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Final height in central precocious puberty after long term treatment with a slow release GnRH agonist

Wilma Oostdijk, Berthon Rikken, Sandra Schreuder, Barto Otten, Roelof Odink, Cathrinus Rouwé, Maarten Jansen, Willem Jan Gerver, Johan Waelkens, Stenvert Drop

Abstract

Objective—To study the resumption of puberty and the final height achieved in children with central precocious puberty (CPP) treated with the GnRH agonist triptorelin.

Patients—31 girls and five boys with CPP who were treated with triptorelin 3.75 mg intramuscularly every four weeks. Girls were treated for a mean (SD) of 3.4 (1.0) years and were followed up for 4.0 (1.2) years after the treatment was stopped.

Results—The rate of bone maturation decreased during treatment and the predicted adult height increased from 158.2 (7.4) cm to 163.9 (7.5) cm at the end of treatment (p < 0.001). When treatment was stopped bone maturation accelerated, resulting in a final height of 161.6 (7.0) cm, which was higher than the predicted adult height at the start of treatment (p < 0.001). Height at the start of treatment was the most important factor positively influencing final height (r = 0.75, p < 0.001). Bone age at cessation of treatment negligibly influenced final height (r = -0.52, p = 0.03). A negative correlation between bone age and height increment after discontinuation of treatment was observed (r = -0.85, p = 0.001). Residual growth capacity was optimal when bone age on cessation of treatment was 12 to 12.5 years. Body mass index increased during treatment and remained high on cessation. At final height, the ratio of sitting height to subischial leg length was normal. Menarche occurred at 12.3 (1.1) years, and at a median (range) of 1.1 (0.4 to 2.6) years after treatment was stopped. The ovaries were normal on pelvic ultrasonography.

Conclusions—Treatment of CPP with triptorelin increases final height, with normal body proportions, and seems to increase body mass index. The best results were achieved in girls who were taller at the start of treatment. Puberty was resumed after treatment, without the occurrence of polycystic ovaries.

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Keywords: central precocious puberty, GnRH agonist, final height.

For more than a decade children with central precocious puberty (CPP) have been treated with gonadotrophin releasing hormone (GnRH) agonists with variable potencies and different routes of administration. GnRH agonist treatment causes hormonal suppression and thereby influences secondary sexual characteristics, growth, and bone maturation.

Some investigators suggest that delaying puberty does not improve final height prognosis, as the decreased rate of epiphyseal maturation is offset by a decrease in growth hormone secretion and growth rate.

Recently, various studies reporting final height and the reversibility of the gonadal axis on discontinuation of treatment have been published. Most of these results are based upon agonist treatment given by daily subcutaneous injections or intranasal application. It was suggested that auxological improvement depends on a continuous and full suppression of gonadal steroid production. Furthermore, improved results have been documented in children with young (bone) ages.

We have shown that treatment of children with CPP with the slow release GnRH agonist triptorelin is effective in continuous suppression of gonadal activity.

In this paper we present the data on the resumption of puberty and the adult height attained in children with CPP who were treated with the slow release GnRH agonist triptorelin.

Methods

PATIENTS

Thirty six patients (31 girls and five boys) with CPP achieved a final height following treatment with triptorelin (D-Trp6-LHRH) at a mean dose of 96 (SD 25) μg/kg intramuscularly repeated every four weeks. The diagnostic criteria for CPP have been described in the past and are summarised as follows: the appearance of secondary sexual characteristics before the age of 8 and 9 years in girls and boys respectively, a bone age/height age ratio > 1, including a pubertal luteinising hormone (LH) response in a GnRH test. Twenty eight patients (25 girls and three boys) received only triptorelin, whereas eight patients (six girls and two boys) were treated with buserelin subcutaneously (27–30 months) or cyproterone acetate orally (6–40 months) before starting triptorelin.

A computerised tomographic (CT) scan or magnetic resonance imaging (MRI) of the brain, or both, was performed in all patients: in...
24 girls CPP was considered idiopathic, in four girls there was infantile encephalopathy, two girls had been operated on for an astrocytoma, and in one girl a hamartoma was found. In none of the girls with organic CPP was growth hormone secretion evaluated. Three of the five boys were suffering from neurofibromatosis, one boy was operated on for an arachnoid cyst, and in one boy the CPP was idiopathic. None of the boys with neurofibromatosis underwent radiotherapy or surgery. Because no growth impairment was observed in these three boys, growth hormone secretion was not evaluated. The boy operated on for an arachnoid cyst was evaluated before GnRH agonist treatment was started, and a normal growth hormone response to arginine stimulation (GH >20 mU/l) was shown.

The patients were treated for at least two years. After discontinuation of triptorelin they were evaluated each subsequent year until the final height was achieved. Final height was considered to be attained when the bone age was $\geq 15.0$ years in girls and $\geq 16.0$ years in boys or when height velocity was $<1.0$ cm/year for a minimum period of one year, or both. Once the final height was attained, pelvic ultrasonography was performed using Acuson equipment (Mountain View, California, USA) with 3.5 or 5 MHz transducers.

**Growth Evaluation**

On initiating the study and subsequently every three months during the treatment and every six to 12 months after the treatment was discontinued, the patients underwent a physical examination including measurement of height, sitting height, and weight, and recording of secondary sexual characteristics. Height was expressed as a quotient using the body mass index (BMI) (weight/height$^2$) and BMI (all at the start of treatment); bone age at the time of discontinuation of treatment; target height, duration of treatment, and chronological age at attainment of final height. Written informed parental consent was obtained before the start of treatment. The study was approved by the ethics committees of the participating university hospitals.

**Results**

During treatment, regression in secondary sexual characteristics occurred because prepubertal oestadiol and testosterone levels were attained in girls and boys, respectively, as has been reported in the past. No local or systemic side effects were observed. One girl developed type 1 diabetes mellitus several months after discontinuation of triptorelin.

**Auxological Evaluation**

No significant differences in chronological age, bone age, height, predicted adult height at the start of treatment, or in final height and target height were found between the group of previously untreated patients compared to the previously treated group (Mann-Whitney U test). Therefore, all data were compiled together.

**Baseline data**

In girls, chronological age at the start of triptorelin treatment was 7.7 (0.8) years (table 1). The onset of puberty as reported by the parents was 6.0 (2.0) years. In boys, chronological age at the start of treatment was 7.9 (1.9) years.

The actual height at the start of treatment in girls is reported in table 1. It was 140.1 (8.4) cm (+2.10 (2.21) SD score) in boys. Patients were treated for a period of 3.4 (1.1) (F) and 4.2 (0.8) (M) years, respectively, and were followed up for 4.0 (1.2) (F) and 3.9 (0.9) (M) years after the discontinuation of treatment.

**Height velocity**

Before initiating treatment the height velocity was 8.0 (2.3) cm/year in girls and 10.4 (4.8) cm/year in boys. In girls, during treatment height velocity decreased to low values in the last year of treatment (table 2). An increase in height velocity...
Correlation between final height and other variables
In girls a positive correlation was shown between final height and height at the beginning and at the end of treatment ($r = +0.75$, $p<0.001$, and $r = +0.84$, $p<0.001$ respectively). A negative correlation was observed between final height and bone age at the end of treatment ($r = -0.52$, $p = 0.03$). Furthermore, a negative correlation between chronological age and bone age at the end of treatment and the height increment after treatment was found ($r = -0.45$, $p = 0.005$, and $r = -0.85, p<0.001$ (fig 2), respectively). No linear correlation was observed between final height and chronological age or bone age at start of treatment or the duration of treatment. A weak correlation was demonstrated between final height and target height ($r = +0.32$, $p = 0.05$).

Stepwise multiple regression analysis revealed that final height ($y$) was influenced most...
significantly by height at the start of treatment ($x_1$). Other significant factors were the bone age at the point of starting of treatment ($x_2$) and at discontinuation of treatment ($x_3$) and the duration of treatment ($x_4$) ($y = +0.91 \times x_1 - 3.6 \times x_2 - 5.0 \times x_3 + 1.5 \times x_4 + 132.5; R^2 = 0.89$; residual SD = 2.46; $p<0.001$).

Twenty four of the 31 girls showed a positive change in predicted adult height (median 5.2 cm, range 0.5 to 10.5 cm) (group A); however, seven of the 31 girls (group B) reached a final height below predicted adult height at start of treatment (median $-2.9$ cm, range $-9$ to $-4.8$ cm). The most important differences between these groups were a younger age at start of treatment in group A (7.5 (0.8) v 8.4 (0.6) years, $p = 0.01$), a higher bone age/chronological age ratio at start of treatment in group A (1.45 (0.15) v 1.29 (0.08), $p = 0.01$), and a longer period of treatment in group A (3.7 (1.1) years v 2.5 (0.7) years, $p = 0.02$).

In 26 girls target height was available. Sixteen of these girls (62%) achieved final height within the target height range, whereas 10 did not.

Change in body proportions and composition
The SD score of the ratio of sitting height to subischial leg length at final height was normal in girls, at 0.07 (0.98) ($n=26$). In 15 girls the ratio $SHLL/SDS_{CA}$ was known during treatment and at final height (table 1). No significant changes were observed.

BMI $SDS_{BA}$ in girls at start of treatment was higher than the reference population and it did not change significantly during and after treatment (table 1). BMI $SDS_{BA}$ at start of treatment was normal and increased during treatment (table 1). It did not change after discontinuation of treatment. The same pattern was observed in boys.

We compared the BMI SD score data of our patients with those from a group of 23 girls treated daily with subcutaneously buserelin, another GnRH agonist, for two years.1 In this buserelin group [chronological age 6.0 (1.7) years; bone age 9.9 (2.2) years at the start of treatment] BMI $SDS_{CA}$ and BMI $SDS_{BA}$ were calculated at the start and at the end of buserelin therapy (BMI at final height was not known in this group of girls). BMI $SDS_{CA}$ at the start and at the end of the treatment was 1.07 (1.00) and 1.30 (1.05) respectively, not significantly different from the equivalent data in our triptorelin group. BMI $SDS_{BA}$ at the start and at the end of buserelin was 0.28 (0.64) and 0.06 (0.75) respectively. BMI $SDS_{BA}$ at the start of buserelin treatment was not significantly different from BMI $SDS_{BA}$ at the start of triptorelin. On the other hand BMI $SDS_{BA}$ at the end of treatment was significantly higher in the triptorelin group than in the buserelin group ($p<0.001$). (There was no difference in bone age at the start of therapy between the two groups.)

Discussion
This study shows that adult height in girls can be increased by treatment with the slow release GnRH agonist triptorelin, although less than would be expected at the point of discontinuation of treatment.

Final height in our girls was 161 (7.0) cm, greater than the final height of untreated girls in the studies of Thamdrup1 (151.3 (8.8) cm, $n = 26$), Sigurjonsdottir and Hayles7 (152.7 (8.0) cm, $n = 34$), and Paul et al$^{11}$ (152.7 (8.6), $n = 93$). The positive change in predicted adult height of 3.5 cm lies between the values of 2.4 to 6.2 cm reported in two other studies where daily subcutaneous preparations were used.11,12

Using triptorelin a greater improvement in predicted adult height of 6.5 (1.4) cm was reported by Brauner et al, despite the fact that the inclusion criteria used were similar to ours.13 The most remarkable positive change in predicted adult height—10 cm—was described by Paul et al in a group of girls treated for 5 years of age.14 This tendency for young patients to obtain a greater positive change in predicted adult height was also observed in our population: in the girls who had a positive change in
predicted adult height of 5.2 cm (range 0.5 to 10.5 cm) (group A), chronological age was significantly lower than in those girls who had a negative change in predicted adult height. Therefore age at the start of treatment appears to play an important role. Only four girls in our group were younger than 7 years of age at the start of treatment; hence we could not show a significant relation between age at start of treatment and the final height obtained.

Height at start of treatment proved to be the most important positive factor influencing final height, as demonstrated by multiple regression analysis and by a positive correlation coefficient (r = +0.75), a characteristic that is also described in normal puberty. Final height showed a weak association with target height, as also shown by Paul et al. Only 62% of the girls with a known target height reached a final height within the target height range. However, this weak association between target height and final height was similarly identified in the normal population (0.45).

In contrast to other studies, no improvement of adult height was demonstrated in boys. The most plausible explanation for this difference is that three out of the five boys were suffering from neurofibromatosis, a condition which itself is associated with growth impairment, and in which the development of complications such as optic nerve glioma may lead to growth hormone insufficiency. On the other hand, in some cases growth impairment has been described independently of growth hormone levels. In our patients growth hormone secretion was not evaluated.

The difference in final height as predicted at the point of discontinuation of treatment and the actual final height attained is caused by a moderate growth acceleration after cessation of treatment and a definite accelerated bone maturation within the same period. The same tendency has been described by Oerter et al. As suggested before, the post-therapy level of gonadotrophins and sex steroids rises more rapidly than in normal puberty, and higher levels are reached in a shorter period of time. This results in a shorter period of growth and eventually a final height that is less than the predicted adult height at the point of discontinuation of treatment.

In addition to height at start of treatment, bone age at the point of discontinuation of treatment is another important negative factor contributing to final height. As shown in fig 2, residual growth capacity is greater before a bone age of 12.0 to 12.5 years. For this reason discontinuation of GnRH treatment might be considered at a bone age of 12 to 12.5 years.

Triptorelin treatment did not influence body proportions, as normal ratios of sitting height to subischial leg length were observed at the point when final height was reached. In girls, BMI SDS, was increased at start of treatment and did not change during or after discontinuation of treatment. On comparison of our data with those of Marti-Henneberg, who described the normal course of BMI during early, normal, and late puberty, the increased BMI SDS at start of treatment appears to be a natural event. Since children with CPP who had been exposed to sex steroids underwent a pubertal growth spurt and subsequently developed accelerated bone age, it would be better to use the index BMI SDS. Using this index, a comparison can be made between the girls with CPP and those with normal age of onset of puberty at similar bone ages. The index was normal at start of treatment and increased during treatment, suggesting that triptorelin increases BMI, as no change in BMI SDS was observed during buserelin treatment.

In our patients a rapid resumption of puberty was observed, menarche occurring 1.1 years (range 0.4 to 2.6 years) after withdrawal of treatment, at a mean age of 12.3 (1.1) years. This is comparable with the results of other investigators. Ultrasound evaluation showed normal ovaries 4.0 (1.2) years after discontinuation of treatment. In contrast to the report by Adams et al, no evidence of polycystic ovaries was found.

We conclude, therefore, that treatment with the depot GnRH agonist triptorelin increases final height in girls without giving rise to abnormal body proportions. The best results are obtained in girls with a greater height at the point of starting treatment. After withdrawal of therapy, resumption of puberty occurs rather rapidly, without evidence for development of polycystic ovaries. Triptorelin seems to increase BMI. From an auxological point of view, discontinuation of triptorelin in girls might be considered at a bone age of 12.0 to 12.5 years.

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