Evaluation of chromosomal insertion loci in the Pseudomonas putida KT2440 genome for predictable biosystems design

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A B S T R A C T

The development of Pseudomonas strains for industrial production of fuels and chemicals will require the integration of heterologous genes and pathways into the chromosome. Finding the most appropriate integration site to maximize strain performance is an essential part of the strain design process. We characterized seven chromosomal loci in Pseudomonas putida KT2440 for integration of a fluorescent protein expression construct. Insertion in five of the loci did not affect growth rate, but fluorescence varied by up to 27-fold. Three sites displaying a diversity of phenotypes with the fluorescent reporter were also chosen for the integration of a gene encoding a muconate importer. Depending on the integration locus, expression of the importer varied by approximately 3-fold and produced significant phenotypic differences. This work demonstrates the impact of the integration location on host viability, gene expression, and overall strain performance.

1. Introduction

Pseudomonas putida has become a desirable target organism for the production of industrial products due to several attributes of its native physiology (Ankenbauer et al., 2020). P. putida has a naturally diverse catabolism with the capacity to metabolize aliphatic, aromatic and heterocyclic compounds in addition to glucose (Schmid et al., 2001), lending to catabolism, with the capacity to metabolize aliphatic, aromatic and heterocyclic compounds in addition to glucose. An unusual cyclic pathway organization for catabolism of hexose sugars provides chemical production. An unusual cyclic pathway organization for catabolism of hexose sugars provides chemical production.

Genetic modification of pseudomonads for industrial bioprocessing requires the ability to stably express heterologous genes and pathways. Furthermore, it is important that heterologous constructs be integrated into the chromosome (Tyo et al., 2009), as plasmid maintenance requires the maintenance of a plasmid-specific selection pressure such as the use of antibiotics. Generating the correct expression stoichiometry of the heterologous genes is equally important and can be challenging. Design parameters used to control protein expression are typically gene copy number (Morlino et al., 1999), promoter (Samuelson, 2011), and terminator type (Curran et al., 2015). However, the chromosomal integration location can affect heterologous protein expression independently of these parameters. The three dimensional structure of the chromosome (Krogh et al., 2018), the order, and the orientation of the genes play a role in determining the expression of each gene (Slager and Veening, 2016). The physical bending and condensing of the chromosome can bridge delocalized genes together (Le et al., 2013), possibly contributing to the rate and order of transcription, and thus, the expression of each gene. DNA supercoiling in response to environmental factors has been shown to affect protein expression (Hulton et al., 1990), and to play a major role in the evolutionary conservation of gene order in bacteria (Sobetzko, 2016). The physical bending and condensing of the chromosome can bridge delocalized genes together (Le et al., 2013), possibly contributing to the rate and order of transcription, and thus, the expression of each gene. DNA supercoiling in response to environmental factors has been shown to affect protein expression (Hulton et al., 1990), and to play a major role in the evolutionary conservation of gene order in bacteria (Sobetzko, 2016). Integrating a heterologous segment of DNA can potentially disrupt this sequence, order and topology, disabling the ability of the host to grow by changing essential gene expression patterns.
Transcription into and out of the heterologous segment may also occur, altering native protein expression. Additionally, appropriate expression of the heterologous DNA could be altered due to interference by regional protein and DNA interactions. Introduction of heterologous DNA into the chromosome would benefit from the characterization of insertion loci in which the expression of the heterologous DNA can be optimized without disrupting the natural sequence and topology of the chromosome that is required for normal cell functioning.

The characterization of chromosomal integration loci for this purpose has been explored to various degrees in several organisms such as Escherichia coli (Sousa et al., 1997; Bassalo et al., 2016), Bacillus subtilis (Sauer et al., 2016), Saccharomyces cerevisiae (Flaggeldt et al., 2009; Yamane et al., 1998), Lactococcus lactis (Thompson and Gasson, 2001), P. putida KT2440 (Dormöse et al., 2019), and Dro sophila melanogaster (Markstein et al., 2008). In this study, we have rationally selected seven insertion loci on the P. putida KT2440 chromosome to test for the expression of a fluorescent reporter gene and the effect of the insertion of the reporter on the growth of the host. A transmembrane transporter protein for the polymer precursor cis,cis-muconate was integrated into these three sites to further verify the effect of the insertion site on protein expression and organizational phenotype.

2. Methods and materials

2.1. Integration site selection

Insertion regions for heterologous gene insertion were manually curated from the P. putida KT2440 genome that were not adjacent to essential genes, were not between genes within the same operon, and were in anti-parallel orientation from the downstream gene. These sites were identified using the BioCyc P. putida KT2440 Tier 3 Pathway/Genome Database (PGDB) (Karp et al., 2019). The selected insertion regions were then categorized based on length, short (<50 bp), medium (>50 bp and <110 bp), or long (>110 bp), and then finally down-selected to a list of 7 insertion regions to encompass a range of all three lengths, representing subsections of the full genome. Integration location sites were named based on the upstream gene identifier.

2.2. Plasmid design and assembly

Plasmids were designed for integration into Pseudomonas putida KT2440 by homologous recombination, inserting the heterologous DNA without removing any endogenous genome sequence. The mKate fluorescent reporter gene and the flanking 400 nucleotide base pair sequences upstream and downstream of each integration site was synthesized by Integrated DNA Technologies (IDT) in Coralville, IA and inserted into the pK18mob-sacB suicide vector carrying a kanamycin resistant cassette and sacB gene which confers sensitivity to sucrose (Elmore et al., 2017; Johnson and Beckham, 2015; Marx, 2008) using NEBuilder HiFi DNA Assembly Master Mix (New England Biolabs (NEB), Ipswich, MA). Plasmid constructs were transformed into NEB S-alpha FF (NEB) Escherichia coli competent cells by heat shock, plated onto LB agar plates supplemented with 50 μg/mL kanamycin sulfate, and incubated at 37 °C for 16–24 h, or until distinct single colony formation was visible. PCR was conducted on single colonies to identify the presence of successfully assembled plasmid transformants using primers synthesized by IDT and the Quick-Load Taq 2X Master Mix (NEB) system. Positive colonies were transferred into LB liquid medium supplemented with 50 μg/mL kanamycin sulfate and incubated at 37 °C for 16–24 h. Surviving colonies were then streaked onto fresh YT-25% sucrose plates and incubated at 30 °C for 48 h. Surviving colonies were then re-streaked onto fresh YT-25% sucrose plates and incubated at 30 °C for 16 h to remove any enduring sacB-containing cells. All final colonies were screened for complete mKate or MucK integration and plasmid backbone excision by colony PCR, and whole genome resequencing. Three lines of each resistant transformant strain were subsequently re-streaked onto LB Plates, followed by long term archiving in 20% glycerol/LB at −80 °C.

2.3. Pseudomonas putida transformation

P. putida KT2440 was used as the wild-type strain. Wild type liquid cultures were grown from plated single colonies in 25 mL of LB medium overnight at 30 °C, shaking at 200 rpm, to saturation. The following day, the cultures were harvested by centrifugation at 10,000 g, washed three times in 15 mL of 10% glycerol, and resuspended in a 50-fold smaller volume of 10% glycerol to generate electrocompetent cells for subsequent transformations.

Transformation was implemented by electroporation in the presence of 500–1000 ng plasmid DNA (0.1 cm cuvette, 1.6 kV, 25 μF, 200 Ω) as previously described (Elmore et al., 2017). Cells were then incubated in 950 μL SOC medium at 30 °C for 2 h, followed by plating on LB agar supplemented with 50 μg/mL kanamycin sulfate at 30 °C for 16 h. Resultant single colonies were re-streaked to single colonies on fresh LB agar +50 μg/mL kanamycin sulfate plates at 30 °C for 16 h to remove any enduring wild-type cells. Resultant colonies were streaked to single colony on YT-25% sucrose agar (10 g/L yeast extract, 20 g/L tryptone, 250 g/L sucrose, 18 g/L agar) for counter-selection against the plasmid backbone, incubating at 30 °C for ~30–48 h. Surviving colonies were then re-streaked onto fresh YT-25% sucrose plates and incubated at 30 °C for 16 h to remove any enduring sacB-containing cells. All final colonies were screened for complete mKate or MucK integration and plasmid backbone excision by colony PCR, and whole genome resequencing. Three lines of each resistant transformant strain were subsequently re-streaked onto LB Plates, followed by long term archiving in 20% glycerol/LB at −80 °C.

2.4. Genome re-sequencing

Genomic DNA was purified from saturated liquid cultures using Qiagen’s DNeasy Blood and Tissue Kit (Qiagen, Valencia, CA) according to the manufacturer’s instructions. Nextera XT libraries (Illumina, San Diego, CA) were produced according to the manufacturer’s protocol (15031942 v03). Final library validation was conducted on an Agilent Bioanalyzer (Agilent, Santa Clara, CA) using a DNA57000 chip and the concentration was determined using an Invitrogen Qubit (Waltham, MA) with the broad range double stranded DNA assay. Barcoded libraries were pooled and prepared for sequencing following the manufacturer’s recommended protocol (15039740v09, Standard Normalization). One paired end sequencing run (2 × 301) was completed on an Illumina MiSeq instrument (Illumina, San Diego, CA) using v3 chemistry.

2.5. Growth and fluorescence assays

Three lines of each transformant strain were inoculated from single colonies into 5 mL of LB and grown overnight at 30 °C, shaking at 200
rpm, for 16 h. Liquid cultures were then centrifuged and washed in the target assay growth medium (M9 + 30 mM glucose). Each well in a black walled, μClear flat-bottom, 96-well plate (Greiner Bio-One, Kremsmünster, Austria) was inoculated at 1%, into a final volume of 100 μL of the assay growth medium. Three replicates of each line, for each strain were measured simultaneously. Each plate was inoculated at 30 °C, with continuous shaking, in a Synergy Mx plate reader (BioTeck, Winooski, VT), measuring OD\text{600 nm} and \text{F_{888, 635 nm}} every 10 min for 48 h. 

Growth rates were calculated using the LINFEST function (fit to 10 measurements) in Microsoft Excel (2016) to determine the slope of the exponential growth phase, at mid-log. Fluorescence readings were normalized to the optical density and reported at mid-log of growth.

2.6. Live cell fluorescence microscopy

The P. putida KT2440 wild type, and P. putida KT2440 strains with mKate inserted at PP_1642, PP_2224, and PP_5322 were selected from single colonies and grown to saturation in LB overnight at 30 °C, shaking at 200 rpm. Saturated cultures were diluted 1:100 into fresh M9 minimal medium supplemented with 30 mM glucose and allowed to grow at 30 °C with continuous shaking until early log phase was reached (OD 0.1–0.2). Cells were imaged using a Zeiss Axio Imager microscope (Carl Zeiss, Gottingen, Germany) with a Canon EOS RebelT1i (Canon, Tokyo, Japan) attached. Images were captured at 240X magnification under bright field light conditions, and with a fluorescent filter. Samples were exposed for approximately 2.5 s.

2.7. Analysis of strain growth on cis,cis-muconate and MucK expression levels

Cis,cis-muconic acid was purchased from ACROS Organics (AC297760100). Stock solutions with a maximum concentration of about 800 mM were prepared by dissolving the powdered reagent in water containing 2 molar equivalents of NaOH. The pH was adjusted to 7.0 with NaOH, as needed, to maintain solubility and prevent isomerization. After filter sterilization, final concentrations were determined by absorbance at 260 nm using an extinction coefficient of 16,900 M⁻¹ cm⁻¹ (Sistrom and Stanier, 1954).

Overnight cultures of wild type P. putida KT2440 and strains carrying the mucK insertion were inoculated into 10 mL LB medium to give a starting culture density of 0.2 OD\text{600 nm} and were grown with shaking at 30 °C until the culture density reached 1.0 OD\text{600 nm}. Cells were harvested by centrifugation and were washed one time with ice cold PBS. The washed cell pellets were weighed and then vortexed into solution. A bicinchoninic acid (BCA) assay (Thermo Scientific, Rockford, IL) was performed to determine protein concentration. Following the assay, dithiothreitol (DTT) was added to the samples to the final concentration of 5 mM and incubated at 60 °C for 30 min with constant shaking at 850 rpm. The samples were alkylated by the addition of iodoacetamide (IAA) to a final concentration of 40 mM and incubated at 37 °C for 1 h with 850 rpm shaking in the dark. The samples were then diluted 10-fold for digestion preparation with 100 mM NH₄HCO₃ and 1 mM CaCl₂ (final concentration). USB trypsin was added to all protein samples at a 1:50 (w/w) trypsin-to-protein ratio and incubated for 3 h at 37 °C. Digested samples were desalted by using 1 mL Discovery C18 SPE columns (Supelco, Bellefonte, PA) using the following protocol: 3 mL of methanol was added for conditioning the column followed by 2 mL of 0.1% trifluoroacetic acid (TFA) in H₂O. The samples were then loaded onto each column followed by 4 mL of 95:5 water-acetonitrile, 0.1% TFA. Samples were eluted with 1 mL 80:20 acetonitrile:water, 0.1% TFA. The samples were concentrated down to ~100 μL using a Speed Vac, and a final BCA was performed to determine the peptide concentration. Samples were diluted to 0.20 μg/μL with nano-pure water for targeted proteomics analysis.

Targeted Proteomics was performed via a Liquid Chromatography (LC) - Selected Reaction Monitoring (SRM) approach. Three fully-tryptic peptides (GTAIGGAYNVGR and GTAAVGYVGR) were spiked into test samples. All the peptides were further compared on BLAST to ensure their uniqueness to target proteins in the organism. Crude (minimum purity) synthetic heavy isotope-labeled (e.g., 13C/15N on C-terminal lysine and arginine) peptides were purchased from New England Peptide (Gardner, MA). Upon receiving, the crude synthetic heavy peptides were mixed together and diluted with 0.1% formic acid in 15% acetonitrile in water to obtain a nominal concentration of 2 pmol/μL for each individual peptide. The heavy peptide mixture stock solution was aliquoted and stored at ~80 °C until further use.

Analysis of the targeted proteomics assay was conducted via LC-SRM. Each SRM precursor-fragment ion pair (i.e., transitions) was first analyzed with LC-SRM by spiking heavy peptides into test samples. Even though all three endogenous peptides can be detected in strains carrying the mucK insertion, two peptides (GTAIGGAYNVGR and LIAGFMADK) of MucK protein were initially selected based on their SRM suitability scores predicted by CONSeQuence (Eyers et al., 2011) and Prego (Searle et al., 2015) software tools. All the peptides were further compared on BLAST to ensure their uniqueness to target proteins in the organism. Crude (minimum purity) synthetic heavy isotope-labeled (e.g., 13C/15N on C-terminal lysine and arginine) peptides were purchased from New England Peptide (Gardner, MA). Upon receiving, the crude synthetic heavy peptides were mixed together and diluted with 0.1% formic acid in 15% acetonitrile in water to obtain a nominal concentration of 2 pmol/μL for each individual peptide. The heavy peptide mixture stock solution was aliquoted and stored at ~80 °C until further use.

To facilitate protein quantification, crude heavy peptide mixture stock solution was spiked in the 0.20 μg/μL peptide samples at a nominal concentration of 50 fmol/μL for each peptide. LC-SRM analysis utilized a nanoACQUITY UPLC® system (Waters Corporation, Milford, MA) coupled online to a TSQ Altis® triple quadrupole mass spectrometer (Thermo Fisher Scientific). The UPLC® system was equipped with an ACQUITY UPLC BEH 1.7 μm C18 column (100 μm i.d. × 10 cm) and the mobile phases were (A) 0.1% formic acid in water and (B) 0.1% formic acid in acetonitrile. 2 μL of sample (i.e., 0.4 μg of peptides) were loaded onto the column and separated using a 110-min gradient profile as follows: (min:flow-rate-μL/min/%B): 0:0.4:1, 6:0.6:1, 7:0.4:1, 9:0.4:6, 99:0.4:13, 70:0.4:22, 80:0.4:40, 85:0.4:95, 91:0.5:95, 92:0.5:95, 93:0.5:50, 94:0.5:95, 95:0.6:1, 98:0.4:1. The LC column was operated at
a temperature of 42 °C. The TSQ Altis™ triple quadrupole mass spectrometer was operated with ion spray voltages of 2100 ± 100 V and a capillary inlet temperature of 350 °C. Tube lens voltages were obtained from automatic tuning and calibration without further optimization. Both Q1 and Q3 were set at unit resolution of 0.7 FWHM and Q2 gas pressure was optimized at 1.5 mTorr. The transitions were scanned with a dwell time of 0.625 msec.

LC-SRM data were imported into Skyline and the peak boundaries were manually inspected to ensure correct peak assignment and peak boundaries. Peak detection and integration were determined based on two criteria: 1) the same LC retention time and 2) approximately the same relative peak intensity ratios across multiple transitions between the light peptides and heavy peptide standards. The total peak area ratios of endogenous light peptides and their corresponding heavy isotope-labeled internal standards were then exported from Skyline as Ratio-to-Standard. For each peptide, the total peak area ratios of individual samples were normalized to the average total peak area ratio of all the samples. For each sample, protein abundance was calculated as an average of the normalized total peak area ratios of all three peptides of a protein.

3. Results and discussion

3.1. Integration site selection and construction

The *P. putida* KT2440 chromosome was computationally screened for potential heterologous DNA integration loci that were not adjacent to essential genes, did not integrate within an operon, and were in an antiparallel orientation relative to the downstream gene [Fig. 1].

From the generated list of possibilities, we down-selected to seven sites for the practical and manageable generation of single reporter strains by homologous recombination [Fig. 2A]. These sites represented different regions of the chromosome and varied in total intergenic length (12–211 bp). The annotated functions of the genes upstream and downstream of each loci are provided in Supplemental Table 1. We designed and synthesized an *mKate* reporter construct driven by the *E. coli* Ptac promoter, which was flanked by terminators on both the 5′ and 3′ ends to insulate the construct from outside transcription readthrough, and to prevent readthrough from the *mKate* gene into the downstream gene [Fig. 2B]. The *mKate* fluorescent reporter construct was introduced into the seven different insertion locations on the *P. putida* KT2440 chromosome by homologous recombination, always in the same orientation (clockwise, as drawn in Fig. 2A).

Whole genome resequencing of the engineered strains showed that each *mKate* construct was integrated into the targeted genome region. Several minor differences were noted relative to the original integration design. For one locus, PP_5388, the *mKate* construct integrated into the gene directly downstream of the target site, PP_5734, which encodes a hypothetical protein of unknown function. For two loci, PP_1035 and PP_1642, there were short base pair deletions from the intergenic region, 40 and 28 bp respectively. Finally, a point mutation occurred in the 5′ terminator of the PP_1035 strain.

3.2. Effects of integration location on growth and expression

Growth and fluorescence of the seven mKate expression strains were measured in M9 minimal medium supplemented with 30 mM glucose. The different integration loci had significant effects on both growth rate and mKate expression. At sites PP_1035, PP_4305, PP_5042, PP_5322, and PP_5388, insertion of mKate did not have a negative effect on growth, and all strains grew similarly to the wild type [Fig. 3]. The specific mKate expression, as measured by Fluorescence/OD, varied up to 27-fold between integration sites, even for locations that did not affect growth. Microscopy of wild-type and engineered cells identified no morphological differences due to fluorescent protein expression, as well as expected levels of cell-to-cell variability in fluorescence (Supplemental Fig. 1).
Potential integration loci such as PP_1642 and PP_2224, which impose significant fitness costs on the host, have obvious tradeoffs. Even among the neutral sites, thus, a 27-fold variation in protein expression could have significant effects on the function of a heterologous pathway. Understanding and accommodating these differences will greatly improve the reliability of pathway design. There are several possibilities that may help explain the variances in host growth or protein expression. Copy number of the genes may be temporarily duplicated during DNA replication (Slager and Veening, 2016), especially close to the origin of replication. This variation may partially explain the expression levels at some sites such as PP_5322 having the highest mKate expression (close to the oriC), and PP_2224 being one of the lowest mKate expression sites (close to the terminus). Although the integration of PP_5388 interrupted the coding sequence of a hypothetical protein, this did not alter normal growth rates.

A previous study (Domröse et al., 2019) used Tn5 transposition to randomly integrate a 21 kb prodigiosin gene cluster across the P. putida KT2440 chromosome. In that example, it was found that the mutants with the highest expression were biased to one specific location, one of seven rrn operons. The authors propose that the optimized insertion at this locus is due to both the redundancy of the rrn operons (preventing interference of normal cell functioning) and the essentiality (high transcription levels) of the operon. However, high expression is not always desirable, and there are many situations where the ability to fine-tune expression through the choice of integration locus would be valuable. In E. coli (Bryant et al., 2014) significant differences were found in the expression of a gfp reporter under the control of a lac promoter integrated at fourteen rationally decided locations. The authors attribute the approximate 300-fold variation in expression across the sites to gene silencing at transcriptionally silent Extended Protein Occupancy Domains (tsEPODs), and transcriptional rate variations due to DNA supercoiling. This work further evidences the importance of the integration location as a major contributor to both maintaining normal cell functioning and generating the quantity of protein expression.

3.3. Effects of integration locus on function of a heterologous transporter

In order to further test the effect of the integration locus on function of a heterologous gene, we integrated a functional protein, a muate transporter, into three of the seven sites and measured the resulting variation in growth and protein expression. The integration sites selected for further characterization each displayed different phenotypes with mKate, such as slow growth and low fluorescence (PP 2224), slow growth and high fluorescence (PP 1642), or wild type growth and medium fluorescence (PP 5042).

cis,cis-mucenate is a potential value-added bioproduct that can be chemically converted into adipic acid for use in nylon synthesis (Johnson et al., 2016). Biologically, cis,cis-mucenate could be further metabolized to products such as β-keto adipic acid, but these pathways will require further optimization. P. putida KT2440 can metabolize cis,cis-mucenate for growth, as it is a key intermediate in the β-ketoacidipate pathway for microbial dissimilation of aromatic compounds (Harwood and Parales, 1996; Ormston and Stanier, 1966). However, KT2440 is not capable of growing on exogenously supplied cis,cis-mucenate, as it lacks import mechanisms for this metabolite. Assimilation of exogenous cis,cis-mucenate would simplify downstream pathway optimization, but will require expression and tuning of a muate transporter.

Here we incorporated a codon-optimized copy of mucK, the gene encoding for the muate importer from Acinetobacter baylyi ADP1 (Williams and Shaw, 1997), into three different insertion site loci. The same plasmids used for mKate insertion were used for mucK, including the homology arms and the insulating terminators. Expression of mucK was driven by the strong, constitutive Pc promoter from the class III integron of Delftia acidovorans C17 (Xu et al., 2013) and the gene 10 ribosome binding site from bacteriophage T7 (Olin et al., 1988). The three strains were then challenged to grow with muate for functional assessment of the insertion sites.

Expression of MucK conferred growth with muate as the sole carbon source, and different growth rates were observed based on the locus of insertion (Fig. 4). We used targeted proteomics (Supplemental Fig. 2) to compare MucK expression levels among strains with mucK inserted at different loci (Fig. 4). Unexpectedly, the insertion site that was correlated with the lowest growth and mKate expression (PP 2224), had the highest growth rate with muate and the lowest MucK expression. The other two sites had the inverse effect in MucK with respect to mKate. The PP 1642 site had low protein expression and higher growth rates with muate, while the PP 5042 site had the highest protein expression and the lowest growth rate. Although high levels of heterologous membrane protein expression can be toxic, growth with glucose was not inhibited by MucK expression (Supplemental Fig. 3). It has been shown that the accumulation of high levels of intracellular muate can be toxic to bacterial cells (Gaines et al., 1996). It
may be that rapid muconate import in the presence of a highly expressed transporter could inhibit cell growth. Indeed, the strain with the slowest growth on muconate displayed the highest levels (approximately 2-fold higher) of MuKc expression.

These experiments demonstrate that variation due to integration locus is not unique to fluorescent protein expression, but also affects functional proteins such as a transporter. In contrast to the previous integration locus analysis (Domrós et al., 2019) where high expression was desirable and integration loci were chosen accordingly, utility of MuKc was highest when expression was moderate. Notably, growth on glucose with the insertion of mKate at PP_2224 and PP_1642 was substantially decreased, where growth on glucose in all three muKc strains was sustained at wild type levels. The expression of mKate at PP_5322 was equivalent to the expression at PP_1642 however the PP_5322 strain was able to maintain wild-type growth rates. The expression of heterologous proteins needs to be tuned for particular purposes, and the choice of integration locus will substantially affect both the final protein expression and, in some cases, cell growth.

4. Conclusion

Chromosomal integration of heterologous genes and pathways is the ideal method for metabolic engineering of industrial strains. In this study, we have selected integration sites for heterologous DNA insertion that are non-proximal to essential genes, non-operon interfering, and in antiparallel orientation to the downstream genes. We measured and characterized a small subset of possible insertion loci for the integration of heterologous DNA, showing that different integration loci can have significant effects on protein expression and the resulting organismal phenotype. This work is an important first step in optimizing the placement of target genes and pathways. Future work is needed to characterize many more sites throughout the genome, test different construct orientations, and examine operons or multigene constructs to determine the biological mechanisms responsible for expression variation. Reliable, predictable expression tuning by choice of integration site will provide additional control for metabolic engineering in this important chassis organism.

CRediT authorship contribution statement

Julie E. Chaves: Investigation, Formal analysis, Validation, Methodology, Visualization, Writing - original draft. Rosemarie Wilton: Investigation, Formal analysis, Validation, Methodology, Visualization, Writing - original draft. Yuqian Gao: Methodology, Investigation, Validation, Formal analysis, Visualization, Writing - original draft. Nathalie Munoz Munoz: Methodology, Investigation, Writing - review & editing. Meagan C. Burnet: Investigation, Writing - original draft. Zachary Schmitz: Investigation, Writing - review & editing. John Rowan: Investigation, Methodology, Writing - review & editing. Leah H. Burdick: Investigation, Writing - review & editing. Joshua Elmore: Conceptualization, Investigation. Adam Guss: Conceptualization, Methodology, Supervision, Project administration, Funding acquisition, Writing - review & editing. Dan Close: Conceptualization, Methodology, Supervision, Project administration. Jon K. Magnuson: Funding acquisition, Writing - review & editing. Kristin E. Burnum-Johnson: Methodology, Supervision, Writing - review & editing. Joshua K. Michener: Conceptualization, Methodology, Investigation, Supervision, Project administration, Funding acquisition, Visualization, Writing - original draft.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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