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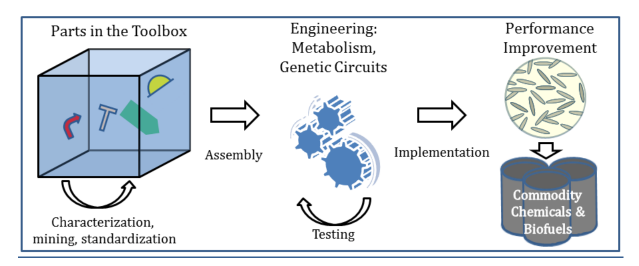
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1	Part by part: synthetic biology parts used in		
2	solventogenic Clostridia		
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Graphical Table of Contents.

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ABSTRACT: The solventogenic Clostridia are of interest to the chemical industry because of their natural ability to produce chemicals such as butanol, acetone and ethanol from diverse feedstocks. Their use as whole cell factories presents multiple metabolic engineering targets that could lead to improved sustainability and profitability of *Clostridium* industrial processes. However, engineering efforts have been held back by the scarcity of genetic and synthetic biology tools. Over the last decade, genetic tools to enable transformation and chromosomal modifications have been developed, but the lack of a broad palette of synthetic biology parts remains one of the last obstacles to the rapid engineered improvement of these species for bioproduction. We have systematically reviewed existing parts that have been used in the modification of solventogenic Clostridia, revealing a narrow range of empirically chosen and nonengineered parts that are in current use. The analysis uncovers elements, such as promoters, transcriptional terminators and ribosome binding sites where increased fundamental knowledge is needed for their reliable use in different applications. Together, the review provides the most comprehensive list of parts used and also presents areas where an improved toolbox is needed for full exploitation of these industrially important bacteria.

Keywords

Clostridium, ABE fermenation, solventogenesis, transcription termination, reporters, promoters

47	ABE: acetone-butanol-ethanol
48	ACE: allele-coupled exchange
49	aSD: anti Shine-Dalgarno sequence, found at the 3' end of 16S rRNA
50	CDS: coding sequence
51	FbFP: flavin-binding fluorescent protein
52	RBS: ribosome binding site
53	RNAP: RNA-polymerase
54	SD: Shine-Dalgarno sequence
55	UTR: untranslated region
56	
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62	we used to compare the overlaps of the outputs of the terminator prediction algorithms.
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Abbreviations

One of the earliest industrial biotechnology processes used by man was the production of acetone by bacterial fermentation. The bacterium used was from the genus *Clostridium*, which are Gram-positive, spore forming, obligately anaerobic organisms¹. In addition to acetone, these processes produced butanol and ethanol, leading to the fermentation being given the name 'ABE' for 'acetone-butanol-ethanol'2. ABE-producing clostridia typically show a biphasic growth pattern, producing acetic and butyric acids during the early stages of growth, and then undergoing a metabolic 'switch', reassimilating the produced acids and producing solvents³. Driven by the demands of industry, many different ABE-producing clostridial isolates were identified; those strains that were used in industry are primarily represented by four species^{4,5}, namely Clostridium acetobutylicum, Clostridium beijerinckii, Clostridium saccharobutylicum, and Clostridium saccharoperbutylacetonicum, although ABE fermentation has been observed in other clostridia⁶⁻⁸. Furthermore, not all butanol-producing clostridia produce the canonical trio of solvent products; for example, Clostridium pasteurianum produces butanol and 1,3-propanediol (instead of acetone) from glycerol⁹. While the ABE process had until recently fallen out of favour due to competition from the petrochemical industry, the necessity of identifying alternative fuels has renewed interest in the production of butanol as a potential biofuel candidate and sustainable commodity chemical¹⁰.

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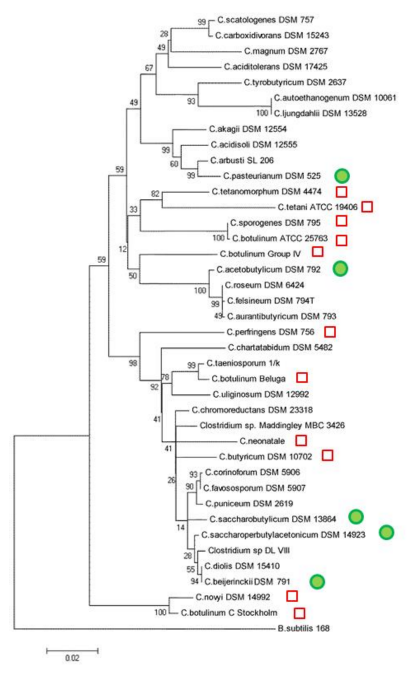
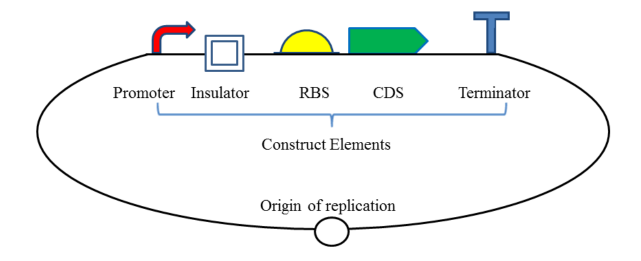


Figure 1. Phylogenetic tree of 16S sequences from selected organisms from the genus Clostridium. The tree was built using Maximum Likelihood method based on the Tamura-Nei model 11 . The bootstrap consensus tree inferred from 1000 replicates 12 is taken to represent the evolutionary history of the taxa analyzed 12 . Evolutionary analyses were conducted in MEGA5 13 . Green circles denote butanol-producing species used in industrial biotechnology (IB), red squares mark risk group 2 species 14 (risk group 3 being highest risk).

A recent comparative genomics study of many industrial saccharolytic strains (those preferring sugar as a carbon source) revealed that the known solventogens fall within two sister clades: one exemplified by *C. acetobutylicum* and one by *C. beijerinckii*¹⁵ (Figure 1). Interestingly, another comparative genomics paper which included more genus-wide species but fewer industrial solventogens supported the split, with the genus' type species *Clostridium butyricum* being more closely related to *C. beijerinckii* and the pathogen *Clostridium tetani* clustering closer to *C. acetobutylicum*¹⁶. Altogether, these findings serve to re-iterate that complex traits within the Clostridia such as pathogenicity are paraphyletic (also see Fig. 1). On the other hand, solventogenesis (of ethanol, butanol or acetone) may be very widespread in the genus, but there has not yet been a definitive comparative study reporting the extent of its conservation to our knowledge, and species and strains certainly vary in their productivity¹⁷. Topics of engineering interest have included improved characteristics such as solvent production^{18,19}, sugar utilisation^{20,21}, growth on alternative feedstocks such as lignocellulose^{22–25} and the production of alternative products^{26–28}.

A shortage of biological parts for the solventogenic Clostridia

To manipulate the solventogenic Clostridia using rational engineering, the development of reliable standard genetic parts for use in synthetic biology is essential. The assembly of these parts into synthetic gene cassettes and larger gene clusters means that they must work together in a consistent and predictable manner to become a useful tool for strain engineering. The basic parts of synthetic biology are the minimal sequence elements with biological function in gene expression (Figure 2), including promoters, ribosome binding sites, transcriptional terminators and other factors in the mRNA that affect stability. Also, synthetic biology tools rely on gene reporters to measure levels of gene expression, and these elements are often combined into plasmids which need their own origins of replication. Work in *Escherichia coli, Saccharomyces cerevisiae* and other model organisms has been transformed in the past decade by the rapid development, characterization and standardization of parts. However, work in these organisms benefits from a legacy of biological knowledge that is not necessarily available when working in non-model organisms, making the engineering of the latter even more



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Figure 2. Features of genetic constructs depicted using symbols from the SBOL v^{29} . Examples of promoters, ribosomal binding sites, reporters, terminators and replicons are described in the main text.

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It is also important to note that, of the existing body of molecular biology research in this genus, the majority has been carried out in *C. acetobutylicum* ATCC 824. Work in this species has benefited from a relatively early development of a reliable electroporation protocol³⁰ and from the publication of the whole genome sequence in 2001, making it the first published clostridial genome³¹. However, *C. acetobutylicum* is only one of many industrially promising solventogenic clostridia. While the engineering of some of these species has historically been hindered by a lack of transformation protocols, these are now available for the transformation of the aforementioned five industrial species^{30,32}-³⁵. In several cases, development of transformation protocols has required the circumvention of restriction systems which degrade incorrectly methylated DNA. Indeed, electroporation of *C. acetobutylicum* only became viable with the expression of the B. subtilis phage $\phi 3T$ I methylase in the E. coli cloning host; this methylates the sequence GCNGC, which would otherwise be cleaved by the C. acetobutylicum Cac824I type II restriction enzyme³⁰. Likewise, transformation of *C. pasteurianum* ATCC 6013 (DSM 525) requires the methylation of CGCG sequences, which has been accomplished by the use of the M.FnuDII³³ or M.BepI³⁶ methylases. Other organisms have more complex restriction systems; in C. saccharobutylicum NCP 262, which has two type I restriction systems, expression of the methylation and specificity domains of these

systems on a plasmid in *E. coli* was sufficient to allow transformation by conjugation³⁴. Some developments have also been made in transforming non-type strains, which may have restriction patterns which differ from those of the type strains. For example, *C. pasteurianum* NRRL B-598 is part of the *C. beijerinckii* cluster¹⁵ but requires the use of a *dam-/dcm-* strain of *E. coli* for successful transformation, suggesting that the type IV system of this strain is particularly important³⁷; conversely, *C. beijerinckii* NCIMB 8052 can be transformed with much greater efficiency even with DNA from a *dam+/dcm+* host³⁸. While an analysis of such developments in the entire genus *Clostridium* would be beyond the scope of this review, recent publications by Pyne et al.³⁹ and Minton et al.⁴⁰ provide a comprehensive review of the development of *Clostridium* strains for genetic engineering. However, it is certain that the range of genetically tractable *Clostridium* species and strains will expand with future research.

The resurgent interest in solventogenic *Clostridium* species suggests that synthetic biology tools are needed and there are clear shortages of particular biological parts with characterized activity available for the engineering of these organisms. We believe this warrants a concerted effort to address the weaknesses in the toolbox. Furthermore, with the establishment of transformation protocols throughout the genus Clostridium, it is also important to consider the potential of adapting existing parts and tools for use in other, less well-developed solventogens. Here, we assess the currently available parts used in synthetic biology projects, starting first with reporters available to measure gene expression, then the promoters, terminators working at the level of transcription and then post-transcriptional factors including mRNA stability, ribosome binding sites, riboswitches and codon usage. Finally we review the replicons currently being used to assemble synthetic gene clusters and briefly provide an overview of methods to introduce these genetic elements onto the chromosome, which has been reviewed elsewhere⁴⁰. Overall, we aim to summarize and systematize the existing biological parts used in the industrial solventogenic *Clostridium* species in order to highlight areas where more research is required and accelerate progress in creating a full range of synthetic biology tools for manipulating these industrially important organisms.

Genetically-encoded reporter systems are the major *in vivo* gene expression measurement techniques available and are required to measure the activity of different parts being tested. The oxygen sensitivity of obligate anaerobes such as the clostridia limits the ease of use of many reporter systems, including some popular fluorescence-based and enzymatically-based reporters, and has led to the development of some novel reporters which are slowly being adopted.

Fluorescent Reporter Proteins

The use of fluorescent reporter proteins is now widespread in biology. Successful use requires the correct folding and maturation of the fluorescent protein to enable detection. The level of signal for the fluorescent protein must be sufficiently high to enable accurate detection as there is no signal amplification as seen in enzymatic reporters.

GFP-family fluorescent proteins

The highly engineered family of Green Fluorescent Proteins (GFPs) now includes variants with improved brightness and photostability and with a range of different colours (different excitation and emission maxima)⁴¹. The major limitation of the GFP-like proteins for anaerobes is the requirement of molecular oxygen for chromophore maturation and fluorescence⁴². This excludes the possibility of real-time gene expression monitoring; however, *in vivo* observations are still possible in some oxygen-tolerant clostridia such as *Clostridium perfringens*⁴³, where yellow fluorescent protein (YFP) was synthesized and fluorescence was developed after exposure of live cells to atmospheric oxygen⁴⁴.

Using a similar principle, it is possible to obtain a snapshot of the protein levels in whole cells by exposing fixed anaerobically grown bacteria to atmospheric O_2 in a process termed 'aerobic fluorescence recovery', enabled by the discovery that GFP-family proteins can undergo maturation in fixed cells. Two studies have demonstrated the technique in *C. difficile* using paraformaldehyde and glutaraldehyde to fix cells expressing codon-optimized variants of CFP (cyan)⁴⁵ and mCherry (red)⁴⁶. This approach works as the fixatives cross-link primary amines (especially lysine residues) which are absent from the GFP-family chromophores. Fixed cells offer an advantage to

the imaging of dying live cells as they more accurately represent normal protein localization (gene expression changes are also likely to occur in dying or metabolically stressed cells)⁴⁶. So far, the technique has not been used to systematically quantify gene expression but rather to label proteins and track their intracellular localization^{46,47}.

Flavin-binding Fluorescent proteins

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220 The flavin-binding fluorescent proteins (FbFPs) are a class of alternative fluorescent

reporters capable of maturation in anaerobic conditions. The FbFPs are small proteins

222 (~11-15kDa) that have oxygen-independent fluorescent properties, using a flavin-

223 mononucleotide (FMN) cofactor as the chromophore.

A pioneering study in 2007 reported the development of three anaerobic fluorescent

reporters derived from the LOV (Light-, Oxygen- or Voltage-sensing) domains of

bacterial proteins: BsFbFP derived from YtvA of Bacillus subtilis, PpFbFP from SB2 of

227 Pseudomonas putida plus an E. coli codon optimized variant of BsFbFP called EcFbFP⁴².

This domain was engineered to emit fluorescence by mutagenesis of a reactive cysteine

in the FMN-binding pocket⁴².

230 These proteins are now commercially available from Evocatal Gmbh under the

trademark name 'evoglow'. Currently listed variants marketed for use in Clostridium

species include Bs1 (monomeric BsFbFP), Bs2 (dimeric BsFbFP) and Pp1 (PpFbFP).

The reporters' functionality in *C. acetobutylicum* was demonstrated by Schulz in 2013;

the highest fluorescence levels were observed for a *C. acetobutylicum* codon-optimized

235 Pp1 (referred to in text as "Pp2"), followed by codon-optimized Bs2 (referred to as

"Bs3"). Interestingly the same constructs gave inverse results in *E. coli*⁴⁸. While Evocatal

237 GmbH offers C-Pp1 and C-Bs2 *Clostridium* codon-optimized reporters with publicly

available nucleotide sequences, it is not clear whether the nucleotide sequences

(reflective of the codon optimization approach) are the same as Pp2 and Bs3. A study in

240 2014 reported the placement of the evoglow Pp1 reporter downstream of the cipP

241 promoter to monitor growth of *Clostridium cellulolyticum* on cellulose⁴⁹. The

242 functionality of the evoglow Bs2 variant has also been demonstrated in C.

pasteurianum⁵⁰ and in Clostridium ljungdahlii⁵¹.

244 Another FbFP example, developed from the *Arabidopsis thaliana* LOV2 domain of the

blue-light receptor protein Phot2, is the improved LOV (iLOV) FbFP⁵². The iLOV FbFP

246 has been further modified for enhanced properties, generating variants such as

photostable iLOV 2.1 (phiLOV 2.1) and others^{53,54}. While phiLOV 2.1 FbFP has improved photostability, it is still substantially lower than that of GFP. A recent study demonstrated the utility of a *C. difficile* codon optimized phiLOV 2.1 in three *Clostridium* species (expression was driven from an engineered strong *xyl* promoter)⁵⁵. Under the test conditions, the three species exhibited varying levels of fold-increase of fluorescence over their background autofluorescence level: 3.2-fold increase in *C. difficile* R20291 (not a *Clostridium sensu stricto* species), 5.6-fold in the pathogenic toxin-producer *Clostridium sordellii* ATCC 9714 and 4.5-fold in the solventogen *C. acetobutylicum* ATCC 824⁵⁵. While the FbFP reporters will be detectable when expression is driven from very strong promoters, further improvements in brightness and photostability will make them more generally useful for multiple applications.

One of the drawbacks of the FbFPs is that all of the available variants are of the same colour; additionally, unfolding caused by translocation has been reported to result in loss of fluorescence due to the loss of the bound flavin cofactor⁵⁴. These limitations prevent the use of FbFPs as the sole reporters for multi-output circuits or secretion. Engineering of FbFPs for different emission spectra has proven to be somewhat complicated; while a simulation study of a theoretical iLOV mutant predicted a red shift increase⁵⁶, experimental results showed that the mutation resulted in an overall brightness decrease and a blue shift of emission⁵⁷. Research is underway to develop more fluorescent derivatives of the LOV domains and to explore the potential for diversifying their colour spectrum^{56,57}. The other limitation of FbFPs is their relative dimness, which can decrease the sensitivity of a reporter system; depending on the measurement instrument used, a promoter's expression level might be below the detection limit. Photostability improvements (as in phiLOV) have resulted in a dimmer mutant⁵³; for applications where a longer exposure time is crucial, the dimmer, stable protein is preferable, but for an automated single measurement application such as flow cytometry, the brighter variant may be more suitable. Overall, all fluorescent protein reporters provide the ability to quantify cell-to-cell heterogeneity of gene expression (when used in flow cytometry or microscopy), which can be very useful, and while there are still some limitations with FbFPs, they have the potential to be a route to reliable *in vivo* real-time expression monitoring in *Clostridium*.

Enzymatic Reporter Proteins

Enzymatic reporters catalyse a (preferably) unique reaction either *in vivo* or *in vitro* and the specific activity is calculated to estimate protein levels and thus gene expression. Enzymatic reporters require the addition of substrates and cofactors to the assay reaction mixture as well as the production of cell lysate. This often means that enzymatic assays involve more preparation steps than fluorescent proteins but can have adjustable sensitivity by varying substrate levels. Also, *in vitro* lysate assays are, by definition, bulk population measurements.

Chloramphenicol acetyltransferase

The first reporter which has been extensively used in the *Clostridium* genus, both *in vivo* and, more quantitatively, in vitro, is the chloramphenicol acetyltransferase (CAT) reporter (encoded by catP). The system was first developed for use in C. perfringens (a medically relevant non-solventogenic bacterium)⁵⁸ and has since been used in *C.* acetobutylicum⁵⁹. Chloramphenicol acetyltransferase is an enzyme that catalyzes the covalent attachment of an acetyl group from acetyl-CoA to chloramphenicol⁶⁰, and is the basis of the chloramphenicol resistance marker found in many bacterial vectors. Transfer of the acetyl group exposes the thiol group of CoA, allowing the progress of the reaction to be observed by addition of 5,5'-dithiobis-(2-nitrobenzoic acid) (DTNB, also known as Ellman's reagent); this compound reacts with the free thiol, releasing 5-thio-2nitrobenzoate, which can be detected by measurement of absorbance at 412 nm^{60, 61}. The assay relies on a continuous spectrophotometric rate determination to calculate specific activity. Potential drawbacks include endogenous activity in chloramphenicol resistant strains (which may be resolved by disruption of the resistance gene) as well as high levels of endogenous non-specific coenzyme A transferases⁶² (knockouts of which would be more laborious and would likely have growth and phenotypic effects) and the cost of the substrate acetyl-CoA.

Carbohydrate hydrolases: β -glucuronidase, β -galactosidase, amylase, endoglucanase

The CAT assay's drawbacks led to the adaptation of a classical $\it E.~coli$ reporter in $\it Clostridium$: the $\it \beta$ -galactosidase enzyme, encoded by $\it lacZ$ (the gene was derived from $\it Thermoanaerobacter~thermosulfurigenes)$ ⁶². To quantify enzyme activity spectrophotometrically, ortho-nitrophenyl- $\it \beta$ -galactoside (ONPG) is used in an $\it in~vitro$ assay and an increase in absorbance at 420 nm due to the release of ortho-nitrophenol is

measured. Similarly, the GUS reporter system, which utilizes β-glucuronidase (E. coli 314 315 gusA), has been used in *C. acetobutylicum* in a fluorimetric assay with a cell lysate⁶³. The 316 fluorimetric assay measures the release of 4-methylumbelliferone (4-MU) after cleavage of 4-MU-β-galactoside or 4-MU-β-glucuronide (by a β-galactosidase or β-glucoronidase, 317 respectively), 4-MU emits light at 460 nm when excited by 365 nm light⁶⁴. Both the β-318 galactosidase and β-glucuronidase reporter systems benefit from the commercially 319 320 available range of fluorometric, spectrophotometric and histochemical substrates and 321 kits, making them an improvement over the CAT assay, although they are still not 322 inexpensive. The endogenous *amyP* gene (encoding an amylase expressed during solventogenesis) 323 324 has been used in *C. acetobutylicum*⁶⁵ as a reporter to study the phenomenon of strain degeneration (loss of solventogenesis) which is often caused by loss of the pSOL1 325 megaplasmid on which amyP is located. A codon-optimized amylase (AmyEopt) has been 326 327 used successfully as a secreted reporter in C. difficile by addition of a zinc metalloprotease PPEP-1 signal sequence⁶⁶. It is noteworthy that the strain used in the 328 above study was not capable of degrading starch under laboratory conditions; use of 329 330 amylase as a reporter in amylolytic strains (such as *C. acetobutylicum* ATCC 824) may require knockout of endogenous amylases to increase signal to background ratio. 331 332 In C. beijerinckii, a secreted endoglucanase (eglA) was cloned from C. saccharobutylicum NCP 262 and used as a reporter for gene expression⁶⁷; assays used were agar plate and 333 334 cell lysate assays which measured substrate (carboxymethylcellulose, CMC) clearance 335 on plates and product (p-nitrophenol from cleavage of p-nitrophenyl cellobioside) accumulation in lysate, respectively. 336 337

Lipase and Alkaline phosphatase

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The lipase encoded by tliA, from Pseudomonas fluorescens SIK W1, has also been used as a reporter in *C. beijerinckii* NCIMB 8052⁶⁸. Lipases are enzymes which hydrolyse the ester bonds found in long-chain acylglycerols, releasing fatty acids⁶⁹. Activity can thus be assessed by an enzymatic assay measuring the cleavage of *p*-nitrophenyl-esters⁶⁹ such as *p*-nitrophenyl decanoate⁶⁸. However, this reporter was only useable in *C. beijerinckii* as this species has a very low endogenous lipase activity. As such, it may not be suitable in other species, such as *C. acetobutylicum*, which has been observed to show inducible lipase activity 70 .

A colorimetric alkaline phosphatase assay was developed and used in *C. difficile* in 2015 based on the *phoZ* gene product from *Enterococcus faecalis*⁷¹. Activity of this gene can be determined by a colorimetric assay with *p*-nitrophenyl phosphate as the substrate. Use in solventogenic clostridia could be limited by native phosphatase activity. In order to examine the suitability of *C. difficile* as a host for this reporter, BLAST analysis was used to screen for *phoZ* homologues, and activity towards 5-bromo-4-chloro-3-indolyl phosphate was tested⁷¹. While the four main industrial strains do not have a *phoZ* homologue, homologues can be found in other species such as *C. pasteurianum* (CLPA RS02340, with 29% identity to *phoZ*).

Luciferase

The luciferase (lucB) reporter was also used in C. acetobutylicum successfully, allowing luminescence detection⁷². However, it also requires oxygen, ATP and luciferin. Notably, the cells used in the luciferase assay were neither lysed nor fixed, but were live cells that were exposed to atmospheric conditions, washed, and kept on ice. This treatment could conceivably introduce changes in gene expression levels prior to measurement. This assay has the lowest background signal level but the requirement for live cell exposure to oxygen may introduce variability. A codon-optimized luciferase ($sLuc^{opt}$) was also successfully secreted in C. difficile using the aforementioned zinc metalloprotease signal peptide⁶⁶.

While there are many reporter choices available to clostridial researchers, we would argue that the multiplicity of reporters used has not helped ease the comparison of data obtained by different laboratories. Altogether, a single reporter has not been established as a community standard; given the drawbacks of each particular system, it is difficult to identify one standout reporter, although our hope would be that improved fluorescence reporters with increased brightness and photostability would be the most useful and enable single cell studies in live cells. Currently we would advocate a choice of more than one reporter, enzymatic and fluorescent, and their systematic and comparative use within the genus. Such an approach could provide insight into the sources of gene expression variability between strains and provide a starting point for future engineering. To achieve this seamlessly, it is advisable to explore translational fusions between reporters; examples from *E. coli* include a FRET pair YFP-FbFP fusion⁷³ and

379 Gemini (lacZ α -GFP) fusion⁷⁴. If successfully applied to the *Clostridium* species, such bi-380 functional reporters have the potential to become a single standard reporter. 381 Additionally, α -peptide complementation strategies using the *E. coli lacZ* gene in *B.* subtilis⁷⁵ and Mycobacterium smegmatis⁷⁶ have been reported but a similar approach in 382 383 Clostridia has not been attempted. Interestingly, the *T. thermosulfurigenes*-derived LacZ 384 protein appears to lack the α -complementation region⁷⁷ and is a dimer⁷⁸ (rather than a 385 tetramer like *E. coli* LacZ). 386 **Transcription** 387 Transcription is the first stage of gene expression and the main stage for regulation of gene expression. Most existing biological parts in *Clostridium* fall within the 388 389 transcription category and are mostly promoters (summarized in Table 1). In contrast, 390 the termination of transcription has been studied relatively little and we present some 391 analysis on the potential to study and improve the parts available for reliable 392 termination. 393 **Transcription initiation** 394 Promoters are DNA elements that are capable of driving transcription by RNA 395 polymerase (RNAP) of downstream regions. Core promoter architecture in bacteria 396 includes the -10 region or TATAAT box (Pribnow sequence), the -35 region and a spacer 397 (with an optimal length of 17bp in *E. coli*) between the two conserved regions⁷⁹. This 398 motif is recognized by the housekeeping sigma factor that provides sequence specificity 399 to RNAP (called RpoD in E. coli and SigA in B. subtilis, C. acetobutylicum and other 400 solventogens)^{80,81}. Both the consensus sequence and sigma factors bear very significant 401 similarities between Firmicutes and Escherichia. In addition, some *E. coli* promoters contain AT-rich UP elements (upstream of -35 402 403 region) that are responsible for recognition by the carboxy-terminal domain of the RNAP α -subunit⁸², an additional *E. coli* promoter feature is the 'extended -10' region⁸³ 404 405 (upstream of the -10 region and within the spacer). Interestingly, near-consensus 406 promoters (including ones with UP elements) have been found to be significantly more 407 common in Firmicutes (including Clostridia) than in other bacteria⁸⁴, a feature that was 408 not explicable through higher AT-content alone. In Clostridia additional sequences with 409 resemblances to UP elements (termed phased A-tracts) have been described⁸⁵ and a 410 conserved extended -10 region that differs from its *E. coli analogue* has also been 411 proposed⁸⁶. The majority of promoters used in the genetic engineering of clostridia have been identified from the transcriptional units of important metabolic genes and most have been shown to be able to drive strong constitutive expression of a gene of interest⁸⁷. The most commonly used constitutive promoter in *C. acetobutylicum* (Table 1) is that of the native thiolase gene (thlA)^{88,89}, which has been used for the expression of genes involved in production of solvents such as butanol⁹⁰ and isopropanol²⁶, the transcriptional regulator gene *tetR*⁹¹ (used a minimal promoter variant-*miniPthI*), and, in a modified form, for the expression of cellulosomal scaffoldins⁹² and glycoside hydrolases^{25,93}. The *C. acetobutylicum* thiolase enzyme catalyzes the condensation of acetoacetyl-CoA from acetyl-CoA which are central metabolic intermediates⁸⁸. The C. acetobutylicum thlA promoter (hereinafter referred to as thlA) is a sigA-dependent promoter (as evidenced by its near-consensus -35 and -10 regions⁸⁹); however, in its normally chromosomal context it is also the subject of regulation by the redox-sensing transcriptional repressor Rex94. Rex inactivation was found to increase native thlA activity in *C. acetobutylicum* about 12-fold⁹⁵. The Rex-binding site has been omitted from the core promoter in commonly used synthetic constructs⁹⁶; yet this binding site is conserved in the promoters of orthologous genes from other solventogens (according to the RegPrecise database⁹⁷ and our promoter region alignments), suggesting that there are additional levels of control for this promoter that could be easily added by addition of the Rex sites if desirable. Indeed, some studies using similar promoters such as thlA and ptb have reported conflicting strength/activity findings, which may well be due to the cloning of regions of varying length (that likely include regulator binding sites such as Rex) and choosing to include the native RBSs or not (such as the ones reported between promoters in the Schulz 2013⁴⁸ and Girbal 2003⁶³ studies). Thiolase promoters have also been used for expression in other clostridia such as *C.* beijerinckii98 and C. pasteurianum99. However, gene expression data (RNAseq from C. beijerinckii) indicates that there are other genes that have higher expression levels than the ones currently used as promoter sources; many of those genes encoded are hypothetical proteins¹⁰⁰. Recently, Yang et al., 2017 constructed a sequence logo of 18 promoters previously identified in *C. acetobutylicum* which revealed a strongly conserved -10 region. Using degenerate oligos to mutagenize the core *thlA* promoter

elements, the authors were able to generate a promoter library of variable strengths,

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including mutations that increased the strength of the promoter, suggesting that *thlA* can be improved further using synthetic biology approaches.

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The use of a constitutive promoter may not always be desirable; it may be preferable to use an inducible promoter, allowing controlled expression of a gene of interest. So far, the only naturally inducible promoters exemplified in a solventogenic clostridium are the Staphylococcus xylosus xylA promoter, which is repressed by XylR in the absence of D-xylose⁶³, and the *C. perfringens bgaL* promoter, which is repressed by BglR in the absence of lactose¹⁰¹. However, constitutive promoters can be made inducible by addition of operator sites (for transcriptional repressors) or by the addition of binding sites for activators, and several such promoters have been developed for use in solventogenic clostridia. LacI-repressible versions of thiolase and ferredoxin promoters have allowed the construction of clostridial expression constructs for genes where expression in *E. coli* would be toxic^{24,25,93}. In *C. acetobutylicum*, the *fac* promoter (Clostridium pasteurianum ferredoxin promoter with lac operator) is able to function as an IPTG-inducible system in the presence of LacI, allowing approximately 10-fold induction⁵⁹. A TetR-repressed, anhydrotetracycline-inducible promoter has also been used in *C. acetobutylicum*, generated by the fusion of the chloramphenicol acetyltransferase promoter pcm with the tetracycline operator $tetO^{91}$. This promoter could achieve up to 313-fold induction, although high levels of anhydrotetracycline were inhibitory to growth. However, operators are known to influence the basal activity of the promoter¹⁰². In fact, a recent study in *E. coli* found the core RpoD promoter too sensitive to sequence context and operator insertions to be a suitable target for forward engineering efforts and turned to extracellular sigma factors (ECFs) with T7 RNAP whose promoter core sequences they found to be more insensitive to operator addition¹⁰³. An alternative strategy - CRISPR-mediated repression of transcription - has been demonstrated in several solventogenic species, namely C. acetobutylicum¹⁰⁴, C. beijerinckii⁹⁸ and *C. pasteurianum*⁵⁰,

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Orthogonal expression systems, i.e. promoters that require other non-native elements for activity (commonly an alternative sigma factor or phage polymerase), first found widespread use in the cloning of genes with toxic products. This approach allows the total repression of genes until they are introduced into the organism of interest. A

commonly used example in *E. coli* is the T7 promoter, which requires the phage T7 polymerase for activity¹⁰⁵. An example from Clostridia is expression driven from the *C.* difficile tcdB promoter that is dependent on the native sigma factor TcdR. The tcdB promoter is highly active in *C. acetobutylicum*, but only when *tcdR* has been integrated into the genome⁵⁹. This enabled the high-level expression of a *mariner* transposon in *C.* acetobutylicum without negative effects on the *E. coli* cloning host. There is a strong case for designing these orthologonal-type systems as simply using a clostridial housekeeping promoter (such as the *thlA* promoter) with a clostridial ribosome binding site would not provide orthogonality in commonly used cloning hosts such as *E. coli* or *B.* subtilis due to the degree of conservation between the transcription and translation initiation systems amongst these organisms; on the contrary, as a rule strong clostridial promoters and RBSs (see translation initiation section) often retain their strength in *E. coli* whereas the opposite is observed more rarely, this is likely due to the on average higher similarity to the Bacteria-wide consensus translation and transcription initiation signals in *Clostridium*^{86,106}. Interestingly, AT-rich DNA can be toxic to *E. coli* (clostridial genomes are very AT-rich) due to transcriptional activity from spurious intragenic promoters and RNAP titration¹⁰⁷. This is exacerbated by the fact that the *E. coli* extended -10 region, which is sufficient alone for transcription initiation in *E. coli*, could also be present by chance in Clostridium-derived sequences, increasing the likelihood of spurious and unpredictable transcription during cloning in *E. coli*.

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Promoter	Comments			
Constitutive				
thlA (C. acetobutylicum)	Widely used for constitutive gene expression in C. acetobutylicum for example expression on C. beijerinckii ald, C. ljungdahlii bdh90,			
	C. acetobutylicum adc, ctfA, ctfB, 26 & tetR (miniPthl variant) 91 . Activity analyzed using GusA 63 and several FbFP 48 reporters			
thl (C. perfringens)	Used for expression of srtA genes from C. acetobutylicum, L. monocytogenes, and B. cereus25.			
fdx (C. sporogenes)	Activity analyzed using CatP reporter ⁵⁹ . Used in the ClosTron system for expression of the Ll.LtrB intron ¹⁰⁸ . Used for expression of			
	$spoA$ integrated into the chromosome for complementation of a $spoA$ mutant 109 .			
Controlled				
thl (thl with double lac operator)	lac -repressed version of $thlA$. Used to express a miniscaffold in $cipc1^{92}$ & weakened version for expression of mannanase $man5K^{93}$.			
thlOid (thl with single lac operator)	A lac -repressible version, used to drive chromosomal expression of the $\it C. cellulolyticum$ glycoside hydrolase $\it xyn10A^{25}$.			
fac (single lac operator, derived from C .	Activity analyzed using CatP reporter ^{59,110} and Pp2 FbFP ⁴⁸ . Formerly used in the ClosTron system for expression of the Ll.LtrB			
pasteurianum)	intron ¹¹⁰ . Used for expression of $codA$ in the C . $acetobutylicum$ knockout vector pMTL-SC7515 ¹⁰⁹ .			
fdxOid (fdx with single lac operator)	Used for chromosomal expression of $\it C.~cellulolyticum~glycoside~hydrolase~\it cel9G^{25}$. Has RBS from $\it C.~acetobutylicum~thlA~promoter$.			
facOid (fac with single lac operator)	Used for chromosomal expression of $\it C.~cellulolyticum~glycoside~hydrolase~\it cel48F^{25}$. Has RBS from $\it C.~acetobutylicum~thlA~promoter$.			
tcdB (C. difficile)	Developed for an orthogonal expression system, requires the exogenous sigma factor TcdR for function ⁵⁹ .			
xylA (S. xylosus)	Repressed by XylR and inducible by D-xylose; assessed with β -glucuronidase reporter giving 17-fold induction with D-xylose 63 .			
pcm (tet01-containing variants)	Repressed by TetR; inducible in the presence of anhydrotetracycline ⁹¹ .			
bgaL (C. perfringens)	Repressed by BgaR and inducible by lactose ¹⁰¹ .			
xyl/tet0 (B. subtilis)	Repressed by TetR ¹¹¹ ; activity in <i>C. acetobutylicum</i> shown by phiLOV 2.1 Opt FbFP expression ⁵⁵			
Fermentation phase-specific				
adc (C. acetobutylicum)	Activity assessed using β -glucuronidase ⁶³ , β -galactosidase ⁷² and Pp2 FbFP ⁴⁸ . Primarily active after onset of solventogenesis.			
ptb (C. acetobutylicum)	Activity assessed using β -glucuronidase ⁶³ , luciferase ⁷² and Pp2 FbFP ⁴⁸ . Active during acidogenesis (not solventogenesis)			
ptb (C. beijerinckii)	Activity assessed through expression of $lacI^{59}$.			
sol (C. acetobutylicum)	Activity assessed using β -galactosidase $\&$ luciferase reporters 72 . Weak expression, primarily active during late exponential phase.			
bdhA (C. acetobutylicum)	Analyzed using β -galactosidase reporter 72 . Comparatively weak expression, primarily active in early exponential phase.			
bdhB (C. acetobutylicum)	Analyzed using β -galactosidase reporter 72 . Primarily active until onset of solventogenesis.			
hydA (C. acetobutylicum)	Activity analyzed using β -glucuronidase ⁶³ and Pp2 FbFP ⁴⁸ . High activity during acidogenesis, decreases to low after phase shift			
adhe2 (C. acetobutylicum)	Activity analyzed using Pp2 FbFP ⁴⁸ . Strong expression during solventogenesis ^{48,112}			

Transcription termination

Bacteria have two distinct mechanisms that function in transcription termination. Both types of terminators are usually located in the 3' end of transcriptional units. Rhodependent terminators rely on the Rho protein that recognizes a target sequence and causes RNA polymerase to fall off of the template DNA¹¹³. The specific DNA recognition sites, *rut* sites, have been used before in genetic circuits¹¹⁴, but not extensively, perhaps due to the relatively poor mechanistic understanding of the process¹¹⁵. The second mechanism, which also has been reported to be more widespread¹¹⁶, is referred to as Rho-independent or intrinsic termination and relies on the folding of a short GC rich hairpin followed by a poly-U transcribed sequence. The hairpin folding and subsequent transcription of the U-tract causes the polymerase to pause and leads to transcript and RNAP dissociation¹¹⁷. This is also facilitated by the fact that ribo-uracil-deoxyribo-adenine is the weakest nucleotide base pair¹¹⁸. Intrinsic terminators are often found downstream of operons; however, they are also involved in transcription attenuation when present within coding regions or downstream of promoters¹¹⁹.

While a number of terminators have been used in the construction of clostridial expression constructs, often derived from clostridial genes such as $adc^{63,120}$, fdx^{109} , CD0164¹⁰⁹ and *thlA*^{90,121}, there have been few published analyses of terminator strengths in clostridia. The *C. pasteurianum fdx* terminator has been shown to be highly effective at preventing read-through inhibition of the replicon from the fac promoter in a clostridial vector⁵⁹, and screening of a selection of terminators in *C. acetobutylicum*²⁵ showed that the E. coli rrnB terminator T1 loop was able to function as an efficient terminator, reducing expression of a downstream gene. However, these analyses only took into consideration the effect of the terminator on a downstream target under the influence of a single promoter. As previously mentioned, the introduction of a terminator may have an effect on expression of an upstream gene by influencing mRNA stability, which is not a desirable feature of a standard part¹²². However, this effect can be quantified in an appropriately designed assay123. Furthermore, the efficiency of termination may increase or decrease depending on promoter activity; this has been recently exploited for the development of a genetic band-pass filter in E. coli¹²⁴. Terminator strength has also been shown to be influenced by the hairpin's proximity to a stop codon or when present within a coding region¹²⁵. A recent study demonstrated

the gradual increase in termination efficiency as distance between the stop codon and hairpin increased¹²⁶. The ribosome was found to repress transcriptional termination when the stop codon and hairpin were in close proximity – termination repression was relieved when insulator sequences of approximately 30 bp were used¹²⁶. Expanding the range of characterized clostridial terminators is clearly necessary for the development of a clostridial synthetic biology toolbox.

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One approach that can be undertaken is the use of algorithms to extract putative terminator sequences from genomes, also known as 'part mining'. Several such bioinformatics tools exist; they rely on seed sequences, secondary RNA structure features or both^{116,127-129}. These bioinformatics tools were developed to aid genome annotations by identifying the 3' ends of operons and may not then detect all the features required by a functional terminator. In order to evaluate the relative difference in sequences retrieved by different programs, we compared the output of three searches performed on the *C. acetobutylicum* ATCC 824 genome (Figure 3A). The three search algorithms that were chosen are as follows: TransTermHP (TTHP)¹²⁸, RNIE¹¹⁶ and WebGeSTer (WG)¹³⁰. TTHP is a widely used tool, while RNIE's authors state that their tool eliminates false positives in comparison to TTHP¹¹⁶. Finally, WebGeSTer was selected because the authors classify the results into different types of intrinsic terminators based on overall secondary structure. They discovered that canonical Utract containing intrinsic terminators (termed 'L-shaped') form the majority of structures found within Firmicutes, whereas they are a minority in the E. coli model where they were first identified¹²⁹.

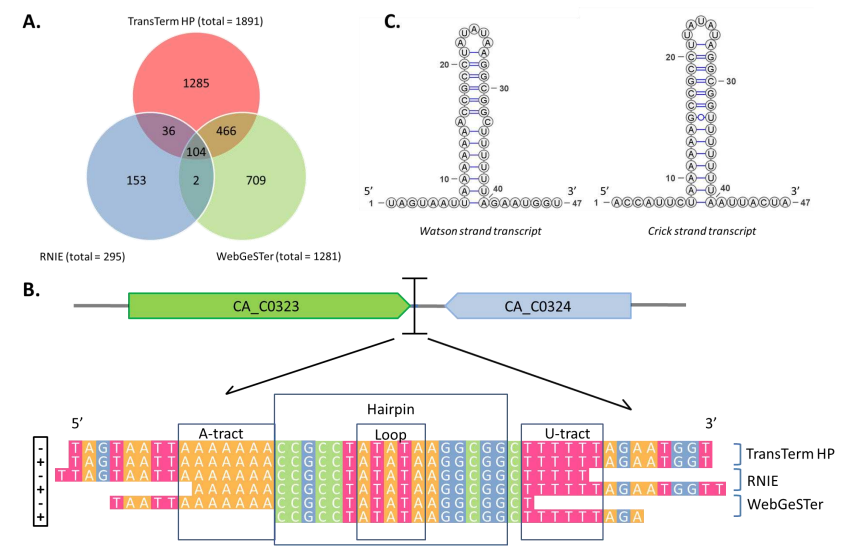


Figure 3. Application of terminator prediction software to C. acetobutylicum. A. Outputs from predicted terminator sets derived from the C. acetobutylicum genome by application of TransTermHP (TTHP), RNIE and WedGeSTer were compared. B. An alignment of predictions of a putative bi-directional terminator (marked by T sign) by all three algorithms (note all three produce separate predictions for plus (+) and minus (-))C. RNA structures predicted by RNAfold^{215,216} for the TransTermHP sequence, drawn with VARNA²¹⁷; it is worth noting that the base-pairing between the A-tract and U-tract may not form until after termination²¹⁸. Terminator lengths: WG: maximum length (max)-48bp, minimum (min) - 28bp, average (avg) - 35.8bp; RNIE: max-45bp, min-34bp, avg-41.7bp, TTHP: max-89bp, min-41bp, avg-57.2bp.

In our analysis, each algorithm was used to produce sets of putative terminator predictions, which were examined to identify overlapping predictions and are presented in Fig. 3A. Using the default settings of the different algorithms, the total number of predictions per program varied considerably, with TTHP and WG each predicting over 1000 terminators while RNIE predicted fewer than 300 (Fig. 3A). Despite this much smaller set, only about half of the predictions were shared with either TTHP and WG sets or both, producing a small set of 104 terminators shared by all three methods, an example of which is illustrated in Fig. 3B, an intergenic region separating two converging coding sequences, CA C0323 (a sensory transduction histidine kinase) and CA C0324 (a tetratricopeptide-repeats-containing protein). This also illustrates a class of terminators with particular utility in synthetic biology, namely bi-directional terminators, which are capable of terminating transcription from both the plus and minus strands. Our analysis finds that 37 bi-directional terminators are supported by all three algorithms, supporting the concepts that there are plentiful targets in clostridial genomes for 'part mining' and future experimental characterisation of these likely strong terminators. An interesting feature of the bi-directional terminators is their hypersymmetry¹³¹ (can be seen in Fig. 3B). This leads to the possibility to form an extended hairpin between the so-called A-tract (can act as U-tract in the reverse transcriptional orientation) and the U-tract (as shown in Fig. 3C). However, as pausing and termination occur at the U-tract¹¹⁷, the formation of an extended hairpin may not be relevant and does not appear to contribute to strength in all terminators¹³². Also, note that in this example the terminator on the minus strand (also referred to as Crick strand) is predicted to have a more stable hairpin (rG and rU form a basepair while rA and rC do not), whereas terminators from both strands are capable of forming an extended hairpin if fully transcribed as a result of base pairing between the A- and Utracts. Combined with having a longer U-tract (Fig. 3C), the Crick strand terminator is possibly the stronger one, interestingly WebGeSTer predicts a smaller terminator downstream of the one depicted in Fig. 3A (not shown). A previous analysis of the C. acetobutylicum genome for terminators using TransTerm¹²⁷, a predecessor of TTHP, revealed the presence of 66 putative bi-directional terminators¹³³, supporting the rationale of combining outputs from multiple algorithms to create a small part list for experimental characterisation.

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Translation

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Translation initiation

Bacterial ribosome-binding sites (RBS) are short sequences located in the 5' 590 untranslated region of messenger RNA (mRNA) transcripts, consisting of a Shine-591 592 Dalgarno sequence (SD), polynucleotide spacer, and a translation initiation codon¹³⁴. 593 Commonly, native promoter-RBS combinations have been used (such as in the 594 pMTL80000 vectors). An alternative strategy is to use a native RBS (such as that of the C. acetobutylicum thlA gene) fused to a new promoter, as in a study from 201625 that 595 596 generated several new hybrid promoters. Others have experimented with the length of the spacer^{135,136}. Yet, in contrast to model organisms (*E. coli* and even *B. subtilis*), there 597 are few published comparisons of modified RBSs for use in solventogenic clostridia. 598 599 Hence the generation and screening of synthetic RBS libraries could be a promising 600 route for optimising the expression of synthetic gene constructs in clostridia. 601 SD sequences provide sequence complementarity for the 3' terminus of the 16S rRNA (known as the anti-Shine-Dalgarno or aSD¹³⁷) which acts as a guide for the ribosome 602 complex enabling mRNA recognition by the translation machinery and translation 603 604 initiation. While most of the knowledge on bacterial translation initiation comes from E. coli work, early studies indicated that the Firmicute B. subtilis requires a longer 605 606 complementary region between the SD and the 16S rRNA to achieve comparable expression levels^{138,139}. An early study on translation initiation revealed that Firmicutes 607 608 have, on average, a higher complementarity of the predicted SD region to the 16S rRNA 609 3' terminus than *E. coli* does¹⁰⁶. Recently, a systematic analysis of SD-aSD pairings in *B.* subtilis and in E. coli confirmed this trend¹⁴⁰. Replicating these studies in solventogens 610 would provide useful information. 611

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The spacer is the mRNA region between the SD and the start codon. The aligned spacer (which is the distance between the start codon and the 5' end of the aSD, base-paired to a SD^{137,141,142} as revealed by sequence alignment) is of particular importance for translation initiation¹³⁷. Defining the length of the aligned spacer precisely is difficult, as interpretations of the extent of the aSD region (starting from the 3' 16S rRNA terminus) vary. The 3' end of the 16S rRNA gene (aSDs) of *B. subtilis* is identical to that of *C. perfringens*¹⁴³ (as well as identical to those of solventogenic clostridia, based on our sequence analysis of published genomes) but to our knowledge there is no reported

experimental validation of the clostridial mature 16S rRNA 3' ends in vivo. Spacers in different species may have different optimal lengths; for example, spacers of *Pyrococcus* abyssi are, on average, roughly 3 nucleotides longer than those of E. coli¹⁴⁴, whereas the spacers in Bifidobacterium longum145 would be considered shorter. According to our definition of the putative clostridial aSD (5' GAUCACCUCCUUUCU 3'), in C. acetobutylicum, the native RBS of the thlA promoter has an 'aligned spacer' of 4 bases. The effect of altering the length of the thlA spacer was recently investigated in two studies conducted in *C. acetobutylicum*^{135,136}. Interestingly Yang *et al.*, 2016 showed that a lengthened thlA RBS spacer with an extra 6 nucleotides (to a total of 14 bases- aligned spacer of 10bp), encoding a Sall restriction site, did not significantly alter reporter expression in comparison to the WT thlA RBS. Shortening the spacer below the WT length resulted in a decrease in expression, while further increases over an aligned spacer length of 10 bases (by the addition of a XbaI site) removed almost all of the expression¹³⁵. Yang et al. (2016) demonstrated the potential benefits of utilising modified spacers in C. acetobutylicum135. Overexpression of the biotin synthesis genes bioY, bioA, and bioB was observed to provide an improvement in growth phenotype and solvent production. The thlA promoter was used to drive expression of bioY, bioD and bioA; replacement of the thlA RBS spacer with a shortened, less 'effective' variant resulted in a further improvement in growth characteristics. These results demonstrate not only the ability of modified RBS to optimize expression of synthetic pathways, but also the necessity of considering the effects of any alterations to the spacer, e.g. the introduction of restriction sites, when generating synthetic constructs. In 2017, Yang et al. generated a library of spacers (a BamHI site preceded the start codon in all cases) by starting with an aligned spacer of 2 bases and progressively increasing its length by 2 bases. After testing 11 variants the authors found that an aligned spacer of 4 bases gave the strongest levels of expression followed by 8, 6 and 10 bases in that order. It is conceivable that not only the length but also the sequence of the spacer might influence translation initiation, for example the introduction of a secondary SD within the primary SD's spacer region. Computational tools to design RBSs exist, such as the RBS calculator, which use biophysical models of RBS recognition and translation initiation, including RNA basepairing between the aSD and SD, spacer length and messenger secondary

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structure^{146,147}. There are reports that *de novo* design produces more accurate results than translation initiation rate prediction of natural sequences for Gram positives¹⁴⁸.

Codon usage during translation

While codon usage is not formally a 'biological part', it is an important feature of coding sequences, the differential frequency of synonymous codons amongst genomes, referred to as codon usage bias^{149,150}, has been shown to strongly influence heterologous protein expression levels^{151,152}.

High genomic AT-content is characteristic of the Firmicutes and is reflected in the nucleotide composition of coding sequences. The *Clostridium* species' codon usage differs from that of other Firmicutes as well as the Proteobacterium *E. coli*¹⁵³ and there are also bioinformatically observable variations within the *Clostridium* genus itself¹⁵⁴ but the significance of the latter in influencing gene expression has not been experimentally verified to our knowledge. Genetically encoded reporters have been used heterologously within the Firmicutes phylum without codon-optimization, for example *Staphylococci* have been sources of reporters and antibiotic resistance genes for Clostridia. On several occasions researchers have successfully used native reporter genes from *E. coli* (*gusA*)⁶³, *T. thermosulfurigenes* (*lacZ*)⁷² and the firefly *Photinus pyralis* (*lucB*)⁷² (we describe these reporters in more detail in the Enzymatic Reporter Protein section). Codon-optimized genes for *C. difficile* have been used in *C. acetobutylicum* ⁵⁵, as well as bespoke *C. acetobutylicum* codon-optimized ones^{24,48}. Researchers have also codon-optimized several GFP-like proteins for use in Firmicutes¹⁵⁵ (with *B. subtilis* in mind) but these have not been used in solventogens to our knowledge.

Codon optimization is not a trivial problem and codon optimization strategies vary considerably. The codon-adaptation index (CAI)¹⁵⁰ has been the historical measure of codon usage bias in an organism while there are others such as the codon bias index and the effective number of codons¹⁵⁶. Interestingly, simply improving a heterologous gene's CAI (making it more like a native gene or a highly expressed native gene) has not been found to correlate with expression levels^{157,158}. Efforts have been made to improve the indices describing codon usage bias and translational efficiency (which codon usage bias is thought to reflect) by studying endogenous gene expression^{159,160}. In addition, condition-specific usage tables have also been reported¹⁶¹. In an alternative strategy, Welch and colleagues developed genetic algorithms to select partial least squares

regression models which revealed that codons predominantly read by tRNAs that are most highly charged during amino acid starvation were good predictors of expression levels. Based on these results the researchers developed proprietary codon optimization algorithms to maximize protein expression^{158,162} which allowed them to predict expression levels in *E. coli*.

While it is clear that there is room for improvement in the heterologous protein expression strategies used in *Clostridium* solventogens, codon optimization strategies themselves are still being developed and the underlying principles are not yet fully understood; progress and existing approaches in the field have been reviewed elsewhere ¹⁶³. A good starting point is for researchers to report the details of the codon optimization strategy undertaken when publishing work containing codon-optimized genes. This way data from heterologous protein expression in solventogens can be compared more reliably.

Post-transcriptional control of gene expression

Tuning gene expression levels in *Clostridium* species has been achieved using control at the RNA level – by either influencing translation or changing RNA degradation rates.

Controlling mRNA stability

Another potential avenue for optimisation of expression levels is the adjustment of mRNA stability. Altering the stability of an mRNA transcript influences the number of transcripts in the cell, thereby affecting the overall rate of translation. In bacteria, a number of factors are associated with mRNA stability, such as secondary structures, RNase recognition sites and polyadenylation, amongst others¹²². The presence of secondary structures at the 5' end of the mRNA has been observed to provide an increase in mRNA stability¹²². In *E. coli*, these structures prevent the binding of RNase E, an endonuclease which binds single-stranded RNA at the 5' end and then scans for cleavage sites. This property has been exploited in *E. coli* by the generation of libraries of synthetic hairpins for introduction into the 5' untranslated region (UTR)¹⁶⁴. While C. acetobutylicum has an RNase E homolog, RNase E/G, it is not certain if this behaves in the same way as *E. coli* RNase E, due to having a different domain organisation 165. Nevertheless, *C. acetobutylicum* also has a homolog of RNase Y¹⁶⁶, which fulfils the role of RNase E in B. subtilis¹⁶⁷. Correspondingly, the utility of 5' hairpins for protection of mRNA has been demonstrated in clostridia; the introduction of 5' stem-loop sequences was confirmed to increase mRNA stability, reporter expression, and expression of the

genes *adhE1* and *adhE2* in both *C. acetobutylicum* and *C. beijerinckii*¹⁶⁸. This effect was much more pronounced during solventogenesis than during acidogenesis. Similarly, the introduction of a terminator hairpin in the 3' UTR can result in improved mRNA stability via inhibition of nuclease activity¹²². Although not fully explored, this principle has been demonstrated in solventogenic clostridia; the expression of a *cat* reporter gene in *C. acetobutylicum* was observed to increase by approximately 36% when the downstream *adc* terminator was replaced by a synthetic terminator, BBa_B1010, from the iGEM registry¹⁶⁸. Additionally, a terminator with activity in the reverse orientation prevents the formation of antisense transcripts which are known to reduce protein expression levels¹⁶⁹.

A completely opposite approach is to reduce mRNA stability by introducing RNase sites into the 3' UTR. This may be useful in the case of proteins which form inclusion bodies when overexpressed. In *E. coli*, sequences derived from the *cat* gene, which contains 28 RNase E sites, have been shown to reduce mRNA stability¹⁷⁰. When combined with two poorly soluble heterologous enzymes, the *cat*-derived 3' UTRs were shown to result in an increase in soluble protein, with concomitant increases in enzyme activity. The authors noted that this improvement could not be observed simply by using a weaker promoter; it was proposed that by limiting the stability of the mRNA, the number of proteins that could be produced from a single transcript was decreased, thus limiting the local concentrations of protein during translation.

Antisense RNAs and Riboswitches

Bacterial antisense RNAs (asRNAs) are transcripts with complementarity to another RNA. Several studies have used asRNAs were used to reduce native gene expression in *C. acetobutylicum*^{169,171} and in *C. pasteurianum*¹⁷², demonstrating that multiple routes to regulate gene expression are available in the solventogenic clostridia. Riboregulators are another class of naturally-occuring and generally trans-activating

asRNA elements that respond to a signal nucleic acid by Watson-Crick base pairing¹⁷³. They have defined sensor and effector domains and have been rationally designed to repress¹⁷⁴ and activate gene expression¹⁷⁵ in *E. coli*. Toehold switches (a synthetic cisregulatory subgroup of riboregulators) that activate gene expression in the presence of

cognate RNAs rely on sequestering the RBS and start codon¹⁷⁶. Like riboswitches, riboregulators are known to be present in *Clostridium* genomes.

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Riboswitches are RNA sequences that are able to bind to a soluble ligand, influencing the properties of the RNA. In nature, riboswitches typically contain a binding domain, or 'aptamer', and an 'expression platform' which mediates the effect¹⁷⁷. Binding leads to a change in conformation of the RNA, leading to formation of a secondary structure which can act as a terminator or an anti-terminator. While riboswitches are typically found in the 5' UTRs of mRNAs, some have been recently determined to control the expression of antisense RNAs or protein-sequestering small RNAs, while yet others have been shown to control access to recognition sequences such as RNase sites¹⁷⁸. The range of applications has been further developed by the creation of synthetic riboswitches. In bacteria, riboswitches have been developed that can influence translation initiation by inhibiting access to the RBS; ligand binding leads to a conformational change or even to self-cleavage, revealing the RBS and allowing translation¹⁷⁹. Riboswitches are found in all taxa and a number have been characterized in the solventogenic clostridia 180-182. However, riboswitches have not yet been used in the engineering of these organisms. As discussed earlier, only a limited range of inducible systems is available for the engineering of clostridia; the use of synthetic riboswitches could be a promising alternative for the creation of controlled promoters.

Replication

Plasmid origins of replication

Four replicons are in routine usage in solventogens: pCB102 (from *C. butyricum*), pBP1 (from *C. botulinum*), pCD6 (from *C. difficile*) and pIM13 (from *B. subtilis*)⁹⁶. The pIM13 replication origin is thought to replicate via rolling-circle replication¹⁸³ while there is evidence that pCD6 replicates in similar fashion to pIP404^{184,185} (which is either theta or strand-displacement). The replication mechanisms of pCB102 and pBP1 are unknown. In *C. acetobutylicum*, two other replicons that have been used are the pAMβ1 replicon and the pUB110 replicon 186; the pUB110 replicon was found to be somewhat more stable than pIM13, whereas the pAMß1 replicon was highly unstable. The *C. beijerinckii* filamentous phage CAK1's origin of replication has been used in *C. beijerinckii* strains¹⁸⁷. Additionally, the development of a replicon specific for *C. saccharoperbutylacetonicum* N1-4 was reported in 2007¹⁸⁸, this replicon is identical to the origin of the endogenous plasmid from *C. saccharoperbutylacetonicum* N1-504¹⁵ . A thermosensitive origin pWV01ts derived from *Lactococcus lactis cremoris*¹⁸⁹ has been shown to work in both *C.* ljungdahlii and C. acetobutylicum⁵¹. Segregation and transformation frequencies are available; however, more work is needed to determine copy number and compatibility groups.

Chromosomal integration

The integration of DNA into the genome, while not a 'part' in itself, is an important consideration for synthetic biology projects. Genomic integration has several advantages over plasmid-based expression strategies, including increased stability, removal of the requirement for antibiotic selection, and standardisation of copy number 190-192. However, there are other factors that must be considered when using chromosomal integration. One implication of the integration position is the copy number effect – genes closer to the origin have a higher copy number than ones near the terminus in exponentially dividing cells due to the mechanism of DNA replication. There is contradictory evidence as to the effects of chromosome location and levels of expression 193,194,195. The cause of some of the apparent transcriptional insulation of chromosomal genes (and thus promoter independence) was found to be due to the gene silencing activity of the HU-protein in *E. coli*194, a protein that is also present in

Firmicutes¹⁹⁶. Genomic engineering in *E. coli* has been carried out extensively; lambda-Red recombineering¹⁹⁷ is well-established, and the utility of the newly developed CRISPR technique has been demonstrated in this species 198,199. However, in the solventogenic clostridia, until recently published examples of genomic integration had been still somewhat limited. An early enabling technology was ClosTron which adapted the Ll.LtrB intron for use in *Clostridium*¹¹⁰. Another method for genomic integration is Allele-Coupled Exchange (ACE), as demonstrated in *C. acetobutylicum*^{24,25,190}. This is a homologous recombination-based method, where homology arms with different lengths are used to control the sequence of recombination events, and the second recombination leads to the generation of a selectable phenotype; currently, this involves either the truncation or repair of the *pyrE* gene or the activation of a promoterless antibiotic resistance gene by integration downstream of a strong chromosomal promoter such as thlA 190. Thus, one potential drawback to this method is that it only allows integration into a limited selection of loci. This drawback can be mitigated by the ability to carry out multiple rounds of iterative ACE, thereby making further genomic integrations into the same locus.

Nevertheless, many new developments have been made regarding the genetic manipulation of solventogenic clostridia. A variety of different allelic exchange-based strategies have been exemplified in *C. acetobutylicum*^{101,109,200,201} and *C. beijerinckii*²⁰²; while most of these studies have focused on the generation of in-frame deletions and subsequent complementation, Al-Hinai et al.¹⁰¹ demonstrate the integration of a heterologous gene through gene replacement. Furthermore, the generation of point mutations through recombineering has been demonstrated in *C. acetobutylicum*²⁰³, suggesting that the integration of DNA through this method may be feasible. Finally, mutant selection via CRISPR has been established in almost all of the main solventogenic species, with published examples in *C. beijerinckii*^{121,204}, *C. acetobutylicum*¹⁰⁴, *C. pasteurianum*²⁰⁵, and *C. saccharoperbutylacetonicum*²⁰⁶. Future developments are certain to expand the range of genomic modifications that can be made in these organisms.

Insulators

A biological part's adjacent sequences can have a profound effect on its behaviour compared to the sequence context in which it was characterized²⁰⁷. This poses a

fundamental challenge to the synthetic biology principle of part creation and characterization²⁰⁸. To counteract this issue in reproducibility genetic engineers have started utilizing a new class of parts called insulators²⁰⁹. Several strategies can be undertaken to insulate a part from its genetic context, and these can be split into two main categories: DNA-level insulators (such as simply using flanking buffer zones of sequence without a biological function or secondary structure) and RNA-level insulators. The latter includes post-transcriptional modification of RNA as well as the commonly used flanking double terminators¹²³ that prevent read-through transcription into synthetic gene constructs. Post-transcriptional insulators consist of inclusion of ribozyme-based insulators or using CRISPR-RNA-processing to decouple the 5' UTR from the coding sequence (CDS)^{208,210}.

Different parts require different types of insulation in order to achieve maximum reproducibility without compromising features such as strength.

Promoters used in synthetic biology projects are often minimal (-35 to -10 region) and without characterized transcriptional start sites; the inclusion of important functional elements such as the UP element has been previously recommended.²¹¹ Accordingly, the addition of an upstream and downstream insulating sequence has increased reproducibility in different genetic contexts²¹². It is worth noting that the strong *Clostridium* promoters that are in widespread use in the *Clostridium* community, *fdx* and *thlA*, are 200 and 150bp respectively. Even though this recombinant *thlA* is longer than a usual minimal promoter, a long 5'UTR contributes to this part's length (recombinant *thlA* is 59bp from its 5' end to the transcriptional start site), while the recombinant 5' end is slightly truncated to exclude a Rex NADH-dependent regulator binding site (as mentioned before). A study that reported the use of a minimal *thlA* promoter in *C. accetobutylicum*, *miniPthl* which has a truncated 5' UTR, did not test activity variation in different genetic contexts⁹¹.

Work in *E. coli* has demonstrated the utility of 'bicistronic RBS', where a leader peptide is translationally coupled with the CDS of interest, in improving reliability and context independence (downstream gene sequence) when a particular 5' UTR is combined with a new coding sequence²¹². An upstream RBS and start codon initiate translation of the leader peptide, the stop codon of which overlaps with the start of the downstream CDS, while the latter's RBS is positioned with the leader peptide. The ribosomes translating the leader peptide unfold the 5' UTR (of the downstream CDS), preventing it from

forming secondary structures with the mRNA of the downstream CDS and thus influencing translation. The same study employed standard transcriptional start site ("+1 promoter"): 5' UTR junctions to minimize (or insulate against) unforeseen effects of combining promoters with new 5' UTRs.

As mentioned above, the observation of ribosomal repression of transcription termination also necessitates the more widespread use of 'distance' insulators of a sequence without emergent function and secondary structure to separate the stop codon and the stem-loop hairpin¹²⁶. Such strategies are yet to be implemented in *Clostridium* engineering projects.

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Discussion

The development of biorenewables as an alternative to petroleum-derived commodity chemicals and fuels has resulted in the emergence of new markets²¹³. Increasing productivity, broadening the range of feedstocks, improving tolerance to solvents and by-products are all existing challenges to achieving higher sustainability and ensuring the economic viability of *Clostridium*-derived biorenewables²¹⁴. *Clostridium* species remain important hosts for the biological production of solvents and their further development relies on the adaptation of novel methodologies such as synthetic biology and metabolic engineering. Much progress has been made in the latter with several projects improving industrially relevant strains; however, the availability of biological parts with known behaviour is one of the limiting factors for the rate and scale of research. The reasons for our anticipation are twofold: first, the physical availability of biological parts streamlines assembly of genetic pathways, and second, knowledge about parts' behaviour is crucial in predicting and analyzing the behaviour of pathways and genetic circuits. We have shown in this review that there are multiple areas where improved knowledge of parts would provide a better toolkit for synthetic biologists using these organisms and consequently improve the rate at which industrial biotechnology and bioenergy process development can improve.

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