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# Robust Learning of Transfer Functions for Single-Cell Transcriptomics Depth Normalization

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Anonymous Authors<sup>1</sup>

## Abstract

Normalization is a critical step in data processing that influences downstream analyses. Normalization aims to adjust for technical variations in data acquisition, facilitating accurate comparisons among observations. In this paper, we identify key challenges in scRNA-seq normalization, including the simplex nature for reads, compositional bias from mRNA population, technical and biological outliers, and non-linear relationship between the input and output. We introduce a new framework to address these challenges by modeling the measurement function and robust learning of parameters. Empirical validation on real datasets demonstrates the effectiveness of the proposed normalization method, RFNorm, in preserving lower-dimensional mathematical structures crucial for cell type and state analysis. This is assessed through the invariance of k-nearest neighbor graphs comparing the performance of RFNorm against established methods.

## 1. Introduction

Many measurements in empirical sciences are subject to perturbations and noise, both technical and intrinsic. In genomics, the development of high-throughput methods like next-generation sequencing (NGS) and single-cell RNA sequencing (scRNA-seq) allows for the analysis of millions of cells and thousands of features per cell under various conditions, presenting new challenges in data processing (Svensson et al., 2017; Amezquita et al., 2020). These include normalization for instrument noise (L. Lun et al., 2016; Bacher et al., 2017; Lopez et al., 2018; Hafemeister & Satija, 2019), data integration across various sources (Korsunsky et al., 2019; Liu et al., 2020; He et al., 2024), and imputation for sparse data (Huang et al., 2018; Van Dijk et al., 2018; Marouf et al., 2020). Normalization, a cornerstone concept in statistics, aims to transform data into a standardized format to enable accurate comparison and analysis. Ideally, normalization should: (1) correct for instrumentation-specific biases (Vallejos et al., 2017), (2) manage statistical distributional properties to support analytical techniques (Ioffe

& Szegedy, 2015), and (3) enable meaningful comparison with respect to hypotheses of interest (Love et al., 2014).

NGS data are prone to instrumentation-specific biases, notably influenced by sequencing depth—the number of total sequenced reads obtained from each cell (Robinson & Oshlack, 2010; Vallejos et al., 2017; Evans et al., 2018; Boeshaghi et al., 2022). A fundamental aspect of NGS is its ability to quantify the presence of RNA or DNA in terms of counts. However, these counts not only depend on the input material but also subject to compositional bias, especially when the library is not sequenced to saturation, a common scenario in scRNA-seq (Love et al., 2014; Evans et al., 2018; L. Lun et al., 2016). Such bias skews the observed counts based on the composition of the RNA population. Furthermore, the intrinsic nature of NGS data is compositional: the abundance of any nucleotide fragment can only be understood in relation to others, constrained by the sequencer’s finite capacity. This limitation frames the data within an Aitchison Simplex rather than a conventional Euclidean space, where an increase in one fragment’s abundance can misleadingly appear as a decrease in another, thereby leading to spurious gene correlations (Su et al., 2023).

Normalization methods must effectively manage the statistical distributional properties of scRNA-seq data to support downstream analysis. These data exhibit inherent complexities such as heteroskedasticity and gene-gene correlations. Heteroskedasticity, the uneven variance across data points, arises from the intrinsic properties of RNA molecules and varies with biological conditions such as environmental influences and cell cycles (Ahlmann-Eltze & Huber, 2023). On the other hand, gene-gene correlations, which emerge from regulatory interactions between genes, reflect essential biological functions and should be preserved (Sun et al., 2021). Therefore, normalization should adjust for heteroskedasticity to stabilize variance across samples, while maintaining the gene-gene correlations. Current normalization methods often treat genes as independent and identically distributed, or group them into bins by expression levels. This approach can distort the true biological signal in data that inherently exhibits a mixed distribution.

Normalization must enable meaningful comparisons. In scRNA-seq, we are interested in biological signals to distin-

guish the functional states of individual cells, track developmental pathways, or identify new cellular phenotypes. A critical and intriguing question emerges: even if we could precisely count the number of mRNA molecules, how do we compare the genomic profiles between two cells? This comparison must address two primary aspects: the compositional nature of cells and the variability from biological factors such as differential expression (DE). Even if a sequencer could perfectly capture every RNA molecule in a cell, cells themselves are inherently compositional due to constraints like volume and energy, which limit RNA synthesis. This is evidenced by observations that smaller cells of the same type contain proportionally less total mRNA (Quinn et al., 2019). Furthermore, optimal normalization should ensure that non-DE genes maintain consistent normalized counts across conditions, while DE genes display read counts that accurately reflect the true differences in transcripts per cell. Additionally, the comparison of molecular profiles can be related to high-dimensional geometric shape analysis in Geometric Morphometrics, such as resistant fit Procrustes superimposition followed by thin-plate spline to visualize morphological deformation between two objects (Zelditch et al., 2012).

In this study, we introduce a framework for normalization and explicitly address properties (1)-(3) by modeling measurement functions to mitigate instrument-induced variations, accounting for out-of-distribution values, and robustly learning the parameters by minimizing empirical risk. Through our proposed normalization framework and simulation study, we highlight the limitations of existing methods, which often assume that a single scalar factor can adequately correct for most technical variations. We introduce an R package called RFNorm, incorporating more realistic assumptions and robust optimization into our model. To evaluate the effectiveness of RFNorm, we assess its performance in preserving low-dimensional mathematical structures as outlined in (Ahlmann-Eltze & Huber, 2023), and verify accuracy and Type 1 error control using DEs in paired Bulk RNA-seq (Squair et al., 2021) and constructed null datasets (Soneson & Robinson, 2018). Our findings show that RFNorm achieves high k-NN graph overlap, effectively preserving biological relationships across various datasets, and performs comparably in identifying marker genes while accurately managing Type 1 error rates comparing with the other two popular scRNA-seq depth normalization methods such as CP10K and Scran.

## 2. Related Works

Normalization can be viewed as a two-step transformation (as in Fig 1b) applied to the raw counts to produce the normalized number (Ahlmann-Eltze & Huber, 2023; Boeshaghi et al., 2022). In this paper, we focus on depth normal-

ization for single-cell RAN-seq data. Put the assumptions of different normalization methods under the same framework enable us to see if it resolves some of the key challenges.

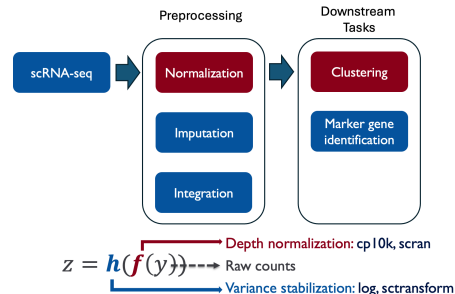


Figure 1. Overview of Normalization in scRNA-seq.

The variance stabilization has a long research history in statistics. Many approaches have been proposed since the era of Bulk RNA-Seq (Anders & Huber, 2010; Robinson & Oshlack, 2010; Love et al., 2014; Hafemeister & Satija, 2019; Laue et al., 2021; Choudhary & Satija, 2022). However, recent research indicates that the simple approach, the logarithm with a pseudo-count ( $\log_1 p$ ), performs as well or better than the more sophisticated alternatives, for instance SCTransform (Hafemeister & Satija, 2019), for variance stabilization in terms of the preservation of low dimensional graphs (Ahlmann-Eltze & Huber, 2023). In this work, we choose  $\log_1 p$  as the variance stabilization function and focus on depth normalization and focusing on the design of depth normalization function.

Most sequencing depth normalization approaches (CP10K (Luecken & Theis, 2019), scNorm (Bacher et al., 2017), BayNorm (Tang et al., 2020)) use the same size factor assumption as SCTransform. However, the scaling transformation has limitations. First, since the observed counts are simplex data, scaling by the sequencing depth does not remove the constant sum constraint and cannot convert closed sequencing data into an “open” unit such as concentration (Quinn et al., 2019). Secondly, transforming only by sequencing depth is a within-sample transformation, whereas a between-sample transformation (Maza, 2016; Evans et al., 2018; Love et al., 2014; L. Lun et al., 2016) would be more appropriate because of the compositional bias. To be more specific, since most of the time we cannot saturate the library during sequencing, the observed counts would be biased by the composition of mRNA population in each cell. Thirdly, the complexity of depth normalization function  $f(y)$  is a unique challenge that has been overlooked.

### 3. Normalization Framework

#### 3.1. Preliminaries

In transcriptomics and single-cell RNA sequencing (scRNA-seq), each biological cell is assumed to contain a certain numbers of measurable RNA molecules across  $G$  different types of transcripts (or genes), denoted as  $n_i = x_1 + x_2 + \dots + x_G$ , where  $n_i$  may vary for each measured cell. The scRNA-seq instruments aim to recover these counts as integer vectors, producing observations  $(y_1, y_2, \dots, y_G)$  for each cell, across potentially tens of thousands or even millions of cells. Normalization in this context is a two-step transformation applied to the raw counts  $y_g$  to produce the normalized number  $z_g$  (Ahlmann-Eltze & Huber, 2023; Boeshaghi et al., 2022). This process can be formally expressed as  $z_g = \mathbf{h}(\mathbf{f}(y_g))$ , where  $\mathbf{h}$  is the variance stabilization function and  $\mathbf{f}$  is the depth normalization function. For example, SCTransform (Hafemeister & Satija, 2019), a widely used normalization method, employs Negative Binomial (NB) regression to model UMI counts. To account for sequencing depth, SCTransform utilizes the link function  $\log \mu = \beta + \log n_i$  in NB regression, where  $n_i$  is the total counts of the cell  $i$ . The link function is designed on the size factor assumption (Vallejos et al., 2017), where the observed counts is the true counts scaled by some cell-specific size factor. Variance stabilization is then achieved using the feature/gene-level Pearson residuals.

#### 3.2. Problem Setup

We propose a framework to characterize the normalization process in scRNA-seq experiments involving  $N$  cells and  $G$  genes. We define three conceptual subspaces: Biological Space  $V_B$ , Measurement Space  $V_M$ , and Normalization Space  $V_N$ , as illustrated in Fig 2. Biological Space is a subspace of Euclidean space  $\mathbb{R}^G$ , representing the true molecular profiles. Measurement Space is a subspace of Aitchison Simplex space because the observed counts are compositional data (Quinn et al., 2019). Ideally, Normalization Space would also be a subspace of Euclidean space  $\mathbb{R}^G$ , representing the post-normalization profiles.

Data in Biological Space  $V_B$  is inherently unknowable to us. We assume that a measurement function  $\phi$  maps vectors from  $V_B$  to  $V_M$ , where  $Y \in V_M$  represents the experimental data collected. A normalization function  $\omega$  is then defined to map vectors from  $V_M$  to  $V_N$ . Ideally,  $\omega$  would serve as the inverse function of  $\phi$ , enabling retrieval of the true biological states. However, inverting this function is complex due to the unknown parameters and nature of  $\phi$ .

Normalization methods thus construct the Normalization Space and formulate  $\omega$  based on an estimated  $\hat{\phi}$  under assumptions about the measurement function. For instance, if the observed counts are assumed to be the true counts

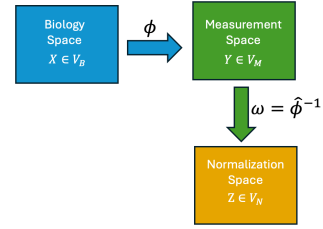


Figure 2. Schema of the normalization framework. Suppose  $X = [\mathbf{x}_i]$ ,  $Y = [\mathbf{y}_i]$ ,  $Z = [\mathbf{z}_i]$  be matrices with dimensions  $G \times N$ . Column vectors  $\mathbf{x}_i$ ,  $\mathbf{y}_i$ ,  $\mathbf{z}_i$  represent observations in their respective subspaces, for instance,  $\mathbf{y}_i = (y_{1,i}, y_{2,i}, \dots, y_{G,i})$ ,  $i \in [N]$ .

scaled by a size factor, then  $\mathbf{y}_i = \hat{\phi}(\mathbf{x}_i) = s \times \mathbf{x}_i$ . The sequencing depth can then be used as the estimator for the size factor, leading to  $\mathbf{z}_i = \omega(\mathbf{y}_i) = 1/s \times \mathbf{y}_i = 10^4/n_i \times \mathbf{y}_i$ , where  $n_i = \sum \mathbf{y}_i$ , where  $n_i = \sum \mathbf{y}_i$  represents the total number of reads in cell  $i$  and  $10^4$  is included for numerical stability. To this end, we integrate CP10K method to our normalization framework.

Our framework serves as a conceptual model to address the three fundamental properties of effective normalization, as outlined in the introduction. By now, it is not a normalization method per se, but rather provides a mindset for designing normalization approaches. Property (1), the correction of instrument-related biases, is approached through the inversion of measurement functions  $\phi$ , which helps to minimize inherent variations in the measurement process. For Property (2), we recommend implementing transformations like the shift-log within the normalization function, stabilizing variance across samples and ensuring statistical integrity. With regard to Property (3), which aims to facilitate meaningful comparisons in alignment with research hypotheses, this mindset guides us to optimize empirical risk for data subsets considered representative and within normal distribution. These strategies underscore the framework’s utility as a foundation for developing robust normalization methods, further discussed in sections 3.3 and 4.2.

#### 3.3. Normalization Paradigms

This section delineates two primary normalization paradigms: between-sample normalization and within-sample normalization. Within-sample normalization aims to scale data within a single sample to reduce internal variability and ensure comparability within that sample. However, the simplex nature of scRNA-seq data and inherent compositional bias often render within-sample normalization suboptimal for comprehensive data analysis, as depicted in Fig 3.3(a-c).

Conversely, between-sample normalization, which we advocate for, adjusts measurements across different samples to

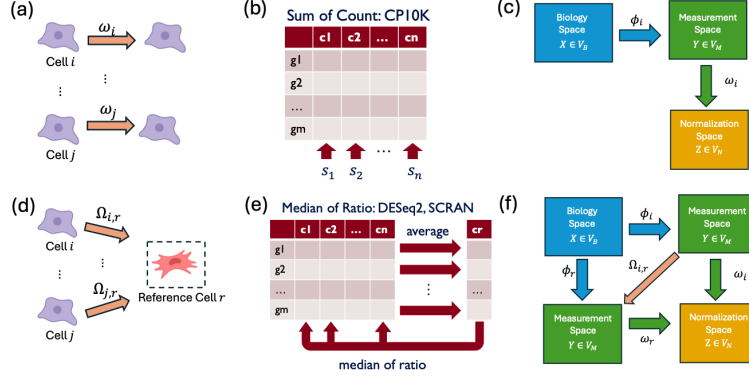


Figure 3. Illustration of normalization paradigms and their components. (a) Represents within-sample normalization where each cell is normalized independently. (b) Shows the application of CP10K which uses the sum of counts across genes for normalization. (c) Depicts the transformation from Biology Space to Measurement Space, then to Normalization Space. (d) Demonstrates between-sample normalization comparing cells to a reference cell. (e) Displays the DESeq2 and SCRAN method where the median of ratio is calculated for normalization. (f) The schema of between-sample normalization and the design of  $\Omega_{i,r} = \omega_i \circ \omega_r^{-1}$ .

account for systematic variations. This paradigm is utilized in methodologies such as EdgeR, DESeq2, and SCRAN. The schema of between-sample normalization in our framework is illustrated in Fig 3.3(d-f). Here, we consider the forms of the measurement functions ( $\hat{\phi}_i, \hat{\phi}_r$ ), and then invert them to derive a composite function  $\Omega_{i,r} = \omega_i \circ \omega_r^{-1}$ . This function maps the counts from a given cell  $\mathbf{y}_i$  to those of a reference cell  $\mathbf{y}_r$ . If we posit that the observed values  $\mathbf{y}$  are related to the true values  $\mathbf{x}$  by an exponential transformation  $\mathbf{y} = A\mathbf{x}^B$ , we can then estimate the relationship  $\mathbf{y}_r = \lambda\mathbf{y}_i^k$  for unknown parameters  $\lambda$  and  $k$ . This allows us to optimize the empirical risk:

$$\arg \min \sum_{g=1}^G \|\Omega_{i,r}(y_{i,g}) - y_{r,g}\|^2 \quad (1)$$

where  $\Omega_{i,r}(\mathbf{y}_i) = \lambda\mathbf{y}_i^k$  and  $\mathbf{y}_i = (y_{i,1}, \dots, y_{i,G})$ .

Optimization of the above function account for instrument-related bias and achieve between-sample normalization. To this end, we transform the measurements in Simplex Space to Euclidean space by composed normalization function  $\Omega$ . To further address property (3), establishing appropriate comparison between two or more cells. Here, we concentrate on the problem that by biological perturbations such as DE, certain values of the RNA measurements may be out-of-distribution. To approach this problem, we propose optimizing the empirical risk for an appropriate subset of the data, assumed to be in-distribution, as discussed in 4.2.

### 3.4. Assumptions of measurement functions

Our framework provides a conceptual model for adjusting the measurement function  $\hat{\phi}$  based on underlying assumptions about the measurement process. In size-factor-

based normalization, commonly used in methodologies like CP10K, DESeq2, EdgeR, and Scran, several critical assumptions underpin the measurement function.

**Assumption 1:** A single scaling factor is sufficient to model the measurement function.

**Assumption 2:** The measurement function is cell-specific.

**Assumption 3:** The majority of genes within each cell share a consistent measurement function.

**Assumption 4:** The measurement function is expected to be monotonic, meaning that increases in input lead to increases in output without any reversal.

CP10K use the assumption (1) and (2) so that the sum of reads is used as the estimator of the size factor. DESeq2 and EdgeR take (3) with (1) and (2) to have a more robust estimation of the size factor. Here we outline the construction of normalization function and more detailed analysis can be found in Appendix. By applying **Assumption 1**, we have  $Y = \hat{\phi}(X) = sX$ . Based on **Assumption 2**, we can construct  $\Omega$  as

$$\Omega(\mathbf{y}_i)_{i,r} = \omega_r^{-1}(\omega_i(\mathbf{y}_i)) = \frac{\mathbf{y}_i}{s_i/s_r}. \quad (2)$$

The reference cell can be the geometry mean of the data (DESeq2) or just any other cell (SCRAN). Applying **Assumption 3**, we have

$$\mathbb{E}\left[\frac{s_i}{s_r}\right] = \mathbb{E}\left[\frac{\sum \mathbf{y}_i}{\sum \mathbf{y}_r}\right] \propto \mathbb{E}\left[\frac{y_{g,i}}{y_{g,r}}\right]. \quad (3)$$

DESeq2, EdgeR, and SCRAN each employ unique strategies to robustly estimate  $\mathbb{E}\left[\frac{y_{g,i}}{y_{g,r}}\right]$ . To this end, we incorporate

these between-sample normalization into our normalization theory framework.

Questioning [Assumption 1](#), we recognize that many instruments exhibit lower detection thresholds and upper saturation limits.<sup>1</sup> We propose that a sigmoidal function may offer a more realistic approximation of the NGS measurement function, especially in scRNA-seq where experimental dropouts often result in zero counts for low input values and the total number of sequences caps the high values. Consequently,  $\phi$  is expected to behave as a piece-wise linear function, registering zero up to a critical lower limit and plateauing at a maximum number beyond an upper saturation limit. In this model, only the input values within a certain “useful range” lead to significant output changes. Within this range, the transfer function is effectively invertible and mimics the scaling transformation typically assumed in size-factor-based normalization. This piece-wise linear behavior can be effectively modeled by a sigmoid function, suggesting that [Assumption 1](#) should be generalized to reflect more complex, realistic behaviors.

It is also crucial to note that all instruments and normalization methods generally assume that the functional relationships are monotonic. Non-monotonic transformations, such as gene-level regularization in SCTransform’s GLM or quantile scaling in scNorm, can disrupt this order, potentially leading to unwanted variations ([Booeshaghi et al., 2022](#)).

## 4. RFNorm: Resistant Fit Normalization

### 4.1. Normalization function

In RFNorm, we consider two hypotheses about the measurement function  $\phi$ : a zero-intercept linear scaling function implemented as RFNorm-L using a linear model through the origin, and a non-linear sigmoid function modeled using the Hill function, designated as RFNorm-NL.

For the nonlinear case, the corresponding normalization function  $\omega_{\text{hill}}$ <sup>2</sup> for the non-linear model is formulated as:

$$\tilde{\phi}(\mathbf{x}) = \frac{a\mathbf{x}^n}{b^n + \mathbf{x}^n}, \quad \omega_{\text{hill}}(\mathbf{y}) = \tilde{\phi}^{-1}(\mathbf{y}) = \left(\frac{b^n \mathbf{y}}{a - b}\right)^{\frac{1}{n}}. \quad (4)$$

Here,  $a$  represents the upper bound of the function,  $b$  describes the  $x$  value at which the response is half-maximal, and  $n$  is the Hill coefficient, indicating the degree of cooperativity.

As a between-sample normalization approach, a mapping

<sup>1</sup>For example, consider the input-output relationship of a light sensor that receives photons of a specific wavelength and generates current.

<sup>2</sup>We analysis three types of common sigmoid functions Hill function, Geompertz function, and Logistic function. Refer to [Appendix A](#) for details.

function  $\Omega$  for every cell  $i$  and reference cell  $r$  can be constructed as:

$$\Omega_{i,r}(\mathbf{y}_i) = \omega_{\text{hill},r}^{-1}(\omega_{\text{hill},i}(\mathbf{y}_i)) = \frac{\mathbf{y}_i}{\beta \mathbf{y}_i + \alpha} \quad (5)$$

where  $\alpha$  and  $\beta$  are reparameterized constants detailed in the [Appendix A.1](#). When  $\beta = 0$ ,  $\Omega$  simplifies to a ratio scaling normalization method.

### 4.2. Resistant Fit for Robust Optimization

Conventionally, parameters in  $\Omega$  are estimated by minimizing the empirical risk, as described in [Eq 1](#). However, this approach is vulnerable to outliers or extreme values, necessitating a robust optimization strategy ([Diakonikolas et al., 2021](#); [Zelditch et al., 2012](#)). Previous methods such as DESeq2 and Scran use median of ratio ([Eq 3](#)) as a form of robust estimate to guard against out-of-distribution values like highly expressed genes and technical zeros. Nevertheless, median discards informative values that are in-distribution and also do not follow a risk-minimization principle. To address this, we propose using a “resistant fit” risk minimization strategy:

$$\arg \min \sum_g^{|B|} \|\Omega_{i,r}(y_{i,g}) - y_{r,g}\|^2 \quad (6)$$

where  $|B|$  is a subset of features that is smaller than  $G$ , the total number of genes. We designate  $B$  as the biological feature set and hypothesize that it contains in-distribution points (features) when mapping one cell to another by [Assumption 3](#). Ideally, we would evaluate all subsets  $B$  such that  $|B|/G = p$  for some fixed proportion  $p$ <sup>3</sup>. However, exploring all  $\binom{G}{|B|}$  possible subsets relative to the reference cell for every cell is computationally prohibitive. Therefore, we implemented an iterative heuristic solution, as detailed in ([Rohlf & Slice, 1990](#)) and ([Zelditch et al., 2012](#)), within the `ResistantFit` function. Initially,  $\Omega_{i,r}$  is fitted to a selected feature set seeded by `InitialFeatureSelection` function. The fitted model is then applied to all gene pairs  $(y_{i,g}, y_{r,g})$ . These pairs are ranked in ascending order based on their residuals relative to  $\hat{\Omega}_{i,r}$ . Only the top-ranked  $|B|$  points are used in the subsequent iteration to fit  $\Omega_{i,r}$ . This iterative process continues until the change in deviations falls below a predefined stopping criterion, as depicted in [Algorithm 1](#). This process iteratively refines the normalization by fitting the resistant fit regression between every cell and the reference cell, thus normalizing the entire dataset.

<sup>3</sup>For example, following the precedent set by EdgeR, which trims 30% of M-values, one might consider  $|B|/G = 0.7$ .

**Algorithm 1** RFNorm Algorithm

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1: Input:  $(\mathbf{y}_1, \mathbf{y}_2, \dots, \mathbf{y}_n)$ 
2:  $\mathbf{y}_r \leftarrow \text{CreateReference}(\mathbf{y}_1, \mathbf{y}_2, \dots, \mathbf{y}_n)$ 
3: for  $i = 1$  to  $n$  do
4:    $\text{ifs} \leftarrow \text{InitialFeatureSelection}(\mathbf{y}_i)$ 
5:    $\mathbf{y}'_i \leftarrow \text{ResistantFit}(\mathbf{y}_i, \mathbf{y}_r, \text{ifs})$ 
6: end for
7: Output:  $(\mathbf{y}'_1, \mathbf{y}'_2, \dots, \mathbf{y}'_n)$ 

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**4.3. Initialization**

In accordance with our [Assumption 4](#) that measurement instruments should preserve monotonic input-output relationships, we initialize the biological feature set  $B$  with a maximally monotonic subset. This strategy involves selecting genes that exhibit a consistent order of expression levels across different cells, effectively reducing noise in the data. This preselection serves as a preliminary data cleaning step, mitigating the risk of normalization artifacts.

We implement this order-preserving strategy by identifying the longest common subsequence (LCS) of genes that display continuous expression between any two cells, using dynamic programming. This ensures that only genes with stable and predictable behavior are included in the initial set  $B$ , which is crucial for the effective application of the normalization process.

**5. Experiments on Simulation****5.1. Compositional Bias**

We adapted the experimental setups from ([L. Lun et al., 2016](#); [Evans et al., 2018](#)) to demonstrate compositional bias in scRNA-seq datasets and to illustrate that within-sample normalization methods such as CP10K are incapable of adequately correcting this bias.

We defined two subpopulations,  $s = \{A, B\}$ , representing different cell types or biological conditions, such as drug treatment or environmental stimulation. For each gene  $g$  in cell  $j$ , the true counts  $x_{gj}$  were sampled from a negative binomial (NB) distribution with mean  $\lambda_s$ . The factor  $\theta_i$  represents cell-specific effects during measurement, such as capture efficiency. The term  $\sum_g \mathbf{x}_i$  in the denominator accounts for the sampling nature of the NGS procedure and is the source of compositional bias and Simplex relations. Here, we simplify the measurement effect as a scaling factor  $\theta_i$ , but we discuss its generalization in sections 3.4.

We assumed each cell has 10,000 genes by the following setup. Parameters  $(\mu_g, \varphi_g, \varphi_l, \mu_l, \varphi_l, \mu_d, \varphi_d)$  are estimated by Splatter ([Zappia et al., 2017](#)) based on ‘‘33k PBMCs from a Healthy Donor, v1 Chemistry’’ from 10x Genomics.

The variable  $\gamma_{is}$  represents the group- and gene-specific

fold-changes, which indicate changes in some genes’ expression levels due to the biological conditions. Differential Expression (DE) involves variations in gene expression levels between different conditions or cell types. Genes are classified as differentially expressed based on their fold-changes relative to control conditions, modeled by  $\gamma_{gs}$  as follows:

$$\begin{aligned} \log_2 \gamma_{gs}(\text{regular}) &= 1.2 \\ \gamma_{gs}(\text{extreme}) &= 10 \\ \gamma_{gs}(\text{non-DE}) &= 1 \end{aligned}$$

In group  $A$ , 10% of the genes were randomly chosen as DE genes. In group  $B$ , 25% were selected as DE genes. Among the DE genes, 85% are regular DEs and 15% are set as extreme responses with 10 times fold-changes. The rest are non-DEs. To this end, the simulation setup can be write as follows.

$$\begin{aligned} \lambda_{g0} &\sim \text{Gamma}(\mu_g, \varphi_g) \\ \lambda_{gs} &= \gamma_{gs} \lambda_{g0} \\ \mathbf{x}_i &\sim \text{NB}(\lambda_s, \varphi_g) \\ \mathbf{y}_i &= \theta_i \mathbf{x}_i / \sum_g \mathbf{x}_i \\ \log_2 \theta_j &\sim \mathcal{N}(\mu_l, \varphi_l) \end{aligned}$$

We simulate three sets of cells  $A1$ ,  $A2$ , and  $B$ .  $A1$  and  $A2$  are replicates of Group  $A$ . We then form two sets of cell pairs. Set  $A/A$  has 100 pair of cells from group  $A1$  and  $A2$ . Set  $A/B$  has 100 pair of cells from group  $A$  and  $B$ . We normalize each group by CP10K. For each gene, we calculate the ratio of expression for each pair of cell. For each non-DE genes, the ratios of groups are shown in the violin plot. We show the violin plot of one gene in Fig 4 (b). Ideally, the same non-DE genes at the groups should have same expression levels. But in Fig 1, we see that the ratio of non-DE genes in group  $A/B$  is higher than group  $A/A$  with  $p\text{-val} < 0.05$ .

**5.2. Outliers**

We employed the same experimental setup as outlined in Section 5.1, with the exception of removing the compositional bias component  $\sum_g \mathbf{x}_i$  to focus exclusively on the impact of outliers when fitting composed normalization function  $\Omega$  between any two pair of cells. So the outliers would be the pairs of genes  $(y_{g,i}, y_{g,j})$ , which are DE genes. Figure 4(c) displays a dot plot of gene expression from two simulated cells, where Cell 1 is from group  $A$  and Cell 2 is from group  $B$ . This experiment was repeated with 100 pairs of cells, and the expression ratios are depicted as a box plot in Figure 4(d).

In this scenario, the ratio of size factors  $\theta_i$  serve as the ground truth for the relationship between non-DE genes

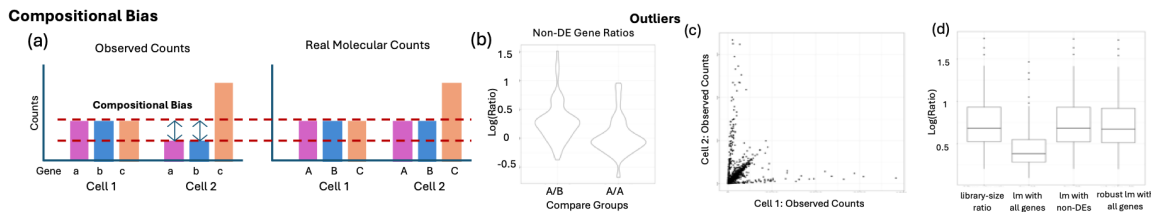


Figure 4. Compositional Bias and Outlier Effects in scRNA-seq Normalization. (a) illustrates the observed counts for three genes in two cells, highlighting the discrepancy between observed and real molecular counts due to compositional bias. (b) presents the distribution of log fold-changes for a non-DE gene between two comparison groups. (c) is a scatter plot illustrating the relationship between observed counts with outliers. (d) shows log fold-change ratios for linear regression and robust linear regression.

across two cells. Estimating this relationship solely based on non-DE genes allows us to accurately recover the true log-ratios. However, fitting a linear model across all genes typically leads to an underestimation of the log fold-change due to the influence of outliers. By employing a robust linear model, such as RFNorm-linear, we successfully align the log-fold-change ratio with the ground truth, demonstrating the efficacy of robust modeling techniques in managing outliers and ensuring accurate data interpretation.

## 6. Apply on Real Data

### 6.1. Invariance of KNN Graph

The efficacy of a preprocessing approach relies on the downstream analysis tasks. The preservation of lower-dimensional mathematical structures, particularly k-nearest neighbor (KNN) graph, is critical in studying various cell types and states (Ahlmann-Eltze & Huber, 2023; Xia et al., 2023). We adopted the Consistent Benchmark from (Ahlmann-Eltze & Huber, 2023) to evaluate RFNorm alongside the other two common used depth normalization methods, CP10K and Scran by comparing the overlap of edges in the KNN graphs post-normalization, as shown in Table 1. It is notable that CP10K might show an inherently higher overlap since count-per-million (CPM) is included in the 22 transformations reviewed in Consistent Benchmark.

The benchmark encompasses three deeply sequenced datasets using msSCRb and Smart-seq3 technologies (Bag-noli et al., 2018; Larsson et al., 2021; Johnsson et al., 2022), each with an average sequencing depth above 50,000 UMIs per cell. A consensus k-NN graph ( $k = 50$ ) is constructed from these datasets after applying 22 transformations, based on the top-50 principal components (PCs) as documented in (Ahlmann-Eltze & Huber, 2023) and Table 1, Common column. To emulate the sequencing depth typical of 10x Genomics, the datasets were downsampled to 5,000 counts, with the results presented in the Reduced rows of Table 1.

RFNorm demonstrates a consistently high overlap in k-NN graphs across all datasets, effectively preserving biological

relationships after normalization. This is evident in both full and reduced datasets, indicating RFNorm’s capability to accurately fit the composed transfer function  $\Omega$ . In full datasets, RFNorm-L and RFNorm-NL yielded results comparable to Scran. Particularly in smaller datasets, such as those from mESC cells, RFNorm-NL matched or even surpassed the performance of CP10K, approaching what is considered ground truth. In reduced datasets, RFNorm generally outperformed both CP10K and Scran in the Fibroblasts dataset, with RFNorm-NL showing superior effectiveness over RFNorm-L. However, in the mESC dataset, Scran proved to be the most effective, likely due to its ability to better estimate size factors in smaller, more distinctly clustered cell populations. In contrast, the lack of clear clustering in the Fibroblasts dataset may reduce Scran’s effectiveness, as detailed in Appendix Fig. 8. Nonlinear models like RFNorm-NL consistently provided better results than linear models, emphasizing their advantage in handling complex biological data. The siRNA KD dataset, with its larger cell count, showcased the scalability of these normalization methods, with RFNorm maintaining high graph overlap even when data formats were reduced.

### 6.2. Marker Gene Identification Validation

To assess the effectiveness of normalization in identifying marker genes, we utilized scRNA-seq data from patients with pulmonary fibrosis and healthy controls, previously analyzed by (Reyfman et al., 2019). From (Squair et al., 2021), marker genes identified in bulk RNA-seq were used as benchmarks to evaluate the accuracy of detecting cell-type-specific markers in scRNA-seq data. We processed subsets of 750, 1000, and 1500 cells from each group using CP10K, Scran, RFNorm-L, and RFNorm-NL, with comprehensive results detailed in Appendix D.

Furthermore, to confirm that the normalization methods accurately control for false positives, we conducted a Type 1 error control test using a “null dataset” derived from “16-cell stage blastomeres” (Deng et al., 2014), following the protocol described by (Soneson & Robinson, 2018). This

Table 1. Comparison of k-Nearest Neighbor (k-NN) Graph Overlap across Different Normalization Methods. The k-NN graphs (k=50) were built based on the top-50 principal components for each normalization method. The downsampled experiments were repeated 5 times.

Dataset	N Cells		Number of Overlapped Edges in KNNs				
			Common	CP10K	SCRAN	RFNorm-L	RFNorm-NL
mESC (mcSCRB)	249	Full	1085	972	941	942	970
		Reduced		782(9.25)	<b>873</b> (10.4)	803(20.33)	812(37.35)
Fibroblasts (ss3)	369	Full	1365	1342	1220	1260	1257
		Reduced		603(22.39)	517(12.88)	767(26.48)	<b>835</b> (35.91)
siRNA KD (ss3)	4298	Full	7194	7187	7186	7189	7187
		Reduced		6832.9(19.26)	6854.5(22.75)	<b>6958.5</b> (18.28)	6499 (29.39)

experiment aimed to verify that the methods do not erroneously identify genes as differentially expressed under controlled conditions. Findings from this test are detailed in Appendix E.

Both sets of analysis demonstrated the comparable performance of RFNorm in maintaining accuracy and managing Type 1 error rates in the detection of biologically relevant markers in scRNA-seq data.

### 6.3. Monotonicity

An effective normalization technique must preserve the data structure by ensuring monotonicity across cells (Booeshaghi et al., 2022). To evaluate how well different normalization methods maintain this property, we employed Spearman’s rank correlation to measure the extent of monotonicity in cellular data from the Fibroblast Dataset (Hagemann-Jensen et al., 2020), both before and after normalization. This metric helps in identifying any potential non-monotonic alterations introduced by normalization techniques. As shown in Figure 6.3, the analysis reveals that techniques such as SCTransform’s GLM regularization and SCNorm’s Quantile scaling may negatively impact the preservation of monotonicity in data structure.

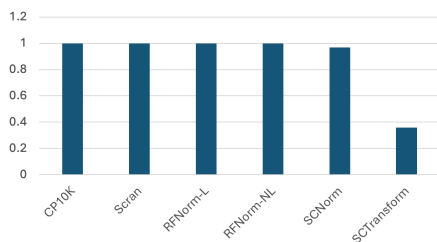


Figure 5. Analysis of Spearman’s rank correlation to assess monotonicity preservation in different normalization methods. Lower values indicate greater disruption of data structure.

## 7. Discussion

Existing literature shows a large number of different methods that have been proposed for normalizing transcriptome data, including those from single cells. However, the methods do not explicitly consider learning the optimal function under an explicit objective function, especially with inclusion of consideration for aligning “comparables” (i.e., gene sets that may be considered samples from a common distribution). Here we propose a new framework that explicitly these issues.

In our study, we established a benchmark to evaluate the effectiveness of depth normalization methods in preserving the structure of k-NN graphs. RFNorm methods consistently showed enhanced performance in maintaining biological relationships across varying complexities and sequencing depths. RFNorm-NL was more proficient in smaller cell populations and in datasets with reduced sequencing depth, whereas Scran showed its strength primarily in the mESC dataset characterized by a limited number of cells and distinct clusters. In the context of larger and more intricate datasets, such as the siRNA KD dataset, a noticeable performance dip in RFNorm-NL within the reduced dataset was observed, suggesting potential areas for refinement in managing large-scale data. Further investigation is warranted to determine how optimal reference cell selection during optimization can minimize information loss in such complex datasets.

Normalization is surprisingly difficult with many subtle issues in practice (e.g., handling abundance of zero observations). Here, we suggest that approaching the problem from the point of view of the natural of the measurement function, optimization of appropriate empirical risk function, and alignment of comparable data (i.e., accounting for out-of-distribution values), provides a broad framework to construct principled methods for normalization.



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## A. Sigmoid Functions

We explored three types of common sigmoid functions. They are Hill function, Gompertz Function, and Logistic function. The corresponding normalization functions are listed in Table 2.

Table 2. Measurement and Normalization transfer functions for different sigmoid functions

Functions	Measurement $\phi(x)$	Normalization $\omega(y)$	Composed Function $\Omega_{i,r}(y_i)$
Hill	$\frac{ax^n}{b^n+x^n}$	$\left(\frac{b^ny}{a-b}\right)^{\frac{1}{n}}$	$\frac{y_i}{\beta y_i + \alpha}$
Geompertz	$L \exp(-b \exp(-kx))$	$-\frac{1}{k} \log\left(\frac{1}{b} \log \frac{L}{y}\right)$	$\frac{L_r}{L_i^\beta} y_i^\beta$
Logistics	$\frac{a}{1+be^{tx}}$	$\frac{1}{t} \left(\log\left(\frac{a}{y} - 1\right) - \log b\right)$	$\frac{y_i}{\beta y_i + \alpha}$

Hill function is commonly used in biochemistry and systems biology to describe the response of a system to a stimulus, typically in terms of the concentration of a substrate. It is especially used to describe cooperative binding of substrates to enzymes or receptors. As discussed in Section 4.1, for scRNA-seq, there are more constrains on the lower end of the range while the observation may fall within the higher-end of the operational range most of the time. So a function with flexible lower end and a long operational range is preferable. As demonstrated in Fig 6, Hill function has the best flexibility and dynamic range to represent sequencing measurement. This is a qualitative check and needs more exploration to make strict conclusion. In table 2, we reported the composed transfer function based on different sigmoid functions after reparameterization. Hill function and logistic function are reduced to the equivalent format.

### A.1. Hill function and Composed normalization function

Set Hill function  $\tilde{\phi}$  as the estimated measurement function, we have

$$\tilde{\phi}(x) = \frac{ax^n}{b^n + x^n} \quad (7)$$

Here  $a$  is the upper bound of the function,  $b$  describes the  $x$  whose response is half-maximal, and  $n$  is the Hill coefficient, which describes the degree of cooperativity.

Consequently, the corresponding normalization function  $\omega_{\text{hill}}$  is

$$\omega_{\text{hill}}(y) = \tilde{\phi}^{-1}(y) = \left(\frac{b^ny}{a-b}\right)^{\frac{1}{n}}. \quad (8)$$

As in Fig.7, we construct the composed mapping function  $\Omega$  for every cell  $i$  and  $r$ . The expression of  $\Omega$  is given by:

$$\Omega_{i,r}(y) = \omega_{\text{hill},r}^{-1}(\omega_{\text{hill},i}(y)) \quad (9)$$

After some reformations (see Appendix A.4), we have

$$\Omega_{i,r}(y_i) = \frac{a_r}{1 + \exp\left[\frac{n_r}{n_i} \log \frac{a_i - y_i}{y_i} - n_r \log \frac{b_r}{b_i}\right]} \quad (10)$$

Set the hill coefficient to be the same, i.e.  $n_r = n_i$ , we have

$$\Omega_{i,r}(y_i) = \frac{a_r (b_r/b_i)^{n_r} y_i}{((b_r/b_i)^{n_r} - 1) y_i + a_i} \quad (11)$$

Without loss of generality, we can reparameterize the function as

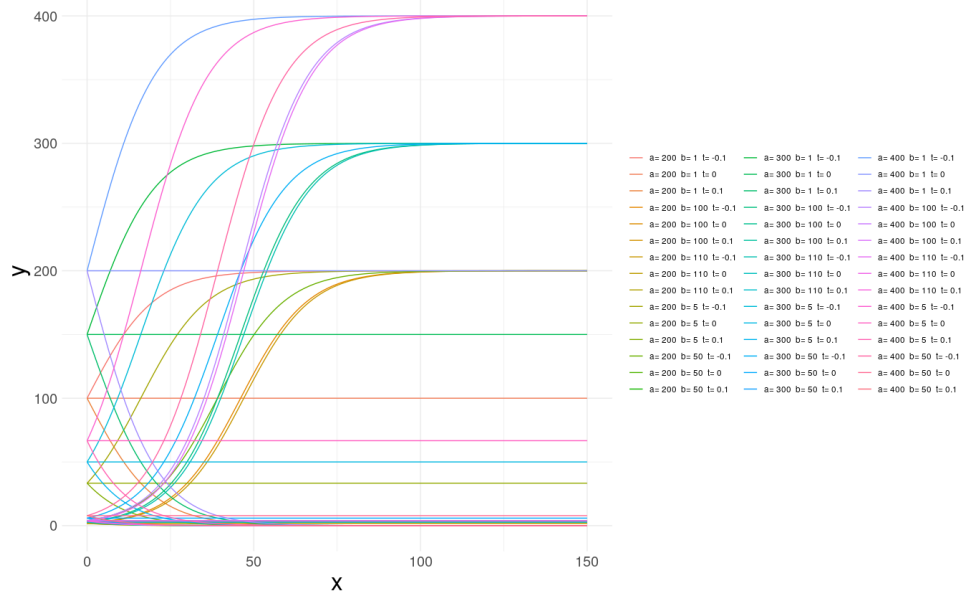
$$\Omega_{i,r}(y_i) = \frac{\lambda_1 y_i}{\lambda_2 y_i + \lambda_3} \quad (12)$$

, where  $\lambda_1 = a_r b_i$ ,  $\lambda_2 = b_i - b_r$ , and  $\lambda_3 = b_r a_i$ . It can be further simplified as

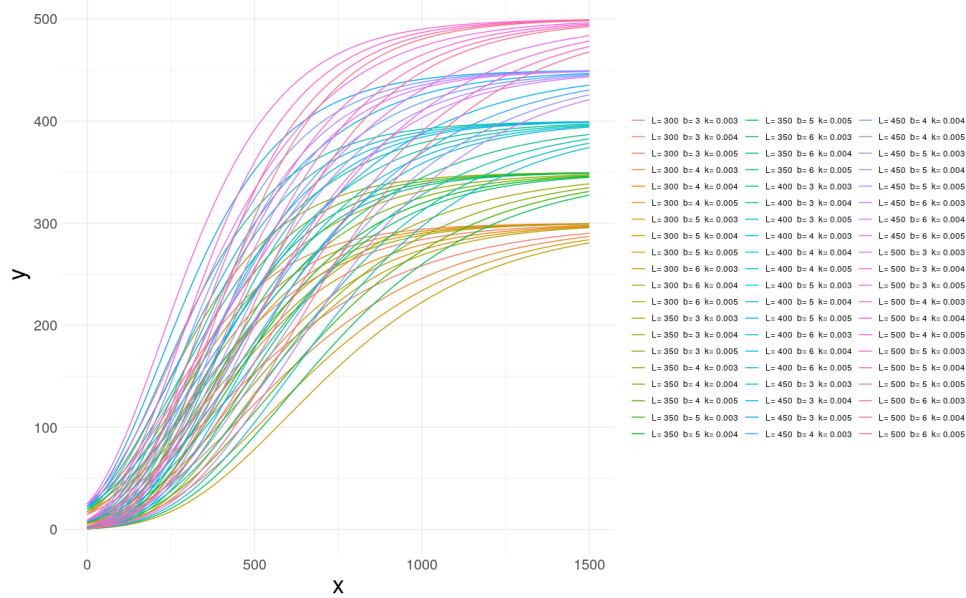
$$\Omega_{i,r}(y_i) = \frac{y_i}{\beta y_i + \alpha} \quad (13)$$

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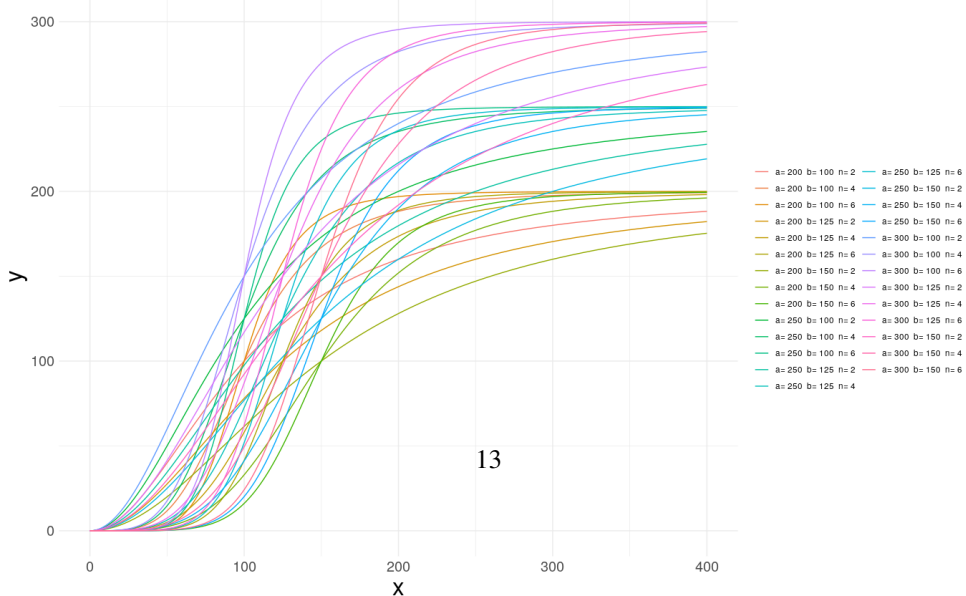
Logistic Function with Different Parameters



Geompertz Function with Different Parameters



Hill Function with Different Parameters



## A.2. Geompertz function and Composed normalization function

Set Geompertz function  $\tilde{\phi}$  as the estimated measurement function, we have

$$\tilde{\phi}(x) = L \exp(-b \exp(-kx)) \quad (14)$$

Here  $L$  is the upper bound of the function,  $b$  describes the horizontal shift, and  $k$  affects the growth rate. Consequently, the corresponding normalization function  $\omega_{\text{geompertz}}$  is

$$\omega_{\text{geompertz}}(y) = \tilde{\phi}^{-1}(y) = -\frac{1}{k} \log\left(\frac{1}{b} \log \frac{L}{y}\right). \quad (15)$$

We construct the composed mapping function  $\Omega$  for every cell  $i$  and  $r$ . The expression of  $\Omega$  is given by:

$$\Omega_{i,r}(y) = \omega_{\text{geompertz},r}^{-1}(\omega_{\text{geompertz},i}(y)) \quad (16)$$

Suppose growth rate  $k$  is non-cell specific,  $k_r = k_i$  and  $Set\beta = \frac{b_i}{b_r}$ . After some reformations, we have

$$\Omega_{i,r}(y_i) = \frac{L_r}{L_i^\beta} y_i^\beta \quad (17)$$

$$\log \Omega_{i,r}(y_i) = \beta \log y_i + \log L_r - \beta \log L_i \quad (18)$$

## A.3. Logistic function and Composed normalization function

Set Logistic function  $\tilde{\phi}$  as the estimated measurement function, we have

$$\tilde{\phi}(x) = \frac{a}{1 + be^{tx}} \quad (19)$$

Here  $a$  is the upper bound of the function,  $b$  describes the horizontal shift, and  $t$  is the growth rate.

Consequently, the corresponding normalization function  $\omega_{\text{logistic}}$  is

$$\omega_{\text{logistic}}(y) = \tilde{\phi}^{-1}(y) = \frac{1}{t} \left( \ln\left(\frac{a}{y} - 1\right) - \ln b \right) \quad (20)$$

We construct the composed mapping function  $\Omega$  for every cell  $i$  and  $r$ . The expression of  $\Omega$  is given by:

$$\Omega_{i,r}(y) = \omega_{\text{logistic},r}^{-1}(\omega_{\text{logistic},i}(y)) \quad (21)$$

Suppose  $\lambda = \frac{t_r}{t_i}$ , after some reformations (see Appendix A.5), we have

$$\Omega_{i,r}(y_i) = \frac{y_i^\lambda}{\frac{b_r}{a_r b_i^\lambda} (a_i - y_i)^\lambda + \frac{1}{a_r} y_i^\lambda} \quad (22)$$

Set the growth rate to be the same, i.e.  $t_r = t_i$  and  $\lambda = 1$ , we have

$$\Omega_{i,r}(y_i) = \frac{y_i}{\frac{a_i b_r}{a_r b_i} + \frac{b_r + b_i}{a_r b_i} y_i} \quad (23)$$

Without loss of generality, we can reparameterize the function as

$$\Omega_{i,r}(y_i) = \frac{y_i}{\beta y_i + \alpha} \quad (24)$$

**A.4. Reformation of composed Hill functions**

$$\begin{aligned}
 \phi_r(y_r) &= \phi_i(y_i) \\
 \left(\frac{b_r^{n_r} y_r}{a_r - y_r}\right)^{\frac{1}{n_r}} &= \left(\frac{b_i^{n_i} y_i}{a_i - y_i}\right)^{\frac{1}{n_i}} \\
 \frac{1}{n_r} \log\left(\frac{b_r^{n_r} y_r}{a_r - y_r}\right) &= \frac{1}{n_i} \log\left(\frac{b_i^{n_i} y_i}{a_i - y_i}\right) \\
 n_r \log b_r - \log \frac{a_r - y_r}{y_r} &= \frac{n_r}{n_i} \log \frac{b_i^{n_i} y_i}{a_i - y_i} \\
 \log\left(\frac{a_r}{y_r} - 1\right) &= n_r \log b_r + \frac{n_r}{n_i} \log \frac{a_i - y_i}{b_i^{n_i} y_i} \\
 y_r &= \frac{a_r}{1 + \exp[n_r \log b_r + \frac{n_r}{n_i} \log \frac{a_i - y_i}{b_i^{n_i} y_i}]} \\
 y_r &= \frac{a_r}{1 + \exp[\frac{n_r}{n_i} \log \frac{a_i - y_i}{y_i} - n_r \log \frac{b_r}{b_i}]}
 \end{aligned}$$

**A.5. Reformation of composed logistic functions**

$$\begin{aligned}
 \omega_r(y_r) &= \omega_i(y_i) \\
 \frac{1}{t_r} \left( \ln\left(\frac{a_r}{y_r} - 1\right) - \ln b_r \right) &= \frac{1}{t_i} \left( \ln\left(\frac{a_i}{y_i} - 1\right) - \ln b_i \right)
 \end{aligned}$$

Suppose  $\lambda_1 = \frac{t_r}{t_i}$ ,

$$\begin{aligned}
 \ln\left(\frac{a_r - y_r}{b_r y_r}\right) &= \lambda_1 \ln\left(\frac{a_i - y_i}{b_i y_i}\right) \\
 \frac{a_r}{b_r y_r} - \frac{1}{b_r} &= \frac{(a_i - y_i)^{\lambda_1}}{b_i^{\lambda_1} y_i^{\lambda_1}} \\
 \frac{a_r}{y_r} - \frac{1}{y_r} &= \frac{b_r (a_i - y_i)^{\lambda_1}}{b_i^{\lambda_1} y_i^{\lambda_1}} + \frac{b_i^{\lambda_1} y_i^{\lambda_1}}{b_i^{\lambda_1} y_i^{\lambda_1}} \\
 \frac{1}{y_r} &= \frac{b_r (a_i - y_i)^{\lambda_1} + b_i^{\lambda_1} y_i^{\lambda_1}}{a_r b_i^{\lambda_1} y_i^{\lambda_1}} \\
 y_r &= \frac{y_i^{\lambda_1}}{\frac{b_r}{a_r b_i^{\lambda_1}} (a_i - y_i)^{\lambda_1} + \frac{1}{a_r} y_i^{\lambda_1}}
 \end{aligned}$$

Assume  $t_r = t_i$ , i.e.  $\lambda_1 = 1$ . We have

$$\begin{aligned}
\frac{a_r - y_r}{b_r y_r} &= \frac{a_i - y_i}{b_i y_i} \\
\frac{a_r}{b_r y_r} - \frac{1}{b_r} &= \frac{a_i}{b_i y_i} - \frac{1}{b_i} \\
\frac{a_r b_i}{y_r} - b_i &= \frac{a_i b_r}{y_i} - b_r \\
\frac{a_r b_i}{y_r} &= \frac{a_i b_r}{y_i} - b_r + b_i \\
\frac{a_r b_i}{y_r} &= \frac{a_i b_r + (b_r + b_i) y_i}{y_i} \\
y_r &= \frac{a_r b_i y_i}{a_i b_r + (b_r + b_i) y_i} \\
y_r &= \frac{y_i}{\frac{a_i b_r}{a_r b_i} + \frac{b_r + b_i}{a_r b_i} y_i}
\end{aligned}$$

## B. Construction of the reference cell

A reference cell need to be selected as the baseline. Different methods make distinct choices in this regard. For example, DESeq2 employs the geometric mean of the dataset as the reference, EdgeR arbitrarily selects one sample for the role, and Scran utilizes an averaged pseudo-cell.

In our proposed approach, we opt for the median of the dataset to serve as the reference cell, denoted as  $y_r$ . We argue that using the median offers a balance, providing a central tendency that is robust to outliers. Unlike the geometric mean, which can be sensitive to zero values, or an arbitrary selection that could introduce bias, the median offers a more stable and unbiased point of reference.

## C. Invariance of KNN Graph



Figure 7. Schema of KNN Consistent Benchmark

mESC dataset (Bagnoli et al., 2018) consist of 249 single mESCs using SCRB-seq. Fibroblasts (Hagemann-Jensen et al., 2020) has 369 individual primary mouse fibroblasts with Smart-seq3. siRNA KD dataset (Larsson et al., 2021) has 4298 fibroblasts siRNA knockdown cells from CAST/EiJ and C57BL/6J mouse strains (5 animals). The tSNE plots of the transformed dataset are in Fig.8.

## D. Concordance with Bulk Data

We assessed the efficacy of cell-type-specific marker detection using scRNA-seq data from patients with pulmonary fibrosis and healthy controls, initially analyzed by (Reyfman et al., 2019) and (Squair et al., 2021). This scRNA-seq dataset was compared against a bulk RNA-seq experiment, using the list of marker genes identified in the bulk data as a benchmark. We analyzed randomly selected subsets of 750, 1000, and 1500 cells from each group after normalizing the data using CP10K, Scran, RFNorm-L, and RFNorm-NL. Differential expression was evaluated using a t-test, with significant findings defined by adjusted p-values below 0.05. According to our analysis, summarized in Table 3, all four methods demonstrated similar performance in detecting differentially expressed genes.



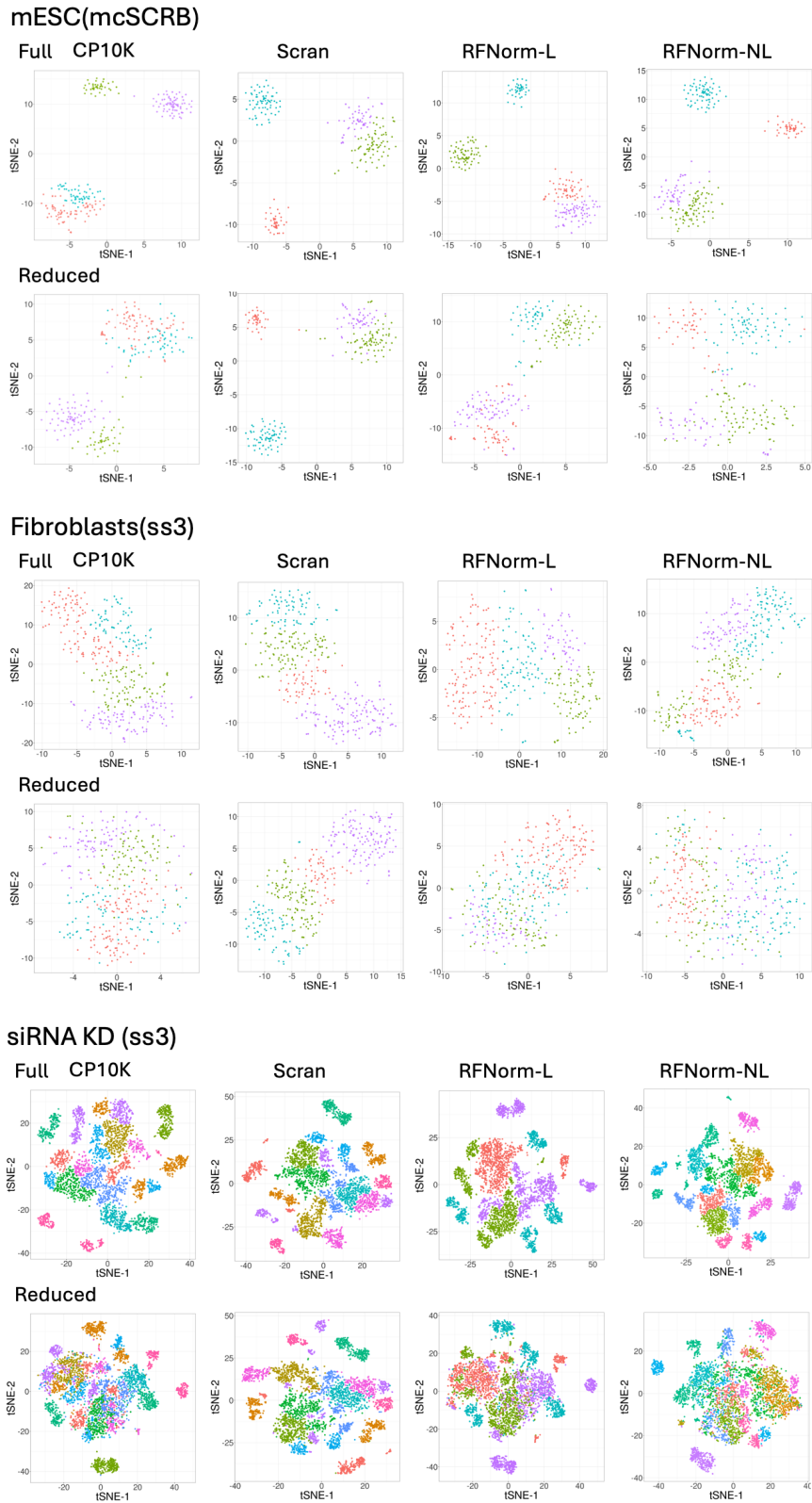


Figure 8. tSNE plots of each dataset after transformations. Cells are colored by clustering using the walktrap clustering algorithm as (Ahlmann-Eltze & Huber, 2023). The reduced dataset is colored by the clusters identified in full dataset.

Table 3. Differential Expression Analysis Results for Various Normalization Methods. Genes with adjusted p-values  $< 0.05$  are considered DE. If the DE is also discovered in the paired bulk-RNA experiment, it is counted as concordance.

N Cells	Type	CP10K	SCRAN	RFNorm-L	RFNorm-NL
1500	# DE	1163.6 (34.98)	1168.4 (24.8)	1206 (34.79)	1166.4 (31.94)
	# Concordance	170.4 (7.19)	169.4 (8.38)	169.4 (7.95)	168.6 (6.58)
1000	# DE	843.6 (50.97)	840.6 (50.48)	884.2 (53.4)	848 (54.55)
	# Concordance	138.4 (11.28)	136.6 (10.11)	139.2 (11.78)	137.6 (11.65)
750	# DE	646.4 (26.14)	648.8 (24.47)	685 (33.41)	653.6 (27.81)
	# Concordance	118.2 (8.22)	117.4 (7.44)	119.6 (7.6)	116 (6.63)

## E. Type 1 Error Control

It is essential that the normalization factors ensure genes with identical expression levels in two different samples are not mistakenly identified as differentially expressed (Robinson & Oshlack, 2010). In our study, we utilized a 'null dataset' comprising 50 cells from the '16-cell stage blastomere', as described in (Deng et al., 2014), following the approach introduced by (Soneson & Robinson, 2018). After applying four different transformations, none of the genes exhibited an adjusted p-value (p-adj) below 0.05, indicating no differential expression. This finding aligns with the observations in (Soneson & Robinson, 2018), where both the t-test and Wilcoxon rank sum test were shown to effectively control Type-1 error across various transformations.