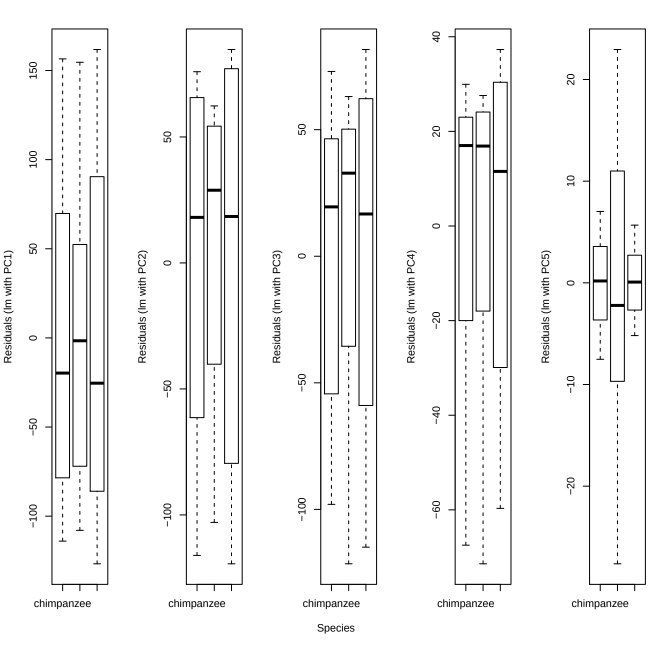
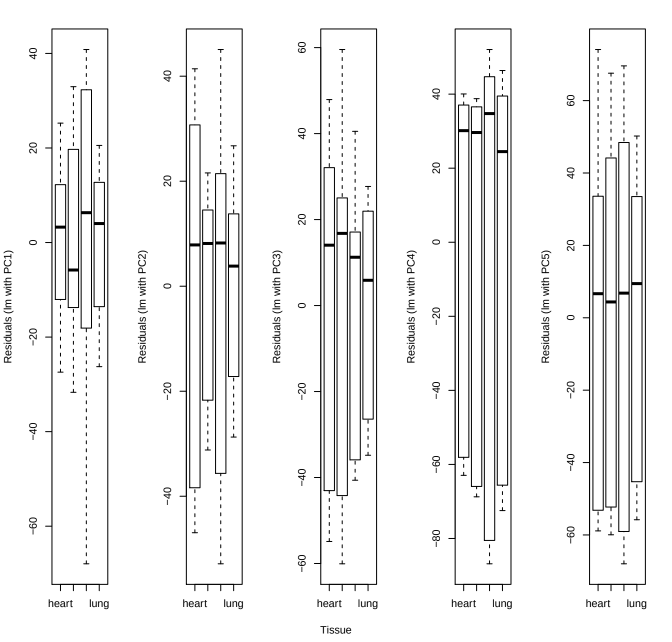
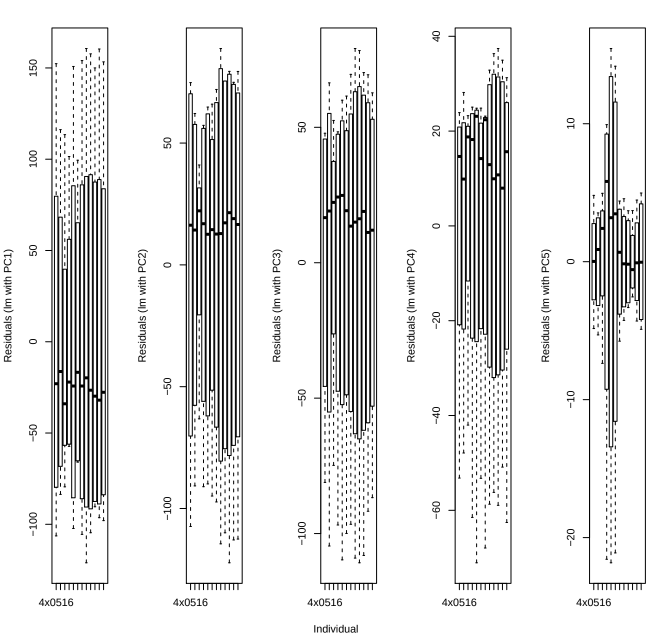
Residuals vs. Species



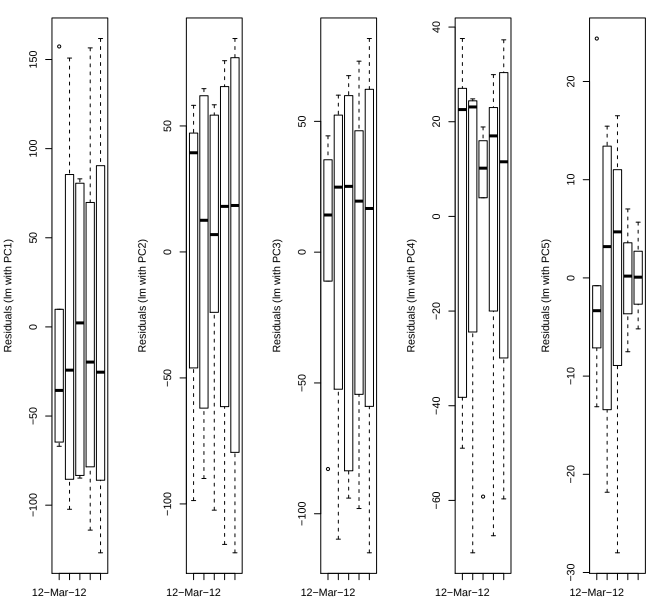
Residuals vs. Tissue



Residuals vs. Individual

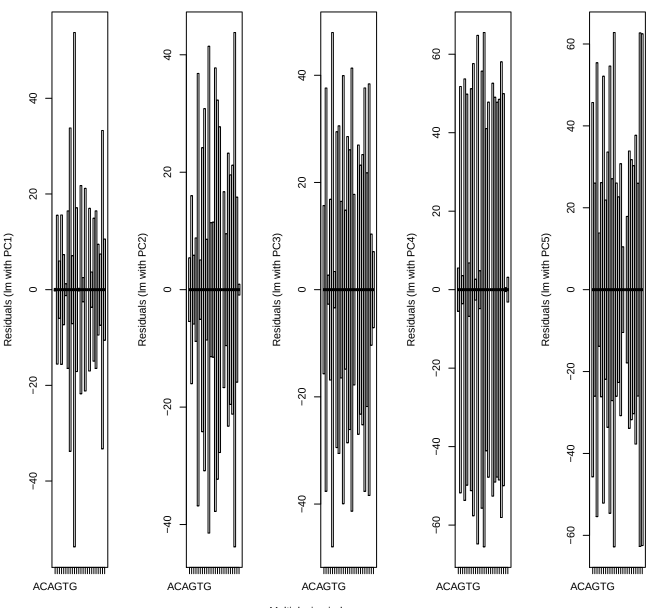


#### Residuals vs. RNA.extraction.date



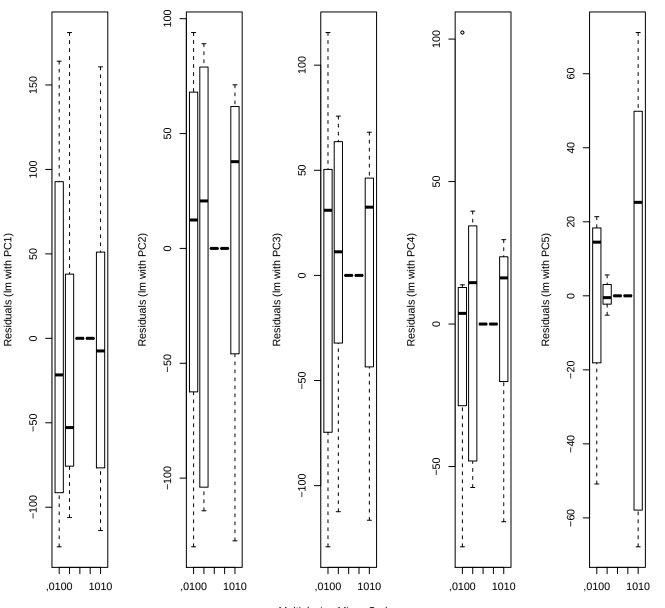
RNA.extraction.date

# Residuals vs. Multiplexing.index.sequence



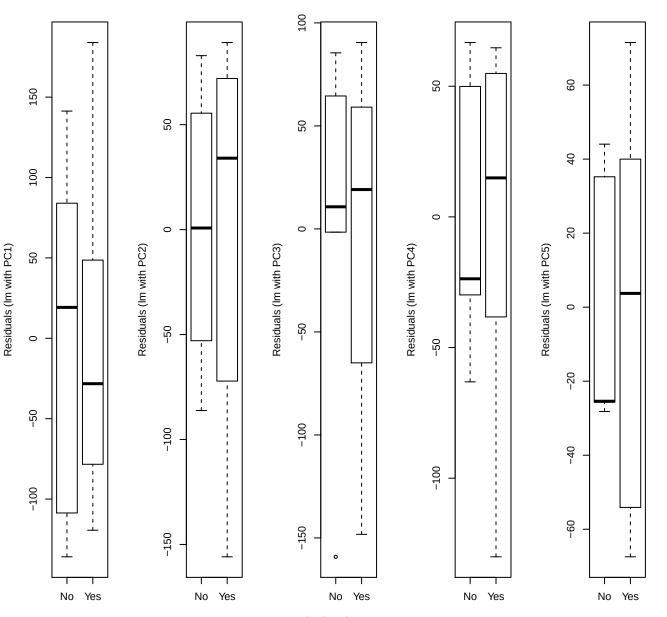
Multiplexing.index.sequence

Residuals vs. Multiplexing.Mixes.Code



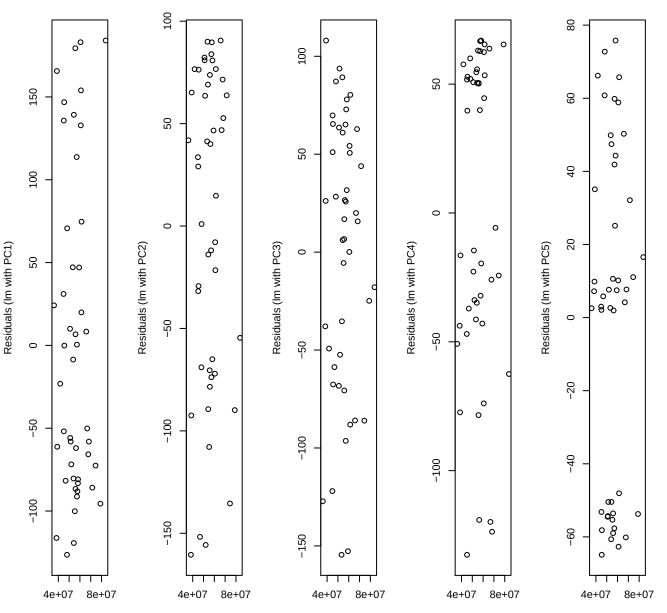
Multiplexing.Mixes.Code

### Residuals vs. Sequencing.location..at.UCGF



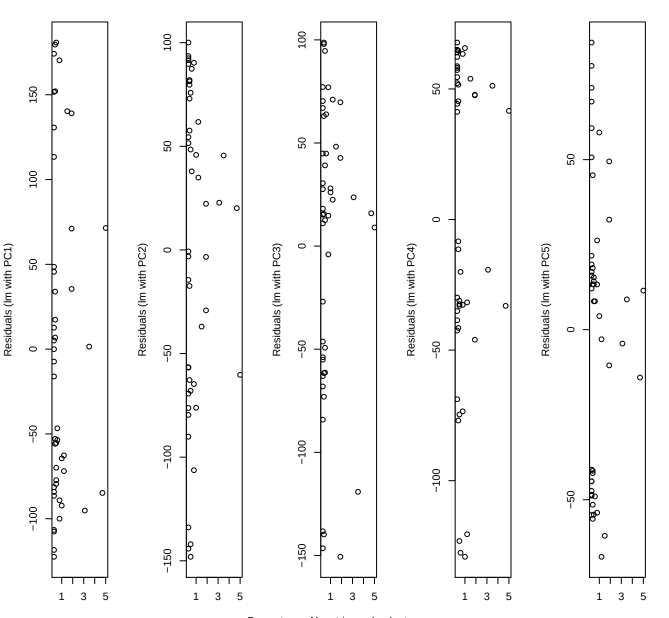
Sequencing.location..at.UCGF

#### Residuals vs. Total.number.of.reads.sequenced



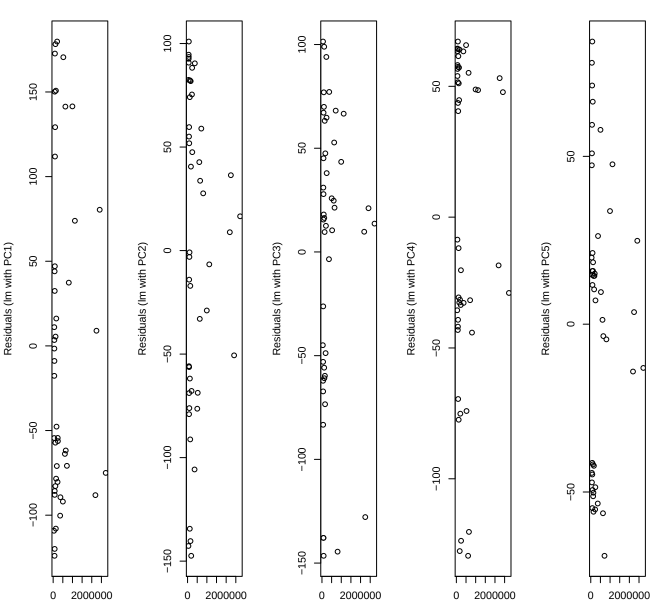
Total.number.of.reads.sequenced

Residuals vs. Percentage.of.bps.trimmed..adapters.

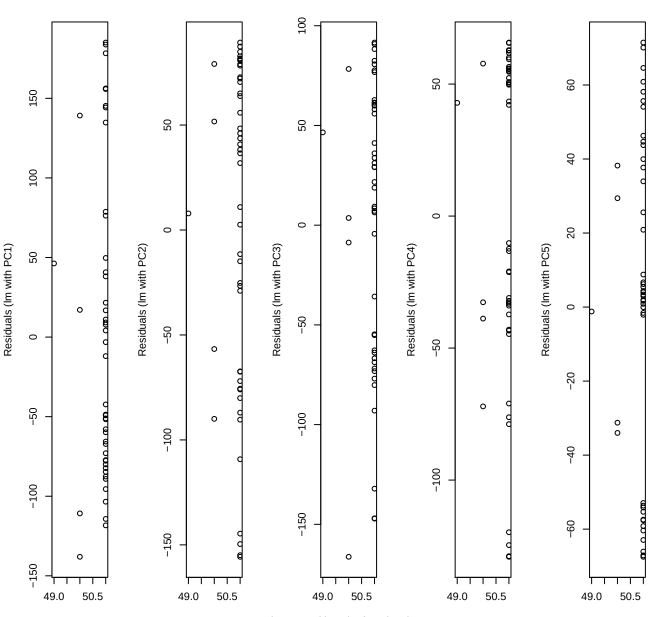


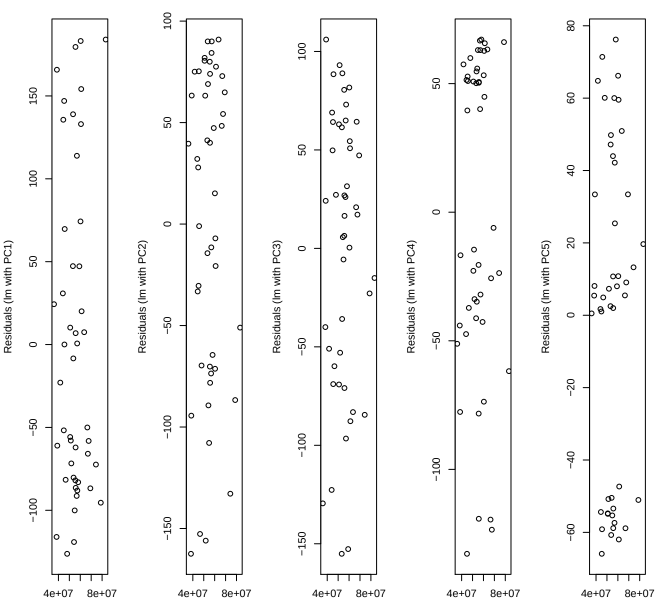
Percentage. of. bps. trimmed.. adapters.

Residuals vs. Number.of.reads.shorter.than.20bp.removed



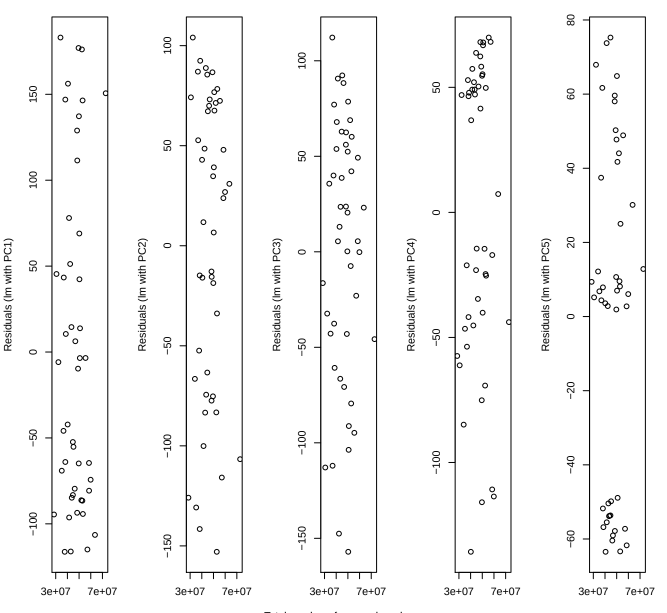
Residuals vs. Maximum.read.length.after.trimming





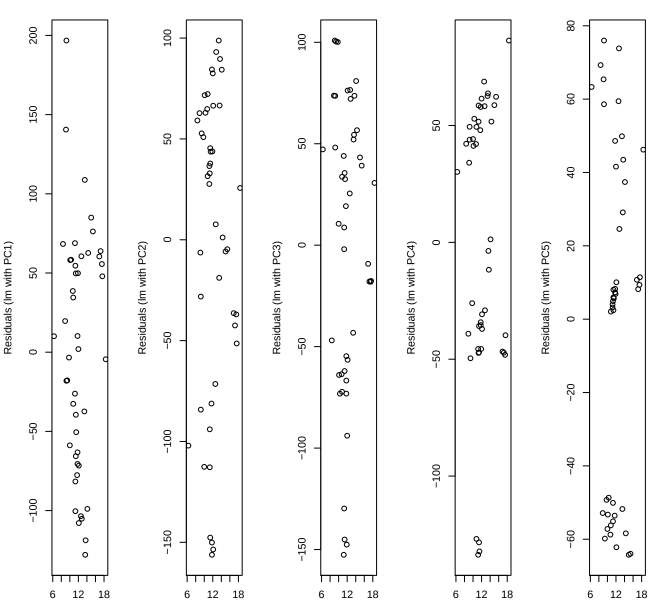
Total.number. of. reads. processed. in. tophat

#### Residuals vs. Total.number.of.mapped.reads

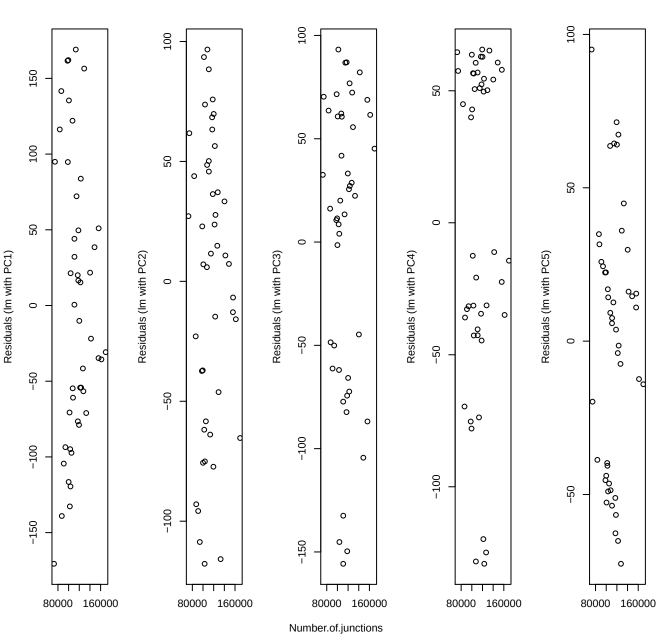


Total.number.of. mapped.reads

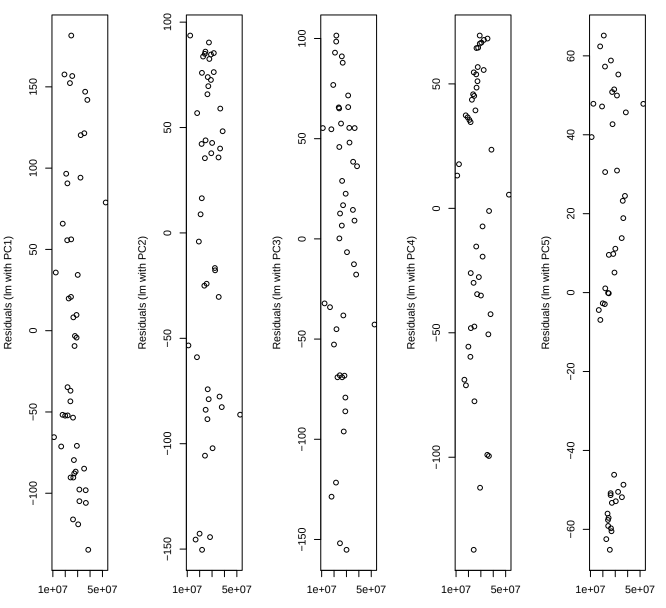
Residuals vs. Percentage.of.mapped.reads.overlapping.a.junction

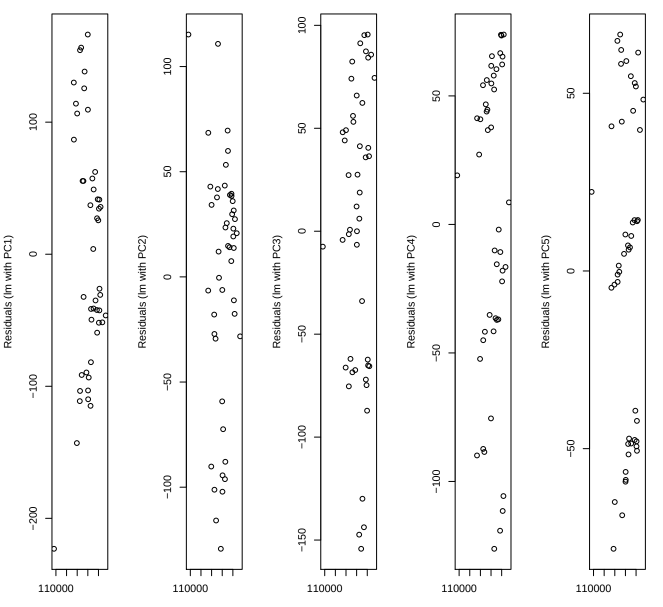


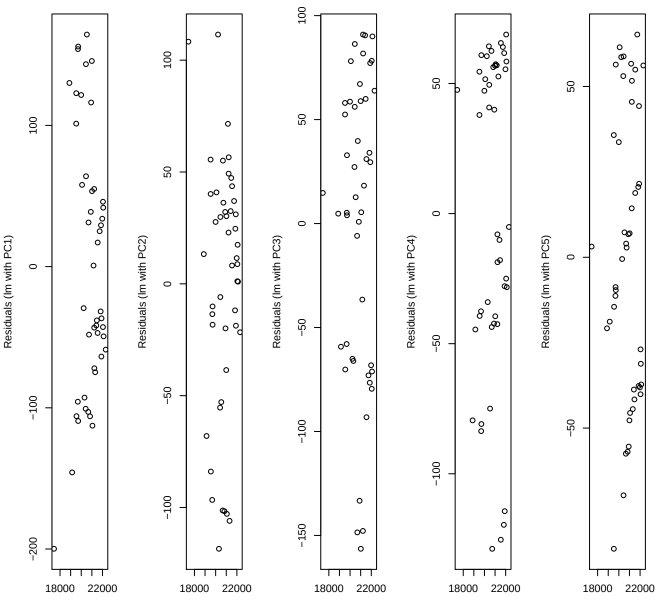
## Residuals vs. Number.of.junctions



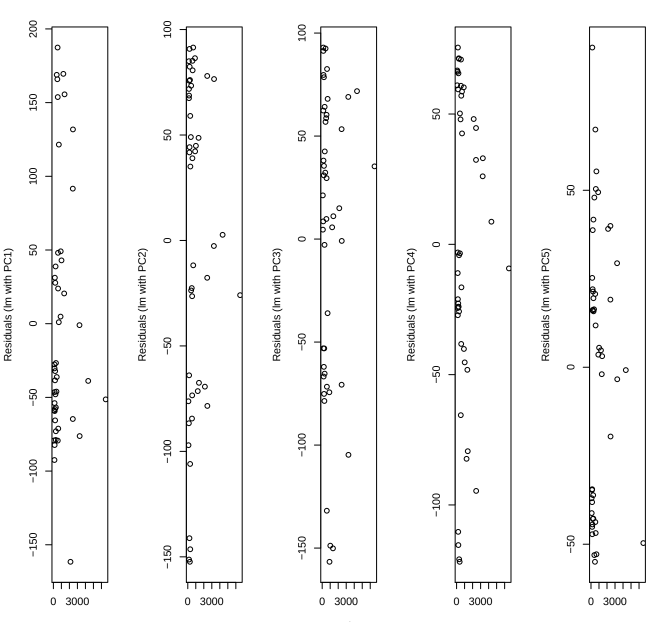
Residuals vs. Number.of.reads.mapped.on.orthologous.exons





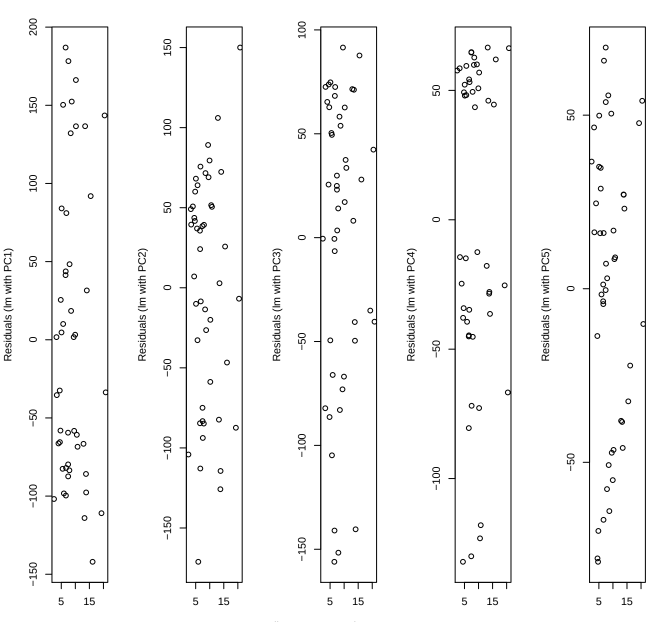


Residuals vs. RNA.concentration..ng.uL.



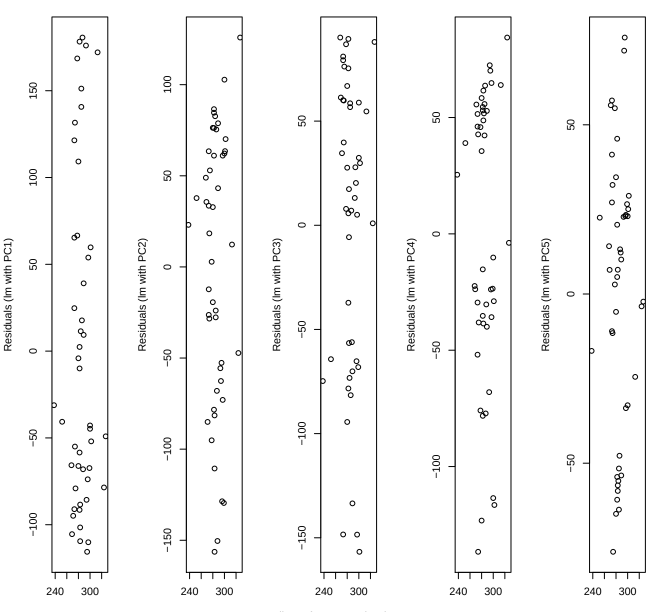
RNA.concentration..ng.uL.

Residuals vs. Library.concentration..ng.uL.



Library.concentration..ng.uL.

Residuals vs. Library.fragments.size..bp.



Library.fragments.size..bp.

# enjamini and Hochbeenjamini and Hochbe

