

Metabolomics Resources

Metabolomics Resources:

1. Metabolomics Databases

1. LC-MS Databases (Liquid Chromatography-Mass Spectrometry): LC-MS research data for download can be accessed done through various repositories and databases where researchers often deposit their datasets for public access. Some prominent sources include:
 1. [Metabolomics Workbench](#): Provides a vast collection of metabolomics data, including LC-MS datasets, along with associated metadata and experimental details.
 2. [MetaboLights](#): A repository for metabolomics experiments, it houses a wide range of datasets, including LC-MS data, and offers tools for visualization and analysis.
 3. [MassIVE](#) (Mass Spectrometry Interactive Virtual Environment): Focuses on MS data, offering access to high-quality datasets including LC-MS, proteomics, and other omics datasets.
 4. [GNPS](#) (Global Natural Products Social Molecular Networking): Particularly useful for natural product research, it hosts LC-MS datasets and enables molecular networking to explore chemical diversity.
 5. [PRIDE Archive](#): While primarily for proteomics data, it also includes some metabolomics datasets, including LC-MS data.
 6. [European Bioinformatics Institute \(EBI\) Metabolomics](#): Offers access to metabolomics datasets, including LC-MS data, along with analysis tools and resources.
 7. [DataCite](#): An open repository that aggregates datasets from various disciplines. You can search for specific LC-MS datasets within its collection.
 8. [BMRB \(Biological Magnetic Resonance Bank\)](#): A repository for NMR spectroscopy data related to biomolecules.
 9. [MetaboLights](#): A repository for metabolomics data, including LC-MS datasets related to metabolomics research.
 10. [Human Metabolome Database \(HMDB\)](#): Contains metabolomic data, including NMR spectra, for human tissues and biofluids.
 11. [SpectraBank](#): A repository for various spectroscopic data, including NMR

spectra.

12. [NMRShiftDB](#): A database specifically focused on organic structures and their NMR spectra.
13. [CCPN \(Collaborative Computational Project for NMR\)](#): Offers tools, software, and access to NMR data.
14. [Chemical Shift Repository \(CSR\)](#): Provides a platform for storing and sharing chemical shift data.
15. [Cambridge Structural Database \(CSD\)](#): Contains a vast collection of small molecule crystal structures, some of which may include associated NMR data.
16. [Spectral Database for Organic Compounds \(SDBS\)](#): Contains NMR, IR, MS, and Raman spectra for various organic compounds.
17. [ACD/NMR Predictors](#): Offers software for NMR prediction and includes access to a database of predicted spectra.

2. NMR Databases

1. [BMRB](#) (Biological Magnetic Resonance Bank): A repository for NMR spectroscopy data related to biomolecules.
2. [MetaboLights](#): While primarily for metabolomics data, it also hosts NMR datasets related to metabolomics research.
3. [HMDB](#) (Human Metabolome Database): Contains metabolomic data, including NMR spectra, for human tissues and biofluids.
4. [SpectraBank](#): A repository for various spectroscopic data, including NMR spectra.
5. [NMRShiftDB](#): A database specifically focused on organic structures and their NMR spectra.
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1. Metabolomics Books

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