

# Structures of $\omega$ repressors bound to direct and inverted DNA repeats explain modulation of transcription

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## ABSTRACT

**Repressor  $\omega$  regulates transcription of genes required for copy number control, accurate segregation and stable maintenance of *inc18* plasmids hosted by Gram-positive bacteria.**  $\omega$  belongs to homodimeric ribbon-helix-helix ( $RHH_2$ ) repressors typified by a central, antiparallel  $\beta$ -sheet for DNA major groove binding. Homodimeric  $\omega_2$  binds cooperatively to promoters with 7 to 10 consecutive non-palindromic DNA heptad repeats ( $5' \text{-A/TATCAC^A/T-} 3'$ , symbolized by  $\rightarrow$ ) in palindromic inverted, converging ( $\rightarrow\leftarrow$ ) or diverging ( $\leftarrow\rightarrow$ ) orientation and also, unique to  $\omega_2$  and contrasting other  $RHH_2$  repressors, to non-palindromic direct ( $\rightarrow\rightarrow$ ) repeats. Here we investigate with crystal structures how  $\omega_2$  binds specifically to heptads in minimal operators with ( $\rightarrow\rightarrow$ ) and ( $\rightarrow\leftarrow$ ) repeats. Since the pseudo-2-fold axis relating the monomers in  $\omega_2$  passes the central C–G base pair of each heptad with  $\sim 0.3$  Å downstream offset, the separation between the pseudo-2-fold axes is exactly 7 bp in ( $\rightarrow\rightarrow$ ),  $\sim 0.6$  Å shorter in ( $\rightarrow\leftarrow$ ) but would be  $\sim 0.6$  Å longer in ( $\leftarrow\rightarrow$ ). These variations grade interactions between adjacent  $\omega_2$  and explain modulations in cooperative binding affinity of  $\omega_2$  to operators with different heptad orientations.

## INTRODUCTION

Gene expression in prokaryotes is primarily regulated by helix-turn-helix proteins that bind specifically to palindromic operators whereas recognition of arrays of direct or inverted

repeats by transcriptional, homodimeric ribbon-helix-helix ( $RHH_2$ ) repressors like  $\omega$  protein is less frequent (1). Structures are known for  $RHH_2$  repressors Arc (2), CopG (3) and MetJ (4) bound to their cognate operators that are bent by 50° to 60°. MetJ<sub>2</sub> binds symmetrically to two to five consecutive 8 bp long palindromic repeats. By contrast, CopG<sub>2</sub> and Arc<sub>2</sub> bind asymmetrically to half sites of palindromic operators that are spaced by 10 and 11 bp, respectively. When bound to these operators, interactions between adjacent  $RHH_2$  contribute to high affinity and cooperative association.

Repressor  $\omega$  is a global regulator of and encoded by broad-host-range and low-copy number plasmids belonging to the *inc18* family that are stably maintained in Gram-positive bacteria (5–7).  $\omega$  was originally isolated from *Streptococcus pyogenes* plasmid pSM19035 where  $\omega_2$  controls promoter regions located upstream of genes involved in plasmid copy number control (*PcopS*), plasmid partitioning (*Pδ*) and post-segregational killing (*Pω*) if the plasmid is lost. These promoters comprise arrays of ten, nine or seven consecutive 7 bp repeats (heptads, symbolized by  $\rightarrow$ ), organized as: *PcopS*, ( $\rightarrow\rightarrow\leftarrow\rightarrow\leftarrow\rightarrow\rightarrow\leftarrow\rightarrow\leftarrow\rightarrow$ ); *Pδ*, ( $\rightarrow\rightarrow\rightarrow\rightarrow\rightarrow\rightarrow\rightarrow\leftarrow\leftarrow$ ) and *Pω*, ( $\rightarrow\rightarrow\leftarrow\rightarrow\rightarrow\leftarrow\leftarrow$ ) (1), see Supplementary Figure 9.

Binding of  $\omega_2$  to a single heptad or to heptads separated by one or more additional base pair is poor ( $k_D > 500$  nM), but tight if operators include at least two consecutive heptads and tightens further with increasing number of heptads. In addition, the affinity depends on heptad arrangement as shown by 6-fold reduced affinity of  $\omega_2$  for diverging repeats ( $\leftarrow\rightarrow$ ) ( $k_D \sim 120$  nM) compared to heptads in direct ( $\rightarrow\rightarrow$ ) or converging ( $\rightarrow\leftarrow$ ) arrangement ( $k_D \sim 20$  nM) (8). Multiple repeat binding sites are also found for eukaryotic operators that interact cooperatively with monomeric and therefore asymmetric transcription factors (1). However, these repeats show different base pair spacings and protein–protein interactions in direct and inverted orientation (9).

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In wild type (wt)  $\omega_2$ , the N-terminal 20 residues of the 71 residues long  $\omega$  monomers are probably unstructured as suggested by secondary structure prediction (10) and were cleaved during crystallization ( $\Delta 20\omega_2$ ). The structure features a typical RHH-fold comprising a 2-fold symmetrical  $\beta$ -sheet with antiparallel pairing of residues 28–32 of each monomer followed by  $\alpha$ -helices  $\alpha 1$  (34–46) and  $\alpha 2$  (51–66) (11).

We describe here the crystal structures of an N-terminal deletion mutant (see Results) with 19 residues removed, hereafter  $\Delta 19\omega$ , in complex with two minimal operators comprising two heptads in  $(\rightarrow\rightarrow)$  and  $(\leftarrow\leftarrow)$  arrangement. It was of interest to elucidate the structural determinants for high specificity, affinity and cooperative binding of  $\omega$  repressor to minimal binding sites and to extrapolate these to natural operators with different heptad arrangements.

Unintentionally, both complexes cocrystallized with free operator DNA. Free  $(\rightarrow\rightarrow)$ -DNA allowed us to compare structural changes in DNA induced by repressor binding, whereas free  $(\leftarrow\leftarrow)$ -DNA was ill-defined in the electron density and could not be fully modeled. The complex between  $\Delta 19\omega_2$  and  $(\leftarrow\rightarrow)$ -DNA dissociated during gel filtration and could not be crystallized.

## MATERIALS AND METHODS

### Plasmid construction

For expression of  $\Delta 19\omega$  in *Escherichia coli*, wt  $\omega$  gene missing the first 19 codons was cloned into NcoI–BamHI-cleaved pET28a (Novagen) to render pET28a- $\Delta 19\omega$ . The described (1) pHP14-borne  $\omega$  gene (pHP14 $\omega$ ) was modified to pHP14 $\Delta 19\omega$  containing promoter  $P\omega$ , the ribosomal binding site and the Met start codon fused to codon 20 of  $\omega$  gene. pHP14 $\omega$  mutants pHP14 $\omega$ Thr29Ala and pHP14 $\omega$ His38Val were generated by site-directed mutagenesis. The plasmids were transferred into *Bacillus subtilis* strain BG511 ( $P\omega: lacZ, recA4$ ) as described (1).

### $\beta$ -galactosidase assay

$\beta$ -galactosidase assays (Table 1) were performed as described (1) except that the centrifuged *B. subtilis* cells were resuspended and lysed by the addition of 0.1% SDS (final concentration 0.0025%) and chloroform.

### Preparation of protein–DNA complexes

$\Delta 19\omega_2$  was expressed in *E. coli* according to (1), the cell paste was resuspended in buffer A [50 mM Tris–HCl (pH 7.5), 50 mM NaCl] and lysed (French Press). The crude extract was processed (1), except that after the phosphocellulose step the fractions were pooled, diluted 5-fold with buffer A and loaded on a POROS 20 HE column (Applied Biosystems).  $\Delta 19\omega_2$  was eluted with 50–1000 mM NaCl gradient in buffer A. Concentrated fractions were gel filtrated on Superdex75 (GE Healthcare) run with buffer B [20 mM Tris–HCl (pH 7.5), 300 mM NaCl].

Complementary oligonucleotides were purified by high-performance liquid chromatography (HPLC), mixed at 1:1 molar ratio, hybridized and purified using a MonoQ column

(GE Healthcare). Eluted DNA was dialyzed against 20 mM Tris–HCl (pH 7.5), 100 mM KCl and 30 mM NaCl.  $\Delta 19\omega_2$  was added at 2.1:1 molar ratio and purified by gelfiltration (SuperdexS75). Fractions of  $\Delta 19\omega_2$ /DNA were concentrated to 10.5 mg/ml for crystallization.

### Crystallization

Using hanging-drop vapor diffusion, crystals with space group C2 grew from drops made of 2  $\mu$ l of  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  solution or 2  $\mu$ l of  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  (bp A9–T6' exchanged by bp G9–C6') solution and 2–3  $\mu$ l precipitant solution [150 mM KH<sub>2</sub>PO<sub>4</sub> (pH 7.0), 2.4 M Na<sub>2</sub>-malonate, 2% 6-amino-caproic acid].

Crystals of  $[\Delta 19\omega_2]_2-(\leftarrow\leftarrow)$  grew in space group P2<sub>1</sub> under similar conditions when precipitant solution contained 150 mM Na/KPO<sub>4</sub> (pH 7.0), 2.4 M Na<sub>2</sub>-malonate and 2 to 3% 2-methyl-2,4-pentanediol. In all cases crystal quality was improved by micro-seeding.

### Data collection, structure determination and refinement

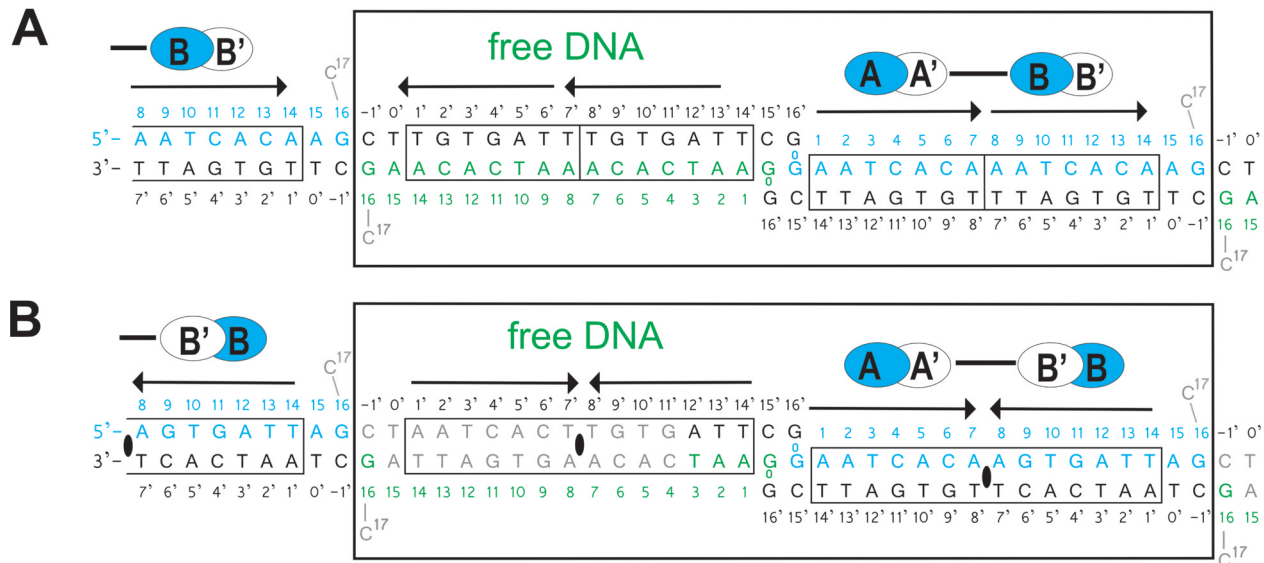
X-ray data were collected at 100 K at the Protein Structure Factory beamline BL14.1 of Free University Berlin at BESSY and processed with HKL2K (12); Table 2.

The structure of  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  was determined by molecular replacement in PHASER (13) with  $\Delta 20\omega_2$  (PDB code 1IRQ) modeled to 8 bp idealized B-DNA. After manual building of  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  and restrained refinement in REFMAC5 (14), F<sub>o</sub>–F<sub>c</sub> maps showed additional electron density for another DNA molecule but not for additional  $\Delta 19\omega_2$ . This second DNA molecule was build manually starting from ideal B-form DNA. In the final model the asymmetric unit consists of one  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  and one free  $(\rightarrow\rightarrow)$ -DNA.  $\Delta 19\omega$  molecules A' and B could be modeled with all residues 19–71, but A and B' only with residues 23–71 and 25–71, respectively, see Figures 1 and 2 for assignment of A, A', B, B'.

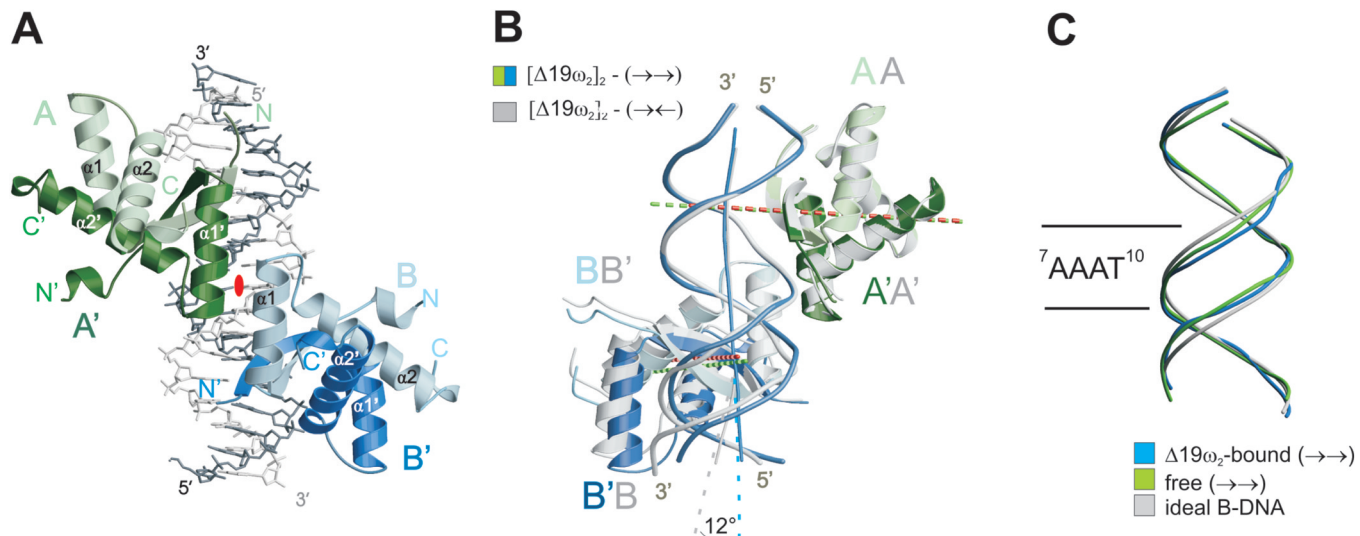
The structure determination of  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  mutant with bp A9–T6' replaced by bp G9–C6' used difference Fourier technique applied to the isomorphous crystal structure of  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$ , see Table 2.

The structure of  $[\Delta 19\omega_2]_2-(\leftarrow\leftarrow)$  was determined in MOLREP (15) using  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  as search model. One  $[\Delta 19\omega_2]_2-(\leftarrow\leftarrow)$  was found, and after restrained refinement sparse electron density indicated only four additional bases for free  $(\leftarrow\leftarrow)$ -DNA that could not be modeled completely, Figure 1B. Molecule B' could be modeled with all residues, B with residues 25–71 and A, A' with residues 22–71.

For refinement of all three structures in Refmac5 TLS groups were assigned and refined for each polypeptide chain and oligonucleotide, see Table 2 for statistics. No non-crystallographic symmetry was used during refinements. Model quality was examined by Whatcheck and Procheck (16) showing that  $\phi$ ,  $\psi$  torsion angles of most amino acids in all three structures are within the most favored, some are in additionally allowed and none are in forbidden areas of the Ramachandran plot. Figures were generated with MOLSCRIPT (17) and Raster3D (18). Analysis of DNA parameters used program Curves (19).



**Figure 1.** DNA used for cocrystallization and arrangement of (A)  $[\Delta 19\omega_2]_2$ - $(\rightarrow \rightarrow)$  and (B)  $[\Delta 19\omega_2]_2$ - $(\rightarrow \leftarrow)$  and free DNAs in the crystal asymmetric units (large boxes). Subunits sharing the same protein-DNA interactions  $\Delta 19\omega_2$ /B shown in blue and  $\Delta 19\omega_2$ /A' in white ovals. Solid bars indicate subunits interacting through helices  $\alpha 1$ ,  $\alpha 1'$ . The pseudo-2-fold axis relating  $\Delta 19\omega_2$  and bound heptads in  $[\Delta 19\omega_2]_2$ - $(\rightarrow \leftarrow)$  indicated by black ellipses in (B). Heptads with sequence 5'-AATCAC<sup>A</sup>/T-3' outlined by arrows, and nucleotides are numbered 0 to 17 and -1' to 16', respectively. Top strands of free DNA in green. Grey nucleotides C17 in both structures and 4 to 15 bp of free DNA in  $[\Delta 19\omega_2]_2$ - $(\rightarrow \leftarrow)$  could not be modeled due to poor electron density.



**Figure 2.** (A) Structure of  $[\Delta 19\omega_2]_2$ - $(\rightarrow \rightarrow)$ . DNA backbone trace in light grey for top strand and dark grey for bottom strand, see Figure 1.  $\Delta 19\omega$  monomers A/A' and B/B' in light and dark green and blue, respectively, helices  $\alpha 1$  and  $\alpha 2$  labeled with white letters. Helices  $\alpha 1'$  of A' and  $\alpha 1$  of B related by a pseudo-2-fold axis perpendicular to the paper plane (red ellipse) form the  $\Delta 19\omega_2$ - $\cdot$  $\Delta 19\omega_2$  interface. (B) Superimposition of  $\Delta 19\omega_2$ A/A' of  $[\Delta 19\omega_2]_2$ - $(\rightarrow \rightarrow)$  (green) and  $\Delta 19\omega_2$ A/A' of  $[\Delta 19\omega_2]_2$ - $(\rightarrow \leftarrow)$  (grey) to show slight positional differences of  $\Delta 19\omega_2$ B/B' associated with palindromic symmetry in  $(\rightarrow \leftarrow)$ . Pseudo-2-fold axes relating monomers in  $\Delta 19\omega_2$  indicated by dashed lines colored green for  $[\Delta 19\omega_2]_2$ - $(\rightarrow \rightarrow)$  and red for  $[\Delta 19\omega_2]_2$ - $(\rightarrow \leftarrow)$ . Helices  $\alpha 1$ ,  $\alpha 1'$  involved in  $\Delta 19\omega_2$ - $\cdot$  $\Delta 19\omega_2$  interactions superimpose well allowing cooperative binding in both complexes. DNA in  $[\Delta 19\omega_2]_2$ - $(\rightarrow \leftarrow)$  is locally kinked by 12° at the centre (bp G11-C4') of heptad A8-T14, and  $\Delta 19\omega_2$  are ~0.6 Å closer (vertical distance between pseudo-2-fold axes) than in  $[\Delta 19\omega_2]_2$ - $(\rightarrow \rightarrow)$ , see text. (C) Phosphate backbone of the 14 bp operator regions of operator DNA free ( $\rightarrow \rightarrow$ ) in green and  $\Delta 19\omega_2$ -bound ( $\rightarrow \rightarrow$ ) in blue superimposed on ideal B-DNA (grey).

## RESULTS

### $\Delta 19\omega_2$ protein

Cocrystallization of wt  $\omega_2$  with operator DNA yielded only crystals with poor X-ray diffraction, but was successful with  $\Delta 19\omega_2$ . *In vitro*,  $\Delta 19\omega_2$  binds specifically to promoter *PcopsS*

with 2-fold lower affinity ( $k_D \sim 12$  nM) (see Supplementary Figure 10) compared to wt  $\omega_2$  ( $k_D \sim 6$  nM) (8), and likewise plasmid-borne  $\Delta 19\omega$  gene product represses *P $\omega$*  utilization *in vivo* 2-fold weaker compared to wt  $\omega$  gene (Table 1). This suggests that even without the N-terminal 19 residues,  $\omega_2$  still binds strongly (only 2-fold weaker) and

specifically to DNA heptads in both, *in vitro* and *in vivo* gene regulation.

### Crystal unit cells contain free and $\Delta 19\omega_2$ bound operator DNA

The crystal structures of  $\Delta 19\omega_2$  bound to two minimal operators formed by 17 bp DNAs with C, G overhangs and comprising direct ( $\rightarrow\rightarrow$ ) and inverted ( $\rightarrow\leftarrow$ ) heptads (Figures 1 and 2) were determined by molecular replacement at 2.45 and 2.6 Å resolution, respectively (Table 2). The asymmetric units of both complexes contain two  $\Delta 19\omega_2$  bound to operator DNAs ( $[\Delta 19\omega_2]_2(\rightarrow\rightarrow)$  and  $[\Delta 19\omega_2]_2(\rightarrow\leftarrow)$ ) which in turn interact with the ends to free operator DNAs ( $\rightarrow\rightarrow$ ) and ( $\rightarrow\leftarrow$ ), respectively, to form pseudo-continuous DNA (Figures 1 and 3).

On the ‘left’ sides of free DNA (Figure 1), nucleotides C17 of free and  $\Delta 19\omega_2$ -bound DNA are not in helical arrangement and not seen in the electron density as they are disordered but were confirmed by MALDI-TOF-spectrometry of dissolved crystals (data not shown). Both,  $[\Delta 19\omega_2]_2$ -bound and free

DNAs, stack with bp G16-C-1' that are related by pseudo-2-fold symmetry.

On the ‘right’ sides of free DNA (Figure 1), the two 3'-G16' overhangs lie in the minor groove of the adjacent duplex and interact with both 5'-G0 to form two consecutive G16'\*(G0-C15') base-triplets (Supplementary Figure 11) with similar geometry as reported (20–22).

In the crystals of  $\Delta 19\omega_2$  bound to ( $\rightarrow\rightarrow$ ) in space group C2 and bound to ( $\rightarrow\leftarrow$ ) in space group P2<sub>1</sub> (Table 2), both  $[\Delta 19\omega_2]_2$ -DNA complexes interact by protein–protein contacts to form layers parallel to the crystallographic *a*, *b* planes. The crystallographic *a*, *b*-axes in space group P2<sub>1</sub> correspond to *b*, *a*-axes in C2 thus reflecting space group and lattice packing similarities. The pseudo-continuous DNA helices (Figures 1 and 3) are oriented in *c*-direction. The doubled *c*-axis in C2 relative to that in P2<sub>1</sub> is due to the C-centering, and the DNA helices are parallel to the *c*-axis at shortest inter-helix distance of ~5 Å in  $[\Delta 19\omega_2]_2(\rightarrow\rightarrow)\equiv(\rightarrow\rightarrow)$  (base-triplets indicated by ≡), possibly stabilized by bridging water. By contrast, in  $[\Delta 19\omega_2]_2(\rightarrow\leftarrow)\equiv(\rightarrow\leftarrow)$  the pseudo-continuous DNA helices are parallel to the crystallographic *a*, *c* plane but inclined at an angle of ~40° towards the *c*-axis and at least ~8 Å apart, and poor electron density (Figure 3B) indicates that they are partially disordered as shown by B factors >100 Å<sup>2</sup>. Consequently, bp 4 to 15 of free ( $\rightarrow\leftarrow$ )-DNA in  $[\Delta 19\omega_2]_2(\rightarrow\leftarrow)\equiv(\rightarrow\leftarrow)$  could not be modeled (Figures 1B and 3B).

In both crystal unit cells (space groups C2 and P2<sub>1</sub>, Table 2), the DNA-bound  $\Delta 19\omega_2$  show minor structural changes compared to the X-ray structure of free  $\Delta 20\omega_2$  (11). This concerns the loop connecting  $\alpha$ -helices  $\alpha 1$  and  $\alpha 2$  [residues 46–48, 2.0 Å root mean square (r.m.s.) deviation for superimposed

**Table 1.** Utilization of  $P\omega: lacZ$  in the presence of  $\omega$  variants in *B. subtilis* cells

Gene provided in trans	$\beta$ -galactosidase activity $P\omega: lacZ$
None	310
pHP14 (control)	298
$\omega$	7
$\Delta 19\omega$	13
$\omega$ Thr29Ala	314
$\omega$ His38Val	52

The  $\beta$ -galactosidase activity is expressed in Miller units.

**Table 2.** Crystallographic data and refinement statistics

	$[\Delta 19\omega_2]_2(\rightarrow\rightarrow)\equiv(\rightarrow\rightarrow)$	$[\Delta 19\omega_2]_2(\rightarrow\leftarrow)\equiv(\rightarrow\leftarrow)$	$[\Delta 19\omega_2]_2(\rightarrow\rightarrow)\equiv(\rightarrow\rightarrow)$ A9-T6' replaced by G9-C6'
Space group	C2	P2 <sub>1</sub>	C2
Unit cell parameters (Å)	44.6, 75.0, 219.4	76.0, 42.5, 103.7	44.6, 76.1, 220.0
$\beta$ (°)	$\beta = 108.8$	$\beta = 107.16$	$\beta = 109.3$
Resolution range (Å)	30.0–2.45	30.0–2.6	30.0–2.9
Observed reflections	83 105	46 759	38 486
Unique reflections	25 244	18 516	14 113
Completeness (%) <sup>a</sup>	97.4 (81.7)	92.8 (72.2)	88.7 (72.1)
$\langle I/\sigma(I) \rangle^a$	12.7 (3.0)	11.7 (3.1)	12.1 (2.5)
R <sub>sym</sub> (%) <sup>a,b</sup>	6.4 (38)	11.3 (29)	10.3 (37)
Refinement statistics			
Total atoms	3189	2609	3134
Solvent atoms	72	51	32
R <sub>work</sub> (%)	22.7	22.5	24.3
R <sub>free</sub> (%)	26.0	25.9	27.4
Root mean square deviations <sup>c</sup>			
Bond lengths (Å)	0.011	0.009	0.014
Bond angles (°)	1.4	1.1	1.5
B factors (Å <sup>2</sup> )	1.6	1.7	1.6
B factors <sup>d</sup>			
$\Delta 19\omega$ A, B, A', B' (Å <sup>2</sup> )	41.6, 43.2, 44.8, 40.2	30.8, 32.3, 33.0, 36.4	38.0, 39.7, 37.6, 43.5
$[\Delta 19\omega_2]_2$ -bound DNA (Å <sup>2</sup> )	44.1, 41.9	34.9, 34.6	41.4, 38.2
Free DNA (Å <sup>2</sup> )	75.8, 82.0	Free DNA incomplete	65.8, 85.3

<sup>a</sup>Values in parentheses refer to the outer resolution shell.

<sup>b</sup>R<sub>sym</sub> =  $(\sum |I_{hkl} - \langle I \rangle|) / (\sum I_{hkl})$ , where  $I_{hkl}$  is the observed intensity and  $\langle I \rangle$  is the average intensity obtained from multiple observations of symmetry-related reflections.

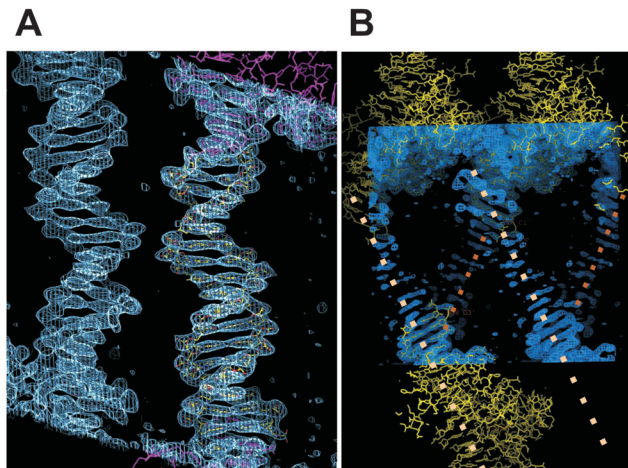
<sup>c</sup>Computed with PROCHECK.

<sup>d</sup>Average B factors calculated for all atoms in each chain.

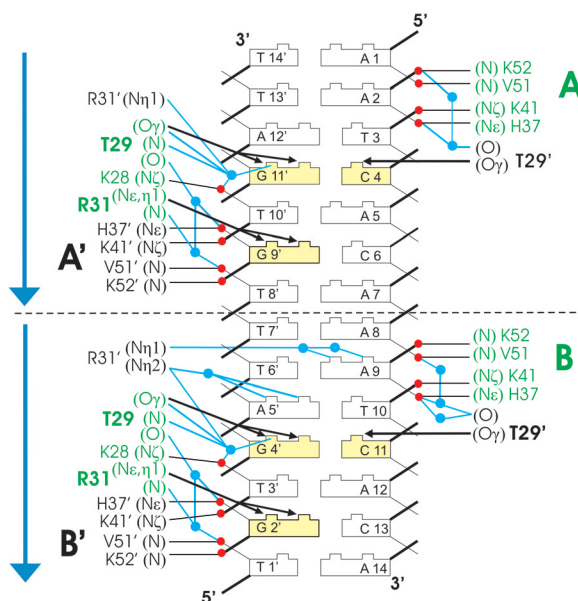
$\alpha$ -Atoms] and to a minor extent the  $\beta$ -sheet (residues 27–32, 0.8 Å r.m.s. deviation).

Since gel permeation chromatography of  $[\Delta 19\omega_2]_2$ -(→→) at 20 mM Tris-HCl (pH 7.5), 150 mM NaCl indicated an apparent molecular mass of 40 kDa (calculated  $M_w = 33.4$  kDa), we assume that under crystallization conditions with 2.4 M Na<sub>2</sub>-malonate, the complexes associate pairwise through base-triplets G16'\*(G0-C15') to form

the complex  $[\Delta 19\omega_2]_2$ -(→→)≡(→→)- $[\Delta 19\omega_2]_2$ . If one  $[\Delta 19\omega_2]_2$ -(→→) of this complex forms layers mediated by protein–protein contacts as in the present structures, it appears that the other  $[\Delta 19\omega_2]_2$ -(→→) has to release  $[\Delta 19\omega_2]_2$  for packing reasons, thereby giving rise to the crystallized  $[\Delta 19\omega_2]_2$ -(→→)≡(→→). The other crystal contact with only DNA–DNA stacking interactions is more flexible than the base-triplets and permits formation of a regular crystal lattice. The same crystallization scenario applies for  $[\Delta 19\omega_2]_2$ -(→←).



**Figure 3.** 2F<sub>0</sub>-F<sub>c</sub> electron density distribution contoured at 1.2  $\sigma$  level; view perpendicular to crystallographic *b*, *c* planes with the *c*-axes vertically oriented. (A) Parallel free DNAs (yellow) in  $[\Delta 19\omega_2]_2$ -(→→)≡(→→) are at ~5 Å shortest distance to each other and stack at both ends with  $\Delta 19\omega_2$ -bound DNAs (magenta). Free DNA is not involved in any further crystal contacts. (B) Discontinuous electron density for free DNA in  $[\Delta 19\omega_2]_2$ -(→←)≡(→←). Modeled complexes of  $[\Delta 19\omega_2]_2$ -(→←) and modeled bases of free DNA (see Figure 1B) in yellow. The central parts of free DNA marked by dotted lines (4 to 15 bp, Figure 1B) could not be modeled due to patchy electron density. The shortest distance between free DNAs is ~8 Å and they are at an angle of ~40°.

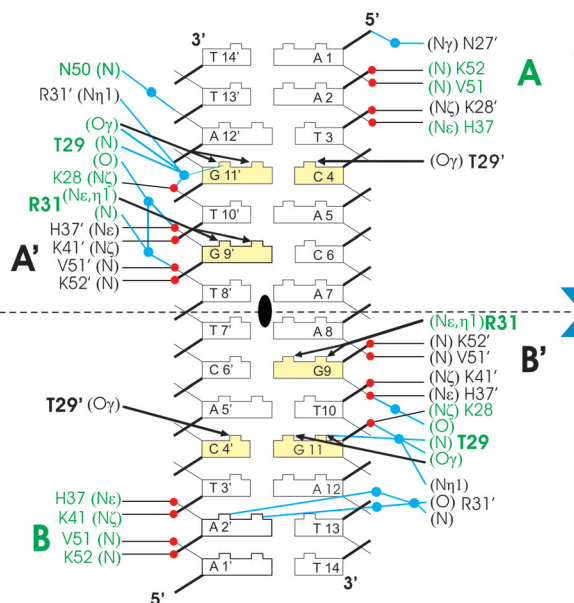


### Protein–DNA interactions

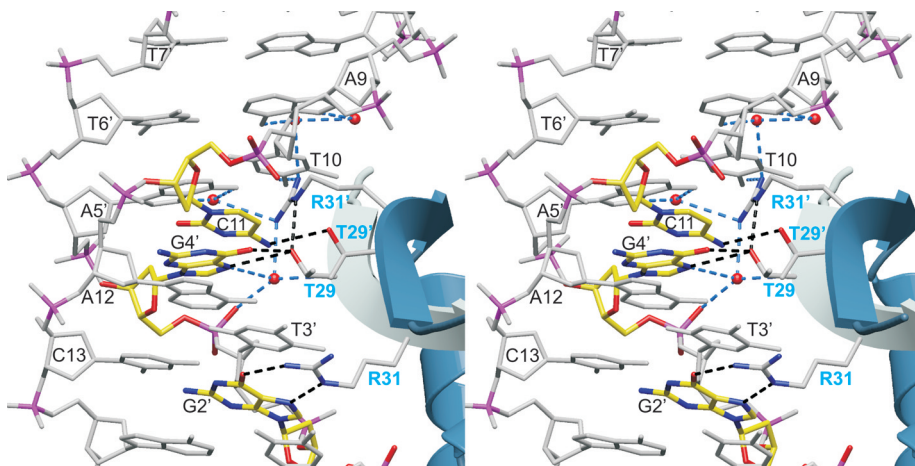
Formation of G16'\*(G0-C15') base-triplets induces distortions at the 5' ends of heptads A1–A7 as indicated by different, partly water-mediated interactions to  $\Delta 19\omega_2$  compared to heptads A8–A14 in (→→) and A8–T14 in (→←), Figures 4 and 5. For this reason, we focus here on protein–DNA interactions for the less distorted dimer  $\Delta 19\omega_2$ B/B' bound to heptad A8–A14 of  $[\Delta 19\omega_2]_2$ -(→→) (Figures 2A and 4(left)). In both structures, for each  $\Delta 19\omega_2$ -bound heptad the direct (not water-mediated) protein–DNA contacts are comparable.

In the major grooves, base pair specific interactions are formed with Thr29 and Arg31 located on the  $\beta$ -sheet. Thr29O $\gamma$  and Thr29'O $\gamma$  of  $\Delta 19\omega_2$ B and B' bind specifically to the central bp G4'-C11, and Arg31N $\epsilon,\eta$  hydrogen bond with base G2' (Figure 4(left)). In contrast, the corresponding Arg31' of  $\Delta 19\omega_2$ B' hydrogen bonds with N $\epsilon'$  to Thr29O $\gamma$  of  $\Delta 19\omega_2$ B and with N $\eta 1',\eta 2'$  through three water molecules to bases G4', A5' and A9, Figures 4 and 5.

To see whether  $\Delta 19\omega_2$  would bind symmetrically to a palindromic heptad featuring two G to provide both, Arg31 and Arg31', with potential binding partners, we determined the 2.9 Å resolution crystal structure of  $[\Delta 19\omega_2]_2$  in complex with a mutated (→→) where bp A9–T6' was replaced by



**Figure 4.** Schematic representation of interactions between  $\Delta 19\omega_2$  and DNA in  $[\Delta 19\omega_2]_2$ -(→→) (left) and  $[\Delta 19\omega_2]_2$ -(→←) (right). The orientation of heptads is indicated by blue arrows and palindromic symmetry by the black ellipse. Yellow bases specifically interact with  $\Delta 19\omega_2$ . Residues labeled green for  $\Delta 19\omega$  A/B and black for  $\Delta 19\omega$  A'/B'. Hydrogen bonds to phosphate oxygens (red) in thin black lines, specific hydrogen bonds to bases in the major groove in black arrows, blue lines indicate water (blue dots) mediated hydrogen bonds. K28 (K28') and Arg31 (Arg31') interact differently in subunits A/B and A'/B'.



**Figure 5.** Stereo view of specific interactions between  $\Delta 19\omega$  B/B' and bound heptad A8–A14 in  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$ . Water molecules; red spheres, hydrogen bonds; dashed lines, phosphorus atoms; magenta, directly contacted bp C11–G4' and base G2' in yellow.

G9–C6' (heptad sequence:  $^8\text{AGTCACA}^{14}$ , Table 2). However, the same interaction pattern was found as in the original heptad. This agrees with similar (2-fold weaker) binding of wt  $\omega_2$  to mutated ( $\rightarrow\rightarrow$ ) with the same replacement in the first heptad ( $^1\text{AGTCACA}^7$ ) compared to the original operator (8). A referee suggested to test the binding of  $\Delta 19\omega_2$  to heptads with pseudo-palindromic symmetry,  $5'\text{-TGTCACA-}3'$ . In view of the binding geometry of  $\Delta 19\omega_2$  to  $^8\text{AGTCACA}^{14}$ , we question whether this would provide novel knowledge. This is because the 5'-AAT- or 5'-AGT-ends of the heptads contact  $\Delta 19\omega_2$  in all cases exclusively through unspecific interactions with phosphate groups or are mediated by water molecules (Figure 4A and B). Hence, base pair exchanges in this part of the heptads should not significantly affect binding of  $\Delta 19\omega_2$ .

Backbone phosphates of all four heptads contact helices  $\alpha 1$  and  $\alpha 2$  of  $\Delta 19\omega$  subunits with pseudo-2-fold symmetry (Figure 4). In  $\Delta 19\omega\text{B}$ , the 5'-phosphate of A9 caps the N-terminus of  $\alpha 2$  by hydrogen bonding to peptide amides of V51 and K52 in a pattern known for RHH<sub>2</sub> proteins and other repressors (4,23), and His37N $\epsilon$ , K41N $\zeta$  of  $\alpha 1$  bind to the 5'-phosphate of T10. Corresponding residues of  $\Delta 19\omega\text{B}'$  interact with 5'-phosphates of G2' and T3'. However, K28N $\zeta$  located on the  $\beta$ -sheet of  $\Delta 19\omega\text{B}$  forms a salt-bridge with the 5'-phosphate of G4' whereas K28'N $\zeta$  on  $\Delta 19\omega\text{B}'$  is  $>5$  Å away from the corresponding 5'-phosphate of C11.

The asymmetry in the binding of each  $\Delta 19\omega_2$  to its particular heptad is reflected by superimposition of C $\alpha$  atoms of monomers (A on A' and B on B', Figure 2A) in each  $\Delta 19\omega_2$  showing 0.6 Å r.m.s. deviation partly associated with structural differences in the loops connecting helices  $\alpha 1$  and  $\alpha 2$  (residues 46–48) and in the  $\beta$ -strands. When  $\Delta 19\omega_2$  dimers are superimposed on each other in the same orientation (A on B and A' on B'), i.e. the bound heptads point in the same direction, r.m.s. deviation of only 0.3 Å confirms that all  $\Delta 19\omega_2$ -heptad interactions are similar.

#### Thr29 is essential for specific operator binding

To test the importance of Thr29 for heptad sequence recognition *in vivo* studies were conducted showing that  $\omega_2\text{Thr29Ala}$  failed to completely repress promoter  $P\omega$

utilization (Table 1). Binding of  $\omega_2\text{Thr29Ala}$  to  $P\omega$  operator DNA embedded in 300 bp DNA was tested by electrophoretic mobility shift assays (EMSA) (Supplementary Figure 10). Protein–DNA complexes formed with  $\Delta 19\omega_2$  but  $\omega_2\text{Thr29Ala}$  required  $\sim$ 100-fold higher concentration (compared to  $\Delta 19\omega_2$ ) that yielded prominent but unspecific binding to  $P\omega$  as confirmed by DNase I footprinting (data not shown).

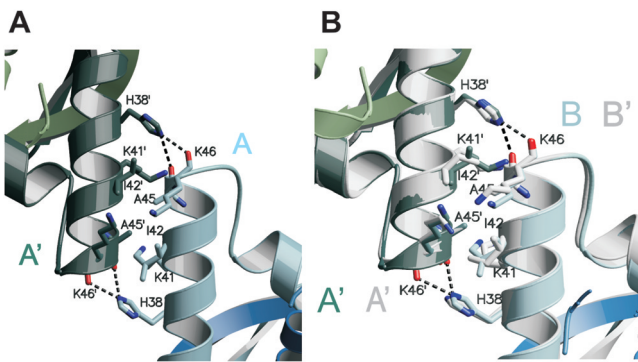
#### $\Delta 19\omega_2\cdots\Delta 19\omega_2$ interactions

Protein–DNA interactions in both complexes bury  $1610 \text{ \AA}^2$  of solvent accessible surface area. Another  $550 \text{ \AA}^2$  are buried by interaction of pseudo-2-fold axis related  $\alpha 1$  helices of adjacent monomers A' and B in  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  and A' and B' in  $[\Delta 19\omega_2]_2-(\leftarrow\leftarrow)$ , respectively (Figure 1 and Figure 2A and B). In detail, bifurcated hydrogen bonds between His38N $\epsilon$  and Ala45' O/Lys46' O are augmented by hydrophobic contacts between Ile42, Ala45 of both subunits (Figure 6). Positions of His38 and His38' are identical in both complexes whereas the hydrophobic side-chain of Ile42 adopts different rotamers without affecting the size of the buried interface. These interactions ensure cooperative binding when several  $\omega_2$  associate with multiple heptad repeats as found in natural operators.

To test the importance of His38, plasmid-borne  $\omega_2\text{His38Val}$  was constructed to remove the bifurcated hydrogen bonds but to maintain the hydrophobic character of the interface.  $\omega_2\text{His38Val}$  repressed  $P\omega$  utilization *in vivo* with  $\sim$ 7-fold lower efficiency than wt  $\omega_2$  (Table 1), indicating the important role of His38 for cooperative binding between  $\omega_2$  and multiple consecutive heptads.

#### Conformation of free and $\Delta 19\omega_2$ -bound DNA

The DNAs in  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  and  $[\Delta 19\omega_2]_2-(\leftarrow\leftarrow)$  are nearly straight B-form with average helical twist of  $36^\circ$  (range  $23^\circ$  to  $43^\circ$ ; Supplementary Figure 12), but show distinct features. The major groove width in  $[\Delta 19\omega_2]_2-(\rightarrow\rightarrow)$  shows strong modulation depending on nucleotide sequence, being  $\sim$ 13 to  $\sim$ 14 Å except for the  $^7\text{AAAT}^{10}$  tract ( $\sim$ 11 to 12 Å) compared to 11.7 Å for ideal B-DNA (24). In contrast, the major groove width in  $[\Delta 19\omega_2]_2-(\leftarrow\leftarrow)$  is more continuous ( $\sim$ 13 to  $\sim$ 14 Å) due to the palindromic symmetry (Figure 7A). In the  $^7\text{AAAT}^{10}$

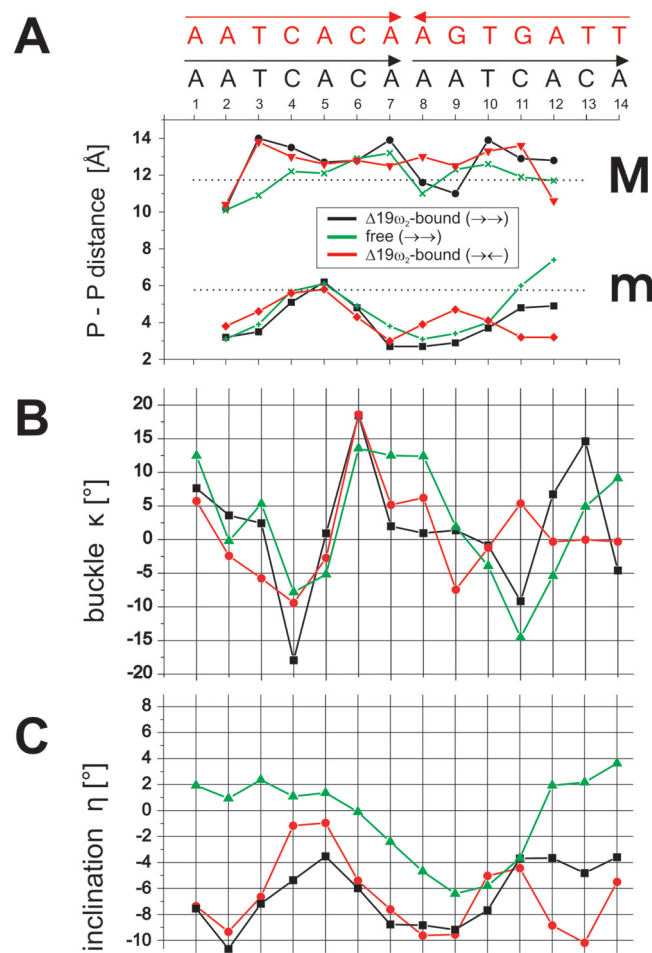


**Figure 6.** Interactions at the  $\Delta 19\omega_2$ ··· $\Delta 19\omega_2$  interface (**A**) between helices  $\alpha 1'$  and  $\alpha 1$  of subunits A' and B of  $[\Delta 19\omega_2]_2$ -(→→), (see Figure 2A). The  $\Delta 19\omega_2$  are related by a pseudo-2-fold axis perpendicular to the paper plane (red ellipse). Residues with contacts  $<4$  Å to neighbor helix are depicted with side-chains, colors as in Figure 2. Bifurcated hydrogen bonds between His38 (His38') and Ala45O/Lys46O to Ala45'O/Lys46'O of the other subunit in dashed lines. (**B**) Superimposition as in Figure 2B with view on the dimer–dimer interface to show that dimer–dimer interactions are comparable in both structures.

tract of  $[\Delta 19\omega_2]_2$ -(→→) and in the  $^7\text{AAGT}^{10}$  tract of  $[\Delta 19\omega_2]_2$ -(→←) the minor groove is narrowed to  $\sim 2.7$  and  $\sim 4$  Å, respectively, compared to 5.7 Å in ideal B-DNA (24) due to an average negative base pair inclination of  $-8^\circ$  (Figure 7C), as frequently found for A-tracts (25). This is associated in  $^7\text{AAAT}^{10}$  of  $[\Delta 19\omega_2]_2$ -(→→) with binding of a single spine of four water molecules, one in each base step as reported in (26). Strong opposite buckles of the C–G bp in the  $^4\text{CAC}^6$  ( $-15^\circ$ ,  $15^\circ$ ) and  $^{11}\text{CAC}^{13}$  ( $-10^\circ$ ,  $15^\circ$ ) segments of  $[\Delta 19\omega_2]_2$ -(→→) (Figure 7B) widen the major groove to  $\sim 13$  Å to accommodate the  $\beta$ -sheet, and adjacent (up- and downstream) minor grooves are narrowed (Figure 7A). Similarly, in  $[\Delta 19\omega_2]_2$ -(→←) the C–G bp in segment  $^4\text{CAC}^6$  show comparable buckles of  $-10^\circ$ ,  $19^\circ$  but in  $^9\text{GTG}^{11}$  the buckles are reduced to  $-8^\circ$ ,  $5^\circ$ . Of the four heptads, three feature two subsequent CA steps that are known for their ability to bend B-DNA through positive roll (20,27,28). However, the CA steps do not exhibit unusual structure except for a moderate slide movement of  $\sim 1$  Å at the central C–G bp where  $\Delta 19\omega_2$  binds to both bases (Supplementary Figure 12).

In the crystal lattice, free (→→)-DNA contacts  $\Delta 19\omega_2$ -bound (→→)-DNA with both ends to form pseudo-continuous helices (Figure 1) and is not distorted by any further crystal contacts (Figure 3A). Interestingly, free (→→)-DNA shows comparable structural features as found for  $\Delta 19\omega_2$ -bound (→→)-DNA (Figure 2C). In the  $^7\text{AAAT}^{10}$  tract of free (→→)-DNA, base pair inclination angles are negative, and the minor groove is narrowed to  $\sim 3.5$  Å whereas the major groove is overall widened to an average of 12.5 Å. Additionally, strong opposite buckles are observed for C–G base pair in both CAC segments that are comparable to  $\Delta 19\omega_2$ -bound (→→)-DNA (Figure 7B). These observations suggest that the described significant deviations from ideal B-DNA are a consequence of nucleotide sequence and not induced by binding to  $\Delta 19\omega_2$  (Figures 2C and 7).

However, conformational changes upon repressor binding concern base pair parameters helical twist, roll and slide (Supplementary Figure 12). In free (→→)-DNA these parameters cluster around values assigned to ideal B-DNA, but



**Figure 7.** Comparison of operator DNA free and in complex with  $\Delta 19\omega_2$ . (**A**) Minor (m) and major (M) groove widths along the central 11 bp of each operator, as measured by shortest P-P distances (less two phosphate group radii, 5.8 Å) indicated by horizontal dotted black lines for ideal B-DNA (24), 5.7 Å and 11.7 Å, respectively. Note minor groove narrowing in the  $^7\text{AAAT}^{10}$  tract of  $\Delta 19\omega_2$ -bound and free (→→) DNA and reduced narrowing in  $^7\text{AAGT}^{10}$  tract of  $(\Delta 19\omega_2)_2$ -(→←). (**B**) Base pair buckle  $\kappa$  ( $^\circ$ ) and (**C**) inclination angles  $\eta$  ( $^\circ$ ) of each operator.

show a negative roll of  $\sim -6^\circ$  and positive slide of  $\sim 1$  Å for the central G11'-C4 and G4'-C11 bp of both heptads, and helical twist angles for bp A5-T10' and A12-T3' decrease to  $\sim 25^\circ$  in  $\Delta 19\omega_2$ -bound (→→)-DNA. Since these deviations are similar for both heptads of  $[\Delta 19\omega_2]_2$ -(→→) they can be attributed to  $\Delta 19\omega_2$  binding. Such comparison is not possible with free (→←)-DNA as it could not be modeled completely.

#### Comparison of $[\Delta 19\omega_2]_2$ -(→→) and $[\Delta 19\omega_2]_2$ -(→←)

The pseudo-2-fold axis relating the monomers in  $\Delta 19\omega_2$  passes with  $\sim 0.3$  Å downstream (5'→3') offset through the central G–C bp of each heptad. Consequently, these symmetry axes are separated by 7 bp in  $[\Delta 19\omega_2]_2$ -(→→) but they are 0.6 Å closer in  $[\Delta 19\omega_2]_2$ -(→←), see Figure 2B. Despite the different dimer–dimer separations helices  $\alpha 1$  and  $\alpha 1'$  forming the  $\Delta 19\omega_2$ .. $\Delta 19\omega_2$  interfaces superimpose well in both complexes, consistent with similar dissociation constants ( $k_D$   $\sim 20$  nM). In contrast, the separation between the two  $\Delta 19\omega_2$  will be  $\sim 0.6$  Å wider in diverging heptads

$[\Delta 19\omega_2]_2$ - $(\leftarrow\rightarrow)$ . Assuming that the interaction pattern between  $\Delta 19\omega_2$  and heptads with  $(\leftarrow\rightarrow)$  orientation is similar as with  $(\rightarrow\rightarrow)$  and  $(\rightarrow\leftarrow)$  heptad orientations, the expected  $\sim 0.6$  Å longer  $\alpha 1\cdots\alpha 1$  contacts are probably less favorable and diminish cooperativity. This agrees with the 6-fold weaker affinity of wt  $\omega_2$  to heptads in  $(\leftarrow\rightarrow)$  arrangement, the finding that  $[\Delta 19\omega_2]_2$ - $(\leftarrow\rightarrow)$  dissociated during gel filtration, and the drastically reduced binding affinity to heptad repeats spaced by one or more additional base pair (8).

Whereas DNA in  $[\Delta 19\omega_2]_2$ - $(\rightarrow\rightarrow)$  is nearly straight, the superimposition in Figure 2B reveals a kink of  $\sim 12^\circ$  at the G11-C4' bp at the centre of the heptad bound to dimer B/B' of  $[\Delta 19\omega_2]_2$ - $(\rightarrow\leftarrow)$ . This kink is associated with  $\sim 12^\circ$  rotation of dimer B/B' of  $[\Delta 19\omega_2]_2$ - $(\rightarrow\leftarrow)$ , the rotational pivot point being located in the  $\Delta 19\omega_2\cdots\Delta 19\omega_2$  interface. By virtue of these two motions, the position and orientation of helix  $\alpha 1'$  of subunit B' remain almost as in  $[\Delta 19\omega_2]_2$ - $(\rightarrow\rightarrow)$  and similar  $\Delta 19\omega_2\cdots\Delta 19\omega_2$  interactions explain comparable dissociation constants of both complexes.

### Structural model of $\omega_2$ -bound to natural operators

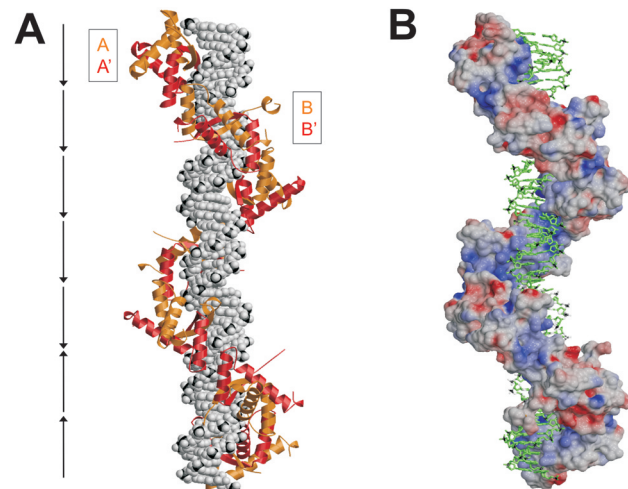
Extrapolation of the structures of  $[\Delta 19\omega_2]_2$ - $(\rightarrow\rightarrow)$  and  $[\Delta 19\omega_2]_2$ - $(\rightarrow\leftarrow)$  allowed modeling of  $\Delta 19\omega_2$  in complex with natural  $P\delta$  promoter (Figure 8). The model implies that wt  $\omega_2$  binds as left-handed matrix to right-handed, straight B-type operator DNA, each  $\Delta 19\omega_2$  being displaced relative to its neighbor by  $\sim 7$  bp and rotated by  $252^\circ$ . Figure 8B shows that the negatively charged sugar-phosphate backbone of DNA faces positively charged surface of  $\Delta 19\omega_2$ .

## DISCUSSION

### Implications of $\Delta 19\omega_2$ -DNA structures for regulation of transcription

The *inc18* family plasmids harbor genes to control their copy number, accurate segregation and stable maintenance during cell division. Since expression of these genes is regulated by the common  $\omega_2$  repressor, a unique mechanism has evolved to fine-tune repressor affinity for the different operators. How this is achieved is shown by the present study. It clearly indicates that the pseudo-symmetric  $\omega_2$  binds with 0.3 Å downstream ( $5'\rightarrow 3'$ ) offset relative to the center G-C base pair of the cognate heptad. Since the operators are nearly straight B-DNA, different heptad numbers and orientations lead to different distances between  $\alpha 1$  helices of adjacent  $\omega_2$ , thereby modulating cooperative interactions between  $\omega_2$  and different operators. The ability to bind to palindromic as well as to non-palindromic operators is a unique feature of  $\omega_2$  and is not shared by other member of the RHH<sub>2</sub> family. We associate this with the interactions between  $\Delta 19\omega_2$  that are related by a pseudo-2-fold rotation axis (Figure 2A and Figure 6A and B) so that they interact comparably, no matter what the orientations of adjacent heptads are,  $(\rightarrow\rightarrow)$ ,  $(\rightarrow\leftarrow)$  or  $(\leftarrow\rightarrow)$ . It is unlikely that the deleted N-termini would contribute to  $\Delta 19\omega_2\cdots\Delta 19\omega_2$  interactions as the N-termini of  $\Delta 19\omega_2$  point away from the  $\Delta 19\omega_2\cdots\Delta 19\omega_2$  interface (Figures 2A and 6A).

When RHH<sub>2</sub> bind to DNA, we have to consider two different binding characteristics, the ‘variable’ binding of flexible side-chains of  $\beta$ -sheets and  $\alpha 1$  helices to major groove and phosphate groups, respectively, and ‘stiff’ capping of N-termini of  $\alpha 2$  helices by rigid main-chain NH hydrogen



**Figure 8.** . Model of seven  $\Delta 19\omega_2$ -bound heptads  $(\rightarrow\rightarrow\rightarrow\rightarrow\rightarrow\leftarrow\leftarrow)$  of the natural promoter  $P\delta$ . (A) DNA in space filling and  $\Delta 19\omega_2$  as orange/red ribbons, (B) DNA in stick and  $\Delta 19\omega_2$  in surface representation colored according to electrostatic potential (negative and positive charges red and blue, respectively). The model is based on the structures of  $[\Delta 19\omega_2]_2$ - $(\rightarrow\rightarrow)$  and  $[\Delta 19\omega_2]_2$ - $(\rightarrow\leftarrow)$ . Repressors form a left-handed protein-matrix winding around the nearly straight operator.

bonding to phosphates. Although Arc<sub>2</sub>, CopG<sub>2</sub> and MetJ<sub>2</sub> bend minimal cognate operators with two repressor binding sites by 50 to 60°, their cores do not rearrange significantly compared to free repressors and are structurally similar [r.m.s. deviations for superimposed C<sub>α</sub> atoms 1.0 to 1.5 Å (3)]. In contrast, Arc<sub>2</sub>, CopG<sub>2</sub> and MetJ<sub>2</sub> superimpose on DNA-bound  $\Delta 19\omega_2$  with higher r.m.s. deviations of 2.3 to 2.8 Å because the  $\beta$ -sheet of  $\Delta 19\omega_2$  protrudes  $\sim 3$  Å less from the repressor surface and is closer to the N-termini of helices  $\alpha 2$  (Supplementary Figure 13). Due to this geometry of  $\Delta 19\omega_2$ , the N-termini of helices  $\alpha 2$  are in the correct position to clamp the phosphate backbones of straight operator DNA when the  $\beta$ -sheet is inserted into the major groove (Figures 2 and 4). In contrast, the other three repressors have to bend DNA around the more protruding  $\beta$ -sheet to place the phosphate backbone in hydrogen bonding distance to N-H groups of the N-termini of their helices  $\alpha 2$ . It is notable that the distances between phosphates bound by  $\alpha 2$  helices is 5 bp in  $\omega_2$  (Figure 4) but 6 bp in the other three RHH<sub>2</sub> since DNA has to follow a longer path when bending around the protruding  $\beta$ -sheet.

### The N-termini of RHH<sub>2</sub> proteins are associated with different functions

The N-terminus in CopG<sub>2</sub> has no obvious function, in MetJ<sub>2</sub> it is involved in binding the corepressor S-adenosylmethionine whereas in free Arc<sub>2</sub> it is disordered but forms a 3<sub>10</sub> helix upon and contributes to DNA-binding. The N-terminal residues in  $\omega_2$  do not contribute to DNA-binding *in vitro* and *in vivo* as shown in the present work. However, we have recently shown that protein  $\delta$ , a homolog to ParA proteins and involved in active plasmid partitioning during cell division, is activated by wt  $\omega_2$  but not by  $\Delta 19\omega_2$  (A. Cicek, F. Pratto, W. Weihofen, J.C. Alonso and W. Saenger, unpublished data). This suggests that  $\omega_2$  is the yet missing ParB protein of *inc18* family plasmids in the known ParA/ParB family of plasmid partitioning systems (29).

## Direct and indirect readout of DNA sequence

Cocrystallized  $\Delta 19\omega_2$ -bound and free ( $\rightarrow\rightarrow$ )-DNA show similar phosphate backbone conformation (Figure 2C) with significant deviations from ideal B-DNA (Figure 7), indicating that conformation is predominantly dependent on DNA sequence and not induced by  $\Delta 19\omega_2$  binding. This provides a good example for the ‘indirect’ readout of local DNA conformation by  $\Delta 19\omega_2$  that depends on the particular nucleotide sequence (30). The combination with ‘direct’ readout resulting from interactions of  $\omega_2$  repressor amino acids with heptad bases increases protein–DNA-binding specificity and affinity. This view is consistent with studies in which the heptad nucleotide sequence was mutated base pair by base pair (8). Mutations at heptad positions 2, 3 and 5 (5'-AATCAC-3') resulted in at least 4-fold weaker binding to wt  $\omega_2$ , although these base pair are not directly contacted by repressor side-chains contrasting direct read out positions 4 and 6 that were much more sensitive against mutations.

## Protein Data Bank accession codes

Atomic coordinates and structure factors have been deposited with the Protein Data Bank under accession codes 2bnw, 2bnz and 2cax.

## SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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*Conflict of interest statement.* None declared.

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