# Genetic-Biochemical Analysis and Distribution of the Ambler Class A β-Lactamase CME-2, Responsible for Extended-Spectrum Cephalosporin Resistance in *Chryseobacterium (Flavobacterium) meningosepticum*

SAMUEL BELLAIS, LAURENT POIREL, THIERRY NAAS, DELPHINE GIRLICH, and PATRICE NORDMANN\*

Service de Bactériologie-Virologie, Hôpital de Bicêtre, Assistance Publique/Hôpitaux de Paris, Faculté de Médecine Paris-Sud, 94275 Le Kremlin-Bicêtre Cédex, France

Received 17 March 1999/Returned for modification 27 June 1999/Accepted 6 September 1999

In vitro synergy between extended-spectrum cephalosporins and either clavulanic acid or cefoxitin was found for Chryseobacterium meningosepticum isolates during a double-disk assay on an agar plate. An extendedspectrum β-lactamase (ESBL) gene from a C. meningosepticum clinical isolate was cloned and expressed in Escherichia coli DH10B. Its protein conferred resistance to most β-lactams including extended-spectrum cephalosporins but not to cephamycins or to imipenem. Its activity was strongly inhibited by clavulanic acid, sulbactam, and tazobactam, as well as by cephamycins and imipenem. Sequence analysis of the cloned DNA fragment revealed an open reading frame (ORF) of 891 bp with a G+C content of 33.9%, which lies close to the expected range of G+C contents of members of the Chryseobacterium genus. The ORF encoded a precursor protein of 297 amino acids, giving a mature protein with a molecular mass of 31 kDa and a pI value of 9.2 in E. coli. This gene was very likely chromosomally located. Amino acid sequence comparison showed that this β-lactamase, named CME-2 (C. meningosepticum ESBL), is a novel ESBL of the Ambler class A group (Bush functional group 2be), being weakly related to other class A β-lactamases. It shares only 39 and 35% identities with the ESBLs VEB-1 from E. coli MG-1 and CBL-A from Bacteroides uniformis, respectively. The distribution of bla<sub>CME-2</sub> among unrelated C. meningosepticum species isolates showed that bla<sub>CME-2</sub>-like genes were found in the C. meningosepticum strains studied but were absent from strains of other C. meningosepticum-related species. Each C. meningosepticum strain produced at least two  $\beta$ -lactamases, with one of them being a noninducible serine ESBL with variable pIs ranging from 7.0 to 8.5.

Chryseobacterium meningosepticum (formerly classified as *Flavobacterium meningosepticum* [45]) is a waterborne saprophytic bacterium. Among *Chryseobacterium* species, *C. meningosepticum* is most commonly associated with infections in humans. It may cause meningitis in newborns and pneumonia and sepsis in immunocompromised patients, especially those hospitalized in intensive care units (3, 39). *C. meningosepticum* is naturally resistant to most  $\beta$ -lactams, including extended-spectrum cephalosporins and carbapenems, with only some isolates remaining susceptible to ureidopenicillins (6).

Phenotype analysis of the  $\beta$ -lactam resistance pattern of a *C.* meningosepticum PINT clinical isolate revealed the presence of a putative extended-spectrum  $\beta$ -lactamase (ESBL) according to the synergy found between clavulanic acid and most extended-spectrum cephalosporins when a double-disk assay was performed on an agar plate (20). Uncommonly, a similar synergy was also found between cephamycins such as cefoxitin or moxalactam and extended-spectrum cephalosporins. Recently, an Ambler class B carbapenem-hydrolyzing  $\beta$ -lactamase has been reported from *C. meningosepticum* CIP 6058 (36). Although the hydrolysis spectrum of this  $\beta$ -lactamase is broad, its presence cannot be responsible for the extended-spectrum cephalosporin resistance profile observed in *C. meningosepticum*. The aim of this work was to analyze both biochemically and genetically the  $\beta$ -lactamase responsible for the observed phenotype and to determine its distribution among nonepidemiologically related *C. meningosepticum* isolates and other *Chryseobacterium* species strains.

## MATERIALS AND METHODS

**Bacterial strains.** The bacterial strains used in this work are listed in Table 1. *C. meningosepticum* PINT was isolated at the Raymond Poincaré Hospital in Garches, France, a suburb of Paris. *C. meningosepticum* AMA and GEO were isolated at the Bicêtre Hospital (Le Kremlin-Bicêtre, France), and both were from tracheoalveolar aspirates. Reference strains were from the Pasteur Institute (Paris, France) and Denmark (7). The *C. meningosepticum* isolates and reference strains were epidemiologically unrelated (data not shown).

*Escherichia coli* DH10B and nalidixic acid- and rifampin-resistant *E. coli* JM109 were used for cloning and conjugation assays, respectively (Table 1). The *Chyseobacterium* sp. strains were identified by standard techniques as described previously (30, 39), and their identities were confirmed with the API 32GN system (bioMérieux, Marcy l'Etoile, France). All strains were stored at –70°C in Trypticase soy (TS) broth (Becton Dickinson, Le Pont de Claix, France) supplemented with 15% glycerol until testing.

Antimicrobial agents and MIC determinations. The antimicrobial agents used in this study were obtained in the form of standard laboratory powders and were used immediately after their solubilization. The agents and their sources have been described elsewhere (32). Antibiotic disks were used for routine antibiograms (Sanofi-Diagnostics Pasteur, Marnes-La-Coquette, France).

MICs were determined by an agar dilution technique on Mueller-Hinton (MH) agar (Sanofi-Diagnostics Pasteur) with an inoculum of 10<sup>4</sup> CFU per spot (8, 23). All drugs were incorporated into MH agar at serial twofold concentrations, and the antimicrobial susceptibilities of all isolates were determined concomitantly. The plates were incubated at 35°C for 18 h. The MICs of  $\beta$ -lactams were determined alone or in combination with a fixed concentration of clavulanic acid (2 µg/ml), tazobactam (4 µg/ml), cefoxitin (0.1 µg/ml), or moxalactam or imipenem (0.05 µg/ml each).

<sup>\*</sup> Corresponding author. Mailing address: Service de Bactériologie-Virologie, Hôpital de Bicêtre, 78 rue du Général Leclerc, 94275 Le Kremlin-Bicêtre Cédex, France. Phone: 33-1-45-21-36-32. Fax: 33-1-45-21-63-40. E-mail: nordmann.patrice@bct.ap-hop-paris.fr.

Strain or plasmid	Relevant genotype or phenotype	Source or reference
Strains		
E. coli DH10B	F' mcrA $\Delta$ (mrr-hsdRMS-mrcBC) $\Phi$ 80 $\Delta$ lacZDM15 $\Delta$ lacX74 deoR recA1 endA1 araD139 $\Delta$ (ara, leu)7697 galU galK1 rpsL nupG	Gibco BRL
E. coli JM109	endÅ1 hsdR17 gyrÅ96 $\Delta$ (lac proÅ) recÅ1 relÅ supE44 thi F' (lacI <sup>q</sup> lacZ $\Delta$ M15 proÅB <sup>+</sup> traD36)	48
Rifampin-resistant E. coli JM109 obtained in vitro	Rifampin-resistant	This study
C. meningosepticum PINT	Extended-spectrum cephalosporin and carbapenem resistant	This study
C. meningosepticum AMA	Extended-spectrum cephalosporin and carbapenem resistant	This study
C. meningosepticum GEO	Extended-spectrum cephalosporin and carbapenem resistant	This study
C. meningosepticum CIP 6058	Extended-spectrum cephalosporin and carbapenem resistant	$IP^a$
C. meningosepticum CIP 6059	Extended-spectrum cephalosporin and carbapenem resistant	IP
C. meningosepticum CIP 7830	Extended-spectrum cephalosporin and carbapenem resistant	IP
C. meningosepticum CIP 7905	Extended-spectrum cephalosporin and carbapenem resistant	IP
C. meningosepticum AB1572	Extended-spectrum cephalosporin and carbapenem resistant	7
C. meningosepticum H01, J100	Extended-spectrum cephalosporin and carbapenem resistant	7
C. indologenes CIP 101026	Carbapenem resistant	IP
C. indologenes PIT	Carbapenem resistant	This study
S. multivorum CIP 103686	Carbapenem resistant	IP
M. odoratus CIP 103105	Carbapenem resistant	IP
Plasmids		
pBK-CMV phagemid	Neomycin and kanamycin resistant	Stratagene
pBR322	Recombinant plasmid containing the 560-bp SspI-PstI internal fragment of bla <sub>TEM-1</sub>	42
pHUC37	Recombinant plasmid containing the 435-bp PstI-NotI internal fragment of bla <sub>SHV-3</sub>	24
pPZ1	Recombinant plasmid containing the 1.1-kb SnaBI fragment of bla <sub>PER-1</sub>	25
pRLT1	Recombinant plasmid containing the 329-bp DraI-XmnI internal fragment of bla <sub>VEB-1</sub>	32
pPL1	Recombinant plasmid containing the 396-bp EcoRI-PvuI internal fragment of bla <sub>OXA-18</sub>	29
pBS1	1.9-kb Sau3AI fragment from C. meningosepticum PINT cloned into pBK-CMV	This study

TABLE 1. Bacterial strains and	plasmids used in this study
--------------------------------	-----------------------------

<sup>a</sup> IP, Institut Pasteur, Paris, France.

Cloning experiments and analysis of recombinant plasmids. Genomics DNAs from *C. meningosepticum* PINT and from other strains were extracted as described previously (25). The *XmnI* restriction endonuclease was from New England Biolabs (Ozyme, Saint Quentin en Yvelines, France), while all the other enzymes used in the cloning experiments were from Amersham Pharmacia Biotech (Orsay, France). Fragments from genomic DNA partially digested with *Sau3AI* were ligated into *BamHI*-restricted phagemid pBK-CMV from Stratagene (Ozyme) (37). Ligation was performed at a 1:2 vector-insert ratio with 200 ng of restricted genomic DNA in a ligation mixture containing 1 U of T4 DNA ligase at 4°C for 18 h. Recombinant plasmids were transformed by electroporation (Bio-Rad Gene Pulser II) into *E. coli* DH10B electrocompetent cells (Gibco BRL, Life Technologies, Cergy Pontoise, France). Antibiotic-resistant colonies (30 µg/ml).

Recombinant plasmid DNA was obtained from 100-ml TS broth cultures grown overnight in the presence of amoxicillin (100  $\mu$ g/ml) at 37°C. Plasmid DNAs were recovered by using Qiagen columns (Qiagen, Courtaboeuef, France). Plasmid mapping was performed after double restriction analysis. Fragment sizes were estimated by comparison with the fragment sizes on a 1-kb DNA ladder (Amersham Pharmacia Biotech).

**Conjugation assays and plasmid content.** Direct transfer of resistance genes into rifampin-resistant *E. coli* JM109 obtained in vitro was attempted by liquid and solid conjugation assays at 30 and  $37^{\circ}$ C. Transconjugants were selected on TS agar plates containing rifampin (200 µg/ml) and amoxicillin (50 µg/ml). Extraction of plasmid DNA from *C. meningosepticum* PINT was attempted by two different methods (10, 15).

**DNA sequencing and protein analysis.** Both strands of the 1.9-kb cloned DNA fragment of recombinant plasmid pBS1 were sequenced with an Applied Biosystems sequencer (ABI 373). The nucleotide sequence and the deduced protein sequence were analyzed with software available over the Internet at the National Center of Biotechnology Information website (http://www.ncbi.nlm.nih.gov) and at Pedro's BioMolecular Research Tools website (http://www.ncbi.nlm.nih.gov) and at Pedro's BioMolecular Research Tools website (http://www.fmi.ch/biology/research\_tools.html). Multiple protein sequence alignments were carried out with the program Clustal W, available over the Internet at the University of Cambridge. A dendrogram was derived from the multiple sequence alignment by a parsimony method with the phylogeny package PAUP (Phylogenetic Analysis Using Parsimony), version 3.0 (44). Among the Ambler class A  $\beta$ -lactamases, 11 were compared to CME-2: PER-1 from *Pseudomonas aeruginosa* RNL-1 (25), VEB-1 from *E. coli* MG-1 (32), CEP-A from *Bacteroides tuifor*-

mis WAL-7088 (40), TEM-3 from Klebsiella pneumoniae CFF104 (41), SHV-2 from Klebsiella ozaenae (11), MEN-1 from *E. coli* MEN (2), L-2 from *Stenotrophomonas maltophilia* 1275 IID (47), TOHO-1 from *E. coli* TUH12191 (13), and NMC-A from *Enterobacter cloacae* NOR-1 (22) as a representative of the serine carbapenem-hydrolyzing  $\beta$ -lactamase group (Bush group 2f) (5), which possesses an extended hydrolysis spectrum toward aztreonam.

**β-Lactamase preparations.** Cultures of *C. meningosepticum* clinical isolates and *E. coli* DH10B harboring recombinant plasmid pBS1 were grown overnight at 37°C in 100 ml of TS broth containing amoxicillin (100 µg/ml) and 4 liters of TS broth, respectively. Bacterial suspensions were pelleted, resuspended in 40 ml of Tris-HCl (50 mM) buffer (pH 8), disrupted by sonification (three times at 50 W for 30 s each time with a Vibra Cell 75022 Phospholyser [Bioblock, Illkirch, France]), and centrifuged at 48,000 × g for 1 h at 4°C. Nucleic acids were precipitated by the addition of spermin (0.2 M; 7% [vol/vol]; Sigma, Saint-Quentin Fallavier, France) for 1 h on ice. This suspension was ultracentrifuged at 100,000 × g for 1 h at 4°C.

Then, the  $\beta$ -lactamase extract from *E. coli* DH10B(pBS1) was loaded onto a preequilibrated Q-Sepharose column (Amersham Pharmacia Biotech). The  $\beta$ -lactamase was recovered in the flowthrough and was subsequently dialyzed overnight against 100 mM phosphate buffer (pH 7.0). The  $\beta$ -lactamase was loaded onto a preequilibrated S-Sepharose column (Amersham Pharmacia Biotech). The enzyme was eluted with a linear NaCl gradient (0 to 1 M) in phosphate buffer (pH 7). The  $\beta$ -lactamase was eluted with NaCl at a concentration of 320 to 350 mM. The fraction containing the  $\beta$ -lactamase activity was dialyzed overnight against 100 mM phosphate buffer (pH 7.0) prior to a 10-fold concentration with a Centrisart-C30 microcentrifuge filter (Sartorius, Goettingen, Germany).

**Isoelectric focusing.** Enzyme preparations from cultures of *C. meningosepticum* clinical isolates and *E. coli* DH10B(pBS1) were subjected to analytical isoelectric focusing on a pH 3.5 to 9.5 Ampholine polyacrylamide gel (Ampholine PAG plate; Amersham Pharmacia Biotech) for 90 min at 1,500 V, 50 mA, and 30 W. The focused  $\beta$ -lactamase was detected by overlaying the gel with 1 mM nitrocefin (Oxoid, Paris, France) in 100 mM phosphate buffer (pH 7.0). Since a metalloenzyme has previously been described in *C. meningosepticum* (36), detection of the pIs of the serine  $\beta$ -lactamases were additionally performed by incubating the enzyme extracts with 100  $\mu$ M clavulanic acid prior to their loading and focusing on an isoelectric focusing gel, followed by nitrocefin detection (28). The pI values were determined and compared to those of known  $\beta$ -lactamases run on the same gels.

**Kinetic measurements.** Purified β-lactamase was used for kinetic measurements ( $k_{cat}, K_m$ ), which were made at 30°C in 100 mM sodium phosphate (pH 7.0) with a Pharmacia ULTROSPEC 2000 spectrophotometer as described previously (17). Various concentrations of clavulanic acid, sulbactam, tazobactam, cefoxitin, imipenem, or moxalactam were preincubated with the enzyme for 3 min at 30°C before testing the rate of benzylpenicillin (100  $\mu$ M) hydrolysis. The 50% inhibitory concentrations (IC<sub>50</sub>s) of these inhibitors were determined. Results were expressed in micromolar units.

**Induction experiments.** To test the inducibility of the serine  $\beta$ -lactamase in *C.* meningosepticum clinical isolates, induction experiments were performed as described previously (31) with cefoxitin (10 µg/ml) or imipenem (1 µg/ml) as the inducer. One unit of enzyme activity was defined as that which was required to hydrolyze 1 µmol of aztreonam per min (aztreonam is not hydrolyzed by metalloenzymes). The total protein content was measured with bovine albumin as the standard (Bio-Rad DC Protein assay kit).

Determination of the β-lactamase relative molecular mass. The relative molecular mass of the β-lactamase from *E. coli* DH10B harboring pBS1 was estimated by sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis analysis. Enzyme extracts and marker proteins were boiled for 10 min in a 1% SDS-3% β-mercaptoethanol solution and were then subjected to electrophoresis on a 12% polyacrylamide gel (25 mA, 4 h) (16). Renaturation of the β-lactamase activity after denaturing electrophoresis and visualization of the β-lactamase on a benzylpenicillin-containing agar gel were performed as described previously (18).

Hybridization experiments. Southern hybridizations were performed with an ECL nonradioactive hybridization kit as described by the manufacturer (Amersham Pharmacia Biotech). Genomic DNA from C. meningosepticum PINT was hybridized with the following DNA probes corresponding to some  $\beta$ -lactamase genes: the 1.1-kb SnaBI fragment from recombinant plasmid pPZ1 for bla<sub>PER-1</sub>, the 450-bp PstI-NotI fragment from recombinant plasmid pHUC37 for blasHV-3 the 560-bp SspI-PstI fragment from plasmid pBR322 for bla<sub>TEM-1</sub>, the 329-bp DraI-XmnI fragment from recombinant plasmid pRLT1 for blavEB-1, the 396-bp EcoRI-PvuI from recombinant plasmid pPL1 for bla<sub>OXA-18</sub>, or the 811-bp PCRamplified fragment internal to  $bla_{CARB-2}$  (12). Two micrograms of denatured DNA was put onto a nylon membrane (Hybond N<sup>+</sup>; Amersham Pharmacia Biotech) and cross-linked with UV light for 2 min with a UV Stratalinker 2400 instrument (Stratagene). Membrane was incubated for 1 h at 60°C in a prehybridization buffer containing 100 µg of salmon sperm DNA per ml, 0.1% SDS, 5× SSC (0.75 M sodium chloride and 0.075 M sodium citrate), 5% dextran sulfate, and a 20-fold dilution of liquid block solution (supplied). Hybridizations were performed overnight at 60°C. Then, two washes were performed successively in the following solutions: 1× SSC–0.1% SDS for 15 min at 60°C and 0.5× SSC-0.1% SDS for 15 min at 60°C. The membrane was blocked and was then incubated with anti-fluorescein-horseradish peroxidase conjugate and finally washed. The signal was generated with a luminol solution and was detected with an autoradiographic film after 15 min of exposure.

Once the ESBL gene was identified, a set of primers was designed (primer 1, [5'-GTA GCT GTT TCT GTT TTG GGG-3'] and primer 2, [5'-CGA GAC CTG GGC AAT GAT TC-3'] at positions 438 to 458 and 1162 to 1181, respectively; (see Fig. 2) in order to attempt PCR amplification of this ESBL gene from the genomic DNAs of the *C. meningosepticum* isolates studied and from isolates of the related species studied (Table 1). In order to analyze the genetic organization of an ESBL-like gene in several *C. meningosepticum* strains, their corresponding genomic DNAs were digested with *Xmn*I and run on a 1% agarose gel. This gel was submitted to Southern hybridization as described above. Since *Xmn*I cut only once, in the middle of the ESBL gene (see below), *DraI-XmnI* and *DraI* restriction digests gave three internal fragments for the ESBL gene used as probes (fragments S1, S2, and S3; see Fig. 1).

Nucleotide sequence accession number. The nucleotide sequence data reported in this paper will appear in the GenBank nucleotide database under the accession no. AF033200.

#### RESULTS

Preliminary hybridization experiments and cloning of the ESBL gene. Genomic DNA from C. meningosepticum PINT failed to hybridize with several probes corresponding to class A  $\beta$ -lactamase genes ( $bla_{SHV-3}$ ,  $bla_{PER-1}$ ,  $bla_{VEB-1}$ ,  $bla_{TEM-1}$ , and bla<sub>CARB-2</sub>) or with a probe for the gene of the clavulanic acid-inhibited extended-spectrum oxacillinase OXA-18. Genomic DNA from C. meningosepticum PINT that had been partially digested with Sau3AI was cloned into the BamHI site of pBK-CMV. Sixteen recombinant E. coli DH10B clones harboring plasmids with inserts that varied in size (1.9 to 12 kb) were obtained after selection on amoxicillin- and kanamycincontaining TS agar plates. A schematic representation was generated for one of them, pBS1, which possesses the smallest insert (Fig. 1).



FIG. 1. Schematic representation of recombinant plasmid pBS1, which encodes the CME-2  $\beta$ -lactamase from *C. meningosepticum* PINT. The thin line represents the cloned insert from *C. meningosepticum* PINT containing the CME-2  $\beta$ -lactamase gene, with the arrow indicating its translational orientation (thick line), and the dotted lines indicate the vector pBK-CMV. The three internal probes (probes S1, S2, and S3) used for hybridization experiments are also represented by double-headed arrows.

β-Lactam resistance phenotype. Analysis of C. meningosepticum PINT by the conventional disk susceptibility assay suggested that its  $\beta$ -lactam resistance phenotype was partially due to the presence of an ESBL. A similar ESBL phenotype was found for E. coli DH10B harboring recombinant plasmid pBS1 (data not shown). This ESBL phenotype was peculiar since synergy was found not only between ceftazidime or cefotaxime and clavulanic acid or tazobactam but also with cefoxitin, moxalactam, or imipenem (data not shown). The MICs of β-lactams for C. meningosepticum PINT indicated that it was resistant to all tested β-lactams with the exception of ureidopenicillins such as piperacillin (Table 2). Similar MICs of  $\beta$ -lactams were obtained for all the C. meningosepticum isolates tested (data not shown). E. coli DH10B harboring recombinant plasmid pBS1 was resistant to the β-lactams tested but remained susceptible to cefepime, cephamycins, and carbapenems, therefore ruling out the role of the  $\beta$ -lactamase gene cloned into pBS1 in the carbapenem resistance of C. meningosepticum PINT. The MICs of the  $\beta$ -lactams were significantly lowered not only in the presence of clavulanic acid or tazobactam, as reported for a classical ESBL phenotype, but also in the presence of cefoxitin at concentrations as low as 0.1 µg/ml or of moxalactam and imipenem, both at 0.05 µg/ml.

Genetic and protein sequence analyses. Conjugation experiments failed to transfer any  $\beta$ -lactam resistance marker from *C. meningosepticum* PINT to rifampin-resistant *E. coli* JM109. However, no control was used in the assays for mating out from *C. meningosepticum* to *E. coli*. No plasmid was detected in *C. meningosepticum* PINT.

DNA sequence analysis of the 1,525-bp region downstream from the 1.9-kb insert of pBS1 revealed an open reading frame (ORF) of 891 bp encoding a 297-amino-acid preprotein (Fig. 2). No sequence typical of a class 1 integron was found within the insert. No ORF likely coding for a regulatory protein was found upstream of the  $\beta$ -lactamase gene.

The overall G+C content of the ORF was 33.9%, which lies close to the expected range of the G+C ratio of members of the *Chryseobacterium* (*Flavobacterium*) genus (37%) (34).

The mature protein (named CME-2 for *C. meningosepticum* ESBL) expressed in *E. coli* DH10B had a molecular mass of 31 kDa (data not shown). Within this protein sequence, seven boxes characteristic of  $\beta$ -lactamases possessing a serine-active site were found (14): box I, positions ABL 45 to 50; box II, positions ABL 70 to 73; box III, position ABL 105; box IV, position ABL 111; box V, position ABL 234 to 236 (Fig. 3). In

TABLE 2. MICs of β-lactams for *C. meningosepticum* PINT, *E. coli* DH10B harboring recombinant plasmid pBS1, and reference strain *E. coli* DH10B

	MIC (µg/ml)							
β-Lactam <sup>a</sup>	C. meningosepticum PINT	<i>E. coli</i> DH10B(pBS1)	E. coli DH10B					
Amoxicillin	256	512	4					
Amoxicillin + CLA	64	4	4					
Amoxicillin + TZB	64	4	2					
Amoxicillin + FOX	128	8	4					
Amoxicillin + MOX	64	16	4					
Amoxicillin + IPM	64	2	4					
Ticarcillin	256	512	4					
Ticarcillin + CLA	64	8	4					
Ticarcillin + TZB	128	4	4					
Ticarcillin + FOX	128	32	4					
Ticarcillin + MOX	128	64	4					
Ticarcillin + IPM	128	4	4					
Piperacillin	32	8	1					
Piperacillin + CLA	16	2	1					
Piperacillin + TZB	16	2	1					
Piperacillin + FOX	16	2	1					
Piperacillin + MOX	32	2	1					
Piperacillin + IPM	32	2	1					
Cephalothin	512	256	2					
Cefoxitin	32	4	1					
Ceftazidime	256	32	0.5					
Ceftazidime + CLA	16	0.5	0.5					
Ceftazidime + TZB	16	0.25	0.5					
Ceftazidime + FOX	16	1	0.5					
Ceftazidime + MOX	64	2	0.5					
Ceftazidime + IPM	64	0.5	0.5					
Cefotaxime	64	4	0.12					
Cefepime	32	0.5	0.03					
Cefepime + CLA	4	0.03	0.03					
Cefepime + TZB	4	0.03	0.03					
Cefepime + FOX	8	0.06	0.03					
Cefepime + MOX	16	0.06	0.03					
Cefepime + IPM	16	0.06	0.03					
Aztreonam	>512	16	0.25					
Aztreonam + CLA	>512	0.12	0.25					
Aztreonam + TZB	>512	0.25	0.25					
Aztreonam + FOX	>512	0.25	0.25					
Aztreonam + MOX	>512	0.5	0.25					
Aztreonam + IPM	>512	0.25	0.25					
Moxalactam	64	0.25	0.12					
Imipenem	32	0.25	0.12					

<sup>&</sup>lt;sup>*a*</sup> CLA, clavulanic acid at a fixed concentration of 2  $\mu$ g/ml; TZB, tazobactam at a fixed concentration of 4  $\mu$ g/ml; FOX, cefoxitin at a fixed concentration of 0.1  $\mu$ g/ml; IPM, imipenem at a fixed concentration of 0.5  $\mu$ g/ml; MOX, moxalactam at a fixed concentration of 0.05  $\mu$ g/ml.

addition, another class A conserved motif, serine-aspartic acidasparagine (SDN), was found at positions 130 to 132 (Fig. 3).

The comparison of CME-2 with other class A  $\beta$ -lactamases revealed weak identity (Fig. 3). The highest degrees of homology were with VEB-1 from *E. coli* MG-1, CBL-A from *B. uniformis*, and PER-1 from *P. aeruginosa* RNL-1 (39, 35, and 34% identities, respectively). A dendrogram was constructed in order to relate CME-2 to representative Ambler class A ESBLs. CME-2 clustered with VEB-1, CFX-A, and CEP-A (Fig. 4).

Biochemical properties of CME-2  $\beta$ -lactamase. Analytic isoelectric focusing revealed that *E. coli* DH10B harboring recombinant plasmid pBS1 produced only one  $\beta$ -lactamase with activity, and it had a pI of 9.2. In the original *C. meningosepticum* PINT isolate,  $\beta$ -lactamases with pI values of 7.6 and 8.3 were found. Only the  $\beta$ -lactamase with a pI of 7.6 corresponded to a serine  $\beta$ -lactamase.

After purification of the CME-2  $\beta$ -lactamase from *E. coli* DH10B(pBS1), the specific activity of 22 mU  $\cdot$  mg<sup>-1</sup> of protein was determined with 100  $\mu$ M benzylpenicillin as the substrate. The overall recovery of CME-2 was 41%, with a 120-fold  $\beta$ -lactamase purification. According to visual inspection of the SDS-polyacrylamide gel, the CME-2  $\beta$ -lactamase gene product was weakly expressed from pBS1 in *E. coli*.

The kinetic parameters of the purified CME-2  $\beta$ -lactamase revealed strong activity against cephalosporins, including extended-spectrum cephalosporins (Table 3), and its activity against aztreonam was also noticeable. CME-2  $\beta$ -lactamase has no detectable activity against piperacillin, cephamycins, and imipenem (Table 3). IC<sub>50</sub> results with benzylpenicillin as the substrate showed that CME-2 activity was strongly inhibited by clavulanic acid (0.05  $\mu$ M), tazobactam (1.5  $\mu$ M), and sulbactam (0.3  $\mu$ M). In addition, unlike other class A ESBLs, CME-2 was inhibited more strongly by cefoxitin (0.004  $\mu$ M), moxalactam (0.002  $\mu$ M), and imipenem (0.015  $\mu$ M).

Distribution of *bla*<sub>CME-2</sub> in *Chryseobacterium* sp. isolates. Using a set of primers designed to amplify a 743-bp internal fragment of  $bla_{CME-2}$  by PCR, positive results were obtained for only five of nine C. meningosepticum strains (C. meningosepticum PINT, CIP 6058, AMA, GEO, and CIP 6059). Negative PCR results were obtained not only for four of nine C. meningosepticum isolates but also for all strains of the C. meningosepticum-related species (Chryseobacterium indologenes, Sphingobacterium multivorum, and Myroides odoratus) (data not shown). However, hybridization experiments with a  $bla_{CME-2}$ -specific internal probe gave positive results for the nine C. meningosepticum isolates and negative results for the strains of the C. meningosepticum-related species. In order to evaluate the distribution of a  $bla_{CME-2}$ -like gene within C. meningosepticum isolates, their DNAs were restricted with *Xmn*I and hybridized with three probes, probes S1 and S2, which are located upstream and downstream from the XmnI site (*Xmn*I cuts in the middle of the  $bla_{CME-2}$  gene), respectively, and S3, a DraI internal fragment, giving a probe specific for the entire  $bla_{CME-2}$  gene (Fig. 1). Six hybridization patterns were obtained for the nine C. meningosepticum isolates (Fig. 5). By using the S1 and S2 probes, C. meningosepticum PINT, CIP 6058, and AMA (pattern I) gave 3- and 1.8-kb fragments, respectively; C. meningosepticum GEO and CIP 6059 (pattern II) gave 2.7- and 1.8-kb fragments, respectively; C. meningosepticum CIP 7830 (pattern III) gave 3.5- and 1.9-kb fragments, respectively; C. meningosepticum CIP 7905 (pattern IV) gave 3.5- and 1.8-kb fragments, respectively; C. meningosepticum AB 1572 (pattern V) gave 3.7-kb fragments with both probes; and C. meningosepticum H01J100 (pattern VI) gave 3.8-kb fragments with both probes. S3 probe hybridizations gave patterns corresponding to the sum of the results obtained after the S1 and S2 probe hybridizations. Only patterns I and II gave positive results after *bla*<sub>CME-2</sub> PCR amplification.

Finally, induction studies with imipenem or cefoxitin as the inducer did not show that these ESBLs from *C. meningosepticum* would be inducible. As assessed by isoelectric focusing analysis, each isolate produced at least two  $\beta$ -lactamases. However, preincubation of the enzyme extracts with clavulanic acid showed that each *C. meningosepticum* isolate produced only one serine  $\beta$ -lactamase with variable pIs ranging from 7.0 to 8.5, with no correlation of the pIs with the hybridization patterns.

1																		G	SATCO	stgaa
10	CAGO	TTG	ACAGI	ACTTO	CAAA	AGAT	TATI	TTAC	CCG	rttC <i>i</i>	ATT	TCCI	'AAGI	AGCTI	ATGTO	CAAA/	GAC	AGGI	GATO	GAAAG
90	CCAT	TCTI	AATAC	CAAC	ACAC	CATTI	TGT	ATCA	TGAI	AAGAT	rGGG <i>I</i>	ACTGO	'AAAI	GATT	TTAC	TAC	GAA	CATO	TCTT	GTTA
170	ATA	28887	AAATI	ATA	TTAT	GAA/	AATO	CATTI	TAAC	AAAA	GTGT/	GAAA	ACTI	ATGTZ	AAG	/GCC1	GAA	CTGGI	AACO	TGCT
250	TTA	\TAA#	AAGGI	TATAC	CGGI	FAAAT	TCTI	TATAT	GAA	rgtt <i>i</i>	ACTAT	TTAT	TAT	GATTI	ATTC	AAAA	ACA	"AAA"	TAG	TGAA
330	ATG	AAA	AAA	ATT	ATA	CTC	CTA	TTT	ATC	TTG	ACA	AGC	CAG	TTG	GTG	CTT	GCT	CAA	CAT	ACT
1	M	K	K	I	I	L	L	F	I	L	T	S	Q	L	V	L	A	Q	H	T
390	TCA	ATA	TTA	AAT	GAT	ATA	AAT	GCT	GTT	ACC	AAA	GAC	AAG	AAA	GCC	ACA	GTA	GCT	GTT	TCT
21	S	I	L	N	D	I	N	A	V	T	K	D	K	K	A	T	V	A	V	S
450	GTT	TTG	GGG	ATA	GAA	AAT	GAT	TTT	CAG	TTT	AGT	AAC	GCC	AAT	GGT	AAT	TTG	AAA	ATG	CCG
41	V	L	G	I	E	N	D	F	Q	F	S	N	A	N	G	N	L	K	M	P
510	ATG	CTG	AGT	GTT	TTT	AAA	TTT	САТ	ATT	GCA	TTG	GCG	GTT	CTA	AAT	CAG	GTA	GAC	AAA	GGT
61	M	L	S	V	F	K	F	Н	I	A	L	A	V	L	N	Q	V	D	K	G
570	AAC	CTT	ACC	TTG	GAT	CAG	AAA	ATA	CTG	ATT	AAA	AAA	TCG	GAT	CTA	TTA	GAA	AAT	aca	TGG
81	N	L	T	L	D	Q	K	I	L	I	K	K	S	D	L	L	E	N	T	W
630	TCA	CCA	CTT	CGT	GAG	AAG	TAT	CCG	GAT	GGA	AAT	GTA	GAA	CTT	ССТ	TTA	AGC	GAA	ATT	ATT
101	S	P	L	R	E	K	Y	P	D	G	N	V	E	L	Р	L	S	E	I	I
690	ACT	TAT	ACT	GTA	GCC	CAA	AGT	GAC	AAC	AAC	GGA	TGC	GAC	ATA	CTA	TTA	AGG	CTA	ATT	GGC
121	T	Y	T	V	A	Q [	S	D	N	N	G	C	D	I	L	L	R	L	I	G
750	GGG	АСТ	AAA	ACT	GTT	CAG	AAA	TTA	ATG	GAT	GTG	AAT	GGT	GTA	AAA	AAC	TTT	CAG	ATA	AAA
141	G	Т	K	T	V	Q	K	L	M	D	V	N	G	V	K	N	F	Q	I	K
810 161	TAT Y P	AAT	GAG	GAA E I	GAA S N	ATG I H	CAT I F	AAA	AAT I I	GAT C C	GTA 7 H	AAA T	ACT I	CTT	TAT ( 1	GCA	AAT I Y	TAC ( 1	ACA	ACT
870	ACA	GCA	TCT	ATG	GTA	AAA	АСТ	CTG	AAA	GCG	TTC	TAT	AAA	GGA	ATG	TTT	TTA	TCA	AAA	AGA
181	T	A	S	M	V	K	Т	L	K	A	F	Y	K	G	M	F	L	S	K	R
930	тсс	ACA	ATT	TTT	CTA	ATG	GAT	ATT	ATG	ACT	AAA	ACC	AAT	ACC	GGA	ATG	TCA	AAG	CTT	CCG
201	s	T	I	F	L	M	D	I	M	T	K	T	N	T	G	M	S	K	L	P
990	GGC	TTG	CTG	CCA	AAA	GTT	AGA	ATG	GCC	AGA	AAA	ACA	GGT	TCT	TCG	GGT	AAA	ATG	AAA	AAC
221	G	L	L	P	K	V	R	M	A	R	K	T	G	S	S	G	K	M	K	N
050	GGA	TTA	ACG	ATT	GCT	GAG	AAC	GAT	TCA	GGA	ATT	GTA	ACT	TTA	GCA	AAT	GGT	AAA	САТ	ТАТ
241	G	L	T	I	A	E	N	D	S	G	I	V	T	L	A	N	G	K	Н	Ү
110	GCA	ATT	GCA	GTA	TTT	GTA	AAG	GAC	TCT	ATG	GAA	AGT	GAG	GAA	GTC	AAT	TGT	GGA	ATC	ATT
261	A	I	A	V	F	V	K	D	S	M	E	S	E	E	V	N	C	G	I	I
170 281	GCC A	CAG Q	GTC V	TCG S	AAA K	ATT I	GTC V	TGG W	GAT D	GCT A	TTA L	AAT N	AAA K	AAA K	AAT N	AAA K	CCC P	<u>taa</u>	TAAT.	AAAAC
233	CGA	AGGI	TCG	TAAAT	TCTI	TAAT	CAAA	LAAAI	'AAA/	AATT	ATG	ACAA	GAA	PATTO	TAGI	AATI	GGTO	CGGG	GAACO	ATGG
313	GTA	LCGGA	ATTO	SCTC	TACT	PTTTC	SCCC2	AACC	GGAI	TTA <i>P</i>	AGTI	AATT	TAG	TAGAT	GTTI	CACA	GGA	AGCGC	TGGA	CAGA
393	GGT	TTA	AACT	CATT?	ACCAC	TAAT	CTGG	ACAG	GAATI	TATTO	CTA	AGGA	AATO	TTAC	AGAA	GAAG	AGA	AGCC	AATA	CACT
473	AGC	AATA	ATCTO	TACI	TTTT <i>I</i>	ACAGO	TTTO	GAAG	ATGO	CTGTI	AAA	GATGC	CGAT	rc 1	525					

FIG. 2. Nucleotide sequence of the 1,525-bp fragment of pBS1 containing the 891-bp coding region for  $bla_{CME-2}$ . The deduced amino acid sequence is designated in single-letter code below the nucleotide sequence. The seven conserved boxes described by Joris et al. (14) and the conserved SDN loop are boxed. The start and stop codons of the gene are in boldface type and are underlined.

## DISCUSSION

1

1

1

The starting point of this study was the observation of a synergy between ceftazidime and cephamycins in a conventional antibiogram disk susceptibility assay performed with a *C. meningosepticum* clinical isolate. The overall  $\beta$ -lactam susceptibility pattern of this *C. meningosepticum* isolate corresponded to that previously described in these bacterial species characterized by resistance to all  $\beta$ -lactams tested, including extended-spectrum cephalosporins, monobactams, and carbapenems, and a moderated susceptibility to piperacillin (6, 20). This profile of resistance to broad-spectrum  $\beta$ -lactams may be explained for all  $\beta$ -lactams except cefepime and imipenem by the hydrolytic properties of the ESBL CME-2. For cefepime and

imipenem the resistance is most likely mediated by the class B carbapenem-hydrolyzing  $\beta$ -lactamase recently identified in *C. meningosepticum* CIP 6058, a strain included in this study (36). The biochemical properties of CME-2 are similar to those of an unsequenced ESBL identified from another *C. meningosepticum* isolate (9). Both enzymes may be classified within Bush group 2be (5). This group includes several  $\beta$ -lactamases which are weakly related to CME-2, i.e., MEN-1, TOHO-1, and L-2. CME-2 activity not only was inhibited by clavulanic acid but was also strongly inhibited by cephamycins and imipenem. This inhibition profile is found in two other class A  $\beta$ -lactamases, VEB-1 from *E. coli* and CEP-A from *B. fragilis* (35). Comparison of their amino acid sequences did not provide evidence of

	1	10	20	D	30		40	50	60	69 70	80
ጥጅΜ-3	MSTOHFR	 VAT.TP	FFAAFCI.P	VFAHPE	 		KLGARV	GYTELDUNS	GKTLESFRPE	[ אידיד 1988 אידיד אידי	VLLCGAV
SHV-2	MRYIR	LCIIS	-LLATLPLA	VHASPQ	PLE	QIKLSES	QLSGRV	GMIEMDLAS	GRTLTAWRAD	ERFPMMSTFK	VVLCGAV
MEN-1	-MVKKSLRQFT	LMATAT	VTLLLGSVP	LYAQTA	DVQQ	KLAELER	QSGGRL	GVALINTAL	NSQIL-YRAD	erfamostsk	VMAVAAV
TOHO-1	MMTQSIRRSM	LTVMAT	LPLLFSSATI	lhaqan	SVQQ	QLEALEK	SSGGRL	GVALINTAI	NSQIL-YRAD	erfamostsk	VMAAAAV
NMC-A	MSLNVKQSR	IAILFS	SCLISISFF	SQANTK	GID	EIKNLET	DFNGRI	GVYALDIGS	GKSFS-YRAN	ERFPLOSSFK	GFLAAAV
L-2	MLARRELOFS	GAAVASSLA	LPLLARAAG	KTAASA	PTDAALT.	AATDFAALEK	AVRGRE	GVTLLDTAS	GRRIG-HRQD	ERFPMOSTFK	SVLAATV
CEP-A	MUKKLIH	LSI15 TATLT	FLLCPAL	VVAQNS	PLET	QLKKAIE	GKKAEL		GODTITVNND	VHEDLLSVER	FHUALAL FHUALAU
CFX-A	MEKNRKKO	IVVLSIALV	CIFILVESL	FHKSATKDS	ANPPLTNV		ACPGEI	GVAVIVN	NRDTVKVNNK	SVYPMMSVFK	VHOALAL
PER-1	MNVIII	KAVVTAS	TLLMVSFSS	FETSAQS	PLLK	EQIESIVI	GKKATV	GVAVWGPDI	LEPLL-INPF	EKFPMQSVFK	LHLAMLV
VEB-1	MKIVKRIL	LVLLS	LFFTIVY	SNAQTD	NLTL	KIENVLK	AKNARI	GVAIFNS-N	EKDTLKINND	FHFPMQSVMK	FPIALAV
CME-2	MKKII	LLFIW	TSQL	VLAQHT	SILN	DINAVTK	DKKATV	AVSVLGIEN	IDFQFSNANGN	LKMPMLSVFK	FHIALAV
								box I		box I	I
	90	1	00 104	110	120	130	135 1	40	150	160	170
	l l	Ī	1	1	Ī	Ī	I I		1	I	1
TEM-3	LSRVDAGQEQL	GRRIHYSON	IDLVKTSP	VTERHLTD-	-GMTVRELCS.	AAITMSDNTA	ANLLLT	-TIGGPKEI	TAFLHNMGDH	V-TRLDRWEP	ELNE-AI
SHV-2	LARVDAGDEQL	ERKIHYRQC	DLVDYSP	VSEKHLAD-	-GMTVGELCA	AAITMSDNSA	ANLLLA	-TVGGPAGI	TAFLRQIGDN	V-TRLDRWEI	ELNE-AL
MEN-1	LKKSESEPNLL	NORVEIKKS	SDLVNYNP	IAEKHVDG-	-TMSLAELSA	AALQYSDNVA	MNKLIS	-HVGGPAS	TAFARQLGDE	T-FRLDRTEF	TLNT-AI
TOHO-1	LKQSESDKHLL	NORVEIKKS	SDLVNYNP	IAEKHVNG-	-TMTLAELGA	AALQYSDNTA	MNKLIA	-HLGGPDK	TAPARSLGDE	T~FRLDRIEF	TLNT-AL
L-2	LSOAFROPALL	NUTRVPVRDA	DLLS-HAP	VTRRHAGK-	-DMTVRDLCR	ATITTSDNGA	ANLIFG	-VVGGPPAL	TAFLESIGDE	V-SRTDRIFF	ELNS-FA
CEP-A	ADYMHHOKOPL	ETRLLIKKS	DLKPDTYSP	LREITYPOGO	IEMSIADLLK	TLOOSDNNA	CDILFN	-YOGGPDAV	NKYLHSLGIR	E-CAVINTEN	DMHK-NL
CBL-A	LDKMDKQSISL	DSIVSIKAS	SOMPPNTYSPI	LRKKFPDQI	FTITLRELMQ	YSISQSDNNA	CDILIE	-YAGGIKHI	NDYIHRLSID	S-FNLSETEL	GMHS-SF
CFX-A	CNDFDNKGISL	DTLVNINRI	KLDPKTWSPI	MLKDYSGPV	ISLTVRDLLR	YTLTQSDNNA	SNLMFK	-DMVNVAQ1	DSFIATLIPR	SSFQIAYTE	EMSA-DH
PER-1	LHQVDQGKLDL	NQTVIVNRA	KVLONTWAP:	IMKAYQGDE	FSVPVQQLLQ	YSVSHSDNVA	CDLLFE	-LVGGPAAI	HDYIQSMGIK	E-TAVVANEA	QMHA-DD
VEB-1	LSEIDKGNLSF	EQKIEITPO	DLLPKTWSP	IKEEFPNG-	TTLTIEQILN	YTVSESDNIG	CDILLK	-LIGGTDSV	QKFLNANHFT	D-ISIKANEE	OMHK-DW
CME-2	LNQVDKGNLTL	DÖKITTIKKS	bor III	box IV	VELPLSEIIT	TVAQSDNNG	CDIFFR	-LIGGINIV	QKLMDVNGVK	N-LOIVINGC	V
			<i></i>	<i></i>							•
	180	190	20	00	210	220 I	230 I	238 I	240 244	250 I	260 I
TEM-3	PNDERDTTMPA	AMATTLRKI	LTG-~ELLT	-LASROOLI	DWMEADKVA-	GPLLRSALPA	G-WFIA	DKSGAS	É-RGSRGI	IAALGP-DGK	PSRIVVI
SHV-2	PGDARDTTTPA	SMAATLRKI	LTSQRLS	-ARSQRQLI	QWMVDDRVA-	GPLIRSVLPA	G-WFIA	IKTGAS	E-RGARGI	VALLGP-NNK	AERIVVI
MEN-1	PGDPRDTTSPR	AMAQTLRNI	TLGKALG	-DSQRAQLV	TWMKGNTTG-	AASIQAGLPA	S-WVVG	TKTGSG	D-YGTTND	IAVIWP-KDR	APLILVT
тоно-1	PGDPRDTTTPL	AMAQTLKNI	TLGKALA	-ETQRAQLV	TWLKGNTTG-	SASIRAGLPK	S-WVVG	DKTGSG	D-YGTTND	IAVIWP-ENH	APLVLVT
NMC-A	PGDERDTSTPA	AVAKSLKTI	LALGNILS	-EHEKETYC	TWLKGNTTG-	AARIRASVPS	D-WVVG	DKTGSCG	A-YGTAND	YAVVWP-KNE	APLIISV
11-2 CFP_3	RGDPRDTTTPA FFCVONWTTPT.	AMAATLQKV	VLGLVLQ.	-LASKQQLA		DACTKAGTC	.⊔-wr.vr ж_улмс	HKTGENG	JAKGOOTGOND	TGFTLL_PDG	HAVSTAV
CBL-A	EAVYRNWSTPS	AMVRLLRTA	ADEKELFS	NKELKDFLV	OTMIDTETG-	ANKLKGMLPA	K-TVVG	HKTGSSDRI	ADGMKTADND	AGLVIL-PDG	RKYYIAA
CFX-A	NKAYSNYTSPL	GAAMLMNRI	FTEGLID	-DEKQSFIF	NTLKECKTGV	DRIAAPLLDK	EGVVIA	HKTGSGYVI	<b>IENGVLAAHND</b>	VAYICL-PNN	ISYTLAV
PER-1	QVQYQNWTSMK	GAAEILKKF	EQKTQLS	-ETSQALLW	KWMVETTTG-	PERLKGLLPA	G-TVVA	HKTGTSG-1	KAGKTAATND	LGIILL-PDO	RPLLVAV
VEB-1	NTQYQNWATPT.	AMNKLLIDT	YNNKNQLLS	-KKSYDFIV	KIMRETTTG-	SNRLKGQLPK	N-TIVA	HKTGTSGIN	I-NGIAAATND	VGVITL-PNG	QLIFISV
CME-2	KTLYANYTTTA	SMVKTLKAP	YKGMFLS	-KRSTIFLN	DEMTKTNTG-	MSKLPGLLPK	VRMA	NKTGSSGKA	KNGLTIAEND	SGIVTL-ANG	KHYATAV
				•			~				
	270	276 280	290								
ጥጽሐ ን	 VTTCCC27M7787	RNROTAFTO	AST.TKHW								
SHV-2	YLRDTPASMAE	RNOOIAGIG	JAALIEHWOR								
MEN-1	YFTOPOPKAES	RRDVLASAA	KIVTNGL								
TOHO-1	YFTQPEQKAER	RRDILAAAA	KIVTHGF								
NMC-A	YTTKNEKEAKH	EDKVIAEAS	SRIAIDNLK-								
L-2	YLQAGAISYEQ	RATVLAQVO	FRIADRLIG-								
CEP-A	FUNDSEADNRE	NSELIAEIS	SKIVYEYVTQ	QTD							
CFX-A	FVRDFRCNKSO	NAN LIARIS	SMIVIDAMK-	TSVKS							
PER-1	* ATOT LOUTOD	A.S.( ) 7 0 4 4 1 1 1									
*	FVKDSAESSET	NEALLAOVA	OTAYOFELK	KLSALSPN							
VEB-1	FVKDSAESSRT FVAESKETSEI	NEAIIAQVA NEAIISDIA	AQTAYQFELK AKITWNYYLN	KLSALSPN K							
VEB-1 CME-2	FVKDSAESSRT FVAESKETSEI FVKDSMESEEV	NEAIIAQVA NEKIISDIA NCGIIAQVS	AQTAYQFELK AKITWNYYLN SKIVWDALNK	KLSALSPN K KNKP							

FIG. 3. Amino acid sequence alignment of CME-2 compared with those of selected class A  $\beta$ -lactamases. The numbering is according to Ambler (1). Roman numerals designate the boxes described by Joris et al. (14): box I, position ABL 45 to 50; box II, positions ABL 70 to 73; box III, position ABL 105; box IV, position ABL 111; box V, position ABL 166; box VI, position ABL 210; box VII, positions ABL 234 to 236. The  $\beta$ -lactamases included in the alignment are TEM-3 from *K. pneumoniae* CFF104, SHV-2 from *K. ozaenae*, MEN-1 from *E. coli* MEN, TOHO-1 from *E. coli* TUH12191, NMC-A from *E. cloacae* NOR-1, L-2 from *S. maltophilia* 1275IID, CEP-A from *B. trigilis* CS30, CFX-A from *B. vulgatus* CLA341, VEB-1 from *E. coli* MG-1, CBL-A from *B. uniformis* WAL-7088, and PER-1 from *P. aeruginosa* RNL-1. Dashes indicate gaps within the alignment.

any particular amino acid identity which may explain this cephamycin inhibition property. This property may help to differentiate *C. meningosepticum* from other *Chryseobacterium* sp. isolates for which a synergy between cephamycins and ceftazidime is not observed.

CME-2 is only distantly related to other class A ESBLs.

However, it belongs to a subgroup of class A  $\beta$ -lactamases, of which VEB-1 from *E. coli* MG-1 is the most closely related to CME-2 (39% amino acid identity).

Although isolated from *C. meningosepticum*, the CME-2  $\beta$ -lactamase does not exhibit features common to class A  $\beta$ -lactamases isolated from some gram-negative species, such as



FIG. 4. Dendrograms obtained for 12 Ambler class A ESBLs by the parsimony method (44). Branch lengths are drawn to scale and are proportional to the number of amino acid changes. The percentages at the branching points (underlined) refer to the number of times that a particular node was found in 100 bootstrap replications (the asterisks indicate uncertainty for nodes with bootstrap values of less than 50%). The distance along the vertical axis has no significance. Abbreviations for  $\beta$ -lactamases are given in the legend to Fig. 3. Percent amino acid identities to CME-2 are indicated in parentheses.

Cys77 and Cys123, which are thought to form a disulfide bridge (33, 38, 46). No cysteine residue is found at these positions in CME-2; instead, an alanine (as for VEB-1, PER-1, CFX-A, CBL-A, CEP-A, L-2, NMC-A, and TOHO-1) and an isoleucine (a leucine in VEB-1, PER-1, CFX-A, and CEP-A), respectively, are found (Fig. 3). Surprisingly, CME-2 possesses a cysteine residue at position 135, close to the SDN motif, as in VEB-1, PER-1, CBL-A, and CEP-A. It is interesting that within the non-TEM and non-SHV class A  $\beta$ -lactamases, cysteine residues are found close to either the SVFK or the SDN conserved motifs at a limited number of positions; position 69 (MEN-1, TOHO-1, NMC-A, and L-2), position 81 (CFX-A), position 123 (L-2), or position 135 (CEPA, CBL-A, PER-1,

TABLE 3. Kinetic parameters of the purified β-lactamase CME-2

Drug	$k_{\text{cat}}$ $(s^{-1})$	$K_m$ ( $\mu$ M)	$k_{ ext{cat}}/K_m \ (\mu \mathrm{M}^{-1} \cdot \mathrm{s}^{-1})$
Cephaloridine	30.6	23.4	1.31
Benzylpenicillin	12.3	7.4	1.66
Amoxicillin	6.7	26.5	0.26
Ticarcillin	1.8	5.8	0.31
Piperacillin	0.3	<1	<u>a</u>
Cephalothin	60.6	41.5	1.46
Cefoxitin	< 0.01		_
Cefuroxime	57.3	81.6	0.70
Cefoperazone	2.6	3.8	0.68
Cefotaxime	12.1	20.8	0.58
Ceftazidime	23.6	125	0.19
Aztreonam	3.2	10.5	0.31
Imipenem	< 0.01	_	_

<sup>a</sup> -, not detectable.



FIG. 5. Autoradiogram after Southern hybridizations of *C. meningosepticum* genomic DNAs. The DNAs were restricted with *Xmn*I, and the probe consisted of the 372-bp *DraI-XmnI* fragment (S1) of pBS1 (A), the 306-bp *XmII-DraI* fragment (S2) of pBS1 (B), or the 667-bp *DraI* fragment (S3) of pBS1 (C) (see Fig. 1 for the locations of S1, S2, and S3). Lanes: 1, *C. meningosepticum* PINT; 2, *C. meningosepticum* CIP 6058; 3, *C. meningosepticum* AMA; 4, *C. meningosepticum* GEO; 5, *C. meningosepticum* CIP 7830; 6, *C. meningosepticum* CIP 6059; 7, *C. meningosepticum* CIP 7905; 8, *C. meningosepticum* AB 1572; 9, *C. meningosepticum* H01J100; 10, *E. coli* DH10B (negative control). The sizes of the DNA fragments may be deduced from the scale (1.5 to 4 kb) shown on the left sides of the gels.

VEB-1, and CME-2). In CME-2, a disulfide bridge may be formed between cysteine 135 and cysteine 276, as evidenced for NMC-A between cysteine 69 and cysteine 238 (43).

The omega loop which goes from positions 169 to 179 is a structural element of class A enzymes. This loop is present in CME-2 but is totally different from those found in TEM or SHV derivatives. In this respect, CME-2 along with VEB-1, PER-1, CBL-A, and CEP-A possesses a histidine in place of an asparagine at position 170 (Fig. 3). This asparagine together with the highly conserved Glu166 and Ser70 is involved in the positioning of the active-site water molecule (19). It is tempting to speculate that this histidine may play a similar role in the catalytic properties of CME-2, but this needs to be confirmed by site-directed mutagenesis.

Many ESBLs are derived from the parental TEM-1, TEM-2, or SHV-1 enzymes by a few amino acid substitutions at position 104, 164, 238, or 240, leading to increased catalytic activities for cefotaxime, ceftazidime, and aztreonam. In the CME-2  $\beta$ -lactamase, two additional amino acids are found at position 104 compared to those found in TEM-1, TEM-2, or SHV-1 derivatives. The Arg104Lys substitution in TEM-3 may correspond to either a glutamic acid, an asparagine, or a threonine in CME-2 (Fig. 3). In the PER-1  $\beta$ -lactamase, additional amino acids (glutamine, asparagine, and threonine) are also found at position 104. PER-1 is the only non-TEM and non-SHV ESBL for which site-directed mutagenesis has so far been performed, but the roles of these amino acids at this position remain unclear (4).

At position 164, some TEM derivatives that possess ESBL properties have a serine or a histidine in place of an arginine. PER-1 possesses an alanine and CME-2 possesses a tyrosine.

Since the Ala164Arg substitution in PER-1 results in a mutant with no detectable activity (4), it would be interesting to study the effect of the Tyr164Arg substitution in CME-2. At positions 238 and 240, TEM-1, TEM-2, and SHV-1 possess glycine and glutamine residues, respectively. CME-2, like PER-1, possesses a serine at position 238, as found for some TEM and SHV ESBLs, and a glycine at position 240.

The G+C ratio of 33.9%, typical of Chryseobacterium species genes, together with negative conjugation and negative plasmid research results, indicated a likely chromosomal location of the  $bla_{CME-2}$  gene. No sequence typical of E. coli or P. aeruginosa promoters was identified upstream of the ATG initiation site. Interestingly, the pI value for CME-2 in E. coli DH10B (9.2) did not correspond to the pI value of the serine  $\beta$ -lactamase (7.6) found in C. meningosepticum PINT, thus indicating a putative difference in the cleavage site of the peptide leader. No regulatory protein gene was found immediately upstream of the  $bla_{CME-2}$  gene. Therefore, CME-2 expression may be either not regulated at all or not regulated like other chromosomally located class A ESBLs such as NMC-A and SME-1 (21, 22). In that respect, induction experiments failed to identify any inducible serine ESBL in C. meningosepticum isolates, whereas inducible L-2 enzymes are found in S. maltophilia (27). However, as found in S. maltophilia (28), several β-lactamases of Ambler class A and class B were identified in each C. meningosepticum isolate (data partially shown), and these may together account for the naturally occurring  $\beta$ -lactam resistance profiles of the *C. meningosepticum* isolates.

The results of PCR with nondegenerated primers for  $bla_{\rm CME-2}$  detection showed that this technique may be too specific for the identification of  $bla_{\rm CME-2}$ -like genes. However, hybridization experiments identified in each *C. meningosepticum* isolate a  $bla_{\rm CME-2}$ -like gene that was present in only one copy. Moreover, they revealed that  $bla_{\rm CME-2}$  gene variants are identified in all *C. meningosepticum* isolates tested, although their  $\beta$ -lactam resistance phenotypes were identical. The presence of a  $bla_{\rm CME-2}$ -like gene in *C. meningosepticum* isolates makes it an identification marker for this bacterial species among *Chryseobacterium* species. Finally, the identification of CME-2  $\beta$ -lactam resistance places an additional clue that class A ESBLs may be a means for naturally occurring  $\beta$ -lactam resistance.

## ACKNOWLEDGMENTS

This work was financed by a grant from the Ministère de l'Education Nationale et de la Recherche (grant UPRES-JE 2227), Université Paris XI, Paris, France.

We thank Esthel Ronco and Brita Bruun for providing us some *C. meningosepticum* clinical isolates and D. Aubert for help in determining kinetic constants.

## **ADDENDUM**

After the work was submitted, we noticed a report from Rossolini et al. (36) showing a very similar enzyme, CME-1, from *C. meningosepticum* CCUG4310. CME-1 displays five amino acid changes with respect to CME-2: valine in CME-1 to isoleucine in CME-2 at position 157, isoleucine to methionine at position 278, and asparagine to lysine at position 295 and deletion of the last two amino acids, lysine and proline, of CME-1 in CME-2. This report explains why the name CME-2 was retained. None of the amino acid differences may be critical to the extended hydrolysis spectrum profile, which was found to be similar in both enzymes.

### REFERENCES

- Ambler, R. P. 1980. The structure of beta-lactamases. Philos. Trans. R. Soc. Lond. B Biol. Sci. 289:321–331.
- Barthélémy, M., J. Péduzzi, H. Bernard, C. Tancrède, and R. Labia. 1992. Close amino-acid sequence relationship between the new plasmid-mediated extended-spectrum beta-lactamase MEN-1 and chromosomally encoded enzymes of *Klebsiella oxytoca*. Biochim. Biophys. Acta 1122:15–22.
- Bloch, K. C., R. Nadarajah, and R. Jacobs. 1997. Chryseobacterium meningosepticum: an emerging pathogen among immunocompromised adults. Report of 6 cases and literature review. Medicine (Baltimore) 76:30–41.
- Bouthors, A. T., N. Dagoneau-Blanchard, T. Naas, P. Nordmann, V. Jarlier, and W. Sougakoff. 1998. Role of residues 104, 164, 166, 238 and 240 in the substrate profile of PER-1 beta-lactamase hydrolysing third-generation cephalosporins. Biochem. J. 330:1443–1449.
- Bush, K., G. A. Jacoby, and A. A. Medeiros. 1995. A functional classification scheme for beta-lactamases and its correlation with molecular structure. Antimicrob. Agents Chemother. 39:1211–1233.
- Chang, J. C., P. R. Hsueh, J. J. Wu, S. W. Ho, W. C. Hsieh, and K. T. Luh. 1997. Antimicrobial susceptibility of flavobacteria as determined by agar dilution and disk diffusion methods. Antimicrob. Agents Chemother. 41: 1301–1306.
- Colding, H., J. Bangsborg, N. E. Fiehn, T. Bennekov, and B. Bruun. 1994. Ribotyping for differentiating *Flavobacterium meningosepticum* isolates from clinical and environmental sources. J. Clin. Microbiol. 32:501–505.
- Fraser, S. L., and J. H. Jorgensen. 1997. Reappraisal of the antimicrobial susceptibilities of *Chryseobacterium* and *Flavobacterium* species and methods for reliable susceptibility testing. Antimicrob. Agents Chemother. 41:2738– 2741.
- Fujii, T., K. Sato, E. Yokota, T. Maejima, M. Inoue, and S. Mitsuhashi. 1988. Properties of a broad spectrum beta-lactamase isolated from *Flavobacterium* meningosepticum GN14059. J. Antibiot. 41:81–85.
- Hansen, J. B., and R. H. Olsen. 1978. Isolation of large bacterial plasmids and characterization of the P2 incompatibility group plasmids pMG1 and pMG5. J. Bacteriol. 135:227–238.
- Huletsky, A., F. Couture, and R. C. Levesque. 1990. Nucleotide sequence and phylogeny of SHV-2 beta-lactamase. Antimicrob. Agents Chemother. 34: 1725–1732.
- Huovinen, P., and G. A. Jacoby. 1991. Sequence of the PSE-1 beta-lactamase gene. Antimicrob. Agents Chemother. 35:2428–2430.
- İshii, Y., A. Ohno, H. Taguchi, S. Imajo, M. Ishiguro, and H. Matsuzawa. 1995. Cloning and sequence of the gene encoding a cefotaxime-hydrolyzing class A beta-lactamase isolated from *Escherichia coli*. Antimicrob. Agents Chemother. 39:2269–2275.
- Joris, B., P. Ledent, O. Dideberg, E. Fonze, J. Lamotte-Brasseur, J. A. Kelly, J. M. Ghuysen, and J. M. Frère. 1991. Comparison of the sequences of class A beta-lactamases and of the secondary structure elements of penicillinrecognizing proteins. Antimicrob. Agents Chemother. 35:2294–2301.
- Kado, C. I., and S. T. Liu. 1981. Rapid procedure for detection and isolation of large and small plasmids. J. Bacteriol. 145:1365–1373.
- Laemmli, U. K., and M. Favre. 1973. Maturation of the head of bacteriophage T4. I. DNA packaging events. J. Mol. Biol. 80:575–599.
- Laurent, F., L. Poirel, T. Naas, E. B. Chaibi, R. Labia, P. Boiron, and P. Nordmann. 1999. Biochemical-genetic analysis and distributions of FAR-1, a class A β-lactamase from *Nocardia farcinica*. Antimicrob. Agents Chemother. 43:1643–1650.
- Massida, O., G. M. Rossolini, and G. Satta. 1991. The Aeromonas hydrophila cphA gene: molecular heterogeneity among class B metallo-β-lactamases. J. Bacteriol. 173:4611–4617.
- Medeiros, A. A. 1997. Evolution and dissemination of beta-lactamases accelerated by generations of beta-lactam antibiotics. Clin. Infect. Dis. 24:S19– S45.
- Moulin, V., J. Freney, W. Hansen, and A. Philippon. 1992. Comportement phénotypique des *Flavobacterium* vis-à-vis de 39 antibiotiques. Med. Mal. Infect. 22:902–908.
- Naas, T., D. M. Livermore, and P. Nordmann. Characterization of an LysR family protein, SmeR from *Serratia marcescens* S6, its effect on expression of the carbapenem-hydrolyzing beta-lactamase Sme-1, and comparison of this regulator with other beta-lactamase regulators. 39:629–637.
- Naas, T., and P. Nordmann. 1994. Analysis of a carbapenem-hydrolyzing class A beta-lactamase from *Enterobacter cloacae* and of its Lys-R type regulatory protein. Proc. Natl. Acad. Sci. USA 91:7693–7697.
- National Committee for Clinical Laboratory Standards. 1993. Methods for dilution antimicrobial susceptibility tests for bacteria that grow aerobically. Approved standard M7-A3. National Committee for Clinical Laboratory Standards, Wayne, Pa.
- Nicolas, M. H., V. Jarlier, N. Honoré, A. Philippon, and S. T. Cole. 1989. Molecular characterization of the gene encoding SHV-3 beta-lactamase responsible for transferable cefotaxime resistance in clinical isolates of *Klebsiella pneumoniae*. Antimicrob. Agents Chemother. 33:2096–2100.
- Nordmann, P., and T. Naas. 1994. Sequence analysis of PER-1 extendedspectrum beta-lactamase from *Pseudomonas aeruginosa* and comparison with class A beta-lactamases. Antimicrob. Agents Chemother. 38:104–114.

- Parker, A. C., and C. J. Smith. 1993. Genetic and biochemical analysis of a novel Ambler class A beta-lactamase responsible for cefoxitin resistance in *Bacteroides* species. Antimicrob. Agents Chemother. 37:1028–1036.
- Patton, R., R. S. Miles, and S. G. Amyes. 1994. Biochemical properties of inducible β-lactamases produced from *Xanthomonas maltophilia*. Antimicrob. Agents Chemother. 38:2143–2149.
- Payne, D. J., R. Cramp, J. H. Bateson, J. Neal, and D. Knowles. 1994. Rapid identification of metallo- and serine β-lactamases. Antimicrob. Agents Chemother. 38:991–996.
- Phillipon, L. N., T. Naas, A. T. Bouthors, V. Barakett, and P. Nordmann. 1997. OXA-18, a class D clavulanic acid-inhibited extended-spectrum β-lactamase from *Pseudomonas aeruginosa*. Antimicrob. Agents Chemother. 41: 2188–2195.
- Pickett, M. J. 1989. Methods for identification of flavobacteria. J. Clin. Microbiol. 27:2309–2315.
- Poirel, L., M. Guibert, D. Girlich, T. Naas, and P. Nordmann. 1999. Cloning, sequence analyses, expression and distribution of *ampC-ampR* from *Morganella morganii* clinical isolates. Antimicrob. Agents Chemother. 43:769–776.
- 32. Poirel, L., T. Naas, M. Guibert, E. B. Chaibi, R. Labia, and P. Nordmann. 1999. Molecular and biochemical characterization of VEB-1, a novel class A extended-spectrum β-lactamase encoded by an *Escherichia coli* integron gene. Antimicrob. Agents Chemother. 43:573–581.
- Pollitt, S., and H. Zalkin. 1983. Role of primary structure and disulfide bond formation in β-lactamase secretion. J. Bacteriol. 153:27–32.
- Richard, C., and H. Monteil. 1983. Isolement, identification, signification clinique du genre *Flavobacterium*. Ann. Biol. Clin. 4:187–198.
- 35. Rogers, M. B., A. C. Parker, and C. J. Smith. 1993. Cloning and characterization of the endogenous cephalosporinase gene, *cepA*, from *Bacteroides fragilis* reveals a new subgroup of Ambler class A β-lactamases. Antimicrob. Agents Chemother. 37:2391–2400.
- 36. Rossolini, G. M., N. Franceschini, M. L. Riccio, P. S. Mercuri, M. Perilli, M. Galleni, J. M. Frère, and G. Amicosante. 1998. Characterization and sequence of the *Chryseobacterium* (Flavobacterium) *meningosepticum* carbapenemase: a new molecular class B β-lactamase showing a broad substrate profile. Biochem. J. 332:145–152.
- Sambrook, J., E. F. Fritsch, and T. Maniatis. 1989. Molecular cloning: a laboratory manual, 2nd ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.

- Schultz, S. C., G. Dabadie-McFarkand, J. J. Neitzel, and J. H. Richards. 1987. Stability of wild-type and mutant RTEM-1 β lactamase: effect of the disulfide bond. Proteins 2:290–297.
- Siegman-Igra, Y., D. Schwartz, G. Soferman, and N. Konforti. 1987. Flavobacterium group IIb bacteremia: report of a case and review of Flavobacterium infections. Med. Microbiol. Immunol. 176:103–111.
- Smith, C. J., T. K. Bennett, and A. C. Parker. 1994. Molecular and genetic analysis of the *Bacteroides uniformis* cephalosporinase gene, *cblA*, encoding the species-specific beta-lactamase. Antimicrob. Agents Chemother. 38:1711–1715.
- Sougakoff, W., S. Goussard, and P. Courvalin. 1988. The TEM-3 β-lactamase, which hydrolyses broad-spectrum cephalosporins, is derived from the TEM-2 penicillinase by two amino-acid substitutions. FEMS Microbiol. Lett. 56:343–348.
- Sutcliffe, J. G. 1978. Nucleotide sequence of the ampicillin resistance gene of Escherichia coli plasmid pBR322. Proc. Natl. Acad. Sci. USA 75:3737–3741.
- 43. Swaren, P., L. Maveyraud, X. Raquet, S. Cabantous, C. Duez, J. D. Pedelacq, S. Mariotte-Boyer, L. Mourey, R. Labia, M. H. Nicolas-Chanoine, P. Nordmann, J. M. Frère, and J. P. Samama. 1998. X-ray analysis of the NMC-A β-lactamase at 1.64 Å resolution, a class A carbapenemase with broad substrate activity. J. Biol. Chem. 41:26714–26721.
- Swofford, D. L. 1989. PAUP (version 3.0): phylogenetic analysis using parsimony. Illinois Natural History Survey, Champaign.
- 45. Vandamme, P., J. F. Bernardet, P. Segers, K. Kersters, and B. Holmes. 1994. New perspectives in the classification of the flavobacteria: description of *Chryseobacterium* gen. nov., *Bergeyella* gen. nov., and *Empedobacter* nom. rev. Int. J. Syst. Bacteriol. 44:827–831.
- 46. Vanhove, M., G. Guillaume, P. Ledent, J. H. Richards, R. H. Pain, and J. M. Frère. 1997. Kinetic and thermodynamic consequences of the removal of the cys-77-cys-123 disulphide bond for the folding of TEM-1 β-lactamase. Biochem. J. 321:413–417.
- Walsh, T. R., A. P. MacGowan, and P. M. Bennett. 1997. Sequence analysis and enzyme kinetics of the L2 serine beta-lactamase from *Stenotrophomonas* maltophilia. Antimicrob. Agents Chemother. 41:1460–1464.
- Yanisch-Perron, C., J. Vieira, and J. Messing. 1985. Improved M13 phage cloning vectors and hosts strains: nucleotide sequences of the M13mp18 and pUC19 vectors. Gene 33:103–119.