**LETTER TO JMG**

**Hypogonadotrophic hypogonadism and cleft lip and palate caused by a balanced translocation producing haploinsufficiency for FGFR1**

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**METHODS**

This study was approved by the Institutional Review Board of Partners Healthcare Inc, encompassing both the Massachusetts General Hospital and the Brigham and Women's Hospital.

**Case report**

The subject is a white man who was aged 24 years at the time of initial diagnosis. He had a history of cleft palate, corrected by surgery. He had no outstanding medical problems other than delayed sexual development and a feminine sounding voice. He had his growth spurt at age 18–19 years, developed sparse armpit hair at age 20, and penile hair at 16–17, but no penile or testicular enlargement. He displayed child-like facial hair, sparse axillary adult appearing hair, and prepubertal chest hair. Based on the presence of cleft palate and hypogonadism, a tentative diagnosis of Kallmann’s syndrome was reached, though the subject did not complain of anosmia. He was prescribed a regimen of testosterone injections, which successfully induced secondary sexual characteristics. At the age of 31, he was seen by a different physician for azoospermia and infertility, and cytogenetic analysis was ordered for the possibility of Klinefelter’s syndrome. The analysis revealed an apparently balanced chromosomal translocation with the karyotype, 46,XY,t(7;8)(p12.3;p11.2). Informed consent for the generation of a lymphoblastoid cell line was obtained in accordance with institutional policies.

**Fluorescent in situ hybridisation analysis**

Breakpoint mapping on chromosome 8 was initiated using clones placed on the cytogenetic map by fluorescent in situ hybridisation (FISH) analysis and on the sequence map by fluorescent in situ hybridisation (FISH) analysis and on the sequence map by sequence tagged sites. Metaphase chromosomes from the patient cell line were prepared for analysis by GTG banding or FISH using standard protocols. Briefly, clones for FISH were

**Key points**

- Kallmann’s syndrome (KS), characterised by hypogonadotrophic hypogonadism and anosmia, can be caused by inactivating mutations of the X linked KAL1 gene, but these mutations account for less than 15% of KS patients. The remaining cases, as well as cases of hypogonadotropic hypogonadism without anosmia, are believed to be caused by mutations at two or more autosomal loci, including a segment of 8p heterozygous for a microdeletion in one KS patient.
- Recently, mutation in FGFR1, the 8p gene encoding fibroblast growth factor receptor 1, has been shown to cause autosomal dominant KS. We report positional cloning of the genomic breakpoints of the balanced reciprocal translocation t(7;8)(p12.3;p11.2) from a male patient with hypogonadotropic hypogonadism and cleft lip and palate. The translocation disrupts FGFR1 (MIM 136350) between exons 2 and 3 and predicts a novel fusion gene product.
- Although various FGFR1 translocations producing fusion proteins have been reported as causes of myeloproliferative disorders, this is the first case in which a constitutional FGFR1 translocation is associated with a developmental disorder.
selected using genome maps provided by the National Center for Biotechnology Information and the University of California Santa Cruz (UCSC) Genomics Bioinformatics Group.\textsuperscript{10,11} Bacterial artificial chromosome (BAC) clones were obtained from CITB-D and RP11 libraries (Invitrogen, San Diego, California, and the Children’s Hospital of Oakland Research Institute) and directly labelled with SpectrumOrange or Green-dUTP (Vysis) by nick translation.

Hybridisations were carried out according to manufacturers’ protocols. Metaphase chromosomes were counterstained with 4,6-diamino-2-phenylindole-dihydrochloride (DAPI), and at least 10 metaphases were analysed using a Zeiss Axioskop microscope. Images were captured with the CytoVision system (Applied Imaging, San José, California, USA). The karyotype, 46,XY,t(7;8)(p12.3;p11.2), was reconfirmed by GTF banding before breakpoint mapping by FISH.

Mapping and cloning of breakpoints

Southern blot analysis of patient lymphoblast genomic DNA with probes D011-A, D011-B, and D011-C to search for altered restriction fragments was carried out using standard protocols. For each lane, 10 μg of genomic DNA from the patient and control were digested with an appropriate restriction enzyme. Fragments were separated on a 1.0% agarose gel and transferred to Hybond-N membrane membranes were washed at 60 °C with 0.15 M NaCl/0.015 M sodium citrate/0.1 % sodium dodecyl sulphate (SDS) for 30 minutes. Autoradiography took place for 16 hours at –70 °C using two intensifying screens. Three hybridisation probes labelled with 32P-dCTP by random priming. Hybridisations were carried out according to manufacturers’ protocols provided. In each experiment, DNA contamination was excluded by the absence of a PCR product in the sample without reverse transcriptase, amplified under the same conditions as the reverse transcribed RNA sample. Nested PCR was carried out using Pfu polymerase (Gibco BRL) with the following primer sets, annealing at 56 °C for 30 seconds with an extension for one minute 40 seconds: TENS1-FGFR1: 5’CTGAGAAGCCCTCGAGTGC3’+5’CAAGATCTGGACATAAGGCAG3’, 5’GGCAAGGCGATCCTGCC3’+5’CTATAGGACATGATTT3’. Then, the der(8) junction fragment was amplified by nested PCR using the primer sets: 5’GCCAGCTCTACAGGTTAAG3’; 5’CCAGGCTCAGGTGAGG3’. Fusion transcript amplification

Total RNA was isolated from patient and control lymphoblastoid cell lines with the RNeasy Mini Kit (Qiagen, Valencia, California, USA). Reverse transcription of total RNA (1 μg) was undertaken by using either random hexanucleotide priming and Superscript II (Gibco BRL, Gaithersburg, Maryland, USA) or the SMART–PCR cDNA synthesis kit (Clontech, Palo Alto, California, USA) according to the protocols provided. In each experiment, DNA contamination was excluded by the absence of a PCR product in the sample without reverse transcriptase, amplified under the same conditions as the reverse transcribed RNA sample. REST was confirmed by using the ABI Prism 377/373 DNA Sequencer (Applied Biosystems, Foster City, California, USA). RESULTS

Delineation of the breakpoint region on 8p11.2

To identify the genes potentially disrupted in the patient, we first mapped the translocation breakpoints using FISH. Two BAC clones selected from the UCSC map as starting clones for FISH—GS1-211B7 and GS1-165D4 from 8p12 and 8p11.2—mapped telomeric and centromeric to the breakpoint,
respectively, showing that the chromosome 8 breakpoint was contained in a 12 Mb region. Subsequent experiments were carried out using BACs chosen from within this region to narrow the candidate region until a breakpoint crossing BAC clone was identified. Seventeen BACs were examined, of which eight were proximal to the breakpoint and nine distal. The final BAC, RP11-100B16, hybridised to the normal chromosome 8, and both der(7) and der(8) chromosomes, indicating that it spans the translocation breakpoint (fig 1B). Additionally two BAC clones that partially

Figure 2 (A) Schematic presentation of positional cloning on chromosome 8. The 24 kb breakpoint region determined by fluorescent in situ hybridisation (FISH) was further narrowed to 1.6 kb by Southern blot hybridisation, which detected junction fragments with five different restriction enzymes. The breakpoint is located between exons 2 and 3 of FGFR1. The FGFR1 gene is not to scale. (B) Southern blot hybridisation of genomic DNA from the translocation patient with 877 bp probe D011-C from intron 2 of FGFR1. Note the detection of five altered fragments caused by the translocation junction, generated by enzymes BamHI, Drai, EcoRV, HindIII, and Sspl. C, genomic DNA from karyotypically normal control; P, patient genomic DNA.
overlap with RP11-100B16—RP11-265K5 and RP11-359P11—mapped distal and proximal to the breakpoint, respectively (fig 2A). Based on the sequence of these BACs, the location of the breakpoint region was confined to ~24 kb of DNA.

**Southern blot hybridisation and cloning of the breakpoints on 8p11.2**

To localise the breakpoint region in 8p11.2 further, three DNA fragments—D011-A, D011-B, and D011-C—were amplified by PCR from the narrowed 24 kb region (fig 2A) and used to probe patient DNA on genomic blots. D011-B and D011-C both detected altered restriction fragments due to the translocation (fig 2B), narrowing the breakpoint to 1557 bp on the BAC restriction map and suggesting that the breakpoint is between exons 2 and 3 of FGFR1 isoform 1 (fig 2A). We cloned and sequenced junction fragments spanning the breakpoints from both derivative chromosomes, which revealed that the translocation is perfectly balanced, without the gain or loss of any sequence. The sequences of the breakpoint regions for the der(7) and der(8) chromosomes are given in fig 3.

**Delineation of the breakpoint region on 7p12.3**

BAC clones RP11-183O1 from 7p22.1 and RP11-341D24 from 7p11.2 were used as starting clones for FISH, and mapped distal and proximal to the breakpoint, respectively, indicating that the chromosome 7 breakpoint was contained in a 49 Mb region. Using randomly selected BACs, the breakpoint region was narrowed to ~1.3 Mb, flanked by RP11-126K7 and RP11-271O10, which map telomeric and centromeric to the breakpoint, respectively. After the breakpoint was cloned and sequenced, on the basis of the chromosome 8 findings (see below) the junction sequence was found to be located in RP11-549I23. This was confirmed by FISH, showing three signals, one each on chromosome 7 and both derivative chromosomes (data not shown).

**TENS1 in 7p12 and FGFR1 in 8p11 are disrupted**

The chromosome 7 breakpoint lies in intron 15 of TENS1 (AF417489, 1445 amino acids, between nucleotides 64702 and 64703 of GenBank entry AC073341, 15665 bp downstream of exon 15), while the breakpoint on chromosome 8 is in intron 2 of FGFR1 (NM_000604, 822 amino acids, between nucleotides 77382 and 77383 of GenBank entry AC087623, 22414 bp downstream of exon 2). The chromosome 8 breakpoint maps within a SINE/Alu repetitive sequence while the breakpoint in TENS1 occurs in unique intronic sequence with no apparent homology to the chromosome 8 breakpoint region.

**Fusion transcript amplification**

The location of the translocation breakpoint predicts two putative reciprocal in-frame fusion transcripts TENS1/FGFR1 and FGFR1/TENS1 (fig 4A). On the derivative chromosome 7, exons 1–15 of TENS1 are predicted to join with exons 3–18 of FGFR1 isoform 1 and to form a 5891 bp TENS1ex1-15/FGFR1ex3-18 transcript that spans 31 exons and encodes 1675 amino acids without frameshift, from the normal TENS1 initiation codon (ATG) in exon 1 to the termination codon (TGA) in exon 31 (corresponding to the normal stop codon in FGFR1 exon 18). The putative fusion protein would consist of the first 883 amino acids of TENS1 joined to the final 792 amino acids of FGFR1 (fig 4, panels A and B).

On the derivative chromosome 8, exons 1-2 of FGFR1 are joined to exons 16–26 of TENS1, predicting a 2546 bp FGFR1ex1-2/TENS1ex16-26 transcript that spans 31 exons and encodes 592 amino acids. Again there is no frameshift, as the start codon and stop codon occur at exon 2 and exon 13, respectively, at the same positions as the corresponding exon 2 of FGFR1 and exon 26 of TENS1. The predicted FGFR1-TENS1 fusion protein would contain the first 30 amino acids of FGFR1 followed by the final 562 amino acids of TENS1 (fig 4, panels A and B).

To establish whether either fusion transcript is expressed in a lymphoblastoid cell line from the patient, we carried out reverse transcriptase PCR (RT-PCR). Only the TENS1-FGFR1 fusion transcript was detected (fig 5), but sequencing revealed the skipping of FGFR1 exon 3, an alternative splicing pattern also seen in several native FGFR1 encoded isoforms. The 5624 bp TENS1ex1-15/FGFR1ex3-18 transcript encodes a fusion protein of 1586 amino acids comprising the first 883 amino acids of TENS1 joined to the final 703 amino acids of FGFR1 (fig 4, panels A and B).

**Mutation analysis of the non-translocated FGFR1 allele**

Mutation analysis of the second non-translocated FGFR1 allele from the patient, done by SSCP and direct sequencing, identified only a heterozygous nucleotide difference, 345 C→T in exon 3, a known SNP (NCBI reference SNP ID: rs2915665) which does not alter the Ser amino acid encoded at this site. Thus the presence of the translocated allele of the FGFR1 results in a disease phenotype without a corresponding coding sequence mutation in the alternate FGFR1 allele.

**DISCUSSION**

The TENS1 locus encodes tensin-like SH2 domain-containing protein 1 (also known as tumour endothelial marker 6, tensin 3), a 1445 amino acid protein named for its similarity with tensin, an actin filament crosslinking protein found in focal adhesions.13 14 TENS1 protein contains a protein tyrosine
phosphatase domain in the amino-terminal region and Src homology 2 (SH2) and phosphotyrosine binding domains near its carboxyl-terminus. FGFR1 encodes several different isoforms of a transmembrane protein, the extracellular moiety of which interacts with fibroblast growth factors (FGFs), setting in motion a cascade of downstream signals, ultimately influencing mitogenesis and differentiation; it is characterised by two or three extracellular immunoglobulin-like loops (depending on inclusion of exon 3), a transmembrane domain, and an intracellular tyrosine kinase domain.

The predicted TENS1/FGFR1 fusion protein lacks the FGFR1 signal peptide and the first Ig-like domain, but contains Ig-like domains 2 and 3, which are sufficient for specific FGF binding, and an intact tyrosine kinase domain region, suggesting the potential for functionality (fig 3B). However, the related KS phenotype in a patient hemizygous for 8p due to the deletion region suggests that the translocation produces hypogonadotropic hypogonadism as a result of haploinsufficiency for FGFR1. Consistent with this view, there was no evidence for an FGFR1 mutation on the second allele in SSCP/sequence analysis in any of 18 exons and splice junctions. The apparent functional hemizygosity for FGFR1 in the translocation patient probably reflects a failure to direct the FGFR1 functional domains to the proper location in the plasma membrane owing to the absence of the appropriate signal peptide and the presence of the large TENS1 moiety. While this work was being completed, Dode et al, and subsequently Sato et al, reported several truncating and missense FGFR1 mutations in KS patients, some with cleft lip and palate, consistent with the haploinsufficiency in

Figure 4 (A) Disruption of genes FGFR1 and TENS1 by t(7;8), resulting in two in-frame fusion genes. White boxes indicate 5’ and 3’ untranslated regions of exons. TENS1 and FGFR1 coding exons are shown as grey boxes and blue boxes, respectively, numbered above the gene in the same colour, whereas exons of fusion genes are numbered in orange below the gene. Note that the stop codons of the two fusion genes are at the same exon locations as in the corresponding wild-type genes, as there is no frameshift. In the amplified TENS1-FGFR1 fusion transcript, exon 3 of FGFR1 is skipped. Notable exons are numbered and the direction of gene transcription is indicated by arrows. The sizes of exons and introns is not to scale. (B) TENS1, FGFR1, FGFR1-TENS1, and TENS1-FGFR1 protein domains. In the amplified fusion protein TENS1-FGFR1, the transmembrane domain (TM), and tyrosine kinase domain (TK) of FGFR1 are not affected by the translocation, but the signal peptide (SP) of FGFR1 and the first immunoglobulin (Ig) like domain are absent. The translocation does not directly disrupt the amino-terminal protein tyrosine phosphatase domain (PTP) or the carboxyl terminal Src homology 2 (SH2) and phosphotyrosine binding (PTB) domains of TENS1, but does segregate the coding sequences for these domains to different fusion transcripts.
the patient reported here.\textsuperscript{a, b, c} However, the absence of frank anosmia in the current patient indicates that sufficient FGFR1 function may have been maintained to prevent the degree of agenesis of the olfactory lobes typical in KS.

As the translocation patient displays no obvious phenotypes distinct from those seen in patients with KS associated FGFR1 point mutations, the disruption of TENS1 does not seem to contribute to the patient’s abnormalities. This suggests that heterozygous inactivation of TENS1 is without dramatic consequence, but the possibility that the predicted fusion proteins effectively provide normal TENS1 function cannot be excluded.

The X linked form of KS is associated with inactivating mutation of the KAL1 gene, encoding anosmin 1, a secreted proteoglycan binding protein with similarities to neuronal cell adhesion molecules.\textsuperscript{17} Anosmin 1 interacts with heparan and chondroitin sulphates to promote cell adhesion and neuronal outgrowth, and has been implicated in the migration of gonadotropin releasing hormone (GnRH) producing neurones and olfactory axonal fibres, though the receptor system through which it acts remains uncertain.\textsuperscript{18} Notably, FGFR1 activation by binding to FGF ligands involves receptor dimerisation that also requires heparan sulphate proteoglycan binding.\textsuperscript{19} Indeed, FGF2 ligand and FGFR1 have been co-crystallised with heparin, and the structure of the complex defined.\textsuperscript{20} The common association with heparan sulphates and the similar effects of KAL1 and FGFR1 inactivating mutations support the suggestion that the FGFR1 signalling pathway participates directly in mediating anosmin 1 function.\textsuperscript{21}

The translocation patient reported here and the KS patients reported by others also support the view that haploinsufficiency for FGFR1 is a cause of cleft lip and palate.\textsuperscript{7, 8, 16} Interestingly, FGFR1 gain of function mutations have previously been associated with the craniosynostosis of Pfeiffer’s syndrome and in the Jackson–Weiss syndrome.\textsuperscript{22, 23} These syndromes can also be caused by mutations in FGFR2, which has also been associated with cleft palate in Apert’s syndrome, indicating that the two receptors may function in the same signalling pathway.\textsuperscript{24} This suggests that FGFR2, located at 10q26, may be an excellent candidate for an additional KS or idiopathic hypogonadotropic hypogonadism locus. Hence it would be of interest to determine whether FGFR2 is disrupted by translocation in a KS patient with a de novo unbalanced der(1)t(1;10)(q44;q26)\textsuperscript{8, 25, 26}

It is noteworthy that a variety of fusion proteins involving FGFR1 underlie the 8p11 myeloproliferative syndrome (EMS)/stem cell leukaemia–lymphoma syndrome, presumably because of constitutive activity of the tyrosine kinase domain.\textsuperscript{27, 28} However, neither Pfeiffer’s syndrome nor the Jackson–Weiss syndrome shows a myeloproliferation defect, despite being likely to mislocalise a portion of FGFR1 containing the tyrosine kinase domain. This reinforces the view that both the fusion partner and the site of the breakpoint are likely to be critical in producing constitutive tyrosine kinase activity in a manner that leads to malignancy. This is the first demonstration that constitutional translocation of FGFR1 can lead to abnormal development rather than to myeloid disorder, and provides a basis for more detailed structure–function comparison of the respective fusion proteins.

**ELECTRONIC DATABASE INFORMATION**

GenBank accession numbers: FGFR1, AC087623, NM_000604; TENS1, AC073341, AF417489.

dbSNP information: rs2915665.

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