Dear Members of the UCLA Research Community:

With the June 2, 2017 update from the Vice Chancellors for Research and Health Sciences about clinical trial registration requirements, the Office of the Human Research Protection Program (OHRPP) has retired the document “Guidelines for Registering in the ClinicalTrials.gov Registry” from its website.

The OHRPP guidelines have been retired in order to streamline sources of information for researchers. Please note the following:

- For guidance, refer to the websites of the Clinical and Translational Science Institute (CTSI) and the Office of Research Policy and Compliance.
- Regulations still require specific consent form language about clinicaltrials.gov. This language remains in the consent templates provided on OHRPP’s website.
- A question about registration remains in the webIRB application. Your answer does not factor into the IRB review but helps the University provide support.

For assistance with reporting, please e-mail the appropriate contact:

- For cancer studies, the Jonsson Comprehensive Cancer Center Office of Regulatory Compliance
- For non-cancer studies, the CTSI Office of Regulatory Affairs

Sincerely,
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