Early evolution of purple retinal pigments on Earth and implications for exoplanet biosignatures

Shiladitya DasSarma1 and Edward W. Schwieterman2,3,4,5

1Department of Microbiology and Immunology, University of Maryland School of Medicine, Institute of Marine and Environmental Technology, Baltimore, MD, USA; 2Department of Earth Sciences, University of California, Riverside, CA, USA; 3NASA Postdoctoral Program Fellow, Universities Space Research Association, Columbia, MD, USA; 4NASA Astrobiology Institute’s Alternative Earths and Virtual Planetary Laboratory Teams and 5Blue Marble Space Institute of Science, Seattle, WA, USA

Abstract

We propose that retinal-based phototrophy arose early in the evolution of life on Earth, profoundly impacting the development of photosynthesis and creating implications for the search for life beyond our planet. While the early evolutionary history of phototrophy is largely in the realm of the unknown, the onset of oxygenic photosynthesis in primitive cyanobacteria significantly altered the Earth’s atmosphere by contributing to the rise of oxygen ~2.3 billion years ago. However, photosynthetic chlorophyll and bacteriochlorophyll pigments lack appreciable absorption at wavelengths about 500–600 nm, an energy-rich region of the solar spectrum. By contrast, simpler retinal-based light-harvesting systems such as the halorhodopsin purple membrane protein bacteriorhodopsin show a strong well-defined peak of absorbance centred at 568 nm, which is complementary to that of chlorophyll pigments. We propose a scenario where simple retinal-based light-harvesting systems like that of the purple chromoprotein bacteriorhodopsin, originally discovered in halophilic Archaea, may have dominated prior to the development of photosynthesis. We explore this hypothesis, termed the ‘Purple Earth,’ and discuss how retinal photopigments may serve as remote biosignatures for exoplanet research.

Background

The major events sparking life on Earth on our 4.6-billion-year-old planet remain enigmatic, although there is general agreement that first life likely arose about 3.7–4.1 billion years ago, during the early Archean or late Hadean eons (Abramov and Mojzsis, 2009; Deamer, 2011; Bell et al., 2015; Knoll, 2015). Evidence for the presence of isoprenoid compounds has been reported in ancient sediments not long after, suggesting the early rise of Archaea (Hahn and Haug, 1986; Ventura et al., 2007). The early rise of Archaea is also suggested by phylogenetic studies, although lateral gene transfers have complicated their interpretation (Lange et al., 2000; Kennedy et al., 2001; Brochier-Armanet et al., 2011; Hoshino and Gaucher, 2018). Stromatolites representing fossilized microbial mats have been estimated to be up to 3.7 billion-years-old (Walter et al., 1980; Vankranendonk et al., 2008; Nutman et al., 2016) and radiocarbon dating has shown 12C enrichment from this early period, consistent with the development of photosynthetic microorganisms (Ohtomo et al., 2014). There is wide agreement that anoxicogenic photosynthesis preceded oxygenic photosynthesis, though the length of the interval for this transition is uncertain (Olson, 2006; Buick, 2008; Rothschild, 2008). Some geochemical proxy records suggest that the earliest oxygenic photosynthesizers may have appeared by ~2.9–3 Ga with geochemical sinks arrests oxygen’s accumulation for a time (Nisbet et al., 2007; Planavsky et al., 2014). Ultimately, because of oxygenic photosynthesis and additional, poorly understood factors, the Earth experienced a Great Oxidation Event about 2.3 billion years ago, which indelibly altered the prevailing chemical conditions of our planet’s atmosphere (Kump, 2008; Lyons et al., 2014; Luo et al., 2016).

What were the important evolutionary events predating the rise of photosynthesis during the early history of life on Earth? Although the events during this very early time are not clear, in this paper, we discuss a speculative hypothesis for early evolution, called the ‘Purple Earth,’ which posits the rise of retinal pigment-based phototrophic life forms on Earth’s surface prior to anoxicogenic and oxygenic photosynthesis. In this view, retinal pigments may have competed with and affected the evolution of photosynthetic pigments and indeed still complements them today in Earth’s oceans and other environments. Early microorganisms employing retinal pigments for generating metabolic energy may have dominated, as halophilic Archaea do today in hypersaline environments, providing a scenario which may serve to guide our search for detectable biosignatures on other worlds.
Early evolution on Earth

During the first half of Earth’s history, stretching over 2 billion years, dramatic and long-lasting evolutionary inventions occurred through processes that we are only beginning to understand (Fig. 1; Deamer, 2011; Knoll, 2015). They include prebiotic evolution and the development of cellularity, the foundation of the last universal common ancestor (LUCA) and evolution of the universal genetic code (Fenchel, 2002). Other factors critical for the success of early life were the evolution of transmembrane potential and chemiosmotic coupling for creating and storing bioenergy, pigments for the capture of light energy for phototrophy and photosynthesis and respiratory chains for anaerobic and aerobic respiration (Zannoni, 2004). In addition, a ‘frozen accident’ has been proposed to establish the genetic code as a universal feature in all extant life on Earth (Crick, 1968; Söll and RajBhandary, 2006). During the earliest period in evolutionary history, well-defined phylogenetic lineages may not yet have been established; instead extensive lateral gene transfers allowed for ready sharing of new innovations until such time when the last common ancestor experienced competitive selective forces and diverged into the three primary ‘Domains of Life’ (Woese, 2002).

Even prior to the evolution of the three Domains, the development of a protocol must have been facilitated by the evolution of a water-tight cell membrane as a permeability barrier, preventing the free diffusion of chemicals into and out of cells, critical for generating and storing cellular energy (Gunner et al., 2013). The intracellular milieu provided a microenvironment in which biomolecular functions, such as the biosynthesis of macromolecules and the genetic code could be established. Transmembrane ion pumps acting as energy transduction and storage systems must have been among the earliest inventions. In one scenario proposed here, a simple light-harvesting system incorporating a retinal pigment allowed light-driven proton pumping and led to a proton-motive gradient. Based on its ubiquity, the transmembrane electrochemical potential (i.e. proton-motive gradient) as well as phosphoric anhydride bonds, such as in adenosine triphosphate (ATP), became established and universal due to their kinetic stability and bioenergetic capabilities in the aqueous environment. Subsequently, retinal as well as a variety of more complex anaerobic and oxygenic light-harvesting systems were invented and resulted in the evolution of diverse phototrophic and photosynthetic microorganisms.

Appearance of purple retinal pigments

The earliest life-forms probably arose in the early Archean or possibly late Hadean Eons, with some molecular clock estimates putting life’s origin as early as 4 Ga (Hedges, 2002). While the exact timing of appearance of retinal pigments is not clear, it may have been a very early metabolic invention coincident with or occurring soon after the development of cellular life. A retinal chromophore bound to a single polypeptide allows a system for phototrophy by forming a chromoprotein, like bacteriorhodopsin, in halophilic Archea dominant in hypersaline environments and proteorhodopsin in pelagic bacteria distributed throughout the oceans (Béjà et al., 2001; Stoeckenius et al., 1979). The absorption of light by this chromoprotein in the 490–600 nm region, a highly energy-rich region of the solar spectrum (Fig. 2), is directly coupled to pumping of protons and the resulting electrochemical gradient chemiosmotically drives ATP synthesis. This type of retinal-dependent phototrophy is considerably simpler albeit less efficient than photosynthesis and it neither results in fixation of carbon nor production of oxygen (Pinhassi et al., 2016). Nevertheless, the widespread distribution of retinal chromoproteins in nature and their unique utilization of the energy-rich, yellow-green region of the spectrum for production of cellular energy suggest their early appearance on Earth.

Evidence for the existence of isoprenoid compounds that are part of the biosynthetic pathway to retinal as well as archaeal lipids in the early history of the Earth has also been provided (Hahn and Haug, 1986; Ventura et al., 2007). It is likely that the evolutionary invention of retinal pigments was coincident with other membrane lipids, which together established the molecular basis for chemiosmotic coupling and phototrophic capabilities (Boucher and Doolittle, 2000). Retinal is produced by a branch of the isoprenoid metabolic pathway leading to carotenoids and branched-chain lipids, which are found in cell membranes (Fig. 3). Retinal pigments occur in both major prokaryotic phylogenetic groups, Archea and Bacteria, as well as in eukaryotes, where they are essential components of the visual system (Ernst et al., 2014). Among the pigments prevalent in nature, retinal has a simple structure compared with many others that are used for photosynthesis and respiration, e.g. chlorophyll and other porphyrins, which may be produced by a branch of the tricarboxylic acid (TCA) cycle, a pathway used by all aerobic organisms (Mailloux et al., 2007). These findings, together with the central position of retinal at the intersection of lipid metabolism and bioenergetics, as well as its widespread distribution suggest that retinal played an important role in the early evolution of life on Earth.

The light-driven proton pumping activity of retinal pigments such as the chromoprotein bacteriorhodopsin in the membrane of an early cell would have allowed the development of chemiosmotic coupling, linking of membrane potential to other transmembrane transport processes and ATP synthesis (Stoeckenius et al., 1979).
et al., 1979). A retinal-based phototrophic system clearly represents one of the simplest bioenergetic mechanisms conceivable, requiring only a single opsin inserted in a membrane vesicle and membrane-potential coupled ATP synthase (Fig. 4). Indeed, such a model phototrophic system, inside out, was established in vitro in the 1970s using haloarchaeal bacteriorhodopsin and mitochondrial ATP synthase in artificial lipid vesicles (Racker and Stoeckenius, 1974). This seminal work was credited with helping to establish the validity of Mitchell’s chemiosmotic coupling hypothesis (Mitchell, 1961) and also forms the foundation of one of the simplest and as proposed, earliest metabolic capabilities in evolution, retinal-based phototrophy.

Early Earth environments would have lacked abundant free O₂ in contrast to highly oxic modern environments and required the production of retinal using a terminal oxidative step in a likely strictly anaerobic environment. A number of potential mechanisms have been proposed for generating such an oxidative potential, such as pyrite-induced aqueous hydrogen peroxide and hydroxide radical formation (Borda et al., 2001; Cohn et al., 2006). Other anaerobic oxidation reactions are also known, such as anaerobic oxidation of methane and ammonium and transformation of isoprenoids by anaerobic microorganisms (Hallam et al., 2004; Hylemon and Harder, 1998; Strous and Jetten, 2004).

Also notable is that modern halophilic Archaea are facultative, rather than obligate aerobes and can respire nitrate and TMAO/ DMSO (Mancinelli and Hochstein, 1986; Müller and DasSarma, 2005). Indeed, Haloarchaea have been shown to engage in phototrophy in microaerobic or anoxic laboratory conditions (Sumper et al., 1976; DasSarma et al., 2012; Laye et al., 2017). Additionally, a considerable amount of evidence suggests that the genes for aerobic respiration were laterally transferred to halophilic Archaea (Kennedy et al., 2001) and their ultimate origin may have been as an anaerobic chemolithoautotrophic methanogen (Nelson-Sathi et al., 2012; Aouad et al., 2018). Hence, haloarchaeal phototrophic metabolism was probably developed well before genes for aerobic respiration were acquired, possibly in Archaea inhabiting hypersaline environments (Stevenson et al., 2015). While these modern haloarchaeal organisms have certainly changed over the eons from the original retinal-based phototrophs, the available evidence illustrates the potential capacity of Haloarchaea to have survived the anaerobic conditions that prevailed on ancient Earth.

In modern halophilic Archaea, the retinal protein bacteriorhodopsin trimers form a hexagonal lattice which can cover a large fraction of the cell surface (Stoeckenius et al., 1979), imparting a bright purple colour to some salt ponds where they dominate (Fig. 5). The resulting purple membrane can be easily isolated using sucrose
density gradients and has been the subject of extensive structural and functional analysis of transmembrane ion translocation (Henderson and Unwin, 1975; Stoeckenius et al., 1979; Krebs and Khorana, 1993; Hirai et al., 2009). Bacteriorhodopsin is a prototype of integral membrane proteins with seven-transmembrane α-helical segments where the retinal chromophore is bound by a Schiff’s base linkage to the ε-amino group of a lysine residue (Bayley et al., 1981). The photobiology of bacteriorhodopsin has been intensively studied, including characterization of the molecular dynamics and role of retinal during photocycling (Hirai et al., 2009). The bacteriorhodopsin resting state is notable for the characteristic colour purple resulting from the strong absorption peak maximum at 568 nm in the yellow-green region of the spectrum.

Spectral complementarity of photopigments
Comparison of the spectrum of bacteriorhodopsin with the major photosynthetic pigments containing chlorophyll and bacteriochlorophylls shows them to be complementary, i.e. the purple pigment absorption peaks in the region with a trough for the green pigments (Fig. 2a). If the evolution of the simpler retinal pigments predated chlorophyll pigments in the evolutionary history, as proposed, it is conceivable that they may have affected the development of the spectral characteristics of evolving chlorophyll pigments (Goldsworthy, 1987; DasSarma, 2006). This may have been the consequence of filtering of light by retinal chromoproteins, resulting in a deficit of wavelengths of light centred around the peak of bacteriorhodopsin absorption in the yellow-green region of the spectrum. The resulting deficit, particularly in a stratified community of microorganisms such as those observed in stromatolites, may explain why chlorophylls and bacteriochlorophylls evolved to absorb relatively little light in the yellow-green energy-rich portion of the electromagnetic spectrum, instead absorbing light primarily in the flanking blue and red regions of the solar spectrum.

Modern stromatolites represent microbial communities with ongoing spectral competition and spectral tuning of chromoproteins (Croce and van Amerongen, 2014). Phototrophic and
photosynthetic microorganisms in microbial mats are commonly stratified based on predictable photosystem characteristics as well as oxygen requirements. In such communities, oxygenic cyanobacteria are found near the surface oxic zone while anaerobic phototrophic and photosynthetic microbes are buried at lower anoxic regions. If these modern stratified microbial communities are like those present in ancient stromatolites, filtering of wavelengths of light would have been an important and pervasive characteristic of microbial communities. Modern microbial communities all over the world support planktonic retinal-containing halophilic Archaea and Bacteria inhabiting brines above photosynthetic mats (Cohen and Rosenberg, 1989). If co-evolution of retinal and chlorophyll photopigments occurred in deep evolutionary history, stratification within such niches may have played an important role in the evolution of spectral properties.

Importantly, modern rhodopsin-based phototrophy is present throughout the oceanic and terrestrial biosphere, including non-hypersaline conditions and environments that may have been common in the distant geologic past. While originally discovered in halophilic Archaea, microbial rhodopsins are common in oceanic planktonic Bacteria (Kandori, 2015). For example, *Pelagibacter ubique* is a widely distributed marine bacterium that produces the retinal chromoprotein, proteorhodopsin, with the ability to use its light-driven proton pumping activity for energy generation. Moreover, the absorption characteristics of proteorhodopsins show spectral variations in oceanic planktonic bacteria isolated from different depths, consistent with spectral tuning (Ranarajan et al., 2007). Rhodopsin-based phototrophy in the ocean may be so widespread as to rival the total light capture of photosynthesizers (Brown, 2014; Gómez-Consarnau et al., 2017). Furthermore, metagenomic analyses have recently uncovered evidence for the widespread presence of rhodopsins in the terrestrial biosphere including in the phyllosphere (leaf surfaces) and even edaphic systems and hypolithic communities in the Antarctic Dry Valley (Atamna-Ismaeel et al., 2012; Guerrero et al., 2017). These findings are consistent with the notion that microorganisms evolve specialized photosystems that make use of any available spectral region with sufficient energy. The widespread presence of microbial rhodopsins in modern environments, along with inefficient chlorophyll absorption in the middle of the visible spectrum where rhodopsin light capture is most efficient, suggests a co-evolution that is consistent with the earlier appearance of retinal-based phototrophy (PBS Eons 2018).

**Rise of photosynthesis**

The rise of anaerobic and oxygenic photosynthesis and retreat of retinal-based life must have occurred in discrete stages, which are not fully understood. For example, the time of appearance of a wide diversity of bacteria with anoxicogenic photosynthetic systems is not precisely known (Jeffrey, 1963; Frigaard et al., 1996; Rothschild, 2008; Chen et al., 2010; Chen and Blankenship, 2011; Croce and van Amerongen, 2014). The purple and green bacteria possessing bacteriochlorophyll may have been an early evolutionary development with photosynthetic reaction centres evolving from electron transport chain components such as cytochromes (Williamson et al., 2011; Mazor et al., 2012). Alternatively, a simplified photosystem may have evolved previously, like those in some heliobacteria (Xiong et al., 1998). In either case, evolution likely first led to the evolution of anoxicogenic photosynthesis with the development of more complex oxygenic photosynthetic membrane systems like those in modern cyanobacteria, including two photosynthetic reaction centres and a host of membrane components, developing later.

With multiple evolutionary steps leading to progressively higher efficiency chlorophyll pigments along with the invention of accessory pigments, photosynthetic microorganisms out-competed retinal-based phototrophic microorganisms in most environments. Evolution of anoxygenic photosynthesizers was followed by oxygenic photosynthesizing cyanobacteria and ultimately eukaryotic algae and plants. Interestingly, a distinct hypothesis that purple sulphur bacteria may have dominated euxinic ocean during the mid-Proterozoic eon has also been proposed, resulting in a second Purple Earth (Brocks et al., 2005; Sanromá et al., 2014), which would have been long after the retreat of retinal-based life dominating the first Purple Earth to ecological niches resembling those of today. The development of eukaryotic algae and complex plants and their spread throughout the terrestrial environment allowed the evolution of land animals and ultimately intelligent life (Catling et al., 2005; Reinhardt et al., 2016). At every step of this progression, it is not clear to what degree evolutionary contingency has played a role and which developments would be inevitable given sufficient time and the appropriate environmental conditions. As a result, the capacity for evolution to generate diverse phototrophic and photosynthetic systems on Earth, even those that do not dominate today, may have considerable implications for the development of novel pigments on other habitable worlds (Johnson et al., 2013).

**Retinal-based phototrophy as an astronomical biosignature**

Regardless of the evolutionary sequence of events leading to retinal phototrophy on Earth, analog photopigments may have arisen independently in other habitable environments in the universe. For example, exoplanets within the habitable zones of most stars would receive ample photon fluxes to power significant levels of (bacteria)chlorophyll or rhodopsin analog phototrophy with some differences in total capacity based on the photospheric temperature and consequent spectral energy distribution of those stars (Kiang et al., 2007b; 2007a; Komatsu et al., 2015; Ritchie et al., 2018). One exception may be the dimmest and reddest M-stars, which produce the least flux in the 400–700 nm wavelength range (Fig. 2b). For these stellar systems, total global productivity may be photon-limited rather than reductant or nutrient limited as it is on Earth (Lehmer et al., 2018). However, FGK stellar systems are the more likely targets for future space-based direct-imaging missions capable of detecting astronomical biosignatures. This is due to the wider angular separation of star and planet in the habitable zone (Stark et al., 2015; 2014) and the productivity of biospheres on planets orbiting these stars would not be photon-limited. It is, therefore, worthwhile to examine what the remote signatures of rhodopsin-like phototrophy would be on exoplanets and how they would compare to those produced by analogs to chlorophyll-based photosynthesis.

The most commonly referenced surface signature of life is the vegetation red edge (VRE), the steep increase in reflectivity of vegetation (primarily green vascular plants) at ~700 nm (Gates et al., 1965; Knipling, 1970). This increase is due to the contrast between the absorption of chlorophyll at red wavelengths and its high albedo at infrared wavelengths due to intracellular scattering. The VRE effect is commonly used to map vegetation by employing broadband observations from Earth-observing satellites (Huete et al., 1994; Tucker et al., 2005). While the VRE has been extensively examined as a possible exoplanet
biosignature (Sagan et al., 1993; Arnold et al., 2002; Des Marais et al., 2002; Seager et al., 2005; Brandt and Spiegel, 2014), its applicability is limited to those planets that have, like Earth, evolved chlorophyll-analog (i.e., red-absorbing, infrared reflecting) powered vegetation with significant continental surface covering fractions.

Even on Earth, green vascular planets have only existed for just the last ~10% of the planet's history, about 470 million years out of 4.6-billion years (Kenrick and Crane, 1997). In order to address the limited time of presence of the VRE, astrobiologists interested in remote biosignatures have begun to consider and catalog surface reflectance signatures from a diverse array of known pigmented organisms including oxygenic and anoxygenic photosynthesizers, rhodopsin-based phototrophs and non-photosynthetic microbes that use pigments as a UV screen or antioxidant, or for other purposes (DasSarma, 2006; Kiang et al., 2007b, 2007a; Cockell, 2014; Hegde et al., 2015; Poch et al., 2017; Schwieterman, 2018; Schwieterman et al., 2018, 2015). The possible existence of a Purple Earth extends and expands the possible biological history of a planet when alternate biosignatures may be detectable, and it also enhances the number of possible evolutionary trajectories for which surface biosignatures may be found.

The photochemical properties of known prokaryotic rhodopsins on Earth are particularly worthy of study as a potential remote biosignature because of their capacity to generate chemical energy using an energy-rich portion of the electromagnetic spectrum. The most consequential difference between rhodopsin and chlorophyll-based phototrophy is the wavelength of maximum absorption. While the absorption peak of chlorophyll a is near 700 nm, bacteriorhodopsin absorption peaks near 570 nm. However, the expression of this signal would differ depending on whether the phototrophic organisms were on dry land or suspended in aquatic environments.

A true bacteriorhodopsin-based analog to terrestrial vegetation would possess a 'green-edge' comparable with the vegetation 'red-edge.' Fig. 6a illustrates the differences between the reflective spectra of a red-edge producing conifer forest (Baldridge et al., 2009) and the green-edge producing Haloarchaea (Schwieterman et al., 2015). While the green colour of the conifer forest results from inefficient absorption in a broad wavelength region centred

---

**Fig. 6.** Surface signatures of retinal and chlorophyll-based phototrophy. (a) Reflectance spectrum of a conifer forest (Baldridge et al., 2009) and a culture of the phototrophic archaeon *Halobacterium* sp. (Schwieterman et al., 2015). (b) Environmental spectrum of a halophile-dominated saltern pond in San Francisco Bay (Dalton et al., 2009). (c) Simulated spectra of planets consisting of 100% sterile ocean, conifer forest, or a halophile-dominated saltern pond under an Earth-like atmosphere generated with a radiative transfer model (Schwieterman et al., 2015).
near 550 nm, the pink colour of halophiles results from their high reflectivity at orange and red wavelengths due to both bacteriorhodopsin and carotenoid pigments such as bacterioruberins (Kushwaha and Kates, 1979; Oren et al., 1992; Oren and Dubinsky, 1994). Green plants are similarly bright in the infrared due to their high reflectivity on the non-visible, long-wavelength side of the VRE. Notably, a wide variety of other organisms have spectral 'edges' at various visible wavelengths. Hegde et al. (2015) conducted an extensive series of reflectance measurements of plated cultures of extremophiles from 350 to 2500 nm showing a distribution of 'edge' features for various phototrophic and non-phototrophic species throughout the UV-visible spectrum. The radiotolerant species Deinococcus radiodurans also possess a ‘green-edge’ in plated cultures due to its primary pigment deinoxanthin (Cockell, 2014; Schwiegerman et al., 2015), which likely functions as an antioxidant. Depending on the environmental context, some of these pigments may also serve as alternative bio-signatures. Rhodopsin-based (and other) phototrophs may have both biological and detectability advantage, however, in that they can harvest light energy for growth and accumulate at the surface of aquatic environments. Consequently, it is also important to consider the spectral differences between plated and suspended cultures, which would map to different planetary environments (e.g., land versus ocean).

The spectral signature of pigmented organisms like Haloarchaea suspended in a lake or ocean would also be affected by the low reflectivity and strong absorption properties of aquatic environments. For example, halophilic communities present saltern ponds possess a peak in brightness near ~680 nm due to increasing reflectivity of bacteriorhodopsin and carotenoid pigments from the green to the red combined with strong water absorption at the reddest wavelengths (Fig. 6b; also see Dalton et al., 2009). Importantly, the brightness of halophilic pigments at orange and red wavelengths confers a detectability advantage over chlorophyll-containing cyanobacteria and algae suspended in water, because chlorophyll’s high infrared reflectivity is counteracted by water vapour absorption, while chlorophyll is most absorptive at wavelengths where water is relatively transparent.

The remote signatures of these phototrophic organisms would further change from the spectral impact (i.e., absorption and scattering) of the overlying atmosphere (Fig. 6c). The spectral signatures of halophilic organisms are again somewhat favoured in this case because of the impact of overlying water vapour absorption nearly coincident with the VRE. Of course, the detectability of these signatures on an exoplanet will also be strongly sensitive to the land covering fraction, cell densities if suspended in water or brine and cloud cover effects (Sanromá et al., 2014; Schwiegerman et al., 2015). The detectability potential of retinal pigments and other halophilic pigment-analogs should be considered when anticipating the variety of potential surface biosignatures of exoplanets.

The capacity to detect surface biosignatures is an ongoing consideration in the design mandate of large, space-based telescopes (Fujii et al., 2018; Schwiegerman et al., 2018) such as the conceived HabEx and LUVIOR/HDST missions (Dalcanton et al., 2015; Mennesson et al., 2016; Rauscher et al., 2016; Bolcar et al., 2017). Direct imaging spectra and spectrophotometry will allow characterization of the surfaces of terrestrial planets in the habitable zone, producing constraints on surface types, including surface biosignatures, providing the cloud covering fraction is sufficiently low (Sanromá et al., 2014, 2013). The detectability potential of retinal photopigments and halophile-analogs suggests wavelengths shortward of the traditional VRE (λ< 700 nm) will be important to observe and analyse for ‘edge’ features suggestive of diverse phototrophic pigments and should be considered in the search for life outside the solar system.

Concluding remarks

Although of enormous scientific interest, our understanding of the early evolutionary history of phototrophic life on Earth has remained limited. We propose here that the biochemical simplicity of retinal-based phototrophy, the spectral complementarity of bacteriorhodopsin pigments with chlorophylls and the newly uncovered widespread diversity of microbial rhodopsins throughout aquatic and terrestrial ecosystems are suggestive of the fundamental role retinal may have played in the early history of life on Earth. We posit here that domination by retinal-based phototrophs in the early history of life may have created the first ‘Purple Earth’ that at some point gave way to modern photosynthesizers before the rise of atmospheric oxygen. If correct, this early phototrophic metabolism would have greatly shaped the evolution of photosynthesis and indeed much of life on Earth. In fact, we know it continues to play a significant role in many environments today.

To test this Purple Earth hypothesis, future work should further explore natural communities of retinal-based phototrophs in diverse environments (e.g. arid, high altitude and polar locations). Additional studies are needed to explore the diversity and light capture capacity of retinal-based phototrophy in modern environments as they may continue to reveal unexpected roles and niches for this metabolism and inform its evolutionary origin. Additionally, future genomic analyses should be designed to consider the importance of the timing of the introduction of aerobic respiration in Haloarchaea in relation to the development of phototrophy during their metamorphosis from anaerobic chemolithoautotrophic methanogens to aerobic phototrophs.

Considering an even broader view, the quest to understand the origin of life and early evolutionary events on our planet has gained increasing urgency with the discovery of thousands of new extrasolar planets, many of which are within the habitable zones of their host stars. Consequently, we may soon have the ability to characterize other potentially living worlds and finally answer the age-old question ‘Are we alone in the universe?’ To realize this goal, however, we need to improve our understanding of major events sparking life on Earth and determine what biosignatures early life produced, especially those which may be detectable by remote sensing.

Simple retinal-based light-harvesting systems like that of the purple chromoprotein bacteriorhodopsin, may potentially serve as remote biosignatures for exoplanet research through the search for brightness peaks about 680 nm like that seen in hypersaline environments on Earth or by spectral ‘edges’ at green-yellow wavelengths (~550 nm) analogous to the traditional vegetation ‘red-edge’ seen at 700 nm. These features are within the wavelength sensitivity window of planned next-generation space-based telescopes capable of directly imaging exoplanets and should be considered in the search for life in the universe.

Acknowledgements

Exobiology research in the S.D. laboratory is supported by NASA grant NNX15AM07G. E.S. is supported by a NASA Postdoctoral Fellowship, administered by the Universities Space Research Association and by the NASA Astrobiology Institute’s Alternative Earths and Virtual Planetary Laboratory teams under Cooperative Agreement Nos. NNA15BB03A and NNA13AA93A, respectively. We thank Priya DasSarma for critical reading of the manuscript.
Huiet A, Justice C and Liu H (1994) Development of vegetation and soil indexes for MODIS-EOS. Remote Sensing of Environment 49, 224–234.

Hylemon PB and Harder J (1998) Biotransformation of monoterpenes, bile acids, and other isoprenoids in anaerobic ecosystems. FEMS Microbiology Reviews 22, 475–488.

Jeffrey SW (1963) Purification and properties of chlorophyll c from Sargassum fluvianum. Biochemical Journal 86, 313.

Johnson N, Zhao G, Caycedo F, Manrique P, Qi H, Rodriguez F and Quiroga I (2013) Extreme alien light allows survival of terrestrial bacteria. Scientific Reports 3, 2198.

Kandori H (2015) Ion-pumping microbial rhodopsins. Frontiers in Molecular Biosciences 2, 52.

Kennedy SP, Ng WV, Salzberg SL, Hood L and DasSarma S (2015) Understanding the adaptation of species NRC-1 to its extreme environment through computational analysis of its genome sequence. Genome Research 11, 1641–1650.

Kennicutt R and Crane PR (1997) The origin and early evolution of plants on land. Nature 389, 33–39.

Kiang NY, Segura A, Govindjee, Blankenship RE, Cohen M, Kandori H, Lehmer OR, Catling DC, Parenteau MN and Hoehler TM (2008) Biotransformation of monoterpenes, bile acids and trimethylamine N-oxide as terminal electron acceptors. Journal of Bacteriology 190, 5198–5203.

Knipling EB (1963) Purification and properties of chlorophyll c from Sargassum fluvianum. Biochemical Journal 86, 313.
Rothschild LJ (2008) The evolution of photosynthesis…again? Philosophical Transactions of the Royal Society B 363, 2787–2801.

Sagan C, Thompson WR, Carlson R, Garnett D and Hord C (1993) A search for life on Earth from the Galileo spacecraft. Nature 365, 715–721.

Sanromé E, Pallé E and García Munoz A (2013) On The effects of the evolution of microbial mats and land plants on the earth as a planet. Photometric and spectroscopic light curves of paleo-Earths. The Astrophysical Journal 766, 133.

Sanromé E, Pallé E, Parenteau MN, Kiang NY, Gutierrez-Navarro aM, Lópe r R and Montañés-Rodriguez P (2014) Characterizing the purple Earth: modeling the globally integrated spectral variability of the Archean Earth. The Astrophysical Journal 780, 52.

Schwieterman EW (2018) Surface and temporal biosignatures. In Deeg H and Belmont J (eds), Handbook of Exoplanets. Cham, Switzerland: Springer International Publishing, pp. 1–29. DOI: 10.1007/978-3-319-30648-3_69-1.

Schwieterman EW, Cockell CS and Meadows VS (2015) Nonphotosynthetic pigments as potential biosignatures. Astrobiology 15, 341–361.

Schwieterman EW, Kiang NY, Parenteau MN, Harman CE, DasSarma S, Fisher TM, Arney GN, Hartnett HE, Reinhard CT, Olson SL, Meadows VS, Cockell CS, Walker SL, Grenfell JL, Hegde S, Rugheimer S, Hu R and Lyons TW (2018) Exoplanet biosignatures: a review of remotely detectable signs of life. Astrobiology 18, 663–708.

Seager S, Turner EL, Schafer J and Ford EB (2005) Vegetation’s Red edge: a possible spectroscopic biosignature of extraterrestrial plants. Astrobiology 5, 372–390.

Segura A, Kredove K, Kasting JF, Sommerlatt D, Meadows V, Crisp D, Cohen M and Mlawer E (2003) Ozone concentrations and ultraviolet fluxes on Earth-like planets around other stars. Astrobiology 3, 689–708.

Söll D and RajBhandary UL (2006) The genetic code - thawing the “frozen accident”. Journal of Biosciences 31, 459–463.

Stark CC, Roberge A, Mandell A and Robinson TD (2014) Maximizing The exoearth candidate yield from a future direct imaging mission. The Astrophysical Journal 795, 122.

Stark CC, Roberge A, Mandell A, Clampin M, Domagal-goldman SD, Michelwain MW and Stapelfeldt KR (2015) Lower limits on aperture size for an exoearth detecting coronagraphic mission. The Astrophysical Journal 808, 149.

Stevenson A, Burkhardt J, Cockell CS, Cray JA, Dijkstraeijhuij J, Fox- Powell M, Kee TP, Kminek G, McGinity TJ, Timmis KN, Timson DJ, Voytek MA, Westall F, Yakimov MM and Hallsworth JE (2015) Multiplication of microbes below 0.690 water activity: implications for terrestrial and extraterrestrial life. Environmental Microbiology 17, 257–277.

Stoeckleinus W, Lozier RH and Bogomolni RA (1979) Bacterio- rhodopsin and the purple membrane of halobacteria. Biochimica et Biophysica Acta (BBA) - Reviews on Bioenergetics 505, 215–278.

Strous M and Jetten MSM (2004) Anaerobic oxidation of methane and ammonium. Annual Review of Microbiology 58, 99–117.

Sumper M, Reitmeier H and Oesterhelt D (1976) Biosynthesis of the purple membrane of Halobacteria. Angewandte Chemie International Edition in English 15, 187–194.

Tucker CJ, Pinzon JE, Brown ME, Slayback DA, Pak EW, Mahoney R, Vermote EF and El Saleous N (2005) An extended AVHRR 8-km NDVI dataset compatible with MODIS and SPOT vegetation NDVI data. International Journal of Remote Sensing 26, 4485–4498.

Vankranendonk M, Philipott P, Lepot K, Bodorkos S and Piranjo F (2008) Geological setting of Earth’s oldest fossils in the ca. 3.5Ga dresser formation, pilbara craton, Western Australia. Precambrian Research 167, 93–124.

Ventura GT, Kenig F, Reddy CM, Schieber J, Frysinger GS, Nelson RK, Dinel E, Gaines RB and Schaeffer P (2007) Molecular evidence of late Archean archaea and the presence of a subsurface hydrothermal biosphere. Proceedings of the National Academy of Sciences 104, 14260–14265.

Walter MR, Buick R and Dunlop JSR (1980) Stomatolites 3,400–3,500 Myr old from the North pole area, Western Australia. Nature 284, 443–445.

Williamson A, Conlan B, Hillier W and Wydrzynski T (2011) The evolution of photosystem II: insights into the past and future. Photosynthesis Research 107, 71–86.

Woese CR (2002) On the evolution of cells. Proceedings of the National Academy of Sciences 99, 8742–8747.

Xiong J, Inoue K and Bauer CE (1998) Tracking molecular evolution of photosynthesis by characterization of a major photosynthesis gene cluster from Heliobacillus mobilis. Proceedings of the National Academy of Sciences 95, 14851–14856.

Zannoni D (ed.) (2004) Respiration in Archea and Bacteria, Advances in Photosynthesis and Respiration. Netherlands, Dordrecht: Springer. doi: 10.1007/978-1-4020-3163-2