

IRB – Waiver of Parental Consent

For studies including participants under the age of 19

Describe exactly what you wish to waive.

Describe why the research involves no more than minimal risk to the subjects.

Describe why the waiver or alteration will not adversely affect the rights and welfare of the subjects.

Describe why the research could not practically be carried out without the waiver or alteration of informed consent.

Will subjects be provided with additional pertinent information after or during the research? If yes, describe how information will be provided to participants.