

drive receptor-mediated siderophore uptake. These chapters also provide a more limited description of iron uptake pathways that do not involve siderophores, such as uptake of heme and ferric citrate.

A second major grouping of six chapters, representing about 30% of the book, focuses on the proteins involved in mammalian iron transport and storage: ferritin, transferrin, and the transferrin receptor. There are three very complementary chapters on ferritin. One chapter by Harrison et al. focuses heavily on crystallographic studies of the protein structure, while a second chapter by Chasteen puts more emphasis on iron binding and oxidation. Both of these chapters include some discussion of bacterioferritin. The third chapter by Powell is devoted exclusively to mineralization and iron core formation, with an interesting discussion of the edge effects associated with the very small iron cores. There are also chapters on transferrin and the transferrin receptor and on the iron responsive element on mRNA's and the genetic control of iron uptake. The final chapter in this group by Crichton and Ward covers the broad topic of iron homeostasis and serves as a nice summary of much of the material covered in the other chapters in this group.

This book also attempts to cover iron transport and storage in plants. Thus there are chapters on the role of both microbial and plant siderophores in plant iron uptake and on plant ferritin. However, the coverage of iron metabolism in plants is not that extensive and somewhat fragmented. Other than a general discussion of the role of proteins and siderophores in limiting the iron available to pathogenic organisms and one chapter on clinical iron chelators, there is not much coverage of clinical issues.

Wesley R. Harris

*Department of Chemistry
University of Missouri—St. Louis
8001 Natural Bridge Road
St. Louis, Missouri 63121*

JM980653E

10.1021/jm980653e

Progress in Medicinal Chemistry, Volume 35.

Edited by G. P. Ellis, D. K. Luscombe, and A. W. Oxford. Elsevier Science B.V., Amsterdam, The Netherlands. 1998. vii + 270 pp. 14.5 × 21 cm. ISBN 0-444-82909-1. \$198.50.

This book is the latest volume in this well-established medicinal chemistry review series. Chapters in this volume include: Modern View of Vitamin D₃ and its Medicinal Uses by M. J. Beckman and H. F. DeLuca (56 pages, 410 references), Neurokinin Receptor Antagonists by C. J. Swain (24 pages, 75 references), Opioid Receptor Antagonists by H. Schmidhammer (49 pages, 207 references), Mechanisms of Bacterial Resistance to Antibiotics and Biocides by A. D. Russell (64 pages, 326 references), and Towards Cannabinoid Drugs - Revisited by R. Mechoulam, L. Hanus, and E. Fride (44 pages, 202 references). The first three chapters cite references up to and including 1996, the last two, 1998. The volume contains a 6-page subject index.

The chapter on vitamin D₃ includes a detailed description of its biosynthesis, metabolism, and current understanding of mechanism of action, followed by a discussion of the therapeutic uses of vitamin D₃ and its analogues. In the last section of the chapter, the authors discuss the potential of vitamin D₃ analogues in the treatment of cancer and immune-related diseases.

The next chapter reveals the therapeutic potential and structural diversity of the many high-affinity neurokinin (NK) receptor antagonists. The authors classify the antagonists as NK₁, NK₂, or NK₃ receptor-selective or as dual NK₁/NK₂ receptor-selective. In the body of the text, useful comparative quantitative receptor affinity and pharmacological data are presented.

The chapter on opioid receptor antagonists begins with a description of receptor antagonism and universal opioid receptor antagonists. The remainder is divided into three sections: μ -opioid, κ -opioid, and δ -opioid receptor-selective antagonists. In each of the three sections, opioid antagonists are classified as peptide or non-peptide and, within these subheadings, as competitive or irreversible/long-acting. Binding affinity and receptor selectivity data are presented in 11 tables. Concise descriptions of the synthesis of the opioid antagonists are presented throughout the text.

The chapter on antibiotic resistance begins with some definitions and in turn discusses the mechanisms of intrinsic resistance, mechanisms of acquired resistance, pathogenic-resistant Gram-positive bacteria, mechanisms of multidrug resistance in Gram-negative bacteria, linked antibiotic-biocide resistance, and overcoming bacterial resistance. The use of drug combinations as a strategy to overcome bacterial drug resistance is only briefly mentioned, and the stereochemistry for some of the structures is not indicated. Despite these minor flaws, this chapter presents some well-organized information on this important topic.

The chapter on cannabinoid drugs describes cannabinoid receptors and the SAR of both the classical cannabinoids and the endogenous ligands, anandamide and 2-arachidonoyl glycerol. This is followed by a discussion of the medicinal properties of the cannabinoids which includes a discussion of the recently discovered oleamide and its potential in the treatment of sleep disorders. An interesting perspective is also provided on the ongoing debate over "medical marijuana."

The quality of the volume is very good. This book will likely be quite useful not only to those familiar with the topics presented but also to those who wish to learn more about these subjects.

Jonathan L. Vennerstrom

*College of Pharmacy
University of Nebraska Medical Center
Omaha, Nebraska 68198-6025*

JM9806547

10.1021/jm9806547

Integration of Pharmaceutical Discovery and Development: Case Histories. Edited by Ronald T. Borchardt, Roger M. Freidinger, Tomi K. Sawyer, and Philip L. Smith. Plenum Press, New York. 1998. xxix + 607 pp. 16 × 23 cm. ISBN 0-306-45743-1. \$125.00.