(1) Will you be following a previously approved project or research programme/line? O Yes, a FERB approved research programme O Yes, a FERB approved research project O Yes, a project which was approved by another Social Science ethical review board (affiliated with Nethics) O Yes, a METC approved project O Yes, an ethical review board of an external organization, please explain 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

Regular observations

• the Netherlands

O Abroad, outside the Netherlands. Please name the country

(1) All possible measures have been taken to prevent subsequent identification of persons
Yes, what measures have been taken (transcription of audio, encoding of video)
O No, explain why not
(2) The observational data are linked to other data, e.g. questionnaire data
● No
O Yes, please explain
(3) What is observed cannot be interpreted as invasive
Correct
O Incorrect, please explain
Participants 1
(1) I will approach
$\mathbf{Z}_{\mathrm{Adults}}$
☑ 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
O Yes, please explain which vulnerable group you are working with and why this study cannot be carried
out without this population.
(3) I am going to approach the participants using
Recruitment websites, where participants register themselves
Friends, family acquaintances
Recruitment agencies that approached the participants
Other, please explain
(4) I use the following resources
Letter/e-mail
O _{SONA systems}
O Other, please explain

Observation

(5) Participants receive PPU or financial compensation in proportion to their efforts
O 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
O An oral procedure prior to data collection (c.f. <u>this description</u>)
(7) This investigation cannot lead to coincidental findings
Correct
O Incorrect, please explain which coincidental findings may appear and how you will deal with this situation
(8) This study does not use deception
Correct
O 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
O Not applicable
• Yes
O No, please explain
In accordance with the instruction document the information letter contains these elements:
(11) Aim of the study
• Yes
O No, please explain
(12) It is emphasized that it concerns student research
● Yes
O No, please explain
(13) Type of tasks, duration, load
Yes
O No, please explain

(14) How the data are handled
• Yes
O No, please explain
(15) Right of removal (unless completely anonymous)
• Yes
O No, please explain
(16) Being able to stop voluntarily at any time without adverse consequences
• Yes
O No, please explain
(17) Contact person for questions
• Yes
O No, please explain
(18) Information letter in understandable language tailored to the target group
• Yes
O No, please explain
(19) Participants will be given the opportunity to remove their personal data
• Yes
O No, please explain
<u>Informed consent</u>
(1) For adults, I ask informed consent
• Actively (via 'wet' signature)
O Online
O Passively (explain why not actively)
O No informed consent asked (explain)

(2) For children up to 12 years old, I ask informed consent from one parent

Actively (via 'wet' signature)
Online
O Passively (explain why not actively)
O No informed consent asked (explain)
(3) For children from 13 to 15 years old, I ask informed consent from one parent and the participant
Actively (via 'wet' signature)
O Online
O Passively (explain why not actively)
O No informed consent asked (explain)
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, because data are not allowed to leave the external institute where the research project takes place
O No, but a copy of the anonymized raw data will be stored on the faculty servers
O Other, please explain
(2) Access to data is limited to student and supervisor(s).
• Yes
O No, please explain
(3) Storage period is in accordance with faculty protocol and/or additional statutory provisions.
• Yes
O No, please explain
(4) Data are not shared with external organization
Correct
O No, please explain
<u>Attachments</u>
(1) Questionnaire
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(2) Information letter(s)
Testdocument uploaden in formed consent.docx deleted
(3) Informed consent form(s)

 $\underline{Test document_uploaden_in\ formed_consent.docx_deleted}$

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

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Signature

- (1) Corona protocols
- \square I declare that I have read the relevant $\underline{UU\ Corona\ protocols}$, and will follow them as I perform the study
- (2) I declare that I have completed the above truthfully, my Solis ID is:

12345