Phosphatidylinositol 3,5-Bisphosphate regulates Ca²⁺ Transport During Yeast Vacuolar Fusion through Activation of the Ca²⁺ ATPase Pmc1

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Running title: PI(3,5)P₂ and Vacuolar Ca²⁺ Flux

Abbreviations

C8, dioctanoyl; PA, phosphatidic acid; PI3P, phosphatidylinositol 3-phosphate; $PI(3,5)P_2$, phosphatidylinositol 3,5-bisphospahte; SNARE, soluble N-ethylmaleimide-sensitive factor attachment protein receptor.

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Summary: During osmotic stress $PI(3,5)P_2$ triggers Ca^{2+} release from vacuoles. Here we show $PI(3,5)P_2$ stimulates Ca^{2+} uptake by vacuoles during fusion, illustrating that it has a dual role in Ca^{2+} transport.

ABSTRACT

The transport of Ca^{2+} across membranes precedes the fusion and fission of various lipid bilayers. Yeast vacuoles during hyperosmotic shock become fragmented through fission events that require Ca^{2+} efflux of their luminal stores through the TRP channel Yvc1. This requires the production of the lipid $PI(3,5)P_2$ by Fab1. Ca^{2+} is also released during vacuole fusion upon trans-SNARE complex assembly, however, the role of $PI(3,5)P_2$ remains unclear. Here we demonstrate that elevated $PI(3,5)P_2$ levels abolish Ca^{2+} efflux during fusion, indicating that $PI(3,5)P_2$ has opposing effects on Ca^{2+} transport in fission versus fusion. Notably, Ca^{2+} efflux was enhanced when $PI(3,5)P_2$ levels were reduced. Importantly, the effect of $PI(3,5)P_2$ on Ca^{2+} flux was independent of Yvc1. Rather, the effect was dependent on the Ca^{2+} pump Pmc1. Vacuoles lacking Pmc1 were resistant to the effects of $PI(3,5)P_2$, while those lacking Yvc1 remained sensitive. Furthermore altering $PI(3,5)P_2$ levels affects the interactions of Pmc1 with the V_0 component Vph1 and the R-SNARE Nyv1. We now propose a model in which elevated $PI(3,5)P_2$ activates continued Pmc1 function to prevent the accumulation of released extraluminal Ca^{2+} .

INTRODUCTION

The role of Ca²⁺ as a regulator of membrane fusion is best understood in the release of neurotransmitters through the fusion of synaptic vesicles with the presynaptic plasma membrane (Südhof, 2012). Upon plasma membrane depolarization an influx of Ca²⁺ into the cytoplasm occurs which subsequently binds to the C2 domains of synaptotagmin-1. Once bound to Ca²⁺, synaptotagmin-1 changes conformation and aids in triggering SNARE-mediated fusion (Chapman, 2008). Ca²⁺ transport across membranes is also a part of endolysosomal trafficking. In *Saccharomyces cerevisiae* vacuolar lysosomes serve as the major Ca²⁺ reservoir where it is complexed with inorganic polyphosphate (Dunn et al., 1994). The transport of Ca²⁺ into the vacuole lumen is mediated through the Ca²⁺/H⁺ exchanger Vcx1, and the ATP driven pump Pmc1, whereas the movement out of the lumen into the cytoplasm is driven by the mucolipin transient receptor potential cation channel (TRPML) family orthologue Ca²⁺ channel Yvc1 (Cunningham, 2011).

Although incomplete, we have a partial understanding of the mechanisms driving the release of Ca^{2+} during intracellular membrane trafficking. During osmotic shock or mechanical stress Yvc1 transports Ca^{2+} from the vacuole lumen to the cytosol that is controlled in part by the formation of $PI(3,5)P_2$ by the PI3P 5-kinase Fab1/PIKfyve (Bonangelino et al., 2002; Denis and Cyert, 2002; Dong et al., 2010; Su et al., 2011). Through unresolved mechanisms this cascade leads to the fragmentation of the vacuole into small vesicles through fission events.

During membrane fusion the release of Ca^{2+} is triggered by *trans*-SNARE pairing (Merz and Wickner, 2004). However, a role for $PI(3,5)P_2$ for fusion related Ca^{2+} efflux has not been tested. Separately, we showed that vacuole fusion was inhibited in the presence of elevated $PI(3,5)P_2$ (Miner et al., 2019). Fusion was blocked by either the addition of an exogenous dioctanoyl form of the lipid (C8- $PI(3,5)P_2$) or through endogenous overproduction by the hyperactive $fab1^{T2250A}$ mutant. Fusion was arrested after the formation of trans-SNARE pairs but

before mixing of the outer leaflets, i.e. hemifusion. The possible role for the TRP Ca^{2+} channel Yvc1 in the inhibition of fusion by $PI(3,5)P_2$ was eliminated, as vacuoles lacking the transporter fused well and were equally sensitive to $PI(3,5)P_2$ relative to wild type vacuoles. Although Yvc1 was not involved in the arrest of fusion, others have shown that Ca^{2+} efflux still occurs in the absence of the channel (Merz and Wickner, 2004). Thus, we postulated that $PI(3,5)P_2$ could still affect Ca^{2+} transport through another mechanism.

Various mechanisms control the involvement of Ca²⁺ transport in membrane fusion including Ca²⁺-binding protein calmodulin and a specific ABC transporter. Calmodulin acts as a sensor and is participates in triggering late fusion events. Calmodulin tightly binding vacuoles upon Ca²⁺ release signals the completion of docking, which stimulates bilayer mixing (Ostrowicz et al., 2008; Peters and Mayer, 1998). Another mechanism that regulates fusion events through modulating Ca²⁺ transport involves the vacuolar Class-C ABC transporters Ybt1 (Sasser et al., 2012b; Sasser and Fratti, 2014).

Here we continued our study of $PI(3,5)P_2$ during fusion and found that elevated concentrations of the lipid blocked the release of Ca^{2+} . Both exogenous $C8-PI(3,5)P_2$ and overproduction by $fab1^{T2250A}$ inhibited Ca^{2+} efflux. In contrast, Ca^{2+} efflux was enhanced when Fab1 was inhibited by Apilimod, or when $PI(3,5)P_2$ was sequestered by ML1-N. This effect was reproduced by the kinase dead $fab1^{EEE}$ mutation. Finally, we found that the effect of $PI(3,5)P_2$ on Ca^{2+} transport occurs through its action on the Ca^{2+} uptake pump Pmc1 and not Yvc1. Thus, we conclude that the observed Ca^{2+} efflux during vacuole homotypic fusion is due in part to the regulation of Ca^{2+} influx by Pmc1.

RESULTS

Ca²⁺ Efflux is inhibited by PI(3,5)P₂

The release of vacuole luminal Ca^{2+} stores occurs upon *trans*-SNARE complex formation in the path towards fusion (Merz and Wickner, 2004). The formation of vacuolar SNARE complexes also relies on the assembly of membrane microdomains enriched in regulatory lipids that include phosphoinositides such as the Fab1 substrate PI3P (Fratti et al., 2004). However, the link between phosphoinositides and Ca^{2+} transport during fusion has remained unclear. By contrast, the dependence of Ca^{2+} efflux on $PI(3,5)P_2$ during vacuole fission under hyperosmotic conditions has been established (Dong et al., 2010). Together, this served as the impetus to explore the role of $PI(3,5)P_2$ in fusion-linked Ca^{2+} transport.

To address this gap in knowledge, we used isolated vacuoles for *in vitro* assays of Ca²⁺ flux. Ca²⁺ transport was monitored through changes in Fluo-4 (or Cal-520) fluorescence as previously demonstrated (Miner et al., 2016; Miner et al., 2017; Miner and Fratti, 2019; Sasser et al., 2012b). Vacuoles were incubated under fusion conditions and reactions were started by the addition of ATP. As seen before, the addition of ATP triggers the rapid uptake of extraluminal Ca²⁺ from the buffer into the vacuole lumen as shown by the decrease in fluorescence (Fig. 1A, black line). Omission of ATP prevents Ca²⁺ uptake and fluorescence remained stable (not shown). Shortly after, and in a SNARE-dependent manner, Ca²⁺ is

released from vacuoles as shown by an increase in fluorescence. As a negative control for SNARE-dependent efflux, select reactions were treated with antibody against Sec17 (α -SNAP), a co-chaperone of Sec18 (NSF) that aids in disrupting inactive *cis*-SNARE complexes to initiate the fusion pathway (Fig. 1A, dotted line) (Mayer et al., 1996).

Here we tested the effects of PI(3,5)P₂ on Ca²⁺ transport. We treated reactions with dioctanoyl (C8) derivatives of PI(3,5)P2, as well as PI3P and phosphatidic acid (PA). PA was included due to its inhibitory effect on vacuole fusion (Starr et al., 2016), while PI3P was included as a precursor to PI(3,5)P₂ and as a pro-fusion lipid (Boeddinghaus et al., 2002; Karunakaran et al., 2012; Karunakaran and Fratti, 2013; Lawrence et al., 2014; Miner et al., 2016). Short chain lipids were used to avoid solubility issues. As a result, seemingly high concentrations of C8 lipids were needed to see an effect. This, however, does not reflect the mol% of the lipids that partition into the membrane bilayer. Collins and Gordon showed that only a small fraction of C8-lipids incorporate into the membrane (Collins and Gordon, 2013). Based on their work we estimate that 100 µM C8-PI(3,5)P₂ correlates to ~1.2 mol% partitioning to the membrane fraction, which is in keeping with what reconstituted proteoliposome studies use in fusion models (Mima and Wickner, 2009). Lipids were added after 10 min of incubation to allow uniform uptake of Ca²⁺. We found that C8-PI(3,5)P₂ blocked Ca²⁺ efflux in a dose dependent manner and as effectively as the α -Sec17 IgG negative control (Fig. 1A, D, red lines). In parallel we found that C8-PI3P only showed a modest inhibition at the highest concentration tested when compared to C8-PI(3,5)P2 (Fig. 1B, D). When C8-PA was tested we found that it did not significantly affect Ca2+ transport (Fig. 1C, D). These data indicate that the Ca2+ efflux associated with vacuole fusion was blocked by PI(3,5)P2. This is in contrast to the stimulatory effect of PI(3,5)P₂ on Ca²⁺ efflux associated with vacuole fission. Importantly the concentrations used do not fully inhibit fusion, suggesting that Ca2+ transport can be uncoupled from fusion by altering the lipid composition of vacuole (Miner et al., 2019).

Inhibition of the PI3P 5-Kinase Fab1/PIKfyve leads to enhanced Ca²⁺ Efflux.

To verify the effect of $PI(3,5)P_2$ on Ca^{2^+} efflux we sought to counter its effects through inhibiting its production. The small molecule Apilimod has been shown to inhibit mammalian PIKfyve (Cai et al., 2013; Dayam et al., 2015). Thus, we first tested if Apilimod could inhibit yeast Fab1 activity. For this we measured the conversion of BODIPY-TMR C6-PI3P to BODIPY-TMR C6-PI(3,5)P₂ by isolated vacuoles. Before we tested the effects of Apilimod, we verified our detection system by using $fig4\Delta$ and $fab1\Delta$ vacuoles that reduce or abolish PI(3,5)P₂ production, respectively. In Figure 2A-B, we show that wild type vacuoles phosphorylated BODIPY-TMR C6-PI3P to make BODIPY-TMR C6-PI(3,5)P₂. The half-life of the product was observed to be short as the levels of BODIPY-TMR C6-PI(3,5)P₂ was significantly reduced after 5 min of incubation. This was in keeping reports by others showing that PI(3,5)P₂ has a short half-life during vacuole fission (Duex et al., 2006a; Duex et al., 2006b; Jin et al., 2008). Next, vacuoles were incubated in the presence of BODIPY-TMR C6-PI3P and treated with either vehicle or Apilimod to inhibit Fab1 activity. Indeed, Apilimod treatment abolished BODIPY-TMR C6-PI(3,5)P₂ generation (Fig. 2C-D).

To visualize the effect of Apilimod on intact vacuoles we performed docking assays where endogenously produced $PI(3,5)P_2$ was labeled with Cy3-ML1-N as described previously (Miner et al., 2019). ML1-N is an N-terminal polypeptide from the endolysosomal mucolipin transient receptor potential (TRPML) Ca^{2+} channel (Dong et al., 2010). ML1-N preferentially binds to $PI(3,5)P_2$ on isolated yeast vacuoles as no labeling is observed with vacuoles from the kinase inactive $fab1^{EEE}$ mutant (Miner et al., 2019). In Figure 2E we show that vacuoles treated with carrier (DMSO) showed distinct $PI(3,5)P_2$ punctate staining as previously reported. When vacuoles were treated with Apilimod, we found that Cy3-ML1-N puncta were eliminated, further illustrating that Fab1 activity was inhibited during the experiment. Some free dye accumulated in the vacuole lumen to give the background fluorescence.

Apilimod alters Ca²⁺ transport

The effects of Apilimod were next tested on Ca^{2^+} efflux. We added a dose response curve of Apilimod after Ca^{2^+} was taken up by vacuoles. The addition of Apilimod enhanced the release of Ca^{2^+} in a dose dependent manner (Fig. 3A-B). This suggests that blocking Fab1 production of $PI(3,5)P_2$ promotes Ca^{2^+} efflux, which compliments the inhibition of Ca^{2^+} release through the addition of C8- $PI(3,5)P_2$ shown in Figure 1. The DMSO carrier control had no effect as shown by the overlapping curve with buffer control. It is important to note that pre-existing $PI(3,5)P_2$ remained on the vacuole upon Apilimod treatment. To compare the effects of new lipid production with the starting pool of $PI(3,5)P_2$ we tested whether physically blocking $PI(3,5)P_2$ with ML1-N would reproduce the effects seen with Apilimod. When ML1-N was added we found that efflux was increased, which partially reproduced the Apilimod affect (Fig. 3C-D). However, higher concentrations ($\geq 1~\mu M$) started to reduce efflux. Unlike Apilimod, high levels of ML1-N was expected to sequester any unbound $PI(3,5)P_2$. This is in agreement with the notion that low levels of $PI(3,5)P_2$ are needed for the fusion pathway to progress, while its absence or excess have deleterious effects on the system (Miner et al., 2019).

Fab1 kinase activity mutants differentially affect Ca2+ transport

Thus far we have used exogenous C8-PI(3,5)P2, chemical inhibition of Fab1, and lipid sequestration to show that Ca2+ efflux is affected by this lipid. To further confirm these findings, we next used vacuoles from yeast that expressed Fab1 mutations that altered kinase activity. First, we used vacuoles that contained the hyperactive Fab1 kinase mutation T2250A (Lang et al., 2017). We found that fab1^{T2250A} vacuoles showed attenuated Ca²⁺ release early in the assay which was followed by delayed release that reached wild type levels (Fig. 4A, red). We propose that the delay in Ca²⁺ release by fab1^{T2250A} vacuoles could be due to an increased time required for Fig4 to consume the elevated PI(3,5)P₂ present on these organelles. This is consistent with the effects of adding exogenous C8-PI(3,5)P2 in Figure 1A and supports a model in which elevated PI(3,5)P2 concentrations suppress Ca2+ efflux. The delay in efflux seen with low concentrations of C8-PI(3,5)P2 was close to what we observed with fab1 vacuoles and suggests that the prolonged inhibition by higher lipid concentrations was due to overwhelming the capacity of Fig4 to consume PI(3,5)P2 in the duration of the experiment. To verify that the decreased Ca2+ seen with fab1T2250A vacuoles was due to directly to changes in PI(3,5)P2 concentrations, we added ML1-N to sequester "excess" lipid and increase Ca2+ efflux. We found that reducing free PI(3,5)P₂ through sequestration partially restored Ca²⁺ efflux by fab1^{T2250A}

vacuoles (Fig. 4B). Although the rescue was incomplete, the trend is consistent with the notion that elevated $PI(3,5)P_2$ reduces Ca^{2+} efflux. ML1-N at concentrations at or above 500 nM reduced Ca^{2+} as seen in Figure 3. In parallel we found that Apilimod restored Ca^{2+} efflux to wild type level and continued to linearly elevated Ca^{2+} release in a dose-dependent manner (Fig. 4C)

To better our sense of how PI(3,5)P₂ concentrations serve as switch to affect Ca²⁺ transport, we used vacuole from yeast that expressed the kinase dead *fab1*^{EEE} mutation (Li et al., 2014). In Figure 4D, we show that *fab1*^{EEE} showed a delay and overall reduction in Ca²⁺ uptake compared to wild type. The eventual release of Ca²⁺ was also delayed, yet robust. Together the altered uptake and release of Ca²⁺ by *fab1*^{EEE} vacuoles reproduces the effects of adding Apilimod. To see if the effects of *fab1*^{EEE} were due to the lack of PI(3,5)P₂, we added exogenous C8-PI(3,5)P₂ and found that the Ca²⁺ efflux of *fab1*^{EEE} vacuoles was shifted to wild type levels (Fig. 4E). In contrast, our previous work showed PI(3,5)P₂ was unable to restore the fusion defect of *fab1*^{EEE} due to vacuolar maturation defects (Miner et al., 2019). This suggests that the regulation of Ca²⁺ efflux by PI(3,5)P₂ is distinct from its role in vacuolar maturation. Collectively with the effect of overproducing PI(3,5)P₂, these data demonstrate that PI(3,5)P₂ can serve as a rheostat to either inhibit or enhance Ca²⁺ release from vacuoles. Furthermore, the effects of PI(3,5)P₂ on Ca²⁺ release appear to be oppositely regulated during osmotic shock versus isotonic conditions, as elevated PI(3,5)P₂ induces Ca²⁺ efflux under hyperosmotic conditions (Dong et al., 2010).

In parallel we also tested the effect of deleting the $PI(3,5)P_2$ 5-phosphatase FIG4 on Ca^{2+} transport. Cells lacking Fig4 have reduced levels of $PI(3,5)P_2$ (Duex et al., 2006b; Duex et al., 2006a), and thus we predicted that it would have an intermediate effect compared to vacuoles with the $fab1^{EEE}$ mutation. As expected $fig4\Delta$ vacuoles showed attenuated Ca^{2+} uptake similar to $fab1^{EEE}$ yet did not show the same delays in Ca^{2+} uptake or efflux (Fig. 4E) Due to the reduced level of $PI(3,5)P_2$ present on $fig4\Delta$ vacuoles we expected that Ca^{2+} efflux would be resistant to added $C8-PI(3,5)P_2$. In accord with the other data, $fig4\Delta$ vacuoles were indeed resistant to $C8-PI(3,5)P_2$ compared to wild type vacuoles (Fig. 4F). This is consistent with our previous finding with $fig4\Delta$ vacuoles and lipid mixing (Miner et al., 2019).

PI(3,5)P₂ acts to regulate Pmc1 during fusion

To hone in on the mechanism by which $PI(3,5)P_2$ regulates Ca^{2+} flux during fusion we generated knockouts of the known vacuolar Ca^{2+} transporters: Pmc1, Vcx1, and Yvc1. As reported by Merz and Wickner, deletion of these transporters was unable to abolish the release of Ca^{2+} during fusion (Merz and Wickner, 2004). In keeping with their findings, we found that Ca^{2+} efflux was close to wild type with each of the deletion strains (Fig. 5A-B). We next asked if treating these deletion strains with C8-PI(3,5)P₂ would inhibit Ca^{2+} flux to the same extent as seen with wild type vacuoles. We found that C8-PI(3,5)P₂ inhibited Ca^{2+} efflux in all the deletion strain vacuoles tested except for $pmc1\Delta$ (Fig. 5A-B). Thus, it was likely that PI(3,5)P₂ affected Ca^{2+} transport through Pmc1. Because Pmc1 transports Ca^{2+} into the vacuole, the effect on the observed efflux was likely due to the hyper-stimulation of Pmc1. It is important to emphasize that deleting YVC1 had no effect on the inhibition of Ca^{2+} flux by PI(3,5)P₂. While Yvc1 is essential for the

 $PI(3,5)P_2$ activated Ca^{2+} transport during vacuole fission, this channel does not appear to play a role during vacuole fusion.

If $PI(3,5)P_2$ was truly acting to trigger Pmc1 activity we hypothesized that the effect should be reversible by the Ca^{2+} pump inhibitor Verapamil (Calvert and Sanders, 1995; Teng et al., 2008). In Fig 5B we summarize data showing that Verapamil blocked the effect of C8- $PI(3,5)P_2$ on Ca^{2+} efflux. This further indicates that the visualized Ca^{2+} efflux could instead be the inhibition of further Ca^{2+} uptake while the cation is released at a constant rate by another mechanism.

To further test the effect of Verapamil on Ca^{2+} transport we treated vacuoles with a dose response curve of the inhibitor at the beginning of the assay. Using wild type vacuoles, we found that Verapamil inhibited Ca^{2+} uptake in a dose dependent manner when added at the beginning of the assay (Fig. 6A-B). Next, we tested if Ca^{2+} uptake continues late in the reaction by adding Verapamil after the initial uptake of Ca^{2+} . As seen in Figure 6C-D, adding Verapamil resulted in a rapid accumulation of extraluminal Ca^{2+} . This suggests that under control conditions, Ca^{2+} uptake continues through the docking stage, 10-15 min into the pathway, after which uptake is either inactivated or simply overcome by Ca^{2+} efflux.

Given the surprising result that Ca^{2+} efflux appeared to be triggered by a reduction in Pmc1 activity we sought to clarify whether the main cause of Ca^{2+} efflux during membrane fusion was through the generation of transient pores as has been proposed. If transient pore formation serves as the primary efflux pathway, then we would only expect Verapamil to trigger Ca^{2+} efflux if fusion was allowed to proceed. Contrary to this explanation, we found that Verapamil was able to trigger a Ca^{2+} efflux event equivalent to standard fusion conditions even when vacuoles were treated with the membrane fusion inhibitor α -Sec17.

PI(3,5)P₂ regulates Pmc1 interactions with the V-ATPase.

Others have previously shown that Pmc1 activity is inhibited in part through its binding to the vacuolar R-SNARE Nyv1 (Takita et al., 2001). We thus tested if added Pl(3,5)P $_2$ would alter the interaction. To accomplish this, we used vacuoles that harbored Pmc1 tagged with HA in order to co-immunoprecipitate Nyv1 (Cunningham and Fink, 1996). We tested Nyv1-Pmc1 interactions in the presence of buffer, C8-Pl(3,5)P $_2$, ML1-N and Apilimod. We found that C8-Pl(3,5)P $_2$ did not alter the interactions between Pmc1 and Nyv1. However, inhibiting Pl(3,5)P $_2$ production with Apilimod reduced the Nyv1-Pmc1 immunoprecipitation (Fig. 7A-B). Although, Pl(3,5)P $_2$ is not known to bind either Nyv1 or Pmc1 directly, it has been shown to bind to the V $_0$ component Vph1 where it stabilizes V $_0$ -V $_1$ holoenzyme assembly (Li et al., 2014). Because the V-ATPase is known to bind Pmc1 we tested if C8-Pl(3,5)P $_2$ would affect Pmc1-Vph1 interactions (Tarassov et al., 2008). Indeed, we found that Pl(3,5)P $_2$ reduced Vph1 interactions with Pmc1. Although, the interactions were not fully disrupted, moderate changes could have large effects on protein function. As a whole, these experiments suggest that the V-ATPase and Pl(3,5)P $_2$ control Pmc1 activity to modulate Ca $^{2+}$ transport across the vacuole membrane during fusion. Although the exact mechanism by which Pl(3,5)P $_2$ levels alters Nyv1-Pmc1-Vph1

interactions remains to be elucidated, these findings are an exciting entrée into a new role for PI(3,5)P₂ in vacuole homeostasis.

DISCUSSION

The yeast vacuole serves as the primary reservoir for Ca^{2^+} , which are translocated across the membrane through multiple transporters including Pmc1, Yvc1, and Vxc1. Although the regulatory mechanisms for each of these transporters are not fully understood, it has been shown that various lipids affect their function in an organelle specific manner. For instance, $PI(3,5)P_2$ regulates Ca^{2^+} release through Yvc1 during the osmotically driven vacuole fission (Dong et al., 2010). Ca^{2^+} is also released from the vacuole during the fusion process upon the formation of *trans*-SNARE complexes (Peters and Mayer, 1998; Bayer et al., 2003; Merz and Wickner, 2004). More recently, we found that elevated levels of $PI(3,5)P_2$ inhibited vacuole fusion at a stage between *trans*-SNARE pairing and outer-leaflet lipid mixing, i.e. hemifusion and full bilayer fusion (Miner et al., 2019). This was observed through adding exogenous C8- $PI(3,5)P_2$ or through using *FAB1* mutations that were either kinase dead or overproduced the lipid. That study established that low levels of $PI(3,5)P_2$ are needed for fusion to occur, but elevated concentrations are inhibitory. At this point it is unclear whether the effects of $PI(3,5)P_2$ are direct or if the ratio with its precursor PI3P and downstream degradation product(s) also play a role. In either circumstance, it is clear that $PI(3,5)P_2$ has a biphasic effect on vacuole fusion.

The biphasic nature of $PI(3,5)P_2$ in vacuole fusion is not limited to this lipid. For instance, PA is needed for reconstituted proteoliposome fusion to occur and for Vam7 binding to vacuoles (Mima and Wickner, 2009; Miner et al., 2016). Yet, inhibiting PA phosphatase activity to produce diacylglycerol (DAG), resulting in elevating PA concentrations, blocks vacuole fusion at the priming stage through sequestering Sec18 from *cis*-SNARE complexes (Sasser et al., 2012a; Starr et al., 2016). Likewise, low levels of phospholipase C (PLC) stimulates vacuole fusion through converting $PI(4,5)P_2$ to DAG, while high levels of PLC potently inhibits fusion (Jun et al., 2004). Finally, while $PI(4,5)P_2$ is well known to stimulate vacuole fusion, an excess of the lipid can also inhibit the pathway (Fratti et al., 2004; Karunakaran and Fratti, 2013; Mayer et al., 2000; Miner et al., 2019). These examples show that the balance of substrate and product lipids can serve to move a pathway forward or arrest its progress.

In this study we continued our investigation on the inhibitory nature of $PI(3,5)P_2$ on vacuole fusion. Our work has identified that elevated levels of $PI(3,5)P_2$ block the net Ca^{2+} efflux observed during fusion vacuole fusion. The effect of $PI(3,5)P_2$ on Ca^{2+} transport was shown by various means including the use of the kinase dead $fab1^{EEE}$ and hyperactive $fab1^{T2250A}$ mutants. While both mutants were previously shown to inhibit hemifusion (Miner et al., 2019), here we see that only $fab1^{T2250A}$ inhibited Ca^{2+} efflux, which is in accord with a model in which excessive $PI(3,5)P_2$ is a negative regulator of the fusion pathway. In contrast, kinase dead $fab1^{EEE}$ vacuoles showed enhanced Ca^{2+} efflux. In both instances, Ca^{2+} flux was partially restored to wild type levels when the amount of free $PI(3,5)P_2$ was increased on $fab1^{EEE}$ vacuoles or reduced on $fab1^{T2250A}$ vacuoles. The increased Ca^{2+} efflux seen with $fab1^{EEE}$ vacuoles observed

here combined with our previous report showing that hemifusion was inhibited by this mutant indicates that Ca²⁺ flux precedes hemifusion.

Another revelation that comes from these experiments is that the effect of PI(3,5)P₂ on Ca²⁺ transport was independent of the TRP channel Yvc1. This is the opposite to what occurs during vacuole fission induced by osmotic stress, when PI(3,5)P2 interacts with Yvc1 leading to the release of Ca2+ from the vacuole lumen (Dong et al., 2010). Our previous work with PI(3,5)P₂ suggested that this lipid inversely regulates fission and fusion. The current work further supports this model and specifies that the effects occur through differentially modulating Ca2+ transport across the vacuole bilayer. Surprisingly, we found that the inhibition of Ca2+ transport by PI(3,5)P₂ was dependent on the Ca²⁺ ATPase Pmc1, which utilizes ATP hydrolysis to move Ca2+ into the vacuole lumen against the concentration gradient. This was evident by the arrest in Ca²⁺ uptake followed by an increase in extraluminal Ca²⁺ when pmc1∆ vacuoles where treated with PI(3,5)P₂. Furthermore, it appears that the regulation is immediate, as treatment with the Fab1/PIKfyve inhibitor Apilimod led to a rapid release of Ca²⁺, suggesting that PI(3,5)P₂ needs to be newly generated, albeit at low levels to regulate Ca2+ transport. Under unencumbered conditions it is likely that PI(3,5)P2 levels are kept in check through the phosphatase activity of Fig4. One of the most striking findings from our study is that the Ca²⁺ efflux associated with fusion appears to be in part through the inactivation of Ca²⁺ influx. Additionally, our data suggests that the primary route of efflux can be uncoupled from membrane fusion as inhibition of SNARE priming was unable to prevent the trigger of Ca2+ efflux following treatment with Verapamil. We therefore conclude that there is a Ca²⁺ efflux route opposed by Pmc1.

The regulation of Pmc1 on yeast vacuoles is unclear relative to other P-type Ca^{2+} ATPases. Notwithstanding, others have shown that that the R-SNARE Nyv1 directly interacts with Pmc1 to reduce Ca^{2+} transport (Fig. 8) (Takita et al., 2001). Although not directly shown, their model posits that increased Nyv1 would proportionally have increased interactions with Pmc1 and reduce activity. We sought to learn if elevated $PI(3,5)P_2$ levels would dissociate the Nyv1-Pmc1 complex to de-repress Ca^{2+} transport. In our hands we found that the Nyv1-Pmc1 interaction was variable when $C8-PI(3,5)P_2$ was added at levels that inhibited Ca^{2+} efflux making it difficult to determine if there was an effect. However, inhibiting $PI(3,5)P_2$ production with Apilimod reduced the amount of Nyv1 that co-isolated with Pmc1, suggesting $PI(3,5)P_2$ can affect complex stability. This regimen of Apilimod treatment resulted in a rapid increase in extraluminal Ca^{2+} . Together, these data bolster the notion that $PI(3,5)P_2$ levels regulate Pmc1 activity.

Neither Pmc1, nor Nyv1 have been shown to bind $PI(3,5)P_2$ directly, which raises the question as to how this lipid affects the complex? While a direct interaction has not been shown it is still possible that an indirect path exists. One of the few bona fide $PI(3,5)P_2$ binding proteins on the vacuole is the V_0 component Vph1 (Li et al., 2014). Their work showed that $PI(3,5)P_2$ stabilized the assembly of the V_1 - V_0 holoenzyme. Because Vph1 binds Pmc1 (Tarassov et al., 2008) and R-SNAREs have been shown to directly bind V_0 (Di Giovanni et al., 2010), we hypothesized that $PI(3,5)P_2$ could alter Vph1-Pmc1 and Pmc1-Nyv1 interactions (Fig. 8). Our

Pmc1 immunoprecipitations showed that Vph1 indeed interacted with Pmc1. Furthermore, we found that adding C8-PI(3,5)P₂ reduced the Pmc1 interaction with Vph1. When we consider that blocking V-ATPase activity results in a decrease in Ca²⁺ uptake by vacuoles (Miseta et al., 1999), we can extrapolate to think that promoting V-ATPase activity through PI(3,5)P₂-Vph1 interactions and promotion of V₁-V₀ complex could inversely promote Ca²⁺ uptake by reducing Vph1-Pmc1 interactions. Our data taken together with the finding that blocking Vph1 activity through addition of α -Vph1 leads to inhibition of Ca²⁺ efflux (Bayer et al., 2003), we suggest a model where Vph1 binds to and inhibits Pmc1 and that the addition of C8-PI(3,5)P₂ or α -Vph1 prevents this association. Thus, we can conclude that PI(3,5)P₂ regulates Pmc1 in part through binding Vph1 and dissociating Pmc1-Vph1. Although this model is incomplete, we believe these data gives further insight into the mechanism behind Ca²⁺ efflux during the fusion process as well as the ability of PI(3,5)P₂ to act as a potent fusion-fission switch.

MATERIALS AND METHODS

Reagents

Soluble reagents were dissolved in PIPES-Sorbitol (PS) buffer (20 mM PIPES-KOH, pH 6.8, 200 mM sorbitol) with 125 mM KCl unless indicated otherwise. Anti-Sec17 IgG (Mayer et al., 1996), and Pbi2 (Slusarewicz et al., 1997) were prepared as described previously. C8-PA (1,2-dioctanoyl-sn-glycero-3-phosphate), C8-PI3P (1,2-dioctanoyl-phosphatidylinositol 3-phosphate), and C8-PI(3,5)P₂ (1,2-dioctanoyl-phosphatidylinositol 3,5-bisphosphate), BODIPY-TMR C6-PI3P and BODIPY-TMR C6-PI(3,5)P₂ were purchased from Echelon Inc. Apilimod was from MedKoo Biosciences and Verapamil was from Cayman Chemical. Both were dissolved in DMSO as stock solutions. Anti-HA antibody was from Thermo-Fisher. GST-ML1-N was produced as described (Dong et al., 2010; Miner et al., 2019). Cy3-maleimide (GE Healthcare) was used to label GST-ML1-N according to the manufacturer's instructions.

Strains

Vacuole Isolation and In-vitro Ca²⁺ Flux Assay

Vacuoles were isolated as described (Haas et al., 1994). Vacuole lumen Ca²⁺ was measured as described (Miner and Fratti, 2019; Miner et al., 2016; Sasser et al., 2012b). *In vitro* Ca²⁺ transport reactions (60 μl) contained 20 μg vacuoles from BJ3505 backgrounds, fusion reaction buffer, 10 μM CoA, 283 nM Pbi2 (inhibitor of Proteinase B), and 150 nM of the Ca²⁺ probe Fluo-4 dextran conjugate MW 10,000 (Invitrogen) or Cal-520 dextran conjugate MW 10,000 (AAT Bioquest). Reaction mixtures were loaded into a black, half-volume 96-well flat-bottom plate with nonbinding surface (Corning). ATP regenerating system or buffer was added, and reactions were incubated at 27°C while Fluo-4 or Cal-520 fluorescence was monitored. Samples were analyzed using a POLARstar Omega fluorescence plate reader (BMG Labtech) with the excitation filter at 485 nm and emission filter at 520 nm for Fluo-4 or Cal-520. Reactions were initiated with the addition of ATP regenerating system following the initial measurement. The effects of inhibitors on efflux were determined by the addition of buffer, inhibitors, or C8-lipids immediately following Ca²⁺ influx. Calibration was done using buffered Ca²⁺ standards (Invitrogen).

PI3P 5-kinase assay and Thin-Layer Chromatography

Fab1 kinase activity was measured with an assay adapted from the detection of Fig4 and Plc1 activity (Rudge et al., 2004; Jun et al., 2004) with some modifications. Kinase reactions (30 μ l) contained 6 μ g vacuoles from BJ3505 backgrounds, fusion reaction buffer, ATP regenerating system, 10 μ M CoA, 283 nM Pbi2, 2 μ M BODIPY-TMR C6-Pl3P, and 1 mM sodium orthovanadate. Reaction mixtures were incubated on ice and immediately quenched with acetone (100 μ l) or incubated at 27°C or on ice for 15 min in the presence of Apilimod or DMSO prior to quenching. Reactions were quenched with acetone after incubation. Following incubation acidic phospholipids were extracted from all reactions (Sasser et al., 2013). TLC plates (Partisil LK6D Silica Gel Plates (60Å), Whatman) were pretreated with 1.2 mM EGTA and 1% potassium oxalate (w/v) in MeOH/Water (3:2) and then dried at 100°C for 30 min prior to use. Dried lipids were resuspended in CHCl3/MeOH (1:1) (40 μ l) and 5 ul was spotted on the plate. Plates were run in CHCl3/acetone/MeOH/AcOH/Water (46:17:15:14:8). Individual channels were loaded with Pl3P and Pl(3,5)P2 standards (Echelon). Imaging of plates was performed using a ChimiDoc MP System (BioRad) and densitometry was determined with Image Lab 4.0.1 software.

Vacuole Docking

Docking reactions (30 µl) contained 6 µg of wild type vacuoles were incubated in docking buffer (20 mM PIPES-KOH pH 6.8, 200 mM sorbitol, 100 mM KCl, 0.5 mM MgCl₂), ATP regenerating system (0.3 mM ATP, 0.7 mg/ml creatine kinase, 6 mM creatine phosphate), 20 µM CoA, and 283 nM Pbi2 (Fratti et al., 2004). Pl(3,5)P₂ was labeled with 2 µM Cy3-GST-ML1-N. Reactions were incubated at 27°C for 20 min. After incubating, reaction tubes were placed on ice and vacuoles were stained with 1 µM MDY-64. Reactions were next mixed with 50 µl of 0.6% low-melt agarose (in PS buffer), vortexed to disrupt non-specific clustering, and mounted on slides for observation by fluorescence microscopy. Images were acquired using a Zeiss Axio Observer Z1 inverted microscope equipped with an X-Cite 120XL light source, Plan Apochromat 63X oil objective (NA 1.4), and an AxioCam CCD camera.

Pmc1-HA immunoprecipitation

Pmc1-HA complex isolation was performed using 20X fusion reactions. Individual reactions were treated with 150 μ M C8-PI(3,5)P₂, 500 nM GST-ML1-N, 500 μ M Apilimod or buffer alone. After 30 min, reactions were placed on ice. Next, reactions were sedimented (11,000 g, 10 min, 4°C), and the supernatants were discarded before extracting vacuoles with solubilization buffer (SB: 50 mM Tris-HCl, pH 8.0, 2 mM EDTA, 150 mM NaCl, 0.5% Tween-20, 1 mM PMSF). Reactions were then nutated for 1 hour at 4°C. Insoluble debris was sedimented (16,000 g, 10 min, 4°C) and 350 μ l of supernatants were removed and placed in chilled tubes. Next, 35 μ l was removed from each reaction as 10% total samples, mixed with 17.5 μ l of 5X SDS loading buffer and 17.5 μ l PS buffer. Equilibrated anti-HA agarose beads (60 μ l) were incubated with the extracts (15 h, 4°C, nutation). Beads were sedimented and washed 5X with 1 ml SB (800 g, 2 min, 4°C), and bound material was eluted with 0.1 M Glycine, pH 2.2 and eluates were mixed with SDS-loading buffer. Protein complexes were resolved by SDS-PAGE and examined by Western blotting.

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Author contributions

GEM and RAF, Conception and design, Data analysis and interpretation, Drafting or revising the article; GEM, KDS, AG, MLS, ECE BCJ Acquisition of data, data analysis

Conflict of Interest

The authors declare that they do not have conflicts of interest with the contents of this article.

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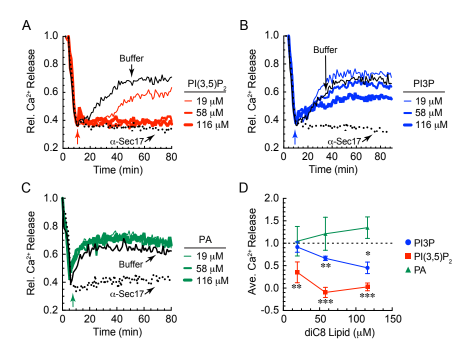


FIGURE LEGENDS

Figure 1. PI(3,5)P₂ blocks Ca²⁺ efflux from vacuoles during fusion. Vacuoles were harvested from BJ3505 and 2X fusion reactions containing 150 μM Fluo-4 dextran. After 10 min of incubation with ATP (arrow) at 27°C to allow for the uptake of Ca²⁺, reactions were treated with C8-PI(3,5)P₂ (A), C8-PI3P (B), or C8-PA (C) at the indicated concentrations. Reactions were further incubated at 27°C and fluorescence was measured every 30 sec for 80 min. Values were normalized to 1.0 representing the extraluminal Ca²⁺ at the beginning of the reaction and expressed as relative (Rel.) values compared to the untreated control. Separate reactions were incubated with 85 μg/ml anti-Sec17 IgG to block SNARE-dependent Ca²⁺ efflux. (D) Average Ca²⁺ efflux from A-C. Error bars are S.E.M. (n=3). *p<0.05, *p<0.01, **p<0.001.

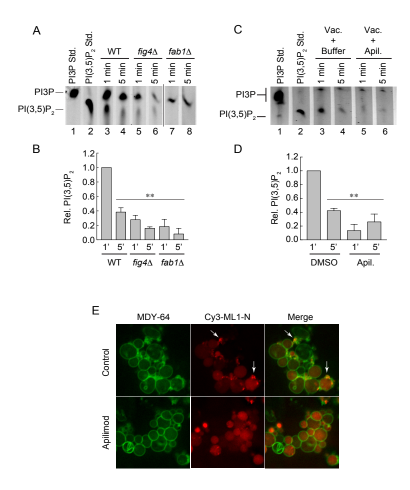


Figure 2. Apilimod inhibits Fab1 activity. (A) Vacuoles were incubated with BODIPY-TMR C6-PI3P at 27°C after which reactions were quenched with acetone and lipids were extracted and resolved by TLC to detect the production of BODIPY-TMR C6-PI(3,5)P₂ by Fab1 activity. Wild type vacuoles were tested in parallel to those from $fig4\Delta$ and $fab1\Delta$ strains. Pure BODIPY-TMR C6-PI3P and C6-PI(3,5)P₂ were used as standards in lanes 1 and 2. (B) Average of three experiments in panel A. (C) Vacuoles were treated with DMSO (carrier) or 500 μM Apilimod. (D) Average of three experiments in panel C. (E) Vacuoles were incubated under docking conditions for 20 min. Endogenous PI(3,5)P₂ was labeled with 2.5 μM Cy3-ML1-N and the total membranes were labeled with 1 μM MDY-64. Vacuoles were treated with Apilimod or DMSO. Arrows point at examples of PI(3,5)P₂ puncta. Error bars are S.E.M. (n=3).

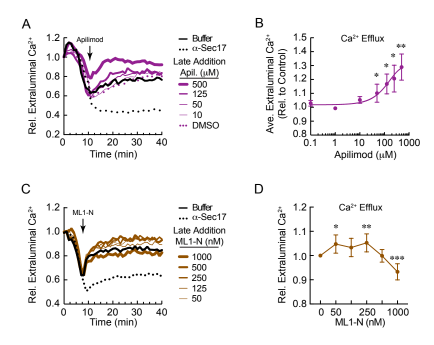


Figure 3. Apilimod modulates Ca²⁺ flux during vacuole fusion. (A) Vacuoles were treated with buffer of anti-Sec17 IgG and incubated for 8-10 min to allow Ca²⁺ uptake. Next, vacuoles were treated with a dose response curve of Apilimod or DMSO (Carrier). Reactions were further incubated for a total 40 min. (B) Quantitation of Ca²⁺ efflux at 30 min normalized to the untreated control. (C) Vacuoles were treated with buffer or anti-Sec17 IgG and incubated for 10 min to allow Ca²⁺ uptake. Next, vacuoles were treated with a dose response curve of GST-ML1-N and reactions were further incubated for a total of 30 min. (D) Ca²⁺ release at 30 min. Values were normalized to the untreated control at 30 min. Error bars are S.E.M. (n=3). *p<0.05, *p<0.01, *p<0.001. Significant differences were in comparison to untreated wild type efflux

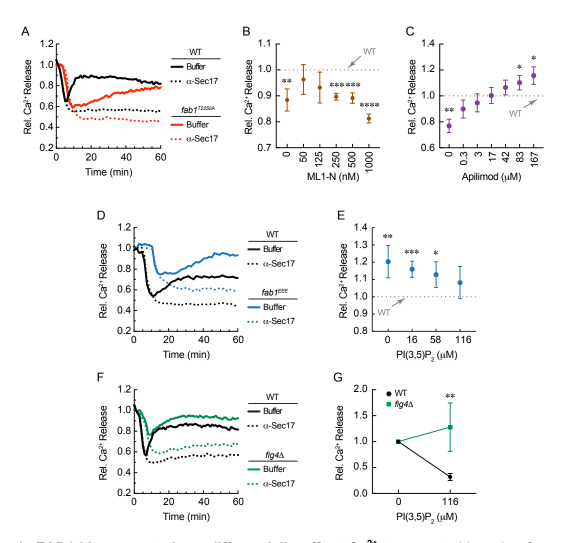


Figure 4. *FAB1* kinase mutations differentially affect Ca²⁺ transport. Vacuoles from wild type yeast were tested for Ca²⁺ transport in parallel with those from $fab1^{T2250A}$ (A), $fab1^{EEE}$ (D) or $fig4\Delta$ (F) yeast. (B) $Fab1^{T2250A}$ vacuoles were treated with a dose response curve of GST-ML1-N after 10 min of incubation. Reactions were then incubated for a total of 60 min. Average Ca²⁺ efflux values at 30 min of incubation were normalized relative to wild type and given a value of 1 (dotted line). (C) $Fab1^{T2250A}$ vacuoles were treated with a dose response curve Apilimod as described for ML1-N in (B). Values were normalized to wild type efflux at 30 min. (E) $fab1^{EEE}$ vacuoles were treated with a dose curve of C8-PI(3,5)P₂. Values were taken at 30 min and normalized wild type efflux without treatment at 30 min. (G) Wild type $fig4\Delta$ vacuoles were treated with buffer or C8-PI(3,5)P₂. Efflux values were taken at 30 min with or without treatment for wild type and $fig4\Delta$ vacuoles Values were normalized to the no treatment condition for each vacuole type. Error bars are S.E.M. (n=3). Significant differences were in comparison wild type. *p<0.05, **p<0.01, ****p<0.001, ****p<0.0001.

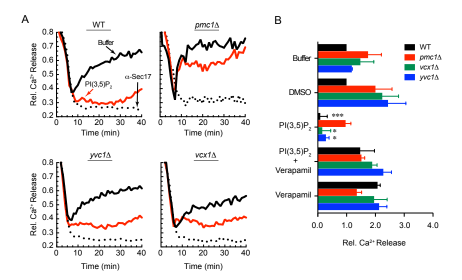


Figure 5. PI(3,5)P₂ regulates Ca²⁺ transport through Pmc1 during fusion. (A) Vacuoles isolated from wild type as well as $pmc1\Delta$, $yvc1\Delta$ and $vcx1\Delta$ strains were used in Ca²⁺ transport assays as described. Reactions were treated with anti-Sec17, buffer or 116 μM PI(3,5)P₂ added after 5 min of incubation. Reactions were incubated for a total of 40 min and fluorescence was measured every 60 sec. (B) Average Ca²⁺ transport levels for multiple experiments with wild type, $pmc1\Delta$, $yvc1\Delta$ or $vcx1\Delta$ vacuoles. After 10 min of incubation reactions were treated with buffer, DMSO, 116 μM C8-PI(3,5)P₂, 500 μM Verapamil or C8-PI(3,5)P₂ and Verapamil together and incubated for a total of 40 min. *p<0.05, ***p<0.001.

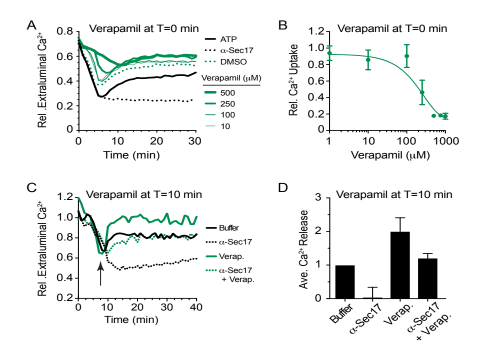
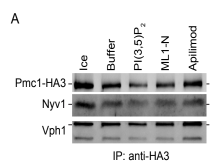


Figure 6. Verapamil inhibits Pmc1 activity. (A) Wild type vacuoles were treated with a dose curve of Verapamil at T=0 min to block Ca^{2+} uptake. Separate reactions were treated with buffer, DMSO or anti-Sec17 as controls. (B) Average Ca^{2+} transport levels for multiple experiments shown in panel A. Reactions were incubated for a total of 30 min and fluorescence was measured every 30 sec. (C) Wild type vacuoles were treated with 500 μM Verapamil or DMSO after 8 min of incubation to allow initial Ca^{2+} uptake. Reactions were analyzed as described in panel A. (D) Average Ca^{2+} transport levels for multiple experiments shown in panel C.



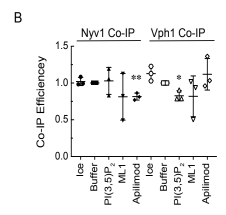


Figure 7. C8-PI(3,5)P₂ enhances Pmc1 interactions with Nyv1. (A) Vacuoles expressing Pmc1-HA3 were incubated in large scale reactions (20X) in the presence of 150 μM C8-PI(3,5)P₂, 500 nM ML1-N, 500 μM Apilimod or buffer alone. Reactions were incubated for 30 min at 27° C. After incubating, the reactions were place on ice and solubilized and processed for immunoprecipitation with anti-HA agarose. Protein complexes were eluted with sodium thiocyanate and resolved by SDS-PAGE and probed by Western blotting for the presence of Pmc1-HA3 and Nyv1. (B) Average quantitation of Nyv1 and Vph1 bound to Pmc1-HA. Values represent concentrations relative to Pmc1-HA3 at time = 30 min. Error bars represent S.E. (n=3). *p<0.05, **p<0.01.

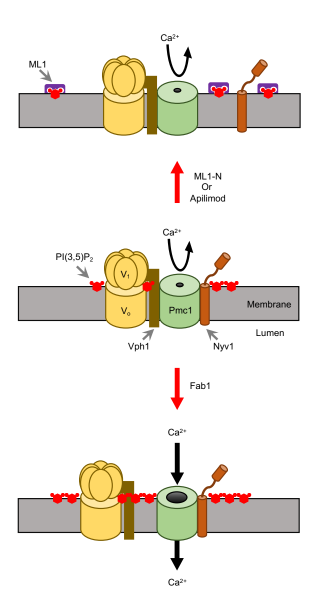


Figure 8. Model for PI(3,5)P₂ regulation of Ca2+ flux during vacuole fusion.

Table S1. Yeast strains used in this study

Strain	Genotype	Source
BJ3505	MATα ura3–52 trp1-Δ101 his3-Δ200	(Jones et al., 1982)
	lys2–801 gal2 (gal3) can1 prb1-∆1.6R pep4::HIS3	
K699	W303-1A, <i>PMC1::HA</i>	(Cunningham and
		Fink, 1996)
RFY74	BJ3505, <i>yvc1∆::kanMX</i> 6	(Miner et al., 2019)
RFY76	BJ3505, <i>fab1∆::kanMX6</i>	(Miner et al., 2019)
RFY78	RFY76, pRS416-FAB1 ^{T2250A}	(Miner et al., 2019)
RFY80	RFY76, <i>pRS416-FAB1^{EEE}</i>	(Miner et al., 2019)
RFY82	BJ3505, <i>fig4∆::kanMX</i> 6	(Miner et al., 2019)
RFY84	BJ3505, <i>pmc1∆::kanMX</i> 6	This study
RFY86	BJ3505, <i>vcx1∆::kanMX</i> 6	This study