

Important Note: This Request for Waiver is not applicable to FDA regulated studies

Request for Waiver of the Requirement to Obtain Signed Consent from Subjects

(Attachment for Human Research Review Application Section VII(3))

- The only record linking the subject and the research would be the consent document and the principal risk would be the harm resulting from breach of confidentiality. (Note: Each subject must be asked whether they want documentation.)
Explain why:

-OR-

- The research presents no more than minimal risk* and involves no more procedures for which written consent is normally required. Explain why:

If documentation is waived, will the subjects be provided with a written statement regarding the research?

- YES - Attach copy of written statement that will be provided.
 NO – Explain below why a written statement is not necessary or appropriate.

**Minimal risks means that the probability and magnitude of harm or discomfort anticipated in the research are not greater in and of themselves than those ordinarily encountered in daily life or during the performance of routine physical or psychological examinations or tests.*