Giemsa Banding Pattern of a Heritable 1q+ Variant Chromosome: A Possible Partial Duplication

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One morphological variant of chromosome 1 in man differs from the more usual form of this chromosome in being longer and more submetacentric. This is due to increased length of the long arm and because there is apposition of chromatids in the paracentromeric region of the long arm (q). This variant, or similar appearing variants of chromosome 1, have been found in 1 per 100 to 1 per 1000 newborns in the United States (Lubs and Ruddle, 1971).

Previous studies have reported 1q+ variants in individuals with a variety of anomalies as well as in phenotypically normal individuals (Cooper and Lawler, 1963; Donahue et al, 1968). Since the same anomaly has never been associated with the variant chromosome in more than one instance, and since phenotypically normal individuals have also been reported carrying the variant chromosome, it can be concluded that it is a normal variant which causes no obvious phenotypic abnormality.

Family studies of 1q+ variants have, together with gene linkage studies, served to assign a series of genes to chromosome 1. These include the loci for the Duffy blood group (Donahue et al, 1968; Ying and Ives, 1968), congenital pulverulant cataract (Renwick and Lawler, 1963/1964), and amylase 2 (Karamyt, Adamek, and Vrba, 1971; Hill, Rowe, and Lovrien, 1972).

We investigated a 3-generation Caucasian family with a variant chromosome 1 (Fig. 1). The proposita was originally ascertained because of mild mental retardation but had no physical anomalies. Other members of the family were phenotypically normal. We designed studies (1) to determine the effect of the hypotonic treatment upon the appearance of the variant; and (2) to learn if the apparent increase in length was due to uncoiling or to the addition of chromosomal material to 1q.

Materials and Methods

Peripheral blood was obtained from each family member. Lymphocytes were cultured and harvested after 72 hr (Moorhead et al, 1960). The fixative was methanol: acetic acid (3:1) and slides were made by flame-drying.

To test the effect of differing hypotonic treatments, the following 3 regimens were tried on different aliquots of the cell suspension: a—0.075 M KCl for 10 min at room temperature; b—0.93% sodium citrate for 10 min at room temperature; and c—distilled water for 30 min at 37°C.

Metaphases from each family member were scored visually for the presence of the 1q+ variant. Criteria for determining that a metaphase manifested the 1q+ variant chromosome were that: (1) it contained a long, submetacentric chromosome 1 due to elongation of 1q; and (2) the suspected variant chromosome showed close apposition of chromatids in the paracentromeric region of the long arm.

The acetic/saline/Giemsa (ASG) procedure was employed to reveal chromosome banding (Sumner, Evans, and Buckland, 1971). Lymphocytes were prepared by regimens a and fixed in methanol: acetic acid (3:1) for 15 minutes. Slides were made by flame drying, then incubated in 2×SSC (0.3 M NaCl and 0.03 M trisodium citrate) at 60°C for 1 hr, rinsed once in distilled water, and stained with a Giemsa solution (Harleco brand, 1 ml in 100 ml, pH 6.5 phosphate buffer, for 12 min or with Gurr's Giemsa 'R66', 1 ml in 50 ml, pH 6.8 phosphate buffer, for 90 minutes).

Results

Family Study. Of the 9 family members, 4 were found to be heterozygous for the 1q+ variant (Figs. 1 and 2). Karyotypes from the 9

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individuals otherwise appeared unremarkable, providing no hint as to the presence of a subtle reciprocal or insertional translocation. Caliper measurements were also normal in all the karyotypes from the 9 family members, except in the case of the 1q+ variant present in the 4 heterozygotes.

Effect of Hypotonic Treatment. A representative metaphase spread is illustrated in Fig. 2. Treatment c (distilled water for 30 minutes) resulted in an increased proportion of cells with an incomplete complement of chromosomes, probably due to increased breakage of the cell membrane. The variation in length and the increased submetacentricity of the 1q+ variant were apparent in essentially all metaphases from each of the 4 heterozygotes, independent of the hypotonic treatment.

Apposition of chromatids was influenced by the type of hypotonic treatment. Between 50 and 100 metaphases from each regimen were scored for this criterion. Apposition in the paracentromeric region of 1q was observed in 25–35% of 1q+ metaphases after KCl treatment (regimen a), 35–45% after sodium citrate (regimen b), and in 50–60% of cells after aqueous hypotonic treatment (regimen c).

The overall chromosome morphology was best and the ASG-banding patterns most distinct after

![Figure 1](image1.png)

**Fig. 1.** Pedigree of the family in which the variant chromosome 1 segregates.

![Figure 2](image2.png)

**Fig. 2.** A representative metaphase from an individual who carries the variant chromosome 1 (arrow) stained with Giemsa without the pretreatment to produce differential banding.
KCl hypotonic treatment (regimen a). This regimen was therefore used to prepare chromosomes for ASG-banding studies.

**ASG-Banding Studies.** The ASG patterns of the usual chromosome 1 and the 1q+ variant chromosome were compared. The banding patterns distal to the prominent secondary constriction on 1q appeared identical. However, differences were observed in the region between the centromere and the secondary constriction. In that region the usual chromosome 1 had a single dark band, whereas the 1q+ chromosome consistently had 2 dark bands with an intervening lightly stained band (Fig. 3). The 2 dark bands on 1q+ were similar in size and stained with the same intensity as the single dark band in this region on the usual chromosome 1 (Fig. 4a). In some cells when 2 bands could be resolved in the region between the centromere and the secondary constriction on the long arm of the usual chromosome No. 1, the same region of the 1q+ variant showed a corresponding increase in the number of subunits.

**Discussion**

Several theories have been advanced to explain the morphological appearance of the variant 1q+. It has been suggested that the variation in length might be due to uncoiling of the normal structure (Donahue *et al.*, 1968; Kamarýt *et al.*, 1971) or to increased length of the normal secondary constriction on chromosome 1 (Ferguson-Smith *et al.*, 1962). Another explanation is that 1q+ represents the addition of chromatin material to the long arm of chromosome 1 (Patau *et al.*, 1961; Cooper and Hernits, 1963; Ying and Ives, 1968).

Our demonstration of an extra band in the variant region of equal size and intensity as the band normally found in this region of chromosome 1 is most consistent with the hypothesis that the increase in length in the variant 1q+ is the consequence of additional material. This hypothesis is further supported by the recent demonstration of a polymorphism in the amount of DNA found in this region (Jones and Corneo, 1971).

Addition of chromosome material could occur by one of 2 mechanisms, either translocation or partial duplication. A translocation could be of 2 types,
insertional or reciprocal. A reciprocal translocation would presumably result in a different banding pattern on 1q+ distal to the break point, which was not observed. We have shown that the extra material occurs in an internal segment of the long arm of the chromosome 1 between the centromere and the often prominent secondary constriction. An insertional translocation is not likely because it would require a minimum of 3 simultaneous chromosome breaks.

The most probable source of extra chromatid material in the variant chromosome 1 is partial duplication (Cooper and Hernits, 1963; Ying and Ives, 1968). One mechanism by which this could have occurred is unequal crossing-over in meiosis. There is evidence for the occurrence of this process on the gene level in humans, for example, in the haemoglobin system (Nance, 1963) and with haptoglobin (Smithies, Connell, and Dixon, 1962). Further evidence for this mechanism would be provided by the observation of the presumed reciprocal product, consisting of a chromosome 1 with an abbreviated long arm and the loss of the normal single dark band.

Although unequal crossing-over should occur randomly, the existence of the partially duplicated product without any accompanying phenotypic abnormalities would be more likely to occur in an area of reiterated DNA sequences. In this connection, it is interesting that with a technique for visualizing centromeric-associated heterochromatin (Arrighi and Hsu, 1971) the variant region on chromosome 1 stains heavily and hence has been postulated to contain highly redundant DNA sequences (Craig-Holmes and Shaw, 1971; Lubs and Ruddle, 1971). Moreover, RNA complementary to satellite II DNA has been found to anneal most prominently to the paracentromeric region of chromosome 1, suggesting that this region is particularly rich in reiterated sequences (Jones and Corneo, 1971). Variation in this region would therefore be consistent with the absence of phenotypic abnormalities in the families in which the variant 1q+ segregates.

Caution should be taken not to extend the results reported here without appropriate studies to other 1q+ variants segregating in unrelated families.

Summary

A 1q+ chromosome variant was found to segregate through 3 generations of a family. The 1q+ variant showed, aside from increased length of the long arm (q1), close apposition of chromatids in the paracentromeric region of q. The frequency with which chromatid apposition was observed depended upon the type of hypotonic treatment. Acetic/saline/Giemsag banding studies revealed extra bands in the paracentromeric region of 1q, consistent with the addition of chromosome material. These findings are most easily explained by partial chromosome duplication due to uneven crossing-over.

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