Please complete this form for Hospital-based Research. Researchers are to complete this form <u>IF</u> they checked "YES" to Question 1.6 in the TRAQ DSS FORM. Check out "*Tips Sheet for Completing TRAQ DSS FORM for Hospital-based Research*" to confirm whether your project is considered "Hospital based-Research".

Information from this form will provide hospital departments the information they need to determine if they can support the study and to ensure smooth and efficient implementation of your research project.

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									FORM. Check out " <i>Tips Shee</i>	
Completing	Hospi	tal Depo	artmen	tal Imp	act &	k Inforn	nation Form"	for	assistance with completing	this
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В.	Briefly describe how your research project will impact the various hospital departments, if applicable.
	If not applicable, please check <u>ALL</u> that apply: NOT APPLICABLE (only research funds will be held in the hospital/hospital research institute. The research project and/or the location of the research team is not within the hospital) NOT APPLICABLE (research will only be occurring in your designated research areas within hospital)
c.	Will hospital inpatients and/or outpatients be recruited to participate in this study?
	Yes No No
D.	If you answered "YES" in Question B, which hospital program(s), service(s) and/or clinic(s) will they be recruited from? Please remember to also select the correct Hospital Operational Director(s) (HOD) under the Approvals tab of your TRAQ DSS FORM prior to submission.
	Is the Program Manager of the hospital program(s), service(s) and/or clinic(s) where your research will be conducted aware of your research proposal? Please be advised that the Program Manager responsible for the day-to-day operation of the program, service or clinic might not be the person with signing authority. Yes No If No, please clarify. Not applicable
E.	If you answered "YES" to Question D, is the Program Manager supportive of any additional work required by hospital staff?
	Yes No If No, please clarify.







F.	Will you verify each hospital inpatients' and/or outpatients' medical records to confirm that they have not removed their consent to be contacted for research before you approach a potential participant or use the participant's personal data from PCS (i.e. chart review)?				
	Yes 🗌	No 🗌	Not applicable		
G.	Will you approach hospital inpatients and/or outpatients about their potential participation in the research project?				
	Yes 🗌	No 🗌	Not applicable		
Н.	. If you answered "YES" to Question G, please identify all individuals who will approach potential hospital inpatients and/or outpatients about their participation in the research project.				
ı.	Please specify the exact hospital resources (staff, equipment, supplies, space, medications, procedures/testing, etc.) needed <u>beyond usual care</u> currently being provided to patients, if applicable.				
	NC NC	search project and OT APPLICABLE (ho	ALL that apply: ly research funds will be held in the hospital/hospital research institute. The /or the location of the research team is not within the hospital) spital resources needed are only usual care/) search will only be occurring in your designated research areas within hospital)		
J.			of these hospital resources (staff, equipment, supplies, space, sting, etc.) will be reimbursed to the individual hospital(s), if		
	NC NC	search project and DT APPLICABLE (ho	ly research funds will be held in the hospital/hospital research institute. The /or the location of the research team is not within the hospital) spital resources needed are only usual care/)		
		JI APPLICABLE (res	search will only be occurring in your designated research areas within hospital)		







١.	for carrying out in individual hospital department(s), if applicable:				
	Study recruitment	Specimen collection (e.g. Blood/Fluids/Tissue/Swabs)		Specimen processing/lab analysis	
	Study documentation	☐ Vitals collection (e.g. BP, HR, RR, WT, HT)		Medication administration	
	Distribution/collection of self-administered questionnaires	Administering questionnaires		Informed consent process	
	Pharmacy medication preparation/storage/monitoring	☐ ECG/EEG/ECT/TMS/EMG		Direct care/exam	
	Data analysis	Other (please indicate below)			
If not applicable, please check <u>ALL</u> that apply: NOT APPLICABLE (only research funds will be held in the hospital/hospital research institute. T research project and/or the location of the research team is not within the hospital) NOT APPLICABLE (only research staff will carry out activities within individual hospital departm NOT APPLICABLE (research will only be occurring in your designated research areas within hospital department (s), if applicable:				rithin the hospital) n individual hospital departments) ed research areas within hospital)	
	Study recruitment	Specimen collection (e.g. Blood/Fluids/Tissue/Swabs)		Specimen processing/lab analysis	
	Study documentation	☐ Vitals collection (e.g. BP, HR, RR, WT, HT)		Medication administration	
	Distribution/collection of self-administered questionnaires	Administering questionnaires		Informed consent process	
	Pharmacy medication preparation/storage/monitoring	☐ ECG/EEG/ECT/TMS/EMG		Direct care/exam	
	Data analysis	Other (please indicate below)			
	If not applicable, please check A	ALL that apply: earch funds will be held in the hos	pital/	hospital research institute. The	
	research project and/or the NOT APPLICABLE (only hos	e location of the research team is	not w within	rithin the hospital) i individual hospital departments)	







Μ.	Will research participants undergo an informed consent process?
	Yes No No Not applicable
N.	If you answered "YES" to Question M, please identify all individuals who will carry out the informed consent process.
Ο.	If you answered "YES" to Question M, please explain how patient confidentiality will be protected, in compliance with applicable privacy legislation, during the consenting process?
Р.	Please describe how the research activities will be coordinated within the existing workflow in individual hospital department(s), if applicable.
If	f not applicable, please check <u>ALL</u> that apply: NOT APPLICABLE (only research funds will be held in the hospital/hospital research institute. The research project and/or the location of the research team is not within the hospital) NOT APPLICABLE (research will only be occurring in your designated research areas within hospital)
Q.	Do all members of your research team hold a Research Hospital Appointment at the hospital location(s) where the research is occurring if they are not paid research employees of the Hospital or hold hospital privileges (i.e. clinicians)?
	Yes No If No, please clarify.







R.	Please provide any additional information that may be relevant to assist hospital operational directors in making a decision about approval for your research project.							
L	research project and/or the	ALL that apply: earch funds will be held in the hospital/hospital research institute. The e location of the research team is not within the hospital) will only be occurring in your designated research areas within hospital)						
S.	Connell 4? Please note: to access the Cer (Pls, study nurses, study coord training. (e.g., WJHCPOR Gene Canada Division 5 training if co	derson Centre for Patient Oriented Research (WJHCPOR) on htre for Clinical Research/Clinical Trials all study team members linators, students and trainees) are required to complete various eral Orientation Training, Good Clinical Practice (GCP) and Health enducting Drug Trials, WJHCPOR Lab training and Queen's Biosafety e Room and/or Freezer Room). Contact Lisa McAvoy for more information regarding training.						
	Yes No No	Not applicable						
	Please check off the rooms/equipment you will be using to carry out your research (rooms marked with an (*) must be booked through KHSC email Outlook Calendar):							
	☐ Interview Room *	Centrifuge Room						
	Exam Room *	Cardiac Monitor						
	☐ Clinical Investigation Unit Infusion Chair *	Clinical Investigation Unit Phase I bed *						
	☐ Minor Procedure Room *	Short term freezer Room (Max. 12 Months)						

REMINDER NOTES:

✓ Some hospital departments may require additional information to be collected before approval will be granted. If additional information is required, the hospital operational director(s)/research director(s) will reach out to you once your TRAQ DSS FORM is submitted and received in their queue.







- ✓ It is important to consult (reach out via email or telephone) with hospital operational director(s)/research director(s) early in your proposal/protocol and budget development to ensure budgets are accurate when applying for grants or negotiating industry contracts and hospital resources are required.
- ✓ If there is urgency for your TRAQ DSS FORM to be reviewed and approved, please reach out to the respective hospital operational director(s)/research director(s) via email or telephone to let them know. TRAQ DSS FORMS are required to be submitted at least 15 business days before any internal/external deadlines to ensure all approvals are in place.
- ✓ Researchers <u>are required to</u> have all necessary certifications (i.e. human ethics, animal care, biohazards, and radiation) and TRAQ DSS FORM approvals in place before commencing research projects. Once all necessary certifications are in place, please upload all approval letters to your TRAQ DSS FORM to ensure all hospital operational director(s)/research director(s) can obtain a copy.
- ✓ Researchers using hospital labs, pharmacy, and/or clinical engineering are required to complete the
 additional study request form and attach to their TRAQ DSS FORM prior to submission under
 "Attachments".





