

Short Tandem Repeats Used in Preimplantation Genetic Testing of B-Thalassemia: Genetic Polymorphisms For 15 Linked Loci in the Vietnamese Population

Dang Tien Truong¹, Ngo Van Nhat Minh¹, Dinh Phuong Nhung¹, Hoang Van Luong¹, Do Quyet¹, Tran Ngoc Anh¹, Trinh The Son¹, Nguyen Thanh Tung¹, Nguyen Thi Thu Ha², Duong Thi Phuong Anh³, Le Hoang³, Nguyen Le Thuy³, Nguyen Thi Hoa³, Nguyen Duy Bac¹, Vu Thi Nga⁴, Toi Chu Dinh⁵

¹Vietnam Military Medical University, Hanoi, Vietnam; ²National Institute of Hematology and Blood Transfusion, Hanoi, Vietnam; ³Tam Anh General Hospital, Hanoi, Vietnam; ⁴Institute for Research and Development, Duy Tan University, Danang, Vietnam; ⁵Department of Human and Animal Physiology, Faculty of Biology, Hanoi National University of Education, Hanoi, Vietnam

Abstract

Citation: Truong DT, Minh NVN, Nhung DP, Luong HV, Quyet D, Anh TN, Son TT, Tung NT, Ha NTT, Anh DTP, Hoang L, Thuy NL, Hoa NT, Bac ND, Nga VT, Chu Drinh T. Short Tandem Repeats Used in Preimplantation Genetic Testing of B-Thalassemia: Genetic Dlymrorphisms For 15 Linked Loci in the Vietnamese Population. Open Access Maced J Med Sci. 2019 Dec 30 7(24):4383-4388.

https://doi.org/10.3889/oamjms.2019.840

Keywords: Preimplantation genetic testing (PGT); β-thalassemia; Short tandem repeat (STRs); Microsatellite

*Correspondence: Toi Chu Dinh. Department of Human and Animal Physiology, Faculty of Biology, Hanoi National University of Education. E-mail: chudinhtoi@hnue.edu.vn

Received: 25-Sep-2019; Revised: 20-Nov-2019; Accepted: 21-Nov-2019; Online first: 20-Dec-2019

Copyright: © 2019 Dang Tien Truong, Ngo Van Nhat

Funding: This research did not receive any financial

Competing Interests: The authors have declared that no competing interests exist

BACKGROUND: β-thalassemia is one of the most common monogenic diseases worldwide. Preimplantation genetic testing (PGT) of β-thalassemia is performed to avoid affected pregnancies has become increasingly popular worldwide. In which, the indirect analysis using short tandem repeat (STRs) linking with HBB gene to detect different β-globin (HBB) gene mutation is a simple, accurate, economical and also provides additional control of contamination and allele-drop-out ADO.

AIM: This study established microsatellite markers for PGT of Vietnamese β-thalassemia patient.

METHODS: Fifteen (15) STRs gathered from 5 populations were identified by in silico tools within 1 Mb flanking the HBB gene. The multiplex PCR reaction was optimized and performed on 106 DNA samples from at-risk families.

RESULTS: After estimating, PIC values were ≥ 0.7 for all markers, with expected heterozygosity and observed heterozygosity values ranged from 0.81 to 0.92 and 0.53 to 0.86, respectively. One hundred percent of individuals had at least seven heterozygous markers and were found to be heterozygous for at least two markers on either side of the HBB gene.

CONCLUSION: In general, a pentadecaplex marker (all < 1 Mb from the HBB gene) assay was constituted for βthalassemia PGT on Vietnamese population.

Introduction

β-Thalassemia is one of the most common monogenic diseases, accounting for 1.5% of the population [1], concentrated in Central and Southern Asia, the Middle East, Northern Africa and the Mediterranean including Vietnam with the carrier. HBB mutations have more than 200 different types which have been knowns [2] lead to insufficient β-globin synthesis. The frequency of Vietnamese carriers ranged from 1.5 to 25.0% depending on ethnic group [3], [4], [5]. PGT-M (Preimplantation genetic testing for monogenic disease) promises to prevent monogenic

disease in children born to at-risk couples by avoiding transferring mutation embryos to women in vitro fertilization. In which, PGT-M for β-Thalassemia has become the most common application monogenic disorders.

There have been various studies that established direct or indirect PGT procedure for β-Thalassemia. Among numerous techniques, microsatellite markers such as STRs have provided many advantages in the indirect analysis. STRs are the repetitive DNA fragments of 2-6 bp which structure is highly conservative, inherited through generations and characteristic for each individual. The STRs is also highly diverse and can be amplified by PCR.

Thus, microsatellite markers linking with \emph{HBB} gene has played an essential role in linkage analysis for β -Thalassemia.

Currently, there have been many typical studies on indirect linkage-based PGT for β -thalassemia published in the world. Wen Wang (2009) combined the Nested-PCR method amplifying STR markers and minisequencing method on nine embryos and concluded five unaffected embryos for transferring [6]. Li Fan (2017) used STR markers to perform PGT on WGA products from 147 day-5 embryos and identified 24 non-mutations, 38 carriers, and 18 mutation embryos [7].

Nevertheless, these direct analysis methods have no probability of controlling contamination and ADO phenomenons, which are considered as the main reasons leading to misdiagnosis in PGT. Thus, β-Thalassemia with wide range of gene and variety of mutations is recommended applying with indirect analysis method. These methods were proved to be sensitive, accurate, reliable and rapid to control the pitfalls of PCR-based PGT, including PCR failure, contamination, and ADO.

Despite the preeminence of STRs, the limited number of researches and available markers for the Vietnamese population, this hampers their utility in linkage-based $\beta\text{-}Thalassemia$ PGT. In this study, we developed a multi-marker panel consisting of 15 STRs for Vietnamese $\beta\text{-}Thalassemia$ patients. The data suggested that the STRs set was qualified to perform PGT-M with high heterozygous values, number of heterozygous markers on each individual and the equal distribution of markers on either side of HBB gene.

Materials and methods

Control Samples for Method Optimization

One hundred six genomic DNA samples were extracted from blood and amniotic fluid of at-risk families at the Vietnam National Institute of Hematology and Blood Transfusion (NIHBT), Vietnam from 07/2016 to 6/2018. DNA was used either to prescreen of microsatellite markers or to determine heterozygosity values of them.

Short tandem repeat

Initial selected STRs were identified based on the STR database and Tandem Repeat Finder provided by Gary Benson (http://tandem.bu.edu/trf/trf.html). DNA sequence within 1 Mb upstream and downstream of the *HBB* gene (11p15.4) (genome assembly GRCh37/hg19, Feb 2009, annotation) was extracted from the UCSC

Genome Browser. Initial selection criteria for the STRs followed Machado 2009 [8]. The first microsatellite markers were subsequently compared and selected for Vietnamese population based on report from populations of Malaysia, China, and India [9]. Primers were designed by Primer3 Tool. UCSC In-silico PCR with downloaded reference DNA sequence (genome assembly GRCh37/hg19, Feb 2009, annotation) and BLAST from NCBI were used to determine and exclude the primer complementing with Alu and nonspecific sequences.

DNA Extraction

DNA was extracted from blood or amniotic fluid by Blood DNA Extraction QIAamp® DNA Mini Kit (Cat No./ID: 51304, QIAGEN) following the optimal instruction from QIAGEN Producer. Purified DNA was qualified and quantified by NanoDrop One Spectrophotometer with criteria: OD A260/A280 between 1.7-2.0 and concentration above 10 ng/ μ l.

PCR amplification

Single PCR: Followed T_m of 15 primers on http://www.operon.com/tools/oligo-analysis-tool.aspx, determined the average theoretical annealing temperature of these 15 primer pairs which is 60°C. Thus, PCR single primer was conducted according to the temperature range set at 55°C-60°C-65°C, and products were analyzed on the agarose gel. Single PCR amplification was performed in a 50 µL reaction consisting of 2 volume μL genomic (concentration: 10-20 ng/µL), 25 µL 2X QIAGEN Multiplex Master Mix, and 0.2 µM of each primer (Table 1). Thermal cycling involved an initial 15 minutes enzyme activation at 95°C, 30 cycles of denaturation at 95°C for 30 seconds, annealing at 60°C for 1 minute 30 seconds, and extension at 72°C for one minute, and a final extension at 60°C for 30 minutes.

Table 1: Singleplex PCR components (Total volume: 50 µL)

Component	Concentrattion	Volume (µL/tube)
HPLC H ₂ O		21
2X QIAGEN Multiplex MasterMix	1 X	25
Forward primer	0.2 µM	1
Reverse primer	0.2 µM	1
DNA template	·	2

Multplex PCR optimization

Multiplex PCR was performed at optimal annealing temperature and primer concentrations initially keeping at the same concentration of 0.2 μ M. Then, each primer concentration was adjusted based on product signal strength by increasing or decreasing 0.05 μ M. The multiplex PCR amplification was performed in 50 μ I reaction consisted of 1X QIAGEN Multiplex PCR Master Mix, 0.05-0.4 μ M of each primer and 100 ng DNA template; followed the protocol: an

initial 15 min enzyme activation at 95°C, 30 cycles of denaturation at 95°C for 30 seconds, annealing at 60°C for 1 min 30 s at the first cycle and 6 additional seconds for every next cycle, and extension at 72°C for 1 min.

Capillary electrophoresis

The PCR products were fluorescently labeled in a 20 μ L mixture consisted of 1X QIAGEN Multiplex MasterMix, 0.2 μ M each fluorescent primer with M13 sequences as in Table 2, following cycling condition: an initial denaturation step at 95°C for 15 min, followed by 5cycles of denaturation at 98°C for 45 s, annealing at 60°C for 1 min, and extension at 72°C for 1 min, and a final extension at 72°C for 5 min.

One μL aliquot of fluorescent PCR product was mixed with 8.5 μL of Hi-Di Formamide (Applied Biosystems, Foster City, CA, USA) and 0.5 μL of GeneScan 500 LIZ size standard, denatured at 95°C for 5 minutes, cooled to 4°C, and resolved in ABI 3130XL Fragment Analyzer (Applied Biosystems). Post-electrophoresis analysis was performed using GeneMapper 5.0 software (Applied Biosystems).

Statistical analysis

Allele frequency, PIC, expected heterozygosity (He) and Observed Heterozygosity (Ho) of the 15 microsatellite markers were calculated using Microsoft Excel.

Results

Identification of STR

Fifteen (15) STRs were identified within 1 Mb around the *HBB* gene (6 upstream STRs and nine downstream STRs) (Table 2). HBB4506 located farthest to HBB gene (0.74 Mb) and all other markers were comparatively closer. Thus, all STRs had high linkage to the HBB gene.

Table 2: Information on initial selected STR

No.	STR	Repeat	Size (bp)	Location with HBB gene	
1	HBB4506	(AC) n	366-398	Downstream	
2	D11S988	(TG) n	103-147	Downstream	
3	HBB4677	(AC) n	172-214	Downstream	
4	D11S2362	(ÀAŤ) n	87-123	Downstream	
5	HBB5089	(AC) n	241-265	Downstream	
6	D11S1243	(TG) n	220-256	Downstream	
7	HBB5138	(AC) n	404-428	Downstream	
8	HBB5178	(TG) n	158-192	Downstream	
9	HBB5205	(AGAT) n	401-449	Downstream	
10	D11S1760	(CA) n	195-241	Downstream	
11	HBB5576	(AAGG) n	327-369	Upstream	
12	HBB5655	(AC)n(AT) n	272-320	Upstream	
13	HBB5820	(AC)n(AG) n	311-331	Upstream	
14	HBB5859	(ATCT) n	375-417	Upstream	
15	D11S1338	(AC) n	137-157	Unstream	

Forward primer had an additional sequence

for fluorescent primer M13 were shown in Table 3.

Table 3: Information of primers for STR amplification

STR	Primer	Sequence (5'-3')	T _m (°C)					
HBB	B1F [*]	GTAAAACGACGGCCAGTGGTTTGACATATCTGTGAG GAAG	71.9					
4506	B1R	GTTTCAGCAAGTAAATAGGGCACTG						
D11S988	B2F**	GGTTTTCCCAGTCACGACGGACAAGAGAAAGTTGAA CATACTG	72.3					
	B2R	GTTTCCACCATTTAAGATGCCAATAAGC	63.2					
HBB	B3F*	GTAAAACGACGGCCAGTGTGTAAAAGGGCCTCTAAT CAG	72.1					
4677	B3R	GTTTCACTGATATACAAATGGCAAAGTG	61.7					
D11S236	B4F [*]	GTAAAACGACGGCCAGTGCTTCCCTRATCTGGAATG AACTC	73.2					
2	B4R	GGGTTTCCCAGTCCTTTTAC	60.4					
HBB	B5F**	GGTTTTCCCAGTCACGACCAATTTCCTTTTCTCCC CTATAC	71.4					
5089	B5R	GTGAGTCTAGCATTTGTCTTGC	60.8					
D11S124	B6F*	GTAAAACGACGGCCAGTGCTGCCCTAATTCTGTCTAC	73.3					
3	B6R	GTTGTGCACYATGAAGATACAC	59.9					
HBB	B7F*	GTAAAACGACGGCCAGTGAGAAATGTCCTTTAGAGA AATACCTTC	71.1					
5138	B7R	GTGGAGAGGAATCTRTTACTG	59.6					
HBB	B8F**	GGTTTTCCCAGTCACGACCGTAATTGCTTTCAGTACC ATTTATG	71.2					
5178	B8R	GATGTATTCGTCAACAGATAAATGG	59.7					
HBB	B9F**	GGTTTTCCCAGTCACGACCCAGGGTAGGTGACATAT AC	73.3					
5205	B9R	GTAACTCAAAAAATGGGACCCAAAC	61.3					
D11S176	B10F**	GGTTTTCCCAGTCACGACACCCTGAGTGTCTTCAAAA CTC	72.9					
0	B10R	GTTTCCAAKACTGCTGCATCATGAC	63.8					
HBB	B11F**	GGTTTTCCCAGTCACGACTCCTTCAGGTAAGAAGGA GC	73.3					
5576	B11R	CTTGAAGAGGCTAGGTGC	59.9					
HBB	B12F**	GGTTTTCCCAGTCACGACTCATTGTTTTGGTAGGTAC TGAAAG	71.4					
5655	B12R	AGTTGTAGTAAGTTTGTCAGGCTA	59.4					
HBB	B13F [*]	GTAAAACGACGGCCAGTGCTGAGATTATTTATACAGC AACACTTG	71.1					
5820	B13R	GTTTCCAGTTATTGGTTGCTTTAGATTAC	63.3					
HBB	B14F**	GGTTTTCCCAGTCACGACCTGTCTATTTCATCTGTCA GCTTC	72.5					
5859	B14R	GTTTAAAGTGTTGGCGTGAGC	60.6					
D11S133	B15F [*]	GTAAAACGACGGCCAGTGAAGGACACACAGATTCAC TTAAAG	71.5					
8	B15R T _m means	GCTACTTATTTGGAGTGTGAATTTC	59.7 66.6					
	ı _m ın c ails		50.0					

^{*} Primer fluorescently labeled with HEX; ** Primer fluorescently labeled with FAM.

Multiplex PCR Optimization

Annealing temperature (T_a) and primer concentration optimization were performed. The results of T_a optimization of 15 primer were shown in Figure 1.

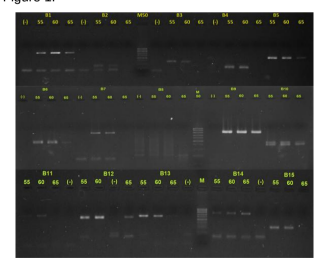


Figure 1: Annealing temperature optimization of 15 STRs. Annealing temperature at 60° C gave clear bands and most uniform signal. The products were electrophoresed on 1% agarose gel, at 120V for 30 minutes. The 10 μ l sample was each well

The optimal annealing temperature showed clear bands and fair signals for all 15 primers (Figure 1). We finally concluded that 60°C was the optimal annealing temperature.

After adjusting primer concentrations (0.05-0.4 μ M), optimal Multiplex PCR gave clear peaks. The PCR products were fluorescently labeled for capillary electrophoresis (Figure 2).

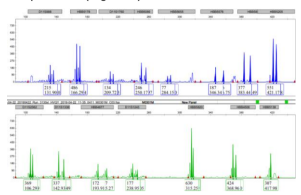


Figure 2: Representative electropherogram of multiplex PCR product after optimization

Polymorphism evaluation

To determine the polymorphism and informativeness of the 15 markers, the allele frequencies, PIC, He, and Ho values of each marker were calculated. Most STRs sequences were successfully amplified in all individuals except D11S1760 (84/106) and HBB5655 (81/106) caused by PCR Failure.

In summary, 270 alleles were observed with 11-28 alleles per marker. Allele frequency ranged from 0.0047 to 0.3255 (Table 4). All marker had PIC values of \geq 0.7 and among these markers ranged, HBB5178 showed the lowest polymorphism (0.79), and D11S1760 showed the highest polymorphism (0.91).

The data showed that He values ranged from 0.81 (HBB5138) to 0.92 (D11S1760), and all markers had Ho values of \geq 0.5, guaranteeing high polymorphic information. The Ho values of markers ranged from 0.53 (HBB5655)-0.86 (HBB5205). Thus, HBB5205 was the most informative marker, while HBB5655 was the least informative marker. Furthermore, the number of heterozygous markers of each individual was also counted.

Data showed 100% of individuals had at least seven heterozygous markers (Figure 4). Also, all were observed to be heterozygous for at least two markers on either side of the *HBB* gene (Figure 5). Based on the results, all 15 markers are high in polymorphism and informativeness for the Vietnamese population.

Table 4: Distribution of observed allele frequencies of 15 microsatellite markers

HBB45	06	D	11S988	H	BB4677	D1	1S2362	НВ	B5089
Allele	Frequency	Allele	Frequency	Allele	Frequency	Allele	Frequency	Allele	Frequenc
364 369 371 372 373 375 377 380 381 382 383 384 385 386 387 388 390	0.0094 0.0283 0.0566 0.0047 0.1651 0.0755 0.0094 0.1651 0.1462 0.066 0.0425 0.1509 0.0142 0.0142 0.0047 0.0047	115 116 120 121 122 123 124 125 126 129 130 131 132 134 137 138 139 140 141 143	0.0472 0.0236 0.0047 0.0142 0.0283 0.0189 0.2783 0.0142 0.1038 0.1274 0.0047 0.0047 0.033 0.0189 0.0613 0.1132 0.0189 0.0189 0.0047 0.0047	173 179 183 184 185 186 187 188 190 191 192 193 194 195 200 201 203 204 205 206 207 208 210 212	0.0047 0.0047 0.0283 0.0047 0.2123 0.0283 0.0047 0.0047 0.0047 0.0047 0.0036 0.0236 0.0236 0.0236 0.0236 0.0236 0.036 0.036 0.047 0.047 0.047 0.047 0.047 0.047 0.047	99 101 103 104 105 106 107 108 119 111 112 113 114 115 116 118 119 121	0.0048 0.0095 0.0095 0.0429 0.0048 0.1095 0.2048 0.0048 0.219 0.0857 0.0095 0.1 0.1095 0.019 0.0286 0.0143 0.0048	242 243 244 248 249 250 251 252 253 254 255 256 257 258 259 260 242 243 244	0.0142 0.0425 0.0189 0.0142 0.0236 0.0991 0.1321 0.0472 0.0896 0.1274 0.283 0.0142 0.0094 0.0189 0.0142 0.0425 0.0425
D11S1	243	н	3B5138	213 HI	0.0047 BB5178	н	BB5205	D11	S1760
Allele	Frequency	Allele	Frequency	Allele	Frequency	Allele	Frequency	Allele	Frequenc
224 235 236 237 238 239 240 241 242 243 244 245 246 247 248 249 250 254	0.0047 0.0047 0.0094 0.0094 0.0283 0.0943 0.1415 0.1651 0.1651 0.1085 0.2406 0.033 0.066 0.0189 0.0283 0.047	404 405 406 414 415 416 417 418 419 420 421 422 423 424	0.0802 0.3208 0.0142 0.0189 0.0047 0.0047 0.0048 0.184 0.1557 0.1368 0.033 0.0142 0.0094 0.0047	164 165 166 167 168 169 170 171 172 173 177 181 183	0.1557 0.1698 0.1179 0.1274 0.0094 0.0142 0.0377 0.3255 0.0189 0.0094 0.0047 0.0047	392 409 413 417 418 420 421 425 426 429 430 431 433 435 437 439 440	0.0047 0.0047 0.0094 0.0802 0.0047 0.0047 0.0047 0.2311 0.0142 0.2358 0.0047 0.2217 0.0189 0.0283 0.066 0.0472 0.0047 0.0047	201 202 203 204 205 206 210 211 213 214 215 216 217 218 219 221 222 226 230 231 277	0.0833 0.0476 0.0893 0.0179 0.0476 0.0119 0.1607 0.006 0.119 0.0417 0.0238 0.006 0.0357 0.006 0.0238 0.006 0.0238 0.0714 0.0119
HBB55			BB5655	HBB5820		HBB5859		D11S1338	
Allele 330 338 339 340 341 342 343 344 345 346 347 348 349 350 351 352 353 354 357 360	Frequency 0.0047 0.0047 0.0047 0.0083 0.0094 0.0425 0.1368 0.0708 0.0425 0.0472 0.1415 0.0142 0.2217 0.0047 0.0142 0.1179 0.0377 0.0377	Allele 283 284 285 286 287 288 289 290 291 292 293 294 315 316 318 325	Frequency 0.0123 0.0988 0.0988 0.1049 0.0556 0.0123 0.0679 0.179 0.1543 0.0247 0.037 0.0247 0.037 0.0244 0.017 0.0864 0.0617 0.0062	Allele 294 313 315 316 317 318 319 320 321 323 324 325 326 327 328 329 330 331 333	Frequency 0.0047 0.0047 0.0047 0.0991 0.2877 0.0472 0.0472 0.0094 0.0189 0.0047 0.0472 0.0044 0.0277 0.0472 0.0044 0.0283 0.0094 0.0283	Allele 374 383 384 387 388 391 392 395 396 397 400	Frequency 0.0047 0.283 0.1604 0.0755 0.0472 0.0991 0.1226 0.0236 0.1274 0.0094 0.0472	Allele 142 143 144 145 146 147 149 150 151 152 153 154 156	y 0.0047 0.1651 0.1274 0.0142 0.0236 0.0047 0.033 0.217 0.0377 0.2311 0.0943 0.033

In general, the study had established a high polymorphic STR panel and relatively close with HBB gene. The panel should be potential to perform PGT for β -thalassemia

Table 5: Observed Heterozygosity, Expected Heterozygous and Polymorphic Information Content values of 15 STRs

Order	Markers	Heterozygous	Total	H _o	He	PIC
1	HBB4506	84	106	0.79	0.883	0.872
2	D11S988	87	106	0.82	0.87	0.86
3	HBB4677	90	106	0.85	0.896	0.888
4	D11S2362	88	105	0.84	0.865	0.851
5	HBB5089	75	106	0.71	0.859	0.847
6	D11S1243	86	106	0.81	0.866	0.853
7	HBB5138	59	106	0.56	0.811	0.789
8	HBB5178	67	106	0.63	0.809	0.785
9	HBB5205	91	106	0:86	0.827	0.805
10	D11S1760	53	84	0.63	0.919	0.914
11	HBB5576	83	106	0.78	0.883	0.873
12	HBB5655	43	81	0.53	0.898	0.89
13	HBB5820	70	106	0.66	0.858	0.846
14	HBB5859	85	106	0.80	0.842	0.825
15	D11S1338	68	106	0.64	0.843	0.824

Discussion

For many years, despite medical advances, the treatment of a $\beta\text{-Thalassemia}$ patient still confronts many obstacles because it is impossible to completely treat the disease. Along with the treatment of individuals with $\beta\text{-Thalassemia},$ the application of PGT-M to help at-risk couples of having healthy babies is an extensively studied field.

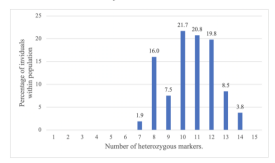


Figure 4: Percentage of individuals for number of heterozygous markers

Direct techniques have been commonly applied in diagnosis. However, for a wide gene such as HBB with a various kind of mutations, direct methods may be ineffective when using Multiplex PCR to simultaneously amplify many mutation sequences, and indirect techniques applying microsatellite markers should be the only option for this situation.

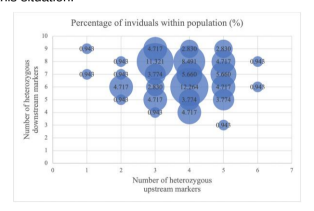


Figure 5: Percentage of individuals heterozygous for different numbers of upstream and downstream flanking microsatellite markers (n = 106)

The application of both direct and indirect techniques will increase the accuracy of the diagnosis. Therefore, linked markers play an essential role in PGT-M β -Thalassemia disease. There have been many studies of microsatellite markers [10], [11], [12], [13], [14], [15], [16], [17], but the number of STRs is still limited, so the detection of new STRs and surveys on different populations are crucial.

In this study, we developed a linked-marker set consisting of 15 STRs used for PGT-M of Vietnamese β-Thalassemia patients. Fifteen published

STR located < 1 Mb distance from HBB were successfully amplified in one PCR amplification on Vietnamese at-risk individuals with high results in Ho, He and PIC estimation. We observed that 100% individuals were heterozygous for at least 7 of the 15 markers and at least 2 of these heterozygous markers were on either side of HBB gene, reliably proving the high polymorphism and informativeness of them in most if not all Vietnamese β -Thalassemia cases.

The studied STRs panel was first established and researched on five populations by Cheng [9]. Our data on Vietnamese cases were consistent with their result and replicated the previous result in the Vietnamese Population Beside the PCR failure happening with D11S1760 (84/106) and HBB5655 (81/106), other STRs were successfully amplified with no contamination and ADO phenomenons. Most previous PGT-M on β-Thalassemia cases included nested-multiplex PCR, however by combining 15 microsatellite markers into a standard multiplex PCR reaction, nested PCR should be removed. For the lack published STRs on Vietnamese available population, establishment of STR set that compatible with Vietnamese population contributed great support to decline the high rate of Vietnamese β-Thalassemia carriers.

STRs contained different polymorphism and informativeness, but all showed the high values of PIC, Ho, He proving their clinical application. Our method had high sensitivity and specificity along with sufficient contamination and ADO monitoring. Microsatellite markers became a reliable tool when using alone in indirect mutation detection or conjunction with direct mutation detection for more precise diagnosis. Besides, this method required highly trained staffs, accurate instrument, and optimal procedures. Therefore, a particular training program needs carrying out before performing this method.

In addition to the previous study from different populations, the STRs panel continuously showed tremendous potential when applying on various kinds of population. Thus, we suggest applying this STRs panel for PGT in clinical cases.

In conclusion, All 15 STR markers were polymorphic and informative with high Ho, He and PIC. At least 7 of 15 markers were informative for 106 studied individuals, and all were observed to be heterozygous for at least two markers on either side of the HBB gene. Thus, these STRs markerhave significant meaning when applied in PGT-M widely.

Authors' contributions

DTT, NVNM, DPN, HVL, DQ, TNA, TTS, NTT, NTTH, DTPA, NLT, HTH, LH and NDB designed and performed experiments, and collected data and

informed consents. DTT, NVNM, DPN, HVL, DQ, TNA, TTS, NTT, NTTH, DTPA, NLT, HTH, LH, NDB, VTN and DTC analyzed and interpreted the results, and edited and corrected the manuscript. DTT, NVNM, VTN and DTC wrote the manuscript. All authors approved the final manuscript.

Ethical approval

This study is approved by the ethics committee of the Tam Anh General Hospital on 21 April 2018 following the Decision No 59/QĐ/BĐKTA by the director of the Tam Anh General Hospital about the establishment of the ethical committee.

Informed consent

The informed consents were signed by patients and their male partners

References

- Higgs DR, Engel JD, Stamatoyannopoulos G. Thalassaemia.
 The lancet. 2012; 379(9813):373-383.
 https://doi.org/10.1016/S0140-6736(11)60283-3
- 2. Giardine B, et al. Updates of the HbVar database of human hemoglobin variants and thalassemia mutations. Nucleic Acids Res. 2013; 42(D1):D1063-D1069. https://doi.org/10.1093/nar/gkt911 PMid:24137000 PMCid:PMC3964999
- 3. Khanh N. The frequency of hemoglobinopathies in Vietnam. Vietnamese J Med. 1993; 8(174):11-16.
- 4. Fucharoen S, Winichagoon P. Hemoglobinopathies in Southeast Asia: molecular biology and clinical medicine. Hemoglobin. 1997; 21(4):299-319. https://doi.org/10.3109/03630269709000664 PMid:9255610
- 5. Bowman JE, editor. Distribution and Evolution of Hemoglobin and Globin Loci: Proceedings of the Fourth Annual Comprehensive Sickle Cell Center Symposium on the Distribution and Evolution of

- Hemoglobin and Globin Loci at the University of Chicago, Chicago, Illinois, USA, October 10-12, 1982. Elsevier Publishing Company; 1983
- 6. Yap C, et al. First successful preimplantation genetic diagnosis in Singapore-avoidance of beta-thalassaemia major. Ann Acad Med Singapore. 2009; 38(8):720-3.
- 7. Fan L, et al. Genetic diagnosis of beta-thalassemia preimplantation using short tandem repeats in human cryopreserved blastocysts. International Journal of Clinical and Experimental Pathology. 2017; 10(7):7586-7595.
- 8. Machado F, Medina-Acosta E. High-resolution combined linkage physical map of short tandem repeat loci on human chromosome band Xq28 for indirect haemophilia A carrier detection. Haemophilia. 2009; 15(1):297-308. https://doi.org/10.1111/j.1365-2516.2008.01866.x PMid:18752533
- 9. Chen M, et al. Identification of novel microsatellite markers< 1 Mb from the HBB gene and development of a single-tube pentadecaplex PCR panel of highly polymorphic markers for preimplantation genetic diagnosis of beta-thalassemia. Electrophoresis. 2015; 36(23):2914-2924. https://doi.org/10.1002/elps.201500146 PMid:26331357
- 10. Rechitsky S, et al. Reliability of preimplantation diagnosis for single gene disorders. Molecular Cellular Endocrinology. 2001; 183:S65-S68. https://doi.org/10.1016/S0303-7207(01)00576-7
- 11. Rechitsky S, et al. Preimplantation genetic diagnosis with HLA matching. Reproductive BioMedicine Online. 2004; 9(2):210-221. https://doi.org/10.1016/S1472-6483(10)62132-3
- 12. Fiorentino F, et al. Short tandem repeats haplotyping of the HLA region in preimplantation HLA matching. European Journal of Human Genetics. 2005; 13(8):953. https://doi.org/10.1038/si.eihg.5201435 PMid:15886713
- 13. Kuliev A, et al. Preimplantation diagnosis and HLA typing for haemoglobin disorders. Reproductive BioMedicine Online. 2005; 11(3):362-370. https://doi.org/10.1016/S1472-6483(10)60845-0
- 14. Fiorentino F, et al. Strategies and clinical outcome of 250 cycles of preimplantation genetic diagnosis for single gene disorders. Human Reproduction., 2005; 21(3):670-684. https://doi.org/10.1093/humrep/dei382 PMid:16311287
- 15. Kuliev A, et al. Preimplantation genetic diagnosis for hemoglobinopathies. Hemoglobin. 2011; 35(5-6):547-555. https://doi.org/10.3109/03630269.2011.608457 PMid:21910603
- 16. Zachaki S, et al. Novel and known microsatellite markers within the β -globin cluster to support robust preimplantation genetic diagnosis of β -thalassemia and sickle cell syndromes. Hemoglobin. 2011; 35(1):56-66. https://doi.org/10.3109/03630269.2010.544620 PMid:21250882
- 17. Durmaz B, et al. Genotyping of β -Globin Gene Mutations in Single Lymphocytes: A Preliminary Study for Preimplantation Genetic Diagnosis of Monogenic Disorders. Hemoglobin. 2012; 36(3):230-243. https://doi.org/10.3109/03630269.2012.675891 PMid:22524255