.1–CHO cells. The AMIGO1 impact on conductance activation was a 1.38-fold acceleration of τ_{0mv} (Fig. 3M) and $G-V \Delta V_{i,Mid} = -5.7$ mV (Table 1). The AMIGO1 impact on voltage sensor activation was a 1.74-fold acceleration of τ_{0mv} (change in α_{0mv} from fit in Fig. 6N) and $Q_{OFF}-V \Delta V_{g,Mid} = -10.8$ mV (Table 2).

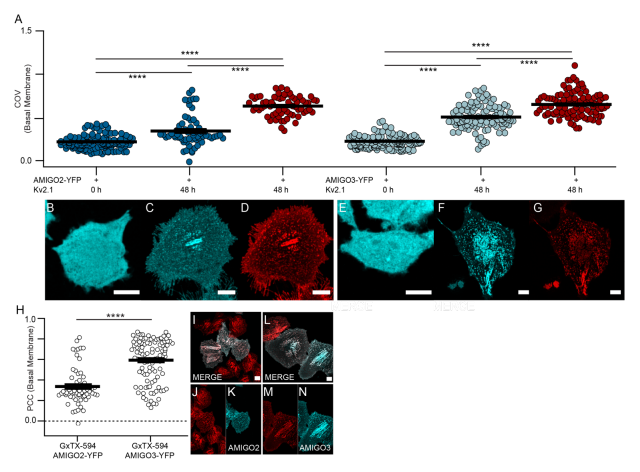


Supplemental Figure 1. AMIGO1, but not SCN1B, modulates Kv2.1 conductance in HEK293 cells. (A) Experimental set up: Whole-cell K⁺ currents from HEK293 cells co-transfected with mKv2.1 and either GFP (red), or AMIGO1-pIRES2-GFP (blue), or SCN1B-pIRES2-GFP (black). (B, C, D) Representative mKv2.1control (14.8 pF), mKv2.1 + AMIGO1 (9.6 pF), or mKv2.1 + SCN1β (10.0 pF) HEK293 cell. Data points from representative cells are bolded in analysis panels. (E, F, G) Normalized G-V relationships for mKv2.1-control, mKv2.1 + AMIGO1, or mKv2.1 + SCN1^β cells. Symbols correspond to individual cells. Lines are 4th order Boltzmann relationships (Eqn. C). (H) Reconstructed 4th order Boltzmann fits using the average $V_{i,Mid}$ and z_i (Table 1). Shaded areas represent $V_{i,Mid} \pm SEM$. (I) Steepness and (J) midpoint of 4th order Boltzmann fits. For the mKv2.1 + AMIGO1 cells, individual $V_{i,Mid}$ and z_i values are displayed in dark or light blue to highlight an increase in variability. Specifically, the standard deviation of $V_{i,Mid}$ increased from ± 3.6 mV in control cells to ± 6.9 mV in mKv2.1 + AMIGO1 cells. We note that the $V_{i,Mid}$ values for mKv2.1 + AMIGO1 cells seemed to partition into two groups: a more negatively shifted group with an average $V_{i,Mid}$ of -13.9 mV (light blue), and a group similar to mKv2.1 alone with an average $V_{i,Mid}$ of -2.5 mV (dark blue). Although all cells analyzed had GFP fluorescence indicating transfection with the AMIGO1-pIRES2-GFP vector, it is possible that some cells were not expressing sufficient AMIGO1 to have a functional effect. Statistics in in Table 1. *: $p = \leq 0.05$, ns: not significant. Bars are mean \pm SEM.

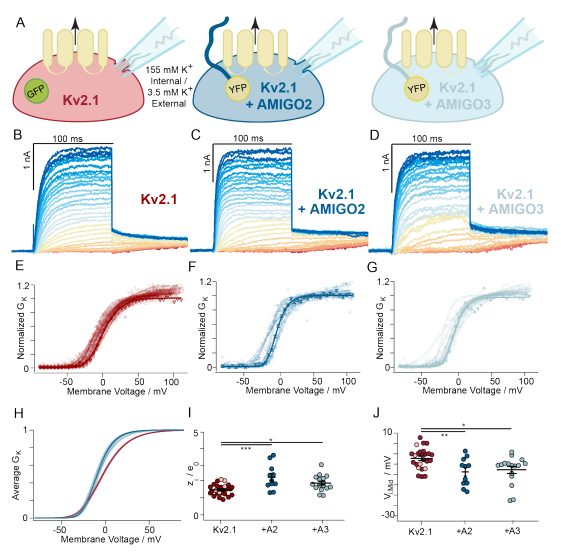


Supplemental Figure 2. Kv2.1 current density ± AMIGO1 in HEK293 and Kv2.1–CHO cells.

AMIGO1 has mixed effects on current density in HEK293 and Kv2.1–CHO cells. Outward current densities normalized by cell capacitance were calculated from mean of the last 10 ms of each voltage step and plotted against the command voltage. Symbols represent individual cells. (**A**, **B**, **C**) HEK293 cells co-transfected with mKv2.1 + GFP, mKv2.1 + AMIGO1–pIRES2–GFP, or mKv2.1 + SCN1B–pIRES2–GFP. To limit the proportion of currents from endogenous voltage-dependent channels (53, 60), we set a minimum outward current density as an inclusion threshold for analysis (65 pA/pF at +85 mV). Of the cells patched, 7 of 18 mKv2.1–control cells, 14 of 28 mKv2.1 + AMIGO1 cells, and 8 of 27 mKv2.1 + SCN1β cells satisfied this inclusion threshold and displayed currents consistent with a Kv2.1 delayed rectifier conductance (*I*_K). Cells that did not meet the inclusion criteria are not plotted making the full variability of current densities is extreme than depicted here. Bolded symbols are exemplars from Supplemental Fig. 1B, C, or D. (**D**) Averages of A, B, and C. (**E**, **F**) Kv2.1–CHO \pm AMIGO1–YFP. Bolded symbols are exemplars from Fig. 3B or 3C. (**G**) Averages of E and F. (**H**, **I**) Kv2.1–CHO \pm AMIGO1–YFP in 100 nM GxTX–594. Bolded symbols are exemplars from Fig. 5B or 5C. Cell symbols matched between E/H and F/I before and after GxTX–594 addition. (**J**) Averages of H and I. (**K**, **L**) Kv2.1–CHO \pm AMIGO1–YFP in 3.5 mM K⁺/100 mM Mg²⁺ external. Bolded symbols are exemplars from Fig. 9B or 9C. (**M**) Averages of E and F. Averaged data are means \pm SEM.

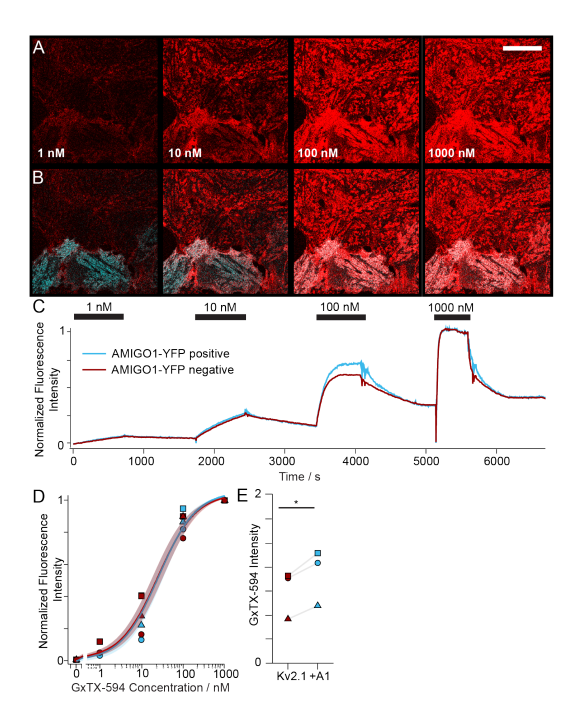


Supplemental Figure 3. Kv2.1 reorganizes and colocalizes with AMIGO homologs in CHO cells. (A) Coefficient of variation of fluorescence from AMIGO2-YFP (dark blue circles), AMIGO3-YFP (light blue circles), or GxTX-594 (red circles). COV from confocal images of glass-adhered membranes (exemplar images in **B-G**). AMIGO2–YFP fluorescence from cells (**B**) not induced for Kv2.1 expression (COV_{A2,0h} = 0.2090 ± 0.0062 , n = 144), (C) induced 48 h for Kv2.1 expression ($COV_{A2,48h} = 0.342 \pm 0.022$, n = 65). (D) GxTX-594 labeling of the cells in C (COV_{A2.48h(GxTX-594)} = 0.631 \pm 0.013, n = 65 cells). AMIGO3–YFP fluorescence from cells (E) not induced for Kv2.1 expression (COV_{A3.0h} = 0.2186 ± 0.0052 , n = 160), (F) induced 48 h for Kv2.1 expression (COV_{A3.48h} = 0.503 ± 0.014 , n = 109). (G) GxTX-594 labeling of the cells in panel F (COV_{A3,48h(GxTX-594)} = 0.650 ± 0.013 , n = 109cells). (H) Costes thresholded, Pearson's colocalization coefficients from cells induced for Kv2.1 expression 48 h prior to imaging. From left to right: $PCC_{A2,GXTX-594} = 0.342 \pm 0.022, \ge 0$ (p < 0.0001, one-tailed, t-test), n = 65; PCC A3.GxTX-594 = 0.597 ± 0.020 , ≥ 0 (p < 0.0001, one-tailed, t-test), n = 108. (I, J, K) Exemplar images where merge overlay (white) shows colocalization between GxTX-594 (red) and AMIGO2-YFP (cyan) or (L, M, N) AMIGO2-YFP (cyan) Arithmetic means and standard errors are plotted. (Statistics) Outliers were removed using ROUT, Q = 1%. An ordinary one-way ANOVA with multiple comparisons was used to evaluate the differences between groups in COV analysis, while a t-test was used to evaluate the PCC data. ****: $p = \leq 0.0001$. Bars are mean \pm SEM. All scale bars are 10 μ m.



Supplemental Figure 4. AMIGO2 and AMIGO3 modulate Kv2.1 conductance in CHO cells.

(A) Experimental set up: Whole-cell K⁺ currents (arrow) from Kv2.1–CHO transfected with GFP (red), rAMIGO2– YFP (dark blue), or rAMIGO3–YFP (light blue). Same voltage protocols and representation as Fig. 3. (**B**, **C**, **D**) Representative Kv2.1–control (5.1 pF), Kv2.1 + AMIGO2 (6.6 pF) or Kv2.1 + AMIGO3 (2.4 pF) cells. (**E**, **F**, **G**) Normalized *G-V* relationships. 5 of the Kv2.1–control cells were recorded from side by side with the Kv2.1 + AMIGO2 cells and Kv2.1 + AMIGO3 cells (light red). There was no statistical difference between these 5 cells and the data previously acquired during Kv2.1 + AMIGO1 recordings for Fig. 3 (assessed by t-test), and data was pooled. Solid lines a 4th order Boltzmann fits (Eqn. C). (**H**) Reconstructed 4th order Boltzmann fits from average $V_{i,Mid}$ and z_i (Supplemental Table 1). Shading $V_{i,Mid} \pm$ SEM. (**I**) Steepness and (**J**) midpoint of fits. Statistics in Table 1. ***: $p = \leq 0.001$, **: $p = \leq 0.01$, *: $p = \leq 0.05$. Bars are mean \pm SEM.



Supplemental Figure 5. AMIGO1 does not impede GxTX–594 binding to Kv2.1.

(A) Fluorescence from Kv2.1–CHO cells transfected with AMIGO1–YFP, induced for Kv2.1 expression for 48 hours and labeled with indicated concentrations of GxTX–594 (red). Scale bar 20 μ m. (B) Overlap (white) between AMIGO1–YFP (cyan) and GxTX–594 fluorescence. (C) Mean fluorescence intensities from ROIs encompassing AMIGO1–YFP positive or negative cells from the concentration-response experiment shown in A. (D) Normalized fluorescence intensity after 500 s at each concentration as in panel C. Symbol shapes represent data from each of 3 experiments. Curves and shaded regions represent the mean \pm SEM of a Langmuir binding isotherm (Eqn. L) fit to individual experiments. $K_d = 27.5 \pm 8.3$ nM without and 27.9 ± 7.2 nM with AMIGO1–YFP. K_d likely is overestimated due to incomplete equilibration at 1 and 10 nM. (E) Cells expressing AMIGO1–YFP had brighter GxTX–594 fluorescence with 1000 nM GxTX–594. Symbols correspond with D.

		G-V fit param	ΔG_{AMIGOX} (kcal/mol)		
	$V_{i,1/2}(mV)$ $V_{i,Mid}(mV)$ $z_i(e_0)$ n				(Eqn. E)
Kv2.1–CHO cells					
rKv2.1 + GFP	-32.5 ± 1.5	-2.0 \pm 1.0 $^{\rm A}$	1.471 ± 0.067 ^D	25	
rKv2.1+ AMIGO2-YFP	-29.7 ± 3.4	-8.7 ± 2.1 ^B	2.25 ± 0.23 $^{\mathrm{E}}$	11	-0.39
rKv2.1+ AMIGO3-YFP	-31.8 ± 2.4	-7.8 ± 1.7 ^C	1.88 ± 0.12 F	16	-0.31

Supplemental Table 1. Fourth order Boltzmann parameters for G–V relationships of AMIGO homologs.

Average $V_{i,1/2}$, $V_{i,Mid}$, and z_i values were derived from a 4th order Boltzmann fits (Eqn. C) of *n* individual cells. All values are given \pm SEM. Ordinary one-way ANOVA test with Dunnett's multiple comparisons p-values: AB: 0.0082. AC: 0.010. DE: 0.0002. DF: 0.026. ΔG_{AMIGO1} from Eqn. E, at $V_{i,Mid}$ for Kv2.1 + GFP.

GxTX(JP) conjugate	AMIGO1– YFP Expression	fitting component	aO	a1	a2	a3	a4	a5	R ²
	- AMIGO	1	0.229	670.4	47.88	11.41	1.075	2.323	0.000
GxTX	- AMIGO	2	0.813	647.0	25.73	21.77	0.631	1.685	- 0.999 - 0.997 - 0.998
Ser13Pra(JP)	+ AMIGO	1	0.893	646.7	23.30	25.63	1.822	0.721	0.997
		2	0.006	-1610	-15206	-1877	4967	461.2	
		1	0.352	594.3	12.11	-11.53	0.568	5.364	0.009
GxTX Lys27Pra(JP)	- AMIGO	2	0.719 608.2 9.71	9.71	59.05	0.359	-0.264	0.998	
		1	0.715	597.8	16.07	18.08	1.578	2.912	0.997
	+ AMIGO	2	0.632	616.3	9.05	26.28	-1.657	1.488	

Supplemental Table 2. Split Pseudo–Voigt fitting parameters.

Fluorescence emission spectra split pseudo-Voigt fitting parameters and root-mean squared values.

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Figure	# Transfections	<i>n</i> per transfection					
Fig. 3	7	peGFP: 5, 2, 2, 4, 1, 2	, 4	+AN	4IGO1: 3, 3, 3, 4, 3, 2, 1		
Fig. 4	6	peGFP: 2, 1, 1, 1, 1,	2	+A	MIGO1: 1, 2, 1, 3, 0, 0		
Fig. 5	5	peGFP: 4, 2, 2, 3, 2		+AMIGO1: 3, 3, 1, 3, 2			
Fig. 6	6	peGFP: 5, 4, 4, 2, 1,	4	+ AMIGO1: 2, 3, 4, 1, 4, 6			
Fig. 7	2	AMIGO1 (-): 6, 5		AMIGO1 (+): 5, 6			
Fig. 9	4	peGFP: 1, 3, 4, 10		+AMIGO1: 5, 5, 7, 6			
Sup. Fig. 1	4	peGFP: 3, 3, 1, 0 +AMIGO		1:4,4,6,0	+SCNB1: 1, 1, 2, 4		
Sup. Fig. 4	5	peGFP: 5, 0, 0, 0, 0 +AMIGO2:		1, 2, 0, 1, 7	+AMIGO3: 1, 7, 5, 0, 3		
		(+peGFP n-values from Fig. 3)					

Supplemental Table 3. N-values for electrophysiology experiments.

Figure	# Transfections	# n values per transfection						
Fig. 1	4	YFP (0 hr): 28,	YFP (1.5 hr):	YFP (48 hr):	YFP (ChR):	GxTX-594 (48	mRuby-ChR	
_		48, 0, 58	25, 55, 42, 95	82, 54, 74, 67	11, 21, 32, 61	hr, AMIGO1):	(AMIGO1):	
						84, 44, 69, 0	20, 16, 32, 60	
Fig. 2	4	AMIGO	1-YFP +GxTX-594	4 (48 hr):	AMIGO1-YFP +ChR-mRuby:			
		85, 41, 69, 0 18, 22, 28, 61						
Fig. 2	3	0 hr: 41, 35, 25 1.5 hr: 38			8, 39, 41	48 hr: 28, 17, 56		
Fig. 8	3	AMIGO1(-)	(GxTX Ser27Pra-J	P): 20, 12, 8	AMIGO1(+) (GxTX Ser27Pra-JP): 39, 20, 13			
	2	AMIGO1(-	-) (GxTX Ser13Pra-	-JP): 15, 55	AMIGO1(+) (GxTX Ser13Pra-JP): 7, 62			
Sup. Fig. 3	2	AMIGO2-YFP	AMIGO2-YFP	GxTX-594 (48	AMIGO3-YFP	AMIGO3-YFP	GxTX-594 (48	
		(0 hr):	(48 hr):	hr, AMIGO2):	(0 hr):	(48 hr):	hr, AMIGO3):	
		28, 116	59, 6	59, 6	117, 43	109, 0	109, 0	
Sup. Fig. 3	2	AMIGO2-YFP +GxTX-594: 64, 1			AMIGO	03-YFP +GxTX-59	4: 108,0	
Sup. Fig. 5	3		AMIGO1(-): 1, 1, 1			AMIGO1(+): 1, 1, 1	l	

Supplemental Table 4. N-values for imaging experiments.