

Figure S2. H3K27ac enrichment density and NCoR1 binding analysis.

(A). Volcano plot showing total number of differentially acetylated regions at 2h and 6h CpG+plC activated DCs compared to unstimulated. (B). Heatmap showing scaled vst of H3K27ac enrichement from ChIP-seq and gene expression from RNA-seq data of plC specific genes in unstimulated, 2h and 6h stimulated condition (C). Boxplot showing distribution of scaled vst H3K27ac enrichmnet on syngergistic, CpG and plC antagoized genes. (D). Scatter plots showing the Super Enhancers (SE) identified at 2hr, and 6hr CpG+plC stimulated cDC1. The CpG/plC specific genes annotated to the SE regions have been marked for each condition. (E). Bar plot showing total number of NCoR1 binding sites identified in unstimulated, 6h CpG, plC and CpG+plC+IFNγ stimulation conditions in cDC1. (F). Percent stacked bar plot showing percent of distribution of NCoR1 binding sites distance relative to TSS in unstimulated, 6h CpG, plC and CpG+plC+IFNγ stimulation conditions in cDC1. (G). Bar plot showing significantly enriched Reactome term from MSigDB for each NCoR1 binding cluster. (H). Heatmap showing odds ratio of overlap of genes associated to each NCoR1 binding cluster and genes classified based on expression as CpG, plC specific and common CpG-plC.