



**Medical College of Wisconsin Cancer Center
New Trial Submission Form**

Principal Investigator:			
Full Protocol Title:			
Patient-friendly Title:			
Planned study site(s):	<input type="checkbox"/> Froedtert <input type="checkbox"/> CHW <input type="checkbox"/> CMH <input type="checkbox"/> SJH <input type="checkbox"/> Drexel <input type="checkbox"/> Community		
Study Overview			
Type of Study	<input type="checkbox"/> MCW Investigator-Initiated <input type="checkbox"/> NCTN/CTN <input type="checkbox"/> External Institutional <input type="checkbox"/> Industry/Pharmaceutical <input type="checkbox"/> Consortium <input type="checkbox"/> Other _____		
	<input type="checkbox"/> Drug <input type="checkbox"/> Device <input type="checkbox"/> Radiation <input type="checkbox"/> Surgical <input type="checkbox"/> Behavioral/Education Intervention <input type="checkbox"/> Observational <input type="checkbox"/> Other _____		
	Scope of trial: <input type="checkbox"/> Local (MCW/community) <input type="checkbox"/> National/Multisite		
	<input type="checkbox"/> Treatment <input type="checkbox"/> Diagnostic <input type="checkbox"/> Epidemiologic/Observational <input type="checkbox"/> Supportive Care <input type="checkbox"/> Device Feasibility <input type="checkbox"/> Ancillary <input type="checkbox"/> Screening <input type="checkbox"/> Health Services Research <input type="checkbox"/> Correlative <input type="checkbox"/> Prevention <input type="checkbox"/> Basic Science <input type="checkbox"/> Other _____		
Phase of Study	<input type="checkbox"/> I <input type="checkbox"/> I/II <input type="checkbox"/> II <input type="checkbox"/> II/III <input type="checkbox"/> III <input type="checkbox"/> III/IV <input type="checkbox"/> IV <input type="checkbox"/> N/A <input type="checkbox"/> Early Phase I <input type="checkbox"/> Other _____	Pilot study? <input type="checkbox"/> Yes <input type="checkbox"/> No	
Authorship	Is authorship likely? <input type="checkbox"/> Yes <input type="checkbox"/> No If yes: <input type="checkbox"/> First/last author <input type="checkbox"/> Middle author Comments:		
Accrual			
Local accrual	Projected annual accrual Overall accrual duration (months) Overall local accrual goal How many patients with this specific disease are seen at our institution per year (include source of data for expected enrollment, e.g. tumor registry, EPIC, CDW, etc.)?		
National accrual	Overall target accrual goal: Current overall enrollment:	Date accrual opened nationally: Expected closed to accrual date:	
Rare disease	<input type="checkbox"/> Check box if annual incidence is ≤ 4 newly diagnosed persons per 100,000 persons in U.S. (rare cancer, rare molecular subtype of common cancer, or unusual clinical situation)		
Competing Trials			
Will this study compete with any currently accruing or pending trials? <input type="checkbox"/> Yes <input type="checkbox"/> No If yes, indicate which trial(s) and describe prioritization plan for enrollment:			



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Funding Source			
<input type="checkbox"/> NCTN/CTN	<input type="checkbox"/> Pharmaceutical	<input type="checkbox"/> MCW Cancer Center	<input type="checkbox"/> There is no funding for this study.
<input type="checkbox"/> Consortium	<input type="checkbox"/> Department	<input type="checkbox"/> Other _____	<input type="checkbox"/> Additional funding is needed.
<input type="checkbox"/> NCI CTEP	<input type="checkbox"/> External Institution	Comments:	
Is the budget negotiable? <input type="checkbox"/> Yes <input type="checkbox"/> No			

For Investigator-Initiated Trials:
 Funding Source: _____ Funding Proposal #: _____
 Has funding been approved? Yes No Amount of award/approved funding: \$ _____

Study Complexity	
No. of Arms <input type="checkbox"/> 1 <input type="checkbox"/> 2 <input type="checkbox"/> 3 <input type="checkbox"/> ≥4	Eligibility Review <input type="checkbox"/> Basic <input type="checkbox"/> Complex/multi-step
Registration/ Randomization <input type="checkbox"/> One step <input type="checkbox"/> Multiple steps	Frequency of Study Tasks <input type="checkbox"/> Daily <input type="checkbox"/> Weekly <input type="checkbox"/> Every 21-30 days or more
Department/Team Impact	<input type="checkbox"/> One or two departments involved – Standard clinical research team <input type="checkbox"/> Three or more departments involved – Complex coordination needed <input type="checkbox"/> Inpatient Care Required
Radiology	Is there an imaging requirement in the protocol? <input type="checkbox"/> Yes <input type="checkbox"/> No If Yes- The requirements are: <input type="checkbox"/> standard <input type="checkbox"/> study-specific For IITs, has a radiologist been identified as a collaborator? <input type="checkbox"/> Yes <input type="checkbox"/> No
Ancillary Studies	<input type="checkbox"/> Banking <input type="checkbox"/> QoL <input type="checkbox"/> PK samples <input type="checkbox"/> Other
Data Collection on Treatment	<input type="checkbox"/> Basic – No AE reporting, batching of data <input type="checkbox"/> Standard – AE reporting and data collection <input type="checkbox"/> Complex – Real time submission, review of source documents for endpoints, etc.
Follow-up Requirements	<input type="checkbox"/> Annual or minimal follow-up <input type="checkbox"/> At each time point of clinical activity <input type="checkbox"/> Complex multiple clinical points
Special Requirements	<input type="checkbox"/> IND application <input type="checkbox"/> Clinicaltrials.gov <input type="checkbox"/> Coordinating center for multi-site study <input type="checkbox"/> Other
Beacon Build needed?	<input type="checkbox"/> Yes <input type="checkbox"/> No

Additional Comments

Disease-Oriented Team approval to send to SRC:

DOT Chair Signature	Date