Brachymesomelic Dysplasia With Peters Anomaly of the Eye Results From Disruptions of the X Chromosome Near the SHOX and SOX3 Genes

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We report on a mother and son affected with an unusual skeletal dysplasia and anterior segment eye abnormalities. Their skeletal phenotype overlaps with the SHOX-related skeletal dysplasias and is intermediate between Leri–Weill dyschondrosteosis (LWD) and Langer Mesomelic dysplasia (LMD). The mother has bilateral Peters anomaly of the eye and was reported as having a new syndrome; the son had severe bilateral sclerocornea. Chromosome analysis showed that the mother has a pericentric inversion of the X chromosome [46,X,inv(X)(p22.3q27)] and the son, a resultant recombinant X chromosome [46,Y,rec(X)dup(Xq)inv(X)(p22.3q27)]. The observed skeletal and ophthalmologic abnormalities in both patients were similar in severity. The additional features of developmental delay, growth retardation, agenesis of the corpus callosum, cryptorchidism and hypoplastic scrotum in the son are consistent with Xq28 duplication. Analysis of the son’s recombinant X chromosome showed that the Xp22.33 breakpoint lies 30–68 kb 5’ of the SHOX gene. This finding suggests that the skeletal dysplasia in both mother and son is allelic with LWD and LMD and results from a novel mis-expression of SHOX. Analysis of the Xq27.1 breakpoint localized it to a 90 kb interval 3’ of the SOX3 gene, supporting a novel role of SOX3 misexpression in the development of Peters anomaly of the eye.

Key words: Peters anomaly; sclerocornea; dyschondrosteosis; mesomelic dysplasia; X-chromosome inversion; SHOX; SOX3

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INTRODUCTION

We previously reported on a patient with a unique mesomelic skeletal dysplasia and Peters anomaly of the eye [Kivlin et al., 1993]. At that time, this combination of findings had not been reported, and we proposed that this patient was affected with a provisionally unique genetic syndrome. To our knowledge, there have been no other reports of this condition. This patient subsequently had a son affected with strikingly similar skeletal and ophthalmologic abnormalities. He had additional congenital
anomalies that prompted high resolution chromosome analyses that showed a pericentric inversion of the X chromosome [46,X,inv(X)(p22.3q27)] in the mother and a recombinant X chromosome in the son [46,Y,rec(X)dup(Xq)inv(X)(p22.3q27)] with deletion of the Xp terminus and duplication of Xq27 → qter.

The skeletal anomalies seen in this mother and son overlap with, but are distinct from, those seen in Leri–Weill dyschondrosteosis (LWD) and Langer mesomelic dysplasia (LMD) and led to the hypothesis that the SHOX gene may be disrupted by the inversion. The short stature homeodomain (SHOX) gene is located in the pseudoautosomal regions of the short arms of the X and Y chromosomes and encodes a homeodomain protein thought to regulate the development of chondrocytes [Marchini et al., 2004]. Deletions and point mutations affecting the SHOX gene cause a range of skeletal abnormalities. Leri–Weill dyschondrosteosis (LWD [MIM 127300]) is a pseudoautosomal dominant disorder characterized by short stature, mesomelia and Madelung deformity. Approximately 65% of LWD is explained by heterozygous mutations of the SHOX gene [Belin et al., 1998; Shears et al., 1998; Schiller et al., 2000; Falcinelli et al., 2002; Flanagan et al., 2002] or by deletions of regions between 30 and 530 kb 3′ of the gene [Benito-Sanz et al., 2006]. These 3′ deletions may affect long-range SHOX regulatory elements. For about 35% of LWD patients the molecular defect has not been identified. Homozygous loss of SHOX has been demonstrated in Langer mesomelic dysplasia (LMD [MIM 249700]) [Belin et al., 1998]. LMD is characterized by more severe mesomelia with short square ulna, bowed radii, bowed tibia and hypoplastic fibulae [Langer, 1967]. Heterozygous mutations in the SHOX gene have also been described in 2–20% of patients with idiopathic short stature [Rao et al., 1997; Binder et al., 2000; Rappold et al., 2002; Stuppia et al., 2003; Huber and Cormier-Daire, 2004], and haploinsufficiency for SHOX is considered etiologically related to the short stature seen in Turner syndrome [Ellison et al., 1997; Rao et al., 1997; Clement-Jones et al., 2000].

Peters anomaly has not been described in association with SHOX-related disorders or other X-chromosome abnormalities, suggesting that the Xq27 breakpoint disrupts a novel gene involved in anterior segment abnormalities. Peters anomaly of the eye encompasses a range of abnormalities including corneal opacity, adherent iris strands and keratolenticular cataract. Peters anomaly is etiologically related to anterior segment mesodermal dysgenesis (ASMD), both abnormalities being caused by mutations in an overlapping set of genes. Mutations causing Peters anomaly and/or ASMD have been found in the genes encoding PAX6 [Jordan et al., 1992; Hanson et al., 1994], PITX2 [Semina et al., 1996; Doward et al., 1999], PITX3 [Semina et al., 1998], FOXC1 [Nishimura et al., 2001; Honkanen et al., 2003], FOXE3 [Semina et al., 2001; Valleix et al., 2006], EYA1 [Azuma et al., 2000], CYP1B1 [Stoilov et al., 1997], and FGFR2 [McCann et al., 2005]. Mutations in the gene encoding B3GALTL cause Peters-Plus syndrome [Lesnik Oberstein et al., 2006], an autosomal recessive condition characterized by anterior chamber defects and a primarily rhizomelic skeletal dysplasia [Van Schooneveld et al., 1984]. Sclerocornea and aphakia are associated with microphthalmia and other anomalies in patients with mutations in the gene encoding SOX2 [Fantes et al., 2003; Ragge et al., 2005].

To better define the genetic etiology of the atypical pattern of dominantly inherited skeletal and ocular anomalies in mother and son, we undertook further studies to localize the breakpoints of the inversion and recombinant X chromosomes in these cases.

MATERIALS AND METHODS

Patients and Samples

Blood was collected with informed consent and approval from the University of Utah Institutional Review Board (IRB) protocol number 7551.

Chromosome Analysis

High-resolution G-banded karyotype analysis was performed on peripheral lymphocytes using standard techniques. Fluorescence in situ hybridization (FISH) was performed using subtelomeric probes for the Xp/Yp and Xq/Yq regions (Vysis, Downers Grove, IL), and with BAC RP11-839D20 (Spectral Genomics, Houston, TX), by standard methods.

Comparative Genomic Hybridization Analysis

Array comparative genomic hybridization (aCGH) analysis was performed using the 1 Mb average resolution SpectralChip 2600 (Spectral Genomics), according to the manufacturer’s protocol. Scanning was performed with Axon’s GenePix 400B microarray scanner and the images were analyzed with SpectralWare 2.0 (Spectral Genomics) for the preparation of ratio plots. For high-resolution CGH, chip 8 of the 8-array Set with an average of 713 bp probe spacing was performed by the manufacturer, and relative ratios (RRs) were determined using the SignalMap software package, version 1.8 (NimbleGen Systems, Madison, WI).

Quantitative Real-Time PCR

Custom TaqMan gene probes for five loci spanning the deletion boundary and for exons 1 and 23 of the PDGFRα gene at 4q12 (used as controls) were designed by ABI (Applied Biosystems, Foster City,
CA). Primer and probe sequences were as follows: P1For: TGCTGCGCTCCTGAG. P1Rev: CCGGTCCCTGAAGATGAAAGGT. P1Fam: CCACGTACAGGACAACAGCTC. M1For: GGTGGCTCAGAGCCTCTAGGA. M1Rev: CATTAAATGAGCGTCTCTCAACTCT. M1Fam: TTTCGGAAAGGAAGTCTG. M2For: CATTGCCATGT-AATCCGGCACTCCCATT. M2Rev: GTTGTCAGAACGCCCTAGCT. M2Fam: CCACTGCGCTGAACACTAA. M3For: GGAGGCTTTAAAAT. M3Rev: GCCCTAGCT. M3Fam: TTGCTAGG. S5For: CCCCCAGGAGGAAAAATGTC. M3Fam: TTTAGG. S5Rev: TCAGGACCGGCGGAAAAAG. S5FAM: CCTTGCGGCCGCGCTG. Genomic DNA was combined with TaqMan probes and Universal master mix and analyzed on an ABI 7900HT RT-PCR system, according to manufacturer specifications. Patient and control reactions were done in triplicate and threshold cycle (Ct) measurements were averaged. RRs were calculated using the ΔΔCt method [Livak and Schmittgen, 2001].

X-Inactivation Studies

The pattern of X-inactivation was determined in the mother’s lymphocytes based on the methylation status of HpaII restriction-enzyme sites closely linked to the polymorphic CAG repeat in the human androgen-receptor (AR) gene, as previously described [Allen et al., 1992].

In Silico Analysis

The March 2006 assembly of the UCSC Human Genome (http://www.genome.ucsc.edu) was used as the primary tool to map probes and genes to genomic sequence. Alignment of deleted probes to the known genomic region surrounding breakpoints was performed using the Sequencher (version 4.5) software package (Gene Codes, Corp. Ann Arbor, MI). Alignment of genomic sequences surrounding the breakpoints to each other was performed using the NCBI BLAST 2 sequences tool (http://www.ncbi.nlm.nih.gov:80/BLAST/).

RESULTS

Clinical Reports

The proband (mother) was described previously [Kivlin et al., 1993]. Briefly, she was a full-term infant with a birth weight of 2,880 g (~1 SD). OFC 36.5 cm (~1.0 SD), length 45.5 cm (~1.7 SD). Shown in Figure 1A, she had minor variation in the shape of her ears with a flat forehead and double hair whors, but a normal jaw. Distal limbs were short and limited extension of the elbow to 150° with ulnar deviation of the wrists was noted. There was camptodactyly of the 4th fingers and rotation of the 5th finger toward the palm bilaterally. The lower limbs showed decreased muscle mass and reduced extension of the knees to 160°. She had bilateral hip dislocation and talipes equinovarus with cavus deformities. Radiographs of the upper extremities (Fig. 1C) showed short, bowed radii and markedly short, thickened ulnae. Lower extremity films (Fig. 1E) showed shortening of the tibiae and fibulae, but to a lesser degree, without fibular hypoplasia. Her ocular anomalies consisted of bilateral corneal clouding that was more severe on the left (Fig. 2A) with bilateral anterior iris synchia and absence of Descemet’s membrane seen on histological exam of the left cornea after transplantation (Fig. 2C).

The proband was seen again at age 20 years then pregnant at 19 weeks gestation. She was 124.5 cm height and blind in her left eye (after two corneal transplants). She was ambulatory over short distances, but relied heavily on assistive devices, including an electric wheelchair. She was developmentally normal, although she had required some resource classes in school. Her partner was of low normal height, with normal extremities. He was affected with Neurofibromatosis type 1 and had mild learning disabilities.

Ultrasound examination of the fetus at 19 weeks revealed lower normal length femora and humeri, short distal extremities, bilateral talipes equinovarus and bilateral ulnar deviation of the hands, consistent with a recurrence of the mother’s skeletal dysplasia in the fetus. By 25 weeks of gestation, all long bones except the humeri were short (3–6 SD below the mean), and the radii and ulnae were thickened and curved. The pregnancy was otherwise uncomplicated.

At 38.5 weeks of gestation, the infant was delivered by primary cesarean for inadequate maternal pelvis. The male infant weighed 2,335 g (~1.9 SD). His length was 36 cm (~3.3 SD). His OFC was 32 cm (~1.7 SD). Similar to his mother, he had brachymesomelia shortening of his extremities, knee contractures, bilateral clubfeet, camptodactyly and ulnar deviation of his wrists. Additionally he had undescended testes, a hypoplastic scrotum, and agenesis of the corpus callosum. There was one café au lait spot on the abdomen, but no other neurocutaneous stigmata. He had significant feeding problems, constipation and severe gastroesophageal reflux, requiring a nissen funduplication and gastrostomy tube placement. Despite these measures, he continued to have persistent severe growth retardation with a weight of 4.5 kg, an OFC of 42 cm and a length of 48 cm (all well below the 3rd centile) at 15 months of age. Cognitive development was also significantly delayed. He smiled at 6 months of age, but never rolled or sat without support. Although visually impaired, he was interactive and would play, but had no words at 22 months of age. Echocardiography, renal ultrasound and thyroid function studies
were all normal. The patient went on to have a complicated medical course with an acute presentation of sepsis and bowel necrosis, requiring total colectomy at 22 months. He had recurrent fevers and required treatment for septic shock. He had multiple respiratory infections with chronic lung disease. He eventually succumbed to these complex medical issues at 22.5 months of age.

His radiographic findings were nearly identical to those of his mother as an infant (Fig. 1D,F). Ocular examination showed complete bilateral sclerocornea (Fig. 2B) with formed anterior chambers and no iris-corneal adhesions by ultrasound biomicroscopy (Fig. 2D). No evidence for microphthalmia was observed. The axial length of the eyes was 22.0 mm OD and 21.7 mm OS.
Chromosome Analyses

The similar presentation of skeletal dysplasia and anterior chamber eye anomalies seen in both mother and son suggested a new autosomal dominant syndrome. However, based on the presence of additional multiple congenital anomalies in the son, chromosome analyses were performed that revealed a large pericentric inversion of the X chromosome in the mother [46,X,inv(X)(p22.3q27)] and a resultant recombinant X chromosome [46,Y,rec(X)dup(Xq)inv(X)(p22.3q27)] in the son (Fig. 3A). The inversion in the mother was found to be de novo based on normal cytogenetic studies done in her parents. FISH was performed using probes from the Xp/Yp and the Xq/Yq subtelomeric regions. Both Xp and Xq probes hybridized to the normal and inverted X chromosomes in the mother (Fig. 3B). On the son’s recombinant chromosome, FISH showed absence of the Xp probe signal (Fig. 3C), and presence of Xq signals on both ends (Fig. 3D), thus documenting a deletion of distal Xp and disomy for distal Xq.

Comparative Genomic Hybridization

We performed aCGH to investigate the possibility of cryptic copy number changes near the breakpoints
of the mother’s inversion, and to better characterize the extent of gains and losses on the son’s recombinant X chromosome. Using the 1 Mb Spectral Genomics array, no gains or losses were detected on the mother’s X chromosome near the inversion breakpoints. The son’s analysis showed a single terminal clone (RP11-839D20) deletion at Xp22.33, and a 12-clone duplication in the Xq27-28 interval. This array mapped the Xp22.33 and Xq27 breakpoints to 5.4 and 7 Mb intervals, respectively.

Next, to map the Xq27.1 breakpoint with higher resolution, aCGH was performed using chip 8 of the 8-chip NimbleGen 713 bp array. The Xp22.33 pseudoautosomal region was omitted from this chip. This analysis localized the Xq27.1 breakpoint to a 13 kb interval that lies 92 kb centromeric of the SOX3 gene (Fig. 4) within a cluster of LINE elements. The SOX3 gene is oriented such that the breakpoint lies 3' of the gene.

Quantitative Real-Time PCR

The aCGH analysis mapped the Xp22.33 breakpoint to a large interval of 5.4 Mb that contains the SHOX gene (Fig. 5), which is mutated in two skeletal dysplasia syndromes with overlap to the phenotypes seen in our patients. To determine if the SHOX gene was disrupted by the inversion breakpoint in the mother or deleted in the son, quantitative real-time PCR (RT-PCR) was used. TaqMan RT-PCR probes were designed at five sites across the interval (Fig. 5A), including a probe within RP11-839D20, known to be deleted by aCGH analysis, and a probe at the 5' end of the SHOX gene. TaqMan probes for two loci at 4q12 were used as controls. The analysis showed that three probes were deleted in the son, but that the 5' end of the SHOX gene was not disrupted. A representative amplification plot is shown in Figure 5B. The same analysis found no deleted probes in the mother. The inversion breakpoint was mapped to the 34 kb interval that lies 33–68 kb distal to the SHOX gene (Fig. 5A).

X-Inactivation Studies

The similarity of skeletal and ocular phenotypes in mother and son raised the possibility of skewed X-inactivation in the mother. X-inactivation studies showed a partially skewed pattern of inactivation, and the AR allele, found to be active on about 75% of the mother’s X chromosomes, was the allele inherited by her son on the recombinant chromosome (data not shown). However, we could not determine whether this allele was present on the inverted chromosome in the mother.

Breakpoint Analysis

The Xq27.1 breakpoint occurs within a 13 kb LINE element cluster, but no homologous LINE elements

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FIG. 3. A. High-resolution chromosome banding (650-band level) analysis of the mother revealed a pericentric inversion of the X chromosome (left) with breakpoints in distal Xp and Xq. The son carried a recombinant X chromosome (right) with deletion of Xp and partial duplication of Xq. Ideograms of the three X chromosomes are shown (center) with arrows marking breakpoints in Xp23.3 and Xq27. FISH in the mother using subtelomeric probes confirmed the presence of both Xpter and Xqter (B) and the deletion of Xpter (C), and duplication of the Xqter (D) in the son.
were found near the Xp22.33 breakpoint in the UCSC browser. Sequence homology has been observed near the breakpoints of recurrent intrachromosomal rearrangements. However, BLAST alignment of 100 kb segments flanking each breakpoint revealed no sequence identity greater than 1 kb in length.

**DISCUSSION**

We describe a mother and son with a strikingly similar syndrome of mesomelic skeletal dysplasia and abnormalities of the anterior segment of the eye associated with disruptions of the X chromosome. The mother's case was previously reported as a new syndrome [Kivlin et al., 1993]. The differential diagnosis of the mother's condition, as discussed in our previous report prior to the discovery of her X chromosome inversion, included Peters plus syndrome, acromesomelic dysplasia and LMD. However, the discovery of X chromosome abnormalities in these individuals with overlapping skeletal and ocular findings would strongly suggest an X-linked etiology. Mapping of the Xp22.3 and Xq27 breakpoints in these cases suggests a role for the **SHOX** gene in the skeletal dysplasia and for the **SOX3** gene in the ocular anomalies, respectively. The findings of severe mesomelic shortening preferentially affecting the upper extremities with short square ulnae and severely angulated radii in our patients resemble the upper extremity findings in LMD. LMD is caused by pseudautosomal recessive loss of the **SHOX** gene and presents with severe mesomelia of the lower limbs with hypoplastic fibulae and curved tibia. Our patients have less severe lower extremity defects, and also lack the severe rhizomelia often seen in LMD. Without sequencing the **SHOX** genes present on the normal X (in the mother) and Y (in the son) chromosomes, we cannot be certain that their skeletal phenotype does not represent a variant form of the recessive LMD. However, this possibility would require that both mother and son inherited additional **SHOX** mutations from each of their unrelated fathers who had normal height and normal limbs. We view this as an extremely unlikely explanation. Disruption of just one copy of **SHOX** would predict a phenotype resembling the pseudautosomal dominant Leri–Weil dyschondrosteosis (LWD). LWD presents with short stature and milder mesomelia of the upper extremities and Madelung deformity. The novel skeletal phenotypes seen in these patients, which affect the upper extremities preferentially like LWD yet lies closer in
A map of Xp22.33 showing all known genes located within 1.5 Mb from the Xpter. aCGH in the son showed deletion of a single BAC clone (RP11-839D20) that lies 240 kb distal to the SHOX gene. Five RT-PCR probes (P1, M1, M2, M3, S5') were designed spanning the interval between this BAC and the SHOX gene. 

B: Plot comparing real-time amplification of control and patient (the son) DNA using probe S5' (left) and probe M2 (right). Amplification of the patient DNA with the M2 probe reaches the threshold (arrow) at roughly one cycle count later than the control DNA relative to amplification of the S5' probe (at the 5' end of SHOX). Control S5' Ct = 22.48; control M2 Ct = 24.57; patient S5' Ct = 22.98; patient M2 Ct = 26.30. The relative ratio (RR) calculated using these Ct values is 0.43 indicating a deletion of the M2 probe in the DNA sample from the patient relative to the SHOX 5' probe. The analysis showed that three probes were deleted in the son (RR = 0.41, 0.50, and 0.43 for the P1, M1, and M2 probes, respectively, normalized to a control probe and compared to a random control individual), but that the 5' end of the SHOX gene was not disrupted (RR = 1.08 and 0.99 for the M3 and S5' probes, respectively).
severity to LMD, could suggest a novel disruption of the SHOX gene. Our finding that the coding region of the SHOX gene is, in fact, not disrupted, supports the hypothesis that misexpression of SHOX, due to a position effect, underlies this skeletal dysplasia. Our findings also suggest that this brachymesomelic skeletal dysplasia is allelic with LWD and LMD, as well as the other SHOX-related skeletal disorders including idiopathic short stature and the short stature of Turner syndrome. Finally, disruption upstream of the SHOX gene could explain a portion of LWD cases in which SHOX coding region mutations are not found, particularly those cases with more severe upper extremity involvement.

SHOX misexpression in these cases could involve either reduced expression due to loss of 5’ enhancers or ectopic expression due to gain of inappropriate enhancers or due to loss of SHOX repressors remains a possibility. The regulatory elements controlling SHOX expression have not been elucidated, but precedence for long-range regulation of the gene exists. Benito-Sanz et al. [2006] reported patients with LWD who harbor deletions in the 3’ region of the gene as far as 500 kb away. Precedence for overexpression causing defects preferentially in the upper extremities exists, as well, in the report by Grigelioniene et al. [2001], who described one patient with severe isolated Madelung deformity and a tandem duplication of the SHOX gene. However, disruptions 5’ of the SHOX gene associated with skeletal phenotypes have not been reported.

Peters anomaly of the eye and bilateral severe sclerocornea have not been reported with SHOX-related disorders. Peters anomaly of the eye and sclerocornea are related conditions in the spectrum of ASMD. Because ASMD is often associated with microphthalmia/anophthalmia (although not in this family), we considered the various loci for microphthalmia located on the X-chromosome as potentially causative. Two Xp-linked syndromes with ocular abnormalities include microphthalmia with linear skin defects (MLSD; MIM 300166), which involves more proximal Xp deletions at Xp22.31, and microphthalmia, cataracts, radiculomegaly and septal heart defects (MCOPS2; MIM 309801) that is caused by mutations in the BCOR gene at Xp11.2. Two Xq-linked disorders, Lenz microphthalmia (MCOP1; MIM 309800) and ANOP1 (MCOP4; MIM 301590), map to Xq27-28 but lie distal to the breakpoint in our patients. This lack of association between X chromosome abnormalities and anterior chambers defects suggested the possibility that a novel gene for Peters anomaly was disrupted at Xq27.1 in our patients. However, it is formally possible that the mother in this case harbors a mutation in one of the genes known to cause Peters anomaly (e.g., PAX6), which she also passed to her affected son, independent of their X-chromosome abnormalities. Sequencing of Peters anomaly genes has not been done.

Mapping of the Xq27.1 breakpoint by aCGH localized the breakpoint in these cases to an interval that lies approximately 90 kb 3’ of the SOX3 gene. No other genes lie within 350 kb. Deletions, duplications and point mutations of the SOX3 gene have all been reported and cause X-linked hypopituitarism with or without mental retardation [Laumonnier et al., 2002; Woods et al., 2005]. A large deletion-insertion occurring 67 kb downstream of the SOX3 gene is associated with X-linked recessive hypoparathyroidism in males [Bowl et al., 2005]. The patients in these reports had no ocular anomalies. However, disruptions of the closely related SOX2 gene are associated with anterior segment anomalies and microphthalmia [Fantes et al., 2003; Ragge et al., 2005]. Further, misexpression of Sox2 in the developing eyes of chick embryos causes abnormalities of the anterior chamber, and Sox2 and Sox3 are functionally similar in their ability to activate lens-specific genes like δ-crystallin [Kamachi et al., 2001; Kondoh et al., 2004]. Therefore, we hypothesize that misexpressed SOX3 could have led to the anterior chamber abnormalities in these patients, again due to a position effect.

The mother had partially skewed X-inactivation in lymphocytes. The SHOX gene is pseudoautosomal, and therefore X-inactivation is irrelevant for the phenotype resulting from its altered expression. On the other hand, the SOX3 gene is subject to inactivation, and both mother and son have a normal copy of this gene, present on the mother's normal X and on the duplicated segment of the recombinant chromosome in the son. Partially skewed X-inactivation in the mother may have contributed to the similarity between her clinical manifestations and those of her son, assuming that the altered allele is in the more frequently active X chromosome.

Blast analysis comparing the breakpoints in this mother and son revealed no homologous repeat DNA to explain the mechanism of inversion. The Xq27.1 breakpoint did occur within a 13 kb LINE element cluster. However, this breakpoint region appears to be distinct from those reported in other disruptions near the SOX3 gene [Bowl et al., 2005; Woods et al., 2005].

In summary, our identification of X chromosome disruptions in this mother and son redefines this syndrome of brachymesomelic skeletal dysplasia with Peters anomaly of the eye. The breakpoints involved lie near genes plausibly related to the observed spectrum of defects (Fig. 6), leading us to speculate that misexpression of both SHOX and SOX3, due to position effects, causes the observed skeletal and ocular abnormalities, respectively. Analysis of these genes in patients with either similar skeletal dysplasias, or anterior segment ocular abnormalities, may confirm this hypothesis.
In particular, we put forward the \textit{SOX3} gene as a possible candidate gene for isolated Peters anomaly and other anterior segment ocular abnormalities.

\section*{ACKNOWLEDGMENTS}

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