DIA 2024 Singapore Annual Meeting

Cultivating Synergies in Clinical Research and the Regulatory Environment to Innovate Healthcare

> 16-17, July 2024 Voco Orchard Road, Singapore

Overview

The rapid evolution of drug development and technology, coupled with regulatory advances have created new opportunities and challenges in healthcare. It is critical for regulators, academia, patients, and industries to collaborate closely to ensure an evolving and robust regulatory ecosystem that enables accelerated and cost-effective access to innovative healthcare solutions. This year, DIA Singapore Annual Meeting aims to bring regulators, patient representatives and industries to share and discuss ways in cultivating synergies in clinical research and the regulatory environment to innovate healthcare in APAC. Come join us and be a part of the discussions to explore the challenges and actions that stakeholders in APAC can adopt to ensure transformations in the healthcare evolution.

Objective

- Receive updates from senior regulators how healthcare products are regulated, what actions/ plans they have for regulating innovative products and clinical trials, and their collaborations with key stakeholders.
- Hear the dialogues between regulators, patient representatives and industries on Clinical Research and the Product Registration topics.
- Receive the information on hot regulatory topics, such as ICH guidelines, AI, and new technologies adopted in R&D.
- ASEAN Townhall: Hear deliberation about the topics discussed at this DIA Singapore from the ASEAN countries regulators.
- Network with regulators, patient representatives, clinical development, pharmacovigilance and regulatory affairs professionals from industries.

Program Chair

- Chair. Finny Liu, MSc, RPh
- APAC Regional Regulatory Policy Lead, Roche

Co-Chair. Helene Sou, MSc, RAC Global Regulatory Policy and Innovation, Takeda Pharmaceutical Company Limited

Program Committee

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- Ellyne Setiawan, MPharm
- Head of Research & Development Quality (Asia
- Pacific), Daiichi Sankyo Singapore Pte. Ltd.

Jack Wong

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- Association (ARPA)
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- Head of Medical Affairs, APAC, BeiGene
- Thean Soo Lo, BPharm, MSc
- Regulatory Affairs Management Consultant, TS Consulting

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- APAC regional medical director, Merck

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8.00 - 8.30 am	Registration					
8.30 - 8.45 am	Opening Remarks					
8.45 am - 10.15 am	Plenary Session - Senior Regulator's Perspectives : What Should the Future Regulatory Ecosystem Look Like? In this session, senior regulators will provide an update on their current regulatory system, initiatives and plan to reflect the changing regulatory landscape. They will also share their thoughts and challenges on hot topics, such as - reliance and working toward regulatory convergence - adopting digital technologies, such as DCT, RWD/RWE, eCTD and e-labeling In the Plenary Panel Discussion, Regulators, Patient and Industry Representative will discuss and make recommendations on new ways of working to ensure that the regulatory system is efficient, sustainable and fitfor-purpose to enable faster approval of innovative healthcare products, as well as what should be considered regarding regulatory agility, alignment and harmonisation.					
Session Chair(s)						
Finny Liu, MSc, RPh APAC Regional Reg Roche, Singapore		Co-Four	im, MBA Ider and CEO D Health Research, Singapore			
8.45 – 9.00 am	Presentation 1 (PMDA)					
	Yuriko Takemura Coordinator, Division of Asia II, O	Office of International Pr	rograms, PMDA JAPAN			
9.00 - 9.45 am	Presentation 2 (MFDS)					
	Heesung Kim, Ph.D ., Director of Biologics Division National Institute of Food and Drug Safety Evaluation (NIFDS), Ministry of Food and Drug Safety (MFDS)					
9.45 - 10.15 am	Presentation 3 (BPOM)					
	Ministry of Health, Republic of In	donesia.	, Pharmaceuticals and Medical Devices od Control - Badan Pengawas Obat dan Makanan (BPOM)			
10.15 – 10.45 am	Tea / Coffee Break					
10.45 – 11.45 am	Panel Discussion					
	Industry representative: Wassim Nashabeh, Ph.D Pharma Technical Regulatory Ger	nentech	Moderator John Lim, Duke-NUS Medical School			
	Patient advocacy: Nidhi Swarup, Crohn's & Colitis Society of Singa	ipore	Panellists: Yuriko Takemura, PMDA Heesung Kim , MFDS Lucia Rizka Andalusia , BPOM			
11.45 – 12.45 pm	Lunch & Network					
12.45 – 1.45 pm	Innovation Hub					
1.45 – 5.30 pm	Session 1. Accelerating and Streamlining Regulatory Processes					
	The recent pandemic highlighted an extraordinary global collaboration among regulators. Utilizing digital tools, establishing new regulatory pathways, and implementing regulatory agility have significantly accelerated and streamlined approval processes for the benefits of patients. In this session, we will provide an update and reflect on various recent innovative regulatory initiatives and pilots, such as reliance, collaborative, and expedited programs. Industry experts will share their experiences and best practices through insightful case studies. The panel discussion with regulators will offer perspectives and explore the potential evolution of these initiatives in the future.					
Session Chair(s) Helene Sou, MSc, R/ Global Regulatory P Takeda Pharmaceuti			egulatory PolicyRegulatory Affairs Management			

1.45 – 1.55 pm	Introduction - Overview of Ways to Accelerate and Streamline I	Degulatory Dracoscos
	Helene Sou, MSc, RAC Global Regulatory Policy and Innovation, Takeda Pharmaceutical Company Limited, Singa	bore
	Session 1a Focus on New	Product Registration
Session Chair(s) Helene Sou, MSc, RAC Global Regulatory Poli Takeda Pharmaceutica		ean Soo Lo, BPharm, MSc gulatory Affairs Management Consultant, Singapore
1.55 – 2.10 pm	Industry sharing Hybrid ACCESS/ORBIS type C pathway: 1st indus	try experience, process, benefits and considerations
	Mi-Young Park, Senior Regulatory Affairs Directo	r, Growth and Emerging Markets, Takeda
2.10 – 2.25 pm	Industry sharing ASEAN Country specific requirements in new pro harmonized to achieve faster registrations	duct registrations: challenges and what can be streamlined/
	Edana Loke, Director, Regulatory Policy and Inte	ligence, Australia, China, Japan, and Asia (JAPAC), Abbvie
2.25 - 3.00 pm	Regulator's sharing SRA's documents/tools and support to enable or	facilitate reliance pathways
	Karen Loft, Indo-Pacific Regulatory Strengthenin Yuriko Takemura, PMDA JAPAN	g Program International Regulatory Branch TGA
3.00 – 3.15 pm	Panel Discussion + Q&A	
	Moderator : Helene Sou, Takeda Pharmaceutical Company Limited, Singar	Panellists : pore Regulators - Karen Loft, TGA Yuriko Takemura, PMDA JAPAN Industry - Mi-Young Park, Takeda Edana Loke, Abbvie
3.15 – 3.45 pm	Tea / Coffee Break	
	Session 1b focus on Post	-Approval Changes
Session Chair(s)		
Sannie S Foong Chong Senior Director, Global MSD International, Sing	l Regulatory Policy Gl	lene Sou, MSc, RAC obal Regulatory Policy and Innovation, keda Pharmaceutical Company Limited, Singapore
3.45 - 4.00 pm	Industry sharing Unleashing the Power of Reliance for PACs: Roch	e's Exciting Journey with 48 NRAs
	Suat Gnoh Por, International Regulatory, Roche	
4.00 - 4.20 pm	Industry sharing ASEAN Country specific requirements for post-a harmonized to achieve more efficiency in regulat	oproval changes: current challenges and what can be streamlined/ ory processes for PACs.
	Sia Lee Yoong, Global Regulatory Policy and Inte	lligence, GlaxoSmithKline Singapore Pte. Ltd
4.20 – 4.55 pm	Regulator's sharing : Recent and Ongoing Improvements and Streamli Philippines , Thailand	ning of Regulatory Processes for PACs:
	PFDA and Thai FDA	
4.55 - 5.25 pm	Panel discussion + Q&A	
	Moderator: Sannie Chong Panellists: Suat Gnoh Por , Roche, Sia Lee Yoong, Regulators ; Mei-Ling Chan , Reviewer, Section o Administration, PFDA (tbc) and Thai FDA (tbc)	GSK, Jeffrey Schnack , Accumulus Synergy New Drug Division of Medicinal Products Taiwan Food and Drug
5.25 – 5.30 pm	Closing Remarks & Day 1 End	

8.30 am - 1.00 pm

Session 2. (Parallel Session)

Drug Development and Innovation in Clinical Research.

Transformative approaches to drug development have the potential to improve efficiency in R&D, bringing new therapies and innovations to market earlier as well as provide better prediction and outcome of patient response. In this session, industry speakers will delve into the use of innovative approaches in drug development, the potential of radiopharmaceuticals in oncology trials, advances in liquid biopsy technology, integration of real-world evidence and the impact of patients' voice in accelerating drug development. Finally, in our dynamic healthcare ecosystem, patient involvement and engagement are no longer optional, they are essential drivers of safer, more effective care. We will explore how patients are at the heart of this transformative journey.

2a Session Chair(s)					
Audrey Ooi, MSc Head- Business Development Clinical Research Malaysia, Malaysia		Vicky Hsu Senior Vice President, Head of Project leadership and Biotech Operations Asia			
2b Session Chair(s)					
Senthil Sockalingam Head of Medical Affair	rs, APAC, BeiGene	Ellyne Setiawan, MPharm Head of Research & Development Quality (Asia Pacific), Daiichi Sankyo Singapore Pte. Ltd.			
8.30 – 8.50 am Innovations in the conduct of early phase		inical trials			
	Aaron Tan, Medical Oncologist, National Car	ncer Centre Singapore			
8.50 – 9.10 am	Opportunities and Challenges in Radioligand Trials in Asia				
	HV Bimba, Senior Clinical Research Medical Advisor Global Drug Development, Novartis Singapore Pte. Ltd.				
9.10 - 9.30 am	Novel functional liquid biopsy: non-invasive circulating tumor cells-derived organoids for anti-cancer drug screening and clinical monitoring				
	Shian-Jiun Shih, CEO and co-founder, Celler	ntia, Inc.			
9.30 – 9.50 am	The use of RWD in accelerating development of an indication				
	Susan Song , Director, Real World Evidence	Growth, Parexel, Singapore,			
9.50 - 10.30 am	Tea / Coffee Break				
10.30 – 11.00 am	Patient's voice in the clinical journey				
11.00 - 11.30 am	Patient's access to clinical trials: What we ca	an do differently?			
	Kate Lawrey, Director and Head of APAC Pa	tient Recruitment, IQVIA RDS East Asia, Singapore			
11.30 – 12.00 pm	Patient Concierge - Clinical trials beyond bo	rders			
12.00 – 1.00 pm	Lunch & Network				

8.30 am – 1.00 pm

Session 3. (Parallel Session) New Regulatory Fields and Trends

Regulatory Affairs has always been an exciting and evolving field, exacerbated with rapid advances in technology. In this session, we will start with exploring AI products regulatory frameworks. Through insightful case studies, we will uncover the intersection of innovation and regulation, offering valuable insights for navigating this dynamic terrain. Then, we will dive into a cloud web-based dossier technology and its fascinating impact on expediting regulatory approval processes in an era where time is of the essence. Next, we will look at creative regulatory pathways and unique strategies employed in Hong Kong and the Greater Bay Area. By embracing unconventional approaches, these regions have carved out distinctive regulatory landscapes, fostering innovation and driving economic growth. Next, we will delve into the regulatory environment for longevity products. Longevity products hold immense promise for improving health and well-being. Our discussions will shed light on the unique challenges and opportunities and essential regulatory considerations for responsibly bringing these transformative products to market. Finally, we will discuss the flexible and innovative regulatory approaches applied in continuous manufacturing of drug substances and products as outlined in the latest ICH Q13. Embracing continuous manufacturing offers unprecedented opportunities for enhancing efficiency, reducing costs, and ensuring product quality.

12.00 – 1.00 pm	Kai Yin Po, Associate Principle Scientist, F	Regulatory Affairs, MSD			
11.30 – 12.00 noon	Continuous manufacturing of Drug Substances and Drug Products: ICH Q13				
	Christine Yuan HUANG, Co-founder, Asia	Longevity Professionals Association (ALPA)			
11.00 - 11.30 am	Longevity Regulatory: how to regulate anti-aging health supplements?				
	Jack Wong, Founder, Asia Regulatory Pro	ofessionals Association (ARPA), Singapore			
10.30 - 11.00 am	Hong-Kong and Greater Bay Area (GBA) Innovative Regulatory Pathway				
9.45 - 10.30 am	Tea / Coffee Break				
	Jeffrey Schnack, Accumulus Synergy, Reg	julatory Policy Lead - Japan & Asia			
9.20 – 9.45 am	Accumulus Synergy & Regulatory Innovation in the Cloud: What Does this Mean for Asia?				
	Greg Michels, CEO, PV.app				
8.55 – 9.20 am	Industry case study: how to regulate AI pr	roducts?			
	TBD				
8.30 - 8.55 am	Regulatory framework for AI products				
Session Chair(s) Jack Wong Founder, Asia Regula Singapore	atory Professionals Association (ARPA),	Finny Liu, MSc, RPh APAC Regional Regulatory Policy Lead Roche, Singapore			

Session 4.

1.00 – 2.30 pm

ASEAN Townhall : What is the Influence of Emerging Regulatory Strategies and Trends on the Surveillance of Medicines after They Have Been Approved?

The ASEAN Town Hall is a key session of DIA Singapore that brings together ASEAN regulators and important industry players such as Clinical Research, Regulatory and Pharmacovigilance professionals. The ASEAN townhall serves as a valuable platform for discussing the current "hot topics" in the evolving regulatory landscape with a key area of focus on addressing operational issues and policy barriers that impact the healthcare sector in the ASEAN region. This collaborative approach is crucial for enhancing the overall quality and accessibility of healthcare services for our patients in need.

This year, the dialogue will revolve around how the current changes or trends in the regulatory landscape (such as risk-based reviews, RWD, use of digital tools etc.) not only bring opportunities for an accelerated development and/or approval timeline for a medicine but also may have an impact on the post-approval monitoring, and pharmacovigilance approaches. We will hear perspectives from ASEAN regulators as well as industry experts.

Speaker: **Asmaa Asim,** RA/PV Lead, South, East & Southeast Asia, Organon Asia **Muzzaffar Halli**, Senior Manager RA/PV, South East Asia, Novo Nordisk

Session Chair(s)

Thean Soo Lo, BPharm, MSc Regulatory Affairs Management Consultant, TS Consultung, Singapore Helene Sou, MSc, RAC Global Regulatory Policy and Innovation, Takeda Pharmaceutical Company Limited, Singapore

2.30 – 3.30 pm APEC Regulatory Harmonization Steering Committee (RHSC) Special Feature: Advancing Regulatory Convergence

Session Chair(s)

Sannie S Foong Chong, Ph.D. Senior Director, Global Regulatory Policy MSD International, Singapore Kum Cheun Wong, PharmD Head Asia Pacific Regulatory & Development Policy, Novartis Asia Pacific Pharmaceuticals Pte. Ltd., Singapore

Participants: Michelle Limoli, FDA / TFDA, PMDA, HSA, Thai FDA, MFDS

3.30 - 4.00 pm	Tea / Coffee Break
4.00 – 5.15 pm	Session 5. Clinical Research Regulations : Critical Aspects that Impacts Clinical Research Practices
	Dive deep into the intricacies of ICH E6 R3 at the final session of DIA Singapore, where we dissect the

updated guidelines with precision, focusing on the critical aspects that impacts clinical research practices. Explore strategies for navigating regulatory landscapes, ensuring compliance, and optimizing clinical research practices. Gain invaluable insights into how the highest standards of quality are maintained through the evolving role of technology and its advances.

Session Chair(s) Senthil Sockalingam Head of Medical Affairs, APAC, BeiGene		Ellyne Setiawan, MPharm Head of Research & Development Quality (Asia Pacific), Daiichi Sankyo Singapore Pte. Ltd.			
4.00 – 4.25 pm	5 pm ICH E6 R3 Data Governance: Moving the needle to the next level of data stewardship.				
	TBD				
4.25 - 4.50 pm	Regulatory landscape of Decentralised Clinical Trials in Asia Pacific				
	Sandy Chan, Associate Director Global Regulatory Policy & Intelligence, Johnson & Johnson				
4.50 – 5.15 pm	Panel with regulators				
	TBD				
5.15 - 5.30 pmClosing Remarks and Conference end					

2024 Singapore Annual Meeting

Event #24652 • July 16-17, 2024 | Voco Orchard Road, Singapore

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