Physical enrichment of transposon mutants from saturation mutant libraries using the TraDISort approach

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ABSTRACT
Transposon-insertion sequencing methods are finding their way into the molecular toolbox of many fields of microbiology. These methods can identify the genomic locations and density of transposon insertions in saturated transposon mutant libraries and can be used to make inferences on gene function. For example, where no insertions or very few insertions are identified within a gene in a mutant library grown under permissive conditions, the gene may be essential. Furthermore, where mutations are enriched or lost in a gene after passaging the library through a selective process, the gene is likely to be involved in phenotypes linked to the process. Typically, a fitness based selection such as a stress condition is used in these experiments and the processed sequencing data are used to identify genes required for fitness under the selection. Our research team recently expanded the utility of the transposon directed insertion sequencing (TraDIS) method by applying a physical separation of a transposon mutant library mediated by fluorescence activated cell sorting, rather than a fitness-based selection. This approach, which we have named “TraDISort” is significant because it allows the study of phenotypes that are not linked to cell survival. The TraDISort approach has a broad range of future applications, in drug development, metabolic engineering and in studies of basic bacterial cell physiology.

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Introduction

The advent of high-throughput sequencing has sparked a new era in microbiology where genome scale experiments have become routine in most laboratories. The impact of these sequencing methods can be clearly seen in advances made to the fields of comparative genomics and metagenomics, and has driven the development of technologies that can elucidate gene function on a genome wide scale, such as transcriptomics, ChIP-seq (chromatin immunoprecipitation followed by sequencing), and most recently, transposon insertion sequencing on a genome wide scale. High-throughput sequencing provided a basis for “transposon insertion sequencing” methods to easily profile high-density libraries of individual random transposon mutants, allowing the insertion sites across the mutant population to be mapped and the relative abundance of each mutant to be determined. Using these methods the insertion site profile of a mutant library that has been subjected to a selection can be compared with the profile of the same library grown without the selection, to infer which genes and genetic elements are acted on either positively or negatively by the selection and thus the possible functions of these genes.

Several sample preparation protocols have been developed for mapping transposon insertion sites, and distinct studies using each method were first published in 2009. These include transposon directed insertion-seq (TraDIS), transposon sequencing (Tn-seq), insertion sequencing (INseq) and high-throughput insertion tracking by deep sequencing (HITS). The specific differences of each method, as
well as advantages and disadvantages are described in several excellent recent reviews.4,5

The majority of transposon-insertion sequencing studies that have been conducted to date have applied fitness based methods to differentially select between mutants in the population, e.g. growth during exposure to a stress condition. These can be conceptualised as massive scale competition experiments between all the mutant strains in the library; a loss of mutations in a particular gene across the library suggests that the gene is important for growth, survival or competitive fitness under the stress, an increase in mutations in a gene suggests that the gene is detrimental to growth, survival or fitness under the stress, and no change in the frequency of mutants for a particular gene suggests that the gene is not important for the fitness of the strain under the stress (Fig. 1).

Due to the huge potential to identify novel gene functions, fitness based transposon sequencing experiments have found a place in the molecular toolbox of many fields of microbiology including microbial ecology, industrial microbiology and medical microbiology. For example, these experiments have been used to identify genes that are involved in small molecule resistance or tolerance, including antibiotics in the opportunistic human pathogens Klebsiella pneumoniae10 and Staphylococcus aureus11 and industrial chemicals in Escherichia coli.12 Transposon sequencing experiments have been used to identify genes required for bacterial colonisation of animals, both in

Figure 1. Overview of a typical transposon insertion sequencing fitness-based experiment. (A) A mutant library in the strain of interest is constructed and plated on media permissive to growth. (B) The mutants are pooled and then cultured in permissive and selective conditions, e.g., laboratory media without or with an antimicrobial, respectively. (C) DNA is isolated from the cultured mutant populations and the transposon insertion sites and frequencies are determined using transposon sequencing. A reduction in the frequency of mutants carrying insertions in a particular gene grown under selective conditions compared with permissive conditions indicates that the gene is required for bacterial fitness under the selective condition (e.g., gene A). An increase in the frequency of mutants carrying insertions in a gene grown under selective conditions relative to permissive conditions suggests that there is an advantage to inactivating the gene under the selection used (e.g., gene B). An equal frequency of mutants with insertions in a gene in populations grown under both selective and permissive conditions suggests that the gene does not influence bacterial fitness under the selective condition (e.g., gene C). For some genes (typically around 10% of annotated genes in bacterial genome) there will be no insertions, or very few insertions in the initial mutant pool and after permissive growth (e.g., gene D). These genes are likely to be essential for bacterial survival even under permissive laboratory growth conditions and typically encode housekeeping functions, such as DNA replication. Overall, the information gathered in these experiments allows hypotheses on gene function to be formulated.
the context of symbionts, such as *Snodgrassella alvi* colonisation of honey bee guts,\(^1\) and in pathogenic interactions, such as *Salmonella enterica* serovar Typhimurium colonisation of food-producing animals\(^\text{14}\) and *Acinetobacter baumannii* colonisation of insect larvae as a virulence model.\(^\text{15}\) Furthermore, transposon sequencing experiments have been used to identify genes required for bacterial transitions between growth states, such as persister cell formation in uropathogenic *E. coli*\(^\text{17}\) and spore formation in *Clostridium difficile*\(^\text{17}\) and *Bacillus subtilis*.\(^\text{18}\) A very recent study used transposon sequencing to examine bacterial fitness under hundreds of unique selective conditions, including carbon and nitrogen utilization and chemical stress conditions, in 25 bacterial strains.\(^\text{19}\) Through this massive study, the authors report the identification of potential phenotypes for close to 8,500 proteins of unknown function, demonstrating the huge potential of these methods for assigning function to novel genes.\(^\text{19}\)

**Physical enrichment for mutants of interest using “TraDISort”**

In contrast to the fitness based mutant selection approaches used in other studies, we recently became interested in whether transposon sequencing methods could be used to directly probe for genes that define physical traits of bacterial cells that may not heavily influence their relative fitness under an easily imposed selective pressure; could we physically separate mutants of interest and then use transposon-insertion sequencing? Fluorescence activated cell sorting (FACS) is a well-established method that can be used to very rapidly screen millions of living cells (or other small particles) for their size, granularity and fluorescence, and sort these cells according to user-defined physical characteristics. Therefore, FACS provided an ideal method to impose a physical gating for mutant cell enrichment.

In our initial study, we set out to identify mutants of the human pathogen *Acinetobacter baumannii* that differentially accumulated the fluorescent dye ethidium bromide.\(^\text{20}\) The UV fluorescence of ethidium bromide increases significantly when it is intercalated into nucleic acids, a property that has been exploited for several decades in the highly sensitive detection of nucleic acids following gel electrophoresis. This property of ethidium bromide also means that it is differentially fluorescent inside and outside of cells—more fluorescent inside due to the high nucleic acid content. Therefore, the fluorescence associated with individual cells can be used as a measure of the amount of ethidium inside the cell. This method has been used extensively to study the function of multidrug efflux pumps that recognize ethidium as a substrate.\(^\text{21}\)

A large population (>100,000) of unique transposon insertion mutants of *A. baumannii* was incubated with a low concentration of ethidium bromide for a sufficient time to allow their intracellular ethidium bromide concentrations to reach equilibrium, i.e., when the rate of accumulation was equal to the rate of efflux. The cells were then subjected to FACS using a BD Influx flow sorter to enrich for sub-populations of mutants that contained the highest and the lowest concentrations of ethidium bromide, the top and bottom 2% of cells based on ethidium bromide fluorescence, respectively. We then used TraDIS sequencing protocols and analyses tools\(^\text{22}\) to profile the transposon insertion sites across these differentially fluorescent populations and compare them to the insertion site profile of the total mutant library pool that had been grown in parallel to FACS. The data showed that insertions in genes encoding various multidrug efflux systems, particularly *adeABC* and *amvA*, were positively and negatively selected in the high and low fluorescent pools, respectively (Fig. 2). This fits with the notion that inactivation of these pumps by transposon insertion would reduce the overall rate of ethidium bromide efflux and result in a higher equilibrium concentration of ethidium in these mutant cells. Therefore, these cells would be far less likely to show low fluorescence and be selected in the low fluorescent pool, but far more likely to show high fluorescence and be selected in the high fluorescent pool. In line with the differential selection for mutations in genes encoding efflux pumps, cells carrying mutations in efflux pump regulator genes were also highly differentially selected. Insertions in efflux pump activator genes showed similar patterns of selection to the efflux pump genes themselves, whereas, insertions in efflux pump repressor genes were much less abundant in the high fluorescent pool, since their inactivation leads to overexpression of efflux pumps, a higher rate of ethidium bromide efflux and a lower equilibrium concentration of ethidium.\(^\text{20}\)

The total amount of ethidium bromide in a cell will reflect not only its equilibrium concentration (i.e., the
sum of accumulation and efflux), but also the total cell volume, where larger cells will have more ethidium bromide, and possibly nucleic acids, and therefore typically higher fluorescence than smaller cells. Additionally, many bacterial strains, including A. baumannii BAL062 used in our initial study, can form aggregates in planktonic culture and the total fluorescence of these particles would be the sum of the aggregated cells. Consequently, we applied a gating procedure during our FACS cell enrichments to exclude any cell aggregates or large cells with division defects before fluorescence based sorting. As a result of this gating we observed a significant reduction in mutants carrying insertions in several cell division genes, and conversely a significant increase in mutants carrying insertions in genes that may promote aggregation, such as the csu type I pilus genes. Thus, another clear application of TraDISort is directly identifying cells involved in regulating and maintaining cellular size and shape, which could also be done with gating alone and without fluorescence.

Collectively, the data generated in these experiments show that flow sorting is a viable approach to enrich for mutants showing altered cellular phenotypes of interest before transposon insertion sequencing. Due to the combination of TraDIS sequencing and flow sorting, we have called this approach “TraDISort”. The TraDISort approach expands the utility of transposon insertion sequencing because it provides an opportunity to study phenotypes that are not directly linked to cell survival or regeneration, i.e.,
where there may be little significant difference in the fitness of mutants within the population, or where the mutants of interest may be less fit than the average.

**Future directions using TraDISort**

Following from our study examining ethidium accumulation into *A. baumannii* cells, TraDISort has significant potential as a new tool for investigating bacterial multidrug efflux pumps. In addition to ethidium bromide, there are many small molecule fluorophores that can be used to monitor the activity of multidrug efflux pumps. These compounds differ in their chemistry and their sites of accumulation within cells, so can be used to examine the function of different sub-sets of multidrug efflux pumps. These compounds differ in their chemistry and their sites of accumulation within cells, so can be used to examine the function of different sub-sets of multidrug efflux pumps. For example, similar to ethidium, Hoechst 33342, and 4′-6-diamidino-2-phenylindole (DAPI) are differentially fluorescent when intercalated into nucleic acids and total cell fluorescence could be used as a proxy for the intracellular concentrations of these dyes. However, whereas ethidium, and Hoechst 33342 are monovalent, DAPI is a bivalent compound so may be recognized by a different set of efflux pumps. Other dyes accumulate in the periplasm of Gram-negative bacteria rather than penetrating the cytosol and could be used in TraDISort to target the functions of efflux pumps that specifically capture their substrates from the periplasm. Finally, the genes involved in controlling the accumulation of fluorescent antibiotics, such as tetracyclines and fluoroquinolones, should be identifiable using the TraDISort approach and may also identify specific toxin transporter systems in addition to efflux. However, protocols are currently being fine-tuned to account for the low fluorescence of these compounds and the lack of differential fluorescence inside and outside bacterial cells. Once efflux systems that recognize these diverse substrates have been identified, TraDISort could further be used to identify the targets for efflux pump inhibitor compounds by treating mutant populations with these compounds in combination with fluorescent efflux pump substrates. Comparison of the mutants selected with and without the inhibitor should identify the pumps being inhibited and potentially the extent of inhibition.

A range of commercially available fluorescent dyes can be used as indicators for the intracellular concentrations of various free ions, such as sodium, iron, zinc and protons. Typically, these dyes undergo a fluorescence change, such as a shift in their excitation or emission spectra, or a shift in their fluorescence intensity when bound to their cognate indicator ion. TraDISort could be used in combination with these dyes to probe for the systems involved in homeostasis of these important ions.

Rather than monitoring fluorescence based on a small molecule fluorophore, the TraDISort approach may also be used in combination with genetic fluorescent reporter constructs. Fluorescent reporter systems incorporating a gene encoding a fluorescent protein have been used for many decades as tools to examine levels of gene and protein expression in various biological systems. Mutant libraries built around strains carrying promoter fusion fluorescent reporter constructs could be used to identify novel regulators that recognize sequence elements in the cloned promoter region. In these experiments mutants showing increased or decreased expression of the fluorescent reporter could be enriched and the insertion sites mapped. The enriched mutants may harbour mutations in the regulators of the gene of interest.

TraDISort has significant potential to be used as a tool in metabolic engineering and synthetic biology. The goal of a typical metabolic engineering project is to develop bacterial strains that can be used as factories to produce a small molecule or molecules of interest, which may be native to the producing strain, or the product of introduced metabolic pathways. A major challenge is to channel metabolic energy toward production of the small molecule(s). The development of a commercially viable production strain typically involves many rounds of rational design, synthetic strain construction and product yield testing. Random mutants, including transposon mutants have been used to assist in the design of high yielding strains. However, these studies have traditionally relied on the isolation and characterization of single isolated mutants, and are thus prone to complications with high numbers of false positives. With recent advances in the development of fluorescent biosensors, saturation transposon mutant libraries could be readily applied in metabolic engineering in a highly-streamlined approach using TraDISort. Fluorescent biosensors responsive to the product of interest would allow mutants that produce higher levels of the product to be enriched by flow sorting approaches in TraDISort and avoid potential for false positives. Such mutants would typically have a lower overall fitness, due to the
burden of channelling more metabolic energy into compound production rather than growth, but may be identified using the TraDISort approach because mutant enrichment is separated from cell fitness.

Conclusions

Transposon insertion sequencing technologies have brought about a new age in genome-wide investigations of gene function. The relative ratios of mutants in a large transposon mutant library can be examined before and after exposure to a selective condition to infer which genes are required for fitness under the selection. Rather than applying a fitness based selection, mutants can also be enriched based on physical characteristics by FACS, as used in the TraDISort approach. By separating mutant enrichment from fitness, TraDISort allows transposon sequencing to be used in a broader range of experiments. For example, some of the most significant impacts of this new technology may be within metabolic engineering and synthetic biology, where fluorescent biosensors could be used with TraDISort to identify mutants that produce superior yields of compounds of interest.

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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