Genetic mapping of a gene causing hypertension in the stroke-prone spontaneously hypertensive rat

This paper describes genetic mapping of a gene causing hypertension in an animal model for primary hypertension in humans. Baseline and sodium loaded systolic and diastolic pressures were measured in a cross between the stroke-prone spontaneously hypertensive rat and the normotensive WKY strain. Using a single sequence repeat markers were analysed for linkage to each of these four blood pressure traits. Significant linkage was found between a marker (RD17) on chromosome 10 and both sodium loaded systolic (lod score = 4.88) and diastolic pressures (lod score = 5.10). Significant linkage was also observed between a marker on chromosome 18 and baseline diastolic pressure.

The locus on chromosome 10 (Bpl) was shown to be closely linked to the gene encoding angiotensin converting enzyme (ACE) which plays an important role in blood pressure homeostasis and is a target of antihypertensive drugs. This suggests the ACE gene as a possible candidate gene causing hypertension. The identification of the gene at Bpl and other genes involved in hypertension will obviously aid the understanding of the aetiology and pathogenesis of this common disorder. In addition, this study shows the possibility of performing genetic dissection in polygenic quantitative traits related to human disease in the mouse or the rat by crossing inbred strains to eliminate problems of heterogeneity and by using sufficient numbers of animals to achieve the statistical power to detect genes affecting quantitative traits.

N S THAKKER

Non-paternity and prenatal genetic screening

Prenatal screening programmes focusing on maternal AFP or fetal chromosomal anomalies are independent of paternity. Genetics service laboratories have long recognised the importance of correct paternity in prenatal testing for mendelian disorders. With population screening for cystic fibrosis now being evaluated, instances of wrong paternity are coming to light. Awareness of wrong paternity in population screening programmes is important not only in terms of evaluating costs and benefit of such schemes but, more importantly, in appropriate management where wrong paternity has altered the a priori risk. Estimates as to the true prevalence of wrong paternity are many. This paper evaluates the data from several sources, scientific and popular, whose estimates of wrong paternity range from 1.4% to 30%. It concludes that reliable data are currently lacking and emphasises the importance of wrong paternity considerations for service planners and those who provide for the context of population screening for cystic fibrosis.

W REARDON

A gene deleted in Kallmann's syndrome shares homology with neural cell adhesion and axonal path-finding molecules

Embryological evidence suggests that the cardinal features of Kallmann's syndrome (hypogonadotropic hypogonadism and anosmia) are related to a defect of neuronal migration at the olfactory placode. Autosomal dominant, recessive, and X linked forms of the disease have all been described. Franco et al have now characterised a gene (KALIG-1, in band Xp22.3) that is a strong candidate for the site of the X linked defect. The methodical collection and mapping of a series of affected patients with terminal Xp deletions or X;Y translocations enabled localisation of the gene to a ~80 kb interval. Making elegant use of standard techniques (identification of Cpg islands, zoo- touts, yeast artificial chromosome and cDNA libraries) the authors identified cDNAs within the critical region and constructed a full length sequence from overlapping clones that translates to a protein of 680 amino acids. Three domains, each comprising about 25 amino acids, show significant homology either to protease inhibitors/neurophysins, or to neuronal cell adhesion molecules/tyrrosine phosphates. The latter proteins have been implicated in cell adhesion and neurite outgrowth, supporting the notion that KALIG-1 may represent the first neuronal migration factor to be identified in humans. The authors do not explain why it appears to be ubiquitously expressed at low levels, but no doubt they are working on that.

ANDREW WILKIE

The phenylketonuria locus: current knowledge about alleles and mutations of the phenylalanine hydroxylase gene in various populations

The screening which detects and the diet which corrects the mental retardation which would otherwise be the untreated newborn with phenylketonuria is a paradigm of preventive medicine. In this detailed review, the authors summarise the molecular analysis of the mutations at the phenylalanine hydroxylase (PAH) locus on chromosome 12 which underlie the disease phenotypes. Using PAH cDNA as a probe shows eight RFLPs which indicate that specific mutations tend to segregate with specific haplotypes. Thirty-one different DNA alterations have been detected to date; only five are deletions while the remaining 26 are single base substitutions leading in most cases to missense mutations. Correlation of a particular activity of the PAH enzyme, and clinical phenotype indicate that not every homozygous mutation leads to PKU, while the phenotype resulting or to neural cell adhesion molecules. There are eight missense mutations which may alter function differences between the haplotype frequencies and their associated mutations north and south of the Alp. As with cystic fibrosis, any possible heterozygote advantage remains controversial.

T R P COLE

Localization of the gene encoding the GABA receptor β3 subunit to the Angelman/Prader-Willi region of human chromosome 15

Prader-Willi syndrome (PWS) and Angelman syndrome (AS) have recently become the focus of considerable interest from clinicians and molecular biologists. These two clinically distinct conditions are associated with cytogenetically similar deletions, 15q11-q13, Prader-Willi critical region (PWCIR), and uniparental disomy (UPD). In PWS the deletions are of paternal origin and UPD is maternal disomy, and vice versa in AS. To date no gene of known function had been mapped to the PWCIR. In this report, a genomic library was screened with bovine GABAα receptor β1 subunit. This identified two clones containing the homologous human GABAα receptor β3 subunit. These clones were used as probes and localised by in situ hybridisation to 15q11-q13. Dosage analysis of a panel of cytogenetically deleted PWS and AS patients identiﬁed hemizygosity for these two probes. Subsequent study of two subjects with unbalanced translocations, one with PWS and the other with AS, showed that both hemizygous regions had the classical markers in the PWCIR but that only the AS patient was deleted for the two new probes. It is believed defects in the GABA receptor function might cause seizures, movement disorders, and uncontrollable behaviour; these are the features of AS but not PWS. The possibility that GABRB3 might play a role in the aetiology of AS is an exciting step forward and ultimately could help explain the high frequency of UPD and higher recurrence risk in AS compared to PWS.

Association of nm23-H1 allelic deletions with distant metastases in colorectal carcinoma

Tumour invasion, lymphatic metastases, and distant metastases have a significant inverse association with patient survival in colorectal cancer. This association is inexact and does not allow accurate classification of patients into risk groups, thereby making prognostication difficult in the individual case. Cohen et al have now sought to identify the prognostic significance of alterations at the nm23-H1 locus at 17q12 in patients with colorectal cancer. The nm23 gene is known to show reduced expression in human breast cancers with nodal metastases. Its possible tumour suppressor role has been further supported by transfection studies. Eleven of the 21 patients entered in this study had deletions with the nm23-H1 probe. These 11 patients had a significantly higher incidence (p=0.03) of distant metastases than had their 10 counterparts who did not show deletion. Does the nm23-H1 locus have a role as a late acting tumour suppressor in colorectal cancer? Have the Cohen et al findings provided for the clinically valuable prognostic measure in colorectal cancer? A larger patient group and longer follow up time than the 25 months in this study will be required to answer these pertinent questions. The preliminary data look encouraging.

JOHN C K BARBER

W REARDON

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