

Internal Medicine

Academic Research Services (ARS)
 Medical Sciences Building (MSB) Room 6111
 Email: imresearch@ucmail.uc.edu



REQUEST FOR REGULATORY SERVICES - NEW PROJECT

INSTRUCTIONS:

1. Please complete and send to IMRegulatory@uc.edu.
2. Attach the final protocol and supporting documents with your initial email request.
3. Your project will be reviewed and placed in our queue.
4. Within 5 business days or less, the regulatory team will communicate a start date for work on your project.

It is imperative that you complete this form thoroughly and completely. All regulatory documents are created utilizing the information provided. **Please take time to correctly answer the facilities and study team information.**

QUESTIONS:

General questions about the form or process: (Helen) Gina Shelton, 513-558-7183 or sheltoHn@ucmail.uc.edu

We look forward to providing you with our services and collaborating with you on this project. Thank you!

Principal Investigator:	<i>Name:</i>	
Sponsor or CRO Regulatory Contact - if applicable **Mandatory for Sponsored Studies**	<i>Name:</i>	
	<i>Email:</i>	<i>Phone:</i>
Individual Completing this Intake Form:	<i>Name:</i>	
	<i>Email:</i>	<i>Phone:</i>
<input type="checkbox"/> Yes <input type="checkbox"/> No	I acknowledge that there may be a non-refundable fee associated with the work for this project. This intake form serves as the PI's approval of the IM Regulatory Service fees. Please review the fees prior to submitting this intake form. ARS IM Regulatory Fee Schedule	
<input type="checkbox"/> Yes <input type="checkbox"/> No	I acknowledge that site selection and ARS review are complete (as applicable) and the project is ready to move forward with IRB Submission. I acknowledge that my regulatory documents will be created utilizing the information in this form and I am ready to proceed with the regulatory process. I also acknowledge that if the information provided is incomplete or inaccurate and all regulatory startup documents need to be revised due to incomplete information, there may be additional fees charged by IMRegulatory services for those revisions.	

SERVICES PROVIDED

Regulatory services will be provided for this initial request and covered under the initial fees for this project.

Study renewals will be billed at the time the Continuing Review is submitted to the IRB of record and will continue to be renewed until the IMRegulatory CRP is notified to close the study.

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PROJECT DETAILS

General	Please Indicate	
Protocol Number		
Study Title		
Sponsor	<input type="checkbox"/> Industry Pharma company, please indicate <input type="checkbox"/> Investigator-Initiated <input type="checkbox"/> Grant/Federal funding <input type="checkbox"/> Other	Sponsor Name:

PROJECT DETAILS- continued

Use the comments column and/or page 6 to provide additional details

General	Please Indicate	Comments/Details
<p>**Mandatory if your study has an ICF IRB mandated: Phone number to be listed on ICF that participants can call 24hr</p>		
IRB of Record	<input type="checkbox"/> UCIRB <input type="checkbox"/> Advarra <input type="checkbox"/> WIRB <input type="checkbox"/> Coprenicus Group (CGIRB) via WIRB <input type="checkbox"/> Other IRB Reliance *	
Source of Funding	<input type="checkbox"/> Federally Funded (NIH/FDA/NIAID etc.)* <input type="checkbox"/> Department Funded - (ex: DOIM awards)* <i>Primarily used for Investigator Initiated Studies</i> <input type="checkbox"/> Industry Funded (Pharmaceutical Companies) <input type="checkbox"/> Foundation Funded * <input type="checkbox"/> No Funding	

* Provide Detail In Comments

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PROJECT DETAILS- continued

Use the comments column and/or page 6 to provide additional details

General	Please Indicate	Comments/Details
Type of Study	<input type="checkbox"/> Drug Study <input type="checkbox"/> Trainee <input type="checkbox"/> Chart Review <input type="checkbox"/> Other <input type="checkbox"/> Device Study <input type="checkbox"/> Humanitarian Use Device (HUD) <input type="checkbox"/> Non-Human Subject Research (NHSR)	
HIPAA Waiver	<input type="checkbox"/> Chart Review (Full) <input type="checkbox"/> Screening/eligibility (partial)	
Facilities Utilized NAME AND ADDRESS OF ANY MEDICAL SCHOOL, HOSPITAL, OR OTHER RESEARCH FACILITY WHERE THE CLINICAL INVESTIGATION(S) WILL BE CONDUCTED ** Please review protocol and make sure you check all that apply	<input type="checkbox"/> University of Cincinnati Medical Center <input type="checkbox"/> West Chester Hospital -7700 University Drive <input type="checkbox"/> UC Health Physician's Office – West Chester <input type="checkbox"/> Hoxworth- 3130 Highland Ave. UC Health <input type="checkbox"/> Holmes- 200 Albert Sabin Way <input type="checkbox"/> UC Health Physician's Office- 222 Piedmont Ave. <input type="checkbox"/> University Endoscopy Center – 9275 Montgomery Rd. <input type="checkbox"/> UCMC Imaging Bldg. – 222 Piedmont Avenue Ste. 1400 <input type="checkbox"/> UC Gardner Neuroscience Institute – 3113 Bellevue Dr.	** Other, please indicate

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PROJECT DETAILS- continued

Use the comments column and/or page 6 to provide additional details

General	Please Indicate	Comments/Details
<p>Local Lab Information</p> <p>NAME AND ADDRESS OF ANY CLINICAL LABORATORY FACILITIES TO BE USED IN THE STUDY</p>	<p><input type="checkbox"/> <u>UCMC Local Lab - Safety Labs</u></p> <p><input type="checkbox"/> <u>IM Division Processing Lab – (UCPC Retrovirology Lab): Shipping/processing lab</u> **Submitter will need to reach out to divisionlabservices@ucmail.uc.edu to complete their intake process if you have not done so already. Before checking this box, please make sure you have secured approval to utilize their services.</p> <p><input type="checkbox"/> <u>ED Lab</u> - **before checking this box, please make sure you have secured approval from the ED lab to utilize their services.</p>	<p>** Other, please indicate</p>
<p>ANCILLARY SERVICES</p>	<p><input type="checkbox"/> <u>UC Health Imaging Services (Any radiology – including DXAs/CT Scan/X-Ray)</u> **Submitter will need to contact Monene Kamm at Research-DiagImaging@UCHealth.com if you have not done so already.</p> <p><small>**NOTE for Reg Team: If this is checked, send to RSC (Rich Anderson) for review of consent risk language and possible RSC committee review.</small></p> <p><input type="checkbox"/> <u>IDS PHARMACY</u> – Submitter should contact ids-pharmacy@uchealth.com</p> <p><input type="checkbox"/> <u>Infectious Diseases Research Pharmacy HOLMES (for ID use only)</u></p>	<p>** Other, please indicate</p>
<p>Supplemental Reviews</p>	<p>Is Institutional BioSafety Committee Review required for your study?</p> <p><input type="checkbox"/> YES <input type="checkbox"/> NO</p> <p><i>* IBC review is required if:</i></p> <ul style="list-style-type: none"> • Human derived materials are handled within a UC Research Lab (does not apply to clinical diagnostic lab) • Introduction of recombinant nucleic acids, infectious agents and/or genetically modified cells into human patients 	<p>** Other, please indicate</p>

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PROJECT DETAILS- continued

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	Name	Role
<p>Project Team</p> <p>INCLUDE ALL SUB-INVESTIGATORS, STUDY COORDINATORS, RESEARCH ASSISTANTS – ANYONE THAT NEEDS TO BE ON YOUR DELEGATION LOG</p>		
<p># of Participants you plan to enroll</p>		
	<p>If this is grant funded with an Industry sponsor component, is there a separate contract with the Industry in addition to the grant? <input type="checkbox"/> YES <input type="checkbox"/> NO <input type="checkbox"/> N/A</p>	

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Additional Information: *if sponsor is an industry Pharma Co., include template CTA injury text below*

Participant Payment YES* NO

** If yes please include the exact language you want in your consent (must match CTA/Budget)*

CTA Injury language for consent form:
(Must match CTA)

Comments and Questions:

**** See Page 7 for additional notes**



UC HEALTH APPROVAL INFORMATION

NOTE: UC HEALTH approval is not part of IMREG services.

Upon IRB approval, IMREG will send the IRB approval documents to the contact listed on this form and the PI. Please use the approval documents to submit for UC HEALTH approval. Enrollment cannot begin without UC HEALTH APPROVAL.

This is required for **all studies including chart reviews and all investigator initiated studies.

Please submit using the REDCap link below for that submission:
<https://survey.uchealth.com/redcap/surveys/?s=RJXFKLN3C7>

RESEARCH FINANCIAL SERVICES (RFS) INFORMATION

REMINDER: Be sure to send the following to Research Financial Services (RFS) to help with your contracting and budgeting:

- Site Selection Letter
- Informed Consent – (draft template is fine)
- Contract
- Protocol
- Budget
- Lab/Pharmacy Manuals

For more information please contact:

Leah Bischoff - bischoh@ucmail.uc.edu

Lisa Schira - shirala@ucmail.uc.edu

Please also remember to work on your **ClinCard** set up if applicable.
For more information, contact OCR.

[https://researchhow2.uc.edu/search?
indexCatalogue=researchhow2%
2Ddev&searchQuery=Greenphire&wordsMode=0](https://researchhow2.uc.edu/search?indexCatalogue=researchhow2%2Ddev&searchQuery=Greenphire&wordsMode=0)