

IRBs are charged with ensuring that risks to subjects are minimized and risks are reasonable in relation to anticipated benefits. When research involves certain hazards, the CW IRB relies on supplemental information obtained from expert review (and approval) of proposed research by applicable safety committees.

If the imaging is being done **solely for research purposes** the research must first be reviewed and approved by the appropriate safety committee. Research that involves the use of X-rays (including DEXA) or CT scan is reviewed by the CW Radiation Safety Research Subcommittee and possibly Wisconsin's Department of Health and Human Services. Research involving MRI or PET/MRI is reviewed by the CW MRI Safety Committee. This safety committee approval must be included in the submission to the IRB.

Even if all the imaging being done as part of the research study is considered **standard of care**, if the imaging will be done at CW/with CW equipment as part of a research study (it is described in the protocol), the CW imaging department needs to be notified/consulted prior to submitting the study to the IRB. This is to review and plan for logistical considerations and is different than a safety review by the safety committee. Beyond any safety considerations, the imaging department needs to be able to plan for staffing, scheduling, use of the equipment and work out logistics in order to be able to support the study efficiently and without disruption to the department.

Appropriate safety committee approval should be sought out and obtained **BEFORE** submitting a project to the CW IRB, and the approval letter must be included with the initial submission. If safety committee approval is indicated, final IRB approval will not be issued until this is secured.

This should be obtained for any new projects, as applicable, or if a project is modified in a way that would make safety committee approval applicable.

	MCW Institutional Biosafety Committee	CW Radiation Safety Research Sub-Committee	State of Wisconsin Radiation Safety Review	CW MRI Safety Committee
How do I know if safety committee review is needed?	<p>When a research project involves use of any biologically hazardous or genetically engineered materials*. IBC approval is no longer required for IRB submissions that deal with only human source material unless the research involves the generation of cell lines, extensive processing of tissue, or work associated with animal studies.</p> <p>IBC review and approval is required for IRB submissions when deferred to non-MCW IRBs.</p> <p>Information on the IBC can be found here</p>	<p>If the imaging is for research only (beyond standard of care), then the radiation safety committee must review and approve before submitting to the IRB. There may be additional review by the State of Wisconsin required as well.</p>	<p>CW Local radiation safety committee will indicate in their approval letter if review by the state is also needed</p>	<p>If your research project involves imaging on a CW MRI scanner for research purposes only (beyond standard of care) the CW MRI Safety Committee must review and approve before submitting to the IRB.</p> <p>If the project involves use of an MRI scanner at Froedtert or MCW this will need to go to the MCW MRI Safety Committee. If the MRI scanner is housed outside of a clinical area there may be additional considerations when used in a pediatric population. If this is the case, contact the CW IRB office early to discuss.</p>
When should I request this review?	<p>Before submission to the IRB. Submissions are required at least two weeks prior to a meeting date in order to be scheduled on that meeting's agenda.</p>	<p>Before submission to the IRB. This should be submitted to the CW Radiation Safety Committee as early in the process as possible for review of both safety and logistics.</p>	<p>Before submission to the IRB, if review is indicated by Radiation Safety Committee.</p>	<p>Before submission to the IRB. This should be submitted to the MRI safety committee as early in the process as possible for review of both safety and logistics.</p>

* infectious agents; biological toxins; all recombinant DNA, both exempt and non-exempt; and all non-human primate derived materials including blood, tissue and bodily fluids.

Kids deserve the best.

<p>How do I request this review?</p>	<p>Applications are submitted through the MCW eBridge system. For direction/instructions on</p>	<p>Send an email to Linda Strain at LStrain@CW.org requesting the review, with documents listed below attached.</p>	<p>Email William Balke at William.Balke@dhs.wisconsin.gov Requesting the review with documents listed below attached</p>	<p>Currently no specific application form. Email Linda Strain at LStrain@CW.org and she will get a review scheduled.</p>
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	submitting an application through eBridge, please visit eBridge Support .			
What documents are needed?	Follow eBridge study application directions and contact MCW Biosafety Officer at 414-955-8060 or IBCsafety@MCW.edu with submission questions.	Research Protocol Informed consent/assent Radiation Procedure Grid	Letter from CW radiation safety Appendix M Variance Research Protocol Informed Consent/Assent	Study Protocol Imaging Guide Any documents that detail the MRI imaging needed (i.e. consent documents)
When are the meetings?	Meetings are held the second Tuesday of every month from 1:30-3:00pm in MFRC 3075, unless otherwise noted	These reviews are done on an ad hoc basis as they are requested	These reviews are done on an ad hoc basis as they are requested; response time is dependent on reviewer schedule.	The MRI Safety Committee meets once per month on the 4th Thursday of the month from 1:00 to 2:00 PM. The meetings are held in the Imaging Conference Room which is in the Imaging Department- first floor of the hospital.
Does the PI or a member of the study team need to be at the review meeting?	No, the PI and/or study team does not need to be present for the review of the submission.	No	NA (No)	The committee would like to have the study coordinator and principal investigator attend the meeting if possible.
How is approval communicated?	Via letter in eBridge	An approval letter will be sent/emailed to the PI/requestor of the review.	Typically, there will be an email response to the submitter followed by a mailing of the hard copy of the decision letter.	Email is utilized for requesting additional information or answering questions. An MRI Safety Review document is emailed to the study coordinator within a few days of the MRI Safety Committee meeting.
How long is the approval valid?	IBC approval is valid for 3 years. If the study changes the way they are using biologicals or adds new	Approval is valid for 3 years. If the study changes the use of radiation, or adds new radiation exposure, this should be re-	As long as the study is under IRB oversight and the subjects or parameters initially approved by DHSS remain unchanged there is	Approval is valid for 3 years. If the study changes the use of MRI, or adds additional MRI imaging, this should be re-reviewed by this safety

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	biologicals, an amendment to the submission must be submitted via eBridge. New work that is added to the amendment may not begin until the amendment has been approved by the IBC.	reviewed by this safety committee and their assessment submitted with the amendment to the IRB.	no need for a re-review by the state for a new letter. An updated approval letter is only needed if something about the radiation use in the study changes.	committee and their assessment submitted with the amendment to the IRB.
How long does approval take?	IBC review times vary. If the submission is time sensitive, please contact the IBC Office at 414-955-8060 or (ibcsafety@mcw.edu) so that appropriate arrangements can be made for the submission.	Varies by study complexity and reviewer schedules – range is a couple of days to a couple of weeks.	Varies depending on study complexity and DHHS reviewer's schedule/availability. Typically this will take a few weeks.	The time needed is based on the time of request in relation to the next scheduled meeting date. Studies are generally reviewed at the next scheduled meeting unless the agenda is full (this would be unusual). The approval document can normally be provided within a few days of the meeting.

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