**PRIDE QUESTIONNAIRE FOR APPROVAL SINGLE STUDY**

(1.1) Title

(1.2) Study type

Single Study/project

(1.3) Division

(1.4) Start date of data collection

# Basic study information

1. Name(s), position(s) and division(s) of the responsible researcher(s): Name Position Division E-mail
2. Name(s), position(s) and division(s) of the executive researcher(s): Name Position Division E-mail
3. Research area/discipline:
4. What is the study's main objective (hypothesis)?
5. Primary funder of the study:
6. Does the study concern a multi-center project, e.g. in collaboration with other universities, a GGZ mental health care institution, or a university medical center?

* Yes
* No

1. 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

# Study details (I)

1. Will the study process personal data?  
    Yes No
2. Does your study exclusively concern the analysis of existing data, document or records? Where can the data be found?

Yes No

1. Are the sources of the existing data, documents or records publicly available?   
   Yes No
2. Will the data be processed by the principal investigator in such a manner that participants can be identified either directly or indirectly (through identifiers (such as a code) linked to them)?  
    Yes No
3. If the study uses de-identified (or pseudonymized) data, does the responsible or executive researchers have access to the key to the code permitting re-identification of the person whose data are being studied?

Yes No Not applicable

1. The research will involve only the use of anonymous survey procedures, interview procedures, the observation of public behavior or other 'light burden research activities' (e.g., classroom observations, collecting teaching materials, or collecting data from Caracal or Osiris (for SoTL-projects)  
    Yes No
2. Will participants be asked to report their own or others' sexual experiences, alcohol or drug use, or suicidal thoughts, and will their identities be known to you?

Yes No

1. Are any participants confined in a correctional or detention facility?  
    Yes No

(9) Will the research procedure harm or discomfort the individual in any way (e.g., research topic, or study activities)

Yes No

# Study details (II)

1. Will participants that are recruited be >16 years?  
    Yes No
2. Will participants that are recruited be mentally competent (wilsbekwaam in Dutch)?  
    Yes No
3. Will participants that are recruited provide active informed consent?

Yes No

1. Will the probability and magnitude of possible harm or discomfort anticipated in the research be greater than ordinarily encountered in daily life or during the performance of routine physical or psychological examinations or tests?  
    Yes No
2. Does the participant population contain vulnerable persons? (e.g., incapacitated, children, mentally challenged, traumatized, pregnant), or a taboo subject (e.g., own or others' sexual activity, hard drug use, suicide thoughts, religious belief, political preference)

Yes No

1. Will participants be subjected to:

Yes No

Inquiries into their sexual behavior or orientation ○ ◉

Inquiries into drug use (also alcohol, smoking, soft drugs) ○ ◉

Assessment of delinquency ○ ◉

Inquiries relating to religious or philosophical belief ○ ◉

Inquiries relating to political opinions ○ ◉

Inquiries into ethnic origin ○ ◉

Inquiries into trade Union membership ○ ◉

Inquiries into violent experiences ○ ◉

Inquiries into personal health ○ ◉

Inquiries into criminal convictions and offences ○ ◉

Shocking images/videos ○ ◉

Deception (information letter does not state real study objective) ○ ◉

Physical pain (electrical/ thermal shocks, noise) ○ ◉

Following orders behaviorally (by force, or outside the context of the lab with possible harmful consequences for the participant or his/her social environment?)

○ ◉

A new technique for data collection? ○ ◉

1. Will data of the following categories be processed:

Yes No

Photo data ○ ◉

Video data ○ ◉

Biological material (buccal, blood, hair) ○ ◉

Yes No

Genetic data ○ ◉

Biometric data (fingerprint, iris or retinal scan, voice recognition and face scan) ○ ◉

Directly identifying data (name, address, date of birth or a combination of those items) ○ ◉

# Study details (III)

1. What is the study's theoretical and practical relevance? (500 words max.):
2. What are the central hypotheses?
3. What is the study's design and procedure? (500 words max.):
4. Optional attachments:

1. What data collection instruments, stimuli and/or manipulations will be used? If data are collected online, please specify the platform used.
2. Optional attachments:
3. Please state which statistical procedures will be used.
4. Will a method be used that may, by coincidence, lead to findings of which the participant should be informed? If so, what actions will be taken in the case of a coincidental finding?  
    Yes No

# Participants

(1) Please provide reasons to justify why this particular group of participants should be subjected to these conditions.

(2) What possible risks could participating in the study hold for participants?:

(3) What measures are implemented to minimize risk for participants?

(4) How does the burden on the participants compare to the study's potential scientific contribution (theory formation, practical usability)?:

(5) What is the number of participants? Provide a power analysis and/or motivation for the number of participants. The current convention is a power of 0.80. If the study deviates from this power, the ERB would like you to justify why this is necessary :

(6) Age category of the participants

1. How will the participants be recruited

1. Please state any specific in- and exclusion criteria and how these are tested.

1. How much time will the prospective participants have to decide as to whether they will indeed participate in the study?:

1. Are the participants fully free to participate and terminate their participation whenever they want and without stating their grounds for doing so?  
    Yes No
2. Will the participants be in a dependent relationship with the researcher?  
    Yes No

(12) Is there an independent contact person or a general email address of a complaint officer to whom the participant can contact? Yes No

1. What time investment and effort will be requested from participants?

1. Will the participants be compensated for their efforts? How (financial reimbursement, travelling expenses, otherwise). What is the amount? How will the compensation be transferred to the participant?  
    Yes No
2. Will this compensation depend on certain conditions, such as the completion of the study?  
    Yes No

# Attachments

* Text (advert) for the recruitment of participants (optional)
* Information letter for participant (required, except for database studies)
* Consent form for participants (required, except for database studies)
* Written or oral feedback information (debriefing text) (optional)
* (Descriptions of) questionnaires (optional)
* (Descriptions of) measurement instruments (optional)
* Miscellaneous documents e.g. data set description (optional)
* Data management plan (required)
* Privacy review (required when personal data will be collected)
* Ethics and privacy quickscan (*optional*)