ZERO-INFLATED QUANTILE RANK-SCORE BASED TEST (ZIQRANK) WITH APPLICATION TO SCRNA-SEQ DIFFERENTIAL GENE EXPRESSION ANALYSIS

BY WODAN LING\textsuperscript{1}, WENFEI ZHANG\textsuperscript{2,*} BIN CHENG\textsuperscript{3} AND YING WEI\textsuperscript{3}

\textsuperscript{1}Public Health Sciences Division, Fred Hutchinson Cancer Research Center, wling@fredhutch.org
\textsuperscript{2}Sarepta Therapeutics, *wenfeizhang@gmail.com
\textsuperscript{3}Department of Biostatistics, Columbia University, bc2159@cumc.columbia.edu; yw2148@cumc.columbia.edu

Differential gene expression analysis based on scRNA-seq data is challenging due to two unique characteristics of scRNA-seq data. First, multimodality and other heterogeneity of the gene expression among different cell conditions lead to divergences in the tail events or crossings of the expression distributions. Second, scRNA-seq data generally have a considerable fraction of dropout events, causing zero-inflation in the expression. To account for the first characteristic, existing parametric approaches targeting the mean difference in gene expression are limited, while quantile regression that examines various locations in the distribution will improve the power. However, the second characteristic, zero-inflation, makes the traditional quantile regression invalid and underpowered. We propose a quantile-based test that handles the two characteristics, multimodality and zero-inflation, simultaneously. The proposed quantile rank-score based test for differential distribution detection (ZIQRank) is derived under a two-part quantile regression model for zero-inflated outcomes. It comprises a test in logistic modeling for the zero counts, and a collection of rank-score tests adjusting for zero-inflation at multiple prespecified quantiles of the positive part. The testing decision is based on an aggregate result by combining the marginal p-values by MinP or Cauchy procedure. The proposed test is asymptotically justified and evaluated with simulation studies. It shows a higher precision-recall AUC in detecting true differentially expressed genes (DEGs) than the existing methods. We apply the ZIQRank test to a TPM scRNA-seq data on human glioblastoma tumors, and exclusively identify a group of DEGs between neoplastic and non-neoplastic cells, which are heterogeneous and have been proved to be associated with glioma. Application to a UMI count scRNA-seq data on cells from mouse intestinal organoids further demonstrates the capability of ZIQRank to improve and complement the existing approaches.

1. Introduction. Differential gene expression analysis is one of the most commonly performed tasks for RNA-seq data with a broad set of applications, such as identifying genes associated with a tumor, understanding phenotypic variation, among many others (Costa-Silva, Domingues and Lopes 2017; Zhang et al. 2014). The traditional RNA-seq experiments measure mRNA transcript abundance averaged over thousands or millions of cells, which have proven useful in many studies. However, a gene may express at substantially different levels in different sub-groups of cells, and the bulk experiments fail to take account of this cell-specific information (Korthauer et al. 2016). As a crucial technology advance, single cell RNA-sequencing (scRNA-seq) measures mRNA transcript abundance in individual cells, and enables us to study the gene-specific expression heterogeneity across cells. This is important for understanding cancer progression and discovering novel cell types (Buettner et al. 2015;

Many methods have been proposed to identify differentially expressed genes (DEGs) in scRNA-seq data. Comprehensive reviews can be found in Soneson and Robinson (2018) and Dal Molin, Baruzzo and Di Camillo (2017). A good scRNA-seq analysis tool should be tailored for two unique features of scRNA-seq data. First, due to biological and technical cell-to-cell variability, a given gene’s expression is often unobserved in a large fraction of cells, which leads to zero-inflation in expression level (Kharchenko, Silberstein and Scadden 2014). Several mixture methods have been developed to accommodate the zero-inflated nature of scRNA-seq data, and lead to substantively improved power compared to more traditional bulk RNA-seq models such as the once-popular DESeq2 (Love, Huber and Anders 2014). For example, MAST (Finak et al. 2015) uses a logistic regression model to model the zero-inflation and a Gaussian linear model to model the positive continuous part. Monocle (Trapnell et al. 2014) fits the data with a generalized additive model (GAM) and takes care of the dropout events using the Tobit model. Second, scRNA-seq data is highly heterogeneous due to both the population and cellular heterogeneity. For example, several works (Kærn et al. 2005; Birtwistle et al. 2012; Singer et al. 2014; Dobrzyński et al. 2012, 2014) report multimodal distributions of scRNA-seq gene expressions. This is probably a result of the multiple stable states among expressed genes (Birtwistle et al. 2012) or comes from a series of biological processes. For example, when a tumor suppressor gene is over-expressed in some cells, it will be slightly adjusted back by another biological process in the human body. Such a negative feedback that causes oscillations has been observed in several studies (Kærn et al. 2005; Monk 2003), which may lead to multiple modes in scRNA-seq gene expressions (Dobrzyński et al. 2012, 2014). The hidden heterogeneity in the scRNA-seq data could be more complex than the existence of multiple modes, which is inherited from the bulk RNA-seq level. For example, Song et al. (2017) reports several eQTLs that only affect the upper tails of gene expressions. Although many aforementioned models consider the inflated zero counts, they rely on parametric models, such as Gaussian model, which could be insufficient to capture the heterogeneity in the scRNA-seq data. The scDD method proposed in Korthauer et al. (2016) utilizes a conjugate Dirichlet process mixture (DPM) of Gaussian to handle the hidden heterogeneity, but fails to accommodate more than two cell conditions and cannot incorporate confounding covariates, which are important considerations for gene expression analyses.

In this paper, we propose a two-part quantile regression model, which fully incorporates the zero-inflated and heterogeneous nature of scRNA-seq data while allows the adjustment of covariates. Quantile regression (Koenker and Bassett 1978) models the conditional distribution of an outcome without any parametric likelihood specification, and is hence a promising non-parametric alternative to detect DEGs with complex and heterogeneous associations. It also offers the flexibility to incorporate confounders. Several earlier works using quantile regression to test genetic associations (Song et al. 2017; Wei et al. 2016) report discoveries in both GWAS and eQTL studies. These genetic association tools from direct quantile regression, however, do not account for zero-inflation. The computation algorithm and theoretical results of quantile regression are built upon the assumption that the conditional distribution of outcomes is absolutely continuous. Since the existence of zero-inflation violates the assumption, quantile regression often fails in producing reliable estimations, and inferences by these existing quantile-based tools may not be valid. Moreover, direct quantile regression cannot capture the dependence of zero proportions on the cell condition and covariates, leading to biased estimations. Thus, these direct quantile-based methods may suffer from the biases, resulting in uncontrolled false positives or underpowered testing. As a remedy, Zhang et al. (2020) uses logistic regression for the zero-inflation, and applies quantile regression to the
scRNA-seq data at shifted quantiles with perturbation to break the probability mass at zero. Though perturbation is a convenient numerical treatment, it could introduce extra noises into the analysis. To address the existing challenges, we propose a rank-score test adjusting the bias caused by zero-inflation based on the proposed two-part quantile model, and establish its asymptotic properties. Finally, we detect DEGs by combining the marginal p-values of the logistic regression for zero counts and the novel rank-score tests over a sequence of quantiles of non-zero expressions. We establish the asymptotic dependence structure of these marginal tests and incorporate it when combining the p-values. This proposed differential test is named zero-inflated quantile rank-score based test (ZIQRank).

Details of ZIQRank are presented in Section 2. Simulation studies to compare it with QRank, linear regression, and existing differential expression analysis methods for bulk and scRNA-seq data can be found in Section 3. It is shown that ZIQRank has a well-controlled false positive rate and a higher precision-recall AUC than the competing methods. In Section 4, we apply the proposed method to a non-UMI TPM scRNA-seq data about glioblastoma in the conquer (consistent quantification of external rna-seq data) repository (Soneson and Robinson 2018), and identify a group of genes that are differentially expressed between neoplastic and non-neoplastic cells. Previous biological investigations warrant the roles of those genes in diagnosis, progression and suppression of glioma. In Section 5, we further use ZIQRank to study a UMI count scRNA-seq data about cells from mouse intestinal organoids. We conclude the paper in Section 6. Technical proofs are relegated to Appendix A.

2. Proposed methods.

2.1. Notations in scRNA-seq data analysis. A scRNA-seq data consists of a random sample of \( n \) cells. For each cell, we sequence \( J \) genes. As a result, we have an \( n \times J \) gene expression matrix \( Y \), whose entry \( Y_{i,j} \) denotes the expression level of the \( j \)th gene in the \( i \)th cell. We then denote the primary covariates of interest, cell conditions, as a \( q \)-dimensional vector \( C_i \). The differential gene expression analysis is to identify genes whose cell-level expressions depend on the cell conditions \( C \). In addition, we collect a set of additional cell characteristics including the intercept, e.g., patient from whom a cell is collected, potentially related clinical features and sequencing information, and denote them as a \( p \)-dimensional vector \( Z_i \).

Throughout the paper, we denote \( Q_{Y_{i,j}}(\tau|X_i) \) as the \( \tau \)th conditional quantile of \( Y_{i,j} \) given \( X_i \), where \( X_i = (Z_i^T, C_i^T)^T \).

2.2. Zero-inflated quantile regression model for individual gene expression. We decompose the conditional distribution of the expression \( Y_{i,j} \) as

\[
P(Y_{i,j} \leq y|X_i) = P(Y_{i,j} = 0|X_i) + P(Y_{i,j} \leq y|X_i, Y_{i,j} > 0)P(Y_{i,j} > 0|X_i)I(y > 0),
\]

where \( P(Y_{i,j} = 0|X_i) \) is the probability of no expression of the \( j \)th gene in the \( i \)th cell, while \( P(Y_{i,j} \leq y|X_i, Y_{i,j} > 0) \) is the conditional distribution of expression level given that the \( j \)th gene is expressed in the \( i \)th cell. Following the decomposition, we consider a two-part model for the zero-inflated expression. First, we model \( P(Y_{i,j} > 0|X_i) \) by a logistic regression model,

\[
\text{logit}\{P(Y_{i,j} > 0|X_i)\} = Z_i^T \zeta_j + C_i^T \gamma_j,
\]

where \( \exp(\gamma_j) \) is the odds-ratio for observing non-zero expression of the \( j \)th gene associated with cell conditions \( C \). Next, we model the conditional distribution \( P(Y_{i,j} \leq y|X_i, Y_{i,j} > 0) \) using a linear quantile regression model,

\[
Q_{Y_{i,j}}(\tau|X_i, Y_{i,j} > 0) = Z_i^T \alpha_j(\tau) + C_i^T \beta_j(\tau),
\]
where $\beta_j(\tau)$ depicts how the $\tau$th quantile of non-zero expression of the $j$th gene differs by cell conditions $C$, and $\alpha_j(\tau)$ captures the contribution of the remaining covariates. Due to the non-parametric nature of quantile regression, the proposed two-part quantile model is a generalization of the MAST method. As the model is proposed for analyzing each gene independently, we focus on the $j$th gene in the rest of Section 2, and omit the subscript $j$ in all the notations for a simpler presentation.

The quantile coefficients $\beta(\tau)$ and $\alpha(\tau)$ can be estimated by minimizing the following loss function

$$\min_{\alpha,\beta} \sum_{i=1}^{n} \rho_\tau(Y_i - Z_i^\top\alpha - C_i^\top\beta)I(Y_i > 0),$$

where $\rho_\tau(u) = u\{\tau - I(u < 0)\}$ is the quantile loss function. Due to the correlation between the indicator $I(Y_i > 0)$ and the loss function $\rho_\tau(Y_i - Z_i^\top\alpha - C_i^\top\beta)$, the existing quantile regression inference tools cannot be applied directly. In addition, those inference tools, if applied directly, will under-estimate the uncertainty that the “positive subset” is observed by chance, and lead to biased tests. In the subsequent section, we adapt the rank-score test (Gutenbrunner et al. 1993) to the proposed zero-inflated quantile model. The rank-score test is robust and computationally efficient. Hence, it is the desired inference tool for genetic associations. In the case of no zero-inflation, Song et al. (2017) has numerically shown that the traditional rank-score test has a well-controlled Type I error on the GTEx multi-tissue gene expression data.

2.3. Rank-score test of $\beta(\tau)$ with zero-inflation. Let $\tilde{C}_i = C_i \cdot I(Y_i > 0)$ and $\tilde{Z}_i = Z_i \cdot I(Y_i > 0)$ as the nominal variables of the cell conditions and remaining covariates. It follows that $\tilde{C}_{n \times q} = (\tilde{C}_1, \cdots, \tilde{C}_n)^\top$, $\tilde{Z}_{n \times p} = (\tilde{Z}_1, \cdots, \tilde{Z}_n)^\top$ are the design matrix associated with $C_i$’s and $Z_i$’s. We further denote $C^* = (I - \tilde{Z}(\tilde{Z}^\top\tilde{Z})^{-1}\tilde{Z}^\top)\tilde{C}$, where $I$ is the $n \times n$ identity matrix. This orthogonal transformation ensures the asymptotic independence between $C^*$ and $\tilde{Z}$.

We construct a rank score for $\beta(\tau) = 0$ by

$$S_{n,\tau}^Q = n^{-\frac{1}{2}} \sum_{i=1}^{n} \psi_\tau(Y_i - \tilde{Z}_i^\top\tilde{\alpha}_n(\tau))I(Y_i > 0)\tilde{C}_i^*, \tag{4}$$

where $\psi_\tau(u) = \tau - I(u < 0)$ is the piece-wise first derivative of the quantile loss function $\rho_\tau(u)$, $\tilde{\alpha}_n(\tau)$ is the minimizer of (3) with $\beta = 0$, and $\tilde{C}_i^*$ is the $i$th row of $C^*$. By design, $S_{n,\tau}^Q$ measures the independent contribution of $C$ onto the $\tau$th quantile of $Y$. When $\beta(\tau) = 0$, $S_{n,\tau}^Q$ is close to a vector of zeros. We also note that the zero-positive uncertainty is incorporated into the rank-score (4).

Finally, with $V_{n,\tau} = n^{-1}(1 - \tau)C^*\tilde{C}^*$, we define the rank-score test statistic at the $\tau$th quantile as

$$T_{\tau}^Q = S_{n,\tau}^{Q\top} V_{n,\tau}^{-1} S_{n,\tau}^Q. \tag{5}$$

Under the conditions outlined in Appendix A.1, we have established the asymptotic distribution of $T_{\tau}^Q$ in the following theorem.

**THEOREM 1.** Under Assumptions 1-4 in Appendix A.1, as $n \to \infty$, we have

(a) At a fixed $\tau$, given $\beta(\tau) = 0$, define

$$\hat{S}_{n,\tau}^Q = n^{-\frac{1}{2}} \sum_{i=1}^{n} \psi_\tau(Y_i - \tilde{Z}_i^\top\tilde{\alpha}_n(\tau))I(Y_i > 0)\tilde{C}_i,$$
we have

\[ S_{n,\tau}^Q \xrightarrow{d} N(0, \Sigma_\tau), \]

where

\[ \Sigma_\tau = \tau(1 - \tau) \left\{ E \left( \tilde{C}_i \tilde{C}_i^\top \right) - E \left( \tilde{C}_i \tilde{Z}_i^\top \right) E \left( \tilde{Z}_i \tilde{Z}_i^\top \right)^{-1} E \left( \tilde{Z}_i \tilde{C}_i^\top \right) \right\} = \tau(1 - \tau)\Sigma_0. \]

Replacing \( \tilde{C}_i \) with \( \tilde{C}_i = \tilde{C}_i - E(\tilde{C}_i | \tilde{Z}_i) \), the decorrelated version \( \tilde{S}_{n,\tau}^Q \) has the same asymptotic distribution but \( \Sigma_0 = E \left( \tilde{C}_i \tilde{C}_i^\top \right) \).

(b) At a fixed \( \tau \), given \( \beta(\tau) = 0 \), we have

\[ T^Q_\tau \xrightarrow{d} \chi^2_q. \]

(c) Let \( S_n^Q = (S_{n,\tau_1}^Q, \ldots, S_{n,\tau_K}^Q) \) be a vector of rank score statistics on a sequence of quantile levels \( 0 < \tau_1 < \cdots < \tau_K < 1 \), given \( \beta(\tau_1) = \cdots = \beta(\tau_K) = 0 \), we have

\[ S_n^Q \xrightarrow{d} N(0, \Sigma), \]

where the \((k,k)\)th diagonal block of \( \Sigma \) is \( \tau_k(1 - \tau_k)E \left( \tilde{C}_i \tilde{C}_i^\top \right) \), which can be approximated by \( V_{n,\tau_k} \), the \((k,l)\)th off-diagonal block is \( (\min\{\tau_k, \tau_l\} - \tau_k \tau_l) E \left( \tilde{C}_i \tilde{C}_l^\top \right) \), which can be approximated by \( n^{-1}(\min\{\tau_k, \tau_l\} - \tau_k \tau_l) \tilde{C}^* \tilde{C}^* \).

All proofs of this section are deferred to Appendix A.2 and A.3.

**Remark:** The proposed rank-score test correcting zero-inflation biases has two major differences compared to the standard one. First, the rank-score statistic (4), in fact, is computed based on the subset of data with positive \( Y_i \)'s. Second, to correct the biases caused by zero-inflation in testing, we incorporate the zero-positive uncertainty in estimating the variance of the rank-score by introducing the zero-truncated nominal covariates. Technically, \( E(\tilde{C}_i \tilde{C}_i^\top) = \mathbb{E}\{C_i C_i^\top P(Y_i > 0 | X_i)\} \). As a consequence, \( V_{n,\tau} \) implicitly incorporates a “propensity score” of each cell, compensating the variability caused by the random status of the gene being observed or not.

2.4. **step-wise algorithm of ZIQRank test in scRNA-seq data.** In differential gene expression analysis, we are interested in testing whether the distribution of the expression level of a gene differs according to the cell conditions \( C \). Following the two-part quantile regression model (1) and (2), the task translates into the following global null hypothesis,

\[ H_0 : \gamma = 0 \text{ & } \beta(\tau) = 0, \forall \tau \in (0, 1). \]

To test the global null hypothesis, we propose a step-wise test procedure, where we first marginally construct a test statistic for \( \gamma \) and the proposed rank-score test statistics (Section 2.3) for \( \beta(\tau) \) on a grid of quantile levels. We then combine the marginal p-values while taking the correlations of the marginal test statistics into account. We call the proposed test as zero-inflated quantile rank-score based test (ZIQRank). We describe the following three steps to implement ZIQRank.

**Step 1. Construct a logistic regression test for \( \gamma \):** Conduct any asymptotically valid test, i.e., Wald test, Rao’s score test or likelihood-ratio test, based on the estimated logistic regression model (1). Denote the test statistic as \( T^L \) and the p-value as \( p^L \).
Step 2. Construct the proposed rank-score test that corrects zero-inflation on a sequence of quantile levels: Following Theorem 1(b), we can compute the p-values $p_{q_k}^Q$ associated with $T_{q_k}^Q$, $k = 1, \ldots, K$ at quantile levels $0 < \tau_1 < \cdots < \tau_K < 1$. Selection of the grid of quantile levels will be elaborated in Section 2.5.1.

Step 3. Combination of marginal p-values: Combine the marginal p-values by MinP (Lee, Wu and Lin 2012; He et al. 2017) or Cauchy combination test (Liu and Xie 2019). Compared to the joint $\chi^2$ test, the two procedures are more appropriate for sparse strong signals, which is a characteristic of scRNA-seq data. The choice between MinP and Cauchy tests on specific data will be discussed in Section 2.5.2.

$T_{ZIQRank-MinP} = \min\{p^L, p_{r_1}^Q, \ldots, p_{r_K}^Q\}$: It uses the minimum p-value as the test statistic, and derives the final p-value by resampling based on the dependence structure of the marginal test statistics. The null hypothesis will be rejected if it is unlikely to observe an even smaller minimum p-value. Let $q_{l_{\min}}^L$ denote the $(1 - T_{ZIQRank-MinP})$th percentile of the distribution of $T^L$, and $q_{l_{\min}}^Q$ denote the $(1 - T_{ZIQRank-MinP})$th percentile of the distributions of $T_{q_k}^Q$, $k = 1, \cdots, K$, which are all $\chi_2^2$. The p-value based on $T_{ZIQRank-MinP}$ is

$$P\{\exists T_{q_k}^Q \geq q_{l_{\min}}^Q, k = 1, \cdots, K \text{ or } T^L \geq q_{l_{\min}}^L | H_0\}$$

$$= 1 - P\{T^L < q_{l_{\min}}^L | H_0\} \cdot P\{\forall T_{q_k}^Q < q_{l_{\min}}^Q, k = 1, \cdots, K | H_0\}$$

$$= 1 - (1 - T_{ZIQRank-MinP}) \cdot P\{\forall T_{q_k}^Q < q_{l_{\min}}^Q, k = 1, \cdots, K | H_0\},$$

where the first equality is based on the asymptotic independence between $T^L$ and $T_{q_k}^Q$ (Appendix A.2). The joint probability $P\{\forall T_{q_k}^Q < q_{l_{\min}}^Q, k = 1, \cdots, K | H_0\}$ can be computed via resampling $S_{n,\tau_k}$’s from the joint limiting distribution of $S_{n,\tau_k}$ under the null hypothesis (Theorem 1(c)), and calculating the realizations of $T_{q_k}^Q$’s with the help of $V_{n,\tau_k}$’s, $k = 1, \cdots, K$.

$T_{ZIQRank-Cauchy} = \tilde{r}_n \tan\{(0.5 - p^L)\pi\} + \sum_{k=1}^K w_k \tan\{(0.5 - p_{r_k}^Q)\pi\}$, where $\tilde{r}_n$ is the observed proportion of zero in $Y_i$’s, and

$$w_k = \frac{\tau_k I(\tau_k \leq 0.5) + (1 - \tau_k) I(\tau_k > 0.5)}{\sum_{k=1}^K \{\tau_k I(\tau_k \leq 0.5) + (1 - \tau_k) I(\tau_k > 0.5)\}},$$

i.e., the sum of all weights is 1, and the p-values associated with central quantiles are assigned with larger weights while those on extreme tails have smaller weights: It uses the weighted average of p-values. After tangent transformation of the p-values, the aggregate statistic follows standard Cauchy distribution under the null hypothesis, regardless of the dependence structure of the marginal test statistics. The p-value based on $T_{ZIQRank-Cauchy}$ is

$$1 - \Phi_{Cauchy}(T_{ZIQRank-Cauchy}).$$

Remark: ZIQRank is an omnibus test, which aggregates all signals of association between the gene expression and cell condition together, regardless of the magnitude and direction. A small p-value indicates that the gene shows differences in either zero proportions or quantiles of the positive part at some levels, or both. When the data is non-normal, such as the zero-inflated scRNA-seq gene expression, combining signals over quantile processes is often more efficient in detecting the difference in comparison to the mean-based approaches, as it captures more heterogeneity among different groups. Similar power gain was reported in Zhao and Xiao (2014).
2.5. Practical considerations of implementing ZIQRank.

2.5.1. Selection of the grid of quantile levels. One issue that affects the testing performance is the selection of a quantile grid. Without prior knowledge or preference, the entire distribution is of interest; thus, a grid covering all typical quantiles is required. Note that if one aims to test whether a particular region is different among distributions, e.g., the tail differences, she/he can specify a grid within the interested interval.

Next, the number of quantiles should be chosen with care. More quantiles might introduce more significant signals, but might also incorporate extra noise. When computing resources are adequate, a tuning process to achieve a satisfactory power while keep Type I error controlled is encouraged. Note that the number of quantile levels cannot exceed the number of the positive observations of the gene, i.e., \( K < (1 - \hat{r}_n)n \). Otherwise, the marginal tests will be highly correlated and the Type I error might be inflated.

2.5.2. Choice between MinP and Cauchy combination tests. MinP and Cauchy combination methods use “local” and “global” principles, respectively, to combine p-values, and have advantages in different scenarios.

When the data has a high degree of heterogeneity, the MinP test will be more powerful because it makes the decision based on the most significant signal. Otherwise, the Cauchy test will be preferred since it pools the uniformly significant or insignificant signals together. This point will be demonstrated by comparing the data characteristics and results of simulation studies (Section 3) and real data applications (Section 4 and 5).

In addition, the MinP test is robust to the extent of zero-inflation, because all the p-values from either logistic or quantile tests are treated equally. By the Cauchy test, however, the result from the logistic component will dominate if the gene expression is highly zero-inflated. This point will be illustrated by comparing the simulation study with cell condition only (Section 3.1) and the simulation study with additional covariates (Section 3.2).

Finally, the Cauchy test is easier to implement and more computationally efficient, because there is no need to estimate the joint probability by resampling based on the dependence structure of marginal test statistics. When computing resources are limited, the Cauchy test will be preferred.

2.5.3. Default settings of ZIQRank in practice. It is practically helpful to specify a default setting of ZIQRank. We recommend a dense quantile grid \( \tau = 0.05, 0.1, \cdots, 0.95 \) for continuous scRNA-seq data (e.g., non-UMI TPM). The grid distributes over the quantile support \((0, 1)\) evenly and thoroughly, helping to fully capture the distributional difference, and is conservative for most scRNA-seq data, which are usually very big. For count data (e.g., UMI count), to ensure reliable estimations and valid inferences, a jittering uniformly distributed with a lower limit 0 and upper limit 1 should be added, i.e., the rank-score (4) becomes

\[
S_{n,\tau}^Q = n^{-\frac{1}{2}} \sum_{i=1}^{n} \psi_\tau \{ W_i - \hat{Z}^*_i \hat{\alpha}_n(\tau) \} I(Y_i > 0) \hat{C}^*_i, \]

where \( W_i = Y_i + U \) and \( U \sim (0, 1) \). Machado and Silva (2005) guarantees that ZIQRank is still valid. To compromise with potential spurious signals by the jittering, a less dense but commonly used grid in quantile analysis \( \tau = 0.1, 0.25, 0.5, 0.75, 0.9 \) is recommended.

MinP test is recommended as the default setting for p-value combination considering the heterogeneity in most scRNA-seq data. Cauchy test will be chosen if the data is less heterogeneous. The rule for heterogeneity is simple and fast to implement. Across genes, calculate (1) the number of crossings of the quantile functions and (2) the coefficient of variation (CV) of non-zero quantile differences between cell conditions. If medians of both measures are larger than 1, the data is considered heterogeneous.

In the following numerical studies, we will use the default setting of ZIQRank to compare with existing approaches. ZIQRank with the default grid but the other combination method will also be examined, for illustration purposes only.
3. Simulation. In this section, we investigate the finite sample performance of ZIQRank in comparisons with existing methods through two simulation studies. Both studies aim to identify DEGs between two cell conditions. The first simulation study only considers the cell conditions without adjusting confounding covariates, named as “unadjusted analysis”. The second one includes one additional covariate, named as “adjusted analysis”.

For comparison, we choose 8 competing methods: (1) QRank, (2) QRank with MinP test, (3) QRank with Cauchy test, (4) linear regression, (5) MAST, (6) Monocle, (7) scDD and (8) DESeq2. Each of them serves as a representative of one group of approaches given the underlying model and characteristics. QRank is a direct quantile-based method ignoring zero-inflation and using a joint $\chi^2$ test. QRanks with MinP / Cauchy tests are presented to remove the effect of various p-value combination methods. However, applying the 3 QRanks directly to the zero-inflated scRNA-seq data is not viable. On the simulated data, their estimations failed in 21 genes (out of 10000), and we observed inflated false positive rates (nearly 25%). To make QRanks applicable in this work, we manually add a small random perturbation to the probability mass at zero, to create “pseudo-continuous” data. MAST and Monocle are parametric scRNA-seq approaches, using Gaussian linear or generalized additive model for the positive part, and consider zero-inflation in the models. scDD is a non-parametric Bayesian method to detect the difference between two cell conditions and cannot adjust additional covariates. DESeq2 is a commonly used method for bulk RNA-seq data, assuming negative binomial distribution, and does not consider zero-inflation.

3.1. Simulation of unadjusted analysis.

3.1.1. Setting of the unadjusted analysis. The simulated data are generated following the simulation framework in scDD R package with modifications. scDD simulates the data for two cell conditions, which mimics the gene expression distributions in a human embryonic stem cell scRNA-seq data, and generates both DEGs and null genes (NGs). Based on the starting data, scDD could cluster and simulate 4 scenarios of DEGs: (1) traditional differential expression (DE), (2) differential proportion of cells within each component (DP), (3) differential modality (DM), and (4) both differential modality and different component means within each condition (DB). Figure 1 plots the density and quantile functions for the 4 DEG scenarios. DE shows a unimodal distribution in density plot and a homogeneous difference in quantile plot, while DP, DM and DB have more than one mode in density plots and demonstrate heterogeneity in quantile differences. Additionally, the two quantile functions in DP form a spindle shape, and those in DB cross at some central quantiles, which are two typical types of differential distribution with minimal mean differences but substantial differences in quantiles. Apart from the overall performance, we aim to examine ZIQRank on each of the 4 types of DEGs, especially those with multi-modal distributions.

However, based on the human embryonic stem cell starting data, scDD generates DEGs with similar percentages of zeros between the two conditions. To introduce extra zero-inflation and more differentiated zero rates between conditions, we simulate DEGs using scDD, then further convert a certain percentage of the lowest non-zero expressions of each DEG to zero. The rationale is that the dropouts of scRNA-seq data are usually due to low gene expression. The selected proportions of the lowest non-zero values for conversion to zero follow a uniform distribution with a lower limit 0 and an upper limit 25%. We do not impose an extra zero-inflation more than 25%, to keep the specific shapes of DE, DP, DM and DB.

For each simulated data, 10000 genes are simulated for two conditions with a sample size of 200 cells each. 8000 NGs are simulated. 2000 genes are simulated as DEGs, with 500 for each of the 4 DEG scenarios. Logarithmic transformation is applied to the simulated data.
Fig 1: Density and quantile functions of the 4 scenarios of differential distributions, DE: traditional differential expression (topleft), DP: differential proportion (topright), DM: differential modality (bottomleft), DB: both differential modality and different component means (bottomright). The dashed lines in quantile plots are the mean expression levels of the two conditions.

To confirm that the simulated data represents the human embryonic stem cell scRNA-seq data, we plot and compare (1) the dropout rates vs. mean and (2) mean vs. variance across genes (Figure S1). It shows that the synthetic data captures and even exaggerates the zero-inflated nature of the real data, and does a satisfactory job in recapitulating the mean-variance relationship. Thus, this simulation is reliable.

ZIQRank with $\tau = 0.05, 0.1, \cdots, 0.95$ quantile grid and MinP procedure is the default, since the simulated data is continuous and highly heterogeneous (median of number of crossings between quantile functions=6, median of CV of quantile difference>1, shown in Figure S2a). The existing methods and ZIQRank are applied to each simulated data. Genes are considered as DEGs when the corresponding adjusted p-values using the Benjamini-Hochberg (BH) procedure are less than 0.05. Precision, defined as the number of true positive calls among all positive calls, and recall, defined as the number of true positive calls among all the true DEGs are calculated. They are summarized into precision-recall (PR) curve and area under the curve (AUC). Also, stratified recall is calculated for each of the 4 DEG scenarios. Stratified precision cannot be calculated, since all the methods except scDD cannot classify the positive calls into the 4 DEG scenarios. False positive rate (FPR), defined as the number of false positive calls among all NGs, is calculated. The simulation process is repeated 10 times, and the results are summarized using means and standard deviations (sd) of the number of DEGs detected and the number of true DEGs detected. We present the PR curves with boxplots of corresponding AUCs, and display the boxplots of stratified recalls and FPRs over the 10 simulation runs.

3.1.2. Results of the unadjusted analysis. ZIQRank has the largest numbers of detected DEGs (1878.10) and correctly detected DEGs (1838.80) (Table 1). Even when the precision-recall trade-off is considered, it yields the highest AUC 0.99. (Figure 2a). In all the 4 DEG scenarios, ZIQRank works best with the highest recall rates (Figure 2b).
Fig 2: Performance of ZIQRank and existing methods in the unadjusted analysis.
Table 1
Summary statistics of the number of DEGs detected, and the number of correctly detected DEGs by ZIQRank and existing methods with adjusted p-value less than 0.05, in unadjusted simulation study.

<table>
<thead>
<tr>
<th>Method</th>
<th>nDEG</th>
<th>sd</th>
<th>ntrueDEG</th>
<th>sd</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZIQRank-MinP</td>
<td>1878.10</td>
<td>9.34</td>
<td>1838.80</td>
<td>9.41</td>
</tr>
<tr>
<td>ZIQRank-Cauchy</td>
<td>1390.00</td>
<td>21.84</td>
<td>1375.10</td>
<td>21.28</td>
</tr>
<tr>
<td>QRank</td>
<td>1636.40</td>
<td>17.91</td>
<td>1600.20</td>
<td>16.11</td>
</tr>
<tr>
<td>QRank-MinP</td>
<td>1825.40</td>
<td>16.02</td>
<td>1802.30</td>
<td>15.10</td>
</tr>
<tr>
<td>QRank-Cauchy</td>
<td>1278.60</td>
<td>16.71</td>
<td>1268.10</td>
<td>15.73</td>
</tr>
<tr>
<td>linear</td>
<td>1365.60</td>
<td>20.35</td>
<td>1349.00</td>
<td>18.74</td>
</tr>
<tr>
<td>MAST</td>
<td>1586.80</td>
<td>8.73</td>
<td>1578.10</td>
<td>8.89</td>
</tr>
<tr>
<td>Monocle</td>
<td>1255.20</td>
<td>24.59</td>
<td>1236.90</td>
<td>23.48</td>
</tr>
<tr>
<td>scDD</td>
<td>1656.50</td>
<td>15.62</td>
<td>1643.30</td>
<td>13.50</td>
</tr>
<tr>
<td>DESeq2</td>
<td>1215.60</td>
<td>25.06</td>
<td>1215.40</td>
<td>25.10</td>
</tr>
</tbody>
</table>

The 1st runner-up is QRank-MinP, which detects 1802.30 true DEGs and has an AUC of 0.98. The outstanding performance is expected, as MinP test is good at picking the most significant signal in the heterogeneous data. Though the AUCs are close, ZIQRank surpasses QRank-MinP in terms of power, especially for DB scenario (DB recall of ZIQRank vs. QRank-MinP=83% vs. 78%). Thus, we can infer that modeling zeros is beneficial, by incorporating the signal from different zero proportions between conditions and amplifying the signals on the non-zero part, especially further separating the crossed distribution functions of DB-type DEGs.

All other methods have inferior AUCs (AUC of scDD=0.96, QRank=0.95, MAST=0.92, etc.). Almost all methods provide satisfactory results for DE and DM scenarios, with stratified recalls higher than 80%. DE and DM scenarios have a clear mean difference between the two conditions, which can be detected by both parametric and non-parametric methods. For DP and DB scenarios, especially DB, ZIQRank has much higher recalls than the parametric methods. This is because the parametric approaches target at detecting the location/mean difference, which is minimal in DP and DB scenarios, while ZIQRank aims to detect quantile difference, which is substantial in the two scenarios.

Though ZIQRank has the highest FPR (Figure 2c), it is still well-controlled below 0.01. Its PR curve and highest AUC also suggest a tolerable false discovery rate (FDR=1-precision). Therefore, though it does not dominate in all aspects, ZIQRank has a non-negligible value in improving the power and complementing the existing methods with controlled false positives.

Overall, ZIQRank performs best in detecting DEGs in terms of PR-AUC in the unadjusted analysis. It improves and complements the existing methods in identifying DP and DB-type multi-modal DEGs, without undermining the power in identifying those DEGs with mean differences, while keeping false positives controlled.

3.2. Simulation of adjusted analysis.

3.2.1. Setting of the adjusted analysis. We further compare the performance of ZIQRank with other methods by including one additional covariate. Similar to Section 3.1.1, we first use scDD R package to generate 10000 genes for two conditions with 200 cells each, including 8000 NGs and 2000 DEGs from the 4 DEG scenarios. Denote the logarithmic transformed expression as $Y_{i,j}$ for the $j$th gene in the $i$th cell. Then, we add one cell-level covariate $Z_i$ into the simulation by $Y'_{i,j} = Y_{i,j} + \alpha_j Z_i$, where $\alpha_j$ is the covariate effect on the $j$th gene. $Z_i$ is simulated from a normal distribution with mean 5 and sd 1.5. $\alpha_j$ is assigned to be zero for randomly selected 75% of the simulated genes; for the remaining genes, $\alpha_j$ is simulated from a uniform distribution with a lower limit of 0 and an upper limit of 0.25. Following
the same procedure in Section 3.1.1, we generate extra zero-inflation by further converting a certain percentage of the lowest non-zero expressions to zero. Similarly, the mean-zeros and mean-variance plots (Figure S1) verify that the simulated data is a fair representation of the human embryonic stem cell scRNA-seq data.

ZIQRank with \( \tau = 0.05, 0.1, \ldots, 0.95 \) quantile grid and MinP procedure is still the default (data is continuous and highly heterogeneous, with median of number of crossings between quantile functions=6, median of CV of quantile difference>1 by Figure S2b). scDD is excluded, as it cannot handle confounding covariates. Genes are considered as DEGs when the BH adjusted p-values are less than 0.05. The numbers of DEGs detected, the PR curves with AUCs, stratified recalls and FPRs are summarized over 10 simulation runs.

### 3.2.2. Results of the adjusted analysis

ZIQRank is the most powerful method as it has the largest number of correctly detected DEGs (1825.10) (Table 2). It also produces the highest AUC (0.98) (Figure 3a). Again, it achieves the highest stratified recalls for all the 4 types of DEGs (Figure 3b).

With a covariate associated with zero proportions and non-zero expressions, we observe more obvious drawbacks of QRanks. The AUCs of QRank, QRank-MinP and QRank-Cauchy are 0.89, 0.94 and 0.85, with excessive false positives shown by their low precisions and high FPRs (Figure 3c). MinP combination helps QRank to boost power, but still inferior to ZIQRank, especially for DB scenario (DB recall of ZIQRank vs. QRank-MinP=81% vs. 78%). The uncontrolled false positives and lower testing power confirm the benefits of modeling inflated zeros when using quantile-based methods in the adjusted analysis.

All other methods have lower AUCs (AUC of MAST=0.92, etc.). For DE and DM scenarios with a substantial mean difference, all methods work well, with recall rates higher than 80%. For DP and DB scenarios, especially DB, with a small difference in mean and large differences in quantiles, ZIQRank dominates the parametric methods.

Note that the performance of ZIQRank-Cauchy improves from the unadjusted analysis (unadjusted vs. adjusted AUC=0.81 vs. 0.95). As discussed in Section 2.5.2, this is because Cauchy test is sensitive to the signals from the zero proportion (shown by larger prevalence of significant logistic p-values than that in unadjusted analysis, Figure S3b vs. S3a).

To sum up, similar to the unadjusted analysis, ZIQRank performs best in terms of PR-AUC in the adjusted analysis. When confounding covariates exist, it still improves and enriches the competing methods by identifying highly heterogeneous DEGs, sustaining the power in identifying DEGs with mean differences, while keeping false positives controlled.

### 4. Analysis of scRNA-seq data about human glioblastoma
(a) Precision-recall curves for all methods (left) and boxplots of area under the curves (right).

(b) Boxplots of stratified recalls for the 4 DEG scenarios.

(c) Boxplots of FPR.

Fig 3: Performance of ZIQRank and existing methods in the adjusted analysis.
4.1. Data source and analysis settings. In this section, we illustrate the performance of ZIQRank in detecting DEGs between neoplastic and non-neoplastic cells, using one scRNA-seq data from a human glioblastoma multiforme study Darmanis et al. (2017) (GSE84465). The dataset was downloaded from conquer repository (Soneson and Robinson 2018). 3584 cells from both the tumor core and the peritumoral brain were sequenced using Smart-seq2 protocol, including 1091 neoplastic cells and 2493 non-neoplastic cells. Cells were collected from 4 patients, named S1, S2, S4 and S6. There are 487 cells from S1, 1169 cells from S2, 1540 cells from S4 and 388 cells from S6.

We apply the following pre-processing steps to the data before analysis: (1) Take the average of TPM of the same gene within each cell, (2) delete the gene if it has positive expressions for one cell condition only, (3) remove the gene if its zero-inflation rate is higher than 97.5%, and (4) take the logarithmic transformation of TPM, i.e., $\log(\text{TPM} + 1)$. After the pre-processing, the dataset includes 22970 genes.

We consider two ways to analyze the data, one is unadjusted analysis, considering only the two cell conditions, and the other is adjusted analysis, which includes a cell-level covariate, the patient to whom the cells belong. All approaches used in simulation studies are applied to the data, except DESeq2. DESeq2 can not be applied here due to its great demand for computing resources, which exceeds the maximum capacity we have. Also, its performance is demonstrated to be worse than other methods in our simulation studies (Section 3). ZIQRank with $\tau = 0.05, 0.1, \cdots, 0.95$ quantile grid and Cauchy test is the default, given the data is continuous and lacks heterogeneity (median of number of crossings between quantile functions=1, median of CV of quantile difference<1, shown in Figure S4a).

4.2. Evaluation of Type I error. Before analyzing the data using different methods, we evaluate their Type I errors on the real scRNA-seq data. Using the pre-processed data, we permute the covariates (cell condition, patient ID) jointly at the cell level to create 50 permuted datasets. The permutation maintains the association between the cell-level covariates, but removes the association between the cell condition and gene expression. Therefore, the permuted datasets are supposed to have no DEGs, and genes with small p-values are considered false positives. We calculate the Type I error as the proportion of genes with nominal p-values of less than 0.01. This evaluation procedure on real data is borrowed from Soneson and Robinson (2018). The Type I error evaluated is essentially a similar concept as FPR with BH procedure in Section 3, but using different criteria to determine the false positives. We use nominal p-values in real data because, unlike simulated data, the true p-value distribution in real data is unknown and may heavily impact the BH adjusted p-values.

As Figure 4 shows, in both analyses, Monocle has a slightly inflated Type I error, while all other methods keep Type I error well-controlled with around 1% genes having p-values smaller than 0.01. Thus, the results of all approaches except Monocle are trustworthy in the following analyses.

4.3. Results on detecting DEGs. We analyze the pre-processed data to identify DEGs using the various methods. The genes with BH adjusted p-values less than 0.01 are considered to be DEGs between neoplastic and non-neoplastic cells. Table 3 provides the numbers of DEGs detected by all approaches in both unadjusted and adjusted analyses.

In the unadjusted analysis, scDD identifies the most DEGs (16817), and ZIQRank detects the second most DEGs (16425). In the adjusted analysis, ZIQRank detects the largest number of DEGs (13777). QRank-Cauchy only detects 14267 and 13014 DEGs, showing that modeling zeros helps the quantile-based tests to capture more signals. All methods detect fewer DEGs in the adjusted analysis compared to the unadjusted analysis. This indicates that the covariate adjustment is necessary when analyzing GSE84465, to reduce false positive genes
due to the confounding effect. Therefore, the interpretation of detected DEGs in the following is based on the adjusted analysis, and we can conclude that ZIQRank has an improved power of identifying DEGs than the others.

Further, to evaluate the various methods, we compare their detected genes to the functional genes in the established pathways, De Novo pathway and Secondary pathway (map05214 of KEGG PATHWAY, Kanehisa and Goto 2000). Amplification, deletion or mutation of the genes in the two paths affect the progression or suppression of glioma. Those genes are considered to be the critical DEGs between neoplastic and non-neoplastic cells based on bi-

Fig 4: Type I error control of ZIQRank and competing methods in analyzing GSE84465.
TABLE 3
Number of DEGs between neoplastic and non-neoplastic cells detected by ZIQRand under the 6 settings and competing methods on GSE84465 for both unadjusted and adjusted analyses.

<table>
<thead>
<tr>
<th>Method</th>
<th>Unadjusted analysis</th>
<th>Adjusted analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZIQRank-MinP</td>
<td>15537</td>
<td>12764</td>
</tr>
<tr>
<td>ZIQRank-Cauchy</td>
<td>16425</td>
<td>13777</td>
</tr>
<tr>
<td>QRank</td>
<td>12968</td>
<td>11792</td>
</tr>
<tr>
<td>QRank-MinP</td>
<td>14670</td>
<td>13589</td>
</tr>
<tr>
<td>QRank-Cauchy</td>
<td>14267</td>
<td>13014</td>
</tr>
<tr>
<td>linear</td>
<td>14239</td>
<td>12345</td>
</tr>
<tr>
<td>MAST</td>
<td>16367</td>
<td>13696</td>
</tr>
<tr>
<td>Monocle</td>
<td>13791</td>
<td>12027</td>
</tr>
<tr>
<td>scDD</td>
<td>16817</td>
<td>-</td>
</tr>
</tbody>
</table>

20 genes from the two pathways are contained in the pre-processed GSE84465 data. ZIQRank detects all of the 20 genes, with the oncogenes, EGFR and CDK4, the tumor suppressors, ARF and Rb, and several others, Shc, Grb2, E2F, GADD45, assigned with super small adjusted p-values that approach 0. The oncogenes, PDGF, PDGFR and MDM2, and tumor suppressor, PTEN, and the rest, TGF\(_{\alpha}\), IGF-1, Raf, mTOR, CDK6, POLK, etc. are also detected by ZIQRank with quite small adjusted p-values at the level of \(10^{-3}\). The competing approaches miss several of the critical genes in the two pathways. The mean-based approaches, MAST, Monocle and linear regression, fail to detect Raf, POLK and CDK6, and the uncorrected quantile-based method, QRank, fails to detect mTOR and CDK6.

Next, ZIQRank exclusively identifies 136 DEGs, while the existing methods fail to detect them. Previous biological investigations show that some of those genes are differentially expressed between neoplastic and regular cells, and play important roles in diagnosis, progression or suppression of glioma. For example, CHD4 is proved to be over-expressed in glioma cells and is a potent suppressor of glioma (McKenzie et al. 2019). We examine the expression profiles of those genes and identify the pattern that highlights ZIQRank’s better detection performance. Figure 5 shows the quantile and violin plots of 4 representative genes (UBR1, PCDHA4, PHLPP1 and UBE2D3). The 4 genes share a similar distributional pattern, having a subtle mean difference between the neoplastic and non-neoplastic conditions, but possessing substantial differences in quantiles. The two quantile functions either cross or form a spindle shape. Note that the 4 genes can be classified as DB or DP scenarios, as described in simulation studies (Section 3). The fundamental properties of ZIQRank can explain its improved power in detecting the two types of DEGs compared to the previous approaches. ZIQRank does not make any particular parametric assumptions on the gene expression (e.g., Gaussian or negative binomial distribution), but captures the quantile differences at various locations of the expression distribution, and detect higher-order associations between the cell condition and gene expression.

For UBR1, the quantile functions of the two cell conditions cross at quantile level 75% with neoplastic cells having higher quantile values below the crossing point, indicating a 75% chance of over-expression in neoplastic cells. This is consistent with Uhlen et al. (2010) and Fazi et al. (2015)’s findings that the expression of UBR1 is moderately higher in glioma cells. The quantile functions of PCDHA4 form a typical spindle shape, which indicates a proportion difference (DP) in high expression state. We divide the gene expression of PCDHA4 into high expression state and low expression state using the cutoff of 1.5 in log-transformed TPM. There is a higher proportion of the neoplastic cells in high expression state (13.74%) compared to the non-neoplastic ones (9.95%), with a p-value of 0.001 by \(\chi^2\) test. The over-expression of PCDHA4 in glioma cells is supported by Uhlen et al. (2010), and PCDHA4 also plays a vital suppression role in other cancers (Tombolan et al. 2016). The violin plot of
PHLPP1 shows a distributional pattern of DB scenario with the neoplastic cells having two modes and the non-neoplastic cells having only one mode. PHLPP1 in neoplastic cells have larger quantile values at higher percentages and smaller quantile values at lower percentages (1.36 at quantile level 0.75 and 0.00 at quantile level 0.5) as compared to non-neoplastic cells (1.10 at quantile level 0.75 and 0.07 at quantile level 0.5). This is a typical example of the tumor suppressor’s negative feedback process discussed in Section 1, and Teng et al. (2016) confirmed that PHLPP1 plays a vigorous suppression role in the inflammatory response of glioma. For UBE2D3, we define the expression in log-transformed TPM above 2.6 to be high expression state, and observe that the proportion of neoplastic cells in high expression state is 64.80%, higher than that of non-neoplastic cells (60.41%), with a p-value of 0.014 by \( \chi^2 \) test. Obacz et al. (2019) confirms the over-expression of UBE2D3 and points out that it controls the recruitment of myeloid cells to glioma.

Besides studying the 4 representative genes from a biological perspective, we define their quantile effect sizes associated with ZIQRank, and numerically illustrate the advantages of ZIQRank over the others on such heterogeneous genes. The effect size, \( \Delta \), is defined as the total area between the two cell conditions’ quantile functions. Since the total area under the quantile function curve equals the mean, the quantile effect size is comparable to mean-based methods. For example, the mean difference of UBR1 is 0.01, but \( \Delta_{UBR1} = 0.09 \), supporting that ZIQRank is more capable of differentiating the complex distributions.

Once a gene is identified, post-hoc analyses on the quantile specific p-values and zero proportion p-value also shed light on how the gene expression distribution differs by conditions. The p-values help depict whether the difference is focused at upper or lower tails or a particular part of the distribution, or whether zero proportions are significantly different between two conditions. For example, rank-score tests on PHLPP1 give small p-values on the 0.3th-0.7th quantiles of the non-zero expression, corresponding to the area between the two crossing points of quantile functions (Figure 5). Moreover, if one aims to confirm the regional difference statistically, she/he can choose a quantile grid within the interested area and use ZIQRank again, as suggested in Section 2.5.1.

In conclusion, ZIQRank improves the power of detecting DEGs, and complements the existing approaches by identifying additional heterogeneous genes. Those genes uniquely detected by ZIQRank are numerically meaningful due to their substantial quantile differences between cell conditions. Moreover, supported by literature, they are biologically crucial by revealing complex biological mechanisms.

4.4. Comparisons on computational performance. Due to the gigabyte size level of scRNA-seq data, computational complexity is a critical measure of scRNA-seq analysis methods. We compare ZIQRank to the existing differential methods by recording the time and memory used in analyzing GSE84465. As presented in Table 4, the time consumed by ZIQRank is 34min, which is comparable to MAST. scDD is the fastest only because we use its fast procedure, while its standard setting is beyond our computing resources. Moreover, scDD cannot be applied in the general adjusted analysis. In terms of memory, ZIQRank entails the least resource of 3G, whereas Monocle, MAST and scDD need 5G, 6G and 8G, and DESeq2 needs more than 32G.

5. Analysis of scRNA-seq data about cells from mouse intestinal organoids.

5.1. Data source and analysis settings. We also demonstrate the advantages of ZIQRank on UMI count data, which is sequenced by CEL-Seq protocol and has different properties from the TPM data in Section 4: (1) the zero values are due to low counts that follow Poisson or over-dispersed Poisson distributions and (2) the data are zero-inflated integers in nature.
Fig 5: Quantile and violin plots of 4 DEGs detected by ZIQRank exclusively on GSE84465.

Table 4

<table>
<thead>
<tr>
<th>Method</th>
<th>Time</th>
<th>Memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZIQRank</td>
<td>34min</td>
<td>3G</td>
</tr>
<tr>
<td>MAST</td>
<td>31min</td>
<td>6G</td>
</tr>
<tr>
<td>Monocle</td>
<td>66min</td>
<td>5G</td>
</tr>
<tr>
<td>scDD</td>
<td>12min</td>
<td>8G</td>
</tr>
<tr>
<td>DESeq2</td>
<td>-</td>
<td>&gt;32G</td>
</tr>
</tbody>
</table>

The dataset was contributed by a study about cells from mouse intestinal organoids Grün et al. (2015) (GSE62270-GPL17021), and downloaded from conquer repository. 2891 cells were collected, including 1547 marker-positive cells and 1344 randomly extracted cells from whole intestinal organoids. We pre-process the data following similar steps, except that we take the average of UMI counts, but round to preserve integer and do not take logarithmic transformation. After the pre-processing, the dataset includes 14545 genes.

We consider unadjusted analysis only, since there are no cell-level covariates other than cell conditions. All methods used in simulation studies are applied to the data, including DESeq2 (our computing resource can handle it on the relatively small data). As DESeq2 assumes negative binomial distribution, it has been popular to analyze such count sequencing data. We also use the count version of Monocle, which assumes negative binomial as well. ZIQRank with \( \tau = 0.1, 0.25, 0.5, 0.75, 0.9 \) quantile grid and Cauchy test is the default, given the data is count and lacks heterogeneity (median of number of crossings between quantile functions=0, median of CV of quantile difference<1, shown in Figure S4b).

5.2. Evaluation of Type I error. We permute the cell conditions to create 50 null datasets, and calculate the Type I error by the proportion of genes with nominal p-values less than 0.01 within each set. As figure 6 shows, all methods except Monocle have Type I error controlled with around 1% genes having p-values smaller than 0.01.

5.3. Results on detecting DEGs. We analyze the pre-processed data to identify DEGs using the various methods. The genes with BH adjusted p-values less than 0.01 are considered to be DEGs between marker-positive and randomly extracted cells. Table 5 provides
Fig 6: Type I error control (by boxplots of FPR from 50 null datasets in unadjusted analysis) of ZIQRank and competing methods in analyzing GSE62270-GPL17021.

### Table 5

*Number of DEGs between marker-positive and randomly extracted cells from mouse intestinal organoids detected by ZIQRank and competing methods on GSE62270-GPL17021.*

<table>
<thead>
<tr>
<th>Method</th>
<th>Unadjusted analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZIQRank-MinP</td>
<td>9661</td>
</tr>
<tr>
<td>ZIQRank-Cauchy</td>
<td><strong>10117</strong></td>
</tr>
<tr>
<td>QRank</td>
<td>5769</td>
</tr>
<tr>
<td>QRank-MinP</td>
<td>6443</td>
</tr>
<tr>
<td>QRank-Cauchy</td>
<td>6436</td>
</tr>
<tr>
<td>linear</td>
<td>5850</td>
</tr>
<tr>
<td>MAST</td>
<td>9915</td>
</tr>
<tr>
<td>Monocle-count</td>
<td>10263</td>
</tr>
<tr>
<td>scDD</td>
<td>8245</td>
</tr>
<tr>
<td>DESeq2</td>
<td>4821</td>
</tr>
</tbody>
</table>

The numbers of DEGs detected by all approaches in the unadjusted analysis. We exclude Monocle-count in the following comparisons, as it has exceedingly inflated Type I error. ZIQRank detects the most DEGs (10117). As QRank-Cauchy only detects 6436 DEGs, we can confirm that modeling zeros is also rewarding for UMI count data, where zeros are due to low counts following Poisson over-dispersed Poisson distributions. MAST detects the second most DEGs (9915), still inferior to ZIQRank because its Gaussian assumption for the non-zero part is inadequate for the complex count distributions. DESeq2 detects the least DEGs (4821), as it misses the inflated zeros, even though it uses negative binomial distribution. Thus, for UMI count data, we can also conclude that ZIQRank has an improved power of identifying DEGs compared to the others. There is no established pathway to further evaluate the detected genes, given that the cells were collected based on markers but not phenotypes.

ZIQRank exclusively identifies 75 DEGs, which have substantial quantile differences between the marker-positive and randomly extracted cells but negligible mean differences. Their patterns are similar to those in Figure 5, thus omitted.
To sum up, on UMI count data, ZIQRank can also improve and enrich the existing methods by detecting more genes with heterogeneous associations with cell conditions.

5.4. Comparisons on computational performance. We compare ZIQRank to the existing differential methods by recording the time and memory used in analyzing GSE62270-GPL17021. As presented in Table 6, ZIQRank is the fastest and still the most economical, using only 6min and 2G. Since a smaller quantile grid is used, both time and memory are improved as compared to Table 4. Monocle-count is the 1st runner-up, entailing 8min and 3.5G. Its drastic improvement from Table 4 is due to the different assumed models, Tobit for continuous data and negative binomial for count data, and the difference in corresponding estimation algorithms. As expected, DESeq2 is the slowest using 44min, and requires the most memory of 10G.

6. Conclusion and discussion. In this paper, we have proposed a zero-inflated quantile rank-score test (ZIQRank) based on a two-part quantile regression model to detect differentially expressed genes in scRNA-seq data. The two-part model models the probability of zero counts by logistic regression, and models the non-zero expression intensity by quantile regressions. A new rank-score test tailored for the two-part quantile regression model was proposed. The ZIQRank test then marginally constructs a logistic regression test statistic on the zero counts, and the novel rank-score test statistics at multiple quantiles of non-zero expressions. Finally, it combines the marginal tests using MinP or Cauchy procedures with the incorporation of test dependency.

The two-part quantile model considers inflated zeros and allows a comprehensive assessment of differentially expressed genes’ distributions, without the restrictions of parametric likelihoods. Both are essential considerations for scRNA-seq analysis, given the zero-inflated and heterogeneous nature of scRNA-seq data. In both simulation studies and real applications, we have shown that ZIQRank outperforms the existing methods in identifying differentially expressed genes in scRNA-seq data. With Type I error well-controlled, it improves the power, and complements the existing methods by revealing extra heterogeneous genes with substantial quantile differences but negligible mean differences among cell conditions. By design, the ZIQRank test can easily incorporate confounding covariates, and is easy to implement and computationally efficient. Hence, it is a generic and practical inference tool for scRNA-seq data. The ZIQRank test can be applied to other zero-inflated data with complex distributional attributes, such as microbiome data.

Results, simulation settings and data examples of this paper are quite consistent with current benchmarking papers. Soneson and Robinson (2018) provides a comprehensive comparison of differential expression analysis methods using the PowSim R package to simulate data and a subset of conquer database as real data examples. Wang et al. (2019) evaluates the performance of several selected methods by using the scDD R package and one scRNA-seq data in Islam et al. (2011). We demonstrate the performance of the proposed ZIQRank using
modified scDD R package and two scRNA-seq data in conquer. Though our paper focused on a different angle, the overlapped part has a very similar conclusion that there is a trade-off between the true positive rate and precision of calling DEGs. Methods with a higher true positive rate tend to show a lower precision due to their introducing false positives, whereas methods with a high precision show a low true positive rate due to identifying few DEGs.

There are various directions to extend ZIQRank. First, the current ZIQRank uses a default setting based on the data characteristics, and its optimal performance can be achieved by tuning the grid of quantile levels. As a progress, one may consider estimating the entire quantile process, and construct a global simultaneous test. The construction of such a test statistic and its asymptotic theory needs to re-established. Second, the current test is conducted on individual genes separately. We can further enhance the power by incorporating the neighboring correlated genes or external functional information into the test.
A.1. Conditions for Theorem 1. Theorem 1 relies on the following sufficient conditions:

**ASSUMPTION 1.** The observations \( \{(X_i, Y_i); i = 1, \cdots, n\} \) are i.i.d. from a joint distribution \( P \), where \( X_i \) is a \( p \)-dimensional vector of covariates including the cell condition \( c_i \).

**ASSUMPTION 2.** For any \( \tau \in (0, 1) \), the conditional distribution function of \( Y_i \)'s given \( Y_i > 0 \), \( \{F_i = F_{Y|Y_i>0}(\cdot|X_i)\} \), are absolutely continuous, with continuous densities \( \{f_i\} \) uniformly bounded away from 0 and \( \infty \) at \( \{F_i^{-1}(\tau|X_i)\} \) with a bounded first-order derivative, \( i = 1, \cdots, n \).

**ASSUMPTION 3.** Eigenvalues of \( \mathbb{E}(\tilde{C}_i\tilde{C}_i^\top) \) and \( \mathbb{E}(\tilde{Z}_i\tilde{Z}_i^\top) \) are bounded away from 0 and \( \infty \).

**ASSUMPTION 4.** There exists a positive constant \( b \) such that \( f_i(F_i^{-1}(\tau|X_i)) = b \) for all \( i \).

Practically, the 4 assumptions easily hold or won’t affect the performance of ZIQRank for most scRNA-seq data. Assumption 1 means that the scRNA-seq data is collected randomly, not longitudinally. Assumptions 2-3 are modified standard regularity conditions (Koenker 2005; Wang and He 2007) to assure the validity of linear quantile regression and quantile rank score test on the positive part. For scRNA-seq data, Assumption 2 implies that every conditional quantile of the non-zero gene expression is uniquely defined, which is satisfied by most commonly used distributions, such as those from the exponential family. Assumption 3 suggests that covariates used in the model are neither redundant (hence no collinearity issue) nor prone to outliers, which is a minimal requirement in most regression models. Assumption 4 postulates a homoscedasticity condition on the non-zero gene expressions of scRNA-seq data. It is a technical assumption, useful for theory derivation. In practice, the rank score is quite robust against deviation from it (Wang 2009; Wei et al. 2006).

A.2. Proof of Theorem 1(a).

**PROOF.** Define

\[
m(\theta^Q(\tau), \theta^L) = \left( \tilde{X}\{\tau - I(Y - \tilde{X}^\top\theta^Q < 0)\}, \ x\{I(Y > 0) - \pi(\theta^L, X)\} \right),
\]

\[
M(\theta^Q(\tau), \theta^L) = \int m(\theta^Q(\tau), \theta^L) dP, \quad M_n(\theta^Q(\tau), \theta^L) = \int m(\theta^Q(\tau), \theta^L) dP_n,
\]

where \( \tilde{X} = (\tilde{Z}^\top, \tilde{C}^\top)^\top \).

By the central limit theorem,

\[
\sqrt{n}M_n(\theta^Q(\tau), \theta^L) \xrightarrow{d} N\left\{0, \begin{pmatrix} \tau(1 - \tau) & 0 \times p \\ 0 & D_{1,\gamma} \end{pmatrix} \right\},
\]

where

\[
D_0 = \mathbb{E}\left\{\tilde{X}_i\tilde{X}_i^\top\right\} - \mathbb{E}\left\{\tilde{Z}_i\tilde{Z}_i^\top\right\} - \mathbb{E}\left\{\tilde{C}_i\tilde{C}_i^\top\right\} = \begin{pmatrix} A & B \\ B^\top & D \end{pmatrix},
\]
Noticing the orthogonality of $\theta^Q(\tau)$ and $\theta^L$ in the asymptotic distribution of $\sqrt{n} M_n(\theta^Q(\tau), \theta^L)$ and for simplicity, denote

$$m_\alpha(\theta^Q(\tau)) = \mathcal{Z}\{\tau - I(Y - \bar{X}^\top \theta^Q < 0)\}, \quad m_\beta(\theta^Q(\tau)) = \mathcal{C}\{\tau - I(Y - \bar{X}^\top \theta^Q < 0)\},$$

and

$$M_{n,\alpha}(\theta^Q(\tau)) = \int m_\alpha(\theta^Q(\tau)) d\mathbb{P}_n, \quad M_{n,\beta}(\theta^Q(\tau)) = \int m_\beta(\theta^Q(\tau)) d\mathbb{P}_n.$$

Since $\hat{\theta}^Q_n(\tau) = (\hat{\alpha}_n(\tau)^\top, \hat{\beta}_n(\tau) = 0)^\top$ is the restricted M-estimator, by Lagrange multiplier method,

$$\sqrt{n} M_{n,\alpha}(\hat{\theta}^Q_n(\tau)) = 0,$$

$$\sqrt{n} M_{n,\beta}(\hat{\theta}^Q_n(\tau)) + \sqrt{n} \hat{\lambda}_n = 0,$$

where $\hat{\lambda}_n$ is the estimator of the Lagrange multiplier parameter.

Let $\Delta$ be a compact neighborhood of the true $\theta^L$, $T$ be a compact subset of interval $(0, 1)$ containing $\Gamma(\tau; X, \theta^L)$, and $\Psi$ is a compact neighborhood of the true $\theta^Q \circ \Gamma(\tau; X, \theta^L)$, where

$$\theta^Q \circ \Gamma(\tau; X, \theta^L) = \theta^Q(\tau_s), \quad \text{and} \quad \tau_s = \Gamma(\tau; X, \theta^L) = \max \left( \frac{\tau - \{1 - \pi(\theta^L, X)\}}{\pi(\theta^L, X)}, 0 \right).$$

The class

$$\mathcal{F} = \left\{ m(\theta^Q(\tau), \theta^L), \tau \times \theta^L \times \theta^Q \in T \times \Delta \times \Psi \right\}$$

is clearly a VC class with a squared integrable envelop function $XX^\top$. Thus, $\mathcal{F}$ is a Donsker class. Define

$$\mathbb{G}_n(\theta^Q(\tau), \theta^L) = \sqrt{n} \{ M_n(\theta^Q(\tau), \theta^L) - M(\theta^Q(\tau), \theta^L) \},$$

The fact that $\mathcal{F}$ is a Donsker class and both $\hat{\theta}^Q_n(\tau)$ and $\hat{\theta}^L_n$ are consistent implies that

$$\mathbb{G}_n(\hat{\theta}^Q_n(\tau), \hat{\theta}^L_n) = \mathbb{G}_n(\theta^Q(\tau), \theta^L) + o_p(1),$$

which implies

$$\sqrt{n} M_n(\hat{\theta}^Q_n(\tau)) = \sqrt{n} M_n(\theta^Q(\tau)) + \nabla M(\theta^Q(\tau))\sqrt{n} (\hat{\alpha}_n(\tau) - \alpha(\tau)) + o_p(1),$$

where,

$$-\nabla M(\theta^Q(\tau)) = \left( \mathbb{E}\left\{ f_i(\bar{X}_i^\top \theta^Q(\tau)) \mathcal{Z}_i \mathcal{Z}^\top_i \right\}, \mathbb{E}\left\{ f_i(\bar{X}_i^\top \theta^Q(\tau)) \mathcal{Z}_i \mathcal{C}^\top_i \right\}, \mathbb{E}\left\{ f_i(\bar{X}_i^\top \theta^Q(\tau)) \mathcal{C}_i \mathcal{Z}^\top_i \right\}, \mathbb{E}\left\{ f_i(\bar{X}_i^\top \theta^Q(\tau)) \mathcal{C}_i \mathcal{C}^\top_i \right\} \right) = b \left( A \quad B \right),$$

with the last equality by Assumption 4.

Plugging (10) into (8) and (9) to get

$$\sqrt{n} M_{n,\alpha}(\theta^Q(\tau)) - bA\sqrt{n} (\hat{\alpha}_n(\tau) - \alpha(\tau)) = o_p(1),$$

$$\sqrt{n} M_{n,\beta}(\theta^Q(\tau)) - bB^\top \sqrt{n} (\hat{\alpha}_n(\tau) - \alpha(\tau)) + \sqrt{n} \hat{\lambda}_n = o_p(1).$$
Multiply (11) by \(-B^\top A^{-1}\) from left and add it to (12), then rearrange terms to yield
\[
\sqrt{n} \hat{\lambda}_n = \sqrt{n} \left\{ B^\top A^{-1} M_{n,\alpha}(\theta^Q(\tau)) - M_{n,\beta}(\theta^Q(\tau)) \right\} + o_p(1)
\]
\[
= \left( B^\top A^{-1}, -I \right) \sqrt{n} \left( M_{n,\alpha}(\theta^Q(\tau)) \right) + o_p(1)
\]
\[
\xrightarrow{d} \left( B^\top A^{-1}, -I \right) \cdot N \left\{ 0, \tau(1 - \tau) \left( \begin{array}{cc} A & B \\ B^\top & D \end{array} \right) \right\}
\]
\[
= N \left( 0, \tau(1 - \tau)(D - B^\top A^{-1}B) \right).
\]

Then, by (9),
\[
\tilde{S}_{n,\tau}^Q(\tilde{\alpha}_n(\tau)) = \sqrt{n} M_{n,\beta}(\tilde{\alpha}_n(\tau)) - \sqrt{n} \hat{\lambda}_n \xrightarrow{d} N \left( 0, \tau(1 - \tau)(D - B^\top A^{-1}B) \right),
\]
as claimed.

\[\square\]

A.3. Proof of Theorem 1 (b) and (c).

**Proof.** Replace \(\hat{C}_i\) by \(\hat{C}_i = \hat{C}_i - E(\hat{C}_i|\hat{Z}_i)\) to obtain \(\tilde{S}_{n,\tau}^Q(\tilde{\alpha}_n(\tau))\), and repeat the same argument as Section A.2, then
\[
\tilde{S}_{n,\tau}^Q(\tilde{\alpha}_n(\tau)) \xrightarrow{d} N \left( 0, \tau(1 - \tau)\Sigma_0 \right),
\]
where
\[
\Sigma_0 = E \left( \hat{C}_i \hat{C}_i^\top \right) - E \left( \hat{C}_i \hat{Z}_i^\top \right) E \left( \hat{Z}_i \hat{Z}_i^\top \right)^{-1} E \left( \hat{Z}_i \hat{C}_i^\top \right).
\]
By definition, \(E(\hat{C}_i \hat{Z}_i^\top) = 0\). Hence, \(\Sigma_0 = E(\hat{C}_i \hat{C}_i^\top)\).

Now we need to prove \(\tilde{S}_{n,\tau}^Q(\tilde{\alpha}_n(\tau)) \xrightarrow{d} N \left( 0, \tau(1 - \tau)\Sigma_0 \right)\), which is equivalent to prove
\[
\tilde{S}_{n,\tau}^Q(\tilde{\alpha}_n(\tau)) - S_{n,\tau}^Q(\tilde{\alpha}_n(\tau)) = o_p(1).
\]

Consider the working model
\[
\hat{C}_i = \xi^\top \hat{Z}_i + \delta_i,
\]
where \(\delta_i\) are i.i.d. random vectors with zero mean and positive definite variance. Then,
\[
\hat{C}_i^* - \hat{C}_i = - (\hat{\xi}_n - \xi)^\top \hat{Z}_i,
\]
where \(\hat{\xi}_n\) is the least square estimator of \(\xi\). By the consistency of \(\hat{\xi}_n\), which is implied by Assumption 3, and the asymptotic tightness of \(\sqrt{n} M_{n,\alpha}(\theta^Q(\tau))\) established in Section A.2, we have
\[
\tilde{S}_{n,\tau}^Q(\tilde{\alpha}_n(\tau)) - S_{n,\tau}^Q(\tilde{\alpha}_n(\tau)) = - (\hat{\xi}_n - \xi)^\top \left\{ \sqrt{n} M_{n,\alpha}(\theta^Q(\tau)) \right\} = o_p(1)O_p(1) = o_p(1),
\]
which completes the proof.

The asymptotic properties of \(S_n^Q = (S_{n,\tau_1}, \cdots, S_{n,\tau_K})^\top\) can be derived as an extension of the result on a fixed quantile. As shown in Koenker (2005), the between-quantile correlation is
\[
\min\{\tau_i, \tau_j\} - \tau_i \tau_j.
\]
Details of this proof are omitted.
Next, since \( \tilde{C}_i^* = \tilde{C}_i - (\hat{\xi}_n - \xi)^\top \tilde{Z}_i \), we have

\[
\frac{\tilde{C}_i^* \tilde{C}_i^*}{n} = \frac{\tilde{C}_i \tilde{C}_i}{n} - 2(\hat{\xi}_n - \xi)^\top \tilde{Z}^\top \frac{\tilde{C}_i}{n} + (\hat{\xi}_n - \xi)^\top \frac{\tilde{Z} \tilde{Z}}{n} (\hat{\xi}_n - \xi)
\]

\[
= \mathbb{E}(\tilde{C}_i \tilde{C}_i^\top) - 2n \mathbb{E} \{ (\hat{\xi}_n - \xi)^\top \tilde{Z} \} + \mathbb{E} \{ \tilde{Z} \tilde{Z}^\top \} (\hat{\xi}_n - \xi)
\]

\[
= \mathbb{E}(\tilde{C}_i \tilde{C}_i^\top) + o_p(1),
\]

which implies that

\[
n^{-1} \tilde{C}_i^* \tilde{C}_i^* \xrightarrow{p} \mathbb{E}(\tilde{C}_i \tilde{C}_i^\top) = \Sigma_0.
\]

Therefore, by Slusky’s Theorem, given \( \beta(\tau) = 0 \),

\[
T^Q_n = S_{n,\tau}^{-1} V_{n,\tau} S_{n,\tau}^Q \xrightarrow{d} \chi^2_q.
\]

Thus, Theorem 1 (b) and (c) are proved.

\[\square\]

**APPENDIX B: COEFFICIENT-BASED TEST**

We also tried the coefficient-based test in the quantile regression adjusting for individual zero-inflation. The asymptotic covariance matrix of \( \hat{B}_n = (\hat{\beta}_n(\tau_1), \ldots, \hat{\beta}_n(\tau_K))^\top \) can be estimated by bootstrap. In each round, we resample from the entire sample, and estimate the set of quantile coefficients, \( \hat{B}_n \), based on the positive part of the bootstrapped dataset. Such a procedure will introduce the zero-positive uncertainty into the estimation, as expected. With \( \hat{B}_n(b), b = 1, \ldots, B \), we can compute the estimated covariance matrix \( \hat{\Sigma}_{\text{coef}} \). Note that we use bootstrap to avoid the computational difficulty in the kernel estimation of the conditional local density.

By some simulation experiments, we find that it has an even higher power compared to the ZIQRank test. However it has two drawbacks. (1) The Type I error is sometimes uncontrolled, while the power is just improved marginally. This is because testing based on coefficients is generally unstable, especially at extreme quantiles (Chen and Wei (2005)). (2) Also, it is computationally intensive because of the resampling. Therefore, we still recommend the ZIQRank test for practical use.

**APPENDIX C: PARTIAL COMBINATION PROCEDURES**

Because of conditional independence, we can first combine the p-values from quantile regression, and further combine it with the p-value from logistic regression by Fisher’s combined probability test. The final test statistic follows a \( \chi^2_4 \) distribution, and it is robust to the extent of zero-inflation regardless of the combination of the quantile p-values.

**REFERENCES**


Supplementary materials. Some figures are deferred here:

Fig S1: Comparisons of simulated and reference data on the mean-zeros and mean-variance relationships. **reference**: a human embryonic stem cell scRNA-seq data that serves as the starting data of simulation; **raw**: the simulated data by scDD package without modification; **unadjusted**: the simulated data with 0-25% extra zero-inflation; **adjusted**: the simulated data with a continuous confounding covariate and 0-25% extra zero-inflation.
(a) Description of quantile difference in a simulated dataset for unadjusted analysis.

(b) Description of quantile difference in a simulated dataset for adjusted analysis.

Fig S2: Summary of quantile difference between cell conditions in simulated datasets, with median of each statistic marked by dashed red vertical lines.
(a) Raw results about zeros on a simulated dataset for unadjusted analysis.

(b) Raw results about zeros on a simulated dataset for adjusted analysis.

Fig S3: Summary of zero rates and p-value from logistic tests in simulated datasets, with median of each statistic marked by dashed red vertical lines.
(a) Summary of quantile difference, neoplastic vs. non-neoplastic cells, in GSE84465.

(b) Summary of quantile difference, marker-positive vs. randomly extracted cells, in GSE62270-GPL17021.

Fig S4: Descriptive statistics about quantile difference between cell conditions in GSE84465 and GSE62270-GPL17021, with median of each statistic marked by dashed red vertical lines.