A microdeletion in 19q13.2 associated with mental retardation, skeletal malformations, and Diamond-Blackfan anaemia suggests a novel contiguous gene syndrome

Dmitry Tentler, Peter Gustavsson, Göran Elinder, Ole Eklöf, Laurie Gordon, Ariane Mandel, Niklas Dahl

Abstract
Diamond-Blackfan anaemia (DBA) is a constitutional red blood cell hypoplasia which may be associated with a variety of developmental abnormalities. A gene for DBA was recently mapped to chromosome 19q13.2 and subsequently cloned. Analysis of 19q marker alleles in DNA of sporadic DBA cases showed de novo microdeletions in three patients also presenting with mental retardation. We have studied one of these patients and characterised the deletion by fluorescence in situ hybridisation (FISH) to extended DNA fibres. The deletion was shown to be continuous over a 3.2 Mb region and the fibre-FISH analysis showed both chromosomal breakpoints. In combination, the clinical and molecular findings suggest a contiguous gene syndrome with a gene locus for mental retardation and, probably, skeletal malformations included in the deletion.

Keywords: chromosome 19q13; microdeletion syndrome; fibre-FISH

Diamond-Blackfan anaemia (DBA, McKusick 205900) is a red cell hypoplasia characterised by subnormal erythropoiesis with absent or decreased red cell precursors in the bone marrow. A subset of patients has associated phenotypic abnormalities including thumb malformations, urogenital abnormalities, atrial or ventricular septal defects, prenatal or postnatal growth retardation, learning difficulties, strabismus, and cataract. At least 10% of patients with DBA have a positive family history for the disorder following either autosomal dominant or autosomal recessive inheritance. One locus for DBA has been mapped to chromosome 19q13.2 in multiplex families and we recently cloned the gene which was shown to encode the ribosomal protein (RP) S19. We have analysed 19q13 marker alleles which flank the RP S19 gene in 50 sporadic cases with DBA. Three subjects also with mental retardation (MR) were found to be hemizygous for several marker loci. One of these patients with microdeletions was recently reported and we present here the second of the three patients. The clinical features of this patient are described as well as the precise mapping of the deletion breakpoints using FISH to extended DNA fibres. The complex phenotype including mental retardation and generalised skeletal malformations suggests a contiguous gene syndrome.

Case report
The patient is a 12 year old male, born at term after an uncomplicated pregnancy to healthy, unrelated parents. There is no family history of skeletal, mental, or haematological disorders. At the age of 2 months he was referred to a paediatric clinic because of pallor. Haemoglobin concentration was 4.8 g/dl and no reticulocytes were detected in peripheral blood. Bone marrow aspirate showed a selective decrease in erythroid precursors but otherwise normal cellularity. Chromosome analysis (G banding) showed a normal 46,XY karyotype and chromosome fragility test with mitomycin C was normal. At 22 months of age the boy had relatively short extremities, slight macrocephaly (+2 SD), and short stature (−3 SD). Major radiological traits recorded at 1 year 10 months of age included 13 pairs of horizontally oriented ribs, mild thoracic platyspondyly, vertebral bone in bone formation, and lack of interpedicular widening with ventrally concave posterior aspects of the lumbar vertebrae (fig 1A). The iliac bodies were narrow, the proximal femoral epiphyses asymmetrical, and the smaller right one in addition slightly irregular (fig 1B). Coxa valga with short and broad femoral necks as well as bow legs with epiphysial flare were also noted (fig 1C). Radiological investigations of the hands were normal.

Moderate psychomotor retardation was diagnosed at the age of 4 years. A cranial CT scan at this age was normal, as were ultrasound and x ray investigations of the heart and lungs. MRI of the liver and spleen at 7 years of age showed iron accumulation.

At 12 years of age he requires transfusions every three weeks. His height is 140 cm (−1.5 SD), weight is 36 kg (−0.5 SD), and head circumference is 58.2 cm (+2 SD). The span of the upper extremities is 131 cm and his sitting height is 73 cm. No follow up x ray has been performed.

Materials and methods
FIBRE-FISH ANALYSIS
In order to determine the deletion breakpoints and for a precise size estimation of the deletion
we used fibre-FISH analysis. Fluorescence in situ hybridisation to extended DNA fibres was performed essentially as described elsewhere. Unfixed linearised DNA fibres were prepared on microscope slides. Ficoll separated lymphoblastoid cells derived from the patient and lymphoblastoid cells from a control were embedded in 1% low melting point agarose at a concentration of 10⁷ cells/ml.

Agarose blocks were incubated at 50°C in 2 mg/ml proteinase K and 1% N-lauroylsarcosine in 50 mmol/l EDTA for 18 hours. A small piece of agarose embedded DNA was placed on a microscope slide and 15 µl of water was added to the agarose block. The slide was heated on a heat block (50 seconds, 100°C) and DNA was extended manually on a slide using the edge of another microscope slide. Extended DNA fibres were denatured using 70% formamide/2 × SSC at 74°C for four minutes followed by an ice cold ethanol series. Cosmid DNA was labelled either with biotin or digoxigenin using nick translation. Double hybridisation was detected using a single layer of FITC-avidin (Vector Laboratories) and rhodamine labelled anti-digoxigenin antibodies (Boehringer Mannheim). Visualisation of probes was performed as described previously.

COSMID CLONES
The probe and interprobe distances in kilobases were based on a detailed physical map of the region. The chromosome 19 cosmids were obtained from the Lawrence Livermore National Laboratory (CA) with their corresponding numbers. In total, nine cosmids were used for fibre-FISH analysis and their relative order from the centromere to the telomere is 15848, 16767, 22692, 16923, 8918, 15894, 26656, 28280, and 29827.

Results
Cosmid clones assigned to the chromosome 19q13 map identified the deletion by FISH to metaphase chromosomes from patient RG. The extremities of the deletion were approximately mapped between, and close to, cosmids 15848 (CYP2B6) on the centromeric side and 13519 (D19S19) on the telomeric side. Additional cosmids in the vicinity of loci CYP2B6 and D19S19 were subsequently hybridised to DNA fibres derived from patient RG and from a normal control, respectively. Two to three cosmids of known relative position and size (38-45 kb) were cohybridised and the clones were distinguished by two colour detection. Hybridisation with cosmid 22692 to DNA fibres derived from the patient showed two different signal patterns corresponding to a length of approximately 40 kb and 15 kb, respectively. Similarly, hybridisation with cosmid 28280 showed two types of signals corresponding to either 40 kb or 20 kb. We then cohybridised the two rearranged cosmids 22692 and 28280 together with the control cosmid 29827 to extended DNA fibres from the patient. The results showed that signals of cosmids 22692 and 28280, normally separated by 3.2 Mb, were located head to tail on approximately half the numbers of fibres (fig 2B). In the same experiment, separate and normal signals from cosmids 28280 and 22692 were also detected (fig 2A). These signals are expected from the non-rearranged fibres. The combined results indicate that the deletion is continuous and that the breakpoints are located within cosmid 22692 (centromeric) and within cosmid 28280 (telomeric).
Discussion
We have characterised a 3.2 Mb microdeletion in chromosome 19q13.2 associated with DBA, mental retardation, and skeletal abnormalities. Fibre-FISH analysis enabled the deletion breakpoints in patient RG to be visualised at a high resolution and the deletion was found to be continuous. The deleted region spans more than 20 known genes\(^{13,14}\) and there are additional expressed sequence tags (ESTs) mapped to 19q13.2. Several gene families are clustered in a region corresponding to the deletion including the CYP2, the CGM, and the PSG gene families. It is known that tandemly repeated and homologous sequences may predispose to recombination events.\(^{15,16}\) However, such a mutation mechanism needs to be verified by DNA sequencing of the deletion breakpoints in our patient.

The combined clinical and molecular findings suggest that the patient has a contiguous gene syndrome. The gene encoding RPS19, previously shown to be mutated in a subset of DBA patients, is included in the deletion. The involvement of other genes in the patient's phenotype is substantiated by the recent clinical presentation of a boy with DBA, psychomotor retardation, and dysplastic changes of bone metaphyses and the vertebral column.\(^7\) This patient carries a deletion of approximately 3 Mb which, although not precisely characterised, overlaps with the one found in our patient. Additional support for a contiguous gene syndrome comes from the investigation of a female with DBA and an associated balanced X:19 translocation which interrupts the RPS19 gene.\(^7\) This female is haploinsufficient for the DBA gene on 19q13.2 and she has normal mental development with no skeletal malformations. In the subset of DBA patients screened for mutations in the RPS19 gene,\(^7\) none of the cases caused by point mutations in RPS19 had psychomotor retardation, while patients with abnormal mental development were found to carry deletions in the 19q13.2 region.\(^8\) The data point to the presence of a gene responsible for mental retardation linked to the RPS19 gene.
Our patient presents with some physical abnormalities, for example, short stature, that have been described previously as frequently associated with DBA. However, there are very few published cases of DBA patients with malformations of the lower limbs, pelvis, and vertebrae. The similar phenotype in our patient may indicate a gene for the skeletal malformations involved in the deletion.

In conclusion, the presence of complex clinical features in our patient suggests the involvement of a gene for mental retardation and, probably, skeletal abnormalities linked to the RPS19 gene. The finding of such uncommon features in a small number of DBA patients may be explained by deletions affecting contiguous genes. The comparison of characterised deletions associated with DBA may further delineate the genomic regions responsible for the complex phenotype of this patient.

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