Hypothesis

The Y specific growth gene(s): how does it promote stature?

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Abstract

Although the presence of a Y specific growth gene(s) (Y growth gene(s)) on Yq has widely been accepted, it remains unknown how this gene promotes stature. In this report, we discuss the growth pattern in normal boys and girls and in patients with growth disorders informative for the Y growth gene(s). The results suggest that the Y growth gene(s) augments statural growth by controlling the sex steroid independent childhood growth pattern.

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Keywords: Y specific growth gene(s); childhood growth.

We have proposed the presence of a Y specific growth gene(s) (Y growth gene(s)) that augments adult height by 8-10 cm independently of the effect of gonadal sex steroids.1,2 This hypothesis is primarily based on white adult height comparisons between patients with pure XY gonadal dysgenesis and those with pure XX gonadal dysgenesis (172.0 cm (SD 7.0)), n=24 v 164.3 cm (SD 7.7), n=22, p<0.01), between patients with complete testicular feminisation syndrome and normal females (172.2 cm (SD 6.5)), n=23 v 162.2 cm (SD 6.0), the British standard, p<0.01), and between normal males and XX males (174.7 cm (SD 6.7)), the British standard, v 166.4 cm (SD 7.4), n=33, p<0.01), in whom the sex chromosome complement is different but the effect of bioactive gonadal sex steroids is similar.1,2 Furthermore, we have localised the Y growth gene(s) to the proximal Yq region between DYS11 and DYS246, by genotype-phenotype correlations in non-mosaic adult male patients with a partial Yq deletion.3 Here, we discuss how the Y growth gene(s) promotes statural growth.

The mean height (supine length during infancy and standing height thereafter) is in principle greater in boys than in girls, except for around the ages of 12 to 14 years when girls become taller than boys because of the earlier onset of the pubertal growth spurt.4 It is noteworthy that the statural difference between the sexes remains only 0.5-2.0 cm in childhood but increases to about 13 cm in adulthood.4 This may raise the possibility that the Y growth gene(s) augments stature in conjunction with gonadal sex steroids.

However, the sex difference in the height gain from childhood to adulthood is not inconsistent with the Y growth gene(s) acting in a sex steroid independent manner. The human growth pattern is divided into three additive and partially superimposed components (infancy, childhood, and puberty) on the basis of mathematical modelling of auxological data.5 Although the infantile growth component tails off by 3–4 years of age, the childhood growth component continues to mature height. Thus, the height gain from childhood to adulthood is the sum of the height increase caused by the extension of the sex steroid independent childhood growth and that added by the sex steroid dependent pubertal growth. According to the Swedish longitudinal growth study, mean childhood growth is expressed by the following second degree polynomial function: boys, Y=-0.18X²+8.53X+71.9 and girls, Y=-0.22X²+8.93X+69.9 (Y: height in cm; X: age in years).6 This sex difference in the childhood growth function produces only 2 cm of height difference at 10 years of age but yields 10 cm of height difference at 20 years of age. Thus, it is assumed that the Y growth gene(s) controls the sex steroid independent childhood growth pattern, leading to the adult height difference of 8-10 cm. The remaining adult height difference between the sexes is explained by the notion that testicular sex steroids have the potential to increase the adult height by 3-5 cm whereas ovarian sex steroids have no major effect on the adult height.1,2

If the above assumption is correct, it is expected that the Y growth gene(s) leads to sex difference in height, which remains hidden in childhood but becomes obvious in adulthood, in disorders lacking a pubertal growth spurt. In this regard, growth data are available in achondroplasia in which sex steroids fail to cause an apparent growth spurt because of severely compromised bone capacity of linear growth. In this disease, the sex difference in the mean height remains only 0.5-1.0 cm in childhood, but becomes gradually clearer during puberty, amounting to about 7 cm in adulthood.6 This growth pattern is consistent...
with the notion that the Y growth gene(s) promotes adult height by controlling childhood growth. Similar growth patterns have also been reported in Laron syndrome and in Down syndrome, although the growth data are insufficient to draw a definite conclusion.

It is also expected that a mutation of the Y growth gene(s) results in a male to female alteration of the childhood growth pattern and, therefore, causes growth deficiency that is hardly recognised in childhood but becomes discernible in adulthood. We have experienced eight 46,XY male patients who exhibit such a unique growth pattern (fig 1). These patients may represent clinical models for a mutation of the Y growth gene(s), although the possibility remains purely speculative at present.

One may argue against the above hypothesis, because several male patients with Yq deletions have short stature which is already obvious in childhood. Such short stature is not explicable by assuming loss of the Y growth gene(s) that is expected to cause a male to female alteration of the childhood growth pattern. However, the Yq deletions are associated with chromosome imbalance (quantitative alteration of euchromatic or non-inactivated region), and it has been suggested that chromosome imbalance could disturb developmental homeostasis or impair cell proliferation, resulting in global non-specific developmental defects including growth failure. In fact, karyotype-phenotype correlations in sex chromosome aberrations are consistent with chromosome imbalance being involved in the development of several features including growth failure. Furthermore, it is possible that the Yq deletions are caused by unbalanced translocations involving the X chromosome or the autosomes, rather than by simple Yq deletions. Indeed, cryptic Xq;Yq translocations have been reported in several retarded patients with Yq- chromosomes. Such unbalanced translocations would result in gross chromosome imbalance and perturbation of multiple non-Y linked genes, exerting a deleterious effect on statural growth. Thus, short stature in boys with Yq deletions would not provide compelling evidence against the hypothesis that the Y growth gene(s) is responsible for the sex difference in the childhood growth pattern.

In summary, we propose that the Y growth gene(s) promotes statural growth by controlling the childhood growth pattern. Furthermore, the Y growth gene(s) might also be involved in fetal growth, since sex difference in fetal body size in eutherian mammals is partly ascribed to the growth advantage brought about by a Y linked gene(s) other than Sry. The hypothesis will be tested when the Y growth gene(s) has been cloned.

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