PHOSPHATIDYLINOSITOL 4,5-BISPHOSPHATE (PIP₂) REGULATES KCNQ3 K⁺ CHANNELS THROUGH MULTIPLE SITES OF ACTION Frank S. Choveau¹, Victor De La Rosa¹, Sonya M. Bierbower¹, Ciria C. Hernandez^{2,3*} and Mark S. Shapiro^{1*}

Department of Cell and Integrative Physiology¹, University of Texas Health at San Antonio, San Antonio,
 Texas; Department of Neurology², Vanderbilt University Medical Center, Nashville, Tennessee, and Life
 Sciences Institute³, University of Michigan, Ann Arbor, Michigan.

10

1

2 3

4 5

6

*Co-corresponding authors:

Mark S. Shapiro, Ph.D.	Ciria C. Hernandez, M.D., Ph.D.
Department of Cell and Integrative Physiology	Life Sciences Institute
University of Texas Health at San Antonio	University of Michigan
STRF, MC 8253	Room 6115
8403 Floyd Curl Drive	210 Washtenaw Ave.
San Antonio, TX 78229 USA	Ann arbor, MI 48109-2216 USA
Tel: (210) 562-4092	Tel: (734) 615-1867
San Antonio, 1X 78229 USA	Ann arbor, MI 48109-2216 USA
Tel: (210) 562-4092	Tel: (734) 615-1867
Fax: (210) 562-4060	Fax: (734) 647-9702
Email: shapirom@uthscsa.edu	Email: ciria@umich.edu

- 13
- 14 Running title: Sturcutural Determiniants of PIP₂ regulation of KCNQ3 channels
- 15
- 16 Keywords: M current, ion channel modulation, potassium channel, lipid signaling, structure
- 17

18 ABSTRACT (201)

- 19 Phosphatidylinositol 4,5-bisphosphate (PIP₂) regulates the function of many ion channels, including M-type
- 20 (KCNQ1-5, Kv7) K⁺ channels; however the molecular mechanisms involved in this regulation remain
- 21 unclear. To identify the sites of action on KCNQ3 channels, we used as our baseline the A315T pore mutant
- 22 (KCNQ3T) that increases channel currents without modifying the apparent affinity of PIP₂ and performed
- 23 extensive mutagenesis in regions that have been suggested to be involved in PIP₂ interactions among the
- 24 KCNQ family. Using the zebrafish (Danio rerio) voltage-sensitive phosphatase to deplete PIP₂ as a probe
- 25 for apparent affinity of the channels, we found that PIP_2 modulates KCNQ channel function through four
- 26 different domains. 1) the A-B helix linker that we previously identified as important for both KCNQ2 and
- 27 KCNQ3, 2) the junction between S6 and the A helix (S6Jx), 3) the S2-S3 linker and 4) the S4-S5 linker. We
- found that PIP_2 interactions within these domains were not coupled to the voltage dependence of activation.
- 29 Extensive homology modeling and docking simulations between the wild-type or mutant KCNQ3 channels
- 30 and PIP₂, correlated with the experimental data. Our results indicate that PIP_2 modulates KCNQ3 channel
- 31 function by interacting synergistically with a minimum of four cytoplasmic domains.

3

32 INTRODUCTION

33 Voltage gated potassium (Kv) channels play critical roles in the function of various tissues including 34 brain, heart and epithelia (Jentsch, 2000). Among Kv channels, KCNQ1-5 (Kv7.1-7.5) channels are 35 regulated by a number of intracellular signaling molecules including phosphatidylinositol 4,5-bisphosphate 36 (PIP₂), which is present in the inner leaflet of the cell plasma membrane at only modest abundance. For 37 some time, it has been known that interactions with PIP₂ regulate M-channel activity (Suh and Hille, 2002; 38 Loussouarn et al., 2003; Zhang et al., 2003; Li et al., 2005; Winks et al., 2005; Gamper and Shapiro, 2007). 39 However, several key questions remain elusive: How and where does PIP₂ regulate KCNQ channels, and are 40 those mechanisms disparate between KCNQ1-containing channels and the others, or do they generalize 41 among KCNQ1-5? To understand the molecular mechanisms by which PIP₂ regulates KCNQ channels, it is 42 necessary to identify the site(s) of PIP₂ action. Kv channels are tetramers of subunits containing six 43 transmembrane domains (S1-S6). The earliest study suggested that PIP₂ interacts with the junction between 44 S6 and the first C-terminal "A helix" (which we call the S6Jx domain) of KCNQ2; thus, replacement of the 45 histidine at position 328 in the S6Jx of KCNQ2 (H367 in KCNQ3, Fig. 1A) by a cysteine reduced the 46 sensitivity of the channel to PIP₂ (Zhang et al., 2003). We identified a "cationic cluster" (K452, R459 and 47 R461 in KCNQ2) in the linker between the A&B helices (A-B linker) of KCNQ2, KCNQ3 and KCNQ4, 48 which were suggested to form electrostatic bonds with the phosphate head groups of PIP₂ molecules 49 (Hernandez et al., 2008). Expanding on those findings, Tinker and co-workers localized a cluster of basic 50 residues (K354, K358, R360 and K362) in the S6Jx of KCNQ1 channels (Thomas et al., 2011). In KCNQ3, 51 the analogous K358, Q362, R364, and K366 residues (Fig. 1A) were suggested to interact with PIP₂. More 52 recent work has suggested two additional domains that interact with PIP₂ and regulate gating, the linker 53 between transmembrane helices S2 and S3 (S2-S3 linker) and between the S4 and S5 helices (S4-S5 linker), 54 domains in KCNQ1 channels whose interactions with PIP₂ were suggested moreover to be important for 55 coupling between the voltage sensor and the gate (Zaydman et al., 2013). Lastly, a recent study based heavily on molecular dynamics simulations suggested state-dependent interactions between PIP2 and the S2-56 57 S3 and S4-S5 linkers of KCNQ2 channels that were weakly coupled to the voltage dependence of activation 58 (Zhang et al., 2013).

59 Lending support for a generalized structural interaction between PIP₂ and the region just distal to the final transmembrane helix of K^+ channels is the crystal structure of PIP₂ bound to the K_{ir}2.2 channel 60 61 (Hansen et al., 2011), which shows a PIP_2 molecule interacting with residues not only in the proximal C-62 terminus as it emerges from the lipid bilayer, but also residues at the distal end of the M2 helix. Thus, it 63 behooved us to more systematically examine all of these regions of a KCNQ channel most amenable to study via a voltage-dependent phosphatase (VSP), which can dephosphorylate nearly all the PIP₂ in the 64 65 plasma membrane within about 500 ms (Murata and Okamura, 2007). This method has been exploited to 66 examine the PIP₂ sensitivity of KCNQ (Falkenburger et al., 2010; Kruse et al., 2012) and TRP (Itsuki et al., 67 2012) channels, among others. Most significantly, unlike reducing PIP2 abundance by stimulating a Gq- and phospholipase C (PLC)-coupled receptors, which could also produce inositol triphosphate (IP₃), Ca^{2+} rises, 68 69 activate protein kinase C and induce other downstream signals, activation of VSP only dephosphorylates 70 PIP₂ to PI(4)P, a singly-phosphorylated lipid that does not allow activation of M channels (Telezhkin et al., 71 2012; Telezhkin et al., 2013; Telezhkin et al., 2012).

72 The Hille group studied KCNQ2/3 heteromers and found that the time constant of dephosphorylation 73 of available PIP₂ in the membrane of a tissue-culture cell is ~ 250 ms (Falkenburger et al., 2010); in that 74 work, while not quantifying the k_{on} or k_{off} of PIP₂, they found a "dwell time" of ~10 ms to be consistent with 75 the modeling of their data, most likely due to the low-affinity of KCNQ2 subunits that determine whether 76 KCNQ2/3 channels are open or closed. Hence, mutations that decrease their apparent affinity of PIP_2 , 77 resulting in "dwell times" necessarily shorter than 10 ms in KCNKQ2-containing channels can not possibly 78 be meaningfully quantified during the decay of the current during the depolarization step to a very positive 79 potential that activates Dr-VSP, since any shorter k_{off} would be wholly confounded by the time required for PIP₂ dephosphorylation. In such a case, only any altered rate of recovery of the current, reflecting an altered 80 81 kon, could be meaningful. Thus, such relatively low PIP₂ apparent-affinity channels are unsuitable for this 82 approach. For these reasons, we choose KCNQ3 homomers as our test channel, due to its extremely high 83 apparent affinity for PIP₂, as manifested by its saturating open probability near unity at saturating voltages, 84 and its maximal depression by M_1 receptor stimulation of only ~40% (Li et al., 2005; Hernandez et al., 85 2009), vs. <0.3, and 90%, respectively, for all other KCNQ isoforms and compositions. Our assumption was

4

that that this channel would be amenable to such analysis using the VSP approach, and that high structural and mechanistic similarity with the other KCNQ subtypes should make our data generalizable among this K^+ channel family. In some experiments we used the alternative assay of quantifying the extent of depression of the current by stimulation of muscarinic M₁ receptors (M₁Rs) co-expressed with the channels (*see below*).

91 In our patch-clamp experiments, we used the well-expressing KCNQ3 A315T (KCNQ3T) channel as 92 a baseline, an inner-pore mutant that increases current amplitudes by >10-fold (Zaika et al., 2008; Choveau 93 et al., 2012), without changing the open probability of the channels or their apparent PIP₂ affinities 94 (Hernandez et al., 2009). We probed the effects of charge neutralization in the S2-S3 linker, the S4-S5 95 linker, the S6Jx domain and the A-B helix linker on changes in the apparent PIP₂ affinity of the channels, as 96 well as their voltage dependence of activation. In addition, homology modeling and PIP₂-docking 97 simulations were performed to seek a structural framework for our experimental results. We find that all the 98 regions tested complement the PIP₂-binding "cationic cluster" previously described in the A-B helix linker 99 for KCNQ2 and KCNQ3 (Hernandez et al., 2008). Whereas the four domains identified here for KCNQ3 as 100 interacting with PIP₂ are conserved with KCNQ1, and likely, KCNQ2, mutations that lower the apparent 101 affinity of the channels for PIP₂ were not correlated with alterations in voltage dependence.

102

134

103 **Results**

104 We choose Danio rerio (Dr)-VSP because it activates at +40 mV, well positive to the saturating 105 voltage for all KCNQ channels. Upon activation of Dr-VSP by depolarization to +120 mV, which 106 dephosphorylates PIP_2 into PI(4)P, quantification of the rate of decay of the current provides an estimate of 107 changes in k_{off} of PIP₂ from the channels due to mutations. We realize that this is an approximation, due to confound of the known rate of Dr-VSP dephosphorylation of PIP₂ by Dr-VSP at that voltage ($\tau \sim 250$ ms). 108 109 However, the deconvolution of those rates is beyond the scope of this paper; moreover, we would need 110 information on the allosteric influence of the binding of one PIP₂ molecule with one subunit on its affinity 111 with another, and the precise number of PIP₂ molecules required for the opening of KCNQ3 homomers, and 112 both sets of data are lacking at this time. Upon the step back to +30 mV, changes in k_{on} of PIP₂ due to 113 mutations were estimated by the rate of recovery of the current. We again realize that this estimate is an 114 approximation due to the confound of the known rate of PI(4)P-5 kinase ($\tau \sim 10s$) (Falkenburger et al., 115 2010). Again, a more sophisticated deconvolution would be extremely difficult without more information, 116 which is not presently available.

117 Besides the measurements described above, we also compared the amplitude of tonic whole-cell 118 currents between cells transfected with KCNQ3T and mutant KCNQ3T channels, and the voltage 119 dependence of activation. The first measurement is based on the correlation between the tonic open 120 probability at the single-channel level, macroscopic current amplitudes and PIP₂ apparent affinity, observed 121 for KCNQ2, KCNQ2/3, KCNQ3 and KCNQ4 channels (Li et al., 2005; Hernandez et al., 2009) and other 122 PIP₂-regulated channels, e.g., GIRK channels as well (Bender et al., 2002; Winks et al., 2005). The voltage 123 dependence of activation is important because whether PIP₂-mediated depression of KCNQ1-containing channels is accompanied by altered voltage dependence is still open to debate (Loussouarn et al., 2003; 124 125 Zaydman et al., 2013) and PIP₂-mediated modulation of KCNQ2/3 channels does not change the voltage 126 dependence of activation (Shapiro et al., 2000; Nakajo and Kubo, 2005; Suh et al., 2006). Whereas the A-B 127 helix linker "cationic" cluster domain identified in PIP₂ interactions with KCNO2 and KCNO3 in our 128 previous work (Hernandez et al., 2009) is not conserved in KCNQ1, the S6Jx, S4-S5 linker and S2-S3 linker PIP₂-interaction domains are conserved, which for KCNQ1 were suggested to form a network of PIP₂-129 130 interacting domains that are involed in voltage-sensor/gate coupling to voltage dependence (Zaydman et al., 2013). We were therefore keen to investigate these issues for the case of KCNQ3, which is found primarily 131 132 in neurons, as opposed to cardiomyocytes or epithelia. With the parameters and assumptions given, we can 133 now present the data.

135 Interactions of PIP₂ with the S2-S3 and the S4-S5 linkers in KCNQ3

Several recent studies have suggested a potential role of the S2-S3 and the S4-S5 linkers in PIP₂KCNQ channel interactions (Park et al., 2005; Zaydman et al., 2013; Zhang et al., 2013; Zhou et al., 2013).
Since these sites are most novels in terms of PIP₂ interactions suggested for KCNQ2-5 channels, we begin
here. For KCNQ1, the interactions with the S2-S3 linker involve R190 and R195, and for the S4-S5 linker,

involve R243, H258 and R259. The sequence alignment of the S2-S3 and the S4-S5 linkers among KCNQ1-140 141 5 channels (Fig. 1A) indicates R190, R195 and H258 residues to be conserved. We tested the effect of 142 charge neutralizing mutations at the analogous positions, R190 and R195, in the S2-S3 linker, and H257 143 (corresponding to H258 in KCNQ1) in the S4-S5 linker of KCNQ3T (Figs. 1A, B, C; Fig. 2). In the S2-S3 linker, the R190Q mutation, but not R195Q, decreased current densities from 197 ± 6 pA/pF to 66 ± 12 144 pA/pF (n = 6, p < 0.001) (Figs. 2A, B; Table 1). Using the Dr-VSP assay, we found the rate of current decay 145 upon depolarization that turns-on Dr-VSP for the R190Q mutant (0.35 ± 0.11 s, n = 8, p < 0.01) to be much 146 faster than for KCNQ3T (0.94 ± 0.13 s, n = 10) (Figs. 2D, E); however, the rates of recovery of the current 147 148 were 7.5 ± 1.7 s and 9.6 ± 1.6 s (n = 10-11) for KCNQ3T and KCNQ3T-R190Q respectively, values not 149 significantly different. The same result was obtained from the analogous mutant R190A (decay = 0.35 ± 2 s, 150 recovery = 7.5 ± 1.8 s; n = 6). Neither response was altered by the R195Q mutation. Either R190 influences 151 Koff for PIP2, but not kon, or our assay is not sensitive enough to detect changes in both rates accurately. As 152 an alternative assay, we turned to the classic M_1R -dependent depression of the current in cells coexpressing 153 M1 muscarinic receptors and KCNQ3T mutants. Since maximal M1R stimulation in tissue-culture cells leads 154 to about an 80% decrease in PIP₂ abundance, rather than to near zero when VSPs are activated (Falkenburger et al., 2010), the maximal depression of KCNQ3 currents is only \sim 30-40%, since enough PIP₂ 155 156 molecules remain in the membrane to keep most KCNQ3 channels PIP₂-bound (Suh et al., 2006; Hernandez 157 et al., 2009). Thus, for such high PIP₂ apparent-affinity channels, a change in that affinity is manifested most 158 in the fractional suppression of the current, not a shift of the dose-response relation of [agonist] vs. current suppression (Fig S1F). In these experiments, we decided to use mutants in which the arginines at positions 159 190 and 196 were mutated to alanines, instead of the highly hydrophilic and bulky glutamines, which can 160 161 interact with PIP₂ via several types of H⁺-bonds, to avoid any such confounding effects. Consistent with previous work, we found the KCNO3T current to be suppressed by a supramaximal concentration (10 µM) 162 of the receptor agonist, oxotremorine methiodide (oxo-M), by only 32.9 ± 8.9 % (n = 3); similarly for cells 163 expressing KCNQ3T-R195A the maximal inhibition was 24.9 ± 5.8 % (n = 7); whereas for KCNQ3T-164 R190A the maximal inhibition was 63 ± 13 % (n = 6; p < 0.05) indicating the R190A mutation to reduce 165 166 PIP₂ affinity, consistent with the Dr-VSP assay (Figs. 2G, S1). Neither the R190Q/A, nor the R195Q/A, mutations affected the voltage dependence of activation (Fig. 2C), suggesting that the apparent affinity of 167 168 PIP_2 for this site to be unrelated to voltage dependence.

169 We found the H257N mutation in the S4-S5 linker (Figs. 1B, C and 2) to result in strongly reduced current densities, from $197 \pm 6 \text{ pA/pF}$ to $30 \pm 3 \text{ pA/pF}$ (n = 7, p < 0.001); the rate of current decay after Dr-170 171 VSP activation was much faster than KCNQ3T (0.58 ± 0.14 s, n = 4, p < 0.05) and the rate of recovery was 172 slightly slower $(11.03 \pm 1.4, n = 2)$ although it was not suitable for analysis in most of the cells recorded, 173 probably due to the astounding shift in the voltage dependence of activation from -34.0 ± 1.9 mV in KCNO3T to 2.5 \pm 2.8 mV in H257N (n = 7-19). Thus we turned again to quantifying the result of M₁R 174 stimulation; for cells co-transfected with M1Rs and the KCNQ3T-H257N mutant, the maximal inhibition 175 was 81.6 ± 7.9 % (n = 4) (Fig. 2G, S1). Together this results indicate that the H257N mutation reduces the 176 177 apparent PIP₂ affinity of the channels.

Since the R243H mutation in the S4-S5 linker was shown to reduce the apparent affinity of KCNQ1 178 for PIP₂ (Park et al., 2005) and R243 is conserved in other KCNQ channels (R242 in KCNQ3) (Fig. 1A), we 179 180 also tested the effect of the R242A mutation on KCNQ3T channels (Fig. 2). This mutant resulted in reduced 181 current densities (146 \pm 16 pA/pF, n = 7, p < 0.001) and slowed rate of current recovery (14.2 \pm 1.7 s, n = 10, p < 0.05) in the VSP assay, nonetheless, the rate of decay upon turn-on of Dr-VSP was not significantly 182 affected (0.77 \pm 0.13 s, n = 10). M₁R stimulation inhibited the current by 64.8 \pm 9.2 % (n = 3; p < 0.01), two 183 184 fold greater than for KCNQ3T (Fig. 2G, S1). These results are consistent with a role of R242 in PIP₂ 185 interactions. This mutation resulted also in a pronounced shift of the voltage-dependence of activation 186 towards more positive potentials ($V_{1/2}$: -4.0 ± 3.2 mV, n = 7) (Fig. 2C). The adjacent mutation R243A was also tested; this mutant displayed reduced current densities as well ($56 \pm 16 \text{ pA/pF}$, p < 0.001), faster rate of 187 188 current decay upon Dr-VSP turn-on (0.59 \pm 0.19 s, p < 0.05) and slowed recovery after Dr-VSP turn-off $(13.1 \pm 1.8 \text{ s}, p < 0.05)$. Surprisingly the voltage-dependence of activation for this adjacent mutant was not 189 affected ($V_{1/2}$: -31 ± 4.7 mV, n = 4). When both arginines where mutated to alanines, the whole-cell current 190 densities were reduced ($65 \pm 11 \text{ pA/pF}$, n = 11, p < 0.001), similar to the R243A single mutant. The rate of 191 192 current decay and recovery after Dr-VSP turn-on or turn-off were significantly affected (0.45 \pm 0.11 s, p < 193 0.01 and 15 ± 1.7 s, p < 0.05; n = 5) to a greater extent than either of the single mutations. The voltage-

dependence of activation of the double mutant displayed the same positive shift as for the R242A single mutant ($V_{1/2}$: -0.2 ± 2.9 mV, n = 5). The M₁R-mediated inhibition of the double mutant was 91.6 ± 2.7 % (n = 4) (Fig. 2G). These results are consistent with an interaction of the KCNQ3 S4-S5 linker with PIP₂, which again seems not to be coupled to the voltage dependence of activation of the channels. Clearly, however, the S4-S5 linker of KCNQ3 itself is coupled to channel voltage dependence, just not in a way that involves PIP₂. A summary of the data is presented in Table 1.

200

201 Interactions of PIP₂ with the S6Jx domain in KCNQ3 channels

202 Three basic residues (K354, R360, and K362) in the S6Jx of KCNQ1, which are conserved in 203 KCNQ3 (K358, R364 and K366) (Fig. 1A), have been found to play a role in PIP₂ interactions (Eckey et al., 204 2014). In addition, Telezhkin and coworkers found the R325A mutation in KCNQ2, homologous to R360 in 205 KCNQ1 and R364 in KCNQ3, to decrease the apparent affinity of the channel for DiC8-PIP₂ (Telezhkin et 206 al., 2013), and early work implicated a role of H328 in KCNQ2, homologous to H367 in KCNQ3 (Zhang et 207 al., 2003). Since the K358, R364, K366 and H367 residues in the S6Jx domain are conserved among KCNQ 208 channels (Fig. 1A), we asked whether PIP₂ interacts with the S6Jx domain in KCNQ3T channels (Fig. 4). We found the R364A mutation to significantly decrease current amplitudes ($72 \pm 8 \text{ pA/pF}$, vs. 197 ± 6 209 210 pA/pF for KCNQ3T, n = 6, p < 0.001); whereas the K358A and K366A mutations did not (195 ± 7 and 187 211 \pm 12 pA/pF, n = 8-9, respectively) (Fig. 4A, B). As before, we measured the responses of each mutant to 212 PIP₂ dephosphorylation by Dr-VSP and the rate of recovery upon Dr-VSP turn-off and found the R364A mutation to result in a much faster decay of the current (0.14 ± 0.02 s, n = 5, p < 0.001) upon activation of 213 214 Dr-VSP, and a much slower recovery of the current $(27.7 \pm 6.9 \text{ s}, \text{n} = 5, \text{p} < 0.001)$ upon its turn off (Fig. 215 3D-F, Fig. S1 C, D, Table 1). The rates of current decay and recovery of K358A and K366A were not 216 significantly altered (Figs. 3D-F, Table 1), nor was the maximal inhibition by M_1R stimulation (25.1 ± 6.8 %, 217 n = 4).

218 We also tested the effect of the K358A and K366A mutations in combination with R364A as the 219 triple mutant KRK-AAA. The KRK-AAA mutant decreased the current amplitude similarly as did R364A (79 ± 11 pA/pF, n = 8), and such channels indicated a similarly reduced apparent affinity for PIP₂ (τ_{decay} = 220 0.29 ± 0.04 s; $\tau_{recovery} = 17.9 \pm 2.6$ s, n = 6-7, p < 0.001) using the Dr-VSP assay, (Fig. 3A, B, D-F; Table 1), 221 222 echoing the results of the single point mutants. None of these single point mutations significantly affected 223 channel voltage dependence (Fig. 3C). Strikingly, however, the KRK-AAA triple-mutation uniquely in this 224 domain resulted in channels with a voltage dependence of activation markedly shifted towards more positive 225 potentials. For KCNQ3T and KCNQ3T-KRK-AAA, the half activation potentials were: -34.0 ± 1.9 and -6.3226 \pm 2.5 (n = 7-19), respectively. We also tested the effects of the H367C mutation on KCNQ3T, which is 227 slightly downstream of R364 in the S6Jx domain. This mutation only slightly reduced current densities (138 228 \pm 5 pA/pF, n = 6, p < 0.01), but significantly increased the rate of decay of the current (0.32 \pm 0.05 s, n = 6, 229 p < 0.01) upon activation of Dr-VSP, and slowed its recovery (36.7 ± 6.9 s, n = 6, p < 0.001) upon Dr-VSP 230 turn-off, indicating an interaction of this residue with PIP₂, as shown for KCNQ2 (Zhang et al., 2003). Such 231 mutant channels displayed no significant shift in the voltage dependence of activation (Fig. 3C, Table 1). Taken together, these results strongly implicate the S6JxA domain of KCNQ3 channels as an important site 232 233 for PIP₂ interactions, as for KCNQ1 channels, and this altered apparent affinity for PIP₂ also seems not to 234 linked an altered voltage dependence of activation.

235

236 The A-B helix linker contributes strongly to the apparent affinity for PIP₂.

237 We previously identified a cluster of basic residues (K425, K432, and R434) in the linker between 238 helices A and B (A-B linker) of both KCNQ2 and KCNQ3 to be critical for PIP₂-mediated control of gating, 239 with mutations of this cluster in KCNQ2 to be somewhat more potent than for KCNQ3 (Hernandez et al., 240 2008). However, a study that deleted the A-B helix domain of KCNQ2 did not find that this deleted domain 241 reduced the PIP₂ apparent affinity for KCNQ2 channels (Aivar et al., 2012). Thus, we tested the importance 242 of this domain of KCNQ3 using the same assays as before. We found that the deletion of the A-B linker (Δ 243 linker) decreased whole-cell current amplitudes by about half ($112 \pm 10 \text{ pA/pF}$, n = 11, p < 0001) (Figs. 4A, 244 B). In cells co-expressing KCNQ3T (Δ linker) with Dr-VSP (Fig. 4D), the rate of current decay upon Dr-245 VSP turn-on was \sim 3-fold faster (0.26 \pm 0.04 s, n = 7, p < 0.001), compared to KCNQ3T (Fig. 4E, Table 1), 246 and the rate of current recovery upon turn-off of Dr-VSP was significantly slower (13.5 \pm 2.2 s, n = 7, p <

7

247 0.05) (Fig. 4F, Table 1). Such data reinforce a critical role of the helix A-B linker in PIP₂ interactions with 248 KCNQ3 channels, correlating with changes in open probability found for the triple (K425E/K432E/R434E) 249 KCNQ3 mutant within in the A-B linker previously studied in excised single-channel patches (Hernandez et 250 al., 2008). Lastly, as for the other PIP₂-interacting domains, the KCNQ3T (Δ linker) did not display any 251 significant shift in channel voltage dependence, with V_{1/2} values for KCNQ3T and KCNQ3T (Δ linker) 252 currents of -34.0 ± 1.9 mV and -32.5 ± 1.5 mV, respectively (Fig. 4C; Table 1).

253 We wondered what the result would be of combining both the RH-AC mutation within the S6Jx 254 domain with the KCNO3T (Δ linker) mutant. To our surprise, such severely mutated channels nonetheless 255 still yielded very small, but observable, PIP₂-dependent currents (Fig. 4). Thus, the whole-cell current 256 density was dramatically decreased from 197 ± 6 to 16 ± 2 pA/pF (n = 8, p < 0.001) (Fig. 4B, Table 1). PIP₂ 257 depletion induced by Dr-VSP rapidly and nearly completely abolished currents from the RH-AC/ Δ linker mutant, with a faster rate of decay upon Dr-VSP turn-on (0.53 ± 0.1 s, n = 6, p < 0.05), and a slower rate of 258 259 recovery upon turn-off of Dr-VSP (45.8 ± 5.2 s, n = 6, p < 0.001), than for KCNQ3T channels (Figs 4D-F; 260 Table 1). The small amplitude of the currents from such severely-mutated channels tested here preclude any 261 significant meaning from comparing data from those channels and those from the RH-AC or the Δ linker 262 mutant alone. They do reinforce the presence of two major PIP₂ interaction sites within the C-terminus of 263 KCNQ3 channels, one in the A-B linker, as previously reported (Hernandez et al., 2008), and the other 264 within the S6Jx domain.

Recently, the first two of a three-lysine cluster located at the end of the B-helix of KCNQ1 (K526, 265 266 K527, K528) have been identified as a critical site where CaM competes with PIP₂ to stabilize the open state 267 of KCNQ1-containing channels (Tobelaim et at., 2017a, b). Since this site is conserved in KCNQ3 (K531, 268 K532 and K533), we independently mutated the three lysines to asparagines and tested them for interaction of PIP2 using our VSP approach. Neither the current decay nor recovery, was altered by any of the three 269 270 mutations (Table 1), indicating that this basic cluster is not involved on PIP₂ binding in KCNO3. Whether 271 this site plays a role on CaM modulation of KCNQ3 channels remain to be determined. It is likely that the 272 involvement of this domain differs between KCNQ1 and KCNQ3.

274 Differences in plasma-membrane expression of KCNQ3T mutant channels do not explain altered current 275 amplitudes

276 Since we use whole-cell current amplitudes as one measure of PIP₂ sensitivity in this study, it was 277 incumbent upon us that we rule out the possibility of differential membrane expression of the mutants suggested to have altered apparent affinity for PIP₂, since this would confound our results. We and others 278 279 have found visualization of membrane proteins tagged with fluorescent proteins under total internal 280 reflection fluorescence (TIRF, evanescent wave) microscopy, which isolates emission from fluorophores 281 within 300 nm of the membrane (Axelrod, 2003), to be by far the most reliable measure of such membrane 282 expression (Bal et al., 2008; Zaika et al., 2008; Boyer and Slesinger, 2010). Under TIRF-illumination, we 283 measured the emission from YFP-tagged WT and mutant KCNQ3T channels expressed in CHO cells (Fig. 5). These data indicate that the decrease of the whole-cell current density is not due to divergent expression 284 285 of mutant KCNQ3T channels in the plasma membrane. In fact, the YFP emission from KCNQ3T (H257N) 286 is even higher than that of KCNQ3T, suggesting that the H257N mutation increases the number of channels 287 at the plasma membrane. Thus, differential membrane abundance of channel proteins does not underlie the 288 differences in macroscopic current amplitudes reported in this study.

289

273

290 PIP₂ is predicted to interact with the S4-S5 linker/S6Jx interface of KCNQ3 channels

291 Our electrophysiological data are consistent with localization of KCNQ3-PIP₂ interactions to four 292 distinct cytoplasmic locations: the A-B helix linker, the S6Jx domain, the S2-S3 linker and the S4-S5 linker. 293 In an attempt to construct a framework of these four sites into a coherent structural model of PIP₂ 294 interactions with the channels, we performed homology modelling and PIP₂ docking simulations for all of 295 the mutants studied in this work. Our overall hypothesis emerging from the experimental data supposes a 296 network of interactions between basic residues located in the S2-S3 linker, the S4-S5 linker, and the S6Jx 297 that, together with the A-B helix linker, govern the PIP₂-mediated regulation of KCNQ3 channel gating. As 298 above, we divide the channel into three basic modules: the voltage-sensor domain (VSD), comprising S2-S4, 299 the pore domain (PD), from S5-S6, and the carboxy terminus, of which the proximal half (up to the end of

the B helix) is the site of several regulatory molecules, and so we call it the regulatory domain (RD). We show models of the VSD, PD and S6Jx based on the co-ordinates of the Kv1.2 channel solved in the activated/open conformation (Khalili-Araghi et al., 2010). R190 and R195 lie within the S2-S3 linker, which is part of the VSD; R242 and H257 lie within the PD, and K358 and R364 are within the S6Jx, which our model predicts also to be in continuous interface with the PD (Figs. 1B, C; K366 and H367 are not displayed). We did not construct a model of PIP₂ binding to the A-B helix linker, due to the lack of a suitable template.

To model the putative network of interactions of PIP₂ with KCNQ3 channels, we first built structural 307 308 models of WT and mutant KCNQ3 channels, and performed PIP₂ docking simulations to the most 309 energetically favourable WT (Figs. 1D, E) and mutant KCNQ3 models (Fig. 6). It is widely thought that 310 positively-charged amino acids are mostly responsible for interactions with PIP₂. Thus, we first simulated 311 the interaction of PIP_2 in the presence of all available positive charges on the protein in the open 312 conformation of WT KCNQ3. In the preferred location for PIP₂ binding in the WT KCNQ3 model (Fig. 1E), 313 the phosphate head-group of PIP₂ is predicted to be directed towards R242 and R243 in the S4-S5 linker and 314 K358 and K366 in the S6Jx, and also predicted to form hydrogen-bond interactions with the nearby residues 315 within the same subunit in both the S4-S5 linker and S6Jx (Fig. 1E, residues in blue in Sub-D). Of note, the acyl tail of PIP₂ is predicted to be directed toward residues in the inner face of S5 (H257) and S6 (F343, 316 F344, L346, and P347) in the neighbouring subunit (Fig. 1E, residues in orange in Sub-C). Thus, PIP₂ 317 318 appears to be cross-linking neighbouring subunits, in analogy with a role for PIP₂ reported for GIRK2 channels (Whorton and MacKinnon, 2011). Taken together, our simulations find that PIP₂ is predicted to 319 320 interact with the S4-S5 linker/S6Jx interface (Fig. 1E), suggesting a mechanistic basis for the effect of 321 mutations in these regions on the favourability for activation; *i.e.*, PIP₂ interactions with the S4-S5 322 linker/S6Jx interface stabilize, and promote opening.

324 Multiple sites of PIP₂ interactions at the VSD-PD interface of mutant KCNQ3 channels

325 In line with previous studies on KCNQ1 and KCNQ2 channels (Zhang et al., 2013; Eckey et al., 326 2014), positively charged residues of the S4-S5 linker (R242 and R243) and S6Jx (K358 and K366) in the 327 same subunit (Fig. 1E, residues colored in blue), and S5 of the neighboring subunit (H257) (Fig. 1E, 328 residues colored in orange) are predicted to be involved in the interactions of PIP₂ with WT KCNQ3. 329 However, our experimental data demonstrate that mainly R190, R242, R243, H257, R364 and H367 are the 330 determinants of PIP₂ interactions, whereas K358 and K366 did not seem important. Therefore, we used our 331 model to ask whether these sites are predicted to alter PIP_2 interactions. We analyzed PIP_2 docking 332 simulations for the following mutants: R190Q, R242A, H257N, R364A, KRK-AAA, H367A, K358A, and 333 K366A (Fig. 6). Unlike WT KCNQ3, PIP₂ docking simulations of R190Q (Fig. 6A), H257N (Fig. 6C) 334 R364A (Fig. 6D), H367A (Fig. 6F), and K366A (Fig. 6H) predict a network of interactions mainly with two 335 positively charged residues of the S4-S5 linker (R242, R243) and one in S6Jx (K358) of the same subunit. 336 Simulations of KRK-AAA (Fig. 6E) and K358A (Fig. 6G) mutants predict that PIP₂ is docked similarly to R242 and R243 of the S4-S5 linker, but in those cases stabilize the network of interactions with H257 in S5 337 338 of the neighboring subunit. Noteworthy for all these mutants, R242 is predicted as a common residue in the 339 network of interactions of PIP₂. Moreover, PIP₂ docking simulations of R242A (Fig. 6B) suggest a network 340 of interactions with R243 of the S4-S5 linker, and two positively charged residues in S6Jx (H363 and K366). 341 Moreover, the R242A, H257N and KRK-AAA mutations are predicted to cause major structural 342 rearrangements in the S4-S5 linker, S5, S6 and S6Jx (Fig. S2). Again, we realize that the experimental data 343 reported little functional effects of charge neutralization of the K358 and K366 residues that might have 344 been predicted to stabilize the interactions of PIP₂ with the channels. However, the simulations of PIP₂ with 345 K358A and K366A (Figs. 6G, H) predict that whereas the orientation of PIP₂ in the inner face of S6Jx is 346 opposite of that predicted for WT channels, the predicted interactions at residues R242 and H257 are 347 predicted to preserve coupling to channel gating by maintaining coupling between the S4-S5 linker and the 348 Alternatively, as stated above, our model may not have such single-residue precision that S6JxA. 349 corresponds to a transmembrane ion channel in situ.

350

323

351 Additional sites of PIP₂ interactions at the S2-S3 interface with KCNQ3 channels

352 Given the lack of correlation between PIP_2 interactions and modification of the voltage dependence 353 of activation observed in our data, we generated additional structural models of KCNQ3 in the closed state

using as a template the co-ordinates of the Kv1.2 channel solved in the resting/closed state (Khalili-Araghi 354 355 et al., 2010). For the modeled closed KCNQ3 channels, the inositol ring of PIP₂ is predicted to be oriented 356 towards K103 in S1, R188 in the S2-S3 linker, and R227 and R230 in S4; whereas the acyl tail of PIP₂ is 357 predicted to form hydrogen bonds with residues in S2 and S4 within the same subunit (Figs. 7A, B, C). To correlate these predictions with function, we performed additional patch-clamp experiments, assaying the 358 359 effect of charge-neutralizing mutations on the apparent PIP₂ affinity of KCNQ3T, again using the Dr-VSP 360 approach. We found that substitution of these positively charged residues with an alanine significantly 361 accelerated the rate of decay of the current upon turn-on of Dr-VSP, compared to KCNQ3T. For KCNQ3T, 362 KCNQ3T-K103A, KCNQ3T-R188A, KCNQ3T-R227A and KCNQ3T-R230A, the rate of decay was 363 respectively 0.84 ± 0.13 s, 0.29 ± 0.05 s, 0.48 ± 0.11 s, 0.18 ± 0.03 s and 0.20 ± 0.03 s, respectively (n = 5-364 11, Fig. 7D). All the point mutants displayed a slower rate of recovery compared to KCNQ3T. We then 365 wondered if combining the K103A and R188A and the R227A and R230A double mutations would result in 366 a synergistically greater reduction in apparent PIP_2 affinity than either mutation alone. We found the rate of 367 current decay upon turn-on of Dr-VSP of KCNQ3T-K103A-R188A to be 2-fold faster (0.39 ± 0.05 s, n = 11, 368 p < 0.05) than that of KCNQ3T channels and intermediate between the K103A (0.29 ± 0.05 s, n = 7) and R188A (0.48 ± 0.11 s, n = 5) mutants, whereas that of the R227A-R230A double mutant was 3-fold faster 369 370 $(0.27 \pm 0.04 \text{ s}, n = 8, p < 0.01)$ than that of KCNQ3T, but slower than those from single R227A and R230A 371 mutants. Finally, both double mutants displayed a slower current recovery after turn-off of Dr-VSP (25.4 \pm 372 3.4 s, n = 11, p < 0.05 and 28.6 \pm 5.0 s, n = 8, p < 0.01) than KCNQ3T, and quite similar to those from 373 single mutants. These data suggest that K103 in S1, R188 in the S2-S3 linker as well as R227 and R230 in 374 S4 play a role in PIP₂ interactions with KCNQ3 but they do not act sinergestically. These data are also 375 consistent with the predictions of our modeling/docking simulations, giving us further confidence in the 376 accuracy of our modeling. Interestingly, R188 is conserved in KCNQ2 but not in other KCNQ channels, 377 suggesting that this residue may also interact with PIP2 in KCNQ2. Unlike R188, R227 is conserved in all 378 KCNQ channels and may also be critical for PIP₂-binding to KCNQ1-5 channels.

380 **DISCUSSION**

379

381 In the present work, we investigate the molecular determinants involved in the regulation of KCNQ3 382 channels by PIP₂. Many studies have investigated the sites of action of PIP₂ on ion channels, including 383 voltage-dependent K⁺ channels (Kv). However, the location of these sites remains controversial. For KCNQ2 and KCNQ3 channels, we have previously highlighted critical PIP₂ interaction domains in the A-B 384 385 helix linker (Hernandez et al., 2008). Others have identified the S6Jx domain as important for KCNQ1-3 (Peroz et al., 2008; Telezhkin et al., 2013; Zaydman et al., 2013), and our results here are in accord with 386 387 those reports. Recent work studying KCNQ1-containing channels has illuminated important PIP₂-interaction 388 domains in the S2-S3 and S4-S5 linkers that play a role in coupling to gating (Zaydman et al., 2013; 389 Zaydman and Cui, 2014; Kasimova et al., 2015). This study is in accord with those findings as well for 390 KCNQ3, in terms of there being additional domains of PIP₂ interactions. Another recent study suggested 391 that the voltage dependence of KCNQ2 channels is regulated via PIP₂ interactions with the S2-S3 and S4-S5 392 linkers (Zhang et al., 2013). We do not find similar results for KCNQ3. Finally, another group recently 393 suggested that deletion of the A-B linker does not affect the apparent affinity of KCNQ2 for PIP₂ (Aivar et 394 al., 2012); however, in retrospect, we wonder if the VSP method is well applicable for such low PIP₂-395 affinity channels, given the extremely brief "dwell time" that PIP₂ must have for them, and correspondingly 396 high k_{off} rate, especially compared to the rate of PIP₂ dephosphorylation. Finally, the current work here, 397 studying KCNQ3, is consistent with our earlier studies implicating the importance of A-B linker domain 398 (Hernandez et al., 2008).

399

400 *Comparison of the regions of KCNQ1-3 channels contributing to PIP₂ interactions.*

The present work, reporting R364 and H367 mutations of KCNQ3T, corresponding to R325A and H328C in KCNQ2, to being also highly involved in PIP₂ interactions, is in accord with previous work on KCNQ2 (Telezhkin et al., 2013; Zhang et al., 2013). For the family of PIP₂-regulated inward rectifier K⁺ (K_{ir}) channels, the JxS6 domain of KCNQ channels is analogous to the C-terminal domain just after M2, which has long been identified as a hot-spot for PIP₂ interactions by mutagenesis studies (Logothetis et al., 2007) and confirmed by the solved crystal structure of PIP₂ bound to GIRK2 channels (Whorton and

407 MacKinnon, 2011). Remarkably, our simulation studies predict that PIP_2 is stabilized between neighbouring 408 subunits in the S6Jx, which is similar to that reported for GIRK2 channels in the similar domain. Hence, we 409 suppose this structural mechanism to be likely conserved among PIP_2 -regulated channels in general. We 410 speculate that the dual A&B helices, both containing calmodulin-binding domains, possessed by KCNQ, but 411 not K_{ir}, channels, endow the A-B linker of KCNQ channels as a more unique site of PIP_2 interactions, for 412 reasons that will likely require more structural studies of these proteins.

Although our results here also show PIP₂ interactions with the S2-S3 and S4-S5 linkers in the VSD of 413 414 KCNQ3, as for KCNQ1, and that small, yet definite PIP₂-sensitive and voltage-gated currents are still 415 produced by KCNQ3T channels mutated to lack interactions with both domains in the C-terminus, we do 416 not find the interactions with the S2-S3 and S4-S5 linkers to be coupled to modifications of voltage 417 dependence. Since the work on KCNQ1 channels showed that such linkage to PIP₂ was not via alterations in 418 the sensitivity of the voltage sensor, but rather to the efficiency of coupling between the voltage sensor and 419 the gating machinery (Zaydman et al., 2013), we hypothesize that the role of PIP_2 interactions in such 420 coupling is probably similar in nature between KCNQ1 and KCNQ3, and likely KCNQ2 as well. 421 Interestingly, a striking difference between KCNQ1-containing channels and KCNQ2-4, is that whereas currents from the latter are depressed by Ca²⁺/calmodulin, those of the former are enhanced (Gamper and 422 423 Shapiro, 2003; Chambard and Ashmore, 2005; Gamper et al., 2005; Shamgar et al., 2006; Zaika et al., 2007; Kosenko and Hoshi, 2013; Sachyani et al., 2014). Given that both critical PIP₂-interaction domains in the C-424 terminus of KCNQ1-3 channels are very likely to be surrounded by Ca²⁺/calmodulin, we are very interested 425 426 to learn the relationship between calmodulin and PIP₂ interactions and voltage-dependent coupling, and the perhaps subtle yet important differences that confer opposite effects of Ca²⁺ loading of calmodulin on the 427 428 function of KCNQ1-containing channels, vs. KCNQ2-4.

429 The basic residues of both S2-S3 and S4-S5 linkers are highly conserved among KCNO channels. In our 430 experiments, K103A, R188A, R190Q, R227A and R230, but not the R195Q or R195A mutations, in S1, the 431 S2-S3 linker and in S4 induced a decrease of the apparent affinity for PIP₂. K162 in the S2-S3 linker of 432 KCNQ2 has been implicated in PIP₂-channel interactions in the closed state, supported by molecular 433 dynamics simulations (Zhang et al., 2013). Our PIP₂ docking simulations of KCNQ3 channels also suggest 434 that PIP₂ interacts with S1 (K103), the S2-S3 linker (R188 and R190) and S4 (R227 and R230) of closed 435 KCNQ3 channels. In the simulations of KCNQ3 (R188A and R190Q), PIP₂ was predicted to interact with 436 the S2-S3 linker and to lose inter-subunit contacts, which might favor channel deactivation. As opposed to 437 previous observations in Shaker and Kv1.2 channels in which the S2-S3 linker has been suggested to 438 interact with PIP₂ preferentially in the closed state (Abderemane-Ali et al., 2012; Rodriguez-Menchaca et al., 439 2012), our experimental results did not show a clear state dependence of $KCNQ3/PIP_2$ interactions. The 440 modeling/ docking simulations are consistent with the opening of KCNQ3 channels involving PIP₂ 441 interactions at the VSD-PD interface, consistent with PIP₂/KCNQ channel interactions involving a complex 442 network of basic residues along the VSD-PD interface and the C-terminus that cooperatively favor opening. 443 They also suggest that a structural mechanism of channel opening involves PIP₂-mediated inter-subunit 444 interactions. Interestingly, such PIP₂-channel interactions have also been described in the crystal structures 445 of Kir2.2 and GIRK2 (Kir3.2) channels, corresponding to the S4-S5 linker, pore domain and the C-terminus 446 in KCNQ channels (Hansen et al., 2011; Whorton and MacKinnon, 2011). Although we do not here find the 447 involvement of PIP₂ interactions with the S4-S5 linker per se to be coupled to voltage dependence of 448 activation, our electrophysiological data and our homology modeling for KCNQ3 are fully in accord with S4-S5 linker and S6 to be critical in the coupling between the VSD and the pore domain, as is generally 449 450 widely seen for voltage-dependent K⁺ channels (Long et al., 2005; Chen et al., 2010; Choveau et al., 2011; 451 Labro et al., 2011; Zaydman et al., 2013).

452 Since only charge neutralizing mutations in the S4-S5 linker (R242A and H257N) and the S6Jx 453 (K358A/R364A/K366A), reduced PIP₂ apparent affinity and shifted the voltage dependence of KCNQ3 454 towards more depolarized potentials, we hypothesize that (i) co-operation between the S4-S5 linker and the S6Jx stabilizes opening of KCNQ3 and (ii) PIP_2 likely plays a role in this coupling, a hypothesis consistent 455 456 with the Kv1.2-2.1 crystal structure in which anionic lipids are bound at the VSD-PD interface of the 457 channel (Long et al., 2007). However, one central question remains unclear as to generality among K^+ 458 channels: Does PIP₂ affect the voltage-sensor movement and by that mechanism, the voltage dependence of 459 Kv channels, or do any effects of PIP₂ on channel voltage dependence generally arise from changes in 460 coupling between the VSD and the PD? In Kv1.2, replacement of an arginine with a glutamine (R322Q) in

461 the S4-S5 linker, which is involved in VSD-PD coupling, affected the channel voltage dependence of 462 activation when PIP₂ was depleted. Moreover, gating current experiments showed that PIP₂ affects the VSD 463 movement of Shaker channels through interactions with the S4-S5 linker (Rodriguez-Menchaca et al., 2012). 464 However, unlike for Shaker, depletion of PIP₂ does not affect VSD movement of homomeric KCNQ1 465 (Zaydman et al., 2013). Different labs has come to divergent conclusions about whether PIP₂-dependent modulation of KCNQ1-containing channels shifts the voltage dependenc of activation, with one group 466 467 positing it does (Loussouarn et al., 2003; Lopes et al., 2005), but another group concluding that it does not 468 (Li et al., 2011; Zaydman et al., 2013). Our data here are in accord with the latter conclusion in the case of 469 KCNQ3 channels, consistent with the conclusions for KCNQ2/3 heteromers (Shapiro et al., 2000; Nakajo 470 and Kubo, 2005; Suh et al., 2006). The presence or absence of KCNE1 subunits is unlikely to alter such 471 conclusions for KCNQ1, since KCNE1 was shown to have no direct impact on VSD activation or pore 472 opening, but rather to affect VSD-PD coupling (Zaydman et al., 2014). Consistent with this, a point 473 mutation (F351A) at the VSD-PD interface had similar effects on KCNQ1 as does inclusion of KCNE1 in 474 the channel. In that work, both KCNE1 and the F351A mutation abolished the "intermediate-open state" of 475 KCNQ1-containing channels, promoting the activated-open states of KCNQ1 by increasing its PIP₂ affinity 476 (Li et al., 2011; Zaydman et al., 2014; Cui, 2016), besides the suppression of inactivation (Hou et al., 2017). 477 We tentatively conclude PIP₂ to not contribute generally to the voltage-dependence of all KCNQ channels, 478 including KCNQ1, as we found for KCNQ3, but more much more likely to the efficiency of VSD-PD 479 coupling. We suspect, but cannot at this point provide evidence, for the underlying reason being the display 480 of two distinct open states of all KCNQ channels (Selyanko and Brown, 1999; Zaydman and Cui, 2014), 481 leading to state transitions, and PIP₂ actions on voltage dependence, differing from those of other Kv 482 channels.

483 Although, we now are in accord with four distinct regions of KCNO1-3 channels interacting with PIP₂, 484 we cannot rule out yet additional PIP₂-binding sites. The distal C-terminus contains basic residues that are 485 conserved in all KCNQ channels, which may also contribute to PIP₂. Our experiments show that the triplet of 486 lysisines (K531, K532 and K533) located at the end of the B-helix of KCNQ3 do not interact with PIP₂. However, R539 and R555 located in the distal C-terminus of KCNQ1 (within the C-helix) were reported to 487 decrease the affinity of the channel to DiC8-PIP₂ (Park et al., 2005), and K526, K527, K528 have been 488 identified as a critical 5th site where CaM competes with PIP₂ to stabilize the open state of KCNQ1-489 containing channels (Tobelaim et at., 2017a, b). The possibility of other PIP₂ interacting sites on the distal 490 491 C-terminus region is intriguing, given the location of the site of phosphorylation of KCNQ3 channels by 492 protein kinase C (PKC) (Hoshi et al., 2003) since such phosphorylation would add a counter-acting negative 493 charge at that locus. This could be a "hot-spot" of PIP₂/PKC cross talk, both of which being affected by 494 stimulation of G_a-coupled receptors. Such a highly-intriguing possibility needs to be carefully examined for all KCNQ2-4 channels, as well as KCNQ2/3 heteromers that underlie most M-type K⁺ currents in the 495 496 nervous system.

497

498 MATERIALS AND METHODS499

500 Cell culture and Transfection

501 Chinese hamster ovary (CHO) cells were grown in 100-mm tissue-culture dishes (Falcon, Franklin 502 Lakes, NJ) in DMEM medium with 10% heat-inactivated fetal bovine serum plus 0.1% 503 penicillin/streptomycin in a humidified incubator at 37° C (5% CO₂) and passaged every 4 days. Cells were 504 discarded after ~30 passages. For patch-clamp and the total internal reflection fluorescent (TIRF) 505 experiments, CHO cells were first passaged onto 35 mm plastic tissue culture dishes and transfected 24h 506 later with FuGENE HD reagent (Promega), according to the manufacturer's instructions. The next day, cells 507 were plated onto cover glass chips, and experiments were performed over the following 1-2 days.

508 Perforated-patch electrophysiology

509 Pipettes were pulled from borosilicate glass capillaries (1B150F-4, World Precision Instruments) using a 510 Flaming/Brown micropipette puller P-97 (Sutter Instruments), and had resistances of 2-4 M Ω when filled 511 with internal solution and measured in standard bath solution. Membrane current was measured with pipette 512 and membrane capacitance cancellation, sampled at 5 ms and filtered at 500 Hz by means of an EPC9 513 amplifier and PULSE software (HEKA/Instrutech). In all experiments, the perforated-patch method of

recording was used with amphotericin B (600 ng/ml) in the pipette (Rae et al., 1991). Amphotericin was prepared as a stock solution as 60 mg/ml in DMSO. In these experiments, the access resistance was typically 7-10 MΩ 5-10 min after seal formation. Cells were placed in a 500 µl perfusion chamber through which solution flowed at 1-2 ml/min. Inflow to the chamber was by gravity from several reservoirs, selectable by activation of solenoid valves (Warner Scientific). Bath solution exchange was essentially complete by <30 s. Experiments were performed at room temperature.

520 Currents were studied by holding the membrane potential at -80 mV, and applying 800 ms 521 depolarizing pulses from 60 mV to -80 mV, every 3 s. KCNQ-current amplitude was measured at 60 mV. 522 To estimate voltage dependence, tail current amplitudes were measured ~ 20 ms after the repolarization at -523 60 mV, normalized, and plotted as a function of test potential. The data were fit with Boltzmann relations of 524 the form: $I/I_{max} = I_{max}/\{1 + \exp[(V_{1/2} - V)/k]\}$, where I_{max} is the maximum tail current, $V_{1/2}$ is the voltage that 525 produces half-maximal activation of the conductance and k is the slope factor. Cell populations were 526 compared using a two tailed t-test. To evaluate the apparent affinity of wild-type and mutant KCNQ3T 527 channels for PIP₂, we used the Dr-VSP cloned into the pIRES-EGFP bicistronic vector, so that transfected 528 cells would express more copies of Dr-VSP than of EGFP. The cells patched were chosen based on their visible EGFP fluorescence as previously described (Falkenburger et al., 2010). Current decay was measured 529 530 at 120 mV, normalized, and plotted as a function of time. Recovery of the current was quantified at 30 mV 531 or 0 mV (which is negative to activation of Dr-VSP) after depolarization to 120 mV or 100 mV. The rate of 532 current recovery was quantified with a single exponential fit as previously described which we realize is an approximation due to the confound of the known rate of PI(4)P-5 kinase ($\tau \sim 10$ s at RT) (Falkenburger et al., 533 534 2010), and the rate of current decay quantified ~ 30 ms after the activation of Dr-VSP at 120 mV with single exponential fits, Finally, the steady-state inhibition of the current by Dr-VSP was quantified by comparing 535 536 current at 30 mV or 0 mV before and after activation of Dr-VSP. Data are given as the mean \pm S.E.M.

The external Ringer's solution used to record KCNQ currents in CHO cells contained (in mM): 160
NaCl, 5 KCl, 2 CaCl₂, 1 MgCl₂ and 10 HEPES, pH 7.4 with NaOH. The pipette solution contained (in mM):
160 KCl, 5 MgCl₂ and 10 HEPES, pH 7.4 with KOH with added amphotericin B (600 ng/ml).

541 Total Internal Reflection Fluorescence (TIRF) microscopy

542 Fluorescence emission from enhanced yellow fluorescent protein (YFP)-tagged KCNQ3T and KCNQ3T 543 mutants (R190Q, R242A, H257N, R364A, KRK-AAA, H367C, Δ linker and RH-AC/ Δ linker) were 544 collected at room temperature using TIRF (also called evanescent field) microscopy. Total internal reflection 545 fluorescence generates evanescent field illumination normal to the interface between two media of differing refractive indices, the cover glass and water in this case, that declines exponentially with distance, 546 547 illuminating only a thin section (300 nm) of the cell very near the cover glass, including the plasma membrane (Axelrod, 2003). All TIRF experiments were performed on a Nikon TE2000 microscope mated to 548 549 a Prairie Technologies laser launch delivery system, as previously described (Bal et al., 2008). Images were not binned or filtered, with a pixel size corresponding to a square of 122×122 nm. The reader should know 550 551 that this system has now been very significantly upgraded.

552

553 Structural homology, simulation and docking models.

554 The human KCNQ3 channel sequence in FASTA format (uniprot ID O43525) was loaded into Swiss-555 PdbViewer 4.10 (Schwede et al., 2003) for template searching against the ExPDB database (ExPASy, 556 http://www.expasy.org/). Then, the structural model for the full length of the *Rattus norvegicus* voltage-557 gated K⁺ channel subfamily A member 2 (Kv1.2; PDB: 3LUT) (Chen et al., 2010) was identified as the best 558 template. The initial sequence alignments between the KCNO3 channel and Kv1.2 were generated with fulllength pairwise alignments using ClustalW (Thompson et al., 1994). Sequence alignments were inspected 559 560 manually to assure accuracy among structural domains solved from the template. Since the turret domain of the KCNO3 subunit was absent in the solved Kv1.2 structure, residues 287-296 were excluded from the 561 562 modeling. The A315T pore mutation was also omitted from the template as, it does not change the apparent 563 PIP₂ affinity of the channel (Hernandez et al., 2009). Full-length multiple alignments were submitted for 564 automated comparative protein modelling implemented in the program suite incorporated in SWISS-MODEL (http://swissmodel.expasy.org/SWISS-MODEL.html). 565

566 Before energy minimization using GROMOS96 (Schuler et al., 2001), the resulting structural models of 567 KCNQ3 subunits were manually inspected, the structural alignments confirmed and evaluated for proper H-

13

bonds and the presence of clashes and missing atoms using estimated using Molegro Molecular Viewer 568 569 (www.clcbio.com). Further structural models were generated by rearrangement of four KCNQ3 subunit models as a tetramer. Coordinates of the Kv1.2 channel in the resting/closed and activated/open states 570 571 (Khalili-Araghi et al., 2010) were used to model the KCNQ3 channel in both forms. The calculated energies 572 for the corresponding KCNQ3 open and closed stated structural models were highly favourable (-35,580 573 KJ/mol and -27,656 KJ/mol, respectively). Neighbourhood structural conformational changes caused by the introduction of single point mutations of the KCNO3 structure were simulated using Rosetta 3.1 (Smith and 574 575 Kortemme, 2008), and implemented in the program suite incorporated in Rosetta Backrub 576 (https://kortemmelab.ucsf.edu). As Rosetta 3.1 does not allow cysteine substitutions, KCNQ3 subunits (WT 577 or mutant) with cysteines were exchanged for alanines. Simulations for single point mutations were carried 578 out for dimers, for which identical mutations were presented in neighboring subunits, excluding distal 579 residues of the C-terminus (404-557).

580 Up to twenty of the best-scoring structures were generated at each time by choosing parameters 581 recommended by the application. The root-mean square (RMS) deviation was calculated between the WT 582 structures and superimposed on the simulated mutant structures. For each mutation, the RMS average over 583 ten low energy structures was computed and conformational changes displayed among neighboring structural domains considered significant for values of RMS > 0.5 Å. PatchDock (Schneidman-Duhovny et 584 585 al., 2005), a molecular docking algorithm based on shape complementarity principles, was used to dock PIP₂ 586 with proposed interacting domains at the interfaces of dimers homology models based upon the Kv1.2 structure. One PIP₂ ligand was simulated docked per subunit, with the structure of PIP₂ used as in the solved 587 588 PIP₂-bound structure of Kv2.2 (Hansen et al., 2011). PatchDock was implemented using an algorithm 589 applied for protein-small ligand docking with a default clustering of 1.5 Å of the RMS as recommended. Before the simulation, a list of residues for three predicted binding sites for PIP₂ in the docking site was 590 derived, as indicated by functional studies, which included domains within the S2-S3 and S4-S5 linkers, and 591 592 the proximal C-terminus. Twenty solutions for the first and the fifth best-scoring simulated mutant were 593 ranked according to the geometric shape complementarity score and the atomic contact energy (-171 594 kcal/mol and -243 kcal/mol for the open and closed states, respectively) (Zhang et al., 1997), and inspected 595 manually to assure accuracy among representative orientations of bound PIP₂. The energy electrostatic 596 interactions for a given docking pose (ligand-protein complex) were analysed using the ligand energy 597 inspector implemented through the Molegro Molecular viewer. The short-range electrostatic interactions (r 598 < 6 Å) between the PIP₂ and residues in WT or mutant were computed and the lowest solutions among the highest geometric score and the right orientation represented here. We prepared the modeling figures using 599 600 Chimera 1.7 (Pettersen et al., 2004).

602 ACKNOWLEDGMENTS

601

603 All authors declare that there is no conflict of interest. We gratefully acknowledge the assistance of Pamela

Reed, Maryann Hobbs and Isamar Sanchez in this work. We also thank Nikita Gamper and Crystal Archer for useful discussions. This work was supported by National Institutes of Health grants R01 NS150305,

606 R01 NS094461and R56 NS153503 to M.S.S.

14

607 **R**EFERENCES

- Abderemane-Ali, F., Z. Es-Salah-Lamoureux, L. Delemotte, M.A. Kasimova, A.J. Labro, D.J. Snyders, D.
 Fedida, M. Tarek, I. Baro, and G. Loussouarn. 2012. Dual effect of phosphatidylinositol (4,5)bisphosphate PIP₂ on Shaker K⁺ channels. *Journal of Biological Chemistry*. 287:36158-36167.
- Aivar, P., J. Fernandez-Orth, C. Gomis-Perez, A. Alberdi, A. Alaimo, M.S. Rodriguez, T. Giraldez, P.
 Miranda, P. Areso, and A. Villarroel. 2012. Surface expression and subunit specific control of steady
 protein levels by the Kv7.2 helix A-B linker. *PloS One*. 7:e47263.
- Axelrod, D. 2003. Total internal reflection fluorescence microscopy in cell biology. *Methods in Enzymology*.
 361:1-33.
- Bal, M., J. Zhang, O. Zaika, C.C. Hernandez, and M.S. Shapiro. 2008. Homomeric and heteromeric assembly of KCNQ (Kv7) K⁺ channels assayed by total internal reflection fluorescence/fluorescence
 resonance energy transfer and patch clamp analysis. *Journal of Biological Chemistry*. 283:30668-30676.
- Bender, K., M.C. Wellner-Kienitz, and L. Pott. 2002. Transfection of a phosphatidyl-4-phosphate 5-kinase
 gene into rat atrial myocytes removes inhibition of GIRK current by endothelin and alpha-adrenergic
 agonists. *FEBS letters*. 529:356-360.
- Boyer, S.B., and P.A. Slesinger. 2010. Probing novel GPCR interactions using a combination of FRET and
 TIRF. *Communicative & Integrative Biology*. 3:343-346.
- 625 Chambard, J.M., and J.F. Ashmore. 2005. Regulation of the voltage-gated potassium channel KCNQ4 in the
 626 auditory pathway. *Pflugers Archiv*. 450:34-44.
- Chen, X., Q. Wang, F. Ni, and J. Ma. 2010. Structure of the full-length Shaker potassium channel Kv1.2 by
 normal-mode-based X-ray crystallographic refinement. *Proceedings of the National Academy of Sciences.* 107:11352-11357.
- Choveau, F.S., C.C. Hernandez, S.M. Bierbower, and M.S. Shapiro. 2012. Pore determinants of KCNQ3 K⁺
 current expression. *Biophysical Journal*. 102:2489-2498.
- Choveau, F.S., N. Rodriguez, F. Abderemane Ali, A.J. Labro, T. Rose, S. Dahimene, H. Boudin, C. Le
 Henaff, D. Escande, D.J. Snyders, F. Charpentier, J. Merot, I. Baro, and G. Loussouarn. 2011.
 KCNQ1 channels voltage dependence through a voltage-dependent binding of the S4-S5 linker to the
 pore domain. *Journal of Biological Chemistry*. 286:707-716.
- 636 Cui, J. 2016. Voltage-Dependent Gating: Novel Insights from KCNQ1 Channels. *Biophysical Journal*.
 637 110:14-25.
- Eckey, K., E. Wrobel, N. Strutz-Seebohm, L. Pott, N. Schmitt, and G. Seebohm. 2014. Novel Kv7.1 phosphatidylinositol 4,5-bisphosphate interaction sites uncovered by charge neutralization scanning.
 Journal of Biological Chemistry. 289:22749-22758.
- Falkenburger, B.H., J.B. Jensen, and B. Hille. 2010. Kinetics of PIP₂ metabolism and KCNQ2/3 channel
 regulation studied with a voltage-sensitive phosphatase in living cells. *Journal of General Physiology*. 135:99-114.
- Gamper, N., Y. Li, and M.S. Shapiro. 2005. Structural requirements for differential sensitivity of KCNQ K⁺
 channels to modulation by Ca²⁺/calmodulin. *Molecular Biology of the Cell*. 16:3538-3551.
- Gamper, N., and M.S. Shapiro. 2003. Calmodulin mediates Ca2+-dependent modulation of M-type K⁺
 channels. *Journal of General Physiology*. 122:17-31.
- 648 Gamper, N., and M.S. Shapiro. 2007. Regulation of ion transport proteins by membrane phosphoinositides.
 649 *Nature Reviews Neuroscience*. 8:921-934.
- Gamper, N., O. Zaika, Y. Li, P. Martin, C.C. Hernandez, M.R. Perez, A.Y. Wang, D.B. Jaffe, and M.S.
 Shapiro. 2006. Oxidative modification of M-type K⁺ channels as a mechanism of cytoprotective neuronal silencing. *EMBO journal*. 25:4996-5004.
- Hansen, S.B., X. Tao, and R. MacKinnon. 2011. Structural basis of PIP₂ activation of the classical inward
 rectifier K⁺ channel Kir2.2. *Nature*. 477:495-498.
- Hernandez, C.C., B. Falkenburger, and M.S. Shapiro. 2009. Affinity for phosphatidylinositol 4,5bisphosphate determines muscarinic agonist sensitivity of Kv7 K⁺ channels. *Journal of General Physiology*. 134:437-448.

- Hernandez, C.C., O. Zaika, and M.S. Shapiro. 2008. A carboxy-terminal inter-helix linker as the site of
 phosphatidylinositol 4,5-bisphosphate action on Kv7 (M-type) K⁺ channels. *Journal of General Physiology*. 132:361-381.
- Hoshi, N., J.S. Zhang, M. Omaki, T. Takeuchi, S. Yokoyama, N. Wanaverbecq, L.K. Langeberg, Y. Yoneda,
 J.D. Scott, D.A. Brown, and H. Higashida. 2003. AKAP150 signaling complex promotes suppression
 of the M-current by muscarinic agonists. *Nature Neuroscience*. 6:564-571.
- Hou, P., J. Eldstrom, J. Shi, L. Zhong, K. McFarland, Y. Gao, D. Fedida, and J. Cui. 2017. Inactivation of
 KCNQ1 potassium channels reveals dynamic coupling between voltage sensing and pore opening.
 Nature Communications. 8:1730.
- Itsuki, K., Y. Imai, Y. Okamura, K. Abe, R. Inoue, and M.X. Mori. 2012. Voltage-sensing phosphatase
 reveals temporal regulation of TRPC3/C6/C7 channels by membrane phosphoinositides. *Channels*.
 669 6:206-209.
- Jentsch, T.J. 2000. Neuronal KCNQ potassium channels: physiology and role in disease. *Nature Reviews Neuroscience*. 1:21-30.
- Kasimova, M.A., M.A. Zaydman, J. Cui, and M. Tarek. 2015. PIP₂-dependent coupling is prominent in
 Kv7.1 due to weakened interactions between S4-S5 and S6. *Scientific Reports*. 5:7474.
- Khalili-Araghi, F., V. Jogini, V. Yarov-Yarovoy, E. Tajkhorshid, B. Roux, and K. Schulten. 2010.
 Calculation of the gating charge for the Kv1.2 voltage-activated potassium channel. *Biophysical journal*. 98:2189-2198.
- Kosenko, A., and N. Hoshi. 2013. A change in configuration of the calmodulin-KCNQ channel complex
 underlies Ca²⁺-dependent modulation of KCNQ channel activity. *PloS one*. 8:e82290.
- Kruse, M., G.R. Hammond, and B. Hille. 2012. Regulation of voltage-gated potassium channels by PI(4,5)P2. *Journal of general physiology*. 140:189-205.
- Labro, A.J., I.R. Boulet, F.S. Choveau, E. Mayeur, T. Bruyns, G. Loussouarn, A.L. Raes, and D.J. Snyders.
 2011. The S4-S5 linker of KCNQ1 channels forms a structural scaffold with the S6 segment
 controlling gate closure. *Journal of Biological Chemistry*. 286:717-725.
- Li, Y., N. Gamper, D.W. Hilgemann, and M.S. Shapiro. 2005. Regulation of Kv7 (KCNQ) K⁺ channel open probability by phosphatidylinositol 4,5-bisphosphate. *Journal of Neuroscience*. 25:9825-9835.
- Li, Y., M.A. Zaydman, D. Wu, J. Shi, M. Guan, B. Virgin-Downey, and J. Cui. 2011. KCNE1 enhances
 phosphatidylinositol 4,5-bisphosphate (PIP₂) sensitivity of IKs to modulate channel activity.
 Proceedings of the National Academy of Sciences. 108:9095-9100.
- Logothetis, D.E., T. Jin, D. Lupyan, and A. Rosenhouse-Dantsker. 2007. Phosphoinositide-mediated gating
 of inwardly rectifying K⁺ channels. *Pflugers Archiv : European Journal of Physiology*. 455:83-95.
- Long, S.B., E.B. Campbell, and R. Mackinnon. 2005. Voltage sensor of Kv1.2: structural basis of
 electromechanical coupling. *Science*. 309:903-908.
- Long, S.B., X. Tao, E.B. Campbell, and R. MacKinnon. 2007. Atomic structure of a voltage-dependent K⁺
 channel in a lipid membrane-like environment. *Nature*. 450:376-382.
- Lopes, C.M., T. Rohacs, G. Czirjak, T. Balla, P. Enyedi, and D.E. Logothetis. 2005. PIP₂ hydrolysis
 underlies agonist-induced inhibition and regulates voltage gating of two-pore domain K⁺ channels.
 Journal of Physiology. 564:117-129.
- Loussouarn, G., K.H. Park, C. Bellocq, I. Baro, F. Charpentier, and D. Escande. 2003. Phosphatidylinositol 4,5-bisphosphate, PIP₂, controls KCNQ1/KCNE1 voltage-gated potassium channels: a functional
 homology between voltage-gated and inward rectifier K⁺ channels. *EMBO Journal*. 22:5412-5421.
- Murata, Y., and Y. Okamura. 2007. Depolarization activates the phosphoinositide phosphatase Ci-VSP, as
 detected in Xenopus oocytes coexpressing sensors of PIP₂. *Journal of Physiology*. 583:875-889.
- Nakajo, K., and Y. Kubo. 2005. Protein kinase C shifts the voltage dependence of KCNQ/M channels
 expressed in Xenopus oocytes. *Journal of Physiology*. 569:59-74.
- Park, K.H., J. Piron, S. Dahimene, J. Merot, I. Baro, D. Escande, and G. Loussouarn. 2005. Impaired
 KCNQ1-KCNE1 and phosphatidylinositol-4,5-bisphosphate interaction underlies the long QT
 syndrome. *Circulation Research*. 96:730-739.
- Peroz, D., N. Rodriguez, F. Choveau, I. Baro, J. Merot, and G. Loussouarn. 2008. Kv7.1 (KCNQ1)
 properties and channelopathies. *Journal ofPphysiology*. 586:1785-1789.

- Pettersen, E.F., T.D. Goddard, C.C. Huang, G.S. Couch, D.M. Greenblatt, E.C. Meng, and T.E. Ferrin. 2004.
 UCSF Chimera--a visualization system for exploratory research and analysis. *Journal of Computational Chemistry*. 25:1605-1612.
- Rae, J., K. Cooper, P. Gates, and M. Watsky. 1991. Low access resistance perforated patch recordings using
 amphotericin B. *Journal of Neuroscience Methods*. 37:15-26.
- Rodriguez-Menchaca, A.A., S.K. Adney, Q.Y. Tang, X.Y. Meng, A. Rosenhouse-Dantsker, M. Cui, and
 D.E. Logothetis. 2012. PIP₂ controls voltage-sensor movement and pore opening of Kv channels
 through the S4-S5 linker. *Proceedings of the National Academy of Sciences*. 109:E2399-2408.
- Sachyani, D., M. Dvir, R. Strulovich, G. Tria, W. Tobelaim, A. Peretz, O. Pongs, D. Svergun, B. Attali, and
 J.A. Hirsch. 2014. Structural basis of a Kv7.1 potassium channel gating module: studies of the
 intracellular c-terminal domain in complex with calmodulin. *Structure*. 22:1582-1594.
- Schneidman-Duhovny, D., Y. Inbar, R. Nussinov, and H.J. Wolfson. 2005. PatchDock and SymmDock:
 servers for rigid and symmetric docking. *Nucleic Acids Research*. 33:W363-367.
- Schuler, L.D., X. Daura, and W.F. Van Gunsteren. 2001. An improved GROMOS96 force field for aliphatic
 hydrocarbons in the condensed phase. *Journal of computational chemistry*. 22:1205-1218.
- Schwede, T., J. Kopp, N. Guex, and M.C. Peitsch. 2003. SWISS-MODEL: An automated protein homology modeling server. *Nucleic Acids Research*. 31:3381-3385.
- 727 Selyanko, A.A., and D.A. Brown. 1999. M-channel gating and simulation. *Biophysical journal*. 77:701-713.
- Shamgar, L., L. Ma, N. Schmitt, Y. Haitin, A. Peretz, R. Wiener, J. Hirsch, O. Pongs, and B. Attali. 2006.
 Calmodulin is essential for cardiac IKS channel gating and assembly: impaired function in long-QT
 mutations. *Circulation Research*. 98:1055-1063.
- Shapiro, M.S., J.P. Roche, E.J. Kaftan, H. Cruzblanca, K. Mackie, and B. Hille. 2000. Reconstitution of muscarinic modulation of the KCNQ2/KCNQ3 K⁺ channels that underlie the neuronal M current. *Journal of Neuroscience*. 20:1710-1721.
- Smith, C.A., and T. Kortemme. 2008. Backrub-like backbone simulation recapitulates natural protein
 conformational variability and improves mutant side-chain prediction. *Journal of Molecular Biology*.
 380:742-756.
- Suh, B.C., and B. Hille. 2002. Recovery from muscarinic modulation of M current channels requires
 phosphatidylinositol 4,5-bisphosphate synthesis. *Neuron*. 35:507-520.
- Suh, B.C., T. Inoue, T. Meyer, and B. Hille. 2006. Rapid chemically induced changes of PtdIns(4,5)P2 gate
 KCNQ ion channels. *Science*. 314:1454-1457.
- Telezhkin, V., J.M. Reilly, A.M. Thomas, A. Tinker, and D.A. Brown. 2012. Structural requirements of
 membrane phospholipids for M-type potassium channel activation and binding. *Journal of Biological Chemistry*. 287:10001-10012.
- Telezhkin, V., A.M. Thomas, S.C. Harmer, A. Tinker, and D.A. Brown. 2013. A basic residue in the
 proximal C-terminus is necessary for efficient activation of the M-channel subunit Kv7.2 by
 PI(4,5)P(2). *Pflugers Archiv*. 465:945-953.
- Thomas, A.M., S.C. Harmer, T. Khambra, and A. Tinker. 2011. Characterization of a binding site for
 anionic phospholipids on KCNQ1. *Journal of Biological Chemistry*. 286:2088-2100.
- Thompson, J.D., D.G. Higgins, and T.J. Gibson. 1994. CLUSTAL W: improving the sensitivity of
 progressive multiple sequence alignment through sequence weighting, position-specific gap penalties
 and weight matrix choice. *Nucleic Acids Research*. 22:4673-4680.
- Tobelaim, W.S., M. Dvir, G. Lebel, M. Cui, T. Buki, A. Peretz, M. Marom, Y. Haitin, D.E. Logothetis, J.A.
 Hirsch, and B. Attali. 2017a. Ca²⁺-calmodulin and PIP₂ interactions at the proximal C-terminus of Kv7 channels. 11:686-695.
- Tobelaim, W.S., M. Dvir, G. Lebel, M. Cui, T. Buki, A. Peretz, M. Marom, Y. Haitin, D.E. Logothetis, J.A.
 Hirsch, and B. Attali. 2017b. Competition of calcified calmodulin N lobe and PIP₂ to an LQT
 mutation site in Kv7.1 channel. *Proceedings of the National Academy of Sciences*. 114:E869-e878.
- Whorton, M.R., and R. MacKinnon. 2011. Crystal structure of the mammalian GIRK2 K⁺ channel and gating regulation by G proteins, PIP₂, and sodium. *Cell*. 147:199-208.
- Winks, J.S., S. Hughes, A.K. Filippov, L. Tatulian, F.C. Abogadie, D.A. Brown, and S.J. Marsh. 2005.
 Relationship between membrane phosphatidylinositol-4,5-bisphosphate and receptor-mediated
 inhibition of native neuronal M channels. *Journal of Neuroscience*. 25:3400-3413.

- Zaika, O., C.C. Hernandez, M. Bal, G.P. Tolstykh, and M.S. Shapiro. 2008. Determinants within the turret and pore-loop domains of KCNQ3 K⁺ channels governing functional activity. *Biophysical journal*. 95:5121-5137.
- Zaika, O., G.P. Tolstykh, D.B. Jaffe, and M.S. Shapiro. 2007. Inositol triphosphate-mediated Ca²⁺ signals
 direct purinergic P2Y receptor regulation of neuronal ion channels. *Journal of Neuroscience*.
 27:8914-8926.
- Zaydman, M.A., and J. Cui. 2014. PIP₂ regulation of KCNQ channels: biophysical and molecular
 mechanisms for lipid modulation of voltage-dependent gating. *Frontiers in Physiology*. 5:195.
- Zaydman, M.A., M.A. Kasimova, K. McFarland, Z. Beller, P. Hou, H.E. Kinser, H. Liang, G. Zhang, J. Shi,
 M. Tarek, and J. Cui. 2014. Domain-domain interactions determine the gating, permeation,
 pharmacology, and subunit modulation of the IKs ion channel. *eLife*. 3:e03606.
- Zaydman, M.A., J.R. Silva, K. Delaloye, Y. Li, H. Liang, H.P. Larsson, J. Shi, and J. Cui. 2013. Kv7.1 ion
 channels require a lipid to couple voltage sensing to pore opening. *Proceedings of the National Academy of Sciences*. 110:13180-13185.
- Zhang, C., G. Vasmatzis, J.L. Cornette, and C. DeLisi. 1997. Determination of atomic desolvation energies
 from the structures of crystallized proteins. *Journal of Molecular Biology*. 267:707-726.
- Zhang, H., L.C. Craciun, T. Mirshahi, T. Rohacs, C.M. Lopes, T. Jin, and D.E. Logothetis. 2003. PIP₂
 activates KCNQ channels, and its hydrolysis underlies receptor-mediated inhibition of M currents.
 Neuron. 37:963-975.
- Zhang, J., M. Bal, S. Bierbower, O. Zaika, and M.S. Shapiro. 2011. AKAP79/150 signal complexes in G protein modulation of neuronal ion channels. *Journal of Neuroscience*. 31:7199-7211.
- Zhang, Q., P. Zhou, Z. Chen, M. Li, H. Jiang, Z. Gao, and H. Yang. 2013. Dynamic PIP₂ interactions with
 voltage sensor elements contribute to KCNQ2 channel gating. *Proceedings of the National Academy* of Sciences. 110:20093-20098.
- Zhou, P., H. Yu, M. Gu, F.J. Nan, Z. Gao, and M. Li. 2013. Phosphatidylinositol 4,5-bisphosphate alters
 pharmacological selectivity for epilepsy-causing KCNQ potassium channels. *Proceedings of the National Academy of Science*. 110:8726-8731.

790 FIGURE LEGENDS

791 Figure 1. Location of the site(s) of PIP₂ action on KCNQ3 channels.

792 (A) Sequence alignments of human KCNQ channels of the putative PIP₂ interaction-domains studied in this 793 work. The residues highlighted in red are conserved basic residues across all KCNQ channels. Structural 794 domains where the putative PIP₂-interacting residues are located are indicated below the alignments as solid 795 lines (α -helices) and non-continuous lines (linkers). (B, C) 3D structural models of the open conformation of 796 the KCNO3 channel in ribbon representation, coloured by subunits as viewed from the membrane plane (B) and the intracellular side (C). Conserved basic residues R190, R195, R242, H257, K358, and R364 tested in 797 798 this study by mutagenesis are shown in gray and mapped onto the channel. (D) Ribbon representations of the arrangement of the VSD-PD interface of a structural subunit model viewed from the outer and inner side 799 (upper panels), and membrane plane (bottom panels). The secondary structure of the channels is colored 800 according to structural domain, as indicated. Side chains of basic residues involved in PIP₂ interactions are 801 802 in color, according to structural domain (grey for the S2-S3 linker and S6Jx, and purple for the S4-S5 linker). 803 The PIP₂ molecule is shown in molecular surface representation within the docking cavity. (E) Expanded 804 view of the most favourable binding model of PIP_2 in the open conformation. Panels show two neighboring 805 subunits (Sub) forming the VSD-PD interface (Sub-C and Sub-D). The docking site enclosed in a red box 806 was enlarged for clarity. In stick representation are the residues forming hydrogen bonds and electrostatic 807 interactions within the interaction site. Residues in blue from the Sub-D enclose the phosphate groups of PIP₂, and residues in orange from the Sub-C enclose the acyl tail of the PIP₂ between sub-C and Sub-D at 808 the S6Jx. The following are the favorable interactions (label in red) predicted to be in the PIP2-docking-809 network (< 6.0 Å, kJ/M): R242 = -12.26, R243 = -4.60, H257 = -1.10, K358 = -4.28, K366 = -5.74. 810 811 Hydrogen bonds are not shown.

812

Figure 2. Effects of charge neutralizing mutations located in the S2-S3 and S4-S5 linkers on KCNQ3T channels.

815 (A) Representative perforated patch-clamp recordings from CHO cells transfected with KCNQ3T or the indicated mutant channels. (B) Bars show summarized current densities at 60 mV for the indicated channels 816 (n = 6-19). (C) Voltage dependence of activation of the tail currents at -60 mV, plotted as a function of test 817 818 potential (n = 5-19). (D) Representative perforated patch-clamp recordings from CHO cells co-transfected 819 with Dr-VSP and KCNO3T or the indicated mutant channels. (E) Bars summarize time constant values from single exponential fits to current decay during Dr-VSP activation (n = 5-10;). (F) Bars summarize time 820 821 constants of single exponential fits to current recovery after Dr-VSP turn-off (n = 5-11). (G) Bars 822 Summarize fractional inhibition after M_1R stimulation for the indicated mutant channels (n = 3-7). * p < 0.05; ** p < 0.01; ***p < 0.001. 823

824

825 Figure 3. Effects of charge neutralizing mutations located in the S6Jx on KCNQ3T channels.

826 (A) Representative perforated patch-clamp recordings from KCNQ3T and mutant channels. (B) Bars show 827 summarized current densities at 60 mV for the indicated channels (n = 6-19). (C) Voltage dependence of 828 activation of the tail currents at -60 mV, plotted as a function of test potential (n = 6-19). (D) Representative 829 perforated patch-clamp recordings from CHO cells co-transfected with Dr-VSP and KCNQ3T or mutant 830 KCNQ3T channels. (E) Bars summarize time constants from single-exponential fits to current decay during 831 Dr-VSP activation (n = 5-10). (F) Bars summarize time constants from single-exponential fits to recovery 832 after Dr-VSP turn-off (n = 5-11). **p < 0.01; *** p < 0.001.

833

834 Figure 4. Effects of the A-B linker deletion on KCNQ3T channels.

835 (A) Representative perforated patch-clamp recordings from cells expressing Dr-VSP and either KCNQ3T, 836 KCNQ3T (Δ linker) or KCNQ3T (RH-AC/ Δ linker). Cells were held at -80 mV and voltage steps were 837 applied from -80 to 60 mV in 10 mV increments every 3s. (B) Bars show summarized current densities at 60 838 mV for the indicated channels (n = 8-19). (C) Shown are the amplitude of tail currents at -60 mV, plotted as 839 a function of test potential from KCNQ3T and KCNQ3T (Δ linker) channels (n = 11-19). (D) Representative 840 perforated patch-clamp recordings from CHO cells co-transfected with Dr-VSP and KCNQ3T or KCNQ3T

841 (Δ linker) or the RH-AC/ Δ linker mutants. (E) Bars summarize time constants from single-exponential fits to

842 current decay during Dr-VSP activation (n = 6-10). (F) Bars summarize time constants from single-843 exponential fits to recovery after Dr-VSP turn-off (n = 6-11). * p < 0.05, *** p < 0.001.

845 Figure 5. TIRF microscopy indicates that mutants in PIP₂-interacting domains result in minor 846 differences in membrane expression of channels.

847 (A) Shown are fluorescent images under TIRF microscopy of CHO cells expressing the indicated YFP-

tagged channels. (B) Bars show summarized emission intensity data for each channel type (n = 32-60).

849

844

Figure 6. Charge neutralizing mutations at the S2-S3 and S4-S5 linkers and S6Jx disrupt PIP₂ interactions of the KCNQ3 channel.

852 Shown are 3D structural models of the most favourable docking PIP₂-docking conformation of the KCNO3 channel after simulation of charge neutralization at the putative PIP₂ binding site residues R190 in the S2-S3 853 linker (A), R242 and H257 in the S4-S5 linker (B and C), and R364, K358-R364-K366 (KRK-AAA) and 854 855 H367 within the S6Jx (D, E, F). As indicated in Fig. 1, binding sites are enclosed in red boxs and enlarged 856 for clarity in the right panels. Upper panels show two neighboring subunits (Sub-C and Sub-D) or a single 857 Sub forming the binding site. The following are the favorable interactions (labeled in red) predicted to be in 858 the PIP₂-docking-network (< 6.0 Å, kJ/M): R190A in panel A, R242 = -4.80, R243 = -2.40, K358 = -4.31; 859 R242A in panel B, R243 = -24.9, H363 = -4.07, K366 = -7.63; H257N in panel C, R242 = -18.8, R243 = -5.41, K358 = -3.11, K366 = -7.52; R364A in panel D, R242 = -12.80, R243 = -4.54, K358 = -3.20; KRK-860 AAA in panel E, R242 = -27.20, R243 = -3.73, H257 = -1.23; H367A in panel F, R242 = -10.3, R243 = -861 862 3.53, K358 = -5.08; K358A in panel G, R242 = -7.43, R243 = -2.57, H257 = -5.42; K366A in panel H, R242 863 = -4.62, R243 = -2.03, H257 = -4.34, K358 = -2.78.

864

865 Figure 7. Effects of charge neutralization of residues predicted within the PIP2 docking site of KCNQ3 in the closed state. (A) Sequence alignments of human KCNO channels show the additional basic residues 866 867 K103, R188, R227, and R230 tested in this study by mutagenesis. Predicted secondary structure of the 868 channel is indicated above the alignments as solid lines (α -helices) and non-continuous lines (linkers). (B) Ribbon representations of the arrangement of the VSD-PD interface of a structural subunit model 869 viewed from the outer and inner side (upper panels), and membrane plane (bottom panels). The 870 871 secondary structure of the channels and PIP2 molecules are shown as in Fig. 1. (C) Expanded view of 872 the most favourable interaction predicted of PIP2 in the closed-channel state. The phosphate group of the 873 PIP2 is oriented toward the S2-S3 linker, whereas the acyl tail is enclosed within the a-helices. The 874 following are the favorable interactions (label in red) predicted to be in the PIP2-docking-network (< 6.0Å): K103 = -4.03, R188 = -1.44, R190 = -1.52, R227 = -3.23, R230 = -5.36. (D) Top, Representative 875 876 perforated patch-clamp recordings from CHO cells co-transfected with Dr-VSP and KCNO3T or the 877 indicated mutants. Cells were held at -60 mV and current decay measured at 100 mV, and recovery of the 878 current measured at 0 mV after the depolarization to 100 mV. Note the larger amplitude of the recovery 879 current in these experiments after turn-off of Dr-VSP, due to the voltage used (0 mV), at which the "leak" 880 current is expected to minimal, compared to +30 mV Bottom, Bars summarize the data from these

881 *experiments (n = 5-11).* * p < 0.05 and ** p < 0.01.

882

TABLE 1

Effects of mutations on the properties and PIP₂-apparent affinities of KCNQ3 channels

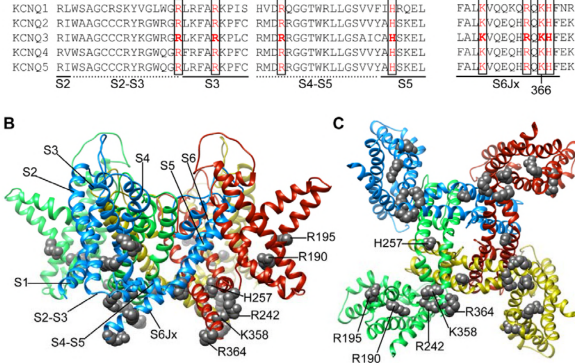
	Structural domain	Channel 1	Function	Rates from D	r-VSP assays
KCNQ3T		Current Density (pA/pF)	V _{1/2} (mV)	τ, decay at +120 mV (s)	τ, recovery at +30 mV (s)
WT	-	197 ± 6	-34.0 ± 1.9	0.94 ± 0.13	7.5 ± 1.7
W 1		(n = 19)	(n = 19)	(n = 10)	(n = 11)
		$66 \pm 12^{***}$	$\textbf{-39.9} \pm 1.9$	0.35 ± 0.11 **	9.6 ± 1.6
R190Q	S2-S3 linker	(n = 6)	(n = 5)	(n = 8)	(n = 8)
R195Q	S2-S3 linker	195 ± 13	-36.0 ± 1.8	0.98 ± 0.22	7.4 ± 1.0
		(n = 6)	(n = 6)	(n = 5)	(n = 5)
R242A	S4-S5 linker	$146 \pm 16^{***}$	-4.0 ± 3.2 ***	0.77 ± 0.13	$14.2 \pm 1.7*$
		(n = 10)	(n = 7)	(n = 10)	(n = 10)
R243A	S4-S5 linker	$56 \pm 16^{***}$	-31 ± 4.7	$0.59\pm0.19*$	13.1 ± 1.8*
		(n = 4)	(n = 4)	(n = 4)	(n = 4)
R242A-R243A	S4-S5 linker	65 ± 11***	-0.2 ± 2.9 ***	0.45 ± 0.11 **	15 ± 1.7*
		(n = 11)	(n = 5)	(n = 5)	(n = 5)
H257N		30 ± 3***	2.5 ± 2.8 ***	$0.58 \pm 0.14*$	11.03 ± 1.4
	S4-S5 linker	(n = 7)	(n = 7)	(n = 4)	(n = 2)
K358A		195 ± 7	-28.8 ± 2.2	0.94 ± 0.25	8.0 ± 1.7
	S6Jx	(n = 9)	(n = 9)	(n = 6)	(n = 6)
R364A		$72 \pm 8***$	-30.1 ± 2	0.14 ± 0.02 ***	27.7 ± 6.9 ***
	S6Jx	(n = 6)	(n = 6)	(n = 5)	(n = 5)
		187 ± 12	-29.6 ± 1.3	0.95 ± 0.12	9.2 ± 1.7
K366A	S6Jx	(n = 8)	(n = 8)	(n = 7)	(n = 6)
KRK-AAA		79 ± 11 ***	-6.3 ± 2.5 ***	0.29 ± 0.04 ***	17.9 ± 2.6 ***
	S6Jx	(n = 8)	(n = 7)	(n = 7)	(n = 6)
H367C	S6Jx	$138 \pm 5^{**}$	$-25.0 \pm 2.1*$	$0.32\pm0.05\textit{**}$	$36.7 \pm 6.9 ***$
115070	SUIX	(n = 6)	(n = 6)	(n = 6)	(n = 6)
$(\Delta \text{ linker})$	C-term	$112 \pm 10^{***}$ (n = 11)	-32.5 ± 1.5 (n = 11)	$0.26 \pm 0.04^{***}$ (n = 7)	$13.5 \pm 2.2*$ (n = 7)
RH-AC/∆ linker	C-term	$16 \pm 2^{***}$ (n = 8)	ND	$0.53 \pm 0.1*$ (n = 6)	$45.8 \pm 5.2^{***}$
KH-AC/A liliker		(11 - 8)		(11 - 0)	(n = 6)
K531N	C-term	193 ± 54	-20.3 ± 6	1.05 ± 0.33	10 ± 0.9
KJJIN V		(n = 8)	(n = 4)	(n = 7)	(n = 7)
K532N	C-term	204 ± 48	-19.9 ± 2.6	1.01 ± 0.32	9.08 ± 1.4
		(n = 8)	(n = 6)	(n = 8)	(n = 8)
K533N	C-term	208 ± 62	-21.1 ± 3.4	0.86 ± 0.12	8.62 ± 0.6
	2	(n = 7)	(n = 7)	(n = 8)	(n = 8)

883 Values represent mean \pm S.E.M. *, ** and *** *P* < 0.05, *P* < 0.01 and *P* < 0.001 (one-way ANOVA with 884 Dunnett's multiple comparisons test) statistically different from wild type (WT). N.D. = not determined.

257

885 Figure 1

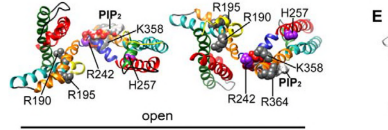
Α



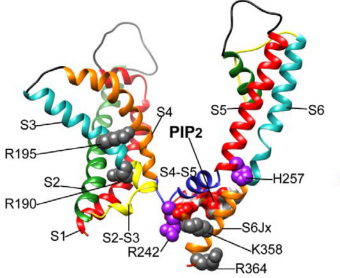
242

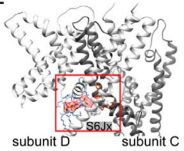
D outer side

inner side



190





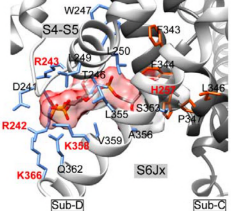
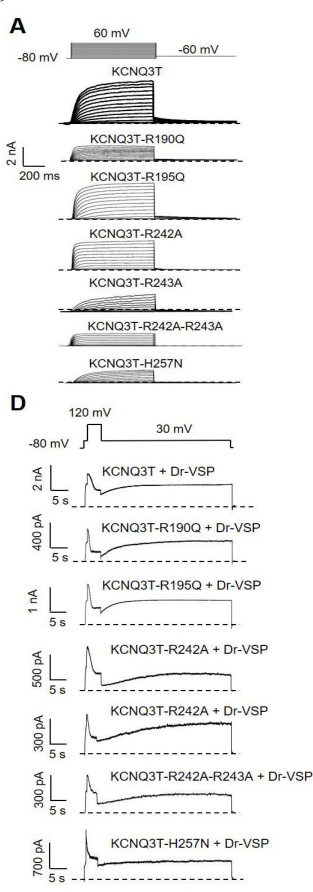
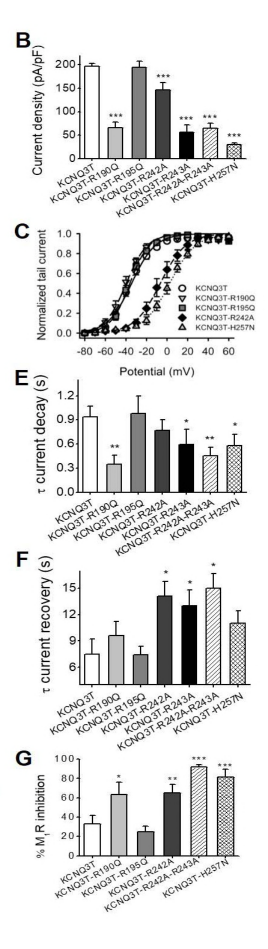
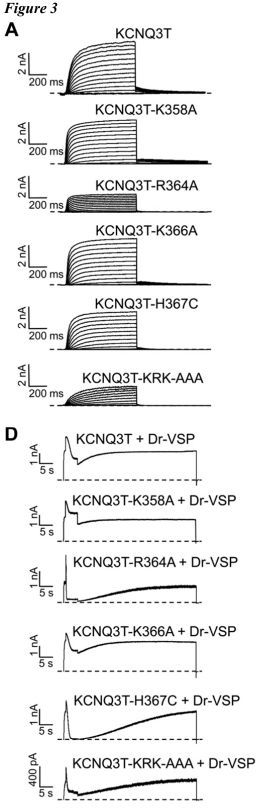


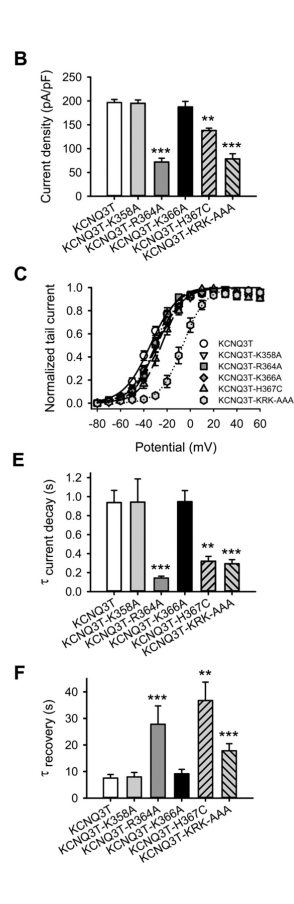
Figure 2



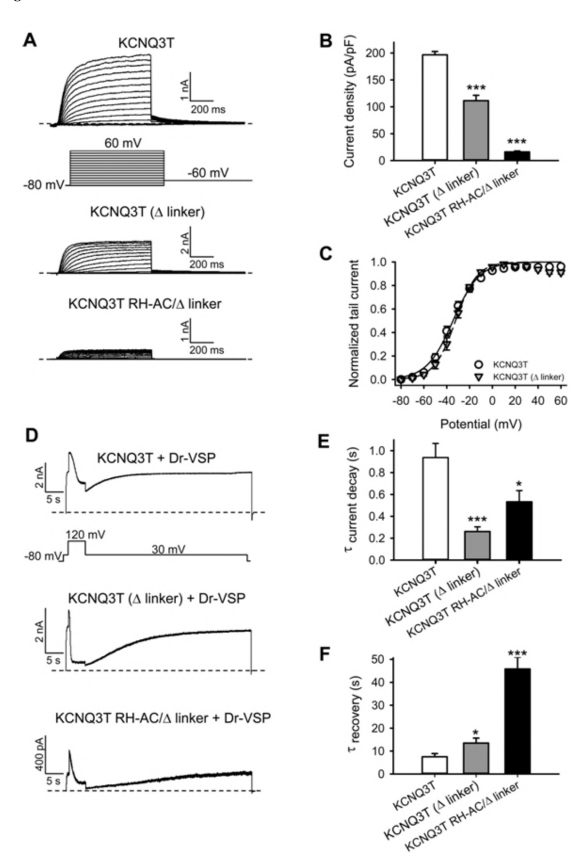


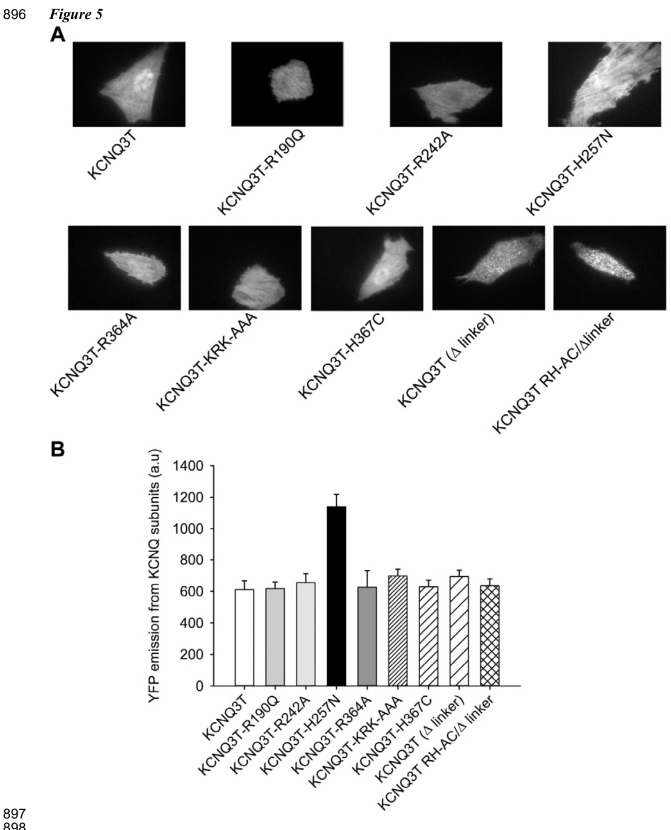
890 Fi

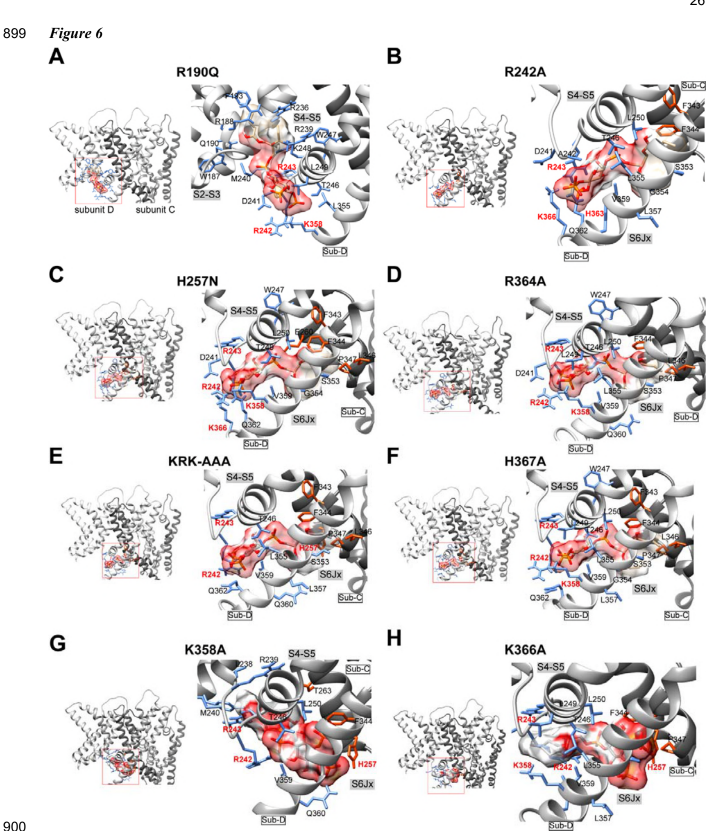


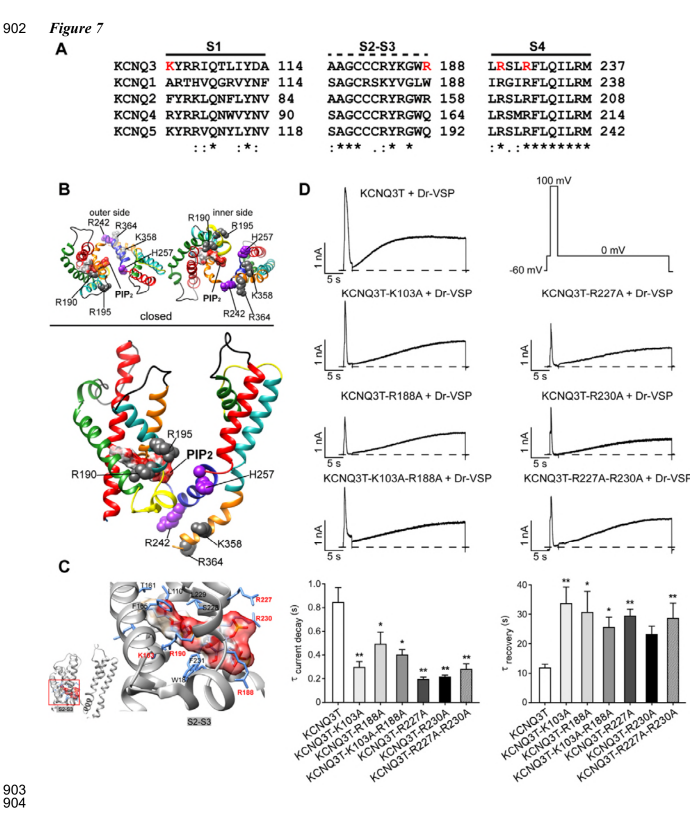


893 Figure 4

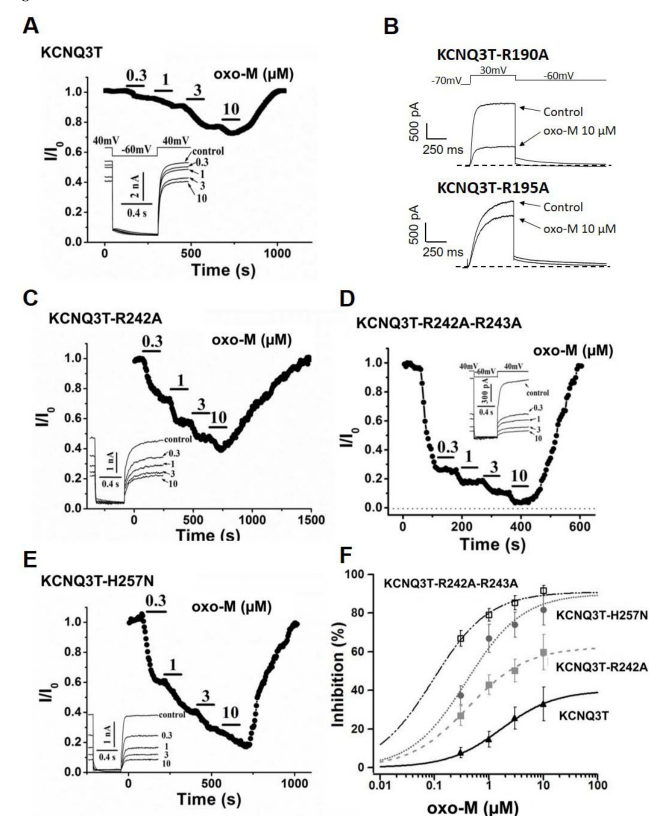








905 Figure S1



906

Figure S1. Effects of charge neutralizing mutations located in the S4-S5 linker of KCNQ3T channels on
 the sensitivity to muscarinic stimulation.

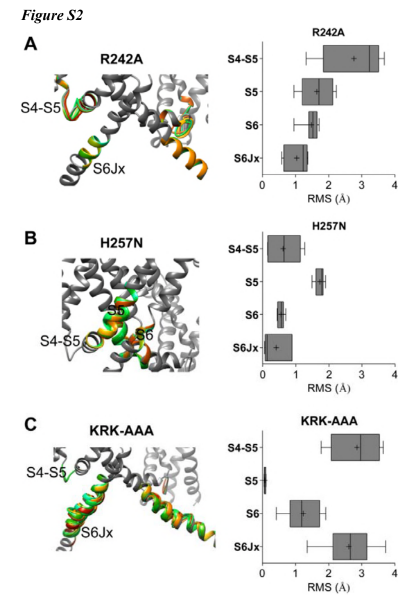
809 KCNQ3T currents during the experiment from stably M_1 receptor-expressing CHO cells transfected with 810 KCNQ3T (A) and the mutants R190A, R195A (B), R242A (C), R242A-R243A (D) and H257N (E). The

911 muscarinic agonist, oxotremorine methiodide (oxo-M), was bath-applied, as indicated. In the insets are 912 shown KCNQ3 current waveforms before and after the application of oxo-M at the concentrations indicated.

913 (F) Concentration dependence of inhibition by oxo-M of the WT (closed triangles), R242A (closed squares),

914 H257N (closed circles), and R242A-R243A (open squares) currents. The line represents the fit of

experimental data by Hill equations. Each point represents the mean \pm SEM from n = 3–7 experiments.



918 919

917

920 Structural perturbations predicted along the S4-S5 linker and the S6Jx are correlated with changes in 921 VSD-PD coupling. Enlarged views of the S4-S5 linker, S5 and the S6Jx domains showing structural 922 rearrangements predicted for the R242A (A), H257N (B) and KRK-AAA (C) mutant channels. The 923 structural elements that differ among the WT and mutant structures are indicated, and shown in gray (WT) 924 and in rainbow (mutant) on the structures. The right panels show the root mean square deviation (RMS) bar 925 plots with the disordered structural elements that differ among the WT and mutant structures. These are 926 represented as interleaved box and whiskers (25-75% percentile, median, and minimum and maximum) by 927 structural elements.