Quantitative evaluation of partial deletions of the DAZ gene cluster

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Abstract. Partial deletions of the DAZ gene cluster are thought to cause spermatogenesis impairment. The presence of homologous copies of this gene in the Y chromosome does not allow PCR to be used for the identification of this abnormality. Hence, sequence family variants (SFV), following amplification of sY581, sY587 and sY586 and subsequent enzymatic digestion with Sau3A, Dral and TaqI, respectively, and the dual fiber fluorescence in situ hybridization (FISH) have been used to this aim. However, SFV is not always able to identify single DAZ gene copy deletions. We report a quantitative real-time PCR application to evaluate partial deletions of the DAZ gene cluster. To accomplish this, we designed a probe on exon 6 of the DAZ gene which is repeated 3 times in DAZ1, once in DAZ2 and DAZ3 and twice in DAZ4. Five normozoospermic healthy men (C1-C5) having 4 DAZ gene copies by SFV were selected. Fiber-FISH confirmed this outcome in C1-C4, but not in C5 who had an incomplete DAZ gene cluster. The men underwent then quantitative real-time PCR and C1 was arbitrarily selected as calibrator for the calculation of the DAZ gene signals because of the lowest variation in the threshold cycles. Real-time PCR identified 7.2±0.05 signals in C2-C4 and 5.4±0.05 signals in C5. The overall coefficient of variation was 1.4±0.2%. The loss of two signals in this subject may relate to a deletion of both DAZ2 and DAZ3 or of DAZ4 gene. Since SFV showed clearly the presence of DAZ2, it may be hypothesized that C5 lacks DAZ4. In conclusion, these data suggested that quantitative real-time PCR seems to be an effective and reproducible technique that can be used to study the DAZ gene cluster. In addition, the probe chosen for this approach may give indication on the DAZ gene copy deleted.

Introduction

The human Y chromosome, the smallest chromosome in humans, contains about 27 genes which have been shown to be necessary for male sexual differentiation and spermatogenesis (1,2). Some of them are present in a single copy, whereas others are members of multigenic families. Some proteins encoded by these genes have a ubiquitous expression, others are expressed prevalently or exclusively at the testicular level (3-6), but their function during spermatogenesis remains to be fully established.

Clinical evidence has accumulated showing that microdeletions of the long arm of the Y chromosome (Yq) are associated with oligozoospermia or azoospermia and, therefore, this Yq tract has been named azoospermia factor (AZF) (for review see ref. 7). Vogt and collaborators (8) reported that Yq microdeletions cluster in three distinct subregions named AZFa, AZFb and AZFc located in the proximal, central and distal segments of Yq11. AZFa and AZFb microdeletions are associated with a specific testicular histological picture, whereas AZFc microdeletions are associated with a variable testicular pathology (9).

The AZFc region is more frequently deleted in men with spermatogenic failure. This region contains several gene families that are expressed exclusively or predominantly in the testis. The best characterized gene family within this region is the deleted in azoospermia gene family (DAZ) (10) which is present in four nearly identical copies, arranged in two clusters with head-to-head orientation (11). Since this structural conformation does not allow PCR gene amplification, to identify partial deletions of the DAZ gene cluster different methodological approaches need to be developed.

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Sequence family variants (SFV) consisting of amplification of sY581, sY587 and sY586 followed by digestion with Sau3A, Dral and TaqI, respectively, has been utilized to identify partial deletions of the DAZ gene cluster in patients with spermatogenic impairment (12). However, SFV using only these three sequence tagged sites (STS) is not able to identify single DAZ gene copy deletions. Indeed, the lack of DAZ1 can not be ascertained because sY581 amplification followed by Sau3A digestion distinguishes only right from left genes in both clusters; sY587 amplification followed by Dral digestion does not distinguish DAZ1 versus DAZ2 deletion; and sY586 amplification followed by TaqI digestion identifies only the deletion of DAZ2.

Therefore, we have developed a different methodological approach to study partial deletions of the DAZ gene cluster. In this study, we report a quantitative real-time PCR application suitable to evaluate partial deletions of the DAZ gene cluster in 5 normozoospermic men (C1-C5) who were also examined by SFV and fiber fluorescence in situ hybridization (FISH).

Materials and methods

Patient selection. Five healthy men aged 37.4 ± 1.8 years (mean ± SEM) with normal sperm density, motility and morphology, were selected for the study. The protocol was approved by the Institutional Review Board and an informed written consent was obtained from each patient. They had no previous history of cryptorchidism, varicocele, exposure to environmental and/or workplace gonadotoxins. Physical examination showed no gynecomastia and a testicular volume ranging from 15 to 20 ml. Laboratory testing showed normal LH, FSH, testosterone, 17ß-estradiol and prolactin serum concentrations ranging from 15 to 20 ml. Laboratory testing showed normal LH, FSH, testosterone, 17ß-estradiol and prolactin serum levels.

Sequence family variants. SFV was performed by amplifying sY581 which is located in intron 3 of the DAZ gene, sY586 located in intron 6 and sY587 located in intron 10, as previously reported (12).

PCR primers and conditions utilized are those described in GenBank (accession numbers: G63906, G63907 and G63908, respectively). PCR products were analyzed by restriction enzyme digestion according to the manufacturer’s instructions.

Digestion of sY581 with Sau3A produces two fragments of 189 and 63 bp in DAZ1 and DAZ4 and three fragments of 130, 59, 63 bp in DAZ2 and DAZ3. Digestion of sY586 with TaqI produces two fragments of 184, 117 bp in DAZ1, DAZ3 and DAZ4, while in DAZ2 the 301 bp amplification product remains undigested. Digestion of sY587 with Dral produces four fragments of 122, 73, 49, 26 bp in DAZ1 and DAZ2 and three fragments of 107, 49, 26 bp in DAZ3 and DAZ4. The digestion products were run on 12% acrylamide-bis-acrylamide gel electrophoresis and visualized by silver staining.

Fiber-FISH. Extended chromatin fibers were prepared as reported (14). Briefly, lymphocytes were fixed in methanol: acetic acid (3:1) solution and spread onto slides which were washed in PBS solution for 1 min. Chromatin fibers were obtained by using an alkaline solution (0.07 M NaOH-ethanol, 5:2) and then fixed by adding a minimal amount of methanol. Slides were then air dried, dipped in a 70%, 95% and 100% ethanol series for 3 min each, air dried and stored at 4°C until hybridization was carried out.

The probes used for fiber-FISH were obtained using a combination of two DAZ-specific cosmids, generously provided by Dr L.G. Brown, Massachusetts Institute of Technology, Cambridge (USA) (11). Cosmid 63C9 contains exons 2-11 spanning almost the entire sequence of the DAZ gene (GB accession number AC000021). Cosmid 46A6 contains exons 8-11 and extended for 35 kb downstream the 3′ end of the DAZ gene (GB accession number AC000022). DNA probes were labeled by nick-translation with dUTP-Cy3 or dCTP-fluorX (Amersham, Milan, Italy).

The hybridization was carried out as previously reported (13). Briefly, DNA fibers were denatured treating each slide with a 70% formamide/2X SSC (pH 7.5) solution at 80°C for 2 min; slides were then dipped in a 70%, 95% and 100% ethanol series for 3 min each and air dried. Probes were precipitated and denatured at 80°C for 5 min. Slides were then mounted with a coverslip and sealed with rubber cement. Hybridization occurred overnight in a dark humidified container at 37°C, after which the coverslip was removed and the slides were immersed in a post-hybridization wash of 50% formamide/2X SSC, 3 times at 37°C for 5 min. Slides were then mounted in DAPI counterstain and antifade and stored in the dark at 4°C, until microscope observation.

The fluorescent signals were evaluated using a Leica DMRXA2 microscope (Leica Microsystems SpA, Milan, Italy) equipped with a single filter-band for DAPI, FITC and Cy3 and a triple filter-band for the three fluorochromes and a digital camera. Camera control and image acquisition were carried out using the Q-fluor software (Leica).

Quantitative real-time PCR. The number of DAZ gene copies was determined by the TaqMan technology, which exploits the 5'-3' nucleolytic activity of the Ampli-Taq polymerase (15). The method uses the TaqMan probe which specifically anneals the template between the PCR primers. The comparative threshold cycle (Ct) method (ΔΔCt), which compares the amplification of the unknown sample (target) with that of a normalizer and a reference control (calibrator), was used to quantify the number of DAZ gene copies.

SRY single gene was chosen as the normalizer because of its localization within the Y chromosome. C1 was arbitrarily chosen as calibrator because he had 4 DAZ gene copies by SFV and fiber-FISH, as C2-C4, but he consistently showed the lowest threshold cycle variability.

Following the analysis of the GenBank sequence NG_004755, we chose an exon 6 probe which is repeated 3 times in DAZ1, once in DAZ 2 and DAZ3 and twice in DAZ4, as results from NCBI BLAST online software. Therefore, when all DAZ gene copies are present seven signals will be expected from quantitative real-time PCR.

DAZ gene probe consisted of 16 base pairs and contained 5'-fluor labeled VIC; it had the following sequence: 5'-VIC-TCC TCC ACC ACA GTT T-MGB. DAZ gene primers had the following sequence: forward CAG CCA GTT CCT TTG...
SRY gene probe consisted of 20 base pairs and contained a fluorescent reporter (FAM); it had the following sequence: 5’-(FAM)-TTT TTC AGG ACA GCA GTA GA-MGB. SRY gene primers had the following sequence: forward AAT TGG CGA TTA AGT CAA ATT CG; reverse TTG ACT ACT TGC CCT GCT GAT C. All probes and primers were purchased from Applera Italia (Monza, Italy).

The reaction mixture for DAZ and SRY real-time PCR was prepared in separate tubes as follows: 50 ng of DNA, 1X TaqMan Universal PCR Master mix (Applera Italia), 200 nM fluorogenic specific DAZ or SRY probe and 900 nM of each specific primer, in a final volume of 50 μl.

The samples were assayed in quadruplicate in three different real-time PCR runs, using an ABI PRISM 7700 (Applied Biosystems/PE, Foster City, USA). After 2 min at 50°C followed by 2 min at 95°C to activate the polymerase, PCR was carried out as 40 cycles of 15 sec at 95°C (denaturation) and 1 min at 60°C (annealing and extension).

The following formulas were applied to calculate the number of DAZ gene signals:

$$
\Delta C_T = C_T (\text{target or calibrator}) - C_T (\text{normalizer})
$$

$$
\Delta \Delta C_T = \Delta C_T (\text{target}) - \Delta C_T (\text{calibrator})
$$

Number of DAZ signals = $2^{-\Delta \Delta C_T} \times 7$

**Statistical analysis.** Results are shown as mean ± SEM. Coefficients of variation (CV) were calculated as ratio between standard deviation and mean number of signals for each man, multiplied by 100.

**Results**

All subjects selected for this study had repeatedly normal sperm parameters and did not present any microdeletion of the AZF region as evaluated by qualitative PCR analysis. Moreover, none of them showed partial deletions of the DAZ gene cluster by SFV (Fig. 1).

Fiber-FISH confirmed the presence of 4 DAZ genes organized in two clusters with head-to-head orientation in subjects C1-C4. Surprisingly, C5 showed lack of one copy of the DAZ gene family (Fig. 2).

C1 was selected as a calibrator for quantitative real-time PCR and therefore the number of DAZ gene signals was set at 7. Subjects C2-C4 had a mean DAZ gene signal of 7.17±0.05, whereas subject C5 had 5.35±0.05 DAZ gene signals (Table I). The lack of two signals in this man suggested the deletion of both DAZ2 and DAZ3 gene copies or of DAZ4 gene. The deletion of DAZ2 gene copy can, however, be ruled out because of the presence of the 301-bp fragment for sY586 which is specific for DAZ2. The mean CVs of DAZ gene signals were 1.5%, 0.7%, 1.6% and 1.8% for C2, C3, C4 and C5, respectively. The overall CV of the 4 subjects was 1.4±0.2%.

**Discussion**

The present study was undertaken to develop a method to identify all possible partial DAZ gene deletions. SFV following amplification of sY581, sY586, sY587 was shown to be ineffective for this purpose, due to the limited number of markers generally used, which does not allow to fully distinguish the various fragments generated from the enzymatic digestion.

Therefore, an underestimation of the frequency of this chromosome abnormality among oligozoospermic or azoospermic patients may result (12).

In addition, it should be pointed out that SFV may in some cases be ineffective because of the reported polymorphic nature of the DAZ genes (12), recently shown to be due to Y-Y gene conversion (16,17).

Quantitative real-time PCR is a reproducible technique useful to study the DAZ region of the Y chromosome. It was able to detect partial deletions of the DAZ gene cluster in the case of the fertile man C5, where SFV failed to identify the abnormality. The deletion of C5 was confirmed by fiber-FISH.

To develop this methodology, we decided to build a probe on exon 6 which, according to the DAZ gene sequence reported in GenBank (accession number NG_004755), is
Figure 2. Fiber-FISH on extended DNA, obtained from subjects C1 and C5. Upper panel: subject C1, utilized as control, shows all 4 DAZ gene copies. Lower panel: the normozoospermic fertile C5 man presents only 3 fluorescent regions of the DAZ cluster.

Table I. Quantitative real-time PCR evaluation of the DAZ gene signals in 5 normozoospermic men.

<table>
<thead>
<tr>
<th>ID</th>
<th>DAZ C&lt;sub&gt;T&lt;/sub&gt;</th>
<th>SRY C&lt;sub&gt;T&lt;/sub&gt;</th>
<th>ΔC&lt;sub&gt;T&lt;/sub&gt;</th>
<th>ΔΔC&lt;sub&gt;T&lt;/sub&gt;</th>
<th>2&lt;sup&gt;ΔΔC&lt;/sup&gt;</th>
<th>No. of DAZ gene signals</th>
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<tr>
<td>Real-time PCR run 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H&lt;sub&gt;2&lt;/sub&gt;O</td>
<td>40±0</td>
<td>40±0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Calibrator - C1</td>
<td>20.17±0.07</td>
<td>22.50±0.10</td>
<td>-2.33</td>
<td>0</td>
<td>0</td>
<td>7</td>
</tr>
<tr>
<td>C2</td>
<td>19.25±0.10</td>
<td>21.60±0.10</td>
<td>-2.35</td>
<td>-0.02</td>
<td>1.01</td>
<td>7.08</td>
</tr>
<tr>
<td>C3</td>
<td>19.29±0.46</td>
<td>21.63±0.15</td>
<td>-2.35</td>
<td>-0.02</td>
<td>1.01</td>
<td>7.08</td>
</tr>
<tr>
<td>C4</td>
<td>20.32±0.12</td>
<td>22.70±0.09</td>
<td>-2.38</td>
<td>-0.05</td>
<td>1.03</td>
<td>7.20</td>
</tr>
<tr>
<td>C5</td>
<td>22.51±0.11</td>
<td>24.59±0.02</td>
<td>-1.97</td>
<td>0.36</td>
<td>0.78</td>
<td>5.46</td>
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<tr>
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<td></td>
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<tr>
<td>H&lt;sub&gt;2&lt;/sub&gt;O</td>
<td>40±0</td>
<td>40±0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Calibrator - C1</td>
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<td>22.89±0.41</td>
<td>-2.49</td>
<td>0</td>
<td>1</td>
<td>7.00</td>
</tr>
<tr>
<td>C2</td>
<td>19.87±0.39</td>
<td>22.40±0.36</td>
<td>-2.53</td>
<td>-0.04</td>
<td>1.03</td>
<td>7.18</td>
</tr>
<tr>
<td>C3</td>
<td>19.48±0.23</td>
<td>22.00±0.30</td>
<td>-2.53</td>
<td>-0.04</td>
<td>1.03</td>
<td>7.18</td>
</tr>
<tr>
<td>C4</td>
<td>21.07±0.01</td>
<td>23.65±0.36</td>
<td>-2.58</td>
<td>-0.09</td>
<td>1.06</td>
<td>7.43</td>
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<tr>
<td>C5</td>
<td>21.94±0.23</td>
<td>23.03±0.07</td>
<td>-2.08</td>
<td>0.41</td>
<td>0.75</td>
<td>5.28</td>
</tr>
<tr>
<td>Real-time PCR run 3</td>
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<td></td>
<td></td>
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<tr>
<td>H&lt;sub&gt;2&lt;/sub&gt;O</td>
<td>40±0</td>
<td>40±0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Calibrator - C1</td>
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<td>22.33±0.44</td>
<td>-2.13</td>
<td>0</td>
<td>1</td>
<td>7.00</td>
</tr>
<tr>
<td>C2</td>
<td>19.28±0.26</td>
<td>21.41±0.36</td>
<td>-2.12</td>
<td>0.01</td>
<td>1</td>
<td>6.97</td>
</tr>
<tr>
<td>C3</td>
<td>19.61±0.28</td>
<td>21.77±0.21</td>
<td>-2.16</td>
<td>-0.03</td>
<td>1.02</td>
<td>7.13</td>
</tr>
<tr>
<td>C4</td>
<td>20.58±0.13</td>
<td>22.77±0.15</td>
<td>-2.19</td>
<td>-0.06</td>
<td>1.04</td>
<td>7.28</td>
</tr>
<tr>
<td>C5</td>
<td>21.50±0.50</td>
<td>23.23±0.25</td>
<td>-1.73</td>
<td>0.40</td>
<td>0.76</td>
<td>5.32</td>
</tr>
</tbody>
</table>

*Mean ± SD; C<sub>T</sub>, threshold cycle; ΔC<sub>T</sub>, difference between DAZ C<sub>T</sub> and SRY C<sub>T</sub> for each subject; ΔΔC<sub>T</sub>, difference between ΔC<sub>T</sub> of C2, C3, C4 or C5 and ΔC<sub>T</sub> C1. C1 was arbitrarily chosen as calibrator because he had 4 DAZ gene copies by SFV and fiber-FISH; he consistently has showed the lowest threshold cycle variability. The fertile normozoospermic man C5 has a partial deletion of the DAZ gene cluster, demonstrated by loss of two signals.
repeated 3 times in DAZ1, once in DAZ2 and DAZ3 and twice in DAZ4.

Therefore, this particular DAZ probe may suggest which gene copy is deleted. Indeed, the loss of three signals more likely indicates the deletion of DAZ1 or DAZ3/DAZ4, the loss of two signals suggests the deletion of DAZ4 or DAZ2/DAZ3 and the loss of one signal the deletion of DAZ2 or DAZ3.

After the description of the first case in a patient with oligozoospermia (18), partial deletions of the DAZ gene cluster have been reported in patients with impaired spermatogenesis (12,19,20). Indeed, patients with oligozoospermia or azoospermia lack one or more DAZ gene copies with a frequency ranging from 4.3% to 9.1%. These studies have not reported deletions in normozoospermic control men.

Recently, a case report showed that the deletion of 2 DAZ gene copies is still compatible with a normal spermatogenesis (21); this deletion was found in 5 infertile brothers with oligozoospermia or azoospermia and in their normozoospermic father.

More recently, Ferras and colleagues (23) found that 20 out of 50 fertile normozoospermic men (40%) had a partial deletion of the DAZ gene cluster, in particular, lacking the DAZ4, suggesting that this gene copy may not play a relevant role during spermatogenesis (23).

By using a quantitative real-time PCR assay developed on DAZ exon 6, we have also found a normozoospermic fertile man (C5) with a partial deletion of the DAZ gene cluster since he had only 5 out of the 7 signals. This deletion most probably involved the DAZ4 and not DAZ2 plus DAZ3 because in the latter case the deletion of DAZ would also be identified through SFV (sY386-TaqI).

Our observation adds further evidence that an intragenic DAZ deletion may be compatible with a normal spermatogenesis, as we have found in our patient C5 who had normal sperm parameters and was naturally fertile.

In conclusion, quantitative real-time PCR appears to be an effective and reproducible technique that can be used to study the genomic composition of the DAZ region. The probe chosen may give indications about the DAZ gene copy loss, which can be further proved by fiber-FISH.

The existence of a DAZ gene copy deletion in a normozoospermic patient suggests that not all DAZ gene copies may have the same role in human spermatogenesis and that partial deletions of this gene may be reported as a polymorphic deletion event in men with normal spermatogenesis.

References