Expression of fatty-acid-modifying enzymes in the halotolerant black yeast

*Aureobasidium pullulans* (de Bary) G. Arnaud under salt stress

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Abstract: Multiple tolerance to stressful environmental conditions of the black, yeast-like fungus *Aureobasidium pullulans* is achieved through different adaptations, among which there is the restructuring of the lipid composition of their membranes. Here, we describe three novel genes encoding fatty-acid-modifying enzymes in *A. pullulans*, along with the levels of their mRNAs under different salinity conditions. High levels of Δ9-desaturase and Δ12-desaturase mRNAs were seen at high salinities, which were consistent with an increased desaturation of the fatty acids in the cell membranes. Elevated levels of elongase mRNA were also detected. Surprisingly, increases in the levels of these mRNAs were also seen following hypo-osmotic shock, while hyperosmotic shock had exactly the opposite effect, demonstrating that data that are obtained from up-shift and down-shift salinity studies should be interpreted with caution.

Key words: *Aureobasidium pullulans*, desaturase, elongase, extremotolerance, halotolerance, salt stress.

INTRODUCTION

The salinisation of irrigated land has become a major agricultural problem, while attempts to breed crops with increased salt tolerance have still not yielded satisfactory results. Studies of the basic adaptation mechanisms of halophilic and halotolerant organisms, and particularly of eukaryotic ones, should help in the breeding of such crops in the future.

*Aureobasidium pullulans* (de Bary) G. Arnaud (*Dothideaceae, Ascomycota*) is a ubiquitous, saprophytic, black, yeast-like fungus. It is often found in the phyllosphere (Andrews *et al.* 1994), the air (Lugauskas *et al.* 2003) and many other, often extreme, environments, such as hypersaline water in solar salters around the World (Gunde-Cimerman *et al.* 2000) and Arctic glacier ice (Gunde-Cimerman *et al.* 2003). *A. pullulans* is of great biotechnological interest because of its production of extracelluar pullulan (Leathers *et al.* 2003), and is also a well known pathogen that can cause a variety of localised, and rarely even, systemic infections (Hawkes *et al.* 2005). *A. pullulans* thrives under many different environmental conditions, and it can tolerate a variety of environmental stresses. It can grow from 4 °C up to 35 °C, and although it thrives best without NaCl, it can tolerate up to 17 % NaCl in its growth medium; it even shows an upward shift in its salinity optimum at high temperatures (Torzilli *et al.* 1985, Zalar *et al.* 1999, Gostinčar unpublished data).

Cell membranes have crucial roles in the adaptation of any organism to environmental extremes, such as low temperature and high salinity. The active restructuring of the membrane lipid composition that can occur in response to environmental changes preserves the suitable dynamic state of the membrane bilayer and restores membrane function following environmental insult (Hazel & Williams 1990). This restructuring of the membrane lipid composition in response to environmental cues can occur either through changes in the products of cell lipid biosynthesis, or through the selective degradation of lipid species with inappropriate properties (Hazel & Williams 1990).

Such cell membrane changes have been particularly well studied in salt-sensitive *Saccharomyces cerevisiae*. As in other eukaryotes, the *de-novo* biosynthesis of saturated fatty acids in these yeast requires acetyl-CoA carboxylase and the fatty-acid-synthase complex. Yeast are only able to synthesise mono-unsaturated fatty acids containing a Δ9 double bond, using a desaturase that is encoded by the OLE1 gene, the expression of which is severely repressed by unsaturated fatty acids (Trotter 2001). Regulation of this gene's expression occurs at the level of OLE1 transcription and via mRNA stability (Gonzalez & Martin 1996). In *S. cerevisiae*, several different enzyme systems have been described that can elongate the fatty-acetyl-CoAs that are formed from *de-novo* synthesis or are acquired from the growth medium, some of which are essential for cell viability (Trotter 2001).

The halotolerant *A. pullulans* has so far been studied only at the level of its membrane composition and fluidity when exposed to hypersaline conditions (Turk *et al.* 2004, 2007). The levels of unsaturated fatty acids in the cell membranes of *A. pullulans* cells grown in 5 % and 10 % NaCl increase significantly due to their enrichment in C18:2Δ9,12 fatty acids. Significant changes in the lengths of the fatty acids were not seen, although such changes have been detected in another black, yeast-like fungus, the halophilic *Hortaea werneckii* (*Horta*) Nishim. & Miyaji (*Turk et al.* 2004). Although *A. pullulans* is more sensitive to salt than *H. werneckii* (but less so than *S. cerevisiae*), it is of interest because of its multiple tolerances to many other types of stress.

The aim of the present study was to find desaturase and elongase genes that are involved in these fatty-acid modifications in *A. pullulans*, to compare them to other known homologous enzymes from different fungi, and to study their differential expression under saline and non-saline conditions. In particular,
we wanted to determine whether these changes in the fatty-acid saturation levels can be explained by the differential expression of the genes encoding desaturases, and whether the expression of the elongase gene changes in cells subjected to salt stress.

**MATERIALS AND METHODS**

**Strain and culture conditions**

The EX-F150 strain of *A. pullulans* was originally isolated from the Sečovlje solar salterns (Slovenia) (Gunde-Cimerman et al. 2000) and has been maintained in the Extremophilic Fungi (Ex Culture Collection of the Department of Biology, Biotechnical Faculty, University of Ljubljana, Slovenia), and as MZKI 8-602 in the Microbial Culture Collection of the National Institute of Chemistry (Slovenia). *A. pullulans* was grown using the standard Yeast Nitrogen Base (YNB) chemically defined liquid medium, both without NaCl and with different concentrations of NaCl (2.5 %, 5.0 %, 7.5 %, 10.0 % and 13.0 %; w/v). The liquid cultures were grown at 28 °C and under constant shaking (180 rpm).

**DNA and RNA isolation**

For DNA isolation, *A. pullulans* was grown in YNB, and the biomass was harvested by centrifugation in the mid-exponential growth phase (OD_{600} = 0.8–1.0). The cell pellet was frozen in liquid nitrogen and homogenised using a mortar and pestle. The DNA was then isolated according to the protocol described by Rozman & Kome (1994).

For RNA isolation, *A. pullulans* was cultured in YNB with different concentrations of NaCl; the biomass was harvested by centrifugation in the mid-exponential growth phase (OD_{600} = 0.8–1.0). The cells that were subjected to osmotic shock were initially grown in YNB without NaCl or with 10 % NaCl (w/v). These cells were also harvested in the mid-exponential growth phase, and then subjected to osmotic shock by resuspension in YNB with 10 % NaCl or without NaCl, respectively (at the same pH and temperature) for specific times (5–120 min); they were then harvested by filtration. Adapted cells that were grown constantly at the final concentration of NaCl were used as an additional end-point reference. The biomass was frozen in liquid nitrogen and stored at −80 °C until further analysis, when they were homogenised in liquid nitrogen using a mortar and pestle. The RNA was isolated using TRI REAGENT™ (Sigma), according to the manufacturer instructions. Possible contaminating DNA was degraded with deoxyribonuclease I (Fermentas). The integrity and purity of the RNA was evaluated spectrophotometrically and with formaldehyde agarose electrophoresis.

**Amplification and sequencing of genes**

Partial sequences of the genes were amplified by polymerase chain reaction (PCR), which was performed as described by Lanišnik Ržner et al. (1999). In all cases, the annealing temperatures were 60 °C. Non-degenerate oligonucleotide primers were constructed on the basis of highly conserved domains in known fungal desaturases: 5'-TAC GAT ACC GAC AAG GCC CCC TA-3' and 5'-GGA ACT CGT GTG GGA GTG TCT A-3' for \( \Delta^9 \)-desaturase, and 5'-CCATCA GAG GAT TGT CCC TCA AAG T-3' and 5'-ATG TTA CCA GTG GCC TTG TGG TGC T-3' for \( \Delta^{12} \)-desaturase. Primers specific for part of the elongase gene were designed using the CODEHOP algorithm (Rose et al. 1998), from protein sequences of known fungal elongases: 5'-GTC ATC TAC ATC ATC ATT TTY GGN GG-3' and 5'-CGG GCC GAC TGG TGG TGG AAC TGG TAY TAC AT-3'. The amplified fragments were recovered from electrophoresis gels using Perfectprep Gel Cleanup (Eppendorf), and then sequenced. Fragments of the elongase gene were cloned into an Escherichia coli plasmid (pGEM™-T Vector System, Promega) prior to sequencing. The same regions were also amplified using cDNA as a template, to confirm the existence of introns indicated by alignment with sequences of homologous genes in the GenBank database. New non-degenerate primers were designed and later used in RT-PCR reactions: 5'-ATC TCC GAC CTC ACG ACG AC-3' and 5'-CTC ACC GAG AGT GAC GAT GG-3 for \( \Delta^3 \)-desaturase gene; 5'-CGG AGA TAC ATT CCC TCG AC-3' and 5'-CCA TGA GAA GTA AGG GAC AAG G-3' for \( \Delta^{12} \)-desaturase gene; and 5'-TGG TAG GTG TGG TGG AAA GC-3' and 5'-CAT ATT TGG CGG CAG AGA GT-3' for the elongase gene.

Southern blotting was performed as described previously (Turk & Plemenits 2002). DNA fragments were amplified as described above (using non-degenerate primers in the case of the elongase gene), and then radioactively labelled and used as probes. The upstream and downstream sequences of all three of these genes were obtained using the GenomeWalker™ Universal Kit (Clontech), according to the manufacturer instructions, using oligonucleotide primers that were specific for the adapters supplied in the kit, and oligonucleotide primers specific for known fragments of the genes. These were as follows: *ApoLE1* upstream 5'-GCC AAG GAG AGT GAC GAT GG-3' and 5'-ATC TCC GAC CTC ACG ACG AC-3' and 5'-CTC ACC CTC AAC GAC AC-3', downstream nested 5'-CGT CAC TCT CCG TGA GGA CTA CCA CAA CT-3'; *ApoODE1* upstream 5'-GTT TCG TTG GAG GTC GTG ATG TGG GAG GAA AA-3', upstream nested 5'-GCT TGT CCG GCC AGA CCA CAA GAA CAA ATG TGA TG-3', downstream nested 5'-CAT CAC ACT TCG TTC GCC CAA CAA CT-3'; *ApELO1* upstream 5'-GAG GTG CAT GAG GTA GAG GCC AGC ACG TCT AA-3', downstream nested 5'-GGC TAC ACC GCA CA-3', downstream nested 5'-ATG TTA CCA CGG CAA GCA CCA CAA CT-3'; 5'-ACT GGC TCG GTG ACC AGC CTT TCG AT-3'; downstream nested 5'-GCA TGA ACT TTC TGC GCC CAA CAA ATG TGA TG-3', downstream nested 5'-CAT CCG AGT CTC ATG GTT GGG CCC CAT TA-3', downstream nested 5'-TGA ACT TCG TTC ACC ACG TCG TCA TGT ACT-3'.

**Gene phylogeny reconstruction**

Homologues of the \( \Delta^3 \)-desaturase and \( \Delta^{12} \)-desaturase and the elongase were identified by BLAST searches (Altschul et al. 1997) against a GenBank non-redundant protein database. In addition to fungal homologues, sequences from other species were added. Protein sequences were aligned using ClustalX (Thompson et al. 1997) and edited in BioEdit software (Hall 1999). Gene trees were generated with MrBayes software applying Bayesian inference (Huelsenbeck & Ronquist 2001, Ronquist & Huelsenbeck 2003). Runs were performed for two million generations with mixed amino-acid models, default temperature and numbers of chains. The trees were sampled every 10 generations. Trees sampled before the analysis reached stationarity of likelihood values and those sampled before the average standard deviation of the split frequencies lowered under 0.5 % were excluded from the final analysis. The stationarity of likelihood values was checked using the Tracer software (Rambaut & Drummond 2007). Enzymes from non-fungal organisms were used as outgroups: *Vibrio fischeri* (\( \Delta^3 \)-desaturases) and *Arabidopsis thaliana* (\( \Delta^{12} \)-desaturases and elongases).
RT-PCR and statistical analysis

Total cDNA was synthesised from the RNA samples isolated as described above, using the RevertAid™ H Minus First Strand cDNA Synthesis Kit (Fermentas). Approximately 12.5 ng of the synthesised cDNA was used as a template for multiplex PCR with each pair of the oligonucleotide primers described above and oligonucleotide primers specific for the 28S rRNA (Lanišnik Rižner et al. 1999). The PCR consisted of 21, 24 and 22 cycles for the Δ⁸-desaturase and Δ¹²-desaturase and for the elongase, respectively. The primer dropping method was used (Wong et al. 1994), with 28S primers added to the reaction for the last 16 PCR cycles. The products of the reactions were separated on agarose gels and stained with ethidium bromide, with the relative abundance of each amplified fragment evaluated by measuring its luminescence with the UN-SCAN-IT software (v. 5.1, Silk Scientific Corporation). The values of given genes were established through comparison of these values between different samples. RT-PCR analyses were carried out in triplicate.

Means and standard deviations were calculated, and the variance due to systematic error was subtracted. The data were analysed for statistically significant differences using a one-way ANOVA test (the variance was checked for homogeneity), followed by the Tukey (HSD) test.

RESULTS

Specific products were successfully amplified using PCR with the oligonucleotide primers specific for parts of the genes encoding a Δ⁸-desaturase, a Δ¹²-desaturase and a fatty-acid elongase. Sequencing of the fragments produced 306 bp, 234 bp and 548 bp partial sequences. Using the GenomeWalker™ Universal Kit, complete coding sequences of three new genes from A. pullulans were obtained. Searches with the BLAST programme (Altschul et al. 1997) showed similarities with several genes for Δ⁸-desaturases and Δ¹²-desaturases and for fatty acid elongases. The genes were named ApOLE1, ApODE12 and ApELO1, respectively. The sequences have been stored in GenBank under the accession numbers DQ901954 (ApOLE1), DQ901955 (ApODE12) and EF123104 (ApELO1).

The coding regions were determined through the alignment of the sequences with known homologous genes obtained from the GenBank database and with cDNA sequences. The deduced ApOle1 protein is 458 amino acids long and the gene contains a 50-bp intron. ApODE12 contains no introns, while the putative ApOde12 protein is of 480 amino acids. Two 52 and 53 bp introns were discovered in the ApELO1 fragment, and their sequences were found to be 50 % identical. The hypothetical ApElo1 protein is of 343 amino acids. Southern blotting of the genomic DNA did not suggest the existence of more than one copy of any of the three genes in the genome of A. pullulans (Fig. 1).

Good convergence of the runs was reached when constructing all three of the gene trees with MrBayes. The likelihood values reached plateaus after approximately 10,000 (Δ¹²-desaturases), 8,000 (Δ²-desaturases) and 6,000 generations (elongases), while the average standard deviations of the split frequencies dropped below 0.5 % after approximately 400,000 (Δ²-desaturases) and 800,000 (Δ¹²-desaturases and elongases) generations. The first 4,000 (Δ¹²-desaturases) and 8,000 (Δ¹²-desaturases and elongases) trees were discarded as burn-in. The posterior probabilities for the amino-acid models were 1 for the Blossom262 model (Henikoff & Henikoff 1992) for Δ¹²-desaturases and Δ²-desaturases, and 1 for the WAG model (Whelan & Goldman 2001) for elongases. All three of the deduced proteins (ApOle1, ApOde12 and ApElo1) clustered with homologous enzymes from fungi belonging to Pezizomycotina.

The relative abundances of the ApOLE1, ApODE12 and ApELO1 genes were studied by RT-PCR. Their profiles for growth at different salinities, and under hyper- and hypo-osmotic shock were analysed separately. One-way ANOVA (α = 0.05) showed significant differences (3.13 ×10⁻⁷< P <0.003) in all cases. The results of the Tukey (HSD) post-hoc testing are shown in Figures 2–4.

The abundance of ApOLE1 mRNA (Fig. 5) was significantly higher at 13 % NaCl than at the lower salinities. When A. pullulans
Fig. 2. Phylogeny of the Δ⁹-desaturase enzymes. The tree was constructed from the alignment of protein sequences found in the GenBank database with the BLAST programme. Analyses were performed with MrBayes software in two runs with two million generations each. Mixed amino-acid models were used, and the first 20% of the trees were excluded from the final consensus tree. Full scientific names of source organisms are followed by gene names (if available) and GenBank accession numbers. Entries marked with an asterisk (*) are available in the database as hypothetical proteins without any assigned functions.

Cells were subjected to osmotic shock, the responses differed considerably. Surprisingly, after hyperosmotic shock, the ApOLE1 mRNA level was reduced, although a significant decrease was detected in only one sample (1 h after the up-shift). All other levels, except that initial prior to up-shift, were significantly lower than the final steady-state levels at 10 % NaCl. In contrast, hypo-osmotic shock led to increased levels of ApOLE1 mRNA. A significant difference was first detected 10 min after the down-shift, and it reached its peak after 30 min. One hour after the down-shift, the ApOLE1 mRNA decreased again, but two hours after the down-shift it was still significantly higher than prior to the osmotic shock.

Nevertheless, the steady-state mRNA levels in the cells not exposed to NaCl was significantly lower than in all of the samples obtained after the hypo-osmotic shock.

The levels of ApODE12 mRNA were generally higher at higher salinities (Fig. 6): the levels were significantly higher at 13 % NaCl as compared to 0.0 %, 2.5 % and 5.0 % NaCl. As seen for the ApOLE1 gene, the levels of ApODE12 mRNA decreased significantly after exposing the cells to hyperosmotic shock. A transient increase in ApODE12 mRNA levels was seen 10 min after the up-shift, which was significantly higher than after 30 min, and 1 h and 2 h later. Hypo-osmotic shock resulted in transiently increased ApODE12 mRNA levels. The peak here was reached 30 min after the down-shift. A significant increase was first seen 10 min after exposing the cells to the shock and 2 h later it could not be detected any more.

The levels of ApELO1 mRNA were highest at 13.0 % NaCl (Fig. 7), and at 7.5 % and 10.0 % NaCl they were also significantly higher than at both 0.0 % and 2.5 % NaCl. After exposing the cells to hyperosmotic shock, the ApELO1 mRNA levels decreased after 5 min, and the decrease remained significant throughout the shock period, even though the steady-state ApELO1 mRNA expression levels at 10.0 % NaCl were significantly higher than all others seen during the up-shift. Hypo-osmotic shock resulted in a significant increase in ApELO1 mRNA levels after 10 min, which then decreased slowly up to 1 h after the shock, when it was still significantly higher than prior to the down-shift.
Fig. 3. Phylogeny of the Δ₁₂-, Δ₁₅-, and ω₃-desaturase enzymes. The tree was constructed from the alignment of the protein sequences found in the GenBank database with the BLAST programme. Analyses were performed with MrBayes software in two runs with two million generations each. Mixed amino-acid models were used, and the first 40% of the trees were excluded from the final consensus tree. Full scientific names of source organisms are followed by the GenBank accession numbers. The presumed functions of the enzymes are marked with full circles ● (Δ₁₂ activity), empty circles ○ (ω₃ activity), squares ■ (Δ₁₅ activity), or combinations thereof. Entries marked with an asterisk (*) are available in the database as hypothetical proteins without assigned function. Presumed large duplications of the ancestor genes are marked with arrows and dashed lines. Other gene duplicates are connected with dotted lines.

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Fig. 4. Phylogeny of elongase enzymes. The tree was constructed from the alignment of the protein sequences found in the GenBank database with the Blast programme. Analyses were performed with MrBayes software in two runs with two millions generations each. A mixed amino-acid model was used, with the first 40 % of trees were excluded from the final consensus tree. Full scientific names of source organisms are followed by gene names (if available) and GenBank accession numbers. Entries marked with an asterisk (*) are available in the database as hypothetical proteins without assigned functions. Presumable large duplication of the ancestor gene is marked with arrows and a dashed line. Other gene duplicates are connected with dotted lines.
Fa t t y-a c i d-M o d iFy i n g e n z yM e s i n A. p u l u l a n s u n d e r S a l t S t r eS S

Fig. 5. ApOLE1 mRNA abundance. A. Semi-quantitative transcriptional analysis by RT-PCR. Samples were prepared from cells grown at different salinities (first column) or subjected to a salinity shift from 0 % to 10 % NaCl (second column) and 10 % to 0 % NaCl (third column). mRNA abundances in cells completely adapted to the final salinity (10 % in the case of an up-shift, and 0 % in the case of a down-shift) are marked with an asterisk (*). Amplified fragments of ApOLE1 and 28S rRNA genes were run on 1 % agarose gel and stained with ethidium bromide. B. ApOLE1 gene expression as detected by RT-PCR. Data represent means ±SD of three independent experiments. Significant differences (Tukey’s HSD) were seen between the value marked with an arrow-head and each of the samples marked with the dot.

Fig. 6. ApODE12 mRNA abundance. A. Semi-quantitative transcriptional analysis by RT-PCR. Samples were prepared from cells grown at different salinities (first column) or subjected to salinity shifts from 0 % to 10 % NaCl (second column) and 10 % to 0 % NaCl (third column). mRNA abundances in cells completely adapted to the final salinity (10 % in case of up-shift and 0 % in case of down-shift) are marked with an asterisk (*). Amplified fragments of ApODE1 and the 28S rRNA genes were run on 1 % agarose gel and stained with ethidium bromide. B. ApODE12 gene expression as detected by RT-PCR. Data represent means ±SD of three independent experiments. Significant differences (Tukey’s HSD) were seen between the value marked with an arrow-head and each of the samples marked with dots.

Fig. 7. ApELO1 mRNA abundance. A. Semi-quantitative transcriptional analysis by RT-PCR. The samples were prepared from cells grown at different salinities (first column) or subjected to salinity shifts from 0 % to 10 % NaCl (second column) and 10 % to 0 % NaCl (third column). mRNA abundances in the cells were completely adapted to the final salinities (10 % in the case of up-shift, and 0 % in the case of down-shift) are marked with an asterisk (*). Amplified fragments of ApELO1 and the 28S rRNA genes were run on 1 % agarose gels and stained with ethidium bromide. B. ApELO1 gene expression as detected by RT-PCR. Data represent means ±SD of three independent experiments. Significant differences (Tukey’s HSD) were seen between the value marked with an arrow-head and each of the samples marked with the dot.
DISCUSSION

The cell membranes of \textit{A. pullulans} contain long-chain fatty acids with double bonds in positions 9 and 12 ($\Delta^9$, $\Delta^{12}$; Turk et al. 2004). We thus assumed that the \textit{A. pullulans} genome must encode the corresponding fatty-acid desaturases as well as a fatty-acid elongase. We also wanted to know if changes in the mRNAs levels of these desaturases and elongases correlate with alterations in fatty-acid composition that have previously been seen at different salinities of the surrounding medium (Turk et al. 2004).

The proteins coded by the new genes that are involved in fatty-acid modifications in \textit{A. pullulans} did not differ significantly from their homologues in other fungi. Analysis of the ApOLE1 sequence revealed the presence of an intron with atypical splice acceptor bases at its 3’ end (TG instead of AG). The high degree of identity between two introns in the ApELO1 gene indicated that they might originate from a relatively recent duplication event and could thus be useful in studies of the evolutionary history of this species.

The essential OLE1 gene was first found in \textit{S. cerevisiae}, and later also in several other fungi (Watanabe et al. 2004). A homologue in \textit{A. pullulans} was therefore expected. The $\Delta^{12}$-desaturases are, on the other hand, less common, and have not been extensively studied (Watanabe et al. 2004). They introduce a cis double bond into fatty acids that already have a cis double bond at the $\Delta^9$ position. This additional double bond further increases the membrane fluidity, although its effect is less pronounced than the effect of the first double bond (Los & Murata 1998). Therefore, this $\Delta^{12}$-desaturase in \textit{A. pullulans} may enable the cell to fine-tune its cell-membrane fluidity and may be one of the reasons for its halotolerance, in contrast to the salt-sensitive \textit{S. cerevisiae} which is not capable of $\Delta^{12}$-desaturation. Like the $\Delta^9$-desaturases, elongases are essential for cell growth (Dittrich et al. 1998). Three different elongases exist in \textit{S. cerevisiae}, each with a different substrate specificity (Trotter 2001). So far, only one elongase homologue has been identified in \textit{A. pullulans}, using degenerate PCR and Southern blotting.

The evolutionary origins of all three of these enzymes from \textit{A. pullulans} in the Pezizomycotina branch as shown by our phylogenetic analysis are not surprising.

Evolution of $\Delta^9$-desaturases also did not show any unusual characteristics with the exception of desaturases from \textit{Mucoromycotina} which clustered with homologous enzymes from Basidiomycota.

Comparisons of the deduced amino-acid sequences of the $\Delta^{12}$-desaturases and $\omega^3$-desaturases indicate several independent duplications of their genes, leading to separate $\Delta^9$ and $\omega^3$ branches; this was previously reported by Damude et al. (2006). According to our analysis, the major duplication occurred before the separation of Pezizomycotina from Saccharomycotina, resulting in a separate $\omega^3$ branch with very few representatives. At least one duplication event could also be detected in \textit{Saccharomycotina} and \textit{Mucoromycotina}.

The phylogenetic analysis of the elongase genes indicates a duplication event early in the evolutionary history of \textit{Saccharomycotina}, which led to two groups of elongase genes, each represented with (at least) one enzyme in most species. Surprisingly, Eto1 and Eto2 from \textit{S. cerevisiae} belong to the same group, and Eto3 to the other one, although Eto2 and Eto3 are more functionally related. Two hypothetical proteins from \textit{Aspergillus oryzae} indicate a possible duplication event in \textit{Pezizomycotina} as well.

Another interesting elongase is a long-chain polyunsaturated fatty acid elongation enzyme from \textit{Mortierella alpina} described by Parker-Barnes et al. (2000), which proved to be quite different from all of the other elongases, and in our analysis it clustered with animal elongase enzymes.

The structural and functional integrity of biological membranes requires the presence of water. Changes in cellular water activities can have profound influences on membrane stability. Cations from salts interact with membrane constituents and can also affect their conformation. Furthermore, abrupt changes in salinity can expose cell membranes to physical stress because of the changes in osmotic pressure (Hazel & Williams 1990). Phenotypic adjustments in the cell-membrane lipid compositions as a response to altered salinity have been observed, among which there are increases in the relative proportions of anionic lipids. Increased levels of unsaturated acyl chains and longer fatty acids have been seen in yeasts and cyanobacteria at higher salinities (Hazel & Williams 1990, Turk et al. 2004), and there is evidence that these changes serve to increase the membrane fluidity at high salt concentrations (Russell 1989, Hazel & Williams 1990).

The high abundance of the desaturase mRNAs seen at high salinities in \textit{A. pullulans} was expected, and it is consistent with an enrichment in the C18:2$\Delta^9,12$ fatty acids in the \textit{A. pullulans} cell membranes at 5 % and 10 % NaCl (Turk et al. 2004). These changes may help to sustain sufficient levels of membrane fluidity at high salt concentrations (Russell 1989, Hazel & Williams 1990). Increased proportions of long chain fatty acids would oppose this effect, but when the \textit{A. pullulans} cell membranes were analysed, no shifts in fatty-acid lengths were detected (Turk et al. 2004). A higher abundance of \textit{ApELO1} mRNA at high salinities might not therefore be associated with alterations in the fatty-acid composition. The reason for this discrepancy might lie in a faster turnover of the long-chain fatty-acid pool at high salinities, a lower efficiency of \textit{ApELO1} due to the high concentrations of compatible solutes, additional undetected elongases with complementary roles, or the postranscriptional control of expression.

The changes in the mRNA levels following osmotic shock were contrary to expectations, considering the observations of the mRNA abundances at different salinities. A similar transient decrease in mRNA abundance after exposure of the cells to hyperosmotic shock was seen previously for the \textit{S. cerevisiae} OLE1 gene. Expression of the \textit{OLE1} gene diminished 45 min after a shift to high osmolarity (Rep et al. 2000), and increased again after 90 min (Yale & Bohnert 2001). In both cases, the changes can probably be associated with the temporary growth arrest caused by the shock, a response that is similar to the previously reported expression of ribosomal proteins (Rep et al. 2000).

Another possibility is that these reactions act as part of a stress-response system in \textit{A. pullulans}. By introducing double bonds into these phospholipid fatty-acyl chains and by decreasing the fatty-acyl chain length in artificial membranes, glycerol permeability is increased (Blomberg & Adler 1992). The immediate changes following the osmotic shock could thus be an emergency response, which in the case of hyperosmotic shock protects the cells from compatible solute leakage until other – possibly slower or energetically more demanding – mechanisms can take over, such as synthesis of other compatible solutes and membrane glycerol transporters, or cell wall melanisation. Indeed, in salt-sensitive \textit{S. cerevisiae}, the ORF’s encoding many known yeast salinity stress response proteins are either unaffected or down-regulated immediately following the up-shift, and gradually induced only later (Yale & Bohnert 2001). Similarly, a decrease in the saturation levels of the fatty acids, and therefore an increase in the permeability of the membranes for molecules such as glycerol could facilitate export of compatible solutes when they are no longer needed.
(Tamas et al. 1999), explaining the transient increases in the levels of the mRNAs encoding fatty acid desaturases.

Finally, many cells deal with abrupt changes in osmotic pressure caused by changes in salinity in a very different way than when they are grown in an environment with constant water activity with this "turgor shock" response somehow having priority over the "salinity shock" response.

The opposite response patterns in mRNA levels following hyper- and hypo-osmotic shock compared to the slow adaptation to growth at different salinities indicates that these two types of stress represent a fundamentally different challenge to cells.

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