# **Systems Biology & Neurobiology**

Homework Report

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### **Abstract**

Deep mutational scanning (DMS) makes use of large-scale mutagenesis to reveal intrinsic protein properties, functions and the consequences of genetic variation. Recently, the CRISPR/Cas9-mediated genomic error-prone editing (CREPE) technology was developed as a high-throughput method for mutating essential genes of *Escherichia coli* [1]. Its authors applied the technology to target rpoB, the gene encoding the  $\beta$  subunit of bacterial RNA polymerase, and used deep sequencing to study resistance against the antibiotic rifampicin. In particular, the authors studied epistastis effects by comparing fitness of double mutants in rpoB with those from the respective single mutations in the presence of rifampicin. In this report, we replicated the aforementioned epistasis study using a simplified dataset provided by A. Choudhury.

### State of the Art

Studying epistasis—be it in human or within bacteria—raises many challenges, as it can rarely be done using observational studies. However, understanding how combinations of mutations affect protein functions and behaviour within cells could give us insights into a huge number of biological processes, from antibiotic resistance to genetic diseases. In this section, we establish a non-exhaustive review of available methods and technologies used for studying epistasis in bacteria.

Deep mutational scanning (DMS) can provide significant insights into the function of essential genes in bacteria. This method couples genotype to phenotype to assess the activities of as many as 1 million mutant versions of a protein in a single experiment [2]. DMS is capable of scoring comprehensive libraries of genotypes for fitness in given environments in a massively parallel fashion. Essential bacterial genes are often targets of interest as they are key to their evolution, and can lead to phenotypes such as antibiotic resistance when mutated.

When the phenotype of interest is the cell's fitness in a specific environment, the presence of genetic interactions between mutations, i.e. epistasis, can constrain the course of evolution. Given the potential impact of epistasis in a variety of biological processes, recent studies have focused on measuring genome-wide levels of epistasis using the multiplex automated genome engineering (MAGE) technology [3]. MAGE was created for large-scale programming and evolution of cells: it simultaneously targets many locations on a chromosome for modification in a single cell or across a population of cells, thus producing combinatorial genomic diversity. It is based on lambda red-mediated recombination of single-stranded oligos to introduce mutations at specific genomic loci [4].

However, the MAGE technology has some limitations, as it was optimized only for a few cell strains and can lead to the accumulation of numerous off-target modifications. A recent study improved on MAGE by using a dominant-negative mutant protein of the methyl-directed mismatch repair (MMR) system, allowing efficient modification of multiple loci, without any observable off-target mutagenesis and prior modification of the host

genome. This improved technology, termed pORTMAGE, was used to achieve a transient suppression of DNA repair in *Escherichia coli*. In addition, pORTMAGE allows comparison of epistatic effects across a wide range of bacterial species [5].

Antibiotic resistance is one of the main topic of study when it comes to understanding the effects of epistasis in bacteria, as the key factors contributing to resistance are yet to be understood. A team of researchers studied how epistatic effects in *Escherichia coli* could be used to perturb the evolution of bacteria towards antibiotic resistance [6]. In this context, robotic lab-evolution platforms can be used to keep population size and selection pressure constant for hundreds of bacteria populations evolving in parallel. Using this method, specific cellular functions that drastically curtail the evolvability of resistance where identified. Using whole-genome sequencing, the team showed that strong negative epistasis was generally underlying these functions.

The CRISPR gene editing technology, already widely used in genome engineering, also allows for investigating how gene expression governs the adaptive pathways available to bacteria during the evolution of resistance. In that sense, the controlled hindrance of adaptation of organisms (CHAOS) approach was recently developed to induce negative epistasis in *Escherichia coli* to deter adaptation [7]. Using a library of deactivated CRISPR-Cas9 devices, the team perturbed the bacterial gene expression and observed that epistatic effects caused large losses of cell fitness in environment containing ciprofloxacin, a clinically-relevant antibiotic. Another team introduced the homologous sequence integration (HoSeI) method to study bacterial genome-wide epistatic phenomena. HoSeI is a genetic marker-less genome editing approach that introduces base substitutions in the target sequence by screening dead or alive cells. It was demonstrated in a strain of *Escherichia coli* to study the effects of epistasis on regulators of bacteria adaptive growth [8].

To generate the libraries of variants that are used in methods involving recombineering, error-prone PCR (epPCR) was developed to perform random mutagenesis. Error-prone PCR protocols are modifications of standard PCR methods, designed to alter and enhance the natural error rate of the polymerase. Taq polymerase is commonly used because of its naturally high error rate. Creating high-quality libraries of random sequences is an important step in this process as it allows variants of individual molecules to be generated from a single-parent sequence. Combined with the synonymous PAM-inactivating mutation (SPM), precise genome manipulation with high efficiency can be achieve in a few steps using CRISPR gene editing methods [9].

Unfortunately, these methods are still failing in scalability, as the vast number of possible epistatic interactions erodes statistical power. Systematically testing interactions is challenging both from a computational and from a statistical point of view, given the large number of possible interactions to consider [10]. Thus, it is of interest to build models to predict genetic traits based on epistasis. Algorithms such as MINED were developed to detect significant pairwise epistasis effects that contribute the most to cell fitness using machine learning approaches [11]. Another study proposed a reinforcement learning approach, EpiRL, where epistasis is modeled as a one-step Markov Decision Process [12]. The use of machine learning in trying to find highly interacted genes could help tackle the challenges raised by the high-dimensionality of epistasis data.

# **Motivations & Hypotheses**

Improvements in DNA synthesis and sequencing have underpinned comprehensive assessment of gene function in bacteria. And currently, the genome mutagenisis techniques and study are using genetic transfer networks to make better predictions of the sequence or word sequence of an entire genome. But low-editing efficiencies and mutational biasing, is a downfall that needs to be attended to. It impacts greatly the quality of the fitness data. In recent years a small number of approaches have also achieved a high degree of effectiveness without mutational annotation.

The aim is to mesure the non-synonymous mutations and not the deletions. In this paper, we propose a novel model that achieves the goal. The idea is to investigate the functional basis of epistasis, and because rpoB plays a central role in transcription, we measured the effects of common rpoB mutations on transcriptional efficiency. Because mutations using the CREPE technology allows the study of combination of mutations. Two mutations are considered to be purely additive if the effect of the double mutation is the amount of the consequences associated with the particular single variations. This occurs whenever genetics are not linked with each other. Simple, component qualities were researched in early stages in the particular background of genes, they are usually fairly rare, along with many genes showing a minimum of some degree of association with epistatic connection.

We address this concept by measuring the particular fitness effect associated with rifampicin resistance mutations in the  $\beta$  subunit of RNA polymerase (rpoB) of Escherichia coli. Epistasis for fitness means that the selective effect of a mutation is conditional on the genetic background in which it appears. Epistasis can be easily seen in nature, the process in which it is involved is still not well known. Furthermore, its consequence on evolution, and its role in natural selection is still incomplete to our knowledge. The mutational path to high fitness genotypes is linked and even supposedly dependent of the genetic background in which novel mutations appear. This background effect is independent of the population as well as any loci of any gene. Sign epistasis has been defined in that the sign of the fitness effect of a mutation is under epistatic control [13].

Thus, using the CREPE method we can assess the fitness effects of individual mutations on the same loci, as well as on diverse location. All this in correlation with the stress to which the bacteria are exposed. We want to explore the theorical and empirical consideration implying the strong genetic constraint on the selective accessibility to high fitness genotype mutation path.

# Methods & Results of the Supporting Article

# Effects of Epistasis in Double Mutants

In this section, we replicated the analyses performed by the authors of the article supporting this report to better understand the impact of epistasis on rifampicin resistance. In

particular, we were interested in comparing the fitness of double mutants, compared to the sum of fitness from the respective single mutations.

### **Data Preprocessing**

The dataset provided by A. Choudhury was obtained by processing the sequencing output from a single biological replicate. Paired Illumina reads were assembled and aligned, and variant counts and amino acid changes were extracted from the qrowdot alignment output. Finally, variants were aggregated by grouping on the mutation positions and summing the read counts. The result is a dataset where each row corresponds to a unique genotype, with columns:

- aa\_change: list of amino acid changes, with original amino acid, position and new amino acid;
- pre: read counts before selection;
- ten: read counts after selection on 10 µg mL<sup>-1</sup> of rifampicin;
- fifty: read counts after selection on 50 µg mL<sup>-1</sup> of rifampicin;
- hundred: read counts after selection on 100 μg mL<sup>-1</sup> of rifampicin.

In this dataset, the wild-type corresponds to the row having the highest pre-selection read counts. The dataset was further processed using code available in the archive attached with this report and on a GitHub repository (https://github.com/sdiebolt/espcisbn-homework). First, non-synonymous mutations were extracted from the aa\_change column.

For each variant, the fitness was estimated as

$$f = \log\left(\frac{C_{i,\text{post}} + 0.5}{C_{\text{wt,post}} + 0.5}\right) - \log\left(\frac{C_{i,\text{pre}} + 0.5}{C_{\text{wt,pre}} + 0.5}\right),\tag{1}$$

where  $C_{i,post}$  and  $C_{i,pre}$  are the variant read counts, respectively post- and pre-selection, for condition  $i \in \{10 \,\mu\text{g mL}^{-1}, 50 \,\mu\text{g mL}^{-1}, 100 \,\mu\text{g mL}^{-1}\}$ , and  $C_{\text{wt,post}}$  and  $C_{\text{wt,pre}}$  are the wild-type read counts, respectively post- and pre-selection [14]. The  $\frac{1}{2}$  constant was added to each count to assist with very small counts. The standard error of this estimate was computed as

$$SE(f) = \sqrt{\frac{1}{C_{i,post}} + \frac{1}{C_{i,pre}} + \frac{1}{C_{wt,post}} + \frac{1}{C_{wt,pre}}}.$$
 (2)

# **Next Steps**

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