

Current Biology, Volume 32

Supplemental Information

**A dicentric bacterial chromosome requires
XerC/D site-specific recombinases for resolution**

Qin Liao, Zhongqing Ren, Emma E. Wiesler, Clay Fuqua, and Xindan Wang

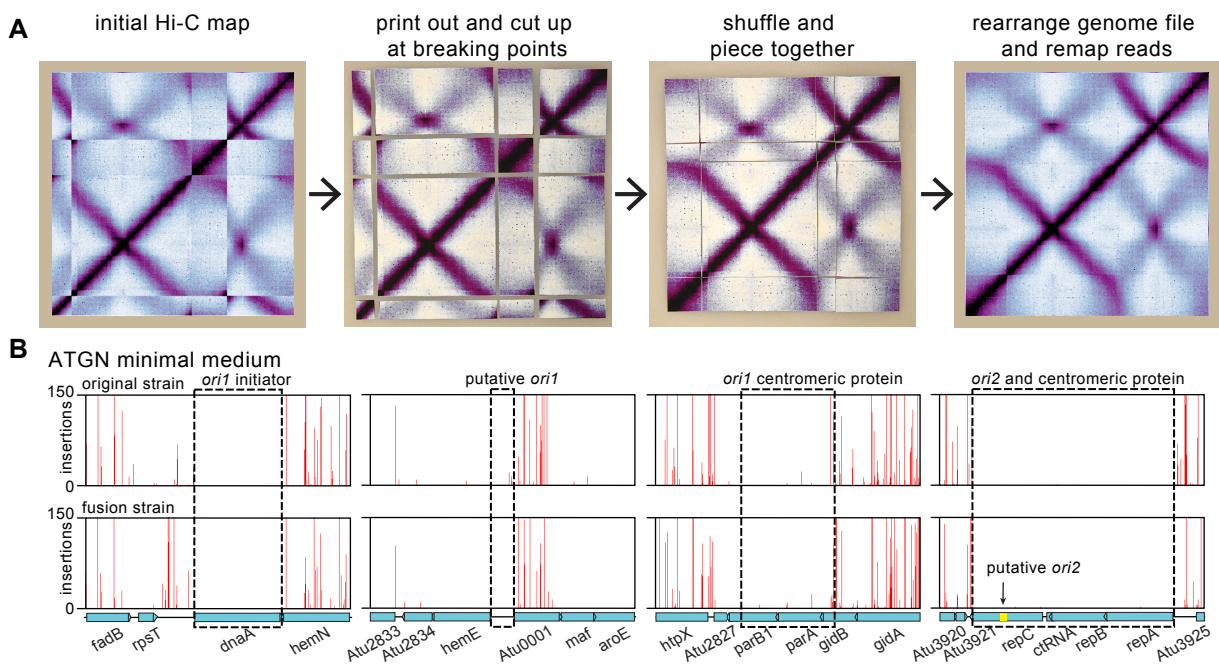


Figure S1. *A. tumefaciens* fusion strain has a dicentric linear chromosome. Related to Figure 1 and Figure 2.

(A) Deciphering the genome rearrangement. The Hi-C map generated by mapping Hi-C reads from *A. tumefaciens* C58 fusion strain to the original C58 reference genome (first panel) was printed out, cut up at breakpoints (second panel), and reassembled to generate a confluent picture (third panel). The pattern indicated that the circular Ch1 and linear Ch2 were fused into a large linear chromosome. When the same Hi-C data were mapped to the rearranged reference sequence, the same pattern was seen (fourth panel).

(B) The initiator proteins, replication origin sequences, and partitioning systems of both *ori1* and *ori2* are required for the survival of the original and fusion strains. Tn-seq plots showing transposon intolerance at indicated regions in the original strain (top) and the fusion strain (bottom). This screen was carried out in minimal ATGN medium.

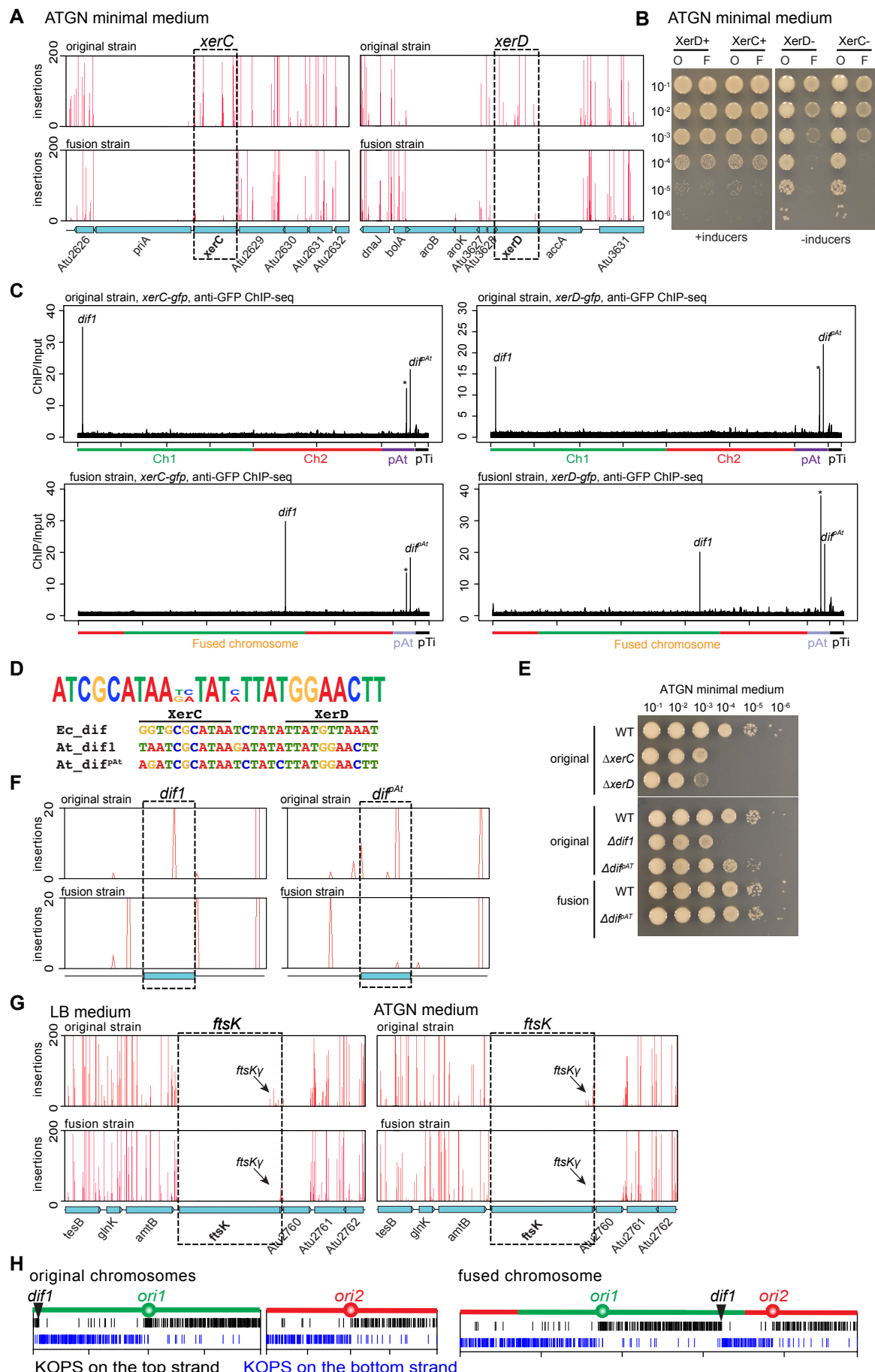


Figure S2. Binding sites of XerC and XerD in the original and fusion strains. Related to Figure 3.

- (A) Tn-seq plots showing transposon insertions at *xerC/xerD* loci in the WT original strain (top) and the WT fusion strain (bottom) growing on ATGN plates.
- (B) Validation of essentiality of *xerD* and *xerC* using depletion constructs, from left to right: AtWX332, AtWX327, AtWX331 and AtWX323. 10-fold serial dilutions of indicated strains were spotted on ATGN plates. The depletion constructs had the endogenous promotor of *xerD* or *xerC* replaced by *Plac-riboswitch*. 2 mM theophylline and 0.5 mM IPTG were added as inducers.
- (C) ChIP enrichment of XerC-GFP and XerD-GFP in the original and fusion strains: top left, AtWX377 (original *xerC-gfpmut3*); top right, AtWX370 (original *xerD-gfpmut3*); bottom left, AtWX367 (fusion *xerC-gfpmut3*); bottom right, AtWX379 (fusion *xerD-gfpmut3*). The original strains were mapped to the original reference genome with *ori1* in the middle of Ch1, and the fusion strains were mapped to the re-arranged reference genome. x-axis shows genome positions and y-axis indicates ChIP enrichment (ChIP/input) in 1-kb bins. Asterisks indicate a systematic peak we observed for every anti-GFP ChIP-seq experiment.
- (D) Top, logo of XerC/XerD binding sequence extracted from ChIP-seq peaks in (C). Bottom, comparison of *A. tumefaciens dif1* and *dif^{At}* with *E. coli dif* with indicated XerC and XerD binding arms.
- (E) 10-fold serial dilutions of the indicated strains spotted on ATGN plates, from top to bottom: AtWX063, AtWX375, AtWX092, AtWX063, AtWX439, AtWX440, AtWX001 and AtWX441.
- (F) Transposon insertions at *dif1* (left) and *dif^{At}* (right) in the WT original strain (top) and the WT fusion strain (bottom) growing on LB plates.
- (G) Tn-seq plots showing transposon insertions at *ftsK* locus in the WT original strain (top) and the WT fusion strain (bottom) growing on LB plates (left) and ATGN plates (right).
- (H) Distributions of FtsK orienting polar sequences (KOPS) in original chromosomes (left) and the fused chromosome (right). KOPS (GGGNAGGG) on the top (Watson) strand and on the bottom (Crick) strand are shown as black and blue bars, respectively.

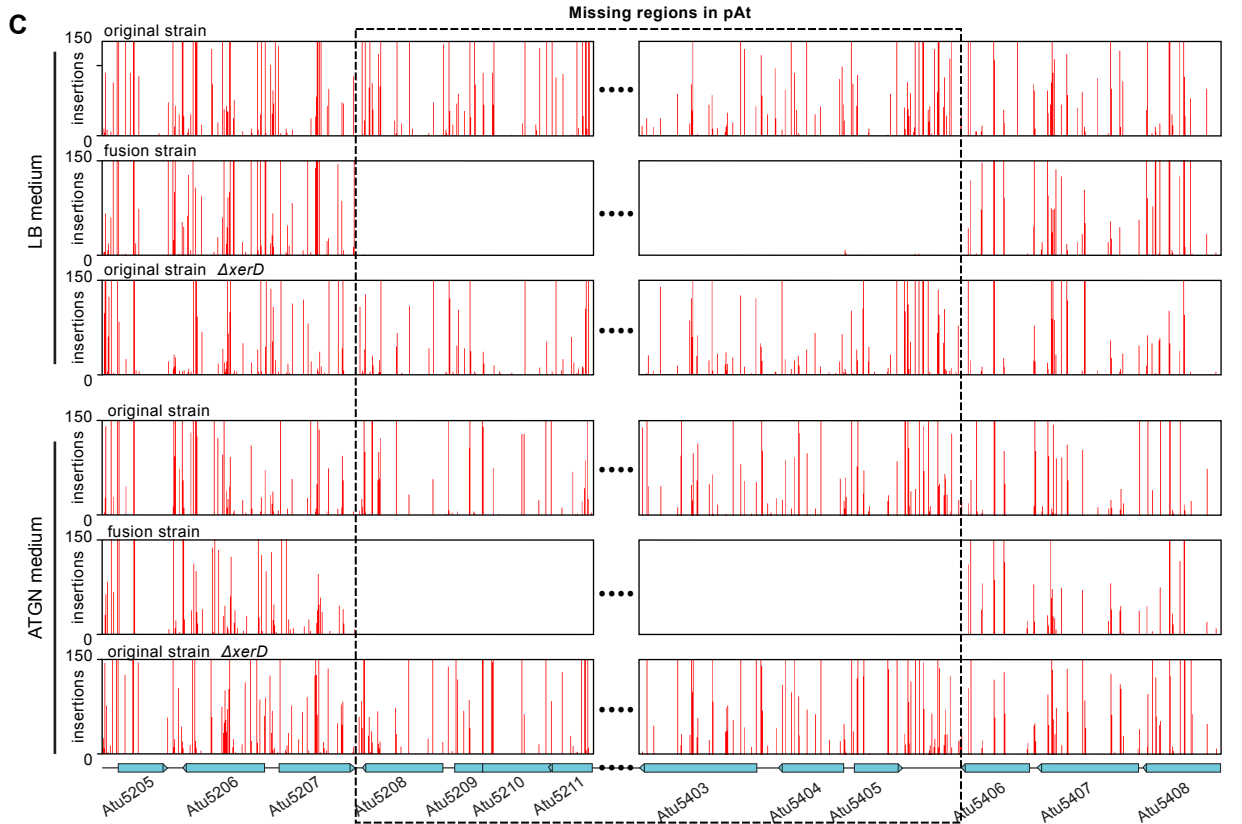
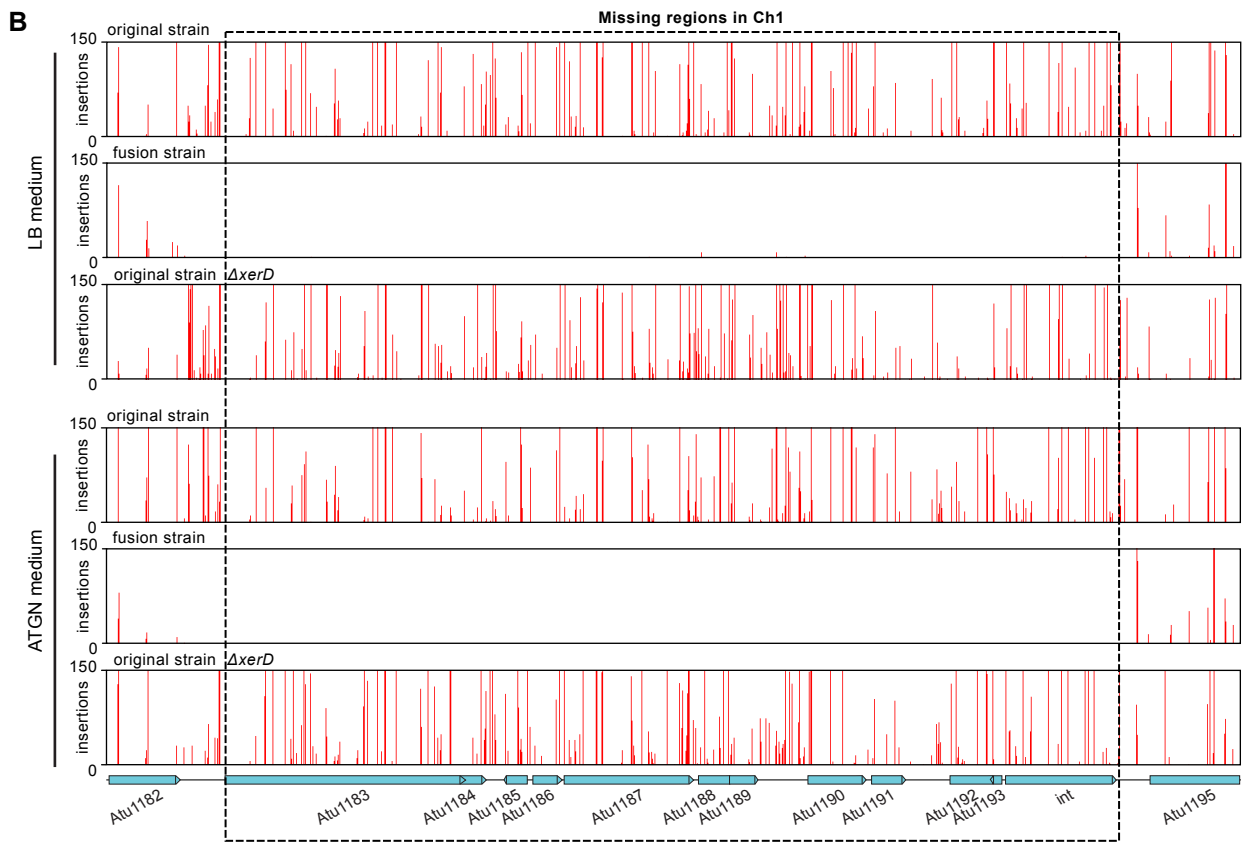
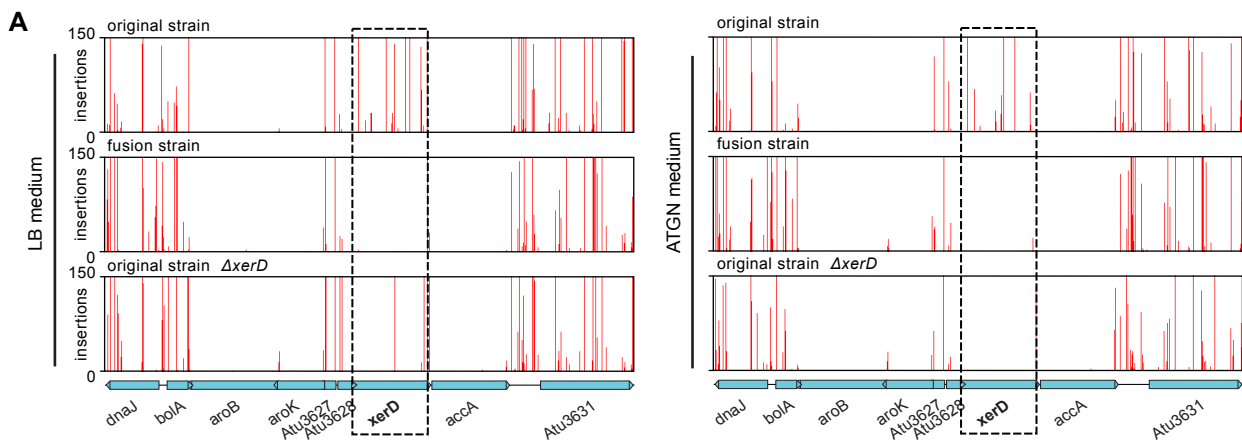
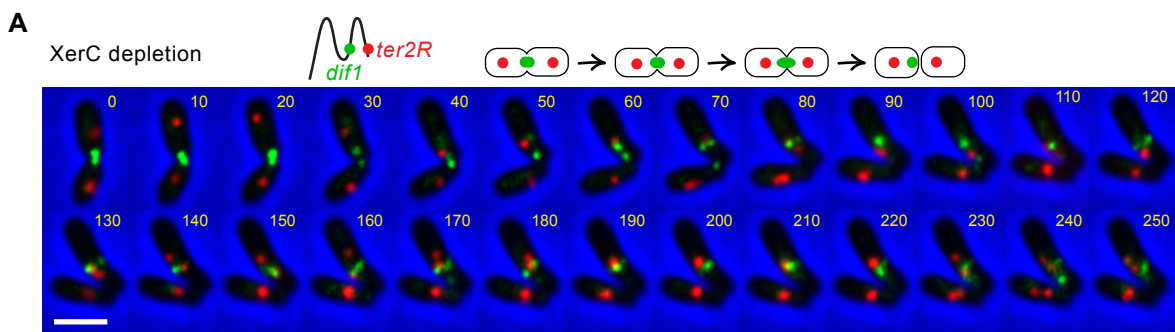


Figure S3. *xerD* is not synthetically lethal with genes that are missing on Ch1 and pAt in the fusion strain. Related to Figure 3. Tn-seq was performed in three different strains: original strain (AtWX063, top), fusion strain (AtWX001, middle), and original $\Delta xerD$ strain (AtWX092, bottom). Cells were growing on LB or ATGN plates as indicated.

(A) Transposon insertions at *xerD* locus. *xerD* was successfully deleted from the original strain as indicated by no insertions at this locus in original $\Delta xerD$ (bottom).

(B) Transposon insertions at the Ch1 region that is missing on the fused chromosome. $\Delta xerD$ cells tolerated insertions at these regions (bottom), suggesting *xerD* is not synthetically lethal with these missing genes.

(C) Transposon insertions at the pAt region that is missing from the fusion strain. $\Delta xerD$ cells tolerated insertions at these regions (bottom), suggesting *xerD* is not synthetically lethal with these missing genes. Only the junctions are shown.



B

cell types	depletion time No. of cells analyzed	WT	XerC depletion					XerD depletion			
		0h 1154 cells	0h 902 cells	3h 1163 cells	6h 1004 cells	12h 828 cells	0h 905 cells	3h 942 cells	6h 980 cells	12h 715 cells	
growing cell with one focus		67.7%	64.5%	65.0%	44.7%	24.0%	53.7%	57.1%	49.0%	20.3%	
growing cell with two foci		4.8%	6.3%	3.6%	4.5%	2.1%	10.5%	7.0%	7.4%	3.5%	
early dividing cell		8.5%	10.9%	11.5%	11.1%	7.0%	14.4%	10.9%	14.4%	4.6%	
deeply constricted cell with one or two foci		12.0%	8.6%	8.1%	22.7%	21.5%	10.4%	13.4%	20.7%	18.3%	
cells without foci		5.9%	7.9%	10.1%	11.9%	38.2%	7.1%	8.9%	6.4%	49.5%	
cell with foci and abnormal cell shape		1.2%	1.8%	1.7%	5.2%	7.2%	4.0%	2.7%	2.0%	3.8%	

Figure S4. Quantitative analysis of *dif1* localization in XerC/D depletion strains. Related to Figure 4.

(A) Time-lapse progression of XerC depletion strain containing *dif1* (green) and *ter2R* (red) on the fused chromosome (AtWX514). *dif1* was visualized as shown in **Figure 4B** whereas *ter2R* was visualized by inserting visualization cassette, *mcherry-parBP1-parSP1*, at 39.8 kb away from *ter2R*.

(B) Quantitative analysis of *dif1* localization patterns in the WT fusion strain (AtWX455), XerC-depleted (AtWX467) and XerD-depleted (AtWX471) fusion strains. Snapshot images of cells growing in LB medium were collected at 0 h, 3 h, 6 h, 12 h upon the depletion. Cells were classified and quantified according to cell cycle and the number of foci. After 6 h depletion, the percentage of deeply constricted cells increased by about two fold. Upon 12 h depletion, the proportion of cells without foci rose by over three fold.

Lab	Isolate name	Configuration	References
L.M. Banta	C58	fusion	Banta Lab collection
P. Brown	C58	fusion	S1
C. Fuqua	C58	fusion	S2
S.K. Farrand	C58	fusion	S3
X. Wang	C58 fusion (AtWX001)	fusion	This study
S.C. Winans	C58	fusion	S4
P. Zambryski	C58	fusion	S5
J.H. Chang	C58 A8	original	Chang Lab collection
S.K. Farrand	C58 C1RS, C58 UIA5, C58 NTL4	original	S6 S7
C. Fuqua	C58	original	S8
S.B. Gelvin	C58	original	Gelvin Lab collection
B. Goodner	C58 BG	original	S9
R. Hallez	C58	original	Hallez Lab collection
C.I. Kado	C58 NT1REB	original	S10
E-M. Lai	C58 A107	original	S11
K.S. Mysore	C58	original	S12
E.W. Nester	C58 A136, C58-3, EN58	original	S13 S14
X. Wang	C58 original (AtWX063)	original	S15
P. Zambryski	C58	original	S16

Table S1. Chromosomal configuration of *A. tumefaciens* strains in different labs. Related to STAR Methods.

Sample name	Figure	Reference	Identifier
401_Wang_HiC_AtWX063_ATGN	1A	S15	GSM5542437
438_Wang_WGS_AtWX063_ATGN_rep2	1A	S15	GSM5689555
500_Wang_HiC_AtWX001_ATGN	1BC, S1A	This study	SAMN27387414
501_Wang_WGS_AtWX001_ATGN	1BC	This study	SAMN27387415
502_Wang_Tnseq_AtWX063_LB	2D, 3A, S2F, S3	This study	SAMN27387416
503_Wang_Tnseq_AtWX001_LB	2D, 3A, S2F, S3	This study	SAMN27387417
504_Wang_Tnseq_AtWX387_LB	3C	This study	SAMN27387418
505_Wang_Tnseq_AtWX398_LB	3C	This study	SAMN27387419
506_Wang_Tnseq_AtWX035_LB	3D	This study	SAMN27387420
507_Wang_Tnseq_AtWX108_LB	3D	This study	SAMN27387421
508_Wang_Tnseq_AtWX063_ATGN	S1B, S2A, S3	This study	SAMN27387422
509_Wang_Tnseq_AtWX001_ATGN	S1B, S2A, S3	This study	SAMN27387423
510_Wang_ChIP_antiGFP_AtWX367_ATGN	S2C	This study	SAMN27387424
511_Wang_input_AtWX367_ATGN	S2C	This study	SAMN27387425
512_Wang_ChIP_antiGFP_AtWX370_ATGN	S2C	This study	SAMN27387426
513_Wang_input_AtWX370_ATGN	S2C	This study	SAMN27387427
514_Wang_ChIP_antiGFP_AtWX377_ATGN	S2C	This study	SAMN27387428
515_Wang_input_AtWX377_ATGN	S2C	This study	SAMN27387429
516_Wang_ChIP_antiGFP_AtWX379_ATGN	S2C	This study	SAMN27387430
517_Wang_input_AtWX379_ATGN	S2C	This study	SAMN27387431
518_Wang_Tnseq_AtWX092_LB	S3	This study	SAMN27387432
519_Wang_Tnseq_AtWX092_ATGN	S3	This study	SAMN27387433

Table S2. Next generation sequencing samples used in this study. Related to STAR Methods.

Strains used in main figures			
Strain	Genotype	Reference	Figure
AtWX001	C58 fusion, wild type	This study	1BCEF, 2BCD, 3A, S1AB, S2AEFGH, S3,
AtWX063	C58 original, wild type	S15	1ADF, 2BCD, 3A, S1B, S2AEFGH, S3
AtWX263	C58 original, carrying pWX970, pSRKKm <i>Plac rfp-repB^{Ch2}</i> (Atu3923/ATU_RS18280) terminator <i>Plac egfp-parB1</i> (Atu2828/ATU_RS13770) terminators	S15	2A
AtWX366	C58 fusion, carrying pWX970, pSRKKm <i>Plac rfp-repB^{Ch2}</i> (Atu3923/ATU_RS18280) terminator <i>Plac egfp-parB1</i> (Atu2828/ATU_RS13770) terminators	This study	2A
AtWX025	C58 fusion, <i>tetRA::a-attTn7 pLac repB^{Ch2} gen, ΔrepB^{Ch2}</i>	This study	2C
AtWX089	C58 original, <i>ΔrepB^{Ch2}</i> (Atu3923/ATU_RS18280)::amp	S15	2C
AtWX192	C58 original, <i>tetRA::gen Ptral-riboswitch-parB1</i> (Atu2828/ATU_RS13770) <i>traR</i>	S15	2C
AtWX194	C58 fusion, <i>tetRA::gen Ptral-riboswitch-parB1</i> (Atu2828/ATU_RS13770) <i>traR</i>	This study	2C
AtWX323	C58 fusion, <i>xerC</i> (Atu2628/ATU_RS12790)::terminator gen <i>LacI</i> terminators <i>Plac riboswitch xerC</i>	This study	3B, S2B
AtWX327	C58 fusion, <i>xerD</i> (Atu3629/ATU_RS16850)::terminator gen <i>lacI</i> terminators <i>Plac riboswitch xerD</i>	This study	3B, S2B
AtWX331	C58 original, <i>xerC</i> (Atu2628/ATU_RS12790)::terminator gen <i>LacI</i> terminators <i>Plac riboswitch xerC</i>	This study	3B, S2B
AtWX332	C58 original, <i>xerD</i> (Atu3629/ATU_RS16850)::terminator gen <i>lacI</i> terminators <i>Plac riboswitch xerD</i>	This study	3B, S2B
AtWX387	C58 fusion, <i>ΔrecA</i> (Atu1873/ATU_RS09160)::amp	This study	3C
AtWX398	C58 original, <i>ΔrecA</i> (Atu1873/ATU_RS09160)::amp	This study	3C
AtWX035	C58 fusion, <i>Δsmc</i> (Atu0801/ATU_RS03945)	This study	3D
AtWX108	C58 original, <i>Δsmc</i> (Atu0801/ATU_RS03945)	S15	3D
AtWX502	C58 fusion, <i>xerC</i> (Atu2628/ATU_RS12790)::terminator gen <i>LacI</i> terminators <i>Plac riboswitch xerC</i> , carrying pWX1076, pSRKKm <i>cymR cuO Plac cuO At parC</i> (Atu1158/ATU_RS05720) <i>At parE</i> (Atu1622/ATU_RS07965)	This study	3E
AtWX503	C58 fusion, <i>xerD</i> (Atu3629/ATU_RS16850)::terminator gen <i>lacI</i> terminators <i>Plac riboswitch xerD</i> , carrying pWX1076, pSRKKm <i>cymR cuO Plac cuO At parC</i> (Atu1158/ATU_RS05720) <i>At parE</i> (Atu1622/ATU_RS07965)	This study	3E
AtWX540	C58 fusion, <i>xerC</i> (Atu2628/ATU_RS12790)::terminator gen <i>LacI</i>	This study	3E

	terminators <i>Plac riboswitch xerC</i> , carrying pWX1080, pSRKKm <i>cymR cuO Plac cuO Ec parE-parC</i>		
AtWX541	C58 fusion, <i>xerD</i> (Atu3629/ATU_RS16850)::terminator gen <i>lacI</i> terminators <i>Plac riboswitch xerD</i> , carrying pWX1080, pSRKKm <i>cymR cuO Plac cuO Ec parE-parC</i>	This study	3E
AtWX455	C58 fusion, <i>ygfp-parB^{MT1}-parS^{MT1}</i> inserted between Atu1460/ATU_RS07195 and Atu1461/ATU_RS07200, 22 kb from <i>dif1</i>	This study	4B, S4B
AtWX467	C58 fusion, <i>xerC</i> (Atu2628/ATU_RS12790)::terminator gen <i>LacI</i> terminators <i>Plac riboswitch xerC</i> ; <i>ygfp-parB^{MT1}-parS^{MT1}</i> inserted between Atu1460/ATU_RS07195 and Atu1461/ATU_RS07200, 22 kb from <i>dif1</i>	This study	4CD, S4B
AtWX471	C58 fusion, <i>xerD</i> (Atu3629/ATU_RS16850)::terminator gen <i>lacI</i> terminators <i>Plac riboswitch xerD</i> ; <i>ygfp-parB^{MT1}-parS^{MT1}</i> inserted between Atu1460/ATU_RS07195 and Atu1461/ATU_RS07200, 22 kb from <i>dif1</i>	This study	4D, S4B

Strains used for strain building or in supplemental figures

Strain	Genotype	Reference	Figure
AtWX367	C58 fusion, <i>xerC-gfpmut3</i> (Atu2628/ATU_RS12790)	This study	S2C
AtWX370	C58 original, <i>xerD-gfpmut3</i> (Atu3629/ATU_RS16850)	This study	S2C
AtWX377	C58 original, <i>xerC-gfpmut3</i> (Atu2628/ATU_RS12790)	This study	S2C
AtWX379	C58 fusion, <i>xerD-gfpmut3</i> (Atu3629/ATU_RS16850)	This study	S2C
AtWX439	C58 original, Δ <i>dif1::amp</i>	This study	S2E
AtWX440	C58 original, Δ <i>dif^{At}1::amp</i>	This study	S2E
AtWX441	C58 fusion, Δ <i>dif1::amp</i>	This study	S2E
AtWX092	C58 original, Δ <i>xerD</i> (Atu3629/ATU_RS16850)::amp	This study	S2E, S3
AtWX375	C58 original, Δ <i>xerC</i> (Atu2628/ATU_RS12790)::amp	This study	S2E
AtWX514	C58 fusion, <i>xerC</i> (Atu2628/ATU_RS12790)::terminator gen <i>LacI</i> terminators <i>Plac riboswitch xerC</i> ; <i>ygfp-parB^{MT1}-parS^{MT1}</i> inserted between Atu1460/ATU_RS07195 and Atu1461 22 kb from <i>dif1</i> ; <i>mCherry-parB^{P1}-parS^{P1}</i> inserted between Atu4854 and Atu4856 39.8 kb from <i>ter2R</i>	This study	S4A

Plasmids used in this study

Plasmid	Description	Reference	
pWX811	pNPTS138 Δ <i>repB^{Ch2}</i> (Atu3923/ATU_RS18280)::amp	This study	
pWX813	pMiniTn7 <i>pLac At repB^{Ch2}</i> (Atu3923/ATU_RS18280)	This study	
pWX832	pNPTS138 Δ <i>smc</i> (Atu0801/ATU_RS03945)	S15	
pWX854	pNPTS138 Δ <i>repB^{Ch2}</i> (Atu3923/ATU_RS18280)::amp	S15	
pWX855	pNPTS138 Δ <i>xerD</i> (Atu3629/ATU_RS16850)::amp	This study	
pWX885	pMiniTn7 <i>PtraI-riboswitch-parB1</i> (Atu2828/ATU_RS13770) <i>traR</i> (Atu6134: ATU_RS23655)	S15	

pWX902	pNPTS138 <i>tetRA::gen PtraI-riboswitch-parB1</i> (Atu2828/ATU_RS13770) <i>traR</i>	S15	
pWX914	pNPTS138 Δ <i>parB1</i> (Atu2828/ATU_RS13770)::amp	S15	
pWX916	pACYC <i>terminator Ppen cfp-parB^{P1}-parS^{P1}</i> (amp)	S17	
pWX923	pACYC <i>terminator Ppen CFP-ParBP1 parSP1 terminators</i>	This study	
pWX926	pNPTS138 <i>Ppen ygfp-parB^{MT1}-parS^{MT1}</i> at Atu1460/ATU_RS07195	This study	
pWX930	pNPTS138 <i>Ppen cfp-parB^{P1}-parS^{P1}</i> at Atu3054/ATU_RS14060	S17	
pWX936	pNPTS138 <i>PT7strong cfp-parB^{P1}-parS^{P1}</i> at Atu3054/ATU_RS14060	S17	
pWX943	pNPTS138 <i>terminator gen lacI terminators Plac-riboswitch</i>	This study	
pWX944	pNPTS138 <i>terminator gen LacI terminators Plac riboswitch xerC</i> (Atu2628/ATU_RS12790)	This study	
pWX945	pNPTS138 <i>terminator gen LacI terminators Plac riboswitch xerD</i> (Atu3629/ATU_RS16850)	This study	
pWX965	pNPTS138 <i>PT7strong ygfp-parB^{MT1}-parS^{MT1}</i> at Atu1460/ATU_RS07195	This study	
pWX968	pNPTS138 <i>PT7strong cfp-parB^{P1}-parS^{P1}</i> at Atu4854	This study	
pWX970	pSRKKm <i>Plac rfp-repB^{Ch2}</i> (Atu3923/ATU_RS18280) <i>terminator Plac egfp-parB1</i> (Atu2828/ATU_RS13770) <i>terminators</i>	S15	
pWX998	pNPTS138 <i>PT7strong mcherry-parB^{P1}-parS^{P1}</i> at Atu4854	This study	
pWX1006	pNPTS138 Δ <i>xerC</i> (Atu2628/ATU_RS12790)::amp	This study	
pWX1007	pNPTS138 Δ <i>recA</i> (Atu1873/ATU_RS09160)::amp	This study	
pWX1008	pNPTS138 <i>xerC-gfpmut3</i> (Atu2628/ATU_RS12790)	This study	
pWX1009	pNPTS138 <i>xerD-gfpmut3</i> (Atu3629/ATU_RS16850)	This study	
pWX1039	pNPTS138 Δ <i>dif1</i> ::amp	This study	
pWX1040	pNPTS138 Δ <i>dif^{PAf}</i> ::amp	This study	
pWX1076	pSRKKm <i>cymR cuO Plac cuO At parC</i> (Atu1158/ATU_RS05720) <i>At parE</i> (Atu1622/ATU_RS07965)	This study	
pWX1080	pSRKKm <i>cymR cuO Plac cuO Ec parE-parC</i>	This study	
PHP45omega	<i>omega-amp cassette</i>	S18	
pNPTS138	<i>oriT sacB kan</i>	S19	
pSRKKm	Broad host-range, <i>Plac</i>	S20	
cumate induction system	pSRKKm <i>cymR cuO Plac cuO msfgfp</i>	S21 gift from Pam Brown	
PLEXRparEC3	<i>E. coli parE parC</i> expressed from IPTG inducible promotor	S22	

Table S3. Bacterial strains and plasmids used in this study. Related to STAR Methods.

oWX2463	gcgcaggggtatcaacaagatgagcgacaaaccgatggc	pWX945
oWX2464	gctagcgaattcgtggatccagatacctcctactgtcgagg	pWX945
oWX2482	agcttctcgcaggataaaaaaaaaaaaaaccggcttcgcgggtacttgg	pWX943
oWX2483	cgcgaaagcggggtttttttccatccccgaaactatgc	pWX944
oWX2484	gcatacgatccggggatggaaaaaaaaaaaaaccggcttcgcg	pWX944
oWX2485	cggaaaggcgggttttttcatgggtgcgccttcgcg	pWX945
oWX2486	cggcaaggcgcacccataaaaaaaaaaaaaccggcttcgcg	pWX945
oWX2497	aagtcaagttgaagggtatacc	sequencing
oWX2491	catcaaacatcgaccacgg	sequencing
oWX2492	gaacgatgccctcattcagc	sequencing
oWX2493	tgcgacatcgataacgtactgg	sequencing
oWX2514	cgcgaagcttcgtcaggatatctcgccaattagaatggcctgt	pWX968
oWX2515	aattcagcatgcgtatctagagttaaatgtggcgtcctgaaag	pWX968
oWX2516	ctgagcaataactagcataacccgcgtgtcgccgcctactatc	pWX968
oWX2517	agcgaattcgtggatccagatatccgtcatcgatcagcaattcgatac	pWX968
oWX2518	tgtcaccacccgtccagaacaagct	sequencing
oWX2519	ggcaaggttcatcaaagacctgcga	sequencing
oWX2584	ttcaagcttaaggaggtggaaacatggcagcaaggagagagaag	pWX998
oWX2585	ctgctcgtcggatccatcgatcgttgcgttgtccatccacc	pWX998
oWX2589	cttcctctccctgtcgaccatgttccacccctttaagcttg	pWX998
oWX2590	ggtggaatggcgaattataaaaaaaatccagaaagaccgttcgaacacgccc	pWX998
oWX2601	cgcgaatccgtcaggatatccagaaagaccgttcgaacacgccc	pWX1006
oWX2602	aaataaacaaaatgggttccgcacgcgtactccattccatcccc	pWX1006
oWX2603	gcctcactgattaaggcattggtaaaaaatctacgacaacgcaccc	pWX1006
oWX2604	ctagcgaattcgtggatccagatatccagcatccaggcgttcgc	pWX1006
oWX2605	ggcgccaaagcttcgtcaggatatcgatgtcggtcatcgaaaaac	pWX1007
oWX2606	aaataaacaaaatgggttccgcggcagcgcacgaggcgtatggatattc	pWX1007
oWX2607	gcctcactgattaaggcattggtaaaaatgttgcgcattccaaaccac	pWX1007
oWX2608	gctagcgaattcgtggatccagatatccgtcagcgatcccgccag	pWX1007
oWX2609	tggcgccaaagcttcgtcaggatatcgatgtcggtttctggccaaac	pWX1008
oWX2610	tccttactggatcccgacatccggcgtgggtggcgttgc	pWX1008
oWX2611	ggatctggaggatcccgatcaaaggag	pWX1008/pWX1009
oWX2612	ttattttgtatagttcatccatgcc	pWX1008/pWX1009
oWX2613	ggcatggatgaactataaaaaaaacggaaatggtaaccgttaaac	pWX1008
oWX2614	gctagcgaattcgtggatccagatagaaaaaggagagaccgggttcgttgc	pWX1008
oWX2615	tggcgccaaagcttcgtcaggatcttcattccggaaaggcgtgc	pWX1009
oWX2616	tccttactggatcccgacatccatcaaggtttgcgttgc	pWX1009
oWX2617	ggcatggatgaactataaaaaaaacggacccggaaaccgcggataatc	pWX1009
oWX2618	gctagcgaattcgtggatccagatcgatgtcgagcatataaacgcg	pWX1009
oWX2624	ggataataccgcgcacatagcag	sequencing
oWX2625	tgcgcaaaactattaaactggcgaac	sequencing
oWX2626	gtataacgactcaactgtggggcc	sequencing
oWX2627	gataatggctgtcgatgttgcacgc	sequencing
oWX2628	ttgacttagagggcgtacgcgttgc	sequencing
oWX2720	tgaccgcgttcgtgttgcgg	sequencing
oWX2796	tggcgccaaagcttcgtcaggatccgggttcgtataagaacgggt	pWX1039
oWX2797	aaataaacaaaatagggttccgcgttgcgttgcacatttc	pWX1039
oWX2798	tagaatgtaccccgatcacaacaaacgcggaaacccttattttttt	pWX1039
oWX2799	aaatccatgggttcaqaqacaaatttaccaatgttcaatcgtqaggc	pWX1039

oWX2800	gcctcactgattaagcattggtaaattgtctctgaaaccttaggattt	pWX1039
oWX2801	gctagcgaaatcgatcccgatctctgcgcggcatttcgacc	pWX1039
oWX2802	tggcgccaagctctctgcaggattcgggtctgtataagaacggtg	pWX1040
oWX2803	aaataaaacaatagggttccgcgaataggaaatgggtacattc	pWX1040
oWX2804	tagaatgtaccaattccctattcgcggaaaccctatttgtttttt	pWX1040
oWX2805	gccctcgcccacatcgccgaaataattaccaatgctaattcgtgaggc	pWX1040
oWX2806	gcctcactgattaagcattggtaaatttccgcgttccgcagggc	pWX1040
oWX2807	gctagcgaaatcgatcccgatccagataagaactgttccgcgttcatgc	pWX1040
oWX2847	ctcacggtaaatggtcgagcgatg	sequencing
oWX2848	tcgtgaggatggccggatgagac	sequencing
oWX2849	gcttgatctcacggtaatggtcg	sequencing
oWX2850	tggttatgctctcgacgagcgctc	sequencing
oWX2912	atcgagtcaaggaagacacatggaaaagatctgtaccccg	pWX1076
oWX2913	catgttccacctcccttcaatggtcacccccaaacttccgcgtcg	pWX1076
oWX2914	ccattgaaaggaggtggaaacatgtggacacagcaacgatcttc	pWX1076
oWX2915	atggcagcagtttccgcgttgc	pWX1076
oWX2920	tcgaatttgcattcgaattgtctcaatccagattatccgcg	sequencing
oWX2922	gcaatcaacttgcctgggtctcc	sequencing
oWX2923	atcggttaagttccacccgcattgc	sequencing
oWX2924	agctgtcgccgcggagcgatag	sequencing
oWX2925	gtgcgggtttcggctgcacattc	sequencing
oWX2926	ttcagggatttcgcattcggccgg	sequencing
oWX2927	gcagcggcaggatggcgtatttgc	sequencing
oWX2936	ttatcgagtcaaggaagacacatgtggcaaaactataacgctgtatg	pWX1080
oWX2939	cgaatttgcattcgaattgtctacttgcgttatcaccgcgtgc	pWX1080
oWX2940	tctgcattgcaggcgttacattctc	sequencing
oWX2941	cagacgaaagagcgttctcttcg	sequencing
oWX2942	cagacgaaagagcgttctcttcg	sequencing
oWX2943	accgcataacctgcgtaaatggc	sequencing
oWX2944	actgcgttcatcttgcacaaactgg	sequencing
oWX2945	agtgtatgcggcagctgtcaag	sequencing
F1 (oWX2212)	tgtctgggttctggaaattcgacgc	check configuration
R1 (oWX2213)	agggtccgtggatagtgttaggc	check configuration
F2 (oWX2214)	cttgatccagaaatgttgcacgc	check configuration
R2 (oWX2215)	ccttgtgaacaacgccttgcaccc	check configuration

Table S4. Oligonucleotides used in this study. Related to STAR Methods.

Supplemental References

- S1. Figueira-Cuilan, W., Daniel, J.J., Howell, M., Sulaiman, A., and Brown, P.J. (2016). Mini-Tn7 Insertion in an Artificial attTn7 Site Enables Depletion of the Essential Master Regulator CtrA in the Phytopathogen *Agrobacterium tumefaciens*. *Appl Environ Microbiol* 82, 5015-5025. 10.1128/AEM.01392-16.
- S2. Morton, E.R., Merritt, P.M., Bever, J.D., and Fuqua, C. (2013). Large deletions in the pAtC58 megaplasmid of *Agrobacterium tumefaciens* can confer reduced carriage cost and increased expression of virulence genes. *Genome Biol Evol* 5, 1353-1364. 10.1093/gbe/evt095.
- S3. Farrand, S.K., and Dessaix, Y. (1986). Proline biosynthesis encoded by the noc and occ loci of *Agrobacterium* Ti plasmids. *J Bacteriol* 167, 732-734. 10.1128/jb.167.2.732-734.1986.
- S4. Fuqua, C., Bурбя, M., and Winans, S.C. (1995). Activity of the *Agrobacterium* Ti plasmid conjugal transfer regulator TraR is inhibited by the product of the traM gene. *J Bacteriol* 177, 1367-1373. 10.1128/jb.177.5.1367-1373.1995.
- S5. Robalino-Espinosa, J.S., Zupan, J.R., Chavez-Arroyo, A., and Zambryski, P. (2020). Segregation of four *Agrobacterium tumefaciens* replicons during polar growth: PopZ and PodJ control segregation of essential replicons. *Proc Natl Acad Sci U S A* 117, 26366-26373. 10.1073/pnas.2014371117.
- S6. Ellis, J.G., Kerr, A., Tempe, J., and Petit, A. (1979). Arginine catabolism: a new function of both octopine and nopaline Ti-plasmids of *Agrobacterium*. *Mol Gen Genet* 173, 263-269. 10.1007/BF00268636.
- S7. Luo, Z.Q., Clemente, T.E., and Farrand, S.K. (2001). Construction of a derivative of *Agrobacterium tumefaciens* C58 that does not mutate to tetracycline resistance. *Mol Plant Microbe Interact* 14, 98-103. 10.1094/MPMI.2001.14.1.98.
- S8. Morton, E.R., Platt, T.G., Fuqua, C., and Bever, J.D. (2014). Non-additive costs and interactions alter the competitive dynamics of co-occurring ecologically distinct plasmids. *Proc Biol Sci* 281, 20132173. 10.1098/rspb.2013.2173.
- S9. Goodner, B., Hinkle, G., Gattung, S., Miller, N., Blanchard, M., Quroollo, B., Goldman, B.S., Cao, Y., Askenazi, M., Halling, C., et al. (2001). Genome sequence of the plant pathogen and biotechnology agent *Agrobacterium tumefaciens* C58. *Science* 294, 2323-2328. 10.1126/science.1066803.
- S10. Chesnokova, O., Coutinho, J.B., Khan, I.H., Mikhail, M.S., and Kado, C.I. (1997). Characterization of flagella genes of *Agrobacterium tumefaciens*, and the effect of a bald strain on virulence. *Mol Microbiol* 23, 579-590. 10.1046/j.1365-2958.1997.d01-1875.x.
- S11. Wu, C.F., Santos, M.N.M., Cho, S.T., Chang, H.H., Tsai, Y.M., Smith, D.A., Kuo, C.H., Chang, J.H., and Lai, E.M. (2019). Plant-Pathogenic *Agrobacterium tumefaciens* Strains Have Diverse Type VI Effector-Immunity Pairs and Vary in In-Planta Competitiveness. *Mol Plant Microbe Interact* 32, 961-971. 10.1094/MPMI-01-19-0021-R.
- S12. Anand, A., Uppalapati, S.R., Ryu, C.M., Allen, S.N., Kang, L., Tang, Y., and Mysore, K.S. (2008). Salicylic acid and systemic acquired resistance play a role in attenuating crown gall disease caused by *Agrobacterium tumefaciens*. *Plant Physiol* 146, 703-715. 10.1104/pp.107.111302.
- S13. Watson, B., Currier, T.C., Gordon, M.P., Chilton, M.D., and Nester, E.W. (1975). Plasmid required for virulence of *Agrobacterium tumefaciens*. *J Bacteriol* 123, 255-264. 10.1128/jb.123.1.255-264.1975.
- S14. Liu, P., and Nester, E.W. (2006). Indoleacetic acid, a product of transferred DNA, inhibits vir gene expression and growth of *Agrobacterium tumefaciens* C58. *Proc Natl Acad Sci U S A* 103, 4658-4662. 10.1073/pnas.0600366103.
- S15. Ren, Z., Liao, Q., Karaboga, X., Barton, I.S., Schantz, E.G., Mejia-Santana, A., Fuqua, C., and Wang, X. (2022). Conformation and dynamic interactions of the multipartite genome in *Agrobacterium tumefaciens*. *Proc Natl Acad Sci U S A* 119. 10.1073/pnas.2115854119.
- S16. Grangeon, R., Zupan, J., Jeon, Y., and Zambryski, P.C. (2017). Loss of PopZ At activity in *Agrobacterium tumefaciens* by Deletion or Depletion Leads to Multiple Growth Poles, Minicells, and Growth Defects. *mBio* 8. 10.1128/mBio.01881-17.

- S17. Ren, Z., Liao, Q., Barton, I.S., Wiesler, E.E., Fuqua, C., and Wang, X. (2022). Centromere Interactions Promote the Maintenance of the Multipartite Genome in *Agrobacterium tumefaciens*. *mBio*, e0050822. 10.1128/mbio.00508-22.
- S18. Fellay, R., Frey, J., and Krisch, H. (1987). Interposon mutagenesis of soil and water bacteria: a family of DNA fragments designed for in vitro insertional mutagenesis of gram-negative bacteria. *Gene* 52, 147-154. 10.1016/0378-1119(87)90041-2.
- S19. Hinz, A.J., Larson, D.E., Smith, C.S., and Brun, Y.V. (2003). The *Caulobacter crescentus* polar organelle development protein PodJ is differentially localized and is required for polar targeting of the PleC development regulator. *Mol Microbiol* 47, 929-941. 10.1046/j.1365-2958.2003.03349.x.
- S20. Khan, S.R., Gaines, J., Roop, R.M., 2nd, and Farrand, S.K. (2008). Broad-host-range expression vectors with tightly regulated promoters and their use to examine the influence of TraR and TraM expression on Ti plasmid quorum sensing. *Appl Environ Microbiol* 74, 5053-5062. 10.1128/AEM.01098-08.
- S21. Choi, Y.J., Morel, L., Le Francois, T., Bourque, D., Bourget, L., Groleau, D., Massie, B., and Miguez, C.B. (2010). Novel, versatile, and tightly regulated expression system for *Escherichia coli* strains. *Appl Environ Microbiol* 76, 5058-5066. 10.1128/AEM.00413-10.
- S22. Madabhushi, R., and Marians, K.J. (2009). Actin homolog MreB affects chromosome segregation by regulating topoisomerase IV in *Escherichia coli*. *Mol Cell* 33, 171-180. 10.1016/j.molcel.2009.01.001.