Inositol polyphosphates regulate and predict yeast pseudohyphal growth phenotypes

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Abstract

Pseudohyphal growth is a nutrient-regulated program in which budding yeast form multicellular filaments of elongated and connected cells. Filamentous growth is required for virulence in pathogenic fungi and provides an informative model of stress-responsive signaling. The genetics and regulatory networks modulating pseudohyphal growth have been studied extensively, but little is known regarding the changes in metabolites that enable pseudohyphal filament formation. Inositol signaling molecules are an important class of metabolite messengers encompassing highly phosphorylated and diffusible inositol polyphosphates (InsPs). We report here that the InsP biosynthesis pathway is required for wild-type pseudohyphal growth. Under nitrogen-limiting conditions that can induce filamentation, InsPs exhibit characteristic profiles, distinguishing the InsP₇ pyrophosphate isoforms 1PP-InsP₅ and 5PP-InsP₅. Deletion and overexpression analyses of InsP kinases identify elevated levels of 5PP-InsP₅ relative to 1PP-InsP₅ in mutants exhibiting hyper-filamentous growth. Overexpression of KCS1, which promotes formation of inositol pyrophosphates, is sufficient to drive pseudohyphal filamentation on medium with normal nitrogen levels. We find that the kinases Snf1p (AMPK), Kss1p, and Fus3p (MAPKs), required for wild-type pseudohyphal growth, are also required for wild-type InsP levels. Deletion analyses of the corresponding kinase genes indicate elevated InsP₃ levels and an absence of exaggerated 5PP-InsP₅ peaks in trace profiles from snf1Δ/Δ and kss1Δ/Δ mutants exhibiting decreased pseudohyphal filamentation. Elevated 5PP-InsP₅:1PP-InsP₅ ratios are present in the hyperfilamentous fus3 deletion mutant. Collectively, the data identify the presence of elevated 5PP-InsP₅ levels relative to other inositol pyrophosphates as an in vivo marker of hyper-filamentous growth, while providing initial evidence for the regulation of InsP signaling by pseudohyphal growth kinases.

Author summary

Changes in metabolite levels underlie important biological processes, including cellular responses to nutrient stress. One such response encompasses the nitrogen stress-induced
transition of budding yeast cells into multicellular filaments, relevant as a model of directional growth and fungal pathogenesis. We report here that a conserved family of charged lipid-derived metabolites, inositol polyphosphates, exhibits characteristic changes as yeast cell form filaments in response to conditions of nitrogen limitation. The ratios of doubly charged inositol pyrophosphates consistently match with the degree of filament formation. Enzymes of the inositol polyphosphate synthesis pathway are required for filament formation, and inositol polyphosphate levels are dependent on kinases that enable wild-type filamentation. Our data indicate that inositol polyphosphates mark filamentous growth states, highlighting a new regulatory role for these ubiquitous eukaryotic second messengers.

Introduction

The transition from unicellular yeast to growth in multicellular filaments is characteristic of many fungi [1–4]. In the budding yeast *Saccharomyces cerevisiae* nitrogen limitation or growth in an alternate carbon source induces cell elongation, unipolar budding, and altered cell-cell adhesion, resulting in the formation of pseudohyphal filaments that presumably benefits the organism as a scavenging mechanism [5–7]. Yeast pseudohyphal growth is relevant as a model of polarized growth, and in the related opportunistic human fungal pathogen *Candida albicans*, filamentous development is required for virulence [8, 9]. Classic studies have identified a core set of conserved signaling modules regulating yeast pseudohyphal growth, including the Kss1p mitogen-activated protein kinase (MAPK) cascade, the AMP-activated kinase family member Snf1p, and the Ras2p/cAMP-dependent protein kinase A (PKA) pathway [10–17]. Genomic screens using loss-of-function mutants and overexpression libraries have identified a broader set of genes required for pseudohyphal growth [18–20], but little is known regarding the changes in metabolite levels that underlie the filamentous growth transition.

Inositol polyphosphates (InsPs or IPs) are highly charged, lipid-derived metabolites that act as second messengers in organisms throughout the eukaryotic kingdom [21]. InsP biosynthesis is initiated by the enzyme phospholipase C, which cleaves the six-carbon cyclitol inositol from phosphatidylinositol 4,5-bisphosphate, yielding diffusible inositol 1,4,5-trisphosphate (InsP$_3$) [22]. InsP$_3$ has long been recognized to induce the early mobilization of calcium from the endoplasmic reticulum in many metazoans, although a similar role is not evident in budding yeast [23]. As indicated in Fig 1A, InsP$_3$ is converted into inositol hexakisphosphate (InsP$_6$) through a series of phosphorylation reactions catalyzed by the InsP kinases Arg82p and Ipk1p [24]. InsP$_6$ is the most abundant InsP with levels ranging from 10–100 μM in most eukaryotes [25]. Pyrophosphorylated InsP molecules containing energy-rich diphosphate bonds are generated from InsP$_5$ and InsP$_6$ through the kinase activities of Kcs1p and Vip1p [26, 27]. Inositol pyrophosphate pools are turned over, such that up to 50% of human InsP$_6$ is converted into pyrophosphate molecules per hour. With this turnover rate, inositol pyrophosphate levels are low, constituting 1–5% of InsP$_6$ levels [28, 29]. Inositol pyrophosphate turnover is in part achieved through the action of three InsP phosphatases in yeast: Ddp1p, Siw14p, and Vip1p, which contains a putative phosphatase domain [30–32].

InsPs, and particularly inositol pyrophosphates, have been found to regulate many cellular processes, encompassing telomere maintenance, insulin signaling, cell migration, endocytosis, mRNA export, and chromatin modification [33–39]. Recent studies in yeast and *Arabidopsis* suggest that inositol pyrophosphates respond to cellular inorganic phosphate levels, acting as sensors for the regulation of phosphate homeostasis in eukaryotes [40]. Interestingly, inositol
Pyrophosphates can generate pyrophosphorylated proteins through the enzyme-independent transfer of their β-phosphoryl group onto phosphoprotein targets [41, 42]. These studies highlight the importance of inositol polyphosphates as metabolite messengers, but questions remain regarding the regulation of InsPs and the breadth of their role in cell signaling.
In previous work, we utilized quantitative phosphoproteomics to identify yeast proteins within kinase-based pseudohyphal growth signaling networks [43]. The data identified several InsP kinases, leading us to consider the possibility that inositol polyphosphate signaling may contribute to pseudohyphal growth, and more broadly, to the cellular response to nitrogen limitation. Our analyses here support this notion, indicating that InsP signaling is required for wild-type pseudohyphal growth and that the respective ratios of inositol pyrophosphates, in particular, change stereotypically in filamentous growth mutants, to the point of being predictive of exaggerated pseudohyphal growth states in a strain that is otherwise competent for filament formation. We further find that pseudohyphal growth kinases of the MAPK and AMPK families are required for wild-type levels of InsPs, suggesting a regulatory link between nutrient-responsive signaling and InsP metabolite messengers.

Results

InsP kinases are required for wild-type pseudohyphal growth

We previously used proteome-wide quantitative mass spectrometry to identify proteins differentially phosphorylated in yeast mutants carrying a catalytically defective allele of a kinase required for pseudohyphal growth [43]. Mutant alleles of eight kinase genes were analyzed in the study (elm1-K117R, fus3-K42R, kss1-K42R, snf1-K84R, ste7-K220R, ste11-K444R, ste20-K649R, and tpk2-K99R). The InsP kinase Arg82p and the inositol pyrophosphate kinases Kcs1p and Vip1p were differentially phosphorylated in the kss1-K42R and snf1-K84R mutants under conditions of reduced nitrogen availability (S1 Table). These results led us to further explore a possible role for InsPs in pseudohyphal growth response.

To assess the functional significance of InsPs in pseudohyphal growth, we generated haploid and homozygous diploid yeast deletion mutants in the filamentous S1278b genetic background for each gene encoding a kinase in the InsP biosynthetic pathway. Haploid deletion mutants were assayed for the ability to invade agar, and surface-spread filamentation was characterized in diploid mutants grown on low-nitrogen medium with a reduced concentration of ammonium sulfate (Fig 1B). Cell morphology of the diploid deletion mutants is indicated in S1 Fig. Invasive and surface-spread phenotypes were consistent for each mutant. Mutants deleted for ARG82, which lack InsPs and inositol pyrophosphates downstream of InsP, exhibited decreased haploid invasive and diploid pseudohyphal development. This is consistent with the observation that diploid cells lacking PLC1 do not form pseudohyphae on low-nitrogen media [44]. Strains deleted for IPK1 and VIP1 showed elevated pseudohyphal growth. Deletion of KCS1 resulted in decreased invasion and surface pseudohyphal growth relative to wild type. Considered together, the deletion analyses indicate that the presence of each InsP kinase is required for wild-type pseudohyphal growth and that perturbation of InsP biosynthesis can result in either decreased or increased pseudohyphal filamentation, with the respective phenotype likely reflecting the accumulation of a given InsP species or set of InsPs. Subsequent data presented here are consistent with this conclusion.

InsP profiles under pseudohyphal growth conditions are distinct and distinguish InsP isoforms

To identify the relative levels of InsP species during pseudohyphal growth, we used the approach of Azvedo and Saiardi [45] in which yeast strains are cultured in media with radiolabeled myo-inositol; labeled inositol is taken up by yeast and subsequently metabolized into downstream InsPs and inositol pyrophosphates. InsP levels were profiled by this method in wild-type filamentous yeast (S1278b) grown in normal media and under pseudohyphal...
growth-inducing conditions in low-nitrogen media with reduced ammonium sulfate. Notable characteristics were evident in the profiles under conditions of low nitrogen (Fig 2A). In particular, two peaks corresponding to InsP$_7$ isoforms were present, whereas only one InsP$_7$ peak has classically been reported in strains grown in media with normal levels of ammonium sulfate. In addition, the peak corresponding to InsP$_5$ was absent under low-nitrogen conditions, and an increased and broad trace corresponding to InsP$_3$ was observed. This broad InsP$_3$ trace presumably indicates InsP$_3$ or InsP$_4$ isoforms that we were not able to characterize with our standard deletion strains. To consider if the changes are specific to pseudohyphal growth or represent a more general response to nitrogen limitation, we determined the InsP profile for a non-filamentous strain (BY4743) on identical low-nitrogen growth media (Fig 2A). The InsP traces exhibited a number of similarities, but in the non-filamentous strain, InsP$_5$ levels did not drop as substantially, and the peaks corresponding to PP-InsP$_4$ and InsP$_8$ were elevated relative to those observed from the filamentous strain on low-nitrogen media.

With two InsP$_7$ isoforms detected in wild type under conditions of low nitrogen availability, we clarified the identity of each isoform peak by profiling InsP levels in homozygous diploid mutants deleted singly for the inositol pyrophosphate kinases VIP1 and KCS1 (Fig 2B). Deletion of VIP1 under low-nitrogen conditions resulted in a strikingly large increase in the second InsP$_7$ isoform peak, presumably corresponding to Kcs1p-produced 5PP-InsP$_5$. Deletion of KCS1 resulted in decreased InsP$_7$ isoforms. The peak corresponding to the second InsP$_7$ isoform that was elevated in vip1$\Delta$/Δ was lost in traces of kcs1$\Delta$/Δ; the peak corresponding to the first InsP$_7$ isoform was decreased but still evident in the trace from kcs1$\Delta$/Δ, and hence is most likely Vip1p-produced 1PP-InsP$_5$. Analysis of kcs1$\Delta$/Δ also identified elevated levels of PP-InsP$_4$ and two profile peaks that were not observed in other InsP kinase pathway mutants that we could not conclusively identify from alignment with our standards. As indicated in Fig 1, the vip1$\Delta$/Δ mutant that accumulates 5PP-InsP$_5$ exhibited exaggerated pseudohyphal filamentation, while the kcs1$\Delta$/Δ mutant that lacks an elevated ratio of 5PP-InsP$_5$ to 1PP-InsP$_5$ was hypofilamentous. Collectively, these data are consistent with the second InsP$_7$ isoform peak corresponding to 5PP-InsP$_5$ and suggest that the relative levels of pyrophosphorylated InsP$_7$ isoforms correlate with pseudohyphal growth phenotypes.

The kinase domain of Vip1p suppresses pseudohyphal growth

InsP levels are established through the actions of both kinases and phosphatases; phosphatases in the InsP biosynthesis pathway (Siw14p, Ddp1p, and Vip1p) are indicated in Fig 3A. Vip1p exhibits both InsP kinase and phosphatase activity. Vip1p contains an amino-terminal RimK/ATP-grasp domain responsible for phosphorylating the 1 position of InsP$_6$ and a C-terminal phosphatase-like domain that acts to dephosphorylate molecules produced by Vip1p itself [31]. To determine if the elevated filamentation in vip1$\Delta$/Δ resulted from loss of its kinase activity or phosphatase activity, we constructed chromosomal point mutations encoding kinase-defective (vip1$^D487A$) or phosphatase-defective (vip1$^H548A$) forms of Vip1p, in which the indicated conserved catalytically important residue was mutated to alanine. Haploid vip1$^D487A$ and diploid vip1$^D487A/D487A$ strains showed increased invasive growth and surface-spread filamentation, respectively (Fig 3B and 3C). Cell morphology images are presented in S1 Fig. The vip1$^D487A$ strains exhibited wild-type filamentous growth. InsP profiles of vip1$^D487A/D487A$ indicated a marked increase in 5PP-InsP$_5$ levels relative to 1PP-InsP$_5$ (Fig 3D). In contrast, the vip1$^H548A/H548A$ mutant exhibited a 5PP-InsP$_5$:1PP-InsP$_5$ ratio of less than 1. Neither point mutant replicated the large increase in InsP$_3$ levels detected in vip1$\Delta$/Δ. From these results, we conclude that the kinase domain of Vip1p, but not its phosphatase-like domain, inhibits pseudohyphal growth, and that mutations impairing Vip1p
Fig 2. Analysis of InsP levels in yeast pseudohyphal growth. A representative profile is shown from at least two independent biological replicates for each strain and growth condition tested. (A) InsP profiles are altered under conditions that induce pseudohyphal filamentation (growth in low-nitrogen SLAD media for eight hours), with distinguishable InsP₆ isoforms. InsP profiles are shown for a diploid wild-type filamentous strain grown under standard conditions (Σ1278b), a wild-type filamentous strain grown in low-nitrogen media for eight hours (Σ1278b SLAD), and a non-filamentous wild-type strain grown in low-nitrogen media for eight hours (BY4743 SLAD). (B) Representative InsP profiles of homozygous diploid vip1Δ/Δ and kcs1Δ/Δ mutants grown in low-nitrogen media compared to wild type. Profile regions corresponding to InsP₅ and InsP₆ are enlarged in the inset box. Each trace indicates counts per min (CPM) as a percentage of total CPM, with background subtracted. Elution fractions are indicated on the X-axis. Values used to generate the InsP traces are provided in S1 Dataset. The mean ratio of 5PP-IP₅ to 1PP-IP₅ is listed with standard deviation for each strain. Individual ratios of InsP₆ isoforms for each tested replicate are provided in S7 Table.

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Fig 3. Mutation of the Vip1p kinase domain results in exaggerated pseudohyphal growth. (A) Diagram indicating phosphatases affecting inositol pyrophosphates. Vip1p phosphatase activity acts redundantly with Ddp1p. (B) Haploid invasive growth phenotypes are shown for strains containing a kinase-defective vip1 allele (D487A) and phosphatase-defective vip1 allele (H548A), respectively. For comparison, haploid wild type and vip1Δ mutants are included. Assays were performed and quantified as described previously. Mean values with standard deviation for three replicates are
kinase activity, but not its phosphatase activity, result in an elevated ratio of 5PP-InsP₅ to 1PP-InsP₅.

To consider the molecular basis of the pseudohyphal growth phenotypes in the vip1 mutants, we assessed mRNA levels of the flocculence gene FLO11. The FLO11 gene encodes a GPI-anchored cell surface flocculin, often used as a molecular marker of pseudohyphal growth. Deletion of FLO11 yields a pseudohyphal growth defect under conditions of low nitrogen, and its complex promoter has been long studied as an important regulatory point in the pseudohyphal response [6]. Transcription of FLO11 is controlled through MAPK signaling (positively through Kss1p and negatively through Fus3p), the PKA pathway, and the Snf1p signaling system. We observed a strong increase in FLO11 mRNA levels in the hyperfilamentous vip1-D487A mutant relative to wild-type under conditions of low nitrogen, but less marked changes in FLO11 mRNA levels in the phosphatase-defective vip1-H548A/H548A strain (S2 Table).

Loss of the InsP phosphatase Siw14p results in elevated 5PP-InsP₅ levels and hyper-filamentous growth

Phenotypic analysis of the homozygous diploid siw14Δ/Δ deletion mutant indicated exaggerated surface-spread filamentation relative to wild type. The ddp1Δ/Δ mutant was wild-type with respect to pseudohyphal growth (Fig 4A). FLO11 mRNA levels were elevated in the hyperfilamentous siw14Δ/Δ mutant relative to wild type under conditions of low nitrogen (S2 Table). InsP profiling of both the siw14Δ/Δ and ddp1Δ/Δ strains under conditions of nitrogen limitation identified elevated levels of 5PP-InsP₅ relative to 1PP-InsP₅ in siw14Δ/Δ and increased levels of 1PP-InsP₅:5PP-InsP₅ in ddp1Δ/Δ (Fig 4B). In both mutants, the reservoir of InsP₆ was depressed, accounting partially for increases in the respective InsP₇ isoforms. The scale in Fig 4B has been adjusted to indicate the change in InsP₆ levels. These results highlight the strong correlation between elevated 5PP-InsP₅:1PP-InsP₅ ratios and hyperactive pseudohyphal growth.

To further consider the effect of perturbing levels of the 1PP-InsP₅ isoform of InsP₇, we generated a homozygous diploid strain of yeast containing a phosphatase-defective allele of VIP1 and a deletion of DDP1. As Ddp1p and Vip1p are both capable of removing the β-phosphate group from position one of PP-InsPs, we expected this mutant to produce highly elevated levels of 1PP-InsP₅. InsP profiling of the vip1-H548A/H548A ddp1Δ/Δ double mutant under conditions of nitrogen limitation identified a trace very similar to that observed in ddp1Δ/Δ under identical conditions. The double mutant exhibited wild-type levels of pseudohyphal growth, also similar to the phenotype of ddp1Δ/Δ (Fig 4A and S1 Fig).

Inositol pyrophosphate kinase overexpression driving elevated 5PP-InsP₅ levels results in elevated pseudohyphal growth

Our results thus far indicate that InsP signaling is required for pseudohyphal growth and that InsP profiles, particularly with respect to the relative levels of 5PP-InsP₅ and other inositol pyrophosphates, change in correlation with the degree of pseudohyphal growth. To establish these points more strongly, we sought to further perturb InsP signaling towards the

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production of specific InsP species for analysis of pseudohyphal growth. Because of the charged phosphate groups decorating the myo-inositol backbone, InsPs cannot be exogenously added for efficient uptake in yeast. Consequently, we utilized gene overexpression and deletion to impact InsP signaling.

Fig 4. Deletion of the SIW14 phosphatase gene results in exaggerated pseudohyphal growth. (A) Surface pseudohyphal filamentation phenotypes of homozygous diploid strains with the indicated InsP phosphatase mutations. Strains were grown on low-nitrogen SLAD medium. Exaggerated pseudohyphal growth in the siw14Δ/Δ mutant is indicated with a “++”; wild-type pseudohypha growth levels are indicated with a “+”. Pseudohyphal growth assays were performed and quantified as described; mean values with standard deviation for three replicates are indicated. Scale bar, 500 μm. (B) Representative InsP profiles of mutated InsP phosphatase strains grown in low-nitrogen minimal media for eight hours. The proportional CPM counts are shown to 1% in order to indicate the changes in IP6 levels. The InsP trace for a wild-type control strain is shown below the plot. Ratios of 5PP-InsP5:1PP-InsP5 are indicated as mean with standard deviation. Ratios of InsP7 in individual replicates are listed in S7 Table.

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To overexpress KCS1 and VIP1, we cloned each gene into a high copy vector such that the coding sequence was expressed under transcriptional control of the ADH2 promoter. Relative to wild type, KCS1 overexpression under conditions of nitrogen limitation resulted in elevated levels of InsP₃, 5PP-InsP₅, and InsP₈, with decreased levels of 1PP-InsP₅ (Fig 5A). KCS1 overexpression mutants exhibited a ratio of 5PP-InsP₃ to 1PP-InsP₃ of over 2, while VIP1 overexpression resulted in a ratio of the 5PP-InsP₃:1PP-InsP₅ isoforms of roughly 0.5. Analysis of pseudohyphal growth phenotypes in the mutants indicated elevated surface-spread filamentation in the KCS1 overexpression mutant under conditions of nitrogen limitation, while the VIP1 overexpression strain exhibited decreased filamentation relative to wild type (Fig 5A and S2 Fig). Deletion of SIW14, IPK1, and VIP1 in a strain carrying the pSGP47-KCS1 plasmid overexpressing KCS1 resulted in hyperfilamentous growth, resembling the background KCS1 overexpression mutant (Fig 5B and S2 Fig). Deletion of KCS1 in a strain overexpressing VIP1 did not change the hypofilamentous phenotype of the parent VIP1 overexpression mutant (S2 Fig). Deletion of SIW14 and IPK1, however, did alter that phenotype, resulting in exaggerated pseudohyphal filamentation relative to wild type (Fig 5B and S2 Fig). In contrast to the background VIP1 overexpression strain carrying the pSGP47-VIP1 construct, InsP profiling of siw14Δ/Δ with pSGP47-VIP1 grown in low-nitrogen media indicated strongly elevated levels of 5PP-InsP₅ relative to 1PP-InsP₅ (Fig 5B). An elevated ratio of 5PP-InsP₅:1PP-InsP₅ was also evident in the siw14Δ/Δ homozygous diploid strain, consistent with the hyper-filamentous phenotype of both mutants. FLO11 mRNA levels were elevated in the hyperfilamentous siw14Δ/Δ strains, with particularly high levels evident in siw14Δ/Δ backgrounds carrying either pSGP47-KCS1 or pSGP47-VIP1 (S2 Table).

Perturbation of InsP signaling bypasses nitrogen limitation to allow pseudohyphal growth

To assess the degree to which InsP signaling can impact pseudohyphal growth independently of other stimuli, we analyzed a set of strains with mutations affecting InsP kinases and/or phosphatases on media with normal levels of ammonium sulfate. In the presence of media with nitrogen levels that are repressive for filamentation, overexpression of KCS1 and deletion of SIW14, individually and in combination, resulted in surface-spread pseudohyphal growth (Fig 6A). InsP profiling of a KCS1 overexpression strain grown in media with normal nitrogen levels indicates two peaks corresponding to the 5PP-InsP₅ and 1PP-InsP₅ isoforms of InsP₇ (Fig 6B). It is notable that the two InsP₇ isoforms are present in this KCS1 overexpression mutant even under conditions with normal nitrogen levels, in contrast to the corresponding wild-type strain under identical conditions.

Pseudohyphal growth kinases are required for wild-type InsP levels

Since kinases and phosphatases in the InsP biosynthesis pathway are essential for wild-type pseudohyphal growth, we investigated the possibility that signaling pathways required for pseudohyphal growth may regulate levels of InsP species. For this study, we focused on the Snf1p kinase pathway and Kss1p and Fus3p MAPKs, as we previously identified InsP kinases as being differentially phosphorylated in strains with kinase-defective alleles of SNF1 and MAPKs [43]. Deletion of TPK2, encoding a catalytic subunit of protein kinase A required for pseudohyphal growth, did not substantially affect InsP levels under conditions of nitrogen limitation.

The homozygous diploid snf1Δ/Δ strain is defective in pseudohyphal growth with no filamentation evident in low-nitrogen media [11, 18, 46]. InsP profiling of snf1Δ/Δ indicated elevated levels of InsP₃, InsP₅, PP-InsP₄, the InsP₇ isoforms, and InsP₈ (Fig 7A). The highly
Fig 5. Overexpression mutants with elevated levels of 5PP-InsP\textsubscript{5} relative to other inositol pyrophosphates exhibit exaggerated pseudohyphal growth. (A) Representative InsP profiles of strains overexpressing VIP1 and KCS1 grown for eight hours in low-nitrogen minimal media. The ratio of InsP\textsubscript{7} isoforms is indicated as mean values with standard deviations shown. InsP\textsubscript{7} ratios in independent replicates are shown in S7 Table. (B) Representative InsP profile of VIP1 overexpression in a siw14\textDelta/\Delta background strain grown in low nitrogen media; a representative InsP profile for siw14\textDelta/\Delta is included also. The InsP trace of the wild-type strain for this analysis is shown in panel A; all strains were grown under identical conditions. Pseudohyphal growth phenotypes are provided for the indicated mutants. Filamentation was quantified as described in Materials and Methods; mean values with standard deviation are indicated. Scale bars, 500 \textmu m.

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elevated levels of InsP$_3$ and the absence of a predominance of the 5PP-InsP$_5$ isoform relative to 1PP-InsP$_5$ and InsP$_8$ are particularly notable. As evidenced in the snf1$\Delta/\Delta$ mutant, a homozygous kss1$\Delta/\Delta$ strain displayed elevated levels of InsP$_3$. The kss1$\Delta/\Delta$ mutant is hypofilamentous and showed roughly comparable levels of the 1PP-InsP$_5$ and 5PP-InsP$_5$ isoforms (Fig 7B), with elevated levels of InsP$_8$. The mating pathway MAPK Fus3 negatively regulates pseudohyphal growth by phosphorylating and targeting the transcription factor Tec1p for destruction; its deletion results in exaggerated invasive and surface-spread filamentation [18, 47]. Consistent with the pattern observed in other mutants with exaggerated pseudohyphal growth, the hyperfilamentous fus3$\Delta/\Delta$ mutant exhibited an elevated ratio of 5PP-InsP$_5$ to 1PP-InsP$_5$ (Fig 7B).

InsP$_8$ levels were also increased in fus3$\Delta/\Delta$.

To determine if InsP signaling in turn impacts MAPK pathway activity, we assessed the phosphorylation state of Kss1p in strains singly deleted for the InsP pathway genes PLC1, IPK1, VIP1, and KCS1 (S3 Fig). Kss1p phosphorylation was not strikingly affected in any of these mutants, suggesting that the MAPK pathway may not be acted upon by InsP signaling; the possibility still exists, however, that MAPK signaling may regulate InsP-based signaling.

### Discussion

InsP signaling has been studied for over two decades, but its functional significance has not been fully elucidated. Owing partially to the labor involved, InsPs have not been profiled over a broadly representative spectrum of growth conditions. Here, we used the genetic workhorse *Saccharomyces cerevisiae* to profile InsP levels under conditions of nitrogen limitation in a filamentous background, and our results indicate a new role for InsP signaling in the pseudohyphal growth response. Perturbation of the InsP biosynthesis pathway altered pseudohyphal growth, and in vivo measurement of InsP levels in these mutants identified characteristic profiles, particularly with respect to the inositol pyrophosphates, that were predictive of...
pseudohyphal growth states. Mutation of genes encoding InsP kinases and phosphatases enabled pseudohyphal growth in the absence of nitrogen limitation. InsP profiles were substantially altered in mutants deleted of SNF1, KSS1, and FUS3, suggesting that these well studied kinase regulators of pseudohyphal growth contribute to the establishment and/or maintenance of wild-type InsP levels under conditions of nitrogen limitation.

Considered collectively, the data suggest the ratio of the InsP₇ isoforms is indicative of the pseudohyphal growth state. Elevated levels of 5PP-InsP₅ relative to the levels of 1PP-InsP₅ were evident in strains that showed exaggerated pseudohyphal growth. This pattern held consistent under conditions of nitrogen limitation in homozygous diploid vip1Δ/Δ, vip1-D487A/ D487A, siw14Δ/Δ, KCS1 overexpression, and fus3Δ/Δ strains. Further, in low-nitrogen media, overexpression of VIP1 in a strain deleted of SIW14 resulted in elevated levels of 5PP-InsP₅ and hyperfilamentation. Collectively, the ratios of InsP₇ isoforms in these hyperfilamentous

Fig 7. The pseudohyphal growth regulatory kinases Snf1p, Kss1p, and Fus3p are required for wild-type InsP signaling. (A) Representative InsP profiles of the wild-type filamentous Σ1278b strain and an otherwise isogenic snf1Δ/Δ mutant grown in low-nitrogen media. (B) Representative InsP profiles are indicated for kss1Δ/Δ and fus3Δ/Δ mutants grown in low-nitrogen media; an InsP profile of the wild-type Σ1278b strain under identical growth conditions is provided for comparison. Mean ratios of the InsP₇ isoforms are shown with standard deviation. InsP₇ ratios of independent replicates are provided in S7 Table.

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mutants is distinct from corresponding ratios in wild-type strains by an independent samples T-test (p<0.001) (S4A Fig). This set of hyperfilamentous mutants exhibit 5PP-InsP₅ to 1PP-InsP₃ ratios that are also distinct from those observed in hypofilamentous mutants (kcs1Δ/Δ, VIP1 overexpression, snf1Δ/Δ, and kss1Δ/Δ mutants) (p<0.001) (S4B Fig). It is interesting that wild-type pseudohyphal growth did not indicate highly elevated levels of 5PP-InsP₅ relative to 1PP-InsP₅ and InsP₆, but instead was characterized by decreased InsP₅ and elevated InsP₇ isoforms, with 1PP-InsP₅ showing the greater increase. It should also be noted that exaggerated levels of PP-InsP₄ and 1,5(PP)₂-InsP₃ in ipk1Δ/Δ promote pseudohyphal growth, although the effects may be observed only upon removal of the higher energy IP₇ and IP₈ pyrophosphates, as we did not observe consistent changes in PP-InsP₄ and 1,5(PP)₂-InsP₃ levels over the mutants we analyzed. Additional InsP profiling in a larger mutant set will likely be necessary to refine the characteristic InsP signatures, but the data are consistent with the notion that InsP profiles are a predictive indicator of mutant pseudohyphal growth states.

These results highlight the importance of inositol pyrophosphates, and particularly InsP₇ isoforms, in the pseudohyphal growth response. It is notable that two peaks corresponding to the InsP₇ isoforms 5PP-InsP₅ and 1PP-InsP₅ are present in profiles of strains grown in low-nitrogen media. Previous analyses of InsP profiles in yeast by HPLC-based fractionation have presented a single peak corresponding to InsP₇; however, these studies were not undertaken using nitrogen-limiting growth media, nor using the 15-minute extended separation gradient we employed here. In fact, our InsP profiles of the wild-type Σ1278b strain under normal growth conditions do indicate a single InsP₇ peak. This single peak had previously been identified from analyses of non-filamentous yeast as 5PP-InsP₅ [26, 33, 45]; however, our fractionation profiles indicate that this peak corresponds to 1PP-InsP₅. Several lines of evidence support this conclusion. First, InsP profiles of kcs1Δ/Δ grown on low-nitrogen media identify a small peak aligning with the one observed in profiles of a wild-type strain under normal growth conditions. This peak most likely corresponds to Vip1p-produced 1PP-InsP₅, as Vip1p is capable of pyrophosphorylating InsP₆ in a strain deleted of KCS1. Second, profiles of vip1Δ/Δ indicate a very large peak aligning with the later InsP₇ isoform fraction, which, by similar logic, likely represents Kcs1p-produced 5PP-InsP₅. Further, an identical peak is observed in profiles of siw14Δ/Δ in which 5PP-InsP₅ accumulates, as this isomer cannot be dephosphorylated in the absence of SIW14 [32]. Thus, the data indicate that we can efficiently separate InsP₇ isomers in yeast under low nitrogen conditions through an extended gradient, with 1PP-InsP₅ fractionating just ahead of 5PP-InsP₅.

Beyond its relevance to yeast pseudohyphal growth, this study is also informative in considering the role of InsPs in the cellular response to nitrogen limitation. Filamentous and non-filamentous strains grown with reduced ammonium sulfate exhibit many similar changes in InsP levels, including elevated levels of pyrophosphorylated InsP₇ isoforms as discussed above. These data suggest that the transition to conditions of decreased nitrogen availability necessitates the increased accumulation of both InsP₇ isoforms, making them detectable by labeling and fractionation protocols. It is additionally interesting that overexpression of KCS1 in normal media results in InsP₇ fractionation peaks corresponding to 5PP-InsP₅ and 1PP-InsP₅, mimicking the profile observed in strains grown under conditions of low nitrogen with associated filamentation. Thus, InsP levels can be manipulated to uncouple pseudohyphal growth from nitrogen limitation, and, further, inositol pyrophosphate abundance may be an intracellular signal of nitrogen conditions, functioning as part of the cellular response in yeast to manage nitrogen limitation.

InsP profiles in this study as well as previous phosphoproteomic data are consistent with a role for AMPK and MAPK signaling in regulating InsP levels under conditions of nitrogen limitation, and the results here provide the first evidence that AMPK and MAPK signaling is
required for wild-type levels of InsPs. Interestingly, inositol polyphosphate multikinase has previously been demonstrated to be capable of binding AMPK [48]. These signaling modules are established regulators of cellular responses to nutrient levels, and functions for Snf1p, Kss1p, and Fus3p in controlling InsP levels would provide an important point of crosstalk between classic cell signaling pathways and metabolic second messengers. In sum, this work further establishes the foundation for additional studies of the regulation and downstream effectors of InsP signaling in cellular stress responses and pseudohyphal growth.

Materials and methods

Strains, plasmids, and media

Strains used in this study are listed in S3 Table. Filamentous yeast strains were derived from the genetic background Σ1278b [5, 49]. Haploid strains were derived from HLY337 and Y825 [50, 51]. Non-filamentous strains were derived from BY4743 [52]. Standard protocols for proper growth and maintenance of yeast have been described previously [53, 54]. DNA was introduced into yeast by standard protocols involving lithium acetate treatment and heat shock [55].

Plasmids for ADH2-based overexpression [56] were derived from pSGP47. The pSGP47 vector contains the S. cerevisiae 2μ origin of replication, resulting in roughly 50 copies per cell. Transcript levels of KCS1 and VIP1 from these overexpression vectors were assessed under normal and low-nitrogen conditions, and the results are presented in S4 Table. The plasmids enable strong overexpression of the KCS1 and VIP1 genes, particularly under conditions of low nitrogen.

S. cerevisiae strains were cultured on YPD (1% yeast extract, 2% peptone, 2% glucose) or Synthetic Complete (SC) media (0.67% Yeast Nitrogen Base without amino acids, 2% glucose, and 0.2% of the appropriate amino acid drop-out mix). Nitrogen limitation conditions for pseudohyphal growth phenotypic assays were achieved using Synthetic Low Ammonium Dextrose (SLAD) media (0.17% Yeast Nitrogen Base without amino acids and without ammonium sulfate, 2% glucose, 50 μM ammonium sulfate, and appropriate amino acids as necessary). For inositol polyphosphate profiles, yeast cultures were grown in minimal media lacking inositol (0.17% Yeast Nitrogen Base without ammonium sulfate, amino acids, and inositol, 2% glucose, 0.5% ammonium sulfate, and appropriate amino acids) and as needed were subsequently grown in SLAD-inositol media (0.17% Yeast Nitrogen Base without amino acids, ammonium sulfate, and inositol, 2% glucose, 50 μM ammonium sulfate, and any required amino acids).

Gene deletions and integrated point mutations

Gene deletions were generated by one-step PCR-based replacement using either KanMX6 or HphMX6 drug-resistance cassettes of pFA6a-KanMX6 or pAG32 [57]. Gene replacement was verified by PCR-based methods as described [58]. Diploid mutants were typically constructed by deleting each desired gene in haploid backgrounds of opposite mating type and subsequently mating the strains, selecting for complementary auxotrophies. The kss1Δ/Δ and fus3Δ/Δ mutants were generated in a diploid cell where each gene copy was deleted by allelic replacement with the indicated drug resistance markers [57]. Overexpression plasmids were constructed by ligation-independent cloning using the parent pSGP47 vector [59, 60]. Integrated point mutations were generated using the URA3-flip-out method as previously described [61, 62].
Filamentous growth assays

For invasive growth assays, cultures of haploid strains were grown overnight with shaking in YPD media at 30˚C, harvested, and adjusted to an optical density of 1 at 600 nm in sterile water. A 5 μl aliquot of the culture was spotted onto YPD plates and allowed to invade the agar for two days. Plates were photographed for surface growth and then gently washed with a light stream of water and photographed again to visualize invasive growth [49, 58, 63]. The degree of invasive growth was quantified as the mean pixel intensity of the washed spot relative to its pre-washed image using the program ImageJ, as described previously [20]. Triplicate replicates were assayed for each haploid strain tested, and results for each independent replicate are provided in S5 Table. Results were binned as follows: hypofilamentous, less than 0.5; wild-type, between 0.5 and 0.7; hyperfilamentous, greater than 0.7.

For surface spread assays, diploid strains were grown overnight in appropriate SC-drop-out media. Saturated cultures were spread onto SLAD medium and incubated at 30˚C for four days [5, 50]. Colonies were imaged using an upright Nikon Eclipse 80i microscope with CoolSnap ES2 CCD (Photometrics). Images were acquired using MetaMorph software (Molecular Devices). The extent of surface-spread pseudohyphal growth was quantified using the approach of Ryan et al. [19] with minor modifications. In brief, the circumference of a defined area of a colony is measured using the NeuronJ program, and the value is compared against the corresponding circumference of the same defined area of a wild-type strain colony under conditions of low nitrogen. The ratio is presented for three replicates (S6 Table), and the average and standard deviation is indicated. Mean ratios of greater than 1.5 are indicated as hyperfilamentous (++), and mean ratios of less than 0.75 are indicated as hypofilamentous (-).

Expression analysis of FLO11, KCS1, and VIP1

Single colonies were inoculated in 4 ml appropriate medium (YPD or SC-Ura). Overnight cultures were diluted 1:1000 in 5 ml YPD or SC-URA and let grow for 16–20 hours. The cells were harvested at 3000g for 5 minutes and washed twice with water. If no nitrogen stress was utilized, the cells were either stored at -80˚C or subjected to RNA extraction right away. If nitrogen stress was utilized, the cells were resuspended in 5 ml SLAD medium and incubated at 30˚C for 8 hours. After incubation, the cells were harvested and washed twice with water. The cells were either stored at -80˚C or subjected to RNA extraction right away. RNA extraction was done using the RiboPure-Yeast kit (Invitrogen) according to manufacturer’s directions. The amount of RNA isolated was determined using a NanoDrop spectrophotometer/fluorometer. For each sample, 1 μg of RNA was converted to cDNA using the Radiant cDNA Synthesis Kit, 1-Step. The resulting cDNA was diluted 1:100 and 2 μl of the diluted cDNA was used as a template for qPCR. qPCR mixes were prepared using the Radiant Green Hi-ROX qPCR kit and run with an Applied Biosystems StepOne Plus qPCR machine. The relative amounts of FLO11, KCS1, and VIP1 expression were calculated using the double delta Ct method with ACT1 as the reference gene.

Primers used for qRT-PCR are as follows: ACT1-qPCR-F: CTGCCGGTATTGACCAAAC T; ACT1-qPCR-R: CGGTGATTTCCTTTTGCATT; FLO11-qPCR-FWD3: GTTGTGTTGCCC AGCGGAGGT; FLO11-qPCR-RVS3: CTACCCACCCCTGTCGGACG; KCS1-qPCR-FWD3: GCATAAATTGGCGGCTCCGGTG; KCS1-qPCR-RVS3: TGGCAGCATGTGGTATTTG; VIP1-qPCR-FWD1: AGAGCTCTTGGGCGGGA; VIP1-qPCR-RVS1: GTGCGGCAGCT TCCCTACCC.
Inositol polyphosphate profiling

HPLC analysis of InsP levels was conducted as previously described [45] with the following modifications. In brief, yeast cultures were grown overnight at 30°C with shaking until fully saturated in rich media. Subsequently, 5 µl of the saturated culture was added to 5 ml of minimal media lacking inositol supplemented with 25 µl of myo[1,2-3H]inositol 1 mCi/ml 30 Ci/mmol (American Radiolabeled Chemicals cat. no. ART02611MC) and allowed to grow overnight at 30°C with shaking at 250 rpm until reaching an OD$_{600}$ of 0.9. Cells were then either harvested and frozen at -80°C until further use or harvested and washed with water prior to being suspended in 5 ml of SLAD-inositol media; this culture was grown for 8 hours at 30°C at 250 rpm. These cultures were harvested and frozen at -80°C until further use. Inositol polyphosphates were extracted by suspending the thawed pellet in 300 µl of 1M perchloric acid with 3mM EDTA and bead beating at 4°C for five minutes. Samples were spun down, and the resulting supernatant was saved and neutralized with the addition of 1 M potassium carbonate and 3 mM EDTA until a pH of 6.0–8.0 was reached. The mixture was allowed to sit on ice for two hours. The sample was again centrifuged, and the clear supernatant was analyzed by HPLC (Hewlett Packard Series 1100) connected to a Partisphere 5 µm SAX cartridge column 125 x 4.6 mm (HiCHROM cat. no. 4621–0505). Inositol polyphosphates were eluted from the column with a gradient from mixing buffer A (1 mM EDTA) and buffer B (1.3 M (NH$_4$)$_2$HPO$_4$, 1 mM EDTA, pH to 3.8 with H$_3$PO$_4$). The gradient was as follows: 0–5 min, 0% buffer B; 5–10 min, 0–10% buffer B; 10–90 min, 10–100% Buffer B; 90–100 min, 100% Buffer B; 100–101 min, 0% buffer B; 101–110 min 0% buffer B. The gradient was run at a flow rate of 1 ml/min and one-1ml fractions were collected every minute for the first 90 minutes. Fractions were mixed with 4 ml of Ultima-Flo AP liquid scintillation cocktail (Perkin-Elmer cat. no. 6013599) and counted using a scintillation counter.

Supporting information

S1 Fig. Cell morphology of IP kinase and phosphatase mutants under conditions that induce pseudohyphal growth. (A) The indicated strains deleted for genes encoding IP kinases were grown in low-nitrogen media along with the wild-type Σ1278b parent strain. Cells were scraped from a colony after growth in low-nitrogen SLAD medium, and the cells were suspended in solution prior to DIC imaging. Cells with length-to-width ratios greater than 2.0 are highlighted with asterisks. Scale bar, 5 µm. (B) Cell morphology of indicated strains mutated for genes encoding IP phosphatases. Cells were grown and imaged as above. Scale bar, 5 µm. (TIF)

S2 Fig. Filamentous growth phenotypes of overexpression mutants perturbing IP biosynthesis. (A) Cells of the indicated strains were scraped from colonies grown on low-nitrogen SLAD medium and imaged by DIC. Cells with length-to-width ratios greater than 2.0 are highlighted with asterisks. Scale bar, 5 µm. (B) Surface-spread filamentation phenotypes of indicated mutant strains. “++” represents hyperfilamentous growth, and “..” indicates an absence of pseudohyphal filamentation. Scale bar, 500 µm. (TIF)

S3 Fig. Kss1p phosphorylation in InsP pathway mutants. Cells were grown overnight in YPD and 30°C. Cells were sub-cultured into either YPD or YP-Gal medium and grown for 6 hours. Protein extracts were made by TCA precipitation and separated by 10% SDS-PAGE. Proteins were transferred to nitrocellulose membrane and blotted with either p44/42 antibody (Cell Signaling Technology, Danvers, MA; #4370) or Kss1 antibody (Santa Cruz Biotechnology, Santa Cruz, CA; #6775) or Pgk1 antibody (Life Technologies; Camarillo, CA; #459250) as
indicated. For secondary antibodies (goat anti-mouse IgG–HRP, Bio-Rad Laboratories, Hercules, CA; #170–6516; goat anti-rabbit IgG–HRP, Jackson ImmunoResearch Laboratories, Inc., West Grove, PA; #111-035-144) were used. Membranes were developed using Western Bright kit from Advansta Inc. (Menlo Park, CA; LPS #K-12045-D20) and imaged using Image-lab software (Biorad Inc.). The ste11Δ mutant is a control for diminished Kss1p phosphorylation. Levels of phosphorylated Kss1p relative to wild-type were estimated by densitometry with normalization to the Pgk1p loading control.

S4 Fig. Hyperfilamentous mutants exhibit 5PP-InsP₅:1PP-InsP₅ ratios distinct from those observed in wild-type and hypofilamentous strains. (A) By independent samples T-test, the difference in ratios of the InsP₅ isoforms between hyperfilamentous mutants analyzed in this study (vip1Δ/Δ, vip1-D487A/D487A, siw14Δ/Δ, KCS1 overexpression, siw14Δ/Δ VIP1 overexpression double mutants, and fus3Δ/Δ strains) and wild-type control strains is statistically significant (p<0.001). (B) Hyperfilamentous mutants indicated above also exhibit a statistically significant difference (p<0.001) in 5PP-InsP₅:1PP-InsP₅ ratios from hypofilamentous mutants (kcs1Δ/Δ, VIP1 overexpression, snf1Δ/Δ, and kss1Δ/Δ mutants) analyzed in this study.

S1 Table. The pseudohyphal growth kinases Snf1p and Kss1p are required for wild-type phosphorylation of IP kinases.

S2 Table. FLO11 mRNA levels in InsP phosphatase mutants.

S3 Table. List of strains used in this study.

S4 Table. mRNA levels of over-expressed KCS1 and VIP1 in high-copy vectors with the ADH2 promoter.

S5 Table. Invasive growth assay datasets.

S6 Table. Pseudohyphal filament circumference of independent replicate colonies.

S7 Table. Ratio of InsP₇ isoforms in independent replicates for indicated strains.

S1 Dataset. Percent CPM values for separated fractions plotted in InsP traces.

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