Computational enhancement of single-cell sequences for inferring tumor evolution

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Abstract

Motivation: Tumor sequencing has entered an exciting phase with the advent of single-cell techniques that are revolutionizing the assessment of single nucleotide variation (SNV) at the highest cellular resolution. However, state-of-the-art single-cell sequencing technologies produce data with many missing bases (MBs) and incorrect base designations that lead to false-positive (FP) and false-negative (FN) detection of somatic mutations. While computational methods are available to make biological inferences in the presence of these errors, the accuracy of the imputed MBs and corrected FPs and FNs remains unknown.

Results: Using computer simulated datasets, we assessed the robustness performance of four existing methods (OncoNEM, SCG, SCITE and SiFit) and one new method (BEAM). BEAM is a Bayesian evolution-aware method that improves the quality of single-cell sequences by using the intrinsic evolutionary information in the single-cell data in a molecular phylogenetic framework. Overall, BEAM and SCITE performed the best. Most of the methods imputed MBs with high accuracy, but effective detection and correction of FPs and FNs is a challenge, especially for small datasets. Analysis of an empirical dataset shows that computational methods can improve both the quality of tumor single-cell sequences and their utility for biological inference. In conclusion, tumor cells descend from pre-existing cells, which creates evolutionary continuity in single-cell sequencing datasets. This information enables BEAM and other methods to correctly impute missing data and incorrect base assignments, but correction of FPs and FNs remains challenging when the number of SNVs sampled is small relative to the number of cells sequenced.

Availability and implementation: BEAM is available on the web at https://github.com/SayakaMiura/BEAM.

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1 Introduction


Many studies have performed single-cell sequencing on tumors to identify clones and their evolutionary relationships (Eirew\textit{ et al.}, 2015; Gawad\textit{ et al.}, 2014; Hou\textit{ et al.}, 2012; Jan\textit{ et al.}, 2012; Li\textit{ et al.}, 2012; Melchor\textit{ et al.}, 2014; Navin, 2015; Potter\textit{ et al.}, 2013; Xu\textit{ et al.}, 2012; Yu\textit{ et al.}, 2014). Thus, single-cell sequencing will be instrumental in revealing the genetic changes that occur during cancer progression, which is a prerequisite for clone identification and the inference of evolutionary relationships among cells and relative timing of mutation events. But, the utility of current single-cell sequencing technologies is limited by many technical issues (Gawad\textit{ et al.}, 2016; Navin, 2014, 2015; Ning\textit{ et al.}, 2014; Wang and Navin, 2015). For example, the low physical coverage of some genomic regions and
positions prevents unambiguous assignment of a nucleotide base to those positions, known as ‘missing bases’ (MBs). Allelic dropout (ADO) events cause false-negatives (FNs) when mutant alleles are present but not amplified. Infidelity of amplification can cause false-positives (FPs) when errors during initial amplification are inherited to subsequent molecules and a ‘mutation’ is identified that was not present in the sampled cell. Sometimes a single-cell cannot be completely separated from other cells, which results in the sequencing of multiple cells together. FP rates ($3 \times 10^{-5}$–$7 \times 10^{-5}$ per homozygous wild-type positions) and ADO rates (0.2–0.4 per heterozygous site) can exceed the rate of occurrence of true mutations (Ross and Markowetz 2016). MBs also occur at frequencies as high as 58% (Hou et al., 2012) in single-cell data sequences (Gawad et al., 2016). All of these problems result in inaccurate single-cell sequences even when high sequencing coverage has been achieved.

Many new methods have been developed to compensate for these issues and allow reliable inference from single-cell sequence datasets. For example, OncoNEM (Ross and Markowetz, 2016) and BitPhylogeny (Yuan et al., 2015) identify clones and their evolutionary relationship (i.e., clone phylogeny). SiFit infers cell phylogeny with the consideration of sequencing errors (Zafar et al., 2017). SCITE (Jahn et al., 2016) and Kim and Simon (2014) methods are designed to infer the order of mutations over time from the single-cell sequences. SCG is designed to deal with the issue of multi-cell sequencing when inferring clone sequences (Roth et al., 2016).

These methods produce corrected single-cell sequences, but they do not report their performance in imputing MBs correctly and reducing FPs and FNs. The primary focus of these current methods has been to improve the quality of biological inferences from error-containing single-cell sequencing data. Consequently, the absolute and relative performance of current methods for reducing the error present in single-cell sequences is not known. Significant improvement in the quality of single-cell sequences will enable use of a large number of sophisticated methods in molecular phylogenetics (Nei and Kumar, 2000) for inferring the evolutionary history of clones, reconstructing ancestral clones, identifying early and late occurring driver mutations, and characterizing inter- and intra-tumor heterogeneity. These standard approaches cannot currently be used for tumor single-cell data, because they are not robust to the presence of high levels of sequence error (Zafar et al., 2017). For example, a widely-used maximum likelihood method (Stamatakis, 2014) produces a cell phylogeny (Fig. 1b) from simulated single-cell sequence data (with MBs, FPs and FNs) that is clearly very different from the true tree (Fig. 1a). In addition to various inconsistencies in the evolutionary relationships, the branch lengths leading to the tips of the phylogeny are extensively overestimated, because all the cells of a clone (same color) are actually identical (Fig. 1a). Consequently, the inferred cell phylogeny shows much greater evolutionary depth, resulting in inflated estimates of tumor heterogeneity and incorrect mapping of mutations. Therefore, single-cell sequences require correction before use in downstream biological analysis.

In this article, we present the performance of four existing methods (OncoNEM, SCG, SCITE and SiFit) in correctly imputing MBs and reducing the numbers of FPs and FNs. We excluded methods that did not produce single-cell sequences, e.g., BitPhylogeny (Yuan et al., 2015) and the method of Kim and Simon (2014). In addition, we propose and test a new method, Bayesian Evolution-Aware Method (BEAM), which employs molecular phylogenetics and a Bayesian prediction framework to improve the quality of single-cell sequences (see Materials and methods). Our testing focused on computer simulated datasets, as knowledge of the true single-cell sequences enables direct assessment of the performance of computational methods (Ross and Markowetz, 2016; Roth et al., 2016). We also analyzed one empirical dataset (Li et al., 2012) to gauge the utility of computational approaches in a real-world scenario and the concordance of the inferences produced.

In the following, we present information on the simulated data used in our evaluation of methods, followed by a description of the BEAM approach and the assumptions, parameters, and accuracy measures used. We then present results from our analyses of simulated and empirical data discuss the patterns observed.

2 Materials and methods

2.1 Generation of datasets by computer simulations

Roth et al. datasets ($R_{1000 \times 50}$ and $R_{100 \times 50}$ datasets): We used the simulator and parameter settings described by Roth et al. (2016) to produce 240 datasets. This simulator first generates a clone phylogeny and then the clone genotypes. A new model phylogeny is
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2.2 Accuracy measurements

We recorded the numbers of missing bases (MBs), false-positives (FPs) and false-negatives (FNs) in the simulated single-cell sequence datasets. The total number of correct positions (with no MB, FP, or FN) were aggregated and divided by the product of the number of cells and the number of SNVs. This quantity is referred to as the initial sequence quality ($Q_0$), which is the same as the mean Hamming distance between the true and the inferred single-cell sequence. After the sequence data was subjected to computational analysis by BEAM, OncoNEM, SCG, SCITE and SiFit, the sequence quality was reassessed ($Q_{BEAM}$, $Q_{OncoNEM}$, $Q_{SCG}$, $Q_{SCITE}$ and $Q_{SiFit}$, respectively).

While the positions containing MBs, FPs and FNs are known for simulated data, no such information exists in the analysis of empirical data and the computational methods must be applied to all the positions. Therefore, we also compared the total numbers of MBs, FPs and FNs before and after the application of a computational method to a dataset. As the number of MBs in the simulated data was larger than FPs or FNs, and thus the accuracy of MB corrections may dominate this measure.

2.3 New method evaluated (BEAM)

The new Bayesian evolution-aware method (BEAM) uses classical molecular evolutionary phylogenetics to impute missing data and detect base assignment errors in the single-cell sequencing data. It is based on the premise that significant evolutionary information is present in the initial cell sequences regardless of base assignment errors. For example, cells from the same clone show a strong tendency to occur in close proximity in the initial cell phylogeny (Fig. 1b), as seen by the location of cells marked by the same color in the true tree (Fig. 1a), despite the presence of a large number of MBs, FPs and FNs in the simulated sequence data. BEAM uses this intrinsic evolutionary information and computes a Bayesian posterior probability (PP) of observing all possible alleles at each SNV position in each single-cell sequence, as described below.

For brevity, we explain BEAM using an example dataset that was generated using the cell phylogeny in Figure 2. It consists of 20 single-cells from eight distinct clones and 200 SNVs. The simulated sequence dataset contained 800 MBs, 429 FPs and 106 FNs (Fig. 3a). For this data, we first infer a cell phylogeny from the observed single-cell sequences by using a maximum likelihood method specifically suited for phylogenetic analysis of SNV data (Stamatakis 2014) (Fig. 3a). This approach does not require the infinite sites assumption, i.e., mutations are allowed to be lost and they may occur at the same genomic position in different cells, which is different from the principle applied in OncoNEM, SiFit and SCITE (Jahn et al., 2016; Ross and Markowetz, 2016; Zafar et al., 2017).

In this example, cells from the same clones (the same color) generally cluster together, but identical cells of a clone can show extensive observed sequence divergence (e.g. brown cells in Figs 2 and 3a). Given this initial cell phylogeny and the initial cell sequences, we estimate PP of each possible base assignment at each position in a cell sequence following equation 5 in Liu et al. (2016). In brief, consider a set of four sequences and their evolutionary tree (Fig. 4). In this tree, $x_1$ to $x_5$ represent the nucleotides at a given position in the tumor cell sequence; $x_6$ is the wild-type base from the normal cell sequence. To estimate PP of a base assignment at $x_1$, nucleotides at the tip nodes are represented by the vector with the constraint that $x_1$ is unknown, that is $x=[?,x_2,x_3,x_4]$. Also, let $y=(y_1,y_2)$ represent the vector of nucleotides $y_1$ and $y_2$ at the two ancestral

generated for every dataset (e.g. Fig. 1a for 1000 cells and 10 distinct clones). To generate a clone phylogeny, new clones are created by accumulating mutations (a mutation rate set to 0.1 per site) until all SNV loci (50) are created. The simulator uses an infinite sites assumption, so no mutations override each other, and introduces loss of heterozygosity at a rate of 0.2 per site in which heterozygous mutants are changed into homozygous mutants without allowing any loss of mutations. A clone genotype is assigned to each cell by sampling from a categorical distribution (clonal prevalence), which is generated from a symmetric Dirichlet distribution with the parameter value of 1.

Datasets with 100 and 1000 cells were produced by Roth et al. (2016) in which doublet single-cell sequencing was simulated by sampling two clone genotypes at different rates: 5%, 10, 20 and 40% of the cells. The simulator generated allelic count data with ADO by sampling from the empirical distribution of SNV frequencies specified in Roth et al. (2016). The depth of coverage at each locus was chosen from a Poisson distribution with a mean of 1000 reads. The number of variant reads was sampled from a Binomial distribution with the parameter selected from an empirical distribution and the depth of coverage sampled from a Poisson distribution. To determine if an allele was present or absent, the Binomial exact test was performed and a p-value threshold of 10⁻⁴ was used. In the resulting data, 3–51% of the observed mutant alleles were FPs and 2–20% of the observed homozygous wild-type alleles were FNs. We randomly assigned a ‘missing’ value (MB) to 20% of the bases.

Ross and Markowetz datasets (M10 × 50 – M50 × 300 datasets): This collection of 690 datasets was generated using the Ross and Markowetz (2016) simulator and parameter settings. In their approach, clone phylogenies were first generated by iteratively adding a branch with a node to an existing node that was randomly chosen from a growing phylogeny (1, 5, 10 and 20 clones). Unobserved clones were then introduced by removing clones that had at least two descendant clones (0, 1, 2, 3 and 4 unobserved clones). A new clone phylogeny was generated for each dataset, and each cell is assigned to a clone with a probability corresponding to its size (10, 20, 30 and 50 cells). Figure 2 shows an example phylogeny of 20 cells used to simulate clone evolution for generating M datasets (M10 × 50 – M50 × 300 datasets). Along the clone phylogeny, true clone genotypes were generated by assigning mutations with a uniform probability (50, 100, 200 and 300 SNVs). Observed genotypes were derived from true genotypes by introducing MBs (10, 20, 30 and 40% of SNVs), FPs (10⁻⁴, 5, 10, 20 and 30% of mutant alleles) and FNs (5, 10, 20 and 30% of wild-type alleles).
An overview of the BEAM approach. (a) The initial cell phylogeny, along with branch lengths, derived from single-cell sequence data simulated using the model tree in Figure 2. The Ross and Markowitz (2016) simulator and parameter settings were used. The number of MBs, FPs and FNs in the initial sequence data are shown in the box. (b) Cell phylogeny produced from the data in which MBs were imputed and FPs and FNs were corrected. The remaining MBs, FPs and FNs are shown. (c) Improved cell phylogeny after recomputing PPs by using the phylogeny in b. (d) The final cell phylogeny produced by BEAM along with the remaining MBs, FPs and FNs. The topology and branch lengths are very similar to those in the model tree shown in Figure 2.

Figure 3. A simple tree used in explaining PP computation. $x_i$ is a base in the single-cell sequence, $y_i$ is a base in the ancestral sequence and $b_i$ refers to a branch length nodes in this tree. Then, the Bayesian posterior probability for a set of nucleotides $(x_1, y_1)$ can be computed as follows (Liu et al., 2016):

$$f(x_1, y_1|x; b) = f(x_1) \times f(y_1|x; b)/f(x; b)$$

(1)

Here, $b$ is the vector of branch lengths $b_1, ..., b_5$. The prior, $f(x_1)$, is the product of the probability of change from its ancestral base $y_1$ and the frequency of the base $y_1$ in the sequence data. The likelihood, $f(x_1|x_1, y_1; b)$, is computed by assuming $x_1$ is unknown, while the denominator is computed for all possible combinations of $x_1, y_1$ and $y_2$. Then, the PP of a nucleotide at $x_1$ is the sum of probabilities of $f(x_1, y|x; b)$ for all possible combinations of $y_1$ and $y_2$. The PP of base A at $x_1$, for example, is given by:

$$PP(x_1 = A) = \sum_{y_1 = A} f(x_1) \times f(y_1|x_1; b)/f(x; b)$$

(2)

We compute PP for each possible base assignment at each position ($i$) in the single-cell sequence of interest, including those for which a base call is available because they could be FPs or FNs. If there are $m$ single-cell sequences and $n$ SNV positions in the dataset, $m \times n$ computations are performed. We assigned the base with PP > 0.7 to the position of interest. When all of the bases show PP < 0.7 at a position, we assign the base or missing value originally observed at that position. To select this cut-off, we compared Q5 between various PP cut-offs (0.6–0.9) on a simulated data and identified PP with the best performance. We found that the use of higher PP cut-offs will increase the number of MBs, whereas a lower PP cut-off will increase the number of FPs and FNs. So, the use of 0.7 as an intermediate cut-off seems to be the best choice. We used this PP cut-off value to analyze all the datasets.

In our example, BEAM improves the single cell sequences by imputing base identities at MB positions by reducing these from 800 to 93, reducing FPs from 429 to 155, and reducing FNs from 106 to 92. Because of these improvements, the inferred cell phylogeny becomes more accurate (Fig. 3b). At the same time, the erroneously long tip branches in the initial phylogeny are shortened via elimination of FPs (Fig. 3a).

In the next step, we use the new cell phylogeny (Fig. 3b) and recompute PPs by applying Equations 1 and 2. Now, MBs decrease from 93 to 14 and FPs and FNs increased slightly (155 to 162 and 92 to 109, respectively; Fig. 3c). The cell phylogeny inferred using these single-cell sequences looks similar to the true phylogeny (Fig. 2). No further iterations are needed after this step, and the last step is to annotate clones. Clone delineations can be done by using the bootstrap procedure to assess the robustness of the branching patterns (Felsenstein, 1985; Nei and Kumar, 2000), merging cells by iterative clustering of nodes along branches (Ross and Markowitz, 2016), or by using a k-medoids clustering approach on a distance...
matrix that is obtained from the latest cell phylogeny (Zafar et al., 2017). BEAM provides the user with an option to assign identical sequences to all the cells of a clone by erasing potentially spurious mutations. In our simulations, we found that it was best to assign identical clone sequences to cells that were connected with effective- zero branch lengths in the phylogeny (default: <2% of SNVs; Fig. 3d). In this case, BEAM will finalize sequences by computing PPs of each possible base assignment and comparing the average PPs between potential bases in order to assign the base with the highest average PP to all the cells from the same clone. Ultimately, BEAM produces refined single-cell sequences as well as the cell phylogeny.

2.4 Options used for analyzing data
We used default or recommended parameters to perform SCG, SCITE, OncoNEM and BEAM analyses. SCG (Roth et al., 2016) was performed with the doublet option. When the status of mutations (presence or absence) within a predicted clone genotype had <0.95 probability, we assigned missing values to those positions. To assign a clone for each cell, SCG additionally computed the probability of having a predicted clone genotype for each cell. Thus, we assigned the predicted clone genotype with the highest probability. When none of predicted clone genotypes had a high probability (>0.01) for a cell, no predicted clone genotype was assigned to the cell. Cells lacking clone genotype assignments were removed, and datasets consisting entirely of cells lacking clone genotype assignments were removed from accuracy considerations as we considered that SCG failed to correct sequences for these datasets.

OncoNEM (Ross and Markowitz, 2016) analyses were performed using true rates of false positive and false negative base assignment errors in the input files (observed sequences). Maximum Bayes factor for which a smaller model was preferred was 10, and the model search stopped when the best scoring tree stabilized for at least 200 iterations. Mutant nucleotides were assigned when the probability of observing the mutation at a given position was >0.95 and missing bases were assigned when the probability of observing the mutant nucleotide was between 0.05 and 0.95. SCITE (Jahn et al., 2016) analyses were performed by giving true rates of false-positive and false-negative detections of mutations. The desired number of repetitions of the MCMC was 1, and the desired chain length of each MCMC repetition was 900,000. Often, mutant nucleotides were not assigned to any cells. When mutant nucleotides were not assigned to any cells, those sites were assigned with wild-type bases. When multiple possible cell phylogenies for a dataset were produced, we used the option to marginalize out the alternatives. We distinguished heterozygous and homozygous mutations for R1000 × 50 and R100 × 50 datasets. When multiple cell sequences were inferred for a single cell, we replaced inconsistent base assignments with MBs. For SiFit (Zafar et al., 2017) analyses, we also input true rates of false positive and false negative detection of mutations. The number of iterations run for each restart was 10,000. Cell genotypes were inferred by inferring the order of mutations along the cell phylogeny predicted by SiFit. LOH rate and deletion rate were set to zero.

2.5 Empirical data
We analyzed a single-cell sequencing dataset from muscle-invasive bladder tumors (Li et al., 2012), which has been analyzed previously in other articles proposing new methods (Ross and Markowitz, 2016; Zafar et al., 2017). We obtained sequenced reads in FASTQ format by using SRA toolkit (v2.8.1) (Leinonen et al., 2011). We mapped these sequenced reads to the human genome sequence (hg18 from UCSC database; https://genome.ucsc.edu/) by using the Burrows-Wheeler alignment tool (BWA v0.7.12) (Li and Durbin, 2009) with aln options. Samtools (v1.3.1) (Li et al., 2009) was used to remove reads with low mapping quality (≤40) when creating BAM files, which were sorted by chromosome coordinate. This initial data processing follows the protocol described in Zafar et al. (2016).

We then used Monovar (Zafar et al., 2016) to call mutations. We performed mpileup in Samtools with the options presented in the instructions for Monovar (i.e. minimum base quality was zero). Monovar analysis was performed with the default or recommended options, i.e. offset for prior probability of false-positive error was 0.002; offset for prior probability of allelic drop out was 0.2; threshold for variant calling was 0.05; and the number of threads used in multiprocessing was 2.

For downstream computational analysis, we selected SNVs in coding regions that were identified by Zafar et al. (2016). Nucleotides identified among the majority of normal cells were assigned as wild-types, and the other bases found in tumor cells were assigned as mutants. Our analyses did not distinguish homozygous and heterozygous mutations. Following Zafar et al. (2016), we assigned missing values to positions with coverage depth less than 6×, in addition to positions where Monovar did not predict a genotype. Lastly, we removed one cell that contained a very large number of missing bases.

3 Results
3.1 Analysis of simulated large datasets
We first present results from the analysis of 120 large datasets that consisted of 1000 cells each with 50 SNVs (R1000 × 50). The initial sequence quality (Q95) of these datasets ranged from 65 to 75% (Fig. 5a), which was caused by 20% MBs, 7–51% FPs and 2–16% FNs. Four of the five methods were able to handle these large datasets and produced refined single-cell sequences of much higher quality than the input (Fig. 5a). The average QBEAM was 89% and varied from 80–95%. The performance was the best when the input sequences contained the fewest errors and the worst when the input sequence error was the largest. SCITE performed similarly well (QSCITE = 90%) and showed very similar trends describing the relationship between input and output quality. While SCG also worked well (QSCG = 89%), it showed much greater variability in performance. SiFit showed the smallest improvements (QSiFit = 80%). Overall, the sequence quality improved for all datasets after a computational method was used.

We found that the correct imputation of MBs was the primary reason for the improvements observed for R1000 × 50 datasets. The fraction of MBs found in the single-cell sequences decreased, on average, by 100, 100 and 96% after the application of BEAM, SCG and SCITE, respectively, and most of the missing data were correctly imputed (88, 88 and 87%, respectively) (Fig. 5b). Although SiFit imputed all of the MBs without producing new MBs, SiFit correctly imputed a much smaller fraction of MBs than the other methods (70% of MBs were correctly imputed). Interestingly, however, no method showed a significant ability to reduce the total number FPs and FNs in this data (Fig. 5c), as the numbers of FPs and FNs in the output were greater than in the input (output/input ratio >1.0). Of all the methods, SiFit showed the worst average ratio (2.18), which can happen because the correction procedure has to be applied to all the bases in the input sequence data as the positions with FP and FN
are not known in the real world data analysis. This causes the creation of many new false positives and false negatives.

Next, we tested the accuracy of computational methods for datasets in which the number of sequenced cells was reduced to 100 (R100 × 50). Again, all methods successfully improved single-cell sequences with very similar output sequence quality (Fig. 6a). Outcomes were similar to those observed for R1000 × 50 datasets, except that SiFit performed much better and OncoNEM produced results. As with the larger dataset (R1000 × 50), the correct imputation of MBs was the primary improvement observed (Fig. 6b), and all of the methods produced sequences in which the numbers of FPs and FNs were similar to or much larger than the error in the input sequence data (Fig. 6c).

In both R1000 × 50 and R100 × 50 datasets, we observed that the identification of FPs and/or FNs was less effective than the imputation of MBs for all the methods (Figs 5c and 6c). We hypothesized that, unlike the imputation of MBs, detection and correction of FPs and FNs was very sensitive to the available cell relationship information that can be gleaned from the initial error-prone single-cell sequencing data. We tested this hypothesis by increasing the number of SNVs used to 500 (R1000 × 500), while keeping the number of errors per SNV the same as in R1000 × 50 dataset. We applied BEAM and SCG to the new collection of datasets and found that the output/input ratio of FPs and FNs for BEAM became much less than 1 (Fig. 7a). That is, BEAM was able to produce sequences with ∼40% fewer FPs and FNs. This improvement over R1000 × 50 datasets is explained by the fact that the initial cell relationships

derived using the input data in BEAM are more accurate when the number of SNVs analyzed is large (compare phylogenies in Figs 7c and 1b). This improvement enables the Bayesian analysis to generate better predictions, because the phylogenetic prior is closer to the truth.

The performance of SCG did not improve, and still created more FP and FN errors in the output (Fig. 7b). This may be because SCG first clusters cells to identify clones and then assigns clone genotypes. This procedure does not appear to benefit from increased evolutionary information in the dataset. Additional analyses support this reasoning, as BEAM, OncoNEM, SCITE and SiFit were able to detect and correct many FPs and FNs when the number of SNVs sampled was large in relation to the number of cells sampled.

### 3.2 Analysis of smaller datasets

Next, we analyzed 690 datasets that were generated by Ross and Markowetz (2016) (M datasets). All contained fewer cells than the R1000 datasets (largest) and R100 datasets (medium-sized), but the M datasets contained the same or larger numbers of SNVs (50, 100, 200 and 300 SNVs) compared to the R datasets. All five methods produced results for all datasets, in which the initial quality of cell sequences was between 65 and 75%. BEAM, SCITE and SiFit increased the quality of the sequences to an average 92% (Fig. 8a), but OncoNEM (82%) was less accurate and SCG performed poorly (65%). OncoNEM and SCG did not impute missing data as accurately as BEAM, SCITE and SiFit (Fig. 8b).
All methods, except SCG, decreased the numbers of FPs and FNs in the analysis of M datasets (Fig. 9), because the ratio of the number of SNVs to the number of cells was greater than that for R datasets. In fact, increasing the number of SNVs provides a proportional increase in the performance of BEAM, OncoNEM and SiFit. Patterns observed for M datasets (Fig. 9) confirm the results for R datasets with 500 SNVs (Fig. 7): BEAM becomes more accurate with larger numbers of SNVs and SCG’s performance does not improve. OncoNEM performed the best in correcting FPs and FNs in M datasets, but achieves this at the expense of producing many MBs (Fig. 8).

As expected, the quality of the inferred sequences became higher as the number of SNVs increased (Fig. 10a). The quality of the inferred sequences was higher when the number of cells was high (Fig. 10b), because all methods utilize similar cell sequences (or clusters of SNVs) to impute MBs and correct FPs and FNs. For example, the PP calculation is affected by the base assignments of neighboring cells, which will become more accurate when a base assignment is supported by larger number of cells from the same clone. Overall, BEAM, SCITE and SiFit provided the most robust results when the number of cells was small, and OncoNEM and SCG were greatly impacted if multiple cells per clone were not sampled. Lastly, the quality of the output sequences was a direct function of the fraction of MBs (Fig. 10c), FPs (Fig. 10d) and FNs (Fig. 10e).

Before proceeding with an analysis of empirical datasets, it is important to note that we did not introduce any loss of mutant alleles in our simulations, e.g. loss of heterozygosity (LOH) and the loss of genomic segments. Such mutations will negatively impact the performance of most methods, as the evolutionary relatedness of sequences will be disturbed by such losses. Therefore, such positions should be detected and removed before applying these computational methods. Also, we found that a high rate of doublet sequencing did not adversely impact the performance of any of the computational methods (Fig. 10f). This result is consistent with the finding in Jahn et al. (2016), who reported that doublet sequencing did not lead to lower accuracy in biological inference, which was a major motivation for the development of SCG (Roth et al., 2016).

3.3 Analysis of an empirical dataset

We applied all five methods to a previously published dataset of a muscle-invasive bladder tumor (Li et al., 2012). This dataset contained 55 cells with 84 SNVs in protein coding regions. The cell
Fig. 10. The relationship of the output sequence quality with the various simulation parameters. (a) Number of SNVs in the data, (b) number of cells sequenced and fractions of (c) missing bases, MBs, (d) false positives, FPs and (e) false negatives, FNs in M datasets. (f) Effect of increasing doublet sequencing rates on the quality of the output sequencing for R datasets.

phylogeny before and after the application of BEAM is shown in Figure 11b. Following the application of BEAM, the cell phylogeny showed high diversity among cells, with no clear demarcation of clones; this is in contrast to the initial cell phylogeny (Fig. 11a) based on BEAM’s clone predictions in Figure 11b. The initial cell phylogeny (Fig. 11a) was transformed into the final cell phylogeny (Fig. 11b) by BEAM, because the final single-cell sequences contained only 6 positions with missing values. This is a major improvement, as the original dataset contained 666 positions with missing values. In imputing MBs, BEAM assigned a mutant base to 337 positions (99% of MBs), BEAM corrected wrong mutant and wild-type base assignments. In these predictions, 105 mutation calls were found to be false positives, 65% of which were also detected by at least two other methods. That is, they were consensus assignments (≥3 out of 5 methods). Also, 183 wild-types were detected to be false negatives, i.e. they should have been mutant assignments. 88% of these were consensus assignments, because at least two other methods suggested the same base assignment. Therefore, we expect that the use of these computational methods will enable better biological conclusions.

4 Discussion

We have reported that computational methods are generally capable of imputing missing bases with high accuracy and, thus, can improve the quality of the tumor single cell sequences. In particular, BEAM, SCG and SCITE produced well in imputing missing bases for datasets with a large number of cells. Our results also confirm Ross and Markowitz (2016)”s conclusion regarding the accuracy of SCG for datasets representing a large number of cells but containing few SNVs. However, we have newly reported that the gain in accuracy is due to correct imputation of missing data, and that SCG does not perform well in correcting FPs and FNs. In fact, all methods require a large number of SNVs to detect and correct FPs and FNs. And, as mentioned earlier, no other methods provided information on their performance in accurately imputing MBs and correcting FPs and FNs. So, our results provide the knowledge of potential errors that each method may produce in actual empirical data analyses, which will be a guide to analyze the inferences of these methods for practitioners.

We have shown that, when the number of SNVs sampled is large, many methods also show good performance in detecting and correcting false positive and false negative mutation assignments. In our analyses, BEAM, SCITE and SCG methods performed very well for datasets containing a small number of cells, both for small and large number of SNVs. These three methods employ molecular phylogenetics, but BEAM is based exclusively on classical molecular phylogenetic methods and applies a Bayesian framework to impute MBs and correct FPs and FNs by using the initial cell sequence phylogeny as a prior. In contrast, SCITE and SCG employ a model that account for sequencing error rates in the process of inferring the evolutionary tree of cells, which also results in improved cell sequences (Jahn et al., 2016; Zafar et al., 2017). SCG produces refined cell sequences by inferring the order of mutations along the inferred maximum likelihood cell phylogeny with given error rates, whereas SCITE uses a Bayesian method to search for a cell phylogeny that intrinsically maps mutations along branches in the phylogeny. On the other hand, BEAM does not determine the order of mutations and considers all possible ancestral base assignments to refine cell sequences.

As BEAM also uses a Bayesian method, we consider BEAM and SCITE to be in the same class of methods, with the difference that more than 93% of BEAM’s assignments for MBs were shared with at least one other computational tool. Therefore, extensive consensus exists among methods. We also examined whether BEAM’s base assignments for MBs were supported by the read count data. As mentioned in an earlier section, Monovar does not assign a base when there are only a few reads. So, we tested whether the wild type base assignments by BEAM were supported by higher read counts than other bases at the examined positions. This was indeed the case, the read counts of wild type alleles assignments averaged 7 times higher. Assignment of mutant alleles to MBs were also supported by 22% larger read counts than the wild type alleles. Therefore, read count data generally supports the assignments made by BEAM.

In addition to imputing almost all MBs in the observed sequences (99% of MBs), BEAM corrected wrong mutant and wild-type base assignments. In these predictions, 105 mutation calls were found to be false positives, 65% of which were also detected by at least two other methods. That is, they were consensus assignments (≥3 out of 5 methods). Also, 183 wild-types were detected to be false negatives, i.e. they should have been mutant assignments. 88% of these were consensus assignments, because at least two other methods suggested the same base assignment. Therefore, we expect that the use of these computational methods will enable better biological conclusions.
SCITE needs to model sequencing error rates, but BEAM does not require this modelling. Also, BEAM decouples the inference of cell phylogenies from the corrections of observed single cell sequences, as compared to joint inference by SCITE. Interestingly, both of them show comparable results, which are among the best. That is, BEAM obviates the need to apply the same sequencing error rate model throughout the tree, which may be preferable because mutational patterns change in cells based on their evolved state (Frank and Nowak, 2004), potentially resulting in heterogeneity of error rate models among clones. An advantage of BEAM over SCITE may be that BEAM performs slightly better when the number of SNVs is large (Fig. 9), because the initial cell phylogeny derived using observed single-cell sequences is expected to be more accurate for datasets with a large number of SNVs.

Our results show that methods that incorporate the cell phylogeny are more powerful than others, especially when the number of cells per clone is small and the number of SNVs is large. This is because BEAM, SiFit and SCITE perform better than OncoNEM, which aims to infer clone phylogeny, and SCG, which employs a mixture model that identifies groups of cells with shared clone genotypes. Because tumor cells descend from pre-existing cells, there is evolutionary continuity in cell sequencing datasets, which enables computational methods to correctly impute missing data and make correct base assignments. We find that molecular evolutionary methods that have been successfully applied for species and strain phylogenetics for decades serve as a strong foundation for phylogenetic approaches with greater power to impute missing data and refine cell sequences for small datasets.

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References


