

Note: Invalid phone numbers and email address if entered in text fields in the form shall not be populated in SIP. **Facility Name** National Hospital Organization Mie Chuo Medical Center THERAPEUTIC AREAS AND PATIENT POPULATION THERAPEUTIC AREA(S) Provide the list of Therapeutic Areas for your Facility: Cardiovascular Diseases **Digestive System Diseases Endocrine System Diseases** Female Urogenital Diseases and Pregnancy Complications Internal Medicine Musculoskeletal Diseases Nervous System Diseases Oncology Pediatrics Respiratory Tract Diseases Sub-Therapeutic Areas: Note: Sub-Therapeutic Areas can be selected online from the Facility Profile in SIP. Other Areas of Expertise: Male Urogenital Diseases, Otorhinolaryngologic Diseases, Wounds and Injuries, Stomatognathic Diseases STUDY PHASE CAPABILITIES Phase I ✓ Phase II ✓ Phase III ✓ 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 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? Academic Industry Investigator Government Other Other Initiated Is your Facility affiliated with a government agency or part of a government funded health service? 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: Japanese 95%, Asian 3%, Others 2%



IRB/ERB/ETHICS COMMITTEE What is the average time (in days) to start a study once you have received the regulatory package?	Less than 30 91-120	30-60 Greater	O 61-90 than 120
Does your Facility perform IRB/ERB/Ethics Committee submissions?		Yes	○ No
Does your Facility have a dedicated department or group to perform IRB/ERB/Ethics Committee submissions?		Yes	No
Department Contact Name	Clinical Trial Managemen	nt Office	
Department Contact Phone Number	+81-59-256-1212		
Department Contact Email Address	317-ch01@mail.hosp.go	o.jp	
Is your Facility able to initiate study activities prior to IRB/El Committee protocol approval?	RB/Ethics	Yes	○ No
What types of IRB/ERB/Ethics Committee does your Facility use? (Select all that apply.)	Local	✓ Centra	l Acting as Local
Does your institution and/or local regulation mandate the case safety reports [e.g., development Safety Update report (DSI suspected unexpected serious adverse reaction (SUSAR) to a local Review Only IRB/ERB/Ethics Committee?	UR),	Yes	No
Are there any other steps that the Sponsor should be aware IRB/ERB/Ethics Committee review and submission?	e of for your	Yes	● No
If Yes, provide details about the role various committees plasite's review and submission process. If you have multiple le explain what drives the decision on which IRB to use.			



Local IRB/ERB/Ethics Committee

IRB/ERB/Ethics Committee Name	National Hospi	tal Organization Mie C	huo Medical Center	· IRB
Street Name and Number	2158-5, Hisai m	nyoujincho		
Building/Floor/Room/Suite				
Additional Address Info				
Country	Japan			
State/Province/Region	Mie			
City	Tsu			
Zip/Postal Code	514-1101			
Registration No.	Registering	Body		
What is the meeting frequency of your Loc IRB/ERB/Ethics Committee?		Weekly Quarterly		Month Monthly
How long before IRB/ERB/Ethics Committee the Submission Packet required?	ee review is	1 week	2 weeks	S
Does the IRB/ERB/Ethics Committee requirements for to release of final approval documents.		Greater t	han 2 weeks Yes	No
Does the IRB/ERB/Ethics Committee requir approval prior to release of final approval		udget	Yes	No

Note: Attachments can be uploaded online from the Facility Profile in SIP.

Note: Additional Local IRB/ERB/Ethics Committees can be added online from the Facility Profile in SIP.

CENTRAL ACTING AS LOCAL IRB/ERB/ETHICS COMMITTEE

Note: Central Acting as Local IRB/ERB/Ethics Committee can be selected online from the Facility Profile in SIP.



REVIEW ONLY IRB/ERB/ETHICS COM	MMITTEE		
IRB/ERB/Ethics Committee Name			
Street Name and Number			
Building/Floor/Room/Suite			
Additional Address Info			
Country	- Select Country -		
State/Province/Region	- Select State -		
City			
Zip/Postal Code			
Registration No.	Registering Boo	dy	
Note: Additional Review Only IRB/ERB/Ethics Committee	s can be added online from the I	Facility Profile in SIP.	
OTHER REVIEW BOARDS			
Does your Facility have other review the study prior to IRB/ERB/Ethics Com	nmittee submission?		Yes No
For example, scientific, radiation safet	ry committees, or oth	ers.	
Review Board Name	Meeting Freque	ency	
	Weekly	Twice a Month	Monthly
	Quarterly	Other	
	☐ Weekly	Twice a Month	Monthly
	Quarterly	Other	



LOCAL LAB

Yes No
National Hospital Organization Mie Chuo Medical Center Clinical Laboratory
2158-5, Hisai myoujincho
Japan
Mie
Tsu
514-1101
l that apply)
CLIA CAP ISO Others Japan Medical Association
om the Facility Profile in SIP.

Note: Additional Local Labs can be added online from the Facility Profile in SIP.



CONSENT AND TRAINING

CONSENT

Does your Facility have a written SOP/Policy/Procedure for: Informed Consent?	O Yes	No
Does your Facility have a written SOP/Policy/Procedure for: Other vulnerable	Yes	● No
populations?		
Does your Facility have a written SOP/Policy/Procedure for: Minor Assent for	Yes	No
pediatric populations?		
Will your Facility require language translations for consents?	Yes	O No
Note : Languages can be selected online from the Facility Profile in SIP.		
If located in the US, has your Facility used or are you able to use the informed	O Yes	O No
consent short form?	O Don't I	Know
	○ Not Ap	oplicable
TRAINING		
Does your Facility have a training program for the research staff?	Yes	O No
Does the course content include GCP?	Yes	O No
Does your Facility use an external program to conduct research training?	Yes	O No
Please provide program course name:	eAPRIN	
Do you have a process or program in place to retrain research staff when a protocol is amended?	Yes	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?	O Yes	No



FACILITY AND EQUIPMENT

FACILITY CAPABILITIES

Can your Facility support patient visits on weekends?	•	Yes	\bigcirc	No
Can your Facility support in-patient admissions for research studies?	•	Yes	\bigcirc	No
Does your study staff have sufficient English knowledge to understand communications in English?	0	Yes	•	No
Does your Facility have access to translators and translation support for study conduct (e.g. consent, study specific instruction)?	O	Yes Not Ap	O plicab	No ole
Does the Facility have storage space for Study-Related materials (e.g. Lab Kits, Patient Materials, etc.)?	•	Yes	0	No
Does your Facility have the ability to collect and store PK/PD specimens?	•	Yes	\bigcirc	No
Does your Facility have the ability to collect PK/PD samples beyond normal business hours?	•	Yes	0	No
Does your Facility typically allow the collection of Pharmacogenomic (PGX) samples for research purposes?	•	Yes	0	No

SIP Facility Profile Form v3.0 Last Updated 05-Nov-2018



EQUIPMENT

✓	CT Scan	Not Applicable Computerized Tomography Scan		
✓	DXA	Dual-Energy X-ray Absorptiometry or Bone Densitometry		
	ECG/EKG	Electrocardiogram		
✓	FLRO	Fluoroscopy		
✓	MRI	Magnetic Resonance Imaging		
\checkmark	MRA	Magnetic Resonance Angiography (MRA)		
\checkmark	MRS	Magnetic Resonance Spectroscopy (MRS)		
\checkmark	MAMMO	Mammography		
\checkmark	NMED	Nuclear medicine (e.g. Bone scan, thyroid scan, thallium cardiac	stress test)	
	PET	Positron Emission Tomography Scan		
\checkmark	X-ray	X-Radiation		
	Other	Other		
Descr	ibe any addii	tional equipment relevant to Clinical Trials:		
GENE	RAL EQUIPN	MENT		
and m	aintenance o	have an SOP or process that ensures routine calibration of general equipment? Examples of general equipment se oximeter, stadiometer, sphymomanomer, etc.?	O Yes	● No
	our Facility de cart)?	have the necessary equipment to treat medical emergencies	Yes	O No



Identify the equipment available at the Facility to support Research studies?

Centrifuge

	3	
	Refrigerated Centrifuge	
✓	Refrigerator (2 to 8 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 No
	Does this equipment provide Min/Max Temperature Monitoring?	Yes No
	How frequently can temperature measurement occur? Check the most frequent	Daily
	measurement your equipment can support.	,
	Does this equipment have back-up power?	Yes O No
	Does this equipment have a temperature alarm?	• Yes • No
	Do you have an SOP which supports calibration of this equipment?	Yes No
✓	Freezer (-20 to -30 Degrees C)	
	Equipment Capabilities: Freezer (-20 to -30 Degrees C)	
	Do you have the ability to generate a temperature monitoring log for this equipment?	Yes No
	Does this equipment provide Min/Max Temperature Monitoring?	Yes • No
	How frequently can temperature measurement occur? Check the most frequent	Less than Daily
	measurement your equipment can support.	
	Does this equipment have back-up power?	• Yes • No
	Does this equipment have a temperature alarm?	Yes No
	Do you have an SOP which supports calibration of this equipment?	Yes No
✓	Freezer (-70 to -80 Degrees C)	
	Equipment Capabilities: Freezer (-70 to -80 Degrees C)	
	Do you have the ability to generate a temperature monitoring log for this equipment?	Yes No
	Does this equipment provide Min/Max Temperature Monitoring?	Yes No
	How frequently can temperature measurement occur? Check the most frequent	Less than Daily
	measurement your equipment can support.	O Vac O Na
	Does this equipment have back-up power?	Yes No
	Does this equipment have a temperature alarm?	Yes No
_	Do you have an SOP which supports calibration of this equipment?	O Yes O No
	Freezer (Liquid Nitrogen -135 Degrees C)	
	Equipment Capabilities: Freezer (Liquid Nitrogen -135 Degrees C)	O v O v
	Do you have the ability to generate a temperature monitoring log for this equipment?	Yes O No
	Does this equipment provide Min/Max Temperature Monitoring?	O Yes O No
	How frequently can temperature measurement occur? Check the most frequent	- Select -
	measurement your equipment can support.	O Yes O No
	Does this equipment have back-up power? Does this equipment have a temperature alarm?	O Yes O No
	Do you have an SOP which supports calibration of this equipment?	O Yes O No
	20 you have an 301 which supports cambration of this equipment.	



COMPUTER CAPABILITIES

Does your Facility have computers which are dedicated to research studies?	Yes	O No		
What type of computer operating system(s) does your institution use to support studies?				
✓ Windows (Windows XP, Windows 7, Windows 8, etc)				
Apple/Mac (OS X Snow Leopard, Mountain Lion, El Captain, etc)				
Unix/Linux (Solaris, Ubuntu, Redhat, etc)				
I don't know				
Other				
What type of internet access does your Facility have?	Cable or DSL			
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)?	No			
Does the Facility have access to local IT support?	Yes			



INVESTIGATIONAL PRODUCT & CONTROLLED SUBSTANCES

INVESTIGATIONAL PRODUCT SHIPPING DETAILS

IP Recipient Name	Mie Chuo Medical Center Clinical Trial Management Office
Street Name and Number	2158-5, Hisai myoujincho
Building/Floor/Room/Suite	
Additional Address Info	
Country	Japan
State/Province/Region	Mie
City	Tsu
Zip/Postal Code	514-1101
Phone Number	+81-59-256-1212
Fax Number	+81-59-256-1212
Email Address	317-ch01@mail.hosp.go.jp



INVESTIGATIONAL PRODUCT STORAGE LOCATION

IP Storage Location Name	Mie Chuo Medical Center Clinical Trial Management Office
Street Name and Number	2158-5, Hisai myoujincho
Building/Floor/Room/Suite	
Additional Address Info	
Country	Japan
State/Province/Region	Mie
City	Tsu
Zip/Postal Code	514-1101
Phone Number	+81-59-256-1212
Fax Number	+81-59-256-1212
Email Address	317-ch01@mail.hosp.go.jp

Note: Additional Investigational Product Storage Locations can be added online from the Facility Profile in SIP.



INVESTIGATIONAL PRODUCT STORAGE EQUIPMENT

Identify the Investigational Product Storage Equipment at your Facility

√	Refrigerator (2 to 8 Degrees C)	
[<u>]</u>	Equipment Capabilities: Refrigerator (2 to 8 Degrees C) Do you have the ability to generate a temperature monitoring log for this equipment? Does this equipment provide Min/Max Temperature Monitoring? How frequently can temperature measurement occur? Check the most frequent measurement your equipment can support. Does this equipment have back-up power? Does this equipment have a temperature alarm? Do you have an SOP which supports calibration of this equipment?	Yes No Yes No Yes No O Yes No Yes No Yes No Yes No Yes No
V Fr	eezer (-20 to -30 Degrees C)	
	Equipment Capabilities: Freezer (-20 to -30 Degrees C) Do you have the ability to generate a temperature monitoring log for this equipment? Does this equipment provide Min/Max Temperature Monitoring? How frequently can temperature measurement occur? Check the most frequent	Yes No Yes No Less than Daily
	measurement your equipment can support. Does this equipment have back-up power? Does this equipment have a temperature alarm? Do you have an SOP which supports calibration of this equipment?	Yes No Yes No Yes No Yes No
✓ Fr	eezer (-70 to -80 Degrees C)	
	Equipment Capabilities: Freezer (-70 to -80 Degrees C)	
	Do you have the ability to generate a temperature monitoring log for this equipment? Does this equipment provide Min/Max Temperature Monitoring? How frequently can temperature measurement occur? Check the most frequent	Yes NoYes No
	measurement your equipment can support.	Daily
	Does this equipment have back-up power? Does this equipment have a temperature alarm? Do you have an SOP which supports calibration of this equipment?	YesNoYesNoYesNo
Fr€	eezer (Liquid Nitrogen -135 Degrees C)	
	Equipment Capabilities: Freezer (Liquid Nitrogen -135 Degrees C) Do you have the ability to generate a temperature monitoring log for this equipment? Does this equipment provide Min/Max Temperature Monitoring? How frequently can temperature measurement occur? Check the most frequent	Yes No Yes No
	measurement your equipment can support.	- Select -
	Does this equipment have back-up power? Does this equipment have a temperature alarm? Do you have an SOP which supports calibration of this equipment?	Yes No Yes No Yes No



INVESTIGATIONAL PRODUCT STORAGE & HANDLING

Is the Investigational Product Storage Room secured with controlled access?	Yes	O No
Do you have the ability to generate a temperature monitoring log for this	Yes	○ No
Investigational Product Storage Room?	<u> </u>	O 1.10
Does the Investigational Product Storage Room provide Min/Max temperature	Yes	O No
monitoring?	res	○ No
Does the Investigational Product Storage Room have back-up power?	Yes	O No
Does the Investigational Product Storage Room have a temperature alarm?	Yes	O No
Do you have an SOP which supports calibration of the temperature	Yes	No
monitoring equipment?		
Does your Facility have the ability to manage on-site or off-site destruction	Yes	○ No
of Investigational Product?		
Does your Facility have a written SOP/Policy/Procedure for destruction of	Yes	No
Investigational Product?	Not Applicable	
Do you provide your Satellite Site(s) with a dedicated inventory of	○ Yes	● No
Investigational Product?	Not Applicable	
Does your Facility have a written SOP/Policy/Procedure to ensure that	Yes	O No
Investigational Product is appropriately maintained during transportation to	Not Ap	plicable
Satellite Site(s)?		
Describe additional Investigational Product Storage & Handling Capabilities:		



PREPARATION AND ADMINISTRATION OF INVESTIGATIONAL PR	RODUCT		
Identify the Investigational Product preparation capabilities at your Fa	acility:		
✓ Extemporaneous Preparation			
✓ Vertical laminar flow hood (chemo/hazardous drugs)			
Glove box (non-vented)			
Horizontal laminar flow hood (non-hazardous drug preparation)			
Glove box (vented to outside)			
Preparation and Administration of Investigational Product			
Is your Facility capable of administering infusions?		Yes	O No
Is your Facility adequately staffed to support studies with both blinded and un-		Yes	O No
blinded Investigational Product?		0 163	O 110
CONTROLLED SUBSTANCES			
Controlled Substances are defined as: A drug or chemical whose manuf	facture, posse	ssion, or use is	s regulated
a government, such as illicitly used drugs or prescription medications the	hat are desigr	nated a Contro	olled Drug.
Does the Facility have the required licenses or registrations	Yes	○ No	
to receive, store, dispense and return controlled substances	ONot Ap	plicable	
as required by local law?			
Is the storage area for controlled substances securely constructed	Yes	ONo	
with restricted access in accordance with local law?	O Not Ap	plicable	
	Yes	No	
Does the Facility have the ability to handle radio-labelled	Yes	O NO	
Investigational Product?			
Does your Facility have the ability to manage on-site or	Yes	○ No	
off-site destruction of controlled substances when appropriate?	○Not Ap	plicable	
ATTACHMENTS			
		1 12 1	

Upload relevant Investigational Product & Controlled Substances documentation including: relevant SOPs for managing or storing Investigational Product(s), IP storage equipment, or licenses/registrations to receive, store, dispense and return controlled substances.

Note: Attachments can be uploaded online from the Facility Profile in SIP.



SOURCE DOCUMENTATION

SOURCE DOCUMENTATION			
SOURCE DOCUMENTS			
What type of source documents will be used? (Select all that app	oly):	✓ Paper	✓ Electronic
Does your Facility have secure storage for patient records?		Yes	○ No
Does your Facility have patient record archiving on-site?		Yes	○ No
Provide Location name and address of any offsite archives.			
ELECTRONIC MEDICAL RECORDS (EMR) /ELECTRONIC HEALT	TH RECORD	OS (EHR)	
Do you have Electronic Health Records (EHR)/ Electronic Medical Records	rds (EMR)?	Yes	O No
What EMR/EHR system do you use?	✓ In-hor	use system	Others
Note: Please select other options for EMR/ EHR used at your Facility online.			
For Facilities with satellite sites, where is the monitor required to access source documents?		Select	
Please list any access limitations/requirements for the Electronic M	<u>1edical Reco</u>	ords:	
None			



MONITORING Check all equipment that will be available to Monitors: None ✓ Phone ✓ Fax ✓ Copy Machines **Internet Access** What Electronic Data Capture (EDC) systems has your staff used for clinical trials? ✓ Oracle Inform ✓ Medidata Rave Oracle Remote Data Capture (RDC) ✓ Others None Describe Other EDC Systems: **CRSCube** ADDITIONAL INFORMATION AND 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** Upload any non-study specific Facility documents that have not been included in other sections of the profile. Lab, IRB/ERB/Ethics Committee, Investigational Product and Controlled Substance documentation should be included in those sections. The document type drop-down list provides examples of the type of documentation to be included in this section. Note: Attachments can be uploaded online from the Facility Profile in SIP.