what we now know of the nucleotide sequence of globin mRNA. Though up to the present still mostly done in rabbits and mice, this work has now entered the realm of human haemoglobin, and comparisons can now be made between some nucleotide sequences of human α and β globin mRNA’s and the homologous regions in the rabbit. The co-ordination of synthesis of α and β chains and of haem and globin are described by Tim Hunt. J. Paul discusses haemoglobin synthesis at the level of chromatin and cell differentiation where the maturation of erythrocytes in mammals proceeds from pluripotent to committed cells destined to become morphologically identifiable red blood cells. There is also a valuable section on erythropoietin.

D. J. Weatherall and J. B. Clegg complement these chapters with an analysis of the molecular basis of the thalassaemias where the genetic systems described previously have gone astray. The clinical management of thalassaemia major is described by B. Modell and this includes intensive transfusion schemes as well as the effects of long-term chelating therapy and the prevention of thalassaemia itself by antenatal diagnosis and abortion.

M. F. Perutz mentions in his introduction that haemoglobin was the first molecule in which the transition from one form to another, fetal to adult, was described which still is one of the simplest manifestations of cell differentiation. If the mechanism of that transition could be subjected to outside control a therapy for sickle cell anaemia and thalassaemia could be developed. It is, therefore, fitting that there is also a final chapter by W. G. Wood on haemoglobin synthesis during human fetal development, where one may hope that in due course it will become possible to control the switch from fetal to adult haemoglobin with all its practical consequences.

This Bulletin should find its way into almost every biologist’s library. Haematologists must be greatly interested in the degree to which haemoglobin has been investigated and can now be integrated with the problems of disease and therapy. Anyone associated with genetics to the slightest degree will find this issue of the British Medical Bulletin indispensable.

HERMANN LEHMANN

The HLA System: An Introductory Survey.


This monograph from the Copenhagen Blood Bank and the Odense Clinical Genetics Unit on the HLA system is admirably clear and up to date. First the components of the system are described, the classic A, B, and C series and the MLC or D locus, the loci for the immune response genes, the loci for the complement components Bf, C2, and C4, and the loci for the Bg and Chido blood groups all close together on chromosome 6. Next the linkage disequilibrium between some of the alleles at the four HLA loci is described, the strongest being between some of those at the C and D loci. References are given to geographical variation; the A1 B8 haplotype, common in Europeans, is rare in other races. Amerindians have few of the well-known HLA antigens. Next the homologous systems in the mouse and rhesus monkey are described. Next a brief account of the biochemistry of the HLA of antigens is given. They are probably glycopeptides embedded in the cell membrane and bound to a molecule of β2-microglobulin. The gene coding for the β2-microglobulin is probably on chromosome 15. Next the importance of the HLA system for organ transplantation is described and the remarkable association of HLA alleles and a variety of diseases. The strongest of these still seems to be that of ankylosing spondylitis and B27, followed by Reiter’s syndrome and anterior uveitis with the same allele. This clearly separates this form of arthritis from other forms. The association of juvenile diabetes with B8 and B15 confirms the view that this form of diabetes is distinct from the more common form with onset usually in middle age. The authors think that the association with B27 must be direct since it has been found in three different races, and different races tend to show different gametic associations. The additive effects of B8 and B15 for juvenile diabetes, in contrast to the absence of such an effect for B13, 17, and 37 for psoriasis, suggests that in the former case the two alleles confer susceptibility by different mechanisms. The authors briefly describe the mechanisms that have been suggested for these associations. Next the authors describe the possible biochemical functions of the HLA system, for example the prevention of tumour formation and the incorporation in the cell membrane of viruses, so making these more easily recognizable by the body defences. Both mechanisms would help to account for the remarkable degree of polymorphism found at the HLA locus. The final chapter gives an outline of the laboratory methods involved in work with HLA.

This is an excellent introduction to a most intriguing development in human genetics.

C. O. CARTER

Biology of Radiation Carcinogenesis