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# Exploitation of the Escherichia coli lac operon promoter for controlled recombinant protein production

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#### **Abstract**

The *Escherichia coli lac* operon promoter is widely used as a tool to control recombinant protein production in bacteria. Here we give a brief review of how it functions, how it is regulated, and how, based on this knowledge, a suite of *lac* promoter derivatives has been developed to give controlled expression that is suitable for diverse biotechnology applications.

**Key words:** *Escherichia coli*, bacterial transcription initiation, *lac* promoter, *tac* promoter, Lac repressor, recombinant protein production

 **Abbreviations Used:** RPP, recombinant protein production; RNAP, RNA polymerase; bp, base pairs; CRE, Core-recognition element; CRP, cyclic AMP receptor protein; Lacl, lactose operon repressor protein; IPTG, isopropyl  $\beta$ -D-1-thiogalactopyranoside; PAR, promoter activity rating; GFP, green fluorescent protein; hGH, human growth hormone; Tat, twin arginine translocon pathway; LB: lysogeny broth; ONPG, o-nitrophenyl- $\beta$ -D-galactopyranose.

#### Introduction

Most of the recombinant protein production (RPP) systems used for expressing proteins in bacteria were constructed in the last century [1-4]. High level RPP provided by these systems enables the synthesis and purification of large amounts of soluble recombinant protein. However, the expression of difficult protein targets (e.g. membrane proteins or proteins secreted out of the cytoplasm), using these RPP systems, may be too high for cells to cope and adequately fold protein, resulting in substantial target degradation or the production of insoluble aggregates (i.e. inclusion bodies) [5-8]. Many "tricks of the trade" can be employed to slow down RPP expression and increase the level of soluble product, e.g. lowering the growth temperature, decreasing the inducer concentration or using a weaker promoter [6-9]. Although such tinkering can be very successful, determining the correct combination of refinements can be time-consuming. This can also be very "hit-and-miss", being dependent on the particular target protein in question [6, 9], and for some induction regimes, for example when using low inducer concentrations, only a proportion of the cells in a culture may in fact express recombinant protein [10-12].

The *E. coli lac* operon promoter was one of the first bacterial promoters to be adopted by biotechnologists for RPP, and it is still used today, especially when *Escherichia coli* is used as the host [6]. Here, we give a brief update of our current understanding of transcript initiation in bacteria, emphasising special features of the *lac* promoter and its regulation. We then review how, based on this information, many *lac* promoter derivatives have now been engineered in order to facilitate controlled RPP expression and avoid the problems that are concomitant with high level overexpression.

#### Transcript initiation and regulation at the E. coli lac operon promoter

Transcript initiation in bacteria takes place when the multisubunit DNA-dependent RNA polymerase holoenzyme (RNAP) interacts with a DNA promoter sequence (Figure 1A). In brief, the RNAP first interacts with double-stranded DNA to form a 'closed complex' in which determinants in different RNAP subunits interact with different promoter sequence elements (Figure 1B) [13]. Thus a determinant in the RNAP  $\alpha$  subunit C-terminal domain interacts with the promoter UP element, a determinant in Domain 4 of the RNAP  $\sigma$  subunit interacts with the promoter -35 element, and a determinant in Domain 3 of the RNAP  $\sigma$  subunit interacts with the extended -10 element (Figure 1B). Following this, Domain 2 of the RNAP  $\sigma$  subunit drives the local unwinding of 13-15 base pairs (bp) of promoter DNA to form the 'open complex', in which the single-stranded DNA template strand is positioned in the active site of the RNAP, such that initiation of DNA-templated RNA synthesis can take place (Figure 1C) [14]. Formation of the initiation-competent 'open complex' from the 'closed complex' is driven by further specific interactions between other RNAP determinants and promoter sequences

[15-17]. Thus, determinants in Domain 2 of the RNAP  $\sigma$  subunit interact with single-stranded bases of the promoter -10 element and with the promoter discriminator element. These interactions involve only the non-template strand of the locally unwound segment of promoter DNA, thereby permitting the single-stranded template strand to access the RNAP active site. The exact position of the template strand in the active site, and the location of the downstream junction between single-stranded and double stranded DNA, is set by other contacts involving amino-acid side-chains of the RNAP  $\beta$  and  $\beta$ ' subunits (which interact with the promoter Core-recognition element, CRE) (Figure 1C) [16].

The activity of any bacterial promoter is set by the formation of the 'closed complex' and the ensuing isomerisation to, and escape from, the 'open complex', as the RNAP copies the template strand to elongate its transcript. In the case of the E. coli lac operon promoter, defects in the UP element, -35 element and extended -10 element hinder 'closed complex' formation but this is remedied by binding of an activatory factor, the cyclic AMP receptor protein (CRP) to a target sequence centred between bp 61 and 62 (referred to as position -61.5) upstream of the transcript start (denoted as +1; see Figure 1D) [18, 19]. A second regulator protein, the lactose operon repressor protein (Lacl) binds to a high affinity target sequence, known as operator O1, centred at position +11 (Figure 1D) [19, 20]. Lacl binding to its target effectively shuts down lac promoter activity, but repression can be broken by the presence of allolactose (a breakdown product of lactose) or by the sugar analogue IPTG (isopropyl β-D-1-thiogalactopyranoside), which both bind to the Lacl repressor and cause it to release operator DNA [20]. Lacl-dependent repression of the lac promoter is supported by Lacl binding to two secondary weaker operators, O3, located at position -82 (Figure 1D), and far downstream, O2 [19-23]. Here, we review how the starting lac promoter has been engineered to make it fit for different biotechnology purposes. We focus on constructions that release the requirement for CRP, base changes in key promoter elements, and the exploitation of the O3 operator.

#### Activator-independent lac promoter derivatives

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140 141 Figure 2A illustrates the organisation of the E. coli lac operon promoter, showing key promoter elements, the DNA site for CRP, and the location of Lacl-binding operators, O1 and O3. Figure 2B shows the base sequence of a typical DNA fragment carrying the lac promoter that might be used in any biotechnology application. The fragment, which carries a useful restriction site at each end, is denoted *lac* O3O1. Potentially, regulation by CRP might be exploitable but, since CRP activity is difficult to control by external cues, many biotechnology applications that use the *lac* promoter have sought to eliminate CRP effects and focussed on regulation by Lacl. One way to do this is by the use of the lac UV5 mutant promoter. This mutant promoter carries a 2 bp change in the promoter -10 hexamer element that creates a consensus -10 promoter element (Figure 2B) [19, 24]. The alternative is to replace lac promoter upstream DNA sequences and this has been done in the tac promoter, which is a chimeric fusion between the upstream elements of the E. coli trp promoter and the downstream elements of the lacUV5 promoter [1, 2]. Figure 2A illustrates both the lacUV5 and tac promoters, and Figure 2B shows the base sequence of DNA fragments carrying these promoters. Note that the tac promoter carries consensus -35 and -10 promoters elements and a single operator for Lacl, O1. For E. coli promoters that are dependent on the housekeeping sigma factor  $\sigma^{70}$  the optimal spacing between the -35 and -10 promoter elements is 17 bp and deviation from this leads to a reduction in promoter activity [25, 26]. Thus, it worth noting that both the tac and lac promoters are in fact suboptimal, having spacers of 16 and 18 bp, respectively (Figure 2B).

Figure 3A-C illustrates the results of simple assays to compare the activities of the tac, lacUV5 and lac promoters. To perform these assays, the tac O1 fragment, the lacUV5 O3O1 fragment and the lac O3O1 fragment were each cloned into a plasmid expression vector (pRW50) such that the promoters controlled transcription of the gene (lacZ) encoding  $\beta$ -galactosidase [27]. Recombinant plasmids were then transformed into a  $\Delta lac$  E. coli host

strain and  $\beta$ -galactosidase expression was measured. As expected the hierarchy of promoter activity was tac > lacUV5 > lac (Figure 3).

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### Modulation of promoter activities using upstream *lac* operator sequences A key feature of Lacl-dependent repression of the *E. coli lac* promoter is the coli lace promoter activities using upstream *lac* operator sequences.

A key feature of LacI-dependent repression of the *E. coli lac* promoter is the contribution of the auxiliary upstream operator, O3, and this has been exploited to tailor promoter activity levels. Data in Figure 3A illustrate how the introduction of certain *lac* operator sequences (*i.e.* O1 or the high affinity "ideal" *lac* operator Oid, see Figure 2A [21, 28, 29]) at position -82 of the *tac* O1 promoter fragment reduced the high activity of the *tac* promoter. Thus, expression from the starting *tac* O1 promoter fragment and each of the *tac* O3O1, *tac* O1O1 and *tac* OidO1 derivatives was induced by IPTG but the introduction of the O1 and Oid operator sequences decreased IPTG-induced expression levels. Similarly, data in Figures 3B and 3C illustrate how the introduction of higher affinity *lac* operators into the upstream region of the *lac*UV5 promoter (*i.e.* in the *lac*UV5 O1O1 and *lac*UV5 OidO1 promoter fragments) or the *lac* promoter (*i.e.* in the *lac* O1O1 and *lac* OidO1 promoter fragments) decreased promoter activity.

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#### The power of lac: combinations make anything possible

In addition to controlling promoter activity by upstream-bound Lacl, activity can be modulated by point mutations in different promoter elements. Data in Figure 3D illustrate how the p34G, p14G, p9A or p8A substitutions (at positions -34, -14, -9 or -8), which make the -35 element, the extended -10 or the -10 element more similar to the respective consensus, can be combined with different operator combinations to produce a suite of IPTG-inducible promoters with a wide range of activities.

To illustrate the use of these promoters in RPP, we selected a subset of 8 promoters from the above suite and gave each a promoter activity rating (PAR) value of PAR1 to PAR8 (Figure 4A). Our rationale for this is that, depending on the target protein being expressed, specific IPTG-induced expression levels could be achieved. To examine this, some of the promoter constructs were introduced into the low-copy-number vector pTorA-GFP [30] and the high-copy-number vector pHAK1 [31], using standard techniques [32-34]. For pTorA-GFP derivatives, each PAR promoter drives the expression of a torA-GFP-6his fusion (GFP, green fluorescent protein), whilst, for pHAK1, each construct expresses a torA-hGH-6his fusion (hGH, human growth hormone). Note that the torA signal sequence in each case will direct the recombinant protein to the Tat (twin arginine translocon) system for periplasmic targeting [7, 30, 31]. Plasmid constructs were transferred into E. coli BL21 cells and RPP was induced in bacterial cultures by the addition of 1 mM IPTG for 3 hrs [7]. Normalised total cellular protein from cells, expressing either TorA-GFP-6His or TorA-hGH-6His, were then analysed by Western blotting. Results in Figures 4B and 4C show that both GFP and hGH were expressed using the PAR promoter constructs. Note that, in some instances, two product bands can be observed, in each case the species with the higher molecular weight still carries the TorA signal sequence, whilst the smaller species has been processed and lacks the TorA moiety [7, 30, 31, 35]. Importantly, using this expression system, inducible expression of different target proteins can be set to specific levels when using both low- and high-copy-number vectors.

#### **Perspectives**

The *lac* promoter and its derivatives have been widely used to express many recombinant proteins to high levels in *E. coli* [1, 2, 7, 36] and many currently used vectors have been designed to optimise expression. However, there are situations where expression must be moderated. For example, the secretion of recombinant biopharmaceuticals out of the *E. coli* cytoplasm into the periplasm is often a preferred industrial strategy, as this minimises downstream processing costs, since the target protein can be purified from the periplasmic contents, with minimal cellular and DNA contamination [37]. For this to be successful, RPP needs to be slowed down so that product is not degraded before it is transported [7].

In this review, we have sought to show how knowledge of the *E. coli lac* promoter can be exploited to produce a suite of derivative promoter fragments to cope with any situation. Previous promoter engineering had focused on altering the *lac* promoter -10 and -35 elements to change the basal promoter expression of constructs [1, 38]. Recent advances in our understanding of transcript initiation in bacteria and its regulation now allow *lac* promoters to be constructed with different operator sequences to alter the induced level of expression. This is possible because, even in the presence of inducer, LacI has some affinity for its operator sequence and, thus, in the induced state the LacI repressor can still remain bound to DNA, as has been observed in single molecule studies [39].

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#### **Author contributions**

D.F.B., S.J.W.B. and C.R. conceived and designed the research programme. D.F.B, R.E.G and K.L.R. performed the experiments. D.F.B. wrote the manuscript with input from all authors.

#### **Competing Interests**

The Authors declare that there are no competing interests associated with the manuscript.

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#### FIGURE LEGENDS

- Figure 1. Transcript initiation at bacterial promoters and regulation at the *lac* promoter (A) Bacterial promoter elements. A sketch showing the organization of different promoter
- elements at a model bacterial promoter with respect to the transcript start site (+1). Elements are denoted by coloured rectangles with their consensus sequence motifs indicated. Note that bacterial promoters usually contain a selection of these elements but rarely contain all of them [40].
- **(B)** Sketch illustrating the key interactions between different domains of the RNAP sigma and alpha subunits and different promoter elements in the initial "closed" complex at a promoter. Note that the DNA duplex remains double-stranded and that the contacts result in Sigma Domain 2 being positioned adjacent to the promoter -10 element.
- **(C)** Sketch illustrating the key interactions between different RNAP determinants and different promoter elements in the transcriptionally-competent "open" complex at a promoter. Note that the DNA duplex around position +1 is unwound, with Sigma Domain 2 making specific base contacts with the single-stranded non-template strand of the promoter -10 element, thereby permitting the single-stranded template strand to enter the RNAP active site.
- **(D)** A schematic representation of the *E. coli lac* operon promoter. The -35 and -10 promoter elements are shown as green and yellow boxes, respectively, the *lac* operator sequences (O1 and O3) are red boxes and the CRP binding site is shown by inverted arrows. The position of each element is given with respect to the start site of transcription (+1). The Lacl tetramer binds to the O1 and O3 operators to form a repression loop, silencing lac operon expression (-ve), whilst CRP activates transcription (+ve).

### Figure 2. Organisation of *lac*, *tac* and *lac*UV5 promoter constructs

- (A) The panel shows schematic representations of the *lac* O3O1, *lac*UV5 O3O1 and *tac* O1 promoters and the important elements involved in their regulation. All numbering is in relation to the promoter transcription start (+1), which is indicated by a bent arrow. The O1 and O3 operator sequences, which bind the LacI repressor, are indicated by red boxes, the -35 promoter elements by green boxes, and the CRP site, within the *lac*UV5 and *lac* promoters, is represented by inverted arrows. The *plac* -10 promoter element is shown as a yellow box, whilst the *plac*UV5 and *ptac* consensus -10 elements are gold. The sequence of the O3, O1 and Oid operators, and the site at which different operator sequences were introduced into the *tac* O1, *lac*UV5 O3O1 and *lac* O3O1 promoters is shown by an arrow.
- **(B)** The panel shows the base sequence of DNA fragments carrying the *lac* O3O1, *lac*UV5 O3O1 and *tac* O1 promoters. All promoter fragments carry the relevant sequences from -92 to +38, in relation to the transcription start, and possess terminal EcoRI and HindIII sites for cloning into the *lacZ* expression vector, pRW50 [27]. The O1 and O3 operator sequences are highlighted in grey, CRP binding targets are underlined and the -35 and -10 promoter elements are in bold and underlined, having also been aligned to the relevant consensus sequence for each element [40]. Note that the *lac*UV5 promoter only differs from the wild-type *lac* promoter by carrying a consensus -10 element (*i.e.* the p8A and p9A substitutions) [24].

#### Figure 3. Expression levels of engineered ptac and plac promoter derivatives

The figure illustrates measured  $\beta$ -galactosidase activities of *E. coli* K-12 strain JM109 ( $\Delta lac$ , lac) cells carrying pRW50 [27, 41] containing different promoter fragments. Promoter fragments were generated using PCR and cloned into pRW50 to create lacZ transcriptional fusions, using standard techniques [27, 33]. Panels (A), (B) and (C) show the effect of introducing the O3, O1 and Oid operator sequences into the upstream region of the tac O1, lacUV5 O3O1 and lac O3O1 promoter fragments, respectively, as displayed in Figure 2. (D) The panel details the effect of introducing point mutations into the -10 and -35 elements of various lac promoter derivatives (*i.e.* lac O3O1, lac O1O1, lacUV5 O3O1, lacUV5 O1O1) to

improve these regions in relation to the extended -10 and -35 consensus sequences (see Figure 2) [42]. By convention, locations are labelled in relation to the plac transcript start point (+1). Note that the lac O1O1 D19 promoter carries a base pair deletion at position -19. The fold increase in expression, in comparison to the weakest promoter (lac O1O1) is indicated for each promoter in the presence of IPTG. In all panels, JM109 cells were grown in LB medium until mid-exponential phase, in the presence or absence of 1 mM IPTG. βgalactosidase activities were determined using o-nitrophenyl-β-D-galactopyranose (ONPG) and a Miller protocol, as in our previous work [43], and are expressed as nmol of ONPG hydrolysed min<sup>-1</sup> mg<sup>-1</sup> dry cell mass. Each activity is the average of three independent determinations and the error bars represent the standard deviation of values.

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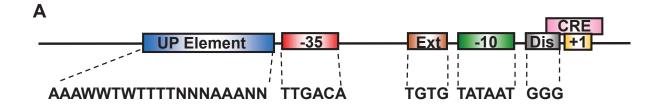
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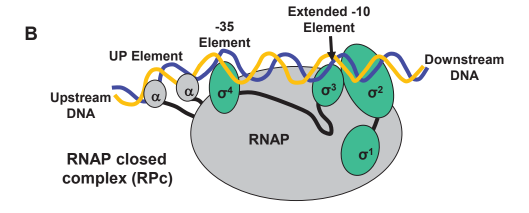
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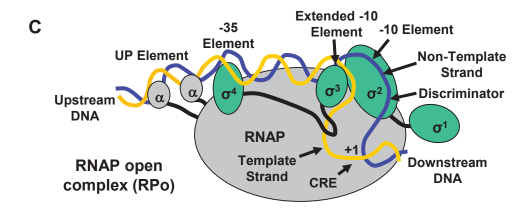
#### Figure 4. The expression of TorA-protein fusions can be set to different levels with **PAR** promoter constructs

(A) Expression from the PAR promoter constructs. The panel shows the β-galactosidase activities measured in the E. coli K-12 strain JM109 ( $\Delta lac$ ,  $lacl^q$ ) containing the lacZexpression vector pRW50 carrying various PAR promoters. PAR promoters (PAR1 to PAR8) were selected from the various tac and lac promoters detailed in Figure 3. β-galactosidase activities are expressed as nmol of ONPG hydrolysed min<sup>-1</sup> mg<sup>-1</sup> dry cell mass. Each activity is the average of three independent determinations and standard deviations are shown for all data points. (B) Expression of a TorA-GFP-6His protein fusion in BL21 cells (Novagen), using pGFP-TorA based vectors [30]. PAR promoters, PAR4 to PAR6 and PAR8 were introduced into the low-copy-number vector pGFP-TorA, such that each promoter drives the expression of a torA-gfp-6his fusion. Note that pGFP-TorA is a derivative of expression vector pEXT22 [44], which has been included as an empty vector control. (C) Expression of a TorA-hGH-6His protein fusion in BL21 cells, using pHAK1 based vectors [31]. PAR promoters, PAR5 to PAR8 were introduced into the high-copy-number vector pHAK1 such that each promoter drives the expression of a torA-hGH-6his fusion. Note that the torA signal sequence directs the GFP-6His and hGH-6His moieties to the Tat translocon for periplasmic targeting [7, 30, 31]. Cells were grown in LB medium until an OD<sub>600</sub> of ~0.4 when RPP was induced by the addition of 1 mM IPTG for 3 hrs. In (B) and (C) expression of TorA-GFP-6His and TorA-hGH-6His was analysed by Western blotting (upper panel), using either anti-GFP antibody (Sigma) or polyclonal anti-hGH antibody, respectively, and a Coomassie blue strained SDS-PAGE gel (lower panel), using normalised total cell protein from BL21 cells, as detailed in [45]. Samples were calibrated by loading Page Ruler Plus prestained markers (Thermo Scientific). Note that, in the Western blot in panel (B), all lanes were from the same blot and have only been separated to aid presentation of the data.

Figure 1







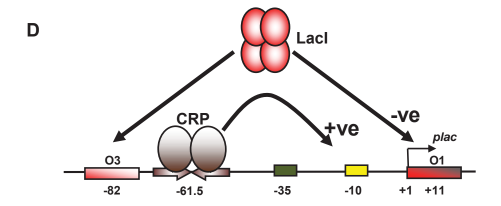


Figure 2

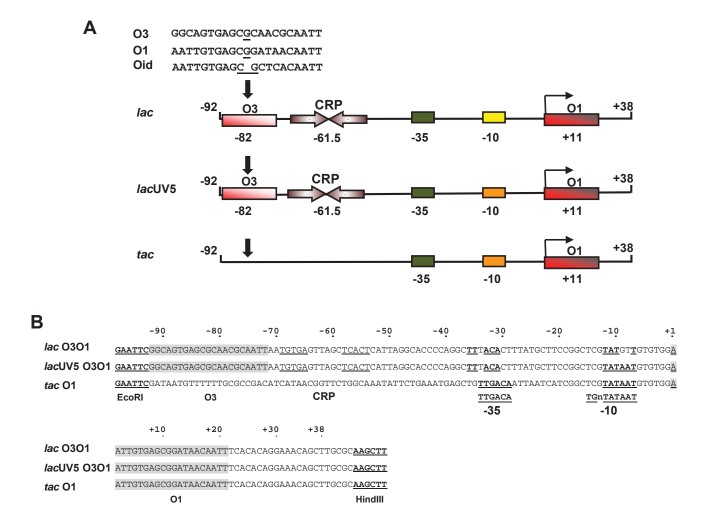
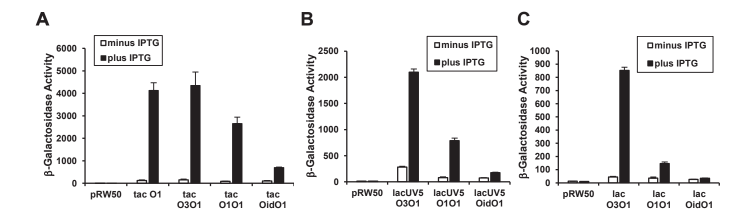


Figure 3



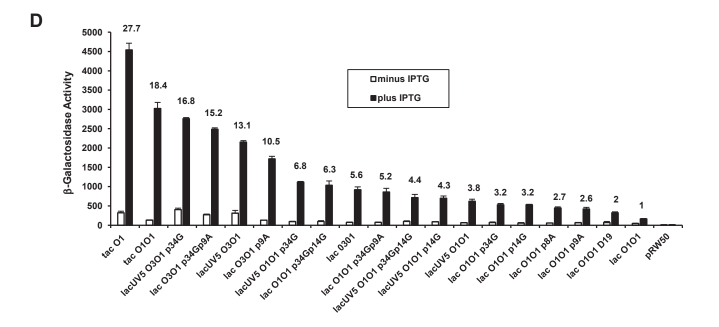


Figure 4

