Mutations in GPAA1, encoding a GPI transamidase complex protein, cause developmental delay, epilepsy, cerebellar atrophy and osteopenia.

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<u>Abstract</u>

Approximately one in every 200 mammalian proteins is anchored to the cell membrane through a glycosylphosphatidylinositol (GPI) anchor. These proteins play important roles notably in neurological development and function. To date, more than 20 genes have been implicated the biogenesis of GPI-anchored proteins. GPAA1 (glycosylphosphatidylinositol anchor attachment 1 protein) is an essential component of the transamidase complex along with PIGK, PIGS, PIGT and PIGU (Phosphatidylinositolglycan biosynthesis class K, S, T and U proteins). This complex orchestrates the attachment of the GPI anchor to the carboxyl terminus of precursor proteins in the endoplasmic reticulum. Here we report bi-allelic mutations in GPAA1 in ten individuals from five families. Using whole exome sequencing, we identified two frameshift mutations (c.981_993del [p.Gln327Hisfs*102] and c.920delG [p.Gly307Alafs*11]), one intronic splicing mutation (c.1164+5C>T) and six missense mutations (c.152C>T [p.Ser51Leu], c.160_161delinsAA [p.Ala54Asn], c.527G>C [p.Trp176Ser], c.869T>C [p.Leu290Pro], c.872T>C [p.Leu291Pro] and c.1165G>C [p.Ala389Pro]). Most individuals presented with global developmental delay, hypotonia, early-onset seizures, cerebellar atrophy, and osteopenia. The splicing mutation was found to decrease GPAA1 mRNA. Moreover, flow cytometry analysis on five available individual samples showed that several GPI-anchored proteins had decreased cell surface abundance in leukocytes (FLAER, CD16, CD59) or fibroblasts (CD73, CD109). Transduction of fibroblasts with a lentivirus encoding the wildtype protein partially rescued the GPI-anchored protein deficiency. These findings highlight the role of the transamidase complex in the development and function of the cerebellum and the skeletal system.

Main text

Initially identified more than three decades ago¹, glycophosphatidylinositol (GPI) anchor proteins (AP) are emerging as a structurally diverse group of membrane-bound glycoproteins with important roles in cellular processes such as embryogenesis, fertilization and neurogenesis. It is estimated that there are about 150 GPI-APs in the human proteome.

The attachment of a GPI anchor to a protein is a conserved posttranslational modification which occurs in the endoplasmic reticulum (ER) and requires at least twelve reactions. Proteins bearing the GPI-anchor exit the ER and are shuttled to the Golgi where they undergo fatty acid modification before being transported to the plasma membrane. Crucial in the GPI biosynthetic pathway are a group of enzymes encoded by five genes (PIGK [MIM:605087], PIGS [MIM: 610271], PIGT [MIM: 610272], GPAA1 [MIM: 603048], and PIGU [MIM: 608528]) forming the GPI transamidase complex. These proteins mediate coupling of the synthesized GPI-anchor to mature proteins with a carboxyl terminal GPIattachment signal peptide² in two principal steps; the cleavage of a C-terminal propeptide from the substrate protein and the formation of an amide bond between the C-terminal residue (ω-site) of the substrate protein and a phosphoethanolamine group of the GPI lipid anchor². Among the five known subunits of the GPI lipid anchor transamidase complex, GPAA1 was the first to be discovered^{3; 4} but its physiological role was only demonstrated recently⁵. This protein is composed of an N-terminal transmembrane domain, a luminal region and a hydrophobic region containing six transmembrane domains in the Cterminus. Recently, Eisenhaber et al. demonstrated that the luminal region of GPAA1 is similar to M28-type peptidases and that it may have one Zn-binding site like some of the M28 family peptidases. This protein was thus suggested to be a catalyst in the second

step to complete the transamidation⁵. Several inherited disorders are known to be caused by GPI biosynthesis defects⁶. However, mutations in GPAA1 have yet to be reported in diseases other than cancer⁷⁻¹⁰. Here, we present functional characterization of GPAA1 mutations in ten individuals with GPAA1 deficiency who manifest clinical features consistent with other inherited GPI-anchor deficiencies, including developmental delay, hypotonia and seizures.

Ten individuals from five unrelated families with bi-allelic GPAA1 (Gene bank: NM_003801.3) mutations were included in this study. Individuals were identified by a search in the Baylor Genetics Laboratory proband whole exome sequencing database, the DECIPHER database¹¹, Genematcher¹², and contacting collaborators. Informed consent was obtained from all families according to protocols approved by the Institutional Review Board at the relevant institutions. Most individuals had global developmental delay including late or absent independent walking and speech abilities. Among the more severely affected individuals, six individuals from families 1, 2, 3 and 5 are unable to walk independently at age 6 and up, and the individual from family 2 has not acquired speech as of age 6. Hypotonia was identified in all cases at an early age. Seizures were present in seven individuals; cerebellar atrophy was found in seven individuals (see MRI in figure 1B) and osteopenia also in five individuals. Further clinical features are summarized in table 1, table S1, and additional MRI images of family 4 can be found in figures S1 to S4.

Exome sequencing showed compound heterozygous mutations in the two siblings of family 1 (Hispanic origin from the USA), a missense mutation c.872T>C [p.Leu291Pro] inherited from the father and a frameshift mutation c.981_993del [p.Gln327Hisfs*102] which is from the mother (Figure 1). Individual 2 (II-2 of family 2 in figure 1, a white family, from the USA) has a missense mutation c.152C>T [p.Ser51Leu] from the mother and an

intronic splicing mutation c.1164+5C>T from the father. He has an unaffected sibling who carries only the c.152C>T [p.Ser51Leu] mutation but no splicing mutation. Two affected siblings from an Egyptian-origin family (family 3) have a frameshift mutation c.920delG [p.Gly307Alafs*11] and missense mutation c.1165G>C [p.Ala389Pro]. A homozygous missense mutation c.527G>C [p.Trp176Ser] was found in two Pakistani siblings from family 4 and a distant cousin, all of whose parents are consanguineous. Two affected siblings from family 5 (a Finnish family) had compound heterozygous mutations, a missense mutation c.869T>C [p.Leu290Pro] inherited from the father and a missense mutation c.160_161delinsAA [p.Ala54Asn] inherited from the mother. DNA from two unaffected siblings was not available for sequencing. These mutations were either not seen in ExAC or noted at very low frequency without homozygotes (Table 2). See figure 1 for the pedigrees and figure 2 for the location of the mutations and the conservation of the substituted amino acids.

Frameshift or splicing mutations that introduce premature stop codons before the last 50 nucleotides of the penultimate exon usually trigger nonsense-mediated mRNA decay (NMD). To determine if the mutations identified in our cohort triggered NMD, we performed real-time PCR on both fibroblasts and lymphoblastoid cell lines (LCLs) of individuals 1a and 1b (II-1 and II-2 of family 1 in figure 1) with the frameshift mutation (c.981_993del) which introduces a stop codon at amino acid 428 (p.Gln327Hisfs*102). We observed a decrease in GPAA1 mRNA up to 50% in both fibroblasts and LCLs (Figure 3A, B). Similarly, individual 2 (II-2 of family 2 in figure 1) with the intronic splicing mutation c.1164+5C>T also has decreased mRNA expression of GPAA1 in LCLs (Figure 3B).

To assess whether these mutations could affect the level of GPI-AP found at the cell surface, we performed flow cytometry analysis on whole blood and LCLs of affected

individuals from families 1, 2 and 4 (samples from families 3 and 5 were not available). Our results show that all tested individuals have GPI-AP deficiency. As shown in Figure 3C, individuals 1a and 1b (II-1 and II-2 of family 1 in figure 1) and 2 (II-2 of family 2 in figure 1) from two families presented with lower level of total GPI-AP in granulocytes as measured by a decrease in fluorescence-labeled aerolysin (FLAER) staining, while lower levels of CD16 were observed only in individuals 2 (II-2 of family 2 in figure 1), 4a and 4b (VII-1 and VII-2 of family 4 in figure 1). In all LCLs examined, cell surface staining also showed decreased FLAER abundance (Figure 3D). In addition, a moderate decrease in CD24 was observed in all individual cell lines (Figure S5).

Since fresh blood samples were not available for the affected individuals in family 3, we also tested the effect of the mutations by transfecting plasmids encoding mutant cDNAs driven by a weak promoter with only TATA box in cells knocked-out for GPAA1. GPAA1-deficient HAP1 cells were generated by CRISPR-Cas9 targeting of the gene. We then cloned human GPAA1 isoform 1 from a Hep3B cDNA library, and ligated to generate C-terminal tagged wild-type and variant pME hGPAA1 HA, then generated by site-directed mutagenesis the different variants. The pME promoter is the SRα promoter (SV40 early promoter + HTLV LTR, enhancer)¹³. We also subcloned the constructs in plasmids with an intermediate strength promoter (pTK using the herpes simplex virus thymidine kinase, or HSV-TK, promoter), and a weak promoter (pTA using the TATA box from the HSV-TK promoter). The pTA promoter activity is similar to that of a promoter-less plasmid. The promoter activities are estimated as pTA:pTK:pME = 1:60:6000.

These results show a decreased activity compared to the wild-type protein only for the p.Ser51Leu and the p.Ala389Pro variants (Figure 4A). In this overexpression experiment, the other variants (p.Trp176Ser and p.Leu291Pro) possibly have sufficient residual

activities to provide a rescue. Similar results were obtained after transfection with plasmids encoding the cDNAs under the control of stronger promoters (see Figures S6 and S7). Western blotting of cell lysates after transfection of cells with a strong promoter (pME) indicates that the p.Ala389Pro variant, located in the second transmembrane domain, leads to protein instability (Figure 4B). This could possibly be due to inadequate incorporation of the protein into the ER membrane.

We also assessed GPI-AP abundance on the cell surface of skin fibroblasts derived from individuals 1a, 1b (II-1 and II-2 of family 1 in figure 1) and 2 (II-2 of family 2 in figure 1) using FLAER, CD73 and CD109 as GPI cell surface markers (Figure S8). Since individuals 1a and individual 2 had a significant downregulation in CD109 and CD73, respectively, we carried out rescue assays of these GPI-APs on fibroblasts of these individuals using a GPAA1-encoding Lv105 lentiviral vector. As shown in Figure 5, CD109 in individual 1a fibroblasts was completely rescued, and a partial rescue of CD73 was observed in individual 2 fibroblasts. Collectively, these findings demonstrate that individuals with GPAA1 mutations have GPI-AP deficiency, which likely causes their clinical manifestations. The protein with the p.Trp176Ser variant might cause a milder biochemical effect since when overexpressed, no decreased activity was noted.

There is clinical overlap between these individuals and those with other GPI biosynthesis defects. Inherited mutations in PIGA [MIM: 311770]¹⁴⁻¹⁶, PIGM [MIM: 610273]¹⁷, PIGL [MIM: 605947]¹⁸, PIGW [MIM: 610275]¹⁹, PIGV [MIM: 610274]^{20; 21}, PIGO [MIM: 610274]^{22; 23}, PIGN [MIM: 606097]²⁴, PIGC [MIM: 601730]²⁵, PIGP [MIM: 605938]²⁶ and PIGG [MIM: 616918]²⁷ were identified as causing GPI-biosynthesis disorders. Similar disorders can also be caused by mutations in PGAP genes (Post GPI Attachment to Proteins, notably

PGAP1 [MIM: 611655]²⁸, PGAP2 [MIM: 615187]^{29; 30} and PGAP3 [MIM: 611801]³¹), which code for a family of proteins involved in structural remodeling of the GPI anchor after its attachment to proteins. The phenotypes of the individuals with GPAA1 mutations in the present study overlap with those identified in other GPI-biosynthesis disorders, such as the epilepsy, hypotonia and developmental delay23; 32; 33, as well as the progressive cerebellar atrophy in individuals with PIGT and PIGN mutations^{24; 34}. The rate and severity of the progressive atrophy is comparable to that seen with PIGT and PIGN deficiency, although the finding is is rarer in PIGN deficiency. The only reported GPI-transamidase complex gene associated with GPI deficiency is PIGT, which causes multiple congenital anomalies-hypotonia-seizures syndrome^{4; 32; 33; 35}, characterized by hypotonia, developmental delay, seizures, dysmorphisms, and malformations of the heart, urinary tract and gastrointestinal system. With relevance to the condition we describe here, most of these individuals also have osteopenia.

All the individuals we describe have normal plasma and/or serum alkaline phosphatase (ALP) levels (Table 1), which can be perturbed in other GPI-AP-synthesis-related disorders. Previous data suggested that the elevated serum ALP observed in some GPI-biosynthesis defects could be explained by cleavage of the signal peptide and secretion of soluble ALP because of abnormal GPI structure. Since GPAA1 is a component of the GPI transamidase complex, mutations in this gene leading to decreased activity might lead to a decrease in serum ALP. This was not observed in our cohort, perhaps because the remaining enzymatic activity is sufficient to maintain normal serum ALP. This hypothesis is supported by the fact that no individual with bi-allelic loss-of-function mutations was identified. With regards to the mechanism leading to osteopenia, perhaps

decreased surface expression of ALP on osteoblasts affects its function and secondarily leads to the osteopenia seen in these individuals.

To date, GPI-transamidase subunits have been proposed to function as oncogenes in breast and bladder cancer⁷⁻⁹. In addition, GPAA1 mRNA was also found to be overexpressed, correlating with increased GPAA1 protein accumulation, in samples from individuals with colorectal cancer¹⁰. It is thought that an overexpression of GPI-biosynthesis genes might lead to increased uPAR (a GPI-anchored protein) which in turn can lead to increased invasiveness and growth of tumor cells³⁷.

Limited therapies are available for GPI-AP deficiency. Since alkaline phosphatase is essential to allow pyridoxal-phosphate to pass through the blood-brain barrier and pyridoxine passes through the blood-brain barrier and is converted to pyridoxal-phosphate, which is important for GABA synthesis, pyridoxine supplementation sometimes helps control the seizures in individuals with GPI-biosynthesis defects including some patients with PIGO and PIGV mutations^{38; 39}. This potential treatment has not yet been tested in our cohort. Future studies of GPAA1 deficiency, such as in mouse models, could allow further pathophysiological studies to explore the cerebellar and bone manifestations, and explore therapeutic avenues.

In summary, we report biallelic mutations in GPAA1 in ten individuals presenting global developmental delay, hypotonia, early-onset seizurescerebellar atrophy., and osteopenia. Moreover, with functional studies, we demonstrate that these individuals have low cell surface GPI-AP levels demonstrating that those mutations cause a defect in the biosynthesis of GPI anchored proteins. Thus, our work expands the group of GPI-

biosynthesis disorders and contributes to the understanding of the role of the GPI transamidase complex in health and development.

Supplemental Data

Supplemental Data include supplemental text, one table, eight figures and can be found with this article online.

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Web Resources

ExAC Browser, http://exac.broadinstitute.org/

GenBank, http://www.ncbi.nlm.nih.gov/genbank/

OMIM, http://www.omim.org/

UniProt, http://www.uniprot.org/uniprot/

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Figure titles and legends

Figure 1. A) Pedigrees of five families with GPAA1 mutations in the study. B) MRI of individual 4b (VII-2 of family 4) at 25 months. T1 sagittal image shows marked reduction in cerebellar volume (arrow). C) Photographs of, starting from the top right, individuals 1a (age 15), 1b (age 10), 3a (age 10), 3b (age 3), and 5 (age 25). A slightly broad and prominent root of the nose is noted in individuals 1b, 3b and 5.

Figure 2. A) Location of the protein variants on the GPAA1 protein (for the splicing mutation, the site of the exon-exon junction is shown). B) Amino acid conservation in vertebrates of the amino acids affected by substitution mutations.

Figure 3. GPAA1 expression by Real time-PCR in individuals from families 1 and 2. RNA extractions from fibroblasts (A) and LCLs (B) of individuals 1a, 1b and 2 were subjected to qRT-PCR. Figures show the results normalized to a reference gene (TBP for fibroblast and GAPDH—for LCLs) from quadruplicate experiments. C) GPI-AP abundance in granulocytes of individuals from familly 1, 2 and 4. Fresh blood samples as well as healthy controls were fixed with 10% formaldehyde, red blood cells were lysed in 0.1% Triton X-100 and the samples were stained with GPI-AP markers (FLAER, CD16) for 1 hour at room temperature. Non-specific binding was washed before analyzing by BD FACScanto II system. The figure shows representative results from experiments done in triplicate. D) GPI-AP abundance in LCLs of individuals from familly 1, 2 and 4. LCLs were established by Epstein-Barr virus immortalization of peripheral blood mononuclear cells (PMBC). Cells were stained with a GPI-AP marker (FLAER) for 1 hour at room temperature. Non-specific binding was washed before analysing the cell surface abundance of GPI-APs by BD FACScanto II system. The figure shows representative results from experiments done in triplicate.

Figure 4. A) Rescue of GPAA1 knockout cells. GPAA1-knockout cells were transiently transfected with a plasmid encoding wild-type or variant GPAA1 under the control of a weak promoter (pTA plasmid) by electroporation. Transfection efficiency was monitored by luciferase assay, and flow cytometry analysis was performed 2 days after transfection. These results show a slightly decreased labelling only for the p.Ser51Leu and the p.Ala389Pro variants (fewer cells with a strong signal), possibly because the residual activities for the other variants are sufficient to provide a rescue in this experiment. B) The p.Ala389Pro variant leads to protein instability. HEK293 cells were transiently transfected with various GPAA1 cDNAs and proteins were analyzed by Western Blotting using an anti-HA antibody (Cell Signaling). Intensities of the bands were normalized with the loading control (GAPDH), and luciferase activities used for evaluating transfection efficiencies. Western blot for the tagged protein indicates that p.Ala389Pro, in the second transmembrane domain, leads to protein instability.

Figure 5. Lentivirus rescue assays in fibroblasts from individuals 1a and 2. Skin fibroblasts derived from individuals 1a and 2 were transduced with GPAA1-expressing-Lv105 lentivirus or empty vector-lentivirus, then transduced, and non-transduced cells were stained with CD109 and CD73 for 1 hour at room temperature. The non-specific binding was washed before analysing the cell surface abundance of GPI-APs by BD FACScanto II system. The figure shows representative results from experiments done in triplicate.

1 <u>Tables</u>

Table 1. Key clinical features. More details can be found in the supplement.

Individual	1a	1b	2	3a	3b	4a	4b	4c	5a	5b
Gender	F	М	M	F	F	F	M	F	M	М
Age at last assessment (yr)	15	10	6y	10.7	3.8	8	5	4	30	25
Height, cm	152.4 (8 th %ile)	127 (4 th %ile)	103.2 (59 th %ile) at 4 y	125 (1 st %ile, -2.3 SD)	95 (15 th %ile)	109 (1 st %ile, -3.3 SD)	98 (1 st %ile, -2.3 SD)	93 (1st %ile -2.2 SD)	165 (– 2 SD)	167 (– 1.8 SD)
Weight, kg	54.4 (58 th %ile)	27.8 (16 th %ile)	15.4 (30 th %ile)	35 (49 th %ile)	14 (19 th %ile)	23.6 (27 th %ile)	17 (23 rd %ile)	15 (31 st %ile)	70 (48 th %ile)	62.4 (31 st %ile)
OFC, cm	53 (14 th %ile)	52 (27 th %ile)	52 (87 th %ile)	49.5 (1 st %ile, -2.3 SD)	48 (19 th %ile)	51 (31 st %ile)	48.6 (4 th %ile)	48 (18 th %ile)	57,2 (50 th %ile)	57,2 (50 th %ile)
Developmen tal delay/ Intellectual disability ^a	++	++	++	++	++	++	++	+	++	++
Hypotonia	+	+	+	+	+	+	+	++	+	+
Seizures	GTC, myocloni c, atonic	GTC	GTC	GTC	GTC, myoclon ic	No	No	No	GTC, myoclonic, atonic, absence	GTC, myoclon ic
Cerebellar atrophy	+	+	- at age 1	+	+	+	+	+	+	+
Nystagmus	+	+	- but cortical visual impairme nt	+	+	+	-	++	+	+

Dysarthria	NA	+	NA	+	NA	NA	NA	+	+	+
Dysmetria	+	+	NA	NA	NA	NA	+	+	+	+
Ataxic gait	+	+	NA	NA	NA	+	+	++	+	+
Spasticity	-	-	-	+	NA	+	+	+	-	-
Osteopenia	+	+	+	+ (Z=- 2.2)	+ (Z=- 2.0)	+	+	+	NA	NA
Dysmorphis ms	None	Slightl	Bitempora	Bitempor al	Slightly broad	Bitempora I	Bitempora	Narrow promine	None	Slightly broad
		broad	narrowing	narrowin	root of	narrowing	narrowing	nt		root of
		root of	with	g, widely	the nose	,	,	forehead		the nose
		the	prominent	spaced		prominent	prominent	antevert		
		nose	forehead,	eyes,		forehead,	forehead,	ed nares		
			anterevet	prominen		anterevet	anterevet			
Diagram	455 (NII)	4.40	ed nares	t ears	004 (NII)	ed nares,	ed nares	004 (NII)	C	400 (NII)
Plasma alkaline	155 (NI)	143 (NI)	223 (NI)	355 (NI)	624 (NI)	176 (NI)	183 (NI)	201 (NI)	Serum alkaline	180 (NI)
phosphatase		(141)							phosphata	
(U/L)									se 465	
(-)									(275-875	
									Ù/L)(NI)	

^aLegend: For the intellectual disability, + refers to mild, ++ to moderate, +++ to severe.; NA, not available; NI, normal; GTC,

⁴ generalized tonic-clonic seizure; OFC, occipitofrontal circumference; F, female; M, male; %ile, percentile; SD, standard deviation

from the mean.

Table 2. Genetic description of the GPAA1 variants.

Family	Genomic variant	cDNA variant	Protein variant	Inheritance	ExAC minor allele frequency
	(hg19)	(NM_003801.3)			
1	chr8:g.145139374T>C	c.872T>C	p.Leu291Pro	Compound	Not present
				heterozygous	
	chr8:g.145139483_145	c.981_993del	p.Gln327Hisfs*102	Compound	Not present
	139495del			heterozygous	
2	chr8:g.145138104C>T	c.152C>T	p.Ser51Leu	Compound	Not present
				heterozygous	
	chr8:g.145139783C>T	c.1164+5C>T	Splicing	Compound	1.661e-05 total, 0.0001226 in
				heterozygous	South Asians. No homozygotes
					detected.
3	chr8:g.145139422del	c.920delG	p.Gly307Alafs*11	Compound	Not present
				heterozygous	
	chr8:g.145139946	c.1165G>C	p.Ala389Pro	Compound	8.681e-06 total, 7.592e-05 in
	G>C			heterozygous	South Asians. No homozygotes
					detected.
4	chr8:g.145138854G>C	c.527G>C	p.Trp176Ser	Homozygous	1.658e-05 total, 0.0001211 in
					South Asians. No homozygotes
					detected.

5	chr8:g.145138112_145	c.160_161delinsAA	p.Ala54Asn	Compound	Not present
	138113delinsAA			heterozygous	
	chr8:g.145139371T>C	c.869T>C	p.Leu290Pro	Compound	Not present
				heterozygous	