	Module1: Introductio	n			Time	
Day 1					22-Aug-16	
Recognize the concept needs	Welcome address		0.25	Professor Dr. Kenji	09:00-09:15	D1-1
of PRD in medical and global	Objective of the course and	Looturo	0.25	Hirayama, Nagasaki	00.45 00.20	
view of health	expectation, Pre-test	Lecture	0.25	University, Japan	09:15-09:30	D1-2
	Overview of product research			Professor Dr. Juntra		
	and development and	Lecture	1.00	Karbwang, Nagasaki	09:30-10:30	D1-3
	stakeholders Break			University, Japan	1030:-10:45	
	Key medical and public health			Professor Dr. Kenji	103010.45	
	issues, and the need for new	Lecture	1.00	Hirayama, Nagasaki	10:45-11:45	D1-4
	products			University, Japan		
	Lunch				11:45-13:00	
	Module 2: Drug Disco	very and	Developn	nent		
	Session 1:	Discove	ry			
	History and overview of drug	Lecture	0.50		13:00-13:30	D1-5
	discovery process	LCOIGIC	0.00		10.00 10.00	D1 3
	The role of Genomics and bioinformatics	Lecture	0.50		13:30-14:00	D1-6
	High throughput screening:					
	Pre-requisite of HITS					
	systems, Assay Development	Lecture	0.50		14:00-14:30	D1-7
	& Validation, Biochemical &	Lecture	0.50		14.00-14.30	טו-/
	Cell-based assay, Assay					
	Readout & Detection				14:30-14:45	
	Break The role of Chemistry in Drug				14.30-14.45	
	Discovery:					
	-Lead Identification,					
Describe the pharmacological	-Lead Generation Libraries	Lecture	0.75	Dr. Nobuhiro Noro,	14:45-16:00	D1-8
process for drug discovery.	(Combinatiorial Chemistry,			GlaxoSmith Kline,		
Identify the process to protect intellectual property	Computational Approches),			K.K., Japan		
intellectual property	-Lead Optimization Pharmacokinetic Issues in					
	Drug Discovery:					
	- Designs and Purposes of					
	DMPK Profiles of Lead &					
	Drug Candidates					
	- DMPK studies in Drug Discovery (Late lead					
	identification & Early lead	Lecture	0.75		16:00-16:45	D1-9
	optimization, Mid lead					
	optimization, Late lead					
	optimization)					
	- Prediction of DMPK and PKPD Correlation - Prediction of Toxicity					
Day 2	- 1 rediction of Toxicity				23-Aug-16	
•	The role of Pharmacology					
	- Pharmacological Evaluations					
	(Selectivity screening,			Professor		
	Pharmacological profiling,	Lecture	0.75	Dr.Yoshimasa	09:00-09:45	D2-1
	Testing in animal models of		00	Tanaka, Nagasaki	15.55 55.70	<i>52</i> 1
	disease, Safety			University, Japan		
	pharmacology) - Examples					
	Session 2 : Precli	nical Dev	elopment			
				Professor Dr. Kesara		
	Overview: Pre-clinical study requiments for human clinical	Lecture	0.50	Na-Bangchang,	09:45-10:15	D2-2
	studies	Lociale	0.00	Thammasat	00.70 10.10	<i>D</i>
			Unive	University, Thailand		
Describe the process of	Assessing of Drug Safety: the role of toxicology					
pharmacological development	- Objective and Type of		1.00	Professor Dr. Kesara Na-Bangchang,		
	Toxicology	l o otrore			10.15 11.15	D0 0
	- Exploratory Toxicology	Lecture	1.00	Thammasat	10:15-11:15	D2-3
	- Regulatory Toxicology			University, Thailand		
	- Toxicity Measures and					
l	Toxicity Test			I		

	Break				11:15-11:30	
	Biopharmaceuticals - Development of Biopharmaceuticals - Type of Biopharmaceuticals - Issues Related to the use of Biopharmaceuticals (Antigenicity, Stability, Drug delivery)	Lecture	0.50	Professor Dr. Kesara Na-Bangchang, Thammasat University, Thailand	11:30-12:00	D2-4
	Lunch				12:00-13:00	
	Drug Discovery in Academia	Lecture	0.50	Professor Dr. Kesara Na-Bangchang, Thammasat University, Thailand	13:00-14:00	D2-5
	Pharmaceutical Development and CMC	Lecture	1.00	Professor Dr. Hiroshi Sasaki, Nagasaki University, Japan	14:00-15:00	D2-6
	Session 3: Clini	cal Devel	opment			
	Overview of clinical development	Lecture	1.00	Professor Dr. Juntra Karbwang, Nagasaki University, Japan	15:00-16:00	D2-7
Day3					24-Aug-16	
	Patents in Drug Discovery: - Publication VS Patents - Patent system - Type of Patent	Lecture	1.00	Professor Dr. Hiroshi Kato, Nihon University, Japan	09:00-10:00	D3-1
	Regulatory Framework	Lecture	1.00	Dr. Hiroshi Yamamoto Nagasaki University, Japan	10:00-11:00	D3-2
	Investigational Phases of Clinical Research (I-IV) and Study Design	Lecture	0.50	Professor Dr. Juntra Karbwang, Nagasaki University, Japan	11:00-11:30	D3-3
	Clinical Development Plan	Lecture	0.50	Professor Dr. Juntra Karbwang, Nagasaki University, Japan	11:30-12:00	D3-4
	Lunch				12:00-13:00	
	Session 4: 1	raditiona	l Medicin			
Underline the importance of traditional medicine in PRD	Introduction of Traditional Medicine and guidance on herbal medicines	Lecture	1.00	Professor Dr. Kiichiro Tsutani, Tokyo Ariake university of medical and health ,Japan	13:00-14:00	D3-5
	Regulation for traditional medicine development	Lecture	1.00	ProfessorDr.Ichiro Arai, Nihon Pharmaceutical University, Japan	14:00-15:00	D3-6
Day4	181				25-Aug-16	
	Pharmacogenomics - Anticipated Benefits - Polymorphisms in Drug Targets - Polymorphisms in ADME - Pharmacogenomics Testing - Pharmacogenomics in Clinical Trials - Examples Break	Lecture	1.50	Dr.Shyh-Yuh Liou, Ono Pharmaceutical Company Limited Head Office, Japan	09:00-10:30	D4-1

	Dhamasa dination in Olivian	Ι		1		
	Pharmacokinetics in Clinical Development - Extrapolation from animal to human pharmacokinetics	Lecture	0.50		10:45-11:15	D4-2
	Pharmacokinetics	Lecture	1.00	Professor Dr. Kesara Na-Bangchang, Thammasat University, Thailand	11:15-12:15	D4-3
	Lunch				12:15-13:00	
	Rev and Exam 1 (Module 2)		1.00	Profeesor Dr. Hirayama or Professor Dr. Kesara	13:00-14:00	D4-4
	Module5: Good Cli	nical Prac	ctice			
	Concept of Good Clinical Practice	Lecture	0.50	Professor Dr. Juntra Karbwang, Nagasaki University, Japan	14:00-14:30	D4-5
Discribe the concepts of GCP, Recognise the principles of Ethics in research and the	Principles of Research Ethics and Ethics codes and Guidance	Lecture	1.00	Professor Dr. Juntra Karbwang, Nagasaki University, Japan	14:30-15:30	D4-6
functions of Ethics Committee	Responsibilities: Sponsor, Monitors, audit, DSMB	Lecture	1.00	Professor Dr. Kesara Na-Bangchang, Thammasat University, Thailand	15:30-16:30	D4-7
Day5					26-Aug-16	
	Module6: Data Ma	nagemen	t Process			
Discribe the concepts of GCP, Recognise the principles of Ethics in research and the functions of Ethics Committee	Data Management Process	Lecture	0.50	Professor Dr. Kesara Na-Bangchang, Thammasat University, Thailand	09:00-09:30	D5-1
	Rev and Exam 5(Module 5 and 6)		1.00	Professor Dr. Juntra Karbwang, Nagasaki University, Japan Professor Dr. Kesara Na-Bangchang, Thammasat University, Thailand	09:30-10:30	D5-2
	Module 3: Vacci	ine Devel	opment			
	Session 1	: Discove	ry			
Describe the principles of basic immunology. Describe the	Historical overview of vaccine Discovery	Lecture	0.50	Professor Dr. Kenji Hirayama, Nagasaki	10:30-11:00	D5-3
process of vaccine discovery	Vaccines for infection control	Lecture	1.00	University, Japan	11:00-12:00	D5-4
	Lunch				12:00-13:00	
	Module 3: Vacci					
D 11 11 1	Session3:Clinic	cal Develo	pment			
Describe the process of vaccine clinical development	Assessment of pre-clinical information	Lecture	1.00		13:00-14:00	D5-5
	Clinical development plan	Lecture	0.50	Dr.Daisuke Tsuzuki	14:00-14:30	D5-6
	Dose selection and regimen	Lecture	1.00		14:30-15:30	D5-7
	Rev and Exam 3 (Module 3)		1.00	Professor Dr. Kenji Hirayama, Nagasaki University, Japan	15:30-16:30	D5-8
Day6					29-Aug-16	
	Field Trip to Hisamitsu Pharmaceutical Co.,Inc		site visit	Mr.Hideyuki Nakano,Hisamitsu Pharmaceutical Co.,Inc	10:00-17:00	D6
Day7					30-Aug-16	
	Module4: Diagno		lopment			
	Discovery and development of diagnostic tools	Lecture	0.50		09:00-09:30	D7-1
	Prototype production and assessment	Lecture	0.50		09:30-10:00	D7-2
	Scale-up, manufacture and control	Lecture	0.50		10:00-10:30	D7-3
	Break				10:30-10:45	

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GIAGHOSHC LOOIS	Development of kits	Lecture	0.50	Dr. Masato Sasaki,	10:45-11:15	D7-4
	Quality assurance/quality control: evaluation of efficacy after application	Lecture	0.50	QIAGEN, Japan	11:15-11:45	D7-5
	Lunch				11:45-13:00	
	Clinical development: validate prototype, manufacture pilot lot, initiate clinical trial	Lecture	0.50		13:00-13:30	D7-6
	Clinical development: Supply chain logistics and production, Statistical	Lecture	0.50		13:30-14:00	D7-7
	Rev and Exam 4 (Module 4)		1.00	Professor Dr. Kenji Hirayama and Professor Dr.Kesara Na-Bangchang	14:00-15:00	D7-8
Day8					31-Aug-16	
	Module 3: Vacci	ne Develo	pment			
	Session 2: Pre-Cli	nical Dev	elopment			
Describe the principles of basic immunology. Describe the process of vaccine discovery	Vaccine Platform Technology and Adjuvant Development (Dr.Arakawa)	Lecture	1.00	Dr. Takeshi Arakawa, University of the Ryukyus, Japan	09:00-10:00	D8-1
	CMC	Lecture	0.50		10:00-10:30	D8-2
Describe the process of pre- clinical development of vaccine	Immunogenicity and protect activity assessment	Lecture	0.50	D. Talaski Asalas	10:30-11:00	D8-3
	Break			Dr. Takeshi Arakawa, University of the	11:00-11:15	
	Safety assessment:Toxicity test in animals: regional complications, systemic toxicity	Lecture	0.75	Ryukyus, Japan	11:15-12:00	D8-4
	Lunch				12:00-13:00	
	Case studies	Group Work	2.00	Professor Dr. Juntra Karbwang, Nagasaki University, Japan Professor Dr. Kesara Na-Bangchang, Thammasat University, Thailand	13:00-15:00	D8-5
Day 9					1-Sep-16	
	Module 7: Post-registration	Activities				
	Overview	Lecture	0.75		09:00-09:45	D9-1
Describe poet registration	How to solve Access to Medicines (ATM) problems in developing countries	Lecture/ Discussi on	0.50	Professor Chitr Sitthi- amorn, Chulalongkorn	09:45-10:15	D9-2
Describe post-registration activities for medicinal products	Improving the quality of now	Lecture	1.00		10:15-11:15	D9-3
	Post-marketing product	Lecture	1.00	University, Bangkok, Thailand	11:15-12:15	D9-4
	vigilance					
	Lunch				12:15-13:00	
Describe post-registration activities for medicinal products	Stakeholders to be involved in	Lecture/ Discussi on	1.00		12:15-13:00 13:00-14:00	D9-5
Describe post-registration activities for medicinal products	Lunch Stakeholders to be involved in making product development work for the intended	Discussi	1.00	Professor Dr. Kenji Hirayama, Nagasaki University, Japan		D9-5
	Stakeholders to be involved in making product development work for the intended beneficiaries	Discussi		· · · · · · · · · · · · · · · · · · ·	13:00-14:00	

CLOSING CEREMONY 11:00-12:00 D10-2