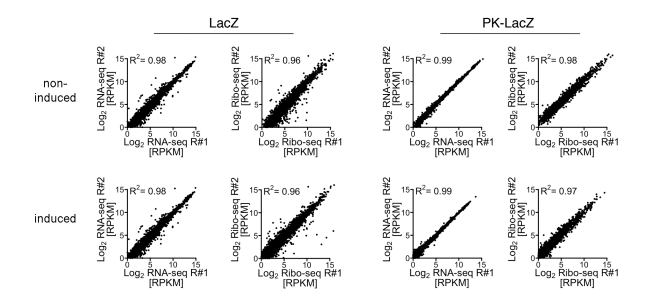
## Supplementary Information for:

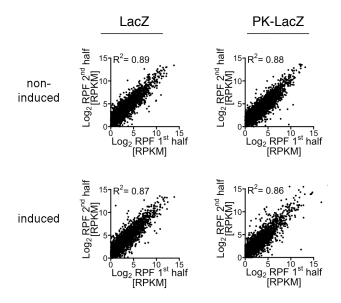
## Absolute quantification of translational regulation and burden using combined sequencing approaches

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**Figure S1: Sequencing data exhibit a good correlation between biological replicates.** Correlation of the RNA-seq and Ribo-seq data of two biological replicates from induced and noninduced cells expressing either LacZ or LacZ-PK. R<sup>2</sup>, Pearson correlation coefficient.



**Figure S2: Comparison of ribosome protected fragments (RPFs) mapping to first and second half of each coding region.** Each point represents a coding region. First 51 bp and last 9 bp not included in the calculation. R<sup>2</sup>, Pearson correlation coefficient.

## Table S1: Measurements of cellular parameters

	Start of induction		End of induction <sup>a</sup>	
Measurement	LacZ	PK-LacZ	LacZ	PK-LacZ
Doubling time (minutes)	85	85	_ <sup>b</sup>	_ <sup>b</sup>
Cell count	1.6 × 10 <sup>9</sup>	1.6 × 10 <sup>9</sup>	1.6 × 10 <sup>9</sup>	1.6 × 10 <sup>9</sup>
Protein mass (grams/cell)	4.13 × 10 <sup>-13</sup>	4.10 × 10 <sup>-13</sup>	4.70 × 10 <sup>-13</sup>	4.30 × 10 <sup>-13</sup>
Mass of LacZ/PK-LacZ as total of cell	-	_	14%	5.8%

a. Induction of constructs lasted 10 min for LacZ and 15 min for PK-LacZ before samples were taken.

b. No measurable growth during the induction period.

Sample <sup>a</sup>			Total # of reads <sup>b</sup>	Multi-mapped reads <sup>c</sup>	Non- coding <sup>c</sup>	Uniquely mapped to mRNA <sup>°</sup>	
PK-LacZ	RNA-seq	non-induced	R#1	66066771	29796556	2876188	33394027
	i vivit-sey		1 1177	00000771	(45.10%)	(4.35%)	(50.55%)
PK-LacZ	RNA-seq	non-induced	R#2	39198111	18025740	1673252	19499119
	NNA-364		I NFL	00100111	(45.99%)	(4.27%)	(49.75%)
PK-LacZ	RNA-seq	induced	R#1	70004102	28545341	1982506	39476255
					(40.78%)	(2.83%)	(56.39%)
PK-LacZ	RNA-seq	induced	R#2	68146152	27782049	1931069	38433034
					(40.77%)	(2.83%)	(56.40%)
PK-LacZ	Ribo-seq	non-induced	R#1	59384989	25736660	3034480	30613849
					(43.34%)	(5.11%)	(51.55%)
PK-LacZ	Ribo-seq	non-induced	R#2	62798001	23567687	6009235	33221079
	Ttibo-seq				(37.53%)	(9.57%)	(52.90%)
PK-LacZ	Ribo-seq	induced	R#1	56568122	27566889	2387216	26614017
FIX-Lacz		Induced			(48.73%)	(4.22%)	(47.05%)
PK-LacZ	Ribo-seq	induced	R#2	54581989	20143633	3541469	30896887
FIX-Lacz					(36.91%)	(6.49%)	(56.61%
LacZ	RNA-seq	non-induced	R#1	192586734	90755510	7758247	94072977
Lacz					(47.12%)	(4.03%)	(48.85%)
LacZ	RNA-seq	non-induced	R#2	53735584	28463851	3382171	21889562
Lauz					(52.97%)	(6.29%)	(40.74%)
LacZ	RNA-seq	induced	R#1	190867334	91396079	6916496	92554759
Lauz	KinA-seq	muuceu	1\#1	190007334	(47.88%)	(3.62%)	(48.49%)
LacZ	RNA-seq	induced	R#2	52249590	25660126	2776669	23812795
Lauz					(49.11%)	(5.31%)	(45.58%)
LacZ	Ribo-seq	non-induced	R#1	183013690	124093781	11347972	47571937
Lacz					(67.81%)	(6.20%)	(25.99%)
LacZ	Ribo-seq	non-induced	R#2	53330842	25579314	4072967	23678561
Lauz					(47.96%)	(7.64%)	(44.40%)
1007	Ribo-seq	induced	R#1	199707963	120352995	7716217	71638751
LacZ					(60.26%)	(3.86%)	(35.87%)
1007	Ribo-seq	induced	R#2	41870343	31443591	1843808	8582944
LacZ					(75.10%)	(4.40%)	(20.50%)

## Table S2: Sequencing and read mapping statistics

a. Biological replicates are denoted "R#1" and "R#2" and "induced" relates to presence of IPTG (1 mM).

b. Reads aligning to more than one sequence including tRNA and rRNA were excluded from the data.

c. Sequencing reads were mapped to the genome of E. coli K-12 MG1655 strain.