

THE 11TH JBF SYMPOSIUM PROGRAM (as of Oct. 2019)

Date: Tue, 25th Feb. – Thu, 27th Feb. 2020 Venue: TOWER HALL FUNABORI, Tokyo, Japan

Day 1: Tuesday, 25th Feb.

12:30-12:45 P	M Opening Remarks			
12:45-14:05	Introduction of AI to bioanalysis and pharmacokinetics			
	 Isn't machine learning difficult if you think it's just a tool? Tatsuya KANEYAMA (Japan Pharmaceutical Manufacturers Association) 			
	 HPLC method development using AI: current practice and recent trend Masaaki SUZUKI (ChromSword Japan) 			
	 Approach for the prediction of ADME parameters using machine learning Koichi HANDA (Teijin Pharma) 			
12:45-14:05	Failure & Trouble Cases and these Solutions of LBA [Parallel Session]			
	 Failure & Trouble Cases and these Solutions of LBA Yohei HAYATA (Shin Nippon Biomedical Laboratories) 			
14:20-15:50	Poster Presentation Part I			
	14:20-15:20 Core-time for public offering presentations (odd number posters)			
16:00-18:00	Practical biomarker measurements in drug development			
	 Draft of points to consider document on biomarker assay validation in Japan Yoshiro SAITO (National Institute of Health Sciences) 			
	"Case study: Validation of an LC-MS method for measuring urinary glycosaminoglycans an it's evaluation as a biomarker in MPSVII" - Julie TAYLOR (Ultragenyx)			
	Biomarker assay validation - FB from EBF discussions on today's challenge of connecting science and regulations - Joanne GOODMAN (AstraZeneca)			
	 Validation of Biomarker Quantification in Japan (DG2019-44) Yoshitaka HASHIMOTO (Ono Pharmaceutical) 			



Day 2: Wednesday, 26th Feb.

9:00-10:40 AM	Aı	nti-drug antibodies in new modality drugs	
		Immunogenicity assessment of biosimilar: FKB327's case - Katsuhiko YAMAMOTO (Fujifilm Kyowa Kirin Biologics)	
	\triangleright	Immunogenicity assessment of peptide therapeutics - Mayurranjan MITRA (Genentech)	
	۶	ADA assessment for therapeutic oligonucleotides - Hideo TAKAKUSA (Daiichi Sankyo)	
		Immunogenicity assessment of gene therapies: Current and future concepts - Lydia MICHAUT (Bioagilytix Europe)	
9:00-10:40 AM	Cl	osed Session: Let's talk to each other about bioanalysis [Parallel Session]	
		Additional registration is required to attend this session.	
10:55-11:25	Po	ster Presentation Part II	
	10:5	5-11:55 Core-time for public offering presentations (even number posters)	
12:35 - 13:35 PM	Ι	Juncheon	
13:45-15:00	ICH M10 Guideline / Draft		
		ICH M10 draft guideline and future perspectives - Akiko ISHII-WATABE and Yoshiro SAITO (National Institute of Health Sciences)	
		ICH-M10: JBF Workshop Report - Proposals from JBF - Takeru YAMAGUCHI (Sumika Chemical Analysis Service)	
		Feedback from ICH M10 discussions in Europe - Philip TIMMERMAN (European Bioanalysis Forum)	
15:30-17:00	Bi	oanalysis for development of DDS drugs	
		A corporate pharmaceutical researcher's expectations for the bioanalysis technologies - Hideo KOBAYASHI (Daiichi Sankyo RD Novare)	
		Case study of bioanalysis in the development of DDS products - Masayoshi SAITO (Mitsubishi Tanabe Pharma)	
		Characterization of Nano-DDS Formulations in Drug Discovery Stage - Eiichi YAMAMOTO (National Institute of Health Sciences)	
17:10-18:10	K	eynote Lecture	
	I	Prof. Yasuhiro Matsumura (National Cancer Center)	
18:30-20:30	Ba	inquet	



Day 3: Thursday, 27th Feb.

9:00-11:00 AM Poster Presentation Part III (JBF Discussion Groups)

- > DG2019-40 Accuracy and precision criteria of method validation for bioequivalence studies
- ▶ DG2019-41 Bioanalysis of Unbound Drug
- DG2019-42 Current situations and issues of multiplex LBA
- DG2019-43 Guide to ADA Analysis: Considerations in Developing Analytical Methods and Conducting Nonclinical/Clinical Studies
- DG2019-44 Validation of Biomarker Quantification in Japan

11:15-12:15 Collaboration Session with Clinical Pharmacology How can Industry Clinical Pharmacologists and Bioanalysts Work Together?

- Expectations to Bioanalysts for Each Measurement Item Ryoko TAKUBO (Chugai Pharmaceutical)
- Analytical Issues in Clinical Studies Risa FUKUSHI (Astellas Pharma)

11:15-12:15Collaboration Session with JSQA [Parallel Session]
How to use computerised system without support data integrity validation by SOP?

Best practice of operation management under data-integrity guidelines - A case study using stand-alone HPLC - Yoshikazu MASAKI (Otsuka Pharmaceutical Factory)

12:30-13:30 PM Luncheon

13:45-15:45 Application of large molecule analytical technologies to new drug development

- Utilization of LC/MS analysis on the immunogenicity research of therapeutic antibodies -Nobuo SEKIGUCHI (Chugai Pharmaceutical)
- Biotransformation evaluation of therapeutic monoclonal antibody using LC-MS
 Kazuhiro KOBAYASHI (Kyowa Kirin)
- > Quantitative analysis of Collategene by Q-PCR Takao KOMATSUNO (Anges)
- Application of quantitative proteomic analysis in drug discovery research
 Tomohiro ANDO (Axcelead)

16:00-17:00 Metabolite evaluation in clinical development

- Application of Tiered Metabolite Quantification Reviewed by Japan Bioanalysis Forum Discussion Group - Makoto NIWA (Nippon Shinyaku)
- Investigation of metabolites and assessment of metabolite exposure in human matrices at the clinical stage Takahiro GOTO (Mitsubishi Tanabe Pharma)
- Tiered Approach for Assessment of Human Metabolites without Authentic Standards
 Satoru KOBAYASHI (Japan Tobacco)

17:00-17:15 Closing Remarks