Asc1 Supports Cell-Wall Integrity Near Bud Sites by a Pkc1 Independent Mechanism

Daniel Melamed, Lavi Bar-Ziv, Yossi Truzman, Yoav Arava*

Department of Biology, Technion – Israel Institute of Technology, Haifa, Israel

Abstract

**Background:** The yeast ribosomal protein Asc1 is a WD-protein family member. Its mammalian ortholog, RACK1 was initially discovered as a receptor for activated protein C kinase (PKC) that functions to maintain the active conformation of PKC and to support its movement to target sites. In the budding yeast though, a connection between Asc1p and the PKC signaling pathway has never been reported.

**Methodology/Principal Findings:** In the present study we found that asc1-deletion mutant (asc1Δ) presents some of the hallmarks of PKC signaling mutants. These include an increased sensitivity to staurosporine, a specific Pkc1p inhibitor, and susceptibility to cell-wall perturbing treatments such as hypotonic- and heat shock conditions and zymolase treatment. Microscopic analysis of asc1Δ cells revealed cell-wall invaginations near bud sites after exposure to hypotonic conditions, and the dynamic of cells’ survival after this stress further supports the involvement of Asc1p in maintaining the cell-wall integrity during the mid-to late stages of bud formation. Genetic interactions between asc1 and pck1 reveal synergistic sensitivities of a double-knock out mutant (asc1Δ/pck1Δ) to cell-wall stress conditions, and high basal level of PKC signaling in asc1Δ. Furthermore, Asc1p has no effect on the cellular distribution or redistribution of Pkc1p at optimal or at cell-wall stress conditions.

**Conclusions/Significance:** Taken together, our data support the idea that unlike its mammalian orthologs, Asc1p acts remotely from Pkc1p, to regulate the integrity of the cell-wall. We speculate that its role is exerted through translation regulation of bud-site related mRNAs during cells’ growth.

Introduction

Asc1p is a member of the WD-40 repeat protein family that adopts a seven-bladed β-propeller structure [1,2]. It was also observed to be a genuine ribosomal protein [1–5], with this function conserved from yeast to human. A cryo-EM study mapped Asc1, as well as its human ortholog, RACK1, to the small subunit head region near the mRNA exit tunnel [1].

RACK1 was initially cloned from rat brain cDNA library as an intracellular Receptor for Activated Protein C Kinase [6], with PKCβII being the preferred binding partner [7,8,9]. Interaction of PKC with RACK1 is thought to hold PKC in an active conformation and to target PKC to appropriate intracellular locations [8,10]. Later studies positioned RACK1 at a central point for multiple cellular functions, as it was found to serve as a scaffold protein for many component from diverse signaling cascades [11–20], with some of them able to interact simultaneously with RACK1, and therefore allow it to integrate inputs from distinct signaling pathways [21].

Based on these observations, it was suggested that RACK1, as a part of the 40 S ribosomal subunit, serves as a docking site for signaling molecules that regulate the activity of translation initiation factors or recruit mRNA-binding proteins to the ribosome [22]. Indeed, it was shown that the mammalian PKCβII interacts with RACK1 while the last associates with translating ribosomes, and that this interaction leads to phosphorylation of the translation initiation factor eIF6, which induce translation initiation [23].

The ability of mammalian RACK1 to compete for localization to the yeast ribosome and to complement phenotypes of asc1 deletion mutant, suggest that these two proteins share similar functions [5]. However, in yeast, the contribution of Asc1p to mRNA translation is not clear. At optimal growth conditions Asc1p is not essential, suggesting that this ribosomal protein is dispensable for the general translation process. In addition, connections between Asc1p and signaling pathways were only recently established. Asc1p was shown to function as G-protein β subunit for the Gz- Gpa1 protein, which is a part of the glucose-stimulated cAMP/PKA signaling pathway, and to inhibit its guanine-nucleotide exchange activity that is required for glucose-signal transmission [24]. In addition, Asc1p was identified as a possible component in the mating pheromone MAPK signaling [25]. However, a connection between Asc1p and the PKC signaling pathway in *Saccharomyces* was never reported.
In the present study, we investigated the relationship between Asc1p and the PKC signaling pathway by following the phenotypes of asc1-deletion mutant and its genetic interactions with PKC1. We point Asc1p as a factor required for the integrity of the cell-wall near the bud site, and suggest that this function is independent of Pkc1p.

Materials and Methods

Yeast strains, plasmids and growth conditions

The following Saccharomyces cerevisiae strains were used: BY4741 (MATa; his3Δ1; leu2Δ0; met15Δ0; ura3Δ0), asc1Δ (MATa, his3Δ1 leu2Δ0 met15Δ0 ura3Δ0; asc1::kanMX4/Euroscarf), pck1Δ (MATa leu2−3 112 ura3−52 tpi1−1 his4 can1r, pkc1Δ; LEU2) and its isogenic wild-type, EG123 (MATa leu2−3 112 ura3−52 tpi1−1 his4 can1r), were kindly provided by Dr. A. Tartakoff [26]. The double knock-out strain asc1Δ/pck1Δ was constructed by sporulation of the mating products of asc1Δ and pck1Δ strains, and selection for germinated spores on synthetic complete media with geneticin (G418) and without leucine. Correct construction of the strain was further verified by Northern blot analysis [27]. The plasmid PDL468 (pGAL-PKC1wt-HA, URA3, CEN) was kindly provided by Dr. M.S. Cyert [29].

Microscopic imaging

To observe cells before and after hypotonic shock treatment, 5 μl of cells were fixed by 4% paraformaldehyde for 10 min and visualized by an Olympus BX61 TRF motorized microscope, equipped with a DP70 digital camera, using a 40 x objective. To follow PKC1-GFP localization, cells were fixed as described, and images were obtained by a Nikon Eclipse 50i microscope with a 100 x immersion objective and recorded by a Nikon DS-5M camera. Images were processed digitally using Adobe Photoshop (Adobe Systems, Inc.).

Zymolase treatment

Cells were grown in YPD plus 1 M NaCl for ~24 hrs to mid-logarithmic phase and concentrated to an OD600 of 5.0 in 2 ml microfuge tube containing 1760 μl YPD supplemented with 1 M NaCl, 200 μl 1 M DTT and 40 μl of 10 mg/ml Zymolase (ImmunoC). At the indicated time points, the turbidity of each sample was measured immediately after the addition of 100 μl aliquot from the reaction tube into a spectrophotometer cuvette already containing 800 μl media and 100 μl of 10% SDS (final conc. 1%), which rapidly lyse cells with severe cell-wall damages. To follow Pck1-GFP re-localization upon cell-wall stress, exponentially growing yeast were cultured in the presence of 0.1 M DTT and 0.2 mg/ml Zymolase for one hour.

Measuring effects of Pck1p mutations on viability

Cells were grown to mid-logarithmic phase in medium lacking uracil and in the presence of 2% glucose at 25°C. Cells were then harvested by centrifugation at 4000 rpm for 4 min at room temperature, washed once with water and diluted in water to a final concentration of 10^6 cells/ml. 100 μl of each sample (~1000 cells)
were seeded on plates lacking uracil and either containing 2% glucose (no induction) or containing 2% galactose and 0.2% sucrose (induction). Colony forming units (CFU) were counted after three days of incubation at 25°C, and the average ratio for glucose/galactose CFU was calculated from three independent repeats.

Cellular fractionation. Cellular fractionation was based on the protocol of Frey et al. [30] for ER-membrane enrichment. Cells were grown to mid-logarithmic phase in YPD media. Cycloheximide was added to a final concentration of 0.1 mg/ml, and cells were harvested (4000 rpm, 4 min, 4°C) and resuspended

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**Figure 2. Response of asc1Δ cells to cell wall stresses.** The indicated strains were grown for 24 hrs at 30°C in liquid YPD media containing 0 M, 0.4 M, 1 M or 1.4 M NaCl (A) or in the presence of 1 M NaCl, 1 M KCl, or 1.5 M Sorbitol (B), or 0.5 M NaCl (C). To impose hypotonic shock, cells were plated in a dilution series on YPD plates containing no supplemental osmolyte and grown either at 30°C ("30°C, Hypotonic shock") or at 37°C ("37°C, Hypotonic shock"). For isotonic control, cells were spotted in a dilution series on YPD plates containing the same osmolyte concentrations as they grew in, and cultured at 30°C ("30°C Control"). Pictures of spotted colonies were taken after 24 hours (left panels) and 48 hours (right panels). To impose calcoflour white stress cells were plated on plates supplemented with 0.5 M NaCl and the indicated calcoflour white concentrations. doi:10.1371/journal.pone.0011389.g002
in 400 µl of ice-cold membrane-fractionation buffer (20 mM Hepes-KOH pH 7.6, 100 mM potassium acetate, 5 mM magnesium acetate, 2 mM dithiothreitol, 0.1 mg/ml cycloheximide and 0.1 mM PMSF) either with or without additional 20 mM EDTA. After the addition of 1 ml glass beads, cells were lysed by two rounds of vigorous vortexing for 90 seconds at 4°C with bead-beater. Recovered lysates were centrifuged for 2 min at 1,200 × g to remove cell debris. The remaining crude lysate was fractionated by centrifugation at 6,000 × g for 20 min at 4°C to cytosol-containing supernatant, and ER membrane-containing pellet [30]. Equivalent amounts of proteins from each fraction were subjected to Western blot analysis.

Western blot analysis. Western analysis was performed as previously described [31]. Anti-HA monoclonal antibody (Covance MMS-101P) was used at a 1:4000 dilution. Rabbit anti-Asc1p, generously provided by Dr. A. Link [5] was used at 1:5000 dilution. Monoclonal anti-Pab1p antibody was a gift from Dr. Mordechai Choder (Technion – Israel Institute of Technology), and was used at 1:10000 dilution. Anti-Hexokinase-HRP conjugated antibody was used at 1:50000 dilution and was kindly provided by Michael Glickman (Technion – Israel Institute of Technology). Anti-mouse IgG-HRP conjugated (Sigma A5906) and Anti-rabbit IgG-HRP conjugated (Sigma A9169), were used at 1:10000 dilution.

Figure 3. Hypotonicity cause rapid reduction in viability of dividing asc1Δ cells. Cells were grown in YPD supplemented with 1 M NaCl either to logarithmic growth phase (A) or to stationary phase (B). Cells were then shifted to media without NaCl to create hypotonic shock. At the indicated time points, dead cells were counted by Methylene Blue (MB) staining.

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Figure 4. asc1Δ cells display aberrant cell-wall morphologies upon exposure to hypotonic conditions. Microscopic observations of wild-type and asc1Δ cells either under sustained growth in rich media (YPD) containing 1M NaCl (A, D), or following one hour exposure to hypotonic shock conditions (shift from 1 M to 0 M NaCl containing YPD media)(B, E), or in hypotonic conditions in the presence of 0.1 mg/ml Cycloheximide (CHX) (C, F). Arrows point to cell-wall deformations sites. Inset in E is a higher magnification of deformed cells.

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Results

asc1Δ strain display phenotypic linkage with the PKC signaling pathway

The mammalian ortholog of Asc1p (RACK1) acts as a receptor for an activated PKCβ isoform [6]. In yeast, the PKC signaling pathway is activated in response to cell-wall stress conditions and controls cell-wall integrity at all stages of cell growth. Accordingly, mutations that interfere with this pathway result in hypersensitivity to various cell-wall perturbing treatments. It is therefore expected that if Asc1p function is directly or indirectly linked to PKC signaling, then loss of asc1 gene will share similar phenotypes with other mutations in the PKC pathway. To study this, we subjected an asc1-deletion mutant (asc1Δ) to staurosporine, a protein kinase inhibitor of which Pkc1p appears to be its primary target [32,33]. Following addition of staurosporine (40 μg/ml) to exponentially growing yeast cultures, asc1Δ strain ceased to proliferate while its parental strain was almost unaffected by the drug (Fig. 1A). Significantly, this growth defect was suppressed by the addition of an osmotic stabilizer (0.8 M NaCl) to the media (Fig. 1B), linking the staurosporine sensitivity of asc1Δ to cell-wall defects resulting from loss of Pkc1p activity.

Additionally, we examined asc1Δ sensitivity to hypotonic stress and to elevated temperatures, two stimuli that damage the fluidity of the cell-wall and activate the PKC signal transduction pathway [34,35]. In order to test the growth response to different strengths of hypotonic shock conditions, cells were grown at 30°C in rich media (YPD) containing different concentrations of NaCl (0, 0.4, 1 or 1.4 M) and after 24 hours serially diluted and seeded on YPD plates containing no NaCl (0 M) (Fig. 2A, left columns). The growth response to elevated temperatures was tested by growing the cells at 37°C (Fig. 2A, middle columns). Monitoring cell growth after 24 and 48 hours demonstrates that following exposure to extreme hypotonic shock conditions (from a 1 M NaCl starting point or above) or to elevated temperature, asc1Δ strain presents delayed resumption of growth and reduced survival, as compared to the wild type strain. The effect on growth was further enhanced when the two stress conditions were combined, with the strongest sensitivity when external osmolarity was reduced from 1 M to 0 M NaCl and growth temperature was 37°C (Fig. 2A dashed-line framed panel). To establish the generality of the sensitivity to hypotonic stress, the same experimental procedure was repeated using KCl and sorbitol (Fig. 2B). Indeed, increased sensitivities to hypotonic conditions and to combined heat shock were demonstrated after the cells were grown with either of the osmolytes, showing that this effect is not restricted to NaCl.

We have also examined the sensitivity of asc1Δ cells to calcoflour white (CFW), a cell-wall damaging agent (Fig. 2C). asc1Δ cells are much more sensitive to CFW than their parental strain (BY4741), as can clearly be seen on the plates supplemented with 100 μg/ml. As expected, cells deleted of Pkc1p are also sensitive to CFW and this can be observed already at a concentration of 50 μg/ml. Note that the parental strain of pkc1Δ (EG123) is much more sensitive to CFW than the parental strain of asc1Δ (BY4741), hence the difference is sensitivity between their progenies.

To follow the dynamics of the loss of viability upon hypotonic shock, we stained asc1Δ with methylene-blue, a dye that stain dead, or physically damaged cells [36,37], at different time points after the cells were subjected to hypotonic shock. Interestingly, while the parental strain remained resistant to methylene-blue staining throughout the experiment, more than half of asc1Δ cells were stained within one minute following the shift to hypotonic conditions with no additional staining at the subsequent time points (Fig. 3A). This result indicates that asc1Δ loss of viability occurs immediately upon exposure to hypotonic treatment, which is consistent with abrupt cell-wall damage. This effect was extremely similar to the one observed for PKC1-deleted cells (pkc1Δ). Yet, the survival of the pkc1Δ strain showed also constant reduction in time (Fig. 3A) and eventually ended with a complete loss of viability (data not shown). When the hypotonic shock was exerted on cells at stationary growth phase, both asc1Δ and pck1Δ were highly resistance to the immediate effect on viability (Fig. 3B). This shows that the two strains share similar vulnerability to hypotonic shock during exponential growth.

Analysis of the cellular morphology of asc1Δ-deletion strain before and after shifting the cells to hypotonic media demonstrated notable cell-wall invaginations that occurred mainly near the sites of the emerging buds (Fig. 4). Importantly, the aberrant cell-wall morphology was not linked to Asc1p function on de-novo synthesis of proteins in response to the hypotonic shock, because arresting the translation process by Cycloheximide had no effect on the morphology of asc1Δ and its parental strain during the hypotonic stress (Fig. 4 compare C to B and F to E). Taken together, these results indicate that Asc1p role in cell-wall metabolism is concentrated mainly in budding sites. Unlike asc1Δ’s response, pkc1Δ cells displayed severe deformations throughout the entire cell contour upon exposure to hypotonic stress conditions (data not shown). This comes in agreement with Pck1p role in regulating and maintaining the integrity of the cell-wall at all steps of cell-cycle.

Genetic interactions suggest that Asc1 and Pkc1 proteins do not act in concert

To further establish the connection between Asc1p and the PKC signaling pathway we tested the effect of two variants of Pkc1 protein on the survival of asc1Δ cells. asc1Δ and its parental strain were transformed with plasmids that promote the expression of an inactive (Pkc1K853R) or a constitutive-active (Pkc1R398A) kinase mutants from a galactose-inducible promoter [32]. We determined the effect of the Pkc1p variants on the viability of each strain by comparing the number of colony forming units (CFU) on plates containing an inducing or non-inducing sugar source (Fig. 5). Over
expressing the wild-type Pkc1p showed no effect on cell survival, in agreement with previous reports [28,32]. However, over-expressing the inactive or the constitutive-active mutants affected the viability of asc1Δ and its parental strain differentially: asc1Δ cells were highly resistant to the deleterious effect of the inactive kinase mutation that was observed in the parental strain (~60% vs. ~2% viability, respectively), and showed increased sensitivity to the constitutive-active form of Pkc1p (~0.15% vs. ~0.6% viability in the parental strain). The simplest explanation for these observations is that asc1Δ cells have higher intrinsic activity of the PKC signaling pathway. The higher PKC activity therefore partially overcome the dominant-negative effect of the kinase-dead mutation and sum up with the constitutive-active mutant to produce a higher toxic effect.

We further constructed a double knockout haploid strain carrying deletions of both ASC1 and PKC1 genes (asc1Δ/pkc1Δ). This strain showed no considerable change in growth rate compared to single deletion mutants of asc1 and pkc1 (Fig. 6A) and presented some of the pke1Δ strain phenotypic hallmarks, such as the existence of giant cells [30] (Fig. 6B), and the necessity for an osmotic stabilizer for growth [35] (Fig. 6C). Yet, it displayed synergistic sensitivities to some cell-wall perturbation treatments: It was unable to grow at elevated temperatures even in the presence of an osmotic stabilizer (Fig. 6C) and showed greater sensitivity to Zymolase (e.g. see the 2.5 min time point at Fig. 6D). This exaggerated sensitivity of the double knockout mutant may indicate that Asc1 and Pkc1 proteins act through different mechanisms to maintain the integrity of the cell-wall.

Asc1p has no effect on Pkc1p localization

One of the outcomes of PKCβII-RACK1 interaction in mammalian cells is the targeting of PKCβII to specific intracellular location, which vary between cell types [6,8,10]. To study whether Asc1p affects the cellular localization of Pkc1p, we fractionated lysates from wild-type and asc1Δ cells to their membranous and cytosolic parts [30] and tested the distribution of Asc1p and Pkc1p between the two compartments. Asc1p appeared to be evenly distributed between the membranous and the cytosolic fractions in wild-type and asc1Δ cells to their membranous and the cytosolic parts [30] and tested the distribution of Asc1p and Pkc1p between the two compartments. Asc1p appeared to be evenly distributed between the membranous and the cytosolic fractions in wild-type and asc1Δ cells [30] (Fig. 7A), similarly as the polysome-associated factor Pab1p. Pkc1p however, appeared almost exclusively in the membranous part (Fig. 7A). This association is independent of Asc1p, as it was unaffected by asc1Δ gene deletion (Fig. 7A). Moreover, EDTA treatment, which leads to disassembly of polysomes and consequently to the release of Asc1 and Pkc1

Figure 6. Phenotypes of asc1Δ/pkc1Δ. A) The indicated strains were grown in rich media in the presence of 0.5 M NaCl to logarithmic growth phase, and diluted to an OD600 of 0.1. Growth was monitored by measuring the OD600 values at the indicated time points. Results of one representative experiment out of three are shown. B) Microscopic morphology of the indicated strains when grown in rich media plus 1 M NaCl. C) Hypotonic shock sensitivity of the indicated strains. Cells were grown to mid-log phase in liquid YPD media supplemented with 0.5 M NaCl, serially diluted and spotted on YPD plates with or without 0.5 M NaCl. Pictures were taken after 48 hrs of incubation at 30 °C. D) Sensitivity to Zymolase. Cells were grown in liquid YPD +0.5 M NaCl to mid-log phase, serially diluted and spotted on YPD plates either with or without added NaCl, or with 0.5 M NaCl. Plates were incubated at 30 °C or at 37 °C. D) Sensitivity to Zymolase. Cells were grown in liquid YPD +1 M NaCl to mid-logarithmic phase and concentrated to an OD600 of 5.0. Samples were then treated with 0.2 mg/ml Zymolase, and at the indicated time points cell-wall sensitivity was determined by adding SDS (final concentration 1%) and measuring the sample’s turbidity.

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in the re-localization of Pkc1p during cell-wall stress. Asc1p has no role in directing Pkc1p to its sites during the budding of the cells' periphery (Fig. 7B). Together, these results suggest that similarly in both strains, forming between five to ten distinct foci at by Zymolase treatment, Pkc1-GFP proteins were re-localized by Pab1p (detected by Hxk1p antibody) was used as a cytosolic marker. B) Pkc1p-GFP was visualized in wild-type and asc1Δ cells grown to mid-log phase either without (“no stress”) or with one hour treatment with Zymolase (“Zymolase stress”).

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Asc1p from the membrane part (Fig. 7A), did not change the membrane association of Pkc1p.

We also followed the localization of a Pkc1-GFP fusion protein that consists of the entire Pkc1p fused through its C-terminus to a green fluorescent protein. Expressing Pkc1-GFP from a high-copy moiety of PKC1 or Asc1p. Pab1p (detected by α-pab1 antibody) was used as a marker for polysomes-associated factor, and Hxk1p (detected by α-Hxk1p antibody) was used as a cytosolic marker. B) Pkc1p-GFP was visualized in wild-type and asc1Δ cells grown to mid-log phase either without (“no stress”) or with one hour treatment with Zymolase (“Zymolase stress”).

Discussion

In mammalian cells RACK1 serves as a scaffold protein for numerous components of diverse signal transduction pathways, of which PKCβII is the most recognized [40]. In this study we show that in the budding yeast, loss of the RACK1 ortholog gene, asc1Δ, results in phenotypes that characterize some PKC signaling mutants. In particular, asc1Δ strain was sensitive to staurosporine (Fig. 1), a specific inhibitor of Pkc1p in yeast [32,33], and showed increased sensitivities to hypotonic- and heat shock conditions and to Zymolase treatment (Figs. 2, 3, 4 and 6D). These are known to provoke a cell-wall stress, which requires a functional PKC signaling for cells to survive.

The dynamics of asc1Δ loss of viability, whereby more than half of asc1Δ cells were damaged within the first minute of exposure to hypotonic conditions, is highly similar to the response of pck1Δ strain (Fig. 3). However, unlike pck1Δ strain that displayed unchanged to slow reduction in viability after the initial drop in survival, asc1Δ strain slowly resumed its growth. This may imply that pck1Δ sensitivity to hypotonic conditions involves two distinct processes, of which asc1Δ susceptibility is linked only to the first, immediate one. Indeed, Levin et al. have shown that PKC1-deleted cells that possessed buds of any size underwent immediate lysis upon transfer to medium lacking osmotic stabilizers, while non budded cells arrested at the early stages of bud formation [33,41]. Therefore, whereas Pck1p function is important to all stages of bud formation Asc1p appears to have a role only in the mid-to late stages of this process.

Several observations in this work suggest that Asc1p is connected to Pck1p by a different mechanism then in higher eukaryotes. First, in mammalian cells, RACK1 serves as a scaffold that mediates the phosphorylation and activation of the MAPK JNK by PKCβII [42]. However, our data suggest that asc1Δ cells contain high basal levels of PKC signaling (Fig. 5), which comes in agreement with the hyper-phosphorylation of the terminal MAPK, Slt2/Mpk1p, in asc1Δ background [25]. This observation raises the possibility that the PKC signaling is activated in asc1Δ strain to compensate for its cell-wall sensitivity due to loss of function of other mechanism. Second, in mammalian cells PKCβII-RACK1 interaction appears to target PKCβII to distinct intracellular locations, which vary between different cell types [6,8,10]. In yeast, full-length Pck1p appears to reside predominantly at the bud-neck of medium to large sized buds and at the tip of small-sized buds, and to become re-localized to the cell’s periphery upon exposure to cell-wall stress [29,39]. Our results show that loss of Asc1p has no effect on Pck1p localization at steady-state growth conditions nor on its re-distribution after cell-wall stress (Fig. 7B), and has no influence on Pck1p fractionation with membrane-compartments (Fig. 7A). Third, while a physical interaction between Asc1 and Pkc1 proteins cannot be ruled out, we were unable to support these by two hybrid analyses or co-immunoprecipitation (data not shown). Moreover, the genetic interactions between the two genes suggest that they do not act in the same pathway. Specifically, double-knockout mutant lacking both asc1 and pck1 genes displayed synergistic sensitivities to cell-wall stress conditions (Fig. 6C, D). Taken together, our observations suggest that in S.cerevisiae, Asc1p contribution to cell-wall integrity is not through the conventional Bck1-Mkk1/2-Mpk1 MAPK module that acts downstream to Pck1p. Rather, Asc1p appears to function in parallel to Pck1p, or to coordinate between the PKC signaling pathway and other cell-wall integrity related mechanisms, to the most.

How might Asc1p affect the integrity of the cell-wall? A likely mechanism is by regulating the translation of mRNAs that encode for cell-wall proteins. Indeed, mutations in Asc1p that were shown to reduce its ability to associate with ribosomes, resulted in also increased sensitivity to calcofluor white, a cell-wall perturbing agent [2]. Based on the immediate lysis of asc1Δ cells upon
hypotonic shock conditions (Fig. 3) that was independent of de-
ovo synthesis of proteins (Fig. 4), we suggest that regulation of
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Author Contributions

Conceived and designed the experiments: DM YA. Performed the experi-
ments: DM LBZ YT. Analyzed the data: DM. Wrote the paper: DM
YA.

References

1. Sengupta J, Nilsson J, Gursky R, Spahn CM, Nisen P, et al. (2004)
Identification of the versatile scaffold protein RACK1 on the eu-
aryotic ribosome by cryo-EM. Nat Struct Mol Biol 11: 957–962.
2. Coyle SM, Gilbert WV, Doudna JA (2009) Direct link between RACK1
function and localization at the ribosome in vivo. Mol Cell Biol 29: 1626–1634.
3. Link AJ, Eng J, Schielitz DM, Carmack E, Mize GJ, et al. (1999) Direct analysis
of protein complexes using mass spectrometry. Nat Biotechnol 17: 676–682.
4. Inada T, Winstall E, Taran SZ, Jr., Yates JR, 3rd, Schielitz D, et al. (2002) One-
step affinity purification of the yeast ribosome and its associated proteins and
mRNAs. RNA 8: 948–958.
5. Gerbasi VR, Weaver CM, Hill S, Friedman DB, Link AJ (2004) Yeast Asc1p
and mammalian RACK1 are functionally orthologous core 40 S ribosomal
proteins that repress gene expression. Mol Cell Biol 24: 8272–8277.
6. Ron D, Chen CH, Caldwell J, Jamieson L, Orr E, et al. (1994) Cloning of an
intracellular receptor for protein kinase C: a homolog of the beta subunit of G
proteins. Proc Natl Acad Sci U S A 91: 839–843.
7. Coskun M, Mochly-Rosen D (1999) Pharmacological modulation of protein kinase
C isozymes: the role of RACKs and subcellular localisation. Pharmacol Res 39:
253–259.
8. Ron D, Jiang Z, Yao L, Vagts A, Diamond I, et al. (1999) Coordination
movement of RACK1 with activated betaIIPKC. J Biol Chem 274:
27039–27046.
9. Stebbins EG, Mochly-Rosen D (2001) Binding specificity for RACK1 resides in
the V5 region of beta II protein kinase C. J Biol Chem 276: 29844–29850.
10. Ron D, Luo J, Mochly-Rosen D (1995) C2-region-derived peptides inhibit
translocation and function of beta protein kinase C in vivo. J Biol Chem 270:
24180–24187.
11. Chang BY, Conroy KB, Machleder EM, Cartwright CA (1998) RACK1, a
receptor for activated protein kinase C and a homolog of the beta subunit of G
proteins. Proc Natl Acad Sci U S A 91: 839–843.
12. Yarwood SJ, Steele MR, Scotland G, Houslay MD, Bolger GB (1999) The
movement of RACK1 with activated betaIIPKC. J Biol Chem 274:
14896–14901.
13. Ceci M, Gaviraghi C, Gorrini C, Sala LA, Offenbauer N, et al. (2003) Release
of eIF6 (p27BP) from the 60 S subunit allows 80 S ribosome assembly. Nature
420: 579–584.
14. Zeller CE, Parnell SG, Dohmann HG (2007) The RACK1 ortholog Asc1
functions as a G-protein beta subunit coupled to glucose responsiveness in yeast.
J Biol Chem 282: 25168–25176.
15. Chasse SA, Flanary P, Parnell SC, Hao N, Cha JY, et al. (2006) Genome-scale
analysis reveals Sec6 as the principal regulator of mating pheromone signaling in
the yeast Saccharomyces cerevisiae. Eukaryot Cell 5: 330–346.
16. Nanduri J, Tartakoff AM (2001) Perturbation of the nucleolus: a novel HOG1-
dependent, Pck1p-dependent consequence of hypertonic shock in yeast. Mol
Cell Biol 11: 1035–1041.
17. Eliyahu E, Purolt L, Melamed D, Scherrer T, Gerber AP, et al. (2010) Tom20
mediates localization of mRNAs to mitochondria in a translation-dependent
manner. Mol Cell Biol 30: 264–294.
18. Gray JV, Ogo JP, Kamadu Y, Stone M, Levin DE, et al. (1997) A role for the
Pck1 MAP kinase pathway of Saccharomyces cerevisiae in bud emergence and
identification of a putative upstream regulator. Embo J 16: 4924–4937.
19. Denis V, Cyert MS (2005) Molecular analysis reveals localization of Saccharomyces
cerevisiae protein kinase C to sites of polarized growth and Pck1p targeting to the
nucleus and mitotic spindle. Eukaryot Cell 4: 36–45.
20. Frey S, Pool M, Seedorf M (2001) Sep160p, an RNA-bonding, polyome-
associated protein, localizes to the endoplasmic reticulum of Saccharomyces
cerevisiae in a microtubule-dependent manner. J Biol Chem 276: 13905–13912.
21. Loya A, Purolt L, Yosefoz Y, Wedles Y, Zuc-Ukelson M, et al. (2000) The 5’-
UTR mediates the cellular localization of an mRNA encoding a short plasma
membrane protein. Rna 14: 1352–1365.
22. Watanabe M, Chen CY, Levin DE (1994) Saccharomyces cerevisiae PKC1
encodes a protein kinase C (PKC) homolog with a substrate specificity similar to
that of mammalian PKC. J Biol Chem 269: 16829–16836.
23. Yoshida S, Ikeda E, Uno J, Mitsuhashi H (1992) Characterization of a staurosporine-
tolerant temperature-sensitive mutant, stt1, of Saccharomyces cerevisiae: NTF1 is allelic to PKC1. Mol Gen Genet 231: 337–344.
24. Kamada Y, Jang US, Petrowski J, Levin DE (1995) The protein kinase C-
coupled MAP kinase pathway of Saccharomyces cerevisiae mediates a novel
aspect of the heat shock response. Genes Dev 9: 1559–1571.
25. Levin DE, Bartlett-Heusser E (1992) Monoclonal in the S. cerevisiae PKC1 gene
display a cell cycle-specific osmotic stability defect. J Cell Biol 116: 1221–1229.
26. Korcak W, Vairo ML (1958) A simple colorimetric method for detecting cell
viability in cultures of eukaryotic microorganisms. Current microbiology
7: 217–221.
27. Paravicini G, Cooper M, Friedli L, Smijh DJ, Carpenier JL, et al. (1992) The
osmotic integrity of the yeast cell requires a functional PKC1 gene product. Mol
Cell Biol 12: 4986–4995.
28. Andrews PD, Stark MJ (2000) Dynamic, Rhe1p-dependent localization of Pck1p
to sites of polarized growth. J Cell Sci 113 (Pt 15): 2085–2093.
29. McCafferty A, Warwicker J, Bolger GB, Houlay MD, Warzyw SD (2002) The
RACK1 scaffold protein: a dynamic cog in cell response mechanisms. Mol
Pharmacol 62: 1261–1273.
30. Levin DE, Bowers B, Chen CY, Kamadu Y, Watanabe M (1994) Dissecting the
protein kinase C/MAP kinase signalling pathway of Saccharomyces cerevisiae.
Cell Biol Mol Biol Res 40: 229–236.
31. Loya A, Purolt L, Yosefoz Y, Wedles Y, Zuc-Ukelson M, et al. (2000) The 5’-
UTR mediates the cellular localization of an mRNA encoding a short plasma
membrane protein. Rna 14: 1352–1365.
32. Watanabe M, Chen CY, Levin DE (1994) Saccharomyces cerevisiae PKC1
encodes a protein kinase C (PKC) homolog with a substrate specificity similar to
that of mammalian PKC. J Biol Chem 269: 16829–16836.
33. Yoshida S, Ikeda E, Uno J, Mitsuhashi H (1992) Characterization of a staurosporine-