Clonality analysis suggests that early-onset acute lymphoblastic leukaemia is of single-cell origin and implies no major role for germ cell mutations in parents

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Summary Childhood leukaemia presenting at a young age has been suspected of resulting from a leukaemogenic mutation in parental germ cells, either spontaneously or due to the exposure of a parent to leukaemogenic environmental hazards, particularly ionizing radiation. Mathematical modelling of leukaemogenesis suggests that any such patient would be especially prone to multiple independent leukaemogenic events leading to multiclonality in terms of cell of origin (analogous to bilaterality in familial retinoblastoma). To test this hypothesis we have carried out a search for multiclonal leukaemogenesis in infant and childhood acute lymphoblastic leukaemia (ALL). We used a polymerase chain reaction-based analysis of the X-linked monoamine oxidase A (MAOA) gene locus to study the clonality of marrow samples obtained from female paediatric ALL patients at the time of disease presentation. We obtained presentation samples from 102 patients of whom 72 were found to be informative at the MAOA locus. These included 20 infant leukaemias (< 1 year at diagnosis). Sixty-six samples were found to be unequivocally monoclonal while the remaining six could not, with certainty, be assigned a clonal origin. We also obtained bone marrow aspirates at first relapse as well as at presentation from eight patients. In each case the same pattern of X-linked allelic inactivation was observed at both time points of the course of the disease. No evidence was found for leukemic multiclonality in any age group at presentation or for leukaemic ‘clone-switching’ in relapse. These findings suggest that both infant and childhood ALL is of single-cell origin and implies that leukaemic predisposition resulting from germ cell mutation is unlikely to have a major role in their pathogenesis.

Keywords: clonality; acute lymphoblastic leukaemia; germ cell mutations

Germ line mutations provide a recognized oncogenic pathway for some paediatric solid tumours, notably retinoblastoma and nephroblastoma (Narod et al, 1991). Tumours resulting from germ line mutations have characteristic young age of onset and propensity to multiclonal genis (e.g. bilaterality in retinoblastoma) whether or not familial associations are apparent. Interest in the possible role of leukaemogenic germ cell mutations in childhood leukaemia was stimulated by the ‘Gardner hypothesis’ (Gardner et al, 1990) postulating increased leukaemia risk in the offspring of fathers receiving gonadal irradiation whilst working at the Sellafield nuclear establishment, although this hypothesis is not supported by additional epidemiological evidence now available (Kinlen, 1993; Little et al, 1995). More recently, Shu et al (1994) reported a statistical association between infant leukaemia and preconceptional exposure of fathers to diagnostic X-rays, and postulated a role for radiation-induced germ cell mutations. The possibility of ‘paternal preconception irradiation’ (PPI) having a causal role in leukaemia was reviewed by the Committee on the Medical Aspects of Radiation in the Environment (COMARE) who concluded that ‘there is no doubt that the hypothesis of PPI causing cancer in offspring can be sustained in principle’ although this did not seem to be the explanation for the increased incidence of childhood leukaemia in the vicinity of the Sellafield nuclear Plant (COMARE, 1996). The existence of vertically transmissible leukaemia-predisposing mutant genes has been demonstrated in transgenic mice (Langdon et al, 1986; Heisterkamp et al, 1990; Griffiths et al, 1992) and it has been shown that these may give rise to independently transformed leukaemic clones (Langdon et al, 1986). In very recent experimental studies, Lord et al (1998) have reported that contamination of male mice with the α-emitter ²³⁹Pu results in an increased susceptibility to leukaemia induction by methyl-nitroso-urea in first-generation offspring.

It seems possible that germ cell mutations induced by environmental factors (e.g. natural background radiation or diagnostic exposures), perhaps followed by subsequent insults, could similarly contribute to the pathogenesis of leukaemias and other neoplasms in human populations. Acute lymphoblastic leukaemia (ALL) is rarely familial but ALL in infants (< 12 months) or very young children (< 24 months) may be suspected as genetically predisposed because of age of onset. Infant leukaemias have distinctive properties: prognosis is poor for ALL (Chessels et al, 1994) and characteristic aberrations at chromosome 11q23 are often observed, usually involving rearrangements of the MLL gene (Piu et al, 1995; Greaves, 1996). Such rearrangements are not heritable and have been shown to originate in utero (Ford et al, 1993; Mahmoud et al, 1995), but this does not exclude the
inheritance of a preceding and possibly predisposing mutation. Very recently, Gale et al (1997) have demonstrated by polymerase chain reaction (PCR) methodology the presence at birth of small numbers of leukaemic cells in neonatal ‘blood spots’ of infants who subsequently developed clinically overt leukaemia. This finding demonstrates the prenatal origin of these leukaemias but does not distinguish preconceptual predisposing events from events occurring exclusively in utero.

Here we report clonality studies which are relevant to the role of germ line mutations in the genesis of ALL. The rationale of the study was to seek evidence of multiclonal origin of early-onset ALL analogous to bilateral presentations of retinoblastoma (Rb) in which one Rb gene is mutated in the germ line, leaving the other allele to be mutated somatically (Knudson, 1971; Narod et al, 1991). ALL is also hypothesized to result from two independent mutations (Greaves and Chan, 1986) although the genetic targets are yet to be identified. It is of course possible that this is an underestimate of mutation number, but it is plausible that only a small number of mutations would be required for a neoplasm which presents in early childhood. We have mathematically modelled ‘two hit’ leukaemogenesis with both mutations occurring somatically, or with one transmissible via the germ line. With parameters chosen to simulate the observed age-incidence of ALL, these studies predict that inheritance of the first mutation would cause multiple independent leukaemogenic transformation in the haematopoietic cells of each individual, typically resulting in ten or more independently arising leukaemic clones by time of diagnosis (Wheldon et al, 1997). Multiclonality following a predisposing germ cell mutation is easily understood as resulting from the large number of somatic cells which need only acquire one further mutation to experience malignant transformation. The predicted number of leukaemic clones (> 10) is greater than the number of independently occurring tumours (about 3) estimated for familial retinoblastoma (Knudson, 1971) because the higher incidence of ALL requires higher mutation rates to be assigned to the two-mutation model (Wheldon et al, 1997). As in Knudson’s model, a small proportion of genetically predisposed leukaemias may present (by chance) as a single clone: however, the present modelling studies (Wheldon et al, 1997) suggest that the fraction of heritably predisposed ALL patients who would present in this way is close to zero. Therefore, we expect multiclonality in almost all presenting ALL cases who have inherited a first mutation (provided of course that the two-mutation model is valid). A clinical corollary of this model is that early-onset leukaemias may be difficult to treat effectively because of repeated emergence of new leukaemic clones.

The investigation reported here provides a test of the multiclonality hypothesis and indirectly of the hypothesis that early-onset leukaemias result from the inheritance of one of two possible mutations.

CLONALITY ANALYSIS BY X-CHROMOSOME INACTIVATION

We tested the multiclonality hypothesis experimentally using X-chromosome inactivation analysis. This is based upon the polymorphic differentiation between maternal and paternal X-chromosomes and the discrimination between active and inactive X-chromosomes, by differential methylation. The first such studies, by Fialkow (1976), utilized the expression of a rare polymorphism at the X-linked, glucose-6-phosphate dehydrogenase gene locus. To improve upon the low informative rate associated with this technique, Vogelstein et al (1987) employed Southern blot analysis of differentially methylated, X-chromosomal loci containing restriction fragment length polymorphisms. However, substantial amounts of intact DNA (5–20 mg) are required for such analyses and sufficient quantities are not readily available from pathological specimens. Therefore, the PCR has been applied to enhance the sensitivity of this approach. Informative X-chromosomal regions include sequences of the genes encoding phosphoglycerate kinase (PGK) (Gilliland et al, 1991), androgen receptor (AR) (Allen et al, 1992) and monoamine oxidase A (MAOA) (Hendriks et al, 1992). The associated heterozygosity rates are 32%, 90% and 75% respectively. Because limited amounts of leukaemic sample material were available, PCR methodology was used in the present study. We based our analysis on the MAOA locus because it has a higher rate of heterozygosity than the PGK locus and its reliability is not limited by incomplete digestion of active alleles: a shortcoming associated with AR-PCR (Mashal et al, 1993).

MATERIALS AND METHODS

Reagents

Taq polymerase, restriction enzymes and dNTPs were purchased from Promega (Southampton, UK). Other reagents, unless otherwise stated, were obtained from Sigma (Dorset, UK).

DNA samples

A total of 102 samples of bone marrow (each containing less than 5 mg of DNA) from female childhood ALL patients were received from the Department of Haematology, Sick Children’s Hospital, Yorkhill, Glasgow; the Department of Haematology and Oncology, Institute of Child Health, London; and the Department of Medicine, Section of Hematology and Oncology, University of Chicago. DNA, purified from bone marrow aspirates of chronic myelocytic leukaemia patients, was acquired from Ms Anne Sproul, Haematology Department, Royal Infirmary, Glasgow. Blood and marrow were obtained originally from patients for diagnostic purposes. DNA was also prepared from the peripheral blood samples of 46 healthy, adult, female and five male volunteers.

High molecular-weight DNA was purified from the mononuclear cell fractions of marrow samples as described previously (Sambrook et al, 1989). Ethanol-precipitated DNA was rehydrated in sterile water and DNA concentration was determined by E_{260} measurements and by fluorescence assay using the Hoeschst 33258 method (Cooper and Stratton, 1991).

Determination of clonality

MAOA-PCR

Individuals whose clonality status is assessable by this method are heterozygous with respect to a GT-dinucleotide/variable number tandem repeat (VNTR) region of the MAOA gene (Hendriks et al, 1992). All DNA samples were screened by amplification at the MAOA locus as described below and those found to be informative were subjected to clonality determination.

The method relies upon the presence of several sites recognized by the methylation-sensitive restriction endonuclease HpaII. These are located 5’ relative to a highly polymorphic region of the MAOA gene. Because inactive X-chromosomes are extensively
methylated at these sites, only the active X-chromosomes are cleaved by \textit{HpaII}. The undigested MAOA alleles are PCR-amplified using primers that flank the 5' site of differential methylation and the 3' hypervariable region, which is composed of a VNTR motif and a GT microsatellite. \textit{SacI} digestion produces fragments of 258–388 bp, depending on the number of VNTR and dinucleotide repeats. If \textit{HpaII} digestion is allowed to proceed to completion, PCR amplification and subsequent \textit{SacI} digestion produce one allelic band from monoclonally derived cell samples, whereas two bands of reduced intensity are generated from polyclonal cell populations.

Two reactions were prepared: (a) 100 ng samples of genomic DNA were digested with 5 units of \textit{HpaII}; (b) the same amount of DNA was incubated with enzyme digestion buffer in the absence of enzyme. All reactions were in a total volume of 10 μl and all incubations were carried out at 37°C for 18 h. \textit{HpaII} digestion was terminated by heat inactivation at 65°C for 5 min. All of the reaction mixture was included in the subsequent PCR amplification.

PCR conditions were a modification of those reported (Hendriks et al., 1992). Briefly, amplification was performed in a total volume of 100 μl containing the following components: 1.5 mM magnesium chloride, 50 mM potassium chloride, 10 mM Tris–HCl (pH 9.0) and 0.1% (v/v) Triton X-100. After initial denaturation at 94°C for 5 min, 35 amplification cycles were performed using a Hybrid Omnigene thermocycler (Teddington, UK). Each cycle comprised 1 min at 94°C, 1 min at 54°C and 2 min at 70°C. The following primers were used: 5'-CAATAAATGTCCTACCTT-3' and 5'-ACATTCTAAGCCTAATC-3' (Oswell, Southampton, UK). After amplification, PCR products were extracted with phenol–chloroform, ethanol precipitated and redissolved in 10-μl containing the following components: 1.5 mM magnesium chloride, 50 mM potassium chloride, 10 mM Tris–HCl (pH 9.0) and 0.1% (v/v) Triton X-100. After initial denaturation at 94°C for 5 min, 35 amplification cycles were performed using a Hybrid Omnigene thermocycler (Teddington, UK). Each cycle comprised 1 min at 94°C, 1 min at 54°C and 2 min at 70°C. The following primers were used: 5'-CAATAAATGTCCTACCTT-3' and 5'-ACATTCTAAGCCTAATC-3' (Oswell, Southampton, UK). After amplification, PCR products were extracted with phenol–chloroform, ethanol precipitated and redissolved in 10-μl digestion buffer containing 8 units of \textit{SacI}. Digestion was carried out for 2 h at 37°C, followed by heat inactivation at 65°C for 5 min. To facilitate allele detection, digests were electrophoresed through 4% (w/v) fine resolution Metaphor agarose gel (FMC Bioproducts, Kent, UK) and stained with ethidium bromide. The relative intensities of the bands were determined using photographic negatives of the stained gels. Densitometry was performed on a Molecular Dynamics computing densitometer (Sevenoaks, Kent, UK) and images were analysed using the Quantity 1P Programme (Pharmacia Herts, UK).

**Southern hybridization**

Clonality assessment, by the analysis of differential methylation at the 5' region of the PGK locus using the \textit{BstXI} polymorphism, was performed as described (Vogelstein et al., 1987). Briefly, 6.25 μg of genomic DNA were digested with \textit{PstI} and \textit{BstXI}. After precipitation and redissolution, half of the DNA was digested with \textit{HpaII} and half was left undigested. Fragments were resolved in 1.0% (w/v) agarose gels and blotted on to Hybond N membranes (Amersham International plc, Bucks, UK) before hybridization to a PGK probe. This was an 800-bp \textit{BamHII/EcoRI} fragment from the 5' end of the PGK gene, labelled to a specific activity of 5 × 10^6 cpn μg⁻¹.

**RESULTS**

To ensure a true assignment of clonality, it was essential that \textit{HpaII} digestion of active MAOA alleles, prior to amplification, proceeded to completion. This was verified by the simultaneous processing of male DNA, whose single active X-allele was not detectable after \textit{HpaII} digestion.

The validity of the determination of the clonal status of ALL samples by MAOA-PCR was confirmed by comparison with clonal assessment using Southern blot analysis of the PGK locus. Three chronic myelocytic leukaemia DNA samples (monoclonal) and two normal peripheral blood DNA samples (polyclonal) were analysed by both methods. The results of the Southern blot analysis were, in every case, in agreement with those obtained by MAOA-PCR.

To study the clonal origin of infant ALL, we attempted to PCR amplify at the MAOA locus, samples of mononuclear cellular DNA from bone marrow aspirates collected at the time of disease presentation. Of the samples from 102 female paediatric ALL patients (age range 1 month to 13 years), 72 were heterozygous at the MAOA polymorphic locus. This is equivalent to a heterozygosity rate of 71%. The age distribution of the different categories of ALL patients is shown in Table 1.

Using the MAOA-PCR-based clonality assessment, 66 out of 72 informative presentation ALL samples were found to be monoclonal. The upper allele was inactive in 29 and the lower allele was inactive in 37. The remaining six samples yielded ambiguous results: the allele intensity ratios in the absence of \textit{HpaII} digestion were approximately unity, whereas after \textit{HpaII} digestion, preferential but incomplete loss of one allele was observed resulting in allele intensity ratios greater than 7:1 or less than 1:5.

To determine whether alteration in clonal status occurred during the course of the disease, another means of assessing the origin of leukemic clones was employed. This involved a comparison of the allele inactivation patterns of DNA isolates from presentation and relapse marrow. Thirteen such pairs of samples (patient ages < 2 years) were obtained. Of these, eight were informative. In every case, the inactive allele in presentation marrow was the same as the inactive allele observed in material obtained at the time of relapse.

**DISCUSSION AND CONCLUSIONS**

To test the possible multiclonality of early onset ALL, we studied clonal origin by MAOA-PCR methodology. Assignment of clonality was possible in samples from 27 informative early-onset patients (0.1–2.0 years) and 39 informative, older children (2.1–12.0 years), whereas six samples gave results which could not be confidently evaluated. This is the largest study of clonality by X-chromosome inactivation so far reported for childhood ALL. All of the evaluable patient samples in both age groups were found to be monoclonal. This demonstration of single-cell origin for childhood ALL is in agreement with the findings of an earlier study of eight female children, aged 2–13 years (Dow et al., 1985). The latter determination of clonality was based upon the analysis of X-chromosome-encoded, glucose-6-phosphate dehydrogenase isoenzymes.

A possible difficulty with the interpretation of clonality analysis in leukaemia is that the disease does not present with individually distinct tumours, and a leukaemic cell population which contains

| Diagnosis | < 1 | 1–2 | > 2–13 |
|-----------|-----|-----|--------|
| cALL      | 5   | 4   | 32     |
| Null ALL  | 12  | 3   | 4      |
| Pre B-ALL | 2   | 0   | 2      |
| T-ALL     | 1   | 0   | 1      |

Table 1 Age distribution and diagnosis of informative ALL patients

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several distinct clones may be numerically dominated at presentation by a single clone – the earliest to arise, or possibly the most rapidly growing – giving a spurious finding of monoclonality, if the co-existing clones fall below the detection threshold of the assay. It is difficult to exclude this eventuality on the basis of a single sample for each patient. However, we had available to us 66 evaluable patients, including 20 infants, and it would be surprising that we were unable to detect clonal co-existence in any of them if true multiclonality were in fact a common occurrence.

An alternative strategy for detection of multiclonal leukaemogenesis requires the provision of sequential marrow samples from the same patient, i.e. at first presentation and at subsequent relapse. This provides the opportunity for ‘clone-switching’ in a multiclonal setting – the clone present at relapse being different from that occurring at first diagnosis, either because the selection of a different (pre-existing) clone has been favoured by therapy or because a new independent clone has arisen subsequent to therapeutic eradication of the initial clone. We obtained presentation and relapse marrow samples from 13 patients < 2 years in this series, eight of whom proved to be informative. In all eight cases, the clone present at relapse did not differ in relation to X-inactivation from the original clone, providing no evidence of multiclonality. It is not possible to calculate precisely the probability of this occurring by chance, because we have no way to quantify the probability that a co-existing minor clone would be resistant to therapy, or that the original dominant clone had been eliminated. Suppose, however, that the original clone were indeed eliminated, and that relapse is due to a newly arising clone (on our mathematical model this is quite a common event). If the newly arising clone had an equal chance of either X-chromosome’s being inactivated, then the probability of observing the same pattern of X-inactivation at presentation and relapse in each of eight patients is 0.58, or 0.008. Despite the restrictive assumptions of the calculation, it seems unlikely that we would have failed to observe ‘clone-switching’ in any of eight cases, if multiclonal leukaemogenesis were usually occurring. A further area of uncertainty concerns the mathematical model which predicts multiclonality were a germ cell mutation to be inherited (Wheldon et al., 1997). We have previously mentioned that more than two multiclonality were a germ cell mutation to be inherited (Wheldon et al., 1997). We have previously mentioned that more than two

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