Informed Consent Form Template Revisions

In response to a new Federal law called the **Genetic Information Nondiscrimination Act (GINA)**, ORI has revised the “What Else Do You Need to Know?” section of the informed consent form (ICF) template to prompt investigators to include appropriate language when the research involves genetic testing and/or collection of genetic information. Sample GINA language is located in the Instructions for Informed Consent Document on ORI’s web site at:

(Medical IRB) [http://www.research.uky.edu/ori/FormsHELP/S2C.htm#GINA](http://www.research.uky.edu/ori/FormsHELP/S2C.htm#GINA) or (Nonmedical IRB) [http://www.research.uky.edu/ori/FormsHELP/S2C_NM.htm#GINA](http://www.research.uky.edu/ori/FormsHELP/S2C_NM.htm#GINA).

Also, the “Do You Have to Take Part in the Study?” section of the ICF template has been revised to provide applicable verbiage when student volunteers are recruited for the research.

Currently, the revisions are reflected only in the English version of the ICF template; translation for the Spanish version of the ICF template is pending.

To ensure you are using the most recent IRB application forms and consent templates, please regularly visit the ORI Human Research Forms web page [http://www.research.uky.edu/ori/human/HumanResearchForms.htm](http://www.research.uky.edu/ori/human/HumanResearchForms.htm) to download the materials applicable to your anticipated submission.

New item requested on the IRB Form A, “General Information Sheet”

ORI has recently revised “Form A”, the General Information Sheet, of the IRB application to include a request for the Principal Investigator’s ‘Link Blue’ ID (i.e, AD Account).

The Link Blue ID is necessary as ORI prepares for a future electronic system. If the PI’s Link Blue ID is not provided at this time, ORI will proceed with processing the IRB application, however, eventually if the Link Blue ID is not provided, the PI won’t be able to log into the electronic IRB system.

Preparing for the Future