scEFSC: Accurate single-cell RNA-seq data analysis via ensemble consensus clustering based on multiple feature selections

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1. Introduction

Advances in next-generation sequencing technologies brought single-cell RNA sequencing (scRNA-seq), which is rapidly generating large amounts of the single-cell RNA sequencing data. The main improvement achieved by scRNA-seq is that it overcomes the drawback of traditional bulk RNA sequencing in averaging gene expression across all cells in a sample [1]. On the contrary, scRNA-seq measures the transcriptome of individual cells, which makes high throughput investigations of tissue samples contributing to reveal heterogeneity of the cells and the molecular basis of phenotypic variation between them. This technology can provide new opportunities to characterize and understand the investigation of complex diseases and the dynamics of cell developmental cycles at the single cell resolution [2]. Therefore, the accurate identification of cell types and their underlying profiles has become an important step in single-cell RNA-seq analysis [3].

Clustering has been proven to be a critical computational method in grouping cells according to transcriptomic characteristics and thereby annotating cell types [4]. Recently, a plethora of clustering algorithms have been used to address the complexities of the scRNA-seq data [5]; for instance, Lin et al. [6] proposed CIDER, a clustering algorithm that reduces data dropouts through an implicit interpolation method using principal coordinate analysis (PCoA) to reduce the dimensionality. Qiu et al. [7] presented the Monocle model for clustering using t-SNE and density peak clustering. Yau et al. [8] proposed the pcaReduce algorithm, which integrates principal component analysis (PCA) and hierar-

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chical clustering to generate a cell state hierarchy in which each clustering branch is associated with a variant principal component. Levine et al. [9] developed the Phenograph algorithm for the analysis of high-dimensional single-cell data, by creating a graph representing the phenotypic similarity between cells. Satija et al. [10] proposed the Seurat algorithm for clustering based on PCA, Shared Nearest Neighbour (SNN) graphs and Louvain's algorithm. Guo et al. [11] proposed the SINCERA algorithm to optimize cell clusters by hierarchical clustering for detecting differentially expressed genes. Grün et al. [12] developed RaceID for identifying rare cell types in complex single cell populations. It is difficult to imagine, however, that each clustering method can be equally effective across all scRNA-seq datasets. Indeed, each clustering algorithm has its own strengths and weaknesses and performances exhibit specific characteristics on particular scRNA-seq datasets [13]. Therefore, this is a challenge for users to decide which clustering algorithm is the most appropriate choice for the scRNA-seq data in hand.

Clustering ensembles have emerged as an effective method for combining solutions from multiple individual clustering algorithms into a consensus result. A number of clustering ensemble models have been developed for the scRNA-seq data; for example, Kiselev et al. [14] proposed the SC3 algorithm to combine the base clustering results obtained using spectral transformation and k-means algorithms into a consensus matrix, and then employing a hierarchical clustering to achieve the final clustering result. Wan et al. [15] presented the SHARP algorithm to cluster scRNA-seq data, using meta-clustering algorithms to obtain the final clustering result. Yang et al. [16] proposed the SAFE-clustering algorithm to first perform an independent clustering using four methods, SC3, CIDR, Seurat, and t-SNE + k-means, and then integrating solutions using three hypergraph-based partitioning algorithms. Geddes et al. [17] developed an autoencoder-based cluster integration framework to obtain random subspace projections from the data and the reducing dimensionality using an autoencoder. Zhu et al. [18] developed the Sc-Gpe algorithm by combining five clustering methods based on single-cell graph partitioning and calculating the probability of cell pairs being classified into the same cluster. Huh et al. [19] presented the SAME-clustering algorithm to generate clustering solutions from multiple methods and selected a subset of maximum diversity to produce improved integrated solutions. However, we find that most clustering algorithms suppose that all features are equally significant in the clustering while in reality, diverse features have different effects on clustering [20]. This is one of the reasons why most clustering algorithms often perform poorly when faced with high-dimensional scRNA-seq data [21]. Therefore, efficient feature selection methods need to be developed for optimizing scRNA-seq data analysis.

To address these challenges, we first carry out a performance evaluation of multiple feature selection methods and nine scRNA-seq clustering algorithms on fourteen real single-cell RNA-seq datasets. Then, we present an accurate single-cell data analysis via Ensemble Feature Selection based Clustering, called scEFSC. The first module proposes removing genes that do not add significantly to the analysis of the scRNA-seq data using multiple unsupervised feature selections. Then, the second module implements different scRNA-seq clustering algorithms on the data generated in the first module to cluster the data, and then combines all the clustering results using a weighting-based meta-clustering method to obtain the final result. We applied scEFSC to the fourteen tested real scRNA-seq datasets compared clustering performance using several evaluation metrics to other scRNA-seq clustering algorithms. Results showed that our proposed scEFSC outperformed the other clustering algorithms. In addition, we realized differential gene expression analysis, gene ontology enrichment and KEGG analysis, demonstrating the biological interpretability of scEFSC.

2. Methods

2.1. Overview of the scEFSC algorithm

In our study, we developed an accurate single-cell data analysis via Ensemble Feature Selection based Clustering, called scEFSC. The input of the proposed scEFSC was the scRNA-seq data, which is composed of an $n \times d$ gene expression matrix $X = \{x_1, x_2, \ldots, x_n\}; x_i = \{x_{i1}, x_{i2}, \ldots, x_{id}\}$, where $n$ is the number of cells and $d$ is the number of genes. To begin, data pre-processing was performed using a log2 transformation to normalize the data and then genes detected in the normalized data in less than 2% of the cells were removed to filter out the low-level expressed genes from the scRNA-seq data. The overall framework of our proposed scEFSC is summarized in Fig. 1. As depicted in this figure, scEFSC consists of three important phases.

In phase A, we first employed a non-negative kernel autoencoder [22] to pre-select 5000 genes to remove insignificant genes. After that, we proposed multiple unsupervised feature selections including Low Variance [23], Laplacian Score [24], SPEC [25], and MCFS [26] to remove genes that do not contribute significantly to the analysis of the scRNA-seq data. We then fed the derived feature subsets into the clustering algorithms. Among them, we can observe that Low Variance is based on statistics; Laplacian Score and SPEC are based on similarity and MCFS is based on sparse learning. The last three methods are extensions of spectral model previously used for scRNA-seq data analysis. Unlike feature extraction methods, these feature selection methods do not change the original representation of the data and are considered to provide better readability and interpretability [27].

In phase B, we applied several different scRNA-seq clustering algorithms to cluster the feature subsets obtained by the multiple feature selection models. Various scRNA-seq clustering methods exist to run in our model. These methods are based on different underlying mathematical formulations as described above, including SC3 [14], CIDR [6], monocyte [7], pcReduce [8], Rphenograph [9], Seurat [10], SHARP [15], SINCERA [11], and RaceID [12]. For each feature subset derived from the different feature selection models, we applied the stated clustering methods to generate cluster labels to finally yield a set of individual cluster labels. In addition, to enhance the diversity of the individual cluster labels in the set, the pairwise Adjusted Rand Index (ARI) was employed to measure the similarity between any two individual clustering labels and then remove the method having similarity with the lowest variance [19]. In phase C, a weighted-ensemble clustering method called wMetaC was used to obtain the final clustering result of the individual cluster labels [15].

2.2. Non-negative kernel autoencoder

Since non-negative kernel autoencoders have been proven to be successful for screening single-cell sequencing data with highly expressed genes [22], in our study we first use non-negative nuclear autoencoders to preselect genes. Indeed, the normalized data is fed to a single-layer autoencoder to remove the insignificant genes, which consists of two important parts: the encoder and the decoder. The autoencoder can be formulated as follows.

\[
f_E(x), f_D(x) = xW_E + b_E \quad (1)
\]

\[
f_D(e), f_D(e) = eW_D + b_D \quad (2)
\]

where $f_E$ and $f_D$ represent the conversion function of the encoder and decoder layers, $x$ is the input of the encoder, $e$ is a restriction of $x$, $e$ is the reduced dimensional data of $x$, $W$ is the weight matrix, and $b$ is the bias vector. The encoder aims to represent the data in a
low-dimensional space, while the decoder tries to rebuild the original input from the reduced data. After that, the encoder weights are imposed as non-negative so that each latent variable is an additive representation of the original features. Then, the non-negative factors of less important features are reduced to zero. Based on the calculated weights, the method retains only those genes with large weight variances that can be considered as important features to characterize the original data. Following the reference [22], we also pre-selected 5000 genes for subsequent algorithm evaluation and analysis.

2.3. Unsupervised Feature Selection Algorithms

After data pre-processing, we employed four unsupervised feature selection algorithms, Low Variance [23], Laplacian Score [24], SPEC [25] and MCFS [26] to further select genes with relevant information, and then obtained four different feature subsets. The detailed section is summarized in the Supplementary Section S1.

Low Variance is a simple and effective unsupervised algorithm for choosing features, consisting of a typical filtering method that
evaluates features by their variance which can be formulated as follows:

$$\sigma^2 = \frac{\sum_{i=1}^{n}(x_i - \mu)^2}{n},$$  \hspace{1cm} (3)

where $x_i$ is the value of the $i$-th sample, $\mu$ is the mean of samples, and $n$ is the number of samples.

Laplacian Score is a filter method with Laplacian Eigenmaps and Locality Preserving Projection. The Laplacian Score first builds a graph and connects the adjacent points. The incidence matrix $S$ can be defined as follows:

$$S_{ij} = \begin{cases} e^{-\|x_i - x_j\|^2}, & \text{if nodes } i \text{ and } j \text{ are connected,} \\ 0, & \text{otherwise}, \end{cases}$$  \hspace{1cm} (4)

where $t$ is a constant. Then, it establishes a diagonal matrix $D$ on the basis of $S$, where the diagonal element $D_{ii}$ of $D$ is the sum of the elements in the $i$-th row of $S$, generates a Laplacian matrix $L = D - S$ according to $S$ and $D$. The Laplacian Score of the $i$-th feature is calculated as follows:

$$L_i = f_i^T L f_i, \hspace{1cm} (5)$$

$$f_i = f_i - \frac{f_i^T D}{1^T D} 1, \hspace{1cm} (6)

$$S_{ij} = e^{-\|x_i - x_j\|^2}. \hspace{1cm} (7)

SPEC, induces the undirected graph $G$ from the data and obtains its adjacency matrix $W = S$. The degree matrix $D$ of the graph $G$ is established by $W$. $D$ is a diagonal matrix and its diagonal element $D_{ii}$ is the sum of the elements in the $i$-th row of $W$. The Laplace matrix $L$ is defined as $L = D - W$. The normalized Laplacian matrix $\bar{L}$ is formulated as $\bar{L} = D^{-\frac{1}{2}} L D^{-\frac{1}{2}}$. The relevance of the features is evaluated by using the spectrum of the graph in SPEC. The scoring function is calculated as follows:

$$\phi_1(F_i) = f_i^T \bar{L} f_i, \hspace{1cm} (8)$$

$$\phi_2(F_i) = \frac{f_i^T \bar{L} f_i}{1 - f_i^T \xi_0}, \hspace{1cm} (9)$$

$$f_i = \frac{D^{1/2} f_i}{||D^{1/2} f_i||}, \hspace{1cm} (10)

$$\phi_1(F_i) = f_i^T \bar{L} f_i, \hspace{1cm} (8)$$

where $F_i$ is the $i$-th feature of data, $f_i$ is the vector formed by the $i$-th feature of data, $\bar{L}$ is the Fourier transform of $L$, and $\xi_0 = D^{1/2} \epsilon$.

MCFS is based on manifold learning and L1-regularized models for feature selection so that the multi-cluster structure of the data can be well preserved. In MCFS, the representation of the data in the embedding space is obtained by solving the generalized feature problem $L y = \lambda D y$. Let $Y = [y_1, \ldots, y_k]$ be the eigenvector corresponding to the smallest eigenvalue, and $K$ be the dimension of the embedding space. The coefficients of linear regression are obtained by addressing the following problems:

$$\min_{a_k} |y_k - X^T a_k|^2 + \beta |a_k|_1, \hspace{1cm} (11)$$

where $a_k$ is the regression coefficient vector, $|a_k|$ is the L1-norm of $a_k$. The importance of each feature is determined by the MCFS score. The MCFS score is obtained as follows:

$$MCFS(j) = \max_{k} |a_{kj}|, \hspace{1cm} (12)$$

where $a_{kj}$ is the $j$-th element of the vector $a_k$.

2.4. Implementation of multiple scRNA-seq clustering methods

After implementing the unsupervised feature selection methods, nine scRNA-seq clustering algorithms, including SC3[14], CIDR[6], moncolle[7], pcaReduce[8], Rphenograph[9], Seurat[10], SHARP[15], SINCERA[11] and RaceID[12], were employed for clustering under those four different feature subsets generated by the above four unsupervised feature selection algorithms. We describe this briefly below, with detailed sections in the Supplementary Section S2.

Table 1 summarizes these single-cell clustering algorithms. For SC3, we used the version 1.18.0 from Bioconductor and employed the number of cell types as the number of clusters $k$ in our proposed model. For CIDR, we used the version 0.1.5 from GitHub (github.com/shibiaowan/CIDR). The Monocle R package was the version 2.18.0 from GitHub (github.com/cole-trapnell-lab/monolol-releas e). For pcaReduce, we used the version 1.0.0 from GitHub (github.com/JustinaZ/pcaReduce) and performed merging based on largest probability. The Rphenograph R package was the version 0.99.1 from GitHub (github.com/JinmingChenLab/Rphenograph). For Seurat, we used the version 4.0.0 from CRAN and the dimension of reduction to use as input was 10. For SHARP, we used the version 1.1.0 from GitHub (github.com/shibiaowan/SHARP). The SINCERA R package was the version 0.99.0 from GitHub (github.com/JinmingChenLab/Rphenograph). For Seurat, we used the version 4.0.0 from CRAN and the maximum number of clusters for the derivation of the cluster number by the saturation of mean within-cluster-dispersion was 20.

2.5. scEFSC: Ensemble Consensus Clustering Based on Multiple Feature Selections

After data preprocessing and non-negative kernel autoencoder, multiple unsupervised feature selection was proposed to select different feature subsets to remove genes that do not contribute significantly to the scRNA-seq data. On this basis, the data generated by the four unsupervised feature selections were clustered by exe-
cutting nine single-cell RNA-seq clustering algorithms separately to finally obtain the underlying set of clustering results.

2.5.1. Ensemble consensus clustering

After obtaining the basic clustering results and removing labels with least diversity, we proposed employing weighted-based meta-clustering (wMetaC) [15] to combine the clustering results from the multiple individual clustering methods. In contrast to traditional cluster-based similarity partitioning algorithms that give equal importance to each instance and each cluster, wMetaC assigns different weights to different instances or pairs of instances and different clusters to enhance the clustering capabilities. With wMetaC, the individual clustering results were transformed into a clustering similarity matrix $S$ whose elements $s_{ij}$ represented the similarity between the $i$th and $j$th cells. Since the weight of each cell pair is determined by the degree of agreement of the co-localization clustering results of the two cells, the similarity matrix $S$ is transformed into a weight matrix $W$ as follows:

$$w_{ij} = s_{ij}(1 - s_{ij}),$$

where $w_{ij}$ is the element of the $i$th row and $j$th column in $W$. When $s_{ij} = 1$ or $s_{ij} = 0$, $w_{ij}$ reaches a minimum value of 0; when $s_{ij} = 0.5$, $w_{ij}$ reaches a maximum value of 0.25. Zero weights were assigned to cell pairs that were easiest to cluster, while the highest weights were assigned to the most difficult pairs to cluster. The weight associated with each cell was then calculated as the cumulative sum of all the intercellular weights associated with the corresponding cell.

Then, the weighted inter-cluster similarity was calculated by dividing the sum of the weights of their overlapping elements by their combined weights. Given two clusters $C_u$ and $C_v$, the inter-cluster similarity in wMetaC was calculated as:

$$S = \frac{\sum_{t \in C_u, j \in C_v} w_{tj} + \delta}{\sum_{t \in C_u, j \in C_v} w_{tj} + \delta},$$

where $w_{tj}$ is the co-localization weight of the $t$th cell and the $j$th cell obtained above, $N$ is the number of cells, and $\delta$ is a very small positive number, with $\delta = 0.01$ used to avoid a zero denominator. Since $\sum_{j=1}^{N} w_{tj}$ summarizes all possible co-localization weights between the $t$th cell and all cells, it is set to the overall co-localization weight of the $t$th cell. The numerator of the inter-cluster similarity formula represents the sum of the co-localization weights of the cells that appear in both the $C_u$ and $C_v$ groups, while the denominator represents the sum of the co-localization weights of the cells that appear in either the $C_u$ or $C_v$ group.

With this, the similarity matrix obtained was clustered using the hierarchical clustering method with the “ward.D”. The correspondence between the clusters was obtained after clustering. Finally, a voting scheme [15] was applied to the clustering results obtained in the previous step and the individual cells were assigned to the clusters to which they fit in the highest proportion, resulting in the final clustering result.

2.5.2. Removing labels with least diversity

Since the diversity of clustering results is beneficial to enhance the performance of the ensemble solution [28], we computed pairwise ARI between clustering labels to quantify the similarity between the results of individual clustering methods to evaluate the diversity. After that, a similarity matrix was constructed by calculating the pairwise similarity between all individual clustering labels, including the clustering labels with a self-similarity value of 1. Then the variance of the similarity vector was evaluated for each clustering labels. Since high pairwise similarity has a high ARI value and a self-similarity value of 1, the similarity vector with the most similar cluster label to the other cluster labels has the smallest variance. Because the clustering labels with the lowest variance contribute least in terms of diversity, we removed clustering labels obtained from the few clustering methods with the lowest variance to obtain the set of clustering labels.

2.6. Evaluation metrics

Comparing the agreement of cell clustering results with the previously published labels is a common method for evaluating the performance of clustering methods for scRNA-seq data analysis. In this study, we used two evaluation metrics: Normalised Mutual Information (NMI) and Adjusted Rand Index (ARI), as summarized in Supplementary Section S3.

3. Results and Discussion

3.1. Data source

We collected fourteen publicly available scRNA-seq datasets containing cell type annotations and gene expression values from various scRNA-seq platforms, which can be downloaded from the Gene Expression Omnibus and Broad Institute Single Cell Portal. All the datasets are from different species, including mouse and human, as well as from different organs, such as brain, lung and kidney. The detailed information on the datasets is summarized in Table 2. Of note, we removed cells labeled ambiguously as “abandoned” and then performed a logarithmic transformation (base 2) to rescale the data if its gene expression values were higher than 100. Indeed, for the 14 scRNA-seq datasets in Table 2, we can see that the number of cells varies from 90 to 13316, the number of genes from 19020 to 55186 and the number of clusters from 4 to 14. To further remove redundant and irrelevant genes and reduce the computational cost of the algorithm, we used a non-negative kernel autoencoder [22] to pre-select 5000 genes for downstream analysis.

3.2. scEFSC is the most accurate among computational methods on the scRNA-seq data

To investigate the advantages of clustering using scEFSC, we benchmarked our proposed scEFSC method alongside the nine scRNA-seq clustering methods on the fourteen published scRNA-seq datasets (Table 2). The clustering results measured by NMI and ARI are summarized in Fig. 2.

The results of the NMI evaluation, are summarized in Fig. 2a. We observe the following: it appears that of the ten single-cell clustering algorithms, scEFSC provided the best clustering results on eleven of the fourteen datasets, and ranked second, slightly below SC3 on the E-MTAB-5061 dataset, and third and fourth on the GSE36552 and GSE84133 datasets; CIDR had the worst performance. Compared to Rphenograph and Seurat that based on the GSE36552 and GSE84133 datasets; CIDR had the worst performance.

The results of the NMI evaluation, are summarized in Fig. 2a. We observe the following: it appears that of the ten single-cell clustering algorithms, scEFSC provided the best clustering results on eleven of the fourteen datasets, and ranked second, slightly below SC3 on the E-MTAB-5061 dataset, and third and fourth on the GSE36552 and GSE84133 datasets; CIDR had the worst performance. Compared to Rphenograph and Seurat that based on the GSE36552 and GSE84133 datasets; CIDR had the worst performance. In addition, compared to the consensus clustering algorithms, SC3 and SHARP, scEFSC was superior to both indicating that multiple clustering is more effective for base clusters in consensus clustering.

In terms of ARI evaluation, as shown in Fig. 2b, we observe broadly similar results to those using the NMI. Our proposed scEFSC performed better than the other nine clustering algorithms, while CIDR and SHARP simultaneously performed the worst. scEFSC clustered best on ten datasets, being second only to SC3 on the GSE71585 and E-MTAB-5061 datasets, second only to SIN-
CERA on the SCP345 dataset and fourth on the GSE84133 dataset. Although CIDER showed a higher performance than most of the other algorithms on the GSE83139 and GSE81861 datasets, it performed particularly poorly on GSE85241, E-MTAB-5061, phs000833.v3.p1, GSE41333 datasets. SHARP performed the worst on the GSE81861 and GSE71585 datasets, and did not perform well on the remaining datasets. SC3 achieved better performance than the others on most scRNA-seq datasets, while scEFSC outperformed SC3 on most datasets. scEFSC was better than community detection method Louvain algorithms, Rphenograph and Seurat. The higher ARI values of scEFSC clustering results over consensus clustering algorithms, SC3 and SHARP, suggest that multiple clustering is more effective in base clustering for consensus clustering.

In summary, our proposed scEFSC algorithm performed well and demonstrated superior clustering performance on fourteen scRNA-seq datasets, improving the clustering performance on the scRNA-seq data. The results also reflect that the combination of multiple clustering models can enhance the performance of consensus clustering.

### 3.3. scEFSC outperforms other ensemble clustering algorithms

We compared scEFSC, SAME[19], scCCESS[43] and RSEC[44] on the fourteen published scRNA-seq datasets as in the previous section. SAME, scCCESS and RSEC are three ensemble clustering algorithms for single-cell RNA-seq data. The clustering results measured by NMI and ARI are summarized in Fig. 3.

Fig. 3a summarizes the clustering results evaluated by NMI. Compared with SAME and RSEC, our proposed scEFSC had the best NMI values on all 14 datasets. Of the fourteen datasets, our proposed scEFSC had better NMI values on twelve datasets compared with scCCESS, they provided the similar results on the GSE41333 dataset, and on the GSE36552 datasets scCCESS had better NMI values than our proposed scEFSC. Our proposed scEFSC performed the best among the four algorithms, and RSEC performed the worst. On the GSE59739, GSE81608, and SCP345 datasets, our proposed scEFSC was superior to the other three algorithms by more than 0.1. Overall, the clustering results showed that our proposed scEFSC outperformed SAME, scCCESS and RSEC on most scRNA-seq datasets, and our proposed scEFSC exhibited excellent clustering performance in comparison with other ensemble clustering algorithms.

3.4. The performance of scEFSC outperforms the scDHA algorithm recently reported in Nature Communications

We compared scEFSC and scDHA[22] on the fourteen published scRNA-seq datasets as in the previous section. scDHA is a scRNA-seq clustering method, which uses a hierarchical autoencoder published in Nature Communications. The scDHA algorithm framework consists of two main components. The first component is a non-negative kernel autoencoder that aims to filter genes that contribute little to the data representation. The second part is a stacked Bayesian autoencoder for data dimensionality reduction. Fig. 4 depicts the clustering results of scEFSC and scDHA as measured by NMI and ARI.

The clustering results of scEFSC and scDHA assessed by NMI are summarized in Fig. 4a. Of the fourteen datasets, our proposed scEFSC had better NMI values on ten datasets compared with scDHA, both algorithms had the same result on the GSE71585 dataset, and on the GSE36552, E-MTAB-5061 and GSE65525 data-sets scDHA had better NMI values than our proposed scEFSC. Our proposed scEFSC significantly outperformed scDHA on the GSE59739 dataset, with an improvement in clustering performance of 0.11. Our proposed scEFSC was superior to scDHA by more than or equal to 0.03 on the GSE81252, GSE81608, GSE85241, GSE60361 and GSE84133 datasets. On the E-MTAB-5061 and GSE65525 datasets, our proposed scEFSC was worse than scDHA.

Fig. 4b summarizes the clustering results for scEFSC and scDHA assessed by ARI. For the fourteen datasets our proposed scEFSC had better ARI values than scDHA on ten datasets, scEFSC and scDHA gave the similar ARI value on the GSE71585 dataset, and on the GSE36552, E-MTAB-5061 and GSE65525 datasets scDHA had better ARI values than our proposed scEFSC. Our proposed scEFSC performed the best among the four algorithms, and RSEC performed the worst. On the GSE59739, GSE81608, GSE85241, GSE65525, GSE60361, phs000833.v3.p1, and SCP345 datasets, our proposed scEFSC was superior to the other three algorithms by more than 0.1.

Overall, the clustering results showed that our proposed scEFSC outperformed SAME, scCCESS and RSEC on most scRNA-seq datasets, and our proposed scEFSC exhibited excellent clustering performance in comparison with other ensemble clustering algorithms.

Table 2

| Source | Organism | cell | gene | class | platform | Ref |
|--------|----------|------|------|-------|----------|-----|
| GSE36552 | Human embryo | 90 | 20214 | 6 | Tang | Nature structural & molecular biology [29] |
| GSE83139 | Human pancreas | 457 | 19950 | 7 | SMARTer | Diabetes [30] |
| GSE81861 | Human tissues | 561 | 55186 | 9 | SMARTer | Nature genetics [31] |
| GSE59739 | Mouse brain | 622 | 23334 | 4 | STRT-Seq | Nature neuroscience [32] |
| GSE81252 | Human liver | 777 | 19020 | 7 | SMARTer | Nature [33] |
| GSE81860 | Human pancreas | 1500 | 39851 | 8 | SMARTer | Cell metabolism [34] |
| GSE71585 | Mouse brain | 1679 | 24150 | 18 | SMARTer | Nature neuroscience [35] |
| GSE85241 | Human pancreas | 2126 | 19140 | 10 | CEL-Seq2 | Cell systems [36] |
| E-MTAB-5061 | Human pancreas | 2209 | 25523 | 14 | Smart-Seq2 | Cell metabolism [37] |
| GSE65525 | Mouse embryo | 2717 | 24175 | 4 | inDrop | Cell [38] |
| GSE60361 | Mouse brain | 3005 | 19972 | 9 | STRT-Seq | Science [39] |
| phs000833.v3.p1 | Human brain | 3042 | 25123 | 16 | Fluidigm C1 | Science [40] |
| GSE84133 | Human pancreas | 8569 | 20125 | 14 | inDrop | Cell systems [41] |
| SCP345 | Human pancreas | 13316 | 21813 | 8 | 10X Genomics | The reference [42] |
Fig. 2. Clustering performance comparison of scEFSC, SC3, CIDR, monocle, pcaReduce, Rphenograph, Seurat, SHARP, SINCERA and RaceID on the fourteen published scRNA-seq datasets. (a) Clustering performance evaluated by NMI. (b) Clustering performance evaluated by ARI.
inferior to scDHA on only a few scRNA-seq datasets. Our proposed scEFSC demonstrated an excellent clustering performance therefore, compared to scDHA.

3.5. Integration of multiple feature selection methods can improve downstream functional analysis

We compared our proposed scEFSC with the scEFSC algorithms with one of the four feature selections on the fourteen scRNA-seq datasets shown in Table 2 using the clustering evaluation metrics NMI and ARI to evaluate the clustering results and summarize the results in Fig. 5.

With the NMI evaluation, the full scEFSC outperforms the other four single-feature selection algorithms on six scRNA-seq datasets including GSE59739, GSE81608, GSE85241, GSE65525, GSE60361, and phs000833.v3.p1. On the GSE36552 and GSE81252 datasets, full scEFSC performed just as well as some of the other four algorithms. On GSE85241, the performance of our proposed scEFSC was only lower than scEFSC having only Low Variance. The performance of full scEFSC on the GSE59739, GSE71585, E-MTAB-5061, GSE84133 and SCP345 datasets, however was not good. In summary, our proposed scEFSC had the best performance, scEFSC with only Low Variance was second best and scEFSC with only MCFS the worst.

Altogether, our proposed scEFSC algorithm exhibited superior clustering performance on the fourteen scRNA-seq datasets, indicating that using multiple unsupervised feature selection algorithms can strengthen the clustering ability of consensus clustering over a single unsupervised feature selection algorithm.

3.6. The clustering performance of scEFSC is affected by different parameters

With ARI evaluation, our proposed scEFSC similarly outperformed the other four algorithms on six datasets including GSE83139, GSE81861, GSE81608, GSE65525, GSE60361, and phs000833.v3.p1. On the GSE36552 and GSE81252 datasets, full scEFSC performed just as well as some of the other four algorithms. On GSE85241, the performance of our proposed scEFSC was only lower than scEFSC having only Low Variance. The performance of full scEFSC on the GSE59739, GSE71585, E-MTAB-5061, GSE84133 and SCP345 datasets, however was not good. In summary, our proposed scEFSC had the best performance, scEFSC with only Low Variance was second best and scEFSC with only MCFS the worst.

Altogether, our proposed scEFSC algorithm exhibited superior clustering performance on the fourteen scRNA-seq datasets, indicating that using multiple unsupervised feature selection algorithms can strengthen the clustering ability of consensus clustering over a single unsupervised feature selection algorithm.

3.6. The clustering performance of scEFSC is affected by different parameters

In terms of parameters, our proposed scEFSC default selects 5000 genes by a non-negative kernel autoencoder and 2000 genes by unsupervised feature selections, using nine clustering algorithms. To verify the superiority of the default parameters, we conducted two sets of experiments on the fourteen scRNA-seq datasets testing these parameters. The first experiment compared the effect
of the number of features selected on scEFSC. The second experiment compared the effect of the number of clustering algorithms in the multiple clustering on scEFSC.

For the first parametric experiment, we compared scEFSC with the proposed default parameters to scEFSC with 10,000 or 5,000 genes selected, and scEFSC with 5,000 or 3,000 genes selected, respectively and the clustering results assessed by NMI and ARI are summarized in Fig. 6a and 6b. NMI and ARI values were broadly similar. scEFSC with default parameters outperformed scEFSC with two feature selections of 10,000 and 5,000 genes on thirteen datasets and was the same on the GSE60361 dataset. Therefore, our proposed scEFSC using a non-negative encoder for 5,000 genes feature selection was effective. In addition, scEFSC with default parameters performed better on eight datasets, the same on three datasets and worse on three datasets, compared to the scEFSC with two feature selections of 5,000 and 3,000 genes. This shows that our proposed scEFSC was effective in selecting 2000 genes using unsupervised feature selection. In conclusion, our proposed scEFSC default setting of 5000 and 2000 genes for feature selection effectively enhanced the clustering performance.

In the second parametric experiment, on the fourteen datasets, we compared scEFSC with our proposed default parameters to scEFSC with eight clustering algorithms and scEFSC with six clustering algorithms. Fig. 6c and 6d summarize the clustering results,
showing the largely consistent results evaluated by NMI and ARI. In the previous comparison of our proposed scEFSC and nine single-cell clustering algorithms, SC3 was second to scEFSC in terms of average clustering results across the fourteen datasets and was better than the other nine single-cell clustering algorithms, while CIDR, SHARP and pcaReduce were the three worst algorithms. The scEFSC using eight clustering algorithms eliminated the SC3 algorithm and the scEFSC using six clustering algorithms excluded CIDR, SHARP and pcaReduce. Our proposed scEFSC clustered better on ten datasets, equally well on the

Fig. 5. Performance comparisons of the different feature selections. NMI and ARI values were multiplied by 100 for comparison purposes. (a) Clustering performance evaluated by NMI evaluation metric. (b) Clustering performance evaluated by ARI evaluation metric.
GSE36552 dataset and worse on three datasets compared to scEFSC using eight clustering algorithms, indicating that SC3 plays an important role in consensus clustering. Furthermore, scEFSC clustered better on eight datasets, the same on the GSE36552 and GSE81252 datasets and worse on four datasets compared to scEFSC using six clustering algorithms, suggesting that CIDR, SHARP and pcaReduce, the three clustering algorithms with the worst average clustering results, also play a positive role in the final clustering result for consensus clustering.

In summary, the selection of 5000 genes with a non-negative kernel autoencoder and 2000 genes with unsupervised feature selections and applying nine clustering algorithms in the multiple clustering method are meaningful in enhancing the clustering advantages of the proposed scEFSC algorithm.

3.7. The running times of scEFSC with other single-cell clustering algorithms

We compared the running time of scEFSC with other single-cell clustering algorithms, including SC3, CIDR, monolcle, pcaReduce, Rphenograph, Seurat, SHARP, SINCERA, and RaceID. All these algorithms were run on Ubuntu 20.04.3 LTS (GNU/Linux 5.4.0–90-generic x86_64). The running time comparisons are summarized in Supplementary Table S3 for fourteen scRNA-seq data. Since scEFSC performs multiple feature selection and multiple clustering, scEFSC has the longest running time compared to other single-cell clustering algorithms.

3.8. Evaluations on a large-scale scRNA-seq dataset GSE109774

Due to the development of scRNA-seq technology, an increasing number of cells are sequenced. Therefore, to demonstrate the scalability of our proposed scEFSC on large datasets, we conducted experiments on the GSE109774 dataset. The GSE109774 dataset is from Mouse tissues with 54,439 cells and 40 celltypes. Since computational resources were limited, we downsampled the large-scale GSE109774 dataset with 16,332 cells (about 30% of the cells). Then we compared scEFSC, SC3, CIDR, monolcle, pcaReduce, Rphenograph, Seurat, SHARP, SINCERA, RaceID, scDHA, SAME, scCCESS and RSEC on GSE109774. Fig. 7 summarizes the performance comparisons under two evaluation metrics NMI and ARI. From the figure, it can be seen that our proposed scEFSC provided better performance than most other algorithms on the large dataset. Compared to SC3, CIDR, monolcle, pcaReduce, Rphenograph, Seurat, SHARP, SINCERA and RaceID, we found that the clustering performance of scEFSC was worse than Rphenograph and SINCERA,
the same as SC3, and better than the other algorithms. SINCERA and Rphenograph had the best clustering performance, CIDR had the worst clustering performance, and SHARP had only better clustering performance than CIDR. Compared to scDHA, the clustering performance of scEFSC was better than scDHA. Compared to SAME, scCCESS and RSEC, the clustering performance of scEFSC was slightly worse than scCCESS, and better than SAME and RSEC. The clustering performance of RSEC was the worst among the three ensemble clustering algorithms, SAME, scCCESS and RSEC. Therefore, our proposed scEFSC is scalable for large-scale scRNA-seq data.

3.9. Functional genomic analysis

Although scEFSC shows superiority in the clustering performance analyses described above, the biological significance of its clustering results is particularly important for the understanding of the biological data. The biological analysis of differentially expressed genes, gene ontology enrichment and KEGG pathway analysis can be performed from the gene expression data and clustering labels obtained from scEFSC. Next, we elucidated the biological significance of scEFSC from the GSE81861 dataset of human colorectal tumour cells.

3.9.1. Differential expressed genes analysis

We used the gene expression matrix obtained from the GSE81861 dataset and the clustering labels from scEFSC to identify differentially expressed genes (DEGs) and marker genes for each cluster. Fig. 8 shows a volcano plot of each cluster analysed for differentially expressed genes, and the top twenty differentially expressed genes are labelled. The top twenty DEGs showed a high-fold change, indicating a greater difference in gene expression from the rest of the cluster and representing the marker genes for that cluster. In this analysis, we found the most differentially expressed genes...
expressed gene was ALDH1A1, which is a member of the ALDH1 gene family. The ALDH1A1 gene is an important marker gene for human colorectal carcinoma stem cells and encodes an enzyme responsible for intracellular aldehyde oxidation, shown to have a function in the early differentiation of stem cells. Its expression profile shows potential as a universal marker for the identification and isolation of normal and cancer stem cells of multiple origins in a variety of tissue types. The remaining genes with large differences were AKR1B10, CYP24A1, and AKR1C3. AKR1B10 is a pathological diagnostic marker for tumours and is highly expressed in normal gastrointestinal epithelial tissues with low or no expression in other normal tissues, and highly significant expression in breast, liver and non-small cell lung cancers. CYP24A1 is a potential oncogene, which is aberrantly expressed in breast, lung and oesophageal cancer tissues and correlates with tumour malignancy and poor patient prognosis, based on its aberrant expression regulated by DNA methylation in some tumour types. AKR1C3 is a member of the AKR1 family and involved in synthesis of a steroid hormone closely associated with the development of many malignancies and also involved in regulating the sensitivity of many anti-tumour drugs.

3.9.2. Gene Ontology (GO) enrichment and KEGG analysis

From the differential gene expression analysis on the GSE81861 dataset, we selected the top twenty differentially expressed genes from each of the nine clusters and removed duplicate genes to obtain 162 genes. These 162 genes were queried by gene ID transformation to 153 genes for gene ontology (GO) enrichment analysis and KEGG pathway analysis to interpret the biological significance of the genes. For the GO enrichment analysis, we obtained 533 enriched GOs, of which 465 were biological processes (BP), 31 were cellular components (CC) and 37 were molecular functions (MF). Fig. 9a shows the top 20 categories of the three GO enrichments, ordered by adjusted p-value. The five most enriched GO biological processes were regulation of lymphocyte activation (GO:0051249), positive regulation of cell activation (GO:0050867), antigen receptor-mediated signaling pathway (GO:0005851), positive regulation of leukocyte activation (GO:0002696) and positive regulation of lymphocyte activation (GO:0051251). We can see that most of the five enriched biological processes were related to signaling pathways and the positive regulation of cell activation, which are highly relevant to cancer.

The top five enriched GO cellular components are MHC class II protein complex (GO:0042613), blood microparticle (GO:0072562), external side of the plasma membrane (GO:0009897), MHC protein complex (GO:0042611) and integral component of lumenal side of endoplasmic reticulum membrane (GO:0071556). The MHC protein complex is closely associated with tumours [45] and there is a clear correlation between MHC-I gene composition and the genes that are mutated in the cancer of this person. The molecule MHC-II [46] may have a greater effect on neoplastic tumours than MHC-I. The remaining three enriched cellular components were all associated with the colorectum. The top five enriched GO molecular functions were alditol: NADP+ 1-oxidoreductase activity (GO:0004032), alcohol dehydrogenase (NADP+) activity (GO:0008106), aldo–keto reductase (NADP) activity (GO:0004033), antigen binding (GO:0023026) and MHC class II protein complex binding (GO:0023026). Expression of MHC-II and related pathway components have been found in cancer cells from colorectal cancer. Tumor-specific MHC-II expression can increase...
the recognition of tumors by the immune system and therefore could play an important role in immunotherapy. Fig. 9c depicts the distribution of genes under the three GO enrichments. (b) Top 9 pathways of KEGG enrichment sorted by adjusted p-values. (c) The network diagram of the top 9 pathways and genes for the KEGG enrichment.

For the KEGG analysis, 9 pathways were matched and 70 genes were successfully annotated. Fig. 9b depicts the 9 KEGG pathways sorted by adjusted p-value. Fig. 9d depicts the network diagram of the top 9 KEGG pathways with genes. This plot shows that genes HLA-DRA, HLA-DRB1 and HLA-DPB1 were matched to most of the KEGG pathways. Human leukocyte antigens (HLA) are the expression products of the major histocompatibility complex (MHC) in humans. HLA plays a key role in the immune effect mechanism of the body against tumors, in which cellular immunity plays a leading role and is regulated by humoral immunity to synergistically kill tumor cells [47]. The top five KEGG pathways are Cysteine and methionine metabolism (hsa00270), Viral myocarditis (hsa05416), Th17 cell differentiation (hsa04659), Asthma (hsa05310) and Wnt signaling pathway (hsa04310). Fig. 10 shows Th17 cell differentiation (hsa04659) and Wnt signaling pathway (hsa04310) provided by the KEGG database. The Th17 cell differentiation belongs to the immune system of biological systems, has five related genes and is highly associated with tumors. Th17 cells are a class of CD4 + effector T cells. Th17 cells and their associated cytokines have been found to be presented in many tumors, taking an important role in inflammation-associated tumors [48]. The Wnt signalling pathway is the part of signal transduction in environmental information processing. This pathway has six related
genes and is closely associated with tumors. Indeed, the Wnt signaling pathway is not only associated with events related to tumor invasion and metastasis, but also plays an essential role in regulating the self-renewal, proliferation and differentiation of tumor stem cells [49].

The GO and KEGG enrichment results reflected that the clustering results of scEFSC can effectively guide the identification of cancer-related biological processes. We can conclude therefore, that the clustering results of scEFSC are biologically significant.

Fig. 10. (a) The KEGG database indicates Th17 cell differentiation (hsa04659). (b) The KEGG database indicates the Wnt signaling pathway (hsa04310).
4. Conclusions

In this study, we propose a clustering method scEFSC based on ensemble feature selection. scEFSC is a method for clustering and analyzing the scRNA-seq data through feature selection and ensemble clustering. First, the feature selection is divided into two steps, the first step is a preliminary feature selection employing an non-negative autoencoder, and the second step uses four unsupervised feature selection algorithms to further select the results achieved by step one. For the high-dimensional sparse single-cell RNA-seq data, feature selection reduces the data dimensionality and selects effective features. Multiple clustering methods are operated on the data after feature selections. Multiple clustering methods are executed on multiple feature subsets to obtain multiple base clustering results. Then, multiple base clustering results are achieved for ensemble clustering with a high degree of variability, which facilitates the generation of better final consensus clustering results. Finally, the least diverse labels are removed and then consensus clustering is performed to obtain the final clustering results.

To demonstrate the effectiveness of the scEFSC clustering result, we tested fourteen publicly available scRNA-seq datasets. The experimental results revealed that our proposed scEFSC significantly outperformed other single-cell clustering algorithms in terms of ARI and NMI evaluation metrics. In addition, we performed differential gene expression analysis, gene ontology enrichment and KEGG analysis on the clustering results of scEFSC to validate the biological interpretability of the clustering results.

CRediT authorship contribution statement

Chuang Bian: Writing – original draft, Methodology, Investigation. Xubin Wang: Methodology, Investigation. Yanchi Su: Investigation. Yunhe Wang: Conceptualization, Writing – review & editing. Ka-chun Wong: Validation, Writing – review & editing. Xiangtao Li: Supervision, Conceptualization, Validation, Writing – review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version, at https://doi.org/10.1016/j.csbj.2022.04.023.

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