

## FACILITY NAME & ADDRESS

Facility Name	Facility Type	Facility Address
Beppu Medical Center	Hospital or Medical Center	1473 Uchikamado Ooaza, Beppu, Oita, Japan, 874-0011

### FACILITY CONTACTS

Primary FPM?	Name	Email Address	Roles
Yes	Yamashita, Manato	yamashita.manato.pg@mail.hosp.go.jp	Facility Profile Manager

### THERAPEUTIC AREAS & PATIENT POPULATION

Therapeutic Area(s)			
Therapeutic Area	Sub-Therapeutic Area		
Allergy			
Anesthesia			
Bone			
Chemically-induced Disorders			
Digestive System Diseases			
Endocrine System Diseases			
Eye Diseases			
Female Urogenital Diseases and Pregnancy Complications			
Fertility			
General Surgery			
Hemic and Lymphatic Diseases			
Immune System Diseases			
Infectious Diseases			
Internal Medicine			
Male Urogenital Diseases			
Mental disorders			
Musculoskeletal Diseases			
Neoplasms			
Nephrology			
Nutritional and Metabolic Diseases			
Oncology			



Therapeutic Area	Sub-Therapeutic Area		
Orthopedics			
Otorhinolaryngologic Diseases			
Pain			
Parasitic Diseases			
Pathological Conditions, Signs and Symptoms			
Pediatrics			
Respiratory Tract Diseases			
Skin and Connective Tissue Diseases			
Stomatognathic Diseases			
Vaccines			
Virus Diseases			
Women's Health			
Wounds and Injuries			
Bacterial Infections and Mycoses			
Nervous System Diseases			
Cardiovascular Diseases			
Congenital, Hereditary, and Neonatal Diseases and Abnormalities			
Other Areas of Expertise			
Study Phase Capabilities			
Phase II; Phase IV			
Other Facility Details Do you have Affiliated Research Sites or Satellite Sites/Clinics? A Satellite Site is a secondary location where the investigator sees No			
clinical trial subjects, usually this is the same investigator who sees subjects at the primary site location.			
What study types does your Facility have experience with?		Industry; Investigator Initiated	
Is your Facility affiliated with a government agency or part of a government funded health service?		No	
Patient Population			
Patient Population Demographics		Pediatrics - Less than or equal to 17; Adults - Ages 18- 64; Geriatrics - Greater than or equal to 65	
Patient Population Comments			



### **IRB/ERB/ETHICS COMMITTEE**

General Questions	
What is the average time (in days) to start a study once you have received the regulatory package?	30-60
Does your facility perform IRB/ERB/Ethics Committee submissions?	Yes
Does your facility have a dedicated department or group to perform IRB/ERB/Ethics Committee submissions?	Yes
Department Contact Name	Clinical
Department Contact Phone Number	+81-977-67-1111
Department Contact Email Address	yamashita.manato.pg@mail.hosp.go.jp
Is your facility able to initiate study activities prior to IRB/ERB/Ethics Committee protocol approval?	Yes
What types of IRB/ERB/Ethics Committee does your Facility use?	Central Acting as Local; Local
Does your institution and/or local regulation mandate the distribution of safety reports [e.g., Development SafetyUpdate Report (DSUR), suspected unexpected serious adverse reaction (SUSAR)] to a local Review only IRB/ERB/Ethics Committee?	Yes
Are there any other steps that the Sponsor should be aware of for your IRB/ERB/Ethics Committee review and submission?	Yes
Other Steps Explain	IRB documents should be submitted at least 2 week prior to the IRB.The approval form is provided to 3 day after the IRB.

### LOCAL IRB/ERB/ETHICS COMMITTEE

Local IRB/ERB/Ethics Committee: National Hospital Organization BEPPU Medical Center Institutional Review Board		
IRB/ERB/Ethics Committee Name	National Hospital Organization BEPPU Medical Center	
	Institutional Review Board	
Address	1473 ooazauchikamado, Beppu, Oita, Japan, 874-0011	
What is the meeting frequency of the IRB/ERB/Ethics Committee?	Monthly	
How long before IRB/ERB/Ethics review is the Submission Packet required?	2 weeks	
Does the IRB/ERB/Ethics Committee require payment prior to release of final approval documents?	No	

Registration#		Registering Body	
NA		NA	
LOCAL IRB/ERB/ETHICS COMMITTEE ATTACHMENTS			
Document Type	Document Name	Description	

No Records

# OTHER REVIEW BOARDS

Does your facility have Other Review Boards that need to approve the study prior to IRB/ ERB/Ethics Committee submission?	)r
example, scientific, radiation safety committees, or others.	

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### Local Lab

Is your Facility using a Local Lab?		Yes
Local Lab: Clinical Laboratory Department		
Lab Name		Clinical Laboratory Department
Lab Contact First Name		Manato
Lab Contact Last Name		Yamashita
Address		1473 ooazauchikamado, Beppu, Oita, Japan, 874-0011
Phone Number		+81-977-67-1111
Fax Number		+81-977-67-6267
Email Address		yamashita.manato.pg@mail.hosp.go.jp
Local Lab Accreditation		Others
		Japanese Association of Medical TechnologistsJananese Committee for Clinical Laboratory Standard
Additional Questions		
Does your Facility have a SOP/written procedure for documer	nting bio-specimen (Sample) processing steps/chain of custody	? No
Do your written procedures ensures that study-specific tempe staff to ensure compliance?	rature bio-specimen storage requirements are known to respon	sible
What is the system or tool that the site currently has or utilizes to document Bio-specimen (Sample) Processing Steps/ Chain of Custody?		of Prefer to use Sponsor provided System/ Tools
Please indicate tissue collection and processing capabilities at your site?		On site collection and Processing
Does your facility has established processes to oversee staff compliance with study-specific lab manual instructions for bio- specimen processing?		No
What are your Facility's capabilities for tissue collection and/o	r processing (embedding)?	
Are LOINC codes available for the Local Lab? (If Yes, you can upload the relevant LOINC list as an attachment in Lab Documentation)		No
Attachments		
Document Type	Document Name	Description
No Records		

### CONSENT & TRAINING

Consent	
Does your Facility have a written SOP/Policy/Procedure for: Informed Consent?	Yes
Does your Facility have a written SOP/Policy/Procedure for: Minor Assent for Pediatric Populations?	Yes
Does your Facility have a written SOP/Policy/Procedure for: Other Vulnerable Populations?	Yes
Will your Facility require language translations for consents?	No
Select the required languages	
If located in the US, has your Facility used or are you able to use the informed consent short form?	Not Applicable



Training	
Does your Facility have a training program for the research staff?	Yes
Does the course content include GCP?	Yes
Does your Facility use an external program to conduct research training?	Yes
Please provide program course name.	eAPRIN
Do you have a process or program in place to retrain research staff when a protocol is amended?	No
Does the study staff that prepares or transports dangerous goods have training that meets the IATA International Air Transport Association (US) or other countries hazardous training requirements for shipping dangerous goods?	No
FACILITY & EQUIPMENT	
Facility Capabilities	
Can your Facility support patient visits on weekends?	Yes
Can your Facility support in-patient admissions for research studies?	Yes
Does your study staff have sufficient English knowledge to understand communications in English?	No
Does your Facility have access to translators and translation support for trial conduct (e.g. consent, trial specific instruction)?	Yes
Does the Facility have storage space for Study-Related materials (e.g. Lab Kits, Patient Materials, etc.)?	Yes
Is the lab kit storage space able to support early phase studies which may require an increased number of kits?	Yes
Does your Facility have the ability to collect and store PK/PD specimens?	Yes
Does your Facility have the ability to collect PK/PD samples beyond normal business hours?	Yes
Does your Facility typically allow the collection of Pharmacogenomic (PGX) samples for research purposes?	Yes
Equipment	
Identify the Diagnostic Equipment available at or near the Facility to support Research studies?	Computerized Tomography Scan; Dual-Energy X-ray Absorptiometry or Bone Densitometry; Magnetic Resonance Imaging; Fluoroscopy; X-Radiation; Magnetic Resonance Angiography; Mammography; Nuclear Medicine (e.g.Bone scan,Thyroid scan,Thallium cardiac stress test); Electrocardiogram
General Equipment	
Does your Facility have an SOP or process that ensures routine calibration and maintenancof general equipment? Examples of general equipment include: scale, pulse oximeter, stadiometer, sphymomanomer, etc.?	Yes
Does your Facility have the necessary equipment to treat medical emergencies (ie. code cart)?	Yes
Equipment Available At The Facility To Support Research Studies	
Identify the equipment available at the Facility to support Research studies?	Refrigerated Centrifuge; Centrifuge; Refrigerator (2 to 8 Degrees C); Freezer (-20 to -30 Degrees C); Freezer (- 70 to -80 Degrees C)
Equipment Capabilities: Refrigerator (2 to 8 Degrees C)	
Do you have the ability to generate a temperature monitoring log for this equipment?	Yes
Does this equipment provide Min/Max Temperature Monitoring?	Yes
How frequently can temperature measurement occur? Check the most frequent measurement your equipment can support.	Daily
Does this equipment have back-up power?	Yes
Does this equipment have a temperature alarm?	Yes
	No

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log for this equipment?	Yes		
g?	No		
the most frequent measurement your equipment can support.	Daily		
	Yes		
	Yes		
nent?	No		
search studies?	Yes		
What type of computer operating system(s) does your institution use to support studies?			
What type of internet access does your Facility have?			
Does your Facility limit or prohibit access and use of external web-based tools or sites for clinical research? (e.g. web portals to submit documents to sponsors or CROs)			
Does the Facility have access to local IT support?			
Does your Facility prohibit the use of an external USB device (e.g. to download and send data from a temperature monitoring device)?			
Business Continuity Plan			
Does your Facility have Business Continuity Plan (BCP) to protect essential business operations which describes how those processes will be performed during a crisis at your Facility?			
Attachments			
Document Name	Description		
	web-based tools or sites for clinical research? (e.g. web portals (e.g. to download and send data from a temperature monitoring otect essential business operations which describes how those		

# INVESTIGATIONAL PRODUCT & CONTROLLED SUBSTANCES

Investigational Product Shippir	ng Details				
IP Recipient Name	Address	Email Address	Phone Number		Fax Number
Beppu Medical Center	1473 ooazauchikamado, Beppu, Oita, Japan, 874-0011	yamashita.manato.pg@mail.hosp.g o.jp			
Investigational Product Storage	e Location				
IP Recipient Name	Address	Email Address	Phone Number		Fax Number
Department of pharmacy	1473 ooazauchikamado, Beppu, Oita, Japan, 874-0011	yamashita.manato.pg@mail.hosp.g o.jp	0977-67-1111		0977-67-6267
Investigational Product Storage	e Equipment				
Identify the Investigational Product Storage Equipment at your Facility			Refrigerator (2 to 8 Degrees C)		
Equipment Capabilities: Refrig	erator (2 to 8 Degrees C)				
Do you have the ability to generate a temperature monitoring log for this equipment?				Yes	
Does this equipment provide Min/Max Temperature Monitoring?			Yes		
How frequently can temperature measurement occur? Check the most frequent measurement your equipment can support.			Daily		
Does this equipment have back-up power?			Yes		
Does this equipment have a temperature alarm?			Yes		
Do you have an SOP which supports calibration of this equipment?			Yes		

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Investigational Product Storage And Handling	
Is the Investigational Product Storage Room secured with controlled access?	Yes
Do you have the ability to generate a temperature monitoring log for this Investigational Product Storage Room?	Yes
Does the Investigational Product Storage Room provide Min/Max temperature monitoring?	Yes
Does the Investigational Product Storage Room have back-up power?	Yes
Does the Investigational Product Storage Room have a temperature alarm?	Yes
Do you have an SOP which supports calibration of this equipment?	Yes
Does your Facility have the ability to manage on-site or off-site destruction of Investigational Product?	Yes
Does your Facility have a written SOP/Policy/Procedure for destruction of Investigational Product?	No
Do you provide your Satellite Site(s) with a dedicated inventory of Investigational Product?	No
Does your Facility have a written SOP/Policy/Procedure to ensure that Investigational Product is appropriately maintained during	Not Applicable
transportation to Satellite Site(s)?	
Describe additional Investigational Product Storage & Handling Capabilities	
Preparation and Administration Of Investigational Product	
Identify the Investigational Draduct proparation conchilition at your Eacility	Extemporaneous Preparation; Vertical laminar flow
Identify the Investigational Product preparation capabilities at your Facility	
	hood (chemo/hazardous drugs)
Is your Facility capable of administering infusions?	
	hood (chemo/hazardous drugs)
Is your Facility capable of administering infusions?	hood (chemo/hazardous drugs) Yes
Is your Facility capable of administering infusions? Is your Facility adequately staffed to support studies with both blinded and un-blinded Investigational Product?	hood (chemo/hazardous drugs) Yes
Is your Facility capable of administering infusions? Is your Facility adequately staffed to support studies with both blinded and un-blinded Investigational Product? Controlled Substances	hood (chemo/hazardous drugs) Yes Yes
Is your Facility capable of administering infusions? Is your Facility adequately staffed to support studies with both blinded and un-blinded Investigational Product? Controlled Substances Does the Facility have the required licenses or registrations to receive, store, dispense and return controlled substances as	hood (chemo/hazardous drugs) Yes Yes
Is your Facility capable of administering infusions? Is your Facility adequately staffed to support studies with both blinded and un-blinded Investigational Product? Controlled Substances Does the Facility have the required licenses or registrations to receive, store, dispense and return controlled substances as required by local law?	hood (chemo/hazardous drugs) Yes Yes Yes
Is your Facility capable of administering infusions? Is your Facility adequately staffed to support studies with both blinded and un-blinded Investigational Product? Controlled Substances Does the Facility have the required licenses or registrations to receive, store, dispense and return controlled substances as required by local law? Is the storage area for controlled substances securely constructed with restricted access in accordance with local law?	hood (chemo/hazardous drugs)   Yes   Yes   Yes

# Attachments Document Type Document Name Description

# No Records

# SOURCE DOCUMENTATION

Source Documents	
What type of source documents will be used?	Paper; Electronic
Does your Facility have secure storage for patient records?	Yes
Does your Facility have patient record archiving on-site?	Yes
Provide Location name and address of any offsite archives	
What type of investigator site file/regulatory binder used (select all that apply)	Paper
What investigator site file (eISF) / eRegulatory system do you use?	
Are monitors able to access eISF/eReg while off-site?	
Please list any access limitations/ requirements for eISF/eReg	



Electronic Medical Records (EMR)/Electronic Health Records (EHR)	
Do you have Electronic Health Records (EHR)/ Electronic Medical Records (EMR)?	Yes
What EMR/EHR system do you use?	Other
For Facilities with satellite sites, where is the monitor required to access source documents?	Main Facility Only
Please list any access limitations/requirements for the Electronic Medical Records.	The Electronic Medical System permits to create a "read-only" user account for CRA which limit to review only study subject's data.
Do you work with a vendor that can electronically exchange data for clinical research from the EHR/EMR?	No
Do you have institutional approval to export data from the EHR/EMR for the clinical research?	
Are monitors able to access EHR/EMR while off site?	No
Does your facility require Sponsor representative to sign any local form (paper or electronic) for access, or any other purpose?	
Monitoring	
Check all equipment that will be available to Monitors:	Phone; Copy Machines; Internet Access
What Electronic Data Capture (EDC) systems has your staff used for clinical trials?	Oracle Inform; Medidata Rave
Does your site/institution and/or local regulations allow remote source data verification of study participant data to support remote monitoring?	No
Which of the following capabilities are available to support remote source data verification? (Check all that apply)	

## ADDITIONAL LOCATIONS

### Additional Locations Add any addresses you wish to be available in the Study Site Profile. These addresses will be available for selection in the following sections of the Study Site Profile -Additional Study Locations - These addresses can be added to your FDA Form 1572, if applicable.

Location Name	Contact Name	Address	Phone Number	Fax Number	E-mail Address
No Records					

### ADDITIONAL INFORMATION & ATTACHMENTS

#### Additional Information

Please provide additional information not captured in other sections of the Facility Profile that you feel is important for Sponsors to know about your Facility. Please reference the section name if applicable.

Facility Attachments		
Document Type	Document Name	Description
No Records		

### ORGANIZATION AFFILIATIONS

Organization Affiliations				
The Organization (s) that requested Affiliation with your Facility/Department are listed below with Affiliation Status				
Organization Name and Address	Organization Affiliation Type	Organization Affiliation Status	Status Date	
No Records				