Characterization of the transactivation and nuclear localization functions of *Pichia pastoris* zinc finger transcription factor Mxr1p

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The zinc finger transcription factor Mxr1p regulates the transcription of genes involved in methanol, acetate, and amino acid metabolism of the industrial yeast *Pichia pastoris* (a.k.a. *Komagataella phaffii*) by binding to Mxr1p response elements in their promoters. Here, we demonstrate that Mxr1p is a key regulator of ethanol metabolism as well. Using transcriptomic analysis, we identified target genes of Mxr1p that mediate ethanol metabolism, including *ALD6*-1 encoding an aldehyde dehydrogenase. *ALD6*-1 is essential for ethanol metabolism, and the *ALD6*-1 promoter harbors three Mxr1p response elements to which Mxr1p binds *in vitro* and activates transcription *in vivo*. We show that a nine-amino-acid transactivation domain located between amino acids 365 and 373 of Mxr1p is essential for the transactivation of *ALD6*-1 to facilitate ethanol metabolism. Mxr1N250, containing the N-terminal 250 amino acids of Mxr1p, localized to the nucleus of cells metabolizing ethanol dependent on basic amino acid residues present between amino acids 75 and 85. While the N-terminal 400 amino acids of Mxr1p are sufficient for the activation of target genes essential for ethanol metabolism, the region between amino acids 401 and 1155 was also required for the regulation of genes essential for methanol metabolism. Finally, we identified several novel genes whose expression is differentially regulated by Mxr1p during methanol metabolism by DNA microarray. This study demonstrates that Mxr1p is a key regulator of ethanol metabolism and provides new insights into the mechanism by which Mxr1p functions as a global regulator of multiple metabolic pathways of *P. pastoris*.

**Pichia pastoris** (a.k.a. *Komagataella phaffii*) is a methylotrophic yeast capable of utilizing methanol as a sole carbon source. The methanol utilization pathway enzyme alcohol oxidase, encoded by *AOX1*, constitutes 30% of the total soluble protein. This strong induction under methanol conditions has led to the development of a methanol-inducible *AOX1*-based promoter for commercial production of heterologous proteins (1, 2). It is a respiratory yeast, suggesting that in high glucose conditions biomass production through tricarboxylic acid cycle is favored over the conventional ethanol generation route followed in other yeasts like *Saccharomyces cerevisiae* because of the increased glycolytic flux. Consequently, the high biomass generation in the bioreactors dramatically increases the product yields, making them industrially more relevant (3). Another remarkable feature of this yeast is the ability to utilize a plethora of carbon compounds as the sole source of carbon. These include glucose, glycerol, ethanol, acetic acid, oleic acid, sorbitol, and amino acids such as glutamate. The genes encoding key enzymes of these metabolic pathways are primarily regulated at the transcriptional level (4, 5). In the case of methanol metabolism, several transcriptional regulators, such as Mxr1p, Trm1p, Mit1p, Rop1p, Mig1p, Mig2p, and Nrg1p, regulate the transcription of *AOX1* gene encoding alcohol oxidase 1 (6–11). Mxr1p regulates the expression of genes of acetate and amino acid metabolism as well and thus functions as a global regulator of central carbon metabolism (4, 5). While Mxr1p primarily functions as a transcriptional activator, it is known to repress the expression of *GT1* encoding glycerol transporter during glycerol metabolism (12).

Transcriptional activators bind to DNA through their DNA-binding domain (DBD) and activate transcription by interacting with proteins of preinitiation complex, either directly or via coactivators through their transactivation domains (TADs) (13, 14). DBDs consist of motifs, such as zinc fingers, helix–turn–helix motif, or basic helix–loop–helix leucine zipper, whereas the TADs consist of amphipathic alpha helices, glutamine-rich domains, or proline-rich domains (15). Nuclear localization of transcription factors is facilitated by nuclear localization signals (NLSs), and the classical NLS is characterized by the presence of monopartite or bipartite NLSs consisting of one or two clusters of basic amino acids separated by a spacer, respectively (16). *P. pastoris* Mxr1p possesses a DBD in the amino terminus consisting of two C2H2 zinc fingers, which exhibit strong homology to the DBD of *S. cerevisiae* Adr1p (6, 17, 18). Mxr1p DBD binds to Mxr1p response elements (MXREs) in the promoters of target genes bearing the consensus sequence 5′-CYCCNY-3′ (4, 5, 18, 19). A TAD enriched in phenylalanine residues present between amino acid residues 246 and 280 of Mxr1p was shown to be functional during methanol metabolism but not ethanol metabolism (20). Serine 215 residue located upstream of this TAD is phosphorylated in cells metabolizing ethanol but not methanol, and interaction of this phosphoserine with 14-3-3 protein inhibits the function of the N-terminal TAD (20).
Key functional domains of Pichia pastoris Mxr1p

Mxr1p is cytosolic in cells metabolizing glucose and localizes to the nucleus when cultured in media containing non-fermentable carbon sources (4, 6). Thus far, the NLS of Mxr1p has not been characterized.

In S. cerevisiae, ethanol is metabolized into acetaldehyde, acetate, and acetyl-CoA by the sequential action of alcohol dehydrogenase, aldehyde dehydrogenase (ALD), and acetyl-CoA synthetase (ACS), respectively. In S. cerevisiae as well as Kluyveromyces lactis, expression of alcohol dehydrogenase 2 is regulated by Adr1p, which is regarded as the homolog of Mxr1p (17, 21). However, the ability of P. pastoris Mxr1p to regulate genes of ethanol metabolism is not known. Considering the existing gaps, the present study was initiated to investigate the regulatory role of Mxr1p during ethanol metabolism, identify the NLS, and characterize the trans-activation functions. Using high-throughput genome-wide RNA-Seq, we have identified novel target genes of Mxr1p essential for ethanol metabolism. Among these, ALD6-1 is essential for ethanol metabolism. Among these, ALD6-1 encoding an ALD, which is downregulated in Δmxr1, was further characterized. We have identified an NLS and a nine-amino acid TAD (9aaTAD) in the amino terminal region of Mxr1p required for the regulation of genes of ethanol metabolism. Finally, we have identified several novel target genes of Mxr1p required for methanol metabolism using DNA microarray.

Results

Transcriptional regulation of ethanol metabolism by Mxr1p

P. pastoris strains used in this study are listed in Table 1. To gain insights into the mechanism of transcriptional regulation of genes of ethanol metabolism by Mxr1p, genome-wide RNA-Seq was carried out. GS115 and Δmxr1 were cultured in YNB, RNA was isolated, and biological replicates were subjected to RNA-Seq. The datasets have been deposited in the Gene Expression Omnibus (GEO) database under accession number GSE168677. A total of 25 to 30 million reads were obtained for each sample after quality trimming. Samples were aligned to the K. phaffii strain CBS 7435 reference genome (https://www.ncbi.nlm.nih.gov/assembly/GCA_900235035.1/). Differentially expressed genes were identified by normalizing the read counts in Δmxr1 to that of GS115 (control). Deletion of MXR1 resulted in significant changes in the transcriptome. Of 5424 genes, 170 were downregulated and 28 were upregulated in Δmxr1 (Supporting information 2). For this analysis, the threshold for statistical significance was considered an adjusted p value of <0.05 and log2 fold change of ±1 for upregulated and downregulated genes. The overall change and the most highly upregulated and downregulated genes are shown in the volcano plot and heat maps, respectively (Fig. 1, B–D). ALD6-1, encoding an ALD, was the most downregulated gene in Δmxr1 (greater than fivefold). Other genes downregulated in Δmxr1 include those encoding peroxisomal proteins, PP7435_Chr2-0527 encoding glutathione transferase involved in detoxification of hydrogen peroxide, lipid metabolism genes (POX1, FAA2, and PCD1 encoding acyl Co-A oxidase,

| Strain                        | Description                     | References |
|-------------------------------|---------------------------------|------------|
| GS115                         | his4                            | (6)        |
| Δmxr1                         | GS115, Ppmxr1A::Zeo3             | This study |
| GS115::Pmxr1A::Zeo3            |                                  | This study |
| Δmxr1-A                       | GS115, HIS4::(P_PpALD6-1-Myc)   | This study |
| Δmxr1-N250                    | GS115, HIS4::(P_PpALD6-1-Myc)   | This study |
| Δmxr1-A-N109                  | Δald6-1, HIS::(P_PpALD6-1-Myc)  | This study |
| Δmxr1-A-N150                  |                                  | This study |
| Δmxr1-A-N250                  |                                  | This study |
| Δmxr1-A-N400                  |                                  | This study |
| Δmxr1-A-N400F*                |                                  | This study |
| Δmxr1-A-N400Q*                |                                  | This study |
| Δmxr1-A-N400Q*                |                                  | This study |
| Δmxr1-A-N81                   |                                  | This study |
| Δmxr1-A-N109                  |                                  | This study |
| Δmxr1-A-N250-M2               |                                  | This study |
| Δmxr1-A-N250-M1               |                                  | This study |
Figure 1. Identification of target genes of Mxr1p during ethanol metabolism by RNA-Seq and characterization of ALD6-1. A, analysis of growth of GS115 and Δmxr1 cultured in YNBE. B, volcano plot of genes differentially regulated in GS115 and Δmxr1 cultured in YNBE. ALD6-1 is the most down-regulated gene in Δmxr1. E, schematic representation of ethanol utilization pathway of Saccharomyces cerevisiae. F, quantitation of ALD and ACS1 mRNAs in GS115 and Δmxr1 cultured in YNBE by quantitative PCR (qPCR). G, analysis of expression of ALD6-1Myc from 1.0-kb ALD6-1 promoter (P) in GS115 and Δmxr1 cultured in YNBE by Western blotting using anti-Myc antibodies. H, analysis of GFP expression from P in GS115 and Δmxr1 cultured in YNBE by Western blotting using anti-GFP antibodies. Quantitation of the data is presented. The intensity of individual bands was quantified and expressed as arbitrary units ± SD relative to controls (n = 3). I, strategy for the generation of Δald6-1 and confirmation of deletion of ALD6-1 by PCR. Gene-specific primers (arrows) amplify ALD6-1 from
long-chain fatty acyl CoA synthetase, and pyrophosphatase, respectively) and GTH1 encoding a high-affinity glucose transporter. Interestingly, ADY2-3 encoding exporter of ammonia was downregulated, whereas MEP2 encoding ammonia transporter was upregulated in Δmxr1. CRC1, CAT2, and YAT1 encoding proteins involved in carnivitate transport were highly upregulated in Δmxr1. Genes encoding acetyl CoA metabolizing enzymes, such as isocitrate lyase, isocitrate dehydrogenase, and isocitrate/isopropylmalate dehydrogenase, were also upregulated in Δmxr1 probably to facilitate metabolism of acetyl CoA via tricarboxylic acid cycle to generate energy in the absence of ALD6-1, the ethanol-metabolizing gene.

**ALD6-1 is indispensable for ethanol metabolism**

In *S. cerevisiae*, ALDs catalyze the conversion of acetaldehyde to acetate, the second step of ethanol metabolism (Fig. 1E). Since ALD6-1 is the most downregulated gene in Δmxr1 cultured in YNBE, we investigated its function and regulation in detail. *P. pastoris* genome harbors four ALD genes annotated as ALD6-1, ALD6-2, ALD4, and ALD2 (Table 2). Quantitation of the four ALD mRNAs by quantitative PCR (qPCR) indicated that ALD6-1 alone was downregulated in Δmxr1 during ethanol metabolism (Fig. 1F). Since ACS1 encoding ACS1 was shown to be downregulated in Δmxr1 during acetate metabolism (4), we examined ACS1 transcript levels in cells metabolizing ethanol. The results indicate that ACS1 mRNA levels are comparable in GS115 and Δmxr1 during ethanol metabolism (Fig. 1F) suggesting that Mxr1p does not regulate ACS1 expression during ethanol metabolism. It is pertinent to mention that *P. pastoris* homologs of transcription factor Cat8 regulate the expression of ACS1 during ethanol metabolism (22). To further investigate regulation of ALD6-1 by Mxr1p, we generated GS115-A and Δmxr1-A expressing Myc-tagged ALD6-1 (ALD6-1Myc) and examined ALD6-1 levels in the lysates of cells cultured in YNBE by Western blotting using anti-Myc epitope antibodies. The results indicate that ALD6-1Myc is downregulated in Δmxr1-A (Fig. 1G). The gene encoding GFP was cloned downstream of 1.0 kb of ALD6-1 promoter (PΔ-GFP), and protein levels in the lysates of GS115 and Δmxr1 cultured in YNBE were examined by Western blotting using anti-GFP antibodies. GFP levels are also downregulated in Δmxr1 (Fig. 1H) suggesting that Mxr1p is likely to regulate ALD6-1 expression at the transcriptional level. To understand the importance of ALD6-1 in ethanol metabolism, gene encoding ALD6-1 was deleted to generate Δald6-1 (Fig. 1I), which exhibits retarded growth when cultured in YNBE (Fig. 1J). Expression of ALD6-1Myc from its own promoter in Δald6-1 (Fig. 1K) readily restores growth in YNBE (Fig. 1L). However, expression of ALD6-1Myc from GAPDH promoter in Δmxr1 (Fig. 1M) does not result in the restoration of growth in YNBE (Fig. 1N). Thus, ALD6-1 has a unique, essential, and nonredundant function during ethanol metabolism. The ability of ALD6-1 to restore growth of Δald6-1 but not Δmxr1 suggests that Mxr1p-regulated genes other than ALD6-1 (Fig. 1, C and D) are essential for ethanol metabolism.

**Identification of MXREs in ALD6-1 promoter (PΔ)**

Since Mxr1p regulates the transcription of target genes by binding to MXREs in their promoters bearing the consensus sequence 5′-CYCCNY-3′ (4, 5), we analyzed 1.0 kb of ALD6-1 promoter (PΔ) and identified three putative MXREs designated as MXRE1, MXRE2, and MXRE3 (Fig. 2A). Since point mutations within the 5′-CYCCNY-3′ motif of AOX1 MXREs abrogate Mxr1p binding (19), 5′-CYCC-3′ sequence was mutated to 5′-CYCA-3′ in the PΔMXREs (Fig. 2B). Oligonucleotides carrying these mutations were synthesized and designated as MXRE1-M, MXRE2-M, and MXRE3-M (Fig. 2B). A recombinant Mxr1p containing 150 N-terminal amino acids including the DBD, which specifically binds to MXREs of AOX1 promoter, was purified from *Escherichia coli* cell lysates (19) and used in an electrophoretic mobility shift assay together with radiolabeled PΔMXREs. The results indicate that Mxr1pN150 binds to wildtype but not the mutant PΔMXREs (Fig. 2C). PΔ, GFP constructs carrying a point mutation in one or all three MXREs were generated (Fig. 2D), transformed into GS115, and GFP levels were examined by Western blotting using anti-GFP antibodies in the lysates of cells cultured in YNBE. The results indicate that mutation of MXRE1, MXRE2, or MXRE3 alone has no significant effect on GFP expression (Fig. 2E). However, GFP expression is significantly reduced when all the three are mutated (Fig. 2F) suggesting that a combination of two MXREs is sufficient for transactivation by Mxr1p from PΔ during ethanol metabolism. Key results obtained thus far are summarized in Figure 2G.

**Mxr1N400 activates ALD6-1 transcription during ethanol metabolism**

Among the various *P. pastoris* transcription factors identified thus far (6–11), Mxr1p has the unique distinction of functioning as a global regulator of multiple metabolic pathways (4, 5, 12). Mxr1p full-length protein (Mxr1FL) consists of 1155 amino acids. A truncated Mxr1p (Mxr1N400) consisting of 400 N-terminal amino acids was shown to activate the transcription of Mxr1p target genes of acetate and amino acid metabolism as well as AOX1 of methanol metabolism (4, 5, 20). Here, we examined the ability of Mxr1N400 to activate
mRNA levels (Fig. 4D) of \( \Delta \text{mxr1} \) indicating that the putative 9aaTAD functions as TAD during ethanol metabolism.

Identification of NLS in \( \text{Mxr1N250} \)

The localization of \( \text{Mxr1N250} \) in the nucleus of cells cultured in YNBE led us to investigate the presence of NLS within 250 N-terminal amino acids. Analysis of the amino acid sequence of \( \text{Mxr1N250} \) by nLS mapper (http://nls-mapper.ib.keio.ac.jp/cgi-bin/NLS_Mapper_form.cgi) revealed the presence of multiple putative NLSs (pNLS-A, pNLS-B, and pNLS-C) (Fig. 4D). To examine their function, \( \text{Mxr1N400} \) carrying a deletion of these putative TADs was generated (\( \text{Mxr1N400}\Delta\text{TAD} \), expressed in \( \Delta \text{mxr1} \), and \( \text{ALD6-1} \) mRNA levels were analyzed by qPCR. The results indicate that \( \text{Mxr1N400} \) but not \( \text{Mxr1N400}\Delta\text{TAD} \) restores \( \text{ALD6-1} \) mRNA levels (Fig. 4G) as well as growth (Fig. 4H) of \( \Delta \text{mxr1} \) indicating that the putative 9aaTAD functions as TAD during ethanol metabolism.

| Gene       | \( \text{P. pastoris CBS7435} \) | UniProt ID | \( \text{P. pastoris GS115} \) | UniProt ID |
|------------|---------------------------------|------------|--------------------------------|------------|
| \( \text{ALD6-1} \) | PP7435_Chr4-0972                 | F2R0E4     | PAS_chrd_0043                  | C4R6P6     |
| \( \text{ALD6-2} \) | PP7435_Chr3-0183                 | F2QUS7     | PAS_chrd_0987                  | C4R6D4     |
| \( \text{ALD4} \)  | PP7435_Chr2-0787                 | F2QSU4     | PAS_chrd2-1_0853               | C4R0W4     |
| \( \text{ALD2} \)  | PP7435_Chr2-0843                 | F2QSZ7     |                                 | C4ROQ6     |

**Table 2** *Pichia pastoris* ALDs

\( \text{ALD6-1} \) transcription during ethanol metabolism. When expressed in \( \Delta \text{mxr1} \), \( \text{Mxr1N400} \) was as efficient as \( \text{Mxr1FL} \) in restoring \( \text{ALD6-1} \) mRNA levels (Fig. 3A). Furthermore, \( \text{Mxr1N400} \) reversed the growth defect of \( \Delta \text{mxr1} \) as efficiently as \( \text{Mxr1FL} \) (Fig. 3B). Thus, 400 N-terminal amino acids of \( \text{Mxr1p} \) are sufficient to activate the transactivation of not only \( \text{ALD6-1} \) but also other \( \text{Mxr1p} \)-regulated genes necessary for ethanol metabolism.

\( \text{Mxr1N400} \) was shown to possess a TAD between amino acid residues 246 and 280 (Fig. 3C) (20). Mutation of three phenylalanine residues (F249, F254, and F278) within this TAD to alanine has been reported to result in the loss of transactivation function of \( \text{Mxr1p} \) (20). To examine the role of these phenylalanine residues in the activation of \( \text{ALD6-1} \) transcription during ethanol metabolism, we generated \( \text{Mxr1N400Q*} \) in which phenylalanine residues were mutated to alanine (Fig. 3C). When expressed in \( \Delta \text{mxr1-A} \), it restores \( \text{ALD6-1}^{\text{Myc}} \) levels as efficiently as \( \text{Mxr1N400} \) (Fig. 3D) indicating that phenylalanine residues are not essential for the transactivation of \( \text{ALD6-1} \) during ethanol metabolism.

Analysis of amino acid sequence of \( \text{Mxr1N400} \) indicated the presence of a glutamine-rich region between 151 and 194 residues (Fig. 3E). Several transcription factors ranging from yeast to humans have been shown to contain glutamine-rich regions within their TADs (15, 23–27). To investigate the role of glutamine residues of \( \text{Mxr1N400} \) in the transactivation of \( \text{ALD6-1} \), we generated \( \text{Mxr1N400Q*} \) as well as \( \text{Mxr1N400F*} \) in which glutamine residues were mutated to alanine, respectively (Fig. 3E). When expressed in \( \Delta \text{mxr1-A} \), both these mutant proteins restore \( \text{ALD6-1}^{\text{Myc}} \) protein (Fig. 3F) as well as mRNA (Fig. 3G) levels in cells cultured in YNBE. Thus, neither phenylalanine residues nor glutamine residues are essential for the transactivation of \( \text{ALD6-1} \) during ethanol metabolism.

To further characterize the N-terminal TAD, we generated \( \text{Mxr1N250} \) and \( \text{Mxr1N150} \) constructs in which GFP was fused to 250 and 150 N-terminal amino acids of \( \text{Mxr1p} \), respectively (Fig. 4A), and these were expressed in \( \Delta \text{mxr1-A} \). Protein expression was confirmed by Western blotting using anti-GFP antibodies (Fig. 4B). \( \text{Mxr1N250} \) and \( \text{Mxr1N150} \) failed to restore \( \text{ALD6-1}^{\text{Myc}} \) protein (Fig. 4C) and mRNA (Fig. 4D) levels in \( \Delta \text{mxr1-A} \) suggesting that the region between 251 and 400 amino acid residues of \( \text{Mxr1p} \) harbors putative TAD(s) essential for ethanol metabolism. Nuclear localization is essential for the regulation of expression of target genes by transcription factors, and it is possible that the inability of \( \text{Mxr1N250} \) to activate \( \text{ALD6-1} \) transcription may be due to the loss of an NLS present between 251 and 400 amino acids. To rule out this possibility, subcellular localization of \( \text{Mxr1N250} \) was examined in cells cultured in YNBE by visualizing direct GFP fluorescence in a confocal microscope. The results indicate that \( \text{Mxr1N250} \) localizes to the nucleus during ethanol metabolism (Fig. 4E and Fig. S1), and the inability of \( \text{Mxr1N250} \) to activate \( \text{ALD6-1} \) expression may be due to the loss of a transactivation function present between 251 and 400 amino acids.

Transcription factors, such as \( \text{Oaf1}, \text{Pip2}, \) and \( \text{Gal4} \), possess a 9aaTAD, which is present in several other transcription factors, such as \( \text{E2A}, \text{MLL}, \text{p53}, \text{NF-κB}, \text{NFAT}, \text{CEBP}, \text{Sox18}, \text{Pdr1}, \text{Gcn4}, \) and \( \text{VP16} \) (28–31). The 9aaTAD interacts with the transcriptional cofactor TAF9 as well as the KIX domain of general transcriptional mediators CBP, p300, or MED15 (28–31). In silico analysis of the amino acid sequence of \( \text{Mxr1N400} \) using the 9aaTAD prediction tool (https://www.med.muni.cz/9aatad/index.php) indicated the presence of a putative 9aaTAD between amino acids 365 and 373 (Fig. 4F). To examine its function, \( \text{Mxr1N400} \) carrying a deletion of this putative TAD was generated (\( \text{Mxr1N400}\Delta\text{TAD} \), expressed in \( \Delta \text{mxr1} \), and \( \text{ALD6-1} \) mRNA levels were analyzed by qPCR. The results indicate that \( \text{Mxr1N400} \) but not \( \text{Mxr1N400}\Delta\text{TAD} \) restores \( \text{ALD6-1} \) mRNA levels (Fig. 4G) as well as growth (Fig. 4H) of \( \Delta \text{mxr1} \) indicating that the putative 9aaTAD functions as TAD during ethanol metabolism.

**Key functional domains of \text{Pichia pastoris Mxr1p}**
**Key functional domains of *Pichia pastoris* Mxr1p**

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**Mxr1N400 is necessary but not sufficient for the activation of Mxr1p target genes required for methanol metabolism**

During methanol metabolism, Mxr1p activates the transcription of *AOX1* encoding alcohol oxidase 1 as well as other methanol utilization (MUT) pathway genes, such as *DHAS*, *FDH*, *PEX5*, *PEX8*, and *PEX14* encoding dihydroxyacetone synthase, formaldehyde dehydrogenase, formate dehydrogenase, peroxins 5, 8, and 14, respectively (6). Furthermore, Mxr1N400 was shown to be sufficient for the activation of *AOX1* during methanol metabolism (20). Interestingly,
Figure 3. Characterization of Mxr1N400 TAD during ethanol metabolism. A, quantitation of ALD6-1 mRNA levels by quantitative PCR (qPCR) in different Pichia pastoris strains cultured in YNBE as indicated. B, analysis of growth of different P. pastoris strains cultured in YNBE. Absorbance at 600 nm of liquid cultures was measured at different time intervals as indicated. C, schematic representation of Mxr1N400 and Mxr1N400F*. The TAD between amino acids 246 and 280 as well as F249, F254, F278, important for transactivation of AOX1 as reported by Parua et al. (20), are shown. In Mxr1N400F*, phenylalanine residues were mutated to alanine residues. D, Western blot analysis of ALD6-1Myc using anti-Myc epitope antibodies in the lysates of different P. pastoris strains as indicated. Cells were cultured in YNBE for 14 h. Quantitation of data is shown. The intensity of individual bands was quantified and expressed as arbitrary units ± SD relative to controls (n = 3). E, schematic representation of Mxr1N400, Mxr1N400Q*, and Mxr1N400F*Q*. Glutamine residues were mutated to alanine residues in Mxr1N400Q*. Both glutamine and phenylalanine residues were mutated to alanine residues in Mxr1N400F*Q*. F, Western blot analysis of ALD6-1Myc using anti-Myc epitope antibodies in the lysates of different P. pastoris strains as indicated. Cells were cultured in YNBE for 14 h. Quantitation of data is shown. The intensity of individual bands was quantified and expressed as arbitrary units ± SD relative to controls (n = 3). G, quantitation of ALD6-1 mRNA levels by qPCR in different P. pastoris strains cultured in YNBE as indicated. Error bars in each figure indicate SD; n = 3. In the bar diagrams, p value summary is mentioned on the bar of each figure. *p < 0.05; **p < 0.005; ***p < 0.0005; and ns, Student’s unpaired t test was done. ns, not significant; TAD, transactivation domain; YNBE, yeast nitrogen base and 1% ethanol.
Figure 4. Characterization of putative N-terminal TAD and NLS in Mxr1N250. A, schematic representation of Mxr1N250 and Mxr1N150 carrying N-terminal GFP fusions. B, Western blot analysis of Mxr1N250 and Mxr1N150 in cells cultured in YNBE using anti-GFP antibodies. C, Western blot analysis of ALD6-1Myc in the lysates of different Pichia pastoris strains as indicated using anti-Myc epitope antibodies. Cells were cultured in YNBE for 14 h. Quantitation of data is shown. The intensity of individual bands was quantified and expressed as arbitrary units ± SD relative to controls (n = 3). D, quantitation of ALD6-1 mRNA levels by quantitative PCR (qPCR) in different P. pastoris strains cultured in YNBE for 14 h. E, live cell confocal imaging of Mxr1N250 cultured in YNBE. Hoechst 33342 was used to stain nuclei. GFP was visualized by direct fluorescence. F, schematic representation of putative nine-amino acid TAD present in Mxr1N400 between amino acids 365 and 373. G, quantitation of ALD6-1 mRNA levels by qPCR in different P. pastoris strains cultured in YNBE for 14 h. H, analysis of growth of different P. pastoris strains cultured in YNBE. I, three putative NLSs (pNLS-A, pNLS-B, and pNLS-C) in Mxr1N250 as predicted by cNLS
Mxr1N400 as well as Mxr1N400f+, Mxr1N400Q+, and Mxr1N400fQ+ restore AOX1 mRNA levels when expressed in Δmxr1 cultured in YNB and 1% methanol (YNBM) (Fig. 5A). Thus, phenylalanine and glutamine residues in Mxr1N400 have no major role in the activation of Mxr1p target genes of ethanol as well as methanol metabolism (Figs. 3G and 5A). However, the ability of Mxr1N400 to regulate transcription of Mxr1p target genes other than AOX1 has not been investigated. Here, we demonstrate that Mxr1N400 activates the transcription of not only AOX1 but also other MUT pathway genes as evident from qPCR analysis of mRNA isolated from cells cultured in a medium containing YNBM (Fig. 5B). The mRNA levels of MUT pathway genes are twofold to fivefold higher in Δmnr1-N400 than those in GS115 because of overproduction of Mxr1N400 from GAPDH promoter as evident from qPCR analysis of MXR1 mRNA levels (Fig. 5C). Surprisingly, Mxr1N400 does not rescue the growth defect of Δmxr1 cultured in YNB, despite activating the transcription of several key Mxr1p target genes of methanol metabolism (Fig. 5D). Thus, in addition to the known Mxr1p targets, that is, MUT pathway genes, Mxr1p regulates the expression of several other genes essential for methanol metabolism whose identity is not known. It is likely that transactivation of these essential genes is facilitated by other putative TAD(s) localized beyond 400 N-terminal amino acids of Mxr1p. Interestingly, the region between 401 and 1155 amino acids harbors several putative 9aaTADs (Fig. 5E and Fig. S7), some of which may be involved in the activation of genes essential for methanol metabolism.

Identification of novel targets of Mxr1p essential for methanol metabolism

To identify genes of methanol metabolism whose transactivation is dependent on Mxr1FL rather than Mxr1N400, RNA was isolated from GS115, Δmxr1, Δmxr1-FL, and Δmnr1-N400 grown in YNB, and biological replicates were subjected to DNA microarray for the identification of differentially expressed genes. In all the comparisons, GS115 was used as a control for clustering differentially expressed genes. To identify significantly regulated genes, a cutoff of p value <0.01 was applied. Microarray datasets have been deposited into the GEO database under accession number GSE146829. Volcano plots and heat maps are used to illustrate overall changes in gene expression.

Comparative analysis of the transcriptome of GS115, Δmxr1, Δmxr1-FL, and Δmnr1-N400 indicates that Mxr1FL is more efficient than Mxr1N400 in restoring the gene expression pattern of Δmxr1 to that of GS115 (Fig. 6, A–C). At least 21 genes are downregulated in Δmnr1, and their expression is restored to almost wildtype levels by Mxr1FL but not Mxr1N400 (Fig. 6B) suggesting that the region beyond 400 amino acids of Mxr1p is required for their transactivation. This is further validated by qPCR of select genes such as HST2 encoding an NAD⁺-dependent protein deacetylase of the silencing information regulator 2 family, FET4-1 encoding low-affinity Fe (II) transporter of the plasma membrane, and cell agglutination protein mam3 (Fig. 6D). HST2 is one of the most downregulated genes in Δmnr1, and its expression is fully restored by the expression of Mxr1FL but not Mxr1N400 (Fig. 6, B and D). Surprisingly, deletion of Mxr1p resulted in the upregulation of genes as well. A cluster of 17 genes is specifically upregulated in Δmnr1, and this is reversed more efficiently by Mxr1FL than Mxr1N400 (Fig. 6C). Thus, the region beyond 400 N-terminal amino acids is required for the regulation of full complement of Mxr1p target genes essential for methanol metabolism, and this explains the need for Mxr1FL to rescue the growth defect of Δmxr1 cultured in YNB.

In addition to the demonstration of differential regulation of genes of methanol metabolism by Mxr1FL and Mxr1N400, the DNA microarray analysis has led to the identification of several novel target genes of Mxr1p in cells metabolizing methanol. Deletion of MXR1 altered the transcriptional landscape drastically (Fig. 6A), and a total of 161 genes with fold ≤–1, 82 genes with fold ≤–1.5, and 32 genes with fold ≤–2 were downregulated in Δmnr1 (Supporting information 3 and Fig. 7A). Interestingly, deletion of MXR1 resulted in upregulation of several genes (Fig. 7B), which is surprising, since Mxr1p is known to function as a transcriptional activator rather than transcriptional repressor. In fact, the number of upregulated genes surpassed that of downregulated genes in Δmnr1, with roughly 222 genes upregulated ≥1-fold, 135 genes ≥1.5-fold, and 79 genes ≥2-fold (Supporting information 3 and Fig. 7B). We have validated the microarray data by qPCR analysis of select genes. These include downregulated genes FBA1-2, CLN2, and POX1 encoding fructose-bisphosphate aldolase, G1/S-specific cyclin, and acyl CoA oxidase, as well as upregulated genes ACS1, and ACS2 and ICL encoding ACS1 and ACS2 and isocitrate lyase, respectively (Fig. 7, C and D). The enriched Gene Ontology (GO) categories for the downregulated genes include RNA-related processes, such as RNA methyltransferase, catalytic activity on RNA, metabolic processes, and transferase activity. In case of upregulated genes, enriched GO categories include primarily carboxylic acid, nucleotide, and amino acid metabolic processes (Fig. S8).

Mxr1p target genes have been shown to possess the 5'-CYCCNY-3' consensus sequence (MXRE) in their promoter regions. Therefore, we examined the presence of MXREs within 1.0-kb promoter region of all the differentially regulated genes.
Cells were cultured in YNBM for 14 h. Error bars in each of the P. pastoris growth of different strains as indicated. Error bars in each MXR1 during methanol metabolism in different strains cultured in YNBM (http://rsat-tagc.univ-mrs.fr/rsat/). Analysis Tools fungi (http://rsat-tagc.univ-mrs.fr/rsat/). Analysis reveals that 82% and 73% of the downregulated and upregulated genes, respectively, harbor at least one MXRE in their promoter (Supporting informations 4 and 5) indicating that these genes may be direct targets of Mxr1p. Number of genes with 1 to 10 MXREs within 1.0 kb of their promoters are indicated in Figure 7, E and F. To examine whether MXR4s of upregulated genes are different from those of downregulated genes, 5'-CYCCNY-3' along with flanking sequences were submitted to MDDLogo (https://weblogo.berkeley.edu/logo.cgi). While the 5'-CYCCNY-3' motif is conserved in both upregulated and downregulated genes, we did not observe enrichment of nucleotides in the flanking sequences (Fig. 7, G and H) making it difficult to derive a consensus sequence specific for upregulated and downregulated genes.

**Discussion**

This study demonstrates for the first time that Mxr1p is required for efficient ethanol metabolism in P. pastoris and in its absence, ALD6-1-mediated conversion of acetaldehyde to acetate is impaired resulting in growth defect. This growth defect is not rescued by other ALDs indicating that ALD6-1 has an important and nonredundant function during ethanol metabolism. Mxr1p binds to three MXREs in ALD6-1 promoter in vitro, and a combination of two of them is sufficient for Mxr1p-dependent expression of a reporter gene from ALD6-1 promoter in vivo during ethanol metabolism. This result is similar to that reported in an earlier study in which two MXREs in the promoter of ACS1 encoding ACS1 were shown to be required for Mxr1p-mediated transactivation during acetate metabolism (4). Mutation of all three MXREs does not completely abrogate GFP expression from ALD6-1 promoter (Fig. 2F) suggesting that transcription factors other than Mxr1p may bind elsewhere in the promoter and contribute to ALD6-1 promoter activity. Thus far, studies have focused on the Mxr1p-mediated repression of genes of MUT pathway such as AOX1 during ethanol metabolism (20, 32). Phosphorylation of serine 215 residue of Mxr1p was shown to be involved in the transcriptional repression of AOX1 during ethanol metabolism (20). Furthermore, acetylation of histones or other transcription factors by acetyl-CoA synthesized during ethanol metabolism may also contribute to the repression of methanol-inducible genes (32). However, no other study except the present study investigated the ability of Mxr1p to regulate the expression of gene(s) essential for ethanol metabolism. While we have focused our attention on ALD6-1, the most downregulated Mxr1p target gene in Δmxr1, it will be interesting to study other Mxr1p targets as well. Overexpression of ALD6-1 in Δmxr1 using a heterologous promoter does not result in restoration of growth (Fig. 1, M and N) indicating that some of the Mxr1p target genes identified in this study by RNA-Seq may also have a key role during ethanol metabolism.

Study of the transactivation functions of Mxr1p during ethanol metabolism is another important aspect of this study. We demonstrate that Mxr1N400 activates the transcription of not only ALD6-1 but also all the Mxr1p target genes required for ethanol metabolism as evident from the fact that growth of Δmxr1 is restored by Mxr1N400 as efficiently as Mxr1FL. We further demonstrate that phenylalanine as well as glutamine residues in Mxr1N400 do not have a role in the transactivation...
Figure 6. Identification of methanol metabolism genes differentially regulated by Mxr1N400 and Mxr1FL. A, volcano plot highlighting significant upregulated and downregulated genes. Genes differentially expressed between GS115 and Δmxr1, GS115 and Δmxr1-FL, and GS115 and Δmxr1-N400 are shown. The y-axis corresponds to $-\log_{10}$ of $p$ value, and x-axis displays the log2 fold change value. Green and red dots represent downregulated and upregulated transcripts, respectively. B, heat map of methanol metabolism genes downregulated in Δmxr1 relative to GS115. When expressed in Δmxr1, Mxr1-FL reverses downregulation more effectively than Mxr1N400 (compare Δmxr1-FL and Δmxr1-N400). Asterisk indicates genes whose expression was validated by quantitative PCR (qPCR). C, heat map of methanol metabolism genes upregulated in Δmxr1 relative to GS115. When expressed in Δmxr1, Mxr1-FL reverses upregulation more effectively than Mxr1N400 (compare Δmxr1-FL and Δmxr1-N400). To identify significantly regulated genes, a cutoff of $p$ value <0.01 was applied. D, validation of DNA microarray data by qPCR of select genes. Error bars indicate SD; n = 2.
of ALD6-1 in cells metabolizing ethanol. The region between 251 and 400 N-terminal amino acids of Mxr1p is essential for ALD6-1 transactivation during ethanol metabolism. Furthermore, 9aaTAD present between 365 and 373 amino acids functions as TAD during ethanol metabolism.

Nuclear localization is essential for Mxr1p to function as a transcription factor. The mechanism by which Mxr1p translocates from cytosol to the nucleus in cells cultured in non-fermentable carbon sources is not known. In this study, we not only identified several putative NLSs using NLS prediction methods.
tools but also experimentally validated them and identified one of them (pNLS-C1) as the actual NLS. Whether other NLSs are present in the region beyond 250 N-terminal amino acids remains to be investigated. NLSs enriched in basic amino acids bind to importin α or importin β, and the resultant protein complex is translocated to the nucleus (33). It will be interesting to examine the role of importin α or β in the nuclear import of Mxr1N250. NLSs are not well characterized in P. pastoris (34), and this is first reported on the detailed characterization of NLS of a P. pastoris transcription factor.

Study of transactivation functions of Mxr1p during methanol metabolism led to interesting results. Phenylalanine residues (F249, F254, and F278) that had previously been reported to be indispensable for transactivation function of Mxr1p by Parua et al. (20) were found to have no role in the activation of either ALD6-1 or AOX1 during ethanol and methanol metabolism, respectively. This discrepancy is primarily because of different experimental systems used in these two studies. It should be noted that Parua et al. (20) examined transactivation by Mxr1p TAD (246–280 amino acids) fused to Gal4-DBD of lacZ gene from a Gal4-responsive promoter in S. cerevisiae metabolizing glucose (20). Our study is physiologically more relevant as we have investigated the role of phenylalanine and glutamine residues of Mxr1N400 in the transactivation of native Mxr1p target genes such as ALD6-1 and AOX1 in P. pastoris rather than a heterologous yeast such as S. cerevisiae. Despite mutation of phenylalanine and glutamine residues, Mxr1N400 still activated AOX1 transactivation indicating that other amino acid residues contribute to the transactivation function of Mxr1N400. In fact, Mxr1N250 failed to activate AOX1 transactivation during methanol metabolism as well (data not shown) suggesting that the region between 251 and 400 amino acids may also be involved in the activation of genes of methanol metabolism.

The fact that Mxr1N400 failed to rescue the growth defect of Δmxr1 during the methanol metabolism despite activating AOX1 as well as other genes known to be involved in MUT pathway suggested that the region beyond 400 N-terminal amino acids is essential for the regulation of other Mxr1p target genes required for methanol metabolism. DNA microarray analysis clearly demonstrated that Mxr1FL restores the expression of several genes downregulated in Δmxr1 more efficiently than Mxr1N400. This is true for genes upregulated in Δmxr1 as well. It will be interesting to examine the role of 9aaTAD identified in this study as well as the putative 9aaTADs present between 401 and 1155 amino acids in the regulation of Mxr1p target genes during methanol metabolism.

This study demonstrates that Mxr1FL regulates the expression of several essential genes for methanol metabolism in addition to the previously reported classical MUT pathway genes. For example, the expression of several genes involved in epigenetic regulation such as histone acetyltransferases and histone deacetylases are regulated by Mxr1p (Supporting information 3 and Fig. 6, B and C). In fact, HST2 encoding a histone deacetylase of the silencing information regulator 2 family is one of the most downregulated genes in Δmxr1 during methanol metabolism. Investigation of the role of HST2 in methanol metabolism is a topic for future research. The upregulation of several genes in Δmxr1 during ethanol as well as methanol metabolism suggests that Mxr1p may function as a transcriptional repressor as well. In fact, 73% of genes upregulated in Δmxr1 cultured in YNBM possess at least one MXRE in their promoters. While the core MXRE (5′-CYYCCNY-3′) was present in the promoters of both upregulated and downregulated genes, sequences flanking the core MXRE did not exhibit high degree of conservation rendering it difficult to derive a consensus sequence specific for upregulated and downregulated genes. Functional characterization of MXREs of genes upregulated in Δmxr1 is a topic of future research. In addition to the identification of Mxr1p as a key regulator of ethanol metabolism, this study provides new insights into the mechanism by which Mxr1p functions as a global regulator of multiple metabolic pathways of P. pastoris.

Key functional domains of Pichia pastoris Mxr1p

During the methanol metabolism, investigation of the role of HST2 in methanol metabolism is a topic for future research. The upregulation of several genes in Δmxr1 during ethanol as well as methanol metabolism suggests that Mxr1p may function as a transcriptional repressor as well. In fact, 73% of genes upregulated in Δmxr1 cultured in YNBM possess at least one MXRE in their promoters. While the core MXRE (5′-CYYCCNY-3′) was present in the promoters of both upregulated and downregulated genes, sequences flanking the core MXRE did not exhibit high degree of conservation rendering it difficult to derive a consensus sequence specific for upregulated and downregulated genes. Functional characterization of MXREs of genes upregulated in Δmxr1 is a topic of future research. In addition to the identification of Mxr1p as a key regulator of ethanol metabolism, this study provides new insights into the mechanism by which Mxr1p functions as a global regulator of multiple metabolic pathways of P. pastoris.

Experimental procedures

Growth media and culture conditions

P. pastoris cells were maintained in nutrient-rich yeast extract, peptone, and dextrose (YPD) agarose plates (1% yeast extract, 2% peptone, and 2% dextrose) and a single colony was cultured overnight in YPD at 30 °C in an orbital shaker at 180 rpm followed by washing with sterile distilled water (twice) and transferred to desired media consisting of 0.67% YNB (YNB without amino acids and with ammonium sulphate) supplemented with a carbon source 1% methanol (YNBM) or 1% ethanol (YNBE). Yeast extract and peptone were purchased from Becton and Dickinson Biosciences, India. E. coli TOP10 was used for plasmid isolation while expression of recombinant proteins was carried out in BL21(DE3) strain of E. coli. Bacterial and yeast transformations were done using the CaCl2 method and electroporation (Gene Pulsar; Bio-Rad), respectively.

Antibodies and other reagents

Anti-phosphoglycerate kinase (PGK) polyclonal antibody was generated by immunizing the rabbit with a purified 6x His-tagged P. pastoris PGK protein. Mouse anti-Myc was purchased from Merck Millipore. Restriction enzymes and T4 DNA ligase were purchased from New England Biolabs. DNA polymerases were purchased from Genei. Oligonucleotides were purchased from Sigma–Aldrich. Nucleotide sequence of primers used in qPCR and RT-PCR will be provided on request.

Growth kinetics

A single colony was inoculated in 5 ml YPD and grown at 30 °C in an orbital shaker at 180 rpm for 18 to 24 h or till late log phase. Cells were pelleted and washed twice with autoclaved milliQ under sterile conditions and resuspended in water. Cells were then inoculated into specific media at an initial absorbance of 0.06 to 0.1 at 600 nm and grown at 30 °C at 180 rpm in an orbital shaker. Absorbance at 600 nm was measured at regular intervals till cells attain stationary phase.
Key functional domains of Pichia pastoris Mxr1p

Subcellular localization studies

Localization of N-terminal GFP-tagged Mxr1 strains was performed by direct fluorescence. For this, cells cultured in desired medium for 14 to 16 h were fixed by incubating with 3.7% formaldehyde in an orbital shaker at 180 rpm at 30 °C for 1 h. Cells were then harvested, washed with 1× PBS (137 mM NaCl, 2.68 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4), followed by spheroplast preparation by incubating with 5 μl of Longlife Zymolyase (G-Biosciences; 1.5 U/ml) in 200 μl of zymolyase buffer (1.2 M sorbitol, 40 mM phosphate buffer, 50 mM MgCl2, and 28.4 mM β-mercaptoethanol) at 37 °C for 30 min. Harvested spheroplasts were washed and resuspended in 1× PBS + 0.05% Tween-20. Smears were prepared and fixed by treatment with methanol and acetone. For nuclear staining, fixed smears were incubated with 1 mg/ml of Hoechst 33342 for 7 min in dark, followed by washing with 1× PBS for 10 min, and mounting on slide and sealing with nail polish. Slides were visualized in confocal microscope (Olympus Fluoview FV3000) with a 100× objective.

Real-time qPCR

P. pastoris cells were cultured for 12 to 14 h, and total RNA was isolated using RNA isolation kit (catalog no. Z3100; Promega) as per the manufacturer’s instructions. RNA was quantified by measuring the absorbance at 260 nm using NanoDrop 2000 spectrophotometers, and 1 μg of DNase-treated RNA was used for complementary DNA (cDNA) preparation. qPCR was carried out using iQ SYBR Green supermix in a StepOnePlus Real-Time PCR system (Thermo Fisher Scientific). The relative mRNA expression levels were obtained using the ΔΔCt method, wherein the Ct value of a sample, which is the Ct value of a gene relative to that of 18S rRNA, is normalized to that of the control.

Western blotting

Cells were lysed using glass-bead lysis method followed by quantification by Bradford assay, separation of protein bands on SDS-PAGE, transfer to polyvinylidene fluoride membrane, and probing with primary and secondary antibodies. ImageJ software (https://imagej.nih.gov/ij/) was applied to quantify all Western blot data. Band intensity of the protein of interest was normalized to that of the loading control, PGK.

Statistical analysis

Student’s t test was carried out using GraphPad Prism 5, software (GraphPad Software, Inc). Data are represented as mean ± SD. p Value summary is mentioned on the bar of each figure where *p < 0.05; **p < 0.005; ***p < 0.0005, and ns, not significant.

RNA-Seq and data analysis

Total RNA was isolated from GS115 and Δmxr1 cultured in duplicates in YNBE for 14 h by Qiagen RNeasy kit according to the manufacturer’s protocol. RNA-Seq was performed using Illumina HiSeq. FastQC and MultiQC software were used to check data quality. The data were checked for sequencing adapter contamination, base call quality distribution, percent of bases above Q20, Q30, and %GC. For all the samples, quality control was above threshold (Q20 > 95%). Fastp was used to process raw sequence reads by removal of adapter sequences and low-quality bases. Bowtie2 aligner was used to align the quality control–passed reads onto indexed K. phaffii CBS 7435 reference genome (GCA_900235035.1_ASM90023503v1). About 94.73% of the reads could be mapped to the reference genome on average. Gene level expression values were obtained as read counts using feature counts software. Expression similarity between the biological replicates was checked by Spearman correlation and principal component analysis. DESeq2 package was used for differential expression analysis. Genes with less than five reads in any of the two samples were removed from further analysis. The read counts were normalized (variance stabilized normalized counts) using DESeq2, and differential enrichment analysis was performed. The protein sequences were annotated against the Uniref100 protein database using BLASTp module of Diamond. The gene names, length, GO annotations, and other additional information were obtained by submitting the Uniref100 Ids to UniProtKB. Genes with absolute log2 fold change ≥1 and adjusted p ≤0.05 were considered significant. The expression profile of differentially expressed genes across the samples is presented in volcano plot and heat map. The enriched GO categories were retrieved from Amigo using S. cerevisiae as organism. The transcriptome datasets generated during the current study are available in the National Center for Biotechnology Information (NCBI) with the accession number GSE168677.

Microarray hybridization and data analysis

For the generation of samples for microarray analysis, GS115, Δmxr1, Δmxr1-FL, and Δmxr1-N400 were cultivated in biological duplicates in YNBM for 14 h. Total RNA extraction was performed using Qiagen RNeasy kit according to the manufacturer’s protocol. The microarray hybridization and scanning were performed at the Agilent-certified microarray facility of Genotypic Technology, Bengaluru, India. The samples for gene expression were labeled using Agilent Quick-Amp Labeling Kit (part number: 5190-0442). The total RNA was reverse transcribed at 40 °C using oligoT primer tagged to a T7 polymerase promoter and converted to double-stranded cDNA. Synthesized double-stranded cDNA was used as a template for cRNA generation by in vitro transcription, and the dye Cy3 CTP (Agilent) was incorporated during this step. The cDNA synthesis and in vitro transcription steps were carried out at 40 °C. Labeled cRNA was cleaned up using Qiagen RNeasy columns (Qiagen; catalog no.: 74106), and quality was assessed for yields and specific activity using the Nanodrop ND-1000. Labeled cRNA sample was fragmented at 60 °C and hybridized on to Agilent P. pastoris Gene Expression Microarray 8X15K AMADID: 085736-One Color Platform. Fragmentation of labeled cRNA and hybridization were done using the Gene Expression Hybridization Kit of...
(In situ) Hybridization kit, part number: 5190-0404; Agilent Technologies. Hybridization was carried out in Agilent’s SureHyb Chambers at 65 °C for 16 h. The hybridized slides were washed using Agilent Gene Expression wash buffers (Agilent Technologies; part number 5188-5327) and scanned using the Agilent Microarray Scanner (part number G2600D; Agilent Technologies). Raw data extraction from images was obtained using Agilent Feature Extraction software. Feature-extracted raw data were analyzed using Agilent GeneSpring GX (version 14.5) software. Normalization of the data was done in GeneSpring GX using the 75th percentile shift method. Significant genes upregulated with fold change ≥1 (logbase2) and downregulated with fold change ≤-1 (logbase2) in the test samples with respect to control sample were identified. Statistical Student's t test p value among the replicates was calculated based on volcano plot algorithm. Differentially regulated genes were clustered using hierarchical clustering based on the Pearson’s coefficient correlation algorithm to identify significant gene expression patterns. Data were analyzed as described in RNA-Seq section. The raw data files are available at the NCBI with an accession number GSE146829.

**Generation of Δmrxr1, Δmrxr1-FL, and Δmrxr1-N400**

Δmrxr1 was generated by disrupting the MXR1 coding region by zeocin resistance cassette (Ze6) in GS115 as has been previously described (4). Δmrxr1-FL and Δmrxr1-N400 were generated by transforming plasmid expressing MxR1FL and MXR1N400 as C-terminal Myc-tagged proteins under GAPDH promoter as described (4).

**Generation of Δald6-1**

A 0.911-kb ALD6-1 promoter region was amplified by PCR from *P. pastoris* genomic DNA using primer pair 5'-GGA CTGTTCATTTAATGGTGATCTGAGCG-3' and 5'-GCT ATGTTGTTGGAGTATCCCCCAGC-3' (962–984 bp of GAPDH promoter in reverse complement, −89 to −114 bp of ALD6-1 promoter in reverse complement). In another PCR, 1.2 kb of zeocin expression cassette was amplified by PCR from pGAPZA vector using the primer pair 5'-CCACCACGATTTCCC AAGGGATGTCGAGCCGATCCCCCAGC-3' (−89 to −114 bp of ALD6-1 promoter, +962 to +984 bp of GAPDH promoter) and 5'-GGAGTGAAGCAATCTCTGATTG ACTTTGCGGTGG-3' (+1467 bp to +1493 bp in reverse complement of 3'-flanking region of ALD6-1, +2137 to +2159 bp in reverse complement of GAPDH promoter). In the third PCR, 872 bp of the 3'-flanking region of ALD6-1 was amplified using the primer pair 5'-GAAGTCAAGGAGTACGTCCACGTCGAGC-3' (+2352 to +2381 bp in the reverse complement of 3'-flanking region of ALD6-1). All the three PCR products were purified and used as a template in the final PCR and amplified using a primer pair 5'-GGACTGTGTAATTGAAGATCGTGCAGCTG ACAG-3' and 5'-GGAAGTGGAGAGGAGGTCTCCGAGA CAACTC-3' to obtain a 3.023-kb product consisting of zeocin expression cassette along with promoter and terminator of ALD6-1. GS115 strain was transformed with Ze6 expression cassette, and zeocin-resistant colonies were selected. Deletion of ALD6-1 was confirmed by PCR using gene-specific primers.

**Generation of GS115-A, Δmrxr1-A, and Δmrxr1-A-N400**

The gene encoding ALD6-1 along with 1.0 kb of its promoter was cloned into pBl3 vector (#25452; Addgene) and transformed into GS115, Δmrxr1, and Δmrxr1-N400. The following primer pair was used: 5'-TCCCCCGGGAGTT GGAAGAGAATCGATCTG-3' and 5'-CCGTCCAGCT ATGGCTCTCGAATGGACTTTCGACTGTC ATGGCTGTAAGC-3' (Xmal and Xhol sites in the primers are underlined). The reverse primer encoded the Myc tag. The PCR product was cloned into pBl3 vector, the recombinant plasmid was linearized using SalI, and transformed into GS115, Δmrxr1, and Δmrxr1-N400. Recombinant clones were selected by plating on YNBD His- plates, and clones expressing Myc-tagged ALD6-1 (ALD6-1Myc) were identified by Western blotting using anti-Myc epitope antibody.

**Generation of GS115-P*α*-GFP and Δmrxr1-P*α*-GFP**

These strains express GFP from 1.0 kb of ALD6-1 promoter and were constructed as described later. Three different PCRs were carried out for the construction of *P*α-GFP expression cassette. In the first PCR, 1 kb ALD6-1 promoter was amplified from genomic DNA of GS115 strain of *P. pastoris* using a primer pair 1F (5'-CGGGATCCATTGAGAGAACAAT AATCAGACT-3' [BamHI site is underlined]) and 1R (5'- CTCCCTTTACTAGCTACTTTACCATGGATAGAAGCTT AAAAGAGGACAGT-3'; +1 to +25 bp of the gene encoding GFP and −1 to −28 bp of *Pα*). In the second PCR, a 714-bp region of *GFP* gene was amplified from pREP1-GFP vector using the primer pair 2F (5'-CACCAGTTGGACAGTGGAGATCTGAGCTG TAAGGAGG-3', −1 to −28 bp of *Pα* and +1 to +25 bp of the gene encoding GFP and 2R (5'-CCGCTTCAGCT CTGATGAGTGGTGAGCTTTCGTTTCGAGC-3'). The Xhol site is underlined. In the third PCR, the PCR products from the first two reactions were used as templates and amplified using the 1F and 2R primers to get the *P*α-GFP expression cassette, which was digested with BamHI and Xhol and cloned into pBl3 (catalog no. 25452; Addgene) to generate pBl3-P*α*-GFP. The generated expression vectors were linearized with Sall and transformed into GS115 and Δmrxr1 strains and plated on YNBD-His- agar plates, and positive colonies were screened using anti-GFP antibody.

**Generation of GS115-P*α*M1-GFP, GS115-P*α*M2-GFP, GS115-P*α*M3-GFP, and GS115-P*α*M4-GFP**

GS115 expressing GFP from 1.0-kb ALD6-1 promoter in which each of the three MXREs (M1, M2, and M3) two MXREs, and all the three MXRES (M4) are mutated were...
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generated as follows: *P*_{ALD6-1}GFP plasmid with MXRE-M1 was generated as follows: *ALD6-1* promoter was PCR amplified using primer pair 1F (5'-CGGGATCCATTGGAGAAGA CAAATGAA TCTGAC-3' [BamHI site is underlined]) and 1R (5'-CCCATCTCCAAAAAGAGGAGCATGATGATGAGAACC GTTGAGGAACTGTCGGTGG-3'). In another PCR, GFP was amplified using primer pair 2F (5'-CACCCATGTTCGCGCTAAGTCTACATCGTTCTTTTGGAGGAGG-3' [mutation is underlined]) and 2R (5'-CCGCCTGAGCTATGTTGTTGGT GCTAGTTTT-3' [the Xhol site is underlined]). In the third PCR, the PCR products from the first two reactions were used as templates and amplified using the 1F and 2R primers. Thereafter, the amplicon was digested with BamHI and XhoI and cloned into ΔGeneration of antibody. Sequence of the synthetic gene will be made available from TWIST Biosciences, wherein the phenylalanine residues at 249, 254, and 278 positions were mutated to alanine in the MXR1 gene 5'-CATGGAC -CATGGAC-3' (Thermo Fisher Scientific). Another synthetic gene, MXR1N400FF*, with phenylalanine residues at 249, 254, and 278 mutated to alanine, and 18 glutamine residues between 150 and 200 amino acids (152, 155, 157, 158, 162, 163, 164, 166, 168, 170, 171, 175, 179, 181, 182, 185, and 194) were mutated to alanine to get MXR1N400F*. This gene was synthesized by GeneArt, with phenylalanine residues at 249, 254, and 278 positions mutated to alanine, and 18 glutamine residues between 150 and 200 amino acids (152, 155, 157, 158, 162, 163, 164, 166, 168, 170, 171, 175, 179, 181, 182, 185, and 194) were mutated to alanine to get MXR1N400F*. This gene was synthesized by GeneArt (Thermo Fisher Scientific). Another synthetic gene, MXR1N400FF*, with phenylalanine residues at 249, 254, and 278 mutated to alanine, and 18 glutamine residues between 150 and 200 amino acids (152, 155, 157, 158, 162, 163, 164, 166, 168, 170, 171, 175, 179, 181, 182, 185, and 194) mutated to alanine in the MXR1N400F* gene 5'-CGGGGTACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3'. In another PCR, the GFP sequence with the N-terminal region of (1311–1329 base pairs) of the GFP gene with an upstream KpnI restriction enzyme site 5'-CGGGGTACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3' (F) and the reverse primer complementary to C-terminal region (1311–1329 base pairs) of *GFP* and overlapping sequence with the N-terminal region of (1–26 base pairs) MXR1 gene 5'-CCTCAAAATGGTGGGAATGGTATCATCACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3' (R). To construct an N-terminal GFP-tagged *Mxr1N150*, first, the GFP sequence was amplified using a common forward primer for amplification of first 150 or 250 bp of *MXR1*, which have an overlapping sequence of C-terminal region (1311–1329 base pairs) of *GFP* and overlapping sequence with the N-terminal region of (1–26 base pairs) MXR1 gene 5'-CATGGAC GAGCTGTACAAGATGAGCAATCTATACATCATTATGAGAACC GTTGAGGAACTGTCGGTGG-3' (K). The reverse primer for amplification of *MXRIN150* is sequence reverse complementary to 432 to 450 bp of *MXR1* gene with a downstream Myc tag sequence, stop codon, and a NotI restriction enzyme site 5'-ATAGTTTTAGCGGCCGCTACTCATGCAATCTATACATCATTATGAGAACC GTTGAGGAACTGTCGGTGG-3' (R1). For amplifying *MXRIN250*, the reverse primer for amplification of reverse primer is reverse complementary to 727 to 750 bp of *MXR1* gene with a downstream Myc-tag sequence, stop codon, and a NotI restriction enzyme site 5'-ATAGTTTTAGCGGCCGCTACTCATGCAATCTATACATCATTATGAGAACC GTTGAGGAACTGTCGGTGG-3' (R2). To get the GFP–MXR fusion product, an overlapping PCR was performed where the amplified GFP and *MXRIN150* or *MXRIN250* PCR products were used as templates, and primer pairs F and R1 or F and R2 were used. This fused PCR product was digested with KpnI and NotI restriction enzymes and cloned under GAPDH promoter in pGHYB (35) vector, followed by transformation in Δmxr1. Positive colonies were screened for expression of Myc-tagged Mxr1N400ΔTAD using anti-Myc epitope antibody.

**Generation of Δmxr1-A-N400F*, Δmxr1-A-N400Q*, and Δmxr1-A-N400FQ***

To introduce mutations in *Mxr1N400*, codon-optimized *MXRIN400* genes were synthesized. *MXRIN400* was synthesized at TWIST Biosciences, wherein the phenylalanine residues at 249, 254, and 278 positions were mutated to alanine residues in *MXRIN400* gene. In the *MXRIN400* gene, 18 glutamine residues between 150 and 200 amino acids (152, 155, 157, 158, 162, 163, 164, 166, 168, 170, 171, 175, 179, 181, 182, 185, and 194) were mutated to alanine to get *MXRIN400Q*. This gene was synthesized by GeneArt (Thermo Fisher Scientific). Another synthetic gene, *MXRIN400FFQ*, with phenylalanine residues at 249, 254, and 278 mutated to alanine, and 18 glutamine residues between 150 and 200 amino acids (152, 155, 157, 158, 162, 163, 164, 166, 168, 170, 171, 175, 179, 181, 182, 185, and 194) mutated to alanine in the *MXRIN400Q* gene 5'-CGGGGTACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3'. In another set of PCRs, Mxr1p sequence was amplified using a common forward primer for amplification of first 150 or 250 bp of *MXR1*, which have an overlapping sequence of C-terminal region (1311–1329 base pairs) of *GFP* and overlapping sequence with the N-terminal region of (1–26 base pairs) *MXR1* gene 5'-CCTCAAAATGGTGGGAATGGTATCATCACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3' (K). The reverse primer for amplification of *MXRIN150* is sequence reverse complementary to 432 to 450 bp of *MXR1* gene with a downstream Myc tag sequence, stop codon, and a NotI restriction enzyme site 5'-ATAGTTTTAGCGGCCGCTACTCATGCAATCTATACATCATTATGAGAACC GTTGAGGAACTGTCGGTGG-3' (R1). For amplifying *MXRIN250*, the reverse primer for amplification of reverse primer is reverse complementary to 727 to 750 bp of *MXR1* gene with a downstream Myc-tag sequence, stop codon, and a NotI restriction enzyme site 5'-ATAGTTTTAGCGGCCGCTACTCATGCAATCTATACATCATTATGAGAACC GTTGAGGAACTGTCGGTGG-3' (R2). To get the GFP–MXR fusion product, an overlapping PCR was performed where the amplified GFP and *MXRIN150* or *MXRIN250* PCR products were used as templates, and primer pairs F and R1 or F and R2 were used. This fused PCR product was digested with KpnI and NotI restriction enzymes and

**Generation of Δmxr1-A-N150, Δmxr1-A-N250, and Δmxr1-A-N250***

To construct an N-terminal GFP-tagged *Mxr1N250* and N-terminal GFP-tagged *Mxr1N150*, first, the GFP sequence was amplified from *pEGFP-C2* (catalog no. 6083-1; Addgene) with the forward primer having N-terminal region (1–18 base pairs) of the GFP gene and an upstream KpnI restriction enzyme site 5'-CGGGGTACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3' (F) and the reverse primer complementary to C-terminal region (1311–1329 base pairs) of *GFP* and overlapping sequence with the N-terminal region of (1–26 base pairs) *MXR1* gene 5'-CCTCAAAATGGTGGGAATGGTATCATCACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3' (R). To construct an N-terminal GFP-tagged *Mxr1N250*, first, the GFP sequence was amplified from *pEGFP-C2* (catalog no. 6083-1; Addgene) with the forward primer having N-terminal region (1–18 base pairs) of the GFP gene and an upstream KpnI restriction enzyme site 5'-CGGGGTACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3' (F) and the reverse primer complementary to C-terminal region (1311–1329 base pairs) of *GFP* and overlapping sequence with the N-terminal region of (1–26 base pairs) *MXR1* gene 5'-CCTCAAAATGGTGGGAATGGTATCATCACCATGATGAGAACC GTTGAGGAACTGTCGGTGG-3' (R). To get the GFP–MXR fusion product, an overlapping PCR was performed where the amplified GFP and *MXRIN150* or *MXRIN250* PCR products were used as templates, and primer pairs F and R1 or F and R2 were used. This fused PCR product was digested with KpnI and NotI restriction enzymes and
cloned under GAPDH promoter in pGHYB (35) vector, followed by transformation of MXRN1250 in Δmxr1 and Δmxr1-A while MXRN1150 in Δmxr1-A. Positive colonies were screened for expression of GFP-tagged Mxr1N250 or Mxr1N150 using anti-GFP antibody.

**Generation of Δmxr1-N62, Δmxr1-N81, and Δmxr1-N109**

To construct an N-terminal GFP-tagged Mxr1N62, N-terminal GFP-tagged Mxr1N81, and N-terminal GFP-tagged Mxr1N109, first the GFP sequence was amplified from pEGFP-C2 (catalog no. 6083-1; Addgene) with the forward primer having N-terminal region (1–18 base pairs) of the GFP gene with an upstream KpnI restriction enzyme site 5'-CGGGGTACCATGGTGAAGGGCGAG-3' (F) and the reverse primer complementary to C-terminal region (1311–1329 base pairs) of GFP and overlapping sequence with the N-terminal region of (1–26 base pairs) MXRI gene 5'-CCAAAAGTTGGGTGATATGCTCTTCTTGTACAG CTCGTCCATG-3'. In another set of PCRs, Mxr1p sequence was amplified using a common forward primer for amplification of first 62, 81, or 109 bp of MXRI, which have an overlapping sequence of C-terminal region (1311–1329 base pairs) of GFP gene followed by N-terminal region (1–26 base pairs) of MXRI gene 5'-CATGGACAGCTGTAACAGATGAG CAATCTACCCAAAATTTTG-3' (F'). The reverse primer for amplification of MXRN62 is sequence reverse complementary to 169 to 186 bp of MXRI gene with a downstream stop codon, and a NotI restriction enzyme site 5'-ATAGTT TACCGGGCGCTATGAGACCTTCTTGTCG-3' (R1). For amplifying MXRN81, the sequence of reverse primer is reverse complementary to 223 to 243 bp of MXRI gene with a downstream stop codon, and a NotI restriction enzyme site 5'-ATAGTTTACGGCGCCCTTACATCGACGCTGAAT TTACG-3' (R2). For amplifying MXRN109, the sequence of reverse primer is reverse complementary to 312 to 327 bp of MXRI gene with a downstream stop codon, and a NotI restriction enzyme site 5'-ATAGTTTACGGCGCCCTTACATCGACGCTGAAT TTACG-3' (R3). To get the GFP–MXR fusion product, an overlapping PCR was performed where the amplified GFP and MXRN62, MXRN81, or MXRN109 GAPDH promoter products were used as templates, and primer pairs F and R1, F and R2, or F and R3 were used. This fused PCR product was digested with KpnI and NotI restriction enzymes and cloned under GAPDH promoter in pGHYB (35) vector, followed by transformation of MXRN1250, MXRN81, and MXRN109, in Δmxr1. Positive colonies were screened for expression of GFP-tagged Mxr1N62, Mxr1N81, or Mxr1N109 using anti-GFP antibody.

**Synthetic genes were cloned in pGHYB (35) with an N-terminal GFP tag under GAPDH promoter followed by transformation in Δmxr1 and screening by Western blotting with anti-GFP antibody. Sequence of synthetic genes will be made available upon request.**

**Data availability**

The RNA-Seq and DNA microarray data are available at NCBI GEO (https://www.ncbi.nlm.nih.gov/geo/) under the accession numbers GSE168677 and GSE146829, respectively.

**Supporting information**—This article contains supporting information.

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**Conflict of interest**—The authors declare that they have no conflicts of interest with the contents of this article.

**Abbreviations**—The abbreviations used are: 9aaTAD, nine-amino-acid TAD; ACS, acetyl-CoA synthetase; ALD, aldehyde dehydrogenase; ALD6-1Myc, Myc-tagged ALD6-1; cDNA, complementary DNA; DBD, DNA-binding domain; GEO, Gene Expression Omnibus; GO, Gene Ontology; MUT, methanol utilization; MXRE, Mxr1p response element; NCBI, National Center for Biotechnology Information; NLS, nuclear localization signal; PGK, phosphoglycerate kinase; qPCR, quantitative PCR; TAD, transactivation domain; YNB, yeast nitrogen base; YNBE, YNB and 1% ethanol; YNBM, YNB and 1% methanol; YPD, yeast extract, peptone, and dextrose.

**References**

1. Cereghino, J. L., and Cregg, J. M. (2000) Heterologous protein expression in the methylotrophic yeast Pichia pastoris. *FEMS Microbiol. Rev.* 24, 45–66
2. van der Klei, I. J., Yurimoto, H., Sakai, Y., and Veenhuis, M. (2006) The significance of peroxisomes in methanol metabolism in methylotrophic yeast. *Biochim. Biophys. Acta* 1763, 1453–1462
Key functional domains of Pichia pastoris Mxr1p

3. Peña, D. A., Gasser, B., Zanghellini, I., Steiger, M. G., and Mattanovich, D. (2018) Metabolic engineering of Pichia pastoris. Metab. Eng. 50, 2–15

4. Sahu, U. and Rangarajan, P. N. (2016) Regulation of acetate metabolism and acetyl Co-A synthetase 1 (ACS1) expression by methanol expression regulator 1 (Mxr1p) in the methylotrophic yeast Pichia pastoris. J. Biol. Chem. 291, 3648–3657

5. Sahu, U. and Rangarajan, P. N. (2016) Methanol expression regulator 1 (Mxr1p) is essential for the utilization of amino acids as the sole source of carbon by the methylotrophic yeast, Pichia pastoris. J. Biol. Chem. 291, 20588–20601

6. Lin-Cereghino, G. P., Godfrey, L., de la Cruz, B. J., Johnson, S., Khungsathiene, S., Tolstorukov, I., Yan, M., Lin-Cereghino, I., Veenhuis, M., Subramani, S., and Cregg, J. M. (2006) Mxr1p, a key regulator of the methanol utilization pathway and peroxisomal genes in Pichia pastoris. Mol. Cell. Biol. 26, 883–897

7. Sahu, U., Krishna Rao, K., and Rangarajan, P. N. (2014) Trim1p, a Zn(II) Cys6-type transcription factor, is essential for the transcriptional activation of genes of methanol utilization pathway, in Pichia pastoris. Biochem. Biophys. Res. Commun. 451, 158–164

8. Wang, X., Wang, Q., Wang, J., Bai, P., Shi, L., Shen, W., Zhou, M., Zhou, X., Zhang, Y., and Cai, M. (2016) Mt1 transcription factor mediates methanol signaling and regulates the alcohol oxidase 1 (AOX1) promoter in Pichia pastoris. J. Biol. Chem. 291, 6245–6261

9. Kumar, N. V., and Rangarajan, P. N. (2012) The zinc finger proteins Mxr1p and repressor of phosphoenolpyruvate carboxykinase (ROP) have the same DNA binding specificity but regulate methanol metabolism antagonistically in Pichia pastoris. J. Biol. Chem. 287, 34465–34473

10. Shi, L., Wang, X., Wang, J., Zhang, P., Qi, F., Cai, M., Zhang, Y., and Zhou, X. (2018) Transcriptome analysis of Δmig1Δmig2 mutant reveals their roles in methanol catabolism, peroxisome biogenesis and autophagy in methylotrophic yeast Pichia pastoris. Genes Genomics 40, 399–412

11. Lee, S. B., Kang, H. S., and Kim, T. S. (2013) Nrg1 functions as a global transcriptional regulator of genes of methanol utilization pathway, in Pichia pastoris. Biochem. Biophys. Res. Commun. 439, 501–505

12. Zhan, C., Yang, Y., Zhang, Z., Li, X., Liu, X., and Bai, Z. (2017) Transcription factor Mxr1 promotes the expression of Aox1 by repressing glycerol transporter 1 in Pichia pastoris. FEMS Yeast Res. 17, 1–10

13. Yang, V. W. (1998) Eukaryotic transcription factors : Identification, characterization. J. Nutr. 128, 2045–2051

14. Mitsis, T., Eftimiaidou, A., Bacopoulou, F., Velachakis, D., Chrousos, G., and Eliopoulos, E. (2020) Transcription factors and evolution: An integral part of gene expression (review). World Acad. Sci. J. 2, 3–8

15. Arnold, C. D., Nemeko, F., Woodfin, A. R., Wienererrother, S., Vlasova, A., Schleiffer, A., Pagani, M., Rath, M., and Stark, A. (2018) A high-throughput method to identify trans-activation domains within transcription factor sequences. EMBO J. 37, 1–13

16. Soniat, M., and Chook, Y. M. (2015) Nuclear localization signals for four distinct Karyopherin-β nuclear import systems. Biochem. J. 468, 353–362

17. Denis, C. L., and Young, E. T. (1983) Isolation and characterization of the positive regulatory gene ADR1 from Saccharomyces cerevisiae. Mol. Cell. Biol. 3, 360–370

18. Kranthi, B. V., Kumar, R., Kumar, N. V., Rao, D. N., and Rangarajan, P. N. (2009) Identification of key DNA elements involved in promoter recognition by Mxr1p, a master regulator of methanol utilization pathway in Pichia pastoris. Biochim. Biophys. Acta 1789, 460–468

19. Kranthi, B. V., Kumar, H. R. V., and Rangarajan, P. N. (2010) Identification of Mxr1p-binding sites in the promoters of genes encoding dihydroxyacetone synthase and peroxin 8 of the methylotrophic yeast Pichia pastoris. Yeast 27, 705–711

20. Parua, P. K., Ryan, P. M., Trang, K., and Young, E. T. (2012) Pichia pastoris 14-3-3 regulates transcriptional activity of the methanol inducible transcription factor Mxr1 by direct interaction. Mol. Microbiol. 85, 282–298

21. Cardarelli, S., D'Amici, S., Tassone, P., Tramonti, A., Uccelli, D., Mancini, P., and Saliola, M. (2016) Characterization of the transcription factor encoding gene, KlADRI: Metabolic role in Kluyveromyces lactis and expression in Saccharomyces cerevisiae. Microbiology (Reading) 162, 1933–1944

22. Barbay, D., Mačáková, M., Sützl, L., De, S., Mattanovich, D., and Gasser, B. (2021) Two homologs of the Cat8 transcription factor are involved in the regulation of ethanol utilization in Komagatella phaffii. Curr. Genet. 67, 641–661

23. Tanaka, M., Clouston, W. M., and Herr, W. (1994) The Oct-2 glutamine-rich and proline-rich activation domains can synergize with each other or duplicates of themselves to activate transcription. Mol. Cell. Biol. 14, 6046–6055

24. Xiao, H., and Jeang, K. T. (1998) Glutamine-rich domains activate transcription in yeast Saccharomyces cerevisiae. J. Biol. Chem. 273, 22873–22876

25. Courey, A. J., Holtzman, D. A., Jackson, S. P., and Tjian, R. (1989) Synergistic activation by the glutamine-rich domains of human transcription factor Spl. Cell 59, 827–836

26. Atanesyan, L., Günther, V., Dichtl, B., Georgiev, O., and Schaffner, W. (2012) Polyglutamine tracts as modulators of transcriptional activation from yeast to mammals. Biol. Chem. 393, 63–70

27. Escher, D., Bodmer-Gravas, M., Barberis, A., and Schaffner, W. (2000) Conservation of glutamine-rich transcription activation function between yeast and humans. Mol. Cell. Biol. 20, 2774–2782

28. Piskacek, S., Gregor, M., Nemethova, M., Grabner, M., Kovarik, P., and Piskacek, M. (2007) Nine-amino-acid transactivation domain: Establishment and prediction utilities. Genomics 89, 756–768

29. Piskacek, M. (2009) Common transactivation Motif 9aaTAD recruits multiple general co-activators TAF9, MED15, CBP and p300. Nat. Preced. https://doi.org/10.1038/npre.2009.2488.2

30. Sandholzer, J., Hoeth, M., Piskacek, M., Mayer, H., and de Martin, R. (2007) A novel 9-amino-acid transactivation domain in the C-terminal part of Sox18. Biochem. Biophys. Res. Commun. 360, 370–374

31. Piskacek, M., Havelka, M., Rezacova, M., and Knight, A. (2016) The 9aaTAD transactivation domains: From Gal4 to p53. PLoS One 11, 1–16

32. Karamanoglou, M., Karamanoglou, F. E., and Inan, M. (2016) Functional analysis of alcohol dehydrogenase (ADH) genes in Pichia pastoris. Biotechnol. Lett. 38, 463–469

33. Truant, R., and Cullen, B. R. (1999) The arginine-rich domains present in human immunodeficiency virus type 1 Tat and Rev function as direct importin β-dependent nuclear localization signals. Mol. Cell. Biol. 19, 1210–1217

34. Weninger, A., Glieder, A., and Vogl, T. (2015) A toolbox of endogenous and heterologous nuclear localization sequences for the methylotrophic yeast Pichia pastoris. FEMS Yeast Res. 15, 4–7

35. Yang, J., Nie, L., Chen, B., Liu, Y., Kong, Y., Wang, H., and Diao, L. (2014) Hygromycin-resistance vectors for gene expression in Pichia pastoris. Yeast 31, 115–125