Statement of Compliance

The University of Toledo Department for Human Research Protections and Institutional Review Boards (IRBs) oversee investigations conducted at the University to assure they adhere to the principles of The Belmont Report, The Declaration of Helsinki, The University of Toledo’s Federalwide Assurance for Protection of Human Subjects, Terms of Assurance, the requirements of FDA regulations 21 CFR Parts 50 and 56, HHS regulations 45 CFR 46, and International Conference on Harmonization (ICH) E6, Good Clinical Practice (GCP), as applicable.

The University of Toledo’s Federalwide Assurance and the corresponding Terms of Assurance provide the framework for the human research subject protection program and associated institutional review boards. Because an understanding of, and compliance with, the FWA and Terms of Assurance is vital to human subject research at UT, the DHRP requires all active research team members to complete training prior to the review of a research protocol.

The University of Toledo IRBs are duly constituted; have written policies and procedures for initial and continuing review of research; prepare written minutes of convened meetings; and retain records pertaining to the review and approval process. The complete DHRP Policies & Procedures document can be accessed on our website.

The University of Toledo:
IORG0000832    Effective through: February 19, 2012
FWA00010686    Effective through: March 26, 2012
Biomed IRB #1   IRB00006058
SBE IRB #2      IRB00001193
UT DHRP website http://www.utoledo.edu/research/RC/HumanSubs_Menu.html

Roland T. Skeel, M.D.
Chair, Biomedical Institutional Review Board