Clinical course, mutations and its functional characteristics of infantile-onset Pompe disease in Thailand

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Abstract

Background: Pompe disease is a lysosomal storage disorder caused by the deficiency of acid alpha-glucosidase (EC. 3.2.1.20) due to mutations in human GAA gene. The objective of the present study was to examine clinical and molecular characteristics of infantile-onset Pompe disease (IOPD) in Thailand.

Methods: Twelve patients with infantile-onset Pompe disease (IOPD) including 10 Thai and two other Asian ethnicities were enrolled. To examine the molecular characteristics of Pompe patients, GAA gene was analyzed by PCR amplification and direct Sanger-sequencing of 20 exons coding region. The novel mutations were transiently transfected in COS-7 cells for functional verification. The severity of the mutation was rated by study of the GAA enzyme activity detected in transfected cells and culture media, as well as the quantity and quality of the proper sized GAA protein demonstrated by western blot analysis. The GAA three dimensional structures were visualized by PyMol software tool.

Results: All patients had hypertrophic cardiomyopathy, generalized muscle weakness, and undetectable or < 1% of GAA normal activity. Three patients received enzyme replacement therapy with variable outcome depending on the age of the start of enzyme replacement therapy (ERT). Seventeen pathogenic mutations including four novel variants: c.876C > G (p.Tyr292X), c.1226insG (p.Asp409GlyfsX95), c.1538G > A (p.Asp513Gly), c.1895 T > G (p.Leu632Arg), and a previously reported rare allele of unknown significance: c.781G > A (p.Ala261Thr) were identified. The rating system ranked p.Tyr292X, p. Asp513Gly and p. Leu632Arg as class “B” and p. Ala261Thr as class “D” or “E”. These novel mutations were located in the N-terminal beta-sheet domain and the catalytic domain.

Conclusions: The present study provides useful information on the mutations of GAA gene in the underrepresented population of Asia which are more diverse than previously described and showing the hotspots in exons 14 and 5, accounting for 62% of mutant alleles. Almost all mutations identified are in class A/B. These data can benefit rapid molecular diagnosis of IOPD and severity rating of the mutations can serve as a partial substitute for cross reactive immunological material (CRIM) study.

Keywords: Lysosomal storage disorder, Pompe disease, Glycogen storage disease type II, Acid alpha glucosidase, GAA
Background

Pompe disease or glycogen storage disease type II (Pompe disease, GSDII, acid maltase deficiency; OMIM; 232,300) is an autosomal recessive lysosomal storage disorder. The disease results from deficiency of alpha glucosidase (GAA) (EC. 3.2.1.20) which catalyzes the hydrolysis of glycogen to glucose, leading to the accumulation of glycogen in lysosomes of skeletal and cardiac muscles, liver and other tissues, with a variety of clinical symptoms [1–3]. The clinical spectrum is classified into infantile-onset Pompe disease (IOPD) and late-onset Pompe disease (LOPD). IOPD is characterized by onset before 12 months with predominant hypertrophic cardiomyopathy, generalized muscle weakness, hepatomegaly, feeding difficulties, and respiratory compromise. Without treatment by enzyme replacement therapy (ERT), patients with IOPD usually succumb to early death by the age of 2 years due to cardiopulmonary failure. LOPD is characterized by slowly progressive proximal muscle weakness and respiratory insufficiency, either in individuals with onset before 12 months without cardiomyopathy, or in individuals with onset after 12 months. LOPD often leads to the necessity of a wheel chair and respiratory support as well as a higher mortality rate compared with healthy individuals [4]. The incidence of Pompe disease has been reported to be about ~1 in 40,000 live births around the world [5].

The GAA gene is located on chromosome 17q25.2–25.3, spanning approximately 20 kb and containing 20 exons. The GAA cDNA has a coding sequence of 2859 bps in length and encodes a polypeptide of 952 amino acids with a calculated molecular weight of 105 kDa [6, 7]. Normally, GAA protein is synthesized as a 110-kDa precursor containing N-linked carbohydrates, and is then modified with mannose 6-phosphate groups [8]. After transport through the Golgi complex and targeting to the endosome/lysosome, the GAA precursor is proteolytically processed and undergoes N-glycan processing, resulting in 95 kDa intermediate forms and finally a 76 and 70 kDa mature protein [9].

Over 550 GAA variants are listed in The Pompe Disease Mutation Database (http://www.pompecenter.nl updated on May 2016) [10]. Nearly 70% of the variants listed are marked as pathogenic, while another 10% of the variants have unknown clinical significance. Genotype-phenotype correlation has been observed in Pompe disease to some extent. Infants who have IOPD with no cross reactive immunological material or CRIM-negative tend to have null mutations, whereas missense and splicing variants may lead to IOPD or LOPD [11, 12]. Some specific mutations are correlated with a specific subtype of Pompe disease, such as c.525delT and c.2482_2646del (exon 18 deletion) being associated with IOPD in the Dutch population [13], p.Asp645Glu associated with IOPD of Taiwanese descendants [14], and c.336-13 T > G associated with LOPD [15, 16]. CRIM status is crucial in prediction of the long-term clinical outcome of ERT and the development of antibodies against the enzyme infused. To prevent such antibody production which may reduce response to treatment, CRIM-negative patients require immunomodulation before the initiation of ERT [3]. Therefore, analysis of CRIM status is recommended for every IOPD patients [3]. Since performing CRIM study in cultured fibroblasts is not practical, a format for severity rating which has been proposed by Kroos et al. [17], is recommended to determine if the pathogenic mutation results in complete absence of GAA (CRIM-negative).

Little information is available on GAA mutations from the Southeast Asian population. Herein, we performed clinical analysis of patients with IOPD, characterization of GAA variants and functional study of the novel mutations identified, and rating its severity.

Methods

Patients

Patients with IOPD based on clinical and/or enzymatic diagnosis during 2000–2018 in the participating hospitals were enrolled in the study. Medical records were reviewed. For genetic analysis, written informed consent was obtained from all participants and the parents of the minor participants, after ethics approval was obtained from the Ramathibodi Hospital Institutional Review Board (protocol ID 06–55–46).

Leukocyte preparation and enzyme assay

Leukocytes were isolated from the whole blood samples as previously published [18]. The assay protocol to measure the acid α-glucosidase activity was performed as previously described [19].

Mutation analysis

Genomic DNA was extracted from the whole blood using QIAamp DNA Mini kit (QIAGEN GmbH). Primers were designed for all 20 exons and flanking exon-intron boundaries of the GAA gene, using primer3 Input version 0.4.0 (http://bioinfo.ut.ee/primer3-0.4.0) [20]. PCR of all exons including the noncoding exon (exon 1) was performed and subsequently subjected to capillary Sanger sequencing in both directions, at Macrogen, Inc. (Seoul, S. Korea). Multiple protein sequence alignment of vertebrate species: Homo sapiens (GenBank accession no. ABI53718.1), Pan troglodytes (GenBank accession no. JAA43384.1), Bos taurus (GenBank accession no. DAA33199.1), Mus musculus (GenBank accession no. AAH10210.1) and Rattus norvegicus (GenBank accession no. AAH61753.1), was performed by using Clustal Omega software (https://www.ebi.ac.uk/Tools/msa/clustalo). In silico analysis tools were used to predict the pathogenicity of newly identified missense mutations.
Functional analysis by expression in COS-7 cells and western blot analysis

**Construction of vectors for GAA mutants and expression in COS-7 cells**

To verify the altered function of three novel nonsense/mis-sense mutations, p.Tyr292X, p.Asp513Gly and p.Leu632Arg and the p.Ala261Thr variants (ClinVar accession number VC000456438.1), site-directed mutagenesis (QuikChange Lightning Site-Directed Mutagenesis Kit; Agilent Technology Inc., Santa Clara, CA, USA) was used to introduce those mutations in wild type GAA sequence in pcDNA3.1/CT-GFP-TOPO plasmid (Invitrogen, Carlsbad, CA, USA). The full-length GAA cDNA of a normal healthy control was PCR-amplified with the primers cDNA-GAA-F: 5′-GCCGTTGTATGCTCTAGAGC-3′ and cDNA-GAA-R: 5′-CTCCAGGTTGCTCAGATCGA-3′, and cloned into the pcDNA3.1/CT-GFP-TOPO. Primers used for mutagenesis are shown in Table 1. The methods for transient transfection of cos-7 cells were as described previously [18]. These cells were used for western blot analysis and measurement of GAA activity. The experiments were performed in triplicate.

**Western blot analysis**

We used the methodology previously described by Ngiw-sara et al., [18]. In the present study, the primary antibody used was a rabbit monoclonal anti-human GAA antibody (dilution 1:1000; #ab137068; Abcam, Cambridge, MA, USA) and the secondary antibody was polyclonal swine anti-rabbit immunoglobulin conjugated with horseradish peroxidase (dilution 1:2000; #P0217; Dako; Agilent Technology Inc., Santa Clara, CA, USA).

**Severity rating of the mutations**

The severity of novel mutations identified and the p.Ala261Thr variant were assigned to class according to the residual enzyme measured at 30°C. For more detail in-formation of severity rating, please refer the original publication by Kroos et al. [17].

### Table 1 Oligonucleotide primers used for site-directed mutagenesis in generation of GAA constructs

| Primer   | Sequence                              |
|----------|---------------------------------------|
| A261T-F  | 5′-ACAGGCTCTACCCGAGCACCCTC-3′         |
| A261T-R  | 5′-GAGGTGCTCGGAGCCCTG-3′              |
| Y292X-F  | 5′-GCAGAACCCTCTAGGGTCTACCC-3′         |
| Y292X-R  | 5′-GGGTGAGACCGGCCCTG-3′               |
| D513G-F  | 5′-CAGGTGCCCTTCGCGCCGTG-3′            |
| D513G-R  | 5′-CAATCCACATGCGGCGGAGGGACTG-3′       |
| L632R-F  | 5′-GCCAGAAATCCCGAGTTTACC-3′           |
| L632R-R  | 5′-GGTTAAACTCTGCCGAATTTCCTGG-3′       |

in culture media (M%), as expressed in cells transfected with mutant compared to wild type GAA; 2) The combination of outcome data in term of quantity and quality of apparent molecular mass of the secreted precursor 110 kDa protein in medium (M110), intracellular precursor 110 kDa (C110), intermediate 95 kDa (C95), and mature 76 kDa (C76) GAA protein, as visualized by SDS-PAGE and immunoblotting. These are given by rating into two digits. The first digit represents quantity data as follow: 1-not detectable, 2-barely detectable, 3-clearly present, but clearly less than normal, 4-normal, and 5-clearly more than normal. The second digit representing quality data is as the following: 1-too little present to judge, 2-apparent molecular mass too low, 3-apparent molecular mass too high, and 4-apparent molecular mass normal. In brief, the severity of mutation was classified into six classes as follows: A-very severe, B-potentially less severe, C-less severe, D-potentially mild, E-presumably nonpathogenic, and F-nonpathogenic. Mutation resulting in no residual GAA activity in the expressed cells (C% = 0–2) and medium (M% = 0–2), and undetectable GAA protein by western blot (M110 = 1,1; C110 = 1,1; C95 = 1,1; C76 = 1,1), in other words; missing 110 KDa band in medium and cell lysate, is classified as class ‘A’. Class ‘B’ mutation yields normal/subnormal amount of the proper sized GAA protein in the expressed cells but not always in the culture media, said otherwise; missing/visible 110 KDa band in medium and visible 110, 95, and 76 KDa bands in cell lysate, as visualized by SDS-PAGE with mutant compared to wild type GAA; 2) The combination of outcome data in term of quantity and quality of apparent molecular mass of the secreted precursor 110 kDa protein in medium (M110), intracellular precursor 110 kDa (C110), intermediate 95 kDa (C95), and mature 76 kDa (C76) GAA protein, as visualized by SDS-PAGE and immunoblotting. These are given by rating into two digits. The first digit represents quantity data as follow: 1-not detectable, 2-barely detectable, 3-clearly present, but clearly less than normal, 4-normal, and 5-clearly more than normal. The second digit representing quality data is as the following: 1-too little present to judge, 2-apparent molecular mass too low, 3-apparent molecular mass too high, and 4-apparent molecular mass normal. In brief, the severity of mutation was classified into six classes as follows: A-very severe, B-potentially less severe, C-less severe, D-potentially mild, E-presumably nonpathogenic, and F-nonpathogenic. Mutation resulting in no residual GAA activity in the expressed cells (C% = 0–2) and medium (M% = 0–2), and undetectable GAA protein by western blot (M110 = 1,1; C110 = 1,1; C95 = 1,1; C76 = 1,1), in other words; missing 110 KDa band in medium and cell lysate, is classified as class ‘A’. Class ‘B’ mutation yields normal/subnormal amount of the proper sized GAA protein in the expressed cells but not always in the culture media, said otherwise; missing/visible 110 KDa band in medium and visible 110, 95, and 76 KDa bands in cell lysate (M110 > =1,1/ C110 > =2,4; C95 > =1,1; C76 > =1,1); and along with 0–2% and 0–5% of wild-type activity in medium and cell lysate, respectively. Class ‘C’ mutation usually results in detectable protein at proper sizes (M110 > =1,1; C110 > =2,4; C95 > =2,4; C76 > =2,4), namely visible 110 KDa band in medium and clearly visible 110, 95, and 76 KDa bands in cell lysate; and 0–10% and 2–10% of enzyme activity in the medium and cells, respectively. Class ‘D’ mutation has normal or less amount of standard sized GAA protein (M110 > =2,4; C110 > =3,4; C95 > =3,4; C76 > =3,4), in other words; presence of 110 KDa band in both medium and cell lysate and presence of 95 and 76 KDa bands in cell lysate; and 5–30% of enzyme activity in the medium and cells. For mutations possessing normal or greater amount of protein of proper molecular mass (M110 > =4,4; C110 > =4,4; C95 > =4,4; C76 > =4,4), otherwise speaking; normal/less than normal presence of 110 KDa band in medium and normal presence of 110, 95, and 76 KDa band in cell lysate, those with the residual enzyme measured at 30–60% in medium and cells are assigned to class ‘E’ while those with the residual enzyme > 60% is considered class ‘F’. For more detail information of severity rating, please refer the original publication by Kroos et al. [17].
Molecular visualization of GAA mutants
The structures of human GAA alone and in complexes with inhibitors [21] were downloaded from the Protein Data Bank (http://www.rcsb.org/pdb/). PyMOL Molecular Graphics System (Schrödinger LLC) was used to display the structural features of GAA protein (PDB: 5NN8) and focusing on the novel mutations identified.

Results
Clinical characteristics
Twelve patients including nine males and three females (10 Thai and 2 from other Asian ethnicities) with IOPD were included in the study (Table 2). Eleven patients were the first affected child of the family. Only one patient (patient 11) had a history of parental consanguinity. Excluding the outlier (patient 2), the average age of onset was 4.1 months (range 2–8) and the average age at diagnosis of IOPD was 8.6 months (range 4–22). Patient 2 was noted to have mild tachypnea at 2 weeks, prompting referral to our hospital at 3–4 weeks of age due to a positive history of two previous siblings affected with cardiomyopathy of unclear etiology.

Table 2
Clinical phenotypes, clinical outcome and family history of 12 IOPD patients

| Patient | Age at onset | Age at diagnosis | Manifestation | ECG | ERT/ Clinical outcome | Parental consanguinity |
|---------|--------------|------------------|---------------|-----|-----------------------|------------------------|
| 1       | 7-8 m        | 9-10 m           | hypotonia, delayed motor and cardiomyopathy | short PR interval, BVH | yes, since 11 m./ death around 3y of age | no |
| 2       | 1-2 m        | 1-2 m            | tachypnea, mild hepatomegaly | short PR interval, BVH, EF 30% | yes, since 4 wk./ favorable outcome at 4 y | no, but two previous deceased children with cardiomyopathy |
| 3       | 3-4 m        | 5-6 m            | dyspnea | short PR interval, BVH | no, died at 13 m | no |
| 4       | 3-4 m        | 7-8 m            | dyspnea, hypotonia, delayed motor milestone | short PR interval, BVH | no/ death at 9 m from severe dehydration (diarrhea) | no |
| 5       | 5-6 m        | 11-12 m          | dyspnea, hypotonia, delayed motor milestone | short PR interval, combined ventricular hypertrophy | death from heart failure and pneumonia at the age of 1.5 y | no |
| 6       | 5-6 m        | 21-22 m          | dyspnea, poor feeding, FTT, delayed motor milestone | short PR interval, BVH, severe MR | no/ death around 2 y. | no |
| 7       | 3-4 m        | 5-6 m            | dyspnea, delayed motor milestone, hepatomegaly | short PR interval, BVH, severe MR, SDD, CK 550 u/L, EF 30% | no/ death around 1-2 y. | no |
| 8       | 1-2 m        | 3-4 m            | tachypnea, hypotonia, mild hepatomegaly | normal PR interval, BVH, SDD, minimal pericardial effusion | no/ death at 13 m. | no |
| 9       | 3-4 m        | 5-6 m            | hypotonia, delayed motor milestone, cardiomegaly | short PR interval; BVH, EF 10%, CK 1,010 u/L | no/ death around 1-2 y. | no |
| 10      | 1-2 m        | 5-6 m            | dyspnea, recurrent aspiration and pneumonia | short PR interval, BVH, SDD | no, died at 10 m. | no |
| 11      | 3-4 m        | 7-8 m            | hypotonia, hepatomegaly | BVH, SDD | no, died around 1-2 y. | yes |
| 12      | 3-4 m        | 5-6 m            | hypotonia, pneumonia with respiratory failure and failed endotracheal extubation | short PR, BVH, mild subaortic obstruction, SDD, EF 13%, CK 966 u/L | yes, since 6m. / ventilator-dependent at 18 m., motor power grade 3 | no |

BVH: Biventricular hypertrophy, CK: Creatine kinase enzyme, ECG: Electrocardiography, EF: Ejection fraction, ERT: Enzyme replacement
Patient 1 received ERT for almost 1 year before the termination of ERT due to deterioration of clinical course and death occurred at 3 years. Patient 2 received ERT since the age of 1 month. Patient 2 was able to sit with support at 8 months, sit unsupported at 12 months and walk with support at 13 months, but could not walk independently until 22 months. The patient said a meaningful word at 14 months and self-fed at 27 months. At the time of this report, patient 2 is 48 months old and has mild myopathic facies; the patient can speak in short sentences and communicate fairly, walk up and down stairs with holding, and attend local preschool. The patient’s weight and height has been around the 25–50 centiles. Patient 2 never experienced significant respiratory/medical problem since the initiation of ERT. CRIM status was not performed before the start of ERT. In addition, glucosidase alpha IgG antibody analysis using enzyme-linked immunosorbent assay (ELISA) at 34 months was negative. Patient 12 received ERT since the age of 6 months, in addition to diuretics and beta-blocker, tube-feeding, and ventilator support. Despite marked improvement of cardiomyopathy and moderate improvement of muscle weakness, patient 12 still has mild macroglossia and is ventilator-dependent at 18-months of age.

**Mutation spectrum**

Biallelic mutations of the GAA gene were identified in all patients. The GAA activity, genotypes and effect on protein phenotype are shown in Table 3. Seventeen mutations were identified, including 12 missense mutations: c.781G > A (p.Ala261Thr); c.877G > A (p.Gly293Arg), c.1003G > A (p.Gly335Arg), c.1099 T > C (p.Trp367Arg), c.1437G > C (p.Lys479Asn), c.1538A > G (p.Asp513Gly), c.1895 T > G (p.Leu632Arg), c.1935C > A (p.Asp645Glu), c.1933G > C (p.Asp645His), c.1941C > G (p.Cys647Trp), c.1942G > A (p.Gly648Ser) and c.2065G > A (p.Glu689Lys); one nonsense mutation: c.876C > G (p.Tyr292X); one splicing mutation: c.1327-2A > G (leading to exon 9 skipping and resulting in an in-frame deletion of 37 amino acid residues); one in-frame deletion mutation: c.2024_2026delACA (p.Asn675del); and two frame-shift mutations: c.1226insG (p.Asp409GlyfsX95).

| Patient | GAA activity (% of normal) | Nucleotide changes | Exons | Effects on coding protein | GAA domains | Inheritance | References (ClinVar accession number) |
|---------|---------------------------|-------------------|-------|--------------------------|-------------|-------------|--------------------------------------|
| 1       | 0                          | c.876C > G        | 5     | p.Tyr292X<sup>a</sup>    | N-terminal β-sheet | FA          | This report (SCV000925966)                      |
|         |                            | c.1003G > A      | 6     | p.Gly335Arg              | N-terminal β-sheet | MO          | Kroos et al. [12]                                   |
| 2       | 0                          | c.1935C > A      | 14    | p.Asp645Glu              | Catalytic GH31 | FA          | Hermans et al. [13]                                  |
|         |                            | c.1933G > C      | 14    | p.Asp645His              | Catalytic GH31 | MO          | Lin&Shieh [15]                                        |
| 3       | 0                          | c.1099T > C      | 7     | p.Trp367Arg              | Catalytic GH31 | FA          | Palmer et al. [16]                                   |
|         |                            | c.1942G > A      | 14    | p.Gly648Ser              | Catalytic GH31 | MO          | Huie et al. [17]                                     |
| 4       | 0                          | c.1226insG       | 8     | p.Asp409GlyfsX95         | Catalytic GH31 | FA          | This report (SCV000925969)                      |
|         |                            | c.2024_2026delACA| 14    | p.Asn675del              | Catalytic GH31 | MO          | Wan et al. [18]                                     |
| 5       | 0                          | c.1538A > G (hom) | 10    | p.Asp513Gly              | Catalytic GH31 | MO, FA      | This report (SCV000925967)                      |
| 6       | 0.43                       | c.781G > A (hom) | 4     | p.Ala261Thr              | N-terminal β-sheet | MO, FA      | ClinVar (SCV000925965)                               |
| 7       | 0.05                       | c.1411_1414delGAGA| 9     | p.Glu471fsX5             | Catalytic GH31 | MO          | Wan et al. [18]                                     |
| 8       | 1.46                       | c.1933G > C      | 14    | p.Asp645His              | Catalytic GH31 | FA          | Lin&Shieh [15]                                        |
| 9       | NA                         | c.1941C > G (hom)| 14    | p.Cys647Trp              | Catalytic GH31 | MO, FA      | Huie et al. [17]                                     |
| 10      | NA                         | c.876C > G (hom)| 5     | p.Tyr292X                | N-terminal β-sheet | MO, FA      | This report (SCV000925966)                      |
| 11      | NA                         | c.1895T > G (hom)| 14    | p.Leu632Arg<sup>b</sup>  | Catalytic GH31 | MO, FA      | This report (SCV000925968)                      |
| 12      | 0.10                       | c.1327-2A > G    | NS8   | Splicing                | -             | MO          | Kroos et al. [11]                                     |
|         |                            | c.1437G > C      | 9     | p.Lys479Asn              | Catalytic GH31 | FA          | Wan et al. [18]                                     |

<sup>a</sup>The activity is expressed as a percent relative to normal control (glucose/hr/mg protein); <sup>b</sup>the mutation creates a BfaI restriction site; <sup>c</sup>the mutant creates SfcI restriction site. PCR-restriction digest with respective enzyme revealed no mutation in 50 healthy control. IOPD infantile-onset Pompe disease, NA not available. Hom is homozygous mutation and FA and MO mean father and mother, respectively.
and c.1411_14delGAGA (p.Glu471fsX5). Among these, the p.Tyr292X, p.Asp513Gly, p.Leu632Arg, and p.Asp409GlyfsX95 variants have not been previously reported.

One novel polymorphism c.858 + 24G > C (SCV000925970) were observed in patient 7. Moreover, the variants c.1726G > A (p.Gly576Ser) and c.2065G > A (p.Glu689Lys) known as common pseudodeficiency alleles among Asian populations were presented in patient 2 (heterozygous for p.Gly576Ser) and patient 6 (homozygous for both p.Gly576Ser and p.Glu689Lys) [26]. The p.Tyr292X and p.Leu632Arg mutations create BfaI and SfcI restriction sites, respectively. These two mutations were not found in 340 control alleles (100 alleles studied by PCR-restriction digest and 240 alleles based on our in-house whole exome database). The p.Ala261Thr and p.Asp513Gly were not present in 240 control alleles in the whole exome database.

The SIFT, Polyphen-2, PredictSNP2, CADD, FATHMM, and FunSeq2 online analyses predicted that the three novel missense mutations p.Ala261Thr, p.Asp513Gly and p.Leu632Arg mutations are damaging/deleterious while the GWAVA analysis program indicated neutral change. Comparative in silico analysis of the p.Asp513Gly and p.Leu632Arg showed evolutionary conservation among vertebrate species of Aspartate513 and Leucine632 residues (Fig. 1a). The position of residue 261 is either alanine or glycine (Fig. 1a), but both these amino acids are conserved in having small, nonpolar side chains, while that of threonine is significantly larger and more polar.

Severity class of the mutations studied

Enzyme activity in cells lysates and medium, quantity and quality of the molecular mass of the GAA mutant proteins, upon transient expression in COS-7 cells, and severity class of the four novel mutations identified and the p.Ala261Thr variant are shown in Table 4 and Fig. 1b.

Compared with wild type GAA, the p.Tyr292X and p.Asp513Gly and p.Leu632Arg mutants displayed enzymatic activities around the level of the negative control from both cell lysate and culture media, whereas the p.Ala261Thr variant exhibited residual activity of approximately 66% in cell extract and 5% in culture medium.

| Species          | Sequence                        | Accession no.          |
|------------------|---------------------------------|------------------------|
| Homo sapiens     | ITGLAEHLS__QVFGDMWLI_VPEILQFN| ABI53718.1             |
| Pan troglodytes  | ITGLAEHLS__QVFGDMWLI_VPEILQFN| JAA43384.1             |
| Bos taurus       | IYGLGEIHI__QVNYGLWLI_ITGMLEFG| DAA31319.1             |
| Mus musculus     | ITGLGEHLS__QVFGDMWLI_VPDILQFN| AAH10210.1             |
| Rattus norvegicus| ITGLGEHLS__QVFGDMWLI_VPEILQFN| AAH61753.1             |

**Fig. 1** GAA protein analysis. a Protein sequence alignment of vertebrate GAA. b Western blot analysis of GAA protein in the wild type and mutants.
Molecular structure of the mutant protein

The recent release of structures of human GAA alone and complexed with inhibitors has been useful and allowed better understanding of the structural effects of the GAA mutations [21]. The GAA structure consists of an N-terminal trefoil Type-P domain, followed by a β-sheet domain, the catalytic (β/α)₈ barrel domain bearing two inserts after β-strands β3 (insert I) and β4 (insert II), and proximal and distal β-sheet domains at the C-terminus. The locations of all identified mutations in this study in the 3D structure are shown in Fig. 2a. The impacts of the novel missense mutations reported in the present study were analyzed based on the 3D structure.

The p.Ala261Thr mutation falls in a loop between the two β-sheets of the N-terminal domain of GAA that interacts tightly with the sixth α-helix of the (β/α)₈ barrel of the catalytic domain (Fig. 2b). The loop therefore helps to stabilize the interaction between the two domains. In the wild type structure, adding methyl and hydroxyl groups of the threonine and alanine will cause clashes with surrounding residues. Although this may be accommodated by small adjustments in the structure, it is likely to destabilize the interaction between the two domains somewhat. This destabilization may account for this mutant presenting only 5% of wild type GAA in medium of transfected cells, since less of the protein may fold properly and exit out of the endoplasmic reticulum.

The p.Tyr292X truncation, located at N-terminal of β-sheet domain, is predicted to eliminate most of the enzyme, including the catalytic domain, which was confirmed by undetectable GAA activity in both cell lysate and culture medium (Fig. 1b). The p.Asp513Gly and p.Leu632Arg substitutions affected the maturation of the 110 kDa precursor to intermediate and mature forms. The 110 kDa precursor was found in culture medium (M) only in p.Ala261Thr but band intensity was obviously less than the wild type GAA, possibly due to an effect on protein transport, while p.Tyr292X, p.Asp513Gly and p.Leu632Arg substitutions interfered with the synthesis of the precursor form completely (Fig. 1b).

The p.Tyr292X, p.Asp513Gly and p.Leu632Arg mutants were assigned to class ‘B’ or potentially less severe mutation. It is unclear whether the p.Ala261Thr mutation should be classified as class D or E since there is much discrepancy between the percentage of normal enzyme activity detected in the cells and medium.

Table 4 Specific activity of GAA mutants in transiently transfected COS-7 cells and severity rating

| Variant         | GAA activity* | Severity rating |
|-----------------|--------------|-----------------|
|                 | Medium | Cell lysates | M110 | C110 | C95 | C76 | M%Wt | C%Wt | Class |
| Wild type       | 60.5   | 631.3 (+ 44.5) | 4.4  | 4.4  | 4.4 | 4.4 | 100  | 100  | -     |
| p.Ala261Thr     | 3.31   | 478.3         | 3.4  | 4.4  | 4.4 | 4.4 | 5.47 | 75.8 | D/E   |
| p.Tyr292X       | 0      | 0             | 1.1  | 2.4  | 2.4 | 2.4 | 0    | 0    | B     |
| p.Asp513Gly     | 0      | 4.8           | 1.1  | 3.4  | 2.4 | 2.4 | 0    | 0.76 | B     |
| p.Leu632Arg     | 0.05   | 0.1           | 1.1  | 3.4  | 2.4 | 2.4 | 0.83 | 0.02 | B     |

*Specific activity of the wild type GAA and its mutants in transiently transfected COS-7 cells (nmol 4-MU/hr/mg protein). M%Wt and C%Wt mean the relative GAA activity normalized by the wild type GAA activity of medium and cell lysates, respectively. M110, C110, C95, C76 are the GAA protein masses detected by western blotting at 100, 95, and 76 kDa from medium (M) and cell lysate (C), respectively. Two digit numbers indicate the quantity and quality of GAA protein; 4 represents as normal while 3, 2 and 1 mean lower quantity and quality of GAA protein, respectively.

compared to the wild type control (Table 4). Western blot analysis demonstrated that the p.Ala261Thr mutant exhibited three major bands: 110 kDa precursor (C110), 95 kDa intermediate form (C95) and 76 kDa mature forms (C76), in its intracellular protein pattern, similar to the wild type GAA. As predicted, the p.Tyr292X mutant peptide resulting from early termination of protein synthesis, migrated as a major form at an aberrant molecular mass of 36 kDa (Fig. 1b). The p.Asp513Gly and p.Leu632Arg substitutions appeared to affect the maturation of the 110 kDa precursor to intermediate and mature forms. The 110 kDa precursor was found in culture medium (M) only in p.Ala261Thr but band intensity was obviously less than the wild type GAA, possibly due to an effect on protein transport, while p.Tyr292X, p.Asp513Gly and p.Leu632Arg substitutions interfered with the synthesis of the precursor form completely (Fig. 1b).

The p.Asp513Gly and p.Leu632Arg mutants were assigned to class ‘B’ or potentially less severe mutation. It is unclear whether the p.Ala261Thr mutation should be classified as class D or E since there is much discrepancy between the percentage of normal enzyme activity detected in the cells and medium.
helices of the $\beta/\alpha$$_8$ barrel. Therefore p.Asp513 has a crucial role in stabilizing the $\beta/\alpha$$_8$ structure that is responsible for holding the two catalytic residues, p.Asp518 and p.Asp616, and the substrate in the exact position for hydrolysis (Fig. 2a and c). Replacing the aspartic residue with glycine (p.Asp513Gly) will result in the loss of this charged H-bonding network, which should significantly destabilize this structure.

The p.Asp409 mutation is located in the loop after the second $\beta$-strand of the catalytic domain; therefore, the frame shift mutation c.1226insG or p.Asp409GlyfsX95 results in truncated peptide lacking most of the catalytic domain, including the two catalytic residues (p.Asp518 and p.Asp616), hence no activity is expected from the residual protein.

The p.Leu632 residue, mutated in p.Leu632Arg, is situated in helix 6 (the helix between the sixth and seventh beta strands of the $\beta/\alpha$$_8$ structure of the catalytic domain and is packed against the proximal $\beta$-sheet domain (Fig. 2a). It is in a hydrophobic environment, surrounded by the side chains of p.Val628, p.Leu636, p.Ala724, p.Pro726 and p.Leu744 (Fig. 2d); therefore, replacing the nonpolar leucine side chain with the highly polar, charged arginine side chain is expected to disrupt the hydrophobic packing of the protein, thereby explaining the lack of enzyme activity.

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Fig. 2 Locations of the mutations in the human GAA structure (PDB: 5NN8, [21]). a The positions of the novel mutations are indicated by residue sticks with blue color and the previously reported mutations are indicated by residue sticks with green color, respectively. b, c and d Close-up view of the position p.Ala261Thr, p.Asp513Gly and p.Leu632Arg, respectively.
Discussion

We described 12 patients with typical presentation of IOPD and fatal outcome early in life, with the exception of three patients who received ERT which modified the natural course of the disease. Patient 1 whose ERT was started when the patient was already on ventilator yielded unsatisfying clinical outcome, ventilator-dependence and complicated pneumonia leading to death. Despite receiving ERT at 4 weeks and having much favorable motor and cardiopulmonary outcomes, patient 2 did not achieve normal milestones. It has been proposed that for better outcome in motor and cognitive function, ERT for IOPD patient should be started before the age of 2 weeks [28, 29]. Without a family history of a previous affected child with Pompe disease, it is quite impossible to make diagnosis and start treatment at such young age. Therefore, it is important that national newborn screening for early diagnosis of IOPD should be implemented which could help the patient achieve full benefit from early treatment with ERT [29]. For patient 12, as expected, the late treatment with ERT (after 1 month) led to an improvement of cardiac manifestation but less effect on respiratory and muscle functions. Moreover, macroglossia in patient 12 may compromise the chance of becoming ventilator-independent.

Hypertrophic cardiomyopathy in the absence of typical short PR-interval and the finding of dilated cardiomyopathy could hinder differential diagnosis of Pompe disease. Most physicians, who are not familiar with Pompe disease, may rely on the presence of short PR along with hypertrophic cardiomyopathy as the diagnostic clue for IOPD, and may not be aware that IOPD may show normal PR interval and dilated cardiomyopathy. We should raise an awareness of these practical points and pitfalls among specialists involved. Despite the fact that the first ERT for Pompe disease became available in 2006, it was not until 2008 that Myozyme was accessible in Thailand, initially by compassionate use and donation from the pharmaceutical company and later commercially available in 2013.

The present study revealed that mutations in the Thai population are more diverse than originally described [30]. Amarinthnukrowh et al. reported the p.Asp645Glu mutation in 8/10 mutant alleles found among Thai patients with IOPD [30]. Including the data from the present study, the prevalence of p.Asp645Glu/His is refined to 32.4% (11/34) and exon 14 was the hot spot for GAA mutation at 50% (17/34), followed by exon 5 (Table 3) [30]. Mutation screening of exons 14 and 5, yielding 62% (21/34) detection rate, may serve as a cost-effective strategy for rapid confirmation of IOPD among Thai and Southeast Asian descendants. In addition, The mutation spectrum of this Southeast Asian cohort are partly similar to those described in Chinese and Taiwanese where the p.Asp645Glu is more common among Chinese and Taiwanese IOPD patients, and as a founder mutation [14]. However, our data was markedly different from those reported in Japanese population where the mutation at p.Asp645 was not found in 38 patients and mutations in GAA-exons 14 and 5 were rarely identified [24]. It is also important that regional/local mutation database along with CRIM and protein status should be established and publicly accessible so that future patients and treating physicians can benefit from this information.

The severity of all novel missense/nonsense mutations described in the present study are in class B-potentially less severe, implying that the mutants may have different enzyme activity in vivo [17]. It has been observed that most of class B variants were associated with severe infantile onset Pompe disease [17]. Among previously published mutations identified in the present study, p.Gly335Arg and p.Gly648Ser have been listed as class ‘B’ mutations [22]. Despite of yielding near absence of enzyme activity in the in vitro system, p.Gly648Ser was found in an adult-onset Pompe patient when it was found in compound with a mild mutation, c.-32-13 T > G [14, 31].

Previous in vitro expression studies of the mutations p.Gly293Arg, p.Trp367Arg, p.Asp645Glu, and p.Cys647Trp have confirmed significant loss of GAA activity and these variants were found in IOPD patients [23, 31–33]. The p.Asp645Glu variant has been shown to cause delayed intracellular transport, deficient phosphorylation and proteolytic processing [25]. Replacement in the equivalent position p.Asp645Asn and p.Asp645His also result in severe loss of the enzyme activity [23, 31]. The p.Asn675del, p.Glu471fsX5 and p.Lys479Asn variants were previously reported in a Chinese patient with IOPD who carried p.Asp645Glu as the second allele [34]. Up to now, all the frame-shift mutations reported in the database at http://www.pompecenter.nl (updated on May 2016) were ranked as very severe. Though the p.Asp409GlyfsX95 was not tested for functional characterization in the present study, it was convincing that this mutant allele likely falls into severe class ‘A’ mutation due to its nature of frameshift and premature truncation. The p.Asp409GlyfsX95 mutant was found in combination with the p.Asn675del in the patient 4, who exhibited complete loss of GAA activity. The c.1327-2A > G (IVS8) mutation, previously reported in an Arabic patient living in the United Kingdom and listed as a class ‘A’ mutation [17], was found in the last of our patients in association with the p.Lys479Asn mutation.

Mutations found in the IOPD patients of this study, were shown to have undetectable/little enzyme activity with the exception of p.Ala261Thr. The p.Ala261Thr mutation yielded partial reduction of enzyme activity and impaired transport (Fig. 2) and was put into class D/E mutation class by the severity rating system. The results of in silico predictive tools are in agreement in
supporting that the residue was potentially damaging. Based on the 3D structure, the position p.Ala261 is located near to the evolutionarily conserved residue p.Glu262 (Fig. 1), with the two residues are in line on the same loop in the N-terminal beta sheet domain. The p.Glu262Lys mutation was previously reported to be a deleterious mutation [35], and it is possible that p.Ala261Thr mutations can be considered to be potentially pathogenic, as found in patient 6, in whom no other deleterious mutation was identified. Notably, patient 6 had classical onset and lethal natural course of IOPD despite of p.Ala261Thr allele being classified as class D/E mutation (Table 2).

Three-fourths (12/16) of the mutations found in this IOPD cohort occurred in the catalytic GH31 domain or resulted in loss of part or all of this domain. The two novel missense mutations, p.Asp513Gly and p.Leu632Arg, were situated in this catalytic domain and were ranked as severe mutations. Homozygous mutation of p.Ala261Thr or p.Gly293Arg which were located in the N-terminal β-sheet domain resulted in little but detectable residual enzyme activity, whereas compound heterozygotes p.Tyr292X/p.Gly335Arg led to complete loss of the GAA activity.

Conclusion
The broad spectrum of GAA mutations described in the present study provide usefulness in confirming the molecular diagnosis of IOPD in the Southeast Asian population of which genetic alterations have been underrepresented in the literature. Almost all the mutations have severity class ‘A’ or ‘B’. The GAA-exons 14 and 5 are the hotspots for rapid molecular diagnosis of IOPD especially when enzymatic diagnosis is not a suitable method such as in case of urgent prenatal diagnosis without known familial mutation. These data can benefit rapid molecular diagnosis of IOPD and severity rating of the mutations can serve as a partial substitute for cross reactive immunological material (CRIM) study.

Abbreviations
CRIM: Cross-reactive immunologic material; EF: Ejection fraction; ERT: Enzyme replacement therapy; GAA: Acid alpha-glucosidase; IOPD: Infantile-onset Pompe disease; LOFD: Late-onset Pompe disease

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Authors’ contributions
LN and DW designed the experiments and prepared the manuscript draft. LN, SN, AK and RC performed enzyme activity analysis and/or mutation study. LN completed molecular cloning and expression study. PS did Western blot analysis. TT, KR, CK, SP and DW provided patient’s blood samples, collected and analyzed clinical data. LN, VC and DW supervised the experiment and the analysis of the experimental data. JKC, DW, and JS had provided critical revision to the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
All relevant data are including in the manuscript. The datasets used and/or analyzed during the current study are available from the corresponding author upon request and on the ClinVar website (https://www.ncbi.nlm.nih.gov/clinvar/) using the ClinVar accession numbers provided in Table 3.

Ethics approval and consent to participate
Written informed consents were obtained from all participants and the parents of the minor participants, following the Ramathibodi Hospital Institutional Review Board (protocol ID 06–55-46).

Consent for publication
Written informed consent to publish was obtained from the parents or legal guardians of the participants whose individual data was detailed in the manuscript.

Competing interests
The authors declare that they have no competing interests.

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