



www.chenomx.com

Metabolite Discovery and Measurement

**SOFTWARE LICENSING OR SERVICES...
THE CHOICE IS YOURS**

Software Licensing

EXTENSIVE REFERENCE LIBRARY

INTUITIVE USER INTERFACE

SUPPORTS MOST NMR DATA FORMATS

QUANTITATIVE RESULTS



REFERENCE LIBRARY

The Chenomx Reference Libraries contain hundreds of fully searchable pH-sensitive compound models. These models are automatically calibrated by adjusting compound line shapes, peak widths, and chemical shifts to better match sample conditions. The libraries have been developed using a combination of mathematical algorithms and actual NMR measurements for suitability for quantitative metabolite measurement.

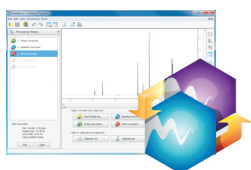
A list of the reference compounds can be referenced from the Chenomx Web site. New entries can be added by Chenomx on request or by users of the software.



PROFILER - Identify metabolites and measure their concentration

Chenomx Profiler is used to identify compounds and quantify their concentrations based on data in an NMR spectrum. Key features include:

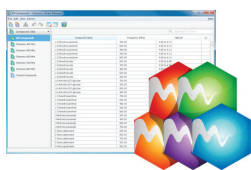
- Computer assisted metabolite assignment
- Spectral binning
- Spectral overlays
- Multiple file concentration exports
- Detailed reference resources for library compounds
- Automated fit of selected metabolites in batches of spectra



PROCESSOR - Process your NMR spectra

Chenomx Processor is used to calibrate a spectrum by specifying pH and CSI. This ensures the proper calibration of our spectral library when identifying and quantifying compounds with Profiler. Chenomx Processor converts various spectrum formats into the Chenomx file format, including Agilent, Bruker, JEOL, JCAMP-DX, Mestrelab, and NMRPipe. It also provides many tools to process your spectra including:

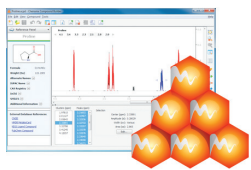
- Phase Correction
- Baseline Correction
- Shim Correction
- Region Deletion
- Automatically process multiple spectra in a batch



LIBRARY MANAGER - Organize and create compound sets

Chenomx Library Manager allows you to create and manage Compound Sets for use in Profiler. Compound Sets allow you to:

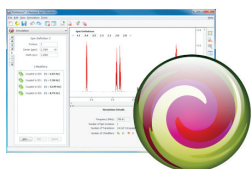
- Customize a list of compounds for a specific biofluid
- Filter and narrow your list of interesting compounds
- Manage your own custom compounds



COMPOUND BUILDER - Add your own compounds and metabolites

Chenomx Compound Builder lets you create your own powerful mathematical models of custom NMR compounds. It includes a number of tools to make this task easier, such as:

- Automatic cluster fit and optimization
- pH curve modeling
- For creating pH-sensitive compound signatures using equation based peak parameters



SPIN SIMULATOR - Generate theoretically accurate metabolite simulations

Chenomx Spin Simulator is a powerful tool for creating and manipulating simulations of compound spin systems. Use Chenomx Spin Simulator to:

- Determine the coupling relationships between nuclei
- Calculate j-coupling constants between nuclei
- See your simulation update in real time

CHENOMX RESEARCH SERVICES offers a cost-effective method for measuring concentrations of many small molecules found in different complex mixtures and biological matrices. Our research services keep repeatability and reproducibility as a priority, and can be used for both large and small studies. Results are fully quantitative, allowing data to be compared across long-term studies. We offer sample analysis, spectral profiling services and/or statistical support.

At the heart of Chenomx's technology platform is a Nuclear Magnetic Resonance (NMR) spectrometer. Chenomx's patented targeted profiling technique easily translates NMR spectra of complex mixtures and biofluids into compound information relevant to the underlying biology. Customers using our research services include some of the world's largest life sciences companies, as well as leading universities in the United States, Europe, and Asia.

Chenomx offers a broad range of services for researchers. Chenomx Research Services are customized for assisting researchers in various ways, including:

- **Study Design**
- **Hypothesis Generation**
- **Hypothesis Testing**
- **Biomarker Pattern Discovery**
- **Personalized Small Molecule Detection**

Applications

Chenomx research services produces a metabolic profile, which can be used to monitor the effects of disease, diet, toxins or pharmaceuticals on an organism. Results from Chenomx research services is particularly valuable in understanding effects on metabolic pathways since it is an inherently quantitative method and provides a global view of the metabolites.

Applications include:

- **Drug metabolism studies** – metabolic pathway analysis, pharmacokinetics
- **Biomarker discovery** – metabolic marker patterns for monitoring or diagnosing disease
- **Cell Metabolism** – cancer, Bio-production and other applications
- **Food and Nutrition** – both food content and effect of nutrition on organisms

Chenomx MetaProfile Results

Chenomx produces a full MetaProfile report describing the mixture analysis results from the complete study. This report breaks down results for each sample, describing small molecules identified in the sample, and describing concentrations of the small molecule found in each sample.

Data is provided to the client as an Excel spreadsheet, suitable for further analysis or transfer to other statistical and analysis systems.

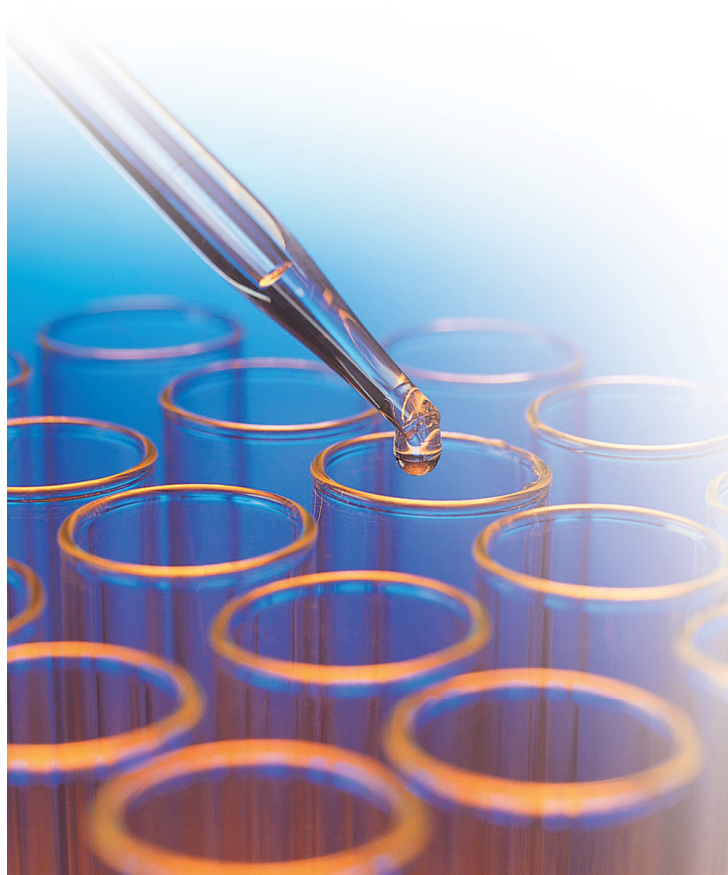
Biomarker Pattern Discovery

Discovery of biomarkers starts with a clear identification and quantification of the components in a mixture. This is offered by Chenomx's patented Targeted Profiling technique. Quantification using Targeted Profiling produces data that may be subjected to a variety of statistics and visualization methods. These methods are key to discovering useful biomarker information.

Diagnostic Model Development

Targeted profiling based on NMR allows you to rapidly develop diagnostic models that you can use directly on experimental data from the same platform. Chenomx gives you access to high-throughput NMR facilities that allow you to create the large data sets necessary to build robust diagnostic models.

Chenomx has years of experience in designing experiments, building diagnostic models, and validating them with clinical trial data. We can help you turn your model into a routine test.





About Us

CHENOMX is a life-sciences company located in Edmonton, Canada. Incorporated in 2000, Chenomx is one of the pioneers in the metabolomics research industry. The mission of the company is to help researchers to better understand biological systems through metabolomics. One of the key values Chenomx provides is a platform for the analysis of biological mixtures using NMR Spectroscopy. This platform includes standardized methods of sample preparation, NMR data acquisition, and Targeted Profiling of spectra produced. Chenomx's customers spend less effort on the development of the NMR methods. Chenomx chose NMR spectroscopy as the measurement technology since it offers a robust, reliable, reproducible method offering quantitative results.

Chenomx is focused on developing the best software tools possible for NMR based metabolomics. This is evident in efforts in developing a library of spectral models of hundreds of metabolites that are present in common research fluids such as urine, blood and cell extracts. Chenomx has meticulously enhanced this library with features that give the most accurate identification and quantification of metabolites. This technology is licensed for use by hundreds of life-science researchers world-wide.

Chenomx leverages the patented technology by offering sample analysis services to researchers world-wide. Physical samples can be sent to Chenomx for turn-key analysis with consulting services to better utilize the results. Spectra collected on client NMR spectrometers can be sent to Chenomx electronically, analyzed by Chenomx scientists and returned with reporting and optional consulting services. The strength of the platform technology reduces sample preparation issues; quantitative results mean that both small and large projects offer valuable results.

The Technology Platform

Chenomx's patented NMR-based metabolomics platform provides a powerful, global analysis of hundreds of metabolites in complex biological mixtures.

Metabolomics

Metabolic profiling, or metabolomics, is the study of patterns in the complement of low molecular weight compounds found in biological fluid samples. Small molecule byproducts of metabolic processes vary in concentration and relative proportion in response to disruptions in the metabolism of an organism, providing insight into the health status of the organism. Genetic abnormalities, enzyme inhibition or activation, pathological conditions, and drug actions can create metabolic disruptions detectable in the metabolite content of biofluids.

Nuclear Magnetic Resonance

Nuclear Magnetic Resonance (NMR) spectroscopy is a powerful tool with a wide range of applications in academia, industry, and medicine. Traditionally, these applications have formed three broad groups: analytical services, structural biology, and medical imaging. NMR has proven to be a versatile and productive technology for metabolomics.

Unlike other analysis platforms, NMR does not rely on separating mixtures before analysis. NMR can simultaneously detect hundreds of metabolites in biological samples, and has been referred to as a "universal detector". NMR's non-destructive, global detection make it a strong choice for any metabolomics study.

Targeted Profiling

Chenomx's patented Targeted Profiling technique addresses one of the more challenging aspects of NMR-based metabolomics: identifying and quantifying metabolites.

Targeted Profiling identifies compounds in a complex biofluid mixture rapidly and accurately. This direct analysis approach differs from previously published methods, which depend on binned spectral areas. Targeted profiling relies instead on libraries of compound signatures modeled to behave like the pure spectra of the individual compounds under comparable experimental conditions. The underlying spectral libraries are collected at a variety of pH conditions to account for variations among sets of samples. Analysis of samples using target profiling directly yields both the identity and concentration of individual compounds in a single operation.

What You'll Need for the Software

- Chenomx runs on multiple platforms including: Windows 8/7/Vista/XP, Mac OS X 10.9-10.7.3, Linux
- Data Acquisition by:
 - 400 through 800 MHz NMR spectrometers manufactured by Agilent, Bruker, JEOL and others
 - Chemical shift indicator solution of a certified concentration is available from Chenomx.



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