

Synthetic Biology, 2020, 5(1): ysaa006

doi: 10.1093/synbio/ysaa006 Advance Access Publication Date: 8 June 2020 Research Article

# Dual UTR-A novel 5' untranslated region design for synthetic biology applications

## Simone Balzer Le<sup>1,2</sup>, Ingerid Onsager<sup>3</sup>, Jon Andreas Lorentzen<sup>1,4</sup>, and Rahmi Lale <sup>1,\*</sup>

<sup>1</sup>PhotoSynLab, Department of Biotechnology, Faculty of Natural Sciences, Norwegian University of Science and Technology, N-7491, Trondheim, Norway,

<sup>2</sup>Present address: Department of Biotechnology and Nanomedicine, SINTEF Industry, N-7465, Trondheim, Norway,

<sup>3</sup>Present address: Diatec Monoclonals, 0349 Oslo, Norway

<sup>4</sup>Present address: Department of Pathology, Stavanger University Hospital, 4011 Stavanger, Norway

\*Corresponding author: E-mail: rahmi.lale@ntnu.no

#### Abstract

Bacterial 5' untranslated regions of mRNA (UTR) involve in a complex regulation of gene expression; however, the exact sequence features contributing to gene regulation are not yet fully understood. In this study, we report the design of a novel 5' UTR, dual UTR, utilizing the transcriptional and translational characteristics of 5' UTRs in a single expression cassette. The dual UTR consists of two 5' UTRs, each separately leading to either increase in transcription or translation of the reporter, that are separated by a spacer region, enabling *de novo* translation initiation. We rationally create dual UTRs with a wide range of expression profiles and demonstrate the functionality of the novel design concept in *Escherichia coli* and *Pseudomonas putida* using different promoter systems and coding sequences. Overall, we demonstrate the application potential of dual UTR design concept in various synthetic biology applications ranging from fine-tuning of gene expression to maximization of protein production.

Key words: 5' untranslated region; transcription; translation; Escherichia coli; Pseudomonas putida

#### Introduction

The DNA region corresponding to the 5' untranslated region of mRNA (UTR) plays a central role in gene expression and protein production. At the DNA level, it is involved in transcript formation due to the interplay between promoter and the initially transcribed sequences (ITS), which covers the first 15 nt in *Escherichia coli* (1–3). At the mRNA level, it influences transcript stability and translation because of secondary structure formation (4–6), interaction with external factors i.e. proteins (7), metabolites (8, 9) and small RNAs (10) in addition to ribosome

binding and translation initiation (11). The translation initiation mainly involves around 15 nt preceding the start codon (12) including the Shine–Dalgarno (SD) sequence as well as the 5' end of the following coding sequence (4).

In bacteria, transcription and translation are coupled and a physical link between transcript formation and transcript turnover (translation and mRNA degradation) has been shown (13). It has been reported that the translation rate is likely to affect the transcription rate which indirectly affects mRNA stability (14–17). The DNA sequence of the 5' UTR has also an

Submitted: 27 September 2019; Received (in revised form): 29 April 2020; Accepted: 29 May 2020

<sup>©</sup> The Author(s) 2020. Published by Oxford University Press.

This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/ licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com

under-recognized role on transcript formation involving nucleotides downstream of the ITS (18, 19). With all the abovementioned characteristics, 5' UTR is one crucial contributor to the maintenance of a fine balance between transcription, transcript stability and translation.

In synthetic biology applications, it is desirable to have a predictive control over the levels of gene expression (20–22). Currently, there are several in silico tools available enabling the design of synthetic 5' UTR sequences for efficient translation initiation (15, 23–25), and these are also applied in combination with a selection of promoters (15, 22, 26). However, because of 5' UTRs sequence proximity both to promoter and coding sequence regions a physical context dependency exists (27), and it has consequently proven difficult to design optimal 5' UTR sequences solely based on translational properties (15). The multiple functionalities overlapping in 5' UTRs, hence represents an optimization challenge for modular design of biological circuitry.

We previously constructed a 5' UTR plasmid DNA library in which a 22 nt stretch, of the 32 nt long 5' UTR DNA sequence upstream of a  $\beta$ -lactamase coding sequence was randomly mutagenized using synthetic degenerate oligos (18). Escherichia coli cells harboring the plasmid DNA library were screened for clones that showed increased ampicillin resistance at the induced state ( $\beta$ -lactamase production positively correlates with the host's ampicillin resistance) with each clone carrying a unique 5' UTR variant. Escherichia coli clones could be categorized into two distinct groups based on their increased ampicillin resistance phenotype: one group of clones with increased reporter translation, high translation rates per reporter transcript; and another group of clones with increased reporter transcription phenotype but not corresponding translation, low translation rates per reporter transcript (18). The 5' UTR variants were identified as a result of screening efforts using a single 5' UTR, and the screening would not allow the identification of specific 5' UTRs with respect to the reporter transcription or translation rate. With these observations, we speculated that the DNA sequence composition of a single 5' UTR leads to a compromise between transcriptional and translational processes; hence it might not be possible to identify a sequence composition that is optimized for both of these processes in a short 5' UTR sequence.

In this study, we designed new functional screening tools (bicistronic artificial operons) to identify 5' UTR variants with desired transcriptional and translational characteristics. We constructed dual UTRs by incorporating the identified 5' UTRs in a single expression cassette. We then demonstrate that the combination of 5' UTRs with transcriptional and translational characteristics can have a synergistic effect on the gene expression and protein production beyond the levels achieved with single 5' UTRs. We demonstrate the functionality of the dual UTR concept in *Escherichia* coli and in an emerging SynBio chassis *Pseudomonas* putida (28).

#### Methods

#### Bacterial strains and growth conditions

Recombinant E. coli DH5 $\alpha$  (Bethesda Research Laboratories), E. coli RV308 (ATCC 31608) and P. putida KT2440 were cultivated in Lysogeny Broth (LB) (10 g/l tryptone, 5 g/l yeast extract and 5 g/l NaCl) or on Lysogeny Agar (LB broth with 15 g/l agar) supplemented with 50 µg/ml kanamycin. Selection of E. coli DH5 $\alpha$  transformants was performed at 37°C, while 30°C was used for

all growth experiments. For the induction of the promoter systems, first overnight cultures were inoculated into fresh medium and were grown until the mid-log phase was reached, and afterwards were induced at the levels specified.

#### **DNA** manipulations

Standard recombinant DNA procedures were performed as described by Sambrook and Russell (29). DNA fragments were extracted from agarose gels using the QIAquick gel extraction kit and from liquids using the QIAquick PCR purification kit (QIAGEN). Plasmid DNA was isolated using the Wizard Plus SV Minipreps DNA purification kit (Promega) or the NucleoBond Xtra Midi kit (Macherey-Nagel). Synthetic oligonucleotides were ordered from Sigma-Aldrich or Eurofins. Restriction cloning was performed according to recommendations from New England Biolabs. PCR reactions were carried out with the Expand High Fidelity PCR System (Roche Applied Science) or Q5 High-Fidelity DNA Polymerase (NEB). Escherichia coli clones were transformed using a modified RbCl protocol (Promega) and P. putida KT2440 was transformed using an electroporation protocol described by Hanahan et al. (30). The constructed plasmids were confirmed by sequencing performed at Eurofins/GATC Biotech using primer 5'-AACGGCCTGCTCCATGACAA-3' for pAO-Tr-, pIB11-(18), pdualUTR-; and primers 5'-CTTTCACCAGCGTTTCTGGGTG-3' and 5'-CAAGGATCTTACCGCTGTTG-3' for pAO-Tn-based constructs.

#### Plasmid constructions

All plasmids are based on the broad-host range mini-RK2 replicon. For the construction of the pAO-Tr plasmid, the bla coding sequence was amplified from the plasmid pIB11 (18) with the primers 5'-GCAGGCGGAATTCTAATGAGGTCATGAACTTATGAG TATTCAACATT-3' and 5'-CTAGAGGATCCCCGGGTACCTTTTCT ACGG-3', introducing the restriction sites EcoRI and BamHI, and was cloned into the pIB22 (31) plasmid as EcoRI-BamHI fragment downstream of the *ce*lB gene. This construction resulted in the plasmid pAO-Tr.

For the construction of pAO-Tn, the *celB* coding sequence and the DNA sequence corresponding to its 5' UTR were PCR amplified from the pAO-Tr plasmid using the primer pair 5'-ACCCCTTAGGCTTTATGCAACAgaaACAATAATAATGGAGTCAT GAACtTATG-3' and 5'-CTTTCACCAGCGTTTCTGGGTG-3'. The resulting PCR product was digested with Bsu36I and EcoRI and re-introduced into pAO-Tr using the same restriction sites leading to pAO-Tn(-1). By this cloning, the additional NdeI and PCII restriction sites were removed (indicated by small letters). The *bla* coding sequence was PCR amplified from pIB11 using the primer pair 5'-cggaattCAACATGTACAATAATaatg-3' and 5'-AGCTAGAGGATCCCCGGGTA-3' and the resulting PCR product was cloned as an EcoRI-BamHI fragment into the pAO-Tn(-1) plasmid resulting in pAO-Tn.

For the individual characterization of Tr- and Tn-UTR's phenotype, 5' UTR DNA sequences were integrated between the unique PciI and NdeI sites in the plasmid pIB11 as annealed pairs of forward and reverse synthetic oligonucleotides.

For the construction of plasmids carrying the dual UTRs, pdualUTR, the native 5' UTR DNA sequence upstream of the *bla* coding sequence in pIB11 was substituted with the annealed oligonucleotides 5'-CATGTACAATAATAATGGAGTCATGAACATAT CTTCATGAGCTCCATTATTATTGTATATGTACAATAATAATGGA GTCATGAACA-3' and 5'-TATGTTCATGACTCCATTATTATTG

#### TACATATACAATAATAATGGAGGTCATGAAGATATGTTCATGAC TCCATTATTATTGTA-3'.

#### For the construction of pdualUTR with the *mCherry* reporter gene, an *E.* coli codon-optimized variant of the *mCherry* coding sequence (a gift from Yanina R. Sevastsyanovich, University of Birmingham), was PCR-amplified using primer pair 5'-GCTGCA TATGGTTTCTAAAGGTGAAGAAG-3' and 5'-GCTCGGATCCTTAT CATTTATACAGTTCGTCCATAC-3' and digested with NdeI and BamHI. The digested fragment was then used to replace the *bla* coding sequence in pdualUTR resulting in pdualUTR-mCherry.

For the construction of all dual UTR combinations annealed synthetic oligonucleotides flanked by PciI and SacI (Tr-dual UTR) and SacI and NdeI (Tn-dual UTR), sticky ends were inserted into pdualUTR or pdualUTR-mCherry using the respective restriction enzyme recognition sites.

For the construction of the  $P_{BAD}$  system, the  $P_{BAD}$  expression cassette was PCR amplified from pSB-B1b (32) using primer pairs 5'-ATGGAGAAACAGTAGAGAGTTG-3' and 5'-TACATGGCTCTGC TGTAG-3'. Each of the five pdualUTR-mCherry vectors were PCR amplified using the reverse primer 5'-CTCCCGTATCGTAGT TATC-3' in combination with one of the forward primers 5'-CAACTCTCTACTGTTTCTCCATAACATGTTACCATGATAATGGA G-3', 5'-CAACTCTCTACTGTTTCTCCATAACATGTTACAATAATA ACGGAGTCATG-3', 5'-CAACTCTCTACTGTTTCTCCATAACATGT AATAAACTAAAGGAGTTATG-3' 5'-CAACTCTCTACTGTTTCTCC ATAACATGTACAATAATAATGGAGTCATGAACATATC-3' each targeting the dual UTR region in r31n47, r50n47, n47r31/n47r50 and wtwt, respectively. The DpnI treated and purified PCR-products with overlapping ends were assembled using in vivo homologous recombination method in E. coli strain DH5a, leading to pdualUTR-PBAD-mCherry (SOM pdualUTR-PBADmCherry.gb). The correct constructs were confirmed by sequencing using the reverse primer binding to the mCherry coding sequence, 5'-GATGTC AGCCGGGTGTTTAAC-3'.

## Generation and screening of 5′ UTR libraries based on pAO-Tr and pAO-Tn

5′ UTR libraries were constructed in pAO-Tr and pAO-Tn by cloning the synthetic degenerate oligonucleotides between their respective NdeI and PciI restriction sites, as described previously (18, 33). After transformation of plasmid DNAs to *E. coli* DH5 $\alpha$ , libraries with ~280 000 transformants (pAO-Tr-based) and ~370 000 transformants (pAO-Tn-based) were generated.

#### $\beta$ -Lactamase expression analysis

β-Lactamase enzymatic assay was performed following the protocol described elsewhere (34). For protein production analysis by SDS-PAGE or western blot, E. coli RV308 was used. Escherichia coli RV308 clones were grown in super broth (32 g/l peptone, 20 g/l yeast extract and 5 g/l NaCl). Expression was induced in the midlog phase, and cultures were harvested 5h after induction with 2 mM m-toluic acid. 0.1 g pellet (wet weight) were washed with 0.9% NaCl and resuspended in 1.5 ml lysis buffer (25 mM Tris-HCl, pH 8.0, 100 mM NaCl, 2 mM EDTA) followed by incubation with 0.2 g/l lysozyme on ice for 45 min and sonication (3 min, 35% duty cycle, 3 output control). After addition of 10 mM MgCl<sub>2</sub> and treatment with 125U Benzonase Nuclease (Sigma-Aldrich) for 10 min, the lysate was centrifuged to separate the soluble supernatant fraction from the pellet. The insoluble pellet fraction was resuspended in 1.5 ml lysis buffer. Both fractions were run on an SDS-PAGE using 12% ClearPAGE<sup>™</sup> gels and ClearPAGE<sup>™</sup> SDS-R Run buffer (C.B.S. Scientific) followed by staining with Coomassie Brilliant Blue R-250 (Merck). Western blot analysis was performed as described elsewhere (29).

#### mCherry production analysis

mCherry activity was determined with an Infinite M200 Pro multifunctional microplate reader (Tecan) by measuring the fluorescence of  $100 \,\mu$ l untreated culture with excitation and emission wavelengths of 584 nm (9 nm bandwidth) and 620 nm (20 nm bandwidth), respectively, and normalization against absorbance reading at OD<sub>600</sub>. Three bacterial colonies from each of the dual UTR constructs were transferred into a 96-well plate (Corning, black, flat bottom, clear) with each well containing 100  $\mu$ l LB with kanamycin. *Escherichia coli* clones were grown until they reached mid log-phase and were induced either with L-arabinose to a final concentration of 0.2% or with *m*-toluic acid at concentrations specified. The induced cultures were grown for 5 and/or 18 h (O/N) as specified in the main text.

#### Transcript analysis

To relatively quantify bla and mcherry transcripts, E. coli RV308 clones were cultivated as described above. For RNA decay studies, E. coli clones with dual UTR constructs were grown until the cultures reached the mid-log phase and were harvested 5 h after induction with 2mM m-toluic acid. Samples for transcript analysis were taken 2, 4, 6, 8 and 10 min after filtration following the wash-out protocol (35). Cell cultures were treated with RNA protect (Qiagen) to stabilize RNA. RNA isolation, DNase treatment, cDNA synthesis and qRT-PCR were performed according to protocols described elsewhere (18). The following primer pairs were used in qPCR: for bla (5'-ACGTTTTCCAATGATG AGCACTT-3' and 5'-TGCCCGGCGTCAACAC-3'); for mCherry (5'-CGTTCGCTTGGGACATCCT-3' and 5'-GATGTCAGCCGGGTGTTT AAC-3'), and for 16S rRNA (5'-ATTGACGTTACCCGCAGAAGAA-3' and 5'-GCTTGCACCCTCCGTATTACC-3'). The primer efficiencies were 94.3, 96.2 and 92.8 for bla, mCherry and 16S rRNA primer pairs, respectively. In each qPCR reactions, non-template controls were performed that show no unspecific amplification. Melting curve analysis was also performed that show a single product for each primer pair.

#### **RBS** calculator details

Translation initiation rates were determined using the reverse engineering function of the RBS calculator (24). The sequence input for the RBS calculator consisted of the 5' UTR DNA sequence (up to 50 nt) and the first 50 nt of the *bla* or *mcherry* coding sequence. 5' UTRs with optimal translational features were generated applying the forward engineering function of the RBS calculator using the following template sequence 5'-GAGCTCCATTA TTATTGTATATGTnnnnnnnnnnnnnnT-3'.

#### Results

### Design, construction and screening of artificial operon plasmid DNA libraries

Initially, we sought to determine whether it would be possible to identify 5' UTR variants that specifically lead either to increased reporter gene expression or protein production. We designed two (bicistronic) artificial operons: pAO-Tr and pAO-Tn (p: plasmid; AO: artificial operon; Tr: transcription; Tn: translation), to identify 5' UTR sequences that specifically lead to increased expression of  $\beta$ -lactamase at the mRNA and protein



Fig. 1 Composition of the artificial operons in plasmids pAO-Tr and pAO-Tn. The inducible XylS/Pm promoter system (indicated by an arrow) drives the expression in both operons, which consists of a 5' UTR; a phosphoglucomutase encoding sequence, celB; a spacer region; a 5' UTR; and a  $\beta$ -lactamase encoding sequence, bla. Two 5' UTR plasmid DNA libraries were constructed using both artificial operons, and the 5' UTR variants identified from pAO-Tr were named Tr-UTRs (a), and 5' UTR variants identified in pAO-Tn were named Tn-UTRs (b). Both artificial operons have transcription terminators on their both ends (indicated by a T). p, plasmid; AO, artificial operon; Tr, transcription; Tn, translation.

levels, respectively. Both artificial operons harbor the phosphoglucomutase encoding sequence, celB, as the first gene; a spacer region; and the  $\beta$ -lactamase encoding sequence, bla, as the second gene (Figure 1). The celB gene was chosen as it can be efficiently transcribed and translated (36), hence would not introduce any undesired restrictions (5); and the bla gene was chosen as host's resistance to ampicillin correlates with the produced amounts of  $\beta$ -lactamase, simplifying the identification of clones with the desired phenotype (18, 37). As for the spacer region, it ensures that the translation of bla only occurs through de novo initiation as opposed to translational read-through (5). The occurrence of de novo initiation was confirmed by eliminating the SD sequence upstream of bla, by replacing the SD sequence GGAG with CCTC, that led to abolished  $\beta$ -lactamase expression (Supplementary Table S1). Expression in both artificial operons is driven by the positively regulated XylS/Pm regulator/promoter system, and a plasmid with the broad-host range mini-RK2 replicon is used as a vector in both constructs (33).

Next, a 34 nt long oligo, with 22 nt degenerate nucleotide composition, was used for the construction of two 5' UTR plasmid DNA libraries. For the transcriptional screening, using pAO-Tr (plasmid maps are provided as Supplementary materials), a plasmid DNA library was constructed by cloning the degenerate oligo upstream of the first gene, *celB* (Figure 1a). Any observed increased expression of  $\beta$ -lactamase (detected as increased ampicillin resistance) would be as a consequence of increased transcription due to the presence of a particular 5' UTR variant. For the translational screening, using pAO-Tn, the same degenerate 5' UTR oligo was cloned upstream of the second gene, *bla* (Figure 1b). Any increased ampicillin resistance observed among the library clones would be as a consequence of increased *de novo* translation of the *bla* gene.

Finally for the screening, the recombinant *E*. coli clones harboring the 5' UTR plasmid DNA libraries ( $\sim$ 280 000 and  $\sim$ 370 000 clones, in pAO-Tr and pAO-Tn, respectively) were plated on agar media containing 0.1 mM *m*-toluic acid (induces transcription from *Pm*) and ampicillin (ranging from 0.5 to 3.5 g/l). Multiple *E*. coli clones were isolated with increased ampicillin resistance phenotype, hypothesized to be either due to increased transcription (led to identification of Tr-UTR variants), or as a

#### (a) Tr-UTR DNA sequences

wt	A	A	С	A	Т	G	Т	-	a	С	a	a	t	a	a	t	a	a	t	g	g	a	g	t	С	a	t	g	a	a	C	A	Т	A	Т	G
LV-2	A	A	С	A	т	G	т	-	•		С	•	•		•		•	С	a	•	•	•		•	t	•	•			•	С	A	Т	A	Т	G
r31	A	A	С	A	т	G	Т	t			С		•	g										•							С	A	Т	A	т	G
r36	A	A	С	A	Т	G	Т	-				•	g	t	•		•		с		•	•		•	a		•		•		C	A	Т	A	Т	G
r50	A	A	С	A	Т	G	Т	t											С										t		С	A	Т	A	Т	G

#### (b) Tn-UTR DNA sequences

wt	A	A	С	A	Т	G	Т	a	с	a	a	t	a	a	t	a	a	t	g	g	a	g	t	С	a	t	g	a	a	С	A	Т	AJ	r G
n24	A	A	С	A	Т	G	Т				t		•	•			t	a									С			С	A	Т	AI	r G
n44	A	A	С	A	Т	G	Т	g										a									С			С	A	Т	<u>A 7</u>	r G
n47	A	A	С	A	т	G	Т		a	t		a		С				a						t						С	A	Т	AT	r G
n58	A	A	С	A	т	G	Т	•			t	•	•		с			a						•			a	t	•	С	A	Т	AT	G



Fig. 2 Nucleotide sequence composition and the phenotype of E. coli clones harboring plasmids with different Tr- and Tn-UTRs. 5' UTR DNA variants were identified by screening pAO-Tr- and pAO-Tn-based 5' UTR libraries for increased ampicillin resistance. The 5' UTR variants r31, r36, r50 are identified from the Tr-UTR library (a); while n24, n44, n47, n58 from the Tn-UTR library (b). The 5' UTR variant LV-2 serves as an internal control that was previously shown to be leading to increase in transcription of the reporter (18). Identical nucleotides are indicated by dots and point mutations are indicated with letters. Nucleotides that are not mutagenized are typed in capital letters including the PciI (ACATGT) and NdeI (CATATG) recognition sequences. The putative SD sequence (ggag) is highlighted in boldface. The ATG start codon (part of the NdeI site) is underlined. The ampicillin resistance phenotypes of the E. coli clones harboring Tr- or Tn-UTR DNA sequences were characterized at the induced state with 0.1 mM mtoluic acid with ampicillin concentrations ranging from 0.5 to 3.5 g/l (c). The low concentration of inducer was used to make sure that resistance levels were in a range allowing us to distinguish moderate phenotypic differences among the clones. Results are presented as averages of the highest ampicillin concentrations at which growth was observed. Error bars point to the next tested ampicillin concentration at which no growth was observed.

consequence of increased translation (led to identification of Tn-UTR variants). Identified clones could grow up to 2.5 g/l ampicillin in the presence of 0.1 mM *m*-toluic acid. To determine the 5' UTR DNA sequences that led to increased ampicillin resistance several clones were selected, plasmid DNAs isolated and sequenced. Among the identified clones (Supplementary Table S2), three Tr-UTRs r31, r36, r50 (Figure 2a) and four Tn-UTRs n24, n44, n47, n58 (Figure 2b) were randomly selected for further characterization along with the wildtype (wt) 5' UTRs, and a previously identified 5' UTR variant, LV-2 (18) (leading to increased transcription), summing up to five 5' UTRs in each category. To ensure that the initially observed ampicillin resistance levels were solely caused by the mutations within the identified 5' UTRs, synthetic oligonucleotides harboring the identified mutations were synthesized and cloned back into pAO-Tr and pAO-Tn, and their ampicillin resistance phenotype were determined (Figure 2c). With the above-mentioned screening and ampicillin resistance characterizations, we concluded that the artificial operons were suitable as a screening tool enabling the identification 5' UTR variants that specifically lead to either increased gene expression or protein production.

#### Design and construction of dual UTR

The screening of the artificial operons gave us access to unique 5' UTR variants. Now having these 5' UTRs, we took a rational design approach and systematically combined them with the objective of benefiting from both the transcriptional and translational characteristic of 5' UTRs in a single dual UTR. A dual UTR consists of two unique 5' UTRs separated by a spacer region: the first 5' UTR proximal to the promoter is where we placed 5' UTRs originating from the transcriptional screening (Tr-UTR); the second 5' UTR is where we placed the 5' UTRs identified from the translational screening (Tn-UTR); while the



**Fig. 3** Schematic view and characterization of ampicillin resistance phenotype in *E*. coli clones harboring plasmids with different dual UTR constructs. The inducible XylS/*Pm* promoter system (indicated by an arrow) drives the expression in all constructs. The dual UTR consists of a Tr-UTR, a spacer region and a Tn-UTR (a). The ampicillin resistance phenotype of the *E*. coli clones harboring each of the 25 dual UTR combinations were characterized using 2 mM *m*-toluic acid to provide full induction (b). Ampicillin concentrations used were ranging from 10 mg/l to 13 g/l. Results are presented as averages of the highest ampicillin concentrations at which growth was observed. Error bars point to the next tested ampicillin concentration at which no growth was observed.

spacer region provides enough space for physical separation of mutations affecting transcription and translation (Figure 3a). With this design concept, we constructed 25 dual UTRs (Figure 3b). In all 25 constructs, the dual UTRs were cloned downstream of the *Pm* promoter, and the entire expression cassette was resting on a plasmid with the broad-host range mini-RK2 replicon.

## Quantification of effects on gene expression and protein production

Initially, the ampicillin resistance phenotype of the *E*. coli clones each harboring one of the 25 dual UTRs were characterized (Figure 3b). The observed ampicillin resistance levels indicated that all the dual UTR constructs were leading to increased reporter gene expression and protein production as compared to expression levels reached with the wtwt dual UTR construct.

A striking finding was that the observed ampicillin resistance phenotype of 17 out of 25 E. coli clones containing the dual UTR constructs was much higher than the resistance levels observed with the E. coli clones containing the individual Tr- and Tn-UTRs on their own (Figure 2c). This initial ampicillin resistance characterization demonstrated that with the dual UTR design concept strong gene expression and high protein production can be achieved. It was also encouraging to observe that high expression levels could be reached by the combination of five Tr- and Tn-UTRs only.

Next, we sought to determine the reporter transcript and protein levels resulting from the dual UTR constructs. Four E. coli clones each harboring one of the dual UTR constructs were grown under induced state and the transcript levels, by relative quantitative real-time reverse-transcription PCR (qPCR); and the enzymatic activities, by  $\beta$ -lactamase enzymatic activity assays, were determined (Figure 4a). The results obtained were in agreement with the design concept that r31, in the r31wt dual UTR construct, led to significant increase in reporter transcript; while n47, in wtn47, led to a significant 10-fold increase in bla transcript and a significant 42-fold increase in  $\beta$ -lactamase enzyme activity compared to the levels of expression reached with the wtwt dual UTR construct. Strikingly, the r31n47 dual UTR led to a significant increase in reporter gene expression and protein production, 46- and 166-fold respectively, compared to the expression levels observed with the wtwt dual UTR construct. For quantitative comparison, we also calculated the translation efficiency for each construct by simply determining the protein-to-transcript ratio using the values presented in Figure 4a. The translation efficiencies were 1, 0.6, 4.1 and 3.6 for the wtwt, r31wt, wtn47 and r31n47 dual UTR constructs, respectively. The observed translation efficiencies were in agreement with the dual UTR design concept that while r31 was leading to increased reporter transcription, hence lower translation efficiency; the n47 was leading to increased reporter translation, hence higher translation efficiency. Taken together, the increased expression observed with the r31n47 dual UTR construct was the result of synergistic effect of 5' UTRs with transcriptional and translational characteristics.

Next, the total cellular protein production was assessed by SDS-PAGE.  $\beta$ -Lactamase could be visualized on the gel in the soluble fraction of sonicated cell lysates from clones harboring the constructs with the wtn47 and r31n47 dual UTRs. The enzyme could also be visualized in the insoluble fraction from the r31n47 dual UTR construct (Figure 4b, upper panel). Specific detection of  $\beta$ -lactamase was also performed by western blotting



Fig. 4 Quantification of *bla* expression levels in *E*. coli clones harbouring plasmids with different dual UTR constructs. (a) Relative *bla* transcript amounts and  $\beta$ -lactamase enzymatic activities (values for the wtwt dual UTR construct were arbitrarily set to 1.0) were determined after 5 hours of growth at the induced state with 2 mM *m*-toluic acid. (b) SDS-PAGE (top) and Western blot (bottom) images of the soluble and insoluble protein fractions of *E*. coli clones producing  $\beta$ -lactamase. Visible  $\beta$ -lactamase bands are highlighted with red rectangles. Error bars indicate the standard deviations calculated from four biological and four technical replicates. The bar values are presented under each bar. Asterisks indicate results of one-tailed T-tests, compared to wtwt construct: \*P-value < 0.04; \*\*P-value < 0.001; \*\*\*P-value < 0.004; \*\*\*P-va



Fig. 5 Quantification of *mCherry* expression levels in E. coli clones harboring plasmids with different dual UTR constructs. (a) Relative *mCherry* transcript amounts and mCherry fluorescent intensities were quantified after 5 h of growth at the induced state with 2 mM *m*-toluic acid. Fluorescence intensities were determined directly from the cultures and normalized with OD<sub>600</sub> values. Values for the wtwt dual UTR construct were arbitrarily set to 1.0. The four images below the panel a show bacterial pellets obtained from the four cultures indicated. Error bars indicate the standard deviations calculated from three biological and three technical replicates. The bar values are presented under each bar. (b) SDS-PAGE image of E. coli clones producing mCherry. Visible mCherry protein bands are highlighted with a red rectangle. Asterisks indicate results of one-tailed T-tests, compared to wtwt construct: 'P-value < 0.02; "'P-value < 0.01; "'P-value < 0.008; """P-value < 0.006; """ + value < 0.000; """ + value < 0.000; "" + va

and the signal strengths correlated with the quantified enzymatic activities (Figure 4b, lower panel).

#### Phenotypic quantification with a red fluorescent protein

The Tn-UTR variants used in the dual UTR constructs were identified using the *bla* gene; therefore, we were interested in testing to what extent the observed increased reporter gene expression was coding sequence (CDS) specific. To assess the potential context dependency, the  $\beta$ -lactamase CDS was substituted with mCherry CDS, encoding for a red fluorescent protein, in wtwt, r31wt, wtn47 and r31n47 dual UTR constructs and mCherry transcript levels, fluorescent intensities and total proteins were quantified. Here, we also calculated the translation efficiencies resulting from the four dual UTR constructs, using the values presented in Figure 5a, and the ratios were: 1, 0.5,

1.2 and 1.4 for the wtwt, r31wt, wtn47 and r31n47 dual UTR constructs, respectively. The phenotypic quantification with mCherry also indicates that r31 leads to a significant increase in gene expression, whereas the n47 leads increase in protein production (Figure 5a). Correspondingly, the mCherry protein could also be easily visualized on an SDS-PAGE, in the soluble fraction only, particularly from the r31n47 dual UTR construct (Figure 5b). These results indicate that the observed significant increase in expression was not CDS-specific. This finding indicates the suitability of the novel 5' UTR design concept for applications aiming for high levels of expression for instance heterologous protein production in bacteria.

#### In silico design of Tn-UTRs

The results reported so far indicate that a combination of 5' UTRs in a dual UTR can lead to strong gene expression and protein production, at least for the two tested selection marker and reporter protein, bla and mCherry in E. coli, respectively. However, context dependency between 5' UTR sequences and CDS cannot be excluded if more genes are to be tested (27). Therefore, we applied a widely used in silico tool, the RBS calculator (24), in designing Tn-UTRs that are specifically designed for the gene of interest to achieve high translation initiation rates (TIR, Supplementary Tables S4 and S5). In the design, the bla and mCherry genes were used as reporters, and E. coli was chosen as the host. In total, six such designed Tn-UTRs were synthesized: three for the  $\beta$ -lactamase and three for the mCherry CDSs (named as dTn-UTRs). The predicted TIR values were 60- to 80-fold higher compared to n47-bla, and 50- to 70fold higher compared to n47-mCherry (Supplementary Table S4). All six dTn-UTRs were cloned into the dual UTR either with the wt or the r31 at the Tr-position (Figure 6). The phenotypic characterizations revealed that all three dTn-UTRs with the bla gene led to a similar increase in expression levels achieved with the n47 combination (Figure 6a). Similar observations were also made for the construct with the *mCherry* gene: Tr-UTR r31 in combination with the dTn-UTRs led to 9- (dTn4) and 46-fold (dTn5 and dTn6) relative increase vs. 58-fold with the Tn-UTR n47 (Figure 6b). These findings indicate that designed 5' UTRs can also be used in the dual UTR context, and even higher expression levels, beyond the predicted levels, can be achieved by the combination of designed Tn-UTRs with Tr-UTRs.

#### Functionality assessment in an alternative host

The broad-host range mini-RK2 replicon and the expression system XvlS/Pm are both known to function in many Gramnegative bacteria (38) including Pseudomonas species. The emerging SynBio chassis, P. putida (28), has the same anti-SD sequence within its 16S rRNA as E. coli. We therefore sought to assess the functionality of the dual UTR concept in P. putida strain KT2440. The dual UTR constructs wtwt, r31wt, wtn47and r31n47 with the mCherry CDS were transferred to P. putida and the mCherry fluorescent intensities were quantified (Figure 7). The strong synergistic effect seen with dual UTR constructs in E. coli was also observed in P. putida. The relative expression levels reached among the dual UTR constructs were weaker in P. putida than the observed relative expression levels in E. coli. The reason for the relative weakness appeared to be due to the reference construct, wtwt, leading to expression of mCherry at much higher levels in P. putida than in E. coli as judged by the stronger bands visualized on the SDS-PAGE gel (Figure 7b). The results obtained in P. putida indicates that the functionality of the dual UTR concept is not only limited to E. coli; hence in principle it can be used in a range of Gram-negative hosts.

## Position specific placement of the Tr- and Tn-UTRs in dual UTR

The conceptualization of dual UTR was the result of a rational design approach. A dual UTR consists of two UTRs: a Tr-UTR,



Fig. 6 Phenotypic characterization of the dual UTR constructs with *in silico* designed Tn-UTRs. (a) The ampicillin resistance phenotype of the *E*. coli clones harboring each of the ten dual UTR combinations were characterized at the induced state with 0.1 mM *m*-toluic acid. The inducible XylS/*Pm* promoter system (indicated by an arrow) drives the expression in both constructs. Error bars point to the next tested ampicillin concentration at which no growth was observed. (b) mCherry fluorescent intensities were quantified after 5 h of growth at the induced state with 2 mM *m*-toluic acid. Fluorescence intensities were determined directly from the cultures and normalized with  $OD_{600}$  values. The fluorescent intensity value for the wtwt dual UTR construct was arbitrarily set to 1.0. Error bars indicate the standard deviations calculated from three biological and three technical replicates. Asterisks indicate results of one-tailed T-tests calculated for the constructs with *mcherry*. The wtTn constructs were compared to the wtwt construct: "P-value < 0.0001; "P-value < 0.0002."



Fig. 7 Quantification of *mCherry* expression levels in *P. putida* clones harboring plasmids with four different dual UTR constructs. (a) Relative mCherry fluorescent intensities were quantified after 5 h of growth at the induced state with 2 mM *m*-toluic acid. Fluorescence intensities were determined directly from the cultures and normalized with  $OD_{600}$  values. The fluorescent intensity value for the wtwt dual UTR was arbitrarily set to 1.0. Error bars indicate the standard deviations calculated from three biological and three technical replicates. The bar values are presented under each bar. (b) SDS-PAGE image of *P. putida* clones producing mCherry. Visible mCherry protein bands are highlighted with a red rectangle. Asterisks indicate results of one-tailed T-tests, compared to wtwt construct: \*P-value < 0.001; \*\*P-value < 0.001. St: Precision Plus Dual Color Protein standard (Bio-Rad); neg ctrl: plasmid-free strain.

placed downstream of the promoter; and Tn-UTR, placed upstream of the start codon. The chosen particular 5' UTRs were rationally placed in Tr- and Tn-UTR positions given their transcriptional and translational characteristics, respectively. This design concept led to the observed synergistic effect resulting in high reporter gene expression and protein production. Observing the synergistic effect on high levels of protein production, we sought to determine whether the enhanced protein production was due to the individual, transcript and translation enhancing, characteristic of the chosen 5' UTRs and whether or not the presence of the functional SD sequence in Tr-UTRs could play a significant role on the observed synergistic effect. To assess the importance of the correct placement of Tr- and Tn-UTRs in dual UTR on the observed phenotype, we constructed dual UTRs by placing the Tn-UTRs, n47 and n58, in the Tr-position; and the Tr-UTRs, r31 and r50, in the Tn-position. After the construction of the plasmids with TnTr dual UTR combinations, we quantified the resulting fluorescent intensities under induced and uninduced states in E. coli (Figure 8). It was intriguing to observe that the dual UTRs with TnTr combinations were not showing the synergistic effect that was seen with TrTn combinations as both the transcript and protein levels were significantly lower. With the observed phenotype of TnTr combinations, we concluded that the presence of a functional SD sequence in Tr-UTR is of no significance and the synergistic effect could only be achieved by the correct placement of the Tr- and Tn-UTRs in dual UTR.

## Functionality assessment with an alternative promoter system

Observing that the dual UTR was functional with another coding sequence, in another host, and the correct positioning of the 5' UTRs was necessary for the observed synergistic effect, we questioned whether the observed increase in gene expression and protein production with dual UTRs was due to an emergent property of the *Pm* promoter. For this functionality assessment, we chose to substitute the XylS/Pm with the AraC/P<sub>BAD</sub> system



Fig. 8 Quantification of *mCherry* expression levels in E. coli clones harboring plasmids with TrTn and TnTr dual UTR constructs. Relative mCherry fluorescent intensities were quantified after 5 h of growth at the induced state with 2 mM *m*-toluic acid. Fluorescence intensities were determined directly from the cultures and normalized with OD<sub>600</sub> values. The fluorescent intensity value for the wtwt dual UTR construct at the uninduced state was arbitrarily set to 1.0. Error bars indicate the standard deviations calculated from three biological and three technical replicates. The bar values are presented under each bar. Asterisks indicate results of one-tailed T-tests, compared to the corresponding TrTn construct (indicated by connecting lines): 'P-value < 0.001; \*\*P-value < 0.001.

in dual UTR constructs, both for the original design of TrTn as well as TnTr combinations. We characterized the phenotype of the E. coli clones by quantifying the fluorescent intensities resulting from each construct under induced and uninduced states after 5 and 18 h (O/N) of growth (Figure 9). The results of this experiment made it clear that the synergistic effect observed in dual UTRs was not an emergent property of the *Pm* promoter as similar expression levels were also reached when we used the  $P_{BAD}$  promoter in combinations. The dual UTRs, both for the TrTn and TnTr combinations. The dual UTR constructs, however, led to much higher background expression from the  $P_{BAD}$  promoter in the absence of induction as compared to the expression levels achieved with the *Pm* promoter.

#### Transcript stability and translational arrest

It has been reported that high translation rate (high ribosome occupancy) occurring on mRNA leads to protection of the mRNA against nucleotide degradation (14–17). Given the high reporter transcript levels observed with the dual UTR constructs, we sought to determine the decay rates of the reporter transcripts originating from the dual UTR constructs, and relative presence of reporter transcripts under translational arrest. For determining the reporter transcript stability, we performed an inducer wash-out assay (35). The wash-out assay relies on the passive diffusion characteristic of the *m*-toluic acid, inducer of the XylS/ Pm system. Escherichia coli clones were grown for 5 h under induced state with 2 mM m-toluic acid. Afterwards the growth medium was replaced with a fresh medium not containing the inducer, and five samples were taken with 2 min intervals. qPCR was performed on the samples and the mRNA decay rates were calculated for the four different dual UTR constructs (Figure 10). The stability of the reporter transcript resulting from the dual



Fig. 9 Quantification of *mCherry* expression levels under the control of the AraC/  $P_{BAD}$  promoter system in E. coli clones harboring plasmids with TrTn and TnTr dual UTR constructs. Relative mCherry fluorescent intensities were quantified after 5 h and 18 h of overnight (O/N) growth at the induced state with L-arabinose to a final concentration of 0.2%. Fluorescence intensities were determined directly from the cultures and normalized with OD<sub>600</sub> values. The fluorescent intensity value for the wtwt dual UTR construct at the uninduced state was arbitrarily set to 1.0. Error bars indicate the standard deviations calculated from three biological and three technical replicates.



**Fig. 10** Quantification of mRNA decay rates in four dual UTR constructs in E. coli. *Escherichia* coli clones were first grown for 5 h under induced state with 2 mM *m*-toluic acid, and afterwards the growth medium was replaced with a fresh medium not containing the inducer, performed for three biological replicates. In total, five samples were taken with 2 min intervals ranging from 0 to 10 min with three technical replicates in each time point. qPCR analyses was performed on the samples and the mRNA decay rates were calculated for the four different dual UTR constructs.

UTRs do not show any significant correlation with the levels of protein production (compared to the mCherry intensity measurements reported in Figure 5). Correlation coefficients for wtwt, r31wt, wtn47 and r31n47 are -0.5, -0.5, -0.3 and 0.59, respectively.

Next, we performed an translational arrest experiment by the addition of chloramphenicol to the growth medium that leads to peptidyl transferase inhibition of bacterial ribosome, hence hinders amino acid chain elongation. For this experiment, the transcript and fluorescence intensities were measured under two conditions: (i) by the induction of expression only with 2 mM *m*-toluic acid, to serve as a control (I); (ii) by the addition of chloramphenicol to a final concentration of  $100 \mu g/$ 



Fig. 11 Quantification of the reporter transcript and mCherry fluorescence intensities under translational arrest. (a) The schematic view of the experimental set-up. *E.* coli clones with different dual UTR constructs were inoculated to a fresh medium and were grown for 2 h. Afterwards, it was either followed by induction of expression only with 2 mM *m*-toluic acid (I) or by the addition of chloramphenicol to a final concentration of 100  $\mu$ g/ml and followed by induction of expression with 2 mM *m*-toluic acid (CI). The quantified relative *mCherry* transcript amounts and mCherry fluorescent intensities were depicted in (b). Fluorescence intensities were determined directly from the cultures and normalized with OD<sub>600</sub> values. Values for the wtwt dual UTR construct at the condition I were arbitrarily set to 1.0. Error bars indicate the standard deviations calculated from three replicates. Asterisks indicate results of one-tailed T-tests, compared to I condition in each construct: '*P*-value < 0.000; \*\**P*-value < 0.0003.

ml and followed by induction of expression with 2 mM m-toluic acid, to assess the accumulation of reporter transcript in the absence of translation (CI) (Figure 11a). The reporter transcript accumulation resulting from the r31n47 dual UTR construct is not negatively affected by the blocking of translation as the transcript levels remain similar as compared to the condition I (Figure 11b). Hence the observed transcript stability, determined by the decay rate experiment, is not due to the high translation observed in the r31n47 dual UTR construct. This same observation also holds true for the wtn47 dual UTR construct. One puzzling observation was the increased fluorescence accumulation resulting from the r31wt construct. The decay rate indicates that the reporter transcript is the least stable compared to the other three reporter transcripts resulting from the dual UTR constructs; however, r31wt still leads to protein accumulation in the presence of chloramphenicol. This observation points to the presence of an unknown biological process that still leads to protein accumulation in the presence of chloramphenicol.

#### Discussion

In this study, we present a novel 5' UTR design named dual UTR. The dual UTR concept both relies and benefits from the

role of 5' UTRs in transcriptional and translational regulation in bacteria. For this study, we design artificial operons to identify 5' UTRs that individually lead to increase in reporter transcription or translation. The combination of 5' UTRs in a single dual UTR leads to a synergistic effect resulting in increased gene expression and protein production, to levels that are beyond the expression levels reached with single 5' UTRs. The observed strong gene expression with dual UTRs represents the importance of the role of 5' UTRs in transcriptional regulation in bacteria. Based on the mRNA decay and translational arrest experiments, we speculate that the reporter transcript accumulations in dual UTR are not, solely, dependent on the translation events, and the Tr-UTRs rather function as an enhancer of transcription. To this end, further research directions will be on the study of single gene expression events by labelled RNA polymerase, and ribosome profiling to understand the role and influence of 5' UTRs in transcript accumulation, especially in relation to ribosome occupancy.

In synthetic biology applications, 5' UTRs are utilized due to their involvement in translational regulation only. Therefore the functional role of 5' UTRs in transcriptional regulation has not been realized in the modular design of biological circuitry. In the circuitry design, different levels of gene expression are achieved either by the use of native promoters, or variants thereof with different strength; or by the use of limited number of regulated promoters via differential induction levels. The use of plasmids with different copy number has also been utilized as an alternative way to achieve varied gene expression. Based on the experimental evidences reported in this study, we envision that the dual UTR concept can also be used as a way to achieve transcriptional regulation in biological circuitry design. This is especially relevant for studies using (unconventional) host microorganisms for which there is a limited number of characterized promoters exist.

The dual UTR design concept is also suitable for applications where the introduction of expression cassettes from plasmidbased systems (multiple copies) to chromosomes (single copy) suffers from the reduction in gene dosage. Hence dual UTRs can be used, even in combination with strong promoters, to achieve maximum transcription rates from a single expression cassette.

To conclude, a 5' UTR involves in complex regulations both at the level of DNA and mRNA. The combination of Tr- and Tn-UTRs in dual UTR leads to a synergistic effect in gene expression and protein production. The functionality of the dual UTR depends on the individual characteristics of the 5' UTRs, and the correct placement of Tr- and Tn-UTRs is crucial for the observed synergistic effect. Dual UTR, as a concept, is functional with different promoters and coding sequences both in *E. coli* and *P. putida*. With these demonstrations, we anticipate that the concept described here are universally applicable to achieve a precise control of expression in Gram-negative bacteria for various synthetic biology applications.

#### SUPPLEMENTARY DATA

Supplementary Data are available at SYNBIO Online.

#### Acknowledgements

Authors would like to thank Laila Berg, for giving valuable technical advice for the library construction and screening; Svein Valla (deceased), for his involvement in an earlier version of the manuscript.

#### Funding

The Faculty of Natural Sciences at the Norwegian University of Science and Technology, Trondheim, Norway (personal PhD stipend that SB received); SINTEF Industry (publication support that SB received); and the Norwegian Research Council with grant numbers 192432 and 192123.

#### **Additional information**

The dual UTR concept has been patented (Application granted on July 24th, 2019).

Conflict of interest statement. None declared.

#### References

- Craig,M.L., Tsodikov,O.V., McQuade,K.L., Schlax,P.E., Capp,M.W., Saecker,R.M. and Record,M.T. (1998) DNA footprints of the two kinetically significant intermediates in formation of an RNA polymerase-promoter open complex: evidence that interactions with start site and downstream DNA induce sequential conformational changes in polymerase and DNA. J. Mol. Biol., 283, 741–756.
- Goldman,S.R., Ebright,R.H. and Nickels,B.E. (2009) Direct detection of abortive RNA transcripts in vivo. Science, 324, 927–928.
- Hsu,L.M., Cobb,I.M., Ozmore,J.R., Khoo,M., Nahm,G., Xia,L., Bao,Y. and Ahn,C. (2006) Initial transcribed sequence mutations specifically affect promoter escape properties. *Biochemistry*, 45, 8841–8854.
- Kudla,G., Murray,A.W., Tollervey,D. and Plotkin,J.B. (2009) Coding-sequence determinants of gene expression in Escherichia coli. Science, 324, 255–258.
- Osterman, I.A., Evfratov, S.A., Sergiev, P.V. and Dontsova, O.A. (2013) Comparison of mRNA features affecting translation initiation and reinitiation. *Nucleic Acids Res.*, 41, 474–486.
- de Smit,M.H. and van Duin,J. (1990) Control of prokaryotic translational initiation by mRNA secondary structure. Prog. Nucleic Acids Res. Mol. Biol., 38, 1–35.
- 7. Hajnsdorf,E. and Boni,I.V. (2012) Multiple activities of RNA-binding proteins S1 and Hfq. *Biochimie*, 94, 1544–1553.
- 8. Mandal,M. and Breaker,R.R. (2004) Gene regulation by riboswitches. Nat. Rev. Mol. Cell Biol., 5, 451–463.
- 9. Coppins,R.L., Hall,K.B. and Groisman,E.A. (2007) The intricate world of riboswitches. *Curr. Opin. Microbiol.*, 10, 176–181.
- 10. Repoila, F. and Darfeuille, F. (2009) Small regulatory non-coding RNAs in bacteria: physiology and mechanistic aspects. *Biol. Cell*, 101, 117–131.
- 11. Kozak, M. (2005) Regulation of translation via mRNA structure in prokaryotes and eukaryotes. *Gene*, 361, 13–37.
- 12. Steitz, J.A. and Jakes, K. (1975) How ribosomes select initiator regions in mRNA: base pair formation between the 3' terminus of 16s rRNA and the mRNA during initiation of protein synthesis in Escherichia coli. Proc. Natl. Acad. Sci. USA, 72, 4734–4738.
- Proshkin, S., Rahmouni, A.R., Mironov, A. and Nudler, E. (2010) Cooperation between translating ribosomes and RNA polymerase in transcription elongation. *Science*, 328, 504–508.
- 14. Yarchuk,O., Jacques,N., Guillerez,J. and Dreyfus,M. (1992) Interdependence of translation, transcription and mRNA degradation in the lacZ gene. J. Mol. Biol., 226, 581–596.
- 15.Kosuri,S., Goodman,D.B., Cambray,G., Mutalik,V.K., Gao,Y., Arkin,A.P., Endy,D. and Church,G.M. (2013) Composability of

regulatory sequences controlling transcription and translation in Escherichia coli. Proc. Natl. Acad. Sci. USA, 110, 14024–14029.

- 16. Boël, G., Letso, R., Neely, H., Price, W.N., Wong, K.-H., Su, M., Luff, J.D., Valecha, M., Everett, J.K., Acton, T.B. et al. (2016) Codon influence on protein expression in *E. coli* correlates with mRNA levels. *Nature*, 529, 358–363.
- 17. Cambray, G., Guimaraes, J.C. and Arkin, A.P. (2018) Evaluation of 244,000 synthetic sequences reveals design principles to optimize translation in *Escherichia* coli. Nat. Biotechnol., 36, 1005–1015.
- 18. Berg, L., Lale, R., Bakke, I., Burroughs, N. and Valla, S. (2009) The expression of recombinant genes in Escherichia coli can be strongly stimulated at the transcript production level by mutating the DNA-region corresponding to the 5'-untranslated part of mRNA. *Microb. Biotechnol.*, 2, 379–389.
- 19. Nouaille,S., Mondeil,S., Finoux,A.-L., Moulis,C., Girbal,L. and Cocaign-Bousquet,M. (2017) The stability of an mRNA is influenced by its concentration: a potential physical mechanism to regulate gene expression. *Nucleic Acids Res.*, 45, 11711–11724.
- Pfleger,B.F., Pitera,D.J., Smolke,C.D. and Keasling,J.D. (2006) Combinatorial engineering of intergenic regions in operons tunes expression of multiple genes. Nat. Biotechnol., 24, 1027–1032.
- 21. Levin-Karp, A., Barenholz, U., Bareia, T., Dayagi, M., Zelcbuch, L., Antonovsky, N., Noor, E. and Milo, R. (2013) Quantifying translational coupling in E. coli synthetic operons using RBS modulation and fluorescent reporters. ACS Synth. Biol, 6, 327–336.
- Mutalik,V.K., Guimaraes,J.C., Cambray,G., Lam,C., Christoffersen, M.J., Mai,Q.-A., Tran,A.B., Paull,M., Keasling,J.D., Arkin,A.P. et al. (2013) Precise and reliable gene expression via standard transcription and translation initiation elements. Nat. Methods, 10, 354–360.
- 23. Na,D., Lee,S. and Lee,D. (2010) Mathematical modeling of translation initiation for the estimation of its efficiency to computationally design mRNA sequences with desired expression levels in prokaryotes. BMC Syst. Biol., 4, 71.
- 24. Salis, H.M., Mirsky, E.A. and Voigt, C.A. (2009) Automated design of synthetic ribosome binding sites to control protein expression. Nat. Biotechnol., 27, 946–950.
- 25.Seo,S.W., Yang,J.-S., Kim,I., Yang,J., Min,B.E., Kim,S. and Jung,G.Y. (2013) Predictive design of mRNA translation initiation region to control prokaryotic translation efficiency. *Metab. Eng.*, 15, 67–74.
- 26.Qi,L., Haurwitz,R.E., Shao,W., DouDNA,J.A. and Arkin,A.P. (2012) RNA processing enables predictable programming of gene expression. Nat. Biotechnol., 30, 1002–1006.

- 27. Cardinale,S. and Arkin,A.P. (2012) Contextualizing context for synthetic biology - identifying causes of failure of synthetic biological systems. *Biotechnol. J.*, 7, 856–866.
- 28. Nikel, P.I. and de Lorenzo, V. (2018) Pseudomonas putida as a functional chassis for industrial biocatalysis: from native biochemistry to trans-metabolism. Metab. Eng., 50, 142–155.
- 29. Sambrook, J., Fritsch, E.F. and Maniatis, T. (1989) Molecular Cloning: A Laboratory Manual, 2nd edn. Cold Spring Harbor Laboratory Press, New York.
- Hanahan, D., Jessee, J. and Bloom, F.R. (1991) Plasmid transformation of Escherichia coli and other bacteria. Methods Enzymol., 204, 63–113.
- 31.Bakke,I., Berg,L., Aune,T.E.V., Brautaset,T., Sletta,H., Tøndervik,A. and Valla,S. (2009) Random mutagenesis of the Pm promoter as a powerful strategy for improvement of recombinant-gene expression. Appl. Environ. Microbiol., 75, 2002–2011.
- 32. Balzer, S., Kucharova, V., Megerle, J., Lale, R., Brautaset, T. and Valla, S. (2013) A comparative analysis of the properties of regulated promoter systems commonly used for recombinant gene expression in Escherichia coli. Microb. Cell Fact., 12, 26.
- 33.Lale,R., Berg,L., Stüttgen,F., Netzer,R., Stafsnes,M., Brautaset,T., Vee Aune,T.E. and Valla,S. (2011) Continuous control of the flow in biochemical pathways through 5' untranslated region sequence modifications in mRNA expressed from the broad-host-range promoter Pm. Appl. Environ. Microbiol., 77, 2648–2655.
- 34. Winther-Larsen,H.C., Blatny,J.M., Valand,B., Brautaset,T. and Valla,S. (2000) Pm promoter expression mutants and their use in broad-host-range rk2 plasmid vectors. *Metab. Eng.*, 2, 92–103.
- 35. Kucharova, V., Strand, T.A., Almaas, E., Naas, A.E., Brautaset, T. and Valla, S. (2013) Non-invasive analysis of recombinant mRNA stability in Escherichia coli by a combination of transcriptional inducer wash-out and qRT-PCR. PLos One, 8, e66429.
- 36. Blatny, J.M., Brautaset, T., Winther-Larsen, H.C., Karunakaran, P. and Valla, S. (1997) Improved broad-host-range RK2 vectors useful for high and low regulated gene expression levels in gram-negative bacteria. *Plasmid*, 38, 35–51.
- 37. Heggeset, T.M.B., Kucharova, V., Nærdal, I., Valla, S., Sletta, H., Ellingsen, T.E. and Brautaset, T. (2013) Combinatorial mutagenesis and selection of improved signal sequences and their application for high-level production of translocated heterologous proteins in *Escherichia coli*. Appl. Environ. Microbiol., 79, 559–568.
- Lale, R., Brautaset, T. and Valla, S. (2011) Broad-host-range plasmid vectors for gene expression in bacteria. Methods Mol. Biol. 765, 327–343.