

CATEGORY	Subcategory	SERVICE NUMBER	PROVIDER	TITLE
<b>Training</b>	<b>GCP and Ethics Committe</b>	F1.1	Clinical Trials Centre, University of Hong Kong	Online Training on research ethics and GCP TRREE
		P1.1	Medical Center Freiburg, Clinical Trials Unit Freiburg	Training course on GCP for Investigators (1.5 days)
		P1.2	Medical Center Freiburg, Clinical Trials Unit Freiburg	Training course on GCP for Principal Investigators (3 days)
		P1.3	Medical Center Freiburg, Clinical Trials Unit Freiburg	Training course on GCP for Study Nurses (3x5 days)
		P1.4	Medical Center Freiburg, Clinical Trials Unit Freiburg	GCP Refresher (0.5 days)
		P1.6	Clinical Trials Centre, University of Hong Kong	Training course on GCP for Clinical Investigators, Study Coordinators and other Study Site Personnel
		P1.7	Clinical Trials Centre, University of Hong Kong	Training course for research ethics committee (REC) members and administrators
		P1.8	Clinical Trials Centre, University of Hong Kong	Training course for Clinical Research Coordinators (CRC)
		P1.11	Istanbul University Center of Excellence for Clinical Research	Training Course on GCP for Study Team
		P1.13	Istanbul University Center of Excellence for Clinical Research	Extended GCP Course to EC/IRB Members
		P1.14	Study Centre Munich, Academic Clinical Research Organisation, TUM School of Medicine	Training Course on GCP (AMG/MPG)
		P1.15	Clinical Trials Center (CTC) University Zurich	Training Course on GCP for Study Team
		P1.16	Clinical Trials Center (CTC) University Zurich	Training Course for CTU Staff (CTU Head, CTU Study Coordinator, CTU Administrator), 2 days

	<b>Medical Devices</b>	P1.5	Medical Center Freiburg, Clinical Trials Unit Freiburg	Medical Device Course (0.5 days)
	<b>Statistics</b>	P1.9	Clinical Trials Centre, University of Hong Kong	Training Course on Basic Statistical Analysis using SPSS for Clinical Investigators, Study Coordinators and other Study Site Personnel
	<b>Management</b>	P1.10	Clinical Trials Centre, University of Hong Kong	Training Course on Planning and Management of Investigator-Initiated Studies (IISs) for Clinical Investigator, Role of Investigators in IISs, Clinical Research Design and Framework for build-up of a CTU
<b>Research infrastructure</b>	<b>Build-up of a CTC/CTU</b>	F2.1	Cambridge Clinical Trials Unit (CCTU)	
		P2.1	Clinical Trials Centre, University of Hong Kong	Consultation / Expert Advice on Build-up of Clinical Trials Center
		P2.4	Istanbul University Center of Excellence for Clinical Research	Build-up of Clinical Trial Unit (CTU) & Consulting /Expert Advice on Build-up of CTU Infrastructure
		P1.12	Istanbul University Center of Excellence for Clinical Research	Training Course for CTU Staff (CTU Head, CTU Study Coordinator, CTU Administrator)
		F2.7	Clinical Trials Center (CTC) University Zurich	Build-up of Clinical Trial Unit (CTU)
		P2.6	Clinical Trials Center (CTC) University Zurich	Consulting / Expert Advice on Build-up of CTU Infrastructure
		P1.16	Clinical Trials Center (CTC) University Zurich	Training Course for CTU Staff (CTU Head, CTU Study Coordinator, CTU Administrator), 2 days
	<b>Build-up of a Phase I Unit</b>	F2.9	Clinical Trials Center (CTC) University Zurich	Build-up of Phase I Unit
		P2.7	Clinical Trials Center (CTC) University Zurich	Consulting / Expert Advice on Build-up of Phase I Unit Infrastructure
		P2.2	Clinical Trials Centre, University of Hong Kong	Consultation / Expert Advice on Build-up of Phase I Unit Infrastructure
<b>CTC/CTU Staffing</b>	F2.2	Cambridge Clinical Trials Unit (CCTU)	CTU staffing	

		F2.8	Clinical Trials Center CTC University Zurich	Staffing of CTU
	<b>Consortium / networks</b>	F2.10	China Medical University Hospital (CMUH)	Introduction of Taiwan Clinical Trial Consortium for Stroke
<b>Trial Management and Operations</b>	<b>IT solutions for trial management</b>	F3.1	Cambridge Clinical Trials Unit (CCTU)	Trial Management IT solutions
		F3.2	Study Centre Munich, Academic Clinical Research Organisation, TUM School of Medicine	Clinical Trial Management Tool (CTMS)
	<b>Trial planning and execution</b>	P3.1	Medical Center Freiburg, Clinical Trials Unit Freiburg	Project Coordination, Biometry, Monitoring, Data Management, Pharmacovigilance
		P3.2	Clinical Trials Centre, University of Hong Kong	Protocol Development for Investigator-Initiated Studies / Industry-Sponsored Studies
		P3.3	Clinical Trials Centre, University of Hong Kong	Study Monitoring for Investigator-Initiated Studies / Industry-Sponsored Studies
		P3.4	Clinical Trials Centre, University of Hong Kong	Data Management for Investigator-Initiated Studies / Industry-Sponsored Studies
		P3.5	Clinical Trials Centre, University of Hong Kong	Statistical Analysis for Investigator-Initiated Studies / Industry-Sponsored Studies
		P3.6	Istanbul University Center of Excellence for Clinical Research	Training, SOPs and Tools for Study Approval Management
		P3.7	Istanbul University Center of Excellence for Clinical Research	Know-how Transfer for Study Protocol Review
		P3.11	Study Centre Munich, Academic Clinical Research Organisation, TUM School of Medicine	CT (budget) Planning, Feasibility; CSP, CSR
P3.14	Istanbul University Center of Excellence for Clinical Research	Project Management Consulting Service for Clinical Trials in Turkey & MENA Region		
P3.17	Hannover Clinical Trials Center (HCTC)	Consulting Service on Sponsor's Responsibility by a Clinical Trials Unit		

		P3.18	China Medical University Hospital (CMUH)	Clinical Trial Feasibility and Study Site Selection
		P3.19	China Medical University Hospital (CMUH)	Monitoring Services in Taiwan
		P3.20	China Medical University Hospital (CMUH)	Study Project Management in Taiwan
		P3.23	China Medical University Hospital (CMUH)	Protocol/ICF development
	<b>Drug safety</b>	P3.16	Medical Center Freiburg, Clinical Trials Unit Freiburg	Drug Safety (premium version)
<b>Quality Management</b>	<b>Quality Assurance</b>	F4.1	Clinical Trials Center (CTC) University Zurich	Quality Assurance (QA): Preparation for Inspection Service
		P4.4	Clinical Trials Center (CTC) University Zurich	Quality Assurance (QA) System Audit Service
		P4.2	Clinical Trials Centre, University of Hong Kong	Study Site Audits
		P4.3	Study Centre Munich, Academic Clinical Research Organisation, TUM School of Medicine	Audit / Inspection Preparation
<b>Supporting tools and services</b>	<b>Randomization tools</b>	F5.2	Study Centre Munich, Academic Clinical Research Organisation, TUM School of Medicine	Randomization Tool
	<b>eCRF, data management and analysis</b>	F5.5	Clinical Trials Center CTC University Zurich	SecuTrial
		P5.7	Clinical Trials Center CTC University Zurich	SecuTrial
		P5.2	Medical Center Freiburg, Clinical Trials Unit Freiburg	Remote Data Entry System for Observational Studies within Europe
		P5.8	China Medical University Hospital (CMUH)	Data Management

		P5.9	China Medical University Hospital (CMUH)	Statistical Data Analysis
		F3.2	Study Centre Munich, Academic Clinical Research Organisation, TUM School of Medicine	Clinical Trial Management Tool (CTMS)
	<b>patient recruitment</b>	F5.4	Clinical Trials Center (CTC) University Zurich	Protocol Optimization for academic trials (IITs) with Patient Recruitment System (PRS)
		P5.5	Clinical Trials Center (CTC) University Zurich	Patient Recruitment System (PRS)
<b>Local Standards and regulations</b>	<b>Europe / Germany</b>	F6.1	Medical Center Freiburg, Clinical Trials Unit Freiburg	Checklists for Trial Application
		F6.2	Medical Center Freiburg, Clinical Trials Unit Freiburg	Consulting for Study design
		F6.3	Medical Center Freiburg, Clinical Trials Unit Freiburg	Consulting for Grant Possibilities
		P6.4	Study Centre Munich, Academic Clinical Research Organisation, TUM School of Medicine	Regulatory Advice and Submissions in Germany
		P6.5	Study Centre Munich, Academic Clinical Research Organisation, TUM School of Medicine	Assistance in Applications for Grants / Federal / State Support Funding Proposals
		P6.7	Hannover Clinical Trials Center (HCTC)	Training / Knowledge Transfer on Local Regulations
		<b>Asia / Japna, Taiwan, China</b>	F6.5	China Medical University Hospital (CMUH)
	P6.8		China Medical University Hospital (CMUH)	Regulatory submission in Taiwan
	P6.9		China Medical University Hospital (CMUH)	IRB/EC Submission in Taiwan
	<b>Turkey</b>	P6.1	Istanbul University Center of Excellence for Clinical Research	Process and Systems for CTUs for Study Submission

		P6.2	Istanbul University Center of Excellence for Clinical Research	Training / Knowledge Transfer for CTUs, P.I.s / Investigators and Study Teams on Local Regulations
		P6.3	Istanbul University Center of Excellence for Clinical Research	Local Regulations regarding EC / IRB Requirements & Processes