

## Metabolomics Resources in the Biological Magnetic Resonance Data Bank (BMRB)



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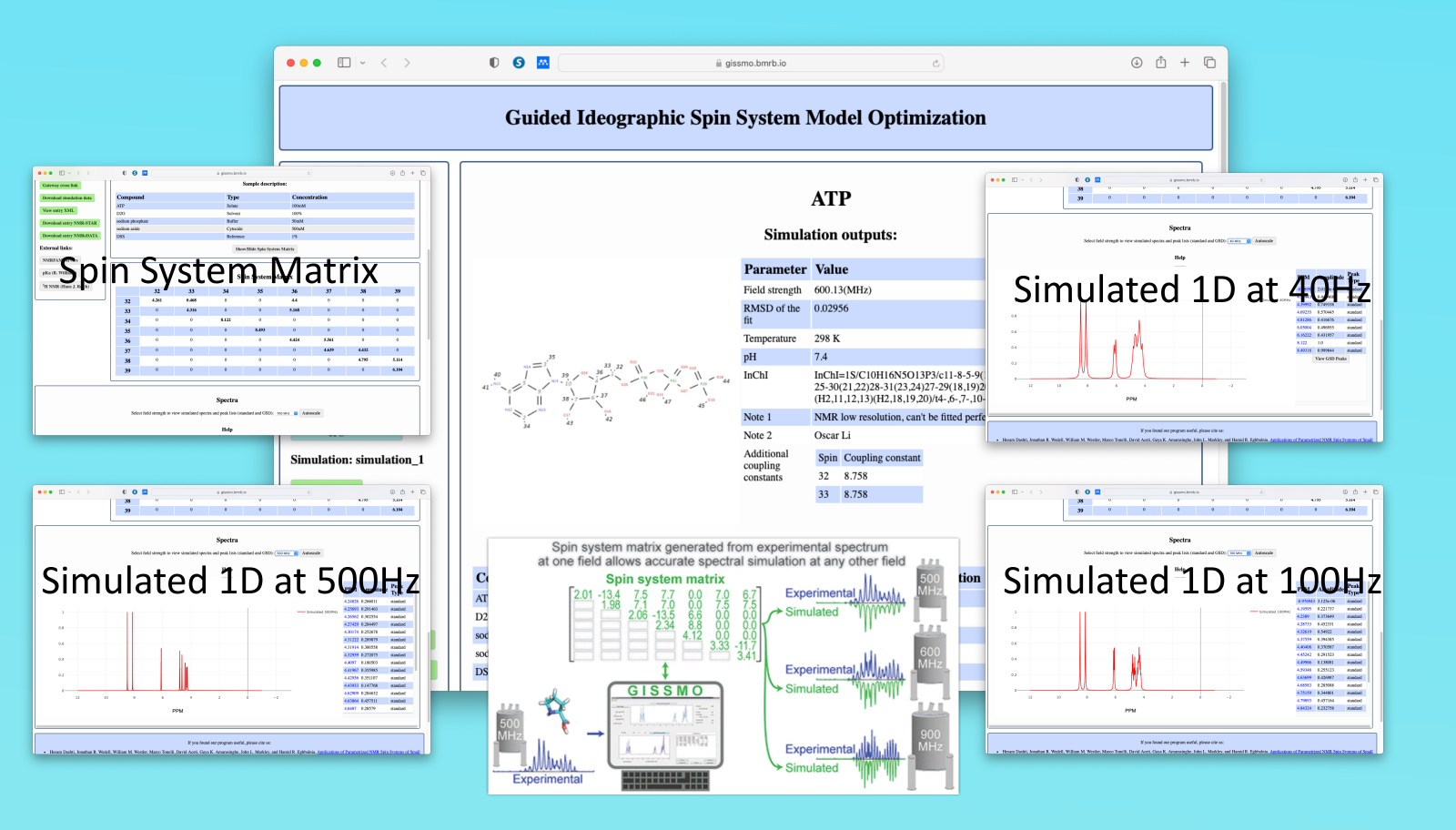
## BMRB small molecules library (https://bmrb.io/metabolomics/)

BMRB<sup>1</sup> provides curated set of reference NMR spectroscopic data for more than 1000 metabolites measured using a standardized experimental protocol

Each reference data set consists of

- Chemical shift assignments
- Sufficient time domain data (FID) from a set of 1D and 2D experiments to analyze the molecule

This library could be used to analyze and identify molecules in bio-fluids and other metabolite mixtures. Peak lists of 1D and 2D experiments can be searched across the database through web interface or using BMRB-API



## **ALATIS Library** (https://alatis.bmrb.io/)

Atom Label Assignment Tool Using InChl String<sup>3</sup> (ALATIS) creates unique InChl identifiers for small molecules through rigorous labeling of their atoms. This numbering system helps to identify molecules across different databases and flawlessly compare the properties of their individual atoms

### ALATIS library includes

YYY NMRbox Software

Software research\_problem= Metabolomics

11 software packages meet the criteria.

Metabolomics toolbox

It's simple! Let's help you get started.

You can search by Research Problem, Software Type, or All Software.

A top class software suite to process your

Tools for processing and analyzing

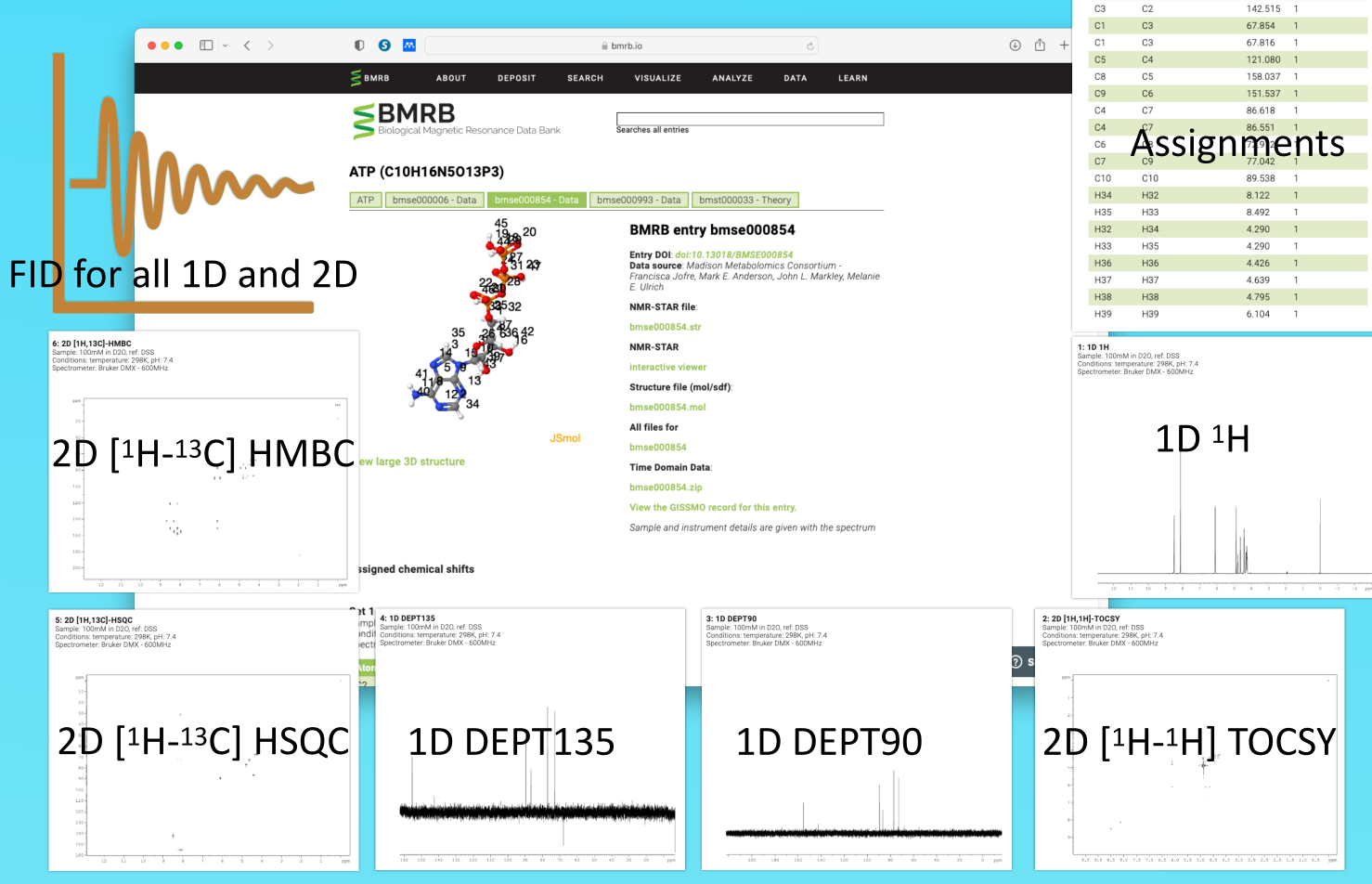
 Unique InChI strings for molecules from various databases like BMRB, PubChem, HMDB and RCSB-PDB Ligand-Exp

or perform a text search

An R package for comprehensive analysis o

Cross-links from PDB entries to BMRB, HMDB and PubChem

BMRB follows ALATIS numbering system in its small molecules database.



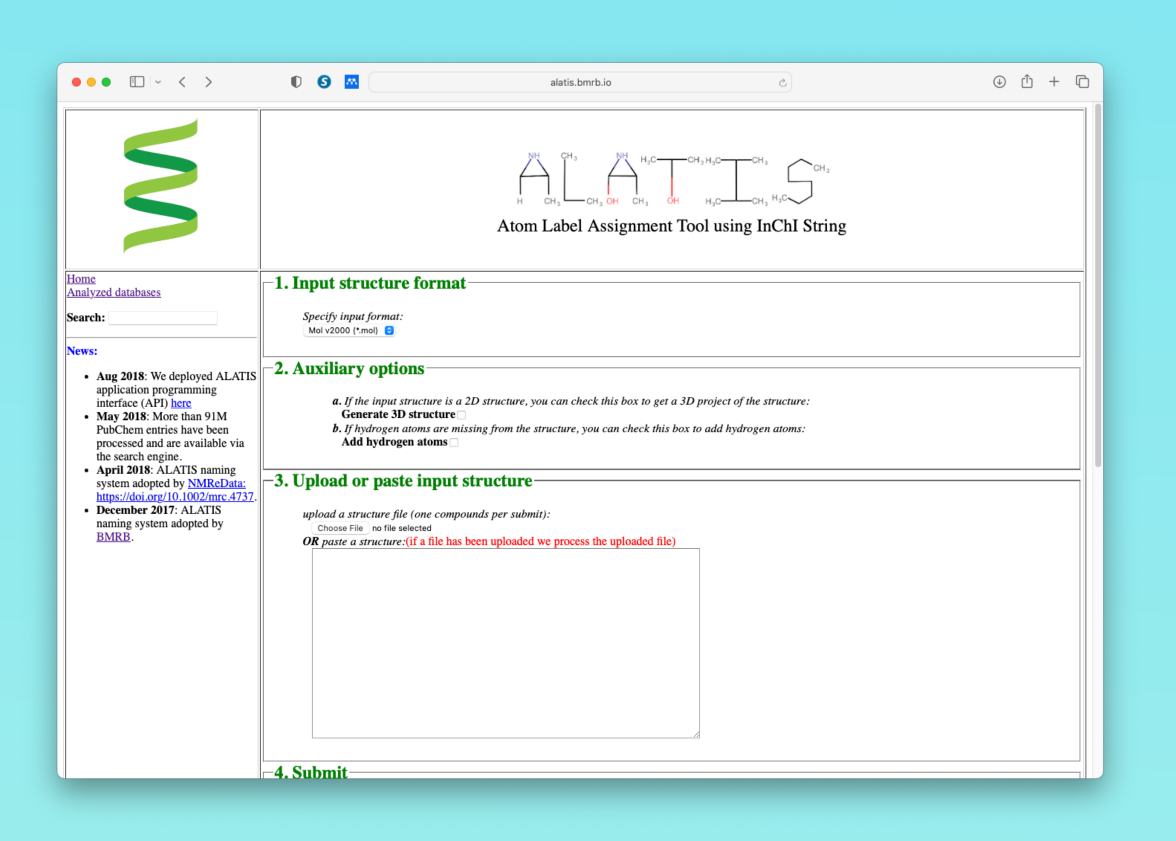
# **GISSMO Library**

(https://gissmo.bmrb.io/)

Guided Ideographic Spin System Model Optimization<sup>2</sup>(GISSMO) enables the efficient calculation and refinement of spin system matrices( chemical shift and coupling constants against experimental 1D-1H spectra of small molecules.

#### GISSMO library includes

- Library of over 1000 parameterized small molecules
- Spin system matrix for every molecule
- Simulated 1D spectra at various field strength (40Hz to 1.3GHz) and peak lists at each field strength



### NMRbox: Software and Computing resource (https://nmrbox.nmrhub.org/)

NMRbox<sup>4</sup> is a resource for biomolecular NMR software. It provides tools for finding the software you need, documentation and tutorials for getting the most out of the software, and cloud-based virtual machines for executing the software. NMRbox is available free for academic use.

NMRbox provides ready to use computing resource along with various preinstalled software tools like ALATIS, BATMAN, dataChord Spectrum Miner, GISSMO, MestReNova(Nova). MetaboAnalystR, Metabolomcis toolbox, MVPACK, MZmine3, RUNNER, tameNMR and MATLAB.

NMRHhub is an umbrella organization which grants access to data collection infrastructure (Network of Advanced NMR: NAN), analysis platform (NMRbox), data resource (BMRB) and Non-uniform sampling and reconstruction project (NUScon) through single portal.

PDF version of this poster



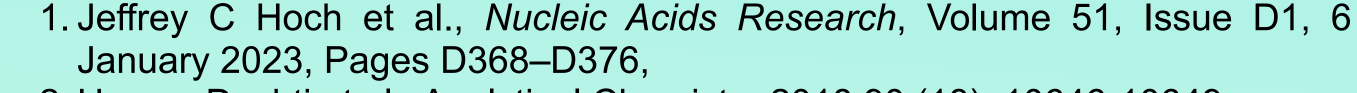












- 2. Hesam Dashti et al., Analytical Chemistry 2018 90 (18), 10646-10649
- 3. Hesam Dashti et al., Sci Data 4, 170073 (2017)
- 4. Jeffrey C Hoch et al.., Biophysical Journal, 112: 1529-1534, 2017.