Heterozygosity of the Complex Corfu $\delta^0\beta^+$ Thalassemic Allele (HBD Deletion and HBB:c.92+5G>A) Revisited

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Abstract: The Corfu $\delta^0\beta^+$ thalassemic allele is a unique thalassemic allele consisting of the simultaneous presence in cis of a deletion of the $\delta$-globin (HBD) and a single nucleotide variant in the $\beta$-globin gene (HBB) in cis has, so far, been described only in individuals of Greek origin. The heterozygosity of Corfu $\delta^0\beta^+$ is detected in 1–2% of the $\beta$-thalassemia carrier population and presents a distinct hematological phenotype of microcytic, hypochromic anemia with normal HbA2 and elevated HbF levels. The study of the Corfu $\delta^0\beta^+$ allele is important for genotype resolution, genetic counseling and prenatal/antenatal diagnosis, and the management of patients.

Keywords: Corfu $\delta^0\beta^+$ thalassemic allele; $\beta$-thalassemia variants; $\beta$-thal hematological phenotype; normal HbA2; high HbF

1. Introduction

$\delta$-thalassemia is caused by defects in the $\delta$-globin (Hemoglobin Subunit Delta, HBD) gene that result in lower HbA2 levels. Isolated $\delta$-thalassemia has no clinical significance but may confound the diagnosis of individuals with $\beta$-thalassemia. $\delta\beta$-thalassemia is a genetically heterogeneous group of disorders in which both the expression of the $\delta$-globin gene and the in cis $\beta$-globin (Hemoglobin Subunit Beta, HBB) gene are affected [1,2].

The unique complex Corfu $\delta^0\beta^+$ allele was first described in one of our patients from the Greek island of Corfu, who presented with non-transfusion dependent thalassemia at the age of 4 years with hemoglobin (Hb) of 9.2g/dL, comprising mainly HbF and low levels of HbA (5.8% with zero HbA2). The parents of the propositus were characterized...
as heterozygotes for “normal HbA₂, type 2 thalassemia”, which has a distinct hematological phenotype, with reduced red blood cells indices, decreased osmotic fragility and an unbalanced α/β-globin synthesis ratio comparable to that of heterozygotes with typical β⁺-thalassemia variants distinguished by increased HbA₂ levels [3,4]. Molecular studies on the propositus identified homozygosity of a complex allele carrying a 7.2 kb deletion δ₀ variant, partially removing δ-globin gene, in cis to IVSI-5G>A β⁺ variant [HBB:c.92+5G>A; NG_000007.3:g.57237_64443del7207] [4,5]. Further studies from our group had, very early, indicated a link between the Corfu δ₀β⁺ allele and a statistically significant raised HbF level [6].

Studying naturally occurring thalassemic variants has significantly improved our understanding of the mechanisms underlying the developmental switching of hemoglobin (Hb) during normal growth [7]. Reactivation of the γ-globin genes (Hemoglobin Subunit Gamma1/2, HBG1/HBG2) for the treatment of thalassemia and sickle cell disease has recently been the focus of many research efforts, with respective gene-editing clinical trials already showing very promising results [8].

To the best of our knowledge, the Corfu δ₀β⁺ allele has been recorded exclusively amongst β-thalassemia heterozygotes of Greek origin, whereas limited data on its presentation have been published. This survey addresses: (i) the relative prevalence of the Corfu δ₀β⁺ allele amongst all β-thalassemia variants prevailing in Greece; (ii) the precise definition hematological phenotype of Corfu δ₀β⁺ heterozygotes and (iii) the evaluation of the specific hematological phenotype in heterozygotes with the Corfu δ₀β⁺ allele compared to heterozygotes with either with β⁺ (IVSI-110 G>A) (HBB:c.93-21G>) or the most common α₀-thalassemia deletion variants found in Greece, i.e., Mediterranean type I (NG_000006.1: g.(23641_23662)(37868_37901)del IthalID:312) and 20.5 Kb (NG_000006.1: g.(18148_18200)_(37868_37901)del IthalID:314).

2. Patients and Methods

2.1. Patients

The molecular basis of β-thalassemia variants (including Corfu δ₀β⁺) was retrospectively ascertained in a total of 2558 Greek β-thalassemia heterozygotes comprising 1264 parents of 682 β-thalassemia patients followed in our Thalassemia Unit and 1294 individuals, mostly of reproductive age, referred for carrier screening between 1992 and 1998. According to the Greek thalassemia prevention program, carrier screening involves the first step of hematological phenotyping, followed by molecular genotyping driven by hematological findings consistent with β-thalassemia heterozygosity.

For the evaluation of the hematological phenotype associated with the Corfu δ₀β⁺ allele, data from 50 heterozygotes were studied and compared to those of 58 heterozygotes with the β⁺ (IVSI-110 G>A) variant and 45 heterozygotes with α₀ deletion variants (Mediterranean type I or 20.5 Kb). For this assessment, data from children, pregnant women, subjects with iron deficiency and subjects with triplicated α or α⁺ variants were excluded.

The Ethics Committee of ‘Aghia Sophia’ Children’s Hospital approved permission for medical review, a waiver of informed consent and the anonymous publication of data, according to the 1964 declaration of Helsinki and its later amendments of October 2013. (Ethical Approval Code, 19027 02/10/2021) (www.wma.net, last access 13 December 2021)

2.2. Methods

Our study is a retrospective analysis of data concerning the characterization of:

(a) The hematological phenotype based on relevant red cell parameters, including Hb (g/dL), MCV (Mean Corpuscular Volume) (fl), MCH (Mean Corpuscular Hemoglobin)(pg), RDW (Red cell Distribution Width) (%), HbA₂ (%) and HbF (%), as measured by standard hematological and biochemical methods; and

(b) The underlying genotype as evaluated with molecular methods and criteria previously described. Molecular genotyping was performed at the Laboratory of Medical
Genetics (LMG), Athens University and included methods specifically applied to detect the Corfu δ^0β^+ variant. [6,9,10]

2.3. Statistical Analysis

Statistical analysis was performed utilizing Graph-Pad Prism version 8. Descriptive statistics were calculated for all phenotype variables in the two groups. For comparison of the hematological phenotype associated with heterozygotes with the Corfu δ^0β^+ variant, the two-tailed non-paired t test was used with the statistical significance level set at p = 0.05. Tukey’s box plots were used for the graphic representation of comparative measurements of RBC indices between the three groups of heterozygotes for Corfu δ^0β^+, α^0 variants and IVS1-110G>A variants, respectively.

3. Results

3.1. Types and Prevalence of HBB Variants

Table 1 summarizes the types and prevalence of HBB variants in 2558 β-thalassemia heterozygotes; the distribution of HBB gene variants is listed in order of frequency. A total of 22 HBB gene thalassemia variants were identified: 10 null variants leading to complete impairment of β-globin synthesis (β^0); 4 with severe reduction in β-globin synthesis (β^+); 5 with mild (β^++); and 3 variants with very mild (so-called silent) reduction in β-chain synthesis (β^sil). The incidence of the “silent” variants was, in general, very low (<1%), except for 1.76% heterozygotes for the HBB:c.-151C>T (−101C>T) variant. The most prevalent variants with an incidence of ~5% and more (considered characteristic for the Greek population) were: β^+ IVSI-110 G>A with an incidence of 40.42%, followed by CD39 (17.67%); IVSI-1G>A (11.96%); IVSI-6 T>C (10.44%); and IVSII-745 G>A (4.93%) accounting for 85.4% (2.175/2.558) of all variants in the cohort. The incidence of the Corfu δ^0β^+ allele was 1.56%.

Table 1. Type and relative incidences of common and rare β thalassemia variants in a cohort from the Greek population of β thalassemia heterozygotes.
β22 c.135delC  CD44 del C
β21 c.114G>A CD37 TGG>TGA
β19 c.*96T>C +1570 T>C
β18 c.316-3C>A IVSII-848 C>A
β17 c.92G>C CD30 AGG>ACG

| N | NM_000518.5 | Thalassemia Variant | Hematologic Phenotype | Number of Cases | Frequency (%) |
|---|-------------|---------------------|----------------------|----------------|---------------|
| 17 | c.92G>C | CD30 AGG>ACG | β0 | 3 | 0.12 |
| 18 | c.316-3C>A | IVSII-848 C>A | β+ | 2 | 0.08 |
| 19 | c.−80T>A | −30 T>A | β++ | 2 | 0.08 |
| 20 | c.*96T>C | +1570 T>C | βsil | 1 | 0.04 |
| 21 | c.114G>A | CD37 TGG>TGA | β0 | 1 | 0.04 |
| 22 | c.135delC | CD44 del C | β0 | 1 | 0.04 |
| **TOTAL** | | | | **2558** | **100.00** |

* Level of suppression of β-globin synthesis: β0 = total; β+ = severe; β++ = mild; βsil = minimal. N: number

3.2. Corfu δ0β+ Hematological Phenotype

To better ascertain the relevant hematological and biochemical indices of the rare Corfu δ0β+ heterozygotes, we included data from 10 additional (50 in total) heterozygotes recruited after 1998. The findings are illustrated in Figure 1. Hemoglobin levels were reduced in both male and female heterozygotes, with a mean of 12.28 ± 1.4g/dL in males and 11.13 ± 0.81 g/dL in females. Furthermore, the hematological indices of MCV, MCH and RDW were outside the normal range, such that MCV and MCH were significantly lower and RDW significantly higher in the Corfu δ0β+ thalassemia heterozygotes (Figure 1). HbA2 levels were within the normal range (2.7 ± 0.5 %), and HbF levels varied widely, ranging between 0.2 and 9.8% (mean: 3.39 ± 2.34%, median: 2.9%) (Figure 1).

![Figure 1](image-url) Dear reviewer, please note that the image URL is placeholder. Please provide the correct URL to access the image. The figure illustrates the definition of the hematological phenotype of Corfu δ0β+; assessment of relevant hematological and biochemical parameters in 50 Corfu δ0β+ thalassemia heterozygotes; shaded areas correspond to normal range.

No significant differences in the severity of relevant red cells indices (Hb, MCV, MCH, RDW) were identified between heterozygotes with Corfu δ0β+, β+ (IVS1-110 G>A) or with αδ-thalassemia heterozygotes (Figure 2). Significant differences were noted for levels of HbA2 and HbF. The Corfu δ0β+ heterozygotes had HbA2 levels in the normal range, in contrast to heterozygotes with IVS1-110 G>A, in which HbA2 levels were increased;
and to α0-thalassemia heterozygotes, in which HbA2 levels were decreased (p < 0.001). Heterozygotes with Corfu δοβ+ had significantly higher HbF levels compared to those with β+ IVSI-110 G>A (p < 0.001). Compared to heterozygotes with α0-thalassemia deletion variants, no differences were found in the hematological phenotype of Corfu δοβ+ heterozygotes, except for the highly significant differences in HbF (p < 0.001) and the lower levels in HbA2 and RDW (Figure 2).

![Tukey's plots representing relevant parameters of hematological phenotypes in 50 Corfu δοβ+ heterozygote (boxplots with lines), 58 with IVSI-110 variant (plain boxplots) and 45 with α0 deletion thalassemia (dotted boxplots). The upper whisker span represents the 75th percentile plus 1.5 times the interquartile distance and the lower the 25th percentile minus 1.5 times the interquartile distance. Dots represent individual values falling beyond the whiskers. Shaded areas correspond to normal range. NS: Non Significant/* p < 0.05, ** p < 0.01, *** p < 0.001 and **** p < 0.0001.](image)

**Figure 2.** Tukey’s plots representing relevant parameters of hematological phenotypes in 50 Corfu δοβ+ heterozygote (boxplots with lines), 58 with IVSI-110 variant (plain boxplots) and 45 with α0 deletion thalassemia (dotted boxplots). The upper whisker span represents the 75th percentile plus 1.5 times the interquartile distance and the lower the 25th percentile minus 1.5 times the interquartile distance. Dots represent individual values falling beyond the whiskers. Shaded areas correspond to normal range. NS: Non Significant/* p < 0.05, ** p < 0.01, *** p < 0.001 and **** p < 0.0001.

4. Discussion

Studies on the molecular basis and worldwide distribution of β-thalassemia variants have identified more than 300 variants with an extremely heterogeneous distribution. Most β-thalassemia variants are rare, whereas usually, only 3–5 variants account for more than 80% of β-thalassemia variants in any given population and follow a population-specific manner.

In this study, the molecular characterization of 2558 β-thalassemia heterozygotes identified 22 β-thalassemia gene variants, five of which were the most common (~85.4%). The incidence of the Corfu δοβ+ variant allele, a unique variant so far reported exclusively in individuals of Greek origin, was 1.56%. In a previous assessment, the Corfu δοβ+ variant was shown to account for ~36% of normal HbA2 type 2 β-thalassaemia heterozygotes [6]. The remaining heterozygotes concerned the coinheritance of either ββ or β+ variants in trans or in cis with δ+ or δ- thal variants or mild β-+ thal variants [11], the detection of which may enable differential diagnosis. It is of interest that neither of the Corfu δοβ+ components, namely the Corfu HBD deletion and the HBB IVSI-5 G>A variant, have so far been observed independently in Greece and Cyprus, suggesting a founder effect for the compound allele [11].

In a similar study on 3769 Greek β-thalassemia heterozygotes, a total of 33 HBB gene variants were identified, of which the same most prevalent five variants covered
90.4% of the heterozygotes. The IVSI-5 G>A variant was detected in 14 β-thalassemia heterozygotes, whereas the δ0 of the Corfu δ0β+ variant was not assessed and may have escaped detection [12].

In this report, the largest group of patients with Corfu δ0β+ heterozygosity is presented. In this respect, we assessed the hematological phenotype of Corfu δ0β+ heterozygotes and compared it to β+-thalassemia heterozygotes with either the most common Greek variant (IVSI-110 G>A), which has similar severity to the IVSI-5 G>A β+ variant of the Corfu δ0β+ allele, or to heterozygotes with deletion type α0 thalassemia variants. As illustrated in Figure 2, hematological red cell indices were similar, whereas HbA2 and HbF levels differed. Corfu δ0β+ heterozygotes showed normal HbA2 in contrast to the significantly increased levels in IVSI-110 G>A heterozygotes and significantly lower in heterozygotes with deletion type α0 thalassemia. In respect to HbF, the majority of Corfu δ0β+ heterozygotes showed considerably elevated HbF levels, in most above 4%, similar to that of classical heterozygotes of the (δβ)0 thalassemia [6].

The clinical and hematomal phenotypes of Corfu δ0β+, either homozygotes or compound heterozygotes with β0 thalassemia variants, have been previously presented in a very small number of patients. All patients had the clinical phenotype of non-transfusion-dependent thalassemia with moderate anemia in childhood (range of Hb 7.2–9.2 g/dL), low HbA (<10%) and high levels of HbF [13]. In contrast, compound heterozygotes of other so-called “type 2 normal HbA2 thalassemic variants” with β0 or β+ variants have the clinical phenotype of thalassemia major, with severe anemia necessitating transfusions in the first years of life [3,9,11].

Gene expression studies to resolve the molecular mechanism of β-globin gene cluster regulation in the original homozygote Corfu δ0β+ patient concluded that the 7.2 kb deleted region, including the HBD gene, contains sequences important for the normal regulation of the HBG1/HBG2, HBD and HBB genes in the cluster. It appears that loss of key sequence motives in the intergenic region between HBG1 and HBD are associated with disrupted (delayed) activation of the HBB and HBD genes and a concomitant increased expression of the HBG1/HBG2 genes in cis [5,13]. Further analysis showed a 1.7kb potential repressor region upstream of HBD within the 7.2 kb Corfu deletion, which contains a possible binding site for the transcriptional repressor protein Bcl11a [14–16]. Bcl11a has been identified as a key regulator of developmental γ-globin silencing [17]. The deleted segment also contains a 250 bp sequence recognized by the chromatin remodeling PYR repressor complex, a potential regulator of hemoglobin switching, whereas chromatin conformation experiments suggested that the segment enables the locus control region of the β-globin gene cluster to activate globin expression in a developmental stage-specific manner [14] (Figure 3). Studies on primary erythroid red cell cultures measuring HBG1/2 and HBB gene transcription steady state mRNA and hemoglobin expression levels in two Corfu δ0β+ homozygotes, four compound β0 heterozygotes and two Corfu δ0β+ heterozygotes disclosed that, in the presence of the Corfu delta deletion, a post-transcriptional mechanism disrupts HBG1/2 gene silencing and potentially induces raised HbF synthesis. Thus, a combination of variants that cause a reduction in adult β-globin synthesis must be present for the Corfu delta deletion to enhance HbF production [13]. This is also supported by a report on two Italian families who carried only the Corfu HBD deletion variant, and in whom the 7.2 kb deleted DNA of the HBD gene was associated with a normal function of the “in cis” HBB and HBG1/2 globin genes [18]. Recent studies using the CRISPR-Cas9 methodology failed to show a consistent increase in HbF and came to similar conclusions that the Corfu HBD deletion requires a simultaneous disruption of the β-globin expression to lead to levels of increased γ-globin expression postnatally [14,16].
β (shaded triangle) involving the entire proposed [19]. In the case of 7,2Kb Corfu deletion, both this functional region and a PYR silencing normal HbA2 and the significantly higher HbF levels. Even in the era of next-generation gene-editing-based treatments may be supported. The hematological phenotype of Corfu insights into the ways that variants induce chromatin reconfiguration and enable reactivation or silencing of genes, like the normal HbA2 in Greece. The milder clinical phenotype in thalassemic patients with genotypes involving the Corfu type 2 normal HbAβ+ allele could be related to a disruption of a binding site involved in HBG1/2 gene silencing and/or HBB gene activation [5,13]. Additional studies on the nature and origin of this, as well as other complex variants, are likely to provide insights into the ways that variants induce chromatin reconfiguration and enable reactivation or silencing of genes, like the β-globin cluster, such that development of respective gene-editing-based treatments may be supported. The hematological phenotype of Corfu β+ heterozygotes is comparable to other β0 and β+ thalassemia heterozygotes, except the normal HbA2 and the significantly higher HbF levels. Even in the era of next-generation sequencing where a wider screening of alleles is applied, the hematological phenotype remains important and should be taken into account, especially in the presence of nor-

Figure 3. Schematic illustration of β-globin clusters (wild type and Corfu δ0β+) regarding the possible impact of BCL11A on chromatin reconfiguration and genes regulation. BCL11A has been known to enable HBG1/2 silencing through promoter interactions, although a longer-range interaction (shaded triangle) involving the entire β cluster and a region upstream HBD in specific has also been proposed [19]. In the case of 7,2Kb Corfu deletion, both this functional region and a PYR silencing complex binding site are impaired, and in the presence of the HBB:c.92+5 G>A variant, unexpected HBG1/2 gene expression (blue arrows) leads to raised HbF levels.

Limitations of the present study include its retrospective nature and the fact that the incidence of the allele was calculated on a partially selected population comprising parents of β thalassemia patients (cascade screening) and subjects from the general population selected for genotyping due to a hematological phenotype indicative of anemia and thalassemia heterozygosity. The cohort consisted of adults of reproductive age (preconception screening), with only a few (<20) children and pregnant women. Carrier screening included both β- and α-gene variants to allow accurate genotyping and proper genetic counseling. For the evaluation of the hematological phenotype only heterozygotes with normal ferritin and iron status were studied. Heterozygotes with any comorbidities and similar red cells changes were excluded.

5. Conclusions

The Corfu δ0β+ double-variant allele accounts for a substantial proportion of β-thalassemic alleles in general and a significant proportion of thalassemic alleles with type 2 normal HbA2 in Greece. The milder clinical phenotype in thalassemic patients with genotypes involving the Corfu δ0β+ allele could be related to a disruption of a binding site involved in HBG1/2 gene silencing and/or HBB gene activation [5,13]. Additional studies on the nature and origin of this, as well as other complex variants, are likely to provide insights into the ways that variants induce chromatin reconfiguration and enable reactivation or silencing of genes, like the β-globin cluster, such that development of respective gene-editing-based treatments may be supported. The hematological phenotype of Corfu δ0β+ heterozygotes is comparable to other β0 and β+ thalassemia heterozygotes, except the normal HbA2 and the significantly higher HbF levels. Even in the era of next-generation sequencing where a wider screening of alleles is applied, the hematological phenotype remains important and should be taken into account, especially in the presence of nor-
mal HbF levels, not exclusive of a Corfu $\delta^0\beta^+$ allele in need of being distinguished from an $\alpha$ thalassemic variant. The precise diagnosis of Corfu $\delta^0\beta^+$ heterozygotes is of great importance, especially in the context of genetic counseling, antenatal diagnosis and the management of patients.

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