Princess Alexandra Hospital - Radiation Oncology Raymond Terrace (ROPART)

Local Services Offered:

Clinical trials site;Investigator Initiated Trials;Satellite Site;Trial Patient Recruitment;Treatment of patients;Completion of study documentation as per ICH GCP and contract

Details: This PDF complements the information found in the CSV generated by this website with additional site information.

Facility Details:	
Please provide your Facility Website.	https://www.metrosouth.health.qld.gov.au/hospital-and-health- centres/princess-alexandra-hospital
What Department is your Trial Site? (Queensland Health HHS- See List of Services and Levels-Clinical Services Capability Framework).	Medical Services
Is your Facility affiliated with a government agency or part of a government funded health service?	Yes
Is your facility/organisation a Life Sciences Queensland (LSQ) Member?	No
Provide the list of Sub-Therapeutic Areas for your Facility.	Radiation; oncology; cancer
Has your Clinical Trial Site been accredited?	Not Applicable
Does your Clinical Trial site undertake any patient recruitment?	Yes
What percentage of Clinical trials undertaken on your site do you meet or exceed the recruitment target?	80-90%
IRB/ERB/Ethics Committee:	
Does your Facility perform HREC (IRB/ERB/Ethics) Committee submissions?	Yes

Does your Facility have a dedicated department or group to perform HREC (IRB/ERB/ETHICS) Committee submissions?	No
HREC Committee Name.	Metro South HHS HREC
Are there any other steps that the Sponsor should be aware of for your IRB/ERB/Ethics Committee review and submission?	No
Does the HREC Committee require payment prior to the release of final approval documents?	Yes
Does the HREC require contract/budget approval prior to release of final approval documents?	Yes
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 paediatric populations?	Yes
Does your Facility have a written SOP/Policy/Procedure for Other vulnerable populations?	Yes
Does your Facility have access to translators and translation support for study conduct (e.g. consent, study- specific instruction)?	Yes
Training:	
Does your Facility have a training program for the research staff?	Yes
Do you have a process or program in place to retrain research staff when a protocol is amended?	Yes

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?	Yes
Facility And Fauinmonts	
Facility And Equipment:	
<u>Facility Capabilities:</u> Can your Facility support in- patient	Yes
admissions for research	
studies?	
Can your Facility support patient	Yes
visits on weekends?	
Is your Facility capable of	Yes
administering infusions?	
Is your Facility adequately staffed to support studies with both blinded and unblinded Investigational Product?	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
Does the Facility have storage space for Study-Related materials (e.g. Lab Kits, Patient Materials, etc.)?	Yes
Equipment:	
Does your Facility have the necessary	Yes
equipment to treat medical emergencies (for example crash/code cart)?	

Does your Facility have an SOP or	Yes
process that ensures routine	
calibration and maintenance of general	
equipment? Examples of general	
equipment include: scale, pulse	
oximeter, stadiometer,	
sphygmomanometer, etc.?	
IT Capabilities:	
Does your Facility have computers	Yes
that are dedicated to research studies?	
What type of computer operating	Windows (Windows XP; Windows 7; Windows 10;
system(s) does your institution use to	etc)
support studies?	
What browser does your facility use?	Chrome
Does the Facility have access to	Yes
local IT support?	
Does your Facility limit or prohibit	Yes
access and use of external web-based	
tools or sites for clinical research (E.g.	
web portals to submit documents to	
sponsors or CROs)?	
Please indicate all equipment that will	Phone;Copy Machines;Internet Access
be available to Monitors	r none, Copy Machines, internet Access
Labs:	
Does your Facility use Local Lab	Yes
Services?	
Please provide Local Lab Name	Pathology Queensland-PAH
IP Storage Details:	
IP Recipient Name	PA Clinical Trials Pharmacy Department
Does the Investigational Product	Yes
Storage Room have back-up power?	
Is the Investigational Product Storage	Yes
Room secured with	
controlled access?	
Is the Investigational Product Storage	Yes
Area securely constructed?	

Does the Facility have the ability to handle radio-labelled Investigational Products?	Yes
Does your Facility have the ability to manage on-site or off- site destruction of controlled substances when appropriate?	Yes
Does your Facility have the ability to manage on-site or off- site destruction of the Investigational Product?	Yes
Does your facility have a written SOP/Policy/Procedure for the destruction of Investigational Product?	Yes
Do you provide your Satellite Site(s) with a dedicated inventory of Investigational Product?	Yes
Does your Facility have a written SOP/Policy/Procedure to ensure that Investigational Product is appropriately maintained during transportation to Satellite Site(s)?	Yes
Source Documents:	
Does your Facility have secure storage for patient records?	Yes
Does your Facility have patient record archiving on-site?	Yes
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?	In-house system