## IRB Tip: Quality Improvement (QI) Projects and IRB Review

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QI activities oftentimes occupy an uncertain territory between clinical care and research. Although most QI activities involve a systematic investigation of some question of interest, many do not involve research as defined in the federal regulations for the protection of human subjects ( 45 CFR 46). In other cases, QI activities are designed to accomplish both a research purpose as well as a non-research purpose and IRB review is required. The following chart formulates some criteria that may be helpful in determining whether a QI project involves a research component and must be reviewed under the regulations for the protection of human subjects:

## QI projects may include a research component if:

- One purpose of the project is to develop generalizable results by testing a hypothesis OR by establishing a clinical practice standard where none exists
- Study procedures involve applying a new intervention that is beyond current standard practice
- Study procedures involve randomizing subjects into different intervention groups
- The project imposes risks or burdens to patients beyond those associated with the standard of practice
- One outcome sought is to generate an analysis that can be applied to other programs, processes or systems


## QI projects do not include a research component if:

- The only purpose of the project is to assess or improve a process, program, or system OR improve performance as judged by established/accepted standards
- The only procedures involve standard practices, interventions or treatments
- The only study procedures involve observing or comparing interventions that are already being done
- The project does not impose additional risks on patients, with the exception of privacy/confidentiality concerns
- The only outcome sought is an improvement in a program, process, or system

