

# On Finding putative PTM (pPTM) Marker Ion in HCD scans using PTM\_MarkerFinder

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## Abstract

Glycopeptides as well as acetylated, methylated and other modified peptides release specific fragment ions during CID (collision-induced dissociation) and HCD (higher energy collisional dissociation) fragmentation. These fragment ions can be used to validate the presence of the PTM (post translational modifications) on the peptides. **PTM\_MarkerFinder**, an R function of the **protViz** package that takes advantage of such marker ions. **PTM\_MarkerFinder** scans the MS/MS spectra in the output of a peptide spectrum match search, e.g., Mascot, for marker ions specific for selected PTMs.

While the software tool has been described by [Nanni, Panse, Gehrig, Mueller, Grossmann, and Schlapbach \(2013\)](#) here we provide a step-by-step guide on how the software can be used.

**Keywords:** MarkerFinder, putative post translational modifications, R.

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## 1. Howto get the software and data

The method for finding the marker ions is contained in the R package **protViz** available through CRAN using <https://cran.r-project.org/package=protViz>. The package requires R ([R Development Core Team 2008](#)) installed.

The minimal data structure requirement for the **PTM\_MarkerFinder** function looks as follow.

```
R> library(protViz)
R> data(HexNAc)
R> str(HexNAc[[1]], nchar.max = 30)

List of 12
 $ peptideSequence   : chr "STMQELNSR"
 $ mascotScore       : num 49.5
 $ modification      : chr "000000000000"
 $ MonoisotopicAAmass: num [1:9] 0 0 0 0 0 0 0 0 0
 $ proteinInformation: chr "zz|ZZ_FGCZCont0219|"
 $ title              : chr "NGlycoFASP_NH" | __truncated__
```

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```
$ pepmass      : num 533
$ charge       : num 2
$ scans        : num 2659
$ rtinseconds  : num 1846
$ mZ           : num [1:150] 101 104 105 110 112 ...
$ intensity    : num [1:150] 369.3 2860 37.3 103.8 190.7 ...
```

Here we have listed the HexNAc data which is included in **protViz**.

**protViz** also provides and perl script `protViz_mascotDat2RData.pl`<sup>1</sup> taking mascot server dat files as input and producing RData output.

```
$ /usr/local/lib/R/site-library/protViz/exec/protViz_mascotDat2RData.pl \
-d=/usr/local/mascot/data/20130116/F178287.dat \
-m=$HOME/mod_file
```

`mascotDat2RData.pl` requires the mascot server `mod_file` keeping all the configured modification of the mascot server.

In theory **PTM\_MarkerFinder** can process the output of any search engine for peptide identification. It is up to the R user writing a wrapper script converting the output of any particular peptide identification search engine to the data structure listed above.

## 2. Finding the Marker Ions

### 2.1. HexNAc – Example

**PTM\_MarkerFinder** can search for any Marker ion series. The next lines define the `HexNAc_MarkerIons`.

```
R> HexNAc_MarkerIons <- c(126.05495, 138.05495, 144.06552,
+     168.06552, 186.07608, 204.08665)
```

The lines below configure the modification information used by the search engine. The HexNAc modification below is described on unimod [http://www.unimod.org/modifications\\_view.php?editid=43](http://www.unimod.org/modifications_view.php?editid=43).

```
R> ptm.0 <- cbind(AA = "-",
+     mono = 0.0, avg = 0.0, desc = "unmodified", unimodAccID = NA)
R> ptm.1 <- cbind(AA='N',
+     mono = 317.122300, avg = NA, desc = "HexNAc",
+     unimodAccID=2)
R> ptm.2 <- cbind(AA='M',
+     mono = 147.035400, avg = NA, desc = "Oxidation",
+     unimodAccID=1)
R> m <- as.data.frame(rbind(ptm.0, ptm.1, ptm.2))
```

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<sup>1</sup>The prefix `protViz_` is used to benefit from the bash tab completion.

`PTM_MarkerFinder` is called.

```
R> S <- PTM_MarkerFinder(data = HexNAc,
+                         modification = m$mono,
+                         modificationName = m$desc,
+                         minMarkerIntensityRatio = 3,
+                         itol_ppm = 20,
+                         mZmarkerIons = HexNAc_MarkerIons)
```

The content of `S` can be seen in the Table below.

scans	mZ	markerIonMZ	markerIonIntensity	markerIonMzError	markerIonPpmError	query	pepmass	peptideSequence	modification
3687	126.06	126.05	9945.00	-0.00	-0.6425764947898	4	713.36	IMNVTTDSLTK	0001000000000
3687	138.06	138.05	1933.00	-0.00	-2.49175522390729	4	713.36	IMNVTTDSLTK	0001000000000
3687	144.07	144.07	412.30	-0.00	-1.59649326794302	4	713.36	IMNVTTDSLTK	0001000000000
3687	168.07	168.07	810.20	-0.00	-2.36811844277867	4	713.36	IMNVTTDSLTK	0001000000000
3687	204.09	204.09	3273.00	-0.00	-1.74435407225623	4	713.36	IMNVTTDSLTK	0001000000000
2540	126.06	126.05	2945.00	-0.00	0.825036336847078	6	490.56	HSFNGNQSTFK	0000001000000
2540	138.06	138.05	759.20	-0.00	-10.3725737215287	6	490.56	HSFNGNQSTFK	0000001000000
2540	144.07	144.07	195.40	-0.00	-0.118001850879316	6	490.56	HSFNGNQSTFK	0000001000000
2540	168.07	168.07	262.90	-0.00	0.916308466469431	6	490.56	HSFNGNQSTFK	0000001000000
2540	186.08	186.08	188.50	-0.00	-2.95577150125756	6	490.56	HSFNGNQSTFK	0000001000000
2540	204.09	204.09	998.40	-0.00	-1.5189603491234	6	490.56	HSFNGNQSTFK	0000001000000
4393	126.06	126.05	13620.00	-0.00	-1.03922824020165	9	891.41	EASGLSDNETEWLK	00000001000000
4393	138.06	138.05	3798.00	-0.00	-0.420122390602973	9	891.41	EASGLSDNETEWLK	00000001000000
4393	168.07	168.07	1526.00	-0.00	-0.642606113437682	9	891.41	EASGLSDNETEWLK	00000001000000
4393	186.08	186.08	1014.00	-0.00	-0.983467730223809	9	891.41	EASGLSDNETEWLK	00000001000000
4393	204.09	204.09	5041.00	-0.00	-1.06817259804309	9	891.41	EASGLSDNETEWLK	00000001000000
2739	126.06	126.05	7327.00	-0.00	-0.690174721011021	10	665.59	NA	NA
2739	138.05	138.05	1963.00	-0.00	-0.311470082107949	10	665.59	NA	NA
2739	144.07	144.07	468.60	-0.00	-0.5344787486255	10	665.59	NA	NA
2739	168.07	168.07	624.30	-0.00	-0.642606113437682	10	665.59	NA	NA
2739	204.09	204.09	2496.00	-0.00	-0.622284313992652	10	665.59	NA	NA

Table 1: Result

```
R> summary(S)
```

scans	mZ	markerIonMZ	markerIonIntensity
2540:6	Min. :126.1	Min. :126.1	Min. : 188.5
2739:5	1st Qu.:138.1	1st Qu.:138.1	1st Qu.: 624.3
3687:5	Median :144.1	Median :144.1	Median : 1526.0
4393:5	Mean :159.5	Mean :159.5	Mean : 2838.1
	3rd Qu.:186.1	3rd Qu.:186.1	3rd Qu.: 3273.0
	Max. :204.1	Max. :204.1	Max. :13620.0
markerIonMzError	markerIonPpmError	query	
Min. :-0.0014320	-0.642606113437682:	2 10:5	
1st Qu.:-0.0003100	-0.118001850879316:	1 4 :5	
Median :-0.0001310	-0.311470082107949:	1 6 :6	
Mean :-0.0002436	-0.420122390602973:	1 9 :5	
3rd Qu.:-0.0000870	-0.5344787486255 :	1	
Max. :-0.0000170	-0.622284313992652:	1 (Other) :14	
pepmass	peptideSequence	modification	
Min. :490.6	EASGLSDNETEWLK:5	0000000010000000:5	
1st Qu.:490.6	HSFNGNQSTFK :6	0000001000000 :6	
Median :665.6	IMNVTTDSLTK :5	0001000000000 :5	

```
Mean      :680.7    NA          :5        NA          :5
3rd Qu.:713.4
Max.     :891.4
```

**Some overview graphics** just an overview of the sample data set HexNAc.

```
R> op <- par(mfrow = c(2, 2), mar=c(4, 4, 4, 1))
R> dump <- lapply(split(S, S$query),
+   function(x){
+     plot(x$mZ, x$markerIonIntensity,
+       type = 'h',
+       col = 'lightblue',
+       cex = 2,
+       ylab = 'intensity', xlab='m/z',
+       xlim = range(c(HexNAc_MarkerIons,
+                     max(HexNAc_MarkerIons)
+                     + 0.1 * (max(HexNAc_MarkerIons) - min(HexNAc_MarkerIons)),
+                     min(HexNAc_MarkerIons)
+                     - 0.1 * (max(HexNAc_MarkerIons) - min(HexNAc_MarkerIons)))),
+       ylim = range(S$markerIonIntensity),
+       log = 'y',
+       main = paste("scan=", unique(x$scans),
+                   "/query=", unique(x$query), sep=''));
+     text(x$mZ, x$markerIonIntensity,
+       round(x$mZ,2), col='red', cex=0.7)
+   }
+ )
R> par(op)
```

Figure 1 displays the output of PTM\_MarkerFinder.

## 2.2. Reshaping the output and export

The R method `reshape` transforms the data frame `S` from a long format to a wide format.

```
R> names(S)[4] <- "mII"
R> S.wide <- reshape(S[,c(1,7,3,4)],
+   direction = 'wide',
+   timevar = "markerIonMZ",
+   idvar = c('scans', 'query'))
R>
```

export as comma separated file

```
R> write.table(S.wide,
+               file = "HexNAc_PTMarkerFinder.csv",
```

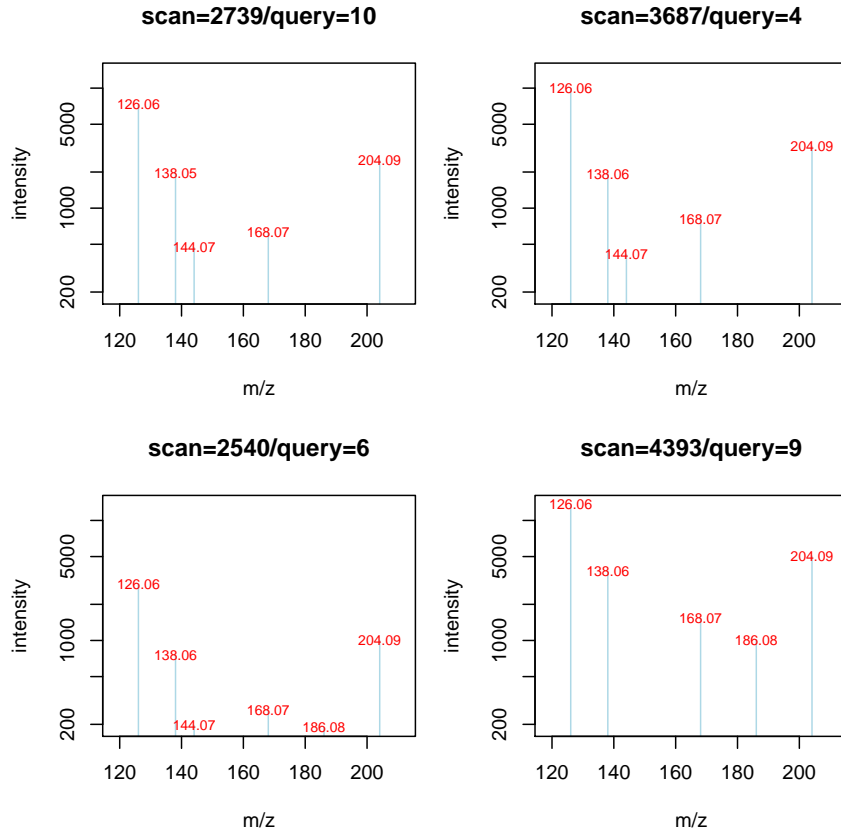


Figure 1: Overview of the marker ions.

scans	query	mII.126.05495	mII.138.05495	mII.144.06552	mII.168.06552	mII.204.08665	mII.186.07608
3687	4	9945.00	1933.00	412.30	810.20	3273.00	
2540	6	2945.00	759.20	195.40	262.90	998.40	188.50
4393	9	13620.00	3798.00		1526.00	5041.00	1014.00
2739	10	7327.00	1963.00	468.60	624.30	2496.00	

Table 2: Result

```
+      sep = ',',
+      row.names = FALSE,
+      col.names = TRUE,
+      quote = FALSE)
```

### 2.3. Visualization of the Result

```
R> # prepare the input
R> d <- list(); d[[1]] <- HexNAc[[3]]; d[[2]] <- HexNAc[[4]]; d[[3]] <- HexNAc[[5]]
R> S <- PTM_MarkerFinder(data = d, modification = m$mono,
+                         modificationName = m$desc,
+                         minMarkerIntensityRatio = 3,
```

```
+      itol_ppm = 20,
+      mZmarkerIons = HexNAc_MarkerIons)
```

The graphics can be seen in Figure 2.

### 3. Demonstration

The user can call the demonstration with

```
R> demo(PTM_MarkerFinder)
```

#### 3.1. Other examples

The following ADP-Ribose marker ions configuration was described by Bilan, Leutert, Nanni, Panse, and Hottiger (2017).

```
R> ADP_Ribose <- c(136.0618, 250.0935, 348.0704, 428.0367)
```

### 4. Session information

An overview of the package versions used to produce this document are shown below.

- R version 3.4.2 (2017-09-28), x86\_64-apple-darwin15.6.0
- Locale: en\_US.UTF-8/en\_US.UTF-8/en\_US.UTF-8/C/en\_US.UTF-8/en\_US.UTF-8
- Running under: macOS High Sierra 10.13.1
- Matrix products: default
- BLAS:  
`/Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRblas.0.dylib`
- LAPACK:  
`/Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRlapack.dylib`
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- Other packages: lattice 0.20-35, protViz 0.2.45, xtable 1.8-2
- Loaded via a namespace (and not attached): codetools 0.2-15, compiler 3.4.2, grid 3.4.2, Rcpp 0.12.12, tools 3.4.2

### References

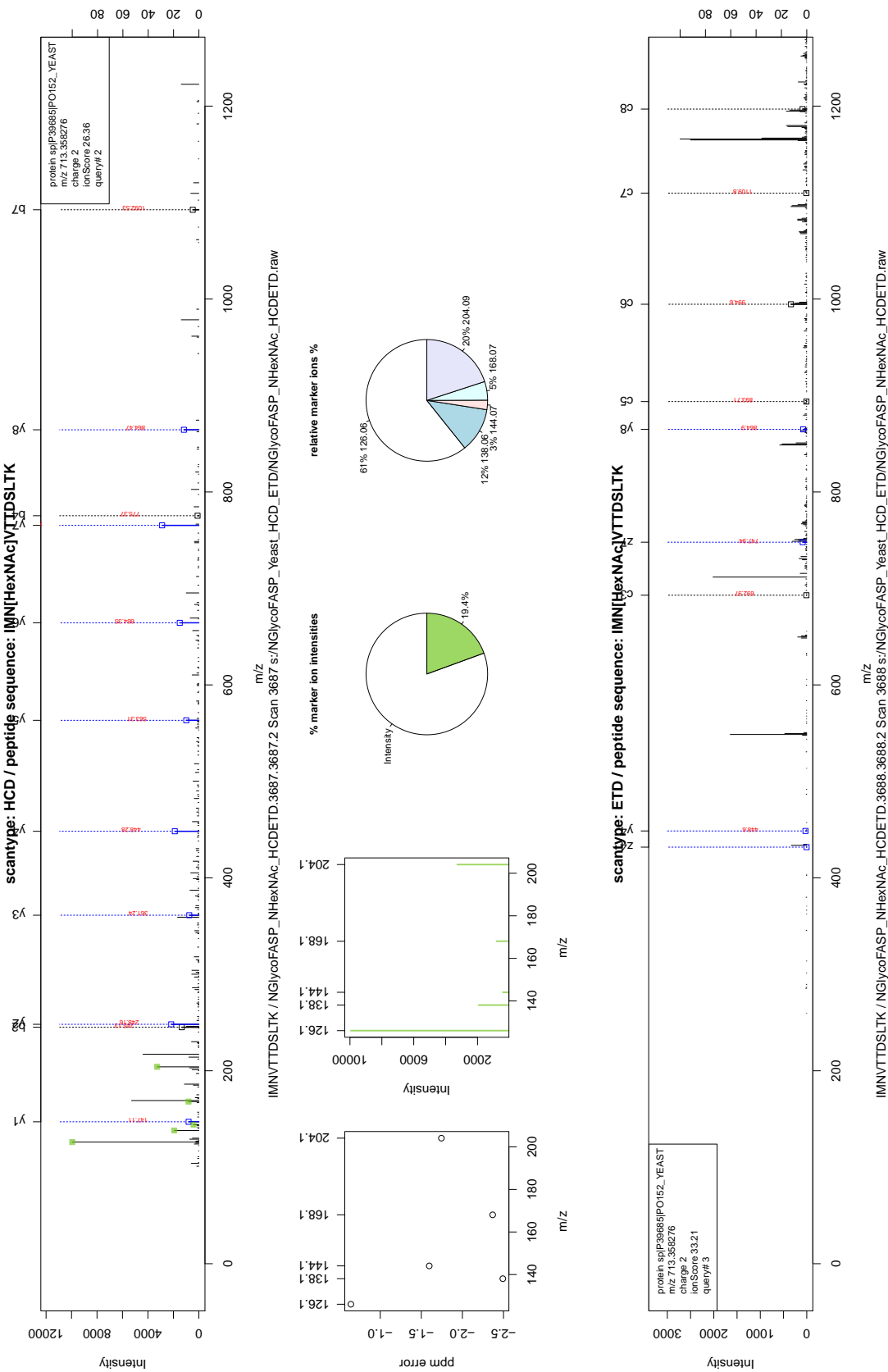


Figure 2: Graphical output of the method.

- Bilan V, Leutert M, Nanni P, Panse C, Hottiger MO (2017). “Combining Higher-Energy Collision Dissociation and Electron-Transfer/Higher-Energy Collision Dissociation Fragmentation in a Product-Dependent Manner Confidently Assigns Proteomewide ADP-Ribose Acceptor Sites.” *Anal. Chem.*, **89**(3), 1523–1530. doi:[10.1021/acs.analchem.6b03365](https://doi.org/10.1021/acs.analchem.6b03365).
- Nanni P, Panse C, Gehrig P, Mueller S, Grossmann J, Schlapbach R (2013). “PTM MarkerFinder, a software tool to detect and validate spectra from peptides carrying post-translational modifications.” *Proteomics*, **13**(15), 2251–2255. doi:[10.1002/pmic.201300036](https://doi.org/10.1002/pmic.201300036).
- R Development Core Team (2008). *R: A Language and Environment for Statistical Computing*. R Foundation for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL <http://www.R-project.org>.

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