

10th Annual PHARMA PACKAGING AND LABELING EAST COAST 2018

21ST - 22ND FEBRUARY, PHILADELPHIA, USA



10th Annual Pharma Packaging and Labeling East Coast Conference

February 21st – 22nd, 2018 | Philadelphia, USA

2018 Speaking Faculty

Patricia Walsh, Director, Head of Global Labeling, **Jazz Pharmaceuticals**

Bob Celeste, Founder, **Center for Supply Chain Studies**

Kashappa-Goud Desai, Associate Fellow, **GlaxoSmithKline**

Christy Wood, Senior Manager Global Labeling, **Jazz Pharmaceuticals**

Kamana Singh, Senior Global Labeling Consultant

Michelle Halliez, Head of Regulatory Labeling, **Sanofi**

Guido Schmitz, Director of Packaging Design, **Bayer**

Shawn R. Ellerman, Assistant Special Agent in Charge, **U.S. Drug Enforcement Administration**

Kathleen Salazar, Director, Global Labeling Operations, **Janssen Pharmaceutical Companies of Johnson & Johnson**

Helen Cocuzza, Senior Manager, Regulatory Affairs, **Foamix Pharmaceuticals**

Matt Sample, Sr. Director, Secure Supply Chain, **BluePoint Laboratories**

Carl Accettura, Vice President, **PharmoRx Therapeutics Inc**

Nicole Quallis, Labeling Manager, **Valeant Pharmaceuticals**

Remon Zakhary, Senior Group Lead Package Design & Development, **Bayer**

Jennifer Nixon Sekawungu, Associate Director Labeling Group, **Janssen Pharmaceuticals**

Niambi Daniels Harris, Director, Commercial QA-GMP, **Daiichi Sankyo**

Eugene Hackett, Director of Corporate Security, **Bristol-Myers Squibb**

Mary Zhang, CBP National Account Manager, **U.S. Customs & Border Protection**

Craig Trautman, CEO, **Intagras**

Peggy Slendorn, Senior Global Account Leader, **CCL Label**

Jackie Leslie, Director Business Development, **BLUE Software**

Arpad Lehoczki, Senior Account Executive, **Global Vision**

Josh Roffman, VP of Product Management, **Loftware**

Kathleen O'Brien, Director of Business Development, **AMPLEXOR Life Sciences**

Ivan Bonzi, Containers B. U. Director, Closures Containers PET Division, **Sacmi Imola S.C**

Orna Bresler, Director of US Account Services, **Perigord**

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	<p>Pharma Packaging and Labeling East Coast 2018 Philadelphia, Pennsylvania Day One: February, 21st 2018</p>
08:15	Registration and refreshments
08:50	Chair's opening remarks – Helen Cocuzza , Senior Manager, Regulatory Affairs, Foamix Pharmaceuticals
09:00	<p>Establishing an accurate global labeling tracking system to ensure your end-to-end labeling is inspection ready</p> <ul style="list-style-type: none"> • Appreciating complexities in tracking labeling changes when managing multiple teams across global sites to minimize challenges encountered due to variations at the individual country level • Recognizing current difficulties in ensuring end to end labeling to determine regulatory repercussion on your company • Evaluating current labeling tracking systems and suggesting improvements to your internal procedures to ensure preparedness for pharmacovigilance inspections • Establishing reporting systems which enable non-compliance changes to be detected quickly and prevent safety update delays and backlogs • Emphasizing improvements in end-to-end labeling to minimize risk and improve safety <p>Patricia Walsh, Director, Head of Global Labeling, Jazz Pharmaceuticals Christy Wood, Senior Manager Global Labeling, Jazz Pharmaceuticals</p>
09:30	<p>Global labeling end-to-end tracking systems</p> <ul style="list-style-type: none"> • Key features that software should provide out of the box • Functionality that is critical to the global regulatory end-to-end tracking process • Implementing end-to-end tracking software in a global regulatory environment • Integration with existing software that are used in other divisions of the organization • Key reports and how to use them <p>Craig Trautman, CEO, Intagras</p>
10:00	<p>Evaluating your inspection readiness strategy to guarantee you are on top of the changing labeling landscape and ready to comply</p> <ul style="list-style-type: none"> • Establishing internal tracking and reporting structures which allow you to react to global guideline alterations without compromising your current timelines • Designing cross departmental reporting systems that allow label amendments in your product portfolios within allocated timelines stipulated by regulators to ensure compliance • Ensuring your labeling documentation is adequately set up to handle inspections by regulators and prevent audits • Developing internal strategies to guarantee labeling information changes are quickly rolled into the supply chain to maximize patient safety • Determining whether internal rearrangements are best achieved through in-house restructuring or outsourcing to guarantee you are prepared for your next inspection <p>Michelle Halliez, Head of Regulatory Labeling, Sanofi</p>

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10:30	<p>Digital solutions for secondary packing</p> <p>Learning how digital printing can relive pain points, solve packing issues, streamline supply, and transform your brand</p> <p>Peggy Slendorn, Senior Global Account Leader, CCL Label</p>
10.45	Morning refreshments and networking
11:15	<p>Understanding the worst drug epidemic in US history: the link between prescription opioid and heroin abuse</p> <ul style="list-style-type: none"> • Explaining the US history drug epidemic to describe the increasing drug abuse in the States • Describing how prescription drug lead to heroin abuses to raise awareness of their seriousness • Highlighting DEA's efforts to reduce the abuse nationwide • Generating solutions to protect the public health <p>Shawn R. Ellerman, Assistant Special Agent in Charge, U.S. Drug Enforcement Administration</p>
11:45	<p>Building a business case for a Label and Artwork Management System</p> <ul style="list-style-type: none"> • Reviewing current label and art management systems and making a business case for a change • Calculating Return on Investment as a result of this change • Aligning new systems to corporate strategies • Exploring the other ROI (Risk of Ignoring) and possible effects on your business <p>Jackie Leslie, Director Business Development, BLUE Software</p>
12:15	<p>Harnessing branding as a story telling tool; remaining relevant in both the present and the future</p> <ul style="list-style-type: none"> • Appreciating brand loyalty as a mechanism to generate a bond between your customer and your company that lasts a lifetime and increases your revenue • Developing a branding strategy with a clear target audience that focuses on generational differences and expectations to ensure your customer loyalty • Recognizing the need for your brand to adapt through time; pinpointing sustainability as a key focus of millennials and the impact this will have on your business model and strategy • Appreciating the role of consistent branding across markets to increase your companies global image • Assessing ways through which branding can differentiate your products from that of the competition to win a competitive advantage <p>Guido Schmitz, Director of Packaging Design, Bayer</p>
12:45	Lunch and networking

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13:45	<p>Sourcing the right contract manufacturers throughout the globe to supply what you need, when you need it</p> <ul style="list-style-type: none"> Assessing logistical complexities of engaging with multiple contract manufactures around the globe and suggesting ways to simplify this process Outlining mechanisms to reduce timelines in finding compliant partners set out by global headquarters to guarantee supplier conformity Establishing partnerships with in-country suppliers to ensure you do not need to ship low value supplies and incur added costs Ensuring packaging components can be produced by your supplier to streamline processes Improving partnerships between CMO's and pharma companies to improve working relationships and prevent process disintegration <p>Remon Zakhary, Senior Group Lead Package Design & Development, Bayer</p>
14:15	<p>Learning why your labeling process is obsolete</p> <ul style="list-style-type: none"> Pinpointing common sources of errors in your labeling process Utilizing technology as a vehicle in improving processes Modernizing the labeling process Unveiling Case Studies to demonstrate ROI <p>Arpad Lehoczki, Senior Account Executive, Global Vision</p>
14:45	<p>DSCSA compliance, serialization, and why it's not just a barcode</p> <ul style="list-style-type: none"> Serialized labeling and packaging, a supply chain paradigm shift towards standardization Presenting product packaging and designing for downstream use beyond distribution Understanding the supply chain impact of FDA's Enforcement Discretion and Grandfathering Guidance Reviewing impacts of serialized exceptions in the supply chain in 2018 and beyond <p>Matt Sample, Sr. Director, Secure Supply Chain, BluePoint Laboratories</p>
15:15	<p>Afternoon refreshments and networking</p>
15:45	<p>Improving cross-departmental relationships and communications to improve labeling process efficiency</p> <ul style="list-style-type: none"> Defining shared cross-departmental priorities to ensure engagement and prevent silos Establishing strong and sustainable connections between packaging and labeling teams and other labeling stakeholders Developing an effective communication strategy to enable early visibility of upcoming labeling changes and projects Using technology to streamline communications across business partners <p>Kathleen Salazar, Director, Global Labeling Operations, Janssen Pharmaceutical Companies of Johnson & Johnson</p>

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16:15	<p>Leveraging enterprise labeling to sustain compliance, protect brand and optimize supply chain performance</p> <ul style="list-style-type: none">• Integrating labeling with “sources of truth” to ensure accuracy and consistency• Automating label variability to keep pace with global regulatory and customer requirements• Creating a workflow and approval process that supports eSignatures, provides audit trail• Extending labeling to suppliers, CMOs, CPOs, etc. to improve visibility and traceability• Supporting converging labeling and artwork management requirements <p>Josh Roffman, VP of Product Management, Loftware</p>
16:45	<p>Uncovering FDA’s final guidance’s on Nonproprietary Naming of Biologics and Biosimilars and what this means for your current labeling strategy</p> <ul style="list-style-type: none">• Appreciating highly complex manufacturing processes required for biologics to determine how the new guidelines will support pharmacovigilance efforts and minimize improper substitution• Uncovering nonproprietary naming and the importance this holds in today’s labeling practices to guarantee you are up to speed with current regulations• Outlining difficulties in ensuring adequately labeled suffixes on all bio-similar products to ensure compliance with strict FDA requirements• Modifying your manufacturing process to ensure it is equipped to cope with additional labeling needs required by the guidelines• Assessing ways to ensure compliance with incoming regulations with regards to electronic prescribed information databases and its effect on biologics <p>Kamana Singh, Senior Global Labeling Consultant</p>
17:15	<p>Chair’s summary and close of day 1 – Helen Cocuzza, Senior Manager, Regulatory Affairs, Foamix Pharmaceuticals</p>

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Pharma Packaging and Labeling East Coast 2018 Philadelphia, Pennsylvania Day Two: February, 22 nd 2018	
08:15	Registration and refreshments
08:50	Chair's opening remarks – Helen Cocuzza , Senior Manager, Regulatory Affairs, Foamix Pharmaceuticals
09:00	<p>Exploiting blockchain technology to ensure compliance with the Drug Supply Chain Security Act (DSCSA)</p> <ul style="list-style-type: none"> Examining blockchain to utilize technology as a tool to improve track and trace, recalls and reimbursements Identifying blockchain strategies as a norm in the industry to simplify internal processes Outlining serialization best practice to enable small and mid-size pharma to ensure compliance Uncovering ways blockchains' evolving nature can improve record security, traceability and prevent cyber-attacks to utilize in your supply chain Harnessing blockchains' power as an alternative way of doing business with your customer base to improve relationship management <p>Bob Celeste, Founder, Center for Supply Chain Studies</p>
09:30	<p>Breaking Cultural Barriers: Localization 101</p> <ul style="list-style-type: none"> Providing an overview of fundamental concepts for translation and localization Unveiling insights into the translation supplier processes for pharma and biotech Identifying the nuances of culture barriers, and how to manage the technicalities of translation. Presenting effective localization strategies that Life Sciences companies can implement to reduce costs and accelerate timelines <p>Kathleen O'Brien, Director of Business Development, AMPLEXOR Life Sciences</p>
10:00	<p>Exploring the role of US CBP and the Center of Excellence and Expertise, how we can assist industry</p> <ul style="list-style-type: none"> Enhancing knowledge on National Account Management Program Uncovering Customs-Trade Partnership Against Terrorism (C-TPAT) program Developing appreciation for Importer Self-Assessment (ISA) program and how this impacts you Addressing Trusted Trader program for your pack and label strategy <p>Mary Zhang, CBP National Account Manager, U.S. Customs & Border Protection</p>
10:30	<p>Compression Blow Forming (CBF), the new way into containers manufacturing</p> <ul style="list-style-type: none"> Discovering the new CBF technology Presenting ways to achieve an innovative blow moulding process merging the best, most valued characteristics of the alternative technologies Demonstrating examples of superior bottle quality Exploiting benefits from CBF technology for your business

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Ivan Bonzi, Containers B. U. Director, Closures Containers PET Division, **Sacmi Imola S.C**

10:45 Morning refreshments and networking

Designing combination products which are high-quality, adaptable and evolve throughout the supply chain

11:15

- Design high-quality combination products by applying quality by design principles
- Assessing current global regulatory landscape for CPs to ensure compliance
- Uncover what are the correct packaging types, materials and designs needed to ensure that deceives are in line with regulatory expectations
- Ensure labels and products are designed in a way which is adaptable as trials develop and move towards the commercial stage
- Highlighting the added costs that current FDA guidelines will have on the industry as a whole to determine cost-saving alternatives
- Maximizing Human Factor Studies (HFS) for CP's effective usage

Kashappa-Goud Desai, Associate Fellow, **GlaxoSmithKline**

Balancing regulatory requirements with brand design to prevent regulatory rejections and make sure you stand out amongst the crowd

11:45

- Recognizing the demands placed on packaging and labeling teams to ensure patient safety information is prioritized
- Exploiting package and label surface area to ensure space is utilized to its fullest capacity and waste is minimized
- Establishing priorities between labeling and branding teams to ensure these do not conflict and result in timeline delays
- Outlining novel packaging designs which both hold all required information while standing out against your competition
- Appreciating how electronic labels can free up packaging space and enable greater creativity to give you a competitive advantage

Helen Cocuzza, Senior Manager, Regulatory Affairs, **Foamix Pharmaceuticals**

Designing packaging which is more secure and appealing both to the regulator and the consumer; pushing the Crayola box to its limits

12:15

- Outlining FDA serialization and changing artwork requirements to ensure regulatory compliance
- Considering patient adherence and patient persistence as part of the packaging and labeling
- Enhancing information flow between regulatory and artwork teams to guarantee documentation used in artwork approval process is a match with regulatory requirements to prevent delays
- Promoting innovative packaging designs to guarantee your products stand out against those of the competition

Carl Accettura, Vice President, **PharmoRx Therapeutics Inc**

12:45 Lunch and networking

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14:00	<p>Speaker Hosted Roundtables Interactive roundtable sessions offer a unique opportunity to come together with your peers to share best practice and develop solutions to critical challenges facing the industry as a whole. Hosted by industry experts and each focused on a single issue, roundtables are an exciting, interactive way to build your personal network and learn from the experience and expertise of others.</p> <p>Each roundtable session lasts for 45 minutes, and delegates may attend up to 2 roundtables</p>
Roundtable 1	<p>Packaging & Labeling Development for New Product Launches Orna Bresler, Director of US Account Services, Perigord</p>
Roundtable 2	<p>Mastering your packaging to ensure your products integrity is guaranteed both while in storage and in transit Niambi Daniels Harris, Director, Commercial QA-GMP, Daiichi Sankyo</p>
Roundtable 3	<p>Discussing FDA's guidance with regards to DSCSA to improve industry standards Nicole Quallis, Labeling Manager, Valeant Pharmaceuticals</p>
Roundtable 4	<p>Establishing best practice to prevent counterfeits entering your drug supply chain Eugene Hackett, Director of Corporate Security, Bristol-Myers Squibb</p>
Roundtable 5	<p>Continuing the conversation on end-to-end labeling to improve patient safety Jennifer Nixon Sekawungu, Associate Director Labeling Group, Janssen Pharmaceuticals</p>
15:30	<p>Chair's summary and close of conference – Helen Cocuzza, Senior Manager, Regulatory Affairs, Foamix Pharmaceuticals</p>