Princess Alexandra Hospital -Haematology

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: In addition to the information contained within csv generated by this website, this PDF provides additional site information

Does your Facility perform	Yes
HREC (IRB/ERB/Ethics)	
Committee submissions?	
Does your Facility have a written	Yes
SOP/Policy/Procedure for	
Informed Consent?	
Does your Facility have a	No
dedicated department or group to	
perform HREC	
(IRB/ERB/ETHICS) Committee	
submissions?	
Does your Facility have the	Yes
necessary equipment to treat	
medical emergencies (for	
example crash/code cart)?	
Is your Facility affiliated with a	Yes
government agency or part of a	
government funded health	
service?	
Does your Facility typically	Yes
allow the collection of	
Pharmacogenomic (PGX)	
samples for research purposes?	
Does your Facility have	Yes
computers that are dedicated to	
research studies?	
Does your Facility have the	Yes
ability to manage on-site or off-	
site destruction of controlled	
substances when appropriate?	
Does the Facility have the ability	Yes
to handle radio-labelled	
Investigational Products?	
Is your Facility capable of	Yes
administering infusions?	
What browser does your facility	Chrome
use?	

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Lab Name	Pathology Queensland-PAH
Local Lab Usage	Yes
Committee Country	Australia
Do you have Electronic Health	Yes
Records (EHR)/ Electronic	
Medical Records (EMR)?	
Life Sciences Queensland (LSQ)	
Member	No
Does your Facility have a written	Yes
SOP/Policy/Procedure for Minor	
Assent for paediatric	
populations?	
Does your Facility have a written	Yes
SOP/Policy/Procedure for Other	
vulnerable populations?	
IP Recipient Name	PA Clinical Trials Pharmacy Department
Does the HREC Committee	Yes
require payment prior to the	
release of final approval	
documents?	
Does your Facility have access to	Yes
translators and translation	
support for study conduct (e.g.	
consent, study-specific	
instruction)?	
Do you provide your Satellite	Yes
Site(s) with a dedicated	
inventory of Investigational	
Product?	
Do you have a process or	Yes
program in place to retrain	
research staff when a protocol is	
amended?	
Does your Facility have a	Yes
training program for the research	
staff?	
Training program for the	Yes
research staff	
Storage Room Backup Power	Yes
Is the Investigational Product	Yes
Storage Room secured with	
controlled access?	
Storage area securely constructed	Yes

What type of computer operating	Windows (Windows XP; Windows 7;
system(s) does your institution	Windows 10; etc)
use to support studies?	
Other areas of expertise	Leukemia; Lymphoma; Multiple myeloma;
-	and Amyloidosis
Sub-Therapeutic areas	Acquired Haemophilia;Acute Lymphoblastic
	Leukaemia;Acute Myelogenous
	Leukaemia;Chronic Lymphocytic
	Leukaemia;Chronic Myelogenous
	Leukaemia;Lymphocytic leukaemia;Mantle
	Cell Lymphoma
Please indicate all equipment	Phone;Copy Machines;Internet Access
that will be available to Monitors	
Has your Clinical Trial Site been	Not Applicable
accredited?	
Has your Clinical Trial site been	Not Applicable
audited?	
If your Clinical Trial Site has	Other
been audited, please select all	
relevant types.	
Does your Clinical Trial site	Yes
undertake any patient	
recruitment?	
What percentage of Clinical	80-90%
trials undertaken on your site do	
you meet or exceed the	
recruitment target?	
EMR/EHR systems	In-house system
Does the HREC require	Yes
contract/budget approval prior to	
release of final approval	
documents?	
Any Additional Process	No