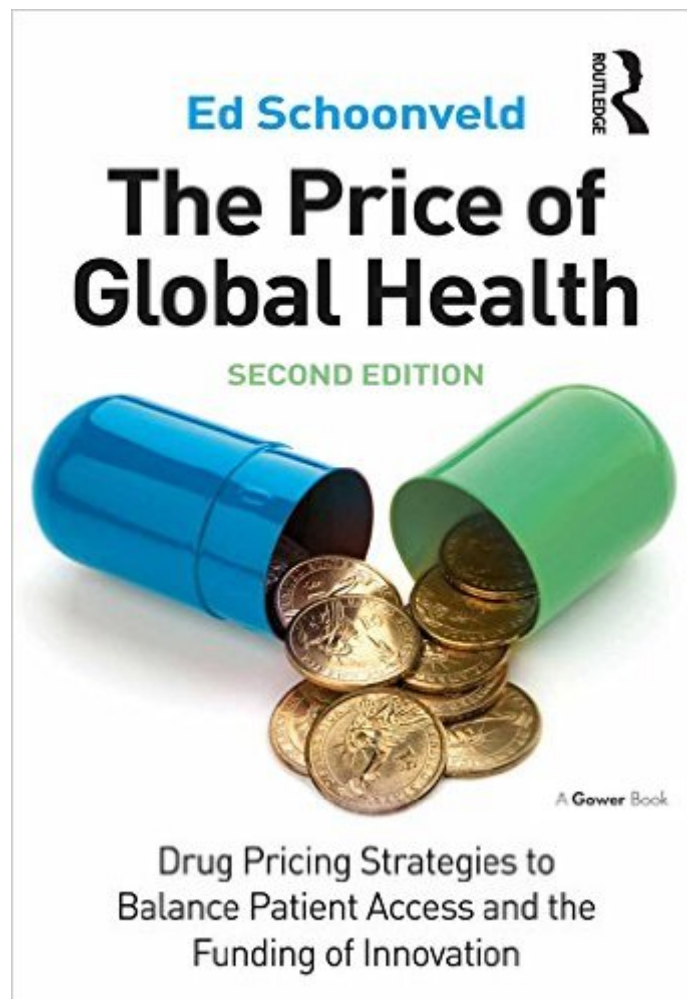


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The Price Of Global Health: Drug Pricing Strategies To Balance Patient Access And The Funding Of Innovation



Synopsis

Public debate on the rising cost of new biotechnology drug treatments has intensified over the last few years as healthcare budget pressures have mounted under a strained economy. Meanwhile, the demand for new, effective medical and drug treatments continues to rise as unhealthy lifestyles cause further increases in diabetes and cardiovascular disease. Global drug pricing is one of the most hotly debated yet least understood aspects of the pharmaceutical industry. How should drug prices be set and what does it mean for patients? Why do governments increasingly get involved, and what is its impact on the global competitive environment? How can a life-saving industry have a poorer image than gun and tobacco industries, whose products are associated with death? Ed Schoonveld explains how pharmaceutical prices are determined in a complex global payer environment and what factors influence the process. His insights will help a wide range of audiences, from healthcare industry professionals to policy makers and the broader public, to gain a better understanding of this highly complex and emotionally charged field. The Price of Global Health is recognized as a valued and unique reference book that covers a complete array of topics related to global pharmaceutical pricing. It contains an in-depth but straightforward exploration of the pharmaceutical pricing strategy process, its underlying market access, general business and ethical considerations, and its implications for payers, physicians and patients. It is a much-needed and invaluable resource for anybody interested or involved in, or affected by, the development, funding and use of prescription drugs. In particular, it is of critical importance to pharmaceutical company executives and other leaders and professionals in commercialization and drug development, including marketing, business development, market access and pricing, clinical development, drug discovery, regulatory affairs, health outcomes, market research and public affairs. The second edition includes new chapters on payer value story development, oncology, orphan drugs and payer negotiations. Furthermore, many country chapters have been substantially updated to reflect changes in the healthcare systems, including the Affordable Care Act in the US, AMNOG in Germany, medico-economic requirements in France and many other country-specific changes. Lastly, almost every chapter has been updated with new examples and illustrations.

Book Information

File Size: 8180 KB

Print Length: 494 pages

Simultaneous Device Usage: Up to 4 simultaneous devices, per publisher limits

Publisher: Routledge; 2 edition (February 24, 2016)

Publication Date: February 24, 2016

Sold by:Â Digital Services LLC

Language: English

ASIN: B01C677KO8

Text-to-Speech: Enabled

X-Ray: Not Enabled

Word Wise: Enabled

Lending: Not Enabled

Enhanced Typesetting: Enabled

Best Sellers Rank: #771,358 Paid in Kindle Store (See Top 100 Paid in Kindle Store) #95

inÂ Books > Business & Money > Industries > Pharmaceutical & Biotechnology #122 inÂ Kindle Store > Kindle eBooks > Business & Money > International > Global Marketing #377 inÂ Books > Business & Money > International > Global Marketing

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Contemporary Drug Information: An Evidence-Based Approach (Gaenelein, Contemporary Drug Information) Lupus Underground: A Patient's Case for a Long-Ignored, Drug-Free, Non-Patentable, Counter-Intuitive Therapy That Actually Works - UVA1 Phototherapy Entry Threats and Pricing in the Generic Drug Industry ASTNA Patient Transport: Principles and Practice, 4e (Air & Surface Patient Transport: Principles and Practice)

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