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Sommario/riassunto

"This book, in 15 chapters, presents fundamental facts alongside practical and clinically-related data. Contributors bring expert perspectives in the field of pharmacokinetics, pharmacodynamics, mathematical and computational science to the coverage. The topics include: optimization of drug dosing regimen in specialized populations, model-based approaches in drug treatment among pediatrics, pharmacotherapeutic considerations in pregnancy and lactation, and therapeutic drug monitoring in special populations. Pavadharini Ravindran cipsupportteam@wiley.com 914443950500"--
