21ST Annual Workshop on Medical Communications
Defining the Future of Medical Communications

Preworkshop Tutorials: March 14, 2010
Workshop: March 15-17, 2010
JW Marriott Camelback Inn Resort & Spa, Scottsdale, AZ, USA

Preworkshop Highlights

Sunday, March 14
• Core Curriculum – Full-day session 8:00 AM-4:30 PM
  Separate registration is required. See page 11.
  This program is specifically designed to meet the needs of individuals new to
  pharmaceutical industry-based medical communications. Those who have
  been in this function for less than 1 year would gain the most from attending.
• Afternoon Tutorials – 1:30-5:00 PM
  Separate registration is required. See page 11.
  – Medical Science Liaisons – Literature Searching/Medical Writing
  – Evidence-based Medicine – Practical Applications of Biostatistics

Workshop Highlights

Monday, March 15
• Opening Session – Health Care Reform Update
• Appropriate data sources: Evidence-based medicine versus alternative sources
• Breakout sessions for field-based medical liaisons (interactions with academia
  and MSL 2009 survey results), contact center (handling of product recalls,
  and emerging technology), and headquarters-based medical communications
  (sharing information between companies, and promotional review)
• Welcoming reception featuring speed networking

Tuesday, March 16
• The future of medical communications
• Leadership and professional development
• Podium and poster pearls
• Breakout sessions on customer insights, compliance, and sales training
• Evening reception featuring professional poster presentations

Wednesday, March 17
• Regulatory Review
• Medical communications experience with Risk Evaluation and
  Mitigation Strategy (REMS)

Who Should Attend
Professionals who work in the following areas:
• Medical communications
• Medical liaisons
• Medical information
• Medical call center environment
• Industry, academia, and government

STEEERING COMMITTEE CHAIRS
Alicia Alexander Cadogan, PharmD
Director, Medical Information
Pfizer Inc

Lynn Bass, PharmD
Senior Regional Medical Liaison
Scientific Affairs, Amgen Inc.

See page 2 for a complete list of Program Committee members.

This program was developed by the DIA Medical Communications SIAC.
CONTINUING EDUCATION

Physician Continuing Medical Education

This activity has been planned and implemented in accordance with the Essential Areas and policies of the Accreditation Council for Continuing Medical Education (ACCME) through the joint sponsorship of Postgraduate Institute for Medicine (PIM) and the Drug Information Association. PIM is accredited by the ACCME to provide continuing medical education for physicians.

Credit Designation
Postgraduate Institute for Medicine designates this educational activity for a maximum of 7.25 AMA PRA Category 1 Credit(s)™. Physicians should only claim credit commensurate with the extent of their participation in the activity. (Conference Tutorials)

Core Curriculum Tutorial: 7.25 AMA PRA Category 1 Credit(s)™; Tutorials #1, 2, 3, and 4: 3.25 AMA PRA Category 1 Credit(s)™ each

Postgraduate Institute for Medicine designates this educational activity for a maximum of 16.25 AMA PRA Category 1 Credit(s)™. Physicians should only claim credit commensurate with the extent of their participation in the activity. (Conference)

Conference: 16.25 AMA PRA Category 1 Credit(s)™

Pharmacist Continuing Education

The Drug Information Association is accredited by the Accreditation Council for Pharmacy Education as a Provider of continuing pharmacy education. Type of Activity: Knowledge

This Program is designated for 11.5 contact hours or 1.15 continuing education units (CEUs). 286-000-10-008-L04-P.

Tutorial #1 is designated for 3.25 contact hours or .325 continuing education units (CEUs). 286-000-10-007-L04-P.

Tutorial #4 is designated for 3.25 contact hours or .325 continuing education units (CEUs). 286-000-10-008-L04-P.

Continuing pharmacy education credits are not available for the following sessions: Plenary Session 1; Breakout Session 2-1; Breakout Session 2-3; Breakout Session 3-1; Speed Networking Reception; Plenary Session 5; Luncheon/Professional Poster Session; Breakout Session 6-2; Plenary Session 7; Resident Poster Session/Reception

Nursing Continuing Education

The Drug Information Association will offer nursing credits for this conference in collaboration with Corexcel.

Corexcel is accredited as a provider of continuing nursing education by the American Nurses Credentialing Center’s Commission on Accreditation.

This conference is designated for a maximum of 15.5 nursing contact hours.

Credit Designation
Core Curriculum Tutorial: 6.75 nursing contact hours; Tutorials #1, 2, 3, and 4: 3.25 nursing contact hours each; Conference: 15.5 nursing contact hours

To receive a statement of credit, participants must attend the program and complete the online credit request process through My Transcript at www.diahome.org. Participants will be able to download a statement of credit upon successful submission of the credit request. Complete details and instructions for accessing My Transcript will be included in the final program.

Disclosure Policy: It is Drug Information Association policy that all faculty participating in continuing education activities must disclose to the program audience (1) any real or apparent conflict(s) of interest related to the content of their presentation and (2) discussions of unlabeled or unapproved uses of drugs or medical devices. Faculty disclosure will be included in the course materials.

Workshop Learning Objectives: At the conclusion of this workshop, participants should be able to:

• Demonstrate core competencies in industry-based drug information practice for both field-based and headquarter-based; verbal and written responses; provision of on-label and off-label information; medical and scientific literature evaluation; regulatory and legal applications; and scientific balance versus fair balance
• Discuss important and recent developments regarding health care reform and regulatory guidances related to medical communications
• Recognize medical communications involvement in job functions such as promotional review, REMS, training, compliance, and referencing
• Assess the current landscape for field-based medical communications, and identify ways to enhance their ability to interact with academicians
• Discuss the do’s and don’ts of dissemination of off-label information
• Identify obstacles and opportunities on the horizon for medical communications, and potential ways to benefit from both

PROGRAM COMMITTEE

Maureen Baldwin, RN, MSN
Associate Director, US Contact Center, Pfizer Inc

Christopher Dadas, PharmD
Director, Medical Affairs, Allergan

Sara Doshi, PharmD
Consultant, Global Medical Information, Eli Lilly and Company

Lesley Fierro, PharmD, MS
Associate Vice President, Medical Information Services, sanofi-aventis

Stacey Fung, PharmD
Senior Manager, Medical Communications, Genentech, Inc.

Leena Jindia, MS, PharmD
Director, Medical Information, Tibotec Therapeutics, Division of Centocor Ortho Biotech Services, LLC

Monica Kwarcinski, PharmD
Senior Director, Medical Services, Purdue Pharma LP

Timothy E. Poe, PharmD
Director, Product Information and Patient Services, GSK Response Center, GlaxoSmithKline

Jennifer L. Riggins, PharmD
Director, Global Medical Customer Solutions, Eli Lilly and Company

Rebecca A. Vermeulen, RPh
Senior Director, Global Medical Customer Solutions, Eli Lilly and Company

Ramineh Zoka, PharmD, MS
Senior Director, Clinical Science Liaison, Centocor Ortho Biotech Services, LLC

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org. Participants will be able to download a statement of credit upon successful submission of the credit request. Complete details and instructions for accessing
8:00-8:30 AM  WELCOME AND INTRODUCTIONS

CHAIRPERSON
Stacey Fung, PharmD
Senior Manager, Medical Communications, Genentech, Inc

Following opening remarks, the Core Curriculum faculty will introduce themselves to the attendees and provide descriptions of their career paths leading to their current roles in Medical Communications. The faculty will then describe their current responsibilities, allowing the audience to begin to see similarities and differences in the practice of Medical Communications across the industry.

FACULTY

Tanya Knight-Klimas, PharmD, Medical Information Manager, MedCommunications
William Lai, PharmD, MBA, Senior Manager, Medical Affairs, Baxter Healthcare Corporation
Carol L. Mitchell, MD, Associate Global Information Consultant, Global Medical Information, Eli Lilly and Company
Nima Patel, PharmD, Cardiovascular Medical Information Senior Manager, Bristol-Myers Squibb
Julia Petses, PharmD, Director, Oncology/Urology Medical Information Services, sanofi-aventis U.S.
Jim R. Wilkinson, PhD, Director, Scientific Affairs, Amgen Inc.
Danielle Ziernicki, PharmD, Director, Global Regulatory Affairs Strategic Policy and Support, Johnson & Johnson Pharmaceuticals Group

8:30-10:00 AM  CORE CURRICULUM SESSION 1

8:30-9:30 AM  Session 1A: Regulatory Environment and Medical Communications Practices

FACULTY
Danielle Ziericki, PharmD

This session introduces the current regulatory environment that influences the activities of a Medical Communications professional. An overview of the pharmaceutical industry and DDMAC regulations will be reviewed along with the rationale for many of the activities common in Medical Communications departments.

Learning Objectives
Attendees will build on their existing knowledge and information presented in Session 1A regarding the regulatory environment that influences Medical Communications practice. Case presentations will be discussed by the faculty and attendees.

9:30-10:00 AM  Session 1B: Regulatory Considerations to Medical Communications Practices

FACULTY
Danielle Ziericki, PharmD
Stacey Fung, PharmD

10:00-10:15 AM  REFRESHMENT BREAK

10:15-11:00 AM  CORE CURRICULUM SESSION 2

Current Practices in Medical Communications

FACULTY
Jim R. Wilkinson, PhD
William Lai, PharmD, MBA
Julia Petses, PharmD

This session will review the growing roles and responsibilities that Medical Communications professionals have and how the group supports healthcare professionals and patients, as well as the commercial success of the business. Topics will include scientific meeting support, publication planning, medical education, promotional review as well as collaboration with internal partners.

11:00 AM-12:00 PM  CORE CURRICULUM SESSION 3

Tricks of Advanced Literature Searching

FACULTY
Carol L. Mitchell, MD

Literature searching is a vital skill for Medical Communications professionals. This session will review “sleek tricks of the trade” for searchers of medical literature. Each attendee will walk away with a new trick that can be applied as soon as the next question is asked.

12:00-1:00 PM  NETWORKING LUNCHEON
(Confirmed Core Curriculum attendees only)

1:00-4:30 PM  CORE CURRICULUM SESSION 4

Common Responsibilities of Medical Communications Staff

This session will discuss the common responsibilities of Medical Communications staff. Routine activities of receiving, researching, and formulating scientific responses to unsolicited questions will be reviewed. Topics will include contact centers, formulary dossiers, compendia, internal team support, promotional review, and scientific meeting support. Role playing and mock examples will be used to reinforce principles that emphasize the importance of our role to the industry and to the customers we serve. Attendees will be presented with real-life scenarios that represent challenges that are common to our roles; groups will be asked to discuss and share their ideal responses to the situations.

Attendees will be divided into two groups – Session A and Session B. Both groups will run concurrently with attendees rotating to the other session after the refreshment break.

Session A

FACULTY
Julia Petses, PharmD
Danielle Ziericki, PharmD
William Lai, PharmD, MBA
Stacey Fung, PharmD (Core Curriculum Chair)

Session B

FACULTY
Tanya Knight-Klimas, PharmD
Jim R. Wilkinson, PhD
Nima Patel, PharmD
Evidence-Based Medicine: Trials and Tribulations

Chairperson and Faculty

C. Daniel Mullins, PhD
Professor, Pharmaceutical Health Services Research Department
University of Maryland School of Pharmacy

Evidence-based medicine (EBM) reflects the practice of medicine that is informed by the best available evidence at the point of medical care decision making. EBM requires the generation and evaluation of valid and reliable evidence for medical decision making. While clinical practice guidelines provide significant support for EBM, often there is new evidence that has emerged since the publication of practice guidelines. Thus, quality health care delivery and informed decision making hinges on appropriate study design and analysis of emerging evidence. A variety of evidence grading systems exist, many of which suggest that (meta analyses of) clinical trials provide the most reliable framework for assessing evidence of effectiveness; however, there are a variety of alternative sources of evidence that inform medical decision making and EBM.

The American Recovery and Reinvestment Act (ARRA) contained $1.1 billion for comparative effectiveness research aimed at providing patients, clinicians, payers and others with evidence-based information to make informed decisions about health care. Increased focus on CER raises new questions concerning how evidence generation, analysis, and synthesis inter-relate and how the communication across various stakeholders interested in assessing the value of new health care technologies and those who participate in decision making regarding utilization can provide feedback to improve both processes. This workshop provides an overview of EBM, CER, and the ability to provide syner-gistic coordination despite their divergent objectives.

Learning Objectives
At the conclusion of this tutorial, participants should be able to:
- Describe sources and hierarchies of evidence used to inform evidence-based medicine
- Compare and contrast study designs and interpretation of results across clinical trials and observation studies
- Demonstrate how comparative effectiveness research will alter how medical evidence is generated and evaluated
- Discuss the government’s dynamic role in assessing the value of drugs and other health technologies

Pithy Prose and Deft Discovery: Not Just Another Writing and Searching Course for the Medical Communications Professional

Chairperson and Faculty

Carol L. Mitchell, MD
Associate Global Information Consultant, Eli Lilly and Company

Faculty

Evelyn Hermes-Desantis, PharmD, BCPS
Clinical Associate Professor, Department of Pharmacy Practice and Administration, Rutgers University

Get the goo out of your writing and put your finger on the information you’re looking for! Two veteran educators, both passionate about the written word and the art of searching, will share their secrets for creating concise yet complete documents in conjunction with maintaining a command of the literature.

Learning Objectives
At the conclusion of this tutorial, participants should be able to:
- Explain what an informationist is and strive to be one
- Recite the four basic design principles and be more aware of the “look” of documents
- Create appropriate medical letters, focusing on the “how” as well as the “what” when communicating data through techniques such as repetition/parallelism, paragraphing, non-text options, and easy-to-scan headings
- Understand five ways in medical letters to “cut out fat” while covering more ground in less space
- Figure out how to develop a Subject Heading-based architecture for content domain monitoring
- Develop an agile and real-time content awareness model for staying abreast of the medical literature by adopting a discipline of frequent daily searching

Target Audience
Medical Communications, medical call center environment, medical liaisons, and medical information professionals.

Medical Science Liaison Forum 2010

Chairperson

J. Lynn Bass, PharmD
Senior Regional Medical Liaison, Amgen Scientific Affairs

Faculty

Marc J. Scheineson, Esq.
Allston & Bird, LLP

Bryan Vaughn
Managing Partner, TriNet Pharma

Beth A. Price
Executive Vice President, The Medical Affairs Company

The activities of the Medical Science Liaison (MSL) continues to evolve beyond the original thought leader development role. MSLs in 2010 are engaging in a variety of collaborative activities with key opinion leaders, Investigators, academicians, and others while facing more restrictive regulatory guidances. MSLs are a crucial and direct conduit from these academic and clinical communities to the sponsor company.

In this tutorial, we will discuss challenges to the modern day MSL role and provide innovative solutions from leaders in this field, representing a variety of companies.

Learning Objectives
At the conclusion of this tutorial, participants should be able to:
- Discuss the current and evolving status of the MSL professional and identify key topic areas influencing MSLs in the 2010
- Discuss key strategies for hiring and retaining the best MSL team
- Discuss key regulatory and compliance guidelines affecting the MSL role
A Review in Practical Applications of Biostatistics

Chairperson and Faculty
Timothy A. Candy, PharmD, MS, BCPS
Senior Manager, Regulatory Affairs, Advertising/Promotion, Baxter Healthcare Corporation

This course is intended to provide a broad overview of the most common terms, topics, and techniques used in biostatistics. There will not be a focus on the specific equations and formulas employed but rather the practical application of these statistical techniques. The course is intended for any professional who has to interpret medical literature or internal study protocols and reports to determine clinical relevance to a specific subject or area of research. Actual examples from the medical literature will be used to help illustrate these concepts. This course will be very beneficial to people who have had no formal education in the area of biostatistics. However, this session can also serve as an excellent refresher course for those individuals who have had some experience with biostatistics.

Learning Objectives
At the conclusion of this tutorial, participants should be able to:
- Discuss common terms and concepts of biostatistics useful in the evaluation of scientific literature
- Describe various concepts within hypothesis testing, one of the most common statistical analysis techniques employed in clinical study designs
- Demonstrate the importance of determining the estimated power of a study design and its relevance to interpreting clinical study results
- Describe common areas of misinterpreting statistical results

DAY 2 | MONDAY, MARCH 15, 2010

REGISTRATION AND CONTINENTAL BREAKFAST
7:00-8:00 AM

WELCOME AND OPENING REMARKS
8:00-8:30 AM
Alicia Alexander Cadogan, PharmD
Director, Medical Information, Pfizer Inc

PLENARY SESSION 1
8:30-10:00 AM
Health Care Reform: An Update
Chairperson
Lesley Fierro, PharmD, MS
Associate Vice President, Medical Information Services, sanofi-aventis

How will health care reform affect the pharmaceutical industry and in particular medical communications? This session will provide up-to-date information about the status of health care reform legislation and current issues. Our keynote speaker, Michael McCaughan, will provide the latest information on topics of interest such as reimbursement, ePrescribing, electronic medical records, comparative effectiveness, REMS, and other issues related to medical communications.

KEYNOTE PRESENTER
Michael McCaughan
Editor-in-chief, FDC/Windhover Biopharma Group

Learning Objectives
At the conclusion of this session, participants should be able to:
- Understand the latest information related to health care reform legislation and its impact on the pharmaceutical industry
- Discuss the potential impact of health care reform on medical communications groups
- Identify opportunities for office- and field-based groups to expand communications with health care providers and payers

REFRESHMENT BREAK
10:00-10:30 AM

SPECIAL PRESENTATION
10:30 AM-12:00 PM
Fair Game: Resources for Medical Communications
Chairperson
Christopher Dadas, PharmD
Central Regional Director, Allergan Medical Affairs

This session will explore some of the appropriate sources of information and materials that can be shared with health care professionals from a pharmaceutical company. We will focus on what MSLs and Medical Information groups use as tools to help educate or respond to medical requests.

Learning Objectives
At the conclusion of this presentation, participants should be able to:
- Discuss current and evolving resources used by medical communications
- Discuss emerging methods of communication that may impact the pharmaceutical industry
- Review regulatory guidelines that allow for appropriate scientific communication

Speaker
Alan Minsk, JD
Partner and Chair, Food & Drug Practice Team, Arnall Golden Gregory LLP

NETWORKING LUNCHEON/DINNER SIGN UPS
12:00-1:00 PM

unless otherwise disclosed, DIA acknowledges that the statements made by speakers are their own opinion and not necessarily that of the organization they represent, or that of the Drug Information Association.

Speakers and agenda are subject to change without notice.

Recording of any DIA tutorial/workshop information in any type of media, is prohibited without prior written consent from DIA.
BREAKOUT SESSION 2-1
Is There Still Room for Strategy in the Field-based Liaison Role within the Current Regulatory Environment?

CHAIRPERSON
Beth Lowenthal

The role of the Medical Liaison has continued to evolve since its inception more than 40 years ago. Each year, new guidances from the FDA, learnings from OIG corporate integrity agreements (CIAs), and academic center guidelines have forced companies to re-evaluate how they deploy their field-based teams and what activities they can safely and compliantly engage in. This session will explore the changes that have occurred and how companies have responded to these changes to ensure scientific exchange and education is preserved between companies and academic centers. The role of strategy will also be explored as it relates to planning and the communication of the value of field based teams to others within the organization. Finally, the session will explore how these changes have and will continue to affect managed care field-based teams.

Learning Objectives

At the conclusion of this session, participants should be able to:

• Review the traditional roles of the medical liaison and how they have evolved over time
• Discuss new changes with institutions and the federal government that have helped to shape how medical liaisons function in their role today
• Explore how these changes have impacted strategy planning at the management level and the challenges that face managers in communicating the value of their teams
• Discuss how these changes have empowered and shaped the role of the managed care liaison

Guy M. Chisolm, III, PhD
Vice Chair, Lerner Research Institute

Beth Lowenthal, PharmD, MBA

Archie Stone, PhD
Director, Medical Science Liaisons, Merz Pharmaceuticals

Peter Sheehan
Business Development, REPtrax offered by deView Electronics

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BREAKOUT SESSION 2-2
Preparing for a Tidal Wave of Calls – How Preparation Can Turn Turmoil into Triumph

CHAIRPERSON
Pete Guillot
President, CenterFirst Consulting, LLC

Medical Information contact centers provide a key interface among the company, the medical community, and customers. When events occur that cause a sudden spike in interest in the company or its products, calls to the contact center can spike. Preparation for the spike is critical. If the calls are handled poorly and customers wait endlessly for less than fulfilling responses, then the initial problem is exacerbated by the response. However, if the center handles the situation with calm and precision, confidence is restored to customers, employees, and the general public.

Learning Objectives:

At the conclusion of this session, participants should be able to:

• Learn how one company prepared for product launch by working closely with other functional areas
• Hear how working with your outsource partner is as critical as working with other internal partners when contingency planning

The Contact Center as Critical Product Launch Tool

Rupa Shah, PharmD
Associate Director, Medical Communications, Bristol-Myers Squibb

Working with a Outsource Partner on Contingency Planning

Paul Biedenbach, PharmD
Director, Operations, PPD

Julia Petses, PharmD
Director, Medical Information Services, Oncology, sanofi-aventis

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BREAKOUT SESSION 2-3
It Takes Two to Tango

CHAIRPERSON
Sara Doshi, PharmD
Consultant, Medical Information
Eli Lilly and Company

Medical Information departments continue to explore how to improve their global operations with technical solutions and human resources. This is increasingly challenging as Medical Information departments also find themselves in the midst of an ongoing sequence of corporate integrations and product acquisitions. These may include interdepartmental changes, large pharma plus large pharma mergers, large pharma plus small pharma mergers, as well as the in-licensing of compounds from various sized companies. This session will review how a Medical Information department adapts when they incorporate new stakeholders for several of these scenarios and for products of various market sizes and at varying points in their lifecycle. Speakers will discuss strategies for sharing data/information, collaborative approaches to problem solving, and aligning talent synergistically.

Learning Objectives

At the conclusion of this session, participants should be able to:

• Identify key learnings from a combined globalization-integration Medical Information technology project
• List factors to consider and potential approaches when working in in-license alliance partnerships of various corporate and market sizes.
• Gain insights into handling a product launch with an alliance partner (ie, building a joint Medical Information team from the ground up) and how the “division of labor” between partners may change or transition over time.

Elke Blaetz, RPh, MS
Global Medical Information Leader, Roche

Phil Naughten, PharmD
Director, Affiliate and Alliance Coordination, Takeda Pharmaceuticals North America, Inc.

Cynthia Larmore, RN, MSN
Associate Medical Information Consultant, Eli Lilly and Company
**BREAKOUT SESSION 3-1**

**Medical Liaison Survey #5: Reassessment of Medical Liaison Program Demographics and Characteristics, Demonstration of Value and Assessment of Resource Utilization**

**CHAIRPERSON**

**Craig Klinger, RPh (not presented)**
Senior Medical Liaison Consultant, Lilly USA, LLC

Medical liaison programs are well established throughout the pharmaceutical industry. They exist in companies of all sizes, and also include biotech and device companies. As the role of the medical liaison expands, the challenges of the role continue to be numerous and differ across companies.

In four previous medical liaison surveys, information was gathered on a wide variety of topics. These topics included team demographics, medical liaison position responsibilities, training programs for medical liaisons (both established and newly hired), establishing the value of medical liaisons within the organization, nontraditional staffing options, compliance issues, roles in the health outcomes environment, competitive intelligence, career strategies, virtual office tools, and clinical trial support.

This 5th annual survey of medical liaison practices will gather data on topics from previous surveys in an attempt to evaluate trends over the past several years. Updated information on medical liaison program characteristics and demographics will be collected, along with information on how ML value is demonstrated to business partners. New data collection will be focused on resources used by medical liaisons to obtain and communicate scientific information.

This survey will be administered to both medical liaisons and medical liaison managers across the industry. Survey results will be presented.

**Learning Objectives**

At the conclusion of this session, participants should be able to:

- Discuss the trends seen over the last several years within the role of the medical liaison with regards to ML characteristics and demographics
- Explain how medical liaison programs are demonstrating value to business partners
- Identify how medical liaisons are obtaining and communicating scientific information to thought leaders

**Craig Klinger, RPh (not presented)**

**J. Lynn Bass, PharmD**

**Chris Marrone, PharmD**
Senior Outcomes Liaison, Lilly USA

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**BREAKOUT SESSION 3-2**

**Managing Traditional Needs with Emerging Technology**

**CHAIRPERSON**

**Nicole Corder, RPh, MBA**
Director, The Lilly Answers Center, Lilly USA, LLC

To advance to the next generation Contact Center, we must embrace innovative technology solutions that will allow us to meet the needs of our customers through a variety of ways. In this session, we will take a look at how to enhance the customer experience through the integration of traditional and emerging communication channels. To consider a variety of customer types, we must look for ways to conquer unique staffing models without sacrificing our regulatory requirements. We will take a look at the Remote Agent program, along with an in-depth view of how to accomplish a quality environment. The purpose of this session will be to demonstrate how we can accomplish the interdependencies that relate to merging our customer needs with technology, but never sacrificing our quality. After the presentations, the audience will be encouraged to participate in an interactive question and answer session.

**Learning Objectives**

At the conclusion of this session, participants should be able to:

- Describe emerging channels to meet our customer’s needs
- Identify implementation challenges for today’s technology advancements
- Learn how to successfully utilize “at home” agents to create flexibility to your staffing model
- Discuss the challenges to maintain a QC/QA environment without limiting the ability to be creative with today’s technology and staffing models

**Emerging Customer Communication Channels**

**Linda Comp**
Senior Vice President, Client Services, Telerx

**Remote Agent Program**

**Bronwyn Binaaxas, RPh**
Contact Center Lead Pharmacist, Rocky Mountain Poison and Drug Center

**Maintaining QC/QA Environment**

**Katherine Cahill Moore, RPh**
Manager, The Lilly Answers Center, Lilly USA, LLC

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**BREAKOUT SESSION 3-3**

**Promotional Review: The Role of Medical Communications**

**CHAIRPERSONS**

**Stacey Fung, PharmD**
Senior Manager, Medical Communications, Genentech, Inc.

**Lois Jessen, MS, PharmD**
Director, US Pharmaceuticals Law and Promotion Compliance, Bristol-Myers Squibb

Promotional review is a critical component to successful marketing of products. Medical Communications can play a key role in ensuring scientific accuracy and clinical relevancy of the content. The promotional review process will be described along with findings from a benchmarking survey of Medical Communications roles in promotional review. Key considerations for conducting an effective review will be discussed. While Medical Communications focuses on scientific evidence, the importance of collaboration in developing effective and compliant material will be discussed. Lastly, an interactive discussion with participants’ input on samples of promotional pieces will take place.

**Learning Objectives**

At the conclusion of this session, participants should be able to:

- Recognize the key areas of focus for Medical Communications review of promotional material
- Describe the various models for promotional review committees and how best to work with the team

**Findings of a Benchmarking Survey on the Role of Medical Communications for Promotional Review**

**Ellen Yang, PharmD**
Medical Communications Fellow, Genentech, Inc.

**Alyson Sous, PharmD**
Postdoctoral Fellow, Bayer HealthCare Pharmaceuticals/ Rutgers University

**Overview of Best Practices for Review of Promotional Materials**

**Lois Jessen, MS, PharmD**
Director, US Pharmaceuticals Law and Promotion Compliance, Bristol-Myers Squibb

Interactive Discussion and Review of Sample Promotional Materials

**Stacey Fung, PharmD**
Senior Manager, Medical Communications, Genentech, Inc.
What Modes of Communication Are Wanted and Needed by Our Customers Today?
Duncan Arbour
Senior Consultant
Blue Latitude

How Can Your Organization Make Needed Changes?
Karla Anderson
Life Sciences Commercial Operations
Price Waterhouse Coopers

Real Experience Implementing a New Platform in Partnership with Sermo
Poonam Bordoloi, PharmD
Senior Manager, Internal Medicine
Medical Information Services
sanofi-aventis

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Senior Manager, Internal Medicine
Medical Information Services
sanofi-aventis

8:00-10:00 AM REFRESHMENT BREAK

10:30-11:30 AM PLENARY SESSION 5
Podium Pearls
CHAIRPERSON
Dominick Albano, PharmD, MBA
Assistant Vice President, Medical Information
Pfizer Inc

This session will offer a unique opportunity for any Medical Communications practitioner (eg, information specialist, medical liaison, manager) to share their successes, challenges, and “pearls of wisdom” on various Medical Communications topics through podium presentations.

11:30 AM–1:00 PM LUNCHEON/PROFESSIONAL POSTER SESSION
Take some time to network with your peers while you view posters prepared by Medical Communications professionals. This is a great opportunity to discuss topics of interest with colleagues and view some of the interesting work being done in Medical Communications.
BREAKOUT SESSION 6-1
Customer Insights: Understanding and Creating Value for Our Customers

CHAIRPERSON
Leena Jindia, MS, PharmD
Director, Medical Information, Tibotec Therapeutics, Division of Centocor Ortho Biotech Services LLC

Consumers and healthcare professionals actively seek information from Medical Communications departments based on their unmet medical needs. We will discuss the results of a customer satisfaction market research study and outline how to implement customer feedback to improve the quality of service. Customer inquiries can be another significant source to gather customer insights. We will discuss the potential strategies for evaluating customer inquiries to generate a deeper understanding of their needs and subsequent application of that understanding to the development of solutions for both the customer and the organization. Next we will highlight the increasing number of Latino consumers and the specific customer service skills needed by contact center representatives to be successful in such interactions.

Learning Objectives
At the conclusion of this session, participants should be able to:

• Understand the evolving needs of our customers in today’s technology era
• How can medical information continue to provide value to its customers
• Describe potential sources of customer insights and its application
• Understand how to analyze voice of customer data to identify insights
• What role does cultural difference play in customer expectations? A focus on the Latino customers and the specific customer service skills needed by contact center representatives to be successful in such interactions.

Evaluating the Quality of Services through a Medical Information Market Research Study
Leena Jindia, MS, PharmD

Mining for Insights: Voice of Customer Analysis
Erica Heverin, PharmD
Associate Director, Medical Communications, Ortho-McNeil Janssen Scientific Affairs, LLC

Cultural Insights: The Latino Perspective
Richard Shapiro
President, ENTREVISTA, a Division of The Center for Client Retention

BREAKOUT SESSION 6-2
Hot Topics in Medical Communications Quality and Compliance

CHAIRPERSON
Monica Kwarcinski, PharmD
Senior Director, Medical Services, Purdue Pharma L.P.

This session will review the results of a survey conducted to identify current quality and compliance practices within pharmaceutical industry Medical Communications departments. An overview of what to expect when an external audit is conducted will also be presented as well as useful information regarding how to mitigate risk in the Medical Communications arena. Lastly, a panel of seasoned colleagues will provide factors to consider when developing and maintaining quality and compliance practices within Medical Communications. After the presentations the audience will be encouraged to participate in an interactive question and answer session.

Learning Objectives
At the conclusion of this session, participants should be able to:

• Describe current quality and compliance practices within pharmaceutical industry Medical Communication departments
• Describe what to expect during an external audit
• Learn how to mitigate risk in Medical Communications
• Discuss factors to consider in developing and maintaining quality and compliance practices within Medical Communications departments

Medical Communications Quality and Compliance Survey Results
Monica Kwarcinski, PharmD

Risk Mitigation in Medical Communications
Mark A. DeWyngaert, PhD, MBA
Managing Director, Huron Consulting Group

PANELISTS
Joyce Martin, PharmD
Senior Manager, Quality Assurance, Compliance, and Training, Medical Communications, Genentech, Inc.

Joseph Tuazon, PharmD
Director/Team Leader, Pfizer Medical Information

BREAKOUT SESSION 6-3
Engaging Medical Communications as a Strategic Partner with Sales Training: Three Models for Success

CHAIRPERSON
Mary Sendi, PharmD
Senior Director, Medical Information, Pfizer Inc

This session was developed for headquarter-based Medical Communications professionals and will address several success models where Medical Communications is partnering in excellence with sales training. This session is designed to describe the concept of medical information leadership beyond responding to medical inquiries and opportunities for headquarter-based Medical Communications professionals to partner with sales training in the learning continuum from development to delivery of field-based product information materials. The session will describe current initiatives within the pharmaceutical industry addressing this topic, address potential future programs and implementation concerns, and propose next steps to build a case for change.

Learning Objectives
At the conclusion of this session, participants should be able to:

• Identify the business outcomes and value of building a strategic partnership between sales training and Medical Communications to deliver field-based product information materials
• Describe how one Medical Communications department has partnered with sales training for the development and review process of learning systems and/or backgrounds
• Describe how one Medical Communications department has partnered with sales training to evaluate and implement innovative programs aimed at delivering high-quality and cost-effective training

Steve Wells
Senior Director, Commercial Learning and Development, Pfizer Inc

Kiumars Q. Vadiei, PhD, RPh, FCP
Senior Director, Global Medical Communications, Clinical Development and Medical Affairs, Shire Pharmaceuticals, Inc.

Seema G. Patel, PharmD
Associate Director, Medical Communications, Ortho-McNeil Janssen Scientific Affairs, LLC
Leading with Passion; Leading through Values
CHAIRPERSON
Alicia Alexander Cadogan, PharmD

A great leader once said, “People don’t care how much you know until they know how much you care.” How is it that some leaders are able to inspire and others cannot? How do leaders recognize dedication and commitment in their employees, and how does that affect working relationships, performance, development, and retention? Whether you are a leader or an individual contributor, your attitude and professional value system have a significant impact on how your team functions and how individuals achieve. In this session, we will learn more about the principles that guide the concept of leading by example, and why this is a critical success factor. We will then hear examples of how the principles of success have shaped or influenced the careers of both leaders and individual contributors in Medical Communications. Lastly, we will share simple steps that each person can perform to help them lead by example every day.

Learning Objectives
Upon completion of this session, participants will be able to do the following:

• Identify the opportunities that each person has to be a leader each day
• Find the passion in their work, and communicate that passion to those with whom they interact
• Focus on values in leadership (integrity, courage, and empathy)
• Relate to the examples shared and apply these examples to their own circumstances

4:30-6:30 PM RECEPTION/RESIDENT POSTER SESSION
CHAIRPERSON
Lesley Fierro, PharmD, MS

Take some time to network with your peers while you view posters prepared by Medical Communications residents and fellows. This is a great opportunity to discuss topics of interest with colleagues and view some of the interesting work being done by up-and-coming industry residents and fellows.

This session will provide the latest information on activities of office-based and field-based Medical Communications personnel that may have important regulatory implications.

Topics will include provision of medical product information in a variety of situations and audiences as well as updates on recent activities by DDMAC and the Office of Inspector General (OIG). Areas of interest around dissemination of scientific literature, medical education, and other recent activities of the FDA pertinent to Medical Communications will be discussed.

This will be an interactive session with opportunity for discussion and questions from the audience.

Learning Objectives
At the conclusion of this session, participants should be able to:

• Discuss recent regulatory actions and the impact on Medical Communications practices
• Discuss how actions of other government agencies affect the pharmaceutical industry and Medical Communications practices

10:00-10:15 AM REFRESHMENT BREAK

10:15-11:45 AM PLENARY SESSION 9
Medical Communications Experience with Risk Evaluation and Mitigation Strategies (REMS)
CHAIRPERSON
Ramineh Zoka, PharmD, MS
Senior Director, Clinical Science Liaison, Medical Affairs, Centocor Ortho Biotech Services, LLC

Risk Evaluation and Mitigation Strategies (REMS) are becoming part of the fabric of pharmaceutical companies to ensure the safe use of their marketed medications. Medical Communications staff plays a critical role in shaping and implementation of this process. During this session, an overview of the key components of REMS will be provided, and examples of how Medical Communications staff have contributed to the successful planning and implementation of REMS programs will be discussed.

11:45 AM-12:00 PM CLOSING REMARKS
J. Lynn Bass, PharmD
Senior Regional Medical Liaison, Scientific Affairs, Amgen Inc.
REGISTRATION FORM
Register online or fax this page to +1.215.442.6199

DIA is a financially independent nonprofit, global, multidisciplinary association that provides a neutral forum for sharing information that optimizes the development and lifecycle management of biopharmaceutical and related products.

21st Annual Workshop on Medical Communications
Event #10006  •  Tutorials: March 14  •  Workshop: March 15-17, 2010
JW Marriott Camelback Inn Resort & Spa, Scottsdale, AZ, USA

Contact Information
Event: JoAnn Boileau, Program Manager, Phone +1.215.442.6175, Fax +1.215.442.6199, or email JoAnn.Boileau@diahome.org
Tabletop Exhibits: Shannon Lewis, Exhibits Manager, Phone +1.215.442.6149, Fax +1.215.442.6199, or email Shannon.Lewis@diahome.org

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TUTORIALS: SUNDAY, MARCH 14  1:30-5:00 pm
#1 Evidence-based Medicine  •  #2 Pithy Prose and Deft Discovery  •  #3 Medical Science Liaison Forum  •  #4 Practical Applications of Biostatistics
Registration is limited to ONE Tutorial. Tutorial registrants cannot register for the Core Curriculum.

CORE CURRICULUM: SUNDAY, MARCH 14  8:00 am-4:30 pm
This is a full-day session and this registration section is limited to the Core Curriculum only. You must complete this section if you wish to attend the Sunday Core Curriculum which is limited to 100 attendees. Your acceptance will be confirmed in writing.

Individuals new to pharmaceutical industry-based medical communications (less than 1 year in this function) would gain the most from attending the Core Curriculum.
I wish to attend the Core Curriculum for an additional fee of US $295  □
I have not attended the Core Curriculum before.
My department is:  □  Contact Center  □  Medical Liaison  □  Medical Information  □  Other

Please check the applicable category:
☐  Academia  ☐  Government  ☐  Industry  ☐  CSO  ☐  Student

TO RECEIVE A TABLETOP EXHIBIT APPLICATION, PLEASE CHECK  □

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Card # ____________________________

Name (printed) ________________________________________________________________

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☐ CHECK: drawn on a US bank payable to and mailed along with this form to: Drug Information Association Inc. P.O. Box 95000-1240, Philadelphia, PA 19195-1240, USA. Please include a copy of this registration form to facilitate identification of attendee.

☐ BANK TRANSFER: When DIA completes your registration, an email will be sent to the address on the registration form with instructions on how to complete the Bank Transfer. Payment should be made in US dollars. Your name and company, as well as the Event ID, # must be included on the transfer document to ensure payment to your account.

TRAVEL AND HOTEL: The most convenient airport is Phoenix Sky Harbor International Airport and attendees should make airline reservations as early as possible to ensure availability. The JW Marriott Camelback Inn Resort & Spa is holding a block of rooms at the reduced rate below until February 20, 2010, for the DIA event attendees. Room availability at this rate is guaranteed only until this date or until the block is filled.

Single $199+  •  Double $199+

Please contact the JW Marriott Camelback Inn Resort & Spa by telephone at +1.800.24CAMEL (22635) and mention the DIA event. The resort is located at 5402 East Lincoln Drive, Scottsdale, AZ 85253, USA.

CANCELLATION POLICY: On or before MARCH 8, 2010
Administrative fee that will be withheld from refund amount:
Member or Nonmember = $200
Government or Academia or Nonprofit (Member or Nonmember) = $100
Tutorial (if applicable) = $50

Cancellations must be in writing and be received by the cancellation date above. Registrants who do not cancel by that date and do not attend will be responsible for the full registration fee paid. Registrants are responsible for cancelling their own hotel and airline reservations. You may transfer your registration to a colleague at any time but membership is not transferable. Please notify DIA of any such substitutions as soon as possible. Substitute registrants will be responsible for nonmember fee, if applicable.

DIA reserves the right to alter the venue, if necessary. If an event is cancelled, DIA is not responsible for any airfare, hotel or other costs incurred by registrants.

Participants with Disabilities: DIA event facilities and overnight accommodations are accessible to persons with disabilities. Services will be made available to sensory-impaired persons attending the event if requested at least 15 days prior to event. Contact the DIA office to indicate your needs.

GROUP DISCOUNTS*: Register 3 individuals from the same company and receive complimentary registration for a 4th! All 4 individuals must register and prepay at the same time - no exceptions. DIA will apply the value of the lowest applicable fee to this complimentary registration; it does NOT include fees for optional events or DIA membership. You may substitute group participants of the same membership status at any time; however, administrative fees may be incurred. Group registration is not available online and does not apply to the already-discounted fees for government or charitable nonprofit/academia. To take advantage of this offer, please make a copy of this registration form for EACH of the four registrants from your company. Include the names of all four group registrants on each of the forms and return them together to DIA.

☐ Please indicate that this form is part of a group registration by checking this box and list below the names of the other three registrants from your company.

1.  __________________________________________
2.  __________________________________________
3.  __________________________________________

Please check the applicable category:
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(Call for registration information)

Last Name  ____________________________
First Name  ____________________________
MI  ☐  Dr. ☐ Mr. ☐ Ms.

Degrees

Job Title

Company

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Mail Stop

City  ____________________________  State  ____________________________  Zip/Postal  ____________________________  Country

email  __________________________________________
Required for confirmation

Phone Number  ____________________________  Fax Number  ____________________________  Required for confirmation


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