

CONGENITAL HEMIPLEGIA. 2000. Edited by Brian Neville and Robert Goodman. Published by Cambridge University Press. 216 pages. C\$89.92 approx.

Congenital hemiplegia, a condition usually diagnosed in relatively early childhood, has been, over the years, and with developing technology, associated with a myriad of etiological factors. These factors can be prenatal or perinatal and while the clinical manifestations and their management may be relatively similar, the actual underlying disease may be as different as perinatal asphyxia, cortical migration disorder/schizencephaly or Sturge-Weber syndrome, to name a few.

In this book, the editors first successfully organize “congenital hemiplegia”, a term initially used in the pre-scan era, into a systematic classification based on clinical, neuro-pathological and MRI-based evidence. This is a major feat. They next address the clinical presentation and physical assessment of children outlining particular “gems” in the physical examination of such patients.

A substantial amount of the remaining chapters cover the therapeutic management of consequences of this condition, paying attention initially to orthopedic issues related to gait and the analysis of the particular pattern of dysfunction at the level of the affected lower extremity. Again, several key points relating to the examination of such patients and to the management options are brought out. They also cover upper extremity involvement and the management of epilepsy syndromes, which are a frequent consequence of congenital hemiplegia usually declaring themselves within the first few years of life. The last few chapters cover the management of emotional, social and educational issues for these children. This is also important because, as the authors point out, most children adapt well to their condition but none are cured of their deficits.

As a Pediatric Neurosurgeon interested in the management of cerebral palsy and spasticity as well as epilepsy, this fresh look at a relatively generous group of conditions under the hat of congenital hemiplegia is timely. The book bridges classical neurology with modern imaging and modern pathological classifications. It provides very helpful management information to the pediatric neurologist, pediatric neurosurgeon, pediatric orthopedist and physiatrist. It is also a very good reference text for psychologists, occupational therapists and physiotherapists dealing with these conditions. I would recommend it strongly for health professionals actively dealing with children with congenital neurological ailments from both the diagnostic and therapeutic ends. It may not be the best reference for the medical student starting to learn child neurology because it deals superficially with several very different diseases under the “umbrella” of congenital hemiplegia.

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DIAGNOSIS AND MANAGEMENT OF PERIPHERAL NERVE DISORDERS. CONTEMPORARY NEUROLOGY SERIES, NUMBER 59. 2001. Edited by Jerry R. Mendell, John T. Kissel, and David R. Cornblath. Published by Oxford University Press. 695 pages. C\$264.00 approx.

The Contemporary Neurology (or “Black Book”) Series of monographs on selected neurological topics has long been known to trainees and practitioners in neurology as an excellent resource for clinical neurology. The latest in this series, number 59, Diagnosis

and Management of Peripheral Nerve Disorders continues that fine tradition. The three principal authors are internationally renowned clinical and scientific experts in this field, and wrote or co-wrote nearly all the chapters in the book, creating a book with an evenhanded approach and consistently high content quality.

Part 1 of the book, Chapters 1 – 8, consists of chapters on general topics such as clinical approach, biopsy, other laboratory testing, and approach to painful neuropathy. Included is a unique chapter on autoantibody testing – a controversial topic in this field which the authors handle well. I might have liked a chapter here on the practicalities of using immune suppressive treatment. Part 2 of the book includes 23 chapters focusing on individual diseases or groups of diseases causing peripheral neuropathy, including two chapters on entrapment neuropathy and plexopathy.

The book has many outstanding features. Probably most striking is the quantity and quality of illustrations. As indicated by the authors in their preface this is an intentional emphasis which proves well worth their efforts. The coverage of topics is very comprehensive. These authors are very active in clinical management of patients, and this is evident in the frequent use of case reports, discussion of clinical dilemmas, and a rational approach to treatment. In some cases (especially with paraprotein associated neuropathies) I might have liked the details of immune suppressive management to have been fleshed out better in some cases. With authors of such experience I would have been interested in their personal opinion in addition to the brief literature review presented.

The book takes the approach of starting with a specific disease, and discussing its manifestations. This is done very well. However, except for a chapter on painful neuropathies there is no discussion from the starting point of a clinical syndrome; the neurology resident looking for a discussion of differential diagnosis of GBS or CIDP will largely be disappointed.

In summary, this is an excellent monograph which highlights the many advances that have been made in understanding, diagnosis and management of peripheral nerve disease in the past decade. When asked by neurology residents or non- subspecialist colleagues what single book they should read cover to cover on the topic of peripheral neuropathy, I will indicate that this is the one.

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CLINICAL DIAGNOSIS AND MANAGEMENT OF ALZHEIMER’S DISEASE. Second Edition. 1999. Serge Gauthier, Ed published by Martin Dunitz, London. 386 pages \$C185.00 approx.

There are some books that one would expect to find on the shelves of medical school and hospital libraries. This volume should be one of those. Physicians and others experienced in the care of individuals with dementia will appreciate how relevant each and every chapter is to the day-to-day management of these disorders.

Our knowledge of the biological substrates of Alzheimer’s disease and of approaches to the clinical management of the disorder has expanded rapidly. The treatment of the topic in this volume is unexcelled in its clarity of exposition and in bringing information of clinical relevance from diverse fields of study to the forefront.

The volume is organised in seven sections including introductory chapters touching on pathophysiology, diagnosis, natural history,