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• 书 讯 •

《世界农药新进展（四）》

张一宾 徐晓勇 张 恽 主编

本书作为 2007 年出版的《世界农药新进展》、2010 年出版的《世界农药新进展（二）》和 2014 年出版的《世界农药新进展（三）》的续篇，保持了内容的连贯性和形式的统一性。本书针对当前世界农药发展的新情况和新特点，按农药的各大类、各地区和主要国家、主要适用作物、主要农药公司市场和新品种做了全面而又系统的介绍，并对新农药品种、专利期已过或即将到期的重要品种，特别是其合成方法进行了重点阐述与分析。此外，还专门介绍了农药剂型加工及研究进展。

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