Ancillary reviews are reviews by other compliance groups or individuals that inform the IRB's review of a new study or a modification to an existing study.

- Ancillary reviews may be assigned by either the researcher or the IRB.
- Once an ancillary review is triggered, researchers should work directly with those entities to ensure compliance.
- The ancillary review in Huron IRB is intended to support compliance across multiple oversight groups and not replace review processes by other compliance groups.
- Ancillary reviews are not assigned by the IRB if a project does not meet the federal definition of Human Subject Research.

The impact of an ancillary review group's approval on the IRB's review process varies.

- Typically, final IRB approval is held until the required ancillary group concludes their review.
- In some instances, the IRB will not initiate its review without documentation of approval by critical review entities.
- The IRB will not hold for the completion of ancillary reviews for studies that meet exempt criteria.
- Documentation of approval by an ancillary review group is provided to the researcher.
- In rare instances, either the ancillary review group or an IRB member may request deviations from the typical review path. An IRB member may recommend holding a submission until an ancillary approval is granted from a key committee **OR** an ancillary review group may recommend IRB review move forward while a required approval is still pending.
- Ancillary reviews that are required for IRB review/approval are not the same requirements for study activation. Study activation requirements are different and managed by Clinical Research Administration.

The tables below highlight the ancillary review groups available and illustrates the typical impact an ancillary review has on IRB review. Please contact the IRB or relevant ancillary review contacts (listed below) with any questions about the ancillary review process or specific requirements.

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	Ancillary Review I	Requirements for Humar	Subject Research Studies	i.	
Committee Name and	Initial study Other Parts of S		ications Study Team Member	Continuing	Training Requirements
Contact	·		, Info	Review	· ·
Cancer Protocol Review &	All cancer related studies	Protocol updates	N/A	N/A	N/A
Monitoring Committee	(retrospective or	require PRMC			
(CPRMC) IRB ID #: IRBAC001	prospective) require Protocol Review and	approval if PRMC is			
IKD ID #: IKDACUUI	Monitoring Committee	listed as an ancillary review committee.			
Sandra Rossi	(PRMC) review and approval	review committee.			
Manager, Research Support	PRIOR to IRB review.				
Sylvester Comprehensive					
Cancer Center	Please submit all study				
	related materials to the				
sandrarossi@med.miami.ed	PRMC via the PRMC				
<u>u or contact via Team</u>	Electronic Submission				
Questions	OPERA (Formerly PES)				
Questions: sccc.prmc@miami.edu					
<u>sccc.princ@mam.edu</u>					
For population science or					
social behavioral study					
specific requirements,					
please					
contact <u>sbs.prmc.startup@</u>					
miami.edu for more					
information.					
Clinical Trial Disclosure	This process facilitates the	N/A	N/A	N/A	N/A
Committee (CTD)	UM's compliance 42 CFR §				
IRB ID #: IRBAC002	11, FDAAA Section 801,				

*Please see training requirements; Approval is not required for personnel removal ** Approval is not required for initial studies

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Ancillary Review Requirements for Human Subject Research Studies					
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Contact	Initial study	Other Parts of Study	Study Team Member Info	Continuing Review	
Karre Wetherington, CCRP, CCA Sr. Clinical Trial Disclosure Associate Regulatory Affairs & Assessment Phone: (305) 243-1107 E-mail: kwetherington@miami.edu	 FDAMA Section 110, CMS, NIH and ICMJE. The CTD Ancillary Committee determines if a study must register on ClinicalTrials.gov. This determination includes: Is it a clinical trial? Who must register the study on ClinicalTrials.gov? Does the protocol have all information needed for registration and reporting? Does the informed consent form have the required CTD language? Clinical Trial Disclosure: Determination and Protocol Registration Policy https://umhs- ummg.policystat.com/policy /token_access/628beb33- 				

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	Ancillary Review F	Requirements for Humar	Subject Research Studies	i	
Committee Name and		Modif	ications	Continuing	
Contact	Initial study	Other Parts of Study	Study Team Member Info	Review	Training Requirements
	<u>c8ef-422e-b7f2-</u>				
	f79acb2dbe1e/				
	Clinical Trial Determination				
	Questionnaire				
Environmental Health and	EHS approval is required for	All modifications that	Adding personnel who	N/A	1) IBC Biosafety Training
Safety Committee (EHS)	studies that collect patient	introduce new risk	will be collecting		every three years for
IRB ID #: IRBAC003	specimens or introduce risk	group 2 agents (or	human specimens,		Clinical Staff via <u>ULearn</u>
	group 2 agents (or higher) or	higher) or any new	processing samples, or		
	any recombinant	recombinant	handling risk group 2		2) UHealth OSHA's
Primary point of contact:	therapeutics.	therapeutics not included in the	(or higher) agents*.		Bloodborne Pathogens, Biomedical Waste, Latex
BSO_Review@miami.edu		previous protocol.			Allergy and TB Training
Secondary Contacts at the		previous protocol.			required every year via
Biosafety Office:	Requirement: Complete the	Requirement:			ULearn
Shane Gillooly	Biosafety Ancillary Risk	Complete the			
Biosafety Manager	Assessment Form and	Biosafety Ancillary			
305-243-3269	upload via the Manage	Risk Assessment Form			
sxg1519@med.miami.edu	Ancillary Reviews activity in	and upload via the			
	IBISResearch-IRB.	Manage Ancillary			
		Reviews activity in			
Quintin James		IBISResearch-IRB.			
qaj3@miami.edu					
Embryonic Stem Cell	Research involving any work	Modifications to the	New personnel*	Yes	Ethical Oversight of HESC
Research Oversight	with the use of human	initially approved			Research. This is an
Committee (ESCRO)	embryonic stem cells and/or	study approved for			online module available
IRB ID #: IRBAC004	their derivatives must be	embryonic stem cells			via ULearn. Individuals

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Committee Name and		Modif	ications	Continuing	
Committee Name and Contact	Initial study	Other Parts of Study	Study Team Member Info	Continuing Review	Training Requirements
	approved by the UM	and/or their			listed or added onto a
Primary point of contact:	Embryonic Stem Cell	derivatives			ESCRO related protocol
Dr. Ellen Kapsalis	Research Oversight				must complete this
Director of Compliance	Committee (ESCRO) prior to	1. update to the			training once. Contact
IACUC / IBC / ESCRO	receipt of IRB approval.	IB or protocol			Liz Meza
		amendment			Senior Regulatory Analyst
ekapsali@med.miami.edu	Requirement: Such	2. protocol			IACUC / IBC / ESCRO
	submissions must be	closure			Imeza@miami.edu for
Secondary Contact:	submitted to	3. RNIs			more information.
	the ESCRO committee				
Liz Meza	outside of the IRB system.	Requirement: Such			
Senior Regulatory Analyst		submissions must be			
IACUC / IBC / ESCRO	Please visit	submitted to			
Imeza@miami.edu	the ESCRO committee	the ESCRO committee			
	website	outside of the IRB			
	at https://www.research.mi	system. The review			
	ami.edu/about/admin-	process for the ESCRO			
	areas/raa/escro/index.html	committee can occur			
		concurrently, but the			
	for further information.	HSRO cannot release			
		the IRB approval until			
		ESCRO signs off.			
Institutional BioSafety	All clinical trial protocols	Modifications to the	Adding personnel for	Yes	Biosafety Training is
Committee (IBC)	that use recombinant DNA,	initially approved	following roles:		required every three
IRB ID #: IRBAC005	synthetic nucleic acid	study approved for	1. Change of PI		years for all research
	materials, or a genetically	the use of			personnel and for new

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	Ancillary Review I	Requirements for Humar	Subject Research Studies		
Committee Name and Contact	Initial study	Modif Other Parts of Study	ications Study Team Member Info	Continuing Review	Training Requirements
Primary point of contact: IBCsupport@miami.edu Secondary Contact: Liz Meza Senior Regulatory Analyst IACUC / IBC / ESCRO Imeza@miami.edu	 modified organism or therapeutic must receive prior approval from the Institutional (IBC). Requirement: Please refer to the documents for Human Gene Transfer Research at Institutional Biosafety Committee (IBC) UResearch University of Miami. Such submissions must be submitted directly to the IBC committee outside of the IRB at IBCsupport@miami.edu. The review process for the IBC and the IRB can occur concurrently, but the HSRO cannot release the IRB approval until IBC signs off. 	recombinant DNA, synthetic nucleic acid materials 1. update to the IB or protocol amendment 2. Closure of study 3. RNIs Requirement : Please refer to the documents for Human Gene Transfer Research at Institutional Biosafety <u>Committee (IBC) </u> <u>UResearch </u> <u>University of Miami.</u>	 sub- investigator study coordinator Individuals administering study product (nurses, etc.) Lab personnel handing study product 		personnel added to the study during the research. Personnel includes anyone involved with the material/agent (whether administrating it or collecting samples or transporting the material across campus). This training is available via ULearn as a module titled "IBC Biosafety Training for Clinical Staff". IBC NIH Guidelines Training (must be completed once) by PIs, sub-investigators, co- investigators. This training is a PowerPoint presentation. Contact IBCsupport@miami.edu for review and credit.

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Committee Name and		Modif	ications	Continuiro	
Committee Name and Contact	Initial study	Other Parts of Study	Study Team Member Info	Continuing Review	Training Requirements
Human Use Radiation Safety Committee (HRSC) IRB ID #: IRBAC006	Protocols where radiation/radioactive materials (not MRI, Ultrasound or Laser Treatment) or radiation	Modifications that introduce radiation/radioactive materials (not MRI, Ultrasound or Laser	N/A	N/A	N/A
Sean O. Wilson Radiation Safety Officer 305-243-6360 sow10@med.miami.edu	producing devices are being used for research purposes	Treatment) or radiation producing devices to the parent study			
Diana R. Hernandez Radiation Control Manager 305-243-6360 drh85@med.miami.edu					
Conflict of Interest Committee (COIC) IRB ID #: IRBAC007 Lory Hayes, Ph.D. Director of DSAM (Disclosures & Scholarly Activities Management) LHayes@miami.edu	The UM COI Committee acts to determine, through a risk-based, case-by-case review, whether a COI is created between a research project and an external relationship. If a COI is found, the COI Committee works with the investigator to develop a management plan.	Modifications meeting criteria for review	Modifications meeting criteria for review	N/A	All investigators are required to complete UM's disclosure process and COI training prior to engaging in Scholarly Activities, repeated annually.
Call the UDisclose System helpline (305-243-0877) with					

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	Ancillary Review Requirements for Human Subject Research Studies				
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	Commission that disale sume		Info	neview	
questions about UM's COI policy or disclosure requirements.	Complete the disclosure process in the UDisclose System.				
Clinical Research Operations and Regulatory Support (CRORS) IRB ID #: IRBAC008 For New PI's: Alina Gjerpen Project Manager, Research and Innovative Medicine Clinical Research Operations and Regulatory Support 305-243-0492 arg136@med.miami.edu	CRORS ancillary review is required for new studies involving an investigator- held IND or IDE and for amendments to the studies.	N/A	N/A	N/A	N/A
For CRORS QC Monitoring: Nicole S. McCullough, MS, CCRP Director, Clinical Research Operations & Regulatory Support (CRORS) University of Miami, Miller School of Medicine Dominion Towers, 1400 N.W 10th Avenue, 1005L	Before the initial ancillary approval, the PI must contact CRORS for a monitoring cost estimate and to discuss the monitoring plan for the study.				

*Please see training requirements; Approval is not required for personnel removal ** Approval is not required for initial studies

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	Ancillary Review F	Requirements for Humar	Subject Research Studies	S	
Committee Name and		Modif	ications	Continuing	
Contact	Initial study	Other Parts of Study	Study Team Member Info	Review	Training Requirements
Ph: 305-243-0493 Fax: 305-243-5392 E-mail: nshank@med.miami.edu					
Data Security Ancillary Committee (DSAC) IRB ID #: IRBAC009	Studies collecting, storing, and transmitting protected health information (PHI).	Modifications meeting criteria for review	N/A	N/A	N/A
Primary point of contact: <u>dsac@miami.edu</u>	Forms: <u>Research Data Security</u> <u>Assessment Form</u>				
Secondary Contact: Joey Casanova, BBA Data Broker Manager 305-243-2631 jcasanova@miami.edu	For information on the review process for data privacy plans, please see <u>SOP-HB-008-01</u>				
Department Review	Based on department. Usually conducted by the Department Chair or the Chair's designee.	Required for PI transfer	N/A	N/A	N/A
UHealth Tower (UHT) ** IRB ID #: IRBAC010	UHT Ancillary Committee approval must be obtained for studies with any research activities at a UHT facility or any studies accessing UHT patient information, before	Modifications that are adding UHT to the study. Requirement: Please complete the UHT	N/A	N/A	N/A

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Committee Name and Contact	Initial study	Other Parts of Study	Study Team Member Info	Continuing Review	Training Requirements
Oncology Protocols Mitchell Diaz MSN, AGPCNP-BC Inpatient Research Nurse Specialist – Hematology/Oncology University of Miami Hospital and Clinics UHealth Tower O (305) 912-4633 mxd1433@miami.edu Danny Pino BSN, RN, OCN, BMTCN Inpatient Research Nurse Specialist – Hematology/Oncology University of Miami Hospital and Clinics UHealth Tower O (305) 912-4633 d.pino2@umiami.edu	using any UHT resources including subject recruitment, facility use, subject interventions such as tests, measurements, drug administration, surgery, or obtaining subject consent. UHT Ancillary Approval is required to determine feasibility of UHT Departments for clinical study protocols Requirement: Please complete the UHT Research Request form in the Link below: <u>https://redcap.miami.edu/s</u> <u>urveys/?s=HKP9E74X7ACHT</u> MDM and upload via the Manage Ancillary Reviews activity in IBISResearch-IRB.	Research Request form in the Link below: https://redcap.miami. edu/surveys/?s=HKP9 E74X7ACHTMDM and upload via the Manage Ancillary Reviews activity in IBISResearch-IRB.			

*Please see training requirements; Approval is not required for personnel removal ** Approval is not required for initial studies

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Committee Name and Contact	Initial study	Other Parts of Study	Study Team Member Info	Continuing Review	Training Requirements
Non-Oncology Protocols					
Carlos Sandoval Director, CTRS Research Operations Miami Miller School of Medicine Office of the Executive Dean for Research - EDR Office: (305)243-8842					
Clinical Translational Research Site (CTRS)** IRB ID #: IRBAC011 Carlos Sandoval Director, CTRS Research Operations Miami Miller School of Medicine Office of the Executive Dean for Research - EDR Office: (305)243-8842	Research protocols using the EDR Clinical Translational Research Site (CTRS) facilities must be reviewed by the CTRS Ancillary Committee. CTRS Provides the following services: Clinical Services (Nursing, Pharmacokinetics, Jackson Memorial Hospital Clinical Research Nursing Support, Laboratory processing, Clinical Study procedures,	N/A	N/A	N/A	N/A

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	In-Patient and Outpatient Clinical Services)				
	Clinical Research Coordinator Services (Consenting, Screening, Recruitment, Study visits, Documentation and data entry)				
	Requirement: Investigators should complete the <u>CTRS Services Requested</u> Form				
Pathology Review Committee (RPSC) ** IRB ID #: IRBAC012	All research studies using human samples including bodily fluids (blood, urine, ascites, saliva), and/or biospecimens not limited to	Required under the following circumstance: Protocol updates	N/A	Protocol updates including human samples	N/A
Omar Aljuboori Translational Research Scientist Department of Pathology 305-243-9453 oxa335@miami.edu	frozen tissue, fresh tissue, or archived tissue (paraffin embedded/FFPE), and/or slides must be reviewed by the Department of	including human samples			

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	Pathology and Laboratory				
Secondary Contact:	Medicine.				
Sophie Egea, PhD					
Director, Clinical Research	Review by pathology				
Department of Pathology	ancillary review committee				
and Laboratory Medicine	is required to process all				
	requests using human				
Research Mailbox:	samples for research.				
DPLMresearch@med.miami.					
<u>edu</u>	Please fill out the form using				
	this link: <u>PRC Form</u> .				
	The Department of				
	Pathology and Laboratory				
	Medicine has 2 human				
	research cores: Translational				
	Research Histopathology				
	Laboratory (TRHL) and				
	Laboratory of Clinical				
	Biosciences (LCB). Pathology				
	Ancillary Review approval is				
	required before placing iLab				
	requests.				
Clinical Research Laboratory	Research using Clinical	N/A	N/A	N/A	N/A
Services (formerly known as	Research Laboratory				
SCCC Research Lab &	Services (formerly known as				

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	Ancillary Review Requirements for Human Subject Research Studies					
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Contact		Other Parts of Study	Study Team Member Info	Continuing Review	Training Requirements	
Satellites- SCCC) ** IRB ID #: IRBAC013 Allie Bivin-Martinez Director, CRLS 305-243-6618 (office) scccresearchlab@med.mia mi.edu	SCCC Research Lab & Satellites) facilities must be reviewed by the CRLS lab staff prior to any research lab utilization. Coordinators should fill out the <u>SCCC Research Lab &</u> <u>Satellites Services Request</u> <u>Form</u> and send it to: scccresearchlab@med.miam i.edu					
	This form must also be uploaded via the Manage Ancillary Reviews activity in IBISResearch-IRB.					
Jackson Health System – Clinical Research Review Committee – (CRRC) ** IRB ID #: IRBAC014 Katuska Barbery, MBA Director of Clinical Research, JHS Office of Research	Approval from the JHS-CRRC must be obtained for studies with any research activities (including recruitment of subjects, facilities use, or subject interventions such as tests, measurements, drug administration, surgery, or consenting subjects) occurring at a JHS facility or any studies that	N/A	N/A	N/A	N/A	

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Ancillary Review Requirements for Human Subject Research Studies					
Committee Name and Contact	Initial study	Modif Other Parts of Study	ications Study Team Member Info	Continuing Review	Training Requirements
Jackson Health System 305-585-7226 Katuska.Barbery@jhsmiami. org	<pre>involve accessing JHS patient information prior to the use of any JHS resources. Please complete the Jackson Clinical Trials Office Application Form and upload to the IRB system via the Manage Ancillary Reviews activity, and send the JHS Study Calendar directly to the CTO. https://jhsmiami.org/jhsoffi ceresearch/ Additional information regarding the JHS CTO approval process can be found at: https://www.ora.miami.ed u/forms-and- rates/forms/clinical- research-forms/index.html</pre>				

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Contact	Initial study	Other Parts of Study	Study Team Member Info	Review	Training Requirements
Schiff Center for Liver	N/A	N/A	N/A	N/A	N/A
Diseases **					
IRB ID #: IRBAC015					
Sonia Carvalho					
Director, Regulatory Support					
scarvalho@med.miami.edu					
(305) 2434639					
SONHS Simulation	N/A	N/A	N/A	N/A	N/A
Hospital **					
IRB ID #: IRBAC016					
Victoria Behar-Zusman					
Professor, School of Nursing					
and Health Studies					
vbehar-zusman@miami.edu					
(305) 2849139					
Christian Ruiz DeGennaro					
Director, Research Support					
<u>cruiz@miami.edu</u>					
JFK Hospital **	N/A	N/A	N/A	N/A	N/A
IRB ID #: IRBAC018					
Jill Kinley, DNP, APRN, CCRC					
Director of Clinical Research					
Jill.Kinley@HCAHealthcare.					

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Contact		Other Parts of Study	Study Team Member Info	Review	Training Requirements	
<u>com</u> (561) 548-1414						
MCA Review Group (ORA)	All Submissions where question #5 in "Study	Revised protocols	PI changes only	N/A	N/A	
Tatyana Vikhlyantseva, Director, Research	Scope" is answered "yes".					
Administration	The purpose of this review is					
TVikhlyantseva@med.miami	to confirm if a Medicare					
<u>.edu</u>	Coverage Analysis (MCA) is					
305-284-3942	needed.					
Bianca Krysztof, Sr.						
Manager, Research						
Administration						
b.gjorgievski@miami.edu 305-284-1735						
505-264-1755						
Amy Gonzalez, Sr. Contract						
and Grants Analyst						
aeg183@miami.edu						
305-284-3509						