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2	Full title: The voltage sensing phosphatase (VSP) localizes to the apical membrane of kidney
3	tubule epithelial cells
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10	Short title: XI-VSP localization in kidney tubules
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18 Abstract

19 Voltage-sensing phosphatases (VSPs) are transmembrane proteins that couple changes in 20 membrane potential to hydrolysis of inositol signaling lipids. VSPs catalyze the dephosphorylation of 21 phosphatidylinositol phosphates (PIPs) that regulate diverse aspects of cell membrane physiology 22 including cell division, growth and migration. VSPs are highly conserved among chordates, and their 23 RNA transcripts have been detected in the adult and embryonic stages of frogs, fish, chickens, mice 24 and humans. However, the subcellular localization and biological function of VSP remains unknown. 25 Using reverse transcriptase-PCR (RT-PCR), we show that both Xenopus laevis VSP (XI-VSP1 and XI-26 VSP2) mRNAs are expressed in early embryos, suggesting that both XI-VSPs are involved in early 27 tadpole development. To understand which embryonic tissues express XI-VSP mRNA, we used in situ 28 hybridization (ISH) and found XI-VSP mRNA in both the brain and kidney of NF stage 32-36 embryos. 29 By Western blot analysis with a VSP antibody, we show increasing levels of XI-VSP protein in the 30 developing embryo, and by immunohistochemistry (IHC), we demonstrate that XI-VSP protein is 31 specifically localized to the apical membrane of both embryonic and adult kidney tubules. We further 32 characterized the catalytic activity of both XI-VSP homologs and found that while XI-VSP1 catalyzes 3-33 and 5-phosphate removal, XI-VSP2 is a less efficient 3-phosphatase with different substrate specificity. 34 Our results suggest that XI-VSP1 and XI-VSP2 serve different functional roles and that VSPs are an 35 integral component of voltage-dependent PIP signaling pathways during vertebrate kidney tubule 36 development and function.

37

38 Introduction

39 Phosphatidylinositol phosphates (PIPs) are lipid second messengers involved in almost all 40 facets of cell biology, including differentiation, proliferation, migration, and polarity (1,2). Many human 41 diseases are linked to mutations in PIP-modifying enzymes, including cancer, peripheral neuropathy, 42 stroke, bipolar disorder, autism and developmental disorders (3–6). As a result, PIP kinases and

43 phosphatases have been extensively studied to understand their roles in these diverse cellular 44 processes and are increasingly viewed as potential therapeutic targets (7–9). Here, we focus on an 45 exceptional member of the PIP-modifying family of enzymes, the voltage-sensing phosphatase (VSP). 46 VSP is a unique PIP-modifying enzyme whose activity is regulated through a voltage sensing 47 domain (VSD) (10). VSDs are composed of four transmembrane helices with the fourth helix (called S4) 48 containing arginines responsible for sensing changes in the electrical membrane potential. In response 49 to membrane depolarizations, the S4 helix changes conformation leading to activation of the cytosolic 50 phosphatase domain (PD), which is homologous to PTEN (phosphatase and tensin homolog deleted on 51 chromosome 10) (10), a well characterized lipid phosphatase. Once VSP is activated, its PD 52 dephosphorylates both the 5- and 3-phosphates from PIPs (10–14). As a result, VSPs provide a direct 53 connection between the electrical signaling and PIP signaling pathways. While electrical signaling 54 pathways are most often discussed in terms of neurons, all cells maintain an ionic gradient that creates 55 a membrane potential. This electrochemical force is utilized to initiate essential cell-signaling functions. 56 For example, pancreatic beta cells use their membrane potential to respond to increasing glucose 57 concentrations by activating ATP-sensitive potassium channels and voltage-gated calcium channels to 58 release insulin (15). Renal tubules also use membrane potentials, specifically activating different 59 potassium channels for regulating cell volume, potassium secretion and tubuloglomerular feedback (16). Since many ion channels are regulated by PIPs (17–19), VSPs are likely to play a role in 60 61 modulating ion channel permeability and their subsequent electrical signaling when channels and VSP 62 are co-expressed.

In line with the idea that VSP serves a fundamental physiological role, VSP orthologs have been identified in several tissues across diverse phyla, including ascidians, newts, salamanders, zebrafish, chicken, amphibians, mice, and humans (10,11,20–26). In particular, VSP mRNA transcripts have been found in the adult nervous systems of sea squirts and the brains of frogs, mice and humans (20,22,27,28) as well as in kidney, stomach, heart, testis and ovary (20,22–25,27–30). In addition to adult tissues, VSP transcripts have been detected in embryonic tissues such as the kidney and eye of

69 zebrafish (31), the kidney, brain and stomach of chicks (21,30), and the brain, spinal cord and eye of 70 mice (25,27,32). The majority of studies investigating VSP localization focused on measuring mRNA 71 expression, with a few studies reporting cellular localization of VSP protein. The first report of the sea 72 squirt VSP (*Ciona intestinalis* VSP, Ci-VSP) found protein expression in sperm tails (10). In chicks, 73 Gallus gallus VSP (Gq-VSP) protein was found in Purkinje neurons throughout the cell, which the 74 authors suggested means Gg-VSP is expressed on the plasma membrane as well as intracellular 75 membranes (30). Lastly, mouse VSP (Mm-VSP) was found in the brains of adult and neonatal mice, 76 specifically in dissociated cortical neurons (32). However, to our knowledge, a connection between VSP 77 activity and PIP-mediated biological processes in native tissues remains unknown.

78 To better understand VSP's biological role, we tested for XI-VSP localization in Xenopus laevis 79 embryos. X. laevis are an allotetraploid species and have two highly similar VSP proteins, termed XI-80 VSP1 and XI-VSP2 (22,33). We found the RNA transcripts for both in multiple stages of embryonic 81 development. The transcript expression patterns differed between the two homologs, suggesting that 82 they fulfill different biological roles. We validated a new VSP antibody, N432/21 (NeuroMab), which 83 recognizes both XI-VSPs and shows that XI-VSP protein is expressed in the brain and kidney of adult 84 and embryonic X. laevis. Furthermore, we found that XI-VSP protein is specifically located on the apical 85 membrane of both embryonic and adult kidney tubules. To probe the function of XI-VSP in these 86 tubules, we tested the catalytic activity of both XI-VSP homologs. We found that they dephosphorylate 87 the 5-phosphate from PIPs, but unlike other VSPs, XI-VSP2 is a significantly weaker 3-phosphatase. 88 From our results, we suggest that by dephosphorylating PIPs in a voltage-dependent manner, XI-VSPs 89 play a fundamental role in kidney development and function.

- 90
- 91 **Results**

92 XI-VSP mRNA transcripts in X. laevis embryos

93 The location and function of a biological molecule are often tightly coupled characteristics. Our 94 previous publication showed adult X. laevis transcript expression in several tissues including testes, 95 kidney, ovary, liver, and brain (22). Although previous studies found VSP transcripts in the embryos of 96 ascidians (28), fish (31), chicken (21,30) and mice (27), expression in X. laevis embryos has not been 97 reported. To better understand XI-VSP's physiological role, we determined the RNA transcript 98 expression of XI-VSP1 and 2 during X. laevis development. We made polyadenylated cDNA libraries 99 from the total RNA from Nieuwkoop and Faber (NF) (34) stage 12-40 embryos and conducted semi-100 quantitative reverse transcriptase PCR (sqRT-PCR) using XI-VSP1 and XI-VSP2 specific PCR primers. 101 We found that both XI-VSP1 and 2 transcripts are expressed in the developing embryo and accumulate 102 by swimming tadpole stages (NF stages >36; Fig 1A-B). Furthermore, we observed a high level of XI-103 VSP1 transcripts in early embryos, which are likely remnants of maternal transcripts in agreement with 104 our previous finding of high level expression of XI-VSP1 in the ovary (Fig 1A) (22). Our results indicate 105 that both XI-VSP1 and XI-VSP2 transcripts are present at early stages of X. laevis development and 106 that the two show different patterns of embryonic RNA expression. These results agree with our 107 previous finding that the transcripts are differentially expressed in adult tissues (22) and further suggest 108 discrete functional roles for each in developing embryos.

109

110 Fig 1. *X. laevis* embryonic stages show VSP mRNA transcripts expression.

(A-B) Semi-quantitative RT-PCR (sqRT-PCR) of a panel of *X. laevis* embryos (NF stage 12-40) using PCR primers specific for XI-VSP1 (A) and XI-VSP2 (B). Both XI-VSP1 and XI-VSP2 transcripts appear to accumulate by stage 36. No bands were seen without reverse transcriptase (not shown). All cDNA libraries were made with equal amounts of total RNA as determined by spectrophotometry and confirmed by agarose gel electrophoresis to visualize ribosomal RNA bands (S1 Fig). sqRT-PCR was repeated at least two times with at least two different embryonic cohorts. The expected PCR amplicon sizes are 389 bp for XI-VSP1 and 478 bp for XI-VSP2. Shown are representative gels.

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We next localized XI-VSP mRNA in embryos by *in situ* hybridization (ISH). We tested NF stage 32-36 *X. laevis* embryos by ISH with an antisense-stand probe against full-length XI-VSP1 mRNA since our sqRT-PCR analysis showed accumulation of mRNA expression at those stages. In agreement with our sqRT-PCR results, we found RNA transcripts in the developing pronephros, brain and brachial arches (Fig 2A), whereas no staining was seen when using a sense-strand control probe (Fig 2B). Since XI-VSP1 and XI-VSP2 mRNAs share 93% identity on the nucleotide level, it is possible this staining is due to transcripts from either XI-VSP1, XI-VSP2, or both.

126

127 Fig 2. VSP mRNA is located in the pronephros and brain of *X. laevis* embryos. (A) *In situ*

128 hybridization (ISH) of whole-mount NF stage 32 embryos. An anti-sense probe against XI-VSPs shows

129 XI-VSP transcript in the proximal pronephritic field (black arrowhead) and brain (white arrowhead) of the

130 embryos. This probe cannot distinguish between XI-VSP1 and XI-VSP2 mRNAs because of the

131 similarity between the two transcripts at the nucleotide level (93%). (B) No staining was observed by

132 ISH in a sibling embryo with a sense control probe. ISH was repeated four times with four different

133 embryonic cohorts. Shown are representative embryos.

134

135 Validating a VSP antibody

136 To determine XI-VSP protein expression in tissues, we validated a mouse monoclonal VSP 137 antibody, N432/21, developed by the NeuroMab facility at the University of California, Davis. We used 138 X. laevis oocytes as a heterologous expression system to determine the specificity of the anti-VSP. 139 Oocytes were injected with 20 ng of VSP RNAs from different species, incubated for 36 hours before 140 being lysed, and the lysates were run on SDS-PAGE gels for Western blotting. The anti-VSP N432/21 141 recognized several species of VSP, including XI-VSP1 (58 kDa), XI-VSP2 (58 kDa), Xenopus tropicalis 142 VSP (Xt-VSP, 58 kDa), FLAG-tagged Ciona intestinalis VSP (Ci-VSP, 66 kDa), and Danio rerio VSP 143 (Dr-VSP, 58 kDa) (Fig 3A). XI-VSPs heterologously-expressed in X. laevis oocytes often displayed a 144 double band at their predicted weight (58 kD), with the XI-VSP2 migrating slightly slower through the

gel than XI-VSP1 and Xt-VSP (Fig 3A-C). The nature of these doublets and the slight difference in
electrophoretic mobility between XI-VSP1, XI-VSP2, Xt-VSP, and Dr-VSP (despite their nearly identical
predicted MWs) is not presently understood. These results demonstrate that anti-VSP N432/21
recognizes both XI-VSP1 and 2 proteins, as well as the most commonly-studied VSP (Ci-VSP) and
VSPs from two other common vertebrate models of development.

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151 Fig 3. X. laevis tissues and embryos show VSP protein expression. (A) Western blot validation of 152 N432/21 anti-VSP in X. laevis oocvtes injected with RNA for Dr-VSP (Dr), FLAG-Ci-VSP (Ci), XI-VSP1 153 (XI1), XI-VSP2 (XI2), Xt-VSP (Xt), or left un-injected (U). All VSPs tested were recognized by the 154 antibody. Un-injected oocytes (U) display no band. Dr-VSP, XI-VSP1, XI-VSP2 and Xt-VSP have a 155 predicted MW of 58 kD while FLAG-Ci-VSP has a predicted MW of 66 kD. The slight difference in 156 electrophoretic mobility between VSPs and their predicted MWs and the nature of the double band for 157 XI-VSP1 and XI-VSP2 (as seen in panels B and C) has not been determined. These results show that 158 anti-VSP N432/21 is specific for VSP and cross-reacts with VSPs from multiple species. (B) Western 159 blot analysis of X. laevis tissues. Lysates from adult kidney (K, 3 µg), testis (T, 30 µg), brain (B, 10 µg), 160 and stage 44 embryo (E, 20 µg) were run against lysates from oocytes injected with RNAs for XI-VSP1 161 (XI1), XI-VSP2 (XI2) or left un-injected (U) and analyzed by Western blot with anti-VSP. A single band 162 of approximately the correct MW (58 kD) was observed in the tissue lysates, indicating the presence of 163 XI-VSP protein in all tissues tested. (C-D) Western blot analysis of X. laevis embryos. Lysates from NF 164 stage 12-40 embryos (30 µg each) were run against lysates from adult kidney (K, 5 µg) and oocytes 165 injected with RNAs for XI-VSP1 (XI1), XI-VSP2 (XI2) or left un-injected (U). Blots were probed either 166 with anti-VSP (C) or anti-actin (D) as a loading control (predicted MW 42 kD). A weak band (potentially 167 corresponding to XI-VSP1) is present only at early embryonic stages 12-20 (red arrowheads), whereas 168 a slightly slower-migrating band (potentially corresponding to XI-VSP2) accumulates at later embryonic 169 stages 36-40. Lysates, gels and blots were repeated three times with either three different adults or 170 three different embryonic cohorts. Shown are representative gels for each.

171

172 Western blots of XI-VSP protein in embryos and adult tissues

173 After validating the VSP antibody, we turned to tissue samples from adult and embryonic X. 174 laevis. Lysates made from adult kidney, testis, brain, and NF stage 44 embryos show a single band of 175 the correct approximate MW (58 kD) by Western blot, indicating that these tissues express XI-VSPs 176 (Fig 3B). The anti-VSP N432/21 recognizes both XI-VSP1 and 2 so we cannot definitively distinguish 177 whether XI-VSP1, XI-VSP2, or both are present. However, their slight difference in electrophoretic 178 mobility makes it possible to infer that XI-VSP2 is predominantly expressed in the kidney while XI-VSP1 179 expression may predominate in the testes, further supporting the likelihood of different roles for the two 180 XI-VSP proteins.

181 Next, we tested for XI-VSP protein expression in X. laevis embryos, NF stages 12-40. Western 182 blots show a faint band at the early embryonic stages, NF 12-20, and darker bands at the later 183 swimming tadpole stages, NF 36-40 (Fig 3C). Based on the slight electrophoretic differences observed 184 in the heterologous system, the slightly lower apparent MW faint bands may represent XI-VSP1. The 185 darker bands have a slightly higher apparent MW, suggesting they represent XI-VSP2. Interestingly, 186 these results suggest that XI-VSP2 is up-regulated as the tadpole develops. This pattern of XI-VSP 187 protein levels agrees with the high level of XI-VSP1 maternal transcripts in the ovary (22) and up-188 regulation of XI-VSP2 RNA during later developmental stages (Fig 1B). Furthermore, from these blots, 189 we suggest post-transcriptional regulation of XI-VSP protein expression in the developing embryo since 190 XI-VSP1 transcripts also are present in later embryonic stages yet are not detectable by Western blot 191 (Fig 3C). Overall, XI-VSP protein is present in both adult and embryonic X. laevis. The differential 192 protein expression between XI-VSP1 and XI-VSP2 is consistent with the differences observed at the 193 RNA level and further suggests the two homologs serve different functional roles.

194

195 Immunohistochemistry of XI-VSP proteins in kidney and brain

196 Since protein accumulation was not detected by immunoblotting until later embryonic stages, we 197 next used whole-mount indirect IHC on NF stage 42 embryos. Because we observed high RNA and 198 protein levels in the kidneys, we initially focused on kidney expression for the IHC experiments. To 199 facilitate kidney identification, we used two transgenic lines that express GFP in the developing kidney, 200 either Pax8:GFP (35) or Cdh17:GFP (36). Whole-mount immunostained Pax8:GFP embryos showed 201 distinct XI-VSP staining in the proximal pronephros, indicating XI-VSP protein is present during kidney 202 development (S2 Fig A-C, A'-C'). This staining was not seen in sibling embryos stained with the 203 secondary antibody alone (S2 Fig A"-C"). Focusing on the pronephritic tubules, we fixed, embedded, 204 and sectioned NF stage 42 Cdh17:GFP embryos for IHC to determine subcellular localization of XI-205 VSPs (Fig 4A-F). Interestingly, we observed XI-VSP staining in the developing pronephritic tubules. 206 specifically localized to the lumenal surface or apical membrane of the tubules (Fig 4A). This is in 207 contrast to the soluble GFP marker that is visible throughout the tubules (Fig 4B). Consecutive sections 208 probed with the secondary antibody alone show the GFP marker without any XI-VSP staining (Fig 4A'-209 F'). We also examined embryos for XI-VSP protein in brain tissue, given the presence of XI-VSP RNA 210 transcripts in embryonic brain (Fig 2A). However, we did not observe any XI-VSP staining in the 211 developing brain, either in whole-mount or in sections (data not shown). This may be due to XI-VSP 212 protein levels being too low to visualize via our IHC methods, or due to masking of the epitope during 213 fixation by XI-VSP binding partners (see discussion). Further studies involving IHC signal amplification 214 and/or antigen retrieval may be needed to distinguish between these possibilities and to localize XI-215 VSP protein in the X. laevis brain. Our findings demonstrate embryonic expression of XI-VSP protein in 216 pronephritic tubules with subcellular localization on their apical membranes, thus, suggesting XI-VSPs 217 play an important role in the development and function of the embryonic X. laevis kidney.

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Fig 4. XI-VSP protein is expressed on the lumenal surface of embryonic pronephroi. NF stage 42 Cdh17:GFP *X. laevis* embryos were sectioned and stained to test for VSP protein. Kidney tubules were identified by the presence of the Cdh17:GFP transgene (B, B'). Sections stained with anti-VSP (A)

showed fluorescence on the lumenal surface (corresponding to the apical membrane) of the proximal
kidney tubule cells (outlined in white). Anti-VSP staining was only observed in the presence of anti-VSP
(A-F) and not in secondary antibody alone control sections (A'-F'). Panels (A, A') anti-VSP or no antiVSP control; (B, B') GFP transgene; (C, C') fluorescence signal overlay of A, B and E; (D, D') bright
field images; (E, E') Hoechst 33342 (to mark nuclei); (F, F') signal overlay of A, B, D and E. IHC was
repeated three times with three different embryonic cohorts. Shown are representative sections. Scale
bar = 25 µm

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230 In addition to embryos, we also tested adult kidney tissue using IHC. Adult male X. laevis 231 kidneys were removed, fixed, embedded, sectioned, and stained with anti-VSP. The sections also show 232 XI-VSP staining on the lumenal surface of the tubules (Fig 5A), which is consistent with the subcellular 233 staining observed in embryonic kidney tubules (Fig 4A). No staining was seen in adult kidney tubules 234 when consecutive sections were probed with secondary antibody alone (Fig 5B). In contrast to our IHC 235 of embryonic kidney sections, we only observed staining in adult kidney sections by anti-VSP after 236 post-fixation antigen retrieval, which suggests a difference in the availability of the VSP antigen in 237 embryonic versus adult kidney tissues. These results suggest a long-term functional role for XI-VSPs in 238 the kidney since they are present from the earliest stages of kidney development and continue into 239 adulthood.

240

Fig 5. XI-VSP protein is expressed on the lumenal surface of adult kidney tubule epithelial cells.
(A) Adult kidney sections were stained with anti-VSP and anti-mouse IgG-Alexa 488. XI-VSP staining is
observed on the lumenal surface (marked with arrowheads) of the kidney tubules (two outlined in
white). This surface corresponds to the apical membrane and not the basolateral membrane of the
epithelial cells. (B) Apical membrane staining was not seen on a consecutive section in the absence of
anti-VSP primary antibody. IHC was repeated on sections from kidneys of three different adult males.
Shown are representative sections. Scale bar = 25 µm

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249 Functional activity of XI-VSPs

250 While VSPs from Ciona intestinalis and Danio rerio function as both 5- and 3-phosphatases (10-251 12,14,37) the XI-VSPs have previously been characterized as only a 5-phosphatase against 252 phosphatidylinositol-4,5-bisphosphate $[PI(4,5)P_2]$ (22). To understand how the XI-VSPs may contribute 253 to kidney function, we examined their phosphatase activity to determine whether both XI-VSP1 and 2 254 dephosphorylate phosphatidylinositol-3,4,5-trisphosphate $[PI(3,4,5)P_3]$ at the 5-phosphate as well as 255 whether they function as 3-phosphatases. Because XI-VSPs are regulated by voltage, we used two 256 electrode voltage clamp (TEVC) to control XI-VSP activation. To monitor all four possible 257 dephosphorylation reactions (Fig 6A), we turned to pleckstrin homology (PH) domain-based optical 258 biosensors. PH domains are well known proteins that bind specifically to certain PIPs (38-40). We 259 chose PH domains from tandem PH domain containing protein 1 (TAPP) and phospholipase C (PLC) 260 because they bind specifically to phosphatidylinositol-3,4-bisphosphate $[PI(3,4)P_2]$ and $PI(4,5)P_2$, 261 respectively, allowing us to monitor both 5- and 3-phosphatase activities (12,14,41,42). To follow 262 $PI(3,4)P_2$, we used the biosensor fTAPP that utilizes the PH domain from TAPP flanked by an N-263 terminal CFP and C-terminal YFP, while the whole sensor is anchored to the membrane through a 264 prenylation site at the C-terminus (Fig 6B) (41–43). When VSP dephosphorylates the 5-phosphate from 265 $PI(3,4,5)P_3$, the TAPP-PH domain of the fTAPP biosensor will bind the $PI(3,4)P_2$ and the resulting 266 conformational change between the CFP and YFP will change the Förster resonance energy transfer 267 (FRET) between them, leading to an increase in the FRET ratio (Fig 6B). As $PI(3,4)P_2$ is depleted when 268 VSP dephosphorylates the 3-phosphate, the TAPP-PH unbinds and the FRET ratio decreases (Fig 6B). 269 Thus, we combine the optical measurements of the biosensors with TEVC to precisely control and 270 monitor catalytic activity of XI-VSPs.

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Fig 6. XI-VSP1 and XI-VSP2 function as voltage-regulated 5- and 3-phosphatases. (A) Schematics of the known VSP reactions. (B) Cartoon representation of fTAPP binding and release with increasing 274 and decreasing PI(3,4)P₂ concentrations. The binding of fTAPP to PI(3,4)P₂ results in a conformational 275 change that increases the FRET signal. Similarly, a reduction of $PI(3,4)P_2$ results in a decrease in the 276 FRET signal. (C) Oocytes were injected with fTAPP and either XI-VSP1, XI-VSP2 or XI-VSP2 C301S 277 (XI2-CS), a catalytically inactivated protein. (left) Averaged fTAPP FRET traces over time during a 278 voltage step from a holding potential of -100 to +160 mV. The FRET signal increases and decreases in 279 the same pulse for XI-VSP1 while it only increases for XI-VSP2. (right) FRET measurements were 280 tested at several voltages and the Δ F/F fTAPP FRET ratio was plotted versus the voltage (RV). The 281 FRET increase (net 5-phosphatase reaction, solid line, closed symbols) was plotted separately from the 282 FRET decrease (net 3-phosphatase reaction, dashed line, open symbols). XI-VSP1 shows robust 283 activity as both a 3- and 5- phosphatase while XI-VSP2 functions as a 5-phosphatase. (D) Oocytes 284 were injected with fPLC and either XI-VSP1, XI-VSP2 or XI2-CS. The ΔF/F fPLC FRET RV shows a 285 FRET decrease (net 5-phosphatase reaction, solid line, closed symbols) for both XI-VSP1 and 2 286 indicating both dephosphorylate $PI(4,5)P_2$. (E) Oocytes were injected with qPLC and either XI-VSP1. 287 XI-VSP2 or XI2-CS. GFP fluorescence measurements were tested at several voltages and the 288 fluorescence voltage relationship plotted showing a fluorescence increase (net 3-phosphatase reaction. 289 dashed line, open symbols). While XI-VSP1 shows significant levels of 3-phosphatase activity against 290 PI(3,4,5)P₃, XI-VSP2 is a much less efficient 3-phosphatase. (F) Averaged fPLC FRET traces over time 291 during a voltage step from a holding potential of -100 to +160 mV for fPLC co-expressed with either XI-292 VSP1 or XI-VSP2. The kinetics of activation are significantly faster for XI-VSP1 than for XI-VSP2. All 293 error bars are \pm SEM., n \geq 8. Data fit with single Boltzmann equations

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When oocytes co-expressed XI-VSP1 and fTAPP, we observed XI-VSP1 inducing an initial increase and subsequent decrease in FRET over time, indicating that it functions as both a 5- and a 3phosphatase (Fig 6C, left). To determine the voltage dependence of activity, we tested several voltages and plotted the FRET versus voltage relationships. To better understand each reaction, we separated the FRET increase from the FRET decrease during our analysis (Fig 6C, right, blue). Note the voltage

300 dependence of the XI-VSP1 5-phosphatase reaction (FRET increase, $V_{1/2}$ = 55 mV) is lower than the 301 voltage dependence of the 3-phosphatase (FRET decrease, $V_{1/2} = 97$ mV) (S1 Table). As a control, we 302 also tested a catalytically inactive XI-VSP2 C301S (XI2-CS) (22) with fTAPP. As expected, a slight, 303 continuous FRET increase was observed. This small signal is expected because it has been previously 304 shown that a low level of endogenous XI-VSP activity is present in X. laevis oocytes (14,22,41,42,44). 305 Interestingly, the same fTAPP experiment with XI-VSP2 only showed a FRET increase (Fig 6C, green) 306 even when we extended the voltage pulse to 60 seconds (data not shown). Additionally, as was 307 previously reported (22), the voltage dependence of XI-VSP2 5-phosphatase activity against 308 $PI(3,4,5)P_3$ is shifted to higher voltages ($V_{1/2} = 92 \text{ mV}$) compared to that of XI-VSP1 ($V_{1/2} = 55 \text{ mV}$) (S1 309 Table). These results indicate that, although the XI-VSP homologs are 93% identical, they do not share 310 identical voltage-dependent functional roles in the cell. 311 To follow PI(4,5)P₂, we used two different biosensors based on the PLC-PH domain, one FRET 312 sensor which utilized the PH domain from PLC instead of TAPP to create fPLC, and a fluorescence 313 sensor where GFP is attached to the N-terminus of PLC, called gPLC. We used fPLC to monitor the 5-314 phosphatase activity during the PI(4,5)P₂ to PI(4)P reaction. When oocytes co-expressed either XI-315 VSP1 or XI-VSP2 with fPLC, we observed robust FRET ratio decreases (Fig 6D), indicating 5-316 phosphatase activity for both homologs, as was previously published (22). We used gPLC to monitor 317 the 3-phosphatase reaction, the $PI(3,4,5)P_3$ to $PI(4,5)P_2$ reaction, because the gPLC sensor is a more 318 sensitive sensor for this reaction than fPLC. Instead of a change in FRET, the gPLC translocates to the 319 membrane when $PI(4,5)P_2$ concentrations increase, resulting in an increase in fluorescence. When 320 oocytes co-expressed XI-VSP1 with gPLC, we observed an increase in fluorescence, indicating the 321 production of PI(4,5)P₂ from the XI-VSP 3-phosphatase reaction (Fig 6E, blue). Unexpectedly, the 322 same experiment with XI-VSP2 also gave a small, but reproducible fluorescence increase (Fig 6E, 323 green), indicating XI-VSP2 is also able to function as a 3-phosphatase. However, we did not observe

any 3-phosphatase activity when using the fTAPP sensor. These results show that XI-VSP2 can

325 remove the 3-phosphate from $PI(3,4,5)P_3$ but not from $PI(3,4)P_2$. Further investigations are needed to

better understand this striking difference in substrate specificity. Interestingly, the kinetics of XI-VSP2 are significantly slower than the kinetics of XI-VSP1 (Fig 6F), as seen by the voltage steps to 160 mV for a 2 second voltage step for XI-VSP1 and a 10 second voltage step for XI-VSP2 (Fig 6F). Overall, our results show that XI-VSP1 can catalyze the same four reactions as other VSPs while XI-VSP2 is restricted by substrate. Since the reactions and kinetics appear to be different between XI-VSP1 and 2, they likely fulfill different functional roles in tissues which correlates with their differential expression patterns.

333

334 **Discussion**

VSPs provide a direct link between the electrical state of a cell and its PIP signaling pathways. By dephosphorylating PIPs in a voltage dependent manner, VSPs could regulate PIP-dependent processes such as cortical cytoskeleton remodeling, ion channel function, G protein-coupled receptor signaling (GPCR) and intracellular calcium signaling (18,19,45). Though the biophysical properties of VSPs have been the subject of intense study since the discovery of Ci-VSP (10), the physiological relevance of this direct link remains unclear.

341 To elucidate XI-VSPs' biological role, we determined the cellular expression pattern of XI-VSP 342 RNA and protein in Xenopus laevis. We extended our previous findings in adult X. laevis tissues (22) by 343 showing both XI-VSP1 and 2 RNA transcripts in developing embryos. Though VSP is seen in the 344 embryonic stages of other species, this is the first study showing XI-VSP RNA and protein in X. laevis 345 embryos. Our findings indicate that XI-VSP RNAs are present at early stages of development, NF 346 stages 12-40, with both XI-VSP1 and 2 RNA levels increasing from NF stage 20 to 40. At the same 347 stages, we observed an increase in protein levels by Western blot, consistent with our RNA detection, 348 indicating an upregulation of the protein during development. Using ISH, we observed XI-VSP RNA 349 transcripts in both the brain and pronephros of NF stage 32-36 embryos. Using IHC, we found XI-VSP 350 protein in the pronephroi of NF stage 42 embryos. Upon sectioning the embryos, we observed anti-VSP 351 staining on the lumenal surface of the pronephritic tubules, corresponding to the apical membrane of

the epithelial cells lining the tubule lumen. Additionally, we sectioned adult kidneys and found the same apical membrane localization in the tubules. To directly test XI-VSP catalytic activation, we combined electrophysiology with optical biosensors to show that both XI-VSPs dephosphorylate the 3- and 5phosphate from PIPs though with different substrate specificities and efficiencies. Combining our localization results with our electrophysiological activity data, we suggest that XI-VSPs have specific functional roles in both the brain and kidney.

358 Concentrating on the kidney, we found XI-VSPs spanning the earliest stages of the developing 359 pronephros into the full adult kidney, suggesting XI-VSPs modulate PIP concentrations on the apical 360 membrane of the kidney epithelial cells in a voltage dependent manner. Kidney tubules are lined with 361 specialized epithelial cells that regulate vertebrate solute homeostasis (46,47). The transepithelial 362 potential difference across these cells contributes to tubular reabsorption of solutes and water (48,49). 363 The lumenal surface of these epithelial cells are decorated with microvilli and primary cilia, both of 364 which function in tubular reabsorption by sensing fluid flow (50–54). Microvilli provide a large absorptive 365 membrane surface for kidney epithelial cells and are actin-dependent dynamic structures with turnover 366 rates in the tens of minutes (55). Heterologously expressed Gg-VSP influences actin-based 367 cytoskeleton rearrangements in cultured chick fibroblasts leading to morphological changes in cortical 368 cell structure (56), suggesting XI-VSPs could modify the actin-based microvilli in kidney epithelial cells. 369 In addition, GPCR signaling originating in microvilli at the apical membrane of proximal kidney tubules 370 is directly influenced by local $PI(4,5)P_2$ concentrations (57), which is a substrate for both XI-VSPs. 371 Likewise, primary cilia are dependent on local $PI(4,5)P_2$ concentrations (58), and importantly, trigger 372 calcium entry due to the mechanical forces from the fluid flow (54). As calcium enters the cells, the 373 resulting depolarization could lead to activation of the XI-VSP on the apical membrane and subsequent 374 regulation of the cortical cytoskeleton, GPCRs, and ion channels during renal tubule development and 375 adult renal function.

Our results are consistent with VSP homologs having a broad role in multiple tissues.
 Specifically, our study agrees with previous studies that also showed embryonic kidney tubule

378 expression of VSP in zebrafish (31), and chicks (21). In particular, Gg-VSP RNA was found on the 379 proximal part of the nephrogenic tubules of HH stage 26 chicks and not the distal or collecting tubules, 380 consistent with our IHC results showing proximal tubule localization in NF stage 32 embryos (S2 Fig). 381 Our findings in X. laevis are also consistent with the findings in chicks in that not all tubules appear to 382 immunostain with our VSP antibody, indicating that the XI-VSPs may be restricted to a functional 383 subset of the developing and adult kidney. Our results go further because we observe clear XI-VSP 384 protein immunostaining on the apical membrane of the tubules where reabsorption of nutrients occurs 385 and where PIP-mediated signaling originates.

386 Not all VSP studies analyzed kidney expression patterns. The human VSP, Hs-VSP1 387 (previously called TPIP and TPTE2), was not examined in kidneys, making human VSP kidney 388 expression unknown (20). In addition to kidney expression, VSP is also expressed in the brain and 389 nervous system of multiple species (20,22,28,30,32). Indeed, we observed XI-VSP RNA transcripts and 390 protein in the brain of adult and embryonic X. laevis. Some reports of VSP expression conflict. 391 particularly regarding expression of mouse VSP, Mm-VSP (previously called PTEN2). Early studies 392 identified Mm-VSP as a testis specific protein (25) while subsequent investigations report expression in 393 the brain (32). While more research is needed, a pattern is emerging with the kidney and brain being 394 the most consistent tissues expressing VSP. Our results support this pattern and a functional role for 395 VSP in both the brain and kidney of multiple vertebrate species.

396 The subcellular localization of VSP is also debated in the field. Prior studies of native VSP protein 397 have found VSP on the plasma membrane of ascidian sperm (10) and of Purkinje neurons in chick 398 cerebellum sections (30). Along with plasma membrane localization, however, the cerebellum sections 399 also showed intracellular staining with a Gg-VSP antibody. A more recent study in mice used 400 dissociated cortical neurons from P0-1 mice and the staining appears throughout the cells (32). In 401 addition, heterologous expression of mouse and human VSP consistently show intracellular membrane 402 localizations (20.25.29). In contrast, our subcellular localization of endogenous XI-VSP protein is on the 403 apical membrane of kidney tubule epithelia. The differences in our results and those from prior studies

may lie in the tissues tested, the antigen retrieval discussed below, or in the antibodies used. Further
experiments are needed to determine whether VSPs function on both intracellular and plasma
membranes. While not tested against either chicken or mouse VSPs, our antibody does cross react
with VSPs from *D. rerio*, *X. tropicalis and C. intestinalis*. As a result, N432/21 may prove to be an
invaluable tool in future studies of VSP's physiological role in vertebrate kidney development and neural
function.

410 It is interesting to note that unlike in fixed embryonic kidney tissue, we only observed staining of 411 adult kidney tissue after post-fixation antigen retrieval. This difference in VSP epitope availability could 412 be due to different roles for VSP in adult versus embryonic kidney tubules, and differential VSP binding 413 partners in these tissues. We recently demonstrated that Ci-VSP forms dimers (42). Since the two XI-414 VSP homologs shown here display different expression in the embryo and adult frog, it is also possible 415 that the difference we observed in VSP epitope availability is due to the presence of XI-VSP

416 homodimers and heterodimers.

417 In addition to our localization experiments, we also tested the electrophysiological characteristics 418 of both XI-VSP homologs. We controlled their activation using TEVC and monitored the production and 419 depletion of different PIPs using optical biosensors. Our previous report indicated that both homologs 420 catalyzed the dephosphorylation of the 5-phosphate from $PI(4,5)P_2$. VSPs from other species have 421 been shown to utilize $PI(3,4,5)P_3$ and $PI(3,4)P_2$ as substrates as well. Here, we tested all three 422 substrates by using PIP sensors that are sensitive to either $PI(3,4)P_2$ or $PI(4,5)P_2$ concentrations 423 (TAPP-PH or PLC-PH respectively), allowing us to monitor the 3-phosphate removal from PI(3,4,5)P₃ 424 and PI(3,4)P₂ as well as the 5-phosphate removal from PI(3,4,5)P₃ and PI(4,5)P₂. We found that XI-425 VSP1 behaves much like other VSPs in that it dephosphorylates both 5- and 3-phosphates. XI-VSP2, 426 on the other hand, is a slower 5-phosphatase and a weaker 3-phosphatase. After modifying our 427 protocols for longer depolarizations to account for the slower kinetics, we observed clear 5-428 phosphatase activity for XI-VSP2 using both fTAPP and fPLC sensors. The voltage-dependence of 429 activation was shifted to higher voltages for XI-VSP2 compared to XI-VSP1. We also observed 3-

- 430 phosphatase activity of XI-VSP2 against PI(3,4,5)P₃, but not against PI(3,4)P₂, even after a one minute
- 431 depolarization. This difference between the XI-VSP homologs further indicates that the two are not
- 432 interchangeable and serve different physiological roles in the cell.

In conclusion, we observe XI-VSP expression in the brain and kidney of *X. laevis* embryos and adults. We localize XI-VSP protein on the apical membrane of kidney tubule epithelial cells, and we characterize both XI-VSPs as 3- and 5- phosphatases with different enzymatic activities. Our results indicate a role for voltage-dependent dephosphorylation of both 5- and 3-phosphates from PIPs on the apical membrane during kidney tubule development and renal function.

438

439 Materials and methods

440 Materials

441 *Ciona intestinalis* VSP (Ci-VSP, NM_001033826) and *Danio rerio* VSP (Dr-VSP,

442 NM 001025458) plasmids were kindly provided by Y. Okamura (Osaka University, Osaka, Japan). X. 443 laevis VSP1 (XI-VSP1, NM 001096603), X. laevis VSP2 (XI-VSP2, NM 001280607), and X. tropicalis 444 VSP (Xt-VSP, NM 001015951) expression plasmids were made in Laurinda Jaffe's laboratory and are 445 available through Addgene (Cambridge, MA). While the original human gene was named 446 transmembrane phosphatase with tensin homology (TPTE), the majority of the literature refer to this 447 protein as the voltage sensing phosphatase or VSP. Here, we follow this more common nomenclature 448 to avoid confusion. fPLC and fTAPP were both kindly provided by E.Y. Isacoff (University of California, 449 Berkeley). GFP-PLC-PH was kindly provided by T. Meyer (Stanford University). Mutations and epitope 450 tags were created by site-directed mutagenesis with Pfu Turbo DNA polymerase (Agilent, Santa Clara, 451 CA) by standard protocols. All DNA constructs were confirmed by sequencing. cRNA was transcribed 452 using SP6 or T7 mMessage mMachine kits (Thermo-Fisher, Waltham, MA). X. laevis oocytes for XI-

- 453 VSP activity assays were purchased from Xenopus 1 (Dexter, MI). Mouse monoclonal anti-VSP
- 454 (N432/21; RRID:AB_2716253) was made by the UC Davis/NIH NeuroMab Facility and is available

455 through Antibodies Incorporated (Davis, CA). Anti-beta actin (clone C4, catalog # sc-47778) was 456 purchased from Santa Cruz Biotechnology (Dallas, TX). 3G8.2C11 antibody was obtained from the 457 European Xenopus Resource Centre (59,60). Goat anti-mouse IgG, light chain specific, conjugated to 458 horseradish peroxidase (HRP, catalog # 115-035-174) was purchased from Jackson ImmunoResearch 459 (West Grove, PA). Goat anti-mouse IgG conjugated to Alexa Fluor 488 or Alexa Fluor 594 was 460 purchased from Thermo Fisher. Anti-digoxygenin (DIG) conjugated to alkaline phosphatase (AP) was 461 purchased from Sigma-Aldrich (St. Louis, MO). Cdh17:GFP and Pax8:GFP transgenic Xenopus laevis 462 frogs were obtained through the National Xenopus Resource (Woods Hole, MA, RRID; SCR 013731) 463 (35,36,60). All animals in this study were used in accordance with protocols approved by the Montana 464 State University IACUC.

465

466 **Reverse-transcriptase PCR**

467 Total RNA was isolated from embryos with TRIzol (Thermo Fisher), quantitated by Nanodrop 468 spectrophotometry and analyzed by ethidium bromide agarose gel electrophoresis (S1 Fig). Two µg of 469 embryo RNA was treated with DNAse I (DNA-free, Thermo Fisher) to remove genomic DNA and 470 cDNAs were reverse-transcribed by priming with oligo-dT using SuperScript IV reverse transcriptase 471 (Thermo Fisher). A parallel control reaction was performed without reverse-transcriptase for each 472 DNAse I-treated RNA. Semi-guantitative PCR was performed on 1/20th of the resulting cDNA using 473 primers for XI-VSP1 (forward: GATGCTGGAAACAACTCCATAGTCC; reverse: 474 GCTGTGTGTGTGGGTCAGAACAC) or XI-VSP2 (forward: GATGCTGGGAACAATTCCGTAGTCA; 475 reverse: GGGTAATAGTACGTTAAGAAGTG) with Tag polymerase in Standard Tag PCR buffer (New 476 England Biolabs, Ipswich, MA) with the thermocycling parameters: 95°C for 30 sec, 37 cycles of 62°C 477 for 20 seconds followed by 68°C for 30 seconds. For each primer pair the forward primer is located 478 upstream of an intron and the reverse primer is located in the 3' UTR to ensure alloallele specificity and 479 amplification of only cDNA. PCR products were analyzed by ethidium bromide agarose gel 480 electrophoresis.

481

482 In-situ hybridization (ISH)

Full-length antisense and sense digoxygenin (DIG)-labeled probes were made from linearized pcDNA3-XI-VSP1 plasmid with SP6 or T7 RNA polymerase and DIG NTPs (Sigma-Aldrich). *In situ* hybridization to detect XI-VSP mRNAs was performed on NF stage 32-36 embryos fixed for 90 minutes in phosphate-buffered 4% paraformaldehyde (pH 7.4) by standard methods (61). Embryos were probed with anti-DIG IgG-AP, stained with NBT-BCIP (Sigma-Aldrich) and imaged with a ProgRes C14plus camera (Jenoptic, Germany) through a 5x objective on an Axioscope.A1 microscope (Zeiss, Germany).

490

491 Defolliculating X. laevis oocytes

492 X. laevis ovaries were purchased from Xenopus 1 (Dexter, MI). Ovaries were processes as 493 described previously (62). Briefly, each ovary was washed once and morselized in Ca²⁺-free (96 mM 494 NaCl, 2 mM KCl, 1 mM MgCl₂, and 10 mM HEPES, pH 7.6). The morselized ovary was washed in Ca²⁺-495 free to remove yolk from lysed oocytes and then digested at room temperature for 30-45 minutes in 2% 496 collagenase (Sigma-Aldrich, catalog # C-0130), 0.1% soybean trypsin inhibitor (Sigma-Aldrich, catalog 497 # T-9003), and 0.1% BSA (Sigma-Aldrich, catalog # A3294) made in Ca²⁺-free. Post-digestion, the 498 oocytes were washed in oocyte wash buffer (34 mM KH₂PO₄, 66 mM K₂HPO₄, 0.1% BSA, pH 6.5) at 499 least 10 times to remove follicles, followed by 10 washes with Ca²⁺-free. The oocytes were then sorted 500 into ND-96 (96 mM NaCl, 2 mM KCl, 1.8 mM CaCl₂, 1 mM MgCl₂, 50 mg/ml gentamicin, 2.5 mM 501 sodium pyruvate, and 10 mM HEPES, pH 7.6) and cultured at 18°C.

502

503 Western Blotting

504Adult Xenopus laevis tissue, oocyte, and embryo lysates were made by homogenization and brief505sonication in lysis buffer (150 mM NaCl, 150 mM dithiothreitol, 0.1% CHAPS, 50 mM Tris, pH 7.4) with

506 protease inhibitors (Roche, Switzerland). X. laevis oocytes were injected with 20 ng VSP RNA and 507 cultured at 18°C for 36 hours before lysis. Protein concentrations were determined by a Quick Start 508 Bradford assay (Bio-Rad, Hercules, CA), Lysates were run on 10% polyacrylamide cels at 200 volts for 509 one hour, transferred to ProTran nitrocellulose membranes (GE Healthcare, Chicago, IL) in sodium 510 borate transfer buffer for 70 minutes at 350 mA and stained with Ponceau-S to evaluate equal loading 511 of protein between the wells. Membranes were blocked in TBS-T block buffer (5% non-fat dry milk, 150 512 mM NaCl, 10 mM, 0.1% Tween-80, 10 mM Tris, pH 7.4) and probed with anti-VSP supernatant at 1:25 513 dilution or anti-beta actin at 1:500 dilution. Blots were then washed in TBS-T, probed with anti-mouse 514 IgG-HRP and developed with ECL reagent (Sycamore Life Sciences, Houston, TX).

515

516 Immunohistochemistry (IHC)

517 Adult Xenopus laevis kidneys and NF stage 40-42 embryos (34) were fixed in phosphate-518 buffered 4% paraformaldehyde (pH 7.4) for 90 minutes followed by sequential equilibration for two 519 hours each in 30% sucrose, 30% sucrose/50% O.C.T (Optimal Cutting Temperature) embedding media 520 (Fisher, Hampton, NH), and 100% OCT. Kidney tissues were then snap-frozen in OCT using a dry 521 ice/ethanol slurry and sectioned at 14 µm onto glass slides coated with 0.5% porcine gelatin, type A 522 (Sigma-Aldrich, catalog # G2500) and 0.05% chromium potassium sulfate. Sections were air-dried 523 overnight at room-temperature and stored at -80°C. Sections were rehydrated with phosphate-buffered 524 saline (PBS; 137 mM NaCl, 27 mM KCl, 100 mM Na₂HPO₄, 18 mM KH₂PO₄, pH 7.4) and subjected to 525 antigen retrieval in 10 mM sodium citrate, 0.05 % Tween-80, pH 6.0 for 20 minutes at 95°C followed by 526 quick cooling on ice. Sections were re-equilibrated with PBS-T (0.1% Triton X-100) and blocked in 527 PBS-T with 10% normal goat serum (Fisher, catalog # 16-210-064). Sections were probed with anti-528 VSP supernatant at 1:25 dilution, washed in PBS-T, and probed with anti-mouse IgG-Alexa Fluor 488 529 or anti-mouse IgG-Alexa Fluor 594 at 1:1000 dilution with 1 µg/mL Hoechst 33342 (Thermo-Fisher) to 530 stain nuclei. Immuno-stained sections were mounted in Vectashield (Vector Labs, UK) under glass 531 coverslips and imaged with a 63x/1.4 NA objective on a Leica SP8 confocal microscope (Leica, Buffalo

532 Grove, IL). Embryo sections were similarly fixed, sectioned, stained and imaged without the antigen 533 retrieval process. Whole-mount embryos were similarly fixed and stained without the antigen retrieval 534 process and with four hours to overnight incubations to allow for antibody penetration. Whole-mount 535 IHC images were captured using Progres Mac CapturePro software on a ProgRes C14plus camera 536 through a Zeiss Plan-neofluar 5x/0.16 NA objective on an Axioscope.A1 microscope. GFP or Alexa 537 Fluor 594 fluorescence in whole-mount immuno-stained embryos was excited by an LED and collected 538 through Zeiss FITC (EX BP 475/40, BS FT 500, EM BP 530/50) or Texas Red (EX BP 560/40, BS FT 539 585, EM BP 630/75) filter sets, respectively. Anti-VSP, anti-3G8 and matching secondary-alone control 540 samples were imaged using the same excitation intensity and digital gain. IHC was repeated a total of 541 three times with tissue from different animals or clutches of embryos.

542

543 Electrophysiology and fluorescence measurement of activity

544 Two electrode voltage clamp (TEVC) was performed as previously described (14). FRET-based 545 PIP sensors (41,42) were used to measure depletion of $PI(4,5)P_2$ or the production and depletion of 546 PI(3,4)P₂. The PIP sensors were designed by adding an N-terminal CFP and a C-terminal YFP to 547 pleckstrin homology (PH) domains. The PH domains were originally derived from phospholipase C 548 (PLC) for $PI(4,5)P_2$ and the tandem PH domain containing protein 1 (TAPP) for $PI(3,4)P_2$, and hence 549 were called fPLC and fTAPP respectively. To measure production of PI(4,5)P₂, the diffusion based 550 sensor GFP-PLC (gPLC) was used which has a GFP was attached to the N-terminus of the PLC-PH 551 domain. All XI-VSP cRNAs (0.8 µg/µl) were mixed with either fPLC cRNA (0.4 µg/µl), fTAPP cRNA (0.4 552 $\mu g/\mu l$) or gPLC cRNA (0.06 $\mu g/\mu l$) and injected into X. laevis oocytes. In all experiments, 50 nl of the 553 cRNA mixtures were injected into oocytes and incubated in ND-96 for 30-40 hours. On the day of the 554 experiments, cells were transferred from ND-96 to ND-96' (ND-96 without gentamicin and sodium 555 pyruvate) containing 8 μ M insulin to promote PI3 kinase activity and up-regulate PI(3,4,5)P₃ levels. 556 A Leica DM IRBE inverted microscope with a Leica HC PI APO 20×/0.7 fluorescence objective 557 was used with a Dagan CA-1B amplifier and illuminated with a Lumen Dynamics X-Cite XLED1 light

558 source. Fluorescence was measured with a ThorLabs photomultiplier tube (PMT). The amplifier and 559 light-emitting diode were controlled by a Digidata-1440A board and Axon™ pClamp™ 10.7 software 560 package (Molecular Devices). For the FRET experiments, light was filtered through a HQ436/20 561 excitation filter and directed to the objective with a 455LP dichroic (Chroma). The microscope cube did 562 not contain an emission filter, because the ThorLabs PMT module contains its own cube. Thus, the 563 emitted light was filtered before the PMTs with a 510-nm dichroic, an HQ480/40 emission filter for CFP, 564 and an HQ535/30 emission filter for YFP (Chroma). For the gPLC experiments, light was filtered 565 through a HQ470/40 excitation filter, an HQ525/50 emission filter and a Q496LP dichroic (Chroma). 566 The voltage protocol consisted of steps from -100 to 180 mV in irregular increments. The length of the 567 voltage pulse was 1.5-2 s for XI-VSP1 and varied for XI-VSP2, 2-10 s for fTAPP, and fPLC, and 3 s for 568 gPLC. Rest periods of 1-3 min between voltage steps were used to allow the cell to recover depleted 569 PIP concentrations before the next voltage step. The resulting FRET or fluorescence was then plotted 570 versus the voltage to generate the fluorescence versus voltage relationship.

571

572 Data analysis

573 For the FRET experiments, AxonTM ClampfitTM 10.7 (Molecular Devices) was used to calculate 574 the FRET ratio. For the FRET or fluorescence increases (fTAPP and gPLC), the Δ F/F,

 $(F_{after} - F_{before})/F_{before}$, was calculated from the pre-pulse baseline to the max signal increase. For the 575 576 fPLC FRET decrease, the Δ F/F was calculated from the pre-pulse baseline to the max FRET decrease. 577 For the fTAPP FRET decrease, the Δ F/F was calculated from the max FRET increase to the max FRET 578 decrease. Calculations were done in Excel (Microsoft) or IGOR Pro (WaveMetrics) and data was 579 plotted in IGOR Pro. When bleaching was observed, it was corrected by fitting to a line and dividing the 580 values by the fit to obtain the final, bleaching-corrected signal. Final voltage dependent curves were 581 plotted with change in FRET or fluorescence on the Y-axis and voltage on the X-axis. The data were fit 582 to single Boltzmann equations. Activity assays were repeated on oocytes extracted on different days

583	from at least three different frogs (biological replicates) until data from a minimum of 8 oocytes were
584	acquired and analyzed. Error bars indicate standard error of the mean (SEM). IHC images were
585	processed with NIS-Elements (Nikon). Fluorescence signals from anti-VSP and matching secondary-
586	alone control images were processed in parallel while applying the same post-image processing
587	parameters in each case. Figures were made either in Illustrator or Photoshop 2018 (Adobe).
588	

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594

595 Author contributions

W. Ratzan and S.C. Kohout conceived and designed the experiments. W. Ratzan, S.E. Killian, V.
Rayaprolu, R. Bradley and S.C. Kohout collected and analyzed data. W. Ratzan, V. Rayaprolu and S.C.
Kohout wrote the manuscript. All authors approved the final version of the manuscript.

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753

- 754 Supporting Information
- 755 **S1 Table. Voltage dependence of activity.**

756

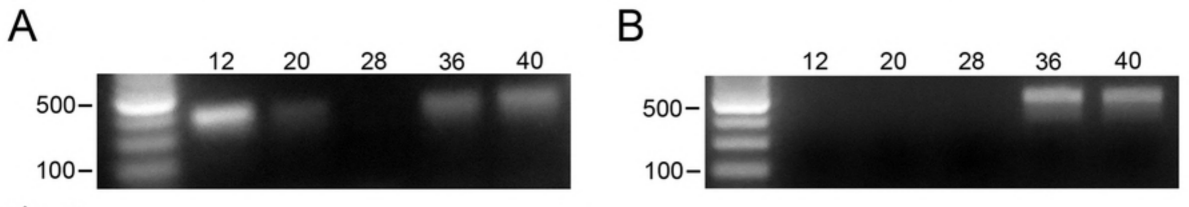
757 S1 Fig. Embryo total RNAs used for sqRT-PCR of XI-VSPs (NF stages 12-40). RNA

concentrations were determined by Nanodrop spectrophotometry and 1 μ g of RNA was run on

- a 2% agarose gel stained with ethidium bromide. The prominent bands seen here are the 28S
- and 18S ribosomal RNAs, indicating that the RNAs are not degraded and that approximately
- requal total RNA was used for subsequent sqRT-PCR analysis.

763 S2 Fig. Whole-mount immunohistochemistry (IHC) of NF stage 42 Pax8:GFP embryos.

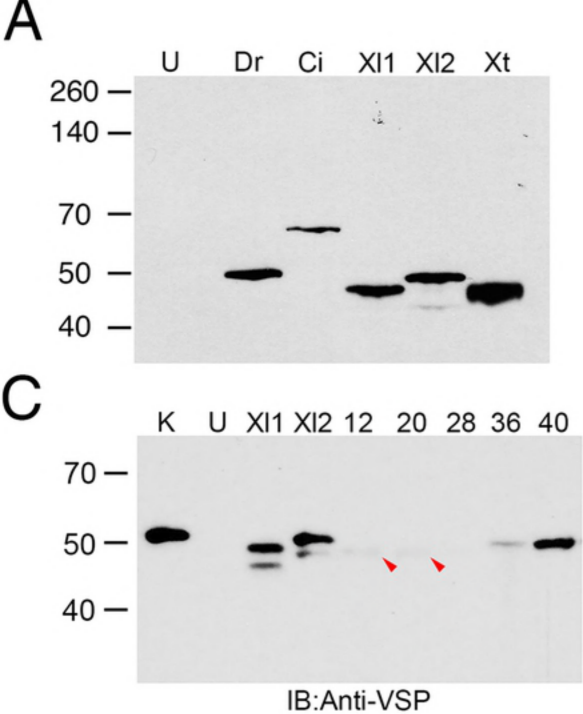
- Embryos stained with 3G8 antibody (A-C), anti-VSP (A'-C') or secondary antibody alone
- control (A"-C"). The Pax8:GFP embryos show strong GFP expression in the proximal
- pronephros (B-B") as confirmed by staining with the proximal pronephros marker 3G8 antibody
- 767 (C). XI-VSP staining of tubules is clearly visible (C') and co-localizes with the GFP, indicating
- 768 XI-VSP in the proximal pronephros. Positive staining is marked with arrowheads, whereas no
- signal above autofluorescence was seen in control animals stained with secondary antibody
- 770 alone (C").

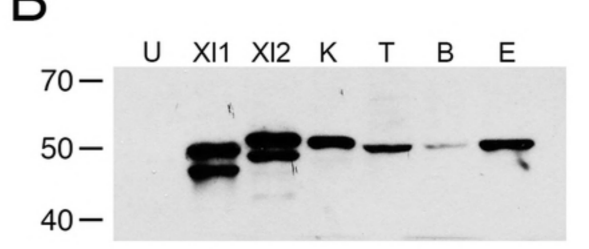


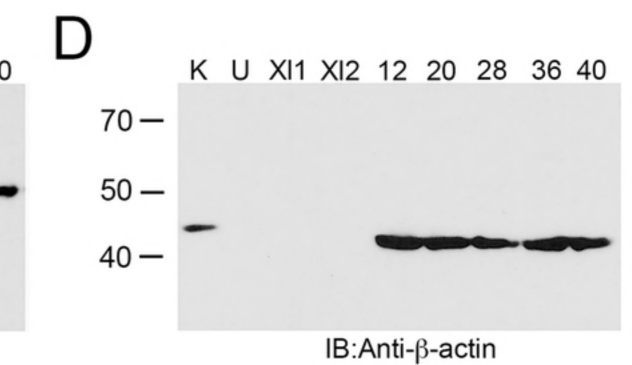












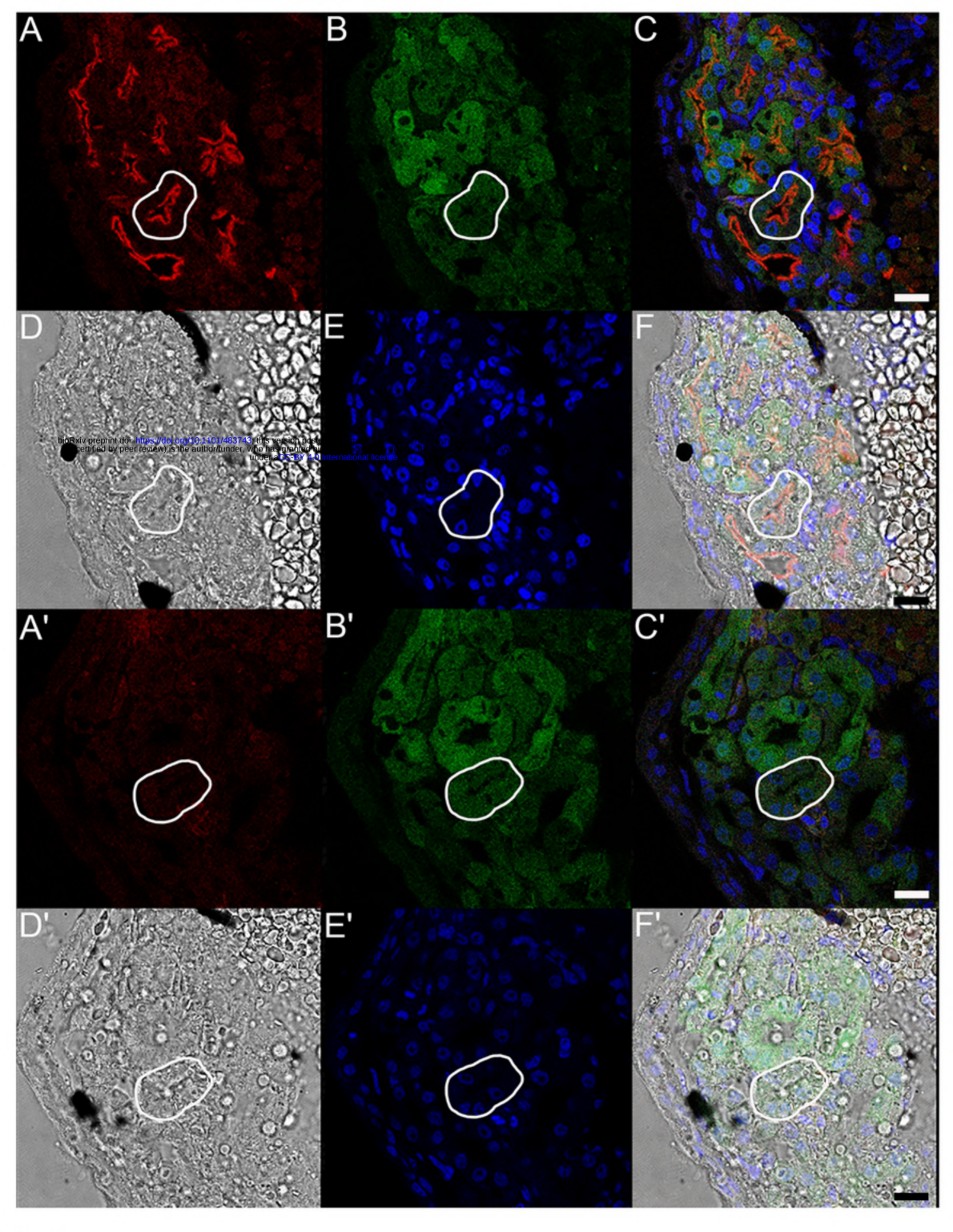
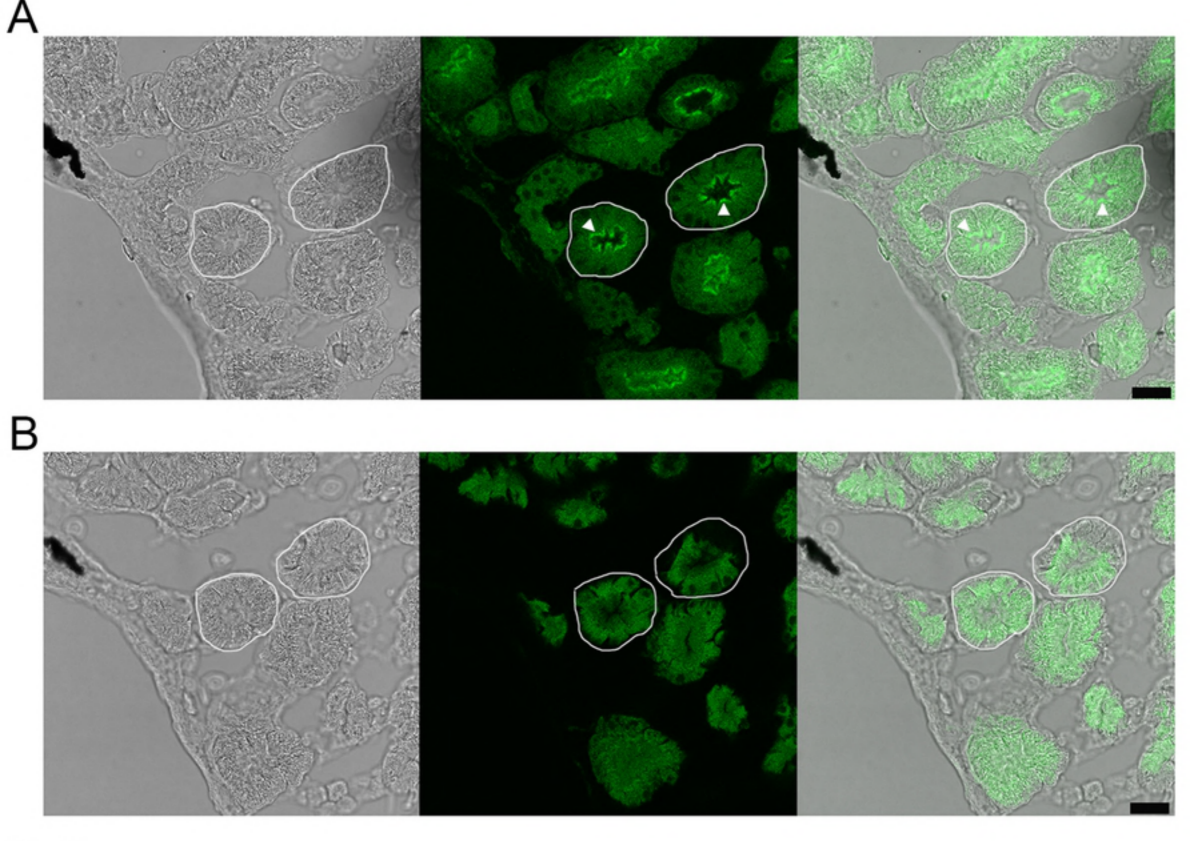
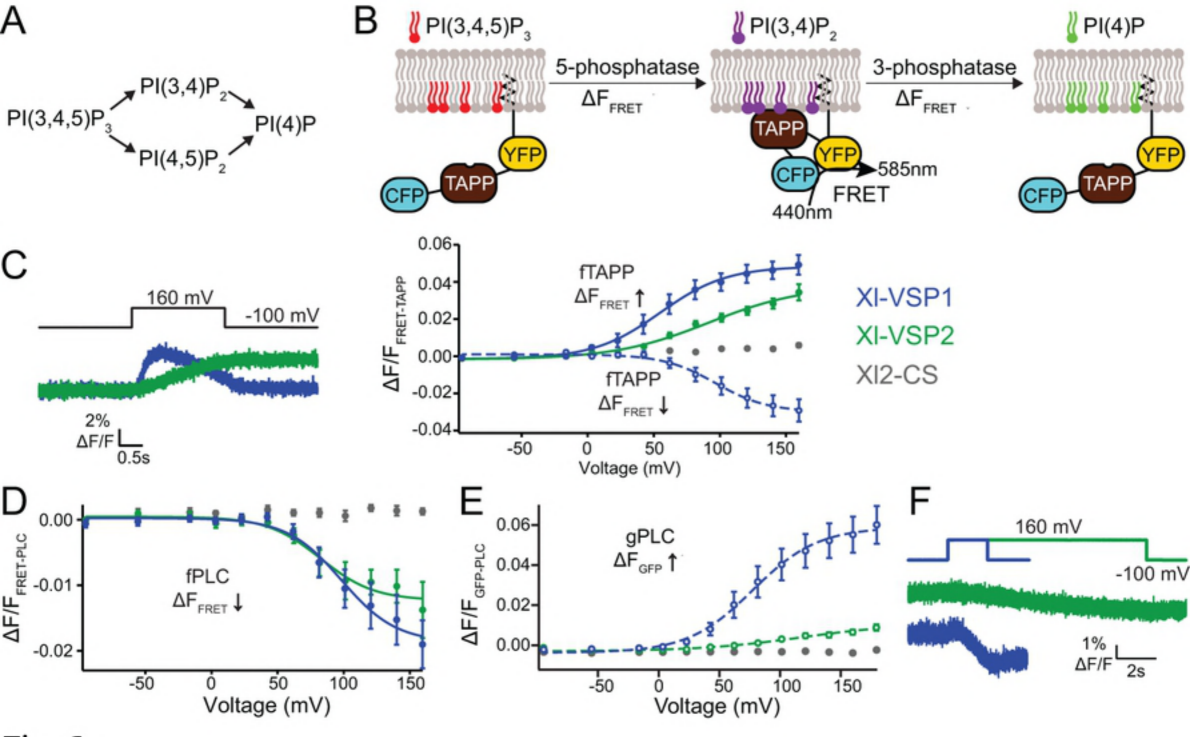


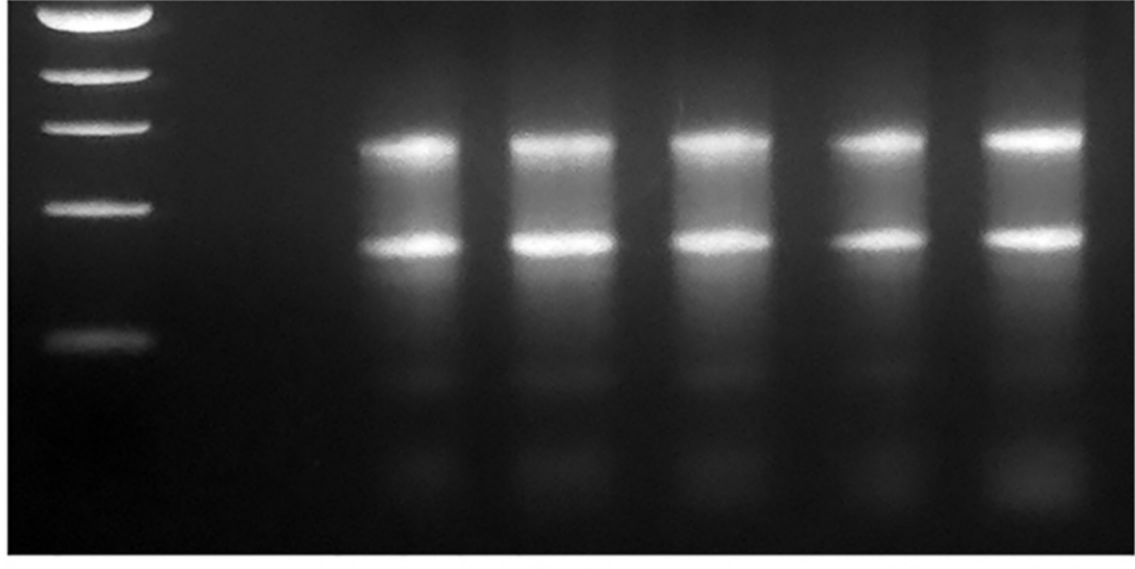
Fig 4





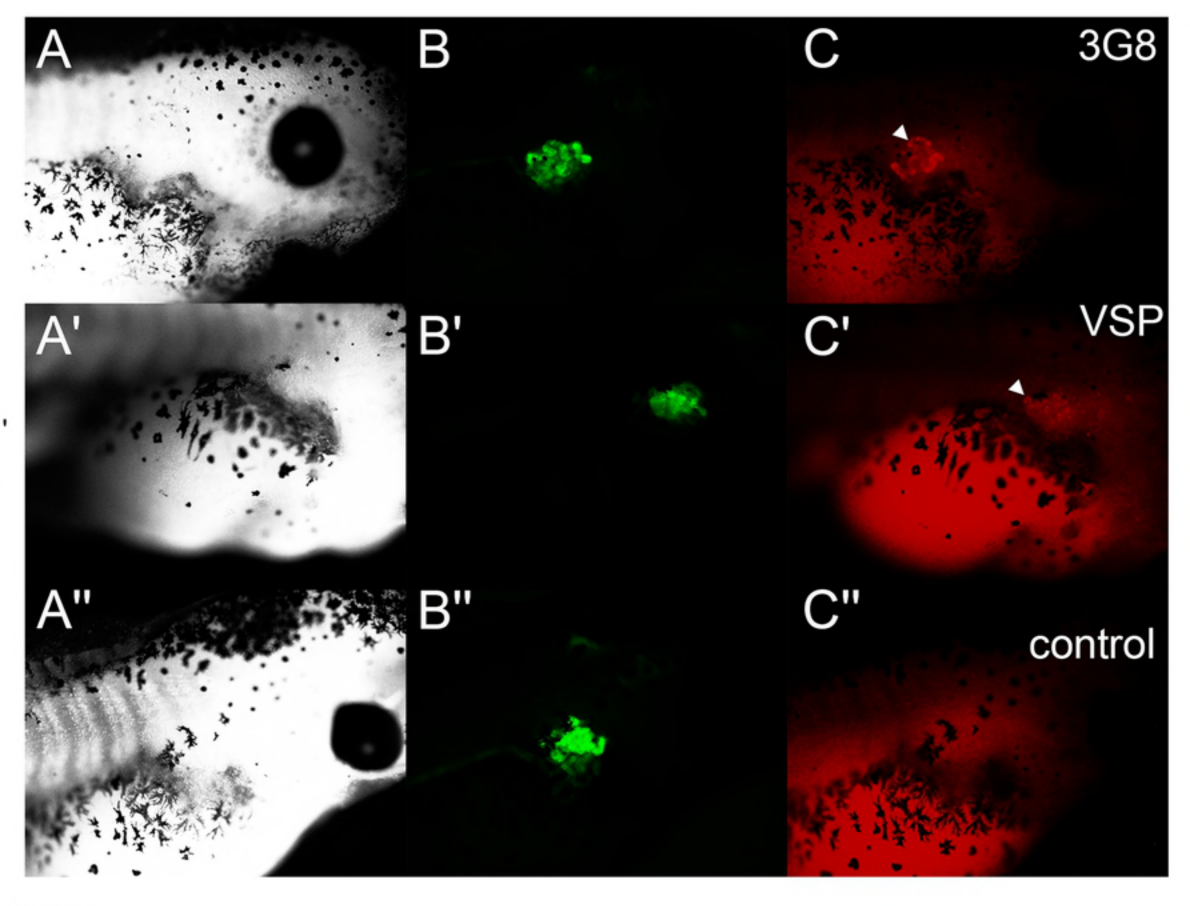
VSP	fTAPP			fPLC		gPLC	
	n	V _{1/2} up	V _{1/2} down	n	V _{1/2} up	n	V _{1/2} down
XI-VSP1	10	55 ± 2	97 ± 3	11	98 ± 5	11	77 ± 3
XI-VSP2	11	$\textbf{92}\pm\textbf{8}$	n/a	11	83 ± 6	9	118 ± 3

S1 Table



12 20 28 36 40

S1 Fig



S2 Fig