

# Dublin City University School of Computing ETHICS COMMITTEE

# NOTIFICATION FORM FOR LOW-RISK PROJECTS AT UNDERGRADUATE OR TAUGHT MASTERS LEVELS

Application Number:			
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Please read the following information carefully before completing your application. Failure to adhere to these guidelines will make your submission ineligible for review.

- Download this form
- > Completed applications must be uploaded to your School of Computing GitLab repo, and must be located in "docs/ethics.pdf".
- > Your supervisor will be notified automatically and must approve your approach initially.
- > The application should consist of one electronic file (PDF) only. The completed application must include this form and also must incorporate all supplementary documentation, especially that being given to the proposed participants e.g consent forms, plain English language statement. It must be proofread and spell-checked before submission.
- ➤ All sections of the application form must be answered as instructed and within the word limits given.

Applications which do not adhere to all of these requirements will not be accepted for review and will require resubmission

Applications must be completed on this form; answers in the form of attachments will not be accepted, except where indicated. No hard copy applications will be accepted. The project <u>must not</u> commence until written approval has been received from the School of Computing Ethics Committee.

PROJECT TITLE	Music Analysis Tool
PRINCIPAL INVESTIGATOR(S)	Dr. Donal Fitzpatrick
The named Principal Investigator is the person with primary responsibility for the research project. In the case of Taught Masters projects and undergraduate projects the supervisor is the Principal Investigator.	
START AND END DATE	02/05/2019
	25/05/2019
LEVEL OF RISK	Notification

Please indicate whether this project requires more
than a notification Justification for your choice is
required under section 3.1

Please confirm that <u>all</u> supplementary information is included in your application (in electronic copy). If questionnaire or interview questions are submitted in draft form, please indicate this by putting (draft) after YES. A copy of the final documentation must be submitted for final approval when available.

My application has been collated as one electronic file which includes the following documentation:			INCLUDED (mark as YES)	NOT APPLICABLE (mark as N/A)
Bibliography				N/A
Recruitment a	dvertisement			N/A
Plain language	e statement/Infor	mation statement	YES	
Informed cons	ent form		YES	
	Data ://www.dcu.ie/site rity_schedule.xls	Security Schedule es/default/files/info/3blank_data_		N/A
Evidence of external approvals related to the research				N/A
Questionnaire/Survey				N/A
Interview/Focus Group Questions				N/A
Debriefing material			N/A	
Other (e.g. loc	al government a	oproval )		N/A

#### Please note:

- 1. Any amendments to the original approved proposal must receive prior SCEC approval.
- 2. As a condition of approval investigators are required to document and report immediately to SCEC any adverse events, any issues which might negatively impact on the conduct of the research and/or any complaint from a participant relating to their participation in the study

#### 1. ADMINISTRATIVE DETAILS

Project Type (select one): Undergraduate Project - Final Year

Undergraduate Project – non-final Year

Taught Masters (Practicum)

(projects at other levels, e.g. PhD or research Masters, should be approved by the University's REC if necessary)

#### 1.1 INVESTIGATOR CONTACT DETAILS

PRINCIPAL INVESTIGATOR(S): Your supervisor and other academic staff who are assisting, it should be clear who is the person who is carrying out the research procedures.

NAME	SCHOOL/UNIT	EMAIL
	School of computing	donal.fitzpatrick@dcu.ie

#### OTHER INVESTIGATORS (STUDENT(S):

NAME	SCHOOL/UNIT	EMAIL
Patrick Ferry	School of computing	patrick.ferry2@mail.dcu.ie

1.2	WILL THE RESEARCH BE UNDERTAKEN ON-SITE AT A Dublin City University CAMPUS ?  YES or NO YES
2.7.)	(If NO, state details of the off-campus location – provide details of the approval to gain access to that location in section
1.3	IS THIS PROTOCOL BEING SUBMITTED TO ANOTHER ETHICS COMMITTEE, OR HAS IT BEEN PREVIOUSLY SUBMITTED TO AN ETHICS COMMITTEE?  YES or NO NO
(If YES,	please provide details and attach copies of approval(s) received etc.)
The in Universet of (https://Code Resea condu	ARATION BY PRINCIPAL INVESTIGATOR(S) Information contained herein is, to the best of my knowledge and belief, accurate. I have read the resity's current research ethics guidelines, and accept responsibility for the conduct of the procedures at in the attached application in accordance with the form guidelines, the SCEC guidelines in the attached application in accordance with the form guidelines, the SCEC guidelines in the attached application in accordance with the form guidelines, the SCEC guidelines of Good Researchsupport/researchethics.shtml), the University's policy on Conflict of Interest, of Good Research Practice and any other condition laid down by the Dublin City University arch Ethics Committee. I have attempted to identify all risks related to the research that may arise in cting this research and acknowledge my obligations and the rights of the participants.
other (	e exists any affiliation or financial interest for researcher(s) in this research or its outcomes or any circumstances which might represent a perceived, potential or actual conflict of interest this should clared in accordance with Dublin City University policy on Conflicts of Interest.
to cor	my co-investigators or supporting staff have the appropriate qualifications, experience and facilities aduct the research set out in the attached application and to deal with any emergencies and gencies related to the research that may arise.
Electro	onic Signature(s):
	al investigator(s): FitzpatrickPatrick Ferry
Print N	ame(s) here: Donal FitzpatrickPatrick Ferry

Date: 02/05/2019

#### 2. PROJECT OUTLINE

2.1	LAY	<b>DESCRIP</b>	TION	Max.	300	words
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Please outline, in terms that any non-expert would understand, what your research project is about, including what participants will be required to do. Please explain any technical terms or discipline-specific phrases.

My project is a webapp that transcribes wav files to midi files by analyses the frequencies. Participants would be asked to upload a file through the webapp and rate the output. The users will use a test account.

#### 2.2 AIMS OF AND JUSTIFICATION FOR THE RESEARCH (Max. 400 words)

State the aims and significance of the project. Where relevant, state the specific hypothesis to be tested. Please provide a brief description of background research, a justification as to why this research project should proceed in that context and an explanation of any expected benefits to the community. NB – all references cited should be listed in an attached bibliography.

The aims would be to test the functionality of the front end, and to evaluate the accuracy of the output.

#### 2.3 DESCRIBE THE METHODOLOGY BEING USED TO ACHIEVE YOUR STATED AIMS

Provide an outline of the proposed method and state who is doing which task – include details of data collection techniques, the tasks participants will be asked to do, the estimated time commitment involved, and how data will be analysed. If the project includes any procedure which is beyond already established and accepted techniques please include a description of it. There should be enough detail provided to facilitate ethical review, but applicants are encouraged to keep it as succinct as possible.

Questionnaire. Observing

#### 2.4 PARTICIPANT PROFILE

Provide the number, age range and source of participants. Please provide a justification of your proposed sample size. Please provide a justification for selecting a specific gender, age, or any other group if this is done in your project.

12 users, no specified age range.

# 2.4(a) PARTICIPANT VULNERABILITY

Are some or all of participants vulnerable in any way? (e.g by virtue of the group they belong to, people who have undergone traumatic or adverse emotional events, people with diminished cognitive ability, power relations between researchers and participants etc.)? If they are, state what this vulnerability (or vulnerabilities) is and justify why this research is being done with such participants.

No			

#### 2.4(b) CHILD PARTICIPANTS (anyone under 18 years old)

If your participants include children, you must confirm that you are in compliance with the research specific guidelines as detailed in "Keeping Children Safe - Policies and Procedures supporting Child Protection at DCU" - available at: <a href="https://www4.dcu.ie/sites/default/files/policy/157%20-%20child\_protection\_handbook\_rev1%282%29%281%29.pdf">https://www4.dcu.ie/sites/default/files/policy/157%20-%20child\_protection\_handbook\_rev1%282%29%281%29.pdf</a>

	Please indicate your compliance with the following guidelines:	Mark here
	We confirm that we have read and agree to act in accordance with the DCU Child	N/a
	Protection policy and procedures	
	We confirm that we have put in place safeguards for the children participating in the	N/a
	research	
	We confirm that we have supports in place for children who may disclose current or	N/a
	historical abuse (whether or not this is the focus of the research)	1
	EXPLAIN HOW PARTICIPANTS ARE TO BE RECRUITED  Please provide specific details as to how you will be recruiting participants. How will people be in:	formed that you
	doing this research? How will they be approached and asked if they are willing to participate? If	rormea that you ' vou are mailin
	phoning people, please explain how you have obtained their names and contact details. If a recruitment	
	to be used, please ensure you attach a copy to this application.	
	I will ask them to participate on the day.	
	PLEASE EXPLAIN WHEN. HOW. WHERE. AND TO WHOM RESULTS WILL BE	DISSEMINATI
	PLEASE EXPLAIN WHEN, HOW, WHERE, AND TO WHOM RESULTS WILL BE INCLUDING WHETHER PARTICIPANTS WILL BE PROVIDED WITH ANY INFORMAT FINDINGS OR OUTCOMES OF THE PROJECT?	DISSEMINATION AS TO T
	INCLUDING WHETHER PARTICIPANTS WILL BE PROVIDED WITH ANY INFORMAT	DISSEMINATI TION AS TO T
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√ES <sub>i</sub>	INCLUDING WHETHER PARTICIPANTS WILL BE PROVIDED WITH ANY INFORMAT FINDINGS OR OUTCOMES OF THE PROJECT?  N/a  ARE OTHER APPROVALS REQUIRED TO GAIN ACCESS TO ANOTHER LOCATION, ETC.?  YES or NO NO  Please specify from whom and attach a copy of the approval documentation. If this is not yet available when this will be obtained.)  HAS A SIMILAR PROPOSAL BEEN PREVIOUSLY APPROVED BY THE DCU SCEC?  YES OR NO	ORGANISATI

# 3. RISK AND RISK MANAGEMENT

3.1	JUSTIFICATION OF STATED LEVEL OF RISK TO RESEARCH PARTICIPANTS
	You must provide a justification for the stated level of risk, as indicated on the cover page of your application. Note that
	the level of risk may be influenced by the vulnerability of the research group, the methods employed and the nature of the
	research itself. For further information on risk levels, please refer to the Levels of Review information on the website:
	https://www.dcu.ie/researchsupport/researchethics.shtml

The survey will be anonymous and I will be only observing.					
The survey will be anonymous and I will be only observing.					

# 3.2 DOES THE RESEARCH INVOLVE:

	YES or NO
<ul><li>use of a questionnaire? (attach copy)?</li></ul>	YES
<ul><li>interviews (attach interview questions)?</li></ul>	NO
<ul> <li>observation of participants without their knowledge?</li> </ul>	NO
<ul> <li>participant observation (provide details in section 2)?</li> </ul>	YES
<ul> <li>audio- or video-taping interviewees or events?</li> </ul>	NO
<ul> <li>access to personal and/or confidential data (including student, patient or client data) without the participant's specific consent?</li> </ul>	NO
<ul> <li>administration of any stimuli, tasks, investigations or procedures which may be experienced by participants as physically or mentally painful, stressful or unpleasant during or after the research process?</li> </ul>	NO
<ul> <li>performance of any acts which might diminish the self-esteem of participants or cause them to experience embarrassment, regret or depression?</li> </ul>	NO
<ul> <li>investigation of participants involved in illegal activities?</li> </ul>	NO
<ul> <li>procedures that involve deception of participants?</li> </ul>	NO
<ul> <li>administration of any substance or agent?</li> </ul>	NO
<ul> <li>use of non-treatment of placebo control conditions?</li> </ul>	NO
<ul> <li>collection of body tissues or fluid samples?</li> </ul>	NO
<ul><li>collection and/or testing of DNA samples?</li></ul>	NO
participation in a clinical trial?	NO
<ul> <li>administration of ionising radiation to participants?</li> </ul>	NO

# 3.3 POTENTIAL RISKS TO PARTICIPANTS AND RISK MANAGEMENT PROCEDURES

Identify, as far as possible, all potential risks to participants (physical, psychological, social, legal, economic, etc.), associated with the proposed research. Please explain what risk management procedures will be put in place to minimise these risks.

None.			

3.4 ARE THERE LIKELY TO BE ANY BENEFITS (DIRECT OR INDIRECT) TO PARTICIPANTS FROM THIS RESEARCH?

YES	NO	
NO		

(If YES, provide details.)			

3.5	ARE THERE ANY SPECIFIC RISKS TO RESEARCHERS?
Example	es include use of dangerous materials, asking certain types of questions, research being undertaken in certain locations, researchers working alone in isolated areas, etc.
	YES or NO
	NO NO
	(If YES, please describe and explain what risk management procedures will be put in place to minimise these risks.)
3.6	DEALING WITH ADVERSE/UNEXPECTED OUTCOMES  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 project.
	autorios onocio lo participante unemiginom inversorionem in the project.
	I will stand by and address any unexpected outcomes accordingly
	T will starte by and address any anoxposited satisfance assortanigly
2.7	HOW WILL THE CONDUCT OF THE PROJECT BE MONITORED?
3.7	explain how the principal investigator will monitor the conduct of the project (especially where several people are involved in
i icasc c	recruiting or interviewing, administering procedures, etc.) to ensure that it conforms with the procedures set out in this
	application. In the case of student projects please give details of how the supervisor(s) will monitor the conduct of the
	project.
	A meeting will be organized to ensure all conforms
3.8	SUPPORT FOR PARTICIPANTS
Depenai	ing on risks to participants you may need to consider having additional support for participants during/after the study.  Consider whether your project would require additional support, e.g., external counselling available to participants.
	Please advise what support will be available.
	Thouse davide what support will be available.
	N/a
	14/4
3.9	DO YOU PROPOSE TO OFFER PAYMENTS OR INCENTIVES TO PARTICIPANTS?
3.5	YES or NO
	NO
	(If YES, please provide further details.)
	in 120, pictor provide further details.

3.10 DO ANY OF THE RESEARCHERS ON THIS PROJECT HAVE A PERSONAL, PHILOSOPHICAL, FINANCIAL OR COMMERCIAL INTEREST IN ITS OUTCOME THAT MIGHT INFLUENCE THE INTEGRITY OF THE RESEARCH, OR BIAS THE CONDUCT OR REPORTING OF THE RESEARCH, OR UNDULY DELAY OR OTHERWISE AFFECT THEIR PUBLICATION?

	YES or NO NO
(If YES,	please specify how this conflict of interest will be addressed.)

<mark>suppor</mark>	e academic qualifications and outline the experience and skills <u>relevant to this project</u> that the PI, other researchers and ar rting staff have in carrying out the research and in dealing with any emergencies, unexpected outcomes, or contingencies th rise. <b>State specifically who will be carrying out the research procedures</b>
An ur	ndergrad student studying computer applications at DCU
5.	CONFIDENTIALITY/ANONYMITY
5.1	WILL THE IDENTITY OF THE PARTICIPANTS BE PROTECTED?  YES or NO YES  (If NO, please explain why.)
IF YO	U ANSWERED YES TO 5.1, PLEASE ANSWER THE FOLLOWING QUESTIONS:
5.2	HOW WILL THE ANONYMITY OF THE PARTICIPANTS BE RESPECTED?
	Please bear in mind that where the sample size is very small, it may be impossible to guarantee anonymity/confidentiality of participant identity. Participants involved in such projects need to be advised of this limitation in the Plain Language Statement/Information Sheet. If you intend to fully anonymize the data, please provide details
	I will prepare a test account to be used so no data is collected on participants and feedback will be anonymously labeled
5.3 Particip	LEGAL LIMITATIONS TO DATA CONFIDENTIALITY pants need to be made aware that confidentiality of information provided cannot always be guaranteed by researchers and
	can only be protected within the limitations of the law - i.e., it is possible for data to be subject to subpoena, freedom of information claim or mandated reporting by some professions. This information should be included in your Plain Languag Statement and Informed Consent Form. Depending on the research proposal and academic discipline, you may need to state additional specific limitations.
	State how and where participants will be informed of these limitations
	Included in plain language statement

4. INVESTIGATORS' QUALIFICATIONS, EXPERIENCE AND SKILLS (Approx. 200 words)

6. REGUI	PERSONAL LATION	DATA -	COMPLIANCE	WITH	THE	GENERAL	DATA	PROTECTION
from the DCU and	data in conjunction d its constituent unit	with other info	ndividual (i.e. the 'Data cormation that is in, or is th teams etc.). Further coo/dp/guides.shtml	s likely to co	ome into,	the possession of	of the 'Data	Controller' (i.e.
6.1	IS PERSONAL YES or NO NO	DATA BEIN	IG PROCESSED AS	S PART O	F THIS	PROJECT?		
	If YES, Please	indicate you	ur compliance with	the follow	ving gu	idelines:		Mark here
	Protection Unit	guidance a	e read and agree nd procedures rega	arding per	sonal d	ata		
			put in place a Per tached it to this app		ta Secu	rity Schedule	(PDSS)	
	Please see th guidance	<mark>e GDPR an</mark>	nd the Research E	thics Pro	ocess s	ection of the	SCEC m	<u>ain webpage</u> fo
			LEASE ANSWER T			QUESTIONS:		
6.2			AL DATA IS BEING include health data, g			lata relating to etl	hnicity/race	of participants, the
rvoto ope	sex lives and/or s			criciro data	arra/or a	ata relating to en	ппонулгаес	or participante, the
6.3	WILL ANONYN YES or NO	IISATION/PS	SEUDONYMISATIO	N OF THE	E PERS	ONAL DATA E	BE UNDER	RTAKEN?
	(If NO, please exp	olain why.)						

# 7. DATA/SAMPLE STORAGE, SECURITY AND DISPOSAL

For the purpose of this section, "Data" includes that in a raw or processed state (e.g. interview audiotape, transcript or analysis). "Samples" include body fluids or tissue samples.

# 7.1 HOW AND WHERE WILL THE DATA/SAMPLES BE STORED?

Note that the SCEC recommends that all data be stored on campus – please justify any off-site storage.

The data will be stored using services such as google forms or survey monkey

## 7.2 WHO WILL HAVE ACCESS TO DATA/SAMPLES?

If people other than the main researchers have access, please name who they are and explain for what purpose.

Investigators only

#### 7.3 HOW LONG IS THE DATA TO BE HELD/RETAINED FOR?

Note that with very few exceptions **personal data** may not be retained indefinitely. It is up to the unit or research team to establish an upper retention limit for each category of personal data under its control.

Until the completion of the project

# 7.4 IF DATA/SAMPLES ARE TO BE DISPOSED OF, PLEASE EXPLAIN <u>HOW, WHEN</u> AND <u>BY WHOM</u> THIS WILL BE DONE?

Note that simply deleting files is not sufficiently secure. The additional steps to be taken to maintain data security should be given.

Personal data must be disposed of in a safe and secure manner at the end of its retention period. If the data is stored in a: a) paper based format then shredding or disposal via a secure bin is recommended; or b) if it is stored in an electronic based format then deletion of the record or full anonymization of the data is recommended. If data/samples are NOT being disposed of, please justify this decision.

The data will be disposed of by Patrick Ferry upon completion of project demo through appropriate methods on Google Forms

8.	FUNDING OF THE RESEARCH
8.1	HOW IS THIS WORK BEING FUNDED, IF IT IS EXTERNALLY FUNDED?  N/A
8.2	PROJECT GRANT NUMBER (If relevant and/or known – otherwise mark as N/A)  N/A
8.3	DOES THE PROJECT REQUIRE APPROVAL BEFORE CONSIDERATION FOR FUNDING BY A GRANTING BODY?  YES or NO NO
8.4.1	HOW WILL PARTICIPANTS BE INFORMED OF THE SOURCE OF THE FUNDING? (e.g. included in the Plain Language Statement)  N/A
8.5	DO THE FUNDERS OF THIS PROJECT HAVE A PERSONAL, FINANCIAL OR COMMERCIAL INTEREST IN ITS OUTCOME THAT MIGHT COMPROMISE THE INDEPENDENCE AND INTEGRITY OF THE RESEARCH, OR BIAS THE CONDUCT OR REPORTING OF THE RESEARCH, OR UNDULY DELAY OR OTHERWISE AFFECT THEIR PUBLICATION?  YES OF NO NO
(If YES,	please specify how this conflict of interest will be addressed.)

# 9. PLAIN LANGUAGE STATEMENT (Attach to this document. Approx. 400 words)

A Plain Language Statement (PLS) should be used in all cases. This is written information in plain language that you will be providing to participants, outlining the nature of their involvement in the project and inviting their participation. The PLS should specifically describe what will be expected of participants, the risks and inconveniences for them, and other information relevant to their involvement. Please note that the language used must reflect the participant age group and corresponding comprehension level – if your participants have different comprehension levels (e.g. both adults and children) then separate forms should be prepared for each group. The PLS can be embedded in an email to which an online survey is attached, or handed/sent to individuals in advance of their consent being sought. See link to sample templates on the website:

https://www.dcu.ie/researchsupport/ethicsapproval.shtml

# PLEASE CONFIRM WHETHER THE FOLLOWING ISSUES HAVE BEEN ADDRESSED IN YOUR PLAIN LANGUAGE STATEMENT/ INFORMATION SHEET FOR PARTICIPANTS:

	YES or NO
Introductory Statement (PI and researcher names, school, title of the research)	YES
What is this research about?	YES
Why is this research being conducted?	YES
What will happen if the person decides to participate in the research study?	YES
How will their privacy be protected?	YES
How will the data be used and subsequently disposed of?	YES
What are the legal limitations to data confidentiality?	YES
What are the benefits of taking part in the research study (if any)?	YES
What are the risks of taking part in the research study?	YES
Confirmation that participants can change their mind at any stage and withdraw from the study	YES
How will participants find out what happens with the project?	YES
Contact details for further information (including SCEC contact details)	YES
Details relating to GDPR Compliance if Personal Data is being sought	YES

	Contact details for further information (including SCEC contact details)	YES
	Details relating to GDPR Compliance if Personal Data is being sought	YES
f any of	these issues are marked NO, please justify their exclusion:	
-		
· <del>-</del>		_
10.	INFORMED CONSENT FORM (Attach to this document. Approx. 300 words)	
10.	INTORNIED CONSENT FORM (Allacir to this document. Approx. 300 words)	
n moot	cases where interviews or focus groups are taking place, an Informed Consent Form is required.	This is an importar
	nt requiring participants to indicate their consent to participate in the study, and give their signature. If y	
	under 18), it is best practice to provide them with an assent form, while their parents/guardians will be	
	Form. In cases where an anonymous questionnaire is being used, it is enough to include a tick box	
	eath the information section for participant), where participants can indicate their consent.	m are questionina
	to sample templates on the website: https://www.dcu.ie/researchsupport/ethicsapproval.shtml	
	· · · · · · · · · · · · · · · · · · ·	
NB - IF	AN INFORMED CONSENT FORM IS NOT BEING USED, THE REASON FOR THIS MUST BE JUSTIF	IED HERE.

# **DUBLIN CITY UNIVERSITY**

### **Informed Consent Form**

Title: Music Analysis Tool Final Year Project University Department: School of Computing Principle Investigator: Dr. Donal Fitzpatrick Investigators Email: donal.fitzpatrick@dcu.ie Principle Investigator Phone Number: 01 700 8929

Other Investigators: Patrick Ferry, patrick.ferry2@mail.dcu.ie, 0833685568

I am aware that all the data gathered will be strictly for research only for the application. There will be no personal data to be stored for this research study and will be disposed when the project is complete.

Please Complete the following (Circle Yes or No for each question)

I have read the Plain Language Statement (or had it read to me)	Yes/No
I understand the information provided	Yes/No
I have had an opportunity to ask questions and discuss this study	Yes/No
I have received satisfactory answers to all my questions	Yes/No

I understand may withdraw from this research study at any time.

I am aware of how the data collected will destroyed and when.

I am aware I will have to complete a questionnaire for this study.

I am aware of how data will be protected until destruction.

I have read and understood the information in this form. My questions and concerns have been answered

•	researchers, ch project.	and I have	a copy	of this	consent	form.	Therefore,	I consent	to take	part	in th
Participant Signature:											
Name	in Block Cap	itals:									

Witness:

Date:

# **DUBLIN CITY UNIVERSITY**

## **Plain Language Statement**

Title: Music Analysis Tool Final Year Project
University Department: School of Computing
Principle Investigator: Dr. Donal Fitzpatrick
Investigators Email: donal.fitzpatrick@dcu.ie
Principle Investigator Phone Number: 01 700 8929

Other Investigators: Patrick Ferry, patrick.ferry2@mail.dcu.ie, 0833685568

This application will take audio files or a youtube link to a song as an input and it will try to transcribe the notes and output to a midi file. Participants will be asked to use the web application front end and listen to midi files which the system will output. They will then be given a questionnaire containing questions about how well the application worked and if any problems arose. They will also be asked to rate the accuracy of the output. No personal data will be stored. Data will be destroyed on completion of the project.

If at any point, during the testing of this project, you feel uncomfortable in participating in this project you may leave. All involvement in this project is entirely voluntary.

There are no risks involved in this study.

If participants have concerns about this study and wish to contact an independent person, please contact:

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