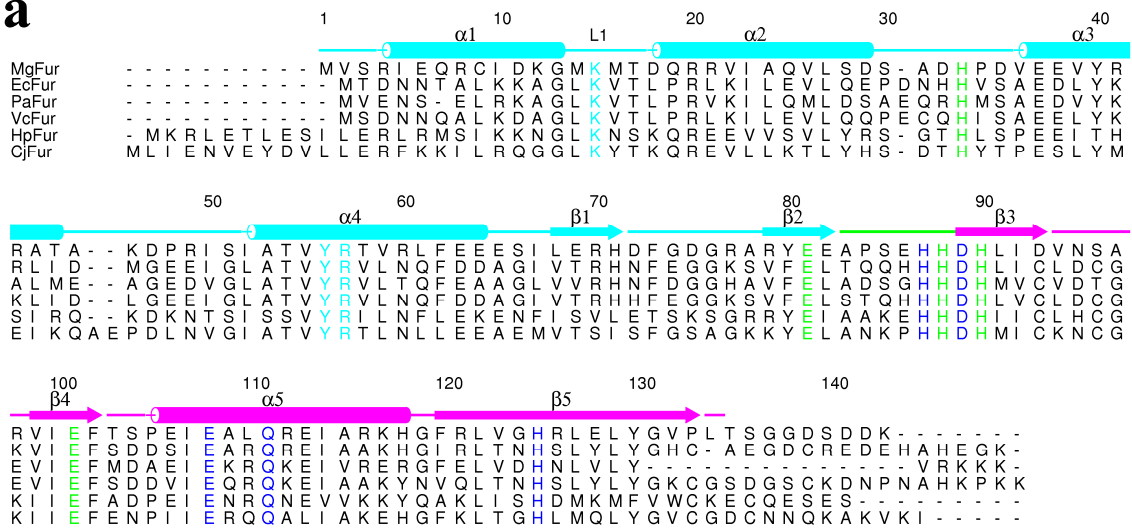
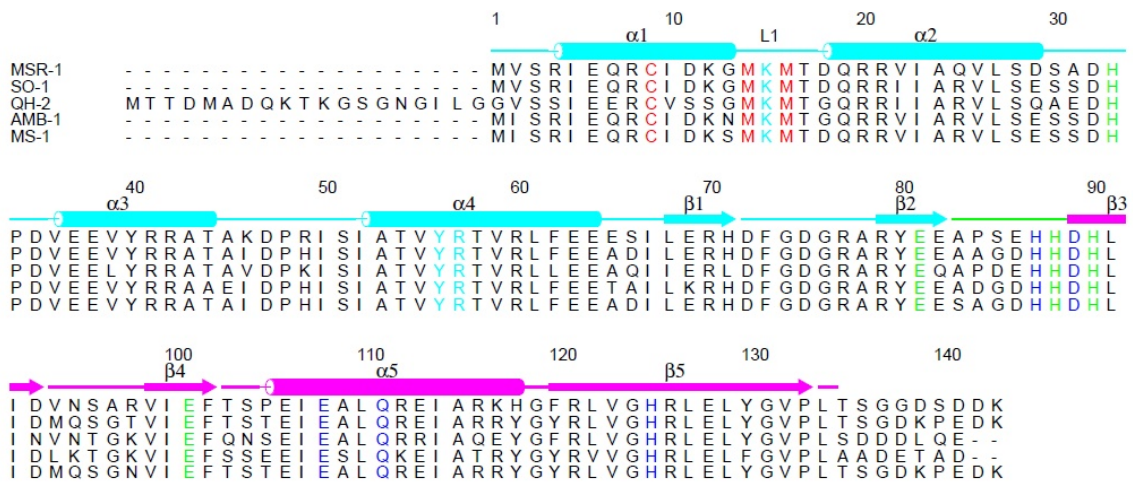
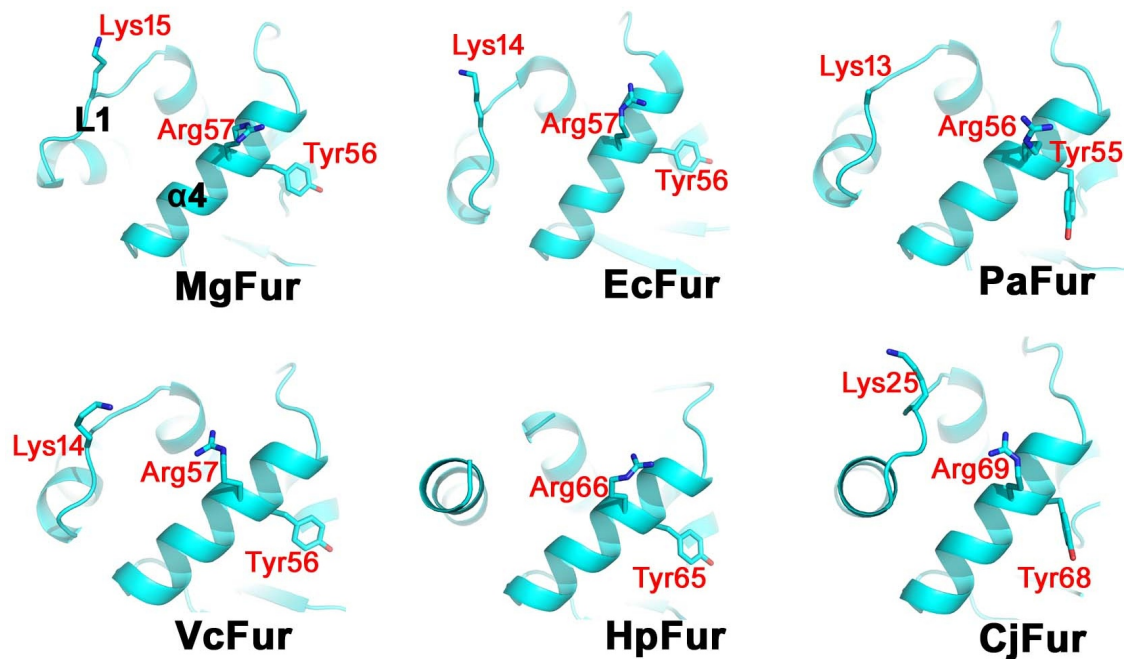
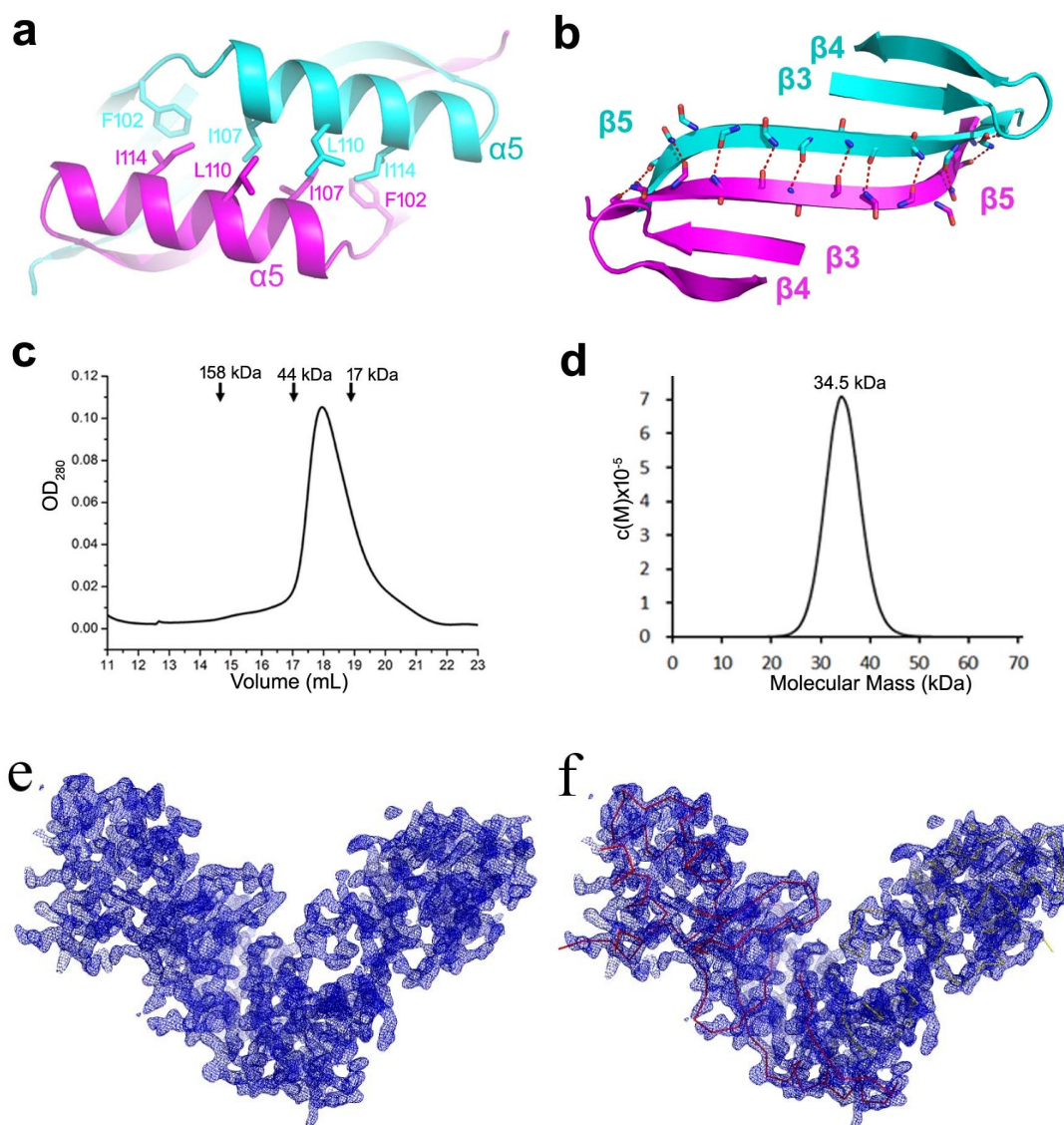


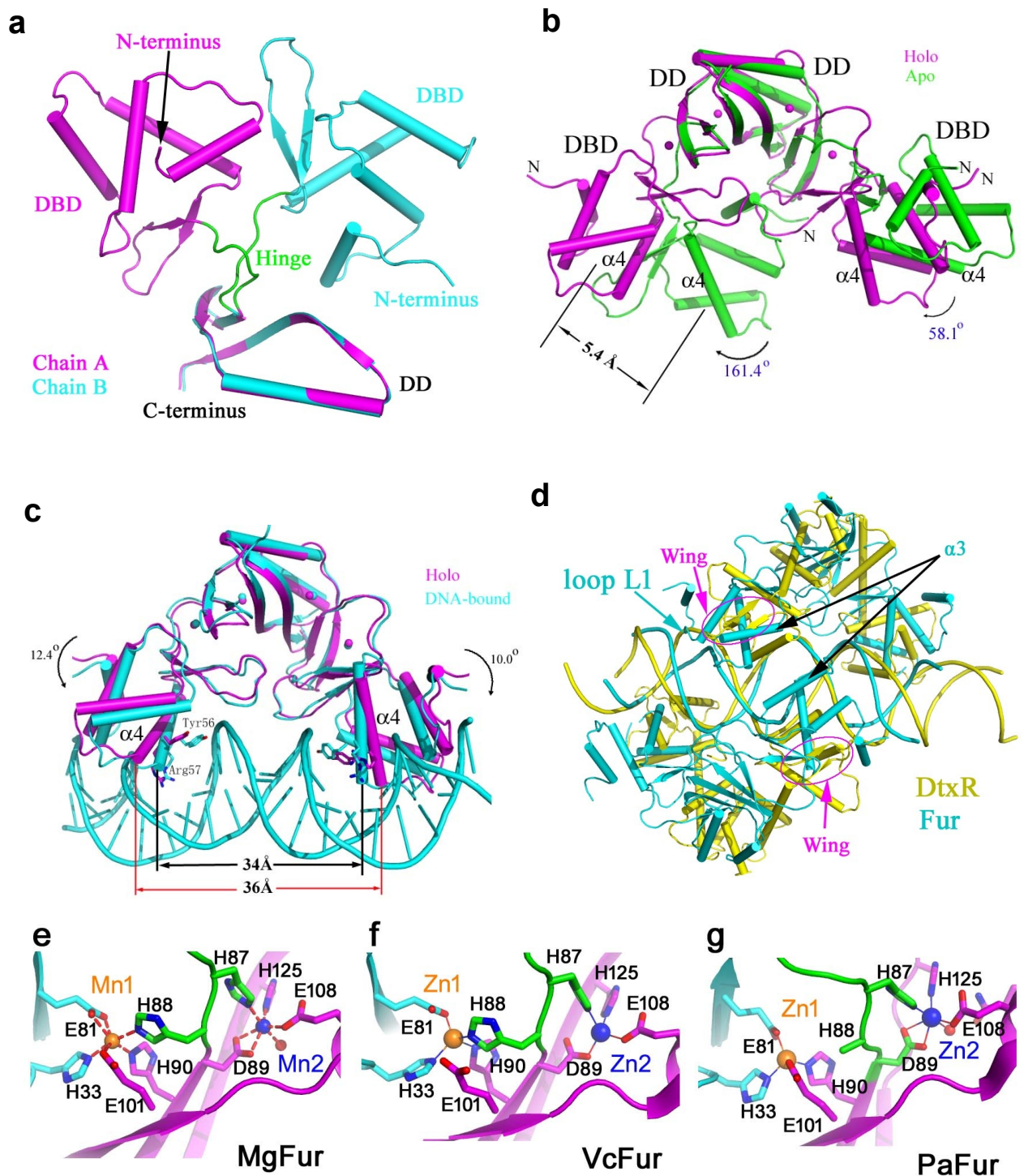
a**b**

C

Supplementary Figure 1. Sequence alignment of prokaryotic Fur proteins. (a) The secondary structure elements of MgFur are shown above the alignment and colored by domains as in Figure 2a. The N-terminal DBD, hinge and C-terminal DD are shown in cyan, green and magenta, respectively. Residues of binding site 1 and binding site 2 are colored in green and blue, respectively. Residues that are important for interacting with the DNA are colored in cyan. (b) Sequence alignment of Fur proteins from five high homology *Magnetospirillum* or *Magnetospira* strains. The sequences used are as follows: *M. gryphiswaldense* MSR-1, *Magnetospirillum* sp. SO-1 (AONQ00000000), *Magnetospira* sp. QH-2 (FO538765), *M. magneticum* AMB-1 (AP007255), *M. magnetotacticum* MS-1 (AAAP00000000). The red colored letters show the conserved sulfur-containing amino acids in *Magnetospirillum* and *Magnetospira* strains. (c) Superimposition of the DBDs of different prokaryotic Fur proteins. The DBDs are shown in the same orientation. Note that the lysine residue was not shown in structures of HpFur due to its flexible conformations in the holo-Fur structures. Sequences and structures used: *M. gryphiswaldense* (Mg, this study), *E. coli* (Ec; PDB ID: 2FU4), *P. aeruginosa* (Pa; PDB ID: 1MZB), *V. cholerae* (Vc; PDB ID: 2W57), *H. pylori* (Hp; PDB ID: 2XIG), *C. jejuni* (Cj; PDB ID: 4ETS).

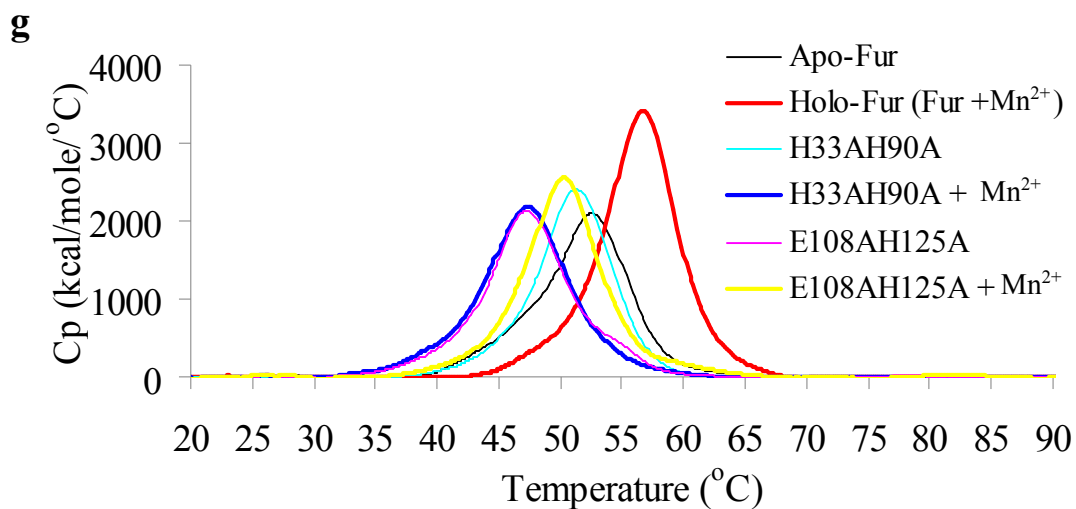
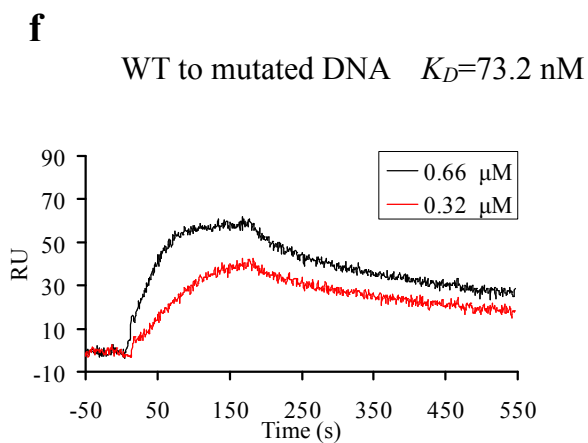
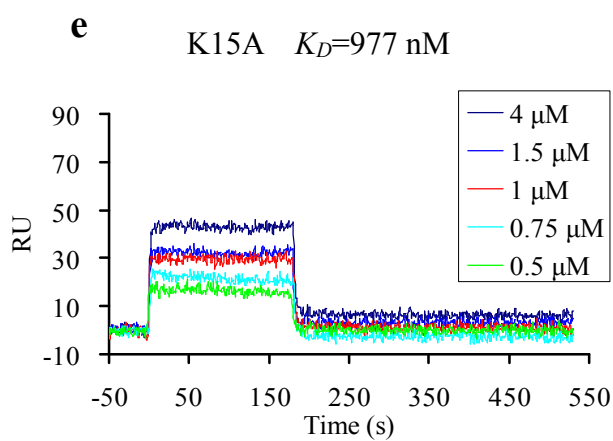
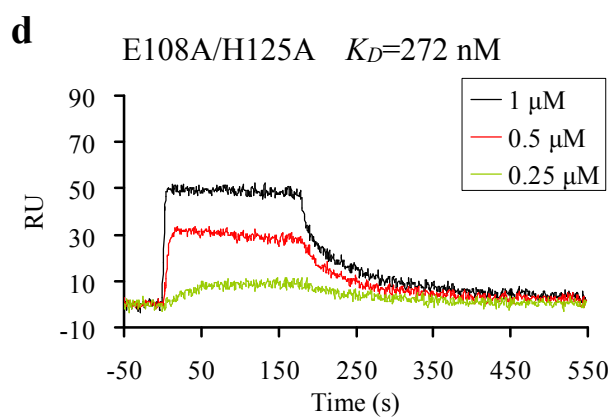
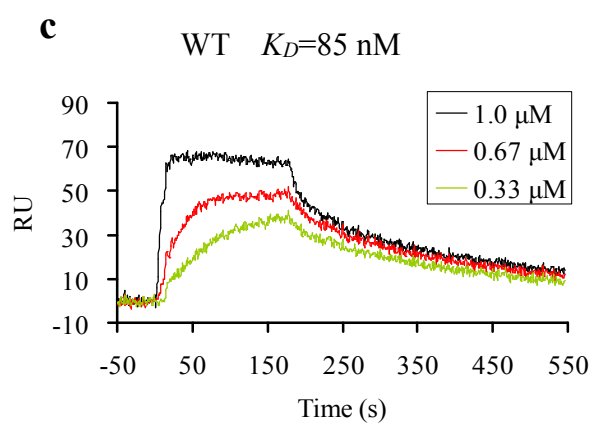
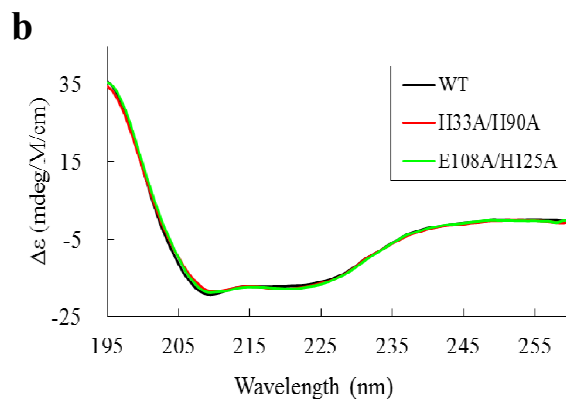
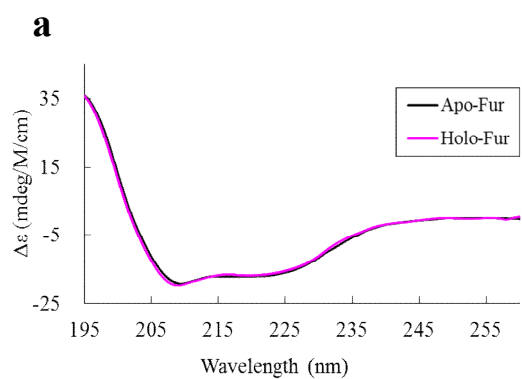


Supplementary Figure 2. Apo-Fur forms a dimer in crystal and solution. (a) Hydrophobic interactions between the two antiparallel helices $\alpha 5$ in the two monomers. (b) Two antiparallel $\beta 5$ strands form hydrogen bonds. (c) Elution profile of apo-Fur by size-exclusion chromatography. The elution volumes of the molecular mass standards are marked at the top of the panel. (d) Analytical ultracentrifugation of apo-Fur, $c(s)$ distribution from sedimentation velocity analytical ultracentrifugation experiments (SV) performed at 0.1 mM protein. Note that the theoretical mass of Fur monomer is 16.4 kDa. (e) Experimental electron density map of holo-Fur. (f) Experimental electron density map of holo-Fur with solved backbone traces of holo-Fur. The two molecules of holo-Fur in one asymmetric unit in the $I222$ space group. The electron density is contoured at 2.0σ . The backbone traces of the two molecules are colored in different colors.



Supplementary Figure 3. Comparison of different states of Fur. (a) Comparison of the two monomer conformations in apo-Fur. (b) Side view of the recognition helix $\alpha 4$ to highlight the conformational change of the DBD. The key residues Tyr56 and Arg57 are shown as sticks and the distances between the two helices $\alpha 4$ in holo-Fur (magenta) and the DNA-bound form (cyan) are labeled. The arrow indicates the movement of the DBD in the DNA-bound state compared to the holo state. In addition, two DBDs decrease their distance from each other after binding to the *feoAB1*

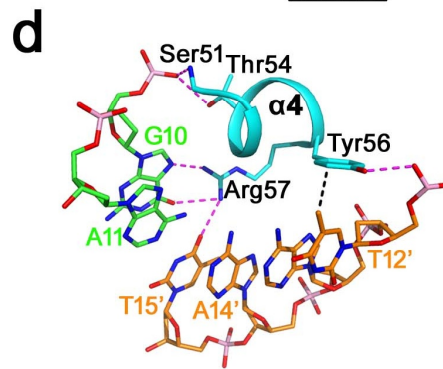
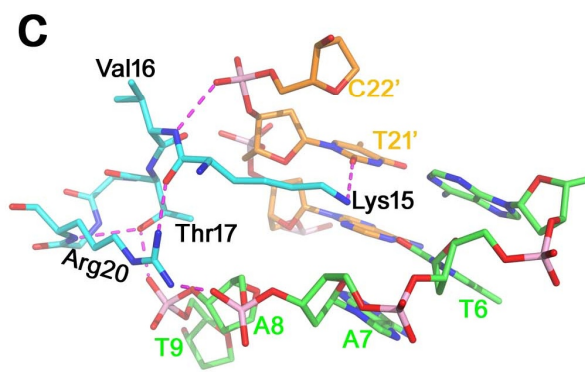
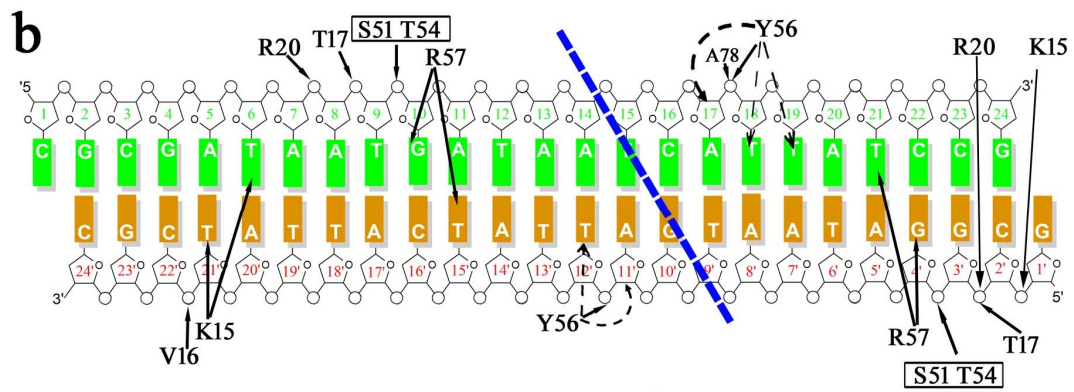
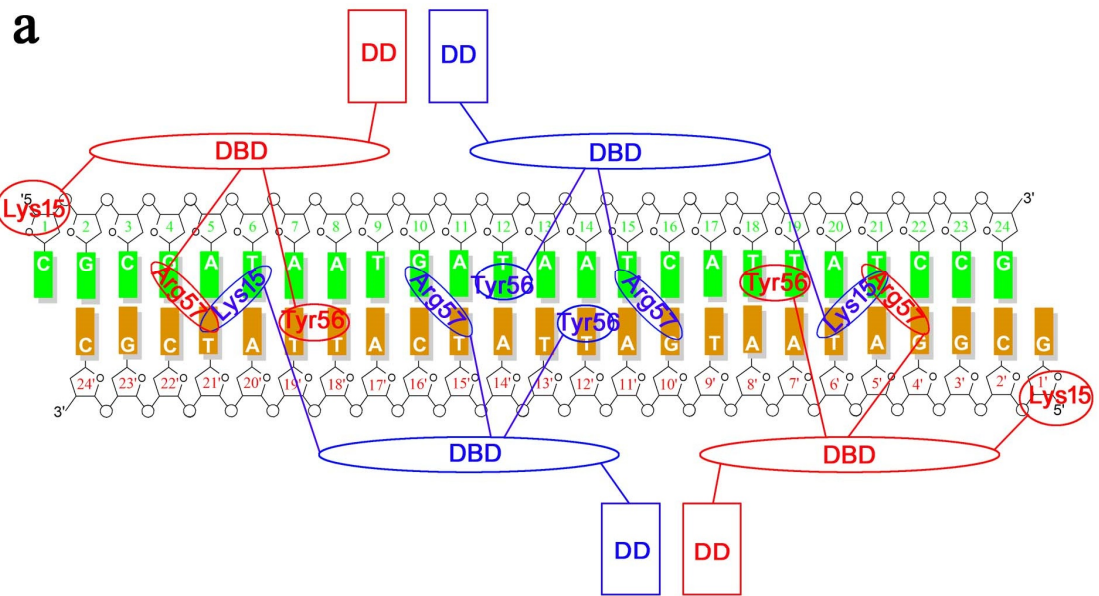
operator. (c) Comparison of holo-Fur and the Fur-*feoABI* operator complex. (d) Comparison of the DtxR-*tox* operator (PDB ID: 1DDN) and Fur-*P. aeruginosa* Fur box complex. Note that the wing of DtxR is shown by a magenta arrow. The wing of DtxR has an anti-parallel β -sheet, which is not found in the corresponding position of Fur. Coordination of the two metal ion binding sites in (e) MgFur, (f) VcFur (PDB ID: 2w57), and (g) PaFur (PDB ID: 1MZB). To make the comparison clear, the residue number is not changed for PaFur (The actual residue number of PaFur is the labeled number minus one).

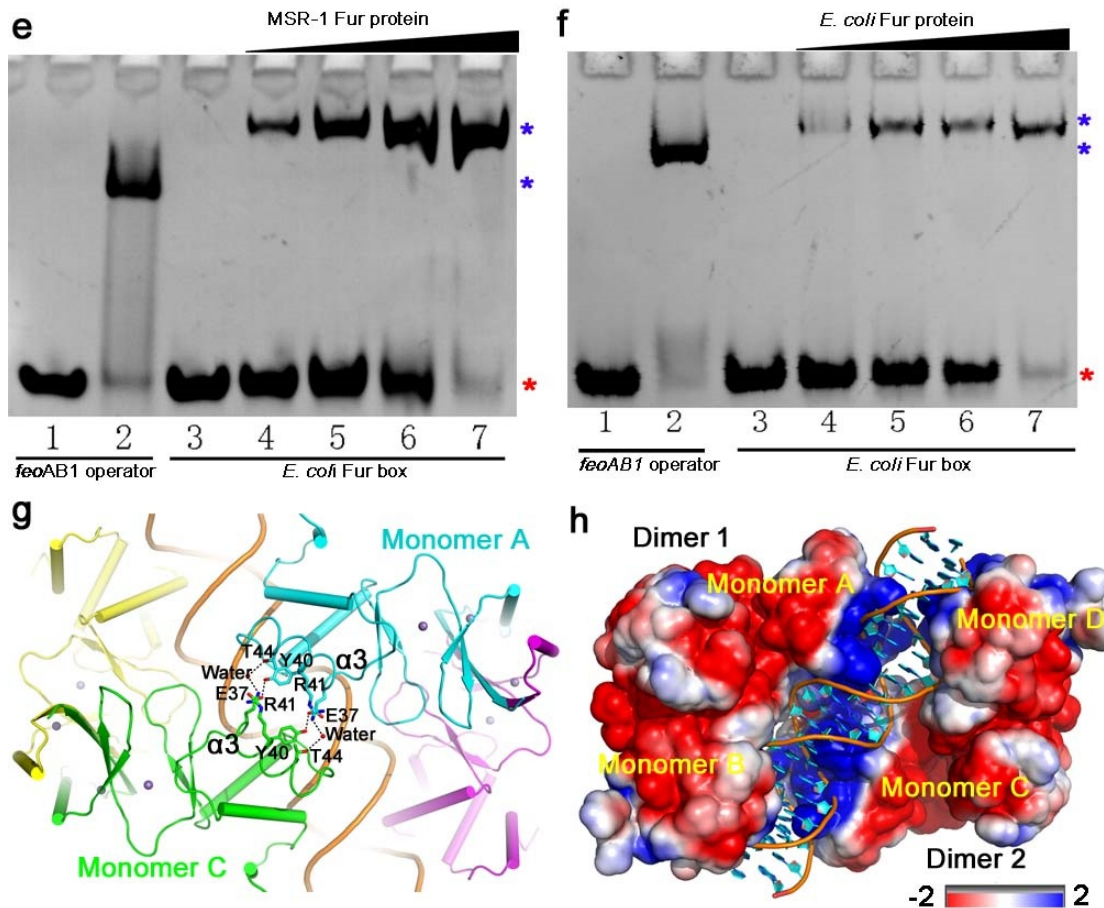


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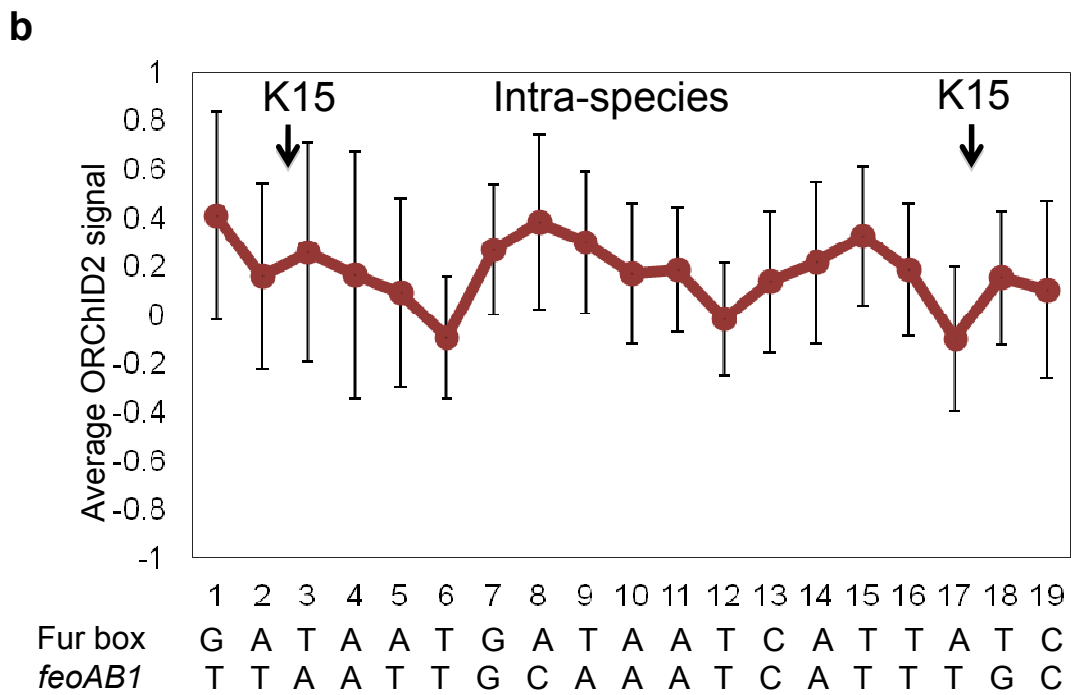
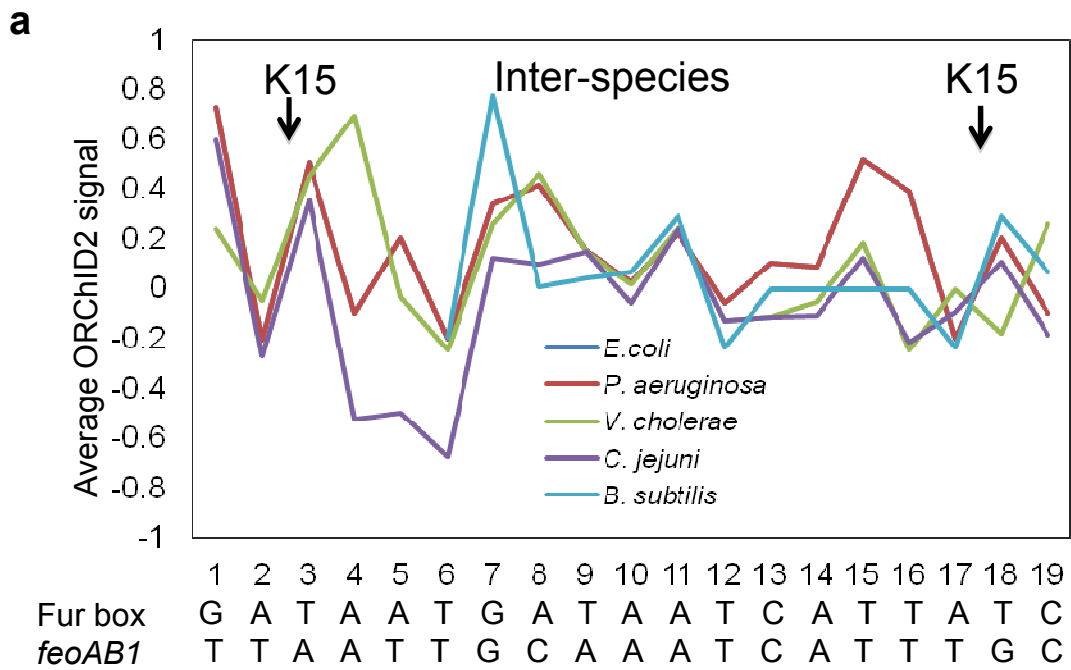
	$t_{1/2}$ (°C)	ΔH (kcal/mol)	T_{onset} (°C)	T_m (°C)
Apo-Fur	7.38	18600	39.42	52.49
Holo-Fur (Apo-Fur + Mn ²⁺)	6.52	26900	44.46	56.68
H33AH90A	6.86	18800	39.2	51.26
H33AH90A + Mn ²⁺	7.54	20000	33.03	47.28
E108AH125A	7.2	19200	33.93	47.33
E108AH125A + Mn ²⁺	6.69	20800	37.61	50.35

Supplementary Figure 4. Properties of Fur and its mutants. (a) CD spectra of apo-Fur and holo-Fur in the presence of 1 mM Mn²⁺. It shows that the addition of metal ions does not alter the secondary structures. (b) CD spectra of the WT, H33A/H90A and E108A/H125A mutants. These data demonstrate that the secondary structures of the two mutants in the metal ion binding sites are similar to that of the WT. (c-f) SPR experiment of Fur-*feoABI* operator binding. Binding of WT Fur and mutants to the *feoABI* operator, determined by surface plasmon resonance (SPR) analysis. The *FeoABI* operator was immobilized onto an NLC sensor chip and subsequently the buffer containing each Fur at the corresponding concentrations was run over the immobilized *feoABI* operator. K_D values calculated from SPR data are also presented. (g) DSC profile of Cp vs. T and thermodynamic binding parameters of (h) Fur mutants with or without Mn²⁺ obtained by subtracting the baseline from the original scan. 1.5 mg/ml protein was used for each sample. All experiments were repeated three times.





Supplementary Figure 5. Interactions between two MgFur dimers and the *P. aeruginosa* Fur box. (a) Schematic diagram of two Fur-DNA interactions. (b) Detailed Fur-DNA interactions between one Fur dimer with DNA. Solid arrows and dashed arrows indicate hydrogen bonds and van der Waals interactions, respectively. Another Fur dimer-DNA interaction is the same because the whole structure is related by a twofold symmetry axis. Close-up view of the interactions of loop 1 (L1) with (c) the minor groove and contacts between helix $\alpha 4$ and (d) the major groove. Magenta dashed lines represent hydrogen bonds and black dashed lines indicate van der Waals interactions. Note that one asymmetric unit contained one Fur dimer and one ssDNA molecule. The structure of one double stranded DNA fragment in complex with two Fur dimers was obtained by a symmetry operation using the twofold axis. (e) MSR-1 Fur protein and (f) *E. coli* Fur protein. Lane 1 and 3 are the control *feoAB1* operator or *P. aeruginosa* Fur box (25 pmol). The Fur protein concentration used in lane 2, lanes 4-7 are 100, 10, 20, 40 and 80 pmol, respectively. The corresponding protein complexes are shown as blue asterisks. Note that the *E. coli* Fur protein contained a His Tag and a TEV site. (g) Detailed interactions between two Fur dimers in Fur-Mn²⁺-*P. aeruginosa* Fur box complex. (h) Electrostatic surface potential of Fur-Mn²⁺-*P. aeruginosa* Fur box ternary complex using the APBS tool of PyMOL (unit: kT/e). The view is that of panel (g) rotated around the y-axis by 180°. The results show that two Fur dimers cooperatively bind to *P. aeruginosa* Fur box independent of protein concentration.

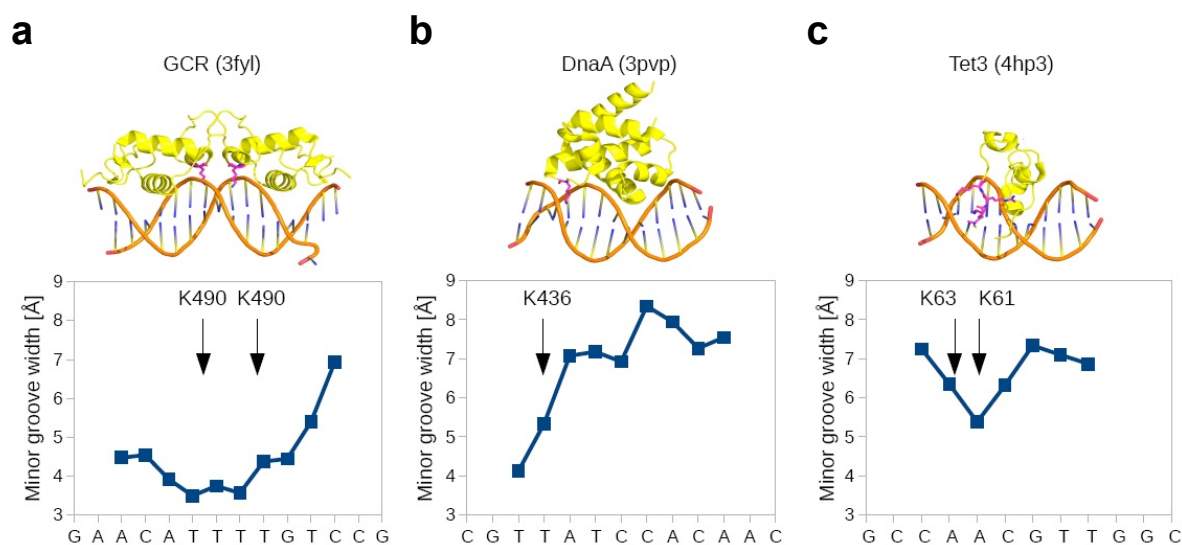


C

Gene	Fur-box element
Consensus	GATAATGATAATCATTATC
pig2	ACGAACGTGAATCATTCTC
pig3_1	GATAATGAGATTGATTATT
pig3_2	GAAAACAATAATCAATCTC
pig4_1	GACATTGAGATTCAATGAC

pig4_2	ATAATTGAGAATCGTTATT
pig6	GAAGATGGTAATTAATTGC
pig7_1	TCCAATGCAAATCAATATC
pig7_2	GCCAATGATATTGATTTGC
pig8	GTAATTGACAATCATTATC
pig9_1	GATAATGAGAATAGTTATT
pig9_2	TGTAATAATAACTATTCTC
pig12	TTTATCGCAACTGATTATC
pig13_1	CTGAATGATAATAATTATC
pig13_2	TCTTCTGATAATTATTATC
pig14	GCAAACGATATTCATCATC
pig17	AAAAACGAGAATTATTTCGC
pig18	GATATTAATACCCATTTCA
pig19	ACGATTGCTAATCAATCTT
pig20	GGAAATGAGAATCATTATT
pig23	CTGAATGAAACCCAATCTT
pig25_1	ACGATTGAAAATCATTATC
pig25_2	TTTTATGATAATGATTTTC
pig31	GTTATTGAGAATCATTGGC
pig32_1	C TTGATGAGAATTATTATG
pig32_2	TAAATTCATAATAATTCTC
fpvA_1	GCTGATCACGATGATGGTC
fpvA_2	GATCACGATGATGGTCTTG
pfeA	TCAAATAACAATCAATATC
fptA	CATAATGATAAGCATTATC
pchR_1	GGAAATGAGATTTATTATC
pchR_2	GAGATTTATTATCATTGGC

Supplementary Figure 6. Hydroxyl radical cleavage intensities of unbound Fur binding sites derived with ORChID2. (a) Consensus Fur-boxes among different species¹⁻⁴ and (b) average of 32 sequences bound by Fur from *P. aeruginosa*¹. The different sequences were aligned to the Fur box based on the conserved nucleotides. Positions where Lys15 would interact with the DNA minor groove, based on the co-crystal structures from this study, are highlighted by an arrow. The sequences used to generate panel (a) are listed in Supplementary Table 1. (c) The Fur-box elements of iron-regulated genes in *P. aeruginosa* were used to generate panel (b). See Methods for ORChID2 references.



Supplementary Figure 7. Lysine is employed to recognize minor groove shape and electrostatic potential

The Fur-DNA ternary complexes presented in this study reveal that lysine can play a key role in DNA shape readout (Figure 6). This finding expands our currently established knowledge that predominantly arginine⁵⁻⁹ and histidine¹⁰ employ this form of protein-DNA readout. Lysine, albeit its abundance at protein surfaces, is in relative terms less frequently found to insert into narrow minor groove regions. A reason for this observation is the smaller volume of its charged side chain, resulting in a smaller Born radius of its head group compared to the ones of arginine and histidine. This results in a higher desolvation cost if its hydration shell was to be removed in order for the lysine side chain to be inserted into a narrow region of the minor groove⁵.

After revealing the use of Lys15 side chains in Fur-DNA recognition, we surveyed the Protein Data Bank, which resulted, compared with arginine, in fewer examples where lysine alone has been observed to intrude a narrow minor groove, such as (a) the *Rattus norvegicus* glucocorticoid receptor DBD homodimer (PDB ID: 3FYL)¹¹, (b) the *Mycobacterium tuberculosis* DnaA-DBD (PDB ID: 3PVP)¹², and (c) the *Xenopus tropicalis* Tet3 CXXC domain (PDB ID: 4HP3)¹³. Among these examples, DnaA binds to origins of replication through the insertion of Lys436 into the minor groove, and when mutated to alanine its binding affinity decreases¹². The same study shows that in other bacteria, however, the respective position is occupied by an arginine instead of lysine, which carries the same positive charge while its desolvation cost is lower than that of lysine⁵.

Taken together, the ternary Fur-DNA complexes presented in this study revealed the use of lysine in recognizing intrinsically narrow regions of the minor groove with enhanced negative electrostatic potential, and the survey of the Protein Data Bank suggests that lysine can replace arginine in certain systems, thus expanding the currently known repertoire of DNA shape readout mechanisms.

Supplementary Table 1. DNA consensus boxes bound by Fur from different species. Conserved nucleotides are shown using colored font.

Species¹⁴	Sequences
<i>E.coli</i> ¹⁵	GATAATGATAATCATTATC
<i>P. aeruginosa</i> ¹	GATAATGATAATCATTATC
<i>V. cholerae</i> ²	GCCAAATGATAATTATTCTCATTGC
<i>C. jejuni</i> ³	TATTTTGATAATTATTATCA
<i>B. subtilis</i> ⁴	TGATAATNATTATCA

Supplementary Table 2. Metal ion–ligand distances in metal ion binding sites of holo-Fur.

Atom	Distance (Å)	Atom	Distance (Å)
Site 1		Site 2	
His-33 NE2	2.24	His-87 NE2	2.16
Glu-81 OE1	2.30	Asp-89 OD1	2.28
Glu-81 OE2	2.20	Asp-89 OD2	2.36
His-88 NE2	2.21	Glu-108 OE1	2.05
His-90 NE2	2.18	His-125 NE2	2.24
Glu-101 OE1	2.17	Water	2.26

Supplementary Table 3. Primers of quantitative real-time RT-PCR.

Gene name	Location	Primer name	Sequence	Product length (bp)
<i>feoB1</i>	EF120624.1	FeoB1-F	AAGGCGATGACGAGGTTCT	238
		FeoB1-R	AAGGTGCGGTTTCAGGAAGA	
<i>rpoC</i>	MGR_3807	RpoC-F	ATCCGTATTTCCATCGCCTCCC	164
		RpoC-R	TTGCCGCACAAGCATTCGT	

Supplementary Table 4. Strains and plasmids used in this study.

Strain and plasmid	Description	Source of reference
Strains		
<i>M. gryphiswaldense</i>		
MSR-1	Wild type	DSM 6361
F4	<i>fur3137</i> -defective mutant, Gm ^r	Ref. 16
F4C	Complement strain of F4, Gm ^r , Tc ^r	Present study
F4M	Complement strain of F4 with Fur3137-C9LM14LM16L, Gm ^r , Tc ^r	Present study
K15A	Complement strain of F4 with Fur3137-K15A, Gm ^r , Tc ^r	Present study
R57A	Complement strain of F4 with Fur3137-R57A, Gm ^r , Tc ^r	Present study
H33AH90A	Complement strain of F4 with Fur3137-H33AH90A, Gm ^r , Tc ^r	Present study
E108AH125A	Complement strain of F4 with Fur3137-E108AH125A, Gm ^r , Tc ^r	Present study
<i>E. coli</i> S17-1	<i>thiendArecAhsdR</i> with RP4-2-Tc::Mu-Km::Tn7 integrated in chromosome, Sm ^r	Ref. 17
Plasmid		
pRK415	Broad host range cloning vector, Tc ^r	Ref. 18
pRKF4M	pRK415 containing <i>tri-fur3137</i> , Tc ^r	Present study
pRKFK15A	pRK415 containing <i>fur3137</i> -K15A, Tc ^r	Present study
pRKFR57A	pRK415 containing <i>fur3137</i> -R57A, Tc ^r	Present study
pRKFH33AH90A	pRK415 containing <i>fur3137</i> -H33AH90A, Tc ^r	Present study
pRKFE108AH125A	pRK415 containing <i>fur3137</i> -E108AH125A, Tc ^r	Present study

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