# Specter: linear deconvolution as a new paradigm for targeted analysis of data-independent acquisition mass spectrometry

# User Guide for Standalone (non-Spark) Mode

Ryan Peckner

April 26, 2018

Specter is an algorithm for the targeted analysis of data-independent acquisition mass spectrometry experiments. It can analyze data from any instrument type and window acquisition scheme. The required user inputs are a DIA data file in centroided mzML format, a spectral library in blib format, and a mass accuracy parameter, specified in parts-per-million.

The raw output of Specter is a .csv file describing the total ion intensities (= sum of fragment ion intensities) of each precursor in the spectral library at each retention time point of the experiment. A snippet of the typical 'SpecterCoeffs' raw output file looks like this:

Scan index	Retention time (s)	Precursor sequence	Precursor charge	Total ion intensity
10032	268.3763	ETLDASLPSDYLK	2	1,569,034
10032	268.3763	NPAADAGSNNASKK	2	3,112,580
10033	268.4273	IVLVDDSIVR	2	722,175

while the 'SpecterQuants' file contains the identifications and quantifications based on these scan-by-scan total ion intensities:

Precursor sequence	Precursor	charge	Quant
ETLDASLPSDYLK		2	148,110,338
NPAADAGSNNASKK		2	32,234,856
IVLVDDSIVR		2	11,768,772

## System requirements

Specter requires Python version  $\geq$  2.7.9 (Specter hasn't been tested with Python 3) and R version  $\geq$  3.2.3, with the packages 'kza',' pracma' and 'moments' installed.

# Setup

Download the contents of the GitHub repository to a location accessible to the machine you'll be using to run Specter. From the command line, cd to this directory. Now create a conda environment called "SpecterEnv" with the commands

- \$ conda env create -f SpecterEnv.yml
- \$ source activate SpecterEnv

Due to a bug in the package pymzml, which Specter uses to parse the mzML mass spec data files, you will need to download the most recent MS bioontology file from http://data.bioontology.org/ontologies/MS/submissions/116/download?apikey=8b5b7825-538d-40e0-9e9e-5ab9274a9aeb, rename it as "psi-ms-4.0.1.obo", and manually move it to the pymzml obo repository located in /usr/anaconda/envs/SpecterEnv/lib/python2.7/site-packages/pymzml/obo (this path may need to be altered depending on where your Anaconda install lives). On a Linux machine, this can be accomplished from the command line via

- \$ wget http://data.bioontology.org/ontologies/MS/submissions/116/download?apikey=8b5b7825-538
  d-40e0-9e9e-5ab9274a9aeb
- \$ mv download?apikey=8b5b7825-538d-40e0-9e9e-5ab9274a9aeb /usr/anaconda/envs/SpecterEnv/lib/
   python2.7/site-packages/pymzml/obo/psi-ms-4.0.1.obo

This issue has been flagged with the pymzml developers but has yet to be resolved.

#### Mass spectrometry data and spectral libraries

All mass spec data has to be provided in centroided mzML format. ProteoWizard's msconvert application is recommended to convert raw mass spec data to mzML and perform centroiding, including for AB SCIEX TripleTOF data in .wiff format. Spectral libraries must be provided in .blib format. Skyline can create a .blib spectral library from most DDA search engine output formats; for a tutorial on how to do this from a pep.xml file, see http://dia-swath-course.ethz.ch/tutorials2014/Tutorial-3\_Library.pdf. The list of supported search engine output formats for conversion to blib by Skyline can be found at https://skyline.ms/wiki/home/software/BiblioSpec/page.view?name=BlibBuild.

**Warning:** When building a spectral library from phosphoproteomics data, make sure to filter your search engine output so that only unambiguously localized phosphosites are included (using a localization score such as the A-score, PTM score, or variable modification location score from Spectrum Mill). This is necessary to avoid conflation of spectra from positional isomers.

## Running a Specter job

Once the SpecterEnv conda environment has been activated, the syntax for running Specter is

where the bracketed arguments are as follows:

- mzMLname: The full path to the mzML file containing the DIA data to be analyzed, without the mzML extension.
- blibName: The name of the blib file containing the spectral library, without the blib extension.
- index: The first or last index of the subset of MS2 spectra to be analyzed (see the next argument).
- StartOrEnd: Should index be interpreted as the first (StartOrEnd = "start") or last (StartOrEnd = "end") index of the spectra to be analyzed? This is useful for breaking jobs into smaller pieces to respect cluster memory constraints. Defaults to "end".
- numProcessors: The number of CPUs Specter will use to parallelize the computations over MS2 spectra. Defaults to the total number of available CPUs.
- instrumentType: This can be one of 'orbitrap','tof', or 'other'. Use of this argument in the first two cases helps avoid certain known issues with mzMLs coming from data converted from these instrument types. Defaults to "orbitrap".
- tol: The instrument mass accuracy, in parts-per-million. Defaults to 10.

For example, the command

would tell Specter to analyze the first 100,000 MS2 spectra in the Orbitrap DIA experiment file PhosphoDIARun1.mzML located in /rpeckner/data/20170501 using the spectral library HumanPhosphoLib.blib located in /rpeckner/libs/. The remaining parameters are the mass tolerance of 20 p.p.m., and 8 CPUs for parallel processing of MS2 spectra. Note that all values except

Specter will create a subdirectory 'SpecterResults' of the directory containing the mzML, into which the output files will be written. These are the following csv files with the common prefix '<mzMLname>\_<bli>blibName>':

- header: The header of the mzML file, containing the scan indices, retention times, and precursor m/z window centers.
- SpecterCoeffs: The raw Specter coefficients for every library precursor and MS/MS scan.
- SpecterCoeffsDecoys: The raw Specter coefficients for both library and decoy precursors and all MS/MS scans.
- SpecterQuants: The final identifications and quantifications of library precursors filtered to a 1% false discovery rate.

You can visualize the calculated chromatograms of individual library precursors in R using ggplot: