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1 Light regulation of resistance to oxidative damage and magnetic crystal biogenesis in

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Abstract

Light–oxygen–voltage (LOV) proteins are ubiquitous photoreceptors that can interact with other regulatory proteins and then mediate their activities, which results in cellular adaptation and subsequent physiological changes. Upon blue-light irradiation, a conserved cysteine (Cys) residue in LOV covalently binds to flavin to form a flavin–Cys adduct, which triggers a subsequent cascade of signal transduction and reactions. We found a group of natural Cys-less LOV-like proteins in magnetotactic bacteria (MTB) and investigated its physiological functions by conducting research on one of these unusual LOV-like proteins, Amb2291, in *Magnetospirillum magneticum*. In-frame deletion of *amb2291* or site-directive substitution of alanine-399 for Cys mutants impaired the protective responses against hydrogen peroxide, thereby causing stress and growth impairment. Consequently, gene expression and magnetosome formation were affected, which led to high sensitivity to oxidative damage and defective phototactic behaviour. The purified wild-type and A399C-mutated LOV-like proteins had similar LOV blue-light response spectra, but Amb2291^{A399C} exhibited a faster reaction to blue light. We especially showed that LOV-like protein Amb2291 plays a role in magnetosome synthesis and resistance to oxidative stress of AMB-1 when this bacterium was exposed to red light and hydrogen peroxide. This finding expands our knowledge of the physiological function of this widely distributed group of photoreceptors and deepens our understanding of the photoresponse of MTB.

Key words: LOV, magnetotactic bacteria, Amb2291, oxidative stress, reactive oxygen species.

52 **Key points**

53 We found a group of Cys-less light–oxygen–voltage (LOV) photoreceptors in magnetotactic bacteria,
54 which prompted us to study the light-response and biological roles of these proteins in these
55 non-photosynthetic bacteria.

56

57 The Cys-less LOV-like protein participates in the light-regulated signalling pathway and improves
58 resistance to oxidative damage and magnetic crystal biogenesis in *Magnetospirillum magneticum*.

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60 This result will contribute to our understanding of the structural and functional diversity of the
61 LOV-like photoreceptor and help us understand the complexity of light-regulated model organisms.

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Introduction

Most bacteria must cope with frequent changes in their environment; hence, they developed various mechanisms to sense and respond to external stimuli for growth and survival. Bacteria can use light as a source of energy when exposed to it, but they can also be damaged by ultraviolet (UV) radiation or light-generated reactive oxygen species (ROS) (Metz et al. 2012). Different types of photoreceptors, such as light-oxygen-voltage (LOV) sensing protein, photoactive yellow protein, phytochrome, sensor of blue light using flavin adenine dinucleotide, cryptochromes, and sensory rhodopsin, are involved in bacterial photosensing (Braatsch and Klug 2004; Losi 2007; Purcell and Crosson 2008; Spudich 2006; Wilde and Mullineaux 2017).

Photoreceptor LOV belongs to a subset of the Per–Arnt–Sim (PAS) family that utilizes internally bound flavin chromophores to sense changes in blue light or redox state (Pudasaini et al. 2017; Taylor and Zhulin 1999; Wu et al. 2013). The flavin chromophore binds non-covalently to LOV in a dark environment. Flavin photon absorption results in the formation of a flavin–C4(a)–cysteinyl adduct with a conserved cysteine (Cys) residue on the PAS-motif, thereby initiating a cascade of structural rearrangements within the LOV core, which are propagated to the domain boundaries (Pudasaini et al. 2015; Salomon et al. 2000). LOV proteins are widely distributed in bacteria. The first bacterial LOV protein from *Bacillus subtilis*, YtvA (Akbar et al. 2001), was initially reported as a positive regulator of general stress response (GSR). It binds to flavin mononucleotide (FMN) and undergoes a photocycle similar to that of the LOV domains of the plant phototropins (Losi et al. 2002). In *Dinoroseobacter shibae*, a blue-light photoreceptor of the LOV superfamily (*DsLOV*) regulates photopigment synthesis (Endres et al. 2015). Purcell et al. suggested that bacterial LOV domain proteins may function as redox sensors via flavin chromophore itself (Purcell et al. 2010). Furthermore, the data hint at a link between *RsLOV* in *Rhodobacter sphaeroides*, carbohydrate metabolism and chemotaxis, and the response of cells to photooxidative stress (Metz et al. 2012). The blue-light-modulating activity of *RsLOV* and *RsAppA* is intended to maximize photosynthesis under favourable conditions and to minimize photooxidative damage (Losi et al. 2015; Metz et al. 2012). In *Brucella abortus*, LOVhK is a cognate sensor histidine kinase of PhyR that functions as a stress sensor (and GSR pathway activator). GSR can be activated by red and blue lights and many other stimuli. Deletion of LOVhK severely impairs cell survival under defined conditions of oxidative and acid stresses (Kim et al. 2014). In addition,

blue-light inhibition of *Listeria monocytogenes* growth is mediated by ROS and is influenced by σ^B and the blue-light sensor Lmo0799 (O'Donoghue et al. 2016).

Previous research has found that replacement of the conserved adduct-forming Cys with alanine (Ala) abolishes the photochemical reaction of LOV (Salomon et al. 2000). Recently, Yee et al. found that by removing the conserved Cys, the LOV protein still undergoes light-induced photoreaction and signal transduction through the signalling capability of neutral semiquinone (NSQ), which is photoreduced from flavin (Yee et al. 2015). Interestingly, naturally occurring LOV proteins with proline (Pro) at the position of the adduct-forming Cys residue can still respond to light. Additional studies are needed to determine whether the light-induced naturally Cys-less LOV undergoing signal transduction is widespread. Notably, we found a group of magnetotactic bacteria (MTB) that possess Cys-less LOV-like proteins in which the Ala occupied the position of the adduct-forming Cys. Moreover, these LOV-like proteins contain an N-terminal putative transmembrane domain. MTB can synthesize intracellular membrane-enclosed, nanosized, and chain-arranged magnetite and/or greigite crystals called magnetosomes (Komeili 2007; Spring and Bazylinski 2002). Magnetosome chains are the magnetic sensors in MTB that act as an internal compass needle and help in the rapid orientation along the geomagnetic field, a behaviour known as magnetotaxis (Faivre and Schuler 2008; Frankel and Blakemore 1989; Komeili 2012). In addition to the magnetotaxis function, magnetosomes in *Magnetospirillum gryphiswaldense* MSR-1 exhibit peroxidase-like activity that reduces intracellular levels of ROS (Guo et al. 2012). Moreover, light irradiation can help *M. magneticum* AMB-1 cells eliminate intracellular levels of ROS (Li et al. 2017). Other studies also revealed the photoresponses of magnetotaxis in some MTB (de Melo and Acosta-Avalos 2017; Li et al. 2017; Qian et al. 2019; Shapiro et al. 2011; Zhou et al. 2011). It is unknown whether and what kind of photoreceptor is involved in this process. Analysis of AMB-1 genome reveals an unusual LOV-like photoreceptor, Amb2291, of which the expected conserved adduct-forming Cys at the location of 399 was actually replaced with Ala. To assess the function of the LOV-like protein, we mutated *amb2291* and analysed the effects of irradiation and hydrogen peroxide (H₂O₂)-induced oxidative stress (photo-oxidative stress) on cell growth and magnetosome biosynthesis in AMB-1 wild-type (WT) cells and *amb2291* mutants. Quantitative reverse transcription polymerase chain reaction (qRT-PCR) and dichlorofluorescein diacetate (DCFH-DA) methods were used to analyse the requirement of *amb2291* in regulating the

expression of antioxidant genes as well as magnetosome formation genes, which scavenge intracellular ROS under photooxidative stress conditions. In addition, deletion of *amb2291* or A399C substitution affects the phototactic behaviour of AMB-1, which further confirms the role of Amb2291 in antioxidant damage. Moreover, we purified the Amb2291 and A399C mutant proteins and analysed their photochemical properties. This study contributes to the elucidation of the photoreaction mechanism and deepens the understanding of the photoresponse of MTB.

Materials and Methods

Strains and culture conditions

Escherichia coli DH5 α was used for general cloning. The WM3064 of *E. coli* was used for intergeneric conjugation between *E. coli* and *M. magneticum*. *E. coli* strains were grown in LB medium supplemented with apramycin (50 μ g/mL) or kanamycin (50 μ g/mL). *M. magneticum* AMB-1 strains (ATCC 700264) were grown on modified enriched *Magnetospirillum* growth medium comprising 5 mL of 0.01 M ferric quinate, 0.68 g of potassium phosphate, 0.12 g of sodium nitrate, 0.74 g of succinic acid, 0.05 g of L-Cys-HCl, 0.2 g of polypeptone, and 0.1 g of yeast extract. In addition, 15 μ g/mL apramycin or kanamycin was used for plasmid selection (Matsunaga et al. 2005; Matsunaga et al. 1991; Yang et al. 2001). *M. magneticum* AMB-1 strains were cultured using a 100 mL Schott flask bottle or a 15 mL cell culture flask and incubated at 30°C.

Identification of the Amb2291 sequence

Amb2291 was identified in the *M. magneticum* AMB-1 genomic database BLAST (Matsunaga et al. 2005). Amino acid sequences were aligned using DNAMAN. Domain was analysed by SMART (<http://smart.embl-heidelberg.de>). The protein accession numbers were as follows: *M. magneticum* AMB-1 (WP_070108660.1, Amb2291); *M. magnetotacticum* MS-1 (KIL98983.1); *M. gryphiswaldense* MSR-1 (WP_106001729.1); *Magnetospirillum* sp. ME-1 (WP_085373061.1); *Magnetospirillum* sp. XM-1 (WP_068434569.1); *H. hochstenium* (WP_008581736.1); *Microcoleus vaginatus* (WP_006631677.1); *Populus trichocarpa* (XP_002315641.1); *Nectria* (XP_003042477.1); *Allium cepa* (ACT22763.1); *Asticcacaulis* sp. YBE204 (WP_023462395.1); *Azospirillum halopraeferens* (WP_029010434.1); *B. subtilis* (O34627.1); *Crocospaera watsonii* (WP_007308072.1); *Gluconobacter frateurii* (WP_023942432.1); *Granulicella mallensis* (WP_014265984.1); *B. abortus*

(WP_002971240.1); *Methylocystis rosea* (WP_018406994.1); *Roseobacter denitrificans* (WP_011568342.1); *Avena sativa* (PDB 2V0U); *Rhodobacter sphaeroides* (WP_009562854.1); and *Dinoroseobacter shibae* (WP_044027762.1).

Purification of protein

The conserved soluble fragment (residues 351–714) of *amb2291* or *amb2291*^{A399C} was cloned into the pET28a plasmid with N-terminal His6-SUMO-tagged fusion proteins. These proteins were expressed in *E. coli* BL21 (DE3) cells and purified as previously described (Li et al. 2019). The expression levels of Amb2291 variants were induced with 10 μ M isopropyl- β -D-1-thiogalactopyranoside for 16 h at 16°C in the dark. For cell lysis, the soluble extract was filtered and incubated with 5 mM FMN on ice for 20 min. After nickel chelate affinity chromatography, the proteins were further purified by size exclusion chromatography on Superdex 200 column in Tris buffer (10 mM Tris/Cl, 200 mM NaCl, pH 8.0). All experiments were carried out in the dark.

Genetic manipulation of AMB-1

All plasmids and primers used in the study are listed in Tables S1 and S2. Deletion of 2045 nucleotides of *Amb2291* (81-2125) was conducted by a CRISPR-Cas9 system using plasmid pCRISPomyces-2 (Cobb et al. 2015). The first step in constructing plasmid pCRISPR-sgRNA*amb2291* was to select the proper sgRNA sequence. sgRNA carrying the 20 nt specific targeting sequence was synthesized (Sangon Biotech, China) and inserted into the pCRISPomyces-2 through a *BbsI* site (Chen et al. 2018; Xie et al. 2014). A 2.0 kb HDR DNA fragment of the editing template was amplified from the purified genomic DNA of AMB-1 and inserted into the pCRISPR-sgRNA*amb2291* with the *XbaI* restriction enzyme.

The vector used for the allelic exchange of WT *amb2291* for *amb2291*^{A399C} was constructed as follows. PCR amplification of *amb2291* with 1.0 kb upstream and downstream sequences was conducted using Q5 high-fidelity polymerase (NEB, United States). The resulting 4.2 kb fragment was incorporated into pBBRI-mcs2. To prevent the sgRNA of the CRISPR plasmid from cutting the replaced *amb2291*^{A399C} in AMB-1, we inserted a nonsense mutation into the sequence of PAM and the sgRNA sequence in the *amb2291*^{A399C} fragment, so that sgRNA will not recognize and Cas9 will not cut *amb2291*^{A399C} in

AMB-1, guaranteeing the construction of the *amb2291*^{A399C} strain. Site-directed mutagenesis of the resulting plasmid by using the NEBuilder HiFi DNA assembly reaction protocol was conducted to introduce the A399C, PAM, and sgRNA sequences of the nonsense mutation (PAM: CCT to aCT and sgRNA sequence: GTTCACCCTGCCCCATGACCC to aTTCACCCTGCCCCATGACCC). The site-directed 4.2 kb fragment was inserted into the plasmid pCRISPR-sgRNA*amb2291* by using the same method as above.

All pCRISPR plasmids were transformed into the *E. coli* strain WM3064 by chemical transformation and transferred into AMB-1 through conjugation (Chen et al. 2018; Philippe and Wu 2010). All deletions were verified by PCR and sequencing.

Irradiation experiment and H₂O₂-induced oxidative stress in AMB-1

We investigated the effect of photooxidative stress on the growth and magnetosome biomineralization of AMB-1 cells. All cells were grown with the exposure to 20 $\mu\text{mol photons/m}^2/\text{s}$ from 625 nm or 450 nm LED light (Fig. S1a, Ouyingzhaoming, China) and were cultured in medium containing H₂O₂ at 0, 5, 7.5, 10, and 15 μM . In general, light exerts a certain decomposition effect on H₂O₂. Therefore, by using the H₂O₂ Quantitative Assay Kit (Sangon Biotech, China), medium with 15 μM H₂O₂ was used to analyse the H₂O₂ concentration changes versus the illumination time. The medium contains no iron ions to prevent the detection of H₂O₂ (Fig. S1b). We also detected the change in the H₂O₂ concentration in the dark, and the value remained unchanged within 20 h. The optical density (OD) and cellular magnetisms (Cmag) of exponentially growing cultures were measured at 600 nm under a 4.5 mT homogeneous magnetic field with a UV-visible spectrophotometer (Unico UV-2800, United States) and an electromagnetic system (Schüler et al. 1995; Zhao et al. 2007).

Quantitative RT-PCR assay

The procedure for qRT-PCR was based on a previously described general protocol (Chen et al. 2018). Analysis was performed with cells in the plateau phase. Total RNA was extracted using an RNAeasy Plus Minikit (Qiagen, Germany). Single-strand cDNA was synthesized with a reverse transcription kit (Takara, Japan). qRT-PCR assay was performed with the SYBR Green Real-time PCR Mix (Thermo Fisher Scientific, United States) on an ABI QuantStudio 6 Flex Real-time PCR System (Applied

Biosystems, United States). The relative expression level was calculated by the $2^{-\Delta\Delta C_T}$ method using the threshold cycle of 16S rRNA for normalization (Livak and Schmittgen 2001). The experiment was repeated at least thrice, with each sample run in triplicate. All primers used for probing each gene in this study are listed in Table S2.

Detection of ROS generation

Intracellular ROS levels were analysed using the DCFH-DA method (He and Hader 2002; Wang et al. 2013). After irradiation and H₂O₂ induction, the cells grew to the late stage of exponential growth. Then, 2 mL of bacterial culture was immediately added to 10 μ L of 10 mM DCFH-DA and incubated at 37°C for 30 min in the dark. The cells were centrifuged at 12,000 rpm for 6 min and washed thrice with a PBS buffer. The fluorescence of all samples was measured using a fluorescence spectrophotometer (Hitachi F-4500, Japan) at an excitation wavelength of 485 nm and emission wavelengths of 500–600 nm.

Motility and phototactic behaviour analyses of AMB-1

AMB-1 motility behaviour was analysed at a frame rate of 33 fps by using a long working distance 40 $\times$ Olympus objective. Cells (10 μ L) were observed on a glass slide with an Olympus microscope. Single cells were tracked, and the average swimming velocities were calculated using the MTrackJ plugin for ImageJ (Meijering et al. 2012; Murat et al. 2015; Zhang et al. 2017). Phototactic behaviour was quantitatively analysed using a modified mini MTB collection vessel as previously described (Li et al. 2017). The cultures (80 mL) with and without H₂O₂ induction were pooled into the reservoir and into the collecting tube. A microaerobic band formed after 30 min of growth in the dark prior to conducting the phototactic experiment. The upper interface of the microaerobic zone is parallel to the collection tube. Then, the device was placed in a cassette, where the collection tube was parallelly exposed to the light beam. Control experiments were conducted in the dark. After 30 min of collection, we measured the OD in the collecting tube and reservoir (OD_{collecting tube} and OD_{reservoir}, respectively). The ratio of (OD_{collecting tube} - OD_{reservoir})/OD_{reservoir} was used to determine the relative numbers of phototactic cells.

Spectrophotometry analyses

The absorbance spectra of purified Amb2291 WT and mutant proteins were determined from 700 nm to

250 nm at room temperature using a UV-visible spectrophotometer. Assays were recorded either in the dark or after exposure to a blue LED lamp (450 nm) or red LED lamp (625 nm) (Ouyingzhaoming, China). Fluorescence excitation and emission spectra were recorded at room temperature with a fluorescence spectrophotometer (Hitachi F-4500, Japan). The emission wavelength was 530 nm, and the excitation was at 450 nm.

Statistical analysis

All statistical analyses were performed using SPSS 22.0 (SPSS, IBM, United States). Hierarchical clustering analysis (HCA) was performed to analyse the values of OD₆₀₀ and Cmag. ROS values and the transcription-level results were analysed using two-tailed Student's t-test. Mann-Whitney U-test was used to analyse phototactic behaviour. One-way analysis of variance was used to investigate the difference in velocities under H₂O₂-induced oxidative stress. Each experiment was repeated three times or more. All data were expressed as the mean±standard deviation. In all cases, statistical tests were considered significant at $p<0.05$.

Results

Genomes of *Magnetospirillum* sp. encode a group of natural Cys-less LOV-like proteins

Genomic analysis allowed us to identify the *amb2291* as a CDS coding a LOV-like protein in AMB-1. This 713-residue protein contains the LOV-like domain encompassing PAS and PAC domains, histidine kinase, and histidine kinase-like ATPase (Fig. 1a). Notably, this LOV domain includes the key residues of the canonical LOVs, such as the signalling relevant glutamine (Gln), the asparagines that interact with the polar side of the isoalloxazine ring, and the arginines that interact with the phosphate moiety of the flavin (Fedorov et al. 2003), but it does not contain the Cys in the highly conserved motif GXNCRFLQ (Fig. 1b). The conserved Cys residue is involved in the photochemical generation of a flavin–Cys adduct upon blue-light absorption by the flavin in LOV photoreceptors (Briggs 2007; Christie et al. 1999). At the Cys position, an Ala substitutes for the strictly conserved Cys in the LOV domain of Amb2291 and the homologues from other *Magnetospirillum* sp. strains MS-1, MSR-1, ME-1, and XM-1 (Fig. 1b). In other naturally occurring Cys-less LOV, this position can be Pro, glycine (Gly), or lysine (Lys). This observation would challenge the photoresponse and physiological function of this group of natural Cys-less, LOV-like proteins in magnetospirilla. Moreover, all *Magnetospirillum* sp.

LOV-like full-length proteins contain transmembrane domains with 1 to 3 predicted transmembrane segments. To assess the function of this group of LOV-like proteins, we constructed an *amb2291* deletion mutant and a site-directed A399C mutant in AMB-1 (substitution of Cys for Ala 399) (as described in “Materials and Methods”).

Characteristics of recombinant Amb2291 and Amb2291^{A399C} proteins

To verify whether the Cys-less Amb2291 has the characteristics of LOV and to investigate the effect of light illumination of Amb2291 or variant, the soluble LOV-like sensory domain of Amb2291 and Amb2291^{A399C} from residues 351 to 714 (Fig. S2) were expressed in *E. coli* BL21 (DE3) as N-terminal His6-SUMO-tagged protein and purified via nickel chelate affinity, SUMO protease treatment and size exclusion chromatography. Recombinant proteins on denaturing gels of electrophoresis have a yellow colour and an apparent size of ~40 kDa (Fig. S3). Based on the peak position of the elution profiles of a Superdex 200-size exclusion column, Amb2291 and Amb2291^{A399C} displayed a molecular weight of approximately 80 kDa (Fig. S3), thereby suggesting the dimer property of these proteins in solution. Amb2291 and Amb2291^{A399C} exhibited a broad absorption band in the blue region that might be composed of sharp peaks near 449 and 475 nm and a shoulder near 420 nm, revealing a mixture of oxidized (450 nm) states of FMN. In addition, the absorption spectrum of Amb2291 had a double-peak structure at 361 and 375 nm, whereas the component at 361 nm was reduced to a minor shoulder in the absorption spectrum of Amb2291^{A399C}. The major UV-A absorption maximum occurred at 378 nm (Fig. 2a). Fig. 2b and c show the fluorescence emission and excitation spectra of the Amb2291 and Amb2291^{A399C} proteins, respectively. Compared with Amb2291, the fluorescence emission spectrum maximum of A399C mutant shows a redshift of 4 nm. The highest peaks of Amb2291 and Amb2291^{A399C} proteins are at 529 and 533 nm, respectively, and those in the excitation spectrum are at 467 nm. Thus, both Amb2291 and Amb2291^{A399C} demonstrated the characteristic properties for the LOV core, but a slight difference was obvious between the two chromophore binding domains.

Fig. 3 shows that dark-state LOV proteins contain an oxidized flavin cofactor that maximally absorbs blue light at 449 nm in the ground state. Blue-light irradiation results in the bleaching of the ground state and in the formation of a photoproduct, and absorption increased at 390 nm (Fig. 3a, b). The results show a faster reaction of the A399C mutant to blue-light illumination compared with the WT

strain (Fig. 3c), demonstrating the efficiency of adduct formation in the presence of Cys. Moreover, similar to the BAT P188C mutant of *H. hochstenium* (Yee et al 2015), Amb2291 and Amb2291^{A399C} appear to form the hydroquinone (HQ) directly with little NSQ intermediate observed on our observed time scale.

Effects of *amb2291* deletion on cell growth under conditions with photooxidative stress

To investigate the role of Amb2291, we performed a CRISPR-Cas9-based gene knockout experiment to construct in-frame deletion of *amb2291* as described in the “Materials and Methods” section. The effects of photooxidative stress on cell growth in AMB-1 WT cells and $\Delta amb2291$ mutant were analysed. AMB-1 cells were treated with different concentrations of H₂O₂ and immediately incubated for 20 h under multiple light ranges (red light 625 nm or blue light 450 nm, 20 μ mol photons/m²/s) or dark conditions. OD at 600 nm was measured to evaluate the oxidative stress effect on growth.

As shown in Fig. 4, the OD values can be grouped into clusters a, b, and c by using the HCA method when the squared Euclidean distance is equal to 4. Without irradiation in the dark, the OD of both AMB-1 WT cells and $\Delta amb2291$ mutant decreased with the increase in the H₂O₂ concentration in the growth media starting from 5 μ M (Fig. 4, black curves). The OD of AMB-1 WT cells drastically decreased from 10 μ M H₂O₂ and continued to decrease with increasing H₂O₂ concentration under blue-light irradiation, while the OD of $\Delta amb2291$ mutant decreased from 7.5 μ M (group c, blue curves in Fig. 4). The reason may be that blue light induces the formation of ROS that adds to the stress induced by H₂O₂ addition. However, red-light irradiation abolished the deleterious effect of H₂O₂ on the WT cells at all analysed concentrations (Fig. 4, red curve, group a). Interestingly, the sensitivity of the $\Delta amb2291$ mutant to H₂O₂ under the red-light irradiation showed no significant change compared with darkness treatment as well as WT cells under darkness (Fig. 4, red dashed-line, group b). Collectively, these results revealed that red light protects the AMB-1 WT cells from the deleterious effect of H₂O₂-generated oxidative stress, which was dependent on the function of Amb2291. For better understanding the response of AMB-1 to photooxidative stress, red light was the main light source in the following studies.

Expression of the *amb2291* gene in AMB-1 cells

To investigate the functional differences between Amb2291 and Amb2291^{A399C}, we constructed an allelic exchange of WT *amb2291* for the *amb2291*^{A399C} mutant. By qRT-PCR analysis, we observed that red-light irradiation increased the expression of the *amb2291* gene in WT cells, regardless of the H₂O₂ treatment (Fig. 5). The deletion of the *amb2291* gene from 81 to 2125 nucleotides substantially enhanced *amb2291* expression, especially under the conditions of red-light irradiation and the presence of H₂O₂ ($p < 0.01$), since the first 80 nucleotides of *amb2291* were still kept in $\Delta amb2291$, but the transcription level is higher than that of WT and *amb2291*^{A399C}. By contrast, site-specific substitution for the conserved Cys, *amb2291*^{A399C} mutation displayed the same phenotype of the WT strain. These results unveil the autoregulation of *amb2291* to a certain extent of the transcription level compared with the higher transcription level in $\Delta amb2291$ caused by the incomplete function of *amb2291*. In addition, the conservation of Ala 399 is not required for the autoregulation, although it is indispensable for the attenuation of photooxidative stress.

Effect of H₂O₂ and light illumination on magnetosome formation in AMB-1 cells

We analysed the effect of photooxidative stress on magnetosome formation by measuring Cmag in AMB-1 WT, $\Delta amb2291$, and *amb2291*^{A399C} mutant cells (Schüler et al. 1995; Zhao et al. 2007). By the HCA method, the results showed that Cmag can be divided into clusters (a) and (b) when the squared Euclidean distance is equal to 4 (Fig. 6a). The Cmag value in WT cells under red-light irradiation and with H₂O₂-induced oxidative stress formed cluster a, which was higher than all those under the other conditions. This result suggested that Amb2291, red light, and oxidative stress contributed to the maximum synthesis of magnetosomes.

We previously reported that magnetosome-formation-associated genes *mms13*, *mms6*, and *mmsF* significantly responded to visible light (Li et al. 2017). To investigate the mechanism by which photooxidative stress influences magnetosome formation in AMB-1 WT and mutant cells, we also performed qRT-PCR to detect the expression levels of *mms13*, *mms6*, and *mmsF*. After 20 h of illumination, the expression levels of *mms13* and *mmsF* were significantly upregulated in WT cells and *amb2291*^{A399C} mutants compared with their dark controls (Fig. 6b, d, $p < 0.05$). Importantly, red-light illumination led to significant upregulation of *mms13*, *mms6*, and *mmsF* in WT cells compared with $\Delta amb2291$ and *amb2291*^{A399C} strains (Fig. 6b, c and d, $p < 0.05$). H₂O₂ treatment also led to significant

upregulation of *mms13* in WT cells compared with $\Delta amb2291$ or *amb2291*^{A399C} mutant strains (Fig. 6b, $p < 0.05$). *mms6* expression was significantly downregulated in the H₂O₂ treatment of $\Delta amb2291$ strains compared with dark controls (Fig. 6c, $p < 0.05$). Red light improved the magnetosome synthesis of WT AMB-1 cells, which was related to the function of Amb2291.

***amb2291* regulates the expression of antioxidant genes and helps AMB-1 scavenge intracellular ROS**

At the end of exponential growth, after 20 h of incubation, we analysed the difference in the expression of the stress-related genes (*oxyR*, *sod*) between the dark and light groups with or without H₂O₂ by qRT-PCR. For WT and mutant cells, regardless of the H₂O₂ treatment, red-light illumination led to significant upregulation of the *oxyR* gene compared with their dark controls (Fig. 7a, $p < 0.01$). By contrast, red-light illumination significantly downregulated the expression of *sod* compared with the dark controls in both the WT cells and the *amb2291* mutants (Fig. 7b, $p < 0.05$). The H₂O₂ treatment without red-light illumination had no obvious effect on *oxyR* and *sod* expression. However, simultaneous red-light illumination and H₂O₂ treatment increased the expression of *oxyR* and *sod* only in the WT strain (Fig. 7, $p < 0.05$).

We further examined the involvement of the *amb2291* gene in controlling intracellular ROS levels by the DCFH-DA method (Fig. 8). The ROS levels of WT cells under photooxidative stress conditions were significantly lower than those of mutant cells ($p < 0.01$), whereas the ROS level significantly increased ($p < 0.01$) in mutant cells under light irradiation compared with those in the dark. The *amb2291* gene helps AMB-1 eliminate intracellular ROS under oxidative stress through the light regulation system.

***amb2291* affects the phototactic behaviour under photooxidative stress**

We analysed the difference in the phototactic behaviour in AMB-1 WT cells and mutants. The phototactic behaviour was quantitatively analysed by a modified mini MTB collection vessel as previously described (Li et al. 2017). To prevent the difference in phototactic behaviour caused by H₂O₂ on the state of bacterial movement, we first examined the speed of bacteria movement under different H₂O₂ concentrations. Low H₂O₂ concentrations (10, 20, 30, and 40 μ M) exhibited

insignificant effects on the velocities of AMB-1 WT cells and mutants. However, with increasing H₂O₂ concentration, the velocities of all cells showed a consistent downward trend (Fig. S4). Thus, we studied the difference in phototactic behaviour between the WT and mutant strains under 0, 10, and 20 μ M H₂O₂ treatments.

Phototrophic behaviour was observed only with red light for all three strains analysed. Phototactic behaviour of the mutant strains all decreased with the increasing concentration of H₂O₂ (Fig. 9, $p < 0.05$). Moreover, the photoresponse of WT cells was significantly stronger than that of $\Delta amb2291$ or $amb2291^{A399C}$ mutants after adding H₂O₂ (Fig. 9, $p < 0.05$). When H₂O₂ was not added, all three strains exhibited a photoresponse. Amb2291 is likely involved in the reduction of the toxicity of H₂O₂ to cells and helped WT cells to actively swim towards light.

Discussion

Flavin-based photoreceptor proteins of the LOV superfamily are ubiquitous in plants, algae, fungi, and prokaryotes (Losi et al. 2015). During photoreaction, the non-covalently bound flavin chromophore forms a covalent adduct with a conserved Cys residue in LOV under blue light to initiate signal transmission to various output domains (Losi 2007). Current studies have reported that some LOV photoreceptors apparently retained biological activity even after the substitution of the Cys thiol for non-reactive side chains (Aihara et al. 2012; Chen et al. 2010; Okajima et al. 2012; Vaidya et al. 2011; Yee et al. 2015). In GenBank, 70 LOV-like proteins contain the strictly conserved LOV domain, except that the adduct-forming Cys is replaced with other residues, including Ala, Pro, Gly and Lys (Yee et al. 2015). In the present study, a group of LOV-like proteins from *Magnetospirillum* sp. in which the conserved Cys residue is substituted with Ala were reported. The Amb2291 and Amb2291^{A399C} mutant proteins in *M. magneticum* AMB-1 were purified. Accordingly, the pale yellow Amb2291 and Amb2291^{A399C} purified from overexpression in *E. coli* indicate the binding of FMN to the chromophore domains and potential partial reduction of FMN, regardless of the presence or absence of the conserved Cys. Combined with Fig. 2, these results demonstrate the characteristics of LOV-like proteins. Then, we examined the spectral change of Amb2291 and Amb2291^{A399C} under blue-light illumination to analyse the photoreaction. Consequently, a faster reaction of Amb2291^{A399C} to blue light was observed compared with the WT strain, which shows that Cys is an effective electron donor to the photoexcited

flavin. Despite of the change in a prolonged illumination studied here cannot be well compared with those in a few minutes, FMN-bound Amb2291 indeed goes through photoreaction as a response to blue light by some mechanism. As the spectrum does not show any change under the same dose of red light (Fig. S5), it demonstrates that the effect of photodamage is negligible. In a conventional LOV photocycle, blue-light absorption drives the formation of a covalent adduct between Cys and flavin, promoting rotation of a conserved glutamine residue to initiate a cascade of structural rearrangements within the LOV core that are propagated to the domain boundaries (Zoltowski and Gardner 2011) and then generating downstream signalling responses. Signal transduction through flavin photoreduction to the NSQ was recently found to be a general mechanism in natural Cys-less, LOV-like regulators. The WT BAT of *H. hochstenium* shows properties similar to LOV proteins, reducing slowly to the NSQ upon blue-light illumination. However, when the WT BAT is converted to a traditional LOV mechanism by mutating the active-site Pro residue to Cys, the BAT P188C mutant shows a faster reaction to blue-light illumination, appears to the fully reduced HQ with little NSQ intermediate observed in their designed time scale (Yee et al. 2015). Interestingly, the absorption changes measured for Amb2291 and Amb2291^{A399C} on exposure to blue light are similar to those of the BAT P188C variant. Although photoreduction of the Cys-less Amb2291 is inefficient, the FMN-bound protein readily undergoes complete chemical reduction to the fully reduced HQ, with little NSQ intermediate observed on the carried-out time scale (Fig. 3a). Moreover, the aromatic residues of phenylalanine and tyrosine close to the FMN-binding domain motif, as revealed in the sequence analysis in Fig. 1, may generate reductive quenching, resulting in a short excited-state life time of Amb2291, which exhibits inefficient photoreduction (Yee et al. 2015).

To analyse the biological functions of *amb2291* under photooxidative stress, we used the CRISPR technique to construct $\Delta amb2291$ and *amb2291*^{A399C} mutants. Growth and magnetosome biosynthesis of AMB-1 WT cells were better than that of $\Delta amb2291$ or *amb2291*^{A399C} mutants. Interestingly, we found that red light protects the WT cells from the deleterious effect of H₂O₂-generated oxidative stress, which depends on the function of Amb2291. Dual sensing of blue light and oxidative stress has been observed in *R. sphaeroides*, wherein the blue-light-modulating activity of RsLOV and RsAppA is intended to affect photosynthesis and minimize photooxidative damage (Losi et al. 2015; Metz et al. 2012). The structural biochemistry of a fungal LOV photoreceptor reveals an evolutionarily conserved

pathway that integrates light and oxidative stress (Lokhandwala et al. 2015). To some extent, the light modulating activity of *amb2291* affected magnetosome synthesis and minimized photooxidative damage. Red-light irradiation and H₂O₂ treatment upregulated the expression of the *amb2291* gene. In addition, the expression results in *amb2291* deletion strains revealed the autoregulation of the *amb2291* gene. Thus, Amb2291 played an important role in photooxidative stress.

Many environmental stimuli, including light, H₂O₂, and hyperthermia, generate high levels of ROS that can damage cellular DNA and proteins, perturb the normal redox balance, and cause oxidative stress (Lushchak 2001; Touati 2000). Therefore, organisms have developed methods to protect themselves from ROS, enabling cells to survive oxidative stress. For example, *oxyR* and *sod* are stress-related genes that regulate the transcription of defence genes via a rapid kinetic reaction in response to oxidative stress (Jo et al. 2015; Lu et al. 2018). In the present study, the expression levels of *oxyR* and *sod* genes were compared between dark and light conditions with or without H₂O₂ treatment. Compared with the mutant controls, simultaneous red-light illumination and H₂O₂ treatment increased the *oxyR* and *sod* expression levels only in the WT strains. This result is inconsistent with the expression of *oxyR* and *sod* genes in the WT strains reported by Li et al. (2017), which may be due to the differences among the concentrations of H₂O₂ and light treatment. Amb2291 is involved in the regulation of oxidative stress genes and is beneficial for resisting the damage caused by photooxidative stress. A previous study showed that singlet oxygen induces carotenoid biosynthesis in *Phaffia rhodozyma* and *Myxococcus xanthus* and is presumed to be light dependent (Hodgson and Murillo 1993; Schroeder and Johnson 1995). Further studies showed that carotenoids can protect bacteria from photooxidative stress damage (Glaeser and Klug 2005). Zeaxanthin accumulates only under photooxidative stress and plays an essential role in protecting the photosynthetic apparatus via the xanthophyll cycle (Demmig-Adams et al. 1999). The Amb2291 protein may also serve the function of both pigments to protect cells from photooxidative stress damage. In addition, the ROS level of WT cells was lower than that of mutant strains under the addition of H₂O₂, which is consistent with the notion that light helps AMB-1 cells eliminate intracellular ROS (Li et al. 2017). These findings further provide new evidence that Amb2291 reduces the increase in ROS through light regulation.

AMB-1 exhibited a photoresponse to red light (Fig. 9), which is consistent with our previous study

(Chen et al. 2011; Li et al. 2017). After H₂O₂ treatment, the number of *amb2291* deletion or *amb2291*^{A399C} mutant cells swimming towards light was significantly less than that of the WT cells. This finding might imply that red light under H₂O₂ conditions reduced the phototaxis of the mutant, which may be related to the increased cytotoxicity of H₂O₂ or ROS in the absence of functional Amb2291. Amb2291 is involved in light regulation to help eliminate ROS under oxidative stress, thereby creating the optimum niche for bacterial growth. It has been shown that Fe₃O₄ magnetosomes in MTB exhibit peroxidase-like activity, which can catalyse the photoFenton reaction according to the following equation to scavenge the intracellular levels of ROS (Guo et al. 2012): $\text{Fe}^{2+} + \text{H}_2\text{O}_2 = \text{Fe}^{3+} + \text{HO}\cdot + \text{OH}^-$ (Goldstein et al. 1993; Touati 2000). Moreover, light irradiation enhances this activity by promoting the synthesis of magnetosome and helps MTB eliminate intracellular ROS (Li et al. 2017). *amb2291* expression was significantly increased with red-light irradiation. Moreover, red-light illumination and H₂O₂ treatment significantly increased the expression of magnetosome formation genes (*mms13*, *mms6*, and *mmsF*) and antioxidative genes (*oxyR* and *sod*) in WT cells compared with those in $\Delta\textit{amb2291}$ and *amb2291*^{A399C} strains. Therefore, Amb2291 participates in the light-regulated signalling pathway and further enhances the synthesis of magnetosomes and resistance to oxidative damage, which helped clear intracellular ROS, thereby improving phototactic behaviour and bacterial growth.

Collectively, Amb2291 showed a similar LOV spectra based on blue-light irradiation but was unable to undergo the canonical photochemistry of LOV domains with the formation of a flavin–Cys covalent adduct. Upon red-light irradiation, Amb2291 improved the resistance to oxidative stress by some mechanism. Our understanding of the downstream physiological role of LOV protein is far less than that of its biochemical properties, because only several of LOV proteins' roles had been identified (Wu et al. 2013). Whether Amb2291 is directly involved in the red-light response needs further confirmation. We propose the following assumptions. First, structural prediction analysis found that Amb2291 has a specific extra membranous domain, while most LOV domains have no transmembrane structure in general, except the one that is found in the cytoplasm and membrane components in *R. sphaeroides* (Hendrischk et al. 2009). We are not sure whether the specificity of this protein structure plays a role. Second, studies have shown that the red/far-red-light-sensing bacteriophytochrome (BphP) can form an integrated signalling network with LOV in response to red, far-red, and blue light (Wu et al. 2013).

Strikingly, genomic analysis revealed the presence of BphP protein in AMB-1, named MagBphP, which is a red/far-red photoreceptor that mediates photosensory responses (Brutesco et al. 2017; Wang et al. 2019). We speculate that Amb2291 and the red/far-red-light-induced MagBphP may form an integrated signalling network; one of the biological significances of light-mediated behaviours in AMB-1 is to avoid environmental stress, such as oxidative stress. Fig. 10 shows the functional model of LOV-like Amb2291. The Cys-less LOV-like Amb2291 protein participates in the light-regulated signalling pathway and improves resistance to oxidative damage and magnetic crystal biogenesis in *M. magneticum*. However, the present work does not provide enough data to allow for a conclusion about the mechanism between Amb2291 and MagBphP signalling pathways. In addition, MTB usually live in aquatic environments and navigate towards sediment. They may move to find better living environments that contain trace oxygen through migration towards light (Wang et al. 2019). One of the major challenges that ancient life may have adapted to is ROS. The ancestral role of intracellular magnetosomes may have helped early life cope with oxidative stress through photoresponse (Lin et al. 2020). In short, Amb2291 acts as an LOV-like protein with a complex sensory role when cellular redox state and/or light were functional input signals.

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Compliance with ethical standards

Conflict of interests The authors declare that they have no conflict of interest.

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753 Figures

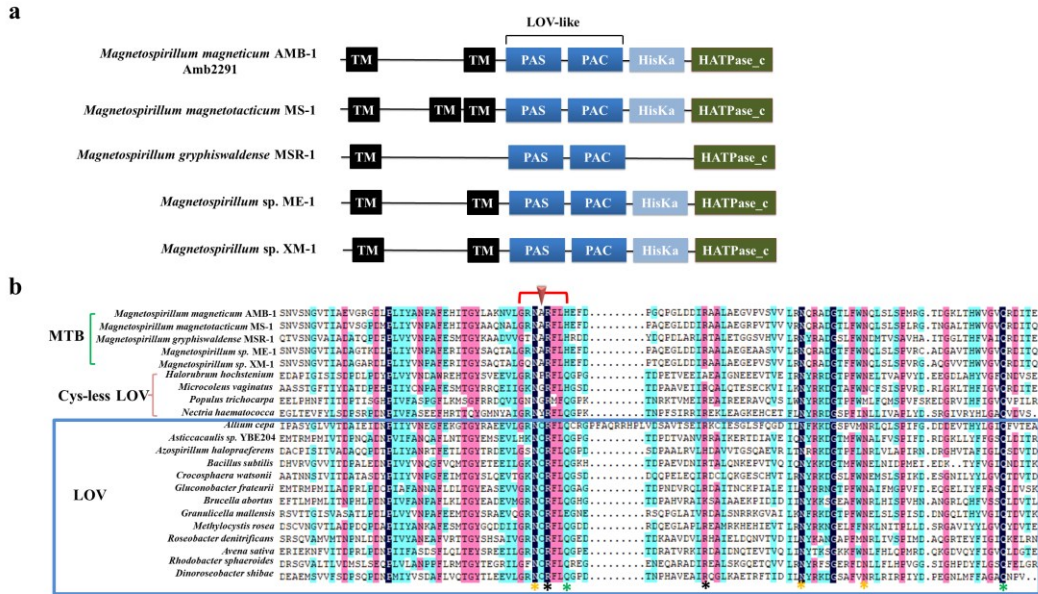


Fig. 1 Domain structures and amino acid sequence alignment of LOV proteins. **a** Domain structures of Amb2291 and other LOV-like proteins of MTB. TM, transmembrane segment; PAS, Per-Arnt-Sim domain; PAC, motif C-terminal to PAS motifs; HisKa, histidine kinase domain; HATPase, histidine kinase-like ATPase; The PAS-PAC domain is similar to LOV. **b** Alignment of selected LOV domains of MTB and similar regions of other Cys-less or LOV proteins. The red brace shows a conserved motif in LOV. The red triangle indicates the position of the conserved Cys involved in photoactivated FMN covalent binding.* Black shows the Arg residues that interact with the phosphate moiety of the flavin, * yellow shows the Asn residues that interact with the polar side of the isoalloxazine ring, and *green shows the Gln residues related to signalling.

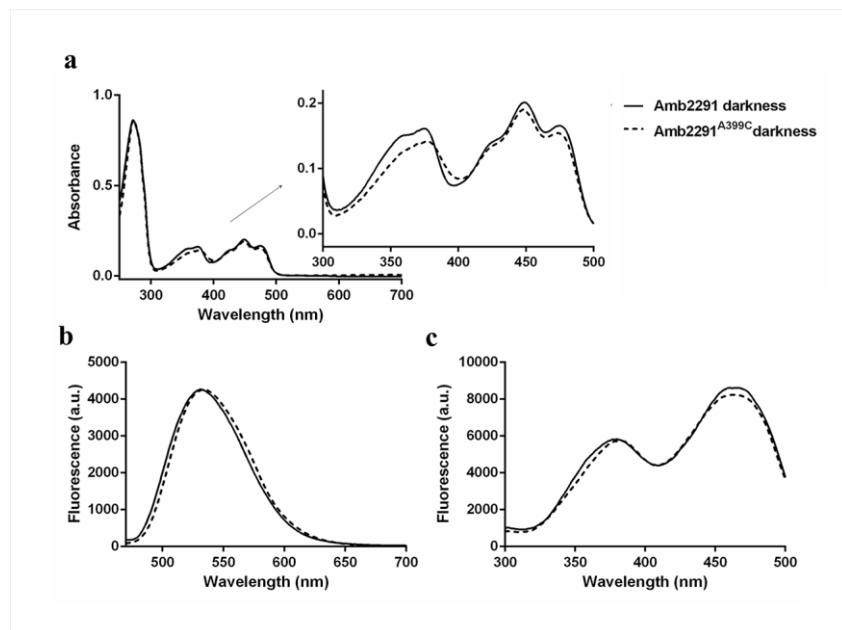
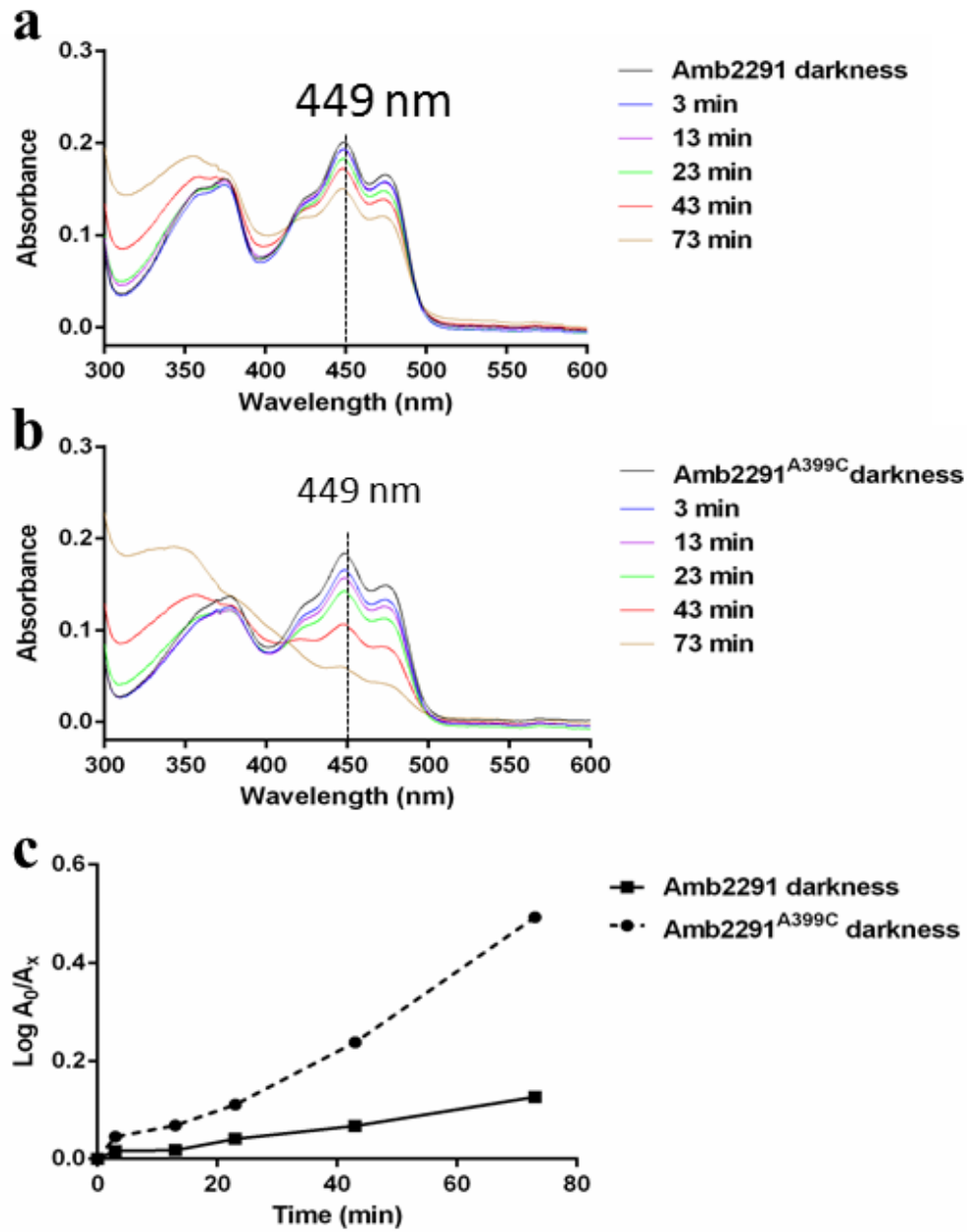


Fig. 2 Spectral properties of purified Amb2291 and Amb2291^{A399C} proteins. **a** Absorption spectra of Amb2291 (solid line) and Amb2291^{A399C} (dashed line) in the dark. **b** Fluorescence emission spectra for Amb2291 and Amb2291^{A399C}. Fluorescence excitation was monitored at 450 nm. **c** Fluorescence excitation spectra for Amb2291 and Amb2291^{A399C}. Emission was at 530 nm.



769

770 **Fig. 3** Photoreduction of purified Amb2291 and Amb2291^{A399C} proteins by blue-light irradiation.

771 Absorbance changes measured for Amb2291 (**a**) and Amb2291^{A399C} (**b**) on exposure to high-intensity

772 blue light (1700 $\mu\text{mol photons/m}^2/\text{s}$). **c** The kinetic data were obtained by following the absorbance

773 changes at 449 nm for Amb2291 and Amb2291^{A399C} proteins. A_0 represents the amount of unreacted

774 flavoprotein at time zero. A_x represents the corresponding amount of unreacted flavoprotein at time x.

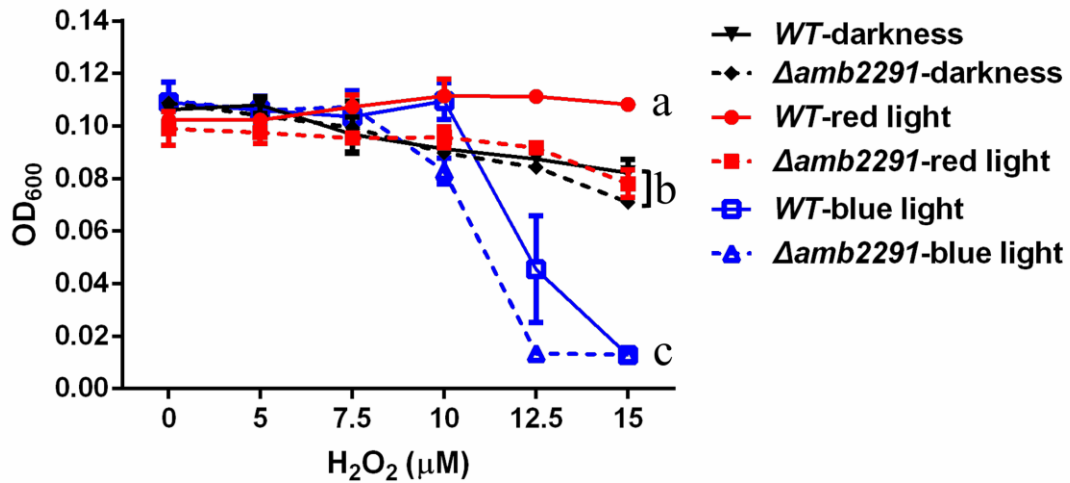


Fig. 4 Evaluation of the H₂O₂-generated oxidative stress effect on AMB-1 cells. The OD values of AMB-1 WT cells (solid lines) and $\Delta amb2291$ mutant (dashed lines) under irradiation or in the dark at different concentrations of H₂O₂ were the average of four independent experiments with standard deviations (error bars). The OD values are grouped into a, b, and c using the HCA method.

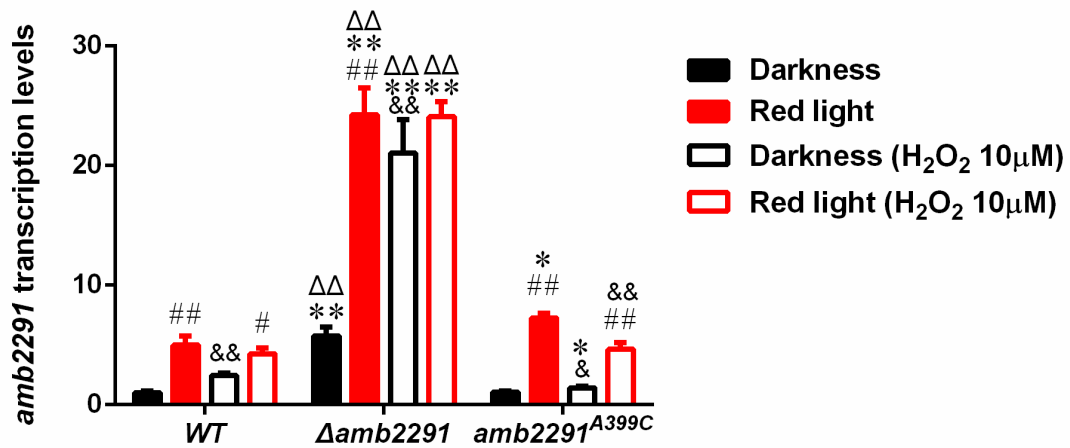


Fig. 5 Relative expression levels of the residual *amb2291* gene determined via qRT-PCR under red light and/or by adding H₂O₂ to the medium ($n=3$). # represents the statistical analyses performed between red light and dark. & shows the statistical analyses between the H₂O₂ group and their respective normal controls. * describes the statistical analyses between the mutants and WT control. Δ shows the statistical analyses between $\Delta amb2291$ and *amb2291*^{A399C} mutant cells.

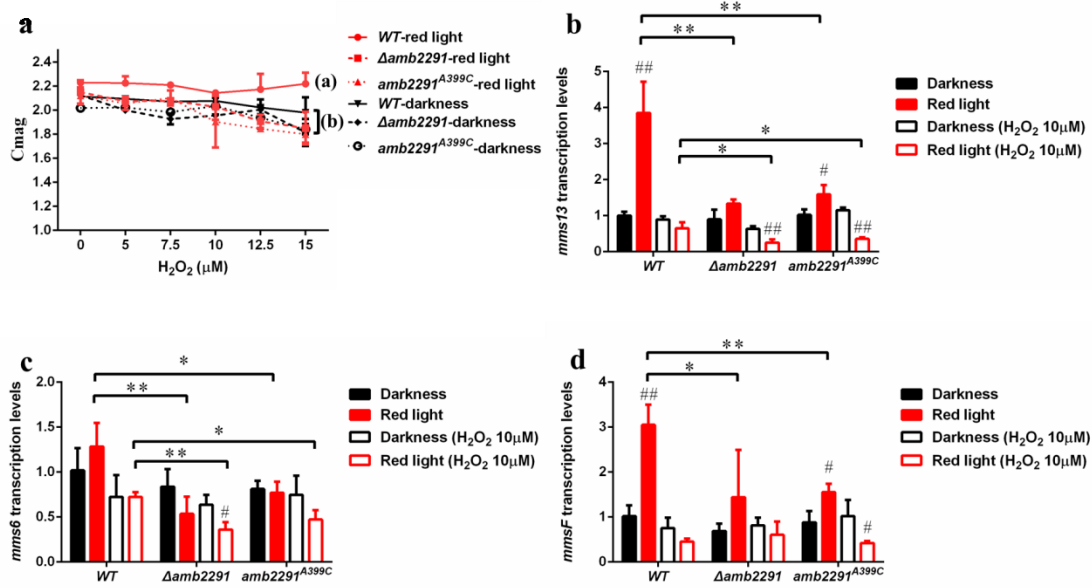


Fig. 6 Effects of *amb2291* deletion or mutant on magnetosome formation under photooxidative stress conditions. **a** Value of Cmag under red-light irradiation or dark at different concentrations of H₂O₂ in AMB-1 cells ($n=4$). The Cmag of each group is shown and analysed using HCA. Relative expression levels of *mms13* (**b**), *mms6* (**c**), and *mmsF* (**d**) determined via qRT-PCR under red light and/or by adding H₂O₂ to the medium ($n=3$). Level of significance of the differences observed between light and dark control is expressed as hashtag sign ($\#p<0.05$ and $\#\#p<0.01$). The significant differences between the two groups depicted by the lines in the graph are indicated by one or two asterisks ($*p<0.05$, $**p<0.01$).

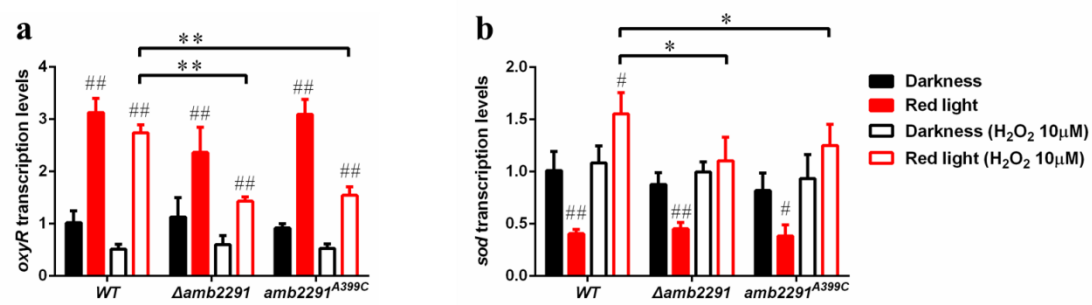


Fig. 7 Relative expression levels of *oxyR* (**a**) and *sod* (**b**) genes determined under red light and/or by adding H₂O₂ in the medium ($n=3$). The level of significance of the differences observed between light and dark control is expressed as hashtag sign ($\#p<0.05$ and $\#\#p<0.01$). The significant differences

between the two groups depicted by the lines in the graph are indicated by one or two asterisks
 (* p <0.05, ** p <0.01).

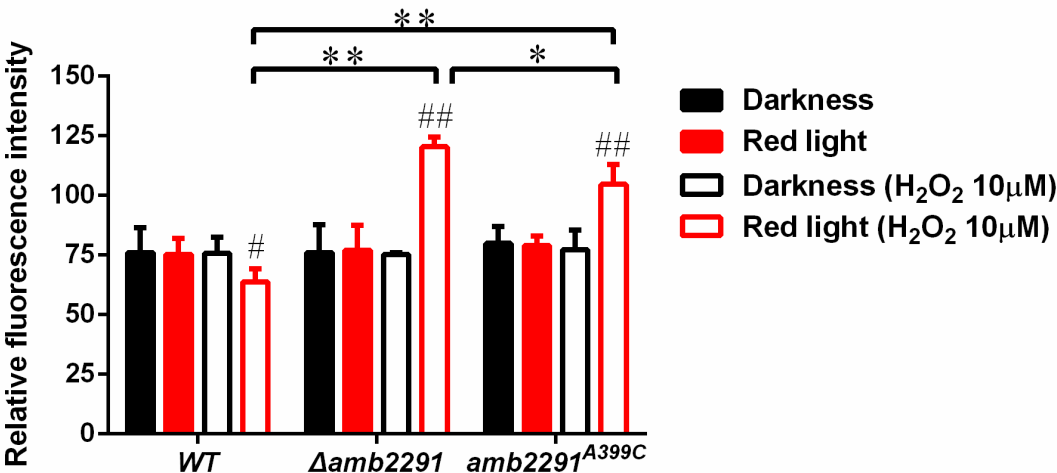


Fig. 8 Effects of Amb2291 on intracellular ROS level in AMB-1 cells ($n=6$). The level of significance of the differences observed between the light and dark control is expressed as hashtag sign (# p <0.05 and ## p <0.01). The significant differences between the two groups depicted by the lines in the graph are indicated by one or two asterisks (* p <0.05, ** p <0.01).

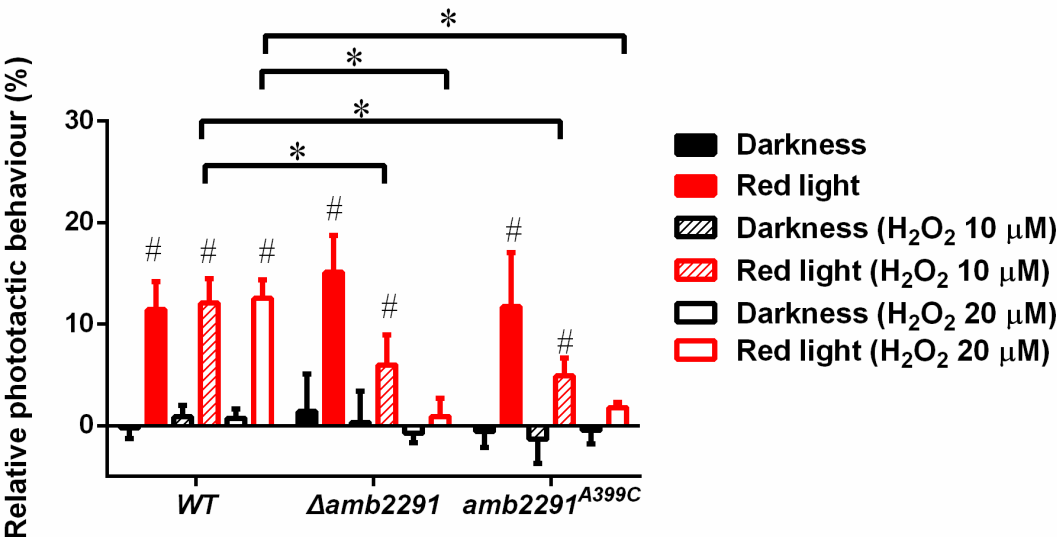


Fig. 9 Effects of *amb2291* deletion or site-directed mutagenesis on the phototactic behaviour of AMB-1 cells. Relative phototactic behaviour (%) of the WT and mutant strains ($n=4$). Level of the significance of the differences observed between light and dark control is expressed as a hashtag sign (# p <0.05). The significant differences between the two groups depicted by the lines in the graph are

indicated by asterisk (* $p<0.05$).

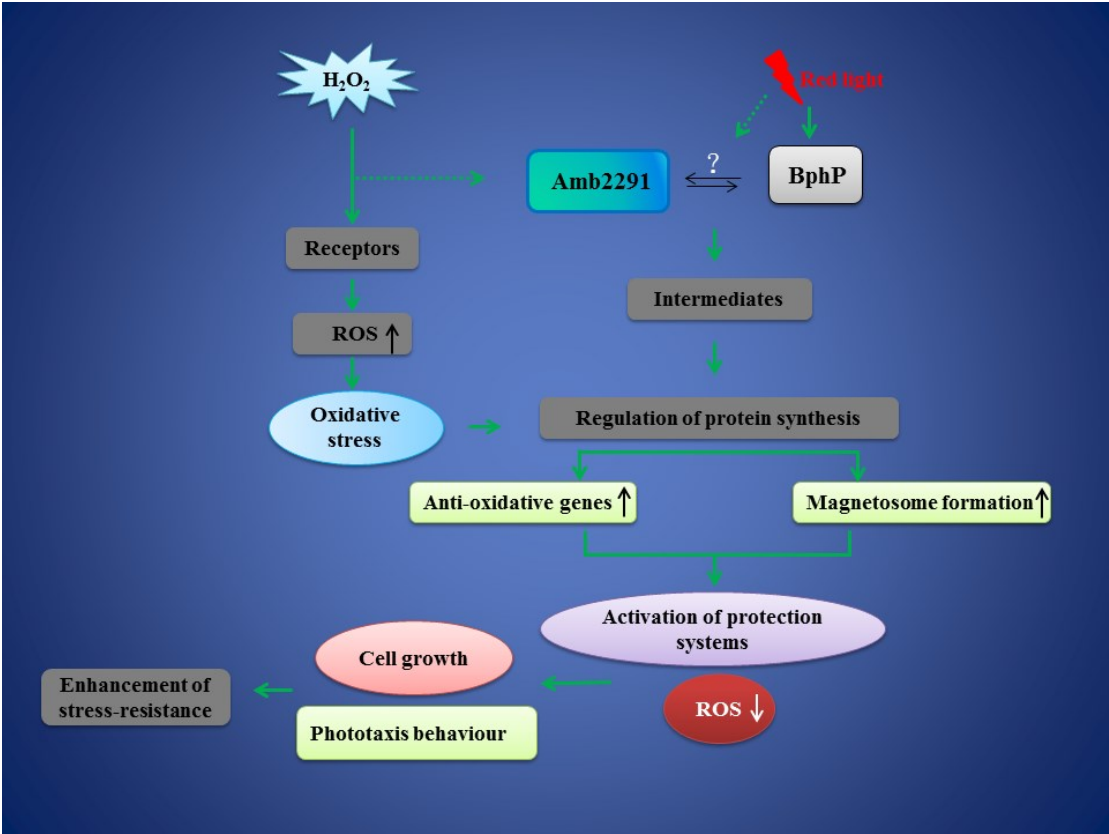


Fig. 10 Functional model of LOV-like Amb2291 protein. Red-light illumination and oxidative stress can regulate the LOV-like protein Amb2291 to enhance the expression levels of antioxidant genes and stimulate the formation of magnetosomes and then reduce intracellular ROS levels. The general function is to protect magnetotactic bacteria (MTB) cells from the deleterious effects of oxidative stress. The arrows show the red-light signal transduction pathway; Amb2291 may interact with red/far-red-light-induced bacteriophytochrome (BphP) by some mechanisms to form the signalling network response to red light. The signal transduction intermediates can be anti-oxidative genes, such as *oxyR* and *sod*, and magnetosome, the magnetite crystal in MTB.