## KHM-1, a Novel Plasmid-Mediated Metallo-β-Lactamase from a *Citrobacter freundii* Clinical Isolate<sup>∇</sup>

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A novel gene,  $bla_{\rm KHM-1}$ , encoding a metallo- $\beta$ -lactamase, KHM-1, was cloned from a clinical isolate of *Citrobacter freundii* resistant to most  $\beta$ -lactam antibiotics. *Escherichia coli* expressing  $bla_{\rm KHM-1}$  was resistant to all broad-spectrum  $\beta$ -lactams except for monobactams and showed reduced susceptibility to carbapenems. Recombinant KHM-1 exhibited EDTA-inhibitable hydrolytic activity against most  $\beta$ -lactams, with an overall preference for cephalosporins.

Acquired metallo- $\beta$ -lactamases (MBLs) produced by gramnegative bacteria, including *Pseudomonas aeruginosa, Acinetobacter* spp., and several enterobacteria, confer resistance to all  $\beta$ -lactams except the monobactams (2). Acquired MBLs are categorized on the basis of amino acid sequences into various types (2, 23). The IMP- and VIM-type enzymes are the most common and are found worldwide (2, 8, 23). Recently, four additional types, SPM, GIM, SIM, and AIM, have been found in Brazil (21), Germany (3), Korea (11), and Australia (24), respectively. We report here on the detection of a novel acquired MBL in a clinical isolate of *Citrobacter freundii* identified in Japan.

*C. freundii* strain KHM243 was isolated in 1997 from a patient with catheter-associated urinary tract infection at Kyorin University Hospital (Tokyo, Japan). *Escherichia coli* K-12 strain W1895 was used as the recipient in conjugation experiments. *E. coli* JM109 (Takara Bio, Shiga, Japan) was used as the host for recombinant plasmids. Plasmid pHSG396 (Takara Bio) was used for the cloning of *bla*<sub>KHM-1</sub> fragments.

Susceptibility to  $\beta$ -lactams was determined by the microdilution method (4). The production of MBL was detected by a double-disk synergy test with disks containing sodium mercaptoacetic acid (MBL production test; Eiken Chemical Co. Ltd., Tokyo, Japan), as described by Arakawa et al. (1).

The transfer of resistance by conjugation was analyzed as described previously (7). *E. coli* transconjugants were selected on Penassay broth agar (antibiotic medium no. 3; Becton Dickinson, Franklin Lakes, NJ) containing rifampin (200  $\mu$ g/ml; Wako Pure Chemical Industries, Ltd., Osaka, Japan) and moxalactam (16  $\mu$ g/ml; Shionogi & Co., Ltd., Osaka, Japan). Plasmid DNA was extracted by an alkaline lysis procedure (9). Plasmid R100 (94.5 kb) (13) from *E. coli* CSH2, plasmid R478 (275 kb) (6) from *E. coli* J53, and three cryptic plasmids (200,

\* Corresponding author. Mailing address: Department of Microbiology, Kyorin University School of Health Sciences, 476 Miyashitacho, Hachioji, Tokyo 192-8508, Japan. Phone: (81) 42 691 0011, ext. 4125. Fax: (81) 42 691 1094. E-mail: moritako@kyorin-u.ac.jp. 60, and 2.4 kb) from *Salmonella enterica* serovar Enteritidis L119 (15) were used as molecular size markers.

PCR analysis specific for class 1 integrons was performed as described previously (12). DNA sequences flanking  $bla_{\rm KHM-1}$  were determined by inverse PCR (16). Briefly, plasmid DNA extracted from *E. coli* transconjugant W1895(pCF243) was digested with EcoRV or XspI (Takara Bio). Self-ligated digests were used as the template for an inverse PCR. The upstream and downstream flanking regions of  $bla_{\rm KHM-1}$  were amplified by inverse PCR with two sets of primers: primers 5'-CGATA TAACAAGAGCTATTTTCAT-3' and 5'-GGTATGCGCTG ACGATTC-3' for the upstream region and primers 5'-GGTG TACAGATAAACGCCG-3' and 5'-TTTATTTGGTGGCTG TTTTGTC-3' for the downstream region.

The KHM-1 MBL from *E. coli* JM109(pKHM-1) was purified with HiTrap Q HP and Superdex 200 columns (GE Healthcare Bio-Sciences KK, Tokyo, Japan), as described by Franceschini et al. (5). During the purification procedure, the presence of  $\beta$ -lactamase activity was monitored with 100  $\mu$ M nitrocefin (Oxoid Ltd., Basingstoke, United Kingdom). The protein concentration was determined with a bicinchoninic acid protein assay kit (Pierce, Rockford, IL). Kinetic analysis was carried out in 50 mM phosphate buffer (pH 7.0) at 25°C with a UV-visible spectrophotometer (V-530; Jasco, Tokyo, Japan). The  $K_m$  and  $k_{cat}$  values and the  $k_{cat}/K_m$  ratio were determined by analyzing  $\beta$ -lactam hydrolysis under initial-rate conditions by use of the Lineweaver-Burk plot.

Antibiotic susceptibility testing showed that *C. freundii* KHM243 was resistant to most  $\beta$ -lactams and showed reduced susceptibility to carbapenems (Table 1). However, KHM243 was susceptible to monobactams (carumonam and aztreonam). The isolate was positive by the MBL production test (data not shown).

*C. freundii* KHM243 has two plasmids, one of approximately 70 kb and one of approximately 200 kb. A conjugation experiment was done with KHM243 and *E. coli* W1895. W1895 transconjugants that were resistant to  $\beta$ -lactams and that contained a 200-kb plasmid, designated pCF243, were obtained. The transconjugant exhibited a profile of susceptibility to  $\beta$ -lactams similar to that of KHM243, although the MICs for

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TABLE 1. MICs of $\beta$ -lactams for <i>C. freundii</i> KHM243, <i>E. coli</i>
W1895(pCF243) transconjugant, E. coli JM109(pKHM-1)
expressing the KHM-1 MBL, and E. coli
host strains

	MIC (µg/ml)				
Antibiotic(s) <sup>a</sup>	C. freundii KHM243	<i>E. coli</i> W1895 (pCF243) <sup>b</sup>	<i>E. coli</i> W1895	<i>E. coli</i> JM109 (pKHM-1) <sup>c</sup>	E. coli JM109
Ampicillin	256	64	8	16	1
Ampicillin- sulbactam	64	64	1	16	0.5
Ticarcillin	>512	>512	2	512	2
Ticarcillin-clavulanic acid	512	>512	4	512	2
Piperacillin	4	16	2	4	0.25
Cephaloridine	512	128	2	64	1
Cefuroxime	>512	>512	2	>512	8
Ceftazidime	>512	>512	0.125	>512	0.063
Cefotaxime	64	>512	0.008	128	0.004
Cefepime	32	>512	0.002	64	0.004
Cefozopran	16	256	0.016	64	0.008
Imipenêm	2	4	0.063	0.5	0.063
Meropenem	4	4	0.004	4	0.004
Aztreonam	0.25	0.063	0.031	0.063	0.031
Carumonam	0.25	0.125	0.031	0.063	0.031
Cefoxitin	512	>512	8	>512	8
Cefmetazole	512	512	0.5	>512	0.25
Cefotetan	128	512	0.125	>512	0.031
Cefbuperazone	128	256	0.063	512	0.031
Cefminox	512	>512	0.125	512	0.25
Moxalactam	256	>512	0.063	>512	0.031
Flomoxef	64	256	0.031	128	0.031

<sup>a</sup> The ratio of the ampicillin to sulbactam was 2:1. The ratio of ticarcillin to clavulanic acid was 15:1.

<sup>b</sup> Natural plasmid carrying the *bla* KHM-1 gene.
<sup>c</sup> Recombinant plasmid constructed by insertion of DNA fragment containing the bla<sub>KHM-1</sub> gene into the cloning vector pHSG396.

some cephalosporins, including cefotaxime, cefepime, and cefozopran, were significantly higher in the transconjugant than in KHM243 (Table 1).

EcoRI-digested fragments of pCF243 were subcloned into pHSG396 and were transformed into E. coli JM109 cells, and transformants were selected on agar medium containing moxalactam (1 µg/ml). Strain JM109 carrying the plasmid that conferred resistance to moxalactam, named pKHM-1, exhibited a profile of susceptibility to β-lactams similar to the susceptibility profiles of KHM243 and the E. coli W1895 transconjugant carrying pCF243 (Table 1). However, the MICs of some antibiotics, including cefotaxime and cefepime, were lower for the transformant carrying pKHM-1 than the transconjugant. This might be explained by insufficient expression of the gene due to insertion of the DNA fragment with a small 5'-flanking region.

pKHM-1 contained an 837-bp insert with a complete open reading frame (ORF) (data not shown). The 726-bp ORF encoded a putative protein of 241 amino acids. The protein was similar to MBLs, such as Uvs123 from an uncultured bacterium (82% identity) (22), IMP-1 (59% identity) (17), and SIM-1 (59% identity) (11) (Fig. 1). The protein was somewhat less similar to VIM-1 (38% identity) (10), GIM-1 (50% identity) (3), and SPM-1 (46% identity) (21) (Fig. 1). We named the ORF encoding the protein bla<sub>KHM-1</sub> and designated the protein KHM-1 (Kyorin Health Science MBL 1). bla<sub>KHM-1</sub> was different from the Citrobacter freundii genome in its GC contents (GC contents, 50.27% and 44.63%, respectively) and codon usage (data not shown). KHM-1 contained amino acid motifs conserved in MBL enzymes, including a zinc-binding motif (HXHXD, residues 97 to 101) and three other residues involved in zinc binding (residues 159, 178, and 217) (Fig. 2) (18, 23).

The DNA sequences flanking *bla*<sub>KHM-1</sub> were determined from 774 bp upstream to 806 bp downstream of it. No sequence homologies for site-specific cointegration events, ORFs, or transmissible elements was detected within the 774-bp upstream of *bla*<sub>KHM-1</sub>. A 360-bp ORF encoding a putative protein of 119 amino acids with 77% identity to hypothetical protein VP1798 of Vibrio parahaemolyticus (14) was located in the fragment 21 to 380 bp downstream of bla<sub>KHM-1</sub>. Strain KHM243 carried a class 1 integron with an array of two gene cassettes, which carried the aadA2 (20) and aac(6')-Iae (19) aminoglycoside resistance determinants; however,  $bla_{KHM-1}$ was not detected in this integron.

Analysis of the purified KHM-1 protein by sodium dodecyl sulfate-polyacrylamide gel electrophoresis showed a single 25kDa band. The activity of KHM-1 against various β-lactams was analyzed with the purified protein. It showed hydrolytic activity against all β-lactams tested except aztreonam (Table 2). Enzymatic activity against aztreonam was undetectable un-

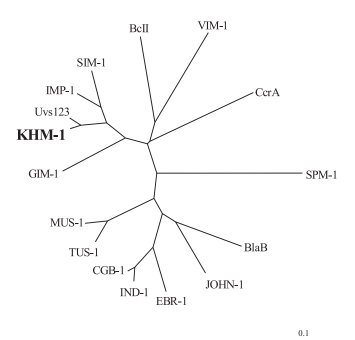


FIG. 1. Dendrogram showing the similarity of KHM-1 to other MBLs. KHM-1 and MBLs from a variety of organisms were tested. The dendrogram was created with the ClustalW program. Branch lengths correspond to the number of amino acid exchanges of the following MBL proteins (GenBank accession numbers, source organism) of BcII (P04190, from Bacillus cereus), BlaB (CAA65601, from Elizabethkingia meningoseptica), CcrA (P25910, from Bacteroides fragilis), CGB-1 (AAL55263, from Chryseobacterium gleum), EBR-1 (AAN32638, from Empedobacter brevis), GIM-1 (CAF05908, from Pseudomonas aeruginosa), IMP-1 (AAB30289, from Serratia marcescens), IND-1 (AAD20273, from Chryseobacterium indologenes), JOHN-1 (AAK38324, from Flavobacterium johnsoniae), MUS-1 (AAN63647, from Myroides odoratimimus), SIM-1 (AAX76774, from Acinetobacter baumannii), SPM-1 (CAD37801, from P. aeruginosa), TUS-1 (AAN63648, from Myroides odoratus), Uvs123 (AAP70377, from uncultured bacterium), and VIM-1 (CAB46686, from P. aeruginosa).

C. freundii KHM-1 Uncultured bacterium Uvs123 S. marcescens IMP-1 A. baumannii SIM-1 P. aeruginosa GIM-1 E. meningoseptica BlaB F. johnsoniae JOIN-1 C. indologenes IND-1 C. indologenes IND-1 E. brevis EBR-1 M. odoratus TUS-1 M. odoratus TUS-1 B. fragilis CcrA B. cereus BCI P. aeruginosa VIM-1 P. aeruginosa VIM-1 P. aeruginosa SPM-1	1 10 20 30 40 50 MKIALVISFGLLFTMMVCADDSLPELDIQKIEDGVŸLŸTAYEKVEGWGLV MKVFLAVSLFLISNLLFAEDPLPELEIKKIEGGVYLYTAYEKVEGWGLV MSKLSVFFIFLFCSIATAAESLPDLKIEKLDEGVYLHTSFEEVNGWGVV MRTLILCLFGTLMTAFAEEAQPDLKIEKIEDGVYLHTSFEEVNGWGVV MKNVLVFLILLVALPALAQGHKPLEVIKIEDGVYLHTSFKNIEGYGLV MKKNISILIALGLFSLQAFGQENPDVKIEKLKDNLYVYTTYNTFNGTKYA MKKSIRFFIVSILLSPFA.SAQVKDFVIEPFIKNNLHIYKTFGVFGGKEYS MKKSIRFFISHLSPLAVSNSLGQSKNSPLQISHLTGDFYVYRTFNDYKGKKIS MKKSIRFFISHLSPLAVSNSLGQSKNSPLQISHLTGDFYVYRTFNDYKGKKIS MKKSIRFFISHLSPLAVSNSLGQSKNSPLQISHLTGDFYVYTYNFPNGVEGKEYS MKKSIRFFISHLSPLAVSNSLGQSKNSPLQISHLTGDFYVYTYNFPNGVEGKEYS MKKSIRFFISHLSPLAVKSNSLGQSKNSPLGISHTGDFYVYRTFNDYKGKKIS MKKSIRFFISHLSPLAVKSNSLGQSKNSPLGISHTGDFYVYRTFOVFGGKEYS MKKSIRFFISHLSPLAVKSNSLGSAFG.QIKFUNDHMYVYTTYQVFGGKEYS MKKSIRFFISHLSPLAVKSNSLGSAFG.QIKFUNDHMYVYTTYQVFGGVEYS MKKSIRFFISHLSPLAVKSSCHSAGSAFAFG.QIKFUNDHMYVYTTYQFFGGVTSS MKKVFILSNLFPVAVMAQKSVKISDDISITQLSDKVYTYVSLAEIEGWGMV MKKNTLLKVGLCVGLLGTIQFYSTISSVQASQKVEKVIKNETGISISQLNKNVWVHTELGSFNG. AVY MTDRDFYS
C. freundii KHM-1 Uncultured bacterium Uvs123 S. marcescens IMP-1 A. baumannii SIM-1 P. aeruginosa GIM-1 E. meningoseptica BlaB F. johnsoniae JOIN-1 C. gleam CGB-1 E. brevis EBR-1 M. odoratus TUS-1 M. odoratus TUS-1 B. fragilis CcrA B. ccrea BeII P. aeruginosa VIM-1 P. aeruginosa SPM-1	60 70 80 90 10 10 10 120 SSNGLVVLDNKNAYLIDTPISATDTEKLVKWIDA.QGFTAKASISTIFITSSTGGIAPLNSKSISKIFTYASK SSNGLVVLDNKNAYLIDTPISATDTEKLVKWIDA.QGFTAKASISTIFITSSTGGIAPLNSKSISTYASK SSNGLVVLDNKAAYLIDTPISATDTEKLVKWIDA.QGFTAKASISTIFISSSTGGIAPLNSKSISTYASK SKQGLVVLDNKAAYLIDTPISATDTEKLVNWIDA.QGFTAKASISTIFISSSTGGIAPLNSKSISTYASK SSNGLVVLDNKAAYLIDTPISATDTEKLVNWIDA.QGFTAKASISTIFISSSTGGIAPLNSKSISTYASK SSNGLVVLDNKAAYLIDTPISATDTEKLVNWIDA.QGFTAKASISTIFISSSTGGIAPLNSKSISTYASK SSNGLVVLDNKAAYLIDTPISATDTEKLVNWIDK.ROYVNASISTISTSSTSSTGGIAPLNSKSISTYASK SSNGLVVLDNKGVVLFDVPWSENSTSSTSSTSSTSSTSSTSSTSSTSTSSTSSTS SNGLVVLDNKGVVLFDVPWSENSTSSTSSTSSTSSTSSTSSTSSTSSTSSTS SNGLVVLTKGVVLFDVPWSENSTSSTSSTSSTSSTSSTSSTSSTSSTS SNGLVVLTKGVVLFDVPWSENSTSSTS SNGLVVLTKGVVLFDVPWSENSTSSTS SNGLVVLTKGVVLFDVPWSENSTS SNGLVVLTKGVVLFDVPWSENSTS SNGLVVTNKGIVLFDVPWSENSTS SNGLVLTKGVVLFDVPWSENSTS SNGLVLTKGVVLFDVPWSENSTS SNGLVLTKGVVLFDVPWSENSTS SNGLVLTKGVVLFDVPWSENSTS SNGLVLTKGVVLFDVPWSENSTS SNGLVLTKGVVLFDVPWSENSTS SNGLVLTKGVVLFDVPWSENSTS SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPWSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVLTKGVVLFDVPMSENST SNGLVVTNGGVENST SNGLVVTVTDEST SNGLVLTKGVVLFDVPMSENST SNGLVVTNGGVENST SNGLVVTVTMENT SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVVLTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVENST SNGLVVTMGGVF
C. freundii KHM-1 Uncultured bacterium Uvs123 S. marcescens IMP-1 A. baamannii SIM-1 P. aeruginosa GIM-1 E. meningoseptica BlaB F. johnsoniae JOIN-1 C. indologenes IND-1 C. gleum CGB-1 E. brevis EBR-1 M. odoratus TUS-1 M. odoratus TUS-1 B. fragilis CcrA B. cereus BCH P. aeruginosa VIM-1 P. aeruginosa SPM-1	139 149 150 160 INNULKING BEQATHERGK, DIALON NY WULKINK HEAFY POAGET DIA INNULKING BEQATHERGK, DIALON NY WULKINK HEAFY POAGET DIA INNULKING KUQATINSPSG, DIALON DIALON DIFTUSION KIEVY POAGET DIA INNULKING KUQATINSPSG, DIALON DIFTUSION DIFTUSION CONTRACTOR DIFTUSION INNULKING KUQATINSPSG, DIFTUSION CONTRACTOR DIFTUSION INNULKING KUPATINI KUTA INNULKING KUPATINI KUTA INNUL
C. freundii KHM-1 Uncultured bacterium Uvs123 S. marcescens IMP-1 A. baumannii SIM-1 P. aeruginosa GIM-1 E. meningoseptica BlaB F. johnsoniae JOIN-1 C. indologenes IND-1 C. gleum CGB-1 E. brevis EBR-1 M. odoratus TUS-1 M. odoratus TUS-1 B. fragilis CcrA B. cereus Bell P. aeruginosa VIM-1 P. aeruginosa SPM-1	170 180 190 200 210 220 230 VYWWLPKCHILFGGCFVKPEGLGNISHAVIKENPASAEKLIIKTYGNATHVVPGHGKUGDASLLEKTR VYWWLPKCHILFGGCFVKPEGLGNISHAVIKENPASAEKLIIKTYGSATHVVPGHGKUGDASLLEKTR VYWWLPKENTIFGGCFVKPEGLGNISHAVIKENPASAEKLIITYGSATHVVPGHGKUGDASLLEKTR VYWWLPKENTIFGGCFVKPEGLGNISHAVIKENPASAEKLIITYGSATHVVPGHGKUGDASLLEKTR VYWWLPKKUIFGGCFVKPEGLGNISHAVIKENPASAEKLIITYGSATHVVPGHGKUGDASLLEKTR VYWWLPKKUIFGGCFVKPEGLGNISHAVIKENPASAEKLIITYGSATHVVPGHGKUGDASLLEKTR VYWWPKKKUIFGGCFVKSAEALDLGYIGBANVENPASAKLIKSKYGAAKLIVPSSEIGDASLLKLTW VYWWPKKKUIFGGCVVKSAEALDLGYIGBANVENPASAKLIKSKYGAAVIVPGHGKUGSGHVEHTL VYWWPFKKNUIFGGCLVKSAAADLGYIGBANVENPASAKLIKSKYGAATHIIPGHDWKGGGHVEHTL VYWWFPKKNUIFGGCLVKSAAADLGYIGBANVENPASAKLIKSKYGAATHIIPGHDWKGGGHVEHTL VYWWFPKKNUIFGGCLVKSAAADLGYIGBANVENPASAKLIKSKYGAATHIIPGHDWKGGGHVEHTL VYWWFPKKNUIFGGCLVKSAAADLGYIGBANVENPASAKLIKSKYGAATHIIPGHDWKGGGHVEHTL VYWWFPKKNUIFGGCLVKSAAADLGYIGBANVENPASAENVENPKKLKSKYGAATHIIPGHDWKGGGHVEHTL VYWFPKKNUIFGGCLVKSAAADLGYIGBANVENPAN VYWFPKKNUIFGGCLUKSAAADLGYIGBANVENPKTILAVKRFKKAKVIIPGHDWKATGHLENTA VYWFPKKKKUIFGGCMLKSAAATHIGNIADGNVIAPPKTILAVKRFKFKAKVIIPGHDWKATGHLENTA VYWFPKKKUIFYGGCLUKSAAADLGYIGBANVENPKTILAVKRFKFKAKVIIPGHDWKATGHLENTA VYWFPKKKUIFYGGCLUKSTAALDLGYIGBANVENPKTILAVKRFKFKAKVIIPGHDWKATGHLENTA VYWFPKKKKUIFGGCMLKSIGNIADAUANINNAPPKTILAVKRFKFKAKVIIPGHDWKATGHLENTA VYWFPKKKKUIFGGCMLKSIGNIADAUANINNAPPTSUENVLKKYFVNINAVVPGHGEVGDFWUNKT
C. freundii KHM-1 Uncultured bacterium Uvs123 S. marcescens IMP-1 A. baumannii SIM-1 P. aeruginosa GIM-1 E. meningoseptica BlaB F. johnsoniae JOHN-1 C. indologenes IND-1 C. gleum CGB-1 E. brevis EBR-1 M. odoratus TUS-1 M. odoratus TUS-1 B. fragilis CerA B. cereus Bell P. aeruginosa VIM-1 P. aeruginosa SPM-1	240 QRAVEALAAKK QRALEALAAKK EQAVKGLNESKKFSKPSN EQAVKGLNESKKFSKPSN EQAVKGLNESKKFPLIN DLAESASNKLMOPTAEASAD DLINEYQOKQKASN KLVDEVLAQKSAGKK ELLNKK DLINKK RILSAYQAQKLKNNKQL RILSAYQAQKLKNNKQL RILSAYQAQKLKNNKQL QIVNQYIESTSKP DLK NVVKAHKNRSVAE KVAEKAYGEMRL

FIG. 2. Multiple-sequence alignments of the amino acid sequence of KHM-1 from *Citrobacter freundii* isolate KHM243 with those of other MBLs. The sequence sources are the same as those indicated in the legend to Fig. 1. Sequence comparison was performed by aligning the protein amino acid sequences by use of the ClustalW program (http://clustalw.ddbj.nig.ac.jp/top-e.html). The residues known to be involved in metal binding are indicated by asterisks. Identical residues are shaded.

Substrate	$K_m \; (\mu \mathrm{M})^a$	$k_{\rm cat}  ({\rm s}^{-1})^a$	$k_{\text{cat}}/K_m$ (M <sup>-1</sup> ·s <sup>-1</sup> )
Penicillin G	$1,340 \pm 56$	$23 \pm 0.9$	$1.7 \times 10^{4}$
Ampicillin	$978 \pm 111$	$19 \pm 2$	$1.9 \times 10^{4}$
Cephaloridine	$4.4 \pm 0.95$	$686 \pm 12$	$1.6 \times 10^{8}$
Cefoxitin	$81 \pm 4$	$1,178 \pm 164$	$1.4 \times 10^{7}$
Cefotaxime	$13 \pm 1.5$	$2,181 \pm 208$	$1.7 \times 10^{8}$
Ceftazidime	$8 \pm 0.4$	$118 \pm 3$	$1.5 \times 10^{7}$
Moxalactam	$71 \pm 8$	$2,794 \pm 260$	$3.9 \times 10^{7}$
Aztreonam	b	_	_
Meropenem	$12 \pm 3$	$0.4\pm0.015$	$3.3 \times 10^{4}$
Imipenem	$268 \pm 53$	$15 \pm 3$	$5.6 \times 10^{4}$

TABLE 2. Kinetic parameters of  $\beta$ -lactamase KHM-1 with various substrates

 $^a$  The  $K_m$  and  $k_{\rm cat}$  values represent the means of three independent experiments  $\pm$  standard deviations.

<sup>b</sup> —, no hydrolysis was detected under conditions with a substrate concentration of up to 1 mM and an enzyme concentration of up to 840 nM.

der the experimental conditions adopted. This activity was inhibited by EDTA but was recovered by addition of Zn<sup>2+</sup> (data not shown). The kinetic parameters, including  $K_m$ ,  $k_{cat}$ , and the  $k_{cat}/K_m$  ratio, were determined for several different  $\beta$ -lactams (Table 2). Relatively higher values of the  $k_{cat}/K_m$ ratio (>10<sup>7</sup> M<sup>-1</sup> · s<sup>-1</sup>), as a result of low values of  $K_m$  and high values of  $k_{cat}$ , were observed with the cephalosporins tested (cephaloridine, cefoxitin, cefotaxime, ceftazidime, and moxalactam); and lower values of the  $k_{cat}/K_m$  ratio (<10<sup>5</sup> M<sup>-1</sup> · s<sup>-1</sup>) were observed with penicillin G, ampicillin, meropenem, and imipenem.

During 1997 and 1998, 104, 13, and 5 clinical isolates of *C. freundii*, *C. koseri*, and other *Citrobacter* spp., respectively, were collected in the hospital and were screened for imipenem resistance. Of these, four isolates of *C. freundii* showed reduced susceptibilities to imipenem (MICs,  $>8 \ \mu g/ml$ ). However,  $bla_{\rm KHM-1}$  was not detected in any of these isolates except the one from the patient infected with strain KHM243. A laboratory-based survey of other isolates of the family *Enterobacteriaceae* is in progress to detect *bla*<sub>KHM-1</sub>.

**Nucleotide sequence accession number.** The nucleotide sequence data for  $bla_{\rm KHM-1}$  and its flanking region from 774 bp upstream to 806 bp downstream reported here have been deposited in the EMBL/GenBank/DDBJ databases under accession number AB443628.

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