Cloning and sequencing of the cellulose synthase catalytic subunit gene of *Acetobacter xylinum*

Inder M. Saxena, Fong Chyr Lin and R. Malcolm Brown, Jr*

Department of Botany, University of Texas at Austin, Austin, TX 78713–7640, USA (*author for correspondence)

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**Abstract**

The gene for the catalytic subunit of cellulose synthase from *Acetobacter xylinum* has been cloned by using an oligonucleotide probe designed from the N-terminal amino acid sequence of the catalytic subunit (an 83 kDa polypeptide) of the cellulose synthase purified from trypsin-treated membranes of *A. xylinum*. The gene was located on a 9.5 kb *Hind* III fragment of *A. xylinum* DNA that was cloned in the plasmid pUC18. DNA sequencing of approximately 3 kb of the *Hind* III fragment led to the identification of an open reading frame of 2169 base pairs coding for a polypeptide of 80 kDa. Fifteen amino acids in the N-terminal region (positions 6 to 20) of the amino acid sequence, deduced from the DNA sequence, match with the N-terminal amino acid sequence obtained for the 83 kDa polypeptide, confirming that the DNA sequence cloned codes for the catalytic subunit of cellulose synthase which transfers glucose from UDP-glucose to the growing glucan chain. Trypsin treatment of membranes during purification of the 83 kDa polypeptide cleaved the first 5 amino acids at the N-terminal end of this polypeptide as observed from the deduced amino acid sequence, and also from sequencing of the 83 kDa polypeptide purified from membranes that were not treated with trypsin. Sequence analysis suggests that the cellulose synthase catalytic subunit is an integral membrane protein with 6 transmembrane segments. There is no signal sequence and it is postulated that the protein is anchored in the membrane at the N-terminal end by a single hydrophobic helix. Two potential N-glycosylation sites are predicted from the sequence analysis, and this is in agreement with the earlier observations that the 83 kDa polypeptide is a glycoprotein [13]. The cloned gene is conserved among a number of *A. xylinum* strains, as determined by Southern hybridization.

**Introduction**

Cellulose, a β-1,4 linked polymer of glucose, is synthesized by all plants, algae, some fungi, and a few bacteria [3]. It is also synthesized in the animal kingdom by tunicates [25] and is reported to be present in humans under certain disease conditions, notably scleroderma [8]. Almost all

The nucleotide sequence data reported will appear in the EMBL, GenBank and DDBJ Nucleotide Sequence Databases under the accession number X54676.
cellulose obtained commercially is produced by plants, where it serves a structural role in the cell wall. Efforts to understand cellulose biosynthesis in plants have been limited by the inability to purify the cellulose synthase and demonstrate in vitro synthesis of cellulose from purified membrane proteins [3]. Among the other organisms that synthesize cellulose, the Gram-negative bacterium *Acetobacter xylinum* is the most studied for its capacity to synthesize cellulose. Cells of *A. xylinum* can utilize a variety of substrates for synthesizing cellulose and convert as much as 50% of supplied carbon to cellulose. The cellulose ribbons intertwine to form a pellicle that floats at the air-liquid interface of stationary cultures. The cellulose is produced in a pure form without being attached to other polysaccharides or proteins. Chemically, *Acetobacter* cellulose is indistinguishable from cellulose obtained from plants, but certain additional features such as strength, hydrophilicity, and the capacity to mold the native pellicle into any desired shape makes it superior for certain applications [26].

Biochemical studies in *A. xylinum* established that UDP-glucose is the substrate for cellulose biosynthesis and that no intermediate is involved in the reaction catalyzed by the enzyme cellulose synthase (UDP-glucose:1,4-β-D-glucosyltransferase; E.C.2.4.1.12). In vitro synthesis of cellulose from cell-free preparations of *A. xylinum* was first demonstrated by Glaser in 1958 [7]. More efficient synthesis of cellulose in vitro was obtained when an activator of cellulose synthase was discovered [21]. The demonstration of cellulose production in vitro from membrane preparations of *A. xylinum* provided a reliable assay for cellulose synthase activity [14]. Cellulose synthase was first purified by product entrapment, and the purified preparation showed two major polypeptides of 93 and 83 kDa on lithium dodecyl sulfate (LDS)-polyacrylamide gel following electrophoresis [13]. The 83 kDa polypeptide has been shown by photoaffinity labelling to be the catalytic subunit of cellulose synthase [15]. Nothing substantial is known about the subunit composition or the nature of the native enzyme, although different molecular weights have been suggested.

In order to understand the nature of the catalytic subunit of *A. xylinum* cellulose synthase and determine its relationship with cellulose synthases from other organisms, we have cloned and sequenced the gene for the cellulose synthase catalytic subunit. The gene was isolated using oligonucleotide probes designed from the N-terminal amino acid sequence of the 83 kDa polypeptide. The amino acid sequence and properties of the polypeptide predicted from the sequence of the cloned gene were found to be in agreement with the N-terminal amino acid sequence and biochemical data available for the cellulose synthase catalytic subunit.

**Materials and methods**

**Materials**

Polynucleotide kinase (10000 U/ml) was obtained from New England Biolabs, Beverly, MA. \( [\gamma{}^{32}P] \)ATP (6000 Ci/mmole), \( [\alpha{}^{32}P] \)dCTP (3000 Ci/mmole), and \( [\beta{}^{35}S] \)dATP (1000–1350 Ci/mmole) were obtained from NEN Research Products, Boston, MA. Restriction endonucleases were obtained from BRL, Gaithersburg, MD. T4 Ligase was from BRL and Promega, Madison, WI. Oligonucleotides were synthesized by Operon Technologies, San Pablo, CA.

**Strains and plasmids**

*A. xylinum* strain ATCC 53582 was used in the present study for obtaining the 83 kDa polypeptide for N-terminal amino-acid sequence and as a source of DNA for cloning the gene. *A. xylinum* strains ATCC 10245, ATCC 23769, and NQ-3 were used for determining DNA homology with the oligonucleotide probes and the cloned cellulose synthase catalytic subunit gene. *E. coli* HB101 was used as the host strain for DNA cloning and JM103 was used as a host for M13 vectors for DNA sequencing. Plasmid pUC18 was used for cloning the *A. xylinum* DNA.
Amino-acid sequencing

For determining the N-terminal amino-acid sequence of the 83 kDa polypeptide, purified cellulose synthase was loaded on a 8% LDS-polyacrylamide gel and the individual polypeptides separated by electrophoresis [13]. The polypeptides from the polyacrylamide gel were electro-photographically transferred [17] to Immobilon-P transfer Membrane (Millipore). The membrane was stained with Coomassie blue and the 83 kDa polypeptide band was cut out with a razor blade. The polypeptide present on the excised membrane band was sequenced using an Applied Biosystems Model 477A Pulse Liquid Phase Sequenator (at the Department of Zoology, University of Texas at Austin).

Southern blotting and hybridization

DNA fragments, obtained following digestion with restriction endonucleases, were separated by agarose gel electrophoresis and transferred to nylon membrane (GeneScreen) using a rapid transfer protocol [20]. For hybridization, oligonucleotides were labelled with $\gamma^{32}$P-ATP and polynucleotide kinase according to standard procedures [16]. Hybridizations were done for 16–24 h at 45 °C in 5 × SSC, 50 mM sodium phosphate (pH 7.0), 10 × Denhardt’s solution, 7% SDS, and 100 µg/ml sheared and denatured salmon sperm DNA. Membranes were washed for 1 h at 45 °C in 3 × SSC, 10 mM sodium phosphate (pH 7.0), and 5% SDS followed by a wash in 1 × SSC and 1% SDS. For higher stringency, membranes were washed in 1 × SSC and 1% SDS at higher temperatures for 5–10 min. Membranes were exposed to X-ray films (X-Omat AR, Kodak) at −80 °C for autoradiography.

DNA manipulations

Total A. xylinum DNA was isolated essentially according to the procedure followed for isolation of DNA from E. coli [24]. Plasmid DNA from E. coli was isolated according to the alkaline lysis procedure [1]. Single-stranded M13 DNA was isolated according to the procedure described in the BRL instruction manual for M13 cloning/dideoxy sequencing. DNA was digested with restriction endonucleases following the manufacturer’s recommendations and the digestion products were analysed by electrophoresis, usually on a 0.7% agarose gel. DNA ligation was done using T4 ligase in buffer supplied by the manufacturer, either for 4 h at room temperature or overnight at 15 °C. For transformation, competent cells of E. coli were prepared according to the procedure of Hanahan [9] and were stored frozen at −80 °C.

DNA sequencing

DNA fragments were cloned in the multiple cloning site of M13mp18 and M13mp19 for sequencing. DNA sequencing of both the strands was performed according to the dideoxy chain-termination procedure [22]. The DNA was labelled with [35S]-dATP using the Sequenase version 2.0 kit from U.S. Biochemical Corp., Cleveland, OH. For a large region, both dGTP and dITP reactions were performed to resolve regions of compression. Sequencing from the end was done using the primer provided in the Sequenase kit. For sequencing regions that were distant from the ends of some clones, synthetic oligonucleotide primers designed from the DNA sequence were used. The DNA sequence was analyzed by using the PC/GENE programs (IntelliGenetics, Inc.).

Results

N-terminal amino acid sequence of the 83 kDa polypeptide

During the purification of cellulose synthase from A. xylinum ATCC 53582, it was observed by LDS-polyacrylamide gel electrophoresis that
trypsin treatment of either the washed membranes or the detergent-solubilized enzyme preparation resulted in the degradation of polypeptides other than the 83 kDa polypeptide and at the same time maintained the cellulose synthase activity [13]. Since trypsin treatment proved to be an efficient method for obtaining relatively pure 83 kDa polypeptide, N-terminal amino acid sequencing of the 83 kDa polypeptide purified from membrane preparations that were treated with trypsin was done. The amino acid sequence obtained is shown below:

Ser-Ser-Thr-Gln-Ser-Glu-Ser-Gly-Met-Ser-Gln-Trp/Leu-Met-Gly-Lys-

**Design of oligonucleotide probes**

For designing oligonucleotide probes from the amino acid sequence, a stretch of 6 amino acids (-Met-Ser-Gln-Trp/Leu-Met-Gly-) was selected from the sequence of the 83 kDa polypeptide that was obtained from trypsin-treated membrane preparation. Eight 17-mer oligonucleotides, each with a unique sequence and having deoxyinosine at the ambiguous position of the serine codons were synthesized. The sequence of the oligonucleotides is given below:

83-1A 5’-ATGTCICAATGGATGGG-3’
83-1G 5’-ATGTCICAGTGATGGG-3’
83-2A 5’-ATGAGICAATGGATGGG-3’
83-2G 5’-ATGAGICAGTGATGGG-3’
83-3A 5’-ATGTCICAACGTGATGGG-3’
83-3G 5’-ATGTCICAGCTGATGGG-3’
83-4A 5’-ATGAGICAACGTGATGGG-3’
83-4G 5’-ATGAGICAGCTGATGGG-3’

I = deoxyinosine

**Identification of *A. xylinum* DNA fragments homologous to oligonucleotide probes**

To determine if the oligonucleotides showed any homology with *A. xylinum* DNA, individual oligonucleotides were used as probes in Southern hybridization with *A. xylinum* ATCC 53582 DNA digested with HindIII. Under conditions of low stringency washing, a single fragment of 9.5 kb showed an intense signal of hybridization with oligonucleotide probes 83-1A and 83-1G, and a faint signal with 83-2G. However, with more stringent washings no signal was observed in membranes which had been probed with 83-1A and 83-2G. Similar washings did not remove the signal from membranes probed with 83-1G, suggesting that a DNA sequence highly homologous to the oligonucleotide 83-1G sequence was present in *A. xylinum*. Probing of HindIII-cleaved DNA from *A. xylinum* ATCC 23769 and a mutant of *A. xylinum* ATCC 53582 defective in cellulose production [23] with oligonucleotide 83-1G similarly showed a single intense band of hybridization (Fig. 1). Less intense hybridization signals were observed when DNA from *A. xylinum* strains NQ-3 and ATCC 10245 were probed with 83-1G, suggesting that the 83-1G sequence may be partly conserved in different strains of *A. xylinum*. DNA from *A. xylinum* strains ATCC
53582 and ATCC 23769 cleaved with restriction enzymes BglII, ClaI, NarI, NruI, and SmaI and probed with 83-1G showed a single band of hybridization in all cases, suggesting that the 83-1G homologous sequence is present in a single copy in the A. xylinum genome.

**Cloning of the cellulose synthase catalytic subunit gene from A. xylinum**

For cloning the cellulose synthase catalytic subunit gene from A. xylinum, DNA from A. xylinum ATCC 53582 was digested with HindIII and the fragments separated on an agarose gel. DNA fragments in the size range of 9.5 kb were extracted from the gel slice and ligated to HindIII-cleaved plasmid pUC18. The ligation mix was used to transform E. coli HB101, and plasmid pools from ampicillin-resistant transformants were screened by hybridization to 83-1G. A plasmid, pIS532, was identified from this screening to carry a DNA fragment homologous to the oligonucleotide 83-1G. pIS532 carried a 9.5 kb HindIII fragment of A. xylinum ATCC 53582. Southern hybridization of pIS532 restriction fragments with 83-1G led to the identification of the region on the 9.5 kb HindIII fragment that carried the gene for the cellulose synthase catalytic subunit.

**N-terminal amino acid sequence analysis**

A region of approximately 3 kb from the plasmid pIS532 was sequenced to obtain the DNA sequence of the cellulose synthase catalytic subunit gene. From the sequence obtained, an ORF of 2169 bp was identified that coded for a polypeptide of 80218 Da (Fig. 2). The N-terminal amino acid sequence of the cellulose synthase catalytic subunit polypeptide that had been obtained from trypsin-treated membrane preparation was identical to the amino acid sequence from position 6 to position 20 of the amino acid sequence deduced from the DNA sequence. In the N-terminal sequence of the cellulose synthase catalytic subunit polypeptide it was unclear if position 12 was occupied by tryptophan or leucine, and since this amino acid position was used in the design of the oligonucleotides, the single codon of tryptophan (TGG) was used in 4 of the oligonucleotides while the other 4 oligonucleotides had the sequence for the most frequently used leucine codon (CTG). From the DNA sequence and the deduced amino acid sequence it is clear that this position is occupied by tryptophan. Since no N-terminal methionine residue was observed in the amino acid sequence of the cellulose synthase catalytic subunit polypeptide that was purified from trypsin-treated membrane preparations, it was uncertain whether this amino acid sequence was the N-terminal end of a processed polypeptide from which a signal sequence had been cleaved, or whether the N-terminal region of the native cellulose synthase catalytic subunit polypeptide had been cleaved by trypsin. Amino acid sequencing of the cellulose synthase catalytic subunit polypeptide obtained from membrane preparations that had not been treated with trypsin was also performed. Even though the sequence obtained showed ambiguities at a number of positions, it unambiguously showed methionine at the N-terminus and the sequence -Glu-Val-Arg-X-X-Thr-Gln- which matches with positions 3 to 9 of the amino acid sequence deduced from the DNA sequence shown in Fig. 2. From the N-terminal amino acid sequence of the cellulose synthase catalytic subunit purified from trypsin-treated membranes and from the amino acid sequence deduced from the DNA sequence, it appears that trypsin cleaved five amino acids from the N-terminal end without in any way affecting the enzyme activity. Such a cleavage would cause no detectable change in the mobility of the polypeptide during polyacrylamide gel electrophoresis, and no such change was indeed observed.

**Sequence analysis of the cellulose synthase catalytic subunit gene**

The overall G + C content of the gene for the cellulose synthase catalytic subunit was found to
CGCTTCCGCACATCAGCATCGTCCATAAGTGAGCGTCGGTGACAGGGGTGTTGCCGATG
AATGGAGTCTCTTGCTGCGAACGCCTGCAGGCGGACCTGGCGTCGCCGCCTCCGGCATCTGA
GCTTTTTTCTTTCTAGGGGATGGGCAGCCACCCCTGATGAAAAAGAGTTTCTTTTCCTCGTGCGT
CCAGCGATCGAGTTCTCAATGATTGCTGACGATCAGTACCACTACCACCGCCTCGCT
CGTCGGACAGGAGGTCTGGTGTTCTGCGGAAACAGAAGAGAATCCTAAGGCGCTATAT
TCAGGGCCAGCCCTGCTGTGTCGTTCTGGTGAATGCGGGGAGGGGACAGACAC
GTCGTCGTCGTCGCGGGCTCTATTCCCCCTTTTGGCTGTGAGTTTCGCTGGGGGAC
CGCCCGCGCTTTGCCGCCGATGTCGCGCATGTCGCTGGGGCTGAGTCTGGGTTTTGAGC
TTTCTCTTTTACAGCATTTTGGAGCGAGTTTTTGTTATGCGAGAGTTCGCTGCTGCAACG

MPEVRSST

6
28
48
68
88
108
128
148
168
188
208
228
248
268
288
308
328
348
368
be 58 mol\%, with values for codon positions 1, 2, and 3 to be 60, 44, and 69 mol\% respectively. That the ATG at position 636 of the DNA sequence is the start codon is suggested from the open reading frame analysis of the complete sequence, since the polypeptide coded by the ORF starting at this position is close to the molecular weight determined for the cellulose synthase catalytic subunit. In this reading frame no other ATG was observed between the stop codon at position 408 and the ATG at position 636. The N-terminal amino acid sequence of the cellulose synthase catalytic subunit from trypsin- treated membranes matched with the deduced
amino acid sequence of this ORF. The presence of sequences corresponding to the E. coli promoter have not been observed upstream of the start codon of the cellulose synthase catalytic subunit gene, nor have they been observed anywhere else in the complete sequence. Preliminary data suggest that the cellulose synthase catalytic subunit gene is not expressed in E. coli when it is under the control of its own promoter. Genes cloned from other Acetobacter species similarly have been shown to carry promoters that do not function in E. coli [5, 6]. A potential ribosome-binding site GGACGAGT is present 5 base pairs upstream of the initiating ATG codon.

Discussion

Cellulose biosynthesis in A. xylinum has been studied in considerable detail at the ultrastructural level. More recently the final step of the pathway and its regulation has become better understood at the biochemical level. Cellulose biosynthesis is believed to be controlled in vivo by a complex multi-component regulatory system, and in vitro by the allosteric activator c-di-GMP [21]. This suggests that cellulose synthase has at least 2 sites, a catalytic site and a regulatory site, to function efficiently. Whether these sites are present on a single polypeptide chain or on separate subunits of an enzyme complex [18] is not very clear at present. The native enzyme has however been suggested to be oligomeric in nature with a $M_r$ in the range of 400–500 kDa [13, 18].

From the DNA sequence, the cellulose synthase catalytic subunit has been predicted to have 723 amino acids and a $M_r$ of 80 kDa. The amino acid composition of the polypeptide (Table 1) shows that it is hydrophobic in nature (Leu + Ala + Ile + Pro + Val + Phe + Met + Trp = 50.2 mol%), in agreement with its observed location in the cytoplasmic membrane [2]. The hydrophobicity plot determined by the method of Kyte and Doolittle [11] shows that the polypeptide is composed of a number of hydrophobic regions (Fig. 3). Analysis of the amino acid sequence by the method of Eisenberg et al. [4] predicts that the polypeptide has 10 membrane-associated helices based on the average hydrophobicity of a 21 residue segment. The positions of the membrane-associated helices were classified with the aid of the hydrophobic moment plot by the method of Eisenberg et al. [4] and using the PC/GENE program Helixmem into monomeric transmembrane, multimeric transmembrane, globular and surface. Of the 6 transmembrane helices in the cellulose synthase catalytic subunit, the first helix was determined to be monomeric transmembrane based on highest hydrophobicity and lowest hydrophobic moment. Such monomeric transmembrane regions are believed to be membrane anchors [4]. From such an analysis, it appears that the cellulose synthase catalytic subunit is anchored in the membrane at its N-terminal end by a single hydrophobic helix. The remaining 5 transmembrane segments have the potential to interact with other helices. The inner boundaries of the 6 transmembrane segments predicted by the method of Eisenberg et al. [4] and determined by the method of Klein et al. [10] are shown in Fig. 2. No signal sequence has

<table>
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<th>Amino acid</th>
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<tr>
<td>Leucine</td>
<td>11.4</td>
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<tr>
<td>Alanine</td>
<td>8.8</td>
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<tr>
<td>Glycine</td>
<td>8.7</td>
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<tr>
<td>Isoleucine</td>
<td>7.1</td>
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<tr>
<td>Proline</td>
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<td>Valine</td>
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<tr>
<td>Arginine</td>
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<tr>
<td>Threonine</td>
<td>5.6</td>
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<tr>
<td>Phenylalanine</td>
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<tr>
<td>Serine</td>
<td>5.6</td>
</tr>
<tr>
<td>Glutamine</td>
<td>4.2</td>
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<tr>
<td>Glutamic acid</td>
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<tr>
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<td>Tyrosine</td>
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<td>Asparagine</td>
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<td>Methionine</td>
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<td>Lysine</td>
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<tr>
<td>Cysteine</td>
<td>1.1</td>
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been observed at the N-terminus of the deduced amino acid sequence; therefore, it is very likely that the cellulose synthase catalytic subunit is anchored in the membrane by a hydrophobic helix. An understanding of the interaction of the cellulose synthase catalytic subunits among themselves and with other putative subunits of the cellulose synthase complex will be important in determining the organization of cellulose synthase in the cytoplasmic membrane and the way this organization influences cellulose production. Such an understanding will be made possible by using gene fusions and mutational analysis with the cloned cellulose synthase catalytic subunit gene for identifying regions of the polypeptide chain involved in interaction with various components of the cellulose synthase structural and regulatory components as well as the cytoplasmic membrane.

The cellulose synthase catalytic subunit was previously characterized as a glycoprotein by lectin affinity chromatography, and Schiff-periodate and fluorescein isothiocyanate-concanavalin A staining analysis [13]. From sequence analysis of the cellulose synthase catalytic subunit polypeptide, 2 N-glycosylation sites (amino acid positions 277 and 294) were predicted based on the consensus sequence Asn-X-Ser/Thr. Even though this consensus sequence is derived from studies with eukaryotic glycoproteins, it has also been shown to be the acceptor sequence for glycosylation in bacteria [12]. The specific role, if any, that glycosylation may play in the organization or function of cellu-
lose synthase activity is not clear. In this respect, it is interesting that a set of integral membrane proteins related to gliding motility in *Cytophaga johnsonae* have been suggested to be glycoproteins [19]. Furthermore, these protein bands were found to shift to lower positions on SDS-polyacrylamide gels in some of the non-motile mutants and return to the wild-type position in some motile revertants, suggesting the role of post-translational modifications, possibly glycosylation, for this variation [19].

Comparison of the predicted cellulose synthase catalytic subunit polypeptide sequence with other sequences in the NBRF protein sequence data base revealed no significant homologies with previously reported proteins. The cellulose synthase catalytic subunit gene however seems to be conserved in the few *A. xylinum* strains that were analyzed for DNA homology by Southern hybridization. Whether this gene is conserved among the other cellulose synthesizing organisms, including higher plants, remains to be studied. If there exists a common enzymatic mechanism for cellulose synthesis, as suggested by immunochi- emical analysis [18], then the *A. xylinum* gene will be a suitable probe for isolating and identifying the corresponding cellulose synthase gene from higher plants.

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**References**


