Rapid, Heuristic Discovery and Design of Promoter Collections in Non-Model Microbes for Industrial Applications.

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Abstract

Well-characterised promoter collections for synthetic biology applications are not always available in industrially relevant hosts. We developed a broadly applicable method for promoter identification in atypical microbial hosts that requires no a priori understanding of cis-regulatory element structure. This novel approach combines bioinformatic filtering with rapid empirical characterisation to expand the promoter toolkit, and uses machine learning to improve the understanding of the relationship between DNA sequence and function. Here, we apply the method in *Geobacillus thermoglucosidasius*, a thermophilic organism with high potential as a synthetic biology chassis for industrial applications. Bioinformatic screening of *G. kaustophilus*, *G. stearothermophilus*, *G. thermodenitrificans* and *G. thermoglucosidasius* resulted in the identification of 636 100 bp putative promoters, encompassing the genome-wide design space and lacking known transcription factor binding sites. 80 of these sequences were characterised in vivo and activities covered a 2-log range of predictable expression levels. 7 sequences were shown to function consistently regardless of the downstream coding sequence. Partition modelling identified sequence positions upstream of the canonical -35 and -10 consensus motifs that were predicted to strongly influence regulatory activity in *Geobacillus*, and Artificial Neural Network and Partial Least Squares regression models were derived to assess if there was a simple, forward, quantitative method for in silico prediction of promoter function. However, the models were insufficiently general to predict pre hoc promoter activity in vivo, most probably as a result of the relatively small size of the training data set as compared to the size of the modelled design space.
Visual Abstract
The predictable control of genetic modules or engineered metabolic pathways is a defining aspiration of synthetic biology\(^1\) requiring thoroughly characterised, robust genetic parts. Although synthetic biology parts and tools of increasing sophistication are available\(^2\)-\(^5\), the majority have been designed for use in a small number of model organisms\(^6\) and characterised only or mainly in these biological contexts\(^7\). Model organisms such as *Escherichia coli* or *Saccharomyces cerevisiae* are invaluable for laboratory-scale, proof-of-principle investigations and are used in some industrial applications\(^8\) but there is a real, practical need to expand the range of microbial chassis available for industrial applications that present more extreme environments for the biocatalyst\(^9\),\(^6\),\(^10\)-\(^13\).

Different control points affect the output of gene networks, including levels of transcription, translation, protein half-life and enzyme kinetics\(^14\). On a practical level, the use of promoters with varied and predictable activation and output characteristics ("strengths") are an essential feature of any synthetic biology toolkit\(^3\),\(^15\),\(^14\) and are particularly useful for balancing differential expression levels in "hard-wired", steady state genetic modules\(^16\). Promoter collections for synthetic biology applications should therefore cover a broad range of recombinant gene expression levels for nuanced tuning of synthetic pathways\(^17\), with individual promoters providing homogeneous, consistent and predictable outputs independently of the associated downstream coding sequence\(^18\).

Conventionally, promoters in atypical chassis may be isolated from upstream of genes or operons\(^15\) that are homologous to well-understood regions in model organisms, or identified using genomic or transcriptomic analyses of the host\(^7\) followed by in-depth characterisation in a range of genetic and environmental contexts. Alternatively, synthetic promoter libraries may be manufactured by mutagenesis of wild-type promoter sequences, again followed by deep analysis of novel activity\(^14\),\(^19\),\(^20\), though this approach tends to reduce, rather than enhance, promoter strength\(^9\),\(^21\)-\(^25\). Finally, recent advances in DNA synthesis have facilitated systematic approaches to promoter and regulatory sequence design by enabling the production and high-throughput screening of comprehensive sequence libraries\(^26\),\(^27\). Due to the scale of DNA synthesis required, however, this approach remains relatively expensive compared to mutagenesis and dependent on ready access to appropriate DNA synthesis facilities.
In this investigation, we used a bioinformatic approach to explore the promoter design space in *Geobacillus thermoglucosidasius*, a metabolically versatile\textsuperscript{11,28-30}, thermophilic microbe\textsuperscript{31} with high potential as a synthetic biology chassis for industrial applications\textsuperscript{6,32}. To date, engineering projects in *Geobacillus* have relied on 1 of 3 endogenous promoter sequences\textsuperscript{11,33,34}, the most widely used being the oxygen-dependent *ldhA* promoter\textsuperscript{9,11,31,35,36}. Mutagenesis-derived, synthetic promoters have also been reported for the genus\textsuperscript{9,37,38}, though their characterisation is limited to single genetic contexts.

Here, we selected 100 putative promoter sequences from the *Geobacillus* core genome encompassing the genome-wide design space and lacking known transcription factor binding sites. The sequences were synthesised, cloned upstream of 2 different reporter CDS and their activities assessed *in vivo*. This process was relatively rapid and resulted in a collection of 7 characterised promoter sequences that displayed a range of activities with low internal variance and that functioned independently of the downstream reporter sequence. Additionally, to better understand the relationship between promoter sequence and activity, the data from the *in vivo* characterisation were used to train and validate a variety of *in silico* models, including Random Forest partition, Artificial Neural Network (ANN) and Partial Least Squares regression (PLS).

The method presented here is broadly applicable to any potential bacterial chassis and could be used to expand synthetic biology tools for other biocatalysts and ultimately enhance our fundamental knowledge of genetic regulation in synthetic and natural systems.
Results & Discussion

Bioinformatic identification of putative promoters from the core genome of 4
Geobacillus species

Different Geobacillus species have the potential to be used as host organisms
for industrial bioproduction\(^6, 8, 33\). We therefore aimed to identify promoters that could
potentially be used across the entire genus. To obtain a suite of promoters that were
representative of the Geobacillus genus, we sequenced and assembled de novo the
genomes of 4 Geobacillus species that were available when the project started; G.
kaustophilus (DSM7263), G. stearothermophilus (DSM22), G. thermodenitrificans
(K1041) and G. thermoglucosidasius (DSM2542). To identify genes that were
common to all 4 Geobacillus species, single-copy coding sequences (CDS) were
clustered into homologous gene families using the GET_HOMOLOGUES software
package\(^39\). To increase calculation robustness, 3 separate clustering algorithms were
used, and the resulting gene families compared. Bidirectional best-hit (BDBH), COG
ingress and OrthoMCL (OMCL) algorithms returned 1,924, 1,914, and 1,902
CDS clusters respectively, with 1,886 homologous clusters being identified by all 3
algorithms (Figure 1A). The core genome of the selected Geobacillus species
therefore contained 1,886 CDS; i.e. a total of 7,544 homologous core CDS.

In prokaryotes, the majority of motifs that affect the initiation of both
transcription and translation occur in the 100 bp sequence window immediately
upstream of the CDS start codon\(^40, 41\). 100 bp sequences from immediately upstream
of the start codon of the 7,544 core CDS were therefore identified as putative
Geobacillus promoter sequences. BPROM software was subsequently used to
classify the 100 bp sequences as putative promoters based on the presence and
nucleotide composition of known conserved functional motifs\(^42\). To isolate sequences
that were likely orthogonal to endogenous regulatory pathways, putative promoters
were screened against BPROMs list of known Transcription Factor Binding Sites
(TFBS, Supporting Table 1), and sequences that contained any known TFBS were
discarded. A phylogeny of the 1,489 putative, generic sequences that remained after
screening was constructed as a representation of the Geobacillus promoter design
space (Figure 1B). Although BPROMs list of E. coli TFBS may not be exhaustively
representative of binding sites that are functional in Geobacillus, the lack of extensive
genus-specific TFBS characterisation in these non-model organisms renders a
genus-specific approach impractical. Given previous successfully applications of
BPROM software for promoter identification\textsuperscript{28}, the utilised list of TFBS was judged likely to provide an adequately generic reference for binding site recognition in \textit{Geobacillus}.

Multiple studies have used promoters isolated from the genomes of bacteriophage for the control of heterologous expression in \textit{E. coli}\textsuperscript{14}. Putative promoters were therefore also identified from the genomes of 2 bacteriophages, \textit{Thermus} phage Phi OH2 and \textit{Geobacillus} phage GBSV1, which were chosen due to their ready availability on the GenBank public database. Intergenic regions of at least 100 bp were identified in both genomes. From these intergenic regions, the 100 bp sequences immediately upstream of the start codon of the adjacent CDS were extracted. The extracted sequences were subsequently analysed using BPROM software to identify putative promoters, and any sequences that contained known TFBS were discarded. 9 putative promoters were identified from \textit{Thermus} phage Phi OH2, and 7 putative promoters were identified from \textit{Geobacillus} phage GBSV1.

In vivo characterisation of putative promoters

A number of studies have considered the effect of genetic context on promoter function in model organisms such as \textit{E. coli} and \textit{S. cerevisiae}\textsuperscript{18,41,43-45}. However, the drive for composable, modular regulatory elements in non-model systems is hindered by the fact that many studies still characterise the function of promoter sequences in a single genetic context. 2 previously published \textit{Geobacillus} synthetic promoter libraries, for example, used only GFP to characterise promoter performance\textsuperscript{9,37}. Putative promoters were therefore characterised upstream of both Dasher GFP and mOrange fluorescent reporters.

A trade-off was required between the desire to empirically explore large portions of the \textit{Geobacillus} promoter design space and the experimental feasibility of characterising large numbers of putative sequences in a host organism with low transformation efficiencies. The promoter phylogeny (Figure 1B) was therefore used to rationally select 100 putative promoters from across the \textit{Geobacillus} promoter design space for \textit{in vivo} characterisation using both reporters.

A sequence alignment of the 100 selected putative promoters revealed a heavily conserved purine-rich region located at the 3' terminus of the 100 bp sequence space (Supporting Figure 1). Given the similarities in both location and
nucleotide composition of the motif to the canonical Shine-Dalgrano sequence\textsuperscript{46}, this
region was identified as the RBS. We therefore changed the terminology, whereby
“promoter” refers to the complete 100 bp sequence, RBS refers to the 15 bp of
sequence at the 3’ terminus of the sequence space and Distal Regulatory Sequence
(DRS) refers to the sequence from -100 to -15 bp upstream of the start codon.

To facilitate potential future applications of the promoter sequences in which
disparate DRS and RBS might be required, the 100 selected putative promoters were
split \textit{in silico} into DRS and RBS parts that were subsequently flanked with type II
restriction cloning affixes (Supporting Table 2). \textit{In vitro} cloning of the DRS and RBS
parts resulted in the insertion of a 4 bp scar sequence at -19 to -16 bp upstream of
the start codon, increasing the length of the promoters to 104 bp. The inclusion of the
scar sequence was empirically shown to have no statistically significant effect on
promoter activity for 20 out of a set of 24 characterised sequences, with significant
alterations in regulatory activity hypothesised to be the result of extreme alterations
to mRNA secondary structure (Supporting Information, Supporting Figure 2).

Of the 100 selected putative \textit{Geobacillus} promoters, 5 \textit{promoter::GFP} and 9
\textit{promoter::mOrange} constructs could not be successfully synthesised. Furthermore,
11 \textit{promoter::mOrange} constructs could not be transformed into \textit{G. thermoglucosidasius}; 80 sequences were therefore characterised \textit{in vivo} upstream of
both reporters (Figure 2A). The characterised sequences covered a 148-fold range of
activity when characterised upstream of GFP, and a 107-fold range of activity when
characterised upstream of mOrange. 45 of the characterised promoters showed
expression levels for both reporter proteins that were not statistically significantly
greater than the negative control, \textit{G. thermoglucosidasius} transformed with the empty
pS797 vector. We therefore defined these 45 sequences as inactive. 19 out of the
100 screened promoters showed statistically significant activity with both reporters; 3
sequences were active with GFP only, and 13 sequences were active with mOrange
only (Figure 2B). A comparison of the codon usage of the 2 reporter proteins showed
them to be broadly comparable (Supporting Figure 3). The discrepancies in gene
expression between the 2 reporters were therefore assumed to be a result of
promoter activity, rather than differential codon utilisation.

To identify the promoters that functioned predictably and independently of the
downstream CDS, K-means clustering was used to group the characterised
sequences into 5 clusters based on their Euclidean distance from the line of
equivalence between GFP and mOrange activity, \( y = x \) (Figure 2C). No correlation in 

\textit{in vivo} activity between the two reporter proteins was observed for the majority of the 

classified sequences; clusters 2 and 4 contained promoters that resulted in 

stronger GFP expression than mOrange expression, whereas clusters 3 and 5 

resulted in stronger mOrange than GFP expression. Clustering identified 13 

promoters (cluster 1) with activity that fell close to the line of equivalence, of which 7 

displayed mean expression levels that were significantly greater than the negative 

control. The characterised \textit{Geobacillus} promoter library therefore contained 7 

functionally composable, active sequences, covering activity levels that were 

between 1.1 and 4.5 times greater than the \textit{G. thermodenitrificans ldhA} positive 

control.

Such functional composability of \textit{cis}-regulatory sequences is crucial if 

information regarding promoter performance derived from laboratory-scale 

characterisation experiments is to be applied to the systematic, scalable, bottom-up 

engineering of increasingly complex synthetic biological systems\textsuperscript{4,18}. The 

development of species-specific insulator mechanisms, that reduce the context-

specificity of regulatory parts through either molecular transcript processing\textsuperscript{47,48} or by 

physically separating genetic regulatory parts to disrupt context-specific mRNA 

secondary structures\textsuperscript{18,41}, is required if the majority of the identified promoters are to 

be used modularly in alternative contexts.

In addition to being functionally composable, promoter sequences for 

synthetic biology applications should ideally yield homogenous, predictable 

expression of the protein of interest at the single-cell level\textsuperscript{49}. Flow cytometry was 

therefore used to analyse the intra-population variation in fluorescence activity of the 

characterised \textit{promoter::reporter} fusions in transformed, clonal cultures. Compared to 

the positive control, the \textit{G. thermodenitrificans ldhA} promoter, 98% of the 

characterised \textit{promoter::GFP} fusions and 73% of the \textit{promoter::mOrange} fusions 

returned lower coefficients of variance, indicating that the majority of the 

characterised sequences offered more predictable regulation of protein expression 

than the current benchmark \textit{Geobacillus} promoter. Furthermore, the 7 promoters that 

functioned independently of coding sequence all returned lower coefficients of 

variation than the positive control \textit{ldhA} promoter (Figure 2D). Although 

subpopulations of cells expressing the reporters were apparent for 4 of the 

characterised promoters, the performance of these promoters was less variable and
therefore more predictable than that of the \textit{ldhA} promoter which has been widely used in studies with potential industrial applications\cite{9,11,31,35,36}.

Analysis of the genes with which the 80 characterised promoters were natively associated in their source genomes showed that the majority of the sequences homogeneously regulate basic cellular functions, and were therefore likely to be constitutive (Supporting Table 3). Cellular functions with which the promoters were natively associated included biosynthesis, cell membrane formation, catabolism, transcription and protein folding. However, 11 of the characterised promoters were natively associated with proteins relating to sporulation, and may therefore result in altered expression levels under sporulation conditions. The failure of the bioinformatic screening to identify and exclude these sequences highlights the limitations of applying bioinformatic tools that were developed in \textit{E. coli} in non-model organisms; as \textit{E. coli} is non-sporulating, a list of \textit{E. coli} TFBS will naturally not contain sporulation-specific TFBS.

\textit{Sequence-function modelling}

Mathematical models with the \textit{pre hoc} capability to determine promoter function could potentially reduce the need for \textit{in vivo} characterisation of large numbers of individual \textit{cis}-regulatory elements. Once a training set of sufficient robustness is established, regulatory elements of the desired strength for a given application could hypothetically be identified from the genome or designed \textit{de novo}, in a manner analogous to tools such as the RBS calculator\cite{3}. To better understand the basis of promoter function in \textit{Geobacillus}, and to assess if there was a simple, forward method for \textit{in silico} prediction of promoter function, statistical learning approaches were used to derive models of the design space.

We used a variety of techniques to mathematically describe the relationship between DNA sequence and function of the promoters characterised above. Partition modelling was used to identify positions within the sequence space that were having the greatest impact on promoter activity, and ANN and PLS models were subsequently used to make quantitative predictions of promoter activity.

\textit{Partition modelling}
Recursive partition modelling is a powerful technique for determining the relationship between a response variable and a set of independent variables without the use of a mathematical model\textsuperscript{50}. Partition models were fit to both the GFP and mOrange characterisation data sets. The number of times each promoter sequence position caused partitions in the data set across 100 random forests was quantified; the larger the number of partitions caused by a sequence position, the more important that position was predicted to be in determining promoter activity.

Sequence positions across the entirety of the sequence space were predicted to strongly influence regulatory activity for both reporters (Figure 3). In particular, sequence positions towards the 5' terminus of the sequence space were predicted to be important in determining promoter activity. This result suggested that UP elements, sequence motifs that are further upstream than the canonical RBS, -10 and -35 motifs and that boost transcription initiation through interactions with the C-terminal domain of the RNA polymerase alpha subunit\textsuperscript{51,52}, are active in Geobacillus.

Artificial Neural Network & Partial Least squares sequence-function modelling

Although the partition models provided useful insights to the relationship between promoter nucleotide sequence and function, they did not provide quantitative predictions of regulatory activity. We therefore applied 2 quantitative modelling approaches, linear Partial Least Squares (PLS) regression and non-linear Artificial Neural Networks (ANN).

To asses the predictive capability of PLS and ANNs when applied to Geobacillus cis-regulatory sequences, models were trained using data derived from the 95 characterised promoter::GFP fusions (Supporting Figure 4). In all instances, each of the 104 nucleotide positions within the promoter sequence was modelled as an individual x variable and GFP fluorescence was used as the response variable, y.

ANNs have previously been shown to return insufficiently accurate predictions when the response surface under investigation is complex and the number of observations in the training data set is small\textsuperscript{53}. Furthermore, although the PLS algorithm was specifically designed to model data sets in which the number of predictor variables is greater than the number of observations in the training set \textsuperscript{54}, the extreme scale of the promoter design space (there are $4^{100}$ potential 100 bp nucleotide sequences) compared to the number of empirically characterised
promoters was thought likely to result in models with limited predictive power. A reduction in the dimensionality of the modelled design space was therefore deemed necessary.

Characterising promoters of shorter length would have immediately reduced the dimensionality of the modelled design space. For example, 50 bp sequences would have been of sufficient length to contain the canonical location of the RBS, -10 and -35 consensus motifs. However, the partition results showed that sequence positions upstream of the -50 position were likely to be important in determining regulatory activity (Figure 3). Sequences of reduced length would therefore not have contained vital upstream regulatory motifs and may therefore have shown reduced activity as compared to the longer sequences.

The results of the partition modelling were therefore used to reduce the dimensionality of the modelled design space. PLS and ANN sequence-function models were derived that modelled GFP fluorescence as a function of varying number of nucleotide positions. Sequence positions were selected in descending order of the number of partitions caused in the 100 partition models (Figure 3). In all instances, model performance was quantified using an independent test set of 10 promoter sequences that were held-back from model training and validation.

The optimum PLS model that was obtained inferred promoter activity as a function of 20 nucleotide positions (Supporting Figure 5). The model returned an $R^2$ value of 0.6024 when applied to the training and validation data sets, and an $R^2$ value of 0.8901 when applied to the test set (Figure 4). These results suggested that the obtained PLS model provided a reasonable fit of the training data and had good predictive power when applied to previously unseen data.

A Design of Experiments (DoE) approach was used to optimise ANN architecture (Supporting Information). In total, over 113,500 single-layer ANNs were fit, varying in terms of the personality of the activation function used, the number of nodes in the hidden layer, the cross validation methodology and the number of promoter sequence positions modelled.

The optimal obtained ANN was an ensemble model that contained 2 constituent ANNs. Each of the constituent models used sigmoidal activation functions with 5 nodes in the hidden layer, and modelled promoter activity as a function of 20
nucleotide sequence positions. The optimal model returned an $R^2$ value of 0.9746 when applied to the training and validation data sets, and an $R^2$ value of 0.9691 when applied to the test set, suggesting a good fit of the training data and strong predictive power (Figure 4). For both ANN and PLS, models that inferred promoter activity as a function of complete 100 bp sequences showed lower predictive accuracy than models of reduced numbers of sequence positions (Supporting Information). This result validated the use of partition modelling to reduce the size of the modelled design space.

**Predicting the function of previously uncharacterised promoters**

To further test the predictive power of the putatively high-performing PLS and ANN models, a secondary test set of previously uncharacterised *Geobacillus* promoters was selected. 10 putative regulatory sequences were selected at random from across the promoter phylogeny (Figure 1A) and characterised in *G. thermoglucosidasius* upstream of GFP. However, despite the strong performance of the 2 models on the primary test set, neither model returned accurate predictions of promoter activity for the selected sequences (Figure 4); the PLS model returned an $R^2$ value of 0.3595 and the ANN returned an $R^2$ value of 0.2283. Consequently, the derived models were insufficiently general to permit accurate predictions of endogenous promoter activity or facilitate rational, forward promoter design.

**Future applications of promoter sequence-function modelling**

The lack of generality shown by the models derived in this investigation was probably the result of the limited number of characterised promoter sequences as compared to the scale of the design space, resulting in training set that does not adequately capture the complexity of the response surface. Although PLS and ANN promoter sequence-function models using comparatively small data sets have been described $^{55-57}$, the promoter libraries used in these studies contained considerable sequence homology, thereby restricting the complexity of the response surface under investigation. If accurate predictive models of more complex promoter design spaces are to be obtained, a training data set that contains several orders of magnitude more promoter sequences than the 80 sequences used here is likely necessary$^7, 26, 43$. However, the scale of the required promoter libraries might be impractical in non-model organisms.
Although high-throughput characterisation of libraries containing thousands of
genetic parts using techniques such as a combination of flow cytometry and
multiplexed DNA or RNA sequencing has been previously described\textsuperscript{7, 26, 43}, such
approaches require the acquisition of large numbers of transformants; approximately
50-fold library coverage is necessary to achieve accurate characterisation of
individual promoters\textsuperscript{43}. However, low transformation efficiencies in many non-model
organisms, including \textit{Geobacillus}, preclude the production of libraries of the required
scale, potentially limiting the usefulness of statistical sequence-function modelling in
these contexts.

\textit{In lieu} of a massive increase in the number of characterised sequences, the
novel bioinformatic approach to promoter identification that was developed in this
investigation, coupled with partition modelling to identify those sequence positions
that are key for determining promoter activity, could be used to provide an initial
screen of the design space in organisms for which understanding of \textit{cis}-regulatory
sequences is limited. This information could subsequently be used for DoE inspired
promoter optimisation in future studies by facilitating the rational design of limited
sequence libraries that vary only at the identified key positions. \textit{In vivo}
characterisation and \textit{in silico} modelling of the designed libraries could potentially
yield models of greater predictive power than those derived here without the need for
a large-scale increase in characterisation throughput.

The models that were derived in this study were based purely on the
statistical likelihood of a given nucleotide occurring at a given position within the
promoter sequence. Measures of biophysical promoter properties, such as mRNA
secondary structures, AT content or the free energy barrier for promoter-RNA
polymerase binding were not included on the basis that unsupervised ANN models
could potentially learn the effect of biophysical promoter properties without specific
terms being explicitly defined in the model. The inclusion of biophysical terms in
future modelling attempts may facilitate the derivation of more accurate predictive
models\textsuperscript{26, 43, 58} by providing more information about promoter function than can be
gleaned from sequence data alone. Alternatively, the use of distance metrics\textsuperscript{59} as
model terms to quantitatively define differences in nucleotide sequence between
promoters might also allow for more accurate mapping of the promoter sequence-
function design space\textsuperscript{60}. 
Finally, although the quantitative sequence-function models derived in this investigation were insufficiently general to determine pre hoc in vivo promoter activity, the potential for statistical modelling to enhance our fundamental knowledge of genetic regulation in complex systems cannot be overlooked. For example, partition modelling of the relationship between nucleotide sequence and in vivo promoter function yielded potentially useful insights into the structure of cis-regulatory elements in Geobacillus; regions of sequence upstream of the likely position of canonical promoter motifs were predicted to be important in determining promoter activity (Figure 3).

Conclusion

We developed a generally applicable method for the identification of constitutive promoters that combines bioinformatic filtering, empirical characterisation and machine learning to expand promoter toolkits in atypical host organisms and increase the understanding of the relationship between DNA sequence and function. The method was used to identify 80 promoters, covering a 2-log range of predictable expression levels, in G. thermoglucosidasius, of which 7 were shown to function consistently regardless of downstream coding sequence. Although sufficiently general in silico models of promoter activity could not be obtained using ANN or PLS, partition modelling identified regions of sequence upstream of the canonical prokaryotic promoter consensus regions that strongly influenced regulatory activity in Geobacillus.

Materials & Methods

Bacterial strains & plasmids

Type strains of Geobacillus kaustophilus (DSM7263), G. stearothermophilus (DSM22) and G. thermoglucosidasius (DSM2542) were obtained from the DSMZ (Brunswick, Germany). Cultures were freeze-dried ampoules and rehydrated as required following the DSMZ standard protocol. G. thermodenitrificans (K1041) was obtained from ZuvaSyntha Ltd. (Hertfordshire, UK).

NEB 5-alpha (New England Biolabs, Massachusetts, United States of America) chemically competent Escherichia coli strain (genotype: fhuA2 D(argF-
lacZ)U169 phoA glnV44 f80D(lacZ)M15 gyrA96 recA1 relA1 endA1 thi-1 hsdR17) was used for microbiological cloning, storage and amplification of plasmid vectors.

*E. coli* S17-1 (genotype: recA pro hsdRm RP4-Tc::Mu::Km::Tn7) was used as the mobilisation host for the conjugal transformation of *Geobacillus* spp. Transfer genes from the RP4 plasmid are integrated into the genome of *E. coli* S17-1, allowing for the conjugal transfer of plasmids containing the requisite mobilisation elements.

All putative promoter sequences were characterised *in vivo* using the pS797 vector (Supporting Figure 6). To facilitate conjugal transformation of *Geobacillus* spp., pS797 contained an origin of transfer (ORI T), comprised of the Nic region and *traJ* gene from the conjugal plasmid RP4. pS797 also contained 2 origins of replication, CoIE and BST1, to allow for propagation in *E. coli* and *Geobacillus* spp., respectively. 2 antibiotic selection markers were also present, allowing for selection by Ampicillin in *E. coli* and by Kanamycin in *Geobacillus*.

Both *E. coli* S17-1 and pS797 were obtained from ZuvaSyntha Ltd. (Hertfordshire, UK).

**Growth media**

All complex growth media were purchased from Becton Dickson UK (Berkshire, UK). *E. coli* cultures were propagated in Lysogeny Broth (LB; 10 g l⁻¹ tryptone, 10 g l⁻¹ NaCl, 5 g l⁻¹ yeast extract). Lennox Lysogeny Broth (LLB; 10 g l⁻¹ tryptone, 5 g l⁻¹ NaCl, 5 g l⁻¹ yeast extract) was used for co-culture of *E. coli* and *G. thermoglucosidasius* during conjugal transformation of *G. thermoglucosidasius*. All *Geobacillus* species were propagated in modified LB (mLB). mLB used a basal composition of LLB, supplemented with 1.05 mM C₆H₉NO₆, 0.91 mM CaCl₂, 0.59 mM MgSO₄ and 0.04 mM FeSO₄.

For all media types, agar was supplemented as required to 15 g l⁻¹. When required, *E. coli* growth media was supplemented with 100 µg ml⁻¹ ampicillin. *G. thermoglucosidasius* growth media was supplemented with 12.5 µg ml⁻¹ kanamycin.

**Bioinformatic identification of putative promoters from the core genome of 4 Geobacillus species**
The genomes of 4 *Geobacillus* species, *G. kaustophilus* (DSM7263), *G. stearothermophilus* (DSM22), *G. thermodenitrificans* (K1041) and *G. thermoglucosidasius* (DSM2542) were sequenced and de novo assembled. Genomes were sequenced using an Illumina MiSeq system, using reads with 300 bp paired end sequencing. The resulting raw sequencing reads were trimmed based on quality score using the fastq-mcf tool and assembled using SPAdes software (Version 3.5). Following assembly, the genome scaffolds were annotated using Prokka software (Version 1.9).

The GET_HOMOLOGUES software package was used to identify gene families with homologues in all 4 of the *Geobacillus* species of interest. To increase calculation robustness, 3 disparate algorithms were used to cluster homologous gene families: Bidirectional best-hit (BDBH), COGtriangles (COG) and OrthoMCL (OMCL). In all instances, the "-t" option was used to isolate only those clusters that contained single-copy proteins. All other software parameters were set as default. Only those clusters that were common to all 3 algorithms were selected for further analysis.

Once identified, the core coding sequences were extracted from the 4 genomes. Output files were parsed, reformatted to GenBank file format and imported into the Artemis genome browser. For each entry, the 100 bp immediately upstream of the start codon was extracted. BPROM software was subsequently used to screen the extracted 100 bp sequences for the presence and nucleotide composition of functional regulatory motifs. Additionally, putative promoters were screened against BPROM's list of known Transcription Factor Binding Sites (TFBS, Supporting Table 2). Any putative promoters containing TFBS were discarded.

The nucleotide sequences of the putative promoters were aligned using MUSCLE software and the resultant alignments were used to construct a phylogenetic tree using FastTree software. Putative promoters were subsequently manually clustered into 21 clades using FigTree software. Putative regulatory sequences sequences were selected at random for in vivo characterisation from these 21 clades. True randomness was achieved by using a random number generator that converted atmospheric noise into numerical values. Initially, those promoters that were selected for in vivo characterisation were manually checked using the Artemis genome browser to ensure that they did not overlap with any
adjacent coding sequences. Later, to expedite this process, BEDTools intersect\textsuperscript{71} was used to identify those putative promoters which were non-overlapping.

Putative promoters were aligned to transcripts of each of the 4 \textit{Geobacillus} species using Bowtie 2 software\textsuperscript{72}. Indexes of the genome files were prepared using the “build” command. Putative regulatory sequences were subsequently aligned to each \textit{Geobacillus} genome using Bowtie 2, with the resultant alignments provided in .sam format. The alignment .sam files were converted to .bam format, sorted and indexed using SAMtools\textsuperscript{73}. The resultant alignments were compared against the 4 selected \textit{Geobacillus} genomes using BEDTools intersect. The “-v” command was used to report only those putative promoters that were non-overlapping with any annotated features in the genome transcripts. Output files were provided in .bam format, and were subsequently converted to FASTA format using bam2fastx software\textsuperscript{74}.

\textit{Bioinformatic identification of putative promoter sequences from bacteriophage}

The genomes of 2 bacteriophages, \textit{Thermus} phage Phi OH2 (NC_021784) and \textit{Geobacillus} phage GBSV1 (NC_008376\textsuperscript{75}) were selected for analysis based on their ready availability from the GenBank database. The retrieved GenBank files were loaded into the Artemis genome browser\textsuperscript{66} and suitable intergenic regions of at least 100 bp length were manually identified. The 100 bp nucleotide sequences immediately upstream of the adjacent CDS were extracted and analysed using BPROM software\textsuperscript{42} to identify putative promoters. Putative promoter sequences were screened against BPROMs list of known TFBS, and any sequences that contained known TFBS were discarded.

\textit{Selection, synthesis and cloning of putative promoters for in vivo characterisation}

Following bioinformatic filtering, putative promoters were synthesised and independently cloned upstream of the coding sequences of 2 reporter proteins, Dasher GFP and mOrange\textsuperscript{76} (Supporting Figure 6). The \textit{Geobacillus} promoter phylogeny (Figure 1B) was used to rationally select putative regulatory sequences for \textit{in vivo} characterisation in \textit{G. thermoglucosidasius}. To maximise the portion of the design space that was empirically explored, at least 2 putative promoters were selected at random from each of the 13 clades of the phylogeny that contained more than 50 sequences. 2 putative promoters were also selected from each of the
analysed phage genomes. Initial characterisation of the bacteriophage promoters showed that only 1 out of the 4 selected sequences was active in *G. thermoglucosidasius* (Supporting Figure 7). This 1 active bacteriophage promoter was added to 99 putative promoters from the *Geobacillus* phylogeny to create a set of 100 putative regulatory sequences.

The 100 selected putative promoters were synthesised and cloned into the pS797 vector (Supporting Figure 6). In all instances, the reporter CDS (GFP or mOrange) was followed by the S718 terminator from the *G. thermodenitrificans* NG80 2-oxoglutarate ferredoxin oxioreductase subunit beta77. Putative regulatory sequences were either directly synthesised upstream of the relevant reporter CDS in pS797 by ATUM (Previously DNA 2.0, California, USA), or were synthesised as double stranded fragments by IDT (Illinois, USA) and cloned *in vitro* upstream of the relevant reporter CDS.

A type II*ls* restriction cloning methodology78,79 was used to join DNA parts. Parts were flanked with unique cloning affixes (Supporting Table 3) containing BsaI restriction sites. Part-specific post-digestion overhangs ensured that digested fragments were only able to ligate in a defined manner. In instances where putative promoters were synthesised by ATUM, the scar sequences that would have resulted from *in vitro* cloning of DRS and RBS were inserted into the sequence *in silico* prior to synthesis.

For *in vitro* cloning, terminator and reporter sequences were synthesised by ATUM in the pJ201 cloning vector. Cloning reactions consisted of 20 fmol of each of the pS797 destination vector and the relevant cloning vectors, with 10 U BsaI restriction endonuclease and 1 U T4 DNA ligase in 2 µl ligation buffer (10x Thermo Scientific FastDigest buffer supplemented with 0.5 mM ATP). Final reactions were made up to 20 µl with ddH2O. Reactions were incubated for 50 cycles of 37 °C for 2 min then 20 °C for 5 min. This was followed by final incubation steps of 50 °C for 5 min then 80 °C for 5 min. 10 µl of the incubated cloning reaction mix was used to transform chemically competent NEB 5-alpha *E. coli*, following the protocol described below. Plasmid construction was verified by diagnostic digest, gel electrophoresis and Sanger sequencing.

*Transformation of chemically competent E. coli*
**E. coli** S17-1 were made chemically competent using a modified version of the protocol described by Hanahan. 5 ml overnight cultures of **E. coli** S17-1 were used to inoculate 40 ml LB at a 1:1000 dilution. Inoculated cultures were incubated at 37 °C, with shaking at 220 rpm, until an OD$_{600}$ of 0.4-0.5 was reached. Cells were harvested by centrifugation at 4,500 g for 8 min at 4 °C and resuspended in 8 ml transformation buffer 1 (TF1: 150 g l$^{-1}$ Glycerol; 30 ml l$^{-1}$ 1 M CH$_3$CO$_2$K pH 7.5; 0.1 M KCl; 0.01 M CaCl$_2$·2H$_2$O. Adjusted to pH 6.4 with CH$_3$COOH, autoclaved, then supplemented with 50 ml l$^{-1}$ filter sterilised 1 M MnCl$_2$·4H$_2$O). Resuspended cells were subsequently incubated on ice for 15 min, and harvested as above. The resulting cell pellet was resuspended in 4 ml transformation buffer 2 (TF2: 150 g l$^{-1}$ Glycerol; 0.075 M CaCl$_2$·2H$_2$O; 0.01 M KCl. Autoclaved, then supplemented with 20 ml l$^{-1}$ filter sterilised 0.5 M MOPS-KOH pH 6.8). 100 µl aliquots of competent cells were flash frozen in liquid nitrogen and stored at -80 °C until required.

For transformation, 100-200 ng plasmid DNA was added to chemically competent **E. coli** of the relevant strain. Samples were incubated on ice for 40 min, then heat shocked at 42 °C for 2 min and incubated on ice for a further 5 min. 700 µl LB was added and the resulting samples were incubated at 37 °C, with shaking at 220 rpm, for 60 min. After incubation, samples were harvested by centrifugation at 4,300 g for 5 min, and 500 µl of the supernatant was removed. The cell pellet was resuspended in the remaining supernatant, 200 µl of which was subsequently plated out onto LB agar plates, with antibiotic selection as required. Plates were incubated at 37 °C for 16 h.

**Conjugal transformation of G. thermoglucosidasius**

Approximately 5 µl of transformed **E. coli** S17-1 was collected from a confluent plate-culture using a microbiological loop, suspended in 600 µl LLB and centrifuged at 4,300 g for 5 min. The supernatant was removed, and the resultant pellet re-suspended in a further 600 µl LLB. Approximately 10-15 µl wild-type **G. thermoglucosidasius** was collected from a confluent plate-culture using a microbiological loop, added to the **E. coli** suspension and re-suspended. The resulting bacterial mix was dispensed onto LLB agar plates, in drops of approximately 10 µl.
LLB plates were incubated at 37 °C for 7 h, followed by incubation at 60 °C for 1 h. The resulting biomass was re-suspended in 1 ml LLB, and used to create dilutions of 1:10 and 1:5 biomass to sterile LLB. 200 µl aliquots of each dilution were spread onto separate mLB agar plates containing 12.5 µg ml$^{-1}$ kanamycin. Plates were incubated at 55 °C for approximately 65 h.

In vivo characterisation of promoter activity

To prepare starter cultures of *G. thermoglucosidasius* for promoter characterisation, transformants were picked and restreaked on mLB agar plates, with antibiotic selection as required. Plates were incubated at 55 °C for 16 h. The resulting biomass was subsequently re-suspended in 5 ml mLB. Bacterial suspensions were then used to inoculate mLB to an OD$_{600}$ of 0.1, with antibiotic selection as required.

3 200 µl sample aliquots per transformant were loaded onto 96-well plates using either a Corbett Robotics CAS-1200 (Qiagen, Netherlands) or a Gilson Pipetmax 268 (Gilson Inc., Wisconsin, USA). To minimise the effect of position dependant bias, to which assays performed in a 96-well plate format can be susceptible$^{81}$, sample aliquots were loaded in a Latin rectangle design; no transformant was represented more than once on any given row or column of the microplate (Supporting Figure 8). 96-well plates with lid covers have been shown to suffer from significant loss of culture in the outermost wells through evaporation$^{82}$. To account for such edge effects, wells at the plate periphery were filled with 200 µl aliquots of sterile growth media. Microplates were incubated using PHMP Thermoshakers (Grant Instruments, UK). Incubation was at 60 °C, with shaking at 800 rpm.

Population-level measurements of culture absorbance and fluorescence were taken using a Tecan Infinite 200 PRO microplate reader (Tecan, Switzerland). For measurements of GFP activity, fluorescence excitation and emission values were 477 nm and 515 nm respectively. For measurements of mOrange activity, excitation and emission values were 546 nm and 576 nm respectively. In both cases, the gain of the instrument was set at 56. Absorbance of all cultures was measured at 600 nm.

Single-cell measurements of fluorescence activity were obtained using a BD FACS Aria II Fluorescence Activated Cell Sorter (FACS), equipped with a 100 µm nozzle. A sheath fluid of Phosphate Buffered Saline was used. Culture fluorescence
was excited at 488 nm and fluorescence intensity was recorded using a 530/30 nm
detector in the case of GFP fluorescence, and a 585/42 detector in the case of
mOrange fluorescence. 100,000 events were recorded per population.

*Promoter sequence-function modelling*

All sequence-function modelling was performed using JMP pro versions 12 &

*Partition modelling*

100 random forest models were generated for each of the GFP and mOrange
characterisation data sets. In all instances, 20% of the available promoter sequences
were randomly selected and withheld from model training to serve as a validation set.
Each random forest contained a maximum of 100 decision trees, with early stopping
if the addition of further trees to the forest did not improve the validation statistic.
Each tree was trained on a data set of 26 randomly selected promoter sequence
positions, drawn with replacement.

To generate partition trees, the selected sequences were divided into groups
that differed maximally in terms of the response of interest. For example, the
maximum difference in expression activity between 2 groups of promoters might be
obtained by splitting the training data into a group of sequences with guanine
residues at the -15 position, and another group where adenine, cytosine or thymine
residues are present at the -15 position (Supporting Figure 9). The resulting sub-
groups were further divided, resulting in the formation of a tree like structure. By
repeating the process multiple times on different, randomly selected portions of the
training data, a “forest” of decision trees was formed. Across the entire forest, the
more times a given factor caused a split in the data set, the better that factor was
predicted to be at explaining variation in the response of interest.

*Selection of an independent test set for PLS & ANN modelling*

To provide an independent test set on which to measure the predictive power
of the derived models, 10 promoter sequences were selected and withheld from
model training and validation. So that the test set contained promoters with a range
of activity levels, the distribution of GFP expression levels of the 95 characterised
sequences was analysed. 2 sequences were subsequently selected at random from the 1st distribution quartile, 5 promoters were selected from the interquartile range and 3 sequences were selected from the 4th quartile.

**Partial Least squares sequence-function modelling**

PLS models were trained that modelled GFP fluorescence as a function of varying numbers of sequence positions. The number of sequence positions modelled was systematically increased from 10 to 50 in increments of 5. Models that fit fluorescence as a function of the complete 104 bp promoters were also generated. For each of the 10 potential groups of $x$ variables, multiple PLS models were fit using the non-iterative linear PLS (NIPALS) algorithm and using either KFold or holdback cross validation to optimise the number of latent variables that were extracted from the original data, with a maximum of 10 latent variables permitted per model. Once trained and validated, the models were used to make predictions of activity for the 10 promoters in the withheld test set (Supporting Figure 5). The optimum model was judged to be the one that returned the highest $R^2$ and lowest Root Average Squared Error (RASE) value when applied to the test set; i.e. the model that had the lowest prediction error.

**Artificial Neural Network sequence-function modelling**

ANNs were fit using the multilayer perceptron algorithm of JMP software with sigmoidal activation functions. Network architecture was optimised using a Design of Experiments approach (Supporting Information).
Figure 1: Bioinformatic identification of putative promoter sequences.

A) Venn diagram showing the number of homologous gene families identified in the genomes of the 4 selected Geobacillus species by Bidirectional best-hit (BDBH), COG triangles (COG) and OrthoMCL (OMCL) clustering algorithms.

B) Phylogeny of putative promoters, rooted at the midpoint. At least 2 putative promoters were selected at random for in vivo characterisation from each of the clades containing > 50 sequences (highlighted in yellow).
Figure 2: *In vivo* characterisation of bioinformatically identified promoter sequences.

Bioinformatically identified putative promoter sequences were synthesised upstream of GFP and mOrange reporter sequences, and promoter activity in *G. thermoglucosidasius* was characterised after 24 h growth. In all instances, the positive control, the *G. thermodenitrificans* *ldhA* promoter is shown in dark grey, and the negative control, *G. thermoglucosidasius* transformed with an empty pS797 vector, is shown in red.

A) Heat map of GFP and mOrange expression levels of the 80 characterised promoters. Each column represents a disparate promoter. To account for differences in intensity between GFP and mOrange fluorescence signals, the mean fluorescence output of each promoter::reporter fusion was normalised to the fluorescence output of the negative control, *G. thermoglucosidasius* transformed to express the empty pS797 vector, at the relevant excitation and emission wavelengths. Regulatory sequences were defined as active if reporter fluorescence was statistically significantly greater than the negative control at the relevant wavelengths.
Significance was determined by ordinary one-way ANOVA with Dunnett’s multiple comparisons test and a significance level of 0.05.

B) Expression levels of the promoters for which fluorescence activity was statistically significant. Bars represent the mean of $n = 3$ independent starter cultures arising from independent transformation events, except in the case of the negative controls, where $n = 14$, and the positive controls, where $n = 11$. Error bars represent standard deviation.

C) GFP and mOrange expression levels are normalised to the negative control. Points represent individual promoter sequences. Promoter groupings were determined by K-means clustering based on the Euclidian distance of the points from the line of equivalence, $y = x$, which is represented by the dashed line.

D) Expression levels of the 7 promoters that functioned consistently regardless of CDS, as determined by flow cytometry. For each promoter::reporter fusion and the negative control, 100,000 events from each of 3 independent starter cultures arising from independent transformation events were combined to form a single “meta” population of 300,000 events. $+$ = ldhA positive control; $-$ = negative control.
Figure 3: Heat map showing the number of data set partitions caused in 100 random forests by individual regulatory sequence nucleotide positions when either GFP or mOrange fluorescence was used as the response variable.

The grey region represents the ACCT cloning scar between the Distal Regulatory Sequence (DRS) and RBS regions. As all of the characterised promoters were identical in these locations, these 4 positions were not included in the partition modelling.
Figure 4: Empirically measured promoter activity levels plotted against activity levels as predicted by the optimum obtained PLS and ANN models.

Points represent individual promoter sequences. Promoters that were used in model training and validation are shown in black, promoters that were part of the primary test set are shown in red, and sequences from the secondary test set are shown in blue. Empirical values are the mean of n = 3 starter cultures arising from independent transformation events. Standard deviation error bars are shown for both primary and secondary test sets, unless hidden by the points. The dashed lines represent the lines of equivalence, where empirically measured and predicted values are equal.
List of Abbreviations

ANN Artificial Neural Network
BDBH Bidirectional Best-Hit
bp Base Pair
CDS Coding Sequence
COG COG triangles
DoE Design of Experiments
DRS Distal Regulatory Sequence
OMCL OrthoMCL
PLS Partial Least Squares
RBS Ribosome Binding Site
TFBS Transcription Factor Binding Site

Supporting Information

The Supporting Information file for this submission contains the following:

- Supporting Text:
  - Analysis of the effect of type IIS restriction cloning scars on the activity of promoter sequences
  - Extended methods for Artificial Neural Network sequence-function modelling

- Supporting Figures:
  - Supporting Figure 1: Visualisation of a sequence alignment of 100 putative promoters used to identify the putative location of the Ribosome Binding Site (RBS).
  - Supporting Figure 2: The effect of cloning scar sequences on promoter activity.
  - Supporting Figure 3: A comparison of codon usage in the GFP and mOrange reporter sequences.
  - Supporting Figure 4: Activity levels of putative promoter sequences characterised upstream of GFP in G. thermoglucosidasius.
  - Supporting Figure 5: $R^2$ and Root Average Squared Error (RASE) values returned by PLS sequence-function models when applied to a test data set.
868  o Supporting Figure 6: Plasmid map of the pS797 expression vector used for in vivo characterisation of putative promoter sequences.
869  o Supporting Figure 7: Initial characterisation of putative promoter sequences isolated from bacteriophage genomes.
870  o Supporting Figure 8: Schematic representation of Latin rectangle 96-well plate layout used in promoter characterisation.
871  o Supporting Figure 9: Schematic representation of a random forest partition model, as applied to promoter sequences.
872  o Supporting Figure 10: Assessing the contribution of ANN model parameters to determining predictive power using a PLS model.
873  o Supporting Figure 11: Model performance statistics for ANNs modelling GFP fluorescence as a function of complete 104 bp promoters.

881  • Supporting Tables:
882  o Supporting Table 1: List of Transcription Factor Binding Sites (TFBS) used in promoter identification.
884  o Supporting Table 2: DNA sequences of cloning affixes used in type II restriction cloning.
886  o Supporting Table 3: Analysis of the native genes with which the characterised promoters were originally associated.
888  o Supporting Table 4: Artificial Neural Network parameters included in the architecture optimisation screening design, and the values specified for each parameter.
891  o Supporting Table 5: DNA sequences of the characterised promoters.

893  **Author Information**
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895  **Author Contributions**
899  J.G., T.P.H., D.A.P. and J.L. designed the study. R.K.T. and T.L. assisted with Bioinformatic analyses. J.G. and C.S. performed the characterisation experiments. J.G. and R.K.T. performed flow cytometry experiments. J.G. analysed the data and performed the sequence-function modelling. J.G. and J.L. wrote the manuscript. All authors commented on and revised the manuscript.
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Data Availability

The sequence data for the 4 Geobacillus spp. used in this study have been submitted to the NCBI Sequence Read Archive and are available under the accession number PRJNA521450.
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