# *Saccharomyces cerevisiae* Elongator mutations confer resistance to the *Kluyveromyces lactis* zymocin

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Kluyveromyces lactis killer strains secrete a zymocin complex that inhibits proliferation of sensitive veast genera including Saccharomyces cerevisiae. In search of the putative toxin target (TOT), we used mTn3:: tagging to isolate zymocin-resistant tot mutants from budding yeast. Of these we identified the TOT1, TOT2 and TOT3 genes (isoallelic with ELP1, ELP2 and ELP3, respectively) coding for the histone acetyltransferase (HAT)-associated Elongator complex of RNA polymerase II holoenzyme. Other than the typical *elp* ts-phenotype, tot phenocopies hypersensitivity towards caffeine and Calcofluor White as well as slow growth and a G<sub>1</sub> cell cycle delay. In addition, TOT4 and TOT5 (isoallelic with KTI12 and IKI1, respectively) code for components that associate with Elongator. Intriguingly, strains lacking non-Elongator HATs  $(gcn5\Delta, hat1\Delta, hpa3\Delta \text{ and } sas3\Delta)$  or non-Elongator transcription elongation factors TFIIS ( $dst1\Delta$ ) and Spt4p (spt4 $\Delta$ ) cannot confer resistance towards the K.lactis zymocin, thus providing evidence that Elongator equals TOT and that Elongator plays an important role in signalling toxicity of the K.lactis zymocin.

Keywords: Elongator/killer yeast/TOT/zymocin

# Introduction

In competing for limited resources, microorganisms have evolved sophisticated strategies to gain selective advantages over their competitors. One of these is the secretion of toxic compounds that results in killing or growth arrest of other species or genera. Well studied cases are the dsRNA-encoded viral KT28 and K1 killer toxins of Saccharomyces cerevisiae (Wickner, 1996), which cause sensitive yeast cells to irreversibly block DNA synthesis and arrest in S phase (Schmitt et al., 1996) or to destroy cytoplasmic membrane function by TOK1 hyperactivation and lethal ion channel formation (Ahmed et al., 1999). Killer dsDNA plasmid-carrying strains of the dairy yeast Kluyveromyces lactis also secrete a zymo-toxin (referred to as zymocin) that inhibits cellular growth of various sensitive yeast genera including S.cerevisiae (Stark et al., 1990; Schaffrath and Breunig, 2000). As judged from fluorescence-activated cell sorter (FACS) analyses, it predominantly causes sensitive budding yeast cells to arrest at the unbudded G<sub>1</sub> stage of the cell cycle with an unreplicated (1n) DNA content (Butler et al., 1991a). Although reminiscent of pre-START arrests induced by the pheromone cascade or displayed by  $cdc28^{ts}$  strains at restrictive growth temperatures (Leberer et al., 1997), the speculation that zymocin might act by antagonizing  $G_1$ cyclin function does not hold true: hyperactive dominant CLN3 alleles are not able to rescue sensitive cells from zvmocin treatment (Butler et al., 1994). Further evidence that the zymocin might act in late G<sub>1</sub> before START is provided by the finding that cells that have been chemically arrested in S phase by hydroxyurea, prior to zymocin treatment, are able to commit one complete round of cell division and get arrested in the new unbudded G<sub>1</sub> cell cycle stage once they have been released from the chemical S block in the continued presence of zymocin (Butler et al., 1991a).

Despite the heterotrimeric  $(\alpha\beta\gamma)$  structure of native holo-zymocin (Stark and Boyd, 1986), intracellular expression of its smallest subunit, the  $\gamma$ -toxin, alone is lethal. Thus, conditional expression of the  $\gamma$  gene from regulatable GAL promoters leads to a galactose-dependent G<sub>1</sub> arrest that mimics the effect of native holo-zymocin (Tokunaga *et al.*, 1989; Butler *et al.*, 1991b). The  $\alpha$  and  $\beta$ subunits, however, are needed for holo-zymocin to act from the cell exterior. The  $\beta$  subunit is predicted to form four C-terminal hydrophobic domains, suggesting cell membrane association, while the  $\alpha$  subunit exhibits an exo-chitinase activity that is essentially required for holozymocin function (Stark et al., 1990; Butler et al., 1991c). Sensitive yeast cells can be rescued from zymocin action by exogenously applying crude chitin preparations (D.Jablonowski and R.Schaffrath, unpublished data) and chitin-deficient S.cerevisiae mutants are resistant towards exo-zymocin (Takita and Castilho-Valavicius, 1993) but not against endogenous y-toxin expression. These observations suggest a role for the  $\alpha$  and  $\beta$  subunits in zymocin docking to sensitive cells, whereas cytotoxicity resides solely within the  $\gamma$ -toxin subunit. Based on their ability to grow in the presence of the holo-form, zymocin-resistant mutants termed skt (sensitivity to K.lactis toxin), iki (insensitive to killer) and kti (K.lactis toxin insensitive), respectively, have been isolated independently (Kawamoto et al., 1990; Butler et al., 1994; Kishida et al., 1996). Sensitivity of these mutants towards intracellular, conditional expression of the  $\gamma$ -toxin can distinguish zymocin binding/uptake (class I) from potential  $\gamma$ -toxin target site mutants (class II). Mutations in the class I genes SKT5 and KTI2 (isoallelic with CHS4 and CHS3, respectively) affect chitin biosynthesis and only lead to exo-zymocin resistance (Kawamoto et al., 1993; Butler et al., 1994), whereas mutations in the class II genes KTI12, IKI1 and IKI3

confer resistance against exo-zymocin and endogenously expressed  $\gamma$ -toxin (Butler *et al.*, 1994; Yajima *et al.*, 1997). Despite these recent advances, the intracellular toxin target site (TOT) still remains elusive and earlier reports that adenylate cyclase, Cdc35p, is involved in zymocin action (Sugisaki *et al.*, 1983) have been ruled out (White *et al.*, 1989).

To understand the zymocin mode of action we isolated several  $\gamma$ -toxin target site (tot) mutants by way of mTn3 transposon mutagenesis and PCR-mediated gene disruption following conditional expression of the  $\gamma$ -toxin from within S.cerevisiae cells. Thus, we identified mTn3:: integrations and re-verified individual knockouts in several TOT genes. The  $tot\Delta$  strains are resistant to both exo-zymocin and endogenous y-toxin expression, providing evidence for true toxin target site mutants. Moreover, loss of TOT gene function renders yeast cells additional phenotypes: slow growth, a G<sub>1</sub> cell cycle delay and sensitivity towards the drugs caffeine, Calcofluor White and 6-azauracil (6-AU). Consistent with the latter, which implies a functional role in transcription elongation, we identified the TOT gene products as constituents of Elongator and components associated with Elongator, a multisubunit complex interacting with elongating RNA polymerase II holoenzyme.

# Results

# Molecular identification of TOT genes and analysis of the tot phenotypes

Among a pool of 100 000 yeast clones carrying insertions of the mini-transposon mTn3::lacZ::LEU2 (mTn3)  $\gamma$ -toxin-resistant clones were screened for by switching on the expression of the  $\gamma$ -toxin from the UAS<sub>GAL1</sub> promoter on 2% galactose plates. Three hundred and twelve clones able to grow on galactose were selected over a period of 7 days. Of these, roughly one-sixth (58 clones) showed  $\beta$ -galactosidase expression in a filter assay, indicating in-frame lacZ fusions. After pHMS14 plasmid rescue, restriction pattern analysis and retransformation into non-mutagenized reporter strain LS20, 41 candidates remained to be subjected to a second viability screen using inducible y-toxin expression from another regulatable promoter (UAS<sub>MET25</sub>) to eliminate promoter-specific false positives related to  $UAS_{GALI}$ . Fifteen clones were able to grow in the absence of added methionine. These were finally subjected to a killer eclipse assay testing exozymocin resistance. Only three candidates passed this test. The TOT1-3 genes disrupted by mTn3 in these three candidates were identified by vectorette PCR and sequencing as YLR384c (ELP1/IKI1/TOT1), YGR200c (ELP2/ TOT2) and YPL086c (ELP3/TOT3).

To verify that zymocin resistance caused by the mTn3 minitransposon insertion was due to inactivation of the *TOT* genes, individual knockouts of *TOT1–3* were constructed by PCR-mediated gene disruption. In addition, two more genes reported previously as zymocin resistance determinants (Butler *et al.*, 1994; Yajima *et al.*, 1997) were individually disrupted and shown to display a *tot* phenotype: *KTI12* (*TOT4*) and *IKI1* (*TOT5*). All these *tot* $\Delta$  cells showed resistance towards endogenously expressed  $\gamma$ -toxin (Figure 1A) as well as against extracellular holozymocin (Figure 1B), indicating that the *tot1–5* $\Delta$  strains

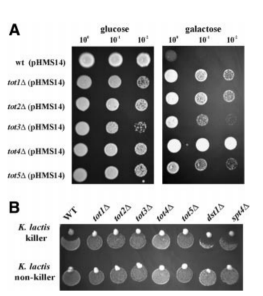
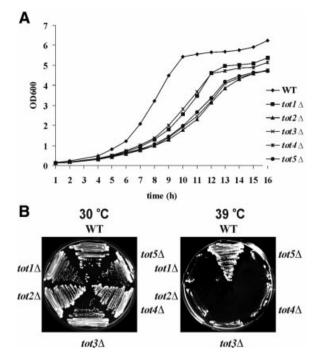


Fig. 1. The *tot* $\Delta$  mutants are resistant towards endogenously expressed  $\gamma$ -toxin and exogenously applied *K.lactis* zymocin. (A)  $\gamma$ -toxin assay. Serial dilutions of toxin-sensitive (wt) and -resistant (*tot1–5* $\Delta$ ) cells transformed with the *GAL1*-driven  $\gamma$ -toxin vector pHMS14 were replica spotted on repressing (glucose) and inducing (galactose) rich medium and grown for 2 days at 30°C. Lack of growth indicates  $\gamma$ -toxin sensitivity. (B) Killer eclipse assay. Wild-type, *tot1–5* $\Delta$  strains, *dst1* $\Delta$  and *spt4* $\Delta$  cells were spotted twice on to YPD medium. A *K.lactis* strain (upper row: killer strain AWJ137; lower row: non-killer strain NK40) was set on to the edge of these spots and incubated for 2 days at 30°C. Zones of inhibition around the zymocin-secreting killer strain (upper row) indicate zymocin sensitivity; lack of growth inhibition equals zymocin resistance.

are true toxin target site mutants. To analyse whether the  $tot 1-5\Delta$  mutations were dominant or recessive, the null mutants were crossed with LF20, the wild-type zymocinsensitive strain of opposite mating type.  $TOT/tot\Delta$ heterozygous diploids obtained in this way were all sensitive towards exo-zymocin and intracellular y-toxin (data not shown), showing that each of the  $tot1-5\Delta$ mutations behaves recessively. Since TOT1, TOT2 and TOT3 were recently identified as ELP1, ELP2 and ELP3, structural genes encoding components of RNA polymerase II Elongator (Otero et al., 1999; Wittschieben et al., 1999; Fellows et al., 2000), mutants lacking transcription factors TFIIS (dst1\Delta) (Archambault et al., 1992) and Spt4p (*spt4* $\Delta$ ) (Hartzog *et al.*, 1998; Wada *et al.*, 1998), both of which affect elongation but are not components of the Elongator complex, were assayed against exo-zymocin.  $dst1\Delta$  and  $spt4\Delta$  cells were found to be as sensitive towards zymocin as the wild-type strain LS20 (Figure 1B), suggesting that zymocin resistance is not generally associated with elongation mutants but involves specifically ELP/TOT gene function. As shown previously by Butler et al. (1994), KTI12 (TOT4) overexpression leads to zymocin resistance, while IKI1 (TOT5) overexpression does not (Yajima et al., 1997). To test the overexpression phenotype of TOT1-TOT4, the genes were overexpressed from their native promoter on multicopy yeast episomal vectors in the wild-type background of strain LS20. TOT4 was the only gene causing  $\gamma$ -toxin resistance in multicopy (data not shown).

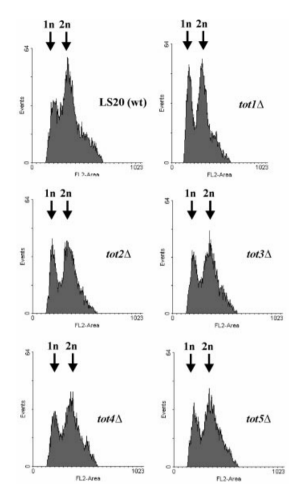
We next analysed the *tot* $\Delta$  cells under various growth conditions. Deletion of each *TOT* gene renders cells a slow growth phenotype (Slg<sup>+</sup>) at 30°C (Figure 2A), which leads



**Fig. 2.** Deletion of *TOT1–5* genes confers slow growth and thermosensitive phenotypes. (**A**) Growth curve of  $tot1-5\Delta$  strains.  $tot1-5\Delta$  cells show a slow growth phenotype and do not reach the biomass of the wild-type reference in stationary phase. Strains were grown in YPD medium (2% glucose) and growth was measured at  $OD_{600}$  over a period of 16 h. (**B**) Ts phenotype of  $tot1-5\Delta$  strains. Strains were streaked on YPD and incubated for 3 days at 30 and 39°C.

to thermosensitivity above 38°C (Figure 2B). Doubling times are lengthened by a factor of 1.5-2.0, resulting in growth rate reductions from  $\mu = 0.5$  (wild type) to  $\mu = 0.25 - 0.3$  (tot1-5 $\Delta$ ). To check whether the Slg<sup>+</sup> phenotype was correlated with a delay in a specific phase of the cell cycle, FACS analysis of exponentially growing  $tot\Delta$  strains was performed together with wild-type strain LS20 as a control (Figure 3). This analysis showed that during exponential growth all  $tot\Delta$  mutant strains had a higher percentage of cells with a 1n DNA content than did wild-type cells (Figure 3). Thus, deletion of any TOT gene leads to a significant delay in the G1 phase of the cell cycle, indicating that the TOT gene products are important for normal cell cycle progression. The thermosensitive phenotype of the *tot* $\Delta$  cells could be partially rescued by addition of 1 M sorbitol to the growth medium; however, the effect was not as striking as with a  $chs3\Delta$  cell wall chitin mutant included as a positive reference strain (data not shown).

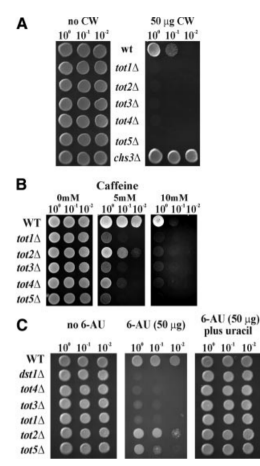
To further check whether the  $tot\Delta$  cells might be affected in cell wall integrity, we tested growth behaviour in the presence of the purine analogue caffeine and the fluorochrome Calcofluor White, a cell wall poison with high affinity for yeast cell wall chitin (Hampsey, 1997). As compared with wild-type strain LS20, all  $tot\Delta$  cells were hypersensitive towards Calcofluor White, indicative of a cell wall defect (Figure 4A). As a Calcofluor Whiteresistant positive control we included the *chs3*\Delta cell wall chitin mutant (Figure 4A). Sensitivity to caffeine was also shown by all  $tot\Delta$  cells; however, to varying degrees (Figure 4B). Thus, the  $tot5\Delta$  strain was more sensitive than



**Fig. 3.** The slow growth phenotype of  $tot1-5\Delta$  strains correlates with a significant delay in the G<sub>1</sub> phase of the cell cycle. FACS analyses of exponential growing  $tot1-5\Delta$  cells show an extended 1*n* peak in comparison with the wild-type strain, indicating a delay in the G<sub>1</sub> to S transition.

 $tot1\Delta$ ,  $tot3\Delta$  and  $tot4\Delta$  cells, while the  $tot2\Delta$  mutant was less sensitive to the drug. This phenotype suggests that cell wall integrity may be affected in all the  $tot\Delta$  cells, presumably by involvement of the Pkc1p MAP kinase signalling pathway, one of the known cellular caffeine targets (Hampsey, 1997).

Inspired by parallel identification of TOT1-3 as transcriptional Elongator genes ELP1-3 (Otero et al., 1999; Wittschieben et al., 1999; Fellows et al., 2000), we checked the effect of TOT gene deletion on transcriptional processes using 6-AU as indicator drug. 6-AU inhibits the nucleotide biosynthesis pathway in yeast, leading to a depletion of UTP and GTP, which in turn affects the efficacy of RNA polymerase II holoenzyme during transcriptional elongation (Exinger and Lacroute, 1992). As shown in Figure 4C, all  $tot\Delta$  strains showed different degrees of sensitivity to 6-AU. A  $dst1\Delta$  strain lacking the elongin TFIIS was used as positive control (Archambault et al., 1992). The strains  $tot1\Delta$ ,  $tot3\Delta$  and  $tot4\Delta$  exhibited nearly the same sensitivity as the  $dstl\Delta$  control. Less sensitivity was shown by  $tot5\Delta$  and  $tot2\Delta$ . All  $tot\Delta$  strains as well as  $dst \Delta$  could grow again on 6-AU when uracil was added to the media. In conclusion, this drug-induced phenotype suggests a role for the putative  $\gamma$ -toxin, TOT, in

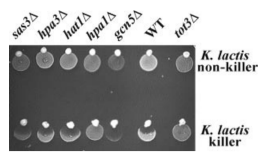


**Fig. 4.** More *tot* phenotypes. (**A**) Calcofluor White sensitivity. Serial dilutions of strains were replica spotted on YPD plates containing no or 50 µg of Calcofluor White (CW). All *tot* $\Delta$  strains show hypersensitivity against Calcofluor White, whereas the positive reference, *chs3* $\Delta$ , displays resistance towards the drug. (**B**) Caffeine sensitivity. Strains were spotted on YPD plates containing up to 10 mM caffeine and grown for 3 days at 30°C. All *tot* $\Delta$  strains show more or less hypersensitivity towards caffeine. (**C**) 6-AU phenotype. Strains were spotted on SD plates containing no 6-AU, 50 µg/ml 6-AU or 50 µg/ml 6-AU plus uracil. Except for the *tot* $\Delta$  mutant, which is mildly affected by the drug, all other *tot* $\Delta$  strains show hypersensitivity towards 6-AU. The *dst* $1\Delta$  mutant served as a positive 6-AU-hypersensitive control strain.

transcript elongation *in vivo*. Moreover, loss of *TOT* gene function has a pleiotropic effect on a yeast cell's performance, leading to a complex *tot* phenotype that includes zymocin and  $\gamma$ -toxin resistance, slow growth, G<sub>1</sub> cell cycle delay and thermosensitivity as well as sensitivity towards the drugs Calcofluor White, caffeine and 6-AU. None of these *tot* phenotypes was severely amplified on combining double *tot1* $\Delta$ *tot2* $\Delta$ , *tot3* $\Delta$ *tot2* $\Delta$ , *tot4* $\Delta$ *tot2* $\Delta$  or *tot5* $\Delta$ *tot2* $\Delta$  mutations in one genetic background (data not shown), indicating that the *TOT1–5* genes are functionally related. Summing up from these phenotypic analyses, TOT can be considered to have an important role for cellular growth.

# The HAT activity of Tot3p is essential for zymocin action

The *S.cerevisiae* genome encodes other histone acetyltransferase (HAT) activities than the one associated with Elp3p (Tot3p) (Wittschieben *et al.*, 1999). To check



**Fig. 5.** Effect of HAT gene deletions on zymocin sensitivity. Strains deleted in the HAT-encoding genes *SAS3*, *HPA3*, *HAT1*, *HPA1/ELP3*, *GCN5* and *TOT3/ELP3* were subjected to a killer eclipse assay essentially as described in Figure 1. Deletion of *TOT3* confers zymocin resistance, whereas the other HAT gene deletions tested are zymocin sensitive.

whether histone acetylation is needed for zymocin action generally, we tested other HAT gene knockouts in *GCN5*, *SAS3*, *HPA1*, *HPA3* and *HAT1* (Kleff *et al.*, 1995; Brownell *et al.*, 1996; Reifsnyder *et al.*, 1996; Brown *et al.*, 2000) using zymocin eclipse assays. As illustrated in Figure 5, all HAT knockouts except for *HPA1*, which is isoallelic to *TOT3/ELP3*, remained zymocin sensitive. Thus, the Elp3p/Tot3p-associated HAT activity appears to be required for zymocin action, whereas other non-Elongator HATs are dispensable for zymocicity.

To check whether the HAT-domain activity of Elp3p (Tot3p) itself is required for y-toxin action, point mutations in ELP3 that change two conserved tyrosine residues to alanine in the B motif of the putative catalytic HAT domain (Y540A and Y541A having <25 and ~35% of wild-type Elp3p-HAT activity, respectively) (Wittschieben et al., 2000) were tested. These mutant alleles were expressed from their native promoters on CEN plasmids (pBOP60-14: Y540A; pBOP60-15: Y541A; and pBOP60-13: wild-type ELP3 allele) in the  $tot3\Delta$  mutant and the LS20 wild-type strains co-maintaining the GAL1-driven  $\gamma$ -toxin vector pHMS14. The resulting strains were checked for complementation of the tot3 $\Delta$ -associated  $\gamma$ -toxin resistance phenotype as well as suppression of  $\gamma$ -toxin sensitivity in wild-type LS20 (Figure 6A) using glucose to galactose shift assays.

Both the Y540A and Y541A mutant ELP3 alleles failed to complement the resistance towards  $\gamma$ -toxin of tot3 $\Delta$ cells while the wild-type ELP3 allele was able to do so (Figure 6A). Furthermore, the Y540A allele (<25% HAT activity) slightly suppressed the sensitivity of wild-type LS20 against  $\gamma$ -toxin, whereas the Y541A allele (~35%) HAT activity) did not (Figure 6A). When overexpressed from multicopy yeast episomal vectors, both alleles (pLF31: Y540A and pLF30: Y541A) suppressed the sensitivity of the wild-type strain LS20 towards  $\gamma$ -toxin, resulting in a resistance phenotype, while the wild-type ELP3 allele did not (Figure 6B). Thus, it appears that a significant reduction of activation of the HAT encoded by ELP3 phenocopies the effect of TOT3/ELP3 gene inactivation on  $\gamma$ -toxin sensitivity, indicating that the HAT catalytic activity of Elp3/Tot3p is required for zymocin action. Moreover, the mutant alleles are semidominant over wild type, indicating that the gene products are incorporated into the Elongator complex, leading to a reduction of Elongator activity.



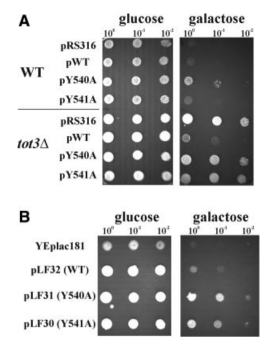


Fig. 6. The HAT activity of Elp3p/Tot3p is essential for  $\gamma$ -toxin action. (A) Wild-type and  $tot3\Delta$  strains containing the *GAL1*-driven  $\gamma$ -toxin vector pHMS14 were transformed with CEN plasmids carrying no insert (pRS316), the wild-type *ELP3* allele (pWT) or mutated *elp3* alleles (pY540A and pY541A). Serial dilutions of the resulting transformants were replica spotted on glucose-repressing and galactose-inducing medium. The mutant *elp3* alleles fail to complement the  $tot3\Delta$ -associated  $\gamma$ -toxin resistance phenotype. In the wild-type background, the Y540A allele slightly suppresses  $\gamma$ -toxin sensitivity. (B) When overexpressed from yeast multicopy plasmids, both the mutant *elp3* alleles (pLF31: Y540A and pLF30: Y541A) suppress the  $\gamma$ -toxin sensitivity of the wild-type strain co-maintaining pHMS14, whereas vector without insert (YEplac181) and the wild-type *ELP3* allele (pLF32) are not able to do so.

#### Co-immunoprecipitation of Tot proteins

Using PCR-mediated one-step tagging in vivo (Knop et al., 1999), all five Tot proteins were C-terminally marked with the c-Myc (Figure 7A) and HA (not shown) epitope tags to analyse gene expression at the translational level and to assess protein-protein interaction using co-immunoprecipitation. As judged from zymocin eclipse assays. TOT1-5 gene tagging had no effect on the biological activity of the individual Tot proteins, i.e. all tagged strains remained as zymocin sensitive as the wild-type reference strain (data not shown). Epitope-tagging identified TOT1-5 as protein-encoding structural genes. Thus, total protein extracts from exponentially grown TOT1-5- $(c-myc)_3$  cells expressed single Tot1-5 polypeptides of estimated molecular weights consistent with the predicted ones that cross-reacted with the anti-c-Mvc monoclonal antibody 9E10 (Figure 7A).

As reported, Elp1p (Tot1p), Elp2p (Tot2p) and Elp3p (Tot3p) associate together within Elongator, a 650 kDa multisubunit complex that interacts specifically with elongating RNA polymerase II holoenzyme (Otero *et al.*, 1999; Wittschieben *et al.*, 1999; Fellows *et al.*, 2000). Consistent with this, we could show by way of co-immunoprecipitation that Tot3p interacts with Tot1p and Tot2p (Figure 7B) and that Tot2p associates with Tot1p and Tot3p (data not shown). Moreover, all three Elongator

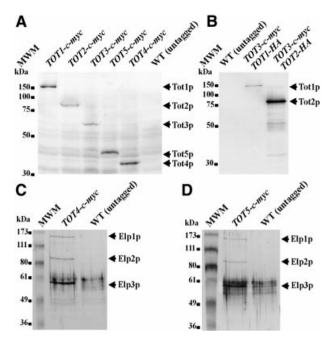


Fig. 7. Tot protein detection and co-immunoprecipitation.  $(\mathbf{A})$  Identification of *TOT1–5* as protein-encoding structural genes. Western analysis: Tot1-5p were tagged individually by one-step in vivo tagging with a triple c-Myc tag. Total protein extracts (50  $\mu g)$  were separated using 10% SDS-PAGE, electroblotted and immunoprobed with the anti-c-Myc antibody 9E10. The position of each tagged Tot protein is indicated by an arrow. (B) Tot3p interacts with Tot1p and Tot2p. Protein extracts from strains expressing c-Myc-tagged Tot3p together with HA-tagged Tot1p or HA-tagged Tot2p were immunoprecipitated using the anti-c-Myc antibody 9E10. Precipitates were then separated using 10% SDS-PAGE, electroblotted and immunoprobed with the anti-HA antibody 12CA5. The positions of HA-tagged Tot1p and Tot2p are indicated by arrows. (C and D) Tot4p (C) and Tot5p (D) interact with Elp1p (Tot1p), Elp2p (Tot2p) and Elp3p (Tot3p). Protein extracts from strains expressing c-Myc-tagged Tot4p (C) or Tot5p (D) were immunoprecipitated using the anti-c-Myc antibody 9E10, separated on a 10% SDS-PAGE gel, electroblotted and immunoprobed by western analysis using polyclonal anti-Elp1p, anti-Elp2p and anti-Elp3p antibodies. The positions of co-immunoprecipitated proteins, Elp1p, Elp2p and Elp3p, are indicated by arrows.

constituents, Elp1p, Elp2p and Elp3p, could also be shown to interact with Tot4p (Figure 7C) and Tot5p (Figure 7D). These co-immunoprecipitations involved precipitation of c-Myc-tagged Tot4p and Tot5p with the anti-c-Myc antibody 9E10 covalently linked to protein A–Sepharose and detection by polyclonal anti-Elp1p, -Elp2p and -Elp3p antibodies (Otero *et al.*, 1999; Wittschieben *et al.*, 1999; Fellows *et al.*, 2000). In conclusion, all *TOT* gene products were found to be associated in a multisubunit complex consisting of (at least) five polypeptides, three of which were recently identified as components of the Elongator complex. Thus, the *S.cerevisiae* Elongator appears to be part of or to represent the putative *K.lactis*  $\gamma$ -toxin target, TOT.

# Gene transcription in response to the K.lactis zymocin

Since the Elp3p-associated HAT activity appears to be essential for Elongator function *in vivo* (Wittschieben *et al.*, 2000), it is highly likely that TOT/Elongator plays a

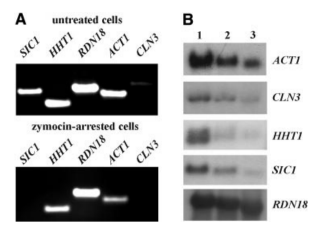


Fig. 8. Gene transcription in response to the *K.lactis* zymocin. (A) Identical amounts of total RNA isolated from untreated and zymocin-arrested cells were subjected to RT–PCR experiments to study the transcription of RNA polymerase I-dependent (*RDN18*) and RNA polymerase II-dependent (*SIC1*, *HHT1*, *ACT1* and *CLN3*) genes by 1% TBE–agarose gel electrophoresis (B) Identical amounts of total RNA (10  $\mu$ g) isolated from untreated cells (lane 1) and cells arrested by zymocin for 3 h (lane 2) and 6 h (lane 3) were subjected to northern blot analysis using probes specific for the *RDN18*, *SIC1*, *HHT1*, *ACT1* and *CLN3* genes.

role in transcript elongation or promoter remodelling by maintaining transcriptionally competent chromatin while piggybacked to the elongating form of the RNA polymerase II holoenzyme (Travers, 1999). To investigate the effect of the K.lactis zymocin on transcriptional processes, identical amounts of total RNA prepared from untreated and zymocin-arrested cells were subjected to RT-PCR experiments using primers specific for RNA polymerase I-dependent and RNA polymerase II-driven genes. As illustrated in Figure 8A, RNA polymerase Idependent transcription of the 18S rRNA gene (RDN18) remained largely unaffected, whereas transcription of the RNA polymerase II-dependent genes (SIC1, CLN3, HHT1 and ACT1) decreased significantly upon treating cells with the K.lactis zymocin. Similarly, this differential display was seen when using conventional northern hybridization studies (Figure 8B). Here, the effect of the K.lactis zymocin on RNA polymerase II-dependent transcription was particularly pronounced with regard to SIC1, CLN3 and HHT1 gene activation. Our observation that RNA polymerase I-dependent transcription remains largely unaffected by the K.lactis zymocin is consistent with the results of a previous study by Butler et al. (1991c) showing that bulk RNA synthesis, measured by incorporation of radiolabelled uracil, is not affected during zymocin treatment. In conclusion, the *K*.lactis zymocin appears to preferentially affect RNA polymerase II-dependent gene transcription.

## Discussion

The *K.lactis* zymocin is toxic against a variety of sensitive yeast genera including *S.cerevisiae*. Earlier reports that the zymocin functions on *S.cerevisiae* by inhibiting the adenylate cyclase, and hence abolishing the roles of cAMP essential for mitotic growth and cell division (Sugisaki *et al.*, 1983), have been disproved (White *et al.*, 1989). Thus, its mode of action still remains unclear. Zymocin-

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resistant S.cerevisiae skt, iki and kti mutants have been isolated previously (Kawamoto et al., 1990; Butler et al., 1994; Kishida et al., 1996). Sensitivity of these towards intracellular, conditional expression of the y-toxin from inducible promoters has distinguished zymocin binding/ uptake (class I) from  $\gamma$ -toxin target site mutants (class II). The presence of as many as 10 distinct kti class II complementation groups suggests that a complex pathway transduces the zymocin's inhibitory effect (Butler et al., 1994). While some of them may be involved in the expression of target(s) inhibited by the  $\gamma$ -toxin, a number of proteins could also participate in the process that is blocked by it. These might act in a biochemical pathway or, alternatively, form a putative target complex containing several components. In favour of the latter, our genetic screens involving mTn3 tagging and PCR-mediated gene disruption to search for potential  $\gamma$ -toxin target (tot) mutants have identified the TOT complex as a putative target. The tot phenotypes common to all TOT gene deletions provide genetic evidence that the specified Tot gene products are functionally related to one another. Thus, in addition to zymocin and  $\gamma$ -toxin resistance, tot1-5 $\Delta$  cells commonly display slow growth and thermosensitive phenotypes as well as G<sub>1</sub> cell cycle delay and hypersensitivity towards the drugs Calcofluor White, caffeine and 6-AU. The fact that none of these phenotypes is severely amplified on combining double  $tot\Delta$  mutations in one background can be genetically interpreted as TOT1-5 genes functioning in the same process. Moreover, our re-identification of ELP1/IKI3, ELP2 and ELP3, structural genes encoding Elongator complex components (Otero et al., 1999; Wittschieben et al., 1999; Fellows et al., 2000), as TOT1, TOT2 and TOT3, respectively, provides substantial evidence for the 'putative target complex' hypothesis. Consistent with this, our co-immunoprecipitation experiments have shown that the putative y-toxin target complex, TOT, consists of two more constituents encoded by KTI12 (TOT4) and IKI1 (TOT5) in addition to the known Elongator components. Since  $tot4\Delta$  and  $tot5\Delta$  cells are phenotypically indistinguishable from  $tot1-3\Delta$  (elp1-3 $\Delta$ ) cells, it may well be envisaged that Tot4p and Tot5p constitute further as yet uncharacterized Elongator components, suggesting that the Elongator complex represents or is part of TOT.

By introducing ELP3 HAT domain mutations into an  $elp3\Delta$  strain, it has been shown recently that the HAT activity associated with Elp3p is essential for Elongator function in vivo (Wittschieben et al., 2000). Thus, in spite of being incorporated into Elongator complexes, the two mutant Elp3 proteins tested (Y540A and Y541A with <25 and ~35% HAT activity compared with wild-type Elp3p, respectively) failed to rescue the cells from the  $elp3\Delta$ associated phenotypes. We have shown here that, similarly, both mutant Y540A and Y541A Elp3 proteins failed to confer sensitivity towards the  $\gamma$ -toxin when re-introduced into our toxin-resistant  $tot3\Delta$  (*elp3* $\Delta$ ) reporter strain. The point mutants behaved as dominant-negative alleles with respect to  $\gamma$ -toxin sensitivity. Thus,  $\gamma$ -toxin sensitivity apparently requires the Elongator-associated HAT activity. Moreover, they conferred  $\gamma$ -toxin resistance when introduced on multicopy episomal vectors into a toxinsensitive ELP3 wild-type reporter strain. Probably, the mutant overproduced Elp3p variants compete for incorporation into Elongator with wild-type Elp3p, thus reducing the concentration of active Elongator complexes. Taken together, these results indicate that not only inactivation of Elongator by way of *TOT1–5* gene deletions, which may ultimately result in non-assembly of functional Elongator complexes, but also *in vivo* assembly of Elongator complexes with significantly reduced HAT activity lead to the *tot* phenotype, including resistance towards the *K.lactis* zymocin.

Since all the TOT genes are non-essential, the zymocin cannot be considered to cause a G<sub>1</sub> arrest by inhibition/ inactivation, direct or otherwise, of Elongator. Instead, sensitive S.cerevisiae cells require a functional Elongator in order to be able to respond to the K.lactis zymocin. This requirement for Elongator is genetically sustained by the observation that cells lacking either non-Elongator HATs  $(gcn5\Delta, hpa3\Delta, hat1\Delta \text{ and } sas2\Delta)$  (Kleff *et al.*, 1995; Brownell et al., 1996; Reifsnyder et al., 1996; Brown et al., 2000) or non-Elongator transcription elongation factors  $(dst1\Delta \text{ and } spt4\Delta)$  (Archambault *et al.*, 1992; Hartzog et al., 1998; Wada et al., 1998) are not able to alter  $\gamma$ -toxin sensitivity in a way the  $tot1-5\Delta$  strains can. This implies that the *K*.lactis zymocin can activate a pathway that is inhibitory to the cell cycle and of which TOT/Elongator is a part.

As for the molecular mode of action of the K.lactis zymocin, our results provide at least two possible models. First, one might envisage direct interaction of the  $\gamma$ -toxin with fully assembled TOT/Elongator complexes in order to promote the observed  $G_1$  cell cycle arrest. Thus,  $\gamma$ -toxinbound TOT/Elongator and free Elongator would compete for a limited supply of a downstream effector molecule that is a key positive regulator of the cell cycle needed for events in late  $G_1$ . Given the fact that Elongator has recently been demonstrated to associate with the CTDhyperphosphorylated form of RNA polymerase II and proposed to be exchanged for Mediator upon CTD dephosphorylation after transcription termination (Otero et al., 1999; Wittschieben et al., 1999), it is conceivable to speculate that RNA polymerase II itself may be a prime candidate for this critical effector protein. In the absence of the y-toxin, RNA polymerase II would associate with TOT/Elongator complexes to acquire full transcriptional activity and to promote a normal G<sub>1</sub> to S transition by activation of START-specific genes. In the presence of the y-toxin, RNA polymerase II would be sequestered and inactivated by the  $\gamma$ -toxin-bound TOT/Elongator complex, leading to a  $G_1$  cell cycle arrest, e.g.  $\gamma$ -toxin-bound TOT/ Elongator complexes associated with RNA polymerase II may prevent exchange for Mediator during or after transcription termination, thus lowering the concentration of RNA polymerase II able to initiate transcription of START-specific genes needed for a proper G1 to S cell cycle progression. Consistent with this model, we found that transcription of RNA polymerase II-dependent genes (SIC1, CLN3, HHT1 and ACT1) was specifically downregulated in zymocin-arrested cells, whereas RNA polymerase I-dependent transcription of the 18S rRNA gene (RDN18) remained largely unaffected. Therefore, we favour the hypothesis that the K.lactis zymocin and its cytotoxic  $\gamma$  subunit have the TOT/Elongator-associated RNA polymerase II holoenzyme as a putative direct target. Alternatively, one might envisage that the  $\gamma$ -toxin target could be a gene product whose transcript is downregulated in strains lacking TOT/Elongator function. To address this question in more detail we are currently designing genome-wide micro-array approaches to identify genes whose transcription is particularly affected in the *tot* mutants.

## Materials and methods

#### Strains, media, DNA constructs and general methods

Routine bacterial transformations used for constructing recombinant DNA, yeast plasmid shuttles and rescues as well as amplification of the mTn3-tagged yeast DNA library involved electroporation of Escherichia coli strains DH5a, XL1-Blue and TOP10F'. These were grown in Luria-Bertani medium supplemented with one or several of the following additions: ampicillin (100 µg/ml), kanamycin (80 µg/ml), X-Gal (80 μg/ml) and isopropyl-β-D-thiogalactopyranoside (IPTG) (50 µg/ml). All yeast strains used or generated in this study are described in Table I. Standard rich and minimal growth media, YEPD and SC, respectively, were prepared essentially as described by Sherman (1991). For phenotypic analysis, these media were supplemented with 6-AU (50 µg/ml), caffeine (1–20 mM) or Calcofluor White (50–1000 µg/ml). For 6-AU assays, Ura+ transformants (YCplac33) (Gietz and Sugino, 1988) of the tot $\Delta$  strains were used. Yeast transformations involved electroporation (Becker and Guarente, 1991) as well as the lithium acetate method (Gietz et al., 1992). Mating type switching of LL20 (MATa, see Table I) to generate MATa cells (LF20, see Table I) was accomplished using HO gene expression from plasmid pHAL24 (YEplac181 carrying the HO gene on a 2.5 kb genomic HindIII fragment) followed by plasmid segregation. To create Ura- derivatives of strain LL20, yeast cells were transformed with a  $ura3\Delta$  deleter plasmid followed by selection against URA3 on SC minimal medium containing uracil and 5-fluoroorotic acid (5-FOA). The deleter plasmid was constructed by autoligation of the YDp-U backbone fragment (Berben et al., 1991) obtained by double digestion with EcoRV and SmaI to delete an internal URA3-borne segment and reconstitute a non-functional ura3A allele. Among Uratransformants (termed LS20, see Table I), ura3-linked auxotrophy was distinguished from ura5 mutants by single copy URA3 gene complementation (YCplac33). To construct the methionine-repressible  $\gamma$ -toxin expression vector pHAL9, the y-toxin gene was fused as an XhoI-SalI restriction fragment from plasmid pARB1 (Butler et al., 1994) to the UAS<sub>MET25</sub> promoter of SalI-cut vector p413MET25 (Mumberg et al., 1994). The latter was obtained by deleting the GFP gene from pGFP-NFUS (Niedenthal et al., 1996) using BamHI and autoligation of the vector backbone. Galactose-inducible expression of the  $\gamma$ -toxin was achieved using pHMS14, a p413-based vector (Mumberg et al., 1995), carrying the UAS<sub>GALI</sub>-\gamma-toxin fusion on a 1.4 kb SstI-EcoRI fragment from pARB1 (Butler et al., 1994; Schaffrath et al., 1997). As a UASGALI vector control, plasmid pHMS22 (p413 carrying the UAS<sub>GAL1</sub> promoter only) was used. Plasmids for overexpression of Tot1p (pFF14), Tot2p (pFF10) and Tot3p (pFF9) were constructed by cloning the coding sequence with UAS and terminator into YEplac195 using SacI-SalI (pFF14), PstI-BamHI (pFF10) and HindIII-SalI (pFF9). Mutated alleles of ELP3/TOT3 (Y540A, Y541A) were cloned for overexpression in YEplac181 by using BamHI-KpnI restriction fragments of pBOP60-13, pBOP60-14 and pBOP60-15 (Wittschieben et al., 2000), generating pLF30 (Y541A), pLF31 (Y540A) and pLF32 (wild type).

#### Isolation and genetic analysis of tot mutants

A yeast transformant population carrying mini-transposon (mTn3) insertions randomly integrated into the genome was constructed as follows. First, strain LS20 was transformed to histidine prototrophy with pHMS14 carrying the  $UAS_{GALI}$ - $\gamma$ -toxin fusion. Several His<sup>+</sup> candidates (termed LS20') were checked for  $\gamma$ -toxin sensitivity by replica plating on to galactose SC his<sup>-</sup> medium, resulting in a Gal<sup>-</sup> phenotype. Next, LS20' was subjected to transposon mutagenesis using electroporation-mediated transformation with the *NotI*-digested mTn3::yeast insertion library (Burns *et al.*, 1994) selecting for the mTn3-based *LEU2* marker on SC his<sup>-</sup>, leu<sup>-</sup> medium. His<sup>+</sup> Leu<sup>+</sup> yeast transformants were subsequently replica plated on galactose medium and incubated for up to 7 days to identify Gal<sup>+</sup> tox<sup>R</sup> candidates. To distinguish genomic mTn3::integrations of the  $UAS_{GALI}$ - $\gamma$ -toxin fusion on pHMS14, total DNA preparations obtained from selected Gal<sup>+</sup> isolates were used for yeast–*E.coli* plasmid rescue and

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Strain	Genotype	Reference
K.lactis		
NK40	MAT $\alpha$ , ade1, ade2, leu2 [k1 <sup>0</sup> k2 <sup>+</sup> ]	Gunge et al. (1981)
AWJ137	$MATa, \ leu2, \ trp1 \ [k1^+ \ k2^+]$	Kämper et al. (1991)
S.cerevisiae		
FY1679-08A	MATa, ura3-52, leu2 $\Delta$ 1, trp1 $\Delta$ 63, his3 $\Delta$ 200, GAL	Euroscarf
FY1646	MAT $\alpha$ , his4-912 $\delta$ , lys2-12 $\delta\delta$ , leu2 $\Delta$ 1, spt4 $\Delta$ ::HIS3	F.Winston
GMY27	MATα, ade2-101, leu2-3,-112, his3Δ200, ura3-52, lys2, gcn5Δ::hisG	Anthony Wright
LPY2121	MATa, $ade2-101$ , $his3\Delta 200$ , $leu2\Delta 1$ , $lys2-801$ , TELadh4::URA3, $ura3-52$ , $trp1\Delta 1$ sas3 $\Delta$ ::HIS3	Darryl Auston
W303-1a	MATa, ura3-1, leu2-3, -112, his3-11, -15, trp1-1, ade2-1, can 100-1	Anne Sutton
RS1236 (SK56)	as W303-1a, but $hat 1\Delta$ ::TRP1	Anne Sutton
RS1392 (YCW2)	as W303-1a, but $hpa1\Delta$ : URA3	Anne Sutton
YRP13	as W303-1a, but $hpa3\Delta$ ::HIS3	Anne Sutton
LL20	MATα leu2-3, -112, his3-11, -15, GAL	NCYC 1445
LS20	as LL20, but <i>ura3</i>	this work
LF20	as LL20, but MATa	this work
LS20'	as LS20 plus pHMS14 (CEN4/HIS3/UAS <sub>GALI</sub> -γ-toxin)	this work
LFY12	as LS20, but <i>tot</i> 4\Delta:: <i>LEU2 GAL</i>	this work
FFY5	as LS20, but tot5\Delta::KILEU2 GAL	this work
FFY6	as LS20, but <i>dst1</i> \Delta:: <i>KlLEU2 GAL</i>	this work
DJY3	as LS20, but <i>chs3</i> \Delta:: <i>KlLEU2 GAL</i>	this work
LFY1a	as LS20, but TOT4-(c-myc) <sub>3</sub> ::SpHIS5	this work
FFY1t	as LS20, but TOT1-(c-myc) <sub>3</sub> ::SpHIS5	this work
FFY2t	as LS20, but TOT2-(c-myc) <sub>3</sub> ::SpHIS5	this work
FFY3t	as FY1679-08A, but TOT3-(c-myc)3::SpHIS5	this work
FFY5t	as FY1679-08A, but TOT5-(c-myc) <sub>3</sub> ::SpHIS5	this work
FFY2-1dt	as FY1679-08A, but TOT3-(c-myc) <sub>3</sub> ::SpHIS5, TOT1-(HA) <sub>6</sub> ::KITRP1	this work
FFY2-3dt	Y2-3dt as FY1679-08A, but TOT3-(c-myc) <sub>3</sub> ::SpHIS5, TOT2-(HA) <sub>6</sub> ::KITRP1	

restriction enzyme analysis, using DNA of starting vector pHMS14 as positive control. Rescued plasmid DNAs identical to the SalI pattern of pHMS14 were retransformed into fresh recipient strain LS20 and checked for tox<sup>S</sup> by conditionally switching on  $\gamma$ -toxin expression on galactose SC (his<sup>-</sup>) plates. Clones that passed this test were next checked for UAS<sub>GALI</sub>specific false positives by using a second conditional  $\gamma$ -toxin expression approach involving the methionine-regulated promoter UAS<sub>MET25</sub> on vector pHAL9. Using this second expression approach, clones that were able to grow under inducing conditions, i.e. in the absence of methionine, were obtained. In addition, the Gal+ Leu+ His+ toxR integrants were screened for in-frame fusions of the start-codon-less lacZ gene carried on the mTn3 portion to yeast coding regions by checking  $\beta$ -galactosidase production on qualitative filter assays essentially as described (Ross-Macdonald et al., 1997). Resistance towards exo-zymocin was assayed using the killer eclipse assay (Kishida et al., 1996). To identify the yeast DNA immediately adjacent to the mTn3 integration site of the mutants, the vectorette PCR approach was used (Ross-Macdonald et al., 1998). PCR products that were specifically amplified from mTn3containing fragments were identified and directly subcloned into vector pCR2.1-TOPO using the topoisomerase cloning kit TOPO TA Version H (Invitrogen). Next, two independent subclones were sequenced for each candidate with the universal M13 reverse (5'-CAGGAAACAGCT-ATGAC-3') and -20 forward primers (5'-GTAAAACGACGGCCA G-3') and analysed using the BLAST and FASTA network services. To analyse whether the mTn3:: marked gene disruptions were recessive or dominant, the individual integrants (MATa, mTn3::LEU2, ura3, pHMS14 [HIS3]) were crossed to LF20 (MATa, leu2, his3, URA3) and diploids selected on SC medium were assayed for  $\gamma$ -toxin sensitivity/resistance by conditionally switching on y expression on galactose medium. For PCRmediated gene targeting (Wach et al., 1997) and construction of defined totA null alleles, the original YDp plasmid set (Berben et al., 1991) was modified with non-Saccharomyces markers to utilize YDp-KlL (K.lactis LEU2: Zhang et al., 1992), YDp-KlU (K.lactis URA3: Längle-Rouault and Jacobs, 1995) and YDp-SpH (S.pombe HIS5: Wach et al., 1997). Knockout primers (Table II) usually consisted of 50 unique nucleotides homologous to the 5'- and 3'-regions of the yeast gene of interest plus a common 21 nucleotide stretch homologous to the multiple cloning site of plasmid pUC9H-STOP (Berben et al., 1991), the backbone of the YDp plasmid set used as yeast marker templates for PCR-mediated gene targeting. For generating the  $tot4\Delta$  allele, a deletion construct, pYF6 (Butler et al., 1994), was alternatively used.

### Flow cytometric determination of cellular DNA content

Cells from exponential growing cultures were fixed in 70% (v/v) ethanol and stored at  $-20^{\circ}$ C. Aliquots (5 × 10<sup>7</sup> cells) of each sample were washed once with 1 ml of 50 mM sodium citrate solution and incubated in the dark for 30 min at 37°C in 1 ml of 1× PBS containing 1 mg of RNase A and 20 µg of propidium iodide. Each sample was analysed using a Becton-Dickinson FACS. The FACS contained a 15 mW argon laser with an excitation wavelength of 488 nm. Fluorescence was measured at 585 nm. Data were collected on 10 000 cells per sample. Under these conditions, fluorescence is considered to be proportional to DNA content (Hutter and Eipel, 1979).

#### Epitope tagging and immunological techniques

Epitopes were fused to genes by using PCR-based one-step *in vivo* epitope-tagging methods and tools as described by Knop *et al.* (1999). For primers used see Table II. For detection of epitope-tagged proteins, 9E10 mouse monoclonal antibody recognizing the c-Myc epitope and 3F10 rat antibody recognizing the HA epitope (Roche) were used as described (Schaffrath and Meacock, 1996). Polyclonal rabbit Elp1, Elp2 and Elp3 antibodies were kindly provided by Jesper Q.Svejstrup (ICRF, South Mimms, UK). Secondary alkaline phosphatase and peroxidase-conjugated antibodies were obtained from Jackson ImmunoResearch. Antibody cross-linking to protein A–Sepharose, preparation of protein extract and co-immunoprecipitation were carried out as described previously (Zachariae *et al.*, 1996) using B60 buffer. Probes were then checked by western analysis. For all protein methods, proteinase inhibitors (Roche) and 0.5–1 mM phenylmethylsulfonyl fluoride were used.

#### Gene transcription analyses

Total RNA was isolated from equal amounts of zymocin-arrested and untreated *S.cerevisiae* LS20 cells using the RNAeasy midi kit (Qiagen) according to the manufacturer's recommendations. Zymocin treatment and arrest were carried out and monitored as described (D.Jablonowski, L.Fichtner, V.J.Martin, R.Klassen, F.Meinhardt, M.J.R.Stark and R.Schaffrath, in preparation). RT–PCR experiments involved equal amounts of total RNA (4 µg) with the RevertAid<sup>TM</sup> kit (MBI Fermentas) for 1 h at 42°C in 20 µl reaction volumes. After first strand cDNA synthesis, 1/20 of the reaction was subjected to PCR (30 cycles) using *Taq* polymerase and oligonucleotide primers (10 µM) (Table II) to amplify fragments specific for the 18S rRNA (*RDN18*; 0.52 kb), the histone H3 (*HHT1*; 0.32 kb), the actin (*ACT1*; 0.44 kb), the G<sub>1</sub> cyclin

Table II. Oligonucleotide primers used in this study			
Name	Description	Sequence	
FF1	FW ko-primer TOT1	5′-AGAAACAGTACAAATGCCTAATGGCTTATGGTTGAACATGACAAGAGTGGCGAC GGCCAGTGAATTCCCGG-3′	
FF2	RV ko-primer TOT1	5'-CAATATGACTCTTAGGGAAATCATGAATCTCTGGAACAGGTATTTCTGGGAGCTT GGCTGCAGGTCGACGG-3'	
FF3	FW ko-primer TOT2	5'-ATGGTGGAATGTATCACTCCCGAAGCCATTTTTATAGGTGCTAACAAGCACGACG GCCAGTGAATTCCCGG-3'	
FF4	RV ko-primer TOT2	5'-CCTCAATCTTGTAATTTTGTCTGCTGGTGTTATATCCTCGTTTAGCTGCGAGCTTGG CTGCAGGTCGACGG-3'	
FF5	FW ko-primer TOT3	5'-AGATGGCTCGTCATGGAAAAGGCCCAAAAACTAACAAAAAAAGCTAGCACGAC GGCCAGTGAATTCCCGG-3'	
FF6	RV ko-primer TOT3	5'-CCAGAAATAACAGAAATTTTCTCTGAACCATGCTCTTCCTTGGCGATTCTAGCTTG GCTGCAGGTCGACGG-3'	
LF13	FW ko-primer TOT4	5′-AAACTAAACAGGCAATTTAGTAAGAAGATGCCACTGGTGCTTTTTACGGGCGACG GCCAGTGAATTCCCGG-3′	
LF14	RV ko-primer TOT4	5'-ATCTCAATTCAAGTTTTTGTTAAGATAATCAGCGAAAAGCGGACCGATCCAGCTTG GCTGCAGGTCGACGG-3'	
FF7	FW ko-primer TOT5	5' CTATTGCTACAGGTAGAACAAGATATAATGGCCAGTTCGTCACATAACCCCGACGG CCAGTGAATTCCCGG-3'	
FF8	RV ko-primer TOT5	5'-AAAAGGATCCTCATATGGATCCTCTTCATCATAATCGTCATCCTTTTCGAGCTTGG CTGCAGGTCGACGG-3'	
FF9	FW ko-primer DST1	5'-GTAGTCAGTCCGCATAAGAGCATTCATCATGGATAGTAAGGAAGTACTGGCGACGG CCAGTGAATTCCCCGG-3'	
FF10	RV ko-primer DST1	5'-TCTGTTACCACATGCTTCACATGTACAGAAAGTGGTCAATGGTTCATCCGAGCTTGG CTGCAGGTCGACGG-3'	
DJ5	FW ko-primer CHS3	5'-TCCGCAGGAAAGAAATTAGAATGACCGGCTTGAATGGAGATGATCCTGATCGACGG CCAGTGAATTCCCGG-3'	
DJ6	RV ko-primer CHS3	5'-GTCTATGCAACGAAGGAGTCACTTTCCTCCTTCCGATTGAGAATATCTTCAGCTTGG CTGCAGGTCGACGG-3'	
S3-TOT1	one-step in vivo tagging TOT1	5'-TACCTGTTCCAGAGATTCATGATTTCCCTAAGAGTCATATTGTTGATTTTCGTACGCT GCAGGTCGAC-3'	
S2-TOT1	one-step in vivo tagging TOT1	5'-CTTTACGAGCACTATAGACAGTAATTTATATAACTAAGAAAATGGTATGCATCGATG AATTCGAGCTCG-3'	
S3-TOT2	one-step in vivo tagging TOT2	5'-GTGTAGGAAGTAGTGATTTGTCCACCCGTATATACTCATTAGCATATGAACGTACGC TGCAGGTCGAC-3'	
S2-TOT2	one-step in vivo tagging TOT2	5'-ATTAACTTATCATCTCTTTTTCACATGAGAAATGATATAGATATTGCATCGATGA ATTCGAGCTCG-3'	
S3-TOT3	one-step in vivo tagging TOT3	5'-ATGGTAAACTAGGATATGAACTAGACGGTCCATACATGTCGAAAAGAATTCGTACGC TGCAGGTCGAC-3'	
S2-TOT3	one-step in vivo tagging TOT3	5'-CTGCTTGGAAAACCGGCCATGTCGGCGGCACATAAAAGTTCTATTTACCTATCGATG	
S3-TOT4	one-step in vivo tagging TOT4	AATTCGAGCTCG-3' 5'-AGGATCGGTCCGCTTTTCGCTGATTATCTTAACAAAAACTTGAATCGTACGCTGCAG	
S2-TOT4	one-step in vivo tagging TOT4	GTCGAC-3' 5'-ATTTCGTCTTGCCATTTACCTTCTGATATTAATCACATGTATATCATCGATGAATTCG	
S3-TOT5	one-step in vivo tagging TOT5	AGCTCG-3' 5'-ACGAAAAGGATGACGATTATGATGAAGAGGATCCATATGAGGATCCCTTTCGTACG	
S2-TOT5	one-step in vivo tagging TOT5	CTGCAGGTCGAC-3' 5'-TAGTTTACATAATCTGGAAGCACTCACTATTTACCATCAGTTTCTACTTTATCGATG	
RDN18 FW	RT–PCR	AATTCGAGCTCG-3′ 5′-CGCGCAAATTACCCAATCCT-3′	
<i>RDN18</i> RV	RT–PCR	5'-GGCAAATGCTTTCGCAGTAG-3'	
ACT1 FW	RT–PCR	5'-CTTCCGGTAGAACTACTGGT-3'	
ACT1 RV	RT–PCR	5'-CCTTACGGACATCGACATCA-3'	
		5'-AGCAAGAAAGTCCACTGGTG-3'	
HHT1 FW	RT-PCR		
<i>HHT1</i> RV	RT–PCR	5'-GAATGGCAGCCAAGTTGGTA-3'	
SIC1 FW	RT–PCR	5'-TTCACAGAACCTAGTCCCTG-3'	
SIC1 RV	RT–PCR	5'-ACTCCTGGCGTCATTTTTCG-3'	
CLN3 FW	RT–PCR	5'-CAATCTACGTCCCCGTTATC-3'	
CLN3 RV	RT–PCR	5'-CGCTCTTTGGAGTAGTAGCA-3'	

(CLN3; 0.61 kb) and the CKI (SIC1; 0.48 kb) genes. Northern blot analysis was carried out according to standard techniques (Sambrook *et al.*, 1989).

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#### References

Ahmed,A., Sesti,F., Ilan,N., Shih,T.M., Sturley,S.L. and Goldstein,S.A. (1999) A molecular target for viral killer toxin: TOK1 potassium channels. *Cell*, **99**, 283–291.

Archambault, J., Lacroute, F., Ruet, A. and Friesen, J.D. (1992) Genetic

#### F.Frohloff et al.

interaction between transcription elongation factor TFIIS and RNA polymerase II. *Mol. Cell. Biol.*, **12**, 4142–4152.

Becker, D.M., and Guarente, L. (1991) High-efficiency transformation of yeast by electroporation. *Methods Enzymol.*, **194**, 182–187.

- Berben, G., Dumont, J., Gilliquet, V., Bolle, P.A. and Hilger, F. (1991) The YDp plasmids: a uniform set of vectors bearing versatile gene disruption cassettes for *Saccharomyces cerevisiae*. *Yeast*, 7, 475–477.
- Brown, C.E., Lechner, T., Howe, L. and Workman, J.L. (2000) The many HATs of transcription coactivators. *Trends Biochem. Sci.*, 25, 15–19.
- Brownell, J.E., Zhou, J., Ranalli, T., Kobayashi, R., Edmondson, D.G., Roth, S.Y. and Allis, C.D. (1996) *Tetrahymena* histone acetyltransferase A: a homolog to yeast Gcn5p linking histone acetylation to gene activation. *Cell*, 84, 843–851.
- Burns, N., Grimwade, B., Ross-Macdonald, P.B., Choi, E.Y., Finberg, K., Roeder, G.S. and Snyder, M. (1994) Large-scale analysis of gene expression, protein localization, and gene disruption in *Saccharomyces cerevisiae. Genes Dev.*, 8, 1087–1105.
- Butler, A.R., White, J.H. and Stark, M.J. (1991a) Analysis of the response of Saccharomyces cerevisiae cells to Kluyveromyces lactis toxin. J. Gen. Microbiol., 137, 1749–1757.
- Butler, A.R., Porter, M. and Stark, M.J. (1991b) Intracellular expression of *Kluyveromyces lactis* toxin  $\gamma$  subunit mimics treatment with exogenous toxin and distinguishes two classes of toxin-resistant mutant. *Yeast*, **7**, 617–625.
- Butler, A.R., O'Donnell, R.W., Martin, V.J., Gooday, G.W. and Stark, M.J. (1991c) *Kluyveromyces lactis* toxin has an essential chitinase activity. *Eur. J. Biochem.*, **199**, 483–488.
- Butler,A.R., White,J.H., Folawiyo,Y., Edlin,A., Gardiner,D. and Stark,M.J. (1994) Two Saccharomyces cerevisiae genes which control sensitivity to G<sub>1</sub> arrest induced by *Kluyveromyces lactis* toxin. Mol. Cell. Biol., 14, 6306–6316.
- Exinger,F. and Lacroute,F. (1992) 6-azauracil inhibition of GTP biosynthesis in Saccharomyces cerevisiae. Curr. Genet., 22, 9–11.
- Fellows, J., Erdjument-Bromage, H., Tempst, P. and Svejstrup, J.Q. (2000) The Elp2 subunit of elongator and elongating RNA polymerase II holoenzyme is a WD40 repeat protein. *J. Biol. Chem.*, **275**, 12896–12899.
- Gietz,R.D. and Sugino,A. (1988) New yeast–*Escherichia coli* shuttle vectors constructed with *in vitro* mutagenized yeast genes lacking six-base pair restriction sites. *Gene*, 74, 527–534.
- Gietz, D., St Jean, A., Woods, R.A. and Schiestl, R.H. (1992) Improved method for high efficiency transformation of intact yeast cells. *Nucleic Acids Res.*, 20, 1425.
- Gunge, N., Tamaru, A., Ozawa, F. and Sakaguchi, K. (1981) Isolation and characterization of linear deoxyribonucleic acid plasmids from *Kluyveromyces lactis* and the plasmid-associated killer character. *J. Bacteriol.*, **145**, 382–390.
- Hampsey, M. (1997) A review of phenotypes in *Saccharomyces* cerevisiae. Yeast, 13, 1099–1133.
- Hartzog,G.A., Wada,T., Handa,H. and Winston,F. (1998) Evidence that Spt4, Spt5, and Spt6 control transcription elongation by RNA polymerase II in Saccharomyces cerevisiae. Genes Dev., 12, 357–369.
- Hutter, K.J. and Eipel, H.E. (1979) Microbial determinations by flow cytometry. J. Gen. Microbiol., 113, 369–375.
- Kämper, J., Esser, K., Gunge, N. and Meinhardt, F. (1991) Heterologous gene expression on the linear DNA killer plasmid from *Kluyveromyces lactis. Curr. Genet.*, **19**, 109–118.
- Kawamoto,S., Arai,N., Kobayashi,M., Kawahara,K., Iwaheshi,H., Tanabe,C., Hatori,H., Ohno,T. and Nakamura,T. (1990) Isolation and characterization of mutants of *Saccharomyces cerevisiae* resistant to killer toxin of *Kluyveromyces lactis. J. Ferment. Bioeng.*, 4, 222–227.
- Kawamoto,S., Sasaki,T., Itahashi,S., Hatsuyama,Y. and Ohno,T. (1993) A mutant allele *skt5* affecting protoplast regeneration and killer toxin resistance has double mutations in its wild-type structural gene in *Saccharomyces cerevisiae*. *Biosci. Biotechnol. Biochem.*, **57**, 1391–1393.
- Kishida, M., Tokunaga, M., Katayose, Y., Yajima, H., Kawamura-Watabe, A. and Hishinuma, F. (1996) Isolation and genetic characterization of pGKL killer-insensitive mutants (iki) from *Saccharomyces cerevisiae*. *Biosci. Biotechnol. Biochem.*, **60**, 798–801.
- Kleff,S., Andrulis,E.D., Anderson,C.W. and Sternglanz,R. (1995) Identification of a gene encoding a yeast histone H4 acetyltransferase. J. Biol. Chem., **270**, 24674–24677.
- Knop,M., Siegers,K., Pereira,G., Zachariae,W., Winsor,B., Nasmyth,K. and Schiebel,E. (1999) Epitope tagging of yeast genes using a PCR-

based strategy: more tags and improved practical routines. *Yeast*, **15**, 963–972.

- Längle-Rouault,F. and Jacobs,E. (1995) A method for performing precise alterations in the yeast genome using a recyclable selectable marker. *Nucleic Acids Res.*, 23, 3079–3081.
- Leberer, E., Thomas, D.Y. and Whiteway, M. (1997) Pheromone signalling and polarized morphogenesis in yeast. *Curr. Opin. Genet. Dev.*, **7**, 59–66.
- Mumberg,D., Muller,R. and Funk,M. (1994) Regulatable promoters of Saccharomyces cerevisiae: comparison of transcriptional activity and their use for heterologous expression. Nucleic Acids Res., 22, 5767–5768.
- Mumberg,D., Muller,R. and Funk,M. (1995) Yeast vectors for the controlled expression of heterologous proteins in different genetic backgrounds. *Gene*, **156**, 119–122.
- Niedenthal,R.K., Riles,L., Johnston,M. and Hegemann,J.H. (1996) Green fluorescent protein as a marker for gene expression and subcellular localization in budding yeast. *Yeast*, **12**, 773–786.
- Otero, G., Fellows, J., Li, Y., de Bizemont, T., Dirac, A.M., Gustafsson, C.M., Erdjument-Bromage, H., Tempst, P. and Svejstrup, J.Q. (1999) Elongator, a multisubunit component of a novel RNA polymerase II holoenzyme for transcriptional elongation. *Mol. Cell*, **3**, 109–118.
- Reifsnyder, C., Lowell, J., Clarke, A. and Pillus, L. (1996) Yeast SAS silencing genes and human genes associated with AML and HIV-1 Tat interactions are homologous with acetyltransferases. *Nature Genet.*, **14**, 42–49.
- Ross-Macdonald, P., Sheehan, A., Roeder, G.S. and Snyder, M. (1997) A multipurpose transposon system for analyzing protein production, localization, and function in *Saccharomyces cerevisiae*. *Proc. Natl Acad. Sci. USA*, 94, 190–195.
- Ross-Macdonald,P., Sheehan,A., Friddle,C., Roeder,G.S. and Snyder,M. (1998) Transposon tagging I: A novel system for monitoring protein production, function and localization. *Methods Microbiol.*, 26, 161–179.
- Sambrook, J., Fritsch, E.F. and Maniatis, T. (1989) Molecular Cloning: A Laboratory Manual. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
- Schaffrath, R. and Breunig, K.D. (2000) Genetics and molecular physiology of the yeast *Kluyveromyces lactis*. *Fungal Genet. Biol.*, **30**, 173–190.
- Schaffrath, R. and Meacock, P.A. (1996) A cytoplasmic gene-shuffle system in *Kluyveromyces lactis*: use of epitope tagging to detect a killer plasmid-encoded gene product. *Mol. Microbiol.*, **19**, 545–554.
- Schaffrath,R., Stark,M.J.R. and Struhl,K. (1997) Toxin-mediated cell cycle arrest in yeast: the killer phenomenon of *Kluyveromyces lactis*. *BIOforum Int.*, 1, 83–85.
- Schmitt,M.J., Klavehn,P., Wang,J., Schonig,I. and Tipper,D.J. (1996) Cell cycle studies on the mode of action of yeast K28 killer toxin. *Microbiology*, **142**, 2655–2662.
- Sherman, F. (1991) Getting started with yeast. *Methods Enzymol.*, **194**, 3–21.
- Stark,M.J. and Boyd,A. (1986) The killer toxin of *Kluyveromyces lactis*: characterization of the toxin subunits and identification of the genes which encode them. *EMBO J.*, 5, 1995–2002.
- Stark, M.J., Boyd, A., Mileham, A.J. and Romanos, M.A. (1990) The plasmid-encoded killer system of *Kluyveromyces lactis*: a review. *Yeast*, **6**, 1–29.
- Sugisaki,Y., Gunge,N., Sakaguchi,K. and Tamura,G. (1983) *Kluyveromyces lactis* killer toxin inhibits adenylate cyclase of sensitive yeast cells. *Nature*, **304**, 464–466.
- Takita,M.A. and Castilho-Valavicius,B. (1993) Absence of cell wall chitin in *Saccharomyces cerevisiae* leads to resistance to *Kluyveromyces lactis* killer toxin. *Yeast*, **9**, 589–598.
- Tokunaga, M., Kawamura, A. and Hishinuma, F. (1989) Expression of pGKL killer 28K subunit in *Saccharomyces cerevisiae*: identification of 28K subunit as a killer protein. *Nucleic Acids Res.*, **17**, 3435–3446.
- Travers, A. (1999) Chromatin modification by DNA tracking. *Proc. Natl Acad. Sci. USA*, **96**, 13634–13637.
- Wach,A., Brachat,A., Alberti-Segui,C., Rebischung,C. and Philippsen,P. (1997) Heterologous *HIS3* marker and GFP reporter modules for PCRtargeting in *Saccharomyces cerevisiae*. Yeast, 13, 1065–1075.
- Wada, T. *et al.* (1998) DSIF, a novel transcription elongation factor that regulates RNA polymerase II processivity, is composed of human Spt4 and Spt5 homologs. *Genes Dev.*, **12**, 343–356.
- White, J.H., Butler, A.R. and Stark, M.J.R. (1989) Kluyveromyces lactis toxin does not inhibit yeast adenylyl cyclase. Nature, 341, 666–668.

- Wickner, R.B. (1996) Prions and RNA viruses of *Saccharomyces* cerevisiae. Annu. Rev. Genet., **30**, 109–139.
- Wittschieben, B.O. *et al.* (1999) A novel histone acetyltransferase is an integral subunit of elongating RNA polymerase II holoenzyme. *Mol. Cell*, **4**, 123–128.
- Wittschieben, B.O., Fellows, J., Du, W., Stillman, D.J. and Svejstrup, J.Q. (2000) Overlapping roles for the histone acetyltransferase activities of SAGA and elongator *in vivo*. *EMBO J.*, **19**, 3060–3068.
- Yajima,H., Tokunaga,M., Nakayama-Murayama,A. and Hishinuma,F. (1997) Characterization of *IK11* and *IK13* genes conferring pGKL killer sensitivity on *Saccharomyces cerevisiae*. *Biosci. Biotechnol. Biochem.*, **61**, 704–709.
- Zachariae, W., Shin, T.H., Galova, M., Obermaier, B. and Nasmyth, K. (1996) Identification of subunits of the anaphase-promoting complex of *Saccharomyces cerevisiae*. Science, 274, 1201–1204.
- Zhang,Y.P., Chen,X.J., Li,Y.Y. and Fukuhara,H. (1992) *LEU2* gene homolog in *Kluyveromyces lactis. Yeast*, **8**, 801–804.

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