A Novel Homeostatic Mechanism Tunes PI(4,5)P₂-dependent Signaling at the Plasma Membrane

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One-Sentence Summary: The enzyme PIP4K functions as both a sensor and negative regulator of PI(4,5)P₂ synthesis by the closely related PIP5K enzymes, tuning the activity of numerous membrane functions.

Abstract: The lipid molecule phosphatidylinositol (4,5)-bisphosphate (PI(4,5)P₂) controls all aspects of plasma membrane (PM) function in animal cells, from its selective permeability to the attachment of the cytoskeleton. Although disruption of PI(4,5)P₂ is associated with a wide range of diseases, it remains unclear how cells sense and maintain PI(4,5)P₂ levels to support various cell functions. Here, we show that the PIP4K family of enzymes that synthesize PI(4,5)P₂ via a minor pathway, also function as sensors of tonic PI(4,5)P₂ levels, inactivating synthesis of the lipid via the major PIP5K-catalyzed pathway when PI(4,5)P₂ levels rise. Perturbation of this simple homeostatic mechanism reveals differential sensitivity of PI(4,5)P₂-dependent signaling to elevated PI(4,5)P₂ levels. These finding reveal that a subset of PI(4,5)P₂-driven functions may drive disease associated with disrupted PI(4,5)P₂ homeostasis.
The lipid molecule PI(4,5)P$_2$ is a master regulator of animal cell plasma membranes (PMs). By recruiting or activating scores of membrane proteins, it controls transport of ions and solutes across the membrane$^{1,2}$, attaches the underlying cytoskeleton$^3$, regulates the traffic of proteinaceous cargo to and from the membrane$^4$, disseminates extracellular signals$^2$, and facilitates the entry, assembly and egress of bacterial and viral pathogens$^{2,5}$. As a result, synthesis of PI(4,5)P$_2$ is essential for life in mammals$^6,7$. Nonetheless, genetic defects occur in humans that either increase or decrease PI(4,5)P$_2$ levels, disrupting cellular physiology in unpredictable ways. These manifest in diseases ranging from cancer$^8$ to kidney disease$^9$ to dysentery$^{10}$. Clearly, there is a central physiological imperative to tightly control PI(4,5)P$_2$ levels for harmonious PM function. A detailed homeostatic mechanism that can sense and maintain PI(4,5)P$_2$ levels, however, has proven elusive.

Most prior work in this area has focused on positive regulation of phosphatidylinositol 4-phosphate 5-kinases (PIP5Ks), the major enzyme responsible for PI(4,5)P$_2$ synthesis (fig. 1A). These enzymes add a phosphate to the 5-OH of its substrate, PI4P$^{11-13}$. However, we reasoned that maintaining tonic PI(4,5)P$_2$ levels in the PM in the presence of abundant PI4P substrate$^{14,15}$ would demand negative feedback. This is especially apparent during lipid re-synthesis after phospholipase C (PLC) activation; PI(4,5)P$_2$ levels plateau despite rapidly rising PI4P levels$^{16-18}$. To elucidate the negative feedback mechanism, we initially, independently, over-expressed all three isoforms of human PIP5K (A-C) or the single isoform from the budding yeast, *Saccharomyces cerevisiae* (Mss4). PI(4,5)P$_2$ levels were measured in live cells using a low-affinity biosensor, Tubby$^{c333H}$ $^{19}$. In all cases, PIP5K over-expression substantially increased PI(4,5)P$_2$ levels (fig. 1A). Surprisingly, catalytic activity of the human enzymes, but not yeast PIP5K, was dispensable for promoting an increase in PI(4,5)P$_2$ levels at the PM (fig. 1A). Conversely, over-expression of the related phosphatidylinositol 5-phosphate 4-kinases (PIP4Ks), which produce PI(4,5)P$_2$ from much less abundant PI5P substrate, actually decreased PI(4,5)P$_2$ levels slightly, also independent of catalytic activity (fig. 1B). These findings are consistent with a previous study reporting the negative regulation of PIP5K by PIP4K$^{20}$. We therefore reasoned that saturation of endogenous, inhibitory PIP4K molecules by PIP5K over-expression, regardless of catalytic activity, would free endogenous, active PIP5K enzyme from negative regulation (fig. 1C). In support of this, over-expression of all three PIP4K isoforms were able to attenuate the elevated PI(4,5)P$_2$ levels caused by PIP5K over-expression (fig. 1D).
To test real-time negative regulation of PIP5K by PIP4K, we triggered PM recruitment of cytosolic, FKBP-tagged PIP4K by chemically induced dimerization (CID) with a membrane targeted FRB domain, using rapamycin\textsuperscript{21}. As shown in fig. 1E, all three isoforms of PIP4K induced a steady decline in PM PI(4,5)P\textsubscript{2} levels within minutes of PM recruitment, independent of catalytic activity. To test for inhibition of PIP5K by PIP4K more directly, we tested activity of purified PIP5K1A on PI4P-containing supported lipid bilayers (SLBs). Addition of PIP4K exhibited delayed inhibition of PIP5K activity (fig. 1F). Once PI(4,5)P\textsubscript{2} reached approximately 28,000 lipids/\mu m\textsuperscript{2} (~2 mol %), PIP5K dependent lipid phosphorylation slowed down, which doubled the reaction completion time (fig. 1F, right). In contrast, we observed no PIP4K dependent inhibition of Mss4 (fig. 1F, inset). These data recapitulate the prior finding that PIP4K only inhibited purified PIP5K in the presence of bilayer-presented substrate\textsuperscript{20}. We therefore hypothesized that inhibition of PIP5K by PIP4K requires recruitment of the latter enzyme to the PM by PI(4,5)P\textsubscript{2} itself.

To probe the interaction of endogenous PIP4Ks with PM PI(4,5)P\textsubscript{2}, we used a split fluorescent protein approach\textsuperscript{22} to add a NeonGreen2 (NG2) tag to PIP4K2C, the most abundant PIP4K in HEK293 cells\textsuperscript{23,24}. Successful integration of the split NG2 tag was evident at both the genomic and protein levels (fig. 2A). As expected, endogenous NG2-PIP4K2C has a mainly cytosolic distribution when viewed in confocal, with a slight enrichment at the cell periphery (fig. 2B). Analysis of the ventral PM by total internal fluorescence microscopy (TIRFM) revealed individual, diffraction-limited and uniform intensity puncta that were dynamically associated with the membrane (fig. 2B). We compared the intensity of these puncta with a PI(4,5)P\textsubscript{2} biosensor tagged with single, tandem-dimer or -trimer mNeonGreen expressed at single molecule levels. This revealed that the NG2-PIP4K2C puncta contained an average of 1.64 NG2 molecules, consistent with dimeric PIP4K2C complexes\textsuperscript{25} containing either one or two NG2-tagged protein copies (fig. 2B).

Given the dynamic PM localization of NG2-PIP4K2C, is this PI(4,5)P\textsubscript{2} dependent? As acute depletion of PI(4,5)P\textsubscript{2} by CID of a PI(4,5)P\textsubscript{2} 5-OH phosphatase and a PM bait\textsuperscript{21} caused rapid depletion of both a high affinity PI(4,5)P\textsubscript{2} biosensor, Tubby\textsuperscript{19}, and endogenous PIP4K2C (fig. 2C), the answer was yes. A PI(4,5)P\textsubscript{2} phosphatase also rapidly depleted bound biosensor and purified PIP4K2A from supported membranes (fig. 2D). Since a relatively small fraction of PIP4K2C is present on the PM at steady state (see confocal images in fig. 2B), could elevated
PI(4,5)P₂ actually enhance recruitment of PIP4K2C? To answer this question, we used CID to acutely recruit a homodimeric mutant PIP5K domain²⁶ to the PM (fig. 2E). This caused rapid increases in PM levels of both PI(4,5)P₂ biosensor and NG2-PIP4K2C, in a manner that was dependent on catalytic activity of the PIP5K. Likewise, over-expression of active yeast Mss4 (but not its inactive mutant) increased PM recruitment of NG2-PIP4K2C (fig. 2F). In PI4P-containing SLBs, Mss4-induced PI(4,5)P₂ synthesis also increased PIP4K2C membrane binding in a highly cooperative manner: PIP4K2C was barely detectable below 2 mol % PI(4,5)P₂, but rose sharply thereafter (fig. 2G). Indeed, analysis of PIP4K2C equilibrium binding on PI(4,5)P₂-containing SLBs also revealed cooperative binding of PIP4K2C above ~2 mol % PI(4,5)P₂ (fig. 2H).

These data suggest that PIP4K2C binds PM PI(4,5)P₂ with relatively low affinity. To test this in live cells, we assessed the kinetics of PM binding during PI(4,5)P₂ re-synthesis after strong PLC activation. Stimulation of over-expressed PLC-coupled muscarinic M3 receptors induced rapid depletion of both NG2-PIP4K2C and PI(4,5)P₂ measured with Tubbyc (fig. 2I, top). Subsequent induction of PI(4,5)P₂ re-synthesis with the muscarinic antagonist atropine revealed much slower rebinding of NG2-PIP4K2C to the PM compared to the Tubbyc PI(4,5)P₂ biosensor; PIP4K2C takes almost twice as long (fig. 2I, bottom). Collectively, these data demonstrate that PIP4Ks are low-affinity PI(4,5)P₂ effectors, poised to sense both decreases and crucially, elevations in PI(4,5)P₂ levels in the PM.

The inhibition of PIP5K activity by PIP4Ks (fig. 1), and the low-affinity PI(4,5)P₂ binding by PIP4K (fig. 2) suggest the following mechanism: when PI(4,5)P₂ levels rise due to PIP5K activity, PIP4K is recruited to the PM, where it can directly bind and inhibit PIP5K. Indeed, whereas PIP5K PM binding is largely unaffected by PIP4K over-expression, all three isoforms of PIP4K are strongly recruited to the PM by co-expression of PIP5Ks (fig. 3A), as previously observed for PIP4K2A²⁷. While this is consistent with a direct interaction between PIP4Ks and PIP5Ks, another possibility exists: the PIP5K dependent increase in PI(4,5)P₂ (fig. 1A) enhances PM recruitment of PIP4K. Prior pull-downs of PIP5K and PIP4K from lysates required cross-linking the proteins when the enzymes may still have been co-localized on the PM²⁰. We therefore sought to distinguish between a direct PIP5K-PIP4K binding interaction versus PI(4,5)P₂-induced co-enrichment on membranes. To this end, we devised an experiment whereby a bait protein (either PIP5K or control proteins) could be acutely localized to domains
of the PM. This was achieved using CID of baits with an endoplasmic reticulum (ER) tethered protein, causing recruitment of the bait protein to ER-PM contact sites (fig. 3B). Enrichment of endogenous NG2-PIP4K2C at ER-PM contact sites was only observed when PIP5K1A was the bait; an unrelated peptide (myristoylated and palmitoylated peptide from Lyn kinase, Lyn11) or Mss4 did not enrich NG2-PIP4K2C (fig. 3B). The use of Mss4 ruled out an effect of enhanced PI(4,5)P$_2$ generation at contact sites, since this enzyme increases PI(4,5)P$_2$ as potently as PIP5K1A (fig. 1A), yet does not cause recruitment of PIP4K2C. Finally, we also demonstrate that PIP4K2C binding to PI(4,5)P$_2$-containing SLBs was greatly enhanced by addition of PIP5K to the membranes, but not by Mss4 (fig. 3C). Clearly, PIP4K enzymes directly interact with PIP5Ks on PI(4,5)P$_2$-containing lipid bilayers.

Synthesizing all of these observations, we propose a simple homeostatic feedback loop that maintains PI(4,5)P$_2$ levels in the PM (fig. 4A): when PI(4,5)P$_2$ levels increase, PIP4K is recruited to the PM in sufficient quantities to inhibit PIP5K, halting further PI(4,5)P$_2$ synthesis. If PI(4,5)P$_2$ levels fall, PIP4K is one of the first PI(4,5)P$_2$ binding proteins to be released (due to its low affinity), causing disinhibition of PIP5K and recovery of PI(4,5)P$_2$. We next sought to test how perturbations of this homeostat would affect physiological function. We could produce graded changes in resting PI(4,5)P$_2$ levels by over-expression of various components of the homeostat: enhanced PIP5K1A expression, either catalytically active or inactive, increases PI(4,5)P$_2$; a myristoylated PIP4K2A retains PM localization even at low PI(4,5)P$_2$, causing sustained reductions in PI(4,5)P$_2$; and a PM-localized PI(4,5)P$_2$ 5-OH phosphatase causes near complete ablation of the lipid. These constructs all show the expected changes in PM PI(4,5)P$_2$ compared to a control, reported by three different PI(4,5)P$_2$ biosensors. Of these, Tubbyc showed the most linear response across all changes in PI(4,5)P$_2$ levels (fig. 4B). We then used these graded changes in steady-state PM PI(4,5)P$_2$ to investigate the concentration requirements for the lipid.

PI(4,5)P$_2$ is the substrate for PLC, the enzyme that cleaves it into second messengers diacylglycerol and inositol (1,4,5)-trisphosphate (IP$_3$), triggering calcium release from ER stores (fig. 4C). Calcium release was indeed reduced by lower PI(4,5)P$_2$ levels, but appeared to be maximal at tonic PI(4,5)P$_2$ levels; it was unaffected by increased PM PI(4,5)P$_2$. This was true for both peak calcium release and total release from stores (assessed by measuring activity in calcium-free medium, fig. 4C). Influx of extracellular calcium was increased by elevated...
PI(4,5)P₂ levels (fig. 4C), consistent with a prior report that store-operated calcium entry is enhanced by increased PIP5K activity. However, IP₃-triggered calcium release appears saturated at resting PI(4,5)P₂. This strongly contrasts with the effects on the other PI(4,5)P₂ signaling pathway, class I phosphoinositide 3-OH kinase (PI3K). Epidermal growth factor (EGF) receptor stimulation activates PI3K, which converts a small fraction of PI(4,5)P₂ to PIP₃ (fig. 4D). Using a sensitive PIP₃ biosensor, we observed PIP₃ production changing proportionately with PI(4,5)P₂, never reaching a saturated level (fig. 4D). This also explains enhanced PI3K signaling reported in PIP4K-null cells. Intriguingly, PIP4Ks were reported to inhibit PI3K/Akt signaling two decades ago, but the mechanism was proposed to be through removal of its PI5P substrate, which was thought to somehow enhance accumulation of PI3K lipid products. The key evidence that it was PI5P that caused the PI3K lipid accumulation came from the observation that it could be recapitulated by the Shigella flexneri effector protein IpgD, which generates some PI5P from PI(4,5)P₂; this and the analogous Salmonella effector SopB both activate the PI3K/Akt pathway. However, it was recently shown that both SopB and IpgD are in fact novel phosphotransferases that directly convert PI(4,5)P₂ into the PI3K signaling lipid PI(3,4)P₂, explaining how these enzymes activate Akt.

In conclusion, our results reveal a remarkably simple homeostatic mechanism that controls PM PI(4,5)P₂ levels (fig. 4A). Perturbation of this homeostasis reveals different sensitivities of PLC and PI3K signaling, with the latter showing enhanced activation with elevated PI(4,5)P₂. This likely explains why the PI3K, and not PLC pathway, drives the phenotype of PIP4K-null fruit flies. More broadly, such differences in the sensitivity of PI(4,5)P₂-dependent PM functions to lipid concentration may go a long way to explaining the phenotypic diversity of diseases associated with dysregulated PI(4,5)P₂ metabolism. For example, they may explain why a selective inhibitor of PI3Kα can correct aberrant kidney function associated with Lowe syndrome models. Indeed, experimental manipulation of PI(4,5)P₂ homeostasis will now afford the ability to determine which of the panoply of PI(4,5)P₂-dependent PM functions are dysregulated by pathological alterations – finally bringing potential therapeutic targets into view.
References


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**Data and materials availability:** All data are available in the main text or the supplementary materials.

Supplementary Materials

**Materials and Methods**

*Cell culture and lipofection*

HeLa (ATCC CCL-2) and HEK293A (ThermoFisher R705-07) cells were cultured in DMEM (low glucose; Life Technologies 10567022) supplemented with 10% heat-inactivated fetal bovine serum (Life Technologies 10438-034), 100 units/ml penicillin, 100µg/ml streptomycin (Life Technologies 15140122), and 1:1,000 chemically defined lipid supplement (Life Technologies 11905031) at 37°C with a humidified atmosphere with 5% CO2. Cells were passaged twice per week diluting 1 in 5 after dissociation in TrpLE (Life Technologies 12604039). 293A cells with endogenous PIP4K2C alleles tagged with split NeonGreen2 (NG2) were generated similarly as described using a protocol we have described. In brief, Platinum Cas9 (Thermo Fisher B25640) was precomplexed with gRNA and electroporated into HEK293NG2-1-10 cells in combination with a single-stranded HDR Template (IDT). Sequences are provided in table 2. The HDR template contains 70 bp homology-arms, the NG2-11 sequence, and a flexible linker in frame with PIP4K2C (ATGACCGAGCTCAACTTCAAGGAGTGCGAAAAGGCCTTTACCAGATATGATGGGTGCGGGC). After recovery, FACS (University of Pittsburgh Flow Cytometry Core) was used to sort NG2-positive cells. These NG2-PIP4K2C cells were cultured under identical conditions to the HeLa and HEK293A cells.

*Chemicals and reagents*
Rapamycin (Thermo Fisher BP2963-1) was dissolved in DMSO at 1 mM and stored as a stock at -20°C, it was used in cells at 1 µM. EGTA (VWR EM-4100) was dissolved in water at 0.5 M and stored at room temperature, it was used in cells at 5 mM. EGF (Corning CB-40052) was dissolved in water at 100 µg/ml and stored as a stock at -20°C, it was used in cells at 10 ng/ml. Carbachol (Thermo Fisher AC10824-0050) was dissolved in water at 50 mM and stored as a stock at -20°C, it was used in cells at 100 µM. Atropine (Thermo Fisher AC226680100) was dissolved in 100% ethanol at 25 mM and stored as a stock at -20°C, it was used in cells at 5 µM.

**Plasmids and cloning**

The EGFP (Aequorea victoria GFP containing F64L and S65T mutations)\(^{35}\), mCherry (Discoma DsRed monomeric variant)\(^{36}\), mTagBFP2 (Entacmaea quadricolor protein eqFP578)\(^{37}\), IRFP713 (Rhodopseudomonas palustris [Rp] bacteriophytochrome BphP2)\(^{38}\) and iRFP670 (RpBphP6 iRFP702 containing V112I, K174M and I247C mutations)\(^{39}\) fluorophores were used in the Clontech pEGFP-C1, -C2, and -N1 backbones as described previously\(^{34}\). Mutated constructs were generated using site-directed mutagenesis using targeted pairs of DNA oligos which were custom made and supplied by Thermo Fisher. New plasmids used in this study were generated using standard restriction-ligation or by using NEBuilder HiFi DNA Assembly (New England Biolabs E552OS). HsPIP5K1A, HsPIP5K1B, Mss4_Kina, and HsPIP4K2C were obtained as human codon optimized synthetic gBlocks (IDT). Otherwise, plasmids were obtained from the sources listed in Table 1. All constructs were sequence verified using Sanger DNA sequencing. Plasmids constructed for this study are available through Addgene.

**Purification of PIP5K1A and Mss4**

Gene sequences encoding human PIP5K1A and yeast Mss4 kinase domain were cloned into a FastBac1 vector to create the following vectors: His6-MBP-TEV-(Gly)5-PIP5K1A (1-546aa) and His6-MBP-TEV-(Gly)5-Mss4 (379-779aa). BACMIDs and baculovirus were generated as previously described\(^{40}\). ES-Sf9 cells were infected with
baculovirus using an optimized multiplicity of infection (MOI), typically 2% vol/vol, that was empirically determined from small-scale test expression. Infected cells were typically grown for 48 hours at 27°C in ESF 921 Serum-Free Insect Cell Culture medium (Expression Systems, Cat# 96-001-01) and then harvested by centrifugation. Insect cell pellets were then washed with 1x PBS [pH 7.2] and centrifuged (3500 rpm for 10 minutes). The final cell pellet was combined with an equal volume of buffer containing 1x PBS [pH 7.2], 10% glycerol, and 2x Sigma protease inhibitor cocktail tablet solution before transferring to the -80°C freezer for storage. For purification, frozen cells were thawed in an ambient water bath and then resuspended in buffer containing 50 mM Na₂HPO₄ [pH 8.0], 10 mM imidazole, 400 mM NaCl, 5% glycerol, 1 mM PMSF, 5 mM BME, 100 µg/mL DNase, and 1x Sigma protease inhibitor cocktail tablet. Cells were lysed using a glass dounce homogenizer. Lysate was then centrifuged at 35,000 rpm (140,000 x g) for 60 minutes in a Beckman Ti-45 rotor at 4°C. High speed supernatant was combined with 6 mL of Ni-NTA Agarose (Qiagen, Cat# 30230) and stirred in a beaker for 1-2 hour(s) at 4°C. Following batch binding, resin was collected in 50 mL tubes, centrifuged, and washed with buffer containing 50 mM Na₂HPO₄ [pH 8.0], 10 mM imidazole, 400 mM NaCl, and 5 mM BME. Ni-NTA resin with His6-MBP-(Asn)10-TEV-(Gly)5-PIP5KA bound was washed in a gravity flow column with 100 mL of 50 mM Na₂HPO₄ [pH 8.0], 30 mM imidazole, 400 mM NaCl, 5% glycerol, and 5 mM BME buffer. Protein elution was achieved by washing the resin with buffer containing 50 mM Na₂HPO₄ [pH 8.0], 500 mM imidazole, 400 mM NaCl, 5% glycerol, and 5 mM BME. Peak fractions were pooled, combined with 200 µg/mL His6-TEV(S291V) protease, and dialyzed against 4 liters of buffer containing 20 mM Tris [pH 8.0], 200 mM NaCl and 2.5 mM BME for 16-18 hours at 4°C. The next day, dialysate was combined 1:1 volumes with 20 mM Tris [pH 8.0], 1 mM TCEP to reduce the NaCl to a final concentration of 100 mM. Precipitate was removed by centrifugation (3500 rpm for 10 minutes) and a 0.22 µm syringe filtration. Clarified dialysate was bound to a MonoS cation exchange column (GE Healthcare, Cat# 17-5168-01) equilibrated with buffer containing 20 mM Tris [pH 8.0], 100 mM NaCl, and 1 mM TCEP. Proteins were resolved over a 10-100% linear gradient (0.1-1 M NaCl, 45 CV, 45 mL total, 1 mL/min flow rate). (Gly)x₁₀-PIP5K1A and
(Gly)$_5$-Mss4 eluted from the MonoS in the presence of 375-450 mM NaCl. Peak fractions containing PIP5K1A were pooled, concentrated in a 30 kDa MWCO Vivaspin 6 centrifuge tube (GE Healthcare, Cat# 28-9323-17), and loaded onto a 24 mL Superdex 200 10/300 GL (GE Healthcare, Cat# 17-5174-01) size exclusion column equilibrated in 20 mM Tris [pH 8.0], 200 mM NaCl, 10% glycerol, 1 mM TCEP. Peak fractions were concentrated to 10-50 µM using a 30 kDa MWCO Amicon centrifuge tube (Millipore Sigma) before snap freezing with liquid nitrogen. PIP5K1A and Mss4 were stored in -80°C as single use aliquots.

**Purification of PIP4K2A**

The gene encoding human PIP4K2A was cloned into a pETM derived bacterial expression vector to create the following fusion protein: His6-SUMO3-(Gly)5-PIP4K2A (1-406aa). Recombinant PIP4KA was expressed in BL21 (DE3) Star E. coli (i.e. lack endonuclease for increased mRNA stability). Using 4 liters of Terrific Broth, bacterial cultures were grown at 37°C until OD$_{600}$=0.6. Cultures were then shifted to 18°C for 1 hour to cool down. Protein expression was induced with 50 µM IPTG and bacteria expressed protein for 20 hours at 18°C before being harvested by centrifugation. For purification, cells were lysed into buffer containing 50 mM Na$_2$HPO$_4$ [pH 8.0], 400 mM NaCl, 0.4 mM BME, 1 mM PMSF (add twice, 15 minutes intervals), DNase, and 1 mg/mL lysozyme using a microtip sonicator. Lysate was centrifuged at 16,000 rpm (35,172 x g) for 60 minutes in a Beckman JA-17 rotor chilled to 4°C. Lysate was circulated over 5 mL HiTrap Chelating column (GE Healthcare, Cat# 17-0409-01) that had been equilibrated with 100 mM CoCl$_2$ for 1 hour, washed with MilliQ water, and followed by buffer containing 50 mM Na$_2$HPO$_4$ [pH 8.0], 400 mM NaCl, 0.4 mM BME. Recombinant PIP4K2A was eluted with a linear gradient of imidazole (0-500 mM, 8 CV, 40 mL total, 2 mL/min flow rate). Peak fractions were pooled, combined with 50 µg/mL of His6-SenP2 (SUMO protease), and dialyzed against 4 liters of buffer containing 25 mM Na$_2$HPO$_4$ [pH 8.0], 400 mM NaCl, and 0.4 mM BME for 16-18 hours at 4°C. Following overnight cleavage of the SUMO3 tag, dialysate containing His6-SUMO3, His6-SenP2, and GGGGGG-PIP4K2A was recirculated for at least 1 hour over a 5 mL
HiTrap(Co²⁺) chelating column. Flow-through containing GGGGG-PIP4K2A was then concentrated in a 30 kDa MWCO Vivaspin 6 before loading onto a Superdex 200 size exclusion column equilibrated in 20 mM HEPES [pH 7], 200 mM NaCl, 10% glycerol, 1 mM TCEP. In some cases, cation exchange chromatography was used to increase the purity of GGGGG-PIP4K2A before loading on the Superdex 200. In those cases, we equilibrated a MonoS column with 20 mM HEPES [pH 7], 100 mM NaCl, 1 mM TCEP buffer. PIP4K2A (pI = 6.9) bound to the MonoS was resolved over a 10-100% linear gradient (0.1-1 M NaCl, 30 CV, 30 mL total, 1.5 mL/min flow rate). Peak fractions collected from the Superdex 200 were concentrated in a 30 kDa MWCO Amicon centrifuge tube and snap frozen at a final concentration of 20-80 µM using liquid nitrogen.

**Purification of PH-PLCδ1 domain**

The coding sequence of human PH-PLCδ1 (11-140aa) was expressed in BL21 (DE3) E. coli as a His6-SUMO3-(Gly)5-PLCδ1 (11-140aa) fusion protein. Bacteria were grown at 37°C in Terrific Broth to an OD₆₀₀ of 0.8. Cultures were shifted to 18°C for 1 hour, induced with 0.1 mM IPTG, and allowed to express protein for 20 hours at 18°C before being harvested. Cells were lysed into 50 mM Na₂HPO₄ [pH 8.0], 300 mM NaCl, 0.4 mM BME, 1 mM PMSF, 100 µg/mL DNase using a microfluidizer. Lysate was then centrifuged at 16,000 rpm (35,172 x g) for 60 minutes in a Beckman JA-17 rotor chilled to 4°C. Lysate was circulated over 5 mL HiTrap Chelating column (GE Healthcare, Cat# 17-0409-01) charged with 100 mM CoCl₂ for 1 hour. Bound protein was then eluted with a linear gradient of imidazole (0-500 mM, 8 CV, 40 mL total, 2 mL/min flow rate). Peak fractions were pooled, combined with SUMO protease (50 µg/mL final concentration), and dialyzed against 4 liters of buffer containing 50 mM Na₂HPO₄ [pH 8.0], 300 mM NaCl, and 0.4 mM BME for 16-18 hours at 4°C. Dialysate containing SUMO cleaved protein was recirculated for 1 hour over a 5 mL HiTrap Chelating column. Flow-through containing (Gly)⁵-PLCδ1 (11-140aa) was then concentrated in a 5 kDa MWCO Vivaspin 20 before being loaded on a Superdex 75 size exclusion column equilibrated in 20 mM Tris [pH 8.0], 200 mM NaCl, 10% glycerol, 1 mM TCEP. Peak fractions containing
(Gly)$_5$-PLCδ1 (11-140aa) were pooled and concentrated to a maximum of 75 µM (1.2 mg/mL) before freezing in liquid nitrogen.

**Purification of OCRL**

The coding sequence of human 5-phosphatase OCRL (234-539aa of 901aa isoform) was expressed in BL21 (DE3) E. coli as a His6-MBP-(Asn)10-TEV-(Gly)$_5$-OCRL fusion protein. Bacteria were grown at 37°C in Terrific Broth to an OD600 of 0.8. Cultures were shifted to 18°C for 1 hour, induced with 0.1 mM IPTG, and allowed to express protein for 20 hours at 18°C before being harvested. Cells were lysed into 50 mM Na$_2$PO$_4$ [pH 8.0], 300 mM NaCl, 0.4 mM BME, 1 mM PMSF, 100 µg/mL DNase using a microfluidizer. Lysate was then centrifuged at 16,000 rpm (35,172 x g) for 60 minutes in a Beckman JA-17 rotor chilled to 4°C. Lysate was circulated over 5 mL HiTrap Chelating column (GE Healthcare, Cat# 17-040901) charged with 100 mM CoCl$_2$ for 1 hour. Bound protein was eluted with a linear gradient of imidazole (0-500 mM, 8 CV, 40 mL total, 2 mL/min flow rate). Peak fractions were pooled, combined with TEV protease (75 µg/mL final concentration), and dialyzed against 4 liters of buffer containing 50 mM Na$_2$PO$_4$ [pH 8.0], 300 mM NaCl, and 0.4 mM BME for 16-18 hours at 4°C. Dialysate containing TEV protease cleaved protein was recirculated for 1 hour over a 5 mL HiTrap Chelating column. Flow-through containing (Gly)$_5$-protein was then concentrated in a 5 kDa MWCO Vivaspin 20 before being loaded on a Superdex 75 (10/300 GL) size exclusion column equilibrated in 20 mM Tris [pH 8.0], 200 mM NaCl, 10% glycerol, 1 mM TCEP. Peak fractions were pooled and concentrated before snap freezing in liquid nitrogen.

**Sortase mediated peptide ligation**

PIP4K2A, PIP5K1A, and PH-PLCδ1 were labeled on a N-terminal (Gly)$_5$ motif using sortase mediated peptide ligation$^{40,41}$. Initially, a LPETGG peptide was labeled with either Alexa488, Alexa647, or Cy5 conjugated to an amine reactive N-Hydroxysuccinimide (NHS) (e.g. NHS-Alexa488). Protein labeling was achieved by combining the fluorescently labeled LPETGG peptide with the following reagents: 50
17 mM Tris [pH 8.0], 150 mM NaCl, 50 µM (Gly)$_5$-protein, 500 µM Alexa488-LPETGG, and 10-15 µM His$_6$-Sortase. This reaction mixture was incubated at 16-18ºC for 16-20 hours, before buffer exchange with a G25 Sephadex column (e.g. PD10) to remove the majority of dye and dye-peptide. The His$_6$-Sortase was then captured on Ni-NTA agarose resin (Qiagen) and unbound labeled protein was separated from remaining fluorescent dye and peptide using a Superdex 75 or Superdex 200 size exclusion column (24 mL bed volume).

**Preparation of small unilamellar vesicles**

The following lipids were used to generated small unilamellar vesicles (SUVs): 1,2-dioleoyl-sn-glycero-3-phosphocholine (18:1 DOPC, Avanti # 850375C), L-α-phosphatidylinositol-4-phosphate (Brain PI(4)P, Avanti Cat# 840045X), L-α-phosphatidylinositol-4,5-bisphosphate (Brain PI(4,5)P$_2$, Avanti # 840046X), and 1,2-dioleoyl-sn-glycero-3-phospho-L-serine (18:1 DOPS, Avanti # 840035C). Lipids were purchased as single use ampules containing between 0.1-5 mg of lipids dissolved in chloroform. Brain PI(4)P and PI(4,5)P$_2$ were purchased as 0.25 mg/mL stocks dissolved in chloroform:methanol:water (20:9:1). To make liposomes, 2 µmoles total lipids were combined in a 35 mL glass round bottom flask containing 2 mL of chloroform. Lipids were dried to a thin film using rotary evaporation with the glass round-bottom flask submerged in a 42ºC water bath. After evaporating all the chloroform, the round bottom flask was flushed with nitrogen gas for at least 30 minutes. We resuspended the lipid film in 2 mL of PBS [pH 7.2], making a final concentration of 1 mM total lipids. All lipid mixtures expressed as percentages (e.g. 98% DOPC, 2% PI(4)P) are equivalent to molar fractions. For example, a 1 mM lipid mixture containing 98% DOPC and 2% PI(4)P is equivalent to 0.98 mM DOPC and 0.02 mM PI(4)P. To generate 30-50 nm SUVs, 1 mM total lipid mixtures were extruded through a 0.03 µm pore size 19 mm polycarbonate membrane (Avanti #610002) with filter supports (Avanti #610014) on both sides of the PC membrane. Hydrated lipids at a concentration of 1 mM were extruded through the PC membrane 11 times.
Preparation of supported lipid bilayers

Supported lipid bilayers were formed on 25x75 mm coverglass (IBIDI, #10812). Coverglass was first cleaned with 2% Hellmanex III (Fisher, Cat#14-385-864) heated to 60-70°C in a glass coplin jar and incubated for at least 30 minutes. We washed the coverglass extensively with MilliQ water and then etched with Pirahna solution (1:3, hydrogen peroxide:sulfuric acid) for 10-15 minutes the same day SLBs were formed. Etched coverglass, in water, was rapidly dried with nitrogen gas before adhering to a 6-well sticky-side chamber (IBIDI, Cat# 80608). SLBs were formed by flowing 30 nm SUVs diluted in PBS [pH 7.2] to a total lipid concentration of 0.25 mM. After 30 minutes, IBIDI chambers were washed with 5 mL of PBS [pH 7.2] to remove non-absorbed SUVs. Membrane defects were blocked for 15 minutes with a 1 mg/mL beta casein (Thermo FisherSci, Cat# 37528) diluted in 1x PBS [pH 7.4]. Before use as a blocking protein, frozen 10 mg/mL beta casein stocks were thawed, centrifuged for 30 minutes at 21,370 x g, and 0.22 µm syringe filtered. After blocking SLBs with beta casein, membranes were washed again with 1mL of PBS, followed by 1 mL of kinase buffer before TIRFM.

Microscopy

For all live-cell imaging experiments, cells were imaged in 1.6 mL of experiment specific imaging media. Base imaging media contained FluoroBrite DMEM (Life Technologies A1896702) supplemented with 25 mM HEPES (pH 7.4) and 1:1000 chemically defined lipid supplement (SF CHIM). Media was then further supplemented with either 10% fetal bovine serum (CHIM) or 0.1% BSA (0.1% BSA CHIM). Alternatively, Ca²⁺ free Ringer’s solution (Ca²⁺ Free) was used, containing 160 mM NaCl, 2.5 mM KCl, 1 mM MgCl₂, 8 mM glucose and 10 mM NaHEPES, pH 7.5. For treatments, 0.4 mL of experiment specific imaging media containing fivefold final concentration of compound was applied to the dish (or 0.5 ml for a second addition).

Confocal imaging was performed on a Nikon TiE A1R platform with acquisition in resonant mode with a 100x 1.45 NA plan-apochromatic objective. The signal-to-noise
ratio was improved by taking 8 or 16 frame averages. Excitation of fluorophores was accomplished using a dual fiber-coupled LUN-V laser launch with 405-nm (BFP), 488-nm (EGFP and NG2), 561-nm (mCherry), and 640-nm (iRFP) lines. Emission was collected on four separate photomultiplier tubes with blue (425-475 nm), green (500-550 nm), yellow/orange (570-620 nm), and far-red (663-737 nm) filters. Blue and yellow/orange channels were recorded concurrently, as were green and far-red. The confocal pinhole was defined as 1.2x the Airy disc size of the longest wave-length channel used in the experiment. Nikon Elements denoising software was used to further enhance the signal-to-noise ratio.

For TIRFM and single-molecule imaging (SMol), a separate Nikon TiE platform coupled with a Nikon TIRF illuminator arm and 100x 1.45 NA plan-apochromatic objective was used. Excitation of fluorophores was accomplished using an Oxxius L4C laser launch with 405-nm (BFP), 488-nm (EGFP and NG2), 561-nm (mCherry), and 638-nm (iRFP) lines. Emission was collected through dual-pass filters (Chroma) with blue (420-480 nm) and yellow/orange (570-620 nm) together, and green (505-550 nm) and far-red (650-850 nm) together. An ORCA-Fusion BT sCMOS camera (Hamamatsu) was used to capture images. For TIRFM, images were captured with 2x2 pixel binning. For SMol, the NG2 channel was excited with 20% power for 50 ms from the 488-nm laser in a 16x16 μm region of the PM. Images were registered in rolling shutter mode with 2x2 pixel binning with a 1.5x magnifier lens.

For all types of imaging, Nikon Elements software was used to acquire all images for all experiments and all data was saved with the ND2 file extension.

Membrane binding and lipid phosphorylation reactions reconstituted on supported lipid bilayers (SLBs) were visualized using an inverted Nikon Eclipse Ti2 microscope using a 100x Nikon (1.49 NA) oil immersion TIRF objective. TIRF microscopy images of SLBs were acquired using an iXion Life 897 EMCCD camera (Andor Technology Ltd., UK). Fluorescently labeled proteins were excited with either a 488 nm, 561 nm, or 637 nm diode laser (OBIS laser diode, Coherent Inc. Santa Clara, CA) controlled with a Vortran
laser drive with acousto-optic tunable filters (AOTF) control. The power output measured through the objective for single particle imaging was 1-2 mW. Excitation light was passed through the following dichroic filter cubes before illuminating the sample: (1) ZT488/647rpc and (2) ZT561rdc (ET575LP) (Semrock). Fluorescence emission was detected on the iXion Life 897 EMCCD camera position after a Nikon emission filter wheel housing the following emission filters: ET525/50M, ET600/50M, ET700/75M (Semrock). All experiments were performed at room temperature (23ºC). Microscope hardware was controlled by Nikon NIS elements.

**Image analysis**

Analysis of all images was accomplished using Fiji software\(^4^2\) using the LOCI BioFormats importer\(^4^3\). Custom macros were written to generate channel-specific montages displaying all x,y positions captured in an experiment in concatenated series. In these montages, individual regions of interest (ROIs) were generated around displayed cells.

For confocal images, the ratio or fluorescence intensity between specific compartments was analyzed as described previously\(^3^4\). In brief, a custom macro was used to generate a compartment of interest specific binary mask through à trous wavelet decomposition\(^4^4\). This mask was applied to measure the fluorescence intensity within the given compartment while normalizing to the mean pixel intensity in the ROI.

For TIRFM images, a minimum intensity projection was used to generate ROIs within the smallest footprint of the cells. Background fluorescence was measured and subtracted from all images at all timepoints. The average pixel intensity in each frame (\(F_t\)) was normalized to the mean pixel intensity in the ROI of the time points before treatment (\(F_{pre}\)) to yield \(F_t/F_{pre}\).

Quantitative data was imported into Prism 8 (GraphPad) for statistical analysis and the generation of graphs and plots. D'Agostino and Pearson normality tests showed data that significantly varied from normal distribution, data were then subjected to a
nonparametric Kruskal-Wallis test. If significant difference was found between sample medians, a post hoc Dunn's multiple comparison test was run.

Representative images were selected based on fluorescence measurements near the median of the sampled population, displayed typical morphology, and robust signal-to-noise ratio. If adjusting brightness or contrast, any changes were made across the entire image.

**Single Molecule Analysis using Thunderstorm**

Mean photon count was estimated using the Fiji ThunderSTORM plugin\(^45\). Either HEK293A cells expressing PH-PLC\(_\delta1\)-mNG2x1-3 or NG2-PIP4K2C cells were imaged using SMol settings. Raw images were run through Fiji using the ThunderSTORM plugin. Settings for molecule localization were determined using a wavelet filter with a local maximum method and integrated Gaussian point spread function. To determine fluorescence intensity per spot, histograms of photon counts, in each condition, were generated using a 5-photon bin size.

**Kinetic measurements of PI(4,5)P\(_2\) production**

The kinetics of PI(4)P phosphorylation was measured on SLBs formed in IBIDI chambers and visualized using TIRF microscopy as previously described\(^40\). Reaction buffer contained 20 mM HEPES [pH 7.0], 150 mM NaCl, 1 mM ATP, 5 mM MgCl\(_2\), 0.5 mM EGTA, 20 mM glucose, 200 \(\mu\)g/mL beta casein (ThermoScientific, Cat# 37528), 20 mM BME, 320 \(\mu\)g/mL glucose oxidase (Serva, #22780.01 Aspergillus niger), 50 \(\mu\)g/mL catalase (Sigma, #C40-100MG Bovine Liver), and 2 mM Trolox (UV treated, see methods below). Perishable reagents (i.e. glucose oxidase, catalase, and Trolox) were added 5-10 minutes before image acquisition. For all experiments, we monitored the change in PI(4)P or PI(4,5)P\(_2\) membrane density using solution concentrations of 20 nM Alexa647-DrrA(544-647) or 20 nM Alexa488-PLC\(_\delta1\), respectively.

**Tables S1 and S2**
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Table S1. Plasmids used in this study.

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Table S2. HDR and gRNA sequences for PIP4K2C.
Fig. 1. Reciprocal regulation of PM PI(4,5)P$_2$ levels by PIP5K and PIP4K. (A) PIP5Ks increase PM PI(4,5)P$_2$ independently of catalytic activity. Cartoon denotes the catalytic activity of PIP5K. Images show equatorial confocal sections of HeLa cells expressing the low affinity Tubby$_{C}^{R332H}$ PI(4,5)P$_2$ sensor (orange), co-transfected with EGFP-tagged catalytically active or dead PIP5K constructs (yeast Mss4 or mammalian A, B, or C isoforms), or EGFP alone as control. Increased PI(4,5)P$_2$ is apparent from increased Tubby$_{C}^{R332H}$ fluorescence in the PM. Box and whisker plots show the mean fluorescence intensity ratio (PM/cell) of the PI(4,5)P$_2$ sensor from >90 cells imaged across at least three independent experiments (boxes displaying median and interquartile range, whiskers representing 10-90% of data and “+” represents mean). (B) PIP4Ks decrease PM PI(4,5)P$_2$ independently of catalytic activity. Cartoon shows the catalytic activity of PIP4K. Images show PI(4,5)P$_2$ sensor in HeLa cells as in A, co-transfected with different PIP4K isoforms (A, B, C), catalytically dead PIP4K2A or a PI(4,5)P$_2$ 5-phosphatase (ptpase) (INPP5E). Box and whiskers from >90 cells imaged across at least three independent experiments as in A. (C) Proposed inhibition of ePIPK5 (endogenous PIP5K) by ePIP4K. With the overexpression of a fluorescently tagged version of PIP5K, regardless of catalytic activity, ePIP4K is sequestered. This relieves endogenous PIP5K from inhibition, increasing PI(4,5)P$_2$ levels. (D) PIP4Ks antagonize PIP5K-mediated PI(4,5)P$_2$ increases. HeLa cells expressing PI(4,5)P$_2$ indicator Tubby$_{C}^{R332H}$ (orange) were co-transfected with the indicated EGFP- or TagBFP2-tagged constructs. Images show confocal equatorial sections of representative cells. Box and whiskers from 90 cells imaged across at least three independent experiments, displayed as in A. (E) PIP4K recruitment acutely inhibits PM PI(4,5)P$_2$ levels. Cartoon schematics show the chemically induced dimerization (CID) system for FKBP-tagged PIP4K isoforms (A, B, C), which dimerize with the PM-anchored FRB-Lyn$_{11}$ upon the addition of rapamycin (rapa). HEK293A cells were transfected with FKBP-tagged proteins, the high affinity PI(4,5)P$_2$ indicator Tubby$_{C}$ and FRB-Lyn$_{11}$. During time-lapse confocal microscopy, cells were stimulated with 1 µM rapa as indicated. Graphs represent mean change in PI(4,5)P$_2$ sensor intensity ratio (PM/cell) ± s.e. for 35-60 cells imaged across three independent experiments. (F) PIP4K2A attenuates the kinetics of PI(4,5)P$_2$ production driven by PIP5K1A, but not Mss4. Kinetics of PI(4,5)P$_2$ production measured on SLBs in the presence of 1 nM PIP5K1A, 20 nM PH-PLC$_{δ1}$, +/- 50 nM PIP4K2A. Inhibition of PIP5K1A activity is delayed until a threshold density of approximately 2% PI(4,5)P$_2$ is created to support membrane recruitment of PIP4K2A. Inset shows kinetics of reactions executed in the presence of 50 nM Mss4, 20 nM PH-PLC$_{δ1}$, +/- 50 nM PIP4K2A. Initial membrane composition: 76% DOPC, 20% DOPS, 4% PI(4)P. Right graphs show the quantification of time required for reactions to reach 95% completion (n = 3 technical replicates).
Fig. 2. PI(4,5)P2 is necessary and sufficient for the PM localization of PIP4K2C. (A) Endogenous tagging of PIP4K2C. Brief cartoon schematic showing the mechanism of endogenous tagging employed for PIP4K2C with NeonGreen2 (NG2). The resulting cell line was termed NG2-PIP4K2C. Cells were genotyped with a mNG specific forward primer and a PIP4K2C specific reverse primer yielding an edited product of ~200bp. Cells were also probed with a PIP4K2C specific antibody showing the expected ~3 kDa shift in weight (arrowhead). (B) Image based characterization of NG2-PIP4K2C. Confocal images display the NG2-PIP4K2C (blue) in cells localized mainly to the cytosol, but slight association of the enzyme to the PM can be seen in the zoomed image. When imaged live by TIRF, dynamic, diffraction limited spots are observed on the membrane (compare differential localization at 0 and +0.3 s). These have an intensity consistent with a mixed population of 1 or 2 mNG molecules when calibrated against single, dimeric or trimeric mNG molecules fused to a PI(4,5)P2 binding domain (graph). This correlates to the mean photon count of a heterogeneously tagged cell population (one or two alleles tagged with NG2111 and dimeric PIP4K2C). (C) Depletion of PI(4,5)P2 causes NG2-PIP4K2C to dissociate from the membrane. Cartoons show the CID system, in TIRF, for FKBP-tagged INPP5E (catalytically active or dead) dimerizing with the PM-anchored Lyn1111-FRB. NG2-PIP4K2C (blue) cells were transfected with FKBP-tagged proteins, the high affinity PI(4,5)P2 indicator TubbyC (orange) and Lyn1111-FRB. During time-lapse TIRF microscopy, cells were stimulated with 1µM rapa, as indicated. Traces represent mean change in fluorescence intensity (F/F₀). (D) Depletion of PI(4,5)P2 causes PIP4K2A to dissociate from SLBs. Imaging chambers containing 50 nM PIP4K2A and 20 nM PH-PLCδ1 at equilibrium with SLBs composed of 96% DOPC and 4% PI(4,5)P2 were visualized by TIRF microscopy. At 30 seconds, 100 nM OCRL was added to catalyze the dephosphorylation of PI(4,5)P2 and membrane dissociation of PIP4K2A and PH-PLCδ1. (E) Acute enrichment of PI(4,5)P2 causes PIP4K2C to increase association with the membrane. Cartoons show the CID system, in confocal, for the interaction of catalytically active or dead FKBP-tagged homo-dimeric PIP5K1C kinase with the PM-anchored Lyn1111-FRB. NG2-PIP4K2C (blue) cells were transfected with FKBP-tagged proteins, the low affinity PI(4,5)P2 indicator TubbyC RS323H (orange) and Lyn1111-FRB. During time-lapse confocal microscopy, cells were stimulated with 1µM rapa, as indicated. Traces represent mean change in fluorescence intensity (change in PM/cell ratio from pre-stimulation levels) ± s.e. of 48-52 cells imaged across at least three independent experiments. (F) Chronic enrichment of PI(4,5)P2 causes NG2-PIP4K2C to associate with the membrane. Cartoons show the expression of catalytically active or dead Mss4. Images show equatorial confocal sections of representative NG2-PIP4K2C cells transfected with Mss4 and the low affinity PI(4,5)P2 indicator TubbyC RS323H. Box and whisker plots show the mean fluorescence intensity ratio (PM/cell) of the PI(4,5)P2 sensor from 88-90 cells images across at least three independent experiments (boxes display median and interquartile range, whiskers represent 10-90% of data and *+ represents mean). (G) Enrichment of PI(4,5)P2 causes dynamic membrane recruitment of purified PIP4K2A. In SLBs, membrane recruitment of 50 nM PIP4K2A monitored during Mss4 catalyzed phosphorylation of PI(4)P. Membranes containing 4% PI(4)P were converted to PI(4,5)P2 using 10 nM Mss4. (H) Purified PIP4K2A localizes to PI(4,5)P2 in a concentration dependent manner. Membrane absorption and equilibration kinetics of 50 nM Alexa488-PIP4K2A measured by TIRF microscopy on SLBs containing 1–4% PI(4,5)P2. PIP4K2A membrane binding exhibited non-linearity with respect to the PI(4,5)P2 lipid density. Quantification of the fold increase in membrane bound PIP4K2A relative to the equilibrium fluorescence intensity of PIP4K2A on a membrane containing 0% PI(4,5)P2. (I) PM localization of PIP4K2C follows resynthesis of PI(4,5)P2. Cartoons show PLC83 mediated loss of PI(4,5)P2 and eNG-PIP4K2C followed by the subsequent reappearance of PI(4,5)P2 and eNG-PIP4K2C. eNG-PIP4K2C (blue) cells were transfected with FKBP-tagged proteins, the high affinity PI(4,5)P2 indicator TubbyC (orange) and Lyn1111-FRB. During time-lapse TIRF microscopy, cells were stimulated with 100 µM carbachol (CCh) and then stimulated with 5 µM atropine as indicated. TubbyC traces represent mean change in fluorescence intensity (F/F₀) ± s.e. of 36-37 cells that were imaged across three independent experiments. The eNG-PIP4K traces represent the mean change in puncta per µm² ± s.e. 40 cells were imaged across at least three independent experiments.
Fig. 3. PIP4K interacts with mammalian PIP5K for inhibition. (A) PIP5K expression increases PIP4K PM localization. The same experimental data set from Figure 1E is used here. HeLa cells expressing PIP5K (green) or PIP4K (blue) were co-transfected with the indicated EGFP- or TagBFP2-tagged isoform constructs. Images show equatorial sections in confocal of representative cells. For box and whisker plots, boxes display median and interquartile range and whiskers representing 10-90% of the data and “+” represents the mean of 90 cells imaged across at least three independent experiments. (B) PIP4K2C interacts with PIP5K1A. Cartoon schematic show the CIDs system for the generation of ER-PM contact sites between ER-anchored FKBP-CyB5 and PM-anchored FRB-tagged constructs. eNG-PIP4K2C (cyan) cells were transfected with FKBP-CyB5, the low affinity PI(4,5)P₂ indicator TubbyΔR332H, and the indicated FRB-tagged construct (magenta). During time-lapse TIRF microscopy, cells were stimulated with 1µM rapa. TIRF images are representative and color-coded to represent fluorescence intensity, as indicated. eNG-PIP4K2C traces represent mean fluorescence intensities (ER:PM/PM) ± s.e. of 32-39 cells imaged across a minimum of three independent experiments. (C) Dynamic PIP5K1A dependent membrane recruitment of PIP4K2A. In the absence of PIP5K, 50 nM PIP4K2A displays a low level of membrane recruitment. The addition of 10 nM PIP5K1A, stimuliates an immediate and steady increase in both PIP4K2A and PIP5K1A membrane localization. Membrane composition: 2% PI(4,5)P₂, 98% DOPC. TIRF images are representative and color-coded to represent fluorescence intensity, as indicated, scale bar is 5 µm. (D) Membrane binding of PIP4K2A is insensitive to yeast Mss4 membrane localization. TIRF microscopy images show the membrane localization of PIP4K2A in the absence and presence of Mss4. Following membrane equilibration of 50 nM PIP4K2A, 10 nM Mss4 was added to the imaging chamber. No appreciable change in PIP4K2A localization was observed during membrane absorption of Mss4. Membrane composition: 4% PI(4,5)P₂ and 96% DOPC. TIRF images are representative and color-coded to represent fluorescence intensity, as indicated, scale bar is 5 µm.
Fig. 4. PI3K, but not calcium signaling, are modulated across all concentration ranges of PI(4,5)P$_2$. (A) Proposed regulation of PIP5K by the low affinity PI(4,5)P$_2$ interaction of PIP4K. The working model for negative feedback of PIP5K via PIP4K resembles the thermostat regulation of temperature. When PI(4,5)P$_2$ levels are high, PIP4K is recruited and held at the PM, via a direct low affinity interaction with PI(4,5)P$_2$. At the PM, PIP4K interacts with and inhibits the catalytic activity of PIP5K, causing reduced PI(4,5)P$_2$ synthesis. (B) PI(4,5)P$_2$ biosensors detect a gradient of lipid levels. HEK293Aa cells were transfected with the indicated fluorescently tagged PI(4,5)P$_2$ modulating proteins (INPP5E, myrPIP4K2A, myrLyn, PIP5K1A catalytic dead or active) and the indicated PI(4,5)P$_2$ biosensor (PH-PLCδ1, Tubby, or TubbyR332H, displayed in orange) for 16-24 hours. Mean fluorescence intensity (PM/cyto) are shown as points with error bars representing s.e. of >120 cells imaged across three independent experiments. (C) PLC-mediated Ca$^{2+}$ signals saturate at tonic PI(4,5)P$_2$ levels. Cartoon schematics of PLC mediated Ca$^{2+}$ signaling and detection. HEK293A cells were transfected with the indicated fluorescently tagged construct and the calcium sensor R-GECO (purple). During time-lapse confocal microscopy (performed with either compete imaging media containing 1.8 mM Ca$^{2+}$ [Ca$^{2+}$] or calcium free Ringer’s media [Ca$^{2+}$-Free]), cells were stimulated with 100 µM CCh as indicated. Traces represent the peak response of mean change in fluorescence intensity (F/F$_{0}$) ± s.e. of >100 cells imaged across a minimum of three independent experiments. The peak response and total area under the curve (AUC) were plotted against the normalized ratio of Tubby$_3$. (D) PI3K mediated PI(3,4,5)P$_3$ synthesis is linearly dependent on PI(4,5)P$_2$ levels. Cartoon schematics show PI3K mediated signaling and detection of PI(3,4,5)P$_3$ upon the addition of EGF. HEK293A cells were transfected with the indicated fluorescently tagged construct and the PI(3,4,5)P$_3$ biosensor, PH-ARNO$^{2G}$-I103Ex2 (aPHx2) (magenta). During time-lapse confocal microscopy, cells were stimulated with 10 ng/mL EGF, as indicated. Traces represent the peak response of mean change in fluorescence intensity (change in PM/cell from pre-stimulation levels) ± s.e. of 35 cells imaged across a minimum of three independent experiments. The peak response and AUC were plotted against the normalized ratio of Tubby$_3$. 

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