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**APPENDIX TO CURRICULUM VITAE**

**PAST SPEAKING ENGAGEMENTS**

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***Understanding FDA Regulation of Mobile Medical Applications.*** RAPS Annual Convention. September 30, 2014, Austin, Texas.

***Update on U.S. Regulation of Biosimilars.*** RAPS Annual Convention. September 29, 2014, Austin, Texas.

***Crisis Management for Regulatory Professionals.*** Regulatory Managers Bootcamp. RAPS Annual Convention. September 28, 2014, Austin, Texas.

***Clinical Trial Liability Concerns for Small Companies.*** Arena International's 2nd Annual Outsourcing in Clinical Trials Southern California, La Jolla, CA, September 23-24, 2014.

***FDA Regulation of Drug and Device Advertising & Promotion.*** Two-Day Course. ComplianceOnline. September 11-12, 2014. Chicago.

***Combination Products, Dietary Supplements, and Veterinary Medicine.*** SDRAN RAC Review Course. August 13, 2014. San Diego.

***Overview of FDA Regulation of Medical Devices.*** DreamIt Philadelphia Healthcare Companies. Webinar. August 11, 2014.

***ANDAs, Orphan Drugs, OTC Drugs and Cosmetics.*** SDRAN RAC Review Course. August 6, 2014. San Diego.

***History of U.S. Regulation.*** OCRA RAC Review Course. August 2, 2014. Irvine, CA.

***FDA Regulation of Labeling and Advertising, Import/Export, and Enforcement.*** OCRA RAC Review Course. August 2, 2014. Irvine, CA.

***FDA Enforcement of GCP Requirements: A Review of Key Warning Letters (2010-2012).*** DIA GCP QA Community. Webinar. July 24, 2014.

***Regulation of Clinical Research in the U.S.*** Barney & Barney In-house Briefing. July 7, 2014. San Diego.

***Leaping the Valley of Death: Keys to Going from the Lab to the Clinic.*** Moderator. DIA Annual Conference, June 17, 2014. San Diego. [no slides; moderated panel]

***Understanding the Sunshine Act.*** DIA Annual Conference. Tutorial. June 15, 2014. San Diego.

***Responding to FDA Warning Letters.*** Duane Morris Webinar. April 30, 2014.

*Global Regulatory Considerations in Clinical Trials.* National Institutes of Health In-House Seminar. March 26, 2014. Bethesda, MD.

*FDA Regulation of Drug and Device Advertising & Promotion.* Course Instructor, Two-Day Course. ComplianceOnline. March 6-7, 2014. San Diego.

*Orphan Drug Regulation.* 2nd Annual Orphan Drugs Research & Commercialization Conference. GTCBio. Feb. 20-21, 2014. San Diego, CA. [no slides; panel discussion].

*U.S. Regulation of Biosimilars.* 2nd Annual Biosimilars And Follow-On Biologics 2014 Americas Conference. February 10-12, 2014. Philadelphia.

*Moderator, Entrepreneur Panel.* 11th BVS La Jolla Biotech Day. Biotech Vendor Services. December 11, 2013. San Diego.

*FDA Regulation of Drug and Device Advertising & Promotion.* Course Instructor, Two-Day Course. ComplianceOnline. November 18-19, 2013. Boston.

*Digital Health and FDA.* Commercializing Software IP: High Tech, Digital Health & Education Conference. Jointly Sponsored by UCSF, UC Berkeley, and the LES Silicon Valley Chapter. November 13, 2013. Berkeley, CA.

*FDA Regulation of Social Media.* MetricStream Webinar. November 6, 2013.

*FDA Enforcement in the Clinical Research Arena.* Compliance2Go Webinar. October 30, 2013.

*Review of FDA's Final Guidance on Mobile Medical Applications.* DIA Annual Conference Program Committee Audio Conference. October 17, 2013. [no slides]

*FDA Enforcement in the Clinical Research Arena.* ACRP San Diego Chapter. September 19, 2013. San Diego.

*Regulation of Combination Products, Dietary Supplements, and Veterinary Products.* SDRAN RAC Review Course. September 4, 2013. San Diego.

*Generic Drug Approvals.* Center for Professional Advancement 3-Day Course. August 13-15, 2013. New Brunswick, NJ.

*Regulation of Generic, OTC and Orphan Drugs, and Cosmetics.* Lecture at SDRAN RAC Review Course. August 7, 2013. San Diego.

*The Gamechanger: The Impact of the Generic Drug User Fee Act.* ACI Legal & Regulatory Summit. July 17 and 18, 2013. New York City.

*Creative Strategies in Dealing With FDA.* PSC Creative Learning Webinar. July 10, 2013.

*Informed Consent: Promise, Pledge, Contract or Platitude?* DIA Annual Conference. June 27, 2013. Boston.

***The De Novo 510(k) Process.*** DIA Annual Conference. June 26, 2013. Boston.

***Regulatory, Clinical and Quality Challenges in Contracting and Due Diligence: The Forgotten Keys to Biopharma Transactions.*** DIA Annual Conference. June 2013. Boston.

***FDA Enforcement.*** DIA Annual Conference. Tutorial. June 23, 2013. Boston.

***FDA Regulation of Social Media.*** OCRA Annual Conference. June 12, 2013. Irvine, Ca.

***Dietary Supplements -- Overview of Key FDA Issues.*** Compliance2Go. Webinar. May 30, 2013.

***Regulatory Convergence: Impact of FCC, HIPAA/Privacy and FDA on Mobile Health and Medical Devices.*** Fx Conferences Audio Conference. May 21, 2013.

***Combination Products -- Regulatory & Quality Challenges.*** Joint ASQ/RAPS SF Area Chapter Annual Conference. May 17, 2013. Santa Clara, California.

***Clinical Trials: Regulatory and Privacy Issues.*** THE BLUEPRINT™ Webinar Series, jointly sponsored by the California Healthcare Institute and Duane Morris LLP. May 14, 2013.

***FDASIA -- Challenges & Opportunities for Drug and Medical Device Companies.*** Compliance2Go. Webinar. April 16, 2013.

***Regulatory Pitfalls in Product Development.*** THE BLUEPRINT™ Webinar Series, jointly sponsored by the California Healthcare Institute and Duane Morris LLP. April 16, 2013.

***Regulatory & Quality Challenges of Virtual Drug Development; Or How to Avoid Getting in Bed With the Devil.*** Strategies for Success in Virtual Drug Development. A BDC/BVS/PGC 2000 Conference. April 15, 2013. San Diego.

***FDASIA -- Update on FDA Implementation.*** American Society for Quality, San Diego Chapter. March 12, 2013.

***Creative Strategies in Dealing with FDA.*** Compliance2Go. Webinar. February 21, 2013.

***Responding to FDA Inspections and Warning Letters.*** Compliance2Go Webinar. February 7, 2013.

***FDA Regulation of Drug and Device Advertising & Promotion.*** Course Instructor, Two-Day Course. ComplianceOnline. January 24-25, 2013. San Francisco.

***FDASIA -- Challenges and Opportunities for FDA-Regulated Industries.*** OCRA. December 4, 2012, Irvine, CA.

***FDA and Social Media.*** Compliance2Go. Webinar. November 27, 2012.

***FDA Update: Impact of FDASIA and the Federal Elections.*** SDRAN. November 15, 2012, San Diego.

***Legal and Regulatory Update – Health Care Reform and FDA Developments.*** BIOCOM. November 13, 2012. San Diego.

***Regulation of Dietary Supplements, Combination Products, and Veterinary Products.*** SDRAN RAC Review Course. August 29, 2012. San Diego.

***Regulation of ANDAs, Orphan Drugs, OTC Drugs and Cosmetics.*** SDRAN RAC Review Course. August 1, 2012. San Diego.

***Responding to FDA Inspections and Warning Letters.*** FxConferences Audio Conference. July 10, 2012.

***Leaping the Valley of Death: Keys to Successfully Going From the Lab to the Clinic for Pharmaceutical Products:*** Drug Information Association (DIA) Annual Conference, Session Chair. June 26, 2012. Philadelphia.

***Biosimilar Regulation and CMOs.*** FierceBiotech Webinar. June 26, 2012.

***FDA Enforcement.*** Pre-conference Tutorial at DIA Annual Conference. June 24, 2012. Philadelphia.

***How Do You Look in Stripes? FDA Enforcement Today.*** Joint FDA/Orange County Regulatory Affairs (OCRA) Annual Conference. June 7, 2012. Irvine, CA.

***The 510(k) Premarket Notification Process.*** FDLI Introduction to Medical Devices Conference. June 5, 2012, Palo Alto, CA.

***Crisis Management for Life Sciences Executives.*** FxConferences Audio Conference. May 22, 2012.

***Internet Issues for Regulatory Professionals -- FDA Regulation of Social Media.*** Orange County Regulatory Affairs Discussion Group. April 17, 2012, Irvine, CA.

***FDA – Creative Strategies in Dealing with The Agency.*** Panel Discussion. March 26, 2012. San Diego.

***Biosimilars -- Wave of the Future or Child of The Privileged Few?*** Licensing Executives Society San Diego Chapter. February 21, 2012.

***The de novo 510(k) Process -- The Impact of the New 2011 FDA Guidance.*** FxTranslations Audio Conference. February 15, 2012.

***Crisis Management for Regulatory Professionals.*** Regulatory Affairs Professionals Society, Rising Leaders Program. Audio Conference. December 15, 2011.

***FDA Enforcement Activities In Clinical Trials Arena.*** FxTranslations Audio Conference. November 3, 2011.

***Get to the Clinic on Time.*** LARTA NIH-CAP Commercialization Workshop, November 1, 2011. Los Angeles.

***An Overview of Issued FDA Warning Letters: What Happened and What Can be Learned.*** ExL Pharma's 2nd Quality Oversight of Clinical Vendors Conference. October 18, 2011. Washington, D.C.

***Regulatory Challenges in Executing Global Clinical Studies.*** The Conference Forum's 2<sup>nd</sup> Annual Executing Global Clinical Trials Conference. September 15, 2011, Philadelphia.

***Regulation of Dietary Supplements.*** Lecture at SDRAN RAC Review Course. August 25, 2011. San Diego.

***Regulation of Generic, OTC and Orphan Drugs, and Cosmetics.*** Lecture at SDRAN RAC Review Course. July 27, San Diego.

***Quality Aspects of Due Diligence for Biopharmaceutical Transactions.*** DIA Annual Conference. June 21, 2011, Chicago.

***FDA Enforcement.*** Tutorial at DIA Annual Conference. June 19, 2011, Chicago.

***Biosimilars.*** Orange County Regulatory Affairs Discussion Group/FDA Annual Educational Conference. June 9, 2011, Irvine, CA.

***Regulatory, Clinical and Quality Challenges in the Regulation of Combination Products.*** The Weinberg Group Inc. Webinar, May 18, 2011.

***Clinical Trial Registries.*** Association of Clinical Research Professionals Annual Conference. May 3, 2011, Seattle.

***IRB Liability.*** Association of Clinical Research Professionals Annual Conference. April 30, 2011, Seattle.

***Overview of FDA Issues for In-Vitro Diagnostics.*** Southern California Biotech Assn. April 13, 2011, Costa Mesa, CA.

***Challenges of Orphan Drug Regulation.*** The Weinberg Group Inc. Webinar, February 23, 2011.

***FDA Enforcement.*** The Weinberg Group Inc. Webinar, December 15, 2010.

***BioSimilar.*** RAPS Annual Conference. October 25, 2010, San Jose, CA.

***FDA Enforcement.*** RAPS Annual Conference. October 25, 2010, San Jose, CA.

***FDA Regulation of Combination Products.*** IIR Combination Products Conference, September 23, 2010, Baltimore, MD.

***Regulation of Generic, OTC and Orphan Drugs, and Cosmetics.*** Lecture at SDRAN RAC Review Course. August 11, 2010, San Diego.

***Regulatory, Quality & Clinical Due Diligence: The Oft-Overlooked Keys to Successful Transactions.*** The Weinberg Group Inc. Webinar, June 23, 2010.

***Informed Consent – Pledge, Platitide or Contract?*** DIA Annual Conference, June 16, 2010, Washington, D.C.

***FDA Enforcement.*** Tutorial at DIA Annual Conference. June 13, 2010, Washington, D.C.

***FDA Enforcement.*** ACI FDA Enforcement Conference. May 25, 2010, Philadelphia.

***510(k) Process.*** BayBio Breakfast Meeting. May 11, 2010, Palo Alto, CA.

***Review of Key 2009 Cases.*** FDLI Annual Conference. April 22, 2010, Washington, D.C.

***CAPA Program.*** OCRA, March 10, 2010, Irvine, CA.

***Drug Development in Today's Regulatory Environment.*** NanoTecNexus Webinar. March 4, 2010.

***Ethical Issues for Clinical Trials.*** ACI International Clinical Trials Conference. February 24, 2010, New York.

***FDA Enforcement.*** FXTranslations Webinar. December 9, 2009.

***How to Respond to a 483.*** SoCalBio FDA Audit Preparedness Workshop. SoCalBio. December 3, 2009, Irvine, CA.

***FDA Enforcement.*** Amylin Pharmaceuticals In-House Lecture. December 1, 2009, San Diego.

***Regulatory Pitfalls in Drug Development.*** American Chemical Society, San Diego Chapter. November 18, 2009, San Diego.

***Drug Safety.*** SDRAN/OCRA Drug Development Conference. November 4, 2009, Carlsbad, CA.

***Webinar on Biosimilars.*** RAPS. October 21, 2009.

***FDA Law.*** Keck Graduate Institute. October 7, 2009, Pomona, CA.

***FDA Enforcement.*** MAGI West Coast Clinical Trials Conference. October 6, 2009, San Diego.

***Biosimilars.*** Foley Life Sciences Day. September 30, 2009, San Diego, CA.

***ANDA vs. 505(b)(2): When and Why.*** The Weinberg Group Inc. Webinar, September 30, 2009.

***Regulatory Update.*** Ophthalmic Drug and Delivery Summit. Pharmaceutical Education Associates. September 22, 2009, San Diego.

***FDA Enforcement.*** The Weinberg Group Inc. Webinar. September 9, 2009.

***Course on Generic Drug Approvals.*** In-House at Teva Parenterals. September 1 and 2, 2009, Irvine, CA.

***Regulation of Generic, OTC and Orphan Drugs, and Cosmetics.*** Lecture at SDRAN RAC Review Course. August 12, 2009, San Diego.

***Drug Development.*** Biotech Vendors Services. July 22, 2009, San Diego.

***Informed Consent.*** DIA Annual Meeting. June 25, 2009, San Diego.

***Clinical Trial Registries.*** DIA Annual Meeting. June 24, 2009, San Diego

***How to Respond to a 483.*** SoCalBio FDA Enforcement Workshop. June 12, 2009, Los Angeles.

***Biosimilars.*** Orange County Regulatory Affairs (OCRA)/FDA Annual Educational Conference. June 10, 2009, Irvine, CA.

***Drug Safety.*** Orange County Regulatory Affairs (OCRA)/FDA Annual Educational Conference. June 10, 2009, Irvine, CA.

***Corporate Health Panel.*** Orange County Regulatory Affairs (OCRA)/FDA Annual Educational Conference. June 9, 2009, Irvine, CA.

***Regulatory Developments for Drug Delivery.*** Third Annual Drug Delivery Summit. Arrowhead Conferences. May 14, 2009, San Francisco.

***Drug Safety: Perspectives on Industry's Duties in the Post-Vioxx Age.*** FDLI Annual Conference. April 22, 2009, Washington, D.C.

***The Future of Biosimilars, BioGenerics, Follow-on Biologics – A Rose by any Other Name?*** New York Biotechnology Association Annual Meeting. April 21, 2009, New York.

***Crisis Management for Senior Regulatory Professionals.*** RAPS Horizons Conference. April 2, 2009, San Francisco.

***FDA Enforcement Issues for Clinical Trials.*** ACI International Clinical Trials Conference. February 26, 2009, New York.

***Roadmap to Emerging Regions -- Clinical Trials in Developing Countries.*** ACI International Clinical Trials Conference. February 26, 2009, New York.

***Overview of FDA Issues for In-Vitro Diagnostics.*** Southern California Biotech Assn. February 13, 2009, Pomona, CA.

***Overview of FDA Issues for Cardiovascular Devices.*** Southern California Biotech Assn. January 28, 2009, Laguna Hills, CA.

***The Forgotten Keys to Bio-Pharma Transactions -- Regulatory, Clinical & Quality Challenges in Contracting and Due Diligence.*** Cambridge Healthcare Institute Second Annual Bridging the Business Development /Alliance Management Interface Conference. November 6, 2008, Boston.

***Regulatory Aspects of Ophthalmic Drug Development.*** Pharmaceutical Education Associates Ophthalmic Drug Delivery Conference. September 22-24, 2008, San Diego

***Regulation of Generic, OTC and Orphan Drug and Cosmetics.*** Lecture at SDRAN RAC Review Course. July 3, 2008, San Diego.

***FDA Enforcement.*** Tutorial at Drug Information Association (DIA) Annual Meeting. June 21, 2008, Boston.

***Panel Discussion on Generic Biologics.*** Licensing Executives Society, San Diego Chapter. May 20, 2008, San Diego.

***IRB Liability.*** Association of Clinical Research Professionals (ACRP) Annual Meeting. April 28, 2008, Boston

***Clinical Trial Registries – Panacea or Pablum?*** Association of Clinical Research Professionals (ACRP) Annual Meeting. April 27, 2008, Boston

***Clearing the US and EU Regulatory Path to Product Approval.*** Swedish American Chamber of Commerce. Swedish-American Entrepreneurial Days. April 9, 2008, San Diego.

***Crisis Management for Regulatory Professionals.*** Regulatory Affairs Professionals Society (RAPS) Horizons Conference. March 28, 2008, San Francisco.

***FDA Regulatory Considerations for the Biomedical Start-Up.*** Israeli Life Sciences Fellows Program, Merage Foundation. February 27, 2008, Irvine, CA.

***The Food & Drug Administration Amendments Act of 2007 – Understanding the Drug Provisions.*** San Diego Regulatory Affairs Network. February 26, 2008, San Diego.

***Keynote Presentation on FDA Regulatory Developments.*** Pharmaceutical Education Associates' 2<sup>nd</sup> Annual Skin Summit Conference, February 20, 2008, Philadelphia.

***Regulatory Considerations for Medical Device Firms.*** NIH-CAP Program. LARTA, February 12 and 13, 2008, via webinar.

***Panel Discussion on Biosimilars.*** BIOCOM Life Sciences Venture Network, February 6, 2008, San Diego.

***International Drug Development.*** Pharmaceutical Education Associates Pipeline to Product Conference, November 30, 2007, Alexandria, VA.

***Regulatory Pitfalls in Drug Development.*** Pharmaceutical Education Associates Pipeline to Product Conference, November 30, 2007, Alexandria, VA.

***Compliance in Clinical Research.*** Eighth Annual Pharmaceutical Regulatory Compliance Congress and Best Practices Forum, November 8, 2007, Washington, D.C.

***FDA Regulatory Strategies for Fast Growing Companies.*** LARTA NIH-CAP Commercialization Training Workshop, October 17, 2007, Marina Del Rey, CA.



***Keynote Presentation on FDA Regulatory Developments.*** Pharmaceutical Education Associates Annual Nasal Drug Delivery Conference, October 4, 2007, Philadelphia.

***Legislative Initiatives.*** RAPS Annual Conference, September 24, 2007, Boston.

***Combination Products – Perspectives on FDA Regulation.*** BVS Orange County Biomedical Day, September 19, 2007, Costa Mesa, CA.

***FDA Regulatory Developments.*** Keynote Presentation at Pharmaceutical Education Associates Annual Ophthalmic Drug Delivery Conference, September 10, 2007, San Diego.

***Regulation of ANDAs, OTC Drugs, Orphan Drug, and Cosmetics.*** Lecture at SDRAN RAC Review Course., August 14, 2007, San Diego.

***Clinical Trial Registries.*** Presented at the University of Southern California Regulatory Science Masters Program, July 27, 2007, Los Angeles.

***Clinical Trial Registries.*** Presented at the American Conference Institute Managing Legal Risks in Clinical Trials Conference, July 16, 2007, San Francisco.

***FDA Enforcement.*** Tutorial at the DIA Annual Meeting, June 17, 2007, Atlanta.

***Keynote Presentation on FDA Regulatory Developments.*** Presented at the Pharmaceutical Education Associates Annual Drug Delivery Conference, June 6, 2007, San Diego.

***The Impact of the Democratic Congress on the Biotech Industry.*** Moderated Panel at the BayBio Annual Meeting, April 26, 2007, San Francisco.

***Guilty Until Proven Innocent: A Look at IRB Liability.*** ACRP Annual Conference, April 23, 2007, Seattle.

***Panel Discussion on Generic Biologics.*** FDLI Annual Conference, April 12, 2007, Bethesda, MD.

***Lifecycle Management for Pharmaceutical Companies: A Generic Perspective.*** Presented at the RAPS Horizons Conference, March 29, 2007, San Francisco.

***Non-Patent Market Exclusivity for Pharmaceuticals Under the Drug Price Competition and Patent Term Restoration Act of 1984 ("Waxman-Hatch").*** San Diego County Bar Association, IP Section, March 19, 2007, San Diego.

***FDA Regulatory Considerations for the Biomedical Start-Up.*** Israeli Life Sciences Fellows Program, Merage Foundation, February 21, 2007, Irvine, CA.

***Alternative Approaches to Drug/Biologics Approvals.*** SDRAN, November 28, 2007, San Diego.

***Informed Consent: Promise, Pledge, Platitude or Contract?*** RAPS Annual Conference, October 18, 2006, Baltimore.

***State Regulation of Clinical Trials.*** 5<sup>th</sup> National Conference on Managing Legal Risks in Structuring & Conducting Clinical Trials, American Conference Institute., September 27-29, 2006, Boston.

***Regulation of ANDAs, Orphan Drugs, OTCs & Cosmetics.*** Lecture at SDRAN RAC Review Course, August 23, 2006, San Diego.

***Key Considerations in Developing Clinical Protocols for U.S. and EU Approval.*** IVT Medical Device Conference, August 15-17, 2006, San Francisco.

***Medical Device Advertising.*** IVT Medical Device Conference, August 15-17, 2006, San Francisco.

***Strategies in Designing Clinicals for Fixed-Combination Drugs.*** DIA Annual Meeting, June 19, 2006, Philadelphia.

***FDA Enforcement.*** Tutorial at DIA Annual Meeting, June 18, 2006, Philadelphia.

***Problems Faced by Device Companies in Navigating FDA Promotional Issues.*** Panel Discussion at the Wilson Sonsini Goodrich & Rosati Medical Device Conference, June 15, 2006, San Jose, CA.

***Clinical Trial Registries: Balm or Bane?*** OCRA/FDA Annual Educational Conference., May 23, 2006, Irvine, CA.

***Product Recalls: A Panel Discussion.*** 3<sup>rd</sup> Annual Medical Device Quality Congress. Management Roundtable and FDANews, May 3, 2006, San Diego.

***Using Clinical Studies to Support Claims for 510(k) Devices.*** RAPS Advertising, Promotion and Labeling Conference, May 2, 2006, Denver.

***The Future of Compliance Governance.*** FDLI Annual Conference, April 7, 2006, Washington, D.C.

***Crisis Management for the Senior RA Professional.*** RAPS Horizons Conference, March 30, 2006, San Diego.

***FDA Regulatory Considerations in Launching Products.*** Women In Technology International (WITI) San Diego Conference, February 14, 2006, San Diego.

***FDA Regulatory Considerations for the Biomedical Start-Up.*** NIH-CAP Workshop, LARTA, October 7, 2005, Newport Beach, CA.

***The “De Novo” 510(k) Process and the Reclassification of Class III Devices.*** 510(k) Workshop, Medical Device Manufacturers Association, October 1, 2005, Boston.

***Specific Payments of Other Sorts -- Understanding SPOOS.*** Clinical Trials – Controlling Costs Conference, Institute for International Research (IIR), September 28, 2005, Philadelphia.

***Regulatory Considerations in Combination Product Development.*** Panel Discussion at the Drug Delivery & Technology Conference, Strategic Research Institute, September 27, 2005, New Brunswick, NJ.

***Specific Payments of Other Sorts – Understanding SPOOS and Clinical Research.*** DIA Annual Meeting, June 29, 2005, Washington, DC.

***FDA Enforcement – What You Need To Know To Avoid – Or Respond.*** Tutorial at DIA Annual Meeting, June 26, 2005, Washington, DC.

***Warning Letters.*** Session Moderator at OCRA/FDA Joint Educational Conference, June 15-16, 2005, Irvine, CA.

***Abbreviated New Drug Applications.*** FDLI Introduction to Biotechnology Conference, June 14-15, 2005, San Francisco, CA.

***California Stem Cell Research And Cures Act.*** RAPS West Coast Conference & Exhibition, March 22-24, 2005, San Francisco, CA.

***Challenges for FDA-Regulated Companies in Addressing Current Corporate Responsibility Trends.*** Workshop on Corporate Responsibility Issues for Regulatory Affairs Professionals. RAPS West Coast Conference & Exhibition, March 22-24, 2005, San Francisco, CA.

***The “De Novo” 510(k) Process and the Reclassification of Class III Devices.*** 510(k) Workshop, sponsored by the Medical Device Manufacturers Association, March 8, 2005, Costa Mesa, CA.

***Strategies in Designing Clinicals for Fixed-Combination Drugs.*** Combination Drug Development Conference sponsored by Barnett International Conferences, March 7-8, 2005, San Diego, CA.

***Recalls by FDA-Regulated Companies.*** Products Liability for FDA-Regulated Firms sponsored by FDLI, January 26-27, 2005, Washington, DC.

***Sarbanes-Oxley – Implications for Life Sciences Companies.*** SDRAN, January, San Diego, CA.

***Financial and Legal Implications of Risk Management: The Broader Picture.*** Developing a Risk Management Strategy: A Hands-On Workshop sponsored by the Pharmaceutical Education & Research Institute (PERI), November 10-11, 2004, Washington, DC.

***Managing Financial Disclosure in Clinical Trials.*** West Coast Drug Development Forum: Challenges in the Development of Therapeutic Products, DIA, October 25-27, 2004, San Francisco, CA.

***FDA Enforcement and Clinical Research.*** Annual Education Symposium, North Texas Chapter, ACRP, October 16, 2004, Dallas, TX.

***Specific Payments of Other Sorts -- Understanding SPOOS and Clinical Research.*** RAPS Annual Meeting, October 12, 2004, Washington, DC.

***Can We Do Better? Innovation in Clinical Trial Agreements.*** 14th International Contracting & Negotiating Clinical Trials, Strategic Research Institute, September 27-28, 2004, La Jolla, CA.

***Case Study: The Generic Drug Scandal.*** Ethics in Regulatory Affairs Seminar, OCRA, August 30, Irvine, CA.

***Regulation of Generic Drugs, OTCs, Orphan Products and Cosmetics,*** Lecture at SDRAN RAC Review Course, August 18, 2004, San Diego, CA.

***FDA Enforcement – What You Need To Know To Avoid – Or Respond.*** Tutorial at DIA Annual Meeting, June 2004, Washington, DC.

***Legal and Regulatory Concerns in the Sourcing of FDA-Regulated Products, Components & Services.*** Center for Professional Advancement In-House Seminar on Vendor & Supplier Qualification, Siemens, January 2004, Concord, CA.

***FDA Enforcement and Compliance.*** SDRAN, October 2003, San Diego, CA.

***State Regulation of Clinical Research.*** Clinical Track of Annual Meeting of the Society of Quality Assurance, October 2003, Washington, DC.

***FDA Regulation of Advertising and Promotion.*** OCRA, September 2003, Irvine, CA.

***Regulation of Generic Drugs, OTCs, and Orphan Products.*** Lecture at SDRAN RAC Review Course, September 10, 2003, San Diego, CA.

***What Every Clinical Director Must Know About FDA Regulatory Compliance.*** Tutorial at DIA Annual Meeting, June 2003, San Antonio, TX.

***Recent Developments in Generic and OTC Drug Regulation.*** Annual Joint Educational Conference , OCRA/FDA, June 2003, Irvine, CA.

***FDA Legal and Regulatory Strategies for Start-up Companies.*** BioMedTrak Program, Tech Coast Angels, March 2003, La Jolla, CA.

***To CRO or Not to CRO.*** Moderator and organizer of the Workshop on Use of Contract Research Organizations in Biomedical Research at the SDRAN IND Conference, February 2003.

***FDA’s “Combination” Product Policy.*** Scripps-BIO 5<sup>th</sup> Annual Drug Development Conference, February 2003, La Jolla, CA.

***Financial Disclosure in Clinical Research.*** San Diego Chapter, ACRP, January 2003, San Diego, CA.

***MDUFMA – A Review of Key Provisions.*** Program on MDUFMA, OCRA, December 2002, Irvine, CA.

***“Specific Payments of Other Sorts:” Sifting Through the SPOOS.*** Barnett-Parexel Conference on Financial Disclosure, November 2002, Philadelphia, PA.

***Legal Issues in Drug Sampling.*** Audioconference sponsored by FDAnews.com, November 2002.

***Financial Disclosure Issues in Clinical Research.*** FDA Regulatory Law Group Breakfast Briefing, entitled “What You Need to Know Before Beginning Your Clinical Trial,” sponsored by Heller, Ehrman, White & McAuliffe, October 2002, San Diego, CA.

***FDA Regulation of Dietary Supplements***, University of Southern California Masters Program on Regulatory Affairs, September 2002, Los Angeles, CA.

***Current Legal Issues Impacting the Generic Drug Industry***. Course on Biotechnology Law, Practicing Law Institute, September 2002, San Francisco, CA.

***The Collateral Legal Consequences of Violating the Food, Drug, and Cosmetic Act – or Why Crime Doesn't Pay***. Association of Medical Diagnostic Manufacturers (AMDM) IVD Conference, September 2002, Del Mar, CA.

***Legal Strategies in Sourcing of FDA-Regulated Goods and Services – Seeking a Win-Win Relationship with Your Contract Manufacturing Organization***. IBC Conference on “Scale-Up: From Bench to Clinic”, August 2002, San Diego, CA.

***Where FDA Leaves Off, Another Agency Picks Up***. Joint FDA Regulatory Law Group/Environmental Law Group Breakfast Briefing, entitled “Beyond FDA – What Every Biomedical Company Must Know About Regulation by Other Federal and State Agencies,” sponsored by Heller, Ehrman, White & McAuliffe. June 2002, Menlo Park, CA.

***What Every Clinical Director Must Know About FDA Regulatory Compliance***. Tutorial at DIA Annual Meeting, June 2002, Chicago, IL.

***Strategies for Success in Dealing with FDA Advisory Panels***. Joint FDA/OCRA Educational Conference, June 2002, Irvine, CA.

***FDA Regulation of Pre-Approval Marketing and Advertising***. FDA Regulatory Law Group Breakfast Briefing sponsored by Heller, Ehrman, White & McAuliffe. April 2002, San Diego, CA.

***FDA Regulation of the Importing and Exporting of Drugs and Devices***. OCRA Conference on Import/Export, March 2002, Irvine, CA.

***The De Novo Petition Process for Medical Devices***. FDA Regulatory Law Group Breakfast Briefing sponsored Heller, Ehrman, White & McAuliffe, November 2001, San Diego, CA and Seattle, WA.

***Ethics in Clinical Research***. Panel Member, Scripps Institute/BIO Joint Conference on Clinical Research, October 2001, San Diego, CA.

***FDA Legal and Regulatory Aspects of Good Clinical Practice***. Society of Quality Assurance (SQA) Annual Meeting, October 2001, San Diego, CA.

***Regulation of Generic Drugs***. Lecture at SDRAN RAC Review Course, October 2001, San Diego, CA.

***FDA Advisory Committees – A Regulatory Overview***. Program on FDA Advisory Committees, SDRAN, September 2001, San Diego, CA.

***Legal Consequences of Violating the Food, Drug, and Cosmetic Act***. Program on Legal Aspects of Recalls, San Diego Regulatory Affairs Network (SDRAN), July 2001, San Diego, CA.

***Legal and Regulatory Strategies in Sourcing of Products, Components and Services for FDA-Regulated Companies.*** Biotechnology Industry Organization (BIO) Annual Conference, June 2001, San Diego, CA.

***FDA Regulation of Imports.*** Food, Drug and Cosmetic Division of the American Society for Quality (ASQ) Conference on “Business Strategies within The Boundaries of the Law,” March 2001, Anaheim, CA.

***FDA Legal and Regulatory Considerations in Drug Development.*** Institute for International Research (IIR) Conference on Drug Discovery, March 2001, San Diego, CA.

***The Collateral Legal Consequences of Violating the Food, Drug, and Cosmetic Act – or Why Crime Doesn’t Pay.*** OCRA Conference on Risk Management, Recalls and Crisis Management, March 2001, Costa Mesa, CA.

***Indemnification in Clinical Research.*** DIA Good Clinical Practice Conference, February 2001, Tucson, AZ.

***International Harmonization of Regulatory Requirements for Biotechnology Products.*** Drug Information Association (DIA) Conference on Biotechnology, February 2001, Dana Point, CA.

***Indemnification in Clinical Research.*** San Diego Chapter of ACRP, November 2000, San Diego, CA.

***Functional Foods: Claims & Labeling.*** Dietary Supplements Conference, RAPS, November 2000, Pasadena, CA.

***An Overview of Global Harmonization of the Regulation of Pharmaceuticals and Medical Devices.*** “Regulatory 101” Seminar, RAPS, November 2000, Pasadena, CA.

***Understanding How to Source Information on FDA Regulatory Activities.*** Panel member, Discussion Presentation at San Diego State University Masters Program on Regulatory Affairs, September 2000. San Diego.

***The Institutional Review Board (IRB) and the Clinical Investigator – Legal/Regulatory Requirements and Perspectives.*** FDA/Orange County Regulatory Affairs (OCRA) Discussion Group Annual Educational Conference, July 2000, Irvine, CA.

***Investigational Device Exemptions (IDE)s.*** Introduction to Device Law Course, FDLI, January 2000, San Diego, CA.

***Device Registration and Listing.*** Introduction to Device Law Course, FDLI, January 2000, San Diego, CA.

***Overview of FDA Regulation of Medical Devices.*** Introduction to Device Law Course, FDLI, January 2000, San Diego, CA.

***Federal Civil and Criminal Laws – How They Impact Medical Device and Drug Companies and Their Employees.*** Compliance with U.S. Regulatory Requirements: FDA Inspections Seminar, RAPS, January 2000, Santa Monica, CA.

***Indemnification in the Clinical Research Context.*** Fall Seminar of the Charlotte Chapter of the Association of Clinical Research Professionals (ACRP), October 1999, Charlotte, NC.

***Legal and Regulatory Aspects for Purchasers of Drug Components, Drugs, and Devices.*** Course on Vendor & Contract Supplier Qualification, CfPA, October 1999, New Brunswick, NJ.

***Global Harmonization Issues and the Protection of Intellectual Property.*** Salud Americas 99 Conference on Latin America's Health Sector Policies, Regulation, and Investment Climate, Institute for the Americas, October 1999, Philadelphia, PA.

***Challenges to Generic Drug Approvals.*** IBC Generic Drug Conference, September 1999, Washington, DC.

***Violations and Enforcement.*** Introduction to Drug Law Course, FDLI, June 1999, Washington, DC.

***Drug Imports and Exports.*** Introduction to Drug Law Course, FDLI, June 1999, Washington, DC.

***Potential Legal Consequences of Product Recalls.*** FDLI Recalls Conference, March 1999, Washington, DC.

***Boosting Awareness of Generics Quality Compared to Brand Name Drug.*** IBC Generic Drug Conference, November 1998, San Diego, CA.

***Health Care Reform and the Generic Drug Industry.*** IBC Generic Drug Conference, January 1994, Orlando, FL.

***Health Care Reform and the Generic Drug Industry.*** IBC Generic Drug Conference, October 1993, Philadelphia, PA.

***FDA Enforcement: A Perspective from Industry on How to Prepare for and Respond to FDA's Knock on Your Door.*** RAPS Annual Conference, October 1993, Washington, D.C..

***Impact of FDA's Regulatory Activities on the Generic Drug Industry: Market Share Through Approvals vs. Market Share Through Attrition.*** International Business Communications (IBC) Conference on Generic Drugs, Competitive Strategies for Pharmaceutical Companies, October 1992, Philadelphia, PA.

***Determining the Regulatory Status of a Drug.*** Seminar on New Drug Applications, RAPS, January 1990, Washington, DC.

***FDA Regulation of Advertising.*** Center for Professional Advancement Course. December 1989. Palm Beach, Florida.

***Importing Drugs.*** McKenna & Cuneo in-house seminar for Embassy Officials on Importing-FDA Regulated Products, July 1989, Washington, DC.

***Impact of Generic Drug Scandal.*** Regulatory Affairs Professional Society (RAPS) Seminar, July 1989, Washington, DC.

***Exclusivity.*** Advanced Drug Law Course, FDLI, October 1988 and May 1989, Washington, DC.

***Orphan Drug Exclusivity.*** Understanding the Orphan Drug Act Seminar, FDLI, October 1988, Washington, DC.

***The Development of Good Manufacturing Practices Under the Food, Drug, and Cosmetic Act.*** Good Manufacturing Practices in the Drug and Allied Industries course, CfPA, June 1987, St. Louis, MO.