

Supporting Information for

High-resolution mapping of Sigma Factor DNA Binding Sequences using Artificial Promoters, RNA aptamers and Deep Sequencing.

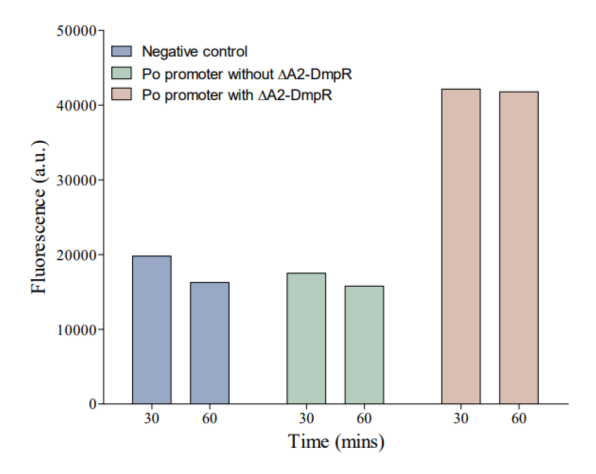
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Fig. S1. Importance of Δ A2-DmpR for *P. putida* σ ⁵⁴-Dependent IVT Reaction

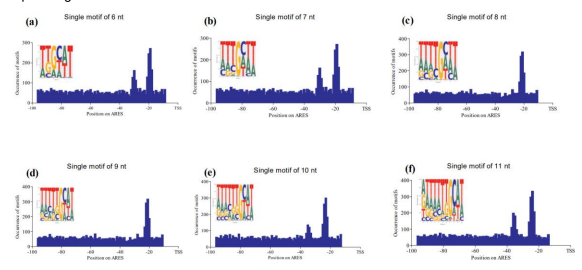


Fluorescence of the Mango aptamer was used to measure RNA synthesis. The results are presented for three experimental groups:

- 1. **Negative Control:** No template added, showing baseline fluorescence.
- 2. **Po Promoter without \triangle A2-DmpR:** Template of the Po promoter without $\triangle A2$ -DmpR, showing fluorescence levels similar to the negative control.
- 3. **Po Promoter with \triangle A2-DmpR:** Template of the Po promoter with $\triangle A2$ -DmpR, showing significantly higher fluorescence levels compared to the negative control.

The data demonstrate that RNA synthesis does not occur in the absence of Δ A2-DmpR, as indicated by the similar fluorescence levels of the negative control and the Po promoter without Δ A2-DmpR. In contrast, significant RNA synthesis, indicated by increased fluorescence, is observed in the presence of both the Po promoter template and Δ A2-DmpR. This highlights the essential role of Δ A2-DmpR in initiating transcription from the σ ⁵⁴ dependent Po promoter.

Fig. S2. Distribution of single motif within a 97 bp Region Upstream of the TSS in the Highest Expressing ARES Subset

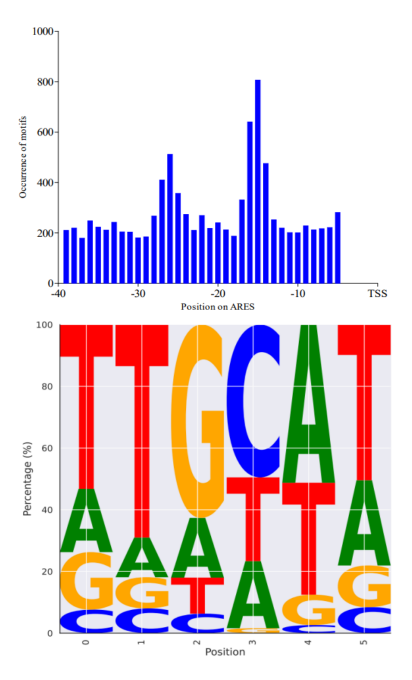


This figure illustrates the distribution of a single conserved motif within a 97 bp region upstream of the Transcription Start Site (TSS) across the highest expressing ARES library for motifs ranging from 6 to 11 nucleotides in length. The x-axis represents the position relative to the TSS, and the y-axis indicates the occurrence frequency of each motif.

- (a) The 6 bp motif (TTGCAT) is identified at the -12 position, consistent with previous core promoter analyses.
- (b) The 7 bp motif shows a sequence extension, adding a T at the beginning (TTTGCTT).
- (c) The 8 bp motif extends further with an additional T (TTTTGCTT).
- (d) The 9 bp motif continues this pattern (TTTTTGCAT).
- (e) The 10 bp motif follows the same trend (TTTTTTGCAT).
- (f) The 11 bp motif (ATTTTTTGCAT) displays the final extension in this analysis.

Each motif is consistently located at the same position upstream of the TSS, with the sequence extension demonstrating high conservation across the lengths studied.

Fig. S3. Distribution of single motif within the 40 bp Region Upstream of the TSS in the Highest Expressing ARES Subset



This figure illustrates the distribution of a single conserved motif within a 40 bp region upstream of the TSS among the highest expressing ARES subset. The x-axis represents the position relative to the TSS, and the y-axis indicates the occurrence frequency of the motif. Analysis reaffirmed the presence of the conserved 6 bp motif (TTGCAT) at the -12 position.

Table S1. Primers used in this study

Primer	Sequence			
name				
	Cloning of double-stranded 200 nt long ARES to pHH100-dBroccoli backbone			
BBa-Prefix- F	GAATTCGCGGCCGCTTCTAGAG			
BBa-Suffix- R	CTGCAGCGGCCGCTACTAGTA			
BB1-F	TCGCACGATATACAGGATTTTGCGGATTTGTCCTACTCAGGAG			
BB1-R	GTCAGCTTTATGCTTGTAAACCGACCGAACAGGCTTATGTC			
BB2-F	CGGTTTACAAGCATAAAGCTG			
BB2-R	GCAAAATCCTGTATATCGTGC			
BB3-F	CGTAATCAAGCCACTTCCTTTT			
BB3-R	GTACGCGTACGGTCTCTGGCATCCGACACCCTGCGTCAATG			
GG-Col-F	GCGGATTTGTCCTACTCA			
GG-Col-R	CTGGATCATCAGAGTATGTG			
For validation	on, 18 ARES, each 200 nt long cloned into the Mango-pSEVA2311 plasmid			
p1-Frag-F	GGCTACGGTCTCCAAGATATCCCGTATAACATTTGTACACTCCGTTATGG			
p1-Frag-R	GGCTACGGTCTCTTGGCAACATTTACATTCAGCAGGCACCAGCGCA			
p1-BB-F	GGCTACGGTCTCTGCCATGTGTATGTGGGCG			
p1-BB-R	GGCTACGGTCTCCTCTTAATTAAAGGCATCAAATAAAACGAAAGG			
p2-Frag-F	GGCTACGGTCTCGACGACTGTCTTAAGGTTAAG			
p2-Frag-R	GGCTACGGTCTCACCACGAGACATATACAATTAC			
p2-BB-F	GGCTACGGTCTCAGTGGATGTAAATGTTGCCATGTG			
p2-BB-R	GGCTACGGTCTCGTCGTTAATTAAAGGCATCAAATAAAACG			
p3-Frag-F	GGCTACGGTCTCCTTCGTGATAATGAGCTGAG			
p3-Frag-R	GGCTACGGTCTCGGGGTAAAATTATATTTAAAGCCG			
p3-BB-F	GGCTACGGTCTCGACCCCTGATGTAAATGTTGCCATGTG			
p3-BB-R	GGCTACGGTCTCCCGAAATTTAATTAAAGGCATCAAATAAAACG			
p4-Frag-F	GGCTACGGTCTCGTAATTGGATTTATTCTACTCTTCC			
p4-Frag-R	GGCTACGGTCTCTCGAAAGAATATCACAATGTAG			
p4-BB-F	GGCTACGGTCTCTTTCGGGGGGTATGTAAATGTTGCCATGTG			
p4-BB-R	GGCTACGGTCTCGATTATAACATTAAATTAAAGGCATCAAATAAAACG			
p5-Frag-F	GGCTACGGTCTCGCCTTTAATTAAGTTGTCCTGATTATGGTAGTACTAGTAG			
p5-Frag-R	GGCTACGGTCTCTTGGCAACATTTACATATGCCCCGCCTCATCATATC			
p5-BB-F	GGCTACGGTCTCTGCCATGTGTATGTGGGCG			
p5-BB-R	GGCTACGGTCTCGAAGGCATCAAATAAAACGAAAGGC			
p6-Frag-F	GGCTACGGTCTCTGAGATTCATTAAATATTATTGGGGAAGAATTGG			
p6-Frag-R	GGCTACGGTCTCATGGACACCACCGTATCCGATG			
p6-BB-F	GGCTACGGTCTCATCCATGTAAATGTTGCCATGTG			
p6-BB-R	GGCTACGGTCTCTCACTTTTAATTAAAGGCATCAAATAAAACG			
p7-Frag-F	GGCTACGGTCTCACCTTTAATTAATAAGCGAGATTCAAGAATTTTTAG			
p7-Frag-R	GGCTACGGTCTCCTAGTATAATTCCATTCCACAG			
p7-BB-F	GGCTACGGTCTCCACTACACGCTGATGTAAATGTTGCCATGTG			
p7-BB-R	GGCTACGGTCTCAAAGGCATCAAATAAAACGAAAG			
p8-Frag-F	GGCTACGGTCTCCAAGGATTTTGAATAAAGTGGTAAAG			
p8-Frag-R	GGCTACGGTCTCACAACATTTACATCCGCTATATGGCTAAATTAG			
p8-BB-F	GGCTACGGTCTCAGTTGCCATGTGTATGTGG			
p8-BB-R	GGCTACGGTCTCCCCTTTAATTAAAGGCATCAAATAAAACG			

p9-Frag-F	GGCTACGGTCTCTGAGTTTACTTCGGGTAAC
p9-Frag-R	GGCTACGGTCTCCCACATGGCAACATTTACATCTACCTATACCACCTAAGAG
p9-BB-F	GGCTACGGTCTCCTGTGTATGTGGGCGTACG
p9-BB-R	GGCTACGGTCTCTACTCCATTTAATTAAAGGCATCAAATAAAACGAAAG
p10-Frag-F	GGCTACGGTCTCAAAAGAGTATGATCCCATTGG
p10-Frag-R	GGCTACGGTCTCGACCATTGATAGGATAGAAAATAAC
p10-BB-F	GGCTACGGTCTCGTGGTTTGAATGTAAATGTTGCCATGTG
p10-BB-R	GGCTACGGTCTCACTTTAATTAAAGGCATCAAATAAAACG
p11-Frag-F	GGCTACGGTCTCTCCTTTAATTAAAATTGTTTTTTTGAATGATATGGTTG
p11-Frag-R	GGCTACGGTCTCTTGATCGATTGCCTATTAAC
p11-BB-F	GGCTACGGTCTCTATCAGATCTTGCGATGTAAATGTTGCCATGTG
p11-BB-R	GGCTACGGTCTCTAAGGCATCAAATAAAACGAAAG
p12-Frag-F	GGCTACGGTCTCAACTAGTATACTATTTCAAAACATATTTG
p12-Frag-R	GGCTACGGTCTCGCAACATTTACATTCCTCATTTTCCATCATC
p12-BB-F	GGCTACGGTTGCCATGTGTATGTGG
p12-BB-R	GGCTACGGTCTCATAGTGTCTTTCTTAATTAAAGGCATCAAATAAAACG
p13-Frag-F	GGCTACGGTCTCACTTAAATAATGTAATACGAGCTCG
p13-Frag-R	GGCTACGGTCTCCATACCTCAGGTTTGC
p13-BB-F	GGCTACGGTCTCGGGAATGTAAATGTTGCCATGTG
p13-BB-R	GGCTACGGTCTCATAAGTTACAACTTAATTAAAGGCATCAAATAAAACG
p14-Frag-F	GGCTACGGTCTCTTACGTAGTTTGTCCAGTATCC
p14-Frag-R	GGCTACGGTCTCCCAACATTTACATCAAAACAGGTGCAGCGGC
p14-BB-F	GGCTACGGTCTCCGTTGCCATGTGTATGTGG
p14-BB-R	GGCTACGGTCTCTCGTAAAATATGAACAATTTTAATTAAAGGCATCAAATAAAACG
p15-Frag-F	GGCTACGGTCTCGCCTTTAATTAATGAACTTATATCTTCTGAACTTTTTTAACGTCTTG
p15-Frag-R	GGCTACGGTCTCGTGGCAACATTTACATGTCGGATACCACCGAGCC
p15-BB-F	GGCTACGGTCTCGGCCATGTGTATGTGGGCG
p15-BB-R	GGCTACGGTCTCGAAGGCATCAAATAAAACGAAAGGC
p16-Frag-F	GGCTACGGTCTCGCAAACTTTAAATGTGCCTCG
p16-Frag-R	GGCTACGGTCTCGACACCCGTCTTGGAATG
p16-BB-F	GGCTACGGTCTCGTGGATGTAAATGTTGCCATGTG
p16-BB-R	GGCTACGGTCTCGTTTGCCCCCGCATCTTAATTAAAGGCATCAAATAAAACG
p17-Frag-F	GGCTACGGTCTCAACCCTCTTTTTGGAATCCTG
p17-Frag-R	GGCTACGGTCTCAATCGGAAGCGAGAAAATCGC
p17-BB-F	GGCTACGGTCTCACGATGTAAATGTTGCCATGTG
p17-BB-R	GGCTACGGTCTCAGGGTATTTTAATTAAAGGCATCAAATAAAACG
p18-Frag-F	GGCTACGGTCTCTTTTCGGCGGATTGATAG
p18-Frag-R	GGCTACGGTCTCTCCCTTGGCTAATGCGAATATAAAAC
p18-BB-F	GGCTACGGTCTCTGGAGATGTAAATGTTGCCATGTG
p18-BB-R	GGCTACGGTCTCTAAGAAATAGTTAAACCTTAATTAAAGGCATCAAATAAAACG
pO54-Frag- F	GGCTACGGTCTCAGCGGCTTAATTTCGCTCG
pO54-Frag- R	GGCTACGGTCTCTCACATGGCAACATTTACATATGTTCATGACTCCATTATTATTGTAC
pO54-BB-F	GGCTACGGTCTCTTGTGTATGTGGGCGTACG
pO54-BB-R	GGCTACGGTCTCACCGCGCTTGCGCAGATTAATTAAAGGCATCAAATAAAACGAAAG
p-Col-F	GTCTTTCGACTGAGCCTTTCG
p-Col-R	CCACATACCAAACCTTCCTTCG
	n of Mango-pSEVA2311 plasmid
Lib1-F	CCTTTAATTAAATGTAAATGTTGCCATGTGTATGTG
Lib2-R	TCCAAGACTAGTACGCGCTACATCCGCTTTAG
Lib3-F	GGATGTAGCGCGTACTAGTCTTGGACTCCTGTTG
Lib4-R	GCAACATTTACATTTAATTAAAGGCATCAAATAAAACGAAAG

Lib7-F	GTCCTACTCAGGAGAGCGTTC
Lib8-R	GCCATGAATGATCCCGAAGG
Sequencing	of ARES plasmid library
SeqDNA	ACACTCTTTCCCTACACGACGCTCTTCCGATCTN 5-7-GACGCAGGGTGTCGGATG
forward	
SeqDNA	GTGACTGGAGTTCAGACGTGTGCTCTTCCGATCTN 5-7-TGGTGCTCGAGCTTGTACAG
reverse	
Sequencing	of in vitro transcribed RNA
SeqRNA	AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCGAG
forward	
SeqRNA1	AGACGTGTGCTCTTCCGATCTCCGACCGTCTCAGATGGACC
reverse	
SeqRNA2	CAAGCAGAAGACGGCATACGAGATGATCTGGTGACTGGAGTTCAGACGTGTGCTCTTCCGATCT
reverse	

Table S2. The sequences and original IDs of the 18 randomly selected ARES, each 200 nt in length, are listed below for the validation experiment.

ID	Original IDs	Sequence
p1	ARES00610543	gatatcccgtataacatttgtacactccgttatggtctgtaataaaatttgggatcgttttttgcgttctaatttacA
'		gatataggctagagtaactactcgcaaacagtgaagatgtgcagtaagggtcaaccggcatagaattgg
		ctcgtgagtacgtcgatctaaagagactggtgtgcgctggtgcctgctga
P2	ARES00171465	cgactgtcttaaggttaagacaatcttattgtgaggtactggggtattggatgcagcatgggtacatttggact
		acattatggatgggggttctgctacagtgtctacttgggacaatatttgcatcactataatgtGcaaaaaga
		aaagtcgggcccttgtttagaaggtatggtcatgtaattgtatatgtctcgtgg
P3	ARES00389971	atttcgtgataatgagctgagttccttgataaagcagtttgtgtaatgaatg
		attgacggctccttataacactccatatgattttgggaacggtgtatgcacaacgctgaggttGgaagtaat
		ggtgattatgatagttaggtggaataaacggctttaaatataattttacccctg
P4	ARES00141878	tgttataattggatttattctactcttcctcataaccgggcttgggaacgaatctagcatcaaacttggcGgag
	7 12000	agatgtcgtttaggagtatgcatatatttctggaaatttcgggatttttgttaaggaatttccagtaggtgttgttc
		gtcaggaatgaaactatagcagctacattgtgatattctttcgggggt
P5	ARES00013532	gttgtcctgattatggtagtactagtagagacaacgtcatcctataaatcgtccaatatataggtgttttggga
' "	71112000010002	actgtcgaattgtgcgtacctatGatcgtcttttccgtaactaagacaagtcagcatattgggggtcgcatgt
		aacgattatcgcacagaagtgagtagtctaggtgatatgatgaggcggggcat
P6	ARES00256315	aagtgagattcattaaatattattggggaagaattggcttgcgagtggaaaaaatgactgatattaggataa
' "	71112000200010	tcttttttgggatcaggtttagcatcagagttagatGagggcaagcaggtgttgattggaggtgcggaacag
		tactataactaccatgcgctgcttgtgggttttagacatcggatacggtggtgtcc
P7	ARES00413161	taagcgagattcaagaatttttagctttatcgtcgtggtgctgtcatgtagtaagta
' '	71112000410101	ctgttgggagtcggggtagttaaagagttgtGaatgatcaaaggtatgtctatgactcagtccatagatcgg
		gaatagtgtggtcttccacctgtcactgtggaatggaat
P8	ARES00523631	aggattttgaataaagtggtaaagttgttgttgctagtgtctgaagatggtatgtttcgtaacgtctgcattgcca
' 0	AINEGUOSESCOT	aaataagtttgagcttatttactttcgggtgaatttgggacaataatttactggttcataaattGggggtaagta
		gttacaatatcagaagtacgttggatgtactaatttagccatatagcgg
P9	ARES00074902	aattgttcatattttacgtagtttgtgtccagtatcccagatacagaccgattagttttgcacgatctgtaaggag
' "	7111200074002	ggatgggattttgcatgccggtacgcttttgatttaaatGggacgaagctggtgcagcgattcttgatcaagt
		atagggagagttagggggtatgtatgtgtactagccgctgcacctgttttg
P10	ARES00116280	gatgcgggggcaaactttaaatgtgcctcgtgaacacgcggttgaacaattgaaatgtgagaggtgaggt
' '	71112000110200	tgtttagtgtcattggaaatgattttcgattaaactgggcgttagtggatGcgtcaaatttgtaaaactaaggtg
		ttgtcgacaatcgtgtcaggaaagctgccttgtaggcattccaagacggtgttgg
P11	ARES00237232	gaaagacactagtatactatttcaaaacatatttgtggaatgaaattgcttatgccgtgatctacagtgaagg
	7 1200020 . 202	gcggcgttctttccggaaagctgttggggacaccagattgatt
		ggtactaaattataacacttaccttgctagctagcggatgatggaaaatgagga
P12	ARES00287266	agagtatgatcccattggttctatagggaatgaaacatttgttaacatacagtggtatgctgatttttgcacggt
		ctgtgcatgtaggccgtctGtctgtaatgctgtgtgaaacaagaatcgtcgagtgcggttaatagggggctt
		gttttcgctttccgatcgcgagaatgttattttctatcctatcaatggtttga
P13	ARES00086654	tgaacttatatcttctgaactttttaacgtcttgttggttaccattagattaaaatagctatagaagcggtagatc
	7 123333333	gtacgttcctcctaagagatgtcaaatatgtattcattacagtggattttgatactttgttgaaagttcgaaaaG
		gtattaagatggttggttagggttgttaaggctcggtggtatccgac
P14	ARES00422720	gttgtaacttaaataatgtaatacgagctcggtagtatctttagcagtatttatcacttggttgg
	7 12000 1221 20	aatgcgaaagaacaggggggaacatgatttttttccaggaggtcacgccaagattgtaaGagggctga
		gagagtcggaaagtgtatacatttaaaggatgtccccgcaaacctgaggtatgga
P15	ARES00952673	ggtttaactatttcttttcggcggattgatagtttaacatctcgtgtgttgatggccaaggtgttagaaggatcgt
		gtcttgagggccgtggattgttctgatgatttttggcaggaaagttaacaaaaagcaaGgatcttatagtatg
		cattatgcttatggtttatattcgcattagccaagggag
P16	ARES00173427	aataccctctttttggaatcctgccatgatcggactttgtctaggtgggagcttttttcaaacgattttagagtaat
		gatgacgtactttttgtatGggctcaattgtgtgtttaggacctttcagccatagtattctcaggctattacagac
		gagtacacttcattgcataactgtaaagggcgattttctcgcttccg
P17	ARES00432513	aattgtttttttgaatgatatggttgcattcatatttacatttacatccatatttgtgtatagcgattttgtaatatggta
		gtctatgatataaatactGattgatattgccgaaacgaggtaaaaaaacatttggtgagccattacatcttttg
		acagatttatcgaagatggttaataggcaatcgatcagatcttgcg
L	1	

P18	ARES00438390	atggagtttacttcgggtaacattggttatatttgagatttgacacatgaccgcactagacttcactacgtgggt ttaaacgtttactggttattttcagattttgtatgaaagttgaatatagggataggGacgctagggctggagtg
		ccgcagcctgataaagccgatacggggatatctcttaggtggtataggtag
pO ⁵⁴	Pos-Control	atgagetetetgegeaagegeggettaatttegetegete

Table S3: Annotation of IVT-generated σ^{54} -RNA polymerase binding motifs in the upstream regions of genes in the *P. putida* genome.

Gene	Product	-24 motifs	-12 motifs	p-value
PP_2783	3-oxoacyl-ACP reductase family protein	TGGGCT	TGGCAT	0,00405
PP_4970	cytochrome c	GTGGCT	GTGTTT	0,00353
PP_1666	DUF2066 domain-containing protein	GTGGTT	GTGTAT	0,003
eutC	ethanolamine ammonia-lyase subunit EutC	TTGGCA	TTGCAT	0,000286
PP_1786	glycosyltransferase	ATGGCA	TTGCAT	0,000286
PP_1030	hypothetical protein	TGGGCT	TTGATT	0,000918
PP_1810	hypothetical protein	TTGGCT	TTGTAT	0,00126
PP_1923	hypothetical protein	ATGGCT	TGGCAT	0,00405
PP_0091	lipoprotein	CTGGCA	TGGCTT	0,00462
PP_1622	peptidoglycan DD-metalloendopeptidase family protein	TTGGTT	TTGCAA	0,00329
PP_4204	type II toxin-antitoxin system MqsA family antitoxin	TTGGCA	GTGTAT	0,003
PP_1394	5-guanidino-2-oxopentanoate decarboxylase	ATGGCT	GTGTTT	0,00353
PP_4577	5-oxoprolinase subunit PxpA	TGGGTA	GTGTTT	0,00353
PP_0859	amidohydrolase	CTGGTA	GTGCTT	0,0015
PP_3521	DMT family transporter	GTGGTT	TTGTAT	0,00126
PP_1210	Dps family protein	TCGGCA	TTGAGT	0,00381
PP_1584	DUF2514 domain-containing protein	TGGGCA	GTGCAT	0,00174
PP_3248	Dyp-type peroxidase	TTGATA	TTGCAA	0,00329
motA	flagellar motor stator protein MotA	TTGGTT	TGGCAT	0,00405
purT	formate-dependent phosphoribosylglycinamide formyltransferase	GTGGCT	TGGCAT	0,00405
pgi	glucose-6-phosphate isomerase	TTGATA	TGGCTT	0,00462
PP_4543	GNAT family N-acetyltransferase	TTGTTA	TTGAGT	0,00381
PP_1159	hypothetical protein	TTGGCT	TTGCAT	0,000286
PP_5488	hypothetical protein	ATGGCT	GTGTAT	0,003
PP_0182	hypothetical protein	ATGGCA	TTGCAT	0,000286
PP_2952	LysR family transcriptional regulator	GTGGTT	TGGCAT	0,00405
PP_3132	oligosaccharide flippase family protein	TGGGCA	TAGCAT	0,00381
PP_5043	PhoPQ-activated pathogenicity-related family protein	TTGGCT	TTGCTT	0,000286
edd	phosphogluconate dehydratase	GTGGCA	TTGTTT	0,00208
PP_4924	S8 family serine peptidase	GTGGCA	TTGTTT	0,00208
PP_0146	TerC family protein	CTGGTA	GTGCAT	0,00174
PP_2420	TonB-dependent receptor	ATGGTT	TTGCGT	0,00267
yegQ	tRNA 5-hydroxyuridine modification protein YegQ	ATGGCT	GTGATT	0,00271
dusA	tRNA dihydrouridine(20/20a) synthase DusA	GTGGTA	GTGAAT	0,00248
tssG	type VI secretion system baseplate subunit TssG	TCGGCA	GTGTTT	0,00353

FIgA flgB flagellar basal body rod protein FIgB TTGGCA TTGCTA 0,0041 flgF flagellar basal-body rod protein FIgF TTGGTT TTGCTT 0,000286 flgG flagellar basal-body rod protein FIgG ATGGCT TTGCAA 0,00329	PP_2433	antitoxin Xre/MbcA/ParS toxin-binding domain-	CTGGCA	TGGCTT	0,00462
PP_1188 dicarboxylate/amino acid:cation symporter TTGCA TTGCTA 0,0041 PP_3989 DNA cytosine methyltransferase ATGGTA TGGCAT 0,00405 PP_2810 DUF1329 domain-containing protein CTGGCA TTGCGT 0,00267 PP_3633 DUF3617 domain-containing protein GTGGTA ATGCAT 0,00491 fligA flagellar basal body P-ring formation chaperone TCGCA TTGCTT 0,000286 fligA flagellar basal body rod protein FlgB TTGGCA TTGCTA 0,0041 fligF flagellar basal-body rod protein FlgG ATGGCT TTGCAA 0,00329 fligE flagellar basal-body rod protein FlgG ATGGCT TTGCAA 0,00329 fligE flagellar bosk-basal body complex protein Flig CTGGCA TTGCAA 0,00329 fligE flagellar bosk-basal body complex protein Flg CTGGCA TTGCAA 0,00329 fligE flagellar basal-body rod protein FlgG ATGGCT TTGCAA 0,00329 FP_2836 fumarylacetoacetate hydrolase family protein GTGGCT TTGCAA <	cls		TCGGCA	ATGCAT	0.00491
PP_3889 DNA cytosine methyltransferase ATGGTA TGGCAT 0,00405 PP_2810 DUF1329 domain-containing protein CTGGCA TTGCGT 0,00267 PP_5363 DUF3617 domain-containing protein GTGGTA ATGCAT 0,00491 flgA flagellar basal body P-ring formation chaperone TCGCA TTGCTT 0,000286 flgB flagellar basal-body rod protein FlgB TTGGTA TTGCTA 0,0041 flgF flagellar basal-body rod protein FlgF TTGGTT TTGCTA 0,00286 fliE flagellar hook-basal body complex protein FliE CTGGCA TTGCTA 0,00329 fliE flagellar hook-basal body complex protein FliE CTGGCA TTGCAA 0,00329 fliE flagellar hook-basal body complex protein FliE CTGGCA TTGCAA 0,00329 fliE flagellar hook-basal body complex protein FliE CTGGCA TTGCAA 0,00329 fliE flagellar hook-basal body complex protein Flig CTGGCA TTGCAA 0,00329 fliE flagellar basal-body robe protein FlgG ATGCCT TTGCAA					•
PP_2810 DUF1329 domain-containing protein CTGGCA TTGCGT 0,00267 PP_5363 DUF3617 domain-containing protein GTGGTA ATGCAT 0,00491 flgA flagellar basal body P-ring formation chaperone FlgA TCGGCA TTGCTT 0,00286 flgB flagellar basal body rod protein FlgB TTGGCA TTGCTA 0,0041 flgF flagellar basal-body rod protein FlgF TTGGTT TTGCAA 0,00329 fliE flagellar hook-basal body complex protein FliE CTGGCA TTGCAA 0,00329 FP_2836 fumarylacetoacetate hydrolase family protein GTGGCT TTGCAA 0,00329 FP_1946 glucose 1-dehydrogenase TTGGCA TTGCAA 0,00329 FP_10486 glucose 1-dehydrogenase TTGGCA TTGCAA 0,00329 PP_10498 GlxA family transcriptional regulator TAGGCA TTGCAA 0,00329 PP_2828 hypothetical protein TAGGTA GTGCAT 0,00174 PP_3592 MurR/RpiR family transcriptional regulator ATGGCA TTGCAA 0,00329	_				, ,
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PP_2209	PP_3288	universal stress protein	TCGGTA	TTGAGT	0,00381
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PP_5722 type II toxin-antitoxin system PemK/MazF CTGGTA TTGAAT 0,000631 family toxin	PP_3778	pyrroline-5-carboxylate reductase	TTGACA	TTGCTA	0,0041
family toxin	PP_1450	HlpA activation/secretion protein HlpB	GTGGCT	TAGCAT	0,00381
xdhA xanthine dehydrogenase small subunit TGGGCA GTGCAT 0,00174	PP_5722	family toxin	CTGGTA	TTGAAT	0,000631
	xdhA	xanthine dehydrogenase small subunit	TGGGCA	GTGCAT	0,00174

Plasmid maps

1) N200 ARES

 $\underline{https://benchling.com/s/seq-wG6faFrPGndrWYAhEMqq?m=slm-wg6jqKmiUQcNzv7zXjAx}$

2) pHH100-dBroccoli construct

https://benchling.com/s/seq-bD8GNHJIVsiaienqUajK?m=slm-NNHWA6VcrijY2MuG6UKe

3) N200-pHH100-dBroccoli construct

https://benchling.com/s/seq-VxVk2xeLeUp684jSTSbb?m=slm-LhFMu9GY5rYQlaGJzR3s

4) Mango-pSEVA2311 construct

 $\underline{https://benchling.com/s/seq-CfaFXRxx76D39VVFn3LS?m=slm-Q88cA84fxmZQlkUefZ72}$