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**The Best Pharmaceuticals for Children Act (BPCA) and the Pediatric Research Equity Act (PREA) were designed to encourage more pediatric studies of drugs used for children. The FDA asked the IOM to review aspects of pediatric studies and changes in product labeling that resulted from BPCA and PREA and their predecessor policies, as well as assess the incentives for pediatric studies of biologics and the extent to which biologics have been studied in children. The IOM committee concludes that these policies have helped provide clinicians who care for children with better information about the efficacy, safety, and appropriate prescribing of drugs. The IOM suggests that more can be done to increase knowledge about drugs used by children and thereby improve the clinical care, health, and well-being of the nation's children.**

**Pediatric Drug Development, Second Edition, encompasses the new regulatory initiatives across EU, US and ROW designed to encourage improved access to safe and effective medicines for children. It includes new developments in biomarkers and surrogate endpoints, developmental pharmacology and other novel aspects of pediatric drug development.**

**Provides a concise yet detailed resource covering all aspects of pharmaceutics, from the scientific fundamentals to the dosage forms and drug delivery systems to drug product analyses. Assists with integrating the science of pharmacy into practice. Chapters from the original parent text Remington: The Science and Practice of Pharmacy 22nd edition were specifically selected to create this new edition. The text pulls heavily from the Pharmaceutics and Pharmaceutical Dosage Forms sections. Various delivery systems and dosage forms are covered as well as parenterals, sterilization processes, and sterile compounding. One chapter addresses pharmaceutical excipients and another discusses pharmaceutical packaging. Pharmaceutical analysis, product characterization, quality control, stability, bioavailability, and dissolution are also covered. Fundamental scientific concepts including thermodynamics, ionic solutions and electrolyte equilibria, tonicity, chemical kinetics, rheology, complex formation and interfacial phenomenon are presented.The text also provides an introduction to pharmacokinetics and pharmacodynamics and the principles of absorption, distribution, metabolism and excretion.In addition, some introductory concepts on drug discovery and drug product approval as well as information resources in pharmacy and the pharmaceutical sciences are presented.**

**Market access is the fourth hurdle in the drug development process and the primary driver for global income of any new drug. Without a strategy in place for pricing, showing value for effectiveness and an understanding of the target purchasers ' needs, the drug will fail to reach its intended market value. Introduction to Market Access for Pharmaceuticals is based on an accredited course in this area, taken from the European Market Access University Diploma (EMAUD), and is affiliated with Aix Marseille University.**

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