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📅 November 28 – 30, 2018 📍 New York Marriott Marquis, New York, NY

**EARN CLE/ETHICS CREDITS**

23<sup>RD</sup> ANNUAL CONFERENCE ON

# DRUG MED DEVICE LITIGATION

Expert Strategies for Leading Products Liability Litigators & In-House Counsel

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Senior Counsel  
Cardinal Health



**Carolyn M. Hazard**  
Senior Vice President, Associate General Counsel – Litigation  
Endo Pharmaceuticals

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Associate General Counsel  
UCB

**Ragan E. Cheney**  
Sr. Vice President, General Counsel  
Corporate Secretary  
Titan Spine

**Greg A. Dadika**  
Associate General Counsel,  
Litigation  
Becton, Dickinson

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Director, Legal Counsel  
Teva Pharmaceuticals USA

**Gina Gencarelli**  
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Par Pharmaceutical

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Senior Counsel  
Bayer U.S.

**Elizabeth Howard**  
Executive VP and General Counsel  
Arbutus Biopharma

**Wendy Hufford**  
Chief Operating Officer, Legal  
Department & Vice President,  
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& Human Resources  
Boehringer Ingelheim USA

**Lisa LeCointe-Cephas**  
Executive Director, Head of  
Global Investigations  
Merck & Co.

**Amanda T. Perez**  
Assistant General Counsel  
Pfizer Inc.

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Head of U.S. Litigation  
Novartis

**Munjot Sahu**  
Counsel – Litigation and  
Legal Compliance  
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**Hon. Ann D. Montgomery**  
Senior Judge  
U.S. District Court, D. Minn.

**Hon. Arnold L. New**  
Coordinating Judge/Complex  
Litigation Center  
Philadelphia Court of  
Common Pleas

**Hon. Patti B. Saris**  
Chief Judge  
U.S. District Court, D. Mass.  
(Boston, MA)

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Business Information in a Global Context

## What's New for 2018?



Extended Networking Breaks



18 Ground Breaking Sessions with the Introduction of 7 Thought Provoking Discussions on the Most Pressing Products Liability Challenges of 2018



2 Panels of Judges Involved in MDLs and Drug & Device Cases



Insights on the Future of Drug and Medical Device Litigation from over 2 Dozen In-House Counsel

400+

ATTENDEES



70+

SPEAKERS



24+

IN-HOUSE SPEAKERS

18+

SESSIONS



## Special Features for In-House Counsel

Designed with the needs of in-house counsel in mind!

> Highlighted features:

- Executive Boardroom Meeting:** Engage with the most sophisticated in-house minds in the industry during an intensive and exclusive 3-hour brainstorming session!\*
- In-House Think Tank Lunch (by Invite Only)** – Discuss the state of the industry candidly with your peers on how members of the defense bar can coordinate their advocacy efforts for 2019.\*
- \*Limited **complimentary passes** available this year for qualified in-house counsel. See page 15 for details.

> Featured sessions on:

- More than just products liability – the latest on pricing transparency initiatives, regulatory updates and patent litigation impacting the industry
- Diversity: Practical ways for incorporating diversity and inclusion into the trial team and litigation strategy
- Enforcement Initiatives and Priorities straight from the enforcers themselves

## Defense Counsel Only War Room

Back by Popular Demand

Open to defense counsel only, join your peers to participate in this unique, interactive networking session that will set the stage for the topics discussed in-depth throughout the event and provide you with valuable takeaways about what your peers from around the country are seeing from the plaintiffs' bar.

See details on page 6

## Participating Companies Include:

Daiichi Sankyo, Inc.

Endo Pharmaceuticals

Medtronic, Inc.

Bayer U.S.

Purdue Pharma L.P.

UCB, Inc.

Eli Lilly

Titan Spine

Becton, Dickinson and Company

Teva Pharmaceuticals USA, Inc.

Cardinal Health

Arbutus Biopharma

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Merck & Co., Inc.

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and more...

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American Conference Institute

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For over 30 years, C5 Group has provided the opportunities that bring together business leaders, professionals and international experts from around the world to learn, meet, network and make the contacts that create the opportunities.

Our conferences and related products connect the power of people with the power of information, a powerful combination for business growth and success.

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ACI: Drug & Medical Device Litigation

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Senior Counsel  
**Cardinal Health (Columbus, OH)**



**Carolyn M. Hazard**  
Senior Vice President,  
Associate General Counsel – Litigation  
**Endo Pharmaceuticals Inc. (Malvern, PA)**

### Speakers



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









**Hon. Susan D. Wigenton**  
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**U.S. District Court, D.N.J. (Newark, NJ)**



# Agenda At-A-Glance

 **New This Year**

	<b>DAY ONE</b> WEDNESDAY, NOVEMBER 28, 2018	<b>DAY TWO</b> THURSDAY, NOVEMBER 29, 2018	<b>DAY THREE</b> FRIDAY, NOVEMBER 30, 2018
<b>7 am</b>		7:00 Registration and Welcoming Breakfast	7:15 Registration and Continental Breakfast
<b>8 am</b>	8:30 Drug and Med Cross-Exam 101 Registration and Continental Breakfast	8:00 Co-Chairs' Opening Remarks	8:00 Co-Chairs' Opening Remarks
<b>9 am</b>	9:00 Drug and Med Cross-Exam 101	8:15 GC and CLO Roundtable	8:15 A View from the Bench
<b>10 am</b>		9:30  MDLs: Examining Their Original Purpose and Attempted Ways at Improving the MDL System	9:30 Enforcers' Roundtable
<b>11 am</b>		10:15 Morning Coffee Break	10:30 Morning Coffee Break
		10:45 A View from the Bench	10:45  Hiring the Right Regional Counsel
<b>12 pm</b>	12:30 In-House Only Think-Tank Lunch	12:00 Networking Luncheon	11:30  What Does It Mean to Be an Innovative Thinker as an Outside Counsel
<b>1 pm</b>	1:45 Registration for Afternoon Sessions	12:00 Networking Luncheon	12:15 Networking Lunch
		1:00  Diversity and Inclusion	1:15  Lay of the Land with Respect to Pricing Transparency, Patent Litigation, and Regulatory Trends
		1:45 Afternoon Breakout Sessions (CHOOSE A OR B) A. The Future of Personal Jurisdiction B. Gaining Control Over Third-Party Financing of Litigation	
<b>2 pm</b>	2:00  Executive Boardroom Meeting (IN-HOUSE COUNSEL BY INVITATION ONLY)	2:30 Afternoon Networking Break	2:00 Opioid Enforcement and Litigation Landscape
	2:30 Defense Counsel Only War Room		2:45 Conference Ends
<b>3 pm</b>		3:00 Afternoon Breakout Sessions (CHOOSE A OR B) A. How to Move Forward When Faced with a Multi-Plaintiff Trial B. Scope of Discovery	
		3:45 Afternoon Breakout Sessions (CHOOSE A OR B) A. State of the Union on Preemption B. Strategies for Minimizing the Effects of Innovator Liability	
<b>4 pm</b>		4:30 Afternoon Breakout Sessions (CHOOSE A OR B)  A. Training the Future Generations of Life Sciences Attorneys  B. Examining the Intersection of Traditional Principles of Product Liability Laws with Digital Health and 3D Printing	
<b>5 pm</b>	5:00 Pre-Registration and Welcoming Cocktail Reception	5:15 Conference Adjourns to Cocktail Party	
	HOSTED BY: McDermott Will & Emery	HOSTED BY: KING & SPALDING	

# DAY ONE

WEDNESDAY, NOVEMBER 28, 2018

## Workshop:

### Drug and Med Cross-Examination 101

9:00 – 12:00 (Registration and Continental Breakfast at 8:30)

**Sean K. Burke**

*Partner*

**Duane Morris (Washington, DC)**

Because a drug or device trial often hinges on the expert witness testimony, knowing how to conduct an effective cross-examination of an expert is crucial in determining the direction of the litigation.

Designed for those who are new to this type of litigation, this intensive 3-hour class will arm you with strategies on how to effectively use key case milestones, such as Daubert/Frye motions, as well as how to cross-examine a science and a regulatory/FDA expert. Points of discussion will include:

- Conveying easy to understand scientific concepts to the jury
- Exacting jury-friendly concessions from the opposing expert
- Highlighting the expert's misuse of statistical data and methodology
- How to undercut attempts to confuse principles of burden of proof with principles of statistical significance
- Using the expert to establish the expertise of the FDA and pervasiveness of FDA review and oversight
- Successfully showing – through cross – that the regulatory/FDA expert lacks sufficient qualifications in the area of proposed testimony
- Using prior reports and testimony to demonstrate that the expert's opinions are boilerplate and full of biases
- Following the money: Using the total dollar figures the expert has made providing testimony against him/her

## Workshop:

### Defense Counsel Only War Room

2:30 – 5:00 (Registration begins at 1:45)

**Patricia A. Barbieri**

*SVP, General Counsel and Secretary*

**Daiichi Sankyo, Inc. (Parsippany, NJ)**

**Max Heerman**

*Principal Litigation Counsel*

**Medtronic (Washington, DC)**

**Open to defense counsel only** – Join your peers for a state-of-the-industry analysis and candid discussion about the latest and greatest in plaintiffs' tactics. In-house and law firm defense counsel are encouraged to participate in this unique, interactive networking session that will set the stage for the topics discussed in-depth throughout the event and provide you with valuable takeaways about what your peers from around the country are seeing from the plaintiffs' bar. Discussion will include:

- Jurisdictional issues
- Discovery, including cross-border discovery
- Successful motions to dismiss and other dispositive motions
- Litigation tactics in aggregated cases: multiple plaintiff trials
- Defending against junk science and using good science/literature to bolster your defense
- Punitive damages: to bifurcate or not bifurcate?
- In-house counsel: how can your clients better serve you?  
Outside counsel: how can your clients be better partners?
- Analysis of the recent wave of cases in device litigation where courts have been excluding evidence of the 510k clearance process
- A survey of the less known but important decisions impacting the drug and device space

5:00 – 6:00

**Pre-Registration  
and Welcoming  
Cocktail Reception**

HOSTED BY:

**McDermott  
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LIMITED  
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PASSES FOR  
IN-HOUSE COUNSEL

12:30

### In-House Think-Tank Lunch\*

2:00 – 5:00 (Registration begins at 1:45)

### **NEW** In-House Executive Boardroom Meeting\*

Top drug and device in-house minds will come together for this exclusive invite-only boardroom meeting to discuss their most difficult challenges and top priorities as they navigate and together think through the future of the healthcare industry.

*\*Open to In-House Counsel and Senior Executives from Drug, Device and Biotech Companies. All interested parties will be pre-qualified before registering for the program.*



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ACI: Drug & Medical Device Litigation

# MAIN CONFERENCE DAY TWO

## THURSDAY, NOVEMBER 29, 2018

7:00

### Registration and Welcoming Breakfast

HOSTED BY: 

8:00

### Co-Chairs' Opening Remarks

**Kailee Goold**

*Senior Counsel*

**Cardinal Health (Columbus, OH)**

**Carolyn M. Hazard**

*Senior Vice President, Associate General Counsel – Litigation*

**Endo Pharmaceuticals Inc. (Malvern, PA)**

8:15

### GC and CLO Roundtable: What Keeps Them Up at Night

**Ragan E. Cheney**

*Sr. Vice President, General Counsel*

*Corporate Secretary*

**Titan Spine (Mequon, WI)**

**Elizabeth Howard**

*Executive VP and General Counsel*

**Arbutus Biopharma (Warminster, PA)**

**Maureen A. Ruane**

*Vice President & Head of U.S. Litigation*

**Novartis (East Hanover, NJ)**

**Richard W. Silbert**

*Vice President, Chief Legal Strategist*

**Purdue Pharma L.P. (Stamford, CT)**

**Jonathan Wasserman**

*Vice President and Associate General Counsel,*

*Litigation & Government Investigations*

**Bristol-Myers Squibb Company (Lawrenceville, NJ)**

Moderator

**Lori G. Cohen**

*Co-Chair, Global Litigation; Chair, Pharmaceutical, Medical Device &*

*Health Care Litigation Group; Chair, Trial Practice Group*

**Greenberg Traurig, LLP (Atlanta, GA)**

In this exclusive session, attendees will have the unique opportunity to hear directly from leading counsel at pharmaceutical and device companies about their greatest products liability challenges. Gain critical insights into the thinking and mindset of key legal decision makers on such topics as:

- Conducting litigation and compliance audits to determine key areas of risk
  - » Factoring in litigation trends and recent product liability law suits and enforcement activity against life sciences companies
  - » Analyzing areas of risk for your company based on internal audits and forecasting the likelihood of a potential lawsuit or enforcement action
  - » Assessing costs of potential litigation
- Communicating areas of potential risk to company's senior executives
- Weighing options to manage an MDL when settlement is not feasible

- Exploring the latest fee arrangement and alternate payment structures with outside counsel
- Balancing the litigation environment with business demands/realities
- How to best address/respond to media scrutiny and attack as well as other public relations issues

9:30



### MDLs: Their Intended Purpose, What Attempts Have Been Made at Improving the MDL System, and Effective Ways for Wrapping Them Up

**John Galvin**

*Partner*

**Fox Galvin (St. Louis, MO)**

**Malini Moorthy**

*Vice President & Associate General Counsel; Head of Global Litigation*

**Bayer U.S. (Pittsburgh, PA)**

**Lana K. Varney**

*Partner*

**King & Spalding (Austin, TX)**

- What is the intended purpose behind MDLs and has that purpose been obscured by practicalities of courts' needing to manage the case dockets?
- What attempts have been made at improving the MDL system?
- Early judicial analysis of inventories in MDLs: Looking at the issue of meritless cases hiding within the MDLs, can any strategies be suggested to the courts as to how to address it?

#### Getting closure: How to wrap up an MDL?

- » Ideas and strategies as to how to manage whatever is left after the settlement (e.g., unpaid claims)
- » How to negotiate and convince Plaintiff's counsel?
- » How to manage client's expectations?
- » What kind of assurances can be had?

10:15

### Morning Coffee Break

10:45

### A View from the Bench: Judicial Insights into Drug and Medical Device Products Liability Litigation

**Hon. Michael J. Davis**

*Senior Judge*

**U.S. District Court, D. Minn. (Minneapolis, MN)**

**Hon. David R. Herndon**

*Judge*

**U.S. District Court, S.D. Ill. (East St. Louis, IL)**

**Hon. Loretta A. Preska**

*Senior Judge*

**U.S. District Court, S.D.N.Y. (New York, NY)**

**Hon. Nancy J. Rosenstengel**

*Judge*

**U.S. District Court, S.D. Ill. (East St. Louis, IL)**

Moderator:

**Andrew T. Bayman (Andy)**

*Partner*

**King & Spalding (Atlanta, GA)**

12:00

Networking Luncheon

HOSTED BY:



1:00



### Practical Ways for Incorporating Diversity and Inclusion into Your Trial Team and Litigation Strategy

**Sonia Chen Arnold**

*Assistant General Counsel*

*Lilly Diabetes*

**Eli Lilly and Company (Indianapolis, IN)**

**Rachel Gallagher**

*Director, Legal Counsel*

**Teva Pharmaceuticals USA, Inc (Horsham, PA)**

**Kim M. Schmid**

*Firm Vice Chair and Executive Managing Partner*

**Bowman and Brooke LLP (Minneapolis, MN)**

**Robert Simpson**

*Partner*

**Shipman & Goodwin LLP**

Having a diverse group of attorneys comprised of individuals of different races, genders, sexual orientations, and generations, which is reflective of the community in which cases are tried, makes for a stronger litigation team with a wealth of perspectives and personal experience. In addition to this common sense rationale for diversity, in-house counsel have espoused a commitment to diversity within their law departments and have made it clear that diversity matters to them when vetting and choosing law firms to represent them. In this session, points of discussion will include:

- Making sure all understand why it makes good business sense to have a diverse litigation team within both companies and law firms
- Moving from an intellectual understanding of the need for diversity to measurable efforts showing recruitment, retention, and advancement
  - » Discussing what diversity initiatives are working and designing sustainable diversity program for life sciences companies and outside law firms representing them
- How law firms and companies can best implement policies that will truly effect change and promote a diverse workforce
  - » What specific evidence of diversity are companies seeking from outside counsel
    - Firm composition overall
    - Partners
    - Breakdowns within teams
- Evaluating a firm's efforts in promoting diversity
  - » Having a written plan and timeline in place to measure diversity efforts
  - » Targeting specific deficiencies within the firm's composition
  - » Putting together a leadership team to develop and mentor diverse talent



I enjoy attending ACI's Drug and Med because of the quality presentations, and the interaction with the lawyers and clients that I've had the opportunity to work with for so many years."

**Goodell DeVries Leech & Dann LLP**

## AFTERNOON BREAKOUT SESSIONS

THURSDAY, NOVEMBER 29, 2018

(CHOOSE A OR B)

### A

1:45

#### The Future of Personal Jurisdiction: Decisions of Note and Interpretations Around the Country

**Halli D. Cohn**

*Partner*

**Troutman Sanders LLP (Atlanta, GA)**

**Sarah Heineman**

*Senior Counsel*

**Bayer U.S. (Pittsburgh, PA)**

**John P. Lavelle, Jr.**

*Partner*

**Morgan Lewis & Bockius LLP (Philadelphia, PA)**

In the aftermath of the *BMS* decision, the discussion will focus on how the decision has been applied by the courts around the country as well as what the next stages are and how Plaintiffs' attorneys have looked to circumvent the decision.

OR

### B

1:45

#### Gaining Control Over Third-Party Financing of Litigation: How are Third-Party Funding Deals Structured and What Efforts Have Been Made to Combat the Trend

**John H. Beisner**

*Partner*

**Skadden, Arps, Slate, Meagher & Flom LLP (Washington, D.C.)**

**Christopher Guth**

*Senior Counsel*

**Bayer U.S. (Pittsburgh, PA)**

**Kelley S. Olah**

*Partner*

**Barnes & Thornburg LLP (Los Angeles, CA)**

- Who gets funded?
- How are these deals structured?
- Potential issues with these deals and ways of addressing them
- Status of the current bill introduced in Congress by Senators Grassley, Tillis, and Cornyn that would require disclosure of third-party litigation financing agreements in civil lawsuits

2:30

Afternoon Networking Break

HOSTED BY:

DrinkerBiddle

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Join the Conversation



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ACI: Drug & Medical Device Litigation



# AFTERNOON BREAKOUT SESSIONS

THURSDAY, NOVEMBER 29, 2018

(CHOOSE A OR B)

## A

3:00

### Best Strategies for Moving Forward When Faced with an Actual Multi-Plaintiff Trial

**Amanda T. Perez**

*Assistant General Counsel*

**Pfizer Inc. (New York, NY)**

**Hildy Sastre**

*Partner*

**Shook Hardy & Bacon L.L.P. (Miami, FL)**

**David B. Sudzus**

*Partner*

**Drinker Biddle (Chicago, IL)**

- Best strategies for arguing against multi-plaintiff trials
- Once faced with this type of a trial, what are the best safeguards that can be put in place?
  - » How to structure limines to preserve the record?

OR

## B

3:00

### Has Proportionality Changed Anything? Impact of Recent Decisions and New Technologies on the Scope of Discovery

**Julie Y. Park**

*Partner*

**Morrison & Foerster LLP (San Diego, CA)**

**Patrick H. Reilly**

*Partner*

**Faegre Baker Daniels LLP (Indianapolis, IN)**

**Munjot Sahu**

*Counsel – Litigation and Legal Compliance*

**Eli Lilly and Company (Indianapolis IN)**

- Latest arguments on discovery
  - » How have the FRCP amendments affected the scope of discovery?
  - » What are the prevailing arguments on plaintiffs' side? Defense side?
- Has proportionality affected discovery in state courts?
  - » How can defendants protect themselves in state courts that don't have the proportionality standard?
- Putting machine learning to work in discovery
  - » What needs to change to make machine learning accepted in discovery?
  - » How comfortable is the industry with machines making distinctions between responsive and privileged documents?

## A

3:45

### State of the Union on Preemption

**Paul J. Cosgrove**

*Partner*

**Ulmer & Berne LLP (Cincinnati, OH)**

**David W. O'Quinn**

*Member*

**Irwin Fritchie Urquhart & Moore LLC (New Orleans, LA)**

**Erica Valenti Visokey**

*Legal Counsel*

**Stryker Corporation (Stamford, CT)**

Moderator

**Terrence J. Dee**

*Partner*

**McDermott Will & Emery (Chicago, IL)**

- Medical device preemption: what is a parallel claim?
- Implied preemption: recent case law
- Examples of successful defense arguments

OR

## B

3:45

### Strategies for Minimizing the Effects of Innovator Liability

**Henninger S. Bullock**

*Partner*

**Mayer Brown LLP (New York, NY)**

**Sean Fahey**

*Partner*

**Pepper Hamilton LLP (Philadelphia, PA)**

**Janet H. Kwuon**

*Partner*

**Reed Smith LLP (Los Angeles, CA)**

Moderator:

**Brennan Torregrossa**

*Vice President and Associate General Counsel*

*Head of Global External Legal Relations Team (GELRT)*

**GSK (Philadelphia, PA)**

- In light of the California Novartis decision, what attempts have been made to combat the effects of innovator liability?
- Given the different decisions in different jurisdictions, how should companies best position themselves?



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# AFTERNOON BREAKOUT SESSIONS

THURSDAY, NOVEMBER 29, 2018

(CHOOSE A OR B)

## A

4:30



### Training the Future Generations of Life Sciences Attorneys to Become the Next Deans of the Products Liability Bar

Wendy Hufford

Chief Operating Officer & Vice President, US Litigation,  
Risk Management & Human Resources  
Boehringer Ingelheim (Ridgefield, CT)

Heidi Levine

Partner  
Sidley Austin LLP (New York, NY)

Sara K. Thompson

Shareholder  
Greenberg Traurig LLP (Atlanta, GA)

- What do clients seek in an associate?
- Recognizing that clients are the ones who are driving the change, what can clients do to help train young attorneys to become the next generation of go to counsel for these types of cases?
  - » Bringing young attorneys to the discussion table with clients
  - » Encouraging candid conversations between clients and their outside counsel as to what is needed to institute the change



5:30

Conference Adjourns to Cocktail Party

## B

4:30



### Predicting Risk and Examining the Intersection of Traditional Principles of Product Liability Laws with Digital Health and 3D Printing

James M. Beck

Senior Life Sciences Policy Analyst  
Reed Smith LLP (Philadelphia, PA)

Erin M. Bosman

Partner  
Morrison & Foerster LLP (San Diego, CA)

Michelle M. Bufano

Partner  
Patterson Belknap Webb and Tyler LLP (New York, NY)

Vernessa T. Pollard

Partner  
McDermott Will & Emery (Washington, DC)

#### IoT

- What is the reasonable standard of care in creating a secure IoT device?
- Is hackability a design defect?
- What is an adequate warning?
- For how long must device manufacturers provide security monitoring and software updates after selling a product?
- Does user failure to download corrective updates act as superseding cause or failure to mitigate?
- Who is liable when a manufacturer cedes control to a third party?

#### 3D Printing

- Digital health information and cyber-privacy — pitfalls that lead to litigation
- Preventive strategies to cyber-privacy issues involving digital health information, and defenses to litigation
- Liability should connected medical devices be compromised
- FDA initiatives in digital health information and regulation of 3D printing
- Potential defendants in product liability litigation over 3D printed prescription medical products
- Electronic information as a “product” in litigation over 3D printed prescription medical products
- Crossing the service/product line — hospital/physician's office product liability over point-of-care 3D printing
- Non-traditional applications of traditional negligence and warranty causes of action in 3D printing cases

HOSTED BY: **KING & SPALDING**

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Join the Conversation



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# MAIN CONFERENCE DAY THREE

## FRIDAY, NOVEMBER 30, 2018

7:15

### Registration and Continental Breakfast

8:00

### Co-Chairs' Opening Remarks

8:15

## A View from the Bench: Judicial Insights into Drug and Medical Device Products Liability Litigation

**Hon. Ann D. Montgomery**

*Senior Judge*

**U.S. District Court, D. Minn. (Minneapolis, MN)**

**Hon. Arnold L. New**

*Coordinating Judge/Complex Litigation Center*

**Philadelphia Court of Common Pleas (Philadelphia, PA)**

**Hon. Patti B. Saris**

*Chief Judge*

**U.S. District Court, D. Mass. (Boston, MA)**

**Hon. Leda Dunn Wettre**

*Magistrate Judge*

**U.S. District Court, D.N.J. (Newark, NJ)**

**Hon. Susan D. Wigenton**

*Judge*

**U.S. District Court, D.N.J. (Newark, NJ)**

Moderator

**Andrea Roberts Pierson**

*Partner*

**Faegre Baker Daniels LLP (Indianapolis, IN)**

9:30

## Enforcers' Roundtable: Priorities with Respect to Consumer Fraud, False Claims, Anti-Kick-Back, and Off-Label

**David M. Eskew**

*Chief, Health Care & Government Fraud Unit*

**U.S. Attorney's Office, District of New Jersey (Newark, NJ)**

**Lisa D. Kutlin**

*U.S. Attorney's Office, Assistant U.S. Attorney*

**Eastern District of New York (Brooklyn, NY)**

**Gregg Shapiro**

*Chief, Affirmative Civil Enforcement*

**U.S. Attorney's Office, District of Massachusetts (Boston, MA)**

**Pat Stein**

*Senior Asst. Attorney General for Health Fraud*

*Consumer Protection Division*

**The Attorney General's Office for the State of Texas (Dallas, TX)**

Moderators:

**Carolyn M. Hazard**

*Senior Vice President, Associate General Counsel – Litigation*

**Endo (Malvern, PA)**

**Sarah Padgitt**

*Senior Counsel / Litigation*

**Baxter International Inc. (Deerfield, IL)**

- The government's perspective on when and why to prosecute: how do enforcers identify companies for investigations?
  - » What techniques are enforcers using these days to investigate manufacturers?
  - » What specific information are enforcers focusing on?
- Practical considerations for in-house and law firm counsel when faced with DOJ or AG action: best practices for responding to a government investigation

10:30

### Morning Coffee Break

10:45

## Hiring the Right Regional Counsel

**Adam C. Bassing**

*Associate General Counsel*

**UCB, Inc. (Smyrna, GA)**

**David L. Ferrera**

*Partner & Chair, Product Liability Practice Group*

**Nutter McClennen & Fish LLP (Boston, MA)**

**Andrew D. Kaplan**

*Partner*

**Crowell & Moring (Washington, DC)**

It is critically important for lead counsel as well as their in-house clients to ensure that they are working with the best regional counsel. At the same time, as a regional counsel, it is important to ensure that you know how to communicate to the lead as well as in-house counsel your skill-set and what you can contribute to the litigation. In this session, attendees will have an opportunity to develop best strategies for evaluating regional counsel, including what questions to ask and what to look for to make sure that any particular regional counsel is the best fit. The session will also include a discussion of what regional counsel can do to make sure they are answering the needs of their potential clients.

## Continuing Legal Education Credits



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You are required to bring your state bar number to complete the appropriate state forms during the conference. CLE credits are processed in 4-8 weeks after a conference is held.

ACI has a dedicated team which processes requests for state approval. Please note that event accreditation varies by state and ACI will make every effort to process your request.

Questions about CLE credits for your state? Visit our online CLE Help Center at [www.americanconference.com/CLE](http://www.americanconference.com/CLE)



1:30



## Lay of the Land with Respect to Regulatory Trends, Pricing Transparency, and Patent Litigation

**Gina Gencarelli**

*Senior Director, Intellectual Property*

**Par Pharmaceutical (Chestnut Ridge, NY)**

**Lisa LeCointe-Cephas**

*Executive Director, Head of Global Investigations*

**Merck & Co., Inc. (Kenilworth, NJ)**

**Julia Post**

*Attorney*

**Covington & Burling LLP (Washington, DC)**

As an in-house counsel, staying abreast of the key rulings and positions government takes as well as what government enforcers plan to do is crucial when looking to assess the risk for a company. In this session, we will look to extrapolate biggest trends and developments with respect to pricing transparency, patent litigation, and most recent regulation within the past year.

2:30

## Opioid Enforcement and Litigation Landscape

**Patricia A. Barbieri**

*SVP, General Counsel and Secretary*

**Daiichi Sankyo, Inc. (Parsippany, NJ)**

**Terry M. Henry**

*Partner*

**Blank Rome LLP (Philadelphia, PA)**

**Wendy West Feinstein**

*Partner*

**Morgan Lewis & Bockius LLP (Pittsburgh, PA)**

- Enforcement and litigation update
- How is opioid MDL different from a typical MDL?
- Interplay of state consumer protection laws with product liability actions
- New theories of tort liability being advanced and how these expansions of traditional confines of tort law may impact other areas of products tort litigation

3:30

## Conference Ends

11:30



## Engaging the Courts in the Right Way: What Does It Mean to Be an Innovative Thinker as an Outside Counsel

**Greg A. Dadika**

*Associate General Counsel, Litigation*

**Becton, Dickinson and Company (Murray Hill, NJ)**

**Jan Dodd**

*Partner*

**Norton Rose Fulbright US LLP (Los Angeles, CA)**

**Kailee Goold**

*Senior Counsel*

**Cardinal Health (Columbus, OH)**

**Stephen E. Marshall**

*Partner*

**Venable LLP (Baltimore, MD)**

Drug and device companies are changing the face of healthcare in a challenging environment every day. Sharing these companies' values around creativity and advancement is going to become critical in order for outside counsel to distinguish themselves. This session will look to illuminate what lawyers/law firms/in-house counsel can do differently to really add value. Among topics to be discussed are:

- Fostering an environment of creativity
  - » How to think about things differently
  - » Questions to ask yourself/client/team
  - » Rewarding and not punishing outside the box thinking
  - » How to advance these ideas with the business
- Improving communication skills
- Candid feedback during and after the matter
  - » Practical tips for doing this

12:30

## Networking Lunch



ACI's Drug and Med conference is informative and attracts top professionals year after year, giving you the opportunity to develop lasting professional and personal relationships."

**Danaher Lagnese PC**



I attend ACI's Drug and Med because it's an opportunity to meet and reconnect with colleagues who are both my peers at other law firms and clients. I found the presentation on attorney advertising to be very helpful."

**DLA Piper LLP**





THE 23<sup>RD</sup> ANNUAL

# DRUG & MED DEVICE

## LITIGATION CONFERENCE

Appreciates The Support Of The Following Organizations



**Greenberg Traurig, LLP (GTLaw)** has more than 2,000 attorneys in 38 offices in the United States, Latin America, Europe, Asia and the Middle East and is celebrating its 50th anniversary. The firm's Pharmaceutical, Medical Device & Health Care Litigation Group is comprised of more than 90 attorneys, in addition to a team of paralegals, nurse paralegals, and trial consultants across the country. The group is an integral part of Greenberg Traurig's

National Litigation Practice, encompassing 600+ attorneys in offices across the United States. Recent recognitions include national rankings for 2017 "Products Liability & Mass Torts" from Chambers USA Guide and national rankings for 2017 "Practice Liability & Mass Torts Defense: Pharmaceuticals and Medical Devices" from The Legal 500 United States, among others.

### KING & SPALDING

**King & Spalding** is an international law firm with more than 1,000 lawyers in 20 offices globally. Our Pharmaceutical & Medical Device Litigation Team helps manufacturers navigate the litigation life cycle, from risk assessments through trial and appeal. Working with more than 300 lawyers and professionals who devote all or a substantial portion of their practices to life-science clients, we know the industry and governing laws better

than anyone. With over 175 lawyers across the world, our Product Liability team, which Law360 named in 2017 as Group of the Year for the fifth consecutive year, leverages knowledge of the complex science and technology behind today's products to deliver litigation victories. We handle the most significant individual, multidistrict, mass tort and class action lawsuits for pharmaceutical and medical device companies. We offer our clients a depth of trial experience that is increasingly unusual in firms of our size. While many firms can "litigate" cases, we try product cases each year in some of the most challenging jurisdictions for corporate clients. For more information, please visit [www.kslaw.com](http://www.kslaw.com).



**Barnes & Thornburg's** extensive drug and medical device practice has been addressing clients' needs in an efficient and results-driven manner for more than 30 years. As national trial counsel in high-stakes pharmaceutical and medical device litigation for Fortune 500 companies, we have the resources and expertise to help you address the evolving challenges you face.



**Bowman and Brooke LLP** is a nationally recognized trial firm with one of the largest product liability practices in the country. The firm's Pharmaceutical and Medical Device Litigation practice is comprised of experienced, nationally recognized trial lawyers serving as national, regional and local counsel in high-profile individual and mass tort litigation. Whether we are preparing and defending company witnesses in the areas of regulatory, drug

safety, clinical trials, medical affairs and marketing, assisting our clients in assessing their product warnings and package inserts to meet FDA compliance, or navigating complex legal challenges, we aggressively defend our clients based on the unique requirements of each case. With a passion and drive for mastering complex medical, scientific, epidemiological, engineering and regulatory issues, Bowman and Brooke's lawyers deliver legal representation that is innovative, cost-effective and complementary to our clients' core business objectives. The firm's attorneys defend a variety of corporate clients, including many Global 500 companies, in widely publicized catastrophic injury and wrongful death matters, and other complex litigation throughout all 50 states. For more information, please visit [www.bowmanandbrooke.com](http://www.bowmanandbrooke.com).



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for understanding our client's business, anticipating client's needs, unprompted communication, legal skills, quality and keeping clients informed. For more information, visit [www.butlersnow.com](http://www.butlersnow.com) or follow Butler Snow on Twitter @Butler\_Snow.



**Drinker Biddle & Reath LLP** is a national law firm with more than 600 lawyers. We handle all types and aspects of products liability litigation and frequently serve as trial counsel and national coordinating counsel in suits defending prescription drugs, over-the-counter drugs and medical devices including orthopedic implants, antibiotics, contraceptives and antipsychotics. For more information, please visit [www.drinkerbiddle.com](http://www.drinkerbiddle.com).



**Faegre Baker Daniels'** product liability lawyers represent pharmaceutical and medical device manufacturers in all 50 states, Canada and Europe. With 750 lawyers and consultants in the U.S., U.K. and China, our firm offers integrated services to help achieve the goals of life science companies ranging from emerging startups to multinational corporations. With a nationwide ranking in Chambers USA 2018, our product liability

litigation team has served as national, regional and local defense counsel in major pharmaceutical and medical device product liability litigation. Our professionals aggressively defend claims in complex mass tort, toxic tort, multidistrict and class action litigation. In addition, we counsel clients on product liability risk management, regulatory compliance, reimbursement and more. Our practice is supported by our national health and life sciences industry team that includes consultants from our advisory and advocacy division based in Washington, D.C., Faegre Baker Daniels Consulting. For more information, please visit [FaegreBD.com](http://FaegreBD.com).



Comfortable before a jury or arguing a complex MDL motion, **Fox Galvin** attorneys serve in a variety of roles on drug and medical device litigation. From our central St. Louis location, we have experience bringing great results and value as lead trial counsel, on discovery teams and as local/liaison counsel.



Founded in 2006, **Golkow Litigation Services** is a global leader in court reporting, litigation support and trial consulting services. Specializing in highly technical real-time transcription for the pharmaceutical and medical device industry, Golkow has provided deposition and video services in all 50 states and over 25 countries. As a preferred provider of court reporting services for hundreds of national MDLs and other consolidated matters,

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## Morgan Lewis

property, litigation, and related issues that our clients face along the product life cycle, from innovation and emerging business issues, through research and development, regulatory approval, product reimbursement, marketing and distribution, to fraud and abuse, product liability and intellectual property litigation, to mergers and acquisitions, collaborations, and outsourcing. Morgan Lewis offers extensive capabilities and decades of experience coordinating complex national litigation, in addition to providing efficient, powerful solutions for the increasingly demanding discovery environment. We are nationally recognized for our leadership and innovation in developing alternative fee structures. For more information, please visit [www.morganlewis.com](http://www.morganlewis.com).

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**Norton Rose Fulbright** is a global law firm. We provide the world's preeminent corporations and financial institutions with a full business law service. We have more than 4000 lawyers and other legal staff based in more than 50 cities across Europe, the United States, Canada, Latin America, Asia, Australia, Africa, the Middle East and Central Asia.

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have turned to Nutter to defend cases in courts throughout the United States and around the world involving allegedly defective medical devices, drugs, industrial materials, and automotive products. The firm's lawyers are known for their client-centric approach and extensive experience in litigation, business and finance, intellectual property, real estate and land use, labor and employment, tax, and trusts and estates.

**Patterson Belknap Webb & Tyler LLP** is based in New York City with approximately 200 lawyers delivering a full range of services across more than 20 practice groups in both litigation and commercial law. More than half of the attorneys at Patterson Belknap are devoted to litigation. Our litigating partners have tried hundreds of cases, including many of the most complex in their fields. We frequently serve as national and regional litigation counsel for the nation's largest pharmaceutical and medical device companies in products liability matters.

## Patterson Belknap

## ReedSmith

science counsel to the top global pharmaceutical and medical device manufacturers in single plaintiff and complex litigation matters, and is regularly recognized for its successes in high-profile product liability and mass tort litigations.

**Reed Smith's** life sciences team is committed to anticipating and solving legal challenges so you can focus on what you do best: improving and saving lives. In addition to our skilled life sciences transactional and regulatory lawyers, our highly-regarded, full-service life sciences litigation practice features a deep bench of seasoned trial lawyers and appellate specialists. For more than 40 years, Reed Smith has served as national, strategic, trial, and



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New York, the firm represents many businesses, institutions, individuals and government entities throughout New England and nationally.

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Our attorneys have extensive first-chair trial experience and regularly serve as national counsel and regional counsel. Our defense strategies involve an in-depth understanding of our clients' products, processes and markets, and exceptional knowledge of the science and technologies underlying these systems. In addition to our strong litigation experience, we provide liability counseling to promote product safety and minimize future liability exposure. We analyze design and manufacturing processes; test protocols and quality assurance programs; advise on insurance coverage; draft product warnings and instructions; review warranty, disclaimer and indemnity language in vendor and customer agreements; develop programs for proper document retention and post-sale product retrofits or recalls; and ensure regulatory compliance.

We are established leaders in the bar. Several of our attorneys hold leadership positions in organizations including the American College of Trial Lawyers, Product Liability Advisory Council (PLAC), Defense Research Institute (DRI), American Bar Association Committee on Products Liability



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
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
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