



















































Plotting parameters:

```
functionName: "plot_region"
                                            queryFiles: c(clip_bam =
      "C:/Users/greenblatt/AppData/Local/Programs/R/R-4.2.1/library/GenomicPlot/extdata/treat chr19.bam",
                                                  chip bam =
   "C:/Users/greenblatt/AppData/Local/Programs/R/R-4.2.1/library/GenomicPlot/extdata/chip_treat_chr19.bam")
                                          centerFiles: c(NarrowPeak =
:/Users/greenblatt/AppData/Local/Programs/R/R-4.2.1/library/GenomicPlot/extdata/test_chip_peak_chr19.narrowPeak
                                                  txdb: NULL
                                           regionName: "narrowPeak"
                                            inputFiles: c(clip_input =
      "C:/Users/greenblatt/AppData/Local/Programs/R/R-4.2.1/library/GenomicPlot/extdata/input_chr19.bam",
                                                  chip_input =
   "C:/Users/greenblatt/AppData/Local/Programs/R/R-4.2.1/library/GenomicPlot/extdata/chip input chr19.bam")
                                                   nbins: 100
            importParams: list(offset = 0, fix_width = 150, fix_point = "start", norm = TRUE, useScore =
                       FALSE, outRIe = TRUE, useSizeFactor = FALSE, genome = "hg19")
                                                verbose: FALSE
                                                  scale: FALSE
                                                heatmap: TRUE
                                                  fiveP: -500
                                                  threeP: 500
                                                 smooth: TRUE
                                                stranded: TRUE
                                                 transform: NA
                                           outPrefix: "test plot region"
                                                  rmOutlier: 0
                                               heatRange: NULL
                                          Ylab: "Coverage/base/gene"
                                            statsMethod: "wilcox.test"
                                                   hw: c(8, 8)
```

nc: 2