

ORIGINAL ARTICLE

Breakpoints around the *HOXD* cluster result in various limb malformations

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Supplementary data are available at <http://www.jmedgenet.com/supplemental>

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Background: Characterisation of disease associated balanced chromosome rearrangements is a promising starting point in the search for candidate genes and regulatory elements.

Methods: We have identified and investigated three patients with limb abnormalities and breakpoints involving chromosome 2q31. Patient 1 with severe brachydactyly and syndactyly, mental retardation, hypoplasia of the cerebellum, scoliosis, and ectopic anus, carries a balanced t(2;10)(q31.1;q26.3) translocation. Patient 2, with translocation t(2;10)(q31.1;q23.33), has aplasia of the ulna, shortening of the radius, finger anomalies, and scoliosis. Patient 3 carries a pericentric inversion of chromosome 2, inv(2)(p15q31). Her phenotype is characterised by bilateral aplasia of the fibula and the radius, bilateral hypoplasia of the ulna, unossified carpal bones, and hypoplasia and dislocation of both tibiae.

Results: By fluorescence in situ hybridisation, we have mapped the breakpoints to intervals of approximately 170 kb or less. None of the three 2q31 breakpoints, which all mapped close to the *HOXD* cluster, disrupted any known genes.

Conclusions: *Hoxd* gene expression in the mouse is regulated by *cis*-acting DNA elements acting over distances of several hundred kilobases. Moreover, *Hoxd* genes play an established role in bone development. It is therefore very likely that the three rearrangements disturb normal *HOXD* gene regulation by position effects.

The *HOX* genes code for a family of highly conserved transcription factors expressed during embryonic development. To date, mutations in four *HOX* genes, namely *HOXA11*, *HOXA13*, *HOXD10*, and *HOXD13*, have been found and all are associated with limb malformations. In particular, it has been shown that in two families a single nucleotide deletion within the second exon of *HOXA11*, which results in a frameshift and a premature stop codon, co-segregates with proximal radial-ulnar synostosis.¹ Mutations in *HOXA13* cause hand-foot-uterus syndrome, a rare dominantly inherited condition (OMIM 140000),² and Guttmacher syndrome (OMIM 176305).³ Recently, a missense mutation in the *HOXD10* gene has been described as the cause of isolated congenital vertical talus, also known as rocker-bottom feet (OMIM 192950), and Charcot-Marie-Tooth disease (OMIM 118220).⁴ The first human limb malformation shown to be caused by *HOXD13* polyalanine tract expansion mutations was synpolydactyly (SPD, OMIM 186000).⁵ Similar pathological polyalanine tract expansions in *HOXD13* have subsequently been reported in other families with SPD.^{6–8} Intragenic frameshift deletions in *HOXD13*, predicted to result in truncated proteins, and also an acceptor splice site mutation and a missense mutation in exon 2 of *HOXD13*, cause an atypical form of SPD.^{9–12} Interestingly, a different missense mutation in the same exon has been found in a family with a dominantly inherited combination of brachydactyly and polydactyly.¹³

Large deletions involving chromosome 2q31.1 have been associated with minor digital anomalies,^{14–15} with major limb defects,^{15–17} or with a combination of severe limb and genital

abnormalities.¹⁸ In contrast, a microdeletion at the 5' end of the *HOXD* cluster that removes *HOXD9* to *HOXD13* and extends 85 kb upstream of *HOXD13* results in an SPD phenotype.¹⁹

Another relevant point to consider is the presence of regulatory mutations affecting gene expression. Spitz *et al* described a balanced t(2;8) translocation that co-segregated with mesomelic dysplasia and vertebral defects.²⁰ The authors mapped the chromosome 2 breakpoint approximately 60 kb downstream from the *HOXD* cluster. Since no known gene is disrupted by the rearrangement, the authors speculated that the translocation affects regulatory sequences of the *HOXD* cluster, resulting in misregulation of *HOXD* gene expression.

In this paper, we report our findings on three unrelated patients with various skeletal malformations and balanced rearrangements involving chromosome band 2q31.

METHODS

Subjects

Proband 1 is a 13 year old boy carrying an apparently balanced chromosome rearrangement t(2;10)(q31.1;q26.3) (schematically depicted in fig 1A). He is the second child of healthy unrelated parents, born after a normal pregnancy and without fetal distress. Clinical examination after birth revealed severe malformation of the hands and feet as well as a dysmorphic face. The hands were short and had six fingers, and absence of the distal phalanges including the nails was

Abbreviations: FISH, fluorescence in situ hybridisation; SPD, synpolydactyly

noted. The feet showed complete absence of toes II–V, and a rudimentary first toe with a missing distal phalanx. x Rays of the hands showed five short metacarpals and six digits, each consisting only of a single phalanx. Metacarpal III was bifurcated at its end, giving rise to two digits. The patient was not able to fully extend his elbows and knees due to contractions. Bilateral inguinal hernias were noted and surgically corrected. x Rays of the thorax showed hypoplasia of the medial ends of both clavicles. Ultrasound of the skull revealed hypoplasia of the cerebellum. Further clinical examinations demonstrated developmental delay, deep set eyes (left more than right), a progressive scoliosis, narrow shoulders, ataxia, coxa valga, and short stature (110 cm at age 6, <3rd percentile) (fig 2A–D). The hands of the patient were surgical corrected and the polydactylous digits removed. The patient has developed well and is currently attending a special school.

Proband 2 is a 23 year old female carrying a balanced de novo translocation $t(2;10)(q31.1;q23.33)$ (fig 1B). She is the first child of healthy parents of Icelandic origin, born at term after an uncomplicated pregnancy. Except for her extremity malformation, her motor, mental, and social development were normal. She has an almost symmetrical limb phenotype consisting of ulnar aplasia, radial shortening, and absence of 3rd to 5th rays (fig 2E). There is hypoplasia of digits 1 and 2 where one epiphysal disc is shared by the two proximal phalangeal bones causing articulation between the fingers. Only one carpal bone is present. Humerus and lower limbs are normal. A slight dextro-convex scoliosis was found at the age of 23 (fig 2F). Her mammary glands are normal.

Proband 3 is a newborn girl with a balanced de novo pericentric inversion $inv(2)(p15q31.1)$. She is the first child of unrelated parents, a 30 year old mother and a 42 year old father of German origin. She was born after an uncomplicated pregnancy in the 40th week of gestation (birth weight 3700 g, length 49 cm, head circumference 37.5 cm). There was no history of any medication or illness during pregnancy. The first radiological examination after birth revealed bilateral aplasia of the fibula and the radius, bilateral hypoplasia of the ulna, unossified carpal bones, and hypoplasia and dislocation of both tibiae.

Breakpoint mapping by fluorescence in situ hybridisation (FISH)

YAC clones from the regions of interest were used for the initial mapping of the breakpoints, followed by fine mapping with smaller clones. FISH was carried out according to standard procedures. Genomic clones were labelled with digoxigenin-dUTP or biotin-dUTP and were hybridised to patient metaphase chromosomes. Signals were detected either by anti-digoxigenin, FITC, or Cy3 conjugated avidin and were visualised by fluorescence microscopy with a CCD

camera (Sensys, Photometrics, Tucson, AZ) and an image analysis program (IPLAB spectrum, Vysis, Downers Grove, IL). The spotted human chromosome 2-specific cosmid library (Livermore) was hybridised with a pool of PCR products selected from the breakpoint region of patient 1. Positive cosmids were obtained as bacterial stocks from the Deutsche Ressourcenzentrum für Genomforschung (RZPD). DNAs were isolated with the Miniprep Kit (Qiagen, Hilden, Germany), and their ends were sequenced with T7 and T3 universal primers.

Southern blotting and breakpoint cloning

Genomic DNAs isolated from patient 1 and control cell lines were digested with *Bam*HI, *Bgl*II, *Hind*III, or *Pst*I restriction enzymes, separated in 1% agarose gels, transferred onto nylon membranes (Roth, Karlsruhe, Germany), and hybridised with [α^{32} P]-dCTP labelled PCR products.

Probes for Southern blot hybridisation were amplified using the following primer sets: 538A12_49750for (5' GCT TCC CAT TGC AGG TGT AAA 3') and 538A12_49750rev (5' ATT ACT GGT CAT CAA TAT CTA GC 3'), 538A12_79400for (5' AAC TCA ACA TAAACT TTT CCA AAG 3') and 538A12_79400rev (5' GAA TGT AAA ATA TAG ACA TTT GAC ATT G 3') (positions 104665–105358 and 110085–110642 of clone RP11-538A12, GenBank accession no. AC016761).

Breakpoint cloning was performed using adaptor ligated PCR as described elsewhere.²¹ *Eco*RI digested patient DNA was PCR amplified with a set of nested primers, AP1 (5' GGA TCC TAA TAC GAC TCA CTA TAG GGC 3') with 84364for (5' CAG ATT GTG ATT AGA TCA GGA G 3') and AP2 (5' TAT AGG GCT CGA GCG GC 3') with 84715for (5' GAC TTA AAA TTG CAG CGT GTG TTT C 3') for der(10), and AP1 with 85206rev1 (5' GTG TAT CTA TCT GAG CTC CAT G 3') and AP2 with 85163rev2 (5' TTC AGC CTT AAG TCA AAA TGT TGG 3') for der(2). Amplified fragments were isolated from 1% agarose gels, subcloned into pGEM-T Easy vector and sequenced using M13 universal primers.

Screening for *HOXD13* mutation in patient 1

Mutation analysis of the *HOXD13* gene in patient 1 included all coding exons and splice sites. Primers used for PCR amplifications are available online as supplementary data available at <http://www.jmedgenet.com/supplemental>.

Computational analysis of the breakpoint regions

Breakpoint regions were analysed in silico with the NIX program to identify known or predicted genes and ESTs. All putative ESTs were screened against entries in GenBank and TIGR databases.

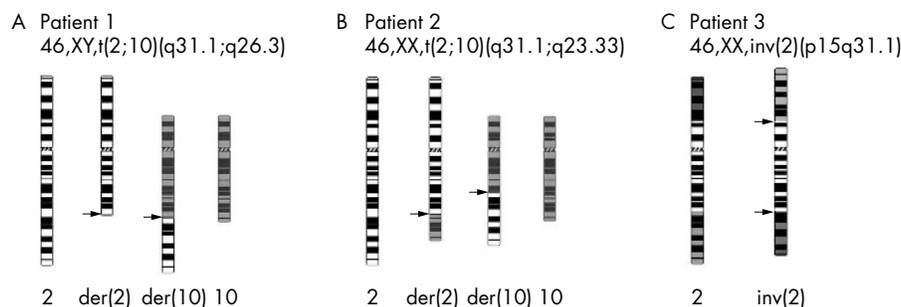


Figure 1 Karyotypes of the three patients with apparently balanced chromosome rearrangements and limb malformations included in this study, and ideograms of the rearranged chromosomes. Arrows point to the breakpoints on der(2), der(10), and inv(2) chromosomes.



Figure 2 (A,B) Pictures of patient 1 at the age of 10. (C) Photo of the right hand of patient 1. (D) x Ray of the right hand of patient 1. Note the shortened metacarpals, webbing, duplications, and shortening of digits; (E) x Ray of the left arm of patient 2 at the age of 10 demonstrating ulnar aplasia, shortened radius, and absent 3rd to 5th rays. Furthermore, there is hypoplasia of digits 1 and 2. (F) x Ray of the spine of patient 2 at the age of 23 revealing a slight dextro-convex scoliosis. (G) Babygram of proband 3. Note severe defects in distal limbs.

RESULTS

Mapping of the chromosome 2q31 breakpoints by FISH

FISH performed on patients' metaphase chromosomes showed that for patient 1, BAC clone RP11-514D19

(GenBank accession number AC016915), which contains *EVX2* and *HOXD8-13*, showed signals on chromosome 2 and on der(10), indicating that the breakpoint lies outside this region (data not shown). Further hybridisation of a series of BAC clones selected from the region proximal to *EVX2* led to the identification of the breakpoint-spanning clone RP11-538A12 (GenBank accession number AC016761) (fig 3A,D). To map the breakpoint more precisely, PCR products generated with primers selected from clone RP11-538A12 were used for screening a chromosome 2-specific cosmid library. Positive clones were hybridised to the patient chromosomes, and clone LLNLc128F0946 showed a signal on der(10), whereas clone LLNLc128A0237, with a partially overlapping sequence, showed a signal on der(2) (fig 3D).

For patient 2, YAC 935E10 (with marker D2S138 and located at 192 cM) was found to span the 2q31 breakpoint (data not shown). Thus, the breakpoint of patient 2 was different from and more distal on chromosome 2q than the breakpoint of patient 1. This was confirmed by the FISH results obtained with BAC clone RP11-387A1 (GenBank accession number AC009336), containing *EVX2* and the complete *HOXD* cluster, which gave signals on chromosome 2 and on der(2) (data not shown). Hybridisation of BAC clone RP11-25L17 (GenBank accession number AC096657) showed split signals on der(2) and der(10), indicating that this clone spans the breakpoint (fig 3B,D). Furthermore, BAC clones RP11-1085F24 and RP11-28M17 were also found to be breakpoint spanning (data not shown). All three breakpoint-spanning BACs have overlapping sequences, with a common region of approximately 22 kb. Therefore, it is most likely that the breakpoint is located within this segment.

For patient 3, YAC clone 888B4 (with markers D2S148 and D2S2173 and located at 190 cM) was found to span the chromosome 2q31 breakpoint (fig 3C), indicating that this breakpoint is different from the two other breakpoints and lies in the region between the *HOXD* cluster and the breakpoint of patient 2. Further FISH studies showed that BAC clone RP11-724O12 (GenBank accession number AC079803) spans the breakpoint (data not shown).

In conclusion, FISH mapping revealed that the chromosome 2q31 breakpoints investigated in the three patients with limb malformations lie outside the *HOXD13*: in patient 1 approximately 390 kb centromeric to *HOXD13*, in patient 2 approximately 1050 kb telomeric to *HOXD13*, and in patient 3 approximately 590 kb telomeric to *HOXD13*.

A similar approach was used to map the breakpoints on chromosome 10. In patient 1, BAC RP11-300B2 (GenBank accession number AL355531) was breakpoint spanning. In patient 2, we were able to narrow the region to approximately 28 kb, between the clones RP11-81C11 and RP11-702G14 and clone RP11-348J12 (GenBank accession number AL358613) (data not shown).

For patient 3, the breakpoint on the short arm of chromosome 2 was narrowed to a region of approximately 170 kb between clones RP11-351H12 (GenBank accession numbers AQ529149 and AQ529148) and RP11-355A23 (GenBank accession number AC009490) (data not shown).

Cloning of the breakpoints of patient 1

In order to localise precisely the chromosome 2 breakpoint of patient 1, we performed Southern blot hybridisation of patient and control DNAs digested with appropriate restriction enzymes. Since there is an approximately 4.5 kb long LINE repeat in the region of interest, we were very restricted with respect to designing probes. The two probes designed from sequences close to the repeat were most informative. Hybridisation of the more telomeric probe (538A12_49750) allowed observation of aberrant restriction fragments in patient DNA digested with *Bam*HI, whereas the more

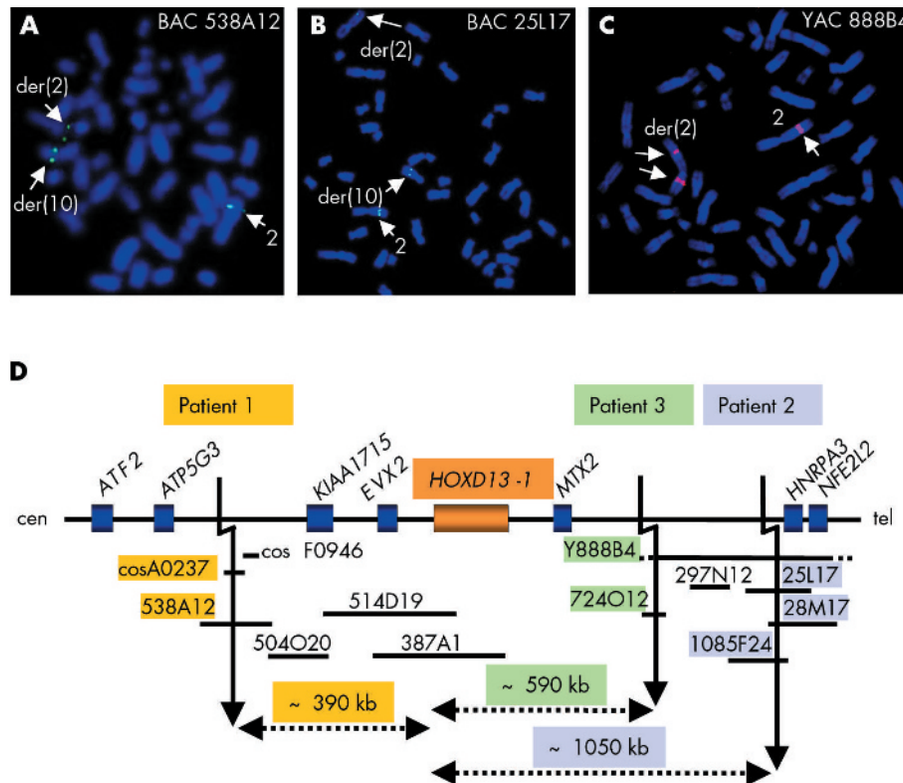


Figure 3 FISH mapping of the 2q31 breakpoints in the patients. (A) Hybridisation of the breakpoint spanning clone RP11-538A12 to patient 1 chromosomes; arrows indicate signals on chromosome 2, der(2), and der(10). (B) FISH results for patient 2 with RP11-25L17, the breakpoint spanning BAC; arrows indicate signals on the normal and the derivative chromosomes. (C) Metaphase showing the breakpoint spanning YAC 888B4 in patient 3 with signals on the normal chromosome 2 and split signals on the inverted chromosome 2. (D) Schematic map of the 2q31.1 band showing the location of the three breakpoints in the vicinity of the *HOXD* complex together with breakpoint-spanning BAC and cosmid clones; known genes from this region are depicted schematically. Approximate distances from the breakpoints to the *HOXD13* gene are indicated by double headed arrows (not drawn to scale).

centromeric probe (538A12_79400) showed aberrant restriction fragments in patient DNA cut with *Bgl*III (fig 4A,B). Mapping of the breakpoint on chromosome 10 gave us additional information about restriction sites on der(2) and der(10) chromosomes, and we found that *Eco*RI should be useful for our further analyses. In order to clone the breakpoint of chromosome 2, we ligated adaptors to *Eco*RI digested patient DNA. Subsequent PCR reactions gave rise to two products, approximately 500 bp long for der(10) and approximately 1.1 kb long for der(2), which were subcloned and sequenced. The breakpoint on chromosome 2 is located between positions 107 910 bp and 107 912 bp of clone RP11-538A12. The nucleotide G at position 107 911 bp is deleted; moreover, on der(2), a 9 bp insertion is present between original chromosome 2 and chromosome 10 sequences (fig 4C).

***HOXD13* is not mutated in patient 1**

The limb malformations of patient 1 resemble those present in patients with SPD and a mutation in *HOXD13*. Our mutation analysis of *HOXD13* showed normal results. Hence, the phenotype is most likely caused by the translocation.

Computational analysis of the breakpoint regions

In silico analysis showed that there is no known gene on the chromosome 2q31 breakpoint-spanning BAC (RP11-538A12) in patient 1. For patient 2, we identified no known gene in the 22 kb long breakpoint interval; the closest gene (*hmRNP3*) is located approximately 45 kb telomeric to the breakpoint region. For patient 3, sequence analysis of the breakpoint-spanning BAC (RP11-724O12) revealed the

presence of three short unspliced ESTs, all derived from either cancer cells or stomach tissues, but none of these ESTs belongs to any known gene.

Our analysis of the chromosome 10 breakpoint region of patient 1 revealed that this breakpoint disrupts the *MGMT* gene, which encodes methylguanine-DNA methyltransferase, an enzyme involved in DNA repair processes. *MGMT* is expressed from the intact chromosome 10 in the patient cell line (data not shown); thus, the breakpoint on chromosome 10 may play no role in the phenotype.

The breakpoint on chromosome 10 in patient 2 was mapped within a 28 kb region of the breakpoint-spanning BAC clone RP11-348J12. This region contains three genes: *SEC15L1* and two cytochrome P450 genes, *CYP26A1* and *CYP26C1*. The product of the human *SEC15L1* gene is highly similar to the yeast protein SEC15, which is one of the components of a multiprotein complex required for exocytosis. To date, there is no indication that *SEC15L1* plays a role in limb development. CYP26 family members of the cytochrome P450 catabolise and contribute to the control of retinoic acid, an important regulator of gene expression during embryonic development.^{22, 23} The *Cyp26a1*^{-/-} knockout mice show caudal truncation, vertebrae transformation, and hindbrain mispatterning. Moreover, in these mice the hindlimbs are fused, whereas no abnormalities of the upper limb structures are visible.²⁴ The absence of lower limb malformations in patient 2 suggests that *CYP26A1* is not involved in the observed upper limb malformations.

The breakpoint region on the short arm of chromosome 2 in patient 3 has been mapped to an interval of approximately 170 kb. This region contains no known gene.

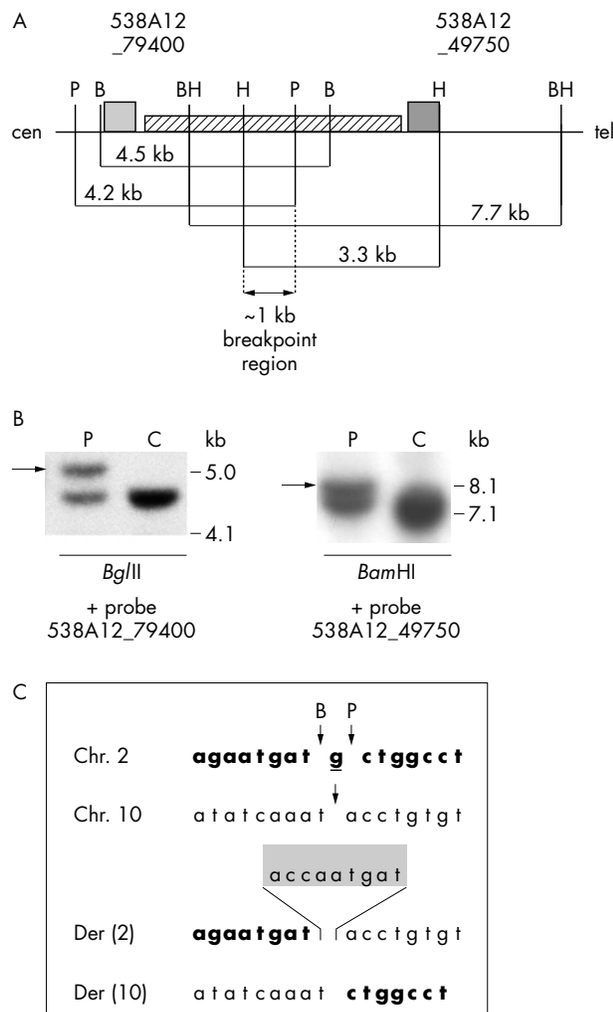


Figure 4 (A) Schematic representation of the breakpoint region on BAC RP11-538A12 and the localisation of probes 538A12_79400 and 538A12_49750 (shown as grey boxes) used for Southern blot hybridisation. The positions and sizes of normal restriction fragments are indicated and enzymes are marked as follows: B, *Bgl*II; BH, *Bam*HI; H, *Hind*III; P, *Pst*I; the striped box between the two probes indicates the LINE repeat. (B) Southern blot analysis with probe 538A12_49750; DNA from patient 1 (P) and a control (C) were digested with *Hind*III or *Bam*HI restriction enzymes; the presence of additional bands in the patient compared to the control (arrows) indicates that the breakpoint is located within the *Hind*III and *Bam*HI restriction fragments. (C) Chromosome 2, 10, der(2), and der(10) sequences in patient 1. Chromosome 2-derived sequences are shown in bold; the underlined guanine residue from chromosome 2 is missing in both derivative chromosomes. The 9 bp insertion of unknown origin within der(2) (shaded box) separates chromosome 2- and chromosome 10-derived sequences.

DISCUSSION

In the present study we have investigated three unrelated patients with severe limb malformations and apparently balanced chromosome rearrangements involving chromosome 2q31 and located close to the *HOXD* complex. It is noteworthy that computational analysis of each breakpoint region in 2q31 provided no indication that any known genes are directly truncated by the breakpoints. However, the patients show phenotypes characteristic for *HOXD* mutations which strongly suggests that the limb abnormalities are the result of disturbed *HOXD* expression through the chromosomal breakpoints rather than a coincidence of affected limb development by other mechanisms.

In patient 1, the phenotype resembles that of patients with SPD. The most common mutation causing SPD is an expansion of the imperfect trinucleotide repeat coding for alanine in the *HOXD13* gene. Sequence analysis of the *HOXD13* genes in patient 1 showed a normal result, which suggests that the phenotype is instead caused by effects of the translocation. It has been proposed that SPD is not only caused by a loss of function of the *HOXD13* gene but also by a combined loss of function of several *HOXD* genes. This conclusion has been drawn from experiments done in the mouse. *Hoxd13* homozygous knock-out mice show a fairly mild phenotype^{25, 26} which resembles neither that of patients with SPD nor that of *spd*h mice.²⁷ Interestingly, mice with a deletion encompassing *Hoxd11*, *Hoxd12*, and *Hoxd13* show a much more severe limb phenotype, comparable to that of the homozygous *spd*h mouse.²⁸ Results obtained by several groups indicate that the posterior genes, *Hoxa13*, *Hoxd11*, *Hoxd12*, and *Hoxd13*, contribute significantly to digit development, and that these genes control this process in a dose-dependent manner, whereas *Hoxa11* and *Evv2*, the latter located at the centromeric end of the *Hoxd* complex, showed only a minor contribution to digit morphogenesis.²⁹ We propose, therefore, that in patient 1 the translocation has an impact on the regulation of *HOXD* expression, which results in the *HOXD* loss of function phenotype. Aberrant gene expression caused by a change of the gene's chromosomal environment, referred to as a position effect, can be the result of various mechanisms, one of which is the separation of regulatory elements from their respective transcription unit, leading to misexpression (reviewed by Kleinjan and van Heyningen^{30, 31}). Intensive studies in the mouse led to the hypothesis that *Hoxd10*, *Hoxd11*, *Hoxd12*, and *Hoxd13* genes, which show very similar expression domains in presumptive digits,³² are under the control of the same digit enhancer that controls their spatial and temporal expression in developing limbs. Furthermore, it has been suggested that this digit enhancer is located centromeric to the *Hoxd* complex.³³ Many attempts have been made to find this enhancer sequence,^{34, 35} but its precise location remains unknown. Introduction of a human PAC clone covering almost the entire *HOXD* cluster and extending approximately 40 kb centromeric to *HOXD13* could not rescue the phenotype in mice which were triple mutants for *Hoxd11/Hoxd12/Hoxd13*, suggesting that the enhancer is located even further away from the cluster.³⁶ More recently, a fragment of human DNA from the region more centromeric to the *HOXD* cluster has been shown to contain transcriptional enhancer activity similar to that controlling both *Hoxd* and *Evv2* genes.³⁷ This DNA segment is part of human BAC clone RP11-504O20 (GenBank accession number AC016751), which partially overlaps with BAC 538A12, the clone spanning the breakpoint in patient 1. Spitz *et al*³⁷ have also analysed BAC 538A12 but could not show any enhancer activity specific for posterior *HOXD* genes. It is thus very likely that the presumed digit enhancer lies close to the breakpoint in patient 1; however, it seems to be neither disrupted nor separated from the *HOXD* cluster by the breakpoint.

In patients 2 and 3, the 2q31 breakpoints are located on the telomeric side of the *HOXD* cluster, and both patients show malformations of more proximal parts of the limbs. In patient 2, the ulnae and the posterior digits in the hands are absent while the radii are hypoplastic. Patient 3 has no radii, ulnae, or fibulae, and shows hypoplastic tibiae. Improper development of the lower arm and lower legs has been described in mice lacking either *Hoxd11* or *Hoxa11* genes, as well as in the double mutants.³⁸⁻⁴² Additionally, in *Hoxd11* knock-out mice, metacarpals, phalanges, and wrist bones are affected.^{39, 40} Compound mutants for *Hoxa10* and *Hoxd11*, as well as *Hoxa10* and *Hoxd10* double knock outs, showed more severe

phenotypes, suggesting that, as for digit formation, zeugopod development is dependent on the correct expression of several posterior *Hoxd* genes.^{43–44} In addition to the knock-out mice, a regulatory mutation in the mouse has been described. *Ulnaless* is a semidominant x ray induced mutation, which changes the level of posterior *Hoxd* expression in limb buds.^{33–35} The phenotype is very severe, with affected zeugopods and almost complete absence of ulnae, which resembles the phenotypes of patients 2 and 3. It has recently been shown that an inversion occurred on chromosome 2 in the *ulnaless* mouse, near the *Hoxd* cluster. The size of the inversion is approximately 770 kb, with one breakpoint centromeric to the *Hoxd* cluster disrupting the *Lnp* gene, and the other breakpoint, which probably does not truncate any gene, telomeric to the *Hoxd* cluster.³⁷ Although the authors highlighted the relevance of the centromeric breakpoint for the *ulnaless* phenotype, our results for patients 2 and 3, together with a recent hypothesis about the early limb control region present on the telomeric site of the *Hoxd* cluster in mouse,⁴⁶ suggest that the *ulnaless* breakpoint telomeric to *Hoxd* may also play a role in the observed zeugopod malformations. In line with this is the translocation t(2;8) described by Spitz *et al*,²⁰ which is also associated with abnormal zeugopod development. In this translocation, the chromosome 2 breakpoint is located closer to the *HOXD* cluster than the breakpoint in patient 3 described in the present study. Similarly to the patients described in this study, however, no obvious gene was disrupted by the rearrangement. These findings indicate that sequences telomeric to the *HOXD* genes also play an important role in the regulation of the entire cluster. In light of this, it is likely that the limb phenotypes in patients 2 and 3 are the result of aberrant *HOXD* expression through the breakpoints telomeric to the *HOXD* cluster.

Other position effect mechanisms can also lead to gene silencing, either by insertion of a gene into a heterochromatic region or by removal of a long range insulator, which normally allows spreading of heterochromatin over long distances. However, it is rather unlikely that one of these mechanisms has caused misregulation of *HOXD* genes in patient 1, since the whole *HOXD* cluster and all currently known regulatory elements have been translocated into a transcriptionally active region at 10q26 containing the *MGMT* gene. Similarly, in patient 2, the breakpoint at 10q23.33 lies within a region containing several genes. Therefore, the phenotypes of these patients are more likely caused by translocation mediated disturbances in other mechanisms; for instance, the rearrangements might have brought the *HOXD* cluster into the vicinity of regulatory elements present near the second breakpoint. Alternatively, another transcription unit from the non-*HOXD* breakpoint regions could compete with the *HOXD* genes for the digit enhancer or for the early limb control region, resulting in misregulation of the *HOXD* genes. Both hypotheses are interesting, however given the present state of knowledge, they are only speculative. At the moment, nothing is known about regulatory elements of *MGMT* or other genes in the vicinity of the breakpoint at 10q26 (*EBF3* and *TXNL2* are located approximately 330 kb and 630 kb telomeric to the chromosome 10 breakpoint in patient 1, and *MKI67* is located approximately 1.4 Mb centromeric to this breakpoint), which could theoretically influence expression of the translocated *HOXD* cluster. Moreover, nothing is currently known about the presence of putative *cis*-acting regulators in the breakpoint regions at 10q23 and 2p15 in patients 2 and 3.

Expression of posterior *HOXD* genes in limb buds is most likely regulated by the interplay between *cis*-acting elements and *trans*-acting factors. To date, two groups of genes, namely *Polycomb* (*PcG*) and *trithorax* (*trxG*) have been implicated in

maintenance of the active or silent state of *Hox* genes in *Drosophila* (for review see Simon⁴⁷) and mammals (reviewed in Schumacher and Magnuson⁴⁸). Results from the last few years suggest that *PcG* genes may also play a role in controlling expression of posterior *Hoxd* genes in mouse limb buds by integrating both local and global regulatory mechanisms.^{49–50} However, it has been shown that binding of different *trans*-acting regulators to DNA might be dependent on intact chromatin architecture.^{51–52} Since it has been proposed that the chromatin structure of any locus is determined by the combination of *cis*-acting elements and the wider chromosomal and nuclear environment,⁵¹ it is plausible that chromosome rearrangements could influence chromatin architecture and thereby result in misregulation of genes. In fact, changes in chromatin structure have been proposed following insertion of some transgenes,⁵³ or in the case of small deletions.⁵⁴

Interestingly, changes in the global chromatin structure might also influence expression of other genes near the breakpoints at 2q31. The closest gene, located only 220 kb telomeric to the breakpoint in patient 1, is *KIAA1715*. This gene is transcribed from the strand opposite that of the *HOXD* cluster. Its mouse homologue, *Lnp*, shows the same expression pattern in limb buds and external genitalia as *Hoxd13* and *Evx2*, which suggests that all three genes are under the control of the same regulatory sequences. In addition, *Lnp* is also expressed in the developing central nervous system in a highly similar pattern to that of *Evx2*, and it has a specific expression domain in the eyes, the heart, and the forebrain.³⁷ The neural enhancer that may activate *KIAA1715* is located in part within the same 40 kb region as the digit enhancer mentioned earlier; therefore, it is possible that both the limb and the neuronal expression domains of *KIAA1715* have been affected by the translocation via a position effect. Since patient 1 has cognitive deficits in addition to limb abnormalities, it is tempting to link the central nervous system phenotype with disturbed expression of the *KIAA1715* gene.

Although theoretically possible, it seems rather unlikely that the breakpoints on chromosome 10 and on the short arm of chromosome 2 cause the phenotypes of the patients reported here. The breakpoint at 10q26 in patient 1 disrupts the *MGMT* gene encoding the DNA-repair enzyme methylguanine-DNA methyltransferase; however, we observed expression of this gene from the intact chromosome 10 in patient lymphoblastoid cell lines. Moreover, *MGMT*^{-/-} knock-out mice are essentially normal,⁵⁵ apart from being somewhat smaller than controls.⁵⁶ For patient 2, three genes, *SEC15L1*, *CYP26A1*, and *CYP26C1*, reside on the chromosome 10-derived breakpoint-spanning clone RP11-348J12; however, misexpression of none of these genes could explain her phenotype. In patient 3, the breakpoint in the short arm of chromosome 2 was mapped to an interval of approximately 170 kb. This interval and also its flanking DNA stretches, including 600 kb of 3' sequence and 500 kb of 5' sequence, harbour no known genes.

In summary, the data presented in this paper are plausible examples of regulatory mutations underlying severe developmental defects. They illustrate the complexity of *HOXD* cluster regulation and highlight the importance of the precise regulation of these genes for proper limb development.

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