

# ProtView: A versatile tool for *in silico* protease evaluation and selection in a proteomic and proteogenomic context

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## ABSTRACT

Tools have been created to generate *in silico* proteome digests with different protease enzymes and provide useful information for selecting optimal digest schemes for specific needs. This can save on time and resources and generate insights on the observable proteome. However, there remains a need for a tool that evaluates digest schemes beyond protein and amino acid coverages in the proteomic domain. Here, we present ProtView, a versatile in-silico

protease/protease combination and digest evaluation workflow that maps *in silico* digested peptides to both protein and genome references, so that the potential observable sections of the proteome, transcriptome and genome can be identified. This supports the identification and quantification of the proteomic evidence of transcriptional, co-transcriptional, post-transcriptional and translational regulations. Benchmarking against biological data comparing multiple proteases shows that ProtView can correctly estimate the relative performances among the digest schemes. ProtView provides this information in a way that is easy to interpret, allowing for digest schemes to be evaluated before carrying out an analysis, in a broader context to optimize proteomic and proteogenomic experiments. ProtView is available at <https://github.com/SSPuliasis/ProtView>.

## INTRODUCTION

Bottom-up proteomics involves using proteases to digest protein mixtures into peptides, which are then analysed by mass spectrometry (MS), allowing the peptides, and therefore the originating protein, to be identified (Aebersold and Mann, 2003). Shotgun proteomics refers to the use of bottom-up methods to identify proteins in complex mixtures. The shotgun proteomics workflow typically begins with the protein sample being denatured, reduced, alkylated, and digested by one or more proteases into peptides, which are then separated by liquid chromatography and identified by tandem mass spectrometry (MS/MS) and database searching (Wu et al., 2002). Database searching determines whether a peptide sequence in a database gives a significant match to each MS/MS spectrum and the degree of matching is assigned a score (Cottrell, 2011). It is paramount that every step is carried out effectively to maximise peptide identification and quantification, eventually maximising protein coverage and quantification. When proteins are digested in the first stage, it is the resulting peptides

that are carried into the subsequent analysis. Therefore, a peptide not generated in the digest cannot be identified in the subsequent analysis.

Trypsin is usually the protease of choice because it is highly specific, cleaving C-terminal to Lysine and Arginine residues (Keil et al., 1992), stable under a wide range of experimental conditions, and generates peptides in the preferred low charge and 7-35 amino acid length range for detection by MS machinery (Swaney et al., 2010), although it is not always the most suitable choice. For example, lysine and arginine are less frequent in membrane spanning protein regions (Kyte and Doolittle, 1982), resulting in fewer detectable peptides per unit length, thus limiting identification and quantification using MS methods when membrane spanning regions are digested with trypsin. Lysine and arginine are also enriched at exon-terminating and junction residues due to their codons (Wang et al., 2018), resulting in trypsin cleavage at splice junctions. This impedes the detection of junction-spanning peptides necessary to identify splice isoforms arising from alternative mRNA splicing. Furthermore, due to enzyme specificity, the perpetual use of any highly specific protease will continuously generate the same sub-sets of peptides. This eventually leads to a 'tunnel vision' display of the proteome in databases and repositories, with regions or even whole proteins that do not produce MS/MS suitable peptides with the protease in question being unidentified and remaining uncharacterised (Tsiatsiani and Heck, 2015).

Digests with different proteases, either to replace or to complement trypsin, have emerged as a way of mitigating the above issues and have proven to be useful in the study of membrane proteins (Fischer and Poetsch, 2006), splice junctions (Wang et al., 2018), N-termini not accessible by trypsin (Soh et al., 2020), and the study of post-translational modifications (PTMs) (Tran et al., 2016). Explorations into alternative enzymes support the argument that there

could be an ideal, non-trypsin centric, digestion scheme for every biological question and type of analysis (Tsiatsiani and Heck, 2015).

Expanding upon this idea, multiple protease digestion strategies can also be brought to bear on the issue of uncharacterised proteins. Combining multiple enzymes can be done in parallel, where peptide information from single protease digests is combined during post MS/MS analysis, or concurrently, where multiple proteases are added to the same sample *in vitro* before MS/MS analysis is carried out. It has been reported that using enzymes in parallel results in a significant increase in sequence coverage compared to single digests of the *Saccharomyces cerevisiae* proteome (Swaney et al, 2010) *Cannabis sativa* buds (Vincent et al., 2019a, Vincent et al., 2019b), human cervical cancer cells (Guo et al., 2014), and human recombinant protein (Choudhary et al, 2003). On the other hand, concurrent digests were reported to increase the number of identified proteins when Trypsin-Asp-N was used on *Schizosaccharomyces pombe* whole cell lysates when compared to trypsin alone (Dau et al., 2020), and be more efficient at yielding fully cleaved peptides and reducing the abundance of missed cleavage in peptides when Trypsin-Lys-C were used on *S. cerevisiae* (Glatter et al., 2012).

The aforementioned studies can make digest scheme recommendations for specific species and types of experiment after carrying out comparisons between digest schemes *in-vitro*. The scope of proteomic analyses is very broad and knowing which digest scheme is better suited to an analysis beforehand can save on costs, time and resources. Programs such as PeptideCutter (Gasteiger et al., 2003) and Rapid Peptides Generator (RPG) (Maillet, 2019) can digest protein sequences with different enzymes *in-silico* to give peptides that will theoretically be generated by a digest. ProteaseGuru (Miller et al., 2021) and Proteogest (Cagney et al., 2003) go a step further and provide interpretations of their digest results to aid in protease selection. Proteogest is a Perl application that allows the user to select a combination of

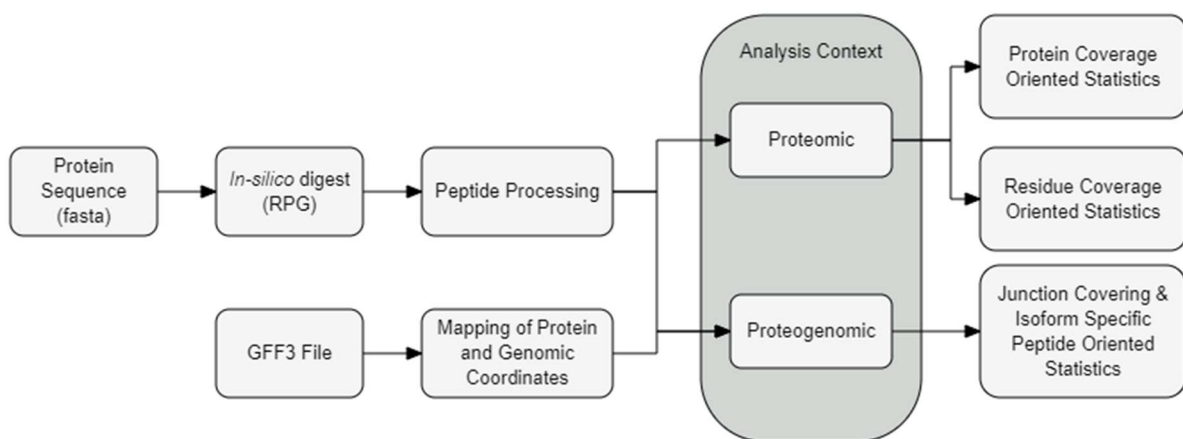
provided or custom modifications, to assess the effects of PTMs on the outcome. ProteaseGuru is a versatile and accessible tool that provides detailed peptide information and includes database PTM annotations and data visualization in the outputs. Nonetheless, there remains a need for such a tool that can also provide information in a wider context, that includes transcriptomic and genomic coverages and regions, e.g. to aid the study of alternative splicing and identify peptides that are unique to individual transcript isoforms, thus allowing the identification and quantification of the effects of transcriptional and translational regulations.

This work introduces ProtView, a method that integrates *in-silico* digestions by Rapid Peptides Generator (RPG) and provides the set of possible peptides that can be identified by each protease, or protease combination, and the variable information that they present, such as peptide length distributions, protein sequence coverage, and amino acid coverage. It also maps the digested peptides back to the genome using coding sequence (CDS) information from the annotations, which can provide detailed locations and information of the digested proteome in transcriptomic and genomic context, enabling analyses such as the identification of splice junction covering peptides, and isoform-unique peptides. It is the first tool that allows for the use of *in silico* proteomic evidence to investigate transcriptional and translational regulations to study the proteomic impact of alternative splicing, polyadenylations, alternative transcriptional starting locations, as well as alternative translational site regulations. We have demonstrated the utility of ProtView, with an analysis on the *Arabidopsis thaliana* proteome. We also compared ProtView predictions with *in-vitro* protease experiments and published proteomic data.

## METHODS

ProtView is a novel computation tool that presents all the *in silico* digested peptides by each protease, or protease combination, and their detailed information on the proteome, transcriptome, and genome, such as peptide length distributions, protein sequence coverage, specific residue coverage, peptides that cover splice junctions, junction coverage, the number of isoform-unique peptides, and genomic coordinates of peptides (Figure 1). All programming was done in Python 3.8 under the GPL v3 license. Details of the program and instructions can be found at (<https://github.com/SSPuliasis/ProtView>).

## PROTVIEW WORKFLOW



**Figure 1.** Outline of the ProtView workflow

### Peptide Processing

Rapid Peptides Generator (RPG) (Maillet, 2019) is incorporated into ProtView because it can process the whole protein database in one go, allows for user-defined proteases, and can generate more information than PeptideCutter (Gasteiger *et al.*, 2005), such as isoelectric point of each peptide. RPG has the option to carry out single or concurrent digests, where a sequence is simultaneously cleaved by multiple enzymes. Based on the output of RPG, ProtView creates parallel enzyme digests by combining peptides from individual single digests.

Duplicates of peptides that are generated by more than one of the single digests (e.g., by proteases with similar cleavage specificities) are removed from the parallel digest, resulting in a non-redundant parallel digest output. The enzyme name in the parallel digest output is the enzymes used separated by '/', whereas the enzyme name for concurrent digests in RPG is the enzymes used separated by '-'. Peptides can then be filtered by amino acid length and/or by content of a specific residue. Filtering for a length of 7-35 amino acids is recommended as the detectable range, although this optional filtering can be set by the user to match the instrumentation or criteria for the experiment. Below shows an example of a peptide generated by RPG and processed by ProtView (Table 1).

**Table 1.** Example of a digested peptide after processing digest results. The table contains information on the enzyme used in the theoretical digest, relative peptide start and cleavage position (end) on the protein sequence, size in amino acids, molecular weight, isoelectric point, peptide sequence, and the protein that the peptide originates from

enzyme	Peptide start	Peptide End	Peptide size	Mol weight	Isoelectric point	Sequence	Parent
Trypsin	1	8	8	991.1297	7.77	MFSNIDHK	AT1G66600.1

### Proteomic Summary Statistics

Protein sequence coverage is the percentage of the original protein sequence digested that is covered by filtered peptides and is calculated as the ratio between the sum of filtered peptide lengths and the total lengths of protein sequences in the FASTA file. Residue coverage is the percentage of an amino acid in the original FASTA sequence file that is covered by the filtered peptides. The total number of peptides generated by each enzyme both before and

remaining after filtering are also presented in the table, alongside their mean and median lengths. Table 2 is a summary table generated by ProtView for a single gene as an example, expandable to multiple genes or a proteome.

Table 2. Summary statistics of a digested protein after processing digest results, with Threonine(T) coverage

enzyme	Total peptides	Mean length	Median length	Filtered peptides	Sequence coverage (%)	T coverage (%)
Asp-N	95	8.9	7	46	70.7	81.2
Glu-C	104	8.1	5	43	80.6	96.9

### **Coding Sequence (CDS) preparation for downstream analyses**

When the genomic coordinate for each protein is available in gene and transcript annotations as coding sequences (CDS) in gff3 format, ProtView assigns a unique ID to each CDS and the adjacent intron preceding it, calculates the length of each intron between CDS, and converts the genomic coordinates of CDS regions to the relative protein sequence coordinates. CDS and intron IDs consist of chromosome, start position, end position, and strand (e.g., Chr1\_24848737\_24848859\_+). While converting CDS genomic coordinates to their relative proteomic coordinates, intron lengths between CDS are added up separately for each isoform, to give cumulative intron length of the introns preceding each CDS within an isoform. To obtain the relative protein coordinates for a CDS on the positive strand, the translation start position of the gene and cumulative intron length (after translation start position) are subtracted from the CDS genomic coordinate increased by one, then divided by 3 to take the triplet code into account. The final number is rounded up to the nearest integer



as shown in the equation below. While for CDS on the negative strand, the same equation can be applied with adjustments shown below.

$$\begin{aligned} & \text{relative protein coordinate (positive strand)} \\ & = \text{RoundUp}\left(\frac{(\text{coordinate} - \text{translation start position of gene} + \text{cumulative intron length} + 1)}{3}\right) \end{aligned}$$

$$\begin{aligned} & \text{relative protein coordinate (negative strand)} \\ & = \text{RoundUp}\left(\frac{(\text{coordinate} - \text{translation start position of gene} + \text{cumulative intron length} + 1)}{3}\right) \end{aligned}$$

Output from this step is saved as two csv files, one for each DNA strand, and allows for the downstream conversion of relative proteomic peptide coordinates to genomic and the identification of peptides covering splice junctions. Table 3 shows an example of the output table for a protein on the positive strand.

**Table 3.** Format of CDSs after processing and extraction from GFF3 files

type	start	end	strand	Parent	CDS ID	Intron ID	Intron length	Cumulative intron length	Protein start	Protein end
CDS	24848737	24848859	+	AT1G66600.1	Chr1_24848737_24848859_+	Chr1_24848651_24848736_+	86	86	83	123

### Genomic coordinates of the peptides

Relative peptide coordinates from the digest output are converted to the outer bounds of their corresponding coordinates on the genome, depending on DNA strand (Figure 2). The digested peptides are compared to CDS on the protein coordinates first, and the information

from CDS that overlap with the digested peptide is used to calculate the genomic coordinates of the peptide. The conversion is shown as the equations below:

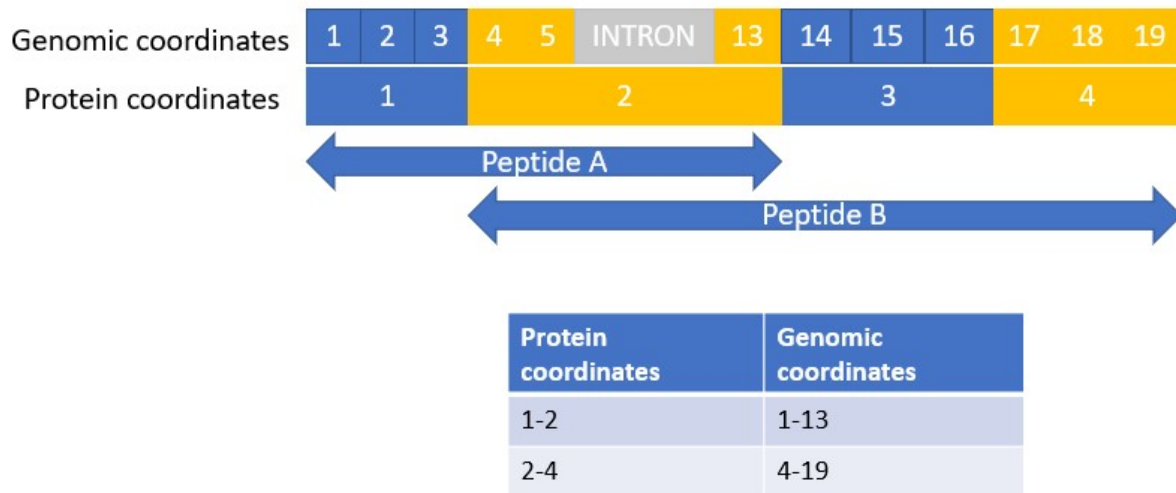
$$\text{genomic start position (positive strand)} = \text{translation start coordinate} + \text{cumulative intron length} + 3 * (\text{protein coordinate} - 1)$$

$$\text{genomic end position (positive strand)} = \text{translation start coordinate} + \text{cumulative intron length} + 3 * (\text{protein coordinate} - 1) + 2$$

$$\text{genomic start position (negative strand)} = \text{translation start coordinate} - \text{cumulative intron length} - 3 * (\text{protein coordinate} - 1) - 2$$

$$\text{genomic end position (negative strand)} = \text{translation start coordinate} - \text{cumulative intron length} - 3 * (\text{protein coordinate} - 1)$$

In the example figure below (Figure 2), amino acids are colored in blue and yellow, and introns in grey. Nucleotide coordinates are numbered within amino acids, with three nucleotides corresponding to one amino acid. Peptides A and B each have the second amino acid as a coordinate, however the number given for the genomic coordinates differs between the peptides due to this amino acid representing the start of peptide B and the end of peptide A, and the script giving outer bounds of genomic positions.

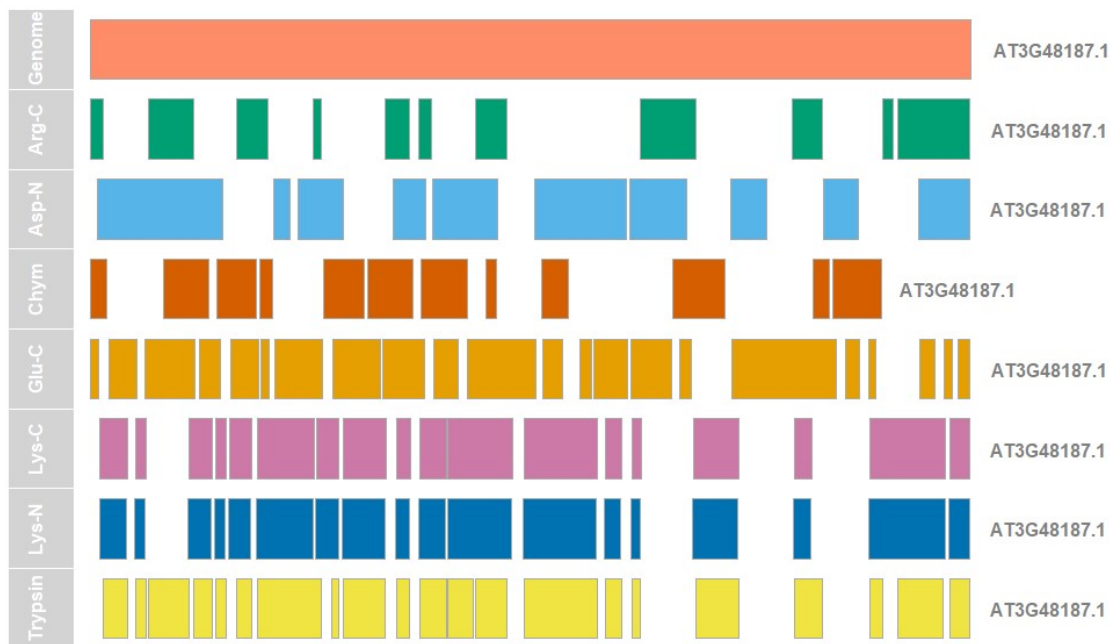


**Figure 2.** Illustration example of relative peptide coordinates and their corresponding coordinates on the genome. The top row of numbers represents nucleotides in the genome, with each triplet corresponding to one amino acid in the protein sequence. The row underneath represents the relative protein coordinates in amino acids (alternating blue and yellow).

The resulting data frame contains the parent isoform, both genomic and protein coordinates for each peptide, and the enzymes used to generate the peptide. Table 4 shows an example of two peptides (SPPVYRTTYLGQHTCKAFGVHD, DNTYGSEMINF), generated by digestion of AT1G66600.1 with Glu-C and Asp-N respectively, and their calculated genomic coordinates. Note that the peptides share the relative protein coordinate 169, but that their genomic coordinates differ after the conversion due to ProtView giving the outer bound of genomic coordinates, depending on whether a coordinate is at the start or the end of a peptide. Genome browser and visualization tools such as the R package Gviz (Hahne and Ivanek, 2016) can be used to visualize peptides mapped onto the genome, as shown in Figure 3.

**Table 4.** Output format of genomic coordinate conversion function of two exemplar digested peptides

Peptide	isoform	Protein start coordinate	Genomic coordinate start	Protein end coordinate	Genomic coordinate end	enzyme
SPPVYRTTYLGQHTCKAFGVHD	AT1G66600.1	148	24849051	169	24849116	Glu-C
DNTYGSEMINF	AT1G66600.1	169	24849114	179	24849146	Asp-N



**Figure 3.** Gviz visualization of genome coverage for the mono-exonic gene AT3G48187.1 using peptides in the 7-35 aa range. The top row represents the genomic sequence. Regions of peptide sequence coverage generated by different proteases are shown in the subsequent rows.

### Junction covering peptide identification & statistics

Another function of ProtView is to identify peptides that span splice junctions. Relative protein splice junction coordinates are determined as the average of the CDS protein coordinates on each side (i.e. 20.5 if a upstream CDS ends at 20 and the downstream begins at

21). For each transcript/proteome isoform, each peptide is then checked against the junction coordinates and is considered as junction-covering if it has at least one amino acid encoded on either side of a splice junction. Positive outcomes are saved in the same format as the digest results, with an additional column for junction location. A summary table is generated from the junction-covering peptides, which includes the number of junction-covering peptides generated by each enzyme, the number of total and unique junctions that a digest scheme covers (to avoid double counting of splice junctions shared between transcripts), and junction coverage percentage, which is the percentage of the total junctions in the transcript isoforms being examined that are covered by digested peptides. Table 5 shows an example junction summary table.

**Table 5.** Junction summary statistics of digested proteins AT1G66600 and AT1G66610

Enzyme	Junction spanning peptides	Unique junctions covered	Total junctions covered	Total junction coverage (%)
Asp-N	2	2	2	33.33333
Glu-C	6	5	6	100

### Isoform-unique peptides

Isoform-unique peptides can be used to identify a protein isoform with certainty and to discriminate between protein/transcript variants with different functions. The isoform-unique peptides come from exons that cover unique regions 1) due to alternative splicing events; 2) N or C terminus of the proteins resulting from different translation start or termination sites. The number of isoform-unique peptides is calculated by removing duplicate peptide sequences

that can be found in more than one isoform from the filtered peptides, giving the number of peptides generated that can only be found in one isoform in each specific digest.

### **Data acquisition & preparation**

The Araport11 *A. thaliana* proteome (Cheng et al., 2017) is used in this manuscript as a dataset to illustrate the range of information that can be given by ProtView. The Arabidopsis protein sequence databases were downloaded from the TAIR database ([https://www.arabidopsis.org/download/index-auto.jsp?dir=%2Fdownload\\_files%2FSequences%2FAraport11\\_blastsets](https://www.arabidopsis.org/download/index-auto.jsp?dir=%2Fdownload_files%2FSequences%2FAraport11_blastsets)), and corresponding GFF3 files from Araport11 (<https://www.araport.org/data/araport11>).

We have also used a number of additional published proteomics data that uses different enzymes for digestion to check the consistency with predictions with ProtView. The Human recombinant DNA derived tissue plasminogen activator protein sequence and *S. cerevisiae* proteome were downloaded from Uniprot (<https://www.uniprot.org/uniprot/P00750>, <https://www.uniprot.org/proteomes/UP000002311>) and digested using the same enzymes as the publications. All *in silico* digests were carried out using Rapid Peptides Generator (RPG) (Maillet, 2019) and filtered with ProtView for a recommended length of 7-35 amino acids.

## **RESULTS AND DISCUSSION**

### **Using ProtView to analyse digests in a proteomic context**

#### **i. Selecting enzymes with the highest protein coverage**

The purpose of the ProtView tool is to guide the choice of protease in *in vitro* experiments, by providing various statistics on peptides generated *in silico*. One of these measurements is the percentage of the original protein sequence that is covered by the digested and potentially identifiable peptides. While the coverage values given by ProtView are expected to be higher

than those obtained in the laboratory, due to the theoretical analysis giving the upper limit of all possible peptides that could be identified, ProtView is useful in being able to show how digest schemes perform relative to one another.

The *A. thaliana* proteome was digested *in silico* by Rapid Peptides Generator (RPG) with Arg-C, Asp-N, Chymotrypsin (high specificity), Glu-C, Lys-C, Lys-N, Trypsin, and pairwise combinations of these proteases in concurrent (represented in this article by '-') and parallel digests (represented in this article by '/'). The resulting peptide data will be used as an example to demonstrate the utility of ProtView. These proteases were chosen because they are highly specific, common alternative proteases, and often paired with trypsin in the literature. Due to the number of possible combinations, this example includes Trypsin in concurrent and parallel combination with Asp-N, Chymotrypsin, and Lys-C. Results for the remaining combinations can be found in the supplementary material. Summary statistics for these digests were calculated using ProtView and can be seen in Table 6, which includes the total number of peptides before and after filtering by length (7-35aa length used here), median and mean lengths of the peptides generated, the number of isoform-unique peptides, and sequence coverage %.

**Table 6.** Summary statistics of in silico digests carried out on *A. thaliana* sorted from highest to lowest protein sequence coverage %

Enzyme	Total peptides	Mean length	Median length	7-35aa range peptides	Sequence coverage (%)	Isoform-unique peptides
Trypsin/Chymotrypsin	4,053,121	10.3	7	1,907,193	88.7	545,769
Trypsin/Asp-N	3,928,784	10.6	7	1,859,247	88.2	525,351
Trypsin/Lys-C	3,017,079	13.8	7	1,442,574	78.6	408,117
Trypsin-Lys-C	2,455,307	8.5	6	1,042,209	69.6	290,657
Trypsin	2,377,488	8.8	6	1,028,635	69.5	287,199
Glu-C	2,599,479	8.0	5	1,049,420	67.8	287,602
Chymotrypsin	1,675,637	12.4	8	878,560	64.1	258,574
Asp-N	1,562,709	13.3	9	834,695	63.3	239,505
Trypsin-Asp-N	3,735,120	5.6	4	1,076,459	59.7	292,529
Trypsin-Chymotrypsin	4,015,741	5.2	4	1,078,818	59.2	291,116
Lys-N	1,380,762	15.1	9	719,633	56.0	207,805
Lys-C	1,377,666	15.1	9	719,763	56.0	207,580
Arg-C	1,173,294	17.8	11	620,534	50.4	182,158



It is not surprising that the total numbers of digested peptides are higher for protease combinations than they are for single proteases, due to combining multiple sets of peptides in the case of parallel digests (‘/’) or increased cleavage sites by using multiple proteases in concurrent digests (‘-’). Despite generating a high number of unfiltered peptides, the concurrent Trypsin-Asp-N and Trypsin-Chymotrypsin combinations give relatively low sequence coverage % in comparison to the other digests, due to many digested peptides being shorter than 7 amino acids and therefore below the filtering threshold. The concurrent Trypsin-Lys-C combination gives a slight increase in sequence coverage (0.4%) when compared to the single tryptic digest, likely due to Lys-C cleaving after Proline, whereas trypsin alone does not (Keil, 1992), and therefore increasing cleavage frequency. This combination is favoured *in vitro* because while both proteases cleave at lysine, Lys-C is more efficient at lysine cleavage than trypsin, and therefore combining them reduces the number of mis-cleaved peptides (Glatter et al., 2012). It should be noted that the order in which the enzymes are added to a sample concurrently doesn’t affect the results *in silico*, however this may not be the case *in vitro*.

## ii. Selecting enzymes with the highest coverage of specific residues

Amino acid composition differs across protein types and families, and the level of post-translational modification differs between amino acids. For example, lysine and arginine, the residues that trypsin cleaves at, are enriched 2.37-fold and 1.95-fold at exon-exon junctions (Wang et al., 2018), but are less frequent in membrane proteins (Kyte and Doolittle, 1982). Acetylation is a PTM that typically occurs on lysine residues. Being able to plan a digest around maximising the coverage of a specific residue may prove useful and options to filter for peptides containing a specific amino acid and calculating amino acid coverage are included in

ProtView, which is shown as the percentage of an amino acid in the original sequence that is covered by peptides after filtering for length.

Table 6 shows residue coverage of Cysteine (C), Serine (S), and Lysine (K) from *in-silico* *A. thaliana* digests to exemplify how much residue coverage % can differ between digest schemes. The digest scheme that gives the highest lysine coverage is Trypsin/Asp-N (80.6%), followed by Trypsin/Chymotrypsin (78.6%), with GluC giving the highest lysine coverage out of the single protease digests (72.8%). This suggests that any of these digests may be favourable to use in analyses focused on the study of acetylation, one of the PTMs that is associated with Lysine. Acetylated Lysine sites are typically not cleaved by trypsin (Garcia et al., 2007), thus peptide length distributions also need to be examined if considering Trypsin/Asp-N or Trypsin/Chymotrypsin. For all three of the residues examined here, parallel protease combinations give the highest coverages, with the exception of Trypsin/Lys-C giving lower lysine coverage (59.1%) due to both of these proteases cleaving at lysine. It should be noted that the protease that gives the highest residue coverage *in silico* may be otherwise unsuitable for use in a certain analysis or require adaptations to the experimental design *in vitro*. For example, in the context of ubiquitination, and despite Glu-C giving high lysine coverage, Glu-C digestion will result in peptides with a STLHLVLRIRGG ubiquitin remnant attached to Lys, causing a +1302.79 Da mass shift that needs to be taken into account in the database search (Warren et al., 2005).

**Table 6.** Residue coverage statistics for *A. thaliana* using Cysteine (C), Serine (S), and Lysine (K)

enzyme	total peptides	mean length	filtered peptides (7-35 aa)	protein sequence coverage (%)	C coverage	S coverage	K coverage
Arg-C	1,173,294	17.8	620,534	50.4	51.5	50.1	51.1
Asp-N	1,562,709	13.3	834,695	63.3	51.7	63.1	65.9
Chymotrypsin	1,675,637	12.4	878,560	64.1	68.6	63.4	63.4
Glu-C	2,599,479	8.0	1,049,420	67.8	74.6	71.7	72.8
Lys-C	1,377,666	15.1	719,763	56.0	58.1	54.9	52.3
Lys-N	1,380,762	15.1	719,633	56.0	58.1	59.9	52.2
Trypsin	2,377,488	8.8	1,028,635	69.5	75.1	72.7	44.6
Trypsin/Asp-N	3,928,784	10.6	1,844,116	87.5	87.7	88.5	80.6
Trypsin/Chymotrypsin	4,053,121	10.3	1,889,594	88.0	91.9	88.9	78.6
Trypsin/Lys-C	3,017,079	13.8	1,432,830	77.9	82.1	79.0	59.1
Trypsin-Asp-N	3,735,120	5.6	1,076,459	59.7	31.9	67.8	30.7
Trypsin-Chymotrypsin	4,015,741	5.2	1,078,818	59.2	66.0	67.6	28.7
Trypsin-Lys-C	2,455,307	8.5	1,042,209	69.6	75.3	73.2	42.3

### Using ProtView to analyse digest outcomes in a transcriptomic context

ProtView provides unique opportunities to examine transcriptomic regulations using proteomic evidence by mapping the digested peptides to the genome reference. ProtView identifies junction-covering peptides and provides junction summary information for each digest scheme. This information is shown in Table 7, consisting of the number of junction-covering peptides, number of junctions covered, unique junctions covered (to avoid double counting of junctions shared between isoforms), and the percentage of junctions that are covered by peptides from each digest after filtering. The function for counting isoform-unique

peptides mentioned previously can be appended to the junction summary table if being examined in a transcriptomic context. The number of isoform-unique peptides was calculated for both the entire sets of peptides generated and the junction-covering peptides.

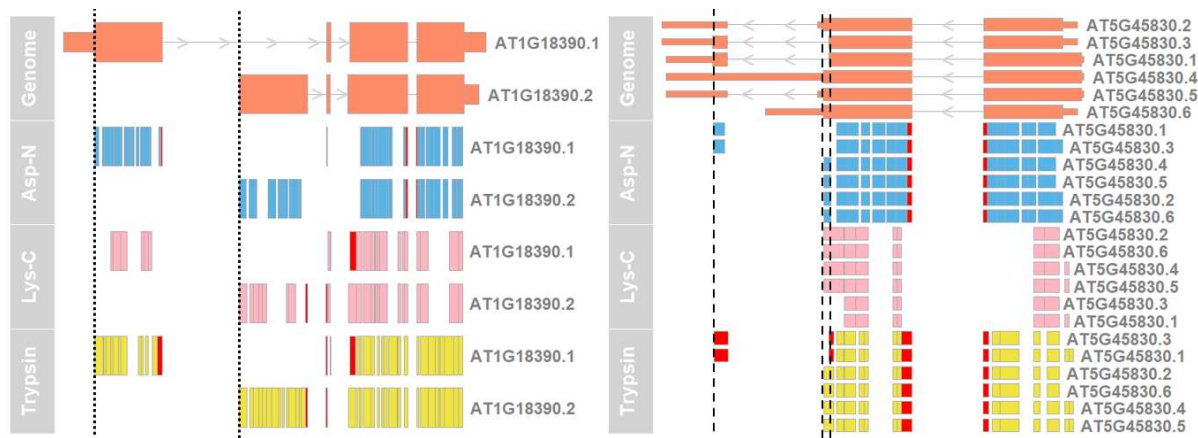
**Table 7.** Junction Summary statistics generated for *A. thaliana* using ProtView

Enzyme	Junction spanning peptides	Unique junctions covered	Total junctions covered	Total junction coverage (%)	Isoform-unique peptides	Junction covering isoform-unique peptides
Trypsin/Chymotrypsin	286442	101973	201015	84.5	545769	65589
Trypsin/Asp-N	276676	99323	195792	82.3	525351	63561
Trypsin/Lys-C	207385	80087	156990	66.0	408117	48464
Chymotrypsin	152914	77761	152914	64.2	258574	34697
Glu-C	147618	74684	147618	62.0	287602	33211
Asp-N	143447	72751	143447	60.3	239505	32777
Trypsin	133528	68070	133528	56.1	287199	30892
Trypsin-Lys-C	133484	68079	133484	56.1	290657	30917
Lys-N	129305	65925	129305	54.3	207805	29634
Trypsin-Asp-N	115813	58815	115813	48.7	292529	26358
Trypsin-Chymotrypsin	114895	58311	114895	48.3	291116	26014
Lys-C	111835	57631	111835	47.0	207580	26656
Arg-C	106708	54338	106708	44.8	182158	24673

Table 7 exemplifies the format of a junction summary table generated by ProtView, with the digest schemes sorted in order of highest to lowest junction coverage %. The output shows that in terms of single enzyme digests, Chymotrypsin, Glu-C, and Asp-N outperform trypsin in

terms of the number of junction-covering peptides generated and junctions covered for the Arabidopsis proteome, further underlining the point that trypsin may not always be the most optimal choice. If carrying out an analysis where maximising splice junction coverage is a priority, these example results suggest that combining chymotryptic peptides in parallel with tryptic peptides can theoretically give a 28.4% increase in junction coverage compared to trypsin alone, in addition to giving the most isoform-unique peptides that can be used to discriminate between protein isoforms.

In addition to the aforementioned overview statistics, ProtView can map peptides onto the genome, allowing the downstream examination and visualization of digested peptides to identify post-transcriptional regulations. For example, AT1G18390 is a gene with two transcript/protein isoforms with alternative transcription start sites. Visualisation of AT1G18390 *in-silico* generated peptides on the genome (Figure 4) shows coverage of the alternative transcriptional start sites (vertical dashed lines) by peptides and allows for isoform-specific and exon-exon junction covering peptides to easily be identified. In this example, trypsin and Asp-N generate peptides that cover the transcription start sites in both isoforms, while the peptides generated by Lys-C do not cover the transcription start site of the AT1G18390.1 isoform. Similarly, AT5G45830 is a gene on the negative strand with alternative stop sites that visually examined on the genome (Figure 5).



**Figure 4.** AT1G18390 and AT5G45830 in-silico generated peptides mapped onto the genome.

The first row shows the genome; thick coloured boxes represent exons, thin coloured boxes represent UTRs, grey lines linking between exons represent introns, and vertical lines show alternative start(AT1G18390) (dotted) or stop(AT5G45830) (dashed) sites. The subsequent rows represent peptides mapped onto the genome, with exon-exon junction-covering peptides in red.

#### Comparing ProtView in silico results with in vitro experimental data

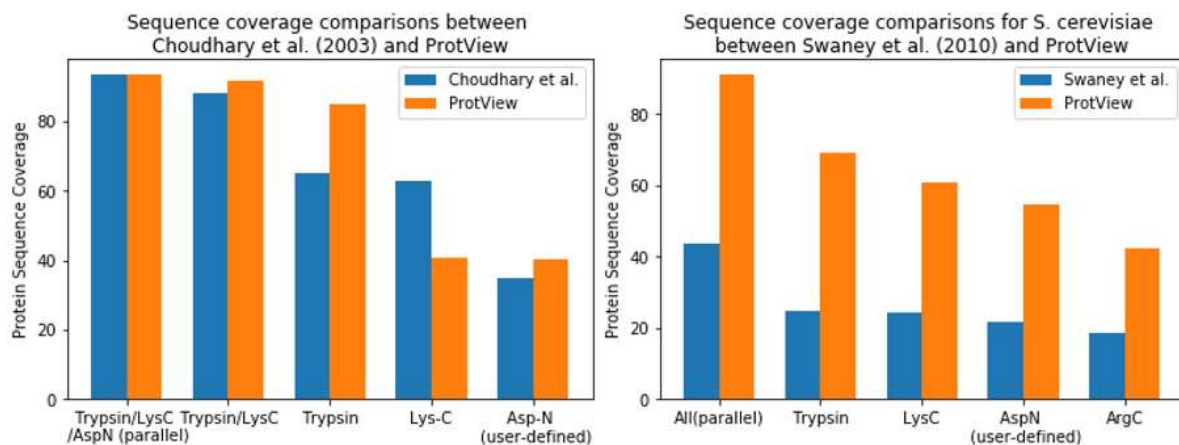
To evaluate how representative the order of protein sequence coverage given by ProtView for different digest schemes is of an *in vitro* experiment, comparisons were carried out between ProtView and two publications that compare protein sequence coverage using different proteases. Choudhary et al. (2003) examined a human recombinant tissue plasminogen activator protein using Trypsin, Lys-C, Asp-N and their parallel combinations. Swaney et al. (2010) examined *S. cerevisiae* digests with Trypsin, Arg-C, Asp-N, Lys-C, and all these proteases in parallel. The proteins and digest schemes used in the publications were digested with RPG and processed with ProtView to give protein sequence coverage

percentages for each digest scheme. Protein sequence coverage values were then compared between those obtained experimentally in the publications and the *in silico* generated values.

In the first comparison, the sequence coverage obtained by Choudhary et al. with Asp-N was much lower than predicted (34.9% for Choudhary et al., 67.1% for Protview), due to the RPG cleavage rules for Asp-N erroneously including cleavage at all Cysteines, despite Asp-N actually cleaving at Cystic acid, a vanishingly rare oxidation form of cysteine in natural protein samples (Paulech et al., 2015). It is highly unlikely therefore that Asp-N would deliver a meaningful number of peptides showing cleavage at "Cysteine". The regions covered by Choudhary et al. do not show cleavage at Cys, thus a user-defined Asp-N (denoted as Asp-N[~C]) cleaving only at Aspartic Acid, was used to repeat the comparison. The highest sequence coverage is obtained by using parallel enzyme digests, with Trypsin/Lys-C/Asp-N[~C] (93.3% for Choudhary et al., 93.4% for Protview), followed by Trypsin/Lys-C (88.2% and 91.8%), Trypsin (65% and 85%), Lys-C (62.8% and 40.6%), and Asp-N[~C] (34.9% and 40.2%). The rankings between the ProtView results and the experimental results are the same, with a spearman correlation of 1 (Figure 5). Two possible explanations were found for the experimental Lys-C coverage being higher than the predicted ProtView value: 1) Peptides above the 35aa filter cutoff length used by ProtView were identified experimentally due to being within the mass range used in the database search. 2) Non-specific cleavage of Lys-C, which can occasionally occur *in vitro* despite Lys-C being highly specific (Raijmakers et al., 2010), but is not considered in the *in-silico* predictions..

In the comparison to Swaney et al. (2010), using user-defined Asp-N rules (Asp-N[~C]), all enzymes in parallel achieved the highest sequence coverage (43.4% for Swaney et al., 91.0% for ProtView), followed by Trypsin (24.5%, 68.9%), Lys-C (24.3%, 60.8%), Asp-N[~C] (21.5%,

54.7%) and ending with Arg-C providing the lowest coverage (18.6%, 42.2%). As in the previous comparison, the spearman correlation value is 1. . The comparisons shows that ProtView can correctly predict the ranking of protease performance and provide rapid pre-analysis to assist in the choice of proteases for addressing a given experimental question.



**Figure 5.** sequence coverage comparisons for human recombinant tissue plasminogen activator between Choudhary et al. (2003) and ProtView and *S. cerevisiae* between Swaney et al. (2010) and ProtView.

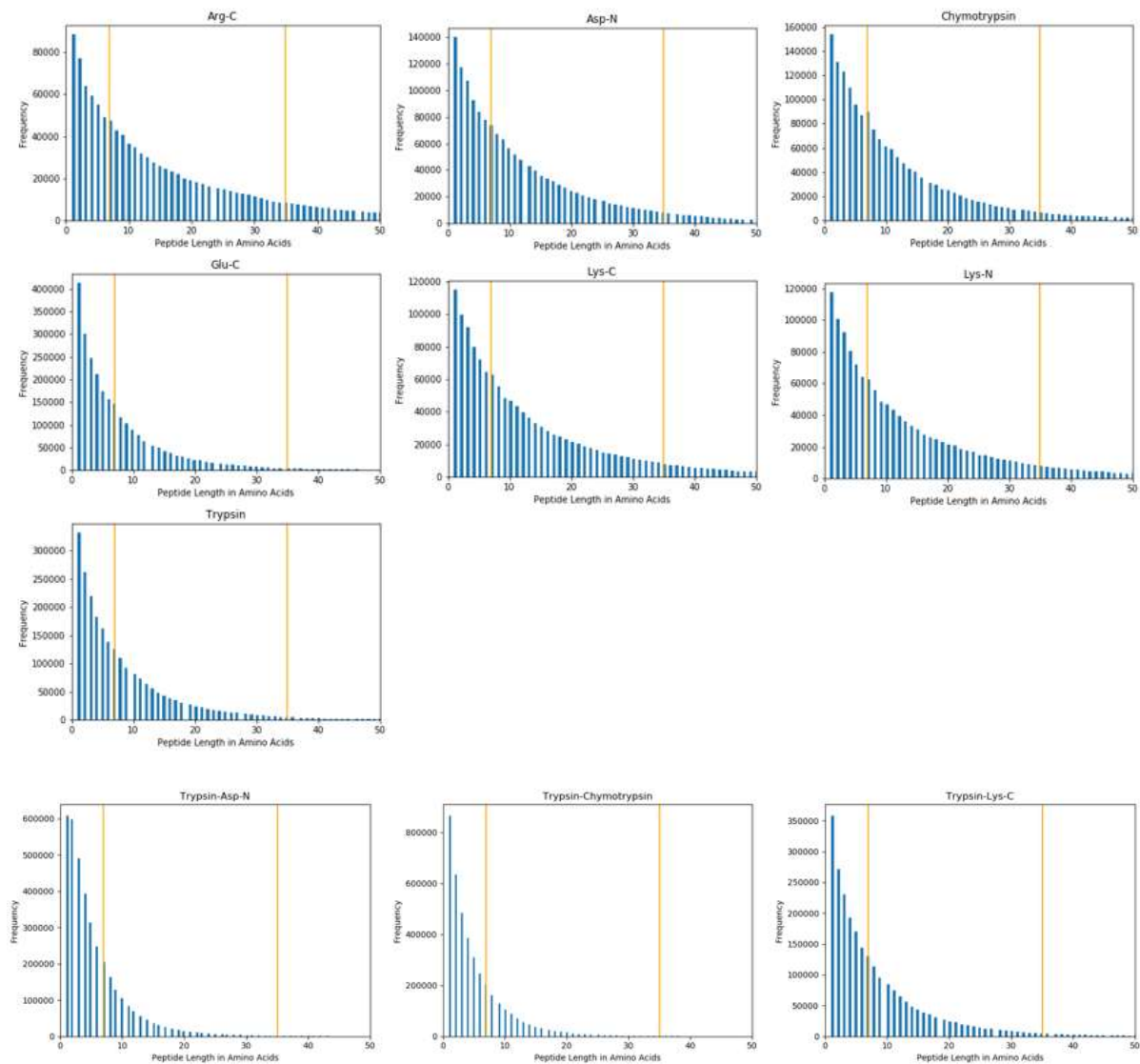
## CONCLUSIONS

Evaluation of digest schemes *in silico* can save on time and resources compared to *in vitro* evaluations. ProtView is a novel software tool for the evaluation of digest schemes, designed to process and analyse *in-silico* digest output by different enzymes and their combinations in multiple contexts. It is clear from our analysis that depending on the focus of the investigation, the ideal choice of enzyme could vary considerably. The enzyme combinations that provide the best protein sequence coverage do not necessarily provide the best view of the proteome in



terms of specific PTMs; Therefore, ProtView is timely to provide a tailored analysis that facilitates the decision and experimental planning process. Preliminary validations show that ProtView can reliably predict the majority of experimentally determined protein sequence coverage orders between digest schemes. ProtView is the first tool that maps the peptides to the reference genome, which allows transcriptomic activities, such as transcriptional (alternative transcriptional start sites and stop sites) and post-transcriptional regulations (alternative splicing), to be studied using proteomic experimental evidence. Mapping peptides to the genome also creates the possibility in the future to integrate sequence variations of different species and their sub-species (e.g cultivars/eco-types/landraces in plants), to derive a list of individualized peptides that is possible to be detected in mass spectrometry-based proteomics experiments.

## Supplementary Material



**Supplementary Figure 1.** Length distributions of *A. thaliana* peptides generated by single enzyme digests, showing lengths up to 50 amino acids. Orange vertical lines indicate the 7-35 amino acid length range used in these analyses and shown in Table 6.

**Supplementary Table 1.** Cleavage specificities of the enzymes used in RPG digests in the benchmark analyses in this article

Residue	R		C		D		F		Y		W		E		K		Exceptions
	C	N	C	N	C	N	C	N	C	N	C	N	C	N	C	N	
Enzyme																	
ArgC	X																
Asp-N				X		X											
Chymotrypsin (high specificity)							X		X		X						FP, YP, WP, WM
GluC					X								X				
Lys-C															X		
LysN																X	
Trypsin	X														X		KP, RP, CKD, DKD, CKH, CKY, CRK, RRH, RRR

**Supplementary Table 2.** Summary statistics of in silico digests carried out on *A. thaliana* for all proteases and pairwise protease combinations ('/' for parallel, '-' for concurrent)

Enzyme	Total peptides	Mean length	Median length	Filtered peptides (7-35 aa)	Protein sequence coverage (%)	Isoform unique peptides
Arg-C	1173294	17.8	11	620534	50.4	182158
Arg-C/Asp-N	2730374	15.3	10	1452634	81.4	420689
Arg-C/Chymotrypsin-high	2848912	14.6	9	1499091	81.6	440727
Arg-C/Glu-C	3772760	11.1	6	1669951	83.4	469753
Arg-C/Lys-C	2550939	16.4	10	1340291	77.8	389723
Arg-C/Lys-N	2547448	16.4	10	1337483	77.8	388979
Arg-C/Trypsin	3009921	13.9	7	1414582	78.7	401303
Arg-C-Asp-N	2606920	8.0	6	1108077	70.1	308649
Arg-C-Chymotrypsin-high	2800617	7.5	5	1104433	69.3	308969
Arg-C-Glu-C	3724458	5.6	4	1064510	60.6	291623
Arg-C-Lys-C	2502642	8.3	5	1050942	69.7	292816
Arg-C-Lys-N	2424059	8.6	6	1050942	69.7	294907
Arg-C-Trypsin	2435752	8.6	6	1054844	69.9	289667

Asp-N	156270 9	13.3	9	834695	63.3	239505
Asp-N/Chymotrypsin-high	322839 9	12.9	8	1709104	86.4	496638
Asp-N/Glu-C	413574 8	10.1	6	1882591	83.3	526570
Asp-N/Lys-C	293426 7	14.2	9	1551948	83.1	446174
Asp-N/Lys-N	294346 0	14.2	9	1554321	83.1	447300
Asp-N-Chymotrypsin-high	305755 2	6.8	5	1026380	63.2	308789
Asp-N-Glu-C	390315 7	5.3	3	1038222	69.6	284585
Asp-N-Lys-C	279981 7	7.4	5	1122250	67.4	304717
Asp-N-Lys-N	289515 3	7.2	5	1094902	68.0	303757
Chymotrypsin-high	167563 7	12.4	8	878560	64.1	258574
Chymotrypsin-high/Lys-C	305329 6	13.7	9	1598319	84.3	466150
Chymotrypsin-high/Lys-N	305006 9	13.7	9	1595540	84.3	465440
Chymotrypsin-high-Lys-C	300498 5	6.9	5	1120839	68.3	309886
Chymotrypsin-high-Lys-N	290946 5	7.2	5	1128967	68.8	311870
Glu-C	259947 9	8.0	5	1049420	67.8	287602
Glu-C/Chymotrypsin-high	427510 7	9.8	6	1927978	87.9	546174
Glu-C/Lys-C	397713 5	10.5	6	1769177	85.2	495169

Glu-C/Lys-N	396799 3	10.5	6	1765807	85.2	494340
Glu-C/Trypsin	497696 4	8.4	5	2078054	90.1	574792
Glu-C-Chymotrypsin-high	422680 1	4.9	3	1057850	56.7	286868
Glu-C-Lys-C	392882 7	5.3	3	1017603	58.0	277973
Glu-C-Lys-N	376001 2	5.5	4	1026685	58.5	280424
Glu-C-Trypsin	494353 7	4.2	3	911196	48.0	247326
Lys-C	137766 6	15.1	9	719763	56.0	207580
Lys-C/Lys-N	274457 4	15.2	10	1439365	57.7	415356
Lys-C-Lys-N	259856 8	8.0	1	665792	52.1	194045
Lys-N	138076 2	15.1	9	719633	56.0	207805
Lys-N/Trypsin	373589 3	11.2	7	1747316	79.5	494614
Lys-N-Trypsin	353495 1	5.9	2	976235	65.4	274177
Trypsin	237748 8	8.8	6	1028635	69.5	287199
Trypsin/Asp-N	392878 4	10.6	7	1844116	87.5	525351
Trypsin/Chymotrypsin-high	405312 1	10.3	7	1889594	88.0	545769
Trypsin/Lys-C	301707 9	13.8	7	1432830	78.0	408117
Trypsin-Asp-N	373512 0	5.6	4	1076459	59.7	292529

Trypsin- Chymotrypsin-high	401574 1	5.2	4	1078818	59.2	291116
Trypsin-Lys-C	245530 7	8.2	6	1042209	69.6	290657

**Supplementary Table 3.** Junction Summary statistics generated for *A. thaliana* using ProtView for all proteases and pairwise protease combinations ('/' for parallel, '-' for concurrent)

enzyme	Junction spanning peptides	Unique junctions covered	Total junctions covered	Total junction coverage (%)	isoform unique peptides
Arg-C	106708	54338	106708	44.8	182158
Arg-C/Asp-N	249948	93812	184954	77.7	420689
Arg-C/Chymotrypsin-high	259622	96985	191119	80.3	440727
Arg-C/Glu-C	254326	95721	189126	79.5	469753
Arg-C/Lys-C	218543	86269	169321	71.1	389723
Arg-C/Lys-N	235754	90064	177340	74.5	388979
Arg-C/Trypsin	211060	83177	163743	68.8	401303
Arg-C-Asp-N	154222	77981	154222	64.8	308649
Arg-C-Chymotrypsin-high	154947	78335	154947	65.1	308969
Arg-C-Glu-C	124541	63249	124541	52.3	291623
Arg-C-Lys-C	133636	68140	133636	56.2	292816
Arg-C-Lys-N	149667	75662	149667	62.9	294907
Arg-C-Trypsin	133794	68187	133794	56.2	289667
Asp-N	143447	72751	143447	60.3	239505
Asp-N/Chymotrypsin-high	295886	103107	203497	85.5	496638

Asp-N/Glu-C	290866	97271	192445	80.9	526570
Asp-N/Lys-C	255045	94945	186872	78.5	446174
Asp-N/Lys-N	272752	98233	193629	81.4	447300
Asp-N- Chymotrypsin-high	154709	77909	154709	65.0	308789
Asp-N-Glu-C	136795	69302	136795	57.5	284585
Asp-N-Lys-C	139246	70694	139246	58.5	304717
Asp-N-Lys-N	150661	76080	150661	63.3	303757
Chymotrypsin-high	152914	77761	152914	64.3	258574
Chymotrypsin- high/Lys-C	264749	98326	193618	81.4	466150
Chymotrypsin- high/Lys-N	281926	101082	199347	83.8	465440
Chymotrypsin-high- Lys-C	140548	71165	140548	59.1	309886
Chymotrypsin-high- Lys-N	153581	77369	153581	64.4	311870
Glu-C	147618	74684	147618	62.0	287602
Glu- C/Chymotrypsin- high	300532	104325	205863	86.5	546174
Glu-C/Lys-C	259453	97618	192543	80.9	495169
Glu-C/Lys-N	276607	99195	195916	82.3	494340
Glu-C/Trypsin	281146	103259	204067	85.7	574792
Glu-C- Chymotrypsin-high	120575	61040	120575	50.7	286868
Glu-C-Lys-C	109317	55888	109317	45.9	277973
Glu-C-Lys-N	119523	60743	119523	50.2	280424
Glu-C-Trypsin	84342	43214	84342	35.4	247326
Lys-C	111835	57631	111835	47.0	207580
Lys-C/Lys-N	241140	68533	134210	56.4	415356



Lys-C-Lys-N	106737	55000	106737	44.8	194045
Lys-N	129305	65925	129305	54.3	207805
Lys-N/Trypsin	262789	89875	177200	74.5	494614
Lys-N-Trypsin	127429	64907	127429	53.5	274177
Trypsin	133528	68070	133528	56.1	287199
Trypsin/Asp-N	276676	99323	195792	82.3	525351
Trypsin/Chymotrypsin-high	286442	101973	201015	84.5	545769
Trypsin/Lys-C	207385	80087	156990	66.0	408117
Trypsin-Asp-N	115813	58815	115813	48.7	292529
Trypsin-Chymotrypsin-high	114895	58311	114895	48.3	291116
Trypsin-Lys-C	133484	68079	133484	56.1	290657

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Sophia Puliassis: Methodology, Software, Writing – original draft, Formal analysis, Validation, Visualisation, Conceptualization. Runxuan Zhang: Conceptualization, Methodology, Supervision, Writing – original draft /Writing – review & editing, Project administration, Validation, Resources, Funding Acquisition. Piers Hemsley: Conceptualization, Supervision, Writing – review & editing, Resources, Funding Acquisition. Dominika Lewandowska: Conceptualization, Supervision, Writing – review & editing.

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