the Pitt-Rogers-Danks syndrome may be caused by a deletion in the same region.

In a recent issue of the Journal of Medical Genetics, Partington et al. reported on a number of patients with deletions or duplications of 4p16.3 in three families: deletion causing Pitt-Rogers-Danks syndrome and duplication resulting in a new overgrowth syndrome, we have serious concerns about (1) their definition of a new overgrowth syndrome and (2) their idea that a triple dose of FGFR3 results in physical overgrowth.

First, we have reAnalysis the data from their table 2 to show that overgrowth is not really a prominent manifestation of duplication involving FGFR3 (table 1). For both height and head circumference, fewer than half of the patients have values ≥90th centile; only with respect to weight do slightly more than half of the patients have values ≥90th centile. On the other hand, a good proportion of patients have values ≤50th centile and even ≤25th. Combining all three parameters (n=76), at the extremes, 28% of patients have values ≥97th centile and 8% have values ≤3rd centile. Although some values for all three growth parameters are large and in the range anticipated to be in that direction, so many patients have middle and lower range values that overgrowth per se does not seem to us to be a particularly prominent manifestation of dup(4p16.3). In fact, when thinking of classical overgrowth syndromes, such as Beckwith-Wiedemann syndrome, Simpson-Golabi-Behmel syndrome, or Robinow-Segovia-Ruvalcaba syndrome with overgrowth frequently present at birth,1 it is difficult to think of dup(4p16.3) as an overgrowth syndrome at all.

Secondly, the function of FGFR3 can be deduced from the Fgrf3−/− knockout mouse,1 which is overgrown with excessively long femora and elongated vertebrae, resulting in a long tail. Thus, the normal function of FGFR3 is to regulate endochondral ossification by "putting the brakes on" growth.

Evidence is accumulating that the known mutations on FGFRs are of the gain of function type. For example, Neilson and Friesel1 made mutations in mRNA and expressed them in Xeno. tyrosine kinase activity compared to their wild type counterparts. FGFR3 mutations for achondroplasia and thanatophoric dysplasia have also been shown to have greatly increased levels of phosphorylated tyrosine kinase activity. Ligand-independent constitutive signalling produced by these mutations results in premature maturation of bones of the skeleton and cranium. This type of activation depends on the particular mutation and its location on the receptor and appears to result from (1) aberrant disulphide bond or hydrogen bonded GFGR dimers (or 2) involvement in the activation loop of the kinase domain.10 Thus, the mutations for short limb skeletal dysplasias on FGFR3 (hypochondroplasia, achondroplasia, and thanatophoric dysplasia) are gain of function mutations that "put the brakes on even more" to various degrees. Fgrf3−/− heterozygous mice have been shown to be normal.11

In conclusion, the idea put forth by Partington et al. that FGFR3 in single dose leads to growth failure and in triple dose to physical overgrowth is not tenable in view of current clinical and experimental evidence.

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Table 1 Rearranged data

<table>
<thead>
<tr>
<th>Centiles (%)</th>
<th>No &gt; 90</th>
<th>≤50</th>
<th>≤25</th>
</tr>
</thead>
<tbody>
<tr>
<td>Height</td>
<td>31</td>
<td>32</td>
<td>29</td>
</tr>
<tr>
<td>Weight</td>
<td>23</td>
<td>52</td>
<td>35</td>
</tr>
<tr>
<td>Head circumference</td>
<td>22</td>
<td>41</td>
<td>32</td>
</tr>
</tbody>
</table>

References

13 Webster MK, Donoghue DJ. Constitutive activation of fibroblast growth factor receptor 3 by the transmembrane deletion mutation found in achondroplasia. EMBO J 1996;15: 520-7.

This letter was shown to Dr Partington et al., who reply as follows.

Thank you for the opportunity of seeing the comments of Cohen and Neri on our recent paper. We were mildly surprised at their confidence not only in dismissing the possibility of a dosage effect of FGFR3 on growth, but also in asserting that all the short stature syndromes associated with mutations in this gene are explained by loss of gain of function. We believe the jury is still out and all will be revealed in good time.

However, we must protest at Cohen and Neri's attempt to extinguish our overgrowth syndrome by the rather cavalier manipulation of the growth data presented in table 2 of our paper. They have chosen to ignore two important points stated in our report: first that II.12 in family 1 is a special case because of severe disease (everyena), which probably limited growth in childhood and, second, that overgrowth "became more obvious in late adolescence and early adulthood".

What Cohen and Neri have done is to take their mixed longitudinal and cross sectional growth data, translate it all into cross sectional data and, neglecting age, treat these measurements in the same way. This does dilute the overgrowth patterns observed. However, if one omits the data on II.12 for the reasons stated and takes the measurements at the oldest age of the 10 remaining subjects (all over the age of 15 years), then all the heights are at or above the 75th centile and five of them are above the 90th centile. In the same way, the head circumference are all at or above the 50th centile with five above the 97th centile. This tells quite a different story from table 1 presented in Cohen and Neri's letter.

Some confirmation of this late growth pattern emerges from the heights recorded on II.11 and II.12 from family 1. Thus, between the ages of 18 and 28 years 5 months, II.11 grew 15 cm and between 17 years 5 months and 28 years II.12 grew 26 cm. Such growth increments are commonplace between the ages of 12 and 16 years, but are quite abnormally after the age of 17 years.

Physical measurements do not convey the whole picture. Clinically, those with overgrowth appear to be big people with large body frames, prominent supraorbital ridges, heavy facial features, and big hands and feet. Some, but not all, of these features are shown in the illustrations in our article.

Lastly, your readers may be interested to know that since our paper was published we have managed to meet III.8 in family 1 again and, on this occasion, were able to make some measurements and take a blood sample. On the age of 33, III.8 is a big woman with a heavy body frame and rather coarse or heavy facial features of which she is self-conscious and does not allow herself to be photographed.

Her height is 180 cm, head circumference 60 cm, and hand length is 20.5 cm. She has mild intellectual handicap and a duplication of 4p16.3.

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Robinow syndrome

I would like to comment on the paper by Sabry et al. in the September issue of the Journal of Medical Genetics. I have seen three patients they diagnosed as having Robinow syndrome in conjunction with a number of unusual abnormalities.

When looking at the photographs of their patients, I doubt whether the diagnosis of Robinow syndrome is correct, especially in the first patient. She has a number of facial characteristics that are indeed seen in Robinow syndrome, such as hypertelorism and...

Letters to the Editor, Book reviews, Notice

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