## **Operon Promoter Landscape**

Operon	Strand	Operon start	Operon end
rrsA-ileT-alaT-rrlA-rrfA	+	4033554	4038659

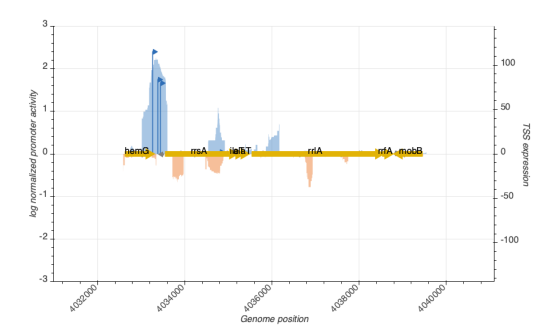


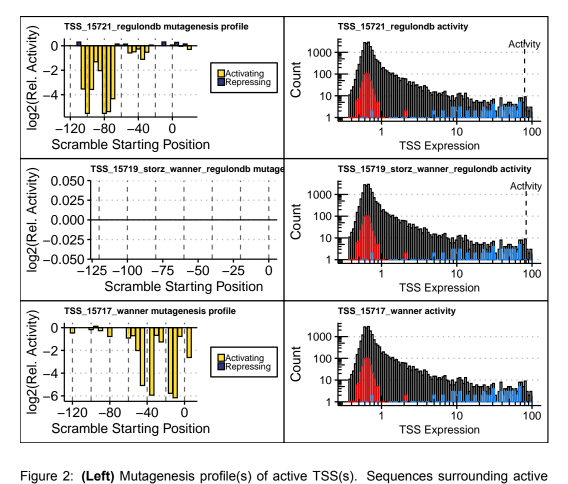


Figure 1: **Promoter activity in rich media (LB) surrounding query operon**. 17,767 previously reported TSSs were evaluated by measuring the promoter activity (right Y-axis) of the 150 bp surrounding the TSS (-120 to +30) to determine which were active or inactive. The genome-wide promoter activity (left Y-axis) was determined by measuring expression of over 300,000 genomic fragments spanning the *E. coli* genome and averaging promoter activity at all nucleotide positions in a strand-specific fashion. Genome coordinates corresponds to *E. coli* genome version U00096.2.

## **TSS Summary**

TSS name	TSS position	Strand	TSS activity	Category
TSS_15721_regulondb	4033439	+	79.6293228	active
TSS_15716_storz	4033204	-	0.6644640	inactive
TSS_15715_storz	4033169	-	0.7520880	inactive
TSS_15719_storz_wanner_regulondb	4033379	+	83.5994409	active
TSS_15722_storz	4033513	-	0.5368769	inactive
TSS_15717_wanner	4033260	+	114.9400357	active

## TSS Scanning Mutagenesis



TSSs were systematically mutated to identify regions controlling expression. Bar height indicates the relative change in promoter activity as a result of scrambling nucleotides within 10 bp regions at 5 bp intervals spanning the promoter. Bar color identifies the region as a putative activator (yellow) or repressor (purple). (**Right**) Dashed line indicates the expression of the indicated TSS relative to all tested TSS sequences. The distributions of expression is shown for all tested TSSs (black), 500 negative controls (red), and a set of constitutive promoters from the BioBrick registry (blue).