

## Protocol behavioural tasks

(1) Will you be following a FERB-approved research programme of your supervisor?

☐ Yes

☒ No

### New data collection

(1) Will you be using:

- ☐ Online collection of questionnaire data
- ☐ Offline collection of questionnaire data
- ☐ Open or semi-open interview (incl. focusgroups)
- ☐ Regular observations
- ☐ Observations recorded on audio, video
- ☐ Naturalistic observation
- ☐ Participant observation (anthropology)
- ☐ Intervention study (not in lab)
- ☒ Behavioural tasks

(2) Where will the study (data collection) be conducted? If this is abroad, please note that you have to be sure of the local ethical codes of conducts and permissions

☒ the Netherlands

☐ Abroad, outside the Netherlands. Please name the country

### Behavioural tasks

(1) The task to be performed by the participant does not present the participant with a probability and magnitude of harm or discomfort greater than ordinarily encountered in daily life.

☒ Correct

☐ Incorrect

(2) Total duration of the task(s) does not exceed 60 minutes per session

- ☒ Correct  
☐ Incorrect

(3) Stimuli that will be presented to participants (you can choose multiple options):

- ☒ Visual stimuli  
☐ Auditory stimuli  
☒ Tactile stimuli (stroking, vibrotactile stimulation)  
☒ Shocks  
☐ Other, please explain

(4) Please provide a short description of the relevant stimulus characteristics.

test

(5) The intensity of the stimuli is below a level that, intended or unintended, makes participants afraid, scared or gives them an unpleasant feeling

- ☒ Correct  
☐ Incorrect

(6) Data that will be collected (you can choose multiple options)

- ☐ Behavioural data (Answers to (short) questions, stimulus classification/preference, response times)  
☒ Motor response data (eye movement coordinates, gripping paths, etc.)  
☐ Electrophysiology (EEG), electromyography (EMG) or skin conductance levels (SCL)  
☐ Other, please explain

(7) Please provide a short description of the experimental setup (site of data collection, whether faculty equipment will be used)

test

(8) My research question concerns (you can choose multiple options):

- ☒ Stimulus characteristics  
☐ Personality characteristics (including mental health)  
☐ Societal subjects (e.g.: political preference, religious subjects)  
☐ Potentially sensitive behavioural subjects (e.g.: sexual behaviour, illegal behaviour)  
☐ Experience of violence, abuse or exploitation  
☒ Motor response data (eye movement coordinates, gripping paths, etc.)  
☐ Substance use (e.g.: drugs, alcohol)  
☐ Other, please explain

## Participants 1

(1) I will approach...

- ☒ Adults
- ☐ Children up to 12 years old
- ☐ Children from 13 to 15 years old
- ☐ Adolescents from 16 to 17 years old

(2) I will approach a vulnerable group

- ☒ No
  - ☐ Yes, please explain which vulnerable group you are working with and why this study cannot be carried out without this population.
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(3) I am going to approach the participants using

- ☒ Recruitment websites, where participants register themselves
  - ☐ Friends, familie, acquaintances
  - ☐ Recruitment agencies that approached the participants
  - ☐ Other, please explain
- 

(4) I use the following resources

- ☐ Letter/e-mail
  - ☒ SONA systems
  - ☐ Other, please explain
- 

(5) Participants receive PPU or financial compensation in proportion to their efforts

- ☒ Yes, please explain
  - ☐ No, please explain
- 

(6) I'm going to ask consent to proceed using...

- ☐ A written procedure prior to data collection
- ☐ Online, by ticking a box
- ☒ An oral procedure prior to data collection (c.f. [this description](#))

(7) This investigation cannot lead to coincidental findings

- ☒ Correct

☐ Incorrect, please explain which coincidental findings may appear and how you will deal with this situation

(8) This study does not use deception

☒ Correct

☐ Incorrect, it does use deception. Please explain

(9) If deception is used, I will ask permission from the participants again immediately after the data collection

☐ Not applicable

☒ Yes

☐ No, please explain

**In accordance with the instruction document the information letter contains these elements:**

(11) Aim of the study

☒ Yes

☐ No, please explain

(12) It is emphasized that it concerns student research

☒ Yes

☐ No, please explain

(13) Type of tasks, duration, load

☒ Yes

☐ No, please explain

(14) How the data are handled

☒ Yes

☐ No, please explain

(15) Right of removal (unless completely anonymous)

☒ Yes

☐ No, please explain

(16) Being able to stop voluntarily at any time without adverse consequences

☒ Yes

☐ No, please explain

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(17) Contact person for questions

- ☒ Yes  
☐ No, please explain
- 

(18) Information letter in understandable language tailored to the target group

- ☒ Yes  
☐ No, please explain
- 

(19) Participants will be given the opportunity to remove their personal data

- ☒ Yes  
☐ No, please explain
- 

### **Informed consent**

(1) For adults, I ask informed consent

- ☒ Actively (via 'wet' signature)  
☐ Online  
☐ Passively (explain why not actively)  
☐ No informed consent asked (explain)
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### **Data management 1**

(1) Data are or will be stored on faculty servers (YODA and/or FSBS research storage) in accordance with faculty protocol

- ☒ Yes  
☐ No, please explain
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(2) Access to data is limited to student and supervisor(s).

- ☒ Yes  
☐ No, please explain

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(3) Storage period is in accordance with faculty protocol and/or additional statutory provisions.

☒ Yes

☐ No, please explain

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(4) Data are not shared with external organization

☒ Correct

☐ No, please explain

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### Attachments

(1) Questionnaire

(2) Information letter(s)

(3) Informed consent form(s)

(9) Miscellaneous documents e.g. data set description (optional)

No files have been uploaded yet

### Signature

(1) Corona protocols

☒ I declare that I have read the relevant [UU Corona protocols](#), and will follow them as I perform the study

(2) I declare that I have completed the above truthfully, my Solis ID is: