


NCI CTRP Registration User's Guide v4.4



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
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CTRP Registration Links

- [NCI Clinical Trials Reporting Program Registration](#)
- [Templates for batch accrual submissions, and other resources](#)

CTRP Resources

- [CTRP Website](#)
- [Dictionary of cancer terms](#)
- [Terminology resources](#)



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Application Support

If you have problems with the program or have suggestions for any of the CTRP User's Guides, contact the NCI Clinical Trials Reporting Office using the information and guidelines provided in [Application Support](#).

About this Guide

This guide provides an overview of NCI Clinical Trials Reporting Program (CTRP) Registration and instructions for using its tools and resources to submit new clinical trials and amend and/or update those currently registered and verified in the CTRP. Additionally it contains instructions for searching for and viewing details of existing registered clinical trials.

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Audience

This guide is designed for members of the NCI clinical research community, who, in their roles as submitters and/or principal investigators, register details about clinical trials for use by the broader scientific community. Separate instructions are provided for Site Administrators.