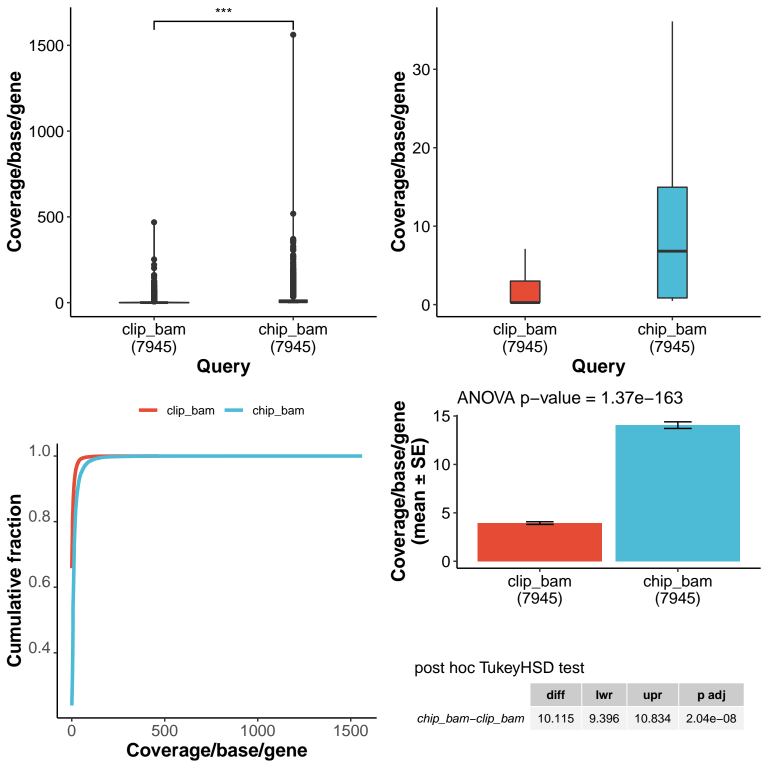
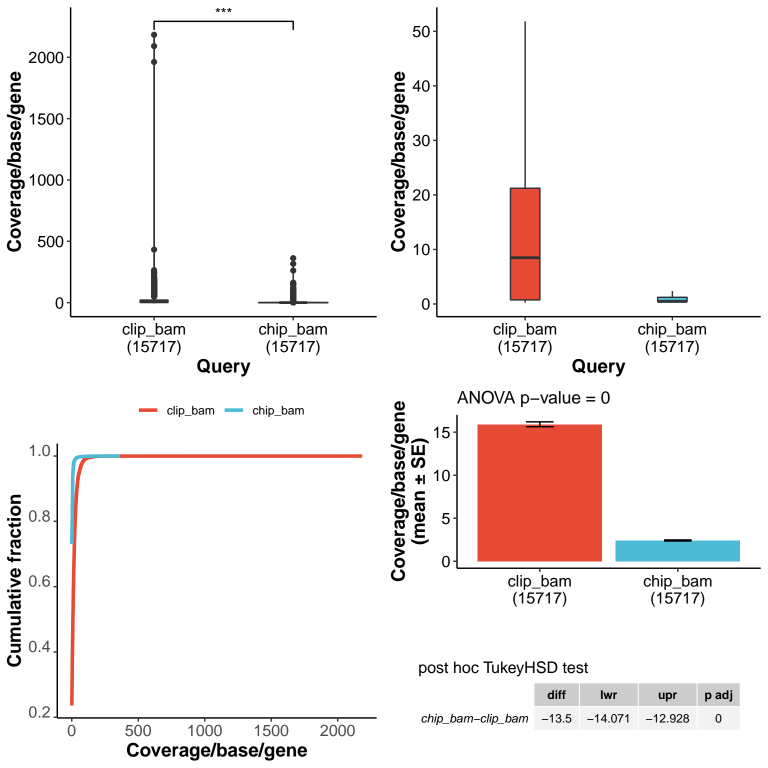


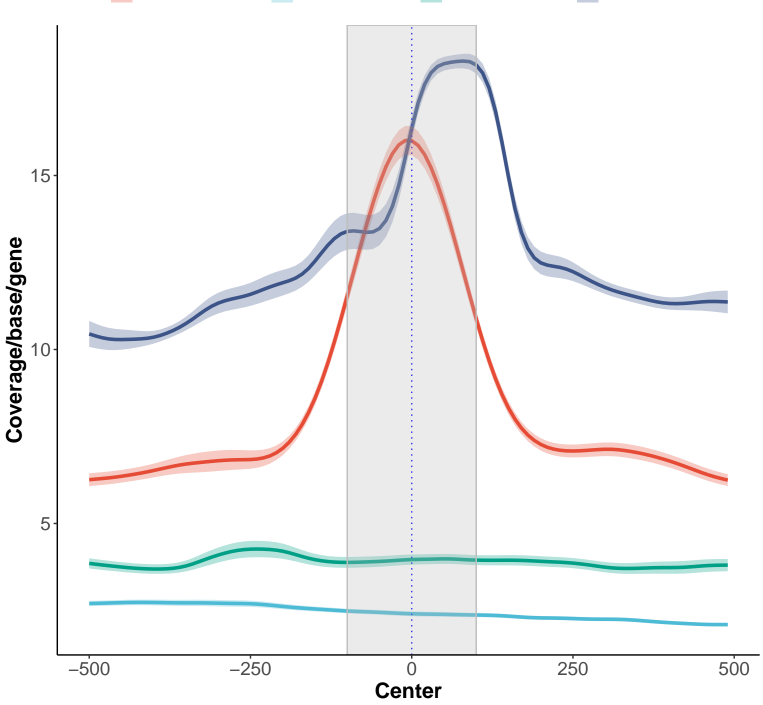
SummitPeak clip_bam chip_bam 16 Coverage/base/gene 4 -250 250 Ö Center -500 500

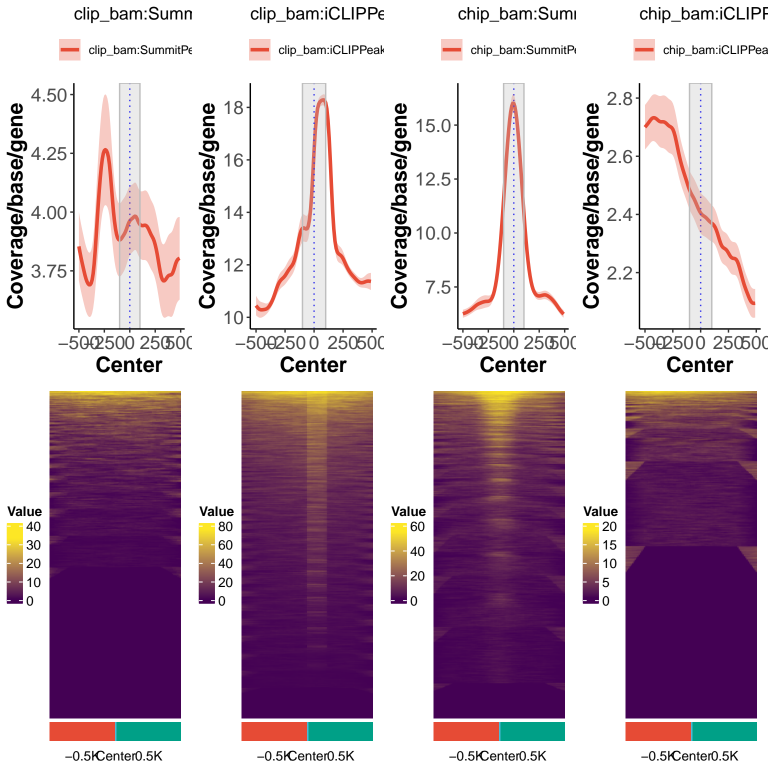


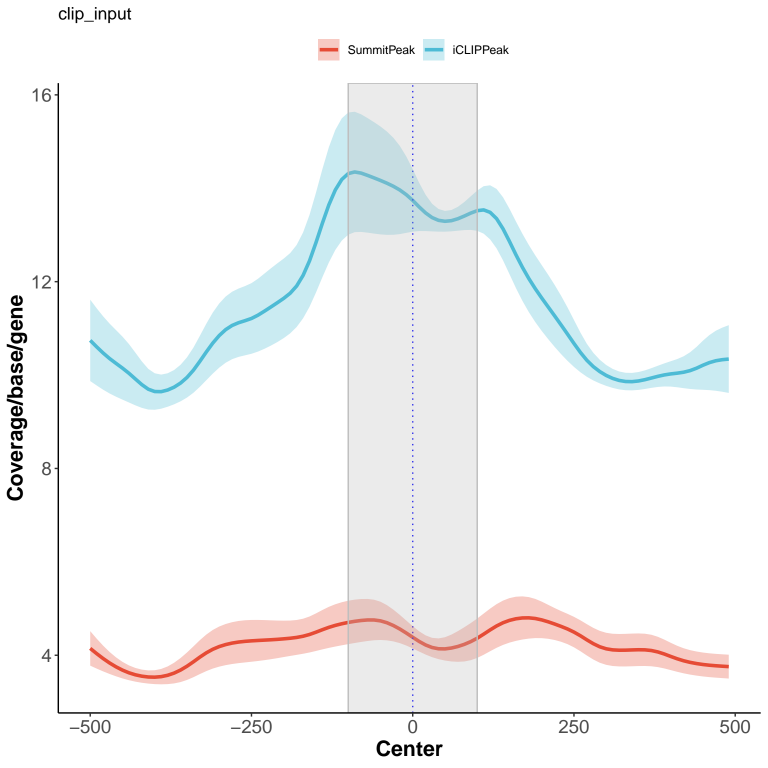
iCLIPPeak clip_bam — chip_bam 15-Coverage/base/gene 5 -250 Center 250 -500 500

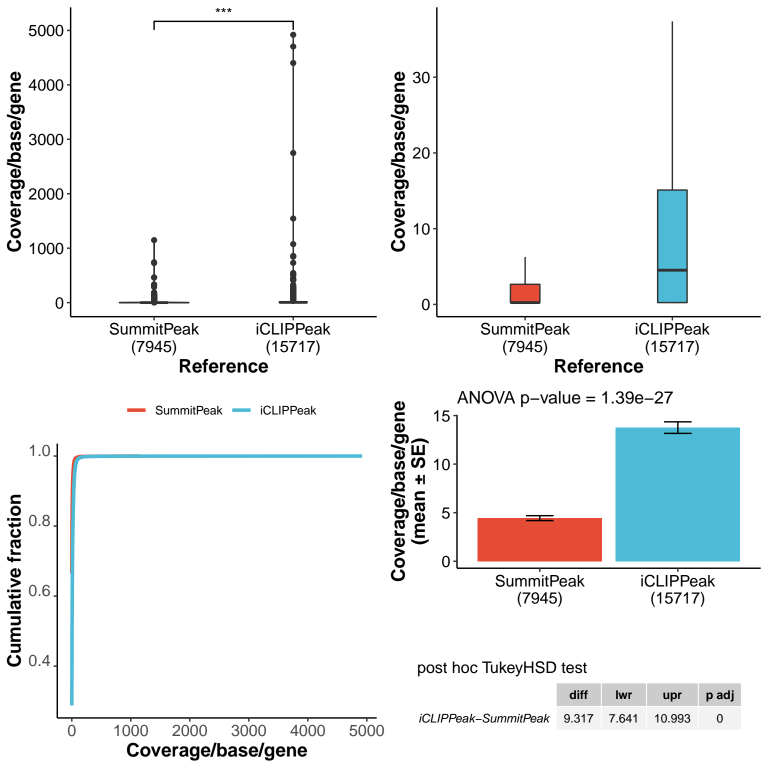


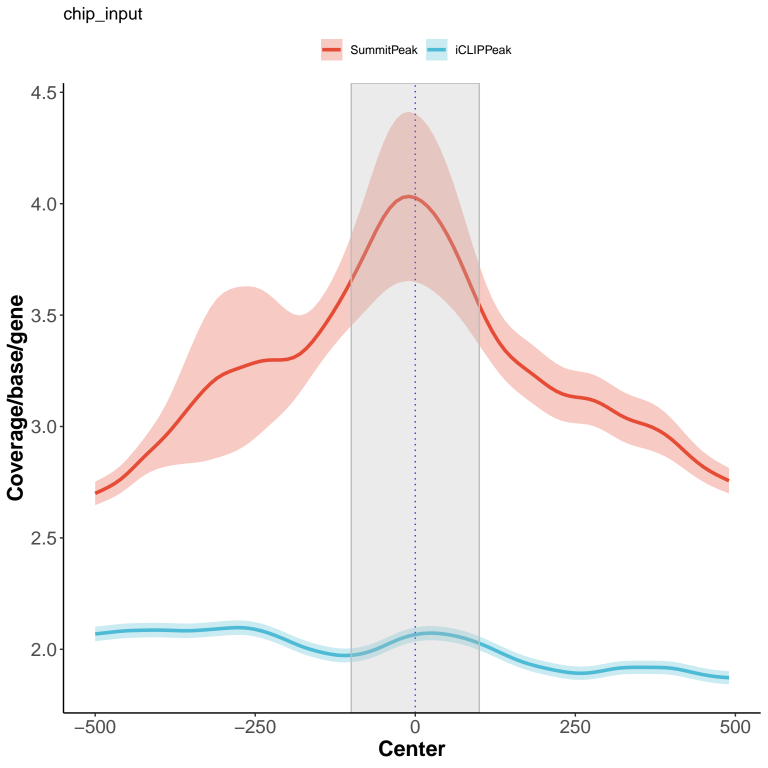
clip_bam:SummitPeak chip_bam:SummitPeak — chip_bam:iCLIPPeak — clip_bam:SummitPeak — clip_bam:iCLIPPeak

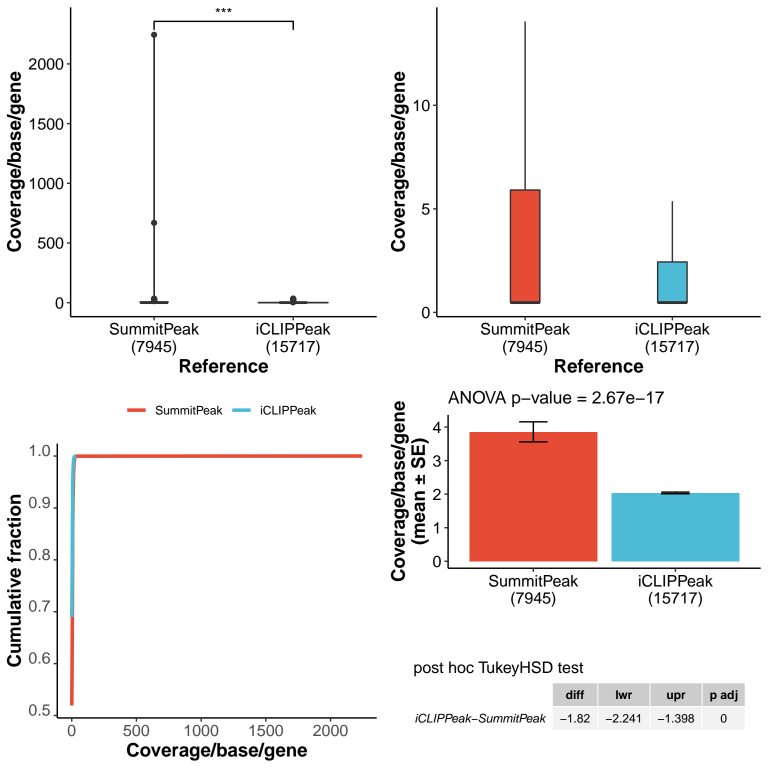




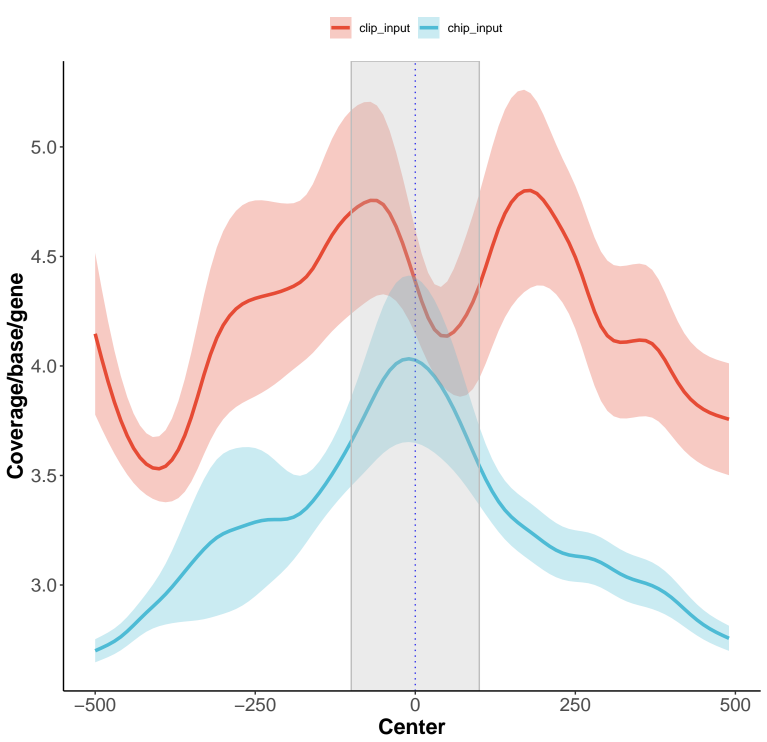


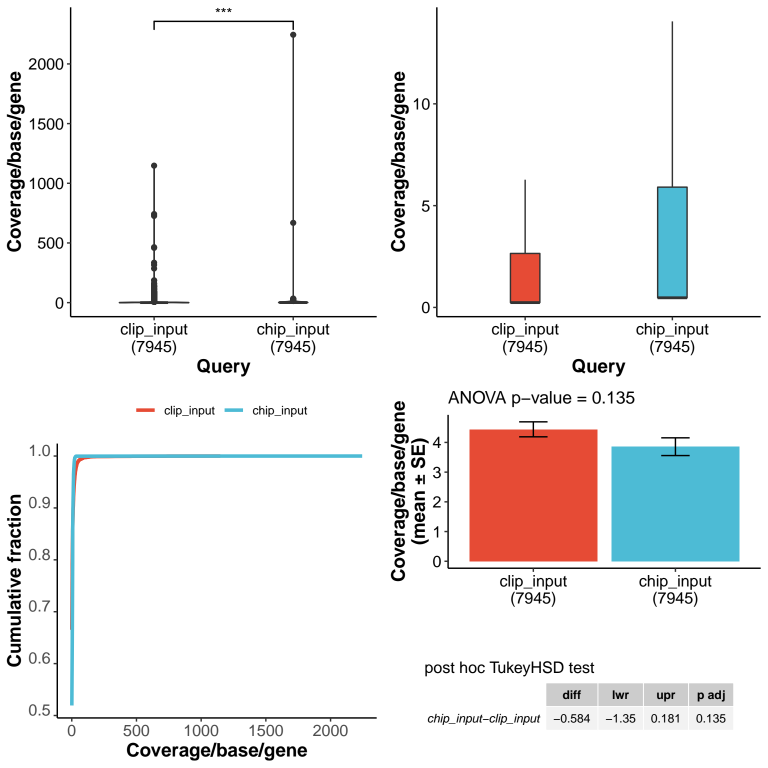


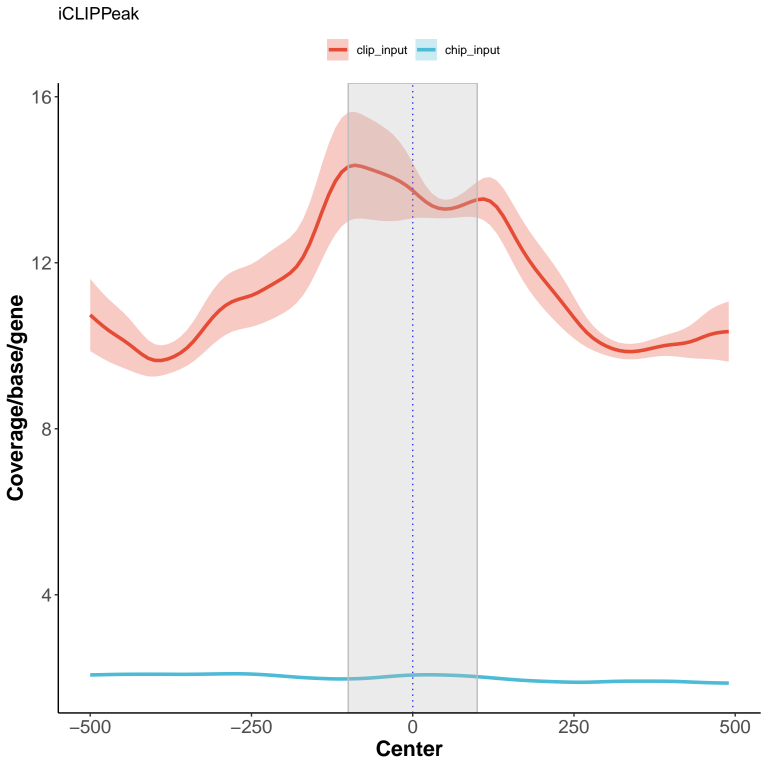


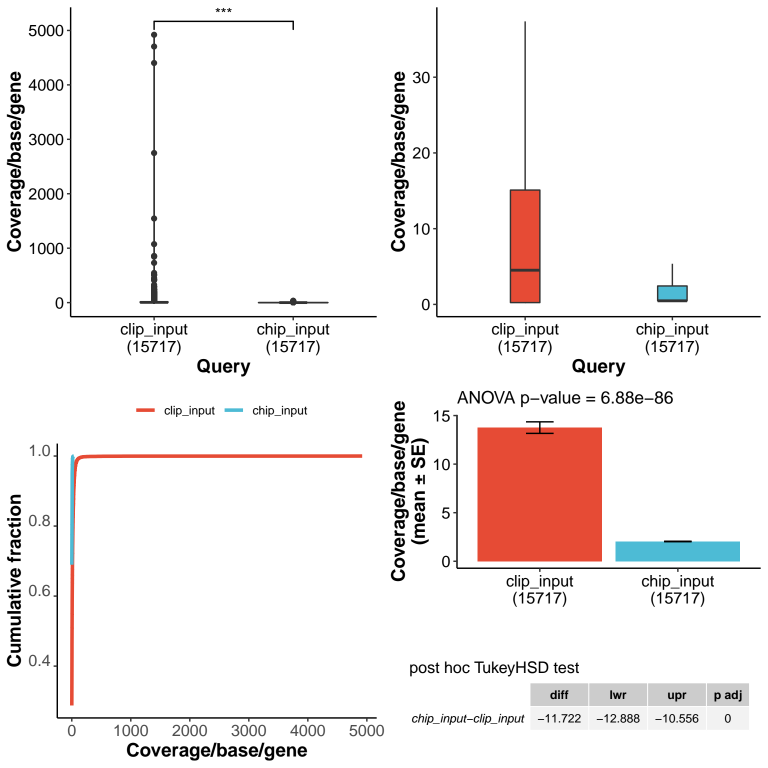


SummitPeak









clip_input:SummitPeak chip_input:SummitPeak chip_input:iCLIPPeak — clip_input:SummitPeak — clip_input:iCLIPPeak 16-12 4

> Ö Center

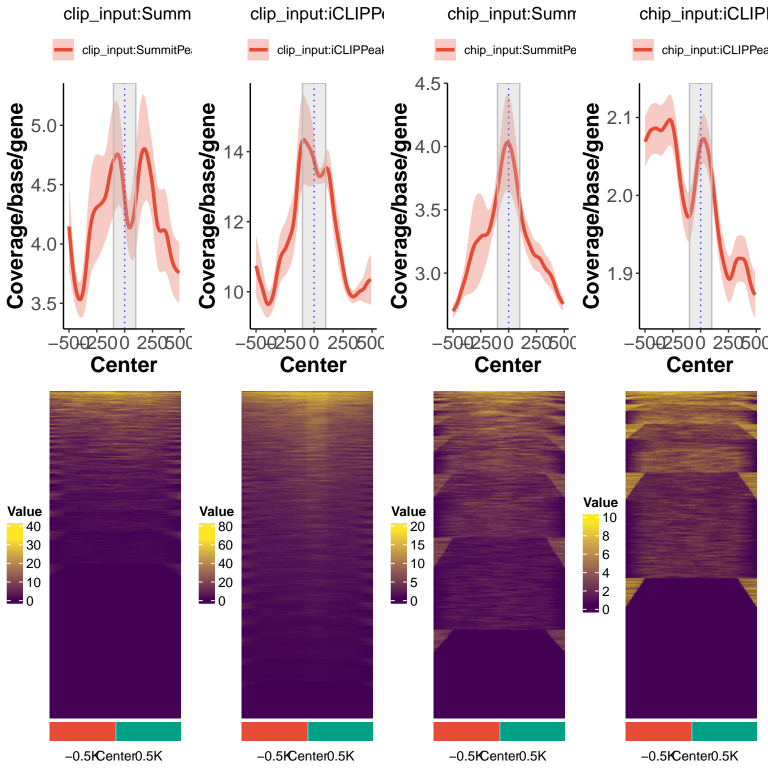
250

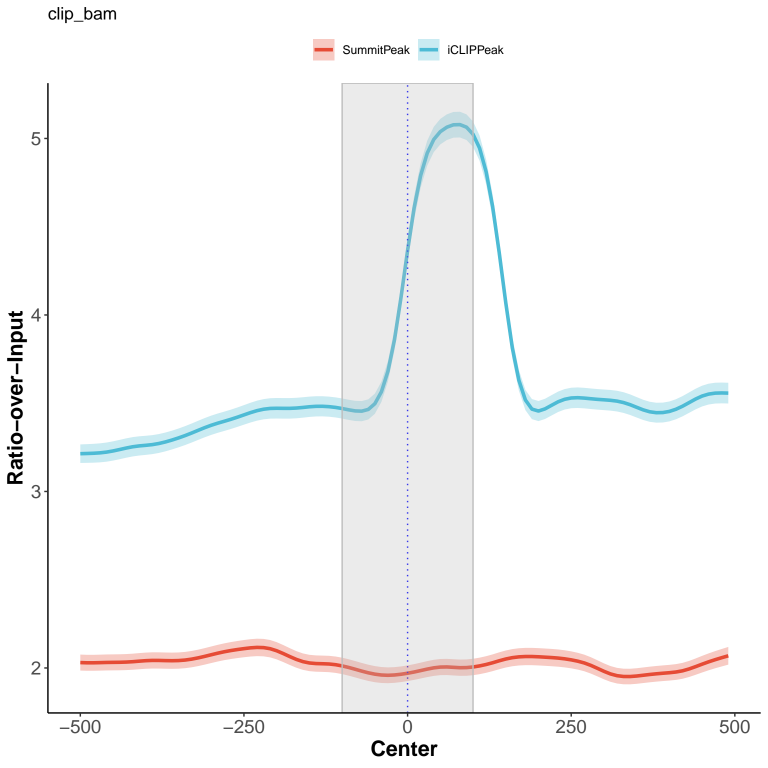
500

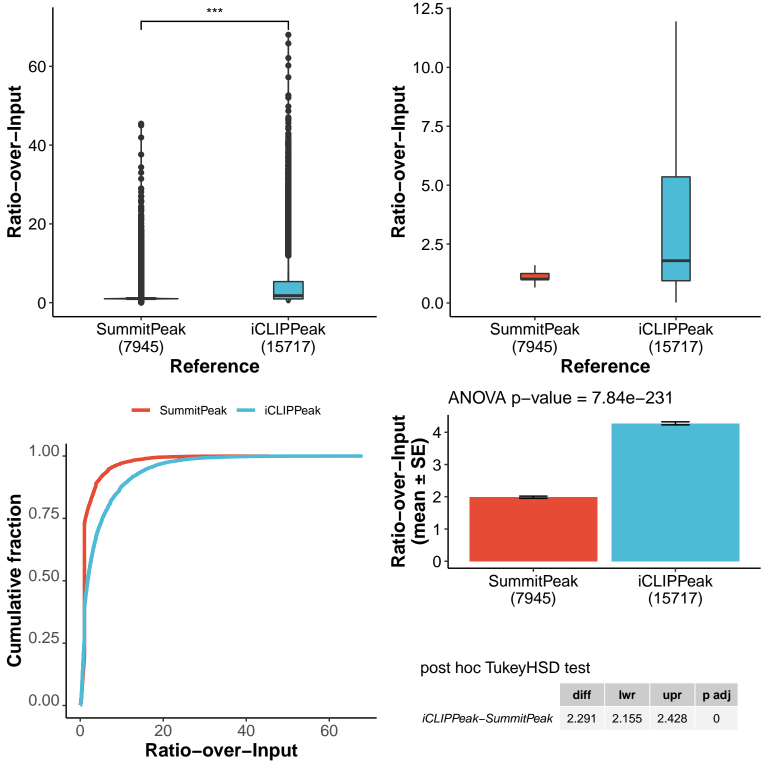
-250

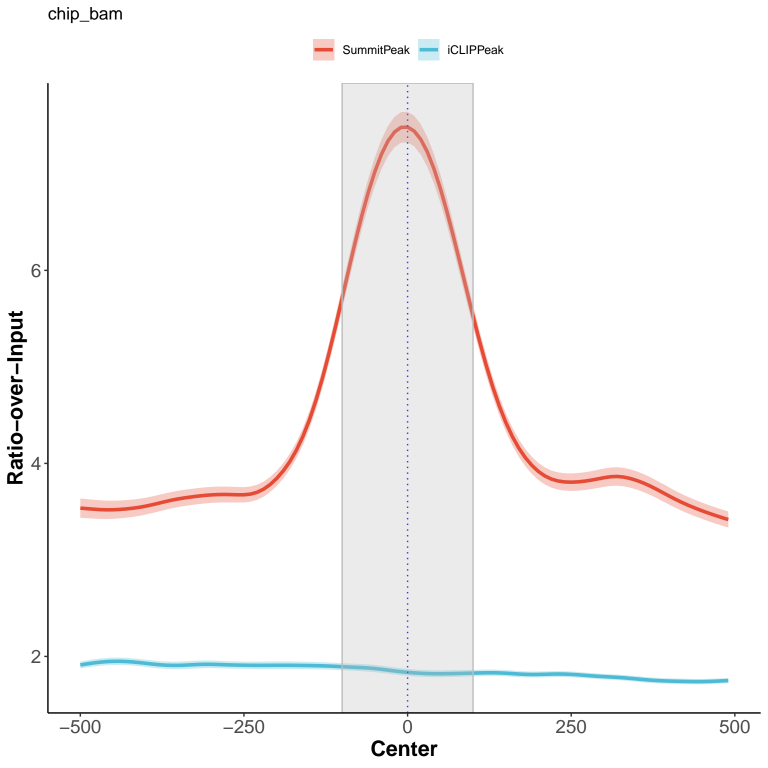
Coverage/base/gene

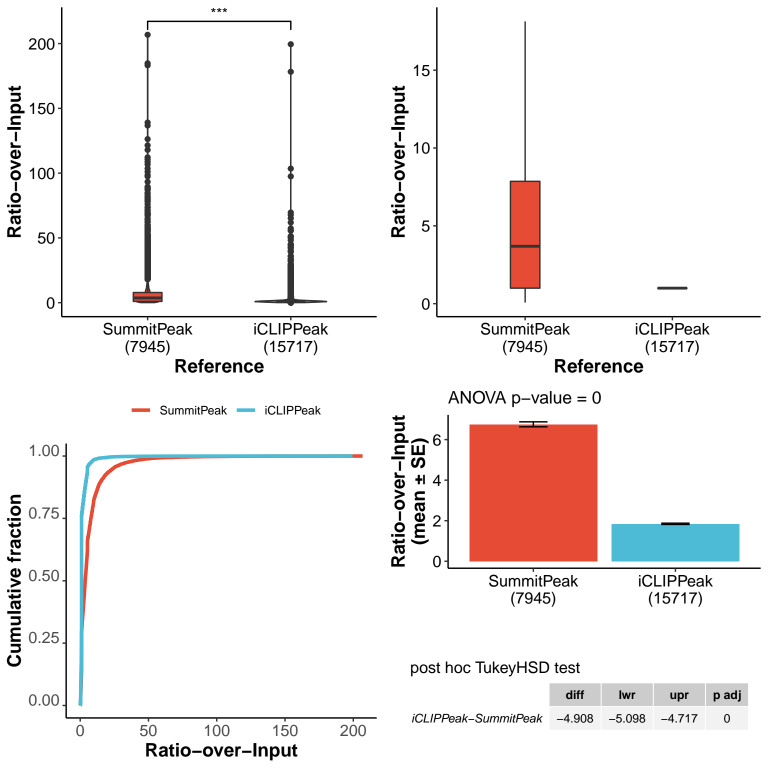
-500



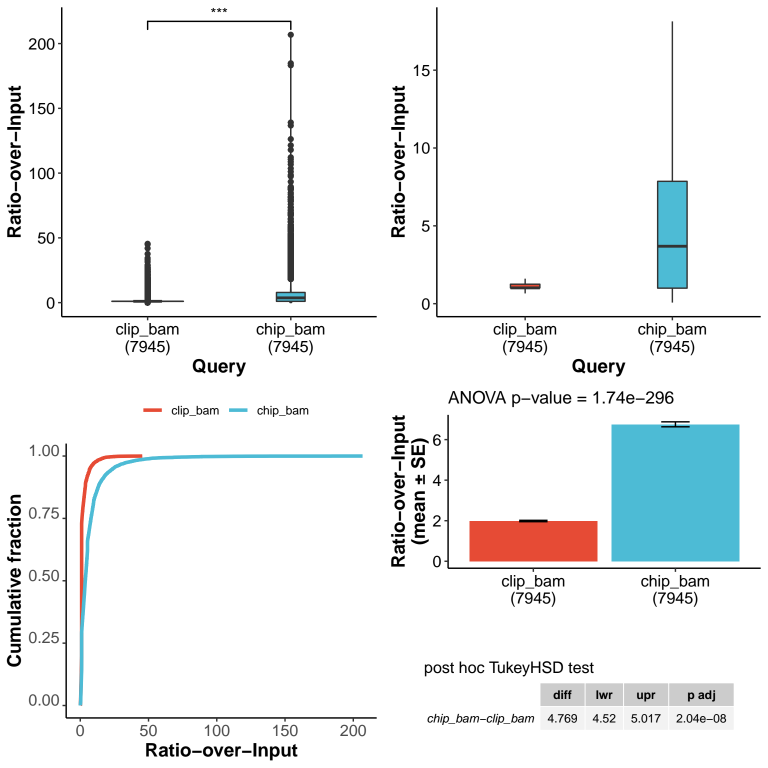


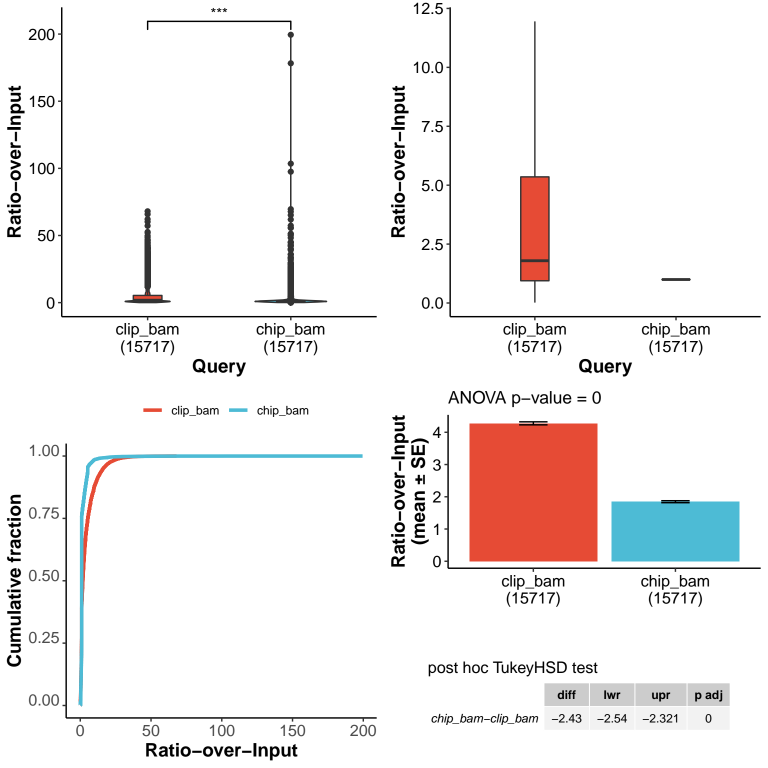




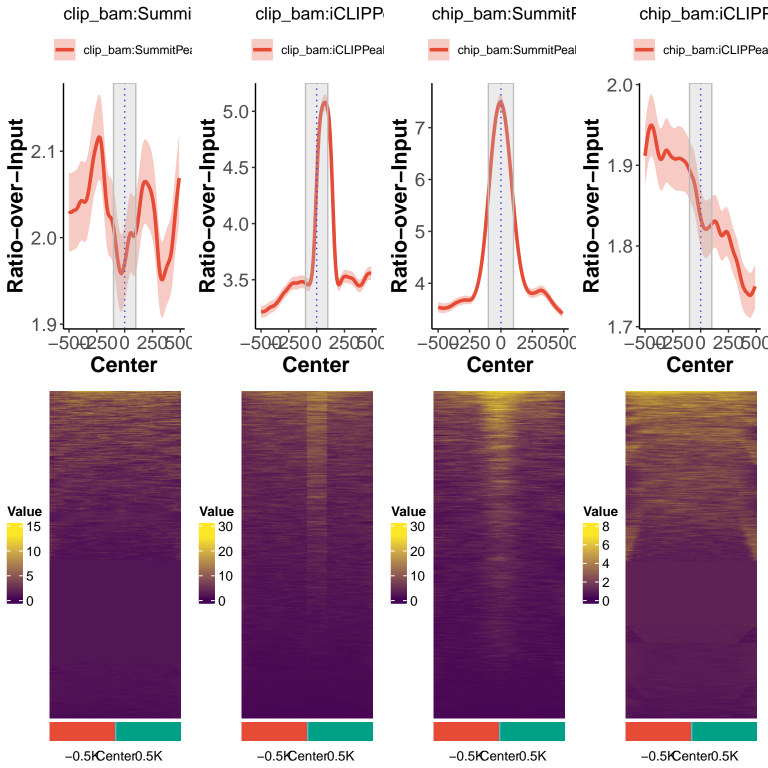


SummitPeak clip_bam chip_bam 6-Ratio-over-Input 2 -250 Center 250 -500 500





clip_bam:SummitPeak chip_bam:SummitPeak chip_bam:iCLIPPeak — clip_bam:SummitPeak clip_bam:iCLIPPeak 6 Ratio-over-Input 2 -250 250 -500 Ö Center 500



Plotting parameters:

```
functionName: "plot_locus"
                               queryFiles: c(clip bam =
  "C:/GREENBLATT/Rscripts/GenomicPlot gh/inst/extdata/treat chr19.bam", chip bam
   = "C:/GREENBLATT/Rscripts/GenomicPlot_gh/inst/extdata/chip_treat_chr19.bam")
                             centerFiles: c(SummitPeak =
  "C:/GREENBLATT/Rscripts/GenomicPlot_gh/inst/extdata/test_chip_peak_chr19.bed",
                                     iCLIPPeak =
  "C:/GREENBLATT/Rscripts/GenomicPlot_gh/inst/extdata/test_clip_peak_chr19.bed")
                                     txdb: NULL
                                  ext: c(-500, 500)
                                   hl: c(-100, 100)
                                    shade: TRUE
                                    smooth: TRUE
importParams: list(offset = 0, fix_width = 150, fix_point = "start", norm = TRUE, useScore =
           FALSE, outRle = TRUE, useSizeFactor = TRUE, genome = "hg19")
                                   verbose: FALSE
                                     binSize: 10
                                   refPoint: "center"
                                    Xlab: "Center"
                             Ylab: "Coverage/base/gene"
                               inputFiles: c(clip_input =
       "C:/GREENBLATT/Rscripts/GenomicPlot_gh/inst/extdata/input_chr19.bam",
                                     chip input =
    "C:/GREENBLATT/Rscripts/GenomicPlot_gh/inst/extdata/chip_input_chr19.bam")
                                   stranded: TRUE
                                   heatmap: TRUE
                                    scale: FALSE
                             outPrefix: "test_plot_locus2"
                                     rmOutlier: 0
                                    transform: NA
                               statsMethod: "wilcox.test"
                                  heatRange: NULL
                                     hw: c(8, 8)
```

nc: 2