Estimating the Number of Essential Genes in Random Transposon Mutagenesis Libraries

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Abstract

Biologists use random transposon mutagenesis to construct knock-out libraries for bacteria. Random mutagenesis offers cost and efficiency benefits over the standard site directed mutagenesis,[8] but one can no longer ensure that all the nonessential genes will appear in the library. In random libraries for haploid organisms, there is always a class of genes for which knockout clones have not been made, and the members of this class are either essential or nonessential. One requires statistical methods to estimate the number of essential genes. Two groups of researchers, Blades and Broman[1] and Jacobs et al.[8] independently and simultaneously developed methods to do this. Blades and Broman used a Gibbs sampler and Jacobs et al. used a parametric bootstrap. We compare the performance of these two methods and find that they both depend on having an accurate probabilistic model for transposon insertion or on having a library with a large number of clones. At this point, we do not have good enough probabilistic models so we must build libraries that have at least five clones per open reading frame to accurately estimate the number of essential genes.

1 Introduction

Scientists are creating knockout clonal libraries for many microorganisms.[5][6][7][9] Usually, researchers follow a site directed mutagenesis approach in which each clone had a predetermined open

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reading frame (ORF) disrupted. Jacobs et al. [8] generated a library for *Pseudomonas aeruginosa* cheaply and efficiently by random transposon mutagenesis. Rather than specifying which gene would be knocked out before mutation, they randomly mutated the genome and determined the location of the gene knockout afterwards.

Each clone in a random mutagenesis library has a single transposon insertion. A transposon is a DNA sequence that can jump within chromosomes and between them. Biologists have built transposons with stop codons that upon insertion in an ORF terminate protein translation. When the insertion is random, one requires statistical methods to help construct the clonal library. Foremost, one would like to know how many clones to make. The number of clones made for the *P. aeruginosa* library was based on the estimated number of essential genes.

Biologists and clinicians are interested in essential genes because they make good candidates for drug targets. [4] Essential genes never appear knocked out in random mutagenesis libraries because bacteria are haploid organisms. As a tautology, once an essential gene is knocked out, the bacterium cannot live. Besides essential genes, there are also nonessential genes that have yet to appear as knockouts in the library because of chance. Jacobs et al. believed that a large number of clones should be created to get a good estimate of the number of the essential genes. They made about about five clones per ORFs and used a parametric bootstrap to estimate the number of essential genes. In a concurrent experiment, Lamichhane et al. [10] used a Gibbs Sampler developed by Blades and Broman [1] and claimed that they only required 0.3 clones per ORF to obtain a good estimate of number of essential genes in *Mycobacterium tuberculosis*.

The *P. aeruginosa* clinical isolate, PAO1, used to create the clonal library has 5,570 ORFs [13] and the *M. tuberculosis* isolate has 4,250 ORFs. Prokaryotic genomes do not have introns so stretches of DNA longer than 300 bp beginning with a methionine codon and lacking a stop codon strongly indicate the existence of an ORF. We assume that all the ORFs and their locations are known.

In this paper, we look at the fit of the probability model underpinning both estimation procedures and compare the accuracy and precision of the parametric bootstrap to the Gibbs sampler. The *M.*
tuberculosis library has 1,403 unique insertions, which corresponds to 1,839 insertions in the PAO1 library for the same coverage of $\frac{1839}{5570} = 0.3X$. The PAO1 library has 30,100 unique insertion locations; a coverage of $\frac{30100}{5570} = 5.4X$ unique insertions per ORF. Both the parametric bootstrap and the Gibbs sampler are based on the same multinomial insertion model. Considering only unique positions, the multinomial model fits the PAO1 data at the 0.3X coverage level. However, if we use all 30,100 insertions to test model fit, we reject the hypothesis that the bin probabilities used in the model are correct. The multinomial model requires the ability to determine the probability that an insertion lands in an individual ORF based solely on the annotation (i.e. ORF length). We show that one can predict the insertion probabilities but not accurately enough to estimate the number of essential genes at coverage levels lower than 5.0X.

We analyze how effectively both methods estimate the number of essential genes at lower than 5.4X coverage by using the empirical distribution of the unique positions in the PAO1 library, i.e. drawing samples without replacement from the list of insertion locations, in spite of the evidence against the multinomial insertion model. We find a bias-variance trade-off in that the parametric bootstrap is more accurate and the Gibbs sampler is more precise. At the 5.4X coverage level, the parametric bootstrap’s estimate of the number of essential genes agrees with the estimate from Gibbs sampler. Once coverage is high enough, we are confident that one can estimate the number of essential genes. The Gibbs sampler exhibits bizarre behavior at coverage levels below 0.3X which we attribute to mis-specification of the insertion probabilities. Through simulation, we compute the necessary coverage for two endpoints: Checking model accuracy and estimating the number of essential genes.

2 Empirical Verification of the Multinomial Model

The data we use for this paper consist of the 42,240 mutants created for the PAO1 library. The locations of 36,154 of them have been mapped and 30,100 land in unique places. Of the unique locations, 27,264 are inside ORFs and the other 2,836 are between them. The inserts hit 4,895 of the 5,570 ORFs internally so there are 675 candidate essential genes. We only consider unique loca-
tions, because there were many more duplicate sites than expected by chance. We feel this is due to local contamination effects created by the high-throughput technology used to create the PAO1 library.

The exact chemical mechanism for transposon insertion is not important for this paper except that biologists putatively maintain that the elements insert randomly and uniformly at specific targets in the genome.\[12\] The elements used in the PAO1 library insert at any base pair and the element used in the *M. tuberculosis* library inserts at TA dinucleotides. Since the PAO1 and *M. tuberculosis* genomes are sequenced and annotated,\[13\] we can locate the exact position of insertion and assume that the location of the transposable element can unambiguously be determined. Biologists engineer the transposons to have stop codons terminating the translation of the protein. However, transposons that landed in the distal portion of the ORF may have only stunted the protein allowing it to function. Hence, there are varying definitions as to what constitutes a knockout. Generally, we use the most liberal definition that an insert landing anywhere within the ORF knocks out the protein function. Lamichhane et al. consider insertions that land in the last 100 bp or 20% of the ORF not to knockout the gene (the “5’80%-3’100-bp” rule). We also look at definitions in which knockouts occur only when the insert landed in the middle 80% or 60% of the ORF.

A multinomial distribution naturally approximates the biology of the transposable elements. We let $k$ be the number of open reading frames in an organism’s genome, $n$ be the number of mutants assayed, and $m$ be the number of essential ORFs. The $m$ is unobservable and is the focus of the estimation procedures. Based on the biology of the transposons used in the PAO1 library, we believe that the probability of hitting a nonessential gene is proportional to its length in base pairs. One caveat that prevents the joint density of number of times the ORFs are hit from being a true multinomial is that the ORFs can overlap and an insertion in the overlap knocks out both genes. Like Blades and Broman,\[1\] we let $\mathcal{G} = (g_1, \ldots, g_k)$ be the vector of zeros and ones indicating whether an ORF is nonessential with 0 indicating essentiality. We note that $\sum g_i = k - m$ and define $X = (x_1, \ldots, x_k)$ to be the number of insertions per ORF in regions that are not shared and $Y = (y_1, \ldots, y_k)$ to be the number of insertions in regions shared between gene $i$ and $i + 1$. $y_k$ is
the number of insertions into the region shared by gene $k$ and the first gene, because prokaryotic chromosomes are circular. We let $(p_1, \ldots, p_k)$ be the probabilities that a transposable element inserts into an ORF given that it is nonessential and let $(q_1, \ldots, q_k)$ be the probability that it inserts into the region shared by gene $i$ and $i+1$. Often it is convenient to model the intergenic region as the $k$th ORF and then the $q$’s have to be changed appropriately. We note that $\sum x_i + y_i = n$ and $\sum p_i + q_i = 1$. Given $\mathcal{G}$, the distribution of $(X, Y)$ is

$$P(X, Y|\mathcal{G}) = \prod_{1 \leq i \leq k} \frac{(p_i g_i)^{x_i} (q_i g_i g_{i+1})^{y_i}}{(\sum_{1 \leq j \leq k} p_j g_j + q_j g_j g_{j+1})^n}. \quad (1)$$

We want to verify that the multinomial insertion model with a class of unhitable essential genes fits the unique locations. We should check that the unique locations occur uniformly and randomly in the genome and that the probability of hitting an ORF can be computed by dividing the length of the ORF by the number of bp in the genome. First, 90.6% of the unique locations are in coding regions which is consistent with the 89.4% of the genome that is actually in coding regions.[13] We plot a histogram of the location of the insertion sites within genes hit (Figure 1). The transposons appear to be inserting at each bp with equal probability even though, according to the Kolomogorov-Smirnov test, the empirical distribution is not consistent with the Uniform distribution ($D = 0.0089$, $p = 0.026$). Unfortunately, we cannot use the intergenic region for testing the distribution of insertion locations since it is difficult to annotate. The region is likely populated with regulatory elements that are not well defined.

We focus on the ORFs and define a modified goodness-of-fit statistic to accommodate the unobserved essential genes. We define $\chi^2_+$ to be the goodness-of-fit statistic on only those ORFs that are hit more than once. From large sample theory, we know that if the number of insertions, $n_i$, is large then

$$\sum_{1 \leq i \leq k} \frac{(x_i - E_i)^2}{E_i} + \sum_{1 \leq i \leq k} \frac{(y_i - E_i)^2}{E_i} \sim \chi^2_{k'-1}. $$

$k'$ is number of ORFs and ORF overlaps that have one or more targets. If one conditions on only a subset of the bins being occupied in a multinomial, then the distribution of counts for these
bins are Multinomial \((n, p^*_1, \ldots, p^*_k)\) with \(p^*_i = p_i / (p_1 + \cdots + p_k)\).

For example, if \(k = 4\) and we condition on only bins 1 and 2 being occupied, then \(p^*_1 = \frac{p_1}{p_1 + p_2}\) and \(p^*_2 = \frac{p_2}{p_1 + p_2}\). If we knew which genes were nonessential then we could test how well the multinomial fits by looking at a \(\chi^2\) on the conditioned multinomial. Define

\[
\chi^{2*} = \sum \left( \frac{x^{*}_i - E^{*}_i}{E^{*}_i} \right)^2 + \sum \left( \frac{y^{*}_i - E^{*}_i}{E^{*}_i} \right)^2
\]

where \(x^{*}_i\) and \(y^{*}_i\) are the counts of insertions in ORFs or overlap regions that have been hit at least once. \(E^{*}_i = n p^*_i\) in which \(p^*_i\) is the recomputed probability of hitting a region given that those regions with at least one insertion were the only ones that could have been hit. We can show that the asymptotic distribution of \(\chi^{2*}\) is \(\chi_{k'-m-1}\) by use of one of the Slutsky Theorems. The distribution is dependent upon the number of essential genes. Using the limiting distribution, we construct a very conservative \(\alpha\)-level test for rejecting the null hypothesis that the joint distribution of the number of times an ORF is hit is Multinomial \((n, p^*_1, \ldots, p^*_k)\). We reject the multinomial model if \(\chi^{2*} > \chi^2_{k'-1,1-\alpha}\), which occurs with probability
Table 1: Goodness-of-Fit for Different Knockout Definitions

| Model                | $\chi^2$ Fit | DOF |
|----------------------|--------------|-----|
| Entire ORF           | 9663.9       | 6324|
| Lamichhane           | 7621.8       | 5570|
| 5’ 10%-3’ 10%       | 8487.5       | 5571|
| 5’ 20%-3’ 20%       | 6733.1       | 5570|

Different models, the modified $\chi^2$ fit, and number of ORFs added to the number of overlaps. Entire ORF counts insertions that land anywhere within an ORF. Lamichhane counts only those that do not land in the last 100 bp or the distal 20%, 5’ 10%-3’ 10% counts those that land in the middle 80%, and 5’ 20%-3’ 20% counts those that land in the middle 60%.

If we compute the $\chi^2$ goodness-of-fit statistic for ORFs hit anywhere internally at least once, we get a value of 9663.9 on 6324 degrees of freedom. Under the null hypothesis that the multinomial insertion model is correct, the 0.95 quantile for the $\chi^2_{6324}$ distribution is 6510.1, and we safely reject the hypothesis that the insertions are following a multinomial distribution whose probabilities are computed from the length of the ORF divided by the number of base pairs in the genome. Model-fitting with 30,100 observations is often dangerous, but we use a conservative test and found a value well within the critical region. We compute the $\chi^2$ goodness-of-fit statistic for other ORF definitions (Table 1) and for each, we reject that the insertion probabilities based on gene length. For the rest of the paper, we use the most inclusive definition of a knockout and count insertions anywhere within the ORF.

To observe the discrepancy between the predicted probabilities and the experimental distribution of hits, we plot the observed number of insertions on gene length in bp for genes that have been hit once next to a simulated set of insertions from the correct multinomial (Figure 2). We see that the coefficient of determination, $R^2$, for the actual observations is 0.45 and for the simulated insertions it is 0.72 meaning that the transposons inserted in some ORFs more often than we expect and into others less. There is no evidence that this phenomenon corresponds with ORF length because the slopes of the linear regressions for both graphs are tantalizingly close: $4.42 \times 10^{-3}$ for the observed data and $4.75 \times 10^{-3}$ for the
simulated data.

For *M. tuberculosis* data set, the number of TA dinucleotides in an ORF divided by the number of TAs in the genome might accurately predict the probability that the transposon inserts into a gene, but our test does not have enough power to detect model failure at the 0.3X coverage level. Using the list of *M. tuberculosis* insertions, we find a $\chi^2$ fit of 854.8 which is not in the 95% rejection region for 4,279 degrees of freedom. This lack of power is due to the low coverage level, not transposon or organismal variation. If we take a sample of 1,839 PAO1 insertions, we find a fit of 1288.1 on 6324 degrees of freedom.

### 3 Comparison of Estimators

Although the probabilities based on the length of the gene are approximate, we can still compare the two methods for estimating the number of essential genes. Here’s a brief description of the parametric bootstrap.\[2\] One can fit the function

$$f(n) = b_0 - b_1 \exp(-b_2 n)$$  \hspace{1cm} (3)

to the cumulative plot of the number of ORFs hit. The parameters $b_0$, $b_1$, and $b_2$ are chosen to minimize the residual standard error between the function and the data. Fitting 30,100 points is computationally intensive, and we want to use exclusively R for all our programming, so we choose 100 equally spaced points including the last one to fit. Hence, for all the insertions, look at the number of different ORFs hit at 301, 602, \ldots, 30100 insertions. One interprets the parameter $b_0$ as the number of nonessential genes.

However, fitting this model is not a standard nonlinear regression, so we compute the bias and variance of the estimated parameters in a different manner. We proceed by assuming that parameters fitted to the actual cumulative plot have the same bias and variance as when they are fitted to the multinomial model without any essential genes. In other words, $k-b_0^* \sim m-\hat{m}$ where $\hat{m}$ is the estimate of the number of nonessential genes. We simulate $(X, Y)$ according to Equation (1) with all $g_i = 1$ by drawing a sample without replacement of size $n$ from the set of targets and placing them in the appropriate ORF. If $b_{0j}^*$ is the $b_0$ fitted to the $j^{th}$ simulated experiment of $l$ experiments,
Bias $\hat{m} \approx \bar{b}_0^* - k$ where $k$ is the number of ORFs in the genome and 
$\text{Var}(\hat{m}) \approx \frac{1}{l-1} \sum_{1 \leq j \leq l} (b_{0j}^* - \bar{b}_0^*)^2$. To compute the $100(1 - \alpha)\%$ confidence interval, we use

$$P \left( \hat{m} + k - b_{0(1-\alpha/2)}^* \leq m \leq \hat{m} + k - b_{0(\alpha/2)}^* \right) \approx 1 - \alpha$$

where $b_{0(x)}^*$ is the $x^{th}$ quantile.

We estimate that there are 382 essential genes with a standard deviation of 15.8. The 95% confidence interval for the number of essential genes is $[340, 412]$.

We find a posterior mean for the number of essential genes of 408 with a 95% credible interval of $[384, 432]$ using the Gibbs sampler. At a coverage of 5.4X insertions per ORF, both estimates of the number of essential genes were basically the same and agree with a basic bioinformatic assessment of gene essentiality. [8]

To see how well our the parametric bootstrap method compared with the Gibbs sampler, we conduct a study using the empirical distribution based on the locations of the 30,100 unique insertion locations in the PAO1 clonal library. We assume that genes were nonessential when an insertion landed anywhere in the entire length of the ORF. We draw 100 samples of a certain coverage without replacement from the 30,100 unique insertion locations and for each sample, we compute the point and interval estimates using both estimation procedures. The means of the 100 samples are displayed in Figure 3.

The Gibbs sampler has tighter credible intervals overall but has more bias than the parametric bootstrap. The credible region at the 20,000 insertions (3.6X coverage) did not cover the estimate of the number of essential genes at 30,100 insertions whereas the 95% confidence interval always did. The Gibbs sampler estimate also exhibits some rather strange behavior at extremely low coverage levels. The most accurate point estimate occurred at 500 insertions or 0.09X coverage. We do not see the immediate drop in the estimate as in Figure 3 when we explore the behavior of the Gibbs sampler by simulating insertions from a simplified, correct mode. Our results are not shown but they are consistent with the simulation studies in Blades and Broman’s technical report. [1] The Gibbs sampler on simulated data behaves like the parametric bootstrap does on the experimental data. Therefore, we have little faith in the accuracy of their method at coverage levels at around 0.1X.
We use the empirical distribution based on the 30,100 unique insertions to find the number of clones needed for a 0.90 probability that the multinomial model would be rejected using the modified $\chi^2$ statistics and for the 95% credible interval found by the Gibbs sampler would cover 408; their best estimate of the number of essential genes. We draw a number of samples without replacement of different coverage levels and determine when the event happened about 90% of the time. We determine that one needs a 2.9X coverage for a 0.9 probability of the $\chi^2$ test rejecting the null hypothesis that the probabilities based gene length are correct. Secondly, one needs a coverage of 5.0X to have a 0.9 probability of covering 408 essential genes using the Gibbs sampler’s 95% credible interval.

### 4 Discussion

Numerous statistical questions arise while creating random transposon mutagenesis clonal libraries. Two groups of researchers developed different methods to estimate the number of essential genes in a prokaryotic organism given the genomic reference sequence and the number and types of clones in the library.

First, both methods are based on a multinomial insertion model that has a substantial number of parameters that must be computed a priori. For each ORF, we need to be able to calculate the probability that a transposon will land in the ORF given that it is nonessential based on the reference sequence. For the transposons used in the PAO1 library, one assumes that each inserts at a nonessential bp of the genome with equal probability. Likewise for the transposon used to build the *M. tuberculosis* library, one assumes that it inserts at each dinucleotide TA with equal probability. It definitely inserts at only TA dinucleotides, and Lamichhane *et al.* present evidence that it inserts at each nonessential TA with equal probability. We have duplicated their analysis for model fit. The PAO1 library transposons have little preference for where they insert in a nonessential genes (Figure 1). 9.4% of the unique insertions land in the 10.6% of the PAO1 genome in intergenic regions whereas 17% of the insertions land in the 13% of the *M. tuberculosis* genome in intergenic regions. The PAO1 library transposons hit duplicate sites more often than one would expect by chance. The majority of identical hits are from experiments run on the same day and many are
from the same plate. This strongly suggests that siblings (mutant strains that divide prior to being plated) and cross-contamination (which may occur at multiple stages from picking colonies through PCR) create the artifact in the data, rather than genuine patterns of exact duplication. Lamichhane’s methodology did not create as many artifacts. Our extreme high-throughput method would be predicted to create a higher proportion of artifacts, most of which would look like exact duplicates.

Extending the model-fit analysis, we compare the observed number of hits with the expected number hits in ORFs that were hit at least once. We reject the model that the probabilities of being hit are determined by gene length. The lengths are predictive of the correct distribution (See Figure 2) but not as accurate as the empirical probabilities computed from a simulation of a multinomial. We see this by comparing the $R^2 = 0.44$ for the actual data with the $R^2 = 0.77$ of the simulated data.

We feel that each gene confers a specific fitness to the cell which causes the difference in coefficients of determination (the $R^2$ values). Some are essential, so cells lacking these genes die. Some are somewhat essential, so cells missing these genes can live but not very well. Finally, some are completely nonessential, so it does not matter whether cells have these genes. The idea that cells lacking different functional genes have different fitness is not novel. How one would estimate these fitnesses is still being studied, especially how to do it from reference sequence alone. Our guess is that it would involve a competitive approach such as the one taken by Gerdes et al. [5].

Second, we compare how well the parametric bootstrap and the Gibbs sampler do at estimating the number of essential genes at lower coverage levels. The parametric bootstrap has less bias but a greater variance (Figure 3). We did a careful exploration of model fit to understand why the Gibbs sampler appears to be hyper-efficient. According to Figure 3, one could analyze 500 insertions, a coverage of 0.09X insertions per ORF and find an accurate estimate of the number of essential genes. We do not see the hyper-efficiency in our simulated data sets or the simulated data sets in Blades and Broman’s technical report [11] leading us to believe that mis-specified insertion probabilities are causing the behavior.

Third, we estimate coverage levels required to achieve various
goals. The most critical determination is the coverage one needs to reject the multinomial insertion model. For PAO1, three unique insertions per ORF are needed. If we are interested in accurate estimation of number of essential genes at sub-saturation coverage levels of around 0.3X, we believe that it is mandatory that one has an accurate probability model. However, this creates a paradox. We cannot know if the probability model is correct unless we achieve at least a modest level of coverage. Lamichhane et al. should have mapped 12,750 insertions to check model sufficiency rather than 1,425 and this would have undermined their goal of looking at a low sub-saturation coverage level. It is possible that a more powerful statistical test will help us identify model inaccuracy. If the transposon used to build the M. tuberculosis library were shown to insert with the correct probabilities in a higher coverage level study, it would become the transposon of choice for low coverage studies.

However, what if the best model that we can find is only partially correct? Right now, we do not know what covariates to use to adjust the probabilities. We could try non-parametric techniques, but these still would probably require high coverage levels. We are beginning to look at probability models that do not assume we know the probability of transposon insertion for an ORF. Alternatively, we could put our effort into only high coverage level studies because with a large number of inserts mapped. We have evidence that both methods find a sensible estimate for the number of essential genes. At the coverage level of 5.4X, we find 382 essential genes using the parametric bootstrap which is similar to the estimate of 408 essential genes using the Gibbs sampler. Both estimates are similar to the count of 350 P. aeruginosa’s genes that are homologous to essential genes in E. coli, as well as matching proportions of essential genes from other studies.

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References

[1] N. J. Blades and K. W. Broman. Estimating the number of essential genes in a genome by random transposon mutagenesis. Technical report, Johns Hopkins University, Department of Biostatistics, 2002.

[2] A. C. Davison and D. V. Hinkley. Bootstrap Methods and their Applications. Cambridge University Press, Cambridge, 1997.

[3] T. S. Ferguson. A Course in Large Sample Theory. Chapman & Hall, London, 1996.

[4] C. Fraser. A genomics-based approach to biodefence preparedness. Nature Rev. Genet., 5:23–33, 2004.

[5] S. Y. Gerdes, M. D. Scholle, J. W. Campbell, G. Balážsi, E. Ravasz, M. D. Daugherty, A. L. Somera, N. C. Krypides, I. Anderson, M. S. Gelfand, and et al. Experimental determination and system level analysis of essential genes in Escherichia coli MG1655. Jour. Bact., 185:5673–5684, 2003.

[6] G. Giaever, A. M. Chu, L. Ni, C. Connelly, L. Riles, S. Veronneau, S. Dow, A. Lucau-Danila, K. Anderson, B. André, and et al. Functional profiling of Saccharomyces cerevisiae genome. Nature, 418:387–391, 2002.

[7] C. A. Hutchison III, S. N. Peterson, S. R. Gill, R. T. Cline, O. White, C. M. Fraser, H. O. Smith, and J. C. Venter. Global transposon mutagenesis and a minimal mycoplasma genome. Science, 286:2165–2169, 1999.

[8] M. A. Jacobs, A. Alwood, I. Thaipisuttikul, D. Spencer, E. Haugen, S. Ernest, O. Will, R. Kaul, C. Raymond, R. Levy, and et al. Comprehensive transposon mutant library of Pseudomonas aeruginosa. Proc. Nat. Acad. Sci., 100:14339–14344, 2003.

[9] K. Kobayashi, S. D. Ehrlich, A. Albertini, G. Amati, K. K. Andersen, M. Arnaud, K. Asai, S. Ashikaga, S. Aymerich, P. Bessieres, and et al. Essential Bacillus subtilis genes. Proc. Nat. Acad. Sci., 100:4678–4683, 2003.
[10] G. Lamichhane, M. Zignol, N. J. Blades, D. E. Geiman, A. Dougherty, J. Grosset, K. W. Broman, and W. R. Bishai. A postgenomic method for predicting essential genes at subsaturation levels of mutagenesis: Application to Mycobacterium tuberculosis. Proc. Nat. Acad. Sci., 100:7213–7218, 2003.

[11] D. J. Lampe, M. E. A. Churchill, and H. M. Robertson. A purified mariner transposase is sufficient to mediate transposition in vitro. EMBO J, 15:5470–5479, 1996.

[12] W. J. Miller and P. Capy. Mobile Genetic Elements. Humana Press, Totowa, NJ, 2004.

[13] C. K. Stover, X. Q. Pham, A. L. Erwin, S. D. Mizoguchi, P. Warrener, M. J. Hickey, F. S. L. Brinkman, W. O. Hufnagle, D. J. Kowalik, M. Lagrou, and et al. Complete genome sequence of Pseudomonas aeruginosa PAO1, an opportunistic pathogen. Nature, 406:959–964, 2000.

[14] J. W. Thatcher, J. M. Shaw, and W. J. Dickinson. Marginal fitness contributions of nonessential genes in yeast. Proc. Natl. Acad. Sci., 95:253–257, 1997.

[15] H. Yu, D. Greenbaum, H. X. Lu, X. Zhu, and M. Gerstein. Genomic analysis of essentiality with protein networks. Trends Genet., 20:227–231, 2004.
Figure 2: Comparison of the observed and theoretical probabilities of insertion. Plot (a) is the observed number of insertions on ORF length. (b) is a simulated number of insertions on ORF length. Both plots only include ORFs that were hit at least once.
Figure 3: Mean point estimates and credible intervals for 100 simulated experiments based on the empirical distribution using the PAO1 insertion list. We draw samples without replacement of various sizes. The dashed, horizontal line is at 382, our estimate of the number of essential genes. The vertical line is the sampling level of the Lamichhane et al. paper. The triangles are the points estimates from the parametric bootstrap method and the circles are point estimates from the Gibbs sampler. The dotted lines form the bounds of the 95% confidence interval and the solid lines the bounds of the 95% credible interval.