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· 消息 ·

《实用儿童脑病学》新书介绍

儿童脑损伤和脑病是危害儿童健康成长和人口素质的主要疾病,严重影响儿童社会交往能力、适应能力、注意力和学习能力,甚至导致癫痫、视听障碍、语言障碍、智力障碍、脑性瘫痪、植物状态等神经伤残。

由陈光福教授主编、人民卫生出版社出版的《实用儿童脑病学》,全书共30章,139万字,460多幅图(部分彩图),涉及儿童神经发育、神经解剖、神经影像学检查、脑功能检查、神经学检查评估、神经保护治疗、神经免疫治疗、神经修复治疗与康复治疗,以及各种儿童脑疾病的病因、发病机制、病理、临床表现、诊断方法与治疗进展等内容。

本书编委为从事儿童神经发育基础、神经影像、神经电生理、神经内科、神经外科、儿童康复和新生儿等专业的专家,专家们历时2年精心撰写,反复修改、补充,力求反映本领域近5年国内外的最新研究进展、诊治指南和专家共识,以及作者的临床经验与研究成果,并对新的探索性治疗应用前景进行了述评和展望。全书图文并茂、条理清晰、内容丰富、可读性强,是一部具有先进性、科学性和实用性的专业参考书。

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