Correlation between thyroglobulin gene polymorphisms and autoimmune thyroid disease

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Abstract. The aim of the present study was to detect thyroglobulin (Tg) gene polymorphisms in a Han Chinese population from the Northern regions of Henan province, China, and to study the correlation between Tg gene polymorphisms and autoimmune thyroid disease (AITD). A total of 270 patients with AITD and 135 healthy controls were enrolled. Genomic DNA was extracted and fluorescence polymerase chain reaction analysis was performed; high-resolution melting curve analysis (HRMA) was used to detect single-nucleotide polymorphisms (SNPs) in exons 10, 12 and 33 of the Tg gene. SNPs were then correlated with AITD. Han people from the Northern regions of Henan displayed four Tg exon SNPs: E10SNP24 T/G, E10SNP158 T/C, E12SNP A/G and E33SNP C/T. Several allele and genotype frequencies differed between the AITD group and the healthy control group (Tg E10SNP: Allele T, P<0.01; allele G, P<0.01; and Tg genotype GG, P<0.01; genotype TG, P<0.01. Tg E12SNP: Allele A, P<0.01; allele G, P<0.01; Tg genotype GG, P<0.01; genotype AG, P<0.01). A statistically significant difference in the frequency of selected Tg SNPs haplotypes was also present between AITD patients and healthy controls (P<0.05). There was no significant difference in haplotypes between various types of AITD (hypothyroidism, hyperthyroidism and Hashimoto's disease). The Tg SNP frequency distribution was significantly different between Han populations of the Northern regions of Henan province and the Xi'an regions of Shaanxi province. The results of the present study suggested that specific Tg gene alleles or genotypes were correlated with AITD; specific Tg SNP haplotypes were associated with hypothyroidism, hyperthyroidism and Hashimoto's disease, and the Tg SNP frequency distribution differed depending on the geographical location of the Han Chinese populations.

Introduction

Autoimmune thyroid disease (AITD) encompasses a range of thyroid conditions, including hyperthyroidism (Graves' disease), Hashimoto's disease (Hashimoto's thyroiditis), idiopathic myxedema and primary hypothyroidism. Genetic factors have an important role in the occurrence of AITD (1). Genetic screening has shown that susceptibility genes for AITD include human leukocyte antigen and the cytotoxic T-lymphocyte-associated antigen-4 (2). Although AITD has been associated with immunomodulatory genes, this does not fully explain the specific autoimmune attack directed towards the thyroid. Therefore, disease-associated antigen-specific genes, including those encoding thyroid peroxidase, thyroid-stimulating hormone receptor and thyroglobulin (Tg) comprise another group of candidate genes which are potentially associated with AITD (2). To date, only few studies on Tg and genes associated with the susceptibility to AITD have been performed, and the findings from different population are not consistent (3-5), possibly due to a range of different detection methods used. Commonly used methods for analyzing differences in gene sequences include single-strand conformation polymorphism (6), denaturing gradient gel electrophoresis (7), denaturing high-performance liquid chromatography (8), temperature gradient capillary electrophoresis (9) and mass spectrometry (10). These methods require separate samples, enzymatic reactions and chemical reactions; these methods are complex, comprise multiple steps, and are time-consuming and challenging to perform. However, the high-resolution melting curve analysis (HRMA) method is relatively simple, fast, accurate, economical and suitable for high-throughput analysis (11-14). HRMA has previously been used to detect variable-number
tandem repeats and displayed consistency with traditional gel electrophoresis (21), a technique which is widely used in genetic testing. To utilize this technique, the present study adopted a protocol for the preparation and use of reagents to extract DNA (22). In the present study, HRMA amplification of small fragments was employed to study Tg gene polymorphisms of a Han Chinese population from the Northern regions of Henan province and examined the correlation between Tg gene polymorphisms and AITD.

Materials and methods

Clinical data. A total of 270 AITD patients (180 females, 90 males) were enrolled at the Department of Nuclear Medicine, Anyang Regional Hospital (Anyang, China) between February 2013 and January 2014. The average age of female and male AITD patients was 37.72±14.30 and 38.04±12.77 years, respectively. A control group comprising 135 healthy individuals (90 females and 45 males) was recruited at Anyang Regional Hospital Medical Center (Anyang, China). The average age of female and male control subjects was 35.68±11.67 and 38.16±11.75 years, respectively. The inclusion criteria were as follows: The diagnosis of AITD was based on typical clinical symptoms, and was confirmed by laboratory tests (90 cases of primary hypothyroidism, 90 cases of Graves’ disease and 90 cases of Hashimoto disease) (23). The control group was age- and gender-matched with the AITD group. The study was approved by the Ethics Committee of Anyang Regional Hospital of Puyang (Puyang, China). Written informed consent was obtained from the patient.

Genomic DNA extraction from blood. Genomic DNA extraction was performed according to the method of Wang et al (22). First, 0.6 ml extraction buffer (guanidine thiocyanate, Tris-HCl, EDTA. Triton X-100) was placed in a 1.5-ml centrifuge tube and 200 µl blood sample was added; the mixture was incubated for 5-10 min at room temperature with continuous gentle mixing to avoid shock and DNA shearing. The mixture was then centrifuged at 13,000 x g for 10 min at room temperature. The supernatant (0.7 ml) was loaded onto a pre-prepared adsorption column (Beijing Solarbio Science & Technology Co., Ltd., Beijing, China), which was placed in a 1.5-ml collection tube. The column was then centrifuged at 12,000 x g for 30 sec at room temperature and the eluate was discarded. Subsequently 500 µl binding buffer was added to the adsorption column, followed by centrifugation at 12,000 x g for 30 sec at room temperature. The eluent was discarded and 600 µl wash buffer (70% ethanol supplemented with 10 mM NaCl) was added to the adsorption column, followed by centrifugation at 12,000 x g for 1 min at room temperature and discarding of the eluent. The washing step was repeated once and the adsorption column was transferred to a fresh 1.5-ml microcentrifuge tube. The column was then centrifuged at 12,000 x g for 30 sec at room temperature. The eluent was discarded and 600 µl wash buffer (70% ethanol supplemented with 10 mM NaCl) was added to the adsorption column, followed by centrifugation at 12,000 x g for 1 min at room temperature and discarding of the eluent. This washing step was repeated once and the adsorption column was transferred to a fresh 1.5-ml microcentrifuge tube. 50 µl Tris-EDTA buffer was added to the column followed by incubation for 1 min at room temperature. The column was then centrifuged at 12,000 x g for 1 min at room temperature, and the eluate containing the DNA was collected and stored at -20°C.

Primer design. As opposed to ordinary PCR primers, HRMA requires the design of primers for the synthesis of PCR products of <150 bp. PCR products were sequenced using separate sequencing primers; the amplified product of PCR using the sequencing primers was >300 bp. Primers were designed according to the published National center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/) gene sequence for the Tg gene. HRMA primers and sequencing primers for the Tg gene are listed in Table I. Primers were synthesized by Sangon Biotech (Shanghai, China).

Table I. PCR primers used in the present study.

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence</th>
<th>PCR product length (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HRM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E10SNP24</td>
<td>F: 5'-CACCTGCTCATTGTTCCTCC-3'</td>
<td>80</td>
</tr>
<tr>
<td></td>
<td>R: 5'-GGCCTGACCGTCTGAG-3'</td>
<td></td>
</tr>
<tr>
<td>E10SNP158</td>
<td>F: 5'-CTCATCCCAAGTGCGAGC-3'</td>
<td>87</td>
</tr>
<tr>
<td></td>
<td>R: 5'-CGTTGTAACCTCAGAACGAC-3'</td>
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<tr>
<td>E12SNP</td>
<td>F: 5'-CGACTGCGTGGAGCACTC-3'</td>
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<td>R: 5'-GCCGGACCTGACAGG-3'</td>
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<tr>
<td>E33SNP</td>
<td>F: 5'-ACCTTCAGGCTGTCCTTTCC-3'</td>
<td>80</td>
</tr>
<tr>
<td></td>
<td>R: 5'-AGTGCAGCATGGGTCGCCG-3'</td>
<td></td>
</tr>
<tr>
<td>Sequencing</td>
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<td></td>
</tr>
<tr>
<td>E10SNP</td>
<td>F: 5'-GCTGATCACCAACTGATGTGTC-3'</td>
<td>481</td>
</tr>
<tr>
<td></td>
<td>R: 5'-CTGAAGTTTCCCGGAAGCCTG-3'</td>
<td></td>
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<tr>
<td>E12SNP</td>
<td>F: 5'-CATACGTGTTTGTTTCTCAGC-3'</td>
<td>389</td>
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<tr>
<td></td>
<td>R: 5'-GCTCCTGTTCACAAAGTGAAG-3'</td>
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<td>F: 5'-GGACAGTATTCTCGAGAGAG-3'</td>
<td>403</td>
</tr>
<tr>
<td></td>
<td>R: 5'-GAACGAGGATAGGAATGCTG-3'</td>
<td></td>
</tr>
</tbody>
</table>

PCR, polymerase chain reaction; HRM, high-resolution melting curve assay; SNP, single nucleotide polymorphism; E10, exon 10; F, forward; R, reverse.
Quantitative PCR conditions and HRMA analysis. Each 20-µl reaction mixture contained 1 µl template, 1 µl forward primer, 1 µl reverse primer, 10 µl 10X HS TaqE mix (KangWei Century Biotechnology Company, Beijing, China), 1 µl 20X EvaGreen (Biotium, Hayward, CA, USA), and 6 µl sterile distilled water. Once mixed, the PCR was performed using a LightCycler 480 RT-PCR machine (Roche Diagnostics, Basel, Switzerland) for fluorescence quantification. The reaction procedure was performed using the following thermocycling conditions: 95°C for 10 min, followed by 40 cycles of 95°C for 10 sec and 60°C for 15 sec, during which fluorescence measurements were collected; then 95°C for 10 sec and 65°C for 15 sec; and finally, the temperature was increased from 60 to 95°C in 1°C-steps. A total of 50 fluorescence measurements were collected to obtain the melting curve. After the PCR and high-resolution melting curves were completed, GeneScanning software (Roche Diagnostics) on the LightCycler instrument was used for analysis of high-resolution melting curves. Through software analysis, DNA samples were assigned to three categories: Mutant homozygous, wild-type homozygous or heterozygous. In each category, the PCR products of selected samples were completely sequenced by Sangon Biotech. The gel electrophoresis was performed on 1% agarose gels (Biowest Agarose; Gene Tech Company, Hong Kong, China). The marker DL2000 (Takara Bio, Inc., Otsu, Japan) was used as the molecular standard.

Statistical analysis. After data from individual samples had been recorded, genotype, allele and haplotype frequencies were calculated. SPSS 16.0 software (SPSS, Chicago, IL, USA) was used for statistical analysis. P-values, odds ratios (OR) and 95% confidence intervals (95% CI) were calculated using the χ² test.

Results

PCR products. Following PCR, the amplified PCR products were analyzed by gel electrophoresis. The PCR product of E10SNP24 was 80 bp in length, the PCR product of E10SNP158 was 87 bp, the PCR product of E12SNP was 93 bp and the PCR product of E33SNP was 80 bp in length. The electrophoresis results showed bands of the expected sizes (Fig. 1).

HRMA. After the PCR was completed, PCR products were analyzed using HRMA. The EvaGreen fluorescent conjugate only labels double-stranded DNA; therefore, as the temperature increased and the DNA gradually unraveled into single strands during the recording of the melting curve, the fluorescence intensity of the mixture was gradually reduced. The DNA sequences of the different genotypes contained different combinations of nucleotide bases, generating PCR products with different melting points and melting curves. The melting curves from different genotypes, which were homozygous for the wild-type allele, while being heterozygous or homozygous for the mutant allele, are represented in graphical form by different colors (Fig. 2). Therefore, the HRMA was able to distinguish between different genotypes; these genotypes were confirmed by DNA sequencing. As an example, the HRMA results for the exon 10 polymorphism, (E10SNP158 T/C), are shown in Fig. 2; other exon genotype maps were similar to that of the exon 10 SNP158 T/C.

Sequencing results. The HRMA graphs were evaluated via their colored curves displayed in red, blue and green, with each color representing a class of sample genotype. One genotype from each category of samples was selected to be sequenced. Every polymorphic locus of four polymorphic loci had three color curves, and 12 sample curve charts were produced. The PCR products of 12 representative samples were randomly selected to be sequenced, encompassing three exons and four polymorphism loci. The sequencing results provided information regarding the 12 genotypes, consisting of four polymorphic loci which each containing wild-type homozygotes, mutant homozygotes and heterozygotes. Through Tg sequencing, four SNPs were identified: In exon 10, the E10SNP24 T/G allele (GG, TG, and TT genotypes) and the E10SNP158 T/C allele (CC, TC and TT genotypes); in exon 12, the E12SNP A/G allele (AA, AG and GG genotypes); and in exon 33, the E33SNP C/T allele (CC, CT and TT genotypes). Representative sequence diagrams for the E33SNP polymorphism (CC, CT or TT) sequenced diagram are shown in Fig. 3A-C.

Statistical results. The frequency distribution of the E10SNP24 (allele T/G, TG and TT genotypes) and the E12SNP (allele A/G, AG and GG genotypes), was significantly different between patients with AITD and healthy controls (P<0.05). The OR values of these genotypes showed a strong correlation with the disease. The E10SNP24 GG and TG genotypes, with OR=0.14 for the GG genotype and OR=11.5 for the TG genotype, and the E12SNP GG and AG genotypes, with OR=4.22 for the GG genotype and OR=0.23 for the AG genotype, were highly correlated with the disease. There were no statistically significant differences in the frequency of E10SNP158 and E33SNP polymorphisms when comparing healthy controls.
Figure 2. E10SNP158T/C HRMA. E10SNP158T/C includes the polymorphism, its dissolution points are different, therefore the dissolution curves are different. The blue curve indicates the E10SNP158CC genotype, the red curve indicates the E10SNP158TC genotype and the green curve indicates the E10SNP158TT genotype. (A) Thumbnail and (B) enlarged drawing of E10SNP158 HRMA segmentation. SNP, single nucleotide polymorphism; E10, exon 10; HRMA, high-resolution melting curve analysis.

Figure 3. Sequencing analysis. The HRMA graphs were evaluated via their colored curves displayed in red, blue and green, with each color representing a class of sample genotype. One genotype from each category of samples was selected to be sequenced. Every polymorphic locus had three color curves. (A) E33SNP CC genotype sequencing maps and polymorphic loci. Exon 33 (20 bases) is a polymorphic locus. The arrow indicates a single peak which was proved to be the homozygous CC genotype. (B) E33SNP TC genotype sequencing maps and polymorphic loci. Exon 33 (20 bases) is a polymorphic locus. The arrow indicates a double peak, which was proved to be the heterozygous TC genotype. (C) E33SNP TT genotype sequencing maps and polymorphic loci. Exon 33 (20 bases) is a polymorphic locus, and the arrow indicates a single peak, which was proved to be the homozygous TT genotype. SNP, single nucleotide polymorphism; E33, exon 33.
with AITD patients (P>0.05) (Table II). Furthermore, a haplotype analysis of the four Tg polymorphic loci, E10SNP24, E10SNP158, E12SNP and E33SNP, was performed. There were seven haplotype combinations of gene polymorphisms that occurred more frequently in AITD patients than in healthy controls, and among these, five haplotype combinations were significantly different (Table III).

The haplotype combination of gene polymorphisms was not significantly different between the different disease sub-groups (P>0.05). Certain gene polymorphism combinations were most likely to occur more frequently amongst AITD patient groups: Disease-linked haplotypes (denoted as E10SNP24 - E10SNP158 - E12SNP - E33SNP) were present in the primary hypothyroidism group (G-T-A-C, T-T-G-C and T-C-G-C), the hyperthyroidism group (G-T-A-C, G-T-G-C and T-C-G-C) and the Hashimoto’s disease group (G-C-G-C, G-T-A-C, G-T-G-C and T-C-G-C); compared with those in the healthy controls, these haplotypes were significantly different.

Table III. Tg haplotype gene SNP portfolio analysis of the frequency between the disease and healthy control groups [n (%)].

<table>
<thead>
<tr>
<th>E10SNP24</th>
<th>E10SNP158</th>
<th>E12SNP</th>
<th>E33SNP</th>
</tr>
</thead>
<tbody>
<tr>
<td>T</td>
<td>T</td>
<td>C</td>
<td>C</td>
</tr>
<tr>
<td>G</td>
<td>G</td>
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</tr>
<tr>
<td>TG</td>
<td>CC</td>
<td>CT</td>
<td>TT</td>
</tr>
</tbody>
</table>

Analysis of haplloid allele combinations were compared between the disease group and the healthy group. *P<0.01, †P<0.05, disease group vs healthy control group. Hypothyroidism (G-T-A-C, OR=0.22; T-T-G-C, OR=4.00; T-C-G-C, OR=5.23); Graves (G-T-A-C, OR=0.22; G-T-G-C, OR=3.57; T-C-G-C, OR=3.00); HT (G-C-G-C, OR=0.57; G-T-A-C, OR=0.24; G-T-G-C, OR=3.50; G-T-G-C, OR=3.59; T-C-G-C, OR=4.45). HT, haplotype; Tg, thyroglobulin; OR, odds ratio; SNP, single nucleotide polymorphism; AITD autoimmune thyroid disease; E10, exon 10.
The present study found four alleles in the Tg gene in a Han Chinese population, as well as twelve associated SNPs and mutations. The HRMA method uses fluorescent probes to examine the differing dissolution profiles of PCR products of different genotypes, based on the different base pairing in homozygous and heterozygous sequences, leading to different melting curves from low to high temperatures, thereby easily distinguishing alleles by detecting a change in the fluorescence signal (15).

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References