

# School of Computing RESEARCH ETHICS COMMITTEE

# APPLICATION FORM FOR ETHICAL REVIEW OF A RESEARCH PROJECT INVOLVING HUMAN PARTICIPANTS WHICH IS IN THE CATEGORY OF NOTIFICATION ONLY

There are 3 generally accepted levels of ethical review for projects carried out in a University or similar setting. These are notification only, expediated and full committee.

This notification only level of review is to approve relatively low-risk research involving human participants, primarily using social science methodologies in which any personal information collected is not of a sensitive nature. The School of Computing Research Ethics Committee has been delegated responsibility by the University to approve ethics submissions from undergraduate and taught Masters projects only, which are in the category of notification only.

Examples of projects in this category include:

- Anonymous surveys in which the topic itself is not likely to elicit significant difficulties for the participants, such as: anonymous internet surveys (e.g. Survey Monkey), street questioning.
- Observation (without audio or visual recording) of public settings where privacy would not normally be expected, such as observing people on streets or at sports events.
- Research carrying no risks beyond those of everyday life (as experienced by the intended participant population), such as asking people's opinions about products or services; asking students about educational experiences; monitoring the impact of daily activities.
- Interviews with public figures, professionals or others in their professional capacity regarding their professional activities.
- Analysis of data (e.g. health records) which have had all identifying information removed by the data holder and been provided to the researcher in accordance with data protection legislation.
- Collection of biological samples which are anonymised and do not require invasive techniques (e.g. hair, nails).

If your project is using data from a public repository like Kaggle or is not generating or using any form of personal data then you do not need research ethics approval, you do not need to complete and to submit this form and your project supervisor should indicate this on the project dashboard.

If your project involves collecting or processing <u>personal data which is of a personal nature</u>, you must first complete the DCU online Data Protection training course and review the <u>"Data Protection – Key Points for DCU Researchers"</u> guidance from the Data Protection Unit to assist you in meeting your legal obligations under GDPR and associated Irish law.

Once you have completed this form (if you need to) you should save it as a PDF file, not WORD, and upload it to the your project dashboard before you start gathering data. It will then be read and assessed by two members of the committee and once two members of the committee approve your submission you will be automatically notified by email and your project can start data gathering.

There are strict deadlines for submitting this form for each class group, undergraduate and taught Masters by which your submission must be made and you will be informed of these deadlines by your course board chair or project co-ordinator. If you do not submit by these deadlines then the research ethics committee is not obliged to approve your submission and when that happens and your project is assessed and graded at the end of the year, you will be awarded 0 for that component of your project.

SECTION 1 – GENERAL DETAILS				
1.1 Project Title				
Prospector – Provisioning	of Containers and V	Ms		
1.2 Applicant Details				
Name	Student or Supervis	or	E-mail	
Alexandru Dorofte	Student		alexandru.dorofte2@mail.dcu.ie	
James Hackett	Student		james.hackett5@mail.dcu.ie	
Stephen Blott	Supervisor		stephen.blott@dcu.ie	
Other Investigators: Including	any external to DCII			
Other investigators. Including	arry external to Doo			
Name	School/Unit/Externa	l Institution	E-mail	
1.3 Key Project Dates				
Proposed start date for data	Proposed end date	for data	Proposed project	
collection	collection		completion date	
15 <sup>th</sup> March 2024	20 <sup>th</sup> April 2024		21st April 2024	
A A Disease in disease which are denote accord				
1.4 Please indicate which academic award				
Undergraduate x	lau	ght Masters 🔲		

### 1.5 Please confirm the location(s) where the research will be carried out

If research will be carried out abroad, you will need to address the ethical challenges raised by this in Section 3 of your application - consult the Conducting Research Abroad document in the Ethics Resources and Guidelines section of the <u>DCU Research Ethics webpage</u>).

Research Conducted Remotely Online	

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Specify from whom the permission is required (e.g. a school Board of Management), and when their written approval will be obtained

Willer approval Will be establed	<i>a</i>		
N/A			

#### SECTION 2 - PROJECT DESIGN AND METHODOLOGY

Research Overview - Please respect the indicated word counts in the following sections and explain all acronyms in full text the first time they appear.

#### 2.1 Provide a brief description of the research (max 250 words):

Please use lay language, include the scientific/theoretical background of study and a justification as to why this research project should proceed in that context

The research study aims to view if a user finds the accessibility of the project usable to their satisfaction. Or to view any potential optimisations that can be made to further satisfy the users' requirements.

#### 2.2 Please state the aims and objectives of the project (max 200 words)

The aims of the project is to deliver on 2 fronts:

- 1<sup>st</sup>: For a user who in this case is a systems administrator where they are given this platform to manage users in a cluster and impose quotas and view systems health all in the comfort of one place Prospector
- 2<sup>nd</sup>. For the user that is given access by the systems administrator to manage their jobs to run their personal projects/services within prospector without the need of an admin to constantly manage them.

#### 2.3 Please confirm your methods of data collection:

Tick all relevant check boxes and provide details for each one, including any devices used to collect data, and whether the data will be anonymous, potentially identifiable or identifiable at point of collection

Method	Describe briefly
Interviews or focus groups	
x Surveys/questionnaires	The applicant will be asked to access Prospector and follow a simple task, and after answer qs regarding their satisfaction and if there was any difficulty in completing the task and provide potential areas to improve.
Audio/video recordings	
Public observations	
Persons in public office	
☐Using existing data (incl.	
secondary data)	
☐Using human derived	
material (biological samples)	
Standard tests	
(educational/personality etc.)	
Standard educational	
practices	
Other (please specify)	

2 / Please confirm who	the participants on this study will be, including group size and
composition:	the participants on this study will be, including group size and
•	graphic characteristics, and state how your proposed sample size was
determined (e.g. power a	
	be computer science students and Redbrick society members.
Our sample size will be	15 students( 3 from each year 1-4, and 3 members from Redbrick)
0 F Disease suding second	
2.5 Please outline your l from and your criteria fo	recruitment process, including where you are sourcing participants
	nvolved, outline the procedures relating to their involvement
vinere gatekeepers are ii	volved, oddine the procedures relating to their involvement
Will ask students within	a Computer Lab if they would like to participate.
	e student is below 18 years of age.
	, c
2 6 Addressing particing	ant vulnerability – if your participants fall into any of the following
	k the relevant tick box/boxes and state below what special
arrangements will be ma	
If your participants are no	ot in any of these categories, tick N/A
X N/A	
Children under 18 year	
	elationships with the researcher (e.g. lecturer-student, therapist-client,
employer-employee)	
	ised or diagnosed intellectual, physical or mental impairment
	stitutions (e.g. prisoners, residents in 24 hr nursing facilities) dergone traumatic or adverse emotional events
Marginalised sections	and cognitive ability
	ed cognitive ability
	s of society
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2.7 Involvement of children under 18 years of age – if your participants are in this category, please confirm compliance with the following:

If your participants are not in this category, tick N/A
x N/A
We confirm that we have read and agree to act in accordance with the DCU Child Protection
policy and procedures (as per the <u>DCU Child Protection Unit webpage</u> )
We confirm that we have put in place safeguards for the children participating in the research
We confirm that we have supports in place for children who may disclose current or historical
abuse (whether or not this is the focus of the research)
We confirm that all requirements will be met prior to commencing the research (e.g. TUSLA
Children First Training completed, Garda Vetting in place)
2.8 Please confirm how the results of the research will be disseminated:  Include a statement on whether the participants will be provided with any information as to the findings or outcomes of the project  They will not be provided any findings or outcomes of the project.

## SECTION 3 - ETHICAL ISSUES AND RISK MANAGEMENT

3.1 Please identify all issues including ethical issues which may arise in the course of this research. What are the potential risks to participants, and how will those risks be addressed or minimised?  Potential risks can be physical, psychological, social, legal, etc. Please include details of any
additional support being provided for participants during/after the study  N/A
3.2 Please identify the potential benefits (direct and/or indirect) to those participating in this research:
Potential benefits should outweigh the potential risks to participants
N/A
3.3 Please describe what measures/protocols you have put in place in the event that there are any unexpected outcomes or adverse effects to participants arising from involvement in the
research: N/A
TV/A
3.4 Do you intend to provide payment or incentives to participants?
Yes No x
If Yes, please consult the REC Guidelines on the Use of Compensation and Incentives (in the Ethics
Resources and Guidelines section of the <u>DCU Research Ethics webpage</u> ) before providing additional details below
details below
3.5 Does this research raise any potential risks for the researchers themselves?
Please consider the location/environment where the research is being conducted, exposure to
distressing data content etc.
Yes No x

If Yes, please describe further and explain what risk management procedures will be put in place to minimise these risks to researchers:
3.6 Does this research raise any potential conflict of interest?
Please consider any potential real <u>or</u> perceived conflicts of interest that might influence the integrity
of the research, or give rise to bias in conducting and reporting the research, or affecting publication (consult the <u>DCU Conflict of Interest Policy</u> for assistance)
Yes No x
If Yes, please identify and explain the steps being taken to address that conflict:
3.7 Please describe how the conduct of the research will be monitored:
Regular oversight by the PI is required to ensure the project conforms to the procedures set out in this application (especially where several people are involved in carrying out the research procedures)
The research will be monitored by the 2 Project partners remotely, where the user will be given access to the project online and answer the questionnaire.

# SECTION 4 - CONFIDENTIALITY AND DATA MANAGEMENT

4.1 Considering your previous response in section 2.3 of the form on data collection, please
confirm whether you are collecting or processing personal data in this research project:
Personal data is any information about a living person, where that person is either identified, or
could be identified from the data itself, or when it is combined with other data. This includes paper
based, electronic and biological samples data. If your data is fully and completely anonymous, it is
<u>not</u> personal data.
Yes No x
If Yes, please confirm your compliance with the following by ticking the checkboxes:
We confirm that we have completed the DCU Data Protection training module on Loop.
We confirm that we have read the "Data Protection – Key Points for DCU Researchers"
guidance on the DCU Data Protection Unit (DPU) website and agree to protect and manage our
data in accordance with same.
We have assessed the degree of risk inherent in the personal data being used in the research
project, and confirm that all DPU GDPR requirements have been met prior to submitting this
application (e.g. completion of Data Protection questionnaire, confirmation that any survey tool
being used is GDPR compliant, that required Data Processing or Sharing Agreements will be in
place, etc.)
pidoc, cto.)
4.2 Data access - please confirm whether access to participant data is confined to the
investigators named on this application:
Yes x No
If No, please name who the other individuals are and why they need access. Any proposed transfer of
data (including outside of the EU) should be detailed here.
4.3 Data storage – please confirm compliance with the following:
Data collected on mobile devices will be protected with a strong password/passphrase at a
minimum, and/or encrypted if the device supports it
·
X Data will be removed from mobile devices as soon as is practicable and stored in a secured
location in DCU (on server or institutional Google Drive)
Paper based data will be held securely in locked cabinets in DCU, with access restricted to the
named researchers
Specific arrangements in relation to biological samples should be stated here:
Any exemptions to the above compliance statements should be justified here:
7 my exempliana to the above compilance statements another be justified field.

4.4 Please confirm who will be research:  Name the relevant DCU investig	·	storage of d	lata generated by the
	Jaio173		
Alexandru Dorofte James Hackett			
4.5 Please confirm how long t		-4- :- 41 "D -4	- Dustastian Kan
For personal data, consult section Points for DCU Researchers" gu			
10 Weeks	didance on the DCO Data Frote	CHOIT OTHE (DE	O) WEDSILE
10 Weeks			
4.6 Please confirm what will h			
Please tick the relevant checkbo			
Archived	Destroyed x	Other	
4.6.1 Archived data			
Please provide the following det	ails:		
Name the DCU staff member	N/A		
responsible for archival and			
future use of data			
Confirm whether the data will	N/A		
be made available to other			
researchers, and if so, how?			
Confirm how the data will be	N/A		
prepared for archive (e.g. will			
datasets be anonymised)			
Confirm where the data will	N/A		
be archived and who will be			
allowed to access it			
4.6.2 Destroyed data			
Please provide the following			
responsibility for data destruction the time of destruction	n il there is no guarantee the s	ludent will nav	e access to the data at
Please justify why the data	Has no research use outside	the scope of th	ne project
will be destroyed	Thas no research use outside	ine scope or ir	ie project.
Name the DCU researcher	Alexandru Dorofte		
responsible for destruction of	James Hackett		
data	- Cameron Francis		
Confirm when the data will be	10 <sup>th</sup> May 2024		
destroyed (specify date)			
Confirm compliance with the	X Electronic data will be over	written/securel	y deleted
following destruction	☐Paper based data will be c		
methods (tick relevant boxes)	☐ Medical samples will be dis		
	relevant DCU approved SOP		

4.6.2 Other - Please explain what will happen to the data if not being archived or destroyed:

N/A			

#### SECTION 5 - PARTICIPANT INFORMATION AND INFORMED CONSENT PROCEDURES

In addition to completing this form you are required to attach, within the single PDF that you submit, a copy of (1) the Participation Information Sheet which you share with your participants and (2) a copy of the Informed Consent Form which your participants sign.

5.1 Please confirm that the following items have been addressed in your Participant Information Sheet which should be shared with all participants whether it involves online or inperson data gathering:

The items below should be used as headings in yourinformation sheet. Note the language used under each item must reflect the participant age group and corresponding comprehension level— if your participants have different comprehension levels (e.g. both adult and child participants) then separate sheets must be prepared for each set. Templates are available via the <u>REC Forms</u>—Applications. Templates and Amendments section of the Research Ethics website.

Introductory Statement (Researcher names and titles, school, title of the research study)

Yes

No

Checklist - tick the relevant check box for each item

What is this research about? Why is this research being conducted?	X	
Why is this research being conducted?		
	X	
Why have you been invited to take part?		
What will happen if you decide to take part in this research study?		
How will your data be used?		
low will your privacy be protected (including any legal limits to confidentiality)?		
What are the benefits of taking part in this research study?		
Vhat are the risks of taking part in this research study?		
Can you change your mind at any stage and withdraw from this study?		
How will you find out what happens with this project?		
Contact details for further information	X	
5.2 Informed Consent Procedures – please confirm whether written consent i	is to be o	btaiı
5.2 Informed Consent Procedures – please confirm whether written consent i Please tick the relevant checkbox	is to be o	btaiı
5.2 Informed Consent Procedures – please confirm whether written consent in Please tick the relevant checkbox  Yes x  No		
5.2 Informed Consent Procedures – please confirm whether written consent in Please tick the relevant checkbox  Yes x  If Yes, describe the procedures by which written consent will be obtained. If you are participants, you will also need to obtain their written assent. Templates are available. Forms - Applications, Templates and Amendments section of the Research Ethics.	re involvin ble via the	g chi
6.2 Informed Consent Procedures – please confirm whether written consent in Please tick the relevant checkbox  Yes x  No  Yes, describe the procedures by which written consent will be obtained. If you are participants, you will also need to obtain their written assent. Templates are available forms - Applications, Templates and Amendments section of the Research Ethics	re involvin ble via the website.	g chi
5.2 Informed Consent Procedures – please confirm whether written consent in Please tick the relevant checkbox  Yes x  No  Yes, describe the procedures by which written consent will be obtained. If you are participants, you will also need to obtain their written assent. Templates are available.	re involvin ble via the website.	g chi
5.2 Informed Consent Procedures – please confirm whether written consent in Please tick the relevant checkbox  Yes x  No  Yes, describe the procedures by which written consent will be obtained. If you are participants, you will also need to obtain their written assent. Templates are available forms - Applications, Templates and Amendments section of the Research Ethics	re involvin ble via the website.	g chi

If you are gathering data from an online process such as Google Form or SurveyMonkey then you should use a page such as the one below, to capture participants' informed consent and your data

gathering should not proceed until participants have completed this form with the appropriate answers.

# Participant - please complete the following (by clicking Yes/No for each question)

I have read the Plain Language Statement (or had it read to me) *	I understand I may withdraw from the Research Study at any point *
Yes	O Yes
O No	O No
I understand the information provided *	I have read and understand the arrangements to be made to protect confidentiality of data, including that confidentiality of information provided is
O Yes	subject to legal limitations *
O No	○ Yes
0	○ No
I have had an opportunity to ask questions and discuss this study *	I have read and understand confirmations relating to any other relevant information as indicated in the PLS $^{\bullet}$
○ Yes	○ Yes
○ No	O No
I understand the information provided in relation to data protection *	I consent to participate in this research study *
O Yes	O Yes
O No	O No
I have received satisfactory answers to all my questions *	
○ Yes	
O No	

#### SECTION 6 - SUBMISSION CHECKLIST AND RESEARCHER DECLARATION

# 6.1 Please confirm all required supplementary documentation to be included in this application within Section 7:

Checklist – tick the relevant check box for each item		s	N/A
Participant Information Sheet/s			
Informed Consent Form/s			X
Informed Assent Form/s			
Recruitment Advertisement			
Questionnaire/Survey			
Interview/Focus Group Questions			
Debriefing Material			
Bibliography			
Approval from another Research Ethics Committee			
Evidence of other external approvals (e.g. Board of Management letter)			
Evidence of internal approvals (e.g. BSC approval review letter)			
Other – provide details here:			

#### 6.2 Signed Declaration

By submitting this form, the applicant (and supervisor) agree to the following:

The information contained herein is, to the best of my knowledge and belief, accurate. I have read the University's current research ethics guidelines, and accept responsibility for the conduct of the procedures set out in the attached application in accordance with the form guidelines, the REC guidance and resources, the University's Conflict of Interest Policy, its Code of Good Research Practice and any other condition laid down by the Dublin City University Research Ethics Committee. I have attempted to identify all risks related to the research that may arise in conducting this research and acknowledge my obligations and the rights of the participants.

I also acknowledge my requirement to be informed as to other duties and legal obligations applying to my research, and to comply with these duties and obligations – this includes being informed about DCU Data Protection guidelines for researchers, DCU Child Protection policy and procedures (where relevant) and DCU Insurance requirements.

I and my co-investigators and/or supporting staff have the appropriate qualifications, experience and facilities to conduct the research set out in the attached application and to deal with any emergencies and contingencies related to the research that may arise. Research will not commence until required consents and approvals are in place.

Electronic Signature(s):
Supervisor: PP Stephen Blott
Print Name here: Stephen Blott
Date: 05/03/2024
Student(s) signature(s):Alexandru Dorofte, James Hackett
Print Name(s) here:Alexandru Dorofte, James Hackett
Date: 05/03/2024

Please attach all required documentation as confirmed by you in the previous section. The application should then be saved as one file in <u>PDF format</u> before submission via the project dashboard.

https://docs.google.com/forms/d/1O6N3tIGSDA4zsWUX9DZNiQR\_69Z5Z1-0BZU0Ll3iJgM/edit