For IRB Office Use
IRB No:
Rec'd:

#### APPLICATION FOR IRB REVIEW OF RESEARCH INVOLVING HUMAN SUBJECTS

(Not for exempt research)

Please complete this application as thoroughly as possible. Your application should include the following:

- 1. A consent form using the current CMU template that the participants and/or parent/guardian will be required to sign.
- 2. A copy of any questionnaires, surveys, images, de-briefings that will be used.
- 3. A copy of any recruitment documents (including advertisements, flyers, letters, invitations, email) to be used;
- 4. A copy of the training certificates for all individuals working on the research unless they are on file with the CMU IRB. Training is available at: <a href="http://www.citiprogram.org">http://www.citiprogram.org</a>. See the <a href="https://www.citiprogram.org">IRB website</a> for details.
- 5. If the PI is a student, the faculty advisor must submit a Faculty Advisor Assurance Form.

Please email all documents to <u>irb-review@andrew.cmu.edu</u>. For assistance call CMU Research Compliance @ 412-268-5460 or email <u>irb-review@andrew.cmu.edu</u>. Additional information and templates are available at <a href="http://www.cmu.edu/osp/regulatory-compliance/human-subjects.html">http://www.cmu.edu/osp/regulatory-compliance/human-subjects.html</a>

1. Protocol			
Title: Pedestrian Circulation Through	out the Carnegie Me	llon Campus	
This is a previously approved stud	y that has lapsed.	Previous IRB No: H	IS
2. Principal Investigator (PI)			
Name: Erika Tang		Department: Statis	stics
Telephone: 949-705-8138	E-mail: ertang@an	drew.cmu.edu	Training Cert. Attached On File
I am a student. If so, please provi	de information abou	ut your faculty advisor b	pelow.
	E-mail: bj20(	@andrew.cmu.edu	Training Cert. Attached On File
Faculty Advisor Name: Brian Junker			
If a student is the PI, the	faculty advisor must c	omplete and submit a Fac	culty Advisor Assurance Form.
If there is someone other than PI to o	orrespond with rega	arding this protocol, ple	ease list below.
Contact Person Name:	Te	lephone:	E-mail:
Business Manager for your departme	nt:	E-mail:	
3. Co-investigators			
Name: Zhiyi Tang	E-mail: zhiyit	:@andrew.cmu.edu	Training Cert.   Attached   On File
Name: Jason Sun	E-mail: jewo	os@andrew.cmu.edu	Training Cert. Attached On File
Name: David Zimmerman	E-mail: dbz@	andrew.cmu.edu	Training Cert.   Attached   On File
Name:	E-mail:		Training Cert. Attached On File
Name:	E-mail:		Training Cert.
Name:	E-mail:		Training Cert. Attached On File
Name:	E-mail:		Training Cert. Attached On File
4. Funding			
Unfunded research		Sponsor/Source: N	I/A

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External Funding	
	CDEV Droposal #4 N / A
	SPEX Proposal #: N/A
│	Oracle String:N/A
Grant Title: N/A	Oracle String.N/A
<u> </u>	ase get it from your department's business manager.
5. Protocol Description	use get it from your department's business manager.
Provide, in lay terms, a summary of your proposed study as outline	ed below. You may attach the protocol to this form if you like.
Purpose of the study. See Attached	
Describe the research procedures (include the activity, locat	ion and time required of the participant) See Attached
Describe the research procedures (include the activity, locat	ion and time required of the participanty. See Attached
Who will be asked to participate?See Attached	
Will questionnaires or surveys be used? X Yes No	
Will tasks be done on a computer? Yes No If yes, ho	w will the tasks be accessed? Remotely via the internet?
In the research lab? Other, please explain:	
Will deception be used? Yes No If yes, describe how	participants will be debriefed. Please include the de-
	participants will be desirered. Flease include the de
briefing material and/or script.	
Will the research be conducted on the CMU campus? X	No If no, please indicate the location(s).
If applicable, please attach documentation of permis	ssion to conduct research in private, non-CMU space.
6. Participants	
Will any of the following classes of vulnerable subjects be in	volved in the proposed study? (check all that apply)
Class	Comments
Pregnant women, human fetuses Yes No Pregnar	nt l
women will not be specifically included or excluded. (See	
http://www.hhs.gov/ohrp/humansubjects/guidance/45cfr46.htm, research that is	
incidental to pregnancy and has no risk to the fetus can only include pregnant wom	en if ALL
aspects of Subpart B are met.)	
Neonates 🔛 Yes 🔀 No	

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Prisoners 🗌 Yes 🔀 No	
Children Yes No	
Individuals with compromised mental status   Yes   No	If yes, indicate how this will be determined.
Will the participants be capable of understanding the nature	of the study and the consent process? X Yes No
If no, explain.	
What is the age range of participants in the proposed study?	7 17-25 years old
How many participants are needed for the study? 2000	How was that number determined?This should represent
	a reasonable number for the traffic flow of students on
	campus who would pass by the described locations in
	the protocol. This amount of data will allow us to
	estimate the travel patterns of the student body of
	Carnegie Mellon on campus.

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What do you estimate the ratio of males to	females be? 6:4 Will this be re	eflective of the local population? 🔀 Yes 🗌
No Will you target a certain population?	Yes No Please explain See	Attached
What do you estimate the percentage of m	inorities will be? 59%	
Please list inclusion and exclusion criteria. S	See Attached	
7. Participant Recruitment		
Describe how participant recruitment will b	pe performed. Include how and	by whom potential participants are
introduced to the study. See Attached		
Check all boxes below that apply.		
CMU directory	Postings, Flyers	Radio, TV
E-mail solicitation Indicate how the em	nail addresses are obtained:	
☐ Web-based solicitation. Specify sites:		
Participant Pool. Specify what pool:Car	ngie Mellon Students walking o	on campus between the hours of 7:30 am to
9:00 pm, Monday through Friday.		

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Other, please specify:
Please attach any recruiting materials you plan to use and the text of e-mail or web-based solicitations you will use.
8. Consent
Do you plan to use consent forms?
If no, you must complete the section below on waiver of informed consent.
If yes, describe how consent will be obtained and by whom.
If participants are minors will assent forms be used?   Yes   No If No, please explain. The study does not intend to
survey minors and will therefore not involve this aspect of consent. If one is selected for the study we will inform
them they unfortunately cannot participate and select another student.
Will the consent form be presented on paper or online? Paper Online
Are you requesting to use a consent format that is different from the CMU model consent? Yes No
If yes, please explain.
Are you requesting a waiver of informed consent?  Yes No
If yes, please explain how each of the elements listed apply to your study:
1. The research involves no more than minimal risk to the subjects;
2. The waiver will not adversely affect the rights and welfare of the subjects;
3. The research could not practicably be carried out without the waiver and ;
4. Whenever appropriate, the subjects will be provided with additional pertinent information after participation.
Are you requesting a waiver of written documentation (signed) of informed consent? X Yes No
If yes, please answer the following questions.
<ol> <li>Will the only record linking the participant and the research be the consent document and the principal risk to the</li> </ol>
participant harm would be from breach of confidentiality? \( \sum \) Yes \( \sum \) No
<ol> <li>Do you consider this a minimal risk study that involves no procedures for which written consent is normally required</li> </ol>

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outside of research? 🔀 Yes 🔛 No
9. Risks and Benefits
Will participants receive intangible benefit from the study? ☐ Yes ☒ No
Discuss the direct and indirect benefits to participants. There are not direct benefits for the individuals. One indirect
benefit is the availability of information about where students tend to walk on campus so that if a person desires,
they can place themselves in ideal situations where they advertise for something.
Discuss the risks to participants. The participants will be asked to give information about their major and other
similar information. There will be no physical or psychological risks to participants.
Discuss how any risks will be managed and/or minimized. The information that the individuals give up will be kept
confidential since all members of the team have taken the CITI course and will abide by all ethical standards.
If deception is involved, please explain.
Indicate the degree of physical or psychological risk you believe the research poses to human subjects (check which one applies).

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proposed research are not greater, in and of themselves, than those ordinarily encountered in daily life of during the performance o routine physical or psychological examinations or tests.  Greater than Minimal Risk: A risk is greater than minimal where the probability and magnitude of harm or discomfort anticipated in the proposed research are greater than those ordinarily encountered in daily life or during the performance of routine physical or psychological examinations or tests.				
Describe how the study fits in this risk level. Students will be asked simply about some general academic information				
that will be kept confidential by the group members.				
<b>10.</b> Participant Compensation and Cost Are participants to be compensated for		No If you what is	the amount, type and source of funds?	
Are participants to be compensated for	tile study: 🔲 Tes 🔼	ino ir yes, what is	the amount, type and source of funds:	
Amount:	Source:		Type (gift card, cash):	
Will participants who are students be of	fered class credit?	] Yes ⊠ No		
Are other inducements planned to recruit participants?   Yes   No If yes, please describe.				
Are there any costs to participants?   Yes   No If yes, please explain.				
Will you compensate participants for inj	ury resulting from pa	rticipation?	No NA If yes, please describe.	
11. Confidentiality and Data Security				
Will personal identifiers be collected? Yes No Will identifiers be translated to a code? Yes No				

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Will recordings be made (audio, video)? ☐ Yes ☒ No If yes, please describe.
Is the information so sensitive that you will obtain a certificate of confidentiality from NIH?   Yes   No
Who will have access to data (surveys, questionnaires, recordings, interview records, etc.)? The principle investigator
and all of the co-investigators. Professor Junker will also have access to the data.
Describe how you will protect participant confidentiality and secure research records (Will they be stored on a secure
computer, locked cabinet, etc?). The data will be stored on secure computers and never open to public viewing by
anyone aside from the principle investigator and the co-investigators.
Describe your process for monitoring data to ensure that study goals are met. (Review of lab notebooks, meetings to
review data, etc.) We will make checks on the data as a group of at least two members to go over responses and insure that the data is considered reasonable and there are no answers which have been incorrectly coded into the data files.
12. Conflict of Interest
Do you or any individual who is associated with or responsible for the design, the conduct of or the reporting of this research have an economic or financial interest in, or act as an officer or director for any outside entity whose interests could reasonably appear to be affected by this research project: Yes No If yes, please provide detailed information to permit the IRB to determine if such involvement should be disclosed to potential research subjects.
13. Cooperating Institutions
Is this research being done in cooperation with any institutions, individuals or organizations not affiliated with CMU?

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☐ Yes ☑ No If yes, please list and describe their role.
Have you received IRB approval from another IRB for this study?   Yes   No  Pending
If yes, please attach a copy of the IRB approval.
If applicable, please provide the name(s) and address(es) of all officials authorizing to access human subjects in
cooperating institutions not affiliated with CMU.
Please attach documentation of approval.

#### Principal Investigator's Assurance Statement for Using Human Subjects in Research

I certify that the information provided in this IRB application is complete and accurate.

I understand that as Principal Investigator, I have ultimate responsibility for the conduct of IRB approved studies, the ethical performance of protocols, the protection of the rights and welfare of human participants, and strict adherence to the studies protocol and any stipulations imposed by Carnegie Mellon University Institutional Review Board.

I understand that it is my responsibility to ensure that the human participants' involvement as described in the funding proposal(s) is consistent in principle, to that contained in the IRB application. I will submit modifications and/or changes to the IRB as necessary.

I agree to comply with all Carnegie Mellon University policies and procedures, as well as with all applicable federal, state, and local laws, regarding the protection of human participants in research, including, but not limited to:

- Ensuring all investigators and key study personnel have completed human subjects training program;
- Ensuring protocols are conducted by qualified personnel following the approved IRB application;
- Implementing no changes in approved IRB applications or informed consent documents without prior IRB approval in accordance with CMU IRB policy (except in an emergency, if necessary to safeguard the well-being of a human participant, and will report to the IRB within 1 day of such change);
- Obtaining the legally effective informed consent from human participants or their representative, using only the currently approved date-stamped informed consent documents, and providing a copy to the participant.
- Ensuring that only IRB-approved investigators for this study obtain informed consent from potential subjects.
- Informing participants of any relevant new information regarding their participation in the research that becomes available.
- Promptly reporting to the IRB any new information involving risks to research participants, including reporting to the IRB, Data Safety and Monitoring Boards, sponsors and appropriate federal agencies any adverse experiences and all unanticipated problems involving risks to human subjects or others that occur in the course of the research.
- If unavailable to conduct research personally, as when on sabbatical leave or vacation, arrangements for another investigator to assume direct responsibility for studies will be made through modification requests to the IRB;
- Promptly providing the IRB with any information requested relative to protocols;
- Promptly and completely complying with IRB decisions to suspend or withdraw approval for projects;

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- Obtaining Continuing Review approval prior to the date the approval for a study expires (approval for the study will automatically expire);
- Maintaining accurate and complete research records, including, but not limited to, all informed consent documents for 3 years from the date of study completion;
- Informing the CMU IRB of all locations in which human participants will be recruited for protocols and being responsible for obtaining and maintaining current IRB approvals/letters of cooperation when applicable;
- Complying with federal, state and local laws and regulations and sponsor terms and conditions; and
- Complying with CMU policies on the responsible conduct of research.

36-303 Group H		
<u>Erika Tang</u>		
Zhiyi Tang		
<u>Jason Sun</u>		
David Zimmerman		2/08/2011
Principal Investigator Name and Signature	Date	

Note: If e-mailed from the PI's CMU e-mail account a hand written signature is not needed. Please type in name and date. If the PI is a student, the faculty advisor must submit a Faculty Advisor Assurance Form.

#### Please email all documents to <a href="mailto:irb-review@andrew.cmu.edu">irb-review@andrew.cmu.edu</a>.

Note: Links to the policies and Federal regulations for the protection of human research subjects (including the Code of Federal Regulations [.CF.R.] Title 45 CFR Part 46 and Title 21 C.F.R. parts 50 and 56) are available on the IRB web page (http://www.cmu.edu/provost/spon-res/compliance/hs.htm).

Comments:			