

## กำหนดการฝึกอบรม

## เรื่อง “การปฏิบัติการวิจัยทางคลินิกที่ดี รุ่นที่ 1”

วันที่ 20-21 เดือน มีนาคม 2562 เวลา 08:30 – 16:30 น.

ห้องท่านผู้หญิงวิริยา ชั้น 5 ตึกสิริกิติ์

วันที่ 20 เดือน มีนาคม 2562

Time	Topic	Speaker
08:30 – 09:00	Register & Opening	
09:00 – 10:30	Overview of Ethical Principles in Human Research and Introduction to GCP <ul style="list-style-type: none"> <li>Development of Ethical Principles for Human Research and Good Clinical Practice (GCP)</li> <li>The International Conference on Harmonization Guidelines for Good Clinical Practice of (ICH GCP)</li> <li>Overview of ICH-GCP</li> <li>Clinical Trial Related Laws &amp; Regulations</li> </ul> Disclosure of Clinical Trial	Pravich Tanyasittisuntorn, M.D.
10:30 – 10:45	Coffee break	
10:45 – 12:00	Introduction to the principles of ethics in human research and GCP (continued)	Pravich Tanyasittisuntorn, M.D.
12:00 – 13:00	Lunch break	
13:00 – 14:45	Training on GCP: Role & Responsibilities <ul style="list-style-type: none"> <li>Role &amp; Responsibilities</li> <li>Investigator Responsibilities</li> <li>Communication with IRB/IEC</li> <li>Compliance with Protocol</li> <li>Sponsor Responsibilities</li> <li>Responsibilities of the Sponsor</li> </ul>	Pravich Tanyasittisuntorn, M.D.
14:45 – 15:00	Coffee break	
15:00 – 15:30	Institutional review board (IRB)/independent ethics committee (IEC) <ul style="list-style-type: none"> <li>Composition and responsibilities of IRB/IEC</li> <li>Application for IRB/IEC review and approval</li> <li>Review process: exemption, expedited and full review</li> </ul>	Nitaya Jeanpan
Time	Topic	Speaker

15:30 – 16:30	<p>Lecture: Informed consent</p> <ul style="list-style-type: none"> <li>• Definition and objectives of informed consent</li> <li>• Informed consent form/patient information sheet: essential elements of information and subjects' comprehension</li> <li>• Conduct of informed consent <ul style="list-style-type: none"> <li>○ Investigator's responsibilities</li> <li>○ Documentation</li> <li>○ Definition of impartial witness and legally acceptable representative</li> </ul> </li> <li>• Consent renewal</li> <li>• Informed consent in vulnerable subjects</li> </ul> <p>Workshop: Informed consent</p>	Nitaya Jeanpan
16:30 – 16:40	Q & A	

Time	Topic	Speaker
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Time	Topic	Speaker
08:45 – 09:00	Register	
09:00 – 10:15	Lecture: Safety reporting <ul style="list-style-type: none"> <li>• Purposes of safety reporting</li> <li>• Safety report terms: adverse event (AE), adverse drug reaction (ADR), severity &amp; seriousness, expectedness &amp; unexpectedness</li> <li>• Reporting process, timelines, and documentation               <ul style="list-style-type: none"> <li>◦ Expedited report</li> <li>◦ Non-expedited report</li> </ul> </li> <li>• Responsibilities of investigator/site staff, sponsor, subjects and IRB/IEC in safety reporting</li> <li>• Data safety monitoring board</li> </ul>	Nitaya Jeanpan
10:15 – 10:30	Coffee break	
10:30 – 11:15	Investigational drug handling <ul style="list-style-type: none"> <li>• Drug label</li> <li>• Drug transportation</li> <li>• Drug storage: storage condition and access control</li> <li>• Drug accountability and documentation</li> <li>• Drug destruction and documentation</li> </ul>	Nitaya Jeanpan
11:15 – 12:00	ICH GCP: Essential Documents <ul style="list-style-type: none"> <li>• Protocol</li> <li>• Case Report Form (CRF)</li> <li>• Source data &amp; Source document</li> <li>• Investigator's Brochure (คู่มือผู้วิจัย)</li> <li>• Informed consent form</li> <li>• Other Essential Documents</li> <li>• Filing And Maintaining Eessential Documents</li> </ul>	Pravich Tanyasittisuntorn, M.D.
12:00 – 13:00	Lunch break	
13:00 – 13:30	<ul style="list-style-type: none"> <li>• Data collection and data management</li> <li>• Informed Consent Process การให้ความยินยอม (หลังได้รับทราบข้อมูล)</li> </ul>	Pravich Tanyasittisuntorn, M.D.
13.30 – 14.00	Subject recruitment, subject retention, and subject compliance <ul style="list-style-type: none"> <li>• Subject recruitment process and investigator's responsibilities</li> <li>• How to develop and implement recruitment plan</li> <li>• Randomization and blinding process</li> <li>• Subject compliance and impacts of the non-compliance</li> <li>• Impacts of subject loss to follow-up and how to retain subjects</li> </ul>	Pravich Tanyasittisuntorn, M.D.

Time	Topic	Speaker
14:00 – 14:15	Coffee break	
14:15 – 15:45	Quality control and quality assurance in clinical trial <ul style="list-style-type: none"> <li>• Definition and purposes of QC &amp; QA</li> <li>• What difference between audit &amp; inspections is</li> <li>• Overview of monitor's responsibilities and monitoring activities</li> <li>• Audit/inspection process and how to respond to /inspection findings/observations</li> <li>• Common audit findings/observations</li> </ul>	Pravich Tanyasittisuntorn, M.D.
15:45 – 16:30	Test	Pravich Tanyasittisuntorn, M.D.
16:30 – 16:40	Q & A	