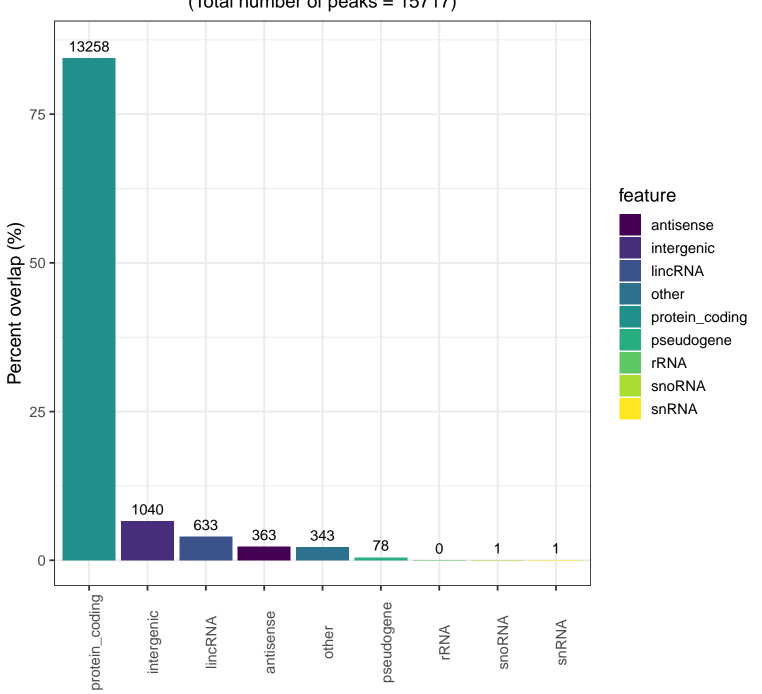
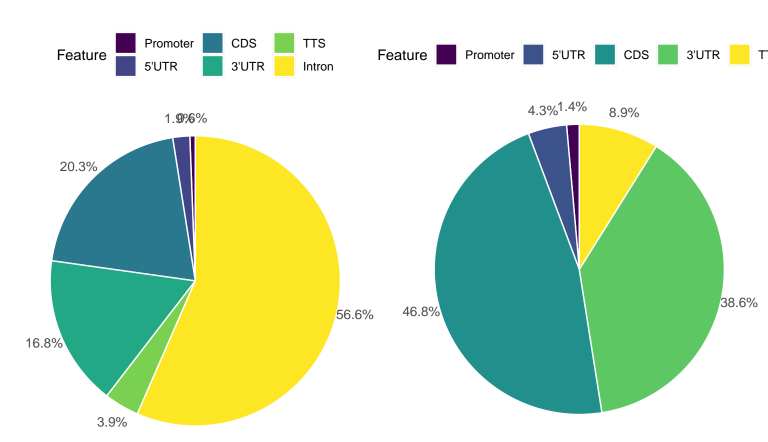
## Annotation of peaks to all type of genes

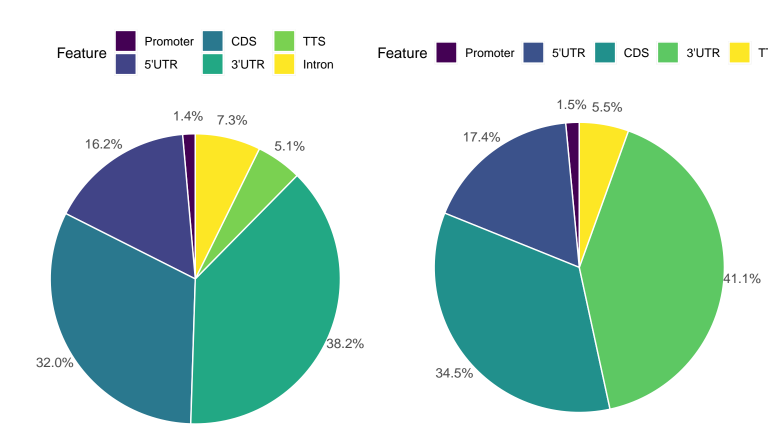
(Total number of peaks = 15717)



## **Absolute counts**



## Length-normalized counts



## Plotting parameters:

functionName: "plot\_peak\_annotation"

peakFile: c(iCLIPPeak =

"C:/GREENBLATT/Rscripts/GenomicPlot\_gh/inst/extdata/test\_clip\_peak\_chr19.bed") gtfFile: "C:/GREENBLATT/Rscripts/GenomicPlot\_gh/inst/extdata/gencode.v19.annotation\_chr19.gtf"

importParams: list(offset = 0, fix\_width = 21, fix\_point = "center", norm = FALSE, useScore =

FALSE, outRle = FALSE, useSizeFactor = FALSE, genome = "hg19")

fiveP: -1000 dsTSS: 0

threeP: 2000

simple: FALSE

outPrefix: "test\_plot\_peak\_annotation2"

verbose: FALSE

hw: c(8, 8)

nc: 2