Evidence for Temporal Regulation of the Two *Pseudomonas cellulosa* Xylanases Belonging to Glycoside Hydrolase Family 11

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*Pseudomonas cellulosa* is a highly efficient xylan-degrading bacterium. Genes encoding five xylanases, and several accessory enzymes, which remove the various side chains that decorate the xylan backbone, have been isolated from the *pseudomonad* and characterized. The xylanase genes consist of *xyn10A*, *xyn10B*, *xyn10C*, *xyn10D*, and *xyn11A*, which encode Xyn10A, Xyn10B, Xyn10C, Xyn10D, and Xyn11A, respectively. In this study a sixth xylanase gene, *xyn11B*, was isolated which encodes a 357-residue modular enzyme, designated Xyn11B, comprising a glycoside hydrolase family 11 catalytic domain appended to a C-terminal X-14 module, a homologue of which binds to xylan. Localization studies showed that the two xylanases with glycoside hydrolase family (GH) 11 catalytic modules, Xyn11A and Xyn11B, are secreted into the culture medium, whereas Xyn10C is membrane bound. *xyn10C*, *xyn10D*, *xyn11A*, and *xyn11B* were all abundantly expressed when the bacterium was cultured on xylan or β-glucan but not on medium containing mannans, whereas glucose repressed transcription of these genes. Although all of the xylanase genes were induced by the same polysaccharides, temporal regulation of *xyn11A* and *xyn11B* was apparent on xylan-containing media. Transcription of *xyn11A* occurred earlier than transcription of *xyn11B*, which is consistent with the predicted mode of action of the encoded enzymes. Xyn11A, but not Xyn11B, exhibits xylan esterase activity, and the removal of acetate side chains is required for xylanases to hydrolyze the xylan backbone. A transposon mutant of *P. cellulosa* in which *xyn11A* and *xyn11B* were inactive displayed greatly reduced extracellular but normal cell-associated xylanase activity, and its growth rate on medium containing xylan was indistinguishable from wild-type *P. cellulosa*. Based on the data presented here, we propose a model for xylan degradation by *P. cellulosa* in which the GH11 enzymes convert decorated xylans into substituted xylooligosaccharides, which are then hydrolyzed to their constituent sugars by the combined action of cell-associated GH10 xylanases and side chain-cleaving enzymes.

The recycling of photosynthetically fixed carbon through the action of microbial plant cell wall hydrolases is a key biological process. Xylan is one of the most abundant plant structural polysaccharides. This heterogencous polymer consists of a backbone of 1,4-linked xylopyranose residues which are decorated with acetyl, arabinofuranosyl, and 4-methyl-b-D-glucuronosyl side chains (6). The backbone xylose polymer is hydrolyzed removed by the action of arabinofuranosidases, mannan side hydrolase families (GH) 10 and 11 (http://afmb.cnrs-mrs.fr/∼cazy/CAZY/index.html [18]). GH 10 xylanases expressed by fungi and *Streptomyces lividans* display more relaxed specificity than GH 11 enzymes from the same microorganisms (7, 8), although it is unclear whether this general view of hemicellulose specificity can be extended to gram-negative prokaryotes. Xylanases, in common with other plant cell wall hydrolases, are encoded by large multigene families (13, 16, 22, 25), although the evolutionary pressures leading to this complex array of genes are not readily apparent.

The majority of plant cell wall hydrolases are not constitutively expressed but are induced by plant structural polysaccharides (9, 20, 30). Several of these enzymes are induced only by their target substrate, while others are synthesized in response to a range of plant structural polysaccharides (9, 26, 33). The general model for induction of microbial plant cell wall-degradative enzymes involves the low-level constitutive expression of a “sensor” enzyme, which generates small oligosaccharides or disaccharides from plant cell wall material that enter the microorganism and elicit the high-level synthesis of plant cell wall hydrolases (26, 30, 33). This model has been demonstrated for *Trichoderma reesei* xylanases; *xyn1* is only induced by xylan or its degradative products, whereas *xyn2* displays low-level constitutive expression and its synthesis is activated by both xylan and cellulose (33). Similarly, it has been proposed that the xylanase encoded by *Streptomyces cyanus* was the constitutively expressed sensor protein that activated the synthesis of the xylan-degrading system in this bacterium (5). Currently, regulatory proteins that control xylanase gene expression have only been identified in filamentous fungi. In *Aspergillus niger* XlnR was shown to be a general transcriptional activator of the xylanolytic system, binding to a consensus heptanucleotide sequence that is also present in the promoters of *Penicillium chrysogenum* xylanase genes (32). Recently, a novel *Trichoderma* transcriptional activator, ACEII, was identified that induced expression of the majority of cellulases and the xylanase encoded by *xyn2* (2).

Although there have been numerous studies on the regulation of xylanase gene expression in fungi, the regulatory mechanisms that induce the synthesis of the corresponding bacterial...
enzymes and the cellular location of these glycoside hydrolases are poorly understood. In this study we investigated the expression patterns and the cellular location of the xylanases produced by the bacterium *Pseudomonas cellulosa*. *P. cellulosa* is an efficient xylan-degrading bacterium (15), which was previously shown to contain genes encoding five xylanases in addition to several side chain-elevating enzymes (4, 11). The bacterium displays xylanase activity when cultured on xylan or cellulose (13, 14, 16, 17, 22, 25); however, it is unclear whether specific subsets of xylanase genes are synthesized in response to different polysaccharides. The cellular location of these enzymes in *P. cellulosa*, apart from Xyn10A (which is extracellular, i.e., secreted into the culture medium when the bacterium is grown on Luria broth [LB] supplemented with xylan [17]), has also not been elucidated, and it is unknown whether the bacterium synthesizes a sensor xylanase. In this report we identify a new *P. cellulosa* GH1 xylanase, Xyn11B, and demonstrate that all of the xylanases from this pseudomonad are induced by the same polysaccharides, although there was evidence for differential temporal expression of the genes encoding the GH11 enzymes. Although one of the highly expressed GH10 enzymes, Xyn10C, is located on the cell membrane, the two GH11 enzymes appear to be the major xylanases in the extracellular milieu. The pattern of xylanase expression, and their cellular location, is discussed in relation to the capacity of the bacterium to use xylan as a carbon and energy source.

### MATERIALS AND METHODS

**Bacterial strains, plasmids, and culture conditions.** The *Escherichia coli* strains used in this study were BL21(DE3)pLysS (Novagen) and XL1-Blue ( Stratagene). The bacteriophage employed comprised a genomic library of *P. cellulosa* (NCIMB 10462) constructed previously in AZAPII (Stratagene [25]). The plasmid vectors used were pCR-Blunt (Invitrogen), pETTib (Novagen), and pRG960bd (31). All *E. coli* strains containing recombinant plasmids were cultured in LB supplemented with 50 μg of ampicillin or kanamycin/mL, as needed, at 37°C unless otherwise stated. *E. coli* cells used to propagate bacteriophages were grown on LB supplemented with 10 mM MgSO₄ and 0.2% (wt/vol) maltose at 30°C before being plated on media supplemented with streptomycin and spectinomycin (both at 50 μg/mL). To introduce the transposon Tn10 into the *P. cellulosa* genome, *E. coli* ST1-1apir harboring pLOFkm (contains a mini-Tn10 element [19]) and the pseudomonad were cultured overnight in LB supplemented with appropriate antibiotics. The cells were pelleted, washed, and then resuspended in phosphate-buffered saline. The cell suspensions were then mixed in a *P. cellulosa*/*E. coli* ratio of 5:1, and a 6-ml portion was filter fated as described previously (3), except that the filters were incubated overnight on Luria agar containing 1 mM IPTG to induce transposase expression, and transconjugants were selected on medium containing 50 μg of rifampin and kanamycin.

**RNA quantification.** RNA was isolated from cultures of *P. cellulosa* grown at 30°C at different stages of the growth phase as described previously (3). Aliquots of RNA (5 μg) were electrophoresed on 1.2% (wt/vol) agarose-formaldehyde denaturing gels (27) and transferred to Hybond N membranes (Amersham-Pharmacia) by using 300 mM sodium citrate buffer (pH 7.2) containing 3 M NaCl, and the nucleic acid was immobilized by UV cross-linking. The filters were then probed with radiolabeled DNA encoding the various xylanase genes. Probe bound to the transcripts was determined by electronic autoradiography (phosphorimagery) by using a Packard Instant Imager. To confirm that the signals were in the linear range of detection, samples which gave the highest and lowest intensity signals were subjected to Northern hybridization with 2.5-, 5-, 10-, and 20-μg RNA samples, and the intensity of the signal of the bound probe was determined. In all cases there was a linear correlation between signal intensity and the amount of RNA electrophoresed. The probes used to detect transcripts derived from *xyn10A* (16), *xyn10B* (22), *xyn10C* (13), *xyn10D* (25), *xyn11A*, and *xyn11B* (Fig. 1, pKE5) encoded the catalytic domains of the respective enzymes, except the *xyn10D* probe, which encoded the full-length xylanase. To distinguish between *xyn11A* and *xyn11B*, probes encoding the xylan-binding modules of *Xyn11A* (pKE3) and *Xyn11B* (pKE4) were used.

To measure transcripts by real-time PCR, 1 μg of total *P. cellulosa* RNA, prepared as described above, was used as a template to synthesize cDNA by using random primers and the Superscript preamplification cDNA synthesis kit (Gibco-BRL). The cDNA was then subjected to PCR with 200 μM concentrations of each transcript-specific primer, 3 mM MgCl₂, and the Light Cycler-DNA SYBR Green 1 Master Mix (Roche Diagnostics). The PCRs were carried out in a Roche Light Cycler instrument with the following temperature parameters: 95°C for 20 min, followed by 50 cycles of 95°C for 15 s, 60°C for 5 s, and 72°C for 10 s. The fluorometric intensity of SYBR Green 1, a specific dye for double-stranded DNA, was measured at the end of each elongation. Fluorescent signals caused by primer dimers and nonspecific background were evaluated by melting-curve analysis and agarose gel electrophoresis, as recommended by the manufacturer. The sizes of the amplified DNA were as follows: *xyn10A*, 380 bp; *xyn10B*, 320 bp; *xyn11A*, 320 bp; *xyn11B*, 238 bp; *xyn10D*, 385 bp; and *xyn10C*, 900 bp.

**Construction of recombinant plasmids.** To clone the 447-bp (463 to 16) and 445-bp (461 to 17) regions immediately upstream of *P. cellulosa*, *xyn11A* and *xyn11B*, respectively, were amplified by PCR. The PCRs were performed with 2 U of the thermostable DNA polymerase Vent (New England Biolabs). The reactions contained 2 mM MgSO₄, 1× Thermopol reaction buffer (New England Biolabs), 0.5 μM concentrations of primers, and 60 ng of target DNA in a final volume of 100 μl. The reactions were subjected to 25 cycles at the following temperatures: 95°C for 1 min, 55°C for 1 min, and 72°C for 2 min. The amplified DNAs were inserted into pCR-Blunt, the Blunt vectors were sequenced, and the nucleic acids were then excised by restricting with BamHI and PspAI (an isoschizomer of SmaI) and cloned into similarly digested pRG960bd to generate pRG960bd-XKE7, which contain the promoter regions of *xyn11A* and *xyn11B*, respectively.

To clone the 5’-region of *xyn11B*, primers were designed which hybridized to the region of *xyn11A* that encoded the family 4 carbohydrate esterase domain of *Xyn11A* and the region of *xyn11B* encoding the C-terminal sequence of the enzyme.
ylink-binding carbohydrate-binding module (CBM) of Xyn11B. The primers were used to amplify the target sequence from *P. cellulosa* genomic DNA, and the 1,728-bp product was cloned into pCR-Blunt to generate pKE2. To clone DNA encoding the xylan-binding CBM, the appropriate region of *xyn11B* was amplified by PCR. The product was cloned into pCR-Blunt to generate pKE4, and the resultant plasmid was used in Northern and Southern analyses of *P. cellulosa* RNA and DNA, respectively. To clone DNA encoding the catalytic domain of Xyn11A, the appropriate region of *xyn11A* was amplified by PCR and the product was restricted with *NdeI* and *XhoI* and cloned into similarly digested pET11b to generate pKE5.

To clone the GAPDH (glyceraldehyde-3-phosphate dehydrogenase) gene (*gpd*) from *P. cellulosa*, the primary structures of the enzyme from 50 prokaryotic and eukaryotic microorganisms were compared. The data revealed a highly conserved sequence at the N and C termini of these proteins, from which primers were designed based on the nucleotide sequence of *gpd* from *Pseudomonas aeruginosa* (M74256). These primers were used in a PCR which generated a 934-bp product that was cloned into the pCR-Blunt and sequenced. Analysis of the sequence revealed that the DNA encoded GAPDH. The amplified DNA was used as a probe in Northern hybridization of *P. cellulosa* RNA as described above.

All of the primers used in the various PCRs described above are available on request.

**Assays.** Protein was measured by the dye-binding method of Sedmak and Grossberg (28) with bovine serum albumin as the standard. Xylanase activity was determined by using a standard reducing sugar assay with dinitrosalicylic acid (24). The reporter enzyme β-glucuronidase was assayed as follows. A 500-μl reaction mixture of 50 mM sodium phosphate buffer (pH 7.0), 1 mM 4-methylumbelliferyl-β-D-glucuronide, 1 mg of bovine serum albumin/ml, and an appropriate dilution of the bacterial cell extract was incubated at 37°C for 10 min and was terminated by the addition of 2.5 ml of 50 mM glycine-NaOH buffer (pH 10.4). The release of the fluorescent reaction product 4-methylumbelliferyl was determined with a RF-1501 spectrophotofluorometer (Shimadzu) by using excitation and emission wavelengths of 365 and 460 nm, respectively.

**Fractionation of *P. cellulosa* and Western analysis.** Stationary cells of *P. cellulosa* were harvested by centrifugation at 6,000 × g for 10 min from a 400-ml culture comprising minimal medium in which wheat arabinoxylan was the carbon source. The cell pellet was then resuspended in 10 ml of 50 mM sodium phosphate-12 mM citric acid buffer (pH 6.5) and sonicated. The cell membranes were then pelleted by centrifugation at 100,000 × g at 4°C for 1 h, and the supernatant, consisting of the cytoplasm and the periplasm (referred to here as the cell extract) was retained for further use. The purity of the different fractions was assessed by measuring the activity of the marker enzymes malate dehydrogenase (cell extract) and NADH oxidoreductase (membrane fraction) as described previously (13). Western analysis of the various fractions was carried out as described previously (13) with antisera raised against the catalytic domains of *xyn11A*, *Xyn10C*, and *Xyn10D*.

**Nucleotide sequence accession number.** The nucleotide sequence for *xyn11B* has been deposited in the GenBank database under accession no. AY065640.

**RESULTS**

**Identification of a novel *P. cellulosa* xylanase gene *xyn11B*.** In previous studies five genes encoding xylanases were isolated from *P. cellulosa*. These genes were designated *xyn10A* (formerly *xynA*), *xyn10B* (formerly *xynB*), *xyn11A* (formerly *xynE*), *xyn10C* (formerly *xynF*), and *xyn10D* (formerly *xynG*), respectively (13, 16, 22, 25). To investigate the regulation of xylanase expression in *P. cellulosa*, it is important that all of the xylanase genes are identified in this bacterium. Exhaustive screening of 10⁶ clones of a λZAPII library of *P. cellulosa* failed to identify any xylanase-expressing clones not previously characterized. Southern hybridization with *xyn11A* as a probe, however, revealed the presence of a sixth xylanase gene, defined as *xyn11B*. The observation that the probe bound to single *BamHI*, *BglII*, *DraI*, and *PstI* restriction fragments may point to linkage between *xyn11A* and *xyn11B*, although this interpretation of the data must be viewed with some caution (Fig. 2). Screening of the genomic library with *xyn11A* led to the isolation of a clone (pKE1) that contained an incomplete open reading frame (ORF), designated *xyn11B*, which exhibited sequence similarity to the 3′ region of *xyn11A*. To clone the missing sequence of the ORF, primers that bound to the 3′ region of *xyn11A* and *xyn11B* (primers were directed against the 3′ region of *xyn11B*, which was not conserved in *xyn11A*), respectively, were used in a PCR of *P. cellulosa* genomic DNA. The resultant DNA product was cloned into pCR-Blunt to generate pKE2. The
sequence of the pseudomonad insert in pKE2 showed that the plasmid contained the missing 5’ region of xyn11B. The physical map of the xyn11A/xyn11B locus is shown in Fig. 1.

Translation of the xyn11B ORF (present in GenBank under the accession no. AY065640) showed that the gene encoded a protein, designated Xyn11B, of M_r 37,889. Comparison of the primary structure of Xyn11B with protein databases indicated that the enzyme had a modular structure. The N-terminal 27 amino acids conform to a typical signal peptide; the 9 N-terminal hydrophilic residues contain several basic amino acids and are followed by a stretch of 18 small hydrophobic residues that are capable of forming an α-helix, which is followed by a potential signal peptidase I cleavage site. The putative signal peptide is followed by 210 amino acids that exhibit 94% sequence identity with the catalytic module of Xyn11A (encoded by xyn11A [25]), demonstrating that this region comprises a catalytic module belonging to GH11. Inspection of the catalytic module of Xyn11A showed that Glu116 and Glu213 were the key catalytic residues of the xylanase since they correspond to key catalytic residues of the GH11 enzyme, which are the catalytic nucleophile and catalytic acids, respectively (21). The GH11 catalytic module is separated by a 20-residue linker sequence rich in serines and glycines from a 110-amino-acid module that displays 57% sequence identity with residues 254 to 362 of Xyn11A. Recent studies have shown that this region constitutes a xylan-binding CBM (H. Xie et al., unpublished data). Thus, Xyn11B is highly homologous to the N-terminal region of Xyn11A, which comprises the xylanase GH11 catalytic module and the xylan-binding CBM, but it lacks the family 4 carbohydrate esterase and the family 10 CBM that are present at the C terminus of Xyn11A (25). The xyn11B ORF is preceded by a typical ribosome-binding sequence, GGAGA, 9 bp upstream of the translational start codon. The sequence immediately downstream of the ORF of xyn11B is GC-rich and contains a region of dyad symmetry that was followed and preceded by a series of thymine and adenine nucleotides, respectively. This region is likely to act as a typical rho-independent transcriptional terminator sequence when either the sense or the antisense strand is transcribed. Upstream of the ORF is a 34-nucleotide sequence that exhibits almost perfect dyad symmetry, although it is not preceded by a succession of thymines, and thus the functional significance of this region is currently unknown.

To determine whether the putative catalytic module of Xyn11B is functional, DNA encoding this region (pKE2; Fig. 1) was transformed into E. coli. The recombinant bacterium, when plated onto LB containing IPTG and either either Blue-Xylan (Sigma Chemical Co.) or Azo-wheat arabinoxylan (Megazyme International Ltd.), displayed xylanase activity as evidenced by the appearance of clear haloes surrounding the bacterial colonies in a blue background.

**Localization of xylanase activity.** To determine the location of the xylanase activity expressed by *P. cellulosa*, the bacterium was cultured on xylan and the cells, in mid-log phase, were fractionated, and the xylanase activity in each fraction was assessed. The data, presented in Table 1, showed that ca. 65% of xylanase activity was present in the culture supernatant and that the remainder was cell associated. It would appear that the majority of cell-associated xylanase activity was on the outer membrane since xylan degradation by whole cells was similar in medium and that was cell-associated was determined. Values are given as means ± the SD where indicated.

**Growth medium**

<table>
<thead>
<tr>
<th>Culture supernatant</th>
<th>Cell associated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oat spelt xylan</td>
<td>0.73 ± 0.07</td>
</tr>
<tr>
<td>Glucuronoxylan</td>
<td>0.65 ± 0.07</td>
</tr>
<tr>
<td>Rye arabinoxylan</td>
<td>0.81 ± 0.10</td>
</tr>
<tr>
<td>Wheat arabinoxylan</td>
<td>0.91 ± 0.11</td>
</tr>
<tr>
<td>β-Glucan</td>
<td>0.21 ± 0.02</td>
</tr>
<tr>
<td>Carbo galactomannan</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Pectin</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>LB</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>LB and oat spelt xylan</td>
<td>0.21 ± 0.04</td>
</tr>
<tr>
<td>Oat spelt xylan/glucose</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

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*P. cellulosa* was grown to mid-log phase, and xylanase activity in the culture medium and that was cell-associated was determined. Values are given as means ± the SD where indicated.

*The assay can detect 10^-2 U of xylanase activity/ml.*
to that by sonicated cell extracts, in which the cell membranes had been disrupted releasing the periplasmic and cytosolic proteins (data not shown). Antisera raised against the xylanases were then used to probe the cellular location of the different enzymes. The data, presented in Fig. 3, showed that Xyn10C was predominantly associated with the cell membrane, although the protein could also be detected in the cell extract, whereas the GH11 xylanases appeared to be present in the culture supernatant; Xyn10D gave no obvious signal on Western analysis (data not shown). The size of Xyn10C, determined by Western analysis, was very similar to its deduced molecular size, indicating a lack of posttranslational modification.

Expression of xylanase activity in different media. *P. cellulosa* was grown on a variety of different media, and the expression of xylanase activity was monitored with time. The data, presented in Table 1, showed that xylan induced the expression of xylanase activity in both the culture supernatant and the cell fraction. Although cellulosic substrates also induced xylanase activity, the level of expression was lower than cells cultured on xylan. No xylanase activity was detected when the bacterium was cultured on LB, mannan, or pectic polysaccharides. When xylan media were supplemented with LB, the level of xylanase expression was greatly reduced, although the inclusion of mannan or pectin in xylan media did not repress the synthesis of these enzymes. No xylanase activity was detected in xylan medium supplemented with glucose until the stationary phase was reached. Assaying the culture supernatant for glucose showed that xylanase activity only appeared when all of the hexasaccharide had been metabolized (data not shown). These data show xylanase activity is not constitutively expressed but is induced by xylan and, to a lesser extent, by β-glucan; however, β-mannans or pectins do not influence the expression of these enzymes.

Regulation of xylanase gene transcripts. The expression of the individual xylanase genes was determined by Northern hybridization. The results obtained from *P. cellulosa* cultured on wheat arabinoxylan are presented in Fig. 4. These data are qualitatively similar to the transcript profile of the xylanase genes when *P. cellulosa* was grown on oat spelt xylan, rye arabinoxylan, and glucuronoxylan (data not shown). The sizes of mRNA and of the respective ORFs (in parentheses) for the various genes were as follows: xyn10A, 1.8 kb (1.83 kb); xyn10C, 1.8 (1.82 kb); xyn10D, 1.2 kb (1.13 kb); xyn11A, 2.0 kb (1.98 kb); and xyn11B, 1.3 kb (1.1 kb). Thus, the lengths of the transcripts were consistent with the sizes of the ORFs encoded by the respective genes. It should be noted that when the region of xyn11A encoding the catalytic domain was used in Northern hybridizations, transcripts of 2.0 and 1.1 kb were detected, a finding consistent with the size of xyn11A and xyn11B mRNA.
Probes that consisted of the regions of xyn11A and xyn11B that encoded the xylan-binding CBMs of the respective enzymes hybridized to 2.0- and 1.1-kb mRNA species, respectively, showing that these probes were specific for transcripts derived from their target genes (Fig. 4). The data revealed that the highest levels of transcripts were from the xyn10D and xyn11B genes, although xyn11A and xyn10C were also efficiently transcribed (Fig. 5). Low levels of mRNA derived from xyn10A were observed, and no expression of xyn10B could be detected. In general, the xylanase genes were transcribed in mid-log phase when cultured on xylan, although only xyn11B expression persisted into stationary phase. The one very clear exception to this pattern was xyn11A mRNA, which appeared in early log phase, and its level had greatly decreased when the other xylanase transcripts were synthesized (Fig. 5). On β-glucan, xylanase mRNA molecules appeared in late log phase, shortly after the cellulase transcripts had reached their respective highest levels (Fig. 6 and 7), and the accumulations of these mRNAs were lower than on xylan. No xylanase transcripts were detected by Northern analysis on LB or on media containing glucose and xylan or when either mannan or pectin were the sole carbon sources (data not shown). To determine whether the lack of transcripts on mannan, LB, or pectin was due to the relatively low sensitivity of Northern hybridization, the RNA preparations were subjected to real-time PCR with primers that were specific for the six xylanase genes. The results were very similar to the Northern hybridization data. Transcripts encoding five of the xylanases were present when the bacterium was cultured on xylan-containing or β-glucan-containing media, but these mRNA species were not detected when the pseudomonad was grown on media containing mannan, pectin, or LB (data not shown). This methodology did not detect xyn10B on any of the media evaluated. To validate that the lack of xylanase transcripts on various media was not due to mRNA degradation, these RNA preparations were probed...
for the presence of transcripts derived from the housekeeping gene *gpd*. The data (examples of which are displayed in Fig. 6 and 7) showed that mRNA encoded by *gpd* could easily be detected, and in all RNA samples the two major rRNA species were clearly visible.

To further probe the regulation of xylanase gene expression, the 450-bp untranslated 5′ regions of *xyn11A* and *xyn11B* were inserted immediately upstream of the β-glucuronidase gene (*uidA*) present in the promotor probe vector pRG960sd, which replicates in a range of gram-negative bacteria (31). The recombinant plasmids, pKE6 and pKE7, which contain the putative *xyn11A* and *xyn11B* promoters, respectively, were introduced into *P. cellulosa* by using either conjugation, with a helper plasmid, or electroporation, demonstrating for the first time that it is possible to transform the gram-negative bacterium. The copy number of the plasmid in the pseudomonad was ca. 1. The level of β-glucuronidase in wild-type *P. cellulosa* or in the bacterium containing pRG960sd was negligible; however, the pseudomonad containing either pKE6 or pKE7 expressed β-glucuronidase activity when cultured in the presence of xylan (Table 2). The addition of glucose to the xylan-containing medium prevented expression of β-glucuronidase in mid-log-phase cultures (Table 2); however, when the hexose sugar had been fully metabolized in stationary phase, substantial reduction in extracellular xylanase activity, the hydro-
lytic capacity of the cell-bound form of the enzyme was similar to that of the wild-type bacterium. Surprisingly, the growth rate of the transconjugants on media containing xylan as the sole carbon source was similar to that of the wild-type bacterium (data not shown). Southern hybridization showed that in these transconjugants the minitransposon had integrated into the \textit{xyn11A}/\textit{xyn11B} locus, mediating a deletion event encompassing both genes, which was confirmed by the absence of \textit{xyn11A} and \textit{xyn11B} transcripts when the transconjugants were cultured on xylan (data not shown). Thus, the reduction in the extracellular xylanase activity was due to the absence of both Xyn11A and Xyn11B.

DISCUSSION

The main objective of this study was to examine the expression of the xylanase genes of \textit{P. cellulosa}. The data showed that, whereas \textit{xyn10C}, \textit{xyn10D}, \textit{xyn11A}, and \textit{xyn11B} were efficiently expressed, only low levels of \textit{xyn10A} transcripts were detected. The ratio of the different xylanase transcripts remained similar irrespective of the culture media, with none of the genes apparently expressed in the absence of xylan or cellulose. These data are in contrast to the xylanase system in \textit{T. reesei}. In this fungus the two major xylanase genes are expressed in response to different nutrients, with \textit{xyn2} displaying significant constitutive expression that is induced by both xylan and cellulose, whereas \textit{xyn1} is only transcribed when media contain xylan (33). In the general model for induction of xylanase expression a sensor enzyme is constitutively expressed (e.g., \textit{T. reesei} \textit{xyn2} [33]) which hydrolyzes xylan into oligosaccharides that enter the bacterium and activate the expression of all of the xylanase genes. It is possible that one or more of the repertoire of \textit{P. cellulosa} xylanases are expressed at extremely low levels in the absence of xylan (i.e., <10 copies per cell, which is the minimum number of transcripts the real-time PCR method can detect) and thus function as the sensor enzyme. Further studies are clearly required to determine the mechanism by which xylan induces expression of the \textit{P. cellulosa} xylanases.

Although the temporal expression of four of the xylanase genes was broadly similar, with maximum levels of transcripts appearing in mid-log phase, \textit{xyn11A}, interestingly, was transcribed in early exponential phase. The expression pattern of \textit{xyn11A} may reflect the higher proportion of extracellular xylanase activity in the culture media, compared to the cell-associated forms of these enzymes, during the early component of the growth phase. Another interesting feature of the expression pattern of \textit{Xyn11B} is that transcripts of the gene persist into the late exponential and early stationary phases, at which point no other xylanase mRNAs were detected. These data may indicate that \textit{xyn11B} mRNA exhibits an unusually high level of stability. The sequential synthesis of the two family 11 enzymes is consistent with their biochemical properties. Both xylanases are secreted into the extracellular culture medium, suggesting that they play a key role in plant cell wall degradation. Furthermore, they both contain xylan- and family 11 xylanase-binding CBMs, indicating that these enzymes target the plant cell wall as their primary substrate. Xyn11A also contains a family 4 xylan acetyl esterase and hydrolyzes acetylated xylan, whereas Xyn11B displays no activity against this decorated xylan. Since ca. 70% of xylose residues in plant cell wall xylans are decorated with acetate groups, the removal of these side chains is an essential prerequisite to xylan hydrolysis. Thus, we propose that the early expression of Xyn11A rapidly removes the acetate groups, making the backbone polysaccharide accessible to xylanase attack, and thus the second major extracellular xylanase, Xyn11B, is expressed at a later stage of the growth phase.

The importance of Xyn11A and Xyn11B in xylan degradation is illustrated by the observation that transposon mutants of \textit{P. cellulosa} with greatly reduced xylanase activity lacked both of these enzymes but expressed the three family 10 xylanases at levels similar to that of the wild-type bacterium. This could reflect the nature of the screen which selected for bacteria with small haloes on Blue-Xylan. Since the secreted enzymes make the largest contribution to halo formation, deletions in Xyn11A and Xyn11B are likely to be isolated from the screen. A rather surprising feature of the mutants is that their growth rates were similar to that of the wild-type bacterium on media in which xylan was the sole carbon source. This suggests that the amount of extracellular xylanase activity is not a limiting factor in the growth of \textit{P. cellulosa} on xylan. It would appear, therefore, that either the hydrolysis of xylooligosaccharides to xylose or the transport of these molecules is the rate-limiting step in the utilization of poorly substituted xylans. In contrast, deletion of any of the three xylanase genes of \textit{Streptomyces lividans} reduced the growth rate of the bacterium, suggesting that in this prokaryote the rate of xylan degradation is the rate-limiting factor (1). It is likely, however, that the loss of the esterase activity of Xyn11A in the \textit{P. cellulosa} mutant would limit its capacity to utilize “natural” highly acetylated xylans. Indeed, the importance of arabinose removal has already been shown to be a limiting factor in the utilization of arabinoxylans as a carbon source by the bacterium (3).

In contrast to the other xylanase genes, no expression of \textit{xyn10B} was detected. It is interesting that \textit{xyn10B} is 120 bp downstream of \textit{abf62A} (formerly \textit{xynC}), a gene encoding a xylan-specific arabinofuranosidase, which is also not expressed in \textit{P. cellulosa} cultured on liquid media containing xylan, arabinobioxyan, or glucuronoxylan (3). These data are somewhat puzzling since these genes encode functional enzymes when expressed in \textit{E. coli}, indicating that they are likely to play a role in hemicellulose degradation in \textit{P. cellulosa}. It is interesting that the enzymes encoded by these genes are the only xylan-degrading glycoside hydrolases to contain family 2a CBMs, which mediate tight and irreversible binding to crystalline cellulose (23). Thus, the primary function of these enzymes could be to attack xylans that are in intimate contact with cellulose, and thus it would be rational for these proteins to be expressed when \textit{P. cellulosa} is presented with complex plant cell walls containing xylan-cellulose composites. This view is consistent with a previous study which showed that an \textit{Aspergillus} xylanase was not expressed in monocultures of the fungus but was produced when the organism was inoculated into compost (10), a highly complex environment. Unfortunately \textit{P. cellulosa} grew very poorly on plant cell walls or insoluble cellulose-xylan composites, precluding the analysis of \textit{xyn10B} expression when the bacterium is presented with complex insoluble substrates.

The transposon mutant lacking \textit{xyn11A} and \textit{xyn11B} expressed similar levels of the other xylanases to wild-type \textit{P. cellulosa}. This finding is in sharp contrast to those of a previous
study on the *Streptomyces lividans* xylanase genes. The bacterium produced three major xylanases, and expression of two of the genes was intimately dependent on the expression of the other (1). The data presented here do not exclude such an interrelationship between the expression of the *P. cellulosa* xylanases, but such a phenomenon does not occur between the family 10 and 11 enzymes.

In *P. cellulosa* the two family 11 xylanases are extracellular, whereas Xyn10C is membrane bound, a finding consistent with its extensive linker sequence between the signal peptide and the catalytic module (25). It was not possible, however, to determine the location of Xyn10D; it is a highly unstable enzyme (data not shown) and was thus probably degraded during processing of the samples for Western analysis. The highly homologous enzyme Xyn10A (which exhibits 96% identity with *P. cellulosa* Xyn10D) from the related bacterium *Cellvibrio mixtus* subsp. *cellulosa* is periplasmic (13), and thus it is likely that Xyn10D is also cell associated. These data are in contrast with the generally held view that aerobic microorganisms secrete plant cell wall hydrolases into the extracellular medium, whereas anaerobic prokaryotes assemble the corresponding enzymes into large multienzyme complexes that are associated with the bacterial cell surface (29). The selective pressure that led to the different locations of the xylanases is an important issue. It is possible that the function of the membrane-bound xylanases may be related to the cell-associated location of the enzymes that remove the arabino and glucurono decorations. The action of the arabinoxylanases and α-glucuronosidases generates linear xylose polymers proximal to the bacterium. The cleavage of these polymers to xylose and small oligosaccharides by cell-associated xylanases would enable the products of hemicellulose degradation to be utilized primarily by *P. cellulosa* rather than by competing bacteria which inhabit these complex ecosystems. Support for this model is provided by the study of Shulami et al. (30) on the gluconic acid gene cluster in *Bacillus stearothermophilus*, which showed that the bacterium synthesized an extracellular xylanase, similar to Xyn10C, that generates short decorated xylooligosaccharides that enter the bacterium and are further degraded to monosaccharides by a battery of intracellular hemicellulases.

**Conclusions.** We have shown here that the synthesis of the different xylanases in *P. cellulosa* is induced by the same poly-saccharides, although there are differences in the temporal expression of the family 11 enzymes. The differences in the temporal expression of Xyn11A and Xyn11B are consistent with their mode of action, with the former enzyme capable of removing acetyl side chains, as well as attacking the xylan backbone, whereas the latter enzyme only cleaves β1,4-xylosyl linkages. The spatial localization of the different xylanases indicates that the various isoforms of these enzymes play quite different roles in the xylan degradation process. The extracellular enzymes mediate initial xylan breakdown, while the cell-associated hemicellulases hydrolyze linear xylooligosaccharides in which the side chains have been removed by membrane-bound glycoside hydrolases. Transposon mutagenesis studies support the view that Xyn11A and Xyn11B play an important role in xylan degradation, although the hydrolysis of the polysaccharide does not appear to be the rate-limiting step in the utilization of xylan as a growth substrate. Finally, this study has generated genetics tools, a random transposon mutagenesis system, and a promoter probe vector in which the reporter gene is regulated by xylan promoters that will be invaluable in the further analysis of the mechanisms regulating the synthesis of the xylan degradative enzymes of *P. cellulosa*.

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