Discussion

The association of a constitutional deletion of the terminal end of q22 (band q13.3–qter) with multiple meningiomas is of great interest. We suspected the presence of intracranial cysts in the patient because of his symptoms, the results of CT scan, and EEG examination. However, meningeal hypoplasia and dysplasia, including meningiomatosis, and an anomalous cyst composed of meningocytic elements were found at necropsy. As is well known, 22q– or monosomy 22 can be commonly observed in cultured meningioma cells. Recently, the relationship between constitutional and somatic chromosomal changes in patients with retinoblastoma and Wilms’s tumour has been shown. These findings tend to support a close relationship between a partial deletion of chromosome 22 and meningeal malformations including tumours. The karyotype of the tumour cells was not investigated in our patient. However, the presence of a constitutional ring chromosome 22, which may lack the distal end of the long arm, supports the suggestion of Zankl and Zang that genetic information involved in the control of cell proliferation is located in this segment. Recently, the human homologue of the simian sarcoma virus oncogene, c-sis, has been assigned to 22q. The proto-oncogene c-sis might be related to the genesis of meningiomas. Furthermore, these chromosome changes may be tissue specific, in the sense that they may predispose specifically to meningiomas, because necropsy revealed no neoplasia in other organs of our patient.

In conclusion, the findings described in this paper suggest that constitutional deletions of chromosome 22 may predispose to tumourigenesis of meningiomas. To confirm this, further neuropathological studies of patients with 22q– or monosomy 22 are required. Clinical examination for congenital anomalies and chromosome analysis should be carried out in all patients with meningiomas.

References


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Case reports

Prenatal diagnosis and follow up of a child with a complex chromosome rearrangement

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SUMMARY A case of de novo, apparently balanced, three way exchange by translocation plus a pericentric inversion is described. The karyotype is 46,XX,t(6;11)(p21;q21),t(11;21)(q21;p13),inv(6)(p21q11) and was ascertained through second trimester amniocentesis. The structural rearrangements appear balanced.

The child was phenotypically normal at birth. Growth and motor development were normal until 30 months, at which time linear growth dropped below the 5th centile. In addition, there was delayed speech development at 2 years of age.

As far as we can determine, this is the first report of a three chromosome exchange including a pericentric inversion ascertained through genetic amniocentesis.

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Case report

The mother was 30 years of age and the father 31 at the time of her pregnancy. There had been two previous normal term pregnancies followed by a pregnancy which produced a term stillborn anencephalic infant. Genetic amniocentesis was performed because of the pregnancy with anencephaly and parental concern. Ultrasound before amniocentesis at 16 weeks' gestation was unremarkable. Alphafoetoprotein analysis was within normal limits for 16 weeks' gestation. Karyotype analysis of cultured amniotic fluid cells revealed the karyotype 46.XX.t(6;11) (p21;q21),t(11;21) (q21;p13),inv(6) (p21q11) (fig 1). Subsequent chromosome studies on peripheral blood samples from each parent showed normal karyotypes in both. Repeat amniocentesis confirmed the original karyotype. After counselling and review of all available data, the parents elected to continue the pregnancy. The infant was born at term by vaginal delivery. Labour and delivery were uncomplicated. Apgar scores were 9 at one and five minutes. The head circumference was on the 50th centile, the length on the 50th centile, and the weight on the 25th centile. No dysmorphic features were noted. Peripheral blood chromosome analysis confirmed the karyotype. During the first year of life, transient episodes of sucrose intolerance were documented. During the first 30 months, linear growth remained between the 5th and 10th centiles. Afterwards linear growth dropped to less than the 5th centile with weight remaining at the 35th centile and head circumference at the 50th centile. Recently, linear growth rate has returned to the 5th centile. No symptoms have been associated with the transient decline in linear growth. CT scan was normal and no deficiency in growth hormone was found. Developmental milestones have been normal: however, her first words were spoken at 2½ years and at 2¾ years there are no complete sentences.

Discussion

It has been possible to identify 45 pregnancies from previous publications concerning kindreds with complex chromosome structural rearrangements occurring either as a new mutation or as a familial multichromosome structural rearrangement.1-17 Both de novo and familial structural rearrangements produced abnormal pregnancies. Fourteen of the 45 pregnancies (approximately 31%) produced liveborn infants who were phenotypically normal, either with a normal karyotype or with a balanced karyotype identical to one parent. Fifteen liveborns (approximately 33%) with multiple anomalies also had unbalanced complex translocations. Fifteen pregnancies (approximately 33%) ended as an

FIG 1  Karyotype showing complex chromosome rearrangement involving chromosomes 6, 11, and 21.
The patient described by Watt and Couzin in 2018 also had speech delay and in addition had short stature. Another patient with complex translocations involving chromosomes 11 and 21 (p11) is translocated to chromosome 21 (p13) (fig. 2). Additionally, a pericentric inversion of chromosome 6 occurred between p21 and q11.

Our patient had a normal phenotype at birth and normal developmental milestones except for speech delay noted at 2½ years. The patient described by Watt and Couzin also had speech delay and in addition had short stature. Another patient with a complex multichromosome translocation had speech delay at the age of 6, an IQ of 82 on the Stanford-Binet scale, and mild dysmorphic features. The three case reports of complex translocations all appear balanced, involve different chromosome pairs, and are associated with speech delay plus variable phenotype abnormalities. We are unable to identify any report of completely normal subjects with de novo translocations involving three or more chromosomes. This may be an important consideration when the abnormal karyotype is ascertained by genetic amniocentesis.

Watt and Couzin and Couzin et al. have suggested that meiosis is likely to be a complicated event in multichromosome translocations. There are at least five breakpoints in the three chromosome exchange described here, including the pericentric inversion. Meiotic configurations will involve chromosome 11 with chromosome 6 or 21. The addition of the pericentric inversion on chromosome 6 further raises the risk for duplication and deficiencies in gametes.

The risk for abnormal reproductive outcome in carriers of complex rearrangements appears to be increased but not absolute. It might be expected that a complex translocation which includes a pericentric inversion carries an even greater risk for abnormal segregates at meiosis. Prenatal fetal assessment will be recommended for any successful pregnancy involving the proband in this report.

References
Case reports


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Primary myelodysplastic syndrome with complex chromosomal rearrangements in a patient with Klinefelter’s syndrome

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SUMMARY A patient with Klinefelter’s syndrome and diabetes mellitus was diagnosed as having myelodysplasia. Cytogenetic analysis of the peripheral blood and the bone marrow cells confirmed the presence of a constitutional 47,XXY chromosome complement. In addition, complex karyotypic abnormalities were present.

Several studies have suggested an increased incidence of extraglandal germ cell tumours,1 carcinoma of the breast,2 and acute myeloid leukaemia3-5 in patients with Klinefelter’s syndrome. In this report we describe a patient with Klinefelter’s syndrome who developed a preleukaemic state (myelodysplastic syndrome). Cytogenetic analysis of the bone marrow and peripheral blood of this patient showed a constitutional 47,XXY chromosome complement. In addition, complex karyotypic abnormalities were present. The significance of these findings is discussed.

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Case report

A male Caucasian, aged 68, presented with a four month history of increasing breathlessness. Eight years previously he had been admitted to another hospital in a hyperosmolar, non-ketotic diabetic coma and at that time he was noted to be tall (185 cm) and to have hypogonadism and prognathism. He had been investigated for acromegaly and was found to have raised levels of FSH and LH and normal levels of TSH, free T4, growth hormone, and prolactin. Skull x-ray showed a normal pituitary fossa. His diabetes had been controlled with oral hypoglycaemic agents and the only other drug therapy he had received was testosterone at six monthly intervals. He was unmarried and had no children. He had no history of exposure to chemicals.

On presentation at this hospital he was noted to be pale and lacking facial and body hair. He had gynaeacomastia and small testes. Full blood count showed Hb 7·5 g/dl, WBC 3·5 \times 10^9/l (neutrophils 46%, lymphocytes 45%, monocytes 3%, eosinophils 6%), and platelets 309 \times 10^9/l. Blood film showed dimorphic red cells, hypogranular neutrophils, and platelet anisocytosis. Bone marrow aspirate showed
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