







Plotting parameters:

```
functionName: "plot_3parts_metagene"
                                 queryFiles: c(chip_query =
       "C:/GREENBLATT/Rscripts/GenomicPlot_gh/inst/extdata/chip_treat_chr19.bam")
                       gFeatures: as.list(environment())[["gFeatures"]]
                                 inputFiles: c(chip_input =
       "C:/GREENBLATT/Rscripts/GenomicPlot_gh/inst/extdata/chip_input_chr19.bam")
                                       scale: FALSE
                                      verbose: FALSE
                                Ylab: "Coverage/base/gene"
handleInputParams: list(offset = 0, fix width = 150, fix point = "start", norm = TRUE, useScore =
             FALSE, outRIe = TRUE, useSizeFactor = TRUE, genome = "hg19")
                                      smooth: TRUE
                                      stranded: TRUE
                           outPrefix: "test_plot_3parts_metagene"
                                     heatmap: TRUE
                                     rmOutlier: TRUE
                                     heatRange: NULL
                                       transform: NA
                                        hw: c(8, 8)
                                           nc: 2
```