glyXtoolMS Usermanual

version 2.0

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'glyXtoolMS' is an open-source software for the analysis of glycopeptide mass spectrometry data.

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# Installation

## OpenMS

For installation of OpenMS visit <https://www.openms.de/> and follow the download/install or build instructions for your operating system.

After installation the following tools should be installed: TOPPAS and TOPPView.

## Python

To run glyXtoolMS, a python 2.7 installation is required, together with the package manager pip. The use of a virtual environment like virtualenvwrapper is recommended if other python versions are/will be installed on the same workstation.

Install python 2.7 from <https://www.python.org/>. The package manager for python will then be installed, too. To check, open a console and type the command “pip”. If it has not been installed, follow the installation instructions on <https://pip.pypa.io/en/stable/installing/#do-i-need-to-install-pip>.

The use of a virtual environment is recommended, in case multiple python installations with different package setups are installed on the computer. For the installation of virtualenvwrapper, please refer to <https://virtualenvwrapper.readthedocs.io/en/latest/>

Virtualenvwrapper can be installed via:

pip install virtualenvwrapper

afterwards a fresh environment can be created using:

mkvirtualenv <envname>

switch into the environment using:

workon <envname>

## glyXtoolMS

glyXtoolMS can be installed using pip in the command line:

pip install glyXtoolMS

The dependencies canvasvg, configparser, lxml ,numpy,pyopenms, pyperclip, and xlwt should then be automatically downloaded and installed.

alternatively the .egg or .wheel can be downloaded from <https://pypi.org/project/glyxtoolms/>

or build manually from https://github.com/glyXera/glyXtoolMS

After the installation of glyXtoolMS, the glyXtoolMS Evaluator should accessible via the console command:

glyXtoolMS

## TOPPAS Script Setup using glyXtoolMS Evaluator

During the first startup of the glyXtoolMS Evaluator, a configuration window for OpenMS will appear(also later available on the Menu/TOPPAS/Configure TOPPAS), since all necessary scripts and tool description files need the be copied from the python package into OpenMS. Please provide the installation path of OpenMS. During the save process all necessary files will be copied over into the OpenMS/share/OpenMS/SCRIPTS/ and OpenMS/share/OpenMS/TOOLS/EXTERNAL/ directories.

Within the same window downloaded TOPPAS workflows (e.g. from the example data sets) can be adapted to the right SCRIPT path.

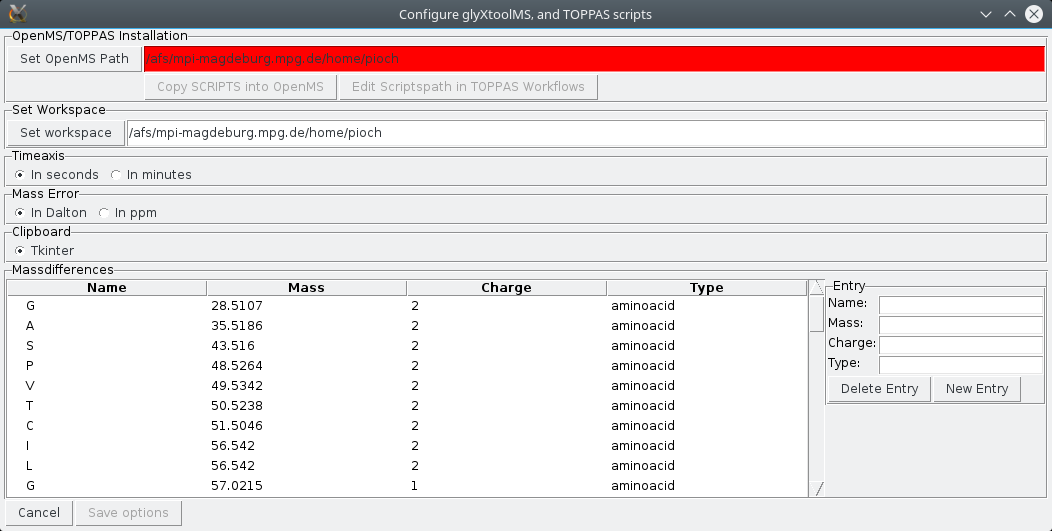


Figure 1: OpenMS Path Configuration

The Configuration Panel can be reached duringthe first startup of the glyXtoolMS Evaluator, or opened via Menu/Program/Configure. Set the Path to your OpenMS Installation.

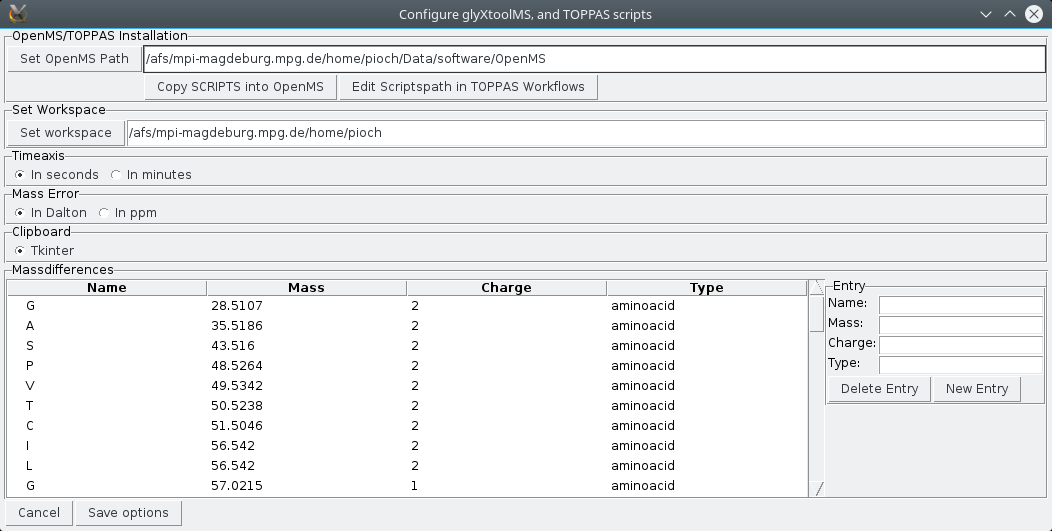


Figure 2: Copy Scripts into OpenMS and edit TOPPAS Workflows

Afterwards the Scripts can be copied over into OpenMS in order to make them available within the TOPPAS pipeline engine. Downloaded TOPPAS workflows can be edited to include the correct scriptpath.

# Analyzing the Example Data Set

The example data sets of human IgG and human fibrinogen used within the manuscript can be downloaded from <https://www.ebi.ac.uk/pride/archive/projects/PXD009716>, containing the raw data files, and – within the zip file – the converted raw mass spectrometry files, the FASTA files, the *N*- and *O*-glycan databases, the TOPPAS workflows, the generated reduced mass spectrometry files, the detected features and the analysis files.

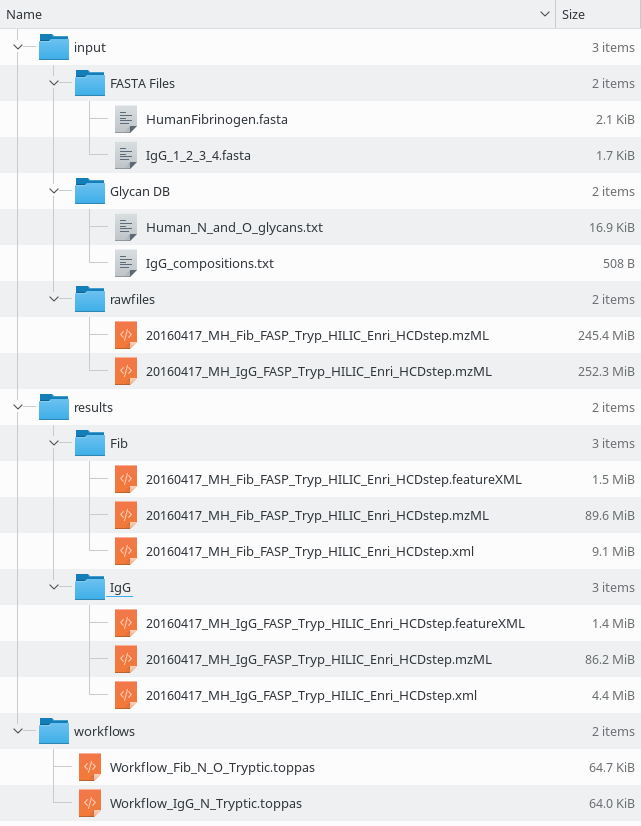


Figure 3: The extracted files from the zip folder within the example data set

The raw data can be analyzed within TOPPAS (see Section 2.1) using the files within the input directory and the corresponding workflow or the provided result files can be viewed/analyzed with the glyXtoolMS Evaluator (see Section 2.2).

## New Analysis with TOPPAS

1. Start the glyXtoolMS Evaluator and edit the workflows like described in Section 1.4
2. Start TOPPAS

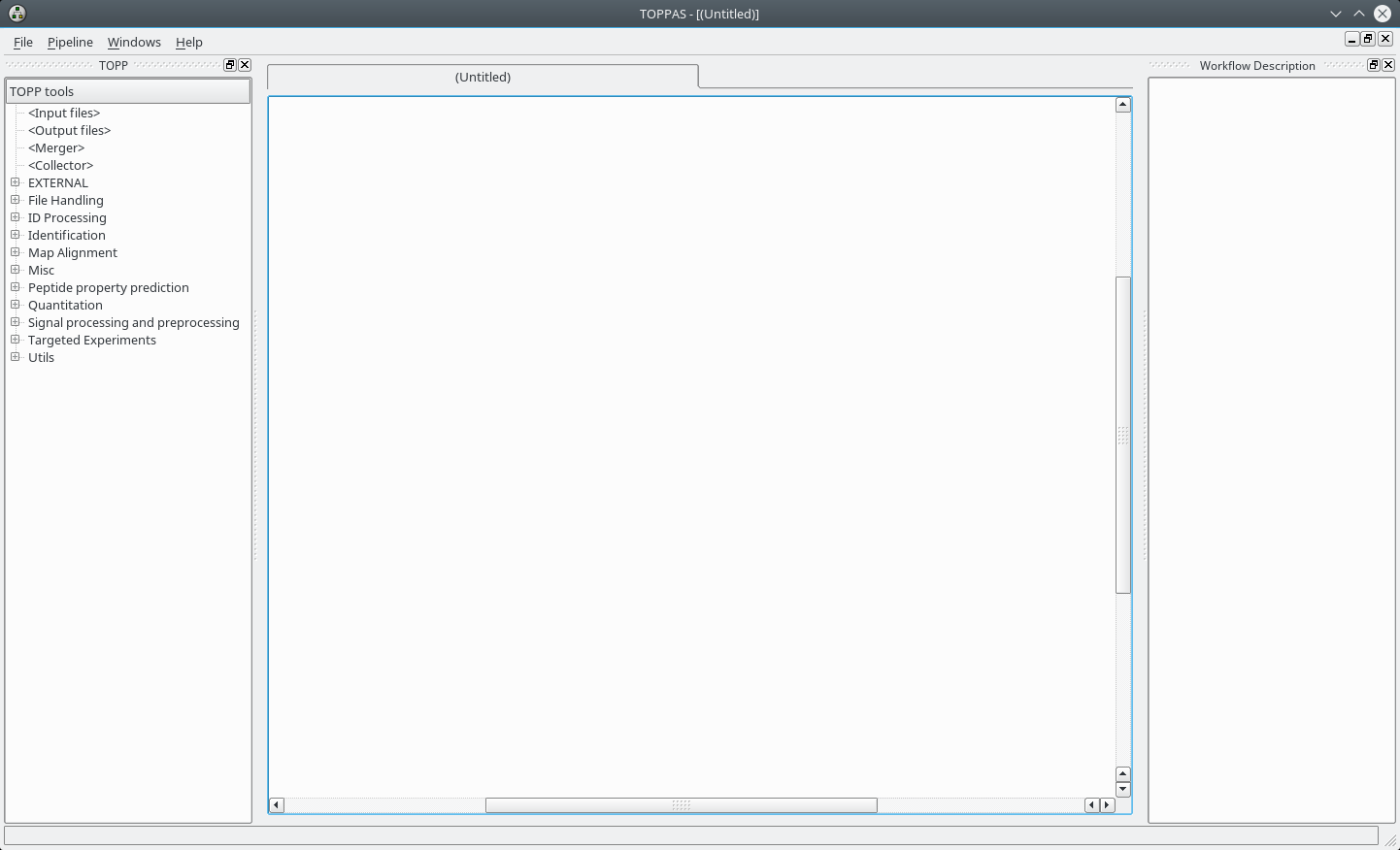


Figure 4: The TOPPAS window

1. Check the existence of the glyXtoolMS scripts. If tools are missing check Section 1.4

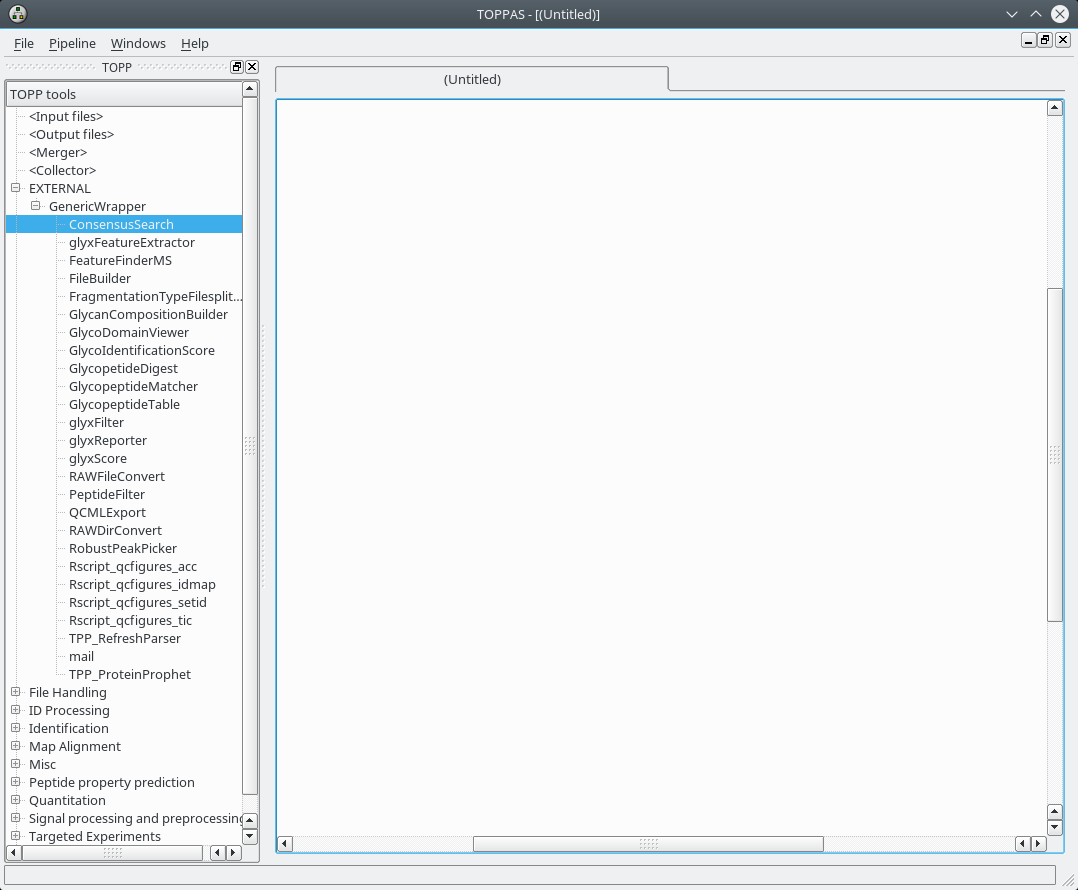


Figure 5: glyXtoolMS scripts are located under “External/GenericWrapper/…”

1. Open the IgG workflow

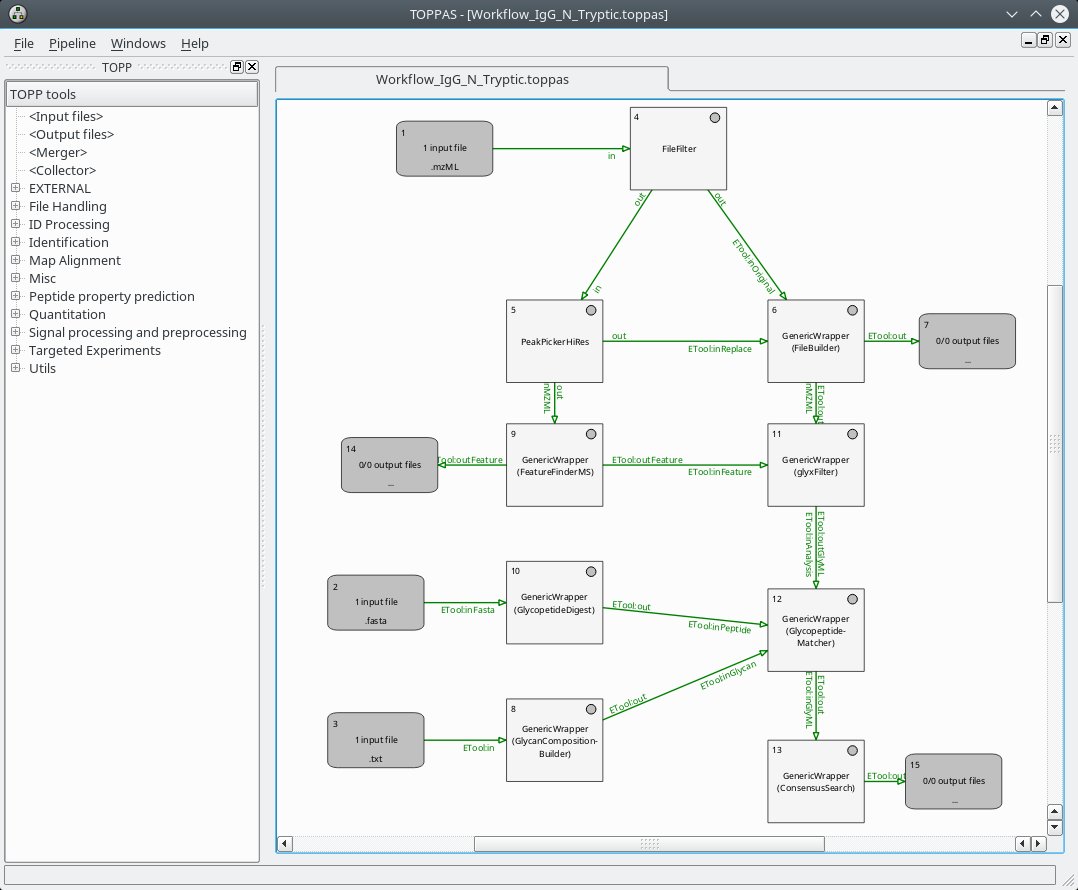


Figure 6: The IgG TOPPAS workflow

1. Double clicking on a tool opens the tool configuration. Check a Generic Wrapper tool for the correct OpenMS Script Path. If not correct, check Section 1.4.

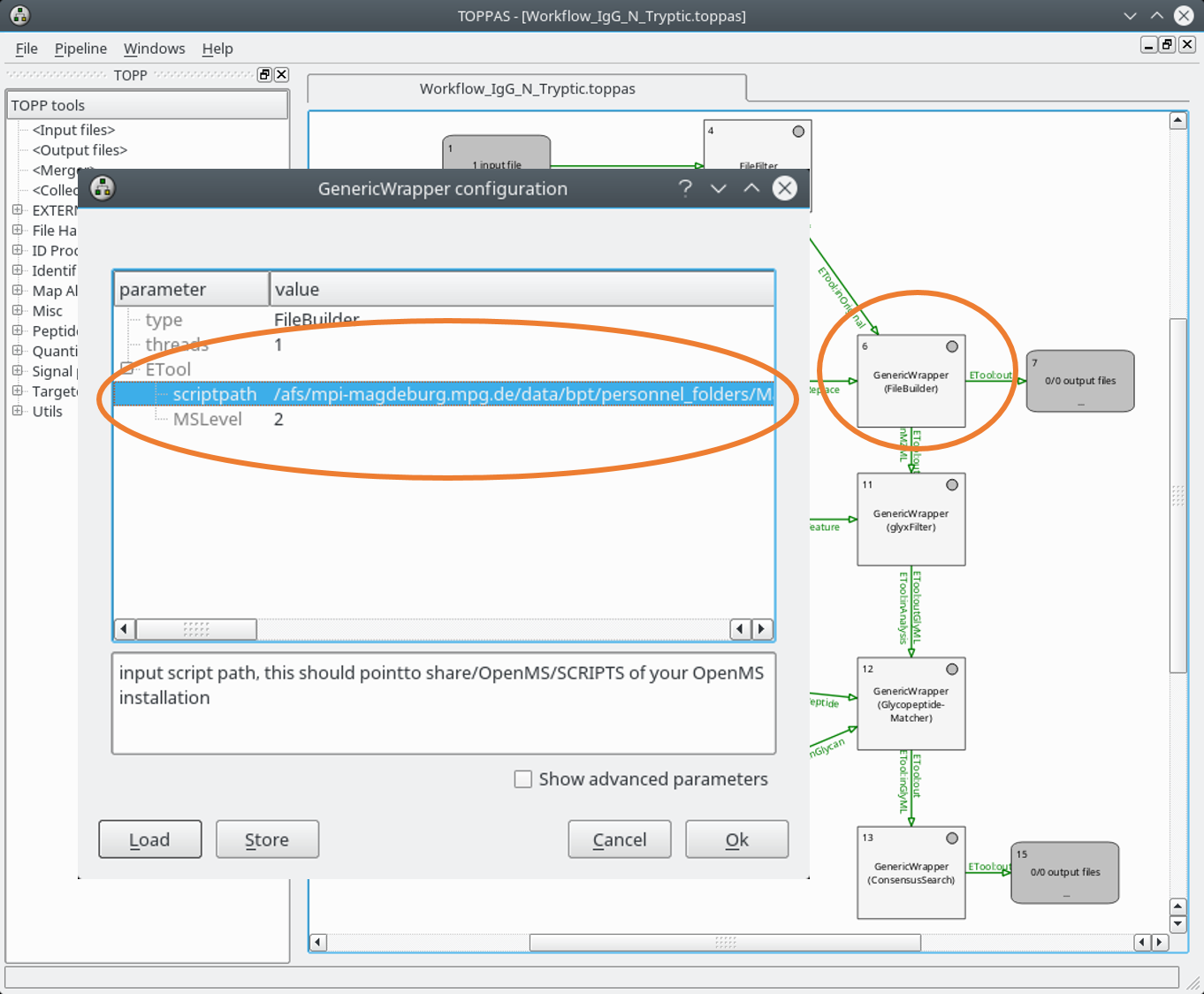


Figure 7: Check OpenMS Path

1. Set the input files:

* A: the IgG mass spectrometry file rawfiles/20160417\_MH\_IgG\_FASP\_Tryp\_HILIC\_Enri\_HCDstep.mzML
* B: the IgG FASTA file from input/FASTA Files/IgG\_1\_2\_3\_4.fasta
* C: the glycan database from input/Glycan DB/Human\_N\_and\_O\_glycans.txt

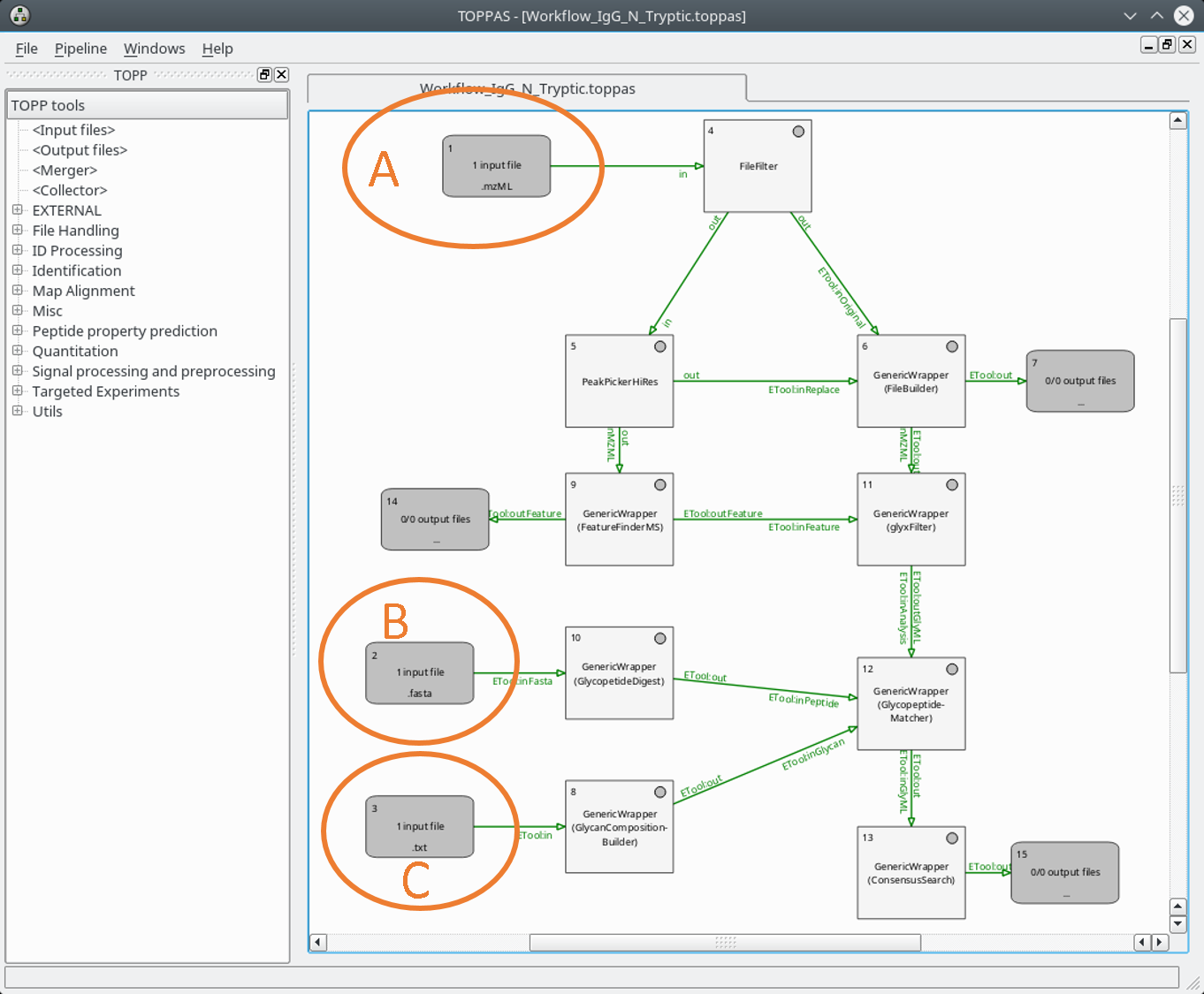


Figure 8: Set input files

1. Run the TOPPAS Pipeline. Select an output folder (TOPPAS will create a folder structure “TOPPAS\_out” containing the files created by the output nodes)

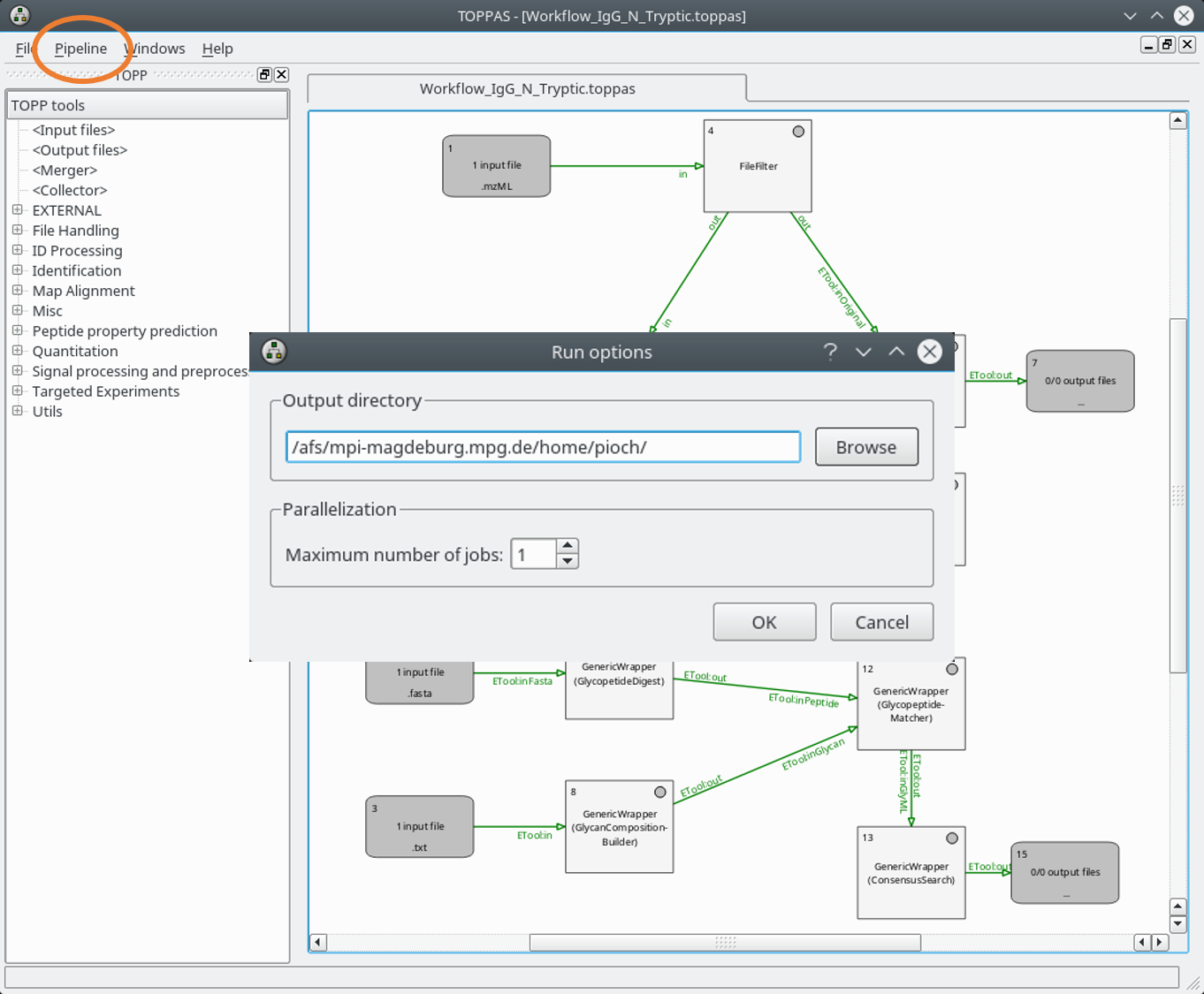


Figure 9: Run the TOPPAS pipeline

1. The resulting analysis files are created during the successful run of the analysis:

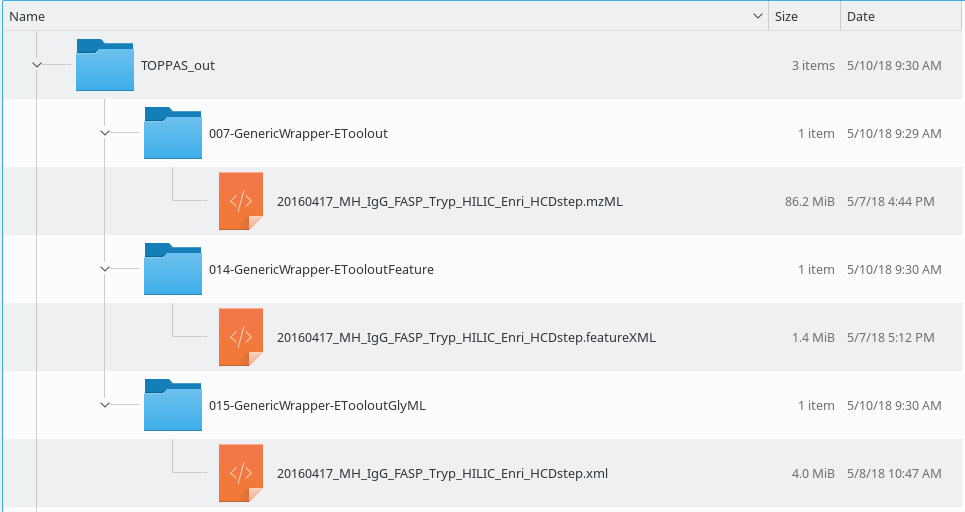


Figure 10: Analysis files created by the IgG TOPPAS workflow

Move all files into a result folder. The created files should correspond to the ones within the example data set stored under “results/IgG”.

## View Analysis with glyXtoolMS Evaluator

1. Start the glyXtoolMS Evaluator via commandline:

glyxtoolms

1. Create new Project with “New Project”. Select the mzML file from the result (it contains continuous MS1 spectra and centroided MS2 spectra) and provide a project name

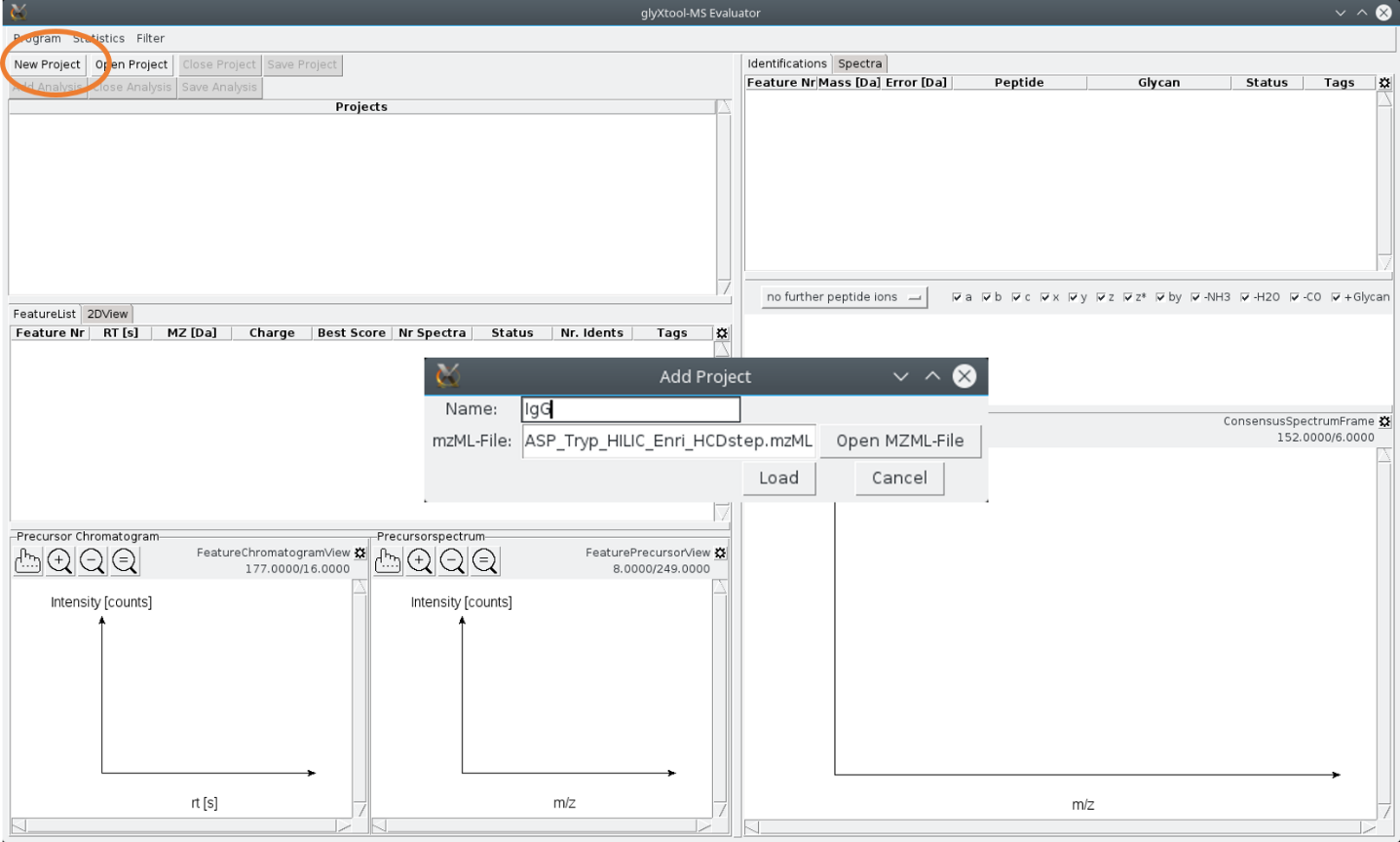


Figure 11: Software Surface

1. Add an Analysis file to the project, by selecting the project, and then using the “Add Analysis Button”. To each Project multiple analysis files can be loaded (originating from the same raw data file). Saving the Project stores a simple file with the project name, its mzML file path and the path to each analysis file. This is only for easy access to the files – saving the project does not save changes to the analysis file!. Analysis files can be saved separately via the “Save Analysis” Button.

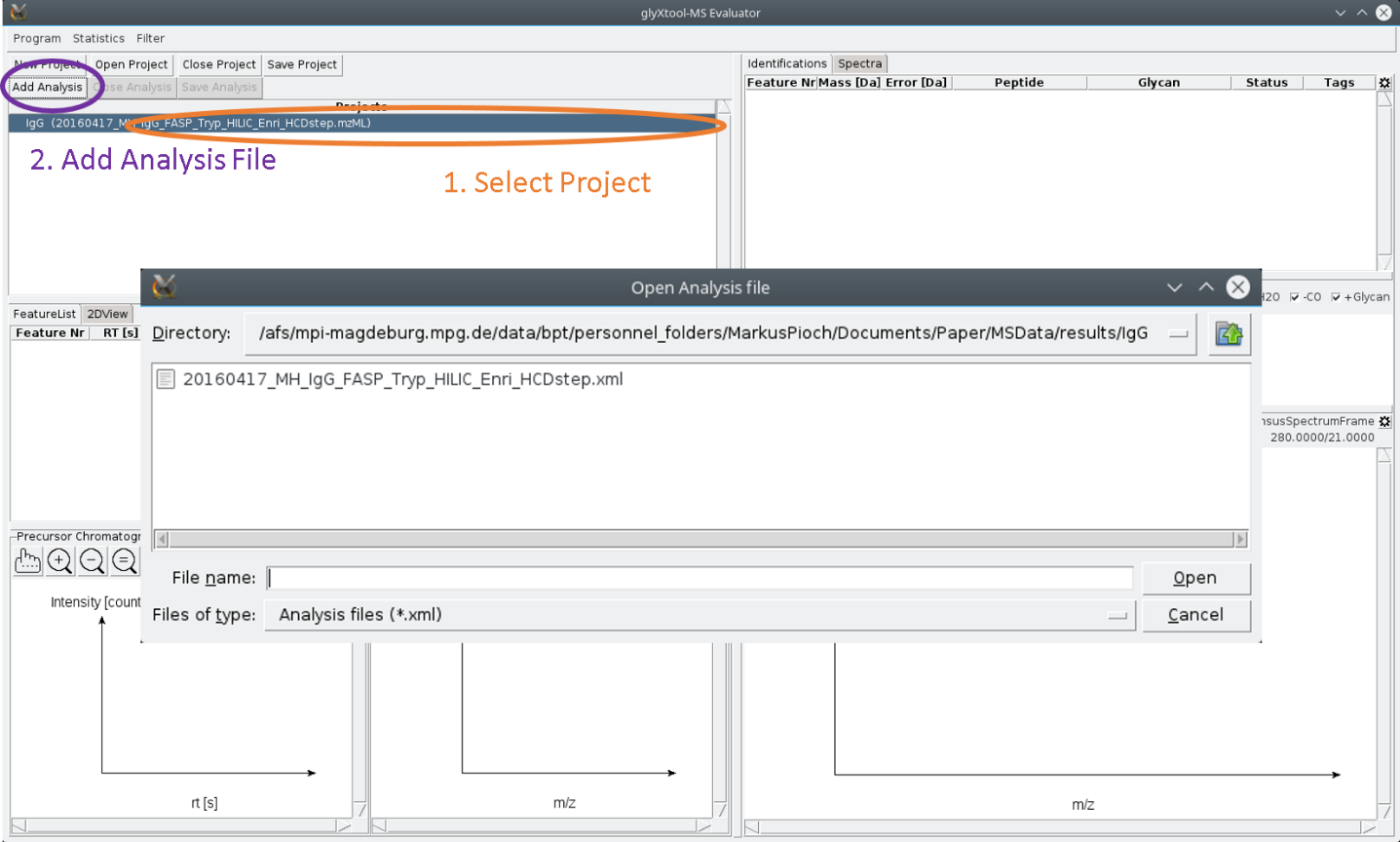


Figure 12: Add Analysis

1. Loaded Analysis File: In the “FeatureList” all features/compounds are listed that were found by the FeatureFinder and with have been identified as potential glycoepetides via the glyXFilter tool. The best Oxonoimion-Score is shown within the table.

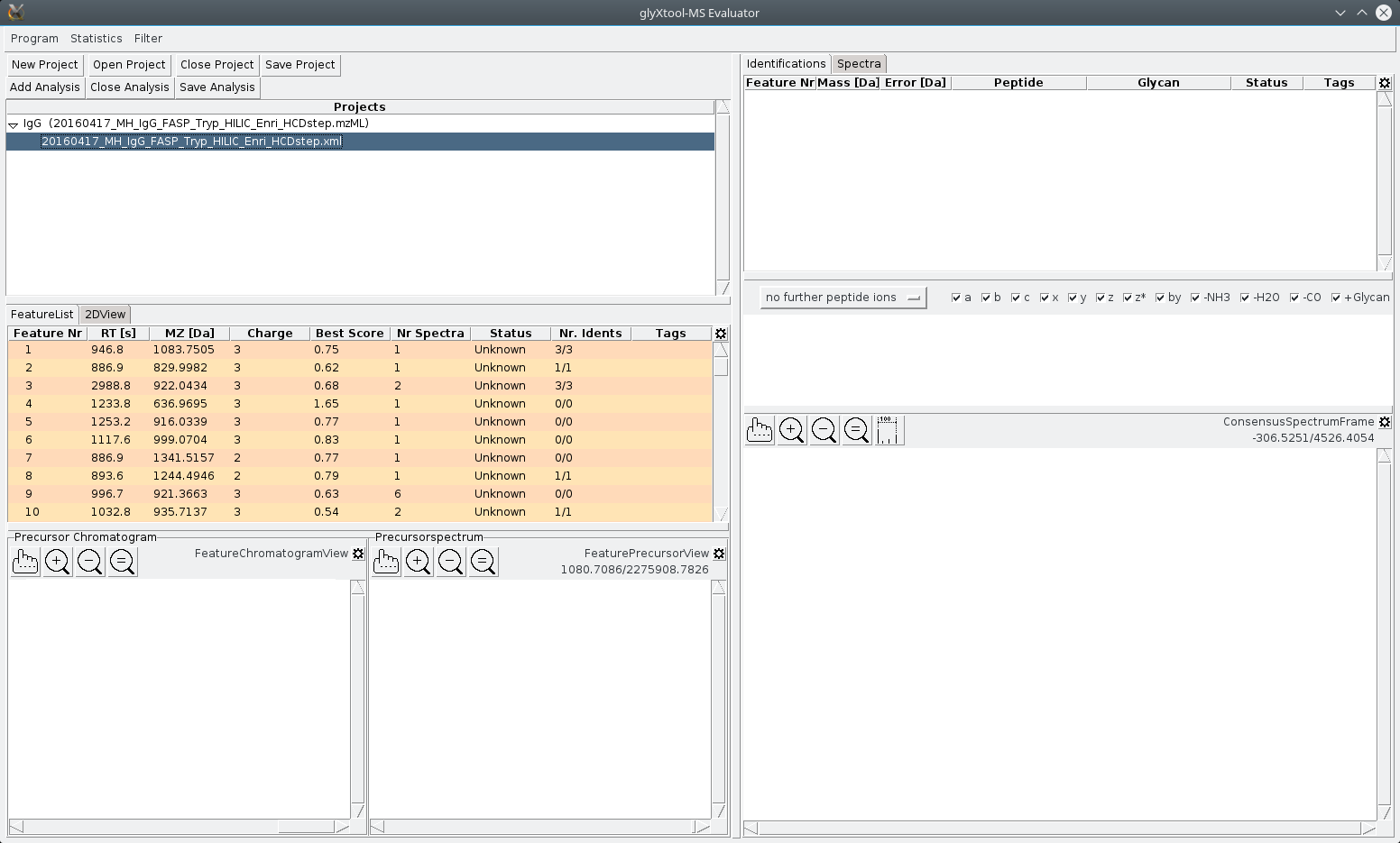


Figure 13: Loaded Analysis file

1. Show the oxonium scoring results for the glyXFilter tool:

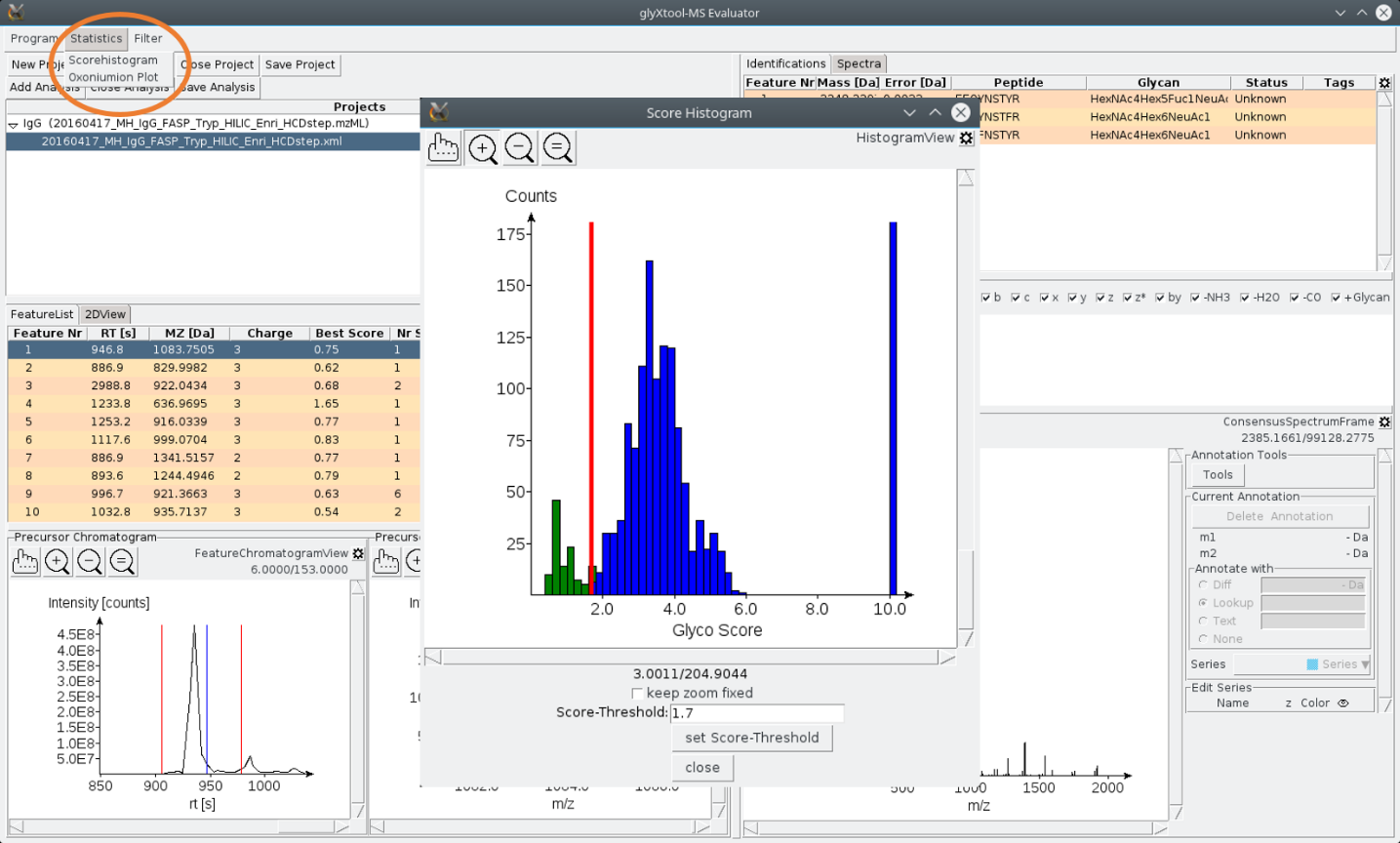


Figure 14: Histogram of Oxonium ion scoring. The threshold should divide two distinct populations of glycopeptids (green) and non glycopeptides (blue)

1. Selecting a Feature within the “FeatureList” shows the extracted ion chromatogram of the monoisotopic precursor peak, the isotopic pattern of the precursor and the consensus spectrum of all fragment spectra within its feature box. On the right side within the Identifications tab – all possible precursor mass matches of theoretical peptides and glycan compositions are shown, generated by the GlycopeptideMatcher tool. The Spectra tab shows the single fragment spectra associated with the feature.

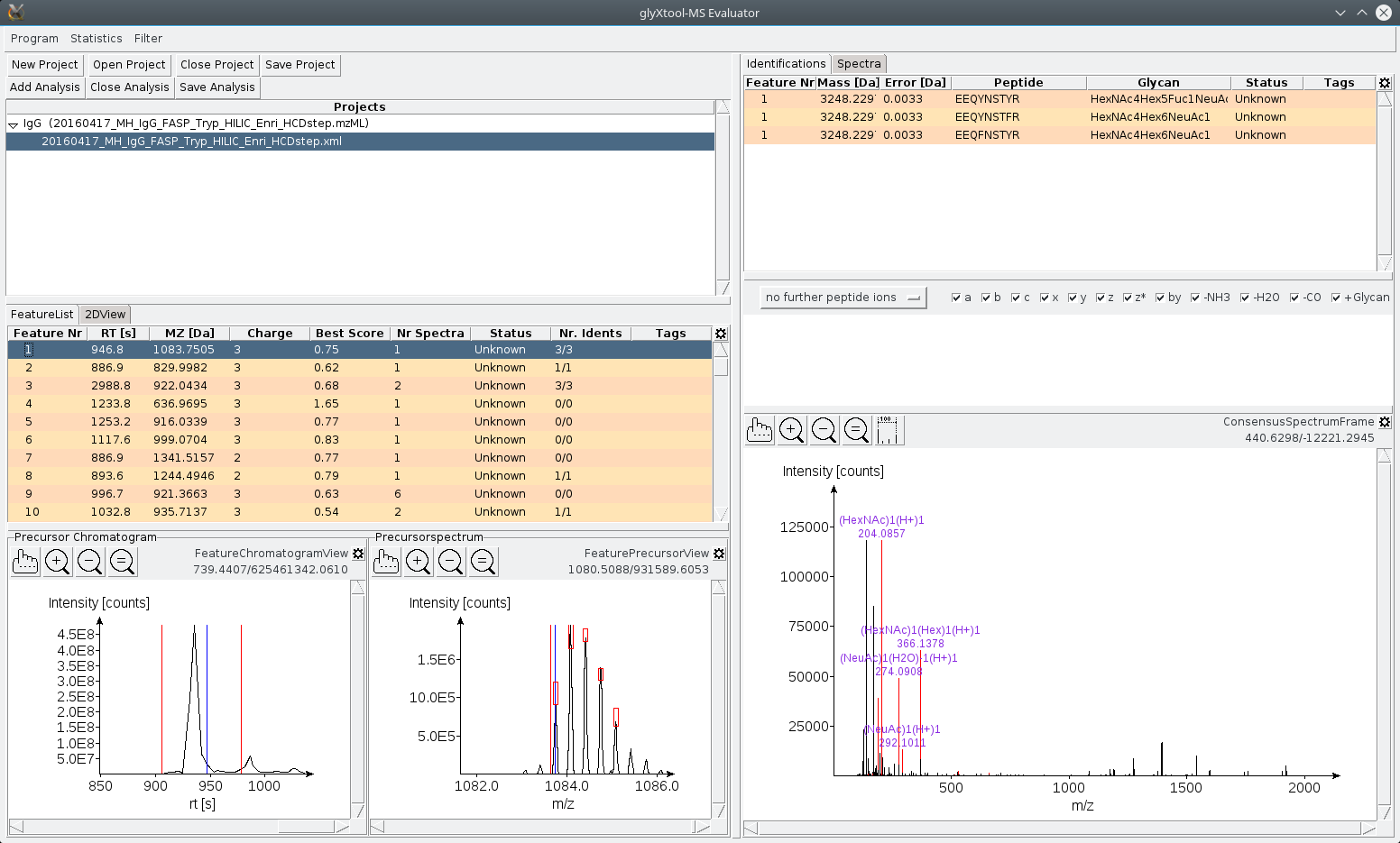


Figure 15: Selecting a feature

1. Selecting an identification within the “Identifications” tab annotates the consensus spectrum with its theoretical ion fragments. Using the “Gear icon” a configuration panel for the consensus spectrum is shown, where ion names and colors can be adapted.

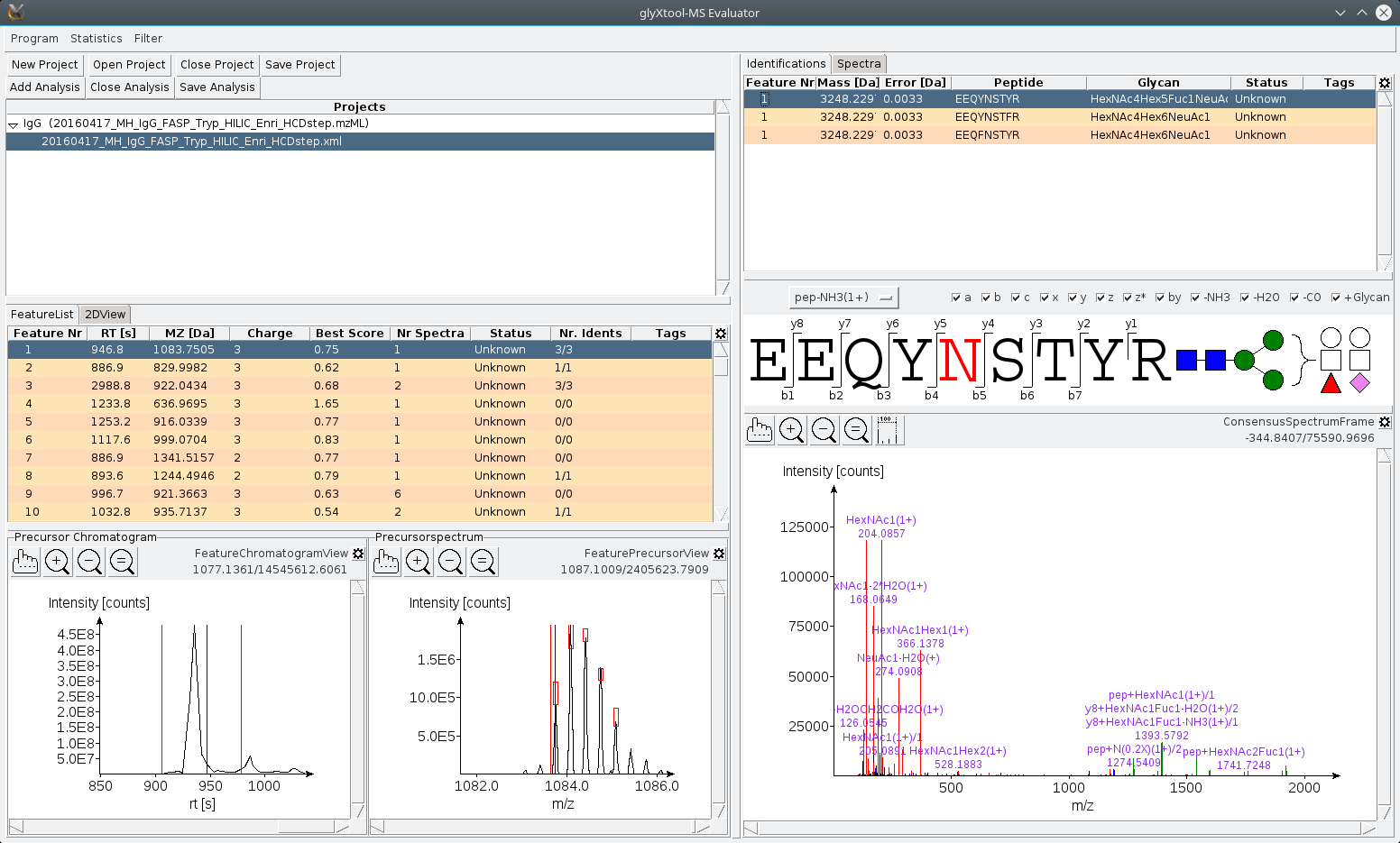


Figure 16: Selecting an identification

1. Selecting a peptide ion shows all occurrences of this ion with the fragment spectrum

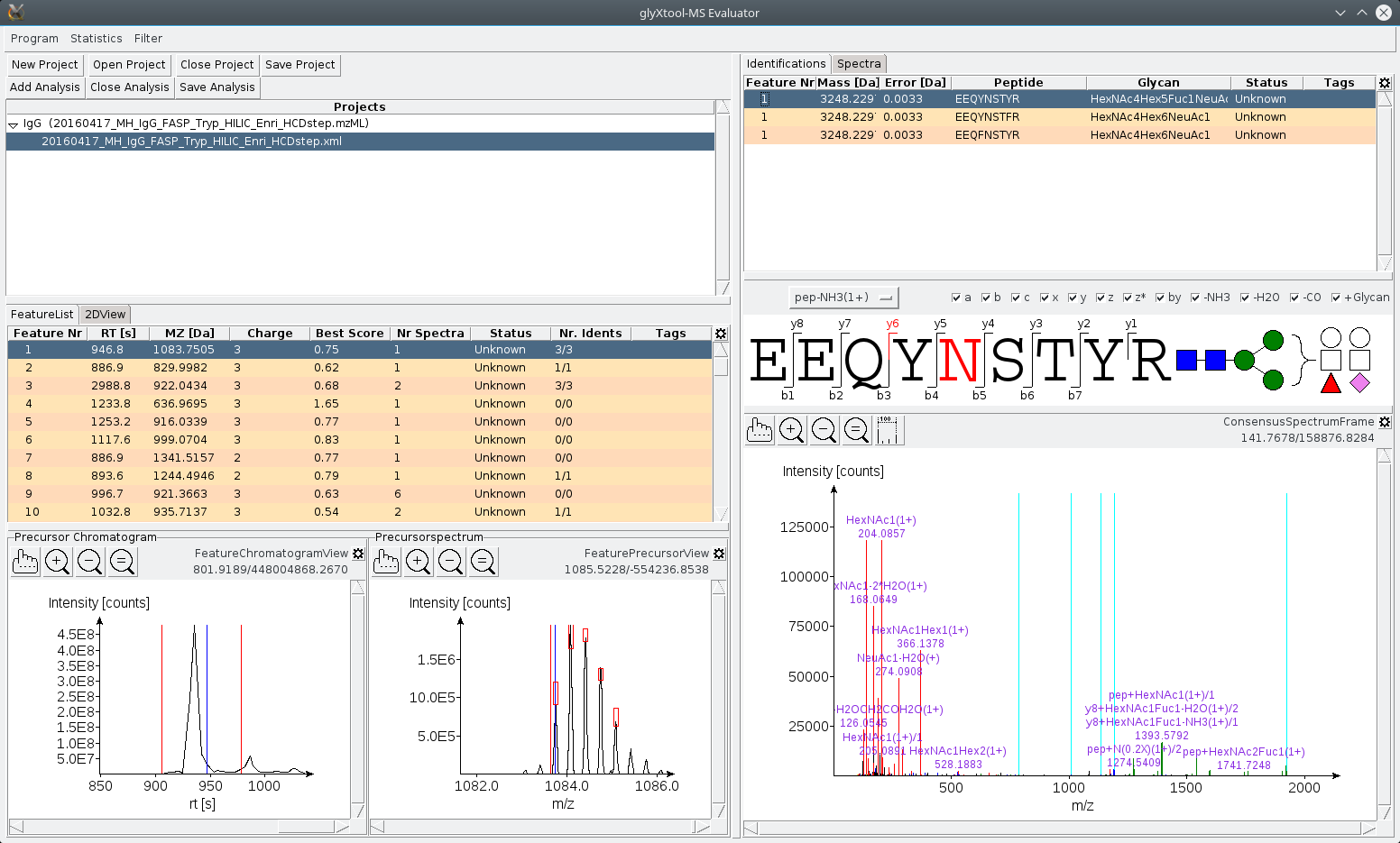


Figure 17: Selecting a peptide ion

1. Right clicking on a selected feature or identification opens a context menu for analysis. A Status of “Unknown”, “Accepted” or “Rejected” can be set, tags can be added, etc.

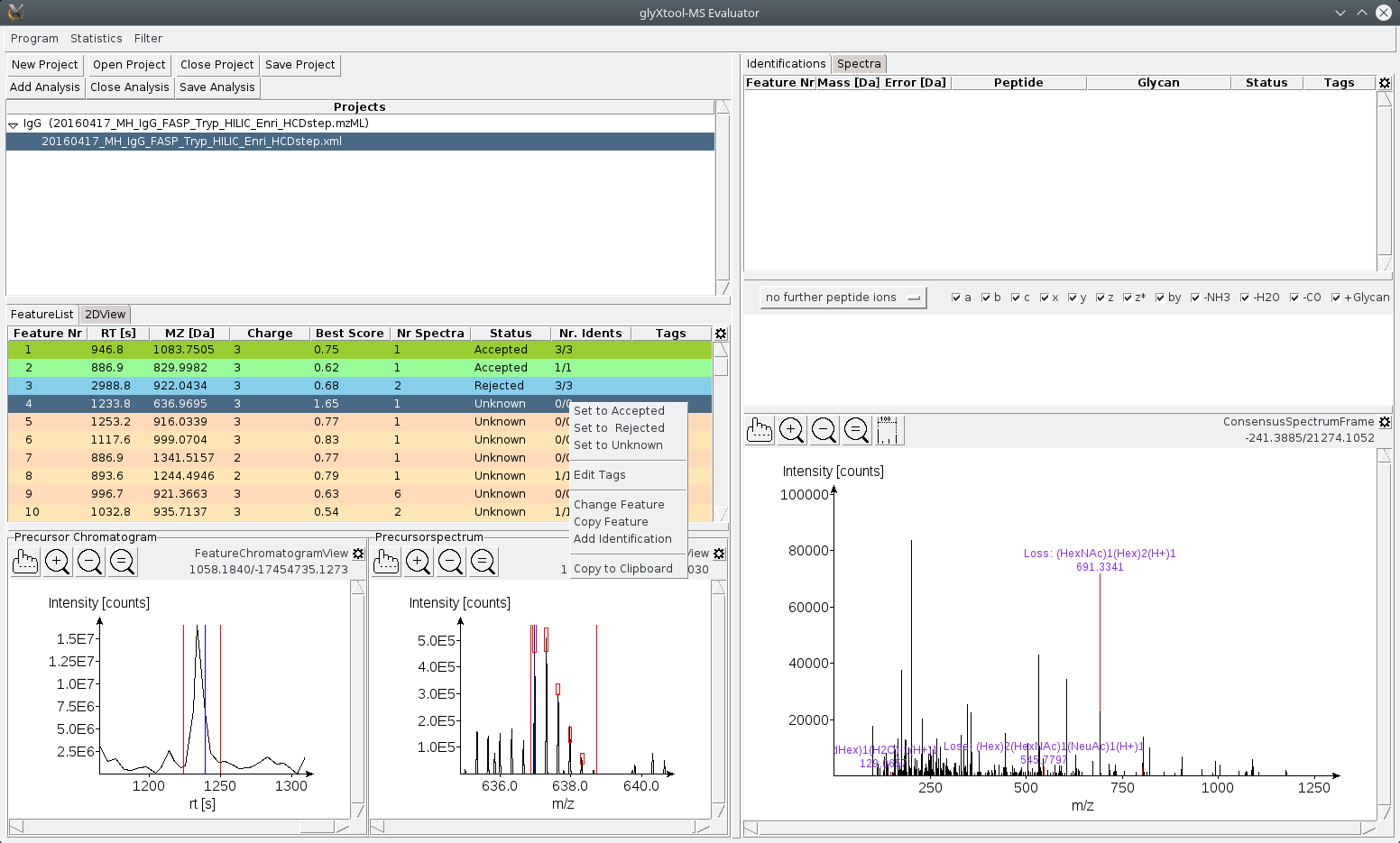


Figure 18: Context menu on either features or identifications

1. Fragment ion annotation series can be added by activating the ruler icon: Either by left clicking on the peak of interest and pulling to the side, or right clicking where a menu shows potentially interesting mass differences to neighbor peaks.

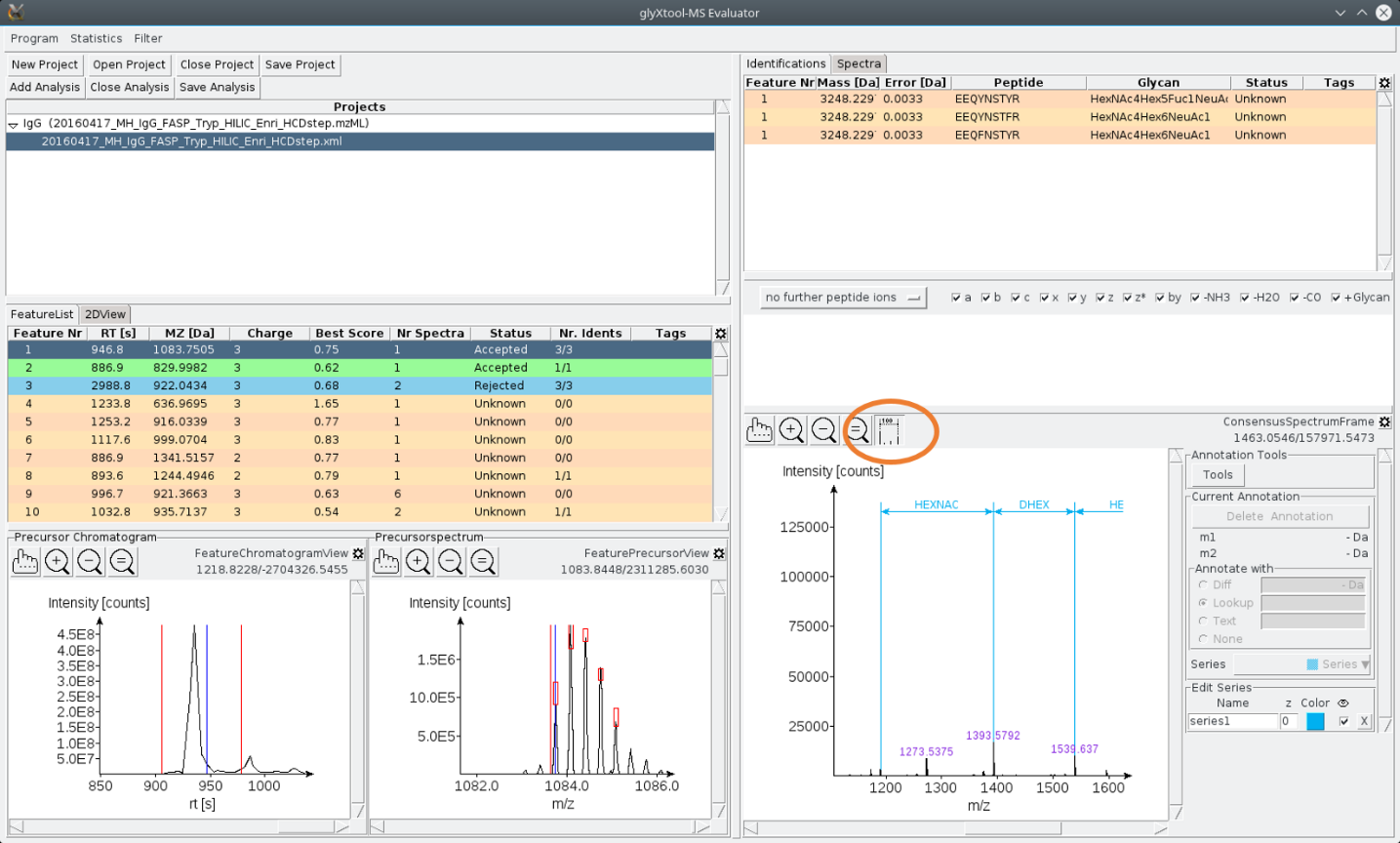


Figure 19: Adding fragment ions annotation series

1. Both the “FeatureList” and the “Identifications” tab support multi selection, if e.g. two features are selected, the identifications are shown for all. By selecting all features with “Cntrl+A” all identifications can be selected, for example.

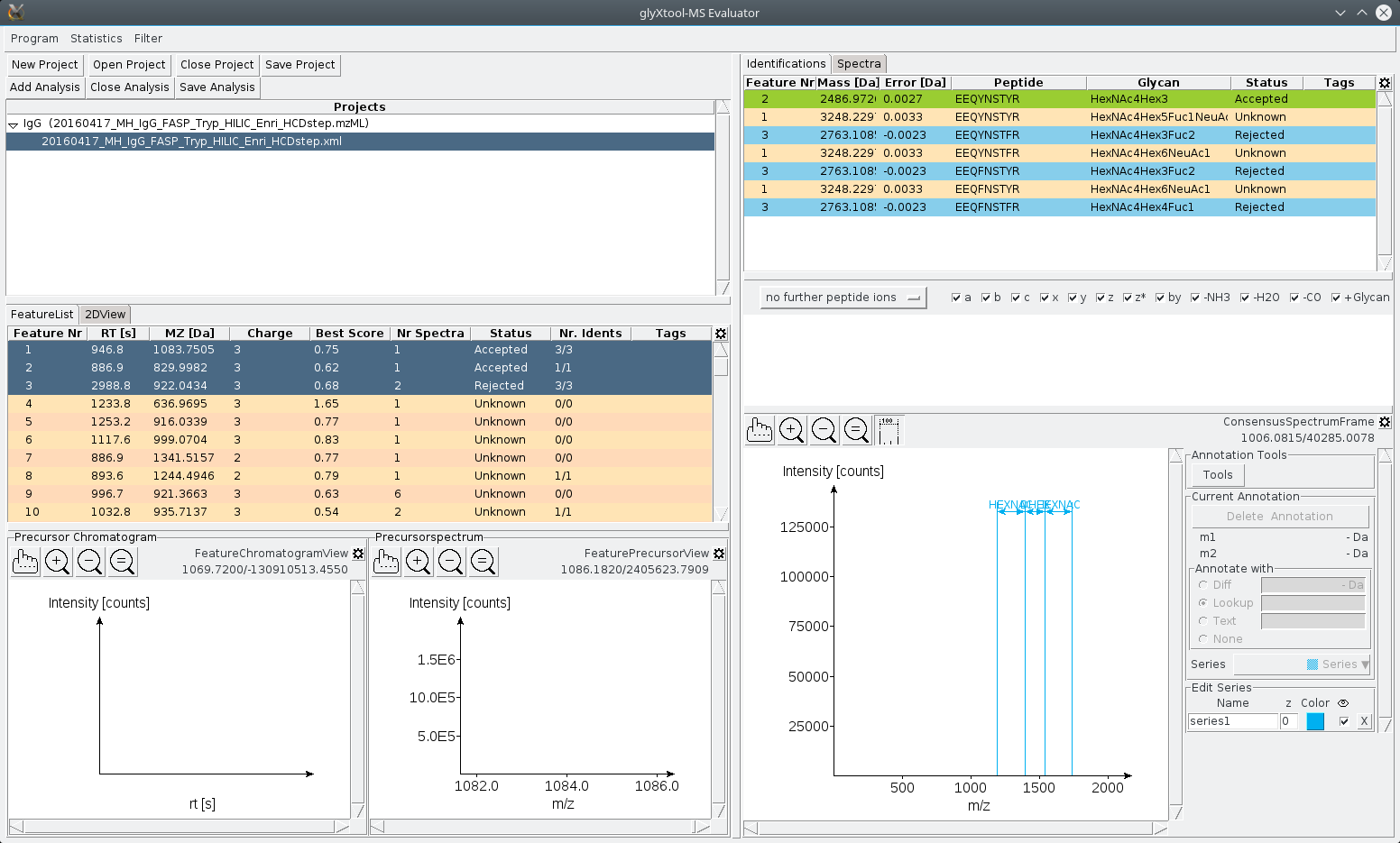


Figure 20: Multiselection of several features displays all corrsponsing identifications within the “Identifications” tab

# TOPPAS tools for glycopeptide analytics

Here, the purpose of each tool and its possible predecessor tools and successor tools are described.

## FeatureFinderMS

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| pot. predecessor tools | → | FeatureFinderMS | → | pot. successor tools |
| MS centroid data in .mzML format  PeakPickerHighRes  PeakPickerWavelet  FileFilter | glyxFilter |

**Purpose**

Finds features around analytes containing at least one fragment spectrum.

**Parameters**

* inMZML: Input mass spectra as centroid data in \*.mzML file format
* outFeature: Feature output file
* tolerance: Mass tolerance in Dalton
* mswindow: maximum mass range of the precursor isotope pattern in dalton
* precursorshift: maximum deviation of the precursor mass from the (average) precursor mass reported within the mass file in dalton
* rtwindow: maximum elution range of the analyte peak in seconds

**Possible Input Nodes**

The tool uses centroided MS1 data. Possible input nodes are the file input node, the various OpenMS Peakpicker nodes or the FileFilter node if the data have to be cropped to a certain elution or mass range

**Possible Output Nodes**

* glyxFilter

**Similar tools**

* FeatureFinderCentroided
* FeatureFinderisotopeWavelet

## FileBuilder

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| pot. predecessor tools | → | FileBuilder | → | pot. successor tools |
| FileFilter  PeakPicker | glyxFilter |
| FileFilter  PeakPicker |

**Purpose**

Replaces the given MS Level spectra in an experiment. In the context of glycopeptide analysis it is used to replace continuous MS2 fragment spectra with their centroided counterpart after peakpicking, while retaining continuous data in the MS1 domain. This is needed as input for the ‘glyXtoolMS Evaluator’ to visualize continuous MS1 data for the precursors.

**Parameters**

* inOriginal: File input of mass spectrometry data in \*.mzML format; All MS level are transferred to the output file except the level provided by the option ‘MSLevel’
* inReplace: File input of mass spectrometry data in \*.mzML format; The spectra matching the given MSLevel option are transferred to the output file
* out: File output in \*.mzML format
* MSLevel: MS level which will be replaced with data from the replacement file

**Possible Input Nodes**

* FileFilter
* PeakPicker
* File input node

**Possible Output Nodes**

* glyXfiler
* File output node

## glyXFilter

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| pot. predecessor tools | → | glyxFilter | → | pot. successor tools |
| FeatureFinderCentroided  FeatureFinderMS\*  FeatureFinderIsotopeWavelt | glyxReporter  Glycopeptide Matcher\*  File Output node (.xml) |
| MS centroid data in .mzML format  PeakPickerHighRes  PeakPickerWavelet  FileFilter  FileBuilder\* |

**Purpose**

The tool searches for glycopeptide evidence in MS2 spectra, based on oxonium ions and neutral losses from the precursor. Reported is a spectrum score between 0.0 and 10.0 for each MS2 spectrum where the lower score signifies a higher glycopeptide probability. The identified glycopeptide fragment spectra are then used to identify glycopeptide features in the FeatureMap. For easier data access in later stages of the analysis pipeline the tool then generates consenus spectra for all identified glycopeptide features. All generated information is finally stored in a \*.xml file.

**Parameters**

* inMZML: Input mass spectra as centroid data in \*.mzML file format
* inFeature: Input feature file as\*.featureXML
* outGlyML: Output file in \*.xml format, containing all scored fragment spectra and all identified glycopeptide features
* createFeatures: (false/true);when no feature could be found within the provided feature map for a given fragment spectra a dummy feature will be generated, if set to true
* hasFucose: (false/true); if true use predefined oxonium ions that contain fucose
* hasNANA: (false/true); if true use predefined oxonium ions that contain N-acetylneuraminic acid
* hasNGNA: (false/true); if true use predefined oxonium ions that contain N-glycolylneuraminic acid
* oxoniumions: Add additional oxonium ions to the search.   
  Format has to be like: (NeuAc)1(H2O)-1(H+)1 with comma separated oxonium ions
* tolerance: Mass tolerance for the oxonium ion search
* toleranceType: Mass tolerance Type (either ppm or Da)
* ionthreshold: Ignores peaks with lower intensity than the given threshold. Set to 0 to include all peaks.
* scorethreshold: Threshold used to identify a fragment spectrum as a glycopeptide. Lower scores signify a higher glycopeptide probability.

**Possible Input Nodes**

* inMZML: needed are centroided MS2 data with sorted peaks after mass. Suitable is the FileBuilder, to generate a suitable input file for the glyXtool Evaulator; PeakPicker or FileFilter if the MS2 data are already centroid data and need to be sorted
* inFeature: All possible FeatureFinder tools

**Possible Output Nodes**

* glyxReporter:
* glycoPeptideMatcher: for matching peptide and glycan composition to the precursor masses of identified glycopeptide features

## GlycopeptideDigest

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| pot. predecessor tools | → | Glycopeptide Digest | → | pot. successor tools |
| Protein sequences in \*.fasta format | GlycopeptideMatcher |

**Purpose**

Generating possible peptide sequences with glycosylation sites from protein sequences via theoretical digest.

**Parameters**

* inFasta: Input file in \*.fasta format containing either protein sequences or peptide squences
* out: \*.xml file containing the generated peptides with glycosylation sites, their possible modifications and the monoisotopic mass of each peptide
* enzymes: The enzyme(s) used for the digest. Currently supported are trypsin, trypsin/P, AspN, Unspecific and NoDigest. The option ‘Unspecific’ cuts after each aminoacid and uses the Nr of missedCleavageSites as the maximum length of the reported peptides. With the option ‘NoDigest’ the provided sequences from the \*.fasta file are used without digest, allowing the user to specify peptides.
* maxNrModifications: Nr of maximum allowed modifications on each peptide. CYS\_CAM and CYS\_CM are excluded.
* modifications: Variable modifications. For each peptide all possible permutations are generated. If e.g. a peptide contains two methionines, three peptides are generated: (0 Oxidations, 1 Oxidation on either methionine and fully oxidized on both residues)
* glycosylation: (N-glycosylation, O-glycosylation). Select which glycosylation site should be checked. Uses the motif N(S|T)(^P) as consensus sequence for N-glycosylation and (S|T) for O-glycosylation
* missedCleavageSites: maximum nr of missed cleavage sites. In case of unspecific digest determines the maximum length of the peptide.

**Possible Input Nodes**

* Input node with \*.fasta file

**Possible Output Nodes**

* GlycopeptideMatcher

## GlycanComposition builder

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| pot. predecessor tools | → | Glycan Composition Builder | → | pot. successor tools |
| File with glycan compositions  Or no file node | GlycopeptideMatcher |

**Purpose**

Provides glycan compositions for the ‘Glycopeptide Matcher’ tool. A given list of glycan compositions can be filtered by the provided ranges if the ‘useAsFilter’ option is set to true, otherwise a list of glycan compositions is calculated in-silico with the given ranges.

**Parameters**

* in: File input
* out: Output file, containing the filtered glycan compostions in an \*.txt file
* useAsFilter: (false/true); If true, filters the glycan compositions from the input file according to the provided monomer ranges. If false disregards content of the input file and calculates all compositon permutations from the given monomer ranges.
* rangeHex: range of hexose within the glycan composition
* rangeHexNAc: range of N-acetylhexoseamine within the glycan composition
* rangeFuc: range of fucose within the glycan composition
* rangeNeuAc: range of N-acetylneuraminic acid within the glycan composition
* rangeNeuGc: range of N-glycolylneuraminic acid within the glycan composition

**Possible Input Nodes**

* File input node. In case the ‘useAsFilter’ is set to false, the content of the file input node is neglected, since the glycan compostion permutations are calculated based on the given ranges. However TOPPAS expects each tool to have an input node to run, thus some file has to be provided to the tool.

**Possible Output Nodes**

* Glycopeptide Matcher

## Glycopeptide Matcher

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| pot. predecessor tools | → | Glycopeptide Matcher | → | pot. successor tools |
| glyxFilter\* | glyxReporter\*  ConsensusSearch  File Output node (.xml) |
| Glycopeptide Digest\* |
| Glycan Composition Builder  File with glycan compositions |

**Purpose**

Matches a given list of peptides and glycan compositions to precursor masses of glycopeptide features.

**Parameters**

* out: Output file, Appends new collected information to the given inAnalysis file
* inAnalysis: Input file containing a glyML analysis file with scored glycopeptide features
* inGlycan: Input file containing a list of glycan compositions to match against
* inPeptide: Input file containing a list of peptide to match against
* tolerance: Mass tolerance for the oxonium ion search
* toleranceType: Mass tolerance Type (either ppm or Da)

**Possible Input Nodes**

1. glyxFilter
2. GlycopeptideDigest
3. Glycan Composition Builder

File with glycan compositions

**Possible Output Nodes**

* glyxReporter
* ConsensusSearch

## Consensus Search

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| pot. predecessor tools | → | Consensus Search | → | pot. successor tools |
| Glycopeptide Matcher | glyxReporter\*  File Output node (.xml) |

**Purpose**

Annotates the consensus spectra of glycopeptide features with peptide fragments based on the theoretical fragments of the peptide sequence suggested by the Glycopeptide Matcher tool.

**Parameters**

* inGlyML: Input analysis file in glyML format
* outGlyML: Output analysis file in glyML format
* tolerance: Mass tolerance for the oxonium ion search
* toleranceType: Mass tolerance Type (either ppm or Da)
* ionthreshold: Intensity threshold for annotating fragment spectra peaks. Set to Zero to ignore intensity.
* pepIons: List of peptide ions to search for

**Possible Input Nodes**

* Glycopeptide Matcher

**Possible Output Nodes**

* glyxReporter

## glyxReporter

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| pot. predecessor tools | → | glyxReporter | → | pot. successor tools |
| glyxFilter  Glycopeptide Matcher  Fragment Search | File Output node (.xls) |

**Purpose**

Converts the collected information stored in the glyML file from the glycopeptide analysis tools into excel sheets.

**Parameters**

* inAnalysis: Input file in glyML format
* outReport: Output as \*.xls file

**Possible Input Nodes**

* glyxFilter
* Glycopeptide Matcher
* Fragment Search

**Possible Output Nodes**

* File output (\*.xls)