Synergism between Inositol Polyphosphates and TOR Kinase Signaling in Nutrient Sensing, Growth Control, and Lipid Metabolism in Chlamydomonas

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The networks that govern carbon metabolism and control intracellular carbon partitioning in photosynthetic cells are poorly understood. Target of Rapamycin (TOR) kinase is a conserved growth regulator that integrates nutrient signals and modulates cell growth in eukaryotes, though the TOR signaling pathway in plants and algae has yet to be completely elucidated. We screened the unicellular green alga Chlamydomonas reinhardtii using insertional mutagenesis to find mutants that conferred hypersensitivity to the TOR inhibitor rapamycin. We characterized one mutant, vip1-1, that is predicted to encode a conserved inositol hexakisphosphate kinase from the VIP family that pyrophosphorylates phytic acid (InsP6) to elucidate. We screened the unicellular green alga Chlamydomonas reinhardtii using insertional mutagenesis to find mutants that conferred hypersensitivity to the TOR inhibitor rapamycin. We characterized one mutant, vip1-1, that is predicted to encode a conserved inositol hexakisphosphate kinase from the VIP family that pyrophosphorylates phytic acid (InsP6). Unexpectedly, the rapamycin hypersensitive growth arrest of vip1-1 cells was dependent on the presence of external acetate, which normally has a growth-stimulatory effect on Chlamydomonas. vip1-1 mutants also constitutively overaccumulated triacylglycerols (TAGs) in a manner that was synergistic with other TAG inducing stimuli such as starvation. vip1-1 cells had reduced InsP7 and InsP8, both of which are dynamically modulated in wild-type cells by TOR kinase activity and the presence of acetate. Our data uncover an interaction between the TOR kinase and inositol polyphosphate signaling systems that we propose governs carbon metabolism and intracellular pathways that lead to storage lipid accumulation.

INTRODUCTION

Photosynthetic cells are defined by their ability to fix and metabolize inorganic carbon (CO₂ or HCO₃⁻). Many photosynthetic organisms can also grow heterotrophically or mixotrophically using externally supplied organic carbon or with a combination of photosynthesis and imported organic carbon (Pringsheim and Wiessner, 1960; Koch, 2004; Harris, 2009a; Morales-Sánchez et al., 2015). The means by which photosynthetic cells coordinate these two modes of carbon metabolism are relatively unexplored (Sheen, 1994; Coruzzi and Bush, 2001; Haldorf and Paul, 2003; Smith and Stitt, 2007), but the elucidation of intracellular carbon regulatory networks has the potential to improve our understanding of photosynthetic cell growth and provide insights that will allow harnessing of photosynthesis for improved yields and storage of fixed carbon.

The unicellular green alga Chlamydomonas reinhardtii is a model organism that has been used successfully to dissect and understand photosynthetic metabolism (Rochaix, 1995; Grossman, 2000; Dent et al., 2005; Minagawa and Tokutsu, 2015; Goodenough, 2015). Chlamydomonas cells can grow photosynthetically and can also metabolize acetate and related compounds to grow mixotrophically or heterotrophically. The relationship between these modes of growth is complex and involves partial suppression of photosynthesis and alterations in gene expression when acetate is present (Helfetz et al., 2000; Boyle and Morgan, 2008; Johnson and Aric, 2012; Roach et al., 2013; Chapman et al., 2015). However, very little is known about intracellular signaling pathways that sense carbon source and control carbon metabolism and utilization in Chlamydomonas.

Target of Rapamycin (TOR) protein kinase is a conserved and essential regulator of eukaryotic cell growth that is thought to integrate external and internal metabolic cues to balance cell growth with energy and nutrient supplies (Menand et al., 2002; Loewith and Hall, 2011; Montané and Menand, 2013; Xiong and Sheen, 2012; Xiong et al., 2013; Rexin et al., 2015). In yeast and metazoans, TOR is found in two different complexes, TORC1 and TORC2, with distinct functions in cell growth and cytoskeletal organization, respectively, while green algae and land plants only...
contain the components of TORC1, namely, TOR, RAPTOR, and LST8 (Diaz-Troya et al., 2008; Loewith and Hall, 2011; Moreau et al., 2012; Maegawa et al., 2015). In plants, the TOR complex plays similar roles as TORC1 in animals and fungi by influencing cytoplasmic protein translation, carbon metabolism, cell growth, cell proliferation, senescence, and autophagy (Deprost et al., 2007; Liu and Bassham, 2010; Caldana et al., 2013; Xiong and Sheen, 2015; Rexin et al., 2015). However, the specific downstream signaling targets and intersecting pathways with TOR kinase in plants are still not completely understood, especially as they relate to control of metabolism. Although the Arabidopsis thaliana TOR gene AtTOR is essential (Menand et al., 2002), conditional silencing resulted in early senescence with a decrease in photosynthesis linked to chlorophyll breakdown (Deprost et al., 2007) along with other metabolite changes resulting from impaired TOR kinase signaling and photosynthetic growth control.

In several species, including Chlamydomonas, TOR can be inhibited by the macrolide drug rapamycin, which has proven to be a pivotal tool for investigating TOR signaling (Crespo et al., 2005; Loewith and Hall, 2011; Rexin et al., 2015). Although Chlamydomonas is sensitive to rapamycin, at saturating drug concentrations cells grow slowly but do not arrest growth completely (Crespo et al., 2005). A screen for rapamycin resistance mutations identified alleles of FKB1 that encode an ortholog of FK506 binding protein FKB12 (Crespo et al., 2005; Vallon, 2005), a conserved peptidyl prolyl isomerase that forms a ternary complex with rapamycin and TORC1 and is required for rapamycin-mediated TOR inhibition (Heitman et al., 1991; Brown et al., 1994; Sabatini et al., 1994). Having a haploid genome and well developed molecular genetic tools, Chlamydomonas is a promising unicellular model organism in which to pursue functional studies of TOR kinase signaling and photosynthetic growth control.

Here, we used insertional mutagenesis in Chlamydomonas to screen for rapamycin-hypersensitive mutants. In doing so, we identified a loss-of-function mutation in VIP1, a gene encoding a conserved protein with predicted diphosphoinositol phosphate kinase activity whose function has not previously been linked to TOR signaling. Levels of InsP₃ and InsP₄, the predicted products of VIP1 kinase, were reduced in vip1-1, suggesting that these InsPs are required for cell growth in conjunction with TOR kinase activity. Unexpectedly, rapamycin hypersensitivity of vip1-1 mutants required the presence of acetate in its growth media and the mutant cells were also found to have altered levels of tricarboxylic acid (TCA) cycle intermediates and increased levels of neutral lipids compared with wild-type cells. Our findings have implications for carbon partitioning and metabolic control by TOR and InsPs in other eukaryotes where these signaling systems are conserved and may provide a means of manipulating carbon flux and increasing yields of valuable storage compounds in algae and crop plants.

RESULTS
Isolation of Rapamycin-Hypersensitive Mutants from Chlamydomonas

A previous screen for rapamycin resistance yielded only fkb1 mutants (Crespo et al., 2005), so we instead screened for rapamycin hypersensitivity among a population of insertional mutants (Galván et al., 2007) generated with the marker aph7 +” that confers resistance to the drug hygromycin (hygR phenotype) (Berthold et al., 2002). We anticipated that rapamycin hypersensitivity might result when growth-related processes that are regulated or co-regulated by TOR are impaired. Our wild-type strains (CC-1690 and CC-1691) and most strains that we have tested showed slow but not completely inhibited growth in the presence of rapamycin, with saturation occurring at 500 nM to 1 μM (Crespo et al., 2005). We transformed and screened around 60,000 insertional mutants in CC-1690 for arrested or reduced growth in 500 nM rapamycin but normal growth in its absence (Figure 1A; see Methods). After retesting initial positives, we found only two mutants that had a consistent rapamycin hypersensitive (rapHS) phenotype, one of which, 266F3 (later found to be caused by insertion in the VIP1 locus and designated vip1-1), is the focus of this study.

Among 600 total progeny from a backcross to a wild-type strain, we found consistent cosegregation of the rapHS phenotype with the hygR insertion (Figure 1B). We next tested whether the rapHS phenotype was dependent on the presence of the FKB12 protein encoded by FKB1. fkb1 vip1-1 double mutants were completely insensitive to rapamycin, indicating that the rapHS phenotype in vip1-1 results from impaired TOR kinase signaling rather than off-target effects of rapamycin (Figure 1C). Further supporting the specificity of the mutant phenotype was our finding that vip1-1 growth had a similar hypersensitivity to two other TOR kinase inhibitors with different inhibitory mechanisms from rapamycin, AZD8055 and Torin1 (Figure 1A) (Thoreen et al., 2009; Chresta et al., 2010).

We used adaptor-mediated PCR to identify a simple insertion of the HygR marker in the 11th coding exon of Cre03.g185500, whose predicted protein is related to a family of diphosphoinositol pentakisphosphate kinases also known as VIP proteins that will be described below (Figure 1D). The 266F3 mutation in Chlamydomonas was therefore designated vip1-1. Vegetative diploids with genotypes VIP1/VIP1, vip1-1/VIP1, and vip1-1/vip1-1 were constructed and tested to reveal that thevip1-1 mutation is a loss-of-function allele that is completely recessive to the wild-type allele (Supplemental Figure 1).

We cloned the ~11-kb wild-type VIP1 genomic locus and used it to construct complementation vectors that contain a paromomycin resistance marker (ParoR) for selection (Sizova et al., 2001) and either unmodified VIP1 (pVIP1) or a version of VIP1 with an N-terminally encoded OLLAS peptide epitope tag (pOL-VIP1) (Park et al., 2008) (Figure 1D; see Methods). Among >500 control transformants of vip1-1 using an empty vector, we never observed complementation of the rapHS phenotype, whereas among transformants that received either pVIP1 or pOL-VIP1, we observed restoration of wild-type or near wild-type growth in the presence of rapamycin at a rate of ~3%, pVIP1 (2/80) and pOL-VIP1 (15/480) (Figure 1A). In transformants complemented with pOL-VIP1, we detected a single band...
on immunoblots probed with anti-OLLAS antibodies migrating at the predicted size of tagged VIP1 (~170 kD) (Figure 1E). The same antibody was used in immunofluorescence experiments on fixed samples to localize VIP1 primarily to the cytoplasm (Figure 1F). We conclude that the rapHS phenotype of vip1-1 is caused by a loss-of-function insertion mutation that either removes or impairs a cytosol-localized VIP1 protein.

VIP1 Encodes a Predicted Diphosphoinositol Pentakisphosphate Kinase

The predicted Chlamydomonas VIP1 protein (CrVIP1) showed high-scoring BLASTP similarity to a conserved family of eukaryotic diphosphoinositol pentakisphosphate kinases of the VIP family (Mulugu et al., 2007; Desai et al., 2014; Laha et al., 2015). VIP proteins contain two conserved domains, an ATP_GRASP domain (pfam 13535) that is predicted to have inositol di-phosphate kinase activity (Mulugu et al., 2007; Fridy et al., 2007; Lin et al., 2009; Pöhlmann et al., 2014) followed by a histidine phosphatase domain (pfam 00328) that in CrVIP1 is interrupted with nonconserved sequences (Figure 2A). A more thorough search revealed a second but divergent VIP-related protein in Chlamydomonas that we designate CrVIP2. To better understand the phylogenetic relationships between CrVIP1, CrVIP2, and other eukaryotic VIP proteins, we constructed a phylogeny using the Chlamydomonas VIP putative paralogs and a diverse set
of VIP homologs from other taxa (Figure 2B). Overall, the phylogeny matches the scenario of at least one VIP gene being present at the base of the eukaryotic radiation and undergoing independent paralogous duplications in several lineages, including chlorophyte algae where the duplication was present in the ancestor of Chlamydomonas and its multicellular relative Volvox carteri. Additional independent duplications were found in the land plants with evidence for at least one event in monocots (Oryza sativa) and another in dicots (Arabidopsis) (Figure 2B; Supplemental Data Set 1).

With the exception of InsP₆ (phytic acid), which accumulates to high levels in many plant seeds, higher order InsPs (InsP₅, InsP₆, InsP₇, and InsP₈) are typically low abundance molecules with diverse roles in intracellular signaling (Raboy, 2003; Barker et al., 2009; Monserrate and York, 2010; Gillaspy, 2011; Wilson et al., 2013; Williams et al., 2015; Livermore et al., 2016). VIP1 is predicted to pyrophosphorylate InsP₆ at the 1,3 positions, generating InsP₇ and possibly 1,3PP-InsP₄ (InsP₈) (Figure 2C) suggests that the same spectrum of InsPs will be produced in Chlamydomonas (Figure 2C). We note that unlike fungi/animals/amoebae, which all contain the predicted InsP biosynthetic enzymes in Chlamydomonas (Figure 2C). The predicted products of CrVIP1, InsP₇, and InsP₈, are usually the lowest abundance InsPs in eukaryotic cells and difficult to detect (Bennett et al., 2006; Losito et al., 2009; Lin et al., 2009; Desai et al., 2014). Sensitive in vivo labeling procedures for InsP detection using radiolabeled 3H-inositol have been successful in other species (Azevedo and Saiardi, 2006; Stevenson-Paulik et al., 2006), but proved unfeasible in Chlamydomonas, which does not take up this precursor. Furthermore, InsPs were not abundant enough in Chlamydomonas lysates to enable utilization of electrophoretic separation and staining detection methods that do not require labeling (Losito et al., 2009). Therefore, we developed a reproducible and sensitive liquid chromatography and tandem mass spectrometry (LC-MS/MS) method for detection and discrimination of different InsP species calibrated using commercial standards and a synthetic InsP (3-deoxy-3-fluoro-α-myo-inositol 1,4,5 trisphosphate [3-fluoro-InsP₃]), as an internal standard for normalization (see Methods). Applying this method to InsPs purified from whole-cell lysates of Chlamydomonas, we compared profiles of wild-type, vip1-1, and complemented vip1-1+pOL-VIP1 strains.

Using LC-MS/MS, we were able to reproducibly detect six major InsP species in wild-type extracts: InsP₃, InsP₄, InsP₅, InsP₆, and InsP₈. Although we currently cannot discriminate between InsP stereoisomers using our method, the conserved repertoire of predicted InsP biosynthetic enzymes in Chlamydomonas (Figure 2C) suggests that the same spectrum of InsPs will be produced in Chlamydomonas as in plants. According to our analysis, we found InsP₅ to be the most abundant species while InsP₃ and InsP₄ were the least abundant (Figure 3A). Moreover, vip1-1 cells had significant defects in accumulation of InsP₅ and InsP₆ that were present at 20 to 30% of wild-type levels and fully or partially restored in complemented strains (Figure 3A, insets). The most abundant InsP species in Chlamydomonas, InsP₆, was somewhat reduced in vip1-1, while levels of three lower order InsPs (InsP₃, InsP₄, and InsP₅) were not significantly altered (Figure 3A). We hypothesize that residual InsP₅ and InsP₆ detected in vip1-1 is due to the activity of CrVIP2 (Figures 2B and 2C) whose mRNA is expressed at <10% of the level of CrVIP1 mRNA under diurnal growth conditions (Zones et al., 2015). Together, our data support the predicted role of Chlamydomonas VIP1 as a diphosphoinositol kinase whose function in InsP₅ and InsP₆ biosynthesis is acutely required for cell growth when the TOR signaling pathway is impaired.

Impaired TOR Signaling Alters InsP Levels

Our results showed a functional interaction between TOR kinase signaling and InsPs; however, they did not indicate whether one signaling pathway is dependent on the other or if they both converge on a common target(s). While tools for direct measurement of TOR kinase activity (i.e., confirmed phosphorylation substrates) are not currently available for Chlamydomonas, we could measure the impact of impaired TOR signaling on intracellular InsP levels. We grew wild-type and vip1-1 cultures under mixotrophic conditions (TAP) and measured InsPs after rapamycin addition for 1 and 12 h. One hour is the earliest time point when growth is measurably reduced by rapamycin, and by 12 h, wild-type cells have begun to acclimate to rapamycin, while vip1-1 mutants are still growth arrested (Figure 3B). For wild-type cells, the response of InsPs to rapamycin involved a transient drop in InsP₅, InsP₆, InsP₇, and InsP₈ at 1 h, with partial recovery after 12 h (Figure 3B). On the other hand, InsP₅, and to some extent InsP₆, became elevated over basal levels in the wild type after 12 h in rapamycin (Figure 3B). For rapamycin-treated vip1-1, we saw no change in InsP₅, but major reductions in other InsP species, including InsP₆, that was reduced by more than 10-fold and a near complete absence of residual InsP₅ and InsP₆ (Figure 3B). Taken together, our results show that impaired TOR signaling leads to changes in InsP levels and suggests that InsPs are downstream of TOR signaling. Moreover, when TOR signaling is impaired in vip1-1 cells, InsP₅ and InsP₆ become nearly undetectable and growth arrests completely, suggesting that these InsPs may play an essential role in cell growth. Additional studies will be required to test whether InsPs also act upstream of TOR kinase to influence its activity.

Interaction between vip1-1 and TOR Signaling Is Carbon Source Dependent

When supplemented with acetate, Chlamydomonas cells generally grow faster while simultaneously downregulating rates of photosynthesis (Heifetz et al., 2000; Chapman et al., 2015). Growth of vip1-1, wild-type, and complemented strains on agar plates or in liquid media with and without acetate and rapamycin were compared (Figure 4). While vip1-1 grew at similar rates as
wild-type or complemented strains under both phototrophic (TP media) and mixotrophic (TAP media) conditions without rapamycin (Figures 4A, 4B, and 4D), its response to rapamycin was acetate dependent (Figures 4A, 4C, and 4E). In media with acetate and rapamycin, growth of \( \text{vip1-1} \) was completely arrested (Figures 4A and 4E) and cell viability was reduced by around 20 to 30% (Figures 4F and 4G), while wild-type and complemented strains showed only transiently slowed growth (Figure 4E) and also had a 20 to 30% reduction in viability (Figures 4F and 4G). However, in media without acetate, \( \text{vip1-1} \) responded to rapamycin similarly to wild-type and to complemented strains with slowed but not arrested growth (Figures 4A to 4C). Overall, our data suggest that defects associated with acetate metabolism may underlie the \( \text{vip1-1} \) phenotype of hypersensitivity to TOR inhibition.

\[ \text{vip1-1} \text{ Partially Uncouples Neutral Lipid Accumulation from Nitrogen Starvation and Acetate Assimilation} \]

The effect of acetate on \( \text{vip1-1} \) prompted us to assess whether the two major storage carbon pools in Chlamydomonas, neutral lipids and starch, were altered in the mutant. For typical wild-type strains, Nile Red stained lipid bodies are most readily detectable in acetate-containing media when cells have been starved for...
nitrogen (TAP -N) (Figure 5A). By contrast, in vip1-1 strains, we always observed lipid bodies, even under conditions where few or no lipid bodies were detectable in wild-type cells, such as in actively growing cultures with nitrogen-replete media (TP and TAP) (Figure 5A). Quantitative measurements of total lipids showed increased levels in vip1-1 strains compared with the wild type under the same growth conditions as described above with significant restoration of wild-type levels in complemented strains. Focusing on mixotrophic conditions (TAP), we found significant reductions of citrate, aconitate, malate, and acetyl-coA in vip1-1 (Figure 7, dark-blue and dark-red bars). vip1-1 cells also responded differently from the wild type to phototrophic conditions where we found 2-fold or greater increases in aconitate, succinate, and fumarate (Figure 7, light-blue and pink bars). As expected, wild-type cells responded to acetate with alterations in levels of TCA cycle intermediates (Heifetz et al., 2000; Boyle and Morgan, 2009; Chapman et al., 2015), but vip1-1 does not undergo the same responses, indicating potential alterations in TCA flux in this mutant that may be tied to altered lipid metabolism and its other phenotypes.

Altered TCA Metabolism in vip1-1 and Modulation of InsPs by Carbon Source

The altered lipid accumulation phenotypes of vip1-1 suggest that the mutant may also have underlying changes in carbon metabolism. Altered metabolite levels can point to altered flux in a corresponding pathway, though they do not provide direct information on rate changes. We focused on the TCA cycle as it is the part of central carbon metabolism most proximal to lipid biosynthesis and to acetate utilization (Figure 7). We grew the wild type and vip1-1 with and without acetate (TAP and TP media) and compared TCA metabolite levels between strains and growth conditions. Focusing on mixotrophic conditions (TAP), we found significant reductions of citrate, aconitate, malate, and acetyl-coA in vip1-1 (Figure 7, dark-blue and dark-red bars). vip1-1 cells also responded differently from the wild type to phototrophic conditions where we found 2-fold or greater increases in aconitate, succinate, and fumarate (Figure 7, light-blue and pink bars). As expected, wild-type cells responded to acetate with alterations in levels of TCA cycle intermediates (Heifetz et al., 2000; Boyle and Morgan, 2009; Chapman et al., 2015), but vip1-1 does not undergo the same responses, indicating potential alterations in TCA flux in this mutant that may be tied to altered lipid metabolism and its other phenotypes.

Modulation of Intracellular InsPs in Response to Growth Conditions and TOR Signaling

The predicted products of VIP1, InsP7, and InsP8 are potential signaling molecules whose levels are perturbed in vip1-1 strains. Moreover, this perturbation impacts central carbon metabolism and lipid accumulation and appears to do so acutely when TOR kinase signaling is also inhibited. These observations suggest that InsPs may be used in Chlamydomonas as signals for responding
to and/or controlling carbon metabolism. If so, the InsP profiles, including the products of VIP1 (InsP₇ and InsP₈), would be expected to change according to growth conditions such as presence/absence of acetate. Indeed, we found that in wild-type cells there were significant changes in InsPs in response to acetate. Wild-type InsP₃ levels were 10-fold lower in TAP versus TP, while levels of InsP₆, InsP₇, and InsP₈ behaved in an opposite manner with moderately higher levels in TAP (Figure 8A). Thus, InsP profiles of wild-type cells, including the predicted VIP1 kinase substrate (InsP₆) and VIP1 kinase products (InsP₇, InsP₈), were modulated by the presence and absence of external acetate.

vip1-1 cells also responded to the presence and absence of acetate but had significant differences from wild-type cells. In the case of InsP₃, vip1-1 cells had near wild-type levels in acetate-containing media but did not show the same degree of elevation in media without acetate (Figure 8A). As expected, the levels of InsP₇ and InsP₈ in vip1-1 were severely decreased under all conditions compared with the wild type. On the other hand, the levels of InsP₆ were much lower in vip1-1 cells grown in TP versus TAP, while in wild-type cells InsP₆ levels changed only modestly between media with and without acetate. In summary, InsPs in Chlamydomonas, including predicted VIP1 kinase substrates and products, showed

Figure 4. Carbon Source-Dependent Growth Inhibition of vip1-1 by Rapamycin.

(A) Growth of the indicated strains (see Fig. 1A) spotted onto agar plates with and without acetate (TAP and TP, respectively) in the absence or presence of 500 nM rapamycin (+rap).

(B) to (E) Growth curves from liquid cultures of strains in (A).

(F) and (G) Viability of the wild type, vip1-1, and vip1-1+pOL-VIP1 in the presence and absence of rapamycin. Colony-forming units (CFUs) were measured at 0, 12, and 24 h and used to calculate viability (CFUs/number of cells plated). Error bars indicate SD from two biological replicates and three technical replicates.
complex acetate-dependent changes in accumulation profiles in a manner that is consistent with InsPs acting as signals for growth metabolism and acetate utilization. Moreover, the same set of InsPs whose levels changed most in response to carbon source in wild-type cells, including InsP_7 and InsP_8, were the most affected by the vip1-1 mutation. Together, our results support a role for InsPs as intracellular signals governing carbon metabolism in Chlamydomonas (Figure 8B).

DISCUSSION

Chlamydomonas VIP1 Provides an Entrée into Inositol Polyphosphate Signaling in Green Algae

Higher order InsPs and the kinases that synthesize them are largely conserved across eukaryotes (Irvine and Schell, 2001; Raboy, 2003; Williams et al., 2015). In fungi, the roles of InsPs have been dissected using mutants that block different steps in the biosynthetic pathway. InsP_7 and InsP_8 are produced by VIP1 and/or KCS1 kinases whose animal homologs are PP-InsP_5K and InsP_6K, respectively, and play diverse roles in cell physiology, including signaling phosphate starvation, telomere length control, DNA repair, autophagy, environmental stress responses, cytoskeletal dynamics, and ribosome biogenesis (Bennett et al., 2006; Shears, 2009; Tsui and York, 2010; Wundenberg and Mayr, 2012; Wilson et al., 2013; Thota and Bhandari, 2015; Livermore et al., 2016).

In plants, InsPs are of special interest because InsP_6 (phytic acid) is agronomically important as a high-abundance phosphate storage compound in seeds (Raboy, 2003), and other InsPs have recently been investigated for potential roles in hormone signaling and plant defense (Tan et al., 2007; Mosblech et al., 2008, 2011; Sheard et al., 2010; Laha et al., 2015). Green algae and land plants encode enzymes required to successively convert InsP_3 to InsP_7 and InsP_8 (Figure 2B) (Loewus and Murthy, 2000; Raboy, 2003; Gillaspy, 2013; Williams et al., 2015), though they are missing the KCS1/InsP_6K family that pyrophosphorylates InsP_6 or InsP_7 on...
position 5 of the inositol ring, an activity which is distinct from the activity of VIP proteins that act on the 1/3 positions (Lin et al., 2009) (Figure 2C). The two VIP paralogs of Arabidopsis (AtVIP1 and AtVIP2; also called AtVIH1 and AtVIH2) can complement a yeast vip1 deletion (Desai et al., 2014) and are required for jasmonate defense signaling (Laha et al., 2015). A QTL for vitamin E levels also mapped to one of the two Arabidopsis VIP genes (At3g01310) (Gilliland et al., 2006), though a connection between InsPs and vitamin E has not been directly established. Other metabolic phenotypes in plants defective for VIP genes have not been reported; however, interaction between InsP7 and SPX domain proteins including regulators of phosphate homeostasis have been recently found (Wild et al., 2016). We note that the media used in our study contain excess inorganic phosphate making phosphate metabolism unlikely to underlie the growth and metabolic defects we observed in vip1-1.

InsPs in green algae were previously examined in two species of Chlamydomonas, C. reinhardtii and C. eugametos (also called C. moewusii), that are quite distantly related (Pröschold et al., 2001). In both species, InsP3 appears to play a conserved role as a second messenger and effector of cytosolic calcium influx that triggers deflagellation (Yueh and Crain, 1993; Quarmby and Hartzell, 1994; Munnik et al., 1998) and is also linked to light-dependent expression of GSA, a gene of the heme/chlorophyll biosynthetic pathway (Im and Beale, 2000). InsP kinase activities that can produce InsP4, InsP5, and InsP6 have been found in extracts from C. eugametos (Irvine et al., 1992), but no roles for these higher order InsPs or evidence of pyrophosphorylated InsPs have been previously reported in green algae. Our identification of a mutation in the C. reinhardtii VIP1 gene that causes defects in InsP7 and InsP8 accumulation provides a tool for dissecting the roles of VIP proteins and pyrophosphorylated InsPs in green algae with possible relevance for other eukaryotic

Figure 6. Increased TAG and Decreased Galactolipids in vip1-1.

(A) TLC separation and iodine staining of different lipid classes from total lipids of wild-type and vip1-1 cells growing in TAP, TAP supplemented with rapamycin for 12 h (+rap), or transferred to TAP-N for 12 h. The left-most lane contains a mixture of standards (Mixstd) whose constituents are abbreviated as follows: PE, phosphatidyl ethanolamine; MAG, monacylglycerol; DAG, diacylglycerol; FFA, free fatty acids; TAG, triacylglycerol. The identification of β-carotene in the fastest migrating position was done separately using a purified standard. Blue arrowhead on right indicates migration of TAGs.

(B) Quantitation of FAMES derived from purified TAGs and normalized by dry cell weight for strains shown in (A). Errors bars indicate so from five replicates.

(C) Relative distribution (percent total) of acyl group types from TAG fraction of wild-type and vip1-1 strains.

(D) and (E) Percentage of FAMES in galactolipid and phospholipid fractions purified from the wild type and vip1-1 and normalized by dry cell weight. Error bars indicate so from at least three biological replicates. Asterisks represent significant differences (P < 0.05) between means evaluated using Student’s t test.
species, including plants that also contain VIP proteins but no KCS1/InsP6K-like proteins (Bennett et al., 2006; Desai et al., 2014; Williams et al., 2015). Despite vip1-1 being a loss-of-function allele, the mutants still have some detectable InsP7 and InsP8 (Figures 3 and 8), indicating that there may be another route for InsP7 and InsP8 synthesis in these cells. A second predicted VIP gene, VIP2 (Figures 2B and 2C), is the best candidate for producing residual InsP7 and InsP8 in vip1-1.

**InsPs as a Metabolic Signal for Carbon Metabolism**

Much attention has already been focused on the metabolic pathways of Chlamydomonas that are responsible for accumulation of TAGs as these are important end products for biofuel and biotechnology applications (Hu et al., 2008; Merchant et al., 2012; Liu and Benning, 2013; Goodenough et al., 2014). Previous studies have assessed the responses of metabolic genes and proteins under non-steady state conditions when TAG accumulates (Miller et al., 2010; Boyle et al., 2012; Msanne et al., 2012; Blaby et al., 2013; Wase et al., 2014) or identified mutants that are defective for TAG accumulation or catabolism (Li et al., 2012; Boyle et al., 2012; Tsai et al., 2014; Xie et al., 2014; Ngan et al., 2015; Kajikawa et al., 2015). The relationship between starch synthesis and TAGs has also been investigated, and it appears that fixed carbon can be shunted toward TAGs when starch synthesis is blocked (Li et al., 2010a, 2010b; Work et al., 2010; Siaut et al., 2011; Goodenough et al., 2014). One recently identified regulatory mutant, tar1, is defective in accumulation of lipid bodies in response to nutrient starvation, and the TAR1 locus was shown to encode a YAK subtype DYRK family protein kinase (Kajikawa et al., 2015). The transcription factors PSR1 and NRR1 have also been postulated as regulators for lipid accumulation during nitrogen starvation (Boyle et al., 2012; Ngan et al., 2015), and the transcription factor CHT7 is required for TAG remobilization when starved cells are returned to nutrient-replete media (Tsai et al., 2014). To date, no putative regulatory mutants have been identified that uncouple TAG accumulation from nitrogen starvation or acetate supplementation as we observed in vip1-1, nor have regulatory mutants been described that cause increased TAG and lipid overaccumulation compared with wild-type cells.

**Figure 7. Altered Profiles of TCA Cycle Metabolites in vip1-1.**

Diagram of TCA cycle with bar graphs showing levels of metabolites that were measured in the wild type and vip1-1 growing mixotrophically in TAP or phototrophically in TP. Solid black arrows show the canonical respiratory TCA cycle where carbon is lost as CO2; gray dashed arrows show the glyoxylate cycle where carbon is conserved for anabolic metabolism; y axes are measured in nmol/gFW. Error bars indicate SD from at least three biological replicates. Asterisks represent significant differences (P < 0.05) between means evaluated using Student’s t test.
InsPs structural isomers (Irvine and Schell, 2001; Siaidi, 2012) and their rapid turnover could make them highly responsive signals for metabolic control.

Although vip1-1 mutants do not appear to suffer from any acute metabolic tradeoffs, there are likely to be fitness defects in this mutant that emerge under specific conditions that have not been tested here. Importantly, the metabolic alterations in vip1-1 that lead to increased TAG levels appear to be synergistic with other conditions that cause increased TAG accumulation such as nitrogen starvation or rapamycin treatment (Figures 6A and 6B). It remains to be determined whether the normal responses to stress and starvation are accelerated in vip1-1 or if the mutant activates a parallel pathway for storage lipid accumulation. Two recent studies implicated the TCA pathway in carbon utilization and lipid accumulation. A Chlamydomonas mutant missing isocitrate lyase, a key enzyme in the glyoxylate shunt within the TCA cycle, was found to have acetate-specific growth defects and to over-accumulate lipids (Plancke et al., 2014). A metabolic modeling study also found that changes in TCA flux were required to maintain metabolic balance when switching from phototrophic to mixotrophic growth (Chapman et al., 2015). Our metabolite profiling results (Figure 7) further suggest that accumulation of lipids in vip1-1 may be coupled to altered flux through the TCA cycle.

A Connection between Inositol Polyphosphates and the TOR Signaling Pathway in Chlamydomonas Is Revealed by vip1-1

Interactions between TOR signaling and InsPs in yeast and animals have been suggested by several studies (Chakraborty et al., 2010; Kim and Guan, 2011) including roles for InsP kinases upstream (Nagata et al., 2005) and downstream of TOR (Worley et al., 2013). These findings are not mutually exclusive, but they do highlight the potential complexity of intracellular signaling pathways where feedback loops can connect signaling systems at different levels. To date there has been no connection made between TOR signaling in photosynthetic eukaryotes and InsPs. Interestingly, we found that rapamycin inhibition, even in wild-type cells, caused changes in InsP levels, meaning that at least part of the rapamycin hypersensitivity we see in vip1-1 could be caused by defects in InsPs that act as effectors of TOR signaling (Figures 3B and 8B). The apparently complete elimination of detectable InsP2 and InsP8 in rapamycin-treated vip1-1 cells is suggestive of an essential role for these InsPs in cell growth. We attempted to assess the effect of vip1-1 on TOR kinase signaling using a commercial antibody that detects S6K (ribosomal protein S6 kinase) phosphorylation by TOR on a conserved site (Thr-389 in human S6K and Thr-932 in Chlamydomonas) and that has been used successfully in Arabidopsis (Xiong et al., 2013) but were unable to detect a specifically recognized cross-reacting protein in Chlamydomonas extracts. Nonconservation with the human S6K phospho-epitope at the −3 and −5 positions with respect to T389 may be partly responsible for this lack of cross-reactivity. Until TOR kinase activity assays are developed for Chlamydomonas, we cannot test the possibility that InsPs are also upstream of TOR signaling and/or function in a parallel pathway (Figure 8B). While the relationships between TOR and InsPs remain to be fully elucidated, the synergism between the two signaling systems that we uncovered may have relevance in other eukaryotes where both are conserved.
METHODS

Chlamydomonas reinhardtii Strains, Culture Conditions, and Viability Test

Strain CC-1690 wild-type MT+ (Sager 21gr) (Sager, 1955) was used as parental strain to generate vip1-1. CC-1691 (Sager 6145c) wild-type MT− was used for segregation/linkage analysis and diploid generation described below. An fkb1 mutant (originally named rap2) contains a disruption in the Cre13.588600 locus and was kindly provided by Jose-Luis Crespo (Crespo et al., 2005). All cultures were maintained on TAP (Tris acetate phosphate) agar plates (Harris, 2009b). TP was prepared as TAP but omitting acetic acid and replacing Tris-base with Tris-HCl, pH 7.4. N-free media was prepared by omitting NH₄Cl. Liquid cultures were grown in 500-mL flasks at 25°C in temperature controlled water baths illuminated continuously from below with ~150 µE m⁻² s⁻¹ LED lights [1:1 fluence ratio red (625 nm) and blue (465 nm)] and bubbled with air or air enriched with 0.5% CO₂ (v/v). Except where mentioned, experiments were done using cultures in exponential phase (1 to 2 × 10⁵ cells/mL) that were maintained in exponential phase through frequent dilution (at least once per day). N-starved cultures were prepared by centrifuging and washing cells twice in the N-free media (TAP-N or TP-N) and then resuspending them in N-free media. Cultures were monitored for cell size and cell concentration using a Multisizer 3 Coulter Counter (Beckman).

Viability experiments were performed using cultures pregrown to mid- to late-exponential phase (2 to 8 × 10⁶ cells/mL) and then diluted into the same media with or without rapamycin to a density of ~2 × 10⁶ cells/mL. Rapamycin (LC Laboratories) was added to a final concentration of 500 nM from a 50 mM stock in DMSO. Viability for each sample was calculated by dividing the number of colony-forming units determined by plating on agar plates by the cell density measured in a Coulter Counter.

Mutant Screening

Insertional mutagenesis (Galván et al., 2007) with a hygromycin resistance gene, aph7′ (Berthold et al., 2002), was used to generate mutants using wild-type parental strain CC-1690. Transformation was performed using electroporation modified from a published method (Shimogawara et al., 1998), as follows: 300 mL cultures in TAP were grown to a density of ~1 × 10⁸/mL. Cells were harvested by centrifugation (4000 rcf, 5 min, room temperature) and resuspended at 3.5 × 10⁷/mL in TAP with 50 mM sorbitol. Next, 1 µg DNA and 300 µL of the cell suspension were placed into 4-mm cuvettes (Bio-Rad) prechilled on ice for 15 min, and then electroporated using a GenePulsar Xcell (Bio-Rad), with settings of 800 V, 25 µF, and no resistance. Transformants were recovered in 10 mL TAP+50 mM sorbitol in low light for 16 h and then concentrated and spread on two TAP agar plates supplemented with 30 µg/mL hygromycin for selection. Transforming DNA was a 1.7-kb gel-purified aph7′ expression cassette (Berthold et al., 2002) digested from plasmid pHyg3 with HindIII (New England Biolabs). After 4 to 5 d, colonies were picked into 96-well plates containing TAP, grown to saturation, and then pinned onto TAP and TAP + rapamycin plates. Rapamycin-hypersensitive mutants were identified by comparing growth on TAP versus TAP + 500 nM rapamycin. AZD8055 (LC Laboratories) and Torin1 (LC Laboratories) were used at 500 nM final concentration diluted from a 50 mM stock in DMSO.

Genetic Analysis

Gamete preparation, mating, and segregation analyses were done as described previously (Harris, 2008b). Random meiotic progeny isolated from among thousands of zygote colonies were individually grown in 96-well microtiter plates and pinned onto agar media plates supplemented with hygromycin or rapamycin. Mating type was scored using PCR genotyping (primers presented in Supplemental Table 1) (Zamora et al., 2004). Vegetative diploid strains (Ebersold, 1967) were generated usingvip1-1 and/or wild-type parental strains of the opposite mating type transformed with either a paromomycin resistance marker containing plasmid, pSI103 (Sizova et al., 2001) or a zeocin resistance marker containing plasmid, pSP124S (Lumbreras et al., 1998). Gametes of opposite mating type and containing different resistance markers were spread onto TAP agar containing 20 and 15 µg/mL zeocin shortly after mixing (1 to 2 h) and vegetative diploid colonies were picked and confirmed using PCR-based genotyping (Supplemental Table 1) (Zamora et al., 2004).

vip1-1 Mapping and Genotyping

Amplification of the pHYG3 insertion junction was performed according to a previously described strategy (O’Malley et al., 2007) using genomic DNA prepared as described (Umen and Goodenough, 2001). Genomic DNA from 266F3 was digested with PvuII, Smal, Stul, and PmlI and blunt-end ligated to annealed adaptor oligonucleotides (Supplemental Table 1). Junction fragments were amplified using nested reactions with adaptor primers AP1 and AP2 along with IMP5h-1 and IMP5h-2, or with IMP3h-1 and IMP3h-2 (Supplemental Table 1). Cloned junction fragments were sequenced and the left border confirmed by amplifying and sequencing a 426-bp fragment amplified with IMPVIFP3 and IMPVIP3R (Supplemental Figure 1 and Supplemental Table 1).

Genotyping was done using 0.5 µL of total genomic DNA prepared using a fast method described at http://www.chlamydocollection.org/methods/quick-and-easy-genomic-dna-prep-for-pcr-analysis/. The VIP1 wild-type allele was scored using VIPWTL and VIP1WTR (Supplemental Figure 1 and Supplemental Table 1), and the vip1-1 allele was scored using VIPWTL and IMPVIPR. PCR was performed using Taq polymerase in a final reaction volume of 20 µL in the presence of 1× Taq buffer (Sambrook and Russell, 2001), 1 µM primers, 80 µM dNTP, 2% DMSO, and 0.5 µL of genomic DNA. PCR conditions were as follows: 96°C for 2 min and then 35 cycles of 94°C for 30 s, 60°C for 30 s, and 72°C for 30 s.

RNA extraction, cDNA synthesis, and RT-PCR were performed as previously described (Fang and Umen, 2008). VIPWTL and VIP1WTR (Supplemental Figure 1 and Supplemental Table 1) were used to amplify cDNAs from the VIP1 locus in the flank region where pHYG is inserted invip1-1 (Supplemental Figure 1). PCR conditions were the same as above.

Cloning and Epitope Tagging of CrVIP1

Chlamydomonas BAC clone 4C23 (Kim et al., 1996) (http://www.genome.clemson.edu) was digested with PsiI and EcoRI and a 14628-bp fragment containing VIP1 was gel purified. The VIP1 fragment was ligated to psi103 (Sizova et al., 2001) digested with SmaI and EcoRI to generate pVIP1. An OLLAS epitope (Park et al., 2008) was inserted into the N terminus of VIP1 as follows. Two fragments (A and B) were amplified from VIP1, fragment A with primers OLLVIP1F and OLLVIP1R, and fragment B with primers OLLVIP2F and OLLVIP2R (Supplemental Table 1). OLLVIP1R and OLLVIP2F share an overlapping sequence that encodes the OLLAS tag sequence (Supplemental Table 1, underlined) and homology with VIP1. Both fragments were amplified using Taq polymerase as described above. pVIP1 and fragments A and B were digested with PfoI (Thermo Scientific) (Supplemental Table 1, lowercase) and recombined using an In-Fusion HD Cloning Kit (Clontech) according to the manufacturer’s instructions to generate pOLLAS-VIP1.

Complementation of vip1-1

vip1-1 was transformed by electroporation as described above and colonies were selected on TAP plates containing 30 µg/mL paromomycin. Transformants were picked and grown in 96-well plates in TAP until growth saturation and then they were pinned onto TAP containing 500 nM rapamycin. Positive clones were identified by loss of rapamycin resistance.
hypersensitivity with growth rates comparable to wild-type strains on rapamycin-containing plates. Complementation rates were calculated after testing paromomycin resistant colonies in rapamycin-containing media. Complementation rates were 15/480 for pOL-VIP1 and 2/80 for pVIP1.

Immunoblotting OLLAS-VIP

Protein extraction and immunoblotting to detect pOLLAS-VIP were performed as described previously (Olson et al., 2010) with the following modifications. Total protein loaded per well was 40 μg; rat α-OLLAS antibodies (NB1-P-06713; Novus Biologicals; Lot A) were used at 1:1000 and HRP-conjugated secondary α-rat antibodies (Thermo Fisher Scientific; 31470) were used at 1:4000.

InsP Extraction and Analysis

The method described below allows separation and relative quantitation of InsP mass isomers InsP3, InsP4, InsP5, InsP6, and InsP7. We cannot currently differentiate between InsP stereoisomers, but methodology to do so using mass spectrometry is under development. For InsP extraction, 300 mL of mid-log phase culture per sample was collected at a cell density of ~2 × 10^9 cells/mL. (Cell density) × (mean cell size) measured using a Beckman–Coulter Multisizer3 was used as a metric for wet biomass. Sample volumes were adjusted slightly where necessary so that all replicates had identical wet biomasses. After harvesting by centrifugation (4000 rcf, 5 min, room temperature), cell pellets were resuspended in 1 mL final volume 5% trichloroacetic acid to extract InsPs and flash frozen in liquid nitrogen. After thawing, the samples were centrifuged at maximum speed for 20 min in a microfuge at 4°C. Each supernatant was then supplemented with 100 μL of 1 μM 3-fluoro-InsP3 (3-deoxy-3-fluoro-3-O-myo-inositol 1,4,5-triphosphate) (Enzo Life Sciences), which served as an internal standard for normalization. Supernatants (33-mL particle size). The column was washed with 1 mL of 25% methanol and then a ramp to 100% A over 6 min, then up to 60% A in 7 min, and finally a ramp to 100% A over 3 min, followed by a hold at 100% A for 4 min. The solvent composition was returned to 100% B over 6 min and the column was reequilibrated for 40 min for a total per sample run time of 70 min. Mass spectrometric data were collected in negative ion mode at a resolution of 70,000 (at m/z 200) with a full scan from m/z 190 to 850 and SIM scans of 483.9 to 513.9, 563.9 to 593.9, 643.9 to 673.9, 723.9 to 753.8, and 803.9 to 833.8. Data were analyzed using the QualBrowser and QuanBrowser applications of Xcalibur (Thermo Fisher Scientific). Data were normalized using the internal standard 3-fluoro-InsP3. Mean data and so were calculated from three biological replicates, each of which had three technical replicates.

Metabolite Analysis

Mid-log phase (1.5 × 10^6 mL^-1) cultures were harvested by fast vacuum filtering onto 50-mm Whatman filter paper (GE Healthcare Life Sciences; 1822-020), which was immediately quenched in liquid nitrogen with the total processing time for each sample kept under 10 s. Fresh weight (FW) for each sample was calculated by determining wet biomass volume (culture density × mean cell size × volume harvested). Packed cells have a density close to 1 mg/mL; therefore, FW in milligrams was equivalent to pellet volume in milliliters. Polar metabolites in frozen samples (5 to 8 mg FW) were extracted with methanol/chloroform/water (7:3:5.6) and analyzed via LC-MS/MS as previously described (Ma et al., 2014) with 0.3 nmole PIPES added to each sample as an internal standard. Metabolite concentrations (nmol gFW^-1) were determined by interpolating from the linear relationship between peak area and the concentrations of standards.

Lipid Analysis

Gas Chromatography-Flame Ionization Detection

Total lipid extraction and fatty acid methyl esters (FAMEs) were obtained from 5 mg of Chlamydomonas freeze-dried pellets using the method reported in Han et al. (2010) and using triheptadecanoin (C17:0 TAG) as an internal standard (Nu-Chek Prep). After evaporation under nitrogen gas, the samples were resuspended in 200 μL hexane and 1 μL was separated by Trace-GC with an FID detector (Thermo Fisher Scientific) fitted with an HP-INNOWax column (0.25-mm inner diameter and 30-m length) (Agilent). The oven temperature was programmed to 100°C (hold 1 min) to 235°C (hold 5 min) in a first ramp of 40°C/min until 185°C and then a second ramp of 7.5°C/min to 235°C. Peak assignments were made using FAME mix C8-C22 (Supelco) and the double-bond positions of the polyunsaturated fatty acids were confirmed by gas chromatography-mass spectrometry (Cahoon et al., 2006).

TLC

Twenty milligrams of freeze-dried pellet was used for extraction of total lipids using a modification of the standard Bligh and Dyer method (Bligh and Dyer, 1959). Four milliliters of CHCl3:methanol (2:1) was added to the sample that was then mixed by vortexing. The samples were heated at 42°C for 30 min followed by addition of 2.5 mL 0.1 n HCL 1 M NaCl and additional mixing by vortexing. Samples were centrifuged for 2 min at 500 rcf at room temperature and then the aqueous (upper) phase was discarded. The organic phase was washed twice with ultrapure water and then dried under nitrogen gas. Samples were resuspended in 1 mL hexane and split into three equal (333 μL) parts. One part was trans-methylated to FAMEs and analyzed by gas chromatography-flame ionization detection (GC-FID) as described above. A second part was used for TLC separation as described by Han et al. (2010) with image acquisition using an Alphaimager 2200 (AlphaInnotec). A mixture of standards was prepared using 18:1 free fatty acid, 18:1 triacylglycerol (TAG), 16:0/18:1 diacylglycerol, and 18:1 monoacylglycerol, 30 μg each, in 200 μL hexane. The third part was used for TAG separation on a silicic acid column with latex beads of 0.5-cm internal diameter (Mitsubishi Kagaku Iatron). Columns were prepared using disposable glass Pasteur pipettes stopped with untreated glass wool and exactly 2 cm of beads. Columns were equilibrated with 4 mL hexane and then 200 μL of sample was loaded onto the column and TAGs were eluted with 4 mL of chloroform. Galactolipids and phospholipids were then eluted using 4 mL of acetone and methanol, respectively. After elution, all the fractions were dried and redissolved in hexane followed by fractionation on TLC plates or analysis by GC-FID, as described above.

Starch Analysis

Starch was measured using a Total Starch Assay Kit (AA/AMG; Megazyme) following the manufacturer’s instructions, but scaled down to 10 mg freeze-dried cell powder as starting material. Mean and so were calculated...
from at least three biological replicates with three technical replicates per sample.

**Fluorescence Microscopy**

**Nile Red Staining**

Lipid body staining was performed as described (Wang et al., 2009). Microscopy was performed with a Leica DMi6000B using a 63× oil immersion objective with differential interference contrast (DIC) optics or wide-field fluorescence collected using a Leica L5 filter cube (excitation BP 480/40; dichroic 505; emission BP 527/30).

**Immunofluorescence**

Immunofluorescence was performed as described previously (Olson et al., 2010) with the following modifications. Primary rat α-OALLS antibody (NB1P-06713, Lot A; Novus Biologicals) was used at 1:100 and secondary α-mouse (Alexa Fluor 488, A-11001, Lot 99C2-2; Thermo Fisher Scientific) was used at 1:1000, as recommended by the manufacturer. Microscopy was performed with a Leica DMi6000B using a 100× oil objective lens with DIC optics or using the following filter cube sets A4: excitation BP 360/40; dichroic 400; emission BP470/40; L5: excitation BP 480/40; dichroic 505; emission BP 527/30. Z stacks through the entire cell were collected for each image (20 × 0.2 μm).

**Phylogenetic Tree Construction**

Accession numbers for VIP1 homolog protein sequences used for alignment and tree construction are listed below. MEGA (Tamura et al., 2013) was used for all tree building steps. Sequences were aligned using MUSCLE (Edgar, 2004). All positions containing gaps and missing data were removed and an unrooted neighboring-joining tree was constructed using the JTT distance model with gamma set to 0.8 and 1000 bootstrap replicates.

**Accession Numbers**

Sequence data from this article can be found for Chlamydomonas genes in Phytozome (https://phytozome.jgi.doe.gov/pz/portal.html) under the following gene IDs: PKB1, Cre13.g868300; CVIP1, Cre03.g183500; CVIP2, Cre01. g0302600; CreTPK1, Cre03.g182100; CreTPK2, Cre01.g018400; CrIPM, Cre12.g555400; CrIP5-2K, Cr04.g229000; and CrVIP1, Cre02.g106500. The remaining sequences can be found at NCBI Protein database or GenBank (http://www.ncbi.nlm.nih.gov). Protein database accession numbers are as follows: Volvox carteri VcVIP1: XP_002485782.1, VcVIP2: XP_002486745.1, Coccomya subellipsosidea CsVIP1: XP_005644985.1; Micromonas pusilla MpVIP1: XP_003058868.1; Ostreococcus tauri OtVIP1: XP_003078214.1, Ostreococcus lucimarinus OlVIP1: XP_001416675; Oryza sativa OsVIP2: NP_001050945; Arabidopsis thaliana AtVIP1: NP_15811280.3 and AtVIP2: NP_001190313.1; Homo sapiens HsVIP1: NP_001243349.1; and HsVIP2: NP_001263206; Thalassiosira pseudonana CCMP1335 TpVIP1: XP_002278804; Pheoadoctylum tricornutum CCAP 1055/1 PtvIP1: XP_002185836.1; Ustilago maydis UmVIP1: XP_011392692; Aspergillus nidulans FGSC A4 AnVIP1: XP_663401.1; Caenorhabditis elegans CeVIP1: NP_740855.2; Schizosaccharomyces pombe SpVIP1: NP_587877.1; Saccharomyces cerevisiae ScVIP1: NP_013514; GenBank; Oryza sativa OsVIP1: EEE55479.1; Drosophila melanogaster DmVIP1: AAT94524.1.

**Supplemental Data**

**Supplemental Figure 1.** vip1-1 insertion structure, genotyping, and dominance testing.

**Supplemental Table 1.** Sequence of primers used for mapping and genotyping VIP1 or vip1-1 and for constructing pOL-VIP1.

**Supplemental Data Set 1.** Alignment used to generate the phylogeny presented in Figure 2B.

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**AUTHOR CONTRIBUTIONS**


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