

University of California, Davis
Health System, Sacramento
Department of Pathology and Laboratory Medicine

CLINICAL AND QUALITY RESEARCH (CQR)

CQR 001

Please allow at least 10 days for initial review. The Clinical and Quality Research Team will follow-up if there are any additional questions or concerns. **Please only use the Hs-Pathresearch@ucdavis.edu to submit this form and for all communications related to this study. Communication outside of this process may significantly delay review of your study.**

A. General Project Information

Study Title: _____

Principal Investigator: _____

IRB Number: _____ IRB Expiration Date: _____

Please attach approved protocol and IRB approval letter. We understand that IRB number and approval may not be available when this form is first submitted but please circle back and provide this information when it becomes available.

Anticipated start date of study: _____

Study duration (years, months, weeks): _____

How many patients will be enrolled at UC Davis? _____

How many patients will be active at any given time (approximately)? _____

B. What Pathology resources are needed? (please check all applicable)

Blood bank

Apheresis

Clinical Testing (if checked, see Part C)

Blood draw: *Frequency per patient* _____

Location: _____

Other (*please describe* _____)

C. Clinical Testing performed at UCDMC*

Please list requested labs: _____

On a **per patient** basis, please describe:

Total number of specimens: _____

Testing Frequency (*i.e. 5 samples per week*): _____

Will any of these tests be used for patient care? yes no

If yes, which labs? _____

**If multiple testing is requested with varying frequencies, please list all tests with the respected frequency. A separate sheet can be added if more room is needed.*

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Was laboratory safety training completed for all involved personnel (if applicable)?

- YES
- NO
- NA

Primary Study Contact

Name: _____

Phone number: _____ E-mail: _____

Study Emergency Contact

Name: _____ Phone number: _____

Electronic signatures accepted.

Study Principal Investigator: _____
Signature _____ Date _____

Study Coordinator (if available): _____
Signature _____ Date _____

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Please direct any questions or concerns to Hs-Pathresearch@ucdavis.edu or 734-2112.

FOR CLINICAL AND QUALITY RESEARCH USE ONLY

- Valid IRB Protocol and Decision Letter on file with the Clinical and Quality Research Team
- Type/Frequency of review determined by Clinical Research Oversight Committee
Type/Frequency: _____

Approval Signatures

FOR CLINICAL AND QUALITY RESEARCH USE ONLY

Clinical Quality Research Assistant: _____
Signature _____ Date _____

Laboratory Section Medical Director: _____
Signature _____ Date _____

Quality Assurance and Operations Manager: _____
(for full review studies only) Signature _____ Date _____