Self-assessment is a valuable activity that can be informative in a variety of settings, including informed consent for clinical research. Now you can proactively discover your informed consent process strengths and areas for improvement using a new Informed Consent Self-Assessment tool!

To support and empower SOM clinical research personnel in conducting safe, efficient, and high-quality research, the CRSO has developed a new Informed Consent (IC) self-assessment tool.

The IC self-assessment tool will enable study teams to ensure that their informed consent processes are conducted in compliance with the IRB-approved consent procedures and federal and institutional requirements for informed consent. It can be used for any study as an additional level of confidence but may be of particular importance for investigator-initiated studies where the investigator has more responsibility and may have less independent monitoring.

The IC self-assessment can be completed by the investigator or a qualified member of the study team. It can be completed at various timepoints during the study for quality assurance or in preparation for a monitoring visit.

Take a moment to explore how the IC self-assessment tool will work for your clinical research study!

**Informed Consent Self-Assessment Tool**

**Coming up!**

The Informed Consent Self-Assessment is the first of several planned self-assessment tools.
Stay tuned for the next self-assessment on training and delegation. If you have a suggestion for a self-assessment tool, please let us know.