Plotting parameters:

functionName: "plot_bam_correlation"

bamfiles: c(chip_input =

"C:/GREENBLATT/Rscripts/GenomicPlot/inst/extdata/chip_input_chr19.bam", chip_treat =

"C:/GREENBLATT/Rscripts/GenomicPlot/inst/extdata/chip_treat_chr19.bam")

binSize: 1e+05

outPrefix: "test_plot_bam_correlation"

importParams: list(offset = 0, fix_width = 150, fix_point = "start", norm = FALSE, useScore =

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

verbose: FALSE

hw: c(8, 8)

nc: 2