Analysing MaxQuant Output with R

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Contents

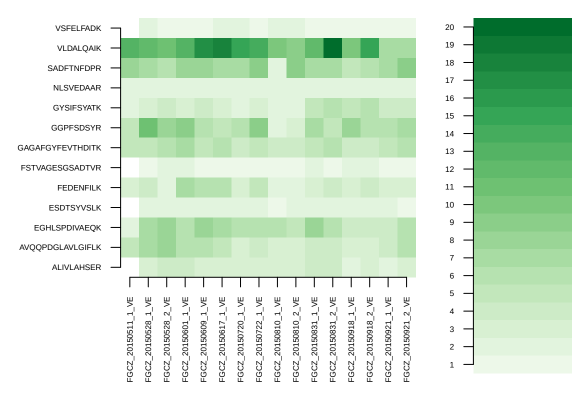
Prepare mrm table	1
Dump maxquant txt files into sqlite database	2
Peptide Evidence	2
Prepare mrm table	

- nr of assigned spectra (MS2) / versus total spectra
- How many H/L target peptides are identified
- Ratio among H/L of the target peptides
- Intensity of heavy or light over the runs

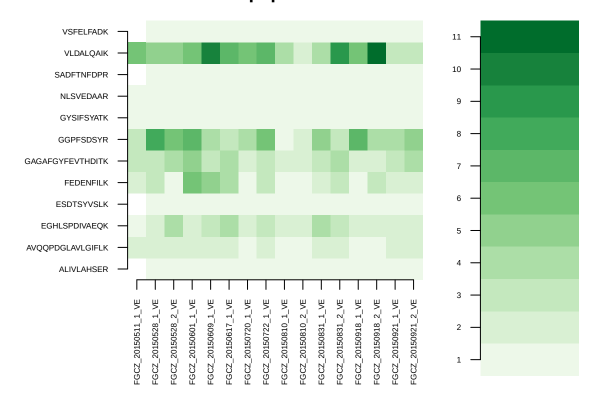
Dump maxquant txt files into sqlite database

Peptide Evidence

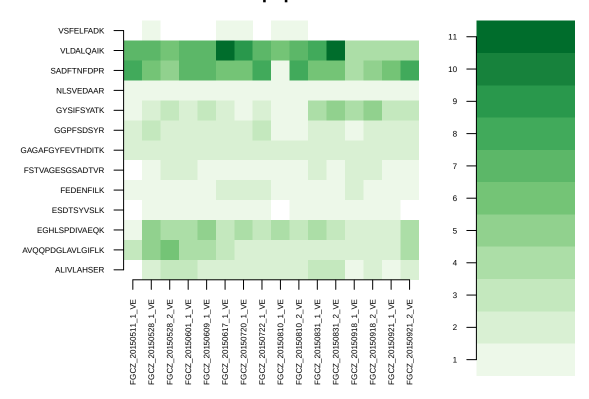
all peptide count

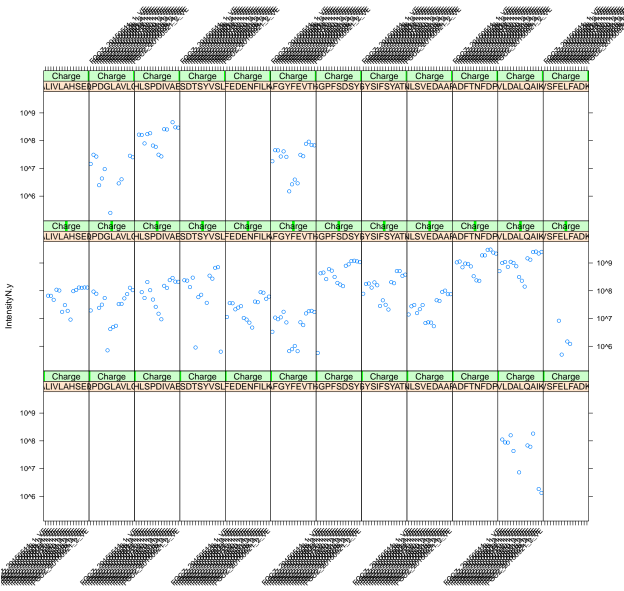


labeled peptide count

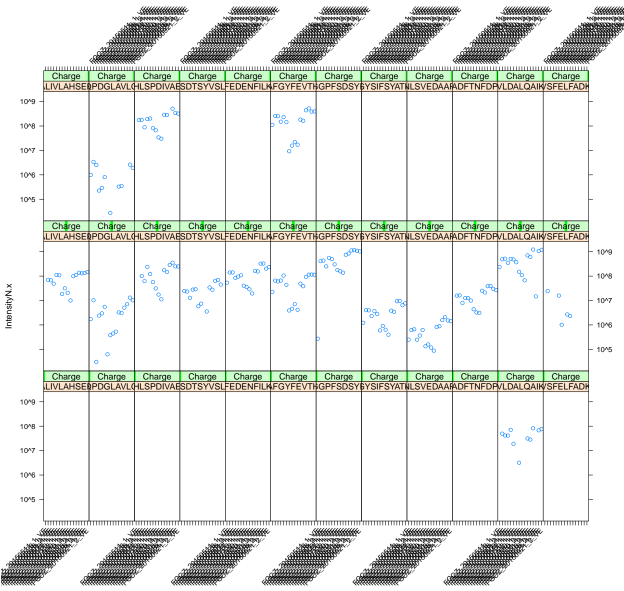


not labeled peptide count

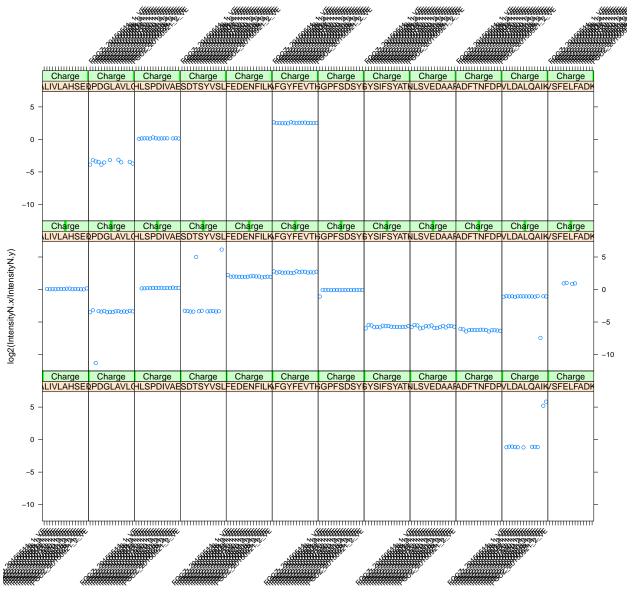




Experiment



Experiment



Experiment

