Research Portal User Manual Researcher

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Introduction

The Ministry of Public Health's role is to create a clear vision for the nation's health direction, set goals and objectives for the country, design policies to achieve the vision, protect the public's health, ensure high quality health research, and monitor and evaluate progress towards achieving those objectives.

A key function of MoPH is oversight research and supporting researchers to ensure that standards are met and performance targets achieved.

In implementation of these technical plans, this document contains the details about the process of accessing the Research web portal as a Researcher.

Application access

Research Department - MoPH Website

Research Department - MoPH Portal

User registration & Accesses

1. For registering as researcher click on "Register as Researcher"

Sign In	
Register as Focal Point Register as Researcher Register as Sponsor Forgot password?	

First name*	Enter Name	
Middle initial	Entet Middle Name	
Last name*	Enter Last Name	
Login id/email (Please use your business email id)*	example@mail.com	
Enter password*	Enter Password	
Confirm password*	Confirm Password	
Telephone	974 XXXXXX	
Institution*	Select	~
Supporting Document *	Please attach file (.PDF Only)	ত্রি Browse
	Et Submit	

Enter all the mandatory details and upload supporting/authorization letter from your institution supporting your role.

Click submit after entering all the details

Thank you for registering with MoPH Research Department. An email has been sent to your registered email id to activate your account.

You will receive an email verification on your registered email within few minutes.

Dear User, Thank you for registering with MoPH Research Portal <u>Please click here to activate your account.</u> Best regards, Research Team, Ministry of Public Health

2. User Login after email verification -

	Login
A LE OF CEL	Registered email address Password
⁷⁷ OP PUBL ^C	Sign In
HEALTH RESEARCH PORTAL	Register as Focal Foint Register as Researcher Register as Sponsor Forgot password?

For registered users -enter registered email and password for Sign In to system.

3. Forgot password

Forgot Password	
Please provide your registered email id and cliv Login id/email (Please use your business email id)	ck on submit, we will send link to your registered email to reset password.
	🖺 Submit



Enter registered email id and 'click' submit, user will get email for reset password.

MoPH Research Portal Reset Password 😕 Inbox 🛪

research@moph.gov.qa

🖹 to me 👻

Dear User,

Please click here to reset password.

Best regards,

Enter new password and click submit button.

Reset your account password

Please enter new password*		
Please confirm new password*		
	E Submit	

Menu access

After successful login as Researcher navigate into landing page.

elcome ! Researche	r , you have logged as Researcher		
Register Clinical Trial	Update Profile	Health Research Registry	Research Training & Workshop
View / register / submit clinical trial	Update profile details like address, contact email etc	View /register a health research	View/register research training
Click to access	Click to access	Click to access	Click to access
Research Proposal	Health Research Data Collaboration	Research Publications	Serious Adverse Event
View/register proposal	Request for health research data from a registered research	Register research publication	Report serious adverse event destails
Click to access	Click to access	Click to access	Click to access

Click on the respective tile to take an action, some are explain below.

4. Update profile

Update researcher profile.

5. Register Clinical trial

Option to submit /report clinical trial to MoPH

6. Health Research registry

Research institutions within the state of Qatar that conduct research involving Human subjects will report to MoPH their research projects.

7. Research Training & Workshop

View/register research training

8. Health Research Data Collaboration

Request for health research data from a registered research / Collaborate for data with others

9. Research Publications

Register research publication

10. Serious Adverse Event Report serious adverse event details

NOTE: From any section/any page in order to go to your home page, please click on "MoPH logo" on the header (top left section).

Update Profile

Update profile



🖺 Add/Save

Register Clinical Trial

View / register / submit clinical trial

Re	ference no.	Study Phase	Investigational product	Description	Submitted on	Status	Action
1 CT	-2019-0009	phase I	Yes	Observational	14/03/2019	Under Process	Comment
2 CT	-2019-0010	phase I	Yes	Observational	19/03/2019	Saved	🕑 Edit 🖾 Comment
СТ	-2019-0004	phase I	Yes	Observational	23/02/2019	Sentback	🖻 Edit 🖾 Comment
СТ	-2019-0016	phase I	Yes	Observational	16/06/2019	Under Process	Comment
СТ	-2019-0011	phase I	Yes	Observational	27/03/2019	Sentback	🕑 Edit 🖾 Comment

Click on "Register/Submit clinical trial" to register a new clinical trial In order to edit a saved application, click on "Edit" on them respective row in the Saved/Submitted grid

Clinical trial general info	ormation
-----------------------------	----------

Before proceeding with the on-line application, the approval by both local registered Ethics committees, Data and Safety Monitoring Boards of the affiliated study sites (when applicable), is required. The Sponsor or the Principal investigator are requested to register all the clinical trial on the registration portal of the MOPH.

The review time is different according to the following:

Category A application: include multi-center clinical studies that have been approved by the FDA-USA, the EMA-European Commission or other regulatory agencies (in the UK, Switzerland, Australia, Canada, Germany, South Africa, Japan). Category A application's permission is usually faster than Category B. The review is usually done within four to six weeks.

Category B applications: include Clinical trials with no approval from the above regulatory bodies. These trials are reviewed on a caseby-case basis. Category B application's review is usually done within six to eight weeks.

In all applications, a support letter initiated by MOPH is required in order for the clinical trials to start.

Read the instructions carefully before proceeding.

Provide the required information in the screen, click on "next arrow icon" and the system will guide you through the application

Clinical trial general information	on
A clinical trial is a prospective biomedical o questions about biomedical or behavioral in treatments, or device.	r behavioral research study of human subjects that is designed to answer specific nterventions such as: drugs, treatments, devices, or new ways of using known drugs,
Is the clinical research, to be reported, defined as a clinical trial (as per criteria mentioned above)?*	● Yes ○ No

Clinical trial general information

	Fields with * are mandatory
Study phase*	○ phase I ● phase II ○ phase IV
	e hanne e hanne e hanne e hanne
Interventional: A type of clinical trial in which part (or no intervention) so that researchers can evalua assignments are determined by the trial's protoco interventions a study where the medicinal product marketing authorization)	icipants are assigned to groups that receive one or more intervention/treatment te the effects of the interventions on biomedical or health-related outcomes. The ol. Participants may receive diagnostic, therapeutic, or other types of t(s) is (are) prescribed in the usual manner in accordance with the terms of the
Observational: a type of clinical trial in which parti biomedical or health outcomes. Participants may i investigator does not assign participants to a spec	icipants are identified as belonging to study groups and are assessed for receive diagnostic, therapeutic, or other types of interventions, but the cific interventions/treatment.
What is the clinical trial's type ?*	 Observational Interventional
What is the Intervention?*	 A new drug; or an approved drug used for a different indication Drug or combination of drug not used as per same usual care settings (route, dosage, target population, indication of use) A new medical deviceor approved medical device with alterate use A new therapeutic intervention (such as cell transplant, gene transfer etc) None of the above
Is it a new investigational product*	○ Yes
Product already approved for use in Qatar and one of the following is diffrent*	○ Yes ● No
Marketed drug	 Indication(s) and clinical use is different Target patient population(s) is different Route (s) of administration is different Dosage regimen(s) is different
Institution	Institution Name-Demo

⇒

Provide the required information click on "next arrow"

1. Study identification, status, description and keywords

	Fields with * a	re mandatory	
ClinicalTrials.gov Identifier	CT-001-2019		
Brief Title*	Human Anatomy		
Acronym*	HMNA		
Overall Recruitment Status*	Recruiting		
Study expected start date(mm/dd/yyyy)*	8/8/2019		
Expected completion date(mm/dd/yyyy)*	8/10/2019		
Actual completion date(mm/dd/yyyy)			
Brief Summary*	Brief Summary about th	e clinical trial	
			/
Detailed Description*	Detailed Description ab	out the clinical trial	
			/
Primary Disease or Condition Being Studied in the Trial or the Focus of the Study*	Primary Disease or Con	dition Being Studied in the Trial or the Focus of the Study	
			/
Keywords*	Primary Disease		

2. Sponsor Details

	Fields with * are mandatory			
Name of organization*	Name of organization			
Name of the person to contact*	Name of the person to contact			
Address*	Address of organization			
Post code				
Telephone number*	97411224455			
Fax number				
Fox Humber				
Email*	organizationofficial@mail.com			
Legal representative of the sponsor in the state of Qatar (if different from the sponsor)				
Name of organization	Name of organization			
Name of the person to contact	Name of the person to contact			
Address	Address of organization			
Telephone number	97411224455			
Fax number				
Email				
Status of the sponson	Select 🔻			
Source of financing if other than the sponsor				

3. Investigator Details

Fields with * are mandatory				
Principal Investigator				
Name*	Name of Principal Investigator			
Organization*	Principal Investigator Organization			
Address	Principal Investigator Organization Address			
Telephone Number*	97411224433			
Fax Number				
Email*	pioffical@mail.com			
Co-Investigator				
Name	Name of Co-Investigator			
Organization	Co-Investigator Organization			
Address	Co-Investigator Address			
Telephone number	97411224433			
Fax number				
Email	cdpioffical@mail.com			

4. Investigational Product Information

Fields with * are mandatory							
Investigational product information							
Trade name*	Enter trade r	ame					
United state adopted name (USAN)	Enter United	state adopted name					
IP has marketing authorization in other countries	OYes N	0					
Country	Select			٣			
	Reco	rd added successfully.					
		bbA 🖺					
Trade name		USAN	Country	Action			
Investigational product trade name		USANAME	Puerto Rico	🔋 Delete			

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5 . Study Design

Fields with * are mandatory					
Interventional Study Design*	Interventional Study Design				
Primary purpose*	 Treatment Diagnostic Supportive Care Screening Health Services Research Basic Science Device Feasibility 				
Additional Primary purpose (if any)					
Interventional Study Model*	 Single Group Parallel Crossover Factorial Sequential 				
Model Description*	Model Description				
Number of Arms*	4				
Is Masking ?*	No				
Allocation*	● N/A (not applicable) ◎ Randomized ◎ Nonrandomized				
No of participant*	45				

6 . Arms, Groups and Interventions

Fields with * are mandatory					
Arm Information*	Arm Information				
Arm Title	Arm Title				
Arm Type	Experimental Active Comparator				
	Placebo Comparator Sham Comparator				
Additional Arm Type (if any)					
Arm Description*	Arm Description				
Interventions*	Interventions				
Interventions Type*	Drug: Including placebo Device: Including sham Biological/Vaccine Sham Comparator				
	Radiation Rehavioral				
	Genetic Dietary Supplement				
	Combination Product Diagnostic Test				
Additional Interventions Type (if any)					
Interventions Name(s)*	Interventions Name				
Other Interventions Name(s)*	Other Interventions Name				
Interventions Description*	Interventions Description				
interventions beautpiton	ne ventora besciption				
Arm or Group/Interventional Cross Reference*	Arm or Group/Interventional Cross Reference				

7 . Outcome Measures

	Fields with	* are mandatory		
t is mandatory to have o	ne primary outcome measures, You can	have multiple outcome	e measures under o	thers.
Outcome measure*	Primary O	thers		
Title*	Enter title			
Description*	Enter descriptio	n		
				1
Time frame*	Enter time frame			
	8	Add		
	Outcome Measu	res added successfully		
Outcome measure	Title	Time frame	Description	Action

8 . Eligibility

Fields with * are mandatory					
Gender Based*	No: Eligibility is not based on gender 🗸 🔻				
Age Limits - The minimum and maximum age of potential participants eligible for the clinical study provided in relevant unit of time					
Minimum Age*	34				
Unit Of Time*	Months v				
Maximum Age*	34				
Unit Of Time*	Years 🔻				
Eligibility Criteria	Eligibility Criteria	1			
		2			

	Fields with * are mandatory
User type*	Central Contact Person 🔻
e name or title, toll-free telepho cation of the study can be addre:	ne number and email address of a person to whom questions concerning enrollment at any sed
First name*	CCP first name
Middle initial	Middle initial
Last name*	CCP last name
Title*	Mr
Email*	ccp-official@mail.com
Telephone*	974112233445
Fax	974 XXXXXX
Position*	Senior
Highest degree*	D. Pharm 🔻

Please chose user type "Central contact person" add the details for the central contact person and click on "Add/Update"

Data added successfully						
🖺 Add/update						
User type	Name	Position	E-mail address	Action		
Central Contact Person	Ccp First Name Ccp Last Name	Senior	ccp-official@mail.com	🕑 Edit 🗍 🛈 Delete		

Please choose "Overall Study official" and provide information about all the study officials and their role

9. Study Contact Person and Facility Information

Fields with * are mandatory					
User type*	Overall Study Officials				
Person(s) responsible for the overall scientific leadership of the protocol, including study principal investigator. Include the following information					
First name*	OSO first name				
Middle initial	Middle initial				
Last name*	OSO last name				
Title*	Mr				
Email*	oso-offical@mail.com				
Telephone*	974112233445				
Fax	974 XXXXXX				
Role*	Study Chair Study Director Study Principal Investigator				
Highest degree*	MD, Ph.D.				

Click on "Add/update" to add the details

Data added successfully

🖺 Add/update

User type	Name	Position	E-mail address	Action
Central Contact Person	Ccp First Name Ccp Last Name	Senior	ccp-official@mail.com	🕑 Edit 🗍 🖞 Delete
Overall Study Officials	Oso First Name Oso Last Name	Study Director	oso-offical@mail.com	🕑 Edit 🗍 🖞 Delete

Facility Information in	a clinical study	
Definition : for each participating facility in a clinical study, add the following information		
Facility information	Select	٣
Unit name		

Please add details about all the participating facilities

	Fields with * are mandatory	
Contract/Agreement with the pharmaceutical company*	Please attach file (.PDF Only)	ত্রি Browse
Contract/Agreement with a CRO (if applicable)*	Please attach file (.PDF Only)	ত্রি Browse
Clinical Trial Protocol*	Please attach file (.PDF Only)	ত্রি Browse
Investigators Brochure*	Please attach file (.PDF Only)	🔯 Browse
Informed Consent (Arabic/English)*	Please attach file (.PDF Only)	ত্রি Browse
Available safety information*	Please attach file (.PDF Only)	🔯 Browse
Case Report Form / All instruments for data collection*	Please attach file (.PDF Only)	ত্রি Browse
Local IRB approval Letter*	Please attach file (.PDF Only)	ত্রি Browse
the clinical trial conducted in other	○ Yes ● No	

Please make sure to attach all the required documents.

Note: if any for the requested documents is not available, please attach a supporting document/clarification for the non-availability of the required document

Fields with * are mandatory					
ls DSMB required*	• Yes O No				
DSMB charter*	Please attach file (.PDF Only)	ि Browse			
DSMB report after initial meeting*	Please attach file (.PDF Only)	🕅 Browse			
Member first name*	Enter member first name				
Member last name*	Enter member last name				
Degree*	Select	~			
Role*	🔿 Chair 🔿 Member 🔿 Alternate				
Affiliated with the institution*	O Yes O No				
Scientific*	⊖ Yes ⊖ No				
Gender*	○ Female ○ Male				
Professional specialty*	Enter professional specialty				
Represented capacity*	Enter represented capacity				
Curriculum Vitae(CV)*	Please attach file (.PDF Only)	ি Browse			
ender*	O Female O Male				
	man and a second second				

Enter professional specialty	
Enter represented capacity	
Please attach file (.PDF Only)	🔯 Browse
Please attach file (.PDF Only)	🕅 Browse
🖺 Add	
on available	
🖺 Add/save 🕻 Submit	
	Enter professional specialty Enter represented capacity Please attach file (.PDF Only) Please attach file (.PDF Only) E Add on available E Add/save C Submit

Please click on "Submit" to submit for review, click on "Add/Save" if you wish to submit later

Health Research Registry

Click on "Register a health research" to register a research

Health Research Registry

								Register a health resear
Saved/	/Submitted application	ons						-
# R	teference no.	Research title	Research type	Fund type	Submitted on	Status	Action	
1 R	R-2019-0007	Regitry	Human Research	Intramural	21/03/2019	Under Process	Comment	
2 R	R-2019-0006	VXCV	Animal Research	Intramural	13/03/2019	Saved	🗹 Edit 🖾 Comment	
Compl	leted applications							-
No a	pplications available							

	Fields with * are mandatory	
Title*	Regitry	
tem number*	1	
Type of fund*	Intramural Fund O Extramural Fund	
Type of research*	Human Research	~
Area*	Basic medical research	~
Name of the lead investigator*	Shouketh A Researcher	~
Contact information of lead investigator	Contact Number : 77103953 ,	

Fields with * are mandatory		
Research involving vulnerable population s)*	Yes	~
Describe the vulnerable populations(s) nvolved in the project *	✓ Children □ Pregnant women □ Cognitively impaired □ Prisoners □ Disabled □ Economically	
Additional research involving vulnerable population (If any)*	N/A	
Name of collaborative institution(s)*	3	
Name of the committee that issued the ethical approval*	IRB-001-2019	
Type of committees*	IRB	~
Type of committee review*	Exempt	~
Date of ethical approval*	3/12/2019	
Start date of research*	3/27/2019	
Date of previous continuing review*	3/27/2019	
Status*	Completed	~

Research plan*	Please attach file (.PDF Only)	🔯 Browse
	Bills575936286	Delete
nstrument used for data collection*	Please attach file (.PDF Only)	🔯 Browse
	Bills575936286	Delete
Publications*	Please attach file (.PDF Only)	छि Browse
	Bills575936286	ଅ Delete
Summary*	Explain Summary	
+	C Add/save	

Research Training and workshop

To register a Research training /workshop click on "Register a training/workshop"

Research training & workshop	
	Register a training/workshop
Saved/Submitted applications	-
No applications available	
Completed applications	-
No applications available	

Fields with * are mandatory				
Title of training/workshop*	ахахх			
Location*	ахах			
Target audience*	ахах			
Training agenda*	ахах			
Start date*	12/3/2018			
End date*	11/6/2018			
.ecturers*	Please attach file (.PDF Only)	ि Browse		
	testDoc	ŵ Delete		

Serious adverse Event

Click on "Serious adverse event" to report a SAE

Serious adverse event report

No applications available

ſ	Reference no.	Clinical trial	Submitted on	Status	Action			
	SAE-2019-0008		24/04/2019	Saved		🗹 Edit	Comment	
2	SAE-2019-0007		17/04/2019	Saved		🗹 Edit	Comment	
3	SAE-2019-0009		29/05/2019	Saved		🖒 Edit	Comment	
4	SAE-2019-0001		07/02/2019	Saved		🕑 Edit	Comment	

Serious adverse event

Clinical trial registered \star

Fields with * are mandatory

Kindly report clinical trial to research@moph.gov.qa. please download clinical trial report form frombelow link

Download clinical trial report form

Clinical trial report form

Particulars of patient					
	Fields with * are mandatory				
Protocol number*					
Site name*					
Pt ID					
Date participant reported the SAE*					
SAF onset date*					
SAE stop date					
Location of SAE					
Was this an unexpected adverse event*	○ Yes ○ No				
Sex	O Male O Female				
Age	Age				
Diagnosis for study participation	Enter diagnosis for study participation				
Brief description of the nature of the	Enter brief description				
SAE*					
	🖺 Add/save				

÷

Fields with * are mandatory		
Category of the SAE*	O Date of death	
	O Congenital anomaly/birth defect	
	○ Life threatening	
	O Required intervention to prevent permanent impairment	
	\bigcirc Hospitalization- initial or prolonged	
	O Disability/incapacity	
	○ Other	
Intervention type	O Medication or nutritional supplement (specify)	
	O Device (specify)	
	○ Surgery (specify)	
	O Behavioral/lifestyle (specify)	
Relationship of event to intervention	\bigcirc Unrelated (clearly not related to the intervention)	
	\bigcirc Possible (may be related to intervention)	
	○ Definite (clearly related to intervention)	
Vas study intervention discontinued due to event	○ Yes ○ No	
What medications or other steps were aken to treat the SAE	What medications or other steps were taken to treat the SAE	
ype of report	○ Initial ○ Follow-up ○ Final	
ist any relevant tests, laboratory	List any relevant tests, laboratory data, and history, including preexisting mec	
3 · · · 3		
lata, and history, including preexisting medical conditions		
taken to treat the SAE Type of report List any relevant tests, laboratory	○ Initial ○ Follow-up ○ Final List any relevant tests, laboratory data, and history, including preexisting mec	

Health Research Data Collaboration

Select the type of collaboration & fill out the complete details, please click on Add/save And please submit after saving to submit it for further processing.

Shared health research data

	Fields with * are manda	tory	
Request type*	 Request for Collaboration 	Request for healtl research data	h
)etails about your request*			h
upporting documents	Please attach file (.PDF (Only) ((C F	Browse
	🖺 Add/save 🕻 Subr	mit	

Research Publications

Research publications

	Fields with * are mandatory		
Research field *	Select		
Name of first author *			
Affiliation of the first author*			
Full list of authors			
Title of publication*			
Journal			
Year of publication *			
Number of pages *			
Upload document *	Please attach file (.PDF Only)		
	Add/save		

Technical Support -

For any technical queries please send an email to - rdhelpdesk@moph.gov.qa