The non-catalytic cellulose-binding domain of a novel cellulase from
*Pseudomonas fluorescens* subsp. *cellulosa* is important for the efficient
hydrolysis of Avicel

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A genomic library of *Pseudomonas fluorescens* subsp. *cellulosa*
DNA, constructed in λZAPII, was screened for carboxymethyl-
cellulase activity. The pseudomonad insert from a recombinant
phage which displayed elevated cellulase activity in comparison
with other cellulase-positive clones present in the library, was
excised into pBluescript SK− to generate the plasmid pC48. The
nucleotide sequence of the cellulase gene, designated *celE*,
revealed a single open reading frame of 1710 bp that encoded a
polypeptide, defined as endoglucanase E (*CelE*), of *M* subsp.
59663. The deduced primary structure of *CelE* revealed an N-terminal
signal peptide followed by a 300-amino-acid sequence that
exhibited significant identity with the catalytic domains of
cellulases belonging to glycosyl hydrolase Family 5. Adjacent to
the catalytic domain was a 40-residue region that exhibited
strong sequence identity to non-catalytic domains located in two
other endoglucanases and a xylanase from *P. fluorescens*. The C-
terminal 100 residues of *CelE* were similar to Type-I cellulose-
binding domains (CBDs). The three domains of the cellulase
were joined by linker sequences rich in serine residues. Analysis
of the biochemical properties of full-length and truncated deriva-
tives of *CelE* confirmed that the enzyme comprised an N-
terminal catalytic domain and a C-terminal CBD. Analysis of
purified *CelE* revealed that the enzyme had an *M* of 56000 and
an experimentally determined N-terminal sequence identical to
residues 40–54 of the deduced primary structure of full-length
*CelE*. The enzyme exhibited an endo mode of action in hydro-
lysing a range of cellulotic substrates including Avicel and acid-
swollen cellulose, but did not attack xylan or any other hemicel-
luloses. A truncated form of the enzyme, which lacked the C-
terminal CBD, displayed the same activity as full-length *CelE*
against soluble cellulose and acid-swollen cellulose, but exhibited
substantially lower activity than the full-length cellulase against
Avicel. The significance of these data in relation to the role of the
CBD is discussed.

INTRODUCTION

It is widely recognized that many cellulases have a modular
structure, comprising a catalytic domain linked to a non-
catalytic cellulose-binding domain (CBD) by flexible linker
sequences rich in hydroxyamino acids [1]. Based on sequence
identity there are at least four families of CBDs in which all
members of a given family are thought to have evolved from a
common progenitor sequence [2]. The role of the CBD in enzyme
function has been investigated in several enzymes. Although
these domains do not appear to affect the activity of fungal
cellobiohydrolases against soluble substrates, they significantly
enhance the capacity of the enzymes to hydrolyse crystalline
cellulose [3]. Knowles et al. [4] suggested that fungal CBDs, in
binding to cellulose chains, disrupt the interchain hydrogen
bonding and funnel the glucose polymers into the active site of
the enzyme. Bacterial CBDs can either exhibit a high affinity for
their ligand, such as the Type-I domains [5], or a low affinity for
cellulose as demonstrated by the CBD of endoglucanase E (*CelE*)
from *Clostridium thermocellum* [6]. It appears that this latter
domain decreases the specific activity and increases the affinity
of the enzyme for soluble cellulase substrates such as β-glucan and
carboxymethylcellulose (CMC). The enzyme exhibits no activity
against insoluble cellulose. In contrast, it is clear that at least
some high-affinity Type-I CBDs enhance the capacity of cellulases
to hydrolyse crystalline cellulose [7,8].

Studies in our laboratories have focused on the molecular
architecture of plant cell wall hydrolases of *Pseudomonas fluores-
cens* subsp. *cellulosa*. This aerobic soil bacterium expresses
multiple endoglucanases and xylanases in addition to enzymes
that cleave xylan side-chain substituents [9,10]. The pseudomonad
also expresses other plant cell wall hydrolases that attack
mannan, galactan and arabinan ([11]; H. J. Gilbert, unpublished
work). To date, the primary structures of two endoglucanases
[12,13], two xylanases [14,15], an arabinofuranosidase [15], an
acetyl xylan esterase [5] and a cellodextrinase [16] have been
determined. Each enzyme has a modular structure comprising
of a catalytic domain, a middle non-catalytic domain of unknown
function and a terminal Type-I CBD. The location of CBDs in
a range of plant cell wall hydrolases that do not attack cellulose
led to the suggestion that in *P. fluorescens* subsp. *cellulosa*, CBDs
could provide a general mechanism by which a consortium of
hydrolases accumulate on the surface of the plant cell wall
resulting in an enhancement, through their close proximity, of
the synergism exerted between the enzymes [12,17]. To
investigate this possibility it is necessary to analyse the relationship
between the structure and function of a cellulase from *P. fluores-
cens* subsp. *cellulosa*. Although two cellulase genes from this bacterium

Abbreviations used: BMCC, bacterial microcrystalline cellulose; CBD, cellulose-binding domain; *CelE*, endoglucanase E; *CelE′*, truncated *CelE*; CMC, carboxymethylcellulose; LB, Luria broth; ORF, open reading frame; PC, 50 mM potassium phosphate/12 mM citric acid buffer (pH 6.5).

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The nucleotide sequence data for *celE* reported in this paper will appear in the EMBL, GenBank and DDBJ Nucleotide Sequence Databases under the accession number X86798.
have previously been cloned, they were poorly expressed in *Escherichia coli*, and we were unable to purify sufficient quantities of the encoded enzymes to evaluate the role of their CBDs in cellulose cleavage. In this report we describe the relationship between the structure and function of a third *Pseudomonas* cellulase that was successfully overexpressed in *E. coli*. The data revealed that the enzyme had a modular structure consisting of an N-terminal catalytic domain, a small non-catalytic domain also found in other pseudomonad cellulases, and a C-terminal Type-I CBD. The cellulase displayed a typical endo-mode of action and exhibited significant activity against insoluble and substantially crystalline cellulose. Removal of the CBD did not affect the activity of the enzyme against soluble or amorphous cellulose, but there was a reduction in the activity of the truncated cellulase against Avicel. It is apparent, therefore, that Type-I CBDs from *P. fluorescens* subsp. *cellulosa* cellulases can play a direct role in the hydrolysis of substantially crystalline substrates; the domain does not simply provide a mechanism by which a repertoire of hydrolases can bind to the plant cell wall.

**MATERIALS AND METHODS**

**Bacterial strains, culture conditions and vectors**

*E. coli* JM83 was cultured in Luria broth (LB) at 37 °C. Phage were propagated in *E. coli* grown on NZY medium. Media were supplemented with 100 μg/ml of ampicillin and 2 μg/ml of 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside to select for *E. coli* transformants and recombinants, respectively. To identify recombinant *E. coli* strains that expressed cellulase activity, bacterial colonies, grown on LB-agar supplemented with medium-viscosity CMC, were stained with 1% Congo Red for 5 min and destained with 1 M NaCl. Cellulase-expressing *E. coli* colonies were surrounded by clear haloes against a red background [18]. The phage and plasmids employed in this work were λZAPII, pBluescriptSK− (Stratagene) and the pMTL series of vectors [19].

**Recombinant DNA methodology**

Plasmid DNA was purified using QiaGen columns (Hybaid) following the instructions of the manufacturer. Transformation of *E. coli*, the general use of nucleic acid-modifying enzymes and Southern hybridization were as described by Sambrook et al. [20]. DNA probes were labelled with [α-32P]dCTP by random priming. *Pseudomonas* genomic inserts in recombinants of λZAPII expressing cellulase activity were excised into pBluescript SK− using VCS-M13 helper phage.

To sequence *celE* a series of nested deletions of the fragment carrying the gene was constructed using the exonuclease III/S1-nuclease kit supplied by Promega. The nested deletions were sequenced by the dideoxy chain termination method of Sanger et al. [21] both manually, using Sequenase (United States Biochemicals) and spin dialysis to purify denatured DNA [22], and by an automated method in which the nested deletions were sequenced using a PRISM ready reaction dye-deoxy terminator cycle, supplied by Applied Biosystems Inc., and the products analysed using an Applied Biosystems automated sequencer (model 373A). Sequences were compiled and ordered using the computer programs of Staden [23]. Both strands of *celE* were sequenced.

**Assays**

Avicel PH105 was obtained from FMC Corporation; bacterial microcrystalline cellulose (BMCC; [8]) was a generous gift from Dr. Bernard Henrissat. Amorphous (acid-swollen) cellulose was prepared from crystalline cellulose as described by Wood [24].

Cellulase, xylanase and mannanase activities were assayed at 37 °C, unless otherwise stated, in 50 mM potassium phosphate/12 mM citric acid buffer (pH 6.5) (PC). The polysaccharides were incubated with enzyme at a final concentration of 0.2% (unless otherwise stated), and the level of reducing sugar released was determined [25]. One unit of enzyme activity was defined as the amount of enzyme that releases 1 μmol of reducing sugar per min. When using insoluble substrates, the residual polysaccharide was removed by centrifugation, before analysing the liquid phase for reducing sugar. Hydrolysis of red debranched arabinan and azurin-cross-linked galactan (both from Megazyme) was determined by culturing *E. coli* harbouring the *celE* gene on substrate-containing agar and looking for the appearance of clear halos against a red background or the solubilization of the blue AZCL dye, respectively. Cellulo-oligosaccharides (200 ng/ml) were incubated with *CelE* and aliquots were removed at regular intervals, boiled for 5 min and applied to a Dionex PA1 column. Separation of cellulooligosaccharides was achieved using a gradient elution of 0–75 mM sodium acetate (in 100 mM NaOH) from 0–16 min, and isocratic elution with 75 mM sodium acetate (in 100 mM NaOH) from 16–25 min. The sugars were detected with a pulsed amperomter. The α or β configuration of the anomeric carbon of cellotriose, generated by the hydrolysis of cellulopentaose by *CelE*, was determined as described previously [26], except that the two anomers of cellotriose were separated on two Spherisorb HPLC columns (S5 ODS2; 25 cm × 4.6 mm) run in series. The oligosaccharides were detected as above after the addition of post-column reagent (300 mM NaOH). Protein was determined by the dye binding method of Sedmak and Grossberg [27] using BSA as the standard. The sizes of polypeptides were determined by SDS/PAGE [28]. Viscometric assays were performed as described by Zhou et al. [29]. The N-terminal sequence of *CelE* was determined by the method of Hunkapillar et al. [30].

**Purification of CelE and CelE’**

*E. coli* XL1-blue harbouring full-length *celE* in plasmid pC48 was cultured overnight in LB broth (3 l) containing ampicillin (100 μg/ml). A cell-free extract was prepared by sonicating the harvested cells in 20 mM Tris/HCl, pH 8.0, and *CelE* contained in the cell-free extract was precipitated between 40% and 60% ammonium sulphate saturation. Precipitated protein was redissolved in a minimum volume of 20 mM Tris/HCl, pH 7.2, and applied to a Sephacryl S-100 HR column (25 mm × 610 mm) which was eluted with 20 mM Tris/HCl, pH 7.2, containing 150 mM NaCl at a flow rate of 0.5 ml/min. Purification of full-length *CelE* was greatly facilitated by the apparent interaction of *CelE* with the Sephacryl matrix, which resulted in retention of the enzyme on the column during the passage of almost a full column volume of the elution buffer. A truncated derivative of *CelE* (CelE’), having the same N-terminal amino acid sequence as the full-length enzyme, but lacking the C-terminal CBD, was generated by treating *CelE* with trypsin. Purified full-length *CelE* was incubated for 16 h at 37 °C in Puck’s saline containing trypsin (0.125 mg/ml) and EDTA (50 μg/ml). SDS/PAGE revealed that *CelE* (*M*ₚ 56000) had been quantitatively converted into a smaller polypeptide (CelE’) with an *M*ₚ of 33000. After dialysing against 20 mM Tris/HCl, pH 8.0, CelE’ was purified by FPLC on Mono Q (Pharmacia) using a 1 ml analytical column and a 40 ml gradient containing from 0 to 1 M KCl in 20 mM Tris/HCl, pH 8.0, at a flow rate of 60 ml/h.
RESULTS

Isolation and characterization of celE

The primary objective of this study was to evaluate whether CBDs from *P. fluorescens* subsp. *cellulosa* cellulases enhance cellulolysis. This was not feasible with the two previously cloned *Pseudomonas* endoglucanase genes, as the gene products were synthesized at low levels in *E. coli*. To determine whether it was possible to isolate a *Pseudomonas* cellulase gene that can be highly expressed in *E. coli*, a genomic library of *P. fluorescens* subsp. *cellulosa* DNA, constructed in AZAPII, was screened for clones that exhibited carboxymethylcellulase activity. A recombinant phage, which appeared to produce high levels of cellulase activity, was selected for further study. The pseudomonad genomic insert was excised from the recombinant phage into pBluescriptSK- to generate the plasmid pC48. Cross-hybridization experiments showed that the cellulase gene (designated celE) encoded by pC48 exhibited no detectable sequence identity with either of the two previously characterized *P. fluorescens* endoglucanase genes. A restriction map of the *Pseudomonas* DNA in pC48 is displayed in Figure 1. Deletion and subcloning experiments located the position of celE within the pseudomonad insert in pC48 between map positions 1.7 and 3.5.

Characterization of CelE

The celE gene product, designated CelE, was purified from *E. coli* cell-free extract and its biochemical properties evaluated. The enzyme had an *M*₅₆ of 56000 (Figure 2) and a broad pH optimum between pH 4.5 and 8.0 (data not shown). The cellulase was also unusually thermostable for an enzyme derived from a mesophilic bacterium; significant thermal inactivation of CelE occurred at temperatures in excess of 60 °C. The N-terminal sequence of the cellulase was DVAPLSVQGNKILAN. Analysis of the substrate specificity of CelE revealed that the enzyme hydrolysed a range of cellulose substrates, exhibiting maximum activity against barley β-glucan (Table 1). The major products generated from CMC hydrolysis were cellobiose and cellobiose (data not shown). Analysis of the activity of the enzyme against cellulo-oligosaccharides showed that the cellulase had no activity against cellobiose or celiotriose, although it hydrolysed longer-chain cellulo-oligosaccharides such as cellotriose and cello-pentaose displaying 2.5 times higher activity against the larger oligosaccharide. The major products generated from cellopentaose were cellobiose and celiotriose, while cello-pentaose was hydrolysed primarily to cellobiose (data not shown).

![Figure 1] Restriction map of recombinant plasmids containing celE

Cleavage sites for restriction endonucleases are as follows: *Bcl* (B), *Cal* (C), *EcoRV* (E), *NcoI* (Nc), *NruI* (N), *PstI* (P) and *SphI* (S). The location and orientation of celE is indicated by an arrow. In pC53 the 5'-truncated derivative of celE was in-frame with the lacZ translational start codon of the vector (pMTL22). The catalytic activity of CelE encoded by the recombinant plasmids is shown by + or –.

![Figure 2] Purification of full-length CelE by gel filtration chromatography on Sephacryl S-100 HR

(a) Column chromatography was performed as described in the Materials and methods section. Endoglucanase activity (■) and protein concentration (–) were measured by quantifying reducing sugar release after fractions were incubated with CMC, and by monitoring *A₂₈₀*, respectively. (b) SDS/PAGE analysis using a 10% (w/v) polyacrylamide gel of cell-free extract from *E. coli* containing pC48 (lane 1), the ammonium sulphate fraction (lane 2) and Sephacryl S-100 HR column fractions 50 (lane 3), 51 (lane 4) and 53 (lane 5). Protein molecular mass markers were run on the extreme left.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Relative activity (%)</th>
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<tbody>
<tr>
<td>Barley β-glucan</td>
<td>100</td>
</tr>
<tr>
<td>Carboxymethylcellulose</td>
<td>45</td>
</tr>
<tr>
<td>Lichenan</td>
<td>120</td>
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<tr>
<td>Xylan</td>
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<tr>
<td>Galactan</td>
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</tr>
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<td>Arabinan</td>
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<td>Galactomannan</td>
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<tr>
<td>Laminarin</td>
<td>NA</td>
</tr>
<tr>
<td>Mannan</td>
<td>NA</td>
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Table 1 Substrate specificity of CelE

The specific activity of purified full-length CelE against barley β-glucan was 350 units/mg of protein. Abbreviation: NA, no activity detectable.
The cellotriose released from cellopentaose by CelE (5 min enzyme reaction) co-chromatographed with the $\beta$ anomer of cellotriose (Figure 3). After incubating these reaction products for a further 1 h at room temperature, cellotriose had mutarotated and significant quantities of the $\alpha$ anomer were observed, confirming that the separation method resolves the two anomeric forms of the trisaccharide. Thus, the cellulase retains the configuration of the anomeric carbon during bond cleavage. CelE therefore hydrolyses glycosidic bonds by a double displacement mechanism. To assess whether CelE was an endo-acting enzyme the reduction of viscosity and release of reducing sugar were determined during incubation with barley $\beta$-glucan. The data (Figure 4) showed that CelE exhibited a typical endo-mode of action, generating a large decrease in viscosity while releasing only small amounts of reducing sugar. The enzyme exhibited no detectable activity against a range of non-cellulosic plant structural polysaccharides including xylans, mannan, galactan and arabinan, or against $\beta$-1,3-linked glucose polymers such as laminarin (Table 1). Although these results indicate that CelE is a typical endo-$\beta$-1,4-glucanase, the enzyme displayed significant activity against both acid-swollen cellulose and crystalline cellulose (Avicel) (Table 2), substrates which are not normally hydrolysed to such a great extent by the majority of endo-glucanases.

**Figure 3** Anomeric configuration of cellotriose generated by CelE

Cellopentaose was incubated with CelE for 5 min (b) and 1 h (c) and the products generated were immediately analysed by HPLC as described in the Materials and methods section. In (d) the oligosaccharides generated in (b) were boiled for 5 min, to mutarotate the $\beta$ anomer, prior to HPLC analysis. (a) Contains cellotriose. The retention times of cellobiose (1), the $\beta$ anomer of cellotriose (2) and $\alpha$ anomer of cellotriose (3) were 6.94 ± 0.06 min ($n = 5$), 10.48 ± 0.03 ($n = 6$) and 10.89 ± 0.03 ($n = 4$), respectively.

**Figure 4** Effect of CelE on the viscosity of CMC

The substrate (2%, w/v) was incubated with CelE in PC buffer, pH 6.5, at 37 °C. Specific viscosity (●) and reducing sugar (■) were measured as described in the Materials and methods section.

**Nucleotide sequence of celE and the primary structure of CelE**

The nucleotide sequence of the region of pc48 containing celE was determined in both strands. The data revealed a single open reading frame (ORF) of 1710 bp encoding a protein of $M_{r}$ 59663 (Figure 5). The codon usage of the ORF was very similar to other *P. fluorescens* plant cell wall hydrolases [5]. Upstream (9 bp) of the putative ATG translational start codon is the sequence AGAGA, which exhibits similarity with the ribosome binding sequences of genes from Gram-negative bacteria. Residues 40–54 of the deduced primary structure of the encoded protein were identical to the experimentally determined N-terminus of mature CelE. The sequence of CelE preceding the mature N-terminus conforms to a typical prokaryotic signal peptide; the hydrophilic N-terminus of the full-length enzyme contains five basic residues and is followed by 15 small hydrophobic residues capable of
forming an α-helix. The $M_r$ of recombinant CelE is similar to the deduced $M_r$ of the mature cellulase (55768).

The primary structure of CelE revealed domains observed in other microbial plant cell wall hydrolases. The 300 N-terminal residues of mature CelE exhibited significant sequence identity with the catalytic domains of endoglucanases belonging to glycosyl hydrolase Family 5 [31,32]. There was substantial sequence identity between residues 377-417 of CelE and the non-catalytic domains, of unknown function, from xylanase A, endoglucanases A and B and cellobiohydrolase C of P. fluorescens subsp. cellulosa plant cell wall hydrolases [2]. The major residues which are conserved in all Type-I CBDs, including four tryptophans and N- and C-terminal cysteines are retained in the CelE CBD homologue. In common with other P. fluorescens subsp. cellulosa plant cell wall hydrolases, the domains of CelE were linked by sequences rich in serine residues.

### Relationship between the structure and function of CelE

To evaluate whether the putative N-terminal catalytic domain homologue constitutes a functional catalytic domain, truncated derivatives of CelE were constructed and their capacity to direct the synthesis of a functional cellulase in E. coli was evaluated. The data presented in Figure 1 demonstrated that truncated CelE lacking 158 C-terminal residues (pC52) retained catalytic activity, but a derivative of the cellulase from which 100 N-terminal residues had been deleted (pC53) did not exhibit endoglucanase activity. These data confirm that the N-terminal 300 residues of mature CelE constitute the catalytic domain. To evaluate whether the C-terminal CBD homologue was biologically active, the capacity of full-length CelE to bind to cellulose was evaluated. The data presented in Figure 7 showed that a polypeptide of $M_r$ 56000, expressed by E. coli containing pC48 bound to Avicel. The polypeptide also bound to acid-swollen cellulose but not to xylan (data not shown). N-terminal sequence analysis of the polypeptide confirmed its identity as CelE. In contrast, a catalytically active but truncated derivative of CelE which had an $M_r$ of 34000 and the N-terminal sequence of DVAPLSVQGNKILAN (corresponding to residues 40-54 of full-length CelE), and was equivalent to the catalytic domain alone (predicted $M_r$ 32879) did not adhere to Avicel. These data suggest that the C-terminal 100-residue domain of CelE constitutes a functional CBD.

To evaluate the importance of the CBD in the capacity of CelE to attack various cellulolytic substrates, the relative activities of full-length CelE and a truncated form of the enzyme, lacking the CBD, against CMC, acid-swollen cellulose, Avicel and BMCC were determined. The data, presented in Table 2, clearly showed that although both forms of the cellulase exhibited similar activities against soluble and amorphous cellulose, the full-length enzyme displayed approximately four times higher activity against Avicel than the truncated derivative. Full-length and truncated CelE exhibited no detectable activity against BMCC.

### DISCUSSION

The data presented in this report describe the cloning and characterization of a highly expressed endoglucanase, designated CelE, from P. fluorescens subsp. cellulosa. The enzyme comprises multiple domains with the catalytic domain derived from glycosyl hydrolase Family 5. In contrast, the catalytic domains of the other two P. fluorescens endoglucanases previously sequenced ([12,13]) are located in glycosyl hydrolase Families 9 and 45, respectively [31,32]. It is apparent, therefore, that the multiple endoglucanases of P. fluorescens resulted from the transfer of several discrete cellulase genes into the pseudomonad. The acquisition of multiple cellulases by lateral gene transfer is a common phenomenon among prokaryotes, while in lower eukaryotes isoforms of endo- and exo-glucanases often arise through gene duplication [32]. Within the classification established for glycosyl hydrolase enzymes, Family 5 is one of the largest groupings and contains enzymes of bacterial and fungal origin. Enzymes of this family are typically endoglucanases with relatively wide substrate ranges, which hydrolyse CMC or barley β-glucan preferentially and display variable but relatively low level activity against acid-swollen cellulose or Avicel. When compared with other endoglucanases belonging to Family 5, CelE from P. fluorescens subsp. cellulosa exhibited much higher activity against acid-swollen cellulose and Avicel. Another unusual feature of CelE is its relatively high thermostability for an enzyme that is derived from a mesophilic soil bacterium. It is possible that this property of the enzyme is a consequence of the same structural elements that confer resistance to proteolytic attack on the catalytic domain. Support for this view is provided by Fontes et al. [33] who showed that there was a clear relationship between the thermostability of plant cell wall hydrolases and their reduced susceptibility to proteolytic attack. It would be rational for CelE to evolve resistance to proteinases as its extracellular location would bring the cellulase into contact with protein-degrading enzymes secreted by other soil bacteria.

Previous studies in our laboratories have identified CBDs in a range of plant cell wall hydrolases expressed by P. fluorescens subsp. cellulosa, several of which do not attack cellulose directly. This led to the hypothesis that in P. fluorescens subsp. cellulosa...
Figure 5  Nucleotide sequence of cefE and the derived amino acid sequence of CefE

The experimentally determined N-terminal sequence of CefE is indicated by underlining. The proposed linker sequences are boxed.
Figure 6 Alignment of CelE with homologous prokaryotic sequences

The middle non-catalytic domain (a) and CBD (b) of CelE were aligned with plant cell wall hydrolyases from P. fluorescens subsp. cellulosa and of CelE, whose residues which showed sequence identity or similarity by all sequence alignments are demarcated by an asterisk. The enzymes compared were as follows: endoglucanase A (EGA; [13]), endoglucanase B (EGB; [12]), xylanase A (XYLA; [14]), xylanase B (XYLB; [15]), cellulodextrinase (CEL; [16]), xylan arabinofuranosidase (XYLD; [15]), and acetyl xylan esterase (XYLE; [5]). The location of the N- and C-terminal residues, in the respective full-length sequences, are indicated.

CeIE plays a direct role in facilitating the hydrolysis of substantially crystalline cellulotic substrates such as Avicel. It could be argued that the CBD is mediating an increase in the hydrolysis of cellulose by bringing the enzyme into intimate contact with its substrate. However, the observation that while the CBD binds to acid-swollen cellulose, it does not enhance the activity of the endoglucanase against this substrate, suggests that the CBD is not causing its effect by simply increasing the concentration of the enzyme at the surface of the polysaccharide. An alternative possibility is that the CBD disrupts the crystalline structure of Avicel facilitating its hydrolysis by CeIE. However, it is apparent that while the cellulase cleaves Avicel it does not attack BMCC. BMCC is among the most highly crystalline forms of cellulose with a crystallinity index (CI) of approximately 80. In contrast, Avicel has a CI of 50 and thus contains amorphous and crystalline regions. Full-length CeE was four times more active against Avicel than the truncated form which lacked the C-terminal CBD. Since CeE has no detectable activity against crystalline cellulose, this suggests that the CBD does not facilitate the hydrolysis of Avicel by disrupting the crystalline structure of the polysaccharide. It would appear, therefore, that the CBD increases the activity of CeE against Avicel by exposing new regions of amorphous cellulose, although the precise mechanism by which the domain mediates increased substrate availability remains to be elucidated. In contrast to the action of the Pseudomonas CBD, Din et al. [34] demonstrated that the isolated CBD from Cellulomonas fimi endoglucanase A could open up the structure of ramie cellulose, making the substrate more accessible to enzyme attack. Similarly, Coutinho et al. [8] demonstrated that removal of the Type-I CBD from C. fimi endoglucanase A diminished the activity of the enzyme against highly crystalline (BMCC), but not amorphous cellulose. Although these data suggest that Type-I CBDs generally increase the capacity of C. fimi cellulases to hydrolyse celluloses which contain a crystalline component, not all Type-I CBDs have the same effect on enzyme activity or exhibit affinity for precisely the same ligands. For example, the C-terminal CBDs of C. fimi xylanase D and Thermomonospora fusca xylanase A exhibit equal affinity for both cellulose and xylan [35,36], while the internal Type-I CBD-homologue of the C. fimi enzyme binds exclusively to xylan [35]. In addition, while fusion of the CBD from T. fusca endoglucanase E to Bacillus subtilis endoglucanase A increased the activity of the enzyme against ball-milled cellulose [7], the cellulase was also more active against amorphous cellulose. Furthermore, the Type-I CBD from P. fluorescens subsp. cellulosa xylanase B did not elicit Avicel hydrolysis when fused to endoglucanases E and A from Cl. thermocellum and Ruminococcus albus, respectively [6]. However, it has been suggested that the lack of activity of the hybrid enzymes against insoluble cellulose could be a consequence of the inability of these catalytic domains to accommodate the insoluble substrate, irrespective of the properties of the CBD [2,7]. It should also be noted that the CBD of C. fimi endoglucanase A greatly increased the activity of the enzyme against BMCC, a substrate not attacked by CeE, even though the pseudomonad enzyme contains a Type-I CBD. This observation supports the view that although CBDs are important in facilitating cellulose hydrolysis, the nature of the active site of endo-acting cellulases also plays a pivotal role in defining the substrates accessible to a specific endoglucanase.

To conclude, it is apparent that the CBD of at least one cellulase from P. fluorescens subsp. cellulosa enhances cellulolysis of Avicel by increasing the exposure of amorphous cellulose to the enzyme, through a mechanism which does not involve the disruption of the crystalline regions of the substrate. This suggests that Type-I CBDs of P. fluorescens cellulases and hemicellulases
do not all play an equivalent role, and argues against the view that the role of the CBDs of the pseudomonad enzymes is to simply increase the synergistic interactions between these plant cell wall hydrolases. However, it remains to be elucidated whether CBDs from *P. fluorescens* subs. *cellulosa* hemicellulases can play an identical role to the CBD from CelE.

We wish to thank the Biotechnology and Biological Sciences Research Council for supporting this work (LE13/138).

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Received 8 December 1994/21 February 1995; accepted 8 March 1995