are also of good quality with the exception of some of the low power shots and I would not be surprised if some of them occasionally find their way into lectures. On the whole, therefore, this book is highly recommended as a reference atlas and text for the diagnostic histopathologist faced with difficult proliferations in the duct system of the breast.

CLIVE WELLS


This is a single author short monograph reviewing the basic pathophysiology of the circulating platelet; it also gives the personal views of the author with respect to the role of platelets in rheology and blood flow. As usual with these publishers, this is a beautifully produced monograph with about 40 excellent black and white photomicrographs of platelets in various forms of activation. The text is aimed primarily at pre-clinical scientists, particularly physiologists and anatomists concerned with the basic concepts of platelet activation. Overall, it is a very readable and well referenced review but does not relate specifically to congenital or acquired disease states. In the last chapter the author discusses his own particular research interest which is the role of platelets in the development of thrombosis under different flow and shear conditions. This text is not specific enough for the general pathologist or haematologist interested in platelets but may primarily be of help to the platelet research worker who is involved in electron microscopy for studying platelet function and also for rheologists interested in the varying concepts of atheroma.

S J MACHIN


Familial adenomatous polyposis (FAP) is the term recommended by the Leeds Castle International Polyposis Group for a condition which serves as an important model for the study of cancer and cancer genetics in general. It is fitting that the person instrumental in stimulating the recent great leap in research activity and enlightenment in the field of polyposis should edit this book, the most important on FAP since Dr HJR Bussey's monograph of 1975. This is a compendium, with each chapter written by an authority on the topic in question. Forty seven chapters cover almost every conceivable aspect of FAP, often including information on wider but related issues. Of interest to pathologists are chapter 4 (Watte and Sohrabi) on all the various inherited colonic polyposis syndromes and chapter 5 (Hamilton) on the pathology of adenomas and malignant polypos. These are generally excellent, although I differ with the statement, "angio-lymphatic involvement is an indication for surgical resection" after carcinoma is found in a colonoepithelial polyp specimen. The later chapters cover specific matters of importance in the counselling of patients; the organisation of polyposis registries, and surgical and laser therapeutic methods. There are chapters on the important topics of extra-colonic manifestations (Bulow), comparative epidemiology with Japanese data (Utsumonita), and retinal lesions, with beautiful colour illustrations. The only omission is specific coverage of the management of adenoma and carcinoma of the ampulla of Vater, although these are alluded to in general terms.

The stature of the contributors has ensured that this is a reference work of the highest order. It should not be regarded as simply a book for the super-specialist; the later chapters are up to the minite (1990) reviews of a wide range of topics of interest and value to pathologists, including the molecular genetics, genesis and biology of neoplasia (such as mucins, lectins, oncogenes, protein kinases and, tissue culture technology). It is a book that libraries, large departments, and interested persons should buy.

IAN C TALBOT


Like any biological population, whether of cells or of whole organisms, the blood platelets are notably heterogeneous with regard to size, density, functional capacity, and to biochemical composition and survival. Platelet experts have argued for many years over the extent to which this heterogeneity is "congenital" or "acquired". It is predominately due to influences on their production from megakaryocytes or to stimuli which they encounter during their lifespan in the circulation? This argument formed the basis of a workshop held under the auspices of the European Society of Clinical Investigation in Graz in April 1988. The present volume contains papers written for publication after the meeting by the chief participants (and also by two groups who were unable to be present at the meeting) and much of the paper will be presented in this issue of The Journal of Pathology.

Although two and a half years have passed between the meeting and the publication of these proceedings, progress in this complex field has not been slow, so as to deprive them of much of their interest. The book provides much food for thought for all those interested in platelet production and ageing in health and in haemorrhagic and thrombotic diseases. Among the position-theoretical concepts advanced here which are still in need of experimental testing, perhaps the most challenging is that of Trowbridge, that platelets are shed from megakaryocytes, not in the bone marrow, as has been generally believed for the past 80 years, but in the pulmonary circulation.

RM HARDIStY

NOTICES

16th European Symposium on Hormones and Cell Regulation
September 23-26, 1991
Mont Ste Odile, Alsace, France
Topics include: nitric oxide and cyclic nucleotides; factors in the development of the nervous system; hormonal factors regulating embryonic development; GTP-binding proteins; receptors and channels; role of protein kinases and phosphoprotein phosphatases in signal transduction.
Further information from: Dr B Hamprecht, Physiologisch-chemisches Institut der Universität, Hoppe-Seyler-Str. 4, D-7400 Tübingen, Germany
Telefax: 49-7071-293361

12th Annual Conference of Australian Dermatopathology Society
Interface Dermatopathology
St Vincent's Hospital, Melbourne, Australia
Friday, 27 September - Saturday, 29 September, 1991
Guest Speaker: Professor AB Ackerman, New York University Medical School. The Conference will highlight clinical and laboratory contributions to diagnosis and will include five minute presentations and a poster display. Participation is invited.
For further information, please contact:
Mrs Ann Dancer, Victorian Medical Postgraduate Foundation Inc (VMFP), PO Box 27, Parkville, Vic 3052, Australia. Telephone: +61 3 347 9633 Fax: +61 3 347 4547

Association of Clinical Pathologists
Junior Membership
Junior membership of the Association is available to medical practitioners who have been engaged in the practice of pathology for a period of less than four years. Junior members are able to remain in this category for a maximum of six years or on the attainment of consultant status. The annual subscription is £24 for those resident in the United Kingdom and £55 for those overseas. The annual subscription may be claimed against tax.
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For Junior Membership apply to: The Honorary Secretary, Association of Clinical Pathologists, School of Biological Sciences, Falmer, Brighton, BN1 9QG. (0273) 678435
Familial Adenomatous Polyposis

Ian C Talbot

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