

Does this proposal involve the use of significant IT resources (beyond basic academic infrastructure); the generation of datasets or digital assets; or a budget with over \$10,000 in IT-related hardware, software, or staff expenditures? (Check [additional requirements](#))

Human Subjects? If yes, indicate "Pending", IRB # or Exemption #: Delayed Onset

NIH-funded Clinical Trial? If yes, investigators and staff involved in the conduct, oversight, or management of clinical trials should be trained in Good Clinical Practice. Training is available through [CITI Program](#). Provide names on the next page.

Will the clinical research study utilize UCLA Health System resources, including but not limited to, any patient care costs? If yes, then a Policy 915 Coverage Analysis is required (contact coverageanalysis@mednet.ucla.edu).

Animal Subjects? If yes, indicate "Pending" or ARC#: Delayed Onset

Human Embryonic Stem Cell Research? If yes, refer to the [Stem Cell Policy and Procedures](#).

Non-UCLA materials/equipment to be used? If yes, indicate type: Source:

Human or primate cells, tissue, or fluids; recombinant or synthetic nucleic acids; potentially infectious materials; exotic plants or plant pathogens; select agents or toxins? For more information, see [IBC website](#).

Use of UC IP? If yes, specify case number:

Yes No Export Control (see [RPC Website](#)) – Does the project involve the following:

Shipping or carrying any tangible object or item to a foreign country?
If yes, specify:

Conducting research or other activities in, taking money to or planning to have money transferred to a foreign country?
If yes, specify:

Training foreign persons in using equipment, technology, or technical data?
If yes, specify:

Traveling to or doing research in a country currently under a US Trade or Economic Embargo (See [OFAC Website](#))?
If yes, specify:

7. Additional Forms Required

Yes No COI ([Disclosure Requirements](#))

Sponsor/Prime Sponsor is Federal [Public Health Service \(PHS\)](#) or agency that has adopted the PHS regulations? If yes, provide names of other investigators on [page 3](#) (See [UCLA Policy 926](#)).

Sponsor/Prime Sponsor is Federal (other than PHS), CIRM or special research programs managed by the UC Research Grants Program Office (RGPO)? If yes, attach COI [Form 740](#) & [Supplement to Form 740](#) (if applicable). See [UCLA Procedure 925.3](#).

Non-Government Sponsor/Prime Sponsor? If yes and project is *Research*, attach [Form 700-U](#), [700-U Addendum](#) and [700-U Supplement](#), as applicable, unless sponsor is [exempt](#). See [UCLA Procedure 925.2](#)

Yes No Industry Sponsored Research

Industry Sponsored Non-Clinical Proposal? If yes, attach [Industry Sponsored Research Checklist](#).

Industry Sponsored Clinical Trial? If yes, view the [Clinical Trials Contracts & Strategic Relations Checklist](#) to determine additional required attachments.

8. Funds Requested

1st Budget Period

Direct Costs (\$): Excluded Direct Costs (\$): F&A Costs (\$): Total Costs (\$):

All Project Periods (*complete only when multiple budget periods are involved*)

Direct Costs (\$): Excluded Direct Costs (\$): F&A Costs (\$): Total Costs (\$):

F&A: F&A Rate (%): F&A Base Type: If Other, specify:

9. Remarks

10. Accepts Responsibility

Approvals: Includes Certifications

The Investigator(s) certifies to the following: (1) that the information submitted within this application is true, complete and accurate to the best of their knowledge; (2) that any false, fictitious, or fraudulent statements or claims may subject the Investigator(s) to criminal, civil or administrative penalties; (3) agrees to accept responsibility for the scientific conduct of the project and to provide the required progress reports if a grant is awarded as a result of the application; and (4) that you are not currently debarred, suspended or ineligible to receive federal or non-federal funds; (5) all Clinical Trials based upon [FDAAA 801](#), will be registered in [ClinicalTrials.gov](#). When multiple Investigators are proposed in an application this assurance must be obtained by all named Investigators.

Principal Investigator (Required) Date

Date

Date

Chair/ORU Director/Dean/Medical Center Director (Required) Date

Date

Date

