clustifyr: an R package for automated single-cell RNA sequencing cluster classification [version 2; peer review: 2 approved]

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Abstract
Assignment of cell types from single-cell RNA sequencing (scRNA-seq) data remains a time-consuming and error-prone process. Current packages for identity assignment use limited types of reference data and often have rigid data structure requirements. We developed the clustifyr R package to leverage several external data types, including gene expression profiles to assign likely cell types using data from scRNA-seq, bulk RNA-seq, microarray expression data, or signature gene lists. We benchmark various parameters of a correlation-based approach and implement gene list enrichment methods. clustifyr is a lightweight and effective cell-type assignment tool developed for compatibility with various scRNA-seq analysis workflows. clustifyr is publicly available at https://github.com/rnabioco/clustifyr

Keywords
Single-cell RNA sequencing, cell type classification, gene expression profile, R package

This article is included in the RPackage gateway.
This article is included in the Bioconductor gateway.

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Amendments from Version 1

The new version of the manuscript includes more detailed introductions and descriptions of the datasets and analyses performed to benchmark clustifyr. Additional functionality in clustifyr is also highlighted, including the ability to identify cell types using marker gene lists and methods to examine cell type classification in the presence of overclustering of the reference or query datasets. Additional software tools were benchmarked against clustifyr and the benchmarking was standardized across multiple datasets for clarity. Lastly we have organized the datasets described in this manuscript into a ExperimentHub resource (clustifyrdatahub) that will be available through Bioconductor. We have included a new table (Table_1.xltx), made edits to Figure 1–Figure 4, and now also provide links to a supplemental table hosted on zenodo (https://doi.org/10.5281/zenodo.3934480).

Any further responses from the reviewers can be found at the end of the article

Introduction

Single-cell mRNA sequencing (scRNA-seq) promises to deliver elevated understanding of cellular mechanisms, cell heterogeneity within tissue, and developmental transitions. A key challenge in scRNA-seq data analysis is the identification of cell types from single-cell transcriptomes. Manual inspection of the expression patterns from a small number of marker genes is still standard practice, which is both cumbersome and potentially inaccurate. Methods that compare cell type expression patterns against robust reference data provide additional confidence in cell type assignments and have the potential to automate and standardize cell type assignment. Unfortunately, current implementations of scRNA-seq suffer from several limitations that further compound the problem of cell type identification. First, only RNA levels are measured, which may not correlate with cell surface marker or gene expression signatures identified through other experimental techniques. Second, due to the low capture rate of RNAs, low expressing genes may face detection problems regardless of sequencing depth. Many previously established markers of disease or developmental processes suffer from this issue, such as transcription factors. On the data analysis front, over or under-clustering can generate cluster markers that are uninformative for cell type labeling. In addition, cluster markers that are unrecognizable to an investigator may indicate potentially interesting unexpected cell types but can be very intimidating to interpret.

For these reasons, investigators struggle to integrate scRNA-seq into their studies due to the challenges of confidently identifying previously characterized or novel cell populations. Formalized data-driven approaches for assigning cell type labels to clusters greatly aid researchers in interrogating scRNA-seq experiments. Currently, multiple cell type assignment packages exist but they are specifically tailored towards input types or workflows. Seurat, a popular toolkit for single cell RNA-seq analysis, implements a mutual nearest neighbor-based method to annotate cell types using another single cell RNA-seq dataset in the Seurat object format. SingleR and scmap provide functionality within the Bioconductor framework to annotate cell types using correlation if provided a reference from bulk-RNA-seq or averaged single cell cluster data. scPred also uses a Bioconductor framework and applies a Support Vector Machine (SVM) model to PCA reduced gene expression data to classify cell types. ACTINN, a neural network-based annotation tool, also relies on existing single cell reference data and operates on files within a command line framework. As more and more approaches to the classification problem are introduced, benchmarking performance and compatibility to sequencing platforms and analysis pipelines becomes increasingly important.

We developed the R package clustifyr, a lightweight and flexible tool that leverages a wide range of prior knowledge of cell types to pinpoint target cells of interest or assign general cell identities to difficult-to-annotate clusters. Here, we demonstrate its basic usage and applications with transcriptomic information of external datasets and/or signature gene profiles, to explore and quantify likely cell types. The clustifyr package is built with compatibility and ease-of-use in mind to support other popular scRNA-seq tools and formats.

Methods

Implementation

clustifyr requires query and reference data in the form of normalized expression matrices, corresponding metadata tables, and a list of variable genes.

```r
library(clustifyr)
library(rnacular)

pbmc_matrix_small[1:5, 1:5]  # query matrix of normalized scRNA-seq counts
cbmc_ref[1:5, 1:5]  # reference matrix of expression for each cell type
pbmc_meta[1:5, ]  # query meta-data data.frame containing cell clusters
length(pbmc_markers_M3Drop$Gene)  # vector of variable genes
```
clustifyr adopts correlation-based methods to find reference transcriptomes with the highest similarity to query cluster expression profiles, defaulting to Spearman ranked correlation, with options to use Pearson, Kendall, or Cosine correlation instead if desired. clustify() will return a matrix of correlation coefficients for each cell type and cluster, with the row names corresponding to the query cluster number and column names as the reference cell types.

```r
res <- clustify(  
  input = pbmc_matrix_small,  
  metadata = pbmc_meta,  
  cluster_col = "seurat_clusters", # column in meta.data with clusters  
  ref_mat = cbmc_ref,  
  query_genes = pbmc_markers_M3Drop$Gene
)
```

Query clusters are assigned cell types to the highest correlated reference cell type, with an automatic or manual cutoff threshold. Query clusters dissimilar to all available reference cell types are labeled as “unassigned”.

Figure 1. Schematic for clustifyr input and output.
res2 <- cor_to_call(
    cor_mat = res,  # matrix of correlation coefficients
    cluster_col = "seurat_clusters",  # column in meta.data with clusters
    threshold = 0.5
)

To better integrate with standard workflows that involve S3/S4 R objects, methods for clustifyr are written to
directly recognize Seurat\textsuperscript{14} (v2 and v3) and SingleCellExperiment\textsuperscript{15} objects, retrieve the required information,
and reinsert classification results back into an output object. A more general wrapper is also included for compat-
ibility with other common data structures and can be easily extended to new object types. This approach also has
the added benefit of forgoing certain calculations such as variable gene selection or clustering, which may already
be stored within input objects.

res <- clustify(
    input = sce_small,  # an SCE object
    ref_mat = cbmc_ref,  # matrix of expression for each cell type
    cluster_col = "cell_type1",  # column in meta.data with clusters
    obj_out = TRUE  # output SCE object with cell type
)

SingleCellExperiment::colData(res)[1:10, c("type", "r")]
#> DataFrame with 10 rows and 2 columns
#>   type                r
#> A2_A1       pDCs 0.814336567702192
#> A2_A10      Eryth 0.665800619720566
#> A2_A11       pDCs 0.682088309107356
#> A2_A12      Eryth 0.665800619720566
#> A2_A2         B  0.634114583333333
#> A2_A3       pDCs 0.814336567702192
#> A2_A4       pDCs 0.814336567702192
#> A2_A5          NK 0.655407634437123
#> A2_A6       pDCs 0.682088309107356
#> A2_A7       pDCs 0.71424223704931

res <- clustify(
    input = s_small3,  # a Seurat object
    ref_mat = cbmc_ref,  # matrix of expression for each cell type
    cluster_col = "RNA_snn_res.1",  # name of column in meta.data containing cell
    clusters
    obj_out = TRUE  # output Seurat object with cell type inserted
    as "type" column
)

res@meta.data[1:5, ]  # output Seurat object with cell type inserted
#>          orig.ident nCount_RNA nFeature_RNA RNA_snn_res.0.8
#> ATGCCAGAACGACT SeuratProject           70           47               0
#> CATGGGCTCTGCATT SeuratProject           85           52               0
#> GAACCTGTAGAAC SeuratProject            87           50               1
#> TGACACTGGATTCTCA SeuratProject         127           56               0
#> AGTCAGACTGCACA SeuratProject          173           53               0
#>                letter.idents groups RNA_snn_res.1 type         r
#> ATGCCAGAACGACT A     g2          0   Mk 0.6204476
#> CATGGGCTCTGCATT A     g1          0   Mk 0.6204476
#> GAACCTGTAGAAC B     g2          0   Mk 0.6204476
#> TGACACTGGATTCTCA A     g2          0   Mk 0.6204476
#> AGTCAGACTGCACA A     g2          0   Mk 0.6204476
In the absence of suitable reference data (i.e. RNA-seq or microarray expression matrices), clustifyr can build scRNA-seq reference data by averaging per-cell expression data for each cluster, to generate a transcriptomic snapshot. Direct reference-building from SingleCellExperiment or Seurat objects is supported as well.

```r
new_ref_matrix <- average_clusters(
  mat = pbmc_matrix_small,
  metadata = pbmc_meta$classified,  # or use metadata = pbmc_meta, cluster_col = "classified"
  if_log = TRUE  # whether the expression matrix is already log transformed
)
new_ref_matrix_sce <- object_ref(
  input = sce_small,  # SCE object
  cluster_col = "cell_type1"  # column in colData with cell identities
)
new_ref_matrix_v3 <- seurat_ref(
  seurat_object = s_small3,  # SeuratV3 object
  cluster_col = "RNA_snn_res.1"  # column in meta.data with cell identities
)
```

Data exploration plotting functions, for dimensional reduction scatter plots and heatmaps, are extended from ggplot2 and ComplexHeatmap packages, featuring colorblind-friendly default colors. Gene list-based methods (clustify_lists()) are also implemented via hypergeometric tests, GSEA, jaccard index, or percentage gene detection by cluster, which provide easy to interpret methods to verify the presence of known positive and negative marker genes.

**Parameters**

**Reference datasets.** Multiple scRNA-seq and other cell type references datasets are provided in an ExperimentHub Bioconductor package (clustifyrdatahub). A description of these datasets and others used for benchmarking and optimizing parameters for clustifyr are provided in Table 1.

**Correlation method.** We benchmarked clustifyr against a suite of comparable datasets, PBMC-bench\textsuperscript{13,16}, generated using multiple scRNA-seq methods on aliquots of peripheral blood mononuclear cells (PBMCs) from two individuals. Additional details about each query and reference dataset are provided in Supplemental Table 1. For each single cell technology, average gene expression profiles were generated from annotated cell types and compared across each platform. Notably, for each reference dataset cross-referenced against all other samples, clustifyr achieved a median F1-score (see Benchmarking Methods) of above 0.94 using Spearman ranked correlation (Figure 2A). Other correlation methods are on par or slightly worse at cross-platform classifications, which is expected based on the nature of ranked vs unranked methods. We therefore selected Spearman as the default method in clustifyr, with other methods also available, as well as a wrapper function to find consensus identities across available correlation methods (call_consensus()).

**Correlation minimum cutoff.** Recognition of missing reference cell types, so as to avoid misclassification, is another point of great interest in the field. From general usage of clustifyr, we find using a minimum correlation cutoff of 0.5 or 0.4 is generally satisfactory. Alternatively, the cutoff threshold can be determined heuristically using 0.8 \* highest correlation coefficient among the clusters. One example is shown in Figure 2B, using PBMC rejection benchmark data modified by the SciBet package\textsuperscript{17}. Megakaryocytes were removed from the reference melanoma immune cells data, but retained in the test data to mimic the situation when the reference data does not contain a rare cell type. clustifyr analysis successfully found the megakaryocytes to be dissimilar to all available reference cell types, and hence left as “unassigned” under the default minimum threshold cutoff.

**Variable gene selection and normalization.** As the core function of clustifyr is ranked correlation, feature selection to focus on highly variable genes is critical. To illustrate the importance of feature selection we used clustifyr to classify pancreatic cell types generated using the inDrops platform using a reference built from a dataset generated on the Smart-Seq2 platform\textsuperscript{18,19}. In Figure 2C, we compare correlation coefficients using all detected genes (>10,000) vs feature selection by the package M3Drop. A basic level of feature selection, e.g. using M3Drop, Seurat VST (default uses top 2,000 variable genes), or simply 1,000 genes with the highest variance in the reference data, is sufficient to classify the pancreatic cells. In the case of other cell type mixtures, especially ones without complete knowledge of the expected cell types, further optimization of clustering and feature selection may be
## Table 1. Collection of datasets used for introducing and benchmarking clustifyr.

A description of single cell RNA-seq, bulk RNA-seq, and microarray datasets used in this study. The datasets available through ExperimentHub are references that were built from raw or downloaded data and can be used with clustifyr. R objects can be accessed using the direct download URLs to the .rda files, or through the clustifyrdatahub ExperimentHub.

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<th>Description</th>
<th># of cell types</th>
<th>Organism</th>
<th>Publication</th>
<th>Source</th>
<th>Data Provider</th>
<th>R object download URL</th>
<th>Bioconductor ExperimentHubID</th>
<th>R object name</th>
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<td>figshare</td>
<td><a href="https://github.com/rnabioco/clustifyrdata/raw/master/data/ref_tabula_muris_facs.rda">link</a></td>
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<td>ref_mouse.rnaseq</td>
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**Data Provider:**
- Bioconductor
- ExperimentHubID
- S3
- ZenoDB
- Allen Brain Institute
- 10x Genomics

**Publication:**
- https://www.cell.com/fulltext/S2405-4712(16)30266-6
- https://doi.org/10.1186/s13059-019-1795-z
- https://doi.org/10.1038/s41586-018-0654-5
- https://doi.org/10.1038/ncomms1404
- https://support.10xgenomics.com/single-cell-gene-expression/datasets

**Source:**
- https://www.cell.com/fulltext/S2405-4712(16)30266-6
- https://doi.org/10.1186/s13059-019-1795-z
- https://doi.org/10.1038/s41586-018-0654-5
- https://doi.org/10.1038/ncomms1404
- https://support.10xgenomics.com/single-cell-gene-expression/datasets

**R object download URL:**
- https://github.com/rnabioco/clustifyrdata/raw/master/data/ref_pan_indrop.rd
- https://github.com/rnabioco/clustifyrdata/raw/master/data/ref_pan_smartseq2.rd
- https://zenodo.org/record/3357167/files/scRNAseq_Benchmark_datasets.zip?download=1
- https://zenodo.org/record/3357167/files/scRNAseq_Benchmark_datasets.zip?download=1

**R object name:**
- rel_pan_indrop
- rel_pan_smartseq2
Figure 2. Parameter considerations for clustifyr. A) Comparison of median F1-scores of different correlation methods for classifying across platforms using the PBMC-bench dataset. B) Heatmap showing correlation coefficients between query cell types and the reference cell types from a rejection test, whereby megakaryocytes were excluded from the reference dataset. The Neg.Cell cluster is megakaryocytes, which is correctly not annotated a different cell type when megakaryocytes are not present in the reference. By default clusters with correlation < 0.50 are assigned as “unassigned” by clustifyr. C) Comparison of correlation coefficients with and without feature selection when comparing average gene expression per cell type between two pancreas scRNA-seq datasets. The “unclassified” cell type was not defined in the Segerstolpe et al. dataset. D) Accuracy (defined as the ratio between the number of correctly classified clusters and the overall number of clusters) and performance were assessed with decreasing query cluster cell numbers using the Tabula Muris as the query dataset and the Mouse cell atlas as the reference dataset. E) Example of overclustering the query data and assigning cell types for data exploration. UMAP of PBMC dataset generated by 10x Genomics with cell types assigned by comparing to reference data from CBMC cells from Stoeckius et al. 2017. F) An assessment of the median F1-score when using single or multiple averaged profiles as reference cell types was conducted using the PBMC-bench test set. The number of reference expression profiles to generate for each cell type is determined by the number of cells in the cluster (n), and the sub-clustering power argument (x), with the formula n^x.
of greater importance. clustifyr does not provide novel clustering, feature selection, or normalization methods on its own, but instead is built to maintain flexibility to incorporate methods from other, and future, packages. We recommend that users use normalized reference and query data and match normalization methods between datasets when possible. We view these questions as fast-moving fields, and hope to benefit from new advances, while keeping the general clustifyr framework intact.

**Minimum cells per cluster.** We next applied clustifyr to a larger general reference set built from the Mouse Cell Atlas and examined cell type classification of another mouse cell atlas, the Tabula Muris dataset. clustifyr assigned cell types with a median accuracy of 1. Using these test datasets we sought to determine the minimum number of query cells necessary in a cluster to obtain accurate cell type annotation. We subsampled the query data (Figure 2D) and as expected, with further downsampling of the number of cells in each query cluster, we observe decreased accuracy. Yet, even at 15 cells per tested cluster, clustifyr still performed well, with a further increase in speed. Based on these results, we set the default parameters in clustifyr to exclude or warn users of classification on clusters containing less than 10 cells. These results also suggest that clustering the query dataset to obtain more refined clusters (e.g., fewer cells per cluster) could be employed to aid in the identification of rarer or less well-defined cell subsets. clustifyr can also be used to classify individual cells, although we do not recommend per cell classification because of the reduced accuracy observed with decreasing numbers of cells per cluster.

**Subclustering.** clustifyr also provides functionality to assess the quality of the cell type annotations. An intentional overclustering and classification function based on k-means clustering (overcluster_test()) is implemented in clustifyr for exploration of cell type annotation at increasing numbers of clusters (Figure 2E). This approach provides a rapid visualization to determine if cell type annotations are stable with varying numbers of clusters. For example, scRNA-seq data from the Seurat PBMC 3k tutorial was reclassified at multiple clustering levels using Cord Blood Mononuclear Cells (CBMCs) as reference, which demonstrated largely stable cell type assignments in the presence of overclustered query data (Figure 2E)\(^9\). When using scRNA-seq data as the reference data, matrices are built by averaging per-cell expression data for each cluster (average_clusters()), to generate a transcriptomic snapshot similar to bulk RNA-seq or microarray data. An additional argument to subcluster the reference single cell clusters is also available, to generate more than one expression profile per reference cell type, in a manner analogous to overcluster_test(), but applied to the reference scRNA-seq dataset. The number of subclusters for each reference cell type is dependent on the number of cells in the cluster (n), and the sub-clustering power argument (x), following the formula \(n^{x}\). This approach does not improve classification in the PBMC-bench data (Figure 2F), whose reference and query clustering are already consistent. However, we envision its utility would greatly depend on the granularity of the clustering in the reference dataset.

**Benchmarking**

Using clustifyr, PBMC clusters from the Seurat PBMC 3k tutorial are correctly labeled using either bulk-RNA seq references generated from processed microarray data of purified cell types\(^9\), the ImmGen database of bulk-RNA-seq\(^21,22\), or previously annotated scRNA-seq results from the Seurat CBMC CITE-seq tutorial\(^14,23\) (Figure 3).

To assess the performance of clustifyr, we used the Tabula Muris dataset, which contains data generated from 12 matching tissues using both 10x Genomics 3’ end seq (“drop”) and Smart-Seq2 (“facs”) platforms. We attempted to assign cell type identities to clusters in “drop” Seurat objects using references built from “facs” Seurat objects, which contain pre-computed variable genes generated by the Seurat mean.var.plot (dispersion z-scores based on expression bins) approach. For each method we used the recommended variable gene selection approach. clustifyr uses variable genes supplied by the user and for benchmarking we used the variable genes stored in the Seurat object. smap calculates variable genes using a modified approach based on M3drop. SingleR selects variable genes by identifying marker genes between clusters. scPred, in contrast, selects informative principal components as a feature selection procedure whereas ACTINN does not perform feature selection for classification.

In benchmarking results, clustifyr is comparably accurate versus other automated classification packages (Figure 4A). Cross-platform comparisons are inherently more difficult, and the approach used by clustifyr is aimed at being platform- and normalization-agnostic. Mean runtime, including both reference building and test data classification, in Tabular Muris classifications was ~ 1 second if the required variable gene list is extracted from the query Seurat object. Alternatively, variable genes can be recalculated by other methods such as M3Drop, to reach similar results (clustifyr (m3drop)).
Figure 3. clustifyr can utilize multiple reference data types. UMAP projections of PBMCs showing the ground truth cell types (A), or cell types called by clustifyr using microarray data from sorted immune cell types (B), bulk RNA-seq from immune cell populations (C) or scRNA-seq data from CBMCs (D).

Signature marker gene lists are an additional reference data type that is commonly used to guide cluster cell type classification. We therefore sought to determine if a gene list enrichment approach could provide comparable classification power to using correlation. clustifyr provides a function clustify_lists() which compares marker genes between query clusters to a list of marker genes per reference cell type. clustify_lists will calculate enrichment with a hypergeometric test, marker overlap with the jaccard index, or use the percent of cells expressing marker genes to annotate cell types. Alternatively, if ranked gene lists are available, Gene Set Enrichment Analysis (GSEA) using the fGSEA package or Spearman ranked correlation can be employed. We find that using gene expression for clustifyr classification had higher accuracy than gene list enrichment using a hypergeometric test or the jaccard index, however this approach could be very useful for datasets without scRNA or bulk RNA-seq data for use as a reference. (Figure 4A).

For scalability benchmarking, we adapted an existing benchmark dataset, scRNAseq_Benchmark subsampling, which contains query and reference data with downsampled numbers of cells from the Smart-seq2 Tabula Muris dataset\(^1\). Once again, clustifyr is accurate and efficient, compared to other developed methods (Figure 4B). As a further comparison, we also examined classification of cell types in murine brain datasets generated by the Allen Institute Brain Atlas, and provided by the scRNAseq_Benchmark pipeline\(^1\). The two murine brain regions contained 34 shared cell types and clustifyr was also able to reach similarly satisfactory cell annotation compared to other annotation methods. (Figure 4C).
**Figure 4. clustifyr accurately and rapidly annotates cell types.**

A) Accuracy and run-time of classifications generated by clustifyr or existing methods using the Tabula Muris dataset to benchmark cell type classifications between datasets generated with the Smart-Seq2 or 10x Genomics sequencing platforms. Each point represents a different tissue comparison. clustifyr (m3drop) indicates clustifyr run using variable genes defined by M3drop, clustifyr_lists (hyper) uses hypergeometric tests to compare marker gene lists, and clustifyr_lists (jaccard) calculates the jaccard index between marker gene lists to annotate cell types.

B) Performance comparison of clustifyr to existing methods with random subsamples of cells from the Smart-Seq2 Tabula Muris dataset. Error bars represent standard error of the mean and are derived from 5 independent subsamples of the dataset.

C) Performance comparison of clustifyr to existing methods testing classification of an Allen Institute Brain Atlas dataset from two murine brain regions that contain 34 cell types. scPred is not shown as it failed with an error on this dataset.

D) Comparing clustifyr to existing methods for rejecting unseen populations using PBMC data. Three reference PBMC datasets were generated that excluded either T-cells, CD4+ T-cells or memory T-cells respectively. The % of rejected indicates the % of the indicated cell type that was not misclassified when the cell type was missing from the reference.
Lastly, we applied clustifyr to a series of increasingly challenging datasets from the scRNAseq_Benchmark\textsuperscript{13} unseen population rejection test (Figure 4D). This test assess how frequently cells will be mis-assigned when the corresponded cell types are not present in the dataset. The PBMC dataset contains different T-cell subsets, which do not often cluster into discrete well-defined cell types solely based on gene expression. Without the corresponding cell type references, 57.5% of T cells were rejected and unassigned. When only CD4+ references were removed, 28.2% of test CD4+ T cells were rejected and unassigned. clustifyr was unable to reject CD4+/CD45RO+ memory T cells, mislabeling them as CD4+/CD25 T Reg instead when the exact reference was unavailable. However, these misclassifications are also observed with other classification tools benchmarked in the scRNAseq_Benchmark study (Figure 4D)\textsuperscript{13}.

**Benchmarking methods**

clustifyr was tested against scmap v1.8.0\textsuperscript{7}, SingleR v1.0.1\textsuperscript{13}, Seurat v3.1.1\textsuperscript{14}, latest GitHub versions of ACTINN\textsuperscript{11} and scPred\textsuperscript{5}, and SVM as implemented in python3 scikit-learn v0.19.1\textsuperscript{20}. scRNA-seq Tabula Muris data was downloaded as seuratV2 objects. Human pancreas data was downloaded as SCE objects. In all instances, to mimic the usage case of clustifyr, clustering and dimension reduction projections are acquired from available metadata, in lieu of new analysis.

An R script was modified to benchmark clustifyr following the approach and datasets of scRNAseq_Benchmark\textsuperscript{13}, using M3Drop\textsuperscript{13} to generate variable genes for clustifyr. R code used for benchmarking, and preprocessing of other datasets, in the form of matrices and tables, are documented in R scripts available in the clustifyr and clustifyrdatahub GitHub repositories.

Classification accuracy was measured using two approaches depending on the datasets compared. For datasets where the query and reference data contain identical cell types, an F1-score, the harmonic mean of the precision and recall, was calculated for each cell type (PBMC-bench, Allen Brain Institute Atlas, and Smart-Seq2 Tabula Muris subsampling). When summarizing classification accuracy across an entire dataset the median F1-score is reported. Datasets with varying cell types in the query and reference data cannot be characterized with an F1-score and instead accuracy, defined as the ratio between the number of correctly classified clusters and the overall number of clusters, is reported (Mouse cell atlas vs. Tabula Muris and Tabula Muris Smart-Seq2 vs. 10x Genomics).

**Operation**

clustifyr is distributed as part of the Bioconductor R package repository and is compatible with Mac OS X, Windows, and major Linux operating systems. Package dependencies and system requirements are documented in the clustifyr Bioconductor repository.

**Conclusions**

We present a flexible and lightweight R package for cluster identity assignment. The tool bridges various forms of prior knowledge and scRNA-seq analysis. Reference sources can include scRNA-seq data with cell types assigned (or average expression per cell type, which can be stored at much smaller file sizes), sorted bulk RNA-seq, and microarray data. clustifyr, with minimal package dependencies, is compatible with a number of standard analysis workflows such as Seurat or Bioconductor, without requiring the user to perform the error-prone process of converting to a new scRNA-seq data structure and can be easily extended to incorporate other data storage object types. clustifyr is designed to perform classification after previous steps of analysis by other informatics tools. Therefore, it relies on, and is agnostic to, common external packages for cell clustering and variable feature selection. We envision it to be compatible with all current and future scRNA-seq processing, clustering, and marker gene discovery workflows. Benchmarking reveals the package performs well in mapping cluster identity across different scRNA-seq platforms and experimental types. As we and others observe\textsuperscript{52}, novel algorithms may not be necessary for cell type classification, at least within the current limitations of sequencing technology and our broad-stroke understanding of cell “types”. Rather, the generation of community curated reference databases is likely to be critical for reproducible annotation of cell types in scRNA-seq datasets.

On the user end, clustifyr is built with simple out-of-the-box wrapper functions, sensible defaults, yet also extensive options for more experienced users. Instead of building an additional single-cell-specific data structure, or requiring specific scRNA-seq pipeline packages, it simply handles basic data.frames (tables) and matrices (Figure 1). Input query data and reference data are intentionally kept in expression matrix form for maximum flexibility, ease-of-use, and ease-of-interpretation. Also, by operating on predefined clusters, clustifyr has high scalability and minimal resource requirements on large datasets. Using per-cluster expression averages results in rapid classification. However, cell-type annotation accuracy is therefore heavily reliant on appropriate selection of the number of clusters. Users are therefore encouraged to explore cell type annotations derived from multiple
clustering settings. Additionally, assigning cell types using discrete clusters may not be appropriate for datasets with continuous cellular transitions such as developmental processes, which are more suited to trajectory inference analysis methods. As an alternative, clustifyr also supports per-cell annotation, however the runtime is greatly increased and the accuracy of the cell type classifications are decreased due to the sparsity of scRNA-seq datasets, and requires a consensus aggregation step across multiple cells to obtain reliable cell type annotations.

To further improve the user experience, clustifyr provides easy-to-extend implementations to identify and extract data from established scRNA-seq object formats, such as Seurat\textsuperscript{14}, SingleCellExperiment\textsuperscript{15}, URD\textsuperscript{16}, and CellDataSet (Monocle)\textsuperscript{30}. Available in flexible wrapper functions, both reference building and new classification can be directly achieved through scRNA-seq objects at hand, without going through format conversions or manual extraction. The wrappers can also be expanded to other single cell RNA-seq object types, including the HDF5-backed loom objects, as well as other data types generated by CITE-seq and similar experiments\textsuperscript{31}. Tutorials are documented online to help users integrate clustifyr into their workflows with these and other bioinformatics software.

**Software availability**

clustifyr is available from Bioconductor: https://bioconductor.org/packages/release/bioc/html/clustifyr.html

Up-to-date source code, and tutorials are available from: https://github.com/rnabioco/clustifyr

Package documentation is also provided at: https://rnabioco.github.io/clustifyr/

 Archived source code as at time of publication and Supplemental Table 1 detailing datasets used in each analysis are available from:

https://doi.org/10.5281/zenodo.3934480\textsuperscript{32}

Data used in examples and additional prebuilt references available from: https://github.com/rnabioco/clustifyrdatahub

License: MIT

**Data availability**

Original raw data used in benchmarking is available from the following sources and additionally described in Table 1.

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<th>Dataset</th>
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Acknowledgements
A previous version of this article is available on bioRxiv: https://doi.org/10.1101/855064.

References

Open Peer Review

Current Peer Review Status: ✔️ ✔️

Version 2

Reviewer Report 24 July 2020

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✔️ Kamil Slowikowski
Massachusetts General Hospital, Boston, MA, USA

Thank you for improving the manuscript!

I have no further comments.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Bioinformatics, computational biology, immunogenomics, scRNA-seq.

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 17 July 2020

https://doi.org/10.5256/f1000research.27827.r67325

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✔️ Keegan Korthauer
1 BC Children's Hospital Research Institute, Vancouver, BC, Canada
2 University of British Columbia, Vancouver, BC, Canada

The authors have addressed all comments and concerns.

Competing Interests: No competing interests were disclosed.
Reviewer Expertise: Statistical genomics, bioinformatics

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.

Kamil Slowikowski
Massachusetts General Hospital, Boston, MA, USA

The authors describe an R package for annotating cell clusters in scRNA-seq datasets. Specifically, the package implements code for computing correlations between the columns of two data matrices. They show that high correlations between unknown cell clusters in the first data matrix and annotated cell types in the second matrix can be used to label the unknown cell clusters. They try varying parameters and show the effects on the results, and they also benchmark the time and accuracy compared to other packages designed to annotate scRNA-seq data.

Details of the code, methods, and analyses are partly provided. Some details seem to be missing (e.g. the functionality for gene lists).

The conclusions about the tool and its performance are partly supported by the findings presented in the article. Some terms such as "medF1-score" and "accuracy" are left undefined, and some results omit some methods (Figure 4A has different methods than B or C). Readers may have difficulty understanding the specific questions that were asked and what results are shown.

Main comments:

1. The clarity of the manuscript can be increased by adding more verbose details about all analyses. Please consider expanding details about each question, the approach, the datasets used, and the results.

2. Please consider adding a table describing the reference datasets used in this article, just like the one shown on one of your GitHub repositories. This should help to summarize which datasets were used for the analyses in this article.

Comments about specific parts of the manuscript are below. Excerpts from the article are shown in "quoted italics" after a bullet point, and my comments are shown directly below the bullet point.

- "A key challenge in scRNA-seq data analysis is the identification of cell types from single-cell..."
transcriptomes. Manual inspection of the expression patterns from a small number of marker genes is still standard practice, which is cumbersome and frequently inaccurate.”

Do we know the accuracy by manual inspection? Is there a reference for this? In the absence of evidence, you might consider weakening the statement to say "may be inaccurate" rather than "is cumbersome and frequently inaccurate". You might consider that many scRNA-seq experiments are done for the purpose of discovering new cell types that have not been well-described in previous published datasets. In this setting, manual inspection is necessary and automated analyses could be inaccurate or misleading.

- “Currently, multiple cell type assignment packages exist but they are specifically tailored towards input types or workflows.”

Please consider naming and describing each method that will be compared to clustifyr in this manuscript, so the reader can assess how the methodology of clustifyr compares to other methods. Which methods are "specifically tailored towards input types or workflows"? Could you give an example to help the reader understand this claim?

**Suggested improvements for Figure 1:**
- In Figure 1, you might consider showing the dimensions of the inputs and outputs. This might help the reader to understand how they relate to each other.
- Should the query and reference data be counts? CPM? Or Log2(CPM + 1)? You might consider elaborating on this.

**Suggested improvements for Figure 2:**
- Please consider rotating Figure 2A, D, and E 90 degrees clockwise to improve legibility.
- Please consider limiting the axes ranges to the data instead of using the range [0, 1].
- Please consider increasing all font sizes in all panels in all figures, including titles, legends, axis text, etc. Some readers might need larger sizes to see clearly.
- Please consider changing the title to "All genes (n = 10,000)" and "M3Drop variable genes (n = 1,000)" in Figure 2C, so we have some sense of the number of genes used to generate each heatmap.
- Please consider showing a graphical representation of the experiment setup for this figure. What is the reference? What is the query? What are their dimensions? What is the main question in this analysis?
- One way to enhance clarity is to add descriptive titles to every figure in every panel (e.g. "Testing different correlation statistics", etc.).
- Please consider adding more details to the legend text for Figure 2 to help readers understand exactly what experiment has been done, what data was used, and what result is shown.
- In Figure 2C, it seems that the y-axis and x-axis have been swapped by mistake. I see that
the y-axis is labeled "ground truth cell type" but it includes "unclassified". I would expect the
category "unclassified" to appear in the "called cell type" axis, but not in the "ground truth

cell type" axis. Are the axes swapped or are they correct? Could you please clarify?

○ In Figure 2E, what does the color indicate? Is it the power argument "n^x" or something

else?

○ The reader may be wondering:
  ○ How many query cells did you use?
  ○ How many clusters were in the query dataset? How many cells per cluster?
  ○ How many reference datasets were used?
  ○ How many clusters were in the reference dataset?
  ○ Were the query and reference datasets acquired from the same tissue sample or

were they completely independent and unrelated?

In the section "Subclustering", please consider adding more details to help the reader avoid
misunderstandings. What exactly is the "sub-clustering power argument (x)"? Please consider
giving a concrete example to help the reader understand this section. Please consider creating a
new figure that helps the reader to understand the "subcluster()" functionality.

What is the PBMCbench data? Is this the same data as mentioned in the section "Correlation
minimum cutoff"?

In the section "Cells per cluster", you might consider introducing the dataset, then introducing the
question that is being addressed, and finally reporting the results. What is the number (15, 8, 4)? Is
the "Mouse Cell Atlas" the same as the "Tabula Muris"? Were these mouse datasets used in the
previous sections? The reader might benefit from an introduction of these datasets.

Suggested improvements for Figure 3:

○ Please consider adding labels "A", "B", "C", "D" to mark each of the four panels, so they can

be referenced clearly.

○ Please consider using the same name consistently in the text and the figure titles. For
example, the figure says "Bulk RNA-seq reference data" but the text says "ImmGen
database". The reader might better understand the results if the same label were used in
both places instead of using two different labels for the same thing.

○ Please consider including the identifiers for readers who wish to find these datasets and
download them. For example, if the datasets are available on NCBI GEO, please consider
including the accession numbers directly in the legend text, or in a table. Check to see if any
other database provides an accession number. If an accession number is not available,
please consider providing the DOI for a publication or a URL for a website that provides the
data. By the way, if any data you are using is not deposited to a permanent repository,
please consider uploading this data to a permanent repository (e.g. Figshare).

In the section describing Figure 4A, please consider these suggested changes:
Please explain what is "clustifyr", "clustifyr_lists", and "clustifyr_m3drop".

How was feature selection performed for each analysis in Figure 4A?

What is the strategy used by scmap?

What is the strategy used by "Seurat"?

What is the strategy used by "SingleR"?

How is clustifyr similar or different?

This section says "Correlation-based clustifyr classification performed better than hypergeometric-based gene list enrichment as implemented in clustify_lists." Please consider explaining the "clustify_lists" algorithm in detail and also consider sharing the quantification of the performance of each approach so the reader can interpret the claim "performed better". Also consider elaborating on "performed better".

What is "scRNAseq_Benchmark subsampling"? Could you elaborate on what this is and why it was used?

Suggested improvements for Figure 4:

Please consider including an overview schematic to help the reader understand which datasets were used for each result.

Please define "accuracy". What is the algorithm for computing this number?

Please define "medF1-score". What is the algorithm for computing this number?

For the lower half of panel B, please consider using a format similar to the one in Figure 2B from Kiselev et al. (2018). For example, please use a log10 axis for time, so readers can see the difference between methods.

Why is "medF1-score" used for Figure 4C and "accuracy" for Figure 4B?

Why does Figure 4A have 6 methods, Figure 4B have 5 methods, and Figure 4C have 3 methods? Is it possible to include all 6 methods for all panels? Could you please comment on the reasons for excluding or including methods in each analysis?

"As we and others observe\textsuperscript{25}, novel algorithms may not be necessary for cell type classification, at least within the current limitations of sequencing technology and our broadstroke understanding of cell “types”. Rather, the generation of community curated reference databases is likely to be critical for reproducible annotation of cell types in scRNA-seq datasets."

I agree that a community curated reference database would be a valuable contribution to the field. You might consider creating a table or other type of descriptive listing that helps the reader to understand all of the references that were used in this article. Consider including tissue source, healthy or disease status, number of cells and genes, technology used for the assay, DOI, data URL, NCBI GEO accession, or any other details that the reader might find helpful.

Thank you for providing a GitHub repository with data files! Please also consider sharing the same data in compressed plain text format (e.g. "file.tsv.gz"). In addition to GitHub, please consider using a specialty service that is funded for the purpose of permanently archiving research data such as NIH Figshare (https://nih.figshare.com). There are other options (Zenodo, Open Science Framework OSF, etc.).

"As an alternative, clustifyr also supports per-cell annotation, however the runtime is greatly
increased and the accuracy of the cell type classifications are decreased due to the sparsity of scRNA-seq datasets, and requires a consensus aggregation step across multiple cells to obtain reliable cell type annotations."

You might consider offering another alternative option. One extreme is to use the cluster averages, while the other extreme is to use single cells. Perhaps there might be a middle ground where clustifyr could automatically use k-means or some other algorithm to form clusters within the user-defined clusters. This would give the user even more flexibility.

After reviewing the code, I can see that there is an "overcluster()" function that seems to do exactly what I suggested. Please consider describing this in the article and showing an example of how it works. In retrospect, I can see that the section titled "Subclustering" was supposed to describe this topic — I misunderstood this section on the first read.

You may want to double-check all of the links in all of your HTML pages. I see three URLs:

- [https://github.com/rnabioco/clustifyrdatahub/](https://github.com/rnabioco/clustifyrdatahub/)
- [https://github.com/rnabioco/clustifyr](https://github.com/rnabioco/clustifyr)
- [https://github.com/rnabioco/clustifyrdata](https://github.com/rnabioco/clustifyrdata)

I can see that the "clustifyrdatahub" repo has code for creating ".rda" files from the reference datasets.

I also see similar scripts at [https://github.com/rnabioco/clustifyrdata/tree/master/data-raw](https://github.com/rnabioco/clustifyrdata/tree/master/data-raw)

Readers might be confused when they see two different repos with similar scripts. You might consider deleting the "clustifyrdatahub" repo if it is not necessary.

I'm happy to see that the data is organized and annotated in the GitHub repo. Specifically, in the GitHub "clustifyrdata" repo, in the "README.md" file, the table shows the name of the reference, the number of cell types, the number of genes, the organism, and a link to the publication. Please consider adding some version of this table to the article, so the reader can understand the scope of this article.

After reviewing the code, I was able to resolve some of my misunderstandings caused by lack of clarity in the terse descriptions in this article. To reduce the chance of misunderstanding by other readers, you might consider clarifying or adding details to the descriptions of functions and results. For example, the article does not mention that GSEA is used to work with gene lists.

References


Is the rationale for developing the new software tool clearly explained?

Yes

Is the description of the software tool technically sound?

Yes
Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Partly

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?
Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Bioinformatics, computational biology, immunogenomics, scRNA-seq.

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 07 Jul 2020

Kent Riemondy, University of Colorado School of Medicine, Aurora,, USA

We thank the reviewer for their detailed suggestions which we believe have substantially improved the clarity of the manuscript. Our responses are indicated in italics below.

The authors describe an R package for annotating cell clusters in scRNA-seq datasets. Specifically, the package implements code for computing correlations between the columns of two data matrices. They show that high correlations between unknown cell clusters in the first data matrix and annotated cell types in the second matrix can be used to label the unknown cell clusters. They try varying parameters and show the effects on the results, and they also benchmark the time and accuracy compared to other packages designed to annotate scRNA-seq data.

Details of the code, methods, and analyses are partly provided. Some details seem to be missing (e.g. the functionality for gene lists).

The conclusions about the tool and its performance are partly supported by the findings presented in the article. Some terms such as "medF1-score" and "accuracy" are left undefined, and some results omit some methods (Figure 4A has different methods than B or C). Readers may have difficulty understanding the specific questions that were asked and what results are shown.

Main comments:
  ○ The clarity of the manuscript can be increased by adding more verbose details about all analyses. Please consider expanding details about each question, the approach,
In an effort to more clearly present clustifyr we have added additional details about each dataset, the questions posed by the analysis, and the conclusions from each analysis.

- Please consider adding a table describing the reference datasets used in this article, just like the one shown on one of your GitHub repositories. This should help to summarize which datasets were used for the analyses in this article.

We have added a table to the main text (Table 1) and a supplemental table that provide additional details about each dataset, and provide a reference of each dataset used in each figure panel.

Comments about specific parts of the manuscript are below. Excerpts from the article are shown in "quoted italics" after a bullet point, and my comments are shown directly below the bullet point.

- "A key challenge in scRNA-seq data analysis is the identification of cell types from single-cell transcriptomes. Manual inspection of the expression patterns from a small number of marker genes is still standard practice, which is cumbersome and frequently inaccurate."

Do we know the accuracy by manual inspection? Is there a reference for this? In the absence of evidence, you might consider weakening the statement to say "may be inaccurate" rather than "is cumbersome and frequently inaccurate". You might consider that many scRNA-seq experiments are done for the purpose of discovering new cell types that have not been well-described in previous published datasets. In this setting, manual inspection is necessary and automated analyses could be inaccurate or misleading.

To our knowledge there has not been a direct study of the accuracy of manual inspection compared to automated methods. We thank the reviewer for noting this point and have weakened this statement accordingly. We also have noted that automated methods can supplement manual inspection of markers to provide additional justification of the discovery of novel cell types.

- "Currently, multiple cell type assignment packages exist but they are specifically tailored towards input types or workflows."

Please consider naming and describing each method that will be compared to clustifyr in this manuscript, so the reader can assess how the methodology of clustifyr compares to other methods. Which methods are "specifically tailored towards input types or workflows"? Could you give an example to help the reader understand this claim?

We have added descriptions of the methodologies used by tools that we compared clustifyr against (see Introduction). We also have noted which tools are tailored towards input types: reference single cell data (Seurat, ACTINN, scPred) or workflows: using Seurat objects (Seurat), using singleCellExperiment (singleR, scPred), or using the command-line (ACTINN).

Suggested improvements for Figure 1:

- In Figure 1, you might consider showing the dimensions of the inputs and outputs. This might help the reader to understand how they relate to each other.

We have added the dimensions to provide clarity.

- Should the query and reference data be counts? CPM? Or Log2(CPM + 1)? You might
consider elaborating on this. 

Clustifyr supports both raw counts or log normalized values. The decision of which to use is left to the user and we recommend using similar normalization as used for the reference matrix, if possible. We have added text (under Variable gene selection and normalization) to provide guidance to the user.

Suggested improvements for Figure 2:

- Please consider rotating Figure 2A, D, and E 90 degrees clockwise to improve legibility.
  - We have amended the figures as suggested.
- Please consider limiting the axes ranges to the data instead of using the range [0, 1].
  - We respectfully decline to implement this suggestion, as we believe restricting the plot to only the range of the data can over-emphasize minor differences in distributions.
- Please consider increasing all font sizes in all panels in all figures, including titles, legends, axis text, etc. Some readers might need larger sizes to see clearly.
  - We have increased the font sizes accordingly.
- Please consider changing the title to “All genes (n = 10,000)” and “M3Drop variable genes (n = 1,000)” in Figure 2C, so we have some sense of the number of genes used to generate each heatmap.
  - We have changed these titles as suggested.
- Please consider showing a graphical representation of the experiment setup for this figure. What is the reference? What is the query? What are their dimensions? What is the main question in this analysis?
  - We have added additional details about the query and reference datasets in the main text, legends, and in the titles of figure panels as an alternative to graphical representations. The questions addressed by each analysis are more clearly stated when introducing each figure panel. We believe that these edits now provide sufficient clarity for the reader to understand the content of each figure.
- One way to enhance clarity is to add descriptive titles to every figure in every panel (e.g. "Testing different correlation statistics", etc.).
  - We thank the reviewer for this suggestion and we have added titles to figure panels that we believe were unclearly presented.
- In Figure 2C, it seems that the y-axis and x-axis have been swapped by mistake. I see that the y-axis is labeled "ground truth cell type" but it includes "unclassified". I would expect the category "unclassified" to appear in the "called cell type" axis, but not in the "ground truth cell type" axis. Are the axes swapped or are they correct? Could you please clarify?
  - The unclassified cell type was annotated as unclassified in the original study, whereas the query dataset contained a cell type (schwann cells) that appears to be similar to the reference data “unclassified”. We have added additional text to the legend to clarify.
- In Figure 2E, what does the color indicate? Is it the power argument "n^x" or
something else?

_The color did not indicate any particular class and therefore was removed._

- The reader may be wondering:
  - How many query cells did you use?
  - How many clusters were in the query dataset? How many cells per cluster?
  - How many reference datasets were used?
  - How many clusters were in the reference dataset?
  - Were the query and reference datasets acquired from the same tissue sample or were they completely independent and unrelated?

_We have added a supplemental table (supplemental table 1) with additional details for each dataset in the manuscript. The tissues samples used for the query and reference datasets were derived from unrelated individuals or mice based on our reading of the original publications for each dataset. An exception was the PBMC-bench dataset, in which multiple single cell technologies were tested using the same aliquot of PBMCs. We have added text to the results section to clarify (under Correlation method)._

In the section "Subclustering", please consider adding more details to help the reader avoid misunderstandings. What exactly is the "sub-clustering power argument (x)"? Please consider giving a concrete example to help the reader understand this section. Please consider creating a new figure that helps the reader to understand the "subcluster()" functionality.

_We have added an additional figure (2E) to demonstrate the utility of the subcluster/overcluster_test functionality._

What is the PBMCbench data? Is this the same data as mentioned in the section "Correlation minimum cutoff"?

_Yes this is the same dataset. We have added additional details to the text to introduce this dataset, as well as additional details provided in table 1._

In the section "Cells per cluster", you might consider introducing the dataset, then introducing the question that is being addressed, and finally reporting the results. What is the number (15, 8, 4)? Is the "Mouse Cell Atlas" the same as the "Tabula Muris"? Were these mouse datasets used in the previous sections? The reader might benefit from an introduction of these datasets.

_We have provided additional text to the manuscript to introduce and describe these datasets to improve clarity about the analyses conducted. The x axis refers to the number of cells per cluster._

Suggested improvements for Figure 3:

- Please consider adding labels "A", "B", "C", "D" to mark each of the four panels, so they can be referenced clearly.

_We have added these labels and referenced them in the updated figure legend._

- Please consider using the same name consistently in the text and the figure titles. For example, the figure says "Bulk RNA-seq reference data" but the text says "ImmGen
database". The reader might better understand the results if the same label were used in both places instead of using two different labels for the same thing.

We have added subtitles to each panel to more clearly reference the datasets in the text.

- Please consider including the identifiers for readers who wish to find these datasets and download them. For example, if the datasets are available on NCBI GEO, please consider including the accession numbers directly in the legend text, or in a table. Check to see if any other database provides an accession number. If an accession number is not available, please consider providing the DOI for a publication or a URL for a website that provides the data. By the way, if any data you are using is not deposited to a permanent repository, please consider uploading this data to a permanent repository (e.g. Figshare).

The publicly available datasets are referenced in the Data Availability section, with additional details now provided in Table 1. GEO accession numbers, DOIs, or URLs are provided, depending on the datasource. Additionally to further ease access to these resources we have organized these datasets into an ExperimentHub (clustifyrdatahub) that is in the process of being submitted to bioconductor.

In the section describing Figure 4A, please consider these suggested changes:

- Please explain what is "clustifyr", "clustifyr_lists", and "clustifyr_m3drop".
- How was feature selection performed for each analysis in Figure 4A?
- What is the strategy used by scmap?
- What is the strategy used by ‘Seurat’?
- What is the strategy used by ‘SingleR’?
- How is clustifyr similar or different?

We have added an additional paragraph to explain the differing clustifyr methods shown in Figure 4A. Feature selection was performed by the Tabula muris investigators using the variable genes selected by Seurat by examining a plot of the gene expression mean vs. variance (mean.var.plot). Seurat and clustifyr use these variable genes, whereas SingleR and scmap define variable genes using differential expression testing or M3Drop respectively. We have added text to explain the feature selection methods used by each benchmarked method.

This section says "Correlation-based clustifyr classification performed better than hypergeometric-based gene list enrichment as implemented in clustify_lists." Please consider explaining the "clustify_lists" algorithm in detail and also consider sharing the quantification of the performance of each approach so the reader can interpret the claim "performed better". Also consider elaborating on "performed better".

We have elaborated on the clustifyr_lists approach for classifying cell types based on gene set enrichment in the text. Additionally we have included a comparison of two approaches that performed best in our benchmarking, using hypergeometric tests, or using the jaccard index and selecting the cell type with the highest index value (Figure 4A).

What is "scRNAseq_Benchmark subsampling"? Could you elaborate on what this is and why it was used?
We have added additional text to the result section to introduce this dataset. This dataset contains random subsets of the tabula muris dataset to enable investigation of performance and accuracy with varying cell numbers.

Suggested improvements for Figure 4:
- Please consider including an overview schematic to help the reader understand which datasets were used for each result.
- We have added descriptive titles and additional text to the results section to describe the datasets and goals of each benchmarking test.
- Please define "accuracy". What is the algorithm for computing this number?
- Accuracy is defined as the ratio between the number of correctly classified clusters and the overall number of clusters for every dataset pair.
- Please define "medF1-score". What is the algorithm for computing this number?
- medF1-score was a shortened term for median F1-score. We have removed all references to medF1-score and replaced with median F1-score. F1-score, the harmonic mean of the precision and recall, is calculated for each cell type. A median F1-score is reported for every dataset pair.
- For the lower half of panel B, please consider using a format similar to the one in Figure 2B from Kiselev et al. (2018). For example, please use a log10 axis for time, so readers can see the difference between methods.
- We have modified the figure accordingly.
- Why is "medF1-score" used for Figure 4C and "accuracy" for Figure 4B?
- A F1-score cannot be calculated when the query and reference datasets contain different cell types. Therefore when comparing datasets with varying cell composition we instead utilized an accuracy metric (as defined above). We have added text to the manuscript that defines accuracy and median F1-score (Benchmarking methods), identifies which datasets were compared with each metric, and provides an explanation of why certain datasets were characterized with accuracy or F1-score.

Why does Figure 4A have 6 methods, Figure 4B have 5 methods, and Figure 4C have 3 methods? Is it possible to include all 6 methods for all panels? Could you please comment on the reasons for excluding or including methods in each analysis?

We agree that it is confusing for differing tools to be shown in different panels. We have therefore benchmarked these methods in a more consistent fashion to enable comparison of each method across different benchmarking tests. One exception however is scPred, which we were unable to successfully run on the Allen Brain Institute atlas data (Figure 4C), which we have noted in the figure legend.
- "As we and others observe, novel algorithms may not be necessary for cell type classification, at least within the current limitations of sequencing technology and our broadstroke understanding of cell “types”. Rather, the generation of community curated reference databases is likely to be critical for reproducible annotation of cell types in scRNA-seq datasets."

I agree that a community curated reference database would be a valuable contribution to the field. You might consider creating a table or other type of descriptive listing that helps the reader to understand all of the references that were used in this article. Consider
including tissue source, healthy or disease status, number of cells and genes, technology used for the assay, DOI, data URL, NCBI GEO accession, or any other details that the reader might find helpful.

**In addition to the dataset details provided in the Data Availability section and the details provided in Table 1, we have also included a supplemental table that references the datasets used in each figure, and an ExperimentHub package allowing direct access to these resources in R.**

Thank you for providing a GitHub repository with data files! Please also consider sharing the same data in compressed plain text format (e.g. "file.tsv.gz"). In addition to GitHub, please consider using a specialty service that is funded for the purpose of permanently archiving research data such as NIH Figshare ([https://nih.figshare.com](https://nih.figshare.com)). There are other options (Zenodo, Open Science Framework OSF, etc.).

The datasets used in this study were all published by other research groups and are hosted in various data repositories including GEO and figshare. As mentioned above we have provided additional methods to access these published and publicly available resources.

○ "As an alternative, clustifyr also supports per-cell annotation, however the runtime is greatly increased and the accuracy of the cell type classifications are decreased due to the sparsity of scRNA-seq datasets, and requires a consensus aggregation step across multiple cells to obtain reliable cell type annotations."

You might consider offering another alternative option. One extreme is to use the cluster averages, while the other extreme is to use single cells. Perhaps there might be a middle ground where clustifyr could automatically use k-means or some other algorithm to form clusters within the user-defined clusters. This would give the user even more flexibility.

After reviewing the code, I can see that there is an "overcluster()" function that seems to do exactly what I suggested. Please consider describing this in the article and showing an example of how it works. In retrospect, I can see that the section titled "Subclustering" was supposed to describe this topic — I misunderstood this section on the first read.

**We have added an additional figure panel (Figure 2E) to illustrate this functionality.**

You may want to double-check all of the links in all of your HTML pages. I see three URLs:

○ [https://github.com/rnabioco/clustifyrdatahub/](https://github.com/rnabioco/clustifyrdatahub/)
○ [https://github.com/rnabioco/clustifyr](https://github.com/rnabioco/clustifyr)
○ [https://github.com/rnabioco/clustifyrdata](https://github.com/rnabioco/clustifyrdata)

I can see that the "clustifyrdatahub" repo has code for creating ".rda" files from the reference datasets.

I also see similar scripts at [https://github.com/rnabioco/clustifyrdata/tree/master/data-raw](https://github.com/rnabioco/clustifyrdata/tree/master/data-raw)

Readers might be confused when they see two different repos with similar scripts. You might consider deleting the "clustifyrdatahub" repo if it is not necessary.

**We apologize to the reviewer for the confusion of multiple data repositories. We have organized...**
clustifyrdata into an ExperimentalHub Bioconductor package at the request of reviewer #1, resulting in overlapping content in the clustifyrdatahub repository. We have mentioned these differences in the documentation of these repositories and added text to the manuscript to point readers to the experimentHub package, which is currently being submitted to bioconductor.

I'm happy to see that the data is organized and annotated in the GitHub repo. Specifically, in the GitHub "clustifyrdata" repo, in the "README.md" file, the table shows the name of the reference, the number of cell types, the number of genes, the organism, and a link to the publication. Please consider adding some version of this table to the article, so the reader can understand the scope of this article.

We have added a table (Table 1) to the main manuscript that contains additional details about each dataset.

After reviewing the code, I was able to resolve some of my misunderstandings caused by lack of clarity in the terse descriptions in this article. To reduce the chance of misunderstanding by other readers, you might consider clarifying or adding details to the descriptions of functions and results. For example, the article does not mention that GSEA is used to work with gene lists.

We have added additional details about the gene list methods (including GSEA) to the article. clustify() and clustify_lists() are the most important functions implemented in clustifyr, which we believe are now sufficiently described in the revised manuscript. Additional package and function level documentation is provided at [https://rnabioco.github.io/clustifyr/](https://rnabioco.github.io/clustifyr/), which we've now provided as a link in the software availability section.

**Competing Interests:** None to declare.
not novel (simple correlation metrics), the software serves to streamline one of the most common procedures in single-cell RNA-sequencing analysis. As detailed below I have some questions regarding the evaluation of the method compared to existing approaches, and a suggestion to more widely distribute the prebuilt references curated as part of the study.

Major comments:
1. The 'unseen population rejection test' is an informative measure. However, it is not clear without going back to the scRNAseq_Benchmark (Abdelaal et al., 2019) how clustifyr's performance compares to other tools. It would be useful to give some quantitative or visualization that conveys this comparison.

2. The approach is aimed at being "normalization-agnostic" as stated in 'Benchmarking' section. However, it's not clear whether this refers to clustifyr in general, or just using the rank correlation setting. If in general, this property should be demonstrated.

3. The benchmarking results provided are very helpful, but it's not clear why only a (differing) subset of the methods was applied to each evaluation (i.e. panels of Figure 4 in particular).

Minor comments:
1. From the description of the method, it seems that if the query dataset is 'over-clustered', meaning a cell-type is incorrectly split into two clusters, clustifyr can return the same cell type assignment for both clusters (provided the correct reference had the highest correlation, and that correlation was above the threshold). Is this correct? If not, please clarify.

2. The prebuilt references in the clustifyrdata github repository has potential utility to researchers who don't already have a reference dataset. It might be a good fit to build these reference datasets as a Bioconductor ExperimentHub package.

References

Is the rationale for developing the new software tool clearly explained?
Yes

Is the description of the software tool technically sound?
Yes

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?
Partly
Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Partly

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

**Reviewer Expertise:** Statistical genomics, bioinformatics

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.

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**Author Response 07 Jul 2020**

Kent Riemondy, University of Colorado School of Medicine, Aurora, USA

We thank the reviewer for their helpful criticisms. Our responses are indicated in italics below.

**Major comments:**
- The 'unseen population rejection test' is an informative measure. However, it is not clear without going back to the scRNAseq_Benchmark (Abdelaal et al., 2019) how clustifyr's performance compares to other tools. It would be useful to give some quantitative or visualization that conveys this comparison.

  
  We agree and have provided an additional figure panel (4E) that provides a visual comparison of clustifyr's performance compared to tools assessed by the scRNAseq_Benchmark.

- The approach is aimed at being "normalization-agnostic" as stated in 'Benchmarking' section. However, it's not clear whether this refers to clustifyr in general, or just using the rank correlation setting. If in general, this property should be demonstrated.

  We are referring the property of rank correlation rather than a specific feature of clustifyr. We have amended the text (subsection: Variable gene selection and normalization) to make this point more clear and provide recommendations that users try to implement the same normalization scheme for reference and query data if possible.

- The benchmarking results provided are very helpful, but it's not clear why only a (differing) subset of the methods was applied to each evaluation (i.e. panels of Figure 4 in particular).

  We agree that the benchmarking would be more clearly presented by providing more complete assessment of methods across each evaluation. We have updated figures 4A,B,C, and D to consistently present clustifyr's performance and accuracy compared to other methods. Of note, we were unable to benchmark scPred when examining the Allen Brain Atlas data (Figure 4C), due to an error that we were unable to troubleshoot.

**Minor comments:**
- From the description of the method, it seems that if the query dataset is 'over-
clustered, meaning a cell-type is incorrectly split into two clusters, clustifyr can return the same cell type assignment for both clusters (provided the correct reference had the highest correlation, and that correlation was above the threshold). Is this correct? If not, please clarify.

The reviewer's comment is correct, clustifyr will assign the cell type with the highest correlation, that meets a minimum cut-off value. For over-clustered query cell types, clustifyr will therefore return the same cell-type label, despite the overclustering. Clustifyr also provides a function (overcluster_test()) to intentionally overcluster the query dataset to potentially identify subpopulations that were grouped into another cell type due to inappropriate query dataset clustering. We have included an additional figure panel (2E) to illustrate this functionality.

- The prebuilt references in the clustifyrdata github repository has potential utility to researchers who don't already have a reference dataset. It might be a good fit to build these reference datasets as a Bioconductor ExperimentHub package.

We thank the reviewer for this suggestion and have built an ExperimentHub package that includes the prebuilt references in the clustifyrdata repository. The package (clustifyrdatahub) has been submitted to Bioconductor.

**Competing Interests:** None to disclose.