SRVX, a sex reversing locus in Xp21.2 → p22.11

Cyrogeneric duplication of the X chromosome in males is a rare event usually characterised by a significant degree of phenotypic abnormality, which can include sex reversal in the presence of an apparently normal Y chromosome. In this paper the authors report two half sibs with maternally inherited cyrogeneric duplications of Xp and sex reversal; the absence of dysmorphic features in mother and children is thought to be because of the relatively small extent of the duplication. Comparison with previous reports allows the putative sex reversing locus (SRVX) to be assigned to a 5–10 megabase fragment between Xp21.2 and Xp22.11 which includes the DMD locus. This regional assignment should help in the isolation of the SRVX gene mutations which may be a cause of sex reversal found in the 90% of sex reversed women with XY gonadal dysgenesis who do not have detectable mutations of the sex determining SRY gene, the sex reversal associated with the dominant condition campomelic dysplasia already mapped to 17q34, or the terminal deletions of 9p or 10q which probably contain recessive sex reversing loci.

JOHN C K BARBER

A worldwide study of the Huntington's disease mutation

One of the most significant consequences of the discovery of the Huntington's disease defect has been an improvement in preclinical and differential diagnosis for this condition. In contrast to the limitations of earlier genetic linkage analysis, assessment of the CAG expansion appears to be a much more accurate and specific diagnostic test. Kremer et al. in the most extensive study to date of this aspect, confirm earlier reports of the high sensitivity and specificity of measuring CAG repeats. Within 565 families from 43 national and ethnic groups were 1007 patients with signs and symptoms compatible with a diagnosis of Huntington's disease. Of these, 995 had an expanded CAG repeat that included from 36 to 121 repeats (sensitivity 98.8%, 95% CI 97.7–99.4). Included among those contributing to the sensitivity estimate were 12 patients with previously diagnosed HD in whom the number of CSG repeats was in the normal range. Re-evaluation of these established that 11 had clinical features atypical of Huntington's disease. In 1581 of 1595 control chromosomes (99.1%), the number of CAG repeats ranged from 10 to 29. The remaining 14 control chromosomes had 30 or more repeats, with two of these chromosomes having expansions of 37 and 39 repeats. An estimate of specificity data was made from 113 subjects with other neuropsychiatric disorders with which Huntington's disease is frequently confused. The number of repeats found in these disorders was similar to the number found on normal human chromosomes and showed no overlap with Huntington's disease (specificity 100%, 95% CI 95.5–100). This study also confirms that the CAG expansion is the molecular basis of Huntington's disease worldwide.

DAVID RAVINE

Huntington disease without CAG expansion: phenocopies or errors in assignment?

Huntington's disease (HD) is associated with an expanded triplet (CAG) repeat within a gene on 4p16.3. Although the paper describing the discovery of the gene was only published in March 1993, several thousand DNA samples have been tested already in various laboratories around the world. In the majority of cases which were diagnosed clinically as having HD, molecular analysis has confirmed the diagnosis. In a few cases, however, there has been no expansion and there can be a variety of reasons for this. Andrew et al report their experience in 1022 clinically affected patients. They found 30 (2.9% of the cohort) who did not have an expanded CAG repeat in the disease range. Ten of the 30 persons with normal sized alleles represented clinical misdiagnosis, six involved a sample mix up, and two arose as clerical errors. The other 12 represent possible phenocopies for HD. It will be interesting to see how many of the patients in the misdiagnosis group (and possibly also the phenocopy group) turn out to have dentato-rubropalidolysian atrophy, the latest triplet repeat to be discovered. This report underlines the importance of confirming the diagnosis in patients at a molecular level whenever possible, before offering presumptive testing to relatives. The small potential for human error can never be completely excluded.

FRANCES FLINTER

Neurofibromatosis type 1: the cognitive phenotype

Learning difficulties are estimated to occur in 30 to 40% of children with NF1. In this study a group from Johns Hopkins University set out to explore whether the presence of the NF1 gene results in a global cognitive deficit as measured by a lowering of IQ, or in a more specific cognitive deficit or learning disability. In addition, they sought to establish whether learning disabilities could be correlated with brain MRI scan findings. Families were included about the study via NF centres and organisations. Of those expressing interest, 12 families with the appropriate structure were chosen. Each comprised of one child with NF1, an unaffected sib, and both natural parents. NF children with known intracranial problems were excluded but family members with known learning difficulties or hyperactivity disorders were not, making some of the results difficult to interpret. A history, physical examination, MRI scan, and a battery of psychological tests were carried out on each person. The results were subject to various statistical analyses. Full scale IQs ranged from 70 to 130 among children with NF1 and from 99 to 139 among unaffected sibs. Scores of NF1 patients ranged from 85 to 114 compared to 80 to 134 in unaffected parents. Children with NF1 showed significant deficits in language and reading compared to sibs without NF1. They also had impaired visuospatial and neuromotor skills. Foci of high signal intensity on T2 weighted MRI scan images were observed in 11 out of 12 NF1 children but in none of the unaffected sibs. They were seen particularly in basal ganglia, cerebellum, brain stem, and subcortical white matter. A statistically significant correlation was found between lowering of IQ and visuospatial deficits and the number of foci seen on scan. The authors postulate that the focal lesions may be causing multiple interruptions of brain pathways leading to cognitive deficits. There was no evidence to suggest that the presence of a focal lesion led to interference with specific function of that part of the brain involved, and as with other MRI abnormalities, further studies on larger patient groups will be needed to establish the nature and significance of these findings.

JILL CLAYTON-SMITH

Mutation of a mutL homolog in hereditary colon cancer

A cell cycle regulator potentially involved in genesis of many tumor types

The proposition that mutations in genes with important functions in the control of normal cellular growth and replication play a key role in the development of cancer finds clear support in both these reports. A DNA mismatch repair gene (hMSH2), recently recognised as a chromosomal gene involved in hereditary non-polyposis colorectal cancer (HNPPC) and microsatellite instability in tumour tissue; this gene's homology to a bacterial equivalent prompted this group to screen a commercial human DNA library for expressed sequences with homology to other bacterial or yeast mismatch repair loci. One of the homologous sequences mapped precisely to 3p21, a site with known linkage to HNPPC. This group then went on to sequence the gene at this site and use RTPCR to identify four different heterozygous germinal mutations. It is tumour cell line no wild type product was found, indicating that the gene designated hMLH1 is a classic tumour suppressor. The identification of specific mutations in hMLH1 will be of immediate benefit to HNPPC families. Mutations in this gene together with its chromosome 2 counterpart hMSH2 may also account for the majority of predisposing mutations in HNPPC which constitutes 4 to 13% of all colon cancers. Investigations of two further mismatch repair genes pulled out by the same screening technique will be awaited with interest. The second group also began