A novel gene from the acidophilic bacterium *Leptospirillum* sp. CF-1 and its role in oxidative stress and chromate tolerance

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**Abstract**

**Background:** Acidophilic microorganisms like *Leptospirillum* sp. CF-1 thrive in environments with extremely low pH and high concentrations of dissolved heavy metals that can induce the generation of reactive oxygen species (ROS). Several hypothetical genes and proteins from *Leptospirillum* sp. CF-1 are known to be up-regulated under oxidative stress conditions.

**Results:** In the present work, the function of hypothetical gene ABH19_09590 from *Leptospirillum* sp. CF-1 was studied. Heterologous expression of this gene in *Escherichia coli* led to an increase in the ability to grow under oxidant conditions with 5 mM K₂CrO₄ or 5 mM H₂O₂. Similarly, a significant reduction in ROS production in *E. coli* transformed with a plasmid carrying ABH19_09590 was observed after exposure to these oxidative stress elicitors for 30 min, compared to a strain complemented with the empty vector. A co-transcriptional study using RT-PCR showed that ABH19_09590 is contained in an operon, here named the "och" operon, that also contains ABH19_09585, ABH19_09595 and ABH19_09600 genes. The expression of the och operon was significantly up-regulated in *Leptospirillum* sp. CF-1 exposed to 5 mM K₂CrO₄ for 15 and 30 min. Genes of this operon potentially encode a NADH:ubiquinone oxidoreductase, a CXXC motif-containing protein likely involved in thiol/disulfide exchange, a hypothetical protein, and a di-hydroxy-acid dehydratase. A comparative genomic analysis revealed that the och operon is a characteristic genetic determinant of the *Leptospirillum* genus that is not present in other acidophiles.

**Conclusions:** Altogether, these results suggest that the och operon plays a protective role against chromate and hydrogen peroxide and is an important mechanism required to face polyextremophilic conditions in acid environments.

**Keywords:** Hypothetical proteins, Chromate, Hydrogen peroxide, Oxidative stress, Och operon, Leptospirillum

**Background**

*Leptospirillum* spp. and other acidophiles face environmental conditions that favor the generation of reactive oxygen species (ROS) [1–4]. Consequently, they may experience oxidative stress, a very deleterious condition where all major cellular components, such as DNA, RNA, proteins, lipids and cofactors can suffer oxidative damage with substantial negative impacts on cellular physiology and activity. *Leptospirillum* sp. CF-1 is an aerobic acidophilic iron-oxidizing bacterium with the capacity to survive in extremely acidic natural or man-made environments, and in high concentrations of dissolved metal(loids) such as Cu, Zn, Al, Cd, Cu, Ni, Cr and As [3, 5]. We have investigated the molecular mechanisms that mediate the antioxidant response in *Leptospirillum* spp., and have established that members of this genus have a highly abundant and efficient thiol/disulfide system [6].
Leptospirillum spp. also possesses antioxidant enzymes, like Dyp-type peroxidases (DyP) [7] and cytochrome c peroxidases (CcP) [8], which are responsible for eliminating the peroxides generated in the cytoplasm and periplasm, respectively. It has also been established that Leptospirillum sp. CF-1 uses a mechanism based on cobalamin (vitamin B_{12}) to protect against oxidative conditions and avoid the induction of oxidative stress [3].

The genome of Leptospirillum sp. CF-1 has been completely sequenced [9], facilitating genomic, transcriptomic and proteomic studies to provide a better understanding of the adaptive mechanisms to extreme environments. Since no genetic systems have been developed for this microorganism to date, omics approaches have become a very important tool in microbial physiology studies. However, genomic data derived from Leptospirillum sp. CF-1 has revealed the existence of a very high number of genes with unknown function (1519 out of 2736; 56%), most of which are present in genetic contexts that are also unknown in terms of function [9]. These hypothetical genes have the regulatory elements required to be transcribed and translated, but the predicted products have no known function or structure, making their biological role and contribution towards adaptation to extreme life impossible to deduce [10]. In the case of strain CF-1, and other strains of the genus, more than half of the annotated genes in its genome still need to be functionally characterized. This proportion is similar to less-characterized microorganisms or extremophiles (>50%) [11], and substantially higher than in model organisms (25–40%; [12]. For example, in cyanobacterial genomes, 30–60% of the putative proteins are encoded by hypothetical genes depending on the species [10]. A similar profile has been described for extremophilic microorganisms; for example, 36% are hypothetical proteins in Acidithiobacillus ferrooxidans and At. thiooxidans [13, 14], and 47% of protein-coding genes from Leptospirillum ferrariphilum are of unknown function [15], a figure that rises to 63% in the case of Ferrovum myxofaciens [16]. Although many hypothetical genes may correspond to non-functional entities called pseudogenes [17], it is also likely that many have functions not yet characterized. As such, they could possess novel biological roles that could allow us to gain a deeper and more comprehensive understanding of the mechanisms involved in adaptation to environmental challenges. Interestingly, transcriptomic and proteomic assays carried out in Leptospirillum sp. CF-1 have shown a large number of hypothetical genes or proteins that were down or up-regulated under oxidative stress conditions (unpublished observations), suggesting a role of these in the adaptation of Leptospirillum spp. to highly oxidative and metal loaded environments.

Bioinformatic studies to characterize hypothetical genes and proteins have traditionally included analyses that search for orthologous genes or sequence similarity to previously characterized proteins, the presence of conserved domains or motifs, the prediction of protein cellular location, and the identification of neighboring genes, among other predictions [18–20]. The utilization of in silico approaches to predict the function of hypothetical genes and proteins has been successfully used in several models of bacterial pathogens [21–23]. However, since most of these analyses are performed based on comparisons with previously-known protein sequences, structures or domains, it becomes evident that there is insufficient data to obtain novel information about the nature, origin and role of predicted hypothetical proteins in less-characterized or extremophilic microorganisms. Thus, in the present work, we have addressed the study of the hypothetical gene ABH19_09590. This gene was heterologously expressed in Escherichia coli to evaluate its ability to confer protection against oxidative stress. Furthermore, neighboring genes from Leptospirillum sp. CF-1 were studied to assess the genetic context of ABH19_09590. The expression profiles of neighboring genes found in this operon were determined to understand their potential role in chromate tolerance.

**Results**

**Selection of hypothetical genes that are upregulated under oxidative stress conditions**

A previous study carried out in Leptospirillum sp. CF-1 showed that oxidative stress induced with ferric ions (260 mM Fe_{2}SO_{4}, Fe^{3+}) for 1 h led to the up-regulation of genes and proteins involved in the synthesis of iron-sulfur centers, protein folding, energy metabolism, synthesis of exopolysaccharides, and metabolism of methionine (data not shown). Interestingly, these experiments also showed changes at the expression level of a number of hypothetical genes and proteins. According to our predictions, 38% of the genes or proteins that were significantly down- or up-regulated corresponded to hypothetical proteins, suggesting that they could be involved in the oxidative stress response of this microorganism and indicating that they could play a pivotal role under environmental stress conditions.

Among all Leptospirillum sp. CF-1 genes that showed statistically reliable regulation of expression levels in transcriptomic assays under ferric ion stress, we chose those whose products were originally annotated as hypothetical proteins and were conserved among other members of the Leptospirillum genus. In this way, we selected four up-regulated genes: ABH19_09590 (4.7 fold-change), ABH19_06740 (3.6 fold-change), ABH19_07700 (2.8 fold-change), and ABH19_010550 (1.8 fold-change).
These genes were evaluated by means of heterologous complementation experiments (described below), to confer protection to *E. coli* against potassium chromate (data not shown). In this initial screen, the ABH19_09590 gene showed a high protective effect, and was thus selected for further analysis and evaluation.

**E. coli** transformed with ABH19_09590 showed tolerance to heavy metals and oxidative stress conditions

Since no molecular tools exist for *Leptospirillum* spp., in order to evaluate the functionality of ABH19_09590 gene from strain CF-1, we measured its ability to confer protection against oxidative stress conditions when heterologously-expressed in *E. coli*. The selected gene was cloned into the arabinose-inducible expression vector pBATopo TA, as described in Methods, and used to transform *E. coli* (Top 10 strain). The transformed cell culture was grown until OD\(_{600}\) 0.5 and expression of the cloned gene was induced with L-arabinose for 1.5 h. Subsequently, the cultures were exposed to oxidative stress culture conditions as described in Methods. The ferric iron initially used to induce oxidative stress in *Leptospirillum* spp. at acidic pH [3], has a high tendency to precipitate and form aggregates with organic components of the circumneutral medium for *E. coli* cultivation therefore potentially affecting the quantification of its biological effect in this bacterium. For this reason in this study we chose potassium chromate and hydrogen peroxide to induce oxidative stress. The effect of these as oxidative stress elicitors in *Leptospirillum* spp. has also been previously reported [3, 24]. In agreement with those previous studies, the exposure of wild type *E. coli* to 5 mM K\(_2\)CrO\(_4\) or 5 mM H\(_2\)O\(_2\) led to a significant decrease in cell growth (K\(_2\)CrO\(_4\); 50%, and H\(_2\)O\(_2\); 63%) at 12 h, as compared to cells that were not exposed to these oxidant agents (100%; Fig. 1). Interestingly, as shown in Fig. 1A and B, heterologous ABH19_09590 transformation exerted a significant positive effect in restoring cell growth in the presence of 5 mM K\(_2\)CrO\(_4\) (81% at 12 h) and 5 mM H\(_2\)O\(_2\) (87% at 12 h). In this way, these data suggest that ABH19_09590 could be exerting a protective role against the oxidative effect of chromate and ROS.

**ABH19_09590 attenuates ROS generation in E. coli**

To determine whether the ABH19_09590 gene product has a suppressive effect on ROS generation, the intracellular ROS levels in *E. coli* exposed to oxidant agents were measured. As shown in Fig. 2, empty vector cells exposed to 5 mM K\(_2\)CrO\(_4\) or 5 mM H\(_2\)O\(_2\) for 30 min showed significantly increased ROS contents (by 335% and 258%, respectively) as compared to control cells that were not exposed to stress (100%). Interestingly, and consistent

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**Fig. 1** Growth of *E. coli* transformed with ABH19_09590 from *Leptospirillum* sp. CF-1. Cells transformed with plasmid pBADTopo/ABH19_09590 were exposed to **A** 5 mM K\(_2\)CrO\(_4\) or **B** 5 mM H\(_2\)O\(_2\). Cell growth was measured each hour at 600 nm absorbance.
with results observed in cell growth, exposure of transformed and induced cells to 5 mM K2CrO4 or 5 mM H2O2 for 30 min, led to a much more minor increase in ROS content (by 120% and 108%, respectively) compared to control cells without exposure (100%). It should be noted that a decrease in intracellular ROS level was not observed in stressed cells harboring the empty pBAD vector. These results suggest that the protein encoded by ABH19_09590 alleviates the cytotoxic effect of oxidative stress elicitors by directly or indirectly scavenging ROS and restoring the redox balance of the cell.

Analysis of the genetic context of the ABH19_09590 gene

In a previous study, we generated a high-quality sequence of the whole genome of strain CF-1 [9]. For this reason, and in order to obtain additional information on the potential role of the ABH19_09590 gene and its encoded protein, we analyzed its genetic context. Immediately upstream from ABH19_09590, genes that encode for a hypothetical protein (ABH19_09575) and for putative NADH dehydrogenase subunit L (NuoL, ABH19_09585) are found, while downstream, genes encoding a hypothetical protein (ABH19_09595), a possible dihydroxyacid dehydratase (DHAD, ABH19_09600) and the translational factor Sua5 (ABH19_09605) were detected. The six candidate genes were analyzed to evaluate co-transcription using RT-PCR, in order to define whether ABH19_09590 is contained in an operon. RT-PCR experiments clearly show that ABH19_09590 is contained in an operon. RT-PCR experiments clearly show that ABH19_09590, ABH19_09595, ABH19_09595 and ABH19_09600 are co-transcribed (Fig. 3), demonstrating that they are expressed and form part of an operon that we have called the “och operon” (oxidative stress and chromate response). The physical genetic arrangement of the och operon, including the flanking non-co-transcribed genes (ABH19_09575 and ABH19_09605) is shown in Fig. 3. The co-transcription of these four genes suggests that they are functionally related and similarly activated in response to physiological signals. Therefore, these results strongly suggest the involvement of the four genes of the och operon in the oxidative stress response of Leptospirillum spp.

The och operon is up-regulated in Leptospirillum sp. CF-1 under oxidative stress

To evaluate the role of the och operon in oxidative defense, the mRNA levels of each of the four genes contained in this operon were evaluated using RT-qPCR assays when Leptospirillum sp. CF-1 was exposed to oxidative stress conditions. Unfortunately, due to the high reactivity between the ferrous ion and hydrogen peroxide, and the subsequent generation of radicals that promote oxidative damage to biomolecules, exposure of iron-oxidizing microorganisms to peroxide usually leads to difficulties in purification of RNA with a quality suitable for performing RT-qPCR experiments. For this reason, oxidative stress induction in Leptospirillum sp. CF-1 was assayed with 5 mM K2CrO4 for 15 and 30 min, without the addition of H2O2. As can be observed in Fig. 4, the transcript levels of the four genes of the och operon increased significantly by 2.8–12.5 fold after 15 min, and by 7.8–23.5 fold after 30 min in cells exposed to potassium chromate. These data strongly suggest a relevant role of the och operon in the physiological oxidative stress response of Leptospirillum sp. CF-1.

In-silico predictions of the proteins encoded by the och operon from Leptospirillum sp. CF-1

To characterize the gene products that make up the och operon, an in-silico analysis was performed: **ABH10_09585** (ochA). The protein encoded by the first gene of the operon from strain CF-1 is predicted to contain 553 amino acids with a theoretical molecular mass of 59.6 kDa and a pl of 9.44. The protein is similar to the membrane-bound subunits L (NuoL)
of NADH:ubiquinone oxidoreductase (54% similarity to NuoL subunit from *Shewanella* sp.). NuoL has a role in proton pumping in complex I of the respiratory chain. However, in many bacterial species this protein is a stand-alone Na⁺/H⁺ antiporter not coupled with oxidoreduction, but displaying a more general role in multiple resistance mechanisms and pH adaptation [25, 26].

**ABH10_09590** (*ochB*) The analysis of the deduced amino acid sequence of ABH10_09590 of *Leptospirillum* sp. CF-1 indicates that the predicted encoded protein possesses 823 amino acids with a theoretical molecular mass of 92 kDa. The prediction shows that this protein harbors five transmembrane domains and lacks domains of known function. However, a DUF2309 domain of unknown function is predicted. DUF2309 is a non-characterized, but widely distributed and conserved domain, spanning 748.4 amino acids on average [27]. Interestingly, this predicted DUF from the deduced ABH10_09590 protein possesses a CXXC active-site motif between amino acids 530 and 533. This motif is highly conserved in most DUF2309 domains and is similar to that reported in iron-sulfur proteins, thioredoxins and oxidoreductases [28]. We postulate that the presence of the CXXC motif may contribute to the formation, isomerization, or reduction of disulfide bonds, or determine other redox functions in the ABH10_09590 protein that could explain the in vivo effect on the growth and ROS concentration under oxidative conditions, when expressed in *E. coli* (Figs. 1 and 2).

**ABH10_09595** (*ochC*) The encoded hypothetical protein is predicted to have 823 amino acids, a mass of 92 kDa and to be located at the cytoplasmic membrane. It does not possess conserved domains or motifs that could give insights into its role in strain CF-1.

**ABH10_09600** (*ochD*) The protein encoded by the fourth gene of the operon corresponds to a putative dihydroxy-acid dehydratase (DHAD), with 558 amino acids and 59.2 kDa. This protein participates in the metabolism of amino acids, and plays a key role in the biosynthesis of valine, leucine and isoleucine from Archaea, Bacteria and Eukarya [29]. Also, it has been shown to be involved in the biosynthesis of D-pantothenate and coenzyme A [30]. More recently, it has been described that up-regulation of DHAD protein in *Staphylococcus* sp. #NIOSBK35 facilitates its survival and growth at high concentrations of Cr(VI), conferring the capability to deal with this strong oxidizing agent [31]. It has been proposed that the up-regulation of DHAD could be indicative of enhanced cellular activity as a strategy to cope with the stress induced by Cr(VI). According to our inspection, dihydroxy-acid dehydratase from *Leptospirillum* sp. CF-1 is a predicted cytoplasmic protein with high similarity to DHAD from *E. coli* (45% identity and 63% similarity).
Interestingly, genes that encode ABH19_09585, ABH19_09590, ABH19_09595, ABH19_09600 proteins are detected in the genomes of a number of other acidophilic microorganisms including *Leptospirillum ferriphilum*, *Acidiferrobacter thiooxydans*, *Acidibacillus sulfuroxidans*, *Acidithiobacillus ferrivorans*, *Acidithiobacillus ferrooxidans*, *Sulfobacillus acidiphilus*, *Ferrovum sp.*, *Ferrithrix thermotolerans*, *Sulfobacillus thermosulfidoxidans*, and *Acidihalobacter prosperus*, among others, with an identity and query cover of over 47% and 74%, respectively (Table 1). However, with the exception of the member from the *Leptospirillum* genus, in these microorganisms the genes are not organized into operons, and appear dispersed throughout the genome.

**Discussion**
Acidophilic bacteria must maintain the intracellular ROS level under control in order to survive the low pH and/or the toxic levels of heavy metals in their environment. This implies that these bacteria must have developed sophisticated mechanisms to reduce the levels of ROS and repair damaged biomolecules. Members of the genus *Leptospirillum* grow under these environmental conditions in the absence of canonical antioxidant systems, namely superoxide dismutase (SOD), catalase (CAT), and glutathione [1, 3]. However, alternative molecular mechanisms to maintain redox homeostasis have been described in *Leptospirillum* spp., including the thiol/disulfide system [6], Dyp-type peroxidases (DyP) [7], cytochrome *c* peroxidases (CcP) [8] and even the use of cobalamin to protect against oxidative conditions [3]. In the present study, we generated experimental evidence that directly or indirectly links the protein encoded by the hypothetic gene ABH19_09590 of *Leptospirillum* sp. CF-1 with protection against oxidative damage. The results showed that *E. coli* cells overexpressing the gene product of the ABH19_09590 gene were able to grow after exposure to the oxidant metal ion Cr(VI) and the oxidative specie *H*₂*O*₂, compared to negative controls. In addition, the measurement of the intracellular ROS levels...
generated by the pro-oxidant agents reinforced the fact that all *E. coli* cultures were exposed to oxidative stress conditions. Interestingly, the culture carrying the plasmid with ABH19_09590 gene had ROS levels similar to those found in control conditions, supporting its antioxidant function.

Additional hints regarding potential biological functions can arise from data concerning the genomic context of hypothetical genes [11]. Thus, we performed a co-transcription analysis to determine whether the ABH19_09590 was part of an operon. The results demonstrated that this gene is co-transcribed with genes encoding NuoL (ABH19_09585), a hypothetical protein (ABH19_09595) and a DHAD (ABH19_09600), altogether conforming the operon denominated “och”. Additionally, in line with our previous unpublished observations regarding the upregulation of ABH19_09595 under oxidative stress (with ferric iron), strain CF-1 exposed to chromate exhibited significantly enhanced mRNA levels of each gene from this cluster, suggesting the up-regulation of the *och* operon under the oxidant conditions encountered in these extreme acidic environments. The *in-silico* analysis of the amino acid sequence of the ABH19_09590 gene revealed the presence of a CXXC active site motif, as found in thioldisulfide oxidoreductases belonging to the thioredoxin superfamily. Their two cysteines usually exist in an oxidized (disulfide bonded) or reduced (unpaired) state, which allow disulfide bonds to generate changes in the redox state and/or conformation of their target proteins [28, 32], which could explain the protective effects observed. The neighboring gene encodes a putative membrane localized NuoL. As a potential proton pump, in the absence of other proteins belonging to complex I of the respiratory chain, NuoL could be involved in pH adaptation [25, 26]. On the other hand, the DHAD protein could confer the ability of the bacterium to deal with Cr(VI) by enhancing cellular activity [31]. Therefore, this *in silico* evidence strongly suggests that the *och* operon could be a response mechanism against oxidative stress, and more specifically to chromate in *Leptospirillum* sp. CF-1.

Comparative genomics has shown that a substantial fraction of the genes in sequenced genomes encode conserved hypothetical proteins that are found in organisms from several phylogenetic lineages and have no known function [11]. Based on a bioinformatic search within the genomes of different acidophilic iron-oxidizing microorganisms, the genes from the *och* operon are apparently conserved, which could suggest their importance for subsisting in extreme bioleaching environmental conditions. However, it is important to highlight that only bacteria from the *Leptospirillum* genus possess an operon-like organization. Therefore, whether the full operon confers better fitness under oxidative conditions to *E. coli*, compared to each gene alone, is a task that should be addressed.

In bacteria, Cr(VI) is imported via anion transporters, and sulfate importers especially seem to represent a major import pathway due to the structural similarity of these two oxyanions [33, 34]. Inside the cell, Cr(VI) generates active intermediates Cr(V) and/or Cr(IV), free radicals, and Cr(III) as the final product. Cr(III) affects DNA replication, causes mutagenesis, and alters the structure and activity of enzymes, reacting with their carboxyl and thiol groups [34–37]. Therefore, ROS are generated during the reduction of Cr(VI) to Cr(III). However, several bacterial mechanisms of Cr(VI) detoxification have been described in detail, such as the chromium resistance (*chr*) operon [33, 38], efflux pumps [38, 39],

| Microorganism                        | Identity (%) | ABH19_09585 | ABH19_09590 | ABH19_09595 | ABH19_09600 |
|--------------------------------------|--------------|-------------|-------------|-------------|-------------|
| *Leptospirillum ferriphilum*         | 100          | 99          | 80          | 99          |
| *Acidiferrobacter thiooxydans*       | 57           | 61          | 54          | –           |
| *Acidibacillus sulfuroxidans*        | 57           | 57          | –           | –           |
| *Acidiithiobacillus ferrireducens*   | 52           | 59          | 54          | 72          |
| *Acidiithiobacillus feroxoxidans*    | 52           | 59          | 54          | 72          |
| *Sulfobacillus acidophilus*          | 51           | 54          | –           | –           |
| *Ferrovum* sp.                       | 52           | 54          | 55          | 72          |
| *Ferrithrix thermotolerans*          | 48           | 56          | –           | –           |
| *Sulfobacillus thermosulfidooxidans* | 47           | 53          | –           | –           |
| *Acidihalobacter prosperus*          | 51           | 57          | 59          | –           |
| Query cover                          | 74–100%      | 98–100%     | 93–99%      | 99%         |
extracellular Cr(VI) reduction [40–43], enzymes involved in the detoxifying processes [44–46], and repair of DNA lesions [47–50]. In the case of acidophilic microorganisms, Cr(VI) precipitation has been identified in *At. ferroxidans*, which can tolerate Cr(III) up to 75 mM during growth on ferrous sulphate [51]. In addition, *Acidiphilum cryptum* JF-5 has the ability to reduce Cr(VI) under aerobic and anaerobic conditions via two mechanisms, a NADPH-dependent chromate reductase and a cytochrome (ApcA) [52]. In strain CF-1, no genes related to Cr(VI) resistance were found by bioinformatic searches. However according to the results generated in this work, we propose that the ABH19_09590 gene product could be part of a new chromate resistant mechanism achieved by direct Cr(VI) reduction, or reduction of the ROS generated.

**Conclusions**

The ABH19_09590 gene from *Leptospirillum* sp. CF-1 has a role increasing tolerance and attenuating ROS generation when transformed *E. coli* are exposed to chromate and hydrogen peroxide. The ABH19_09590 gene encodes a DUF2309 domain-containing protein likely involved in thiol/disulfide exchange reactions. This gene forms part of the och operon that also contains genes encoding a NuoL subunit of NADH:ubiquinone oxidoreductase (ABH19_09585), a hypothetical protein (ABH19_09595) and a dihydroxy-acid dehydratase (ABH19_09600). The genes from the och operon are up-regulated in strain CF-1 under chromate exposure. Finally, the och operon is a unique genetic determinant from members of the *Leptospirillum* genus and may represent an important mechanism needed to face polyextremophilic conditions of acid environments. Additional efforts are required to elucidate the role of hypothetical genes in extremophilic microorganisms.

**Methods**

**Bacterial strains and growth conditions**

*Leptospirillum* sp. CF-1 was grown as described previously by Ferrer et al. [3]. Briefly, it was grown in 9 K BR medium containing 18.4 g/L ferrous sulfate (FeSO₄·7H₂O) at 37 °C with constant stirring at 180 rpm. *E. coli* Top 10 was grown in Luria–Bertani (LB) under aerobic conditions at 37 °C and constant stirring at 180 rpm. The growth of all recombinant *E. coli* strains was estimated by measuring optical density (OD) at 600 nm.

**DNA isolation**

*Leptospirillum* sp. CF-1 was grown until the late exponential phase. Cells were harvested by centrifugation at 8000 ×g for 15 min and washed once with acid water (10 mM H₂SO₄) and twice with 10 mM sodium citrate pH 7.0. Genomic DNA 1 was isolated using the Wizard® Genomic DNA Purification Kit (Promega) according to the manufacturer’s instructions and stored at −20 °C until further use.

**Cloning of ABH19_09590 gene in an expression vector**

The ABH19_09590 gene was cloned in the pBADTopo TA expression vector (Thermo Fisher Scientific, Waltham, MA, USA) to generate pBADTopo/ABH19_09590. The cloning was performed using the PCR product obtained from ABH19_09590 gene with genomic DNA from *Leptospirillum* sp. CF-1 as a template and specific primers (Table 2). The ligation product was obtained according to the manufacturer’s instructions, and then introduced into *E. coli* Top 10 using the CaCl₂ transformation protocol [53]. To select transformants, the transformation mix was plated on LB-Agar media containing 100 µg/mL ampicillin.

**Heterologous expression assay**

Wild-type *E. coli* Top 10, or cells carrying pBadTopo/ABH19_09590 or pBadTopo alone (empty vector) were grown until OD₆₀₀ = 0.2–0.3. Gene expression was induced with 0.2% L-arabinose during 1.5 h. Afterwards, cultures were diluted (OD₆₀₀ = 0.2–0.3) with fresh medium containing 0.2% L-arabinose and treated with 5 mM K₂CrO₄ or 5 mM H₂O₂. Controls with non-exposed cells were included. Bacterial growth was monitored hourly by spectrophotometry at OD₆₀₀ for 12 h.

**Determination of ROS Levels**

The oxidant-sensitive probe H₂DCFDA (2′,7′-dichlorodihydrofluorescein diacetate) [54] was used to determine the intracellular level of total ROS according to [3]. For this purpose, 15 mL of wild-type *E. coli* Top 10, or cells harboring pBadTopo/ABH19_09590, or pBadTopo alone were cultured overnight in LB media containing 100 µg/mL ampicillin. Afterwards, cultures were centrifuged and resuspended in fresh medium with 100 µg/mL ampicillin until OD = 0.5. Subsequently, cell cultures were induced with 0.2% L-arabinose during 1.5 h, and further exposed to oxidative stress with 5 mM K₂CrO₄ or 5 mM H₂O₂ for 30 min. Controls with non-exposed cells were included. After stress induction, cells were collected by centrifugation, disrupted by sonication and treated with H₂DCFDA fluorescent probe as described in [3]. The emitted fluorescence values were normalized to the respective protein concentration that was determined as described by [55].
RNA isolation and cDNA synthesis

*Leptospirillum* sp. CF-1 was grown until the late exponential phase. Cells were harvested by centrifugation at 8000 ×g for 15 min and washed once with acid water (10 mM H₂SO₄) and twice with 10 mM sodium citrate pH 7.0. Washed cells were suspended in 9 K BR medium (without Fe₂SO₄) and incubated with 5 mM K₂CrO₄ for 15 or 30 min, or 5 mM H₂O₂ for 30 min. Cells were collected by centrifugation at 8000 ×g for 10 min, and washed twice with 10 mM sodium citrate pH 7.0. RNA was isolated using the RNeasy Mini Kit (Qiagen). DNA was removed by DNase I treatment (New England Biolabs) according to the manufacturer’s instructions. cDNA synthesis was carried out with the AffinityScript qPCR cDNA Synthesis kit (Agilent Technologies). The reaction mixture of 20 µl contained First Strand master mix, 0.1 µg/µl random primers, Affinity Script RT/RNase Block enzyme mixture and 1 µg RNA. cDNA synthesis was carried out at 25 °C for 5 min, 42 °C for 15 min and then the enzyme was inactivated at 95 °C for 5 min. cDNA was stored at −80 °C until further use.

### Determination of gene co-transcription of och operon:

#### RT-PCR reaction

Primers for PCR reactions were designed using the available gene sequences of *Leptospirillum* sp.CF-1 [9]. The GoTaq® DNA Polymerase kit (Promega) was used for PCR amplification according to the provider’s instructions. The characteristics of each primer pair and size of amplicons are shown in Table 2.

#### Determination of relative levels of RNA: quantitative PCR reaction

Primers of *Leptospirillum* sp. CF-1 designed for qPCR reactions are shown in Table 2. The KAPA SYBR FAST qPCR kit (Kapabiosystems) was used for qPCR amplification according to the manufacturer’s instructions. The qPCR conditions were an initial denaturation at 95 °C for 5 min, followed by 40 cycles of denaturation (95 °C for 30 s), annealing (60 °C for 20 s) and extension (72 °C for 10 s). All reactions were performed in a StepOne Real-Time PCR system (Applied Biosystems). The relative abundance of each gene versus a constitutively expressed gene (16S rRNA gene) was determined by the comparative Ct (ΔΔCt) method.

#### In-silico analysis of gene products

Predicted amino acid sequences derived from genes were used to perform a BlastP [56] search of the NCBI non-redundant data base. Only the best hits were accepted as evidence of putative orthologs. Translated proteins were further characterized using bioinformatics tools: primary structure similarity relations were determined using ClustalW 1.8 [57], structural motif predictions were

### Table 2

| Gene                     | Sequence                        | Tm  | Amplicon (bp) |
|--------------------------|---------------------------------|-----|---------------|
| Cloning                  |                                |     |               |
| ABH19_09590 (F)          | ATG AAC ACA CAC TGC CCG GAC AC | 61  | 2469          |
| ABH19_09590 (R)          | CTTC ACC GCG GGC CAC ATT        |     |               |
| RT-PCR                   |                                |     |               |
| ABH19_09575 (F)          | GCAG TAA GAG AGG AGA CAC       | 61  | 587           |
| ABH19_09585 (R)          | TTG TAC GGG GTC GTC GTG         |     |               |
| ABH19_09585 (F)          | CTG GAC TCA CTA CAT GCA AGA GT | 60  | 294           |
| ABH19_09585 (R)          | GAT TAC CGT CTT TCC TCC         |     |               |
| ABH19_09590 (F)          | ACC TCA TCG AAA ACG AAT GG     | 61  | 388           |
| ABH19_09590 (R)          | GAT ACG GCC CTC TTT TTT CAC    |     |               |
| ABH19_09595 (F)          | GTC GCT TGA GAT GAT GTG         | 60  | 392           |
| ABH19_09600 (R)          | ACA ATC GGG CAA GGA AAG AT     |     |               |
| ABH19_09600 (F)          | GTG CCG GAA GAG GAAT TT        | 60  | 286           |
| ABH19_09605 (R)          | CAG ACG TAC ACG ACC CC         |     |               |
| RT-qPCR                  |                                |     |               |
| ABH19_09585 (F)          | ATC ATC TAC ACG GGA GCC TTT    | 60  | 110           |
| ABH19_09585 (R)          | GTC AGG GCA ATG GAA AGA GT     |     |               |
| ABH19_09590 (F)          | ACA ACG TTT GGA ACC GAT GG     | 60  | 105           |
| ABH19_09590 (R)          | GAT CTT CTT CTA GAG ATG        |     |               |
| ABH19_09595 (F)          | ACC TGG AGT GTG CTG GTG         | 60  | 111           |
| ABH19_09595 (R)          | GTG ACT TCT CTC CAG TCT        |     |               |
| ABH19_09600 (F)          | GTC CCG TCT TGT TCG CAG CT     | 60  | 104           |
| ABH19_09600 (R)          | CC GTC TCC CAC AAG GAT TTT     |     |               |

(F) Forward, (R) Reverse
were determined using Prosite [58] and peptide domain predictions were determined using ProDom [59]. Theoretical isoelectric points (pI) and molecular weights (MW) were predicted using Compute pI/MW [60], whilst bacterial protein subcellular localization was predicted using PSORTb v3.0 [61].

Statistical analysis
Statistical analysis was performed using the one-way ANOVA test followed by Tukey’s, using GraphPad Prism 5. The differences were considered to be significant at P < 0.05.

Abbreviations
ROS: Reactive oxygen species; SOD: Superoxide dismutase; CAT: Catalase; DHAD: Dihydroxy acid dehydratase; DyP: Dyp-type peroxidases; CcP: Cytochrome c peroxidase; OD: Optical density; LB: Luria–Bertani; H2DCFDA: 2′,7′-Dichlorodihydrofluorescein diacetate.

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Author contributions
JRA and MR designed the studies and performed the experiments. JRA and GL wrote the paper and generated the figures. RCh and AF contributed to the discussion of results and to the revision of the manuscript. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

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