

## Transcriptional Regulation of *xyn2* in *Hypocrea jecorina*

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Received 24 May 2002/Accepted 10 November 2002

The xylanase system of the filamentous fungus *Hypocrea jecorina* (*Trichoderma reesei*) consists of two specific xylanases, Xyn1 and Xyn2, which are simultaneously expressed during growth on xylan but respond differentially to low-molecular-weight inducers. Using *in vivo* footprinting analysis of xylan-induced and noninduced mycelia, we detected two adjacent nucleotide sequences (5'-AGAA-3' on the noncoding strand and 5'-GGGTAAATTGG-3', referred to as the xylanase-activating element [XAE], on the coding strand, respectively) to bind proteins. Among these, binding to the AGAA-box is only observed under noninduced conditions, whereas binding to XAE is constitutive. Electrophoretic mobility shift assay with heterologously expressed components of the *H. jecorina* Hap2/3/5 protein complex and the cellulase regulator Ace2 suggests that these two transactivators form the protein complex binding to XAE. *H. jecorina* transformants, containing correspondingly mutated versions of the *xyn2* promoter fused to the *Aspergillus niger* *gox4* gene as a reporter, revealed that the elimination of protein binding to the AGAA-box resulted in a threefold increase in both basal and induced transcription, whereas elimination of Ace2 binding to its target in XAE completely eliminated transcription under both conditions. Destruction of the CCAAT-box by insertion of a point mutation prevents binding of the Hap2/3/5 complex *in vitro* and results in a slight increase in both basal and induced transcription. These data support a model of *xyn2* regulation based on the interplay of Hap2/3/5, Ace2 and the AGAA-box binding repressor.

$\beta$ -1,4-Xylans are heteropolysaccharides that have a backbone of  $\beta$ -1,4-linked xylopyranosyl residues, to which side groups such as D-glucuronic acid, L-arabinose, *p*-coumaric acid, and ferulic acid are attached and which constitute 20 to 35% of the roughly 830 Gt of annually formed renewable plant biomass (33). Enzymes capable of degrading the xylan backbone are formed by various microorganisms and comprise endoxylanases (EC 3.2.1.8) and  $\beta$ -xylosidase (EC 3.2.1.91) (15). Among these, the xylanases of the ascomycete *Hypocrea jecorina* (anamorph *Trichoderma reesei*) have received strong attention because of their application in the pulp and paper and feed industry (4).

Although the biochemistry of xylan degradation by *H. jecorina* has now been studied in detail (2, 10, 23, 25), the mechanism by which the fungus regulates the formation of its xylanases is still largely unsolved (44). *H. jecorina* forms two specific endo- $\beta$ -1,4-xylanases, Xyn1 and Xyn2 (EC 3.2.1.8), and the respective genes (*xyn1* and *xyn2*) have been cloned (27, 34). Expression of *xyn1* is induced by D-xylose and is repressed by glucose in a Cre1-dependent manner (18, 44), whereas the expression of *xyn2* is partially constitutive and further induced by xylobiose, xylan, cellulose, and sophorose (44). However, the regulatory circuits governing *xyn1* and *xyn2* gene expression

have not yet been elucidated. In *Aspergillus niger*, expression of the xylanolytic system is regulated by the zinc binuclear cluster type transcriptional regulator XlnR (36), in which it appears to be a central regulator, since it controls the expression of more than 10 genes not only involved in the degradation of xylan but also in xylose metabolism and cellulose degradation (6, 9, 35). The cloning of a *xlnR* homologue of *H. jecorina* has not yet been reported, but nucleotide motifs resembling the consensus for XlnR binding (5'-GGCAAA-3') are present in the *xyn1* and *xyn2* 5'-upstream sequences (unpublished data). On the other hand, Aro et al. (1) recently reported that a *H. jecorina* mutant, in which the gene encoding the cellulase regulator *ace2* had been disrupted, exhibited reduced expression of *xyn2*.

Using promoter deletion analysis, we previously reported that a 55-bp fragment of the *xyn2* promoter contains all of the information necessary for regulating *xyn2* gene expression (44). Here we identify the nucleotide sequences within these 55 bp that are essential for binding of proteins and responsible for *xyn2* regulation by using both *in vitro* and *in vivo* strategies. Furthermore, we show that both basal transcription and induction of *xyn2* depends on the binding of the Hap2/3/5 complex (42) and Ace2 to an undecameric motif (5'-GGGTAAATTG G-3'; the xylanase-activating element [XAE]) and that this binding is counteracted by an as-yet-unknown DNA-binding protein (complex) binding to an AGAA-box immediately upstream of XAE.

### MATERIALS AND METHODS

**Microbial strains and plasmids.** *H. jecorina* QM9414 (ATCC 26921) and *H. jecorina* RUT C30 (ATCC 56756) were used throughout the present study. *H. jecorina* TU-6 (8), a *pyr4*-null mutant of QM9414, was used as recipient strain for *pyr4*-mediated cotransformation experiments. The strains were maintained on

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TABLE 1. Oligonucleotides used throughout this study<sup>a</sup>

Name	Sequence	Nucleotide range (reference)
CKT087	5'-ATCGGAGTCGACTCGCATCCG-3'	-1081 to 1059 (43)
CKT088	5'-ATGCTAGCGTTGATGTCTTCTGCTTCAGC-3'	-22 to -1 (43)
Prxyn2f	5'-TGATGAAAGGAGAACAACCTTCTAGACTGGGTAATTTGGTCAAT-3'	-249 to -207
Prxyn2f	5'-AGCGGCTGGCCATTGACCAATTTACCCAGTCTAGAAGTTGTTCTCC-3'	-196 to -241
Prxyn2M2f	5'-TGATGAAAGGAGAACAACCGGAGAGACTGGGTAATTTGGTCAAT-3'	-249 to -207
Prxyn2M2r	5'-AGCGGCTGGCCATTGACCAATTTACCCAGTCTCTCCGTTGTTCTCC-3'	-196 to -241
Prxyn2bf	5'-CTAGACTGGGTAATTTGGTCAATGGCCAGCCGCTCGGCCGTGCGGAGACGAGGCA-3'	-229 to -175
Prxyn2br	5'-AGCTTGCCCTCGTCTCCGCACGGCCGAGCGGCTGGCCATTGACCAATTTACCCAGT-3'	-171 to -225
Prxyn2bM3f	5'-CTAGACTGGGTAATTTGGTCAATGGCCAGCCGCTCGGCCGTGCGGAGACGAGGCA-3'	-229 to -175
Prxyn2bM3r	5'-AGCTTGCCCTCGTCTCCGCACGGCCGAGCGGCTGGCCATTGACCTTTTACCCAGT-3'	-171 to -225
Prxyn2bM4f	5'-CTAGACTGGGTTTATTGGTCAATGGCCAGCCGCTCGGCCGTGCGGAGACGAGGCA-3'	-229 to -175
Prxyn2bM4r	5'-AGCTTGCCCTCGTCTCCGCACGGCCGAGCGGCTGGCCATTGACCAATAAACCCAGT-3'	-171 to -225
Prxyn2bM5f	5'-CTAGACTGTGTAATTTGGTCAATGGCCAGCCGCTCGGCCGTGCGGAGACGAGGCA-3'	-229 to -175
Prxyn2bM5r	5'-AGCTTGCCCTCGTCTCCGCACGGCCGAGCGGCTGGCCATTGACCAATTTACACAGT-3'	-171 to -225
<i>xyn2</i> P1n	5'-CTTCTCATATGTATCGGAGCAGAAG-3'	-357 to -333
<i>xyn2</i> P2n	5'-ATCTATGTGAGCTGACTCGAGACGGC-3'	-326 to -301
<i>xyn2</i> P3n:	5'-AGCTGACTCGAGACGGCTGAGACAGCA-3'	-317 to -291
<i>xyn2</i> P1c	5'-CTAGGAGTTGTTGTGTCTTTTG-3'	-25 to -46
<i>xyn2</i> P2c	5'-GTTGTTTAGGTTTGGCTTCTTG-3'	-83 to -104
<i>xyn2</i> P3c	5'-CAGGTTTGGCTTCTTGAGTCTTTAGG-3'	-89 to -114
Prxyn2af	5'-TGATGAAAGGAGAACAACCTTCTAGACTG-3'	-249 to -222
Prxyn2ar	5'-TGACCAGTCTAGAAGTTGTTCTCTTTTC-3'	-222 to -245
Prxyn2aM2f	5'-TGATGAAAGGAGAACAACCGGAGACTG-3'	-249 to -222
Prxyn2aM2r	5'-TGACCAGTCTCTCCGTTGTTCTCTTTTC-3'	-222 to -245
Prxyn1.1f	5'-TTGGCAGGCTAAATGCGACATCTTAGCCGGA-3'	-430 to -400
Prxyn1.1r	5'-TGCATCCGGCTAAGATGTCGCATTTAGCTG-3'	-396 to -421

<sup>a</sup> Oligonucleotides already used in earlier studies are referenced. Positions of the oligonucleotides in the respective promoter are given. Underlined letters indicate bases added for labeling or generating restriction enzyme sites, and double underlined bases indicate introduced point mutations.

malt agar which contained 5 mM uridine in the case of TU-6. *Escherichia coli* JM109 (41) was used for the propagation of vector molecules. Plasmids pFG1 (8), pAT3 (34), and pSJ3 (17) were obtained from our department stock.

**Fungal growth, induction of xylanases, and preparation of cell extracts.** *H. jecorina* was grown on glucose and xylan or induced by sophorose and xylobiose, respectively, as described previously (18, 44). Xylan from oat spelts (Sigma, Steinheim, Germany) was used throughout. Cell extracts were prepared as described previously (30).

**Construction of pLW reporter plasmids.** The pLW reporter plasmid series was developed from plasmid pSJ3 by fusing the *H. jecorina xyn2* 5' noncoding regions (-1 to -850) to the *gox4* (glucose oxidase-encoding) structural gene of *A. niger* as reporter. To construct pLW-WT, primers CKT087 and CKT088 (Table 1) were used to amplify a 1,081-bp fragment from the *xyn2* 5' noncoding sequences, thereby also generating an additional *NheI* terminal site at the 3' end. Amplification was performed with *Taq* polymerase (Promega, Madison, Wis.) and pAT3 as the template DNA in a Biometra thermocycler by applying 30 cycles of 1 min at 95°C, 1 min at 56°C, and 1 min at 74°C. The PCR product was cloned into pGEM-T (Promega) to yield pLW1, sequenced, and digested with *Sall* and *NheI*. Due to a *Sall* restriction site within the amplified sequence and the gained *NheI* site at the 3' end, a 0.8-kb fragment was released and thereafter fused to *gox4* of pSJ3 by replacing the *nag1* 5'-noncoding region (cut with *XhoI/XbaI*). To yield pLWm1, a four-primer PCR mutagenesis strategy was followed: in a first step, two overlapping fragments containing the intended point mutations were amplified with the primers CKT087/*xyn2*prpm2r and CKT088/*xyn2*prpm2f (Table 1), respectively, by using pAT3 as a DNA template and Herculase (Stratagene, La Jolla, Calif.) lacking terminal transferase activity. Derived fragments were thereafter subjected as a template to a second PCR with primers CKT087 and CKT088 by using *Taq* polymerase. PCR protocols and further processing of the derived amplicon essentially followed the procedure described for constructing pLW-WT.

All further mutations of the *xyn2* 5'-noncoding region were performed in pLW1 as a template and thereafter transferred into pSJ3 as described above. To obtain pLWM2, pLW1 was cleaved with *XbaI* and *HindIII*; protruding ends were blunted with the mung bean exonuclease (Promega) by applying 0.12 U (37°C, 30 min)/μg of DNA and thereafter religated. To construct pLWM3, pLWM4, and pLWM5, the synthetic oligonucleotides Prxyn2bM3f and Prxyn2bM3r; Prxyn2bM4f and Prxyn2bM4r; and Prxyn2bM5f and Prxyn2bM5r (Table 1) containing the respective point mutations were annealed, thereby producing appro-

prate 5'-protruding ends for an insertion into the *XbaI*- and *HindIII*-cleaved plasmid pLW1.

**Isolation and manipulation of nucleic acids.** Genomic DNA was isolated as described previously (8). After electrophoretic separation, DNA was blotted onto Hybond-N membranes (Amersham Pharmacia Biotech, Little Chalfont, United Kingdom) and hybridized at 64°C for 20 h according to standard protocols (28). Washing was performed with 0.2× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate)-0.1% (wt/vol) sodium dodecyl sulfate (SDS) at 68°C (2 times for 15 min each time). Standard methods were used for plasmid isolation, restriction enzyme digestion, and random priming (28).

**DNA transformations.** *E. coli* transformations were carried out according to standard techniques (28). Transformation of *H. jecorina* TU-6 was carried out according to an optimized protocol for particle bombardment (11), applying cotransformation of pFG1 with the respective reporter constructs. For each construct, at least 30 uridine prototrophs were purified to mitotic stability, and their DNA was isolated and examined for integration and copy number by Southern analysis. Cotransformation frequency varied from 90 to 100%, yielding copy numbers between 1 and 5, integrated into ectopic loci.

**Determination of gene copy number and integration locus in transformants.** Southern hybridization was carried out as described by Sambrook et al. (28). Chromosomal DNA was digested with *PstI* and hybridized with a [ $\alpha$ -<sup>32</sup>P]dCTP-labeled 440-bp fragment bearing 312 bp of the *xyn2* promoter and 128 bp of the *gox4* gene. The endogenous *xyn2* gene and the integrated vector copies were quantified by imager analysis, and the values obtained were normalized to the length of the labeled probe.

**Glucose oxidase assay.** Glucose oxidase activity was assayed as described previously (17). One unit of activity is defined as the amount of enzyme which oxidizes 1 μmol of glucose per min at pH 5.8 and 25°C.

**EMSA.** Oligonucleotides used for electrophoretic mobility shift assay (EMSA) were annealed with their complementary oligonucleotides (Table 1) and end labeled with [ $\alpha$ -<sup>32</sup>P]dCTP by using Sequenase version 2.0 (Amersham Pharmacia Biotech). The resulting double-stranded oligonucleotides were purified by non-denaturing polyacrylamide gel electrophoresis (PAGE). The binding assay and PAGE were performed essentially as described previously (30). Binding was achieved by incubating 100 μg of protein of the cell extract with 5 ng of labeled fragment (15 min, 0°C). For competition experiments, unlabeled synthetic oligonucleotides were used in a 50- or 100-fold molar excess. Unlabeled oligonucleotides were annealed with the complementary synthetic oligonucleotide as

described by Strauss et al. (31). After annealing, double strands were filled in by using Sequenase version 2.0 (Amersham Pharmacia Biotech). Fragments bearing DNA-binding regions of *A. niger* XlnR and *H. jecorina* Hap2, Hap3, Hap5 and Ace2 were expressed as glutathione *S*-transferase (GST) fusion proteins in strain BL21 as described by van Peij et al. (36), Zeilinger et al. (42), and Aro et al. (1), respectively. The resulting GST fusion proteins and the proteins after thrombin cleavage were used in EMSAs at the following concentrations: GST-XlnR (0.1, 0.2, 0.5, and 1  $\mu$ g), XlnR (0.05, 0.1, 0.2, and 0.5  $\mu$ g); Hap2, Hap3, and Hap5 (0.5  $\mu$ g each); GST-Ace2 (1  $\mu$ g); and Ace2 (0.5  $\mu$ g). Oligonucleotide PRxyn1.1 (Table 1) was used as a positive control for EMSA with XlnR.

**In vivo genomic footprinting via ligation-mediated PCR.** Methylation of genomic DNA was performed at 30°C in a shaking water bath by transferring 18-ml aliquots of *H. jecorina* cultures grown on glucose, glycerol, or xylan for the period indicated to 100-ml Erlenmeyer flasks and pulsing them with 40  $\mu$ l of DMS in 2 ml of 200 mM methyl ethanesulfonate MES buffer (pH 5.5) for 2 min. Methylation was stopped by addition of 50 ml of ice-cold TLE $\beta$  buffer (10 mM Tris [pH 8], 1 mM EDTA, 300 mM LiCl, 2% [vol/vol]  $\beta$ -mercaptoethanol). Mycelial samples were filtered and washed twice with 50 ml of TLE $\beta$  buffer, and genomic DNA was extracted according to a standard protocol (8). The extracted methylated DNA was cleaved at methylated guanine and adenine residues by incubation of 20  $\mu$ l of DNA with 1.25  $\mu$ l of 0.5 M HCl for 1.5 h on ice. After precipitation with ethanol, the DNA was dissolved in 48  $\mu$ l of bidistilled water and incubated at 90°C for 30 min with 2  $\mu$ l of 1 M NaOH, followed by a second precipitation with ethanol. Finally, DNA was dissolved in 20  $\mu$ l of Tris-EDTA. In vitro methylation and cleavage of genomic DNA was performed as described by Mueller and Wold (22). Methylated and cleaved DNA was analyzed by ligation-mediated PCR as described by Garrity and Wold (5), and as modified by Wol-schek et al. (38), by using *Vent* polymerase (NEB, Beverly, Mass.). To visualize the noncoding strand, the primers xyn2P1n, xyn2P2n, and xyn2P3n were used, and to visualize the coding strand, the primers xyn2P1c, xyn2P2c, and xyn2P3c were used.

**Chromatin analysis.** Micrococcal nuclease (MNase)-based mapping of chromatin organization was carried out as described previously (7). Mycelia were harvested by filtration, pressed dry with filter paper, frozen in liquid nitrogen, and ground to a fine powder. Then, 200-mg portions of the mycelial powder were suspended in 2 ml of nuclease digestion buffer (15 mM Tris-HCl [pH 7.5], 250 mM sucrose, 60 mM KCl, 15 mM NaCl, 0.05 mM CaCl<sub>2</sub>, 3 mM MgCl<sub>2</sub>, 0.5 mM dithiothreitol). Next, 200- $\mu$ l aliquots of the digestion mixture were incubated with MNase for 5 min at 37°C by using various MNase concentrations of between 2 U (100 U/g of mycelium) and 0.01 U (0.5 U/g of mycelium). The reaction was terminated by adding 200  $\mu$ l of 40 mM EDTA-2% SDS, followed by two rounds of phenol-chloroform extraction and ethanol precipitation. The resuspended DNA was treated with RNase A and passed through a Sephacryl S-200 microspin column (Amersham Pharmacia Biotech). For the preparation of the naked DNA control, 20  $\mu$ g of purified chromosomal DNA were digested with appropriate concentrations of MNase as described above and then processed as for the chromatin samples.

**Indirect end-labeling analysis.** Indirect end labeling was carried out as described previously (39). After secondary restriction enzyme digestion with *Nsi*I, the samples were electrophoresed in 1.2% agarose gels in 1 $\times$  TAE, transferred to nylon membranes (Hybond-N<sup>+</sup>; Amersham Pharmacia Biotech) in 10 $\times$  SSC, and hybridized at 42°C for 20 h in a hybridization solution containing 50% formamide, 5 $\times$  SSPE, 5 $\times$  Denhardt solution, 0.5% SDS, 10% (wt/vol) dextran sulfate, and 200  $\mu$ g of denatured herring sperm DNA/ml. Labeling of the specific probe (0.4-kb *Hind*III/*Nsi*I fragment isolated from plasmid pAT3) was performed by random priming. The autoradiographs were analyzed by calculating the  $R_f$  values for each band.

## RESULTS

**Identification of in cis-acting elements involved in the regulation of xyn2 expression.** The *xyn2* transcript is abundantly accumulating during growth on xylan, but a basal level of expression can also be observed during growth on glycerol and glucose, two carbon catabolite-repressing carbon sources (Fig. 1A). Regulation of *xyn2* transcription is not subject to carbon catabolite repression but is strictly dependent on induction since addition of the disaccharide xylobiose triggers its expression similarly on glucose and glycerol in a Cre1 negative background (strain *H. jecorina* RUT C-30; Fig. 1B). All of this

regulation is conferred by a short 55-bp stretch of the *xyn2* promoter (44). To identify the in cis-acting regulatory motifs within this 55-bp *xyn2* promoter fragment (−235 to −180), we performed in vivo genomic footprinting of the respective nucleotide area by using mycelia pregrown on glycerol and then replaced on medial containing glucose, xylan, or glycerol as carbon source. Under all three conditions, a clear, distinct protection pattern was detected on the noncoding strand (Fig. 1C), whereas no protection was observed on the coding strand (data not shown). Among the protected nucleotides, A<sub>−215</sub> within the ATTGG (CCAAT) motif was protected under all conditions, thus corroborating previous in vitro data on the potential involvement of the CCAAT-box in *xyn2* regulation (44). In addition, one purine base upstream of the CCAAT-box (−219) was also protected under all conditions. Further upstream at −231 to −228, an AGAA tetranucleotide motif was fully protected under noninducing conditions, whereas only a weak protection of G<sub>−229</sub> within this motif was evident under inducing conditions. Furthermore, an increase in the degree of methylation of two guanine bases located immediately upstream of the AGAA-box (G<sub>−232</sub> and G<sub>−235</sub>) was also detected only under noninducing conditions.

These data revealed that a part of the *xyn2* promoter is constitutively bound by DNA-binding proteins. This would imply that the *xyn2* promoter is permanently accessible to its in trans-acting factors. To prove this directly, we investigated the chromatin structure of *xyn2* by MNase treatment by using indirect end labeling with an 0.4-kb *Hind*III/*Nsi*I fragment of the *xyn2* gene as a probe (Fig. 1D). MNase treatment yielded a ladder of nucleosome-specific bands with a nucleosomal repetition length of 170 ( $\pm$ 9) bp upstream of the previously identified regulatory sequences (above −235). This array of at least two positioned nucleosomes is followed by a nucleosome-free region, emphasized by identical MNase patterns as in the naked DNA control, which spans the detected regulatory nucleotide motifs and the TATA-box and reaches even into the structural gene. No differences were evident between induced and noninduced conditions, thus implying that the TATA-box is permanently accessible, which is in accordance with the constitutive transcription of *xyn2*.

**In vitro characterization of protein binding to the AGAA-box.** To verify the binding of regulatory proteins to the identified nucleotide motifs, EMSAs were carried out by using cell extracts prepared from *H. jecorina* mycelia grown on glucose or xylan. Synthetic oligonucleotides (Table 1 and Fig. 2A), covering the whole promoter area for which methylation protection had been observed were used as probes. The 54-bp oligonucleotide Prxyn2, resembling the wild-type *xyn2* promoter area between −196 to −249, yielded one major protein-DNA complex with increased mobility under inducing conditions. This is in accordance with the findings that the AGAA-box and two adjacent 5' guanines are only bound by proteins under noninducing conditions (Fig. 2B, lanes 1 to 3). More direct evidence was obtained by EMSAs with oligonucleotide Prxyn2a, which represents a *xyn2* promoter fragment spanning from −222 to −249 and thus only bearing this motif (Fig. 2C, lanes 1 to 3). Only extracts from glucose-grown mycelia resulted in complex formation. To provide evidence that binding to the AGAA-box is sufficient to account for the differences in mobility observed in the binding of cell extracts from induced-



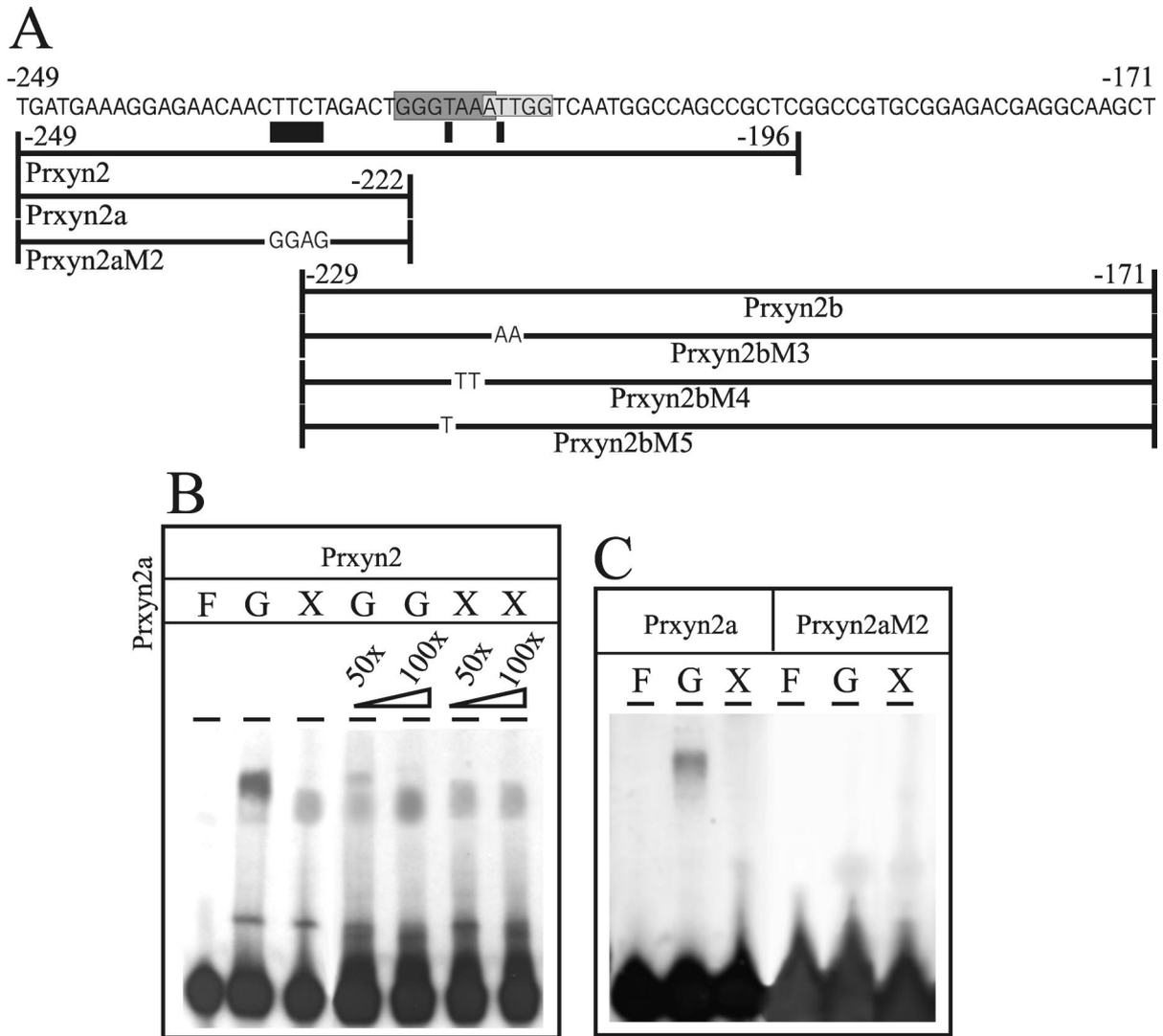


FIG. 2. (A) Schematic drawing of the oligonucleotides used in EMSAs. The given sequence spans the whole area of the previously identified *xyn2* 5'-noncoding region responsible for regulation of transcription. Potential regulatory elements are shaded, whereas the new motif identified by methylation protection in vivo footprinting is underlined. (B and C) EMSA analysis with 100  $\mu$ g of cell extract derived from *H. jecorina* mycelia replaced on either 1% (wt/vol) glucose (G, repressing conditions) or 1% (wt/vol) xylan (X, inducing conditions) and 5 ng of labeled oligonucleotides as probes: Prxyn2 (B) and Prxyn2a and Prxyn2aM2 (C). "F" indicates free probe; unlabeled Prxyn2a was used as a competitor in 50- or 100-fold molar excess (B, lanes 4 to 7).

and noninduced conditions to the wild-type promoter fragment Prxyn2, competition experiments with 50- and 100-fold molar excesses of unlabeled Prxyn2a over labeled Prxyn2 were performed; these experiments showed that the competing oligonucleotide titrates away the protein(s) binding to the AGAA-box from the labeled wild-type promoter fragment, causing the complexes from induced- and noninduced conditions to migrate with similar mobility in an EMSA (Fig. 2B).

In order to investigate whether the AGAA-box is sufficient and essential for this protein binding, EMSAs were performed with a mutated version of Prxyn2a, in which the AGAA-box was mutated to CTCC, whereas the 5'-upstream Gs had been left intact (oligonucleotide Prxyn2aM2; Table 1 and Fig. 2A). These data show that a replacement of the AGAA sequence in the noncoding strand by CTCC completely impaired protein

binding (Fig. 2C, lanes 4 to 6). We conclude from these data that the AGAA-box is the essential core motif responsible for protein binding in this area of the *xyn2* promoter and that binding under induced and noninduced conditions differs only in the binding of protein(s) to the AGAA-box under noninducing conditions.

**The *xyn2* promoter contains a regulatory element that is bound by Hap2/3/5 and Ace2.** The permanent protection of A<sub>-215</sub> within the CCAAT-box of the *xyn2* promoter is consistent with previous findings of constitutive formation of a protein-DNA complex with this motif (44). Interestingly, the inverted CCAAT-box (ATTGG) in the *xyn2* promoter is joined 5' by a GGGTAA motif, which closely resembles the binding site for the xylanase activator XlnR (36) and the cellulase regulator Ace2 (1). Both motifs together (5'-GGGTAAATT

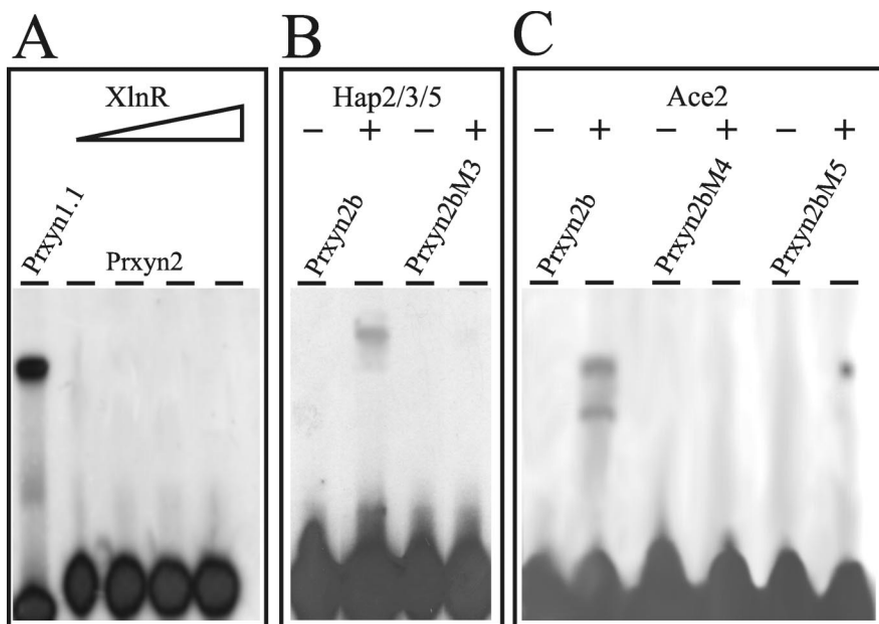


FIG. 3. (A) Analysis of the involvement of a XlnR-like factor in binding to the *xyn2* 5'-noncoding region. EMSA analysis with labeled oligonucleotide Prxyn2 as a probe and increasing amounts of recombinant GST-XlnR (lanes 2 to 4; 0.05, 0.1, 0.2, and 0.5  $\mu$ g) is shown. As a positive control, oligonucleotide Prxyn1.1 and 0.1  $\mu$ g of fusion protein were mixed in the binding reaction (lane 1). (B) Reconstitution of the CCAAT-binding activity on the *xyn2* promoter with recombinant GST-Hap2/3/5 (0.5  $\mu$ g each). EMSA was performed with labeled oligonucleotide Prxyn2b (lanes 1 and 2) and Prxyn2bM3 (lanes 3 and 4). (C) Analysis of the involvement of Ace2 in binding to the *xyn2* 5' regulatory region. EMSA analysis with labeled oligonucleotide Prxyn2b (lanes 1 and 2) and with oligonucleotides Prxyn2bM4 and Prxyn2bM5 (lanes 3 to 6) as probes and 1  $\mu$ g of recombinant GST-Ace2. -, Free probe; +, addition of the respective fusion proteins to the binding reaction.

GG-3') also bear a striking resemblance to an inverted version of the cellulase-activating element (CAE) in the *H. jecorina* *cbh2* promoter (5'-ATTGGGTAATA-3'), which is responsible for the induction of *cbh2* gene transcription by cellulose and sophorose (43).

To investigate whether XlnR or Ace2 bind to the 5'-GGTAAATTGG-3' motif, EMSAs with a heterologously expressed fragment of *A. niger* XlnR (36) containing the zinc-finger domain were performed with oligonucleotide Prxyn2. However, in contrast to a positive control experiment with oligonucleotide Prxyn1.1 (spanning a fragment of the *xyn1* upstream regulatory region bound by XlnR [C. Wacenovskiy and R. L. Mach, unpublished data]), complex formation was not observed with the GST fusion protein (Fig. 3A) or with the XlnR zinc-finger domain alone, irrespective of the amount of protein used. In contrast, by using the overexpressed zinc-finger domain of Ace2 (1), both the GST-Ace2 fusion (Fig. 3C, lanes 1 and 2) and the zinc-finger region alone showed protein-DNA binding in an EMSA. Oligonucleotides bearing mutations previously identified to decrease protein binding to the CAE (AA<sub>-217/-218</sub> to TT; PRxyn2M4 [43]) and a G<sub>-221</sub>-to-T mutation (essential for Ace2 binding to the *cbh1* promoter; Prxyn2M5 [1]) led to a complete loss of Ace2 binding. We thus conclude that Ace2 binds to the GGTAAT motif 5'-adjacent of the inverted CCAAT-box in the *xyn2* promoter.

Recently, Zeilinger et al. (42) demonstrated that the Hap2/3/5 protein complex binds to the inverted CCAAT-box (ATTGG) within the CAE of *cbh2*. In order to prove that this is also the case in the *xyn2* promoter, oligonucleotide PRxyn2b and the *H. jecorina* Hap subunits 2, 3, and 5 were subjected to

EMSA. DNA-protein complex formation was clearly observed and abolished by introduction of a TT<sub>-214/-215</sub>-to-AA mutation within the CCAAT-box (oligonucleotide Prxyn2M3; Fig. 3B). From these data it can be concluded that the CAE-like motif in the *xyn2* promoter is bound by the Hap2/3/5 protein complex and Ace2 but not XlnR.

**Mutations impairing protein binding in vitro lead to a loss of distinct regulatory functions in *xyn2* gene expression in vivo.** To investigate whether the protein-binding motifs identified above indeed confer regulation of *xyn2* transcription in vivo, the promoter mutations studied above were introduced into 850 bp of the 5'-noncoding region of the *xyn2* promoter (pLW-WT) and fused to the *A. niger* *gox4* (glucose oxidase-encoding) gene as a reporter. Except for a deletion from -171 to -229, which eliminates all putative elements identified (pLW-M2), the mutations inserted essentially resemble those used for the analysis of protein-DNA complex formation in vitro (pLWM1 = PRxyn2aM2; pLWM3 = PRxyn2M3; pLWM4 = PRxyn2M4; pLWM5 = PRxyn2M5; Fig. 4A). The different constructs were introduced into *H. jecorina* TU-6 by cotransformation with plasmid pFG1 (8) employing biolistic bombardment (11).

Three transgenic strains bearing ectopic single-copy integrations of the wild-type construct (LW-WT) or each mutated construct (LWM1-5) were investigated (Fig. 4B). As expected, deletion of the complete nucleotide area between -171 to -229 resulted in a loss of transcription both on noninducing (glycerol), as well as on inducing (xylan), carbon sources. However, the same effect was obtained when mutation 4 (pLWM4), which interfered with the binding of Ace2, was introduced,

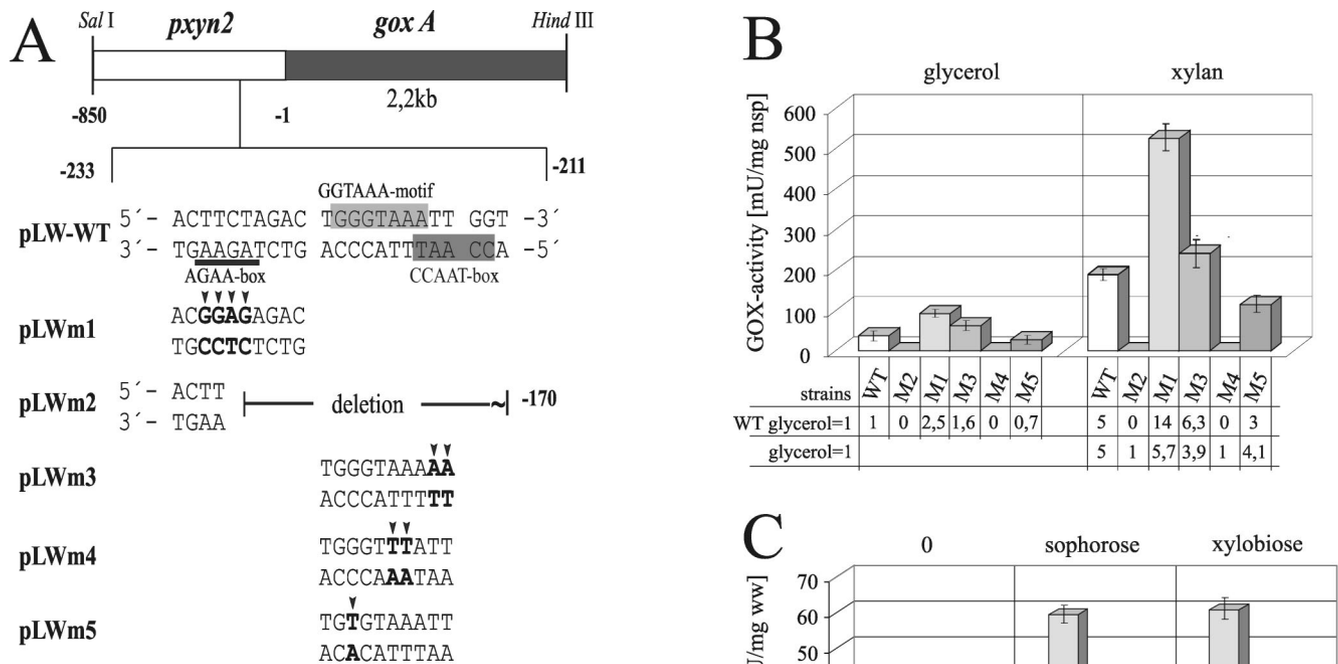


FIG. 4. (A) Schematic representation of the reporter vectors used for transformation and the respective mutations in the *xyn2* 5'-non-coding region. Point mutations are indicated by arrows; putative motifs are boxed or underlined. (B) Comparison of glucose oxidase (GOX) activity given in milliunits per mg of NaOH-soluble protein measured after cultivation of the strains LW-WT, LWM1, LWM2, LWM3, LWM4, and LWM5 on glycerol (non inducing conditions) and xylan (inducing conditions) for 24 h. The results are the means of two independent experiments performed with three different single-copy integration strains; error bars indicate the standard deviation. Factors representing regulatory switches were either calculated based on the GOX activity obtained from the wild-type reporter strains (WT glycerol = 1) or from the respective reporter strains (glycerol = 1) grown on glycerol. (C) Comparison of GOX activities given in milliunits per milligram (wet weight) of replaced mycelia, measured 12 h after replacement of the strains LW-WT, LWM1, LWM3, and LWM5 on media without a carbon source (0) or supplemented with either sophorose or xylobiose in a final concentration of 2 mM. The results are given as the means derived from an identical experimental set up as in panel B.

indicating that AA<sub>-217/-218</sub> is essential for transcription. Mutation of the AGAA-box (pLWM1) exhibited a striking effect, since it not only increased the basal transcriptional level on glycerol but drastically increased the inducibility by xylan. The mutation within the CCAAT-box (pLWM3) also increased the basal level of transcription on glycerol but had a less-pronounced effect on the inducibility by xylan. Mutation 5 (pLWM5), which in vitro impairs the binding of Ace2 to the *xyn2* promoter, resulted in a moderate (20 to 40%) decrease of *xyn2* gene transcription under both noninducing and inducing conditions.

**Sophorose- and xylobiose-mediated induction involve the same nucleotide motifs.** We have previously shown that the expression of *xyn2* can be induced by both xylan and cellulose and their respective soluble degradation products, xylobiose and sophorose (44). Since transcription of the cellulase genes (*cbh1* and *cbh2*), which are also induced by cellulose and sophorose, is not triggered by xylan or xylobiose, we investigated whether

induction of *xyn2* expression by cellulose and xylan works via the same promoter motifs. To this end, we studied the inducibility of *xyn2* expression by xylobiose and sophorose in the various promoter mutants described above. As shown in Fig. 4C, both sophorose and xylobiose resulted in roughly similar levels of *xyn2* induction, which was 2.5-fold over the control (glycerol). A mutation of the AGAA-box (LWM1) led to a dramatic and comparable increase in *xyn2* expression with both xylobiose and sophorose. Similar to the behavior on xylan, a mutation in the CCAAT-box (LWM3) had only a small stimulatory effect on the inducibility of *xyn2* by xylobiose or sophorose. Also, in accordance with the behavior on xylan, a mutation in the Ace2-binding site (LWM5) reduced *xyn2* gene expression by about half. These data show that induction of *xyn2* by sophorose and xylobiose strictly follows the pattern observed on xylan, and the nucleotide motifs are therefore responsible for the induction by both xylan and cellulose.

## DISCUSSION

By a combination of in vitro and in vivo techniques, two adjacent *cis*-acting motifs on the noncoding strand of the *xyn2* promoter (5'-AGAA-3' and 5'-GGGTAAATTGG-3', respectively) were identified as responsible for the regulation of *xyn2* gene expression. Although the latter motif in analogy to the CAE in the *cbh2* promoter (43), named XAE, is essential for both the basal and induced expression of *xyn2*, the former

is only bound under noninducing conditions and is therefore a repressor of *xyn2* gene expression.

Results from *in vitro* protein-binding studies, *in vivo* footprinting, and chromatin analysis consistently showed that XAE within the *xyn2* promoter is constitutively bound by its *trans*-acting factors. In addition, results from EMSA analysis with heterologously expressed and purified DNA-binding proteins suggest that these *trans*-acting factors are the Hap2/3/5 protein complex (42) and Ace2 (1). Although a role for the Hap2/3/5 complex has been assumed from EMSA analysis with cell extracts (44), the nature of the other binding partner was unknown. The binding site [(GG)GTAATA] does not exactly match the consensus of any of the known DNA-binding proteins but has high similarity to those proposed for the xylanase regulator XlnR (36) and the cellulase regulator Ace2 (1).

Here we now provide evidence that Ace2 but not XlnR binds to this sequence and that the effect of two mutations in the Ace2-binding motif within XAE (a G-to-T mutation in the second guanine and a AA-to-TT mutation) on binding *in vitro* are perfectly reflected by the impact of these mutations on *xyn2* gene expression *in vivo*. The importance of individual nucleotides within XAE for binding of Ace2 noted here thereby strongly contrasts with the findings of Aro et al. (1), who reported that a simultaneous mutation in the TAA triplet only partially reduced binding, whereas mutating all three nucleotides of GGC eliminated DNA binding. Thus, the specific nucleotide exchanges performed in the present study produced essentially the opposite effect. The fact that the AA-to-TT mutation completely abolished both basal and induced *xyn2* expression indicates that Ace2 acts as a general rather than as a specific transactivator in *H. jecorina*. The use of cell extracts of *H. jecorina* in EMSA analysis revealed that impairment of Ace2 binding also affected binding of the Hap2/3/5 protein complex, whereas impairment of binding of the latter had no effect on Ace2 binding (E. Würleitner and R. L. Mach, unpublished data).

The Hap2/3/5-binding CCAAT motif in XAE is a common *cis*-acting element found in the promoter and enhancer regions of a large number of eukaryotic genes (20), including filamentous fungi (3), and has been shown to be involved in maintaining a transcriptionally active chromatin structure (24). We have recently shown that the CCAAT-box within the CAE of *H. jecorina* *cbh2* is involved in nucleosome assembly (S. Zeilinger, M. Leonhartsberger, M. Pail, R. L. Mach, and C. P. Kubicek, unpublished data) and that a mutation of CCAAT to CCCTT within CAE results in a 30% decrease in gene transcription (43). However, in the *xyn2* promoter, a similar mutation in the CCAAT-box of XAE resulted in a small but reproducible increase both in basal and in induced *xyn2* gene transcription, indicating that the Hap2/3/5 complex partially mediates *xyn2* repression. Negative regulation by the human Hap homologue NF-Y has been observed in some cases (e.g., topoisomerase II $\alpha$ , the type II transforming growth factor  $\beta$  receptor, and the varicella-zoster virus ORF62 [12, 13, 21]) and, more recently, also for the *lysF* promoter of the fungus *A. nidulans* (37). In hematopoietic cells, repression of the protein-tyrosine phosphatase gene by NF-Y was shown to be dependent on its interaction with the histones (40). A similar mechanism does not apply for the Hap2/3/5-mediated repression of *xyn2*, since chromatin analysis exhibited that the *xyn2* promoter is permanently accessible. An alternative mechanism has been found in the *SHP-1* gene promoter of MCF7 cells, where activating and

repressing effects of the CCAAT-box are due to position-dependent competition with binding of the transactivator COUP-TFII (16). However, we also consider this explanation less likely since the binding sites for Hap2/3/5 and Ace2, albeit partially overlapping, are located on opposite strands of the DNA. Rather, we consider it more likely that the Hap2/3/5 complex is involved in positioning the 5' nucleosomes, which in its absence therefore would move downstream and partially interfere with the binding of the AGAA-binding repressor protein.

The major difference in nucleotide protection between induced and noninduced conditions was the binding of an as-yet-unidentified protein to the AGAA-box. To the best of our knowledge, no such consensus sequence has so far been reported for other fungal promoters, although these four nucleotides form the core of the glucocorticoid receptor element (5'-AGAACA-3' [26]), which occurs in two inverted copies, separated by 3 bp (29). Interestingly, the AGAA-box of the *xyn2* promoter was also accompanied by an inverted copy (5'-AGAAcaacTTCT-3'), leading us to assume that the respective factor may actually bind to both boxes. However, mutation of the 3' TTCT-box had no effect on the binding to AGAA on the noncoding strand (Würleitner and Mach, unpublished), and this hypothesis was thus refuted.

The fact that the mutation in the AGAA element resulted in a >3-fold increase in *xyn2* gene transcription under both induced and noninduced conditions renders the *xyn2* promoter a potentially useful alternative for protein expression in *H. jecorina*. Currently, the *cbh1* promoter is almost exclusively used for these purposes, since the corresponding protein Cel7A (CBH I) accounts for >60% of the total protein secreted by the fungus (14). Average secreted amounts of Xyn2 are approximately one-fifth of the amount of Cel7A (32), implying that a *xyn2* promoter with a mutated AGAA motif may lead to protein yields almost as high as from the *cbh1* promoter.

We have previously reported that *xyn2* is subject to induction both by cellulose and by xylan and some of its corresponding degradation products (i.e., sophorose and xylobiose, respectively [44]). The results from the present study show that induction by both systems acts via the same promoter motif, which is also constitutively bound by its *trans*-acting factors, thus implying that specificity of induction does not involve a specific DNA-binding protein and rather must be due to signaling mechanisms. The relief from repression by the AGAA-binding protein alone is not sufficient to explain induction because both the induced and the basal transcription of *xyn2* are increased in the AGAA-to-GAGG mutant. Furthermore, an EMSA with cell extracts showed that the protein complex binding XAE exhibits the same migration property under induced and noninduced conditions and thus most likely consists of the same proteins. This argues against the possibility that a further protein would bind to the Hap2/3/5-Ace2-XAE complex under inducing conditions and thereby stimulate transcription. Rather, our working hypothesis is that the induction by xylan and/or cellulose may be mediated by covalent modification of Ace2, e.g., by phosphorylation. The identical migration of the XAE-protein complex from induced and noninduced conditions would not contradict this assumption, since the excess of oligonucleotide used likely masks the introduction of a phosphate group into Ace2 and thus will not give rise to a difference in the electrophoretic mobility. Our previous findings that *xyn2* gene transcription can be inhibited by

calmodulin antagonists, which led us to propose the involvement of a  $\text{Ca}^{2+}$ -calmodulin-dependent protein kinase in its induction (19), and the abundance of protein kinase targets in Ace2, including those for calmodulin-dependent protein kinase II (predicted by "PhosphoBase" [15]; <http://www.cbs.dtu.dk/databases/PhosphoBase/predict/predict.html>) would be in accordance with this speculation.

#### ACKNOWLEDGMENTS

This study was supported by a grant from Fonds zur Förderung Wissenschaftlicher Forschung to R.L.M. (P-12918 MOB).

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