

## **Site Profile Form**

**Purpose of Site Profile form:** The intent of the Site Profile form is to capture site capabilities that are collected during site qualification and not to replace current individual pre-study activities. The intent is to reduce the administrative burden on sites associated with completing multiple forms requesting the same or similar information. The form is not intended to capture study specific or therapeutic specific information.

The form will be in an electronic format, with drop down or check boxes to keep the form simple and easy to use. There will be free text input boxes for providing any necessary explanations. Site should keep a copy of the completed form on file.

If additional text is needed in any of responses, use an asterisk and enter at the bottom of the form.

1. COMPLETED BY:
Full Name: Renzo Yokoyama
Date Completed: May 12, 2017 Role: Center for Clinical Research Advisor to the Director, Scretariat, DM, CCRC
Investigator Name:
2. SITE DETAIL:
Institution Name: Okinawa Prefectural Chubu Hospital
Address (Location): 281, Aza-miyazato
City: Uruma State/Region/Province: Okinawa
Country: Japan Postal Code: 904-2293
Type: Hospital Public
Therapeutic Area: ☐ Auto immune ☐ Cardiovascular ☐ Critical Care ☐ Dermatology ☐ Infectious Disease ☐ Men's Health ☐ Metabolic/ Endocrine ☐ Musculoskeletal ☐ Neuroscience ☐ Oncology ☐ Osteoporosis ☐ Pain ☐ Pediatrics ☐ Psychiatry ☐ Respiratory ☐ Vaccines ☐ Virology ☐ Women's health Other:  Trial phase capabilities: ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐
Do you have affiliated research sites or satellite sites/clinics? X Yes No
Which different sponsor type(s) do you have research experience? 💢 Industry 🗀 Academic 🦵 Investigator Initiated 🦳 None
Ethnicity of patient population - please break down your population by % of ethnicity
Japanease(95%)
Demographics of patient population: 🗵 Pediatric 🗵 Adult Other comments:
Is your site affiliated with a government agency or part of a government funded health service? Yes X No
Site Contacts: Primary site contact for clinical trials
First Name: Renzo Phone: +81-98-973-4111 Fax: +81-98-973-4124
Surname: Yokoyama Email: yokoyama_renzo@hosp.pref.okinawa.jp



	AL COMMITTEE REVIEW PROCESS  - This section is only applicable if the site is directly resp	ancible for performing the others	ammittae submission
	:/Ethics committee	7	ommittee submission.
IND/ END		City: Uruma	
Name:	Okinawa Prefectural Chubu Hospital Institutional Review Board	Province: Okinawa	
Address:	281, Aza-miyazato	Country: Japan	
	281, Aza-miyazato	Postal Code: 904-2293	
IRB/ERB	/Ethics committee registration number (if applicable)		thics committee type: Local Kocal Central/acts as lo
N/A		Name: Renzo Yokoyama	Local P.S Centraly acts as to
IRB/ERB If yes, ple to the rig	tht of the form	Phone: +81-98-973-4111  Email: chubu_research@hosp.pref	
steps are steps are approva - IRB/ERB - Amount - Amount	provide a general outline of the steps required to obtain e dependent on one another, and/or if they can be come e covered, in addition to any other applicable administral, scientific review committees, etc.) / Ethics committee(s) meeting schedule/frequency cof time in advance of an IRB/ERB/ Ethics committee meeting to of time following an IRB/ERB/Ethics committee review you recur local IRB/ERB/Ethics committee require payment of any fees	pleted in parallel or in sequence. Fative steps required at your site (exhaus all documentation must be submitted eive written confirmation of approval	Please ensure that the followin xample – contract/budget
- Amoun - Amoun - Does y	eeting schedule/frequency: The second Tuesday of each rat of time in advance of an IRB that all documentation mus at of time following an IRB review you receive written configuration local IRB require payment of any fees ahead of subm	t be submitted: 15 business days be rmation of approval: Wthin 7 buisined ission or prior to the release of the fil	ss days nal approval documents?: None
Please pr steps are	this section is only applicable if the site is NOT responsible ovide a general outline of the steps required to obtain a dependent on one another, and/or if they can be compreview committees, or other, but excluding ethical compactors in N/A or please, explain.	approval for a study at your institut eleted in parallel or in sequence (ex	tion/site, including whether an ample- contract/budget appro
	MED CONSENT ur site have a written SOP, policy/procedure for Informa	nd Consont?	⊠ Yes No
	r Assent for pediatric populations?		⊠ Yes
	·vulnerable populations?		∑ Yes
Will you	r site require language translations for consents		⊠ Yes
	nat languages will be required? Please list. Japanese		
	UALIFICATIONS/TRAINING ur site have a training program for the research staff?.	5	⊠ Yes □ No
	ur site have a training program for the research staff ne course content include GCP?		Yes X No
Does yo	our site use an external program to conduct research tra	aining? If yes, please provide progr	ram course name:   Yes  X
Does yo	ur program have a provision for training staff when upd	lates to protocols occur?	☐ Yes        No



6. FACILITIES AND EQUIPMENT				
LOCAL LAB:				
Name/Details: Local Lab				
Phone: +81-98-973-4111	Fax: +81-98-973-4124	Email: chubu_	research@hosp.pr	ef.okinawa.jp
Local lab accreditation GLF	CLIA CAP ISO	other LIMA JAMT O	MA	
Does the study staff that prepare	s or transports dangerous goods have Association (US) or other countries ha rous goods?	training that meets t	he	□ No ⊠ N/A
EQUIPMENT:			1 163	
	routinely?			No
	ency available?			No
	ers for biological sample storage?		•	•
-	logical sample storage?		* **	∑-70
				□ No
_	for refrigerators?			No
	for freezers ?lable?			No
	er outage of refrigerators and freezer:		The second secon	No
				No
	pment is out of range for refrigerators		•	⊠ No
Do you have				No
•		nternational phone li		No
	cess lab samples? uge for processing lab samples?			
Do you have reingerated centrific	ige for processing ian samples:		····· 🔀 Yes	┌─ No
COMPUTER CAPABILITY:				
	moutors for the research studies?		⊠Yes	No
What is your current browser and	mputers for the research studies?	••••••••••••	1^ tes	[ NO
Safri Ver.10.1 (12603.1.30.0.34),A				
2 CH2-00 - C-00 C-00 - C-00 C-00 C-00 C-00 C	valls?		× Yes	No
	nternet access?			
			CONTROL CONTROL	No
boes your site have wheless life	rnet capabilities?	***************************************	······ 🔀 Yes	No
OTHER:				
PK/PD capability?	***************************************		🗵 Yes	No
Lab hours to accommodate PK/PI	D studies beyond (8-5, M-F)?	••••••	🔀 Yes	No
Is your site open on weekends?			× Yes	No
Are you able to admit research su	ubjects to an in-patient setting for res	earch purposes?	X Yes	Γ <sup>™</sup> No
DIGITAL DIAGNOSTIC CAPABILITI	ES:			A
	ray 🔀 DXA 🗍 Other (please list)			
STORAGE FACILITIES:				
	ge secured to protect patient privacy?		🔀 Yes	No
Are the archiving facilities on site		provide name and lo		
The tire dremaing facilities off site	i po res i no, ii offsite	provide name and 100	anon miorination.	
Is there storage area on site for st	tudy related materials, ex. Lab kits or	other items?	X Yes	□ No
	,			, 140



7. INVESTIGATIO	NAL PRODUCT (IP)		
Ship to address:	281, Aza-miyazato, Uruma-city, Okinawa, 904-2293, Japan Okinawa Prefectural Chubu Hospital		
Primary	Ph	00.070.4444	
Contact:  Center	for Clinical Research Phone:  +81	98-973-4111	
Email: chubu_	research@hosp.pref.okinawa.jp +81	-98-973-4124	
Storage location	the same as the shipping address? (if study specific skip)	⊠ Yes	No
Infusion capabili	ty?	⊠Yes	No
IP-STORAGE AND			
Is the IP storage	area secured with controlled access?	▼ Yes	No
Is the temperatu	re monitoring available for the following? $oximes$ Room temp $oximes$ Refrigera	ator Fre	ezer
Please detail ter	nperature device capabilities (for example -min/max), frequency for monitoring.		
min/max, every	dav	-	
	re monitoring alarmed in the event that there is an excursion?	▼ Yes	No
	plan in the event of a power outage or equipment failure?	▼ Yes	No
	uately staffed to perform both blinded and un-blinded roles, in case un-blinded		
drug monitoring	is required?	× Yes	No
8 OUESTIONS S	PECIFIC TO DESTRUCTION OF IP		
	ave the capability to destroy IP on site/arranged directly via sub-contractor?	▼ Yes	□ No □ N/A
<u>-</u>	ave a written SOP/policy/procedure for IP destruction?	▼ Yes	□ No □ N/A
IP – SATELLITE S		, ,,	
	site(s) have a dedicated inventory?	Yes	□ No 区 N/A
Do you have a d	rug transportation procedure for satellite sites?	Yes	□No 区N/A
9. QUESTION SP	ECIFIC TO CONTROLLED SUBSTANCES		
	ve the regulatory required licenses or registrations to receive, store, dispense		
	olled substances as required by local law?	▼ Yes	□No □N/A
	ity for controlled substances is securely constructed with restricted access to	57 V	Car Carlo
•	diversion?capability?	⊠ Yes ⊠ Yes	□No □N/A □No □N/A
	ave the capability to destroy IP on site for controlled substances?		No N/A
,	,	X	
	UMENTATION/CRFS/SITE MONITORING		
	its: Are site source documents Paper Electronic Electro	Both	
-password mana- restrict access	agement to medical records		
Will monitors ha	ve access to $\overline{\boxtimes}$ Phone $\overline{\boxtimes}$ Fax $\overline{\boxtimes}$ Copy machines	⊠ Interne	et access
CRFs			
	data systems has your staff used for clinical trials? 💢 Inform 🧮 Medid	ata Rave	
Other, ple	ase list		



out your site. Ple	ease reference section nu	mber if applicable:			
			727		

Please provide any additional information not captured elsewhere on this form, that you feel is important that we should know