Research Article

Error Analysis of Deep Sequencing of Phage Libraries: Peptides Censored in Sequencing

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Next-generation sequencing techniques empower selection of ligands from phage-display libraries because they can detect low abundant clones and quantify changes in the copy numbers of clones without excessive selection rounds. Identification of errors in deep sequencing data is the most critical step in this process because these techniques have error rates >1%. Mechanisms that yield errors in Illumina and other techniques have been proposed, but no reports to date describe error analysis in phage libraries. Our paper focuses on error analysis of 7-mer peptide libraries sequenced by Illumina method. Low theoretical complexity of this phage library, as compared to complexity of long genetic reads and genomes, allowed us to describe this library using convenient linear vector and operator framework. We describe a phage library as $N \times 1$ frequency vector $n_i$, where $n_i$ is the copy number of the $i$th sequence and $N$ is the theoretical diversity, that is, the total number of all possible sequences. Any manipulation to the library is an operator acting on $n$. Selection, amplification, or sequencing could be described as a product of a $N \times N$ matrix and a stochastic sampling operator ($Sa$). The latter is a random diagonal matrix that describes sampling of a library. In this paper, we focus on the properties of $Sa$ and use them to define the sequencing operator ($Seq$). Sequencing without any bias and errors is $Seq = Sa I_N$, where $I_N$ is a $N \times N$ unity matrix. Any bias in sequencing changes $I_N$ to a nonunity matrix. We identified a diagonal censorship matrix ($CEN$), which describes elimination or statistically significant downsampling, of specific reads during the sequencing process.

1. Introduction

In vitro selection experiments—such as phage display [1, 2], RNA display, SELEX, and DNA aptamer selection [3, 4]—employ large libraries, from which $10^2$–$10^6$ active sequences are identified through iterative rounds of selection and amplification. With the recent emergence of deep sequencing, it became possible to extract a large amount of information from the libraries before and after selection [5–10]. Deep examination of the library is a promising technique for direct evaluation of binding capacities of all binding sequences from one panning experiment. Deep sequencing also allows the characterization of unwanted phenomena in selection, such as amplification bias [6, 11].

Analysis of $10^6$ reads by deep sequencing gave rise to a large number of errors that were not present in the analysis based on the small number of sequences obtained using the Sanger method. Analysis of errors in information-rich datasets is a problem with over 50 years of history; correction of digital data made of bits or words is a topic of intense research in communication theory [12]. As phage display operates with limited digital sets, data analysis techniques from the communication theory could be applied to phage display. For example, Rodi and coworkers used a positional frequency matrix to calculate the informational content or Shannon entropy of each sequence [13]. This approach could be used to distinguish potential fast growing sequences from potential hits [14]. With the introduction of deep sequencing, the problem of error analysis in phage display becomes identical to a classical information theory problem: “reproducing at one point, either exactly or approximately, a message selected at another point” [15]. The “message” is the sequence information stored in the library. Sequencing process transmits this information and makes either stochastic or predictable errors. Understanding the sources of errors during sequencing could provide mechanisms for bypassing
them, for correcting the errors, and for maximizing the amount of useful information received from sequencing.

There are over 10,000 published literature reports that contain the terms “deep sequencing” or “next generation sequencing” or any of the trademark names such as “Illumina” (reference: ISI database). Among these reports, less than 10 published reports describe sequencing of phage-displayed libraries [5–7, 9, 10, 16–19]. Deep sequencing efforts in the literature are largely focused on genome assembly and metagenomic analyses. The error analysis techniques tailored for genome assembly cannot be used directly for analysis of phage libraries because the data output from phage library sequencing is very different from the genome assembly. In genome assembly, genomic DNA is shredded into random fragments and sequenced. The genome is then assembled from these fragments in silico. Although multiple fragments cover each area of the genome, the probability to observe two identically shredded fragments is very small. Two exact sequences, thus, could be considered amplification artifacts and removed by error analysis software. On the contrary, in phage-display sequencing, the reads are exactly of the same length. Duplication of the same read is important for validation of the accuracy of this read. Some researchers focus exclusively on reads that have been observed multiple times and discard singleton reads as erroneous [5]. Within each library, the copy numbers of sequences range continuously by six or more orders of magnitude [5, 6, 9]. Some phage clones are observed in the entire library only a few times; other clones could be present at copy number of 100,000 per sequencing run [5, 6, 9]. Unlike multiple cells with identical genomes, each screen is unique: identical set of sequences with identical copy numbers cannot be obtained even if the screen is repeated due to stochastic number of the screen that contains low copy number of binding clones [20].

Metagenomic analyses of microorganisms recovered from environmental samples [21, 22], also known as “microbiome” [23] and “viriome” analyses [24], encountered similar problems to those observed in phage library analysis: the concentration of species observed in a particular sample is unequal [25]. The abundance of species might range by a few orders of magnitude [26]. It is possible that error analysis tools developed in the above areas could find use in phage display sequencing. For example, there are multiple published algorithms for removing errors from low copy number reads to ascertain that low copy number sequences are new species and not sequencing errors (e.g., see [27–29] and references within). Metagenomic analysis is usually more complex than analysis of phage-display libraries. First, in metagenomics, the bacterial or viral genes must be assembled from short reads de novo. Second, there is no simple relationship between phylogenetic classification of “species” and the observed DNA sequence. Third, the exact number of species in the environment is unknown. On the other hand, sequencing of phage-displayed peptide libraries has none of these problems: (i) it requires no assembly steps because each sequence is covered by one read; (ii) a unique DNA sequence defines a unique “species”; and (iii) the theoretical complexity in synthetic libraries is known exactly. For small libraries, such as the library of 7-mer peptides, the complexity, (20), is within the reach of next-generation sequencing. We see phage-displayed peptide libraries as an ideal model playground for the development of optimal error analysis and error correction protocols. It is possible that error analysis developed from phage libraries analysis could then be used in other areas such as genomic and metagenomic analyses.

The errors in sequencing could be divided into “annotated” and “invisible.” The “annotated” errors that originate from misincorporation of nucleotides are annotated using Phred quality score [30]. These annotated errors are removed during the processing (see below). Examples of “invisible” errors are sequence-specific frame shifts that lead to emergence of truncated reads during the illumina sequencing [31]. Invisible errors could also originate during the preparation of the libraries for sequencing. Examples are removal of AT-rich fragments during purification of dsDNA [32] and erroneous incorporation of nucleotides during PCR [33, 34]. Mutations have the most significant impact on the observed diversity of the library. There are 63 ways to misspell a 21-mer-nucleotide sequence with a one-letter error (point mutation). The large dynamic range in concentrations of clones in the phage library exacerbates the problem. Clones that are present in high abundance—10\(^5\) copies per read—are more prone to yield errors [6]. For example, we observed that random point mutations convert several short sequence with a copy number of 10\(^{-5}\) to a library of sequences with copy numbers ranging from 1 to 10\(^5\) [11]. In attempt to unify error analysis into one convenient theoretical framework, we generalized all errors as follows. All errors either lead to disappearance of particular sequence or its conversion to another sequence of the same length. Errors, thus, operate within a finite sequence space, and it should be possible to use elementary linear algebra to generalize most processes that lead to errors.

### 2. Theoretical Description

See Table 1.

#### 2.1. Operator Description of the Phage-Display Library and Selection Process

In our previous reports, we described the phage library as a multiset, or a set in which members can appear more than once [35]. This description also simplifies the analysis of the errors in these libraries. The multiset description represents a library with N theoretical members as an ordered set of N sequences and N \(\times\) 1 copy number vector (n) with positive integer copy numbers (Figure 1(a)). Any manipulation of a phage library—such as erroneous reading or selection—changes the numbers within the copy number vector. All manipulations to the multiset, thus, could be described by operators (Op) that convert vector \(n_1\) to another vector \(n_2\) as \(n_2 = \text{Op} n_1\) (Figure 1(c)). For an N \(\times\) 1 vector, the operator is \(N\times N\) matrix. If elements are selected or eliminated independently of one another, the \(N\times N\) matrix is diagonal (Figure 1(d)). This approach is uniquely convenient for libraries of short reads. For example, a library of 7-mers contains exactly \(20^7 = 1.28 \times 10^9\) peptides and is described...
Sel is the copy number vector for naïve library, Sel is the copy number vector after panning, and Pan is a panning operator. In standard phage display, the Pan operator is a complex product of all manipulation steps (binding, amplification, dilutions, etc.). If a screen uses no sampling procedure:

(1) \[ \text{Sel} = \text{Pan \ Naive}, \]

where Naive is the copy number vector for naïve library, Sel is the copy number vector after panning, and Pan is a panning operator. In standard phage display, the Pan operator is a complex product of all manipulation steps (binding, amplification, dilutions, etc.). If a screen uses no sampling procedure:

(2) \[ \text{Pan} = {\mathbf{1}}_{\text{Sa}} \text{K}_a, \]

select = \[ {\mathbf{1}}_{\text{Sa}} \text{K}_a \text{Naive}, \]

where \( \text{K}_a \) is a deterministic "association" operator, which contains association constants for every phage clone present in the library. Description of such operator is beyond the scope of this paper and we recommend consulting other reports that attempted to generalize the selection procedure [20]. Another operator in (3) is a sampling operator \( {\mathbf{1}}_{\text{Sa}} \), which describes stochastic sampling of the library with m sequences to yield a sublibrary with \( \mathbf{f} \) m-sequences, where \( \mathbf{f} \in [0, 1] \) is a sampling fraction. \( {\mathbf{1}}_{\text{Sa}} \) operator has the following properties, which emanate from physical properties of the sampling procedure:

(1) \[ {\mathbf{1}}_{\text{Sa}}, 0 = 0 \text{ (sampling does not create new} \]

sequences from nonexisting sequences).
Any phage library is a multiset defined by sequence set and copy number vector
\[ \text{Lib} = [S, n] \]

\[ \text{Lib} = \left\{ \begin{array}{c} \text{sequence}_1 \text{sequences}, n_1, n_2, \ldots, n_k \end{array} \right\} \]

- \( S \): number of all possible sequences in the library
- \( n \): copy number of the \( i \)th sequence in the library

Any manipulation of library is an operator acting on the \( n \) vector (copy number vector)

Example: sampling of the library

Sampling operator \( (Sa) \) acts on \( n \) vector

\[ \begin{array}{c} 1_n = \begin{pmatrix} n_1 \\ n_2 \\ \vdots \\ n_k \end{pmatrix} \\ \text{Lib} = [S, n] \end{array} \]

\[ \begin{array}{c} 2_n = Sa \cdot 1_n \end{array} \]

Some elements present in 1st library are "gone" from the 2nd library

Properties of \( Sa \):
1. \( Sa(\cdot \cdot \cdot(sa(sa(n))) \cdot \cdot \cdot) = sa(n) \)
2. \( 1_k \sum_s sa(n) \rightarrow n, \text{as } k \rightarrow \infty \)

Multiset description of the phage display selection

Theoretical library

\[ T = [S, 1] \]

Synthesis of nucleotides

\[ S = [S, m_n] \]

The \( i \)th sequence \( S_i \) has copy number \( m_n \)

Naive library

\[ N = [S, n_{\text{ naïve}}] \]

Ligation, transformation, growth

\( N \) cannot be measured. They have to be approximated from the "observables".

Selected (panned) library

\[ P = [S, m_{\text{pan}}] \]

Panning, amplification

Sequencing

P and N cannot be measured. They have to be approximated from the "observables":

\[ \begin{array}{c} d_{n} = [m_{\text{ naïve}}, n] \\ \text{seq} = \{1, n = 0 \text{ Kronecker}0, n \neq 0 \text{ delta} \} \end{array} \]

Implementation—the entire code is <30 lines in MatLab—the script allows rapid calculation of the results of \( Sa \) for a multiset of reasonable size (several million sequences, Figures 4 and 5) on a desktop computer.

We evaluated the behaviors of \( 0.5 Sa \) for several multisets. The probability to observe a specific solution is described in Figure 3(b). Individual solutions can be represented as lines with nodes on XY-plane, where each node represents one element of the multiset (Figures 3(d) and 3(e)). The most probable solutions reside near the "expected solution" (represented as dotted line), and the probability to observe a solution where many elements deviate from the probable solution is low (Figure 3(e)). Graphical representation of the solutions highlights that sampling could lead to deviation of the frequency of the individual elements of the multiset; for example, Figure 3(e) describes >2 fold deviation from...
Analysis of all probable solutions of multiset (Figure S2). This redundancy makes the calculations solutions (e.g., solutions that have equal probability in $B = \{S_a\}$) converge rapidly. Each step, thus, is an operator changing the $n$ vector. (b) If we ignore bias during preparation, operators could be approximated as unity vectors, and sequencing could be represented as a product of sampling and analysis operators. (c) Analysis operator ($A_n$) is a binary decision matrix, which describes what sequences are and are not considered as errors. Decisions, such as removal of sequences or correction of sequences, are the most important because they decide which “observed” sequences are considered “real.” To make the analysis of the selection process meaningful, the same $A_n$ operator should be used in all analyses.

![Diagram](image_url)

Figure 2: Operator description of the deep sequencing process. (a) A library of phage must be processed before deep sequencing. Each step involves sampling, which is either a deliberate partitioning of the sample or random loss of the sample. Each sample preparation step could (and does) introduce bias in sequence abundance. Each step, thus, is an operator changing the $n$ vector. (b) If we ignore bias during preparation, operators could be approximated as unity vectors, and sequencing could be represented as a product of sampling and analysis operators. (c) Analysis operator ($A_n$) is a binary decision matrix, which describes what sequences are and are not considered as errors. Decisions, such as removal of sequences or correction of sequences, are the most important because they decide which “observed” sequences are considered “real.” To make the analysis of the selection process meaningful, the same $A_n$ operator should be used in all analyses.

The sampling operator is critical in phage display because sampling of libraries occurs in every step of the selection and the preparation of libraries for sequencing. The stochastic nature of sampling operators makes two identical screens “similar within a confidence interval.” Solving (1) exactly is not possible, but it should be possible to estimate the solution within a confidence interval. Consider

$$\text{Sel} \in \left[ k_{\text{Naive}} \right],$$

where $k_{\text{Naive}}$ and $k_{\text{Naive}}$ are diagonal matrices of the upper and lower confidence intervals for the association constants.

A simulation of the behavior of the $A_n$ operator (Figures 3 and S3) suggests that the relative sizes of the confidence intervals might be impractically large when the copy numbers of sequences are <10.

Multiple sampling events of the $A_n$ operator yield a normal distribution for each element of the vector (Figure 3). Fitting this normal distribution could yield a “true” value of the process. This process is identical to the extrapolation of the average from the normal distribution of noisy data. Multiple algorithms for such extrapolation exist for one- and multidimensional stochastic processes [38, 39]. We believe that $A_n$ behaves as a one-dimensional stochastic process and it might be possible to extrapolate the true value of the sampling from 7 to 10 repeated instances of $A_n$ (i.e., the number of data sufficient to fit an 1D normal distribution).

The necessary practical steps towards solving (3) or (10) are the following. (i) Eliminate or account for any bias not related to binding (e.g., growth bias). (ii) Repeat the screen
Figure 3: (a) Testing the sampling operator implemented as random indexing function using a model multiset. (b) In 100,000 trials, we observed 22 unique solutions from which 14 resided in a 95% confidence interval. Solutions with 0 and 1 copies of element A were found at equal abundances (“redundant solutions”). (c) Representation of the most probable solution as a line with 4 nodes; “probability to find the solution; dotted line is an expected “average solution” for 50% sampling. (d) The 5th most probable solution; (e) least probable solution deviates the most from the average; (f) combination of all solutions. Red thick lines describe the most probable solutions; thin blue lines describe the least probable solutions. (g) Sampling of larger multisets yields more possible solutions (here, 2957 in 5000 trials). (h) All solutions of the sampling represented as lines. (i) Probability to observe a particular copy number after sampling. While (h) is the most accurate representations of the confidence intervals, the thin blue lines describe solutions outside the confidence interval; this representation is impractical due to large number of redundant solutions in larger multisets. In (i), confidence interval could be extrapolated from distributions of individual copy numbers (e): red dots are on or outside the confidence interval.

several times. (iii) Measure all copy numbers of all sequences, including zero values, with high confidence. Requirement (i) has been an ongoing effort in our group [11, 40] and other groups [13, 41–43]; for review see [11, 44]. Deep sequencing makes it simple to satisfy requirement (ii) and obtain multiple instances of the same experiment. For example, we described the Illumina sequencing method that allows using barcoded primers to sequence 18 unrelated experiments in one deep
sequence number of data in one point

- >1000 copies
- 100–1000 copies
- 10–100 copies
- 1–10 copies

Copy number before the sampling

Copy number after the sampling

Number of data in one point

- >1000 copies
- 100–1000 copies
- 10–100 copies
- 1–10 copies

Figure 4: (a) Testing the sampling operator using a large multiset made of 1000 unique elements with 1000 different copy numbers. Images describe linear and log-scale representation of the confidence interval of the sampling operator. Solutions beyond this interval were not observed in 5000 trials. Dotted line represents an overestimate of the 99.9% confidence interval (for details, see Figure S4). Most probable outcomes of the Sa operator have either zero or one unique sequence beyond this interval. This line is used in subsequent sections (Figures 5 and 6). We note that distributions of the copy numbers have well-defined shape; according to central limit theorem, it is a normal distribution. With enough replicas, it should be possible to extrapolate the center of this distribution, define the solutions explicitly, and bypass the stochastic nature of the Sa operator.

sequencing experiment [45]. We recently scaled this effort to 50 primer sets and evaluated the performance replicas of simple selection procedures (in preparation).

The measurement of the copy numbers of sequences is a separate problem that can be described using the same sampling operators and bias operators that describe how the library is skewed by each preparation step. For example, isolation of DNA by gel purification disfavors AT-rich sequences, whereas PCR favors sequence with within specific GC-content range [32]. The real sequence abundance in any phage library (\(n_{real}\)), hence, has to be derived from the observed sequence abundance (\(n_{obs}\)) by solving this equation:

\[
\begin{align*}
\text{(7)}
\text{obs}_n &= (f_{5} \text{ Sa } A_n)(f_{4} \text{ Sa } \text{Seq})(f_{3} \text{ Sa } \text{PCR}) \\
&\times (f_{2} \text{ Sa } \text{Is})(f_{1} \text{ Sa } \text{Gr}) \text{real}_n.
\end{align*}
\]

In this equation, each operator in brackets describes a bias at a particular step. \(\text{Sa}\) describes sampling at that step, and \(f_1–f_5\) describe the sampling fractions. The bias in growth (Gr), isolation (Is), PCR amplification (PCR), and sequencing (Seq) could be related to the nucleotide sequences. The An analysis operator is a matrix that describes retaining, discarding, or correcting the sequence (Figure 2(b)). An ideal An operator could compensate for the biases introduced by another operator (Figure 2(c)). To define such operator, (7) could be potentially solved using repeated sequencing of a well-defined model library. In the next applied section, we examine the real deep sequencing data and identify conditions under which these operators could be at least partially defined.

2.2. Analysis or the Error Cutoff in Deep Sequencing Reads. All next-generation sequencing techniques provide quality score (Phred Score) for every sequenced nucleotide. In Illumina sequencing, this score is related to the probability of the nucleotide being correct [46]. In low throughput Sanger sequencing, the Phred score monotonously decreases with read length and the mechanisms that yield errors in capillary electrophoresis are well understood. Common practice in Sanger sequencing is to discard all reads after the first nucleotide with a Phred score of 0. In next-generation sequencing, the filtering of the reads is usually more stringent as follows.

(A) Discard reads that have at least one read that has score lower than “cutoff.”
(B) Discard reads that had cumulative Phred score lower than cutoff.
The reads have been parsed to identify adapters and barcodes.

(b) Multiset view of the intermediate library. The library contains subsets that have low, medium, and high quality reads. Error filtering of this intermediate library to eliminate any read with Phred score below 30 yields a high quality library of reads $30^L$. (c) Mean accuracy of the reads in the library after error filtering ranges from 95% to 99.6%. Even for very low-quality cutoff, Phred > 1, the average read quality is 95%. (d) Distribution of cumulative read accuracy in libraries processed using different cutoffs. (e) Linear plot of the data presented in (d) with zoom in on the region with >90% cumulative accuracy.

(C) Use a combination of A and B (accept reads with minimal cutoff and minimal cumulative score).

Many of the error analyses in the area of deep sequencing are designed for genetic reads, which have variable lengths and unknown sequence throughout the whole read. Analysis of the reads in a phage-display library is a simpler problem because phage-derived constant adapter regions flank the variable reads. Identification of the adapter region is a necessary first step in the analysis. Reads, in which the adapters cannot be mapped, cannot be used. We designed algorithms to recover reads, even if adapters were hampered by truncation, deletion, or mutation [6]. We observed that the reads flanked by the erroneous adapters had a significantly higher error rate than reads flanked by “perfect” adapters (unpublished). Example of mapping of flawed reads in Illumina sequencing is provided in ERROR_TAG_data0001.txt (see Section 4). In the remaining sections, we analyze the population of the sequences preceded by a “perfect” adapter to identify possible sequence-specific biases.

We analyzed a typical library sequenced by Illumina using various cutoffs (Figure 5). We analyzed a 33-bp segment of the library that contained variable seven amino acids and a constant region and GGGS terminus. A simple cutoff that discards reads with Phred <1 nucleotides yields library
termed $1n$, which had an average 95% accuracy of the 33-nucleotide read. Reads that do not contain Phred = 0 nucleotide rarely contain multiple low-quality reads. The $1n$ library was bimodal: 80% of the reads had overall accuracy of 99%, very few reads with accuracy 5–90%, and significant number of reads with accuracy of 1% (Figures 5(d) and 5(e)). These observations suggest that reads can be divided into (i) reads free of errors and (ii) reads with multiple errors.

An example of a more stringent cutoff is elimination of reads with Phred < 13 nucleotides; this process yielded a library $13 n$ in which every nucleotide had > 95% confidence. The number of total reads in $13 n$ was 10% less than number of reads in $1n$; that is; sum($13 n$) = 0.9sum($1n$). The average accuracy of the reads in the $13 n$ library was 99.2%. Theoretically, the 0.95 confidence cutoff in a 33-nucleotide read could yield reads with accuracy as low as (0.95)$^{13} = 18\%$. In practice, the probability to find reads with multiple nucleotides of 95% accuracy was vanishingly small. Specifically, among 500,000 reads, the lowest observed cumulative accuracy was 77%. Such a result, for example, could be obtained in a sequence that has 27 “perfect” nucleotides and 5 nucleotides with a Phred = 13 score: (1)$^{37}$ (0.95)$^5 = 0.77$. Applying the most stringent cutoff to eliminate all reads with a Phred < 10 yielded a library $30 n$ in which every nucleotide had 99.9% confidence. The average confidence of the reads improved slightly from 99.2% to 99.6%. The number of total reads in $30 n$ was 30% less than number of reads in $13 n$; that is, sum($30 n$) = 0.7sum($13 n$). It was not clear whether such cutoff is an improvement or a detriment for analysis. In the next section, we examined how frequency of the members of the library changed upon application of each error cutoff.

2.3. Example of Error Analysis: Sequence-Specific Censorship during Phred Quality Cutoff

If errors occur by random chance, they should be uniformly distributed in all sequences. Removal of erroneous read, in that case, should be identical to sampling of the library by $1 Sa$ operator, where f is the sampling fraction. For example, consider the removal of Phred < 13 nucleotides from an unfiltered library (process denoted as $1n \rightarrow 13 n$). From the experiments, we know that sum($13 n$) = 0.9sum($1n$); if errors were distributed in sequences at random, the $1n$ and $13 n$ vectors should be related as

$$13 n = 0.9 Sa (1n).$$  

(8)

The solutions should reside within a confidence interval

$$13 n \in [\log C_{hi} C].$$

(9)

If errors occur preferentially in specific reads, the frequency of these reads should occur beyond the confidence interval of the $0.9 Sa$. This process could be described by a diagonal matrix $Bias$ as

$$13 n = 0.9 Sa (Bias(1n)).$$

(10)

The elements of the diagonal matrix $Bias$ = $\|B_{ij}\|$ could be estimated as follows:

$$13 n_i \in [\log C_{ij} ^{hi} C_{ij}], \quad B_{ij} = 1,$$

(11)

$$13 n_i < \log C_{ij}, \quad B_{ij} = \frac{13 n_i}{(0.913 n)},$$

(12)

Figure 6(c) describes the representative solution of the $0.9 Sa(1n)$ (green dots) and the confidence interval (blue lines). Supplementary Scheme S3 describes the script that calculated this interval from multiset $1n$, described as a plain text file PhD7-Amp-0F.txt, using 10,000 iterative calculations of $0.9 Sa(1n)$. This calculation required ~2 hours on a desktop computer. Confidence interval was estimated as the minimum and maximum copy number found after 10,000 iterations. In this approximation of the confidence interval, for sequences with the copy number < 10 before sampling, it was impossible to determine whether the sequence disappeared due to random sampling or due to bias. The values of $Bias$ operator cannot be defined for these sequences and it could be assumed to be 1 (see (11)). For copy number > 10, however, sequence-specific bias can be readily detected. We observed that the removal of Phred < 13 reads yielded a multiset in which a large number of sequences deviated beyond the confidence interval (Figure 6(d)). Their sequences could be readily extracted by comparing the vector $13 n$ with the vector of the lower confidence intervals $\log C$ (see (12)). The solution of the $Bias$ can be illustrated graphically (Figure 6(e)). Top 30 censored sequences are listed in Table S1; the other sequences can be found in the supplementary information (file PhD7-Amp-0F-13F-CEN.txt).

We performed similar calculations for $1n \rightarrow 30 n$ and $13 n \rightarrow 30 n$ processes. The latter process is the most interesting because $13 n$ library has all nucleotides within acceptable confidence range (> 95%) and the distribution of cumulative quality suggested that errors, on average, do not cluster in one read (Figure 5). The $13 n \rightarrow 30 n$ conversion eliminated 30% of the reads, and copy numbers of many sequences deviated significantly from the random sampling; these sequences are represented by green dots outside the blue confidence interval in Figure 6(h). Top 30 sequences are listed in Table S2. The censorship is not only sequence-specific, but also position-specific. In sequences that had been censored during the $13 n \rightarrow 30 n$ process, lower quality reads clustered around 3-4 specific nucleotides (supplementary information Figure S5).

The mechanism that leads to the disappearance of censored sequences is not currently clear. We attempted to identify common motifs in censored sequences using two approaches: (i) clustering and principal component analyses based on Jukes-Cantor distance between sequences and (ii) identification of motifs using multiple unique sequence identifier software (MUSI) [17]. These approaches could not detect any property common to censored reads, which would make them significantly different from the other, noncensored reads. Still, we hypothesize that the observed
Sequencing results
Removal of errors

\[ \text{Ph} > 30 \]
\[ L = [S, 0n] \]

\[ 30n = \text{Bias Sa} \]

30n=
Bias Sa

All nucleotides

\[ 0L = [S, 0n] \]

\[ 30L = [S, 30n] \]

Phred > 30

Ph > 13

Ph > 10

Ph > 0

Ph > 30

Ph ≥ 1

Ph > 30

Ph > 13

Ph > 0

Ph > 30

Ph ≥ 1

Ph > 30

Ph > 13

Ph > 0

Ph > 30

Ph ≥ 1

Ph > 30

Ph > 13

Ph > 0

Ph > 30

Ph ≥ 1

Figure 6: (a) Operator and multiset description of the error filtering procedure. Applying a Phred > 30 cutoff to library filtered by Phred > 1 cutoff \( (1n) \) yields a subpopulation of the library \( (30n) \). If errors are sequence-independent, the \( 1n \rightarrow 30n \) process should be identical to random sampling \( (30n = Sa1n) \). Any sequence-specific bias \( (\text{Bias}) \) should be detected as deviation from \( Sa1n \). (b) Progressive sampling with more stringent cutoff. (c) Theoretical \( Sa1n \) and theoretical 99.9% confidence interval (blue). (d) Observation of statistically significant deviation from \( Sa \) operator: dots beyond the blue line represent sequences prone to bias. Red dots represent sequences that disappeared after \( 1n \rightarrow 30n \) process or during \( Sa1n \) sampling. (e) Magnitude of the bias range from 5 to 100-fold. (f) Bias in sampling of Phred > 30 data from Phred > 1 data ((f) is theory, (g) is observed). (h) Bias upon sampling of Phred > 30 data from Phred > 13. Many sequences were lost in this sampling and this loss was statistically significant beyond the 99.9% interval. This result shows that some sequences have propensity to harbor low- and medium-quality reads. Distribution of the errors is sequence specific.

censorship represents sequence-specific errors, which occur in every time such sequence passes through the Illumina analyzer. For example, the sequences listed in Tables S1 and S2 and supplementary files were censored in five independent experiments, which were pooled and processed simultaneously in one Illumina run. Analysis of other instances of Illumina sequencing performed by other groups could help prove (or disprove) that censorship is indeed sequence-specific.
and experiment-independent. Sequence-specific censorship during Illumina analysis has been described in other publications [46]. The observations presented above suggest that reading of some sequences in phage libraries does not yield an accurate copy number. Even if these sequences were enriched due to binding, their apparent copy number in sequencing would be decreased due to sequencing bias. If the magnitude of bias is known, however, such error could be corrected. We anticipate that other biases could be calculated for these and other libraries in similar fashion. Their calculation extends beyond the scope of this paper and it will be performed in our next publication.

3. Discussion

3.1. Significance and Transformative Potential of Library-Wide Error Correction. In the Medicinal Chemistry field, structure-activity relationships (SAR) and pharmacophores are built using both positive and negative observations. It is the negative results that bear the most significance in these studies because they allow mapping of the range of conditions under which particular structure no longer works. For example, SAR of an R group of a ligand might be built on the following observations. A ligand binds to the target when the R group in the specific position is methyl or ethyl; changing R to iso-propyl and tert-butyl ablates the binding. This concludes that the R group must be a small alkyl group. An analogous situation is found in SAR of peptide ligands; the most important information from alanine scan mutagenesis is loss of function because it helps identifying the important residues. Interestingly, loss-of-binding conclusions are never applied to phage-display. The phage-display field is driven by positive results. Most publications report and follow up only on sequences enriched in the screen and consider only large copy numbers interesting. All papers focus on sequences that were found. Very few papers in phage display ask why other sequences were not found.

One of the reasons why phage display is not used for SAR-type analysis is because negative observations in phage library cannot be determined with high confidence. From a practical point of view, measuring zero with high confidence requires the largest number of observation (the highest depth of sequencing). The payoff, however, is immense: one screen with “confident zeros” could potentially yield SAR for every possible substitution of every possible amino acid. We refer to this (theoretical) possibility as “Instant SAR,” and its condensed theoretical form is described in (3) or (9) and (10). This paper demonstrates that the depth of sequencing is not the only problem towards this goal. Accurate estimate of negative results requires complete characterization of the origins of errors in sequencing which yield false negative values by censoring certain sequencing. Other types of censorship, such as growth bias, should be characterized and eliminated as well. As the phage display field is currently focused on positive results, the need for optimal error corrections and recovery of erroneous reads is low. With the rise of SAR-type applications in phage display, error correction will be recognized as the most significant barrier because it could lead to improper assignment of low frequencies and negative results. Improved error correction strategies could assign a lower confidence to the sequence instead of eliminating the errors and labeling them as confident zero. Proper mathematical framework, possibly similar to the one used in this paper, could be then used to carry all confidence intervals through calculations to yield reliable SAR-type data.

We note that the framework described in this paper is suitable for the analysis of the selection from libraries in which the diversity of the libraries before and after selection could be covered entirely by deep sequencing. With the current depth of sequencing, it corresponds to medium-scale libraries of \(\sim 10^6\) random members and affinity-matured libraries that contain \(\sim 10^6\) point mutations. We are in the process of generating these medium-scale libraries and running selection procedures that will allow us to apply and refine our framework. In the future, as technical capabilities and depth of sequencing increase, the process would be applicable to larger libraries as well.

4. Methods

4.1. Generation of Z-Bars and Other Visualization Techniques. Sequencing of the libraries has been described in our previous publications [6, 47]. All data visualization in this paper was done by MATLAB scripts; raw *.eps output from MATLAB scripts subject to minor postprocessing in Adobe Illustrator to adjust fonts relative dimensions of plots. Core scripts are described in the supplementary information. Other scripts are available in our previous publication [47]. Illumina files used for the analysis can be found in the directory at http://www.chem.ualberta.ca/~derda/mathbiology/; the file ERROR_TAG_data0001.txt is an example of error-tagged reads; PhD7-Amp-xxF.txt is the library filtered with xx Phred cutoff (xx = 1, 13 and 30); file PhD7-Amp-13F-30F-CO.txt describes confidence intervals for Phred(13) to Phred(30) filtering process; other files with "-CO.txt" extension describe confidence intervals of other processes. Supplementary Figures S1–S4 and Schemes S1 and S2 describe MATLAB implementation of the Sa operator.

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