Dicentric chromosome in the bone marrow of a child with megakaryoblastic leukaemia and Down’s syndrome

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SUMMARY A two year old girl with Down’s syndrome (constitutional karyotype: 47 + 21), presenting with pancytopenia, developed acute megakaryoblastic leukaemia (AMKL). Her bone marrow contained an abnormal clone with a novel dicentric chromosome derived from chromosomes 5 and 7 (karyotype 46, XX, −5, −7, + dic (5; 7) (p 13; p 11·2), +21. This case provides further evidence for a connection between chromosome 21 and this unusual form of childhood leukaemia, and raises questions about the loss of short arm material from chromosomes 5 and 7 compared with the more usual monosomy or long arm loss.

Case report

A two year old caucasian girl with Down’s syndrome presented with a three week history of anorexia, lethargy, and bleeding from the gums. There were no relevant abnormalities on physical examination apart from pallor and bruising. The initial blood count was haemoglobin 4·1 g/dl, white cells 3·31 × 10⁹/l (neutrophils 0·86 × 10⁹/l, lymphocytes 2·28 × 10⁹/l, monocytes 0·17 × 10⁹/l, no blast cells), and platelets 17 × 10⁹/l. A bone marrow aspirate and trephine biopsy specimen showed hypocellularity and pronounced fibrosis with scanty blast cells present. These cells were negative for the lymphoblast markers CD10, DR, and TdT, and had the appearance of megakaryoblasts. Initial treatment was supportive, with red cell and platelet transfusions and antibiotics. A repeat bone marrow trephine biopsy specimen taken one month later showed increased cellularity and included numerous cells with characteristic megakaryoblastic morphology. These cells stained positively with monoclonal antibodies to platelet glycoproteins IB and IIb/IIIa. Acute megakaryoblastic leukaemia (AMKL) was diagnosed. She was treated with daunorubicin, cytosine arabinoside, etoposide, cyclophosphamide, thioguanine and intrathecal methotrexate, and 13 months after diagnosis was in remission.

Cytogenetic studies

The constitutional karyotype ascertained from stimulated lymphocyte cultures was 47 + 21. Trypsin G banding analysis of metaphases from the initial bone marrow aspirate showed a clonal karyotype with a dicentric chromosome resulting from a translocation between the short arm of one chromosome 5 and the short arm of one chromosome 7 (figure). The clonal karyotype was interpreted as 46, XX, −5, −7, + dic (5; 7) (p 13; p 11·2), +21. Lymphocyte cultures stimulated by phytohaemagglutinin and exposed to mitomycin C or mustine hydrochloride had chromosome aberrations in the normal range of

Figure Abnormal dicentric chromosome (dic) shown paired with normal homologues of chromosomes 5 and 7: p—short arm; q—long arm; c—centromere positions.

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frequency, thus excluding Fanconi’s anaemia. Examination of polymorphic variation in the short arms of chromosomes 21 by G and Q banding was unremarkable.

Discussion

AMKL (FAB classification, M7') was, until recently, considered to be a rare form of acute non-lymphocytic leukaemia (ANLL), characterised by the abnormal proliferation of megakaryoblasts. The development of methods for immunological identification of blast cells should, however, increase its recognition. In two recent reports AMKL represented 8% of all leukaemia cases, unselected for age, when antibodies to platelet glycoprotein IIB/IIIA complex and factor VIII/von Willebrand factor complex were used to characterise the cells of megakaryocyte/platelet lineage. It seems likely that many of the previously reported cases of childhood acute myelofibrosis will now be classified as AMKL.

The clinical findings in 20 cases of AMKL in children have been summarised; these may be added eight more cases, and there is also a recent report of Fanconi’s anaemia associated with AMKL. Down’s syndrome was present in five of the above cases; all of these developed AMKL before the age of 4 years. Another patient had a constitutional ring 21 chromosome but was phenotypically normal.

Together with the present report there are 18 childhood cases of AMKL in which karyotype analysis of bone marrow has been performed, including three of the patients with Down’s syndrome. Multiple acquired chromosomal anomalies were found in 15 cases; the most common change was an extra chromosome 21, detected in six of the patients with a constitutionally normal karyotype.

Down’s syndrome is well known to be over represented in the leukaemic population, occurring in 2–5% of unselected leukaemic children and possibly in up to 15% of those presenting with ANLL before the age of 2 years. The proportion of children with AMKL who have Down’s syndrome is higher than in other forms of ANLL (six of 30). An association between childhood acute myelofibrosis and Down’s syndrome has also been described. The occurrence of an additional chromosome 21 in the leukaemic cells of six of 15 of the other cases of AMKL is also particularly common, even though an additional chromosome 21 is a common acquired karyotypic change in acute leukaemia, both in normal children, and in those with Down’s syndrome. Together, these findings provide evidence for a closer association between AMKL and the presence of additional copies of chromosome 21 than occurs in other forms of leukaemia, although bias due to the small number of cases cannot be excluded.

The occurrence of the dic (5; 7) in the case reported here is of interest for two reasons. Firstly, dicentrics affecting two non-homologous chromosomes are relatively uncommon in leukaemia, probably because they are inherently unstable. Secondly, the leukaemic clone was effectively monosomic for most of the short arms of chromosomes 5 and 7. Although loss of material from these two chromosomes is a well recognised phenomenon in ANLL and myelofibrosis, especially when related to previous cytotoxic treatment or radiotherapy, this loss almost always affects either the long arms alone (5q−, 7q−) or the whole chromosome (monosomy 5, 7). In only two previous reports of AMKL have similar abnormalities been described, one with monosomy 7, the other with monosomy 7p resulting from the formation of an isochromosome 7q.

The cause of the association between Down’s syndrome and ANLL in early childhood remains a mystery. As the relative risk compared with that of the normal population falls progressively with age, the predisposition to ANLL may be associated with only a small subgroup of patients with Down’s syndrome. The observation of transient leukaemoid reactions in some infants with Down’s syndrome and the association with the trisomy 21 cells in mosaics suggest that the extra chromosome 21 may be a primary trigger for the oncogenic process. Those mosaics which arise from a post-zygotic mitotic error, are homozygous for two of their chromosomes 21 in the trisomic cells; it would be interesting to determine whether homozygosity for a critical part of chromosome 21 is the rule in children with Down’s syndrome who develop ANLL.

References


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