Figure S4. Analysis of **H3K27me3** and **H3K27ac** ChIP-Seq data from ENCODE project. **H3K27me3** (a) and **H3K27ac** (b) data set from human mammary epidermal cells (HMEC) was taken from UCSC Genome Browser and analyzed following the same pipeline used for Std PAT-ChIP and EPAT-ChIP (LRC) data sets. Pie charts depicting the distribution of peaks across genomic features with relative percentage values shown on the right. Promoters are defined as -3 Kb to +3 Kb relative to the TSS while Downstream as -3 Kb relative to the end of 3’ UTR region.