HIGHLIGHTS

WEB WATCH

Sequence data wanted!

 http://www.gene.ucl.ac. uk/nomenclature/ workshop/virtual.html

 http://hgvbase.cgb.ki.se Can you help annotate the human genome? There are ~1,500 human genes for which the chromosome location is known but the actual coding sequence has not yet been identified. The HUGO Gene Nomenclature Committee (HGNC) is now looking to the scientific community to provide the sequences of these genes. To find a list of the genes in question and to submit data, visit their Virtual Gene Nomenclature Workshop web site. The deadline for data submission is 30 June 2003. This initiative is part of the HGNC's valiant project to provide unique symbols for the estimated 30,000 human genes. Now is your chance

to take part in this quest! The Human Genome Variation Database (HGVbase) provides another opportunity for researchers to contribute data to help annotate the genome. It aims to provide a comprehensive catalogue of the variation in the human genome. All types of polymorphisms can be submitted online regardless of chromosomal location, allele frequency or phenotypic effect. The web site is curated using both manual and automated curation tools and the quality of data is checked before entry. The database is run by an EU consortium, consisting of teams from the European Molecular Biology Laboratory, Germany, the European Bioinformatics Institute, UK, and the Karolinska Institute, Sweden. It provides a resource for those interested in the effect of genetic variation on drug responses and disease susceptibility. Indeed, data from HGVbase have already been used to identify genetic polymorphisms that are associated with benzene poisonina.

Catherine Baxter

CANCER GENETICS

Making sense of missense

BRCA1 mutations have been linked to an increased risk of breast and ovarian cancer. Most of the highly penetrant alleles identified encode truncated proteins but missense alleles have also been recorded their impact on cancer susceptibility is particularly difficult to assess because of low penetrance and the lack of adequate functional assays. In their paper, Fleming *et al.* describe an evolutionary approach to identify the missense alleles that are most likely to be associated with a disease phenotype.

The rationale behind this study is that mutations in functionally important amino acids are most likely to be associated with an increased risk of cancer. These key amino acids can be identified on the basis of their conservation in mammals, or from evidence of recent positive selection in the human lineage. To find conserved BRCA1 regions, the authors aligned Genbank sequences for exon 11 of BRCA1 from 57 mammals. Considering five amino-acid sites at a time, regions were defined as conserved if the first and last residues were fixed or conservative (that is, identical in all species or all residues sharing similar biochemical properties) and at least 80% of the sites were also fixed or conserved. Seven out of eight conserved regions were located in regions known to interact with other proteins. In addition, a conserved stretch of amino acids was identified in a region of unknown function that is also conserved in BRCA1 homologues from clawed frog and chicken.

If site conservation is a good indicator of its functional importance, one would predict that missense mutations affecting fixed sites,



or resulting in non-conservative substitutions at conservative sites, are most likely to be associated with a disease phenotype. In fact, 38 of the 139 documented missense alleles

HUMAN GENETICS

LD orienteering

Alleles at different loci are not those in linkage disequilibrium (LD) occur together in populations more often than is expected. The extent to which LD occurs in the human genome, and how this affects variation, will determine how easy it will be to map complex-trait loci through whole-genome association studies. Now, Stumpf and Goldstein use computer simulations to examine the block-like structure of LD and conclude that it is time to abandon the general idea of an average extent of LD in the human genome.

Initial assessments of how to use LD in genetic-association studies assumed a uniform recombination rate and an idealized demographic population history. But, theory is seldom exactly like real life. There has been much evidence that recombination rates vary across the genome, sometimes resulting in a block-like genome structure. Stumpf and Goldstein set out to model the effects of recombination hotspots, as well as demography, on the extent of the LD. To assess the interactions between recombination hotspots and demography the authors used simulations of populations that

undergo several bottle-necks that create LD and observed how the associations decayed under different intensities of recombination hotspots. The results show that the probability of the LD block-like structure is intimately linked with demography — severe bottlenecks delay the block-like genome structure whereas relatively high intensities of hotspots maintain it for long periods of time.

Not satisfied with simulations alone, Stumpf and Goldstein turned to real data. Having considered models of populations with different demographic histories — Europeans, Finns and Georgian Jews — the authors conclude that as a result of a strong interaction between demography and hotspots, the block-like structure of LD might be present in some



in exon 11 fall into one of these categories. The authors argue that these sites should be the priority for future studies, along with three missense mutations in residues that show signs of recent positive selection in human and primate lineages. Interestingly, these three amino acids lie in the region of BRCA1 that interacts with RAD51— a protein involved in double-stranded break repair.

The authors present a promising approach for prioritizing the study of missense mutations in BRCA1, as well as in other genes associated with heritable diseases. Indeed, the authors show that this method can be used to predict the ß-globin amino-acid residues that, when mutated, are associated with various globin pathologies. The true test of this approach, however, will be to determine how many of the 41 missense mutations highlighted by Fleming et al. are associated with an increased risk of breast and ovarian cancer.

Catherine Baxter Catherine Ba

(10.1073/pnas.0237285100) FURTHER READING Narod, S. A. Modifiers of risk of hereditary breast and ovarian cancer. *Nature Rev. Cancer* **2**, 113–123 (2002) WEB SITE

Elaine A. Ostrander's lab: http://www.gs. washington.edu/faculty/ostrander.htm

populations but absent from others. On this basis, they suggest that different statistical methods will be needed for association studies in different populations and/or gene regions. They also suggest that, depending on their demographies, different populations might be suitable for different stages of association mapping. The LD debate is far from over, but it is clear that we must be cautious about extrapolating from one gene region, and one population, to another.

Magdalena Skipper

Beferences and links

ORIGINAL RESEARCH PAPER Stumpf M. P. H. & Glodstein D. B. Demography, recombination hotspot intensity and the block structure of linkage disequilibrium. *Curr. Biol.* **13**, 1–8 (2003) **FURTHER READING** Ardlie, K. G. *et al.* Patterns of linkage disequilibrium in the human genome. *Nature Rev. Genet.* **3**, 299–309 (2002) WEB SITE

Centre for Population Genetics and Human Health: http://popgen.biol.ucl.ac.uk/people.html



IN BRIEF

EVOLUTIONARY GENOMICS

Parallel changes in gene expression after 20,000 generations of evolution in *Escherichia coli*.

Cooper, T. et al. Proc. Natl Acad. Sci. 100, 1072-1077 (2003)

Parallel changes in independent evolutionary lineages in response to an environmental challenge are clear indicators of adaptive evolution. Cooper *et al.* used DNA macroarrays to examine the changes in gene-expression in two lineages of *E. coli* in response to a glucose-limiting medium. The authors go on to identify a specific mutation accounting for many of the 59 genes that changed expression in parallel, raising the possibility that this might provide a general strategy for identifying the genes involved in adaptation.

PLANT GENETICS

Direct measurement of the transfer rate of chloroplast DNA into the nucleus.

Huang, C. et al. Nature 5 February 2003 (10.1038/nature01435)

We know that plant chloroplast genes can move to the nuclear genome, but how often does this happen? To answer this question, Huang *et al.* engineered tobacco chloroplast genomes with a gene that confers kanamycin resistance only if it is transferred to the nucleus. The authors found 16 out of 250,000 plants with independent nuclear insertions: a rate high enough to significantly impact nuclear genome organization and gene function, and to have implications for the design of genetically modified crops.

GENOMICS

Ringlike structure of the *Deinococcus radiodurans* genome: a key to radioresistance?

Levin-Zaidman S. et al. Science 299, 251-256 (2003)

D. radiodurans can withstand ionizing radiation at doses that are lethal to all other organisms. In this paper, Levin-Zaidman *et al.* describe the unusual 'toroidal' conformation of its genome. This tightly packed structure might confer resistance to ionizing radiation by enabling broken DNA strands to be held together tightly, facilitating template-independent repair.

TECHNOLOGY

A genomics-guided approach for discovering and expressing cryptic metabolic pathways.

Zazopoulos, E. A. et al. Nature Biotech. 21, 187-190 (2003)

Taking advantage of the fact that bacterial metabolic genes are clustered, Zazopoulos *et al.* developed a high-throughput method for identifying metabolic loci independently of their expression. First, using a genomic library, the authors generated 700-bp random genome-sequence tags (GSTs). Second, GSTs of interest were identified on the basis of sequence homology to the biosynthetic genes that are present in microbial databases. Selected GSTs were then used as probes to identify new metabolic genes (and their neighbours) from individual cosmids or BACs.