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## **OPEN** Three Distinct Glutamate **Decarboxylase Genes in Vertebrates**

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Gamma-aminobutyric acid (GABA) is a widely conserved signaling molecule that in animals has been adapted as a neurotransmitter. GABA is synthesized from the amino acid glutamate by the action of glutamate decarboxylases (GADs). Two vertebrate genes, GAD1 and GAD2, encode distinct GAD proteins: GAD67 and GAD65, respectively. We have identified a third vertebrate GAD gene, GAD3. This gene is conserved in fishes as well as tetrapods. We analyzed protein sequence, gene structure, synteny, and phylogenetics to identify GAD3 as a homolog of GAD1 and GAD2. Interestingly, we found that GAD3 was lost in the hominid lineage. Because of the importance of GABA as a neurotransmitter, GAD3 may play important roles in vertebrate nervous systems.

Glutamate decarboxylases (GADs) are essential for the conversion of glutamate to  $\gamma$ -aminobutyric acid (GABA), the predominant inhibitory neurotransmitter in central nervous systems<sup>1</sup>. GADs are members of the Group II pyridoxal-5'-phosphate-dependent decarboxylases, which includes decarboxylases that operate on several different substrates<sup>2</sup>. Two GAD proteins found in vertebrate species, GAD67 and GAD65, are encoded by the paralogous genes GAD1 and GAD2, respectively<sup>3</sup>. While both GADs synthesize GABA and are co-expressed in most vertebrate GABAergic neurons, GAD1 synthesizes cytoplasmic GABA that is used for extrasynaptic and metabolic purposes and GAD2 regulates the vesicular pool for release<sup>4-6</sup>. Nevertheless, GAD1 and GAD2 sequences are highly similar to each other, and they share a common intron-exon organization, indicating a common origin<sup>7</sup>.

The evolutionary history of GAD genes is long and diverse. Genes with homology to GAD arose before the evolution of eukaryotes<sup>8</sup>. Genes encoding GAD are found, for example, in Escherichia coli<sup>9</sup>, Saccharomyces cerevisiae<sup>10</sup>, Drosophila melanogaster<sup>11</sup>, and Caenorhabditis elegans<sup>12</sup>. Furthermore, GABA signaling via membrane receptors elicits hyperpolarization in plants as well as mammals, suggesting conserved or convergent roles for the product of GAD enzymatic activity<sup>13</sup>. In most vertebrate species, only two GAD genes have been described. Another gene in the GAD family, GAD-like 1 (*GADL1*) resembles *GAD1* and *GAD2* in sequence, but is expressed in mouse skeletal muscles and kidney rather than in the brain<sup>14</sup>. There have also been some hints of greater diversity in vertebrate GAD genes.

In addition to the teleost gad1 and gad2 genes, a third gene, gad3, was found in brain cDNA of the abyssal grenadier (Coryphaenoides (Nematonurus) armatus), a benthic teleost fish<sup>15</sup>. A similar gad3 sequence was subsequently identified in the brain cDNA of goldfish (*Carassius auratus*)<sup>16</sup>. The sequences of goldfish and abyssal grenadier gad3 are clearly related to gad1a, gad1b, and gad2 sequences, but their evolutionary history remained unknown<sup>16</sup>. Furthermore, no gad3 genes were reported in any species other than grenadier and goldfish. This absence remained an anomaly, since the goldfish (order Cypriniformes), is very distantly related to the abyssal grenadier (order Gadiformes). Recent teleost phylogenies indicate that the Ostariophysians, of which Cypriniformes including goldfish are members, diverged from the Euteleosts, which include the abyssal grenadier, over 250 million years ago<sup>17</sup>. Thus, the conservation of a gad3 gene in these two divergent species suggested that gad3 was present in an early teleost ancestor. Because the teleost lineage is known to have experienced a whole-genome duplication early in its evolution, one reasonable possibility could therefore have been that gad3 was a teleost-specific gad paralog<sup>18-21</sup>.

Since the original identification of gad3 from teleost brain cDNA, many comparative genomic resources have become available. The sequencing of teleost and other vertebrate genomes has been accompanied by the development of databases and software for analyzing the conservation of genes. Sarcopterygii species with sequenced genomes include primitive fishes, e.g. elephant shark<sup>22</sup> and coelacanth<sup>23</sup>, as well as tetrapods, e.g. chicken<sup>24</sup>, dog<sup>25</sup>, human<sup>26</sup>, Tasmanian devil<sup>27</sup>, Chinese softshelled turtle<sup>28</sup>, and Xenopus<sup>29</sup>. Actinopterygii species with sequenced genomes include the spotted gar<sup>30</sup> as well as teleosts like fugu<sup>19</sup>, medaka<sup>31</sup>, tilapia<sup>32</sup>, and zebrafish<sup>33</sup>.

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Species	Scientific Name	GAD1	GAD2	GAD3
Chicken	Gallus gallus	ENSGALT00000043162	ENSGALT00000012268	
Burton's mouthbrooder	Astatotilapia burtoni	Gad1a: XM_014332345 Gad1b: XM_014340384	XM_005932121	XM_005950266
Coelacanth	Latimeria chalumnae	ENSLACT00000014577	ENSLACT00000011268	ENSLACT0000005682
Dog	Canis familiaris	ENSCAFT00000049584	ENSCAFT0000006929	ENSCAFT0000000144
Elephant shark	Callorhinchus millii	SINCAMT0000000719	SINCAMT00000011054	SINCAMT0000005039
Fugu	Takifugu rubripes	Gad1a:ENSTRUT00000045798 Gad1b:ENSTRUT00000020549	ENSTRUT0000024751	ENSTRUT00000021119
Abyssal grenadier	Coryphaenoides armatus	AF043268	AF043267	AF043269
Human	Homo sapiens	ENST00000358196	ENST00000376261	ENST00000592477*
Medaka	Oryzias latipes	Gad1a:ENSORLT00000021605 Gad1b:ENSORLT00000011550	ENSORLT00000016248	
Spotted Gar	Lepisosteus oculatus	ENSLOCT0000009532	ENSLOCT0000009370	ENSLOCT0000015874
Tasmanian Devil	Sarcophilus harrisii	ENSSHAT00000013524	ENSSHAT00000015741	ENSSHAT0000004379
Nile tilapia	Oreochromis niloticus	Gad1a:ENSONIT00000011023 Gad1b:ENSONIT00000023558	ENSONIT0000008095	ENSONIT0000008040
Chinese softshelled turtle	Pelodiscus sinensis	ENSPSIT00000002371	ENSPSIT00000019780	ENSPSIT00000019627
Xenopus	Xenopus tropicalis	ENSXETT00000040862	ENSXETT00000040531	ENSXETT00000012900
Zebrafish	Danio rerio	Gad1a:ENSDART00000140425 Gad1b:ENSDART00000003008	ENSDART00000021609	ENSDART00000109561

Table 1. Vertebrate GAD1, GAD2, and GAD3 transcript sequence IDs. \*pseudogene transcript sequence.

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We used recently generated genomic resources to ask whether *gad3* is present and expressed in species other than the goldfish and abyssal grenadier. Our results revealed a surprisingly broad conservation of *GAD3* in mammals, reptiles, birds, and amphibians, as well as fishes.

#### Methods

Throughout this paper, we use standard gene nomenclature. For fishes, gene symbols are lowercase and italicized and protein symbols are capitalized. For other vertebrates, human conventions are used: gene symbols in all capitals and italicized, protein symbols in all capitals.

Vertebrate sequence data (Table 1) for *GAD1*, *GAD2*, and *GAD3* homolog transcripts were downloaded from Ensembl genomes for the following species: chicken (*Gallus gallus*), coelacanth, fugu, human, medaka, spotted gar, Tasmanian devil, tilapia, Chinese softshell turtle, xenopus, zebrafish<sup>34</sup>. Transcript DNA sequences for elephant shark were retrieved from the elephant shark Ensembl server. Transcript cDNA sequences for grenadier were retrieved from NCBI<sup>15,35</sup>. The spotted gar genome shares extensive similarity with both tetrapod and teleost genomes, so we chose to focus on this species for sequence alignment, phylogenetics, and intron/exon structure comparisons<sup>30</sup>. Additionally, we obtained sequences for fruitfly (*Drosophila melanogaster*) *GAD1*: NM\_079190, sea urchin (*Strongylocentrotus purpuratus*) *GAD*: XM\_779763, amphioxus (*Branchiostoma floridae*) *GAD*: XP\_002592141, and tunicate (*Ciona intestinalis*) *GAD*: ENSCINT00000004013.

In addition to the species listed in Table 1, we identified several other tetrapod species with *GAD3* genes. These included: Orangutan: ENSPPYG00000009199, Rhesus: ENSMMUG00000001554, Rabbit: ENSOCUG00000022124, Horse: ENSECAG00000009017, Platypus: ENSOANG00000002106, Lizard: ENSACAG00000008555.

**Sequence Alignment.** Both DNA and amino acid sequences were aligned using MAFFT v7.017<sup>36,37</sup> (by translation alignment for CDS sequences); algorithm E-INS-I; scoring matrix: BLOSUM62; gap open penalty: 1.53; offset value: 0.

**Model Testing.** MEGA 6 software was used to compare 24 DNA evolution models for the aligned GAD CDS sequences<sup>38</sup>. A generalized time-reversible plus gamma (GTR + G + I) model had the lowest BIC score (Bayesian Information Criterion) and AICc value (Akaike Information Criterion, corrected), so it was used for subsequent phylogenetic analyses. In this model, non-uniformity of evolutionary rates among sites is modeled by estimating a discrete Gamma distribution (+G) of rates and by assuming that certain sites are evolutionarily invariable (+I).

**Bayesian Phylogenetic Inference.** MrBayes  $3.2.6^{39}$  was used to infer phylogenetic relationships between GAD homologs based on aligned nucleotide CDS sequences, and was accessed via the CIPRES web portal<sup>40</sup>. In MrBayes, the GTR + G + I model of evolution was used; with default settings except for the following specified parameters: nruns(number of runs) = 2; ngen(number of generations) = 1000000; samplefreq = 500; nchain (number of chains) = 8; temp(chain heating temperature) = 0.1; savebrlens = yes; burninfrac(fraction of initial generations discarded) = 0.25.

Diagnostics of the MCMC sampling were carried out using Tracer v1.6 (http://tree.bio.ed.ac.uk/software/ tracer/). The effective sample size (ESS) for each parameter was >300 for each run, allowing adequate sampling of the Markov chain.

The tree file generated using MrBayes was visualized using FigTree v1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/).

**Synteny.** Gene synteny for *GAD3* genes was compared to the syntenic region near *GAD3* using Genomicus<sup>41</sup>. Orangutan was used as a reference species for cross-species synteny and protein similarity to highlight

Spotted Spotted

	1	10	20		30	40	50	60	70	
Gar Gad1	MASSAPSS	SSGGAPI	dp <b>n</b> stnir <b>i</b>	PST <b>T</b> YDT	WCGVAHG	CTRKLGMK	ICGFLQRN	NSLEDKSRIV	SSLKERQSS	K N L F S CE
Gar Gad2	MASHGFWS	F <b>G</b> TENC	GG <b>N</b> NSSQS <b>I</b>	Y S <b>T</b> PRA	WCQAAQK	FTGGLGSK	LCALL	- SVGEGEKTI	D <b>S</b> TTKQ <b>Q</b> GA	-TLETCE
Gar Gad3	M								EKVKS	DKHL <b>T</b> K <b>E</b>
	80	90	10	0	110	120	130	14	) 1	50
Gar Gad1	NTDKDSRH	r <b>r</b> a é t	D		FSNLFAR	DLLPAKNG	EEPTMQFL	LEVVDILLNY	VRKTFERST	KVLDFHH
Gar Gad2	- CSKPCNC	CSKTNV	D		FSVLYST	DLLPATDG	DLATITFL	QEIVDILLAY	IVKTFDRST	KVIDFHY
Gar Gad3	NNKAESIW	I G E K R Q	DAICIENDO	CVVNGTKN	FSTIYST	DLLPTKNG	EEPTKHFL	LEVVNILLNY	VKKSFDRSS	KVLDFHY
	160	17	70	180	190	200	)	210	220	230
Gar Gad1	PHQLLEGN	IEGFNL	ELSEQPESI	EQILVDC	RDTLKYG	VRTGHPRF	FNQLSSGL	DIIGLAGEWI	TSTANTNMF	TYEIAPV
Gar Gad2	PNEL I	QTNNWI	ELSDEPETI	DDILLNC	RATLKYA	IKTGHPRY	FNQLSTGL	DMVGLAADWI	TSTANTNMF	TYEIAPV
Gar Gad3	PHQLKEGI	<b>D</b> GFSL	DLPDQPENI	EQVLVDC	RDTLKYG	VKTGHPRF	FNQLSSGL	DIIGLAGEWI	TSAANTNIF	TFEISPV
	240		250	260	270	)	280	290	300	310
Gar Gad1	FVLMEQLI	LKKMRI	EMIGWPGGI	GDGIFSP	GGAISNM	YSVMAARY	KYFPEVKT	K G M 🗛 A V P K L 🗉	LFTSEHSHY	SIKKAGA
Gar Gad2	FVLLEYVT	LKKMRI	EIIGWPEGN	GDGIFSP	GGAISNM	YAMLVARF	KMFPEVKE	KGMSAVPRLV	AFTSEHSHF	SIKKGAA
Gar Gad3	FILMEEVI	LKKMQI	EKIGWPAEI	RDGIFSP	GGSISNL	YSVLLARY	HFFFFKT	KGMAAVPRLA	L F T S E H S H Y	SIRKAAA
	32	20	330	340		350	360	370	380	390
Gar Gad1	ALGEGTEN	IVILLK(	DERGRVIE	ADLEAKI	IDAKQKG	HVPLFVNA	TAGTTVYG	AFDPIHDIAC	ICEKYNLWL	HVDGAWG
Gar Gad2	ALGIGTDS	SVILIKV	/ DERGKMIE	SDLERRI	VEAKQKG	FVPFFVSA	TAGTTVYG	AFDPLIAIAC	ICRKHKVWM	HVDGAWG
Gar Gad3	VLGIGTEN	<b>VFMVK</b>	DERGKMII	SELEFSI	LKAKVMG	SVPFYVNA	TAGTTVYG.	AFDPL <mark>S EIA</mark> D	VCEKHNLWM	HVDASWG
		400	410	4	120	430	440	450	460	
Gar Gad1	GCLLMSRK	KHRHKLS	GIERANSV	TWNPHKM	MGVLLQC	SAILVREK	GILQGCNQ	MCAGYLFQQD	KQYDVTYDT	GDKAIQC
Gar Gad2	GSLLMSRK	KHRWKL	GVERANSV	′ ТѠNPНКМ	MSVPLQC	SALLVREE	<u>G LM Q</u> R CN Q.	MHACYLFQQC	KHYDLSYDT	GDKALQC
Gar Gad3	GCLLMSKF	KHSVKLI	KGIERAISV	ТШИРНКМ	MGIPLQC	SAILVRKK	GLLQSCNQ	LCAEYLFQPD	KHYDVSYDT	GDKIIQC
	470	480	49	0	500	510	520	53	) 5	40
Gar Gad1	<u>GRHVD</u> FF	FWLMWI	K A K G T <mark>V</mark> G F E	QQINKCL	ELSEYLY	TKIKNREG	YEMVF DGE	PQHINVCFWY	IPPSLRVMP	DCEERRE
Gar Gad2	GRHVDIFF	LWLMWI	RAKGTVGFE	AQIDKCL	ELSEYLY	NKIKNRDG	YEMVFDGK	PQHINVCFWY	LPPGLRYVE	DKEERMR
Gar Gad3	GRHVDAFF	FWLMWI	KAKGTEGFE	AQINKCL	QNAQHFY	NELKKRDD	FELVYKSE	PEHSNVCFWY	IPPSLKQSP	H G V <b>BR</b> N M
	550	50	50	570	580	590	)	600	610 6	518
Gar Gad1	R L HKVA PK	IKARMN	MESGTTMV	YQPQGDK	VNFFRMV	ISNPAATK	SDIDFLID	EIERL	GQD	L
Gar Gad2	RLHKVAP	IKARMM	MEYGTTMVS	YQPQGEK	VNFFRMV	ISNPAATF	DIDFLIE		GQD	L
Gar Gad3	КЬНВVАРК	IKAKMI	IEEGTTMVC	YQPLGEN	VNFFRCV	FSNPATNK	SDVDFLIE	PFTSK	GTVSFN <mark>GQ</mark> -	F.

**Figure 1.** Alignment of predicted protein sequences translated from spotted gar (*Lepisosteus oculatus*) **GAD genes.** Black: amino acids similar in all three sequences; Gray: similar to one corresponding residue; White: not similar to either corresponding residue. Sequence similarity was calculated using BLOSUM62 matrix with threshold = 1.

conservation of synteny and GAD3 protein sequence in vertebrates despite the absence of *GAD3* in some hominids. For comparing primate synteny, simiiformes (last common ancestor of simians) was used as the reference taxon. Synteny data, protein similarity, and species images were downloaded from Genomicus.

### Results

We found previously uncharacterized *GAD3* genes in many vertebrate genomes, including diverse fishes and tetrapods. A teleost *gad3* transcript was found in a transcriptome library generated from testis tissue from *Astatotilapia burtoni*: (>comp56037\_c0\_seq1\_indA\_testis). Zebrafish *gad3* has been previously referred to with the identifier *zgc:163121*. Interestingly, *GAD3* had already been annotated in the *Xenopus tropicalis* genome as *GAD1.2*. It appears, however, that it has not yet been studied in *Xenopus*.

**Sequence Similarity.** Gad3 predicted protein sequence from spotted gar (*Lepisosteus oculatus*) is more similar to Gad1 (Pairwise Identity: 60.5%) than to Gad2 (Pairwise Identity: 53.9%) (Fig. 1). Gad1 and Gad2 share 67.1% pairwise identity. The N-terminal domain, which is quite variable between Gad1 and Gad2, is truncated and highly divergent in Gad3. The N-terminal 92 amino acids (aa) of Gad1 align to the N-terminal 84 aa of Gad2. Gad3 has 42 aa aligning in this range, only 27 of which align to Gad1 and Gad2 sequence (with a 15 aa gap). These 27 aa of Gad3 have: 23.1% pairwise identity with Gad1, 11.5% pairwise identity with Gad2.

**Phylogeny.** Pyridoxal 5'-phosphate (PLP)-dependent decarboxylase genes include Glutamate Decarboxylase-Like 1 (*GADL1*), Cysteine Sulfinic Acid Decarboxylase (*CSAD*), and histidine decarboxylase (*HDC*), in addition to GADs. Therefore, we tested the phylogenetic relationship of *GAD3* to other genes in this group, using *HDC* as an outgroup for *GAD*, *GADL1*, and *CSAD* genes<sup>42</sup>. A neighbor-joining tree of aligned predicted amino acid sequences from the spotted gar (*Lepisoteus oculatus*) place *gad3* most closely related to the *gad1/gad2* clade (Fig. 2).

A phylogenetic tree of vertebrate *GAD1*, *GAD2*, and *GAD3* nucleotide coding sequences was generated using MrBayes (Fig. 3). In insects, *GAD1* is the single homolog of vertebrate GAD genes (insect *GAD2* is homologous to vertebrate *CSAD* and *GADL1*). Therefore we chose *Drosophila melanogaster GAD1* as the outgroup for the vertebrate and deuterostome GAD genes.

**Exon-intron Structure.** Spotted gar *gad1* and *gad2* each have 16 exons (Fig. 4). Spotted gar *gad3* has 17 exons. While both *gad1* and *gad2* have coding sequence beginning in exon 1, *gad3* coding sequence (CDS) begins in the second exon (exon 2). The predicted coding sequence of *gad3* has a gap (does not align) with the 5' CDS sequence found in *gad1* and *gad2* exon 1, exon 2, and part of exon 3. The 3' portion of the *gad3* CDS is included on exon 17, while *gad1* and *gad2* stop codons are found in exon 16.

Aside from these differences, *gad3* exon structure is largely similar to *gad1* and *gad2*. All of the exon junctions from exon 3 to exon 16 are in identical locations for all three gad genes. In our alignment of the three *gad* genes, the only gaps introduced in *gad3* are located in exon 2 and exon 17.

**Synteny.** In spotted gar, *gad1* and *gad2* are located adjacent to the myosin genes *myo3b* and *myo3a*, respectively. Similarly, in humans *GAD1* is located near *MYO3B* on chromosome 2, and *GAD2* is located adjacent to



**Figure 2.** GAD3 is closely related to GAD1 and GAD2, and more distantly related to other members of the PLP-dependent decarboxylase gene family. A phylogenetic tree of aligned PLP-dependent decarboxylase amino acid sequences from the spotted gar was generated using neighbor-joining and 2000 bootstrap iterations. Percent bootstrap support for nodes are shown. The scale bar (bottom) indicates substitutions per site.



**Figure 3.** Phylogenetic tree of *GAD1*, *GAD2*, and *GAD3* nucleotide sequences. This consensus tree was generated using MrBayes with *Drosophila melanogaster GAD1* as the outgroup. Nodes are labeled with posterior probabilities. Distinct gene lineages are indicated by colors. The scale bar (bottom) indicates substitutions per site.

MYO3A on chromosome 10. On the other hand, gad3 is not located near a myosin gene in the genome of spotted gar. The genes located adjacent to spotted gar gad3 are mc4r and cdh20. This syntenic block of genes is conserved



**Figure 4.** Shared exon-intron structure of GAD genes from spotted gar (*Lepisosteus oculatus*). (A) Maps of the three gad genes showing 16 exons in *gad1* and *gad2*, and 17 exons in *gad3*. The direction of transcription is from left to right. Genomic distance spanned and strand on which the gene is located are indicated for each gene. (B) Aligned spotted gar *gad1*, *gad2*, and *gad3* CDS regions annotated with position of exons. Bases colored black indicate disagreements with the consensus sequence of the three gad genes.

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across many vertebrate species, and represents the inferred ancestral state of the bony vertebrates (euteleostomi) (Fig. 5).

**GAD3 Conservation and Gene Loss.** GAD3 predicted protein sequence is highly conserved in diverse vertebrate genomes (Fig. 6). Yet primates appear to have experienced varying degrees of gene loss at the *GAD3* locus (Fig. 7). We identified a human transcript (ENST00000592477) with homology to *GAD3* (Table 1), derived from a pseudogene located in the human genome in the conserved *GAD3* syntenic position between *MC4R* and *CDH20* (Fig. 7). Similarly, gorilla (*Gorilla gorilla*) *GAD3* is annotated as a pseudogene in Ensembl (ENSGGOG0000027455). Although macaque (*Macaca mulatta*) and orangutan (*Pongo pygmaeus*) predicted GAD3 protein sequences share relatively high pairwise identity (84.5%), the *GAD3* genes in these two species appear to have large insertions (or deletions) in their predicted coding sequence.

#### Discussion

We identified a novel glutamate decarboxylase homolog, *GAD3*, found in many vertebrate genomes. We provide phylogenetic and intron/exon structural evidence that *GAD3* is an ancient paralog of *GAD1* and *GAD2*. The conserved chromosomal synteny of *GAD3* in vertebrates supports an ancient origin for this gene. Surprisingly, *GAD3* was lost in the hominid lineage. Taken together, the phylogenetic analyses, comparisons of gene structure, and synteny data suggest that *GAD3* arose via gene duplication of a protovertebrate *GAD* homolog, likely before the duplication of another paralog which gave rise to *GAD1* and *GAD2*.

**GAD3 Evolution.** Although our data do not rule out the possibility of a local duplication that gave rise to *GAD3*, they are consistent with an origin of *GAD3* in an early vertebrate via whole-genome duplication. Whole-genome duplication is thought to have played a major role in early vertebrate evolution<sup>43-45</sup>. Following genome duplication, these duplicated gene pairs (ohnologs) experienced a range of outcomes including non-functionalization, sub-functionalization, and neo-functionalization<sup>46,47</sup>. Sub-functionalization may happen via protein changes<sup>48</sup> or via regulatory element loss<sup>49</sup> in which ancestral expression domains are differentially lost in different genes<sup>50</sup>. For example, recent evidence indicates that duplication of a corticotropin-releasing hormone



**Figure 5.** *GAD3* genes are found in a conserved syntenic region in vertebrates. Compared to orangutan *GAD3* for reference, other species including representatives of mammals, other tetrapods, and Euteleostomi (bony vertebrates) in general have similar chromosomal positions of *GAD3* genes. The central green pentagons (surrounded by a vertical rectangle) represent the *GAD3* genes. For each species, *GAD3* and 10 flanking genes on each side are represented by colored pentagons. The pentagons point in the direction of transcription, and each color identifies a set of orthologous genes. To the right of the figure, both species and chromosome are indicated for each *GAD3* ortholog. Figure modified from Genomicus PhyloView output<sup>41</sup>.

(CRH) gene in an early vertebrate led to a broadly expressed CRH1 and a CRH2 with expression restricted to a single hindbrain nucleus<sup>51</sup>. Like our recent analyses of CRH genes, the discovery of GAD3 as a conserved vertebrate gene relied on freely available genomic resources, pointing to the likelihood that many gene families have unannotated homologs remaining to be found in sequenced genomes<sup>51,52</sup>.

**GAD3 Function.** GAD3 is phylogenetically closer to GAD1 and GAD2 than to GADL1, but nonetheless it is possible that its enzymatic functions differ from those of GAD1 and GAD2. The absence of much of the N-terminal region, which regulates intracellular localization of GAD1 and GAD2 proteins, suggests that GAD3 protein may have different localization<sup>5</sup>.

Little is known regarding the function of GADL1 enzyme, though polymorphisms are linked to differential response to lithium treatment for bipolar disorder<sup>53</sup>. Mammalian GADL1 does not appear to have glutamate decarboxylase activity, despite its name. Instead, it catalyzes the decarboxylation of aspartate, cysteine sulfinic acid, and cysteic acid to  $\beta$ -alanine, hypotaurine, and taurine, respectively<sup>14</sup>. Recently, GADL1 and CSAD were found to have preference for cysteine sulfinic acid as a substrate<sup>42</sup>. Future studies of GAD3 biochemical substrates will be necessary to address the possibility of substrates other than glutamate.

Intriguingly, zebrafish *gad3* (referred to as *zgc163121*) mRNA expression was significantly downregulated by treatment with dexamethasone, a glucocorticoid agonist, in 25hpf larval zebrafish, as measured by microarray and qPCR<sup>54</sup>. In the deep-sea fish in which *gad3* was first described, the armed grenadier, *Coryphaenoides* (*Nematonurus*) *armatus*, *gad2* mRNA levels were found to be expressed in the brain in a sexually dimorphic manner, i.e. higher in male hypothalamus than in female, but no differences were found in *gad3* levels<sup>35</sup>. In the goldfish, however, *gad3* mRNA levels in the telencephalon were highest in sexually mature fish of both sexes during the breeding period<sup>55</sup>. Since the specific role of *gad3* is unknown in any taxa, the full range of factors that regulate *gad3* expression in the brain, and potentially elsewhere, awaits further investigation.

**Loss in hominids.** The loss of *GAD3* in both chimpanzees and humans appears to have been preceded by changes to *GAD3* sequences in other hominids. Predicted gorilla, orangutan, and gibbon *GAD3* transcripts appear to be truncated relative to fish *gad3* sequence, but it may be that not all the exons in these sequences are fully annotated in Ensembl. Glutamate metabolic pathways appear to have been under positive selection in hominids, as seen for example in the origin of glutamate dehydrogenase 2 (*GLUD2*) by retroposition of *GLUD1*<sup>56</sup>.

Gene losses have played major roles in human evolution<sup>57</sup>. For example, loss of L-gulonolactone oxidase (*GULO*) makes humans and other Haplorhini susceptible to scurvy, a vitamin C deficiency. Despite conferring this disadvantage, GULO gene loss has occurred in multiple mammalian lineages, including guinea pigs<sup>58</sup> and some bats<sup>59</sup>.





**Figure 6.** GAD3 protein sequence is highly conserved across Euteleostomi. Predicted protein sequences of GAD3 orthologs were compared using Genomicus<sup>41</sup>. The degree of similarity to the reference sequence (orangutan, in green) is indicated by the color of the block, according to the scale shown at bottom.



**Figure 7.** *GAD3* is lost in hominids. The central green pentagons (surrounded by a vertical rectangle) represent the *GAD3* genes. Human and chimpanzee genomes have no functional gene at the location corresponding to *GAD3* in other primates. Humans have a pseudogene at this location. A thick blue line between two genes indicates a "gap", i.e. a gene lost relative to the ancestral Simiiformes genome. A thin blue line between two genes indicates a "break" in the continuity of the alignment, i.e. a gene added or lost relative to the ancestral Simiiformes genome. Figure modified from Genomicus AlignView<sup>41</sup>, which relies on data and images from Ensembl<sup>34</sup>.

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Hominids are not the only lineage that has lost *GAD3*. Rodent genomes, including mice, rats, and squirrels also appear to be missing *GAD3* homologs. The absence of *GAD3* in both humans and mice likely explains why this gene was not discovered sooner, since many investigators choose to focus on these two species.

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### **Author Contributions**

Both authors had full access to all of the data in this study and take responsibility for its collection and analysis. Study concept and design: B.P.G. and K.P.M. Acquisition of data: B.P.G. Analysis and interpretation of data: B.P.G. and K.P.M. Drafting of the manuscript: B.P.G. Critical revision of the manuscript for important intellectual content: K.P.M. Obtained funding; K.P.M. Administrative, technical, and material support: K.P.M.

### Additional Information

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