



Plotting parameters:

```
functionName: "plot_5parts_metagene"
                              queryFiles: c(NarrowPeak =
"C:/GREENBLATT/Rscripts/GenomicPlot_gh/inst/extdata/test_chip_peak_chr19.narrowPeak",
                                     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")
                  gFeatures_list: as.list(environment())[["gFeatures_list"]]
                                    inputFiles: NULL
importParams: list(offset = 0, fix width = 100, fix point = "center", norm = FALSE, useScore
          = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19")
                                    verbose: FALSE
                                     transform: NA
                                     smooth: TRUE
                                     scale: FALSE
                                    stranded: TRUE
                        outPrefix: "test_plot_5parts_metagene1"
                                    heatmap: TRUE
                                   heatRange: NULL
                                      rmOutlier: 0
                              Ylab: "Coverage/base/gene"
                                      hw: c(8, 8)
                                         nc: 2
```