A Systematic Evaluation of Single Cell RNA-Seq Analysis Pipelines

Library preparation and normalisation methods have the biggest impact on the performance of scRNA-seq studies

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Abstract

The recent rapid spread of single cell RNA sequencing (scRNA-seq) methods has created a large variety of experimental and computational pipelines for which best practices have not been established, yet. Here, we use simulations based on five scRNA-seq library protocols in combination with nine realistic differential expression (DE) setups to systematically evaluate three mapping, four imputation, seven normalisation and four differential expression testing approaches resulting in $\sim 3,000$ pipelines, allowing us to also assess interactions among pipeline steps. We find that choices of normalisation and library preparation protocols have the biggest impact on scRNA-seq analyses. Specifically, we find that library preparation determines the ability to detect symmetric expression differences, while normalisation dominates pipeline performance in asymmetric DEsetups. Finally, we illustrate the importance of informed choices by showing that a good scRNA-seq pipeline can have the same impact on detecting a biological signal as quadrupling the sample size.

Introduction

Many experimental protocols and computational analysis approaches exist for single cell RNA sequencing (scRNA-seq). Furthermore, scRNA-seq analyses can have different goals including differential expression (DE) analysis, clustering of cells, classification of cells and trajectory reconstruction¹. All these goals have the first analysis steps in common in that they require expression counts or normalised counts. Here, we focus on these important first choices made in any scRNA-seq study, using DE-inference as performance read-out. Benchmarking studies exist only separately for each analysis step, which are library preparation protocols^{2,3}, alignment^{4,5}, annotations⁶, count matrix preprocessing^{7,8} and normalisation⁹. However, the impact of the combined choices of the separate analysis steps on overall pipeline performance has not been quantified. In order to achieve a fair and unbiased comparison of computational pipelines, simulations of realistic data sets are necessary. This is because the ground truth of real data is unknown and alternatives, such as concordance analyses are bound to favour similar and not necessarily better methods.

To this end, we integrated popular methods for each analysis step into our simulation ¹⁶ framework powsimR¹⁰. As the basis for simulations, powsimR uses raw count matrices ¹⁷ to describe the mean-variance relationship of gene expression measures. This includes ¹⁸ the variance introduced during the experiment itself as well as extra variance due to the ¹⁹ first to computational steps of expression quantification. Adding differential expression ²⁰ then provides us with detailed performance measures based on how faithfully DE-genes ²¹ can be recovered. ²²

One main assumption in traditional DE-analysis is that differences in expression are 23 symmetric. This implies that either a small fraction of genes is DE while the expression of the majority of genes remains constant or similar numbers of genes are up- and 25 down-regulated so that the mean total mRNA content does differ between groups¹¹. 26 This assumption is no longer true when diverse cell types are considered. For example, 27 Zeisel et al.¹² found up to 60% DE genes and differing amounts of total mRNA levels 28 between cell types. This issue of asymmetry is conceptually one of the characteristics 29 that distinguishes single cell from bulk RNA-seq and has not been addressed so far. 30 Therefore, we simulate varying numbers of DE-genes in conjunction with small to large 31 differences in mRNA content including the entire spectrum of possible DE-settings. 32

Realistic simulations in conjunction with a wide array of scRNA-seq methods, allow us not only to quantify the performance of individual pipeline steps, but also to quantify interdependencies among the steps. Moreover, the relative importance of the various steps to the overall pipeline can be estimated. Hence, our analysis provides sound recommendations regarding the construction of an optimal computational scRNA-seq pipeline for the data at hand.

Results

The starting point for our comprehensive pipeline comparison is a representative selection of scRNA-seq library preparation protocols (Figure 1A). Here, we included one fulllength method (Smart-seq2¹³) and four UMI methods^{14,15,2,16}. The UMI strategies 42 encompass two plate-based (SCRB-seq, CEL-seq2) and the most common non-commercial 43 and commercial droplet-based protocols (Drop-seq, 10X Chromium). CEL-seq2 differs 44 from SCRB-seq in that it relies on linear amplification by *in vitro* transcription, while 45 SCRB-seq relies on PCR amplification using the same strategy as 10X Chromium (see 46 Ziegenhain et al.^{17,2} for a detailed discussion). We then combine the library preparation 47 protocols with three mapping approaches 18,19,20 and three annotation schemes 21,22,23 48 resulting in 45 distinct raw count matrices (Online Methods). We simulated 27 distinct 49 DE-setups per matrix, each with 20 replicates, resulting in a total of 19,980 simulated 50 data sets (Figure 1B). 51

Genome-mapping quantifies genes with high accuracy

We first investigated how expression quantification is affected by different alignment methods using our selection of scRNA-seq experiments. For each of the three following strategies we picked one the most popular methods (Supplementary Figure S2): 1. 55

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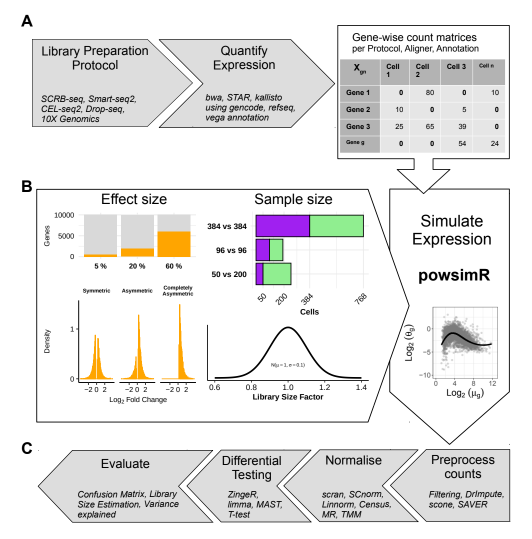


Figure 1. Study Overview

A) The data sets yielding raw count matrices. We use scRNA-seq data sets from Ziegenhain et al.² and Zheng et al.¹⁶ representing 5 popular library preparation protocols. For each data set, we obtain multiple gene count matrices that result from various combinations of alignment methods and annotation schemes (see also Supplementary Figure S1 and S2, and Supplementary Table S1 and S2). B) The simulation setup. Using powsimR Vieth et al.¹⁰ distribution estimates from real count matrices, we simulate the expression of 10,000 genes for two groups with 384 vs 384, 96 vs. 96 and 50 vs. 200 cells, where 5%, 20% or 60% of genes are DE between groups. The magnitude of expression change for each gene is drawn from a narrow gamma distribution ($X \sim \Gamma(\alpha = 1, \beta = 2)$) and the directions can either be symmetric, asymmetric or completely asymmetric. To introduce slight variation in expression capture, we draw a different size factor for each cell from a narrow normal distribution. C) The analysis pipeline. The simulated data sets are then analysed using combinations of four count matrix preprocessing, seven normalisation and four DE approaches. The evaluation of these pipelines focuses on the outcome of the confusion matrix and its derivatives (TPR, FDR, pAUC, MCC), deviance in library size estimates (RMSE) and computational run time.

alignment of reads to the genome using splice-aware alignment (STAR¹⁸), 2. alignment to the transcriptome (BWA¹⁹) and 3. pseudo-alignment of reads guided by a transcriptome (kallisto²⁴).We then combined these with three annotation schemes including two curated schemes (RefSeq²¹ and Vega²³) and the more inclusive GENCODE²² (**Supplementary Table S2**).

First, we assessed the performance by the number of reads or UMIs that were aligned 61 and assigned to genes (Figure 2A and Supplementary Figure S3). Alignment rates 62 of reads are comparable across all scRNA-seq protocols. Assignment rates on the other 63 hand show some interaction between mapper and protocol. All mappers, aligned and 64 assigned more reads using GENCODE as compared to RefSeq annotation, whereas 65 the pseudo-aligner kallisto profited most from the more comprehensive annotation 66 and here in particular the 3'UMI protocols. Generally, STAR in combination with 67 GENCODE aligned (82-86%) and assigned (37-63%) the most reads, while kallisto 68 assigned consistently the fewest reads (20-40%). BWA assigned an intermediate fraction 69 of reads (22-44%), but - suspiciously - these were distributed across more UMIs. As 70 reads with the same UMI are more likely to originate from the same mRNA molecule 71 and thus the same gene, the average number of genes with which one UMI sequence 72 is associated, can be seen as a measure of false mapping. Indeed, we find that the 73 same UMI is associated with more genes when mapped by BWA than when mapped by 74 STAR (Figure 2B). This indicates a high false mapping rate, that probably inflates the 75 number of genes that are detected by BWA (Figure 2C and Supplementary Figure **S4**). 77

This said, it remains to be seen what impact the differences in read or UMI counts obtained through the different alignment strategies and annotations have on the power to detect DE-genes.

As already indicated from the low fraction of assigned reads, kallisto has the lowest mean expression and the highest gene dropout rates (**Figure 2D** and **Supplementary Figure S7**) and, as expected from a high fraction of falsely mapped reads, BWA has the largest variance. To estimate the impact that these statistics have on the power to detect DE-genes, we use the mean-variance relationship to simulate data sets with DE-genes (**Figure 2D,E**). As previously reported², UMI protocols have a noticeably higher power than Smart-seq2 (**Figure 2F**). Moreover for Smart-seq2, we find that kallisto especially with RefSeq annoation performs slightly better than STAR, while for UMI-methods STAR performs better (**Figure 2F** and **Supplementary Figure S9**).

In summary, using BWA to map to the transcriptome introduces noise, thus considerably reducing the power to detect DE-genes as compared to genome alignment using STAR or the pseudo-alignment strategy kallisto, but given the lower mapping rate of kallisto STAR with GENCODE is generally preferable.

Many asymmetric expression changes pose a problem without ⁹⁴ spike-in data. ⁹⁵

The next step in any RNA-seq analysis is the normalisation of the count matrix. The main idea here is that the resulting size factors correct for differing sequencing depths. In order to improve normalisation, spike-ins as an added standard can help, but are not feasible for all scRNA-seq library preparations. Another avenue to improve normalisation would be to deal with sparsity by imputing missing data prior to normalisation as

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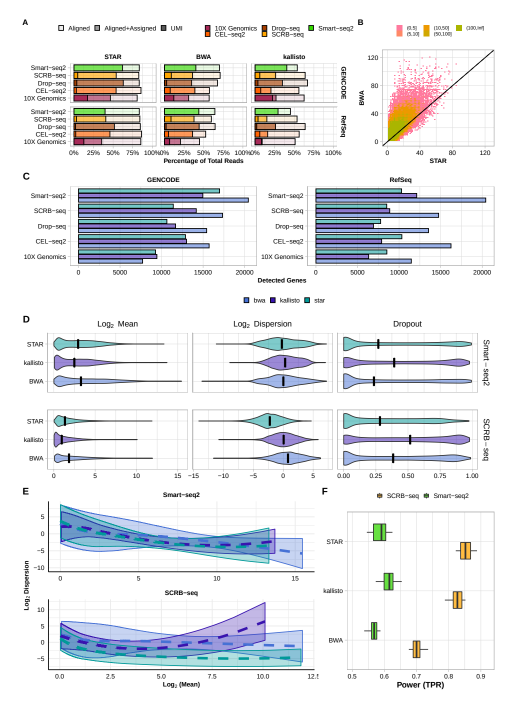


Figure 2. Expression Quantification.

A Read alignment and assignment rates per library preparation protocol stratified over aligner and annotation. The lighter shade represents the percentage of the total reads that could be aligned and the darker shade the percentage that also was uniquely assigned (see also Supplementary Figure S3). For comparability, cells were downsampled to 1 million reads/cell, with the exception of 10X Genomics data that were only sequenced to on average 60,000 reads/cell. Hence, these data are farther from saturation and have a higher UMI/read ratio. \mathbf{B} Number of genes per UMI with >1 reads for BWA and STAR alignment using the SCRB-seq data set and GENCODE annotation. Colours denote number bins of UMIs. C Number of genes detected per Library Preparation Protocol stratified over Aligner and Annotation (i.e. at least 10 % nonzero expression values) (see also Supplementary Figure S4). D Estimated mean expression, dispersion and gene dropout rates for SCRB-seq and Smart-seq2 data using STAR, BWA or kallisto alignments with GENCODE annotation (see also Supplementary Figure S7). \mathbf{E} Mean-dispersion fitting line applying a cubic smoothing spline with 95% variability bands for SCRB-seq and Smart-seq2 data using STAR, BWA or kallisto alignments with GENCODE annotation (see also Supplementary Figure S8). **F** The effect of quantification choices on the power (TPR) to detect differential expression stratified over library preparation and aligner. The expression of 10,000 detected genes over 768 cells (384 cells per group) were simulated given the observed mean-variance relation per protocol. 5% of the simulated genes are differentially expressed following a symmetric narrow gamma distribution. Unfiltered counts were normalised using scran. Differential expression was tested using limma-trend (see also Supplementary Figure S9).

discussed in the next chapter (Figure 1C). To begin with, we compare how much the 101 estimated size factors deviate from the truth. As long as there is only a small proportion 102 of DE-genes or if the differences are symmetric, estimated size factors are not too far 103 from the simulated ones and there are no large differences among methods (Figure 3A 104 and **Supplementary Figure S12**). However with increasing asymmetry, size factors 105 deviate more and more and the single cell methods scran^{25} and SCnorm^{26} perform 106 markedly better than the bulk methods TMM²⁷, MR²⁸ and Positive Counts as well 107 as the single cell method Linnorm²⁹. Census³⁰ is an outlier in that it has a constant 108 deviation of 0.1, which is due to filling in 1 when library sizes could not be calculated. 109

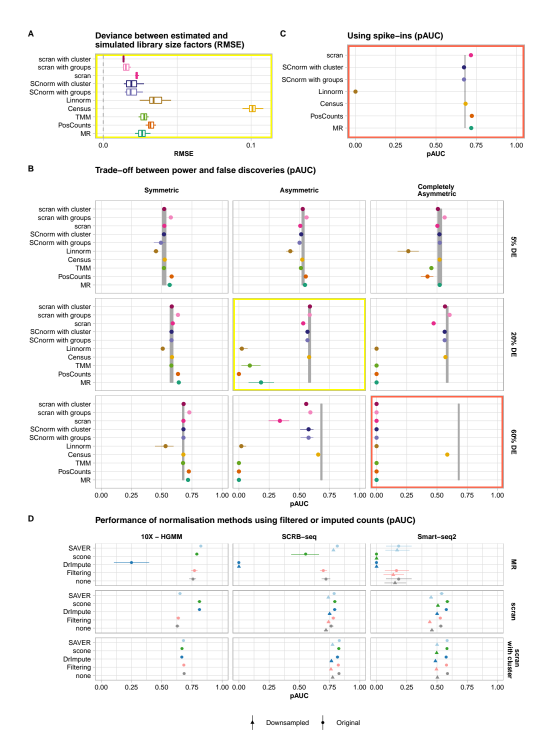
To determine the effect of these deviations on downstream analyses, we evaluated the 110 performance of differential expression inference using different normalisation methods 111 (Figure 3B and Supplementary Figure S15). Firstly, the differences in the TPR 112 across normalisation methods are only minor, only Linnorm performed consistently 113 worse (Supplementary Figure S13). In contrast, the ability to control the FDR 114 heavily depends on the normalisation method (Supplementary Figure S14). For 115 small numbers of DE-genes or symmetrically distributed changes, the FDR is well 116 controlled for all methods except Linnorm. However, with an increasing number and 117 asymmetry of DE-genes, only SCnorm and scran keep FDR control, provided that cells 118 are grouped or clustered prior to normalisation. In our most extreme scenario with 119 60% DE-genes and complete asymmetry, all methods except Census loose FDR control. 120 SCnorm, scran, Positive Counts and MR regain FDR control with spike-ins for 60% 121 completely asymmetric DE-genes (Supplementary Figure S14). Given similar TPR 122 of the methods, this FDR control determines the pAUC (**3B**,**C**). 123

Since in real data it is usually unknown what proportion of genes is DE and whether cells contain differing levels of mRNA, we recommend a method that is robust under all tested scenarios. Thus, for most experimental setups scran is a good choice, only for Smart-seq2 data without spike-ins, Census might be a better choice.

Imputation has little impact on pipeline performance.

If the main reason why normalisation methods perform worse for scRNA-seq than for bulk data is the sparsity of the count matrix, reducing this sparsity by either more stringent filtering or imputation of missing values should remedy the problem³¹. Here, we test the impact of frequency filtering and three imputation approaches (DrImpute³², scone³³, SAVER³⁴) on normalisation performance. Note, that we use the imputation or filtering only to obtain size factor estimates, that are then used together with the raw count matrix for DE-testing.

We find that simple frequency filtering has no effect on normalisation results (Figure 136 **3D**). Performance as measured by pAUC is identical to using raw counts. In contrast, 137 imputation can have an effect on performance and there are large differences among 138 methods. Imputation with DrImpute and scone rarely increased the pAUC and occa-139 sionally as in the case of SCRB-seq with MR normalisation, the pAUC even decreased 140 by 100% and 76%, respectively due to worse FDR control relative to using raw counts 141 (Supplementary Figure 18). In contrast, these two imputation methods achieved an 142 appreciable increase in pAUC together with scran normalisation, $\sim 28\%$, 4% and 9% for 143 10X Genomics, SCRB-seq and Smart-seq2 data, respectively. SAVER on the other hand 144 never made things worse, irrespective of data set and normalisation method but was 145





The data in panels A-C are based on Smart-seq2 data, all panels are based on two groups of 384 cells, STAR alignment with GENCODE annotation was used for quantification. A The root mean squared error (RMSE) of estimated library size factors per normalisation method is plotted for 20% asymmetric DE-genes (see also Supplementary Figure S12). B The discriminatory ability determined by the partial area under the curve (pAUC) based on DE testing with limma-trend for normalisation without spike-ins per DE-pattern. The grey ribbon indicates the pAUC given simulated size factors (see also Supplementary Figure S13-S15). C Using spike-ins for normalisation for 60% completely asymmetric DE-genes. D Effect of preprocessing the count matrix for 20% asymmetric DE-genes without spike-ins. Counts were either left asis ('none'), filtered or imputed prior to normalisation. The derived scaling factors were then used for normalisation and DE testing was performed on raw counts using limma-trend (see also Supplementary Figure S16-S18). This procedure was applied to the full count matrix (circle) and to the count matrix downsampled to 10% of its original sequencing depth (triangular). Missing data points are due to failing imputation runs with the sparser data.

able to rescue FDR control for MR normalisation of UMI data, even in a completely asymmetric DE-pattern. 147

These observations suggest that data sets with a high gene dropout rate might 148 benefit more from imputation than data sets with a relatively low gene dropout rate 149 (Supplementary Figure S16-18). In order to further investigate the effect of im-150 putation on sparse data, we downsampled the Smart-seq2 and SCRB-seq data, which 151 were originally based on 1 million reads/cell, to make them more comparable to the 152 10X-HGMM data with on average of 60,000 reads/cell. A radical downsampling to 10%153 of the original sequencing depth decreases the number of detected genes for SCRB-seq 154 by only 1%, suggesting that the original RNA-seq library was sequenced to saturation. 155 In contrast, the Smart-seq2 data were much less saturated at 1 million reads/cell: Down-156 sampling reduced the number of detected genes by 34%. However, the relative effect of 157 imputation on performance remains small. This is probably due to the fact that the 158 main effect of downsampling is a reduction in the detected genes, which also cannot be 159 imputed. Thus, if a good normalisation method is used to begin with (e.g. scran with 160 clustering), the improvement by imputation remains relatively small. 161

Good normalisation removes the need for specialised single cell 162 DE-tools.

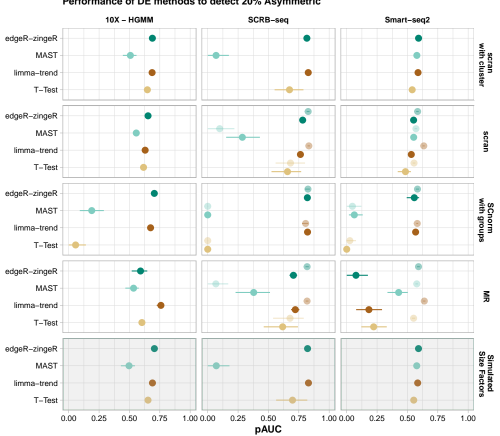
The final step in our pipeline analysis is the detection of DE-genes. Recently, Soneson ¹⁶⁴ and Robinson³¹ benchmarked 36 DE approaches and found that $edgeR^{27}$, MAST³⁵, ¹⁶⁵ limma-trend³⁶ and even the T-Test performed well. Moreover, they found that for ¹⁶⁶ edgeR, it is important to incorporate an estimate of the dropout rate per cell. Therefore, ¹⁶⁷ we combine edgeR here with zingeR³⁷. ¹⁶⁸

Both edgeR-zingeR and limma-trend in combination with a good normalisation reach similar pAUCs as using the simulated size factors (Figure 4). However, in the case of edgeR-zingeR this performance is achieved by a higher TPR paid while loosing FDR control (see Supplementary Figure S20), even in the case of symmetric DE-settings (Supplementary Figure S22-S24).

Nevertheless, we find that DE-analysis performance strongly depends on the nor-174 malisation method and on the library preparation method. In combination with the 175 simulated size factors or scran normalisation, even a T-Test performs well. Conversely, in 176 combination with MR or SCnorm, the T-Test has an increased FDR (Supplementary 177 Figure S20). SCnorms bad performance with a T-Test was surprising given SCnorms 178 good performance with limma-trend (Figure 3B). One explanation could be the rela-179 tively large deviation of SCnorm derived size factors (Figure 3A and Supplementary 180 Figure S12) which inflate the expression estimates. 181

Furthermore, we find that MAST performs consistently worse than the other DE-tools when applied to UMI-based data, but -except in combination with SCnorm- it is doing fine with Smart-seq2 data. Interestingly, Census normalisation in combination with edgeR-zingeR outperformed limma-trend with Smart-seq2 (Supplementary Figure S25).

In concordance with Soneson and Robinson³¹, we found that limma-trend, a DE-tool developed for bulk RNA-seq data showed the most robust performance. Moreover, library preparation and normalisation appeared to have a stronger effect on pipeline performance than the choice of DE-tool.



Performance of DE methods to detect 20% Asymmetric

Figure 4. Evaluation of DE tools.

The expression of 10,000 genes over 768 cells (384 cells per group) were simulated given the observed mean-variance relation per protocol. 20% of the simulated genes are differentially expressed following an asymmetric narrow gamma distribution. Unfiltered counts were normalised using simulated library size factors or applying normalisation methods. Differential expression was tested using T-Test, limma-trend, MAST or edgeR-zingeR. The discriminatory ability of DE methods is determined by the partial area under the curve (pAUC) for the TPR-FDR curve (see also Supplementary Figure S19-S21).

Normalisation is overall the most influential step.

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Because we tested a nearly exhaustive number of $\sim 3,000$ possible scRNA-seq pipelines, 191 starting with the choice of library preparation protocol and ending with DE-testing, 192 we can estimate the contribution of each separate step to pipeline performance for our 193 different DE-settings (Figure 1 B). We used a beta regression model to explain the 194 variance in pipeline performance with the choices made at the seven pipeline steps 1) 195 library preparation protocol, 2) spike-in usage, 3) alignment method, 4) annotation 196 scheme, 5) preprocessing of counts, 6) normalisation and 7) DE-tool as explanatory 197 variables. We used the difference in pseudo- R^2 between the full model including all 198 seven pipeline steps and leave-one-out reduced models to measure the contribution of 199 each separate step to overall performance. 200

All pipeline choices together (the full model) explain ~ 50% and ~ 60% of the variance in performance, for 20% and 60% DE-genes, respectively (**Figure 5A**). Choices of preprocessing the count matrix contribute very little ($\Delta R^2 \ll 1\%$). The same is true for annotation ($\Delta R^2 \ll 2\%$) and aligner choices ($\Delta R^2 \ll 5\%$). For aligner and annotation, it is important to note that these are upper bounds, because our simulations do not include differences in gene detection rates (**Figure 2C**).

Surprisingly, the choice of DE-tool only matters for symmetric DE-setups ($\Delta R_{\text{DE}=0.2}^2 =$ 207 15%; $\Delta R_{\text{DE}=0.6}^2 = 11\%$), and the choice of library preparation protocol has an even bigger 208 impact on performance for symmetric DE-setups ($\Delta R_{\text{Symmetric}}^2 = 17 - 29\%$) and addi-209 tionally for 5% asymmetric changes ($\Delta R_{5\%}^2$ Asymmetric = 17%). Normalisation choices 210 have overall a large impact in all DE-settings ($\Delta R^2 = 12 - 38\%$), where the importance 211 increases with increasing levels of DE-genes and increasing asymmetry. Spike-ins are 212 only necessary if many asymmetric changes are expected and have little or no impact 213 if only 5% of the genes are DE or the changes are symmetric (**Figure 5A**). Moreover, 214 for completely asymmetric DE-patterns, the regression model did not converge without 215 normalisation and spike-ins, because their absence or presence alone pushed the MCCs 216 to the extremes. 217

For the best performing pipeline SCRB-seq + STAR + GENCODE + SAVER218 imputation + scran with clustering + limma-trend, using 384 cells per group instead 219 of 96 improves performance only by 6.5-8%. Sample size is more important if a naive 220 pipeline is used. For SCRB-seq + BWA + GENCODE + no count matrix preprocessing 221 + MR + T-Test the performance gain by increasing sample size is 10-12% and even 222 worse, for many asymmetric DE-genes, lower sample sizes occasionally appear to perform 223 better (Figure 5B and Supplementary Figure S21). Next, we tested our pipeline 224 on publicly available 10X Genomics data set containing the expression profiles of approx. 225 1000 human peripheral mononuclear blood cells $(PBMC)^{16}$. First, we classified the cells 226 using SingleR³⁸ into the celltypes available in the Blueprint Epigenomics Reference³⁹ 227 distinguishing Monocytes, NK-cells, CD8+T-cells, CD4+T-cells and B-cells (Figure 228 **5C,D**). We applied the previously defined good (STAR + qencode + SAVER imputation)229 + scran with clustering + limma-trend) and naive (BWA + gencode + no preprocessing 230 + MR + T-Test) pipeline to identify DE-genes between the cell types. Cross-referencing 231 the identified DE-genes with known differences in marker gene expression 39 , we find 232 that the good pipeline always identifies a higher fraction of the marker genes as DE 233 than the naive pipeline (Figure 5E). Comparing NK-cells and CD8+ T-cells, the good 234 pipeline identifies 148 known markers as DE, while the naive pipeline finds only 54. The 235

diminished separation between those two cell-types using the naive pipeline is already visible in the UMAP (Figure 5D).

In summary, we identify normalisation and library preparation as the most influential choices and the observation that differences in computational steps alone can significantly lower the required sample size nicely illustrates the importance of bioinformatic choices. 240

Discussion

Here we evaluate the performance of complete computational pipelines for the analysis of 242 scRNA-seq data under realistic conditions with large numbers of DE-genes and differences 243 in total mRNA contents between groups (Figure 1). Furthermore, our simulations allow 244 us not only to investigate the influence of choices made at each pipeline step separately, 245 but also to estimate the relative importance and interactions between different steps 246 of an entire scRNA-seq analysis pipeline. We implemented all assessed computational 247 methods and more in powsimR, so that users can easily evaluate pipeline performance 248 given their own data and expected DE-settings. 249

Beginning with the creation of the raw count matrix, we find that transcriptome 250 mapping with BWA¹⁹ appears to recover the largest number of genes. However, many 251 of these are probably due to falsely mapped reads, also increase expression variance 252 which ultimately results in a lower sensitivity (Figure 2C-F). In contrast, the pseudo-253 alignment method kallisto²⁴ appears to assign reads precisely, but looses a lot of reads 254 leading to a lower mean expression. Finally, a genome mapping approach using the 255 splice-aware aligner STAR¹⁸ in conjunction with GENCODE annotation recovers the 256 most reads with high accuracy (Figure 5F). 257

Concerning the preprocessing of the count matrix, we found in concordance with Andrews and Hemberg⁴⁰ that in particular for sparse data such as 10X, SAVER³⁴ imputation before normalisation improves performance, while filtering genes has no effect with our data sets and combinations of normalisation and DE-testing methods. 260

The choice that had the largest impact on performance throughout all tested DE-262 settings is the choice of normalisation method. Only for symmetric changes, the choice 263 of library preparation protocol had a slightly larger impact than normalisation. In 264 line with Evans et al.¹¹, we found that normalisation performance of bulk methods 265 and also some of the single cell methods declined with asymmetry (Figure 3B). In 266 particular, for 60% completely asymmetric DE-genes only Census retained FDR control. 267 Unfortunately, Census is not recommended for the use with UMI-counts. Thus, for 268 UMI-counts and 60% completely asymmetric changes, only the use of spike-ins could 269 restore test performance. In the debate about the usefulness of spike-ins 41,17 , we land on 270 the pro side: Our simulations clearly show that spike-ins are useful in DE-testing settings 271 with asymmetric changes which is likely to be a common phenomenon in scRNA-seq 272 data. Due to good performance across DE-settings and its speed (Supplementary 273 Figure S22) we would recommend scran with prior clustering as the best choice for 274 normalisation (Figure 5F). 275

The choice in DE-testing method, our final pipeline step had relatively little impact 276 on overall pipeline performance. A good normalisation prior to DE-testing alleviates the 277 need for more complex and thus vulnerable methods, such as for example MASTs hurdle 278 model which implicitly assumes that the CPM values were generated from zero inflated 279 negative binomial count distribution. Indeed, in Vieth et al.¹⁰ we showed that also 280

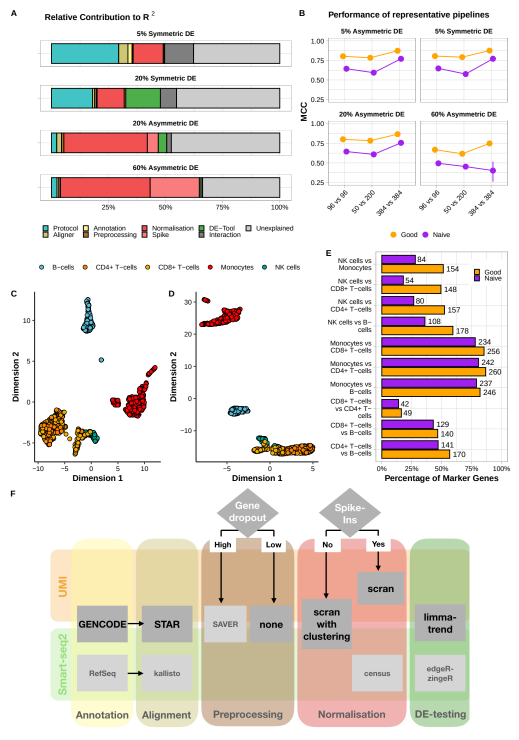


Figure 5. Evaluation of analysis pipeline.

A, **B** The expression of 10000 genes over 768 cells were simulated and 5%, 20% or 60% of the genes were differentially expressed following a symmetric or asymmetric narrow gamma distribution. This simulation setup was applied to protocols, alignments, annotations, preprocessing of counts, normalisation and DE tools. For each analysis set, the Matthew Correlation Coefficient was averaged over 20 simulations and rescaled to [0,1] interval. The MCC was used as a response variable in beta regression models with log-log link function. **A** The contribution of each covariate in the full model (Protocol + Aligner + Annotation + Preprocessing + Normalisation + DE-Tool). **B** Performance according to sample size, 1 good and 1 naive pipeline (see also Supplementary Figure S21). **C**, **D**, **E** The expression of ~ 1000 human PBMcs profiled with 10X Genomics were processed using the good and naive pipeline. Cell types were identified with SingleR classification using the Blueprint Epigenomics Reference. Cell types are represented in a UMAP, for good **C** and naive **D** pipeline, respectively. True marker genes, i.e. given by the reference, per pairwise comparison of cell types for the good and naive pipeline are given in **E** where genes needed to have a adjusted p-value < 0.1, absolute log2 fold change threshold (> 0.1) and expressed in at least 10% of the cells to be considered. **F** Pipeline recommendations for UMI and Smart-seq2 data.

scRNA-seq data fit a negative binomial distribution rather well and that the previously reported zero-inflation in scRNA-seq data is mainly due to amplification noise which is removed in UMI-data. Hence, it is not surprising that in concordance with Soneson and Robinson³¹, we find that relatively straight forward DE-testing methods adapted from bulk RNA-seq perform well with scRNA-seq data.

Finally, we want to remark that paying attention to the details in a computational pipeline and in particular to normalisation pays off. The effect of using a good pipeline as compared to a naively compiled one has a similar or even greater effect on the potential to detect a biological signal in scRNA-seq data as an increase in cell numbers from 96 to 384 cells per group (Figure 5B).

Online Methods

Single Cell RNA-seq Data Sets

The starting point for our comprehensive pipeline comparison is the scRNA-seq library preparation (Figure 1 A). In our comparison, we included the gene expression profiles of 294 mouse embryonic stem cells (mESC) as published in Ziegenhain et al.² (Supplementary 295 **Figure S1**). We selected four data sets for our comparison: Smart-seq 2^{13} a well-based 296 full-length scRNA-seq protocol, CEL-seq2¹⁵ a well-based 3' UMI-protocol using linear 297 amplification, SCRB-seq a well-based 3' UMI-protocol with PCR amplification^{42,2} 298 and Drop-seq¹⁴ a droplet-based 3' UMI-protocol. In addition, 92 poly-adenylated 299 synthetic RNA transcripts of known concentration designed by the External RNA 300 Control Consortium (ERCCs)⁴³ were spiked in for all methods except Drop-seq. All raw 301 cDNA sequencing reads were cut to a length of 45 bases and downsampled to one million 302 cDNA reads per cell (Supplementary Table S1 and Supplementary Figure S1). 303

Finally, we added a 10X Chromium data set sequencing mouse NIH3T3 cells¹⁶, ³⁰⁴ yielding ~ 400 good cells with on average $\sim 60,000$ reads/cell and another 10X data set ³⁰⁵ analysing $\sim 1,000$ human peripheral blood mononuclear cells (PBMCs). ³⁰⁶

These choices of library preparation protocols cover the diversity in current protocols without imposing partiality due to biological differences and technical handling.

Gene Expression Quantification

For genome mapping and quantification of the UMI-data with a splice-aware aligner, 310 we used the $zUMIs^{44}$ (v.0.0.3) pipeline with STAR¹⁸ (v.2.5.3a) and the mouse genome 311 (Mus_musculus.GRm38) together with annotation files (gtf) for GENCODE (vM15), 312 Vega (VEGA68) and RefSeq (Release 85) (Supplementary Table S2). zUMIs is a 313 fast and flexible pipeline for processing scRNA-seq data where cell barcode or UMI reads 314 with low sequence quality reads are filtered out prior to UMI collapsing by sequence 315 identity which yields identical count results as UMI-tools^{45,44}. For Smart-Seq2 we 316 use the same pipeline settings as in zUMIs, simply omitting the UMI collapsing step 317 (Supplementary Table S3). 318

For transcriptome alignment, we downloaded transcriptome fasta files corresponding ³¹⁹ to the annotations listed above. We used BWA¹⁹ (v0.7.12) to align the scRNA-seq reads ³²⁰ to these transcriptomes. We only removed reads that aligned equally well to transcripts ³²¹ of different genes as truly multi-mapped. The remaining reads were tallied up per gene. ³²²

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For UMI data, the reads were collapsed per gene by identity, similar to the strategy recommended in zUMIs.

For kallisto²⁴ (v0.43.1), a transcriptome-guided pseudo-alignment method, we followed the recommended quantification procedure to yield abundance estimates per equivalence class. To be comparable with other alignment methods, the counts per equivalence class were collapsed per gene. The counts of equivalence classes representing multiple genes were filtered out. For SCRB-seq, CEL-seq2, Drop-seq and 10X Genomics libraries, we chose the UMI-aware quantification option. The ERCC spike-in sequences were appended to the genome or transcriptome sequences for quantification. 320

Simulations

We used powsimR to estimate, simulate and evaluate single cell RNA-seq experiments 333 ¹⁰. PowsimR has been independently validated for benchmarking DE-approaches³¹ and 334 consistently reproduces the mean-variance relationship and dropout rates of genes of 335 scRNA-seq data (see also Supplementary Figure 28). The gene expression quantification 336 using three different aligners in combination with three annotations per library prepara-337 tion protocol produced 45 count matrices. These count matrices are the basis for our 338 estimation in powsimR. Genes needed at least one read or UMI count in at least one cell 339 to be considered in the estimation for simulation parameters. Since we 10 and others 46,47 340 have found previously, we assume that UMI counts follow a negative binomial distribution 341 and only Smart-seq2 needs the inclusion of zero-inflation. To simulate spike-in data, 342 we added an implementation of the simulation framework for pure technical variation 343 of spike-ins described in Kim et al.⁴⁸ to powsimR. The parameters required for these 344 simulations were estimated from 92 ERCC spike-ins in the SCRB-seq, CEL-seq2 and 345 Smart-seq2 data, respectively². To evaluate the effect of differing sequencing depths, we 346 added a new module to powsimR that estimates the degree of PCR amplification for 347 UMI data. This allows the user to downsample a read count matrix by binomial thinning 348 as implemented in edgeR $thinCounts()^{27}$ and then to reconstruct the corresponding 349 UMI count matrix base on the estimated PCR amplification rates. 350

For a detailed evaluation of the pipelines, we simulated two groups of cells for pairwise 351 comparisons with the following three sample size setups: 96 vs. 96, 384 vs. 384 or 50 vs. 352 200 cells (Figure 1B). For simplicity, we kept the number of genes that we simulated 353 constant at 10,000. To introduce slight variation in expression capture, we draw a different 354 size factor for each cell from a narrow normal distribution $(X \sim N(\mu = 1, \sigma = 0.1))$ 355 (Figure 1B). This distribution fits the considered data sets well, irrespective of the 356 applied library preparation method. Furthermore, the two groups of cells can have 357 5%, 20% or 60% differentially expressed genes. To capture the asymmetry of observed 358 expression differences, we considered three setups of DE-patterns: symmetric (50% up-359 and 50% down-regulated), asymmetric (75% up- and 25% down-regulated) or completely 360 asymmetric (100% up-regulated). The magnitude of expression change is drawn from a 361 narrow gamma distribution $(X \sim \Gamma(\alpha = 1, \beta = 2))$ defining the log2 fold change, which 362 is then added to the sampled mean expression. The combination of these parameters 363 results in a total of 27 DE-setups that were then applied to the parameter estimates 364 from 37 different count matrices to simulate 20 replicates for each setting, producing a 365 total of 19,980 simulated data sets. 366

These data sets were then analysed by a nearly exhaustive number of combinations 367

of four imputation strategies (scone, SAVER, DrImpute), gene dropout filtering (remove 368 genes with more than 80% zero expression values) together with seven normalisation 369 approaches (TMM, MR, Linnorm, Census, Linnorm, scran, SCnorm) with or without 370 spike-ins, depending on library preparation protocol and method (**Figure 1C**). Nor-371 malisation factors were then derived as described in Soneson and Robinson³¹ and used 372 in conjunction with the raw count matrices for DE-testing using four representative 373 approaches (T-Test, limma-trend, edgeR-zingeR, MAST). The resulting p-values were 374 corrected for multiple testing with Benjamini-Hochberg FDR and we applied a threshold 375 level of 10% to define positive test results. All these steps were seamlessly implemented 376 into powsimR (github: https://github.com/bvieth/powsimR). In total we analysed 2,979 377 different RNA-seq pipelines. 378

Evaluation metrics

To evaluate the normalisation results, we determined the root mean squared error 380 (RMSE) of a robust linear model using the difference between estimated and simulated 381 library size factors as response variable in rlm() implemented in R-package MASS⁴⁹ (v.7.3-51.1) (Supplementary Figure S10)⁹. 383

All other measures are based on the final results of an entire scRNA-seq pipeline 384 ending with DE-testing. Knowing the identity of the genes that were simulated to show 385 differing expression levels and the results of the DE-testing, we used a number of metrics 386 related to the confusion matrix tabulating the number of true positives, false positives, 387 true negatives and false negatives. We define the power to detect differential expression 388 with the TPR $(TPR = \frac{TP}{TP+FN})$. The false discovery rate is defined as $FDR = \frac{FP}{FP+TP}$. 389 We combine these two measures in a TPR versus FDR curve to quantify the trade-off 390 between true and false discoveries in a genome-wide multiple testing setup as advocated 391 by 50 . We then summarise these curves by their partial area under curve (pAUC) of 392 TPR versus observed FDR that still ensures FDR control at the nominal level of 10%393 (Supplementary Figure S11). This way of calculating the AUC is ideal for data with 394 relatively high true negative rates as the partial integration does not punish methods 395 that are over-conservative, i.e. that stay way below the nominal FDR. 396

To summarise the whole confusion matrix in one representative value we chose 397 the Matthews Correlation Coefficient $(MCC = \frac{TP*TN - FP*FN}{\sqrt{(TP+FP)(TP+FN)(TN+FP)(TN+FN)}})$, because it is a balanced measure ensuring a reliable comparison of method performance 398 399 across all DE-settings^{50,51}. As for the pAUC, we calculated the maximal value of MCC 400 where the cutoff still ensured FDR control at the nominal level of 10%. 401

To quantify the relative contribution of each step in the analysis pipeline, we used the 402 MCC as a response variable in a beta regression model implemented in R-package betareg 403 $(v.3.1-1)^{52}$ with each individual pipeline step. Because the MCC assumes the extremes 404 of 0 and 1 in some DE-settings, we applied the recommended transformation, namely 405 $MCC_{transformed} = \frac{MCC*(n-1)+0.5}{n}$, where n is the sample size⁵³. The contribution is 406 then given by the difference between the full model $pseudo - R^2$ containing all covariates 407 versus a model leaving one step out at a time. This is then scaled to the total variance 408 explained to give relative ΔR^2 percentages. 409

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Data Availability

The scRNA-seq data used in this manuscript are all publicly available, and they are summarised in Supplementary Table S1. The SCRB-seq, Smart-seq2, Drop-seq, CELseq2 data are available at the Gene Expression Omnibus (GEO) under accession code GSE75790. The HGMM and PBMC data sets are available at 10x Genomics's official website (https://support.10xgenomics.com/single-cell-gene-expression/datasets).

Code Availability

The software and code used are summarised in Supplementary Table S3 and S4. ⁴¹⁷ A compendium containing processing scripts and detailed instructions to reproduce ⁴¹⁸ the analysis for this manuscript is available from the following GitHub repository ⁴¹⁹ (https://github.com/bvieth/scRNA-seq-pipelines). ⁴²⁰

Author Contributions

B.V. and I.H. conceived the study. B.V. prepared and analysed the scRNA-seq data. B.V. implemented and conducted the simulation and evaluation framework. S.P. and C.Z. helped in data processing and power simulations. W.E. and I.H. supervised the work and provided guidance in data analysis. B.V., I.H., and W.E. wrote the manuscript. All authors read and approved the final manuscript.

Acknowledgements

This work was supported by the Deutsche Forschungsgemeinschaft (DFG) through	428
LMU excellent, SFB1243 (Subproject A14/A15) and DFG grant HE 7669/1-1. C.Z. is	429
recipient of an EMBO long-term fellowship (ALTF 673-2017).	430

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