Identification, Cloning, and Expression of the CAMP-Like Factor Autotransporter Gene (cfa) of Bartonella henselae

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The CAMP reaction was first described by Christie et al. (R. Christie, N. E. Atkins, and E. Munch-Petersen, Aust. J. Exp. Biol. 22:197–200, 1944) as the synergistic lysis of sheep red blood cells by Staphylococcus aureus sphingomyelinase and CAMP factor (cohemolysin), a secreted protein from group B streptococci. We observed a CAMP-like reaction when Bartonella henselae was grown in close proximity to S. aureus on 5% sheep blood agar. This study describes the cloning, sequencing, and characterization of a CAMP-like factor autotransporter gene (cfa) from B. henselae. A cosmid library of B. henselae ATCC 49793 was constructed using SuperCos1 in Escherichia coli XL1-Blue MR. Cosmids were screened for the CAMP reaction, and a quantitative cohemolysis microtiter assay was developed using purified sphingomyelinase. Cosmid clones with the strongest cohemolytic reaction had similar restriction enzyme patterns. A DNA fragment that expressed the cohemolysin determinant was subcloned in a 7,200-bp StuI-BamHI fragment which contained a 6,024-bp open reading frame. The reaction had similar restriction enzyme patterns. A DNA fragment that expressed the cohemolysin determinant was subcloned in a 7,200-bp StuI-BamHI fragment which contained a 6,024-bp open reading frame. The deduced amino acid sequence showed homology to the family of autotransporters. The autotransporters are a group of proteins that mediate their own export through the outer membrane. They contain an N-terminal passenger region, the α-domain, and a C-terminal transporter region, the β-domain. The α-domain contains four, nearly identical 42-amino-acid repeats and showed homology to the family of RTX (repeat in toxin) hemolysins. The concentrated supernatant of the recombinant strain expressed a protein with a molecular mass of 180 kDa on sodium dodecyl sulfate-polyacrylamide gel electrophoresis consistent with the calculated molecular weight of the secreted α-domain. In conclusion, we have characterized a novel secreted cohemolysin autotransporter protein of B. henselae.

Bartonella henselae is an extremely fastidious, pleomorphic, gram-negative rod that has been found to be associated with cat scratch disease, bacillary angiomatosis, and bacillary peliosis (9, 12, 28, 48). Of these syndromes, cat scratch disease is the most common and affects an estimated 25,000 people in the United States annually, the majority of which are children (25).

Bartonella spp. are the only bacterial pathogens for humans that engage in erythrocyte parasitism (30, 38) and require erythrocytes or heme supplements in order to grow in vitro (42). Most B. henselae isolates require more than 7 days of incubation before growth can be detected. Typically they are plated on freshly prepared enriched (chocolate- or blood-containing) medium incubated at 35 to 37°C with 5 to 10% CO2 and >40% humidity. There are two types of colony morphology for B. henselae. They can appear as dry, irregular whitish raised colonies or moist circular tan colonies, often in the same culture. B. henselae colonies are not hemolytic on blood agar (47). We, however, observed a synergistic lysis of the red cells when B. henselae was grown in close proximity to S. aureus on 5% sheep blood agar. Cohemolysis (CAMP-like reaction) has not been previously described for B. henselae. The CAMP reaction, as originally described in 1944 by Christie, Atkins, and Munch-Peterson (whose initials give rise to the eponym CAMP) (8), is the synergistic lysis of sheep erythrocytes by Staphylococcus aureus sphingomyelinase C (beta-toxin) and CAMP factor (cohemolysin), a secreted 25.3-kDa protein from group B streptococci (Streptococcus agalactiae) (36). The CAMP reaction is performed with erythrocytes whose cell membranes contain at least 45 mol% sphingomyelin. Human erythrocytes have a sphingomyelin:cholesterol ratio of 1:5 and are known not to undergo CAMP hemolysis (8).

The CAMP reaction occurs in a sequential two-step process. The first reaction is the hydrolysis of membrane sphingomyelin and phospholipids by the action of a sphingomyelinase or phospholipase (11). In the second reaction, the CAMP factor interacts with the metastable red cell membrane, leading to cell lysis (3). A number of other gram-positive and gram-negative bacteria are known to react positively in the CAMP test, including Listeria monocytogenes (33), Listeria seeligeri (37), Rhodococcus equi (34), Pasteurella spp. (15), Aeromonas spp. (14), certain Vibrio spp. (29), group G streptococci (44), and Actinobacillus pleuropneumoniae (17).

Since the cohemolysin of B. henselae causes lysis of red blood cells, it is considered a potential virulence factor. To facilitate studying the role of the CAMP reaction in the pathogenesis of B. henselae infection, we cloned and characterized the CAMP-like factor of B. henselae. The deduced amino acid sequence of this gene is homologous to the family of autotransporters. The autotransporters are a family of diverse proteins secreted by gram-negative bacteria that mediate their own export through the outer membrane. They contain an N-terminal passenger region, the α-domain, and a C-terminal transporter region, the β-domain. This study describes the cloning,
sequencing, and characterization of a gene encoding a novel CAMP-like factor (Cfa) of *B. henselae* that possesses the characteristics of an autotransporter virulence protein.

**MATERIALS AND METHODS**

**Bacterial strains and plasmids.** Characteristics of the bacterial strains and plasmids used in this study are described in Table 1.

**Media.** *E. coli* strains were routinely grown in Luria-Bertani broth (LB). Kanamycin (50 μg/ml) was added as appropriate. *S. aureus* and *L. monocytogenes* were routinely grown on Columbia blood agar base (Difco, Detroit, Mich.) containing 5% defibrinated sheep blood (Difco) at 37°C.

**DNA manipulations and cloning.** Standard methods were followed for molecular biological techniques (41). Oligonucleotides were synthesized at the Huntsman Cancer Center Peptide and DNA facility.

**CAMP test.** The CAMP test for cohemolytic activity was performed on fresh *B. henselae* ATCC 49739 strain was streaked on the surface of the blood agar and then incubated for 7 days at 35°C under 5% carbon dioxide. After 7 days of growth, the sphingomyelinase-strain was streaked on the surface of the blood agar and then incubated for 7 days perpendicular and juxtaposed to the cultures were centrifuged 5 min at 10,000 rpm (3,000 × g) at 4°C. A microtiter plate was constructed using a SuperCos1 cosmid vector kit (Stratagene, La Jolla, Calif.).

**Construction of *B. henselae* cosmid library.** A cosmid library of *B. henselae* ATCC 49739 was constructed using a SuperCos1 cosmid vector kit (Stratagene, La Jolla, Calif.).

**B. henselae* genomic DNA was partially digested with BamHI. The digested DNA was sized to yield 30- to 40-kb fragments, dephosphorylated, and then ligated into the BamHI site of SuperCos1, previously linearized with XbaI. Packaging of the cosmids into phage and their subsequent infection in the *E. coli* strain XLI-Blue MR were performed as described by the manufacturer.

**Preparation and analysis of outer membrane, cytoplasmic, and secreted proteins.** Enriched outer membrane proteins and cytoplasmic proteins were prepared by modification of the procedure by Hantke (21). Exponentially grown cells were pelleted for 10 min at 4°C at 10,000 rpm (8,000 × g) in a Beckman JA-20 in a 50-ml tube. The pellet was resuspended in 1 ml 0.2 M Tris, pH 8.0, and transferred to a microcentrifuge tube. The bacteria were pelleted at 16,000 × g for 2 min at 4°C. After decanting of the supernatant, the pellets were resuspended in 50 μl of 0.2 M Tris (pH 8.0) on ice. In sequence on ice, 100 μl of 0.2 M Tris (pH 8.0), 1 μl sucrose, 10 μl of 10 mM EDTA (pH 8.0), 10 μl of lysozyme (2 mg/ml), and 320 μl of distilled water were added with gentle mixing after each addition. The mixture was incubated for 10 min at room temperature. Then 10 μl of DNase (bovine pancreatic DNase I, 1 mg/ml in 0.15 M NaCl, 50% glycerol; Sigma) and 500 μl of 2% Triton X-100, 10 mM MgCl2, 50 mM Tris (pH 8.0) were added. The solution was centrifuged at 16,000 × g for 30 min at 4°C. The supernatants (cytoplasmic proteins) were suspended in 2× Laemmli sample buffer. After removal of the supernatants, the pellets were washed four times with ice-cold distilled water and then resuspended in 100 μl of 2× electrophoresis sample buffer.

For purification of supernatant (secreted proteins), 100-ml cultures were incubated for 16 to 18 h at 37°C on a rotary shaker at 225 rpm. Culture supernatants were recovered by centrifugation and filtered. Ammonium sulfate was dissolved to a final concentration of 60% saturation. The precipitate was centrifuged and dissolved in 3 ml of 50 mM Tris-HCl, pH 7.5, and this preparation was dialyzed against 50 mM Tris-HCl (pH 7.5). This resulted in an approximately 10-fold concentration of the original supernatant. The protein preps were analyzed by sodium dodecyl sulfate (SDS)-4 to 15% polyacrylamide gel (Ready Gel; Bio-Rad Laboratories, Inc. Hercules, Calif.) and stained with Coomassie blue.

**Western blot.** Ten microliters of proteins (approximately 6.5 mg/ml) were loaded to each well of a 4 to 15% SDS-polyacrylamide gel electrophoresis (PAGE). Electrophoresis was carried out at 200 V for 30 to 60 min. The separated proteins were then transferred to a nitrocellulose membrane at 100 V for 1 h using the Mini-Trans-Blot electrophoretic transfer cell (Bio-Rad Laboratories). The membranes were then blocked overnight at 4°C with 3% dry milk in Tris-buffered saline (TBS) solution. Serum samples were diluted 1:200 in a diluent wash (3% dry milk in TBS).

The serum positive for antibodies against *B. henselae* used for Western blotting was pooled from 68 cases of suspected cat scratch disease cases that were
submitted to ARUP laboratories (Salt Lake City, Utah) for confirmative diagnosis. These sera had immunoglobulin G (IgG) titers ranging from 1:512 to 1:4,096 for *B. henselae*, as determined by indirect immunofluorescent assay (IFA) (Focus Technologies, Cypress, Calif.). The IFA titer of the pooled specimen was 1:1,024. The *B. henselae* antibody-negative sera used as a control for Western blotting had titers of <1:64 by IFA. The procedures followed were in accord with the ethical standards established by the University of Utah and are in accord with the Helsinki Declaration of 1975. Specimens were collected under approval by the University of Utah Institutional Review Board (IRB no. 11343). Specimens were stored at −20°C until testing commenced and were then stored at 2 to 8°C while the evaluations were performed.

The serum was preadsorbed with *E. coli* XL1-Blue (containing SuperCos1 vector) whole-cell lysate. The diluted primary antibody solution was incubated with a 1:10 (vol/vol) dilution of the lysate for 30 min with shaking at 37°C to reduce background antibodies reacting against *E. coli* proteins.

After preadsorption, the sera were reacted with the membrane containing the protein samples for 3 h on a rocking platform. The membrane was then washed three times, changing the diluent wash every 5 min. A 1:10,000 dilution of alkaline phosphatase goat anti-human IgG (γ-chain-specific) conjugate (Sigma) was then added to the membranes. The membrane was then incubated for 1 h and washed as before, before the addition of 5-bromo-4-chloro-3-indolyl phosphate/nitroblue tetrazolium (BCIP/NBT) substrate (Sigma). This reaction was carried out for 5 to 10 min and then stopped by the addition of distilled water.

**DNA sequencing.** The DNA sequence was determined by the AB 3700 96 capillary DNA Analyzer from Applied Biosystems. Synthetic oligonucleotides used as primers for DNA sequencing were synthesized by the Huntsman Cancer Center DNA peptide facility, University of Utah.

**DNA and protein database searches.** The National Center for Biotechnology Information Services were used to consult the SwissPROT, GenBank, and EMBL databases with the BLAST algorithm (1, 20). Prediction of the signal peptide cleavage site was performed using the SignalP 3.0 Server, http://www.cbs.dtu.dk/services/SignalP/ (2).

**Nucleotide sequence accession number.** The GenBank accession number for the sequence presented in this article is AY695890.

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**RESULTS**

**Demonstration of the CAMP effect (cohemolysis) in a *B. henselae* ATCC 49793 strain.** We analyzed *B. henselae* for cohemolytic activity by growing the organism on fresh Columbia 5% sheep blood agar plates for 7 days at 35°C under 5% carbon dioxide. After 7 days of growth, a sphingomyelinase-producing *S. aureus* indicator strain was streaked vertically on the surface of the blood agar plate, and the *B. henselae* strain was streaked horizontal to the *S. aureus* streak and incubated an additional 18 h at 37°C. The *B. henselae* strain showed distinct cohemolytic zone of increased hemolysis within the diffusion zone (a) of the *S. aureus* strain after 18 h of growth of the *S. aureus* strain (Fig. 1, lane 1).

**Cloning and sequencing a cohemolysin gene from *B. henselae* ATCC 49793.** To clone the cohemolysin determinant in *E. coli* we constructed a cosmid library of *B. henselae* ATCC 49793 in SuperCos1 cosmid. Three-hundred cosmid clones were screened by the CAMP test. Eighteen cosmid clones exhibited at least a slight CAMP effect. Four cosmid clones showed the strongest CAMP activity similar to *Listeria monocytogenes* CAMP reaction and had similar restriction patterns when digested with HindIII, EcoRI, or BglII. One of these cosmids, pCML75, containing a 29.7-kb *B. henselae* chromosomal insert, was restriction mapped and sequenced (Fig. 2). DNA fragments of cosmid pCML75 were subcloned and tested for cohemolytic activity. Cohemolytic activity was isolated with a 7.2-kb fragment on one end of the DNA insert, from a StuI restriction site to the BamHI site within the multiple cloning site of a plasmid derivative of SuperCos1 (Table 1), pCML76 (Fig. 2).

Figure 1 illustrates examples of the CAMP effect from various bacterial species and the cosmid clones and plasmid sub-
clones of *B. henselae* DNA in *E. coli*. The *B. henselae* strain exhibited CAMP activity, with an area of hemolysis within the sphingomyelinase diffusion zone. The control strain of group B streptococcus showed a large distinct zone of complete hemolysis, and the *L. monocytogenes* control strain showed a mild CAMP reaction. The *E. coli* XL1-Blue strain containing pCML75 with the 29.7–kb insert of *B. henselae* chromosomal DNA in SuperCos1 showed mild cohemolysis. The *E. coli* XL1-Blue strain containing plasmid pCML76 with the 7.2-kb subclone demonstrated moderately strong CAMP activity.

The *B. henselae* cohemolysin is homologous to two protein families, RTX hemolysins and autotransporters. Sequence analysis of the 7.2-kb subclone demonstrated a 6,024-bp open reading frame coding for 2,008 amino acids. The nucleotide sequence is identical to nucleotide sequence at locus tag BH13030 of the recently sequenced *B. henselae* Houston strain-1 genome (GenBank accession no. YP034038). The gene was named cfa (for CAMP-like factor autotransporter). The open reading frame begins 160 bp downstream of a StuI restriction site. A putative Shine-Dalgarno sequence is located just 3 bp upstream from the initiating methionine. A signal sequence of 33 amino acids is predicted using the Signal P program.

When the deduced amino acid sequence was subjected to a BLAST similarity search the midportion of the protein and carboxy-terminal end of the protein shows separate homologies with calcium dependent RTX (repeat in toxin) hemolysins and the family of autotransporters, respectively. This homology suggests that like other autotransporters, the protein has three distinct domains: a signal peptide, a secreted α-domain, and a transporting β-domain. The precursor form of Cfa, including the leader sequence, presumed secreted α-domain and the transporting β-domain has a predicted molecular mass of 214,260 and calculated isoelectric point of 4.96. The presumed secreted α-domain has a calculated molecular mass of 179,493 and a calculated isoelectric point of 4.75. The predicted outer membrane transporting β-domain has a calculated molecular mass of 31,179 and isoelectric point of 8.98. In Fig. 2, the grey area and vertical bars in the arrow representing Cfa shows the region of homology of the translated protein with RTX family of (repeat in toxin) hemolysins and toxins. A range of 21 to 28% identity and 36 to 48% similarity is seen with a number of RTX toxins and hemolysins of *Chromobacterium violaceum* (GenBank accession no. AAQ57990), *Novosphingobium aromaticivorans* (ZP00303863), *Magnetospirillum magnetotacticum* (ZP00055704), *Nitrosomonas europaea* (CAD84072), and *E. coli* (AE016756), from amino acids 748 to 970. Figure 3A shows a comparison of part of this region of Cfa with the RTX hemolysins of *Chromobacterium violaceum* and *Nitrosomonas europaea*.

From amino acid 824 to 991, there are four nearly identical tandem glycine- and serine-rich amino acid repeats comprised of the following 42 amino acids in single letter code: P-S-S-V-E-T-S-I-T-P-T-A-V-S-G-D-N-S-A-G-P-V-G/R-G-E/G-Q-S-A-S/P-V-A-R-S-E-S (D-G-G-V-T-V-V-L-S). Slashes and boldface lettering indicate the variant amino acids. The last 9 amino acids of the 42-amino-acid sequence (in parentheses) are only repeated three times instead of four. Although Cfa shows some similarity to RTX toxins and contains an area of glycine-rich repeats, the repeats are not similar to the glycine-rich repeat consensus sequence described for RTX family of toxins L/A-F-X-G-X-G-N/D-D-X (46).

A BLAST search for “short nearly exact matches” against amino acids 3 to 33 of the 42 listed above reveals 52% identity and 58% similarity to a repeating amino acid sequence found
in Streptococcus gordonii platelet binding protein (serine-rich repeat protein) GspB (AAL13053).

The last 286 amino acids of the deduced amino acid sequence of cfa (from 1723 to 2008) shows homology to the family of autotransporters. A range of 23 to 28% identity and 38 to 46% similarity is seen with a number of autotransporter proteins including E. coli YchA (BAA97898), Bordetella pertussis Vag8 (AAC31247), Salmonella enterica serovar Typhimurium ShdA (AAD25110), and Yersinia pestis YapH (CAC14227). Figure 3B shows a comparison of part of this region of Cfa with the autotransporters, E. coli YchA and B. pertussis Vag8.

Expression of the cohemolysin protein in E. coli. We developed a cohemolysin microtiter assay to quantitatively measure the amount of red blood cell hemolysis by unconcentrated bacterial supernatants in the presence of sphingomyelinase. We derived our protocol based on previously published hemolysis assays (19) and protocols for the pretreatment of tissue culture cells with sphingomyelinase to sensitize cells to the effects of toxins (35). Red blood cells untreated and treated with sphingomyelinase were tested in tandem to compare the hemolytic versus the cohemolytic activity of the bacterial supernatants (Table 2).

Panel A of Fig. 4 shows the results of a microtiter hemolysis assay of the bacterial supernatants without pretreatment of sheep red blood cells with sphingomyelinase. No agglutination of the sheep red cells was observed for any of the bacterial culture supernatants. Hemolytic activity of the culture supernatant from E. coli containing the original cosmid clone, pCML75 (Fig. 4A, row E), was minimal to none at a 1:2 dilution. The bacterial supernatant from L. monocytogenes showed slight hemolysis at an endpoint titer of 1:4 (Fig. 4A, row A). The bacterial supernatant from E. coli containing plasmid subclone pCML76 (cfa gene and its promoter region) showed lysis of the sheep red blood cells up to an endpoint titer of 1:8 (Fig. 4A, row F), as observed by a reduction in the size of the red blood cell pellet. No detectable hemolysis was observed with the bacterial supernatant of E. coli containing the subclone pCML77 (pCML76 with an internal 5.6-kb EcoRV-EcoRV deletion of cfa) or with the bacterial supernatant of E. coli containing the cosmid cloning vector SuperCos1.

Panel B of Fig. 4 shows the results of a microtiter cohemolysis assay of the bacterial supernatants with pretreatment of the sheep red blood cells with sphingomyelinase (0.025 U/ml) for 30 min. A trace amount of hemolysis was observed directly
around the red blood pellet of all the treated cells, including the red cell controls without bacterial supernatant (Fig. 4B, row B), indicating a slight hemolytic effect with sphingomyelinase treatment alone. Hemolysis was observed to an endpoint titer of 1:64 for the *L. monocytogenes* positive control (Fig. 4B, row A). Hemolytic activity of the culture supernatant from *E. coli* containing the original cosmid clone, pCML75 (Fig. 4B, row E), was enhanced with sphingomyelinase treatment to an endpoint titer of 1:32. The bacterial supernatant from *E. coli* containing plasmid subclone pCML76 (*cfa* gene and its promoter region) also showed enhanced lysis with sphingomyelinase pretreatment with lysis of the sheep red blood cells up to an endpoint titer of 1:128 (Fig. 4B, row F), as observed by a reduction in the size of the red blood cell pellet. No hemolysis compared to negative control was observed with the bacterial supernatant of *E. coli* containing the subclone pCML77 (pCML76 with an internal deletion of *cfa*) or with the bacterial supernatant of *E. coli* containing the cosmid cloning vector SuperCos1.

We also performed the microtiter assay on human and feline erythrocytes (Table 2). Human erythrocytes have been reported not to undergo CAMP hemolysis (8). We observed only slight hemolysis of the human erythrocytes without pretreatment with sphingomyelinase at a 1:2 dilution of the *L. monocytogenes* supernatant. Hemolysis was enhanced in human erythrocytes treated with sphingomyelinase to an endpoint titer of 1:8 for *L. monocytogenes*. No detectable hemolysis of the untreated or treated human erythrocytes was observed with any of the *E. coli* strains containing plasmid clones. The hemolysis and cohemolysis microtiter assay using feline erythrocytes showed titers that were two- or fourfold lower than what was observed with sheep erythrocytes.

**Protein and Western blot analysis of the cohemolysin protein in *E. coli***. We chose to examine the expression of Cfa in *E. coli* by SDS-PAGE and Western blot due to the slow growth of *B. henselae* in liquid and on solid media. Concentrated culture supernatants, outer membrane proteins, and cytoplasmic proteins from *E. coli* containing the SuperCos1 cloning vector and the plasmid pCML76 containing the *cfa* clone was subjected to SDS-PAGE and Western blot analysis (Fig. 5). A protein band of an apparent molecular mass of 180 kDa was observed in the concentrated supernatant of the *E. coli* containing the *cfa* clone (pCML76; Fig. 5A, lane 2) not seen in the concentrated supernatant of the *E. coli* strain containing SuperCos1 (Fig. 5A, lane 1). A similar size band was observed in the original cosmid clone pCML75 (data not shown). The observed 180-kDa band is consistent with the molecular weight of the predicted mature secreted protein (α-domain) of Cfa. Slightly higher molecular weight bands are observed in the cytoplasmic proteins of both the *E. coli* containing SuperCos1 and pCML76 (Fig. 5A, lanes 5 and 6). The β-domain of Cfa with the predicted molecular mass of 31.2 kDa would be expected to be observed in the outer membrane of the bacterial cell. The outer membrane protein preparations of the *E. coli* containing SuperCos1 and *E. coli* containing pCML76, however, appeared identical with no additional proteins bands observed in this region.

A Western blot of the same proteins, probed with sera pool from 68 patients with cat scratch disease (IgG IFA titer ranging from 1:512 to 1:4,096 for *B. henselae*) showed reactivity against the *E. coli* proteins in addition to the 180-kDa protein band unique to the concentrated bacterial supernatant of *E. coli* containing the *cfa* clone (Fig. 5B, lane 2). The same Western blot probed with sera negative for *B. henselae* anti-
A diverse group of toxins, proteases, and phospholipases have been identified as CAMP factors, including phospholipase C of *L. monocytogenes* (33). Apx toxins of *A. pleuropneumoniae* (17), RTX toxins of *Pasteurella aerogenes* (31), cholesterol oxidase of *Rhodococcus equi* (34), and O-sialoglycoprotein endopeptidase of *Riemerella anatipestifer* (10).

The cohemolysin Cfa of *B. henselae* is homologous to the family of autotransporters. Autotransporters (or proteins with type V secretion pathways) have three common characteristics, an N-terminal signal sequence for translocation across the inner membrane, a passenger domain (α-domain) that is transported to the cell surface, and a conserved C-terminal transporter (β-domain) that facilitates secretion across the outer membrane through a beta-barrel porin structure (24).

Although the β-domains of autotransporters are highly conserved, the passenger α-domains are widely divergent, reflecting their many different functions (22). Virulence determinants are often secreted to the bacterial cell surface or released into the outside environment. Numerous diverse virulence factors have been classified as autotransporter proteins including toxins, proteases, lipases, hemagglutinins, and adhesions (22). A cohemolysin or hemolysin has yet to be characterized as an autotransporter protein.

The α-domain of Cfa shows some homology to the RTX (repeats in structural toxin) toxins, a class of pore-forming toxins found among various species of *Pasteurella* (31, 32) and *Actinobacillus* (6, 7, 18) that play an important role in pathogenesis. Pax of *P. aerogenes* (31) and the ApxD, ApxII, and ApxIII RTX toxins of *A. pleuropneumoniae* (17) have a cohemolytic phenotype. The RTX toxins, however, have a type I secretion system in which secretion requires three accessory proteins which comprise a channel spanning both the inner and outer membranes (13). Although Cfa has a series of glycine-rich amino acid repeats in the α-domain, these repeats do not fit the repeat consensus sequence described for the RTX family of toxins (46). Therefore, without the consensus repeats and the operon consisting of the genes involved in transport and post-translational modification, Cfa cannot be considered an RTX protein.

On Western blot, the secreted α-domain of Cfa was recognized by a sera pool of 68 patients with suspected cat scratch disease, positive for *B. henselae* antibodies by IFA. Cfa could be a potential candidate for immunodiagnostic test for the identification of individuals with *B. henselae* antibodies. Further studies need to be performed to establish the sensitivity and specificity of this antigen in diagnostic tests for *B. henselae* antibodies.

The carboxy-terminal domain of Cfa is highly homologous to the β-domains of autotransporters, which form porin structures in the outer membrane to facilitate the secretion of the passenger domain. Several *B. henselae* hypothetical proteins (BH06590, BH13160, BH13180) also show a high degree of homology with the C-terminal end of Cfa. Based on the homology of Cfa with these β-domains, an outer membrane protein of approximately 31 kDa should be visualized on examination of the outer membrane protein of the cfa clone pCML76 (Fig. 5A, lane 4). No new protein, however, was identified on SDS-PAGE of the outer membrane different than the control strain containing the cloning vector SuperCos1 (Fig. 5A, lane 3). The protein may be either rapidly degraded or poorly expressed and thus not detected with Coomassie blue stain. Conversely, since the cfa gene was expressed in *E. coli* and not *B. henselae*,...
the protein may not be processed properly in this heterologous host. Because of the difficulty with absorbing out anti- *E. coli* antibodies, we also cannot rule out that there is not a reaction of the antisera with a protein corresponding to a 31-kDa band corresponding to the C-terminal protein. The lack of detection of the β-domain outer membrane in SDS-PAGE has been previously noted in other autotransporters, including the serine protease A of *Neisseria meningitidis* (45). Although the *B. henselae* CAMP factor shows homology to regions from *Shigella* protein is secreted in this manner. In membrane preparations, we do not yet have proof that the 180-kDa were unable to show the C-terminal protein in outer membrane preparations, because we previously noted in other autotransporters, including the secretion of the antisera with a protein corresponding to a 31-kDa band from the National Institute of Allergy and Infectious Diseases to further study the CAMP reaction for the prompt, presumptive identification of *Streptococcus agalactiae* (Lancefield group B) in clinical material. J. Clin. Microbiol. 1:171–174.


