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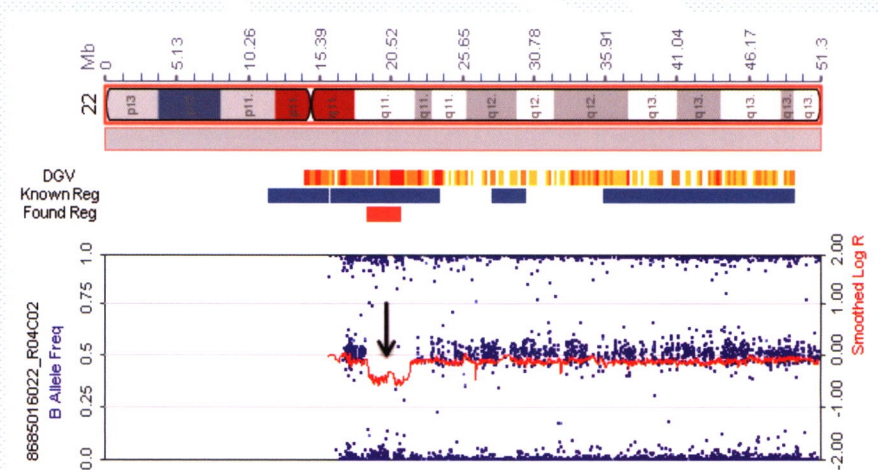
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注:箭头指向的区域表示22q11.2微缺失区域。

图(P420) 病例2诊断为22q11.2微缺失综合征的SNP-array结果

Figure(P420) SNP-array analysis results analysis diagram of case 2 diagnosed with 22q11.2 microdeletion syndrom

中山大学 主管

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杂志简介：

《分子诊断与治疗杂志》于2009年5月创刊，是由中山大学主管，《中国家庭医生》杂志社有限公司主办，中山大学达安基因股份有限公司承办的面向国内外公开发行的分子医学权威性刊物，也是我国第一份以分子诊断与治疗学科为主要内容的分子医学专业学术期刊。本刊以传播分子医学理念、报道分子医学前沿、倡导分子医学创新、促进分子医学进步为办刊宗旨。主要刊登我国医学分子诊断和分子治疗相关领域中的基础理论研究、临床实践和技术方法的最新研究成果以及国内分子医学新技术、新理论和新进展等方面的文章。目前有编委及审稿专家80余人。在编委会和审稿专家及编辑部的共同努力下，于2016年被收录为“中国科技核心期刊”（中国科技论文统计源期刊）。

目前有“述评”、“论著”、“综述”、“前沿进展”、“专家笔谈”、“讲座”、“动态”和“专栏”等栏目。涵盖基因诊断、免疫诊断、核酸分子诊断、蛋白分子诊断、信号分子诊断、分子影像诊断、诊断仪器应用、基因治疗、核酸分子治疗、蛋白分子治疗等内容。

