RAS/Cyclic AMP and Transcription Factor Msn2 Regulate Mating and Mating-Type Switching in the Yeast *Kluyveromyces lactis*\textsuperscript{v}

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In response to harsh environmental conditions, ascomycetes produce stress-resistant spores to promote survival. As sporulation requires a diploid DNA content, species with a haploid lifestyle, such as *Kluyveromyces lactis*, first induce mating in response to stress. In *K. lactis*, mating and mating-type switching are induced by the DNA-binding protein Mts1. Mts1 expression is known to be upregulated by nutrient limitation, but the mechanism is unknown. We show that a ras2 mutation results in a hyperswitching phenotype. In contrast, strains lacking the phosphodiesterase Pde2 had lower switching rates compared to that of the wild type (WT). As Ras2 promotes cyclic AMP (cAMP) production and Pde2 degrades cAMP, these data suggest that low cAMP levels induce switching. Because the MTS1 regulatory region contains several Msn2 binding sites and Msn2 is a transcription factor that is activated by low cAMP levels, we investigated if Msn2 regulates MTS1 transcription. Consistently with this idea, an msn2 mutant strain displayed lower switching rates than the WT strain. The transcription of MTS1 is highly induced in the ras2 mutant strain. In contrast, an msn2 ras2 double mutant strain displays WT levels of the *MTS1* transcript, showing that Msn2 is a critical inducer of *MTS1* transcription. Strains lacking Msn2 and Pde2 also exhibit mating defects that can be complemented by the ectopic expression of Mts1. Finally, we show that *MTS1* is subjected to negative autoregulation, presumably adding robustness to the mating and switching responses. We suggest a model in which Ras2/cAMP/Msn2 mediates the stress-induced mating and mating-type switching responses in *K. lactis*.

The three cell types in *S. cerevisiae* are the a- and α-haploid cell types and the a/α-diploid cell type. Diploids arise from a mating reaction, i.e., the cellular and nuclear fusion of haploids with opposite mating types (28). A single locus (*MAT*) determines the mating type, which encodes either the a (*MATα*) or α gene (*MATa*). Each haploid genome also contains two additional loci that encode mating-type information, and they are known as the cryptic mating-type loci (*HMLα* and *HMRa*) because they are not expressed. Mating-type switching in *S. cerevisiae* is induced by an endonuclease called HO (homothallicism), which cuts the *MAT* locus in a position that shares homology with the cryptic mating-type loci (29, 38). A gene conversion then exchanges the genes present in the *MAT* locus for the genes present in one of the cryptic mating-type loci.

In *S. cerevisiae*, mating-type switching is a thoroughly characterized process which has contributed significantly to basic knowledge of gene conversion (26) and transcriptional regulation (17, 37). Mating-type switching in *S. cerevisiae* is regulated by controlling the transcription of the *HO* gene. *HO* transcription is repressed in *MATa/MATα* diploids (by the a1/α2 repressor) and induced in the G1 phase of the cell cycle (by SBF) (2). In addition, a lineage-specific regulation was described in which only mother cells express HO, since daughter cells contain a repressor of *HO* transcription (Ash1) (16). These regulatory mechanisms lead to a switching rate of ~0.5/generation in haploid *S. cerevisiae* strains, resulting in a rapid return to the diplophase. Consistently, *S. cerevisiae* strains isolated from natural sources are diploid.

The function of mating-type switching in *S. cerevisiae* is to facilitate a rapid return to the diploid state. An obvious advantage of being diploid is that survival in harsh environmental conditions is enhanced through the formation of spores. Spores form after meiosis, and this developmentally distinct form of yeasts is resistant to many types of stresses, including heat, drought, and toxic chemicals (15). However, other ascomycetes grow predominantly as haploids in nature. One example of this is the distantly related ascomycetes *Kluyveromyces lactis* (42). Hence, in yeasts where the haplophase dominates there are additional mechanisms that ensure survival in a harsh environment.

Mating-type switching also has been observed in *K. lactis* (6, 27). *K. lactis* relies on gene conversion and cryptic mating-type loci for completing switching, but the inducer of switching is different from that of *S. cerevisiae*. In *K. lactis*, mating-type switching is induced by a DNA-binding protein called Mts1 (mating-type switch 1). *MTS1* is orthologous to the *S. cerevisiae* RME1 gene, which has a role in repressing meiosis-specific genes during vegetative growth. Mts1 binds to several sites in both the *MATα* and *MATa* loci. Mts1 itself shares no homology with DNA nucleases but appears to stimulate the excision of a mobile DNA element present in the *MATα* locus. The current model posits that the mobilization of the *MATα* mutator-like element (MULE) induces switching. Consistently with this idea, the transposase-like protein encoded by the α3 MULE is essential for switching from *MATα* to *MATa*. Moreover, the mutational inactivation of the catalytic DDE motif of α3 abolishes switching. The mechanism underlying *MATα* to *MATα* switching remains unknown, but it probably involves a hairpin-
generating nuclease. Hairpin intermediates can be observed in the MATa locus in genetic backgrounds that are compromised for hairpin resolution.

Switching rates in K. lactis have never been accurately determined, but it is clear that the rates are lower than those in S. cerevisiae (27). The regulation of switching also seems different between yeasts, as switching in K. lactis is induced in nutrient-limited conditions. Interestingly, MTS1 transcription is induced in nutrient-limited conditions as well (6, 9), leading to a model in which switching is controlled through the regulation of MTS1 expression. Apart from stimulating switching, Mts1 also induces mating through the direct activation of genes essential for mating (9). Hence, nutrient limitation stimulates both mating and switching. Mating-type switching followed by mating can then promote spore formation upon nutrient limitation, albeit with some delay compared to that of diploid S. cerevisiae.

Stress-induced gene transcription in S. cerevisiae is partly controlled by a set of redundant zinc finger-containing transcription factors called Msn2 and Msn4 (34, 44). These factors bind the stress response element (STRE) AGT, which is present in the regulatory regions of many stress-induced genes (11). In S. cerevisiae, large-scale analyses showed that Msn2/4 activate about 200 genes (22). The activation of several genes promotes a phenomenon known as cross-protection, i.e., that transient exposure to one type of stress will provide protection from future exposure to different stress conditions. There is also evidence from other ascomycetes that orthologs of Msn2/4 is induced in nutrient-limited conditions as well (6, 9), leading to a model in which switching is controlled through the regulation of MTS1 expression. Apart from stimulating switching, Mts1 also induces mating through the direct activation of genes essential for mating (9). Hence, nutrient limitation stimulates both mating and switching. Mating-type switching followed by mating can then promote spore formation upon nutrient limitation, albeit with some delay compared to that of diploid S. cerevisiae.

In this study, we demonstrate that mating-type switching in K. lactis is regulated by the Ras/cAMP pathway and Msn2. This regulation is mediated by controlling the transcription of the MTS1 gene. The transcription of MTS1 links mating-type switching with mating, as both processes are directly induced by Mts1. We propose that this regulation of mating-type switching and mating serves as an adaptation to the mostly haploid lifestyle of K. lactis.

MATERIALS AND METHODS

Yeast strains. The strains used in this study are listed in Table 1. Unless noted otherwise, all gene deletions/insertions were generated using a one-step gene disruption/methodology (32) with a kanMX, NAT, or URA3 PCR fragment amplified from pFA6a-KanMX (5), pAG25 (23), or pRS306. Genetic manipulations were confirmed by DNA blots or locus-specific PCR. Sequences of oligonucleotides used are available on request. SAY724 (MATa hmla1::kanMX nej1::LEU2) was generated through a cross between SAY130 (MATa hmla1::kanMX) and SAY509 (MATa nej1::LEU2). SAY975 (meta2::NAT hmla1::kanMX nej1::LEU2) was generated in SAY724. SAY990 (meta2::NAT hmla1::kanMX was obtained by crossing SAY724 with SAY119. The strains SAY1131 (pde2::kanMX), SAY1379 (pat::3::LEU2) and SAY1554 (meta2::kanMX mts1::LEU2) were generated in SAY1539, whereas SAY1069 (meta2::NAT hmla1::kanMX leu2::SPT8) was generated in SAY1069. SAY1089 (pat::3::LEU2) was generated by amplifying a MTS1-containing PCR fragment with 50 bp of flanking DNA corresponding to the MTS1 5’ end and introducing it into SAY572. SAY1069 (pat::3::LEU2; ob- tained by selection) was generated by the random transformation of linearized pMB18 into SAY990. The genetic selection is described in more detail below.

S. cerevisiae TEF2. Cloning was performed using standard methods (41). Plasmids pPMB35 (6) and pCJ118 (13) are described elsewhere. pPMB18 (pRS405-ADH1-pTEF2) was generated in two steps. First, AKP13 was generated by a three-factor cloning using an XbaI-BamHI PCR fragment containing the S. cerevisiae ADH1 promoter (bp –492 to –13 relative to the start codon), an XhoI-BamHI PCR fragment containing the S. cerevisiae TEF2 promoter (bp –1000 to –10 relative to the start codon), and a Ssph-I-Xhol-digested pRS405 (6). Second, the ADH1-TEF2 promoters were released from pRS405 using a NotI-Xhol digest and ligated into NotI-Xhol-cut pRS405 (LEU2), generating pPMB18.

Media and standard methods. Protocols for DNA and Western blotting, DNA/RNA preparations, and the transformation of yeast and bacteria, as well as the composition of growth media for yeast and bacteria, were published previously (4, 41, 43). For protein blasts, the primary anti-TAP antiserum (Open Biosystems) and the anti-Pgk1 antiserum (a gift from Per Ljungdahl) both were diluted 1:1,000. The PCR-based determination of the MAT genotype was described before (6).

RT-qPCR. Reverse transcriptase quantitative PCR (RT-qPCR) procedures and primers used for amplifying the ACT1 control cdNA were described before (7). The primers used for amplifying the MTS1 cDNA were 5’ TCCACAAAA ACCAAAAGGC and 5’ TTCTTTGGCAACCGGAGGTCC.
Analysis of chromatin immunoprecipitation (ChIP) results. Primary data published previously (9) were used in combination with the MochiView software (30) to scan the *K. lactis* genome for Mts1-Myc binding sites.

**Genetic selection.** pPMB18 (330 μg) was linearized with BamHI and introduced into SAY990. Eleven independent transformations with 30 μg of linearized plasmid resulted in approximately 1.3 × 10^7 colonies on SC-Leu plates. The colonies were replica printed to SC-Leu plates containing 150 μg/ml G418 (designated SC-Leu + 150 μg/ml G418). Colonies growing on SC-Leu + 150 μg/ml G418 plates were transferred to fresh plates (containing SC-Leu + 150 μg/ml G418) and then patched on SC-Leu and replica printed on a yeast extract-peptone-dextrose (YPD) plate containing a lawn of the test mating proficient strain SAY44 (MATα). The mating plates were further replica printed on SD+Ura to score mating-proficient colonies. Inverse PCR (32) followed by DNA sequencing were performed to identify the genomic position of pPMB18 in the mating-proficient strains. The pPMB18 insertion in the *RAS2* gene (C13387g) corresponded to amino acid 73 (out of 284).

**RESULTS**

*K. lactis* switches mating type with a rate of 6 × 10^-4 events/generation in rich medium. *K. lactis* strains grown under laboratory conditions exhibit stable mating types and hence switch mating type infrequently (3). Previously, we have observed low-level switching when strains were grown in nutrient-limited conditions (6), but we lacked a reliable assay for measuring switching rates. Measuring switching rates using quantitative PCR is unfeasible, as *MAT*, *HMLα*, and *HMRα* all are flanked by identical repetitive sequences (L and R), such that long amplicons would be necessary to distinguish between *MATα* and *MATα*. To establish a non-PCR-based assay measuring switching rates, we generated a strain that contained an inser-

tion of the *URA3* gene downstream of the *MATα* locus (Fig. 1A). The *URA3* gene was inserted downstream of the *MATα* gene without disrupting the open reading frame (ORF). We predicted that mating-type switching would lead to the elimination of the *URA3* gene. Because *ura3* strains are resistant to 5-fluoro-
orotic acid (5-FOA), mating-type switching rates could be determined by measuring the fraction of cells that became 5-FOA^-.

As a control, we also generated a strain with an insertion of the *URA3* gene downstream of the *HMRα* gene. *HMRα* acts as a donor during switching rather than as a recipient, the *HMRα*:URA3 locus should be lost at a very low rate. The strains were inoculated into rich medium from a plate selecting for the insertion (SC-Ura), grown for six to eight generations, and then plated on 5-FOA and YEPD plates. The average 5-FOA/YEPD ratio was 4.6 × 10^-3 (standard errors from the means [SEM], 1.2 × 10^-3) for the *MATα*:URA3 strain in 11 independent experiments (Fig. 1B). For the *HMRα*:URA3 strain the average 5-FOA/YEPD ratio was only 3.0 × 10^-6, which is 1,000-fold lower than that for the *MATα*:URA3 strain. We tested the mating type of eight 5-FOA-resistant segregants obtained from the *MATα*:URA3 strain by *MAT*-specific PCR and found that they had the *MATα* genotype. Hence, the 5-FOA^- strains had switched their mating type and the *MATα*:URA3 strain preferentially used the *HMLα* locus as the donor.

If this assay accurately measured switching, then it should be possible to block switching using the appropriate mutations. To test this idea, we determined switching in a *MATα*:URA3 *mts1Δ* mutant strain. Since Ms1 is critical for the induction of switching, we expected this strain to have a drastically reduced switching rate. In the *mts1Δ* mutant strain, the average 5-FOA/ YEPD ratio was 9.7 × 10^-6, which is comparable to the ratio in the *HMRα*:URA3 strain. We conclude that the assay accurately measured switching frequencies, with a mating-type switching frequency of cells grown in rich medium of approximately 6 × 10^-4 events/generation.

**FIG. 1.** Genetic assay for measuring switching rates. (A) Schematic of the test strain used for measuring switching rates. The *S. cerevisiae* *URA3* gene was inserted downstream of the *MATα* gene. Upon mating-type switching, the *MATα*:URA3 locus is replaced by the sequences present at *HMLα* (most frequently) or *HMRα*, resulting in 5-FOA resistance. (B) The ratios of 5-FOA/YEPD plating efficiencies were determined in strains SAY1488 (*MATα*:URA3), SAY1554 (*MATα*:URA3 *mts1Δ*:LEU2), and SAY1489 (*HMRα*:URA3). Error bars represent the standard errors from the means calculated from 11 (SAY1488), 3 (SAY1554), and 3 (SAY1489) independent experiments.

**Identification of Ras2 as a regulator of mating-type switching.** To identify genes that regulate switching in *K. lactis*, we performed a genetic selection for mutations that increased switching rates. A previous selection (6) showed that a physical translocation of a *kanMX* gene, encoding G418 resistance, from a silent locus into the expressed *MAT* locus would lead to its expression. We used a strain in which the *HMLα* gene was exchanged for the *kanMX* gene. In addition, the strain contained a deletion of the *MATα* gene, and since the *MATα* gene is necessary for mating, the resulting strain was sterile (Fig. 2A). In *K. lactis*, illegitimate recombination frequencies are high, making it possible to use exogenously added DNA as a mutagen. Hence, the *mutα2Δ::MAT hmlα::kanMX* strain was mutagenized using the random insertion of a plasmid containing a LEU2 selectable marker into the genome. We next selected for G418-resistant colonies and tested if such isolates could mate with a *MATα* tester strain. The goal of this procedure was to obtain mutations that resulted in constitutive switching, first switching to *mutα1Δ::kanMX*, becoming G418 resistant, and then switching to *MATα*, becoming fertile. Among the isolated mutants was a strain that showed poor
growth, and DNA blotting using a MAT-specific probe revealed that the strain formed a mixture of MATa and mataΔ::kanMX genotypes (Fig. 2B). An inverse PCR procedure showed that the plasmid insertion was in the 5’ end of the RAS2 gene (C13387g), resulting in a disruption of the RAS2 ORF. RAS2 encodes a small GTPase with a central role in growth regulation in all eukaryotes. In fungi, Ras proteins are orthologs, and in directed experiments (data not shown) we postulated that the Ras2 activity will lower cAMP levels and result in decreased transcription and switching, and that high levels of cAMP regulate switching.

These data suggest that low levels of cAMP induce Msn2 and Pde2 regulate mating.

**PDE2 is required for normal switching rates.** Because compromised Ras2 activity will lower cAMP levels and result in hyperswitching, we tested if higher cAMP levels would inhibit switching. To obtain a strain with increased cAMP levels, we generated a strain with a deletion of the PDE2 gene (A3619g). Pde2 is a cAMP phosphodiesterase that degrades cAMP, hence limiting intracellular cAMP levels. We predicted that pde2 mutants would have lower switching rates, the phenotype opposite that of ras2.

The average 5-FOA/YEPD ratio in the pde2 strain was 1.0 × 10^{-3} (SEM, 2.3 × 10^{-4}), indicating a 5-fold reduced switching rate compared to that of the wild type (Fig. 3A). These results are consistent with a role for cAMP in the regulation of switching.

**MSN2 also is required for normal switching rates.** We attempted to identify the transcription factor that is responsible for the cAMP-dependent regulation. Two key transcription factors responding to cAMP levels in S. cerevisiae are Msn2 and Msn4. The activity of the paralogous Msn2/4 proteins is inactivated by PKA-dependent phosphorylation (18, 25), leading to reduced transcription from Msn2/4-activated genes when levels of cAMP are high. Hence, a K. lactis Msn2/4 ortholog was an excellent candidate for mediating a cAMP-dependent regulation of switching. BLAST searches of the K. lactis genome using S. cerevisiae Msn2/4 as queries identified a single Msn2/4 ortholog (F26961g) which had the highest similarity to Msn2.

We deleted the MSN2 gene and measured switching rates in the resulting strain. In the msn2 strain the 5-FOA/YEPD ratio was only 3.5 × 10^{-4} (SEM, 1.2 × 10^{-4}), demonstrating a 13-fold decreased switching rate compared to that of the WT. Hence, Msn2 regulates switching in K. lactis.

**Transcription of MTS1 is regulated by Msn2, Ras2, and Pde2.** The Mts1 protein induces mating-type switching in K. lactis by binding to sites in both the MATa and MATa loci. Since we previously determined that the transcription of MTS1 was induced in nutrient-limited conditions (6), we hypothesized that Ras2/Pde2/Msn2 exert their effects on switching through regulating the transcription of the MTS1 gene. Consistently with this idea, Msn2 binds the stress response element (STRE) in S. cerevisiae, which has the sequence AGGGG. In the 3.2-kb intergenic region upstream of the MTS1 ORF there are five consensuses STREs and six close matches to the STRE. We investigated the expression of the MTS1 gene normalized to the expression of ACT1 (Fig. 3B) by RT-qPCR. In the ras2 mutant strain, MTS1 transcription was upregulated 20-fold compared to that of the isogenic wild-type strain, explaining the hyperswitching phenotype. In contrast, MTS1 mRNA levels in the pde2Δ and msn2Δ mutant strains were reduced 5- and 3-fold, respectively, compared to that of their parental strain. To test if the increased MTS1 transcription in the ras2 mutant required Msn2, we measured MTS1 transcription in a ras2 msn2 double mutant strain. This strain expressed levels of MTS1 comparable to those of the wild type, indicating that Msn2 acts downstream of Ras2 in the regulation of switching. These data suggest that low levels of cAMP induce MTS1 transcription and switching, and that high levels of cAMP repress MTS1 transcription and switching.

**Msn2 and Pde2 regulate mating.** Previously, an expression microarray experiment comparing a wild-type strain to an mts1Δ strain revealed that 20 genes were downregulated (>2-fold) and 52 genes were upregulated (>4-fold) in the mts1Δ strain (9). Specifically, Mts1 directly activated the transcription of a group of genes required for mating, which is consistent with the observation that mts1Δ strains are sterile. Because Mts1 directly activates genes required for mating, we anticipated that the Ras2/cAMP/Msn2 pathway would regulate mat-
ing in addition to its role in mating-type switching. We tested this notion by comparing the mating efficiency of \( \text{MAT}^{a} \text{msn2} \) and \( \text{MAT}^{a} \text{pde2} \) to that of an isogenic wild-type \( \text{MAT}^{a} \) strain. The strains were mixed with a \( \text{MAT}^{a} \) tester strain on medium that only allowed the growth of diploid cells (SC-Ade-Leu). Both the \( \text{pde2} \) and \( \text{msn2} \) genes were required for efficient mating, as the formation of diploids was severely reduced in the mutant strains (Fig. 3C). We predicted that the mating defect of \( \text{msn2} \) and \( \text{pde2} \) were the result of defective Mts1 induction. To test this idea, we introduced a \( p \text{GAL-MTS1} \) plasmid into these strains, expecting that ectopic Mts1 expression would complement the mating defect. Ectopic Mts1 expression indeed improved mating in the \( \text{msn2} \) and \( \text{pde2} \) strains (Fig. 3C). Hence, Msn2/Pde2 regulates both mating and mating-type switching in \( K. \text{lactis} \).

**Autoregulation of MTS1 transcription.** Others recently published a chromatin immunoprecipitation analyzed genome-wide by hybridization to a microarray (designated ChIP-on-chip) using epitope-tagged Mts1 (9). By further examining this data set, we discovered that a prominent Mts1 binding peak was present upstream of the \( \text{MTS1} \) gene. In the left panel, mRNA was prepared from strains SAY1488 (WT), SAY1546 (\( \text{pde2} \)), and SAY1539 (\( \text{msn2} \)). In the right panel, mRNA was prepared from SAY900 (WT), SAY1069 (\( \text{ras2} \)), and SAY1664 (\( \text{msn2}\text{kanMX ras2} \)). The level of expression in the wild-type strains was set to 1.0. The data were normalized to the expression of the \( \text{ACT1} \) gene using the comparative threshold cycle (Ct) method. Error bars indicate the maximum and minimum values obtained in two separate experiments. (C) Mating tests of the \( \text{MAT}^{a} \) strains WT (SAY572), \( \text{msn2}\text{kanMX} \) (SAY1379), and \( \text{pde2}\text{NAT} \) (SAY1131) to the \( \text{MAT}^{a} \) tester strain (WM52). Diploids were selected on SC-Ade-Lys.
region and that the overexpression of Mts1 represses MTS1 transcription, we conclude that MTS1 transcription is negatively autoregulated.

**DISCUSSION**

In this study, we demonstrate that mating-type switching is regulated through the RAS/cAMP pathway in *K. lactis*. We suggest a first model (Fig. 4C) in which Ras2 senses nutrient levels, thereby regulating cAMP production. High cAMP levels repress the activity of the Msn2 transcription factor. We assume that Msn2 activity is regulated by PKA in *K. lactis* similarly to Msn2 regulation by PKA in *S. cerevisiae*. An alignment of *Saccharomyces* and *Kluyveromyces* Msn2 reveals that residues in the NLS predicted to be targets of PKA in *S. cerevisiae* are partly conserved in *K. lactis* (data not shown). However, we have not been able to demonstrate that the subcellular localization of Msn2 changes in ras2 mutants compared to that in the WT.

When Ras2 is inactive or its function is compromised, Msn2 activates the transcription of Mts1, which in turn promotes mating-type switching. Given the presence of Msn2 binding sites (STREs) in the MTS1 regulatory region, we reason that the role of Msn2 in this aspect is direct. It turns out that the epitope tagging of Msn2 renders the protein inactive (data not shown), which has prevented us from demonstrating Msn2 binding to the MTS1 promoter by ChIP. We tested another cAMP-responsive candidate protein for the regulation of MTS1 transcription, called Gis1, but the deletion of GIS1 did not affect switching rates (data not shown). A prediction from this model is that MTS1 induction should rely on Msn2. This prediction was confirmed (Fig. 3B), as the absence of Msn2 suppressed the 20-fold overexpression of MTS1 observed in the ras2 mutant background. However, the ras2 msn2 double mutant expressed more MTS1 than the msn2 single mutant did (Fig. 3B). Hence, it is possible that Ras2 also affects Mts1 expression in an Msn2-independent manner.

The intergenic region representing the upstream region of MTS1 and the downstream region of the next ORF (*PRX1*) is unusually large (3.2 kb), considering that the average intergenic region in *K. lactis* is only about 500 bp (20). We speculate that this reflects a complex regulation of MTS1 transcription, giving room for binding sites for multiple regulatory proteins. Apart from the Msn2 binding sites presumably mediating a cAMP-dependent response, there is also a binding site for the a1/a2 heterodimer (9). The a1/a2 heterodimer represses the transcription of MTS1 in diploids or haploids with defects in the silencing of the cryptic mating-type loci. In *S. cerevisiae*, several genes required for mating are directly repressed by a1/a2 and are collectively known as haploid-specific genes. However, *K. lactis* haploid-specific genes are directly induced by Mts1 and not repressed by a1/a2. As MTS1 is repressed by a1/a2, the final output in *S. cerevisiae* and *K. lactis* is the same, but during the evolution of ascomycetes a rewiring of the regulation has occurred.

We also show that Mts1 is autoregulated, mediating the
repression of its own transcript (Fig. 4). Negative autoregulation in other systems has been shown to reduce cell-to-cell fluctuations in steady-state levels of transcription factors (8) and also to speed up the response times of transcriptional networks (40). Hence, negative autoregulation by Mtsl is likely to add robustness to the mating and switching responses.

We have shown previously that Mtsl directly regulates switching through binding sites in the \textit{MATa} locus. Mtsl also plays a pivotal role in regulating mating by inducing haploid-specific genes (9). Hence, Mtsl induces both mating and switching, but presumably an individual cell must do either. We hypothesize that this choice is regulated. It seems as if switching should be the secondary choice, as switching is costly and results in an inbreeding of the resulting offspring. A nutrient limitation-induced mating reaction, on the other hand, is more direct and also opens up the possibility for outcrossing. Hence, if there are cells of the opposite mating type in the vicinity, it would make sense to mate rather than switch. An attractive possibility is that mating pheromones inhibit switching (obviously pheromones induce mating), which is an idea that will be interesting to test in the future.

It is interesting that the regulation of switching in \textit{K. lactis} is completely different from the regulation of switching in \textit{S. cerevisiae}. In homothallic \textit{S. cerevisiae} strains, switching occurs in mother cells in the G1 phase of the cell cycle, resulting in a very high switching rate. In contrast, in \textit{K. lactis} switching rates are relatively low (6 \times 10^{-4}) in rich medium (Fig. 1B) and are regulated by nutrient availability (6, 27). Because the \textit{K. lactis} genome contains an HO pseudogene in a syntenic position compared to the \textit{S. cerevisiae} genome, the common ancestor probably used an HO-mediated switching mechanism. After their separation, \textit{K. lactis} evolved a novel switching mechanism that acquired a different type of regulation. We argue that the differential regulation of switching in \textit{K. lactis} has evolved as a consequence of the haplophase being the predominant life cycle. Natural isolates of \textit{K. lactis} strains are haploid, in contrast to \textit{S. cerevisiae}, where natural isolates are diploid. The diplophase is unstable in \textit{K. lactis}, probably explaining the inclination for a haploid DNA content. Hence, \textit{K. lactis} growing in nature cannot immediately sporulate in response to nitrogen and carbon starvation but must mate and form diploids first. Therefore, the Ras/cAMP regulation of mating and switching in \textit{K. lactis} may promote survival during starvation by ensuring the formation of resilient spores.

Cyclic AMP signaling controls a wide range of responses in fungi (10, 19). A few examples are the pseudohyphal growth response in \textit{S. cerevisiae}, which is controlled by intracellular cAMP concentrations (33). Morphological changes in fungi such as \textit{Candida albicans} (21) and \textit{Cryptococcus neoformans} (1) are also regulated by Ras/cAMP, and these morphological switches are critical for pathogenicity. Hence, our data add to an extensive list of responses to cAMP signaling in fungi, but to our knowledge this work is the first link between Ras/cAMP and mating-type switching.

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REFERENCES

1. Alspaugh, J. A., L. M. Cavallo, J. R. Perfect, and J. Heitman. 2000. \textit{RAS1} mediates filamentation, mating and growth at high temperature of \textit{Cryptococcus neoformans}. Mol. Microbiol. 36:352–365.
2. Andrews, B. J., and I. Herskowitz. 1989. Identification of a DNA binding factor involved in cell-cycle control of the yeast HO gene. Cell 57:21–29.
3. Andrews, B. J., A. Kep, J. O. Sjöstrand, and J. R. Pringle. 2000. \textit{Kluyveromyces lactis} Sir2p regulates cation sensitivity and maintains a specialized chromatin structure at the cryptic alpha-locus. Genetics 156:91–91.
4. Aubele, F. M. 1999. Short protocols in molecular biology: a compendium of methods from current protocols in molecular biology. Wiley, New York, NY.
5. Bähler, J., et al. 1998. Heterologous modules for efficient and versatile PCR-based gene targeting in \textit{Schizosaccharomyces pombe}. Yeast 14:943–951.
6. Barseoum, E., P. Martinez, and S. U. Åström. 2010. Alpha3, a transposable yeast cell to environmental changes. Mol. Biol. Cell 21:4241–4257.
7. Barseoum, J. O. Sjöstrand, and S. U. Åström. 2010. Ume3 is required for the \textit{MATa/MATa}bpha cellular identity and transcriptional silencing in \textit{Kluyveromyces lactis}. Genetics 184:999–1011.
8. Becskei, A., and L. Serrano. 2000. Engineering stability in gene networks by low-amplification. Nature 405:590–593.
9. Booth, L. N., B. B. Tuch, and A. D. Johnson. 2010. Intercalation of a new tier of transcription regulation into an ancient circuit. Nature 468:959–963.
10. Borst-Walmesly, M. L., and A. R. Walmesly. 2000. cAMP signalling in pathogenic fungi: control of dimorphic switching and pathogenicity. Trends Microbiol. 8:133–141.
11. Boy-Marcotte, E., M. Perrot, F. Bussereau, H. Boucherie, and M. Jacquet. 1998. Msn2p and Msn4p control a large number of genes induced at the diauxic transition which are repressed by cyclic AMP in \textit{Saccharomyces cerevisiae}. J. Bacteriol. 180:1044–1052.
12. Brock, D., et al. 1985. Differential activation of yeast adenylate cyclase by wild-type and mutant RAS proteins. Cell 41:763–769.
13. Chen, X. J. 1996. Low- and high-copy-number shuttle vectors for replication in the budding yeast \textit{Kluyveromyces lactis}. Gene 172:131–136.
14. Chen, X. J., and G. D. Clark-Walker. 1994. sir2 mutants of \textit{Kluyveromyces lactis} hypersensitive to DNA-targeting drugs. Mol. Cell. Biol. 14:4501–4510.
15. Coluccio, A. E., R. K. Rodriguez, M. J. Kerman, and A. M. Neiman. 2008. The yeast spore wall enables spores to survive passage through the digestive tract of \textit{Drosophila}. PLoS One 3:e2873.
16. Cosma, M. P. 2004. Development-specific repression of \textit{Saccharomyces cerevisiae} HO: Ash1 is the commander. EMBO Rep. 5:953–957.
17. Cosma, M. P., T. Tanaka, and K. Nasmyth. 1999. Ordered recruitment of transcription and chromatin remodeling factors to a cell cycle- and developmentally regulated promoter. Cell 97:299–311.
18. De Vere, V., W. Reiter, A. Ballarini, G. Ammerer, and C. Brocard. 2005. A dual role for PPI1 in shaping the Msn2p-dependent transcriptional response to glucose starvation. EMBO J. 24:4115–4123.
19. D’Andrea, A. L., and J. H. McCusker. 2005. Engineering stability in gene networks by intercalation of a new tier of transcription regulation into an ancient circuit. Nature 468:959–963.
20. Dujon, B., et al. 2004. Genome evolution in yeasts. Nature 430:35–44.
21. Feng, Q., E. Summers, B. B. Tuch, and G. Fink. 1999. Ras signaling is required for serum-induced hyphal differentiation in \textit{Candida albicans}. J. Bacteriol. 181:6339–6346.
22. Gasch, A. P., et al. 2000. Genomic expression programs in the response of \textit{Saccharomyces cerevisiae} to environmental changes. Mol. Biol. Cell 11:4241–4257.
23. Goldstein, A. L., and J. H. McCusker. 1999. Three novel dominant drug resistance cassettes for gene disruption in \textit{Saccharomyces cerevisiae}. Yeast 15:1531–1553.
24. Görner, W., et al. 1999. Nuclear localization of the C2H2 zinc finger protein Msn2p is regulated by stress and protein kinase A activity. Genes Dev. 12:586–597.
25. Görner, W., et al. 2002. Acute glucose starvation activates the nuclear localization signal of a stress-specific yeast transcription factor. EMBO J. 21:135–144.
26. Haber, J. E. 1998. Mating-type gene switching in \textit{Saccharomyces cerevisiae}. Annu. Rev. Genet. 32:561–599.
27. Herman, A., and H. Roman. 1966. Allele specific determinants of homothallicism in \textit{Saccharomyces lactis}. Genetics 53:727–740.
28. Herskowitz, I. 1989. Life cycle of the budding yeast \textit{Saccharomyces cerevisiae}. Microbiol. Rev. 52:536–553.
29. Herskowitz, I., J. Rine, and J. N. Strathern. 1992. Mating-type determination and mating-type interconversion in \textit{Saccharomyces cerevisiae}, p. 583–656. In E. W. Jones, J. R. Pringle, and J. R. Bouchard (ed.), \textit{The molecular and cellular biology of the yeast Saccharomyces, vol. 2}. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
30. Homann, O. R., and A. D. Johnson. 2010. MochiView: versatile software for ChIP-on-chip and cDNA motif analysis. BMC Biol. 8:89.
31. Jacquet, M., G. Renault, S. Lallet, J. De Mey, and A. Goldbeter. 2003. Oscillatory nucleocytoplasmic shuttling of the general stress response tran-
scriptional activators Msn2 and Msn4 in *Saccharomyces cerevisiae*. J. Cell Biol. 161:497–505.

32. **Kege, A., P. Martinez, S. D. Carter, and S. U. Åström.** 2006. Genome wide distribution of illegitimate recombination events in *Kluyveromyces lactis*. Nucleic Acids Res. 34:1633–1645.

33. **Lorenz, M. C., and J. Heitman.** 1997. Yeast pseudohyphal growth is regulated by GPA2, a G protein alpha homolog. EMBO J. 16:7008–7018.

34. **Martinez-Pastor, M. T., et al.** 1996. The *Saccharomyces cerevisiae* zinc finger proteins Msn2p and Msn4p are required for transcriptional induction through the stress response element (STRE). EMBO J. 15:2227–2235.

35. **Mayordomo, I., F. Estruch, and P. Sanz.** 2002. Convergence of the target of rapamycin and the Snf1 protein kinase pathways in the regulation of the subcellular localization of Msn2, a transcriptional activator of STRE (stress response element)-regulated genes. J. Biol. Chem. 277:35650–35656.

36. **Medvedik, O., D. W. Lamming, K. D. Kim, and D. A. Sinclair.** 2007. MSN2 and MSN4 link calorie restriction and TOR to sirtuin-mediated lifespan extension in *Saccharomyces cerevisiae*. PLoS Biol. 5:e261.

37. **Nasmyth, K.** 1993. Regulating the HO endonuclease in yeast. Curr. Opin. Genet. Dev. 3:286–294.

38. **Nickoloff, J. A., E. Y. Chen, and F. Heffron.** 1986. A 24-base-pair DNA sequence from the *MAT* locus stimulates intergenic recombination in yeast. Proc. Natl. Acad. Sci. U. S. A. 83:7831–7835.

39. **Roetzer, A., et al.** 2008. *Candida glabrata* environmental stress response involves *Saccharomyces cerevisiae* Msn2/4 orthologous transcription factors. Mol. Microbiol. 69:603–620.

40. **Rosenfeld, N., M. B. Elowitz, and U. Alon.** 2002. Negative autoregulation speeds the response times of transcription networks. J. Mol. Biol. 323:785–793.

41. **Sambrook, J., and D. W. Russell.** 2001. Molecular cloning: a laboratory manual. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.

42. **Schaffrath, R., and K. D. Breunig.** 2000. Genetics and molecular physiology of the yeast *Kluyveromyces lactis*. Fungal Genet. Biol. 30:173–190.

43. **Schiestl, R. H., and R. D. Gietz.** 1989. High efficiency transformation of intact yeast cells using single stranded nucleic acids as a carrier. Curr. Genet. 16:339–346.

44. **Schmitt, A. P., and K. McEntee.** 1996. Msn2p, a zinc finger DNA-binding protein, is the transcriptional activator of the multistress response in *Saccharomyces cerevisiae*. Proc. Natl. Acad. Sci. U. S. A. 93:5777–5782.

45. **Smith, A., M. P. Ward, and S. Garrett.** 1998. Yeast PKA represses Msn2p/Msn4p-dependent gene expression to regulate growth, stress response and glycogen accumulation. EMBO J. 17:3556–3564.

46. **Toda, T., et al.** 1985. In yeast, RAS proteins are controlling elements of adenylate cyclase. Cell 40:27–36.