MicroReview

Shedding light on a Group IV (ECF11) alternative σ factor

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Summary

This year marks the 50th anniversary of the discovery of σ70 as a protein factor that was needed for bacterial RNA polymerase to accurately transcribe a promoter in vitro. It was 25 years later that the Group IV alternative σs were described as a distinct family of proteins related to σ70. In the intervening time, there has been an ever-growing list of Group IV σs, numbers of genes they transcribe, insight into the diverse suite of processes they control, and appreciation for their impact on bacterial lifestyles. This work summarizes knowledge of the Rhodobacter sphaeroides σE-ChrR pair, a member of the ECF11 subfamily of Group IV alternative σs, in protecting cells from the reactive oxygen species, singlet oxygen. It describes lessons learned from analyzing ChrR, a zinc-dependent anti-σ factor, that are generally applicable to Group IV σs and relevant to the response to single oxygen. This MicroReview also illustrates insights into stress responses in this and other bacteria that have been acquired by analyzing or modeling the activity of the σE-ChrR across the bacterial phylogeny.

Introduction

The study of Group IV or extracytoplasmic function (ECF) σ factor function has provided many new insights into the cell biology, stress responses and signaling pathways across the bacterial phylogeny (Staron et al., 2009; Feklístov et al., 2014), and provided strategies to allow for targeted control of gene expression in native and heterologous hosts (Rhodius et al., 2013; Pinto et al., 2019). This contribution will review what is known about the ECF11 sub-family of Group IV σs (Staron et al., 2009). It will focus on the founding member of the ECF11 sub-family, the Rhodobacter sphaeroides σE protein, its cognate cytoplasmic anti-σ ChrR (Newman et al., 1999; 2001), and their role in a stress response to singlet oxygen, a reactive oxygen species (ROS) encountered by a variety of cells (Anthony et al., 2005; Ziegelhoffer and Donohue, 2009). It will summarize lessons learned about Group IV σ factor function by studying this system, and highlight unanswered questions about this response.

Singlet oxygen (1O2) is a ROS

Prior to the introduction of molecular oxygen (O2), organisms had a limited metabolic and regulatory repertoire. However, when photosynthetic cells acquired the ability to produce O2, they altered the Earth’s atmosphere and influenced the forms of life that inhabited the planet (Raymond et al., 2003; Kerr, 2005). In particular, the accumulation of atmospheric O2 allowed evolution of pathways like aerobic respiration that couple the four-electron reduction of O2 to formation of a proton gradient (Kerr, 2005). One advantage to aerobic respiration is the large amount of energy that is conserved as O2 is reduced to water (Gennis, 1986; Brzezinski and Gennis, 2008; Borisov et al., 2011; Bueno et al., 2012; Soo et al., 2019). However, there are other, potentially deleterious, consequences to life in the presence of O2. One trade off to accumulation of atmospheric O2, or its use as a terminal electron acceptor, is formation of different ROS (Rosner and Storz, 1997; Schulz et al., 2000; Mittler et al., 2004; Frick et al., 2015; Taverne et al., 2018).

When one electron is sequentially transferred to O2 (Fig. 1), Type I ROS (superoxide, hydrogen peroxide, or hydroxyl radicals) are formed (Rosner and Storz, 1997). Each Type I ROS can damage biomolecules, kill cells or trigger the onset of debilitating diseases (Schulz et al., 2000; Taverne et al., 2018). Consequently, considerable effort has been invested into determining the stress response(s) to these ROS (Rosner and Storz, 1997;
In contrast, less is known about how cells respond to the Type II ROS $1O_2$ (Ragàs et al., 2013; Dogra et al., 2018). $1O_2$ is formed when energy transfer from an excited, triplet state donor, to $O_2$ alters the distribution of electrons in its outer orbital (Fig. 1). Enzymes which detoxify superoxide or $H_2O_2$ are ineffective against $1O_2$ due to differences in outer orbital electron organization between these Type 1 and Type II ROS (Ziegelhoffer and Donohue, 2009). Indeed, there are no known enzyme-catalyzed systems for $1O_2$ detoxification (Davies, 2004).

The outer orbital electron organization of $1O_2$ makes it a strong oxidant (~900mV energy difference between $1O_2$ and $O_2$). $1O_2$ is known or predicted to peroxidize and eventually cleave unsaturated bonds in olefins, oxidize amino acid side chains or nucleic acid bases and cleave peptide or phosphodiester bonds (Nymann and Hynninen, 2004; Godley et al., 2005). Thus, it is not surprising that $1O_2$ can also inhibit growth or kill cells (Anthony et al., 2005; Ziegelhoffer and Donohue, 2009; Lemke et al., 2014).

**Biological formation of $1O_2$**

Major cellular sources of $1O_2$ include the enzymes NADH oxidase, myloperoxidase or chloroperoxidase (Kochevar, 2004; Davies, 2004; Godley et al., 2005). Light energy capture by photosynthetic pigments is another significant source of $1O_2$ (Fig. 1). In the light reactions of photosynthesis, photons excite chlorophyll pigments to a high-energy state (Cogdell, 2000; Frank and Brudvig, 2004; Kochevar, 2004; Triantaphylides and Havaux, 2009). Normally, these excited (triplet state) pigments transfer energy to a reaction center (in bacteria) or photosystem (in cyanobacteria, algae and plants) resulting in light-driven oxidation of this membrane enzyme. However, at a significant frequency, energy transfer from light-excited photopigments to $O_2$ generates $1O_2$ (Cogdell, 2000; Frank and Brudvig, 2004; Kochevar, 2004; Uchoa et al., 2008; Triantaphylides and Havaux, 2009).

$1O_2$ has a high reactivity, so it is predicted to have a short cellular half-life (~100 ns), not travel far its site of synthesis, and produce localized damage (Kochevar, 2004).
phototrophs, $^1\text{O}_2$ formation initiates a process called photodioxidative stress (Triantaphylides and Havaux, 2009; Ziegelhoffer and Donohue, 2009) that can inactivate integral photosynthetic membrane enzymes (Clogg, 2000; Fryer et al., 2002; Frank and Brudvig, 2004; Kochevar, 2004; Szabó et al., 2005), peroxidize or cleave nearby olefins (carotenoids or unsaturated fatty acids), destroy bilayer integrity and function (Girotti and Kriska, 2004; Ramel et al., 2012; Lemke et al., 2014), signal changes in nuclear gene expression from the organelle (chloroplasts) where it is generated in eukaryotic phototrophs or trigger apoptosis (Danon et al., 2005; Foyer and Noctor, 2005).

$^1\text{O}_2$ promotes a bacterial transcriptional response

We uncovered a role for an ECF11 Group IV $\sigma$ factor in a $^1\text{O}_2$ stress response by studying the photosynthetic bacterium *Rb. sphaeroides* (Anthony et al., 2005; Dufour et al., 2008; Ziegelhoffer and Donohue, 2009; Nam et al., 2013). In the laboratory, photosynthetic growth of *Rb. sphaeroides* is often achieved by incubating cells anaerobically in the light (Tavano and Donohue, 2006; Donohue and Kiley, 2011), so $^1\text{O}_2$ is not formed under these conditions. However, we discovered that *Rb. sphaeroides* mounts a transcriptional response to $^1\text{O}_2$ either when pigmented cells are exposed to light and $\text{O}_2$ or when non-pigmented cells are exposed to the photosensitizer methylene blue, light and $\text{O}_2$ (Anthony et al., 2005), two conditions that are well known to produce this ROS (Fig. 2). The master regulator of this transcriptional response to $^1\text{O}_2$ in *Rb. sphaeroides* is the Group IV $\sigma$ factor, $\sigma^E$ (Anthony et al., 2005; Campbell et al., 2007; Greenwell et al., 2011).

In many cells, the ability of carotenoids to quench $^1\text{O}_2$ is generally accepted to be a major route of detoxification of this ROS (Clogg, 2000; Frank and Brudvig, 2004; Kochevar, 2004). However, quenching by carotenoids must not provide complete protection against $^1\text{O}_2$ since this ROS can inactivate proteins, and oxidize membrane fatty acids and other olefins (Rinalducci et al., 2004; Kochevar, 2004; Nishiyama et al., 2004; Estevam et al., 2004; Ramel et al., 2012; Lemke et al., 2014). In addition, $^1\text{O}_2$ formation kills *Rb. sphaeroides* $\Delta\sigma^E$ cells, demonstrating the essential role of the $\sigma^E$-dependent transcriptional response to this ROS (Anthony et al., 2005).

The *Rb. sphaeroides* $\sigma^E$-dependent pathway is not activated by superoxide, $\text{H}_2\text{O}_2$ or hydroxyl radicals (Anthony et al., 2005; Greenwell et al., 2011). However, we and others found that the organohydroperoxide tert-butylhydroperoxide ($\text{t-BOOH}$) increases *Rb. sphaeroides* $\sigma^E$ activity (Lourenco and Gomes, 2009; Greenwell et al., 2011; Nam et al., 2013). Based on studies with model compounds, $^1\text{O}_2$ oxidization of biomolecules could form organohydroperoxides in the membrane (Stief, 2003; Davies, 2004; Kochevar, 2004; Watabe et al., 2007; Triantaphylides and Havaux, 2009). Thus, activation of the *Rb. sphaeroides*...
ChrR is a negative regulator of \( \sigma^E \) activity

Group IV \( \sigma \)s typically bind to a cognate anti-\( \sigma \) factor that is co-transcribed with the \( \sigma \) factor structural gene (Staron et al., 2009). \( \text{Rbs. sphaeroides} \) \( \sigma^E \) follows this paradigm since ChrR, its cognate anti-\( \sigma \), which forms a complex with \( \sigma^E \), prevents it from transcribing target genes (Newman, 2001; Newman et al., 2001; Anthony et al., 2004; Campbell et al., 2007), is encoded by the \( \text{rpoEchrR} \) operon (Newman et al., 1999). However, unlike many other Group IV anti-\( \sigma \) factors, which are integral membrane proteins (Staron et al., 2009), \( \text{Rbs. sphaeroides} \) ChrR is a cytoplasmic protein (Newman et al., 2001; Anthony et al., 2003; 2004; Campbell et al., 2007). Indeed, \( \text{Rbs. sphaeroides} \) ChrR was the founding member of the ECF11 family of Group IV anti-\( \sigma \)s (Staron et al., 2009).

How ChrR blocks \( \sigma^E \) activity

Key insights into the ECF11 family came from solving the three-dimensional structure of the \( \text{Rbs. sphaeroides} \) \( \sigma^E \)-ChrR complex (in collaboration with the Darst lab) (Campbell et al., 2007). In the \( \sigma^E \)-ChrR complex, the \( \text{Rbs. sphaeroides} \) \( \sigma^E \) fold is similar to that of other \( \sigma \)s, including \( \text{Escherichia coli} \) \( \sigma^70 \) and \( \sigma^E \) (Campbell et al., 2002; 2003), consisting of two \( \alpha \)-helical domains (\( \sigma \) regions 2 and 4) connected by a short domain 2-4 linker (Fig. 3).

In this structure, ChrR contains two major structural elements that are connected by a flexible linker. The ChrR N-terminal domain makes extensive contacts with those \( \sigma^E \) regions predicted to bind RNA polymerase and promoter DNA (Fig. 3), so this part of ChrR was called the anti-sigma domain (ChrR-ASD) to denote how it could block \( \sigma \) factor function.

Zinc binding to the ChrR-ASD is needed to inhibit \( \sigma^E \) activity

\( \text{Rbs. sphaeroides} \) ChrR and \( \text{Streptomyces coelicolor} \) RsrA were founding members of the ZAS anti-\( \sigma \) proteins, a family of zinc-dependent anti-sigma factors that each bind a zinc metal in their N-terminal ASDs (Paget et al., 2001a; Paget and Buttner, 2003; Bae et al., 2004; Zdanowski et al., 2006). It was known that zinc binding was required for ChrR to inhibit \( \sigma^E \) activity (Newman et al., 2001). In the \( \sigma^E \)-ChrR complex, zinc is tetrahedrally coordinated to amino acid side chains in the ChrR-ASD (Fig. 3) that were predicted to be involved in zinc binding based on studying mutant ChrR proteins containing single alanine substitutions at these positions (Newman et al., 2001).

Conservation of structure and function among different Group IV anti-\( \sigma \)s

The structure of the \( \sigma^E \)-ChrR complex contributed to developing an early model for how many anti-sigma factors could inhibit function of Group IV \( \sigma \)s (Campbell et al., 2002). This model was based on the unexpected finding that the ChrR-ASD and the N-terminal domain of the \( \text{E. coli} \) anti-sigma factor RseA (RseA-ASD) both contain similar \( \alpha \)-helical bundles despite the lack of significant primary amino acid sequence similarity between these proteins (Campbell et al., 2007). Each ASD contains one structurally conserved helix, helix IV, which interacts with region 2.1 of its cognate Group IV \( \sigma \) in their respective complexes (Fig. 4, right). Based on this, we proposed that many other ECF anti-sigma factors will use a region...
structurally related to ASD helix IV to bind region 2.1 of their cognate ECF $\sigma$-s and block RNA polymerase binding (Campbell et al., 2007).

We also made the unexpected finding that the region of structural similarity between the ChrR- and RseA-ASDs includes the ChrR zinc binding site (Fig. 4, left). The ability of both anti-$\sigma$-s to adopt a similar fold shows that either zinc–protein interactions (in ChrR) or protein–protein interactions (in RseA) can stabilize the ASD helical bundle (Campbell et al., 2007).

The other structurally conserved helices in the ASD of each anti-sigma factor (helices I-III) interact with different regions of their cognate sigma factors, providing a way for each anti-$\sigma$ to recognize specific partner proteins (Campbell et al., 2007). When these observations were combined with comparative genomics, it predicted that the ASDs of ChrR homologs, as well as many other Group IV anti-$\sigma$-factors that have lower degrees of amino acid identity and thus fall into other ECF subfamilies (Staron et al., 2009), could adopt a similar fold when bound to their cognate $\sigma$ factor (Campbell et al., 2007). Subsequent structural analysis of additional complexes has revealed that, while the ASD conformation has been highly conserved among anti-sigma factors, the mechanism of inhibition of $\sigma$ factor activity is unique for each cognate pair examined (Sineva et al., 2017).

**The C-terminal domain of ChrR is needed to release $\sigma^E$ in the presence of $^1O_2$**

The ChrR C-terminal domain binds a 2$^{nd}$ zinc atom within a structural element that adopts an overall fold similar to proteins in the cupin superfamily (Khuri et al., 2001), so we called this the ChrR cupin-like domain (ChrR-CLD, Fig. 3). In structurally characterized proteins that contain a CLD, it can have enzyme activity (isomerases) or bind a ligand (Khuri et al., 2001). The overall CLD fold and the residues known to bind zinc in the $\sigma^E$-ChrR structure are predicted to exist in many other ChrR homologs (Campbell et al., 2007) and a variety of ZAS that are members of other ECF subfamilies (Rajasekar et al., 2016).

The ChrR-CLD had little contact with $\sigma^E$, leading us to propose that this region was unnecessary for formation of the $\sigma^E$-ChrR complex. As predicted, a truncated ChrR protein lacking the CLD (ChrR85) inhibited $\sigma^E$ activity, but cells containing ChrR85 did not mount a transcriptional response to $^1O_2$ (Campbell et al., 2007), predicting that the ChrR-CLD is needed to activate the response. When we analyzed function of ChrR variants containing amino acid substitutions in the CLD zinc ligands, we identified side chains that are (147Glu and 177His) and are not (187Cys and 189Cys) needed for $^1O_2$ or the organoperoxide like t-BOOH (see below) to increase $\sigma^E$ activity (Greenwell et al., 2011).

$^1O_2$ stimulates ChrR turnover

Some Group IV anti-$\sigma$-factors, including others that bind zinc, are reversibly modified by an inducing signal (Paget and Buttner, 2003; Antelmann and Helmann, 2011). However, there is no known mechanism for reversible protein modification by $^1O_2$ (Davies, 2004). Instead, we found that $^1O_2$ promotes ChrR proteolysis (Nam et al., 2013), releasing $\sigma^E$ so it can bind RNA polymerase (Anthony et al., 2004) and directly activate transcription (Ziegelhoffer and Donohue, 2009; Dufour et al., 2010; 2012).

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**Fig. 4.** Structural similarity between the *Rb. sphaeroides* ChrR- & *E. coli* RseA-ASD. The left panel shows the structural similarity between helices I-III and the displacement of helix IV of the ASD of *Rb. sphaeroides* ChrR (red) and *E. coli* RseA (white). The blue sphere is the Zn$^{2+}$ atom in the ChrR-ASD. The right panel shows that, despite this displacement of helix IV in the ASD of ChrR (red) and RseA (white), it interacts with a structurally conserved part of region 2 in the cognate Group IV sigma factors (region 2 of the *Rb. sphaeroides* and *E. coli* $\sigma^E$ proteins are both shown in green). Figures modified from (Campbell et al., 2007).
There is precedent for regulated proteolysis of a Group IV anti-σ factor during a stress response, since cleavage of *E. coli* RseA is initiated by a protease cascade (including DegS and YaeL) that responds to envelope stress (Alba et al., 2002). After DegS and YaeL cleavage of RseA in its membrane spanning region, housekeeping proteases complete degradation of this anti-σ, releasing the *E. coli* σ^F^ so it can activate transcription (Chabert et al., 2007). Homologs of extra-cytoplasmic proteases that cleave RseA have been reported to promote ChrR turnover in vivo (Nuss et al., 2013). However, direct proteolytic cleavage of ChrR has yet to be reported and it is not clear how membrane- or periplasmic-localized proteases promote direct or indirect degradation of this cytoplasmic anti-σ. Indeed, it is possible that 1O_2^\* initiates a conformational change in ChrR that can make it protease-susceptible since this ROS can remove zinc from a synthetic peptide that contains the metal ligands and mimics the fold found in the ChrR-ASD (Chabert et al., 2019). There is precedent for zinc release playing such a regulatory role in the bacterial chaperone Hsp33 and the ZAS anti-σ factor, RsrA, that each respond to oxidative stress signals (Jakob et al., 2000; Kim et al., 2001; Raman et al., 2001; Paget et al., 2001a; 2001b; Paget and Buttner, 2003; Zdanowski et al., 2006; Rajasekar et al., 2016), so the precise role(s) of the ChrR-ASD zinc in the response to 1O_2^* is unknown.

Other proteins have been shown to be needed for ChrR turnover in the presence of 1O_2^* (Nam et al., 2013; Nuss et al., 2013). However, these proteins lack significant amino acid sequence similarity to proteases but some catalyze synthesis of an unusual furan-containing fatty acid (Lemke et al., 2014). 1O_2^* formation promotes destruction of furan-containing fatty acids, so it has been proposed that peroxidation of membrane bound olefins can act as a second messenger to stimulate the activity of one or more proteases that initiates degradation of ChrR in response to this ROS (Lemke et al., 2014).

Organohydroperoxides and 1O_2^* promote ChrR turnover by different mechanisms

While the presence of either 1O_2^* or an organohydroperoxide like t-BOOH increases σ^E^ activity (Glaeser et al., 2011; Nam et al., 2013), they appear to inactivate ChrR by different mechanisms. By comparing the ability of either 1O_2^* or t-BOOH to increase σ^E^ activity in cells containing wild-type ChrR, a truncated ChrR85 protein, variant ChrR proteins with single amino acid changes (Greenwell et al., 2011), and host mutants with defects in furan fatty acid synthesis (Nam et al., 2013), one can genetically separate the effects of 1O_2^* and t-BOOH on σ^E^ activation. To explain this observation, it has been proposed that ChrR inactivation in the presence of 1O_2^* (requires a ChrR-CLD) or t-BOOH (occurs in cells lacking an intact ChrR-CLD) do not occur by identical mechanisms (Greenwell et al., 2011; Nam et al., 2013).

The biological response to 1O_2^*

Identifying the members of a transcriptional regulon is often instrumental to understanding functions needed during a stress response (Guisbert et al., 2008). By combining computational (phylogenetic clustering), in vitro (transcription assays) and in vivo (gene fusions, global gene expression or chromatin immunoprecipitation) analyses, we found that 1O_2^* activated ~160 genes (Fig. 5), with the majority of them the indirect result of this ROS activating a σ factor cascade that includes RpoH_{II}, one of two *Rb. sphaeroides* σ^E^ homologs (Green and Donohue, 2006; Dufour et al., 2008; 2012; Ziegelhoffer and Donohue, 2009). Indeed, a combination of *in vitro* and *in vivo* studies showed that < 10% of the 1O_2^* activated genes (~13/160 genes) were directly transcribed by σ^E^-containing RNA polymerase (Anthony et al., 2005; Green and Donohue, 2006; Dufour et al., 2008; 2012; Dufour and Donohue, 2012). Others predict that a larger number of genes are part of the σ^E^-dependent response to 1O_2^* (Glaeser et al., 2007; Nuss et al., 2009; Berghoff et al., 2009), but many studies often do not distinguish direct and indirect effects of 1O_2^* on downstream gene expression. The direct targets of *Rb. sphaeroides* σ^E^ encode proteins that could prevent or remove damage from lipid peroxidation, enzymes that can repair mutations, electron transport metalloproteins, another alternative sigma factor RpoH_{II} and proteins of unknown function (Anthony et al., 2005; Green and Donohue, 2006; Watabe et al., 2007; Dufour et al., 2008; 2012; Ziegelhoffer and Donohue, 2009). The finding that rpoH_{II} transcription is absolutely dependent on σ^E^ (Fig. 5) predicted that 1O_2^* activated a transcriptional cascade and that both σ^E^ and RpoH_{II} have a role in this stress response (Anthony et al., 2005; Green and Donohue, 2006; Watabe et al., 2007; Dufour et al., 2008; 2012). As predicted, 1O_2^* is bactericidal to cells lacking σ^E^ or RpoH_{II} (Anthony et al., 2005; Green and Donohue, 2006; Nuss et al., 2010). Additional direct σ^E^ target genes are needed for rapid ChrR proteolysis, while others encode proteins that could potentially reduce products of olefin oxidation, prevent oxidation of unsaturated fatty acids, serve as electron carriers or repair damaged macromolecules (Dufour et al., 2008; Ziegelhoffer and Donohue, 2009; Lemke et al., 2014). The ~145 direct RpoH_{II} target genes (Fig. 5) encode bioenergetic enzymes that contain oxidant-sensitive metal centers (NADH dehydrogenase, etc.), metalloenzymes that synthesize cofactors for bioenergetic enzymes.
(tetrapyrroles, quinone, etc.) and glutathione-dependent enzymes that can repair oxidized macromolecules (Dufour et al., 2012; Dufour and Donohue, 2012). Numerous $\sigma^E$ and RpoH$_{II}$ targets have no known function (Dufour et al., 2008; 2012; Dufour and Donohue, 2012), illustrating how little is known about the cellular and biological response to $1O_2$ and organoperoxide stress.

Unlike *E. coli*, which contains a single heat shock $\sigma$ factor (Guisbert et al., 2008), *Rb. sphaeroides* contains two homologs, RpoH$_I$ and RpoH$_{II}$ (Karls et al., 1998; Green and Donohue, 2006). $1O_2$ is not bacteriocidal to $\Delta$RpoH$_I$ cells, and RpoH$_I$ activity is increased during heat shock (Karls et al., 1998; Green and Donohue, 2006; Dufour et al., 2012). Indeed, many of the $\sim$130 genes that are directly transcribed by RpoH$_I$ encode homologs of typical heat shock proteins (Green and Donohue, 2006; Dufour and Donohue, 2012), so it appears that its primary role is in thermal adaptation, similar to that of *E. coli* $\sigma^{32}$ (Guisbert et al., 2008). However, many of the 45 genes which are directly transcribed by both RpoH$_{II}$ and RpoH$_I$ (Fig. 5) encode proteins that could act in both $1O_2$ and heat stress (Green and Donohue, 2006; Dufour et al., 2012; Dufour and Donohue, 2012).

Often, members of a stress regulon are part of a homeostatic loop that is needed to activate the response (Ades et al., 2003; Guisbert et al., 2008). This appears to be true for $1O_2$ stress, since mutants lacking $\sigma^E$ target genes that produce furan-containing fatty acids are defective in increasing activity of this $\sigma$ factor when they are exposed to this ROS (Nam et al., 2013). However, cells lacking other ECF11 regulon members have normal activation of $\sigma^E$ activity and rates of ChrR proteolysis in the presence of $1O_2$ (Hendrischek et al., 2007; Nam et al., 2013; Nuss et al., 2013).

**Conservation of the ECF11 system across the bacterial phylogeny**

Selective pressures experienced by cells in nature can dictate a relationship between signals and regulated genes, so the function of a given regulon may have evolved to accommodate variance in environmental conditions across cells with different lifestyles or habitats. For example, a comparative analysis of the *E. coli* $\sigma^E$ regulon in nine $\gamma$-proteobacteria revealed the existence of a ‘core regulon’ that encodes functions involved in envelope stress, plus an ‘extended regulon’ that includes functions related to pathogenesis or symbiosis, and led to the proposal that host-microbe interactions also activate this stress response (Rhodius et al., 2006).

A similar phylogenetic analysis of $\sigma^E$-ChrR homologs across bacterial divisions also suggested that this system evolved prior to the divergence of the $\alpha$- and $\gamma$-proteobacteria, and shows that it includes species which have both photosynthetic and non-photosynthetic lifestyles (Dufour et al., 2008). By analyzing these genomes for genes orthologous to those transcribed by $\sigma^E$ and promoters that contain the motif recognized by this Group IV alternative $\sigma$, it was found that many of the direct *Rb. sphaeroides* $\sigma^E$-ChrR regulon members were present and predicted to contain a $\sigma^E$ promoter in these diverse species. The $\sigma^E$ targets that were most conserved across species, which comprise a so-called ‘core $\sigma^E$-ChrR regulon’ of $\sim$8 genes, include the *rpoEchrR* operon and genes...
involved in synthesis of furan fatty acids that are required for ChrR turnover in the presence of 1O2 (Dufour et al., 2008). Therefore, it is possible that the photosynthetic and non-photosynthetic species which contain σE-ChrR homologs both encounter 1O2 in nature (Dufour et al., 2008). The observation that proteins required for ChrR turnover in the presence of 1O2 are conserved members of the core σE regulon gene suggests that a similar homeostatic feedback loop activates this stress response in other bacteria (Dufour et al., 2008; Nam et al., 2013; Lemke et al., 2014). As predicted by these phylogenetic analyses, the Caulobacter crescentus σE-ChrR system was rapidly activated by 1O2 and organic hydroperoxides and exhibited a slower response to other inducers (Lourenco and Gomes, 2009), suggesting other signals or pathways activate the ECF11 regulon in this and other species.

Comparative genomics also identified another group of genes directly transcribed by σE that are not highly conserved among bacterial species and constitute an ‘extended σE-ChrR regulon’ (Dufour et al., 2008). This extended σE-ChrR regulon contains numerous genes of unknown function, illustrating the potential to reveal new biology by elucidating their function. There are also gene sets which are only part of the extended σE-ChrR regulon in selected bacteria, suggesting they encode functions associated with the lifestyle or ecological niche of these organisms (Dufour et al., 2008).

Future directions

Like other optogenetic circuits (Zhao et al., 2018), one advantage of studying Rh. sphaeroides σE-ChrR is the ease of controlling production of the stimulating signal, 1O2, by the presence or absence of light (Anthony et al., 2005). In addition, biochemical, genetic, genomic and computational methods were combined to reveal control principles of this system, define processes that are impacted by 1O2 formation, and predicted the properties of σE-ChrR networks in other bacteria that contain ECF11 proteins (Newman et al., 1999; Anthony et al., 2003; Campbell et al., 2007; Dufour et al., 2008; 2012; Greenwell et al., 2011; Dufour and Donohue, 2012; Nam et al., 2013; Lemke et al., 2014).

Despite the knowledge accumulated by studying Rh. sphaeroides σE-ChrR, major gaps remain in our understanding of important aspects of its function. For example, to understand how Rh. sphaeroides σE activity is increased, information is needed on the events and proteins that regulate ChrR turnover in the presence of 1O2. Other needs include insight into a direct interaction of 1O2, peroxidation products of fatty acids, or other biomolecules with the ChrR-ASD and ChrR-CLD, the protease(s) that degrade ChrR, and the signal transduction pathway used to promote turnover of a cytoplasmic anti-σ factor by a membrane ROS. In addition, identifying the function of genes that are directly transcribed by σE-containing RNA polymerase but only found in selected species (extended members of the σE regulon) can provide needed insight into stress response functions associated with lifestyles or ecological niches of these bacteria.

It is crucial to point out that many of the above questions illustrate knowledge gaps for other Group IV alternative σs, so it is likely that answers obtained by analyzing ECF11 proteins will have broad applicability to other regulatory networks. In this way, analysis of the Group IV alternative σs will continue to illuminate new features of biological processes across the bacterial phylogeny.

Accession numbers

The atomic coordinates for proteins discussed are deposited in the Protein Data Bank (http://wwpdb.org/) under ID codes 2Q1Z, 2Z2S, and 1OR7. The gene expression and chromatin immunoprecipitation data sets discussed can be found in the Gene Expression Omnibus through series accession number GSE39806 (https://www.ncbi.nlm.nih.gov/geo/).

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