## Skyline DDA Search for MS1 Filtering

The Skyline Targeted Mass Spectrometry Environment provides informative visual displays of the raw mass spectrometer data you import into your Skyline documents. These displays allow you to manipulate the data by performing tasks such as refining the peptides and transitions you are measuring, and tuning integration boundaries. Originally developed to analyze quantitative assays from selected reaction monitoring (SRM – also referred to as multiple reaction monitoring or MRM) mass spectrometry, Skyline has been expanded to extract time-intensity chromatograms from MS1 spectra for use in peptide quantification experiments involving mass spec runs with data dependent MS/MS.

The Skyline MS1 Full-Scan Filtering supports importing data sets from discovery-type proteomics experiments where mass spectrometers were operated in data dependent acquisition (DDA) mode. After importing the raw data, new and previously existing Skyline features facilitate quantifying peptide precursor MS1 signal across many replicate acquisitions. This mode may also be used for visualizing and better understanding quantitative outputs from other "Label Free" quantitative tools, because of the exceptional data visualization plots in Skyline.

This tutorial will cover the following areas critical to making effective use of Skyline MS1 Filtering when you want Skyline to perform peptide spectrum matching for the MS/MS spectra in your DDA data:

- Setting up a Skyline document for MS1 filtering
- Performing a DDA search on raw data to find potential quantitative targets

Skyline aims to provide a vendor-neutral platform for targeted mass spectrometry investigation. It can import raw data for MS1 filtering from the instrument vendors Agilent, Bruker, SCIEX, Shimadzu, Thermo-Scientific and Waters, making the expertise you gain here transferrable to any mass spectrometry lab with an instrument from one of these vendors.

## **Getting Started**

To start this tutorial, download the following ZIP file:

https://skyline.ms/tutorials/DdaSearchMs1Filtering.zip

Extract the files in it to a folder on your computer, like:

C:\Users\brendanx\Documents

This will create a new folder:

C:\Users\brendanx\Documents\DdaSearchMS1Filtering

If you have been using Skyline prior to starting this tutorial, it is a good idea to revert Skyline to its default settings. To do so:

• Start Skyline.

• On the Start Page, click Blank Document which looks like this:



- On the **Settings** menu, click **Default.**
- Click **No** on the form asking if you want to save the current settings.

The document settings in this instance of Skyline have now been reset to the default.

Since this tutorial covers a proteomics topic, you can choose the proteomics interface by doing the following:

• Click the user interface control in the upper right-hand corner of the Skyline window, and click **Proteomics interface** which looks like this:



Skyline is operating in proteomics mode which is displayed by the protein icon **\*** in the upper right-hand corner of the Skyline window.

You could start editing this blank document in a number of ways, but for this tutorial you will use a sequential set of forms called a wizard that will walk you through the steps of searching mass spectrometer data dependent acquisition (DDA) data files, setting up targets, and importing chromatograms from those files.

Before starting the DDA search, you need to change the internal standard Skyline uses by default:

- On the **Settings** menu, click **Peptide Settings**.
- Click on the **Modifications** tab.
- Set the Internal standard type to "none"

The Peptide Settings form should now look like:

1 Peptide Settings	Х
Digestion Prediction Filter Library Modifications Quantification	
<u>Structural modifications:</u> ✓ Carbamidomethyl (C) <u>E</u> dit list	
Max variable mods: Max losses:	
Isotope label type: heavy ~ Isotope modifications:	
E <u>d</u> it list	
I <u>n</u> temal standard type: none	
OK Cancel	

• Click the **OK** button in the **Peptide Settings** form.

## Searching DDA Files to Load Peptides into a Skyline Document

You can use the **Import Peptide Search** wizard to run a peptide search on the MS/MS spectra in DDA data files.

First, do the following to save your new document:

- Click the **Save** button on the toolbar (Ctrl-S).
- Navigate to the DdaSearchMS1Filtering folder you created for this tutorial.
- In the File name field, enter 'DdaSearchMS1FilteringTutorial.sky'.
- Click the **Save** button.

Now, initiate the Import Peptide Search wizard as follows:

• On the File menu, choose Import and click Peptide Search.

Skyline should present a form that looks like this:

Spectral Library <ul> <li>Build</li> <li>Use existing</li> </ul> Start from:             Search results (build library directly) <ul> <li>Result files:</li> <li>File</li> <li>Score</li> <li>Type</li> <li>Threshold</li> <li>Remove Files</li> </ul>
Build Use existing     Start from:     Search results (build library directly)     Result files:     File     File     Score     Type     Threshold     Remove Files
Start from: Search results (build library directly) Result files: File File File Score Type Score Threshold Remove Files Start from: Add Files Remove Files
Search results (build library directly)          Result files:
Result files:          File       Score Type       Score Threshold       Add Files         Remove Files
File Score Type Score Threshold Remove Files
Remove Files
iRT standard peptides:
None V
Include ambiguous matches
Workflow     DDA with MS1 filtering
O DIA
O PRM
Finish Next > Cancel

In the **Start from** dropdown list, the **Search results (build library directly)** option works on the output from DDA search engines (e.g. pepXML files from Comet, .dat from Mascot) and the **DDA raw (search and build library)** option works on the raw data (e.g. RAW, WIFF, \*.d, mzML, mzXML). The mz5 files for this tutorial are centroided to make them faster to download than the original profile Thermo RAW files produced by the mass spectrometer.

Add the included DDA mz5 files to the search by doing the following:

- In the Start from dropdown list, select DDA raw (search and build library).
- Click the Add Files button.

- Select all the mz5 files in the DdaSearchMS1Filtering folder you created for this tutorial.
- Click the **Open** button.

The wizard form should now look like this:

🗽 Import Peptide Search	×
Spectral Library	
Build O Use existing	
Start from:	
DDA raw (search and build library) $\qquad \checkmark$	
Files to search: Cut-off score: 0.95	
File	Add Files
QE_140221_01_UPS1_100fmolspiked.mz5	Remove Files
QE_140221_02_UPS1_300fmolspiked.mz5	
QE_140221_03_UPS1_600fmolspiked.mz5	
iRT standard peptides:	
None	
Include ambiguous matches	
Workflow	
DDA with MS1 filtering	
O DIA	
O PRM	
Finish Next >	Cancel

• Click the **Next** button.

A form should appear asking you how to handle the prefix shared by the three mz5 files:

Import Results	$\times$
The files you have chosen have a common prefix and suffix. Would you like to remove some or all of the prefix or suffix to shorten the names used in Skyline?	
O Do not remove	
<u>R</u> emove	
Common prefix:	
QE_140221_0	
Common <u>s</u> uffix:	
spiked.mz5	
Replicate <u>n</u> ames: 1_UPS1_100fmol 2_UPS1_300fmol 3_UPS1_600fmol	
OK Cancel	

• Click the **OK** button.

The wizard will advance to the **Add Modifications** page, where it lists all of the amino acid modifications in the document that you might want to include in the DDA search. It is important to distinguish here between fixed and variable modifications: fixed (sometimes called static) modifications are always applied to the specified amino acids. For example, Carbamidomethyl C is usually treated as a fixed modification because all cysteines in the data are expected to be alkylated. Oxidation M is almost always treated as a variable modification because oxidation is hit-or-miss depending on sample handling. Skyline's search always treats isotopic labels as variable, but you can change whether other modifications are treated as fixed or variable by clicking the **Edit modifications** button. You can also add modifications to the document from this page. Since the document was reset to defaults, the list starts with only Carbamidomethyl (C):

1 Import Peptide Search			×
Add Modifications			
Specify all modifications you want to include in the search:			
Carbamidomethyl (C) = C[57] (fixed)			
Edit modifications			]
	< Back	Next >	Cancel

These data are SILAC labelled, so you will need to add heavy label modifications here. To add them, do the following:

- Click the Edit modifications button.
- Click the Edit heavy modifications menu option.
- Click the Add button in the Edit Isotope Modifications form.
- In the Name field of the Edit Isotope Modification form, enter "Label:13C(6)15N(2) (C-term K)".
- Click the down arrow at the right side of the **Name** field and click the item with the same name. This will populate the specificity and composition fields from Unimod.

The Edit Isotope Modification form should now look like this:

ta Edit Isotope Modification	×
<u>N</u> ame: Label:13C(6)15N(2) (C-term K)	OK Cancel
Amino acid: Teminus:	
Chemical fomula	
☑ 13C ☑ 15N 🗌 180 🗌 2H	
Monoisotopic mass: Average mass: 8.014199 7.941847	
Relative retention time:	
Matching	

• Click the **OK** button

Add a second isotope modification by doing the following:

- Click the Add button on the Edit Isotope Modifications form.
- From the **Name** dropdown list of the **Edit Isotope Modification** form, choose "Label:13C(6)15N(4) (C-term R)".

The **13C** and **15N** checkboxes are checked automatically to tell Skyline to use <sup>13</sup>C for all carbon atoms and <sup>15</sup>N for all nitrogen atoms present in an Arginine molecule, for a total mass shift of 10 Daltons (6x <sup>13</sup>C + 4x <sup>15</sup>N).

The Edit Isotope Modification form should now look like this:

<u>1</u> Edit Isotope Modificatio	on		×
<u>N</u> ame: Label:13C(6)15N(4) (C-term R)	1	~	OK Cancel
Amino acid: Terminus:	]		
Chemical formula			
🗹 13C 🗹 15N	180	🗌 2H	
Monoisotopic mass: 10.008269 Relative retention time:	A <u>v</u> erage mass 9.928665		
Matching ~			

Skyline automatically calculates both the monoisotopic and average masses, approximately 8 Daltons for Lysine (K) and 10 Daltons for Arginine (R) that will result from using 13C and 15N in these amino acid residues. To finish adding the heavy modifications:

- Click the **OK** button in the **Edit Isotope Modification** form.
- Click the **OK** button in the **Edit Isotope Modifications** form.
- Check the checkboxes in the Add Modifications list for the "Label:13C(6)15N(2) (C-term K)" and "Label:13C(6)15N(4) (C-term R)" modifications you just created.

Now you will add Oxidation (M) as a structural modification:

- Click the Edit modifications button.
- Click the Edit structural modifications menu option.
- Click the Add button in the Edit Structural Modifications form.
- In the Name field of the Edit Structural Modification form, enter "Oxidation (M)".
- Click the down arrow at the right side of the **Name** field and click the item with the same name. This will populate the specificity and composition fields from Unimod.
- Click the **OK** button in the **Edit Structural Modification** form.
- Click the **OK** button in the **Edit Structural Modifications** form.
- Check the checkbox in the Add Modifications list for the "Oxidation (M)" modification you just created.
- Also make sure the checkbox for "Carbamidomethyl (C)" is checked, as it should be because you chose the default settings.

At this point the **Add Modifications** page should look like:

<u>1</u> Import Peptide Search			×
Add Modifications			
Specify all modifications you want to include in the search:			
Carbamidomethyl (C) = C[57] (fixed)     Label:13C(6)15N(2) (C+erm K) = K[8] (isotopic label)     Label:13C(6)15N(4) (C+erm R) = R[10] (isotopic label)     Oxidation (M) = M[16] (variable)			
Edit modifications			
	< Back	Next >	Cancel

• Click the **Next** button.

The wizard will advance to the **Configure Full-Scan Settings** page.

• In the Mass Accuracy field, enter "20".

The other fields on this page should default to values you can use for this tutorial, and the wizard should look like this:

🗽 Import Peptide Search				×
Configure Full-Scar	Settings			
Precursor charges:				
MS1 filtering				
Count ~	Precursor mass analyzer: Centroided ~			
Peaks:	Mass Accuracy:			
Use high-selectivity extrac	ction			
Retention time filtering <ul> <li>Use only scans within</li> </ul>	5 minutes of MS/MS IDs			
	ans			
		< Back	Next >	Cancel

- See the MS1 Full Scan Filtering tutorial (page 9) to learn more about these settings.
- Click the **Next** button.

This should bring you to the **Import FASTA** page in the wizard. For this tutorial you will use a human protein FASTA with the sequences from Sigma-Aldrich's Universal Proteomics Standard (UPS) prepended at the top (so that Skyline uses those accession numbers for any peptides that are shared between UPS and non-UPS proteins). To select the FASTA:

- Click the **Browse** button.
- Select the '2014\_01\_HUMAN\_UPS.fasta' file in the folder you created for this tutorial.
- Click the **Open** button in the **Open FASTA** form.

The wizard should now look like this:

import Peptide Search	×
Import FASTA (required)	
Enzyme:       Max missed cleavages:         Trypsin [KR   P]       0         FASTA records begin with '>' and have the protein name followed by the optional protein description.	
/lineTester Results\DdaSearchMs1Filtering\DdaSearchMs1Filtering\2014_01_HUMAN_UPS.fasta Brow	ise
< Back Next > Ca	ncel

• Click the **Next** button.

The wizard will advance to the **Adjust Search Settings** page. Here you can set the most important parameters for the DDA search. For this tutorial, do the following:

- In the **MS1 tolerance** field, enter "5". (Note that when you leave the text box the form will assume you mean ppm and set the unit box accordingly.)
- In the **MS2 tolerance** field, enter "10".

The form should now look like this:

Import Peptide Search					2
Adjust Search Set	tings				
Search engine: MSAmanda V					4
MS1 tolerance: 5 MS2 tolerance:	Unit: ppm v Unit:	/			
10 Fragment ions:	ppm 、				
MS2 analyzer: Default		· ·			
Max variable mods:					
Additional Settings					
			< Back	Next >	Cancel

• Click the **Next** button to start the search.

The **DDA Search** page will show you the progress of the search. You may also cancel the search by clicking the **Cancel Search** button on this page.

earch progress:	Show timestamp
nput files contain 106460 ms2 spectra Searching spectra 1 - 10000 of 106460 spectra Starting spectra 1 - 10000 of 106460 spectra Starting search against fwd database Loading database Loading database done Search against fwd database finished. Starting search against decoy database Loading database Loading database done Search against decoy database finished. Search against decoy database finished. Search finished for 10000/106460 spectra Starting search against fwd database Search against fwd database finished. Starting search against fwd database Search against decoy database finished. Starting search against decoy database Search against decoy database finished. Starting search against decoy database Search finished for 20000/106460 spectra Starting spectra 20001 - 30000 of 106460 spectra Starting search against fwd database Search finished for 20000/106460 spectra Starting search against decoy database finished. Starting search against decoy database Search against decoy database finished. Starting search against decoy database Search against decoy database finished. Starting search against decoy database Search against decoy database finished. Search finished for 30000/106460 spectra Search against decoy database Search against decoy database Search finished for 30000/106460 spectra Starting search against fwd database Search against decoy database Search against decoy database Search finished for 30000/106460 spectra Starting search against fwd database	
Constituent	

After the search is finished:

• Click the **Finish** button.

Skyline will start to build a spectral library from the search results. After the library build is completed, a message box will pop up to warn you that there were some spectra that could be interpreted as multiple different peptides with equal confidence, and that those peptides will be ignored:

Skyline >	×
The library built successfully. Spectra matching the following peptides had multiple ambiguous peptide matches and were excluded: AAAPTLSEHFR[+10.00827] AAELIANSLATAGDGLIELR AALLSTQTLPAPSPTSPPGGILK[+8.0142] AALQSQENLIHEAR[+10.00827] AAVLQWTSVTAQVTFR ADGESC[+57.02146]SASM[+15.99492]/YQEGK ADGESC[+57.02146]SASM[+15.99492]YQEGK ADHQPLTEASYVNLPTIALC[+57.02146]NTDSPLR ADHQPLTEASYVNLPTIALC[+57.02146]NTDSPLR ADHQPLTEASYVNLPTIALC[+57.02146]NTDSPLR ADHEMAVSYLK[+8.0142] ADVLAFPSSGFTDLAEIVSR[+10.00827] AEASPSSAAVSSLIEK AELMEILDLHR[+10.00827] AEASPSSAVSSLIEK AFASLR AFATVR AFATVR AFATVR AFATVR AFATVR AFAVTR AFATVR AFAVTR AFAVTR AGSIVLK[+8.0142] AGTVIANVAK AGSIVLK[+8.0142] AGTVIANVAK AGSIVLK[+8.0142] AIELGIYPAVDPLDSTSR AIDVSVM[+15.99492]VGGSTLAK AIELEK[+8.0142] AIELER AGLER AGSSEGAQSSLOEVYHK[+8.0142] AIELDNANAVYFC[+57.02146]NR AIEMELR AIGLER AIGSEG AIESDQEQGR[+10.00827] ALVDLEPGTM[+15.99492]DSVR[+10.00827] ALVDLEPGTM[+15.99492]DSVR[+10.00827] ALVDLEPGTM[+15.99492]DSVR[+10.00827] ALASR[+10.00827] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALAPTWEQLALGLEHSETVK[+8.0142] ALEINSIAPIVIFASNR	
ALGEGAAWWQSQR	

• Click the **OK** button to dismiss the message.

Skyline will then start to import the library into your document. When that is finished it will ask you to set criteria for including proteins in the document:

Associate Proteins				?	×
Organize all documer	nt peptides into associ	ated proteins or protein grou	ups.		
Protein parsimony op	otions oups <sup>?</sup>				
<u>S</u> hared peptides are	: <sup>2</sup>				
Assigned to first prot	ein		$\sim$		
Find minimal prote	ein list that explains all proteins <sup>2</sup> ptein: sults:	peptides <sup>-</sup>			
	Mapped	Unmapped	Targets		
Proteins	19,749	68,778	6381		_
Peptides	12,800	0	12,800		
Shared Peptides	44,862		0		
	6381 proteir	ns, 12,800 peptides, 25,376	oprecursors, 76,12 OK	8 trans Cano	sitions cel

Skyline can merge proteins that match the same set of peptides into protein groups and find a minimal list of proteins that explain all the peptides. You can read more about protein grouping and parsimonious protein inference in this documentation page: <u>Skyline Protein Association</u>. For this tutorial, leave the checkboxes for creating protein groups and finding a minimal list of proteins unchecked.

- Select Assigned to first protein from the Shared peptides are: dropdown list.
- Click the **OK** button.

## **Configuring Skyline to View the Imported Data**

After the proteins are imported into the document, you will see the main Skyline window with the UPS proteins at the top of the **Targets** view. You should see **5,219** proteins there (counted in the status bar). Skyline will also begin extracting chromatograms with progress displayed in an **Importing Results** form. You can wait several minutes for chromatogram extraction to complete or you can continue with all but the last step below during the import.

- Click on the [+] icon next to the third protein P01112ups|RASH\_HUMAN\_UPS.
- Click on the [+] icon next to that protein's third peptide R.TGEGFL<u>CVFAINNTK.S.</u>

- Click on that peptide's first precursor 835.9140++ and the chromatogram for that precursor and the MS/MS spectrum for that peptide will appear. (Note that the bold, underlined residue "<u>C</u>" in the peptide sequence indicates a carbamidomethyl cysteine).
- If you do not see the MS/MS spectrum, on the View menu, click Library Match.
- If you do not see as many annotated peaks in the Library Match view as in the image below, on the View menu, choose Ion Types and check B and Y.
- If you do not see the entire chromatogram for the peptide, on the **View** menu, choose **Auto-Zoom** and click **None** (Shift-F11).
- Right-click the chromatogram graph, choose Peptide ID Times, and click Aligned if it is unchecked.

Your Skyline window should look something like this:



The document is now fully configured for MS1 Filtering with three DDA runs imported. You will see that the chromatogram in this view is approximately 10 minutes in length (133 to 143 minutes), due to the **Use only scans within [5] minutes of MS/MS IDs** setting chosen in the import wizard. Note that when the Skyline document is set up for MS1 Filtering, in the place where you would see product ion transitions (e.g. y-ions) for triple quadrupole SRM experiments, you will now see the different precursor isotope peaks, such as for peptide TGEGFL<u>C</u>VFAINNTK: precursor – 835.914++, precursor [M+1] - 836.4154++, and precursor [M+2] – 836.9164++.

To configure a few other features that will be helpful in general, especially to visualize certain MS1 Filtering data, perform the following steps:

• On the Settings menu, make sure Integrate All is checked.

This tells Skyline to treat all chromatograms in a peak group (here precursor ions M, M+1 and M+2) as integrating together, regardless of whether peaks appear to be co-eluting with the largest peak. It no longer impacts the integrated peak areas as it once did.

• On the View menu, choose Peak Areas and click Replicate Comparison.

A window with a graph like this should appear:



You can dock the **Peak Areas** window to your desired location by doing the following:

- Click and hold down the left mouse button, and then drag it until the mouse cursor is above the Library Match view.
- When a set of 5 icons in the shape of a cross appears, move the mouse into the lower icon and release the left mouse button to divide the space on the right edge of the Skyline window between the **Peak Areas** and **Library Match** views.
- Move the Library Match view below the Targets view by the same process.
- On the View menu, choose Auto-Zoom and click Best Peak (F11).
- On the **View** menu, choose **Arrange Graphs** and click **Column**.
- Right-click a chromatogram graph, and click **Legend** to turn off the legend in the chromatogram graphs.

Now your Skyline window should look something like this:



From this point you can learn more about working with DDA data by consulting the <u>MS1 Full Scan</u> <u>Filtering tutorial</u>.