

Portable Proteomics Pipeline (P3)

MSGF Benchmark

Daniel Kristiyanto (daniel.kristiyanto@pnnl.gov)

May 26, 2016

1. Files:

```
## [1] "TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f11.mzML"
```

```
## [1] "ID_003632_9011437E.fasta"
```

2. Identification:

MSGFPlus.jar was downloaded from <https://omics.pnl.gov/software/ms-gf>. Identification was run multiple times (within or without Docker), and resulted the same results.

Identification with different switches resulted different results.

```
# Command Line (Without Switch)
```

```
java -Xmx8000M -jar P3/MSGFPlus.jar -s \  
TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f11.mzML -o \  
TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f11_no_switch.mzid \  
-d ID_003632_9011437E.fasta
```

```
# Command Line (2)
```

```
java -Xmx8000M -jar P3/MSGFPlus.jar -s \  
TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f11.mzML -o \  
TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f11.mzid \  
-d ID_003632_9011437E.fasta \  
-t 10ppm -m 0 -inst 1 -e 1 -ti -1,2 -ntt 2 -tda 1 -minLength 6 \  
-maxLength 50 -minCharge 2 -maxCharge 5 -n 1 -thread 7 \  
-mod MSGFDB_Mods.txt -minNumPeaks 5 -addFeatures 1
```

```
## [1] "TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f11_no_switch.mzid"
```

```
## [2] "TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f11.mzid"
```

3. Result from PNNL

```
## [1] 12695
```

```
##      Scan MSGFDB_SpecEValue      Peptide
## 1 11612      1.481722e-36 IQDLTGLDVSTAEELQVANYGVGGQYEPHFDFAR
## 2 14441      8.834771e-36 LTPFIIQENLNLALNSASAIGCHVNVNIGAEDLR
```

```
##      Scan MSGFDB_SpecEValue Peptide
## 12947 3823      0.04856372 SPAGGG
## 12948 4099      0.05022644 PGGAGM
```

4. Result from MSGF+ without any switches (sorted by SpecE-Value)

```
##      scan number(s) ms-gf:denovoscore ms-gf:specevalue
## X9740.1      10815      277      4.434865e-30
## X8340.1      9275      234      2.157325e-27
## X10711.1     11883      157      1.386474e-25
##
##      pepseq
## X9740.1 SHSTEPGLVLTGQGQDVGGQLGLGENVMER
## X8340.1 DLYANTVLSGGTTMYPGIADR
## X10711.1 ILGGVISAISEAAAQYNPEPPPPR
```

```
##      scan number(s) ms-gf:denovoscore ms-gf:specevalue
## X13036.1     14441      381      4.861374e-08
## X10464.1     11612      279      5.246882e-08
##
##      pepseq
## X13036.1 KETDLKQIQTLIQGTQTRLKYSQNELEMIKK
## X10464.1 AGISEAQLTDAETSKLIYDFIEDQGGLEAVRQEMR
```

Reading the MZID file manually

```
java -Xmx2000M -XX:+UseConcMarkSweepGC -cp \
../P3/MSGFPlus.jar edu.ucsd.msjava.ui.MzIDToTsv -i \
TCGA_13-1489_42-2590_36-2529_117C_W_PNNL_B2S5_f11_no_switch.mzid \
-showQValue 1 -showDecoy 1 -unroll 1
```

```
##      ScanNum      Peptide      Protein SpecEValue
## 1      10815 R.SHSTEPGLVLTGQGQDVGGQLGLGENVMER.K ref|NP_001041660.1 4.43e-30
## 2      10815 R.SHSTEPGLVLTGQGQDVGGQLGLGENVMER.K ref|NP_001041659.1 4.43e-30
## 3      10815 R.SHSTEPGLVLTGQGQDVGGQLGLGENVMER.K ref|NP_001041664.1 4.43e-30
```

5. Results with switches

```
##      scan number(s) ms-gf:denovoscore ms-gf:specevalue
## X11197.1     12418      321      6.104450e-34
## X9050.1      10056      258      4.553317e-33
```

```
## X9060.1          10067          288      9.340615e-33
##                                     pepseq
## X11197.1 VYLASLETLDNGKPFQESYALDLDEVIK
## X9050.1      CLCLPSYVGALCEQDTETCDYGWHK
## X9060.1      CLCLPSYVGALCEQDTETCDYGWHK

##          scan number(s) ms-gf:denovoscore ms-gf:specvalue      accession
## X13036.1      14441          345      2.196462e-06 ref|NP_060250.2
## X10464.1      11612          279      2.280634e-06 ref|NP_000487.1
##                                     pepseq
## X13036.1 FILPNVSTPVSDAFKTQMELLQAGLSRTPTR
## X10464.1  GDYKDSSDFGAPHPQVQSVRRIRDMQWHQR
```

Reading the MZID file manually

```
##      ScanNum
## 1      12418
## 2      10056
## 3      10056
##
##                                     Peptide
## 1      R.+144.102VYLASLETLDNGK+144.102PFQESYALDLDEVIK+144.102.V
## 2 R.+144.102C+57.021LC+57.021LPSYVGALC+57.021EQDTETC+57.021DYGWHK+144.102.F
## 3 R.+144.102C+57.021LC+57.021LPSYVGALC+57.021EQDTETC+57.021DYGWHK+144.102.F
##      SpecEValue
## 1      6.10e-34
## 2      4.55e-33
## 3      4.55e-33
```

7. Filtering (PNNL -> Docker)

a. Spec-Evalue 10^{-10}

From PNNL

There are 4633 rows remained.

From Docker

There are 1665 rows remained.

```
## [1] "=="Head=="
```

```
##          Scan MSGFDB_SpecEValue      Peptide ms-gf:specvalue
## 3641 12418      6.104450e-34 VYLASLETLDNGKPFQESYALDLDEVIK      6.104450e-34
```

```
## 2717 10056      4.553317e-33    CLCLPSYVGALCEQDTETCDYGWHK      4.553317e-33
##                                     pepseq
## 3641 VYLASLETLDNGKPFQESYALDLDEVIK
## 2717      CLCLPSYVGALCEQDTETCDYGWHK
```

```
## [1] "==Tail=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide ms-gf:specevalue
## 2508 9513      2.871564e-13    NIIGSSPVADFSAIK      8.381110e-05
## 223  1927      5.243567e-25    CCYDGACVNNDETCEQR      9.379988e-05
##                                     pepseq
## 2508 LVDETEGQCGETDPNSEMPR
## 223   HECCFRYHCTGCCFR
```

Differences

Data from Docker is matched with from PNNL using scan number as ID. MSGFDB_SpecEValue, Peptide are values from PNNL. ms-gf:specevalue, pepseq are the corresponding values from Docker Container. There are 4621 of rows in total with 1227 exact match.

325 with the same peptide identification and different spec-evalue, 3069 of different peptide identification as well different spec-evalues (12 of them are unmatched values).

```
## [1] "==Unmatched=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide ms-gf:specevalue pepseq
## 2291 8856      1.119096e-15    CSEGVFLLTTPR      NA    <NA>
## 1416 6555      6.590530e-15    CQSLQEELDFRK      NA    <NA>
```

```
## [1] "==Different Spec eValues=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide
## 3127 11078      5.495878e-31    MGLDPACQVDIICGDHLLEQYQTLR
## 2548 9602      2.682754e-30    WSGPLSLQEVDEQPQHPLHVTYAGAAVDELGK
##      ms-gf:specevalue      pepseq
## 3127      7.059070e-31    MGLDPACQVDIICGDHLLEQYQTLR
## 2548      1.788561e-30    WSGPLSLQEVDEQPQHPLHVTYAGAAVDELGK
```

```
## [1] "==Different Peptides identification and different Spec Evalues=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide ms-gf:specevalue
## 2595 9730      6.005831e-22    FFDHSGTLVMDAYEPEISR      2.231272e-20
## 4204 13789      4.811341e-14    NAQMAQSPILLGGAASTLLQNR      6.118720e-19
##                                     pepseq
## 2595 DLYANTVLSGGTTMPGIADR
## 4204 IQTQLNLIHPDIFPLLTFR
```

b. Spec-Evalue 10^{-15}

From PNNL

There are 2511 rows remained.

From Docker

There are 767 rows remained.

```
## [1] "=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide ms-gf:specvalue
## 1763 12418      6.104450e-34 VYLASLETLDNGKPFQESYALDLDEVIK      6.104450e-34
## 1170 10056      4.553317e-33  CLCLPSYVGALCEQDTETCDYGWHK      4.553317e-33
##
##      pepseq
## 1763 VYLASLETLDNGKPFQESYALDLDEVIK
## 1170  CLCLPSYVGALCEQDTETCDYGWHK
```

```
## [1] "=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide ms-gf:specvalue
## 49 1876      1.449447e-17  QNEAAVGNCAEHMR      3.858078e-05
## 50 1927      5.243567e-25  CCYDGACVNNDTCEQR      9.379988e-05
##
##      pepseq
## 49  DSFMRHRGH CER
## 50  HECCFRYHCTGCCFR
```

Differences

Data from Docker is matched with from PNNL using scan number as ID. MSGFDB_SpecEValue, Peptide are values from PNNL. ms-gf:specvalue, pepseq are the corresponding values from Docker Container. There are 2511 of rows in total with 572 exact match.

170 with the same peptide identification and different spec-evalue, 1769 of different peptide identification as well different spec-evalues (0 of them are unmatched values).

```
## [1] "=="
```

```
## [1] Scan      MSGFDB_SpecEValue Peptide      ms-gf:specvalue
## [5] pepseq
## <0 rows> (or 0-length row.names)
```

```
## [1] "=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide
## 1437 11078      5.495878e-31      MGLDPACQVDIICGDHLEQYQTLR
## 1057 9602      2.682754e-30 WSGPLSLQEVD EQPHPLHVTYAGAAVDELGK
##      ms-gf:specevalue      pepseq
## 1437      7.059070e-31      MGLDPACQVDIICGDHLEQYQTLR
## 1057      1.788561e-30 WSGPLSLQEVD EQPHPLHVTYAGAAVDELGK
```

```
## [1] "==Different Peptides identification and different Spec Evalues=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide ms-gf:specevalue
## 1091 9730      6.005831e-22 FFDHSGTLVMDAYEPEISR      2.231272e-20
## 123 3378      2.191136e-20 ASGNYATVISHNPETKK      2.189761e-14
##      pepseq
## 1091 DLYANTVLSGGTTMYPGIADR
## 123      NKPGPYSSVPPPSAPPPK
```

c. Spec-Evalue 10^{-20}

From PNNL

There are 936 rows remained.

From Docker

There are 223 rows remained.

```
## [1] "==Head=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide ms-gf:specevalue
## 571 12418      6.104450e-34 VYLASLETLDNGKPFQESYALDLDEVIK      6.104450e-34
## 327 10056      4.553317e-33 CLCLPSYVGALCEQDTETCDYGWHK      4.553317e-33
##      pepseq
## 571 VYLASLETLDNGKPFQESYALDLDEVIK
## 327 CLCLPSYVGALCEQDTETCDYGWHK
```

```
## [1] "==Tail=="
```

```
##      Scan MSGFDB_SpecEValue      Peptide ms-gf:specevalue
## 19 3273      1.342643e-21 ESCNPQDCVPEVDENCKDK      2.671039e-05
## 12 1927      5.243567e-25 CCYDGACVNNDTCEQR      9.379988e-05
##      pepseq
## 19 AEGAYYDNHNSVVYLDHGENGEK
## 12      HECCFRYHCTGCCFR
```

Differences

Data from Docker is matched with from PNNL using scan number as ID. MSGFDB_SpecEValue, Peptide are values from PNNL. ms-gf:specvalue, pepseq are the corresponding values from Docker Container. There are 936 of rows in total with 165 exact match.

54 with the same peptide identification and different spec-evalue, 717 of different peptide identification as well different spec-evalues (0 of them are unmatched values).

```
## [1] "==Unmatched=="
```

```
## [1] Scan          MSGFDB_SpecEValue Peptide          ms-gf:specvalue
## [5] pepseq
## <0 rows> (or 0-length row.names)
```

```
## [1] "==Different Spec eValues=="
```

```
##      Scan MSGFDB_SpecEValue          Peptide
## 450 11078      5.495878e-31      MGLDPACQVDIICGDHLEQYQTLR
## 284  9602      2.682754e-30 WSGPLSLQEVDEQPQHPLHVTYAGAAVDELGK
##      ms-gf:specvalue          pepseq
## 450      7.059070e-31      MGLDPACQVDIICGDHLEQYQTLR
## 284      1.788561e-30 WSGPLSLQEVDEQPQHPLHVTYAGAAVDELGK
```

```
## [1] "==Different Peptides identification and different Spec Evalues=="
```

```
##      Scan MSGFDB_SpecEValue          Peptide ms-gf:specvalue
## 299  9730      6.005831e-22      FFDHSGTLVMDAYEPEISR      2.231272e-20
## 401 10627      4.148874e-23 GMSLNLEPDNVGVVVFVFGNDK      7.877905e-10
##
##      pepseq
## 299 DLYANTVLSGGTTMYPGIADR
## 401  GNAEPDPPVRLFQIHGNDK
```