Analysis of a terminal Xp22.3 deletion in a patient with six monogenic disorders: implications for the mapping of X linked ocular albinism

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Abstract
The molecular characterisation of chromosomal aberrations in Xp22.3 has established the map position of several genes with mutations resulting in diverse phenotypes such as short stature (SS), chondrodysplasia punctata (CDPX), mental retardation (MRX), ichthyosis (XLI), and Kallmann syndrome (KAL). We describe the clinical symptoms of a patient with a complex syndrome compatible with all these conditions plus ocular albinism (OA1). He has a terminal Xp deletion of at least 10 Mb of DNA. Both the mother and sister of the patient are carriers of the deletion and show a number of traits seen in Turner's syndrome. The diagnosis of ocular albinism was confirmed in the patient and his mother, who shows iris translucency, patches and streaks of hypopigmentation in the fundus, and macromelanosomes in epidermal melanocytes. By comparative deletion mapping we can define a deletion interval, which locates the OA1 gene proximal to DXS143 and distal to DXS85, with the breakpoints providing valuable starting points for cloning strategies.

Interstitial and terminal deletions in Xp22.3 are associated with monogenic disorders such as short stature, chondrodysplasia punctata, mental retardation, X linked ichthyosis, and Kallmann syndrome, which can occur singly or together. Male patients with deletions in Xp22.3 are nullisomic for this region and therefore they show various phenotypes according to the length of the deletion involved. While genes associated with X linked ichthyosis, and Kallmann syndrome have been cloned, those for chondrodysplasia and mental retardation have not. Patients manifesting contiguous gene syndromes, including ocular albinism (OA1), have been published, but these reports lack either hard clinical evidence or clear molecular data.

The physical map of Xp22.3 comprises more than nine million base pairs (Mb), including the whole of the pseudoautosomal region. A large number of deletions and translocations have been assembled in order to narrow cloning intervals for genes in this chromosomal region. However, the gene position for ocular albinism remains unresolved. Two alternative orders have been suggested: the putative OA1 locus has been placed distal and proximal to the locus DXS143 by linkage analysis.

Here we describe a patient with a large Xp22.3 deletion who has ocular albinism in addition to short stature, chondrodysplasia punctata, mental retardation, ichthyosis, and Kallmann syndrome. Clinical investigation of the family was followed by cytogenetic analysis and comparative deletion mapping.

Methods
PATIENTS
A detailed clinical description of the patient (W1) with symptoms of mental retardation, ichthyosis, and Kallmann syndrome was reported by Pike et al. A recent ophthalmological examination has shown no signs of ocular albinism (normal vision, normal fundus, no nystagmus). Growth assessment was performed according to tables published in the Netherlands.

CYTOGENETIC ANALYSIS
High resolution chromosomal analysis of peripheral lymphocytes of the patient BK and his mother were performed according to Yunis using amethopterin as the synchronising agent and Giemsa-Trypsin banding. The inactivation pattern of the X chromosome in the mother was investigated using bromodeoxyuridine as the synchronising agent and fluorochrome-photolysis-Giema staining.

DNA ANALYSIS
DNA extraction, Southern blotting, and pulsed field gel electrophoresis (PFGE) were carried out as described previously. DNA probes used for hybridisation are summarised in the table.

Case reports
The patient investigated in this study (BK) is 9 years old and shows signs of a complex syndrome involving the eyes, brain, skeletal system, and skin. He was born after two miscarriages as the second child of unrelated parents. He has two healthy brothers and a sister.
Placental insufficiency had been assumed in the 37th week of gestation after measuring low levels of oestadiol. Delivery was spontaneous in the 40th week of pregnancy. Hypertriglyceridaemia and a large heart were noted soon after birth.

Growth parameters measured were far below normal levels. Weight at birth was 2400 g (mean = 4 SD), length 40 cm (mean = 5 SD), and head circumference 32 cm (mean = 3 SD). The patient’s height at the age of 8½ years was 102 cm (mean = 5 SD), the upper/lower ratio was 1:4 (normal 1:01), and the head circumference was 51.5 cm (mean = 1.5 SD).

Dysmorphic features suggested the diagnosis of chondrodysplasia punctata which was confirmed soon after birth by radiographs showing multiple stippled calcifications of the epiphyses, predominantly in the paravertebral region and the neighbourhood of the carpals and tarsal bones. Metacarpophalangeal pattern profile analysis (at the age of 5) showed shortening of metacarpals 1 to 4 and pronounced brachyteleactyly. Ulnar deviation of the wrist was noted which was caused by hypoplasia of the distal ulna. Dysmorphic features of the face included a broad, flat nose with anteverted nares and a small lateral groove, short columella, pouting upper lip, and hypoplasia of the maxilla. Lateral x-ray of the skull at the age of 15 months confirmed the maxillonasal dysplasia and absence of the anterior nasal spine, the ‘Binder phenotype’:30,31 Psychomotor retardation became evident at the age of 3 months. He started to stand at 3½ years and to walk independently after his sixth birthday. He did not learn to speak. He recognises persons familiar to him and he shows pleasure at their touch.

Ichthyosis was diagnosed at the age of 5 and hyperkeratosis was fully developed at the age of 7. At the age of 8 years, Kallmann syndrome was assumed because of microgenitalia and cryptorchidism in addition to mirror movements. His sense of smell could not be assessed.

Ocular albinism was diagnosed after an ophthalmological investigation under anaesthesia at the age of 7 months. A poorly pigmented fundus, aplasia of the macula and choroid, and a horizontal nystagmus were noted. Iris translucency, ‘mud splattered fundus’, and a reduction of pigmentation epithelium confirmed the carrier status of his mother (fig 1A,B). This diagnosis was confirmed by showing the presence of macromelanosomes in an electron microscopic study of a skin punch biopsy (fig 1C).
Results of cytogenetic and DNA investigations

High resolution chromosomal analysis showed the karyotype 46,Y,del(X)(p22.3) in the patient (700 bands, 30 metaphases) and the karyotype 46,X,del(X)(p22.3) in his mother (700 bands, 50 metaphases) (fig 3).

Genomic DNA of patients BK and W1 was characterised by Southern and PFGE analysis. Twelve DNA probes were used from Xp22.3 ranging from ANT3 in the pseudoautosomal region to p71-7A (DXS69), which maps more than 10 Mb away from the telomere (M Wappenaar, personal communication). In an attempt to identify junction fragments at the proximal breakpoints, PFGE experiments were performed using the following probes: P45 (DXS410), 782 (DXS85), and p71-7A (DXS69). The results of these hybridisation experiments are summarised in the table.

The large deletion in patient BK corresponded to absent hybridisation signals with all probes used except 782 (DXS85) and p71-7A (DXS69). In contrast to patient W1, the probe P45 (DXS410) was deleted in BK (fig 4). Furthermore, altered MluI and NotI fragments were detected with probe 782 (fig 5), which were not found in 10 control females and in patient W1. Unaltered fragments were identified in both patients with probe p71-7A, indicating that it maps proximal to probe 782. These data suggest that the critical interval for OA1 is located between DXS143 and DXS85, segregating with the locus DXS410.

Discussion

This patient, with a contiguous gene syndrome, showed a deletion of more than 10 Mb. Cytogenetically, the deletion comprises most of band Xp22.3. On DNA analysis, the most distal probe tested and deleted is ANT3, which maps 1-3 Mb away from the telomere in the pseudoautosomal region. Nine other Xp22.3 probes analysed are deleted including P45 (DXS410) which maps more than 10 Mb away from the telomere (M Wappenaar, personal communication). Ten genes are assigned to the deleted region. They include four genes which so far have not been associated with human disease: a gene for adenine nucleotide translocase (ANT3), a gene for a colony stimulating factor receptor (CSF), a gene for the leucocyte antigen MIC2, and the gene GS1. Six other genes include those with mutations leading to short stature, chondrodysplasia punctata, mental retardation, ichthyosis, Kallmann syndrome, and ocular albinism. Patient BK shows symptoms of all these disorders. Additional phenotypes could be obscured by this complex clinical picture. The so far unexplained hypertriglyceridaemia,
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should result from unbalanced gene expression within the deleted region where several genes have been shown to escape X inactivation.1

The typical features of chondrodysplasia punctata are present in patient BK. They include alterations of epiphyseal bones, brachytelephalangy, and the peculiar dysmorphic features, a manifestation mimicked by warfarin embryopathy.30,37 No signs of chondrodysplasia punctata were described in patient W1.16 This is in concordance with positive hybridisation results for MIC2 (table) and for a probe which maps distal to MIC2.

Of all the symptoms in this case, mental retardation is the most general. The number of missing transcripts in the deletion described could contribute to reduced cerebral function, a hypothesis which gains support from the observation that the extent of mental retardation seems to correspond to the length of the terminal deletions described. Patient BK is severely mentally retarded but a milder expression has been described in patient W1 (table) who has a much smaller deletion.16

The diagnosis of ichthyosis and Kallmann syndrome is consistent with the absent hybridisation signals for both the STS and the KAL-X cDNA. Although the endocrinological status has not been assessed, the mirror movements recorded are pathognomonic of this condition (our observation and B Heye, personal communication).

Two linkage analyses, published recently, map the OA gene proximal to KAL-X.13,38 Single recombinants analysed in a large Newfoundland kindred define a critical interval for the OA1 gene between the loci DXS143 and DXS85.15 For deletion mapping of the OA gene, special emphasis has been placed on the ocular condition in patient BK and his family. Ocular symptoms observed in the patient, his mother, and his sister are consistent with the diagnosis of X linked ocular albinism. Nettleship-Falls type of ocular albinism is characterised by poor visual acuity, nystagmus, macular hypoplasia, and hypopigmentation of the retina as well as the iris.39,40 Carrier females show a 'mud splattered' appearance of the fundus with hyperpigmented streaks and most of them have iris translucency and macromelanosomes.41 Both the aberrant fundus appearance and macromelanosomes were found in the patient's mother who carried the Xp22.3 deletion on one X chromosome.

In contrast, no symptoms of ocular albinism were observed in patient W1. This patient presents mental retardation, ichthyosis, and Kallmann syndrome because of a deletion involving the loci distal as well as proximal to the STS and KAL-X genes (table). Both patients are deleted for DXS143 which, according to the physical map of this region, maps proximal to the KAL-X gene. The locus DXS410 is deleted in BK, but not in W1. The junction fragments seen with probe 782 (DXS85) in BK indicate a breakpoint close to this probe. Probes more proximal give rise to unaltered fragments in both patients. The two proximal breakpoints of patients BK and W1 therefore define a deletion interval which

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**Figure 5** Junction fragments in patient BK detected by the probe 784 (DXS85). High molecular DNA from patients BK and W1 was digested with the rare cutters NotI (N), MluI (M), and BstHI (B), respectively, and blotted after PFGE onto a nylon membrane. The filter was hybridised with probe 784 and washed under stringent conditions (60°C, 0.1 × SSC). Exposure time was four days. Fragment lengths (kb) are indicated.

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for example, could suggest a gene affecting lipid metabolism.

The presence of a growth gene(s) has been postulated after analyses of karyotypes from X chromosome rearrangements at the tip of Xp.19-36 The critical region for the putative growth gene(s) comprises the pseudoautosomal region between the telomere and locus DXYS17 which maps about 2 Mb proximal to the telomere.11 The deletion described includes ANT3, which maps to the critical region. The height of patient BK remained far below the 3rd centile (mean −5 SD), much shorter than in patients with chondrodysplasia punctata.36 Moderate short stature of female carriers with Xp22.3 deletions has been reported by Curry et al.36 and Ballabio et al.2 Both carrier women investigated had short stature, as well as several symptoms typically seen in Turner's syndrome. These include a low posterior hair line and pigmented naevi in the mother, a short neck and hypoplasia of the distal ulna in the sister, as well as abnormal palmar dermatoglyphs in both and Madelung deformity in the mother. All these symptoms
should contain the OA1 gene (fig 4). The minimum distance of this interval measures 400 kb because of the length of the MluI fragment detected in W1 with probe P45 (DXS410). The maximum interval is determined by the distance between the loci DXS143 and DXS85. This distance is unlikely to be larger than 1 to 2 Mb: DXS143 has been mapped 9-5 Mb from the centromere.4 This distance is already close to the cytogenetically estimated length of about 10 to 12 Mb (6 to 7%) for the whole of Xp22.3 (164 Mb = 100% for the whole X).44 DXS69, which maps proximal to DXS85, is also located within band Xp22.3.15

Deletion mapping is undertaken under the assumption that no chromosomal aberrations other than plain deletions have taken place. Most of these are characterised in the Xp22.3 region by DNA markers today agree with this assumption.1 There are no indications of a complex rearrangement in patient BK from cytogenetic and DNA analysis. One explanation for the discrepant results on the location of the OA1 gene is a complex chromosomal rearrangement in one of the patients published recently.1 Analysis of terminal Xp deletions in this study places the OA1 gene proximal to DXS143 and distal to DXS85, thus providing starting points for cloning of the OA1 gene.

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