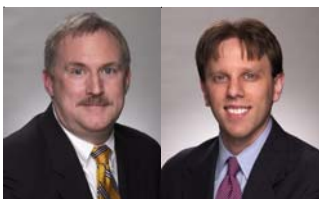


Noteworthy...

at Saul Ewing LLP

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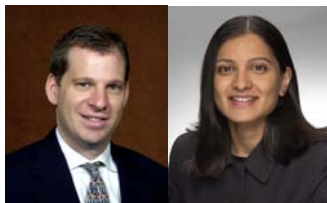
Life Sciences Practices



New Medical Device Compliance Handbook. Partner Mark Levy, with assistance from Associate Randy Undercofler, drafted a Compliance Training Handbook for Medical Device Sales Representatives. The handbook provides an overview of public and private compliance guidelines and the relevant laws and regulations that affect medical device companies and their sales representatives. The informative handbook was published by ePharmaceuticals, a division of HCPPro.



Notable Award and Presenter. Partner John Reiss was presented with the prestigious Yerger award by the New Jersey HFMA Chapter for his long-standing involvement with the organization. In March 2006, John presented a seminar entitled, “NJ HFMA Governance Survey Results/Hot Governance Issues and Best Practices” at the New Jersey HFMA Quarterly meeting, “Corporate Compliance - The Right Thing to Do and How to Do It Right!”



Medical Staff Briefing Q&A. Partner Bruce Armon and Associate Anjali Kharod have recently been quoted in the Medical Staff Legal Advisor Question & Answer section, a weekly publication prepared by HCPPro. Bruce and Anjali’s answers are directed towards hospital credentialing professionals.

Biotechnology | Healthcare | Medical Devices | Pharmaceuticals