Fine-Tuning Modulation of Oxidation-Mediated Posttranslational Control of *Bradyrhizobium diazoefficiens* FixK2 Transcription Factor

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Abstract: FixK2 is a CRP/FNR-type transcription factor that plays a central role in a sophisticated regulatory network for the anoxic, microoxic and symbiotic lifestyles of the soybean endosymbiont *Bradyrhizobium diazoefficiens*. Aside from the balanced expression of the fixK2 gene under microoxic conditions (induced by the two-component regulatory system FixLJ and negatively auto-repressed), FixK2 activity is posttranslationally controlled by proteolysis, and by the oxidation of a singular cysteine residue (C183) near its DNA-binding domain. To simulate the permanent oxidation of FixK2, we replaced C183 for aspartic acid. Purified C183D FixK2 protein showed both low DNA binding and in vitro transcriptional activation from the promoter of the fixNOQP operon, required for respiration under symbiosis. However, in a *B. diazoefficiens* strain coding for C183D FixK2, expression of a fixNOQP′-lacZ fusion was similar to that in the wild type, when both strains were grown microoxically. The C183D FixK2 encoding strain also showed a wild-type phenotype in symbiosis with soybeans, and increased fixK2 gene expression levels and FixK2 protein abundance in cells. These two latter observations, together with the global transcriptional profile of the microoxically cultured C183D FixK2 encoding strain, suggest the existence of a finely tuned regulatory strategy to counterbalance the oxidation-mediated inactivation of FixK2 in vivo.

Keywords: CRP/FNR proteins; in vitro transcription; microarrays; microoxia; protein–DNA interaction; rhizobia; symbiosis

1. Introduction

Nitrogen (N) is an essential nutrient for all living organisms on Earth. Biological nitrogen fixation (BNF) and denitrification represent two crucial pathways in the biogeochemical N cycle, maintaining the global balance of combined N (reviewed in [1–3]). Rhizobia are important contributors to BNF, a process that is highly relevant for both agronomy and the environment, since it reduces the need for chemical fertilizers in agriculture. They consist of a large group of α- and β-proteobacteria that can establish symbiotic associations with leguminous plants (reviewed in [4]). Importantly, they express the nitrogenase enzyme, which catalyzes the reduction of N2 to ammonium inside nodules located at the roots and occasionally on the stems of the plant partner (reviewed in [5,6]). During the symbiotic interaction, rhizobia are challenged to respond and adapt their physiology to a battery of signals. These include oxidative stress generated by the plants or the low partial pressure of free oxygen within the nodules (microoxia) (reviewed in [6–9]). Microoxia is needed for
the expression and functionality of nitrogenase and also the cbb3-type high-affinity terminal oxidase essential for bacterial respiration within the nodules (reviewed in [5,6,9–11]).

*Bradyrhizobium* species are the most widely employed diazotrophs as inoculants for soybean crops in agriculture [12]. In addition to being an efficient nitrogen fixer, *B. diazoefficiens* [13] is the only rhizobial species known for its ability to carry out complete denitrification, both in free-living and in symbiotic conditions ([14]; reviewed in [15,16]). In this bacterium, a complex regulatory network formed by two interconnected cascades (FixLJ-FixK2-NnrR and RegSR-NifA) controls the expression of genes required for microoxic, denitrifying and symbiotic modes of life ([17]; reviewed in [18]). The FixLJ-FixK2-NnrR cascade is oxygen-sensitive and activation by the two-component system FixLJ occurs at a concentration ≤ 5% O2, where the phosphorylated FixJ response regulator induces the expression of several genes, including *fixK2* (reviewed in [18]). The FixK2 protein plays a crucial role in this regulatory network, since it provides the link with the RegSR-NifA cascade and is also involved in the activation of hundreds of genes [19]. Among them, the *fixNOQP* operon encoding the high-affinity terminal oxidase *cbb3* genes involved in structural and accessory components of denitrification or regulatory genes (e.g., *rpoN1, fixK1* and *nnrR*) are included.

FixK2 is a member of the cyclic AMP receptor protein (CRP)/fumarate-nitrate reductase regulator (FNR) superfamily of bacterial transcription factors, which includes proteins that respond unevenly to a wide spectrum of environmental and intracellular cues (reviewed in [20–22]). This class of proteins has been described to control functions such as photosynthesis, virulence, carbon source utilization, nitrogen fixation and various modes of respiratory electron transport. CRP/FNR-type regulators have a fairly low similarity, but retain a well-conserved domain structure. This common protein architecture comprises an amino-terminal sensing domain linked via a long α-helical region (required for dimerization of the active homodimer) to a helix-turn-helix (HTH)-type DNA-binding domain at the carboxy-terminus (reviewed in [21]). This HTH motif recognizes and interacts with a palindromic DNA sequence located at distinct coordinates within the promoter region of regulated target genes (reviewed in [20,21]). In the case of FixK2, the consensus DNA recognition sequence is an imperfect 14-base-pair palindrome (TTGA/C-N6-T/GCAA, FixK2 box) [23,24].

Within CRP/FNR-type proteins, the transcriptional output to environmental and intracellular stimuli results from the interaction between a signaling molecule and the sensing domain. This induces a conformational change required for productive binding of the active dimer to the recognition sequence located at regulated gene promoters (reviewed in [25]). Signal perception can be through a direct response via a chemical modification of the protein or by binding to a specific prosthetic group or an effector molecule (reviewed in [21]).

Unlike most of the CRP/FNR superfamily members, the existence of a cofactor in modulating FixK2 transcription activation is unknown. Instead, *fixK2/FixK2* expression and FixK2 activity are subjected to complex transcriptional and posttranscriptional regulation (reviewed in [18]). Further to induction by the FixLJ system in response to microoxia, expression of the *fixK2* gene is auto-repressed by its own product by an as yet unidentified mechanism [26,27]. FixK2 is also controlled at a posttranslational level by oxidation [28] and by proteolysis, by both specific cleavage and also general degradation mediated by the ClpAP1 chaperone-protease system [29]. In addition, we recently observed that *fixK2* is among ~90 genes regulated at a posttranscriptional level in response to microoxia [30].

Oxidation-mediated posttranslational regulation of FixK2 occurs at the level of its single cysteine residue (C183), which resides in proximity to the DNA-binding domain [23,28]. The oxidation of this cysteine triggers protein inactivation either through the formation of dimers via an intermolecular disulfide bridge, or through the modification of cysteine to sulfenic, sulfenic or sulfonic acid variants, which inactivates the protein due to a steric hindrance effect and also to electrostatic repulsion with target promoters [23,28]. In vivo, FixK2 posttranslational oxidation might be relevant for the rapid cessation of transcrip-
tional activity in response to reactive oxygen species (ROS) produced at several stages of
the symbiotic interaction with soybeans (at the early stage of root hair infection, during
endosymbiotic respiration and at late nodule senescence) (reviewed in [7,8,31]).

The aim of this work was to advance our understanding of the mechanism under-
pinning the oxidation-mediated posttranslational control of the FixK$_2$ regulatory protein,
both in vitro and in vivo. Our hypothesis was that if C183 in FixK$_2$ was exchanged to
aspartic acid, this semi-conservative replacement (due to both its size and charge) would
permanently mimic FixK$_2$ overoxidation (e.g., sulfenic/sulfinic acid cysteine derivatives).
This stable modification might help to better unravel the consequences of FixK$_2$ oxidation
in vivo, especially regulation associated with transient bursts of ROS during symbiosis. We
characterized the DNA-binding properties, in vitro transcription (IVT) activation activity
and oligomeric state of recombinant C183D FixK$_2$. The effect of C183D FixK$_2$ was also ana-
lyzed in strains cultivated under free-living, microoxic conditions as well as in symbiosis
with soybean plants. Together, our results reveal a fine-tuning mechanism in B. diazoefficiens
to compensate for FixK$_2$ inactivation in response to cellular oxidizing conditions.

2. Results

2.1. Assessing the Impact of C183D Exchange in FixK$_2$ on In Vitro Transcription Activation
Activity and Protein–DNA Interaction Ability

Transcriptional regulation mediated by the FixK$_2$ protein is affected, among other
factors, through an oxidation-mediated posttranslational control (reviewed in [18]). The
C183 residue in FixK$_2$ plays a central regulatory role because it is sensitive to ROS, giving
rise to overoxidized species of the protein, i.e., sulfenic, sulfinic and sulfonic acid derivatives.
In order to mimic FixK$_2$ overoxidation, we performed a cysteine to aspartic acid replacement
and subsequent functional analyses of the C183D FixK$_2$ protein variant. In this context, its
performance was compared with that of the genuine FixK$_2$ protein [32], and with that of a
C183S FixK$_2$ derivative, which is oxidation-resistant [24]. All these proteins were previously
purified as untagged variants using the intein-mediated purification with an affinity chitin-
binding tag (IMPACT) methodology (New England Biolabs (NEB), Hitchin, UK).

The ability of the C183D FixK$_2$ protein to activate transcription in vitro in collaboration
with B. diazoefficiens RNA polymerase (RNAP) was monitored in a multiple-round IVT acti-
vation assay using the template plasmid pRJ8816, which harbors the fixNOQP operon pro-
moter cloned upstream of the B. diazoefficiens rrn transcriptional terminator (Figure 1) [33].
Importantly, this plasmid allows simultaneous analysis of both FixK$_2$-dependent (fixNOQP
transcript, 243 nucleotides [nt]) and FixK$_2$-independent (control transcript, 107 nt) transcrip-
tional responses elicited by B. diazoefficiens RNAP (Figure 1). The FixK$_2$ protein efficiently
activated transcription at 0.5 µM (Figure 1, lane 2), which increased at higher concentrations
(Figure 1, lanes 3 and 4). In contrast, the C183D FixK$_2$ derivative triggered low levels of
transcription from the fixNOQP promoter even when 2.5 µM of the protein was present
in the reaction (Figure 1, lane 7). However, the C183S FixK$_2$ variant showed higher levels
of transcription activation activity than the FixK$_2$ protein, reaching saturation at 0.5 µM
(Figure 1, lane 8).
Figure 1. IVT activation from the fixNOQP promoter mediated by different FixK2 protein derivatives. Plasmid pRJ8816 harboring the fixNOQP promoter cloned upstream of the B. diazoefficiens rrr transcriptional terminator was employed as template for multiple-round IVT activation assays with B. diazoefficiens RNAP holoenzyme. A series of concentrations of FixK2 protein variants were added to the reactions: lane 1, no protein (-); lanes 2, 5 and 8, 0.5 µM; lanes 3, 6 and 9, 1.25 µM; lanes 4, 7 and 10, 2.5 µM. The positions of the fixNOQP transcript and the FixK2-independent transcript (used as control for the experiments) are depicted on the right. Each panel refers to different sections of the same gel. Shown are the results of a typical experiment that was performed at least twice.

nt, nucleotides.

Since FixK2 belongs to the CRP/FNR-type transcription factor family, which act as functional dimers, the solution oligomeric state of C183D FixK2 was analyzed by size-exclusion chromatography (SEC) and compared to those of native FixK2 and the C183S FixK2 derivative (Figure 2). These experiments were performed to determine whether the diminished transcription efficiency of the C183D FixK2 protein variant could be attributed to an altered oligomeric state. Importantly, prior to SEC, each protein derivative preparation was analyzed by denaturing sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and showed a purity over ~95% for the band that corresponds to the predicted molecular mass of FixK2 (~25.6 kDa) (Figure S1). During non-denaturing individual SEC experiments for the three protein variants, chromatographic elution profiles showed a concentration-dependent behavior, with retention volumes ranging from the apparent molecular weight of the dimer (~52 kDa) to that of the monomer (~26 kDa) (Figure 2), as previously described for the N-terminally tagged wild-type protein [33]. The three proteins showed a monomer–dimer equilibrium; however, the proportion of the dimeric fraction with respect to the monomeric fraction was higher for the native FixK2 protein (Figure 2A) compared to C183S FixK2 and C183D FixK2 (Figure 2B and 2C, respectively) at similar concentrations. The reason for this difference might be related to the susceptibility of the wild-type derivative to the formation of disulfide bridges via C183. However, C183S FixK2 (Figure 2B) and C183D FixK2 (Figure 2C), which are devoid of cysteine residues, both showed similar monomer–dimer profiles despite their contrasting performance in transcriptional activation assays from the fixNOQP promoter. Therefore, the impaired IVT
activation activity observed for the C183D FixK\textsubscript{2} derivative is unlikely to be solely related to different oligomeric behavior.

**Figure 2.** Comparative SEC of native FixK\textsubscript{2} and C183S and C183D FixK\textsubscript{2} variants at different protein concentrations. Elution profiles were monitored at 280 nm following chromatography of FixK\textsubscript{2} loaded at 2.5, 5, 10, 20 and 30 \(\mu\)M (native, upper panel); C183S FixK\textsubscript{2} at 2.5, 5, 10, 20 and 40 \(\mu\)M (C183S, middle panel); and C183D FixK\textsubscript{2} at 2.5, 5, 10, 20 and 40 \(\mu\)M (C183D, bottom panel). The dashed lines show the calculated elution volume for the theoretical \(M_w\) of the monomeric (~26 kDa) and dimeric forms (~52 kDa).

To evaluate whether or not the C183D mutation in FixK\textsubscript{2} affects the DNA-binding capacity of the protein, electrophoretic mobility shift DNA assays (EMSA) were performed. Target DNA for these experiments was generated by PCR amplification of the promoter region of the \(\text{fixNOQP}\) operon. We found that a FixK\textsubscript{2}–DNA complex was readily detected when 0.25 \(\mu\)M of C183S FixK\textsubscript{2} protein was included in the reaction (Figure 3A, gel at the top). However, a concentration at least 16-fold higher (i.e., 4 \(\mu\)M) of the C183D FixK\textsubscript{2} protein was required to detect any interaction with DNA (Figure 3A, gel at the bottom), as determined by free-DNA disappearance, since the protein–DNA complexes apparently did not enter the gel at such protein concentration of this protein. Furthermore, a similar DNA mobility shift with each individual protein was only detected at a concentration approximately 32-fold higher of the C183D FixK\textsubscript{2} protein (8 \(\mu\)M) with respect to the C183S FixK\textsubscript{2} variant (0.25 \(\mu\)M) (Figure 3A), again determined by equivalent free DNA disappearance.
Figure 3. In vitro interaction of C183S and C183D FixK2 derivatives with the fixNOQP promoter tested by EMSA (A) and surface plasmon resonance (SPR) (B) approaches. (A) A 90-bp PCR fragment containing the FixK2 box at 20 nM was incubated with increasing concentrations (0 to 12 µM) of FixK2 protein variants, indicated at the top of each gel. Lower bands show free DNA, while upper bands correspond to the protein–DNA complexes. The molecular marker GeneRuler™ 1 Kb Plus DNA Ladder (Thermo Fisher Scientific, Waltham, MA, USA) is shown on the first lane. (B) A biotinylated double-stranded oligonucleotide containing the FixK2 box from the fixNOQP promoter was immobilized on a streptavidin (SA) sensor chip by biotin–streptavidin binding. The sensorgrams with the relative resonance units (RU) of the interaction with DNA of C183S and C183D FixK2 protein variants at 250 nM are shown. Data of the C183D FixK2 protein did not allow us to calculate any kinetic/affinity parameters.

The DNA-binding properties of the C183D FixK2 variant were also determined by employing the surface plasmon resonance (SPR) methodology (Figure 3B). In these assays, the FixK2 box located within the fixNOQP promoter was immobilized on a streptavidin (SA) sensor chip and the binding kinetics and affinity were analyzed by monitoring the response in resonance units (RU) vs. time. In line with the EMSA results, purified C183D FixK2 interacted poorly with DNA (Figure 3B). Further, neither affinity nor kinetic parameters
could be calculated as they were beyond the Biacore range and non-specific interactions were detected at high protein concentrations. This was in contrast with the results of a previous study performed with the C183S FixK2 derivative, which showed that FixK2–DNA interaction takes place at the nanomolar range and fitted well to a kinetic model for the interaction of one protein dimer per DNA molecule [24].

2.2. In Vivo Effects of C183 to Aspartic Acid Replacement in FixK2

To determine the effect of substituting C183 with aspartic acid in FixK2 in a cellular context, we performed a series of in vivo experiments. Firstly, we measured the β-Galactosidase activity of a chromosomally integrated fixNOQP'− lacZ fusion in a B. diazoefficiens strain encoding C183D FixK2 (C183D-fixK2) compared to the wild-type and ΔfixK2 strains, both used as controls (Figure 4). All strains were cultured under microoxic conditions (0.5% O2) for 48 h. An induction of approximately 600 Miller units (MU) was observed in the wild type, while, as expected, only basal levels were detected in the ΔfixK2 strain (Figure 4). However, expression of the fixNOQP'− lacZ in the C183D-fixK2 strain was similar to that observed for wild-type cells, suggesting that, in vivo, other mechanisms counterbalance the impaired transcriptional output of the C183D FixK2 protein observed in vitro.

![Figure 4](image)

**Figure 4.** Expression data for a chromosomally integrated fixNOQP'− lacZ fusion in different B. diazoefficiens backgrounds. Wild-type, C183D-fixK2 and ΔfixK2 strains were cultivated for 48 h microoxically (0.5% O2). β-Galactosidase values are means ± standard errors of a representative experiment performed with two parallel cultures assayed in quadruples. The experiment was repeated at least twice. WT, wild type.

Since FixK2 also directly or indirectly regulates the expression of genes involved in the denitrification process in B. diazoefficiens [19,34,35], we investigated whether the C183D mutation in FixK2 affects denitrifying growth (anoxia with nitrate as terminal respiratory electron acceptor) (Figure 5). Again, the C183D-fixK2 strain showed growth profiles that were similar to the wild type rather than the ΔfixK2 strain, where denitrifying growth is abolished.
The fixNOQP operon, employed as an archetypical target to monitor FixK activity [33], encodes the cbb3 high-affinity terminal oxidase, required for bacterial respiration within root nodules. To investigate the ability of C183D FixK2 to support the plant-endo- symbiotic interaction, we performed plant infection tests with soybeans inoculated with the wild type and the C183D-fixK2 and ΔfixK2 strains at two time-points: at 25 days post-inoculation (dpi), when maximal nitrogen fixation activity has been observed, and at 32 dpi, which corresponds to a late bacteroidal development stage [36] (Table 1).

### Table 1. Symbiotic phenotype of different B. diazoefficiens strains on soybean plants. Shoot dry weight (SDW), nitrogen shoot content (N), nodule number per plant (NN), nodule dry weight per plant (NDW), dry weight per nodule (NDW/NN) and leghemoglobin content in nodules (Lb) were determined at 25 and at 32 days post-inoculation (dpi). WT, wild type.

| Parameters          | WT     | ΔfixK2 | C183D-fixK2 |
|---------------------|--------|--------|-------------|
| **25 dpi**          |        |        |             |
| SDW (g)             | (0.54 ± 0.13) | (0.59 ± 0.10) | (0.47 ± 0.11) |
| N (mg)              | (12.60 ± 4.0)  | (4.90 ± 1.2)  | (12.20 ± 3.90) |
| NN                  | (38.30 ± 4.5)  | (34.50 ± 3.5)  | (32.20 ± 6.70) |
| NDW (mg)            | (38.67 ± 7.58) | (16.83 ± 1.94) | (32.50 ± 6.63) |
| NDW/NN (mg)         | (1.03 ± 0.24)  | (0.49 ± 0.03)  | (1.03 ± 0.22) |
| Lb (mg Lb · g NFW⁻¹) | (11.83 ± 0.59) | (0.11 ± 0.02)  | (10.08 ± 0.35) |
| **32 dpi**          |        |        |             |
| SDW (g)             | (0.92 ± 0.14)  | (0.68 ± 0.13)  | (0.77 ± 0.01)  |
| N (mg)              | (18.60 ± 5.90) | (5.00 ± 1.30)  | (21.00 ± 5.80) |
| NN                  | (31.30 ± 13.60)| (50.70 ± 13.60)| (27.80 ± 4.00) |
| NDW (mg)            | (38.17 ± 5.04) | (25.33 ± 6.31) | (32.83 ± 4.96) |
| NFW/NN (mg)         | (1.44 ± 0.65)  | (0.50 ± 0.05)  | (1.21 ± 0.28)  |
| Lb (mg Lb · g NFW⁻¹) | (11.51 ± 0.24) | (0.15 ± 0.02)  | (11.23 ± 0.71) |

Shown are the average values ± standard deviation of one representative experiment out of at least three repetitions (n = 6 plants per strain at harvest point).

No significant phenotypic differences, neither at 25 nor at 32 dpi, were observed in the C183D-fixK2 strain compared to the wild type with regard to several parameters relevant for...
plant-endosymbiotic efficacy, such as shoot dry weight (SDW), nitrogen shoot content (N), nodule number per plant (NN), nodule dry weight per plant (NDW), dry weight per nodule (NDW/NN) and leghemoglobin content in nodules (Lb) (Table 1). This contrasted with the phenotype of the plants inoculated with the ΔfixK2 strain, in which N, NDW/NN and Lb values were severely diminished (Table 1), which is in line with previous studies [24,26].

2.3. Appraisal of the Impact of the C183D Mutation on a Wider FixK2-Mediated Control Landscape

In order to reconcile and further understand the in vitro and in vivo results obtained with C183D FixK2, which suggested that, in cells, other mechanisms may compensate for the low DNA-binding capacity and IVT activation activity of the modified protein, a series of additional assays were performed. Firstly, we analyzed the abundance of FixK2 by Western blot of crude extracts from cells grown under microoxic free-living conditions and from soybean bacteroids (Figure 6A,B, respectively). Steady-state levels of FixK2 were approximately 2–3-fold higher in the C183D-fixK2 strain than in the wild type (Figure 6A, lane 2 vs. lane 1). A similar profile was also observed in soybean bacteroids extracted from nodules at 25 and at 32 dpi (Figure 6B, lanes 2 and 4 vs. lanes 1 and 3, respectively).

**Figure 6.** Expression of fixK2 at protein (A,B) and transcriptional (C) levels. Steady-state levels of FixK2 protein in cells cultivated under microoxic free-living conditions (A) and in soybean bacteroids collected at 25 and 32 dpi (B). Immunodetection was performed with a polyclonal FixK2 antibody [28]. (A) 60 µg of crude extract of wild-type (lane 1) and C183D-fixK2 strains (lane 2) both cultivated microoxically (0.5% O2). (B) 10 µL of soybean bacteroid crude extract of wild-type (lanes 1 and 3) and C183D-fixK2 strains (lanes 2 and 4). Apparent molecular mass of FixK2 is shown on the left. Representative results of at least three independent biological replicates are shown. (C) β-Galactosidase activity from a chromosomally integrated fixK2/-lacZ fusion in B. diazoefficiens wild-type, C183D-fixK2 and ΔfixK2 strains. Cells were cultivated for 48 h microoxically (0.5% O2). Values are the means ± standard errors of a typical experiment performed with two parallel cultures assayed in quadruples. The experiment was repeated at least twice. WT, wild type.

Based on these results, we also monitored whether the C183D FixK2 modification affected the expression of the fixK2 gene itself. Here, we measured β-Galactosidase activity...
from a fixK2- lacZ fusion integrated into the chromosome of the B. diazoefficiens C183D-fixK2 strain when cultivated microoxically (Figure 6C). In line with the increased levels of FixK2 protein observed in the immunodetection experiments, expression of fixK2 was around three-fold higher in the C183D-fixK2 strain compared to those values observed in wild-type cells. This induction profile for the fixK2- lacZ fusion was similar to that observed in the ΔfixK2 strain (Figure 6C; [18,26,27]) and therefore indicated that de-repression of fixK2 auto-regulation also occurred in the C183D-fixK2 strain.

In order to examine whether other, more global mechanisms could be involved in the C183D FixK2 phenotype in vivo, a global transcriptional analysis of the B. diazoefficiens C183D-fixK2 strain was performed and compared with that of the wild type, both grown under microoxic conditions. For this purpose, we employed the well-validated B. diazoefficiens custom-made GeneChip [37]. This comparative transcriptomic profile showed that 104 genes showed differential expression in the C183D-fixK2 strain, with 26 genes being upregulated and 78 genes downregulated (Table S1, Datasheet A; Figure 7). As expected, we found the fixK2 gene within the group of upregulated genes, and a relative change of fivefold was observed. However, among the downregulated genes in the C183D-fixK2 strain background, a series of legitimate FixK2-activated targets such as fixNOQP, fixGHIS and napEDABCD were not present. Similarly, the expression of genes encoding other CRP/FNR-type transcription factors under positive control of FixK2 (i.e., nnrR, fixK1; bll2109, bll3466) did not change.

**Figure 7.** Workflow of microarray data analyses of the C183D-fixK2 strain. Labels of the comparisons between specific transcription profiles are depicted alongside the circles. The total number of differentially expressed genes is indicated in parentheses. Up/down arrows refer to increased and decreased gene expression. The group of genes with differential expression in the C183D-fixK2 strain (dark grey circle, left) showed an overlap of 50 genes (light grey circle, middle) with those in ΔfixK2 strain (white circle, right; [19]), both grown microoxically (0.5% O₂) and compared with the wild type grown in the same conditions. Within the overlap, 47 genes showed downregulated expression in both the C183D-fixK2 and ΔfixK2 strains, which includes 37 genes organized in mono- or polycistronic transcriptional units that harbor a putative FixK2 box within the promoter region (26 putative transcriptional units, see Table 2).
Table 2. List of the 37 genes belonging to 26 putative FixK2 box-associated transcription units whose expression is downregulated in both the C183D-fixK2 and ∆fixK2 strains in comparison to the wild type (WT), both cultured microoxically (0.5% O2).

| Query     | FC (C183D-fixK2 vs. WT) | FC (∆fixK2 vs. WT) | Locus_Tag | Gene Name | Product                     | Position | Motif                     | Predicted Operon Structure |
|-----------|--------------------------|---------------------|-----------|-----------|----------------------------|----------|---------------------------|---------------------------|
| bll0330   | −2.4                     | −11.0               | Bdaspc4_01315 | -         | DNA-binding response       | −106     | TTGACCTGGATCAA             | -                         |
| bll0818   | −2.1                     | −9.3                | Bdaspc4_03880 | -         | hypothetical protein       | −66      | TTGATCCCCGGTCAA            | -                         |
|blr1289    | −3.2                     | −23.1               | Bdaspc4_06390 | -         | oleate hydratase           | −37      | TTGATCCAGCGCAA             | -                         |
| bll2517   | −3.2                     | −10.2               | Bdaspc4_12930 | -         | acetate/propionate family  | −89      | TTGACCTCACGCAA             | bll2518-bll2517           |
| blr3115   | −9.6                     | −30.6               | Bdaspc4_16100 | -         | MBL fold metallo-hydrolase | −74      | ATGATCTGGGTCAA             | blr3115                   |
| bll3117   | −2.4                     | −6.6                | Bdaspc4_16110 | -         | thymidine phosphorylase    | −287     | TTGACGTATCGCAA             | -                         |
| blr3815   | −2.2                     | −7.6                | Bdaspc4_19720 | -         | HAD family hydrolase       | −69      | TTGAGGTGCATCAA             | bll4240-bll4241           |
| blr4240   | −3.1                     | −25.1               | Bdaspc4_22005 | -         | pyridoxamine 5’-phosphate  | −75      | TTGACCTAGCGCAA             | -                         |
| blr4241   | −2.9                     | −83.3               | Bdaspc4_22010 | -         | cytochrome c               | −38      | TTGACCTCCGTCAA             | -                         |
| bll4412   | −3.2                     | −20.7               | Bdaspc4_22980 | -         | efflux RND transporter     | −72      | TTGACGTGGTCGCA             | -                         |
| blr4634   | −2.8                     | −20.2               | Bdaspc4_24260 | -         | periplasmic adaptor subunit| −48      | TTGACCTGCATCAA             | blr4652-blr4653-blr4654   |
| bll4635   | −2.5                     | −29.4               | Bdaspc4_24265 | groL5, groEL5 | translational machinery    | −150     | TTGCCGCTAGGTCAA            | -                         |
| blr4637   | −2.6                     | −111.5              | Bdaspc4_24275 | hspC2     | Hsp20/alpha crystallin     | −86      | TTGAGCAAAATCAA             | -                         |
| blr4644   | −3.2                     | −20.9               | Bdaspc4_24320 | -         | universal stress protein   | −72      | TTGATTTGCGTCGCA            | -                         |
| blr4645   | −2.8                     | −10.6               | Bdaspc4_24325 | -         | host attachment protein    | −69      | TTGATCGGGATCA              | -                         |
| bll4652   | −3.1                     | −95.2               | Bdaspc4_24370 | -         | nitroreductase             | −48      | TTGACCTGACATCAA            | blr4652-blr4653-blr4654   |
| bll4653   | −2.8                     | −16.8               | Bdaspc4_24375 | dnaJ      | J domain-containing protein| -        | -                          | -                         |
| blr4654   | −2.8                     | −30.0               | Bdaspc4_24380 | -         | hypothetical protein        | -        | -                         | -                         |
| bll4655   | −2.5                     | −14.2               | Bdaspc4_24385 | ppsA      | phosphoenolpyruvate        | −47      | TTGACCTGCCTCA              | -                         |
| bsr6066   | −4.0                     | −92.6               | Bdaspc4_31980 | -         | hypothetical protein        | −105     | TTGACCTGTCTCAA             | bsr6066-blr6067           |
| blr6067   | −2.7                     | −20.9               | Bdaspc4_31985 | -         | phage holin family protein | -        | -                         | -                         |
Table 2. Cont.

| Query a | FC (C183D-fixK₂ vs. WT) b | FC (ΔfixK₂ vs. WT) c | Locus_Tag d | Gene Name e | Product f | Position g | Motif h | Predicted Operon Structure i |
|---------|-------------------|-------------------|-------------|-------------|-----------|-----------|--------|-----------------------------|
| bll6073 | -3.5              | -27.9             | Bdiaspc4_32015 | *phaC2*     | probable poly-beta-hydroxybutyrate polymerase | -81 | TTGATGCAGCTCAA | - |
| bll6074 | -2.7              | -90.9             | Bdiaspc4_32020 | -           | CBS domain-containing protein | -143 | TTGAGCTGCATCAA | - |
| bll6525 | -2.1              | -7.7              | Bdiaspc4_34395 | -           | hypothetical protein | -22 | TTGATCTGCATCAA | - |
| bll7086 | -2.3              | -97.1             | Bdiaspc4_37390 | *hemN₂*     | oxygen-independent coproporphyrinogen III oxidase | -140 | TTGCAGGACCGCAA | - |
| bsr7087 | -3.2              | -53.8             | Bdiaspc4_37395 | -           | hypothetical protein | -115 | TTGCAGCAGCGCAA | bsr7087-bll7088 |
| bll7345 | -2.9              | -16.8             | Bdiaspc4_38745 | -           | - | -76 | TTGATCCGATCAA | - |
| bll7986 | -2.1              | -5.6              | Bdiaspc4_42230 | -           | HlyD family efflux transporter periplasmic adaptor subunit | -66 | CTGATCTAAATCAA | bll7988-bll7987-bll7986 |
| bll7987 | -2.5              | -17.4             | Bdiaspc4_42235 | -           | ABC transporter permease | - | CTGATCTAAATCAA | bll7988-bll7987-bll7986 |
| bll7988 | -3.3              | -33.1             | Bdiaspc4_42240 | -           | ABC transporter ATP-binding protein | - | CTGATCTAAATCAA | bll7988-bll7987-bll7986 |
| bll7989 | -2.6              | -5.3              | Bdiaspc4_42245 | *mat*       | methionine adenosyltransferase | -203 | TTGAGCCATGCAG | - |
| bll7990 | -3.2              | -19.7             | Bdiaspc4_42250 | -           | hypothetical protein isoprenylcysteine carboxyethyltransferase family protein | - | TTGATCTCGGCAA | bll7992-bll7991-bll7990 |
| bll7991 | -2.8              | -22.8             | Bdiaspc4_42255 | -           | - | - | TTGAGGGATGCAA | - |
| bll7992 | -2.7              | -23.0             | Bdiaspc4_42260 | DUF2933 domain-containing protein | -59 | TTGATCTCGGCAA | bll7992-bll7991-bll7990 |
| bll7993 | -2.8              | -8.5              | Bdiaspc4_42265 | -           | hypothetical protein | -60 | TTGAGGGATGCAA | - |

a Best blast hit in the *B. diazoefficiens* USDA 110 genome ([38]; GenBank acc. # NC_004463.1; RefSeq annotation as from January 2016). Direct FixK₂ targets as defined in [19] or validated by IVT are shaded in grey.
b Fold change (FC) values of gene expression in the C183D-fixK₂ strain in comparison to the WT, both grown under microoxic conditions.
c FC values of gene expression in cells of ∆fixK₂ in comparison to wild-type cells, both grown under microoxic conditions; [19].
d Nomenclature of *B. diazoefficiens* 110 spc₄ genes according to the NCBI annotation (GenBank acc. # CP032617); [30].
e Gene name according to the NCBI annotation with modifications (boldfaced) (GenBank acc. # CP032617); [30].
f Protein/gene product according to the NCBI annotation with modifications (boldfaced) (GenBank acc. # CP032617); [30].
g Position of the first nucleotide of the motif relative to the annotated translational start site of the associated gene.
h Predicted putative FixK₂ binding site.
i Operon structure prediction as previously described; Ref. [19].
The comparison of the C183D-fixK2 strain profile with the previously published transcriptional data of the ΔfixK2 strain under microoxic conditions [19] revealed a partial overlap between both groups of genes (Figure 7). In particular, while 54 genes, mainly represented by hypothetical and unknown proteins, were specific for the C183D-fixK2 strain profile (Table S1, Datasheet B), a further group of 50 genes were present in both profiles (Figure 7; Table S1, Datasheet C). Of this subset, 47 genes were downregulated (i.e., activated by FixK2), and specifically, 37 of them were organized into 26 transcriptional units, with each harboring a putative FixK2 binding site (Figure 7; Table 2). Furthermore, this includes 10 genes belonging to the set defined as putative direct FixK2 targets [19], and, in particular, the hspC2, ppsA, phaC2, hemN2 and bsr7087 genes, which were previously validated by IVT activation assays (compiled by Cabrera and coworkers [24]; Table 2). These observations demonstrate that the expression of certain FixK2-dependent targets is not counterbalanced in the C183D-fixK2 strain background.

3. Discussion

FixK2 is one of 16 CRP/FNR-type proteins present in the genome of B. diazoefficiens [39] but is distinguished among this family of regulators since it is capable of activating the transcription of the genes it regulates in collaboration with the RNAP of B. diazoefficiens in vitro, without any identifiable effector molecule [33]. Alternatively, different levels of regulation have been described for FixK2: (i) it is integrated into a complex regulatory network that responds to low oxygen formed by two interlinked cascades (FixLJ-FixK2 and RegSR-NifA), where fixK2 expression is balanced through FixLJ-mediated activation and FixK2-triggered auto-repression (direct or indirect by an unknown mechanism) ([17,19,26,27]; reviewed in [18]); (ii) the activity of FixK2 is modulated at posttranscriptional [30] and posttranslational levels (reviewed in [18]). This latter mode of regulation involves proteolysis by specific cleavage, and by general degradation mediated by the ClpAP1 chaperone-protease system [29] and oxidation at the level of residue C183 in response to oxidizing agents ([28]; reviewed in [18]).

Computational analyses of bacterial CRP/FNR family members performed by Matsui and coworkers [22] proposed that these proteins evolved from an ancestral FNR protein involved in nitrogen fixation. Although FixK-type proteins are part of the FNR group, they lack the [Fe-S] ligand-binding motif characteristic for FNR-type proteins. As with FixK2, other examples within the CRP/FNR protein family of regulatory proteins capable of activating gene transcription without the need of a cofactor are known and include (i) SdrP of Thermus thermophilus HB8, which is involved in the supply of nutrients and energy, redox control and the polyadenylation of mRNA. This protein not only is active in vitro without any cofactor but also lacks a putative binding pocket for a cofactor in its crystal structure [40]. (ii) PrfA of the human pathogen Listeria monocytogenes is capable of binding to its target DNA with low affinity without a cofactor [41] but its activity is modulated by carbon source availability in L. monocytogenes cells [42]. Recently, it was confirmed that reduced glutathione is the ligand for PrfA, both in vivo and in vitro [43,44]. (iii) FNR of Acidithiobacillus ferrooxidans ATCC23270 has low affinity for its [Fe-S] cofactor to allow a better transition between both aerobic and anaerobic environments [45]. (iv) Vfr of Pseudomonas aeruginosa can activate the transcription of some of its target genes in the absence of a cofactor (reviewed in [46]). In addition to cofactor-mediated modulation, regulation of targets by L. monocytogenes PrfA depends on the steady-state levels of this transcription factor in cells, which is subject to transcriptional, translational and posttranslational control [47]. Similarly, the activity of Escherichia coli FNR has also been shown to be modulated by protein levels through the degradation of the monomeric apo-protein by the ClpXP proteolytic system under oxic conditions [48,49]. All these antecedents, together with the key role of FixK2 in the microoxic metabolism of B. diazoefficiens, both in free-living conditions and in symbiosis, as well as in denitrification [19,26,34], support the possibility of the existence of alternative mechanisms for this protein to respond to intracellular and environmental stimuli.
The crystal structure of C183S FixK2 in complex with its genuine DNA-binding site (FixK2 box) present at the promoter of the fixNOQP operon [23] revealed why the C183 residue of FixK2 plays such a key role in its posttranslational control by oxidation. This is due to its proximity to the DNA-binding domain and its susceptibility not only to the formation of disulfide bridges but also to the generation of overoxidized sulfenic, sulfinic and sulfonic acid species, which result in electrostatic repulsion and steric hindrance [28]. Specifically, C183 interacts directly with the adenine located in position 7 of strand W [23], which is located immediately before thymine in position 8, which establishes hydrophobic interactions with the L195 residue of the HTH DNA-binding motif of FixK2.

In our work, we have analyzed whether the exchange of C183 for an aspartic acid residue can simulate the permanent oxidation of FixK2. To explore how the C183D mutation may affect FixK2–DNA interaction in silico, we modeled a battery of protein derivatives (i.e., FixK2, C183S FixK2, C183D FixK2 and the sulfenic, sulfinic and sulfonic FixK2 variants) with the double-stranded FixK2 box DNA sequence from the fixNOQP promoter (Figure 8). According to these predictions, the replacement of C183 by aspartic acid causes the acquisition of a free negative charge and consequently an electrostatic repulsion with the phosphate groups of both the adenine 6 and adenine 7 bases of strand W of the target DNA [23]. Furthermore, the presence of the oxygen atom from the aspartate branched side chain also gives rise to steric hindrance due to the proximity of this atom to the bases described above, reducing the intermolecular distances of 7.1 and 4.9 angstroms (Å), to 4.5 and 3.5 Å, respectively (Figure 8A and Figure 8C). Thus, the C183D FixK2 derivative–DNA interaction likely mimics that of the sulfenic-derived cysteine (Figure 8D) and the sulfinic-derived cysteine (Figure 8E) due to the size and charge of each radical, respectively, rather than the most oxidized sulfonic-derived cysteine (Figure 8F).

Figure 8. Modeling of different FixK2 protein variants with the double-stranded DNA containing the FixK2 box present at the fixNOQP promoter. Shown are the protein–DNA distances between the negatively charged oxygens of the phosphate group of the nitrogenous bases adenine 6 and adenine 7 of the strand W of DNA [23], and cysteine (A), serine (B), aspartic acid (C), cysteine–sulfenic acid (D), cysteine–sulfinic acid (E) and cysteine–sulfonic acid (F) residues of FixK2. The predictions of the 3D models of FixK2 and C183D FixK2 were obtained with the Pymol 2.2.3 program (https://pymol.org/2/; accessed on 13 October 2021), using the C183S FixK2–DNA structure as a template (http://wwpdb.org/; code 4I2O; accessed on 7 June 2021). Visualization of molecular structures and interactions was performed using the Discovery Studio Visualizer program, version V20.1.0.19295 (BIOVIA, Waltham, MA, USA), which also allowed the modeling of sulfenic, sulfinic and sulfonic acid derivatives of FixK2. Distances in angstroms (Å) are represented by dashed lines; adenine 6 on the left; adenine 7 on the right.
The oxidation-mediated FixK$_2$ inactivation similarity of the C183D FixK$_2$ derivative was first analyzed in vitro. As expected, purified C183D FixK$_2$ showed a low DNA-binding ability, determined by both SPR and EMSA approaches (Figure 3). This may also have affected the interaction with the RNAP polymerase and holocomplex conformation required for transcriptional output, as an impaired IVT activation capacity (a reduction of approximately 75%) for the C183D FixK$_2$ protein derivate was observed in comparison to the FixK$_2$ and C183S FixK$_2$ variants (Figure 1). Furthermore, the monomer–dimer equilibrium of the oligomeric state of the C183D FixK$_2$ protein variant appeared to be shifted more to the monomeric form in comparison to that of the native FixK$_2$ protein (Figure 2). However, as this profile was fairly similar to that of the oxidation-insensitive C183S FixK$_2$ protein, which interacts effectively with DNA and is fully active (Figure 1), it cannot be taken as the main factor to explain its deficiency in both DNA-binding capacity and IVT activation activity.

Despite the results found in vitro, intriguingly, the *B. diazoefficiens* C183D-fixK$_2$ strain showed a wild-type phenotype with regard to the expression of a fixNOQP$^{-}$-lacZ fusion under microoxic conditions (Figure 4), its denitrifying growth behavior (Figure 5) and its symbiotic performance with soybeans (Table 1). This was in contrast with the phenotype of a ΔfixK$_2$ strain [24,26] and indicated the existence of alternative mechanisms in *B. diazoefficiens* cells, which compensates for the in vitro characteristics of the C183D FixK$_2$ protein variant. To test this hypothesis, we then monitored the steady-state levels of C183D FixK$_2$ protein in both *B. diazoefficiens* cells grown under free-living microoxic conditions and in soybean bacteroids isolated from nodules at 25 and 32 dpi. In all conditions tested, the abundance of the C183D FixK$_2$ protein was higher (approximately 2–3 fold) than the wild-type protein (Figure 6), which could explain the absence of a phenotype of the *B. diazoefficiens* C183D-fixK$_2$ strain in our in vivo assays.

In order to obtain a global overview of the effect of the C183D replacement in FixK$_2$, a transcriptomic profile of the *B. diazoefficiens* C183D-fixK$_2$ strain grown under microoxic conditions was next performed and compared to that of the wild type. Some remarks are here mentioned. A high proportion (920 out of 970) of the genes belonging to the FixK$_2$ regulon did not show differential expression in the C183D-fixK$_2$ strain (Figure 7). This group includes other genes encoding CRP/FNR-type regulators whose expression is activated by FixK$_2$, such as bll2109, bll3466, fixK$_1$ and mnrR [19]. This finding, together with the increased abundance of the C183D FixK$_2$ protein (Figure 6), might be the rationale for the compensated expression of genes belonging to the FixK$_2$ regulon in the C183D-fixK$_2$ strain. In this context, we should not overlook that the C183D FixK$_2$ variant did not seem to mimic the inactive, most oxidized, sulfonic acid derivative of the FixK$_2$ protein (Figure 8), which might contribute to the mild phenotype of the *B. diazoefficiens* C183D-fixK$_2$ strain.

Regardless of these arguments, 104 genes still showed differential expression in the C183D-fixK$_2$ strain in comparison with the wild type. Interestingly, 47 genes belonging to this group are under the positive control of FixK$_2$, and 37 of them are organized in 26 transcriptional units that contain a FixK$_2$ binding site within their promoter region (Figure 7; Table 2). This set includes direct targets compiled in [24], such as *hemN$_2$*, *phbC$_2$*, *ppsA*, bll4637 or bsr7087, but neither the fixNOQP operon encoding the *cbb*$_3$ high-affinity terminal oxidase nor the *napEDABC* genes encoding the periplasmic nitrate reductase involved in denitrification were present. These observations indicate that the overexpression of C183D FixK$_2$ is not sufficient to compensate for the FixK$_2$-mediated activation of transcription for all its targets. However, the inspection of the FixK$_2$ boxes associated with the 26 transcription units, as well as the neighbor nucleotides (positions 6 and 7 of strand W of the fixNOQP promoter DNA; [23]), did not reveal a conserved pattern that could offer a plausible reason for this differential behavior of the C183D FixK$_2$ protein with respect to the activation of the expression of direct targets.

Within the group of genes differentially expressed in the C183D-fixK$_2$ strain, around half (54 out of 104; Table S1) were not part of the FixK$_2$ regulon. Among them, we did not find induction of those encoding other CRP/FNR-like proteins that could also counter-
balance the constrained behavior of the C183D FixK2 variant. Instead, we encountered a large proportion of genes that code for hypothetical or unknown proteins, which makes it difficult to conduct a more comprehensive study.

Importantly, in accordance with the fixK2'-lacZ fusion data determined under microoxic conditions (Figure 6), we found increased expression of the fixK2 gene in the C183D-fixK2 strain in comparison with the wild type. This enhanced expression was also previously found in the ΔfixK2 strain [18,26,27], which is an indication that FixK2 negatively regulates its own expression (directly or indirectly) by an unknown mechanism. Reutimann and coworkers [27] proposed that this control is likely indirect, where the FixK2 protein may be involved in the activation of its own repressor or an activator of the fixK2 repressor gene.

As de-repression of the fixK2 gene still occurred in the C183D-fixK2 strain, we surveyed the list of genes that appeared to be downregulated in both C183D-fixK2 and ΔfixK2 regulons to identify possible candidates. None of the remaining regulatory genes previously proposed (i.e., blr1216, bsr4636, blr7666) ([27]; reviewed in [18]) appeared in such groups of genes (Table S1). Nevertheless, we found a predicted response regulator gene, bll0330, which harbors a putative FixK2 binding site within its promoter region (Table 2). Although its expression is also under the positive control of the response regulator FixJ, it was previously overlooked as it is not induced under microoxic conditions [19]. The functional analysis of this gene in the context of fixK2 negative auto-regulation would be interesting to pursue; however, we believe that this is beyond the scope of this paper.

4. Materials and Methods
4.1. Strains, Plasmids, and Primers

A detailed description of the plasmids and bacterial strains used in this work is compiled in Table 3. Table S2 describes primer names and sequences employed in this study.

| Strain or Plasmid | Description | Resistance | Source or Reference |
|-------------------|-------------|------------|---------------------|
| **Strains**       |             |            |                     |
| E. coli           |             |            |                     |
| DH5α              | supE44 ΔlacU169 (qpx80 lacZ ΔM15)                  |            | Bethesda Research   |
|                   | hisD17 recA1 endA1 gyrA96 thi-1 relA1             |            | Laboratories Inc.,  |
|                   |                                                        |            | Gaithersburg, MD, USA |
| S17-1             | thi pro recA hisD17 RP4Tc::Mu Km::Tn7             | Tp' Sm' Spc' | [50]               |
|                   | fluA2 lacZ::T7 gene1 [lon] ompT sulA11             |            |                     |
|                   | R(mcr-73::miniTn10-TetO2)2 [dcm]                  |            |                     |
|                   |                                                        |            |                     |
| ER2566            |                                                        |            | NEB, USA            |
|                   | R(zgb-210::Tn10-TetO2) endA1                      |            |                     |
|                   | Δ(mcrC-mrr)114::T110                               |            |                     |
| **B. diazoefficiens** |             |            |                     |
| 110spc4           | Wild type (WT)                                    | Cm' Spc'  | [51]               |
| 9043              | ΔfixK2                                             | Cm' Spc' Sm' | [26]               |
| 1255              | C183D-fixK2                                        | Cm' Spc'  | This work          |
| 3604              | WT::fixNOQP-lacZ                                    | Cm' Spc' Te' | [52]               |
| 9043-3603         | ΔfixK2::fixNOQP-lacZ                                 | Cm' Spc' Te' | This work          |
| 1255-3603         | C183D-fixK2::fixNOQP-lacZ                           | Cm' Spc' Te' | This work          |
| 1109              | WT::fixK2'::lacZ                                    | Cm' Spc' Te' | This work          |
| 9043-1109         | ΔfixK2'::fixK2'::lacZ                                | Cm' Spc' Te' | [18]               |
| 1255-1109         | C183D-fixK2'::fixK2'::lacZ                          | Cm' Spc' Te' | This work          |
### Table 3. Cont.

| Strain or Plasmid | Description | Resistance | Source or Reference |
|-------------------|-------------|------------|---------------------|
| **Plasmids**      |             |            |                     |
| pTXB1             | Expression vector for the IMPACT protein purification system. It codes for a C-terminal thiol-cleavable Mxe GyrA-intein–chitin-binding domain (CBD) under T7 promoter control | Amp<sup>r</sup> | NEB, USA |
| pBBR1MCS-2        | lacPOZ mobRP4, low-copy-number cloning vector | Km<sup>r</sup> | [53] |
| pK18mobsacB       | Mobilizable pUC18 derivative, mob, sacB | Km<sup>r</sup> | [54] |
| pRJ0051           | [pTXB1] with a 715-bp NdeI/SpeI fragment encoding C183S FixK<sub>2</sub>-intein fused in frame with the CBD of the vector | Amp<sup>r</sup> | [32] |
| pRJ0053           | [pTXB1] with a 715-bp NdeI/SpeI fragment encoding FixK<sub>2</sub>-Intein fused in frame with the CBD of the vector | Amp<sup>r</sup> | [32] |
| pRJ8848           | [pUC19] with a 2.288-kb SalI fragment encoding C183S FixK<sub>2</sub> | Amp<sup>r</sup> | [23] |
| pMB1250           | with a 2.288-kb SalI fragment encoding C183D FixK<sub>2</sub> | Amp<sup>r</sup> | This work |
| pMB1251           | [pBBR1MCS-2] with a 1.843-kb BamHI-Xbal fragment from pMB1250 | Km<sup>r</sup> | This work |
| pMB1253           | [pTXB1] with a 715-bp NdeI/SpeI fragment from pMB1251 encoding C183D FixK<sub>2</sub> | Amp<sup>r</sup> | This work |
| pRJ9041           | [pUC19] with a 2.288-kb SalI fragment encoding FixK<sub>2</sub> | Amp<sup>r</sup> | [33] |
| pMB1256           | [pRJ9041] with a 3.965-kb NotI fragment from pMB1251 | Amp<sup>r</sup> Km<sup>r</sup> | This work |
| pMB1254           | [pMB1256] Religation of a 4.974-kb BglII fragment | Amp<sup>r</sup> | This work |
| pMB1255           | [pK18mobsacB] with a 1.849-kb BamHI fragment from pMB1254 | Km<sup>r</sup> | This work |
| pRJ3603           | [pSUP202pol2] ‘blr2761, blr2762 and fixNOQP’-‘lacZ on a 8.261-kb XhoI fragment | Tc<sup>r</sup> | [52] |
| pRJ9054           | [pSUP202] fixJ, blr2758 and fixK<sub>2</sub>-‘lacZ on a 4.434-kb NsiI/DraI fragment | Tc<sup>r</sup> | [26] |
| pMB1109           | [pRJ9054] fixK<sub>2</sub>-‘lacZ with a 136-bp SmaI fragment deletion within the blr2758 coding region | Tc<sup>r</sup> | [18] |

### 4.2. Media and Growth Conditions

*E. coli* cells were typically grown in Luria–Bertani (LB) medium [55] at 37 °C overnight. When needed, antibiotics were added at the following concentrations (in µg·mL<sup>−1</sup>): ampicillin, 200; kanamycin, 30; spectinomycin, 25; streptomycin, 25; tetracycline, 10.

*B. diazoefficiens* strains were routinely cultured oxically at 30 °C under rigorous shaking (170 rpm) in a peptone–salt–yeast extract (PSY) medium [19,51]. Microoxic cultures (0.5% O<sub>2</sub> in PSY medium), and under denitrifying conditions (anoxia in yeast extract–mannitol [YEM] medium supplemented with 10 mM KNO<sub>3</sub>; [56]) were essentially prepared as described previously [24]. The initial optical density (OD) at 600 nm of the cultures was 0.02, except for those employed in β-Galactosidase assays, which was 0.2, since not all the strains showed the same growth behavior. In the microoxic cultures, the gas phase was exchanged in cycles of 8/16 h. Antibiotic concentrations in *B. diazoefficiens* cultures were as follows (in µg·mL<sup>−1</sup>): chloramphenicol, 15 (solid medium); kanamycin, 200 (solid medium), 100 (liquid medium); spectinomycin, 200 (solid medium), 100 (liquid medium); tetracycline 100 (solid medium), 50 (liquid medium).

### 4.3. Strain and Plasmid Construction

A *B. diazoefficiens* strain that encodes a C183D FixK<sub>2</sub> protein variant was constructed using a markerless mutagenesis approach based on the sacB-based methodology [54,57]. Firstly, C183 in FixK<sub>2</sub> was exchanged by aspartic acid using site-directed mutagenesis and...
plasmid pRJ8848 as a template, and oligonucleotides fixK_2_mut59 and fixK_2_mut60, yielding plasmid pMB1250. A 1.843-kb BamHI/XbaI fragment from pMB1250 was then cloned into the corresponding sites of the pBBR1MCS-2 vector, thus resulting in plasmid pMB1251. Next, a 3.965-kb NotI fragment from pMB1251 was inserted into the linearized NotI pRJ9041 plasmid, to give rise to plasmid pMB1256. This plasmid was subsequently cut with BglII, and recirculation of a 4.974-kb fragment yielded plasmid pMB1254. Finally, to construct plasmid pMB1255, a 1.849-kb BamHI fragment derived from plasmid pMB1254 was cloned into the suicide vector pK18mobsacB. Plasmid pMB1255 was then transferred to E. coli S17.1 cells, which were employed in biparental conjugation with B. diazoefficiens wild type. Single recombination transconjugants were selected by kanamycin resistance, followed by double recombination selection by sucrose resistance, as described elsewhere [57]. The genomic organization of the resulting markerless strain encoding a C183D FixK_2 derivative (strain 1255) was verified by PCR and sequencing using specific primers (Table S2).

In order to construct a plasmid that expresses a C183D FixK_2 derivative fused at its C-terminal region with the Mxe GyrA-intein–chitin-binding domain (CBD) expressed under the control of the T7 promoter, a 727-bp PCR-amplified fragment from pMB1251 with the oligonucleotides fixK_2_mut19 and fixK_2_mut58 was restricted with NdeI and SpeI and subsequently cloned in frame into the pTXB1 vector (NEB, Hitchin, UK), thus resulting in plasmid pMB1253. The correctness of the plasmid pMB1253 sequence was verified by sequencing with suitable primers (Table S2).

To construct B. diazoefficiens C183D FixK_2 encoding strains harboring either a fixNOQP’- ‘lacZ or a fixK_2’- ‘lacZ translational fusion, plasmids pRJ3603 and pMB1109 were transferred from E. coli S17.1 cells via biparental conjugation into the chromosome of the 1255 strain. Transconjugants were selected by tetracycline resistance and further verified by PCR and sequencing, yielding strains 1255-3603 and 1255-1109, expressing fixNOQP’- ‘lacZ and fixK_2’- ‘lacZ fusions, respectively.

Plasmid and genomic DNA isolation was performed using the Qiagen Plasmid Kit (Qiagen, Germantown, MD, USA) and REALPURE Genomic DNA (Durviz, Valencia, Spain), respectively.

4.4. β-Galactosidase Activity Assays

Expression of fixNOQP’- ‘lacZ and fixK_2’- ‘lacZ fusions in B. diazoefficiens cells grown under microoxic conditions was analyzed by measuring β-Galactosidase activity. Cells cultivated for 48 h were first permeabilized and subsequently used for the assays, as previously described [55,57]. The absorbance at 420 nm of the enzymatic reactions and at 600 nm of the cultures was recorded in a plate reader (SUNRISE Absorbance Reader; TECAN, Männedorf, Switzerland) using the Xfluor4 software (TECAN, Männedorf, Switzerland). These data were used to calculate the specific activity of β-Galactosidase in Miller units (MU).

4.5. Plant Infection Test and Physiological Analyses

Plant inoculation and growth experiments were performed essentially as described previously [58]. Soybean seeds (Glycine max L. Merr., cv. Williams 82, harvest at October 2011) were firstly surface-sterilized and germinated at 30 °C for 48 h in darkness. After germination, seeds were sown in 0.25 L pots containing sterile vermiculite and 50 mL of modified Jensen N-free solution, as indicated earlier [58]. The seedlings were then inoculated independently with cell suspensions of each strain in sterile saline solution (0.9% w/v NaCl) at an OD_600 of 0.5 (~10^5 cells mL^-1), prepared from oxically grown cultures collected at stationary phase (OD_600 ~1). Plants were then cultivated under controlled conditions with an initial irrigation with modified Jensen medium followed by sterile deionized water until harvest at 25 and 32 dpi.

The plant physiology parameters nodule number per plant (NN), nodule dry weight (NDW) per plant, dry weight per nodule (NDW/NN) and shoot dry weight (SDW) were measured after harvesting as described by Tortosa and coworkers [58]. For bacteroid isolation and additional analyses, a minimum of 1 g of fresh nodules randomly collected
from at least 3 plants were stored at −80 °C after quick freezing in liquid nitrogen. SDW was recorded after 3 days at 70 °C; samples were ground to less than 0.5 mm for nitrogen (N) determination. N content in SDW was measured by the Dumas method using the LECO TruSpec CN Elemental Analyzer [59].

For leghemoglobin (Lb) determination in the nodular fraction, 0.5 to 1.0 g nodules were manually homogenized by using a cooled porcelain pestle and mortar with 6 mL of buffer solution (50 mM Na₂HPO₄ · 2H₂O/NaH₂PO₄ · 2H₂O, pH 7.4, 0.02% w/v K₃Fe(CN)₆, and 0.1% w/v NaHCO₃) and 0.1 g of polyvinyl poly(vinylpolypyrrolidone) (PVPP) according to the methodology described in previous studies [58]. Then, the extract was centrifuged at 12,000 × g at 4 °C for 20 min. Lb content was fluorometrically determined after an acidic reaction at 120 °C during 30 min, according to LaRue and Child [60]. After cooling the samples, the fluorescence in each tube was measured with a spectrophotofluorometer (Shimadzu Scientific Instruments, Kyoto, Japan) (λexcitation = 405 and λabsorption = 600 nm). Non-autoclaved tubes containing acidic nodular fraction were used as a control.

4.6. Overexpression and Purification of Non-Tagged FixK₂ Protein Variants

Non-tagged FixK₂ protein derivatives were purified with the IMPACT system (NEB, Hitchin, UK) according to the protocol detailed in [24]. In brief, E. coli ER2566 cells individually transformed with plasmids pRJ0051, pRJ0053 and pMB1253 were grown in 500 mL of LB medium at 37 °C until an OD₆₀₀ of 0.3. The cultures were then incubated for 1 h at 30 °C up to an OD₆₀₀ of 0.8, before addition of 0.1 mM IPTG for the induction of overexpression of the individual recombinant proteins. After incubation for 16 h at 16 °C, cells were collected and employed for protein purification [24]. Fractions of the different purification steps were collected and analyzed by Blue Coomassie-stained 14% SDS-PAGE, as described by Laemmli [61]. Cell pellets were resuspended in loading dye (62.5 mM Tris HCl pH 6.8, 2% SDS, 10% glycerol, 50 mM dithiothreitol [DTT], 0.01% bromophenol blue) in a proportion of 100 µL per mL of OD₆₀₀ = 1 and subsequently boiled at 95 °C for 10 min and centrifuged at 12,000 × g for 5 min before loading. For desalting, protein fractions from the affinity chromatography were pooled and buffer-exchanged by passing them through a prepacked Sephadex G-25M column (PD-10; Cytiva Europe GmbH, Cornellá de Llobregat, Spain) equilibrated with the suitable buffer for each further assay (IVT activation activity, SEC, EMSA, SPR).

4.7. In Vitro Transcription Activation Assay

IVT activation experiments were basically performed as described in previous studies [24,33,62]. Essentially, 20 µL reactions containing the basic transcription components, 1 µg of B. diazoefficiens RNAP, 750 ng of plasmid pRJ8816 that harbors the promoter of the fixNOQP operon [33] and different concentrations (0, 0.5, 1.25 and 2.5 µM) of individual protein derivatives (i.e., FixK₂, C183S FixK₂ and C183D FixK₂) were incubated at 37 °C for 30 min. Transcription products were monitored with a PhosphorImager (Molecular Dynamics, Massachusetts, MA, USA) and signal intensities were evaluated with the Image Lab™ software (Bio-Rad, California, CA, USA).

4.8. Size-Exclusion Chromatography Experiments

Analytical SEC experiments of the FixK₂ protein derivatives were performed at room temperature on a Superdex 200 10/300 GL column (Cytiva, Little Chalfont, UK) using an ÅKTA PURE protein purification system (Cytiva, Little Chalfont, UK). After equilibrating the column with elution buffer (40 mM Tris-HCl, pH 7.0, 150 mM KCl 0.1 mM EDTA), 100 µL protein samples were injected and separated at a flow rate of 0.75 mL.min⁻¹. Absorbance was recorded at 280 nm. The following proteins were used as standards for calibration (Figure S2): conalbumin (75 kDa), ovalbumin (43 kDa), carbonic anhydrase (29 kDa), ribonuclease A (13.7 kDa) and aprotinin (6.5 kDa) (Cytiva, Little Chalfont, UK). Gel filtration experiments were repeated at least three times with independent preparations of each protein at a range of at least five concentrations. The UNICORN™ system control
software (Cytiva, Little Chalfont, UK) was employed to program the chromatography runs and for preliminary analyses of the data by adjusting for injection times.

4.9. Electrophoretic Mobility Shift DNA Assays

Stable FixK$_2$–DNA interaction was tested electrophoretically. First, 15 $\mu$L reactions containing 10 ng of purified 90-bp PCR fragment spanning the promoter region of the fixNOQP operon (Table S2) and different protein concentrations, from 0 to 12 $\mu$M, in modified IVT buffer (40 mM Tris-HCl pH 8, 10 mM MgCl$_2$, 0.1 mM EDTA, 0.1 mM DTT, 150 mM KCl, 0.4 mM K$_3$PO$_4$), were incubated for 30 min at room temperature. Reactions were mixed with one sixth volume of loading dye (30% glycerol in modified IVT buffer supplemented with bromophenol blue) and loaded onto a 6% non-denaturing polyacrylamide–0.5X Tris-Borate EDTA (TBE) gel. After running the electrophoresis for 40 min at 180 V, gels were incubated in a 1X SYBR-Gold (Invitrogen, Waltham, MA, USA) solution in 0.5X TBE for 30 min. Finally, UV-induced signals were detected by a Gel Doc XR+ System (Bio-Rad, California, CA, USA) and quantified with the Quantity One and Image Lab software (Bio-Rad, California, CA, USA).

4.10. Surface Plasmon Resonance Analyses

FixK$_2$–DNA interaction ability was analyzed by SPR using a Biacore X100 Biosensor (Cytiva Europe GmbH, Cornellá de Llobregat, Spain) with SA sensor chips according to the methodology described by Cabrera and coworkers [24]. All buffers were previously filtered and degassed. The biotinylated double-stranded fixNOQP promoter region was synthesized by annealing complementary primers (Table S2), leaving the biotinylated primer at 10 $\mu$M. Then, the double-stranded oligonucleotide was diluted at 5 nM in immobilization buffer (Tris-HCl 10 mM pH 7.5, 50 mM NaCl, 1 mM EDTA) and captured at 100 RU in a sensor chip. Protein–DNA interaction assays were carried out in running buffer (40 mM Tris-HCl pH 7.0, 150 mM KCl, 0.1 mM EDTA) supplemented with 0.005% Tween 20 at 25$^\circ$C. The analyte was injected in both flow cells at 40 $\mu$L/min during 120 s of contact time, followed by 120 s of dissociation. In a first round, the analyte was diluted in running buffer from 0 to 250 nM in a random order, with at least one duplicate of a low concentration analyte after a higher concentration. Range of protein concentration was extended up to 3 $\mu$M in further experiments. The sensor surface was regenerated with injections of 0.2% SDS at 30 $\mu$L/min during 60 s. The number of trials, computer support and data analysis and quantification were as described earlier [24].

4.11. Immunoblot Detection of FixK$_2$

Steady-state levels of FixK$_2$ protein were monitored in B. diazoefficiens cells grown under microoxic conditions and in soybean bacteroids by immunoblotting using a polyclonal antibody against FixK$_2$ [28]. At least three biological replicates of 300 mL of microaerobically grown cultures (0.5% O$_2$) at mid-exponential phase (OD$_{600}$ of 0.45–0.58) were collected (5000× g, 7 min, 4 $^\circ$C), washed with fractionation buffer (40 mM Tris-HCl pH 7.0, 150 mM KCl) and resuspended in 1.5 mL of the same buffer containing 0.2 mM 4-[2-Aminoethyl]benzenesulfonyl fluoride hydrochloride (AEBSF). Cell suspensions were disrupted by three passes through a cold French pressure cell (SLM Aminco, Jessup, MD, USA) at approximately 120 MPa, and subsequently centrifuged (27,000× g, 30 min, 4$^\circ$C) to obtain total cell-free extracts.

Isolation of bacteroids from soybean nodules inoculated with the different strains was performed as described elsewhere [14,58]. In short, 0.8 to 1 g of nodules per strain and condition were employed. After extraction, bacteroids were resuspended in 2 mL of 50 mM Tris-HCl pH 7.4. Cell density of bacteroid suspensions was determined and adjusted to an equal OD$_{600}$ with the same buffer. Then, aliquots were taken, centrifuged at 12,000× g for 5 min and resuspended in six-fold-diluted SDS loading dye (350 mM Tris-HCl pH 6.8, 10% SDS, 30% glycerol, 620 mM DTT, 0.01% bromophenol blue) in a proportion of 20 $\mu$L per mL.
of OD$_{600} = 1$. Finally, they were boiled at 95 °C for 10 min and centrifuged at 12,000× g for 5 min before loading.

Conditions of SDS-PAGE and Western blotting were similar to those described in previous studies [30,62]. Samples were resolved in 14% SDS-PAGE, and subsequently transferred to nitrocellulose membranes using a Trans-Blot Turbo System (Bio-Rad, California, CA, USA). A rabbit-derived polyclonal antibody against FixK$_2$ [28] at a 1:1000 dilution was used as primary antibody, while a horseradish peroxidase (HRP)-conjugated goat anti-rabbit IgG (Bio-Rad, California, CA, USA) at a 1:3500 dilution was employed as secondary antibody. Visualization of the signals was performed with a ChemiDoc XRS instrument (Universal Hood II, Bio-Rad California, CA, USA). The Quantity One and Image Lab software programs (Bio-Rad, California, CA, USA) were employed for image analyses.

4.12. Determination of Protein Concentration

Protein concentrations of samples employed in Western blot assays, as well as of purified recombinant proteins, were determined using the Bio-Rad reagent (Bio-Rad, California, CA, USA) and bovine serum albumin (BSA) as the standard protein for the calibration curve. The concentration of purified proteins used in this study is referred to in the dimeric form.

4.13. Microarray Sample Preparation and Data Analyses

For microarray experiments, B. diazoefficiens cultures were grown to mid-exponential phase (OD$_{600}$ of 0.45 to 0.58). Cell harvest, isolation of total RNA, cDNA synthesis, fragmentation, labeling and conditions for hybridization with a custom-designed B. diazoefficiens Gene Chip BJAPETHa520090 (Affymetrix, Santa Clara, CA, USA) were as described in previous studies [19,36,37].

For these experiments, 1.8 µg of labeled fragmented cDNA was hybridized to the arrays. A minimum of three independent biological samples were analyzed. Signal intensities detection, normalization and analyses were done with Affymetrix Expression Console software version 1.4.1 (Affymetrix, Santa Clara, CA, USA). Transcriptome analysis Console 3.1 software (Affymetrix, Santa Clara, CA, USA) was used for comparative analyses. Normalized intensities (MAS 5.0 algorithm) were compared between conditions using one-way between-subject ANOVA (ANOVA p-value < 0.05). Only genes that passed the statistical tests and where the change in expression (measured as n-fold change [FC]) was $\geq 2$ or $\leq -2$ in comparisons between two strains were considered as differentially expressed.

4.14. Biocomputing Analyses

In silico analyses of the interaction of the battery of FixK$_2$ protein derivatives with DNA were performed based on the structure of the FixK$_2$–DNA complex ([23]; http://wwpdb.org/, entry PDB 4I2O; accessed on 7 June 2021). The prediction of the 3D models of FixK$_2$ and C183D-FixK$_2$ was obtained with the Pymol 2.2.3 program (https://pymol.org/2/; accessed on 13 October 2021), using the C183S FixK$_2$–DNA structure as a template. The visualization of molecular structures and interactions was performed using the Discovery Studio Visualizer program version V20.1.0.19295 (BIOVIA, Waltham, MA, USA), which also allowed modeling and predictions with derivatives of FixK$_2$ proteins that harbor specific mutations or alterations in the oxidation state.

5. Conclusions

The main goal of this work was to better understand FixK$_2$-dependent regulation essential for low-oxygen metabolism (microoxia) of the model denitrifying plant-endosymbiotic bacterium B. diazoefficiens. Microoxia has been recognized as an essential signal for both nitrogen fixation and denitrification.

Our intention was to explore whether cells could be pre-primed for ROS defense through modification of the single redox active cysteine (C183) in the FixK$_2$ transcription factor. Our functional study of a C183D FixK$_2$ variant, simulating permanent overoxidation of the protein, reveals the existence of a cellular mechanism to counteract inactivation that
boosts FixK2 levels through transcriptional and posttranscriptional means, giving rise to wild-type phenotypes in both free-living cells and soybean bacteroids.

We believe that our research provides a platform to undertake further synthetic biology approaches to modify rhizobial FixK-type proteins and improve the durability of the symbiotic interaction and fitness in response to oxygen. This could be applied to enhance the productivity and sustainability of soybean crops, which will contribute to global food security, human health and the environment.

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