

Executable Analysis Document Supporting Proteomics Component of the Manuscript: “Widespread Abrogation of Triplet Translation Continuity and Stop Codon Function in *Euplotes*”

Alexei V. Lobanov, Stephen M. Heaphy, Anton A. Turanov, Maxim V. Gerashchenko, Sandra Pucciarelli, Raghul R. Devaraj, Fang Xie, Vladislav A. Petyuk, Richard D. Smith, Lawrence A. Klobutcher, John F. Atkins, Cristina Miceli, Dolph L. Hatfield, Pavel V. Baranov, Vadim N. Gladyshev

Tue 21 Jun 2016

Contents

1	Introduction	1
2	Post MS/MS Search Analysis Steps	2
2.1	Prerequisites	2
2.1.1	Downloading Datasets	2
2.1.2	Reading Frameshift Marks	2
2.2	Processing of MS/MS Search Results	3
2.2.1	Trypsin Digest Fractionated by SCX	3
2.2.2	Trypsin Digest Fractionated by HPRP	6
2.2.3	Glu-C Digest Fractionated by HPRP	8
2.3	Compendium of Peptides Covering Frameshift Locations	10
3	Manual Validation	11
4	Session Information	19

1 Introduction

The vignette describes and reproduces all the steps that aimed to confirm frameshifts in the *Euplotes crassus* proteome. The global 8M urea soluble proteome was digested using conventional trypsin protocol and alternatively with Glu-C protease under high pH (7.5) conditions. The latter restricts specificity of Glu-C cleavages to C-terminal of glutamic acid (E). The peptides resulting from trypsin digest were fractionated using two different approaches: with strong cation exchange (SCX) and high pH reverse phase (HPRP) chromatographies. The peptides from Glu-C digest were fractionated using HPRP only.

The datasets were deposited to PRIDE and available by this link <http://dx.doi.org/10.6019/PXD004333>. Summary of the datasets shown in the table below:

Dataset Prefix	Digestion Enzyme	Fractionation Chromatography Type
Euplotes_1_SCX	trypsin	SCX
Euplotes_1_HPRP_1	trypsin	HPRP
Euplotes_1_HPRP_2	Glu-C (pH 7.5)	HPRP

Preprocessing of the raw files prior MS/MS searches was done in two steps. First, the raw files were processed with

[DeconMSN](#) to correct for wrong assignments of monoisotopic peaks. The parameters are as follows:

```
DeconMSN.exe -I35 -G1 -F1 -L6810 -B200 -T5000 -M3 -XCDTA
```

At the second step the peak files were processed with [DtaRefinery](#) to perform post-acquisition recalibration of parent ion mass-to-charge ratios. The peak lists (concatenated dta files in this case) were searched using [MS-GF+](#) tool against 6-frame translated *Euplotes Crassus* genome concatenated with tentatively frameshifted sequences and common contaminants. The 6-frame translated FASTA file, DtaRefinery and MS-GF+ parameter files are available in extdata folder of the EuplotesCrassus.proteome package.

For example:

```
fpath <- system.file("extdata",
                    "MSGFDB_GluC_StatCysAlk_10ppmParTol.txt",
                    package="EuplotesCrassus.proteome")
cat(readLines(fpath, n=12), sep = '\n')
## #Parent mass tolerance
## # Examples: 2.5Da or 30ppm
## # Use comma to set asymmetric values, for example "0.5Da,2.5Da" will set 0.5Da to the left (expMass<th
## PMTolerance=10ppm
##
## #Max Number of Modifications per peptide
## # If this value is large, the search will be slow
## NumMods=3
##
## #Modifications (see below for examples)
## StaticMod=C2H3N1O1, C, fix, any, Carbamidomethyl # Fixed Carbamidomethyl C (alkylation,
```

2 Post MS/MS Search Analysis Steps

2.1 Prerequisites

2.1.1 Downloading Datasets

To download the datasets we will take advantage of [rpx](#) R package. Note, this step may take awhile (10-30 min) depending on the speed of the internet connection. However, if they are downloaded the script will use the available datasets instead of downloading them again.

```
library(rpx)
id <- "PXD004333"
px <- PXDataset(id)
repoFiles <- pxfiles(px)
mzids <- grep('*msgfplus.mzid.gz', repoFiles, value=T)
system.time(pxget(px, mzids))
##      user      system elapsed
##    1.093      6.608    580.919
```

2.1.2 Reading Frameshift Marks

The FASTA files containing 595 sequences with frameshifts available as a part of this package and available as `system.file("extdata", "Euplotes_Crassus_frameshifts.fasta", package="EuplotesCrassus.proteome")`. There is an additional FASTA file with frameshift locations marked with exclamation mark !.

```
library(Biostrings)
fasta_clean <- readAAStringSet(
  system.file("extdata",
              "Euplotes_Crassus_frameshifts.fasta",
              package="EuplotesCrassus.proteome"),
  format="fasta", nrec=-1L, skip=0L, use.names=TRUE)
fasta_marks <- readAAStringSet(
  system.file("extdata",
              "Euplotes_Crassus_frameshifts_with_mark.fasta",
              package="EuplotesCrassus.proteome"),
  format="fasta", nrec=-1L, skip=0L, use.names=TRUE)
length(fasta_clean)
## [1] 595
```

2.2 Processing of MS/MS Search Results

2.2.1 Trypsin Digest Fractionated by SCX

For processing of MS/MS identification we will use [MSnID](#) R package. First step is to read the LC-MS/MS datasets corresponding to 25 SCX fractions.

```
library(MSnID)
trypscpx <- grep('Euplotes_1_SCX_.*msgfplus.mzid.gz', repoFiles, value=T)
trypscpxPrj <- MSnID()
system.time(trypscpxPrj <- read_mzIDs(trypscpxPrj, trypscpx, backend = 'mzR'))
##      user      system elapsed
## 11.634    4.734   37.345
```

Assess the peptide termini for their corresponding cleavage patterns. We will leave peptides that resulted only from proper trypsin cleavage events. That is we won't allow peptide resulting from irregular cleavages.

```
trypscpxPrj <- assess_termini(trypscpxPrj, validCleavagePattern="[KR]\\.[^P]")
trypscpxPrj <- apply_filter(trypscpxPrj, "numIrregCleavages == 0")
```

Note, that for this project we are interested only in peptides covering the sites of the frameshifting events. So if a peptide identification can be explained by a regular protein sequence we are not interested in pursuing this identification. The protein/accession names of normal (non-frameshifted) sequences starts with Contig or Contaminant. If the FASTA entry sequence is a results of the frameshift event if starts with comp. Therefore in the code below we retain only peptide-to-spectrum matches that can appear only due to frameshifted sequences.

```
## Rule on how to split the names.
## Contig + Contaminants - main piece
## comp - sequences with frameshifts
trypscpxPrj.main <- apply_filter(trypscpxPrj, "!grepl('comp', accession)")
trypscpxPrj.fmsh <- apply_filter(trypscpxPrj, "grepl('comp', accession)")
## if peptide matches to the main piece we don't care about it
trypscpxPrj.fmsh <- apply_filter(trypscpxPrj.fmsh,
                                "!(peptide %in% peptides(trypscpxPrj.main))")
show(trypscpxPrj.fmsh)
## MSnID object
## Working directory: "."
## #Spectrum Files: 25
## #PSMs: 442 at 58 % FDR
## #peptides: 348 at 67 % FDR
```

```
## #accessions: 291 at 66 % FDR
```

Setting-up and optimizing filtering options for MS/MS identifications. Since the number of peptides mapping frameshifted sequences is rather low we will loosed up the FDR of the identification up to 5%, however, then follow-up with manual spectra validation.

```
trypscxPrj.fmsh$mme.ppm <- abs(mass_measurement_error(trypscxPrj.fmsh))
trypscxPrj.fmsh$score <- -log10(trypscxPrj.fmsh$`MS.GF.SpecEValue`)
trypscxPrj.fmsh <- apply_filter(trypscxPrj.fmsh, "mme.ppm < 10")

filtr <- MSnIDFilter(trypscxPrj.fmsh)
filtr$mme.ppm <- list(comparison="<", threshold=5.0)
filtr$score <- list(comparison=">", threshold=8.0)
#' pre-optimization with brute-force approach
filtr.grid <- optimize_filter(filtr, trypscxPrj.fmsh, fdr.max=0.05,
                             method="Grid", level="peptide", n.iter=20000)
evaluate_filter(trypscxPrj.fmsh, filtr.grid)

##           fdr    n
## PSM      0.02970297 104
## peptide  0.03703704  56
## accession 0.04166667  50
```

```
#' fine tune with optimization using simulated annealing technique
filtr.sann <- optimize_filter(filtr.grid, trypscxPrj.fmsh, fdr.max=0.05,
                              method="SANN", level="peptide", n.iter=20000)
evaluate_filter(trypscxPrj.fmsh, filtr.sann)

##           fdr    n
## PSM      0.02941176 105
## peptide  0.03636364  57
## accession 0.04081633  51
```

```
trypscxPrj.fmsh <- apply_filter(trypscxPrj.fmsh, filtr.sann)
show(trypscxPrj.fmsh)
## MSnID object
## Working directory: "."
## #Spectrum Files: 18
## #PSMs: 105 at 2.9 % FDR
## #peptides: 57 at 3.6 % FDR
## #accessions: 51 at 4.1 % FDR
```

Finally we will extract only those peptides that exactly span the frameshift sites. That is their sequences should be present/identifiable in normal FASTA file, however missing in the file with frameshifts masked with the exclamation mark !.

```
#' extract only those that map frameshift sites
library(dplyr)
pepSeq <- unique(trypscxPrj.fmsh$pepSeq)
pepSeqMapped_to_clean <- pepSeq %>%
  sapply(grep, x=fasta_clean) %>%
  sapply(length) %>%
  subset(>.0) %>%
  names
pepSeqMapped_to_with_marks <- pepSeq %>%
  sapply(grep, x=fasta_marks) %>%
  sapply(length) %>%
```

```
subset(>0) %>%
  names
pepSeqFmsh_trypscx <- setdiff(pepSeqMapped_to_clean, pepSeqMapped_to_with_marks)
print(pepSeqFmsh_trypscx)
## [1] "SAQEEQDDEVIIDDQNPLLEDDLQIDEPEQK" "WTPIDLPSSEITFVQGIQTVTGAGDPSMK"
## [3] "ESNHNNNDITNKNEIAYILR" "KKKQEENNLKR"
```

Reporting extra information on the peptide sequences spanning frameshift sites: dataset, scan, charge, score, and mass measurement error.

```
meta_tryp_scx <- trypscxPrj.fmsh %>%
  apply_filter('pepSeq %in% pepSeqFmsh_trypscx') %>%
  psms %>%
  select(spectrumFile,MS.GF.SpecEValue,mme.ppm,spectrumID,chargeState,peptide) %>%
  rename(SpecEValue = MS.GF.SpecEValue, charge = chargeState, `MME (ppm)`=mme.ppm) %>%
  mutate(spectrumFile = sub('_msgfplus.mzid.gz',' ',spectrumFile))
library(xtable)
print(xtable(meta_tryp_scx, display = c('d','s','e','f','s','d','s')),
      include.rownames=FALSE,
      comment = FALSE,
      size='scriptsize',
      floating = F)
```

spectrumFile	SpecEValue	MME (ppm)	spectrumID	charge	peptide
Euplotes_1_SCX_10_13Nov09_Falcon_09-09-14	3.41e-15	0.30	index=6106	3	K.SAQEEQDDEVIIDDQNPLLEDDLQIDEPEQK.V
Euplotes_1_SCX_10_13Nov09_Falcon_09-09-14	3.41e-15	0.30	index=6106	3	K.SAQEEQDDEVIIDDQNPLLEDDLQIDEPEQK.V
Euplotes_1_SCX_12_13Nov09_Falcon_09-09-14	1.53e-21	0.08	index=8908	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_SCX_12_13Nov09_Falcon_09-09-14	1.07e-20	1.10	index=8896	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_SCX_12_13Nov09_Falcon_09-09-14	7.29e-19	1.10	index=8897	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_SCX_12_13Nov09_Falcon_09-09-14	2.17e-15	0.94	index=8895	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_SCX_18_13Nov09_Falcon_09-09-15	9.27e-17	0.11	index=5912	2	K.ESNHNNNDITNKNEIAYILR.Y
Euplotes_1_SCX_20_13Nov09_Falcon_09-09-15	2.23e-11	0.70	index=10317	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_SCX_22_13Nov09_Falcon_09-09-15	4.36e-10	3.76	index=9720	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_SCX_23_13Nov09_Falcon_09-09-15	2.47e-09	1.64	index=9440	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_SCX_24_13Nov09_Falcon_09-09-15	3.42e-10	8.85	index=2127	3	R.KKKQEENNLKR.K

2.2.2 Trypsin Digest Fractionated by HPRP

All the processing steps are conceptually the same as in the section above.

```
tryphprp <- grep('Euplotes_1_HPRP_1_*.msgfplus.mzid.gz', repoFiles, value=T)
tryphprpPrj <- MSnID()
system.time(tryphprpPrj <- read_mzIDs(tryphprpPrj, tryphprp, backend = 'mzR'))
##      user  system elapsed
##  7.895   4.458  36.343
```

```
tryphprpPrj <- assess termini(tryphprpPrj, validCleavagePattern="[KR]\\.[^P]")
tryphprpPrj <- apply_filter(tryphprpPrj, "numIrregCleavages == 0")
```

```
tryphprpPrj.main <- apply_filter(tryphprpPrj, "!grepl('comp', accession)")
tryphprpPrj.fmsh <- apply_filter(tryphprpPrj, "grepl('comp', accession)")
tryphprpPrj.fmsh <- apply_filter(tryphprpPrj.fmsh,
                                "!(peptide %in% peptides(tryphprpPrj.main))")
show(tryphprpPrj.fmsh)
## MSnID object
## Working directory: "."
## #Spectrum Files: 24
## #PSMs: 511 at 49 % FDR
## #peptides: 399 at 62 % FDR
## #accessions: 293 at 78 % FDR
```

```
tryphprpPrj.fmsh$mme.ppm <- abs(mass_measurement_error(tryphprpPrj.fmsh))
tryphprpPrj.fmsh$score <- -log10(tryphprpPrj.fmsh$`MS.GF.SpecEValue`)
tryphprpPrj.fmsh <- apply_filter(tryphprpPrj.fmsh, "mme.ppm < 10")

filtr <- MSnIDFilter(tryphprpPrj.fmsh)
filtr$mme.ppm <- list(comparison="<", threshold=5.0)
filtr$score <- list(comparison=">", threshold=8.0)
filtr.grid <- optimize_filter(filtr, tryphprpPrj.fmsh, fdr.max=0.05,
                             method="Grid", level="peptide", n.iter=20000)
evaluate_filter(tryphprpPrj.fmsh, filtr.grid)
##           fdr    n
## PSM      0.02631579 195
## peptide  0.04504505 116
## accession 0.07142857 75
```

```
filtr.sann <- optimize_filter(filtr.grid, tryphprpPrj.fmsh, fdr.max=0.05,
                             method="SANN", level="peptide", n.iter=20000)
evaluate_filter(tryphprpPrj.fmsh, filtr.sann)
##           fdr    n
## PSM      0.02604167 197
## peptide  0.04504505 116
## accession 0.07142857 75
```

```
tryphprpPrj.fmsh <- apply_filter(tryphprpPrj.fmsh, filtr.sann)
show(tryphprpPrj.fmsh)
## MSnID object
## Working directory: "."
## #Spectrum Files: 23
## #PSMs: 197 at 2.6 % FDR
## #peptides: 116 at 4.5 % FDR
```

```
## #accessions: 75 at 7.1 % FDR
```

```
library(dplyr)
pepSeq <- unique(tryphprpPrj.fms$pepSeq)
pepSeqMapped_to_clean <- pepSeq %>%
  sapply(grep, x=fasta_clean) %>%
  sapply(length) %>%
  subset(>0) %>%
  names
pepSeqMapped_to_with_marks <- pepSeq %>%
  sapply(grep, x=fasta_marks) %>%
  sapply(length) %>%
  subset(>0) %>%
  names
pepSeqFms$tryphprp <- setdiff(pepSeqMapped_to_clean, pepSeqMapped_to_with_marks)
print(pepSeqFms$tryphprp)

## [1] "FFAAPEK" "ELAFLKRAQEIGLEPYNEYHGKKK"
## [3] "VVQEGNTNVKK" "WTPIDLPSSEITFVQGIQTVTGAGDPSMK"
## [5] "IIQNFQINTVFEDLDEIMQTQVQR" "KSSKACEEERRKR"
## [7] "LINDLTNDK" "LISELTSEK"
## [9] "IVENFNK" "LSQEHLISYISL"
## [11] "LINDLTNDKANLK"
```

```
meta_tryph_hprp <- tryphprpPrj.fms %>%
  apply_filter('pepSeq %in% pepSeqFms$tryphprp') %>%
  psms %>%
  select(spectrumFile, MS.GF.SpecEValue, mme.ppm, spectrumID, chargeState, peptide) %>%
  rename(SpecEValue = MS.GF.SpecEValue, charge = chargeState, `MME (ppm)` = mme.ppm) %>%
  mutate(spectrumFile = sub('_msgfplus.mzid.gz', '', spectrumFile))
library(xtable)
print(xtable(meta_tryph_hprp, display = c('d', 's', 'e', 'f', 's', 'd', 's')),
  include.rownames=FALSE,
  comment = FALSE,
  size='scriptsize',
  floating = F)
```

spectrumFile	SpecEValue	MME (ppm)	spectrumID	charge	peptide
Euplotes_1_HPRP_1_04_17Nov09_Falcon_09-09-14	7.58e-11	0.08	index=3031	1	R.FFAAPEK.I
Euplotes_1_HPRP_1_04_17Nov09_Falcon_09-09-14	2.44e-09	0.00	index=3046	2	R.FFAAPEK.I
Euplotes_1_HPRP_1_05_17Nov09_Falcon_09-09-14	1.46e-09	5.31	index=8245	3	R.ELAFLKRAQEIGLEPYNEYHGKKK.T
Euplotes_1_HPRP_1_06_17Nov09_Falcon_09-09-14	5.54e-10	2.21	index=759	2	K.VVQEGNTNVKK.L
Euplotes_1_HPRP_1_08_17Nov09_Falcon_09-09-14	5.93e-22	2.11	index=8644	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_HPRP_1_08_17Nov09_Falcon_09-09-14	2.18e-21	0.78	index=8638	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_HPRP_1_08_17Nov09_Falcon_09-09-14	3.05e-21	2.11	index=8646	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_HPRP_1_08_17Nov09_Falcon_09-09-14	4.19e-16	0.82	index=8639	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_HPRP_1_09_17Nov09_Falcon_09-09-14	1.19e-21	0.70	index=8806	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_HPRP_1_09_17Nov09_Falcon_09-09-14	1.20e-21	1.57	index=8812	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_HPRP_1_09_17Nov09_Falcon_09-09-14	5.49e-20	1.64	index=8802	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_HPRP_1_09_17Nov09_Falcon_09-09-14	4.33e-15	1.53	index=8810	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A
Euplotes_1_HPRP_1_16_22Nov09_Falcon_09-09-14	4.51e-21	0.33	index=10684	2	K.IIQNFQINTVFEDLDEIMQTQVQR.H
Euplotes_1_HPRP_1_16_22Nov09_Falcon_09-09-14	1.36e-11	1.25	index=10678	3	K.IIQNFQINTVFEDLDEIMQTQVQR.H
Euplotes_1_HPRP_1_18_17Nov09_Falcon_09-09-15	5.08e-09	2.64	index=13785	2	K.KSSKACEEERRKR.E
Euplotes_1_HPRP_1_20_17Nov09_Falcon_09-09-15	1.91e-11	0.00	index=3425	1	K.LINDLTNDK.A
Euplotes_1_HPRP_1_22_17Nov09_Falcon_09-09-15	6.65e-11	1.67	index=3600	2	K.LISELTSEK.S
Euplotes_1_HPRP_1_22_17Nov09_Falcon_09-09-15	2.55e-10	0.78	index=3602	1	K.LISELTSEK.S
Euplotes_1_HPRP_1_22_17Nov09_Falcon_09-09-15	1.89e-09	0.49	index=2595	2	K.IVENFNK.I
Euplotes_1_HPRP_1_23_17Nov09_Falcon_09-09-15	3.01e-13	1.01	index=2200	2	K.LSQEHLISYISL.L
Euplotes_1_HPRP_1_24_17Nov09_Falcon_09-09-15	2.45e-16	1.41	index=2709	2	K.LINDLTNDKANLK.D

2.2.3 Glu-C Digest Fractionated by HPRP

All the processing steps are conceptually the same as in the section above. The only substantial difference is the specification of the enzyme digestion rule.

```
gluchprp <- grep('Euplotes_1_HPRP_2_.*msgfplus.mzid.gz', repoFiles, value=T)
gluchprpPrj <- MSnID()
system.time(gluchprpPrj <- read_mzIDs(gluchprpPrj, gluchprp, backend = 'mzR'))
##      user  system elapsed
##   5.821    4.043   34.076
```

```
gluchprpPrj <- assess termini(gluchprpPrj, validCleavagePattern="E\\.[^P]")
gluchprpPrj <- apply_filter(gluchprpPrj, "numIrregCleavages == 0")
```

```
gluchprpPrj.main <- apply_filter(gluchprpPrj, "!grepl('comp', accession)")
gluchprpPrj.fmsh <- apply_filter(gluchprpPrj, "grepl('comp', accession)")
gluchprpPrj.fmsh <- apply_filter(gluchprpPrj.fmsh,
                                "!(peptide %in% peptides(gluchprpPrj.main))")

show(gluchprpPrj.fmsh)
## MSnID object
## Working directory: "."
## #Spectrum Files: 24
## #PSMs: 555 at 67 % FDR
## #peptides: 440 at 80 % FDR
## #accessions: 297 at 89 % FDR
```

```
gluchprpPrj.fmsh$mme.ppm <- abs(mass_measurement_error(gluchprpPrj.fmsh))
gluchprpPrj.fmsh$score <- -log10(gluchprpPrj.fmsh$`MS.GF.SpecEValue`)
gluchprpPrj.fmsh <- apply_filter(gluchprpPrj.fmsh, "mme.ppm < 10")

filtr <- MSnIDFilter(gluchprpPrj.fmsh)
filtr$mme.ppm <- list(comparison="<", threshold=5.0)
filtr$score <- list(comparison=">", threshold=8.0)
filtr.grid <- optimize_filter(filtr, gluchprpPrj.fmsh, fdr.max=0.05,
                             method="Grid", level="peptide", n.iter=20000)
evaluate_filter(gluchprpPrj.fmsh, filtr.grid)
##              fdr  n
## PSM          0.02222222 46
## peptide      0.03448276 30
## accession    0.05000000 21
```

```
filtr.sann <- optimize_filter(filtr.grid, gluchprpPrj.fmsh, fdr.max=0.05,
                             method="SANN", level="peptide", n.iter=20000)
evaluate_filter(gluchprpPrj.fmsh, filtr.sann)
##              fdr  n
## PSM          0.02222222 46
## peptide      0.03448276 30
## accession    0.05000000 21
```

```
gluchprpPrj.fmsh <- apply_filter(gluchprpPrj.fmsh, filtr.sann)
show(gluchprpPrj.fmsh)
## MSnID object
## Working directory: "."
## #Spectrum Files: 18
## #PSMs: 46 at 2.2 % FDR
```



```
## #peptides: 30 at 3.4 % FDR
## #accessions: 21 at 5 % FDR

library(dplyr)
pepSeq <- unique(gluchprpPrj.fmsH$pepSeq)
pepSeqMapped_to_clean <- pepSeq %>%
  apply(grep, x=fasta_clean) %>%
  apply(length) %>%
  subset(>0) %>%
  names
pepSeqMapped_to_with_marks <- pepSeq %>%
  apply(grep, x=fasta_marks) %>%
  apply(length) %>%
  subset(>0) %>%
  names
pepSeqFmsH_gluchprp <- setdiff(pepSeqMapped_to_clean, pepSeqMapped_to_with_marks)
print(pepSeqFmsH_gluchprp)

## [1] "NFNKITGKEQEEEE"          "SVNRENLDNEKLINDLTNDKANLKDIVFDLMFE"
## [3] "NLDNEKLINDLTNDKANLKDIVFDLMFE"    "NKIRFFAAPEKIFE"
## [5] "MQDEEILKSIEESKLEQEKEEEKNE"      "VYLGLMEEYE"

meta_gluc_hprp <- gluchprpPrj.fmsH %>%
  apply_filter('pepSeq %in% pepSeqFmsH_gluchprp') %>%
  psms %>%
  select(spectrumFile, MS.GF.SpecEValue, mme.ppm, spectrumID, chargeState, peptide) %>%
  rename(SpecEValue = MS.GF.SpecEValue, charge = chargeState, `MME (ppm)`=mme.ppm) %>%
  mutate(spectrumFile = sub('_msgfplus.mzid.gz', '', spectrumFile))
library(xtable)
print(xtable(meta_gluc_hprp, display = c('d', 's', 'e', 'f', 's', 'd', 's')),
  include.rownames=FALSE,
  comment = FALSE,
  size='scriptsize',
  floating = F)
```

spectrumFile	SpecEValue	MME (ppm)	spectrumID	charge	peptide
Euplotes_1_HPRP_2_06_22Nov09_Falcon_09-09-15	6.80e-07	2.95	index=13369	2	E.NFNKITGKEQEEEE.Y
Euplotes_1_HPRP_2_08_25Nov09_Falcon_09-09-15	3.78e-17	0.19	index=9982	3	E.SVNRENLDNEKLINDLTNDKANLKDIVFDLMFE.K
Euplotes_1_HPRP_2_08_25Nov09_Falcon_09-09-15	3.33e-07	0.57	index=9974	4	E.SVNRENLDNEKLINDLTNDKANLKDIVFDLMFE.K
Euplotes_1_HPRP_2_09_17Nov09_Falcon_09-09-17	5.74e-16	0.44	index=10771	3	E.NLDNEKLINDLTNDKANLKDIVFDLMFE.K
Euplotes_1_HPRP_2_09_17Nov09_Falcon_09-09-17	5.03e-07	1.11	index=10770	4	E.NLDNEKLINDLTNDKANLKDIVFDLMFE.K
Euplotes_1_HPRP_2_12_17Nov09_Falcon_09-09-17	2.09e-09	0.43	index=3933	3	E.NKIRFFAAPEKIFE.T
Euplotes_1_HPRP_2_12_17Nov09_Falcon_09-09-17	1.62e-07	0.07	index=3930	2	E.NKIRFFAAPEKIFE.T
Euplotes_1_HPRP_2_15_17Nov09_Falcon_09-09-17	2.83e-07	1.61	index=1758	2	E.MQDEEILKSIEESKLEQEKEEEKNE.E
Euplotes_1_HPRP_2_21_22Nov09_Falcon_09-09-17	2.17e-07	0.10	index=6671	1	E.VYLGLMEEYE.A
Euplotes_1_HPRP_2_22_22Nov09_Falcon_09-09-17	2.12e-08	0.88	index=6753	1	E.VYLGLMEEYE.A

2.3 Compendium of Peptides Covering Frameshift Locations

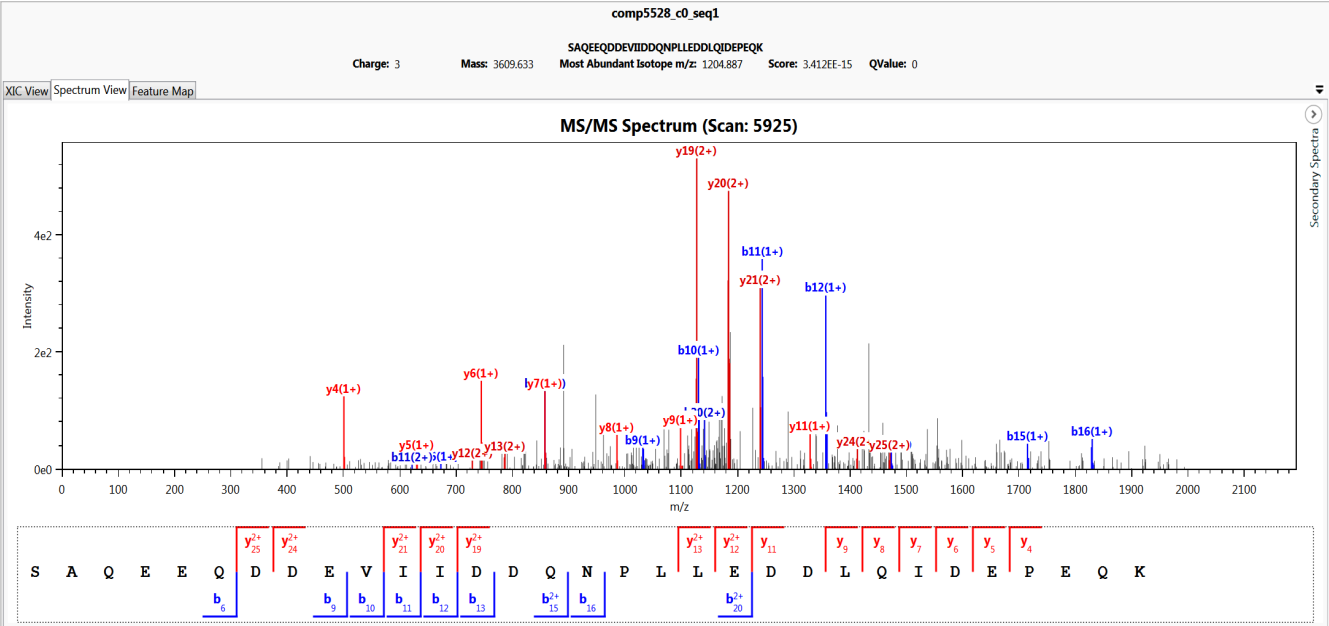
Final set of peptides and corresponding references to LC-MS/MS datasets and spectra. Overall, **4**, **11**, and **6** unique peptide sequences spanning the frameshift sites were identified in trypsin/SCX, trypsin/HPRP, and 'Glu-C/HPRP' experiments, respectively.

spectrumFile	SpecEValue	MME (ppm)	spectrumID	charge	peptide	experiment
Euplotes_1_SCX_10_13Nov09_Falcon_09-09-14	3.41e-15	0.30	index=6106	3	K.SAQEEQDDVEIDDQNPILLEDLDLQIDEPEQK.V	trypsin/SCX
Euplotes_1_SCX_10_13Nov09_Falcon_09-09-14	3.41e-15	0.30	index=6106	3	K.SAQEEQDDVEIDDQNPILLEDLDLQIDEPEQK.V	trypsin/SCX
Euplotes_1_SCX_12_13Nov09_Falcon_09-09-14	1.53e-21	0.08	index=8908	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/SCX
Euplotes_1_SCX_12_13Nov09_Falcon_09-09-14	1.07e-20	1.10	index=8896	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/SCX
Euplotes_1_SCX_12_13Nov09_Falcon_09-09-14	7.29e-19	1.10	index=8897	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/SCX
Euplotes_1_SCX_12_13Nov09_Falcon_09-09-14	2.17e-15	0.94	index=8895	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/SCX
Euplotes_1_SCX_18_13Nov09_Falcon_09-09-15	9.27e-17	0.11	index=5912	2	K.ESNHNNDITNKNEIAYLR.Y	trypsin/SCX
Euplotes_1_SCX_20_13Nov09_Falcon_09-09-15	2.23e-11	0.70	index=10317	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/SCX
Euplotes_1_SCX_22_13Nov09_Falcon_09-09-15	4.36e-10	3.76	index=9720	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/SCX
Euplotes_1_SCX_23_13Nov09_Falcon_09-09-15	2.47e-09	1.64	index=9440	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/SCX
Euplotes_1_SCX_24_13Nov09_Falcon_09-09-15	3.42e-10	8.85	index=2127	3	R.KKKQEENLKR.K	trypsin/SCX
Euplotes_1_HPRP_1_04_17Nov09_Falcon_09-09-14	7.58e-11	0.08	index=3031	1	R.FFAAPEK.I	trypsin/HPRP
Euplotes_1_HPRP_1_04_17Nov09_Falcon_09-09-14	2.44e-09	0.00	index=3046	2	R.FFAAPEK.I	trypsin/HPRP
Euplotes_1_HPRP_1_05_17Nov09_Falcon_09-09-14	1.46e-09	5.31	index=8245	3	R.ELAFKRAQEIGLEPYNEYHGKKK.T	trypsin/HPRP
Euplotes_1_HPRP_1_06_17Nov09_Falcon_09-09-14	5.54e-10	2.21	index=759	2	K.VVQEGNTNVKK.L	trypsin/HPRP
Euplotes_1_HPRP_1_08_17Nov09_Falcon_09-09-14	5.93e-22	2.11	index=8644	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/HPRP
Euplotes_1_HPRP_1_08_17Nov09_Falcon_09-09-14	2.18e-21	0.78	index=8638	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/HPRP
Euplotes_1_HPRP_1_08_17Nov09_Falcon_09-09-14	3.05e-21	2.11	index=8646	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/HPRP
Euplotes_1_HPRP_1_08_17Nov09_Falcon_09-09-14	4.19e-16	0.82	index=8639	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/HPRP
Euplotes_1_HPRP_1_09_17Nov09_Falcon_09-09-14	1.19e-21	0.70	index=8806	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/HPRP
Euplotes_1_HPRP_1_09_17Nov09_Falcon_09-09-14	1.20e-21	1.57	index=8812	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/HPRP
Euplotes_1_HPRP_1_09_17Nov09_Falcon_09-09-14	5.49e-20	1.64	index=8802	2	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/HPRP
Euplotes_1_HPRP_1_09_17Nov09_Falcon_09-09-14	4.33e-15	1.53	index=8810	3	R.WTPIDLPSSEITFVQGIQTVTGAGDPSMK.A	trypsin/HPRP
Euplotes_1_HPRP_1_16_22Nov09_Falcon_09-09-14	4.51e-21	0.33	index=10684	2	K.IIQNFQINTVFEDLDEIMQTQVQR.H	trypsin/HPRP
Euplotes_1_HPRP_1_16_22Nov09_Falcon_09-09-14	1.36e-11	1.25	index=10678	3	K.IIQNFQINTVFEDLDEIMQTQVQR.H	trypsin/HPRP
Euplotes_1_HPRP_1_18_17Nov09_Falcon_09-09-15	5.08e-09	2.64	index=13785	2	K.KSKACEEERRK.R	trypsin/HPRP
Euplotes_1_HPRP_1_20_17Nov09_Falcon_09-09-15	1.91e-11	0.00	index=3425	1	K.LINDLTNDK.A	trypsin/HPRP
Euplotes_1_HPRP_1_22_17Nov09_Falcon_09-09-15	6.65e-11	1.67	index=3600	2	K.LISELTSEK.S	trypsin/HPRP
Euplotes_1_HPRP_1_22_17Nov09_Falcon_09-09-15	2.55e-10	0.78	index=3602	1	K.LISELTSEK.S	trypsin/HPRP
Euplotes_1_HPRP_1_22_17Nov09_Falcon_09-09-15	1.89e-09	0.49	index=2595	2	K.IVENFNK.I	trypsin/HPRP
Euplotes_1_HPRP_1_23_17Nov09_Falcon_09-09-15	3.01e-13	1.01	index=2200	2	K.LSQEHLISYIS.R	trypsin/HPRP
Euplotes_1_HPRP_1_24_17Nov09_Falcon_09-09-15	2.45e-16	1.41	index=2709	2	K.LINDLTNDKANL.D	trypsin/HPRP
Euplotes_1_HPRP_2_06_22Nov09_Falcon_09-09-15	6.80e-07	2.95	index=13369	2	E.NFNKITGKEQEEEEE.Y	Glu-C/HPRP
Euplotes_1_HPRP_2_08_25Nov09_Falcon_09-09-15	3.78e-17	0.19	index=9982	3	E.SVNRENLDNEKLINDLTNDKANLKDIVDFLMFE.K	Glu-C/HPRP
Euplotes_1_HPRP_2_08_25Nov09_Falcon_09-09-15	3.33e-07	0.57	index=9974	4	E.SVNRENLDNEKLINDLTNDKANLKDIVDFLMFE.K	Glu-C/HPRP
Euplotes_1_HPRP_2_09_17Nov09_Falcon_09-09-17	5.74e-16	0.44	index=10771	3	E.NLDNEKLINDLTNDKANLKDIVDFLMFE.K	Glu-C/HPRP
Euplotes_1_HPRP_2_09_17Nov09_Falcon_09-09-17	5.03e-07	1.11	index=10770	4	E.NLDNEKLINDLTNDKANLKDIVDFLMFE.K	Glu-C/HPRP
Euplotes_1_HPRP_2_12_17Nov09_Falcon_09-09-17	2.09e-09	0.43	index=3933	3	E.NKIRFFAAPEKIFE.T	Glu-C/HPRP
Euplotes_1_HPRP_2_12_17Nov09_Falcon_09-09-17	1.62e-07	0.07	index=3930	2	E.NKIRFFAAPEKIFE.T	Glu-C/HPRP
Euplotes_1_HPRP_2_15_17Nov09_Falcon_09-09-17	2.83e-07	1.61	index=1758	2	E.MQDEILKSIEESKLEQEQQEEKKNE.E	Glu-C/HPRP
Euplotes_1_HPRP_2_21_22Nov09_Falcon_09-09-17	2.17e-07	0.10	index=6671	1	E.VYLGLMEEYE.A	Glu-C/HPRP
Euplotes_1_HPRP_2_22_22Nov09_Falcon_09-09-17	2.12e-08	0.88	index=6753	1	E.VYLGLMEEYE.A	Glu-C/HPRP

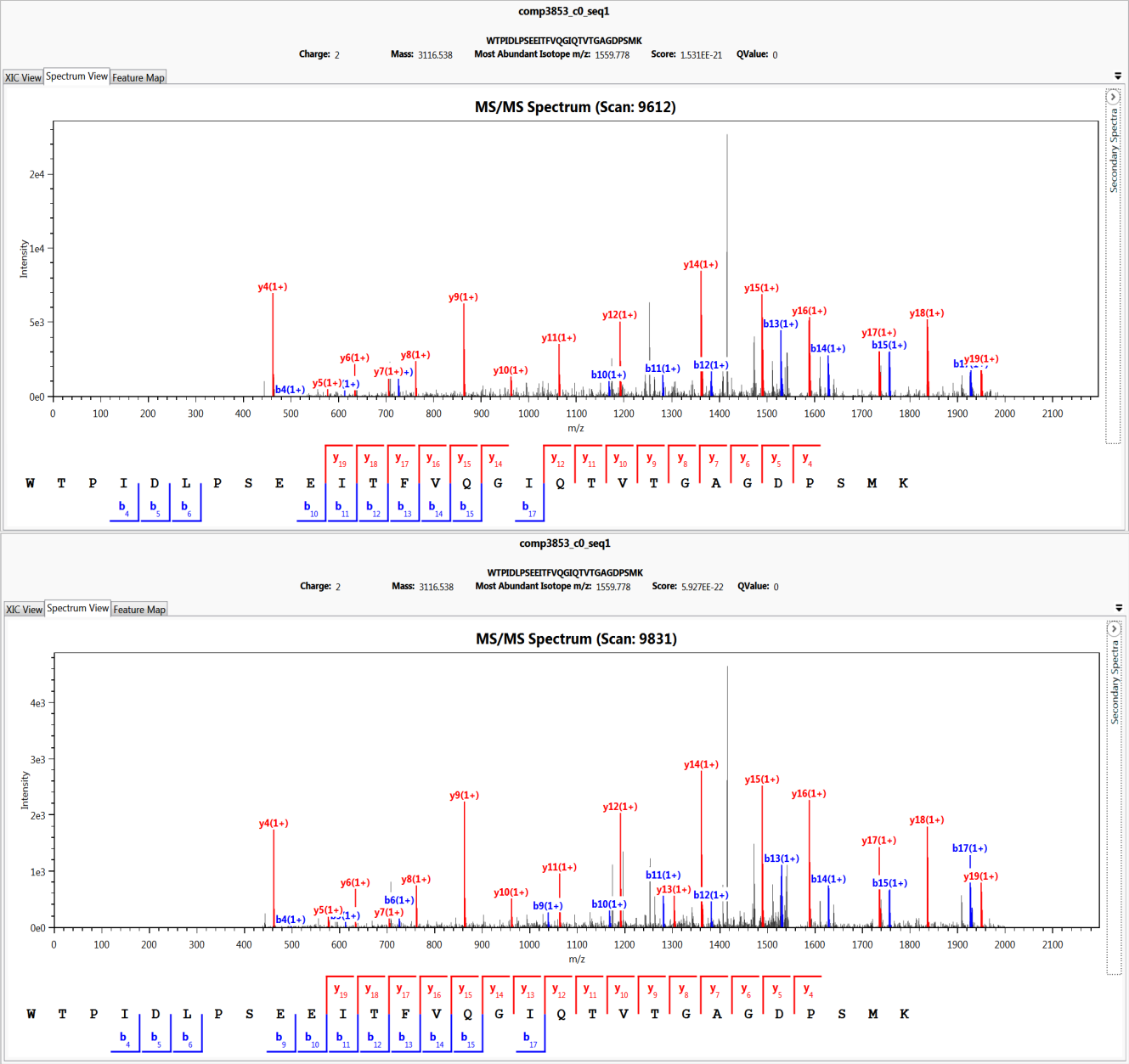
3 Manual Validation

Manual validation was performed by LCMSSpectator. The spectra that have passed the consensus opinion of 5 independed experts are shown below. Necessary raw and mzIdenML files to reproduce the analysis are available at <http://dx.doi.org/10.6019/PXD004333>. Note, the MS/MS scan number is not the same identifier as spectrumID in the table above.

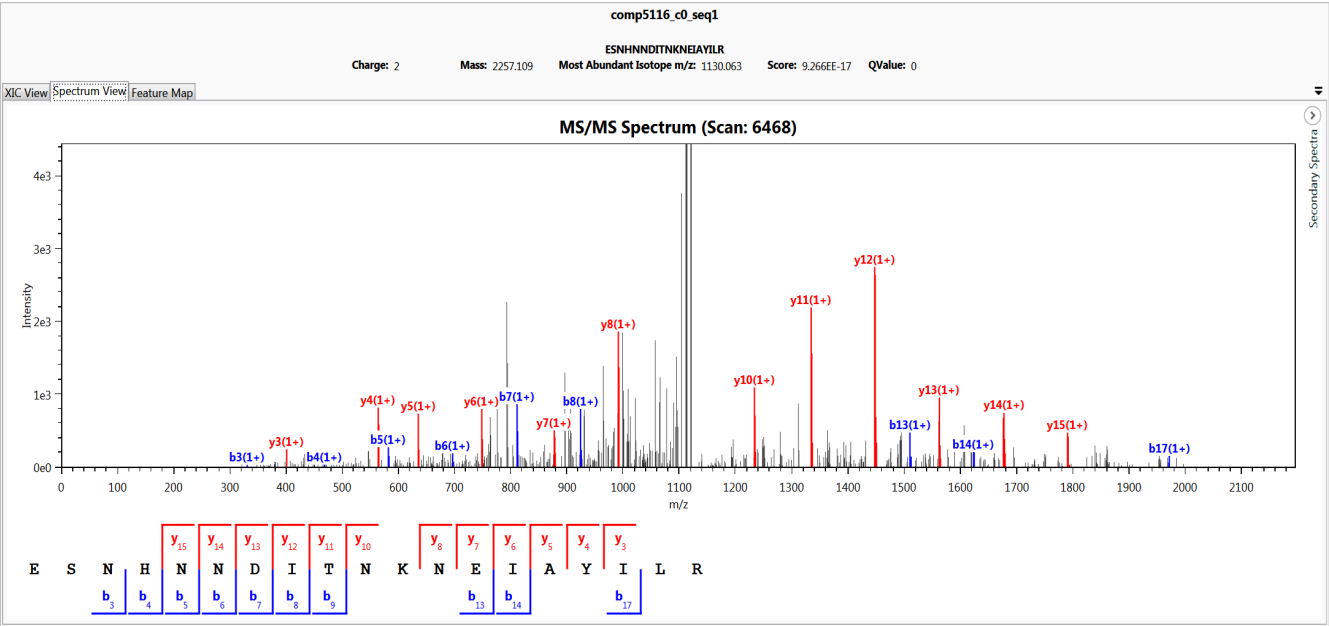
SAQEEQDDEVIIDDQNPLLEDDLQIDEPEQK



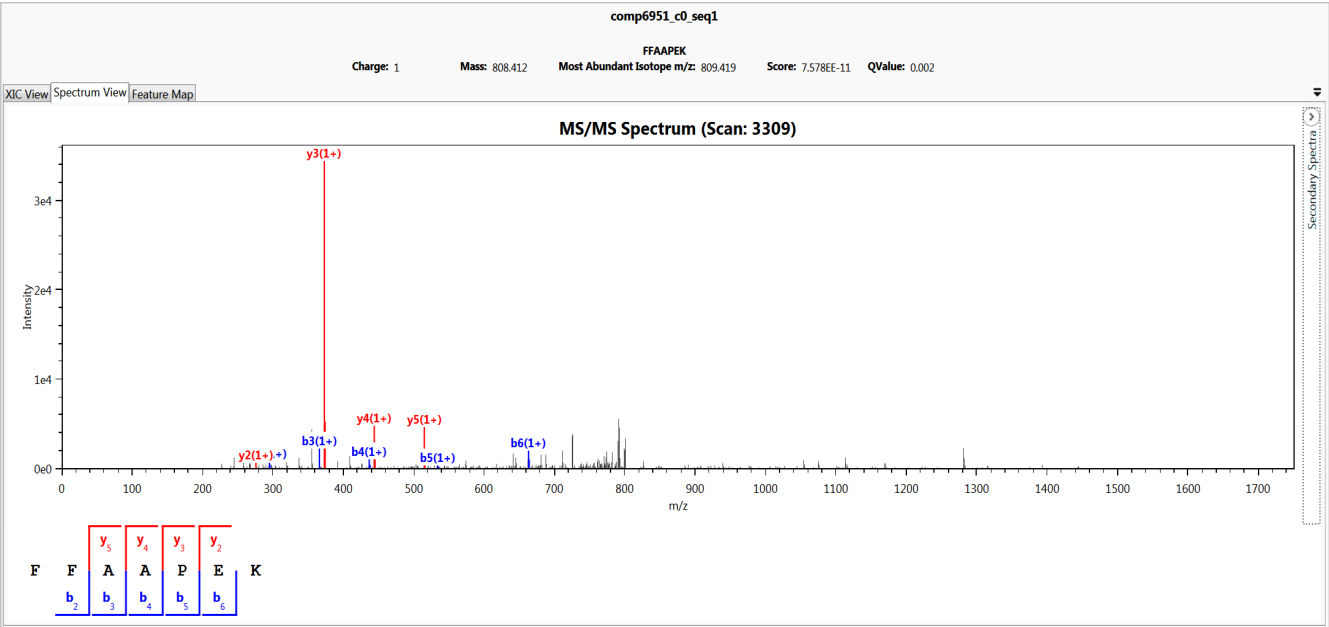
WTPIDLPSSEITFVQGIQTVTGAGDPSMK



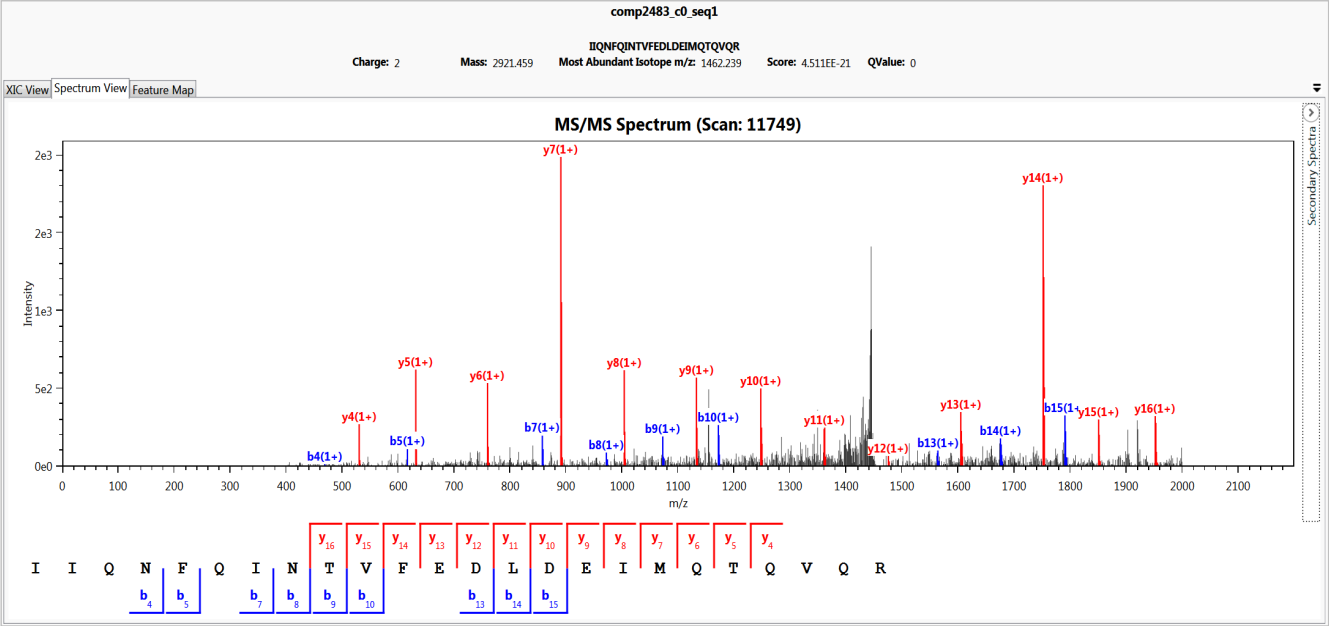
ESNHNNNDITNKNEIAYILR



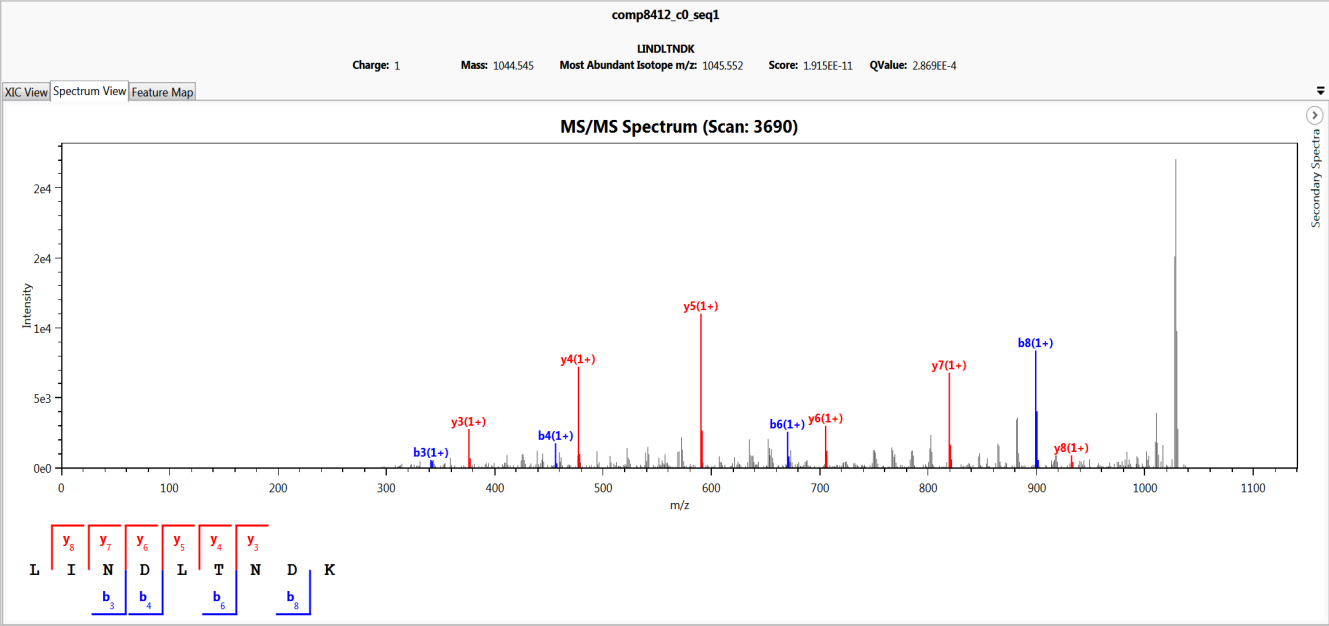
FFAAPEK



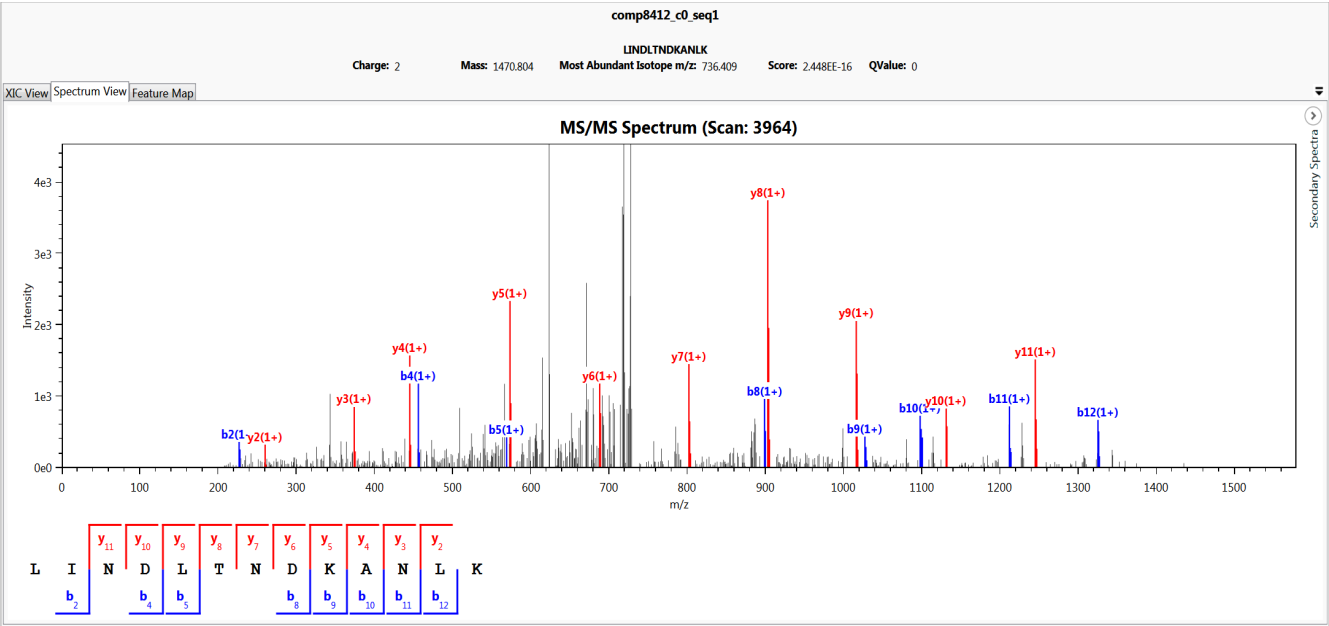
IIQNFQINTVFEDLDEIMQTQVQR



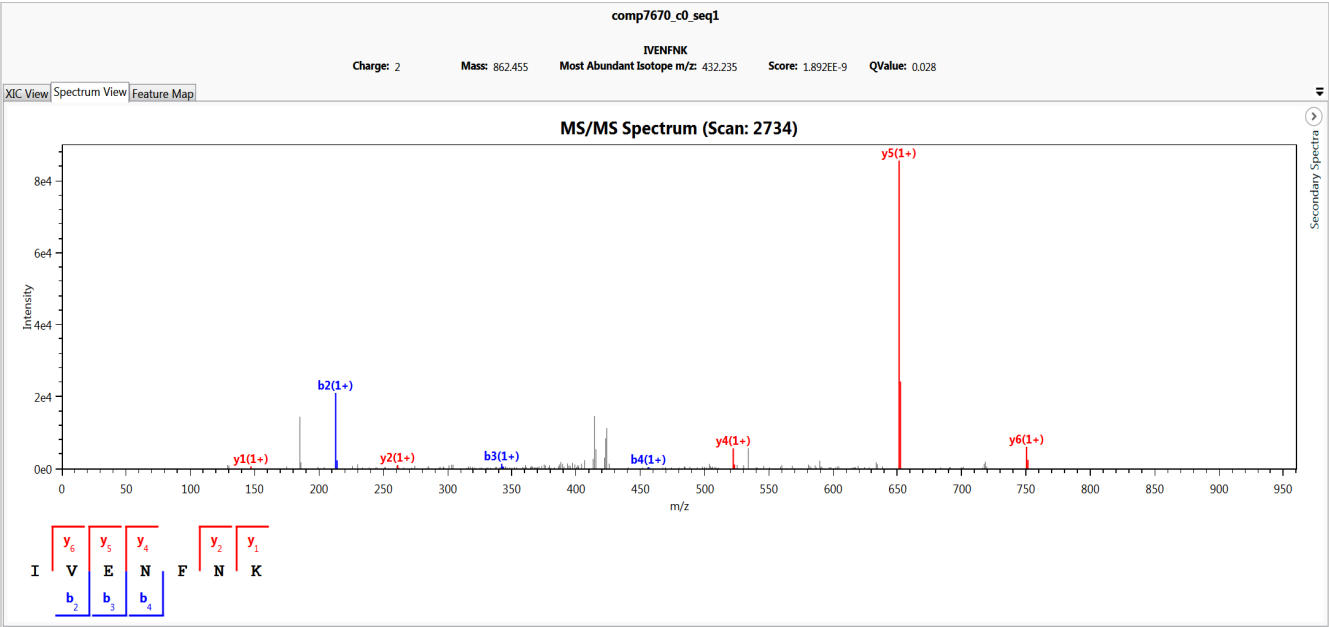
LINDLTNDK



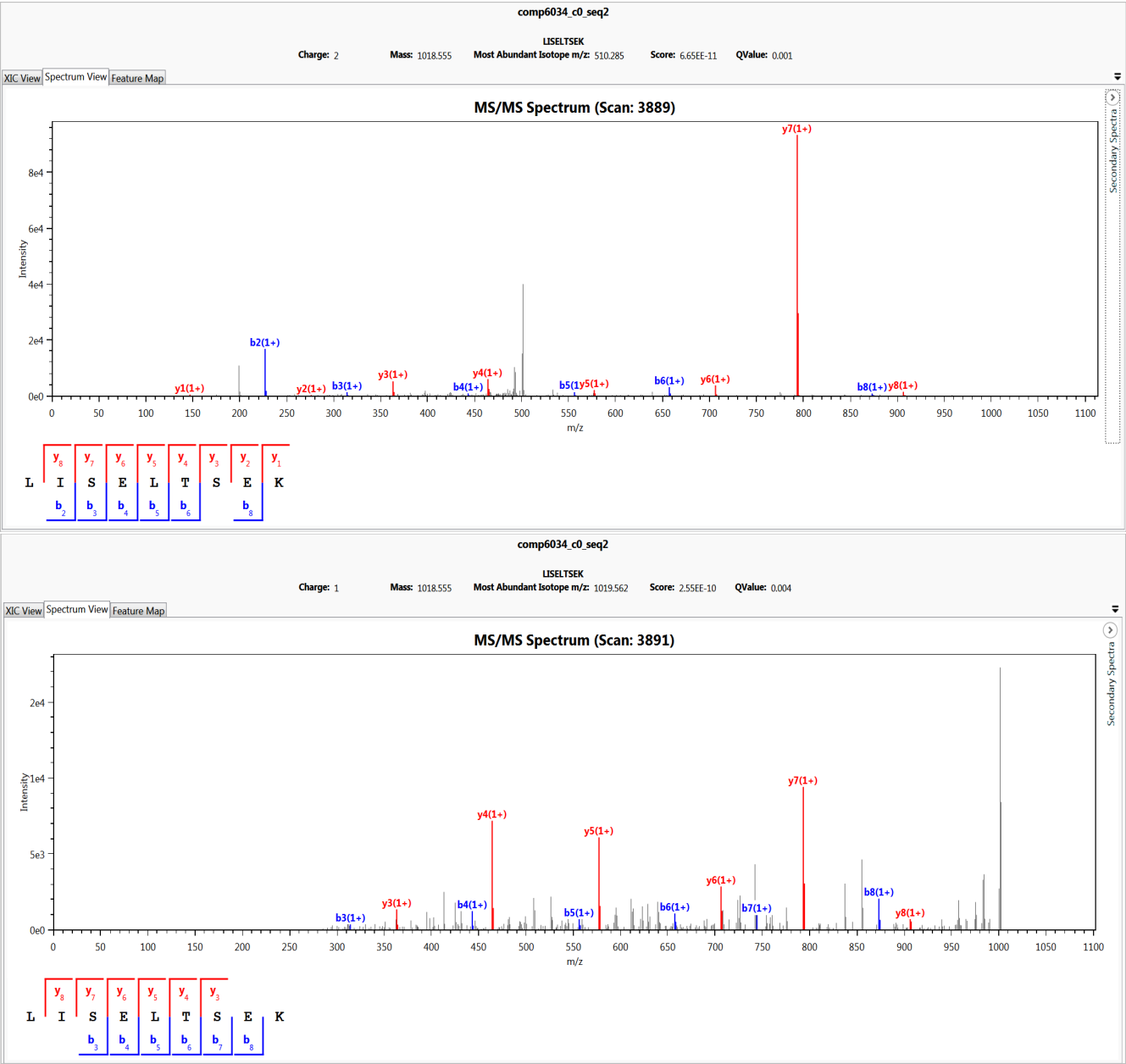
LINDLTNDKANLK



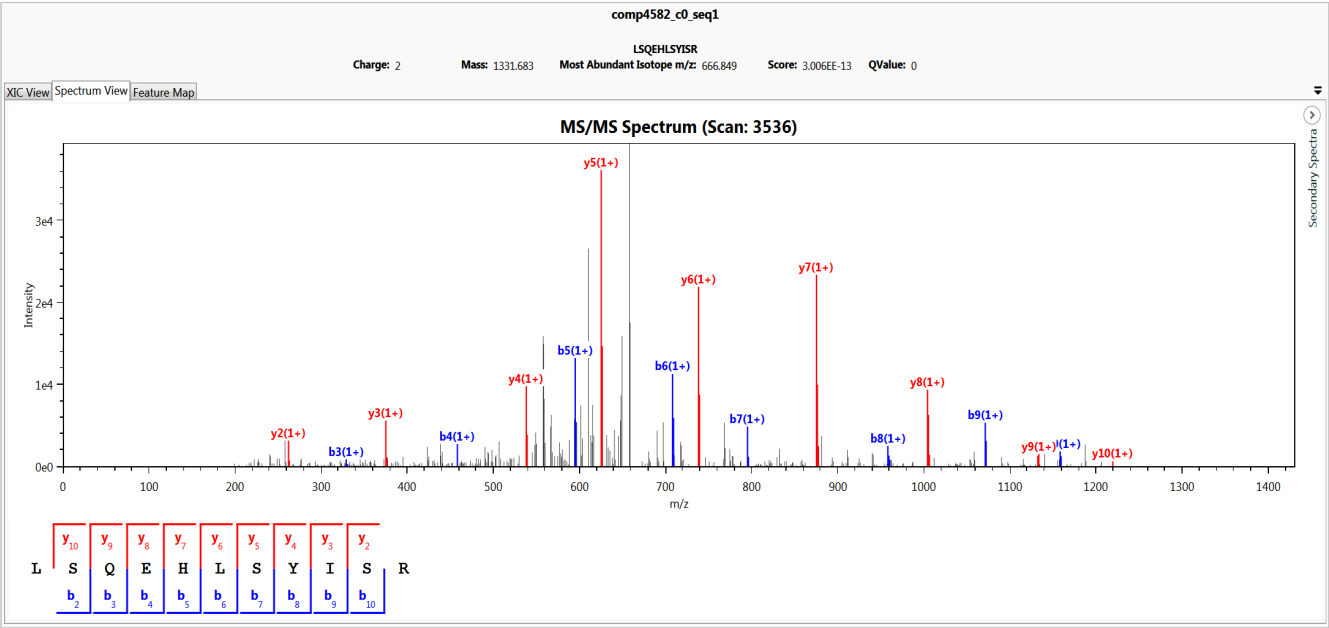
IVENFNK



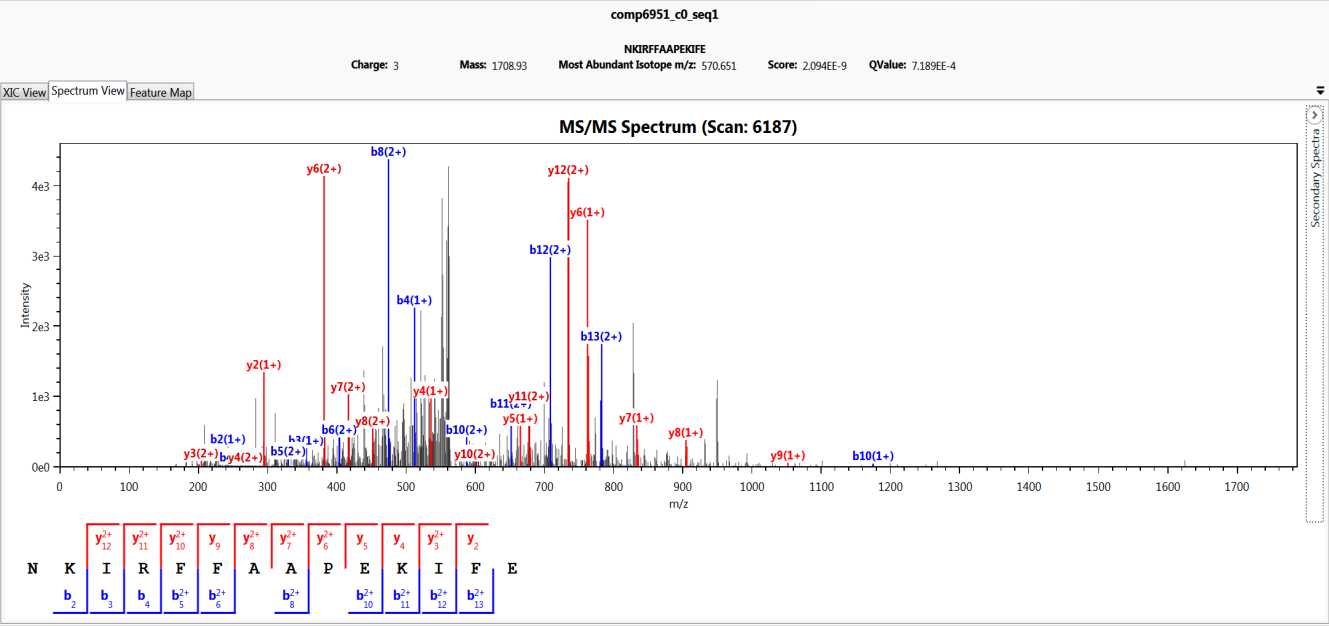
LISELTSEK



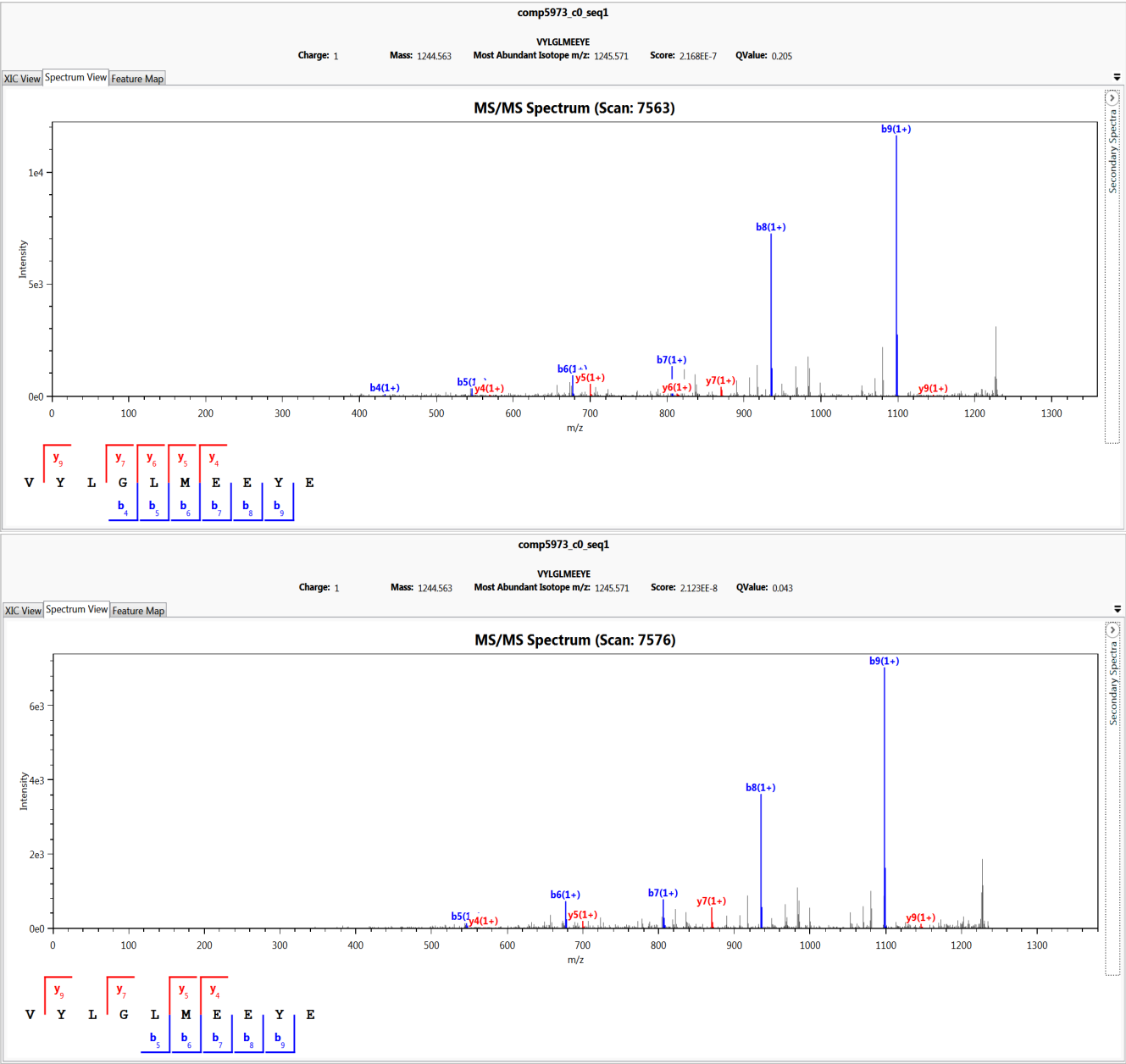
LSQEHLSYISR



NKIRFFAAPEKIFE



VYLGLMEEYE



4 Session Information

All software and respective versions used in this document, as returned by sessionInfo() are detailed below.

- R version 3.2.4 (2016-03-10), x86_64-apple-darwin13.4.0
- Locale: C/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, stats4, utils
- Other packages: BiocGenerics 0.16.1, BiocStyle 1.8.0, Biostrings 2.38.4, IRanges 2.4.8, MSnID 1.7.3, Rcpp 0.12.5, S4Vectors 0.8.11, XVector 0.10.0, dplyr 0.4.3.9000, knitr 1.12.3, rpx 1.6.0, xtable 1.8-2
- Loaded via a namespace (and not attached): Biobase 2.30.0, BiocInstaller 1.20.3, BiocParallel 1.4.3, DBI 0.4-1, MALDIquant 1.14, MSnbase 1.18.1, ProtGenerics 1.2.1, R.cache 0.12.0, R.methodsS3 1.7.1, R.oo 1.20.0, R.utils 2.3.0, R6 2.1.2, RCurl 1.95-4.8, XML 3.98-1.4, affy 1.48.0, affyio 1.40.0, assertthat 0.1, bitops 1.0-6, chron 2.3-47, codetools 0.2-14, colorspace 1.2-6, compiler 3.2.4, data.table 1.9.6, digest 0.6.9, doParallel 1.0.10, evaluate 0.8.3, foreach 1.4.3, formatR 1.3, futile.logger 1.4.1, futile.options 1.0.0, ggplot2 2.1.0, grid 3.2.4, gtable 0.2.0, highr 0.5.1, htmltools 0.3.5, impute 1.44.0, iterators 1.0.8, lambda.r 1.1.7, lattice 0.20-33, lazyeval 0.2.0, limma 3.26.9, magrittr 1.5, munsell 0.4.3, mzID 1.8.0, mzR 2.4.1, pcaMethods 1.60.0, plyr 1.8.4, preprocessCore 1.32.0, reshape2 1.4.1, rmarkdown 0.9.5, scales 0.4.0, stringi 1.1.1, stringr 1.0.0, tools 3.2.4, vsn 3.38.0, yaml 2.1.13, zlibbioc 1.16.0