Book Reviews


A misleading title—the book records papers published at the 1973 Boston Conference on Diagnosis, Genetics and Management of Birth Defects, but is not a systematic review or bibliography of malformation syndromes. The chapter headings show it to include several general papers on nomenclature or classification of syndromes; these, by Warkany, McKusick, Smith, Herrmann, and Opitz, are of interest, and the last named particularly thorough in discussion of basic concepts: inborn error, dysplasia, malformation, etc., but, together with a check list for facial malformation syndromes, they account for only 34 pages (17%) of the book. The remainder is given up to papers of limited scope: case and family reports, and reviews of rare syndromes; and 7% of the book is virginal paper. It does not live up to expectations engendered by the publishers' puff '...the latest and most complete examination of malformation syndromes available...', and is not a good investment at the English equivalent of $14.50.

R. H. LINDBENBAUM


The once neglected field of bone dysplasias is now receiving widespread interest, and several excellent books have recently appeared which provide general reviews of the subject, despite our fragmentary knowledge of most of these disorders. This volume, the report of a conference held in Boston in 1973, does not attempt to cover the field systematically; however, the contents in general complement rather than duplicate the earlier volume on the subject produced by The National Foundation, and the standard of most contributions is high. Abundant radiographs and photomicrographs, reproduced with fair quality, add greatly to the value of the book, and the price is not excessive by present standards.

Several contributions deserve special mention. The opening chapter by Rimoin describes histological studies on bone and cartilage biopsies which in the past have been remarkably lacking and often confusing due to erroneous clinical diagnosis. Findings of particular interest to the medical geneticist are the identical appearance of changes in the mild 'variant' of diastrophic dwarfism to those of the classical disease, suggesting the same basic defect and possibly allelism; by contrast those familial cases of thanatophoric dwarfism studied histologically showed an entirely different appearance to those seen in isolated cases, suggesting that heterogeneity may be present, and that most isolated cases represent new dominant mutations.

The article by Spranger and Langer on spondyloepiphyseal dysplasias is valuable for its clarity in what seems at times a hopelessly confused area. They differentiate a number of disorders which can be considered well delineated, and separate them from the large remaining pool of conditions where classification is at best provisional. Pozanski and Garn tackle another neglected area, the quantitative measurement of radiological bone changes. They outline a series of key measurements and ratios which are likely to prove valuable, and which hopefully may replace the somewhat 'impressionistic' and subjective interpretation so often used at present.

Remaining chapters of the book give reviews and case reports on specific dysplasias such as Kniest syndrome, thoracic dysplasia and related disorders, and the tricho-phino-phalangeal syndrome. The increasing integration of histological, biochemical, and genetic data with clinical and radiological information shows that progress is steadily being made towards a fuller understanding of this complex group of inherited disorders.

PETER S. HARPER


This small March of Dimes volume is the text of the conference in Boston, presumably 1974, on syndromes mostly associated with chromosomal anomalies. O. J. Miller gives a clear account of the new banding techniques and W. R. Breg the application of these new methods to identify the individual chromosomes involved in conditions such as Down's, Edward's, and Patau's syndromes, and to identify chromosomes involved in translocations. He notes for example that the usual D/G translocation in Down's syndrome is between 14 and 21; that the deleted chromosome seen in chronic myeloid leukaemia is 22; that the D chromosome with the long arm deletion associated with mental retardation in retinoblastoma is 13. E. Early and W. Stanley report the first instance of an in vitro cell culture system (in the rabbit), in which non-disjunction of acrocentric chromosomes occurs spontaneously. P. Neurath describes the current status of computer analysis of chromosomes. He estimates that one needs to be doing 10,000 karyotypes a year to justify running a computer system with a scanner.

The second half of the book is concerned with a variety
of non–chromosomal genetic topics: cleft lip and palate; the present status of treatment of the mucopolysaccharidoses; new studies on mucolipidosis III; Menkes syndrome; the prune belly syndrome; familial cardiac lipidosis.

While an individual volume of these Boston conferences is perhaps not worth buying, the series may provide a useful continuous summary of advances in clinical genetics.

C. O. CARTER


Volume X in the series is an excellent indicator of the continuing expansion of medical genetics and its increasing practical importance. There are seven chapters whose subjects include the mutation rate in man (John Edwards), biochemical polymorphisms in animals (M. J. Siciliano, D. A. Wright, and Charles R. Shaw), inborn errors of the thyroid (John B. Stanbury), mucopolysaccharidoses (Elizabeth F. Neufeld), control of Tay Sachs' disease (M. M. Kaback, R. S. Zeiger, L. W. Reynolds and Marguerite Sonneborn), the XYY conundrum (D. S. Borgaonkar and S. A. Shah), and finally the evergreen topic of ethical issues in genetics (Robert M. Veatch). It is perhaps invidious to select from among consistently good articles, and one's own interests naturally colour one's selection. The chapter by Siciliano, Wright, and Shaw describing biochemical polymorphisms in animals is particularly fascinating. Having kept tropical fish for a number of years and observed at considerable expense their frequent mortality, I was greatly enlightened by the account of a single gene model for melanoma occurring in F1 hybrids of certain swordtails and platyfish. The descriptions of biochemical markers in these chromosomally undistinguished fish show the way that massive and apparently selectively neutral polymorphism can provide convenient markers for mapping. As the authors correctly point out animal models may vary in their relevance to man from being totally meaningless to being completely satisfying. For the most part their values lie in suggesting mechanisms or experiments which may be relevant in man and provide experimental systems which cannot be directly approached for ethical reasons. Borgaonkar and Shah have produced a nicely balanced account of the XXY chromosome male and discussed its merits as a syndrome. Though they are unable to resolve the issue, they provide a most valuable review and bibliography. Elizabeth Neufeld discusses the mucopolysaccharidoses and mucolipidoses storage disease with particular clarity. The other chapters are excellent, as I indicated earlier, and the whole volume deserves to be read from cover to cover.

R. HARRIS

Announcement

The Dr. Heinz Karger Memorial Foundation invites the submission of papers on the following subjects: 1976—an original research paper on 'Methods for the early diagnosis of genetic disorders'. 1977—an original research paper on 'Molecular biology of metabolic diseases'.

Conditions: Manuscripts shall not exceed 20 typewritten pages, including illustrations, tables, and bibliography. Manuscripts marked 'Competition' must reach the publishers, S. KARGER AG, Arnold-Böcklin-Strasse 25, CH-4011 Basel (Switzerland), not later than 28 February 1976 and 1977. The manuscript must be typewritten on one side only, double-spaced, and is to be submitted in quadruplicate, and in accordance with the instructions contained in The Manuscript (Rules for the preparation of manuscripts and bibliographies of scientific papers). This booklet can be obtained free of charge from the publishers if the request is marked 'Competition'. Language: English, German, or French. Publication: The winning papers will be published in English in one of the Karger journals. The award for the prizes will be SFr. 7000.00 each. The Council of the Foundation will judge the papers and confer the prizes.