In many studies sponsors wish to collect information concerning the health status of a research subject's partner when that partner becomes pregnant. Even though pregnancy may be an exclusion criterion for subjects, or the protocol requires the use of birth control measures, pregnancy may occur. In this case the collection of information about the pregnant partner (before and/or after parturition) or child (following delivery) may only be obtained pursuant to documentation of informed consent and HIPAA authorization from the pregnant partner. This procedure does not imply that consent of the pregnant partner and documentation of permission to collect health information makes the pregnant partner an enrolled subject in the main part of the research study. Rather, this procedure provides ethical protection for the privacy and welfare of the pregnant partner.