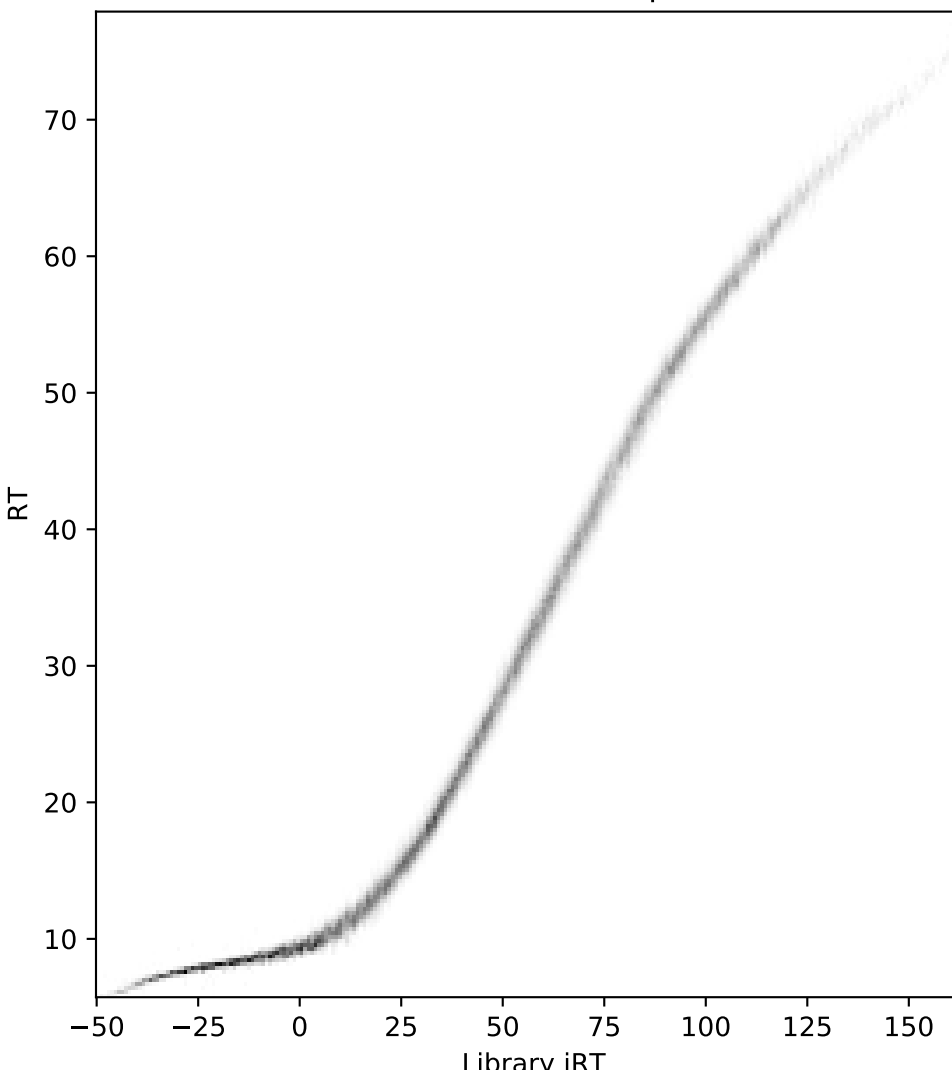
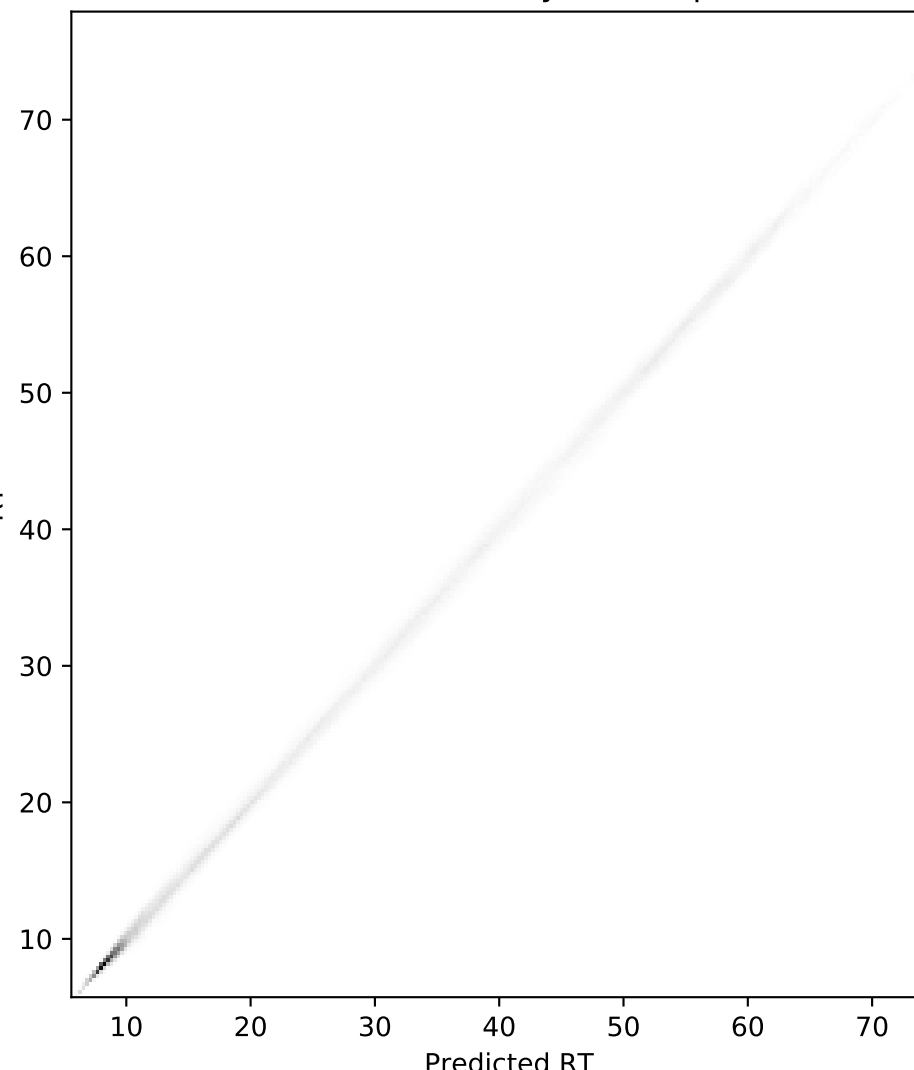


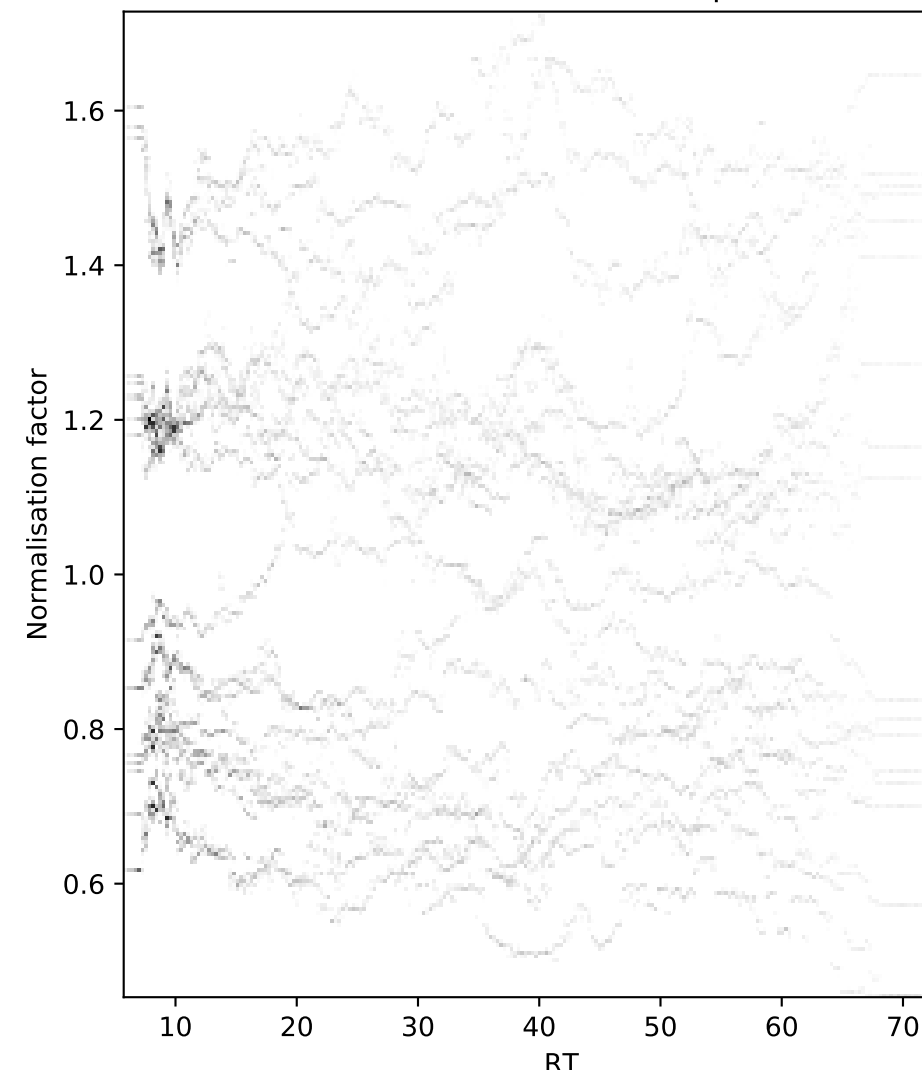
Retention times heatmap, all runs

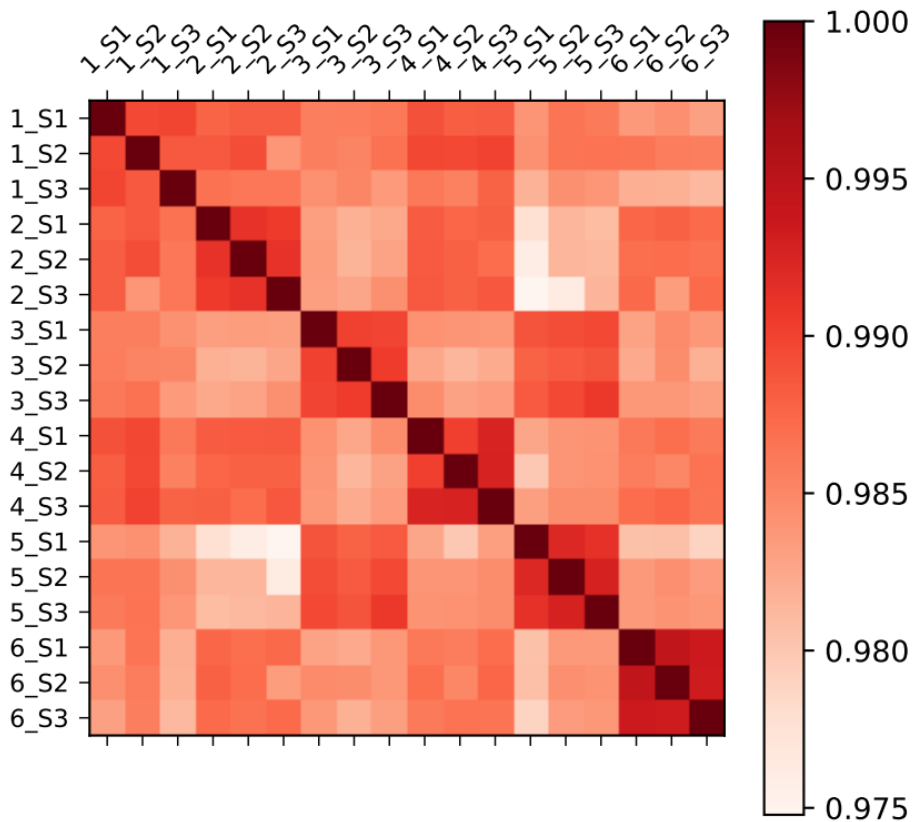


Retention time accuracy heatmap, all runs



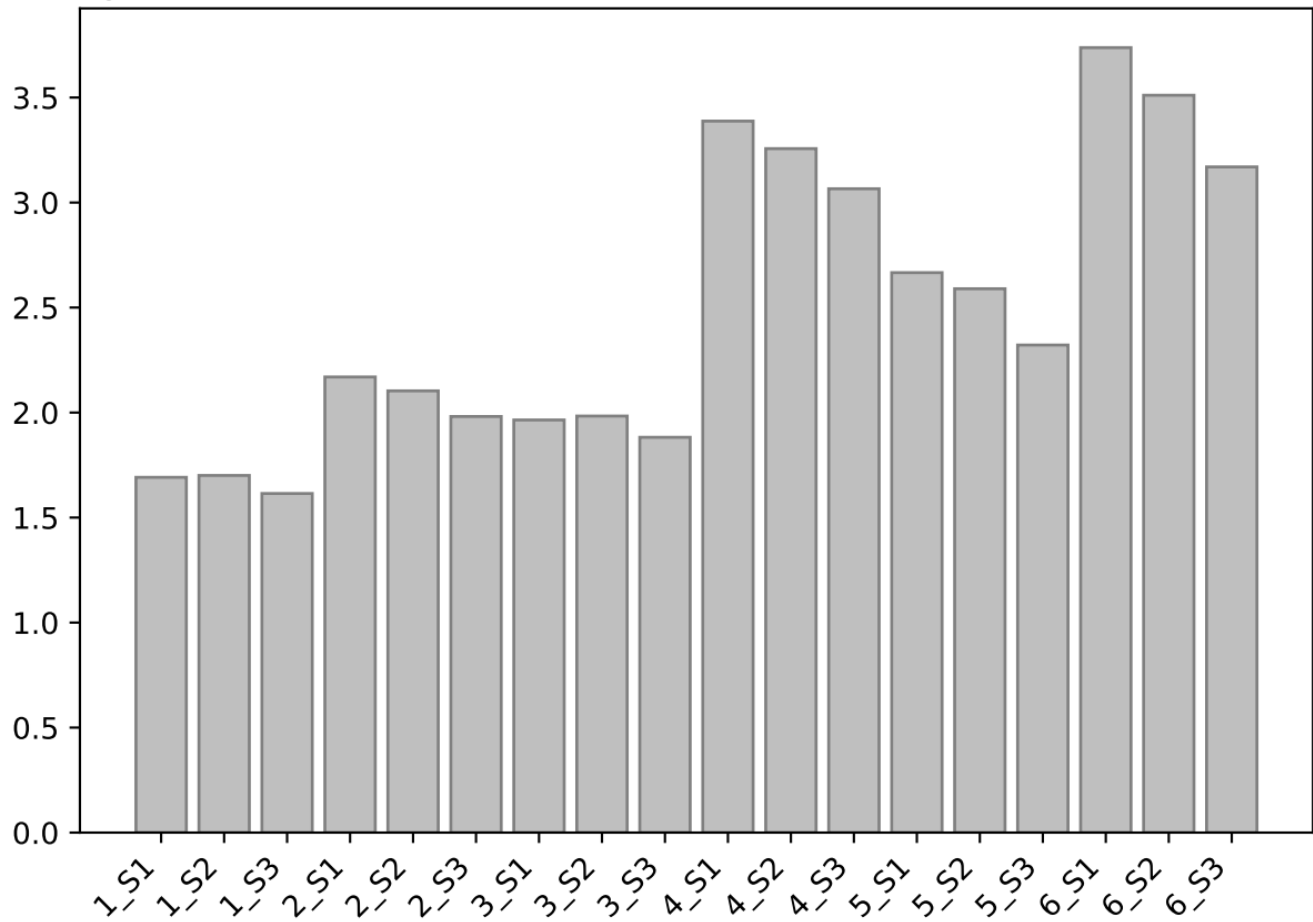
Normalisation factor heatmap, all runs



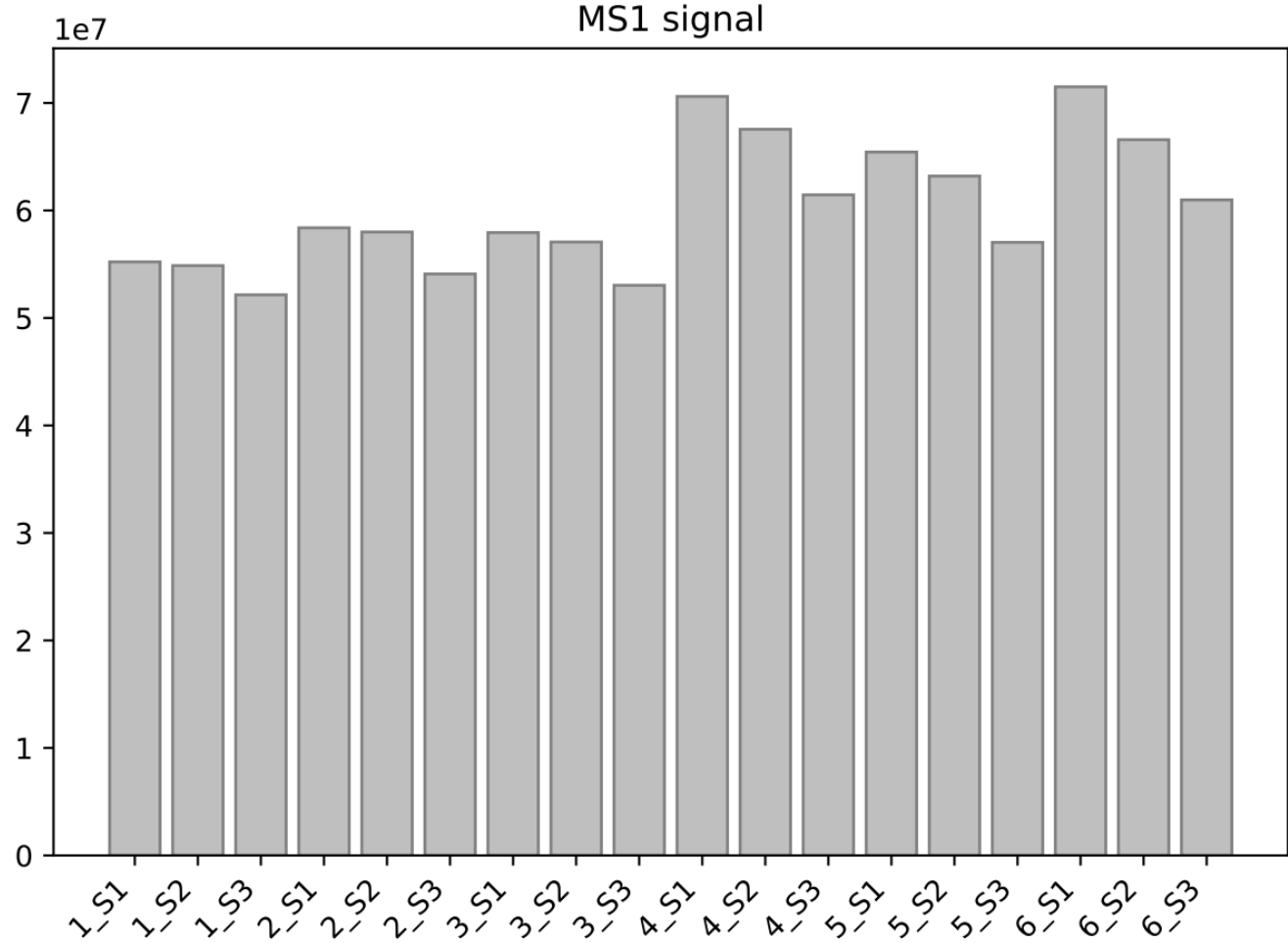


1e7

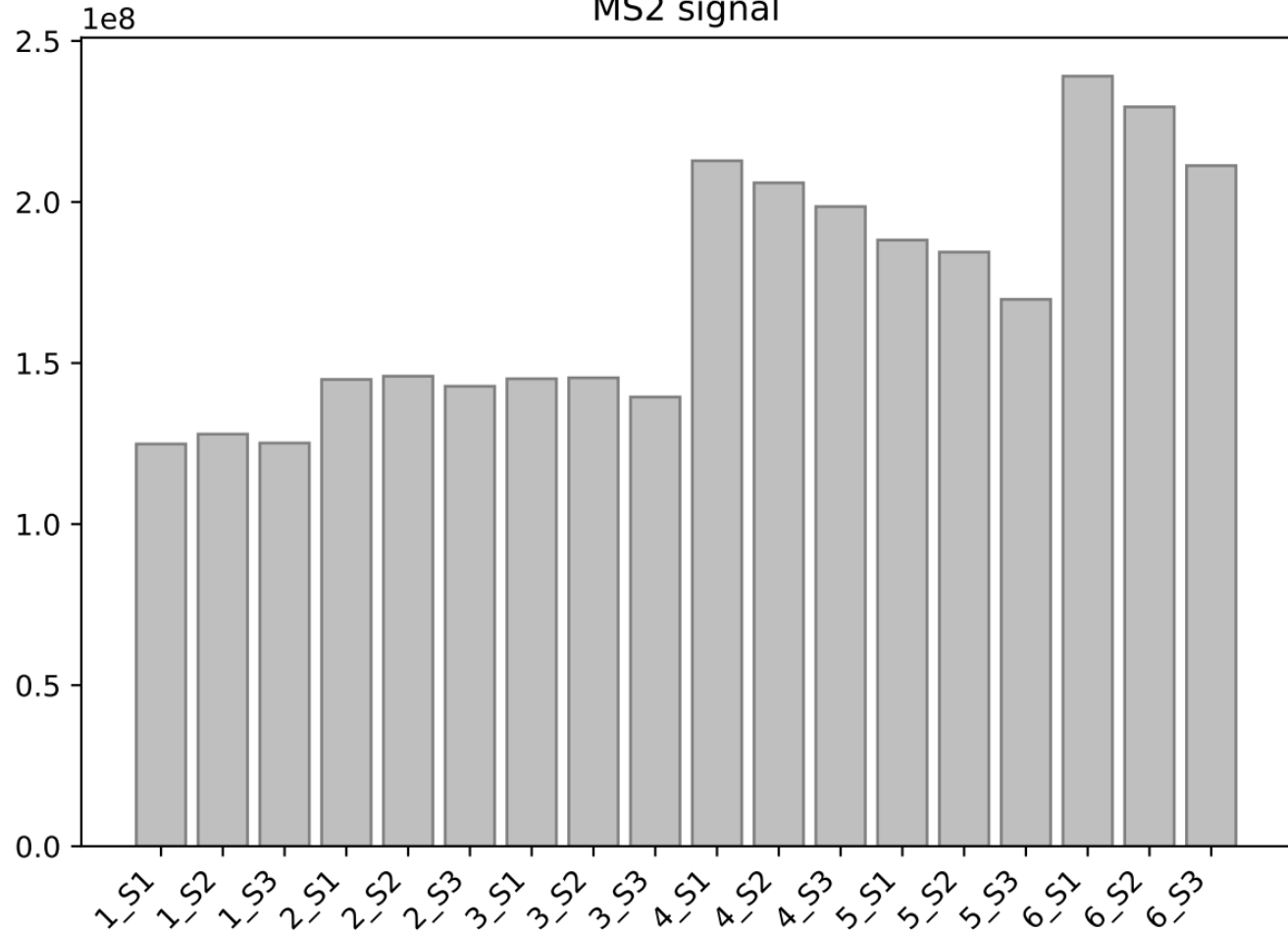
## Total quantity, 1% FDR



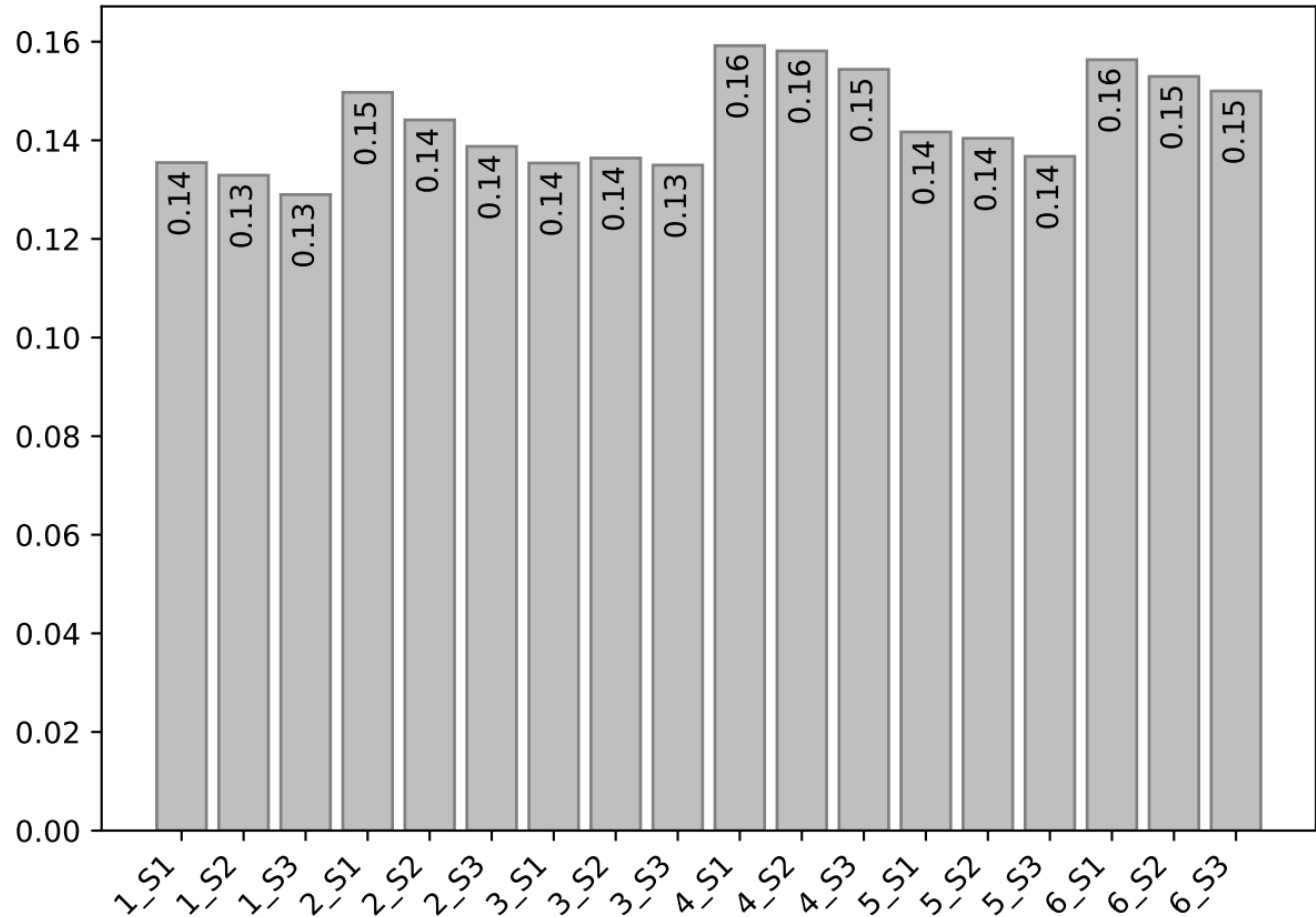
# MS1 signal



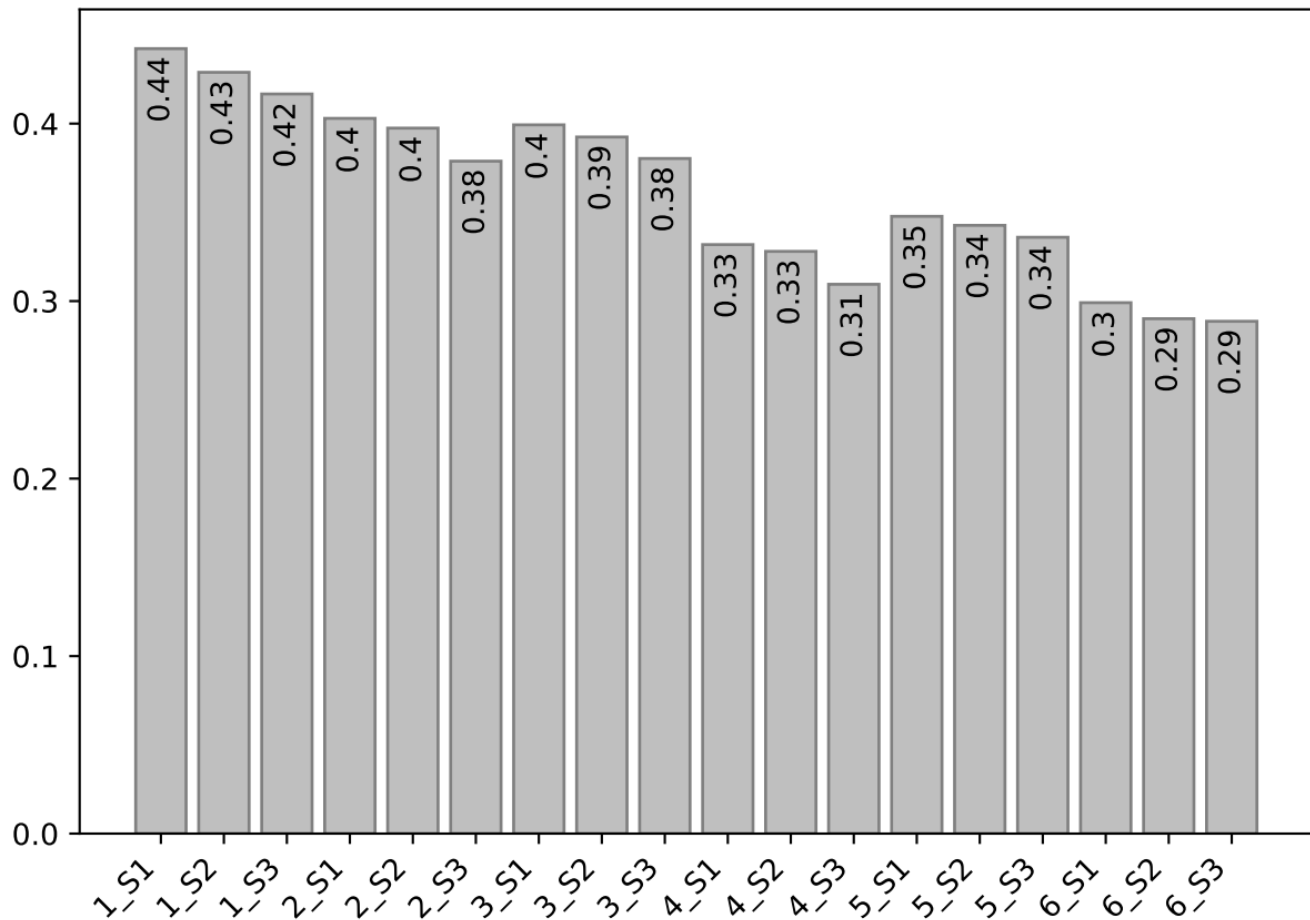
MS2 signal



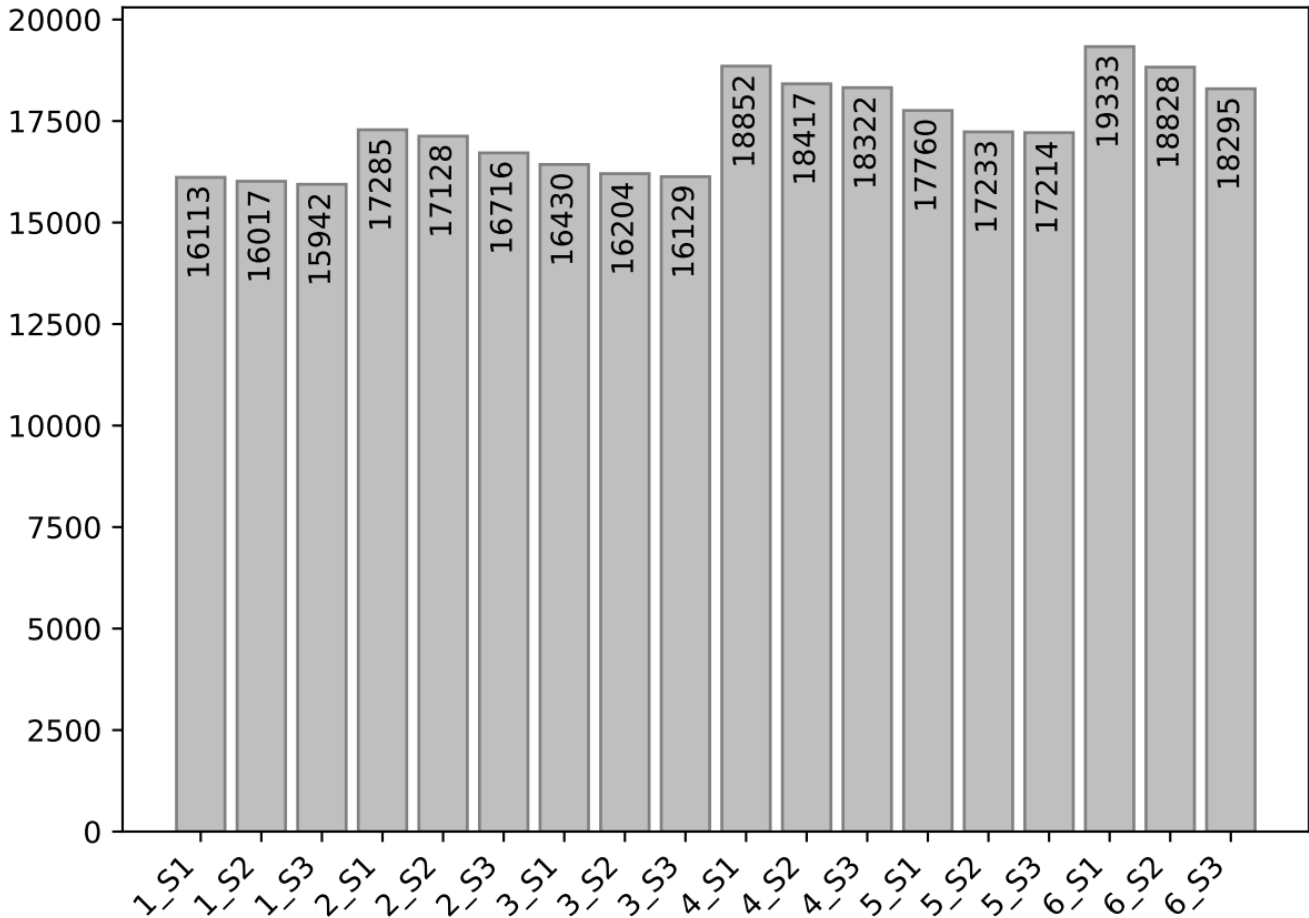
# Total quantity/MS2 signal ratio



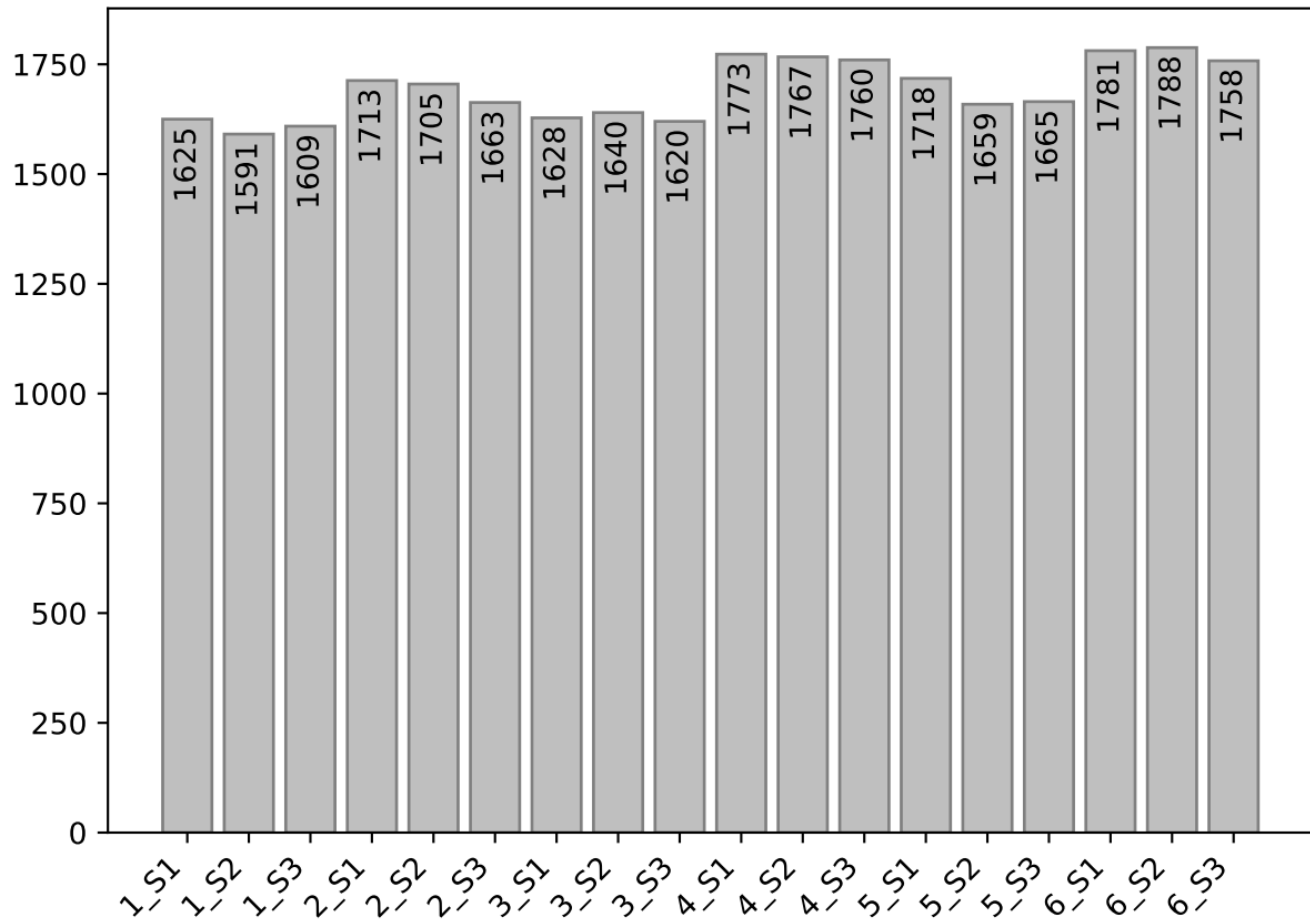
### MS1/MS2 signal ratio



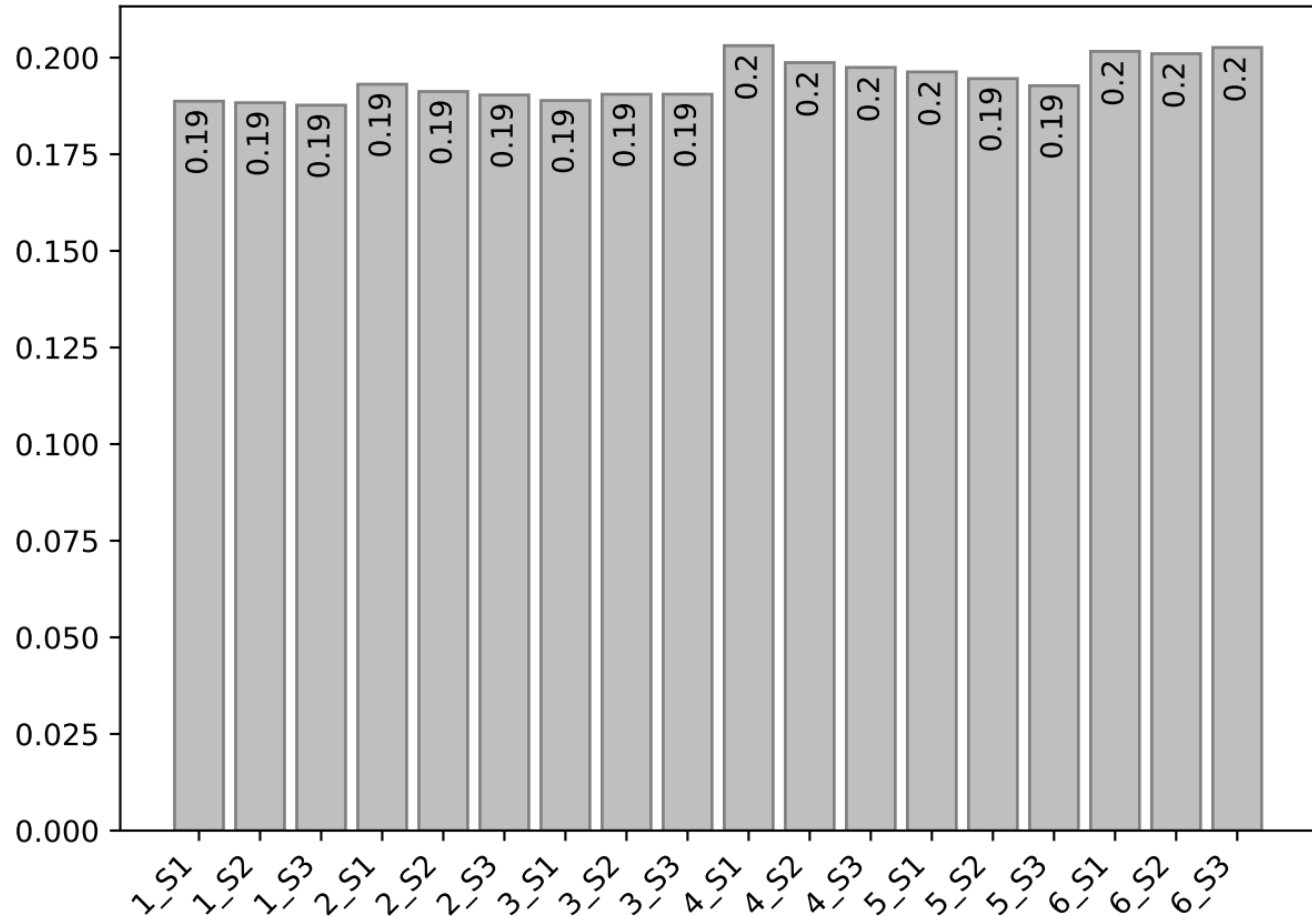
# Precursors, 1% FDR



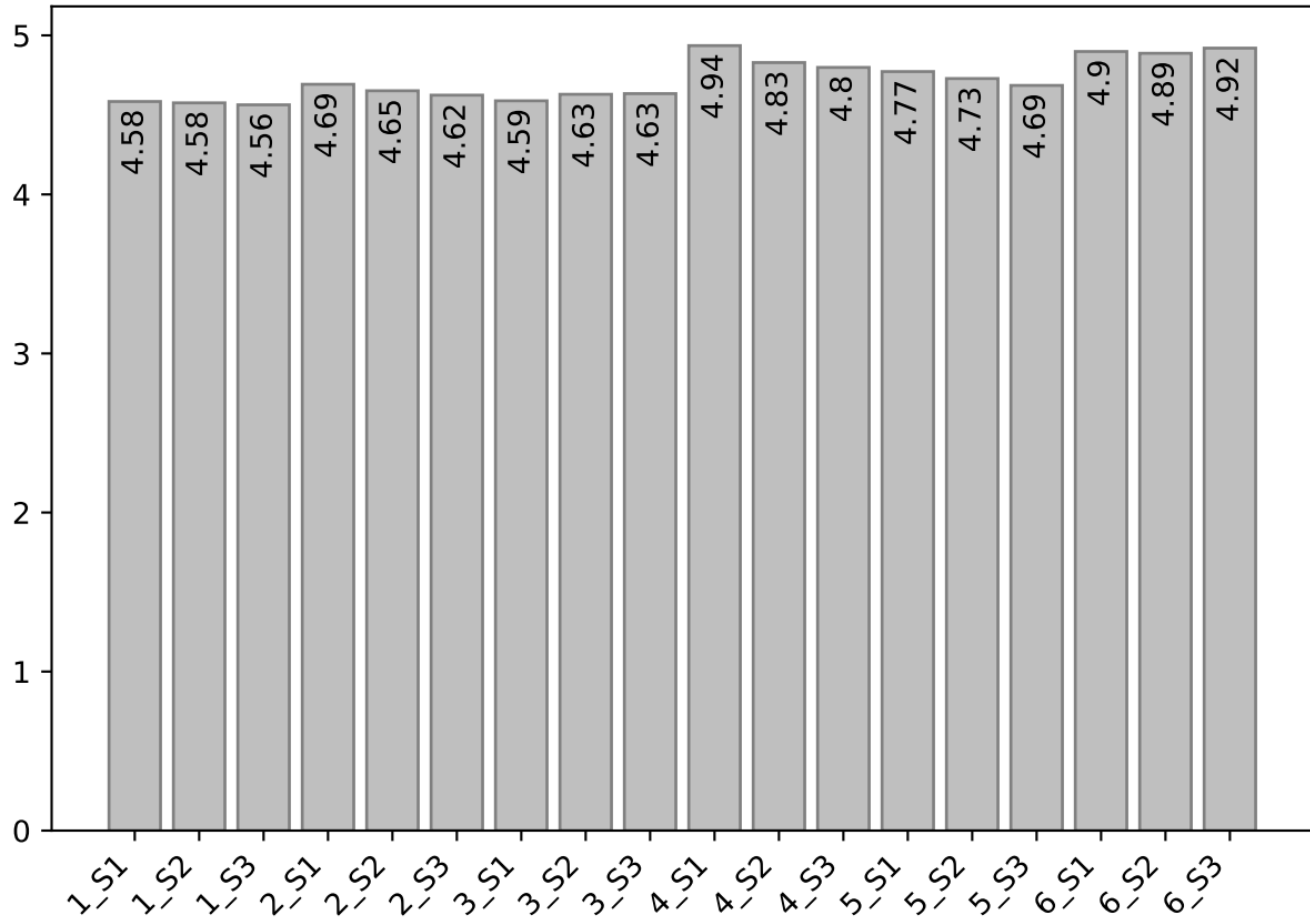
# Unique proteins, 1% protein-level FDR



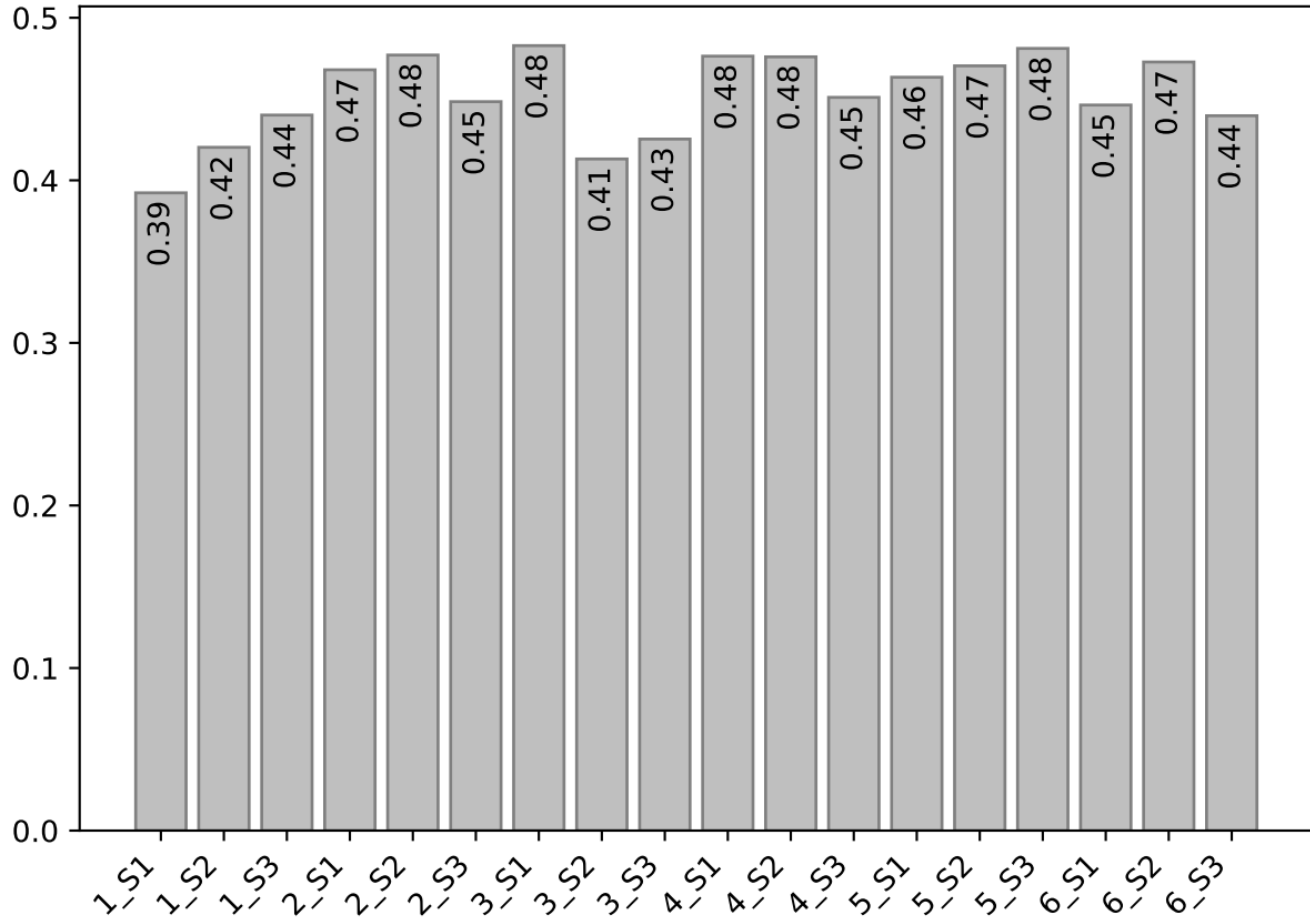
### Mean peak FWHM, in minutes



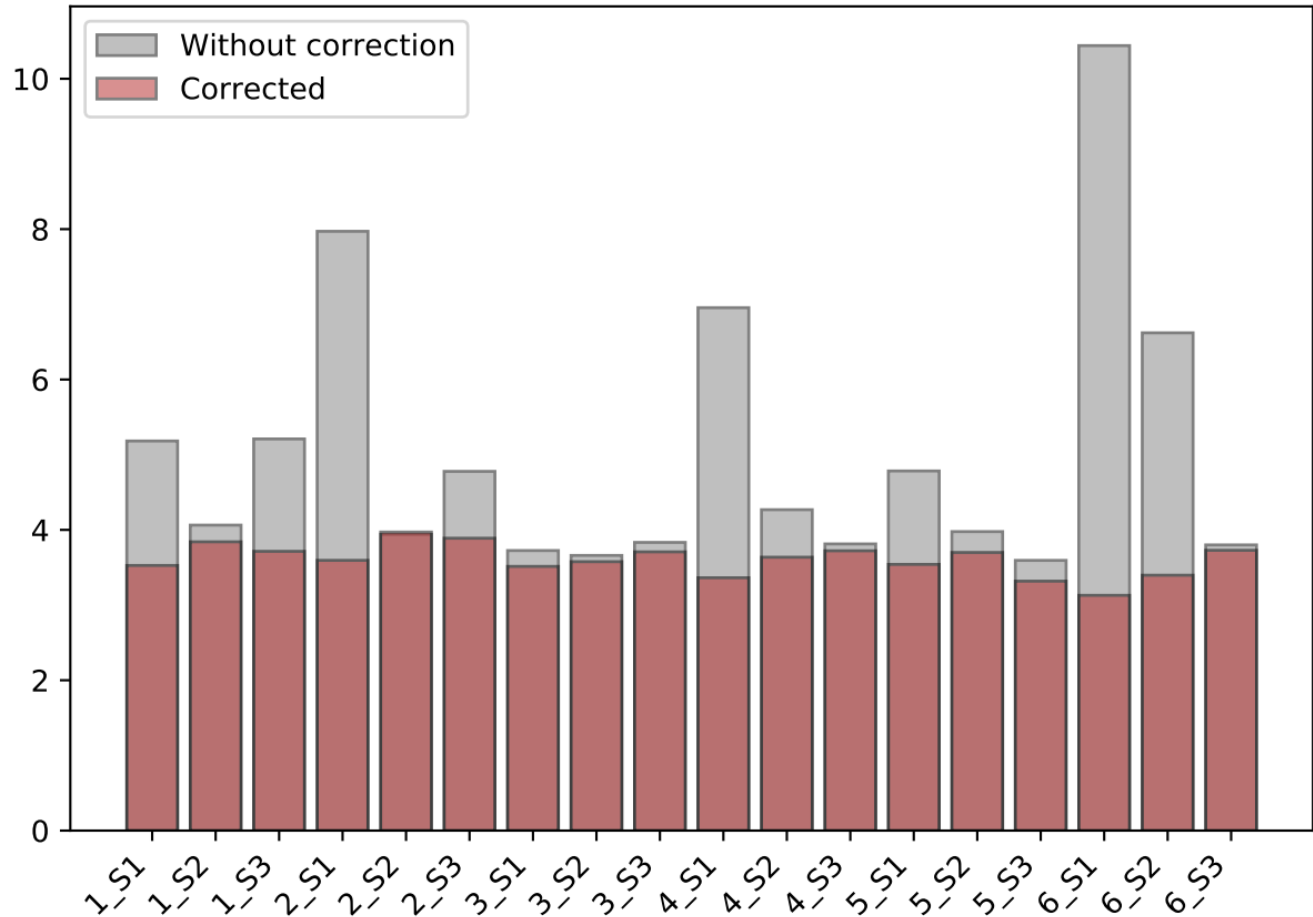
# Mean peak FWHM, in MS2 scans



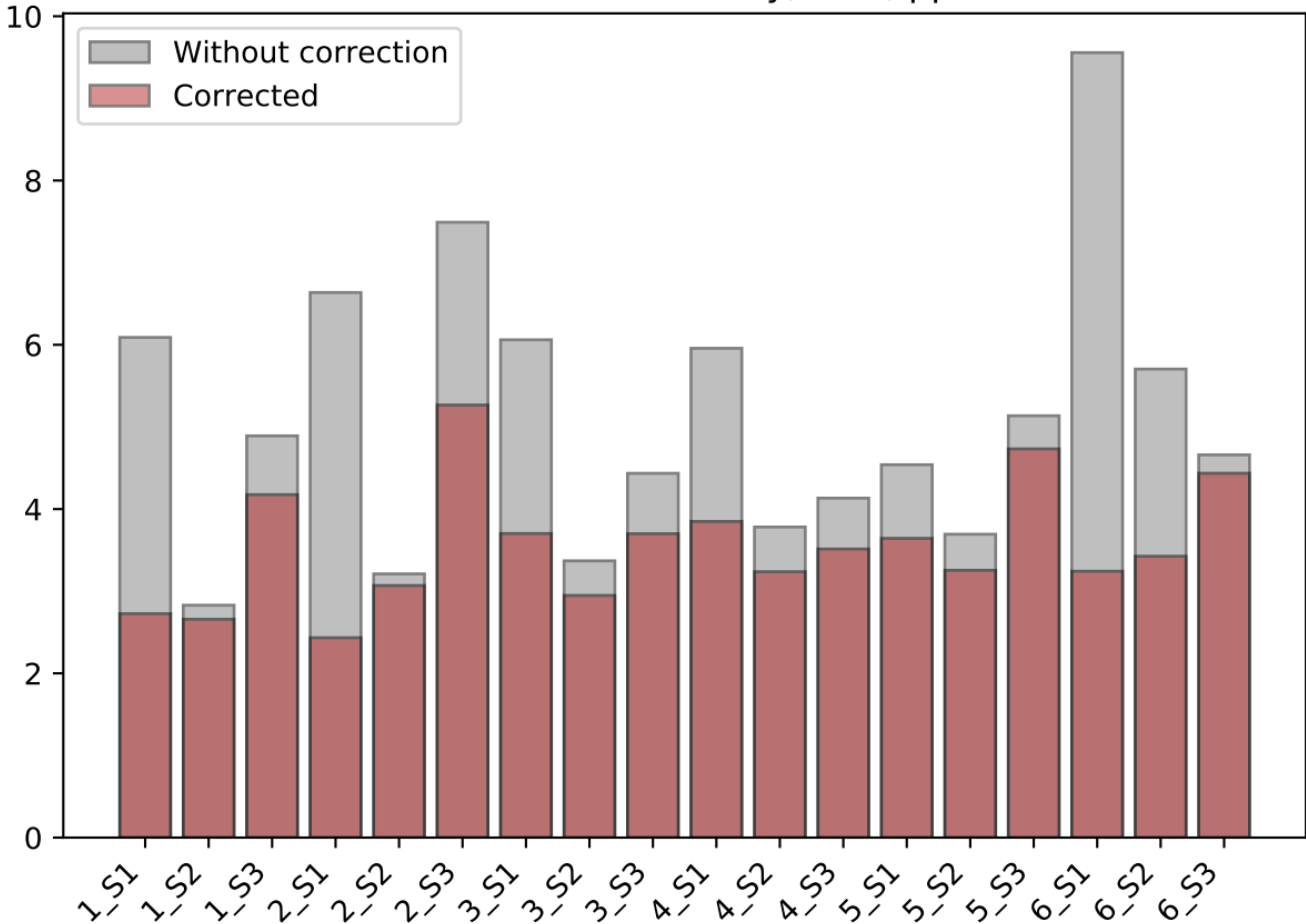
# Median RT prediction accuracy, minutes



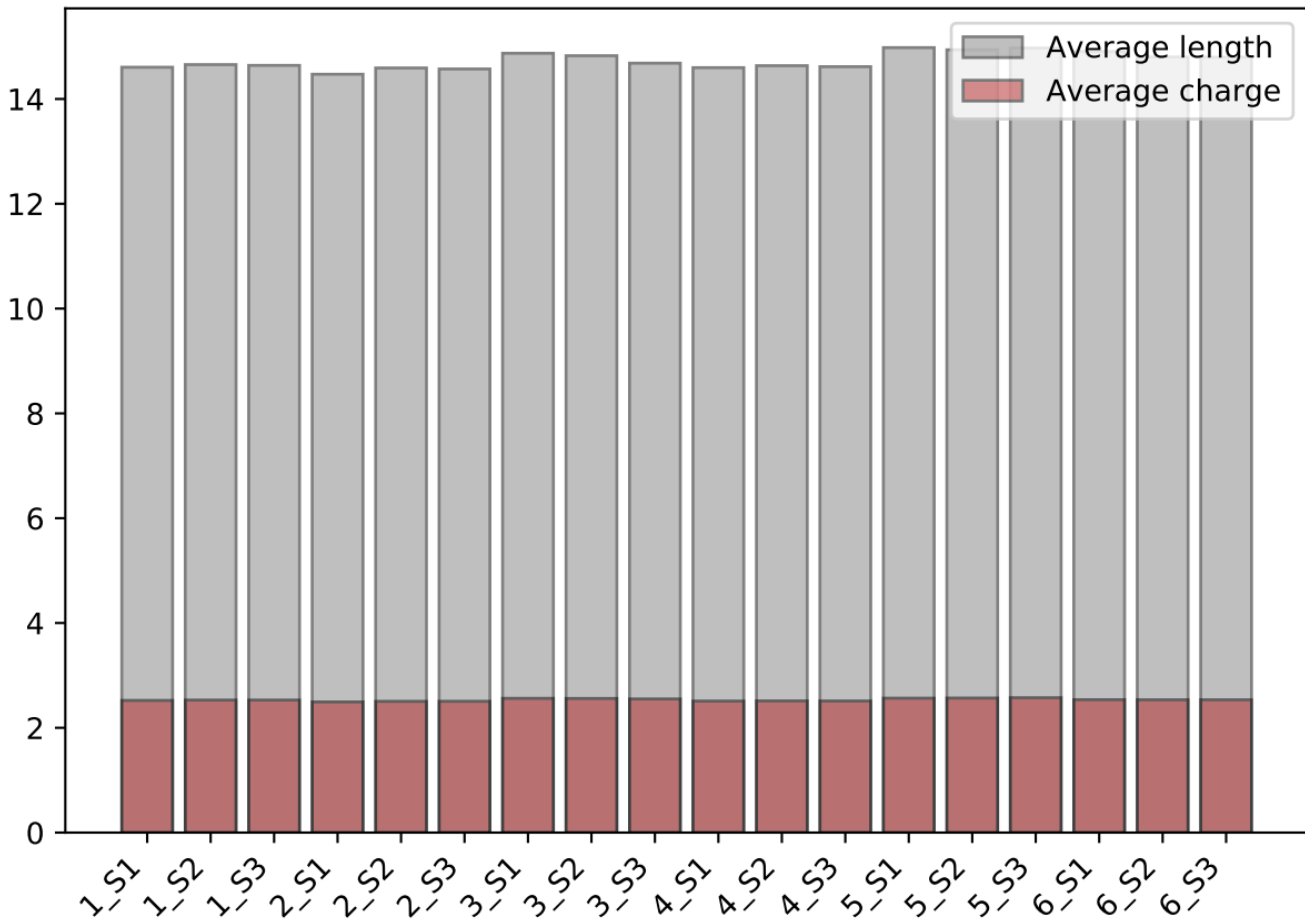
Median mass accuracy, MS2, ppm



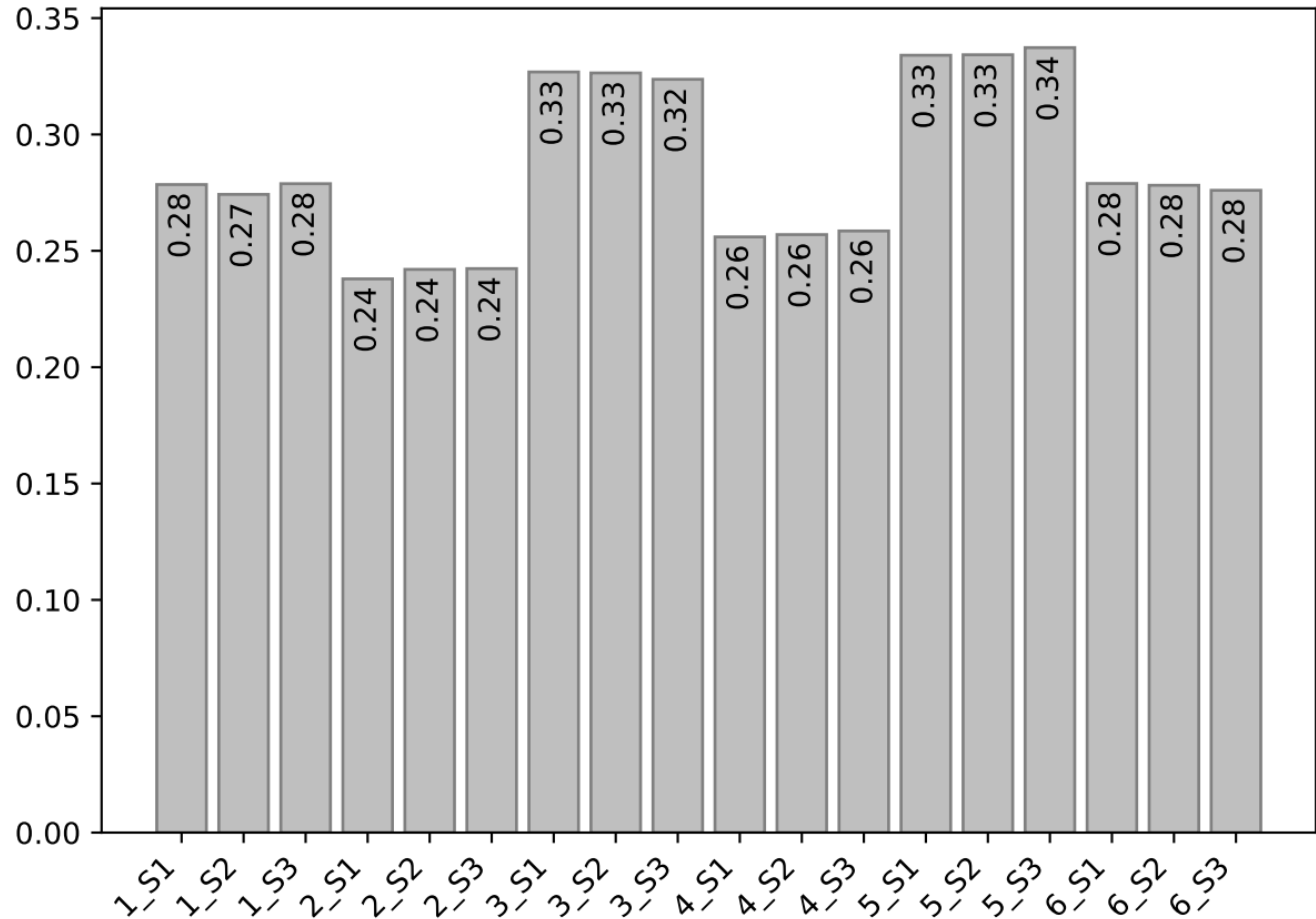
Median mass accuracy, MS1, ppm



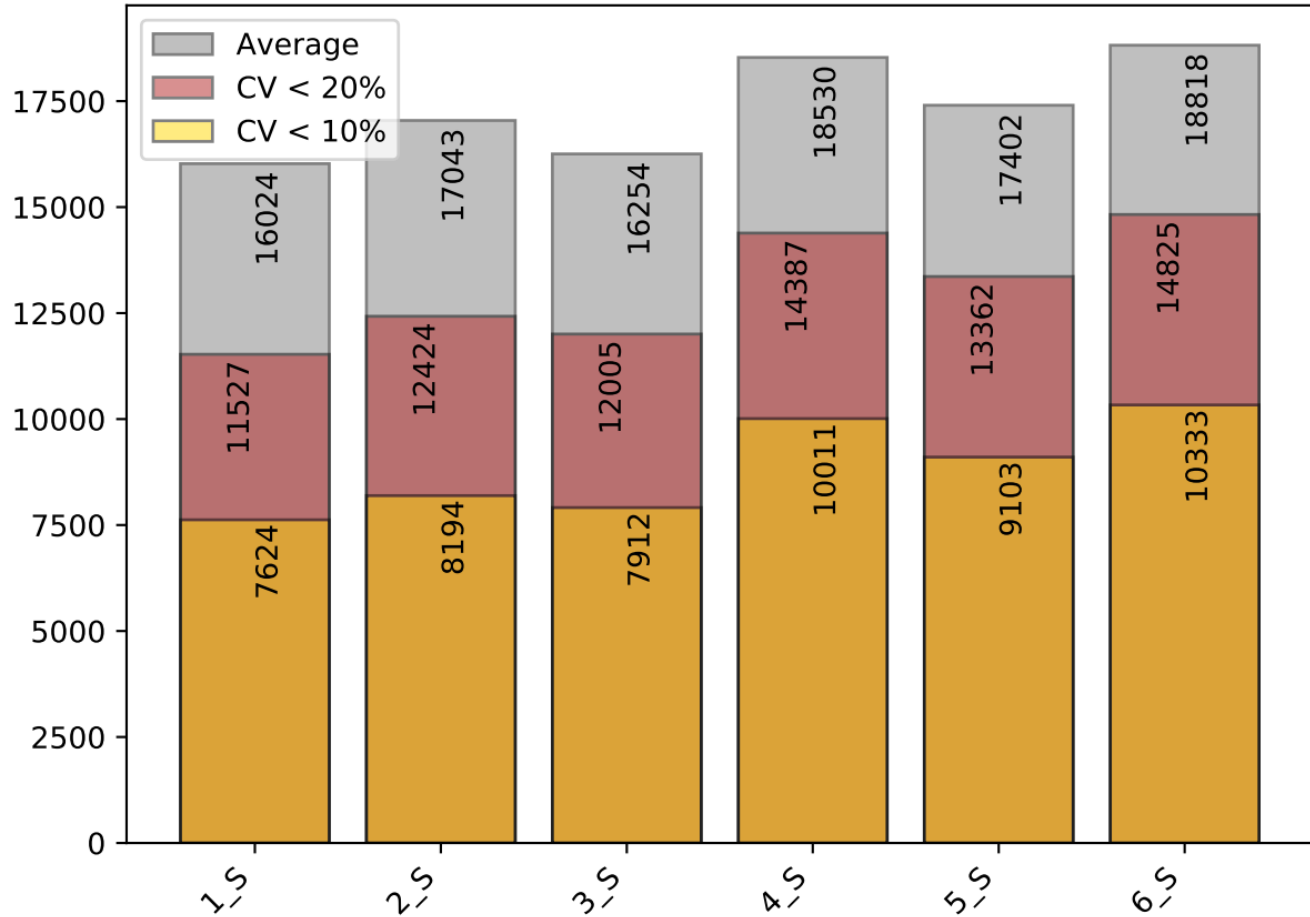
## Peptide characteristics



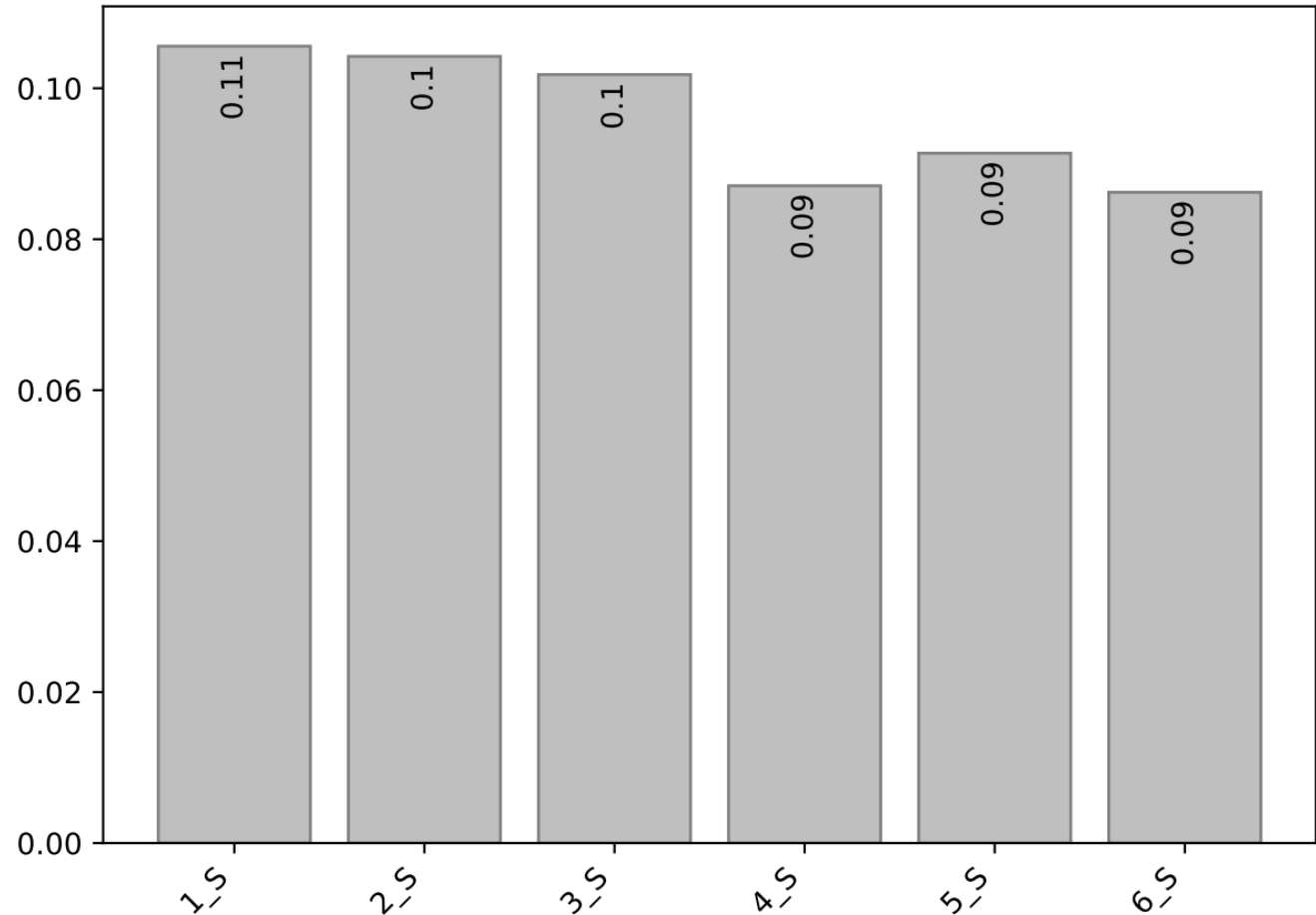
## Average missed tryptic cleavages



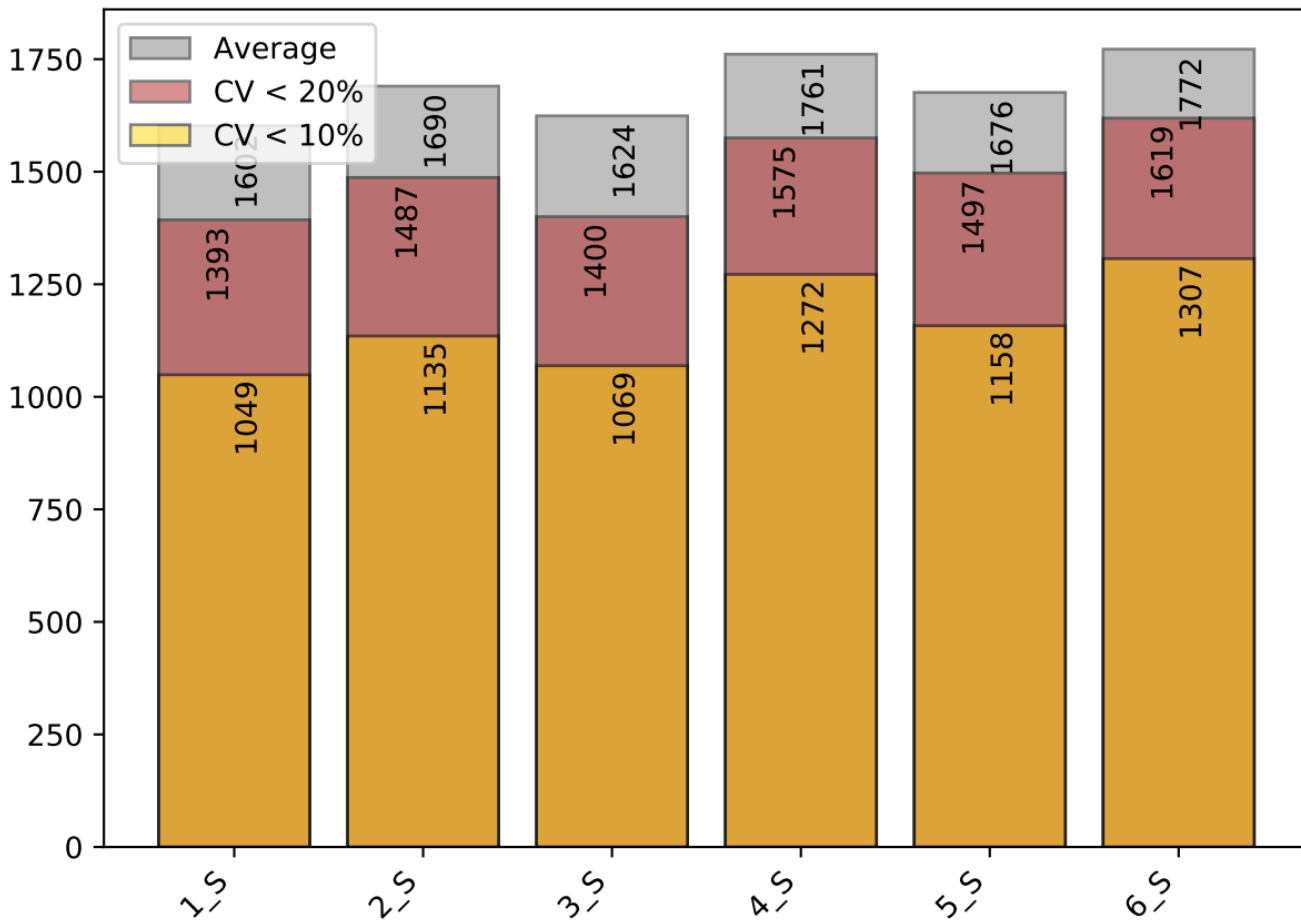
# Precursors, 1% FDR



### Median precursor CV, 1% FDR



# Protein groups, 1% FDR



### Median protein group CV, 1% FDR

