Data and text mining

mspire: Mass spectrometry proteomics in Ruby

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ABSTRACT

Summary: Mass spectrometry based proteomics stands to gain from additional analysis of its data, but its large, complex data sets make demands on speed and memory usage requiring special consideration from scripting languages. The software library 'mspire'—developed in the Ruby programming language—offers quick and memory-efficient readers for standard XML proteomics formats, converters for intermediate file types in typical proteomics spectral-identification work flows (including the Bioworks .srfl format), and modules for the calculation of peptide false identification rates.

Availability: Freely available at http://mspire.rubyforge.org

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Supplementary information: Additional data models, usage information, and methods available at http://bioinformatics.icmb.utexas.edu/mspire

1 INTRODUCTION

The analysis of mass spectrometry (MS) proteomics data is challenging on many fronts. Data sets are complex, with information spanning multi-level hierarchies, and they are also very large—files are often of near gigabyte size. Access to MS proteomics data is increasing with the advent of standardized formats such as mzXML and repositories such as PeptideAtlas (Desiere et al., 2006), but its analysis remains no less daunting. Strongly typed languages (e.g., C/C++ and Java) are well suited for intensive computational tasks, but less so for exploring landscapes of computational possibilities. Scripting languages (e.g., Python, Perl and Ruby) are ideal for quick prototyping and the exploration of new ideas but can be too slow or memory inefficient for large data sets. Thus, a need exists for scripting language tools capable of dealing with the size and complexity of MS proteomics data.

Ruby is a full-featured programming language created with inspiration from Perl, Python, Smalltalk, and Lisp. It is object oriented and remarkably consistent in its design. Ruby's syntax encourages the use of blocks and closures which lend flexibility and conciseness to programming style. Also, while it is powerful, Ruby is relatively easy to learn, making it a natural first programming language for budding bioinformaticians. Ruby does not have the same degree of support for scientific computation as Python (e.g., NumPy and PyLab), but it is building significant momentum in this area (e.g., SciRuby at http://sciruby.codeforpeople.com). These features encouraged our use of Ruby in the creation of a high-level library supporting MS proteomics analysis.

A few libraries/tools exist for working with MS proteomics data outside of Ruby. InSilicoSpectro, the only other scripting language library, is an open-source library written in Perl for “implementing recurrent computations that are necessary for proteomics data analysis”. While there is some overlap with the work described here (e.g., in silico protein digestion), that library is currently geared towards the support of the Phenyx and Mascot search engines and low-level spectral computation (Colinge et al., 2006) while mspire is geared towards supporting Thermo's Bioworks software (SEQUEST) and downstream analysis such as false identification rate determination. The ProteomeCommons.org IO framework also has the ability to read/write and convert common data formats (Falkner et al., 2007), but this library is written in Java and does not provide any higher-level language tools.

2 FEATURES

Mspire is a software package for working with mass spectrometry proteomics data as outlined in Fig. 1A.

2.1 Memory Usage and Speed

Mspire relies on several memory-saving techniques that are critical for working with large data files. Large quantities of objects are implemented as Arrayclass (http://arrayclass.rubyforge.org) objects, providing highly efficient memory usage (Fig. 1B) while preserving accessor behavior common to typical Ruby objects.

By default, spectra from mass spectrometry file formats (mzXML and mzData) are decoded into memory-efficient strings and are only completely cast when spectral information is accessed. An option is also available for storing only byte indices of spectral information that can be used for fast, random access of spectra or for reading files of essentially unlimited size.

REXML, Ruby’s standard library XML parser, can be far too slow when reading large XML files generated in MS proteomics. Mspire can use either XMLParser or LibXML (both of which have C/C++ bindings) for rapid parsing of large files.

Performance reading and then accessing two spectra across thousands of mzXML files from the PeptideAtlas is shown in Fig. 1C. Late evaluation of a spectrum allows files to be read at ~ 20 MB/sec with no file-size limit.

2.2 Reading MS proteomics data formats

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2.3 Bioworks SEQUEST Results Files (.srf)

Bioworks previously produced separate text files for each spectrum but now outputs a single SEQUEST results file (.srf) for each set of searches. This increases the speed of a search, decreases disk space usage, and is much easier to work with in file system operations. Unfortunately, because the output is binary, accessing its contents can be difficult and downstream analysis tools (outside of Bioworks) do not currently support this format.

We created a reader for .srf files using the Ruby “unpack” function. It extracts both spectral information and SEQUEST results. The reader is fast and also works across platforms because it does not rely on any vendor software libraries.

2.4 Reading/writing spectral identification formats

Even when derived from the same upstream data source, formats for working with spectra identifications can vary widely. We designed readers and writers for common downstream spectral-identification software formats for SEQUEST-based data: pepXML files which are used in the Trans-Proteomic Pipeline (Protein Prophet) and also the .sqt format, which can be used with DTSelect and Percolator (Kall et al., 2007).

Readers are tailored to their respective format so that users can not only extract format-specific information easily but also implement a common interface so that users can easily extract information shared across these formats.

2.5 Determining false identification rates

Bioworks software support for determining false identification rates (FIRs) is currently nonexistent, and so downstream tools are necessary. Mspire supports peptide FIR determination from target-decoy DB searches (both the creation of decoy databases and the summary of search results), PeptideProphet and Percolator. Known biases in sample content can also be used to establish an FIR.

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REFERENCES


