

Transcription and Analysis of Polymorphism in a Cluster of Genes Encoding Surface-Associated Proteins of *Clostridium difficile*

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Recent investigations of the *Clostridium difficile* genome have revealed the presence of a cluster of 17 genes, 11 of which encode proteins with similar two-domain structures, likely to be surface-anchored proteins. Two of these genes have been proven to encode proteins involved in cell adherence: *slpA* encodes the precursor of the two proteins of the S-layer, P36 and P47, whereas *cwp66* encodes the Cwp66 adhesin. To gain further insight into the function of this cluster, we further focused on *slpA*, *cwp66*, and *cwp84*, the latter of which encodes a putative surface-associated protein with homology to numerous cysteine proteases. It displayed nonspecific proteolytic activity when expressed as a recombinant protein in *Escherichia coli*. Polymorphism of *cwp66* and *cwp84* genes was analyzed in 28 strains, and transcriptional organization of the three genes was explored by Northern blots. The *slpA* gene is strongly transcribed during the entire growth phase as a bicistronic transcript; *cwp66* is transcribed only in the early exponential growth phase as a polycistronic transcript encompassing the two contiguous genes upstream. The putative proteins encoded by the cotranscribed genes have no significant homology with known proteins but may have a role in adherence. No correlation could be established between sequence patterns of Cwp66 and Cwp84 and virulence of the strains. The *cwp84* gene is strongly transcribed as a monocistronic message. This feature, together with the highly conserved sequence pattern of *cwp84*, suggests a significant role in the physiopathology of *C. difficile* for the Cwp84 protease, potentially in the maturation of surface-associated adhesins encoded by the gene cluster.

Clostridium difficile, a gram-positive spore-forming anaerobic bacterium, is a significant nosocomial enteric pathogen, causing pseudomembranous colitis (PMC) and many cases of antibiotic-associated diarrhea (AAD) (29). The two main virulence factors are exotoxins, toxins A and B, both of which damage the human colonic mucosa and are potent tissue-damaging enzymes (3, 23). *C. difficile* takes advantage of the disturbance of the normal colonic flora, following antibiotic treatment, to colonize the gastrointestinal tract.

C. difficile has been shown to adhere in vitro to a variety of cultured cell lines, including Vero (37) and Caco-2 cells (9, 14). Interest for adherence determinants has recently risen, and numerous studies have characterized factors involved in the adherence and colonization processes, such as S-layer proteins (5, 6, 7), the adhesin Cwp66 (37), flagella (34), the heat-shock protein GroEL (17), and hydrolytic enzymes (28, 31).

C. difficile expresses on its surface an S-layer, which forms a regular two-dimensional array visible by electron microscopy (7). Each strain carries an S-layer, which is composed of two distinct proteins, one of high molecular weight called P47 and another of low molecular weight called P36. Both of these subunits are encoded by the *slpA* gene and are produced from the posttranslational cleavage of a precursor (6, 19). P36, encoded by the variable 5'-terminal part of the *slpA* gene, is

immunodominant, and its variability could play a role in the antigenic variation of the bacteria (7, 8, 19). P47 is encoded by the conserved 3'-terminal part of *slpA*, displays significant homology to the cell wall-anchoring domain of the autolysin CwlB of *Bacillus subtilis* (6, 7, 19), and shows strong and specific binding to gastrointestinal tissues and some extracellular matrix proteins (collagen I, thrombospondin, and vitronectin) (5). Recently the sequences of the variable region of the *slpA* gene were found to be strictly identical within a given serogroup (except for serogroup A) but divergent between serogroups (20).

Cwp66 (clostridial wall protein of 66 kDa) is a surface-associated protein with a two-domain structure. The N-terminal part of the protein presents homology to the cell wall-anchoring domain of the autolysin CwlB of *B. subtilis*; the C-terminal domain is cell surface exposed. Cwp66 has been shown to mediate adherence of *C. difficile* to Vero cells (37).

Since antibodies directed against P47 (5), P36 (19), and Cwp66 but also against the flagellar cap FliD (34) and GroEL (17) partially inhibit adherence of *C. difficile*, it seems very likely that the binding of *C. difficile* to host cells involves several proteins.

The genes encoding Cwp66 and the S-layer precursor are located close to each other in a 37-kb DNA fragment (Fig. 3 in reference 19) in the *C. difficile* genome (www.sanger.ac.uk/Projects/C_difficile/). This genetic locus carries 17 open reading frames (ORFs) in the same orientation, 11 of which encode proteins which present a two-domain structure, as described above: a domain homologous to the cell wall-anchoring do-

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TABLE 1. Strains of *C. difficile* used in this study

Category or strain	Toxinotype	Clinical origin	Geographic origin	GenBank accession no.	
				<i>cwp66</i>	<i>cwp84</i>
Serogroup A					
RefA = W1194					
ATCC 43594 ^a	A+B+	PMC ^b	Belgium		AY212121
68750	A+B+	PMC	France	AY211599	AY212122
Kohn	A-B-	Asymptomatic child	United Kingdom		AY212123
9354	A+B+	Adult AAD ^c (AIDS patient)	France		AY212124
Serogroup B					
RefB = 1351					
ATCC 43593 ^a	A-B-	Asymptomatic neonate	Belgium		AY212125
Ex560	A-B-	ND ^d	Belgium	AY211601	AY212126
CO109	A-B-	ND	France	AY211600	AY212127
Serogroup C					
RefC = 545					
ATCC 43596 ^a	A+B+	PMC	Belgium	AY211602	AY212128
C253	A+B+	AAD	Italy	AY211603	AY212129
630	A+B+	PMC	United Kingdom	AY211604	AY212130
1075	A-B-	ND	Belgium	AY211605	AY212131
Serogroup D					
RefD = 3232					
ATCC 43597 ^a	A-B-	Asymptomatic adult	Belgium		AY212132
93136	A-B-	Asymptomatic child	France		AY212133
Serogroup G					
RefG = 2022					
ATCC 43599 ^a	A+B+	Asymptomatic neonate	Belgium		AY212134
95938	A+B+	PMC (relapse)	France	AY211606	AY212135
Serogroup H					
RefH = 2149					
ATCC 43600 ^a	A+B+	PMC	Belgium		AY212136
93369	A+B+	Asymptomatic adult	France		AY212137
53444	A-B-	ND	ND		AY212138
90204	A+B+	Asymptomatic child	France		AY212139
Serogroup I					
RefI = 7322					
ATCC 43601 ^a	A-B-	Asymptomatic neonate	Belgium		AY212140
56026	A-B-	Asymptomatic child	Belgium	AY211607	AY212141
Serogroup K					
RefK = 4811					
ATCC 43602 ^a	A-B-	Asymptomatic neonate	Belgium	AY211608	AY212142
94416	A+B+	PMC (relapse)	France	AY211609	AY212143
Serogroup X					
RefX = 5036					
ATCC 43603 ^a	A-B-	Asymptomatic neonate	Belgium	AY211610	AY212144
36678	A-B+	ND	Belgium	AY211611	AY212145
Diverse					
79685	A+B+	PMC	France	AY211612	AY212146
57207	A-B+	ND	ND		AY212147
CD 268	A+B+	Isolated from rabbit	Italy		AY212148

^a Reference strain for the serogroup (Prof. Delmée, Université catholique de Louvain, Belgium).

^b PMC, pseudomembranous colitis.

^c AAD, antibiotic-associated diarrhea.

^d ND, not determined.

main of CwlB, present in either the N-terminal or the C-terminal part, and a second domain (named the functional domain) displaying remote homologies with different enzymes or structural proteins from gram-positive bacteria (4, 6, 19).

The aim of this study was to characterize this putative virulence cluster by focusing on three genes: *slpA*, *cwp66*, and *cwp84*. The *cwp84* gene, located immediately downstream from *cwp66*, encodes a putative 84-kDa protein with a characteristic signal peptide, whose functional domain displays homologies with several cysteine proteases, and shows the conserved Pept_C1 domain of the papain family. Since extracellular proteases have been described to be virulence factors in many

bacteria (22), we cloned and expressed the corresponding gene to permit functional studies. We studied the transcription of *slpA*, *cwp66*, and *cwp84* genes by Northern blotting to determine whether these genes are expressed as part of an operon. We also investigated the polymorphism of *cwp66* and *cwp84* genes in different strains of *C. difficile* in an attempt to see if a correlation between pattern of sequences and virulence exists.

MATERIALS AND METHODS

Bacterial strains and growth conditions. Twenty-eight *C. difficile* isolates belonging to nine different serogroups were studied (Table 1). The strains include the nine reference strains for specific serogroups (11), clinical isolates from

TABLE 2. Primers used in this study

Purpose, relevant gene, or primer name	Oligonucleotide sequence (5' → 3')	Position (bp)
Probe generation		
<i>slpA</i>		
slpA-NB1	GCT GCT CCT GTT TTT GCT	+54/+71
slpA-NBR1	CTT CTT TTG CAT TTA TAA C	+828/+810
<i>cwp66</i>		
cwp66-R1	AAT CCA TCA TCT GTA GCG	+1818/+1801
cwp66-NB1	CTC AAA TTG GTG GCT TAG G	+915/+933
cwp66-NBR1	ATG GCT CTT CAT CTG TTG G	+1712/+1694
<i>cwp84</i>		
cwp84-NB1	CTC TAG ATG GAG TAG AAA CT	+104/+123
cwp84-NBR1	GAC CAG CAT ATT CAA GTT G	+1030/+1012
<i>gdh</i>		
gdh-NB1	GAT GTA AAT GTC TTC GAG ATG	+13/+33
gdh-NBR1	GGT CCA TTA GCA GCC TCA C	+974/+956
RT-PCR		
<i>slpA</i>		
orf6-RT1	AGA TGG ACA AGT GTA TGC	+1545/+1562
slpA-RTR1	CAG TCG TTT TTA ACT ACA G	+115/+97
slpA-RT1	ATG GTG GAA CTA ACT TAG	+2077/+2094
orf7-RTR1	TTG CTC ATC TGC TTT GTC	+41/+24
<i>cwp66</i>		
orf8-RT1	AGT TGA ATT GAC AGT AAT AC	+1824/+1843
orf9-RTR1	CTG TGC ATA ATA TGA CAT GT	+60/+41
orf9-RT1	GCT ATA GGA TAT CAT TCA G	+576/+594
slpA-RTR1	GAT AAA GCA TCT GCT ATG G	+195/+177
Amplification of functional gene domains		
<i>cwp84</i>		
orfE-ATG	GGG GTA AAC ATG AGA AAG	-9/+9
cwp84-TAA	GGA ACT CCA TTT ACT ACT G	+1178/+1150
<i>cwp66</i>		
cwp66-1	AGC AGT GGG TGT ATT AGC	+805/+822
cwp66-R1	AAT CCA TCA TCT GTA GCG	+1818/+1801
cwp66-NB1	CTC AAA TTG GTG GCT TAG G	+915/+933
cwp66-NBR1	ATG GCT CTT CAT CTG TTG G	+1712/+1694
cwp66-ATG	CGA AAG AAT TAG GAG GTA AGA	-35/-15
cwp66-TAA	TAT GTA TGT AAT GAT TGA TTT GC	+1992/+1970

PMC, AAD, or asymptomatic carriers, and one strain of animal origin. Clostridial strains were grown under anaerobic conditions in TGY (tryptone glucose yeast infusion broth; Difco Laboratories) and on Columbia agar plates supplemented with 4% horse blood (Biomerieux).

The BL21 *Escherichia coli* strain, used as a host for *cwp84* cloning, was grown on Luria broth or brain heart infusion (BHI) agar or in broth (Difco Laboratories), supplemented with 100 µg of ampicillin/ml to maintain the pGEX plasmid.

Serratia sp. and *Streptococcus pyogenes* strains, used as positive controls in proteolytic assays, were grown on BHI agar or in broth, at 37°C, under aerobic conditions.

Typing of strains. Typing of strains belonging to serogroup C was done by PCR ribotyping and randomly amplified polymorphic DNA (RAPD) analysis with primers AP3, AP4 (2), and APRB11 (10), as described previously.

Cloning of *cwp84* into the vector pGEX-6P-1 and protein expression. To clone the *cwp84* gene into the pGEX-6P-1 expression vector (Amersham Biosciences), two oligonucleotide primers, pGEXcwp84-*EcoRI* (5'GGGTAGAATTCAGAAAGTATAAATCA3') and pGEXcwp84-*XhoI* (5'TCTCTCGAGTCACTATTTTCTAAAAG3'), incorporating an *EcoRI* and *XhoI* site, respectively (und erlined), were used to amplify by PCR the full-length coding region of the *cwp84* gene of the 79685 strain. The resulting 2.4-kb DNA fragment was digested with the two enzymes and ligated (1 U of T4 ligase; Invitrogen) between the *EcoRI* and *XhoI* sites of pGEX-6P-1. Sequencing of the insert was done, with primers pGEX-3' and pGEX-5' (Amersham Biosciences) and internal primers, to ensure that no sequence mismatch occurred during the cloning. The plasmid carrying an in-frame fusion between *gst* and *cwp84* was transformed into *E. coli* BL21 (Amersham Biosciences). Subsequent protein expression and purification steps were performed by induction of the *tac* promoter with 0.1 mM isopropyl-β-D-T

hiogalactopyranoside (IPTG), following by a single-step affinity chromatography employing glutathione-Sepharose-4B, as described in protocols from Amersham Biosciences. Purification attempts were done with or without cysteine protease inhibitors, such as leupeptin and E64 (Sigma). Before cleavage from the glutathione *S*-transferase (GST), purified fractions were tested for the presence of recombinant protein by immunoblotting, as previously described (37), with antibodies directed against GST (Amersham Biosciences) and against the N-terminal part of Cwp66, which is 56% homologous to the C-terminal part of Cwp84, as shown by amino acid sequence alignment.

Proteolytic assays. Proteolytic assays were done with the clone *E. coli* BL21(pGEXΩ*cwp84*) and the strain without insert BL21(pGEX) as a negative control. *Serratia* sp. and *S. pyogenes* were used as positive controls for gelatinase and cysteine protease activity, respectively. Assays were performed on gelatin, skimmed milk, and azocoll. To test for gelatinase activity, clones were grown aerobically for 24 h in BHI broth, and then 0.1 mM IPTG and a photographic film containing gelatin and charcoal were added, and cultures were continued for 24 h under anaerobic conditions. Gelatinase activity was assessed by release of charcoal in the culture medium after lysis of gelatin. Caseinase activity was determined on skimmed milk supplemented or not with 0.1 mM IPTG by the measure of the clear halo around the streak culture. Azocoll degradation was determined by measuring optical density at 520 nm (OD₅₂₀) after 48 h of culture following by 6 h of contact with azocoll, as previously described (28).

RNA manipulations. Total RNA extraction was performed at the beginning (OD₆₀₀, ~0.3) and the middle (OD₆₀₀, ~0.7) of the exponential growth phase and during the stationary phase from *C. difficile* strain 630 (18-h culture) according to the Trizol (Invitrogen) procedure previously described (13), with some modifications. Briefly, after bacterial lysis in prelysis buffer containing 50 mM

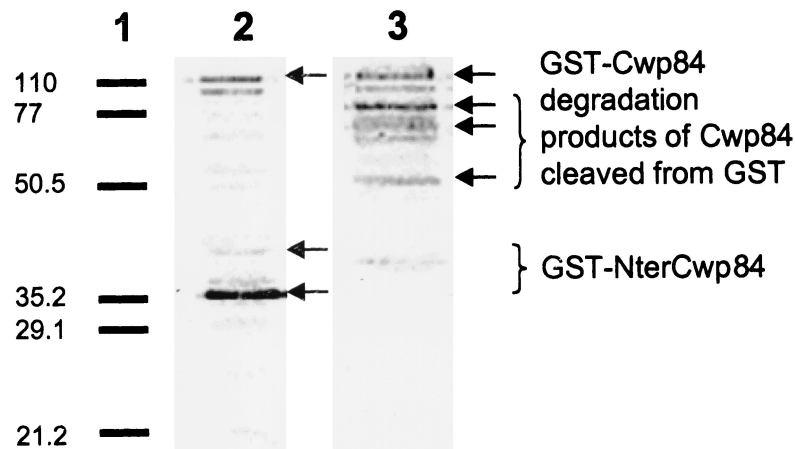


FIG. 1. Immunoblot analysis of purified fraction GST-Cwp84. Equivalent amounts of the purified fraction were separated by SDS-12% polyacrylamide gel electrophoresis, transferred to a Hybond-P membrane (Amersham Biosciences), and incubated either with anti-GST antibody (lane 2) or anti-Cwp66Nter (lane 3). Revelation was done with phosphatase alkaline-conjugate antigoat antibodies or phosphatase alkaline-conjugate antirabbit antibodies, respectively, with nitroblue tetrazolium-5-bromo-4-chloro-3-indolylphosphate (Invitrogen). Lane 1, low-range SDS-polyacrylamide gel electrophoresis standard (Bio-Rad). Size markers are given in kilodaltons on the left.

Tris (pH 8.0), 25% sucrose, and 10 mg of lysozyme/ml and RNA extraction with trizol, RNA was precipitated in absolute ethanol and stored until utilization at -20°C . Before every manipulation, an aliquot was removed, washed with 70% ethanol, and dissolved in RNase-free water at 60°C . The RNA concentration was measured optically at 260 nm. Fifteen to thirty micrograms of RNA was electrophoresed in 20 mM guanidine thiocyanate (15; I. Podglajen, personal communication)-1.8% agarose gel in $1\times$ Tris-borate-EDTA at 65 V for 4 to 5 h in a horizontal gel electrophoresis apparatus. RNA was then transferred directly to Nylon membrane Hybond-N+ (Amersham Biosciences) by the downflow capillary method.

(i) **Generation of probes.** PCR products obtained with primers described in Table 2 were purified and labeled using the ECL direct labeling and detection system (Amersham Biosciences), according to the manufacturer's instructions.

(ii) **Northern blot.** Membranes were prehybridized for 1 h at 42°C in hybridization buffer (Amersham Biosciences) prior to the addition of probe. Hybridizations were performed overnight at 42°C and followed by two washes (20 min each) in 6 M urea-0.4% sodium dodecyl sulfate (SDS)-0.5 \times , 0.2 \times , or 0.1 \times SSC (1 \times SSC is 0.15 M NaCl plus 0.015 M sodium citrate), depending on the probe used, and two washes in 2 \times SSC (10 min each). Detection was carried out with the ECL direct labeling and detection system (Amersham Biosciences). A probe for the glutamate-dehydrogenase *gdh* gene of *C. difficile* was generated for use as a positive control for the hybridization experiments, as previously described (16).

RT-PCR. RNA were treated by RNase free-DNase (Amersham Biosciences). Reverse transcription (RT)-PCR was performed using the SuperscriptOne-Step RT-PCR with the *Platinum Taq* kit (Invitrogen), as recommended by the manufacturer, with 100 ng of RNA. Primers used were those designed to generate the probes for Northern blots and some others chosen to check the polycistronic nature of mRNA (Table 2). Simultaneously, PCR was performed on samples with the same oligonucleotides to exclude false-positive amplification from residual DNA.

Analysis of cotranscribed ORFs. Homologies of cotranscribed ORFs with known proteins were searched with BLAST (www.ncbi.nlm.nih.gov/BLAST). Particular motifs and secondary structures were analyzed with the PROSITE and SOPMA programs (www.pbil.univ-lyon1.fr/).

PCR amplifications. DNA from the 28 isolates was extracted according to the protocol provided in the DNeasy tissue kit (Qiagen). The primers used for amplification of the functional regions of the *cwp66* and *cwp84* genes (corresponding to their C-terminal and N-terminal domains, respectively [Fig. 3 in reference 19]), are listed in Table 2. PCR amplification was performed in a Perkin-Elmer GeneAmp PCR System 2400 thermocycler, with one bead of Ready-To-Go PCR kit (Amersham Biosciences), each primer at a final concentration of 0.4 μM , and 100 ng of genomic DNA (in a 25- μl reaction mixture). Initial denaturation was carried out at 95°C for 5 min, following by 35 cycles of amplification: denaturation at 95°C (1 min), annealing at 50 or 55°C (depending on primers used) for 1 min, and extension at 72°C for 1 min, 1 min 30 s, or 2 min.

An additional step of extension for 10 min at 72°C was performed at the end of the amplification. Samples (3 μl) of amplified products were analyzed by electrophoresis in a 1.0% (wt/vol) agarose gel.

***cwp66* and *cwp84* sequencing.** PCR products with the expected size were purified with the High Pure PCR purification kit (Roche). Automatic DNA sequencing was performed on the two strands with the Big Dye Terminator cycle-sequencing kit (Applied Biosystems) and analyzed with an ABI PRISM 310 genetic analyzer (Perkin-Elmer). Initial sequencing was carried out with the same primers as used for PCR, and more sequence was acquired by the DNA walking strategy.

Nucleotide and protein sequence alignments were performed with the DNA CLUSTAL W program (www.ebi.ac.uk/clustalw/).

Nucleotide sequence accession numbers. The GenBank nucleotide sequence accession numbers of the functional domain of the genes *cwp66* and *cwp84* from the *C. difficile* strains studied are given in Table 1.

RESULTS

Expression of Cwp84 and proteolytic assays. In an attempt to produce a recombinant protein corresponding to the *cwp84* ORF in the *cwp* gene cluster of *C. difficile* (4, 6, 19), encoding a potential protease, the *cwp84* gene of the strain 79685 was expressed as an in-frame fusion with GST in *E. coli*. The *E. coli* clone pGEX Ω *cwp84* presented nonspecific proteolytic activity on gelatin and skimmed milk compared to the negative control (data not shown). A light degradation of azocoll could be observed for this clone, as measured by the mean of OD_{520} of three distinct experiments: OD_{520} for *E. coli* pGEX Ω *cwp84* = 0.074; OD_{520} for *E. coli* pGEX = 0.022; OD_{520} for *Serratia* sp. = 0.379; OD_{520} for *S. pyogenes* = 0.109.

Purification of the recombinant protein was carried out with 0.1 mM IPTG induction, since use of 0.5 and 1.0 mM concentrations led to formation of inclusion bodies. Immunoblot analysis of purified fractions from various assays showed that the two antibodies used (anti-GST and anti-Cwp66 N-ter) revealed one very faint band of 110 kDa, the size of which corresponds to that of the fusion protein GST-Cwp84 (Fig. 1). Numerous other bands were recognized alternatively with the two antibodies. These bands, especially those specifically recognized by anti-Cwp66 N-ter, probably correspond to degradation products of Cwp84. A very tight band of approximately

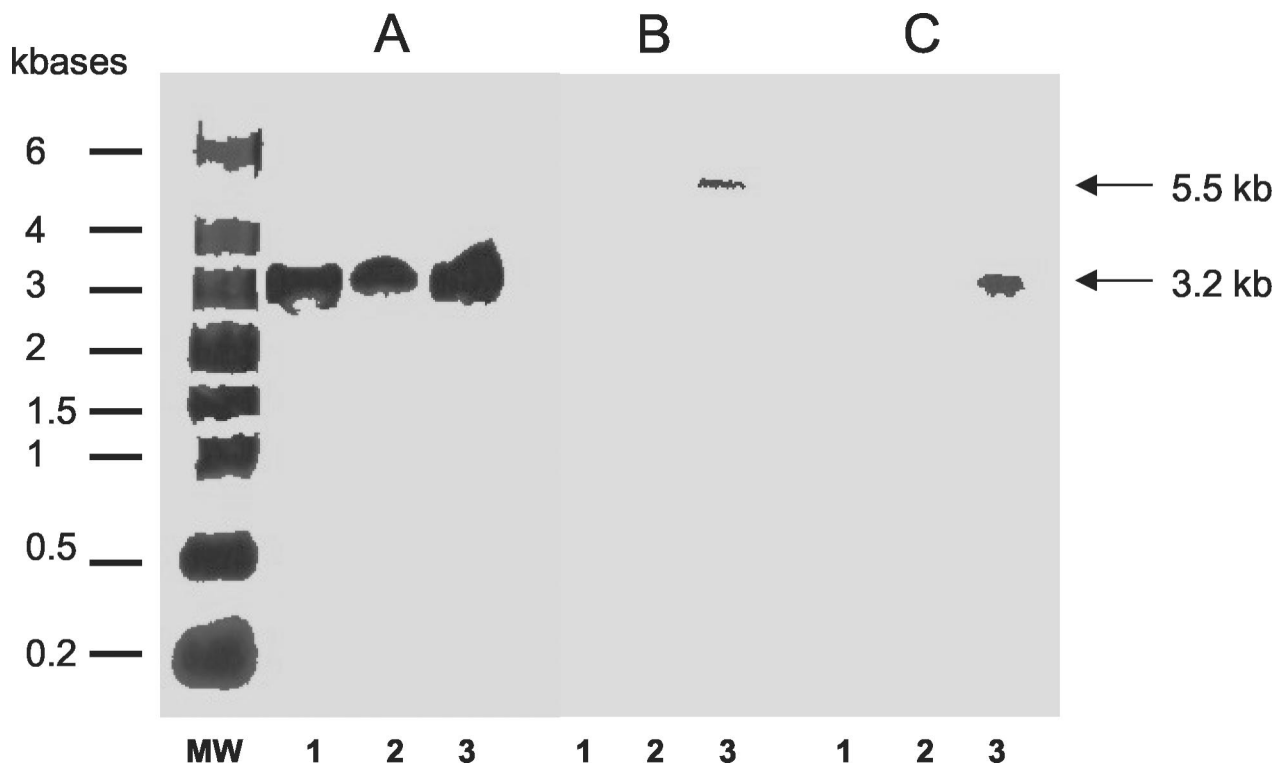


FIG. 2. Transcriptional analysis of *cwp* cluster. Hybridization DNA–membrane-immobilized RNA on strain 630; 15, 30, and 25 μ g of total RNA were used for hybridization with probes specific to *slpA* (A), *cwp66* (B), and *cwp84* (C), respectively. MW, RNA molecular weight marker (Sigma); lanes 1A, 1B, and 1C, RNA from stationary growth phase (18-h culture); lanes 2A, 2B, and 2C, RNA from the middle of the exponential growth phase (OD_{600} of ~ 0.7); lanes 3A, 3B, and 3C: RNA from the beginning of the exponential growth phase (OD_{600} of ~ 0.3). Sizes of transcripts are indicated with arrows. The probe specific for the transcript of the *gdh* gene hybridized with an estimated 1.3-kb transcript in the sample corresponding to the different growth phase, as expected (16) (data not shown).

35 kDa was detected only by anti-GST, which may correspond to the GST (29 kDa) bound to a small N-terminal part of Cwp84. Cysteine protease inhibitors used did not prevent degradation of the protein of interest. Purified fractions did not have proteolytic activity on nonspecific substrates, and various modifications in purification conditions were not more successful in inducing this activity.

Transcriptional analysis of the *cwp* cluster. The organization and structure of the genes in the *cwp* cluster suggest that some of the genes could be cotranscribed and form an operon.

Therefore, the transcription of several genes of the cluster was investigated by Northern blotting and RT-PCR.

Hybridizations with *slpA*-specific probe are shown in Fig. 2A. Transcripts of *slpA* were detected during all phases of growth and were estimated to be 3.2 kb, whereas size of the *slpA* gene is only 2,160 bp. RT-PCR with primers encompassing intergenic regions between *orf6* and *slpA* and between *slpA* and *orf7* was performed (Fig. 3). Amplifications were positive with intergenic primers encompassing *orf6* and *slpA*, indicating the bicistronic feature of the *slpA* transcript (Fig. 4A).

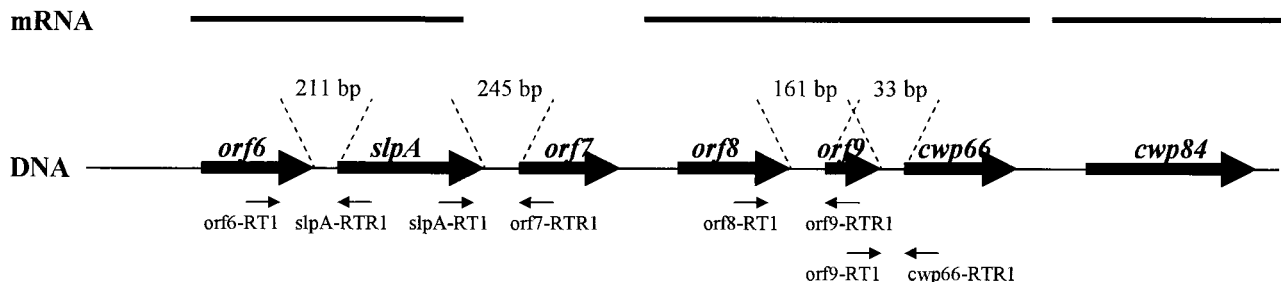


FIG. 3. Genetic organization of the DNA cluster carrying the *slpA*, *cwp66*, and *cwp84* genes from the genome of *C. difficile* 630. The thick horizontal arrows represent the open reading frames with their names indicated above. Thick lines above represent transcripts that are monocistronic (for *cwp84*) or polycistronic (for *orf6-slpA* and *orf8-orf9-cwp66*). The sizes of intergenic regions, determined from analysis of the available genome sequence of strain 630, are indicated between the dotted lines. Thin arrows below DNA indicate the localization of primers used in intergenic RT-PCR.

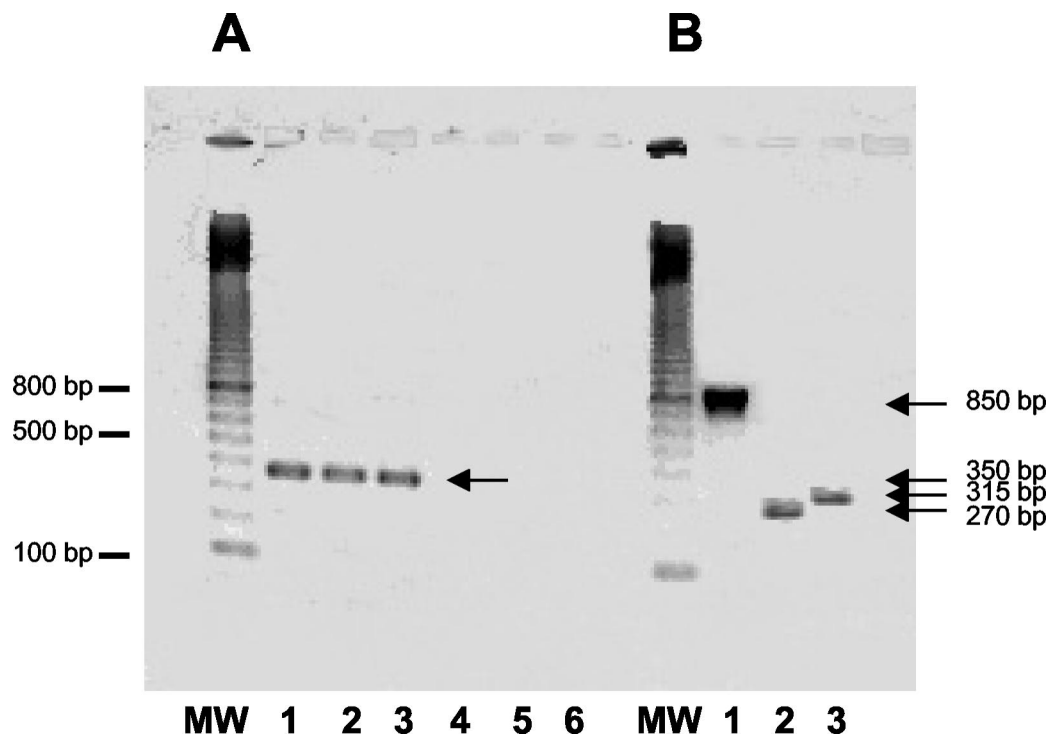


FIG. 4. RT-PCR analysis of the genes located upstream from *slpA* (A) and *cwp66* (B) in strain 630. (A) MW, molecular weight marker (100-bp ladder; Amersham Biosciences). Lanes 1 to 3, amplification between *orf6* and *slpA* with *orf6*-RT1/*slpA*-RTR1 primers, demonstrating the presence of an intergenic mRNA. Lanes 4 to 6, no amplification was obtained with the primers *slpA*-RT1/*orf7*-RTR1, showing absence of intergenic RNA messenger. RNA was extracted from the stationary phase (lanes 1 and 4), middle exponential growth phase (lanes 2 and 5), and beginning of the exponential growth phase (lanes 3 and 6). (B) MW, molecular weight marker (100 bp ladder). Lanes 1 to 3, RNA extracted from the beginning of the exponential growth phase. Lane 1, *cwp66*-NB1/*cwp66*NBR1 primers; lane 2, *orf8*-RT1/*orf9*-RTR1 primers; lane 3, *orf9*-RT1/*cwp66*-RTR1 primers.

Hybridizations with a probe specific for the *cwp66* gene functional domain showed the presence of a transcript only at the beginning of the exponential growth phase (Fig. 2B). The size of the transcript was estimated to 5.5 kb, whereas the *cwp66* gene size is only 1.8 kb. However, this size is compatible with a polycistronic transcript encompassing the two genes immediately upstream of *cwp66* in the sequenced genome of strain 630 (Fig. 3). Primers encompassing regions between *orf8* and *orf9* and between *orf9* and *cwp66* gave positive results in RT-PCR, confirming the occurrence of a polycistronic transcript *orf8-orf9-cwp66* (Fig. 4B).

Hybridizations with a probe specific for *cwp84* showed the presence of a prominent transcript in the early exponential growth phase (Fig. 2C). The size was estimated to be 3.2 kb, corresponding to a monocistronic transcript. The presence of this transcript was confirmed by RT-PCR (data not shown).

Analysis of the putative Orf6, Orf8, and Orf9. The deduced peptidic sequence of *orf6* shows a 528-amino acid protein with a two-domain structure similar to that described for the other proteins encoded by the genes of the cluster. The cell wall-anchoring domain, located on the N-terminal part, has 53% homology with the cell wall-anchoring domain of CwlB. The predicted secondary structure of the C-terminal part is characterized by a mostly α -helical conformation without any particular motif and does not show any homology with known proteins. The deduced peptidic sequence of *orf8* shows again a two-domain structure. The cell wall-anchoring domain, 52%

homologous to the autolysin CwlB of *B. subtilis*, is located in the C-terminal part. The predicted secondary structure of the functional domain shows a strikingly α -helical conformation (71%) with a putative transmembrane segment but shares no significant homology with known proteins. Orf9 does not possess the two-domain structure. Its predicted secondary structure is also rich in α -helices (44%). The deduced protein has no significant homology with known proteins.

Polymorphism of *cwp84* and *cwp66* genes. The functional region of the *cwp84* gene was easily amplified and sequenced from the 28 strains studied. This domain shows a high degree of conservation, since there were only 51 polymorphic nucleotide sites over 1,043 nucleotides sequenced, including 14 that would result in amino acid replacements (Fig. 5). The nucleotide sequences were 100% identical within each serogroup of C, D, I, K, and X. In contrast, the four strains belonging to serogroup A each had a unique *cwp84* functional region sequence, as previously observed for the S-layer precursor *slpA* (20).

Amplification of the C-terminal functional domain of the *cwp66* gene was successful only for 14 strains (RefC, 630, C253, 1075, CO109, Ex560, 68750, 56026, 95938, RefK, 94416, RefX, 36678, and 79685), and sequencing revealed numerous nucleotide changes, which resulted in nine distinct deduced peptidic sequences. The deduced peptidic sequences were 100% identical within serogroup C (four strains), serogroup X (two strains), and two strains of serogroup B (RefB could not be

	17							
RefC	TVCFLIVSTI	PVSAENHKT	DGVETA EYSE	SYLQYLEDVK	NGDTAKYNGV	IPFPHEMEGT		60
RefH	-----	-----	-----	-----	-----	-----		
RefG	-----	-----	-----	-----	-----	-----		
Ex560(B)	-----	-----	-----	-----	-----	-----		
RefA	--Y-----	-----	-----	-----	-----	-----		
RefD	-----	-----K---	-----	-----	-----	-----		
9354(A)	-----	-----N---	-----	-----	-----	-----		
				<i>Cysteine 116</i>				
RefC	TLRNKGRSSL	PSAYKSSVAY	NPMDLGLTTP	ARNQGSLN <u>T</u> C	WSFSGMSTLE	AYLKLKGYGT		120
RefH	-----	-----	---N---	-----	-----	-----		
RefG	-----	-----	-----	-----	-----	-----		
Ex560(B)	-----	-----	---N---	-----	-----	-----		
RefA	-----	-----	---N---	-----	-----	-----		
RefD	-----	-----	---N---	-----	-----	-----		
9354(A)	---S-S---	--S-----	---N---	-----	-----	-----		
RefC	YDLSEELRW	WATGGKYGWN	LDDMSGSSNV	TAIGYLTAWA	GPKLEKDIPY	NLKSEAQGAT		180
RefH	-----	-----	-----	-----	-----	-----		
RefG	-----	-----	-----	-----	-----	-----		
Ex560(B)	-----	-----	-----	-----	-----	-----		
RefA	-----	-----	-----	-----	-----	-----		
RefD	-----	-----	-----	-----S-	-----	-----		
9354(A)	-----	-----	-----	-----	-----	-----		
RefC	KPSNMDTAPT	QFNVTDVVRL	NKDKETVKNA	IMQYGSVTSG	YAHYSTYFNK	DETAYNCTNK		240
RefH	-----	-----	-----	-----	-----	-----		
RefG	E-----	-----	-----	-----	-----	-----		
Ex560(B)	-----	-----	-----	-----	-----	-----		
RefA	-----	-----	-----	-----	-----	-----		
RefD	-----	-----	-----	-----	-----	-----Y-		
9354(A)	R-----	-----	-----	-----	-----	-----		
	<i>Histidine 262</i>			<i>Asparagine 287</i>				
RefC	RAPLN <u>H</u> AVA	IVGWDDNYSKD	NFASDVKPES	<u>N</u> GAWLVKSSW	GEFNSMKGFF	WISYEDKTL		300
RefH	-----	-----	-----	-----	-----	-----		
RefG	-----	-----	-----	-----	-----	-----		
Ex560(B)	-----	---N---	-----	-----	-----	-----		
RefA	-----	-----	-----	-----	-----	-----		
RefD	--T-----	-----	---A---	-----	-----	-----		
9354(A)	-----S-	-----	-----	-----	-----	-----		
						370		
RefC	TDTDNYAMK	SVSKPDSDKM	YQLEYAGLSK	IMSNKVTAAN	VDFFSRDSEK	LDSV		354
RefH	-----	-----	-----	-----	-----	-----		
RefG	-----	-----	-----	-----	-----	-----		
Ex560(B)	-----	-----	-----	-----	-----	-----		
RefA	-----	-----	-----	-----	-----	-----		
RefD	-----	-----	-----	-----	-----	-----		
9354(A)	-----	-----	-----	-----	-----	-----		

FIG. 5. Polymorphism of the functional domain of Cwp84, from amino acid 17 to 370. The seven sequences represents the different alleles recovered from the 28 strains studied. Only amino acid differences from serogroup C reference strain (ATCC 43596) are indicated. Identical amino acids are represented by dashes. RefC, same sequence than following strains (serogroup): 630(C), C253(C), 1075(C), Kohn(A), 95938(G), 53444(H), 57027, RefI, 56026(I), RefK, 94416(K), RefX, 36678(X). RefH, same sequence as the following: 93369(H), 90204(H), 68750(A), and 79685(S3). RefG, same sequence as strain CD268. Ex560(B), same sequence as CO109(B). RefA, same sequence as RefB. RefD, same sequence as 93136(D).

amplified). The genes sequenced from strains 56026 (serogroup I) and 95938 (serogroup G) were highly divergent from the strain 630 *cwp66* gene. It is noteworthy that the Cwp66 sequence of serogroup C seems to diverge significantly from

the other patterns of sequence: percentages of homology for serogroup C are of 61% with serogroup B strains, 62% with serogroup X strains, 59% with serogroup K strains, and 50% with strain 95938 and 56026. On the contrary, there is a high

	286										
630(C)	VLAGNKLDTT	QKDVLNTKI	DKVTQIGGLG	NENVVEDILD	IQEETKYTVE	TIDELNAAIK					60
Ex560(B)	--A-I	-----	-----	--EA-KS-I-	M--K-----	-----V---					
79685(S3)	I-----	-----	-----	--DA-KS-V-	M--K-----	--E---V---					
36678(X)	-----A	-----V-	-----	--DA-KS-V-	M--K-----	--E---V---					
630(C)	RADANDIIKF	KPEKEKTINN	SFSIETKKTV	TIELDGRYRQ	TITLDIPNGK	FNNYAEIEGG					120
Ex560(B)	K-----V-I-	E---DTN-SD	--K-A-N-AI	-V-F--VFKK	S--I-M---D	VK-FG--SDD					
79685(S3)	K-----V-I-	E---DTN-SD	--K-A-N-AI	-V-F--VFK-	S--I-M---D	VK-FG--SDD					
36678(X)	K-----V-I-	E---DTN-SD	--K-A-N-AI	-V-F--VFK-	S--I-M---D	VK-FG--SDD					
630(C)	VKLKNIKNES	LVNKGSIQDL	DIYDENGCKI	ENESSGEIWF	VTIVEEANDV	YIVNSGDITK					180
Ex560(B)	MRID---KGT	-I-E---GI	---SK-----	--TND-D--I	I--DAD-K--	--E-D-----					
79685(S3)	IRID---KGT	-I-E---GI	---SK-----	--T-D-D--I	I--DAD-K--	--E-D-----					
36678(X)	IRID---KGT	-I-E---GI	---SK-----	--T-D-D--I	I--DAD-K--	--E-D-----					
RefK		-E---GI	---SK-----	--K-D-D--L	I---S--RG-	----D---S-					
630(C)	ISNNSSTII	RNSGNIDTVT	GKKEPAISGN	KPKVNDTEKE	TKAARGLNPR	VEACSVPKKD					240
Ex560(B)	----APGV--	K---K--L-N	-NEQ-----K	--TT----YN	DER----SVS	TKP--I-E-N					
79685(S3)	----APGV--	K---K--L-N	-NEQ-----K	--TT----YN	DER----SVS	TKP--I-E-N					
36678(X)	----APGV--	K---K--L-N	-NEQ-----K	--TT----YN	DER----SVS	TKP--I-E-N					
RefK	---SC-DV--	K---K-NL--	-TE-----K	--IT----YD	DER-H--SVK	T----T-Q-N					
630(C)	YVMITIPNSP	KDSRYKIYYR	VVYNKPYAMD	VGDKINIGEW	TVAPTDEEPF	LEKAKNGCYV					300
Ex560(B)	R-RV--SSE-	-S-----	--ED--S--Y	--E--SVRN-	DL-SKSDGS-	V-----S-I					
79685(S3)	R-RV--SSE-	-S-----	--ED--S--Y	--E--SVRS-	DL-SKSDGS-	V-----S-I					
36678(X)	R-RV--SSE-	-S-----	--ED--S--Y	--E--SVRS-	EL-SKSDGS-	V-----S-I					
RefK	-IIV--SSK-	-N-N-A----	--GD--S--Y	--E---PRD-	YSVSKSDDS-	I-----S-I					
			610								
630(C)	EAVEVNTSTK	EVSRWGRINA	TDDGF								325
Ex560(B)	-V--I----N	K-----S-V	-----								
79685(S3)	-V--I----N	K-----S-V	-----								
36678(X)	-V--I----N	K-----S-V	-----								
RefK	-V--I---NN	R-----SSS	----L								

FIG. 6. Polymorphism of the functional domain of Cwp66, from amino acids 286 to 610. The five sequences represents the various alleles recovered from 12 strains studied (the sequences of strains 56026 and 95938 are too divergent to appear in this alignment). Only differences from strain 630 (serogroup C) are indicated. Identical amino acids are represented by dashes.

degree of homology (>90%) among serogroups B, X, and K (Fig. 6).

Typing of strains. Typing of strains belonging to serogroup C was done to check for the nonclonality of these strains, since the sequences of *cwp84* and *cwp66* were perfectly identical among the four strains of this serogroup. PCR ribotyping generated only two profiles, one for the three toxigenic strains and one corresponding to the nontoxigenic strain. In contrast, patterns of RAPD obtained with three different primers readily differentiated the three toxigenic strains (data not shown). A higher level of discrimination for RAPD compared to PCR ribotyping has already been described for toxigenic strains belonging to serogroup C (2, 36).

DISCUSSION

The cluster carrying genes studied in this study was previously characterized (6, 19). Two of these genes encode well-characterized surface-exposed proteins involved in adherence of *C. difficile* to host cells: adhesin Cwp66 (37) and S-layer proteins (5, 7). The transcription of some of these genes has been recently studied by RT-PCR (6), but no information on their transcriptional organization was available in the beginning of our study. So, the first aim of our study was to analyze transcription of the three genes, *slpA*, *cwp66*, and *cwp84*, to investigate if an organization in operon exists in this cluster as has been described for the toxigenic element of *C. difficile* (16).

Transcriptional analysis of *slpA*, encoding the precursor of the S-layer proteins, revealed that this gene is strongly transcribed during the whole growth phase, reflecting the fact that S-layer proteins are the major surface proteins (7, 8). More surprisingly, the putative 57.5-kDa protein encoded by *orf6* may also be strongly expressed. This protein may be of importance for the bacterium, but no putative role could be assessed in view of its structure and homologies. DNA sequence analysis showed that a ribosome binding site exists just upstream from *orf6* (AGGAGG) and also sequences which could be part of a promoter: TATAAA (−10) and TTTTAG (−35). Intriguingly, we also found putative promoter sequences immediately upstream of *slpA*. As has been demonstrated for toxins A and B (13, 16), a monocistronic transcript of *slpA* could also exist in conditions other than those used in this work, increasing production of the S-layer precursor, which could be important for the adaptation of *C. difficile* to its environment.

The *cwp66* gene is transcribed only in the early exponential phase, at a low level, as a polycistronic transcript encompassing three contiguous *orf* genes in the following order: *orf8-orf9-cwp66* (Fig. 3). Cotranscription of *orf9* and *cwp66* is not surprising because the intergenic region is short (<50 bp) and there is no putative promoter upstream from *cwp66*. *orf8* and *orf9* are each preceded by likely promoter consensus sequences at −10 and −35. We are unable to explain this simultaneous transcription, since the putative proteins encoded by *orf8* and *orf9* do not exhibit any homology to known proteins. Nevertheless, one hypothesis could be that the two or three proteins could associate *in vivo* to form a complexed adhesin, as has been already described for *Porphyromonas gingivalis* (27). In this bacterium, the associated proteins (cysteine protease, adhesin, and hemagglutinin) are processed from a large polyprotein encoded by a single gene (26). It is possible that *Orf8*, which displays a putative transmembrane segment, could anchor the adhesin in the cytoplasmic membrane. The fact that *cwp66*, considered as a putative colonization factor, is transcribed only in the beginning of the exponential phase, while no transcription of toxins could be detected (13, 16), is somewhat surprising. But this phenomenon has already been described for *Staphylococcus aureus*, in which cell wall-associated adhesins, such as the fibronectin binding-proteins, are expressed during the exponential phase and repressed postexponentially when synthesis of exoproteins is induced (30, 39).

In a recent publication, Calabi and Fairweather (4) found no polycistronic RNA transcript for this cluster. This discrepancy could be explained first by the different strain studied and, second, by different culture conditions (especially for *slpA*, which could be transcribed from its own promoter), but lack of details about the experiments undertaken by Calabi and Fairweather prevents us from drawing definitive conclusions.

As the Cwp66 protein has already been demonstrated to be involved in adherence of *C. difficile* to cultured cells and could therefore be involved in pathogenesis of the bacterium, it seemed interesting to compare sequences of this gene from different clinical isolates in order to establish virulence profiles. Unfortunately, although six different couples of primers were used, we obtained a specific amplification product only for 14 strains. The absence of amplification of the 3' part of *cwp66* in some strains, regardless of serogroup or toxinotype, could be explained by the high variability of this domain, as previously

described by dot blot experiments (37). However, we cannot exclude the possibility that this domain could be deleted in some strains. The sequence of *cwp66* does not display any obvious correlation with serogroup or virulence. However, it is noteworthy that the four nonrelated serogroup C strains shared the same conserved *cwp66* gene sequence, which is divergent from those of the other strains: this serogroup is known to contain the most outbreak-related strains (36), and conservation of this adhesin could be of importance for colonization and dissemination of these strains.

To further our understanding of the role of all the component genes of this cluster, we investigated *cwp84*, a gene located just downstream from *cwp66*. The anchoring domain of Cwp84 is located in the C-terminal part, and the N-terminal domain displays significant homologies to cysteine proteases of archaea (e.g., *Methanosarcina mazei*, GenBank number NC_003901), eukaryotes (e.g., papain, GenBank number M15203), and bacteria (e.g., PepC, *Streptococcus thermophilus*, GenBank number Q56115), especially around the active amino acids of the enzymes. Alignments allow us to determine the putative catalytic triad: cysteine in position 216, histidine in 262, and asparagine in 287. *C. difficile* is not currently considered a proteolytic bacterium, but some studies have shown that this bacterium displays some surface-associated proteolytic activity (28, 32), and most proteolytic strains have been shown to be the most virulent in the hamster model (31). Moreover, surface-associated proteolytic activity seems to be mainly due to a thiol-protease (32). Proteases are a well-known virulence factor for some important pathogenic bacteria (22), and especially the cysteine proteases produced by *Porphyromonas gingivalis* or *Streptococcus pyogenes* (21, 35).

Cloning of the *cwp84* gene into an expression system in an *E. coli* strain with deletions of major proteases demonstrated non-specific proteolytic activity of this protein on gelatin, skimmed milk, and azocoll. Purification of this protease failed, likely due to an autocatalytic process of the protease, leading to the cleavage between the GST and the catalytic domain of Cwp84. This phenomenon has been described for other extracellular cysteine proteases (38), such as SpeB, a well-characterized virulence factor of *S. pyogenes* (12). The fact that leupeptin and E64 do not inhibit the autocatalytic process of Cwp84 is somewhat surprising, but other cysteine proteases, like Lys-gingipain and clostripain, respectively, are also not inhibited by these molecules. Further purification attempts with different strategies are in process.

The transcription of the *cwp84* gene could be detected only in the early exponential phase, but in a pronounced fashion. Sequence analysis of the functional domain of the *cwp84* gene indicated that this gene is highly conserved, particularly the amino acids potentially involved in the active site. No correlation between Cwp84 sequence patterns and serogroups or virulence profiles was found. In fact, strains from different serogroups or with different toxinotypes shared the same peptidic sequence. It is noteworthy that SpeB and gingipains, cysteine proteases involved in virulence of *P. gingivalis*, have been shown to be highly conserved and expressed among all the strains (24, 25, 33). Taken together, the highly conserved pattern of this gene and its strong expression, at least in the strain studied and in our experimental conditions, indicate that Cwp84 might have an important function in the physiology of

C. difficile. A tempting hypothesis would be that Cwp84 acts as a maturation protease for various cell surface-associated proteins, as it has been shown for SpeB in *S. pyogenes* (1) and the gingipains in *P. gingivalis* (18). In particular, Cwp84 could play a role in the processing of the S-layer precursor or of Cwp66, for which some cleavage products have been detected in surface extracts of *C. difficile* (37).

The study of the transcription of the genes located on this putative virulence cluster is an important step in the characterization of the colonization process by *C. difficile* and will be further confirmed by in vivo experiments, in which transcription of genes coding for adhesins and toxins will be studied in a *C. difficile* monoxenic mouse model.

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