"A critical guide for successfully conducting clinical trials" 05th November 2020, Virtual Conference (Time Zone – IST)

Key Speakers Include



BANOTH VENKATESWARLU Assistant Drugs Inspector, Central Drug Standard Control Organization (CDSCO)



CHIRAG TRIVEDI Director & Head of Clinical Study Unit Sanofi-Aventis



PRASANNA GANAPATHI Associate Vice President – Global Clinical Sciences Mylan Laboratories



SHUBHANGI DESAI Director - Global Clinical Trial Management Abbott (Singapore)



YASMIN SHENOY Director-Regulatory Affairs Sanofi-aventis



ARUN BHATT Consultant - Clinical Research & Development



MILIND ANTANI Leader, Pharma and Healthcare Nishith Desai Associates



SHREEKANT SAPATNEKAR Director - Clinical Research Lilavati Hospital & Research Centre



SANDESH SAWANT Director and Head Clinical Trials Cipla



MURTUZA BUGHEDIWALA Associate Director - GCO Johnson & Johnson



S.R.SALUNKHE Former Assistant commissioner FDA Maharashtra



RAJENDRA JANI Senior Subject Expert & Advisor Clinical Research Consultant



Novartis

MURUGANANTHAN KRISHNAN Country Monitoring Head - Global Development Operations, Global Drug Development,

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CEO - QbD International, United Nations Adviser, Member Editorial Board Journal of Generic Medicines, England



ANISH DESAI Director IntelliMed Healthcare Solutions



SRIRUPA DAS Director - Medical Affairs Abbott

RANJIT BARSHIKAR



SUTAPA BANDYOPADHYAY NEOGI **Professor**, International Institute of Health Management Research (IIHMR)



ANANT PATIL Asst Professor Department of Pharmacology Dr DY Patil Medical College



PRATIKSHA PALAHE Head NFB National facility for Biopharmaceuticals



ARUN GUPTA Head Medical Affairs & Clinical Research Dabur Research & Development Centre



JYOTSNA PATWARDHAN Head Development QA Novartis



REBU NINAN Director - Healthcare Consulting GlobalData Healthcare

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"The sessions was informative on products and the discussions was fantastic. It has given a better idea on where the industry is heading. To start a new era of clinical trials, this seems to be a promising start for the industry. The second innings for the clinical trials seems promising technically as well as operationally"

Sr. Manager - Global RA, Abbott



VAIBHAV SALVI Head - Project Management and Strategic Initiatives, Clinical Study Unit, Sanofi



KARAN THAKKAR Senior Manager Pfizer



PRASHANT A. PANDYA DGM-Global Strategic Sourcing - Scientific Affairs, Mylan Laboratories



Key Speakers Include

SAKHARAM GARALE Founder & CEO **RENOVARE Healthcare Solutions**



SUJAY KULKARNI **Business Partner/ Medical Expert Novartis**



SANDEEP JAGTAP Assist. General Manager - Clinical R & D **Mylan Laboratories**

WHO ATTENDS?



PRANJAL BORDOLOI Vice President - Clinical, Medical Affairs & Pharmacovigilance, Veeda Clinical Research



VISHWAS SOVANI **Founder Director** Pharmawisdom

PRASHANT BODHE Director CliniSearch



EXHIBITOR'S



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"Very well structured summit. Adequate knowledge on the topic of clinical research, regulations, guidelines, amendments, etc are well discussed"

ICRI (Institute of Clinical Research, India)

AGENDA AT A GLANCE

CONFERENCE INTRODUCTION

We are glad to announce the **11th Annual Clinical Trials Summit 2020 to be held in Mumbai, India during 28th May 2020.** The global Contract Research Organization (CRO) market size was estimated at US\$ 34.5 billion in 2018 and is projected to reach US\$ 55.3 billion by 2024, growing at a CAGR of 8.2% during 2019 to 2024. Indian clinical trials market size is expected to reach US\$ 3.15 billion by 2025. It is projected to register a CAGR of 8.7% over the forecast period.

Increasing cost of drug development is expected to drive the growth. Drug maker and sponsor companies are under pressure to replace the revenue loss caused by generics, increasing patent expiry, number of partnerships to identify biologics, and growing R&D costs, which has made drug development more expensive and complex. In addition, growing pressure on market players to follow stringent timelines has increased the demand for outsourcing of research activities.

This Conference brings together Researchers, Doctors, Principle Investigators, Clinical research sites, CROs, CMOs, Investors, and senior executives from Biopharma, Medical devices and Pharmaceutical industries around the globe to discuss, reflect on and develop their ideas. It offers many opportunities for professional contact and development

The **11th Annual Clinical Trials Summit 2020** will provide opportunities for everyone to learn, gain insight and new skills, and also, there will be many opportunities to network and meet new peoples from industry and patient's clinical organizations, **11th Annual Clinical Trials Summit 2020** hope to lead to new successful collaborations in the future. It is definitely our aim and ambition for every participant to return home somehow enriched, both professionally and on a personal level.

KEY THEMES DISCUSSED

- Current key changes and challenges for trials in India.
- Challenges while growing your research development.
- Discuss the various principles and methods for implementing the project life cycle at each important phase.
- Setting up the best position to sustain an agile procedure for your study design
- Addressing biomarker integration into a protocol while remaining agile.
- Planning and managing an adaptive clinical trials Challenges and the best practices to achieve
- Discussing on the flexibility to redesign clinical trials at intermediate stage.
- Current evolution of clinical trials: Addressing challenges for the future?
- Discussing the major challenges with global trials How can they be overcome?
- Discussion and development of functional processes in living organisms
- Provide clinical research and construct foundations for biomedical research and forms of study.
- Required advancement of clinical trials and new medicines
- Discussing the pharmaceutical industry's financed portion.
- Discussing future Current challenges and overview to look out for while collaborating with the CROs and Sponsorships company
- Establishing an effective and quality collaboration between Sponsors and CROs and if it fails, what are the costs and negative results of a failed partnership?
- Clinical studies and patients assessment
- Identification of study participants and evaluation of physiological or health outcomes.
- New drugs and clinical trial rules to prepare for regulatory inspection and to improve the quality and lifespan of patients
- Be part of a major networking opportunity

WHO SHOULD ATTEND AND WHO YOU'LL MEET

CEO's, CTO's, CIO's, Presidents, Vice Presidents, Directors Heads & Managers of:

Clinical Research & Development, Clinical Research Services, Clinical Operations, Clinical Data Management, Clinical IT, Clinical Trials, Medical Affairs, Regulatory Affairs, Compliance, Quality control / Assurance/GCP, Clinical Study Design, Safety Surveillance, Subject Recruitment, E-Clinical Systems

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WHY SHOULD YOU ATTEND?

Get more from the event, with a **broader scope bringing the whole communications value chain together**. Enjoy and make the best out of our **dedicated networking 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.

WHY EXHIBIT?

Make Sales Debut new products Profile your brand Meet new business partners Develop key relationships Educate pharma and biotech companies



🗧 CERTIFICATION 🎡

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

"A critical guide for successfully conducting clinical trials"

05th November 2020, Virtual Conference (Time Zone - IST)



Conference was very informative and the positive of the conference is Mr. Bangarurajan sir. Regulatory perspectives were very much good and clarified. Very much *happy for the conference*

Research Associate, The Himalaya Drug Company

DAY ONE - 05th November 2020

09:20 - Chairperson opening remarks

RANJIT BARSHIKAR

CEO - QbD International, United Nations Adviser Member Editorial Board Journal of Generic Medicines, England

MARKET OVERVIEW & ANALYSIS

09:30 - Challenges while growing your research development.

- · Discuss the various principles and methods for implementing the project life cycle at each important phase.
- Discuss key factors affecting income from operations.
- Key factors that can make profit margins or break them.
- Analytics to help monitor critical measures of development.

.....

10:00 - Clinical Trials in India: A peek into the future

- The evolution of clinical trials in the decade starting 2010
- Current landscape for clinical trials in India
- What does the future hold for us
- What do we do to enhance patient centricity in clinical trials

CHIRAG TRIVEDI

Director & Head of Clinical Study Unit Sanofi-aventis

10:30 - Morning Coffee/Tea & Discussion

CHALLENGES & OPPORTUNITIES

10:50 - DISCUSSION WITH EXPERTS: Planning and managing an adaptive clinical trial - Challenges and the best practices to achieve

- Discussing on the flexibility to redesign clinical trials at intermediate stage.
- With cross-over research, how adaptive designs can work
- Understanding the logistical barriers that must be over come in order to use adaptive designs within existing trial frameworks.

Addressing the review attempts to clarify the variations between several common types of adaptive designs proposed.

- What are the main sources of the funding and how regulatory agencies are solving these limitations.
- How do you handle the contracting? What are the extra costs?
- What advice would you have on to persuading people that the implementation and designs does not pose a risk?
- Adaptive clinical study designs in global trials

Moderator:

RANIIT BARSHIKAR

CEO - QbD International, United Nations Adviser Member Editorial Board Journal of Generic Medicines, England

Panellists:

RAJENDRA JANI Senior Subject Expert & Advisor **Clinical Research Consultant**

SHREEKANT SAPATNEKAR **Director - Clinical Research** Lilavati Hospital & Research Centre

SANDESH SAWANT

Director and Head Clinical Trials Cipla

ARUN GUPTA

Head Medical Affairs & Clinical Research **Dabur Research & Development Centre**

SANDEEP JAGTAP

Assist. General Manager - Clinical R & D **Mylan Laboratories**

ANANT PATIL

Asst. Professor Department of Pharmacology Dr DY Patil Medical College

- 11:30 DISCUSSION WITH EXPERTS: Current evolution of clinical trials: Addressing challenges for the future?
- Discussing the major challenges with global trials -How can they be overcome?
- What are the main challenges faced in the field of clinical trials and how tackle those issues.
- Challenges associated with balancing the desire for external validity, pragmatic trials, precision medicine, operational complexity and the expense of clinical trials

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"The sessions was informative on products and the discussions was fantastic. It has given a better idea on where the industry is heading. To start a new era of clinical trials, this seems to be a promising start for the industry. The second innings for the clinical trials seems promising technically as well as operationally"

Senior Business Analyst, HCL Technologies

AGENDA AT A GLANCE

DAY ONE - 05th November 2020

- Addressing the complexities of future clinical trials to be more practical, applicable, reliable and the collection of more meaningful data.
- Enhanced visibility of clinical trials and improving portunity for the future usefulness of trial results and the efficiency of their conduct
- Collaborative efforts to drive in the right direction

Moderator:

PRANJAL BORDOLOI Vice President – Clinical, Medical Affairs & Pharmacovigilance Veeda Clinical Research

Panellists:

PRASANNA GANAPATHI Associate Vice President – Global Clinical Sciences Mylan Laboratories

MURTUZA BUGHEDIWALA Associate Director - GCO Johnson & Johnson

SUJAY KULKARNI Business Partner/ Medical Expert Novartis

KARAN THAKKAR Senior Manager Pfizer

SUTAPA BANDYOPADHYAY NEOGI **Professor**, International Institute of Health Management Research (IIHMR)

12:10 - Discussion and development of functional processes in living organisms

• Provide clinical research and construct foundations for biomedical research and forms of study.

Pre-clinical testing and medical treatment evaluation.
Select applicants based on admission into clinical studies.

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• Required data for preclinical studies

12:40 - Networking luncheon

Afternoon Chair Person

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13:50 - Required advancement of clinical trials and new medicines approach

- Discussing the pharmaceutical industry's financed portion.
- Understanding the essential issues of morality and safety.
- Controlling excessively clinical research results.

Sponsorships company

• Questions frequently asked about commonly per formed scholastic clinical research.

14:20 – DISCUSSION WITH EXPERTS: Discussing future – Current challenges and overview to look out for while collaborating with the CROs and

- Establishing an effective and quality collaboration between Sponsors and CROs and if it fails, what are the costs and negative results of a failed partnership?
- Failing partnerships between a pharma company and its CROs What is the reason? Is it financial or the research development time? How can we avoid those situations?
- How to build a success partnership between pharmaceutical companies and CROs in order to per form clinical trials effectively. What is the right effort to shape a positive relationship from both the sponsor and CRO.
- What are some of the obstacles that often occur in partnerships with sponsors and CRO? How to ensure that your team is able to handle these situations effectively?
- What are the needs of different sponsors and how CRO must appreciate and approach?
- What to keep in mind while strategic partnership is a balancing act between the sponsor's need for flexibility and the CRO's need for standardization.

Moderator:

PRASHANT BODHE Director CliniSearch

Panellists:

SHUBHANGI DESAI Director - Global Clinical Trial Management Abbott (Singapore)

MURUGANANTHAN KRISHNAN

Country Monitoring Head - Global Development Operations, Global Drug Development, Novartis

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"Its a good conference and the approach of new ideas get a merge in single pool without any barriers."

Research Associate, Lupin Bioresearch

AGENDA AT A GLANCE

REBU NINAN Director - Healthcare Consulting GlobalData Healthcare

PRASHANT A. PANDYA DGM-Global Strategic Sourcing - Scientific Affairs Mylan Laboratories

SAKHARAM GARALE Founder & CEO RENOVARE Healthcare Solutions

VAIBHAV SALVI Head - Project Management and Strategic Initiatives Clinical Study Unit, Sanofi

PRATIKSHA PALAHE Head, NFB National facility for Biopharmaceuticals

15:10 - Afternoon Tea/Coffee

15:30 - DISCUSSION WITH EXPERTS: Sponsor centric to Patient centric - Improvising patient recruitment their involvement, engaging them

- Patient recruitment and retention in clinical trials What will persuade and impact the patient?
- Ensure patient protection and delivering high quality data
- Overcoming challenges and barriers
- Protecting data and patient privacy
- Patient-Centric Enrolments Planning and Engagement
- Complying with regulatory guidance?
- RWE/RWD , Economic evidence

Moderator:

S.R.SALUNKHE Former Assistant commissioner FDA Maharashtra

Panellists:

ARUN BHATT Consultant – Clinical Research & Development

ANISH DESAI Director IntelliMed Healthcare Solutions VISHWAS SOVANI Founder Director

DAY ONE - 05th November 2020

Founder Director Pharmawisdom

16:10 - DISCUSSION WITH EXPERTS: New drugs and clinical trial rules - Being ready for regulatory inspections

- Current key changes and challenges for trials in India
- What are the current challenges that the researchers must know before conducting a clinical trials in India? In the current scenario, how to solve these challenges?
- Current Scenario protocol and testing procedure for authorizing a new drug before it is used on a patient?
- How to be prepared on inspections and documentations during the inspections
- Discussing about the validity of clinical trial permission to initiate a clinical trial?
- Staying on top of recent regulatory updates

Moderator:

MILIND ANTANI Leader, Pharma and Healthcare Nishith Desai Associates

Panellists:

BANOTH VENKATESWARLU Assistant Drugs Inspector Drug Standard Control Organization (CDSCO)

YASMIN SHENOY

Director-Regulatory Affairs Sanofi-aventis

SRIRUPA DAS Director - Medical Affairs Abbott

JYOTSNA PATWARDHAN

Head Development QA Novartis

17:00 - Chairperson's closing remarks and end of conference

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"Topic was very good huge on current seminar, Location is very good to Aelequase, Speaking was good to deliver current situation, Very on panel discussion and due and Answer session"

Sr. CRA, Lambda Therapeutic Research

AGENDA AT A GLANCE

Features of our Virtual Conference

NETWORKING

Lobby – Here at the lobby, all attendees can see the other participants. You can choose to start a conversation privately at any time with any of the other co-participants– For more details – check out the links (YouTube videos in the last page)



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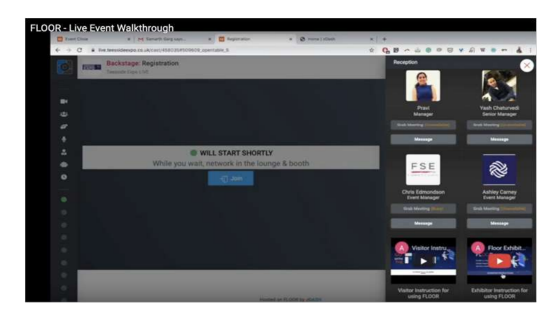
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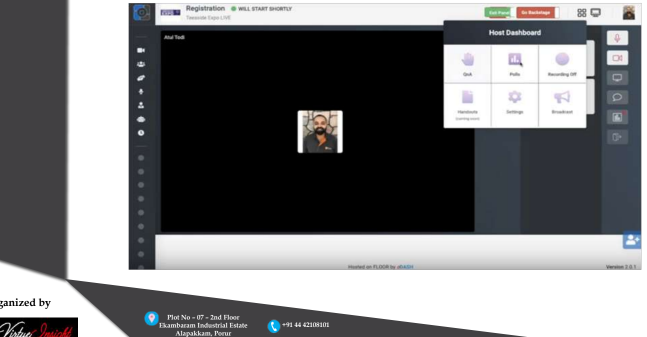
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Sr. CRA, Lambda Therapeutic Research

Reception – Should you have any questions to the organisers, you can find them at the reception - For more details - check out the links (YouTube videos in the last page)



Q&A, Polls & Handouts- We can have Q&A from the audience at the end of every session as usual and also have polls and handouts done - For more details - check out the links (YouTube videos in the last page)



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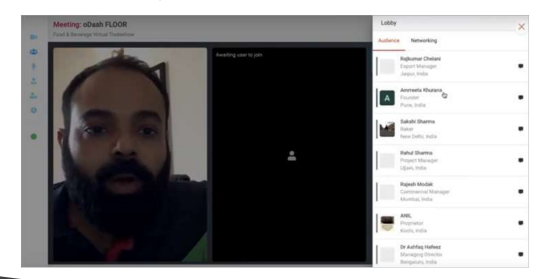
LIVE STREAMING

Solo Presentations & Panel Sessions– Interactive panel sessions and solo presentations sessions - For more details – check out the links (YouTube videos in the last page)



SPONSORS & EXHIBTORS

Exhibitors– Exhibitors have booths where they can start a conversation with any of the attendees and also attend to the attendees who visit their stall - For more details – check out the links (YouTube videos in the last page)



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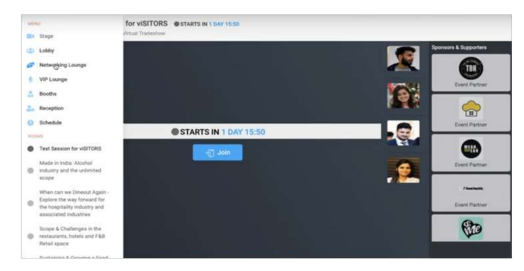
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Sponsors– Sponsors can have speaking slot sessions and their logos would be visible in all sessions for their branding purposes - For more details – check out the links (YouTube videos in the last page)



Links to YouTube videos of the conference webinar platform

Live Event Walkthrough - https://www.youtube.com/watch?v=KRX5j3gQeF0 Exhibitor Instructions - https://www.youtube.com/watch?v=uOvH46TeYrw Visitor Instructions - https://www.youtube.com/watch?v=c4WSfp9RFP0

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REGISTER ONLINE :

AGENDA AT A GLANCE

Link : https://www.bookmytrainings.com/catalogue/event/75086-11th-annual-clinical-trials-summit-2020 For Multiple Bookings - Photocopy this form and send it to bookings@virtueinsight.com

RESERVATION PRICING:

Early Bird Discount Price

Cost per delegate (Valid till 05th October 2020) -

Fee: INR 07,000 + GST(18%)

Standard Rate

Cost per delegate (Valid From 06th October 2020)-

Fee: INR 11,000 + GST(18%)

CERTIFICATION

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

Discounted Rate for Bulk Booking of More Than 5 Delegates

Please email us at bookings@virtueinsight.com

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ForenameSurname	•••
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I confirm that I have read & agree to the terms and conditions of booking..... (Please Tick)

Signature

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REGISTRATION FORM

Should you have any questions on bookings, Please feel free to contact us.

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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.

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.

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 INR 5,000

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.

Video: If you cannot attend the conference, you can still purchase the Video of the virtual conferences for INR 5,000 + Tax

Indemnity: Virtue Insight reserves the right to make alterations to the conference/executive briefing content, timing, speakers or venue without notice. The event may be postponed or cancelled due to unforeseen events beyond the control of Virtue Insight. If such a situation arises, we will reschedule the event.