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<td><strong>Author(s)</strong></td>
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C C So, A C Y So, A Y Y Chan, et al.

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Detection and characterisation of \(\beta\)-globin gene cluster deletions in Chinese using multiplex ligation-dependent probe amplification

C C So, 1 A C Y So, 1 A Y Y Chan, 1 S T Y Tsang, 1 E S K, Ma, 2 L C Chan 1

ABSTRACT

Background: Deletions in the \(\beta\)-globin cluster causing thalassaemia and hereditary persistence of fetal haemoglobin (HPFH) are uncommon and difficult to detect. Data in Chinese are very scarce.

Aims: To use a recently available technique to investigate the frequencies and nature of \(\beta\)-globin cluster deletions in Chinese.

Methods: 106 subjects with phenotypes of thalassaemia or HPFH and suspected to have deletions in the \(\beta\)-globin cluster were studied. A commercially available kit employing multiplex ligation-dependent probe amplification (MLPA) was used to screen for deletions. Gap PCR and direct nucleotide sequencing were used to characterise deletions detected.

Results: 17 deletions in the \(\beta\)-globin cluster were found in 17 patients: 8 of Chinese (\(\alpha\)\(\beta\)\(\beta\))\(\alpha\) thalassaemia, 7 of Southeast Asian (Vietnamese) deletion and 2 of Thai (\(\gamma\)\(\beta\))\(\beta\) thalassaemia. The only type of deletion detected in \(\beta\)-thalassaemia was Chinese (\(\gamma\)\(\beta\))\(\beta\) thalassaemia. The deletional form of HPFH was rarely seen in only 1 case of Thai (\(\gamma\)\(\beta\))\(\beta\) thalassaemia. Deletions presenting as \(\beta\)-thalassaemia trait and raised HbF were all of the Southeast Asian (Vietnamese) deletion type. When these deletions were co-inherited with classical \(\beta\)-thalassaemia mutations in compound heterozygous states, the phenotypes could be very variable.

Conclusions: In the Chinese population, there are only relatively few types of deletions seen in the \(\beta\)-globin cluster. MLPA is a fast and effective way of screening for these deletions. Characterisation of these deletions allows the development of simpler and more specific PCR-based tests for routine diagnostic use. Accurate prediction of phenotype is not always feasible. The molecular defects in many cases of HPFH still await discovery.

Defects in the \(\beta\)-globin gene cluster at chromosome 11p15.4 cause thalassaemias, haemoglobinopathies and fetal haemoglobin (HPFH). Deletions greater than 1 kb are seen in deletional HPFH and most \(\delta\)-\(\beta\)-thalassaemias. Deletions are distinctly uncommon in \(\beta\)-thalassaemias, which are usually due to single nucleotide substitutions or insertion/deletion of a few nucleotides in the \(\beta\)-globin gene.\(^{1}\) Screening for larger deletions is technically difficult. Routine PCR-based techniques target only known deletions, while direct nucleotide sequencing of promoters, exons and exon-intron junctions is not helpful in this setting as a normal allele is present. Deletions can be screened by Southern blotting, array comparative genomic hybridisation (CGH), quantitative PCR, fluorescence in-situ hybridisation with tiling probes and conventional cytogenetics. However, these techniques all have disadvantages of low resolution, high cost, poor throughput, or complex test design and data readout. Because of these limitations, the prevalence and nature of \(\beta\)-globin gene cluster deletions in many populations remains uncharacterised. This hinders the development of a comprehensive diagnostic algorithm and the provision of genetic counselling and prenatal diagnosis.

Multiplex ligation-dependent probe amplification (MLPA) is a recently described method that can detect mid-size deletions down to a few hundred bases or single exon level.\(^{2}\) Each probe consists of two oligonucleotides that bind adjacent to each other at the target sequence. A ligation reaction takes place and all intact probes are amplified in a PCR using one single set of labelled primers as all probes have the same primer recognition ends. Amplification products differ in size, which can be separated by electrophoresis using a standard sequencer. The peak height/area of a probe represents the amount of amplification product, which is in turn proportional to the copy number of target sequence recognised by the probe in the sample. This technique is fast and reliable and it can be applied to many hereditary and acquired diseases caused by gene dosage changes.\(^{3}\)

We used a commercial MLPA kit to screen a group of Chinese patients suspected to have deletions in the \(\beta\)-globin gene cluster. Deletions detected were further characterised and correlated with phenotypic data. The study results provide information on the prevalence of these deletions in the Chinese population and thus facilitate the development of simple and specific testing protocols.

METHODS

Subjects, phenotyping and globin genotyping

A total of 106 Chinese subjects suspected to have deletions in the \(\beta\)-globin gene cluster were identified in our records from 1998 to 2008; 102 were archive clinical samples sent for globin phenotyping and genotyping, and 4 patients were recruited from an ongoing genetic association project. Clinical and haematological data, including blood counts and haemoglobin (Hb) study results, were retrieved. HbA2 and HbF levels were measured using cation exchange high performance liquid chromatography (1998 to June 2003, Variant; July 2003–06, Variant II, BioRad, Hercules, California, USA). DNA was extracted from peripheral blood samples collected in EDTA using standard phenol/chloroform method. Routine \(\alpha\)-globin genotyping (screening for \(\alpha\)\(\alpha\), \(\alpha\)\(\delta\), \(\alpha\)\(\beta\) and \(\alpha\)\(\gamma\)) \(\beta\)-globin
Multiplex ligation-dependent probe amplification

Probes and reaction mixture for ligation and PCR were purchased from MRC-Holland (SALSA MLPA kit P102 HBB, MRC-Holland, Amsterdam, the Netherlands). Twenty-five probes spanning a 68.7 kb region of the \(-\)globin gene cluster from 5′ of the locus control region to 3′ of the \(-\)globin gene were used. HS1-4 of the \(-\)globin locus control region and all coding genes (HBE1, HBG2, HBG1, HBD, HBB) were targeted. Four DNA quantity control probes were included to safeguard adequacy of test DNA, and another ligation-dependent probe targeted at chromosome 2q14 was used to control for successful ligation reaction. Fourteen other control probes targeting different chromosomal sites other than 11p15.4 were present in the probemix for result normalisation in each sample. The SALSA MLPA P102 HBB probemix was designed by Helena Duarte and Jan Schouten at MRC-Holland. A 200 ng aliquot of sample DNA was used for each subject. Tests were performed according to the manufacturer’s instructions using a standard thermocycler (GeneAmp PCR System 9700, Applied Biosystems, Foster City, California, USA). Amplification products were separated by electrophoresis in an ABI-3130XL Genetic Analyzer (Applied Biosystems) and results analysed by GeneMapper v3.7 (Applied Biosystems). A normal control sample was run in every batch of patient samples.

MLPA data analysis

Data from GeneMapper were exported to a web-based program for analysis (Coffalyser MLPA DAT v5.4, MRC-Holland). Global normalisation was used, where all peak areas were normalised by dividing each peak area by the combined peak area of all peaks in a particular sample. From the 26 normal samples, the mean and SD of the normalised peak area for each probe was calculated. Deletion of a probe recognition sequence in a patient sample was defined as a normalised peak area below 3 SD of the mean normalised peak area for that probe in the normal controls. Heterozygous deletion (copy number of one) resulted in a 35–50% reduction of normalised peak area of a probe. Homozygous deletion (zero copy) was seen as complete absence of a peak.

Characterisation of deletions detected by MLPA

The extent of deletions detected by MLPA was compared to known \(-\)globin gene cluster deletions reported in Southeast Asians. Gap PCR was performed to screen for candidate deletions using the respective flanking primers (table 2). Details of reaction conditions are available as supplementary data on the website.

RESULTS

Performance of MLPA

No loss of probe amplification signal (no deletion) was detected in any of the 26 normal control samples. Deletion with an extent consistent with Chinese (\(^{\gamma 8}\)) thalassaemia was detected in the positive control sample previously confirmed to have this deletion. The DNA quantity controls and ligation-dependent probe amplification (screening for deletion of CTTT in codons 41/42, IVSII-nt 654 (C→T), nt -28 (A→G), codon 17 (A→T), codon 43 (G→T) and codons 71/72 (+A)), detection of anti-3.7 and anti-4.2 \(\alpha\) globin gene triplication, and direct nucleotide sequencing of \(\alpha\)- and \(\beta\)-globin gene were performed as previously described in diagnosis at all 106 subjects. A gap PCR for \(\beta\)-thalassaemia intermedia or major phenotype, but a \(\beta\)-globin gene mutation was only found in one of the parents, which was presumed but not confirmed. A deletion of the other \(\beta\)-allele could not be excluded.

Twenty-six archive normal DNA samples with normal red cell indices and haemoglobin study results, and one archive sample with known Chinese (\(^{\gamma 8}\)) thalassaemia deletion were used as negative and positive controls respectively for validation of the MLPA kit.

Table 1  Classification of 106 patients by phenotype and routine globin genotyping results

<table>
<thead>
<tr>
<th>Patient category</th>
<th>Case no.</th>
<th>MCV (fl)</th>
<th>MCH (pg)</th>
<th>HbA2 (%)</th>
<th>HbF (%)</th>
<th>Deletion detected by MLPA</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>β-thalassaemia trait, no (\beta)-globin gene mutation detected</td>
<td>4</td>
<td>60.0–83.5</td>
<td>19.6–27.8</td>
<td>4.0–5.8</td>
<td>0.3–1.0</td>
</tr>
<tr>
<td>II</td>
<td>δβ-thalassaemia trait, no (\beta)-globin gene mutation detected</td>
<td>8</td>
<td>59.6–79.8</td>
<td>17.5–27.5</td>
<td>2.3–2.8</td>
<td>6.8–21.8</td>
</tr>
<tr>
<td>III</td>
<td>HPFH trait, no (\beta)-globin gene mutation detected</td>
<td>10</td>
<td>63.1–112.2</td>
<td>20.4–38.0</td>
<td>1.0–3.5</td>
<td>5.2–64.5</td>
</tr>
<tr>
<td>IV</td>
<td>β-thalassaemia trait with raised HbF, no (\beta)-globin gene mutation detected</td>
<td>9</td>
<td>58.3–80.6</td>
<td>19.2–27.8</td>
<td>3.7–6.0</td>
<td>6.4–45.3</td>
</tr>
<tr>
<td>V</td>
<td>β-thalassaemia trait with raised HbF, one common (\beta)-globin gene mutation detected</td>
<td>16</td>
<td>58.3–75.3</td>
<td>17.8–26.7</td>
<td>1.6–6.6</td>
<td>6.9–90.0</td>
</tr>
<tr>
<td>VI</td>
<td>β-thalassaemia intermediate/major, (\beta)-globin gene mutation detected in only one parent</td>
<td>28</td>
<td>50.4–97.4</td>
<td>12.4–31.2</td>
<td>0.3–7.5</td>
<td>0.8–99.5</td>
</tr>
<tr>
<td>VII</td>
<td>β-thalassaemia intermediate/major, presumably homozygous, no family study</td>
<td>31</td>
<td>63.7–91.2</td>
<td>19.4–29.7</td>
<td>0.0–8.1</td>
<td>0.3–90.0</td>
</tr>
</tbody>
</table>

*Patient had concomitant iron deficiency anaemia.
†Two patients with low MCV had concomitant \(\alpha\)-thalassaemia trait (\(\alpha\)9-deletion); 1 patient with normal MCV and low MCH had heterozygous Hb Constant Spring mutation.
‡Most patients had received red cell transfusion.

MCH, mean corpuscular haemoglobin; MCV, mean corpuscular volume; MLPA, multiplex ligation-dependent probe amplification.
Deletion detected by MLPA

A total of 17 heterozygous deletions were detected in 17 patients, accounting for 16% of 106 subjects studied. All deletions were characterised by gap PCR using primers for known β-globin gene cluster deletions reported in Southeast Asians. Figure 1 shows the extent of the three types of deletion found, in relation to the main regulatory elements at the β-globin gene cluster. Table 3 shows clinical phenotype, haematological data and the final β-globin cluster genotype of these 17 patients.

The commonest type was Chinese (‘γδβ’)-thalassaemia (n = 8), followed by Southeast Asian (Vietnamese) deletion (n = 7) and Thai (‘γδβ’)-thalassaemia (HPFH-6) (n = 2). Deletions were mostly found in group II (δβ-thalassaemia trait, 4/8), group IV (β-thalassaemia trait with raised HbF, 5/9) and group VI (β-thalassaemia intermedia/major with β-globin gene mutation detected only in one parent, 5/28). Together they accounted for 14 of 17 deletions detected. Deletions were distinctly uncommon in group I (β-thalassaemia trait, 0/4), group III (HPFH trait, 1/10) and group V (heterozygous β-globin gene mutation with raised HbF, 0/16) (table 2).

In the heterozygous state (groups I to V), the types of deletion detected within each group were homogeneous. All four deletions found in group II (δβ-thalassaemia trait) were Chinese (‘γδβ’)-thalassaemia. All five deletions found in group IV (β-thalassaemia trait with raised HbF) were Southeast Asian (Vietnamese) deletion. The single case of deletional HPFH found in group III was Thai (‘γδβ’)-thalassaemia (HPFH-6). On the other hand, different types of deletions were seen in patients with β-thalassaemia intermedia/major in groups VI and VII, where the deletions occurred in the compound heterozygous state with a co-existing β or β′ mutation.

Of note, 2 of 31 β-thalassaemia intermedia/major patients in group VII, who were previously diagnosed to have homozygous β-globin gene mutation using routine globin genotyping methods, were found to harbour a β-globin gene cluster deletion.

DISCUSSION

This is the largest study reporting to date on globin gene deletion in thalassaemia patients using MLPA as a screening tool. Excluding group VII (patients presumably labelled as homozygous for common β-thalassaemia mutations without family study) where the pre-test probability for deletion is the lowest, the deletion detection rate in our cohort is only 20% (15/75). A further 9 patients with δβ-thalassaemia of the Chinese (‘γδβ’)-type were detected in our laboratory by gap PCR since 2003 during the study period. Inclusion of these 9 cases in our cohort would only have increased the prevalence to 28.5% (24/84). This is still substantially lower than the 61% reported in a previous Dutch study using the same method, although sample selection criteria are different. These findings suggest that there may be major differences in the prevalence of β-globin cluster deletion among different ethnic groups.

Within our Chinese cohort, relatively few types of deletion are present in defined phenotypic groups. Chinese (‘γδβ’)-thalassaemia is the only deletion detected in δβ-thalassaemia. In the heterozygous state, Southeast Asian (Vietnamese)
Deletions are not detectable in many thalassaemic subjects in our patient cohort. The underlying molecular defects leading to thalassaemia remain elusive. Non-deletional mutations in the regulatory regions in β-globin cluster can be targets for further investigation, such as the β-locus control region, the Ikaros transcription factor binding site immediately upstream of δ-globin gene which is implicated in γ/β-globin gene switching, the 3′HS-1 regulatory sequence for γ-silencing and the further 3′ erythroid-specific enhancer element. We have excluded common triplicated α-globin gene configurations by multiplex PCR, although rarer forms of α-globin gene duplication causing an intermedia phenotype in some β-thalassaemia heterozygotes are still possible. The very low detection rate in patients suspected to harbour deletional HPFH is unexpected, as many of them have significantly raised Hbf above 10% to even 50%. Further studies to look for non-deletional types of HPFH mutations within the β-globin cluster such as γ-promoter mutations are indicated. Other quantitative trait loci for Hbf expression, including chromosome 6q23 and 2p15 region have also been shown in various populations, including Chinese.

Our study clearly shows a role for MLPA in deletion screening in the β-globin cluster. The technique is fast and easy to perform, requiring only standard instruments in a routine molecular diagnostic laboratory. Results analysis is simple. It is relatively inexpensive when compared to other techniques such as Southern blotting, array CGH or quantitative PCR. However, the potential pitfalls must be borne in mind. Theoretically, small deletions of β-globin cluster not targeted by the MLPA probes can be missed, as consecutive probes are not continuous or overlapping. Loss of isolated single probes must therefore be interpreted with caution. One has to exclude polymorphism at binding sites of probes and use alternative methods to confirm any suspected small deletions.
Large deletions in the β-globin gene cluster are relatively uncommon in Chinese, particularly in the group of hereditary persistence of fetal haemoglobin.

A narrow spectrum of deletions is observed, with each deletion showing a characteristic phenotype.

However, an accurate genotype-phenotype prediction is not always possible in the compound heterozygous states.

Multiplex ligation-dependent probe amplification is a good screening method for unknown β-globin gene cluster deletions.

In conclusion, using MLPA as a screening tool, relatively few types of deletions are seen in the β-globin gene cluster in Chinese. Characterisation of these deletions allows the development of simpler and more specific PCR-based tests for their detection, a strategy that has been successfully employed for detection of common α-globin gene deletions using a multiplex gap-PCR approach. However, accurate genotype-phenotype correlation is not always feasible. Moreover, the molecular defects in many cases of HPFH will await discovery. Whether these observations can be generalised depends on results of similar large scale studies in other Chinese populations.

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Competing interests: None.

Ethics approval: Not applicable. Reason: All 102 patient samples were initially taken with proper consent for globin study during diagnostic workup which includes phenotyping and genotyping. The four subjects recruited from an ongoing genetic association study have given written consent and ethical approval has been obtained for genotyping.

Provenance and peer review: Not commissioned; externally peer reviewed.

REFERENCES


