MCK1 is a novel regulator of myo-inositol phosphate synthase (MIPS) that is required for inhibition of inositol synthesis by the mood stabilizer valproate

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Abstract

Myo-inositol, the precursor of all inositol compounds, is essential for the viability of eukaryotes. Identifying the factors that regulate inositol homeostasis is of obvious importance to understanding cell function and the pathologies underlying neurological and metabolic resulting from perturbation of inositol metabolism. The current study identifies Mck1, a GSK3 homolog, as a novel positive regulator of inositol de novo synthesis in yeast. Mck1 was required for normal activity of myo-inositol phosphate synthase (MIPS), which catalyzes the rate-limiting step of inositol synthesis. mck1Δ cells exhibited a 50% decrease in MIPS activity and a decreased rate of incorporation of [¹³C₆]glucose into [¹³C₆]-inositol-3-phosphate and [¹³C₆]-inositol compared to WT cells. mck1Δ cells also exhibited decreased growth in the presence of the inositol depleting drug valproate (VPA), which was rescued by supplementation of inositol. However, in contrast to wild type cells, which exhibited more than a 40% decrease in MIPS activity in the presence of VPA, the drug did not significantly decrease MIPS activity in mck1Δ cells. These findings indicate that VPA-induced MIPS inhibition is Mck1-dependent, and suggest a model that unifies two current hypotheses of the mechanism of action of VPA—inositol depletion and GSK3 inhibition.

Introduction

Myo-inositol is the precursor of all inositol compounds, including phosphoinositides, inositol phosphates, inositol sphingolipids, and glycosylphosphatidylinositol. Inositol compounds are critical for nearly all cellular processes [1, 2] and are essential for the viability of eukaryotes. The pivotal role of inositol is underscored by the link between perturbation of inositol metabolism and human neurological disorders [3, 4]. Lithium, a widely used mood-stabilizing drug, was found to induce inositol depletion by inhibiting inositol monophosphatase activity [5].
Inositol depletion was thus proposed as a possible therapeutic mechanism of action of the drug [6]. Consistent with the inositol depletion hypothesis, the mood-stabilizing drug valproic acid (VPA), a short chain branched carboxylic acid with no structural similarity to lithium, was also reported to deplete intracellular inositol [7, 8]. Elucidating how inositol metabolism is regulated and how mood-stabilizing drugs deplete inositol will greatly benefit the understanding of both essential cellular functions and the pathologies underlying neurological and other illnesses associated with perturbation of inositol metabolism.

In the yeast *Saccharomyces cerevisiae*, inositol is synthesized *de novo* from glucose 6-phosphate (G-6-P) in a two-step reaction that has been well characterized. The rate-limiting enzyme myo-inositol-3-phosphate synthase (MIPS), which is encoded by *INO1*, converts G-6-P to inositol-3-phosphate (I-3-P) [9, 10]. In the second step, inositol-3-phosphate is dephosphorylated to myo-inositol by inositol monophosphatase [11, 12]. *INO1* is the most tightly controlled phospholipid biosynthesis gene. The transcription of *INO1* is regulated by the transcriptional repressor Opi1 in response to extracellular inositol levels [13]. When inositol is available in the growth medium, Opi1 translocates to the nucleus, where it represses *INO1* transcription and inhibits inositol synthesis. When inositol is limiting, Opi1 is excluded from the nucleus, and *INO1* transcription is derepressed by Ino2 and Ino4 for the *de novo* synthesis of inositol. In addition to the Ino2-Ino4-Opi1 regulatory circuit, *INO1* transcription in yeast and mammalian cells is regulated by inositol pyrophosphate synthase. Interestingly, yeast *INO1* is positively regulated by inositol pyrophosphate synthase, while mammalian cells are negatively regulated by this enzyme [14, 15]. In addition to transcriptional control of *INO1*, inositol synthesis is regulated by posttranslational modification of MIPS. Both yeast and human MIPS contain conserved phosphorylation sites, which regulate the enzymatic activity of MIPS [16, 17]. Inositol synthesis is thus highly regulated in both yeast and mammalian cells by multiple mechanisms at the level of *INO1*.

In light of the key role of MIPS in the regulation of inositol synthesis, elucidating the mechanism whereby MIPS is affected by the inositol-depleting drug VPA is of obvious importance. Yeast cells treated with therapeutically relevant concentrations of VPA exhibited decreased levels of inositol-3-phosphate and inositol, consistent with decreased MIPS activity [7]. Both yeast and human MIPS purified from yeast cells grown in the presence of VPA exhibited decreased enzymatic activity, consistent with covalent modification of the enzyme [17, 18]. However, *in vitro* activity of purified yeast or mammalian MIPS was not affected by VPA, indicating that inhibition is indirect [8, 18]. Interestingly, mutation of two phosphorylation sites in MIPS decreased VPA-induced MIPS inhibition [17], suggesting that inhibition of the enzyme by VPA may be mediated by a phosphorylation cascade.

While the inositol depletion hypothesis has stimulated considerable research into the mechanisms of action of mood stabilizing drugs, inhibition of GSK3 has also been proposed as a therapeutic mechanism of action. Lithium was first reported to cause GSK3β inhibition in *Xenopus* [19]. Inhibition of GSK3 by lithium was also observed in other model systems [20–24]. Interestingly, VPA has also been shown to inhibit GSK3 in several studies [25–28], suggesting that GSK3 inhibition may account for the therapeutic effect of mood-stabilizing drugs. GSK3 is a serine/threonine kinase that exerts regulatory functions in many cellular events [29–31]. GSK3α and GSK3β, which contain highly conserved amino acid sequences in their kinase domains, are two major GSK3 isoforms expressed in mammalian cells [32, 33]. GSK3β is predominant expressed in the brain. The alteration of GSK3β-mediated signaling pathways is associated with neuronal disorders and cancer [30, 31, 34, 35]. Consistent with the GSK3 inhibition hypothesis, GSK3 inhibitors exhibited mood-stabilizing effects in animal studies. In rodents, GSK3 inhibitors AR-A014418 and L803-mts exhibited anti-depressive effects in forced swim tests [36–38]. GSK3 inhibitors also antagonized amphetamine-induced
hyperactivity, a rodent model of mania [37, 39]. Therefore, similar to inositol depletion, GSK3 inhibition is a common outcome of VPA and lithium and is associated with mood-stabilizing effects.

Although the inositol depletion and GSK3 inhibition hypotheses of mood stabilization have been suggested independently based on seemingly unrelated studies, our findings suggest that they may be linked by a common mechanism [40]. In light of published findings that VPA causes inositol depletion by indirectly inhibiting MIPS [18], that VPA inhibits GSK3 (23–26), and that yeast cells lacking all four GSK3 homologs (MCK1, MSD1, MRK1 and YGK3) exhibit inositol depletion [41], we hypothesized that VPA inhibits MIPS by a mechanism involving negative regulation of one or more GSK3 homologs, thereby causing inositol depletion. In this report, we demonstrate for the first time that the GSK-3 homolog MCK1 (but not MSD1, MRK1 or YGK3) is required for the optimal synthesis of inositol. Importantly, VPA does not inhibit MIPS in mck1Δ cells, indicating that Mck1 is a downstream target of VPA in MIPS inhibition.

**Materials and methods**

**Yeast strains**

All strains used in this study were in the W303 background. The WT strain was crossed with the isogenic gsk3Δ (mck1Δmrk1Δmds1Δygk3Δ) mutant for the generation of haploid spores, from which isogenic mck1Δ, mrk1Δ, mds1Δ and ygk3Δ strains were derived. The GSK3 genotype of each mutant strain was confirmed by PCR. For determination of MIPS enzymatic activity, yeast strains that harbor the His-Xpress tagged MIPS gene at the INO1 locus were constructed. To do so, the INO1 gene was first replaced by a KanMX cassette, which was subsequently replaced by an N-terminal His-Xpress tagged INO1 cassette cloned from the pRD-INO1 plasmid [17].

**Growth media**

Yeast cells were grown at 30°C, 37°C or 38°C in synthetic complete (SC) medium, which contained glucose (2% w/v), adenine (20 mg/liter), arginine (20 mg/liter), histidine (20 mg/liter), methionine (20 mg/liter), tryptophan (20 mg/liter), leucine (60 mg/liter), lysine (200 mg/liter), threonine (300 mg/liter), ammonium sulfate (0.2% w/v), inositol-free Difco vitamin mix, vitamin-free yeast base, plus agar (2% w/v) for solid medium. Inositol (75 μM) and VPA (1 mM) were added separately as indicated.

**Measurement of intracellular inositol levels**

Intracellular inositol levels were determined using the method of Maslanski and Busa with modification [42]. Briefly, cells were lysed in dH2O containing 1X protease inhibitor by vortexing with acid-washed glass beads at 4°C. Cell extracts were mixed with 7.5% perchloric acid and centrifuged at 10,000 g for 10 min at 4°C. Supernatants were collected and titrated with ice cold KOH to pH 7. Samples were clarified by centrifugation and loaded onto columns containing 1 ml AG 1-X8 resin/H2O (1:1) mixture. Inositol was eluted with 5 ml dH2O, and eluates were dried in an oven at 70°C and stored at -80°C. Prior to assay, samples were dissolved in dH2O. Inositol content in samples was measured as described previously [43].

**In vivo assay of the rate of inositol de novo synthesis**

Cells were grown in SC I+ medium to the mid log phase, washed twice with dH2O, transferred to SC I- medium, and incubated for 1 h or 3 h. [13C6]-glucose was added to a final concentration
of 0.2%. After 15 min, cells were harvested and lysed in dH$_2$O containing 1X protease inhibitor by vortexing with acid-washed glass beads at 4˚C. Cell extracts were mixed with acetonitrile to a final concentration of 90% and centrifuged at 16,000 g for 10 min at 4˚C to remove precipitated protein. Supernatants were collected and stored at -20˚C for the analysis of $^{13}$C labeled inositol-3-phosphate ($^{[13}$C$_6$]-I-3-P) and inositol ($^{[13}$C$_6$]-Inositol) by LC-MS. $^{[13}$C$_6$]-I-3-P and $^{[13}$C$_6$]-Inositol were quantified by LC-MS as follows: Samples were subjected to HPLC on a Diamond Hydride column (Cogen HPLC Columns, Microsoft Technology Corporation, 2 mm x 150 mm, 4μ, 100 Å) using gradient elution. The gradient between mobile phases A: acetonitrile-aqueous ammonium formate (9:1 v/v) and B: acetonitrile-0.1% aqueous formic acid (1:9 v/v) was: 0 min– 95% A, 2 min– 80% A, 4 min– 80% A, 7 min– 30% A, 9 min– 30% A. At the end of each chromatographic run, the mobile phase composition was changed to initial conditions and equilibrated for 5 min before the start of the next sample analysis. The flow rate was 0.2 ml/min and the column was maintained at 35˚C. The eluent was directly introduced to the mass analyzer (QTRAP5500, Sciex) via a Turbolon Electrospray ionization source. The following parameters were used for the mass analyzer: Ion spray voltage: -4500 V, Curtain gas: 35, Source temperature: 600˚C, GS1 & GS2: 45, and EP & CXP: 10. $^{[13}$C$_6$]-I-3-P and $^{[13}$C$_6$]-inositol were monitored by Multiple Reaction Monitoring with the following transitions, declustering potentials (DP), and collision energies (CE). $^{[13}$C$_6$]-I-3-P m/z 265.1 to 79 at DP: -110 V and CE: -25 V; $^{[13}$C$_6$]-inositol m/z 185.1 to 89 at DP: -96 V and CE: -21 V. Under these conditions, the retention times for $^{[13}$C$_6$]-I-3-P and $^{[13}$C$_6$]-inositol were 4.7 and 3.6 min, respectively. The detection limits were 0.1 ng and the quantitation limits were 0.4 ng on the column for each analyte.

MIPS activity assay

Cells expressing His-Xpress tagged MIPS were grown to the early stationary phase and lysed in buffer (50 mM Tris-HCl pH 7.5, 0.6 M sorbitol, 0.3 M NaCl, 1X protease inhibitor and 1X phosphatase inhibitor) by vortexing with acid-washed glass beads at 4˚C. MIPS protein was purified from cell extracts using the PureProteome™ Nickel Magnetic Bead System (Millipore). Purified MIPS protein was dialyzed (1 mM Tris acetate pH 8.0, 0.05 M dithiothreitol, 0.025X protease inhibitor and 0.1X phosphatase inhibitor) and concentrated with an Amicon Ultra-0.5 Centrifugal Filter (Millipore). MIPS protein concentration was determined by Bradford assay. Enzymatic activity of 3μg purified MIPS was determined by enzyme-coupled colorimetric assay [44].

Immunoblotting

Cell extract was obtained as described above and protein concentration was determined by the Bradford assay (Bio-Rad). Extract corresponding to 30–40 μg protein was analyzed by SDS-PAGE on a 10% gel. Immunoblotting was performed using primary antibodies against the Xpress tag (R901-25, mouse, Thermo Fisher), β-actin (mAbGEa, mouse, Thermo Fisher), followed by corresponding secondary antibodies.

Results

Deletion of MCK1 leads to inositol depletion

We have previously shown that yeast cells lacking all four GSK3 homologs (gsk3Δ) exhibit decreased inositol synthesis [45]. To identify the GSK3 gene required for inositol synthesis, isogenic mutants were constructed with all combinations of gsk3 mutations. As seen in Fig 1A, a nearly 50% drop in intracellular inositol levels was observed in mck1Δ cells, similar to the 60% decrease in inositol levels seen in gsk3Δ cells. The presence of only the MCK1 gene (i.e., triple mutant mrk1Δmds1Δygk3Δ) was sufficient for optimal inositol synthesis. Consistent with
decreased intracellular inositol levels, mck1Δ cells exhibited decreased growth at 36˚C, which was rescued by supplementation of inositol (Fig 1B). These findings indicate that Mck1 is a positive regulator of inositol synthesis.

**Mck1 regulates the rate of inositol de novo synthesis**

We wished to determine if decreased intracellular inositol levels in mck1Δ cells reflected a decreased rate of synthesis of inositol in vivo. To do so, cells were grown in I⁺ medium,
washed, and transferred to I− medium for 3 h. [13C6]glucose was then added, and synthesis of [13C6]-inositol-3-phosphate and [13C6]-inositol (designated as [13C6]-I-3-P and [13C6]-inositol, respectively) was detected by LC-MS. As seen in Fig 2, mck1Δ and gsk3Δ cells exhibited significantly lower levels of [13C6]-I-3-P and [13C6]-inositol compared to WT cells. These results indicate that Mck1 is required for optimal de novo synthesis of inositol.

### Mck1 regulates MIPS activity

MIPS catalyzes the rate-limiting step of inositol de novo synthesis. To ascertain if decreased inositol synthesis in mck1Δ cells resulted from decreased enzymatic activity of MIPS, isogenic strains expressing His-Xpress tagged MIPS in the genomic site were constructed (Fig 3A). Compared to WT, a 50% decrease in activity was observed in MIPS purified from mck1Δ cells (Fig 3B), consistent with the loss of a positive regulator of MIPS in the mutant. MIPS activity in the mrk1Δmds1Δygk3Δ triple mutant was similar to that of WT, in agreement with the finding of WT inositol levels in the mutant (Fig 1A). WT and mutant cells contained similar levels of MIPS protein (Fig 3C), indicating that the deletion of MCK1 does not affect MIPS synthesis. Taken together, these findings indicate that deletion of MCK1 results in inositol depletion due to decreased activity of MIPS.

### VPA-induced MIPS inhibition is Mck1 dependent

In light of our previous reports that VPA indirectly inhibits MIPS [8, 18], our current finding that Mck1 is a positive regulator of MIPS activity suggests a mechanism in which VPA causes inositol depletion by direct or indirect inhibition of Mck1. If VPA inhibition of MIPS activity were mediated by Mck1, then VPA would not cause a further decrease in MIPS activity in the mck1Δ mutant. To address this possibility, MIPS activity was measured in mck1Δ cells grown in the presence or absence of VPA. As shown in Fig 4, WT cells treated with VPA exhibited more than a 40% decrease in MIPS activity. However, VPA did not significantly decrease MIPS activity in mck1Δ cells. These findings suggest that the effects of loss of Mck1 and VPA treatment are not additive, and that VPA-induced MIPS inhibition is Mck1-dependent.
Discussion

The current study demonstrates for the first time that de novo synthesis of inositol is positively regulated by Mck1, which is required for normal activity of the enzyme (MIPS) catalyzing the rate-limiting step of inositol synthesis. Our findings also identify Mck1 as a possible downstream target of VPA.

Among mutants of the four GSK3 homologs, mck1Δ, mrk1Δ, mds1Δ, and ygk3Δ, only mck1Δ cells exhibited decreased intracellular inositol levels, as well as inositol dependent growth at elevated temperature (Fig 1). The mrk1Δmds1Δygk3Δ triple mutant containing only the wild type MCK1 homolog had increased intracellular inositol levels (Fig 1B). These findings indicate that MCK1 is the GSK3 homolog that is necessary and sufficient for normal intracellular inositol levels. Somewhat increased levels of inositol in mrk1Δmds1Δygk3Δ cells suggests that one or more of these genes may negatively regulate inositol synthesis.

MIPS purified from mck1Δ cells exhibited WT levels of MIPS protein but decreased enzymatic activity (Fig 3), suggesting that MCK1 affects enzymatic activity, but not synthesis, of MIPS. Consistent with decreased MIPS activity, the rate of inositol synthesis was decreased.

Fig 3. MIPS activity is decreased in mck1Δ cells. (A) Strains expressing His-Xpress tagged MIPS were constructed by knocking out the INO1 gene with the KanMX gene, which is then replaced with the His-Xpress INO1 fusion gene. (B) Cells expressing the His-Xpress tagged MIPS were grown in SC medium to the early stationary phase. Enzymatic activity of MIPS protein purified from cell extracts was determined as described in “Experimental procedures” (2 independent experiments with 3 replicates each). Values shown are mean ± SEM. (C) MIPS protein levels are not changed in gsk3 mutant cells. 30 g protein from total cell extract were resolved with 10% SDS-PAGE and analyzed by Western blot.

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in Δ cells (Fig 2). These findings strongly support a regulatory role of Mck1 in controlling inositol synthesis. We have previously shown that MIPS is a phosphoprotein with at least three potential phosphosites. While Mck1 has GSK3 kinase enzymatic activity, it does not directly phosphorylate MIPS (data not shown). However, the possibility remains that Mck1 indirectly affects MIPS activity by activating a phosphorylation cascade that phosphorylates the enzyme.

As both MCK1 ablation and VPA lead to similar cellular effects, including inositol depletion [7, 8] and MIPS inhibition (Fig 4), we considered the possibility that they are components of a single mechanism. In support of this, VPA did not significantly decrease MIPS activity in Δ cells (Fig 4). This finding suggests that Mck1 acts downstream of VPA to regulate MIPS. We have previously shown that VPA causes increased phosphorylation of MIPS (17). It is possible that VPA affects MIPS phosphorylation via an Mck1 phosphorylation cascade.

This study demonstrates for the first time that Mck1 is required for VPA-mediated MIPS inhibition, and for optimal inositol synthesis in vivo. These findings have implications for understanding the mechanisms that control inositol synthesis and VPA-induced inositol depletion, and suggest that two common outcomes in published reports of VPA effects, inositol depletion and GSK3 inhibition, are intrinsically related. We suggest that inositol depletion and GSK3 inhibition may be mechanistically inter-related (Fig 5). Accordingly, inositol depletion results from decreased MIPS activity caused by VPA induced inhibition of Mck1/GSK3. Inositol-containing molecules are involved in the regulation of growth, protein secretion, apoptosis, and many other cellular events [3, 46, 47]. The alteration of inositol homeostasis affects hundreds of signaling pathways and leads to perturbation of numerous cellular functions [48, 49], among which are those that are related to the mood-stabilizing effects of VPA.
Author Contributions

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