Adolescent growth and pubertal progression in the Silver-Russell syndrome

P S W DAVIES,* R VALLEY,† AND M A PREECE*

*Department of Growth and Development, Institute of Child Health, London, and †Department of Anaesthesia, Hospital for Sick Children, Toronto, Canada

SUMMARY The pattern of growth and development of 18 adolescent children with the Silver-Russell syndrome was studied. Mature height was about −3.6 standard deviation scores in both sexes. This is comparable to the height reduction at diagnosis, which has been reported previously. The pattern of puberty and adolescent growth was essentially normal and occurred at a marginally earlier time than normal, although in this small series the differences were not significant. In the girls there was a tendency to gain subcutaneous fat after puberty. This trend was not apparent in the boys.

Of the recognisable growth disorders presented to the paediatrician one of the more common is the Silver-Russell syndrome. The characteristic physical features of this syndrome were described independently but almost simultaneously.1 2 Such patients are usually thin, with short stature noticeable from birth. There are a number of dysmorphic features such as a small and triangular face, lowset ears, and clinodactyly.

The work of Tanner et al,3 described in detail the natural history of growth in this disorder. As these authors stated, however, few of their patients had reached maturity and thus a quantitative description of the later stages of longitudinal growth was not possible. Since that time a number of the patients have reached physical maturity, and sufficient auxological data have been collected to allow a longitudinal analysis of physical growth in this syndrome.

Methods

The criteria adopted by Tanner et al1 for the diagnosis of Silver-Russell syndrome were: (a) stature that was two standard deviations below the 50th centile for height on British standards; (b) birth weight when adjusted for sex, length of gestation, birth order, and maternal height two standard deviations or more below the 50th centile on the Tanner-Thomson standards4; and (c) absence of any other recognisable syndrome, and of chromosomal, endocrine, metabolic, chondrodysplastic, or psychiatric disturbances that would account for the short stature.

The clinical records and existing growth data of the patients who had satisfied those criteria were reviewed. The original diagnosis of Silver-Russell syndrome was reassessed. In addition to the criteria adopted by Tanner et al,3 patients had to exhibit the classical features of this syndrome—for example, clinodactyly, a triangular face, and lowset ears. Patients to be included in this study were then selected by virtue of firstly, having stopped growing in the sense that they had grown less than 1 cm in the last year of measurement and secondly, that their growth had been measured for at least three years before the onset of puberty. These selection criteria yielded a total of 18 patients, nine of each sex.

Using the longitudinal growth data produced from these patients it was possible to describe the mean growth curve of a number of linear dimensions such as height and sitting height. The growth curves of such dimensions have a characteristic and similar shape in normal individuals. When attempting to describe the ‘typical’ or ‘mean’ growth curve of a specific group of individuals, however, account must be taken of the variability in timing, duration, and intensity of biological events; in particular the adolescent growth spurt. If this variation in the tempo of growth is not accommodated the growth curve found by simply taking the mean of individuals’ measurements at different ages will not be representative of the true mean curve. In such a growth curve the duration of the adolescent growth spurt is lengthened and its intensity reduced.

In the study reported here, as we wished to study specifically the growth period around puberty, special attention was paid to this phase-effect. In the
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past this problem has been overcome by centering individual growth curves on discernable common events notably the age at peak height velocity (PHV). An alternative approach is to apply a mathematical model to the individual growth curves and use the mean values of the mathematical parameters that describe the individual curves to produce a mean constant curve.6

The growth curve for many anthropometric variables can be defined by the model described by Preece and Baines that used five parameters, each being associated with meaningful biological events.7 This mathematical model is:

\[ h = h_1 - \frac{2(h_1 - h_0)}{\exp[s_0(t-\theta)] + \exp[s_1(t-\theta)]} \]

where \( h \) is height at age \( t \), \( h_1 \) is final or adult height, \( s_0 \) and \( s_1 \) are rate constants, \( \theta \) is a time constant, and \( h_0 \) is height at age=0. It is an improvement on previous models that only considered parts of the growth curve8 9 or produced many, sometimes biologically meaningless, parameters.10 This method of growth modelling was applied to the longitudinal data of each child for stature, sitting height, and subischial leg length.

A constant component of the Silver-Russell syndrome is that patients are thin, with skinfold thicknesses being appreciably below normal. By the nature of their growth curves measurements of skinfolds do not easily lend themselves to curve fitting. In order to describe the changes in these measurements that occur throughout growth, standard deviation scores (SDS) were calculated for the skinfold measurements at the tricep and subscapular sites, allowing for age and sex differences. These scores were determined by: 

\[ \text{SDS} = \frac{X - \bar{x}}{Sx} \]

where \( X \) is the measurement, \( \bar{x} \) is the mean of the measurement, and \( Sx \) is the standard

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Table 1  Biological parameters derived from the Preece-Baines curves pertaining to stature

<table>
<thead>
<tr>
<th></th>
<th>Boys Mean (SD)</th>
<th>Girls Mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult height (cm)</td>
<td>150·7 (2·8)</td>
<td>142·0 (6·2)</td>
</tr>
<tr>
<td>Size at peak height velocity (cm)</td>
<td>135·7 (3·2)</td>
<td>127·5 (5·5)</td>
</tr>
<tr>
<td>Age at peak height velocity (years)</td>
<td>13·6 (1·0)</td>
<td>11·6 (1·4)</td>
</tr>
<tr>
<td>Velocity at peak height velocity (cm/year)</td>
<td>8·3 (1·5)</td>
<td>8·0 (1·4)</td>
</tr>
</tbody>
</table>

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Fig 1  Mean constant curves for height attained in boys and girls plotted on British standards.
deviation of the measurement. The standardising data were taken from the data of Tanner and Whitehouse.\textsuperscript{11} Standard deviation scores for both skinfold sites were calculated from the data existing closest to three years before the age at PHV, at age of PHV, and three years after this event.

A disturbance of pubertal events has been cited as occurring in some cases of Silver-Russell syndrome.\textsuperscript{1} To obtain some indication of the pattern of development of sexual characteristics for the children in this study the ages of attainment of puberty stages were recorded for each child.\textsuperscript{12}

Results

Table 1 shows the means and standard deviations of some of the biological parameters derived from the Preece-Baines curve, which were obtained from the individual fits to the stature data of the boys and girls.

In both sexes adult height was well below the third centile for normal British children. Indeed, the standard deviation scores of adult height for boys and girls was $-3.61$ and $-3.58$, respectively. Fig 1 shows the mean constant curves for stature for both sexes. Fig 2 shows the height velocity curves derived from these and plotted on standards for normal British children.

Table 2 shows the derived biological parameters for sitting height and subischial leg length for both sexes. Adult sitting height was $-3.29$ standard

<table>
<thead>
<tr>
<th>Boys</th>
<th>Girls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sitting height</td>
<td></td>
</tr>
<tr>
<td>Adult sitting height (cm)</td>
<td>82.7 (1.6)</td>
</tr>
<tr>
<td>Size at peak sitting height velocity (cm)</td>
<td>74.3 (1.4)</td>
</tr>
<tr>
<td>Age at peak sitting height velocity (years)</td>
<td>14.0 (1.0)</td>
</tr>
<tr>
<td>Velocity at peak sitting height velocity (cm/year)</td>
<td>5.1 (1.0)</td>
</tr>
<tr>
<td>Subischial leg length</td>
<td></td>
</tr>
<tr>
<td>Adult subischial length (cm)</td>
<td>68.4 (2.2)</td>
</tr>
<tr>
<td>Size at peak leg length velocity (cm)</td>
<td>61.4 (2.0)</td>
</tr>
<tr>
<td>Age at peak leg length velocity (years)</td>
<td>13.0 (0.9)</td>
</tr>
<tr>
<td>Velocity at peak leg length velocity (cm/year)</td>
<td>4.7 (1.3)</td>
</tr>
</tbody>
</table>

![Fig 2](Image) Mean constant curves for height velocity in boys and girls plotted on British standards.
deviation scores below the mean in the boys and
-3.08 for the girls. Adult subischial leg length was
-3.06 standard deviation scores below the mean in
the boys and -2.64 below the mean in the girls.

The mean values of the standard deviation scores
at the triceps and subscapular skinfold sites in both
sexes are shown in table 3. In all but one case the
mean skinfold standard deviation score is below the
average for normal British children.

Mean age of attainment for each stage of pubertal
development and mean age of menarche are shown
in table 4. The centile position of these ages as
compared with normal British children is shown in
the third column. As would be expected the girls
show pubertal changes at a slightly earlier age than
the boys.

Table 3 Mean standard deviation scores of two skinfolds
(SEM)

<table>
<thead>
<tr>
<th></th>
<th>Triceps</th>
<th>Subscapular</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boys</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 years before peak</td>
<td></td>
<td></td>
</tr>
<tr>
<td>height velocity</td>
<td>-0.13 (0.17)</td>
<td>-0.03 (0.18)</td>
</tr>
<tr>
<td>At peak height velocity</td>
<td>-0.23 (0.24)</td>
<td>-0.04 (0.17)</td>
</tr>
<tr>
<td>3 years after peak</td>
<td></td>
<td></td>
</tr>
<tr>
<td>height velocity</td>
<td>-0.11 (0.25)</td>
<td>-0.09 (0.20)</td>
</tr>
<tr>
<td>Girls</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 years before peak</td>
<td></td>
<td></td>
</tr>
<tr>
<td>height velocity</td>
<td>-0.50 (0.17)</td>
<td>-0.25 (0.14)</td>
</tr>
<tr>
<td>At peak height velocity</td>
<td>-0.33 (0.20)</td>
<td>-0.15 (0.18)</td>
</tr>
<tr>
<td>3 years after peak</td>
<td></td>
<td></td>
</tr>
<tr>
<td>height velocity</td>
<td>-0.13 (0.31)</td>
<td>+0.03 (0.17)</td>
</tr>
</tbody>
</table>

Table 4 Attainment of puberty stages and age of menarche
(SEM). The centile position of each age is shown in the third
column. Conventionally centile positions below the 50th
centile indicate delayed attainment of a puberty stage

<table>
<thead>
<tr>
<th>Stage</th>
<th>Mean age of attainment</th>
<th>Centile position</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Boys</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pubic hair 2</td>
<td>12.52 (0.35)</td>
<td>~50th</td>
</tr>
<tr>
<td>Pubic hair 3</td>
<td>13.58 (0.28)</td>
<td>~50th</td>
</tr>
<tr>
<td>Pubic hair 4</td>
<td>14.57 (0.35)</td>
<td>~50th</td>
</tr>
<tr>
<td>Genitalia 2</td>
<td>12.58 (0.33)</td>
<td>25th-50th</td>
</tr>
<tr>
<td>Genitalia 3</td>
<td>13.68 (0.41)</td>
<td>~25th</td>
</tr>
<tr>
<td>Genitalia 4</td>
<td>14.88 (0.45)</td>
<td>10th-25th</td>
</tr>
<tr>
<td>Girls</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pubic hair 2</td>
<td>11.60 (0.46)</td>
<td>~50th</td>
</tr>
<tr>
<td>Pubic hair 3</td>
<td>12.51 (0.49)</td>
<td>~50th</td>
</tr>
<tr>
<td>Pubic hair 4</td>
<td>13.04 (0.60)</td>
<td>~50th</td>
</tr>
<tr>
<td>Breast 2</td>
<td>11.47 (0.61)</td>
<td>~50th</td>
</tr>
<tr>
<td>Breast 3</td>
<td>12.61 (0.50)</td>
<td>25th-50th</td>
</tr>
<tr>
<td>Breast 4</td>
<td>13.22 (0.66)</td>
<td>~50th</td>
</tr>
<tr>
<td>Menarche</td>
<td>12.98 (0.59)</td>
<td>~50th</td>
</tr>
</tbody>
</table>

Discussion

The aim of this work was to describe some of the
features of longitudinal growth in the Silver-Russell
syndrome. For a number of patients sufficient
auxological data have been collected to produce
growth curves and to fit a mathematical model to
those curves. Therefore, the mean constant curve
for a number of body dimensions can be described.
Standard deviation scores were calculated for tricep
and subscapular skinfold measurements and the age
of attainment of puberty stages were also recorded.

The major presenting feature of this syndrome—
short stature—affects both sexes to the same degree.
The slight difference in the adult height standard
deviation scores between boys and girls is not
significant (p>0.05). The mean height standard
deviation score at referral found by Tanner et al1
was -3.58, remarkably similar to the reduced adult
height found in this study. This indicates that there
is little catch up growth in the Silver-Russell
syndrome during childhood and adolescence. It may
also confirm the observation of Tanner et al3 that
growth essentially proceeds normally in childhood;
the patient, however, always remains small in
comparison with his or her peers. In contrast to this
it has been reported in a recent publication that five
out of 15 patients diagnosed as having Silver-Russell
syndrome experienced catch up growth.15 It should
be noted, however, that a number of the patients
described by these workers would not have satisfied
the diagnostic criteria adopted in our study.

The Preece-Baines curve produces a number of
biological parameters pertaining to the adolescent
growth spurt. An abnormal pattern and timing of
pubertal events was originally reported by Silver et
al as one of the features of the syndrome.1 Age at
PHV in both sexes was about six months earlier than
for normal British children. Nevertheless, as the
standard deviation of the timing of this event is
about 0-9 of a year in both sexes,14 the age at
PHV found in this work is well within normal limits.
The actual velocity of growth at this time, 8-3
cm/year in the boys and 8-0 cm/year in the girls, is
also well within normal expectations, although slightly
below the mean velocity found for British
children. Other biological parameters derived from
the mean constant curve such as the age at take off
(the beginning of the adolescent growth spurt) and
the velocity at take off are also within normal
ranges, although again slightly reduced. Consequently,
all these factors tend to indicate an essentially
normal adolescent growth spurt that is reduced
slightly in magnitude and occurs slightly early. This
description of the adolescent growth spurt is
supported by the height velocity curves shown in fig 2.
The standard deviation scores for adult sitting height and subischial leg length would suggest that leg length is reduced to a lesser degree than sitting height. This difference, however, is not significant at the 5% level and therefore we cannot conclude that adults with Silver-Russell syndrome are disproportionate. In normal British children leg length reaches its peak velocity on average about six months earlier than the trunk. Also, the spurt at adolescence in sitting height is greater than in the lower limbs. In the group of children included in this study peak leg length velocity is roughly one year in advance of peak sitting height velocity. This difference is significant at the 5% level in the boys but not in the girls. The girls have a greater peak velocity in leg length than in sitting height and although the difference between the peak velocities is not significant, this pattern is very similar to that seen in normal adolescents.

The values of the mean standard deviation scores for the triceps and subscapular skinfold measurements shown in table 3 are similar to those reported for the triceps skinfold site alone by Tanner et al. The mean standard deviation scores are closer to the mean for the normal population in this present group than is usually found in younger children with Silver-Russell syndrome. The data presented here support the description that body composition tends to normality in later childhood and adolescence.

The mean age of attainment of puberty stages as shown in table 4 shows that for the group of children in this study the mean ages are similar to those found in normal British children, with the ages lying on or close to the 50th centile. Mean age of menarche in the nine girls studied here is remarkably close to the mean age of 13-0 years usually cited for normal British girls. Age of attainment of genitalia stages 2 to 4 in boys occur progressively later in comparison with normal British children. The 50th centile for time spent between these stages is 1-9 years in normal children, while in this study the time spent between the stages was 2-3 years. Nevertheless, this value is still within the normal range. Thus we must conclude that there is a normal pubertal development of sexual characteristics in this syndrome. The belief that there is a disturbance of pubertal development in Silver-Russell syndrome stems from early descriptions of the growth disorder. Since that time more information has become available regarding normal variation in timing of pubertal events and it now seems that adolescent development is essentially normal. Nevertheless, there are still contradictory reports in the literature pertaining to the normality of pubertal events. In a recent report a potential relation between intrauterine growth retardation and an early adolescence has been cited, while the timing of pubertal events has been reported as normal in an equally recent work.

It has been postulated that in the Silver-Russell syndrome intrauterine growth retardation results in a reduction in total body cell mass, and that after birth growth proceeds normally with the child always remaining small in comparison with his or her peers. This study would certainly lend support to the normality of physical growth. Puberty occurs essentially normally both in terms of the adolescent growth spurt and the development of sexual characteristics.

The characteristic appearance of many patients with this syndrome aids in diagnosis, however, the prognosis with regard to adult stature is poor. Human growth hormone has been shown to have variable and short term effects on growth velocity, with some patients showing growth acceleration. Further trials over longer periods are now required and any long term benefit can be judged against the data reported here.

References

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Correspondence to Mr PSW Davies, Dunn Nutrition Unit, Downhams Lane, Milton Road, Cambridge CB4 1XJ.

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