# Analysing MaxQuant Output with R

#### FGCZ

#### 30 September 2015

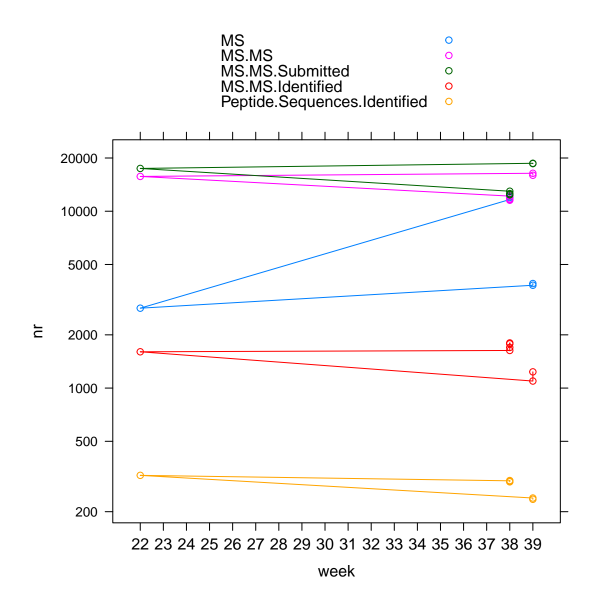
## Contents

Prepare mrm table	1
Analysed Dataset is IMP_DIV	2
Looking at measurement error	3
Looking at MSQC 1 peptide counts	4
Look at Intensities (by species)	5
Looking at intensities msqc1	6
Looking at retention time	8
Log fold change	10

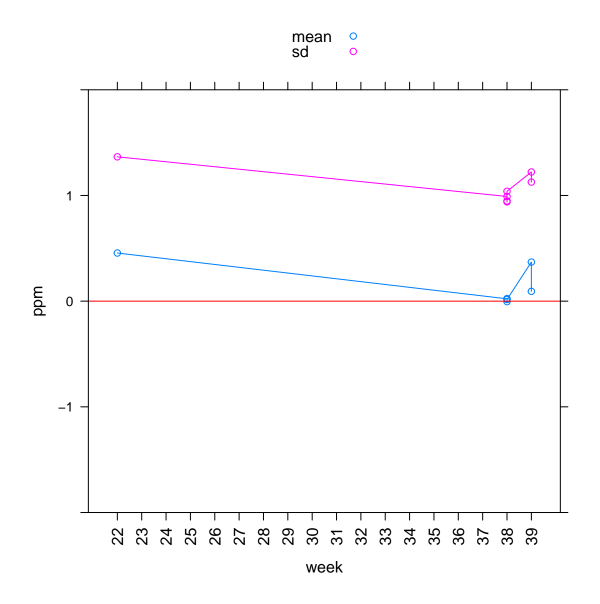
## Prepare mrm table

- 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

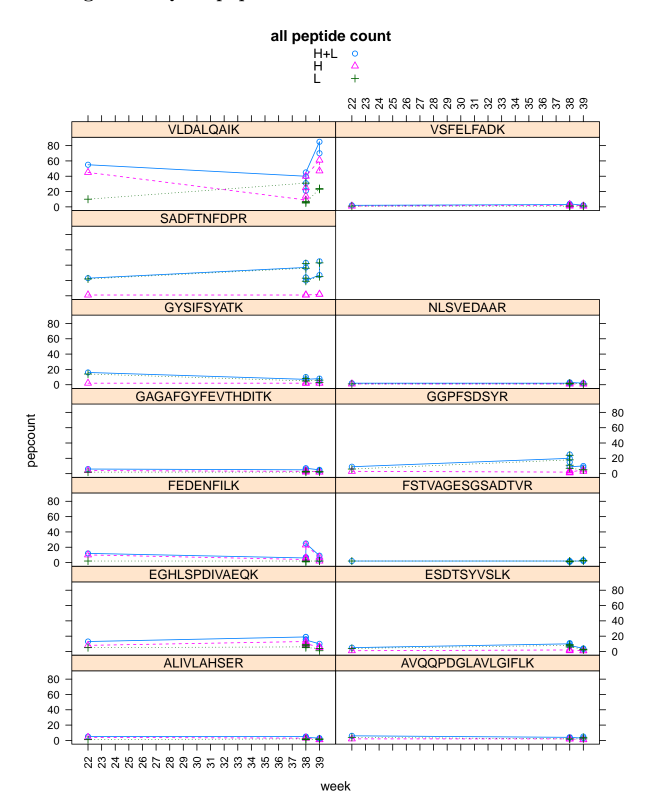
## Analysed Dataset is IMP\_DIV



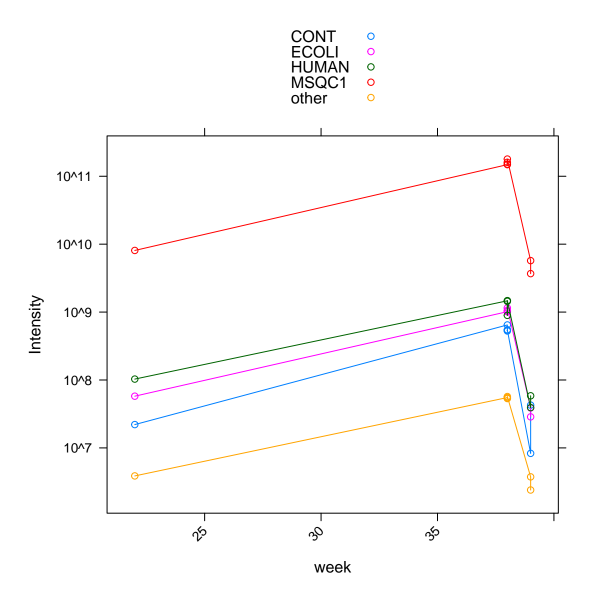
# Looking at measurement error



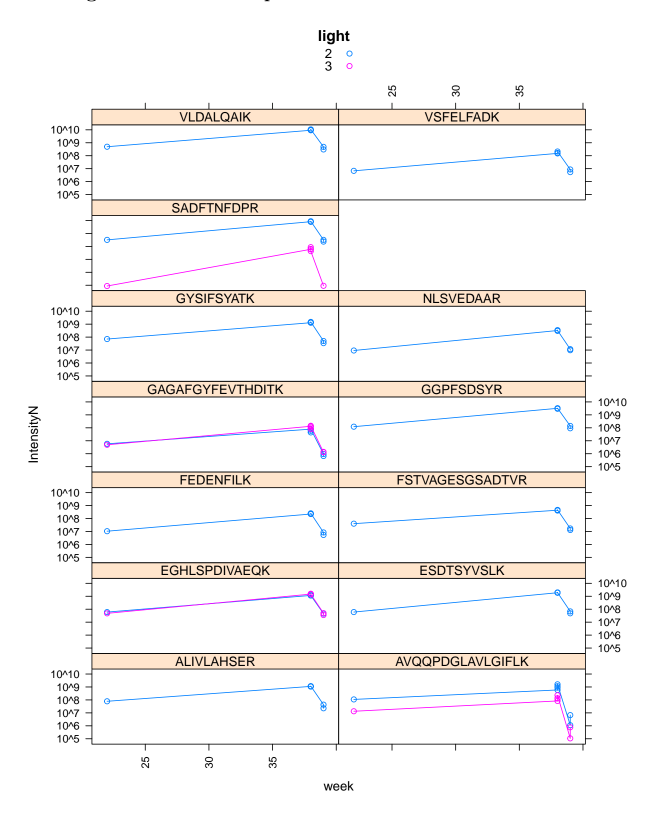
## Looking at MSQC 1 peptide counts

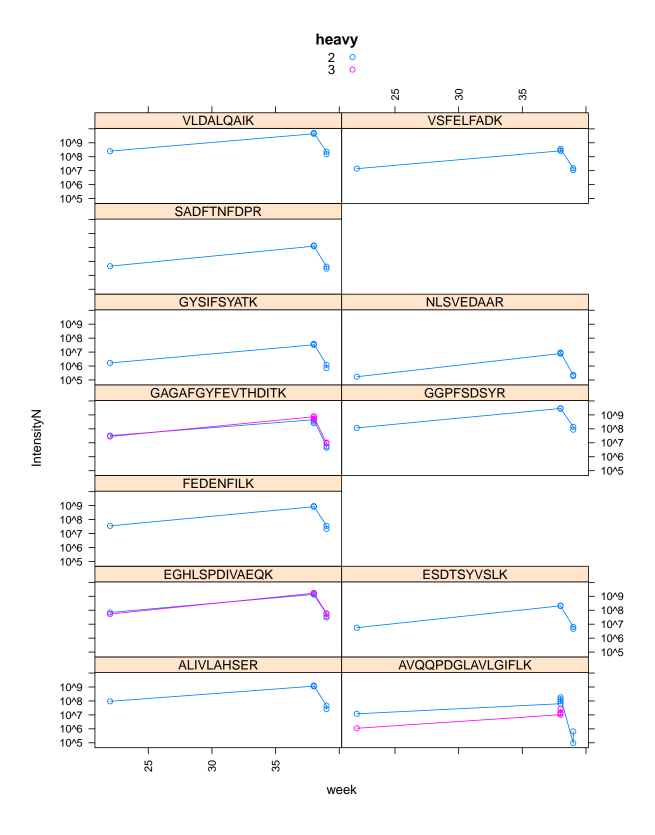


# Look at Intensities (by species)

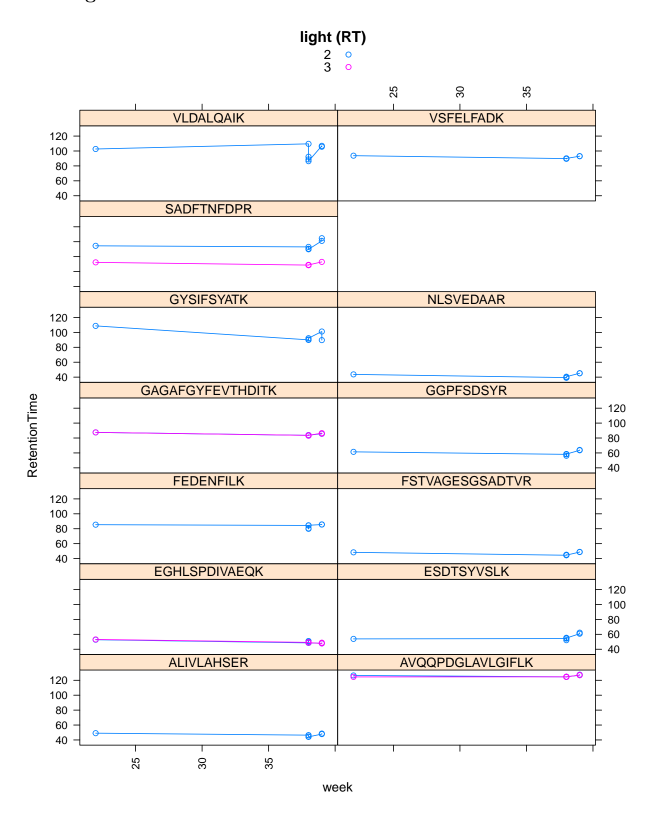


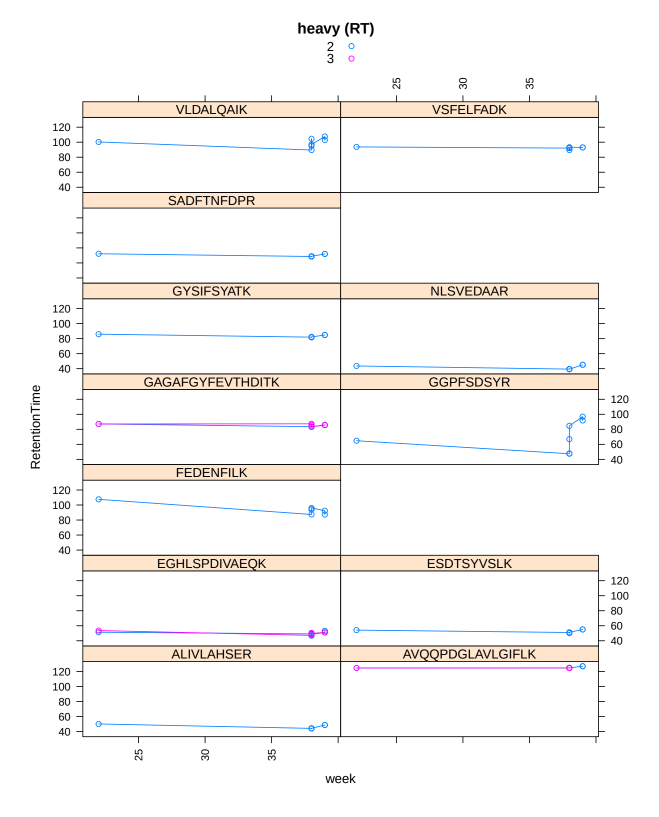
## Looking at intensities msqc1





## Looking at retention time





## Log fold change

## Warning in inner\_join\_impl(x, y, byx, byy): joining factors with ## different levels, coercing to character vector

