
DOI: [http://dx.doi.org/10.1021/acssynbio.6b00065](http://dx.doi.org/10.1021/acssynbio.6b00065)

Copyright:

The final publication is available at ACS Publications via [http://dx.doi.org/10.1021/acssynbio.6b00065](http://dx.doi.org/10.1021/acssynbio.6b00065)

Date deposited:

18/07/2016

Embargo release date:

20 May 2017

This work is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported License

Newcastle University ePrints - eprint.ncl.ac.uk
Effect of genome position on heterologous gene expression in *Bacillus subtilis*; an unbiased analysis

Christopher Sauer¹,²,⁴, Simon Syvertsson¹,⁴, Laura C. Bohorquez³, Rita Cruz¹,², Colin R. Harwood¹, Tjeerd van Rij², Leendert W. Hamoen¹,³,⁵

¹ Centre for Bacterial Cell Biology, Institute for Cell and Molecular Biosciences, Newcastle University, Richardson Road, NE2 4AX Newcastle, United Kingdom
² DSM Biotechnology Center, P.O. Box 1, 2600 MA Delft, The Netherlands
³ Bacterial Cell Biology, Swammerdam Institute for Life Sciences, University of Amsterdam, De Boelelaan, 1081 HZ Amsterdam, The Netherlands
⁴ Both authors contributed equally
⁵ Corresponding author: Leendert W. Hamoen, University of Amsterdam, l.w.hamoen@uva.nl

Keywords: *Bacillus subtilis*, gene expression, genome location, transposon, GFP, LacZ
A fixed gene copy number is important for the in silico construction of engineered synthetic networks. However, the copy number of integrated genes depends on their genomic location. This gene dosage effect is rarely addressed in synthetic biology. Two studies in *Escherichia coli* presented conflicting data on the impact of gene dosage. Here, we investigate how genome location and gene orientation influences expression in *Bacillus subtilis*. An important difference with the *E. coli* studies is that we used an unbiased genome integration approach mediated by random transposon insertion. We found that there is a strong gene dosage effect in fast growing *B. subtilis* cells, which can amount to a 5-fold difference in gene expression. In contrast, gene orientation with respect to DNA replication direction does not influence gene expression. Our study shows that gene dosage should be taken into account when designing synthetic circuits in *B. subtilis* and presumably other bacteria.
INTRODUCTION

*Bacillus subtilis* is widely used as bacterial model system and for the production of enzymes and vitamins. It is genetically tractable, easily culturable and regarded as safe. This has resulted in the development of a wide array of genetic tools for use in both academic and industrial environments, and has made the organism a popular cloning chassis in synthetic biology. The natural genetic transformation system of *B. subtilis* facilitates the integration of DNA into the genome by homologous recombination. In contrast to the use of plasmids as genetic carriers, genome integration assures that heterologous genetic material is stably maintained at a copy number that reflects that of the chromosome. The latter is important when genetic modules (e.g. biobricks) are used for the construction of engineered synthetic metabolic pathways and regulatory networks. However, since DNA replication originates at a fixed position on the genome (origin of replication), and rapidly growing bacteria initiate new rounds of replication before the previous rounds have been completed, the copy number of integrated genes will fluctuate. As a result, gene dosage rises with increasing proximity to the origin of replication. Another aspect that might affect the activity of integrated genes is the transcriptional orientation. The majority of native genes are co-directional with DNA replication in order to reduce collisions between RNA polymerase and the DNA replication machinery. It is therefore possible that the orientation of integrated genes will influence expression. In this study, we set out to determine to which extent genome location and gene orientation influences heterologous gene expression in *B. subtilis*.

Several groups have studied the effects of genome location on gene expression using the model organism *Escherichia coli*. By varying the position of a reporter gene on the genome, Block and coworkers have shown that expression is only influenced by gene dosage and not by the orientation of the gene relative to the direction of DNA replication. They found that in rapidly growing cultures of *E. coli* the maximum expression levels were approximately 9-fold higher for gene locations close to the origin of replication compared to locations close to the replication terminus. More recently, Bryant and coworkers reported that gene expression can vary up to 300-fold for a reporter fusion that was integrated at different locations on the *E. coli* genome. Moreover, they concluded that gene dosage
had only a minor influence on expression, and that the observed differences are related to specific features involved in chromosome organization. The discrepancy between these reports may therefore reflect the particular locations chosen for the integration of the reporter fusions.\textsuperscript{12, 13} To prevent this complication, we developed an unbiased reporter integration approach for \textit{B. subtilis}.

Instead of choosing the insertion sites of reporter genes, we used a transposon-mediated random insertion approach and selected clones exhibiting a wide range of expression activities. Following this approach, we found that expression of a \textit{lacZ} reporter gene can differ up to 5-fold based on its chromosomal location. This difference in expression correlated strongly with the location of the reporter cassette relative to the origin of replication, and was not influenced by gene orientation with respect to the DNA replication direction. Thus, gene dose is an important factor influencing heterologous gene expression in \textit{B. subtilis}, and this should be taken into account when designing \textit{in silico} synthetic regulatory circuits, and also when expressing commercially valuable heterologous proteins.
RESULTS & DISCUSSION

Unbiased chromosome insertion

To be able to make an unbiased assessment of the influence of genomic location on gene expression in *B. subtilis*, we made use of the Mariner transposon. This transposon only requires a TA dinucleotide sequence for transposition and consequently it is known to insert randomly into the genome. To determine the influence of gene orientation on expression, a bidirectional reporter cassette was constructed comprising the constitutively active $P_{veg}$ promoter and the inducible $P_{spac}$ promoter, fused, respectively to *gfp* and *lacZ* reporter genes (Figure 1). To control the $P_{spac}$ promoter, the *lacI* repressor gene was integrated into the genome at the *aprE* locus. Transcriptional read-through from and into adjacent genes was avoided by the inclusion of strong bidirectional terminators at both ends of the cassette (Figure 1). These bidirectional *B. subtilis* terminators were identified using the WebGeSTer database. The reporter cassette was subsequently cloned into the Mariner transposon. Despite the size of the cassette (~6 kb), the transposon was still active, albeit with a transposition frequency that was an order of magnitude lower than the wild type transposon (Supporting Information Table S1).

Following transposition, cells were grown on nutrient agar plates containing X-gal and IPTG, and colonies showing a range of color intensities, from dark blue to almost white, were selected at random for detailed analysis. To ensure that the transposon insertions did not influence growth, the growth rates of the selected clones were measured in microtiter plates. Fourteen clones that showed normal growth rates (Supporting Information Figure S1) were selected for further analyses. The locations of the transposon insertions on the *B. subtilis* genome are shown in Figure 2 and detailed genetic information about these loci is provided in the Supporting Information (Figure S2).

Effect of genome position

The fourteen transposon insertion strains were grown in LB medium at 37 °C in the presence of 1 mM IPTG, and samples were taken during exponential growth ($OD_{600}$~0.5) to determine β-galactosidase activity and fluorescence intensity (GFP) measurements. Figure 3A shows the β-galactosidase activity
and GFP fluorescence intensity with respect to the locations of the transposons on the genome. The orientation of the reporter genes is indicated by arrowheads. The trend lines show a clear correlation between chromosome location and gene expression for both reporters. The maximum difference in expression between a gene located close to the replication origin (oriC) or to the terminus (ter) is approximately 5-fold. The differences in GFP expression levels are clearly visible by eye (Figure 3B). The experiment was carried out in triplicate and a representative set of data is shown in the Supporting Information (Figure S3A).

To determine whether the observed activities were influenced by read-through from adjacent genes, β-galactosidase activities were also measured in the absence of IPTG. In the eight transposon clones tested, the absence of IPTG resulted in very low β-galactosidase activities, maximally of 6 Miller Units (Supporting Information Figure S4), indicating that the reporter cassette is transcriptionally isolated by the flanking bidirectional terminators, and therefore not influenced by transcriptional read-through from adjacent genes.

It has been reported that in B. subtilis the copy number of a gene close to oriC can be ~5 times higher than that of a gene close to the ter (ori:ter ratio) when growing exponentially (~2.5 doublings/h) in LB medium. To examine the relation between expression and gene dosage, DNA copy numbers of the different loci were determined using qPCR (Figure 3C). We found a comparable ori:ter ratio as described before, and a distribution of DNA copy numbers of the different loci that reflects the trends of lacZ and gfp expression, indicating that gene dosage is a major factor influencing gene expression in fast growing cells.

**Outliers**

Despite the clear relationship between expression levels and the genomic location of the reporters, a few transposon insertions deviated from the trend lines (see arrows Figure 3A). For example, the transposon integrated close to oriC (~1°) showed a much lower β-galactosidase activity than expected, even though the sequences of the lacZ gene and Pspac promoter did not reveal any mutations. This transposon was inserted into the non-essential ribosomal gene rmO-23S that is part of the rmO-16S operon (Supporting Information Figure S2). Possibly, the terminator of the transposon cassette is not
fully able to prevent read-through from the strongly transcribed rmO-16S operon. The transposon insertion at 332° (spsC) showed a significantly lower than expected GFP activity for reasons that are currently unknown (Figure 3A).

**Effect of transcription direction**

Like most other bacteria, the orientation of genes on the *B. subtilis* genome is strongly correlated with the direction of DNA replication.\(^7, 9\) To determine whether gene orientation affected expression, β-galactosidase activities and GFP fluorescence intensities were analyzed with respect to their orientation, as indicated in Figure 4. The expression levels were plotted against the distance of the transposon insertion sites from the origin of replication. The outliers at 1° and 332° were discarded in this analysis. The resulting trend lines are comparable (Figure 4), indicating that gene orientation did not have a strong influence on expression levels. One replicate is shown in Supporting Information Figure S3B-C. At first sight this might seem surprising considering the strongly biased orientation of the genes on the genome, however, this bias is likely related to the negative effect of RNA polymerases colliding with the DNA replication machinery, resulting in less efficient DNA replication rather than effecting transcription efficiency.

**Effect of growth rate reduction**

*B. subtilis* grows considerably slower in minimal medium containing glucose and amino acids than in nutritionally rich LB medium.\(^21, 22\) This is also reflected in a lower ori:ter ratio which in minimal medium approaches 3:1.\(^20\) If expression differences are indeed due to gene dosage effects, this should be reflected in a reduced expression ratio when grown in minimal medium. To test this, four transposon strains were selected; two with the reporter cassette close to the terminus (181° and 191°) and two with the reporter cassette close to the origin of replication (5° and 342°). The strains were grown in LB medium and minimal medium with mean generation times at 37°C of 27 min and 45 min, respectively. Indeed, growth in minimal medium reduced the difference in lacZ expression from around 3.5-fold to 2.5-fold compared to LB (Figure 5). However, the gfp marker, for unknown reasons, did not show a clear reduction in expression.
Murray and Koh have shown that the ori:ter ratio is similar when *B. subtilis* is grown in LB medium at 30°C and 37°C, despite the mean generation time at 30°C (50 min) being virtually double that at 37°C (27 min). This appears to be reflected in the unchanged expression activities of both marker genes (Figure 5).

**Expression differences in stationary phase**

Since DNA replication is markedly reduced when exponential growth ceases, the differential expression of different transposon strains should decrease as well. However, when we measured β-galactosidase activities and GFP intensities during stationary phase, the differential expression profiles with respect to location were still observed (Figure 6A & B. Replicate shown in Figure S5). Both β-galactosidase and GFP are stable proteins and a possible explanation might be that the observed trends are still visible in the stationary growth phase as a consequence of the difference in reporter protein expression during logarithmic growth. To test this, *lacZ* was induced only in the stationary phase (Figure 6A) and, as expected, the observed β-galactosidase levels were much smaller compared to the induction of *lacZ* during logarithmic growth (compare Y-axis scales). More importantly, the difference in *lacZ* expression between the transposon strains was almost completely abolished (Figure 6C). To confirm that this effect is a consequence of a reduction in gene dosage, the DNA copy numbers of the different loci were determined using qPCR (Figure 6D). Again the distribution of DNA copy numbers followed closely the trend observed for *lacZ* expression.

**Conclusion**

Our unbiased selection approach shows that gene dosage as a consequence of multiple replication forks plays a considerable role in gene expression in *B. subtilis*, which is in full agreement with the study of Block and coworkers, but does not support the conclusions from the study by Bryan and coworkers. Similar to Block and coworkers, we also found that the orientation of reporter genes has no influence on expression levels. The importance of gene dosage in *B. subtilis* is not restricted to heterologous proteins as was shown in two recent studies that revealed the impact of chromosomal location of genes involved in the regulation of motility, biofilm development and sporulation.
Clearly, gene dosage effects have to be taken into account when locating synthetic circuits in bacteria. Moreover, a potential 5-fold increase in enzyme production represents a commercially significant improvement. In fact, our transposon reporter system might be adapted to improve the production of enzymes that are expressed from the genome, since we show that even large DNA constructs (at least 6 kb) can be randomly inserted into the *B. subtilis* genome by the Mariner transposon. In light of the broad host-range of the Mariner transposon, such strategy is especially useful for production organisms for which no convenient genome recombination system is available.
METHODS

Bacterial strains and growth conditions

For cloning purposes E. coli DH5α was used. The GFP and β-galactosidase experiments were performed in B. subtilis strain W168 (lacA::tet aprE::lacI, cat). Cells were grown in Luria-Bertani (LB) medium with the addition of 10 µg/ml kanamycin for overnight cultures. No antibiotics were added when cells were assayed. Minimal medium was based on Spizizen’s minimal medium and consists of 2 g/l (NH₄)₂SO₄, 14 g/l K₂HPO₄, 6 g/l KH₂PO₄, 1 g/l sodium citrate, 2 g/l MgSO₄, 5 g/l glucose, 2 g/l tryptophan, 0.2 g/l casamino acids, and 2.2 g/l ammonium ferric citrate. Pre-cultures were performed by growing cells from the overnight culture in fresh media until they reached exponential phase. Then all cultures were diluted to OD₆₀₀ ~ 0.05 and grown until the samples reached an OD₆₀₀ ~ 0.5 and 3.0.

Plasmid and strain construction

A unique cloning site was created in pMarB by linearizing the plasmid with the PCR primer pair oSS330/oSS331 (Supporting Information Table S1). The PCR product was digested with MunI and ligated, resulting in plasmid pSS121. The reporter cassette was assembled using splicing by overlap extension PCR. GFP was amplified from pSG1729-gfp using oligo primer pair oSS346/oSS258, Pvet was amplified from chromosomal DNA of B. subtilis W168 with primer pair oSS259/oSS260, and lacZ was amplified from pMUTIN4 with primer pair oSS261/oSS262. The Pspac promoter was introduced by overhangs in primers oSS260 and oSS261. The three fragments were joined stepwise using oSS346 as the terminal gfp primer, and oSS347 as the terminal lacZ primer, which also introduced BamHI sites at both ends of the final construct. The components were first assembled into pUC19. The bidirectional transcriptional terminator ezrA/braB was PCR amplified from the chromosomal DNA of B. subtilis W168 using primer pair oSS344/oSS345 containing SacI restriction sites, and subsequently cloned into the SacI site of pUC19 resulting in pSS116. The BamHI site of pSS116 was used to introduce the Pvet-gfp/Pspac-lacZ fragment (pSS117). A bidirectional transcriptional terminator was incorporated into the Sphi site of pSS117 by amplifying the ywoG/ywoF terminator using primer pair oSS348/oSS349 (pSS118). Both terminators were identified using the WebGeSTer database. EcoRI and HindIII sites
were used to excise the reporter cassette from pSS118 and for cloning into *MunI* digested pSS121, after the insert and vector were blunt-end following a treatment with the Klenow fragment of DNA polymerase I. The resulting transposon vector was called pSS125 (Supporting Information Figure S6).

**Transposon selection**

The transposon procedure was based on that described by Lampe and Le Breton.14,15 *B. subtilis* W168 was transformed with pSS125 and transformants were selected on erythromycin containing plates at 30°C. A few colonies were taken and cultured in LB medium for 3-5 hours at 30°C and spread on nutrient agar plates with kanamycin at several dilutions from 10⁻³ to 10⁻⁶. The plates were incubated overnight at 50°C, the non-permissive temperature for the plasmid, favoring the integration of the $P_{veg^-}gfp/P_{spac^-}lacZ$ fragment and the kanamycin-resistance cassette by the plasmid-encoded transposase. Undesired strains that still contained the plasmid were identified via the screening of plates containing erythromycin positive clones. Strains which were only resistant to kanamycin were used for further experiments. Transposon insertion efficiencies were determined by comparing the number of colonies with the desired kanamycin resistance, revealing a successful integration of the expression cassette, with the number of colonies having both kanamycin and erythromycin resistance, reflecting the presence of the plasmid (Supporting Information Table S2). Colonies were plated on nutrient agar plates containing 40 µg/ml of X-gal and 10 µM IPTG to select clones by color.

To determine the transposon integration sites, arbitrary PCR was used as described by Knobloch.27 In short, PCR reactions were performed using GoTaq polymerase (Promega) and primer pairs Arb1/MarB1 or Arb1/MarB2. The thermocycler program began with an initial denaturation at 95°C for 5 min, 6 cycles at 95°C for 1 min, 30°C for 1 min, 72°C for 1 min, 30 cycles at 95°C for 30 sec, 50°C for 30 sec, 72°C for 1 min and a final extension at 72°C for 7 min. In the second PCR step primer pairs Arb2/MarB1 or Arb2/MarB2 used 2 µl of the first reaction as a template. For the 2nd PCR the following programme was used: 95°C for 5 min, 6 cycles at 95°C for 30 sec, 50°C for 30 sec, 72°C for 30 sec, 30 cycles at 95°C for 30 sec, 58°C for 30 sec, 72°C for 30 sec, and a final extension at 72°C for 7 min. PCR products were sequenced using primer MarB1N or MarB2N.
β-galactosidase and GFP assays

To measure β-galactosidase activities, cells were harvested and flash frozen using liquid nitrogen and stored at -80°C. β-galactosidase activity assays were performed according to the standard protocol of Miller,\textsuperscript{28} whereby cells were first lysed by adding 20 mg/ml lysozyme at room temperature for 30 min.

To determine GFP levels, cells were washed in PBS after sampling. A sample of 0.3 µl was applied to a GeneFrame (AbGene, Surrey, UK) containing 1.25% agarose in PBS, supplemented with 2 µg/ml of DAPI. For microscopic measurements, a Zeiss 200M microscope was used with the following excitation filters and exposure times: GFP (470/525) for 500 ms; DAPI (350/460) for 1000 ms. Microscopy images were analyzed using in-house software (NucTracer) that uses the DAPI stained nucleoid as identifier for the region of interest to measure GFP fluorescence in the cell. NucTracer is based on the ImageJ plugin ObjectJ that supports graphical vector objects identifying images on a transparent layer.\textsuperscript{29, 30}

DNA copy numbers analysis

DNA copy number analysis was performed as described by Murray and Koh.\textsuperscript{20} In short, chromosomal DNA was isolated during exponential (OD\textsubscript{600} ~0.5) and stationary phase (OD\textsubscript{600} ~3.0). As a control for single copy number, DNA from \textit{B. subtilis} spores was isolated. Regions of origin (oriC) and terminus (ter) of replication and the transposon locations were amplified using the primers listed in Supporting Information Table S2. qPCR was performed using iQ™ SYBR® green supermix (Bio-Rad) in a C1000 thermal cycler (CFX96 real-time system, Bio-Rad). A relative quantification analysis (ΔΔC\textsubscript{q}) was performed by determining quantification cycles (C\textsubscript{q}) to calculate the ratios of transposon location compared to ter and by normalizing to the ori:ter ratio of chromosomal DNA of spores that contain one chromosome reflecting an equivalent ratio of 1. Two loci were excluded from the analysis (1° and 241°) due to primer inefficiencies during qPCR.

SUPPORTING INFORMATION

Table S1: Transposon insertion efficiency of the expression cassette comparing pSS125 to pMarB
Table S2: Primers used in this study
Figure S1: Growth rate measurement of transposons
Figure S2: Overview of the chromosomal transposon insertion sites
Figure S3: Effect of chromosomal location and transcription direction on gene expression
Figure S4: Transcriptional read-through measured by β-galactosidase assays in the absence of IPTG
Figure S5: Expression differences in stationary growth
Figure S6: Plasmid map of pSS125

ACKNOWLEDGEMENTS
C.S. and R.C. were supported by the European Commission funded Marie Curie Initial Training Network ATRIEM (project No. 317228). S.S. was supported by a BBSRC DTG PhD studentship. L.C.B was supported by the European Commission funded Marie Curie Innovative Training Network AMBER (project No. 317338), and L.W.H was supported by an NWO STW-Vici (12128) grant.
REFERENCES


FIGURE LEGENDS

Figure 1. Schematic overview of the Mariner transposon containing the bidirectional reporter cassette. The housekeeping SigA-dependent P_{veg} promoter and the inducible P_{spac} promoter are fused to gfp and lacZ, respectively, and located between strong intrinsic transcription terminators (T) to prevent read-through from and into adjacent genes. The transposons are selected by kanamycin resistance (kan^{R}). The inverse terminal repeats (ITR) used by the transposase are indicated. The lacI gene coding for the repressor of the P_{spac} promoter is integrated at the genomic aprE locus (not indicated).

Figure 2. Chromosomal location of selected transposon mutants. Arrows indicate the orientation of the gfp marker and the location is given in degrees (1° ≡ 11,708 base pairs). The origin of replication (oriC) and replication terminus (ter) are indicated. In B. subtilis the ter region is located approximately 178.7° on the B. subtilis genome. DNA replication directions are indicated.

Figure 3. Effect of chromosomal location on gene expression. Cells were grown in LB medium at 37°C and samples were taken during exponential growth (OD_{600} ≈ 0.5). (A) β-galactosidase activities (blue) and GFP fluorescence (red) are plotted relative to the chromosomal locations (degrees) of the transposons. Error bars indicate standard deviations of two technical replicates in case of β-galactosidase and at least 100 cells for GFP. The triangles indicate transcriptional direction of lacZ or gfp. Two outliers are marked by grey arrows. Goodness of fit for β-galactosidase with R^2 = 0.86 (excluding locus at 1°) and GFP with R^2 = 0.70. (B) Visual differences in GFP expression illustrated by fluorescence micrographs of four different transposons. Scale bar 5 µm. (C) DNA copy numbers of transposon locations determined by qPCR. Error bars indicate standard deviations of three biological replicates. Goodness of fit with R^2 = 0.78.

Figure 4. Effect of gene orientation on gene expression. β-galactosidase activities (A) and GFP fluorescence intensities (B) from Figure 3 are plotted against the distance of the transposon insertion
sites from the origin of replication (oriC). Triangles indicate the transcriptional direction of the reporter genes. Reporter genes that are transcribed in the same direction as DNA replication are shown in blue and genes that are transcribed against the DNA replication direction are shown in red. Error bars indicate the standard deviation of two technical replicates in case of β-galactosidase and at least 100 cells for GFP. Goodness of fit for β-galactosidase with $R^2 = 0.60$ (blue), $R^2 = 0.91$ (red) and for GFP with $R^2 = 0.82$ (blue), $R^2 = 0.66$ (red).

**Figure 5.** Influence of growth conditions on gene expression. The fold expression differences of β-galactosidase and GFP under different growth conditions were calculated by dividing the average expression levels of two loci close to the origin by that of two loci close to the terminus of replication. Cells were grown in LB medium (blue) and minimal medium (green) at 37°C, and in LB medium at 30°C (red). Error bars indicate the standard deviation of three biological replicates.

**Figure 6.** Reporter gene expression in stationary growth. (A) Transposon strains were grown in LB at 37°C and lacZ expression was induced with 1 mM IPTG at the start of growth ($t_0$) after which cells were harvested 3.5 hours later ($t_1$), or at the beginning of the stationary phase ($t_1$) after which cells were harvested 2 hours later ($t_2$). (B) β-galactosidase activities (blue) and GFP fluorescence (red) are plotted against chromosomal location (degrees) of the transposons when 1 mM IPTG was added at the start ($t_0$) of growth. Error bars indicate the standard deviation of two technical replicates in case of β-galactosidase and at least 100 cells for GFP. Goodness of fit for β-galactosidase with $R^2 = 0.77$ (excluding locus at 1°) and GFP with $R^2 = 0.80$. (C) β-galactosidase activities when 1 mM IPTG was added in the beginning of the stationary growth phase ($t_1$). Average values with standard deviations of three independent biological replicates are shown. Direction of the triangles indicates transcriptional direction of lacZ or gfp. Goodness of fit for β-galactosidase with $R^2 = 0.34$. (D) DNA copy numbers of transposon locations determined by qPCR. Error bars indicate standard deviations of three biological replicates. Goodness of fit with $R^2 = 0.42$. 
Figure 2

Bacillus subtilis
W168

oriC

Replication

ΔrecJ 241°
ΔyqeT 224°
ΔyqhS 217°
ΔbshB2 181°
Δyof 181°
Δbxa 196°
ΔspeA 131°
Δycg 52°
ΔckB 31°
Δtc 5°
ΔsuG 342°
Δspsc 332°
Δvmb 308°
ΔmO-23S 1°
Figure 3

**A**

![Graph](Image)

- **β-gal activity (100-MU)**
- **GFP (AU)**

**B**

![Images](Image)

- **P_{wgr/gfp}**
- Chromosome position (°)

**C**

![Graph](Image)

- **DNA copy number**
- Chromosome position (°)
Figure 5