BUILDING A WORLD-CLASS DVERTISING AND PROMOTION

AN INTERACTIVE WORKSHOP PRESENTED BY PHILLYCOOKE CONSULTING AND FDANEWS

YOUR EXPERT SPEAKER:



DALE A. COOKE Owner, PhillyCooke Consulting

Mr. Cooke's practice specializes in helping FDA-regulated companies develop compliant promotional tactics and improve the promotional review. He is the author of Effective Review and Approval of Digital Promotional Tactics and is currently at work on a book about compliant social media usage for prescription product manufacturers.

"Dale is easy to listen to. The material covered is comprehensive. The sessions covered what I believe is beneficial to those responsible for ad/promo review"

> — Tim Williams, VP of Regulatory Affairs, CR Bard

APRIL 27-28, 2016

CHICAGO MARRIOTT OAK BROOK OAK BROOK, IL

2 Days, 200 Tips For Improving Your Advertising and Promotion Review Program

FDA marketing scrutiny no longer is limited to magazine and TV ads. Now the agency is poking around, checking signage in tradeshow booths... checking in on Twitter and Facebook...and listening to the physicians and other healthcare professionals you've paid to speak or train.

Come to Chicago in April for two days of intense learning. You'll arrive back home with a bag full of tricks and tips to keep all your marketing efforts squeaky-clean.

- Understanding Pre-Approval Communications Don't get on the FDA's or SEC's radar screens before your product is even approved. Learn how to properly disclose information and remain in compliance.
- How To Maximize Disease Awareness Communications Take away valuable tips and tricks for using disease awareness communications pre- and post-approval.
- Hurray! You're Approved Building the most aggressive but compliant campaign from first day of approval to commercial launch.
- Assuring Your Promotions Meet FDA Off-Label Standards Successfully navigating 4 major traps that can earn you a warning letter fast.
- Itching To Do More With Social Media? Discover how to get your message out there ... without crossing the line.







FDANEWS BUILDING A WORLD-CLASS ADVE

DAY ONE

8:00 a.m. - 9:00 a.m.

Registration and Continental Breakfast

9:00 a.m. - 9:45 a.m.

Pre-approval Communications

 How to meet your SEC requirements for disclosing information while not running afoul of FDA pre-approval promotion prohibitions

9:45 a.m. - 10:30 a.m.

Disease Awareness Communications

- A review of FDA's help-seeking guidance
- Keys for using disease awareness communications prior to approval. Essential information for continuing communications efforts post-approval compliantly.

10:30 a.m. - 10:45 a.m.

Break

10:45 a.m. - 11:15 a.m.

From Day of Approval through Commercial Launch

- Understanding the timeline and key dates for communications
- Minimizing the pain, while maximizing the impact of initial promotional communications

11:15 a.m. 12:00 p.m.

Essential Advertising & Promotion Regulations

 A review of all of the requirements for product promotion, including product name usage, fair balance, directions for use

12:00 p.m. - 1:00 p.m.

Lunch

1:00 p.m. - 1:45 p.m.

Format-Specific Promotional Requirements

- DTC television promotion
- Brief summary requirements for print promotion

1:45 p.m. - 2:30 p.m.

Substantial Evidence & Other Standards

 A review of the substantial evidence standard, what fails to meet that standard, and when other standards apply

2:30 p.m. - 2:45 p.m.

Break

2:45 p.m. - 4:00 p.m.

Off-Label Information

- Avoiding off-label promotion
- Scientific exchange exemption
- Responding to unsolicited requests
- Distributing off-label reprints

4:00 p.m. - 4:30 p.m.

The Promotional Review Process

- An overview of a the standard promotional review process
- Essential traits for any effective process
- Key decisions in establishing or improving performance
- Effective use of metrics for evaluating performance

4:30 p.m.

Session Wrap-Up, End of Day One

DAY TWO

8:30 a.m. - 9:00 a.m.

Continental Breakfast

9:00 a.m. - 9:45 a.m.

Integrating Digital Promotion

 Key considerations for evaluating the compliance of digital promotional tactics and their integration into the overall promotional mix

9:45 a.m. - 10:15 a.m.

Social Media Part 1

 Overview of the platforms, issues, and a review of the status of FDA guidance

10:15 a.m. - 10:30 a.m.

Break

10:30 a.m. - 12:30 p.m.

Social Media Guidances

 A review of the three social media guidances released in 2014: 2253 Filing, Presenting Risk Information in Space-Limited Contexts, & Correcting Misinformation on Third-party

12:30 p.m. - 1:30 p.m.

Lunch

1:30 p.m. - 3:15 p.m.

Promotional Review Board Practicum

 Workshop participants will apply the lessons from the earlier part of the workshop to specific product promotions. They will work in teams to evaluate specific promotional tactics, determine what (if any) parts of the promotion are problematic, and how to provide direction to a brand marketer to make the promotions compliant.

3:15 p.m. - 3:30 p.m.

Break

3:30 p.m. - 4:15 p.m.

Continuing Regulatory Intelligence: Staying Abreast of Ad/Promo News

 Ad/Promo is an area of ongoing developments. This session will cover the most prominent sources for keeping up with these developments.

4:15 p.m. - 4:30 p.m.

Wrap-up and Adjourn Workshop

"Dale was very engaging and informative. I like the interactive portion where we reviewed actual ads."

- Workshop Attendee

ERTISING AND PROMOTION REVIEW PROGRAM

WHO WILL BENEFIT?

- Advertising and marketing managers
- Social media teams
- Promotion review committee members
- Medical affairs
- Continuing education and tradeshow organizers
- Regulatory compliance officers
- PRC coordinators
- Legal counsel
- Compliance
- Executive management
- Outside ad agencies and marketing consultants

"[Dale is] an animated speaker who seems to know something about everything and has no shortage of opinions. This is potentially very dry and dull material. Dale brings out the exciting and humorous aspects especially well. It's an engaging two days."

— Michael Benedetto, **Editorial Group Leader, FCB Health**

"This is normally a dry topic, but with Dale it was anything but dry. Dale gave one of the best, most engaging presentations. Dale is highly knowledgeable and also very entertaining."

— Ellen Derrico, Global Head, **Market Development - Life** Sciences & Healthcare, QlikTech

"As a fellow regulatory professional, Dale is one of the most deeply knowledgeable experts I've heard on this complicated subject. As the pharmaceutical industry is experiencing, digital promotional tactics can trip up even experienced regulatory professionals, but Dale showed how basic principles can be applied to develop compliant promotional materials."

> - Kathleen Koons, Sr Regulatory Affairs Manager, **DJA Global Pharmaceuticals Inc.**

Course Binder Materials:

Full slides from the PowerPoint presentations Reference documents:

FDA Advertising & Procedural Guidances

- Help-Seeking Guidance
- DTC Broadcast Guidance & Q&A
- FDAAA Pre-Dissemination Review Requirements
- Product Name Guidance (All three versions from 1999, 2012, and 2013)
- Presenting Risk Information Guidance
- Social Media Guidances
 - Postmarketing Submissions Requirements
 - Responding to Unsolicited Requests for Off-label Information
 - Presenting Risk Information in Space-limited Contexts Correcting Thirdparty Misinformation
- Distributing Off-Label Reprints

Relevant Sections of Code of Federal Regulations

Form 2253

PhRMA Principles on DTC Advertising

PhRMA Principles on Interactions with Healthcare Professionals

Pre-approval Promotion Checklist

Effective Review & Approval Process Checklist

Keys to Evaluating Promotional Review Systems List of Key Resources for **Continuing Education**

Articles by Dale Cooke

- Developing Compliant Search Engine Marketing Campaigns (Publication Date of September 2014)!
- Industry Standards for Linking Disease Awareness Websites to Product Promotion!
- Patient Testimonial Videos: FDA Actions on Risk Information Presentation!
- Presenting Risk Information on Websites!
- · Where Things Stand on FDA Guidance on Social Media (Publication Date of September 2014)

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□ Yes!			
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HOTEL INFORMATION INFORMATION:

April 27-28, 2016

Chicago Marriott Oak Brook 1401 West 22nd Street Oak Brook, IL 60523

Toll free: (800) 228-9290 Tel: +1 (630) 573-8555

www.marriottoakbrook.com

Room rate: \$159 plus 9% tax Reservation cut-off date: April 5, 2016

TEAM DISCOUNTS:

Significant tuition discounts are available for teams of two or more from the same company. You must register at the same time and provide a single payment to take advantage of the discount. Multi-attendee discounts are available and will be calculated at check out.

2-4 attendees – 10% 5-6 attendees – 15% 7-9 attendees – 20% 10+ attendees – 25%

TUITION:

Tuition includes all conference presentations, conference materials, two breakfasts, two lunches and refreshments.

CANCELLATION AND SUBSTITUTION:

Written cancellations received at least 21 calendar days prior to the start date of the event will receive a refund — less a \$200 administration fee. No cancellations will be accepted — nor refunds issued — within 21 calendar days from the start date of the event. A credit for the amount paid may be transferred to any future FDAnews event. Substitutions may be made at any time. Noshows will be charged the full amount. In the event that FDAnews cancels the event, FDAnews is not responsible for any airfare, hotel, other costs or losses incurred by registrants. Some topics and speakers may be subject to change without notice.