

RpoH_{II} Activates Oxidative-Stress Defense Systems and Is Controlled by RpoE in the Singlet Oxygen-Dependent Response in *Rhodobacter sphaeroides*^{∇†}

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Photosynthetic organisms need defense systems against photooxidative stress caused by the generation of highly reactive singlet oxygen (¹O₂). Here we show that the alternative sigma factor RpoH_{II} is required for the expression of important defense factors and that deletion of *rpoH_{II}* leads to increased sensitivity against exposure to ¹O₂ and methylglyoxal in *Rhodobacter sphaeroides*. The gene encoding RpoH_{II} is controlled by RpoE, and thereby a sigma factor cascade is constituted. We provide the first in vivo study that identifies genes controlled by an RpoH_{II}-type sigma factor, which is widely distributed in the *Alphaproteobacteria*. RpoH_{II}-dependent genes encode oxidative-stress defense systems, including proteins for the degradation of methylglyoxal, detoxification of peroxides, ¹O₂ scavenging, and redox and iron homeostasis. Our experiments indicate that glutathione (GSH)-dependent mechanisms are involved in the defense against photooxidative stress in photosynthetic bacteria. Therefore, we conclude that systems pivotal for the organism's defense against photooxidative stress are strongly dependent on GSH and are specifically recognized by RpoH_{II} in *R. sphaeroides*.

Anoxygenic photosynthetic bacteria are widespread in aquatic habitats and show remarkable metabolic versatility that allows adaptation to changing environmental conditions, e.g., light intensity and oxygen concentration. *Rhodobacter* spp. have been studied extensively in regard to their adaptation to different environmental conditions (20, 31, 50). *Rhodobacter sphaeroides* can perform aerobic respiration. If the oxygen tension in the environment drops, photosynthetic complexes are formed, which are organized in an intracytoplasmic membrane system. This differentiation process is independent of light. In the presence of light and the absence of oxygen, anoxygenic photosynthesis enables *R. sphaeroides* to grow, while in the dark, anaerobic respiration or fermentation can be carried out. The redox-dependent formation of photosynthetic complexes is controlled by several regulatory systems in *R. sphaeroides* (31, 49). When *R. sphaeroides* grows at intermediate oxygen concentrations, blue-light illumination leads to a repression of photosynthesis genes (5, 32, 44). Most likely, this response reduces the risk of ¹O₂ formation. Under anaerobic conditions, the expression of photosynthesis genes is stimulated by light (5, 21).

Even when pigmented, *R. sphaeroides* grows well in the presence of oxygen and high light intensities, although bacteriochlorophylls and their precursors represent potent photosensitizers. Only recently, BChl *a* in the photosynthetic reaction center was shown to generate ¹O₂ (4, 46), which has also been

measured in intact *R. sphaeroides* cultures harboring or lacking carotenoids (16). Upon addition of the photosensitizer methylene blue, growth transiently slows down in the presence of light but then continues at rates almost as high as those of controls in the dark (16). This implies an adaptive response to ¹O₂. Although, as in other organisms, carotenoids play a major role in the defense against photooxidative stress in *R. sphaeroides* (8), the carotenoid content shows little change in the presence of ¹O₂ (16). However, the mRNA levels for a glutathione (GSH) peroxidase and a putative Zn-dependent hydrolase were strongly increased by ¹O₂ (16). The upregulation of the genes encoding RpoE and RpoH_{II} (previously called σ^{38}) and of some genes with putative RpoE and RpoH_{II} target sequences upon blue-light illumination suggested a role for these sigma factors in the organism's response to photooxidative stress (6). While RpoE belongs to the ECF (extracytoplasmic function) family of sigma 70 factors (36), RpoH_I and RpoH_{II} are members of the sigma 32 family of proteins (25). Dissociation of RpoE from the anti-sigma factor ChrR activates expression of genes involved in the response to ¹O₂ (2, 3). A putative RpoE target sequence upstream of the *rpoH_{II}* gene suggested that its expression is under RpoE control (3, 6).

Recently, we have identified cytoplasmic proteins of *R. sphaeroides* which show increased synthesis rates upon ¹O₂ exposure (17). While this response was RpoE dependent for some of the proteins, it was independent of RpoE for others. There was only a small overlap of RpoE-dependent genes, as detected by transcriptome analysis (3), and of genes encoding proteins whose expression is dependent on RpoE (17). We wanted to test the hypothesis that RpoH_{II} directly activates some of the RpoE-dependent genes and that RpoH_{II} plays an important role in the ¹O₂ response. To this end, we constructed and characterized an *rpoH_{II}* deletion mutant. Proteins exhibiting altered synthesis in the *rpoH_{II}* mutant were identified by

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TABLE 1. Strains and plasmids

Strain or plasmid	Description	Source or reference
Strains		
<i>E. coli</i>		
S17-1	<i>recA pro hsdR</i> RP4-2-Tc::Mu-Km::Tn7 <i>tra</i> ⁺ Km ^r Sp ^r	45
JM109	<i>recA1 supE44 endA1 hsdR17 gyrA96 relA1 thi(lac-proAB)</i>	New England Biolabs
<i>R. sphaeroides</i>		
2.4.1	Wild type	47
2.4.1(pRK415)	Wild type harboring pRK415, Tc ^r	This study
Δ <i>chrR</i>	<i>chrR</i> mutation in 2.4.1, Tp ^r	36
TF18	<i>rpoE chrR</i> mutation in 2.4.1, Tp ^r	43
2.4.1 Δ <i>rpoH</i> _{II}	2.4.1 <i>rpoH</i> _{II} :: Ω Sp ^r /St ^r	This study
2.4.1 Δ <i>rpoH</i> _{II} (pRK415)	2.4.1 Δ <i>rpoH</i> _{II} harboring pRK415, Tc ^r	This study
2.4.1 Δ <i>rpoH</i> _{II} (pRK2.4.1 <i>rpoH</i> _{II})	2.4.1 Δ <i>rpoH</i> _{II} harboring pRK2.4.1 <i>rpoH</i> _{II}	This study
2.4.1 Δ <i>rpoH</i> _{II} (pRK2.4.1 <i>rpoH</i> _{II} RSP0600)	2.4.1 Δ <i>rpoH</i> _{II} harboring pRK2.4.1 <i>rpoH</i> _{II} RSP0600	This study
Plasmids		
pPHU281	Suicide plasmid for <i>R. sphaeroides</i> , Tc ^r	23
pHP45 Ω	Ap ^r Sp ^r Sm ^r ; source of Ω Sp ^r /St ^r cassette	41
pRK415	Tc ^r	26
pPHU2.4.1 <i>rpoH</i> _{II} :: Ω Sp/St	pPHU281 with Sp ^r /St ^r cassette, flanked by the upstream and downstream region of <i>rpoH</i> _{II}	This study
pRK2.4.1 <i>rpoH</i> _{II}	pRK415 harboring a 1.0-kb fragment containing <i>rpoH</i> _{II} flanked by the 104-bp upstream and 11-bp downstream regions	This study
pRK2.4.1 <i>rpoH</i> _{II} RSP0600	pRK415 harboring a 1.8-kb fragment of <i>rpoH</i> _{II} and RSP0600 flanked by the 104-bp upstream region of <i>rpoH</i> _{II} and the 14-bp downstream region of RSP0600	This study
pDrive cloning vector	Ap ^r Km ^r	Qiagen

proteome analysis, and changes in expression of *rpoH*_{II} and RpoE-dependent genes were monitored in response to ¹O₂ exposure. It was previously shown that RpoH_{II} and RpoH_I are able to recognize similar promoter regions (19); however, RpoH_I does not complement RpoH_{II} under conditions of ¹O₂. Our data reveal that RpoH_{II} has a major role in ¹O₂ stress defense and acts downstream of RpoE in a sigma factor cascade.

MATERIALS AND METHODS

Bacterial strains and growth conditions. *R. sphaeroides* strains were grown at 32°C in minimal salt medium containing malate as the carbon source (14). Aerobic growth conditions with a concentration of 160 to 180 μ M of dissolved oxygen were established by gassing cultures with air in flat glass bottles or by continuous shaking of Erlenmeyer flasks at 140 rpm with a culture volume of 20%. In semiaerobic cultures, a volume of 80% in Erlenmeyer flasks and shaking at 140 rpm led to a dissolved-oxygen concentration of approximately 25 μ M. For heat shock experiments, aerobic cultures were shifted to 42°C in a water bath. When necessary, trimethoprim (50 μ g ml⁻¹) or spectinomycin (10 μ g ml⁻¹) was added to liquid and solid growth media, which contained 1.6% agar. Antibiotics were omitted from precultures, cultures, and agar plates used for *R. sphaeroides* strains during stress experiments and inhibition zone assays (see below) to ensure identical culture conditions. *Escherichia coli* strains were grown at 37°C in LB medium with continuous shaking at 180 rpm or on solid growth medium.

Construction of an *R. sphaeroides rpoH*_{II} deletion mutant. *R. sphaeroides* strain 2.4.1 Δ *rpoH*_{II} was generated by transferring the suicide plasmid pPHU2.4.1*rpoH*_{II}:: Ω Sp^r/St^r (Table 1) into *R. sphaeroides* 2.4.1 and screening for the insertion of the spectinomycin resistance/streptomycin resistance (Ω Sp^r/St^r) cassette into the chromosome by homologous recombination. Parts of the *rpoH*_{II} gene of *R. sphaeroides* 2.4.1, together with upstream and downstream sequences, were amplified by PCR using the oligonucleotides 2.4.1*rpoH*_{II}_knockout-up_EcoRI, 2.4.1*rpoH*_{II}_knockout-up_BamHI, 2.4.1*rpoH*_{II}_knockout-down_BamHI, and 2.4.1*rpoH*_{II}_knockout-down_SphI (see Table S1 in the supplemental material). The PCR fragments obtained, including the promoter region of *rpoH*_{II}, were cloned into pPHU281 (23) by using the appropriate restriction endonucleases. Then, the Ω Sp^r/St^r cassette obtained from plasmid pHP45 Ω (41) was inserted into the

BamHI restriction site to obtain the plasmid pPHU2.4.1*rpoH*_{II}:: Ω Sp^r/St^r. The plasmid pPHU2.4.1*rpoH*_{II}:: Ω Sp^r/St^r was transferred into *E. coli* strain S17-1 (45) and mobilized into *R. sphaeroides* wild-type strain 2.4.1 by biparental conjugation. Conjugants were selected on malate minimal medium agar plates containing 10 μ g spectinomycin ml⁻¹.

Southern blot analysis of chromosomal DNA isolated from spectinomycin-resistant and tetracycline-sensitive conjugants were carried out to confirm that the double crossover of the Ω Sp^r/St^r cassette into the *R. sphaeroides* chromosome had occurred. For this purpose, the linearized plasmid pPHU281, the *rpoH*_{II} upstream region also used for construction of the deletion mutant, and the Ω Sp^r/St^r cassette were radioactively labeled by nick translation and used as probes. By insertion of the Ω Sp^r/St^r cassette, 473 bp of the 879-bp *R. sphaeroides rpoH*_{II} gene was deleted. DNA of the *rpoH*_{II} mutant did not hybridize with linearized plasmid DNA of pPHU281, and positive signals of the appropriate sizes were obtained at the same positions by hybridization with the *rpoH*_{II} upstream region and the Ω Sp^r/St^r cassette.

Complementation of the *R. sphaeroides rpoH*_{II} deletion mutant 2.4.1 Δ *rpoH*_{II}. For complementation of 2.4.1 Δ *rpoH*_{II}, a 1.0-kb PCR fragment containing the entire *rpoH*_{II} gene, along with 104 bp of the upstream sequence and 11 bp of the downstream sequence, was amplified by using the oligonucleotides RSP0601_UP_EcoRI and RSP0601_DWN_EcoRI. The PCR fragment obtained was cloned into the pDrive vector (Qiagen, Hilden, Germany). Digestion of the pDrive vector containing the insert with PstI and XbaI, followed by cloning with the same restriction sites into plasmid pRK415 resulted in plasmid pRK2.4.1*rpoH*_{II}. This plasmid was subsequently transformed in *E. coli* S17-1 and conjugated with strain 2.4.1 Δ *rpoH*_{II} to obtain the complemented strain 2.4.1 Δ *rpoH*_{II} (pRK2.4.1*rpoH*_{II}). In addition, we complemented strain 2.4.1 Δ *rpoH*_{II} with a plasmid harboring a 1.8-kb fragment which contained the entire *rpoH*_{II} and RSP0600 genes, along with 104 bp of the upstream sequence of *rpoH*_{II} and 14 bp of the downstream region of RSP0600. The fragment was amplified using the oligonucleotides RSP0601_UP_EcoRI and RSP0600_DWN_EcoRI. This fragment was cloned into pRK415 and transferred to strain 2.4.1 Δ *rpoH*_{II} with the same strategy as that used for constructing 2.4.1 Δ *rpoH*_{II}(pRK2.4.1*rpoH*_{II}) to obtain strain 2.4.1 Δ *rpoH*_{II}(pRK2.4.1*rpoH*_{II}RSP0600).

Measurement of sensitivity to ¹O₂. For inhibition zone assays, 0.05-ml volumes of exponential-phase *R. sphaeroides* cultures (optical density at 660 nm of 0.5), grown under semiaerobic conditions, were diluted into 5 ml of prewarmed top agar (0.8% agar) and layered onto minimal malate medium agar plates. Filter

paper disks were placed on the surface of the agar, and 5 μ l of 10 mM methylene blue solution (Sigma-Aldrich, Seelze, Germany) was applied to the filter disks. Zones of inhibition were measured after incubation for 48 h at 32°C under a fluorescent tube (model NL 36 W/860 daylight; Spectralux Plus, Radium, Wipperfurth, Germany). Inhibition zone assays were also performed in the dark with filter paper disks containing 5 μ l of 10 mM methylglyoxal, 200 mM H₂O₂, and 500 mM paraquat to generate superoxide.

High light and photooxidative-stress conditions. High light and photooxidative-stress conditions were established as described previously (16). In brief, cultures were grown under semiaerobic conditions overnight to obtain pigmented cells. Cultures were diluted to an optical density at 660 nm of 0.2 and allowed to double once under aerobic growth conditions in darkened flat glass bottles. High light conditions were generated by illumination with 800 W m⁻² of white light, and for photooxidative stress, ¹O₂-producing methylene blue was added to liquid cultures at a final concentration of 0.2 μ M.

Radioactive labeling of proteins. Samples of 7 ml were retrieved from *R. sphaeroides* cultures, 15 μ Ci of L-[³⁵S]methionine (GE Healthcare, München, Germany) was added, and incubation was performed for 10 min under the experimental conditions described above. Samples were cooled on ice after incubation, and cells were harvested by centrifugation at 10,000 \times g for 10 min at 4°C and stored at -20°C until further processing.

Gel-based proteome analysis. Proteome analysis performed by two-dimensional (2D) gel electrophoresis followed the procedure described previously (17). In brief, for the extraction of soluble proteins, harvested cells were washed and disrupted by sonication. Intact cells and cell debris were removed by centrifugation, and the supernatant was used for ultracentrifugation at 100,000 \times g. The radioactive label was quantified in the colorless supernatant by adding 10- μ l aliquots to 1 ml of rotiszint scintillation cocktail (Roth, Karlsruhe, Germany) and measured in a Beckmann LS-6500 scintillation counter (Beckmann Coulter, Fullerton, CA). For 2D gel electrophoresis, protein samples containing 1.5 \times 10⁶ cpm were treated with RNase A (Qiagen) and RQ1 DNase I (Promega, Madison, WI) to remove nucleic acids. Proteins were precipitated using trichloroacetic acid, and the protein pellets were dried and then solubilized in sample buffer (17). Then, samples were applied to immobilized pH gradient strips (ReadyStrip; Bio-Rad, Hercules, CA). After isoelectric focusing, proteins were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis, and gels were fixed, dehydrated, dried, and exposed to phosphorimaging screens for 48 h. Phosphorimages were read with a Molecular Imager FX (Bio-Rad) set to a resolution of 100 μ m.

Protein spots on digital phosphorimages were compared using Delta2D version 3.3 software as described by the manufacturer (Decodon, Greifswald, Germany).

Extraction of RNA. Cell samples from growth experiments were rapidly cooled on ice and harvested by centrifugation at 10,000 \times g in a cooled centrifuge. For 5' rapid amplification of cDNA ends (RACE), total RNA was isolated using hot phenol and quantified by photometric analysis at a wavelength of 260 nm. Samples were treated with 1 U of RNase-free DNase I (Invitrogen) per 1 μ g RNA to remove contaminating DNA. After DNase I treatment, the RNA was purified by standard procedures using a mixture of phenol-chloroform-isoamyl alcohol and chloroform-isoamyl alcohol before being precipitated with sodium acetate and ethanol. For real-time reverse transcription-PCR (RT-PCR), total RNA was isolated by Total RNA isolation reagent (TRIR; ABGene, Epsom, United Kingdom) as described by the manufacturer. After extraction, DNase I treatment and purification of RNA samples were carried out as described above. Contamination by any remaining DNA was detected by PCR amplification of RNA samples, using primers targeting *gloB* (RSP0799) (see Table S1 in the supplemental material), as described previously (16).

5' RACE. For the determination of 5' mRNA ends using 5' RACE, 5 μ g of total RNA was reverse transcribed into cDNA by using 20 U of avian myeloblastosis virus reverse transcriptase (Promega) and 12.5 pmol of gene-specific primers (*gloA*-RACE-1, *gst*-RACE-1, RSP3075-RACE-1, and 2.4.1*rpoZ*-B [see Table S1 in the supplemental material]). RT was performed for 1 h at 50°C and then 15 min at 60°C with all primers in one reaction mixture. cDNA was then purified with a High Pure PCR product purification kit (Roche). The purified cDNA was poly(A) tailed using 20 U of terminal transferase (Fermentas). Poly(A) tailing was performed for 30 min at 37°C, and the terminal transferase was then heat inactivated at 70°C for 10 min. The poly(A)-tailed cDNAs served as templates for the first of two PCRs. The first PCR was performed using the oligo(dT) anchor primer and a gene-specific RACE-1 primer. For the second PCR, the PCR anchor primer and the gene-specific RACE-2 primer (nested PCR) were used. Resulting PCR products were cloned into the pDrive cloning vector (Qiagen) and sequenced with the respective gene-specific RACE-2 primer.

Real-time RT-PCR. Primers employed for analyzing the relative expression of target genes, using real-time RT-PCR, are listed in Table S1 in the supplemental material. For normalization of mRNA levels, the *rpoZ* gene, which encodes the ω subunit of RNA polymerase of *R. sphaeroides*, was used (39). Conditions for real-time RT-PCR were previously described in detail (16, 17). For real-time RT-PCR, a final concentration of 4 ng μ l⁻¹ of total RNA was applied using a one-step RT-PCR kit (Qiagen), and Sybr green I (Sigma-Aldrich) was added at a final dilution of 1:50,000 to the master mixture to detect double-stranded DNA. The relative expression of target genes was calculated relative to the expression of untreated samples and relative to that of *rpoZ* (40). PCR efficiencies were 2.02 for *rpoZ*, 1.98 for *ggt*, 2.31 for *gloB*, and 2.02 for *gpx* (16, 17). Additional PCR efficiencies were determined experimentally using serial dilutions of RNA, as follows: 2.04 for *rpoE*, 2.09 for *rpoH_{II}*, 2.0 for *rpoH_I*, 2.01 for *gloA*, 2.02 for *groEL*, 1.98 for RSP3075, and 2.0 for RSP3537.

Statistical analysis. Statistical analysis of the comparison of *C_p* (crossing point) values obtained for individual genes under different stress conditions, using real-time RT-PCR, was performed with Student's *t* test using Microsoft Excel 2003 (Microsoft, Redmond, WA). Significance levels (*P* values of ≤ 0.1 , ≤ 0.05 , and ≤ 0.01) are indicated in the figure legends. Statistically significant differences in protein synthesis were tested with the Student *t* test option available in Delta2D software. Proteins affected by ¹O₂ were included in the analysis if changes in protein synthesis were statistically significant as previously described (16).

RESULTS

RpoH_{II} contributes to the defense of *R. sphaeroides* against ¹O₂. We constructed an *rpoH_{II}* deletion mutant of *R. sphaeroides* as described in Materials and Methods. Inhibition zone experiments revealed a significantly higher sensitivity to ¹O₂ exposure for the *rpoH_{II}* mutant than for the isogenic wild-type strain 2.4.1. The diameter of the zone of inhibition observed for wild-type strain 2.4.1 was 2.6 \pm 0.1 cm (mean \pm standard deviation) (*n*, 3); and it was increased in the *rpoH_{II}* mutant to 3.2 cm (*n*, 3) when 10 mM methylene blue was applied in the presence of light for the generation of ¹O₂. Inhibition zones observed for the wild type and for the *rpoH_{II}* mutant that harbored plasmid pRK415 were not significantly different from those described above (see Table S2 in the supplemental material). The deletion of *rpoH_{II}* in strain 2.4.1 Δ *rpoH_{II}* was complemented in *trans* by the expression of *rpoH_{II}* on a low-copy-number plasmid yielding strain 2.4.1 Δ *rpoH_{II}*(pRK2.4.1*rpoH_{II}*). To control for polar effects of the *rpoH_{II}* deletion on the RSP0600 gene located downstream of *rpoH_{II}*, a second plasmid containing *rpoH_{II}* and RSP0600 was used for complementation of the *rpoH_{II}* mutant, resulting in strain 2.4.1 Δ *rpoH_{II}*(pRK2.4.1*rpoH_{II}*RSP0600). Inhibition zones of the two complemented *rpoH_{II}* mutants formed upon ¹O₂ exposure were not significantly different from those obtained with the wild type (see Table S2 in the supplemental material). The diameters measured 2.5 \pm 0.1 cm (*n*, 3) for the former and 2.4 \pm 0.1 cm (*n*, 3) for the latter strain. From these observations, we conclude that RpoH_{II} is involved in the response of *R. sphaeroides* to ¹O₂ and that the higher sensitivity of the *rpoH_{II}* mutant is not due to polar effects of the resistance cassette on the downstream gene. Further inhibition zone tests revealed that there were no significant differences in resistance to hydrogen peroxide or superoxide generated by the addition of paraquat between the *rpoH_{II}* mutant and the wild type (see Table S2 in the supplemental material).

The *rpoH_{II}* gene is triggered by exposure to ¹O₂. In order to assess in detail the factors that trigger the expression of *rpoH_{II}*, we used real-time RT-PCR to quantify the relative expression levels of the *rpoH_{II}* gene in wild-type and mutant strains under

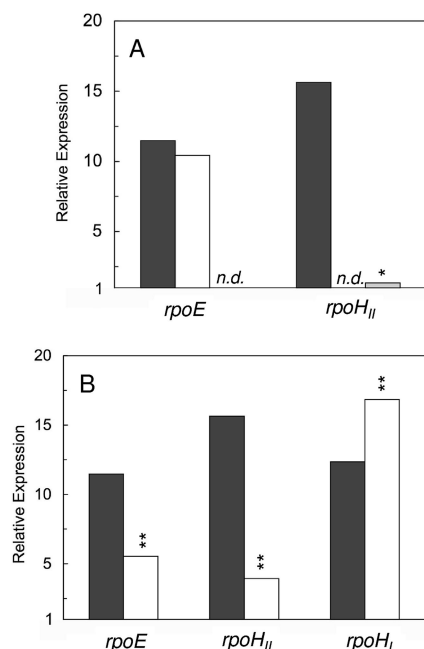


FIG. 1. Exposure to 1O_2 and heat shock affect relative expression levels of sigma factors $rpoE$ and $rpoH_{II}$. (A) Levels of relative expression are shown for sigma factors $rpoE$ and $rpoH_{II}$ in response to 1O_2 exposure in wild-type strain 2.4.1 (black bars), 2.4.1 $\Delta rpoH_{II}$ (white bars), and $rpoE$ *chrR* mutant strain TF18 (gray bar). (B) The expression levels of $rpoE$ and $rpoH_{II}$ were compared to that of $rpoH_I$ in wild-type strain 2.4.1 under conditions of 1O_2 exposure (black bars) and heat shock (white bars). Exposure to 1O_2 was performed for 7 min, and heat shock for 30 min at 42°C. Values for relative expression levels represent the increase in gene expression compared to that of the control at time point 0 min and were normalized to mRNA levels determined for $rpoZ$. Mean values for three independent experiments are shown, and statistically significant differences in expression for mutants relative to that of wild-type cultures were determined using Student's *t* test. Levels of significance are as follows: *, $P \leq 0.01$; **, $P \leq 0.05$. n.d., not detectable.

different growth conditions. A 15-fold increase in the $rpoH_{II}$ mRNA level was observed 7 min after the onset of 1O_2 exposure in the wild type but not in strain TF18, which lacks RpoE and its anti-sigma factor ChrR (Fig. 1A). This result is in agreement with the RpoE-dependent expression of $rpoH_{II}$ (3). Under the same conditions, the mRNA level of $rpoE$ was increased 10- to 12-fold in the wild type and $rpoH_{II}$ mutant, confirming earlier findings that $rpoE$ is triggered by 1O_2 exposure (3, 17).

RpoH_{II} is a member of the sigma 32 family of proteins, implying a role in heat shock response. Like RpoH_I of *R. sphaeroides*, RpoH_{II} can complement the temperature-sensitive phenotype of an *E. coli* $rpoH$ mutant (19). We therefore compared the mRNA levels of $rpoH_I$ and $rpoH_{II}$ and also of $rpoE$ after exposure to 1O_2 and heat shock (Fig. 1B). To verify that our growth conditions initiated a heat shock response, relative expression levels of the known heat shock gene *groEL* (25) were monitored. The *groEL* levels were increased eightfold in both the wild type and the $rpoH_{II}$ mutant after heat shock (data not shown). This result is in agreement with the heat shock response observed previously for *R. sphaeroides* (25) and with the finding that the $rpoH_{II}$ mutant is not heat

sensitive. The $rpoE$ mRNA level increased about 12-fold upon 1O_2 exposure and about 5-fold after heat shock (Fig. 1B). Under the same conditions, $rpoH_{II}$ mRNA levels increased by a factor of 15 in response to 1O_2 but only by a factor of 4 after heat shock. In comparison, the $rpoH_I$ mRNA levels were increased 12- and 17-fold in response to 1O_2 and heat shock, respectively. Like the response of $rpoH_{II}$ to 1O_2 , the response of $rpoH_{II}$ to heat shock was RpoE dependent. These data demonstrate a partial overlap of the 1O_2 response and the heat shock response in *R. sphaeroides*. The pronounced increase in $rpoH_{II}$ mRNA levels under photooxidative-stress conditions over those in response to heat shock hints at $rpoH_{II}$'s prominent role in response to 1O_2 exposure.

Deletion of $rpoH_{II}$ affects the synthesis of soluble proteins upon 1O_2 exposure. Protein synthesis patterns in strain 2.4.1 $\Delta rpoH_{II}$ under photooxidative stress were clearly different than the patterns observed in the wild type (Fig. 2; Table 2). Overall, most of the proteins (76%) characterized by the proteome analysis as RpoE dependent (category 1) (17) and conditionally RpoE dependent (category 2; less induction in a mutant lacking RpoE) showed dependency on RpoH_{II}, whereas RpoE-independent proteins (category 3) were not affected by the lack of RpoH_{II}.

Products of the genes *gloA*, *ggt*, *adh*, RSP2314, RSP3075, RSP3076, and RSP3164 with a predicted RpoH_{II} target sequence (6) did not show a significant response to 1O_2 in the $rpoH_{II}$ mutant (Table 2). In total, 25 proteins previously shown to be dependent or conditionally dependent on the presence of RpoE (17) were synthesized at lower or undetectable levels in the $rpoH_{II}$ mutant, as indicated by blue protein spots in superimposed 2D gels and decreased relative spot intensities (Fig. 2; Table 2). Other proteins depending on RpoH_{II} in category 1 are encoded by the genes *gst* (RSP1591), *eda*, *prxQ*, *rpsB*, RSP0423, and RSP2268. In category 2, proteins encoded by the genes *bfr*, *cysE*, *gloB* (RSP0799), *gloB* (RSP2294), *gst* (RSP1397), *pgl*, *pqqL* (RSP1096, RSP1097), RSP0663, RSP1415, RSP1507, and *catA* were significantly affected by the deletion of $rpoH_{II}$ (Fig. 2; Table 2). In the complemented strain 2.4.1 $\Delta rpoH_{II}$ (pRK2.4.1 $rpoH_{II}$), most RpoH_{II}-dependent proteins were synthesized in amounts that were similar to that of the wild-type strain 2.4.1. For Ggt, PqqL (RSP1097), and RSP1507, however, the amounts of synthesized proteins were restored to less than 50% of that of the wild type (Table 2). The amounts of proteins synthesized in strain 2.4.1 $\Delta rpoH_{II}$ (pRK2.4.1 $rpoH_{II}$ RSP0600), which also contains the RSP0600 gene downstream of $rpoH_{II}$, were similar to those of the wild-type strain, with the exceptions of Gst, RSP2314, and PqqL (RSP1096) (Table 2).

Only one protein that increased in synthesis upon 1O_2 exposure was encoded by a gene with a preceding RpoE target sequence: RSP1852, encoding a hypothetical protein related to UV endonucleases (3, 6, 17). A second gene, RSP1464 (*dsbG*, encoding a periplasmic disulfide isomerase), overlaps with RSP1465, which as yet has no function assigned and is preceded by a putative RpoE target sequence (17). Both proteins were synthesized in the $rpoH_{II}$ mutant when it was exposed to 1O_2 (Fig. 2; Table 2), which underlines their regulation in an RpoE-dependent but RpoH_{II}-independent manner.

Additional RpoE-dependent proteins not affected by the deletion of $rpoH_{II}$ were found in category 2 (encoded by RSP0355, RSP0558, RSP1410, *sucC*, *sucD*, and *hisH*). The

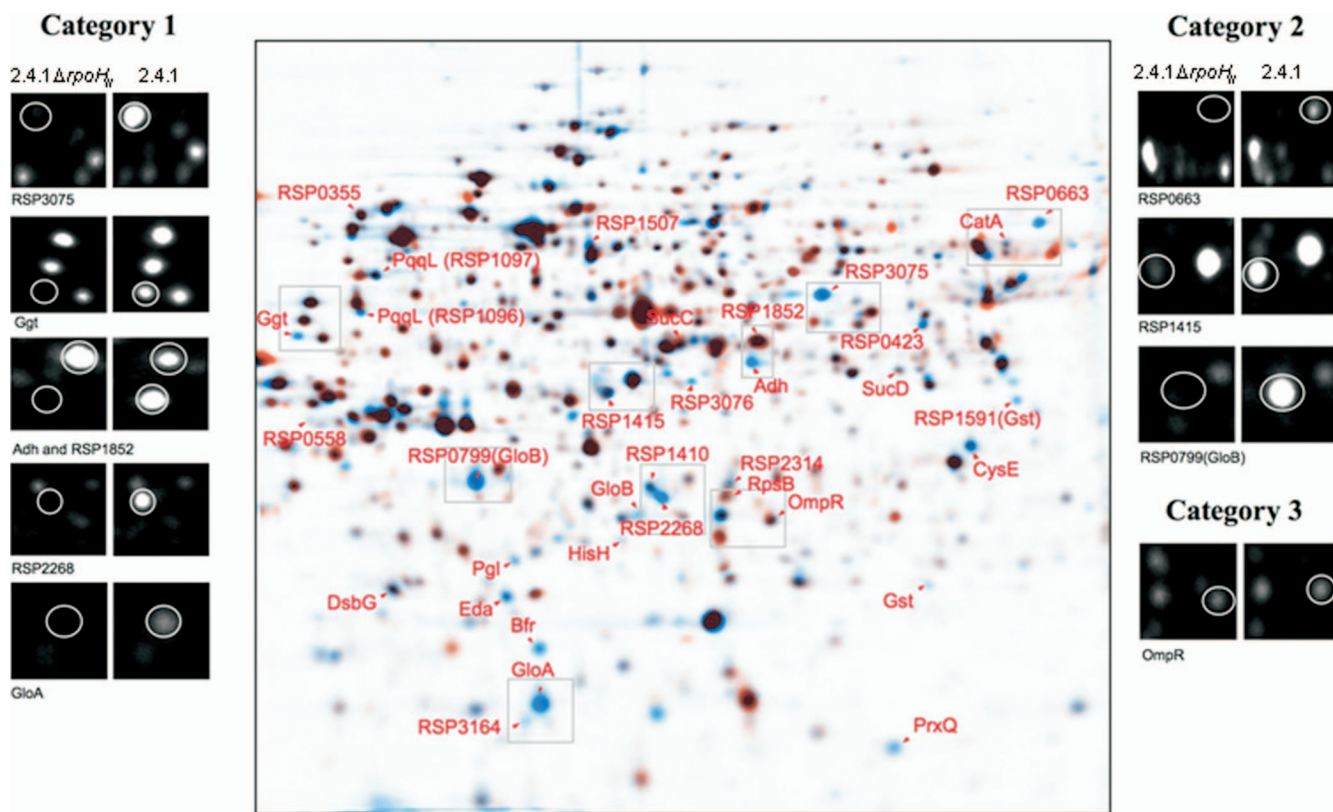


FIG. 2. Superimposed 2D gel electrophoresis images indicating differences in synthesis of soluble proteins between wild-type strain 2.4.1 and strain 2.4.1 Δ rpoH_{II} in response to ¹O₂ exposure. Protein extracts were prepared from cells labeled in vivo with L-[³⁵S]methionine during exponential growth in the presence of methylene blue (0.2 μ M) and high light (800 W m⁻²). Superimposed images were generated by combining the digitalized autoradiograms in Delta 2D software. For both strain 2.4.1 and strain 2.4.1 Δ rpoH_{II}, three gels of independent experiments were fused and overlaid. Blue spots represent proteins from strain 2.4.1, and orange spots represent proteins from strain 2.4.1 Δ rpoH_{II}. Therefore, in superimposed images, blue spots indicate proteins not synthesized in strain 2.4.1 Δ rpoH_{II}. Black spots indicate proteins synthesized similarly in both strains, and orange spots represent proteins exhibiting increased synthesis in strain 2.4.1 Δ rpoH_{II}. Protein numbers labeled with RSP prefixes have hypothetical functions.

synthesis of proteins previously classified as RpoE independent (category 3) was also not altered by the deletion of rpoH_{II}. This is reflected by, e.g., the synthesis of OmpR, which is not altered in strain 2.4.1 Δ rpoH_{II}, as depicted in superimposed 2D gels (Fig. 2; Table 2).

RpoH_{II}-dependent gene expression. In order to determine the role of RpoH_{II} in the expression of strictly RpoH_{II}-dependent genes, we compared mRNA levels of some of the genes in the rpoH_{II} mutant with those in the rpoE-chrR mutant strain TF18 and in the wild-type strain 2.4.1 by real-time RT-PCR (Fig. 3). We included gpx, for which we did not detect a protein in our proteome analysis. Relative mRNA levels of gpx were increased upon exposure to ¹O₂ (16) and blue light, and the gene carries a putative RpoH_{II} target sequence in the upstream region (6). Significantly increased mRNA levels (increased by factors of 3 to 10) were observed for the genes gloA, gloB (RSP0799), adh, ggt, gst (RSP1591), gpx, and RSP3075 in the wild type after 7 min of ¹O₂ exposure (Fig. 3). In contrast, no significant increase in mRNA levels of the above-mentioned genes was observed for the rpoH_{II} mutant. Interestingly, we observed a small increase in mRNA levels in the case of all tested genes, except for gst (RSP1591) in strain TF18. Complementation of the rpoH_{II} mutant with plasmids harboring rpoH_{II}

or rpoH_{II} and RSP0600 restored the relative expression, upon ¹O₂ exposure, of all genes mentioned above. For all of the investigated genes except gst (RSP1591), the relative expression was even higher than that of the wild type, but significantly increased values were observed only for gloB (RSP0799) (Fig. 3).

Because the rpoH_{II} mRNA level was also slightly increased by heat shock (Fig. 1B), mRNA levels of the RpoH_{II}-dependent gene gst (RSP1591) measured under photooxidative stress (Fig. 3) were compared to those in response to heat stress (data not shown). Exposure to heat failed to induce gst (RSP1591) in both the wild-type 2.4.1 and the rpoH_{II} mutant cultures, which supports the specific role of RpoH_{II}-dependent gene expression in response to ¹O₂ exposure.

Identification of putative RpoH_{II}-specific target sequences. To confirm that transcription initiates at the predicted target sequences for some of the RpoH_{II}-dependent genes, we mapped 5' ends of the mRNA by 5' RACE (Fig. 4). For this purpose, we chose gloA, gst (RSP1591), and RSP3075 to investigate genes that strongly depend on RpoH_{II} (Fig. 3).

5' RACE was performed with RNA extracts from the wild type and strain 2.4.1 Δ rpoH_{II} after they were exposed to ¹O₂. To assess the relative changes in expression, cDNA synthesis was performed with gene-specific primers in the same reaction (see

TABLE 2. Comparison of relative protein spot intensities after ¹O₂ exposure

Category and locus tag	Protein	Function	Relative protein spot intensity (%) ± SD ^a		
			2.4.1Δ <i>rpoH</i> _{II}	2.4.1Δ <i>rpoH</i> _{II} (pRK2.4.1 <i>rpoH</i> _{II})	2.4.1Δ <i>rpoH</i> _{II} (pRK2.4.1 <i>rpoH</i> _{II} RSP0600)
Category 1					
RSP0392†	GloA	Predicted lactoylglutathione lyase: glyoxalase I	8 ± 4	116 ± 12	98 ± 11
RSP3272†	Ggt	γ-Glutamyltranspeptidase	0.0 ± 0.0	38 ± 5	29 ± 9
RSP3537†	Adh	Zn-containing alcohol dehydrogenase	0.0 ± 0.0	169 ± 25	148 ± 2
RSP2314†		Predicted oxidoreductase	11 ± 5	63 ± 16	105 ± 7
RSP3075†		Hypothetical protein	9 ± 1	74 ± 0.7	95 ± 2
RSP3076		Hypothetical protein	0.0 ± 1	161 ± 11	152 ± 9
RSP1591	Gst	Predicted glutathione S-transferase	0.0 ± 0.0	93 ± 2	30 ± 10
RSP2645	Eda	KDPG/KHG bifunctional aldolase	39 ± 23	97 ± 6	107 ± 16
RSP2973	PrxQ	Peroxiredoxin	20 ± 9	159 ± 37	91 ± 23
RSP2860	RpsB	30S ribosomal protein S2	192 ± 44	93 ± 9	110 ± 5
RSP0423		Predicted oxidoreductases related to aryl-alcohol dehydrogenases	30 ± 3	111 ± 33	90 ± 40
RSP2268		Predicted metallo β-lactamase	0.0 ± 0.0	113 ± 6	109 ± 11
RSP1852		Hypothetical protein	147 ± 40	108 ± 14	117 ± 1
RSP1464	DsbG	Putative periplasmic thiol-disulfide interchange protein of the DsbA family	82 ± 26	94 ± 24	81 ± 11
Category 2					
RSP3164†		Ferredoxin-like protein	0.0 ± 0.0	74 ± 10	73 ± 21
RSP3342	Bfr	Bacterioferritin	0.0 ± 0.0	147 ± 31	160 ± 15
RSP2481	CysE	Serine acetyltransferase	35 ± 13	62 ± 6	59 ± 18
RSP0799	GloB	Putative hydroxyacylglutathione hydrolase: glyoxalase II	7 ± 4	63 ± 3	58 ± 6
RSP2294	GloB	Putative hydroxyacylglutathione hydrolase: glyoxalase II	2 ± 1	95 ± 1	87 ± 11
RSP1397	Gst	Glutathione S-transferase	15 ± 13	106 ± 49	112 ± 5
RSP2735	Pgl	Inner membrane subunit of ABC sugar transporter	32 ± 15	112 ± 13	104 ± 4
RSP1096	PqqL	Putative zinc peptidase	7 ± 5	76 ± 8	51 ± 0.8
RSP1097	PqqL	Putative zinc peptidase	42 ± 12	47 ± 19	39 ± 0.5
RSP0663		Formate-tetrahydrofolate ligase	6 ± 4	81 ± 54	51 ± 41
RSP1415		Putative polysaccharide deacetylase	0.4 ± 0.0	78 ± 0.3	79 ± 4
RSP1507		NADH-dependent aldehyde dehydrogenase	4 ± 1	35 ± 15	49 ± 9
RSP2779	CatA	Catalase	37 ± 17	129 ± 55	117 ± 49
RSP0355		Putative serine protease involved in heat shock response, exhibiting chaperone function	96 ± 25	75 ± 36	65 ± 3
RSP0558		Possible ribosomal protein L11: predicted methyltransferase	83 ± 43	70 ± 0.4	65 ± 0.6
RSP0967	SucC	Succinyl-coenzyme A synthetase; β subunit	120 ± 53	98 ± 5	109 ± 6
RSP0966	SucD	Succinyl-coenzyme A synthetase; α subunit	127 ± 59	90 ± 9	88 ± 25
RSP1410		Hypothetical protein	49 ± 46	48 ± 3	49 ± 2
RSP2241	HisH	Phosphoribosyl-ATP pyrophosphatase	133 ± 58	100 ± 0.1	100 ± 0.1
Category 3					
RSP0847	OmpR	Two-component transcriptional regulator	117 ± 66	131 ± 32	126 ± 3

^a Protein spot intensities ± standard deviations (SD) were normalized to values observed for the wild-type strain 2.4.1 and are shown for the *rpoH*_{II} mutant strain 2.4.1Δ*rpoH*_{II} and for the complemented mutants harboring the plasmid pRK2.4.1*rpoH*_{II} or pRK2.4.1*rpoH*_{II}RSP0600. Category 1 includes RpoE-dependent proteins, category 2 contains conditionally RpoE-dependent proteins, and category 3 contains RpoE-independent proteins (17). Boldface proteins depend on RpoH_{II}; †, predicted RpoH_{II} target sequence (6). Mean values of three independent experiments are depicted for the *rpoH*_{II} mutant and for the two complemented mutants.

Table S1 in the supplemental material). For all three genes, specific products were PCR amplified from wild-type cultures. Amplification of PCR products obtained from RNA extracts of strain 2.4.1Δ*rpoH*_{II} was much weaker for *gloA* and *gst* (RSP1591) and missing for RSP3075 in comparison to those for the wild type (Fig. 4). A primer specific for *rpoZ*, a gene used for normalization in the real-time RT-PCR approach described previously (39), yielded PCR products from both strains of very similar intensities (Fig. 4). These data strongly support the RpoH_{II}-dependent expression of genes encoding proteins not synthesized upon ¹O₂ exposure in strain 2.4.1Δ*rpoH*_{II}. 5' ends were determined by sequencing the am-

plified DNA fragments after cloning them into the pDrive vector. Putative -10 and -35 RpoH_{II} target sequences were found for all three genes upstream of the 5' end of the respective PCR products (Fig. 4).

Those putative target sequences were similar to those of previously predicted RpoH_{II}-dependent promoters (6) but exhibited differences in the -35 and -10 regions compared to target sequences located upstream of the start codon of several heat shock-triggered genes and recognized by both RpoH_I and RpoH_{II} (19). The -35 region putatively recognized by RpoH_{II} during ¹O₂ exposure contains not only TTG but also TTT or TGG and is flanked by GC-rich positions (Fig. 4). In contrast,

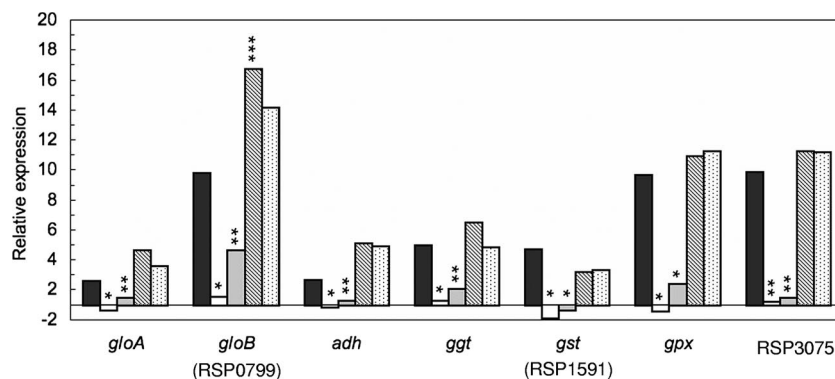


FIG. 3. Selected functional genes depending on RpoH_{II} for expression, which contained a putative RpoH_{II} promoter (6). Relative expression levels of *gloA*, *gloB* (RSP0799), *adh*, *ggt*, *gst* (RSP1591), *gpx*, and RSP3075 in strain 2.4.1 (black bars), 2.4.1Δ*rpoH*_{II} (white bars), *rpoE chrR* mutant strain TF18 (gray bars), strain 2.4.1Δ*rpoH*_{II}(pRK2.4.*rpoH*_{II}) (hatched bars), and 2.4.1Δ*rpoH*_{II}(pRK2.4.*rpoH*_{II}RSP0600) (stippled bars) were determined after 7 min of incubation under photooxidative-stress conditions. Relative expression was determined by real-time RT-PCR. Levels of significance are indicated as follows: *, $P \leq 0.01$; **, $P \leq 0.05$; and ***, $P \leq 0.1$.

the conserved -10 region consists of CTAGCT for all three genes investigated and is separated from the corresponding -35 region by 15 to 16 bp. For comparison, the RpoH_I-specific promoter P1 of *dnaK* is shown, which differs in the -10 region by the exchange of GC with TA (Fig. 4).

Changes in methylglyoxal sensitivity. The proteome analysis approach indicated that several proteins lacking in the *rpoH*_{II} mutant upon ¹O₂ exposure are involved in the degradation of methylglyoxal (Table 2). This toxic nucleophile accumulates

during the disorder of sugar metabolism and oxidative stress and is spontaneously formed within cells (24). Differences between the *rpoH*_{II} mutant and the wild type in sensitivity to methylglyoxal were observed in inhibition zone assays. Inhibition zones observed for the wild type were 3.2 ± 0.1 cm in diameter and were significantly increased to 3.6 ± 0.1 cm for the *rpoH*_{II} mutant. Strains harboring plasmid pRK415 yielded no significantly different inhibition zone values (see Table S2 in the supplemental material). The inhibition zone diameters

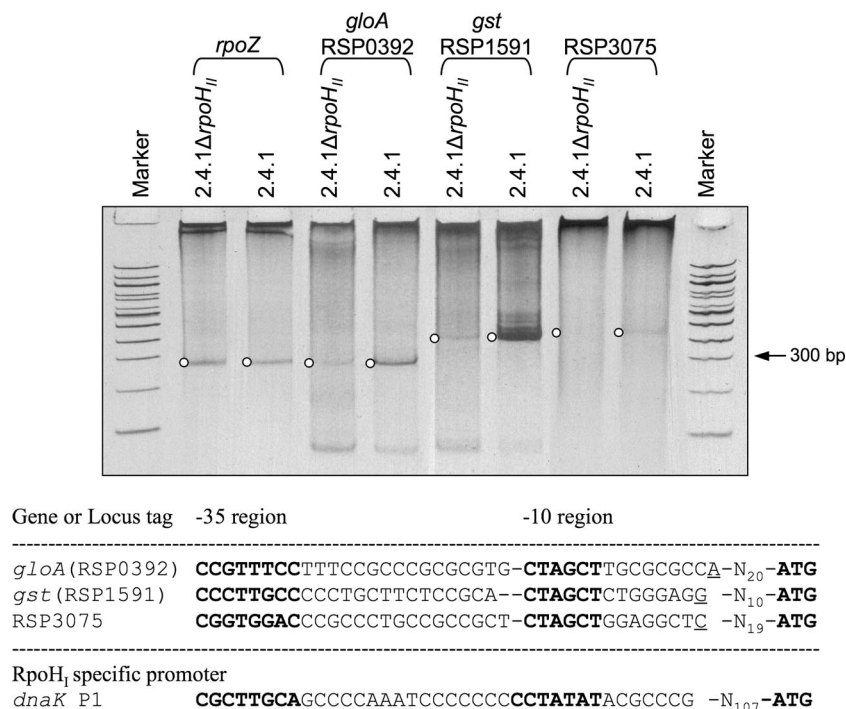


FIG. 4. Separation of 5' RACE products obtained from RNA extracts of wild-type and *rpoH*_{II} mutant cultures after 10 min of photooxidative stress. PCR products obtained after the second PCR (nested) were separated on a 10% polyacrylamide gel and stained with ethidium bromide. Upstream of the 5' ends of the sequences corresponding to the depicted DNA bands, RpoH_{II} target sequences were found and are depicted as aligned sequences below the gel image. DNA marker lanes, 100-bp ladder. In the alignment, transcription start sites are underlined and putative -35 and -10 regions are printed in bold letters. The *dnaK* P1 promoter sequence is shown for comparison and is recognized only by RpoH_I in vitro (19).

yielded by the two complemented *rpoH_{II}* mutants were not significantly different from that of the wild type and measured 3.2 ± 0.1 cm and 3.3 ± 0.1 cm for strains 2.4.1Δ*rpoH_{II}*(pRK2.4.1*rpoH_{II}*) and 2.4.1Δ*rpoH_{II}*(pRK2.4.1*rpoH_{II}*RSP0600, respectively.

DISCUSSION

A functional hierarchy is established by the RpoE-RpoH_{II} sigma factor cascade. Here we demonstrate that RpoH_{II} acts downstream of RpoE in a sigma factor cascade. This is shown by the lack of *rpoH_{II}* expression in the *rpoE*-deficient mutant TF18 under conditions of ¹O₂ exposure (Fig. 1A). *rpoH_{II}* was one of the genes identified among the few that are preceded by an RpoE target sequence in the promoter region (3, 6). Interestingly, a rather small number of genes belonging to the RpoE regulon exhibit a conserved RpoE target sequence (3, 6, 17). Their functions are restricted to DNA repair upon UV damage (photolyase, *phrA*), energy metabolism (cytochrome A, *cycA*; the cyclopropane/cyclopropene fatty acid synthesis RSP1091 protein), gene regulation (*rpoE*, *rpoH_{II}*, and the *tspO*-like RSP1409 protein), transformation of thiols (RSP3184 and *dsbG*), and hypothetical functions (RSP1465 and RSP1852). Therefore, the function of proteins encoded by genes exhibiting a conserved RpoE target sequence are rather different than those of the RpoH_{II} regulon (see below), which provides evidence that a functional hierarchy exists in the response to ¹O₂ exposure.

Our findings show that most of the genes encoding proteins that directly or conditionally depend on RpoE but do not contain a conserved RpoE target sequence are indeed controlled by RpoH_{II} (Fig. 2 to 4; Table 2). Unlike time-oriented sigma factor cascades that regulate, e.g., sporulation, the RpoE-RpoH_{II} cascade controls a functional hierarchy characterized by (i) the rapid induction of RpoH_{II}-dependent protein synthesis upon ¹O₂ exposure (17) and (ii) the function of proteins encoded by genes preceded by RpoE and RpoH_{II} target sequences (Table 2).

Slightly increased mRNA levels of RpoH_{II}-dependent genes were observed upon ¹O₂ exposure in the TF18 strain lacking *rpoE* but not in the *rpoH_{II}* mutant, except for *gloB* (RSP0799) (Fig. 3). This finding indicates that the expression of *rpoH_{II}* may be triggered to a small extent by further molecular factors acting directly on *rpoH_{II}* that have yet to be identified. Although *rpoH_I* transcript levels were increased by ¹O₂ exposure up to 12-fold, the synthesis of typical heat shock proteins such as HslO was increased only 2- to 3-fold compared to that of control conditions (17), and spot intensities of additional heat shock proteins such as GroEL and DnaK were not increased (data not shown). In *Agrobacterium tumefaciens*, the activity of RpoH_I is controlled by the chaperone DnaK, which releases RpoH_I under heat shock conditions (35). It remains to be elucidated if RpoH_I of *R. sphaeroides* is controlled by a mechanism similar to that in *A. tumefaciens*.

Analysis of the *rpoH_{II}* mutant and the complementation with plasmids harboring *rpoH_{II}* or *rpoH_{II}* and RSP0600 show that sensitivity to ¹O₂ and methylglyoxal is caused by the deletion of *rpoH_{II}*. The amounts of proteins synthesized in the complemented strains are not restored to wild-type levels for all proteins, e.g., those encoded by *ggt* and *gst* (Table 2). However,

mRNA levels were restored for both genes and were not different for the two complemented strains.

RpoH_{II} controls a functional response to ¹O₂ exposure. The role of RpoH_{II}-type sigma factors, which are widely distributed among organisms in the *Alphaproteobacteria* (19), is largely unknown with respect to environmental response and function of the controlled genes. The proposed function of genes under RpoH_{II} control may explain the adaptive response to photooxidative stress observed for *R. sphaeroides* (17). ¹O₂ reacts with a variety of cellular molecules, such as proteins, lipids, DNA, and RNA, and forms partially oxidized reaction products, e.g., protein and lipid peroxides, which are toxic to cells (11, 22). Our data show that several genes encoding proteins involved in GSH-dependent and -independent detoxification of ¹O₂ reaction products are controlled by RpoH_{II} (Table 2).

The first group of RpoH_{II}-dependent proteins is involved in the GSH-dependent detoxification of reaction products formed from cellular macromolecules interacting with ¹O₂. Gst (RSP1591 and RSP1397, encoding glutathione *S*-transferase) catalyzes the detoxification of a wide range of molecules that harbor an electrophilic carbon, sulfur, or nitrogen atom and may reduce organic peroxides (22). Mechanistically, conjugates of the electrophilic substrates are formed with GSH. Those conjugates may be further metabolized by Ggt (RSP3272, encoding γ -glutamyltransferase) and aminopeptidases, which subsequently cleave the glutamyl and glycine residues (22). In *R. sphaeroides*, Ggt may represent a periplasmic protein that also ensures regeneration of GSH and redox homeostasis. The aminopeptidase function is unclear but may be carried out by the RpoH_{II}-dependent putative Zn-dependent peptidases encoded by RSP1096 and RSP1097, which harbor a PqqL peptidase motif. PQQ was shown to be a factor for survival against oxidative stress in vitro (33) and in a bacterial system such as that of *E. coli* (27).

The second group of RpoH_{II}-dependent proteins is involved in the degradation of peroxides. The gene product of *prxQ* (RSP2973, encoding a peroxiredoxin) was shown to reduce H₂O₂ and cumene hydroperoxide in vitro, and the deletion of *prxQ* leads to an increased sensitivity to exposure to ¹O₂ and other reactive oxygen species (48). Gpx (RSP2389, encoding GSH peroxidase) was induced by organic hydroperoxides in *Chlamydomonas reinhardtii* (15, 30), and bovine Gpx degraded protein peroxides in vitro (10, 34). Although mRNA levels were increased upon ¹O₂ exposure (16), Gpx was not detected on 2D gels of soluble proteins (Fig. 2) and may therefore be membrane associated in *R. sphaeroides*. Control of *prxQ* and *gpx* expression by RpoH_{II} may be crucial for organic peroxide degradation upon photooxidative stress. Synthesis of CatA (previously called CatE) is decreased in the absence of RpoH_{II} under photooxidative stress. The CatA synthesis induced by RpoH_{II} may reflect (i) the known ¹O₂ sensitivity of catalases or (ii) the need for increased CatA levels to degrade H₂O₂ generated from ¹O₂ reaction products in the cytoplasm.

The third group of proteins controlled by RpoH_{II} is involved in methylglyoxal removal. Methylglyoxal is generated under conditions of carbohydrate metabolism disorder (28) and amino acid breakdown and may accumulate under oxidative stress (24). This strong nucleophile rapidly reacts with macromolecules such as proteins and nucleic acids (9). Therefore, intracellular accumulation needs to be avoided to prevent cellular damage (24). Synthesis of proteins involved in GSH-

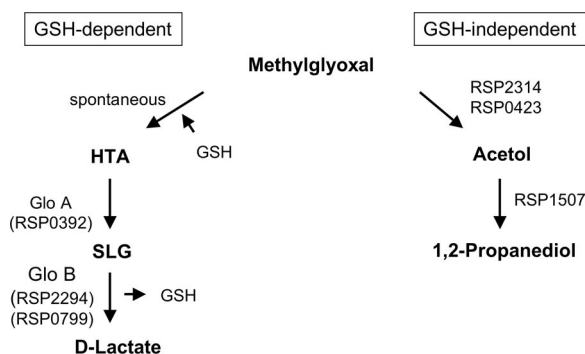


FIG. 5. Role of selected RpoH_{II}-dependent proteins in GSH-dependent and GSH-independent degradation of methylglyoxal. The putative functions of the depicted proteins were inferred from comparisons to homologous proteins based on BLAST search results. HTA, hemithioacetal; SLG, *S*-D-lactoylglutathione. For the functions of the depicted proteins, see Table 2.

dependent and -independent pathways of methylglyoxal degradation are controlled by RpoH_{II} in *R. sphaeroides* (Table 2). GSH-dependent degradation of methylglyoxal is carried out by GloA and GloB (RSP0799 and RSP2294, respectively), which represent the glyoxalase system forming D-lactate as an end product (Fig. 5). Degradation of methylglyoxal independently of GSH is performed by methylglyoxal reductase, aldo-keto reductases, and alcohol dehydrogenases, thereby forming lactaldehyde and 1,2-propanediol (Fig. 5). The oxidoreductases YeaE and YghZ found in *E. coli* have been shown to degrade methylglyoxal in vitro in an NADPH-dependent manner (29). A BLAST search (1) revealed the aldo-keto reductases related to DkgA, YeaE, and YghZ in *R. sphaeroides*: the RpoH_{II}-dependent RSP0423 and RSP2314 oxidoreductases (27 to 46% identity) (Fig. 5). The acetol formed by those oxidoreductases may be degraded by the predicted RSP1507 aldehyde dehydrogenase to 1,2-propanediol, which may then be further metabolized by the RSP3537 product, a predicted alcohol dehydrogenase (not shown in Fig. 5). The control of methylglyoxal-degrading pathways by RpoH_{II} is supported by an increased sensitivity of the *rpoH_{II}* mutant to methylglyoxal. Further studies of the functions of those proteins need to be performed to verify their role in the response to ¹O₂, with respect to methylglyoxal detoxification in *R. sphaeroides*.

Our data show that several of the RpoH_{II}-dependent proteins make use of GSH-dependent and -independent reactions to elicit a cellular response to prevent ¹O₂-caused damages. GSH obviously plays a central role in the defense of *R. sphaeroides* against ¹O₂, which exceeds direct scavenging of ¹O₂ by GSH, as proposed previously (11, 13). Interestingly, 5-methyltetrahydrofolate exhibits the capacity to prevent photosensitizing reactions and may act as a ¹O₂ scavenger (37). A putative formate-tetrahydrofolate ligase (RSP0663) is under the control of RpoH_{II}. Alternatively, 5-methyltetrahydrofolate would need to be regenerated under photooxidative stress, because it is specifically damaged by ¹O₂ and acts as a cofactor of, e.g., photolyases and cryptochromes (42).

Additional proteins controlled by RpoH_{II} that are putatively involved in oxidative-stress responses are Bfr (bacterioferritin) and the RSP3164 (ferredoxin-like) protein, which represent

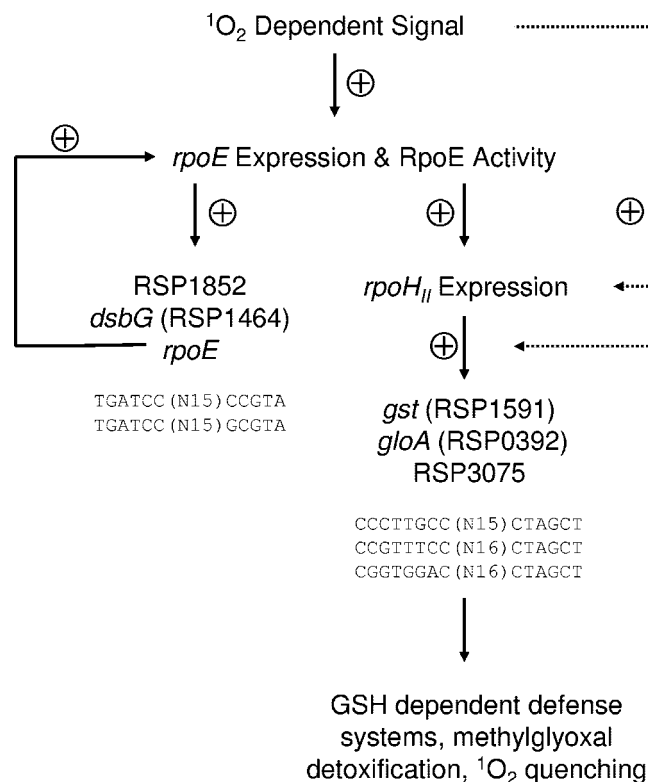


FIG. 6. The current model displays the role of RpoH_{II} in the RpoE-dependent response to ¹O₂ in *R. sphaeroides*. Solid arrows indicate positive effects in the regulatory cascade triggered by ¹O₂. Dotted arrows indicate the functions of hypothetical expression factors involved in the RpoE-RpoH_{II}-dependent gene induction. Representative target sequences for RpoE (3, 17) and the putative target sequences for RpoH_{II} identified in this study (Fig. 4) are depicted.

iron-binding proteins. The control of intracellular iron concentration is pivotal to preventing the formation of hydroxyl radicals by the reaction of iron(II) with peroxides via the Fenton reaction (18). Ferredoxins are involved in cellular redox reactions and may serve as electron donors for the detoxification of reactive oxygen species. In conclusion, the RpoH_{II} regulon defined by the investigation of the *rpoH_{II}* mutant by using a proteome approach revealed a functionally sharply defined and newly assigned role for RpoH_{II} in controlling genes in *R. sphaeroides* in response to ¹O₂ exposure.

Current model of RpoE-RpoH_{II}-controlled gene expression under photooxidative stress. Our current model of gene regulation upon photooxidative stress displays the interaction of the *rpoE* and *rpoH_{II}* gene products (Fig. 6). The recognition of ¹O₂ or ¹O₂-caused damage by *R. sphaeroides* is not clear to date. However, a recently recognized Zn-binding site in the anti-sigma factor ChrR may be involved in the recognition of ¹O₂ (7), which may ultimately lead to the dissociation of the RpoE/ChrR complex. The binding of RpoE to specific target sequences triggers the expression of a small number of genes, including *rpoE* itself and *rpoH_{II}* (3, 6). Thereby, expression and activity of RpoE are increased and *rpoH_{II}*-dependent genes are induced. The functions of genes that harbor an RpoE target sequence did not explain the adaptation of *R. sphaeroides* to photooxidative stress. In contrast, genes that harbor an RpoH_{II}

target sequence encode defense systems responding to oxidative stress. GSH-dependent and GSH-independent mechanisms controlled by RpoH_{II} remove organic peroxides generated by the reaction of ¹O₂ with macromolecules, and accumulation of toxic by-products such as methylglyoxal is prevented. Genes involved in the regeneration or production of ¹O₂-specific scavengers or metabolites that are especially prone to ¹O₂ reaction, such as GSH (13) and 5-methyltetrahydrofolate (37), are under the control of RpoH_{II}.

RpoH_{II}-like sigma factors have so far been recognized only in the *Alphaproteobacteria* (19) and may control rather specific functions. In *Brucella melitensis*, the gene BMEI0280 (*rpoH1*) represents the *R. sphaeroides* *rpoH_{II}* homolog. This gene is not involved in the response to heat stress but may be important for the chronicity of *B. melitensis* infection (12). In *Sinorhizobium meliloti*, *rpoH_{II}* is induced during stationary phase, and an *rpoH_{II}* mutant showed that the resulting strain is not heat sensitive (38). Because several *Alphaproteobacteria* have two or even three *rpoH* homologs, it is reasonable to assume that one of the two heat shock sigma factors was recruited for functions other than defense against heat shock. Our regulon analysis represents the first in vivo study of an RpoH_{II}-type sigma factor by using an *rpoH_{II}* mutant background and indicates that it is crucial for the regulation of defense systems against photooxidative stress in *R. sphaeroides*. Furthermore, our data indicate that the ¹O₂-dependent signal triggers the recognition of RpoH_{II}-specific target sequences and that the RpoH_{II}-dependent response is to some extent independent from RpoE.

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REFERENCES

- Altschul, S. F., W. Gish, W. Miller, E. W. Myers, and D. J. Lipman. 1990. Basic local alignment search tool. *J. Mol. Biol.* **215**:403–410.
- Anthony, J. R., J. D. Newman, and T. J. Donohue. 2004. Interactions between the *Rhodobacter sphaeroides* ECF sigma factor, σ^E , and its anti-sigma factor, ChrR. *J. Mol. Biol.* **341**:345–360.
- Anthony, J. R., K. L. Warczak, and T. J. Donohue. 2005. A transcriptional response to singlet oxygen, a toxic byproduct of photosynthesis. *Proc. Natl. Acad. Sci. USA* **102**:6502–6507.
- Arellano, J. B., Y. A. Yousef, T. B. Melø, S. B. B. Mohamad, R. J. Cogdell, and K. R. Naqvi. 2007. Formation and geminate quenching of singlet oxygen in purple bacterial reaction center. *J. Photochem. Photobiol. B* **87**:105–112.
- Braatsch, S., M. Gomelsky, S. Kuphal, and G. Klug. 2002. A single flavoprotein, AppA, integrates both redox and light signals in *Rhodobacter sphaeroides*. *Mol. Microbiol.* **45**:827–836.
- Braatsch, S., O. V. Moskvina, G. Klug, and M. Gomelsky. 2004. Responses of the *Rhodobacter sphaeroides* transcriptome to blue light under semiaerobic conditions. *J. Bacteriol.* **186**:7726–7735.
- Campbell, E. A., R. Greenwell, J. R. Anthony, S. Wang, L. Lim, K. Das, H. J. Sofia, T. J. Donohue, and S. A. Darst. 2007. A conserved structural module regulates transcriptional responses to diverse stress signals in bacteria. *Mol. Cell* **27**:793–805.
- Cogdell, R. J., T. D. Howard, R. Bittl, E. Schlodder, I. Geisenheimer, and W. Lubitz. 2000. How carotenoids protect bacterial photosynthesis. *Philos. Trans. R. Soc. Lond. B* **355**:1345–1349.
- Cooper, R. A. 1984. Metabolism of methylglyoxal in microorganisms. *Annu. Rev. Microbiol.* **38**:49–68.
- Davies, M. J. 2004. Reactive species formed on proteins exposed to singlet oxygen. *Photochem. Photobiol. Sci.* **3**:17–25.
- Davies, M. J. 2003. Singlet oxygen-mediated damage to proteins and its consequences. *Biochem. Biophys. Res. Commun.* **305**:761–770.
- Delory, M., R. Hallez, J.-J. Letesson, and X. De Bolle. 2006. An RpoH-like heat shock sigma factor is involved in stress response and virulence in *Brucella melitensis* 16M. *J. Bacteriol.* **188**:7707–7710.
- Di Mascio, P., T. P. A. Devasagayam, S. Kaiser, and H. Sies. 1990. Carotenoids, tocopherols and thiols as biological singlet molecular oxygen quenchers. *Biochem. Soc. Trans.* **18**:1054–1056.
- Drews, G. 1983. *Mikrobiologisches praktikum*. Springer Verlag, Heidelberg, Germany.
- Fischer, B. B., R. I. L. Eggen, A. Trebst, and A. Krieger-Liszskay. 2006. The glutathione peroxidase homologous gene GpxH in *Chlamydomonas reinhardtii* is upregulated by singlet oxygen produced in photosystem II. *Planta* **223**:583–590.
- Glaeser, J., and G. Klug. 2005. Photo-oxidative stress in *Rhodobacter sphaeroides*: protective role of carotenoids and expression of selected genes. *Microbiology* **151**:1927–1938.
- Glaeser, J., M. Zobawa, F. Lottspeich, and G. Klug. 2007. Protein synthesis patterns reveal a complex regulatory response to singlet oxygen in *Rhodobacter*. *J. Proteome Res.* **6**:2460–2471.
- Goldstein, S., D. Meyerstein, and G. Czapski. 1993. The Fenton reagents. *Free Radic. Biol. Med.* **15**:435–445.
- Green, H. A., and T. J. Donohue. 2006. Activity of *Rhodobacter sphaeroides* RpoH_{II}, a second member of the heat shock sigma factor family. *J. Bacteriol.* **188**:5712–5721.
- Gregor, J., and G. Klug. 1999. Regulation of bacterial photosynthesis genes by oxygen and light. *FEMS Microbiol. Lett.* **179**:1–9.
- Happ, H. N., S. Braatsch, V. Broschek, L. Osterloh, and G. Klug. 2005. Light-dependent regulation of photosynthesis genes in *Rhodobacter sphaeroides* 2.4.1 is coordinately controlled by photosynthetic electron transport via the PrrBA two-component system and the photoreceptor AppA. *Mol. Microbiol.* **58**:903–914.
- Hayes, J. D., J. U. Flanagan, and I. R. Jowsey. 2005. Glutathione transferases. *Annu. Rev. Pharmacol. Toxicol.* **45**:51–88.
- Hübner, P., J. C. Willison, P. M. Vignais, and T. A. Bickle. 1991. Expression of regulatory *nif* genes in *Rhodobacter capsulatus*. *J. Bacteriol.* **173**:2993–2999.
- Kalapos, M. P. 1999. Methylglyoxal in living organisms: chemistry, biochemistry, toxicology and biological implications. *Toxicol. Lett.* **110**:145–175.
- Karls, R. K., J. Brooks, P. Rossmessl, J. Luedke, and T. J. Donohue. 1998. Metabolic roles of a *Rhodobacter sphaeroides* member of the sigma 32 family. *J. Bacteriol.* **180**:10–19.
- Keen, N. T., S. Tamaki, D. Kobayashi, and D. Trollinger. 1988. Improved broad-host-range plasmids for DNA cloning in Gram-negative bacteria. *Gene* **70**:191–197.
- Khairnar, N. P., H. S. Misra, and S. K. Apte. 2003. Pyrroloquinoline-quinone synthesized in *Escherichia coli* by pyrroloquinoline-quinone synthase of *Deinococcus radiodurans* plays a role beyond mineral phosphate solubilization. *Biochem. Biophys. Res. Commun.* **312**:303–308.
- Kim, I., E. Kim, S. Yoo, D. Shin, B. Min, J. Song, and C. Park. 2004. Ribose utilization with an excess of mutarotase causes cell death due to accumulation of methylglyoxal. *J. Bacteriol.* **186**:7229–7235.
- Ko, J., I. Kim, S. Yoo, B. Min, K. Kim, and C. Park. 2005. Conversion of methylglyoxal to acetol by *Escherichia coli* aldo-keto reductases. *J. Bacteriol.* **187**:5782–5789.
- Leisinger, U., K. Rufenacht, B. Fischer, M. Pesaro, A. Spengler, A. J. Zehnder, and R. I. Eggen. 2001. The glutathione peroxidase homologous gene from *Chlamydomonas reinhardtii* is transcriptionally up-regulated by singlet oxygen. *Plant Mol. Biol.* **46**:395–408.
- Mackenzie, C., J. M. Eraso, M. Choudhary, J. H. Roh, X. Zeng, P. Bruscella, A. Puskas, and S. Kaplan. 2007. Postgenomic adventures with *Rhodobacter sphaeroides*. *Annu. Rev. Microbiol.* **61**:283–307.
- Masuda, S., and C. E. Bauer. 2002. AppA is a blue light photoreceptor that antirepresses photosynthesis gene expression in *Rhodobacter sphaeroides*. *Cell* **110**:613–623.
- Misra, H. S., N. P. Khairnar, A. Barik, K. Indira Priyadarshini, H. Mohan, and S. K. Apte. 2004. Pyrroloquinoline-quinone: a reactive oxygen species scavenger in bacteria. *FEBS Lett.* **578**:26–30.
- Morgan, P. E., R. T. Dean, and M. J. Davies. 2004. Protective mechanisms against peptide and protein peroxides generated by singlet oxygen. *Free Radic. Biol. Med.* **36**:484–496.
- Nakahigashi, K., H. Yanagi, and T. Yura. 2001. DnaK chaperone-mediated control of activity of a σ^{32} homolog (RpoH) plays a major role in the heat shock response of *Agrobacterium tumefaciens*. *J. Bacteriol.* **183**:5302–5310.
- Newman, J. D., M. J. Falkowski, B. A. Schilke, L. C. Anthony, and T. J. Donohue. 1999. The *Rhodobacter sphaeroides* ECF sigma factor σ^E and the target promoters *cycA* P3 and *rpoE* P1. *J. Mol. Biol.* **294**:307–320.
- Offer, T., B. N. Ames, S. W. Bailey, E. A. Sabens, M. Nozawa, and J. E. Ayling. 2007. 5-Methyltetrahydrofolate inhibits photosensitization reactions and strand breaks in DNA. *FASEB J.* **21**:2101–2107.
- Oke, V., B. G. Rushing, E. J. Fisher, M. Moghadam-Tabrizi, and S. R. Long. 2001. Identification of the heat-shock sigma factor RpoH and a second RpoH-like protein in *Sinorhizobium meliloti*. *Microbiology* **147**:2399–2408.
- Pappas, C. T., J. Sram, O. V. Moskvina, P. S. Ivanov, R. C. Mackenzie, M. Choudhary, M. L. Land, F. W. Larimer, S. Kaplan, and M. Gomelsky. 2004. Construction and validation of the *Rhodobacter sphaeroides* 2.4.1 DNA microarray: transcriptome flexibility at diverse growth modes. *J. Bacteriol.* **186**:4748–4758.
- Pfaffl, M. W. 2001. A new mathematical model for relative quantification in real-time RT-PCR. *Nucleic Acids Res.* **29**:e45.
- Prentki, P., A. Binda, and A. Epstein. 1991. Plasmid vectors for selecting

- IS1-promoted deletions in cloned DNA: sequence analysis of the omega interposon. *Gene* **103**:17–23.
42. **Sancar, A.** 2003. Structure and function of DNA photolyase and cryptochrome blue-light photoreceptors. *Chem. Rev.* **103**:2203–2238.
 43. **Schilke, B. A., and T. J. Donohue.** 1995. ChrR positively regulates transcription of the *Rhodobacter sphaeroides* cytochrome *c*₂ gene. *J. Bacteriol.* **177**: 1929–1937.
 44. **Shimada, H., K. Iba, and K.-i. Takamiya.** 1992. Blue-light irradiation reduces the expression of *puf* and *puc* operons of *Rhodobacter sphaeroides* under semi-aerobic conditions. *Plant Cell Physiol.* **33**:471–475.
 45. **Simon, R., M. O'Connell, M. Labes, and A. Puhler.** 1986. Plasmid vectors for the genetic analysis and manipulation of *Rhizobia* and other gram-negative bacteria. *Methods Enzymol.* **118**:640–659.
 46. **Uchoa, A., P. Knox, R. Turchielle, N. Seifullina, and M. Baptista.** 2008. Singlet oxygen generation in the reaction centers of *Rhodobacter sphaeroides*. *Eur. Biophys. J.* **37**:843–850.
 47. **van Niel, C. B.** 1944. The culture, general physiology, morphology, and classification of the non-sulfur purple and brown bacteria. *Bacteriol. Rev.* **8**:1–118.
 48. **Wakita, M., S. Masuda, K. Motohashi, T. Hisabori, H. Ohta, and K.-I. Takamiya.** 2007. The significance of type II and PrxQ peroxiredoxins for antioxidative stress response in the purple bacterium *Rhodobacter sphaeroides*. *J. Biol. Chem.* **282**:27792–27801.
 49. **Zeilstra-Ryalls, J., M. Gomelsky, J. M. Eraso, A. Yeliseev, J. O'Gara, and S. Kaplan.** 1998. Control of photosystem formation in *Rhodobacter sphaeroides*. *J. Bacteriol.* **180**:2801–2809.
 50. **Zeilstra-Ryalls, J. H., and S. Kaplan.** 2004. Oxygen intervention in the regulation of gene expression: the photosynthetic bacterial paradigm. *Cell. Mol. Life Sci.* **61**:417–436.