Interestingly, the previously reported patient with a different but overlapping deletion of chromosome 12 long arm (q13.3–q21.1) and the subject of this report have only a few features in common (table). The deletion in each includes the portion 12q13.3 to q21.1. It is unclear whether the small difference in the portion of 12q deleted accounts for the fact that they do not resemble each other to a greater extent. The delineation of a syndrome associated with interstitial deletion of 12q must await reports of additional cases.

The detection of subtle chromosome abnormalities such as this show the need for minimum standards for routine studies of those at risk for chromosome abnormalities. Our impression is that these minimum standards should include well banded chromosomes of the 450 to 550 band stage.

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A third case of de novo partial trisomy 4p

More than 30 cases of partial trisomy 4p have been described.1 The majority of these cases were the result of a parental translocation and only two cases had parents with normal karyotypes.

The proband was born spontaneously at term after a pregnancy complicated by mild toxaemia. Paternal age was 32 years and maternal age 29 years. Birth weight was less than 2500 g. Failure to thrive was mentioned in the early medical records, but generally physical health has been good. His height developed along the 25th centile. At 16 years of age he started to grow excessively with a final height, at the age of 21, of 201 cm. Psychomotor development has always been slow and speech never developed.

At the age of 23 years the proband was examined by one of us (JWEO). He was a tall, restless, mentally deficient man with an occipitofrontal circumference of 53 cm (−2 SD). His face (fig 1) was elongated and rather narrow with a prominent glabella, heavy eyebrows with synophrys, a mild mongoloid slant, midfacial hypoplasia, bulbous nose, mild macrostomia, high palate, and a long, receding chin. The ears showed a slightly prominent antihelix and attached earlobes. There were cubiti valgi, bilateral short third to fifth metacarpals, short first metatarsals, and a short fourth metatarsal on the left. Secondary sexual characteristics were normal, although the testes were rather small in size.

Examination by an endocrinologist provided no additional information; ophthalmological examination was normal. The EEG showed diffuse abnormalities, especially of the brain stem. Radiographs showed a normal skull, shortened metacarpals, and hypoplastic first ribs. The degree of mental deficiency was considered to be moderate, but was difficult to assess because of behavioural problems.

Cytogenetic analysis was performed on peripheral blood lymphocytes, using GTG and RBA banding techniques. Fourteen cells from the proband were examined and showed 46 chromosomes in all cells with a partial duplication of chromosome 4: 46,XY,dup(4)(p12→p15.2) (fig 2). The parental karyotypes were normal.

Apart from the excessive height the phenotypic abnormalities of the proband are consistent with trisomy 4p, as reviewed by Gonzalez et al.1 Only two cases of de novo interstitial trisomy 4p have been reported. In a third case,2 a chromosome 1 was also involved and therefore, in the strictest sense, this was not isolated trisomy 4p.

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FIG 2  (a) Partial karyotype of the proband. (Above) GTG banding. (Below) RBA banding. The arrows indicate the duplicated segment. (b) Idiogram of chromosome 4.

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