Impaired pubertal growth in acute lymphoblastic leukaemia

M Uruena, R Stanhope, J M Chessells, A D Leiper

Abstract
The growth of 182 patients who were long term survivors of childhood acute lymphoblastic leukaemia was retrospectively analysed. All remained in first remission and were treated with either 1800 or 2400 cGy of cranial irradiation. None had been treated with either testicular or spinal irradiation. Ninety three (51 boys, 42 girls) were treated with 2400 cGy and 89 (42 boys, 47 girls) were treated with 1800 cGy cranial irradiation. All patients were treated with standard chemotherapy including intrathecal methotrexate in similar dose regimens in either group. Mean age (SD) at diagnosis in the group treated with 2400 cGy was 4.8 (2.6) years and mean age in the group treated with 1800 cGy was 6.5 (3.3) years. Mean height SD score at diagnosis in the 2400 cGy group was +0.29 and final height achieved was −0.63. Mean height SD score at the start of treatment in the group treated with 1800 cGy was +0.40 and mean final height was −0.53. There was a similar reduction in height SD score in both groups during the pubertal growth spurt.

The decrement in height SD score was greater when treatment was administered at less than 7 years of age in either dose regimen, both in prepubertal and pubertal growth. However, the decrease in height SD score was found to be greater in girls than boys. There was a trend in both sexes for the onset of puberty to be at a younger age with a lower treatment dose of radiotherapy. However, in girls treated with the lower dose regimen there was a significant reduction in the mean age of onset of puberty which was 9.9 years. Our data suggest that girls treated at less than 7 years of age have a severe impairment of pubertal growth, which is probably a combination of the dual endocrinopathy of premature puberty and growth hormone insufficiency.

Long term survival for children with acute lymphoblastic leukaemia has improved enormously in recent years, partly due to the introduction of treatment to the central nervous system,1 2 which is given as a combination of intrathecal methotrexate and cranial irradiation.3 Because of the morbidity of cranial irradiation, the dose administered has been reduced in recent years from 2400 cGy in 15 fractions over three weeks to 1800 cGy in 10 fractions over two weeks.4

Cranial irradiation may cause pituitary dysfunction, of which growth hormone insufficiency is the most common endocrinopathy. This is related to both the administered dose of irradiation and the method of fractionation.5 6 The effect of the irradiation is delayed and progressive. It has been reported that regimens of low dose cranial irradiation may cause growth hormone insufficiency and growth failure7 8 in children with acute lymphoblastic leukaemia. The degree of growth suppression and whether this requires growth hormone replacement treatment has been controversial.9 10 Recent data have even suggested that final height is not altered in children treated for acute leukaemia.11 Low dose cranial irradiation in children with acute lymphoblastic leukaemia has been reported to be associated with central premature and precocious puberty,12-14 predominantly affecting girls.15 However, the significance of this for ultimate height is unknown.

Between 1970 and 1980 the dose of cranial irradiation administered was 2400 cGy but this was superseded in 1981 by a dosage of 1800 cGy. During the last 20 years treatment has altered in that induction and consolidation has become more intensive. However the basic drugs used, including intrathecal methotrexate and maintenance treatment (6-mercaptopurine, methotrexate, prednisolone, and vincristine), have essentially remained unaltered. We have been able to compare the onset of puberty and the pubertal growth spurt in children treated by either of these two irradiation dose regimens.

Patients and methods

PATIENTS
Data from 182 patients were analysed in this study from a total cohort of 307 patients who were survivors of acute lymphoblastic leukaemia diagnosed at the Hospital for Sick Children between 1970 and 1986.16 All patients were in first remission, had been off treatment for two years or more, and had attained the onset of puberty at the time of this study. None had received gonadal or spinal irradiation. Twelve children (seven treated with 1800 cGy, five treated with 2400 cGy) were treated with growth hormone and/or gonadotrophin releasing hormone (GnRH) analogue during the course of this study and were excluded, as well as those with dysmorphic syndromes or abnormal karyotypes.

Ninety three patients (51 boys, 42 girls) were treated with 2400 cGy cranial irradiation and 89 (42 boys, 47 girls) were treated with 1800 cGy. Cranial irradiation was usually given within eight weeks of diagnosis or occasionally delayed until the patient was 2 years of age. Chemotherapy regimens have been previously
The numbers of patients at diagnosis, onset of puberty, and final height are indicated in fig 1. Because the group treated with 2400 cGy was before 1980, a much larger number of patients had attained final height than in the lower dose group.

METHODS
Clinical records of the patients were examined retrospectively including height obtained at diagnosis (equivalent to the onset of treatment), the onset of puberty, and final height. Stature was expressed as SD scores with comparison of data from Tanner et al.20 21 No adjustment was made for secular trend in height. The onset of puberty was taken as the acquisition of breast development in girls (B2) and the development of either stage 2 genitalia and/or 4 ml testicular volume in boys. Normal data for the ages of attainment of sexual maturation were from Marshall and Tanner.22 23 The patients were divided into two groups according to the two dose regimens of cranial irradiation used as well as the age at onset of acute lymphoblastic leukaemia. Statistical measurements were performed using paired and independent t test.

Results
Data for change of height SD score during puberty are shown in fig 1. The pattern of reduction of height SD score during both prepuberty and puberty was similar in both groups. However, this was not significant in the 1800 cGy irradiation group, which probably reflects the smaller proportion of patients who had attained final height. Age at onset of puberty was greater by 0-4 years in the children treated with the higher dose regimen but this may be a reflection of a secular trend.

The decrement in height SD score, especially during puberty, was greater when cranial irradiation was administered at less than 7 years of age in both dose regimens (fig 2). In contrast, children treated for acute lymphoblastic leukaemia at the age of 7 years or above were more...
Figure 3  Height SD score in 93 (51 boys, 42 girls) and 89 (42 boys, 47 girls) patients treated for acute lymphoblastic leukaemia with 2400 cGy and 1800 cGy cranial irradiation respectively, before and after 7 years of age. Horizontal bars represent 1SD. The number of patients are indicated.

Figure 4  Age at the onset of puberty in 69 girls (B2) and 63 boys (4 ml testicular volume) treated for acute lymphoblastic leukaemia under the age of 7 years with either of two dose regimens of cranial irradiation (2400 or 1800 cGy). Vertical bars represent 1SD. Data for normal girls and boys from Marshall and Tanner 1969 and 1970 respectively. 

Discussion

The main difficulty in the interpretation of this study arises from the fact that we have compared different treatment groups from 1970 until 1986, and this study is in series and not in parallel. No adjustment has been made for secular change in either stature or for the timing of puberty. However we believe that, even considering these difficulties, our data are relevant. Furthermore the exclusion of 12 patients who received endocrine treatment for
growth failure increases the significance of our findings. Broomhall and colleagues demonstrated in a large number of patients that children with acute lymphoblastic leukaemia were significantly taller than the normal population with a mean SD score of +0.49.24 We found that at diagnosis our patients were taller than average. A possible explanation is that leukaemia is a disease of the higher social classes and the latter have greater than average stature.25 It is appreciated that the treatment of brain tumours with high dose cranial irradiation is associated with a high incidence of growth hormone insufficiency/deficiency.2 In our data the decrement in final height was similar with both 2400 cGy and 1800 cGy cranial irradiation. Our observations are in agreement with Shalet in that the initial endocrinopathy associated with cranial irradiation is growth hormone insufficiency.3 It has been reported by previous authors that growth hormone secretion in young children is more sensitive to damage by a given dose of irradiation than in older children.26 The effects of irradiation on pituitary function tend to be delayed and progressive over several years.3 Certainly children treated with cranial irradiation under the age of 7 years have a higher incidence of growth failure than older children26 27 and our data are in agreement with this.

Cranial irradiation of either 2400 or 1800 cGy affects growth in prepuberty as well as in puberty. In normal children growth in prepuberty is a continuous deceleration, and there is evidence of a progressive decrease in growth hormone secretion until the onset of the growth spurt.28 During the pubertal growth spurt growth is dependent on both sex steroids and growth hormone and one without the other produces an inadequate growth acceleration.29 It has been suggested that growth hormone secretion in children treated with low dose cranial irradiation is predominantly decreased during puberty rather than prepuberty.28 Our data suggest that overall, prepubertal growth is affected almost as much as pubertal growth, but when the groups were subdivided by age, a different picture emerged. There was a significant decrease in height SD score between the onset of puberty and final height attainment in both radiation groups diagnosed under the age of 7 years, but not during prepubertal growth in the 1800 cGy group. This suggests that the larger dose of 2400 cGy affected both prepubertal and pubertal growth but 1800 cGy given at a young age predominantly caused loss of the growth spurt. Of course, the majority of the population with leukaemia are diagnosed under the age of 7 years and indeed in this study the onset of leukaemia was below this age in 72% of the children. The latter situation of impairment of the pubertal growth spurt in those children receiving 1800 cGy at less than 7 years of age will thus be the commonest prevailing picture at the present time.

Recent data have suggested that low dose cranial irradiation is associated with precocious or early puberty in girls rather than boys and our present analysis support this. Of interest from our study is the suggestion that earlier puberty is the result of the lower dose regimen of 1800 cGy rather than 2400 cGy cranial irradiation. Certainly early puberty could be a contributory reason as to why growth impairment in the girls was greater than in the boys. Alternatively it may also be that growth hormone secretion is more adversely affected in girls than boys in the same way that gonadotrophin secretion is affected differently between the sexes. We know no other supportive data for such a hypothesis. The most important single growth hormone treatment for children treated at a young age as they show the greatest degree of growth retardation. As well as the possibility of growth hormone treatment for this group, it may be advantageous to therapeutically manipulate the timing of pubertal maturation using a GnRH analogue. There is now some evidence to suggest that a combined regimen of both growth hormone and GnRH analogue may be advantageous.

In conclusion, we suggest that all children who are treated for acute lymphoblastic leukaemia with cranial irradiation, especially girls treated at a young age, have their growth carefully monitored both during prepuberty and especially during puberty. Certainly the growth failure induced by cranial irradiation at a low dose regimen may be significant and require appropriate treatment. Our data indicate which groups of children by age and sex are likely to require special attention.

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... and while on the subject

Accepted treatment for Kawasaki disease is with aspirin and intravenous gammaglobulin. An American multicentre randomised trial reported in 1986 showed that coronary artery dilatations were less common after treatment with gammaglobulin and aspirin than after aspirin alone.¹ In that trial the gammaglobulin was given as four consecutive daily doses of 400 mg/kg body weight. Now the same workers have shown that a single infusion of 2g/kg is equally if not more, effective (Jane W Newburger and colleagues, New England Journal of Medicine 1991;324:1633–9). Clinical and laboratory indices of inflammation also resolved quicker with the single large dose.

An editorial in the same issue of the journal points out that there are still unresolved questions.² The main end point in these trials has been coronary dilatation of any severity seen on echocardiography. Dilatation of less than 6 mm is apparently relatively benign and it is the giant aneurysms which lead to myocardial infarction and death. The effect of gammaglobulin on mortality is not known. Nevertheless in the most recent trial none of the 254 children who had normal coronary arteries when first seen and who received the single large infusion of gammaglobulin developed giant aneurysms. All of the patients in this study were treated within 10 days of the onset of the disease. The effect of treatment later in the course of the disease in unknown but it seems reasonable to treat if the disease process is still active.²

The role of aspirin is uncertain. Continuous low dose aspirin seems indicated if there is persisting coronary abnormality but whether initial high dose treatment is necessary is not known.

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