# Genome analysis SCAN-ATAC-Sim: a scalable and efficient method for simulating single-cell ATAC-seq data from bulk-tissue experiments

# Zhanlin Chen (1,2,<sup>†</sup>, Jing Zhang<sup>3,\*,<sup>†</sup></sup>, Jason Liu<sup>1</sup>, Zixuan Zhang<sup>4</sup>, Jiangqi Zhu<sup>4</sup>, Donghoon Lee<sup>5,6</sup>, Min Xu<sup>7</sup> and Mark Gerstein<sup>1,2,\*</sup>

<sup>1</sup>Department of Molecular Biophysics and Biochemistry, Yale University, New Haven, CT 06520, USA, <sup>2</sup>Department of Computer Science, Yale University, New Haven, CT 06520, USA, <sup>3</sup>Department of Computer Science, University of California, Irvine, CA 92617, USA, <sup>4</sup>School of Electronic Engineering and Computer Science, Queen Mary University of London, London E1 4NS, UK, <sup>5</sup>Department of Genetics and Genomic Sciences, New York, NY 10029, USA, <sup>6</sup>Department of Psychiatry, Icahn School of Medicine at Mount Sinai, New York, NY 10029, USA and <sup>7</sup>Computational Biology Department, Carnegie Mellon University, Pittsburgh, PA 15213, USA

\*To whom correspondence should be addressed.

<sup>†</sup>The authors wish it to be known that, in their opinion, the first two authors should be regarded as Joint First Authors.

Associate Editor: Anthony Mathelier

Received on June 6, 2020; revised on October 17, 2020; editorial decision on November 30, 2020

# Abstract

**Summary:** scATAC-seq is a powerful approach for characterizing cell-type-specific regulatory landscapes. However, it is difficult to benchmark the performance of various scATAC-seq analysis techniques (such as clustering and deconvolution) without having *a priori* a known set of gold-standard cell types. To simulate scATAC-seq experiments with known cell-type labels, we introduce an efficient and scalable scATAC-seq simulation method (SCAN-ATAC-Sim) that down-samples bulk ATAC-seq data (e.g. from representative cell lines or tissues). Our protocol uses a consistent but tunable signal-to-noise ratio across cell types in a scATAC-seq simulation for integrating bulk experiments with different levels of background noise, and it independently samples twice without replacement to account for the diploid genome. Because it uses an efficient weighted reservoir sampling algorithm and is highly parallelizable with OpenMP, our implementation in C++ allows millions of cells to be simulated in less than an hour on a laptop computer.

Availability and implementation: SCAN-ATAC-Sim is available at scan-atac-sim.gersteinlab.org. Contact: zhang.jing@uci.edu. or mark@gersteinlab.org

Supplementary information: Supplementary data are available at Bioinformatics online.

# **1** Introduction

High-resolution single-cell assay for transposase-accessible chromatin using sequencing (scATAC-seq) techniques reveal transcriptional landscapes in a cell-type-specific manner (Buenrostro et al., 2015). Numerous scATAC-seq data analysis approaches, such as those used for calling and defining active regions in various cell types, clustering and deconvolution, have been published (Bravo Gonzalez-Blas et al., 2019; Fang et al., 2019; Liu et al., 2019; Schep et al., 2017; Xiong et al., 2019; Zamanighomi et al., 2018). However, it has been difficult to evaluate the efficacy of these techniques because we do not have *a priori* knowledge of gold-standard cell types. One way to evaluate these analysis methods is to simulate scATAC-seq with ground-truth labels. With this approach, the analysis methods can be benchmarked against one another with quantifiable parameters that affect the separability of different cell types. There are three major challenges in simulating realistic scATACseq data. First, each open chromatin region can only be captured zero, one or two times in a diploid genome, resulting in at most two reads at one locus in scATAC-seq. Second, similar to bulk ATACseq data, many reads in scATAC-seq come from non-peak, background regions. Third, it is computationally expensive to simulate a dataset with millions of cells in order to evaluate the performance of an analysis method on large datasets. Currently, there are two existing approaches for simulating scATAC-seq data, and both approaches are limited (Table 1).

The first approach randomly samples reads from a curated set of bulk ATAC-seq data (Fang *et al.*, 2019). Bulk ATAC-seq has dramatic variations in the signal-to-noise ratio between experiments. However, cells in scATAC-seq experiments undergo similar procedures, resulting in less background variation. Directly sampling the bulk ATAC-seq can introduce cell-type-specific backgrounds such as

Table 1. Comparison between SCAN-ATAC Sim and previous methods and a brief list of parameters that control the simulation

Features	Direct down sampling	Peak region sampling	SCAN-ATAC Sim
Read-level simulation	Yes	No	Yes
Flexible signal-to-noise	No	No	Yes
Diploid genomic constraint	No	Yes	Yes
Short runtime	No	No	Yes
Long flag	Default value	Long flag	Default value
-cell_number	10 000	-min_frag	1000
-signal_to_noise	0.7	-max_frag	20 000
-frag_num	3000	-extend_peak_size	1000
-variance	0.5	-bin_size	1000

bias from batch effects, confounding downstream analyses. Moreover, it may extract more than two reads for a single genomic locus, thus violating the diploid nature of scATAC-seq experiments. Lastly, direct down-sampling is inefficient. We replicated the method, and it used 26.6 h to simulate one million cells.

The second method simulates individual cells by selecting foreground peaks with a strictly fixed signal-to-noise ratio (Xiong *et al.*, 2019; Zhang et al., 2020). In reality, over half of the reads in scATAC-seq come from background regions. Even though it utilizes drop-out ratios and adheres to the diploid constraint, peak region sampling ignores the background; thus, it is limited in representing real data. Further, this method does not simulate reads. Rather, each cell is represented by simulated foreground peaks. Analysis methods that construct a cell-by-bin read coverage matrix cannot be evaluated using a dataset simulated from this method due to the lack of read-level information. Lastly, repeated sampling from a binomial distribution for all peak regions also limits the efficiency when scaled to millions of cells.

We aim to address these challenges. The lack of a standard, representational synthetic scATAC-seq dataset motivated SCAN-ATAC-Sim, which offers an improvement in simulation quality and reduction in runtime compared to both previous approaches. Our command line software, implemented in C++ with OpenMP parallelization, takes BAM files from bulk ATAC-seq experiments as input and outputs sampled reads for each cell based on user-provided parameters such as the number of reads per cell, total cell number and signal-to-noise ratio (Table 1).

#### 2 Materials and methods

SCAN-ATAC Sim consists of two main steps: data preprocessing and single-cell simulation. Briefly, the process starts with BAM files of bulk ATAC-seq experiments for desired cell types (Fig. 1a). The data preprocessing step defines a cell-type-specific foreground from the merged peaks and a unified background (Fig. 1b). For each cell, the single-cell simulation step samples the foreground and background regions twice without replacement with the probability proportional to the read coverage (Fig. 1c) and randomly selects one read for each sampled region (Fig. 1d). Then, reads from both the foreground and background are combined to form reads in one cell (Fig. 1e). This single-cell simulation step is then repeated for a large number of cells as specified by the user-provided parameter.

#### 2.1 Data preprocessing

The foreground regions are defined by merging peaks from various cell types. Then, the background regions are defined as the complement of the 1 kb-extended foreground divided into bins of fixed sizes (Fig. 1a). The paired and de-duplicated reads from each cell type are intersected with the foreground region to obtain cell-type-specific foreground reads, and unified background weights are created by combining background reads from all cell types (Fig. 1b).



Fig. 1. (a) Bulk-tissue ATAC-seq reads are partitioned into foreground and background based on overlap with merged peaks. The number in each region indicates the read coverage. (b) The cell-type-specific foreground reads are separated, and a unified background is created by combining background reads arcsos all cell lines. (c) The regions are sampled without replacement, with the read coverage as weights for the foreground and background, for paternal and maternal alleles. (d) Reads are sampled from the selected regions using a uniform distribution. (e) All sampled reads are combined to form reads covering a cell. (f) chr1 visualization of bulk and simulated ( $\rho = 0.4$ , f = 1k, c = 100k) CLP cells is shown in the Integrative Genome Browser. (g) Performance for region number  $N_e$  versus cell number is shown for four cores. (h) Percentage of peaks from simulated CLP cells that overlap with CLP bulk peaks is shown, demonstrating the relationship between the signalto-noise ratio and the cell-type specificity of the simulation

#### 2.2 Single-cell simulation

First, for a cell of cell type *c*, the parameter  $N_c$  determines the total number of reads (or regions, since we only sample one read from each region) in an individual cell.  $N_c$  can vary with a log-normal distribution so that every cell has a slightly different number of reads. We also designate a user parameter for the signal-to-noise ratio  $\rho$ . We use  $N_c$  and  $\rho$  to calculate the allocation of foreground and background reads in a cell so that  $N_c = N_c^F + N_c^B$  (1).

$$N_{c}^{F} = \rho * N_{c} N_{c}^{B} = (1 - \rho) * N_{c}$$
(1)

A high signal-to-noise ratio will allocate more reads for foreground regions, thereby making the cell types more separable. Once an allocation between foreground and background is made, the allocations are further halved to mimic reads coming from maternal  $(N_{c,M}^F, N_{c,M}^B)$  and paternal  $(N_{c,P}^F, N_{c,P}^B)$  alleles. Next, SCAN-ATAC Sim performs an efficient two-step sampling to generate representa-tive regions. First, for each cell,  $N_{c,M}^F$  and  $N_{c,P}^F$  regions are separately sampled in two independent trials. Each trial samples without replacement and uses read coverage from the corresponding cell type as weights (Fig. 1c). Representative background regions are generated in a similar manner, with averaged weights from all cell types. Second, one read is randomly sampled from each selected region with equal probability (Fig. 1d). Hence, for any given region in the genome, we can guarantee that no more than two reads are sampled from the maternal and paternal trials because each trial samples without replacement. Lastly, the foreground and background reads are merged for a total of  $N_c^F + N_c^B = N_c$  reads in one cell (Fig. 1e). This process is repeated many times to simulate a massive number of cells.

Specifically, if  $r_{i,c}^F$  represents the total number of reads in the  $i^{th}$  foreground region for cell type c, then the probability of sampling region i can be calculated as  $p_{i,c}^F = r_{i,c}^F / \sum_i r_{i,c}^F$ . In contrast, we calculate a uniform background sampling probability for the background region. For instance, if  $r_{i,c}^B$  represents the read count in the *i*th background region for cell type c, then the uniform background sampling rate for the *i*th background region is  $p_i^B = \sum_i r_{i,c}^B / \sum_i \sum_i r_{i,c}^B$ .

# **3 Results**

# 3.1 Comparing simulated scATAC-seq with bulk ATAC-seq

We analyzed how the simulation mimics cell-type specificity. We simulated and aggregated common lymphoid progenitor (CLP) cells under various signal-to-noise ratios. The pileup signal distribution of simulated scATAC-seq was similar to bulk ATAC-seq (Fig. 1f). We also called peaks from the bulk and simulated data. The amount of overlap between simulated and bulk peaks increased as the signal-to-noise ratio increased (Fig. 1h), indicating that the signal-to-noise ratio contributed to the cell-type specificity of chromatin accessibility, which can be used to vary the difficulty of the benchmarking task. Furthermore, the simulation retained cell-type specificity across different cell types (Supplementary Fig. S2).

#### 3.2 Analyzing simulated scATAC-seq data

Next, we analyzed the simulated scATAC-seq data using SnapATAC (Fang *et al.*, 2019). Under high signal-to-noise ratios, SnapATAC clustered and labeled cells with high accuracy (Supplementary Fig. S3). As the signal-to-noise ratio decreased, the separability between certain cell types decreased in the cell clustering. The read coverage also influenced the analysis outcome, with more reads per cell increasing the foreground signal and separability (Supplementary Fig. S4).

#### 3.3 Complexity and runtime

There are two major computational challenges to simulating scATAC-seq data. First, up to millions of representative foreground

and background regions can be selected with varying probabilities without replacement for one cell. Second, one experiment can contain tens of thousands to millions of cells. To address these challenges, we used two techniques to implement a highly efficient and scalable software for the sampling procedures mentioned in Section 2.3.

First, to improve the efficiency of single-cell sampling, we implemented a reservoir-sampling algorithm to select the representative regions. If *n* represents the number of regions and *m* represents the number of regions to be selected, weighted reservoir sampling without replacement can be performed with  $O(n) + O(m \cdot \log(n/m))O(\log m)$ , as compared to  $O(n \cdot m)$  for traditional weighted sampling methods (Efraimidis and Spirakis, 2006). Especially in our case where  $n \gg m$ , weighted reservoir sampling approaches O(n).

Second, we further parallelized our method in a cell-wise fashion with multicore parallelism using OpenMP, which offers a scalable improvement in runtime. By utilizing both approaches, we enabled the sampling of millions of cells in less than an hour. We demonstrate that SCAN-ATAC-Sim achieves a scalable speed-up for a cell group of four cell types. The runtime does not change with signalto-noise ratio  $\rho$  but varies linearly with both cell and region number  $N_{\epsilon}$  (Fig. 1g).

# **4** Discussion

We introduce a new software tool for constructing a labeled scATAC-seq dataset from bulk ATAC-seq data via guided downsampling. This tool is useful for evaluating the efficacy of single-cell data analysis techniques by simulating scATAC-seq data while adhering to various biological constraints. The acceleration obtained via multicore parallelism permits the simulation of millions of cells in less than an hour. Moreover, this tool is highly scalable and offers a space-time tradeoff to match the rate of growth in the number of cells sequenced for scATAC-seq.

# Funding

This work was supported by the National Institutes of Health [U01MH116492], National Institutes of Mental Health [K01MH123896] and National Institute of General Medical Sciences [R01GM134020].

Conflict of Interest: The author(s) declare(s) that there is no conflict of interest.

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