Inferring Cancer Progression from Single-Cell Sequencing while Allowing Mutation Losses

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Abstract

**Motivation:** In recent years, the well-known Infinite Sites Assumption (ISA) has been a fundamental feature of computational methods devised for reconstructing tumor phylogenies and inferring cancer progressions. However, recent studies leveraging Single-Cell Sequencing (SCS) techniques have shown evidence of the widespread recurrence and, especially, loss of mutations in several tumor samples. While there exist established computational methods that infer phylogenies with mutation losses, there remain some advancements to be made.

**Results:** We present SASC (Simulated Annealing Single-Cell inference); a new and robust approach based on simulated annealing for the inference of cancer progression from SCS data sets. In particular, we introduce an extension of the model of evolution where mutations are only accumulated, by allowing also a limited amount of mutation loss in the evolutionary history of the tumor: the Dollo-\(k\) model. We demonstrate that SASC achieves high levels of accuracy when tested on both simulated and real data sets and in comparison with some other available methods.

**Availability:** The Simulated Annealing Single-Cell inference (SASC) tool is open source and available at https://github.com/sciccolella/sasc.

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**Supplementary information:** Supplementary data are available at Bioinformatics online.

1 Introduction

Recent developments in targeted therapies for cancer treatment rely on the accurate inference of the clonal evolution and progression of the disease. As discussed in several recent studies (Morrisay and Garzia, 2016; Wang et al., 2016), understanding the order of accumulation and the prevalence of somatic mutations during cancer progression can help better devise these treatment strategies.

Most of the available techniques for inferring cancer progression rely on data from next-generation bulk sequencing experiments, where only a proportion of observable mutations from a large amount of cells is obtained, without the distinction of the cells that carry them. In recent years, many computational approaches have been developed for the analysis of bulk sequencing data with the purpose of inferring tumoral subclonal decomposition and reconstructing tumor phylogenies (evolutionary trees) (Strino et al., 2013; Jiao et al., 2014; Hajirasouliha et al., 2014; Yuan et al., 2015; Popic et al., 2015; Malikic et al., 2015; El-Kebir et al., 2016; Marraus et al., ...
Single-Cell Sequencing (SCS) technologies promise to deliver the best resolution for understanding the underlying causes of cancer progression. However, it is still difficult and expensive to perform SCS experiments with a high degree of confidence or robustness. The techniques currently available are producing datasets which contain a sizeable amount of noise in the form of false negatives from allelic dropout, and missing values due to low coverage. Another issue that these technologies suffer from is the presence of doublet cell captures. However, such issues are slowly fading away and the state-of-the-art in preprocessing steps for removing such artifacts is quite advanced (DePasquale et al., 2019). Hence, we believe that more immediate issues, such as the lack of accuracy reflected in the high dropout and false negative rates inherent to the technology, call for methods that are able to infer cancer progression from this data produced by current SCS techniques.

Various methods have been recently developed for this purpose (Jalan et al., 2016; Ross and Markowitz, 2016; Zafar et al., 2017, 2019), some of them introducing a hybrid approach of combining both SCS and VAF (bulk sequencing) data (Ramazzotti et al., 2017; Malikic et al., 2018; Salehi et al., 2017). Most of these methods, however, rely on the Infinite Sites Assumption (ISA), which essentially states that each mutation is acquired at most once in the phylogeny and is never lost. One reason being that such a simplifying assumption leads to a computationally tractable model of evolution, namely, the problem of finding a perfect phylogeny (Gusfield, 1991). This model is safe to use in settings such as the evolution of natural populations, and tends to be the norm more than the exception in this setting (Kimura, 1969). Cancer progression, however, is a fairly extreme situation, where the evolution is very fast, under attack from the immune system, and with a high mutation rate. As a result, studies of SCS data are beginning to reveal phenomena that cannot always be explained with a perfect phylogeny (Kuipers et al., 2017; Brown et al., 2017). Some papers (Kuipers et al., 2017) reveal widespread recurrence and loss of mutations, while large deletions on several branches of a tree can span a shared locus (Brown et al., 2017), thus a given mutation may be deleted independently multiple times.

In this work we propose a novel and more general model to explain the above phenomena, which is not unnecessarily held back by strict adherence to the ISA. Some recent methods are beginning to appear, which have the same objective in mind, such as TRaIT (Ramazzotti et al., 2017), SiFiT (Zafar et al., 2017) and SPhYR (El-Kebir, 2018); in detail, TRaIT accounts for violations of the ISA by accommodating for convergent evolution, SiFiT accounts for both mutation recurrence and loss without specifying a particular model of evolution, and, on the other hand, SPhYR (independently from our paper) utilizes the same phylogeny model used in this work, thus allowing deletions of mutations. In our approach, we use the Dollo model (Farris, 1977; Rogozin et al., 2006), one of the models that is more general than the perfect phylogeny model, to allow the loss of point mutations. In particular, while the Dollo model still constrains that a mutation can only be acquired at most once, it allows a number of independent losses of the mutation. Once we depart from an ideal, error-free, perfect phylogeny model (Gusfield, 1991), we lose its convenient computational tractability: in fact allowing errors or missing data results in an NP-hard problem. Adopting the more general Dollo model is only going to exacerbate the problem. However, if we restrict the number of losses of any mutation to 1 or 2 (rather than strictly 0), the resulting solution space is still small enough to explore a sizable portion of it in a reasonable amount of time, in practice. Moreover, from a biological point of view, one would not expect a mutation to be lost more than a few times, since it is not likely that mutations are widely lost (Kuipers et al., 2017; Brown et al., 2017). Furthermore, all the currently available methods assume that the false negative rate is the same for all mutations. While this is suitable for samples coming from DNA (i.e., scDNA-seq) data, the false negative rate of mutations in samples coming from RNA (i.e., scRNA-seq) data can vary a because of differing levels of gene expression. Since our approach is suitable for both types of data, that is, a suitable parameter setting can be found for modeling the progression of cancer from samples coming from either DNA or RNA data, to accommodate the latter, our approach also allows a different false negative rate for each mutation: it is one of the first methods with this feature. In fact, to the best of our knowledge, the only other paper with a similar feature has appeared very recently (Wu, 2019); in that paper, different false negative and false positive rates are allowed for each mutation and for each cell. At the same time, mutation losses are not allowed. SciPhi (Singer et al., 2018) also allows different rates, but it is essentially a phylogeny-aware mutation caller, not a tool designed to infer tumor phylogenies.

Here we introduce the Simulated Annealing Single-Cell Inference (SANS) tool, a maximum likelihood phylogeny search framework that allows deletion of mutations, by incorporating the Dollo parsimony model (Farris, 1977; Rogozin et al., 2006). We show that our approach is competitive with the state-of-the-art tools for inferring cancer progression from SCS data, while being the only tool to correctly identify important driver mutations in some real datasets, as verified by the manually curated progression scenarios for these data.
on different factors. In particular, for scRNA-seq data, the varying expression levels of different genes can easily lead to different false negative rates for each mutation, since a highly expressed gene will have significantly higher coverage than an under-expressed gene, resulting in a more accurate SNV call for that particular gene. On the other hand, a gene which is less expressed is likely to have a lower coverage, leading to a less accurate presence/absence estimation in the cells. We assume that these errors occur independently across all the (known) entries of I. Namely, if \( E_{ij} \) denotes the final \( n \times m \) output matrix, i.e., the binary matrix without errors and noise estimated by the algorithm, then \( \alpha_j \) denotes the false negative rate of mutation \( j \); and \( \beta \) denotes the false positive rate, similarly to (Jahn et al., 2016; Ross and Markowetz, 2016; Ziaie et al., 2017; El-Kebir, 2018). Hence, for each entry of \( E_{ij} \) the following holds:

\[
P(I_{ij} = 0|E_{ij} = 0) = 1 - \beta \quad P(I_{ij} = 0|E_{ij} = 1) = \alpha_j \quad P(I_{ij} = 1|E_{ij} = 0) = \beta \quad P(I_{ij} = 1|E_{ij} = 1) = 1 - \alpha_j.
\]

We aim to find a matrix which maximizes the likelihood of the observed matrix I (Jahn et al., 2016) under the probabilities of false positives/negative and missing entries. Different from previous works, our model also accounts for losses of mutations, thus we define the prior probability \( P(I_{ij}) = \gamma_j \) — independent from the previous ones — of losing mutation \( j \) and the set of variables \( c_j \) for \( j = 1, \ldots, m \) that denotes the total number of losses for mutation \( j \) in the evolutionary history. In practice, we expect that a researcher might able to determine that some mutations \( j \) are very unlikely to be lost, therefore setting \( \gamma_j = 0 \).

However, we are interested in the reconstruction of the evolutionary history of the input cells, thus the resulting matrix E should contain clones (groups of cells with the same mutations) that can be explained by an evolutionary process of the mutations. This restriction motivates the introduction of the concept of phylogenetic tree, or simply phylogeny.

A (cancer) phylogeny T on a set \( C \) of \( m \) mutations and \( n \) cells (affected by these mutations) is defined as a rooted tree whose internal nodes are labeled by the mutations of \( C \), while the leaves are labeled by the cells (see Fig. 1A). Notice that the labeling must satisfy some restrictions depending on the evolutionary model that we consider. For example, in a perfect phylogeny, no two nodes have the same label. This is an alternative, but essentially equivalent, definition of classical character-based phylogeny, where the tree T is defined on a set of characters and where leaves have no label and represent different species.

The state of a node \( z \) is defined as the set of mutations that have been acquired but not lost in the path from the root to \( z \). The state of each leaf \( l \) of T is naturally represented by a binary vector of length \( m \), called genotype profile, that we denote \( D(T,l) \), where \( D(T,l)_i = 1 \) if and only if the leaf \( l \) has the mutation \( j \) and 0 otherwise (see Fig. 1B).

We say that the tree \( T \) encodes a matrix E if there exists a mapping \( \sigma \) of the rows (cells) of \( E \) to the leaves of \( T \) such that \( E_{ij} = D(T, \sigma_i)_{j} \) for each row \( i \) of \( E \), where \( \sigma_i \) denotes the image of row \( i \) through the mapping \( \sigma \). Informally, \( \sigma_i \) is the node in the phylogenetic tree corresponding to the node where the cell \( i \) is attached. Notice that the matrix \( E \) is fully characterized by the pair \( (T, \sigma) \) (see Fig. 1C). Thus, our problem can be expressed as finding the tree \( T \) that maximizes the following objective function:

\[
\max_{\sigma} \sum_{j} m \left( \gamma_j \log(1 - P(L(j))) + \sum_{i} \log(P(I_{ij}|D(T, \sigma_i))) \right)
\]

We point out that the values assigned to the unknown entries of the input matrix do not factor into the objective function, that is \( P(I_{ij} = 0|E_{ij} = 0) = P(I_{ij} = 0|E_{ij} = 1) = P(I_{ij} = 0|E_{ij} = 1) = 1 - \alpha_j \).

2.2 Introduction of the Dollo-k model

The Dollo parsimony rule allows that, in a phylogeny, any single mutation is uniquely introduced in the evolutionary history, but deletions of the mutation can occur any number of times. A restricted version of the Dollo model can be obtained by bounding the number of deletions for each mutation. We denote as Dollo-k the evolutionary model in which each mutation can be acquired exactly once and can be lost at most \( k \) times. The special cases, Dollo-0 and Dollo-1, correspond to the perfect (Gusfield, 1991) and persistent (Bonizzoni et al., 2017; Della Vedova et al., 2017; Bonizzoni et al., 2012) phylogeny models, respectively. The phylogeny reconstruction problem under a Dollo-k model is NP-complete (Goldberg et al., 1996) for any \( k > 1 \).

Since the Dollo evolutionary model allows back mutations, we introduce a new type of node label in the phylogenetic tree, to express mutational losses. For each mutation \( p \) we create \( k \) new mutations \( p^+_{i} \) for \( i \in \{1, \ldots, k\} \), representing the possible number of losses. As a result, no two different nodes have the same label. Additionally, we impose that all nodes labeled by a mutation loss \( p^- \) are descendants of the node labeled by the gain of mutation \( p \). Consequently, the vector \( D(T, \sigma) \) which expresses the genotype profile of a row \( i \) will have a 1 in mutations acquired but never lost in the path from the root to the parent \( \sigma_i \) of the leaf \( i \).
that the tree of (Fig. 1D) is a Dollo-1 phylogeny. We stress that, unlike the case of the perfect phylogeny, when deletions are introduced, we might have more than one tree that is a solution. For example, switching the labels of nodes $b^-$ and $d^-$ in Fig. 1 produces a different tree which is still a solution of the proposed input matrix when the Dollo model is considered. Moreover, the set of ancestral relationships between those two mutations is opposite in both representations. An increase of the number of cells and mutations, coupled with the noise caused by false calls and missing entries, expands the solution space of this problem, increasing the number of different cancer progression phylogenies which equally explain the same input.

Our model. The model we employ in this work is the Dollo-$k$ model, with the added restriction that there are at most $d$ total mutation losses in the entire progression. In addition to $k$, this $d$ is a user supplied parameter. Note that, with a maximum $d$ of total losses in the progression, it means that the variable $c$ is subject to (1) $c_j \leq k \cdot \gamma_j$ and (2) $\sum_j c_j \leq d$.

Only a small number of mutation losses in each tumor have been reported (Kuipers et al., 2017), therefore we expect small values of $k$ and $d$ to be used in practice. Most precisely, we believe that $k \leq 2$ and $d \leq 5$ in almost all cases. If the number of mutation is not too small, setting $d \leq 5$ essentially implies $k \leq 1$, hence making the parameter $k$ mostly irrelevant. Still, we have decided to keep it because it guarantees that some degenerate trees are never computed. We recall that our model also has the $\gamma_j$ parameters, that is, the prior probability of losing mutation $j$.

2.3 Simulated Annealing

As mentioned before, the fact that (1) we can flip entries and that (2) we want to find the maximum likelihood tree, makes the phylogeny reconstruction problem under the Dollo-$k$ model computationally hard for any $k > 0$. For this reason, in this paper we consider the Simulated Annealing (Kirkpatrick et al., 1983) (SA) approach in order to find a tree which maximizes the likelihood of an incomplete input matrix and that satisfies the Dollo-$k$ phylogeny model, where $k$ is given as input.

SA is a random search technique which explores the region of feasible solutions, searching for an optimal one. As all other meta-heuristic strategies, it is not guaranteed that SA finds the optimal value of the objective function in a finite number of steps; nevertheless, unlike other deterministic search methods which can be trapped into local optima, SA has been designed to overcome this drawback and converge to a global optimum. The basic idea of the algorithm is to perform a random search which accepts, with some probability, changes that do not necessarily improve the objective function. At each step, the probability of moving to some state with a smaller value changes according to a parameter called the temperature, which continuously decreases as the exploration evolves. In the first iterations of the algorithm execution, the temperature is very high, and it is possible (with a fairly high probability) to accept a move into a state with a lower objective value, but as temperature decreases, the probability of moving also decreases. At the end, when the temperature is sufficiently low, the algorithm becomes a local search method, hence unable to escape a local optimum.

2.3.1 Neighborhood topology

An essential element of a SA approach that we must provide how the algorithm search process can move from a given state to another. In our particular framework, we attempt to find a tree, thus we must define the neighborhood of a phylogenetic tree in the feasible region, and the algorithm moves from a tree to one of its neighbors. The choice of neighborhood is crucial in the algorithm definition since it determines how feasible solutions are explored, hence ultimately determining whether or not the algorithm converges.

In our approach, the notion of neighborhood is operational, that is, two phylogenetic trees are neighbors if one can obtained from the other via some operation from a set we will define shortly. For the sake of clarity, we introduce some notation:

- **Swap node labels**: given two nodes $u, v \in T$ such that neither is an ancestor of the other, we swap the labels rooted in $u$ and $v$ as a new child of $v$ by adding the edge $(v, u, \rho(v))$.

- **Add a deletion**: given two nodes $u, v \in T$ such that $v$ is an ancestor of $u$, we insert a node $v^-$ that represents a loss of mutation $v$. The new node is made the parent of $u$.

- **Remove a deletion**: given a node $u \in T$, labeled as a loss, we simply remove it from the tree $T$. All children of $u$ are added as children of $\rho(u)$ and the node $u$ is then deleted.

- **Subtree Prune and Reattach**: given a tree $T$ and two internal nodes $u, v \in T$ such that either $u$ is an ancestor of the other, we prune the subtree rooted in $u$ and we reattach it as a new child of $v$.

Moreover, in the SA search processes, we have that, given a tree and a valid tree operation, the probability of accepting the new solution is $\min(1, \exp(-\Delta E/T))$, where $\Delta E$ is the possible change in the likelihood function after performing the operation, and $T$ is the current temperature. The cooling process follows a geometric decay with a factor (cooling rate) $\alpha$ of $10^{-5}$, i.e., the temperature at the $i$-th iteration is equal to $T_i = (1 - \alpha) T_{i-1}$, and $T_0 = 10^4$. The SA process stops when the temperature drops below a lower bound set at $10^{-3}$.

Since mutation losses are not as frequent as mutation gains, our approach allows to set an upper bound on $d$; the total number of deletions of the resulting tree. For example, in a Dollo-$k$ model we can consider only trees where each mutation
is lost at most $k$ times, but there are at most $d$ nodes associated to mutation losses.

2.4 Visualization

Alongside the main tool, we produced a post-processing tool, called SASC-viz, that can be used to perform processing and filtering operations after the computation of the main tool. Notice that the following operations do not change the actual evolutionary history computed, but only how it is displayed.

- **Collapsing simple paths**: when this option is activated, all simple non-branching paths are collapsed, i.e. if a node has only one child, then such node is merged with its child;
- **Collapsing low-support paths**: when this option is activated, if the support of a node $i$ is lower than a specified value, then $i$ is merged with its parent $p(i)$, where the support $s_i$ of a mutation $i$ is computed on the output tree as follows: let $s(i)$ be the set of nodes in the subtree rooted at $i$, and let $C(i)$ be the number of cells assigned to the node $i$. Then the mutation support $s_i$ is:

$$s_i = \sum_{u \in s(i)} C(u) / \sum_{v \in s(i)} (C(v) - C(p(i)))$$

We used SASC-viz to produce the pictures of the real dataset analyzed. In particular, Fig. 6 and 7 are obtained by activating the collapsing simple paths option. Fig. 5 was produced with more enabled options: by collapsing simple paths and collapsing low-support paths with threshold 5%.

3 Results

Results on simulated data

We have tested our method on simulated data, where the ground truth phylogeny is known. We recall that it is possible, however, that a completely different tree achieves a better likelihood on the input data than the one obtained via simulation. This problem is essentially unavoidable, since generating a progression that is the unique solution for the corresponding SCS input matrix would require the contrived addition of artifacts to both the desired tree and the input matrix. These artifacts would likely be so artificial that the resulting instance would not satisfy even the basic assumptions on cancer progression.

Generating simulated datasets

To test the methods, we run three different experiments with increasingly sophisticated models, according to the parameter settings of Table S1. In the first experiment, we explore a model with the possibility of mutation losses, a phenomenon which has been evidenced by (Kuipers et al., 2017); the second experiment is produced with a model using only different (mutation-specific) false negative rates, as seen in the real data we use. For the third experiment, we combine the previous two to extend the simulation to the most general model in which mutations have the possibility to be lost, and each have a specific false negative rate.

The false negative error rate distribution of the real datasets is obtained by analyzing the raw data of MGH36 and MGH694 from (Tirosh et al., 2016) and comparing the mutation frequencies in the scRNA-seq data to the matching Whole Exome Sequencing (WES) from bulk RNA sequencing, to deduce the drop out frequency. We have analyzed more than 2000 mutations and observed that the distribution of the real data are consistent with a Beta distribution, which we have used as a base for the generation of our simulated data.

The values of different false negative rates are randomly chosen from a Beta distribution $B(\alpha, \beta)$ with parameters $\alpha, \beta < 1$, to better simulate the values found in real datasets; the different values of the prior probabilities of mutation losses are produced using a Triangular distribution with lower limit $a$, upper limit $b$, and mean $c$, indicated as $T(a,b,c)$. Such distribution is usually used when only the mode, upper and lower bounds are known in a population as proxy for a fair estimation of real-case scenarios. A detailed description of the method used to simulate the datasets is available in the supplementary material.

Evaluating the simulated datasets

For each of the three experiments, we measure the accuracy of SASC with two scores based on standard cancer progression measures used in various studies (Malikic et al., 2017; Jahn et al., 2016), i.e. Ancestor-Descendant and Different-Lineage accuracies; a novel parsimony-based score based on the difference between the number of flips, i.e., changes from 0 to 1 and from 1 to 0, estimated by some tool to correct the input; and the actual number of flips introduced by the simulation process to induce the noise. Lastly, we evaluate the trees using the edit distance measure of (Karpov et al., 2018). A detailed explanation of all the measures is available in the supplementary material. Note that none of the above mentioned metrics explicitly measures the ability of tools to correctly infer ISA violations.

Additionally to the aforementioned measures, we provide two accuracy measures for the estimation of false negatives: (a) an accuracy of the estimation of the average false negative rate in the simulations and (b) the value of the average, over the 50 trees, of the Mean Squared Error (MSE) over the set of estimations, for each mutation, of the mutation-specific false negative rates. Note that (b) gives an indication also of the variance of the estimation of false negative rates, which is important when these rates are heterogeneous, and far from being normally distributed — something we see in real data that we use here and that is due to varying gene amplification and expression levels.

Results of the simulation experiments

We now detail the evaluation of the last experiment, which shows the most interesting results. Due to lack of space, we refer to the Supplementary materials for a detailed discussion on the other two, less general, experiments.

We decided to compare SASC against SCITE (Jahn et al., 2016), SiFit (Zafar et al., 2017) and SPlyR (El-Kebir, 2018). While B-SCITE (Malikic et al., 2017) is a clear improvement over SCITE, it combines single-cell data with bulk sequencing data — since we do not manage the latter kind of data, a fair comparison is not feasible. For the same reason, we do not compare against TRaIT (Ramaszotti et al., 2017) and PhiSCS (Malikic et al., 2019). OncoNEM (Ross and Markowitz, 2016) was excluded because it is not able to complete the execution on datasets as large as the ones used in the simulations. Each of the tools is properly run with millions
This experiment shows the results when the datasets contain both heterogeneous false negatives and deletions based on scRNA-seq error model, thus complementing the other experiments. SASC outperforms any other tool in every considered measure (Fig. 2 and 3) and it also shows the best estimation of the false negative rates in terms of average and MSE (Fig. 4). It is particularly interesting to notice the drop in performance of SPhyR when it is forced to employ the Dollo model, since this is the only experiment with mutation losses involved. It is also very clear that SASC outperforms all the available methods when it deals with heterogeneous false negative rates and mutation losses. It also interesting to notice that SASC shows a much higher accuracy than the other two tools that allow mutational losses — SiFit and SPhyR — when such losses are present in the dataset.

Results on real cancer data

We test SASC on four different datasets, comprising both scDNA-seq and scRNA-seq sequencing data. Due to lack of space, the figures of the trees inferred by the other methods are displayed in the Supplementary Materials. Since SiFit exhibited poor performances on the simulated datasets, it is excluded in the comparison on real datasets.

Oligodendroglioma IDH-mutated tumor

We test SASC on an oligodendroglioma IDH-mutated tumor; in particular, on cancer MGH36 (Tirosh et al., 2016), consisting of 77 SNVs, distinguished from PCR false positives using matched WES, over 579 cells. Fig. 5 shows the tree computed by SASC and the distribution of the false negative rates (shown in the bottom-right corner plot). The distribution stresses the necessity of a method that considers heterogeneous false negative rates, since there are two spikes of rates (at roughly 0.1 and 0.9), i.e., it is highly bimodal, and using the average of the rates would not be an accurate representation. In this particular tumor, no deletion was expected: this is confirmed by the inferred tree.

For the dataset MGH36 from (Tirosh et al., 2016), there is no manually curated tree to compare the results of the tools, thus we report the number of false negatives and false positives inferred by the methods, this is the number of flips from 0 to 1 and from 1 to 0 respectively from the input to the output. The rationale for this score is to report a parsimony score of the algorithms; a comparison of the likelihood values will not be fair, since SASC uses a different formula than the other tools. Such score is shown in Table S5; SASC introduces the lowest number of false negatives to obtain the solution, albeit being
very close to SCITE (Fig. S11), while SPhyR (Fig. S14) infers the highest number.

**Childhood Acute Lymphoblastic Leukemia**

Furthermore, we test **SASC** on Childhood Acute Lymphoblastic Leukemia data from (Gawad et al., 2014). In particular, we focus on Patient 4 and Patient 5 of this study, given their large amount of both cells and mutations, as well as their complexity. Data on Patient 4 consists of 78 somatic Single Nucleotide Variants (SNVs) over 143 cells, while Patient 5 is affected by 104 somatic SNVs over 96 cells. The original study estimated an allelic drop-out rate of less than 30%. Since the trees in (Gawad et al., 2014), determined using expectation maximization on a multivariate Bernoulli distribution model, are manually curated and of high quality, we select them as the ground truth.

To ensure the absence of doublets, i.e. noise produced by error due to the fact that two cells are sequenced instead of a unique cell, we preprocess the input using the Single-Cell Genotyper (SCG) tool (Roth et al., 2016). SCG is a statistical model which removes all cells of the datasets that are likely to be doublets.

Fig. 6 shows the tree inferred by **SASC** for Patient 4; **SASC** correctly infers the tree structure obtained in the study, as well as the size of the subclonal population. The driver mutations are correctly detected, and mutations COL5A2, SDPR and TRHR are inferred as deletions. Furthermore, boldfaced and colored mutations indicate the correctly placed specific driver mutations for the subclone of the same color. It is interesting to notice that, in the original study, the violet subclone does not have mutations COL5A2 and TRHR; these particular mutations are in fact deleted in the clone. This solution was found assuming a Dollo-1 phylogeny model with no restriction on the total number of deletions in the cancer progression.

In Fig. 7, the inferred solution for Patient 5 of the same study is shown. As in the previous dataset, our inferred tree perfectly supports the hypotheses proposed in the original sequencing study: in fact, it correctly infers the topology of the tree, as well as the placement of driver mutations. Boldfaced mutations are the driver mutations for the tree or the subclone with the same color. This solution was found assuming a Dollo-1 phylogeny model with a restriction of 10 deletions in the cancer progression. As described in the section “Simulated Annealing”, such values for $k$ and $d$ were empirically found to give the best likelihood.

Since the original study (Gawad et al., 2014), provides manually curated trees we can compare **SASC**, SCITE and SPhyR to them.

SCITE is run using the same setting used for **SASC**, i.e. the proposed values of false positive and false negative rates. Fig. S9 shows the tree inferred by SCITE for Patient 4; the tree structure is similar to the one proposed in the paper but it presents more clones. Furthermore, we highlighted in red all driver mutations that were not correctly detected, and in blue mutations that define a subclone and should be in the same subtree. Fig. S10 shows the tree inferred by SCITE for Patient 5 of the same study; the tree topology is correctly inferred, however mutations highlighted in red are driver mutations that were not correctly detected.

SPhyR is run using the same setting used for **SASC**, i.e. the proposed values of false positive and false negative rates and assuming a Dollo-1 model. Fig. S12 shows the tree inferred by SPhyR for Patient 4; the tree structure is similar to the one proposed in the paper. The drivers and distinct subclones are also correctly placed. Fig. S13 shows the tree inferred by SPhyR for Patient 5 of the same study; the tree topology is correctly inferred, however it infers a large number of mutation losses, which is very unlikely and it is probably due to the fact that deletions are used, in this case, to correct false negatives at no cost in terms of likelihood function.

**Medulloblastoma**

Lastly we test the methods on Medulloblastoma patient ICH1031 from (Hovestadt et al., 2019) consisting of 96 mutations over 330 cells. Given the complexity and the dimensions of the trees, we display them in the Supplementary Materials. **SASC** and SPhyR computed the solution using a Dollo-2 phylogeny model.

Fig. S15 shows the tree inferred by **SASC**, which reported a total of 2 mutation losses. Both trees inferred by **SASC** and SCITE (Fig. S16) express, as expected, various mutations correlated to the TUBB gene. On the other hand SPhyR (Fig. S17) inferred a total of 56 mutations over the 96 present in the sample. Furthermore, similarly to the previous experiment, SPhyR inferred a total of 24 mutational losses, which is very unlikely for so many losses to be present in a single sample, since evidence from (Kuipers et al., 2017) suggests that this phenomenon is extremely rare. It is more likely that, also in this case, deletions are used to correct false negatives at no cost in terms of likelihood. Lastly, while **SASC** and SCITE each finished its computation in less than 2 hours, SPhyR took more than 24 hours.

**4 Conclusion**

We have presented **SASC** and we have shown that it is an accurate tool for inferring intra-tumor progression and subclonal composition from both scDNA-seq and scRNA-seq data. **SASC**
Fig. 6. The tree inferred by SASC for Patient 4 of the Childhood Lymphoblastic Leukemia data from (Gawad et al., 2014). Different clones are indicated with different colors. Red nodes indicate deletions of mutations, while boldfaced mutations are the mutations indicated as driver in the original sequencing study. Mutations in bold and colored are driver mutations for the clone with the same color. Mutations are clustered by collapsing simple linear paths. The picture was drawn using the SASC-viz post-processing tool.

Fig. 7. Tree inferred by SASC for Patient 5 of the Childhood Lymphoblastic Leukemia data from (Gawad et al., 2014). Different clones are indicated with different colors, while the red-colored nodes indicate deletions of mutations, and mutations highlighted in bold are the mutations indicated as driver in the original sequencing study. Mutations boldfaced and colored are driver mutations for the same colored clone. Mutations are clustered by collapsing simple linear paths. The picture was drawn using the SASC-viz post-processing tool.

We have tested SASC on three simulated datasets, and we have shown that SASC is able to outperform all tools when there are mutation losses, while being competitive with SCITE and SPhyR when there are no mutation losses.

We have tested SASC on three real datasets. SASC has inferred a likely phylogeny tree structure, correctly identifying the driver mutations and the decomposition of the clones. Furthermore, it has solved those large datasets in adequate runtime.

A particularly interesting example is given by the inferred tree in Fig. 7. The corresponding input dataset in this case contains more than 5000 conflicts between mutations — each conflict is a pair of mutations witnessing a violation of the Infinite Sites Assumption (ISA). With only a slight relaxation of the ISA — the Dollo-1 model — SASC is able to infer an accurate solution with a total of only 8 deletions, while perfect phylogeny methods would require a large number of changes to the entries in the input just to produce a feasible solution.

A future extension could be the inclusion of coverage information from the reads, as in Monovar (Zafar et al., 2016) and SciΦ (Singer et al., 2018), since it will also have an impact on the false negative rates. Another direction is towards even more general models, for example, allowing each mutation to appear more than once in the tree. Also in this case, special attention must be paid to keeping the model sufficiently restricted so that computation time does not explode, and inferred trees are still relevant from a biological point of view.

The need for a model that allows mutation losses has been established in (Kuipers et al., 2017), but no clear consensus on the model that is most suited to represent the true evolution of tumors has been reached so far, to the best of our knowledge. In our paper, we introduce and follow a restricted version of the Dollo-k model, where the number of mutations in each site and the number of overall mutations is limited — even though our method can be used also in a more relaxed setting. Determining which of the possible models is going to be the basis for effective and efficient tumor phylogeny inference is something that needs to be explored in the future, but it will likely need the development of different methods, and a deeper understanding of the models.

All the data produced and the experimental settings are publicly available and reproducible at SASC’s repository https://github.com/sciccolella/sasc.

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