Documented Halomonas Expressed Protein Index, Quantified Protein Profile and Spectral Library Producing Makefile

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Abstract

Documentation of a makefile to reproduce protein index, protein quantification and spectral libraries for *Halomonas Bisphagenesis* TD01. A complete commentary on the accompanying makefile script to enable reproduction of libraries from raw mass spectrometry data files.

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Introduction

This document describes a data processing pipeline which generates data reported in paper “Baseline Proteomics Characterisation of Biomanufacturing Organism Halomonas Bluephagenesis”. The pipeline is scripted in gnu-make and describes every process between raw data and the package submitted to PRIDE repository (https://doi.org/10.6019/PXD028156).

Introduction to Pipeline

Data was acquired as MS in which the instrument alternates between a low energy scan and a collision energy “ramp”. The low energy scan captures precursor ions, the ramp captures product ion spectra. Data of this kind is not supported by opensource tools such as openms or trans-proteomics pipeline for spectra generation. So the Waters proprietary tools are used.

The proprietary tools that manage the pipeline as follows.

- Mass spectrometry data files are converted to feature lists by apex3d.exe.
- Precursor and product ions are matched by elution profile and combined into conventional spectra peak lists by peptide3d64.exe.
- Spectra are searched against a fasta database by iadb.exe.

It is possible to output additional information at each stage of the process. For example apex3d.exe is able to output a .csv file of all identified features and peptide3d64.exe can output a conventional .mgf file which may be submitted to other search engines. The final output of iadb.exe is a set of three .csv files summarising the outcome at a protein, peptide and fragment level.

After peakpicking data is searched and processed along two separate tracks. Protein expression is quantified using the “top three” method incorporated into Waters “iadb” search. Spectra obtained from the peptide3d64.exe are separately extracted as .mgf files and processed through the X!Tandem search engine and trans-proteomics-pathway to produce spectral libraries.

Introduction to Make

The analysis pipeline is executed using gun-make also called make. The program make is well documented(https://www.gnu.org/software/make/manual). Make is principally used to compile software but is also useful to document bioinformatics pipelines. Make is a useful tool for scripting the “many-to-many” operations in which multiple mass spectrometry data files are each processed into a series of intermediate files followed by “many-to-one” operations where multiple intermediate files are combined into a summary output. Make facilitates writing commands for each operation in a single place for application to multiple files which ensures consistency and facilitates modification and experimentation with the pipeline as a whole.

The principle of make is that the program is supplied with a set of “rules” for converting “prerequisites” into “targets” by means of a “recipe”. In this case the targets are a set of spectral libraries and the source files are MS datasets, fasta protein sequence libraries etc. Recipes take the general form:

```
target : prerequisite
  program -in prerequisite -out target
```

Recipes can be written in any order and make is able to chain them together to convert source files to target files via intermediate files. The following make file has been written with recipes in the sequence in which they would be run through the pipeline. When executed; make will parallelise the processing to speed up job completion.

Make is able to substitute text for macros which are defined myMacro:=text or myMacroList:=element_01 element_02 and then called ${myMacro} or ${myMacroList}. It also has a set of built in functions that are
called in the general form: $(function_name \ variable, param, param...)$ which are able to manipulate variables, for example to change a file suffix, or change or remove a directory path. In this way it is possible to write rules convert raw data to intermediate files through an analysis pipeline without explicitly Terminal writing out all intermediate file names. Each recipe used for this project are briefly described and links to documentation supplied as encountered in the make file.

**System Setup**

This section describes the programs that must be installed in order to run this pipeline.

**PC Setup**

This pipeline was developed on a PC with specifications:

- **Processor** Intel i7-7820X 3.6GHz 8 cores.
- **RAM** 128 GB
- **Operating System** Windows 10 2004 Build 19041.1165

Windows is required for Waters’ proprietary tools. The **peptide3d** program and process consumes a lot of RAM, script detailed below prevents more than one instance of **peptide3d** running simultaneously to prevent the system running out of RAM. It is possible the pipeline will fail if run on a system with substantially less RAM than was used in development.

**Install All Required Programs**

1. If Waters executables (**apex3d**, **peptide3d**, **merge** and **iadb**) outlined above are not otherwise available download Progenesis QI for Proteomics and install, as this software installs the required executables. Here the executables from Waters’ PLGS
2. Download Rtools and install, this includes gnu tools **sed**, **awk**, **grep**, etc.
3. If not present on your system download and install Trans-Proteomic Pipeline (TPP) this work was tested with v6.0.0.
4. If not present on your system download and install x!tandem this work was tested with version ALANINE (2017.02.01). As an alternative the version of x!tandem distributed with the Trans-Proteomic Pipeline (TPP) may be used, however at present this is the older Jackhammer TPP (2013.06.15.1 - LabKey, Insilicos, ISB) version which was not used.
5. If not present on your system download and install seqkit this work was tested with v0.12.0.
6. If not present on the system download and install openMS this work was tested with v2.4.0.
7. A python distribution is required to run the msproteomicstools package. This work was tested with miniconda for python 3.7 with msproteomicstools installed as per the instructions given on the website. The minimal python installation enables just the requirements for this process to be installed with the versions required to replicate this work. Miniconda may be installed alongside other instances of python allowing this work to be repeated without disrupting a pre-existing system. During installation note the target directory and adjust if required. For single user installation this will look like: `C:\Users\<user>\AppData\Local\Continuum\miniconda3`, for all users it may be `C:\ProgramData\Miniconda3`. Note `C:\ProgramData\` is a hidden directory on windows 10, navigate to it directly by entering path into windows explorer or set windows explorer to show hidden files.
8. Install required python packages. To access the python command line, open a windows command window; navigate to the **Miniconda3\Scripts** directory; and type **activate** into the command line. Required packages will need to be installed from pip.org through the python command line. The following commands entered into the anaconda prompt should install the required packages in the
versions used here. Alternatively the latest versions of these programs might be installed although that might require adjustments to processing commands below.

```
pip install numpy==1.15.3
pip install pymzml==0.7.5
pip install Biopython==1.72
pip install Cython==0.29.2 --install-option="--no-cython-compile"
pip install msproteomicstools==0.8.0
```

No further interaction with the python environment should be required.

**Executing Make**

Gnu-make is executed from the command line. It is easiest to start a new cmd window in the project directory, set the PATH variable to just the location of gnu-make and then execute gnu-make on the script:

```
PATH="C:\Program Files\gmake"
gnumake-4.3-x64.exe -rR -j 8 -l 8 -f ./IntegratedAnalysis.mk --output-sync all
```

The command line is documented on the gmake manual. Briefly: -rR turns off internal rules for compiling software; -j indicates the number of concurrent jobs permitted, select a lower number if limited processors or RAM available; -l checks load is below value specified before starting a new job, this provides some protection against overloading RAM which causes Waters programs to crash; -f indicates the file name of the makescript to run; --output-sync puts the output of each job on the console after the job finishes, if not set the output of concurrently running tasks will be output concurrently and consequently unintelligible; all is the name of the “target” required by this build, alternatives are available for testing, see below.

**Documented Makefile for Pipeline**

The following code blocks are combined into a single gnu-make script to run the pipeline.

**Set Shell**

Each recipe in the make file is run in a “shell”, that is a separate command line environment. On windows there is a choice of shell. This script used windows native executables from Waters to process the data and so the windows cmd.exe shell was chosen. It is likely that the script could be rewritten to run in the Windows Subsystem for Linux if required. However in this case all the dos commands such as copy and ren could need to be replaced with unix alternatives and paths changed to unix format.

```
SHELL=C:/Windows/System32/cmd.exe
```

Paths to Executables

Explicit control of software versions is assured by assigning the full path of each executable to a macro for each of the programs required for the pipeline. These must be edited to be correct for the system on which
the make script is to be run. Setting them explicity should also support reproducability since the version of software used here may be installed along side subsequent releases of the software being used in future active research. Checking these are correct also acts as a check list to ensure system is set up correctly.

# Paths to the Waters executables
apex_path="C:\PLGS3.0.3\lib\apex3d\Apex3D64.exe"
peptide3d_path="C:\PLGS3.0.3\lib\apex3d\Peptide3D.exe"
IADB_path="C:\PLGS3.0.3\bin\iaDBs.exe"
mergeFractions_path="C:\PLGS3.0.3\bin\MergeFractions.exe"

# Paths to the sed and awk (gawk) used to process text files during library build
sed_path="C:\Rtools\bin\sed.exe"
awk_path="C:\Rtools\bin\gawk.exe"

# Path to 7 zip to zip data files for deposition.
zip7_path="C:\Program Files\7-Zip\7z.exe"

# Paths to spectrast etc
spectrast_path=\"C:\Program Files\OpenMS-2.4.0\share\OpenMS\THIRDPARTY\SpectraST\spectrast.exe\"
python_path="C:\ProgramData\Miniconda3\python.exe"
spectrast2tsv_path="C:\ProgramData\Miniconda3\Scripts\spectrast2tsv.py"
TargetedFileConverter_path="C:\Program Files\OpenMS-2.4.0\bin\TargetedFileConverter.exe"
perl_path="C:\TPP\perl\bin\perl.exe"
mayu_path="C:\TPP\bin\Mayu.pl"

# Paths to additional bioinformatics programs used in this analysis
DecoyDatabase_path="C:\Program Files\OpenMS-2.4.0\bin\DecoyDatabase.exe"
seqkit_path="C:\Program Files\seqKit_v0.12.0\seqkit.exe"

### TPP bits
Tandem2XML_path="C:\TPP\bin\Tandem2XML.exe"
xinteract_path="C:\TPP\bin\xinteract.exe"
PeptideProphet_path="C:\TPP\bin\PeptideProphetParser.exe"
perl_path="C:\TPP\perl\bin\perl.exe"
tandem_path="C:\Program Files\tandem-win-17-02-01-4\bin\tandem.exe"
msconvert_path="C:\Program Files\ProteoWizard\ProteoWizard 3.0.20002.36ad78f14\msconvert.exe"
Peptide3D2MGF_path="C:\PROGRA~1\Waters\Symphony\utils\PEPTID-1.EXE"

### Powershell
Powershell_path="C:\Windows\System32\WindowsPowerShell\v1.0\powershell.exe"

Macros

In make macros are rules to generate a required instructions prior to execution. They may be used to generate output file names from an input file name, or vice versa. They may be used to generate entire recipe comands.

The following macro tests to see if required target directories exist and creates it if not. It is used at the start of most recipes to ensure the target directory is available.

define makeDir
if not exist $(subst /,\,,$(abspath $(dir ${1})))) mkdir $(subst /,\,,$(abspath $(dir ${1}))))
The following two macros turn relative file paths into absolute file paths, this allows relative paths to facilitate portability of the script but absolute paths to be provided to programs which require them. They also convert between unix-like forward slash “/” and windows like back slash “\” for those that need them.

```bash
define winify
$(subst /,\,$(abspath ${1}))
endef

define unixify
$(subst \,/,$(abspath ${1}))
endef
```

The following pair of macros between them find the position of a string within a gmake word list. The slightly esoteric formulation is a recursive macro _pos, which nibbles through a word list until it removes the matching element combined with pos which counts the remaining elements. The code enables the pep3d process to be run one file at a time as run concurrently multiple pep3d processes will consume all the RAM causing the processes to crash.

```bash
_pos = $(if $(findstring $1,$2),$(call _pos,$1,$(wordlist 2,$(words $2),$2),x $3),$3)
pos = $(words $(call _pos,$1,$2))
```

The following macro calls powershell to obtain a UUID. This is used to pass a UUID to the IADB search.

```bash
UUID=$(shell ${Powershell_path} -Command "[guid]::NewGuid().ToString()"
```

### Mass Spectrometry Raw Data Files

The raw data acquired on the Xevo has been stored on a local hard drive.

Waters data are stored as multiple files inside a folder. Make can’t treat a directory as a pre-requisite so each file is specified as the _HEADER.TXT file held in each of the directories and this element is removed when passed to the apex3d program.

The raw data file paths are concatenated into a single list of files so they can all be separately processed by a single command. This ensures consistency in data processing. Raw data directory paths will need to be adjusted to the location of downloaded raw data for re-use.

```bash
rawDirComps:=_CHRO001.DAT _CHRO002.DAT _CHRO003.DAT _CHRO004.DAT _CHRO005.DAT \
_CHROMS.INF _extern.inf _FUNC001.DAT _FUNC001.IDX _FUNC001.STS \
 FUNC002.DAT _FUNC002.IDX _FUNC002.STS _FUNC003.DAT _FUNC003.IDX \
 FUNC003.STS _FUNCTNS.INF _HEADER.TXT _INLET.INF

rawDirCompsNoDat:=_CHROMS.INF _extern.inf _FUNC001.IDX _FUNC001.STS \
_FUNC002.IDX _FUNC002.STS _FUNC003.IDX \
_FUNC003.STS _FUNCTNS.INF _HEADER.TXT _INLET.INF

rawDirEcoli_wcl=E:/massSpecData/xevo_FBRH_MR_Pro/Data/20_02_03_EcoliFractions/

rawFilesEcoli_wcl:= Ecoli_F01.raw Ecoli_F02_300ng.raw Ecoli_F03_300ng.raw \
 Ecoli_F04_300ng.raw Ecoli_F05_300ng.raw \
 Ecoli_F06_300ng.raw Ecoli_F07_300ng.raw \
 Ecoli_F08_300ng.raw Ecoli_F09_300ng.raw \
 Ecoli_F10_300ng.raw Ecoli_F11_300ng.raw \
```
Ecoli_F12_300ng.raw Ecoli_F13_300ng.raw \
Ecoli_F14_300ng.raw Ecoli_F15_300ng.raw \
Ecoli_F16_300ng.raw Ecoli_F17_300ng.raw \
Ecoli_F18_300ng.raw Ecoli_F19_300ng.raw \
Ecoli_F20_300ng.raw Ecoli_F21_300ng.raw \
Ecoli_F22_300ng.raw Ecoli_F25_300ng.raw \
Ecoli_F26_300ng.raw Ecoli_F27_300ng.raw \
Ecoli_F28_300ng.raw Ecoli_F29_300ng.raw \
Ecoli_F30_300ng.raw Ecoli_F31_300ng.raw

# Ecoli_F23_300ng.raw Ecoli_F24_300ng.raw

rawDirHalomonas_wcl=E:/massSpecData/xevo_FBRH_MR_Pro/Data/20_02_12_HaloFracs/

rawFilesHalo_wcl:= 05_Halo_F1.raw 06_Halo_F2.raw 07_Halo_F3.raw \
08_Halo_F4.raw 09_Halo_F5.raw 10_Halo_F6.raw \
11_Halo_F7.raw 12_Halo_F8.raw 13_Halo_F9.raw \
14_Halo_F10.raw 15_Halo_F11.raw 16_Halo_F12.raw \
17_Halo_F13.raw 18_Halo_F14.raw 19_Halo_F15.raw \
20_Halo_F16.raw 21_Halo_F17.raw 22_Halo_F18.raw \
23_Halo_F19.raw 25_Halo_F21.raw 26_Halo_F22.raw \
27_Halo_F23.raw 28_Halo_F24.raw 29_Halo_F25.raw \
30_Halo_F26.raw 31_Halo_F27.raw

rawFilesEcoliWCLPaths:=$(addsuffix /_HEADER.TXT, $(addprefix ${rawDirEcoli_wcl}, ${rawFilesEcoli_wcl}))
rawFilesHaloWCLPaths:=$(addsuffix /_HEADER.TXT, $(addprefix ${rawDirHalomonas_wcl}, ${rawFilesHalo_wcl}))

rawFiles_All_wcl:=${rawFilesEcoliWCLPaths} ${rawFilesHaloWCLPaths}

############
# Edit for secretome

rawCellFracDir_1:=E:/massSpecData/xevo_FBRH_MR_Pro/Data/20_08_26_Hal_Ecoli_CellFracs/
rawCellFracDir_2:=E:/massSpecData/xevo_FBRH_MR_Pro/Data/21_02_19_MemPeri/

## Membrane

rawEcoli_Periplasm_Fracs_1:= mbpssmr9_01_EP_MSe_2uL.raw mbpssmr9_06_EP_MSe_2uL.raw mbpssmr9_11_EP_MSe_2x.raw
rawHalo_Periplasm_Fracs_1:= mbpssmr9_03_HP_MSe_2uL.raw mbpssmr9_08_HP_MSe_2uL.raw mbpssmr9_13_HP_MSe_2x.raw

rawEcoli_Periplasm_Fracs_2:= mbpssmr9_EP.raw

rawFilesEcoliPeriPaths:=$(addsuffix /_HEADER.TXT, $(addprefix ${rawCellFracDir_1}, ${rawEcoli_Periplasm_Fracs_1})) \
$(addsuffix /_HEADER.TXT, $(addprefix ${rawCellFracDir_2}, ${rawEcoli_Periplasm_Fracs_2}))
rawFilesHaloPeriPaths:=$(addsuffix /_HEADER.TXT, $(addprefix ${rawCellFracDir_1}, ${rawHalo_Periplasm_Fracs_1})) \
$(addsuffix /_HEADER.TXT, $(addprefix ${rawCellFracDir_2}, ${rawHalo_Periplasm_Fracs_2}))
rawFiles_All_peri:=${rawFilesEcoliPeriPaths} ${rawFilesHaloPeriPaths}

### Edit for secretome

rawSereteEcoliDir:=E:/massSpecData/xevo_FBRH_MR_Pro/Data/21_01_29_Secretome/

rawCellFracDir:=E:/massSpecData/xevo_FBRH_MR_Pro/Data/21_02_19_MemPeri/

rawEcoliSec:= mbpssmr9_ecoli_secretome_2uL_02.raw \ 
            mbpssmr9_ecoli_secretome_5uL_01.raw \ 
            mbpssmr9_ecoli_secretome_8uL_03.raw \ 
            mbpssmr9_ecoli_secretome_highConc_5uL_01.raw

rawHaloSec:= 21_02_19_SecretomeHalo_10uL.raw 21_02_19_02_SecretomeHalo.raw

rawFilesEcoliSecPaths:=$(addsuffix /_HEADER.TXT, $(addprefix ${rawSereteEcoliDir}, ${rawEcoliSec}))

rawFilesHaloSecPaths:=$(addsuffix /_HEADER.TXT, $(addprefix ${rawCellFracDir}, ${rawHaloSec}))

rawFiles_All_sec:=${rawFilesEcoliSecPaths} ${rawFilesHaloSecPaths}

With all the raw data specified they are grouped for convenience into lists of all files, and of each bacteria species separatly.

rawFiles_All:=${rawFilesEcoliSecPaths} ${rawFilesHaloSecPaths} ${rawFilesEcoliPeriPaths} ${rawFilesHaloPeriPaths} ${rawFilesEcoliWCLPaths} ${rawFilesHaloWCLPaths}

rawFiles_Ecoli:=${rawFilesEcoliSecPaths} ${rawFilesEcoliPeriPaths} ${rawFilesEcoliWCLPaths}

rawFiles_Halo:=${rawFilesHaloSecPaths} ${rawFilesHaloPeriPaths} ${rawFilesHaloWCLPaths}

rawDir:=${rawFilesEcoli_wcl} ${rawFilesHalo_wcl} ${rawEcoli_Periplasm_Fracs_1} ${rawHalo_Periplasm_Fracs_1} ${rawEcoli_Periplasm_Fracs_2} ${rawEcoliSec} ${rawHaloSec}

### Intermediate Files

Lists of intermediate files are useful for setting put the rules for converting intermediate files. Here the names of the required intermediate files are determined from the raw data file names. Apex3d output files are put into sub dir apex, peptide3d to sub dir pep3d and iadb files to subdir iadb. The useful outputs of iadb, the tables of protein, peptide and fragment level summary data, are output to sub dir iadb\tables although that is handled in the recipes.

apexDir:=./apex/
pep3dDir:=./pep3d/
pep3dMergedDir:=./pep3d/merge/
iadbDir:=./iadb/
iadbTabDir:=./iadb/tables/
TPP:=./tpp_mayu/
# Apex paths:
apexPaths:=$(addprefix ${apexDir},${notdir $(rawFiles_All:.raw/_HEADER.TXT=_Apex3D.bin)})
#$info first apex $(firstword $apexPaths))

# Apex paths WCL:
apexPaths_wcl:= $(addprefix ${apexDir},${notdir $(rawFiles_All_wcl:.raw/_HEADER.TXT=_Apex3D.bin)})

# Apex paths periplasm:
apexPaths_peri:=$(addprefix ${apexDir},${notdir $(rawFiles_All_peri:.raw/_HEADER.TXT=_Apex3D.bin)})

# Apex paths periplasm:
apexPaths_sec:=$(addprefix ${apexDir},${notdir $(rawFiles_All_sec:.raw/_HEADER.TXT=_Apex3D.bin)})

# Pep3d Paths:
pep3dPaths:=$(addprefix ${pep3dDir},${notdir $(rawFiles_All:.raw/_HEADER.TXT=_Pep3D_Spectrum.bin)})
pep3dAll:=${pep3dPaths} ${pep3dMergedFiles}

# Merged pep3dFiles
pep3dMergedFiles := ./pep3d/merged_halo_wcl_Pep3D_Spectrum.bin \
    ./pep3d/merged_ecoli_wcl_Pep3D_Spectrum.bin

# IADB paths
## Merged files:
iadb_mg_bin:=$(abspath $(addprefix ${iadbDir},${notdir $(pep3dMergedFiles:_Pep3D_Spectrum.bin=_IA_workflow.bin)}))
iadb_mg_tabFrag:=$(abspath $(addprefix ${iadbTabDir},${notdir $(pep3dMergedFiles:_Pep3D_Spectrum.bin=_IA_final_fragment.csv)}))
iadb_mg_tabPep:=$(abspath $(addprefix ${iadbTabDir},${notdir $(pep3dMergedFiles:_Pep3D_Spectrum.bin=_IA_final_peptide.csv)}))
iadb_mg_tabProt:=$(abspath $(addprefix ${iadbTabDir},${notdir $(pep3dMergedFiles:_Pep3D_Spectrum.bin=_IA_final_protein.csv)}))
iadb_all:=$(sort ${iadb_mg_bin} ${iadb_mg_tabFrag} ${iadb_mg_tabPep} ${iadb_mg_tabProt})

# mzXML
pep3dEcoliPaths_mzXML:=$(addprefix ./pep3d/,$(notdir $(rawFiles_Ecoli:.raw/_HEADER.TXT=_Pep3D_Spectrum.mzXML)))
pep3dHaloPaths_mzXML:=$(addprefix ./pep3d/,$(notdir $(rawFiles_Halo:.raw/_HEADER.TXT=_Pep3D_Spectrum.mzXML)))
al_mzXML:=${pep3dEcoliPaths_mzXML} ${pep3dHaloPaths_mzXML}

# x!tandem
all_tandem: = $(al_mzXML:.mzXML=.t.xml)

# pepXML
pep3dEcoliPaths_pepXML:=$(addprefix ./pep3d/,$(notdir $(rawFiles_Ecoli:.raw/_HEADER.TXT=_Pep3D_Spectrum.pep.xml)))
pep3dHaloPaths_pepXML:=$(addprefix ./pep3d/,$(notdir $(rawFiles_Halo:.raw/_HEADER.TXT=_Pep3D_Spectrum.pep.xml)))
al_pepXML:=${pep3dEcoliPaths_pepXML} ${pep3dHaloPaths_pepXML}

# mzid
all_mzid:=${al_mzXML:.mzXML=.t.xml.mzid)

# interact
all_interact:=./pep3d/interact_ecoli_all.pep.xml ./pep3d/interact_halo_all.pep.xml

# prophet
all_prophet:=./pep3d/interact_all_ecoli_prophet.pep.xml ./pep3d/interact_all_halo_prophet.pep.xml
# mayuFiles
mayuOut:= ./tpp_mayu/mayuOut_all_halo_psm_protFDR0.05_t_1.07.csv \
./tpp_mayu/mayuOut_all_ecoli_psm_protFDR0.05_t_1.07.csv \
./tpp_mayu/mayuOut_all_halo_main_1.07.csv \
./tpp_mayu/mayuOut_all_ecoli_main_1.07.csv \
./tpp_mayu/mayuOut_all_ecoli_plot_psm_protFDR0.04_t_1.07.csv \ 
./tpp_mayu/mayuOut_all_ecoli_plot_main_1.07.csv \
./tpp_mayu/mayuOut_all_halo_plot_psm_protFDR0.04_t_1.07.csv \ 
./tpp_mayu/mayuOut_all_halo_plot_main_1.07.csv

The spectral libraries output in various formats defined here:

nonConsLib:= ./tpp_mayu/ecoli_all.splib ./tpp_mayu/halo_all.splib
ConsLib:= ./tpp_mayu/ecoli_all_cons.splib ./tpp_mayu/halo_all_cons.splib ./tpp_mayu/ecoli_SWATHatlas2020.splib
TsvLib_pk:=$(ConsLib:.splib=_pk.tsv)
TsvLib_os:=$(ConsLib:.splib=_os.tsv)
targetSplib:= ${nonConsLib} ${ConsLib} ${TsvLib_pk} ${TsvLib_os}
uploadSplib:= ${nonConsLib} $(nonConsLib:.splib=.sptxt) ${TsvLib_pk} ${TsvLib_os}

Fasta Files
The .fasta protein database files are specified here.

Also specified are a pair of .fasta databases of common contaminants to draw-off likely contaminating spectra and a file containing a the peptides set as retention-time and quantitative standards concatenated into a single pseudo-protein.

# organism fasta files
fasta_ecoli=D:/fasta/Ecoli_MG1655_UP000000625.fasta
fasta_halo=./ini/halAndy/Halomonas_Hal1chrom.feature_protein.fasta

# iRT and contaminant fasta files
fasta_iRT=./ini/irtfusion_quant.fasta
fasta_iRT_single=./ini/irtfusionSingleProt.fasta
fasta_cRAP=./ini/crap.fasta
fasta_MPI=./ini/contaminants.fasta
fasta_NoDumCont=./ini/NoDup_cont.fasta

# derived fasta files used in the analysis
fastaForSearchEcoli=./ini/fastaForSearchEcoli.fasta
fastaForSearchHalo=./ini/fastaForSearchHalo.fasta
fastaForSearchEcoli_decoy=./ini/fastaForSearchEcoli_decoy.fasta
fastaForSearchHalo_decoy=./ini/fastaForSearchHalo_decoy.fasta
fastaForSpectrastEcoli=./ini/fastaForSpectrastEcoli.fasta
fastaForSpectrastHalo=./ini/fastaForSpectrastHalo.fasta
fastaForSearchHalo_xtand=./ini/fastaForSearchHalo_xtand.fasta
fastaForSearchEcoli_xtand=./ini/fastaForSearchEcoli_xtand.fasta
fastaForSearchHalo_xtand_decoy=./ini/fastaForSearchHalo_xtand_decoy.fasta
# List of fasta files to be included in the data submission

uploadFasta:= ${fastaForTandem} ${fastaForSearchEcoli} ${fastaForSearchHalo} ${fasta_halo} ${fasta_ecoli}

The specified *.fasta files are combined so that:

- The two contaminant databases are combined and given a common cont_ prefix to protein names.
- The contaminant and iRT databases are combined with the strain specific database for halomonas C3001.
- All annotation apart from accession number is stripped from the sequence annotation.

The *.fasta files are also converted to a table format for easy import and use in R.

## Specify Upload Packet

This section specifies the files which are to be uploaded to Pride and copies them to a specific folder for upload.

uploadDir:=D:/HaloLibForUpload/

```makefile
#uploadRaw_zip:=$(addsuffix /_HEADER.TXT,$(addprefix ${uploadDir},$(notdir $(rawFiles_All:.raw/_HEADER.TXT=.raw.zip))))
#uploadRaw:=$(foreach raw,${rawDir},$(addprefix ${uploadDir},$(foreach,comp,${rawDirComps},$(addsuffix ${comp},${raw}))))
#thisMakeFile:=$(lastword $(MAKEFILE_LIST))

#uploadRawDir:=$(foreach raw,${rawDir},$(addprefix ${uploadDir},$(raw:.raw=.raw.zip)))
uploadRaw:=$(foreach comp,${rawDirCompsNoDat},$(addsuffix /${comp},${uploadRawDir}))

# $(info allRawComp: ${uploadRaw})
```

intermediateFilesForUpload:= ${uploadFasta} ${uploadSplib} ${all_pepXML} ${all_interact} ${all_prophet} ${all_mzXML} ${thisMakeFile}

uploadFileList:=$(addprefix ${uploadDir},$(notdir ${intermediateFilesForUpload}))

upload_mzid:= $(addprefix ${uploadDir},$(notdir $(all_mzid:_Pep3D_Spectrum.t.xml.mzid=.mzid)))

uploadTargets := ${uploadFileList} ${upload_mzid} ${uploadRaw}

## Specify Targets

A critical part of a makefile is specification of the targets. Providing recipes exists to build intermediate and final target files from input files all these targets will be produced. The targets specified in all will be built. This is assured by making the special variable .PHONY dependent on all because all pre-requisites of .PHONY must always be rebuilt. The target .PHONY is the first target defined in the file and an invocation of make on the makefile will built that target if no other target is given on the command line. Other targets may be specified on the gnu-make invoking command line, the targets buildSet for example buildApex are provided for trouble shooting to enable each step of the process to be built in turn. The special target .ONESHELL is also set here to specify that all lines in a recipe be run in the same shell invocation rather than each in a
separate invocation. Running in a single shell is important for a few recipes where variables are set in one line and used in subsequent lines in the recipe.

```
allTargets := $(sort ${apexPaths}) ${pep3dPaths} ${iadb_all} ${targetSplib} ${mayuOut}
allUpload := ${uploadTargets} ${rawFiles_All_zip}

.PHONY: all \
buildMZID buildInteract buildProphet buildMayu buildSpplib buildUpload \
buildTandem buildApex buildPep3d testBuildApex testBuildPep3d testBuild_mzXML \
testBuildTandem buildIadb buildPep3dMerge buildUploadRaw buildUploadInt

all: ${mayuOut} ${allTargets} ${allUpload} ${spectrastTargets}

$(info all targets: $(sort ${allTargets} ${allUpload}))

.ONESHELL:

buildFasta : ${uploadFasta}
buildApex : ${apexPaths}
buildPep3d : ${pep3dPaths}
buildPep3dMerge : ${pep3dMergedFiles}
buildIadb : ${iadb_all}
buildTandem : ${all_tandem}
buildMZID : ${all_mzid}
buildInteract : ${all_interact}
buildProphet : ${all_prophet}
buildMayu : ${mayuOut}
buildSpplib : ${targetSplib}
buildUpload : ${uploadTargets}
buildUploadRaw : ${uploadRaw}
buildUploadInt : ${uploadFileList}

apexTestFile:=$(lastword ${apexPaths})
pep3dTestFile:=$(lastword ${pep3dPaths})
mzXMLTestFile:= ./pep3d/31_Halo_F27_Pep3D_Spectrum.mzXML
tandemTestFile:= ./pep3d/31_Halo_F27_Pep3D_Spectrum.t.xml

testBuildApex:${apexTestFile}
testBuildPep3d:${pep3dTestFile}
testBuild_mzXML:${mzXMLTestFile}
testBuildTandem:${tandemTestFile}
```

**Recipes**

**Generate Fasta Files**

```bash
decoy_string=reverse_
cont_string=cont_

$(fasta_ecoli:.fasta=_clean.fasta) : ${fasta_ecoli}
 ${sed_path} -e \'s/\[:blank:]/\n/g\' $< > $0
```
$(fasta_cRAP:.fasta=_clean.fasta) : ${fasta_cRAP}
    $(sed_path) -e 's/\'/>/\'$
    $< > $@

$(fasta_MPI:.fasta=_clean.fasta) : ${fasta_MPI}
    $(sed_path) -e 's/\'/>/\'$
    $< > $@

$(fasta_NoDumCont) : $(fasta_MPI:.fasta=_clean.fasta) $(fasta_cRAP:.fasta=_clean.fasta)
    type $(call winify, $\$)
    $(seqkit_path) rmdup --by-seq --dup-num-file ./ini/duplicatedCont.txt -o $(call winify, .\ini\tmp\$(notdir $\$))
    del $(call winify, .\ini\tmp\$(notdir $\$))

${fastaForSearchEcoli} : $(fasta_ecoli:.fasta=_clean.fasta) ${fasta_NoDumCont} ${fasta_iRT}
    type $(call winify, $\$)
    $(call winify, $@)

${fastaForSearchHalo} : ${fasta_halo} ${fasta_NoDumCont} ${fasta_iRT}
    type $(call winify, $\$)
    $(call winify, $@)

${fastaForSearchEcoli_xtand} : $(fasta_ecoli:.fasta=_clean.fasta) ${fasta_NoDumCont} ${fasta_iRT_single}
    type $(call winify, $\$)
    $(call winify, $@)

${fastaForSearchHalo_xtand} : ${fasta_halo} ${fasta_NoDumCont} ${fasta_iRT_single}
    type $(call winify, $\$)
    $(call winify, $@)

${fastaForSpectrastEcoli} : $(fasta_ecoli:.fasta=_clean.fasta) ${fasta_iRT_single}
    type $(call winify, $\$)
    $(call winify, $@)

${fastaForSpectrastHalo} : ${fasta_halo} ${fasta_iRT_single}
    type $(call winify, $\$)
    $(call winify, $@)

./ini/HaloFasta.tsv: ${fastaForSearchHalo}
    $(seqkit_path) fx2tab $< > $@

./ini/EcoliFasta.tsv: ${fastaForSearchEcoli}
    $(seqkit_path) fx2tab $< > $@

${fastaForSearchHalo_decoy} : ${fastaForSearchHalo}
    $(DecoyDatabase_path) -in $\$ -out $\$ -decoy_string ${decoy_string}

${fastaForSearchEcoli_decoy} : ${fastaForSearchEcoli}
    $(DecoyDatabase_path) -in $\$ -out $\$ -decoy_string ${decoy_string}
Set Apex, Peptide3d and Mayu Variables

It is possible to use makefile variables to set the values of switches on executables. For full executables for Apex and Peptide3d see Apex Feature Picking and Peptide3d Spectra Generation. However during pipeline refinement the following parameters to which the process is quite sensitive were taken out to this section for ease of adjustment.

**Apex Settings** The Apex3d program has many settings which are set below:

```bash
# Whole cell lysate APEX settings
bCSVOutput_wcl:=1
apex_stRT_wcl:=12
apex_endRT_wcl:=80
le3DThreshCounts_wcl:=10
he3DThreshCounts_wcl:=25
chromFWHMMin_wcl:=0.45
apex_binThresh_wcl:=650
apexTrackSNRThreshold_wcl:=2.25
driftFilterScaleFactor_wcl:=0.7
msFilterScaleFactor_wcl:=0.7
apexTrackFilterWide_wcl:=0
apexTrackSNRMeasFWHM_wcl:=1
minScansPerPeak_wcl:=7
msResolution_wcl:=20000
noFilteredIons_wcl:=1
bEnableTempCal_wcl:=1

# Periplasm apex settings
bCSVOutput_peri:=1
apex_stRT_peri:=12
apex_endRT_peri:=80
le3DThreshCounts_peri:=10
he3DThreshCounts_peri:=25
chromFWHMMin_peri:=0.4
apex_binThresh_peri:=550
apexTrackSNRThreshold_peri:=2.0
driftFilterScaleFactor_peri:=0.7
msFilterScaleFactor_peri:=0.7
apexTrackFilterWide_peri:=0
apexTrackSNRMeasFWHM_peri:=2.5
minScansPerPeak_peri:=6
msResolution_peri:=20000
```
noFilteredIons_peri:=1
bEnableTempCal_peri:=1

### Apex - from HaloSpecLib

| Setting                                  | Value  |
|------------------------------------------|--------|
| bCSVOutput_sec                           | 1      |
| apex_startRT_sec                         | 12     |
| apex_endRT_sec                           | 80     |
| le3DThreshCounts_sec                     | 10     |
| he3DThreshCounts_sec                     | 25     |
| chromFWHM_min_sec                        | 0.4    |
| apex_binThresh_sec                       | 550    |
| apexTrackSNRThreshold_sec               | 2      |
| chromFilterScaleFactor_sec              | 0.7    |
| driftFilterScaleFactor_sec              | 0.7    |
| msFilterScaleFactor_sec                 | 0.7    |
| apexTrackFilterWide_sec                 | 0      |
| apexTrackSNRMeasFWHM_sec                | 1      |
| minScansPerPeak_sec                     | 6      |
| msResolution_sec                        | 20000  |
| noFilteredIons_sec                      | 1      |
| bEnableTempCal_sec                      | 1      |

#### Peptide3D Settings

As with Apex3d Peptide3d has a many settings. A few of these were adjusted during pipeline development and are set here:

| Setting              | Value |
|----------------------|-------|
| minHEIntenThreshold  | 10    |
| minLEIntenThreshold  | 25    |
| msResolution         | 20000 |
| disableSingleHE      | 0     |

#### Peak Picking

The Apex3d and peptide3d algorithms between them work to pick peaks from the raw data, de-isotope them, relate precursor and product ions and produce spectra peaklists for database searching. I am not entirely clear how the tasks are split. I think Apex does the initial peak picking and feature detection and peptide3d associates precursor and product ions and determines peptide mass from multiply charged precursors and outputs a de-isotope peak list. I need to find some documentation to properly understand this.

#### Apex Feature Picking

The Apex algorithm is run with settings closely mirroring those run by PLGS. The major difference is substantial effort put into optimising the low-energy and high-energy thresholds that appear critical in getting high numbers of peptides hits.

For whole cell lysate:

```bash
define rawApex_rule
${1} : ${2}
   $(call makeDir,${1})
   ${apex_path} -pRawDirName $(call winify,${2}) \ 
     -pRawDirName $(dir $(call winify,${1})) \ 
     -outputDirName $(dir $(call winify,${1}))user \ 
     -bEnableLockMass 1 \ 
```
-noFilteredIons ${noFilteredIons} \ 
-bCSVOutput ${bCSVOutput_wcl} \ 
-lockmassZ1 556.2771 \ 
-lockMassToleranceAMU 0.25 \ 
-le3DThresholdCounts ${le3DThreshCounts_wcl} \ 
-he3DThresholdCounts ${he3DThreshCounts_wcl} \ 
-startingRTMin ${apex_stRT_wcl} \ 
-endingRTMin ${apex_endRT_wcl} \ 
-chromFWHMMin ${chromFWHMMin_wcl} \ 
-binIntenThreshold ${apex_binThresh_wcl} \ 
-WriteXML 0 \ 
-isADC 1 \ 
-bEnableCuda 0 \ 
-adcCalc2.128 1 \ 
-apexTrackSNRThreshold ${apexTrackSNRThreshold_wcl} \ 
-apexTrackMaxIterations 10 \ 
-chromFilterScaleFactor ${chromFilterScaleFactor_wcl} \ 
-driftFilterScaleFactor ${driftFilterScaleFactor_wcl} \ 
-msFilterScaleFactor ${msFilterScaleFactor_wcl} \ 
-apexTrackFilterWide ${apexTrackFilterWide_wcl} \ 
-apexTrack3DCombine 0 \ 
-4DReportFilterIntensity 1 \ 
-apexTrackSNRMeansFWHM ${apexTrackSNRMeansFWHM_wcl} \ 
-startingMassAmu 265 \ 
-endingMassAmu 2000 \ 
-msResolution ${msResolution_wcl} \ 
-minScansPerPeak ${minScansPerPeak_wcl} \ 
-bEnableTempCal ${bEnableTempCal_wcl} 

endef

$(foreach rawFile, ${rawFiles_All_wcl},$(eval $(call rawApex_rule,
 $(addprefix ${apexDir},$(notdir $(rawFile:.raw/_HEADER.TXT=_Apex3D.bin))),
 $(rawFile:.raw/_HEADER.TXT=.raw))))

For periplasm:

define rawApex_rule
${1} : ${2}
  $(call makeDir,${1})
  $(call prepapex_rule,${apex_path} \ 
  -pRawDirName $(call winify,${1}) \ 
  -outputDirName $(dir $(call winify,${1})):user \ 
  -bEnableLockMass 1 \ 
  -noFilteredIons ${noFilteredIons} \ 
  -bCSVOutput ${bCSVOutput_peri} \ 
  -lockmassZ1 556.2771 \ 
  -lockMassToleranceAMU 0.25 \ 
  -le3DThresholdCounts ${le3DThreshCounts_peri} \ 
  -he3DThresholdCounts ${he3DThreshCounts_peri} \ 
  -startingRTMin ${apex_stRT_peri} \ 
  -endingRTMin ${apex_endRT_peri} \ 
  -chromFWHMMin ${chromFWHMMin_peri} \ 

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define rawApex_rule
${1} : ${2}
  $(call makeDir,${1})
  ${apex_path} -pRawDirName $(call winify,${2})
    -outputDirName $(dir $(call winify,${1}))
    -outputUserDirName $(dir $(call winify,${1}))user
    -bEnableLockMass 1
    -noFilteredIons ${noFilteredIons}
    -bCSVOutput ${bCSVOutput_sec}
    -lockmassZ1 556.2771
    -lockMassToleranceAMU 0.25
    -le3DThresholdCounts ${le3DThreshCounts_sec}
    -he3DThresholdCounts ${he3DThreshCounts_sec}
    -startingRTMin ${apex_stRT_sec}
    -endingRTMin ${apex_endRT_sec}
    -chromFWHM ${chromFWHMMin_sec}
    -binIntenThreshold ${apex_binThresh_sec}
    -WriteXML 0
    -isADC 1
    -bEnableCuda 0
    -adcCalc2.128 1
    -apexTrackSNRThreshold ${apexTrackSNRThreshold_sec}
    -apexTrackMaxIterations 10
    -chromFilterScaleFactor ${chromFilterScaleFactor_sec}
    -binIntenThreshold ${apex_binThresh_peri}
    -WriteXML 0
    -isADC 1
    -bEnableCuda 0
    -adcCalc2.128 1
    -apexTrackSNRThreshold ${apexTrackSNRThreshold_peri}
    -apexTrackMaxIterations 10
    -chromFilterScaleFactor ${chromFilterScaleFactor_peri}
    -binIntenThreshold ${apex_binThresh_peri}
    -WriteXML 0
    -isADC 1
    -bEnableCuda 0
    -adcCalc2.128 1
    -apexTrackSNRThreshold ${apexTrackSNRThreshold_peri}
    -apexTrackMaxIterations 10
    -chromFilterScaleFactor ${chromFilterScaleFactor_peri}
    -4DReportFilterIntensity 1
    -apexTrackSNRMeasFWHM ${apexTrackSNRMeasFWHM_peri}
    -startingMassAmu 265
    -endingMassAmu 2000
    -msResolution ${msResolution_peri}
    -minScansPerPeak ${minScansPerPeak_peri}
    -bEnableTempCal ${bEnableTempCal_peri}
eendf

$(foreach rawFile, ${rawFiles_All_peri},$(eval $(call rawApex_rule,
  $(addprefix ${apexDir},$(notdir $(rawFile:.raw/_HEADER.TXT=_Apex3D.bin))),
  $(rawFile:.raw/_HEADER.TXT=.raw))))

For secretome:

define rawApex_rule
${1} : ${2}
  $(call makeDir,${1})
  ${apex_path} -pRawDirName $(call winify,${2})
    -outputDirName $(dir $(call winify,${1}))
    -outputUserDirName $(dir $(call winify,${1}))user
    -bEnableLockMass 1
    -noFilteredIons ${noFilteredIons}
    -bCSVOutput ${bCSVOutput_sec}
    -lockmassZ1 556.2771
    -lockMassToleranceAMU 0.25
    -le3DThresholdCounts ${le3DThreshCounts_sec}
    -he3DThresholdCounts ${he3DThreshCounts_sec}
    -startingRTMin ${apex_stRT_sec}
    -endingRTMin ${apex_endRT_sec}
    -chromFWHM ${chromFWHMMin_sec}
    -binIntenThreshold ${apex_binThresh_sec}
    -WriteXML 0
    -isADC 1
    -bEnableCuda 0
    -adcCalc2.128 1
    -apexTrackSNRThreshold ${apexTrackSNRThreshold_sec}
    -apexTrackMaxIterations 10
    -chromFilterScaleFactor ${chromFilterScaleFactor_sec}

Peptide3d Spectra Generation  The peptide3d algorithm is run with as close as possible to the PLGS defaults. Some changes have been made. There was an issue with peptide3d “hanging” with some files. The process would consume all the RAM and never complete. Documentation on this from Waters suggested altering some settings as noted in the code below.

The recipe uses the pipe | as part of its dependency instruction. All dependencies after this symbol are “order-only” dependencies. This means they will not force re-running of a recipe if they themselves are updated, however if they are to be updated in this run they must be rebuild prior to the current target. The feature is used here to ensure each pep3d file in the complete list of pep3d files has all subsequent pep3d files in the list as order only dependencies. The result is that only a single pep3d file may be produced at a time to prevent this particular operation overwhelming available RAM and crashing the process.
Merge  The fractionated whole cell lysate peak lists produced by peptide3d are merged prior to search and quantification by IADB.

define pep3dFileFromRaw
   $(addprefix ./pep3d/,$(notdir $(1:.raw/_HEADER.TXT=_Pep3D_Spectrum.bin)))
endef

merge_halo_wcl.ini : $(call pep3dFileFromRaw,${rawFilesHaloWCLPaths})
   $(file >$@) $(foreach bin,$^,$(file >>$@,$(call winify,${bin})))

merge_ecoli_wcl.ini : $(call pep3dFileFromRaw,${rawFilesEcoliWCLPaths})
   $(file >$@) $(foreach bin,$^,$(file >>$@,$(call winify,${bin})))

./pep3d/merged_halo_wcl_Pep3D_Spectrum.bin : merge_halo_wcl.ini
   ${mergeFractions_path} -inputFileSpectrums $(call winify,$<) \
   -outputUserDirName ".\pep3d\merge_halo_wcl" \ 
   -WriteXML 0 \ 
   -outputFile $(call winify, $@)

./pep3d/merged_ecoli_wcl_Pep3D_Spectrum.bin : merge_ecoli_wcl.ini
   ${mergeFractions_path} $< \ 
   -outputUserDirName ".\pep3d\merge_ecoli_wcl" \ 
   -WriteXML 0 \ 
   -outputFile $(call winify, $@)

IADB Search Whole Cell Lysate Merged Files

The search is run through the Waters search system iadb.exe. The program requires an input of spectra from a Waters .bin file; a .fasta database to search; and a .xml file specifying the search parameters. The latter was exported from the PLGS gui and edited below.

define searchSets
<?xml version="1.0" encoding="UTF-8"?>
<?xml version="1.0" encoding="UTF-8"?>
<WORKFLOW_TEMPLATE TITLE="FBRH_BacteriaSearch" UUID="${UUID}" WORKFLOW_TEMPLATE_ID="_15731280752780_5292366780985168">
   <PROTEINLYNX_QUERY TYPE="Databank-search">
      <DATABANK_SEARCH_QUERY_PARAMETERS>
         <SEARCH_ENGINE_TYPE VALUE="PLGS"/>
   </DATABANK_SEARCH_QUERY_PARAMETERS>
</PROTEINLYNX_QUERY>
</WORKFLOW_TEMPLATE>
The search execution is set up below. A rule is created to search every given pep3d file and output a matching search result.

```bash
define IadbFromPep3d
$(abspath $(addprefix ${iadbDir},$(notdir $(1:_Pep3D_Spectrum.bin=_IA_workflow.bin))))
```

```bash
define pepCSVfromPep3d
```

params.xml : 
```bash
$(file > params.xml,${searchSets})
```
$(abspath $(addprefix ${iadbTabDir},$(notdir $(1:_Pep3D_Spectrum.bin=_IA_final_peptide.csv))))
endef

define protCSVfromPep3d
$(abspath $(addprefix ${iadbTabDir},$(notdir $(1:_Pep3D_Spectrum.bin=_IA_final_protein.csv))))
endef

define fragCSVfromPep3d
$(abspath $(addprefix ${iadbTabDir},$(notdir $(1:_Pep3D_Spectrum.bin=_IA_final_fragment.csv))))
endef

define search_rule
$(call IadbFromPep3d,${1}) $(call fragCSVfromPep3d,${1}) $(call pepCSVfromPep3d,${1}) $(call protCSVfromPep3d,${1}) $(call makeDir,$(call IadbFromPep3d,${1})) $(call makeDir,$(call fragCSVfromPep3d,${1})) $(IADB_path) -paraXMLFileName params.xml -pep3DFileName $(call winify,${1}) -proteinFASTAFileName $(call winify,${2}) -proteinForAbsQuan "iRT_Quant" -protAmountForAbsQuanInFMol 434 -WriteXML 0 -allowHeMultiZ 1 -bAutoCalibrate 1 -fragmentTypes BY -bEnablePPMCalc 1 -protFalsePositiveRateMax 100 -outputDistractionProteins 1 -outputUserDirName $(call winify,$(dir $(call IadbFromPep3d,${1})))
endef

$(eval $(call search_rule,./pep3d/merged_ecoli_wcl_Pep3D_Spectrum.bin,$(fastaForSearchEcoli_decoy)))

$(eval $(call search_rule,./pep3d/merged_halo_wcl_Pep3D_Spectrum.bin,$(fastaForSearchHalo_decoy)))

$(eval $(call search_rule,./pep3d/merged_ecoli_all_Pep3D_Spectrum.bin,$(fastaForSearchEcoli_decoy)))

$(eval $(call search_rule,./pep3d/merged_halo_all_Pep3D_Spectrum.bin,$(fastaForSearchHalo_decoy)))
TPP Pipeline for Spectral Library

The trans-proteomic pipeline is used to process .mgf files produced by peptide3d above into a spectral library. The results of the TPP are cross referenced with the iadb search to yield a high quality list of quantified proteins.

X!Tandem Search  The first step is to convert .mgf files into mzXML files which can be searched by X!tandem:

```bash
define mzXML_Rule
\${1} : \${2}
    $(call makeDir,\${1})
    \${msconvert_path} \${2} --mzXML -o $(dir \${1}) --outfile $(notdir \${1})
endef

$(foreach mzXMLfile, \${all_mzXML},
    $(eval
        $(call mzXML_Rule,$\n            \${mzXMLfile},$\n            \${mzXMLfile:_Pep3D_Spectrum.mzXML=_Pep3DAMRT.mgf}$\n        )
    ))
```

Each of the resulting .mzXML files is then searched with x!tandem as below. The process is a little complex. For each mzxml file a copy is made of the standard xtandem parameter file. This copy is then edited with required spectrum file, target file name and fasta search file. After the search is complete, which outputs a tandem xml and an mzid file, the paramater file is deleted.

```bash
define Xtandem_Rule
\${1} $(1:.t.xml=.t.xml.mzid) & \${2} \${3} ./ini/default_input.xml ./ini/Xtandem_params.xml \${fastaForTandem}
    $(call makeDir,$\{1\})
    copy $(call winify, ./ini/Xtandem_params.xml) $(call winify, $(1:.t.xml=_params.xml))
    \${sed_path} -i "s|spectrumFile|$\{subst \,\,\,$\{2\}|" $(1:.t.xml=_params.xml)
    \${sed_path} -i "s|outFile|$\{subst \,\,\,$\{1\}|" $(1:.t.xml=_params.xml)
    \${sed_path} -i "s|taxonomyForSearch|$\{subst \,\,\,$\{3\}|" $(1:.t.xml=_params.xml)
    $(tandem_path) $(1:.t.xml=_params.xml)
    del /F $(call winify, $(1:.t.xml=_params.xml))
    if exist sed* del sed*
endef

$(foreach specFile, \${pep3dEcoilPaths_mzXML},
    $(eval
        $(call Xtandem_Rule,$\n            $(specFile:.mzXML=.t.xml),$\n            $(specFile),$\n            ./ini/taxonomy_ecoli.xml$\n        )
    ))
```

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The `.t.xml` files are then converted into `.pepXML` files to be processed by the rest of the TPP:

```bash
define Xtandem_pep_xml_Rule
${1}: ${2}
  $(call makeDir,${1})
  Tandem2XML_path $(call winify,${2}) $(call winify,${1})
endef

$(foreach pepXML, ${pep3dEcoliPaths_pepXML},
  $(eval 
    $(call Xtandem_pep_xml_Rule,
      ${pepXML},
      $(pepXML:.pep.xml=.t.xml) 
    )
  )
)

$(foreach pepXML, ${pep3dHaloPaths_pepXML},
  $(eval 
    $(call Xtandem_pep_xml_Rule,
      ${pepXML},
      $(pepXML:.pep.xml=.t.xml) 
    )
  )
)
```

**Interact and peptide prophet** The `.pepXML` files are then combined into single `.pepXML` files using TPP `xinteract`:

```bash
define xinteract_Rule
${1} : ${2}
  $(call makeDir,${1})
  if exist $(call winify,${1}), del /F $(call winify,${1})
  xinteract_path --eT --MONO --E${3} --17 --nP --D${3} --d${decoy_string} --N${1} ${2}
endef
```

```bash
$(eval 
  $(foreach pepXML, ${pep3dEcoliPaths_pepXML},
    $(call Xtandem_pep_xml_Rule,
      ${pepXML},
      $(pepXML:.pep.xml=.t.xml) 
    )
  )
)

$(foreach pepXML, ${pep3dHaloPaths_pepXML},
  $(call Xtandem_pep_xml_Rule,
    ${pepXML},
    $(pepXML:.pep.xml=.t.xml) 
  )
)
```

The `.t.xml` files are then converted into `.pepXML` files to be processed by the rest of the TPP:
The scores for the peptides are then adjusted using peptide prophet:

```bash
define peptideProphet_Rule
${1} : ${2}
 $(call makeDir,${1})
 if exist $(2:.pep.xml=_tmp.pep.xml) del /F $(2:.pep.xml=_tmp.pep.xml)
 if exist $(call winify,${1}), del /F $(call winify,${1})
 copy $(call winify,${2}) $(call winify,${2:.pep.xml=_tmp.pep.xml})
 if exist ${1} del /F ${1}
 $(PeptideProphet_path) $(call winify,${2:.pep.xml=_tmp.pep.xml}) \ 
 ACCMASS PPM MINPROB=0.05 DECOY=${decoy_string} \ 
 DECOYPROBS NONPARAM NONTT IGNORECHG=1 IGNORECHG=4 \ 
 IGNORECHG=5 IGNORECHG=6 IGNORECHG=7
 ren $(call winify,${2:.pep.xml=_tmp.pep.xml}) $(notdir ${1})
 endef
```

```bash
$(eval \
 $(call peptideProphet_Rule,$
      ./pep3d/interact_all_ecoli_prophet.pep.xml,$
      ./pep3d/interact_all_ecoli_all.pep.xml)\ 
 )
```

```bash
$(eval \
 $(call peptideProphet_Rule,$
      ./pep3d/interact_all_halo_prophet.pep.xml,$
      ./pep3d/interact_halo_all.pep.xml)\ 
 )
```

**Mayu FDR Estimation**  Mayu is a tool that estimates peptide and protein false discovery rates. Mayu is actually run twice here. One time with settings defined in `mayu_Rule` to find the correct score to set for spectra import into spectrast. The second time with settings defined in `mayu_Rule_plot` making calculations to higher FDR to produce data for figures presented in the paper.

```bash
define mayu_Rule
${1} ${2} &: ${3} ${4}
 $(call makeDir,${1})
```
${perl_path} ${mayu_path} \
-A $(call winify,${3}) \
-M $(subst _psm_protFDR0.05_t_1.07.csv,,$(call winify,${1})) \
-C $(call winify,${4}) \
-E ${decoy_string} \
-G 0.02 \
-H 49 \
-I 1 \
-P protFDR=0.05:t
endef

$(eval \
$(call mayu_Rule,\n./tpp_mayu/mayuOut_all_halo_psm_protFDR0.05_t_1.07.csv,\n./tpp_mayu/mayuOut_all_halo_main_1.07.csv,\n./pep3d/interact_all_halo_prophet.pep.xml,\n${fastaForSearchHalo_xtand_decoy})
)

$(eval \
$(call mayu_Rule,\n./tpp_mayu/mayuOut_all_ecoli_psm_protFDR0.05_t_1.07.csv,\n./tpp_mayu/mayuOut_all_ecoli_main_1.07.csv,\n./pep3d/interact_all_ecoli_prophet.pep.xml,\n${fastaForSearchEcoli_xtand_decoy})
)

define mayu_Rule_plot
${1} ${2} &: ${3} ${4}
$(call makeDir,${1})
${perl_path} ${mayu_path} \
-A $(call winify,${3}) \
-M $(subst _psm_protFDR0.04_t_1.07.csv,,$(call winify,${1})) \
-C $(call winify,${4}) \
-E ${decoy_string} \
-H 100 \
-PmFDR \
-I 1 \
-G 0.05 \
-P protFDR=0.04:t
endef

$(eval \
$(call mayu_Rule_plot,\n./tpp_mayu/mayuOut_all_halo_plot_psm_protFDR0.04_t_1.07.csv,\n./tpp_mayu/mayuOut_all_halo_plot_main_1.07.csv,\n./pep3d/interact_all_halo_prophet.pep.xml,\n
Spectrast and Spectral Library Generation  The integrated search results are now converted into a spectral library with spectrast. This takes pepXML files generated above, filters them to control FDR against the mayu result files and applies an iRT index.

```bash
define spectrast_Rule
$(1) $(1:.splib=.sptxt) &: $2 $3 $4
    if not exist $(dir $1) mkdir $(dir $1)
    $$ (eval mayuCutoff := $(shell ${awk_path} -F, "BEGIN{min=1}{if($$5<=min) min=$$5}END{print min}" $4)
    $$ (file > $$((dir $4)mayuCutoff.txt,$$mayuCutoff))
    ${spectrast_path} -V \
        -cN$(call unixify,$(basename $(1))) \n        -cICID-QTOF \n        -c_IRR \n        -c_IRT$(call unixify,$(3)) \n        -cP$${mayuCutoff} \n        -co \n        -c_RDY"${decoy_string}" \n        $(call unixify,$(2))
endef

# WCL
$(eval $(call spectrast_Rule,\
    ./tpp_mayu/halo_all.splib,\
    ./pep3d/interact_all_halo_prophet.pep.xml,$(RTindex),\
    ./tpp_mayu/mayuOut_all_halo_psm_protFDR0.05_t_1.07.csv))

$(eval $(call spectrast_Rule,\
    ./tpp_mayu/ecoli_all.splib,\
    ./pep3d/interact_all_ecoli_prophet.pep.xml,$(RTindex),\
    ./tpp_mayu/mayuOut_all_ecoli_psm_protFDR0.05_t_1.07.csv))
```

# WCL

```bash
$(eval $(call spectrast_Rule,\
    ./tpp_mayu/halo_all.splib,\
    ./pep3d/interact_all_halo_prophet.pep.xml,$(RTindex),\
    ./tpp_mayu/mayuOut_all_halo_psm_protFDR0.05_t_1.07.csv))

$(eval $(call spectrast_Rule,\
    ./tpp_mayu/ecoli_all.splib,\
    ./pep3d/interact_all_ecoli_prophet.pep.xml,$(RTindex),\
    ./tpp_mayu/mayuOut_all_ecoli_psm_protFDR0.05_t_1.07.csv))
```
define spectrastCons_Rule
${1} $(1:.splib=.sptxt) & : ${2} ${3}
    if not exist $(dir ${1}) mkdir $(dir ${1})
    ${spectrast_path} -V \
    -cN$(basename ${1}) \n    -cICID-QTOF \n    -c_ANN \n    -cf"iRT<190" \n    -cu \n    -c_NPK7 \n    -cD${3} \n    -cAC \n    ${2}
endef

$(eval $(call spectrastCons_Rule,./tpp_mayu/ecoli_all_cons.splib,./tpp_mayu/ecoli_all.splib,${fastaForSpectrastEcoli}))
$(eval $(call spectrastCons_Rule,./tpp_mayu/ecoli_SWATHatlas2020.splib,./litData/EcoliSWATHatlas2020/Ecoli_consensus.sptxt,${fastaForSpectrastEcoli}))
$(eval $(call spectrastCons_Rule,./tpp_mayu/halo_all_cons.splib,./tpp_mayu/halo_all.splib,${fastaForSpectrastHalo}))

The spectral libraries produced by spectrast are now converted into .tsv format for ease of use with other DIA tools such as openSWATH and peakview.

define spectrast2tsv_Rule
${1} : ${2}
    if not exist $(dir ${1}) mkdir $(dir ${1})
    ${python_path} ${spectrast2tsv_path} \
    -l 100,2000 \
    -s b,y \
    -x 1,2 \
    -g -17.03,-18.0 \
    -o 3 \
    -n 20 \
    -p 0.05 \
    -d -e \
    -k openswath \
    -a ${1} \n    ${2}
endef

$(foreach splib2conv, ${ConsLib},
    $(eval 
        $(call spectrast2tsv_Rule, 
            $(basename ${splib2conv})_os.tsv, 
            $(basename ${splib2conv}).sptxt, 
        )
    )
)
define spectrast2tsv_Rule
${1} : ${2} 
  if not exist $(dir ${1}) mkdir $(dir ${1})
  ${python_path} ${spectrast2tsv_path} \
    -l 100,2000 \
    -s b,y \
    -x 1,2 \
    -g -17.03,-18.0 \
    -o 3 \
    -n 20 \
    -p 0.05 \
    -d -e \
    -k peakview \
    -a ${1} \n    ${2} 
endef

$(foreach splib2conv, ${ConsLib},\n  $(eval \n    $(call spectrast2tsv_Rule,\n      $(basename ${splib2conv})_pk.tsv,\n      $(basename ${splib2conv}).sptxt,\n      )\n  )\n)

Create Upload Package

Copy raw data from the various data directories to the upload

define zip7raw
$(foreach comp,${rawDirCompsNoDat},${addsuffix /${comp},${uploadDir}$(notdir $(1:.raw/_HEADER.TXT=.raw.zip))}$zip7_path) u $(call winify,$(dir $(firstword $(foreach comp,${rawDirComps},${addsuffix /${comp},${uploadDir}$(notdir $(1:.raw/_HEADER.TXT=empty.raw.zip))})))
  if exist $(call winify,$(dir $(firstword $(foreach comp,${rawDirComps},${addsuffix /${comp},$(1:.raw/_HEADER.TXT=.raw))))
  $(foreach comp,${rawDirComps},${addsuffix /${comp},${uploadDir}$(notdir $(1:.raw/_HEADER.TXT=empty.raw.zip)))))
endef

rawFileTest:=$(firstword ${rawFiles_All})

$(info testUpload: $(rawFileTest:.raw/_HEADER.TXT=.raw))

#$(foreach raw, ${rawFiles_All},${info $(call zip7raw,$(raw)))

$(foreach raw, ${rawFiles_All},\
  $(eval \n    $(call zip7raw,\
      ${raw}\n    )\n  )\n)

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define copyToUpload_MZID
${1} : ${2}
    copy /y $(subst /,\,\$(abspath ${2})) $(subst /,\,\$(abspath ${1}))
endef

# # $(info MZID: ${all_mzid})

$(foreach file, ${all_mzid},\
    $(eval \
        $(call copyToUpload_MZID,\
            $(addprefix ${uploadDir},$(notdir $(file:_Pep3D_Spectrum.t.xml.mzid=.mzid))),\
            ${file}\
        )\
    )\
)

define copyToUpload
$(addprefix ${uploadDir},${(notdir ${1})) : ${1}
    copy /y $(call winify, ${1}) $(call winify,${uploadDir})
endef

$(foreach file, ${intermediateFilesForUpload},\
    $(eval \
        $(call copyToUpload,\
            ${file}\
        )\
    )\
)