



Health Research Ethics Application Form V.22

Project	Info.
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File No: Ref No: -1

Project Title:

Principal Investigator: ()

Start Date: End Date: Keywords:

Question	Answer
Is the PI conducting research on behalf of Island	
Health or external?	
If the PI is not from Island Health, please	
provide the name of the Island Health	
collaborator. All studies must have at least one	
Island Health affiliated team member.	
If PI is from Island Health what is their	
department?	
If External Researcher, do they have Island	
Health affiliation/privileges?	
Study nickname or acronym (if applicable):	
Type of funding for this research study; if for-	
profit funded, please complete the funding tab in	
this application.	
Provide name of the funding agency,	
department or industry sponsor (clinical trials).	
For funded studies, please provide name of the	
institution where the funds will be held:	
Enter any applicable information about your	
funding which is not already included (including	
funding applied for but not yet received).	
Is the study funded by the US Department of	
Health and Human Services (DHHS)?	
If yes, please indicate which DHHS funding	
agency.	
If this submission is part of an academic	
program please provide the name of the	
institution, supervisor, and program.	

Please describe how you will disseminate the	
results of the research study. Include if and how	
you will target specific knowledge users, and	
any plans to report results back to participants.	
If participants will not receive a report of study	
results, please explain why not.	
Do you consent to being contacted by a	
member of the Island Health Research and	
Capacity Building team regarding the	
development of dissemination strategies?	
Identify where the research will be carried out at	
Island Health (hospital, department, clinical	
area, health centre, etc.).	
Name the Island Health hospital(s) involved:	
Name the Island Health health centre(s)	
involved:	
Name the Public Health Unit(s) involved:	
Will data be sent outside of Island Health?	
(transferred)	
Will the study require any non-standard devices	
to be connected to Island Health's network?	
If yes, please describe the device, its technical	
safeguards, and who will be using it.	
Project Team Info.	

Project Team Info.

Principal Investigator

Prefix:
Last Name:
First Name:
Affiliation:
Position:
Email:
Phone1:
Phone2:
Fax:
Primary Address:
Institution:
Country:

Common Questions

Comments:

1. 1. How to Complete the Application

#	Question	Answer
	The Health Research Ethics Board (HREB)	
1.1	reviews research that is predominantly	
	behavioural or social sciences related.	
1.2	Mandatory Fields	
1.3	Which Tabs MUST be completed	
1.4	Who Can Submit	
1.5	Documents	

2. 2. Review Process and Timelines

#	Question	Answer
2.1	Institutional Approval	
2.2	Timelines	
2.3	After Initial Approval	

3. 3. Principal Investigator and Study Team

#	Question	Answer
3.1	Other than the PI, list all names and	
	primary affiliation required to be on the	
3.1	Certificate of Approval. (e.g. Dr. Jane Doe,	
	UBC)	
	Enter the Principal Investigator's secondary	
3.2	appointments or affiliations (including Post-	
3.2	Secondary and Health Authorities) if	
	applicable	
	Describe each study team member's (co-	
	investigator, staff, research assistant,	
	external supervisor, consultant, etc.) role in	
3.3	the study e.g. statistician, supervisor,	
	advisor, student, etc. Ensure each	
	individual is entered in the first box of this	
	tab.	
	All Investigators and study team members	
	are required to update their TCPS 2	
3.4	training to the CORE-2022 version. Please	
3.4	confirm that all study team members,	
	including the Investigator, have completed	
	this training.	
	Please describe any special training	
3.5	requirements or qualifications required for	
	the study team to conduct the study.	

4. 4. Study Funding Information

#	Question	Answer
4.1	Please provide the funding title (if different	
4.1	than the project title). (N/A if not applicable)	
4.2	Please list the type of funding for this	
4.2	research study.	
4.3	If other, please describe	
	Please provide the name of the funding	
4.4	agency, department or industry sponsor.	
	(N/A if not applicable)	
	For funded studies, please provide name of	
4.5	the institution where the funds will be held.	
	N/A if not applicable	
	Enter any applicable information about	
1.6	your funding which is not already included	
4.6	(including funding applied for but not yet	
	received).	
4.7	Is the study funded by the US Department	
4.7	of Health and Human Services (DHHS)?	
4.0	If yes, to above, please indicate which	
4.8	DHHS funding agency.	

5. 5. Conflict of Interest

# Question	Answer
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	-	
	Study Related Conflict of Interest Conflicts	
	of Interest (COIs) in research are situations	
	where someone's personal interests	
	(financial, career, or other) could	
	compromise or could be perceived to	
	compromise the objective conduct of	
	research or integrity of the data. Conflicts	
	of interest can arise naturally from an	
	Investigator's engagement inside and	
	outside the health authority, and the mere	
	existence of a COI or the perception of a	
	COI does not necessarily imply wrongdoing	
	on anyone's part. Nonetheless, real and	
	perceived COI must be recognized,	
5.1	disclosed, and assessed. This question	
	asks Investigators to disclose COIs that	
	may relate to the research study that is the	
	subject of the REB application. Do the	
	Principal Investigator, Co-Investigators	
	and/or their related parties have any	
	personal interest(s) that could compromise	
	or reasonably be perceived to compromise	
	the objective conduct of the research or the	
	integrity of the data generated by the	
	study? Personal interests may include	
	business, commercial or financial interests,	
	dual roles (e.g. PI and Doctor), as well as	
	personal matters and career interests.	
	You have answered yes, please complete	
	this section. Refer to the Island Health	
5.2	Conflict of Interest Policy and Disclosure	
0.2	form for next steps and attach the	
	Disclosure form once completed to the	
	application.	
	Please describe the conflict of interest	
5.3	(COI) including dollar value where	
	applicable.	

	Do any of the researchers conducting this	
	study occupy more than one role with	
	respect to potential participants (e.g. acting	
	as both a researcher and a therapist,	
5.4	health care provider, caregiver, teacher,	
3.4	advisor, consultant, supervisor, manager,	
	student, or employer, etc.) that may create	
	a real, potential, or perceived conflict of	
	interest that could affect the integrity of the	
	research?	
5.5	If yes, please describe	
	Please advise how you propose to manage	
5.6	any actual, perceived, or potential COI	
	outlined above.	
5.7	If applicable, please identify mitigation for	
5.7	any possible 'power over' relationships.	

6. 6. Study Type

#	Question	Answer
	Indicate whether your application is a	
6.1	Retrospective Chart Review or Health-	
	Behavioural	
	Please enter any other locations where the	
	research will be conducted under this	
6.2	Research Ethics Approval (e.g., Name of	
0.2	privately owned clinic, community centre,	
	school, classroom, participant's home, in	
	the field - provide details).	

7. 7. Review Type and Risks

#	Question	Answer
	Relationship to Previous Ethics	
	Applications: If this proposal is closely	
	linked to any other proposal	
7.1	previously/simultaneously submitted, enter	
	the Institution or Health Authority name and	
	associated Research Ethics Board study	
	number of that proposal.	
	If applicable, please describe the	
7.2	relationships between this proposal and the	
1.2	previously/simultaneously submitted	
	proposal listed above.	

	Are you aware of any rejection of this study	
	by any Research Ethics Board? If yes,	
7.3	please provide known details and attach	
	any available relevant documentation	
	under Attachments.	
	Peer Review Has the research proposal	
	received any independent	
7.4	scientific/methodological peer review? All	
	above minimal risk studies require a peer	
	review.	
7.5	External peer review details including	
7.5	name of individual:	
7.6	Internal peer review details including name	
7.0	of individual:	
	If this research proposal has NOT received	
7.7	any independent scientific/methodological	
'	peer review, explain why no review has	
	taken place:	
	After considering the level of risk your	
	research involves and the vulnerability of	
7.8	your study population, please tick one box	
	below that best represents the overall level	
	of risk.	
	Provide an explanation for the assessment	
7.9	of research risk and group vulnerability	
	reported above.	
	Does your application fall under minimal	
	risk? (eg. was it assigned an overall risk	
7.10	level of 1 on the minimal risk matrix. Please	
	see the risk matrix included in the guidance	
	document.)	
	Does this study require review and	
7.11	approval by another Canadian REB outside	
'.''	of Research Ethics British Columbia	
	(REBC)?	

8. 8. Summary of Study and Recruitment

#	Question	Answer
	Provide a brief statement about the project	
8.1	written in lay language. Do not exceed 100	
0.1	words and do not cut and paste directly	
	from the study proposal.	

	Summarize the research proposal including	
8.2	study purpose, hypothesis, study	
	population, and research method.	
	Inclusion Criteria - Describe the	
8.3	participants being selected for this study,	
	and list the criteria for their inclusion.	
	Exclusion Criteria - Include details if	
8.4	otherwise eligible participants will be	
0.4	excluded due to other characteristics. If no	
	exclusion criteria are applicable, enter n/a.	
	Recruitment Provide a detailed	
8.5	description of the steps you will use to	
0.0	recruit participants. Include: Who will	
	contact the prospective participants	
	Recruitment Provide a detailed	
	description of the steps you will use to	
8.6	recruit participants. Include: By what	
	means will recruitment be done (e.g.,	
	public posting, third party recruitment etc.)?	
	Recruitment Provide a detailed	
8.7	description of the steps you will use to	
	recruit participants. Include: How will	
	prospective participants be identified	
	Recruitment Provide a detailed	
8.8	description of the steps you will use to	
	recruit participants. Include: all site specific	
	information.	
	Recruitment Provide a detailed	
	description of the steps you will use to	
8.9	recruit participants. Include: Attach all	
	materials, including letters of initial contact,	
	posters, scripts and advertisements in the	
	attachments tab.	
	Use of records: If existing records (e.g.	
	health records, clinic databases,	
	registration details, etc.) will be used	
8.10	access information about potential	
	participants, please describe how	
	permission to access the information, and	
	to collect and use the information, will be	
	obtained.	

	Summary of Study Procedures Describe	
8.11	briefly in a step-by-step manner what the	
	researcher will be doing with participants,	
	after then have been recruited and	
	consented.	
	Research Types Select all that apply to	
	your study. Please review the research	
8.12	methods descriptions before responding. If	
	none apply, please select 'None of these	
	Methods'	
8.13	If other, please describe:	

9. 9. Participant Information and Consent Process

#	Question	Answer
9.1	How much time will a participant be asked	
3.1	to dedicate to the project?	
	Describe what is known about the risks of	
9.2	the proposed research for participants and	
	how it will be mitigated.	
	Describe any potential benefits to the	
9.3	participant that could arise from	
	participation in the proposed research.	
	If your research involves an identified	
9.4	group or community, outline the likely	
	impacts of the research on the community.	
	Specify how potential participants will be	
	invited to take part in the study. Include	
9.5	details of where the consent will be	
	obtained and documented, and under what	
	circumstances.	
	If applicable please describe the	
9.6	community consent process. If no	
9.6	community consent is being sought, please	
	justify.	
	Describe any reimbursement and	
	incentives (e.g., meals, parking,	
	medications) or payments/gifts-in-kind	
9.7	(e.g., honoraria, gifts, prizes, credits) to be	
	offered to the participants. Provide full	
	details of the amounts, payment schedules,	
	and value of gifts-in-kind.	
	Obtaining Consent - Include details of	
9.8	where and when consent will be obtained	
3.3	and how it will be documented.	
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	If you are asking for a waiver or an	
	alteration of the requirement for participant	
	informed consent please justify the waiver	
9.9	or alteration and confirm that the study	
	meets the criteria below in yellow box.	
	Please address each criterion individually.	
	How long after being provided with detailed	
	information about the study will the	
9.10	participant have to decide whether or not to	
	participate? Provide your rationale for the	
	amount of time given.	
	Will every participant have the capacity to	
9.11	give fully informed consent on their own	
	behalf?	
9.12	If no, please provide the details of the	
0.12	nature of the incapacity:	
	If a participant does not have the capacity	
9.13	to give fully informed consent, who will	
	consent on their behalf?	
	If a participant does not have the capacity	
9.14	to give fully informed consent, will they be	
	able to give assent to participate?	
0.45	If Yes, explain how assent will be sought.	
9.15	Please be sure to attach copies of the	
	assent form under the Attachments tab. Describe any situation in which the	
	•	
9.16	demonstration of ongoing consent for this	
	research might be appropriate, and how	
	this would take place. What provisions are planned for	
	participants, or those consenting on a	
	participant's behalf, to have special	
9.17	assistance, if needed, during the consent	
	process (e.g., consent forms in Braille, or in	
	languages other than English).	
	Describe any restrictions regarding the	
	disclosure of information to research	
	participants (during or at the end of the	
9.18	study) that the funder/sponsor has placed	
	on investigators, including those related to	
	the publication of results.	
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10. 10. Number of Participants and Locations for Behav ...

#	Question	Answer

10.4	Does this research focus on Indigenous	
10.1	peoples, communities or organizations?	
	Will the research be conducted on	
	Indigenous reserves, Métis settlement(s),	
10.2	or lands governed under a self-government	
	agreement or an Inuit or First Nations land	
	claims agreement?	
	Do any of the criteria for participation	
	include membership in an Indigenous	
10.3	community, group of communities, or	
	organization, including urban Indigenous	
	populations?	
	Does the research seek input from	
10.4	participants regarding a community's	
10.1	cultural heritage, artifacts, traditional	
	knowledge or unique characteristics?	
	Will Indigenous identity or membership in	
10.5	an Indigenous community be used as a	
	variable for the purposes of analysis?	
100	Will the results of the research refer to	
10.6	Indigenous communities, peoples,	
	language, history or culture?	
	Community Engagement If you	
	answered yes to questions above, have	
10.7	you initiated or do you intend to initiate an	
	engagement process with the Indigenous	
	collective, community or communities for	
	this study? If you answered Yes please describe the	
	process that you have followed or will	
	follow with respect to community	
100	engagement. Include the role or position of	
10.8	those consulted, including their names if	
	appropriate. Attach any documentation of	
	consultations (i.e. formal research	
	agreement, letter of approval, email	
	communications, etc.) below. No community consultation or engagement	
	If you answered no to the previous	
	1	
10.9	question, briefly describe why community	
10.9	engagement will not be sought and how	
	you can conduct a study that respects	
	Indigenous communities and participants in	
	the absence of community engagement.	

	If your research involves an identified	
10.10	group or community, outline the likely	
	impacts of the research on the community.	
	If applicable please describe the	
10.11	community consent process. If no	
10.11	community consent is being sought, please	
	justify.	
10.12	Registration for Publication of Clinical	
10.12	Trials	
10.13	If yes, please enter the following	
10.13	information:Has the study been registered?	
10.14	Authorized Registry used:	
10.15	Clinical Trial unique identifier:	
	Number of Participants How many	
10.16	participants will take part in the entire study	
	(eg. World-Wide)?	
	How many participants will take part at	
10.17	institutions covered by this Research	
	Ethics approval?	
10.18	Principal Investigator and Research Team	
10.16	Experience:	

11. 11. Security of Data and Confidentiality of Person ...

#	Question	Answer
11.1	Please select all tools for data collection.	
11.2	If other, please describe:	
11.3	Please describe who on the study team will	
11.5	be doing the data collection:	
	It is the PI's responsibility to ensure that all	
	members of the study team who will be	
	accessing data are made aware of their	
11.4	responsibilities concerning privacy and	
	confidentiality. Explain who will have	
	access to the data at each stage of	
	processing and analysis.	
	Describe how the identity of the	
	participants will be protected both during	
11.5	and after the research study, including how	
	the participants will be identified on data	
	collection forms.	
	If a study code/key/master list will be	
	created to link each participant to the data	
11.6	being retained, please describe who is	
	keeping the list, where, and what	
	safeguards there are to protect the list.	

11.7	Will any personal health information or	
11.7	personal identifiers be collected?	
	If yes, please describe what personal	
11.8	identifying information will be collected, and	
	justify the need for it to be collected.	
	How and where will data be stored? (E.g.,	
11.9	computerized files, hard copy, videotape,	
11.9	audio recordings, personal electronic	
	communications device, other.)	
	If data will be sent outside of the Institution	
	where it originated, please describe the	
11.10	type of data to be transferred, who the data	
	will be transferred to, where the data will	
	transferred, and how the data will be sent.	
	If data will be received from other sites,	
11.11	please describe the type of data, where it	
''''	will be received from, and how the data will	
	be received.	
	If data will be linked to any other data	
	source (including a biorepository) please	
11.12	identify the data set, how the linkage will	
11.12	occur, and explain how confidentiality	
	regarding the shared information will be	
	preserved.	
	Describe any data that will be sent to,	
11.13	accessed from, or stored outside of	
11.13	Canada. Include details of where it will be	
	sent to or accessed from, and the purpose.	
11.14	Describe the safeguards in place to protect	
11.14	the confidentiality and security of the data:	
	If any data or images are to be kept on the	
11.15	Web, what precautions have been taken to	
	prevent them being copied?	
	Describe what will happen to the data at	
11.16	the end of the study (including how long	
''''	the study data will be retained, when and	
	how the data will be destroyed)	
	If there any plans for future use of either	
11.17	data or audio/video recordings please	
11.17	provide details, including who will have	
	access and for what purposes, below.	

11.18	Is this application for research requiring	
	access to clinical charts OR data from	
	registries or databases such as	
	PopDataBC or Pharmanet?	
11.19	If data will be collected from health records	
	at Island Health, please identify, by name,	
	who will be accessing the health record.	
11.20	Insert the date range of the charts/data to	
	be included in the research:	

12. 12. Retrospective Chart Review

#	Question	Answer
12.1	Is this a retrospective chart review study for	
	which participant consent will be obtained?	
12.2	Describe how permission to access the	
	medical records and to collect and use	
	these records will be obtained.	
	Briefly describe the type of data that you	
12.3	intend to collect (e.g., disease, diagnosis,	
12.3	outcome, demographic, aggregate,	
	personal-level).	
12.4	Number of Records/Patient Charts	
	Are you collecting and retaining personally	
12.5	identifiable information to be a part of the	
	data set?	
	Indicate what personally identifying	
12.6	information you will be collecting and	
12.0	retaining as part of the dataset. Include a	
	justification of why it is required.	
12.7	Please explain if and why the identifiable	
12.7	information essential to the research.	
	Please explain how the use of the	
	identifiable information without the	
12.8	participants consent is unlikely to adversely	
	affect the welfare of the participants to	
	whom the information relates.	
	Please explain how the researchers will	
400	take appropriate measures to protect the	
12.9	privacy of individuals and to safeguard the	
	identifiable information.	
12.10	Please explain how the researchers will	
	comply with any known preferences	
	previously expressed by individuals about	
	any use of their information.	
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	Please explain why it is impossible or	
12.11	impracticable to seek consent from	
	individuals to whom the information relates.	
	Please describe how the researchers will	
12.12	obtain any other necessary permissions for	
12.12	secondary use of the information for	
	research purposes.	
	Describe the risks associated with the	
	possible disclosure of the data. Include any	
12.13	foreseeable circumstances where	
	disclosure of identifying data may be	
	required by law.	
	Describe how the identity of the	
	participants will be protected both during	
12.14	and after the research study, including how	
	the participants will be identified on data	
	collection forms.	
	Explain who will have access to the data at	
	each stage of collection, processing and	
	analysis, and indicate whether a current list	
12.15	of the names of study personnel (including	
	co-investigators) and their delegated tasks	
	will be maintained in the study file. If a list	
	will not be maintained, please explain why.	
	Describe how and where the data will be	
12.16	stored (e.g., computerized files, hard copy,	
	video-recording, audio-recording, personal	
	digital device, other)	
	Describe what will happen to the data at	
	the end of the study, including how long the	
	data will be retained and where, when and	
12.17	how the data will be destroyed, and what	
	plans there are for future use of the data,	
	including who will have access to the data	
	in the future and for what purpose.	
12.18	Will data be transferred out of the custody	
	and control of Island Health?	
12.19	If data will be transferred out of Island	
	Health, please describe a) the type of data	
	to be transferred, b) who the data will be	
	transferred to, c) where the data will be	
	transferredd) how the data will be sent.	
12.20	Do you plan to link the data to any other	
	data?	

	If yes to data linkage:a) Identify the data
	set, b) how the linkage will occur, c)
	provide a list of data items in the other
12.21	database. d) identify what personal
	information will be used to link the
	databasese) how confidentiality regarding
	this shared information will be preserved.

13. 13. Attestations

I attest that the information provided in this form is accurate and up to date at the time	
13.1 form is accurate and up to date at the time	
of submission.	
I agree to conduct the study in accordance	
with the Tri-Council Policy Statement:	
Ethical Conduct for Research Involving	
Humans – TCPS 2 (2022)	
13.3 I agree to conduct the study in accordance	
with the REB approved documents.	
I have read, understood, and agree to	
abide by the Island Health policies and	
procedures regarding the conduct of	
13.4 research: specifically Policy 25.2 Free and	
Informed Consent in Research, Policy 25.3	
Research Integrity Policy, and (if	
applicable) 705 Research Finance Policy	
13.5 I agree that Island Health may conduct a	
compliance audit of this study.	
I will monitor and abide by all provincial	
and institutional safety requirements in	
order to uphold the rights and welfare of	
research participants as COVID-19	
13.6 circumstances evolve. I will ensure that	
additional safety requirements such as	
contact tracing, affirming vaccination status	;
of participants, and other measures are	
performed if or when necessary.	
Principal Investigator Signature: By signing	
this application electronically, I understand	
that my electronic signature has the same	
legal effect and can be enforced in the	
same way as a written signature. Please	
enter your name.	

14. Please submit an operational application to pair w ...

#	Question	Answer
14.1	If Island Health staff/physicians will be	
	involved in the recruitment of participants	
	for the study, please describe what the	
	involvement will entail.	
14.2	If Island Health staff/physicians will be	
	involved in any other part of the conduct of	
	the study, please describe what that	
	involvement will entail.	
14.3	Please identify all departments where	
	personnel will be requested to support the	
	study. If department name is unknown,	
	please identify the type of support required.	