Report for PEP Section in mzTab File example_4

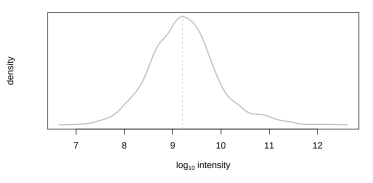
The PEP section of the mzTab file contains 1,335 quantified peptide features.

Peptides of Interest

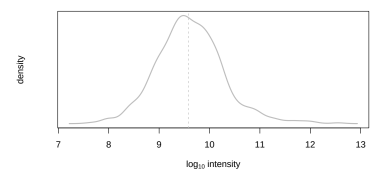
Sequence	Accession	Charge	Retention Time	m/z
SSAAPPPPPR	STD_01	2	1659.92	493.77
HVLTSIGEK	STD_03	2	2127.71	496.29
IGDYAGIK	STD_05	2	3096.71	422.74
TASEFDSAIAQDK	$STD_{-}06$	2	4266.53	695.83
SAAGAFGPELSR	STD_07	2	4457.27	586.80
ELGQSGVDTYLQTK	STD_08	2	5741.14	773.90
GLILVGGYGTR	$STD_{-}09$	2	6431.53	558.33
GILFVGSGVSGGEEGAR	P52209	2	6780.92	796.41
GILFVGSGVSGGEEGAR	P52209	2	6781.34	801.41
SFANQPLEVVYSK	$STD_{-}11$	2	6787.30	745.39
ELASGLSFPVGFK	$STD_{-}14$	2	9083.08	680.37

Proteins of Interest

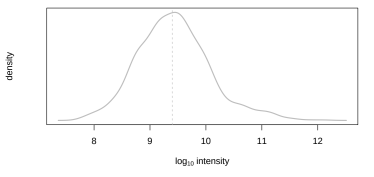
Sequence	Accession	Charge	Retention Time	m/z
INQEELASGTPPARFPK	O15117	3	4675.29	618.99



(a) peptide abundances 1, median (intensity) = 1,605,469,952



(b) peptide abundances 2, median (intensity) = 3,819,539,968



(c) peptide abundances 3, median (intensity) = 2,497,959,936

Figure 1: peptide abundance distributions. $\,$

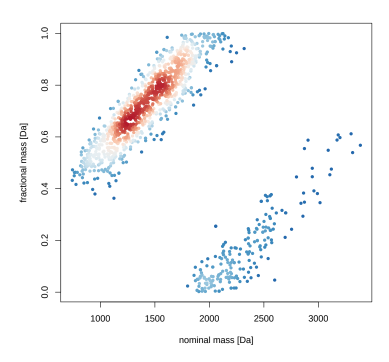


Figure 2: Kendrick nominal fractional mass plot

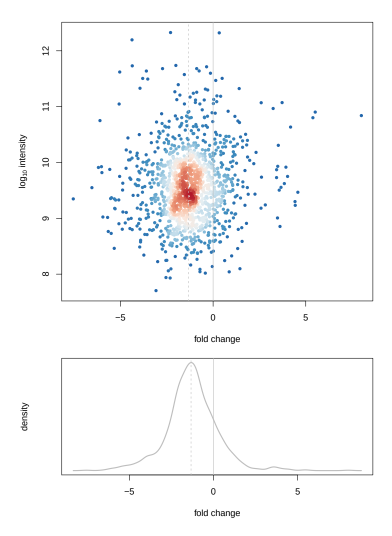


Figure 3: Fold changes of peptide abundances 1 and 2. $\mathrm{median}(\mathrm{fc}) = -1.3328 \qquad \mathrm{sd}(\mathrm{fc}) = 1.5445$

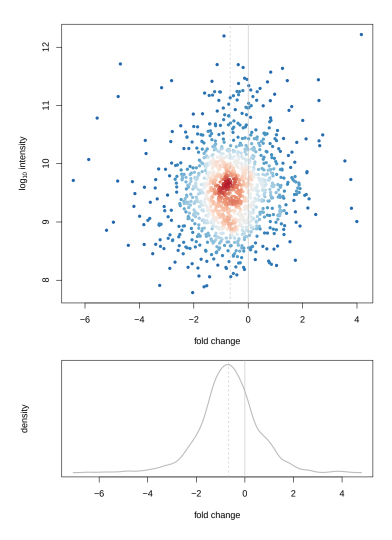


Figure 4: Fold changes of peptide abundances 1 and 3. $median(fc) = -0.6641 \qquad sd(fc) = 1.1804$

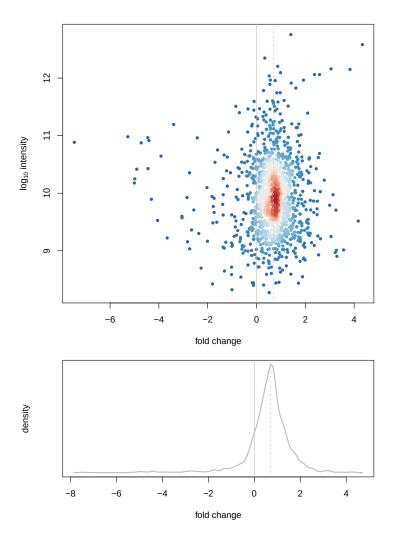


Figure 5: Fold changes of peptide abundances 2 and 3. $median(fc) = 0.6958 \qquad sd(fc) = 0.9636$

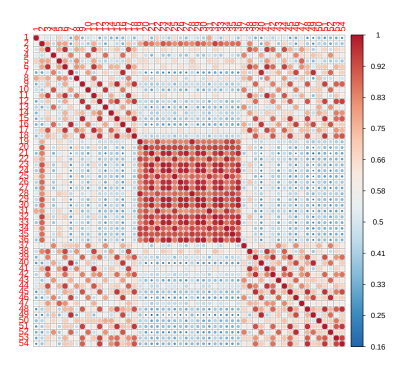


Figure 6: Correlation of all peptide abundances.