Gene expression

ESCO: single cell expression simulation incorporating gene co-expression

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Abstract

Motivation: Gene-gene co-expression networks (GCN) are of biological interest for the useful information they provide for understanding gene-gene interactions. The advent of single cell RNA-sequencing allows us to examine more subtle gene co-expression occurring within a cell type. Many imputation and denoising methods have been developed to deal with the technical challenges observed in single cell data; meanwhile, several simulators have been developed for benchmarking and assessing these methods. Most of these simulators, however, either do not incorporate gene co-expression or generate co-expression in an inconvenient manner.

Results: Therefore, with the focus on gene co-expression, we propose a new simulator, ESCO, which adopts the idea of the copula to impose gene co-expression, while preserving the highlights of available simulators, which perform well for simulation of gene expression marginally. Using ESCO, we assess the performance of imputation methods on GCN recovery and find that imputation generally helps GCN recovery when the data are not too sparse, and the ensemble imputation method works best among leading methods. In contrast, imputation fails to help in the presence of an excessive fraction of zero counts, where simple data aggregating methods are a better choice. These findings are further verified with mouse and human brain cell data.

Availability and implementation: The ESCO implementation is available as R package ESCO. Users can either download the development version via github (https://github.com/JINJINT/ESCO) or the archived version via Zenodo (https://zenodo.org/record/4455890).

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

A synchronization between gene expression leads to gene co-expression. Cell heterogeneity, due to cell type or cell cycle, can generate correlations between genes that are highly expressed in similar cells. Alternatively, any form of gene cooperation within a cell type, such as gene co-regulation, also results in co-expression. To differentiate these two settings, we refer them as the gene co-expression across heterogeneous cell groups and gene co-expression within homogeneous cell groups respectively, throughout this article. Understanding gene co-expression in the former setting helps with cell-type identification, and in the latter setting, it helps detect gene regulation relationships and can further provide insights into genetic disorders (Pang et al., 2020; Poliouδakis et al., 2019).

Single-cell RNA sequencing (scRNA-seq), a recent breakthrough technology that paves the way for measuring transcription at single cell resolution to study precise biological functions, allows us to target gene co-expression within homogeneous cell groups for the first time. Indeed, early statistical models argued that genes within homogeneous cell groups were independent (Quinn et al., 2018). However, they overlooked the investigations from the biological end, which reveal that correlation arises due to the stochastic nature of gene expression and gene regulation dynamics (Raj et al., 2006). scRNA-seq data present many challenges for co-expression analysis, due to the sparsity of counts, which include many zeros, mainly arising from low capture and sequencing efficiency in the data collecting process. Sparsity occurs in both a gene- and a cell-specific manner and is observed to have the greatest impact on genes that have low expression.

An ever-growing literature attempts to address these challenges using imputation and other denoising methods. To systematically benchmark these methods, we require realistic simulation tools to
construct a ground truth for scRNA-seq data with realistic technical noise; however, currently there is a paucity of methods for this purpose.

Numerous scRNA-seq simulators using both non-parametric and parametric approaches have been proposed during recent years, e.g. Splat (Zappia et al., 2017), SymSim (Zhang et al., 2019a), PROSSTT (Papadopoulos et al., 2019) and SERGIO (Dibaenia and Sinha, 2020). Each of those methods focuses on producing realistic marginal behavior of gene expression, and successfully modeling these features, as well as capturing cell type heterogeneity. But, those simulators either ignore gene co-expression, or they generate it in a way that is hard to benchmark. Real data clearly display gene co-expression within homogeneous cell groups (Supplementary Fig. S1A) and gene co-expression across heterogeneous cell groups (Supplementary Fig. S1B). By contrast, almost all gene pairs show no correlation for simulated data generated using Splat, even without the challenge of added technical noise (Supplementary Fig. S1C). While the data simulated by SymSim may show a modest level of gene co-expression (Supplementary Fig. S1D left panel), that correlation arises from the cell type confounding, rather than true gene-gene interaction (Supplementary Fig. S1D right panel). PROSSTT, shares a similar issue with SymSim, in that it also introduces co-expression via a random dot product model. SERGIO, on the other hand, directly approximates the biological gene expression process via a series of differential equations with gene regulation relationship as constrains, therefore it is able to introduce gene co-expression based on real gene-gene interactions. However, it is hard to anticipate the final level of co-expression from the imposed gene regulation relationship, hence it is difficult to systematically benchmark the outcome.

Here, we propose a new simulation tool, Ensemble Single-cell expression simulation incorporating gene CO-expression, ESCO, which is constructed as an ensemble of the best features among current simulators to preserve the marginal performance, while allowing easily incorporating co-expression structure among genes using a copula. Particularly, ESCO allows realistic simulation of a homogeneous cell group, heterogeneous cell groups, as well as complex cell group relationships such as tree and trajectory structure, together with a flexible input of co-expression. As for technical noise, ESCO integrates the parametric and non-parametric approaches in current literature and gives the user flexibility to choose. In order to mimic a specific real dataset, ESCO can estimate all the hyperparameters in a feasible way for both a homogeneous cell group or heterogeneous cell groups. ESCO is implemented in the R package ESCO, which is built upon the R package Splat (Zappia et al., 2017), in order to provide a unified software framework.

2 Materials and methods

2.1 Models

Despite their differences, current simulation approaches arguably follow a general flowchart (Fig. 1). For example, Splat (Zappia et al., 2017) simulates scRNA-seq data using a hierarchical model in which the gamma-Poisson distribution imposes a mean and variance trend; SymSim (Zhang et al., 2019a) is based on a similar hierarchical model with gene kinetics guiding the hyperparameter selection, a non-parametric approach to introduce more realistic noise, and a focus on tree-structured heterogeneity; PROSSTT (Papadopoulos et al., 2019) aims to simulate realistic cell trajectories using a model based on Brownian motion; SERGIO (Dibaenia and Sinha, 2020) starts from the gene regulation relationship and solves a series of stochastic differential equations given by gene kinetics to impose those regulations. The more complex non-parametric modeling tends to fit data better than parametric modeling, given that the aim is to mimic data for which the model has already been trained. However, this approach is not practical for producing simulated data similar to a new dataset. For example, the non-parametric methods like SymSim and SERGIO use grid search over a large number of tuning parameters. By contrast, the parametric Splat approach can be tuned to data by fitting a one-step statistical regression model. ESCO also follows the general flowchart in Figure 1, but it aims to incorporate the best features from the existing methods. Figure 2 illustrates the superiority of ESCO, as it allows simulation of scRNA-seq data with various cell heterogeneity patterns and customized gene co-expression patterns. The correlation pattern input is successfully replicated in the simulated data, both within and between homogeneous cell groups. In this section, we elaborate on the specific simulation models that ESCO adopts, following the framework outlined in Figure 1. More detailed descriptions of the simulation models and the time complexity for simulating large complex data are provided in Supplementary Notes S1 and S2, respectively.

Base expression level We simulate base expression level in an empirical way that allows inputting any density function, either non-parametric or parametric. Particularly, we denote the base expression level for gene $g$ as $\lambda_g$, and we let $\lambda_g \sim \Lambda$ for all $g$.

Extrinsic variation The heterogeneity of cell groups is driven by the differential expressed (DE) behavior among certain gene sets across groups. Therefore we implement the cell group heterogeneity, i.e. the extrinsic variation, via modeling the behavior of DE genes ($G^{DE}$). We use the random dot product model to introduce this heterogeneity by imposing a DE factor generated separately on the otherwise homogeneous gene expression means. Particularly, we generate the different cell group structures we want, via modeling the DE factor $f^g_k$ for gene $g$ in cell group $k$ in each of the following ways.

A. Discrete cell groups: In order to generate clear and distinguishable cell groups, we randomly split the set of DE genes into subsets, each is identified as marker genes for a cell group. Then we simulate the DE factor for each marker gene set as a LogNormal random variable with different mean and variance indexed by group identity.

B. Tree-structured cell groups: We utilize the idea in SymSim (Zhang et al., 2019a), which makes the DE factor of similar cell groups more related to each other. Particularly, we generate the DE factor from a multivariate normal distribution, where the covariance matrix is given by the tree structure of the data. Additionally, in order to assure the identifiability of different cell groups, we introduce extra heterogeneity via strengthening the DE factor for a small proportion of DE genes, which are identified as marker genes in this setting (different from those in the discrete cell group setting).

C. Continuous cell trajectories: We utilize the idea in PROSSTT (Papadopoulos et al., 2019), which uses Brownian motion to generate the DE factors, so that the smooth cell heterogeneity can be generated.

Finally, we generate the base expression with an adjustment of library size for each gene $g$ in cell $c$ as

$$\lambda_{gc} = L_c \lambda_{gc}' \sum_{g} \lambda_{gc}'$$ for each cell $c$,

where $\log L_c \sim \mathcal{F}_L$, and $\lambda_{gc}', \sim \lambda_{gc}^{\text{DE}}$ if $g \in G^{DE}$, with $k(c)$ denotes the group identity of cell $c$; otherwise $\lambda_{gc} \sim \lambda_g$. 

Intrinsic variation

Marginal distribution: Gene expression in individual cells is an inherently stochastic process (Raj et al., 2006). If the gene regulation is ignored, this process is just a simple two state birth-death process. The steady-state distribution for this stochastic process in most cases turns out to be a Gamma-Poisson, Beta-Poisson or Poisson, which is justified from the theoretical biochemistry aspect (Gruen et al., 2014; Kim and Marioni, 2013), the experimental data sampling aspect (Quinn et al., 2018), and also the common observations from the data. Splat (Zappia et al., 2017) and PROSSTT (Papadopoulos et al., 2019) utilize the negative binomial model in the simulation of marginal gene expression; while SymSim (Zhang et al., 2019a) uses a Beta-Poisson instead; SERGIO (Dibaenia and Sinha, 2020) simulates the gene expression via solving the series of ordinary differential equation functions following the literature about gene kinetics with regulation (Schaffter et al., 2011).

ESCO adopts the negative binomial model, since it is widely accepted in the literature and enjoys support from biochemistry,
experimental data sampling and empirical observations. Particularly, following Splat (Zappia et al., 2017), we can naturally enforce a mean-variance trend by simulating the Biological Coefficient of Variation (BCV) for each gene. BCV is defined as the square root of the standard deviation divided by the mean, i.e. the square root of the coefficient of dispersion. It has been pointed out (McCarthy et al., 2012) that one should not assume a common dispersion for all the genes, as a gene-specific variation is often detected in RNA-seq case studies. Splat simulates BCV as a weighted sum of a common dispersion and a gene-specific dispersion, such that some information can be shared across genes to benefit the estimation, while preserving the gene-specific variation.

**Co-expression:** The gene expression (either the truth or the observed) is not necessarily independent even within cells of the same type, resulting from gene regulation. Characterizing the joint distribution requires solving the steady distribution of multiple correlated stochastic processes, which usually does not have a closed-form solution and requires large computational power (Dibaeinia and Sinha, 2020; Pratapa et al., 2020). Since the marginal distribution of gene expression is understood fairly well, naturally, we think of using the copula to model the gene dependence. This idea is shown to be successful in Inouye et al. (2017) to model bulk RNA-seq data.

A copula is defined by a joint cumulative distribution function (CDF), \( C(\mathbf{u}) : [0,1]^p \rightarrow [0,1] \) with uniform marginal distributions. One of the most popular copula models is the Gaussian copula, which is defined simply as: Delete

\[
C_{\Sigma}^{Gauss} = N_2(\Phi^{-1}(u_1), \Phi^{-1}(u_2), \ldots, \Phi^{-1}(u_p))
\]

where \( \Phi^{-1} \) denotes the inverse function of standard normal CDF, and \( N_2 \) denotes the joint CDF of a multivariate normal random vector with zero means and correlation matrix \( \Sigma \). Due to the well-known consistency between \( \Sigma \) and the empirical Pearson correlation matrix, the Gaussian copula allows for directly interpretable dependence simulation, and therefore is adopted by ESCO.

**Technical noise:** Currently, there are mainly two single cell library preparation protocols: (i) full-length mRNAs profiling without the use of UMIs (e.g. with a standard Smart-Seq protocol); and (ii) profiling only the end of the mRNA molecule with the addition of UMIs (e.g. 10x Chromium). The former protocol is usually applied for a small number of cells and with a large number of reads per cell, providing full information on transcript structure. The latter is normally applied for many cells with shallower sequencing, and it is impacted less by amplification and gene length biases. We focus on the UMI-based protocol in this article because it is usually less biased with greater sparsity.

There currently exist two approaches to simulate the technical noise: one is based on data generating process, and the other is based on data visualization and fitting. As an example of the former, SymSim (Zhang et al., 2019a) uses the empirical approximation of the major steps in the experimental procedures such as mRNA capture, PCR amplification, RNA fragmentation and sequencing, to directly imitate the technical noise. On the other hand, Splat (Zappia et al., 2017) simulates the technical noise by adopting a zero-inflation model, where the zero-inflation probability relates to the gene expression level in a way that comes from the observed trend in the real data.

There are both pros and cons with regard to these two approaches. The empirical approach facilitates the generation of more realistic noise, but suffers from finding appropriate configuration to match a particular dataset (actually, SymSim uses a grid search to do the matching). In contrast, the parametric approach allows a one-step estimation of the parameters from the real data, but can suffer from poor goodness-of-fit due to the mismatch of
ESCO can simulate scRNA-seq data of various cell heterogeneity and gene co-expression. (A) The simulation results for one homogeneous cell group consisting of 200 cells and 500 genes. The first panel displays the heatmap of log2 transformed normalized simulated expression data, where rows represent genes and columns represent cells; 30% of genes are chosen to be co-expressed genes, and the rest are independent genes. The following displays depict, in order, the given correlation structure for co-expressed genes, the simulated correlation structure among those co-expressed genes without noise, and that with technical noise, and the simulated correlation structure for independent genes. (B) The simulation results for three discrete heterogeneous cell groups consisting of 500 cells and 1000 genes. 30% of the genes are chosen to be cell-type DE genes and presumably co-expressed, among which each marks one cell type. Another 10% of genes are chosen to be housekeeping genes, and also presumably co-expressed. The rest are independent non-DE genes. The first display shows the heatmap of log2 transformed normalized simulated data, where different gene types (rows) and cell types (columns) are marked with color bars on the margin. The following displays depict, in order in each row, the given correlation structure for both marker genes of Group2 and co-expressed housekeeping genes, the simulated correlation structure among those co-expressed genes without noise, and that with technical noise; and, at the end of each row the simulated correlation structure among all DE genes across all cells, and that among all independent genes across all cells, with corresponding gene types marked with a color bar on top. (C) The simulation results for five heterogeneous cell groups that follow a tree structure given in the first panel. We simulate 1000 cells and 2000 genes: 30% of genes are chosen to be DE genes and presumably co-expressed, among which 5% are markers; the rest are independent non-DE genes. The second panel shows the heatmap of log2 transformed normalized simulated data. Different cell types are marked with color bars on the column margin, together with the hierarchical clustering of cells. The following displays depict, in order, the resulting correlation structure among all marker genes across all cells, and that among all independent genes across all cells, with corresponding gene types marked with a color bar on top; the given correlation structure for co-expressed marker genes of Neuron1 cells, and the resulting correlation structure among those co-expressed genes. (D) The simulation results for five heterogeneous cell groups that follow a smooth cell trajectory structure given in the top left panel. There are 1000 cells and 2000 genes: 30% of genes are chosen to be DE genes and presumably co-expressed and share the same correlation structure within each branches, and the rest are independent non-DE genes. The following displays depict, in order, the UMAP for the first two dimensions of the simulated data, the heatmap of log2 transformed normalized simulated data for all DE genes in one continuous path (i.e. branches 1 → 2 → 5), with branch ID marked with a color bar on top; the given shared correlation structure for the DE genes, and the resulting correlation structure simulated of those genes within each branch.
models. Therefore, ESCO integrates both procedures and gives users the freedom to choose between the two.

2.2 Estimation
ESCO facilitates mimicking any particular dataset, consisting of either homogeneous or heterogeneous cell groups, by estimating the hyperparameters from the data. Through learning the parameters in the parametric model, this approach fits data as well as possible, given the limitations of the parametric choice, as illustrated by comparing mouse brain cells (Zeisel et al., 2015) with simulated outcomes. A good match is obtained for mean, variance of expression, UMAP of cells, percent zero outcomes and co-expression patterns (Supplementary Fig. S2).

Next, we elaborate on our specific estimation strategies. Recall that ESCO takes a hierarchical modeling approach, paired with a copula. As such, an empirical Bayesian approach to parameter estimation would be appropriate. However, it is usually infeasible to compute the solution. Therefore, we follow Splat and estimate the parameters in each layer separately. Particularly, we assume the data are already normalized (i.e. no batch effect arises due to technical reason) and have disjoint marker gene sets across cell types, and consider the three estimation tasks in the following.

Estimating the heterogeneity: We have introduced three types of heterogeneity of gene expression (discrete, tree and trajectory), but we only present an estimation procedure for the discrete one here, and leave the estimation of the more complex structure of the other two models to future work. Nevertheless, ESCO is usable for these two models provided the tree structure and trajectory information are available from side information. When the tree/trajectory information is not available, in contrast with SymSim and PROSSIT, we caution against using a grid search to chose model parameters due to the difficulty in determining a good ‘match’ in these complex heterogeneity cases. SymSim and PROSSIT use summary statistics, such as global mean and variance, as standards for a good ‘match’, but two datasets can have similar mean and variance and totally different cell heterogeneity structure.

Following our modeling of the discrete heterogeneous cell groups, we first split all the genes to DE and non-DE genes based on their Area Under the Curve (AUC) scores in cell group prediction using SC3 (Kiselev et al., 2017), provided that we already have the true cell group annotation. Particularly, we use 0.7 as our cutting threshold of the AUC score, i.e. classifying the genes with AUC score no less than 0.7 as DE genes and the others as non-DE genes.

We then use the DE genes to estimate the DE factors. Particularly, we divide those DE genes into marker genes for each cell group based on their classification result from SC3 (Kiselev et al., 2017). We assume that the mean distribution of marker genes in their marked cell group follows the same distribution in the other cell group and a DE factor that follows LogNormal distribution indexed by the cell type. Therefore, we estimate the DE factor for marker genes of cell group $k$ via fitting a LogNormal distribution on the ratio of their sample mean within cell group $k$ and those outside cell group $k$.

Estimating the intrinsic variation
Marginal: As for estimating the parameters related to marginal intrinsic variation, we follow the technique used in Splat (Zappia et al., 2017), with a few refinements. We allow non-parametric fitting of the library size distribution and base mean distribution, which can be done quickly by computing the empirical CDF and also later on sampled from using Metropolis-Hastings sampling due to the univariate nature. One may refer to Zappia et al. (2017) for further details about the estimation procedure for other marginal parameters included in the algorithm, such as BCV and outlier.

Copula: To circumvent challenges due to technical noise and sparse counts, we cluster similar cells and form metacells (Baran et al., 2019) and then estimate $\Sigma$ in the Gaussian copula model. As an integrated version of the original real data, the size of metacells must be carefully selected so that the technical variation can be reduced, while some biological variation can be preserved. We refer the reader to the source paper of MetaCell (Baran et al., 2019) for further details.
former is assessed by computing the Adjusted Rand Index (ARI) and the latter using Area Under the Curve (AUC). We evaluate ARI and AUC for each imputation method under a range of sparsity levels (i.e., zero proportion) for the marker genes within cell groups, the housekeeping genes across cell groups and the DE genes across cell groups (Fig. 3B).

To compute ARI we choose the number of clusters that maximizes the score, calculated over a range of clusters numbers (2–9). To calculate AUC we label gene pairs as connected or un-connected based on the co-expression significance in permutation testing of the simulated truth. We then assess the prediction accuracy (AUC scores) of connections for each imputation method using their estimated co-expression. All the results are averaged over 10 replicates.

We observe the following results (Fig. 3B). (i) Generally, imputation helps (beat the un-imputed raw data, depicted by the bold dashed black line) recovering both gene co-expression within homogeneous cell groups and gene co-expression across heterogeneous cell groups, but fails to help much with gene co-expression within homogeneous cell groups when facing excessive sparsity (>90% zeros), while tends to introduce spurious gene co-expression across heterogeneous cell group when facing moderate sparsity (~60–80% zeros). (ii) As for a comparison among different methods, there is no universal winner for all settings, but the ensemble method, depicted by the bold black line, provides the best or close to the best performance across almost all settings we considered.

In the following section, we aim to verify our findings of imputation using real data. It is conjectured that the co-expression of marker genes in the mouse brain will be similar to that of the human brain. Therefore, we expect the recovered gene correlation from a dataset measuring mouse brain will follow a similar pattern to those from the dataset measuring the human brain. Particularly, we use Zeisel data (Zeisel et al., 2015) for the mouse brain and Velmeshev data (Velmeshev et al., 2019) for the human brain. The Zeisel data have deeper sequencing for single cells and consequently are less noisy, with less sparsity, compared with the Velmeshev data, which have a much greater number of nuclei sampled, each with fewer reads. Therefore, we can see the influence of the sparsity level on gene co-expression by directly comparing
these two datasets. We select five common cell types in both datasets and use the Zeisel data as the benchmark. We evaluate the correlation matrix of marker genes before and after imputation of Zeisel data, across cell types and within one cell type (i.e. interneurons). Figure 3C(a) plots both the gene co-expression across heterogeneous cell groups and gene co-expression within homogeneous cell groups before and after imputation with EnImpute method (Zhang et al., 2019b) using Zeisel data, while Figure 3C(b) plots the same results but using the Velmeshev data. We can see that for the Zeisel data (moderate level of sparsity), imputation enhances the gene co-expression pattern both within homogeneous and across heterogeneous cell groups. In contrast, for the Velmeshev data (excessive sparsity), imputation fails to help much to recover the gene co-expression across heterogeneous cell groups pattern, while failing utterly for gene co-expression within homogeneous cell groups, which is expected, as it is a harder task. This investigation supports some of our findings of imputation, i.e. imputation can generally help, but may fail as sparsity levels increase to a very high level.

3.3 Data aggregating is a better way to recover GCN with excessive sparsity

Despite the excessive sparsity in the Velmeshev data, these data have the advantage of abundant numbers of cells, which inspired us to explore another approach for recovering gene co-expression: data aggregation that utilizes the abundance of measured cells. We introduce two methods below, a simple heuristic (AOB) and a complex algorithm (BigScale).

Averaging over cell bags. If one has successfully assigned the cell type labels, one may be able to use the simple procedure of averaging gene expression over random splits within cell types, and then compute the gene co-expression based on those averaged values (Polioudakis et al., 2019). We will refer to this procedure as AOB (Averaging Over Bags). The only tuning parameter here is the bag size, which should be chosen carefully so that we can mitigate the influence of sparsity and other noise, while still maintaining some variability among samples.

Pre-clustering and transforming the expression value. More recently, a method called BigScale (Iacono et al., 2019) was developed for the problem of recovering GCN in a similar, but more complex way. This algorithm first clusters cells sharing highly similar transcripts together, and then treats them as biological replicates to evaluate the noise and an indirect measure of correlation. This method works well when there is a sufficiently large number of cells for meaningful cell clusters to form, but it is computationally challenging.

We find both methods work well in recovering gene co-expression across heterogeneous cell groups [Supplementary Fig. S3C(c)], though neither successfully recover gene co-expression within homogeneous cell groups. Future efforts are needed to recover these subtle signals.

4 Discussion

In this article, we propose a new scRNA-seq simulator, ESCO, which borrows the good features of the current state of art simulators in an ensemble, while for the first time, allowing both interpretable and controllable gene co-expression generation. Specifically, ESCO allows realistic simulation of various cell group structure, ranging from simple homogeneous cell groups to tree-structured discrete cell groups to continuously changing cell trajectories, together with gene co-expression. ESCO outperforms other methods as it preserves the highlights of all the other existing simulators in one R package, including the hierarchical semi-parametric modeling of homogeneous groups from Splatter, the tree-structure generation from SymSim and the trajectory generation from PROSTT, all while interacting gene-gene interactions. Specifically, ESCO allows the flexible generation of both gene co-expression across heterogeneous cell groups arising from a cell group structure and gene co-expression within homogeneous cell groups arising from gene-gene interaction in one functional cell group, which have been overlooked and underdeveloped in other methods.

There is still much room for future work in this area. The efficient estimation of the hyperparameters from the real data in the tree-structured cell group and continuous cell trajectories scenario still needs improvement. Currently, most simulators rely on a grid search of parameters to find parameters that fit a particular data, but these parameter choices do not extend to new settings, and it is extremely challenging to simulate data similar to new datasets. The ability to simulate realistic batch effects in various settings is also not satisfactory in the current methods. ESCO, which mimics Splat in this regard, shares this shortcoming. A careful, deep-dive to produce realistic batch effects is needed.

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