A. Complete the UH Research Application Form (the following 3 pages)

Completion and review of the application ensures department and administrative approval is obtained.

- Projects that do not meet the federal definition of research = non-human subjects research (NHSR) or activities determined by the Institutional Review Board (IRB) as not human subjects research **DO NOT** require a UH Research Application Form. The UH research process: [https://med.umkc.edu/ora/conducting-research-at-UH/](https://med.umkc.edu/ora/conducting-research-at-UH/)

- The Principal Investigator (PI) or co-PI must be a member of the current UH Medical Staff or UH employee.

- All researchers and study personnel must complete the CITI education course for the protection of human research subjects ([Group 1-Biomedical](https://med.umkc.edu/ora/human_subjects/)) and renew every 3 years.

- Attach the appropriate items (study protocol, consent form, etc., listed on pg. #3) with the Application Form.

- To avoid bias and assure objectivity in research, for all sponsored research projects - researchers and staff shall disclose any potential conflicts and significant financial interests in accordance with the UH Code of Conduct, the sponsor’s requirements, and University of Missouri System conflict of interest disclosure requirements.

- For funded research projects, please forward the sponsor’s draft study agreement/contract and budget to Research Administration for review and negotiation as soon as the contract drafts are received.

- Research policies are posted on the UH internal ULink site: [tmcmed.sharepoint.com/](https://tmcmed.sharepoint.com/) on the UH network.

Office of Research Administration e-mail = [SOMResearch@umkc.edu](mailto:SOMResearch@umkc.edu) or call (816) 235-6015. **The office is located in the UMKC School of Medicine** on the 4th floor (M4-308).

B. Obtain approval from the Institutional Review Board (IRB)

Per an assurance with the U.S. Office of Human Research Protections, and institutional policies - research conducted at UH requires approval by the appropriate IRB. The IRB will not provide final approval to conduct the research until verification of administrative approval at UH has been obtained.

*Most* research at UH is reviewed by the UMKC IRB: [https://ors.umkc.edu/services/compliance/irb](https://ors.umkc.edu/services/compliance/irb) The UMKC IRB website has additional info about collaborative IRB review for projects that involve multiple institutions. The UMKC IRB utilizes an electronic IRB submission system.

Research requiring access, review, use, recording, or disclosure of any patient protected health information at UH also requires review to ensure compliance with HIPAA Privacy Rule requirements. The Privacy Rule requirements will be reviewed during IRB review.

*Please contact the IRB directly for questions about reviews, submission forms, or meeting information.*

You may submit/apply to A & B above at the same time.
1. Research Protocol Title:

2.  □ New project  □ On-going project  Date of planned study initiation

3. Research site  □ UH-HSD  □ UH-LW  □ UH-BH  □ UMKC SOM

4. Is this a sponsored research project?  □ Yes  □ No  Study Sponsor or separate research organization (if any)

5. Principal Investigator
   Phone  Pager  Email

   Faculty Mentor (if this is a resident’s or student’s research project)

6. Study Coordinator(s)
   Phone  Pager  Email

   List all other staff that will work directly on this project:

7. Please indicate who will pay the costs of treatment in the event a study participant suffers an injury during the conduct of this research project.

8. Indicate the study protocol procedures that are not ‘standard of care’ for this research project.

9. Will any advertisements (newspaper, radio, TV, internet, flyers, posters, etc.) be utilized?  □ Yes  □ No
   If Yes, the ad(s) requires approval by the IRB. In addition, UH Public Relations must approve the info as well.
10. Will study participants receive compensation for study participation?  □ Yes  □ No

If Yes, please indicate compensation method:

☐ Check or UH Cash Office stipend from UH research study account
☐ Gift certificate or gift card. Indicate source:
☐ Other:

*Please note, the amount and method of compensation also require IRB approval.*

11. Is the study sponsor providing any study recruitment incentives or bonuses to the site that are not mentioned in the study agreement/contract or budget?  □ Yes  □ No

If Yes, please describe:

12. Attach copies of these items with this application:

(1) ☐ Current Research Study Protocol.
(2) ☐ Study Contract / Grant Application (if applicable).
(3) ☐ Sponsor’s study Budget (if applicable).
(4) ☐ UH Research Expense Worksheet for funded projects (if applicable) – “internal budget”.
(5) ☐ Informed Consent form (if applicable). *Send the final IRB approved version when available.*
(6) ☐ For sponsored projects, every project staff member that has a significant financial interest or potential conflict of interest shall submit a Disclosure Form.
(7) ☐ Investigator’s Drug Brochure for pharmacy review (if applicable).
(8) ☐ Review by Privacy Board and IRB Approval/Determination* (send IRB approval when available).

* If the IRB submission is approved before review and approval of the UH Research Application, the IRB may not release the approved Informed Consent and send final IRB approval until this Application has received administrative approval at UH.

Please obtain the signatures of the appropriate individuals (items #14 – #16 on the next page) before submitting this Application Form.

13. As the Principal Investigator or UH employee/workforce member signed below, we/I certify that we/I have reviewed the applicable policies and requirements:

(1) The Belmont Report: Ethical Principles and Guidelines for the Protection of Human Subjects of Research,
(2) U.S. DHHS regulations for the Protection of Human Subjects at 45 CFR Part 46,
(3) US Food & Drug Administration (FDA) Regulations located throughout 21 CFR, and
(4) The relevant UH institutional policies and procedures for the protection of human research subjects, clinical trials, research privacy, and research integrity.

See CenterPoint (UH policies page) and https://www.trumed.org/professional-education/research-administration/

<table>
<thead>
<tr>
<th>Principal Investigator (Signature)</th>
<th>Date</th>
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</thead>
<tbody>
<tr>
<td>Co-Principal Investigator (Signature)</td>
<td>Date</td>
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<tr>
<td>(UH Medical Staff Member or Employee if PI is not)</td>
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<td>Inter Office Mail Address</td>
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Research Administration (816) 235-6015 or HSDResearch@umkc.edu
### Laboratory Approval

14. Any lab tests/procedures (central or local), including Point-of-Care testing, to be performed for this study project?

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<th>Yes</th>
<th>No</th>
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Please indicate the **local** lab tests requested to be performed AND any point-of-care tests (i.e., urine pregnancy test, blood glucose, etc.) to be performed by study staff.

### Pharmacy Approval

15. Any use of a drug (approved or investigational) or drug-eluting device?

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If **Yes**, list the investigational product:

Comments

### Department Approval

16. 

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<tr>
<th>Department Manager</th>
<th>Date</th>
<th>Department Chair</th>
<th>Date</th>
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(Program Director if resident’s study)

Comments

### Administration Approval

17. Financial and Administrative Review

| Estimated Revenue $_____ |

Funds will be administered by:  

| UH | UMKC | Other_____ |

Comments _____

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<tr>
<th>Director of Research Administration</th>
<th>Date</th>
<th>UH Legal Counsel</th>
<th>Date</th>
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(As to Legal Form and Insurance)  
(Not applicable for non-sponsored research studies)