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Updated By: Amy R Price @ 10-Oct-2017 05:04:20 PM

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A. Instructions

Thank you for using eRIA for your <u>IRB</u> application. Please read all of the following instructions and complete each step carefully.

- 1. Complete all required questions in this form. A detailed instruction guide is available here. It is recommended that you save your work often (using the Save button above) as this system will not automatically retain information entered.
- Attach all related documents such as advertisements, assent forms/scripts, surveys, consent forms/scripts, grants, interview guides, questionnaires and evidence of training in the Supporting Documents section of this form.
- 3. Check the Locked checkbox to validate all mandatory questions have been answered. Clicking Locked saves the electronic form and verifies all mandatory responses are present. If a message appears indicating mandatory questions have not been completed, then provide responses as indicated. Repeat this step until the "Form Complete Successful" message appears.
- 4. Click the Close button above to close this form and return to the submission package window.
- 5. Upon return to the submission package window, click the Submit button (located on the right, below the application status). Wait for the system to complete the operation. Once your application is submitted the status will change to either Intake Review or Pending PI Review and Assurance.
- 6. Click the Done button to close the main window and return to the Portal.

Icon Legend:

* Required field

Add an entry
Delete an entry
Sedit or choose a
value

Phover over this icon for Help

□Upload a file
□Remove a value or file
□View an uploaded file

B. General Information

Protocol Number 10087

Submission Number 10087-01

*(1) Study Title

Using Amazon Mechanical Turk to create and norm stimulus materials

*(2) Is your research being funded by any of the U.S. Department of Health and Human Services (DHHS) agencies listed here?

Yes

*(3) Select your study sponsor from the list below.

First click the $^{f d}$ icon to the right. Then click the $^{f N}$ icon to select your funder. If you have more than one funder, click the

 \clubsuit icon to add a second funder, then click the riangle icon to select the second funder. Repeat as necessary.

If your study is internally funded by Princeton University or is unfunded, please choose "Princeton University" or "Internal" (either choice is fine).



*(4) Type of Research

Use the icon below to specify if this is a Junior Project, Senior Thesis, Faculty Research, etc.

Postdoctoral Research

C. Research Personnel

(Add)

Name

Hasenfratz, Liat

☐ Principal Investigator (PI)

The PI is the person who is ultimately responsible for the conduct of the study. For student-initiated research, the PI is typically the student's faculty advisor. However, the PI does not have to be your faculty advisor. For the eligibility criteria to serve as a PI at Princeton University, please see this document.

ヹEmail Communication

The IRB will send all communications to the PI. If you would like this person to also receive IRB communications about this study, please check this box.

Role in Research: Researcher

IRB Compliance Training

This information is automatically populated based on data received from the <u>CITI Program</u> or recorded by the IRB Office

Certification	Begin	End
CITI - IRB - Social & Behavioral Research Investigators	22-Feb-2017	

 $\overline{}$

Name

Hasson, Uri

☑ Principal Investigator (PI)

The PI is the person who is ultimately responsible for the conduct of the study. For student-initiated research, the PI is typically the student's faculty advisor. However, the PI does not have to be your faculty advisor. For the eligibility criteria to serve as a PI at Princeton University, please see this document.

Role in Research: Principal Investigator

IRB Compliance Training

This information is automatically populated based on data received from the <u>CITI Program</u> or recorded by the IRB Office

Certification	Begin	End
NIH - NIH IRB Training	03-Oct-2008	

 $\overline{}$

Name

Price, Amy R

☐ Principal Investigator (PI)

The PI is the person who is ultimately responsible for the conduct of the study. For student-initiated research, the PI is typically the student's faculty advisor. However, the PI does not have to be your faculty advisor. For the eligibility criteria to serve as a PI at Princeton University, please see this document.

■Email Communication

The IRB will send all communications to the PI. If you would like this person to also receive IRB communications about this study, please check this box.

Role in Research: Researcher

IRB Compliance Training

This information is automatically populated based on data received from the $\underline{\textit{CITI Program}}$ or recorded by the IRB Office

Certification	Begin	End
CITI - IRB - Social & Behavioral Research Investigators	08-Dec-2016	

n-Princeton Personnel list all individuals from non-Princeton institutions who will be working with a chapter of the project (i.e., investigators from Penn, Columbia, Harvard, etc.).	you on this
e save your work before proceeding to the next section of this form. Other t be saved.	

E. Study Design

The IRB asks the following questions because it must confirm that the study meets certain criteria for approval found in the federal regulations. If you are interested in learning more about these criteria, please visit our resources page.

* Objectives

(1) Using language understandable to a non-scientific layperson, provide a summary of the purpose of your research project. Include a description of the background and rationale for the study, explain the research design, research methodology, hypotheses and goal(s). The IRB understands that sometimes hypotheses and rationale emerge as data is collected, but please do your best to specify the following:

- · List the problems to be addressed
- Describe what is to be learned
- Identify the specific objectives of the proposed research

Research on the neurobiology of language typically investigates the neural representations of isolated individual words or individual sentences, however in real life we interpret the meaning of words using the context of many prior words. Our objective is to obtain norming data for our text stimuli that will better characterize the stimuli to be used in future experiments. This protocol only aims to collect norming data on the stimuli, rather than collect any actual neural data using the stimuli. The aim of this norming data is to provide us with information on the predictability of certain words when given a set of prior words. This will provide an empirical validation for our stimulus materials, rather than relying solely on our intuitions about what should come next. For example, if subjects read the words "She sat at her desk and started writing with her ______", most people would think that the word that should come next in the story is "pencil" or "pen". We will ask participants to respond with the word they believe comes next in the sentence by typing their answers into a blank box. Based on people's responses, we will be able to empirically characterize each of the words in the story.

Research Components

The research involves the following components (check all that apply):

*(2) Questionnaires/Surveys/Interview Guide/Topics of Conversation

Yes **I**No□

Upload the Questionnaires/Surveys/Interview Guide/Topics of Conversation Guide in section N ("Supporting Documents") of this form. The IRB cannot make a risk determination of the study without reviewing the proposed questions.

st(3) Ethnographic/Field Research/Participant Observation/Embedding in the field

Yes 🗖 No🗹

*(11) Analysis of tissues or specimens

Yes 🗆 No🗹

*(12) Audio/Video Recording

Yes 🗆 No🗹

*(13) Data Analysis Only (the research is solely limited to data analysis- no interaction or intervention with the subjects)

Yes No

*(14) Clinical Trials

Yes □ No**ਤ**

*(15) Hazardous Agents

Yes □ No**ਤ**

* Study Procedures

(19) Describe in detail the procedures that will be used to achieve the objectives of the research project. Remember to attach the <u>survey</u>, questionnaire, interview guide, and topics of conversation in the Supporting Documents section, if applicable.

We will collect this norming data using Amazon Mechanical Turk. We will not receive any personal or identifiable information about subjects from Mechanical Turk. We will receive basic demographics about each participant: age, native language, gender, racial and ethnic categories. Amazon Mechanical Turk does not provide any other additional information about the participant. Subjects will be reimbursed for their time at a rate of \$6 per survey. In the survey, they simply read 10 words at a time and then type the word that they believe comes next based on those ten words. We will have approximately 400 participants fill out the survey.

* Number of Subjects

(20) Indicate the maximum number of subjects to be accrued across all sites. If a total number cannot be provided, please provide an estimate. If the activity is solely limited to data analysis, indicate the number of subjects' records that will be analyzed.

400

Study Populations

The regulations require that additional protections be given to the subject populations listed below. If you indicate "Yes" to any of these questions, the application will guide you to resources you should consult.

*(21) Does your study involve children?

No

*(22) Does your study involve <u>prisoners</u>?

No

*(23) Does your study involve pregnant women, human fetuses or neonates?

No

* Inclusion and Exclusion Criteria

(24) Describe the criteria that define who will be included in your study and who will be excluded from your study.

The only requirements will be that the subjects are age 18-65, and that they are native English speakers since the survey is examining the statistics of natural language. Participants are drawn from the diverse Amazon Mechanical Turk community, which represents all minorities and both genders.

Multi-Site Research

*(25) Is this is a multi-site study?

☐ Yes **☑**No

Recruitment Methods

*Describe how you will locate, contact, and invite subjects to participate in your study. If you will be using recruitment materials such as advertisements or invitations (printed, audio, visual, online), submit them in the **Supporting Documents** section of this form.

We will be using Amazon Mechanical Turk to collect these surveys. Our survey will be located on a list of many possible surveys that subjects can choose to complete. We will not perform any additional recruiting such as advertisements or invitations.

*(29) Describe the amount and timing of any payments to subjects.

Subjects will take between 25-45 minutes to complete the survey and they will receive \$6 for the completing the survey. These payment rates reflect standard researcher practices for MTurk studies.

Resources

*(30) Describe your process to ensure that all research personnel are adequately informed about the protocol, the research procedures, and about their duties and functions.

The research personnel involved have reviewed and agreed to the protocol we are proposing. They will receive all future updates to the protocol.

*(31) Describe the availability of medical, psychological, or other resources that subjects may need as a result of participating in your study and explain how these resources will be supplied.

This is a minimal risk study. Participants are not eligible for psychological or medical resources.

F. Risks to Subjects

*(1) List the reasonably foreseeable <u>risks</u>, discomforts, hazards, or inconveniences to the subjects related to the subjects' participation in the research.

We do not foresee many risks, discomforts, hazards, or inconveniences to the subjects related to the subjects' participation in the research. Amazon Mechanical Turk is a webservice where subjects complete the surveys at their leisure on their own computer, and no identifiable information is given to us regarding the subjects. Subjects are typing responses to the words using their keyboard. If they find reading unpleasant or typing unpleasant, then they may find that inconvenient and they can choose to stop the survey at any time. Amazon Mechanical Turk is used by thousands of researchers, and most surveys involve reading or typing so subjects who are choosing to complete surveys on Mechanical Turk are most likely subjects who will not find reading or writing as a inconvenience or a discomfort.

*(2) Describe the probability, magnitude, duration, and reversibility of the risks. Consider physical, psychological, social, legal, and economic risks.

We are not aware of any risks.

*(3) Describe any financial costs (for example, medical tests, tolls, parking) that subjects may be responsible for because of participation in the research.

There are no financial costs that subjects will be responsible for because of participation in the research

*(4) Please describe the medical treatment and <u>compensation</u> that will be provided to subjects in the event of an injury caused by your research. If medical treatment and <u>compensation</u> will not be provided to subjects, indicate that.

Subjects are reading words and typing word responses using the keyboard. We are not aware of any medical treatment or compensation that will be necessary for the subjects, and therefore we will provide no medical treatment or compensation to the subjects.

*(5) Do you think the study poses more than Minimal Risk to subjects?

☐ Yes 🗹 No

G. Potential Benefits

IRB must determine that the risks to subjects are reasonable in relation to the expected benefits. Describe the direct benefits to individual subjects, as well as to society collectively, including contributions to the collective knowledge base of a scientific discipline.

*(1) Describe the potential <u>benefits</u>, if any, that individual subjects may experience from taking part in the research. Include the probability, magnitude, and duration of the potential benefits.

We are not aware of any potential benefits that individual subjects may experience.

*(2) What is the value of this study to society?

The value of this study to society is that we will learn more about the cognitive processes of language and how they are represented in the human brain. Language is one of the most fundamental human skills and we know little about how the human brain processes individual words or groups of words. The results from this survey will allow us to perform analyses to discover how groups of words are represented in the human brain.

H. Withdrawal of Subjects

 f^* (1) Describe anticipated circumstances under which subjects may be withdrawn from the research without their consent. This survey is simple and straightforward to complete. We do not anticipate many situations in which subjects are withdrawn from the research without their consent. It may be that some subjects who complete the survey have an unidentified language disorder in which case their responses may deviate from the group average. If their average responses across the entire survey deviate by more than 4 standard deviations from the mean, then their responses may be eliminated from the group results. Please save your work before proceeding to the next section of this form. Otherwise, your work will not be saved.

I. Confidentiality

st (1) Is your study federally funded and will you collect identifiable, sensitive information? Examples of "sensitive" information include research on mental health and research on the use and effect of alcohol and other psychoactive drugs.

☐ Yes 🗹 No

Please refer to the following resources:

- Research Data Security Guidelines
 Protecting Your Research Data -- Summary Table
- Research Data Management Guidelines

Given the above guidelines, describe the procedures to secure and maintain the confidentiality of data in the following areas:

*(2) Desktop and laptop computers used in the study (see row 3 of the <u>table</u>):

Subjects complete the surveys using Amazon Mechanical Turk on their own computers. No identifiable, sensitive information is given to us by Amazon Mechanical Turk. Our lab computers are password protected.

*(3) File server(s) used in the study (see row 4 of the <u>table</u>):

The file servers to be used in this study require authentication for access to data; file server is configured in a manner consistent with University security practices.

*(4) Removable media used in the study (see row 5 of the <u>table</u>):

*(5) Paper forms used in the study (see row 6 of the <u>table</u>):

*(6) Cloud storage services used in the study (see row 7 of the <u>table</u>):

*(7) Encryption methods used in the study (see row 8 of the <u>table</u>):

*(8) Data transfer methods used in the study (see row 9 of the <u>table</u>):

(1) Describe the steps that will be taken to protect subjects' privacy interests. his data is very private. The subjects complete the surveys online in the privacy of their own ome. The data that we receive from Amazon Mechanical Turk has no identifiable or personal information associated with it.		
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lease save your work before proceeding to the next section of this form. Otherwise, your work dill not be saved.	information associated with it.	
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ill not be saved.	Please save your work before proceeding to the next section of this form. Other	erwise, vour work
	will not be saved.	iwise, your work

. Consent	
(1) Will subjects consent?	
▼ Yes □ No	
The IRB recommends that you use the Princeton University IRB consent form <u>template</u> . Tonsent form template contains approvable language and its use will minimize delays in review process.	
k (2) Will any information about the study be withheld during the consent process (a.k.a. deception)?	•,
☐ Yes ☑No	
(3) Will subjects sign the consent form?	
☐Yes ☑No	
You must request a waiver of written documentation of consent by providing sufficient nformation such that the IRB can justify that the following criteria are met:	
• The research involves no procedures for which written consent is normally required out	side of
the research context, or The only record linking the subject and the research would be the consent document; principal risk of a signed consent document would be the potential harm resulting from breach of confidentiality; and each subject will be asked whether the subject wants documentation linking the subject with the research, and the subject's wishes will gover	n a
We are respectfully requesting that the IRB consider a waiver of written documentation of at this time. We are only collecting indirect identifiers from the Amazon Mechanical Turk participants such as gender, native language, age, racial and ethnic categories. We are in are collecting any information that would allow us to identify Amazon Mechanical Turk part a future date. We also have no interest in being able to identify participants at a later and are only using the Amazon Mechanical Turk to create and norm stimulus materials for contential use in future experiments. Therefore, the only record linking the subject and the research would be the consent document, and the principal risk of a signed consent document would be the potential harm resulting from a breach of confidentiality. Each subject will be whether the subject wants documentation linking the subject with the research, and the subject will govern. Subjects will be given all of the contact information for us (the research well as the Princeton IRB contact information, in case they have any questions or concernation. The research involves no procedures for which written consent is normally required of the research context.	n no way rticipants date or ne ument ne asked ubject's chers) as
(4) Are you obtaining <u>consent</u> of non-English-speaking subjects?	
Yes No	
(9) Subjects are adult students or employees.	
☐Yes ☑ No	
^k (11) Subjects are adults who are <u>cognitively impaired</u> . □ Yes ☑No	
^k (13) Subjects are adults who are unable to <u>consent</u> . ☐ Yes ☑ No	

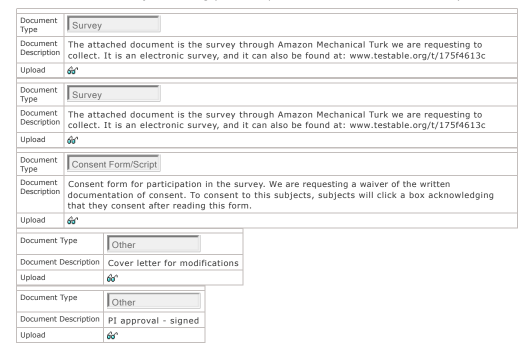
will not be saved.

L. International Research
*(1) Will this project be conducted in whole, or in part, at a location outside the United States? Yes No
Please save your work before proceeding to the next section of this form. Otherwise, your work will not be saved.

M. Conflict of Interest	
*(1) Does the PI or any research personnel have a financial or other potential conflict of interest affecting objectivity in the study? For more information, please review the COI website for details. Yes \square No \square	
Please save your work before proceeding to the next section of this form. Otherwise, your work will not be saved.	

N. Supporting Documents

Attach supporting documents here. Supporting documents include advertisements, assent forms/scripts, surveys, consent forms/scripts, grants, interview guides, questionnaires and verification of human subjects training (if not completed via CITI and while at Princeton).



O. PI Assurance

By electronically approving this protocol and submitting it to the IRB, the PI agrees to follow the following obligations outlined in Princeton University SOP 207: Obligations of the Principal Investigator for Human Subjects Research

- 1. If the PI has any questions about whether s/he is conducting research involving human subjects, the PI must contact the IRB before commencing the study.
- 2. Principal investigators must not commence research until the PI has the IRB approval letter and obtained all other required approvals, such as approvals of departments or divisions that require approval of the use of their resources.
- PIs must conduct the research in accordance with the most recent protocol approved by the IRB.
- 4. PI must protect the rights, safety, and welfare of subjects involved in the research.
- 5. The PI must comply with all requirements and determinations of the IRB.
- 6. The PI must use sound study design in accordance with the standards of his/her discipline and design studies in a manner that minimizes risks to subjects.
- 7. The PI must ensure that there are adequate resources to carry out the research safely. This includes, but is not limited to, sufficient investigator time, appropriately qualified research team members, equipment, and space.
- 8. PIs are required to provide verification of human subjects training. This requirement is described in greater detail in SOP 202.
- 9. The PI must ensure that research staff are qualified including, but not limited to, appropriate human subjects training, education, expertise, credentials, protocol requirements and privileges, to perform procedures and duties assigned to them during the study.
- 10. PIs must not make modifications to the research without prior IRB review and approval, unless necessary to eliminate apparent immediate hazards to subjects.
- 11. The PI must report the information items listed in <u>SOP 206: Reporting Requirements for Investigators</u> to the IRB within 5 business days of learning of the information.
- 12. PIs must submit continuing reviews to avoid a lapse in approval of their study. If approval of the research expires, the PI must stop all research activities immediately and contact the IRB.
- 13. PIs must close the research (end the IRB's oversight) when all the following criteria are met:
 - The protocol is permanently closed to enrollment and
 - o All subjects have completed all protocol-related interventions and interactions and
 - No additional identifiable private information about the subjects is being obtained and
 - The PI's analysis of private identifiable information is completed.
- 14. Unless the IRB approved a protocol to include the following populations, the PI must not enroll the following subjects in the study:
 - Adults unable to consent
 - o Children
 - Neonates of uncertain viability: a newborn that may not be viable (able to survive after delivery, given the benefit of available medical therapy to the point of independently maintaining heartbeat and respiration)
 - Nonviable neonates: a newborn that is not viable
 - o Pregnant women
 - o Prisoners
 - Individuals unable to speak English
- 15. When consent, permission, or assent are required by the IRB, the PI must ensure that it is obtained and
- 16. The PI must retain research records (including signed consent documents) for three years after completion of the research. "Completion of the research" means when the definition of human subjects research is no longer met: the protocol is permanently closed to enrollment; and all subjects have completed all protocol-related interventions and interactions; and no additional identifiable private information about the subjects is being obtained; and analysis of private identifiable information is completed. Completion of the research is typically evidenced by the PI submitting a closure form or the study's lapse (expiration) of approval.
- 17. If the PI is a lead investigator of a multi-site study, the PI must manage information that is relevant to the protection of subjects across all study sites, such as reporting Unanticipated Problems Involving Risks to Subjects or Others to the IRB(s); reviewing interim study results; and securing approval of modifications from the IRB(s) before their implementation.
- 18. For studies regulated by a federal department or agency, the PI must follow the additional obligations of that federal department. These agencies include, but are not limited to, the Department of Defense; Department of Energy; Department of Justice; Environmental Protection Agency; Education Department; Federal Drug Administration; and the National

Institutes of Health.

f * Select the appropriate response below.

I am not the PI

You are welcome to save this document and return later to complete and submit it. If you're ready to submit it, you have two submission options:

- a. If the PI will review documents in this electronic system, you can advance this submission to the PI for his/her review or
- b. If the PI will not review documents in this electronic system, you can submit this to the IRB Office. If you choose this option, you'll have to include an attachment of a hardcopy of this form signed by the PI.
- ${}^{\displaystyle igstyle *}$ Given the options above, select the desired response below.

Attach the signed PI approval and advance this submission to the IRB

*After completing the steps below, please upload a scan or picture of the signed PI Approval Form here.

66

Please follow these steps to complete and then submit your application to the IRB.

- 1. Click the link P. PI Approval Form Initial Application.
- 2. Print the form.
- 3. Have your PI sign the printed form on the signature line.
- 4. Attach/upload the scan or image file to this application (using the Upload icon above).
- 5. Check the Locked checkbox to validate all mandatory questions have been answered. Clicking Locked saves the submission form and verifies all mandatory responses are present. If a message appears indicating "Incomplete mandatory field(s) found", please provide responses as indicated. Repeat this step until you have answered all the mandatory questions. Checking the Locked checkbox when all mandatory questions are answered closes the form automatically and returns you to the submission package window.
- 6. Upon return to the submission package window, click the Submit button (located on the right, below the application status). Wait for the system to complete the operation. Once your application is submitted the status will change to Submit to IRB.
- 7. Click the Done button to close the submission package window and return to the Portal.

P. PI Approval Form - Initial Application	
Protocol Number:10087	
Submission Number:10087-01	
PI:Hasson, Uri	
Study Title:	
Using Amazon Mechanical Turk to create and norm stimulus materials	
Date: 28-Sep-2017	
By signing this form and submitting it to the IRB, I am agreeing to follow the obligations outlined in Princeton University SOP 207: Obligations of the Principal Investigator for Human Subjects Research.	I
PI Signature :	
Date:	



IRB Office- Hide Form: ☐ Yes ☐ No

Research Integrity & Assurance 87 Prospect Ave, 3rd Floor Princeton, NJ 08544 Telephone (609) 258-0865

IRB Website | Email the IRB Office | Help

Appendix 1 EForm Name: IRB Application Page: N. Supporting Documents **Section**: Upload **Question**: survey example.docx File Name:

The survey is completed electronically. Each question is presented on its own screen. Once the answer is typed into the box and the participant presses enter, the survey moves to the next screen with a new question. There are approximately 945 questions for the entire survey. It will take approximately 25-45 minutes to complete the survey and you will receive \$6 in compensation. An example of a single screen is below.

To view the complete electronic survey, visit: www.testable.org/t/175f4613c

image on screen until you type your answer and press ENTER	
I began my illustrious career in journalism in the Bronx	
	image on screen until you type your answer and press ENTER

Appendix 2 EForm Name: IRB Application Page: N. Supporting Documents **Section**: Upload **Question**: File Name: survey example.docx

The survey is completed electronically. Each question is presented on its own screen. Once the answer is typed into the box and the participant presses enter, the survey moves to the next screen with a new question. There are approximately 900 questions for the entire survey, and an example of a single screen is below.

To view the complete electronic survey, visit: www.testable.org/t/175f4613c

Example	:	
1/935		
	image on screen until you type your answer and press ENTER	
	I began my illustrious career in journalism in the Bronx	

Appendix 3 EForm Name: IRB Application Page: N. Supporting Documents **Section**: Upload **Question**: consent_mturk_survey.docx File Name:



TITLE OF RESEARCH: Using Amazon Mechanical Turk to create and norm stimulus materials

PRINCIPAL INVESTIGATOR: Uri Hasson (hasson@princeton.edu)

RESEARCHER: Amy Price (aprice@princeton.edu)

PRINCIPAL INVESTIGATOR'S DEPARTMENT: Psychology and Neuroscience

You are being invited to take part in a survey being conducted by the Hasson Lab in the Department of Psychology at Princeton University. Before you decide to participate in this study, it is important that you understand why the research is being done and what it will involve. Please take the time to read the following information carefully, as it contains important information about the study and your rights as a research participant. Please email the researcher at aprice@princeton.edu if there is anything that is not clear or if you need more information.

Purpose of the research:

You have been asked to participate in a study to learn more how we make predictions about upcoming words in a sentence. To understand how easily you can predict an upcoming word in a story, you will read a sequence of words and then try to predict the next word in the story by typing that word into the blank box in the survey. There is no right or wrong answer for each question and you will not receive feedback for your answers.

Study Procedures:

If you choose to participate in this study, you will be asked to read text and type words. Your total expected time commitment for this study is 25 - 45 minutes. There are 945 questions in the survey. You will be asked to pay careful attention to the text as your read the story, as you may be asked to answer questions about the story.

Benefits and Risks:

Benefits: There are no direct benefits to participating in this experiment. However, you may draw satisfaction from the knowledge that information generated by this study may help promote better understanding of how we communicate with each other and process complex stimuli. This knowledge may eventually lead to improvements in the diagnosis, treatment and cure of neurological and psychiatric illness.

Risks: The risks involved in this study are minimal. If you experience any discomfort with the stimuli, please let us know by email. You may end the study at any time.

Confidentiality:

All records from this study will be kept confidential. Your responses will be kept private, and we will not include any information that will make it possible to identify you in any report we might publish. Research records will be stored securely in a locked cabinet and/or on password-protected computers. The research team will be the only party that will have access to your data. Anonymous data (without your name) may be presented at research meetings and published in research journals.

Compensation:

You will be paid \$6 for participation in the survey.

I understand the information that was presented and that:

- A. My participation is voluntary, and I may withdraw my consent and discontinue participation in the project at any time. My refusal to participate will not result in any penalty.
- B. I do not waive any legal rights or release Princeton University, its agents, or you from liability for negligence.
- 3. I hereby give my consent to be the subject of your research.

Click the box if you agree:

Date:

Who to contact with questions:

1. PRINCIPAL INVESTIGATOR:

Uri Hasson, Professor of Psychology

Phone: 609-258-3884

Email: hasson@princeton.edu

2. RESEARCHER:

Amy Price, Postdoctoral Fellow Email: aprice@princeton.edu

3. If you have questions regarding your rights as a research subject, or if problems arise which you do not feel you can discuss with the Investigator, please contact the Institutional Review Board at:

Assistant Director, Research Integrity and Assurance

Phone: (609) 258-8543 Email: <u>irb@princeton.edu</u>

Appendix 4 **EForm Name**: IRB Application N. Supporting Documents Page: **Section**: Upload **Question**: cover letter for modifications.docx File Name:

MEMORANDUM 10/4/2017

To: Institutional Review Board

From: Amy Price, postdoctoral fellow Hasson lab

Re: Modifications required for 10087 Using Amazon Mechanical Turk to create and

norm stimulus materials

Thank you for your timely review and modifications request!

We have corrected the following:

- 1) Sections E(19) and (20) of the application now show the same number of subjects.
- 2) We have added a consent form (attached to the new submission).
- 3) The survey now states that it contains 945 questions. The duration for completing the 945 questions is 25-45 minutes which is the accurate duration.

We have also modified the demographic information we are requesting to collect. Due to NIH requirements, our funding requires that in addition to gender, age, and native language, we will also need to collect racial and ethnic categories from the participants. We have updated the 'Study Procedures' section to include this request.

Thank you, Amy Price

Appendix 5 **EForm Name**: IRB Application N. Supporting Documents Page: **Section**: **Question**: Upload PI approval.pdf File Name:

P. PI Approval Form - Initial Application
Protocol Number: 10087
Submission Number:10087-01
PI:Hasson, Uri
Study Title:
Using Amazon Mechanical Turk to create and norm stimulus materials
Date: 28-Sep-2017
By signing this form and submitting it to the IRB, I am agreeing to follow the obligations outlined in Princeton University SOP 207: Obligations of the Principal Investigator for Human Subjects Research.
PI Signature :
Date: 10/10/2017

Appendix 6	
EForm Name: IRB Application	
Page:	O. PI Assurance
Section:	
Question: File Name:	After completing the steps below, please upload a scan or picture of the signed PI Approval Form here. PI signature IRB initial submission form.pdf
r ne Name:	ri signature iko initiai suomission torm.pui

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