Analysis of DAZL SNP260 and SNP386 in infertile Chinese males using multi-analyte suspension array

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Abstract. The aim of the present study was to investigate the association between two single nucleotide polymorphisms (SNPs) and infertility in Chinese males using multi-analyte suspension array (MASA). A total of 196 male patients with azoospermia or severe oligospermia (sperm density <5x10⁶/ml, non-obstructed) who had a normal karyotype and no azoospermia factor microdeletions were recruited, along with 40 healthy, fertile males as controls. Two SNPs of the deleted in azoospermia-like (DAZL) gene, SNP260 and SNP386, were genotyped by allele-specific primer extension (ASPE) combined with MASA technology. The SNP260A>G and SNP386A>G mutations were found in the males with infertility. The SNP260, but not the SNP386, mutation was detectable in the control group. The mutation rates in the controls and patients were 2.5 and 3.06% for SNP260, and 0 and 2.04% for SNP386, respectively. A χ² analysis did not identify any significant differences in the frequency of either mutation between the fertile and infertile males. In conclusion, the combination of ASPE and MASA methods for SNP genotyping was high-throughput, accurate and cost-efficient. The method was applied to detect SNP polymorphisms in the DAZL gene; and neither the A260G nor the A386G polymorphism of DAZL appeared to be involved in male infertility in the Chinese population.

Introduction

Male infertility is considered to be associated with azoospermia factor (AZF), one of several genes located in the Yq11 chromosomal region. The most common mutations identified in AZF are microdeletions in AZFc (60%) (1,2). The members of the deleted in azoospermia (DAZ) gene family are the primary regulators of proliferation in early germ cells and are important candidate genes for male infertility in the AZFc region (3). There are three genes in the DAZ gene family: DAZ, DAZL and BOULE. The DAZ-like (DAZL) gene is located in the 3p24 chromosomal region and is an autosomal homolog of DAZ; 83% of the cDNA coding regions in DAZ and DAZL are similar (4-6).

A previous study suggested that DAZL is important in the sperm production process (7). However, there are conflicting studies regarding whether DAZL mutations impact male infertility. Following completion of the human genome project, single nucleotide polymorphisms (SNPs) have been intensively investigated. As genetic markers, SNPs have been associated with pharmacogenomics and disease susceptibility. Two studies (8,9) have described A>G transitions at the SNP260 and SNP386 nucleotide positions of DAZL, and SNP386 was found to be associated with susceptibility to spermatogenic failure in the Taiwanese population. However, studies in other countries, including Italy, India and Japan, have indicated that there is not an association between the two SNPs and spermatogenic impairment (10-13). Thus, the role of the SNP260A>G and SNP386A>G transitions in male infertility is controversial (12-16). To the best of our knowledge, there have been no studies regarding DAZL SNPs in the Chinese population. Therefore, in the present study, the distribution of the DAZL A260G and A386G SNPs in Chinese males was investigated.

The methods for SNP genotyping are advancing. The available technology includes: Restriction fragment length polymorphism (RFLP), Taqman, high performance liquid chromatography and single-strand conformation polymorphism analysis (17). The most common methods used currently in China to detect SNPs are RFLP and DNA sequencing; however, these techniques are limited in their applications due to the high costs in time and resources.

In the present study, a high-throughput, low-cost and low-consumption method that combined allele-specific primer extension (ASPE) and multi-analyte suspension array (MASA)
technology was applied (18) to characterize the DAZL SNP260 and SNP386 mutations in a Chinese population sample.

Materials and methods

Ethics statement. This study was approved by: The Ethics Committee of Chongqing Institute of Science and Technology for Population and Family Planning, the Research Ethics Committees of the Key Laboratory of Birth Defects and Reproductive Health, and the Ethics Committee of the Human Sperm Bank of Chongqing (Chongqing, China). All ethics approvals were provided in compliance with the Declaration of Helsinki (World Medical Association, 2000). All participants included in this study provided informed consent.

Patients and sample collection. A total of 196 males with azoospermia (non-obstructed) or severe oligospermia (sperm density <5x10^6/ml), but with a normal karyotype and no AZF microdeletions, were recruited from the Affiliate Hospital of Sichuan Genitalia Hygiene Research Center (Chengdu, China). Forty samples from fertile controls comprised of healthy volunteers were collected from the Human Sperm Bank of Chongqing. Whole blood (5 ml) was obtained from all patients and control subjects. Peripheral blood lymphocytes were isolated from the whole blood and stored at -20˚C until required for analysis.

Coupling of oligonucleotides to microspheres. The sequences of cZipcodes for SNP260 and SNP386 (Table I) were selected online through Luminex Corporation (Austin, TX, USA). The 5' ends of all cZipcodes were C-12-linked and amino-modified. To couple the cZipcodes to carboxylate beads, 2.5 million carboxylate beads, suspended in 25µl 0.1 mol/l 2-(N-morpholino)ethanesulfonic acid (pH 4.5), were mixed with 0.5 nmol amino-modified cZipcodes. Subsequently, 1.25 µl 10 mg/ml fresh 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride was added to the microsphere/oligo mixture and incubated in the dark for 30 min at room temperature, and this step was repeated twice. The microsphere/oligo mixture was mixed occasionally during incubation to ensure that the microspheres remained in suspension. The bead mixture was washed with 0.5 ml 0.02% Tween-20 and 0.5 ml 0.1% SDS. The beads were finally resuspended in 50 µl Tris-EDTA (pH 8.0) at 4˚C in the dark.

DNA isolation and polymerase chain reaction (PCR). Genomic DNA was extracted from peripheral blood lymphocytes using a DNA isolation kit (Youjing Corp., Seoul, Korea). PCR was performed in a total volume of 25 µl containing 150 ng genomic DNA, 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 1.5 mM MgCl_2, 200 µM dNTPs, 5 pmol of each primer (Table II) and 1.5 U DNA polymerase. The PCR cycling conditions were as follows: 94˚C for 2 min, followed by 30 cycles of 94˚C for 30 sec, 55˚C for 1 min and 72˚C for 2 min.

ASPE. To remove unincorporated dNTPs and primers from the PCR reaction, 2 U Shrimp alkaline phosphatase and 1 U Exonuclease I were added to 10 µl of the pooled PCR products. The mixture was incubated at 37˚C for 15 min, and then the enzymes were inactivated by heating at 95˚C for 20 min. The ASPE reaction comprised 10 µl enzyme-treated PCR products, 40 mM Tris-HCl (pH 8.4), 100 mM KCl, 25 mM MgCl_2, 0.5 pmol of each SNP oligonucleotide, 100 µM of each dNTP (including biotin-labeled dCTP) and 3 U Tsp DNA polymerase in a total volume of 20 µl. The PCR cycling conditions were as follows: 96˚C for 2 min, followed by 30 cycles of 94˚C for 30 sec, 55˚C for 1 min and 72˚C for 2 min.

Hybridization of ASPE products to the microspheres. A total of 3,000 probe-coupled microspheres of each type were added

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**Table I. Sequences of Capture probes, ZipCodes and cZipCodes (5’>3’).**

<table>
<thead>
<tr>
<th>SNP</th>
<th>Capture probe</th>
<th>ZipCode</th>
<th>cZipCode</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNP260A</td>
<td>CTGGCCTCTCTGGAGATGGT</td>
<td>CTACAAACAAAACATAATTACAA</td>
<td>TGATAATGTTTGTGGTTTTGTAG</td>
</tr>
<tr>
<td>SNP260G</td>
<td>CTGGCCTCTCTGGAGATGGC</td>
<td>CTTTAATCTTATACATTTATCA</td>
<td>TGATAAAGTAGGATTTAAAAGG</td>
</tr>
<tr>
<td>SNP386A</td>
<td>GCAAAGAAGCTCTAATCTTCTCAGT</td>
<td>TCAACAATCTTTTTTACATCAAATC</td>
<td>ATTGATTGAAAGAGTTGTTGGA</td>
</tr>
<tr>
<td>SNP386G</td>
<td>GCAAAGAAGCTCTAATCTTCTCAGG</td>
<td>TCAATCATTACACTTTTTCAACAAT</td>
<td>ATTTGTTAAAGAGTTGTTGGA</td>
</tr>
</tbody>
</table>

SNP, single nucleotide polymorphism.

**Table II. Primer sequences (5’>3) and the sizes of the PCR amplification products.**

<table>
<thead>
<tr>
<th>PCR product</th>
<th>Primer sequence</th>
<th>Product length (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNP260A Forward</td>
<td>CCTGAGCCTGAACTAACCTTAGAATG</td>
<td>225</td>
</tr>
<tr>
<td>SNP260A Reverse</td>
<td>AATATACCTTGCTGTTGTC</td>
<td></td>
</tr>
<tr>
<td>SNP386 Forward</td>
<td>GGGAGAAATTGTACATCATCG</td>
<td>198</td>
</tr>
<tr>
<td>SNP386 Reverse</td>
<td>AAAATTACTCACCCCTTTGACAC</td>
<td></td>
</tr>
</tbody>
</table>

PCR, polymerase chain reaction; SNP, single nucleotide polymorphism; bp, base pairs.
to the ASPE products and 1X tetramethylammonium chloride was used to obtain a final volume of 50 µl. The mixture was heated at 90˚C for 10 min and then hybridized at 53˚C for 5, 10, 15, 30 and 45 min to optimize the hybridization time. The hybrid products were labeled with 200 ng fresh streptavidin-R-phycoerythrin at 55˚C for 5 min. The fluorescent signal was measured using the Luminex 100 system (Luminex Corporation).

RFLP and DNA sequencing for verification of the accuracy of the ASPE. All samples identified as SNP mutants using ASPE were sequenced using the alternate methods. Samples randomly selected from the non-mutated infertile males and control patients were also used. The RFLP reaction consisted of 17.3 µl purified PCR products, 0.2 µl bovine serum albumin, 2 µl 10X PCR buffer, and 0.5 µl restriction endonuclease DdeI for SNP260 or AluI for SNP386. The total reaction volume was 20 µl. Following incubation at 37˚C for 4 h, the enzyme products were analyzed using polyacrylamide gel electrophoresis. PCR products were first cloned (TOPO TA cloning kit; Invitrogen Life Technologies, Carlsbad, CA, USA) according to the instructions of the manufacturer, followed by DNA sequencing. The identification of the obtained sequence was verified by using alignment search tool (BLAST) analysis (www.ncbi.nlm.nih.gov).

Statistical analysis. Statistical analysis was performed using SPSS software, version 13.0 (SPSS, Inc., Chicago, IL, USA). The χ² test was used to evaluate the genotype distribution and allele frequencies of the polymorphisms. P<0.05 was considered to indicate a statistically significant difference.

Results

Assay optimization. To assess the association between DAZL mutations and male infertility, the present study aimed to develop a bead-based platform for SNP genotyping of DAZL. A schematic of the ASPE technology applied is shown in Fig. 1. For the SNP genotyping, two capture probes were designed for each SNP site. One primer was complementary to the wild-type sequence and the other was complementary to the mutation. A thermostable polymerase was used to extend the capture probe by incorporation of dNTPs, and one of which was biotin-labeled (Fig. 1). Extension only occurred if the 3' nucleotide of the probes had annealed to the template DNA. A DNA sequence, termed Zipcode, was added at the 5'-end portion of the capture probe. The Zipcode hybridized to its complementary sequence, termed cZipcode, which was coupled to a specific fluorescent microsphere (Fig. 1). The sequences of the Zipcode and cZipcode pairs for each SNP are shown in Table I. With the extension of the capture probe,
the template DNA was labeled with biotin. Following hybridization and dye incorporation, the signals were captured and analyzed by Luminex 100.

To optimize the hybridization time, samples of the mixture were incubated for 5, 10, 15, 30 and 45 min. Allowing samples to hybridize for 30 min achieved the best specificity and highest fluorescence intensity (Fig. 2). For the multiplex bead array screening, a cut-off value was established based on a fluorescence intensity of 100. Results with fluorescence intensity <100 were classified as negative. Results classified as positive were at least 10-fold greater than the negative reading. The results presented had to meet the aforementioned requirements to be considered valid.

SNP260 assayed by ASPE, RFLP and DNA sequencing. In the 196 infertility patients and 40 controls, the heterozygote mutation was not identified using MASA. Only one (2.50%) of the 40 controls and six (3.06%) of the 196 patients were observed

![Image](image-url)

**Figure 3. Results of the multi-analyte suspension array, restriction fragment length polymorphism and DNA sequencing.**

(A) The fluorescence intensity of the blank, wild-type and homozygous mutations of SNP260 and SNP386. (B) Electrophoretic analysis of the PCR products of deleted in azoospermia-like digested by Ddel or AluI; lanes 1 and 6, marker; lane 2, SNP260 PCR products (225 bp); lanes 3 and 4, SNP260 PCR products with wild-type digested by Ddel (217, 3 and 5 bp; the latter two were run off the gel); lane 5, SNP260 PCR products with homozygous mutation digested by Ddel (155, 62, 5 and 3 bp; the latter three were run off the gel); lane 7, SNP386 PCR products (198 bp); lane 8, SNP386 PCR products with wild-type digested by AluI (118 and 80 bp); lane 9, SNP386 PCR products with homozygous mutation digested by AluI (105, 80 and 13 bp, the last was run off the gel). (C) Representative DNA sequence analysis of SNP260 showing the wild-type sequence. (D) Representative SNP260 DNA sequence analysis showing the homozygous mutation. (E) Representative DNA sequence analysis of SNP386 showing the wild-type sequence. (F) Representative SNP386 DNA sequence analysis showing the homozygous mutation. PCR, polymerase chain reaction.

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**Table III. Relative prevalence of mutations in DAZL SNP260 and SNP386 in infertile males and controls.**

<table>
<thead>
<tr>
<th>Mutation site</th>
<th>Infertility (n=196)</th>
<th>Control (n=40)</th>
<th>$\chi^2$</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNP260A&gt;G</td>
<td>1 (2.50)</td>
<td>0 (0.00)</td>
<td>0.18</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>SNP386A&gt;G</td>
<td>6 (3.06)</td>
<td>4 (2.04)</td>
<td>0.05</td>
<td>&gt;0.05</td>
</tr>
</tbody>
</table>

DAZL, deleted in azoospermia-like; SNP, single nucleotide polymorphism.
to have a homozygous mutation (Table III). The results of the wild-type and homozygous mutations detected by MASA are shown in Fig. 3A. The results obtained using ASPE were consistent with the results obtained using RFLP (Fig. 3B) and DNA sequencing (Fig. 3C-F).

**SNP386 assayed by ASPE, RFLP and DNA sequencing.** In the enrolled patients, no heterozygote mutations in SNP386 were detected (Table III). Four (2.04%) of the 196 infertility patients were identified to have the homozygous SNP386 mutation (Table III). All the controls were wild-type homozygous (Table III). However, χ² analysis did not indicate a statistically significant difference between the groups. The results of the wild-type and homozygous mutations detected by MASA are shown in Fig. 3A. The ASPE results were again consistent with those of the RFLP (Fig. 3B) and DNA sequencing (Fig. 3C-F).

**Association between the DAZL mutation and infertility in males.** The mutation rates of SNP260 and SNP386 in the controls and patients were low: 0.00 and 2.50% for SNP260 and 2.04 and 3.06% for SNP386, respectively. The frequencies of the SNP mutations in the control and infertile males were similar (P>0.05). This was inconsistent with results reported in a study of controls and infertile male patients from Taiwan (0.86 and 7.39%, respectively, for SNP386) (8,9).

**Discussion**

**DAZL, DAZ and BOULE** are the three members of the DAZ gene family which encode RNA-binding proteins associated with impaired spermatogenesis. The DAZ gene family contributes to spermatogenesis in a number of different species. Houston et al (19) found that DAZL knockout reduced the number of testicular stem cells in mice and caused spermatogonia to halt in meiosis. They also demonstrated that mice lacking BOULE exhibited impaired spermatogenesis, but the symptoms were corrected by reimplantation of the Xenopus xDAZL gene. Sleva et al (20) have also reported that the human DAZL gene partially restored function in mice lacking DAZL. Lin et al (21) have shown that males with Sertoli-cell-only syndrome exhibited low levels of expression of DAZL. All these studies demonstrate that the DAZL gene is important during spermatogenesis.

Spermatogenesis is a complex process regulated by several genes located on autosomal and sex chromosomes (22). Approximately 10% of infertile males have complete or partial deletions of the DAZ gene cluster (23,24). The hypothesis that DAZL is involved in spermatogenesis is supported by studies demonstrating its testis-specific expression and its high homology to DAZ (25,26). Studies have indicated that DAZL may be involved in germ cell development. For example, in Caenorhabditis elegans, inactivation of DAZL was associated with meiotic arrest in oogenesis (27). In mice, knockout of the DAZL homolog led to the loss of germ cells in males and females (28). Additionally, in humans, low mRNA transcript levels of the DAZL gene were identified in infertile males with testicular failure (29).

However, there have been few studies that establish the association between the human DAZL gene mutation and male infertility. Two studies (8,9) have identified two SNP mutations in DAZL in the Taiwanese population: SNP260A>G in exon 2 and SNP386A>G in exon 3. SNP386A>G was identified in 7.39% of infertile males compared with 0.86% of fertile males. By contrast, studies in India, Japan and Italy observed that SNP386A>G was not associated with male infertility (10-13). Notably, Bartoloni et al (10) found the SNP386 mutation in <1/316 males with azoospermia and severe oligospermia. It is possible that the mutation may only be associated with infertility in males of Asian descent. However, the present study, conducted using Chinese male participants, also revealed no association between male infertility and the SNP386 mutation. This suggests the SNP386 mutation may be associated with infertility in only the Taiwanese population. Alternatively, the mutation site may be located in non-specific sites of the gene for the RNA-binding protein, and would therefore not likely be important in the sperm production process. To further understand the association between SNP386 and male infertility, additional studies at the protein level are required.

To date, there are four methods compatible with MASA that can be used to genotype SNPs: (i) Single base chain extension (SBCE), (ii) ASPE, (iii) oligonucleotide ligation assay and (iv) direct DNA hybridization (DH) (30). Lee et al (31) compared the accuracy, efficiency and costs of these methods, and showed that SBCE and ASPE are the most accurate. However, SBCE is also the most expensive while DH is the cheapest.

There are numerous advantages of incorporating ASPE and MASA technology to perform targeted high-throughput genetic characterization. ASPE lends itself to this application as it is: Highly specific, cost-efficient, versatile and not labor-intensive. Using ASPE and MASA, the fluorescent signal from the positive results is always ~10-fold higher than that from the negative results. The high signal-to-noise ratio makes it easy to interpret the results analyzed using a Luminex 100. Furthermore, the experiments can be conducted in <8 h. The combined technique is also easily scaled and sample throughput may be increased using 96-well plates instead of single tubes (18). It is also possible that by taking advantage of the full bead array, up to 50 SNP sites could be analyzed in a single tube simultaneously. This would minimize consumption of Tsp DNA polymerase and dNTPs, particularly biotin-labeled dCTP. In the full multiplex assay, the more SNP sites genotyped, the less each reaction costs. In traditional MASA formats, the microspheres are directly coupled to specific probes. Thus, each set of beads is only able to genotype one site. However, in assays combining ASPE and MASA, the microspheres are coupled to a non-specific cZipcode. The cZipcode-coupled microspheres can be used to detect almost any SNP linked to a Zipcode tag and specific capture probes.

In the present study, a potential platform was established for SNP research. The results suggested that SNP260 and SNP386 are not linked to azoospermia and severe oligospermia in Chinese males. However, studies are required to understand the important role of DAZL in spermatogenesis.

The most common methods to detect SNPs currently used in China are based on RFLP or DNA sequencing. However, the current methods are expensive and labor-intensive. In the present study, 196 male patients with azoospermia or severe
oligospermia (sperm density <5x10^9/ml, non-obstructed) that had a normal karyotype and no AZF microdeletions were recruited, along with 40 healthy, fertile controls. SNP260 and SNP386 of DAZL were genotyped using ASPE combined with MASA technology. The combined method for SNP genotyping was high-throughput, accurate and cost-efficient. The technique was successfully applied to detect polymorphisms in the DAZL gene. However, no significant differences were identified between the frequencies of the mutations in SNP260 or SNP386 in the infertile and fertile males (P>0.05). The A260G and A386G polymorphisms of DAZL appeared to not affect male fertility in the Chinese population.

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References


