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Synthetic biology of cyanobacteria: unique challenges and opportunities

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INTRODUCTION

Cyanobacteria have garnered a great deal of attention recently as biofuel-producing organisms. Their key advantage over other bacteria is their ability to use photosynthesis to capture energy from sunlight and convert CO₂ into products of interest. As compared with eukaryotic algae and plants, cyanobacteria are much easier to manipulate genetically and grow much faster. They have been engineered to produce a wide and ever-expanding range of products including fatty acids, long-chain alcohols, alkanes, ethylene, polyhydroxybutyrate, 2,3-Butanediol, ethanol, and hydrogen. These processes have been reviewed recently (Gronenberg et al., 2013) and will not be covered in detail in this review. Rather, we will look toward how the techniques of the emerging field of synthetic biology might bear fruit in improving the output of such engineered strains. Due to the low price of commodity goods like fuels and platform chemicals, it is critical to maximize the productivity of engineered strains to make them economically competitive. We believe that the tools of synthetic biology can help with this challenge.

Specifically, this review will cover systems, parts, and methods of analysis for synthetic biology. Synthetic biology requires a well-characterized host or “chassis” strain that can be genetically manipulated with ease and predictability. Ideally, the host should grow quickly and tolerate a range of environmental conditions. The host should be simple to cultivate using readily available laboratory equipment and inexpensive growth media. Simple, rapid, and high-throughput techniques should be available for procedures like DNA/RNA isolation, metabolomics, and proteomics. To achieve modular, “plug-and-play” modification of the host strain, its metabolism and regulatory systems must be well-characterized under a wide variety of relevant conditions. Since cyanobacterial biofuel production processes will need to use sunlight as an energy source to be economically and environmentally useful, the day/night cycle will be particularly relevant; the intermittent nature of this energy source will be a key engineering challenge. We will discuss which cyanobacterial chassis have been used and their relative merits and unique traits. Ultimately, the hope is that one of these strains might be developed to become a “green E. coli” for which a wide variety of genetic parts and systems are available for easy modification. Next, we will discuss the critical issue of how gene expression can be controlled in cyanobacteria. Compared with other systems, there are few examples of simple and effective controllable promoters in cyanobacteria. We will also discuss methods for analysis of gene expression using light-emitting reporters and for global analysis of metabolism using either constraint-based modeling or measurement of ¹³C labeling.
GENETIC MODIFICATION OF CYANOBACTERIA

Several strains of cyanobacteria are known which are readily amenable to genetic modification (See Table 1). Such modifications can be performed either in cis (through chromosome editing) or in trans (through plasmid addition) and synthetic biology experiments have used both approaches. We discuss advantages and disadvantages of each approach, as well as recent technical developments below. While even the best cyanobacterial model systems are still far from being a “green E. coli,” many tools are already available and more are being developed. The future holds great promise for this field.

GENETIC MODIFICATION IN CIS: CHROMOSOME EDITING

Cis genetic modification is the most common approach in cyanobacterial synthetic biology. This approach takes advantage of the capability of many cyanobacterial strains for natural transformation and homologous recombination (see Table 1) to create insertion, deletion, or replacement mutations in cyanobacterial chromosomes. Traditionally, strains have been transformed with selectable markers linked to any sequence of interest and flanked by sequences homologous to any non-essential sequence on the chromosome (See Figure 1).

This strategy allows the creation of targeted mutations to the chromosome, but sometimes raises concerns about segregation in polyploid strains. However, once segregated, such mutations can be stable over long time periods even in the absence of selective pressure from added antibiotics (Liu et al., 2011; Wang et al., 2013). While such stability is desirable, systems that create major metabolic demand, by for example redirecting flux into biofuel-producing pathways, will face greater selective pressures for mutation or loss of heterologous genes.

Recently, several methods have been developed that allow the creation of markerless mutations in cyanobacterial chromosomes (Figure 1B). Two of these methods operate on a similar principle: First, a conditionally toxic gene is linked to an antibiotic resistance cassette and then inserted into the chromosome.

Table 1 | Model strains of cyanobacteria for synthetic biology.

| Strain          | Genetic methods                         | Ideal growth temp (°C) | Doubling time (h) | Metabolisms                      | Genome-scale models? | Notes                                                   | References                  |
|-----------------|-----------------------------------------|------------------------|-------------------|---------------------------------|-----------------------|--------------------------------------------------------|----------------------------|
| *Synechocystis* sp. PCC 6803 | Conjugation, natural transformation, Tn5 mutagenesis, fusion PCR | 30                     | 6–12               | Mixotrophic, autotrophic        | Yes                   | Extensive systems biology datasets are available | Heidorn et al., 2011 |
| *Synechococcus elongatus* PCC 7942 | Conjugation, natural transformation, Tn5 mutagenesis | 38                     | 12–24              | Autotrophic                     | No                    | A model strain for the study of circadian clocks | Chen et al., 2012 |
| *Synechococcus* sp. PCC 7002 | Conjugation, natural transformation | 38                     | 3.5                | Mixotrophic, autotrophic        | Yes                   | Among the fastest-growing strains known | Xu et al., 2011 |
| *Anabaena variabilis* PCC 7120 | Conjugation, natural transformation | 30                     | >24                | Mixotrophic, autotrophic        | No                    | Nitrogen-fixing, Filamentous | Zhang et al., 2007 |
| *Leptolyngbya* sp. Strain BL0902 | Conjugation, Tn5 mutagenesis | 30                     | ~20                | Autotrophic                     | No                    | Filamentous, Grows well in outdoor photo-bioreactors in a broad range of conditions | Taton et al., 2012 |
with selection for antibiotic-resistant mutants. Next, a second transformation is carried out in which the resistance cassette and toxin gene are deleted, and markerless mutants are selected which have lost the toxic gene. This principle has been used in cyanobacteria with the *B. subtilis* levansucrase synthase gene sacB, which confers sucrose sensitivity (Lagarde et al., 2000) as well as with *E. coli* mazF, a general protein synthesis inhibitor expressed under a nickel-inducible promoter (Cheah et al., 2013). This latter system has advantages for cyanobacterial strains that are naturally sucrose-sensitive. Either method allows the reuse of a single selectable marker for making multiple successive changes to the chromosome. In addition to these methods, a third system operates on a similar principle—a cyanobacterial strain that is streptomycin resistant due to a mutation in the *rps12* gene can be made streptomycin-sensitive by expressing a second heterologous copy of wild type *rps12* linked to a kanamycin (or another antibiotic) resistance cassette as well as any sequence of interest. Streptomycin-resistant, kanamycin-sensitive markerless mutants can be recovered in a second transformation (Takahama et al., 2004). Although this method can also be used to make successive markerless mutants, it requires a background strain that is streptomycin-resistant due to an altered ribosome. Thus, it may not be an ideal method for synthetic biology studies that seek to draw conclusions about translation in wild-type systems. For the ability to transfer any translated genetic parts or parts involved in translation (such as ribosome binding sites) to other strains, this mutation could be problematic. A possible advantage of this system is that both selections are positive selections, whereas the sacB or mazF systems require a negative selection in their second transformation. Care must be taken to ensure that sucrose resistance is due to loss, as opposed to mutation, of the counter-selectable marker. Recombinase-based systems including Cre-LoxP [in *Anabaena* sp. PCC7120, (Zhang et al., 2007)] or FLP/FRT [in *Synechocystis* sp. PCC6803 and *Synechococcus elongatus* PCC7942, (Tan et al., 2013)] have also been used to engineer mutants that lack a selectable marker. However, these methods leave a scar sequence, meaning that the final chromosomal sequence is not completely user-specifiable and also that multiple mutations using this technique in the same cell line may potentially lead to undesirable crossover events or other unexpected results. Until recently, it has been difficult to create mutants at high throughput in cyanobacterial strains, as transposon-based methods developed for use in other strains can work poorly in cyanobacterial hosts. However, libraries can be created in other strains and subsequently transferred to a cyanobacterial host via homologous recombination. A Tn7-based library containing ∼10,000 lines was recently created to screen for strains with increased polyhydroxybutyrate (PHB) production (Tyo et al., 2009) and a similar approach has been taken for finding mutants in circadian clock function in *Synechococcus* 7942 (Holtman et al., 2005) and later extended to include insertions into nearly 90% of open reading frames in that strain (Chen et al., 2012). Chromosomal DNA fragments were first cloned into a plasmid library in *E. coli* and then the library was mutagenized with Tn7 before homologous recombination back into the cyanobacterial host strain. This could be an especially valuable approach for the validation of genome-scale models of cyanobacterial metabolism (see below).

**GENETIC MODIFICATION in trans: FOREIGN PLASMIDS**

Although transgene expression in cis is the most common approach in cyanobacterial research, genes are also routinely expressed in cyanobacteria in trans (Huang et al., 2010; Landry et al., 2012; Huang and Lindblad, 2013). In synthetic biology and metabolic engineering of other prokaryotes, this is by far the more common approach, and has led to such standardized approaches as “Bio-Brick” assembly in which standardized genetic “parts” such as promoters, ribosome binding sites, genes, and terminators can be readily swapped in and out of standard plasmids (http://partsregistry.org). This move toward standardization of genetic parts is a critical aim for synthetic biology, independent of the chassis organism or method of transformation. However, a limited number of plasmids are available for expression in cyanobacterial hosts. Plasmid assembly for expression in cis or in trans in cyanobacterial hosts has generally been performed in *E. coli* because of the longer growth times that would be associated with assembling vectors in cyanobacterial hosts (Figure 2A). This requires broad host range plasmids. However, with the rise of in vitro assembly methods such as SLIC (Li and Elledge, 2007), Gibson assembly (Gibson et al., 2009), CPEC (Quan and Tian, 2009), fusion PCR (Szewczyk et al., 2007), and Golden Gate (Engler and Marillonnet, 2011), this limitation may become less important over time (Figure 2B). These next-generation cloning methods have been reviewed elsewhere (Hilson et al., 2012) and will not be covered here. Fusion PCR has been used to construct linear DNA fragments for homologous recombination in cyanobacterial chromosomes (Nagarajan et al., 2011), but to our knowledge replicative vectors for cyanobacteria have so far not been constructed without the use of a helper heterotrophic strain. Techniques for in vivo assembly of plasmids that have been developed for yeast (Shao and Zhao, 2009) may be adaptable to cyanobacteria because of their facility for homologous recombination (Figure 2C). Such an improvement could greatly speed

![FIGURE 2 | DNA assembly methods. (A) Traditionally in cyanobacterial synthetic biology, plasmids are assembled in vitro and then propagated in *E. coli* before being transformed into cyanobacteria. (B) More recently, methods have been developed for in vitro assembly and direct transformation via fusion PCR. (C) Another recent method has been developed for in vivo plasmid assembly via homologous recombination in yeast which may also be applicable in certain cyanobacterial strains.](http://www.frontiersin.org)
up the process of making cyanobacterial mutant strains, either for modification in cis or in trans. The major technical challenge for such an approach is that the long time after transformation required to isolate cyanobacterial mutants (typically 1 week or more) means it is critical to have high-fidelity assembly methods to avoid a time-consuming screening process.

Although shuttle vectors do exist for cyanobacteria, there has been little characterization of their copy numbers in cyanobacterial hosts, and the lack of replicative vectors with varied copy numbers limits the valuable ability to control the expression level of heterologous genes by selecting their copy number (Jones et al., 2000; Dunlop et al., 2011). Plasmids derived from RSF1010 appear to have a copy number of 10–30 (or ~1–3 per chromosome) in Synechocystis sp. PCC 6803 (Ng et al., 2000; Huang et al., 2010), but copy numbers of other broad host-range plasmids have not been quantified to date. Endogenous plasmids of cyanobacteria have also been used as target sites for expression of heterologous genes in Synechococcus sp. PCC 7002 (Xu et al., 2011). This strain harbors several endogenous plasmids whose copy numbers range from ~1 to 8 per chromosome, with an approximate chromosome copy number of 6 per cell. Synechocystis sp. PCC 6803 also has plasmids whose copy numbers span a similar range [from ~0.4–8 per chromosome (Berla and Pakrasi, 2012)]. The origins of replication from these plasmids constitute a source of genetic parts that could be used to generate cyanobacterial expression plasmids having a range of copy numbers, and which could potentially be modified to create higher or lower-copy plasmids that are compatible with existing plasmids in various cyanobacterial systems. The range of shuttle vectors that have been used in cyanobacterial hosts has been recently reviewed (Wang et al., 2012). While many tools are available for genetic modification of these biotechnologically promising strains, opportunities abound to develop new and improved tools that will allow research to proceed faster.

**UNIQUE CHALLENGES OF THE CYANOBACTERIAL LIFESTYLE**
Organisms that survive using sunlight as a primary nutrient face unique challenges. These must be better understood and addressed to fulfill the biotechnological promise of cyanobacteria through synthetic biology.

**LIFE IN A DIURNAL ENVIRONMENT**
A primary goal of synthetic biology in cyanobacteria is to use photosynthesis to convert CO₂ into higher-value products such as biofuels and chemical precursors. To make such a process economically and environmentally feasible will require using sunlight as a primary energy source. While some cyanobacteria are facultative heterotrophs, their key advantage over obligate heterotrophic bacteria is photosynthesis. Unlike heterotrophic growth environments where carbon and energy sources can be provided more uniformly both in space and time, sunlight will only be available during the day and will be attenuated as it passes through the culture. Under certain conditions, cultures may be able to take advantage of a “flashing light effect” to integrate spatially uneven illumination by storing chemical energy when in bright light near the reactor surface and using that energy to conduct biochemistry during time spent in the dark away from the reactor surface. This ability will depend on light intensity, mixing rates, reactor geometry, and likely other factors. Certain diazotrophic cyanobacteria can even use daylight to continue growth during the night. *Cyanothoece sp.* ATCC 51142 [along with several other strains (Taniuchi et al., 2008; Latysheva et al., 2012; Pfreundt et al., 2012)] is a unicellular diazotrophic cyanobacterium that performs photosynthesis and accumulates glycogen during the day, and then during the night breaks down its glycogen reserves to supply energy for nitrogen fixation. Thus, these strains spread out the energy available from sunlight over a 24-h period. This process involves a genome-wide oscillation in transcription, with more than 30% of genes oscillating in expression between day and night (Stockel et al., 2008). To take full advantage of sunlight, synthetic systems must be created that are capable of responding appropriately to this challenging dynamic environment. It has recently been shown that biofuel-producing strains that dynamically tune the expression of heterologous pathways in response to their own intracellular conditions produce more biofuel and exhibit greater stability of heterologous pathways (Zhang et al., 2012). As challenging as the design of such a system was for batch heterotrophic cultures, it will be even more challenging in production environments that include a diurnal light cycle.

While not all strains exhibit as complete a physiological change between day and night as *Cyanothoece* 51142, all cyanobacteria do have a circadian clock that adapts them to their autotrophic lifestyle. The cyanobacterial circadian clock is anchored by master regulators KaiA, KaiB, and KaiC, which act by cyclically phosphorylating and dephosphorylating each other (Akiyama, 2012). While the circadian rhythm can be reconstituted in *vitro* using the three Kai proteins in the presence of ATP (Nakajima et al., 2005), the accurate maintenance of this clock in *vivo* depends on proper protein turnover (Holtman et al., 2005), on codon selection in the kaiBC transcript (Xu et al., 2013), on transcriptional feedback (Teng et al., 2013), and on the controlled response of the entire program of cellular transcription to the output of the KaiABC oscillator. While disturbing rhythmicity can lead to strains that grow better under constant light, the circadian clock is adaptive for strains living in a dynamic environment (Woelfle et al., 2004; Xu et al., 2013). Therefore, integrating synthetic gene circuits such as biofuel production processes into the circadian rhythm of cyanobacterial hosts will likely lead to both improved production and improved strain stability in outdoor production environments.

**REDIRECTING CARBON FLUX BY DECOUPLING GROWTH FROM PRODUCTION**
While redirecting carbon flux is a challenge in all metabolic engineering efforts, it has been suggested that stringent control of fixed carbon partitioning among central metabolic pathways poses a major limitation to chemical production especially in photosynthetic organisms (Melis, 2013). During the growth phase, it may be true that carbon partitioning is tightly controlled by any number of mechanisms including metabolite channeling or simply high demand for metabolic intermediates. However, biofuel production during non-growth phases (Atsumi et al., 2009; Liu et al., 2011; Wang et al., 2013) demonstrates that under appropriate conditions, cyanobacterial hosts can produce biofuel...
compounds with higher selectivity, since biofuel can be produced by metabolically active cells even in the absence of growth. Enhancing their productivity in this phase is a major opportunity for cyanobacterial synthetic biologists to overcome these limits on carbon partitioning. Capturing this opportunity will require designing complete metabolic circuits that remain highly active during stationary phase.

RNA-BASED REGULATION

Recently, regulation of gene expression through RNA mechanisms has received great attention across bacterial clades (Selinger et al., 2000; Sharma et al., 2010; Mitschke et al., 2011). While these mechanisms of regulation may be important in all bacteria, their prominence is perhaps the greatest in the cyanobacteria and may help these diurnal organisms adapt to their highly dynamic environment: in a recent dRNA-seq study, many of the most highly expressed RNAs belonged to families of non-coding RNAs which are present in nearly all sequenced cyanobacteria, but not in any other organisms (Gierga et al., 2009; Mitschke et al., 2011). While their high expression in Synechocystis 6803 suggests functional importance for non-coding RNAs, few have clearly elucidated functions to date. syr1 overexpression has been shown to lead to a severe growth defect in Synechocystis 6803 (Mitschke et al., 2011). Another small RNA, isiR, has a critical function in stress response in Synechocystis 6803. isiR binds to the mRNA (isiA) for the iron-stress inducible protein, which when translated, forms a ring around trimers of photosystem I, preventing their activity and thus oxidative stress in the absence of sufficient iron (Duhring et al., 2006). The binding of isiR to isiA appears to result in rapid degradation. This particular arrangement allows a very rapid and emphatic response to iron repletion in cyanobacteria, since a large pool of isiA transcripts can be quickly silenced and marked for degradation by transcription of the antisense isiR. Although little is so far known about the generality of this type of regulation, the dynamics of this response might also be effective to use for synthetic systems in cyanobacteria that live in the presence of light as an intermittently available but critical nutrient.

While non-coding RNA has received a lot of recent attention, two-component systems make up the most widely studied family of environmental response regulators in cyanobacteria. Many of these systems have known functions in response to diverse environmental stimuli such as nitrogen, phosphorous, CO2, temperature, salt, and light intensity and quality (Ashby and Houmard, 2006; Montgomery, 2007). Many of the most widely-used systems in the construction of synthetic biological devices (such as the ara and lux clusters) use 2-component systems, and even combine 2-component systems with non-coding RNA to control system dynamics (Waters and Bassler, 2006). As synthetic biology advances into the construction of more and more complex systems, there will be a growing need to understand and use all of the different mechanisms available for control of gene expression and enzyme activity in cyanobacteria.

PARTS FOR CYANOBACTERIAL SYNTHETIC BIOLOGY

While cyanobacteria are promising organisms for biotechnology, synthetic biology tools for these organisms lag behind what has been developed for E. coli and yeast (Heidorn et al., 2011). Furthermore, synthetic biology tools developed in E. coli or yeast often do not function as designed in cyanobacteria (Huang et al., 2010). Here, we discuss inducible promoters and reporters in cyanobacteria, and cultivation systems that will allow their testing at increased throughput. Refining such systems will make cyanobacterial synthetic biology more user-friendly, a central goal for developing the “green E. coli.”

INDUCIBLE PROMOTERS

Creation of synthetic biology systems that predictably respond to a specific signal often depends upon inducible promoters for transcriptional control. An ideal inducible promoter will have the following properties: (1) It will not be activated in the absence of inducer. (2) It will produce a predictable response to a given concentration of inducer or repressor. This response may be digital (i.e., on/off) or graded change with different concentrations of inducer/repressor. (3) The inducer at saturating concentrations should have no harmful effect on the host organism. (4) The inducer should be cheap and stable under the growth conditions of the host. Finally, (5) the inducible system should act orthogonally to the host cell’s transcriptional program. Ideal transcriptional repressors should not bind to native promoters and if non-native transcriptional machinery is used (such as T7 RNA polymerase) it should not initiate transcription from native promoters. Promoters must perform as ideally as possible in order to be used in the construction of more complex genetic circuits (Moon et al., 2012).

Many common inducible promoters in cyanobacteria respond to transition metals. These have often been the basis of metal detection systems (Erbe et al., 1996; Boyanapalli et al., 2007; Peca et al., 2007, 2008; Blasi et al., 2012). Cyanobacteria balance metal intake for the organisms’ needs against potential oxidative stress and protein denaturation (Michel et al., 2001; Peca et al., 2008) via tightly regulated systems. As shown in Table 2, cyanobacteria’s metal-responsive promoters frequently show greater than 100-fold dynamic range. For example, the promoter for the Synechocystis sp. PCC 6803 gene, coaA, was induced 500-fold by 6 μM Co2+ (Guerrero et al., 2012), and Pmfb from Synechococcus elongatus PCC 7942 was induced 300-fold by 2 μM Zn2+ (Erbe et al., 1996). The most responsive cyanobacterial promoters reported were Pmfb from Synechocystis sp. PCC 6803, responding 1000-fold to 0.5 μM Ni2+ (Peca et al., 2007), and PisaAB also from Synechocystis sp. PCC 6803, repressed 5000-fold by 30 μM Fe3+ following depletion (Kunert et al., 2003).

While the sensitivity of these promoters to low concentrations of ions may seem like an advantage, in practice it can make them difficult to use. Glassware must be thoroughly cleaned according to special protocols to remove trace metals and cells often have to be starved for extended periods, inducing stress responses, to use such inducible systems. Additionally, promoters endogenous to a chassis strain are woven into a complex, incompletely understood regulatory system. In this system, promoters are activated by multiple inducers, such as PcoaT (Co2+ and Zn2+) and PziaA (Cd2+ and Zn2+), both from Synechocystis sp. PCC 6803 and inducers can also activate multiple promoters, such as Cd2+ inducing ziaA and isiA (Blasi et al., 2012).
Table 2 | Inducible promoters used in cyanobacterial hosts.

| Promoter | Source | Inducer/repressor and concentration | Expression host | Expressed gene | Dynamic range | Measure of expression | References |
|----------|--------|------------------------------------|-----------------|---------------|---------------|----------------------|------------|
| **METAL-INDUCIBLE PROMOTERS** | | | | | | | |
| **ArsB** | Synechocystis sp. PCC 6803 | Inducer AsO$^{2-}$ 720 Mm | Synechocystis sp. PCC 6803 | arsB | 100-fold RT-PCR | Blasi et al., 2012 |
| **ZiaA** | Synechocystis sp. PCC 6803 | Inducer Cd$^{2+}$ 2 μM | Synechocystis sp. PCC 6803 | ziaA | 10-fold RT-PCR | Blasi et al., 2012 |
| **coat** | Synechocystis sp. PCC 6803 | Inducer Co$^{2+}$ 6 μM | Synechocystis sp. PCC 6803 | Gene encoding EFE from Pseudomonas syringae | 500-fold 48 nL ethylene mL$^{-1}$ h$^{-1}$ | Guerrero et al., 2012 |
| **coat** | Synechocystis sp. PCC 6803 | Inducer Co$^{2+}$ 6.4 μM | Synechocystis sp. PCC 6803 | coaR + luxAB | 70-fold 70 RLU$^a$ | Peca et al., 2008 |
| **nrsB** | Synechocystis sp. PCC 6803 | Inducer Co$^{2+}$ 3 μM | Synechocystis sp. PCC 6803 | nrsB | 10-fold RT-PCR | Peca et al., 2007 |
| **coat** | Synechocystis sp. PCC 6803 | Inducer Co$^{2+}$ 1 μM | Synechocystis sp. PCC 6803 | coaT | 10-fold RT-PCR | Peca et al., 2007 |
| **petE** | Synechocystis sp. PCC 6803 | Inducer Cu$^{2+}$ 0.5 μM | Synechocystis sp. PCC 6803 | Gene encoding EFE from Pseudomonas syringae | 5-fold 28 nL ethylene mL$^{-1}$ h$^{-1}$ | Guerrero et al., 2012 |
| **petE** | Synechocystis sp. PCC 6803 | Inducer Cu$^{2+}$ 3 μM | Anabaena sp. PCC 7120 | hetP | 4.5-fold 8% heterocyst frequency | Higa and Callahan, 2010 |
| **petE** | Synechocystis sp. PCC 6803 | Inducer Cu$^{2+}$ 0.3 μM | Anabaena sp. PCC 7120 | hetN (prevents heterocyst formation) | Qualified but not quantified | 0% heterocysts from 10% uninduced | Callahan and Buikema, 2001 |
| **isiAB** | Synechocystis sp. PCC 6803 | Repressor Fe$^{3+}$ 30 μM | Synechocystis sp. PCC 6803 | isiAB + gfp | 5000-fold From 5000 RFU$^a$ | Kunert et al., 2003 |
| **idiA** | Synechococcus elongatus PCC 7942 | Repressor Fe$^{3+}$ 0.043 mM | Synechococcus elongatus PCC 7942 | luxAB | 170-fold Luminescence (5.3 × 10$^6$ cpn) | Michel et al., 2001 |
| **isiAB** | Synechococcus sp. strain PCC 7002 | Repressor Fe$^{3+}$ 100 nM | Synechococcus sp. strain PCC 7002 | luxAB from Vibrio harveyi | 2-fold From 0.012 RLU cell$^{-1}$ s$^{-1}$ | Boyanapalli et al., 2007 |
| **nrsB** | Synechocystis sp. PCC 6803 | Inducer Ni$^{2+}$ 0.5 μM | Synechocystis sp. PCC 6803 | nrsB | 1000-fold RT-PCR | Peca et al., 2007 |
| **nrsB** | Synechocystis sp. PCC 6803 | Inducer Ni$^{2+}$ 5 μM | Synechocystis sp. PCC 6803 | nrsB | 400-fold RT-PCR | Blasi et al., 2012 |
| **nrsB** | Synechocystis sp. PCC 6803 | Inducer Ni$^{2+}$ 6.4 μM | Synechocystis sp. PCC 6803 | nrsR + luxAB | 50-fold 50 RLU$^a$ | Peca et al., 2008 |
| **Smt** | Synechococcus elongatus PCC 7942 | Inducer Zn$^{2+}$ 2 μM | Synechococcus elongatus PCC 7942 | luxCDABE from Vibrio fisheri | 300-fold 325,000 cps luminescence | Erbe et al., 1996 |
| **ziaA** | Synechocystis sp. PCC 6803 | Inducer Zn$^{2+}$ 5 μM | Synechocystis sp. PCC 6803 | ziaA | 40-fold RT-PCR | Peca et al., 2007 |
| **ziaA** | Synechocystis sp. PCC 6803 | Inducer Zn$^{2+}$ 4 μM | Synechocystis sp. PCC 6803 | ziaA | 40-fold RT-PCR | Blasi et al., 2012 |
| **coat** | Synechocystis sp. PCC 6803 | Inducer Zn$^{2+}$ 3.2 μM | Synechocystis sp. PCC 6803 | coaR + luxAB | 25-fold 25 RLU$^a$ | Peca et al., 2008 |
| **coat** | Synechocystis sp. PCC 6803 | Inducer Zn$^{2+}$ 5 μM | Synechocystis sp. PCC 6803 | coaT | 10-fold RT-PCR | Peca et al., 2007 |

(Continued)
Table 2 | Continued

| Promoter | Source | Inducer/repressor and concentration | Expression host | Expressed gene | Dynamic range | Measure of expression | References |
|----------|--------|------------------------------------|-----------------|---------------|---------------|----------------------|------------|
| coat     | Synechocystis sp. PCC 6803 | Inducer Zn\(^+\) 4 μM | Synechocystis sp. PCC 6803 | coaT | 8-fold | RT-PCR | Blasi et al., 2012 |
| Smt      | Synechococcus elongatus PCC 7002 | Inducer Zn\(^+\) 2 μM | Synechocystis sp. PCC 6803 | Gene encoding EFE from Pseudomonas syringae | 2-fold | 2 nL ethylene mL\(^{-1}\) h\(^{-1}\) | Guerrero et al., 2012 |
| zieA\(^8\) | Synechocystis sp. PCC 6803 | Inducer Zn\(^+\) 3.5 μM | Synechocystis sp. PCC 6803 | hydA1 from Chlamydomonas reinhardtii | Qualified but not quantified | 109 nmol H\(_2\) mg Chl\(^{-1}\) min\(^{-1}\) | Berto et al., 2011 |
| **METABOLITE-INDUCIBLE PROMOTERS** | | | | | | | |
| tetR\(^b\) | E. coli | Inducer aTc 10³ ng/per ml | Synechocystis sp. strain ATCC27184 | eYFP | 290-fold | >10,000 RFU | Huang and Lindblad, 2013 |
| trp-lac | E. coli | Inducer IPTG 100 μM | Synechococcus elongatus PCC 7942 | invA and glf genes from Zymomonas mobilis | 160-fold for fructose + 30-fold for glucose | 160 μM fructose + 30 μM glucose | Niederholtmeyer et al., 2010 |
| Trc | E. coli | Inducer IPTG 1 mM | Synechococcus elongatus PCC 7942 | uidA from E. coli | 36-fold | 340 nmol MU min\(^{-1}\) mg protein\(^{-1}\) (β-Glucuronidase activity) | Geerts et al., 1995 |
| A1lacO-1 | E. coli | Inducer IPTG 1 mM | Synechocystis sp. PCC 6803 | Gene encoding EFE from Pseudomonas syringae | 8-fold | 170 nL ethylene mL\(^{-1}\) h\(^{-1}\) | Guerrero et al., 2012 |
| trc20 | E. coli | Inducer IPTG 2 mM | Synechocystis sp. PCC 6803 | Gene encoding GFPmut3B | 4-fold | 12 RFU\(^a\) (relative to PlacI) | Huang et al., 2010 |
| trc10 | E. coli | Inducer IPTG 2 mM | Synechocystis sp. PCC 6803 | Gene encoding GFPmut3B | 1.6-fold | 101 RFU\(^a\) (relative to PlacI) | Huang et al., 2010 |
| LlacO1\(^c\) | E. coli | Inducer IPTG 1 mM | Synechococcus elongatus PCC7942 | alsS (B. subtilis), alsD (A. hydrophila), and adh (C. beijerinckii) | 1.6-fold | 1.6 (relative to sADH and ALS activity) | Oliver et al., 2013 |
| Trc\(^d\) | E. coli | Inducer IPTG 1 mM | Synechocystis sp. PCC 6803 | Gene encoding EFE from Pseudomonas syringae | No significant difference | 170 nL ethylene mL\(^{-1}\) h\(^{-1}\) | Guerrero et al., 2012 |
| **MACRONUTRIENT-INDUCIBLE PROMOTERS** | | | | | | | |
| psbA2 | Synechocystis sp. PCC6803 | Inducer light 500 μmol photons m\(^{-2}\) s\(^{-1}\) | Synechocystis sp. PCC6803 | ispS from Pueraria montana (kudzu) | Qualified but not quantified | ~50 mg isoprene g DCW\(^{-1}\) d\(^{-1}\) | Lindberg et al., 2010 |
| psbA2\(^8\) | Synechocystis sp. PCC 6803 | Inducer light 50 μEm\(^{-2}\)s\(^{-1}\) | Synechocystis sp. PCC 6803 | hydA1 from Chlamydomonas reinhardtii | Qualified but not quantified | 130 nmol H\(_2\) mg Chl\(^{-1}\) min\(^{-1}\) | Berto et al., 2011 |
| psbA1 | Anabaena sp. PCC 7120 | Inducer light 30 μEm\(^{-2}\)s\(^{-1}\) | Anabaena sp. PCC 7120 | hetR from E. coli | Qualified but not quantified | 17% heterocyst frequency | Chaurasia and Apte, 2011 |
| nirA | Synechococcus elongatus PCC 7942 | Inducer/Repressor NO\(_2\)/NH\(_4\)+ 176 mM/176 mM | Synechocystis sp. PCC 6803 | Gene encoding p-hydroxyphenylpyruvate dioxygenase from Arabidopsis thaliana | 25-fold | 250 ng tocopherol mg DCW\(^{-1}\) | Qi et al., 2005 |

(Continued)
Thus, these promoters fall short according to criteria 2, 3, and 5 described above.

While few good choices have so far been available for inducible promoters in cyanobacteria, it will be helpful to understand the differences in the cellular machinery of *E. coli* and cyanobacteria in order to adapt existing systems for use in a cyanobacterial “green *E. coli*.” First, RNA polymerase (RNAP) is structurally different between *E. coli* and cyanobacteria. In cyanobacteria the β’ subunit of the RNAP holoenzyme is split into two parts, as opposed to one in most eubacteria, creating a different DNA binding domain (Imamura and Asayama, 2009). Being photosynthetic, circadian, and sometimes nitrogen-fixing, cyanobacteria also employ three sets of interconnected σ factors that are different than those used by *E. coli* (Imamura and Asayama, 2009). Guerrero et al. (2012) looked at the variation in the −35 and −10 regions of PAlacO−1 and Ptrc. PAlacO−1 is not inducible in *Synechocystis* sp PCC 6803 and had the “standard” bacterial structure in these regions while Ptrc−1, which produced an 8-fold response to IPTG in the same host, had a different structure in both regions. They postulated that *Synechocystis* 6803’s σ factors had different selectivity for these two regions. In fact, by systematically altering the bases between −10 and the transcription start site, a library of TetR-regulated promoters with improved inducibility were created in *Synechocystis* sp. strain ATCC27184 (a glucose-tolerant derivative of *Synechocystis* 6803). The best performing promoter induced a 290-fold change in response to 1 μg/ml aTc (Huang and Lindblad, 2013). This work demonstrates the improvements that can be seen when modifying parts to work in a particular chassis. However, the light-sensitivity of the inducer aTc required the use of special growth lights that may have had other effects on photoautotrophic metabolism. Further studies that follow in this vein of using well-characterized synthetic biology parts and modifying them to function optimally in a particular cyanobacterial chassis are likely to bear fruit.

The lack of inducibility seen in lac-derived promoters in cyanobacteria could also be a function of inadequate transport of IPTG into cells. Concentrations of IPTG above 1 mM have been shown to induce lac-derived promoters in organisms without an active lactose permease, like many cyanobacteria. By introducing an active lactose permease into *Pseudomonas fluorescens*, inducibility was boosted five times at 0.1 mM IPTG (Hansen et al., 1998). Evolving the Lac repressor for improved inducibility is another strategy. Gene expression improved ten times with 1 μM IPTG through rounds of error prone PCR and DNA shuffling (Satya Lakshmi and Rao, 2009). Strength of expression and inducibility may also vary between different cyanobacterial strains. IPTG caused as much as a 36-fold response using the trc promoter in *Synechococcus elongatus* PCC 7942, but little or no response in *Synechocystis* sp. PCC 6803 (See Table 2). Phylogenetic analysis of σ factors from six different cyanobacterial strains, including *Synechocystis* 6803, showed *S. elongatus* 7942 to be distinctive. *S. elongatus* 7942 has σ factors that are unique to marine cyanobacteria as well as a group 3 σ factor similar to those from the heterocyst-forming *Anabaena* sp. PCC 7120 (Imamura and Asayama, 2009). Understanding these strain-specific differences will enhance the synthetic biologist’s ability to design promoters with ideal characteristics in their chassis of choice. This relates to the ability to take up inducers as well as the optimal characteristics of inducers (as in the light-sensitivity of aTc) as described above.

REPORTERS

Characterization of synthetic biological circuits depends on a reporting method to track the expression, interaction, and position of proteins. Preferably the reporter should be detected without destruction of the organisms or additional inputs. Bacterial luciferase and fluorescent proteins are the most common non-invasive reporters. The lux operon is frequently used for reporting in cyanobacteria (Michel et al., 2001; Mackey et al., 2007; Peca et al., 2008) and is well-suited for real time reporting of gene expression due to the short half-life of the relevant enzymes (Ghim et al., 2010). The superior brightness of fluorescent proteins makes them more ideal for subcellular localization via microscopy or for cell-sorting methods. Fluorescent proteins are produced in an array of colors and also do not require additional substrates. Their use in cyanobacteria is somewhat complicated by the fluorescence of the organism’s photosynthetic pigments, but Cerulean, GFPmut3B (a mutant of green fluorescent protein) and EYFP (enhanced yellow fluorescent protein) have all been

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**Table 2** | Continued

| Promoter | Source | Inducer/repressor and concentration | Expression host | Expressed gene | Dynamic range | Measure of expression | References |
|---------|-------|-----------------------------------|-----------------|---------------|---------------|---------------------|------------|
| nirA    | *Synechococcus* elongatus PCC 7942 | Inducer/Repressor NO$_3^-$/NH$_4^+$ | *Synechococcus* elongatus PCC 7942 | cmpABCD | 5-fold | 260 nmol HCO$_3^-$ mg Chl$^{-1}$ | Omata et al., 1999 |
| Nir     | *Anabaena* sp. PCC 7120 | Inducer/Repressor NO$_3^-$/NH$_4^+$ | *Anabaena* sp. PCC 7120 | nir | Qualified but not quantified | 250 mg labeled proteins for NMR L$^{-1}$ | Desplancq et al., 2005 |

*In the presence of 5 μM DCMU, which inhibits the PSII-dependent oxygen evolution.

*Grown in the dark on 5 mM glucose.

*Leaky production of 2,3-butanediol, no IPTG, and 1 mM IPTG similar.

*Plac variants had differential expression early in growth phase but dynamic range was reduced as growth proceeded.

*RFU = Relative Fluorescence Units; RLU = Relative Luminescence Units.
used successfully in cyanobacteria as reporters of gene expression (Huang et al., 2010; Heidorn et al., 2011; Landry et al., 2012; Huang and Lindblad, 2013).

Bacterial luciferase luminesces upon oxidation of reduced flavin mononucleotide (Meighen, 1993). Fluorescent proteins also require oxygen to correctly-fold and fluoresce (Hansen et al., 2001). The light-dark cycle of nitrogen-fixing cyanobacteria provides temporal separation of the oxygen-sensitive nitrogenase from oxygen-evolving photosynthesis (Golden et al., 1997). During the dark cycle, respiration reduces intra-cellular oxygen levels so that nitrogenase can function. Therefore, neither bacterial luciferase nor traditional fluorescent proteins can likely be used to study cyanobacteria in their dark cycle or to report on synthetic biology systems that operate in these oxygen-depleted conditions. Using blue light photoreceptors from Bacillus subtilis and Pseudomonas putida, oxygen-independent flavin mononucleotide-binding florescent proteins have been devised (Drepper et al., 2007). With an excitation wavelength of 450 nm and an emission wavelength of 495 nm, they should perform well in cyanobacteria, although no data supporting this has been published yet. Functionality of these new fluorescent proteins was also improved by replacing a phenylalanine suspected of quenching with serine or threonine, resulting in a doubling of the brightness (Mukherjee et al., 2012). This expanding variety of easily readable reporter systems will be extremely valuable for cyanobacterial synthetic biology.

CULTIVATION SYSTEMS

To date, most synthetic biology and metabolic engineering work in cyanobacteria has been performed using simple, low-tech cultivation methods such as shake flasks or bubbling tubes grown under standard fluorescent light sources. Often, laboratory incubators have simply been retrofitted by the addition of fluorescent light sources available in home improvement stores. However, as light and CO₂ are major nutrients for cyanobacteria, it is critical to properly standardize the inputs of these resources to reliabily characterize biological parts. It is also critical to increase the throughput of cyanobacterial growth systems to be able to screen the large numbers of variants that can be generated by combinatorial methods, as is routinely performed by growing heterotrophic bacterial cultures in 96-well plate format. Growth of cyanobacteria in 6-well plates can be routinely performed in our lab and by others (Huang and Lindblad, 2013) along with 24-well plates (Simkovsky et al., 2012), but growth in 96-well plates is poor, limiting assay throughput and requiring more space in lighted chambers under consistent illumination, which is often a limitation. Simple, low-cost systems to reproducibly grow many cyanobacterial cultures in parallel are necessary.

GENOME-SCALE MODELING AND FLUXOMICS OF CYANOBACTERIA

A primary aim of cyanobacterial synthetic biology is the production of particular metabolites as biofuels or platform chemicals. As such, better understanding the metabolic phenotypes of wild-type and synthetic strains is a critical aim. While cyanobacterial metabolomics have been recently reviewed (Schwarz et al., 2013), here we describe recent progress in genome-scale modeling and fluxomics of cyanobacteria. These approaches can help guide the creation of synthetic strains with desirable metabolic phenotypes such as biofuel overproduction via in silico prediction or in vivo measurement of metabolic fluxes (See Figure 3). Specific to cyanobacterial systems, we highlight a number of challenges including complexity of modeling the photosynthetic metabolism and performing flux balance analysis (FBA), poor annotations of important metabolic pathways, and unavailability of in vivo gene essentiality information for most cyanobacteria. Finally, we focus on recent advancements in this area.
CHALLENGES
Incorporating photoautotrophy into metabolic models

FBA is a tool to make quantitative in silico predictions about metabolism (Fell and Small, 1986; Savinell and Palsson, 1992; Varma et al., 1993; Orth et al., 2010). An FBA model incorporates the stoichiometry of all genome-encoded metabolic reactions and assumes steady-state growth, such as during exponential phase. This assumption leads to a model that consists of a system of algebraic equations which state that the rate of producing any given metabolite is equal to the rate of consuming that metabolite. A solution to this system of equations is a possible answer to the question “what are all the metabolic fluxes in this system?” Since there are usually more reactions than metabolites, this system of equations is underdetermined and has many possible solutions. Therefore, one has to pick a solution that satisfies a biological objective, such as maximal growth, energy production, or byproduct formation (Varma and Palsson, 1994). For this purpose, a model will also include upper and lower bounds of fluxes that constrain the model to produce physically and biologically reasonable solutions.

Success of FBA greatly depends on the quality of the metabolic network reconstruction as well as the availability of regulatory constraints under a given environmental or growth condition. For instance, constraints can be added that disable or limit fluxes due to known regulatory constraints or substrate availability (Zomorrodhi et al., 2012). For cyanobacteria, the major challenges to develop a genome-scale metabolic model and subsequently perform FBA are the same ones faced by these organisms in their diurnal environment: how to incorporate light and how to differentiate light and dark metabolisms. Although it has been nearly a decade since publication of the first study applying FBA to cyanobacteria, it is only recently that models have incorporated complete descriptions of the light reactions of photosynthesis (Nogales et al., 2012). In so doing, these authors were able to highlight the critical importance of alternate electron flow pathways to growth under diverse environmental conditions, and to identify differences in metabolism during carbon-limited and light-limited growth. However, debate remains among photosynthesis researchers about the exact form of the light reactions (Heyes and Hunter, 2005; Kopceca et al., 2013). This uncertainty about the exact stoichiometry of metabolism is a challenge for the predictive power of FBA in photosynthetic systems. While FBA requires the assumption of a pseudo-steady state, all cyanobacteria must alternate between day and night metabolisms during a diurnal cycle. A recent model (Saha et al., 2012) of Cyanothece sp. ATCC 51142 utilizes proteomic data to model the diurnal rhythm of this strain, which fixes carbon during the day and nitrogen during the night (see section Unique Challenges of the Cyanobacterial Lifestyle).

Incompleteness of genome annotation

Genome scale models are built starting with an annotated genome sequence (see Figure 3), which allows prediction of which metabolic reactions are available in a given strain. However, genome annotation is constantly evolving, and open questions remain about important metabolic reactions in cyanobacteria.

The understanding of several key pathways in cyanobacteria has been recently revised. Zhang and Bryant (2011) identified enzymes from Synechococcus 7002 that can complete the TCA cycle in vitro and have homologues in most cyanobacterial species, which were previously thought to possess an incomplete TCA cycle. Based on this information, Synechocystis 6803 model iSyn731 (Saha et al., 2012) allows for a complete TCA cycle including these reactions. However, using flux variability analysis (Mahadevan et al., 2002; Mahadevan and Schilling, 2003) it was determined that this alternate pathway is not essential for maximal biomass production [unpublished results, (Saha et al., 2012)]. Fatty acid metabolism in cyanobacteria has unique properties that have been recently uncovered due to increased interest in these pathways for biofuel production. Both Synechocystis sp. PCC 6803 and Synechococcus elongatus PCC 7942 contain a single candidate gene annotated for fatty acid activation. While in both organisms the gene is annotated as acyl-CoA synthetase, it shows only acyl-ACP synthetase activity instead (Kaczmarzyk and Fulda, 2010). Further analysis also shows the importance of acyl-ACP synthetase in enabling the transfer of fatty acids across the membrane (Von Berlepsch et al., 2012). Quinone synthesis is another pathway with conflicting annotations. Cyanobacteria contain neither ubiquinone nor menaquinone (Collins and Jones, 1981). Despite the lack of ubiquinone within cyanobacteria, a number of cyanobacterial genomes contain homologs for six E. coli genes involved in ubiquinone biosynthesis (Sakuragi, 2004). Given these homologous genes it is probable that plastoquinone, a quinone molecule participating in the electron transport chain, is produced in cyanobacteria using a pathway very similar to that of ubiquinone production in proteobacteria. Wu et al. (2010) showed that Cyanothece 51142 contains an alternative pathway for isoleucine biosynthesis. Threonine ammonia-lyase, catalyzing the conversion of threonine to 2-ketobutyrate, is absent in Cyanothece 51142. Instead, this organism uses a citramalate pathway with pyruvate and acetyl-CoA as precursors for isoleucine synthesis. An intermediate in this pathway, namely ketobutyrate, can be converted to higher alcohols (propanol and butanol) via this non-fermentative alcohol production pathway. These active areas of research will help to better define cyanobacterial metabolism and allow the generation of models that can more accurately predict cellular phenotypes. While newer fluxomics techniques can yield powerful results in well-characterized strains, developing a “green E. coli” will also require expanded knowledge of biochemistry that to date can only come from older methods of single gene or single protein analysis.

Fewer mutant resources to test model accuracy

The quality or accuracy of any genome-scale metabolic model can be tested by contrasting the in silico growth phenotype with available experimental data on the viability of single or multiple gene knockouts (Thiele and Palsson, 2010). Any discrepancies between model predictions and observed results can aid in model refinement (Kumar and Maranas, 2009). For model strains besides cyanobacteria, concerted efforts to create complete mutant libraries have led to improvements in metabolic modeling. To the best of our knowledge, extensive in vivo gene essentiality data are available only for Synechocystis 6803 among the cyanobacteria in...
the CyanoMutants database (Nakamura et al., 1999; Nakao et al., 2010), but only for \( \sim 119 \) genes, compared with 731 genes associated with metabolic reactions in a recent genome-scale model (Saha et al., 2012). Thus, only a small subset of the model predictions on gene essentiality can be evaluated using available data for *Synechocystis* 6803, and the proportion is much lower for any other strain. While a genome-wide library of knockout mutants has been created in *Synechococcus* 7942 (Chen et al., 2012) and genome-scale models have yet had genome-scale models generated for their metabolism. Thus, stoichiometric models are emerging as a valuable tool for use across model cyanobacterial systems.

### 13C MFA analysis

While *in silico* models are great tools for generating hypotheses on how to use synthetic biology interventions to alter metabolism, they need to be complemented by fluxomics methods that allow *in vivo* measurement of metabolic fluxes to assess these interventions. Such a suite of tools allows the closure of the design-build-test engineering cycle in synthetic biology (Figure 3). To this end, Young et al. (2011) have developed a method to measure fluxes in autotrophic metabolism via dynamic isotope labeling measurements. In this approach, cultures are fed with a step-change from naturally labeled bicarbonate to NaH\(^{13}\)CO\(_3\) and the labeling patterns of metabolic intermediates are followed over a time-course to determine relative rates of metabolic flux. Previous studies (Yang et al., 2002) have also assessed metabolic fluxes under mixotrophic growth conditions, using a pseudo-steady-state approach in which cells are fed with \(^{13}\)C labeled glucose and metabolic fluxes are inferred from labeling patterns of proteinogenic amino acids. These studies have been extremely useful in identifying fluxes that exist *in vivo*, but have previously been regarded as wasteful or futile cycles, such as the oxidative pentose phosphate pathway and RuBP oxygenation. Comparisons between flux measurements (Young et al., 2011) and flux predictions (Saha et al., 2012) for *Synechocystis* 6803 have revealed the necessity of additional regulatory information for accurate *in silico* predictions of phenotype. These modeling and fluxomics efforts have resulted in deeper understanding of the metabolic capabilities of the modeled strains and of cyanobacteria in general.

### CONCLUSIONS

Cyanobacterial synthetic biology offers great promise for enhancing efforts to produce biofuels and chemicals in photoautotrophic hosts. While several cyanobacterial chassis strains have been used in synthetic biology efforts, the tools for their manipulation and analysis need greater development to unlock this potential and develop a “green *E. coli.*” Metabolic modeling is a complementary tool that can help guide the creation of synthetic strains with desirable phenotypes. By developing the tools for strain manipulation and control, synthetic biologists can unlock a bright future for the biotechnological use of abundant light and CO\(_2\).

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