Candida albicans Tpk1p and Tpk2p isoforms differentially regulate pseudohyphal development, biofilm structure, cell aggregation and adhesins expression

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Abstract

Candida albicans undergoes a reversible morphological transition from single yeast cells to pseudohyphal and hyphal filaments. In this organism, cAMP-dependent protein kinase (PKA), coded by two catalytic subunits (TPK1 and TPK2) and one regulatory subunit (BCY1), mediates basic cellular processes, such as the yeast-to-hypha transition and cell cycle regulation. It is known that both Tpk isoforms play positive roles in vegetative growth and filamentation, although distinct roles have been found in virulence, stress response and glycogen storage. However, little is known regarding the participation of Tpk1p and/or Tpk2p in pseudohyphal development. This point was addressed using several C. albicans PKA mutants having heterozygous or homozygous deletions of TPK1 and/or TPK2 in different BCY1 genetic backgrounds. We observed that under hypha-only inducing conditions, all BCY1 heterozygous strains shifted growth toward pseudohyphal morphology; however, the pseudohypha : hypha ratio was higher in strains devoid of TPK2. Under pseudohypha-only inducing conditions, strains lacking TPK2 were prone to develop short and branched pseudohyphae. In tpk2Δ/tpk2Δ strains, biofilm architecture was markedly less dense, composed of short pseudohyphae and blastospores with reduced adhesion ability to abiotic material, suggesting a significant defect in cell adherence. Immunolabelling assays showed a decreased expression of adhesins Als1p and Als3p only in the tpk2Δ/tpk2Δ strain. Complementation of this mutant with a wild-type copy of TPK2 restored all the altered functions: pseudohyphae elongation, biofilm composition, cell aggregation and adhesins expression. Our study suggests that the Tpk2p isoform may be part of a mechanism underlying not only polarized pseudohyphal morphogenesis but also cell adherence. Copyright © 2011 John Wiley & Sons, Ltd.

Keywords: Candida albicans; PKA; pseudohyphal development; biofilm architecture; adhesins

Introduction

Candida albicans is a pathogenic fungus that exists in multiple cell forms, including yeast, pseudohyphae and hyphae. Although pseudohyphae superficially resemble true hyphae, it is becoming increasingly clear that there are fundamental differences between the two forms in the organization of the cell cycle and the septin cytoskeleton (Sudbery et al., 2004). Pseudohyphal cells are highly branched, ellipsoidal in form and have constrictions at cell junctions where hyphal cells...
have parallel sides and lack constrictions at their septa. Other differences include nuclear division and the presence of a specialized structure only present in hyphal cells, the Spitzenkörper, which is important for concentrating and delivering secretory vesicles to the cell tip (Braun et al., 2000; James et al., 2006). Because of the extensive phenotypic differences between hyphal and pseudohyphal cells, it is generally believed that expression of different gene sets specifies the morphology, although definitive evidence for this is lacking. C. albicans yeast cells can be induced to form hyphae or pseudohyphae by a variety of environmental signals, such as the presence of serum, 35–37 °C growth temperature and neutral pH (Odds, 1988; Sudbery, 2001). Although the mechanism underlying pseudohyphae formation is still unknown, several authors have identified specific proteins involved in the process (Li et al., 2006; Sherwood and Bennett, 2008; Trunk et al., 2009). A recent report by Carlisle et al. (2009) demonstrated a continuous transition from yeast to pseudohyphae, controlled by the dosage-dependent expression of a filament-specific transcriptional regulator, Ume6p.

In C. albicans, the cAMP/cAMP-dependent protein kinase (PKA) pathway mediates basic cellular processes, including dimorphic transitions. Positive roles have been established for both catalytic isoforms Tpk1p and Tpk2p in hyphae formation (Bockmühl et al., 2001; Cloutier et al., 2003). Jung and Stateva (2003) have demonstrated that in liquid media, constitutive activation of the cAMP pathway following deletion of PDE2, encoding the high cAMP-affinity phosphodiesterase, inhibits hyphal but not pseudohyphal growth. In addition, it was reported that overexpression of the transcription factor EFG1 results in a pseudohyphal growth form rather than true hyphae (Stoldt et al., 1997; Tebarth et al., 2003). More recently, we have shown that heterozygous strains for the regulatory subunit of PKA (BCY1) in a wild-type and in a null TPK2 genetic background promote pseudohyphal growth in several true hyphae-inducing liquid media (Giacometti et al., 2006). In this work, we investigated pseudohyphal development of several PKA mutants having heterozygous or homozygous deletions of TPK1 and/or TPK2 in different BCY1 genetic backgrounds. We provide genetic evidence that deletion of TPK2 but not TPK1 upholds pseudohyphal growth under true hyphae-inducing conditions. Under pseudohyphae-inducing conditions, tpk2 Δ/tpk2 Δ strains were severely impaired in pseudohypha extension. We also show that in these strains biofilm architecture is structurally weakened. Immunolabelled hyphae indicated that expression of adhesins Als1p and Als3p is diminished in the tpk2 Δ/tpk2 Δ mutant in comparison with the tpk1 Δ/tpk1 Δ strain. Our work shows that, although not required for initial germ-tube formation, Tpk2p is necessary for pseudohypha extension. Furthermore, our data indicate that Tpk2p plays an important role in biofilm formation and cell adhesion to various supports, including silicone surfaces used in medical devices.

**Materials and methods**

**Chemicals**

Reagents were obtained from the following sources: calcofluor white (CFW) and kemptide (LRRASLG) were from Sigma; phosphocellulose paper P-81 was from Whatman; [32P]ATP and [3H]cAMP were from New England Nuclear; polyvinylidenedifluoride (PVDF) membranes (Immobion-P) were from Millipore; MDY4-64, Alexa Fluor 488 and concanavalin A (Alexa Fluor 594) were from Molecular Probes–Invitrogen; all restriction enzymes used were from Promega; ‘Complete Mini’ protease mix was from Roche. All other chemicals were of analytical grade.

**Strains, media, and culture conditions**

We performed the studies with new and previously obtained C. albicans PKA mutant strains, all derived from the wild-type strain CAI4, as detailed in Table 1. Yeast cells were cultured at 30 °C in YPD (1% yeast extract, 2% peptone, and 2% dextrose) or in SD minimal medium (Sherman et al., 1986).

In all strains used in this study, the URA3 gene was re-established with the CIP10 vector (Murad et al., 2000), ensuring URA3 expression at the neutral RPS10 locus.

**Heat stress sensitivity test**

Sensitivity to heat shock was assessed as previously described (Giacometti et al., 2009). Briefly, cells of wild-type and PKA mutants from the stationary
phase were streaked out on YPD plates and incubated at 50°C for 30 min, 1 h and 2 h. At these time points, the plates were shifted to 30°C and growth was analysed after 2 days.

Filamentation assays

*C. albicans* inoculated from a freshly grown YPD plate was cultured for 16 h at 30°C in liquid YPD, at which point the culture was 95% unbudded and the cell density was 2 × 10⁸ cells/ml. To promote hypha-only formation, the culture was diluted (to 1 × 10⁷ cells/ml) with pre-warmed liquid YPD medium (37°C) supplemented with 10% FBS or in pre-warmed minimal medium (37°C) containing 10 mM GlcNAc (Shepherd *et al*., 1980). The cells were then cultured at 37°C. Alternatively, to promote pseudo-hypha-only formation, the cells were incubated in liquid YPD at 35°C (Sudbery, 2001). Chitin staining was done by directly adding 1 μg calcofluor white (CFW) to 100 μl of cell suspension, followed by 15 min incubation and washing. Vacuoles were stained with the lipophylic dye MDY4-64 (Cole *et al*., 1998). Nuclei were stained with DAPI. The cells were visualized with an Olympus BX50 fluorescence microscope. Images were taken with a Cool SNAP-Pro colour digital camera kit with Image Pro Plus software (Media Cybernetics).

**Anti-Als immunolabelling of cultured germ tubes**

The experiments were performed essentially as described by Coleman *et al*., (2009). Solid YPD cultures were stored at 4°C for no more than 1 week before a fresh plate was prepared. Starter cultures were grown to saturation by inoculating a single colony into 10 ml YPD liquid medium and incubating for 16 h at 30°C. For germ tube assays, yeast cells from the saturated starter culture were washed with phosphate-buffered saline (PBS) and suspended at a density of 5 × 10⁶ cells/ml in 3% paraformaldehyde. Fixed *C. albicans* cells were washed three times with PBS prior to and between each step in the immunolabelling protocol. Cells were resuspended in 15 μl of normal goat serum for 15 min at room temperature to block non-specific antibody binding. The cells were incubated in 18 μg/ml anti-Als1 and anti-Als3 monoclonal antibodies in PBS for 60 min at 4°C on a rotating mixer and then in 3 μg/ml Alexa Fluor

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**Table 1. *C. albicans* strains used in this study**

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<tr>
<th>Strain</th>
<th>Genotype</th>
<th>Source or reference</th>
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<tr>
<td>CAH</td>
<td>ura3Δ::λimm434/ura3Δ::λimm434</td>
<td>Forzi and Irwin, 1993</td>
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<tr>
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<td>Same as CAH but RPS10/tps10Δ::Clp10</td>
<td>Giacometti <em>et al</em>., 2009</td>
</tr>
<tr>
<td>RGS5</td>
<td>Same as CAH but Bcy1Δ::Cat RPS10/tps10Δ::Clp10</td>
<td>Giacometti <em>et al</em>., 2009</td>
</tr>
<tr>
<td>RGI1.1</td>
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<td>This study</td>
</tr>
<tr>
<td>RIU1.1</td>
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<td>This study</td>
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<tr>
<td>I1H6-4a</td>
<td>Same as CAH but tpk1Δ::hisG/tpk1Δ::hisG</td>
<td>Boekmühl <em>et al</em>., 2001</td>
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<td>RS1u</td>
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<tr>
<td>RGI2</td>
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<td>This study</td>
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<td>RGI2.1u</td>
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<td>This study</td>
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<td>R2U2.1</td>
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<td>RS2u</td>
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</tr>
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<td>RSI1u</td>
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<tr>
<td>BBA1u</td>
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<td>This study</td>
</tr>
<tr>
<td>ECI1u</td>
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<td>HPY421</td>
<td>tpk2Δ::Glp47/tpk2Δ::Glp47 TPK1/tpk1Δ::hisG/tpk1Δ::hisG Δ::URA3-dpl200</td>
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<tr>
<td>HPY321</td>
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Table 2. Primers used in this study

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<th>Name</th>
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<td>BCY1KO5</td>
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<td>CTAAGTAGTAAAATGAGTGAGAGGGAGAGACAGAGA</td>
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<td>BCY1KO3</td>
<td>Reverse</td>
<td>GCGAGAGGGGGAGTAAAAGTTTTCCCAAGTACGACGTT</td>
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<tr>
<td>URA3ver5</td>
<td>Forward</td>
<td>TTCCGAGGCTTGGCGTAATCAT</td>
</tr>
<tr>
<td>BCY1ver3</td>
<td>Reverse</td>
<td>GGAAAGCAAAAAGTGTAAACAA</td>
</tr>
<tr>
<td>RT1-BCY1</td>
<td>Forward</td>
<td>ATGTCTAATCTCAACAACA</td>
</tr>
<tr>
<td>RT2-BCY1</td>
<td>Reverse</td>
<td>TTAATGACCAGCAGTTGG</td>
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<td>RT1-TPK1</td>
<td>Forward</td>
<td>AAGAAGTTCAAGATGTTGACTTAT</td>
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<tr>
<td>RT2-TPK1</td>
<td>Reverse</td>
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<td>RT1-TPK2</td>
<td>Forward</td>
<td>GAGTTTAGACCGTTACAGTGG</td>
</tr>
<tr>
<td>RT2-TPK2</td>
<td>Reverse</td>
<td>ACTGCTGATTTAGCAAGAG</td>
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<tr>
<td>RT1-ACT1</td>
<td>Forward</td>
<td>CCCAGCTTGGCGGTAGGCGGCT</td>
</tr>
<tr>
<td>RT2-ACT1</td>
<td>Reverse</td>
<td>GTGGTGACAAATGTGGAGACCA</td>
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</table>

488-conjugated donkey anti-mouse IgG under the same conditions. Wet mounts of immunolabelled cells were examined using an Olympus FV300 confocal scanning laser microscope with a fluorescein isothiocyanate protocol and with excitation at 488 nm (Argon laser). The lenses used included Zeiss UplanFl 20x/0.5 and UplanApo 40x/1.0 water immersion objectives. Images were captured and processed using Photoshop v 5.5 software (Adobe Systems, San Jose, CA, USA).

DNA manipulations

DNA purifications were performed with Qiagen affinity columns, following the manufacturer’s recommendations. Bacterial plasmid DNA was isolated by the alkaline lysis method (Sambrook et al., 1989) or using the QIAprep Spin Miniprep Kit (Qiagen). Yeast genomic DNA was isolated according to Adams et al. (1997). DNA modifying enzymes were used according to the manufacturers’ recommendations.

Generation of *C. albicans* tpk1 Δ/tpk1 Δ BCY1/bcy1 Δ mutant strain

 Knockout of one allele of the *BCY1* gene was generated using the PCR-based adaptation (Wilson et al., 1999) of the sequential URA-Blaster technique (Fonzi and Irwin, 1993), as previously described in Giacometti et al. (2009). Specific primers BCY1KO5 and BCY1KO3 were designed to generate the PCR deletion construct BCY1::dpl200-URA3-dpl200. The products of 10 PCR reactions were pooled and used to transform CAI4-derived strain tpk1 Δ/tpk1 Δ (IIHH6-4a), following the protocol described by Wilson et al. (1999). This technique allowed us to obtain strain tpk1 Δ/tpk1 Δ BCY1/bcy1 Δ (RG12). URA3 transformants were grown on uridine-deficient SD solid medium, and proper genomic insertion of the transforming cassette was determined by a PCR-based analysis of transformed colonies, using a set of primers combining a forward oligo internal to the URA3 cassette (URA3ver5) and a reverse one external to the modified region (BCY1ver3). From 21 independent isolations for mutant tpk1 Δ/tpk1 Δ BCY1/bcy1 Δ, 13 colonies showed heterozygous loss of the *BCY1* allele. Three tpk1 Δ/tpk1 Δ BCY1/bcy1 Δ clones rendered identical phenotypes in the characterization assays. All primers used are detailed in Table 2.

To evaluate whether *URA3* prototrophy located in the modified gene of interest affected the observed phenotypes, we performed all tests in the entire set of mutants expressing *URA3* at the neutral RPS10 locus. To that end, the *URA3* marker was recycled by selection on SD medium plus 5-FOA (1 mg/ml) and uridine (50 µg/ml), and all *ura3* mutants were transformed with *Stu* I-digested CIp10 plasmid (Murad et al., 2000) in order to avoid potential problems associated with ectopic expression of *URA3* (Brand et al., 2004).

Preparation of cell extracts and enzymatic assays

Yeast cells (1–2 × 10⁷) from stationary phase were suspended in 500 µl 10 mM sodium phosphate
buffer, pH 6.8, containing 1 mM EGTA, 1 mM EDTA, 10 mM 2-mercaptoethanol and one tablet of ‘Complete mini’ protease mix per 10 ml. All manipulations were thereafter performed at 4 °C. Cells were lysed by disruption with glass beads as described previously. The resulting suspension was spun down in a microfuge at maximum speed for 30 min and the supernatant was centrifuged for 45 min at 100,000 × g. This second supernatant was used immediately for enzymatic assays. PKA activity was measured as previously described (Giacometti et al., 2009), using [γ-32P]ATP as the phosphate donor. PKA-specific activity was expressed as pM 32P incorporated to kemptide per min and per mg of protein. The reactions were carried out under conditions of linearity with respect to the amount of extract and the time of incubation.

RNA isolation and semi-quantitative RT–PCR

Total RNA was isolated from stationary growth cells by the hot-phenol method (Auskubel et al., 1994). Isolated RNA was then treated with DNase at 37 °C for 30 min. The SuperScript First-Strand Synthesis Kit for RT–PCR (Invitrogen) was then used to synthesize cDNA. RNA concentration was measured spectrophotometrically and 2 µg were added to the cDNA synthesis reaction. OligodT (Invitrogen) was used to prime the cDNA synthesis reaction. One-tenth volume of the final cDNA product was added to PCR reactions specific for each gene. Primer sequences for BCY1 (RT1-BCY1 and RT2-BCY1), TPK1 (RT1-TPK1 and RT2-TPK1) and TPK2 (RT1-TPK2 and RT2-TPK2) are detailed in Table 2. Samples were denatured at 94 °C for 2 min, followed by 15–30 cycles of (94 °C for 45 s, 55 °C for 45 s and 72 °C for 30 s). The levels of amplified products were determined at several cycle intervals to ensure that samples were analysed during the exponential phase of amplification. We performed reactions without reverse transcriptase to control for the presence of contaminating DNA. A 900 bp PCR product amplified with RT1-ACT1 and RT2-ACT1 primers from C. albicans ACT1 was used as internal mRNA loading control. Transcripts were quantified using ImageJ (Abramoff et al., 2004).

Biofilm formation

Development of biofilms was performed using a modification of a protocol described by Chandra et al. (2001). Briefly, C. albicans strains were grown in SD medium plus 50 mM glucose, pH 7.0, at 30 °C for 16 h with orbital agitation. Cells were then centrifuged and resuspended in PBS to OD965 = 1.0 and 100 µl of cells were allowed to adhere for 90 min at 37 °C to wells of a sterile 96-well polystyrene microtitre plate (Falcon) coated with 10% FBS. Cells that did not adhere were removed by washing twice with 200 µl PBS. SD medium supplemented with glucose was then added to each well and the biofilms allowed to form at 37 °C for up to 48 h.

For crystal violet staining, each well was washed twice with PBS, dried for 1 h at room temperature and stained with 110 µl 0.4% aqueous crystal violet solution for 45 min. The wells were then washed four times with 200 µl MilliQ sterile water and destained with 200 µl 95% ethanol.
45 min, 100 µl destaining solution was withdrawn from each well and transferred to a new plate that was measured with a microtitre plate reader (Multiskan MS, Labsystems, Finland) at 540 nm. The absorbance values of the controls were subtracted from the values of the test wells to minimize background interference. Three independent experiments were performed with 10 replicates for each strain.

For the XTT reduction assay, an XTT solution was prepared (1 mg/ml in PBS) and filter-sterilized through a 0.22 µm pore size filter. Menadione solution (0.4 mM in acetone) was filtered and mixed with XTT solution at a ratio of 1:5 by volume before the assay. After biofilm formation, the wells were washed three times with 200 µl PBS. To each well, 200 µl PBS and 12 µl XTT–menadione solution were added. The culture plate was incubated in the dark for 2 h at 37°C, then 100 µl of each sample were transferred to a new plate and measured with a microtitre plate reader at 492 nm. The absorbance values of the controls were subtracted from the values of the test. The experiment was performed three times, using 10 replicates for each strain.

To analyse the adhesion and biofilm formation of C. albicans strains to a relevant medical device, we evaluated the ability of strains to adhere and form biofilms on a catheter silicone surface. In this experiment, the strains were grown overnight in YPD at 30°C, diluted to OD600 = 0.5 in 2 ml Spider medium (Liu et al., 1994) and added to a sterile 12-well plate containing a silicone elastomer square, pretreated overnight with 10% FBS. The inoculated plate was incubated at 37°C for 90 min with shaking to allow the initial adhesion of cells. The squares were washed with 2 ml PBS and placed into 12-well plates containing 2 ml fresh Spider medium and incubated at 37°C for an additional 48 h to allow biofilm formation. The biofilm mass on the silicone substrate was observed by concanavalin A staining, as described by Kuhn et al. (2002). Cells were visualized with an Olympus FV300 confocal scanning laser microscope with a rhodamine-fluorescein isothiocyanate protocol [excitation at 543 nm (HeNe laser) and emission at 560 nm]. The lenses used included Zeiss UplanFI ×20/0.5 and UplanApo ×40/1.0 water immersion objectives. Images were captured and processed using Photoshop v 5.5 software.

Measurement of pseudohyphal length and constrictions

Quantitative cell measurements of microscopic images were made using ImageJ version 1.42I software. The average diameters and the intersepta distances were determined from measurements of CFW stained cells; more than 200 individual pseudohyphal filaments per strain were measured. Pseudohyphal lengths and diameters were traced directly from images and constriction frequency was measured by assessing: (a) the average numbers of compartments from the pseudohyphal apex to the first constriction; and (b) the percentage of subapical pseudohyphal compartments that had a branch or lateral bud.

Results and discussion

Absence of Tpk2p isoform enriched pseudohyphae population under hypha-only inducing conditions

Previous work from our laboratory showed that strains lacking a BCY1 allele in a wild-type and in TPK2 null genetic background exhibited a mixture of true hyphae and pseudohyphae in several liquid inducing media (Giacometti et al., 2006). Therefore, it seemed relevant to examine the germinal phenotype of a heterozygous BCY1 mutant in a background devoid of TPK1 isoform, since a recent report from our laboratory showed that Tpk1p and Tpk2p isoforms have distinct roles in the stress response pathway and in glycogen accumulation (Giacometti et al., 2009).

With this in mind, we performed the heterozygous disruption of C. albicans BCY1 in the strain tpk1Δ/tpk1Δ (see Material and methods). The phenotype and biochemical characteristics of the tpk1Δ/tpk1Δ BCY1/bcy1Δ (RG12.1u) strain are summarized in Table 3 and Figure 1. PKA activity was routinely measured at the stationary phase because levels of TPK1 and TPK2 mRNA expression are at the maximum at this stage, allowing the most discriminating comparisons (Souto et al., 2006; Giacometti et al., 2009).

The vacuolar morphology of strain tpk1Δ/tpk1Δ BCY1/bcy1Δ resembled that previously described in strain tpk2Δ/tpk2Δ BCY1/bcy1Δ (Giacometti et al., 2006), with the cytoplasm occupied by a single abnormally large vacuole. Both tpk null
strains harbouring both BCY1 alleles showed the same vacuolar phenotype as wild-type RGI4 (Figure 1A). The \( \text{tpk1}^\text{Δ} \Delta \text{tpk1}^\text{Δ} \text{BCY1}/\text{bcy1}^\text{Δ} \) mutant strain exhibited a relatively high PKA-specific activity (Table 3), provided by the more abundant Tpk2p isoform (Cloutier et al., 2003; Souto et al., 2006), and a low cAMP-binding activity (Figure 1D), which correlated well with a diminished mRNA and Bcy1p expression (Figure 1B, C, E). The fact that cAMP binding of \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \text{bcy1}^\text{Δ}/\text{bcy1}^\text{Δ} \) strain was negligible reinforces the idea that all binding activity corresponded to the Bcy1p subunit.

The \( \text{tpk1}^\text{Δ} \Delta \text{tpk1}^\text{Δ} \text{BCY1}/\text{bcy1}^\text{Δ} \) mutant was extremely sensitive to heat exposure (Figure 2, streak 4), due to the presence of the less regulated Tpk2p isoform, which we have reported to be involved in conferring thermosensitivity to the cell (Giacometti et al., 2009). Identical results were obtained with three independently isolated clones (data not shown).

We then characterized the ability of \( \text{tpk1}^\text{Δ} \Delta \text{tpk1}^\text{Δ} \text{BCY1}/\text{bcy1}^\text{Δ} \) mutant strain to shift from yeast to hyphal growth (YPD medium plus 10% FBS at 37°C) and compared it with previously obtained \( \text{BCY1}^\text{Δ}/\text{Delta1} \) strains (Giacometti et al., 2006). The results obtained are shown in Figure 3. Under this hypha-only inducing condition, the wild-type strain exhibited about 90–97% hyphae and 1–3% pseudohyphae. In contrast, we found that the other strains differed in the proportion of pseudohyphae formed. Similarly to other heterozygous \( \text{BCY1}^\text{Δ}/\text{Delta1} \) mutant strains, \( \text{tpk1}^\text{Δ} \Delta \text{tpk1}^\text{Δ} \text{BCY1}/\text{bcy1}^\text{Δ} \) produced a mixture of hyphae and pseudohyphae. However, strain \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \text{BCY1}/\text{bcy1}^\text{Δ} \) exhibited the higher proportion of pseudohyphae; even strain \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \text{BCY1}/\text{bcy1}^\text{Δ} \) harboured both \( \text{BCY1}^\text{Δ}/\text{Delta1} \) alleles, also produced a significant amount of pseudohyphae, very probably due to the lower expression of \( \text{BCY1}^\text{Δ}/\text{Delta1} \) in this strain (Giacometti et al., 2009; Souto, PhD Thesis, 2006). Complementation of the \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \) mutant with a wild-type copy of \( \text{TPK2} \) increased the levels of hyphae formation and reduced pseudohyphae proportion, suggesting a key role for Tpk2p in the hyphae/pseudohyphae-forming decision. Although the effect was less noticeable, \( \text{tpk1}^\text{Δ} \Delta \text{tpk1}^\text{Δ} \) mutant restored with a wild-type copy of \( \text{TPK1} \) also reduced the already low levels of pseudohyphae.

A careful observation of cell population revealed considerable differences between the \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \) and \( \text{tpk1}^\text{Δ} \Delta \text{tpk1}^\text{Δ} \) strains. As can be seen in Figure 4A, the pseudohyphae were shorter and exhibited more lateral buds and branches in those strains devoid of \( \text{TPK2} \) alleles. Also, in these strains the number of constrictions per filament increased, while compartment length was diminished (Figure 4B). In contrast, strains harbouring \( \text{TPK2} \), including complemented \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \Delta \text{TPK2} \) strain, showed extended filaments with fewer constrictions. As expected, DAPI staining of pseudohyphae revealed one nucleus per compartment (Figure 4C). The above data showed that under hypha-only inducing conditions low Bcy1p levels promoted the development of pseudohyphae; while absence of Tpk2p impaired pseudohyphae elongation.

### Pseudohypha elongation depends on the expression of Tpk2p isoform

In view of the results described above, morphological differences between strains of the whole set were assessed under pseudohyphae-only inducing conditions (YPD medium at 35°C). Cell morphology was monitored after 2 h. As can be seen in Figure 5A, all strains were able to develop pseudohyphal growth. However, striking differences in cell morphology could be observed. Thus, all strains lacking \( \text{TPK2} \) alleles formed chains of budding yeast cells that remain attached to each other (see strains \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \), \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \text{TPK1}/\text{tpk1}^\text{Δ} \), \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \text{BCY1}/\text{bcy1}^\text{Δ} \) and \( \text{tpk2}^\text{Δ} \Delta \text{tpk2}^\text{Δ} \text{bcy1}^\text{Δ}/\text{bcy1}^\text{Δ} \)). The \( \text{TPK2}/\text{tpk2}^\text{Δ} \)
strain showed a mixture of long and short pseudohyphae. Complementation of the \( \text{tpk2}^{\Delta} / \text{tpk2}^{\Delta} \) mutant with a wild-type copy of \( \text{TPK2} \) restored the elongation capability of pseudohyphae, confirming that Tpk2p was required for elongation of pseudohyphae. In contrast, wild-type, homozygous and heterozygous \( \text{tpk1}^{\Delta} / \text{tpk1}^{\Delta} \) mutants developed pseudomycelia formed by elongated filaments with fewer compartments between constrictions. The average compartment lengths of the \( \text{tpk2}^{\Delta} / \text{tpk2}^{\Delta} \) mutants were approximately 50% shorter than \( \text{tpk1}^{\Delta} / \text{tpk1}^{\Delta} \) and wild-type (Figure 5B).

A study by Crampin et al. (2005) revealed that different molecular mechanisms drive polarized
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Figure 2. Effect of heat shock on growth of tpk1 Δ/tpk1 Δ BCY1/bcy1 Δ mutant. Yeast cells from wild-type RGI4, BCY1/bcy1 Δ (RG65), tpk1 Δ/tpk1 Δ (RS1u), tpk1 Δ/tpk1 Δ BCY1/bcy1 Δ (RG12.1u), tpk2 Δ/tpk2 Δ (RS2u) and tpk2 Δ/tpk2 Δ BCY1/bcy1 Δ (BBA1u) mutants from the stationary stage of growth were streaked out on YPD plates. The control plate was incubated at 30 °C (left panel); the heat-shocked plate was held at 50 °C for 2 h and then shifted to 30 °C (right panel). Growth was analysed after 2 days.

Figure 3. Hyphae and pseudohyphae formation from strains RGI4, BCY1/bcy1 Δ (RG65), tpk1 Δ/tpk1 Δ (RS1u), tpk1 Δ/tpk1 Δ BCY1/bcy1 Δ (RG12.1u), tpk1 Δ/tpk1 Δ::TPK1 (HPY321), tpk2 Δ/tpk2 Δ (RS2u), tpk2 Δ/tpk2 Δ BCY1/bcy1 Δ (BBA1u) and tpk2 Δ/tpk2 Δ::TPK2 (HPY421) under hypha-only inducing conditions. Strains were grown in YPD at 30 °C overnight and about 1 × 10^7 cells were transferred to liquid YPD plus 10% FBS. Cells with hyphae (grey bars) or pseudohyphae (empty bars) were scored as a percentage of the total germinated cells (± SD from three independent experiments). At 2 h post induction 80–90% of the cells had germinated.

Our results indicated that Tpk2p but not Tpk1p played an important role in pseudohypha elongation.

TPK2 deletion results in a disorganized and weakened biofilm structure

It was reported by Bockmühl et al. (2001) that tpk2 Δ/tpk2 Δ but not tpk1 Δ/tpk1 Δ mutants were deficient in invading agar. Also, Park et al. (2005) described the reduced capacity of tpk2 Δ/tpk2 Δ mutant to invade and damage oral epithelial cells. Furthermore, several reports have shown that filamentation is not strictly required for biofilm formation, but is definitively involved in the complex architecture of the biofilm (Baillie and Douglas, 1999). In view of these facts, we hypothesized that strains lacking the Tpk2p isoform would show an altered biofilm development. Therefore, we quantified biofilm production of tpk1 Δ/tpk1 Δ and tpk2 Δ/tpk2 Δ strains carrying one or both BCY1 alleles by the violet crystal assay (Figure 6A, grey bars) and using the XTT reduction method (empty growth in hyphae as compared to pseudohyphae.)
Figure 4. Pseudohyphae characterization of wild-type and PKA mutant strains under hypha-only inducing conditions. RG4, BCY1/bcy1Δ (RG65), tpk1Δ/tpk1Δ (RS1u), tpk1Δ/tpk1Δ BCY1/bcy1Δ (RG12.1u), tpk2Δ/tpk2Δ (RS2u), tpk2Δ/tpk2Δ BCY1/bcy1Δ (BBA1u) and tpk2Δ/tpk2Δ::TPK2 (HPY421) strains were grown in YPD at 30°C overnight and about 1 × 10⁷ cells were transferred to liquid YPD plus 10% FBS. (A) Two hours after induction, pseudohyphae were stained with CFW; scale bar = 10 µm. (B) Measurement of pseudohyphae length, number of constrictions per filament and compartment length (µm). ∗p < 0.01. Error bars represent standard error (SE). (C) DAPI staining of RG4, tpk1Δ/tpk1Δ (RS1u) and tpk2Δ/tpk2Δ (RS2u) pseudohyphae; scale bar = 5 µm.
Figure 5. Pseudohyphal germinative behaviour of wild-type and PKA mutant strains. (A) Stationary-phase yeast cells from wild-type RGI4 and mutants BCY1/bcy1Δ (RG65), TPK1/tpk1Δ (R1U1.1), TPK1/tpk1Δ BCY1/bcy1Δ (RG11.1), tpk1Δ/tpk1Δ (RS1u), tpk1Δ/tpk1Δ BCY1/bcy1Δ (RG12.1u), tpk1Δ/tpk1Δ::TPK1 (HPY321), TPK2/tpk2Δ (R2U2.1), tpk2Δ/tpk2Δ (RS2u), tpk2Δ/tpk2Δ TPK1/tpk1Δ (RS11u), tpk2Δ/tpk2Δ BCY1/bcy1Δ (BBA1u), tpk2Δ/tpk2Δ bcy1Δ/bcy1Δ (EC1u) and tpk2Δ/tpk2Δ::TPK2 (HPY421) were induced to germinate for 2 h at 35°C in fresh YPD medium; scale bar = 10 μm. Insets on the corners show germinated cells stained with CFW. (B) Measurement of pseudohyphae length, number of constrictions per filament and compartment length (μm). *p ≤ 0.01. Error bars represent SE.
Figure 6. Biofilm formation and metabolic activity in BCY1/bcy1Δ (RG65), tpk1Δ/tpk1Δ (RS1u), tpk1Δ/tpk1Δ BCY1/bcy1Δ (RG12.1u), tpk1Δ/tpk1Δ::TPK1 (HPY321), tpk2Δ/tpk2Δ (RS2u), tpk2Δ/tpk2Δ BCY1/bcy1Δ (BBA1u) and tpk2Δ/tpk2Δ::TPK2 (HPY421) in comparison to wild-type RGI4. (A) Biofilm production in 96-well plates was quantified using crystal violet staining (grey bars). Biofilm metabolic activity was quantified using the XTT reduction assay (empty bars). The mean ± SD was determined from three independent experiments, measuring 10 replicates for each strain. (B) Biofilm cell morphology was observed at ×40. Pseudohypha constrictions are indicated by black arrowheads. (C) CSLM of biofilm formed on silicone squares stained with concanavalin A, observed at ×100 water immersion; scale bar = 50 µm.

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bars) to quantify metabolic activity. Growth of the \( \text{tpk2} \Delta / \text{tpk2} \Delta \) strain resulted in a weakened structure that was obvious immediately when viewing the biofilms in the 96-well plate in which they were grown. The mature biofilm was dislodged easily from the bottom of the well and readily crumbled into many pieces when the 96-well plate was moved from the incubator. The 48 h biofilm mass from \( \text{tpk2} \Delta / \text{tpk2} \Delta \) and \( \text{tpk2} \Delta / \text{tpk2} \Delta \) was significantly less than that of the rest of the strains.

As judged by light microscopy (Figure 6B), the three morphologies were present in biofilms from all strains tested. However, strains \( \text{tpk2} \Delta / \text{tpk2} \Delta \) and \( \text{tpk2} \Delta / \text{tpk2} \Delta \) failed to form highly extended filaments, while strains \( \text{BCY1/bcy1} \Delta \), \( \text{tpk1} \Delta / \text{tpk1} \Delta \), \( \text{tpk1} \Delta / \text{tpk1} \Delta \) and \( \text{tpk1} \Delta / \text{tpk1} \Delta / \text{TPK1} \) and wild-type strain were able to form mycelia composed of long tubular cells. Strains lacking Tpk2p isoform showed parallel filaments in the mutant biofilm that contrasted in appearance with the intertwining hyphae and pseudohyphae present in the \( \text{tpk1} \Delta / \text{tpk1} \Delta \) and in RGI4 biofilm structure. Reintegration of the \( \text{TPK2} \) wild-type allele in strain \( \text{tpk2} \Delta / \text{tpk2} \Delta \) restored the production of long filaments similar to those of the reference wild-type strain.

The results described above prompted us to assess the ability of these strains to adhere to an abiotic support, such as squares of silicone catheter material. As can be seen in Figure 6C, images from confocal scanning laser microscopy (CSLM) inspection of silicone slices revealed that the biofilm morphology of \( \text{tpk1} \Delta / \text{tpk1} \Delta \) and RGI4 strain was similar, composed of a dense population of predominantly mature filamentous forms. In contrast, in a background of catalytic activity provided exclusively by Tpk1p, the biofilm formed was markedly less dense, composed of short pseudohyphae, few true hyphae and yeast cells (see mutant \( \text{tpk2} \Delta / \text{tpk2} \Delta \)). Reconstitution of the \( \text{TPK2} \) wild-type allele in strain \( \text{tpk2} \Delta / \text{tpk2} \Delta \) restored biofilm architecture and adhesion ability. The fragile biofilm formed by the \( \text{tpk2} \Delta / \text{tpk2} \Delta \) strain could be the consequence of the morphological abnormalities observed under conditions promoting hyphae and pseudohyphae.

The filamentation and adherence defects for \( \text{tpk2} \Delta / \text{tpk2} \Delta \) strain are in agreement with the reduction in virulence of \( \text{tpk2} \) null strain in a mouse intravenous and mouse oropharyngeal infection model \textit{in vitro} (Park \textit{et al.}, 2005). These findings suggested a significant defect in cell adherence of \( \text{tpk2} \Delta / \text{tpk2} \Delta \) strains, very probably due to a different expression pattern of genes involved in cell–cell interaction, such as adhesins, and/or cell to surface adhesion.

Adhesins Als1p and Als3p are poorly expressed in a \( \text{tpk2} \)-null background

In \textit{S. cerevisiae}, filamentation and cell aggregation are both subjected to regulatory pathways that converge on Flo11p, an effector of flocculation. Different pathways are known to regulate the expression of \textsc{flo} genes in yeast, among them the Ras1-cAMP signalling cascade (Verstrepen and Klis, 2006). Downstream, Tpk2p activates Flo8p and represses Sfl1p, which encodes a suppressor of flocculation (Conlan and Tzamarias, 2001). Sfl1p is regulated by Tpk2p phosphorylation (Robertson and Fink, 1998).

In \textit{C. albicans}, \textsc{als} genes encode a family of cell-surface glycoproteins, some of which control adhesion to host surfaces (reviewed in Hoyer \textit{et al.}, 2008). It has been shown that the expression of Als1p and Als3p, both effectors of filamentation, function redundantly to promote biofilm formation (Nobile \textit{et al.}, 2008) and are also controlled by the transcription factor Efg1p (Fu \textit{et al.}, 2002; Zhao \textit{et al.}, 2004), which in turn is believed to be regulated by Tpk2p (Lengeler \textit{et al.}, 2000; Sonneborn \textit{et al.}, 2000).

Since adhesion to surfaces and cell–cell interaction are closely related, it was of interest to assess the ability of Tpk1p and Tpk2p isoforms to mediate flocculation (see Figure 7). When grown as blastospores in SD minimal liquid medium, the decreased degree of clumping in the \( \text{tpk2} \Delta / \text{tpk2} \Delta \) mutant was dramatic (Figure 7A). Under the hypha-inducing condition, wild-type \textsc{roi4}, \( \text{tpk1} \Delta / \text{tpk1} \Delta \) and reintegrated \( \text{tpk2} \Delta / \text{tpk2} \Delta / \text{TPK2} \) strain flocculated extensively, forming large aggregates of cells that rapidly sedimented to the bottom of the tube; while the \( \text{tpk2} \Delta / \text{tpk2} \Delta \) mutant flocculated minimally under this condition (Figure 7A, B).

In view of the described defect in flocculation of the \( \text{tpk2} \) null strain, we investigated the cell surface localization of Als1p and Als3p adhesins, using indirect immunofluorescence with anti-Als1p and anti-Als3p monoclonal antibodies (see Figure 8).
As can be seen, a faint staining for both proteins was detected on the surface of \textit{tpk2 Δ/tpk2 Δ} germ tubes after 1 h of growth in YPD plus 10% FBS. This staining was more intense on the hyphal surface of wild-type, \textit{tpk1 Δ/tpk1 Δ} and reintegrated \textit{tpk2 Δ/tpk2 Δ::TPK2} strains. In these strains occasional areas of brighter staining were observed at the hyphal tip. No immunofluorescence was observed in blastospores or in pseudohyphae from wild-type or \textit{tpk} mutants (not shown). Although the above results are only qualitative, they suggest that isoform Tpk2p is predominately controlling cell aggregation. Considering that \textit{EFG1} is regulated by Tpk2p (Lengeler et al., 2000; Sonneborn et al., 2000) and that \textit{efg1} null mutants failed to express \textit{ALS1} (Fu et al., 2002), it is reasonable to assume that in \textit{tpk2 Δ/tpk2 Δ} strain Efg1p is less activated and consequently Als proteins are poorly expressed. These results are in line with those of Nobile et al. (2006), who found that a null mutant of the transcription factor \textit{BCR1} exhibited a reduced expression of both \textit{ALS1} and \textit{ALS3}, which seems to be critical for its biofilm defect.

Our recent genetic and biochemical studies of PKA mutants allowed us to uncover several Tpk functional specificities. Thus, in a previous report we showed that Tpk1p and Tpk2p, despite a high level of amino acid sequence identity and overlapping roles in viability, are not redundant in stress and glycogen metabolism (Giacometti et al., 2009). The results presented here further indicate that Tpk2p plays a major role in regulating pseudohyphal filament extension, biofilm formation and cell adherence.

A more detailed understanding of these regulatory mechanisms will no doubt allow us to gain broader insight into the relationship between fungal morphology and virulence.
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Figure 8. C. albicans cells from strains RGI4, tpk1Δ/tpk1Δ (RS1u), tpk2Δ/tpk2Δ (RS2u) and tpk2Δ/tpk2Δ::TPK2 (HPY421) labelled with either anti-Als1 or anti-Als3 MAb and Alexa Fluor 488 conjugated secondary antibody. (B) Als1p detection through CSLM of the surface of germ tubes grown in YPD + 10% FBS for 1 h. Als3p detection is shown in (D). (A, C) The same images as (B) and (D), respectively, illuminated with white light only. Scale bar = 10 µm

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