



Journal of Enzyme Inhibition and Medicinal Chemistry

ISSN: 1475-6366 (Print) 1475-6374 (Online) Journal homepage: https://www.tandfonline.com/loi/ienz20

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To cite this article: Ayşenur Eminoğlu, Daniela Vullo, Aycan Aşık, Dilşat Nigar Çolak, Claudiu T. Supuran, Sabriye Çanakçı & Ali Osman Beldüz (2016) Cloning, expression and biochemical characterization of a  $\beta$ -carbonic anhydrase from the soil bacterium *Enterobacter* sp. B13, Journal of Enzyme Inhibition and Medicinal Chemistry, 31:6, 1111-1118, DOI: 10.3109/14756366.2015.1100176

To link to this article: https://doi.org/10.3109/14756366.2015.1100176



Published online: 26 Oct 2015.

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### Journal of Enzyme Inhibition and Medicinal Chemistry

www.tandfonline.com/ienz ISSN: 1475-6366 (print), 1475-6374 (electronic)

J Enzyme Inhib Med Chem, 2016; 31(6): 1111–1118 © 2015 Informa UK Limited, trading as Taylor & Francis Group. DOI: 10.3109/14756366.2015.1100176

#### **RESEARCH ARTICLE**

## Cloning, expression and biochemical characterization of a $\beta$ -carbonic anhydrase from the soil bacterium *Enterobacter* sp. B13

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#### Abstract

A recombinant carbonic anhydrase (CA, EC 4.2.1.1) from the soil-dwelling bacterium *Enterobacter* sp. B13 was cloned and purified by Co<sup>2+</sup> affinity chromatography. Bioinformatic analysis showed that the new enzyme (denominated here B13-CA) belongs to the  $\beta$ -class CAs and to possess 95% homology with the ortholog enzyme from *Escherichia coli* encoded by the *can* gene, whereas its sequence homology with the other such enzyme from *E. coli* (encoded by the cynT gene) was of 33%. B13-CA was characterized kinetically as a catalyst for carbon dioxide hydration to bicarbonate and protons. The enzyme shows a significant catalytic activity, with the following kinetic parameters at 20 °C and pH of 8.3:  $k_{cat}$  of  $4.8 \times 10^5 \text{ s}^{-1}$  and  $k_{cat}/K_m$  of  $5.6 \times 10^7 \text{ M}^{-1} \times \text{s}^{-1}$ . This activity was potently inhibited by acetazolamide which showed a  $K_1$  of 78.9 nM. Although only this compound was investigated for the moment as B13-CA inhibitor, further studies may reveal new classes of inhibitors/activators of this enzyme which may show biomedical or environmental applications, considering the posssible role of this enzyme in CaCO<sub>3</sub> biomineralization processes.

#### Introduction

Carbonic anhydrases (CAs) [EC 4.2.1.1] are Zn(II)dependent metalloenzymes which catalyze the reversible hydration of CO<sub>2</sub> to HCO<sub>3</sub><sup>-</sup> and H<sup>+1</sup>. CAs have important functions in the eukaryotes as well as in prokaryotes, such as pH regulation, biosynthetic reactions, respiration, CO<sub>2</sub> transport, photosynthesis, among others<sup>2</sup>, and are known to be among the fastest enzymes described so far. Bacterial CAs have an essential role in the life cycle of these organisms, as well as in the regulation of calcium carbonate mineralization, at least in some bacterial species. Indeed, CaCO<sub>3</sub> mineralization by bacteria is crucial for soil and ground water remediation, sequestration and capture of atmospheric CO<sub>2</sub>, as well as sand and soil strengthening and consolidation<sup>3</sup>.

CAs are encoded by six evalutionary unrelated families, the  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$ ,  $\zeta^4$  and  $\eta$ -CAs<sup>5</sup>, which are present in organisms all over the phylogenetic tree. Among them, the  $\alpha$ -,  $\beta$ -,  $\gamma$ - and  $\delta$ -CAs are broadly distributed in many microorganisms<sup>1</sup>, whereas the  $\eta$ -CAs were described only in protozoa for the moment<sup>5</sup>.

To date, two different  $\beta$ -CAs have been reported in *Escherichia coli*, the most investigated bacterium for the molecular biology viewpoint. One such enzymes is encoded

#### Keywords

β-Carbonic anhydrase, *Enterobacter* sp., kinetics

Tavlor & Francis

Taylor & Francis Group

#### History

Received 26 August 2015 Revised 21 September 2015 Accepted 22 September 2015 Published online 22 October 2015

by the *cynT* gene and was the first  $\beta$ -CA identified in this bacterium, as a product of the cyn operon, being shown to possess sequence homology with higher plants  $\beta$ -CAs and to function in the process of cyanate utilization as a nitrogen source by this bacterium<sup>6,7</sup>. The second one is *can* (previously called *yadF*), which has 30% amino acid sequence homology with  $CynT^8$ . It was reported that can is essential for the growth of E. coli under atmospheric CO<sub>2</sub> conditions<sup>2</sup>. It has been also shown that expression of can is susceptible to the increase of cell density and to the rise in the medium temperature. In E. coli, CA activity is normally provided by the expression of both these enzymes, although detailed data regarding their localization within the bacterium are missing for the moment. In fact, bicarbonate/CO2 are necessary during the normal growth of this bacterium and for the biosynthesis of several biomolecules involved in vital functions. It has been suggested that can is conserved throughout Enterobacteriaceae genera (the type of bacteria to which E. coli also belongs)9. However, only four  $\beta$ -CA genes have been identified in *Enterobacteriaceae* so far, i.e. cynT, can (previously yadF), cah and mig-5<sup>9</sup>. Recently,  $\beta$ -CAs from two *Enterobacter* species were phylogenetically compared with the *E. coli*  $CAs^{10}$ , but only one of these enzymes has been investigated in detail for the moment, i.e. the one encoded by the cynT gene, for which the X-ray crystal structure was reported by Cronk et al.8

Although ultimately there are many studies on various bacterial  $\beta$ -CAs, the investigation of *Enterobacteriaceae* is

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Figure 1. Phylogenetic analysis of 16S rDNA similarities of *Enterobacter* sp. B13 based on the BLAST result using the neighbor-joining method. Scale bar represents 0.01 substitutions per nucleotide position. The organisms and GeneBank accession numbers of analyzed sequences are given in parenthesis.

rather limited with only one *E. coli* enzyme being investigated in some detail, as mentioned above. Here, we describe a new  $\beta$ -CA from a bacterium belonging to this bacteria family, more precisely from an *Enterobacter* sp. B13 recently isolated from soil in Northern Turkey.

#### Materials and methods

0.01

#### Sampling site, culture conditions and microorganisms

The bacterial specimens were isolated from soil and mud samples which were collected from a riverside nearby Trabzon, Sürmene, Turkey. The samples were diluted with dH<sub>2</sub>O and filtrated. The filtrate inoculated in TSA plates and incubated at different temperatures. Selected colonies which grew at 37 °C were then inoculated to modified B-4 agar plates to monitor the CaCO<sub>3</sub> decomposition properties of the grown bacteria. Among these isolates, clone B13 showed CO<sub>2</sub> hydratase activity<sup>11</sup> and was subsequently used as a source of genomic DNA in this study.

#### Identification of the bacterial strain by 16S rRNA analysis

For identification of the strain, the 16S rRNA gene was amplified via PCR from genomic DNA with eubacterial universal primers and subsequently sequenced (Macrogen Inc., Seoul, Korea).

## Cloning of *Enterobacter* sp. B13 *can* and construction of the recombinant expression vector

The genomic DNA of Enterobacter sp. was obtained with Wizard Genomic DNA Purification Kit (Promega, Madison, WI) and was used as a template for the CA gene. The amplification process was performed by PCR with two primers in order to amplify the Enterobacter sp. B13 can with NdeI (Biolab) and BamHI (Thermo Scientific, Waltham, MA) recognition sites (CANdeI; 5'-CCA TAT GAA CGA CAT AGA TAC-3" and BamHI CAR5'-CGG ATC CTT ATT TAT GGT TNA CGT GC-3') with the following conditions: initial denaturation steps at 95 °C for 3 min, denaturation at 95 °C for 45 s, followed by annealing at 51 °C for 1 min and primer extension at 72 °C for 1 min 20 s, followed by a step at 72 °C for 5 min, for 35 cycles. The PCR products were cloned into pGEM-T Easy vector (Promega) and sequenced (Macrogen Inc., Seoul, Korea). After the sequence verification, both the positive clone and the expression vector pET-15b (Novagen) were digested with NdeI-BamHI, then excised from the agarose gel (Qiaquick Gel Extraction Kit, Qiagen, Venlo, The Netherlands) and ligated (Thermo Scientific T4 Ligase, Thermo Scientific, Waltham, MA).

Construction and cloning of the recombinant plasmid was carried out in *E. coli* JM101 (Novagen, Podenzano, Italy) as described in the literature<sup>12</sup>. Transformants were checked by enzymatic digestion.



Figure 2. SDS–PAGE analysis (15%) of *Enterobacter* sp. B13-CA (stained with coomassie brilliant blue). Lane 1, protein molecular weight markers (NEB, P7710S), lane 2, purified *Enterobacter* sp. B13-CA.

## Overexpression in *E. coli* BL21 (DE3) *pLysS* and purification of the $\beta$ -CA

The recombinant vector designed as pB13 can was transformed into E. coli BL21 (DE3)pLysS (Novagen) for overexpression and sequenced for correctness. A single colony, harboring the recombinant plasmid was chosen and cultured overnight in Luria Broth (LB) medium containing 50 µg/ml ampicillin. This culture was used to inoculate 200 ml LB-ampicillin medium and incubated at 37 °C with vigorous shaking. When OD<sub>600</sub> reached the mid-log phase,  $\sim 0.6-0.8$ ,  $0.5 \text{ mM ZnSO}_4$  were added to the culture and expression was induced by 0.1 mM isopropyl  $\beta$ -Dthiogalactopyranoside (IPTG) for 24 h at 22 °C, for the production of recombinant B13-CA. Cells were then collected by centrifugation at 9000 rpm for 10 min at 4 °C. The pellet was resuspended in the lysis buffer (30 mM Tris-SO<sub>4</sub> (pH 8.0), 0.1% Triton X-100, 200 mM NaCl, 20 mM imidazole and 0.2 mg/ml lysozyme) and sonicated. After centrifugation at 7000 rpm for 20 min at 4 °C, the supernatant containing N-terminally His-tagged protein was purified by Co<sup>2+</sup> (TALON Metal Affinity Resin, Clontech), preequilibrated with the wash buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub> (pH 8.0), 0.1% Triton X-100, 300 mM NaCl and 20 mM imidazole). After transferring, the lysate was incubated in the His beads in 15 ml falcon tubes overnight at +4 °C. Following the two washing

steps at pH 8.0 and one at pH 7.0 by gentle shaking at +4 °C, the CA was eluted with 250 mM imidazole in the same buffer (pH 7.0). SDS-PAGE (15%) was run to evaluate the purity of the protein. Purified B13-CA was dialyzed overnight in dialyis buffer (30 mM Tris–SO<sub>4</sub>, pH 8.0, 0.5 mM ZnSO<sub>4</sub>, 1 mM dithiothreitol-DTT) at +4 °C. After dialysis, the concentration of the protein was measured with both NanoDrop Spectrophotometer 2000 and at 595 nm, with bovine serum albumin (BSA) as a standard.

#### Zymography

Zymography analysis of the CA activity was performed as described in the literature<sup>13</sup>. To determine the esterase activity, zymogram staining was performed with minor modifications as described previously<sup>14</sup> at two different temperatures (+4 °C and also at RT) on SDS-PAGE (15%). In brief, followed by electrophoresis the gels were incubated in 100 mM Tris–HCl (pH 7.5) including 0.5% Triton X-100, for 6 h at +4 °C and for 4 h at RT. After incubation, the gels were rinsed in 100 mM Tris–HCl, containing 100 mM  $\alpha$ -naphthyl acetate ( $\alpha$ -NAc), dissolved in 2 ml acetone, and 100 mM Tris–HCl (pH 7.5) buffer containing 20 mg Fast Red Salt (two solutions were added to the gels at the same time) until the bands become visible.

#### **Bioinformatic analysis**

The nucleotide sequence of *Enterobacter* sp. B13 *can* was submitted to GeneBank under accession number KT184504. The identification of 16S rDNA sequence was determined by the EzTaxon identification tool (http://www.ezbiocloud.net/eztaxon) against 16S rRNA sequences found in the database, i.e. strains with validly published prokaryotic names<sup>15</sup>. *Haemophilus influenzae* ATCC was selected as outgroup member. Aamino acid resemblance was determined by using the DELTA-BLAST server (http://blast.ncbi.nlm.nih.gov/Blast.cgj).

For predicting the estimated molecular weight and to perform the multiple amino acid sequence alignment, Compute pI/Mw tool of ExPASy<sup>16</sup> and ClustalW<sup>17</sup> were used, respectively. Protein family and conserved domains were identified by Conserved Domain Search Database (http://www.ncbi.nlm.nih.gov/cdd). Determination of evolutionary familiarities was performed by Mega 6<sup>18</sup>.

#### CO<sub>2</sub> hydrase activity assay of B13-CA

An Applied Photophysics stopped-flow instrument was used for assaying the CA catalyzed CO<sub>2</sub> hydration activity<sup>19</sup>. Phenol red (at 0.2 mM) was used as indicator, working at the absorbance maximum of 557 nm with 10 mM TRIS (pH 8.3) as buffer and 0.1 M NaClO<sub>4</sub> (for maintaining constant ionic strength), at 20 °C, following the CA-catalyzed CO<sub>2</sub> hydration reaction for a period of 10–100 s (the uncatalyzed reaction needs around 60–100 s in the assay conditions, whereas the catalyzed ones are of around 6–10 s). The CO<sub>2</sub> concentrations ranged from 1.7 to 17 mM for the determination of the kinetic parameters. The uncatalyzed rates were determined in the same manner and subtracted from the total observed rates. Enzyme concentrations in the assay system were about 16 nM for all the enzymes considered in the present study.

#### **Results and discussion**

## 16S rDNAsequence analysis and characterization of the bacteria

The result of the microbial growth on the B-4 agar plates led to the observation that the B13 bacterial isolate was effectively



0.05

Figure 3. Evolutionary relationship analysis, by the neighbor-joining method, of 21 known CAs from prokaryotes, in comparison to *Enterobacter* sp. B13-CA (accession numbers are given in parenthesis). Scale bar represents 0.05 substitutions per nucleotide position.

#### DOI: 10.3109/14756366.2015.1100176

Figure 4. Multiple amino acid sequence alignment of 21 known β-class CAs: Enterobacter sp. B13-CA, E. coli str. K-12 substr. MG1655can, Ralstonia eutropha can, Enterobacter sp. RS1 yadF, Klebsiella pneumoniae can, H. influenzae can, C. freundii yadF, P. aeruginosa PAO1 cynT, P. aeruginosa PAO1 CA, Synechococcus elongates PCC 7942 icfA, B. suis 1330 CA, E. coli str. K-12 substr. MG1655 cynT, M. thermautotrophicus cab, S. enterica mig-5, P. carotovorum cah, H. pylori cynT, H. influenzae Rd KW20 CA, Burkholderia pseudomallei CA, P. gingivalis CD, V. cholerae CA, S. pneumoniae GA17570 CA and M. tuberculosis Rv1284. Triangles indicates the zinc ion-binding residues. Other active side conserved residues were shaded in blue (red letters). The figure was drawn with ESPript.

	0, 1		-	
	20 30	40	50 6	50
Ent_spBI3_CA	EEDPGFFGKLAQA	ONPRE LWIGSDER	VP. AERLTGLEPCE	PEVHR
E.COLL_CAN	EEDPGFFERLAQA	OKPRF LWIGSDSR	VP. AERLTGLEPCE	LEVHR
R.eutropha_can	AEDPTFFMRLANQ	CAPEYLWIGSDER	VP ANQILGLAPER	VEVHR
Ent_spksi_yadr	EEDPGFFERLAQA	ONPRE LWDG SDSR	VP. AERLTGLEPER	LEVHR
K.pneumoniae_can	EEDPGFFERLSQT	CRERFLWIG SISR	VP. AERLTGLEPGE	
A.Influenzae_can	EENSTIFKELADH	OND DELWIG SPSR	VP. AERLINLEPER	
C. Freundii_yadr	EEDPGFFETLTHA	ONPRELWIG SSER		
P.aeruginosa_PAO1_CynT	PARSQLERSLATE	CAPRALFIA SISK	VVPELLIQREPER	
P.aeruginosa_PAOI_CA	QEDPDIFAKLARQ	OUDDUI DI DOCO	TP DNI TROCCMC	TUTT
S.elongatus_PCC_/942_ICIA	FYDDEVECDIICC	ODDEELMIC COODD		
B.Suis_1330_CA	EXDPETESKLLSS			TUVIR
M thermoutet rephicus ash	NODEBEBDI CDI	LEDVI CITTOMOCO		
M. thermautotrophicus_cab	NODERERDLSDLK	HSPRICIIT M SR		ARVIK
S.enterica_mig-5	HDILAQKENSIAG	COTNTDON NO		TINSR
P. carocovorum_can	LSPDFSLCETGKN	C C C C C C C C C C C C C C C C C C C		NNGHI
H. pylori Cynt	DIASGIKGPLPMP	P S HIAIVA M AR		ANVIR
H.Influenzae_Rd_Kw20_CA	LENSTIFKELADH	OT PHILWIG SISK	VP. AEKLINLEP E	
B.pseudomailei_CA	ADDPQIFSRLADQ	CAPEILWIGSSER	VP. ANQIIGLPPE	VEVHR
P.gingivalis_CD	RDLNAQAVAGLEG	OFFEATILS		
V. Cholerae_CA	AETPEYFARLARG	ONPDF LWIG ADSR	VP. AERLIGLYSCH	LIVHR
S.pneumoniae_GA17570_CA	YVALHGQLNLPLK	<b>PKTRVAIVT<u>emps</u>r</b>	LH VAQALGLALGL	DAHILR
	70	80	90 100	
Ent en B13 CA	NVANLUTHTOLN	CLSVVOVAVDV	EVENTITIC	GVOAA
E coli can	NVANLUTHTDIN.	CLSVVOVAVDV		
B eutropha can	NTANUTAUCDIN.	CHOVY ZINVDV	FVFUTTTCCIVCIC	GUODA
Ent en RS1 vadE		ALAVIOTAVEV	EVENIIICG YG G	GVQAA
Line_spnor_yade	NUANI, UTHTOIN	ALAVIOFAVEV	LEVENIIICGHYGOG LKVRHITVVGHYGOG	GVQAA GVKVA
V proumonine can	NIANVIAHSDLN. NVANLVIHTDLN.	ALAVIOFAVEV	LEVEHILICG YGGG LKVRHITVVG YGGG LEVEHILICG YGGE	GVQAA GVKVA
K.pneumoniae_can	NVANLVIHTDLN. NVANLVIHTDLN. NVANLVIHTDLN.	ALAVIOFAVEV CLSVVQYAVDV CLSVVQYAVDV CLSVVQYAVDV	LEVEHILICGHYGC LEVEHILICGHYGC LEVEHILICGHYGC LEVEHILICGHYGC LEVEHILICGHYGC	GVQAA GVKVA GVQAA
K.pneumoniae_can H.influenzae_can C.freundii uadF	NIANVIAHSDLN. NVANLVIHTDLN. NVANLVIHTDLN. NVANQVIHTDFN.	ALAVIQEAVEV CLSVVQYAVDV CLSVVQYAVDV CLSVVQYAVDV CLSVVQYAVDV	EVEHILICGIYG LEVEHILICGIYG LEVEHILICGIYG LEVEHILICGIYG LEVEHILICGIYG LKIEHILICGIYG	GVQAA GVKVA GVQAA GIHAA
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K.pneumoniae_can H.influenzae_can C.freundii_yadF P.aeruginosa_PAO1_CynT B.aeruginosa_PAO1_C2	NVANLVIHTDLN. NVANLVIHTDLN. NVANLVIHTDLN. NVANLVIHTDLN. NAGNIVPGYGPQP	ALAVIQ AVEV CLSVVQ AVDV CLSVVQ AVDV CLSVVQ AVDV CLSVVQ AVDV G.GVSASVE AVAV	EVEHIIICG YG EVEHIIICG YG EVEHIIICG YG KIEHIIICG YG KIEHIIICG YG GVGDIVVCG SD GVGDIVVCG SD	GVQAA GVKVA GVQAA GIHAA GVQAA AMG.A
K.pneumoniae_can H.influenzae_can C.freundii_yadF P.aeruginosa_PAO1_CynT P.aeruginosa_PAO1_CA C.cloratur_DCG_7042_icfb	NVANLVIHTDLN. NVANLVIHTDLN. NVANLVIHTDLN. NVANLVIHTDFN. NAGNIVPGYGPQP NVANVVIHTDLN.	. ALAVIQ AVEV CLSVVQ AVDV CLSVVQ AVDV CLSVVQ AVDV CLSVVQ AVDV G.GVSASVE AVAV CLSVIQ AVDV	EVEHIIICG YG C EVEHIIICG YG C EVEHIIICG YG C KIEHIIICG YG C KIEHIIICG YG C GVGDIVVCG YG GVGDIVVCG YG KVKHILVTG YG C	GVQAA GVKVA GVQAA GIHAA GVQAA AMG.A GVRAS
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	110	120	130	140
Ent_spB13_CA	VENTELG L	IDNWLLHIRDI	WFKHSSLLGEME	.QERRLDTLCEL
E.coli_can	VENPELGL	INNWLLHIRDI	WFKHSSLLGEME	.QERRLDTLCEL
R.eutropha_can	LKRERIG L	<b>ADN<mark>W</mark>LRHVRDV</b>	ADKHEAYLGTLI	REDDAHTRLCEL
Ent_spRS1_yadF	NPELG L	IDNWLLHIRDI	WFKHSSLLGEME	.QERSLDTLCEL
K.pneumoniae_can	VENPELG L	IDNWLLHIRDI	WFKHSSLLGEME	. EDRRLDTLCEL
H.influenzae_can	MADKDLG L	INNWLLHIRDI	WFKHGHLLGKLS	S. PEKRADMLTKI
C.freundii_yadF	IENKEQG L	IDNWLLHIRDI	WFKHSSLLGEME	. QERRMDTLCEL
P.aeruginosa_PAO1_CynT	IASCACLDQLPA	VAG <mark>W</mark> LHHAEAA	R. AMNSAHEHAS	. EAARLDALVRH
P.aeruginosa_PAO1_CA	LHNDQLGL	IDG <mark>W</mark> LRSIRDL	AYEYREHLEQLE	TEEERVDRLCEL
S.elongatus_PCC_7942_icfA	LKLNQLQEDMPL'	VYD <mark>W</mark> LQHAQAT	RRLVLDNYSGYE	. TDDLVEILVAE
B.suis_1330_CA	MDGYGHGI	IDNWLQPIRDI	AQANQAELDTIE	ENTQDRLDRLCEL
E.coli_cynT	IASCQCMDHMPA	<b>VSHWLRYADSA</b>	R. VVNEARPHSI	. LPSKAAAMVRE
M.thermautotrophicus_cab	EDLIVSR	MRE	LGVEEEVIENFS	SIDVLNPVGDEEE
S.enterica_mig-5	IDNAELG N	LTG <mark>L</mark> LDEIKPA	IAKTEYSGERKO	SNYDFVDAVARK
P.carotovorum_cah	NADGALT	VIA <mark>L</mark> MFQEGAA	NPQLATAWQQIE	ARVDQAEDVRTP
H. pylori CynT	DDDFKR	<mark>.</mark> <del>.</del>	AIQDETGIRPT	WS.PESYPDAVE
H.influenzae_Rd_KW20_CA	MADKDLG L	INNWLLHIRDI	WFKHGHLLGKLS	S. PEKRADMLTKI
B.pseudomallei_CA	LHNRRVG L	ADN <mark>W</mark> LHHVQDV	REKHAALLEDWE	LGEARYRRLIEL
P.gingivalis_CD	IKGVEMGN	ITSLMEEIKPS	VEATQYMGERTY	ANKEFADAVVKE
V.cholerae_CA	IDNPQLG L	INNWLLHIRDY	YLKHREYLDKME	AEDRSDKLAEI
S pneumoniae GA17570 CA	NEPFOE		YLKEELGVDVS	DODFLPFODIEE

	150	160	170	180	190
Ent_spB13_CA	NVMEQVYN	LGHSTIMQS	AWKRGQKVSI	HGWAYGIH	DGLLRNLEVTAT
E.coli_can	NVMEQVYN	LGHSTIMQS	AWKRGQKVTI	HGWAYGIH	DGLLRDLDVTAT
R.eutropha_can	NVIEQVNN	VCQTTVLQD	AWSRGQAVTVI	IGWVYG <mark>VS</mark>	DGLLRDLGMAAS
Ent_spRS1_yadF	NVMEQVYN	LGHSTIMRS	AWKRGQKVSVI	HGWAYGIH	D <mark>GLL</mark> RNLYVTAT
K.pneumoniae_can	NVMEQVYN	LGHSTIMQS	AWKRGQKVTI	HGWAYGIH	DGLLRDLDVTAV
H.influenzae_can	NVAEQVYN	LGRTSIVKS	AWERGQKLSL	HGWVYDVN	DGFLVDQGVMAT
C.freundii_yadF	NVMEQVYN	LGHSTIMRS	AWKRGQKVTI	HGWAYGIH	DGLLRDLDVTAT
P.aeruginosa_PAO1_CynT	NVIAQLAN	LRTHPCVAR	ALEQGR . LNL	HGWVYDIE	SGRIDALDGAS.
P.aeruginosa_PAO1_CA	NVIQQVAN	VSHTSIVQN	AWHRGQSLSVI	HGCIYGIK	D <mark>GLW</mark> KNLNVTVS
S.elongatus_PCC_7942_icf#	NVLTQIEN	LKTYPIVRS	RLFQGK . LQII	GWI <mark>Y</mark> e <mark>ve</mark>	SGEVLQISRTSS
B.suis_1330_CA	SVSSQVES	LSRTPVLQS	AWKDGKDIIV	HGWMYNLK	DGLLRDIGCDCT
E.coli_cynT	NVIAQLAN	LQTHPSVRL	ALEEGR . IAL	HGWVYDIE	SGSIAAFDGAT.
M.thermautotrophicus_cab	NVIEGVKR	LKSSPLIP.	ESIGVI	HGLIIDIN	T <mark>GRL</mark> KPLYLDED
S.enterica_mig-5	NVELTIEN	IRKNSPVLK	QLEDEKKIKI	<b>JGSMYHLT</b>	G <mark>G</mark> K <mark>V</mark> EFFEV
P.carotovorum_cah	VAIQALLP	TSLN.YYRF	SGSLTTPPCSI	EGIRWLVL	D <mark>H</mark> P <mark>V</mark> TASAEQIS
H. pylori CynT	DVRQSLRR	IEVNPFVT.	KHTSLI	RGFVFDVA	T <mark>G</mark> KLNEVTP
H.influenzae_Rd_KW20_CA	NVAEQVYN	LGRTSIVKS	AWERGQKLSL	IGWVYDVN	D <mark>GFL</mark> VDQGVMAT
B.pseudomallei_CA	NVIEQVVN	VCRTTIVND	AWARGQPLTVI	HALVYGVH	D <mark>GRM</mark> RNLGMAVS
P.gingivalis_CD	NVIQTMDE	IRRDSPILK	KLEEEGKIKI	GAIYEMS	T <mark>G</mark> K <mark>V</mark> HFL
V.cholerae_CA	NVAEQVYN	LANSTVLQN	AWERGQAVEVI	HGFVYGIE	DGRLEYLGVRCA
S.pneumoniae_GA17570_CA	SVREDMQL	LIESPLIP.	DDVIIS	SGAI <mark>Y</mark> N <mark>VD</mark>	TGSMTVVEL

promoting CaCO<sub>3</sub> precipitation, presumably due to the CA activity present in the bacterium (data not shown). The EzTaxon identification results of 16S rDNA sequence of the B13 strain showed that it possesses a high percentage similarity with bacteria belonging to Enterobacteriaceae. Among various such bacteria, strain B13 was 99.04% identical to Enterobacter xiangfangensis strain 10-17, 98.63% to Enterobacter asburiae strain JCM6051, 98.56% to Enterobacter cloacae strain ATCC 13047, 98.45% to Enterobacter cowanii CIP 107300, 98.41% to Klebsiella michiganensis W14, 98.23% to Escherichia hermannii GTC 347, 97.90% to Escherichia vulneris ATCC 33821, to 97.89% Citrobacter youngae CECT 5335, to 97.77% Enterobacter

Table 1. Kinetic parameters for the CO<sub>2</sub> hydration reaction catalyzed by the human cytosolic isozymes hCA I and II ( $\alpha$ -class CAs)<sup>48</sup> and the bacterial  $\beta$ -CAs: PgiCAb (from *P. gingivalis*)<sup>28</sup>, HpyCA (from *H. pylori*)<sup>35</sup>, LpCA1 and LpCA2 (from *L. pneumophila*)<sup>24</sup> and B13-CA (from *Enterobacter* sp. B13).

Enzyme	Class	$k_{\text{cat}} (\text{s}^{-1})$	$k_{\text{cat}}/K_{\text{m}}$ (M <sup>-1</sup> ×s <sup>-1</sup> )	<i>K</i> <sub>I</sub> (acetazolamide) (nM)
hCA I	α	$2.0 \times 10^{5}$	$5.0 \times 10^{7}$	250
hCA II	α	$1.4 \times 10^{6}$	$1.5 \times 10^{8}$	12
PgiCAb	β	$2.8 \times 10^{5}$	$1.5 \times 10^{7}$	214
НруСА	β	$7.1 \times 10^{5}$	$4.8 \times 10^{7}$	40
LpCA1	β	$3.4 \times 10^{5}$	$4.7 \times 10^{7}$	76.8
LpCA2	β	$8.3 \times 10^{5}$	$8.5 \times 10^{7}$	72.1
B13-CA	β	$4.8  imes 10^5$	$5.6 \times 10^{7}$	78.9

All the measurements have been made at 20 °C, pH 8.3 by a stopped flow  $CO_2$  hydrase assay method<sup>19</sup>. Acetazolamide (AAZ) inhibition data are also provided.

*ludwigii* EN-119T, 97.73% to *Enterobacter soli* LF7a, 97.67% to *Cedecea davisae* DSM 4568, 97.63% *Cedecea neteri*, 97.60% to *Erwinia aphidicola* GTC 1688, 97.59% to *Enterobacter aerogenes* KCTC 2190 (Figure 1).

#### Purification of Enterobacter sp. B13-CA

The recombinant B13-CA was purified in concentrations ranging between 0.04 and 2.4 mg/ml from 250 ml cultures. The purified enzyme was subjected to SDS-PAGE analysis (Figure 2) which led to the observation of a distinct single band, with the predicted molecular mass of 24.95 kDa (deduced from the amino acid sequence). Data of the SDS-PAGE (Figure 2, lane 2) showed that the purified B13-CA had indeed a molecular weight (for the monomer) of 25 kDa.

#### Biochemical characterization of Enterobacter sp. B13-CA

According to the zymogram analysis, *Enterobacter* sp. B13 CA did not reveal any esterase activity. Furthermore, no esterase activity was observed by using a spectrophotometric assay with  $\alpha$ -naphthyl acetate as substrate (data not shown). This is to be expected, since esterase activity was not reported for any  $\beta$ -class CAs, although many putative activated esters have been investigated as possible substrates<sup>20</sup>. In fact only the  $\alpha$ -<sup>20,21</sup> and  $\eta$ -CAs<sup>22</sup> possess esterase activity with activated esters (such as 4-nitrophenyl- or  $\alpha$ -naphthyl acetate) as substrates.

#### **Bioinformatic analysis**

Considering the sequence of the can gene of Enterobacter sp. B13, which contains 663 bp, it encodes a protein of 220 amino acid residues. Based on DELTA-BLAST results of this amino acid sequence, Enterobacter sp. B13 can shows 33% homology with E. coli cynT and 95% with can (from the same bacterium, i.e. E. coli). Phylogenetic analysis of amino acid resemblance revealed that B13-CA is a member of the  $\beta$ -CA class and has a close relationship with can genes identified in other bacteria belonging to Enterobacteriaceae (Figure 3). A conserved domain database (CDD) (http://www.ncbi.nlm.nih.gov/Structure/cdd/ wrpsb.cgi) search of Enterobacter sp. B13 can sequences showed that the active and ion binding site residues of B13-CA are well conserved, as in all  $\beta$ -CAs investigated in detail, such as among others the enzymes from H. influenzae, Citrobacter freundii, Pseudomonas aeruginosa, Synechococcus elongatus, Brucella suis, E. coli, Methanothermobacter thermautotrophicus, Salmonella enterica, Pectobacterium carotovorum, Helicobacter

*pylori, Porphyromonas gingivalis, Vibrio cholerae, Streptococcus pneumoniae, S. mutans* and *Mycobacterium tuberculosis Rv1284*<sup>23-42</sup> (Figure 4).

#### Catalytic activity of B13-CA

In fact there are two types of  $\beta$ -CAs in organisms all over the phylogenetic tree, which are defined by their pH-catalytic activity profile as well as active site structural configuration<sup>41–52</sup>. Type I  $\beta$ -CAs display catalytic activity over a broad pH range (6.5–9.0) with the active site zinc tetrahedrally coordinated by three amino acid residues (two Cys and one His) and a hydroxide/water. In contrast, type II β-CAs are catalytically active only at a pH 8 and higher where they adopt a functional active site configuration like that of type I. In fact, below pH 8 they are conformationally selfinactivated by the addition of a fourth amino acid (an Asp residue) which coordinates to the Zn(II) ion as the fourth ligand, and displacing the zinc bound solvent<sup>39,52</sup>. At pH>8, the Asp copordinated to Zn(II) makes a salt pair interaction with a conserved Arg residue, which "opens" the active site in the sense that a water molecule/hydroxide ion takes the place of the Asp residue, generating thus the nucleophile responsible for the catalytic activity of these enzymes<sup>39,48–52</sup>.

As seen from Figure 4, where an alignment of the amino acid sequence of B13-CA with that of other 20 bacterial  $\beta$ -CAs is presented, the new enzyme described here has all the features of a catalytically effective  $\beta$ -CA: (i) the putative Zn(II) ligands, Cys42, Asp44 (in case B13-CA is a type II  $\beta$ -CA), His 98 and Cys101; (ii) The catalytic dyad involved in activation of the zinc-coordinated water molecule/hydroxide ion for catalysis<sup>53</sup>, represented by the Asp44–Arg46 residues, which resembles in a way the activation of the water molecule in aspartic proteases<sup>53</sup>.

As seen from Table 1, in which we compared the catalytic activity of the new enzyme reported here, B13-CA with that of other  $\alpha$ - and  $\beta$ -class CAs from various organisms, B13-CA shows a significant activity as catalyst for the hydration of  $CO_2$  with formation of protons and bicarbonate. Indeed, B13-CA possesses the following kinetic paramaters at 20 °C and pH of 8.3:  $k_{cat}$  of  $4.8 \times 10^5$  s<sup>-1</sup> and  $k_{cat}/K_m$  of  $5.6 \times 10^7$  M<sup>-1</sup> × s<sup>-1</sup>. In fact the catalytic activity of the new enzyme is in the same range as those of other  $\beta$ -CAs, such as those from *P. gingivalis* (PgiCAb)<sup>28</sup>, H. pylori (HpyCA)<sup>35</sup>, or Legionella pneumophila (LpCA1 and LpCA2)<sup>24</sup>, recently characterized by one of our groups. Furthermore, the enzyme was inhibited by the clinically used<sup>54–57</sup> sulfonamide CA inhibitor (CAI) acetazolamide (AAZ) with an inhibition constant of 78.9 nM, in the same range as the human (h) hCA I isoform, HpyCA, LpCA1 and LpCA2 (which were inhibited with inhibition constants ranging between 40 and 250 nM). Only the physiologically dominant isoform hCA II was better inhibited by this drug, with a  $K_I$  of 12 nM (Table 1).

#### Conclusions

A recombinant CA (EC 4.2.1.1) from the soil-dwelling bacterium *Enterobacter* sp. B13 was cloned and purified by Co<sup>2+</sup> affinity chromatography. Bioinformatic analysis showed the new enzyme (denominated here B13-CA) to belong to the  $\beta$ - CA class and to possess 95% homology with the ortholog enzyme from *E. coli* encoded by the can gene, whereas its sequence homology with the other such enzyme from *E. coli* (encoded by the *cynT* gene) was of 33%. B13-CA was characterized kinetically as a catalyst for carbon dioxide hydration to bicarbonate and protons. The enzyme showed a significant catalytic activity, with the following kinetic paramaters at 20 °C and pH of 8.3:  $k_{cat}$  of  $4.8 \times 10^5 \text{ s}^{-1}$  and  $k_{cat}/K_{m}$  of  $5.6 \times 10^7 \text{ M}^{-1} \times \text{s}^{-1}$ . This activity was potently inhibited by acetazolamide which showed a  $K_{I}$  of 78.9 nM. Although only this compound was investigated for the moment as

B13-CAI, further studies may reveal new classes of inhibitors/ activators of this enzyme<sup>58–60</sup>, which may show biomedical or environmental applications.

#### **Declaration of interest**

The authors do not have any conflicts of interest.

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