A systematic performance evaluation of clustering methods for single-cell RNA-seq data [version 2; peer review: 2 approved]

Angelo Duò 1,2, Mark D. Robinson 1,2, Charlotte Soneson 1,2

1Institute of Molecular Life Sciences, University of Zurich, Zurich, 8057, Switzerland
2SIB Swiss Institute of Bioinformatics, Zurich, 8057, Switzerland

Abstract
Subpopulation identification, usually via some form of unsupervised clustering, is a fundamental step in the analysis of many single-cell RNA-seq data sets. This has motivated the development and application of a broad range of clustering methods, based on various underlying algorithms. Here, we provide a systematic and extensible performance evaluation of 14 clustering algorithms implemented in R, including both methods developed explicitly for scRNA-seq data and more general-purpose methods. The methods were evaluated using nine publicly available scRNA-seq data sets as well as three simulations with varying degree of cluster separability. The same feature selection approaches were used for all methods, allowing us to focus on the investigation of the performance of the clustering algorithms themselves. We evaluated the ability of recovering known subpopulations, the stability and the run time and scalability of the methods. Additionally, we investigated whether the performance could be improved by generating consensus partitions from multiple individual clustering methods. We found substantial differences in the performance, run time and stability between the methods, with SC3 and Seurat showing the most favorable results. Additionally, we found that consensus clustering typically did not improve the performance compared to the best of the combined methods, but that several of the top-performing methods already perform some type of consensus clustering.

All the code used for the evaluation is available on GitHub (https://github.com/markrobinsonuzh/scRNAseq_clustering_comparison). In addition, an R package providing access to data and clustering results, thereby facilitating inclusion of new methods and data sets, is available from Bioconductor (https://bioconductor.org/packages/DuoClustering2018).

Keywords
Clustering, Single-cell RNA-seq, RNA-seq, Benchmarking, Clustering methods
Amendments from Version 1

We thank the reviewers for their constructive comments. In response, we have made the following modifications to the manuscript:

- Clarified the rationale for including the selected data sets and methods
- Included two additional clustering methods; RaceID2 and monocle
- Exchanged the Venn diagrams in Supplementary Figure 2 for UpSet plots
- Investigated the scalability of each method by subsampling of the largest data set
- Clarified the use of random seeds by the different methods
- Increased the size of Figure 5B

In addition, the text has been clarified in several places. Detailed responses to all points raised by the reviewers are available below.

To provide easy access to the data and clustering results from our study, and thereby simplify inclusion of additional clustering methods and data sets in the comparison, we now provide an R package (available from https://bioconductor.org/packages/DuoClustering2018) leveraging the ExperimentHub framework from Bioconductor including accessor functions to retrieve all necessary data and result objects, as well as plotting functions to generate various types of plots illustrating the performance of the methods.

See referee reports

Introduction

Recent advances in single-cell RNA-seq (scRNA-seq) technologies have enabled the simultaneous measurement of expression levels of thousands of genes across hundreds to thousands of individual cells\textsuperscript{18}. This opens up new possibilities for deconvolution of expression patterns seen in bulk samples, detection of previously unknown cell populations and deeper characterization of known ones. However, computational analyses are complicated by the high variability, low capture efficiency and high dropout rates of scRNA-seq assays\textsuperscript{19}, as well as by strong batch effects that are often confounded by the experimental factor of interest\textsuperscript{19}.

Given a collection of single cells, a common analysis task involves identification and characterization of subpopulations, e.g., cell types or cell states. With lower-dimensional single-cell assays such as flow cytometry, cell type detection is often done manually, by visual inspection of a series of two-dimensional scatter plots of marker pairs ("gating") and subsequent identification of clusters of cells with specific abundance patterns. With large numbers of markers, such strategies quickly become unfeasible, and they are also likely to miss previously uncharacterized cell populations. Instead, subpopulation detection in higher-dimensional single-cell experiments such as mass cytometry (CyTOF) and scRNA-seq is often done automatically, via some form of clustering. As a consequence, a large number of clustering approaches specifically designed for or adapted to these types of assays are available in the literature\textsuperscript{11}.

While extensive evaluations of clustering methods have been performed for flow and mass cytometry data\textsuperscript{20,21}, there are to date fewer such studies available for scRNA-seq. The latter is complicated by the large number of different data generation protocols available for scRNA-seq, which in turn has a big effect on the data characteristics. Menon\textsuperscript{22} specifically evaluated three methods (Seurat\textsuperscript{11}, WGCNA\textsuperscript{18} and BackSPIN\textsuperscript{19}), illustrating their different behavior in low and high read depth data. A recent paper\textsuperscript{20} compared 12 clustering tools on scRNA-seq data sets from the 10x Genomics platform, showing that different methods generally produced clusterings with little overlap. An overview of several different types of clustering algorithms for scRNA-seq data is given by Andrews and Hemberg\textsuperscript{19}.

Here, we extend these initial studies to a broader range of data sets with different characteristics and additionally consider simulated data with different degrees of cluster separability. We evaluate 14 clustering algorithms, including both methods specifically developed for scRNA-seq data, methods developed for other types of single-cell data, and more general approaches, on a total of 12 different data sets. In order to focus on the performance of the clustering algorithms themselves, we use the same preprocessing approach (specifically cell and gene filtering) for all methods, and investigate the impact of the preprocessing separately. In addition to investigating how well the clustering methods are able to recover the true partition if the number of subpopulations is known, we evaluate whether they are able to correctly determine the number of clusters. Further, we study the stability and run time of the methods and investigate whether performance can be improved by generating a consensus partition based on results from multiple individual clustering methods, and the impact of the choice of methods to include in such an aggregation.

We observed large differences in the clustering results as well as in the run times of the different methods. SC3 and Seurat generally performed favorably, with Seurat being several orders of magnitude faster. In addition, Seurat typically achieved the best agreement with the true partition when the number of clusters was the same, while other methods, like FlowSOM, achieved a better agreement with the truth if the number of clusters was higher than the true number. Finally, we show that generally, combining two methods into an ensemble did not improve the performance compared to the best of the individual methods.

Given the high level of activity in methods research for pre-processing, clustering and visualization of scRNA-seq data, it is expected that many new algorithms (or new flavors of existing ones) will be proposed. In order to facilitate re-assessment as new innovations emerge and to provide extensibility to new methods and data sets, all (filtered and unfiltered) data sets as well as all clustering results are accessible via an R/Bioconductor package, leveraging the Bioconductor ExperimentHub framework (https://bioconductor.org/packages/DuoClustering2018). In addition, the complete code used to run all analyses is available on https://github.com/markrobinsonuzh/scRNAseq_clustering_comparison. The current system uses a Makefile to run a set of R scripts for clustering, summarization and visualization of the results.
**Methods**

**Real data sets**

Three real scRNA-seq data sets were downloaded from *conquer* and used for our evaluations: GSE60749-GPL13112 (here denoted *Kumar*), SRP073808 (*Koh*) and GSE52529-GPL16791 (*Trapnell*). These data sets were chosen to represent different degrees of “difficulty” in the clustering task. In particular, the *Trapnell* data set was not generated with the aim of detecting subpopulations, but rather to investigate a continuous developmental trajectory. Nevertheless, it was included in our evaluation as an example of a data set where the phenotype designated as the “true” cluster labels (see below) may not represent the strongest signal present in the data. Table 1 and Supplementary Figure 1 give an overview of all data sets used in this study. For each of the data sets from *conquer*, the gene-level length-scaled TPM values (below referred to as “counts” since they are on the same scale as the raw read counts) and the phenotype were extracted from the MultiAssayExperiment object provided by *conquer* and used to create a SingleCellExperiment object. We also estimated transcript compatibility counts (TCCs) for each of these data sets using kallisto v0.44, and used these as an alternative to the gene-level count matrix as input to the clustering algorithms.

The selected cell phenotype was used to define the “true” partition of cells when evaluating the clustering methods. For the *Kumar* data set, we grouped the cells by the genetic perturbation and the medium in which they were grown. For the *Trapnell* data set we used the time point (after the switch of growth medium) at which the cells were captured, and for the *Koh* data set we used the cell type annotated by the data collectors (obtained through FACS sorting). We note that the definition of the ground

---

**Table 1. Overview of the data sets used in the study.**

<table>
<thead>
<tr>
<th>Data set</th>
<th>Sequencing protocol</th>
<th># cells</th>
<th># features</th>
<th>Median total counts per cell</th>
<th>Median # features per cell</th>
<th># subpopulations</th>
<th>Description</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Koh</td>
<td>SMARTer</td>
<td>531</td>
<td>48,981</td>
<td>1,390,268</td>
<td>14,277</td>
<td>9</td>
<td>FACS purified H7 human embryonic stem cells in different differentiation stages</td>
<td>24</td>
</tr>
<tr>
<td>KohTCC</td>
<td>SMARTer</td>
<td>531</td>
<td>811,938</td>
<td>1,391,012</td>
<td>66,086</td>
<td>9</td>
<td>FACS purified H7 human embryonic stem cells in different differentiation stages</td>
<td>24</td>
</tr>
<tr>
<td>Kumar</td>
<td>SMARTer</td>
<td>246</td>
<td>45,159</td>
<td>1,687,810</td>
<td>26,146</td>
<td>3</td>
<td>Mouse embryonic stem cells, cultured with different inhibition factors</td>
<td>23</td>
</tr>
<tr>
<td>KumarTCC</td>
<td>SMARTer</td>
<td>263</td>
<td>803,405</td>
<td>717,438</td>
<td>63,566</td>
<td>3</td>
<td>Mouse embryonic stem cells, cultured with different inhibition factors</td>
<td>23</td>
</tr>
<tr>
<td>SimKumar4easy</td>
<td>-</td>
<td>500</td>
<td>43,606</td>
<td>1,769,155</td>
<td>29,979</td>
<td>4</td>
<td>Simulation using different proportions of differentially expressed genes</td>
<td>29</td>
</tr>
<tr>
<td>SimKumar4hard</td>
<td>-</td>
<td>499</td>
<td>43,638</td>
<td>1,766,843</td>
<td>30,094</td>
<td>4</td>
<td>Simulation using different proportions of differentially expressed genes</td>
<td>29</td>
</tr>
<tr>
<td>SimKumar8hard</td>
<td>-</td>
<td>499</td>
<td>43,601</td>
<td>1,769,174</td>
<td>30,068</td>
<td>8</td>
<td>Simulation using different proportions of differentially expressed genes</td>
<td>29</td>
</tr>
<tr>
<td>Trapnell</td>
<td>SMARTer</td>
<td>222</td>
<td>41,111</td>
<td>1,925,259</td>
<td>13,809</td>
<td>3</td>
<td>Human skeletal muscle myoblast cells, differentiation induced by low-serum medium</td>
<td>25</td>
</tr>
<tr>
<td>TrapnellTCC</td>
<td>SMARTer</td>
<td>227</td>
<td>684,953</td>
<td>1,819,294</td>
<td>66,864</td>
<td>3</td>
<td>Human skeletal muscle myoblast cells, differentiation induced by low-serum medium</td>
<td>25</td>
</tr>
<tr>
<td>Zhengmix4eq</td>
<td>10xGenomics GemCode</td>
<td>3,994</td>
<td>15,568</td>
<td>1,215</td>
<td>487</td>
<td>4</td>
<td>Mixtures of FACS purified peripheral blood mononuclear cells</td>
<td>5</td>
</tr>
<tr>
<td>Zhengmix4uneq</td>
<td>10xGenomics GemCode</td>
<td>6,498</td>
<td>16,443</td>
<td>1,145</td>
<td>485</td>
<td>4</td>
<td>Mixtures of FACS purified peripheral blood mononuclear cells</td>
<td>5</td>
</tr>
<tr>
<td>Zhengmix8eq</td>
<td>10xGenomics GemCode</td>
<td>3,994</td>
<td>15,716</td>
<td>1,298</td>
<td>523</td>
<td>8</td>
<td>Mixtures of FACS purified peripheral blood mononuclear cells</td>
<td>5</td>
</tr>
</tbody>
</table>
truth constitutes an intrinsic difficulty in the evaluation of clustering methods, since it is plausible that there are several different, but still biologically interpretable, ways of partitioning cells in a given data set, several of which can represent equally strong signals. Many public droplet-based data sets contain cell type labels, but these are typically inferred by clustering the cells using the scRNA-seq data, and thus any evaluation based on these labels risks being biased in favor of methods similar to the one used to derive the labels in the first place. By using ground truths that are defined independently of the scRNA-seq assay, we thus avoid artificial inflation of the signal that could result if the truth was derived from the scRNA-seq data itself.

In addition to the data sets from *conquer*, we obtained UMI counts from the Zheng data set, generated by the 10x Genomics GemCode protocol, from https://support.10xgenomics.com/single-cell-gene-expression/datasets. We downloaded counts for eight pre-sorted cell types (B-cells, naive cytotoxic T-cells, CD14 monocytes, regulatory T-cells, CD56 NK cells, memory T-cells, CD4 T-helper cells and naive T-cells) and combined them into three data sets, with a mix of well-separated (e.g., B-cells vs T-cells) and similar cell types (e.g., different types of T-cells) and uniform and non-uniform cluster sizes. For the data set denoted Zhengmix4eq, we combined randomly selected B-cells, CD14 monocytes, naive cytotoxic T-cells and regulatory T-cells in equal proportions (1,000 cells per subpopulation). For the Zhengmix4uneq data set, we combined the same four cell types, but in unequal proportions (1,000 B-cells, 500 naive cytotoxic T-cells, 2,000 CD14 monocytes and 3,000 regulatory T-cells). For the Zhengmix8eq data set, we combined cells from all eight populations, in approximately equal proportions (400–600 cells per population). For these data sets, we used the annotated cell type (obtained by pre-sorting of the cells) as the true cell label.

**Simulated data sets**

Using one subpopulation of the *Kumar* data set as input, we simulated scRNA-seq data with known group structure, using the *splatter* package\(^1\) v1.2.0. We generated three data sets, each consisting of 500 cells, with varying degree of cluster separability. For the *SimKumar4easy* data set, we generated 4 subpopulations with relative abundances 0.1, 0.15, 0.5 and 0.25, and probabilities of differential expression set to 0.05, 0.1, 0.2 and 0.4 for the four subpopulations, respectively. The *SimKumar8hard* data set consists of 4 subpopulations with relative abundances 0.2, 0.15, 0.4 and 0.25, and probabilities of differential expression 0.01, 0.05, 0.05 and 0.08. Finally, the *SimKumar8hard* data set consists of 8 subpopulations with relative abundances 0.13, 0.07, 0.1, 0.05, 0.4, 0.1, 0.1 and 0.05, and probabilities of differential expression equal to 0.03, 0.03, 0.03, 0.05, 0.05, 0.07, 0.08 and 0.1, respectively. The GitHub repository (https://github.com/markrobinsonuzh/scRNAseq_clustering_comparison) contains a link to a *countsimQC* report\(^1\), comparing the main characteristics of the simulated data sets to those of the underlying *Kumar* data set.

**Data processing**

The *scater* package\(^1\) v1.6.3 was used to perform quality control of the data sets. Features with zero counts across all cells, as well as all cells with total count or total number of detected features more than 3 median absolute deviations (MADs) below the median across all cells (on the log scale), were excluded. Depending on the availability of manual annotation, we filtered out cells that were classified as doublets or debris. The *scater* package was also used to normalize the count values, based on normalization factors calculated by the deconvolution method from the *scran* package\(^1\) v1.6.2, and to perform dimension reduction using PCA\(^3\) and t-SNE\(^4\). Either the raw feature counts or the log-transformed normalized counts were used as input to the clustering algorithms, following the recommendations by the authors (see Figure 4 for a summary of the input values used for each method).

**Gene filtering**

We evaluated three methods for reducing the number of genes provided as input to the clustering methods. For each filtering method, we retained 10% of the original number of genes (with a non-zero count in at least one cell) in the respective data sets. First, we retained only the genes with the highest average expression (log-normalized count) value across all cells (denoted *Expr* below). Second, we used *Seurat*\(^5\) to estimate the variability of the features and retained only the most highly variable ones (HVG). Finally, we used *M3Drop*\(^6\) to model the drop-out rate of the genes as a function of the mean expression level using the Michaelis-Menten equation (M3Drop). The gene-wise Michaelis-Menten constants are computed and log-transformed, and the genes are then ranked by their p-value from a Z-test comparing the gene-wise constants to a global constant obtained by combining all the genes. After filtering, we used *scran* to renormalize each data set, excluding cells with negative size factors. *Supplementary Figure 2* shows the overlap between the retained genes with the different filtering methods, for each of the 12 data sets, and *Supplementary Table 1* provides the number of cells retained after each type of filtering.

**Clustering methods**

Fourteen clustering methods, publicly available as R packages or scripts, were evaluated in this study (see Table 2 for an overview). We included general-purpose clustering methods, such as hierarchical clustering and K-means, as well as methods developed specifically for scRNA-seq data, such as *Seurat* and *SC3*, and methods developed for other types of high-throughput single-cell data (*FlowSOM*). The collection of methods include representatives for most types of algorithms commonly used to cluster scRNA-seq data. The type of the underlying clustering algorithm for the different methods is shown in Figure 4.

All methods except *Seurat* allow explicit specification of the desired number of clusters (k). *Seurat* instead requires a resolution parameter, which indirectly controls the number of clusters. For each data set, we ran each method with a range of k values (from 2 to either 10 or 15, depending on the true number of subpopulations in the data set). We ran *Seurat* with a range of resolution parameter values, yielding approximately the range of k values evaluated for the other methods. A subset of the methods provide an estimate of the true number of clusters; we record this estimate for comparison with the true number of
subpopulations. For each choice of k (or resolution), we ran each method five times, allowing us to investigate the intrinsic stability of the obtained partitions. Note that the data is the same for all five instances, and thus only the stochasticity of the clustering method affects our stability evaluation. All parameter values except for the number of clusters were set to reasonable values following the authors’ recommendations or the respective manuals (Table 2). Gene and cell filtering within the clustering methods were disabled whenever possible, since these steps were performed in a uniform way during the preprocessing and gene selection steps.

**Evaluation criteria**

In order to evaluate how well the inferred clusters recovered the true subpopulations, we used the Hubert-Arabie Adjusted Rand Index (ARI) for comparing two partitions\(^9\). The metric is adjusted for chance, such that independent clusterings have an expected index of zero and identical partitions have an ARI equal to 1, and was calculated using the implementation in the mclust R package v5.4. We also used the ARI to evaluate the stability of the clusters, by comparing the partitions from each pair of the five independent runs for each method with a given number of clusters.

We used a normalized Shannon entropy\(^6\) to evaluate whether the methods preferentially partitioned the cells into clusters of equal size, or whether they preferred to generate some large and some small clusters. Given proportions \(p_1, \ldots, p_N\) of cells assigned to each of \(N\) clusters, the normalized Shannon entropy is defined by

\[
H = -\sum_{i=1}^{N} p_i \log p_i / \log N
\]

Since the true degree of equality of the cluster sizes varies between data sets, we subtracted the normalized entropy calculated from the true partition to obtain the final performance index.

To evaluate the similarities between the partitions obtained by different methods, we first calculated a consensus partition from the five independent runs for each method, using the clue R package\(^{19}\) v0.3-5. Next, for each data set and each imposed number of clusters, we calculated the ARI between the partitions for each pair of methods, and used hierarchical clustering based on the median of these ARI values across all data sets to generate a dendrogram representing the similarity among the clusters obtained by different methods. To investigate how representative this dendrogram is, we also clustered the methods based on each data set separately, and calculated the fraction of such dendrograms in which each subcluster in the overall dendrogram appeared.

Finally, we investigated whether clustering performance was improved by combining two methods into an ensemble. For
each data set, and with the true number of clusters imposed, we calculated a consensus partition for each pair of methods using the clue R package, and used the ARI to evaluate the agreement with the true cell labels. We then compared the ensemble performance to the performances of the two individual methods used to construct the ensemble.

**Results**

**Large differences in performance across data sets and methods**

The 14 methods were tested on real data sets as well as simulations with a varying degree of complexity (Table 1) and across a range of the number of subpopulations. Focusing on the agreement between the true partitions and the clusterings obtained by imposing the true number of clusters showed a large difference between data sets as well as between methods (Figure 1; a summary across different numbers of clusters can be found in Supplementary Figure 3).

As expected, excellent performances were achieved for the well-separated data sets with a strong difference between the groups of cells (Kumar, KumarTCC and SimKumar4easy). When filtering by expression or variability, close to all methods achieved a correct partitioning of the cells in these data sets, while the M3Drop filtering led to a poorer performance for the simulated data set. All methods failed to recover the partition of the cells by time point in the Trapnell data sets, where the ARIs were consistently below 0.5. This indicates that there are other, stronger, signals in this data set that dominate the clustering.

We note that the M3Drop filtering consistently led to a worse performance for the simulated data sets, while the performance was more similar to the other filterings for the real data sets, which may indicate that the simulated dropout pattern is not consistent with the one being modeled by the M3Drop package. Due to negative size factor estimates, a larger number of cells had to be excluded in the Zhengmix data sets after the M3Drop filtering compared to the expression or HVG filtering (Supplementary Table 1). At most just over 20% of the cells in the expression and HVG filtering and up to approximately 40% of the cells for the M3Drop filtering were excluded, making a direct comparison between the filterings difficult. Furthermore, the genes retained in the M3Drop and expression filterings showed a low degree of overlap in many of the data sets (Supplementary Figure 2). Overall, only small differences were seen between the results for the data sets containing gene abundances and those containing transcript compatibility counts (TCCs).

While none of the methods consistently outperformed the others over the full range of the imposed numbers of clusters in all data sets, SC3 and Seurat often showed the best performance. These methods were also the only ones that achieved a good separation of the cell types in the droplet-based Zhengmix data sets, which have a much higher degree of sparsity and a larger number of cells than the other data sets. This is consistent with a previous study showing that Seurat performed better than other types of algorithms on data with low read depth. Generally, the performance of Seurat was also not strongly affected by the gene filtering approach (except for the simulated data sets), while other methods, like SAFE, were more sensitive to the choice of input genes for some data sets. FlowSOM showed a poor performance for the true number of clusters (see Supplementary Figure 4 for an illustration, together with a selection of other data set/method combinations with poor ARI values). However, if the number of clusters was increased, the performance of FlowSOM improved considerably, and if the methods instead were compared at the number of clusters that gave the optimal performance for each method, FlowSOM showed a better performance (Supplementary Figure 5). RtsneKmeans, a general-purpose method, showed a higher average performance across the data sets and filterings than many of the clustering methods.
algorithms specifically developed for scRNA-seq data. Compared to SC3 and Seurat, RtsneKmeans showed poorer performance for the SimKumar8hard and Zhengmix4uneq data sets. The subpopulations in these data sets are nested in the t-SNE space, explaining the difficulty in clustering for the K-means algorithm (Supplementary Figure 1).

We also investigated whether the number of detected features per cell differed between the clusters, using a Kruskal-Wallis test52. No strong association was found for the simulated data sets (Supplementary Figure 6), indicating that there is low inherent bias in the clustering algorithms. For most of the real data sets, we found highly significant differences in the number of detected features between cells in different clusters. However, it is unclear whether this represents a technical effect or a biological difference between the cell populations.

**Run times vary widely between methods**

We measured the elapsed time for each run, using a single core and excluding the time to estimate the number of clusters if this was done via a separate function. Since the run times are strongly dependent on the number of features and cells in a data set, we represent them as normalized run times, by dividing with the time required for RtsneKmeans for the same data set (Figure 2A). Seurat was the fastest method, while pcaReduce, SAFE and SC3 were the slowest, sometimes by a large margin. Clustering only half of the cells with SC3 and predicting the class of the others with a Support Vector Machine (SC3svm) gave slightly shorter run times than applying the SC3 clustering to all cells. The method could potentially be accelerated by using a lower proportion of cells as a training subset. A detailed overview of the run time and the dependence on the number of clusters is given in Supplementary Figure 7 and Supplementary Figure 8. Apart from SC3 and SC3svm, the imposed number of clusters did not affect the run time.

Plotting the run time versus the ARI for a subset of the data sets (excluding the ones with the strongest signal, where all methods found the correct clusters, and the TCC data sets) (Figure 2B) further illustrated the variability between the methods. Interestingly, Seurat was generally the fastest method, especially for the droplet-based data sets, but at the same time provided among the best partitionings of the data.

The scalability of the methods was investigated by subsampling the largest data set (Zhengmix4uneq) and plotting the run time as a function of the number of cells (Supplementary Figure 9). The majority of the methods showed a linear increase in run time as a function of the number of cells, while CIDR and RaceID2 scaled worse. The run time of SC3 and SC3svm, and to some extent SAFE, showed more complex patterns since these methods reduce the number of random starts of the Kmeans algorithm drastically if the number of cells exceeds 2,000.

**High stability between clustering runs**

Figure 1 illustrated the average performance of each method across the five runs on each data set, for the true number of clusters. By comparing the partitions obtained in the individual runs, we could also obtain a measure of the stability of each method (Figure 3A). CIDR, monocle, RaceID2, PCAHC, TSCAN, ascend and Seurat returned the same clusters in all five instances for all data sets, while the stability of the other methods depended on the data set. TSCAN and monocle set the random seed to a fixed value internally, which explains the high stability of these methods. Seurat, SC3 and RaceID2 allow the user to set the random seed via an input argument, and we explicitly set this to different values in the five independent runs. Again, the stability was lower for the simulated data sets after gene filtering by M3Drop (note that the same genes were
used in all five runs), indicating that the selection of genes may be suboptimal.

A summary of the variability both within and between the different filterings is shown in Supplementary Figure 10. It is worth noting that comparing the performances between the different filtering approaches is difficult for two reasons: first, the variability of the clustering runs for a given filtering might exceed the variation between the filterings, and second, filtering with M3Drop led to the exclusion of a large number of cells in the Zhengmix data sets, and these cells can not be used for the comparison. For the stable methods CIDR, TSCAN, ascend and PCAHC, the type of filtering had a relatively large impact on the clustering solutions, and often filtering on the mean gene expression and the gene variability gave more similar clusters than filtering with M3Drop. The stochastic methods showed both a high variability between the individual runs for a given filtering and between runs with different filterings.

**Qualitative differences between cluster characteristics**

By computing the Shannon entropy for the various partitions, we obtained a measure of the equality of the sizes of the clusters (Figure 3B). Since the true degree of cluster size uniformity as well as the number of clusters are different between data sets, we compared the normalized Shannon entropy of the clusterings to that of the true partitions. Thus, a positive value of this statistic indicates that a method tends to produce more equally sized clusters than the true ones, and a negative value instead indicates that the method tends to return more unequal cluster sizes, e.g., one large cluster and a few small ones. Most methods gave cluster sizes that were compatible with the true sizes for most data sets (a statistic close to 0), while especially...
FlowSOM was more variable, and often tended to group the cells into one large cluster and a few very small ones (see Supplementary Figure 4 for an example). One consequence of this was that FlowSOM often showed higher ARI values for a larger number of clusters, while the performance of many of the other methods decreased with increasing k (Supplementary Figure 3). These methods tended to have more equally sized clusters for larger numbers of clusters than the true number, leading to a higher disagreement between the true classification and the clusterings (the entropy across the range of k is shown in Supplementary Figure 11).

The optimal number of clusters can differ from the “true” one

Above, we investigated the performance and stability of the methods when the true number of clusters (the number of different labels in the partitioning considered as the ground truth) was imposed. Whether this number of clusters actually provided the highest ARI value (i.e., the best agreement with the ground truth) mainly depended on the difficulty of the clustering task (Figure 3C), and the choice of method. No method achieved the best performance at the annotated number of clusters in all the data sets, although generally, the methods reached their maximum performance at or near the annotated number of clusters. The notable exception was FlowSOM, which required a relatively large number of clusters to reach its maximal performance.

SC3, CIDR, ascend, SAFE and TSCAN all have built-in functionality for estimating the optimal number of clusters. In most cases, the estimated number was close to the true one; however, ascend and CIDR had a tendency to underestimate the number of clusters, while SC3 and TSCAN instead tended to overestimate the number (Supplementary Figure 12). The tendency of SC3 to overestimate the cluster number is consistent with a previous publication. The agreement with the true partition at the estimated number of clusters is shown in Supplementary Figure 13. SC3 is still the best-performing method in this situation.

Inconsistent degree of similarity between methods

The similarity between each pair of methods was quantified by means of the ARIs for each pair of consensus clusterings (across the five runs of each method for each data set and number of clusters). Figure 4 shows a dendrogram of the methods obtained by hierarchical clustering based on the average ARI values across all data sets for the true number of clusters. The numbers shown at the internal nodes indicate the fraction of the corresponding dendrograms from the individual data sets where a particular subcluster could be found. In general, the groupings of the methods shown in the dendrogram were unstable across data sets, indicated by the low stability fractions of all subclusters. This is consistent with previous studies showing generally poor concordance that varied across data sets. Even SC3 and SC3svm had surprisingly different clusterings; in less than a third of the data sets, these two methods showed the most similar clusterings. In addition, no apparent association between the similarity of the clusterings and the type of input or the dimension reduction or underlying type of clustering algorithm was seen (Figure 4).

Ensembles often don’t improve clustering performance

Next, we investigated whether we could improve the clustering performance by combining methods into an ensemble. For each pair of methods, we generated a consensus clustering and evaluated its agreement with the true partition using the ARI. In general, the performance of the ensemble was worse than the better of the two combined methods, and better than the worse of the two methods (Figure 5A), suggesting that we would obtain a better performance by choosing a single good clustering method rather than combining multiple different ones. This is
largely consistent with a recent study evaluating the combination of four methods (SC3, CIDR, Seurat, tSNE+Kmeans), where the ensemble performance was generally on par with the best individual method \(^4\). It is still possible that an ensemble method could provide a general improvement over a given single method, since it is unlikely that the same method will be the best performing in all conceivable data sets. In fact, among the methods we evaluated, both SC3 and SAFE combine multiple individual methods to achieve the final clustering result. Studying individual combinations in more detail, we observed that combining SC3 or Seurat with almost any other method led to a worse performance than obtained by these methods alone (consistent with the observation that they were among the methods giving the best performance). On the other hand, methods like CIDR, FlowSOM and TSCAN could often be improved by combining them with another method (Figure 5B).

**Discussion and conclusions**

In this study, we have evaluated 14 clustering methods on both real and simulated scRNA-seq data. There were large differences in the ability of the methods to recover the annotated clusters, and performance was also strongly dependent on the degree of separation between the true classes. SC3 and Seurat, two clustering methods developed specifically for single-cell RNA-seq data, delivered the overall best performance, and were the only ones to properly recover the cell types in the droplet-based data sets. There was, however, a large difference in the run time, with SC3 being several orders of magnitude slower than Seurat. Another difference between these two methods is that SC3 includes a method for estimating the number of clusters (which has a tendency towards overestimation), while Seurat will determine the number of clusters based on a resolution parameter set by the user.

The same preprocessing steps and fixed gene sets were used for all clustering methods. This enabled us to investigate the impact of the clustering algorithm itself, rather than entire pipelines or workflows. The selection of the filtering approach had an impact on the majority of the methods and resulted in different clustering solutions. Specifically for the more difficult data sets there was a higher dissimilarity. However, this did not necessarily affect the performances of the methods.
The stability of clustering algorithms can be evaluated by generating perturbed subsamples of the data set and redoing the clusterings. These subsamples can be created in several ways, e.g., by random subsampling with or without replacement, by adding noise to the original data or by simulating technical replicates. Freytag showed that SC3, Seurat, CIDR and TSCAN were stable under cell-wise perturbations. In our study, we evaluated the methods with respect to their sensitivity to random starts. Overall, the methods showed a high degree of stability across all data sets, except for the simulated data sets in combination with the M3Drop filtering, where the stochastic methods showed a decrease in stability. This may be due to a disagreement between the mean-dropout relationship in the simulated data and the one assumed by M3Drop, leading to a suboptimal gene selection.

The evaluated methods are based on a broad spectrum of approaches for dimensionality reduction and clustering. We note that the majority of the methods use PCA or PCoA for dimension reduction or Euclidean distances as the distance metric. Thus, no clear advice on the type of algorithm that is best suited for clustering single-cell RNA-seq data can be made based on our results. In fact, the two best-performing methods, SC3 and Seurat, rely on very different underlying clustering algorithms.

We investigated the impact of changing the imposed number of clusters for the different methods, which revealed that a subset of the methods, in particular FlowSOM, consistently showed a better agreement with the true subpopulations if the number of clusters was increased beyond the true number. The reason for this appears to be that FlowSOM tends to split off a few very small clusters. In addition to the number of clusters, most methods rely on other hyperparameters. In this study, we have fixed these to reasonable values. However, additional investigations into the effect of these hyperparameters on the results would be an interesting direction for future research.

### Data availability

The R/Bioconductor data package DuoClustering2018 provides full access to the filtered (and unfiltered) data sets, the clustering results from our study and functions for summarizing the performance of different scRNA-seq clustering methods. Additionally, helper functions and descriptions for the evaluation of new methods and data sets are provided.

Archived R scripts as at time of publication are available from https://doi.org/10.5281/zenodo.1314743

### Supplementary material

Supplementary File 1: PDF file containing Supplementary Figures 1–13 and Supplementary Table 1.

Click here to access the data

### References

   [PubMed Abstract] [Publisher Full Text]
   [PubMed Abstract] [Publisher Full Text]
   [PubMed Abstract] [Publisher Full Text] [Free Full Text]
   [PubMed Abstract] [Publisher Full Text] [Free Full Text]
   [PubMed Abstract] [Publisher Full Text] [Free Full Text]
   [PubMed Abstract] [Publisher Full Text] [Free Full Text]
7. Svensson V, Vento-Tormo R, Teichmann SA: Exponential scaling of single-cell


41. Assaf R: Identifying cell populations with scRNASeq. Mol Aspects Med. 2018; 59: 114–122. Published Abstract | Publisher Full Text


Open Peer Review

Current Peer Review Status: ✔ ✔

Version 2

Reviewer Report 11 September 2018

https://doi.org/10.5256/f1000research.17687.r38139

© 2018 Freytag S. This is an open access peer review report distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Saskia Freytag
Department of Medical Biology, University of Melbourne, Parkville, Vic, Australia

I am satisfied with the changes the authors have made.

Furthermore, I want to commend the authors for making the data accessible through an R package to facilitate further benchmarking of single cell clustering methods.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 10 September 2018

https://doi.org/10.5256/f1000research.17687.r38138

© 2018 Fan J. This is an open access peer review report distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Jean Fan
Department of Biomedical Informatics, Harvard Medical School, Boston, MA, USA

The authors have thoroughly addressed my comments. Based on the new revised results, I have the following minor comment:

- The authors note that "We found substantial differences in the performance, run time and stability between the methods, with SC3 and Seurat showing the most favorable results."

Please clarify what metric(s) you are using to assess "most favorable results" ie. "most favorable results in
terms of overall accuracy in cell-type identification", "in terms of run time", "in terms of stability", etc. Based on your new Supplemental Figure 9, it seems that SC3 would not be favorable in terms of runtime/scalability.

With this minor comment addressed, I find this article now suitable for indexing in F1000 and look forward to the authors’ future follow-up work.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

---

**Author Response 21 Sep 2018**

**Angelo Duò**, SIB Swiss Institute of Bioinformatics, Zurich, Switzerland

Thank you for your comment. SC3 and Seurat show the highest accuracy when the true number of clusters is used as the input parameter. Also, SC3 has the highest median ARI when the estimated number of clusters is imposed. Similarly, using the number of clusters that gives the maximum performance, SC3 evaluates best. These two methods have both the lowest variability within and between the different filterings. Seurat is stable with regards to random starts and at least for the datasets filtered on the average expression and the variability. Finally, whereas Seurat was generally the fastest method, we agree that SC3 is currently not favorable in terms of the runtime or scalability.

**Competing Interests:** No competing interests were disclosed.

---

**Reviewer Report 03 August 2018**

https://doi.org/10.5256/f1000research.17093.r36545

© 2018 Freytag S. This is an open access peer review report distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

**Saskia Freytag**

Department of Medical Biology, University of Melbourne, Parkville, Vic, Australia

**Overview**

The authors present comprehensive benchmarking of clustering tools in R on real and simulated single-cell RNA-seq datasets. Their work includes performance, stability and run time analysis. Furthermore, they also investigate whether combining results from different methods increases performance.
Major comments

- Throughout the entire manuscript the authors should make it clear that only clustering tools available in R were investigated. This is important, as there are quite a number of popular python applications for clustering of single cell RNA-seq data available.
- Like Jean Fan, I am concerned about the appropriateness of the Trapnell et al. dataset and the Zheng et al. 10x datasets. Furthermore for the Zheng et al. dataset, I would like to know why the authors did not use all 10 pre-sorted cell populations available? Furthermore, how did the authors choose which cell populations to combine for their Zhengmix4 and Zhengmix8 datasets?

Minor comments

- The authors show nicely that Seurat is not very strongly affected by gene filtering. Could this be a result of its clustering approach being based on the 500 most variable genes?
- On page 7 in the paragraph “Run Times vary widely between methods” the authors use Adjusted Rand Index instead of its already introduced abbreviation
- Could the size of Figure 5 be increased?
- Why did some methods get raw and some methods log-transformed normalized counts?
- Consider changing Supplementary Figure 2 to a visual representation that represents size differences between sets, like UpSetR plots.
- On page 10 the authors say: "In addition, no apparent association between the similarity of the clusterings and the type of input or dimension reduction or underlying type of clustering algorithm was found." Could the authors explain in more detail how this analysis was performed.
- On page 6, the authors speculate that there are stronger signals that dominate clustering in the Trapnell et al dataset that are not time points. What could these be? Have the authors investigated cell cycle?

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

Competing Interests: No competing interests were disclosed.
I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 31 Aug 2018

Charlotte Soneson, SIB Swiss Institute of Bioinformatics, Zurich, Switzerland

Thank you for reviewing our manuscript and for your constructive comments. Below are point-by-point responses to the individual comments.

Throughout the entire manuscript the authors should make it clear that only clustering tools available in R were investigated. This is important, as there are quite a number of popular python applications for clustering of single cell RNA-seq data available.

This has been clarified in the Abstract as well as in the Methods part of the text. Some of the most widely used clustering methods implemented in Python (e.g., scanpy) implement the same or similar clustering methods as those evaluated in this study, and could thus be considered to be implicitly investigated. Also, the evaluation system we provide (via the code in the GitHub repository and the associated data package) is not strictly limited to methods implemented in R; other methods can be included e.g. using system() calls.

Like Jean Fan, I am concerned about the appropriateness of the Trapnell et al. dataset and the Zheng et al. 10x datasets. Furthermore for the Zheng et al. dataset, I would like to know why the authors did not use all 10 pre-sorted cell populations available? Furthermore, how did the authors choose which cell populations to combine for their Zhengmix4 and Zhengmix8 datasets?

We agree that the Trapnell data set was not generated with the purpose of finding cell types - however, we still find it useful to illustrate the performance of the methods in a data set where the “true clusters” (defined as the time point at which the cells where collected) do not represent the main/strongest signal in the data (see e.g. the t-SNE plots in Supplementary Figure 1). We have clarified this in the “Methods-Real data sets” section of the revised paper.

For the Zhengmix data sets, our aim was to generate data sets with a mix of well-separated (e.g., B-cells vs T-cells) and similar cell types (e.g., different types of T-cells). In addition, we wanted to investigate if the number of cell populations and/or the equality of the population sizes had an impact on the performance. The included cell type combinations were selected to allow us to address these questions; however, given the richness of this data set, there are certainly many more possible combinations to explore. We have expanded the description in the “Methods-Real data sets” section a bit to highlight these goals.

The authors show nicely that Seurat is not very strongly affected by gene filtering. Could this be a result of its clustering approach being based on the 500 most variable genes?

In all our investigations, we preselect the genes that are used as input for each clustering algorithm using three different variable selection methods, and internal variable selection or filtering steps are disabled. Specifically, for Seurat we perform the PCA using all the genes remaining after our filtering, and the clustering is then performed in the principal component space. Thus, the stability of Seurat should be affected in the same way as that of the other methods by the selection of variables.
On page 7 in the paragraph “Run Times vary widely between methods” the authors use Adjusted Rand Index instead of its already introduced abbreviation

*Thanks for noticing this, we now use the abbreviation also here.*

Could the size of Figure 5 be increased?

*We have increased the size of Figure 5B.*

Why did some methods get raw and some methods log-transformed normalized counts?

*The methods are based on different distributional assumptions and underlying models, affecting the type of values that are most suitably used as input. We followed the recommendations of the authors of the respective methods, and the type of input used for each method is summarized in Figure 4.*

Consider changing Supplementary Figure 2 to a visual representation that represents size differences between sets, like UpSetR plots.

*We have replaced the Venn diagrams in Supplementary Figure 2 with UpSet plots.*

On page 10 the authors say: "In addition, no apparent association between the similarity of the clusterings and the type of input or dimension reduction or underlying type of clustering algorithm was found." Could the authors explain in more detail how this analysis was performed.

*This conclusion is drawn based on Figure 4, where no association between the clustering of methods by cluster similarity and any of the method characteristics can be seen. This has been clarified in the “Results-Inconsistent degree of similarity between methods” section of the revised paper.*

On page 6, the authors speculate that there are stronger signals that dominate clustering in the Trapnell et al dataset that are not time points. What could these be? Have the authors investigated cell cycle?

*We have not explicitly investigated the interpretation of the strongest signal in the Trapnell data set. However, Supplementary Figure 1 suggests that the annotation that we used to define the “true” clusters (the time at which the cells were collected) does not fully explain the grouping of the cells in the t-SNE visualization (in particular, the T12 and T24 groups are intermingled). As noted above, the main purpose of including this data set was to investigate the behaviour of the various methods in a data set where the clusters were less apparent.*

**Competing Interests:** No competing interests were disclosed.
Overview

Duo et al compare multiple single-cell RNA-seq clustering approaches on real and simulated single-cell RNA-seq datasets.

Major comments

- Quite a number of single-cell RNA-seq datasets are available for benchmarking but only a few were explored here. While an exhaustive interrogation of all single-cell RNA-seq datasets available is beyond the scope of this paper, it would be worthwhile for the readers if the authors could comment briefly on the appropriateness of the datasets used here in terms of their cell-type diversity or other factors that may impact benchmarking. As the authors note, a method's performance is inherently tied to the degree to which the tested subpopulations are truly (or artificially) transcriptionally distinct. In particular, I am concerned about the appropriateness of the Trapnell dataset, as it was originally intended for pseudotime/trajectory inference and may not even contain discrete transcriptional subpopulations. The poor performance as noted in Figure 1 for this dataset may simply arise from different methods cutting along this continuous trajectory in different ways. Similarly, for the Zheng 10x datasets, since each cell-type was sorted and sequenced separately, there is inevitably some degree of confounding of cell-type specific effects with batch effects that could make clustering much easier.

- As datasets get bigger, the scalability of each method will be an important consideration. The authors provide a preliminary look into this via the different run time of each method in Figure 2, but how this run time depends on the number of cells is unclear. Readers will be interested in whether some methods scale better than others. It is worth having an additional figure of run time as a function of number of cells (via downsampling cells and then extrapolating to larger datasets) to fully capture the scalability of each method.

- With regard to the stability between cluster runs, some methods may internally set various random seeds to ensure reproducibility. Please double check that the stability observed in Figure 3 is not simply the result of which methods uses random seeds. If a method does use an (or likely multiple) internal random seed, the seed must be changed to accurately assess stability.

Minor comments

- There are quite a number of single-cell RNA-seq clustering approaches and the list keeps growing (https://github.com/seandavi/awesome-single-cell). Only a fraction is represented in this comparison. While an exhaustive comparison of all methods is beyond the scope of this paper, the authors should comment briefly on how these particular 12 clustering algorithms were chosen.

- While nearly all methods assessed use dimensionality reduction as a first step, it is unclear why some
were allowed to reduce to 30 dimensions while others 50. It seems that particularly as datasets get larger with presumably more cell-types captured in each datasets, we will likely want to increase the number of PCs to fully capture the variation present in the data. While the authors have left the investigation into the effects of the number of PCs to future research, they should briefly note the reason for the choice of PCs used for each method.

Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Partly

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Partly

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Charlotte Soneson, SIB Swiss Institute of Bioinformatics, Zurich, Switzerland

Thank you for reviewing our manuscript and for your constructive comments. Below are point-by-point responses to the individual comments.

Quite a number of single-cell RNA-seq datasets are available for benchmarking but only a few were explored here. While an exhaustive interrogation of all single-cell RNA-seq datasets available is beyond the scope of this paper, it would be worthwhile for the readers if the authors could comment briefly on the appropriateness of the datasets used here in terms of their cell-type diversity or other factors that may impact benchmarking. As the authors note, a method’s performance is inherently tied to the degree to which the tested subpopulations are truly (or artificially) transcriptionally distinct. In particular, I am concerned about the appropriateness of the Trapnell dataset, as it was originally intended for pseutotime/trajectory inference and may not even contain discrete transcriptional subpopulations. The poor performance as noted in Figure 1 for this dataset may simply arise from different methods cutting along this continuous trajectory in different ways. Similarly, for the Zheng 10x datasets, since each cell-type was sorted and sequenced separately, there is inevitably some degree of confounding of cell-type specific effects with batch
effects that could make clustering much easier.

There is indeed a large (and increasing) number of public scRNA-seq data sets available, generated with many different types of protocols. However, the main issue (especially with droplet-based data sets) is that no independent annotation of the cells is available, which implies that they are not suitable for unbiased benchmarking like we are doing here. Many public droplet-based data sets do contain “cell type labels”, but these are typically inferred by clustering the cells based on the scRNA-seq data itself, and thus any evaluation risks being biased in favor of methods similar to the one used to derive the labels in the first place. This is the main reason behind the selection of these data sets. We agree that the Trapnell data set was not generated with the purpose of finding cell types - however, we still find it useful to illustrate the performance of the methods in a data set where the “true clusters” (defined as the time point at which the cells were collected) do not represent the main/strongest signal in the data (see e.g. the t-SNE plots in Supplementary Figure 1). For the Zheng data set, it’s true that there could be confounding with batch effects, and ambiguous cells may be excluded, which would also make clusters more distinct. For our Zhengmix data sets, we therefore included both very different (e.g., B-cells and T-cells) and more similar (e.g., different types of T-cells) cell types (Supplementary Figure 1). We have expanded the discussion in the “Methods-Real data sets” section of the revised paper to clarify these issues.

As datasets get bigger, the scalability of each method will be an important consideration. The authors provide a preliminary look into this via the different run time of each method in Figure 2, but how this run time depends on the number of cells is unclear. Readers will be interested in whether some methods scale better than others. It is worth having an additional figure of run time as a function of number of cells (via downsampling cells and then extrapolating to larger datasets) to fully capture the scalability of each method.

Thanks for pointing this out. We have included a plot illustrating the scalability, investigated by downsampling of the largest data set, in Supplementary Figure 9.

With regard to the stability between cluster runs, some methods may internally set various random seeds to ensure reproducibility. Please double check that the stability observed in Figure 3 is not simply the result of which methods uses random seeds. If a method does use an (or likely multiple) internal random seed, the seed must be changed to accurately assess stability.

Two of the methods (TSCAN and monocle) set random seeds internally and do not allow these to be changed by the user. Other methods (SC3, Seurat and RaceID2) set a random seed but let the user specify it. For these methods, we explicitly set the random seed to different values in the five runs. We have clarified this in the “Results-High stability between clustering runs” section of the revised text.

There are quite a number of single-cell RNA-seq clustering approaches and the list keeps growing (https://github.com/seandavi/awesome-single-cell). Only a fraction is represented in this comparison. While an exhaustive comparison of all methods is beyond the scope of this paper, the authors should comment briefly on how these particular 12 clustering algorithms were chosen.

The methods were chosen to represent the most common types of algorithms used for clustering of scRNA-seq data. We have tried to include the most widely used methods, but also to include methods from tangential fields as well as more traditional clustering methods to serve as a
baseline. We have clarified this in the text.

While nearly all methods assessed use dimensionality reduction as a first step, it is unclear why some were allowed to reduce to 30 dimensions while others 50. It seems that particularly as datasets get larger with presumably more cell-types captured in each datasets, we will likely want to increase the number of PCs to fully capture the variation present in the data. While the authors have left the investigation into the effects of the number of PCs to future research, they should brief note the reason for the choice of PCs used for each method.

We extracted 50 principal components for the methods that performed an additional dimension reduction (by t-SNE), and 30 principal components for methods where the clustering was done in the principal component space. The only exception was FlowSOM; this was unintentional and has been harmonized in the revised version to use the same number of PCs as the rest of the methods.

**Competing Interests:** No competing interests were disclosed.