A novel mutation deep within intron 7 of the GBA gene causes Gaucher disease

Anna Malekkou1,2 | Ioanna Sevastou1 | Gavriella Mavrikiou1 | Theodoros Georgiou1,2 | Lluisa Vilageliu3 | Marina Moraitou4 | Helen Michelakakis4 | Chrystalla Prokopiou5 | Anthi Drousiotou1,2

Abstract

Background: Mutations in the GBA gene that encodes the lysosomal enzyme acid β-glucocerebrosidase cause Gaucher disease (GD), the most common lysosomal storage disorder. Most of the mutations are missense/nonsense, however, a few splicing mutations within or close to conserved consensus donor or acceptor splice sites have also been described. The aim of the study was to identify the mutation(s) in a Cypriot patient with type I GD.

Methods: The genomic DNA of the proband was screened for nine common mutations using Polymerase chain reaction–restriction fragment length polymorphism (PCR–RFLP) analysis. All exons and exon-intron boundaries, and the 5'UTR and 3'UTR regions of the GBA gene, were investigated by Sanger sequencing. RNA analysis was performed using standard procedures, and the abnormal transcript was further cloned into pGEM-T-Easy plasmid vector and sequenced. The relevant intronic region was further sequenced by the Sanger method to identify the genetic variant.

Results: A novel point mutation, g.12599C > A (c.999 + 242C > A), was detected deep in intron 7 of the GBA gene. This type of mutation has been previously described for other diseases but this is the first time, as far as we know, that it is described for GD. This mutation creates a new donor splice site leading to aberrant splicing and resulting in the insertion of the first 239nt of intron 7 as a pseudoexon in the mRNA, creating a premature stop codon.

Conclusion: This study expands the mutation spectrum of GD and highlights the importance of RNA sequencing for the molecular diagnosis of patients bearing mutations in nonexonic regions.

KEYWORDS
Cypriot, deep intronic mutation, Gaucher disease, GBA, glucocerebrosidase
1 | INTRODUCTION

Gaucher disease (GD) is the most common lysosomal storage disorder. GD follows an autosomal recessive mode of inheritance and is commonly due to loss-of-function mutations in the GBA gene (OMIM: 606,463), encoding glucocerebrosidase (GCase, E.C.3.2.1.45), and more rarely due to mutations in the PSAP gene, encoding saposin C. GCase is responsible for the degradation of glucosylceramide (Glucer) to glucose and ceramide inside the lysosomes (Brady, Kanfer, Bradley, & Shapiro, 1966). GCase deficiency results in lysosomal accumulation of Glucer in macrophages, known as ‘Gaucher cells’, which are key players in the pathophysiology of the disease. Activated macrophages secrete chitotriosidase, which reaches very high values in plasma and is used as a marker for disease progression and for monitoring response to treatment (Hollak, van Weely, van Oers, & Aerts, 1994). A common polymorphism in exon 10 of the CHIT1 gene causes deficiency of chitotriosidase, with no clinical consequences. Gaucher patients homozygous for this polymorphism are monitored using other biomarkers such as PARC-CCL18.

GD patients are generally classified into three distinct types based upon the absence or presence of neurological symptoms (Sidransky, 2012); type 1 (nonneuronopathic), type 2 (acute neuronopathic), and type 3 (subacute neuronopathic). The most common type is the nonneuronopathic type 1 (OMIM#230800) which presents with systemic manifestations such as hepatosplenomegaly, anemia, thrombocytopenia, and bone abnormalities. Type 2 (OMIM#230900) and 3 (OMIM#2301000) GD are characterized by manifestations of the central nervous system, with type 2 being more severe, with life-threatening neurological disease in infancy.

The GBA gene is located on chromosome 1q21 and consists of 11 exons and 10 introns spanning a sequence of 7.6 Kb (Ginns et al., 1985). A highly homologous pseudogene (GBAP), which shares 96% coding sequence similarity with GBA, is located 16 Kb downstream of the GBA gene (Horowitz et al., 1989). The GBAP gene contributes significantly to the generation of mutations in the GBA gene due to gene-pseudogene rearrangements (Tsuji et al., 1987), something that complicates the molecular diagnosis for GD. More than 400 different mutations have been identified so far throughout the GBA gene, including point mutations, insertions, deletions, splice-site mutations, and “complex mutations” due to recombination events of the GBA gene and the GBAP pseudogene (The Human Gene Mutation Database, http://www.hgmd.org). DNA analysis in Gaucher patients usually starts by screening for the common mutations in the population, and if negative this is followed by Sanger sequencing. More recently, a sequencing method using long reads on the Oxford Nanopore minION platform was shown to be able to detect most disease causing variants with the added advantages of phasing and intronic analysis (Leija-Salazar et al., 2019).

GD is rare in Cyprus with only three unrelated patients having been diagnosed with type I GD in the last 30 years. Two patients were found to be compound heterozygotes, with the genotype N370S/L444P. The molecular investigation of the third patient was the subject of this study.

2 | MATERIALS AND METHODS

2.1 | Ethical compliance

This work is covered by the code of Ethics of the Cyprus Institute of Neurology and Genetics. All persons who were included in this study gave their informed consent.

2.2 | Case Report

The proband is a Greek Cypriot lady who was referred to the Hematology clinic at the age of 42, as she was found to have hepatosplenomegaly (liver 8 cm, spleen 10 cm). Her hematological parameters were within normal ranges except for elevated acid phosphatase (20.6 mg/dl, normal range: 1–4.7 mg/dl), thrombocytopenia (platelets 8 × 10⁴/mm³) and hypergammaglobulinemia. Myelogram and bone marrow biopsy showed the characteristic “Gaucher cells”. Diagnosis was confirmed by measuring GCase activity in isolated leukocytes. The proband also had a very low value of plasma chitotriosidase (chito) and was found to be homozygous for the 24 bp duplication of the CHIT1 gene. The proband did not develop any neurological symptoms and was classified as Type 1 GD. The patient started enzyme replacement therapy (ERT) at the age of 53 with a biweekly dosage of 20 IU/kg. After 1 year of treatment, hemoglobin increased from 11.5 g/dl to 12.4 g/dl, and platelets increased from 47,000/mm³ to 54,000/mm³. The size of the liver and spleen was reduced by 30% and 40% respectively. The patient continues to receive ERT without any side effects. Treatment is monitored by measuring the PARC-CCL18 biomarker in plasma.

2.3 | Enzyme measurements

EDTA plasma and a lysate of isolated white blood cells from peripheral blood were used for the determination of chitotriosidase and GCase activities respectively. The activities were measured using artificial fluorescent substrates as previously described (Mavrikiou, Petrou, Georgiou, & Drousiotou, 2016; Wenger, Clark, Sattler, & Wharton, 1978).
2.4 | DNA and RNA analysis

Details of methods and kits used can be found in Tables S1-S4 in Appendix S1.

Genomic DNA from the proband, family members and control individuals was extracted from peripheral blood. Total RNA was isolated from the patient’s whole blood and control individuals, and cDNA was synthesized from 1 μg of total RNA. PCR amplification was conducted with the DNAs and cDNAs that were obtained. For restriction enzyme analysis, specific DNA fragments of the GBA gene were amplified by PCR, using specific primers that prevent the amplification of GBAP, followed by the appropriate restriction enzyme digestion (Table S2 in Appendix S1). For cloning, the desired amplified PCR fragments were purified from agarose gel and subcloned into pGEM-T-Easy plasmid vector (Promega). Randomly selected individual clones were grown in LB medium at 37°C overnight and the plasmid DNAs were isolated. The amplified PCR products and the cDNA plasmid inserts were bidirectionally sequenced using specific primers (Table S3 in Appendix S1). The resultant sequences were aligned and compared to the GBA genomic (NG_009783.1) or cDNA (NM-000157.4) sequences from Genbank reference sequence (NC_000001.11) using the NCBI Blast tool. Bioinformatics prediction analysis for possible changes in splice sites was performed using the NNSplice program from the Berkeley Drosophila Genome Project (http://fruitfly.org/seq_tools/splice.html).

RFLP analysis was performed to confirm the mutation. Nested PCR was performed using EX7_F and N370S_R followed by EX7_F and IN7_R primers (Table S4 in Appendix S1). The product was digested with the Hpy188III restriction endonuclease for 2 hours at 37°C and this was followed by electrophoresis in an agarose gel. RFLP analysis was also used to screen for the mutation in 100 samples from normal individuals.

3 | RESULTS

3.1 | Enzyme results

The diagnosis of GD in the proband was confirmed by the measurement of GCase activity which was found to be 1.7 nmol mg⁻¹ hr⁻¹ (normal range 5.4–16.8 nmol mg⁻¹ hr⁻¹). Plasma chitotriosidase activity was also very low, 1.4 nmol ml⁻¹ hr⁻¹ (normal range 9.5–44 nmol ml⁻¹ hr⁻¹), and the patient was found to be homozygous for the 24 bp duplication polymorphism (c.1049_1072dup24) in exon 10 of the CHIT1 gene. For this reason, the treatment of this patient is monitored by measuring the PARC-CCL18 biomarker in plasma. The values for the GCase and chitotriosidase activity for the proband and other family members are shown in Figure 1.

3.2 | Screening for common mutations

The genomic DNA of the patient was first screened for nine common mutations in the GBA gene: N370S, L444P, D409H, R463C, 55bpdel, IVS10-1G > A, IVS6-2A > G, R120W, and Y108C, by PCR–RFLP analysis. No mutation was detected.

**FIGURE 1** Family pedigree. Black symbols represent affected persons and symbols with a dot, carriers. The levels of GCase and chitotriosidase activity can be seen underneath each individual. Normal ranges: GCase: 5.4–16.8 nmol protein⁻¹ hr⁻¹, Chito: 9.5–44 nmol ml⁻¹ hr⁻¹
3.3 | Sanger sequencing

We then proceeded to sequence all exons of the GBA gene, including exon–intron boundaries and the 5′UTR and 3′UTR regions. No mutation was identified.

3.4 | cDNA analysis

To determine whether aberrant RNA splicing occurred, total RNA was isolated and the cDNA was generated by reverse transcriptase. Different PCR amplicons of cDNA were generated and then bidirectionally sequenced. Gel electrophoresis of the cDNA amplicon spanning exons 6–10 yielded an additional band of higher molecular weight (Figure 2a). The DNA corresponding to the two bands was separately extracted from the gel and cloned into a pGEM-TEasy vector and the clones were sequenced. The clone corresponding to the lower molecular weight fragment was identical to the normal transcript sequence (WT). However, the higher molecular weight transcript revealed an insertion of 239 bp between exons 7 and 8. After blast analysis of this sequence, this insertion was found to be identical with the first 239nt of intron 7 (Figure 2b). This finding suggested the presence of a genomic mutation leading to alternative splicing. Subsequent amplification and sequencing of the gDNA region spanning exon 7 to exon 8 revealed a novel mutation at nucleotide position g.12599 (Chr1: 155,237,099, GRCh38.p13), deep within intron 7, resulting in a C to A substitution. The proband was homozygous for this mutation.

In silico analysis using the NNsplice program, searching for potential splice site sequences in the amplified PCR product (spanning exon 7 to exon 8 including intron 7), confirmed the creation of a new donor splice site (score 1.0). The novel mutation creates a new splice donor site leading to the insertion of the first 239 nucleotides of intron 7 in mRNA, resulting in a premature stop codon.

The mutation was also found in the heterozygous state in one of the parents (the other parent is not alive), one sister, and four children of the proband, and was absent in controls (Figure 3). It is noteworthy that the parents of the proband originate from the same village.

3.5 | RFLP analysis

The mutation g.12599C > A abolishes a cleavage site for the Hpy188III restriction endonuclease: the patient showed only a single band at 753 bp, while the normal control showed two DNA fragments at 418 bp and 335 bp. RFLP analysis was used to screen for the mutation in 100 normal Greek Cypriots, originating from all major geographic areas of Cyprus. All samples were negative for the mutation.

4 | DISCUSSION

Approximately 10% of about 80,000 mutations reported in the human gene mutation database (HGMD) affect splicing. About
600 mutations that affect the pre-mRNA splicing process have been documented in patients with lysosomal storage disorders (Dardis & Buratti, 2018). Most of these splicing mutations occur within or close to conserved consensus donor (+1/+2) or acceptor (-1/-2) splice sites (Krawczak, Reiss, & Cooper, 1992) and can be detected by Sanger sequencing of PCR amplicons of exon and exon–intron boundary regions. Alterations in splice sites usually cause aberrant splicing due to disruption of recognition and interaction of the spliceosome with these sequences during mRNA splicing. Splicing mutations have also been reported in a variety of other positions that are fairly distant from splice sites (Lewandowska, 2013), such as point mutations within the branch point sequence. Instances of pathogenic mutations in deep intronic regions, other than branch point sites, have been published. Usually, these deep intronic mutations are located more than 100 bp away from exon–intron boundaries and are associated with aberrant splicing, and most commonly, with pseudoexon inclusion due to activation of a cryptic splice site. Deep intronic mutations leading to pseudoexon inclusion have been documented in multiple diseases with increasing frequency: in patients with neurofibromatosis (Cunha et al., 2016), Duchenne muscular dystrophy (Trabelsi et al., 2014), HPRT deficiency (Corrigan, Arenas, Escuredo, Fairbanks, & Marinaki, 2011), as well as several hereditary tumor syndromes, like melanoma (Harland, Mistry, Bishop, & Bishop, 2001), retinoblastoma (Dehainault et al., 2007), and breast cancer (Anczukow et al., 2012).

In the present study we have identified a novel mutation deep in intron 7 of the GBA gene in a Cypriot patient with type 1 GD. This is the first time that this type of mutation is described for GD. This mutation creates a cryptic splice donor site, leading to the creation of an aberrant transcript with an insertion of the first 239nt of intron 7 between exons 7 and 8. This transcript contains a 239 bp pseudoexon with a premature stop codon that may cause mRNA degradation due nonsense mediated mRNA decay (NMD) (Maquat, 2004). This may explain why the ° of the mutant transcript in the patient is relatively small compared to the WT transcript. Moreover, the c.999 + 242C > A mutation does not abolish the generation of WT mRNA, as it was found to be present in our patient, who is homozygous for this mutation; this may explain the relatively mild phenotype of the patient and the late onset of the disease. We have also developed an appropriate screening test, that utilizes Hpy188III restriction endonuclease digestion, for confirmation and for future efficient identification of the g.12599C > A (c.999 + 242C > A) mutation.

Mutations in the GBA gene have been shown to be associated with an increased risk for developing Parkinson’s disease, with the heterozygote carriers having about a fivefold higher risk (Gan-Or et al., 2015). In this respect it is interesting to note that the mother of our proband developed Parkinson’s disease, raising the possibility that the g.12599C > A (c.999 + 242C > A) mutation might act as a risk factor for Parkinson’s disease.

In conclusion, this study expands the mutation spectrum of GD and highlights the importance of RNA sequencing for the molecular diagnosis of patients bearing mutations in non-exonic regions.

ACKNOWLEDGMENTS

The authors thank the family members and healthy individuals who participated in this study. This study was funded by the Cyprus Institute of Neurology and Genetics.

CONFLICT OF INTEREST

The authors declare no conflicts of interest.

AUTHOR CONTRIBUTIONS

Anna Malekkou contributed to the design of the study, the execution of most of the experiments, the interpretation of the results, and the writing of the manuscript. Ioanna Sevastou, Gavriella Mavrikiou, Theodoros Georgiou, and Lluisa Vilageliu contributed to the experimental work and the interpretation of the results. Marina Moraitou and Eleni Michelakakis contributed to the experimental work, the interpretation of the results,
and writing of the manuscript. Chrystalla Prokopiou provided the clinical information on the patient. Anthi Drousiotou designed and supervised the study and contributed to the writing, revising, and editing of the manuscript. All authors approved the final version of the manuscript.

**ORCID**

Anna Malekkou https://orcid.org/0000-0001-5918-9213

Anthi Drousiotou https://orcid.org/0000-0001-5971-7637

**DATA AVAILABILITY**

The variant of this study has been submitted to ClinVar database (http://www.ncbi.nlm.nih.gov/clinvar/), reference accession number SCV00930452. Other data are available from the corresponding author upon request.

**REFERENCES**

Anczukow, O., Buisson, M., Leone, M., Coutanson, C., Lasset, C., Calender, A., … Mazoyer, S. (2012). BRCA2 deep intronic mutation causing activation of a cryptic exon: Opening toward a new preventive therapeutic strategy. *Cancer Research*, 18(18), 4903–4909. https://doi.org/10.1158/1078-0432.CCR-12-1100

Brady, R. O., Kanfer, J. N., Bradley, R. M., & Shapiro, D. (1966). Demonstration of a deficiency of glucocerebrosidase-cleaving enzyme in Gaucher’s disease. *The Journal of Clinical Investigation*, 45(7), 1112–1115. https://doi.org/10.1172/JCI105417

Corrigan, A., Arenas, M., Escuero, E., Fairbanks, L., & Marinaki, A. (2011). HPRT deficiency: Identification of twenty-four novel variants including an unusual deep intronic mutation. *Nucleosides, Nucleotides & Nucleic Acids*, 30(12), 1260–1265. https://doi.org/10.1080/15257770.2011.590172

Cunha, K., Oliveira, N., Fausto, A., de Souza, C., Gros, A., Bandres, T., … Cappellen, D. (2016). Hybridization capture-based next-generation sequencing to evaluate coding sequence and deep intronic mutations in the NFI1 gene. *Genes*, 7(12), https://doi.org/10.3390/genes71201233

Dardis, A., & Buratti, E. (2018). Impact, characterization, and rescue of pre-mRNA splicing mutations in lysosomal storage disorders. *Genes*, 9(2), https://doi.org/10.3390/genes9020073

Dehainault, C., Michaux, D., Pagès-Berhouet, S., Caux-Moncoutier, V., Doz, F., Desjardins, L., … Houdayer, C. (2007). A deep intronic mutation in the B1R gene leads to inframe sequence exonsion. *European Journal of Human Genetics*, 15(4), 473–477. https://doi.org/10.1038/sj.ejhg.5201787

Gan-Or, Z., Amshalom, I., Kilarski, L. L., Bar-Shira, A., Gana-Weisz, M., Mirelman, A., … Orr-Urtreger, A. (2015). Differential effects of severe vs mild GBA mutations on Parkinson disease. *Neurology*, 84(9), 880–887. https://doi.org/10.1212/WNL.000000000001315

Ginns, E. I., Choudary, P. V., Tsuji, S., Martin, B., Stubblefield, B., Sawyer, J., Barranger, J. A. (1985). Gene mapping and leader polypeptide sequence of human glucocerebrosidase: implications for Gaucher disease. *Proceedings of the National Academy of Sciences*, 82(20), 7101–7105. https://doi.org/10.1073/pnas.82.20.7101

Harland, M., Mistry, S., Bishop, D. T., & Bishop, J. A. (2001). A deep intronic mutation in CDKN2A is associated with disease in a subset of melanoma pedigrees. *Human Molecular Genetics*, 10(23), 2679–2686. https://doi.org/10.1093/hmg/10.23.2679

Hollak, C. E., van Weely, S., van Oers, M. H., & Aerts, J. M. (1994). Marked elevation of plasma chitotriosidase activity. A novel hallmark of Gaucher disease. *The Journal of Clinical Investigation*, 93(3), 1288–1292. https://doi.org/10.1172/JCI17084

Horowitz, M., Wilder, S., Horowitz, Z., Reiner, O., Gelbart, T., & Beutler, E. (1989). The human glucocerebrosidase gene and pseudogene: Structure and evolution. *Genomics*, 4(1), 87–96. https://doi.org/10.1016/0888-7543(89)90119-4

Krawczak, M., Reiss, J., & Cooper, D. N. (1992). The mutational spectrum of single base-pair substitutions in mRNA splice junctions of human genes: Causes and consequences. *Human Genetics*, 90(1–2), 41–54. https://doi.org/10.1007/BF00210743

Leija-Salazar, M., Sedlazeck, F. J., Toffoli, M., Mullin, S., Mokretar, K., Athanasopoulou, M., … Proukakis, C. (2019). Evaluation of the detection of GBA missense mutations and other variants using the Oxford Nanopore MinION. *Molecular Genetics & Genomic Medicine*, 7(3), e564. https://doi.org/10.1002/mgg3.564

Lewandowska, M. A. (2013). The missing puzzle piece: Splicing mutations. *International Journal of Clinical and Experimental Pathology*, 6(12), 2675–2682.

Maquat, L. E. (2004). Nonsense-mediated mRNA decay: Splicing, translation and mRNP dynamics. *Nature Reviews Molecular Cell Biology*, 5(2), 89–99. https://doi.org/10.1038/nrm1310

Mavrikiou, G., Petrou, P., Georgiou, T., & Drousiotou, A. (2016). Chitotriosidase deficiency in the Cypriot population: Identification of a novel deletion in the CHIT1 gene. *Clinical Biochemistry*, 49(12), 885–889. https://doi.org/10.1016/j.clinbiochem.2016.03.013

Sidransky, E. (2012). Gaucher disease: Insights from a rare Mendelian disorder. *Discoverie Medicine*, 14(77), 273–281.

Tnabelsi, M., Beugnet, C., Deburghgrave, N., Commere, V., Orhant, L., Leturcq, F., & Chelly, J. (2014). When a mid-intronic variation of DMD gene creates an ESE site. *Neuromuscular Disorders*, 24(12), 1111–1117. https://doi.org/10.1016/j.nmd.2014.07.003

Tsuji, S., Choudary, P. V., Martin, B. M., Stubblefield, B. K., Mayor, J. A., Barranger, J. A., & Ginns, E. I. (1987). A mutation in the human glucocerebrosidase gene in neuronopathic Gaucher’s disease. *The New England Journal of Medicine*, 316(10), 570–575. https://doi.org/10.1056/NEJM198703053161002

Wenger, D. A., Clark, C., Sattler, M., & Wharton, C. (1978). Synthetic substrate beta-glucosidase activity in leukocytes: A reproducible method for the identification of patients and carriers of Gaucher’s disease. *Clinical Genetics*, 13(2), 145–153.

**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

How to cite this article: Malekkou A, Sevastou I, Mavrikiou G, et al. A novel mutation deep within intron 7 of the GBA gene causes Gaucher disease. *Mol Genet Genomic Med*. 2020;8:e1090. https://doi.org/10.1002/mgg3.1090