Not para-, not peri-, but centric inversion of chromosome 12

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Abstract
A 39 year old male with primary infertility was diagnosed as having Klinefelter syndrome by conventional cytogenetic analysis, which also showed an abnormal chromosome 12. Fluorescence in situ hybridisation (FISH) analysis of the aberrant chromosome using a 12 specific centromeric probe showed a break in the alphoid repeats followed by an inversion within the short arm, resulting in a pseudodicentric chromosome. Further FISH analyses using telomeric and subtelomeric probes showed that the other breakpoint was in the subtelomic region of the short arm. The karyotype is designated 47,XXY,inv(12)(p10p13.3). To our knowledge this is the first report of a case of "centric inversion".

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Following the discovery of the specificity of centromeric alpha satellite sequences for each chromosome,1 fluorescence in situ hybridisation (FISH) using these sequences became an important tool in resolving chromosome aberrations where conventional banding techniques were inadequate. FISH applications include identifying the chromosome origin of marker chromosomes and determining the localisation or involvement of the centromere and telomere in inversions and translocations.2-4 Using FISH, we have identified the breakpoints of a unique inversion involving the short arm of chromosome 12 in a patient with Klinefelter syndrome.

Case report
A 39 year old male with infertility was referred to our centre suspected of having Klinefelter syndrome. His testes were hypoplastic and he had azoospermia. The parents of the proband had died, but were non-consanguineous and phenotypically normal. There was no history of infertility in the family. The proband’s two brothers and two sisters, who had two or three children each, would not cooperate in the investigation.

Materials and methods
Metaphase chromosomes prepared from PHA stimulated lymphocytes and cultured skin fibroblasts were analysed by standard Giemsa staining and GTL and CBG banding. The commercial FISH probes (Oncor Inc) used were biotin or digoxigenin labelled D12Z3 (chromosome 12 centromere specific), biotin labelled “All Human Centromere”, and digoxigenin labelled “All Telomere” probes. In addition, 100 ng digoxigenin labelled DOP-PCR product of the YACs y922c8 (1390 kb), y968f7 (1580 kb), and y771h4 (1100 kb) mapping to the subtelomeric region of the short arm of chromosome 12 were used. Labelling and hybridisation were carried out according to Kievit et al.5

Results
Giemsa staining and GTL banding of the initial PHA stimulated lymphocyte chromosomes of the patient suspected of having Klinefelter syndrome showed 47 chromosomes, including an extra X chromosome and a chromosome 12 with an aberrant short arm, in all the cells analysed (fig 1). FISH analysis of chromosome 12 using D12Z3 showed two signals, one located at the original centromere and the other in the telomeric region (fig 2). The intensity of these two signals were similar to one another, though weaker than the centromeric signal on the normal chromosome 12. These findings indicated a centromeric fission followed by inversion of the whole short arm of chromosome 12.

Chromosomes were further analysed for the primary constriction and to determine whether the inversion involved the whole short arm or a part of it. With CBG banding, the aberrant chromosome was shown to contain two heterochromatin regions, one located in the original centromeric region and the other in the telomeric region. The primary constriction was always at the telomeric site in 75 lymphocytes and 25 skin fibroblasts analysed. Simultaneous hybridization with biotin labelled “All Centromere” and digoxigenin labelled “All Telomere” probes showed that the telomeric sequences were not inverted. Further analysis of the telomeric region was carried out using the YAC probes y922c6, y968f7, and y771h4, mapping to a region 2 cM, 5 cM, and 15 cM from the telomere, respectively (fig 2). The FISH signals for y922c8 and y968f7 were stationary, while the signal for y771h4 had moved to the original centromeric region of the abnormal chromosome 12. This result indicated that the distal breakpoint has occurred between the YACs y968f7 and y771h4, 5-15 cM from the telomere.

The karyotype of the patient is designated 47,XXY,inv(12)(p10p13.3) and the extended karyotype is 47,XXY,inv(12)(p10→p13.3::p10→p13.3::q10→qter).
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Discussion
In a patient with Klinefelter syndrome we have identified an inversion of the short arm of chromosome 12, where one of the breakpoints was within the centromeric alpha satellite sequences and the other in the subtelomeric region, 5-15 cm from the telomere (Fig 2). Chromosome 12 is one of the chromosomes in which inversions have been reported frequently, including 14 pericentric and 19 paracentric inversions.\textsuperscript{7-12} However, an inversion with a centromeric breakpoint has not been reported previously, either for chromosome 12 or for any other human chromosome. We therefore suggest the term "centric inversion" for this unique case.

Since there is great individual variability in the number of alpha satellite repeats of a specific centromere, the absolute amount of alpha satellite DNA cannot be critical for the function of the centromere.\textsuperscript{11} This is supported by the detection of partial deletion of the alpha satellite sequences without any effect on mitotic stability or meiotic segregation\textsuperscript{14,15} and studies suggest that as little as 140 kb of alpha satellite DNA may be sufficient for centromere function.\textsuperscript{14} Furthermore, alpha satellite repeat sequences could be split into two without functional loss of centromere activity.\textsuperscript{15,17,18} In the present case, although it is not possible to deduce the exact amount of the alphoid sequences, the intensity of the FISH signals was almost equal. Either of the centromeres could thus be expected to have sufficient alphoid sequences for normal centromere function. CBG banding results showed that the primary constriction was always in the telomeric region, suggesting that only one of the centromeres was active. However, FISH results contradicted those of CBG banding, as in some

Figure 1  Karyotype of the patient with 47,XXY,inv(12)(p10p13.3) showing the inverted chromosome 12 and the extra X chromosome.

Figure 2  Schematic representation of the short arm of chromosome 12 and the FISH results. The breakpoints are represented by vertical arrows. The proximal breakpoint of the inversion is within the alpha repetitive centromeric region, as shown by the presence of two signals on the inverted chromosome 12 using D12Z3 as FISH probe (C). The distal breakpoint of the inversion is localized to the subtelomeric region using YAC 9687 (A) and YAC 771h4 (B) as FISH probes. The hybridisation signal of YAC 9687 (A) remains in the telomeric region on the inverted chromosome 12, while the signal of YAC 771h4 has moved to the centromeric region (B), indicating that the breakpoint has occurred in a region 5-15 cm away from the telomere.
metaphases the condensed fluorescent signal indicative of the primary constriction was observed in the centromeric region. To define the activity of the centromeres, studies with antibodies specific to CENP-C and CENP-E are required.

Aneuploidy events, including Klinefelter syndrome, have been reported in conjunction with para- and pericentric inversions, implying a possible interchromosomal effect increasing the risk of non-disjunction. In one patient, Klinefelter syndrome and paracentric inversion of chromosome 12 occurred simultaneously, as in the present case, though involving the long arm. Various degrees of other infertility problems have also been reported in several male inversion carriers, suggesting an effect of inversions on infertility. In the present case, though, the sterility is likely to be the result of Klinefelter syndrome.

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