Nationwide carrier detection and molecular characterization of β-thalassemia and hemoglobin E variants in Bangladeshi population

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Abstract

Background: β-thalassemia is one of the most common inherited blood disorders in the world and a major deterrent to the public health of Bangladesh. The management of thalassemia patients requires lifelong frequent blood transfusion and the available treatment options are unsatisfactory. A national policy on thalassemia prevention is mandatory in Bangladesh. However, precise and up-to-date information on the frequency of β-thalassemia carriers are missing due to lack of accurate diagnostic approaches, limited access to information and absence of national screening program. This study aims to determine the nationwide carrier frequency of hemoglobin E (HbE) and β-thalassemia and mutation spectrum among the carriers using molecular, hematological and biochemical methods.

Methods: The study enrolled a total of 1877 individuals (60.1% male and 39.9% female) aged between 18 and 35 years. Total sample size and its division-wise breakdown were calculated in proportion to national and division-wise population. Venous blood was collected and subjected to CBC analysis and Hb-electrophoresis for each participant. Serum ferritin was measured to detect coexistence of iron deficiency anemia with thalassemia carrier. DNA-based High Resolution Melting (HRM) curve analysis was performed for confirmation of carrier status by mutation detection.

Results: Of 11.89% (95% CI, 10.43–13.35) carriers of β-globin gene mutations, 8.68% (95% CI, 7.41–9.95) had HbE trait (ETT) and 2.24% (95% CI, 1.57–2.91) had beta-thalassemia trait (BTT). Among eight divisions, Rangpur had the highest carrier frequency of 27.1% (ETT-25%, BTT-2.1%), whereas Khulna had the lowest frequency of 4.2% (ETT-4.2% only). Moreover, α-thalassemia, HbD trait, HbE disease, hereditary persistence of HbF were detected in 0.11, 0.16, 0.43 and 0.16% participants, respectively. HRM could identify two individuals with reported pathogenic mutations in both alleles who were erroneously interpreted as carriers by hematological indices. Finally, a total of nine different mutations including a novel mutation (c.151A > G) were detected in the β-globin gene.

Conclusions: Since carrier frequency for both HbE and β-thalassemia is alarmingly high in Bangladesh, a nationwide awareness and prevention program should be made mandatory to halt the current deteriorating situations. Mutation-based confirmation is highly recommended for the inconclusive cases with conventional carrier screening methods to avoid any faulty detection of thalassemia carriers.

Keywords: β-thalassemia, Carrier frequency, HRM curve analysis, Detection accuracy, Novel mutation, National policy

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**Background**

β-thalassemia, characterized by reduced or absent β-globin chain synthesis, is one of the most common inherited blood disorders in the world and hence a major deterrent to public health. Although widespread, the major at-risk populations are mainly from Mediterranean, Middle East and Southeast Asian countries including Bangladesh [1, 2]. WHO reported that approximately 1.5% of global population are carriers of β-thalassemia and 68,000 children are born each year with various thalassemia syndromes [3]. However, precise and up-to-date data on the frequency of β-thalassemia carriers and patients are missing due to lack of accurate diagnostic approach, limited access to information and absence of national screening program in many of the thalassemia-inflicted countries.

Although thalassemia carriers are usually healthy, the patients with β-thalassemia major and HbE/β-thalassemia require lifelong blood transfusion and iron chelation therapy for survival [4]. Cardiac failure due to iron overload and transfusion-related infections have remained the leading causes of deaths of patients with thalassemia [5, 6]. Still, there is no cure for thalassemia except allogeneic bone marrow transplantation (BMT). Also, BMT therapy is too expensive to afford for majority of the world population and the outcome is often unsatisfactory. Due to lack of regular treatment and optimal health care facilities, most thalassemia patients die before adulthood in developing countries [7]. However, many at-risk populations in Cyprus, Greece, Sardinia, Iran etc. have successfully reduced the births of β-thalassemia children by adopting and implementing various preventive measures including nationwide awareness, carrier screening and prenatal diagnosis combined with genetic counseling [8–11].

Although Bangladesh has gained tremendous success in fulfilling Millennium Development Goal-4 by reducing under-5 mortality, there has been an increasingly rapid transition in the burden of disease from primarily communicable to non-communicable diseases. Thalassemia is the single most inherited monogenic blood disorder in Bangladesh and it causes not only substantial morbidity and deaths but also inflicts emotional and financial burden on the family and society [12]. An estimation using limited prevalence data has shown that roughly 33/10,000 newborns are being born each year with thalassemia in Bangladesh [12]. However, thalassemia patient-care and support facilities are barely available in most public and private hospitals. Moreover, health awareness on thalassemia remains highly inadequate among mass population. So, in addition to adoption of proper patient management strategy, prevention by premarital screening and/or prenatal diagnosis should be a useful approach for reducing the risk of thalassemia [7–9, 11]. However, considering socio-religious and financial issues as well as health risk associated with prenatal diagnosis and abortion of the affected fetus, pre-marital screening followed by genetic counseling is arguably the best approach to prevent thalassemia in Bangladesh. In order to weigh the future situation and implement an appropriate policy to tackle thalassemia, precise and up-to-date information on current carrier status is required. The last thalassemia carrier screening was conducted in 2005 on 735 school-going children in Bangladesh [13]. Moreover, the hematological and biochemical methods which are commonly used for screening purposes have limitations as these often end up in false positive and false negative results [14, 15]. In this regard, supplemental molecular methods have been in wide use for their accuracy in carrier screening and predicting severity of the thalassemia patients including their treatment, prognosis and overall management [4, 11]. Recently, the Government of Bangladesh has launched a National Thalassemia Prevention Program. As a part of that strategy, this study was conducted using funds from Non Communicable Disease Control (NCDC) Program, Director General of Health Services, MOHFW, Government of Bangladesh and Rotary Club of Dhaka North. In addition to conventional methods, the study applied DNA-based approaches to determine the accurate status of carriers and also incidence of the at-risk cases with both mutant alleles of HBB gene, which might be responsible for transition from asymptomatic to symptomatic non-transfusion dependent thalassemia in future. Accordingly, the study was conducted on a total of 1877 marriageable-aged participants. Also, a methodical approach was used to accurately determine the division-wise distribution of thalassemia carriers across Bangladesh.

**Methodology**

**Study population**

This cross-sectional study enrolled a total of 1877 participants (aged between 18–35 years) from March 2018 to February 2019 from 10 different (4 universities, 4 medical colleges, and 2 business organizations) institutions of Dhaka city with the intent to conduct awareness campaign and screen young unmarried adults. In addition, these institutes are attended by the students and employees from all 8 administrative divisions across Bangladesh. Upon obtaining written informed consent, approximately 5.0 mL of venous blood was collected from each participant via standard venipuncture in EDTA tube. The study was ethically approved by Bangladesh Medical Research Council (BMRC). After completion of the thalassemia screening tests, each participant received the report through email. Those who were found to be carriers of mutations in β-globin gene, were further given an opportunity to receive additional information and counseling.
Sample size calculation

Total sample size was calculated using the following formula and division-wise sample size was calculated in proportion to national and division-wise population size according to Bangladesh Bureau of Statistics [16].

\[
n = \frac{(\alpha + \beta)^2 \left( p_1(1-p_1) + p_2(1-p_2) \right)}{(p_2-p_1)^2} \times \text{design effect}
\]

Where,  
\( \alpha \) = the probability of type I error
\( \beta \) = the probability of type II error (power of the test)
\( p_1 = 10.2\% \) previously estimated prevalence [13]
\( p_2 = 15\% \) expected prevalence

Considering 95% confidence interval (CI), power 80% and a design effect of 2.5, the estimated sample size was 1875. To eliminate bias and for proper representation of each administrative division, samples were collected proportionately to the population size of that division by quota sampling method.

Analysis of hematological parameters

About 2.0 ml of collected whole blood were used for CBC (Complete Blood Count) analysis to determine RBC indices including hematocrit, mean corpuscular volume (MCV), mean corpuscular hemoglobin (MCH), mean corpuscular hemoglobin concentration (MCHC) and Red cell distribution width (RDW) using automated Hematology analyzer (Sysmex kx-21, Sysmex Corporation, Kobe, Japan).

Hemoglobin electrophoresis

Hemoglobin electrophoresis was performed on Sebia CAPILLARYS-2 Flex Piercing (Sebia, Lisses, France) using Capillarys Hemoglobin (E) kit to measure HbA, HbA2, HbF and other abnormal Hb variants following manufacturer’s instructions.

Serum ferritin assay

Serum ferritin was assayed on miniVIDAS® Immunoassay Analyzer (bioMérieux, USA) using VIDAS® FERRITIN kit (bioMérieux SA, Marcy, France) following manufacturer’s instructions.

Molecular analysis

Real-time PCR-based high resolution melting (HRM) curve analysis

Genomic DNA was extracted from whole blood using QIAGEN flexigene® DNA kit (Qiagen, Hilden, Germany) according to manufacturer’s guidelines. Real-time PCR and HRM curve analysis using Precision Melt Analysis™ Software (BioRad) were performed on Bio-Rad CFX96 Real-Time System. This HRM method was previously developed for mutation screening in the \( \beta \)-globin gene of Bangladeshi and regional population of thalassemic endemic countries [2, 17, 18]. This high throughput approach enables to screen mutation(s) in unknown specimens in the presence of reference samples without nucleotide sequencing as well as to screen a large number of samples in a quick and cost-effective manner.

Sanger DNA sequencing

In this study, once the mutation positive specimens with a new HRM patterns other than the references were identified, nucleotide sequencing was carried out for those samples to identify the mutation. Sanger DNA sequencing using ABI PRISM-310 software version 3.1.0 (Applied Biosystems) was performed following the polymerase chain reactions (PCR) targeting the mutational hot-spot region of HBB gene for Bangladesh (exon1, intron 1 and a portion of exon 2 of beta-globin gene) and the PCR products purification using the MinElute® PCR purification kit (Qiagen) following the manufacturer’s instructions. Then the retrieved sequence results were compared with the reference sequences (NC_000011.10) for confirmation of the mutation.

Statistical analysis

The comparison of sensitivity and specificity between traditional methods and molecular approach were performed using https://www.openepi.com/DiagnosticTest/DiagnosticTest.htm with 95% CI. The CI for an observed proportion was calculated using Stata software (version 14.2). With the known genotype frequency, the number of expected newborns with thalassemia was calculated by Hardy-Weinberg equation [19].

Results

Of the 1877 participants, male to female ratio was 1.5:1 and their average age was 23.4 ± 5.02 (mean ± SD) years (Table 1). The participants were from both rural and urban origins. About 4.32% of the participants had consanguineous parents. Moreover, although all the participants had general education, only 68.14% of them knew the term ‘thalassemia’, whereas 62.3% had no prior knowledge of the disease etiology, severity and risk factors etc. before attending the awareness program, which was arranged as a part of this study.

Screening for thalassemia carriers based on MCV, MCH and hemoglobin electrophoresis

The red blood cell count (RBC) and the hematological indices are important in the diagnosis of asymptomatic carriers as almost all kinds of thalassemia carriers show microcytic hypochromic parameters with apparently normal hemoglobin level. Mean corpuscular volume (MCV) and mean corpuscular hemoglobin (MCH) are the two most widely used RBC indices for detecting microcytic hypochromic anemia. In the present study,
β-thalassemia traits was overlooked because of low level of HbA2.

For Group 1 samples, serum ferritin was measured to confirm IDA and HRM was then performed to detect any coexistence of β-globin gene mutation with IDA. Absence of β-globin gene mutation confirmed that there was no carrier in this group and thus none of the β-thalassemia traits was overlooked because of low level of HbA2.

From the borderline suspected Group 2, a participant with 3.5% HbA2 generated a HRM curve pattern different from the wild type cluster (without mutation in HBB gene) and also did not match with any of the HRM curves previously established for all the reported mutations in Bangladesh [2]. Sanger sequencing identified and confirmed the suspected mutation as c.151A > G.
(ACT>GCT; Thr > Ala) in the HBB gene and upon BLAST with databases it was found to be a novel mutation, thereby confirming the carrier status of this participant (Fig. 1).

Identification of a single β-globin gene mutation in 41 out of 46 participants of group-3 could confirm their carrier status. However, of the rest five samples, three having 3.7% HbA2, one with 4% and one having 3.6% HbA2 turned out to be normal in HRM curve analysis and the HRM results were further confirmed by Sanger sequencing. All of these five participants had lower MCH values than the cutoff value (< 27 pg) and three of them showed MCV within normal range i.e. higher than the cutoff value (> 80 fl). Table S1 (Additional file 1: Table S1) shows the hematological features of these 5 participants having HbA2 > 3.5% without mutation in HBB gene.

Molecular analysis was also able to identify 2 participants (out of 165 HbE carriers based on Hb electrophoresis) with compound heterozygous mutations (c.79G > A + c.92 + 5G > C); one in Hb-E allele and another one in the trans-allele to HbE. These participants had relatively high levels of fetal hemoglobin (HbF of 2.8 and 11.4%), which also could justify the presence of mutations in both alleles because presence of a mutation in the allele which is trans to HbE allele (c.79G > A mutation) induces higher levels of HbF production. Hemoglobin indices of the two samples have been summarized in Table 4. However, homozygous c.79G > A was identified in all 8 participants of Group 5, thus confirming their HbE disease status. Therefore, the study identified a total of 163 HbE carriers, 42 β-thalassemia carriers and 8 participants with HbE disease.

Next we wanted to compare the sensitivity and specificity between traditional hematological analysis and HRM-based molecular approach (Table 5). Compared to the molecular method with 100% sensitivity and 100% specificity, combination of CBC and Hb electrophoresis tests showed 99.55 (95% CI, 97.51–99.92) sensitivity and 99.82 (95% CI, 99.47–99.94) specificity. Furthermore, only Hb electrophoresis gave 5 false positive and one false negative results, whereas the combination of CBC and Hb electrophoresis resulted in faulty detection in 4 cases.

**Mutation spectrum in the HBB gene of the study participants with thalassemia traits**

As shown in Table 6, out of nine different mutations including a novel mutation, the most common mutation was c.79G > A (CD 26/ HbE) (73.42%) followed by c.92 + 5G > C (14.41%).

**Frequency of thalassemia carriers and other hemoglobinopathies among Bangladeshi population**

ETT was found to be the most prevalent trait with a frequency of 8.68% (163/1877) followed by 2.24% BTT
and; thus ETT and BTT together comprised of a total frequency of 10.92% (205/1877). Moreover, participants with HbD trait, asymptomatic HbE disease, suspected NTD HbE-β-thalassemia, hereditary persistence of fetal hemoglobin (HPFH) and α-thalassemia trait were also identified (Table 7). Altogether, 11.89% (223/1877) participants were carriers of abnormal hemoglobin genes.

Contribution of consanguinity to increase thalassemia carrier frequency in the country

There were 81 (4.32%; 95% CI, 3.4–5.24) participants of consanguineous parents. The carrier frequency among the participants with history of consanguinity was 23.5% (19/81), whereas it was almost half (11.4%, 204/1796) among the children of non-consanguineous parents. The highest consanguinity was observed in Rangpur division (8/140; 5.7%), which also had the highest carrier frequency among the eight administrative divisions of Bangladesh. The findings suggest that consanguinity contributes significantly to the increased rate of thalassemia in Bangladesh.

Distribution of β-thalassemia and HbE carriers across eight divisions of Bangladesh

We found that the frequency of ETT was higher than that of the BTT across all divisions except Barisal (Fig. 2). The ETT frequency varied from as low as 4.2% (95% CI, 1.65–6.75) in Khulna Division to as high as 25% (95% CI, 17.83–32.17) in Rangpur. Conversely, the highest BTT frequency was found in Barisal Division (3.9%; 95% CI, 0.57–7.23). Unexpectedly, we could not detect any participants with BTT in Khulna Division. The highest frequency of BTT plus ETT was found in Rangpur division (27.1%; 95% CI, 19.74–34.46) followed by Rajshahi Division (16.4%; 95% CI, 11.22–21.58).

Discussion

This is the first thalassemia carrier screening study conducted among young individuals of marriageable-age in Bangladesh. The study aimed to determine (1) nationwide carrier frequencies of β-thalassemia and HbE traits more precisely using molecular approach as supplementation to hematological and electrophoretic indices for rectification of the false positive or false negative cases (2) distribution

| Table 4 Hemoglobin indices of the two participants containing compound heterozygous mutation |
|---|---|---|---|---|---|---|---|---|---|---|
| Participant No. | Age (Years) | Hb (g/dl) | MCV (fl) | MCH (pg) | RDW (%) | HbA (%) | HbF (%) | HbE (%) | HbA2 (%) |
| 1 | 20 | 9.7 | 69.6 | 20.8 | 19 | 63.5 | 11.4 | 22.1 | 3 |
| 2 | 21 | 9.9 | 66 | 19 | 19.3 | 79.3 | 2.8 | 14.3 | 3.6 |
| Screening method          | Mean sensitivity, % (95% CI) | Mean specificity, % (95% CI) | Positive predictive value, % (95% CI) | Negative predictive value, % (95% CI) | Diagnostic accuracy, % (95% CI) | True +ve (n) | False +ve (n) | True −ve (n) | False −ve (n) |
|--------------------------|-----------------------------|-----------------------------|-------------------------------------|--------------------------------------|----------------------------------|--------------|---------------|--------------|---------------|
| MCV + MCH only           | 96.04 (92.64–97.9)          | 59.76 (57.37–62.17)         | 24.72 (21.98–27.67)                | 99.1 (98.29–99.52)                  | 64.14 (61.95–66.28)               | 218          | 664           | 986          | 9             |
| Hb electrophoresis       | 99.55 (97.51–99.92)         | 99.7 (99.29–99.87)          | 97.81 (94.97–99.06)                | 99.94 (99.66–99.99)                 | 99.68 (99.3–99.85)                | 223          | 05            | 1648         | 01            |
| MCV + MCH and Hb Electrophoresis | 99.55 (97.51–99.92) | 99.82 (99.47–99.94)        | 98.67 (96.17–99.55)                | 99.94 (99.52–99.96)                 | 99.79 (99.66–99.99)               | 223          | 03            | 1650         | 01            |

CI: Confidence Interval, MCV: Mean corpuscular volume, MCH: Mean corpuscular hemoglobin
of division-wise carrier frequencies, and (3) the mutation spectrum in the β-globin gene of the carriers.

Carrier frequency of ETT plus BTT was 10.92% (95% CI, 9.51–12.33), where ETT had the highest frequency (8.68%; 95% CI, 7.41–9.95) followed by BTT (2.24%; 95% CI, 1.57–2.91). A previous study conducted in 2005 by Khan et al. reported an average frequency of 10.2% in Bangladeshi population with 6.1% ETT and 4.1% BTT [13]. This difference in frequencies between these two studies can be explained by the fact that previous studies used only conventional hematological approaches which often give false positive and false negative results [14, 15], whereas the present study applied molecular approach to avoid faulty detection of any carriers. Moreover, this study was conducted on a larger number of participants and the number of total participants as well as division-wise enrollment was enumerated statistically in proportion to the population size of each division for a precise apprehension of distribution of carrier frequency. Other studies conducted in neighboring India and Sri Lanka reported the comparable BTT frequencies of 2.68–4.05% and 2.2%, respectively, where the ETT frequencies were 3.4 and 0.5%, respectively [24–26]. However, our study showing an alarmingly high ETT frequency (8.68%) is supported by the previous study in Bangladesh [13]. Among eight administrative divisions, the highest carrier frequency was detected in Rangpur Division, where the HbE carrier frequency was 25%. Previous study showed the highest thalassemia carrier frequency in Rajshahi Division because the newly created administrative region of Rangpur was the part of Rajshahi Division during that study.

The first line of screening for identification of thalassemia carriers is a complete blood count followed by a measurement of HbA2 and HbF proportions. Since screening by hematological indices may result in misdetection of carriers due to factors like co-inheritance of α-thalassemia, mutations in β- and δ-globin genes, mutations in KLF1 gene and iron deficiency anemia [15, 27, 28], this study applied a comparatively cheaper and easy-to-perform DNA-based HRM curve analysis method to confirm and characterize mutations in the β-globin gene. This method supplements the conventional hematological and electrophoretic parameter based approaches for identification of the thalassemia carriers and patients [2]. For example, although the parameters MCV ≥ 80 fl and MCH ≥ 27 pg are usually considered as a negative indicator for HbE trait, our study using electrophoresis had identified two HbE carriers with MCV ≥ 80 fl and MCH ≥ 27 pg and a case having MCV > 80 fl and MCH < 27 pg as HbE carrier, indicating the shortcomings of hematological indices in screening of HbE carriers. Similarly, five participants with HbA2 > 3.5%, which is a widely used indicator of beta thalassemia trait, turned out to be normal by HRM curve analysis and Sanger sequencing, thereby further demonstrating shortfalls of Hb electrophoresis in detection of thalassemia carriers. The higher HbA2 might be caused by mutations in the KLF1 gene leading to borderline high HbA2 and thus may result in false positive findings in Hb electrophoresis [15]. Moreover, a number of studies reported frequent detections of high HbA2 levels in healthy individuals without any mutation in the HBB gene [29, 30]. On the other hand, specimens with HbA2 level in the borderline range (3.3–3.5%) with low MCV and/or low MCH must be subjected to DNA-based analysis to determine the carrier status of the participants. Notably, borderline HbA2 level might result from coinheritance of β-globin gene mutations with iron deficiency anemia and α-thalassemia traits that usually lower the level of HbA2 to normal or borderline range in the β-thalassemia carriers [31]. In this study, all the specimens with abnormal hematological indices were tested using high resolution melt (HRM) curve analysis to confirm the

Table 6: Mutation spectrum of β-globin gene in the β-thalassemia and HbE carriers in Bangladeshi population

| SL No. | Mutation Pattern | Number of samples, n (%; 95% CI) |
|--------|-----------------|-----------------------------------|
| 1      | c.79G > A (HbE) | 163 (73.42; 67.62–79.22)          |
| 2      | c.92 + 5G > C   | 32 (14.41; 9.8–19.02)             |
| 3      | c.79G > A + c.79G > A | 8 (3.61; 1.16–6.06)           |
| 4      | c.92 + 130G > C | 01 (0.45; 0.43–1.33)             |
| 5      | c.151A > G     | 01 (0.45; 0.43–1.33)             |
| 6      | c.126_129delCTTT| 03 (1.35; 0.16–2.86)             |
| 7      | c.27_28insG     | 02 (0.90; 0.34–2.14)             |
| 8      | c.47G > A      | 03 (1.35; 0.16–2.86)             |
| 9      | c.79G > A + c.92 + 5G > C | 02 (0.90; 0.34–2.14)         |

*Novel mutation; not reported in Bangladeshi population and also globally [23]

Table 7: Distribution of thalassemia carriers and other Hb variants among the study participants

| Types of thalassemia carriers and other Hb variants | Number of participants, n | Frequency %, (95% CI) |
|-----------------------------------------------------|---------------------------|----------------------|
| HbE Trait (ETT)                                     | 163                       | 8.68 (7.41–9.95)     |
| β-thalassemia trait (BTT)                           | 42                        | 2.24 (1.57–2.91)     |
| HbE disease                                        | 08                        | 0.43 (0.13–0.73)     |
| α-thalassemia trait                                 | 02                        | 0.11 (0.04–0.26)     |
| HbD trait                                          | 03                        | 0.16 (0.02–0.34)     |
| Suspected NTD HbE/β-thalassemia                    | 02                        | 0.11 (0.04–0.26)     |
| HPFH                                                | 03                        | 0.16 (0.02–0.34)     |
| Total carriers of mutations in one or both alleles of globin genes | 223                       | 11.89 (10.43–13.35) |

CI Confidence Interval, Hb Hemoglobin, HPFH Hereditary persistence of fetal hemoglobin, NTD Non-transfusion dependent
presence of mutation in β-globin gene and if the molecular tests were not performed, about 5 in every 1000 carriers of the β-thalassemia and HbE variants would have been missed and around 1.8% cases could have been interpreted erroneously. Therefore, although the combination of MCV, MCH and Hb electrophoresis resulted in high sensitivity and specificity, the DNA-based approaches like HRM curve analysis and Sanger sequencing had been proved to be very useful to avoid false positive and false negative results by detecting mutations in the β-globin gene, and thereby confirming the true thalassemia carrier status of the participants. In a previous study, we described the advantages and cost-effective nature of this approach over other DNA-based screening methods like Denaturing High Performance Liquid Chromatography (DHPLC), Single Strand Conformational Polymorphism (SSCP) and Denaturing Gradient Gel Electrophoresis (DGGE), Amplification Refractory Mutation System (ARMS) PCR, Sanger nucleotide sequencing etc. [2]. Notably, this real time PCR-based high throughput HRM curve analysis is easy to perform and time-saving as there is no need of post PCR amplification processes like PCR product purification, gel electrophoresis etc. All these advantages offered by the HRM-based techniques make it an ideal candidate for molecular screening of thalassemia in countries of thalassemia belt and resource limitation.

Fig. 2 Map showing the frequency of β-thalassemia and HbE carriers across eight administrative divisions of Bangladesh
More importantly, this study identified two clinically asymptomatic individuals with pathogenic mutations in both alleles of HBβ gene using HRM method, who were detected as carriers by Hb electrophoresis. Although the association between genotype and phenotype is established for both α- and β-thalassemia, differentiation into various phenotypes of thalassemia is mostly based on clinical signs and symptoms. However, the same mutations i.e. c.79 G > A + c.92 + 5G > C have been reported as pathogenic in patients with mild to severe form of HbE-beta thalassemia [32, 33]. Hence, although currently these two individuals who seem to be carriers without any clinical manifestations, they might be at risk of developing non-transfusion dependent thalassemia (NTDT) in future. In our ongoing study on the role of various genetic modifiers on clinical heterogeneity of thalassemia patients in Bangladesh, the age of first transfusion of NTDT patients ranged from 13 to 60 years (unpublished data). It should also be mentioned that patients with hemoglobin E/beta-thalassemia show different phenotypic variability at different stages of development [34]. Since numerous factors have been identified to be associated with disease severity of the NTDT patients, identification of NTDT patients is vital for prognosis because increased intestinal iron absorption in such patients increases the risk of thrombotic disease, pulmonary hypertension, sudden cardiac arrest, and liver damages etc. [34–36]. Thus the significance of diagnosis of NTDT is crucial because a timely treatment intervention will curtail the progression of disease severity and thus prevent an untimely death. All these aspects emphasize on the requirement of molecular-based carrier screening which is the ultimate confirmation of a carrier status. Furthermore, the study identified 9 different mutations including a novel mutation (c.151A > G) in the betaglobin gene of the carriers. Further studies are needed to know the pathogenesis of this novel mutation.

With the current ETT plus BTT carrier frequency of 10.92%, 9176 babies are born with thalassemia each year (according to Hardy-Weinberg equation), thereby further worsening the situations of thalassemia patients [13, 19]. Moreover, the study found that the thalassemia carrier frequency was almost double among the children with history of consanguineous marriage, a common socio-culture of this region and thus consanguinity contributes to increased burden of thalassemia. Even though majority of the participants were university/college going students, 62% of them did not know the disease etiology and about 32% did not have any knowledge about the disease prior to enrolment suggesting that the knowledge regarding thalassemia is quite insufficient among the mass population.

At present, the yearly medical cost required for thalassemia patient ranges from $1632 to $3960 in Bangladesh and there is neither a national insurance facility nor a subsidized or free treatment system from the government [12], suggesting a severe health, economic and emotional burden to the nation and thus adoption of a national thalassemia prevention strategy is a demand of time. Several thalassemia endemic countries have set up comprehensive national prevention programs, which include public awareness and education; carrier screening using molecular diagnostics, genetic counseling and prenatal diagnosis [37, 38]. Effectiveness of such prevention program in Sardinia is evidenced by a reduction in the birth rate with thalassemia major from 1:250 live births to 1:4000 and such success is also achieved by other countries including Cyprus, Iran, and Turkey [10, 38, 39]. Although carrier screening and counseling are being performed on a voluntary basis in some countries, countries like UAE, Saudi Arabia, Jordan, Cyprus, Iran and Turkey are performing compulsory premarital screening for thalassemia to discourage marriage between two carriers [38]. These strategies may help guide health policy makers of Bangladesh to adopt an appropriate thalassemia prevention strategy considering the available resources, religious values and social culture.

In summary, as prevention of thalassemia is far cheaper and better than treatment and currently no affordable cure is available, an immediate and concerted action on thalassemia prevention should be made mandatory in Bangladesh. A massive awareness program targeting general population and an intensive educational program for health personnel including physicians, nurses, health and family planning workers should be carried out nationwide promptly. Also, appropriate screening methods combining hematological, electrophoretic and molecular approaches associated with genetic counseling should be required in existing hospitals and health facilities. The information of this study will be helpful in several ways, such as measuring nationwide carrier frequency with accuracy and grasp the gravity of the situation, identifying the at-risk population and thus prioritizing them, and necessities and benefits of molecular-based carrier screening. Moreover, this study demonstrated the feasibility and usefulness of cost-effective HRM approach in resource limited settings which can be followed in other countries of thalassemia-belt for detection of HBβ gene mutations and confirmation of the carrier status.

**Conclusion**

This study highlights that adoption of a molecular screening method for detection of mutations in the HBβ gene could overcome the shortcomings of the conventional methods, in particular, for prenatal and newborn screening and for confirmation of the inconclusive cases by the traditional approaches. With the current carrier frequency, HbE/β-thalassemia will be posing a tremendous threat to the public health of Bangladesh if necessary measures like awareness program for mass population and medical personnel and; establishment of...
carrier screening facilities aligned with genetic counsel-
ing in health centers and hospitals across the country,
are not implemented immediately. Lastly, the HRM-
based cost-effective molecular methods can be initiated
in other thalassemia-prone countries and help in fighting
these non-curable and life-threatening disorders.

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s13023-020-1294-z.

**Additional file 1: Table S1.** Hematological features of the participants having high HbA2 level (HbA2 > 3.5%) without mutation in beta globin gene of hemoglobin.

**Abbreviations**

BMT: Bone Marrow Transplantation; BTT: Beta-thalassemia trait; CBC: Complete Blood Count; CI: Confidence Interval; EDTA: Ethylenediamine tetra acetate acid; ETT: HbE Trait; fl: Femtolitre; Hb: Hemoglobin; HbB: Hemoglobin beta subunit gene; HPFH: Hereditary Persistence of Fetal Hemoglobin; HRM: High Resolution Melting; IDA: Iron Deficiency Anemia; KLF1: Kupfip-like factor 1; MCH: Mean Corpuscular Hemoglobin; MCV: Mean Corpuscular Volume; NTD: Non Transfusion Dependent; Thalassemia; PCR: Polymerase Chain Reaction; pg: Pictogram; WHO: World Health Organization

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**Authors’ contributions**

FAN, GMB, MT and MH conducted the laboratory work, data analysis and manuscript draft writing. NS, SK, WIH, MR and SKG assisted in clinical information collection, interpretation and to improve the manuscript. KM, FQ, SSQ, HUS designed the study plan and supervised the overall project. All authors read and approved the final manuscript.

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**Availability of data and materials**

All relevant data are within the paper. Further information is available from the authors on request.

**Ethics approval and consent to participate**

This study was approved by the National Ethics Research Committee (NERC) of Bangladesh Medical Research Council (BMRC). Registration no # 102 03 04 2018.

**Consent for publication**

All the participants described in this article provided written informed consent for participation in the study and for publication of the results. All the authors have read and approved the paper for publication.

**Competing interests**

The authors declare that they have no competing interests.

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**References**

1. Flint J, Harding RM, Boyce AJ, Clegg JB. The population genetics of the haemoglobinopathies. Baillie’s Clin Haematol. 1998;11(1):1–51.
2. Islam MT, Sarker SR, Sultana N, Begum MN, Bhuyan GS, Talukder S, Muraduzzaman A, Alauddin M, Islam MS, Biswas PP. High resolution melting curve analysis targeting the HBB gene mutational hot-spot offers a reliable screening approach for all common as well as most of the rare beta-globin gene mutations in Bangladesh. BMC Genet. 2018;19(1):1.
3. Modell B, Darlison M. Global epidemiology of haemoglobin disorders and derived service indicators. Bull World Health Organ. 2008;86:480–7.
4. Galanello R, Origa R. Beta-thalassemia. Orphanet J Rare Dis. 2010;5(1):11.
5. Aeppos A, Farmakis D, Deftereos S, Tsironi M, Tassiopoulos S, Mouysakis I, Karagiorga M. Thalassemia heart disease: a comparative evaluation of thalassemia major and thalassemia intermedia. Chest. 2005;127(5):1523–30.
6. Zurilo M, De Stefano P, Borgia-Pignatti C, Di Palma A, Melevedi C, Piga A, Di Gregorio F, Burattini M, Terzoli S. Survival and causes of death in thalassaemia major. Lancet. 1989;344(8563):27–30.
7. Haghighpanah S, Nasirabadi S, Rahimi N, Faramarzi H, Karimi M. Sociocultural challenges of beta-thalassaemia major birth in carriers of beta-thalassaemia Iran. J Med Screen. 2012;19(3):109–11.
8. Xu X, Zhou Y, Luo G, Liao C, Zhou M, Chen P, Lu J, Jia S, Xiao G, Shen X. The prevalence and spectrum of α and β thalassemia in Guangdong Province: implications for the future health burden and population screening. J Clin Pathol. 2004;57(5):517–22.
9. Voksiardou E, Ladi E, Kattamis A, Hasapopoulos E, Economou M, Kourakli A, Maragkis K, Kontogianni K, Laffoneatis S, Yvert E, A national registry of haemoglobinopathies in Greece: deduced demographics, trends in mortality and affected births. Ann Hematol. 2012;91(10):1451–8.
10. Bozkurt G. Results from the North Cyprus thalassemia prevention program. Hemoglobin. 2007;31(2):257–64.
11. Barnett AN, Saminathan R, Chozian M. Thalassaemia screening and confirmation of carriers in parents. Best Pract Res Clin Obstet Gynaecol. 2017;39:27–40.
12. Hossain MS, Raheem TA, Ferdous S, Nahar N, Islam S, Arifuzzaman M, Razzaque MA, Alam R, Aziz S. Thalassemias in South Asia: clinical lessons learnt from Bangladesh. Orphanet J Rare Dis. 2017;12(1):93.
13. Khan W, Banu B, Amin S, Selimuzzaman M, Rahman M, Hossain B, Sanvardi G, Siddya S, Iqbal A, Rahman Y. Prevalence of beta thalassemia trait and Hb E trait in Bangladeshi school children and health burden of thalassemia in our population, DS (Child) HJ. 2005;21(1):1–7.
14. Harthoorn-Lasthuizen E, Lindemans J, Langenhuysen MC. Influence of iron deficiency anaemia on haemoglobin A2 levels: possible consequences for β/Thalassaemia screening. Scand J Clin Lab Invest. 1999;59(1):65–70.
15. Perese L, Satta S, Moi P, De Fanti F, Marunza L, Soliano MC, Barello S, Caio A, Galanello R. KLF1 gene mutations cause borderline HbA2. Blood. 2011;118(6):1454–8.
17. Boonyawat B, Monsereenusorn C, Traivaree C. Molecular analysis of beta-globin gene mutations among Thai beta-thalassemia children: results from a single center study. Appl Clin Genet. 2014;7:253.

18. George E, Teh L, Rosli R, Lai M, Tan J. Beta Thalassaemia mutations in Malays: a simplified cost-effective strategy to identify the mutations. Malaysian J Med Health Sci. 2012;28(1):1–8.

19. Wigginton JE, Cutler DJ, Abecasis GR. A note on exact tests of hardy-Weinberg equilibrium. Am J Hum Genet. 2005;76(5):887–93.

20. Viprakasit V, Origa R, Fucharoen S. Genetic basis, pathophysiology and diagnosis. In: Guidelines for the Management of Transfusion Dependent Thalassaemia (TDT). 3rd ed. Cyprus: Thalassaemia International Federation, TIF publishers; 2014.

21. Giordano P. Strategies for basic laboratory diagnostics of the hemoglobinopathies in multi-ethnic societies: interpretation of results and pitfalls. Int J Lab Hematol. 2013;35(5):465–79.

22. Stephens A, Angastiniotis M, Baysal E, Chan V, Fucharoen S, Giordano P, Hoyer J, Mosca A, Wild B. Haematology ICFTSo: ICSH recommendations for the measurement of haemoglobin A2. Int J Lab Hematol. 2012;34(1):1–13.

23. Patrinos GP, Giardine B, Riemer C, Miller W, Chui DH, Anagnou NP, Wajcman H, Hardison RC. Improvements in the HbVar database of human hemoglobin variants and thalassemia mutations for population and sequence variation studies. Nucleic Acids Res. 2004;32(suppl_1):D537–41.

24. Madan N, Sharma S, Sood S, Colah R, Bhatia H. Frequency of β-thalassemia trait and other hemoglobinopathies in northern and western India. Indian J Hum Genet. 2010;16(1):16.

25. Nagar R, Sinha S, Raman R. Haemoglobinopathies in eastern Indian states: a demographic evaluation. J Community Genet. 2015;6(1):1–8.

26. de Silva S, Fisher C, Premawardhena A, Lamabadasuriya S, Peto T, Perera G, Old J, Clegg J, Olivieri NF, Weatherall DJ. Thalassaemia in Sri Lanka: implications for the future health burden of Asian populations. Lancet. 2000;355(9206):786–91.

27. Harteveld CL, Higgs DR, α-thalassaemia. Orphanet J Rare Dis. 2010(5):1–13.

28. Galanello R, Ruggeri R, Addis M, Paglietti E, Cao A. Hemoglobin A2 in iron deficient 8-thalassemia heterozygotes. Hemoglobin. 1981;5(6):613–8.

29. Figueiredo MS. The importance of hemoglobin A2 determination. Rev Bras Hematol Hemoter. 2015;37(5):287–9.

30. Yang Z, Chaffin CH, Easley PL, Thigpen B, Reddy W. Prevalence of elevated hemoglobin A2 measured by the CAPILLARYS system. Am J Clin Pathol. 2009;131(1):42–8.

31. Denic S, Agarwal MM, Al Dabbagh B, El Essa A, Takala M, Showqi S, Yassin J. Hemoglobin a2 lowered by iron deficiency and α-thalassemia: should screening recommendation for β-thalassemia change? ISRN Hematol. 2013;2013:859294.

32. George E. HbE β-thalassaemia in Malaysia: revisited. J Hematol Thromb Dis. 2013;1(101):2.

33. Miri-Moghaddam E, Bahrami S, Naderi M, Bazi A, Karimipoor M. Molecular characterization of β-thalassemia intermedia in Southeast Iran. Hemoglobin. 2016;40(3):173–8.

34. Musallam KM, Rivella S, Vichinsky E, Rachmilewitz EA. Non-transfusion-dependent thalassemias. Haematologica. 2013;98(6):833–44.

35. strains I, Taher A, Vichinsky E, Musallam K, Cappellini M, Taper A. Non-transfusion-dependent thalassemias: an update on complications and management. Int J Mol Sci. 2018;19(1):182.

36. Taher A, Vichinsky E, Musallam K, Cappellini M-D, Viprakasit V. Guidelines for the management of non transfusion dependent thalassaemia (NTDT). Nicosia. Thalassaemia International Federation; 2013.

37. De Santis V, Kattamis C, Canatan D, Soliman AT, Elsayed F, Karimi M, Dar S, Wall Y, Yassin M, Soliman N. β-thalassemia distribution in the old world: an ancient disease seen from a historical standpoint. Mediterr J Hematol Infect Dis. 2017;9(1):e2017018.

38. Cao A, Kan YW. The prevention of thalassemia. Cold Spring Harbor perspectives in medicine. 2013;3(2):a011775.

39. Caò A, Cristina Rosati M, Galanello R. Control of β-thalassemia by carrier screening, genetic counselling and prenatal diagnosis: the Sardinian experience: In: Ciba Foundation Symposium 197-Variation in the Human Genome: Variation in the Human Genome: Ciba Foundation Symposium 197: 2007. USA: Wiley Online Library; 2007. p. 137–55.

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