Juvenile rheumatoid arthritis and del(22q11) syndrome: a non-random association

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Abstract
Del(22q11) is a common microdeletion syndrome with an extremely variable phenotype. Besides classical manifestations, such as velocardiofacial (Shprintzen) or DiGeorge syndromes, del(22q11) syndrome may be associated with unusual but probably causally related anomalies that expand its phenotype and complicate its recognition. We report here three children with the deletion and a chronic, erosive polyarthritis resembling idiopathic cases of juvenile rheumatoid arthritis (JRA).

Patient 1, born in 1983, initially presented with developmental delay, facial dysmorphism, velopharyngeal insufficiency, and severe gastro-oesophageal reflux requiring G tube feeding. From the age of 3 years, he developed JRA, which resulted in severe restrictive joint disease, osteopenia, and platyspondyly. Patient 2, born in 1976, had tetralogy of Fallot and peripheral pulmonary artery stenosis. She developed slowly, had mild dysmorphic facial features, an abnormal voice, and borderline intelligence. JRA was diagnosed at the age of 5 years. The disorder followed a subacute course, with relatively mild inflammatory phenomena, but an extremely severe skeletal involvement with major osteopenia, restrictive joint disease (bilateral hip replacement), and almost complete osteolysis of the carpal and tarsal bones with phalangeal synostoses, leading to major motor impairment and confinement to a wheelchair. Patient 3, born in 1990, has VSD, right embryotoxon, bifid uvula, and facial dysmorphism. She developed JRA at the age of 1 year. She is not mentally retarded but has major speech delay secondary to congenital deafness inherited from her mother.

In the three patients, a del(22q11) was shown by FISH analysis. These observations, and five other recently published cases, indicate that a JRA-like syndrome is a component of the del(22q11) spectrum. The deletion may be overlooked in those children with severe, chronic inflammatory disorder.

Keywords: juvenile rheumatoid arthritis; del(22q11) syndrome

Del(22q11) syndrome is a newly recognised contiguous gene syndrome resulting from a (usually) submicroscopic deletion in the proximal part of the long arm of chromosome 22, encompassing in its phenotype Shprintzen velocardiofacial syndrome, Takao conotruncal anomalies face syndrome, DiGeorge anomaly, some instances of Opitz GBBB syndrome type 2, and isolated outflow tract defects (such as truncus arteriosus, tetralogy of Fallot, or interrupted aortic arch). The incidence could be as high as 1/3500 births.

The spectrum of clinical anomalies associated with 22q11 monosomy is remarkably variable. Recently, juvenile rheumatoid arthritis (JRA) has been reported in association with del(22q11) in five children. We report here three further cases of this association, indicating that the presence of a JRA-like disorder may not be fortuitous, but rather causally related to the presence of the deletion.

Case reports
PATIENT 1
This girl is the second child of unrelated parents, born after an uneventful pregnancy. Clinical investigations at birth showed a tetralogy of Fallot, which was partially surgically corrected at the age of 10 months, pulmonary valvular insufficiency, and peripheral pulmonary stenosis. No other malformations were present. Assessment of calcium metabolism and immunodeficiency are not available from that time, but suspicion of DiGeorge syndrome was never raised in her medical records.

At the age of 4 years (fig 1A), she was admitted for painful inflammatory swelling of the knees, for which she was treated with indomethacin and salicylates. Sedimentation ratio was repeatedly high. There was no hepatosplenomegaly. Calcium homeostasis, serum parathormone and thyrocalcitonin, and lumphocyte counts were normal. The clinical course was notable for multiple relapses of the arthritis during the next 15 years, affecting all large and small joints, usually with an important inflammatory component but no fever (“subacute” JRA). Therapy was initially limited to anti-inflammatory drugs. Corticosteroids were used from 9 to 18 years and d-penicillamine from 12 to 14 years. There were no ocular manifestations of iridocyclitis or uveitis. At the age of 12, transient non-viral hepatitis was noted. Progressive right ventricular insufficiency was treated with digitalis. Despite corticoids, progressive destruction of the hips and knees led her to become wheelchair bound. Because of progressive flexion deformity, she required tenotomies around the hips and knees at the age of 15. At the age of 20, hip replacement was performed. There was major involvement of the hands and
wrists. Immune investigations repeatedly showed hypergammaglobulinemia, circulating IgA immune complexes, and absence of rheumatoid factor, anti-DNA antibodies, and antinuclear antibodies. HLA typing was A3A9-B35-DR1DR5.

When we saw her at the age of 22, she was non-ambulatory. Ankylosis of the elbows and finger retraction severely restricted her ability in fine motor skills. She was of apparently low normal intelligence. She had a large nose, narrow palpebral fissures, narrow mouth (fig 1B, C), and high palate. She had a high pitched, nasal voice. IQ testing was not performed. A skeletal survey showed kyphoscoliosis, generalised arthropathy (including the sacroiliac joints), severe osteopenia, carpal osteolysis, and secondary synostosis of the metacarpals and phalanges (fig 2). She had attended normal school until the age of 10 and was switched thereafter to special schooling. Poor academic achievement was attributed to her physical problems.

Del(22q11) was shown on spread chromosomes hybridised with the specific DNA probe D22S75 and the control probe D22S39 (Oncor), using standard FISH techniques, as recommended by the manufacturer.

The mother suffered recurrent psychiatric problems and had abandoned her daughter during infancy. Features noted in family photographs suggest that she probably also had the deletion. None of the relatives had ankylosing spondylitis, inflammatory bowel disease, uveitis, psoriasis, or psoriatic arthritis (and the same is true for the other two cases).

PATIENT 2

This boy is the second child of healthy, unrelated parents. His neonatal course was characterised by low birth weight, hypotonia, and poor feeding. Language development was delayed. Clinical examination at the age of 2 years showed a coarse, expressionless (“myopathic”) face (fig 3A), brachycephaly, with normal OFC, flat nasal bridge,

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Figure 1  Patient 1. (A) Front view in infancy. (B, C) Appearance at the age of 22: note prominent nose with large tip and narrow palpebral fissures.

Figure 2  Patient 1. (A) Severe hand deformities. (B) X ray of the hand at the age of 22 showing generalised osteolysis, osteoporosis, and secondary bone fusion.
Figure 3 Patient 2. (A) Front view in infancy. (B, C) Appearance at the age of 12: note prominent nose with large tip and narrow palpebral fissures.

PATIENT 3

This child, born in 1990, was the first girl of unrelated French parents. The mother suffered congenital unilateral deafness. Birth weight was 2670 g, length 46 cm, and OFC 34 cm. Clinical investigations at birth showed VSD and camptodactyly of the toes. There were no feeding problems during the first year of life and psychomotor development was not delayed. Severe bilateral deafness was diagnosed at 9 months. At the age of 1 year, she presented with acute polyarthritis affecting large and small joints with mildly increased CRP. JRA was diagnosed and required sylacitate and later corticosteroid treatment. She later attended a school for deaf children. She was considered to be of normal intelligence.

When first evaluated aged 7, several dysmorphic features were noted: ridged metopic suture, narrow, upward slanting palpebral fissures, large nose with bulbous tip, small mouth, high vaulted palate with bifid uvula, crowded teeth, small ears, and right posterior embryotoxon. She showed painful ankylosis of the wrists, elbows, ankles, knees, fingers, and toes, with secondary retractions of the small joints. Calcium homeostasis, circulating vitamin A, and PTH were normal. X-rays showed osteopenia (confirmed by osteodensitometry) and generalised JRA arthropathy. Immunological investigations showed normal lymphocyte count and the absence of rheumatoid factor and anti-DNA and antinuclear antibodies. CD3, CD4, CD8, and helper/suppressor ratio were in the normal range. HLA genotype was A2A3-B13B62-CW6-DR13-DQ6.

Karyotyping with probe D22S75 (Oncor) showed a 22q11 deletion. The mother was not deleted; the father was not tested.

Discussion

Del(22q11) is a multiple congenital anomaly (MCA) syndrome, with a remarkably variable and continually expanding phenotype that merges Shprintzen velocardiofacial syndrome, most cases of DiGeorge sequence, some cases of Robin sequence, and rare, often atypical instances of the CHARGE association. Although phenotypic expression of the disorder has become increasingly better recognised, relatively few data have been recorded on immune dysfunction beyond the neonatal period, during which manifestations of DiGeorge syndrome (with thymic hypoplasia and immunodeficiency) may be observed in roughly 10% of cases. T cell dysfunction has been observed in the milder cases of DiGeorge syndrome. Some case reports mention autoimmune disorders in the syndromes known to belong to the del(22q11) spectrum. A single older patient with DiGeorge syndrome has been reported with autoimmune Graves disease. Two adults with del(22q11) were recently reported with adult onset idiopathic thrombocytopenia.
associated in one case with chronic leucopenia. A similar pattern of infantile thrombocytopenia followed by haemolytic anaemia at the age of 8 was mentioned in an abstract. In the report of Pinchas-Hamiel et al., a child with DiGeorge syndrome suffered autoimmune haemolytic anaemia (responding to corticosteroids), thrombocytopenia, and chronic, granulomatous, probably autoimmune hepatitis.

The exact classification of the rheumatoid disease in our cases was the subject of much debate during the preparation of this manuscript. American authors (who prefer the term JRA) and European authors (who often use the term "juvenile chronic arthritis (JCA)) appear to have slightly different definitions of these disorders. The most recently adopted international nomenclature uses the term juvenile idiopathic arthritis (JIA). Whatever name is chosen (JRA, JCA, or JIA), it requires that the arthritis is idiopathic, that is, not associated with any other recognisable condition. As, before the discovery of the microdeletion, the three patients were considered to have bona fide JRA by several rheumatologists in the three university hospitals where they were followed, and as a causal relationship with del(22q11) has still to be proven, we suggest that the erosive polyarthritis observed in association with del(22q11) should be described as a JRA-like disorder. Synovial puncture was not performed, so we have no idea of the cytological appearance of the articular fluid.

A JRA-like disorder has been reported in five children with del(22q11). In the report of Rasmussen et al., onset was at 7 and 6 years, respectively. Both had positive antinuclear antibodies and one of them was positive for rheumatoid factor. HLA typing was not performed (S Rasmussen, personal communication). Interestingly, case 2 of Rasmussen et al. inherited the deletion from her father and had two deleted sibs: neither of them developed JRA. In the report of Sullivan et al., two boys and one girl were described with JRA and del(22q11). Onset was at 17 months, 19 months, and 5 years, respectively. One had positive antinuclear antibodies and all were negative for rheumatoid factor. They each had two HLA alleles associated with an increased risk of polyarticular JRA. The three children showed immunological anomalies: all three had impaired response to mitogens, two had IgA deficiency, the two youngest patients had abnormal CD4:CD8 T cell ratio, and the third one an increased ratio. In each case, as in our patients, the course was "subacute", polyarticular and there were no ocular complications nor systemic features such as rash, organomegaly, or lymphadenopathy and, apparently, only mild increase of inflammatory markers.

JRA is a common autoimmune disease of childhood. Its incidence is about 1/10 000 and its prevalence 1/1000. Although most cases are sporadic, many multiplex pedigrees have been observed in which JRA shows familial aggregation compatible with partially penetrant autosomal dominant inheritance. One third to one half of the familial transmission is accounted for by linkage to the HLA loci with predisposing alleles such as DR5, DR8, DQ2,1, DRB1 (particularly DRB1*0401), and DQB1, whereas linkage to DR1 or DR4, often observed in adult onset rheumatoid arthritis, is rarely noted in the paediatric population with positive rheumatoid factor. No linkage has been shown with chromosome 22.

The pathogenesis of JRA is complex, but central role is attributed to T cells, as the only known function of DR is to present peptides of the CD4 T lymphocytes. Abnormal suppressor activity and abnormal immunoglobulin production are part of JRA and point to an abnormal T cell function. As T cell dysfunction is a major component of DiGeorge syndrome, the association of autoimmune phenomenon with del(22q11) may not be coincidental, although the exact pathogenic mechanism remains to be solved. Rasmussen et al. noted that in their two families other members had joint problems. They speculated that JRA could occur for genetic reasons unrelated to the del(22q11).

We would rather suggest that JRA is pathogenically related to del(22q11), the T cell dysfunction component of DiGeorge sequence being a possible adjuvant for developing JRA in those who have other predisposing factors (such as specific HLA subtypes). But the HLA typing of JRA positive del(22q11) cases is not available in all cases, but in our case 1, as in that of Sullivan et al., the expected DR1 group was present.

Del(22q11) syndrome is currently considered to be a contiguous gene syndrome. The deletion usually spans 3 Mb, but some patients have much smaller, nested deletions. The critical region is about 480 kb in size. The difference in deletion size or modifying genetic loci or both may account for a part of the diversity of expression of the disorder, but the genomic basis of del(22q11) syndrome is still unknown. Six genes and several unidentified ESTs have been mapped in the critical region. These include the clathrin heavy chain gene CLTD, the G SCL, a goosceoid-like homeobox gene expressed in early human development, the mitochondrial citrate transport protein CTP, the HIRA, a protein related to yeast repressors of histone transcription genes, and TMVCF, a transmembrane protein of unknown function. Other genes have been mapped within the larger deletion, but outside the critical region: TUPLE1, a putative transcription factor; ZNF74, a zinc finger gene, a homologue of Drosophila dishevelled gene, DGC6, a gene with homologies to Drosophila gonadal protein gdl, the γ1 chain of human laminin, T10, a sequence expressed during embryonic development, and several other less well defined genes (DGC5, rpL28, ARV, and V1-41). None of these genes appears to play an important role in T cell immunity.

The incidence of JRA in del(22q11) is unknown. Sullivan et al. gave an estimate of 4/1000 for their local series of cases, but JRA has never been mentioned in older reviews on DiGeorge syndrome. This is not mentioned in the recent collaborative European study. This could be an indication of the rarity of this complication, but we feel it is more likely that the
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several complications of JRA may obscure the classical phenotype of del(22q11) and prevent its recognition. The incidence of del(22q11) among JRA patients is currently unknown. Short stature and failure to thrive, which are associated with del(22q11) syndrome, can be attributed to JRA. Facial anomalies in del(22q11) are quite subtle and can be blurred by the effects of corticotherapy. It is likely that the long periods of illness and their interference with normal schooling may be considered responsible for school difficulties, behavioural problems, and mild intellectual impairment. For these reasons, referral for clinical genetic evaluation is probably infrequent so that del(22q11) is more likely to be missed in this population of children.

We would strongly recommend careful re-evaluation of all children with JRA who show velocarpyangeal insufficiency, learning disabilities, or heart anomalies for possible 22q11 deletion. In reviewing papers reporting chronic arthritis with multisystemic involvement, we wonder whether some cases may have del(22q11). This could certainly be the case for patients 2 and 3 in a “new” syndrome reported by Coffin, consisting of mental retardation, arthritis, deafness, and facial dysmorphism. Chronic, infantile, neurological, cutaneous, and articular (CINCA) syndrome (Lorber syndrome) designates series of isolated patients reported with polyarthritis, developmental problems, failure to thrive, and hearing loss. Some of them could have del(22q11). In an ill-defined syndrome labelled camptodactyly-arthritis, some patients may also have the deletion.