

RESEARCH ARTICLE

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Assessment of databases to determine the validity of β - and γ -carbonic anhydrase sequences from vertebrates



Reza Zolfaghari Emameh^{1*} , Marianne Kuuslahti², Hassan Nosrati³, Hannes Lohi^{4,5,6} and Seppo Parkkila^{2,7}

Abstract

Background: The inaccuracy of DNA sequence data is becoming a serious problem, as the amount of molecular data is multiplying rapidly and expectations are high for big data to revolutionize life sciences and health care. In this study, we investigated the accuracy of DNA sequence data from commonly used databases using carbonic anhydrase (CA) gene sequences as generic targets. CAs are ancient metalloenzymes that are present in all unicellular and multicellular living organisms. Among the eight distinct families of CAs, including α , β , γ , δ , ζ , η , θ , and ι , only α -CAs have been reported in vertebrates.

Results: By an in silico analysis performed on the NCBI and Ensembl databases, we identified several β - and γ -CA sequences in vertebrates, including *Homo sapiens*, *Mus musculus*, *Felis catus*, *Lipotes vexillifer*, *Panholops hodgsonii*, *Hippocampus comes*, *Hucho hucho*, *Oncorhynchus tshawytscha*, *Xenopus tropicalis*, and *Rhinolophus sinicus*. Polymerase chain reaction (PCR) analysis of genomic DNA persistently failed to amplify positive β - or γ -CA gene sequences when *Mus musculus* and *Felis catus* DNA samples were used as templates. Further BLAST homology searches of the database-derived “vertebrate” β - and γ -CA sequences revealed that the identified sequences were presumably derived from gut microbiota, environmental microbiomes, or grassland ecosystems.

Conclusions: Our results highlight the need for more accurate and fast curation systems for DNA databases. The mined data must be carefully reconciled with our best knowledge of sequences to improve the accuracy of DNA data for publication.

Keywords: Carbonic anhydrase, Contamination, Curation, Database, DNA, Sequencing

Background

Carbonic anhydrases (CAs) are metalloenzymes that are classified into eight evolutionarily distinct families, including α , β , γ , δ , ζ , η , θ , and ι [1–4]. These enzymes catalyze the hydration of carbon dioxide to bicarbonate and protons and are involved in various biochemical pathways, such as gluconeogenesis, ureagenesis and photosynthesis, and other physiological functions, such

as pH homeostasis, electrolyte transfer and calcification [5].

There are 12 α -CA isozymes, including CA I-IV, CA VA and VB, CA VI, CA VII, CA IX, and CA XII-XIV, that are expressed in humans [6]. Interestingly, CA XV is the only active CA isozyme known to date that is expressed in several vertebrate species but is lost in human and chimpanzee genomes [7]. In addition to the 13 mammalian α -CA isozymes, there are three acatalytic CA-related proteins (CARPs), including CARP VIII, CARP X, and CARP XI, with crucial physiological roles [8–11]. α -CAs have been reported from many organisms, including both prokaryotes and eukaryotes [12].

* Correspondence: zolfaghari@nigeb.ac.ir

¹Department of Energy and Environmental Biotechnology, National Institute of Genetic Engineering and Biotechnology (NIGEB), Tehran 14965/161, Iran
Full list of author information is available at the end of the article



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Although β -CAs are present in archaea, bacteria, plants, fungi, protozoans, and insects, there are no reports of β -CAs in any vertebrate species [13, 14]. Similarly, γ -CAs are present in many prokaryotes and eukaryotes, such as plants and fungi, whereas they do not exist in any vertebrates according to the current knowledge [15, 16]. Incomplete β -CA gene sequences have been identified in the genome of the cephalochordate *Branchiostoma floridae* (the Florida lancelet), but whether they represent a pseudogene or an incompletely sequenced active gene has not been determined [17]. Some annotated β - and γ -CA sequences present in databases have been linked to vertebrate genomes, but in fact, they might have originated from either gut microbiota or other normal flora or even from environmental bacterial contamination. Kraken and Taxoblast are two recently designed ultrafast programs to identify contaminant DNA sequences from metagenomic and genome sequencing databases [18, 19]. The main limitation of both methods is the lack of accessibility to a computer or server with enough RAM for quick operation while performing genome blast homology searches.

In this study, we first searched for β - and γ -CAs in vertebrates using in silico tools. The results obtained from the NCBI and Ensemble databases led us to perform polymerase chain reaction (PCR) amplifications using mouse and cat genomic DNA as templates. The results indicated that the “vertebrate” β - and γ -CA sequences detected from databases were presumably derived from gut microbiota, environmental microbiomes, or grassland ecosystems. This finding emphasizes the importance of fast and accurate biocuration of database sequences.

Results

Identification of β - and γ -CAs

The BLASTP program from the NCBI database identified β -CA protein sequences from some vertebrates, including *Lipotes vexillifer* (XP_007454654.1), *Pantholops hodgsonii* (XP_005974256.1), *Homo sapiens* (SJM31717.1), and *Oncorhynchus tshawytscha* (XP_024266887.1). In addition, the TBLASTN program of Ensembl genome browser 95 identified the genomic location for a β -CA gene in *M. musculus*, strain NOD/ShiLtJ (genomic location: LVXS01065484.1: 870–1430), *Hippocampus comes* (genomic location: LVHJ01039623: 18–230*), and *Hucho hucho* (genomic location: QNTS01034426:189–644*). The aforementioned methods identified γ -CA protein sequences from some vertebrates, including *L. vexillifer* (XP_007452618.1), *P. hodgsonii* (XP_005961532.1), *H. sapiens* (SJM34589.1), *F. catus* (XP_004001159.1), and *Rhinolophus sinicus* (XP_019578089.1). Additionally, the genomic location was identified for a γ -CA gene in *Xenopus tropicalis*

(genomic location: GL180697.1: 4765–5075) and *H. comes* (genomic location: LVHJ01047219:4–240*) (Fig. 1 and Table 1). The multiple sequence alignment (MSA) analysis showed that the predicted polypeptide sequences would contain highly conserved amino acids, which are considered important for the classical β -CA (Fig. 2) and γ -CA (Fig. 3) enzymes.

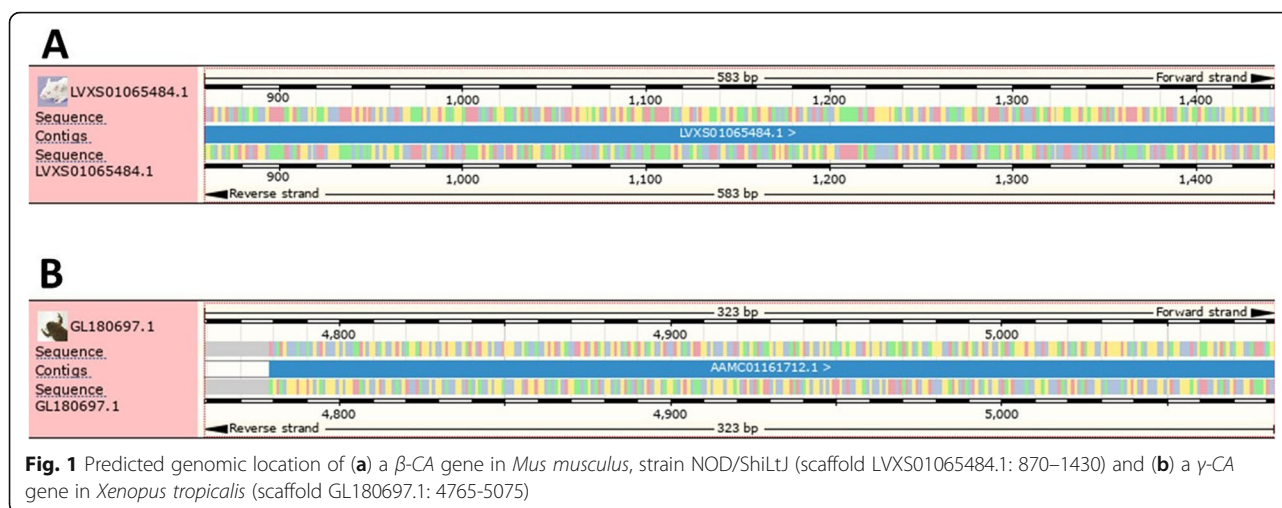
Our further analysis revealed that the genomic organization of the coding genes for the “vertebrate” β - and γ -CA proteins was consistent with the single exonic pattern of coding genes in prokaryotes. In addition, the BLAST homology search analysis decrypted the high percentage of identities (73–100%) between the predicted β - and γ -CA protein sequences of vertebrates and some other organisms, which mostly involved prokaryotic species (Table 1).

Molecular analysis of β - and γ -CA genes from vertebrates

To investigate whether β -CA or γ -CA genes are truly present in vertebrate genomes, we performed PCR using DNA samples extracted from ear punching specimens of *M. musculus* and whole blood of *F. catus*. The first round PCRs with low stringent conditions showed some positive signal for the primer pairs P1 and P3 of *F. catus* and P5 and P8 of *M. musculus* (Fig. 4a). Estimation of the PCR product size was conducted based on the product length from Table 2. Because the signal remained weak in most cases, we performed the second round PCR using the PCR amplicons from the first round PCR as templates. The results of the second round of PCR are shown in Fig. 4b. The sequencing results revealed that none of the sequenced PCR products represented the predicted β -CA gene from *M. musculus* or the γ -CA gene from *F. catus*.

Discussion

CA genes are widely distributed in species of all life kingdoms. Despite this general concept, β - and γ -CA genes have never been reported in vertebrate genomes to the best of our knowledge based on previous literature. Our survey on the β - and γ -CA gene sequences of vertebrates presented in public databases in 2017–2020 revealed, however, that some sequences were or are still available, such as β -CA genes from *L. vexillifer* and *M. musculus*, as well as γ -CA genes from *L. vexillifer*. Some data were removed in 2019–2020, such as β -CA genes from *P. hodgsonii* and *H. sapiens*, as well as γ -CA genes from *P. hodgsonii*, *X. tropicalis*, *H. sapiens*, *F. catus*, and *R. sinicus*. Some new sequences appeared and were annotated on databases in 2019–2020, including β -CA genes from *H. comes*, *H. hucho*, and *O. tshawytscha*, as well as the γ -CA gene from *H. comes*. At first glance, the reports of “vertebrate” β - and γ -CA genes in databases raised our interest as a potentially novel discovery, but

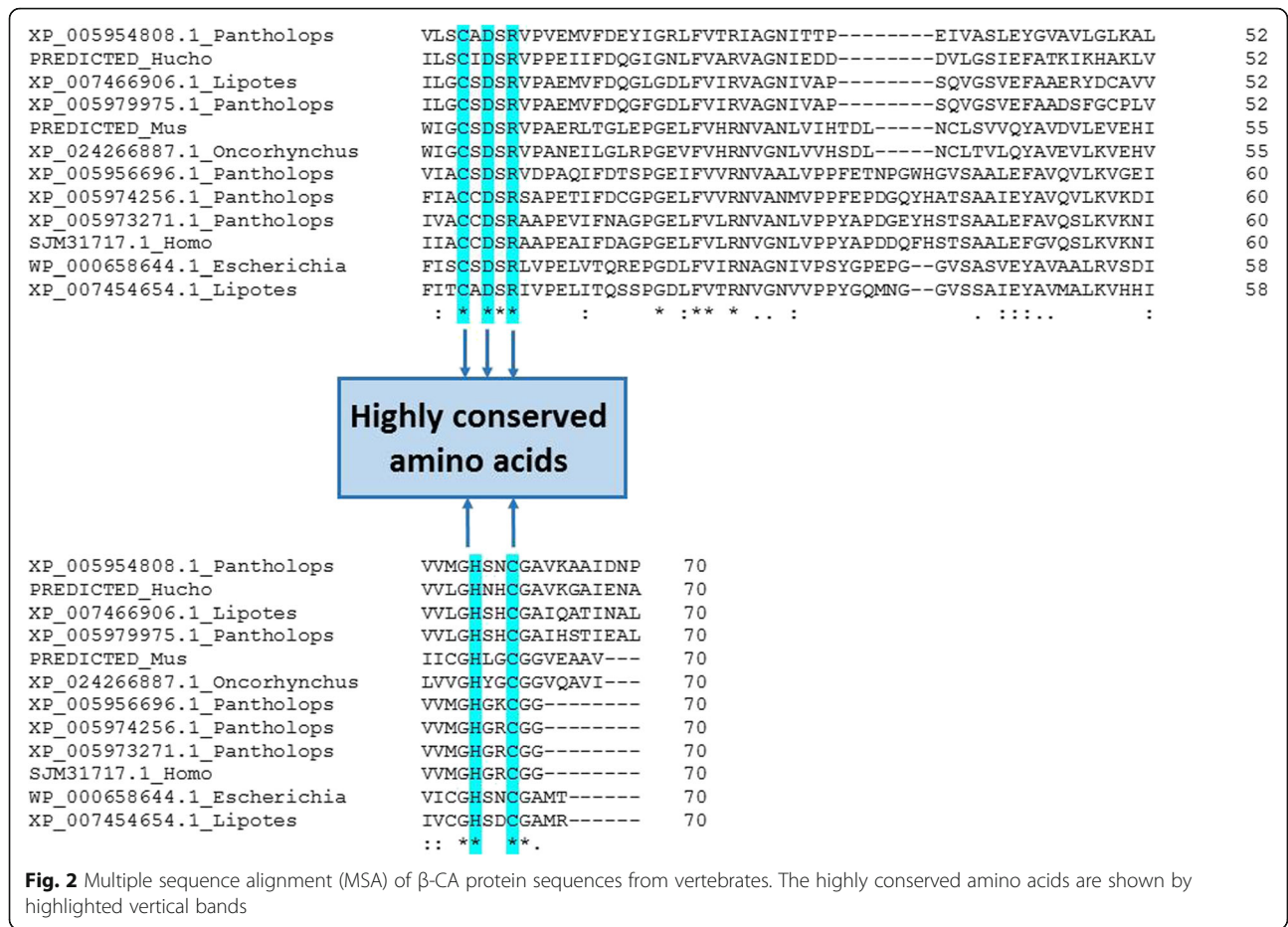
**Table 1** Identified β - and γ -CAs from vertebrates

Type of CA	NCBI IDs	Vertebrate species	Status in database		73–100% identical species	Exon count
			2017–2018	2019–2020		
β-CA	XP_007454654.1	<i>Lipotes vexillifer</i> (extinct Yangtze River dolphin)	A	A	<i>Pseudomonas</i> sp.	1
	XP_007466906.1				<i>Acinetobacter</i> sp.	
	XP_005974256.1	<i>Pantholops hodgsonii</i> (Tibetan antelope)	A	D	<i>Agrobacterium</i> sp. <i>Rhizobium</i> sp.	1
	XP_005956696.1				<i>Sphingobium</i> sp.	
	XP_005973271.1				<i>Mesorhizobium</i> sp.	
	XP_005979975.1				<i>Acinetobacter</i> sp.	
	XP_005954808.1				<i>Sphingobium</i> sp.	
	LVXS01065484.1: 870–1430 ^a	<i>Mus musculus</i> , strain NOD/ShiLJ (house mouse)	A	A	<i>Serratia</i> sp.	ND
	SJM31717.1	<i>Homo sapiens</i> (Human)	A	D	<i>Mesorhizobium delmotii</i>	1
	LVHJ01039623:18–230 ^a	<i>Hippocampus comes</i> ^b (Tiger tail seahorse)	U	A	<i>Muricauda</i> sp. (87.3%)	ND
QNTS01034426:189–644 ^a	<i>Hucho hucho</i> (Huchen or Danube salmon)	U	A	<i>Flavobacterium</i> sp. (73.7%)	ND	
XP_024266887.1	<i>Oncorhynchus tshawytscha</i> (Chinook salmon)	U	A	<i>Hydrogenophaga</i> sp.	1	
γ-CA	XP_007452618.1	<i>Lipotes vexillifer</i> (extinct Yangtze River dolphin)	A	A	<i>Pseudomonas</i> sp.	1
	XP_007465530.1					
	XP_005974442.1	<i>Pantholops hodgsonii</i> (Tibetan antelope)	A	D	<i>Caulobacter</i> sp.	1
	XP_005977566.1				<i>Delftia</i> sp. (98%)	
	XP_005974267.1				<i>Acinetobacter</i> sp.	
	GL180697.1: 4765–5075 ^a	<i>Xenopus tropicalis</i>	A	D	Comamonadaceae bacterium	ND
	SJM34589.1	<i>Homo sapiens</i> (Human)	A	D	<i>Mesorhizobium delmotii</i>	1
	XP_004001159.1	<i>Felis catus</i> (domestic cat)	A	D	<i>Acidovorax</i> sp. (97%)	1
	XP_019578089.1	<i>Rhinolophus sinicus</i> (Chinese rufous horseshoe bat)	A	D	<i>Brassica</i> sp. (94%)	1
	LVHJ01047219:4–240 ^a	<i>Hippocampus comes</i> (Tiger tail seahorse)	U	A	Bacteroidetes bacterium (93.7%)	ND

Abbreviations: ND Not defined, A Available, D Discontinued, U Unavailable (Supplementary file 1)

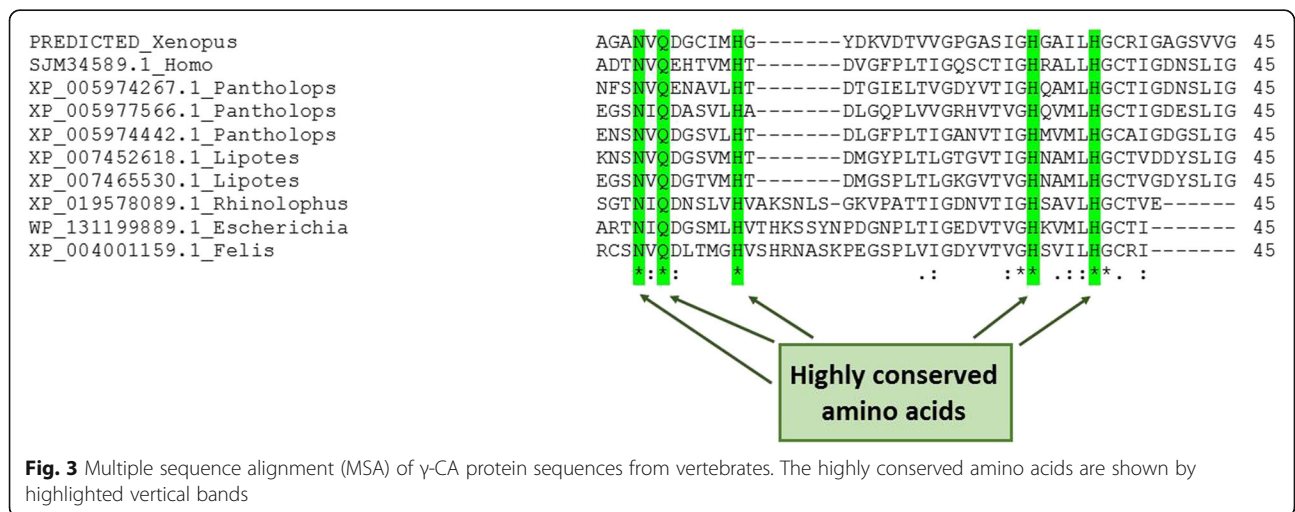
^a: Genomic location in the Ensembl genome browser 95

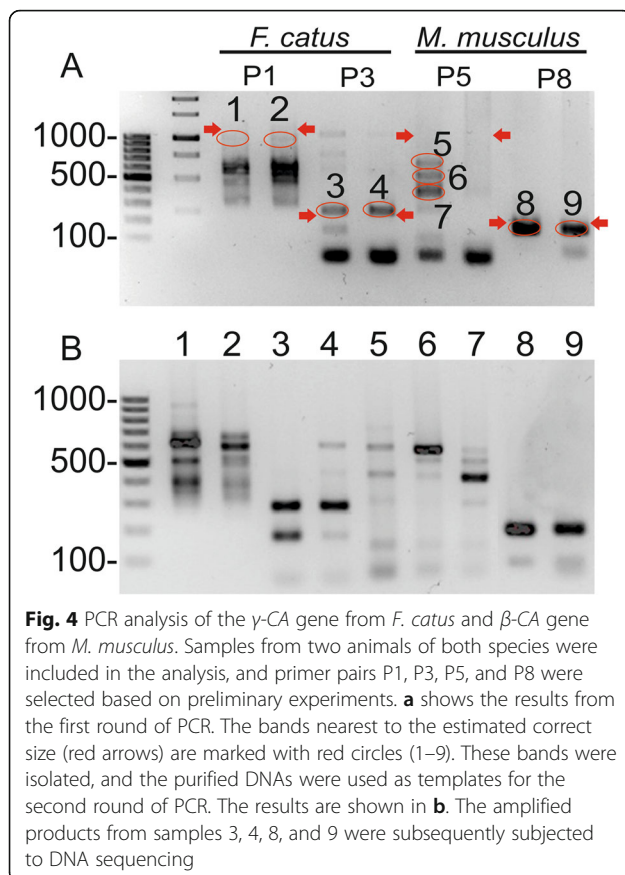
^b: The sequencing shows only the first highly conserved sequence (CXDXR)



enthusiasm gradually dissipated as most data were discontinued in 2019–2020. The BLAST homology search analysis of the predicted “vertebrate” β - and γ -CA protein sequences filtered with the “prokaryota” keyword defined that the discontinued β - and γ -CA genes belonged to prokaryotes. The most striking false-positive

sequences in databases were originally annotated as human β - and γ -CAs, which we defined by the BLAST homology search as *Mesorhizobium delmotii* enzymes instead of human origin (Table 1). Our results suggest that the predicted “human” β - and γ -CAs were derived from bacterial contamination of human DNA samples





that caused false interpretation during sequencing. As a sign of improved accuracy, these false-positive data were removed from databases in 2019–2020.

Another piece of evidence for the bacterial contamination of DNA samples is the contamination of *H. comes* sample with *Muricauda* sp. and *Bacteroides* sp., both of which are abundantly present in seawater sediments [20, 21]. In addition, DNA samples of salmon fishes (*H. hucho* and *O. tshawytscha*) can be contaminated with gut microbiota or egg-associated bacterial species, such as *Flavobacterium* sp., *Pseudomonas* sp., and *Hydrogenophaga* sp. [22, 23]. *Comamonadaceae* bacterium from gut microbiota may represent the main source of bacterial contamination for the DNA samples of *X. tropicalis* [24]. Notably, due to the living habitat of *R. sinicus* in meadows, scrubs, and grasslands and feeding in these important ecosystems, the contamination of the bat DNA sample was mainly derived from plant species, such as *Brassica* sp. (cruciferous vegetables), instead of contamination from gut microbiota.

The exon count of the predicted “vertebrate” β - and γ -CA genes suggested the presence of only a single exon in each case. This finding also supported the idea that prokaryotes from gut microbiota and environmental microbiome are the major source of contaminants that led to unexpected sequencing results from vertebrate DNA samples [25]. This idea was further supported by our PCR analysis of both mouse and cat genomic DNA samples combined with DNA sequencing, which consistently failed to identify any β - or γ -CA sequences in mice and cats.

Table 2 Designed primers for the β - and γ -CA genes

CA family	Vertebrate species	Primer pairs	Product length (bp)	
γ -CA	<i>Felis catus</i> (cat)	P1	Forward: 5'- AGATAACTACTTCACATCTGACA -3' Reverse: 3'- ATACAGGGCTGGGTGCCT -5'	1089
		P2	Forward: 5'- GGTGATTGGCGACTACGTGA - 3' Reverse: 3'- CTCAGTCGGTTAGGTGGCTG - 5'	625
		P3	Forward: 5'- GCGCGTGAAGAACAACCTACC - 3' Reverse: 3'- GTGTTCAAGTTCGCTCATCGG - 5'	217
		P4	Forward: 5'- AAGCGGCAACCTCTACATCG -3' Reverse: 3'- CGTGAGGTAGGCAGTAGACG -5'	341
β -CA	<i>Mus musculus</i> (Mouse)	P5	Forward: 5'- TGATAATGCCGATGGTCGTG -3' Reverse: 3'- AGTAGCCATGGCCTTGCGAT -5'	1023
		P6	Forward: 5'- TGGATTTTCCGGCACCGTTA -3' Reverse: 3'- CGGGTCTTCTTGTGATGT -5'	441
		P7	Forward: 5'- ACATCAGCAAGGAAGACCCG -3' Reverse: 3'- CACAATACGTCAAGGCGCTG -5'	391
		P8	Forward: 5'- GCTGCACATCCGTGATCTCT -3' Reverse: 3'- GGATCCCATACCCCAACCG -5'	191

It is clear that a significant amount of incorrect sequence data on both β -CA and γ -CA genes remain in public databases. Some existing examples are β -CA genes of *L. vexillifer*, *M. musculus*, *H. comes*, *H. hucho*, and *O. tshawytscha* and γ -CA genes of *L. vexillifer* and *H. comes*. The present findings highlight the importance of database curation efforts to achieve a higher degree of accuracy within a shorter revision time.

Conclusions

Online databases are important sources of information for mining genomic and proteomic data of living organisms. Unfortunately, these databases also include misannotated data to some extent due to microbial or other contamination. We used β - and γ -CA gene sequences as bioinformatic tools to demonstrate such contamination in various species. Our findings emphasize the importance of fast and reliable curation for achieving better quality and more accurate genomic and proteomic data.

Methods

Identification of β - and γ -CAs

In the first step, the β - and γ -CA protein sequences from *Escherichia coli* (NCBI IDs: WP_000658644.1 and WP_131199889.1, respectively) were used as the query in the Basic local alignment search tool (BLAST) for sequence similarity search analysis through the BLASTP program (<https://blast.ncbi.nlm.nih.gov/Blast.cgi?PAGE=Proteins>) of NCBI database [26] and TBLASTN program of Ensembl genome browser 95 (<https://asia.ensembl.org/Multi/Tools/Blast?db=core>) [27]. We filtered the results using “vertebrata” as the organism name, in which the BLASTP program only searched for β - and γ -CA protein sequences within vertebrates. Additionally, we applied the scientific name of defined vertebrates as the filter in the TBLASTN program of Ensembl genome browser 95. The obtained β - and γ -CA protein sequences were aligned using the Clustal Omega algorithm (<https://www.ebi.ac.uk/Tools/msa/clustalo/>) [28].

In the second step, we performed a BLAST homology search analysis on the obtained β - and γ -CA protein sequences from vertebrates, in which the results were filtered against “prokaryota” as the organism name. Afterward, exon counts were performed to detect β - and γ -CA gene sequences from vertebrates through the gene analysis program of the NCBI database.

Molecular analysis of β - and γ -CA genes from vertebrates

We designed eight primer pairs using Primer-BLAST for molecular detection of the β -CA gene from *Mus musculus* (Mouse) and the γ -CA gene from *Felis catus* (cat) (four primer pairs for each CA gene) identified through bioinformatic methods (Table 2) [29].

The ear blood samples of one *M. musculus* and 1 ml EDTA-blood samples of one privately-owned *F. catus* were collected under the permission of the animal ethical committee of the County Administrative Board of Southern Finland (ESAVI/8321/04.10.07/2017 for the mouse and ESAVI/7482/04.10.07/2015 for the cat) for molecular detection of the predicted β -CA gene of *M. musculus* and γ -CA gene of *F. catus*. In the Tampere University’s animal facility, mice are routinely earmarked and the same samples were used for genotyping purposes in another project. Written consents were collected from the participating cat owners and samples were collected as a part of the ongoing feline genetic research at Dr. Lohi’s laboratory. Cats visited a veterinary clinic for a routine sample collection. Genomic DNA was extracted from white blood cells using a semiautomated Chemagen extraction robot (PerkinElmer Chemagen Technologie GmbH, Baeswieler, Germany) according to the manufacturer’s instructions. The DNA concentrations were measured using a Qubit fluorometer (Thermo Fisher Scientific, Waltham, Massachusetts, USA) and a Nanodrop ND-1000 UV/Vis Spectrophotometer (Nanodrop Technologies, Wilmington, Delaware, USA), and samples were stored at -20°C . Polymerase chain reaction (PCR) was performed according to the protocol used by Zolfaghari Emaeh R et al. [30]. PCR amplification was run on a thermocycler (Bioer XP Cycler, Hangzhou Bioer Technology Co. Ltd., Hangzhou, China) according to the following details: 95°C (3 min), [95°C (15 s), 60°C (15 s), 72°C (15 s)] \times 40 cycles, 72°C (2 min). The amplified products were run on a 1.6% agarose gel and purified using a NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel). The second round of PCR was run as previously described, and the selected PCR amplicons (Fig. 4; samples 3, 4, 8, and 9) were treated with Exo I and Fast AP enzymes and sequenced using ABI PRISM BigDye[®] Terminator v3.1 Cycle Sequencing kit and 3500xL Genetic Analyzer (Applied Biosystems, Inc., Foster City, CA, U.S.A.).

Supplementary information

Supplementary information accompanies this paper at <https://doi.org/10.1186/s12864-020-6762-2>.

Additional file 1. Supplementary file 1

Abbreviations

BLAST: Basic Local Alignment Search Tool; CA: Carbonic anhydrase; CARP: CA-related protein; MSA: Multiple sequence alignment; NCBI: National Center for Biotechnology Information; PCR: Polymerase chain reaction

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Authors' contributions

RZE, HL, and SP designed the study. RZE carried out the search and data mining related to the β - and γ -CA sequences from vertebrates and bacteria from various databases. HN helped in the data mining and preparing the files. RZE performed the in silico studies and designed the experimental assays. MK performed the PCR experiments and sequencing. HL organized the collection of blood sample from cat and with SP helped in planning the experimental methods. By his expertise in genetics HL helped to reach the major conclusions of the study. RZE drafted the first version of the manuscript. All authors participated in writing further versions and read and approved the final manuscript.

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Availability of data and materials

No novel DNA, RNA, and protein sequence data related to β - and γ -CAs were produced in this study to be annotated in the databases. The analyzed datasets used in the current study were collected from NCBI database including XP_007454654.1, XP_007466906.1, XP_005974256.1, XP_005956696.1, XP_005973271.1, XP_005979975.1, XP_005954808.1, SJM31717.1, XP_024266887.1, XP_007452618.1, XP_007465530.1, XP_005974442.1, XP_005977566.1, XP_005974267.1, SJM34589.1, XP_004001159.1, and XP_019578089.1 as well as Ensembl genome browser including LVXS01065484.1: 870–1,430, LVHJ01039623:18–230, QNTS01034426:189–644, GL180697.1: 4,765–5,075, and LVHJ01047219:4–240.

Ethics approval and consent to participate

The ear samples of *M. musculus* and the blood samples of *F. catus* were collected under the permission of the animal ethical committee of the County Administrative Board of Southern Finland (ESAVI/8321/04.10.07/2017 for the mouse and ESAVI/7482/04.10.07/2015 for the cat). Written consent was collected from the participating cat owners.

Consent for publication

Not Applicable.

Competing interests

The authors declare that they have no conflict of interests.

Author details

¹Department of Energy and Environmental Biotechnology, National Institute of Genetic Engineering and Biotechnology (NIGEB), Tehran 14965/161, Iran. ²Faculty of Medicine and Health Technology, Tampere University, FI-33520 Tampere, Finland. ³Department of Materials Engineering, Tarbiat Modares University, Tehran, Iran. ⁴Department of Veterinary Biosciences, University of Helsinki, 00014 Helsinki, Finland. ⁵Department of Medical and Clinical Genetics, University of Helsinki, 00014 Helsinki, Finland. ⁶Folkhälsan Research Center, 00290 Helsinki, Finland. ⁷Fimlab Laboratories Ltd. and Tampere University Hospital, FI-33520 Tampere, Finland.

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