Towards Novel Amino Acid-Base Contacts in Gene Regulatory Proteins: AraR – A Case Study

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Abstract

AraR is a transcription factor involved in the regulation of carbon catabolism in Bacillus subtilis. This regulator belongs to the vast GntR family of helix-turn-helix (HTH) bacterial metabolite-responsive transcription factors. In this study, AraR-DNA specific interactions were analysed by an in vitro missing-contact probing and validated using an in vivo model. We show that amino acid E30 of AraR, a highly conserved residue in GntR regulators, is indirectly responsible for the specificity of amino acid-base contacts, and that by mutating this residue it will be possible to achieve new specificities towards DNA contacts. The results highlight the importance in DNA recognition and binding of highly conserved residues across certain families of transcription factors that are located in the DNA-binding domain but not predicted to specifically contact bases on the DNA. These new findings not only contribute to a more detailed comprehension of AraR-operator interactions, but may also be useful for the establishment of a framework of rules governing protein-DNA recognition.

Introduction

Protein–DNA binding is a process fundamental to life as it masters many genetic activities such as transcription, recombination, DNA replication and repair. The specific interaction between transcription factors and their cognate DNA sites is critical for regulation of gene expression in cells. Understanding how these different proteins are able to find and bind selectively to only one, or just a small number, specific sequence(s) out of the millions of nucleotides present in a genome is a major goal of molecular biology. The recognition principles of protein–DNA interfaces are guided by the complex interplay of noncovalent interactions [1,2,3,4]. In general, DNA recognition follows two paradigms, direct and indirect readout. In the case of direct readout, proteins form contacts such as, hydrogen bonds and van der Waals contacts, mainly in the major, and to a lesser extent also the minor, groove of the DNA to the edges of the base pairs to probe the DNA sequence [1,2,3,4]. Indirect readout occurs through protein contacts to the DNA that depend on base pairs that are not directly contacted by the protein in which the sequence-dependent deformability or structural differences between DNA molecules contribute to their discrimination. A DNA-protein “recognition code”, although of great utility in molecular biology, remains elusive and improbable. While it is clear that a single recognition code does not exist there is some evidence for the existence of a degenerated code whereby one group of bases displays tendency to interact with a certain group of amino acids [4,5,6]. In recent years, researchers have addressed this issue by strengthening a comprehensive framework of the rules governing protein–DNA interactions. Different strategies have been described for the construction of Zinc-fingers (ZFs) and TAL (transcription activator-like) proteins with new binding specificities [7,8]. Nevertheless, there is not a simple one-to-one correspondence between protein and DNA sequences, thus direct readout alone is insufficient to justify the specificities of protein-DNA interactions.

AraR is a homodimeric transcription factor involved in the regulation of carbon catabolism in Bacillus subtilis. The protein displays a chimeric organization, consisting of two functional domains with different phylogenetic origins [9,10]: a small N-terminal DNA-binding domain (DBD) comprising a winged helix–turn–helix (HTH) motif belonging to the GntR family of transcriptional regulators [11] and a larger C-terminal domain homologous to that of the GalR/LacI family of bacterial regulators and sugar-binding proteins [12]. Recently, the three-dimensional crystal structure of the AraR C-terminal domain [13] and the DNA-binding domain [14] were independently solved. AraR typifies one of the GntR-subfamilies of proteins (reviewed in [15]). The GntR superfamily is one of the largest groups of HTH bacterial metabolite-responsive transcription factors (Pham family: PF00392; Prosite Family PS05949) and GntR-like regulators are widespread in bacteria and are known to control many fundamental cellular processes, such as primary metabolism, motility, development, antibiotic production, antibiotic resistance, plasmid transfer and virulence (reviewed in [15]).
The control in gene expression exerted by AraR is modulated by the presence of the inducer L-arabinose. Binding of AraR to L-arabinose leads to induction of expression of the ara regulon (Figure 1), which is composed of at least thirteen genes. The products of these genes include the regulator itself, extracellular and intracellular catabolic enzymes involved in the degradation of arabinose-, galactose- and xylose-containing polysaccharides, uptake of these sugars into the cell and further catabolism of L-arabinose and arabinose oligomers [9,16,17,18]. In the absence of inducer, AraR recognizes and binds at least eight palindromic operator sequences (aka boxes), located in the five known arabinose-inducible promoters (Figure 1). Three of these promoters contain two aka boxes: the promoter of the araABDLMNQP-abfA operon (boxes ORA1 and ORA2), of araE/ORF1 and ORF2 and of abf2 (ORX1 and ORX2). In the cases of the genes araR and abfA, a single box is present (ORR1 and ORR2) (Figure 1). AraR binding to the promoters displaying two boxes is cooperative, requiring in phase and properly spaced operators, and involves the formation of a small loop in the DNA. These two mechanistically diverse modes of action of AraR result in distinct levels of transcriptional regulation, as cooperative binding to two aka boxes results in a high level of repression while interaction with a single operator allows a more flexible control [10,18,19].

Previous studies have mapped the functional domains of AraR and characterized the C-terminal region involved in effector binding and dimerization [20]. Moreover, guided by molecular modelling we identified amino acids potentially involved in DNA binding and the effect of their substitution revealed key residues necessary for the DNA binding and regulatory activity in vivo and in vitro [21]. In addition, important bases for AraR-DNA interactions in both arms of the palindromic operator sequences were also identified [21]. In this work we studied AraR-DNA specific interactions using methodologies designed to detect direct or indirect interactions between the atoms/residues of the interacting partners, both in vitro and in vivo. AraR mutant proteins displaying a moderate effect in AraR-DNA interaction and single point mutations in the operator DNA leading to partial derepression of gene expression were probed. The results obtained provide valuable information concerning the specific interaction of AraR-DNA and insights into the binding of GntR regulators in general.

Materials and Methods

Strains and growth conditions

Escherichia coli DH5α (Gibco BRL) was used as host for routine molecular cloning work. E. coli strains were grown in LB [22] medium and the antibiotics ampicillin (100 µg ml⁻¹) and tetracycline (12 µg ml⁻¹) were added when appropriated. B. subtilis strains used in this study (Table 1) were grown in liquid LB or C-minimal medium [23] and chloramphenicol (5 mg ml⁻¹), kanamycin (10 mg ml⁻¹) or erythromycin (1 mg ml⁻¹) were added when appropriated. The B. subtilis and E. coli cells were transformed as described previously [7]. The Amy phenotype was tested by detection of starch hydrolysis on tryptose blood agar base medium (Difco) plates, containing 1% (w/v) of potato starch, with an I₂–KI solution as described previously [9]. The Thr phenotype was determined by growth on Spizizen minimal medium [24] supplemented with 2% (w/v) of glucose, 0.2% (w/v) potassium glutamate, 3 mM MgSO₄, and 2% (w/v) agar.

DNA manipulation and construction of plasmids

DNA manipulations were carried out as described by Sambrook et al. [25]. Restriction enzymes were purchased from MBI Fermentas and used according to the manufacturer’s instructions. DNA was eluted from agarose gels with GFX gel band purification kit (Amersham Pharmacia Biotech). DNA sequencing was performed with ABI PRIS BigDye Terminator Ready Reaction Cycle Sequencing kit (Applied Biosystems). PCR amplifications were done using high-fidelity Phusion DNA polymerase (Finnzymes) and the resulting products purified by QIAquick PCR purification kit (Qiagen). For the construction of plasmids pMI35 and pMI36, bearing substitutions E30A and Y5F, respectively, the mutated araR alleles were amplified by PCR with primers ARA1 and ARA73 (Table 2), using as template chromosomal DNA from strains IQB568 and IQB571 [21], respectively. The PCR products were digested with EcoRI-BamHI (or EcoRI-BglII) and independently IQB568 and IQB571 [21], respectively. The PCR products were subcloned into the respective pLS30 sites [20]. The obtained

![Figure 1](https://example.com/figure1.png)

Figure 1. The arabinose (ara) regulon comprises thirteen genes located in three different regions of the chromosome. The genes are represented as black arrows pointing at the direction of transcription. The AraR repressor, in the absence of the effector molecule - arabinose - binds to palindromic sequences (AtT/AktGaCGTACaaA/TtT consensus depicted, bottom left) found in the promoter region of the ara genes. The AraR protein is shown as a dimer. The eight AraR boxes are represented as white rectangles. Binding to the different operators may either be cooperative or uncooperative.

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plasmids were then digested with ScaI, which allows the occurrence of a double crossover recombination event at \textit{amyE} locus of the \textit{B. subtilis} chromosome (Table 1).

| Strain  | Genotype                        | Source | Details |
|---------|---------------------------------|--------|---------|
| 168T    | Prototroph                      | F. E. Young | [9] |
| IQB 215 | \textit{ΔaraR}:k1m}           |        |         |
| IQB 568 | \textit{ΔaraR}:k1m} \textit{araAB}:lacZ \textit{erm}:amyE::\textit{araR} E30A cat | [21] |         |
| IQB 571 | \textit{ΔaraR}:k1m} \textit{araAB}:lacZ \textit{erm}:amyE::\textit{araR} Y5F cat | [21] |         |
| IQB 761 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} cat | pLS30—IQB215 |         |
| IQB 778 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} E30A cat | pM135—IQB215 |         |
| IQB 774 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} YSF cat | pM136—IQB215 |         |
| IQB 771 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM137—IQB761 |         |
| IQB 790 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM148—IQB761 |         |
| IQB 772 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} YSF cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM146—IQB761 |         |
| IQB 773 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} YSF cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM146—IQB761 |         |
| IQB 779 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} E30A cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM137—IQB787 |         |
| IQB 798 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} E30A cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM148—IQB787 |         |
| IQB 796 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} E30A cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM145—IQB787 |         |
| IQB 797 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} E30A cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM146—IQB787 |         |
| IQB 795 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} YSF cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM137—IQB774 |         |
| IQB 791 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} YSF cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM148—IQB774 |         |
| IQB 792 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} YSF cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM145—IQB774 |         |
| IQB 793 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} YSF cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM146—IQB774 |         |
| IQB 927 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM164—IQB787 |         |
| IQB 928 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM164—IQB787 |         |
| IQB 929 | \textit{ΔaraR}:k1m} \textit{amyE::\textit{araR}} cat · \textit{thrC::ōx1 T6} wat-lacZ \textit{erm} | pM163—IQB787 |         |

*The arrows indicate transformation and point from donor DNA to recipient strain.

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1Transformation was carried out with linearized DNA.

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Table 2. Oligonucleotides used in this work.

| Primer | Sequences (5' → 3') | Complementary sequence |
|--------|----------------------|------------------------|
| ARA1   | (−39) TAAGGTAACATTATTGGCCG (−22) | pSN32 (fwd) |
| ARA73  | (+77) CCTCACCAGTAGTGTACCC (+60) | pSN32 (rev) |
| ARA87  | (−207) AAAATACCGATTACGGCATCG (−186) | abD (fwd) |
| ARA262 | (−37) GATGACAAGCATATATTAGCAATATT (−13) | araABDLMNQP-abfA |
| ARA263 | (+90) CCCCCCTCTATAAGAAATCCGCC (+68) | araABDLMNQP-abfA |
| ARA542 | (−75) TAAATACAGCTACAATAAT (−54) | ORX1 T6→G (fwd) |
| ARA541 | (−54) ATATTTGACGTTCTGATTTA (−75) | ORX1 T6→G (rev) |

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β-Galactosidase assays

*B. subtilis* strains were grown in C-minimal medium supplemented with 1% (w/v) casein hydrolysate in the presence and absence of 0.4% (w/v) L-arabinose, as previously reported [9]. Samples of cell culture were collected and analysed 2 h after the addition of L-arabinose, β-Galactosidase activity was measured using the substrate *p*-nitrophenyl-β-D-galactoside (ONPG) and expressed in Miller units, the ratio of β-galactosidase activity in the presence and absence of inducer was taken as a measure of AraR repression in the analysed strains (Repression Index) as described previously [9].

Electrophoretic mobility shift assay (EMSA)

DNA fragments carrying the operator sequences ORA1A2, wild-type and mutants ORA1 A1→C, G1→T, C16→G, and T16→G were amplified by PCR, with primers 3A262 and 3A263, using plasmids pLM51, pLM61, pLM62 and pLM50 [21], respectively, as template. Overexpression and protein purification of the AraR wild-type and mutant variants (Y5F and E30A) were performed as described previously [20].

The assays were performed as described in Franco et al. [21]. DNA fragments were radiolabelled with [γ-32P] dATP using T4 Polynucleotide Kinase. The protein-DNA binding reaction was carried out in a volume of 10 μl containing 12.5 mM HEPES-KOH pH 7.6, 10 mM MgCl2, 0.5% (w/v) BSA, 1 mM DTT, 10% glycerol (v/v), 200 mM NaCl, 4 mM Na2HPO4, 4 mM NaH2PO4, 0.4 mM EDTA, a 200-fold molar excess of competitor DNA (polydIdC), 1 nM of labelled DNA and increasing concentrations of wild-type or mutant AraR proteins, and incubated at room temperature for 30 min. The reaction mixtures were then submitted to electrophoresis on a native 8% polyacrylamide gel containing Tris-glycine buffer (25 mM Tris, 200 mM glycine, pH 8.9) and run at 100 V for ~1 h. Gels were vacuum dried and exposed on a Phosphorimager screen before analysis with a Molecular Dynamics Storm 860 Imager and ImageQuant version 5.0.

The determination of the dissociation constants, K_d values, was obtained using the GraphPad Prism software and the “one site total binding” model, following the equation Y = Bmax X/K_d[X]+ NS X, with X = AraR concentration, Y = bound protein, Bmax is the maximum specific binding and NS is the slope of nonspecific binding. Concentrations of AraR were determined assuming a pure dimeric protein. Differences between K_d were analyzed by Mann Whitney U test using SPSS software, P<0.05 was considered as the level of statistical significance. The value 0.057 (Table 3) was considered moderate evidence against the null hypothesis [H0: On average there is no difference in binding affinity of the two DNA fragments (mutant DNA fragment vs wild-type DNA fragment)]. The association constant (K_assoc) is calculated from K_d = 1/K_assoc and the Gibbs free energy (ΔG°) by ΔG° = −RT ln K_assoc.

Results

Probing amino acid-base contacts *in vitro*

In a previous study aimed at understanding the specific properties of the interaction AraR-operator sequences, we substituted amino acids, in or near the winged-HTH motif, which according to the model were predicted to contact DNA [20,21], and the effects of these substitutions on the ability of AraR to function *in vivo* and on the DNA-binding affinities *in vitro* were determined [20,21]. Conversely, mutational analysis of the AraR-binding sites was used to determine the base-specific requirements for transcriptional regulation *in vivo* and DNA binding *in vitro*. These experiments showed that specific AraR residues and operator bases are crucial to achieve a high level of regulatory activity, while others display variable contributions to DNA binding. In order to characterize in detail the AraR-DNA specific interaction we used the loss-of-contact approach [27]. In this study we initially used an *in vitro* missing-contact probing [28,29] using electrophoretic mobility shift assay (EMSA) to determine the binding affinities of AraR and mutant proteins to a DNA fragment bearing the promoter of the metabolic operon with two operators (ORA1-ORA2) and the same fragment comprising single base pair substitutions in the ORA1 box (AATTGTTCTGTACAAAAT). The rationale of these experiments was as following: a certain amino acid alteration leads to an increase in K_d for the wild-type operator (Figure 2A); if this increment is the consequence of a lost direct or indirect interaction between that particular amino acid and a specific base, when we use a DNA fragment with a substitution in that particular base we expect no major effect in the K_d, when compared to the wild-type DNA, because a particular contact had already been lost and quantified (Figure 2B); in contrast, if the amino acid exchanged is not involved in contacts with the specific mutated base we will expect an additional increase in K_d (Figure 2C).

This methodology, in addition to indicating residues directly involved in contacts with bases may also reveal amino acids whose presence is important to maintain the overall structural arrangement of the protein even though they do not directly contact bases in the DNA. For the experiments we chose AraR mutant proteins, AraR Y5F and E30A, which displayed a moderate effect in AraR-DNA interaction both *in vivo* and *in vitro*, and base pair substitutions leading to partial derepression *in vivo*, A1→C, G1→T, T16→G and T16→G and T16→G [21]. The results of the EMSA are summarized in Figure 3 and the calculated K_d values are shown in Table 3. The AraR wild-type protein showed a statistical significant decrease in the affinities for a DNA fragment bearing the promoter of the metabolic operon with two operators (ORA1- ORA2), when we compared the wild-type DNA fragment to the same fragment mutated in the ORA1 box. Previously, we have shown that binding of AraR to ORA1-ORA2 is cooperative and a single point mutation in either ORA1 and ORA2 causes an almost complete loss of AraR regulation in vivo [10,19]. Similarly, in vitro a single-point mutation in ORA1 reduces dramatically the apparent affinity of AraR for the second operator ORA2 [10].

The AraR E30A protein displayed a decrease in the affinity for all mutated operators except for the T16→G operator (Table 3). In fact, AraR E30A showed no additional significant decrease in the affinity, relative to the wild-type operator, when the T16→G operator mutant was used (Figure 3 and Table 3). As T6 in ORA1 is important for protein binding [21], and the T16→G mutation did not reduce the binding affinity of AraR E30A, this suggests that this operator substitution did not further affect the loss of contact of AraR E30A. The K_d of the mutant AraR Y5F for the operator mutations tested revealed a significant a reduction in the affinity compared to the wild-type for G1→T and T6→G, but not for A1→C or T16→G (Figure 3 and Table 3). This could indicate that Y5 might be relevant for the contact of AraR with T16 and A1 of ORA1. Because these nucleotides are located in opposite positions in the palindromic sequence of the operator, this observation suggests that Y5 of one monomer is important for the interaction with A1, while the other contacts T16. However, the crystal
Table 3. Thermodynamic parameters of AraR-DNA interaction reactions.

| Protein  | Dissociation constant $K_d$ ($\times 10^{-8}$ M)$^a$ | DNA fragment araABDLMNPO-abfA promoter | $\Delta G^c$ of the AraR-DNA association reactions$^d$ |
|----------|--------------------------------------------------|-----------------------------------|-----------------------------------------------|
|          | $OR_A$ WT | $OR_A, T_6\rightarrow G$ | $p^b$ | $OR_A, T_6\rightarrow G$ | $p^b$ | $OR_A, A_7\rightarrow C$ | $p^b$ | $OR_A, G_5\rightarrow T$ | $p^b$ |
| AraR WT  | 2.7±0.5   | 5.6±0.4           | 0.016 | 11.0±3.3                  | 0.016 | 7.4±1.6                  | 0.038 | 34.5±9.5                  | 0.008 |
| AraR E30A| 19.8±5.3  | 25.4±6.7          | 0.412 | 34.7±6.4                  | 0.057 | 67.5±12.9                | 0.029 | 74.8±18.7                 | 0.057 |
| AraR Y5F | 27.4±3.2  | 47.6±9.0          | 0.016 | 35.5±11.1                 | 0.214 | 33.1±7.6                 | 0.343 | >200                      | 0.016 |

$^a$Dissociation constant ($K_d$) of binding of AraR and AraR mutants (Y5F, E30A) to a wild-type araABDLMNPO-abfA promoter (WT) and mutants ($OR_A, T_6\rightarrow G$, $T_6\rightarrow G$, $A_7\rightarrow C$, $G_5\rightarrow T$) calculated by densitometric quantification of the bands corresponding to free DNA and protein-DNA complex by EMSA (see Materials and Methods). The values represent the average and standard deviation of at least three (three to seven) independent assays, with an intrinsic error <31%.

$^b$p-value (Mann Whitney U test) for each pairwise comparison, mutant DNA fragment vs wild-type DNA fragment. Mutated operators that did not show statistical significant variation ($p>0.05$) when compared to the wild-type operator are highlighted in bold.

$^c$Gibbs free energy is calculated from the equilibrium association constant $K_d$ (see Materials and Methods).

$^d$Variation of Gibbs free energy = $\Delta G^c$ mutant -- $\Delta G^c$ wild-type. ND = not determined.

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structure of the AraR-DNA binding domain bound to ORA1 [14] showed Y5 interacting with the DNA backbone near nucleotide T6 (see below).

In summary, the results obtained in vitro suggest that AraR residue E30 may play an important role in the interaction of the protein with the T6 nucleotide.

In vivo validation of protein-DNA interactions

Since the experimental conditions used to derive Kd values bear little resemblance to intracellular situations, the in vitro results were confirmed by in vivo assays. For this, we constructed B. subtilis strains in order to confront the different araR alleles and mutant DNA operator sequences in the same cell. The different araR alleles were ectopically introduced at the amyE locus of an araR null mutant background. Additionally, a transcriptional fusion between the araA promoter, carrying the ORA1-ORA2 operators, and the E. coli lacZ gene, was generated and ectopically introduced at the B. subtilis thrC locus (Figure 4). This genetic system allows us to measure the regulatory activity of the native and mutant proteins over distinct promoters (wild-type and mutated) fused to the lacZ reporter gene by determination of the levels of accumulated β-galactosidase. In previous studies we have shown that in these conditions the cellular level of both mutant proteins AraR E30A and AraR Y5F is comparable to that seen with wild-type AraR, ruling out the possibility of deregulation originated by degradation of the repressor [21]. The results of the confrontation of the different araR alleles and the various promoters in the series of strains constructed are summarized in Table 4.

The analysis of repression index of the wild-type AraR with the different promoter fragments showed a decrease in the regulatory activity when a mutated box ORA1 was used, compared to the wild-type ORA1-ORA2. The mutation ORA1 T16→G displayed the higher deregulation, while ORA1 A1→G and T6→G exhibited similar less drastic effects. These results are comparable to those obtained in the in vitro assays (Table 3). The dissociation constant of the mutant Y5F suggested that this amino acid might interact with two nucleotides in the operator sequence, T16 and A1 (Table 3). However, the in vivo analysis does not corroborate the hypothesis (Table 4), as mutations at position T16 and A1 have a drastic effect in the regulatory activity of mutant Y5F (IQB792 and IQB793; Table 4). The in vivo results are in agreement with the crystal structure of the AraR-DNA binding domain bound to ORA1 [14] that revealed Y5 interacting with the DNA backbone near nucleotide T6, thus this residue is not involved in direct or indirect contact with T16 and A1 (discussed below).

The EMSA assays indicated that residue E30 could be relevant for the interaction of the AraR protein with the T6 nucleotide (Table 3), although both the N-terminal AraR model [21] and the N-terminal AraR-ORA1 structure [14] suggest non-specific contacts of E30 to the DNA backbone (discussed below). This observation was supported by the in vivo data because the regulatory activity of mutant AraR E30A over the mutant ORA1 T6→G-lacZ promoter fusion is 2.7-fold higher (strain IQB790, Table 4) than that observed for the wild-type promoter ORA1 WT-lacZ (strain IQB779, Table 4). Furthermore, the lower level of expression observed in the strain bearing the mutant AraR E30A and the mutant ORA1 T6→G-lacZ promoter fusion (strain IQB790, Table 4), both in the presence and absence of inducer, compared to that obtained in the strain harbouring the wild-type AraR regulator and the mutant ORA1 T6→G-lacZ promoter fusion (strain IQB790, Table 4) suggests a stronger interaction of the E30A protein towards the mutated DNA promoter.

Overall the in vivo results highlight the importance of amino acid E30 in the regulatory activity AraR and in the contact of the protein with the nucleotide T6 in ORA1.

Residue E30 is important for the AraR regulatory activity in distinct promoters

As T6 is a well-conserved nucleotide in the consensus signature of the AraR DNA binding site, present in all AraR operators characterized so far (Figure 1), to establish that E30 is an important amino acid for the AraR contact to the thymine at position 6 we assayed this effect in the context of a different promoter. The abf2 gene is regulated by cooperative binding of AraR to two in-phase operators ORX1-ORA1 similarly to that observed in the arabinose metabolic operon promoter [18]; Figure 1). Thus, using the same strategy the wild-type ORX1 (ATACATACCGTACAAAT) and mutant ORX1 T6→G abf2-lacZ fusions were constructed and introduced at the B. subtilis thrC locus.

The analysis of the regulatory index exerted by the native AraR in the strain IQB927 showed no effect of ORX1 T6→G mutation when compared to the wild-type promoter (strain IQB926, Table 5). On the other hand, mutant AraR E30A leads to a complete loss of the regulation of the wild-type abf2-lacZ promoter.
promoter fusion *abf2*, showing once again the importance of this amino acid in the regulatory mechanism of this transcription factor. Nevertheless, the confrontation of the mutant E30A with mutation T6R (strain IQB929, Table 5) leads to an increase in the regulatory activity when compared to the wild-type promoter (strain IQB928, Table 5). Therefore, the T6R single nucleotide change partially suppresses the loss of regulation caused by the E30A amino acid substitution pointing out that E30 is an important amino acid for the AraR contact to the thymine at position 6 of both operator sequences OR\textsubscript{A1} and OR\textsubscript{X1}.

**Figure 3. Analysis of operator mutations on AraR–DNA affinity in vitro by EMSA.** AraR wild type left column; AraR E30A middle column; and AraR Y5F mutant right column. The indicated amounts of AraR protein were used in the binding reactions, AraR was incubated with the 5'-end labelled probe (1 nM) bearing the wild-type or mutant operators OR\textsubscript{A1}/OR\textsubscript{A2} and the protein-DNA complexes were resolved on native 8% polyacrylamide gels. The mutation in each DNA fragment is depicted. doi:10.1371/journal.pone.0111802.g003

**Discussion**

The sequence-specificity of DNA recognition by proteins should be viewed in a complete framework. At the atomic level the specificity of DNA-binding proteins is mainly accomplished through direct hydrogen bond and hydrophobic interactions between specific amino acid side chains and functional groups of nucleotide bases in the major and minor groove [1,2,3,4,30]. Nevertheless these direct or water-mediated hydrogen bonds are insufficient to completely explain the specificity of many DNA-
binding proteins. In addition to the chemical complementarity between protein and DNA atoms, it is required a structural complementarity along the networking surfaces of the protein and DNA molecules [31]. The use of genetic methods to identify amino acid base pair contacts in a specific protein-DNA complex is a complementary approach to the X-ray diffraction and to two-dimensional nuclear magnetic resonance spectroscopic (2D NMR) analyses. Furthermore, the construction and analysis of single amino acid substitutions is the only method to determine the apparent binding free energy contribution and the apparent specificity free energy contribution of an amino acid-base pair contact [27 and references therein].

![Figure 4. Genetic organization of the reporter B. subtilis strains. The circle illustrates the B. subtilis chromosome and the location of the amYE, araE/araR, and thrC loci indicated in degrees. The construction containing the wild-type or mutant araR alleles placed at the amYE locus is represented in the top left. The araR-null genetic background is depicted in middle left. The regulatory activity exerted by the araR alleles over the wild-type or mutant araA promoter sequences is measured by a promoter lacZ fusion placed at the thrC locus (bottom left).](image)

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**Table 4.** Regulatory activity of the wild-type AraR protein and mutants E30 and Y5 over an araA-lacZ promoter fusion (wild-type and mutated variants).

| araR allele | Strain | araA'-lacZ | −ara | +ara | R.I. |
|------------|--------|------------|------|------|------|
| araRwt     | IQ8771 | OR_{A1A2} wt | 17.5±2.2 | 1032.4±134.3 | 59.4±7.4 |
|            | IQ8790 | OR_{A1}, T_{P}→G | 99.5±12.8 | 1308.7±113.2 | 13.4±1.7 |
|            | IQ8772 | OR_{A1}, T_{P}→G | 453.9±15.3 | 1815.7±133.2 | 4.0±0.2 |
|            | IQ8773 | OR_{A1}, A_{P}→C | 84.4±5.37 | 1205.6±72.9 | 14.3±0.0 |
| E30A       | IQ8779 | OR_{A1A2} wt | 431.8±38.2 | 1490.9±104.3 | 3.5±0.1 |
|            | IQ8798 | OR_{A1}, T_{P}→G | 82.3±7.1 | 773.8±74.5 | 9.4±0.8 |
|            | IQ8796 | OR_{A1}, T_{P}→G | 1583.2±61.8 | 1429.2±170.9 | 0.9±0.1 |
|            | IQ8797 | OR_{A1}, A_{P}→C | 1413.0±189.9 | 1429.3±187.3 | 1.0±0.0 |
| Y5F        | IQ8775 | OR_{A1A2} wt | 172.8±0.56 | 1462.9±33.5 | 8.5±0.2 |
|            | IQ8791 | OR_{A1}, T_{P}→G | 1670.2±24.2 | 1409.2±34.1 | 0.8±0.0 |
|            | IQ8792 | OR_{A1}, T_{P}→G | 1890.7±63.5 | 1454.7±112.0 | 0.8±0.1 |
|            | IQ8793 | OR_{A1}, A_{P}→C | 1362.4±30.3 | 1497.3±121.7 | 1.1±0.1 |

^{a}_β\text{-Galactosidase activities of the B. subtilis strains grown in the absence (−ara) or presence (+ara) of arabinose. Values represent the average and standard deviation of at least three independent experiments, each assayed in duplicate. MU Miller units.}

^{b}R.I. (Repression Index) indicates the regulatory activity, calculated as the ratio between values obtained in the presence and in the absence of inducer.

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The GntR family members, in general, possess a DNA binding at the N-terminus of the protein and an effector-binding and/or oligomerisation domain at the C-terminus (Pfam family: PF00392; Prosite Family PS03949; [15]). The DNA-binding domain is conserved throughout the GntR family, consisting of a 3-helical bundle core with a small beta-sheet (wing), winged-HTH motif. Despite the vast number of GntR family members sequences deposited in databases there are only a few crystal structures available to examine in detail structure/function relationships. AraR is a transcription factor that typifies one of the sub-families of the GntR group, and recently the three-dimensional crystal structure of the AraR C-terminal domain [13] and the DNA-binding domain [14] were separately and independently determined. In this work, AraR was used to characterize specific interactions with the DNA by an in vitro missing-contact probing and posterior validation in vivo. In the in vitro fragment the results obtained in vitro with the AraR wild-type protein correlate well with those previously obtained in in vivo experiments [19], except for the mutation G52→T that showed a more accentuated decrease in the affinity measured in vitro than the loss of regulation observed in vivo [21]. Moreover, the data obtained in vitro in this study with the AraR wild-type protein are consistent with those previously observed in vivo using a different genetic system [21]. Although, The in vitro EMSA analysis using AraR mutant Y5F and the different DNA fragments bearing point mutations in the ORA1 operator suggested that residue Y5 could be important for protein contacts with two nucleotides in opposite sites of the operator palindromic sequence, T16 and A1 (Table 3), however the in vivo results do not corroborate this hypothesis (Table 4). The in vivo results validate the data of the crystal interaction studies with mutant T6 [14,21]; and Figure 5A). The core of HTH motif is comprised by two α-helices, H2 and H3, spaced by a short four-residues turn (T) in between. In AraR E30 belongs to H2, the stabilizing helix, while R41 and 45 to H3, the recognition helix. The angle between H2 and H3 is typically of 120°, however it can vary between 100° and 150° [34]. Since E30 interacts with R41 and R45, this interaction is crucial to settle the geometry and spatial arrangement of H2 and H3, and protein docking on DNA by the recognition helix, H3 (Figure 5A and B). The role of the E30 is not only the interaction with the DNA but is also to limit the rotation of the recognition helix. In the E30A mutant, R41 and R45 are no longer interacting with E30, moreover this alanine substitution impairs the contacts of this residue with the DNA backbone (Figure 5C). As a result, the regulatory activity of the mutant protein decreases in the presence of the wild-type ara operon promoter, which does not occur in the presence of mutant ORA1 T6→G promoter as a consequence of a spatial orientation of H2 and H3 (Table 4). On the other hand, enrichment of the operator DNA with another guanine, T6→G, could lead to a significant alteration in DNA conformation. In fact, the exocyclic 2-amino groups of the guanines are crucial elements in DNA structure and recognition, as they are known to exert a substantial influence on DNA bending, flexibility and intrinsic curvature [35,36,37,38]. Therefore if the functional groups in the protein do not correctly juxtapose with those in the DNA, protein-DNA complex stability is impaired, which seems to be the case of the wild-type AraR interaction with the mutated operator T6→G. An amino acid not directly involved in contacts with bases, such as

### Table 5. Regulatory activity of the wild-type AraR protein and mutant E30A over an abf2-lacZ promoter fusion (wild-type and mutated variant).

| araR allele | Strain    | abf2-lacZ | −ara | +ara | R.I. |
|------------|-----------|-----------|------|------|------|
| araR wt    | IQB926    | ORX12, wt | 7.2±1.1 | 119.4±16.6 | 16.7±0.2 |
|            | IQB927    | ORX12 T6→G | 27.2±3.08 | 450.4±66.9 | 16.6±1.7 |
| E30A       | IQB928    | ORX12, wt | 155.8±12.9 | 133.3±14.7 | 0.9±0.1 |
|            | IQB929    | ORX12 T6→G | 148.2±21.8 | 336.7±40.5 | 2.3±0.2 |

*β-Galactosidase activities of the B. subtilis strains grown in the absence (−ara) or presence (+ara) of arabinose. Values represent the average and standard deviation of at least three independent experiments, each assayed in duplicate. MU Miller units.

R.I. (Repression Index) indicates the regulatory activity, calculated as the ratio between values obtained in the presence and in the absence of inducer.

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E30, placed within or adjacent to the DNA binding domain can therefore indirectly affect the affinity of the protein to DNA by properly modulating the protein conformation, allowing a correct alignment between the functional groups of the protein and the DNA.

Although there is no 'recognition code' between amino acids and nucleotides, they possess some preferential interactions, for instance arginines are known to interact favourably with guanines [4,5,6]. Thus, we propose that the effect observed in vivo of the recovery of regulation in the double mutant E30A OR_A1 T6R is due to the loss of interaction between E30, and R41 or R45, which results in a conformational change that allows a proper arrangement between the functional groups of the protein and the new operator DNA composition. R41 and R45 became free to establish new interactions with the nucleotides, not only the G at position 5, but also with the new G at position 6 (Figure 5D). Thus, the E30A mutation results in a better contact of the latter residues (R41 or R45) with G5 and the mutated G6 adjusting to the new DNA sequence, as observed by the increased regulatory activity of the mutant protein in the presence of the mutated operators (OR_A1 and OR_X2) when compared to the native protein (Table 4 and Table 5).

Our results provide information beyond the pairwise analysis, the data highlight and demonstrate that residues that are not involved in specific interactions with nucleotides, but act as linker residues by positioning other amino acids in the correct 3D context of a nucleoprotein complex, can be as important for the protein-DNA interaction as residues making direct contact with DNA bases, and have a crucial role in the modulation of DNA recognition. Furthermore, we show that by manipulating these residues it is possible to redesign the specificity of protein–DNA interactions.

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Author Contributions

Conceived and designed the experiments: ILC ISF IS-N. Performed the experiments: ILC ISF. Analyzed the data: ILC IS-N. Wrote the paper: ILC ISF IS-N.
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