Cloning, Expression, and Primary Structure of a Chlamydia trachomatis Binding Protein

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The gene encoding an 18,000-dalton eucaryotic cell-binding protein of Chlamydia trachomatis serovar L2 was cloned into Escherichia coli, and the nucleotide sequence of a 1,658-base-pair PstI restriction endonuclease fragment encoding this protein was determined. The recombinant chlamydial gene consists of a 486-base-pair open reading frame encoding a polypeptide of molecular weight 18,314. The resultant polypeptide, comprising 162 amino acids, possesses a highly charged carboxy-terminal end. The expression of this recombinant protein is under the control of a vector promoter. The recombinant 18,000-dalton protein possessed the same eucaryotic cell-binding characteristics as did the native chlamydial 18,000-dalton protein when electrophoresed and transferred to nitrocellulose. Polyclonal antibodies to the recombinant protein exhibited neutralizing activity.

Chlamydiae are obligate intracellular parasites of eucaryotic cells which have been linked to an expanding spectrum of human disease and constitute an important public health problem (6, 19). The infectious form of the organism, elementary bodies (EB), attaches to host cells and, once ingested, initiates the subsequent developmental cycle consisting of intracellular, metabolically active reticulate bodies (25).

Recently, two proteins of molecular ratios (Mr) 18,000 and 31,000, associated with the extracellular EB have been found to bind to eucaryotic cells (5, 28). These Chlamydia trachomatis proteins are of particular interest since the attachment of the pathogen to its host cell represents a critical point of interaction and is necessary for successful invasion. Antibodies raised against these chlamydial cell-binding proteins inhibit association of EB to host cells and possess neutralizing activity as well (26, 28). These data suggest a protective role for such antibodies. The eucaryotic cell-binding proteins are present only on EB and not on the intracellular, noninfectious reticulate bodies. Moreover, the major 18,000-dalton binding protein is present on all C. trachomatis strains which have been examined to date (5, 26) and is a surface-exposed component. The biological importance of the 18,000-dalton protein, coupled with our interest in examining potential C. trachomatis vaccine candidates, prompted us to use a genetic approach to prepare quantities of this protein for subsequent characterization. We describe here the molecular cloning and expression in Escherichia coli of an 18,000-dalton C. trachomatis binding protein and report its complete nucleotide sequence. We named this protein chlanectin because of its cell adhesion properties.

MATERIALS AND METHODS

Organisms. C. trachomatis serovar L2 (L2/434/Bu) was used for preparation of chromosomal DNA. Chlamydiae were grown in HeLa 229 cells for 48 h and purified as described previously (9). E. coli JM83 and JM105 were used as recipients for DNA transformation or transfections (24), and E. coli P678-54 (1) was used as material for minicell preparations. The plasmids used were pUC8, pUC118, and pUC119, which were provided by J. Vieira. E. coli was grown routinely in 2× YT medium (1.6% tryptone, 1% yeast extract, 0.5% sodium chloride). JM83 cells harboring pUC8 or transformants were grown in 100 μg of ampicillin per ml.

DNA enzymes. The restriction enzymes used, T4 DNA ligase, and the large fragment of E. coli DNA polymerase I (Klenow fragment) were purchased from New England BioLabs, Inc., or Boehringer Mannheim Biochemicals. These enzymes were used in conditions specified by the manufacturers.

DNA isolation and manipulations. Chromosomal DNA was isolated from purified EB as described previously (7, 27). Plasmid vector DNA was isolated by the procedure of Birnboim and Doly (2) and purified by cesium chloride gradient centrifugation in the presence of ethidium bromide (17). Chromosomal DNA was partially digested with restriction endonuclease PstI and sized on 0.8% agarose gels (11). DNA fragments (4 to 6 kilobases [kb]) were pooled and ligated to PstI-digested pUC8. Competent JM83 cells were prepared by calcium chloride and transformed with ligated DNA. White ampicillin-resistant recombinant colonies were identified on isopropyl-β-D-thiogalactopyranoside and X-Gal plates.

Expression of the recombinant 18,000-dalton protein. Positive recombinants were grown in YT broth. The cells were harvested, suspended in electrophoresis buffer (0.0625 M Tris, 2% sodium dodecyl sulfate [SDS], 10% glycerol, 0.001% bromophenol blue, 5% β-mercaptoethanol), and boiled for 3 min. Proteins were separated by 12.5% SDS-polyacrylamide gel electrophoresis (SDS-PAGE) (10) and either stained with Coomassie blue or transferred to nitrocellulose (23). Blots were then reacted with 125I-labeled HeLa membranes as described previously (28) or hyperimmune rabbit antiserum to recombinant protein, followed by probing with 125I-labeled protein A. HeLa cell membranes were isolated by a protocol followed earlier (28) and labeled with 125I by the lactoperoxidase method (14).

Analysis of recombinant proteins. E. coli minicells harboring pCT161/18 or pUC8 were grown to stationary phase and purified by sucrose density centrifugation. The cells were labeled with [35S]methionine (4), and then labeled polypep-
was determined by the dideoxy nucleotide chain termination method (18) with M13 mp18 and M13 mp19 recombinant clones (12). The DNA fragments used for sequencing were generated by restriction endonuclease digestion, followed by electrophoresis on 5% polyacrylamide gels. Subclones were subsequently generated, on the basis of sequencing information, either through single or double digests with restriction endonucleases. The 17-base-pair (bp) oligonucleotide primer was purchased from the Regional DNA Synthesis Facility, University of Calgary. Amersham Corp. provided [γ-32P]ATP (3,000 Ci/mmol).

RESULTS

Identification of recombinant chlanectin. A recombinant DNA library comprising 4- to 6-kb fragments of C. trachomatis serovar L2 DNA was constructed by using the expression vector pUC8. From 300 recombinants identified on the basis of ampicillin resistance and insertional inactivation of the lacZ gene, one clone, designated pCT161, encodes a polypeptide of Mr 18,000, identical in size to a chlamydial eucaryotic cell-binding protein. The chimeric pCT161 contains approximately 4.7 kb of C. trachomatis DNA. Complete digestion of pCT161 with the restriction endonuclease PstI generated three fragments, one corresponding to vector DNA and the other two representing the insert. The latter fragments were 3 and 1.7 kb. Subcloning experiments revealed that the sequence encoding the 18,000-dalton protein is confined to the small 1.7-kb PstI fragment (Fig. 1). This subclone, designated pCT161/18, was used in further experiments.

Expression and biological features of the recombinant 18,000-dalton protein. Evidence that the recombinant 18,000-dalton polypeptide is the C. trachomatis binding protein was derived from its binding to 125I-labeled HeLa cell membranes (Fig. 2, lane 5). Labeled HeLa cells also reacted with native

![FIG. 1. Construction of chlanectin expression plasmids. A linearized physical map of insert sequences and their properties is shown. The expression of the recombinant polypeptides, as monitored by their binding to 125I-labeled HeLa cell membranes or by immunoblot assay using hyperimmune rabbit antiserum to cloned chlanectin, is indicated on the right. The cross-hatched bar indicates the protein-coding region of the chlanectin gene on the cloned DNA insert, and the arrow shows the direction of transcription from the vector promoter. Only the relevant restriction sites are shown. Abbreviations: P, PstI; H, HaeIII; X, XbaI.](http://jb.asm.org/)

![FIG. 2. Identification of the chlanectin gene product synthesized by pCT161/18 both in vivo and in vitro. (A) Coomassie blue-stained SDS-PAGE analysis of the polypeptides synthesized by the vector pUC8 and the constructed plasmid pCT161/18. Lanes 1 to 4 represent standard molecular weight markers, pCT161/18 in E. coli K-12 strain JM83, pUC8 in JM83, and serovar L2 EB, respectively. (B) Parallel blot of the gel shown in panel A reacted to 125I-labeled HeLa cell membranes; lanes 5 to 7 are identical to lanes 2 to 4. (C and D) Gel electrophoresis pattern of 35S-labeled gene products synthesized in minicells (C) and the in vitro transcription-translation coupled system (D). Lanes 8 and 10 represent pCT161/18, and lanes 9 and 11 represent pUC8-directed polypeptides. (E) Parallel blot of an unlabeled gel similar to the gel shown in panel D reacted to 125I-labeled HeLa cell membranes (lanes 12 and 13). 18 K, 18,000-dalton protein.](http://jb.asm.org/)

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**TABLE 1**

<table>
<thead>
<tr>
<th>Vector</th>
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FIG. 1. Construction of chlanectin expression plasmids. A linearized physical map of inserted sequences and their properties is shown. The expression of the recombinant polypeptides, as monitored by their binding to 125I-labeled HeLa cell membranes or by immunoblot assay using hyperimmune rabbit antiserum to cloned chlanectin, is indicated on the right. The cross-hatched bar indicates the protein-coding region of the chlanectin gene on the cloned DNA insert, and the arrow shows the direction of transcription from the vector promoter. Only the relevant restriction sites are shown. Abbreviations: P, PstI; H, HaeIII; X, XbaI.

Polypeptides were analyzed by SDS-PAGE, followed by fluorography with En3Hance (New England Nuclear Corp.) to increase the autoradiographic signal.

In vitro transcription translation studies were performed with 1 μg of DNA and 5 μCi of [35S]methionine with E. coli S-30 extract (Boehringer Mannheim). Labeled polypeptides were resolved by SDS-PAGE followed by fluorography and examined by autoradiography. In some experiments, the in vitro-synthesized gene products were reacted to 125I-labeled HeLa cells. In these experiments, [35S]methionine was deleted from the reaction and replaced by unlabeled methionine. The products were resolved and transferred to nitrocellulose after electrophoretic separation and then reacted to 125I-labeled HeLa cells as previously described (28).

**Localization of recombinant protein.** The inner and outer membrane fractions of E. coli JM83 harboring pCT161/18 were separated by the procedure of Dougan et al. (29). The French-pressed pellet (78,000 × g) was dissolved in 3 mM EDTA and layered on 44% (wt/vol) sucrose-3 mM EDTA (pH 7.2) and then centrifuged at 78,000 × g overnight. The upper layer, consisting of inner membrane fraction, was diluted threefold in 3 mM EDTA (pH 7.2), while the outer membrane pellet fraction was suspended and layered on 52%-56% (wt/vol) sucrose-EDTA gradients. After centrifugation at 78,000 × g for 12 h, the band on the 56% sucrose layer was collected and diluted threefold in 3 mM EDTA. Both inner and outer membrane fractions were pelleted and repurified through sucrose twice by centrifugation at 176,000 × g for 1 h.

The periplasmic proteins were isolated by the cold-shock procedure of Dougan et al. (3), while the cytoplasmic proteins were prepared from spheroplasts obtained as described above. The spheroplasts were dissolved in cold water and French pressed three times at 12,000 lb/in². Unlysled cells were removed by centrifugation at 8,000 × g for 15 min, while the supernatant was centrifuged for 1 h at 100,000 × g to remove any crude membrane fraction.

**Neutralization assay.** Hyperimmune rabbit antiserum to recombinant 18,000-dalton protein purified from SDS-PAGE samples was raised as described previously (7, 28). These antisera were then used in neutralization assays as described earlier (26, 28). In each assay, normal rabbit serum was used as a control.

**Sequence determination.** The nucleotide sequence of both strands of the 1.7-kb PstI fragment of plasmid pCT161/18 was deduced from the dideoxy nucleotide chain termination method (18) with M13 mp18 and M13 mp19 recombinant clones (12). The DNA fragments used for sequencing were generated by restriction endonuclease digestion, followed by electrophoresis on 5% polyacrylamide gels. Subclones were subsequently generated, on the basis of sequencing information, either through single or double digests with restriction endonucleases. The 17-base-pair (bp) oligonucleotide primer was purchased from the Regional DNA Synthesis Facility, University of Calgary. Amersham Corp. provided [γ-32P]ATP (3,000 Ci/mmol).
EB binding proteins (lane 7) but not with any polypeptides from the host E. coli strain harboring pUC8 (lane 6). Thus, pCT161/18 directs the production of a protein in E. coli which possesses the same eucaryotic cell-binding determinants as the native chlamydial binding protein. The presence of this overproduced protein on gels could be visualized with Coomassie blue stain (lane 2) but was absent with pUC8-derived products (lane 3). The chimeric plasmid pCT161/18 was transformed into the E. coli minicell strain P678-54. The recombinant plasmid encodes a protein of M₆ approximately 18,000, which was absent from minicells containing pUC8 (Fig. 2C). The recombinant gene product was also identified by using an E. coli-derived coupled transcription-translation system. Figure 2D shows the synthesis of an 18,000-M₆ polypeptide using pCT161/18, compared with pUC8, with which no similar polypeptide was visualized. Expression of chlamectin relative to that of β-lactamase was greater in an in vitro system (Fig. 2D) than in in vivo systems (Fig. 2A and C). Unlabeled proteins synthesized by the in vitro transcription-translation system bound to ¹²⁵I-labeled HeLa membranes on nitrocellulose blots. E. coli cells were fractionated as described above and separated on 12.5% polyacrylamide gels. This analysis revealed that most of the chlamectin was in the outer membrane fraction, as is the case with the native C. trachomatis binding protein (28).

Neutralization of chlamydial infectivity by antisera. Synthesis of chlamectin by cloned and subcloned DNA fragments was examined by immunoblot assay using hyperimmune rabbit antisera to the cloned 18,000-dalton protein (Fig. 1). This antisera was also used in neutralization assays. An inoculum of 3.5 × 10⁴ inclusion-forming units produced approximately 50% infection of HeLa cell monolayers. Preincubation of EB with antisera before inoculation was approximately 90% inhibitory, compared with 80% with antibodies to the native 18,000-dalton protein of EB. Normal rabbit serum also exerted a slight inhibitory effect (approximately 10%). Antiserum raised to whole EB react only weakly to the recombinant 18,000-dalton protein in immunoblots, compared with a strong reaction with native EB proteins as reported earlier (28).

Cellular localization of chlamectin. The cellular localization of chlamectin was examined in E. coli harboring pCT161/18 by the immunoblot technique using polyclonal antibodies to recombinant chlamectin, as well as by its binding characteristics to ¹²⁵I-labeled HeLa membranes on nitrocellulose blots. E. coli cells were fractionated as described above and separated on 12.5% polyacrylamide gels. This analysis revealed that most of the chlamectin was in the outer membrane fraction, as is the case with the native C. trachomatis binding protein (28).

Nucleotide sequence. The entire 1,658-bp PstI fragment of plasmid pCT161/18 was sequenced in both directions by using the dyeoxy method. The DNA sequence of the chlamectin gene and the deduced primary structure of the protein encoded by this gene are shown in Fig. 3. The complete C. trachomatis 18,000-dalton cell-binding protein was located on this 1,658-bp fragment. The proposed trans-
Amino acid sequence. The open reading frame of the recombinant gene was translated and, beginning at the initial methionine, comprises 162 amino acids, specifying a polypeptide of Mr 18,314. The derived chlamectin sequence contains highly charged residues toward the carboxy-terminal end. The presence of two short clusters of positively charged arginines—one between amino acids 146 and 148 and another between amino acids 156 and 159—is striking. The sequence contains two cysteine residues that are adjacent to each other. There is a short hydrophobic region toward the amino-terminal end which is preceded by two positively charged amino acids. The protein has a homogeneous distribution of charged, polar, and hydrophobic residues without clustering of hydrophobic groups, although chlamectin exhibits a 43% hydrophobicity overall.

Promoter studies. Sequences similar to the E. coli consensus promoter were not found upstream of the Shine-Dalgarno region. Sequence data demonstrated that the chlamectin gene in pCT161/18 is located in the same orientation as the lacZ promoter of pUC8. To test the dependence of the chlamydial gene on the lacZ promoter, the 650-bp PsrI-XbaI (nucleotides 1 to 650) fragment was subcloned into the PsrI-XbaI sites of pUC118 and pUC119. Expression of a biologically active gene product based on its binding characteristics to $^{125}$I-labeled HeLa membranes was obtained only with pUC119, where the open reading frame is in the same orientation as the lac promoter (Fig. 1).

**DISCUSSION**

The data presented here indicate that we isolated an E. coli recombinant that expresses an 18,000-dalton C. trachomatis binding protein identical in size and activity to a native C. trachomatis polypeptide. The deduced molecular size of the recombinant chlamectin (18,314) agrees closely with the published value of 18,000 (5, 28) for the native protein. $^{125}$I-labeled HeLa cells react with this recombinant polypeptide as well as they react with native EB binding proteins, reported earlier (5, 26, 28).

The ability of the cloned pCT161/18 plasmid to direct the synthesis of chlamydial protein was examined in E. coli minicells, as well as in an S-30 in vitro transcription-translation system. Polypeptides with similar molecular sizes were expressed in both systems, although the ratio of 18,000-dalton protein synthesis relative to that of $\beta$-lactamase was significantly higher in the in vitro system. Low levels of expression by minicells may be due to regulatory functions which are inoperative in an in vitro system (8). Further, the ability of in vitro products to bind to $^{125}$I-labeled HeLa cells may suggest either that the protein is fully active with normal structure and function, as has been observed with cloacin DF13 (8), or, if final conformation is not achieved, that the overall conformation of the protein is not crucial for eliciting its cell-binding properties.

The cellular localization of chlamectin on the outer membrane of E. coli and the location of the native binding protein on the outer membrane of C. trachomatis suggest the presence of a leader peptide (21). Most bacterial proteins inserted into the outer membrane have an amino-terminal extension with one or more basic residues, including one Arg or Lys (13). The presence of a hydrophobic region consisting of five amino acids preceded by a charged amino acid may represent a signal sequence although the characteristic hydrophobic core of 12 to 20 amino acids representing a typical signal sequence is lacking.

Antibodies directed against the recombinant 18,000-dalton protein exert a neutralizing effect in HeLa cell infection assays, presumably by binding to chlamectin, which then becomes unavailable for subsequent attachment to host cells. These findings agree with earlier results with polyclonal antisera to native chlamydial 18,000- and 31,000-dalton polypeptides (28). The weak cross-reactivity of recombinant chlamectin with polyclonal antibodies to whole EB may represent posttranslational modification in chlamydiae or alternate mechanisms for assembly of protein which are lacking in E. coli.

The entire PsrI fragment located on pCT161/18 was sequenced in both directions. The sequence revealed an open reading frame beginning at nucleotide 159 and extending up to nucleotide 644, comprising 162 amino acids. A sequence, AGGA, 10 positions upstream probably represents the ribosome-binding site (20). However, no consensus promoter sequences (~10 and ~35) similar to E. coli promoters were detected. The possibilities that the promoter is present far upstream of the start codon or that chlamydial promoters are unlike those of E. coli cannot be addressed without additional molecular and biochemical characterization. Recently, major outer membrane protein genes sequenced from both serovars L2 and L1 also failed to reveal consensus promoter sequences comparable to E. coli promoters (15a, 22). However, although chlamectin uses the vector promoter for its expression, a fused gene product was not produced, since stop codons were found in all three reading frames preceding the start codon. The absence of a functional promoter on the 650-bp PsrI-XbaI fragment was confirmed by the lack of expression of this insert in pUC118, where the lacZ promoter is oriented opposite to the inserted gene.

Hydropilicity has been used extensively to predict the antigenicity of various proteins and peptides. On the basis of the hydrophilic-algorithm model of Parker et al. (15), it was found that amino acids 20 to 30 and 125 to 162 are mostly surface accessible and hence antigenic. Whether any cell attachment role is played by these residues remains undis- determined. Moreover, the presence of a Gly-Asp-Ser-Val amino acid residue between amino acids 115 and 118 and Arg-Gly-Arg-Ser-Pro residues between 132 and 136, where four of five amino acids resemble the cell attachment domain of fibronectin (16), is notable. However, a computer-assisted comparison between chlamectin and E. coli revealed no significant homology. Interestingly, the presence of hydrophobic and homogenous distribution of polar, hydrophobic, and charged residues on chlamectin bear some similarities to those of the serovar L2 major outer membrane protein as reported recently by Stephens et al. (22).

The cloning and expression of the gene for the functional part of the chlamydial binding protein chlamectin in E. coli will facilitate further structural analysis. We should have sufficient quantities of the 18,000-dalton protein to study its structure-function properties, and especially its eucaryotic cell-binding ability, in greater detail. The deduced amino acid sequence of chlamectin, derived from nucleotide sequencing, will assist in identification of the domain respon-
sible for cell binding. This domain may be a candidate for future chlamydial vaccine development.

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LITERATURE CITED