**Course Content**

**GenBank Database: Description and Scope**

**The NCBI Derivative Databases: RefSeqs**

**Database Searching Using Entrez**

**The NCBI Structures Database**
- The Molecular Modeling Database
- Viewing Structures and Structural Alignments with Cn3D

**Similarity Searching Using NCBI BLAST**
- Local Alignment Statistics
- Scoring Systems
- Using WWW BLAST
- PSI-BLAST / PHI-BLAST
- RPS-BLAST (CDD Search)
- Specialized BLAST pages

**Genomic Resources at NCBI**
- Complete Genomes in Entrez
- Higher Genome Resources
  - RefSeq and LocusLink
  - UniGene
  - Variation Data (SNPs)
- The NCBI Draft Genome
- The Map Viewer
- Mouse and Other Genomes

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**A Field Guide to GenBank and NCBI Molecular Biology Resources**

Molecular sequence databases are an important part of nearly all areas of biological and biomedical research. To assist life scientists in using these data and tools, the National Center for Biotechnology Information (NCBI) at the National Library of Medicine offers, at your institution, a one- to two-day training course entitled, “A Field Guide to GenBank and NCBI Molecular Biology Resources.” An overview and list of scheduled locations for the training course is at: [www.ncbi.nlm.nih.gov/Class/FieldGuide/](http://www.ncbi.nlm.nih.gov/Class/FieldGuide/)

This training course is provided free of charge, and the NCBI will pay all travel expenses of the instructors.

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**Course Description**

The course consists of a 3-hour morning lecture followed by a 2-hour instructor-led computer workshop in the afternoon. During the afternoon session, students will get hands-on experience with the resources at the NCBI site and will learn how to apply these resources in their specific research area. The course provides practical information about GenBank, RefSeq and the integrated genome resources, as well as the other databases maintained at the NCBI. It also provides instruction on effective use of Entrez and BLAST, the two main database search tools.

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**Intended Audience**

The course is designed for principal investigators, postdoctoral fellows, graduate and advanced undergraduate students and others in the life sciences who work with biological sequence data. Both experienced and novice users of NCBI’s tools and resources will benefit.

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**Requirements for Hosting the Course**

- A pre-registered audience of at least 50 people
- An auditorium or similar setting with computer projection capabilities
- A multi-workstation computer classroom with internet access and standard web browsers for each machine
- An instructor’s workstation with projection capabilities

**Note:** Multiple hands-on sessions can be offered depending on the number of workstations available and the number of people registered for the course. If necessary, additional sessions can be added the following day.

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**Instructors**

Peter S. Cooper, Susan Dombrowski, Wayne Matten, Rana Morris, and Eric Sayers are bioinformatics training specialists with the NCBI User Services staff.

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