A *SPGY* copy homologous to the mouse gene *Dazla* and the *Drosophila* gene *boule* is autosomal and expressed only in the human male gonad

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We have isolated a series of human testis poly(A) cDNA clones by cross-hybridization to *SPGY1*, a Y gene homologous to DAZ. Their sequence analysis revealed an identical nucleotide composition in different ‘full-length’ clones, suggesting that all were encoded by the same gene. We mapped this gene to the short arm of chromosome 3 and designated it *SPGY1α* (*SPGY* like autosomal). Comparison of the *SPGY* cDNA sequences with the cDNA sequences of *DAZ* and *SPGY1* revealed two prominent differences. The tandem repetitive structure of 72 bp sequence units (DAZ repeats) is absent. *SPGY1α* contains only one 72 bp sequence unit. Downstream of it, a specific 130 bp sequence domain is present which is absent in *DAZ* and *SPGY1* but present in the mouse gene *Dazla* and in the *Drosophila* gene *boule*. *SPGY1α* encodes an RNA binding protein expressed only in the human male gonad. The data presented give strong evidence that not *DAZ* but *SPGY1α* is the functional human homologue of *Dazla* and *boule*.

INTRODUCTION

In eukaryotes, RNA binding proteins play important roles in various aspects of the splicing process and in the nuclear export of primary transcripts, in cytoplasmic RNA stability and during translation (1). One well characterized group of RNA binding proteins contain one or more copies of the so-called RNA recognition motif (RRM) in their N-terminal regions. The RRM domain consists of ~80 amino-acids (2). Most RRM proteins are expressed constitutively. However, cell type-specific RRM proteins also exist and seem to be functional in the brain (3,4), during sex determination (*Drosophila*: 5,6) and in the germ line (in *Drosophila*, for spermatogenesis, 7; in the mouse, for spermatogenesis and oogenesis, 8,9; in human, for spermatogenesis, 10–12). The recent discovery of the *Drosophila* gene *boule* with a high homology of its RNA binding domain to that of the human gene *DAZ* (13) suggests that RNA binding proteins in the germ line are highly conserved and might fulfil essential regulatory functions during the complex process of spermatogenesis. Gene products of *boule* function during the male meiotic cell cycle at the G2–M transition, and loss of *boule* function results in azoospermia (13). Therefore, it was argued that *DAZ* (because of its distinct sequence homology to *boule*) must have an essential function during the meiotic cell cycle in human spermatogenesis. However, this is inconsistent with the fact that the total deletion of *DAZ* can be inherited from father to son (14) and that deletion of *DAZ* is not always correlated with azoospermia (13). Therefore, *DAZ* deletions seem to be compatible with development of motile spermatozoa, and the primary mutation effect in men with deletion of *DAZ* is subfertility but not azoospermia (14,16). These results suggested that the functional human homologue of *boule* had not yet been isolated and that *DAZ* might not have the same function as *boule* in spermatogenesis (17).

The mouse homologue of *boule* is located on chromosome 17 (9). Because of its distinct sequence homology to *DAZ*, it was designated *Dazla* (*DAZ* like autosomal). Therefore, if a human homologue of *boule*/*Dazla* exists, it is expected to be autosomal and located on the human homologue of mouse chromosome 17, chromosome 6 (17,18).

We have analysed this hypothesis experimentally by screening different cDNA libraries prepared from adult human testis poly(A) RNA with the cDNA probe *SPGY1*, a Y gene homologous to *DAZ* (11). *SPGY1* is member of the *SPGY* (spermatogenesis gene on the Y) locus in distal Yq11 containing several *DAZ*-related genes, in addition to *DAZ* itself (12,16). Their transcripts are characterized by different numbers of a tandem repetitive 72 bp sequence unit (*DAZ* repeat). *DAZ* contains seven such repeats (11), *SPGY1* contains 12 (12,14).

Sequence analysis of *SPGY1* homologous cDNA clones identified a subclass of clones with a common sequence structure absent in *DAZ* or *SPGY1* but present in *boule* and *Dazla*. Two features were most prominent. The 72 bp sequence unit (*DAZ* repeat) was not present in a tandem repetitive structure but as a single sequence unit like in *Dazla* and *boule*. Adjacent to it, a 130 bp sequence domain was analysed which is absent in *DAZ* and

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SPGY1 but present in Dazla and boule. Here we present the isolation of the complete cDNA sequence of this SPGY gene and the sequence of its putative translation product. It maps to the short arm of chromosome 3 and therefore is the first SPGY copy mapped to an autosome. We designated it as SPGYLA (SPGY like autosomal). It encodes RNA binding protein only expressed in the human male gonad. Our results strongly suggest that not DAZ but SPGYLA is the functional human homologue of Dazla and boule in human spermatogenesis.

RESULTS

A SPGY copy with specific cDNA structure is located on chromosome 3

During analysis of the genomic sequence structure of SPGY genes by DNA blot experiments, we found that not all SPGY genes could be located in the AZFc region in distal Yq11. Azoospermic men with deletion of AZFc retained a cross-hybridizing genome fragment also present in the female genome (12). The same was observed in blots with the DAZ probe (11). We were able to identify this non-Y gene copy by analysis of a series of cDNA clones isolated by homology to SPGY1. They were characterized by the absence of all but one of the tandem repetitive 72 bp sequence units (DAZ repeats) found in DAZ (11) and in SPGY1 (12,14), and the presence of a specific sequence unit of 130 nucleotides (sequence position 762–891 in full-length cDNA clone CT355 submitted to the GenBank database; accession no. U66726) which is absent in the sequence of DAZ, SPGY1 and other cDNA clones with tandem repetitive 72 bp sequence units. All cDNA clones containing this sequence structure (one 72 bp sequence unit plus the specific 130 bp sequence unit downstream of it) were identical, so presumably were derived from one SPGY gene. It was mapped by fluorescence in situ hybridization (FISH) analysis to the short arm of chromosome 3 (Fig. 1). As other SPGY copies (analysed so far) were mapped to the AZFc region in distal Yq11 (14,16), the new autosomal copy was designated as SPGYLA (SPGY like autosomal). We scrutinized the mapping position of SPGYLA by PCR analysis of a panel of human–hamster hybrid cell lines containing overlapping subsets of human chromosomes. With the SPGYLA-specific primer pair (LA1-for/LA1-rev), a PCR product was present in all hybrid cells containing human chromosome 3.

When we searched for homology of the SPGYLA-specific 130 bp sequence domain in different databases, a sequence of the same length was identified in the mouse gene Dazla (sequence position 932–1061 in Dazla clone pSG4; accession no. U38690) and a sequence of similar length in the Drosophila gene boule (position 1362–1485 in boule sequence; accession no. U51858). This homology was confirmed at the protein level. Translation of SPGYLA into its putative protein sequence revealed homology to the protein of Dazla along its whole length of 295 amino acids (Fig. 2a). Comparison of these proteins with the boule-encoded protein revealed a high sequence similarity in the regions of their RNA recognition motif (RRM) but also downstream of it, e.g. along the 24 amino acids encoded by the conserved 72 bp sequence unit (DAZ repeat) as described already for boule/Dazla by Eberhart et al. (13). In Figure 2b and c, we display these homologous sequence domains together with that of the DAZ protein. We conclude that SPGYLA most likely also encodes an RNA binding protein.

$S P G Y 1$ is expressed only in adult human testis

Gene copies of the SPGY locus in distal Yq11 are expressed only in human testis (12,16), but Dazla is expressed in the mouse female gonad as well (9). We analysed the tissue expression pattern of SPGYLA by RNA blot and RT–PCR analyses. In the RNA blot experiment, we used a probe of SPGYLA containing only its 3′-untranslated region (UTR) in order to exclude potential cross-hybridizations to testis transcripts of other SPGY genes. A strong hybridization signal was visible with the SPGYLA 3′ UTR probe to a testis poly(A) RNA population of 2.9 kb length (Fig. 3a). This length is similar to the length of the isolated cDNA clone CT355 (2924 bp), suggesting that it corresponds to a full-length testis transcript of the corresponding SPGYLA gene. Additional homology to testis RNA of 2.1 kb was indicated by a weak second hybridization signal. RNA pools prepared from male kidney and male liver in the same experiment did not reveal any cross-hybridization signals. To confirm the testis specificity of SPGYLA transcription in males and to analyse its presence in female tissues, we performed RT–PCR with the SPGYLA-specific primer pair (LA1-for/LA1-rev). An RT–PCR product was only visible in testis tissue and was absent in all female tissues analysed (Fig. 3b). The tissue expression pattern of SPGYLA therefore resembles the pattern of DAZ and SPGY1, but not the pattern of the mouse gene Dazla.

A copy of SPGYLA translocated to the Y chromosome during primate evolution

Male-specific genomic DNA fragments cross-hybridizing to the Dazla probe were only observed in the genome of human and gorilla (9). As SPGYLA and Dazla sequences are homologous to
Figure 2. (a) Comparison of the putative translation products encoded by SPGYLA and Dazla. The SPGYLA-encoded protein is homologous to the mouse Dazla-encoded protein (89% identity and 93% similarity) along the whole sequence. Homology of the RRM domains (b) and 24 amino acid domains (c) encoded by the conserved 72 bp sequence units (DAZ repeat) between SPGYLA, Dazla, DAZ and boule. The 24 amino acid domain of the DAZ-encoded protein was translated from the 72 bp consensus sequence unit (11). The highly conserved protein boxes of the RRM domain: RNP-1 (an octapeptide) and RNP-2 (a hexapeptide) are marked. Both peptide boxes are identical in the SPGYLA-, Dazla- and DAZ-encoded proteins, and in the boule-encoded protein with two conservative amino acid exchanges. Protein sequences were aligned by the software tool PRETTYBOX highlighting identical amino acids by black boxes. Conservative amino acid exchanges are marked by grey boxes according to a matrix scale based on the Dayhof table (in Atlas of Protein Sequence and Structure, 1979, pp. 353–358).

each other, we expected a similar pattern of cross-hybridization in zoo-DNA blots with the human SPGYLA probe. Consequently, male-specific hybridization fragments were observed in the genome of chimpanzee (Pan troglodytes) and gorilla (Gorilla gorilla) (data not shown). However, in a zoo-DNA blot containing genomic DNA samples of a new world monkey (Platyrrhini: Callithrix jacchus) and of two old world monkeys (Catarrrhini: Macaca mulatta, Macaca nemestrina), male-specific SPGYLA cross-hybridizing fragments were only present in the DNA of the old world monkeys but not in the DNA of the new world monkey (Fig. 4). Cross-hybridization signals in the mouse genome, the cattle genome and the wallaby genome (Macropus eugenii) marked the evolutionary conservation of the SPGYLA sequence structure.

Male-specific genomic SPGYLA fragments were not only present in macaques but also in other Catarrrhini. Examination of corresponding blot patterns in baboons (Papio hamadryas; an old world monkey) and tamarins (Saguinus fusicollis; a new world monkey) presented the same basic result. Cross-hybridization to male-specific fragments was only present in the DNA of the old world monkey but not in the DNA of the new world monkey. These results suggest that during primate evolution translocation of a SPGYLA copy to the Y chromosome had occurred after divergence of the lineage of Platyrrhini and before divergence of the lineage of Catarrrhini. This time interval ranges between 36 and 55 million years ago (19).

DISCUSSION

SPGYLA is the human homologue of Dazla and boule

It is well known that genes expressed in different tissues create testis-specific RNAs by alternative splicing. Moreover, genes expressed in testis and transcribed pre-meiotically might be translated only post-meiotically (20–22). This indicates that in the male germ line, the availability of functional gene products
their protein structure. RRM domains consist of one copy of the highly conserved RRM in the N-terminal part of proteins involved in such processes. They were described for Testis-specific RNA binding proteins are strong candidates to be regulated primarily by post-transcriptional processes. (7,13), mouse (8,9,23) and human (10–12). All contain at least one copy of the highly conserved RRM in the N-terminal part of their protein structure. RRM domains consist of ∼80 amino acids.

Two so-called RNP boxes (RNP-1, an octapeptide; RNP-2, a hexapeptide) can be matched in RRM domains of RNA binding proteins in yeast as in human. This suggests that RRM domains have a very ancient origin and that the mode of their interaction with specific RNAs is highly conserved. The specificity of the RNA–protein binding reaction is thought to be achieved by auxiliary sequence blocks in the RRM domain and the C-terminal part of the RRM protein structure (2,24). By these criteria, we suppose that the protein of SPGYLA presented here is the functional human homologue of the protein encoded by the mouse gene Dazla and most likely also of the protein encoded by the Drosophila gene boule.

SPGYLA- and Dazla-encoded proteins have similar molecular weights (SPGYLA, 33 178 Da; Dazla, 33 312 Da) and the sequences are 89% identical (93% similar) along their total length (Fig. 2a). In contrast to this, the DAZ-encoded protein (with one consensus 72 bp sequence unit encoding 24 amino acids) has a higher molecular weight (44 300 Da) and is homologous to the Dazla-encoded protein only along 201 amino acids. Their C-terminal peptides are different because of the presence of the 130 bp sequence unit in the protein-coding region of SPGYLA and Dazla and its absence in DAZ. Therefore, our supposition is that not DAZ but SPGYLA is the functional human homologue of the mouse gene Dazla.

A functional homology between the boule- and SPGYLA-encoded proteins is also most likely but, due to the large evolutionary distance between Drosophila and human, it is difficult to judge it by sequence comparison. Both are RNA binding proteins with different conserved domains including the RRM domain (Fig. 2b). Moreover, the cDNA structure of boule is homologous to the cDNA structure of Dazla and SPGYLA. All three contain one homologous 72 bp sequence unit and the 130 bp sequence unit downstream of it. A functional homology of all three proteins is therefore most likely.

**Figure 3.** Analysis of SPGYLA expression in human tissues: (a) RNA blot hybridization performed with a SPGYLA-specific 3′ UTR probe prepared as described in Materials and Methods. The RNA blot contained poly(A) RNA of human testis (ts), of male kidney (ki) and of male liver (lv). A strong hybridization signal was observed only in testis tissue to a RNA population of 2.9 kb; a second weak testis RNA signal was obtained at 2.1 kb. RNA marker lengths are indicated at the left in kb (GIBCO-BRL: 15620). (b) RT–PCR with the SPGYLA-specific primer LA1-for/LA-1rev was performed on RNA samples of different male tissues (ts: testis; br: brain; pr: prostate; sp: spleen; he: heart) and female tissues (ov: ovary; ki: kidney; lv: liver; lu: lung). The first lane on the left (M) shows fragments of a 1 kb DNA ladder marker (Gibco-BRL: 15615). SPGYLA homologous transcripts (identified by the 161 bp RT–PCR product) were only observed in testis tissue. In a parallel control experiment, the same RNA samples were analysed for transcripts of the constitutively expressed human β-actin gene. Primers actin-forward/actin-reverse amplified a homologous RNA product (identified by the 525 bp RT–PCR product) in each tissue analysed.

**Figure 4.** Hybridization pattern of SPGYLA probe CT355 to a zoo-DNA blot. Male (m) and female (f) genomes of mouse, Xenopus, cattle, Macropus eugenii (wallaby), Callithrix jacchus (Ca. j.), Macaca mulata (Ma. m.), Macaca nemestrina (Ma. n.), and human were restricted with BamHI. Cross-hybridization of CT355 is observed to genomic DNA fragments of all species except Xenopus. The weak signals in the wallaby genomes are due to a low amount of DNA in these lanes. Male-specific cross-hybridization fragments are observed only in the genomes of both macaques and of human. Hybridization stringency was normal as described earlier (14). Filter wash after hybridization was done in 2× SSC, 0.5% SDS at 65°C. The human blot filter was exposed for a shorter time in order to visualize the male-specific BamHI fragment. Positions of λ-HindIII marker fragment lengths are the same and indicated at the left in kb.

**SPGYLA is located on the short arm of chromosome 3**

Mapping of human genes homologous to mouse genes of mouse chromosome 17 show that they are clustered on the short and long
arm of human chromosome 6 (18). Therefore, it was suggested recently that the functional human homologue of \textit{Dazla} should also be located on chromosome 6 (17). However, FISH analysis of the location of \textit{SPGYLA} on human chromosomes does not support this prediction. A distinct hybridization signal was only observed on the short arm of chromosome 3 (Fig. 1). No additional FISH signals were observed on chromosome 6. This result was confirmed by mapping of \textit{SPGYLA} on a human–hamster hybrid cell panel. It is intriguing to note that, despite its sequence homology to \textit{SPGY} genes in distal Yq11, the \textit{SPGYLA} probe does not hybridize on the Y chromosome. We explain this result by the presence of a 1.9 kb long 3′ UTR domain in the \textit{SPGYLA} cDNA structure. This sequence region is specific for this gene and represented 66% of the length of the \textit{SPGYLA} FISH probe used.

No evidence exists for the presence of a male fertility locus on chromosome 3 by genetic or molecular linkage analysis (25). Comparative mapping of DNA loci of human chromosome 3 to mouse chromosomes indicates that this human chromosome is composed of chromatin domains of different mouse chromosomes (3, 6, 9, 14 and 16) (25), and now also of 17.

\textbf{Function of \textit{SPGYLA} in human spermatogenesis}

\textit{SPGYLA} is only expressed in the human male gonad. Its transcription product is polyadenylated and has a length of 2.9 kb. The second weak cross-hybridizing RNA signal (2.1 kb) observed in the testis RNA blot is difficult to explain at the moment. Cross-hybridization to testis transcripts of other Y chromosomal \textit{SPGY} genes was excluded by using a \textit{SPGYLA}-specific 3′ UTR probe. Moreover, \textit{DAZ} and \textit{SPGY} hybridized to testis RNA of 3.5 kb (11,12). It is interesting to note that the mouse homologue \textit{Dazla} also hybridized to two different testis RNA populations (9). Our result might therefore reflect the use of an alternative polyadenylation site in \textit{SPGYLA} transcripts. A second possibility may be that a divergent \textit{SPGYLA} copy is present on the same or another autosome or on the Y chromosome as discussed below.

The expression pattern of \textit{SPGYLA} is different from the pattern of its mouse homologue \textit{Dazla} which is expressed also in the female gonad (9). We have examined the expression of \textit{SPGYLA} in human ovaries of different ages (all before menopause) in order to exclude a possible age-dependent inactivation of the \textit{SPGYLA} gene in the female gonad. However, we were not able to detect any \textit{SPGYLA} transcripts in any ovary tissue by RT–PCR. We cannot exclude expression of \textit{SPGYLA} in certain ovarian stages not available from human females for medical reasons. We also cannot exclude the possibility that the ovarian tissues analysed do not show \textit{SPGYLA} transcription because of their pathological aetiology.

The function of \textit{Dazla} in the male and female gonad is not yet known. The function of \textit{boule} most likely occurs during the male meiotic cell cycle at the G2–M transition (13). Loss of \textit{boule} results in azoospermia. Therefore, if \textit{SPGYLA} is the functional homologue of \textit{boule}, its function is expected to be at human male meiosis, and \textit{SPGYLA} mutations should cause azoospermia.

\textbf{SPGYLA and the SPGY locus}

It is attractive to speculate that Y genes of the \textit{SPGY} locus in distal Yq11 containing a repetitive 72 bp sequence unit (like \textit{DAZ} and \textit{SPGY1}) might have evolved from a functional ancestor \textit{SPGYLA} copy on the Y chromosome with one 72 bp sequence unit. We were able to show that each 72 bp sequence unit is a separate exon unit in the \textit{SPGY} locus (Vogt et al., in preparation). However, molecular analysis of all \textit{SPGY} genes in distal Yq11 is an essential prerequisite to distinguish between \textit{SPGY} gene structures containing repetitive 72 bp exon units and \textit{SPGY} genes containing only one 72 bp exon unit like \textit{SPGYLA}. This work is in progress.

Our analysis of the presence of \textit{SPGYLA} in different primates suggests that a copy of \textit{SPGYLA} was translocated to the Y chromosome at a time interval between 36 and 55 million years ago, after divergence of Platyrrhini and before divergence of Catarrhini. Although, we have not yet confirmed the presence of \textit{SPGYLA} on the Y chromosome of the macaques and baboon by FISH analysis, the male-specific fragments of their genomic DNA blot patterns can be explained most easily by the presence of at least one \textit{SPGYLA} copy on their Y chromosomes. The presence of a \textit{SPGYLA} copy on the human Y chromosome, therefore, also cannot yet be excluded. However, because we detect no FISH signal with the \textit{SPGYLA} probe on the human Y chromosome, its sequence structure is expected to be divergent from that of the autosomal \textit{SPGYLA} gene described here.

\textbf{MATERIALS AND METHODS}

\textbf{DNA analysis}

Hybridization of filter discs (duplicates) prepared from plated testis poly(A) cDNA libraries (Clontech: HL1010b; HL3042b) with the probe \textit{SPGY1} was performed as described by Quertermous (26). Sequence analyses of selected cDNA clones were performed on the Pharmacia A.L.F. express using the Sanger dideoxy chain termination method (Autoread kit, Pharmacia) and the strategy of primer walking on overlapping subfragments from both DNA strands. The sequence of the full-length clone CT355 was submitted to the GenBank database; (accession no. U66726). For sequence alignment analyses, we used the PRETTYBOX and PRETTYPLOT software tools written by Peter Rice, Informatics Division, The Sanger Centre, Cambridge, UK, and part of the HUSAR software package of the German Cancer Institute, Heidelberg.

Genomic DNA samples from peripheral blood lymphocytes of human individuals and animals, and DNA samples of hybrid cell lines were prepared using standard protocols. Conditions used for restriction, blotting, probe labelling and hybridization were the same as described previously (14). Genomic PCR for analysis of the specific 130 bp sequence region in CT355 (\textit{SPGYLA}) was performed with 200 ng of DNA and the primer pair: LA-1for (5′-ATGCCACAGTTGCCCTG-3′) and LA-1rev (5′-AAACCCCGTGAAAGTTCCCA-3′) using the Expand ™ Long Template PCR system (Boehringer). Conditions for amplification were: 10× 1 min at 97°C, 1 min at 65°C, 5 min at 68°C followed by 15 cycles of 1 min at 94°C, 1 min at 65°C, 5 min at 68°C. The final elongation time was 5 min at 68°C. The template was denatured for 5 min at 98°C.

\textbf{RNA analysis}

RNA samples were extracted from the different tissues indicated in the text using the standard guanidium isothiocyanate procedure (27) or ordered from Clontech (male tissues: brain, cat.
no. 64020; heart, cat. no. 64025-1; spleen, cat. no. 64034-1; prostate, cat. no. 64038; female tissues: kidney, cat. no. 63047; liver, cat. no. 64022-1; lung, cat. no. 64023). Poly(A) RNA was purified on oligo(dT) columns (Pharmacia). For RNA blot experiments, RNA samples (20 μg) were electrophoresed on 1% agarose gels containing 2.2 M formaldehyde as denaturing agent. Before blotting, RNAs in the gel were partially hydrolysed in 50 mM NaOH/10 mM NaCl and neutralized in 0.1 M Tris/10× SSC buffer (pH 7.5). Blots were pre-hybridized (5 h) and hybridized (overnight) in a 50% formamide buffer (5× SSPE, 10× Denhardt blocking solution, 2%SDS); 20× SSPE is: 3 M NaCl, 0.2 M NaH2PO4, 0.2 M EDTA (pH 7.4). For the RNA blot experiment we used a SPG/UTR-specific 3′ UTR probe in order to exclude potential cross-hybridizations to testis transcripts of other SPG/UTR genes. The 3′ UTR probe was prepared by amplifying the last 1644 nucleotides of CT355 (positions 1277–2921) by PCR using the CT355 forward primer 355P1 (5′-GAAAATAGATT-TATCTGCAGG-3′) and a vector primer (λ-gtl1rev: 5′-TTGA-CACCAAGACACTGTAA TG-3′) conditions: 25× 1 min at 94°C, 1.5 min at 55°C, 5 min at 72°C. Final extension was for 10 min at 72°C. Denaturation was for 5 min at 94°C. After hybridization, filters were washed under stringent conditions (2× SSC, 0.1% SDS, 65°C). RT–PCR experiments were performed from cDNA aliquots prepared with the SUPERScript™ II system of Gibco-BRL from all RNA samples in the indicated testis. The LA-1orf/LA-1rev primer pair was used for analysis of RNA tissue specificity with the following cycling conditions: 30× 1 min at 94°C, 1 min at 55°C, 1.5 min at 72°C. The final extension was for 5 min at 72°C. Denaturation was for 5 min at 94°C. The length of the RNA amplification product is 161 bp (see Fig. 2b). As a positive control in all RT–PCR experiments, we used actin-forward (5′-ATCTTCGCTGCTGACCTG-3′) and actin-reverse (5′-GCTGTACCCACATCTGGCTG-3′), which are two primers of a human β-actin gene (28), under the same cycling conditions. They are expected to amplify an RNA product of 525 bp in each tissue analysed.

**FISH analysis**

Fluorescence in situ hybridization with CT355 on human male metaphase chromosomes was carried out as described by Köhler and Vogt (29). Both probes were biotinylated by nick translation after a limited DNase I fragmentation. Hybridization signals were amplified once using a biotinylated anti-avidin antibody (Vector Laboratories). Fluorescent signals were analysed in a Zeiss Axiophot microscope with an aligned filter set for DAPI (LP450–490, BP365, FT395) and FITC (LP515–565, BP450–490, FT510). The Axiophot was coupled to a cooled charge-coupled device (CCD) camera for further image processing using the software package ‘Gene JOIN’ as described by Ried et al. (30).

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