MCW Guidance: Guidelines for Documentation of Human Subject Research

This document was created to provide guidance to the investigator and research team regarding the expectations for documentation of Human Subject Research Related Activities. The guidance document provides an overview of the Human Research Protection Program's (HRPP) expectations regarding the documentation of research related activities for studies regulated by FDA regulatory requirements, studies regulated by HHS regulatory requirements, and for those studies not subject to FDA or HHS regulatory requirements. The HRPP's expectations are based upon regulatory requirements, state law, Medical College of Wisconsin's policies and the HRPP SOPs.

In the table the left column lists various examples of research related documentation and in the subsequent columns to the right are the HRPP's expectations for documentation to be included in the research records for FDA governed studies with and without ICH GCP E6, HHS regulated studies and finally studies that are not governed by FDA or HHS regulatory requirements but are subject to MCW policies and SOPs.

These guidelines were reviewed and are supported by the MCW Human Research Advisory Committee in the efforts of promoting transparent information regarding the conduct of human research projects.

Guidelines for Documentation of Human Subject Research

Documentation	FDA governed	FDA	HHS governed	No federal regulatory	
	studies with	governed	studies (federally	oversight, subject to MCW	
	ICH GCP E6	studies	funded)	policies and SOPs	
Regulatory file	Х	Х	Х	X	