



Towards a Systems Biology Approach to Understanding the Lichen Symbiosis: Opportunities and Challenges of Implementing Network Modelling

Hadi Nazem-Bokaei^{1,2,3*}, Erik F. Y. Hom⁴, Andrew C. Warden², Sarah Mathews⁵ and Cécile Gueidan^{1*}

OPEN ACCESS

Edited by:

Enrica Pessione,
University of Turin, Italy

Reviewed by:

Tomislav Cernava,
Graz University of Technology, Austria
Christoph Scheidegger,
Swiss Federal Institute for Forest,
Snow and Landscape Research
(WSL), Switzerland

*Correspondence:

Hadi Nazem-Bokaei
Hadi.Nazem-Bokaei@csiro.au
Cécile Gueidan
Cecile.Gueidan@csiro.au

Specialty section:

This article was submitted to
Systems Microbiology,
a section of the journal
Frontiers in Microbiology

Received: 14 February 2021

Accepted: 09 April 2021

Published: 03 May 2021

Citation:

Nazem-Bokaei H, Hom EFY,
Warden AC, Mathews S and
Gueidan C (2021) Towards a Systems
Biology Approach to Understanding
the Lichen Symbiosis: Opportunities
and Challenges of Implementing
Network Modelling.
Front. Microbiol. 12:667864.
doi: 10.3389/fmicb.2021.667864

¹ CSIRO Australian National Herbarium, Centre for Australian National Biodiversity Research, National Research Collections Australia, NCM, Canberra, ACT, Australia, ² CSIRO Land and Water, Canberra, ACT, Australia, ³ CSIRO Synthetic Biology Future Science Platform, Canberra, ACT, Australia, ⁴ Department of Biology and Center for Biodiversity and Conservation Research, The University of Mississippi, University City, MS, United States, ⁵ Department of Biological Sciences, Louisiana State University, Baton Rouge, LA, United States

Lichen associations, a classic model for successful and sustainable interactions between micro-organisms, have been studied for many years. However, there are significant gaps in our understanding about how the lichen symbiosis operates at the molecular level. This review addresses opportunities for expanding current knowledge on signalling and metabolic interplays in the lichen symbiosis using the tools and approaches of systems biology, particularly network modelling. The largely unexplored nature of symbiont recognition and metabolic interdependency in lichens could benefit from applying a holistic approach to understand underlying molecular mechanisms and processes. Together with ‘omics’ approaches, the application of signalling and metabolic network modelling could provide predictive means to gain insights into lichen signalling and metabolic pathways. First, we review the major signalling and recognition modalities in the lichen symbioses studied to date, and then describe how modelling signalling networks could enhance our understanding of symbiont recognition, particularly leveraging omics techniques. Next, we highlight the current state of knowledge on lichen metabolism. We also discuss metabolic network modelling as a tool to simulate flux distribution in lichen metabolic pathways and to analyse the co-dependence between symbionts. This is especially important given the growing number of lichen genomes now available and improved computational tools for reconstructing such models. We highlight the benefits and possible bottlenecks for implementing different types of network models as applied to the study of lichens.

Keywords: systems biology, network modelling, signalling, metabolic model, lichen symbiosis

INTRODUCTION

Lichens are often seen as a typical example of successful and sustainable symbiotic interactions between micro-organisms (Ahmadjian, 1993; Honegger, 1998). With the long evolutionary history of these fungal-algal associations (Gueidan et al., 2011; Prieto and Wedin, 2013; Lutzoni et al., 2018; Nelsen et al., 2019) and their multiple origins within the evolution of fungi (Gueidan et al., 2008; Schoch et al., 2009; Nelsen et al., 2020), lichens have colonised and diversified greatly in most terrestrial and some aquatic environments, including the most inhospitable niches (Kappen, 2000; Sadowsky and Ott, 2016). They are a discrete but inherent part of most of our landscapes, including both natural and man-made. This success stems from their ability to act as self-sustainable ecosystems, for which an evolutionary modularity (i.e., selection of the most fitted partners for a particular environment) has allowed adaptation to a broad range of habitats. Because of their slow growth, they particularly excel in colonising harsh habitats in which competition with faster growing micro-organisms is low. As such, they have adapted to surviving on nutrient-poor substrates and under drastically fluctuating environmental conditions, and play key roles in their ecosystems. In the future, lichen adaptations and their natural ecological flexibility may prove to be key to the successful responses of lichens to climate change.

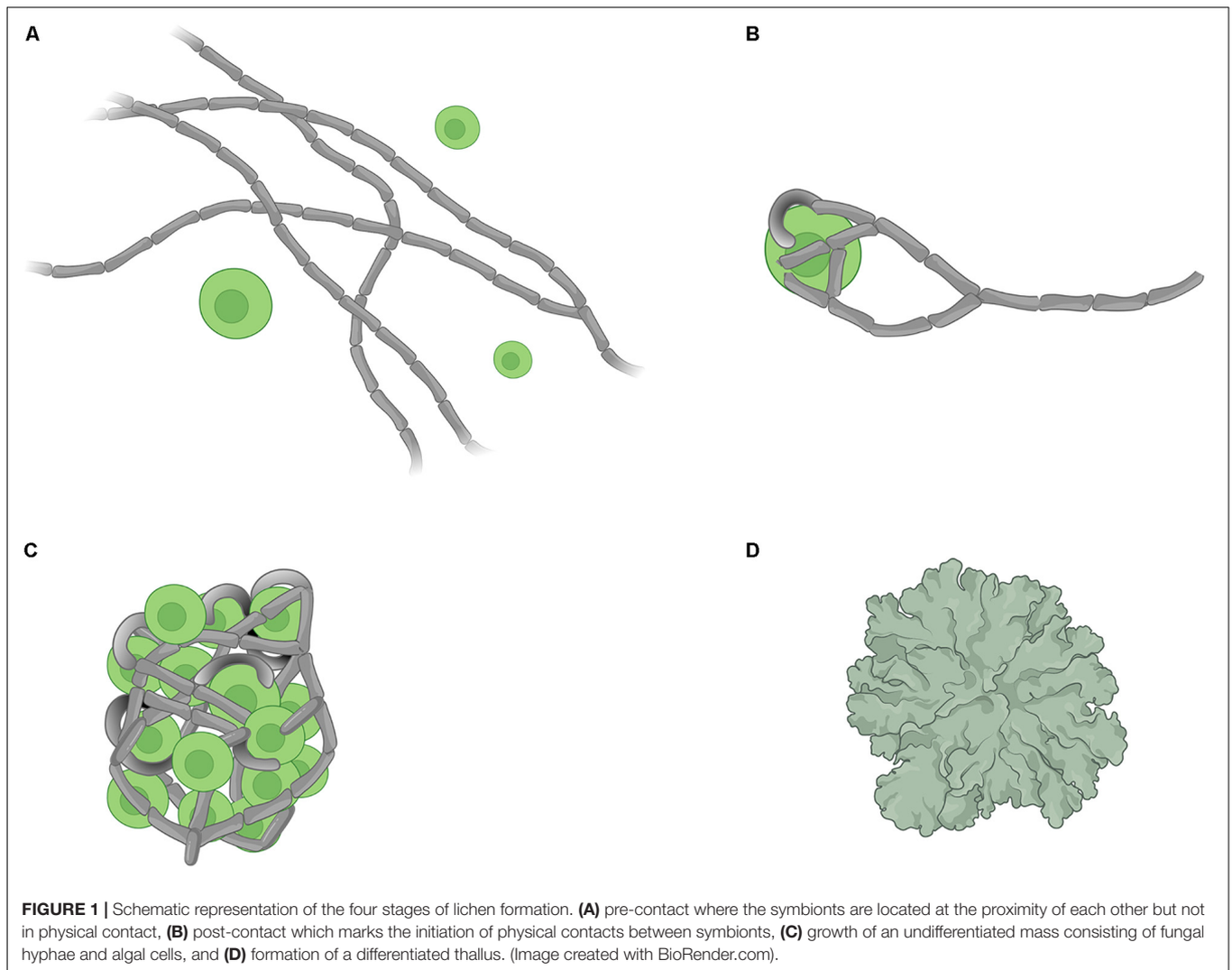
The lichen symbiosis is no longer perceived to be the simple union of a fungal partner (i.e., mycobiont) and a microalgal partner (i.e., photobiont), either an alga (i.e., chlorolichen) or a cyanobacterium (i.e., cyanolichen). Instead, previous studies have shown that lichens harbour a diverse microbiome (e.g., Petrini et al., 1990; Hofstetter et al., 2007; Grube et al., 2009; Hodgkinson and Lutzoni, 2009), and more recent studies corroborate lichens as multi-symbioses, i.e., complex multi-species associations including bacteria and other fungi or algae (Spribille et al., 2016; Onut-Brannstrom et al., 2018; Tuovinen et al., 2019; Smith et al., 2020; Leiva et al., 2021). In such symbioses, each partner contributes to the association: the primary mycobiont provides shelter and minerals to the photobiont, while the photobiont provides organic carbon fixed from atmospheric CO₂ via photosynthesis (Nash, 2008a) as well as nitrogen if it is a cyanobacteria. Additional bacteria, algae, and/or fungi have also been shown to serve certain functions in the lichen symbiosis (Cernava et al., 2017; Smith et al., 2020; Tagirdzhanova et al., 2021), although much more remains to be explored. Additionally, the levels of dependence and specificity of some of these microbes to the symbiosis are still debated (Grube et al., 2015; Kono et al., 2017; Jenkins and Richards, 2019; Lendemer et al., 2019; Smith et al., 2020). Lichens demonstrate unique physiological properties and ecosystem functions (Porada et al., 2014). All lichens contribute to atmospheric carbon fixation, with global net carbon uptake by both lichens and bryophytes predicted to be 0.34–3.3 Gt carbon per year (Palmqvist, 1995; Green et al., 2008; Palmqvist et al., 2008; Porada et al., 2013). Cyanolichens are capable of both carbon and nitrogen fixation (Dahlman et al., 2004; Nash, 2008b; Porada et al., 2017). Lichens grow on various substrates (including rocks,

trees, and soil), can survive extreme temperatures, tolerate desiccation (poikilohydric) and high levels of UV radiation, and form morphologically diverse structures (Beckett et al., 2008; Kranner et al., 2008). Many lichens produce unique specialised/secondary metabolites, including depsides, xanthenes and dibenzofurans, some of which have been shown to have medicinal properties (Fahselt, 1994; Elix and Stocker-Worgotter, 2008; Calcott et al., 2018).

The establishment of the lichen symbiosis, or “lichenisation,” has been described as a four-stage process (Ahmadjian et al., 1978): (A) a pre-contact phase (chemical interactions between symbionts but no physical contact), (B) a post-contact phase (with chemical and physical interactions), (C) a phase of growth characterised by an un-differentiated mass, and (D) a phase of differentiation that leads to a stratified thallus (Figure 1). Because mycobionts grow relatively slowly, the application of classical experimental microbiology techniques and co-culture/resynthesis experiments to the understanding of the development and functioning of the lichen symbiosis has lagged. Despite some recent studies focusing on early stages of lichenisation (Joneson et al., 2011; Armaleo et al., 2019; Kono et al., 2020), the molecular basis of fungal-algal interactions during lichenisation remains mostly uncharacterised, and processes involved in signalling and metabolic interplays between the symbionts are poorly understood. Contemporary systems biology approaches may facilitate tackling long-standing questions about the lichen symbiosis.

Systems biology is the study of living systems through the joint application of advanced high-data-volume generating technologies (e.g., ‘omics’) and computational tools (e.g., multi-scale or constraint-based modelling) to gain a more holistic understanding of the inter-dependencies of system components and underlying system complexity. Hypotheses are generally tested using iterative cycles of ‘wet’ (lab-based) and ‘dry’ (simulation-based) experiments, by which systems-level data are generated, analysed, and then used to inspire new insights and hypotheses about the biological system at hand (Kitano, 2002a,b). For instance, applying systems- and genome-level approaches to the legume-rhizobium symbiosis has greatly enhanced the knowledge on the underlying mechanisms of symbiotic interactions at molecular level, moving us one step closer to improving agricultural crop yields through the development of more efficient symbiotic N₂ fixation processes (diCenzo et al., 2019). A similar systems biology approach has not yet been applied to the study of the lichen symbiosis.

In this review, we summarise the current knowledgebase of signalling and recognition mechanisms in the lichen symbiosis. We then discuss the modelling of signalling networks as a tool to extend our understanding of such mechanisms in lichens. We review the literature on lichen metabolism and propose that modelling fluxes in metabolic networks could be a powerful tool for providing insights into lichen metabolism in particular, and the metabolic interplays between symbiotic partners in general. We provide a broad overview of metabolic network models and their applications in addition to a review of some of the symbiotic systems that have been studied through the lens of metabolic



models. Finally, the opportunities and challenges of modelling both signalling networks and metabolic fluxes are discussed.

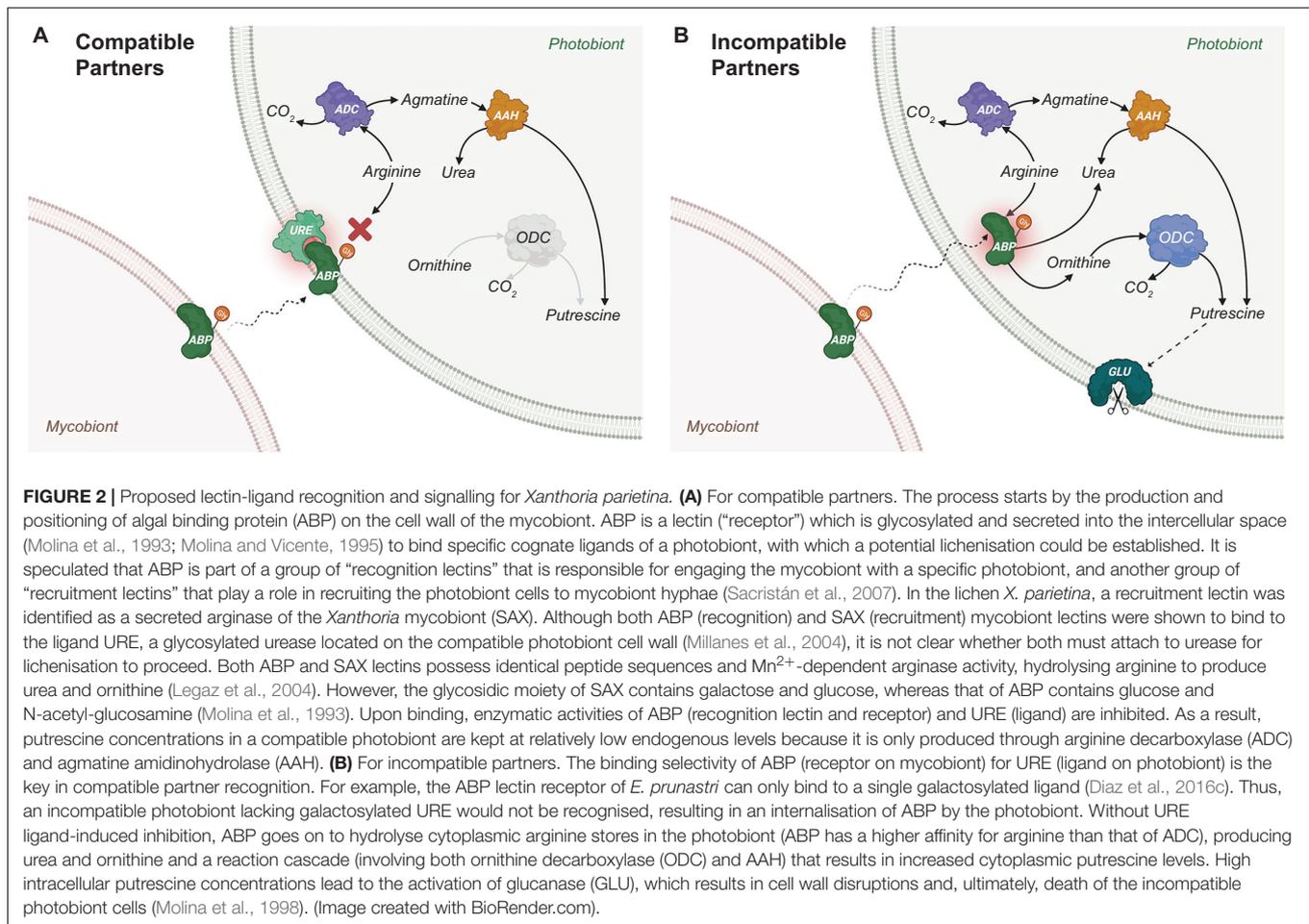
SIGNALLING AND RECOGNITION PATHWAYS IN THE LICHEN SYMBIOSIS

Distinct small molecules are produced by lichen symbionts during symbiosis that are absent when mycobiont and photobiont are grown separately (Green and Smith, 1974; Elshobary et al., 2015). Whether symbiont signalling and recognition processes in lichens are driven initially by those small molecules, or whether recognition processes are initiated by other regulatory mechanisms is not known. The available data for molecules with potential roles in signalling and/or recognition mechanisms during lichen symbiosis are summarised in **Table 1**. So far, there is no direct evidence confirming the production of compounds with a potential role in signalling and/or recognition during lichenisation by inhabiting fungi or bacteria. Several studies have shown that signalling between lichen symbionts can be initiated as early as the pre-contact stage of lichenisation

(Joneson et al., 2011; Meessen and Ott, 2013; Piercey-Normore and Athukorala, 2017; Armaleo et al., 2019). At present and for a few reasons, it is difficult to propose universal signalling models that initiate lichen symbiosis. Firstly, there is no single signalling molecule with a known or proposed role that has been studied across different lichens. Secondly, signalling pathways of those molecules with putative recognition roles have not been elucidated. Thirdly, lichens have likely evolved independently in several fungal lineages (Gueidan et al., 2008; Schoch et al., 2009), suggesting that the nature of these signalling pathways might differ depending on the species of interest. Nonetheless, owing to advances in genetic and analytical tools, several studies have begun to uncover mechanistic details underlying partner signalling and recognition at various stages of lichenisation (Meessen et al., 2013; Meessen and Ott, 2013; Athukorala et al., 2014; Athukorala and Piercey-Normore, 2015).

Lectin-Ligand Signalling in Lichens

Lectins are glycoproteins that occur ubiquitously across all domains of life (Kennedy et al., 1995). Lectins have also been isolated and characterised from both chlorolichens and



cyanolichens (Table 1). Possessing versatile carbohydrate-binding site(s), lectins can act as receptors and/or bind/agglutinate cells that may facilitate further interfacial communication between cells. The glycosidic moieties of lectins synthesised by the mycobiont may contain various combinations of carbohydrate groups that bind to specific ligands from the photobiont. In this atypical receptor-ligand system, lectins from mycobionts act as receptors for photobiont-attached ligands. A proposed mechanism of photobiont recognition and recruitment by the mycobiont is illustrated in Figure 2, based on the extensive studies of the lichens *Xanthoria parietina* and *Evernia prunastri* (Bubrick and Galun, 1980; Bubrick et al., 1981; Perezurria and Vicente, 1989; Vicente and Perezurria, 1989; Rodriguez and Vicente, 1991; Molina et al., 1993, 1998; Molina and Vicente, 1995, 2000; Legaz et al., 2004; Millanes et al., 2004). Lectins characterised from other lichens also have been proposed to have roles in the establishment and/or maintenance of compatible symbiotic relationships (Table 1).

In several lichen associations (including *X. parietina* and *E. prunastri* shown in Figure 2), the ligand for lectin receptors has been identified as urease, which is bound to the cell wall of the photobiont (Molina et al., 1993; Millanes et al., 2004; Diaz et al., 2009). In the lichen *Cladonia rangiferina*, a

urease-like recognition-related protein (RR1) was characterised and speculated to act as a ligand on the cell wall of the compatible photobiont of this lichen association (Athukorala et al., 2014; Athukorala and Piercey-Normore, 2015). Urease is produced by several lichens (presumably by the photobiont) and is secreted into the culture medium under laboratory conditions (Perezurria et al., 1989, 1993). The secretion of urease into the medium is hypothesised to be the consequence of its transfer from the photobiont to the mycobiont, depending on the nitrogen content of the mycobiont as well as the water content of the lichen thallus (Perezurria et al., 1989). However, it is not clear, whether the secreted ureases play a role similar to that of membrane-bound urease.

The lectin recognition and signalling mechanism summarised for chlorolichens in Figure 2 can be true of all or some cyanolichens (Sacristán et al., 2007; Vivas et al., 2010; Díaz et al., 2016a). Díaz et al. (2015), Díaz et al. (2016b) showed that actin- and myosin-like proteins produced by the cyanobacterial photobiont *Nostoc* of the lichen *Peltigera canina* is involved in the chemotactic movement of photobiont cells towards the lectin of the mycobiont. The process also involves a contractile protein and ATPase of photobiont, which creates a series of contraction-relaxation steps that result in photobiont

TABLE 1 | Molecules produced by different lichen symbionts with proposed roles in symbiotic signalling and recognition.

Molecule	Chemical class	Proposed role	Mycobiont	Photobiont°	References
Produced by the mycobiont					
Algal binding protein (ABP)	Glycoprotein	Plays a role in recognition of photobiont ligand	<i>Xanthoria parietina</i> ¹	<i>Trebouxia sp.?</i>	Molina et al., 1993; Molina and Vicente, 2000
Cyanobacterium-binding protein (CBP)	Possibly a glycoprotein	Plays a role in the first step of the recognition of compatible symbionts in a cyanolichen	<i>Peltigera canina</i> ²	<i>Nostoc sp.</i>	Diaz et al., 2009
Galectin LEC-1 and LEC-2	Glycan-binding proteins	Plays a role in recognition of photobiont ligand	<i>Scytinium palmatum</i> ⁴	<i>Nostoc sp.</i>	Vivas et al., 2010
			<i>Peltigera membranacea</i> ²	<i>Nostoc sp.</i>	Manoharan et al., 2012; Miao et al., 2012
<i>Nephroma laevigatum</i> agglutinin (NLA)	Possibly a glycoprotein	Functions as a determinant of specificity at the initial stage of symbiont interaction	<i>Nephroma laevigatum</i> ³	<i>Nostoc sp.</i>	Kardish et al., 1991
<i>Peltigera membranacea</i> agglutinin (PMA)	Glycoprotein	Functions in the recognition process between symbionts	<i>Peltigera membranacea</i> ²	<i>Nostoc sp.</i>	Lehr et al., 1995
Phytohemagglutinins	Glycoprotein	May be involved in the initial stages of the symbiosis establishment	<i>Peltigera canina</i> ²	<i>Nostoc sp.</i>	Lockhart et al., 1978
			<i>Peltigera polydactyla</i> ²	<i>Nostoc sp.</i>	Lockhart et al., 1978
Phytolectin	Glycoprotein	May be involved in the recognition or initial interactions between compatible lichen symbionts	<i>Peltigera horizontalis</i> ²	<i>Nostoc sp.</i>	Petit, 1982
			<i>Peltigera canina</i> var. <i>canina</i> ²	<i>Nostoc sp.</i>	Petit et al., 1983
Secreted arginase of <i>Evernia</i> (SAE)	Hydrolytic enzyme	Plays a role in recognition of photobiont ligand (e.g., urease)	<i>Evernia prunastri</i> ⁵	<i>Trebouxia excentrica</i>	Legaz et al., 2004
Secreted arginase of <i>Xanthoria</i> (SAX)			<i>Xanthoria parietina</i> ¹	<i>Trebouxia sp.?</i>	Molina et al., 1993; Molina and Vicente, 2000
			<i>Xanthoria parietina</i> ¹	<i>Pseudotrebouxia aggregata</i>	Legaz et al., 2004
<i>Xanthoria</i> -protein	Glycoprotein	May have role in initiation of lichen resynthesis and discriminate between photobionts	<i>Xanthoria parietina</i> ¹	<i>Trebouxia sp.</i>	Bubrick and Galun, 1980; Bubrick et al., 1981
			<i>Variospora aurantia</i> ¹	<i>Pseudotrebouxia sp.</i>	Bubrick and Galun, 1980
			<i>Flavoplaca citrina</i> ¹	<i>Pseudotrebouxia sp.</i>	Bubrick and Galun, 1980
Produced by the photobiont					
Chitinase	Hydrolytic enzyme	Regulates controlled parasitism between the symbionts	<i>Cladonia rangiferina</i> ⁶	<i>Asterochloris sp.</i>	Athukorala and Piercey-Normore, 2015
Cyclo-L-leucyl-L-tyrosyl (CLT)	Cyclic dipeptide*	Not known	<i>Romjularia lurida</i> ⁸	<i>Asterochloris sp.</i>	Meessen et al., 2013
Cyclo-L-tryptophyl-L-tryptophyl (CTT)	Cyclic dipeptide*	Promotes the germination rate of mycobiont <i>in vitro</i> after 30 days	<i>Gyalolechia bracteata</i> ¹	<i>Trebouxia sp.</i> , Cl.1, sbgr.1	Meessen et al., 2013
		Not known	<i>Gyalolechia fulgens</i> ¹	<i>Trebouxia sp.</i> , Cl.1, sbgr.1	Meessen et al., 2013
		Not known	<i>Thalloidima sedifolium</i> ⁷	<i>Trebouxia sp.</i> , Cl.1, sbgr.1	Meessen et al., 2013
		Not known	<i>Xanthoria elegans</i> ¹	<i>Trebouxia sp.</i>	Meessen et al., 2013
Indole-3-carbaldehyde (ICA)	Phytohormone precursor	Decreases the germination rate of mycobiont <i>in vitro</i>	<i>Gyalolechia bracteata</i> ¹	<i>Trebouxia sp.</i> , Cl.1, sbgr.1	Meessen et al., 2013
		Not known	<i>Gyalolechia fulgens</i> ¹	<i>Trebouxia sp.</i> , Cl.1, sbgr.1	Meessen et al., 2013
		Not known	<i>Thalloidima sedifolium</i> ⁷	<i>Trebouxia sp.</i> , Cl.1, sbgr.1	Meessen et al., 2013
Rhamnose	Deoxy sugar	Not known	<i>Xanthoria elegans</i> ¹	<i>Trebouxia sp.</i>	Meessen et al., 2013
		Decreases the germination rate of mycobiont <i>in vitro</i>	<i>Gyalolechia bracteata</i> ¹	<i>Trebouxia sp.</i> , Cl.1, sbgr.1	Meessen et al., 2013
Ribitol	Sugar alcohol	Acts as pre-/post-contact signal molecule	<i>Cladonia grayi</i> ⁶	<i>Asterochloris sp.</i>	Joneson et al., 2011
		Overcomes the growth arrest of the mycobiont and promotes mycelium growth [#]	<i>Gyalolechia bracteata</i> ¹	<i>Trebouxia sp.</i> , Cl.1, sbgr.1	Meessen et al., 2013

(Continued)

TABLE 1 | Continued

Molecule	Chemical class	Proposed role	Mycobiont	Photobiont ^o	References	
Urease	Hydrolytic enzyme	Serves as a ligand for different lichen lectins	<i>Xanthoria parietina</i> ¹	<i>Pseudotrebouxia aggregata</i>	Millanes et al., 2004	
			<i>Evernia prunastri</i> ¹	<i>Trebouxia excentrica</i>	Millanes et al., 2004	
Produced by the symbiosis as a whole						
(The experiment settings did not allow to attribute the compound to either the mycobiont or the photobiont)						
1-aminocyclopropane-l-carboxylic acid (ACC)	Phytohormone precursor	Affects differentiation and regulates interactions in the lichen thallus	<i>Cladonia rangiferina</i> ⁶	Unidentified	Ott et al., 2000	
			Not known	<i>Usnea longissima</i> ⁵	Unidentified	Ott et al., 2000
			Not known	<i>Parmelia saxatilis</i> ⁵	Unidentified	Ott et al., 2000
			Not known	<i>Usnea sphacelata</i> ⁵	Unidentified	Ott et al., 2000
			Not known	<i>Peltigera polydactyla</i> ²	Unidentified	Ott et al., 2000
			Not known	<i>Peltigera canina</i> ²	Unidentified	Ott et al., 2000
			Not known	<i>Nephroma resupinatum</i> ³	Unidentified	Ott et al., 2000
Abscisic acid (ABA)	Phytohormone	Affects differentiation and regulates interactions in the lichen thallus	<i>Scytinium palmatum</i> ⁴	<i>Nostoc sp.</i>	Vivas et al., 2010	
			<i>Cladonia rangiferina</i> ⁶	Unidentified	Ott et al., 2000	
			Not known	<i>Cladonia arbuscula</i> ⁶	Unidentified	Ott et al., 2000
Indole-3-acetic acid (IAA)	Phytohormone	Affects differentiation and regulates interactions in the lichen thallus	<i>Cetraria islandica</i> ⁵	Unidentified	Ott et al., 2000	
			<i>Cladonia rangiferina</i> ⁶	Unidentified	Ott et al., 2000	
			Not known	<i>Peltigera hymenina</i> ²	Unidentified	Ott et al., 2000
			Not known	<i>Cetraria islandica</i> ⁵	Unidentified	Ott et al., 2000
			Not known	<i>Cladonia arbuscula</i> ⁶	Unidentified	Ott et al., 2000
Not known	<i>Ramalina duriae</i> ⁷	<i>Trebouxia sp.</i>	Epstein et al., 1986			

^o*Trebouxia*, *Pseudotrebouxia*, and *Asterochloris* are eukaryotic algae (Chlorophyta, *Trebouxiophyceae*) and *Nostoc* is a prokaryotic alga (cyanobacteria).

Fungal lineages are as follows: ¹ Ascomycota, Lecanoromycetes, Teloschistales, Teloschistaceae; ² Ascomycota, Lecanoromycetes, Peltigerales, Peltigeraceae; ³ Ascomycota, Lecanoromycetes, Nephromataceae; ⁴ Ascomycota, Lecanoromycetes, Peltigerales, Collemataceae; ⁵ Ascomycota, Lecanoromycetes, Lecanorales, Parmeliaceae; ⁶ Ascomycota, Lecanoromycetes, Lecanorales, Cladoniaceae; ⁷ Ascomycota, Lecanoromycetes, Lecanorales, Ramalinaceae; ⁸ Ascomycota, Lecanoromycetes, Lecideales, Lecideaceae.

[#]Ribitol in these pre-contact experiments (mycobiont and photobiont separated by a membrane) was added in concentrations of 0.05, 0.8, and 2.0%w/v to the culture media (water agar and malt yeast agar). Ribitol was not identified as photobiont exudate in these experiments, as seen in other studies (Richardson et al., 1968).

^{*}Belong to the class of diketopiperazines (DKPs) with potential applications as antitumor, antiviral, antifungal, and antibacterial properties.

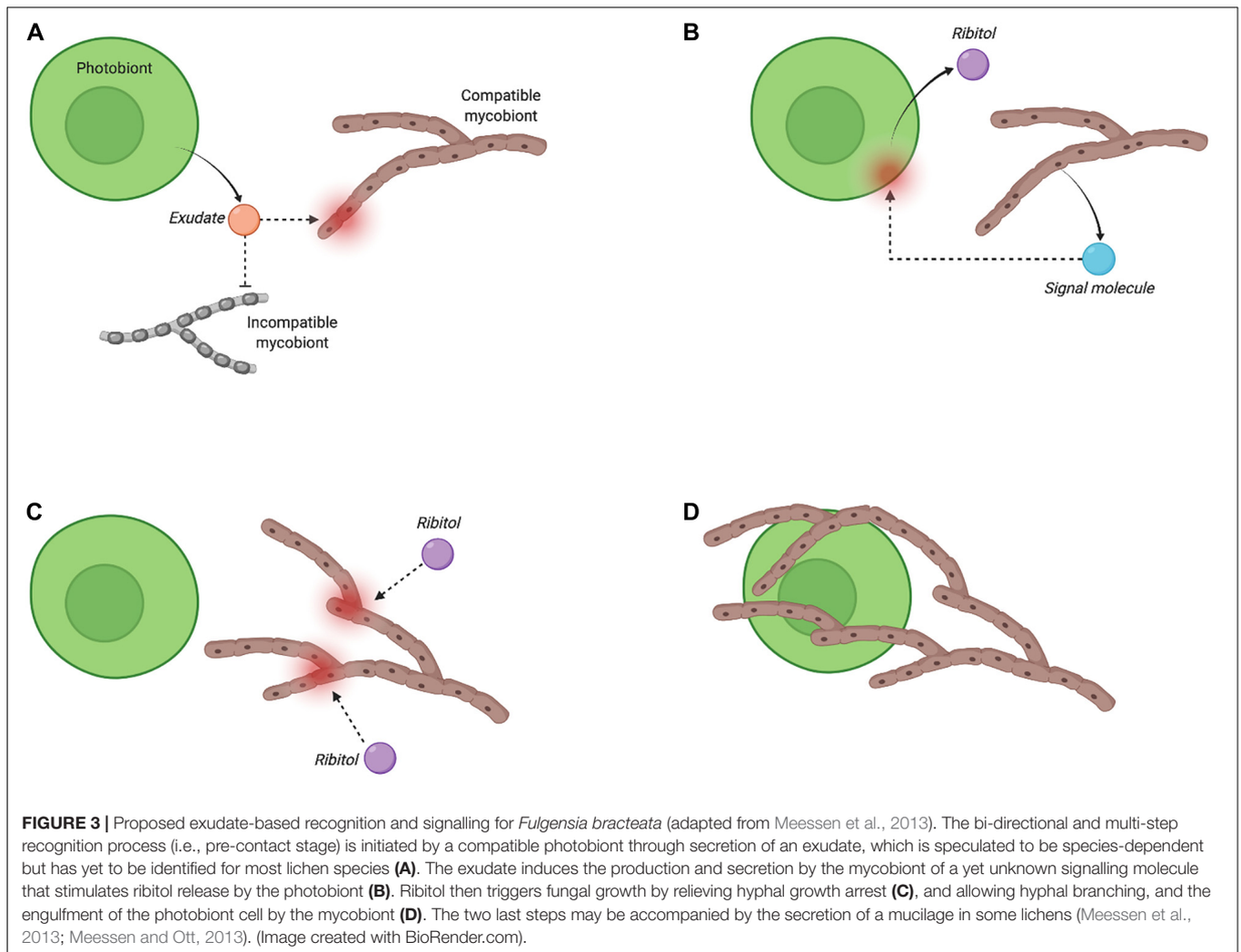
movement towards mycobiont lectin (Diaz et al., 2011). Upon cell contact of photobiont and mycobiont, desensitisation occurs and photobiont contractile motility stops. It is yet unknown if a similar type of chemotaxis applies to chlorolichens.

It is speculated that mycobionts of some lichens not only rely on lectin-ligand recognition mechanisms for establishing the initial photobiont contact, but that these mechanisms might also be involved in further replication and growth of young photobiont cells within the lichen thallus (Díaz et al., 2016a). The factors triggering the initiation of symbiont recognition through lectin-ligand binding and the ways in which symbionts increase the probability of association have been poorly understood, although several hypotheses have been proposed (Díaz et al., 2016a). For example, the photobiont could secrete a yet unknown diffusible compound that is sensed by a compatible mycobiont to trigger mycobiont lectin biosynthesis. Mycobionts may also produce multiple lectins with competing specificities for different photobionts, which may also be a strategy for rejecting incompatible photobionts. We could test some of these hypotheses using a systems biology approach, for example, through time-course

analysis of coupled gene expression and metabolome profiles of lichen co- and mono-cultures to identify candidate genes and molecules with potential signalling roles. Armaleo et al. (2019) recently pursued a transcriptome study exploring the differential expression of genes involved in symbiosis and signalling between *Cladonia grayi* and its algal partner *Asterochloris glomerata*. While only a snapshot in time, this work provided unprecedented insights into the complexity of responses underlying lichen symbioses.

Exudates Signalling in Lichens

Carbohydrate release and translocation from photobiont to mycobiont of a lichen was first proposed in the mid-1960s by Drew and Smith, who used radioactive isotope tracing to estimate the proportion of labelled carbon in sodium [¹⁴C]-bicarbonate fixed to [¹⁴C]-glucose by the cyanobacterial symbiont (*Nostoc*) of *Peltigera polydactyla* compared with its free-living and cultured forms (Drew and Smith, 1967a,b). Carbohydrate movement from photobiont to mycobiont has been investigated for more than 30 additional lichens and is reviewed elsewhere (Smith et al., 1969). The results of these studies showed that glucose and



sugar alcohols are the main forms of carbohydrates released by cyanobacterial and microalgal photobionts, respectively, and that they are translocated to the respective mycobionts. In the absence of a symbiotic relationship, the levels of carbohydrate released by the photobionts decline significantly or drop to zero. Following these initial studies, the importance of carbohydrate release by lichen photobionts gained a renewed interest in efforts to uncover the molecular mechanisms behind the early stage of lichenisation (Joneson et al., 2011; Meessen et al., 2013; Meessen and Ott, 2013; Athukorala et al., 2014; Athukorala and Piercey-Normore, 2014; Armaleo et al., 2019). A possible exudate signalling model based on the release and movement of ribitol is shown in Figure 3, and is largely based on independent studies observing ribitol release in the cultures of *Gyalolechia bracteata* (Meessen et al., 2013; Meessen and Ott, 2013) and *Cladonia grayi* (Joneson et al., 2011). Although the exact nature of the secreted molecules in this exudate-signalling model has not been fully elucidated, it is speculated that an exchange of carbon and nitrogen could be the driver for uniting symbionts in the first place. Hom and Murray (Hom and Murray, 2014) showed that co-culturing of model fungi

Saccharomyces cerevisiae, *Aspergillus nidulans*, or *Neurospora crassa* with the alga *Chlamydomonas reinhardtii* could facilitate mutualistic interactions through exchanging carbon and nitrogen under specific growth conditions; their results also suggest that carbon released by mycobiont respiration (as CO₂) could be recaptured by the photobiont for efficient carbon recycling within the lichen symbiosis (Schwartzman, 2010). Thus, the need for nutrient exchange between species could trigger the initiation of symbiotic interaction in lichens. Signalling network modelling, discussed in the following section, is one approach to generate insights on how specific exudate compounds could play a role in the overall flow of signals through the proposed ‘exudates signalling’ mechanism.

Signalling Network Modelling: Challenges and Opportunities for the Lichen Symbiosis

A signalling network consists of a series of ‘signals’ and ‘receptors’ whose relationships are determined by the signal transduction mechanisms governing the network. These signals and receptors

could be any or combination of enzymes (e.g., kinases), organic substances (e.g., ATP), inorganic molecules (e.g., phosphates), or other proteins or biomolecules. Reactions connecting these molecules frame the underlying signalling mechanisms and the goal of signalling network modelling would be to predict such interactions and the emergent cascade of signalling events that can explain or predict the behaviour of the signalling network. Signalling network models are often divided into descriptive and predictive subtypes. Descriptive models are usually simpler and provide a qualitative overview of the signalling pathway structure (i.e., topology of signal molecules and reactions), whereas predictive models may capture kinetics of the signalling pathway (i.e., reaction rates) and be capable of estimating system behaviours under new perturbations. The application of diverse descriptive and predictive modelling to signalling networks has been reviewed elsewhere (Hyduke and Palsson, 2010; Morris et al., 2010; Terfve and Saez-Rodriguez, 2012; Rother et al., 2013; Lavrik and Samsonova, 2016; Antebi et al., 2017). The scope and choice of signalling network modelling approach vary with the complexity of the network being explored. For example, some of the most detailed and comprehensive predictive signalling models have been developed for complex but known signalling networks of human B-cells (Papin and Palsson, 2004), prostate cancer cells (Dasika et al., 2006; Vardi et al., 2012), and Toll-like receptors (TLRs) functioning in immune system (Li et al., 2009).

In symbiotic systems, signalling pathways have been a topic of particular focus for legumes-rhizobia and plants-root fungi (mycorrhiza) symbioses (Bonfante and Genre, 2010; Bonfante and Requena, 2011; Oldroyd, 2013; Venkateshwaran et al., 2013; Mohanta and Bae, 2015; Martin et al., 2017; Poole et al., 2018; Clear and Hom, 2019). However, modelling the signalling networks in these systems has not received much attention, perhaps due largely to the knowledge gap in certain key signalling steps. For example, in the common symbiotic signalling “SYM” pathway, which shares similar signalling steps between arbuscular mycorrhizal and rhizobial symbioses, it remains unclear how symbiosis receptor kinases (SYMRK) transmit signals to downstream cation channelling proteins (i.e., CASTOR/POLLUX) located in the nucleus (Huisman and Geurts, 2020). Moreover, the precise mechanisms for how plants discriminate between arbuscular mycorrhiza and rhizobia symbionts are still unknown, although signalling pathways functioning in parallel to the SYM seem likely to be involved. Modelling signalling networks could represent a complementary approach to fill such gaps by simulating system behaviours with proposed/candidate mechanisms implemented by which symbionts transduce signals and communicate.

Faced with the paucity of detailed mechanistic knowledge on signalling networks in lichens (despite several potential signal molecules identified; see **Table 1**), the modelling of signalling networks in lichens suffers from similar challenges as those of other symbiotic systems and no models have yet been reported. Nevertheless, given the recent availability of ‘omics’ data for a variety of lichens (Mittermeier et al., 2015; Wang et al., 2015; Armaleo et al., 2019), there are now new opportunities to

develop signalling models of lichens. For instance, a proteomics approach could enable measuring lectin and urease levels of lichen cultures at pre- and post-contact stages informing the relative abundances of these proteins. The proteome profile of such cultures could also indicate the presence/absence of other specific proteins at the corresponding stages of lichenisation that may correlate with lectin/urease activity levels and provide deeper insights into how the recognition process initiates. A signalling pathway model could be developed to explore the link between putrescine biosynthesis and lectin production in repression of cell wall disruption of compatible photobiont as described in **Figure 3**.

METABOLIC INTERPLAY IN THE LICHEN SYMBIOSIS

The literature on lichen metabolism has been largely focused on understanding the exchange of key nutrients between symbionts (Lines et al., 1989; Kono et al., 2020; ten Veldhuis et al., 2020) and identifying lichen secondary metabolites and their biosynthetic pathways (i.e., metabolite profiling) (Fahsel, 1994; Aubert et al., 2007; Elix and Stocker-Worgotter, 2008; Mittermeier et al., 2015; Bertrand et al., 2018b; Brakni et al., 2018; Calcott et al., 2018; Kuhn et al., 2019; Goga et al., 2020; **Figure 4**). In the 1960s, observations of carbohydrate storage and translocation between the symbionts of *Peltigera polydactyla* (Smith and Drew, 1965; Drew and Smith, 1967a,b) together with a series of similar studies on other lichens (Smith et al., 1969) established the foundations for studying the metabolic interplay in lichens. The primary aim of those studies was to identify the form of carbon translocated between lichen symbionts, as explained in the previous sections. Next to nothing is known about the metabolic program and gene expression in lichen symbionts following carbohydrate exchange and assimilation. Most metabolic studies in lichens have concentrated on understanding the overall carbon and nitrogen economy in lichens, mainly with respect to overall carbon fixation, carbon sinks, lichen growth, and nitrogen fixation by cyanolichens (Honegger et al., 1993; Dahlman et al., 2004; Nash, 2008b; Palmqvist et al., 2008). Eisenreich and colleagues (Eisenreich et al., 2011) suggested that using ‘omics’ methods together with isotope labelling experiments (increasingly referred to as ‘fluxomics’) could enhance our understanding of lichen metabolic pathways, although this has yet to be fully realised to study lichen metabolism at a systems-level.

Thus, despite of being broadly recognised that carbohydrates and inorganic molecules are exchanged between lichen symbionts, a systems-level molecular understanding of metabolism is still lacking for lichens, including their primary symbionts and auxiliary partners. This lack has left key features of metabolism unexplored, including, for example, central aspects of carbon metabolism with respect to lichen compartmentalisation or the role of cell wall components and biosynthesis on the growth and metabolite exchange between symbionts. A systems-level understanding of lichen metabolism will become more likely in near future in light of the recent insights on lichen microbiota composition and functions within

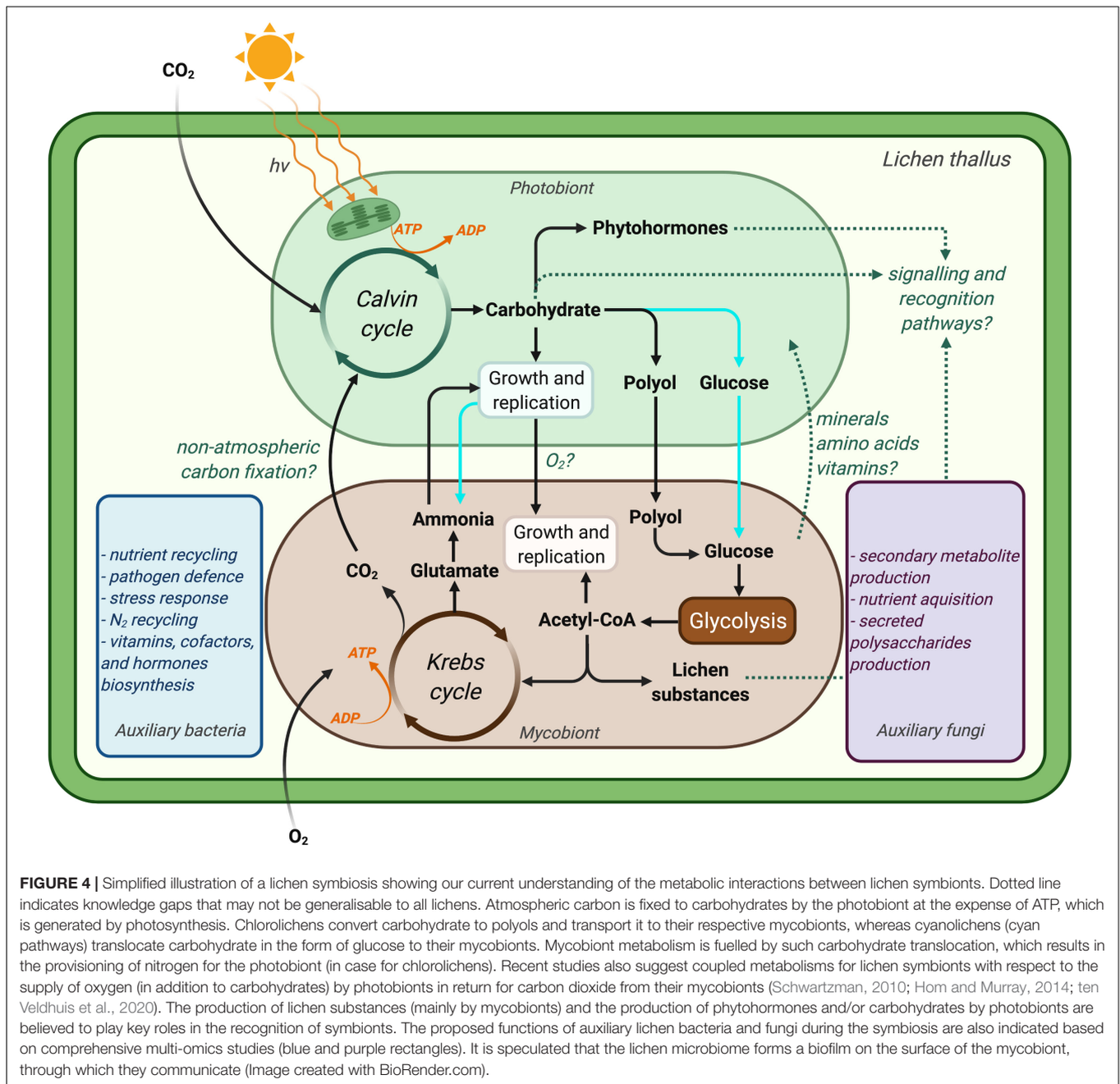


FIGURE 4 | Simplified illustration of a lichen symbiosis showing our current understanding of the metabolic interactions between lichen symbionts. Dotted line indicates knowledge gaps that may not be generalisable to all lichens. Atmospheric carbon is fixed to carbohydrates by the photobiont at the expense of ATP, which is generated by photosynthesis. Chlorolichens convert carbohydrate to polyols and transport it to their respective mycobionts, whereas cyanolichens (cyan pathways) translocate carbohydrate in the form of glucose to their mycobionts. Mycobiont metabolism is fuelled by such carbohydrate translocation, which results in the provisioning of nitrogen for the photobiont (in case for chlorolichens). Recent studies also suggest coupled metabolisms for lichen symbionts with respect to the supply of oxygen (in addition to carbohydrates) by photobionts in return for carbon dioxide from their mycobionts (Schwartzman, 2010; Hom and Murray, 2014; ten Veldhuis et al., 2020). The production of lichen substances (mainly by mycobionts) and the production of phytohormones and/or carbohydrates by photobionts are believed to play key roles in the recognition of symbionts. The proposed functions of auxiliary lichen bacteria and fungi during the symbiosis are also indicated based on comprehensive multi-omics studies (blue and purple rectangles). It is speculated that the lichen microbiome forms a biofilm on the surface of the mycobiont, through which they communicate (Image created with BioRender.com).

the lichen symbiosis (Spribille et al., 2016; Cernava et al., 2017; Smith et al., 2020).

Rhizobiales have been found to be a dominant bacterial order in the microbiome of various terrestrial or marine lichens examined to date (Grube et al., 2009; Hodkinson and Lutzoni, 2009; Hodkinson et al., 2012; Erlacher et al., 2015). Specifically, Rhodospirillales were found to be common in chlorolichens, and Sphingomonadales and Bacteroidetes in cyanolichens (Hodkinson et al., 2012; Graham et al., 2018; West et al., 2018). Several factors are believed to influence lichen-associated bacterial community composition and diversity. These include the nature of lichen secondary metabolites (driven mainly by the type of primary mycobiont), large-scale

geography, growth type, and the type of primary lichen photobiont (Grube et al., 2009; Hodkinson et al., 2012; Aschenbrenner et al., 2016). Some of these auxiliary bacteria were thought to be able to fix atmospheric nitrogen and, as cyanobacterial photobionts, might play an important role as a nitrogen source for the lichen symbiosis (Hodkinson and Lutzoni, 2009). Additionally, cyanobacterial lichens, which often grow in nitrogen-limited environments, were shown to harbour a diversity of bacteria that would otherwise not grow in such nitrogen-limited environments (Hodkinson et al., 2012). Apart from nitrogen fixation, meta-omics (e.g., meta-genomics, meta-transcriptomics, and meta-proteomics) studies have revealed functional roles for the microbiome of

the lichen *Lobaria pulmonaria*, including: nutrient recycling in the decaying parts of the lichen thallus, pathogen defence, detoxification processes, protection against oxidative stress, biosynthesis of vitamins, cofactors, and hormones, activation of ketone metabolism during dehydration, and upregulated transcription of transport systems, tRNA modification and various porins during hydration (Cernava et al., 2015; Grube et al., 2015; Aschenbrenner et al., 2016; Sigurbjornsdottir et al., 2016; Cernava et al., 2017; Cernava et al., 2019). The role of these auxiliary bacteria is thus critical to the maintenance and functioning of the lichen symbiosis.

The large diversity of lichen-associated fungi has been revealed through culture-dependent methods first (Petrini et al., 1990; Arnold et al., 2009), then meta-omics data analyses (Spribille et al., 2016; Smith et al., 2020). The low biomass of these auxiliary fungi relative to the primary mycobiont and the inability to culture them have prevented a detailed analysis of their functional roles in the lichen symbiosis. However, based on the analysis of meta-genome of the lichen *Alectoria sarmentosa*, a recent study showed that auxiliary fungi (two basidiomycete yeasts) may play roles in producing secreted extracellular polysaccharides, lichen nutrient acquisition, and secondary metabolite production (Tagirdzhanova et al., 2021). They are therefore also likely to play an important role in the maintenance and functioning of the lichen symbiosis.

Although meta-omics analyses of lichen microbiomes have provided invaluable insights on the diversity and function of multi-species lichen symbioses, constraint-based metabolic modelling could potentially enable a deeper understanding of the multi-species metabolic interplay. For example, by applying a systems biology approach using genome-scale metabolic reconstructions for 773 human gut bacteria (AGORA), a more sophisticated understanding of the interactions between the host and gut microbiome was achieved, revealing how system responses depended upon the metabolic potential of each component species and the nutrients available (Magnusdottir et al., 2017). The AGORA framework confirmed that a high fibre diet (usually linked to a healthy microbiome) would result in higher proportion of commensal and mutualistic pair-wise interactions between gut microbes. This framework was able to show how the host-microbiome operates mechanistically and indicate how many positive interactions are sufficient to maintain a healthy gut community. A similar systems-level understanding of lichens could help in understanding the metabolic interdependency for symbiotic establishment and maintenance, and in predicting the role of associated lichen microbes and lichen responses to environmental changes or likely environmental niches. This would also aid in re-creating/re-synthesizing lichens *in vitro* and using them for biotechnological applications.

Genome-Scale Metabolic Flux Modelling: Challenges and Opportunities for the Lichen Symbiosis

Genome-scale metabolic network models simulate the metabolism of a living cell as a collection of hundreds to

thousands of biochemical reactions (forming metabolic pathways of an organism) and enable quantitative and gene-grounded predictions of phenotypes under different growth conditions (Varma and Palsson, 1994; Covert et al., 2001). This set of reactions is framed as a set of ordinary differential equations, in which the number of variables and equations are defined by the number of metabolites and reactions, respectively. Solving this system of equations under a given set of assumptions (e.g., net zero system flux or “flux balance”) allows for determining optimal fluxes for each reaction in the metabolic network. Specific constraints describing the physico-chemical, environmental, regulatory, and/or topological conditions of the metabolic network can be imposed to identify optimal flux distributions consistent with these assumptions (Price et al., 2004). Such constraint-based metabolic modelling enables a wide range of applications including, but not limited to, predicting cellular functions (e.g., energy production) (Edwards et al., 2001; Orth and Palsson, 2012; Bordbar et al., 2014), identifying optimal strains and culture media conditions for specific applications (Pharkya et al., 2004; Nazem-Bokaei and Senger, 2015), formulating metabolic/strain engineering strategies (Burgard et al., 2003; Chung et al., 2010; Kim and Reed, 2010; Ranganathan et al., 2010; Rocha et al., 2010; McAnulty et al., 2012; Yen et al., 2013; Kim et al., 2019), identifying drug targets (Kim et al., 2011, 2012; Angione, 2019; Gu et al., 2019), producing natural/non-natural chemicals and precursors (Yim et al., 2011; Ye et al., 2014; Nazem-Bokaei et al., 2016; Wei et al., 2017; Nazem-Bokaei and Maranas, 2018; Biz et al., 2019; Gu et al., 2019), creating knowledgebases of metabolic, genomic, and biodiversity information (Kumar et al., 2012; Pabinger et al., 2014; King et al., 2016; Nazem-Bokaei et al., 2017; Norsigian et al., 2020), and studying syntrophic/symbiotic communities (see below). **Table 2** lists select examples of two-species metabolic models that have been studied.

Techniques developed for the characterisation of metabolic interactions among members of microbial communities based on genome-scale metabolic modelling can be classified into two main groups: lumped (also called enzyme soup, mixed bag, or metagenome-scale modelling (Chan et al., 2017a)) and compartmentalised (Biggs et al., 2015; Henry et al., 2016; Zomorodi and Segre, 2016). The analysis of interactions in a microbial community can be performed under steady-state or dynamic conditions. While an extensive description of these techniques and their implementation can be found elsewhere (Biggs et al., 2015; Zomorodi and Segre, 2016; Chan et al., 2017a; Ang et al., 2018; Garcia-Jimenez et al., 2021) and is beyond the scope of this review, it is worth broadly covering the general aim of each technique. The lumped modelling approach seeks to find optimal conditions that benefits the whole community (e.g., mutualistic symbiosis) by neglecting boundaries between members of the community (Taffs et al., 2009; Henry et al., 2016). The compartmentalised modelling approach, on the other hand, retains boundaries between members while also allowing individual members to share a compartment and transfer metabolites. For example, the compartmentalised modelling approach enables considering a member-level objective towards achieving a community-level

TABLE 2 | Select two-species metabolic network models that have been constructed and analysed¹.

Partners/symbionts ²	Community modelling approach ³	Key outcomes of the study	References
<i>Desulfovibrio vulgaris</i> (r: 89) <i>Methanococcus maripaludi</i> (r: 82)	Compartmentalised; steady-state	This is the first study on modelling mutualistic interactions between a sulphate-reducing bacterium and a methanogen using a compartmentalised approach. Using relatively small metabolic networks of the two microbes, a syntrophic methanogenesis was simulated when <i>D. vulgaris</i> produced hydrogen, carbon dioxide, and acetate, which were utilised by the methanogen.	Stolyar et al., 2007
<i>Geobacter sulfurreducens</i> (c: 2, g: 588, r: 727) <i>Rhodoferrax ferrireducens</i> (c: 2, g: 744, r: 762)	Compartmentalised; dynamic	This work analysed the dynamics of growth between two bacteria competing for uranium bioremediation.	Zhuang et al., 2011
<i>Scheffersomyces stipites</i> (c: 3, g: 814, r: 1371) <i>Saccharomyces cerevisiae</i> (c: 8, g: 904, r: 1412)	Lumped; dynamic (s: 3588)	In this study a co-culture simulating lignocellulosic feed breakdown for biofuel production was analysed using metabolic models of <i>S. cerevisiae</i> converting hexose and <i>S. stipites</i> converting pentose part of the synthetic feed into ethanol.	Hanly and Henson, 2013
<i>Geobacter metallireducens</i> (c: 2, g: 987, r: 1284) <i>Geobacter sulfurreducens</i> (c: 2, g: 837, r: 1085)	Compartmentalised; steady-state (t: 36)	A multi-omics approach was used in this study to understand electron flow mechanisms between the two bacteria. Results suggested that while <i>G. metallireducens</i> could respond only to syntrophic changes at transcriptomic level, <i>G. sulfurreducens</i> responded at both transcriptomic and genomic levels.	Nagarajan et al., 2013
<i>Bifidobacterium adolescentis</i> (g: 452, r: 699) <i>Faecalibacterium prausnitzii</i> (g: 484, r: 713)	Compartmentalised; steady-state	This study demonstrated that through modelling only two representatives of human gut microbiome, <i>B. adolescentis</i> and <i>F. prausnitzii</i> , the growth of the latter is severely affected when acetate production by the first microbe became limited.	El-Semman et al., 2014
<i>Salmonella enterica</i> <i>Escherichia coli</i> K12 strain	Compartmentalised; dynamic	Community modelling confirmed growth of <i>E. coli</i> on lactose minimal media was feasible only in co-culture with <i>S. enterica</i> , which received acetate and produced methionine in return.	Harcombe et al., 2014
<i>Escherichia coli</i> K strain (c: 3, g: 1260, r: 2073) <i>Escherichia coli</i> L strain (c: 3, g: 1260, r: 2073)	Compartmentalised; dynamic (t: 2)	Auxotrophy was studied using two mutants of <i>E. coli</i> , in which one grew with leucine and produced lysine that was assimilated by the other strain.	Zhang and Reed, 2014
<i>Ketogulonigenium vulgare</i> (c: 3, g: 663, r: 2073) <i>Bacillus megaterium</i> (c: 3, g: 1055, r: 2073)	Compartmentalised; steady-state (t: 453)	In this study an artificial consortium was constructed to analyse the production of vitamin C and other metabolites (e.g., 2-keto-l-gulonic acid) during two-step fermentation process	Ye et al., 2014
<i>Leptospirillum ferriphilum</i> (r: 87) <i>Ferroplasma acidiphilum</i> (r: 71)	Compartmentalised; steady-state	In this work, a bacteria-archaea mixed culture was modelled to study bioleaching (oxidizing iron)	Merino et al., 2015
<i>Chlamydomonas reinhardtii</i> (c: 10, g: 1080, r: 2191) <i>Saccharomyces cerevisiae</i> (c: 8, g: 750, r: 1266)	Compartmentalised; dynamic (t: 2)	The goal of this study was to feed process models with metabolic models of algal-fungal co-culture for optimizing biodiesel production. The alga produced oxygen for the yeast and in return received carbon dioxide secreted by the yeast. This study is an example of creating artificial symbiosis through exchange of key metabolites between an alga and a fungus, which could lead to higher biodiesel production compared with single cultures of the alga.	Gomez et al., 2016
<i>Thermosynechococcus elongatus</i> BP-1 (g: 583, r: 917) <i>Meiothermus ruber</i> strain A (g: 729, r: 1163)	Lumped and compartmentalised; steady-state (s: 1707)	The lumped model showed highest overall consistency between predicted fluxes and measured gene expression data. However, this approach provided no information on the potential interactions between the two members of consortia. The gap-filled compartmentalised model provided the best performance among all models with respect to predicting key metabolites interacting between the two bacteria.	Henry et al., 2016
<i>Medicago truncatula</i> (c: 6, g: 3403, r: 2909) <i>Sinorhizobium meliloti</i>	Compartmentalised; steady-state (t: 20)	The community model predicted the preferred uptake of ammonia over nitrate when both present in excess. At dark and when ammonia is limiting, the model predictions were in favour of nitrate uptake. The symbiotic model predicted amino acid cycling which is shown to be essential for nitrogen fixation for some rhizobial strains.	Pfau et al., 2018
<i>Nitrosomonas europaea</i> (g: 578) <i>Nitrobacter winogradskyi</i> (g: 579)	Compartmentalised; dynamic (t: 25)	Aerobic co-culture of two model nitrifying bacteria was used to study the dynamics of nitrification in agricultural settings	Mellbye et al., 2018
<i>Phaeodactylum tricorutum</i> (c: 6, g: 1027, r: 4456) <i>Pseudoalteromonas haloplanktis</i> (c: 2, g: 721, r: 1322)	Lumped; dynamic (s: 3588)	This work demonstrates the advantages of using metabolic models to simulate a diatom-bacteria co-culture to study the effect of changes in growth parameters on the co-culture to represent ocean food ecosystem. Using a linear community-level biomass objective function, a multi-compartment model was built, and then, converted into a dynamic, constraint-based, model of co-culture. Simulating this synthetic ecosystem revealed that the growth of the diatom was negatively affected by the growth of the bacterium due to the shortage of phosphate and sulphate.	Fondi and Di Patti, 2019

¹ Community metabolic models developed to study interactions among more than two organisms in any microbiota was excluded in this table for simplicity. For further information on larger communities of microbes the reader is referred to the text and these reviews (Zomorodi and Segre, 2016; Ang et al., 2018; Chan et al., 2017a; Gu et al., 2019).

² Numbers in parenthesis indicate the number of compartments (c), genes (g), and reactions (r), if available, captured in the respective metabolic model of the symbiont.

³ Numbers in parenthesis indicate the number of inter-species transporters (t) or shared reactions (s), when available, captured in the respective community metabolic model.

objective by imposing a constant growth rate across all members for a community to ensure co-existence and stability (Chan et al., 2017b). Although computationally more expensive, the compartmentalised modelling approach also allows for the study of different types of species-species interactions (e.g., parasitism) (Zomorodi and Maranas, 2012). A dynamic modelling approach enables predictions of changes in metabolites and biomass over time within the community and relies on kinetic data of uptake reactions. The dynamic approach has been extended to enable spatial analysis of communities, as in the COMETS (Computation Of Microbial Ecosystems in Time and Space) framework, which coupled metabolic with diffusion modelling and was applied to understand metabolite exchange within a three-member microbial community (Harcombe et al., 2014).

To our knowledge, no genome-scale metabolic network model has yet been constructed for any lichen association or its symbionts. With the first genomes of mycobionts (Park et al., 2013a,b, 2014a,b; Armstrong et al., 2018; Bertrand et al., 2018a; Wang et al., 2018) and photobionts (Armaleo et al., 2019) of several lichens assembled and more foreseen to come, it is a timely opportunity to understand the lichen symbiosis through the lens of genome-scale metabolic models. Since little is known about the metabolic response of lichens to different environmental conditions (e.g., light intensity, water content, nutrient availability, etc.), developing a metabolic network model could shed invaluable insights on symbiosis at the molecular level. Furthermore, the available computational tools for modelling community interactions could allow for predicting the role of a specific symbiont on the performance of a lichen under a known environmental perturbation (e.g., nutrient limitation). A lichen metabolic model could be used as the framework for the integration of 'omics' data obtained for lichens to test multiple hypotheses including, for example, the regulatory effect of different carbohydrates on the growth and exchange of metabolites between lichen symbiont. Since *in vitro* lichen re-synthesis is still hampered by the complexity of the lichenisation process, metabolic modelling could highlight potential metabolites that may need to be exchanged between symbionts as well as the metabolic pathways that may lead to successful differentiation and growth. Moreover, metabolic modelling could be used to examine the potential for symbiosis between various combinations of mycobionts and photobionts, and provide insights into the evolution of the lichen symbiosis. Validating predictions of flux distribution by community metabolic models could be a challenge, due to multi-compartmental nature of lichen symbiosis and difficulties in measuring fluxes through each compartment *in vivo*. However, recent advances in the field of metabolic flux analysis now make it possible to resolve fluxes by carefully designing the isotope labels and tracing them across different compartments (Schwechheimer et al., 2018). Another practical challenge for the development of lichen metabolic models may pertain to the characterisation of the cellular composition of individual lichen symbionts. For example, many lichen mycobionts grow slowly, making it experimentally difficult to obtain sufficient cell mass needed to formulate a 'biomass' reaction in a metabolic model representing cellular growth. Moreover, due to the lack of

data specific to the metabolic pathways of lichens, the model curation process may be patchy, with irreconcilable gaps and network disconnects. However, metabolic models for lichens could be reconstructed by leveraging the ever-increasing number of high-quality metabolic models becoming available for not-too-distantly related filamentous fungi, microalgae, or cyanobacteria (Brandl and Andersen, 2015; Gomez et al., 2016; Santos-Merino et al., 2019).

CONCLUSION AND FUTURE PERSPECTIVES

Lichens, although historically well-known and iconic symbioses, still bear a sense of mystery as our understanding of the signalling networks and pathways responsible for their symbiotic establishment and maintenance is still in its infancy. Two signalling mechanisms were reviewed in this article but many more could be explored with the aid of techniques such as untargeted metabolomics. Signalling/metabolic network modelling approaches could support the field of experimental lichenology by providing insights into: (1) the signalling molecules and the roles they play at different stages of lichenisation, (2) how lichen symbionts benefit from the symbiosis with regards to carbon, nitrogen, and other limiting nutrients or environmental conditions, (3) which conditions allow lichens to produce secondary metabolites and the genes that are involved, and (4) how lichens manage to accumulate and tolerate high levels of toxic metals. Advances in DNA sequencing technologies in recent years have significantly reduced the cost of generating genome sequences. At the same time, improvements in high performance computing and development of more biologist-friendly tools for modelling and analysing 'genome-scale' metabolic networks have enabled the exploration of metabolically-coupled microbial communities. Combining these genome resources and systems biology tools could open up a whole new era for the study of the lichen symbiosis.

AUTHOR CONTRIBUTIONS

HN-B and CG conceptualised and wrote the manuscript. HN-B and CG designed and created figures. EFYH, ACW, and SM revised the manuscript. All authors read and approved the final manuscript.

FUNDING

This work was supported by funding from the CSIRO Synthetic Biology Future Science Platform (FSP), Grant # OD-206013. EFYH was funded in part by NSF grants #1541538 and #1846376.

ACKNOWLEDGMENTS

We would like to thank Colin Scott (CSIRO) as well as the reviewers for critical reading and suggesting constructive improvements to the manuscript.

REFERENCES

- Ahmadjian, V. (1993). *The Lichen Symbiosis*. Hoboken, NJ: John Wiley & Sons.
- Ahmadjian, V., Jacobs, J. B., and Russell, L. A. (1978). Scanning electron-microscope study of early lichen synthesis. *Science* 200, 1062–1064. doi: 10.1126/science.200.4345.1062
- Ang, K. S., Lakshmanan, M., Lee, N. R., and Lee, D. Y. (2018). Metabolic modeling of microbial community interactions for health, environmental and biotechnological applications. *Curr. Genom.* 19, 712–722. doi: 10.2174/1389202919666180911144055
- Angione, C. (2019). Human systems biology and metabolic modelling: a review from disease metabolism to precision medicine. *Biomed. Res. Intern.* 2019:8304260.
- Antebi, Y. E., Nandagopal, N., and Elowitz, M. B. (2017). An operational view of intercellular signaling pathways. *Curr. Opin. Syst. Biol.* 1, 16–24. doi: 10.1016/j.coisb.2016.12.003
- Armaleo, D., Muller, O., Lutzoni, F., Andresson, O. S., Blanc, G., Bode, H. B., et al. (2019). The lichen symbiosis re-viewed through the genomes of *Cladonia grayi* and its algal partner *Asterochloris glomerata*. *BMC Genom.* 20:605. doi: 10.1186/s12864-019-5629-x
- Armstrong, E. E., Prost, S., Ertz, D., Westberg, M., Frisch, A., and Bendiksby, M. (2018). Draft genome sequence and annotation of the lichen-forming fungus *Arthonia radiata*. *Genome Announc.* 6:e0281-18.
- Arnold, A. E., Miadlikowska, J., Higgins, K. L., Sarvate, S. D., Gugger, P., Way, A., et al. (2009). A phylogenetic estimation of trophic transition networks for Ascomycetous fungi: are lichens cradles of symbiotrophic fungal diversification? *Syst. Biol.* 58, 283–297. doi: 10.1093/sysbio/syp001
- Aschenbrenner, I. A., Cernava, T., Berg, G., and Grube, M. (2016). Understanding microbial multi-species symbioses. *Front. Microbiol.* 7:180. doi: 10.3389/fmicb.2016.00180
- Athukorala, S. N. P., Huebner, E., and Piercey-Normore, M. D. (2014). Identification and comparison of the 3 early stages of resynthesis for the lichen *Cladonia rangiferina*. *Can. J. Microbiol.* 60, 41–52. doi: 10.1139/cjm-2013-0313
- Athukorala, S. N. P., and Piercey-Normore, M. D. (2014). Recognition between *Cladonia rangiferina* and the algal partner during resynthesis. *Bot. Botaniq.* 92, 640–640.
- Athukorala, S. N. P., and Piercey-Normore, M. D. (2015). Recognition- and defense-related gene expression at 3 resynthesis stages in lichen symbionts. *Can. J. Microbiol.* 61, 1–12. doi: 10.1139/cjm-2014-0470
- Aubert, S., Juge, C., Boisson, A. M., Gout, E., and Bligny, R. (2007). Metabolic processes sustaining the reviviscence of lichen *Xanthoria elegans* (Link) in high mountain environments. *Planta* 226, 1287–1297. doi: 10.1007/s00425-007-0563-6
- Beckett, R. P., Kranner, I., and Minibayeva, F. V. (2008). “Stress physiology and the symbiosis,” in *Lichen Biology*, 2nd Edn, ed. T. H. Nash (Cambridge: Cambridge University Press), 134–151. doi: 10.1017/cbo9780511790478.009
- Bertrand, R. L., Abdel-Hameed, M., and Sorensen, J. L. (2018a). Lichen biosynthetic gene clusters. Part I. Genome sequencing reveals a rich biosynthetic potential. *J. Nat. Prod.* 81, 723–731. doi: 10.1021/acs.jnatprod.7b00769
- Bertrand, R. L., Abdel-Hameed, M., and Sorensen, J. L. (2018b). Lichen biosynthetic gene clusters Part II: Homology mapping suggests a functional diversity. *J. Nat. Prod.* 81, 732–748. doi: 10.1021/acs.jnatprod.7b00770
- Biggs, M. B., Medlock, G. L., Kolling, G. L., and Papin, J. A. (2015). Metabolic network modeling of microbial communities. *Wiley Interdiscip. Rev. Syst. Biol. Med.* 7, 317–334. doi: 10.1002/wsbm.1308
- Biz, A., Proulx, S., Xu, Z., Siddhartha, K., Mulet Indrayanti, A., and Mahadevan, R. (2019). Systems biology based metabolic engineering for non-natural chemicals. *Biotechnol. Adv.* 37:107379. doi: 10.1016/j.biotechadv.2019.04.001
- Bonfante, P., and Genre, A. (2010). Mechanisms underlying beneficial plant-fungus interactions in mycorrhizal symbiosis. *Nat. Commun.* 1:48.
- Bonfante, P., and Requena, N. (2011). Dating in the dark: how roots respond to fungal signals to establish arbuscular mycorrhizal symbiosis. *Curr. Opin. Plant Biol.* 14, 451–457. doi: 10.1016/j.pbi.2011.03.014
- Bordbar, A., Monk, J. M., King, Z. A., and Palsson, B. O. (2014). Constraint-based models predict metabolic and associated cellular functions. *Nat. Rev. Genet.* 15, 107–120. doi: 10.1038/nrg3643
- Brakni, R., Ahmed, M. A., Burger, P., Schwing, A., Michel, G., Pomares, C., et al. (2018). UHPLC-HRMS/MS based profiling of algerian lichens and their antimicrobial activities. *Chem. Biodiver.* 15:e1800031. doi: 10.1002/cbdv.201800031
- Brandl, J., and Andersen, M. R. (2015). Current state of genome-scale modeling in filamentous fungi. *Biotechnol. Lett.* 37, 1131–1139. doi: 10.1007/s10529-015-1782-8
- Bubrick, P., and Galun, M. (1980). Proteins from the lichen *Xanthoria parietina* which bind to phycobiont cell walls correlation between binding patterns and cell wall Cyto Chemistry. *Protoplasma* 104, 167–173. doi: 10.1007/bf01279379
- Bubrick, P., Galun, M., and Frensdorff, A. (1981). Proteins from the lichen *Xanthoria parietina* which bind to phycobiont cell walls localization in the intact lichen and cultured mycobiont. *Protoplasma* 105, 207–211. doi: 10.1007/bf01279219
- Burgard, A. P., Pharkya, P., and Maranas, C. D. (2003). OptKnock: a bilevel programming framework for identifying gene knockout strategies for microbial strain optimization. *Biotechnol. Bioeng.* 84, 647–657. doi: 10.1002/bit.10803
- Calcott, M. J., Ackerley, D. F., Knight, A., Keyzers, R. A., and Owen, J. G. (2018). Secondary metabolism in the lichen symbiosis. *Chem. Soc. Rev.* 47, 1730–1760. doi: 10.1039/c7cs00431a
- Cernava, T., Aschenbrenner, I. A., Soh, J., Sensen, C. W., Grube, M., and Berg, G. (2019). Plasticity of a holobiont: desiccation induces fasting-like metabolism within the lichen microbiota. *ISME J.* 13, 547–556. doi: 10.1038/s41396-018-0286-7
- Cernava, T., Erlacher, A., Aschenbrenner, I. A., Krug, L., Lassek, C., Riedel, K., et al. (2017). Deciphering functional diversification within the lichen microbiota by meta-omics. *Microbiome* 5:82.
- Cernava, T., Muller, H., Aschenbrenner, I. A., Grube, M., and Berg, G. (2015). Analyzing the antagonistic potential of the lichen microbiome against pathogens by bridging metagenomic with culture studies. *Front. Microbiol.* 6:620. doi: 10.3389/fmicb.2015.00620
- Chan, S. H. J., Simons, M., and Maranas, C. D. (2017a). Computational modeling of microbial communities. *Syst. Biol.* 163–189. doi: 10.1002/9783527696130.ch6
- Chan, S. H. J., Simons, M. N., and Maranas, C. D. (2017b). SteadyCom: predicting microbial abundances while ensuring community stability. *PLoS Comput. Biol.* 13:e1005539. doi: 10.1371/journal.pcbi.1005539
- Chung, B. K., Selvarasu, S., Andrea, C., Ryu, J., Lee, H., Ahn, J., et al. (2010). Genome-scale metabolic reconstruction and in silico analysis of methylotrophic yeast *Pichia pastoris* for strain improvement. *Microb. Cell Fact.* 9:50. doi: 10.1186/1475-2859-9-50
- Clear, M. R., and Hom, E. F. (2019). The evolution of symbiotic plant–microbe signalling. *Annu. Plant Rev.* 2:684.
- Covert, M. W., Schilling, C. H., Famili, I., Edwards, J. S., Goryanin, I. I., Selkov, E., et al. (2001). Metabolic modeling of microbial strains in silico. *Trends Biochem. Sci.* 26, 179–186. doi: 10.1016/s0968-0004(00)01754-0
- Dahlman, L., Persson, J., Palmqvist, K., and Nasholm, T. (2004). Organic and inorganic nitrogen uptake in lichens. *Planta* 219, 459–467.
- Dasika, M. S., Burgard, A., and Maranas, C. D. (2006). A computational framework for the topological analysis and targeted disruption of signal transduction networks. *Biophys. J.* 91, 382–398. doi: 10.1529/biophysj.105.069724
- Díaz, E., Sánchez Elordi, E., Santiago, R., Vicente Córdoba, C., and Legaz González, M. E. (2016a). Algal-fungal mutualism: cell recognition and maintenance of the symbiotic status of lichens. *J. Vet. Med. Res.* 3:1052.
- Díaz, E. M., Ampe, C., Van Troys, M., Vicente-Manzanares, M., Legaz, M. E., and Vicente, C. (2016b). An actomyosin-like cytoskeleton in the cyanobiont (*Nostoc* sp.) of *Peltigera canina*. *Phytochem. Lett.* 16, 249–256. doi: 10.1016/j.phyto.2016.05.005
- Díaz, E. M., Cutrona, C., Sanchez-Elordi, E., Legaz, M. E., and Vicente, C. (2016c). Direct and cross-recognition of lichenized *Trebouxia pumalyi* (Chlorophyta, Trebouxiophyceae) and *Nostoc* Vaucher ex Bornet (Cyanobacteria, Cyanophyceae) by their homologous and heterologous fungal lectins. *Braz. J. Bot.* 39, 507–518. doi: 10.1007/s40415-016-0268-9
- Díaz, E. M., Sacristan, M., Legaz, M. E., and Vicente, C. (2009). Isolation and characterization of a cyanobacterium-binding protein and its cell wall receptor in the lichen *Peltigera canina*. *Plant Signal. Behav.* 4, 598–603. doi: 10.4161/psb.4.7.9164
- Díaz, E. M., Vicente-Manzanares, M., Legaz, M. E., and Vicente, C. (2015). A cyanobacterial beta-actin-like protein, responsible for lichenized *Nostoc* sp motility towards a fungal lectin. *Acta Physiol. Plant.* 37:249.
- Díaz, E. M., Vicente-Manzanares, M., Sacristan, M., Vicente, C., and Legaz, M. E. (2011). Fungal lectin of *Peltigera canina* induces chemotropism of compatible

- Nostoc* cells by constriction-relaxation pulses of cyanobiont cytoskeleton. *Plant Signal. Behav.* 6, 1525–1536. doi: 10.4161/psb.6.10.16687
- diCenzo, G. C., Zamani, M., Checucci, A., Fondi, M., Griffiths, J. S., Finan, T. M., et al. (2019). Multidisciplinary approaches for studying rhizobium-legume symbioses. *Can. J. Microbiol.* 65, 1–33. doi: 10.1139/cjm-2018-0377
- Drew, E. A., and Smith, D. C. (1967a). Studies in physiology of Lichens 7. Physiology of *Nostoc* Symbiont of *Peltigera polydactyla* compared with cultured and free-living forms. *New Phytol.* 66, 379–388. doi: 10.1111/j.1469-8137.1967.tb06017.x
- Drew, E. A., and Smith, D. C. (1967b). Studies in physiology of Lichens.8. Movement of glucose from alga to fungus during photosynthesis in thallus of *Peltigera polydactyla*. *New Phytol.* 66, 389–389. doi: 10.1111/j.1469-8137.1967.tb06018.x
- Edwards, J. S., Ibarra, R. U., and Palsson, B. O. (2001). In silico predictions of *Escherichia coli* metabolic capabilities are consistent with experimental data. *Nat. Biotechnol.* 19, 125–130.
- Eisenreich, W., Knispel, N., and Beck, A. (2011). Advanced methods for the study of the chemistry and the metabolism of lichens. *Phytochem. Rev.* 10, 445–456. doi: 10.1007/s11101-011-9215-3
- Elix, J. A., and Stocker-Worgotter, E. (2008). “Biochemistry and secondary metabolites,” in *Lichen Biology*, 2nd Edn, ed. T. H. Nash III (Cambridge: Cambridge University Press), 104–133. doi: 10.1017/cbo9780511790478.008
- El-Semman, I. E., Karlsson, F. H., Shoaie, S., Nookaew, I., Soliman, T. H., and Nielsen, J. (2014). Genome-scale metabolic reconstructions of *Bifidobacterium adolescentis* L2-32 and *Faecalibacterium prausnitzii* A2-165 and their interaction. *BMC Syst. Biol.* 8:41. doi: 10.1186/1752-0509-8-41
- Elshobary, M. E., Osman, M. E. H., Abushady, A. M., and Piercey-Normore, M. D. (2015). Comparison of lichen-forming cyanobacterial and green algal photobionts with free-living algae. *Cryptogam. Algal.* 36, 81–100. doi: 10.7872/crya.v36.iss1.2015.81
- Epstein, E., Sagee, O., Cohen, J. D., and Garty, J. (1986). Endogenous auxin and ethylene in the lichen *Ramalina duriaei*. *Plant Physiol.* 82, 1122–1125. doi: 10.1104/pp.82.4.1122
- Erlacher, A., Cernava, T., Cardinale, M., Soh, J., Sensen, C. W., Grube, M., et al. (2015). Rhizobiales as functional and endosymbiotic members in the lichen symbiosis of *Lobaria pulmonaria* L. *Front. Microbiol.* 6:53. doi: 10.3389/fmicb.2015.00053
- Fahselt, D. (1994). Secondary biochemistry of lichens. *Symbiosis* 16, 117–165.
- Fondi, M., and Di Patti, F. (2019). A synthetic ecosystem for the multi-level modelling of heterotroph-phototroph metabolic interactions. *Ecol. Model.* 399, 13–22. doi: 10.1016/j.ecolmodel.2019.02.012
- Garcia-Jimenez, B., Torres-Bacete, J., and Nogales, J. (2021). Metabolic modelling approaches for describing and engineering microbial communities. *Comput. Struct. Biotechnol. J.* 19, 226–246. doi: 10.1016/j.csbj.2020.12.003
- Goga, M., Elečko, J., Marciněinová, M., Ručová, D., Baėkorová, M., and Baėkor, M. (2020). “Lichen metabolites: an overview of some secondary metabolites and their biological potential,” in *Co-Evolution of Secondary Metabolites*, eds J. M. Merillon and K. Ramawat (Cham: Springer), 175–209. doi: 10.1007/978-3-030-16814-8_6
- Gomez, J. A., Hoffner, K., and Barton, P. I. (2016). From sugars to biodiesel using microalgae and yeast. *Green Chem.* 18, 461–475. doi: 10.1039/c5gc1843a
- Graham, L. E., Trest, M. T., Will-Wolf, S., Miicke, N. S., Atonio, L. M., Piotrowski, M. J., et al. (2018). Microscopic and metagenomic analyses of *Peltigera ponojensis* (Peltigerales, Ascomycota). *Intern. J. Plant Sci.* 179, 241–255. doi: 10.1086/696534
- Green, T. G. A., Nash, T. H., and Lange, O. L. (2008). “Physiological ecology of carbon dioxide exchange,” in *Lichen Biology*, 2nd Edn, eds T. G. A. Green, T. H. Nash, and O. L. Lange (Cambridge: Cambridge University Press), 152–181. doi: 10.1017/cbo9780511790478.010
- Green, T. G. A., and Smith, D. C. (1974). Lichen Physiology XIV. Differences between lichen algae in symbiosis and in isolation. *New Phytol.* 73, 753–766. doi: 10.1111/j.1469-8137.1974.tb01303.x
- Grube, M., Cardinale, M., De Castro, J. V., Muller, H., and Berg, G. (2009). Species-specific structural and functional diversity of bacterial communities in lichen symbioses. *ISME J.* 3, 1105–1115. doi: 10.1038/ismej.2009.63
- Grube, M., Cernava, T., Soh, J., Fuchs, S., Aschenbrenner, I., Lassek, C., et al. (2015). Exploring functional contexts of symbiotic sustain within lichen-associated bacteria by comparative omics. *ISME J.* 9, 412–424. doi: 10.1038/ismej.2014.138
- Gu, C., Kim, G. B., Kim, W. J., Kim, H. U., and Lee, S. Y. (2019). Current status and applications of genome-scale metabolic models. *Genome Biol.* 20:121.
- Guaidan, C., Ruibal, C., De Hoog, G. S., and Schneider, H. (2011). Rock-inhabiting fungi originated during periods of dry climate in the late devonian and middle triassic. *Fungal Biol.* 115, 987–996. doi: 10.1016/j.funbio.2011.04.002
- Guaidan, C., Villasenor, C. R., De Hoog, G. S., Gorbushina, A. A., Untereiner, W. A., and Lutzoni, F. (2008). A rock-inhabiting ancestor for mutualistic and pathogen-rich fungal lineages. *Stud. Mycol.* 61, 111–119. doi: 10.3114/sim.2008.61.11
- Hanly, T. J., and Henson, M. A. (2013). Dynamic metabolic modeling of a microaerobic yeast co-culture: predicting and optimizing ethanol production from glucose/xylose mixtures. *Biotechnol. Biofuels* 6:44. doi: 10.1186/1754-6834-6-44
- Harcombe, W. R., Riehl, W. J., Dukovski, I., Granger, B. R., Betts, A., Lang, A. H., et al. (2014). Metabolic resource allocation in individual microbes determines ecosystem interactions and spatial dynamics. *Cell Rep.* 7, 1104–1115. doi: 10.1016/j.celrep.2014.03.070
- Henry, C. S., Bernstein, H. C., Weisenhorn, P., Taylor, R. C., Lee, J. Y., Zucker, J., et al. (2016). Microbial community metabolic modeling: a community data-driven network reconstruction. *J. Cell. Physiol.* 231, 2339–2345. doi: 10.1002/jcp.25428
- Hodkinson, B. P., Gottel, N. R., Schadt, C. W., and Lutzoni, F. (2012). Photoautotrophic symbiont and geography are major factors affecting highly structured and diverse bacterial communities in the lichen microbiome. *Environ. Microbiol.* 14, 147–161. doi: 10.1111/j.1462-2920.2011.02560.x
- Hodkinson, B. P., and Lutzoni, F. (2009). A microbiotic survey of lichen-associated bacteria reveals a new lineage from the Rhizobiales. *Symbiosis* 49, 163–180. doi: 10.1007/s13199-009-0049-3
- Hofstetter, V., Miadlikowska, J., Kauff, F., and Lutzoni, F. (2007) Phylogenetic comparison of protein-coding versus ribosomal RNA-coding sequence data: A case study of the Lecanoromycetes (Ascomycota). *Mol. Phylogenet Evol.* 44, 412–426. doi: 10.1016/j.ympev.2006.10.016
- Hom, E. F. Y., and Murray, A. W. (2014). Niche engineering demonstrates a latent capacity for fungal-algal mutualism. *Science* 345, 94–98. doi: 10.1126/science.1253320
- Honegger, R. (1998). The Lichen symbiosis - What is so spectacular about it? *Lichenologist* 30, 193–212. doi: 10.1017/s002428299200015x
- Honegger, R., Kutasi, V., and Ruffner, H. P. (1993). Polyol patterns in 11 species of aposymbiotically cultured lichen mycobionts. *Mycol. Res.* 97, 35–39. doi: 10.1016/s0953-7562(09)81109-x
- Huisman, R., and Geurts, R. (2020). A roadmap toward engineered nitrogen-fixing nodule symbiosis. *Plant Commun.* 1:100019. doi: 10.1016/j.xplc.2019.100019
- Hyduke, D. R., and Palsson, B. O. (2010). Towards genome-scale signalling network reconstructions. *Nat. Rev. Genet.* 11, 297–307. doi: 10.1038/nrg2750
- Jenkins, B., and Richards, T. A. (2019). Symbiosis: wolf lichens harbour a choir of fungi. *Curr. Biol.* 29, R88–R90.
- Joneson, S., Armaleo, D., and Lutzoni, F. (2011). Fungal and algal gene expression in early developmental stages of lichen-symbiosis. *Mycologia* 103, 291–306. doi: 10.3852/10-064
- Kappen, L. (2000). Some aspects of the great success of lichens in Antarctica. *Antarct. Sci.* 12, 314–324. doi: 10.1017/s0954102000000377
- Kardish, N., Silberstein, L., Fleminger, G., and Galun, M. (1991). Lectin from the lichen *Nephroma laevigatum* Ach localization and function. *Symbiosis* 11, 47–62.
- Kennedy, J. F., Palva, P. M. G., Corella, M. T. S., Cavalcanti, M. S. M., and Coelho, L. C. B. B. (1995). Lectins, versatile proteins of recognition - a review. *Carbohydr. Polym.* 26, 219–230. doi: 10.1016/0144-8617(94)00091-7
- Kim, H. U., Kim, S. Y., Jeong, H., Kim, T. Y., Kim, J. J., Choy, H. E., et al. (2011). Integrative genome-scale metabolic analysis of *Vibrio vulnificus* for drug targeting and discovery. *Mol. Syst. Biol.* 7:460. doi: 10.1038/msb.2010.115
- Kim, H. U., Sohn, S. B., and Lee, S. Y. (2012). Metabolic network modeling and simulation for drug targeting and discovery. *Biotechnol. J.* 7, 330–342. doi: 10.1002/biot.201100159
- Kim, J., and Reed, J. L. (2010). OptORF: optimal metabolic and regulatory perturbations for metabolic engineering of microbial strains. *BMC Syst. Biol.* 4:53. doi: 10.1186/1752-0509-4-53

- Kim, M., Park, B. G., Kim, E. J., Kim, J., and Kim, B. G. (2019). In silico identification of metabolic engineering strategies for improved lipid production in *Vibrio vulnificus* by genome-scale metabolic modeling. *Biotechnol. Biofuels* 12:187.
- King, Z. A., Lu, J., Drager, A., Miller, P., Federowicz, S., Lerman, J. A., et al. (2016). BiGG Models: a platform for integrating, standardizing and sharing genome-scale models. *Nucleic Acids Res.* 44, D515–D522.
- Kitano, H. (2002a). Computational systems biology. *Nature* 420, 206–210.
- Kitano, H. (2002b). Systems biology: a brief overview. *Science* 295, 1662–1664. doi: 10.1126/science.1069492
- Kono, M., Kon, Y., Ohmura, Y., Satta, Y., and Terai, Y. (2020). In vitro resynthesis of lichenization reveals the genetic background of symbiosis-specific fungal-algal interaction in *Usnea hakonensis*. *BMC Genom.* 21:671. doi: 10.1186/s12864-020-07086-9
- Kono, M., Tanabe, H., Ohmura, Y., Satta, Y., and Terai, Y. (2017). Physical contact and carbon transfer between a lichen-forming *Trebouxia* alga and a novel *Alphaproteobacterium*. *Microbiol. SGM* 163, 678–691. doi: 10.1099/mic.0.000461
- Kranner, I., Beckett, R., Hochman, A., and Nash, T. H. (2008). Desiccation-tolerance in lichens: a review. *Bryologist* 111, 576–593. doi: 10.1639/0007-2745-111.4.576
- Kuhn, V., Geisberger, T., Huber, C., Beck, A., and Eisenreich, W. (2019). A facile in vivo procedure to analyze metabolic pathways in intact lichens. *New Phytol.* 224, 1657–1667. doi: 10.1111/nph.15968
- Kumar, A., Suthers, P. F., and Maranas, C. D. (2012). MetRxn: a knowledgebase of metabolites and reactions spanning metabolic models and databases. *BMC Bioinform.* 13:6. doi: 10.1186/1471-2105-13-
- Lavrik, I. N. S., and Samsonova, M. G. (2016). The systems biology of signaling pathways. *Biophysic* 61, 78–84.
- Legaz, M. E., Fontaniella, B., Millanes, A. M., and Vicente, C. (2004). Secreted arginases from phylogenetically far-related lichen species act as cross-recognition factors for two different algal cells. *Eur. J. Cell Biol.* 83, 435–446. doi: 10.1078/0171-9335-00384
- Lehr, H., Fleminger, G., and Galun, M. (1995). Lectin from the lichen *Peltigera membranacea* (Ach) Nyl - characterization and function. *Symbiosis* 18, 1–13.
- Leiva, D., Fernandez-Mendoza, F., Acevedo, J., Caru, M., Grube, M., and Orlando, J. (2021). The bacterial community of the foliose macro-lichen *Peltigera frigida* is more than a mere extension of the microbiota of the subjacent substrate. *Microb. Ecol.* doi: 10.1007/s00248-020-01662-y. Epub ahead of print.
- Lendemer, J. C., Keepers, K. G., Tripp, E. A., Pogoda, C. S., McCain, C. M., and Kane, N. C. (2019). A taxonomically broad metagenomic survey of 339 species spanning 57 families suggests cystobasidiomycete yeasts are not ubiquitous across all lichens. *Am. J. Bot.* 106, 1090–1095.
- Li, F., Thiele, I., Jamshidi, N., and Palsson, B. O. (2009). Identification of potential pathway mediation targets in toll-like receptor signaling. *PLoS Computat. Biol.* 5:e1000292. doi: 10.1371/journal.pcbi.1000292
- Lines, C. E. M., Ratcliffe, R. G., Rees, T. A. V., and Southon, T. E. (1989). A C-13 Nmr-study of photosynthate transport and metabolism in the lichen *Xanthoria calcicola* Oxner. *New Phytol.* 111, 447–456. doi: 10.1111/j.1469-8137.1989.tb00707.x
- Lockhart, C. M., Rowell, P., and Stewart, W. D. P. (1978). Phytohemagglutinins from nitrogen-fixing lichens *Peltigera canina* and *Peltigera polydactyla*. *FEMS Microbiol. Lett.* 3, 127–130.
- Lutzoni, F., Nowak, M. D., Alfaro, M. E., Reeb, V., Miadlikowska, J., Krug, M., et al. (2018). Contemporaneous radiations of fungi and plants linked to symbiosis. *Nat. Commun.* 9:5489.
- Magnusdottir, S., Heinken, A., Kutt, L., Ravcheev, D. A., Bauer, E., Noronha, A., et al. (2017). Generation of genome-scale metabolic reconstructions for 773 members of the human gut microbiota. *Nat. Biotechnol.* 35, 81–89. doi: 10.1038/nbt.3703
- Manoharan, S. S., Miao, V. P., and Andresson, O. S. (2012). LEC-2, a highly variable lectin in the lichen *Peltigera membranacea*. *Symbiosis* 58, 91–98. doi: 10.1007/s13199-012-0206-y
- Martin, F. M., Uroz, S., and Barker, D. G. (2017). Ancestral alliances: plant mutualistic symbioses with fungi and bacteria. *Science* 356:eaad4501. doi: 10.1126/science.aad4501
- McAnulty, M. J., Yen, J. Y., Freedman, B. G., and Senger, R. S. (2012). Genome-scale modeling using flux ratio constraints to enable metabolic engineering of clostridial metabolism in silico. *BMC Syst. Biol.* 6:42. doi: 10.1186/1752-0509-6-42
- Meessen, J., Eppenstein, S., and Ott, S. (2013). Recognition mechanisms during the pre-contact state of lichens: II. Influence of algal exudates and ribitol on the response of the mycobiont of *Fulgensia bracteata*. *Symbiosis* 59, 131–143. doi: 10.1007/s13199-012-0219-6
- Meessen, J., and Ott, S. (2013). Recognition mechanisms during the pre-contact state of lichens: I. Mycobiont-photobiont interactions of the mycobiont of *Fulgensia bracteata*. *Symbiosis* 59, 121–130. doi: 10.1007/s13199-013-0232-4
- Mellbye, B. L., Giguere, A. T., Murthy, G. S., Bottomley, P. J., Sayavedra-Soto, L. A., and Chaplen, F. W. R. (2018). Genome-scale, constraint-based modeling of nitrogen oxide fluxes during coculture of *Nitrosomonas europaea* and *Nitrobacter winogradskyi*. *mSystems* 3:e0170-17.
- Merino, M. P., Andrews, B. A., and Asenjo, J. A. (2015). Stoichiometric model and flux balance analysis for a mixed culture of *Leptospirillum ferriphilum* and *Ferroplasma acidiphilum*. *Biotechnol. Prog.* 31, 307–315. doi: 10.1002/btpr.2028
- Miao, V. P. W., Manoharan, S. S., Snaebjarnarson, V., and Andresson, O. S. (2012). Expression of lec-1, a mycobiont gene encoding a galectin-like protein in the lichen *Peltigera membranacea*. *Symbiosis* 57, 23–31. doi: 10.1007/s13199-012-0175-1
- Millanes, A. M., Fontaniella, B., Garcia, M. L., Solas, M. T., Vicente, C., and Legaz, M. E. (2004). Cytochemical location of urease in the cell wall of two different lichen phycobionts. *Tissue Cell* 36, 373–377. doi: 10.1016/j.tice.2004.06.007
- Mittermeier, V. K., Schmitt, N., Volk, L. P. M., Suarez, J. P., Beck, A., and Eisenreich, W. (2015). Metabolic profiling of alpine and ecuadorian lichens. *Molecules* 20, 18047–18065. doi: 10.3390/molecules201018047
- Mohanta, T. K., and Bae, H. (2015). Functional genomics and signaling events in mycorrhizal symbiosis. *J. Plant Interact.* 10, 21–40. doi: 10.1080/17429145.2015.1005180
- Molina, M. C., Stocker-Worgotter, E., Turk, R., Bajon, C., and Vicente, C. (1998). Secreted, glycosylated arginase from *Xanthoria parietina* thallus induces loss of cytoplasmic material from *Xanthoria* photobionts. *Cell Adhesion Commun.* 6, 481–490. doi: 10.3109/15419069809010796
- Molina, M. C., and Vicente, C. (1995). Correlations between enzymatic-activity of lectins, putrescine content and chloroplast damage in *Xanthoria parietina* phycobionts. *Cell Adhesion Commun.* 3, 1–12. doi: 10.3109/15419069509081274
- Molina, M. C., and Vicente, C. (2000). Purification and characterization of two isolectins with arginase activity from the lichen *Xanthoria parietina*. *J. Biochem. Mol. Biol.* 33, 300–307.
- Molina, M. D., Muniz, E., and Vicente, C. (1993). Enzymatic-activities of algal-binding protein and its algal cell-wall receptor in the lichen *Xanthoria parietina* - an approach to the parasitic basis of mutualism. *Plant Physiol. Biochem.* 31, 131–142.
- Morris, M. K., Saez-Rodriguez, J., Sorger, P. K., and Lauffenburger, D. A. (2010). Logic-based models for the analysis of cell signaling networks. *Biochemistry* 49, 3216–3224. doi: 10.1021/bi902202q
- Nagarajan, H., Embree, M., Rotaru, A. E., Shrestha, P. M., Feist, A. M., Palsson, B. O., et al. (2013). Characterization and modelling of interspecies electron transfer mechanisms and microbial community dynamics of a syntrophic association. *Nat. Commun.* 4:2809.
- Nash, T. H. (2008a). *Lichen Biology*. Cambridge: Cambridge University Press.
- Nash, T. H. (2008b). “Nitrogen, its metabolism and potential contribution to ecosystems” in *Lichen Biology*, 2nd Edn, ed. T. H. Nash (Cambridge: Cambridge University Press), 216–233. doi: 10.1017/cbo9780511790478.012
- Nazem-Bokaei, H., Gopalakrishnan, S., Ferry, J. G., Wood, T. K., and Maranas, C. D. (2016). Assessing methanotrophy and carbon fixation for biofuel production by *Methanosarcina acetivorans*. *Microb. Cell Fact.* 15:10.
- Nazem-Bokaei, H., and Maranas, C. D. (2018). A prospective study on the fermentation landscape of gaseous substrates to biorenewables using *Methanosarcina acetivorans* metabolic model. *Front. Microbiol.* 9:1855. doi: 10.3389/fmicb.2018.01855
- Nazem-Bokaei, H., and Senger, R. S. (2015). ToMI-FBA: a genome-scale metabolic flux based algorithm to select optimum hosts and media formulations for expressing pathways of interest. *Aims Bioeng.* 2, 335–374. doi: 10.3934/bioeng.2015.4.335

- Nazem-Bokaei, H., Yen, J. Y., Athamneh, A. I. M., Apte, A. A., Mcanulty, M. J., and Senger, R. S. (2017). SyM-GEM: a pathway builder for genome-scale models. *Adv. Biochem. Biotechnol.* 2:141. doi: 10.29011/2574-7258.000041
- Nelsen, M. P., Lucking, R., Boyce, C. K., Lumbsch, H. T., and Ree, R. H. (2019). No support for the emergence of lichens prior to the evolution of vascular plants. *Geobiology* 18, 3–13. doi: 10.1111/gbi.12369
- Nelsen, M. P., Lucking, R., Boyce, C. K., Lumbsch, H. T., and Ree, R. H. (2020). The macroevolutionary dynamics of symbiotic and phenotypic diversification in lichens. *Proc. Natl. Acad. Sci. U.S.A.* 117, 21495–21503. doi: 10.1073/pnas.2001913117
- Norsigian, C. J., Pusarla, N., Mcconn, J. L., Yurkovich, J. T., Drager, A., Palsson, B. O., et al. (2020). BiGG models 2020: multi-strain genome-scale models and expansion across the phylogenetic tree. *Nucleic Acids Res.* 48, D402–D406.
- Oldroyd, G. E. D. (2013). Speak, friend, and enter: signalling systems that promote beneficial symbiotic associations in plants. *Nat. Rev. Microbiol.* 11, 252–263. doi: 10.1038/nrmicro2990
- Onut-Brannstrom, I., Benjamin, M., Scofield, D. G., Heidmarsson, S., Andersson, M. G. I., Lindstrom, E. S., et al. (2018). Sharing of photobionts in sympatric populations of *Thamnomelia* and *Cetraria* lichens: evidence from high-throughput sequencing. *Sci. Rep.* 8:4406.
- Orth, J. D., and Palsson, B. (2012). Gap-filling analysis of the iJO1366 *Escherichia coli* metabolic network reconstruction for discovery of metabolic functions. *BMC Syst. Biol.* 6:30. doi: 10.1186/1752-0509-6-30
- Ott, S., Krieg, T., Spanier, U., and Schieleit, P. (2000). Phytohormones in lichens with emphasis on ethylene biosynthesis and functional aspects on lichen symbiosis. *Phyton Annale. Rei Bot.* 40, 83–94.
- Pabinger, S., Snajder, R., Hardiman, T., Willi, M., Dander, A., and Trajanoski, Z. (2014). MEMOSys 2.0: an update of the bioinformatics database for genome-scale models and genomic data. *Datab. J. Biol. Datab. Curat.* 2014:bau004.
- Palmqvist, K. (1995). Uptake and fixation of CO₂ in lichen photobionts. *Symbiosis* 18, 95–109.
- Palmqvist, K., Dahlman, L., Jonsson, A., and Nash, T. H. (2008). “The carbon economy of lichens,” in *Lichen Biology*, 2nd Edn, ed. T. H. Nash (Cambridge: Cambridge University Press), 182–215. doi: 10.1017/cbo9780511790478.011
- Papin, J. A., and Palsson, B. O. (2004). The JAK-STAT signaling network in the human B-cell: an extreme signaling pathway analysis. *Biophys. J.* 87, 37–46. doi: 10.1529/biophysj.103.029884
- Park, S. Y., Choi, J., Kim, J. A., Jeong, M. H., Kim, S., Lee, Y. H., et al. (2013a). Draft genome sequence of *Cladonia macilenta* KoLRI003786, a lichen-forming fungus producing biruloquinone. *Genome Announc.* 1:e0695-13.
- Park, S. Y., Choi, J., Kim, J. A., Yu, N. H., Kim, S., Kondratyuk, S. Y., et al. (2013b). Draft genome sequence of lichen-forming fungus *Caloplaca flavorubescens* Strain KoLRI002931. *Genome Announc.* 1:e0678-13.
- Park, S. Y., Choi, J., Lee, G. W., Kim, J. A., Oh, S. O., Jeong, M. H., et al. (2014a). Draft genome sequence of lichen-forming fungus *Cladonia metacoralifera* Strain KoLRI002260. *Genome Announc.* 2:e01065-13.
- Park, S. Y., Choi, J., Lee, G. W., Park, C. H., Kim, J. A., Oh, S. O., et al. (2014b). Draft genome sequence of *Endocarpon pusillum* strain KoLRL000583. *Genome Announc.* 2:e0452-14.
- Perezurria, E., Avalos, A., Guzman, G., and Vicente, C. (1993). Urease production and secretion by 3 antarctic lichen species. *Endocytobios. Cell Res.* 9, 239–243.
- Perezurria, E., and Vicente, C. (1989). Purification and some properties of a secreted urease from *Evernia prunastri* thallus. *J. Plant Physiol.* 133, 692–695. doi: 10.1016/s0176-1617(89)80074-4
- Perezurria, E., Vicente, C., and Filho, L. X. (1989). Screening of urease production and secretion by 7 species of finnish lichens. *Biochem. Syst. Ecol.* 17, 359–363. doi: 10.1016/0305-1978(89)90048-3
- Petit, P. (1982). Phytolectins from the nitrogen-fixing lichen *Peltigera horizontalis* - the binding pattern of primary-protein extract. *New Phytol.* 91, 705–710. doi: 10.1111/j.1469-8137.1982.tb03349.x
- Petit, P., Lallemand, R., and Savoye, D. (1983). Purified phytolectin from the lichen *Peltigera canina* var *canina* which binds to the phycobiont cell-walls and its use as cytochemical marker insitu. *New Phytol.* 94, 103–110. doi: 10.1111/j.1469-8137.1983.tb02726.x
- Petrini, O., Hake, U., and Dreyfuss, M. M. (1990). An analysis of fungal communities isolated from fruticose lichens. *Mycologia* 82, 444–451. doi: 10.2307/3760015
- Pfau, T., Christian, N., Masakapalli, S. K., Sweetlove, L. J., Poolman, M. G., and Ebenhoh, O. (2018). The intertwined metabolism during symbiotic nitrogen fixation elucidated by metabolic modelling. *Sci. Rep.* 8:12504.
- Pharkya, P., Burgard, A. P., and Maranas, C. D. (2004). OptStrain: a computational framework for redesign of microbial production systems. *Genome Res.* 14, 2367–2376. doi: 10.1101/gr.2872004
- Piercey-Normore, M. D., and Athukoral, S. N. P. (2017). Interface between fungi and green algae in lichen associations. *Botany* 95, 1005–1014. doi: 10.1139/cjb-2017-0037
- Poole, P., Ramachandran, V., and Terpolilli, J. (2018). Rhizobia: from saprophytes to endosymbionts. *Nat. Rev. Microbiol.* 16, 291–303. doi: 10.1038/nrmicro.2017.171
- Porada, P., Poschl, U., Kleidon, A., Beer, C., and Weber, B. (2017). Estimating global nitrous oxide emissions by lichens and bryophytes with a process-based productivity model. *Biogeosciences* 14, 1593–1602. doi: 10.5194/bg-14-1593-2017
- Porada, P., Weber, B., Elbert, W., Poschl, U., and Kleidon, A. (2013). Estimating global carbon uptake by lichens and bryophytes with a process-based model. *Biogeosciences* 10, 6989–7033. doi: 10.5194/bg-10-6989-2013
- Porada, P., Weber, B., Elbert, W., Poschl, U., and Kleidon, A. (2014). Estimating impacts of lichens and bryophytes on global biogeochemical cycles. *Glob. Biogeochem. Cycles* 28, 71–85. doi: 10.1002/2013gb004705
- Price, N. D., Reed, J. L., and Palsson, B. O. (2004). Genome-scale models of microbial cells: evaluating the consequences of constraints. *Nat. Rev. Microbiol.* 2, 886–897. doi: 10.1038/nrmicro1023
- Prieto, M., and Wedin, M. (2013). Dating the diversification of the major lineages of ascomycota (Fungi). *PLoS One* 8:e65576. doi: 10.1371/journal.pone.0065576
- Ranganathan, S., Suthers, P. F., and Maranas, C. D. (2010). OptForce: an optimization procedure for identifying all genetic manipulations leading to targeted overproductions. *PLoS Computat. Biol.* 6:e1000744. doi: 10.1371/journal.pcbi.1000744
- Richardson, D. H., Hill, D. J., and Smith, D. C. (1968). Lichen physiology XI. Role of alga in determining pattern of carbohydrate movement between lichen symbionts. *New Phytol.* 67, 469–486. doi: 10.1111/j.1469-8137.1968.tb05476.x
- Rocha, I., Maia, P., Evangelista, P., Vilaca, P., Soares, S., Pinto, J. P., et al. (2010). OptFlux: an open-source software platform for in silico metabolic engineering. *BMC Syst. Biol.* 4:45. doi: 10.1186/1752-0509-4-45
- Rodriguez, M., and Vicente, C. (1991). Water status and urease secretion from 2 ecotypes of *Xanthoria parietina*. *Symbiosis* 11, 255–262.
- Rother, M., Munzner, U., Thieme, S., and Krantz, M. (2013). Information content and scalability in signal transduction network reconstruction formats. *Mol. Biosyst.* 9, 1993–2004. doi: 10.1039/c3mb00005b
- Sacristán, M., Vivas, M., Millanes, A. M., Fontaniella, B., Vicente, C., and Legaz, M. E. (2007). “The recognition pattern of green algae by lichenized fungi can be extended to lichens containing a cyanobacterium as photobiont,” in *Communicating Current Research and Educational Topics and Trends in Applied Microbiology*, Vol. 1, ed. A. M. éndez-Vilas (Cham: Springer), 213–219.
- Sadowsky, A., and Ott, S. (2016). Symbiosis as a successful strategy in continental Antarctica: performance and protection of *Trebouxia* photosystem II in relation to lichen pigmentation. *Polar Biol.* 39, 139–151. doi: 10.1007/s00300-015-1677-0
- Santos-Merino, M., Singh, A. K., and Ducat, D. C. (2019). New applications of synthetic biology tools for cyanobacterial metabolic engineering. *Front. Bioeng. Biotechnol.* 7:33. doi: 10.3389/fbioe.2019.00033
- Schoch, C. L., Sung, G. H., Lopez-Giraldez, F., Townsend, J. P., Miadlikowska, J., Hofstetter, V., et al. (2009). The ascomycota tree of life: a phylum-wide phylogeny clarifies the origin and evolution of fundamental reproductive and ecological traits. *Syst. Biol.* 58, 224–239.
- Schwartzman, D. W. (2010). Was the origin of the lichen symbiosis triggered by declining atmospheric carbon dioxide levels? *Biol. Lichens Symbios.* 105, 191–196.
- Schwechheimer, S. K., Becker, J., and Wittmann, C. (2018). Towards better understanding of industrial cell factories: novel approaches for (13)C metabolic flux analysis in complex nutrient environments. *Curr. Opin. Biotechnol.* 54, 128–137. doi: 10.1016/j.copbio.2018.07.001
- Sigurbjornsdottir, M. A., Andresson, O. S., and Vilhelmsson, O. (2016). Nutrient scavenging activity and antagonistic factors of non-photobiont lichen-associated bacteria: a review. *World J. Microbiol. Biotechnol.* 32:68.

- Smith, D., Muscatine, L., and Lewis, D. (1969). Carbohydrate movement from autotrophs to heterotrophs in parasitic and mutualistic symbiosis. *Biol. Rev.* 44, 17–85. doi: 10.1111/j.1469-185x.1969.tb00821.x
- Smith, D. C., and Drew, E. A. (1965). Studies in physiology of lichens v. translocation from algal layer to medulla in *Peltigera polydactyla*. *New Phytol.* 64, 195–200. doi: 10.1111/j.1469-8137.1965.tb05390.x
- Smith, H. B., Dal Grande, F., Muggia, L., Keuler, R., Divakar, P. K., Grewe, F., et al. (2020). Metagenomic data reveal diverse fungal and algal communities associated with the lichen symbiosis. *Symbiosis* 82, 133–147. doi: 10.1007/s13199-020-00699-4
- Spribile, T., Tuovinen, V., Resl, P., Vanderpool, D., Wolinski, H., Aime, M. C., et al. (2016). Basidiomycete yeasts in the cortex of ascomycete macrolichens. *Science* 353, 488–492. doi: 10.1126/science.aaf8287
- Stolyar, S., Van Dien, S., Hillesland, K. L., Pinel, N., Lie, T. J., Leigh, J. A., et al. (2007). Metabolic modeling of a mutualistic microbial community. *Mol. Syst. Biol.* 3:92. doi: 10.1038/msb4100131
- Taffs, R., Aston, J. E., Brileya, K., Jay, Z., Klatt, C. G., McGlynn, S., et al. (2009). In silico approaches to study mass and energy flows in microbial consortia: a syntrophic case study. *BMC Syst. Biol.* 3:114. doi: 10.1186/1752-0509-3-114
- Tagirdzhanova, G., Saary, P., Tingley, J. P., Diaz-Escandon, D., Abbott, D. W., Finn, R. D., et al. (2021). Predicted input of uncultured fungal symbionts to a lichen symbiosis from metagenome-assembled genomes. *Genome Biol. Evol.* 2021:evab047.
- ten Veldhuis, M. C., Ananyev, G., and Dismukes, G. C. (2020). Symbiosis extended: exchange of photosynthetic O₂ and fungal-respired CO₂ mutually power metabolism of lichen symbionts. *Photosynthes. Res.* 143, 287–299. doi: 10.1007/s11220-019-00702-0
- Terfve, C., and Saez-Rodriguez, J. (2012). Modeling signaling networks using high-throughput phospho-proteomics. *Adv. Exp. Med. Biol.* 736, 19–57. doi: 10.1007/978-1-4419-7210-1_2
- Tuovinen, V., Ekman, S., Thor, G., Vanderpool, D., Spribile, T., and Johannesson, H. (2019). Two basidiomycete fungi in the cortex of wolf lichens. *Curr. Biol.* 29, 476–483. doi: 10.1016/j.cub.2018.12.022
- Vardi, L., Ruppin, E., and Sharan, R. (2012). A linearized constraint-based approach for modeling signaling networks. *J. Comput. Biol.* 19, 232–240. doi: 10.1089/cmb.2011.0277
- Varma, A., and Palsson, B. O. (1994). Stoichiometric flux balance models quantitatively predict growth and metabolic by-product secretion in wild-type *Escherichia coli* W3110. *Appl. Environ. Microbiol.* 60, 3724–3731. doi: 10.1128/aem.60.10.3724-3731.1994
- Venkateshwaran, M., Volkening, J. D., Sussman, M. R., and Ane, J. M. (2013). Symbiosis and the social network of higher plants. *Curr. Opin. Plant Biol.* 16, 118–127. doi: 10.1016/j.pbi.2012.11.007
- Vicente, C., and Perezurria, E. (1989). Production and secretion of urease by *Evernia prunastri* thallus and its symbionts. *Endocytobios. Cell Res.* 6, 87–97.
- Vivas, M., Sacristan, M., Legaz, M. E., and Vicente, C. (2010). The cell recognition model in chlorolichens involving a fungal Lectin binding to an algal ligand can be extended to cyanolichens. *Plant Biol.* 12, 615–621.
- Wang, Y., Yuan, X., Chen, L., Wang, X., and Li, C. (2018). Draft genome sequence of the lichen-forming fungus *Ramalina intermedia* strain YAF0013. *Genome Announc.* 6:e0478-18.
- Wang, Y., Zhang, X., Zhou, Q., Zhang, X., and Wei, J. (2015). Comparative transcriptome analysis of the lichen-forming fungus *Endocarpon pusillum* elucidates its drought adaptation mechanisms. *Sci. China Life Sci.* 58, 89–100. doi: 10.1007/s11427-014-4760-9
- Wei, S. S., Jian, X. X., Chen, J., Zhang, C., and Hua, Q. (2017). Reconstruction of genome-scale metabolic model of *Yarrowia lipolytica* and its application in overproduction of triacylglycerol. *Bioresour. Bioprocess.* 4:51.
- West, N. J., Parrot, D., Fayet, C., Grube, M., Tomasi, S., and Suzuki, M. T. (2018). Marine cyanolichens from different littoral zones are associated with distinct bacterial communities. *PeerJ* 6:e5208. doi: 10.7717/peerj.5208
- Ye, C., Zou, W., Xu, N., and Liu, L. (2014). Metabolic model reconstruction and analysis of an artificial microbial ecosystem for vitamin C production. *J. Biotechnol.* 182–183, 61–67. doi: 10.1016/j.jbiotec.2014.04.027
- Yen, J. Y., Nazem-Bokaei, H., Freedman, B. G., Athamneh, A. I. M., and Senger, R. S. (2013). Deriving metabolic engineering strategies from genome-scale modeling with flux ratio constraints. *Biotechnol. J.* 8, 581–594. doi: 10.1002/biot.201200234
- Yim, H., Haselbeck, R., Niu, W., Pujol-Baxley, C., Burgard, A., Boldt, J., et al. (2011). Metabolic engineering of *Escherichia coli* for direct production of 1,4-butanediol. *Nat. Chem. Biol.* 7, 445–452.
- Zhang, X., and Reed, J. L. (2014). Adaptive evolution of synthetic cooperating communities improves growth performance. *PLoS One* 9:e108297. doi: 10.1371/journal.pone.0108297
- Zhuang, K., Izallalen, M., Mouser, P., Richter, H., Risso, C., Mahadevan, R., et al. (2011). Genome-scale dynamic modeling of the competition between *Rhodospirillum rubrum* and *Geobacter* in anoxic subsurface environments. *ISME J.* 5, 305–316. doi: 10.1038/ismej.2010.117
- Zomorodi, A. R., and Maranas, C. D. (2012). OptCom: a multi-level optimization framework for the metabolic modeling and analysis of microbial communities. *PLoS Computat. Biol.* 8:e1002363. doi: 10.1371/journal.pcbi.1002363
- Zomorodi, A. R., and Segre, D. (2016). Synthetic ecology of microbes: mathematical models and applications. *J. Mol. Biol.* 428, 837–861. doi: 10.1016/j.jmb.2015.10.019

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 Nazem-Bokaei, Hom, Warden, Mathews and Gueidan. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.