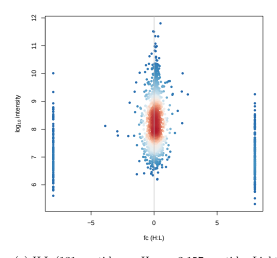
## Triple TAILS Report example

This report is based on 9,294 quantified peptide features provided in the mzTab input file. Some peptide sequences have been quantified multiple times (multiple charge states, different fractions). In the following analysis, we will choose the most intense peptide quantifications and focus on the subset of 3,858 (41.51%) unique peptide sequences. 1,378 of those peptides are mapped very close to N- or C-terminus (less than 6 amino acid positions) which leaves us with 2,480 quantified peptides.

	number of peptides with	number of peptides with	number of peptides with	total number
	$\mathrm{fc}  o -\infty$	fc unchanged	$\mathrm{fc} \to +\infty$	of peptides
H:L	131	2,157	192	2,480

Table 1: Number of peptides for which both or none of the two labelled forms were detected (fc unchanged), or only one of them (fc  $\to \pm \infty$ ).



(a) H:L (131 peptides no Heavy, 2,157 peptides Light and Heavy, 192 peptides no Light)

Figure 1: Fold change vs log intensity for the set of 2,480 unique peptides. Infinite fold changes (fc  $\to \pm \infty$ ) are mapped to fc =  $\pm 8$ . The colour gradient reflects the density of the scatter plot.

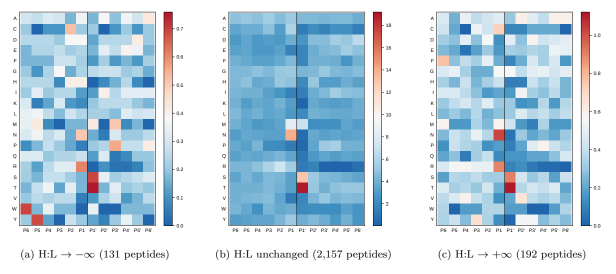


Figure 2: Amino acid frequencies fingerprints for the nine subsets listed in Table 1. For example, figure (g) depicts the fingerprint of peptides detected in light (L) but absent in medium (M).