

"Uniting industry leaders to analyse advanced commercial developments & to identify successful strategies of Biosimilars"

23rd & 24th June 2021, Virtual Conference (Time Zone - BST)

AGENDA AT A GLANCE

Key Speakers Include



CLAUDIA LOUATI Policy Advisor



FREDRIK SUNDBERG Global Director, Strategic Technology Partnership, Cytiva (Formerly GE Life Sciences)



RAJESH DESIKAN
Vice President & Head, US Marketing,
Oncology & Immunology Biosimilars
Fresenius - Kabi



JULIE MARECHAL JAMIL
Director Biosimilar Policy & Science
Medicines for Europe



MATTHEW TURNER
Senior Director Government Affairs and
Policy Biosimilars Europe, Asia, Latam &
Canada, Fresenius Kabi



CECIL NICK Vice President Parexel



ANNA AILLERIE Brand Management Lead, Europe Dr Reddy's Laboratories SA



MIGUEL NAVARRETE OLMEDO Hospital & Biosimilars Commercial Director STADA Arzneimittel



LOUIS BOON CSO Polpharma Biologics



HANMANT BARKATE Vice President & Head Medical Services (India, MEA), Glenmark



SWEETY MATHEW Regulatory Affairs Biocon



NIKLAS EKMAN
Head of the Biological Section, Finnish
Medicines Agency (Vice-Chair of the
Biosimilar Working Party (BMWP), EMA)



OMAR ALI Pharmacist Consultant QIPP Adviser Payer Network



RENE ANOUR Senior Clinical Expert/Head of National Scientific Advice, Austrian Medicines & Medical Devices Agency (AGES)



BER OOMEN Executive Director, ESNO (European Specialist Nurses Organisations)



MICHEL MIKHAIL International Expert in Biosimilars Global Expert in Regulatory Affairs



LENNEKE DE WINTER Senior Scientist USP Polpharma Biologics



JAKOB LANGE Senior Director Delivery Systems Ypsomed



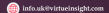
ANDREAU SOLDEVILA Founder & CEO Syna Therapeutics



INGRID SCHWARZENBERGER Senior Regulatory Consultant, Independent Consultant (Former Head Global Regulatory Policy, Sandoz)









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BERT THOMAS Senior VP, Business Development Bio -Thera Solutions



SANDY EISEN Chief Medical Officer Frontline Pharma Consulting Ltd



ZIQUN HAN Director Zen Medical Science



MARIE MANLEY Partner, Head of the UK Life Sciences Sidley Austin



ALEXANDER ROUSSANOV Life Sciences Regulatory & Privacy Lawyer Arnold & Porter



ROBERT A. JOHNSTONE Board Member International Alliance of Patients Organisations



JOHAN DE MUNTER Assistant Nurse Manager Cancer Center University Hospital Ghent, President, European Oncology Nursing Society

Plus more COMING SOON.....

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Key Speakers

Conference Info Day One

Day Two

Booking Details

CONFERENCE INTRODUCTION

Global biosimilars market size is expected to grow from USD 35.7 billion by 2025 from USD 11.8 billion in 2020, at a CAGR of 24.7%. This industry is experiencing significant growth due to the rising incidence of chronic diseases and the increasing demand for biosimilars due to their cost-effectiveness. The growth in the market may be attributed to the cost-effectiveness of the biosimilars when compared to reference biologics coupled with the patent expiration of the many blockbuster biologic drugs. Increasing investment by the companies for the development of biosimilars will also be the key factor driving the market. According to a recent report, as many as nine drugs in the biologics category have either gone off patent or will do so by 2025. Their total revenue was \$62 billionin 2018. This creates a major opportunity for their respective biosimilars. It is estimated that revenue of these biosimilars will grow by 24 per cent annually for seven years to \$13.3 billion in 2025. in the US and Europe. That offers a big opportunity

Our 15thBiosimilars Congregation 2021 will provide insight into the current state of play in the EU and stimulate debate, in a multi-stakeholder setting, on the vital role of biosimilar medicines in the sustainability of healthcare systems. Beyond a comprehensive multi-stakeholder setting, on the vital role of biosimilar medicines in the sustainability of healthcare systems. Beyond a comprehensive outlook of key European market access policies, our speakers will outline the key recent developments in regulatory science and regulatory policy in the EU and other international jurisdictions. Special emphasis will be placed on strengthening the link between regulators and medical communities as an essential basis for greater understanding and acceptance of biosimilar medicines. This Biosimilars conference will focus on multiple aspects of Biosimilar product development to successfully deliver safe, Biosimilar products to the market place. By attending this conference, you will gain a comprehensive outlook on the key issues surrounding Biosimilars. This event will provide an important platform for Biosimilars stakeholders to discuss and share best practices in furthering Biosimilars development.

It gives me great pleasure in welcoming all of you to the Virtue Insight's 15thBiosimilars Congregation 2021.



🌟 CERTIFICATION 🥋



E-Certificate of attendance would be provided to attendees on request, upon completion of conference

KEY THEMES

- Strategies for market access and expansionby identifying key changes and future projections
 Understanding the biosimilars opportunity for pharma companies
 Consequences of Brexit & this pandemic situation on Biosimilars
 Current Challenges and Opportunities for future-Strategies in developing Biosimilars
 A Clinician's Guide to Biosimilars in Oncology: understanding the Science of Extrapolation and Interchangeability
 Biosimilars Pricing & Market access Bringing it faster into market

- GMP, GCP, QC & R&D

 Current challenges and opportunities strategies to develop Biosimilars

 Payer perspective on biologics and Biosimilars

 Biosimilar Interchangeability: The newest regulation

- Biosimilar Physicians and Patients perspective CMC, Preclinical and clinical considerations for Biosimilars and Follow-on Biologics
- Impact of Technology
- Commercial landscape & market access for Biosimilars: Predicts to prepare for a successful tomorrow

 Hear case studieson biosimilars drug development from pre-clinical to clinical and the various testing required such as immunogenicity and bio-similarity tests
- Research-based industry Biosimilar strategies
- Considerations for the analytical similarity assessments when designing a Biosimilar development program
- Determining the right investments & potential returns from Biosimilars
- Latest developments in regulation to increase speed of entry and compliance
- Future of next generation biosimilars
- Be part of a major networking opportunity

WHO SHOULD ATTEND

CSOs, CMOs, Vice Presidents, Presidents, Heads, Directors, Team Leaders, and Senior Scientists from the following roles:

Biopharmaceuticals/ Biotherapeutics, Follow on Biologics/Follow on Proteins/Biosimilars, Biologics/Biotechnology/ Biogenerics, Legal Affairs, Intellectual Property, Health Economics, Pricing and Reimbursement, Clinical Immunology, Principal Scientist, Chief Scientific Officer, Process Control and Analytical Technologies, Analytical Characterisation, Regulatory Compliance, Pharmacovigilance, Drug Safety & Risk Management, Quality Affairs/ Quality Control, New Product Development, Process Science, Portfolio Management, Research & Development, Business Operations, Scientific Affairs, Commercial Affairs, Marketing

WHY SHOULD YOU ATTEND

Get more from the event, enjoy and make the best out of our dedicated networking drinks time, meet the leading international vendors showcasing the products of tomorrow in the co-located exhibition. **Expand your knowledge** of the latest business models and strategies in the high-level conference. Whether you are on the branded or generic side, you cannot afford to miss this opportunity to benchmark your tactics and strategies against the industry leaders who will be the first to traverse the pathway. Devise an immediate action plan for your biosimilar prosecution and litigation strategies in light of the barriers to entry, research and development costs, and regulatory hurdles, which are balanced against an enormous potential for increased profit margins.







identify successful strategies of Biosimilars"

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DAY ONE - 23rd June 2021

09:30 - Welcome Address & Virtual Conference Platform Moderator: **Instructions LOUIS BOON CSO** Polpharma Biologics **MARKET OVERVIEW & ANALYSIS** Panellists: 09:40 - The biosimilar business case - A Growth formula for generics biosimilars **CLAUDIA LOUATI Policy Advisor** • Generics and Biosimilars: Industrial Strategy **FDA** Globalization of Biosimilars GMP, GCP, QC & R&D JULIE MARECHAL JAMIL Innovation and Technology for Biosimilar Development Director Biosimilar Policy & Science Licensing of biosimilars **Medicines for Europe** **MATTHEW TURNER** Senior Director Government Affairs and Policy PAYER'S PERSPECTIVE Biosimilars Europe, Asia, Latam & Canada, Fresenius Kabi 10:20 - Biosimilars - Bringing it into the market quickly RENE ANOUR Senior Clinical Expert/Head of National Scientific Advice Strategies in overcoming obstacles in Biosimilar Austrian Medicines & Medical Devices Agency (AGES) development Effective strategies for product design **JOHAN DE MUNTER** How Payers are aligning biosimilars? Assistant Nurse Manager Cancer Center University Global impact of biosimilars over generics Hospital Ghent, President, European Oncology Nursing Requirements for product development program Society Bridging the 'uncertainty gap' between payers & pharma - the shifting paradigm **SANDY EISEN** What to expect in the next 2 years? **Chief Medical Officer** Frontline Pharma Consulting Ltd **OMAR ALI Pharmacist Consultant QIPP Adviser Payer Network 12:00 - Topic TBC** **BER OOMEN** 11:00 - Morning Coffee/Tea & Discussion **Executive Director ESNO (European Specialist Nurses Organisations) CHALLENGES & OPPORTUNITIES** 12:40 - Networking luncheon 11:20 - Keynote Panel Discussion: Understanding the biosimilars opportunity for pharma companies

PATIENT'S PERSPECTIVE

13:50 - Analysing Physicians and Patients perspective

- National and International developments in biosimilar medicines
- Physicians education Challenges
- Importance of Physician and Patients inputs to shape the international standards for biosimilars
- Encouraging physicians Policies

AT A GLANCE

Conceptualised By





• Latest developments, Trends and Future of Biosimilars

Issues to overcome to increase uptake of biosimilars

Generate enough interest and enthusiasm for biosimilars

Lack of stakeholder confidence - what does this lead to?

Looking at sustaining growth through pandemic Current Challenges and Research trends in Biosimilars &

Biologics





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AGENDA AT A GLANCE

DAY ONE - 23rd June 2021

Physicians / pharmacist collaboration **Panellists:** Harmonizing global standards to ensure safety and **ROBERT A. JOHNSTONE** efficacy of biosimilars **Board Member International Alliance of Patients Organisations** 14:30 - Reducing time to market: Fast track process MIGUEL NAVARRETE OLMEDO development towards biosimilarity Hospital & Biosimilars Commercial Director **STADA Arzneimittel** Accelerated high-titer cell line generation Upstream process modulation to obtain biosimilarity **BERT THOMAS** Senior VP, Business Development Biosimilar cell line portfolio for out-licensing **Bio -Thera Solutions** LENNEKE DE WINTER Senior Scientist USP **ANNA AILLERIE** Polpharma Biologics Brand Management Lead, Europe Dr Reddy's Laboratories SA MICHEL MIKHAIL 15:10 - Afternoon Tea/Coffee **International Expert in Biosimilars Global Expert in Regulatory Affairs** 15:30 - Solution Provider Presentation 16:40 - End of Day 1 of conference. For sponsorship opportunities please contact

COMMERCIALISATION & MARKET ACCESS

16:00 - Panel Discussion: Commercial landscape & market access for Biosimilars: Predicts to prepare for a successful tomorrow

- Comparison of US/EU biosimilar developments, policies and guidelines
- The impact of Biosimilars on the competitive landscape of biological products
- Challenges and obstacles faced by manufacturers in developing biosimilars
- Bringing the next generation of Biosimilars to the market
- Ensuring market access and reimbursement
- Evidence generation will be the key to future success
- Stakeholders approach in successfully bringing Biosimilars to the market

Moderator:

RAJESH DESIKAN

info.uk@virtueinsight.com

Vice President & Head, US Marketing, Oncology & Immunology Biosimilars, Fresenius - Kabi

FOR SPONSORSHIP OPPORTUNITIES:-

Sponsorship or exhibition is the best way to speed network with decision makers. The world leader speakers in our conferences attract niche delegates from all over the world. This would be a wonderful opportunity to reach the right audience and save money and time on all your other advertising gimmicks. To give you an advertising edge we constantly update the industry pioneers via emails/news letter about the event and advertise the event via different forms of media.

Sponsorship Enquires - info.uk@virtueinsight.com







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AGENDA AT A GLANCE

DAY TWO - 24th June 2021

09:30 - Welcome Address & Virtual Conference Platform	BUSINESS MODELS
Instructions	12:00 - Solution Provider Presentation
•••••	For sponsorship opportunities please contact
MANUFACTURING	info.uk@virtueinsight.com
09:40 – Overcoming Development Challenges for Biosimilars with Effective Bioprocessing and Analytics Quality by Design	12:20 - Solution Provider Presentation
 Current market and manufacturing challenges Cost-effective manufacturing approaches, quality-by-design and rapid quality control 	For sponsorship opportunities please contact info.uk@virtueinsight.com
Novel analytics and regulatory strategies for bringing next generation Biosimilars to market	12:40 - Networking luncheon
FREDRIK SUNDBERG Global Director, Strategic Technology Partnership	
Cytiva (Formerly GE Life Sciences)	CLINICAL
10:20 - Pricing & Market access	13:50 - A Clinician's Guide to Biosimilars in Oncology: Understanding the Science of Extrapolation and Interchangeability
 Best Practices for a competitive Market Impact of pharma pricing over innovation Biosimilars influence on pricing and reimbursement Market access success rate Biosimilar Milestone 	HANMANT BARKATE Vice President & Head Medical Services (India, MEA) Glenmark
	14:20. Learning complisition and minuting
11:00 - Morning Coffee/Tea & Discussion	14:30 - Leanbio capabilities and pipeline
	Lean Development for BiosimilarsLean bioproduction
	Cost effective CMC developmentReduced non clinical and clinical program
CLINICAL	ANDREAU SOLDEVILA
11:20 - Clinical Data Requirements for Biosimilars: Have the regulators got it right?	Founder & CEO Syna Therapeutics
Residual uncertainty after demonstrating similarity at the structural and biological activity level	
 Are studies in patients needed? What concerned are addressed by patient data The position of different regulatory agencies	15:10 - Afternoon Tea/Coffee
CECIL NICK Vice President Parexel	15:30 - Self-injection devices for biosimilars - overview and market trends

• Introduction to devices for self injection with market

overview







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DAY TWO - 24th June 2021

- Particular needs for the biosimilar area including IP aspects
- Recent trends from the market place 1 larger volumes
- Recent trends from the market place 2 carbon neutrality and renewable materials

JAKOB LANGE

Senior Director Delivery Systems Ypsomed

REGULATION OVERVIEW & UPDATE

16:10 - Panel Discussion: The developing regulatory framework in advanced and developing markets

- Market and regulatory developments in the Europe and globally
- Predicting the post Brexit changes in biosimilars regulation in UK
- EMA's act on switching & interchangeability?
- How regulators, payers and policy makers take initiatives to make healthcare more sustainable
- · Collaboration with HTA's for patients benefit
- CMC regulatory considerations for Biosimilar products development
- Regulatory changes necessary to maximize biosimilars potential
- The way forward

Moderator:

LOUIS BOON CSO Polpharma Biologics

Panellists:

INGRID SCHWARZENBERGER

Senior Regulatory Consultant, Independent Consultant (Former Head Global Regulatory Policy, Sandoz)

NIKLAS EKMAN

Head of the Biological Section, Finnish Medicines Agency (Vice-Chair of the Biosimilar Working Party (BMWP), EMA)

SWEETY MATHEW

Regulatory Affairs Biocon

ZIQUN HAN

Director

Zen Medical Science

MARIE MANLEY

Partner, Head of the UK Life Sciences Sidley Austin

ALEXANDER ROUSSANOV

Life Sciences Regulatory & Privacy Lawyer Arnold & Porter

16:50 - End of conference

FOR DELEGATE REGISTRATIONS:-

Our potent conference agenda delivering the latest information and the world class leaders as speakers attract delegates to attend from around the world. We aim for our attendees to be equipped with knowledge of latest developments & enable them to network with the industry key personnel.

Delegate Registration - info.uk@virtueinsight.com









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AGENDA AT A GLANCE

REGISTER ONLINE:

Link: https://www.virtueinsight.com/pharma/15th-Biosimilars-Congregation-2021-Virtual-Conference/products/

For Multiple Book	cings - Photocopy this form and send it to info.uk@v	virtueinsight.com	
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Surname		FOR BANK TRANSFER:	
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Email		TERMS AND CONDITIONS:	
	How to Pay one of the following payment options)	Payment terms: Virtue Insight requires the full amount to be paid before the conference. We may refuse entry to delegates who have not paid their invoice in full.	
EARLY BIRD PRIO 1 Delegate @ £300 +	RESERVATION PRICING: CE +VAT (Valid Till 17th May 2021)	Cancellations: Delegates and vendors are subject to the following charges and refunds upon withdrawal or cancellation between 2-3 month's prior 75% cancellation fee/ 25% refund. Less than 2 months prior to the event Full cancellation fee / No refund.	
3 Delegates @ £700 STANDARD RAT	+VAT (Valid Till 17th May 2021)	Administration Fee: If you cancel your participation (once confirmed) and haven't paid the attendance fee you will be liable to pay an administration fee of £200	
-	+VAT (Valid From 18th May 2021) 0 +VAT (Valid From 18th May 2021)	Substitutions/Name Change: If you are unable to attend you may nominate, in writing, another delegate to take your place at any time prior to the start of the event. This can be done at no extra cost.	
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