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Exploring the optimization of autoencoder design for imputing single-cell RNA sequencing data



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ABSTRACT

Autoencoders are the backbones of many imputation methods that aim to relieve the sparsity issue in single-cell RNA sequencing (scRNA-seq) data. The imputation performance of an autoencoder relies on both the neural network architecture and the hyperparameter choice. So far, literature in the single-cell field lacks a formal discussion on how to design the neural network and choose the hyperparameters. Here, we conducted an empirical study to answer this question. Our study used many real and simulated scRNA-seq datasets to examine the impacts of the neural network architecture, the activation function, and the regularization strategy on imputation accuracy and downstream analyses. Our results show that (i) deeper and narrower autoencoders generally lead to better imputation performance; (ii) the sigmoid and tanh activation functions consistently outperform other commonly used functions including ReLU; (iii) regularization improves the accuracy of imputation and downstream cell clustering and DE gene analyses. Notably, our results differ from common practices in the computer vision field regarding the activation function and the regularization strategy. Overall, our study offers practical guidance on how to optimize the autoencoder design for scRNA-seq data imputation.

1. Introduction

Single-cell RNA-sequencing (scRNA-seq) enables the measurement of genome-wide gene expression at the single-cell level [1–3]. State-of-the-art scRNA-seq technologies can measure tens of thousands of genes and up to millions of cells [4], allowing the investigation of cell-to-cell heterogeneity [5], the identification of cell types [6], and the inference of cell state transitions [7]. A notable characteristic of scRNA-seq data is the high proportion of zeros (i.e., high sparsity). Depending on the sequencing platform and sequencing depth, the zero proportion can range from 50% to more than 90% [8]. Two types of zeros exist in scRNA-seq data: biological zeros and non-biological zeros [9]. Biological zeros indicate the actual absence of gene expression in cells, while non-biological zeros are originated from technical biases and noise in scRNA-seq experiments [10]. Without spike-ins or prior biological knowledge, it is difficult to distinguish between these two types of zeros in scRNA-seq data [8].

The high sparsity of scRNA-seq data hinders data analysis. Many

computational methods have been developed to reduce data sparsity, and they can be divided into three broad categories [8]. First, model-based imputation methods use probabilistic models to describe gene expression distributions in scRNA-seq data (e.g., scImpute [11] and BISCUIT [12]). These methods aim to first distinguish between biological zeros and non-biological zeros based on gene expression distributions and then only impute the identified non-biological zeros. Second, data-smoothing methods modify each cells' gene expression levels based on its similar cells' gene expression levels (e.g., MAGIC [13] and DrImpute [14]). Two cells may be defined as similar if they share common neighboring cells in a low-dimensional space. Unlike model-based imputation methods, data-smoothing methods do not distinguish non-biological zeros and alter all gene expression levels, including the nonzeros. Third, data-reconstruction methods first learn a latent low-dimensional space of cells and then reconstruct cells in the original high-dimensional space (e.g., ZIFA [15] and DCA [16]). Most data-reconstruction methods use the reconstructed data to replace all zeros in the original data but keep the original nonzeros unchanged.

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Among the data-reconstruction methods, autoencoder-based methods are popular for their imputation accuracy and scalability [17, 18]. Specifically, an autoencoder is a neural network that contains an encoder for dimension reduction and a decoder for data reconstruction. Then the reconstructed data are regarded as the imputed scRNA-seq data. Despite the popularity of autoencoders in the scRNA-seq field, there lacks a formal discussion on how to optimize the autoencoder design, including the neural network architecture, the activation function, and the parameter regularization [8]. Existing autoencoder-based methods either adopt the wisdom in other fields (e.g., computer vision) or design autoencoders with limited justification.

Here, we conduct an empirical study to explore how to optimize the autoencoder design for imputing scRNA-seq data. Our study consists of three investigations. The first investigation is the optimization of autoencoder design for imputation accuracy. In detail, we generate 36 semi-synthetic scRNA-seq datasets with artificial zeros (whose original, non-missing values are known) by applying three masking schemes to 12 real scRNA-seq datasets. Then, we train autoencoders with varying depths (depth means the number of layers) and widths (width means the number of nodes per layer), seven activation functions, and two regularization strategies on each semi-synthetic dataset. Next, we evaluate the trained autoencoders' imputation accuracies in terms of the normalized root mean squared error (NRMSE) and the Pearson correlation coefficient, which are calculated between the imputed values and the original values of the artificial zeros. Based on the evaluation results, we find the autoencoder designs that lead to the best imputation accuracies.

The second investigation is the optimization of autoencoder design for cell clustering. In detail, we train autoencoders with the aforementioned designs on 20 real scRNA-seq datasets containing curated cell type information. Next, to examine the impact of autoencoder design on downstream cell clustering, we apply the trained autoencoders for imputation and evaluate the clustering accuracies on the imputed datasets in terms of the adjusted Rand index (ARI) and the adjusted mutual information (AMI).

The third investigation is the optimization of autoencoder design for differentially expressed (DE) gene identification. In detail, we simulate 20 synthetic datasets with ground-truth DE genes by applying the scDesign simulator [19], which is trained on 20 real scRNA-seq datasets (see Methods). Then we train autoencoders with the aforementioned designs on these synthetic datasets. Next, to examine the impact of autoencoder design on downstream DE gene analysis, we apply the trained autoencoders for imputation and evaluate the DE gene identification accuracies on the imputed datasets in terms of the precision, recall, and true negative rate (TNR).

In the above three investigations, we consider many choices of neural network depths, widths, activation functions, and regularization strategies. Since a full combination of those design aspects is computationally infeasible, we adopt a sequential, greedy search strategy to explore the best design. In each investigation, we first examine a full combination of neural network depths and widths, the two most important aspects, with the other design aspects fixed as in common practices (see Results). Based on the optimized combination of width and depth, we next optimize the activation function. Finally, given the optimized width, depth, and activation function, we optimize the regularization strategy.

Our results yield the following guidelines about autoencoder design for scRNA-seq data imputation (Table 1). First, deeper and narrower autoencoders lead to more accurate imputation, cell clustering, and DE gene identification, yet the benefit of depth saturates at 10 hidden layers. Second, the sigmoid and tanh activation functions consistently have the best performance in all evaluations. Third, parameter regularization is critical to the performance of autoencoder-based imputation methods. In particular, weight decay regularization is more capable of improving cell clustering and DE gene analysis, while dropout regularization shows superiority in improving the overall imputation accuracy

Table 1

Guidelines of autoencoder design for scRNA-seq data imputation. Each row represents one of four design characteristics. Each column represents one of three evaluation metrics. Each entry corresponds to the marginally optimized design characteristic under each evaluation metric.

Autoencoder design characteristics	Evaluation metrics		
	Imputation accuracy	Cell clustering	DE gene identification
Number of hidden layers Number of units per hidden layer	≥ 10 32	≥ 10 Robust	\geq 5 Robust
Activation function	Sigmoid or Tanh	Sigmoid or Tanh	Sigmoid or Tanh
Regularization strategy	Dropout	Weight decay	Weight decay

for the artificial zeros. Moreover, the optimal degree of regularization is dataset-specific.

Our findings suggest that many autoencoder-based imputation methods have used suboptimal autoencoder designs, including shallower and wider autoencoders and the rectified linear unit (ReLU) activation function. Moreover, our findings highlight the importance of using empirical benchmarking to optimize autoencoder designs, and more generally, neural network designs, in bioinformatics research.

2. Results

2.1. Impacts of autoencoder architecture (depth and width) on imputation accuracy

We collect 12 real scRNA-seq datasets to evaluate the overall imputation accuracy of a variety of autoencoder architectures. These datasets cover a wide range of cell types, sequencing depths, zero proportions, and experimental platforms (Supplementary Table S1). We apply three masking schemes (i.e., random masking, double exponential masking, and medium masking, which reflect different degrees of dependence of missingness on the actual values; see Methods) to these 12 real datasets, obtaining 36 masked datasets. To evaluate the imputation accuracy on each masked dataset, we calculate the NRMSE and the Pearson correlation coefficient between the masked values and the imputed values (Methods), referred to as the "imputation NRMSE" and the "imputation correlation," respectively, in the following text.

We build autoencoders of various architectures by increasing the depth (i.e., the number of hidden layers) from 1 to 15. For each depth, we set the width (i.e., the number of hidden units per layer) to 32, 64, 128, or 256. All hidden layers are fully connected with the same width. In total, we have 60 autoencoder architectures, corresponding to 15×4 depth-width combinations. We choose the ReLU function [20] as the activation function and train the autoencoders by the Adam optimization algorithm [21] (Methods). For each autoencoder architecture and each masked dataset, we set 10 random seeds in the autoencoder training to obtain 10 autoencoders, whose imputation NRMSEs and imputation correlations are averaged to represent the overall imputation accuracy of the autoencoder architecture on the masked dataset.

Under the random masking scheme, Fig. 2a–b show the impacts of autoencoder depth-width combinations on the imputation NRMSE and the imputation correlation. First, deeper autoencoders achieve lower imputation NRMSEs and higher imputation correlations. The benefit of depth is more prominent when the autoencoder has no more than 10 layers. Second, narrower autoencoders (with 32 hidden units per layer) typically have higher imputation accuracy than wider autoencoders (with 64 or more hidden units per layer) of the same depth. This finding is consistent with the observation that deeper and narrower neural networks perform better in computer vision tasks (e.g., image classification and object detection) [22,23]. Under the double exponential masking scheme, we observe a similar relationship between the



Fig. 1. Autoencoder and the measurement of imputation accuracy. a, The basic structure of an autoencoder [24]. b, The introduction of artificial zeros by masking. c, The training of an autoencoder for imputation. The reconstructed data are the autoencoder's output during the training process. d, The calculation of an autoencoder's imputation accuracy on masked values. The imputed data are the final autoencoder's output after the training process.

autoencoder depth-width combinations and the imputation accuracy, except for the dataset bmmc (Supplementary Fig. S1). However, the relationship no longer holds under the median masking scheme: while deeper autoencoders still have better imputation accuracy on certain datasets, the width does not have a significant impact on the imputation accuracy (Supplementary Fig. S2). Among the three masking schemes, random masking has the highest imputation accuracy, followed by double exponential masking and then median masking. Note that under median masking, the imputation accuracy is low on most datasets, indicated by many larger-than-one imputation NRMSEs and close-to-zero imputation correlations (Supplementary Fig. S2).

A possible explanation of the results is that, among the three masking schemes, random masking best preserves gene-gene correlations. Since random masking is performed independent of gene expression levels, the gene-gene correlations in the masked data are unbiased estimates of the gene-gene correlations in the original, unmasked data. Unlike random masking, double exponential masking assumes an inverse relationship between a gene's masking probability and average expression (i.e., the probability of masking increases as a gene's average expression level decreases); hence, the gene-gene correlations in the masked data are no longer unbiased estimates of the gene-gene correlations in the original data. While random masking and double exponential masking are both stochastic schemes, median masking is a deterministic scheme where the 50% lowest expression levels of all genes are masked as zeros. Hence, median masking distorts gene-gene correlations to the greatest extent among the three masking schemes. Under the hypothesis that gene-gene correlations play a crucial role in determining the imputation accuracy of an autoencoder, it is unsurprising that the best imputation accuracy is achieved under the random masking scheme, followed by the double exponential masking scheme. Hence, the following analysis on imputation accuracy will focus on random masking and double exponential masking.

2.2. Impacts of activation function on imputation accuracy

An activation function is a nonlinear transformation applied to generate the hidden units in an autoencoder [25]. It provides an autoencoder with the capacity to learn complex nonlinear relationships among features, e.g., nonlinear interactions among genes in scRNA-seq data. ReLU is a widely used activation function in autoencoder-based imputation methods, motivated by its success in computer vision [25]. However, justification is lacking for using ReLU to impute scRNA-seq data, and no empirical comparison has been done between ReLU and other activation functions.

Here, we train autoencoders with seven activation functions, including sigmoid, tanh, ReLU, LeakyReLU (with two hyperparameter settings) [26], ELU [27], and SELU [28], to evaluate the impacts of activation functions on the imputation accuracy (Methods). For each activation function, we impute 24 masked datasets (the aforementioned 12 scRNA-seq datasets after random masking or double exponential masking) by training 20 autoencoders using 20 random seeds on each masked dataset. Fig. 3 and Supplementary Fig. S3 compare the seven activation functions' resulting imputation NRMSEs and imputation correlations under the random masking and double exponential masking schemes. Surprisingly, the most popular ReLU function is not the top performer. Instead, the sigmoid and tanh functions outperform the other activation functions on all datasets under both masking schemes. Additionally, the sigmoid and tanh functions result in significantly smaller variances of the imputation NRMSEs and the imputation correlations than the other functions do, indicating that the sigmoid and tanh functions lead to more stable imputation accuracy. Between sigmoid and tanh, although they have similar performance, sigmoid has higher imputation accuracy on the datasets pbmc and human_mix and more stable imputation accuracy on the datasets mbrain, pbmc, human_mix, and mouse_cortex.

The comparison of the seven activation functions reveals three insights. First, unlike ReLU that introduces sparsity into hidden layers (i.e.,



Fig. 2. Impacts of the autoencoder depth and width on the imputation NRMSE (a) and the imputation correlation (b) based on the random masking scheme. Each point is the average of the results obtained from 10 random seeds used for autoencoder training.

many hidden units are zeros) [26], sigmoid and tanh generate dense hidden layers (i.e., many hidden units are nonzeros). Our empirical results suggest that dense hidden layers lead to higher imputation accuracy for scRNA-seq data. Second, Leaky ReLU, ELU, and SELU are modified forms of ReLU that output a close-to-zero negative value when the input is negative (Methods). They generate pseudo-sparsity in the hidden layers to avoid the "dead ReLU" problem (i.e., most hidden units are zeros, causing the autoencoder to stop learning) [29]. However, our empirical results show no consistent improvement of these modified ReLU functions over ReLU in terms of imputation accuracy, though these modified ReLU functions lead to dense hidden layers as sigmoid and tanh do. A possible interpretation is that sigmoid and tanh have continuous derivatives, while the modified ReLU functions have discontinuous derivatives. Third, we do not observe vanishing gradients or exploding gradients [30], two common problems in training deep neural networks using sigmoid or tanh. We hypothesize that appropriate preprocessing and normalization of scRNA-seq data might help stabilize gradients in the training process.



Activation Function

Fig. 3. Impacts of the activation function on the imputation NRMSE (a) and the imputation correlation (b) based on the random masking scheme. Sg: sigmoid; Th: tanh; RL: ReLU; LRL: LeakyReLU ($\alpha = 0.01$); LRL.2: LeakyReLU ($\alpha = 0.2$); EL: ELU; SEL: SELU. Each boxplot shows the results obtained from 20 random seeds used for autoencoder training.

2.3. Impacts of regularization on imputation accuracy

Some autoencoder-based imputation methods use weight decay or dropout as the regularization strategy [31,32]. However, the selection of the regularization strategy and the corresponding hyperparameters (i.e., λ in weight decay and p in dropout, Methods) is mostly ad hoc [8]. To examine the impact of regularization on imputation accuracy, we train

autoencoders with weight decay or dropout as the regularization strategy (Methods) to impute the aforementioned 24 masked datasets (the 12 real scRNA-seq datasets after random masking or double exponential masking). We vary $\lambda \in [1e-7, 5e-4]$ and $p \in [0.01, 0.4]$ and compare the resulting imputation NRMSEs and imputation correlations (Methods; Figs. 4 and 5; Supplementary Figs. S4 and S5). All autoencoders have the same architecture and activation function: 10 fully connected hidden



Fig. 4. Impacts of the weight decay regularization on the imputation NRMSE (a) and the imputation correlation (b) based on the random masking scheme. Each point is the average of the results obtained from 10 random seeds used for autoencoder training.

layers, 32 hidden units per hidden layer, and the sigmoid activation function. For each dataset and regularization strategy (with a given value for the corresponding hyperparameter), we use 10 random seeds to train 10 autoencoders and average the resulting imputation NRMSEs and imputation correlations.

Under random masking, the weight decay strategy barely improves the imputation accuracy except on datasets mouse_spleen and human_mix. Larger values of λ even lead to worse imputation accuracy, suggesting over-regularization (Fig. 4). In contrast, the dropout strategy with a proper dropout rate p improves the imputation NRMSEs on six datasets and the imputation correlations on 11 datasets. The optimal range of p depends on the evaluation metric: 0.02–0.2 for imputation NRMSE and 0.2–0.4 for imputation correlation (Fig. 5). Hence, p = 0.2 is a reasonable choice.



Fig. 5. Impacts of the dropout regularization on the imputation NRMSE (a) and the imputation correlation (b) based on the random masking scheme. Each point is the average of the results obtained from 10 random seeds used for autoencoder training. Some y-axis tick values appear to be identical due to the four-digit resolution, reflecting the negligible variation of y-axis values across dropout rates.

Under double exponential masking, both the weight decay and dropout strategies improve the imputation accuracy (Supplementary Figs. S4 and S5). Specifically, the weight decay strategy reduces the imputation NRMSEs on six datasets and increases the imputation correlations on 11 datasets (Supplementary Fig. S4). The optimal range of λ

depends on the evaluation metric: 1e-7–5e-5 for imputation NRMSE and 5e-6–5e-4 for imputation correlation. Hence, 5e-6–5e-5 is a reasonable range for λ . The dropout strategy exhibits a stronger positive impact on the imputation accuracy, with the imputation NRMSEs reduced on 10 datasets and the imputation correlations increased on all 12 datasets

(Supplementary Fig. S5). The optimal range of the dropout rate p is similar to that under random masking.

We summarize the impacts of regularization on the imputation accuracy from three perspectives. First, compared with the weight decay strategy, the dropout strategy is more capable of improving both the imputation NRMSE and the imputation correlation under both random masking and double exponential masking. Second, both the weight decay and dropout strategies are more effective under double exponential masking than random masking. As discussed in "Impacts of autoencoder architecture (depth and width) on imputation accuracy," double exponential masking masks small nonzero values in the scRNAseq data matrix, thus distorting gene-gene correlations. In contrast,



Activation Function

Fig. 6. Impacts of the autoencoder design on cell clustering accuracy measured by ARI. a, Depth and width. b, Activation function. The dash lines in (a) show the cell clustering performance without imputation. Each point in (a) is the average of the results obtained from five random seeds used for autoencoder training. Each boxplot in (b) shows the results obtained from 10 random seeds used for autoencoder training.

random masking does not have this issue. Hence, we hypothesize that regularization has a bigger benefit under double exponential masking because regularization's improvement of robustness is more needed after double exponential masking's distortion on gene-gene correlations. Third, the optimal hyperparameter for each regularization strategy largely depends on the dataset and the masking scheme, making it impossible to find a universally optimized hyperparameter value Interestingly, compared with the imputation NRMSE, the imputed correlation requires a higher degree of regularization to be optimized, under both the weight decay and dropout strategies. Future research is required to find the reason underlying this phenomenon.

2.4. Impacts of autoencoder design (architecture, activation function, and regularization) on cell clustering

The ultimate goal of imputation is to improve downstream analysis through the enhancement of signals in the sparse scRNA-seq data [33]. We collect 20 real scRNA-seq datasets containing annotated cell types to examine the impact of autoencoder design on cell clustering (Supplementary Table S2). Here, the datasets are different from those used in the evaluation of the imputation accuracy because evaluating clustering accuracy does not require the knowledge of non-missing values. Specifically, we first conduct K-means clustering on each original dataset and calculate the adjusted Rand index (ARI) and the adjusted mutual information (AMI) to measure the clustering performance (Methods). Note that we choose K-means clustering instead of the more popular graph-based Louvain or Leiden clustering in the Seurat package [34] because we want to set K, the number of clusters, to the number of cell types for fair evaluation. While setting K to a specific number is natural for K-means clustering, it requires manual tuning for Louvain and Leiden clustering and is too labor-intensive for our evaluation. Second, we train autoencoders with various architectures, activation functions, and regularization strategies to impute the 20 datasets. Finally, we conduct K-means clustering on each imputed dataset and calculate the corresponding ARI and AMI.

Fig. 6a and Supplementary Fig. S6a show the impact of autoencoder architecture on cell clustering. Similar to the previous analysis, we increase the depth of autoencoders from 1 to 15 and set the width to 32, 64, 128, or 256 for each depth. All hidden layers in each autoencoder are fully connected with the same width. In total, we have 60 $(15\times\,4)$ depth-width combinations. We use the sigmoid activation function because of its superior performance in the previous imputation accuracy evaluation. On each dataset, we use five random seeds for autoencoder training to obtain five imputed datasets, on which we calculate the imputation ARIs and the imputation AMIs; then we report the average imputation ARI and the average imputation AMI. Our results, which are consistent between ARIs and AMIs, show that deeper autoencoders lead to a larger improvement in cell clustering accuracy than their shallower counterparts do. The benefit of depth saturates after 10 hidden layers. Unlike the autoencoder depth, the autoencoder width has no obvious impact on cell clustering accuracy (Fig. 6a; Supplementary Fig. S6a). Surprisingly, imputation does not always improve cell clustering accuracy: imputation only improves ARIs on eight datasets and AMIs on four datasets, regardless of autoencoder architectures.

Fig. 6b and Supplementary Fig. S6b show the impact of activation functions on cell clustering. Similar to the previous analysis, we train autoencoders with seven activation functions, including sigmoid, tanh, ReLU, LeakyReLU (with two different hyperparameters), ELU, and SELU. For each activation function, we impute the 20 scRNA-seq datasets using 10 autoencoders trained under 10 random seeds. All autoencoders have 10 fully connected hidden layers with 32 hidden units per layer, an optimized architecture based on our previous analysis. We observe that sigmoid and tanh outperform other activation functions on all datasets in terms of both imputation AMI and imputation ARI. They also lead to more stable cell clustering accuracy than other activation functions do. The performance of sigmoid and tanh is similar except for datasets mouse_cortex and Klein, where tanh has a slight advantage over sigmoid.

Fig. 7 and Supplementary Fig. S7 show the impact of regularization on cell clustering. Similar to the previous analysis, we use the weight decay and dropout strategies and vary $\lambda \in [1e-7, 5]$ and $p \in [0.01, 0.4]$ (Methods). Based on our previous analysis, all autoencoders have 10 fully connected hidden layers, with 32 hidden units per layer, and the sigmoid activation function. Interestingly, the weight decay strategy significantly improves cell clustering accuracy-with the optimized weight decay hyperparameter λ 's (mostly between 0.01 and 0.1), all 20 datasets have improved ARIs, and 18 datasets have improved AMIs. However, the dropout strategy does not lead to same improvementwith the optimized dropout hyperparameter *p*'s (in a broad range), only eight datasets have improved ARIs, and only four datasets have improved AMIs. Hence, unlike the previous imputation accuracy evaluation, which prefers the dropout strategy, here the weight decay strategy is preferred. However, we do not want to over-interpret the results because the autoencoders we use do not consistently lead to better ARIs and AMIs, suggesting that imputation might not be needed for cell clustering, consistent with our previous report [9].

2.5. Impacts of autoencoder design (architecture, activation function, and regularization) on DE gene analysis

The signal enhancement in scRNA-seq data by imputation is expected to benefit another important downstream analysis—the identification of differentially expressed (DE) genes. To examine the impact of autoencoder design on DE gene analysis, we utilize the simulator scDesign [19] to generate 20 synthetic datasets with ground-truth DE genes (Methods). Each synthetic dataset is generated by learning the distributions of genes' expression levels in one real scRNA-seq dataset (20 real datasets in total; Supplementary Table S3). These real datasets (and their synthetic counterparts) cover a wide range of biological and technical conditions. We use synthetic data in this analysis since the ground-truth DE genes are unknown in real scRNA-seq datasets.

After simulation, we apply the MAST method [35] to the pre-imputed synthetic datasets to identify DE genes and calculate the corresponding precision, recall, and true negative rate (TNR), which are considered as the baseline accuracies. Next, to impute each synthetic dataset, we train autoencoders with various architectures, activation functions, and regularization strategies. Finally, we apply MAST to each imputed dataset and calculate the corresponding precision, recall, and TNR (Methods).

Fig. 8a, Supplementary Fig. S8a, and Supplementary Fig. S10a show the impacts of autoencoder architecture on the recall, precision, and TNR. The settings of depth, width and activation function for autoencoders are the same as in the evaluation of cell clustering (1-15 fullyconnected hidden layers; 32, 64, 128, or 256 hidden units per layer; sigmoid activation function). On each synthetic dataset, we set five random seeds in the autoencoder training to obtain five imputed datasets, whose precisions, recalls, and TNRs are averaged to represent the autoencoder's performance. First, deeper autoencoders lead to better recalls on 10 datasets, while the benefit of depth saturates after five layers (Fig. 8a). In contrast, width has no significant improvement on the recall. Unexpectedly, 11 synthetic datasets show no improvement in recall after imputation, regardless of the autoencoder architecture. Second, depth and width have no significant impact on the precision, except for the datasets Interneurons, Epithelial_cells, and astrocytes, where deeper autoencoders lead to slightly higher precisions (Supplementary Fig. S8a). Overall, 19 synthetic datasets have improved precisions after imputation. Third, the impact of depth and width on the TNR is limited, because all TNRs are already close to one before imputation, and most TNRs are increased by less than 0.05 after imputation (Supplementary Fig. S10a). Altogether, our results suggest that autoencoder-based imputation benefits the precision instead of the recall or the TNR for the identification of DE genes.



Fig. 7. Impact of the autoencoder regularization strategy on cell clustering accuracy measured by ARI. a, Weight decay regularization. b, Dropout regularization. The dash lines show the cell clustering performance without imputation. Each point is the average of the results obtained from five random seeds used for autoencoder training.

Fig. 8b, Supplementary Fig. S8b, and Supplementary Fig. S10b show the impacts of the activation function on the recall, precision, and TNR. Again, we train autoencoders with seven activation functions; for each function, we impute the aforementioned 20 synthetic datasets using 10 autoencoders trained by setting 10 random seeds. All autoencoders have 10 fully connected hidden layers with 32 hidden units per layer. In terms of the recall, sigmoid and tanh outperform the other activation functions on 15 datasets (Fig. 8b). The advantage of sigmoid and tanh is similar for the precision—they outperform the other activation functions on 12 datasets (Supplementary Fig. S8b). All activation functions have similar performance based on the TNR, which is consistently close to one (Supplementary Fig. S10b). Overall, sigmoid and tanh have similar



Fig. 8. Impacts of the autoencoder design on DE gene identification accuracy measured by recall. **a**, Depth and width. **b**, Activation function. The dash lines in (**a**) show the recall without imputation. Each point in (**a**) is the average of the results obtained from five random seeds used for autoencoder training. Each boxplot in (**b**) shows the results obtained from 10 random seeds used for autoencoder training.

performance, and they both provide a more stable improvement in the identification of DE genes than the other activation functions do.

Fig. 9, Supplementary Fig. S9, and Supplementary Fig. S11 show the impacts of regularization on the recall, precision, and TNR. Again, we add either weight decay or dropout to autoencoders and adjust the corresponding hyperparameter, as in the previous analysis. All autoencoders have 10 fully connected hidden layers, with 32 hidden units per layer, and the sigmoid activation function. We observe that, compared

to dropout, weight decay exhibits a stronger improvement in recall, precision, and TNR: weight decay outperforms the no-regularization autoencoders on 20, 12, and 20 datasets, respectively, with the optimal hyperparameter λ 's (in the range of [5e-6, 0.05]) (Fig. 9a, Supplementary Fig. S9a, and Supplementary Fig. S11a); in contrast, dropout outperforms the no-regularization autoencoders on 17, 6, and 17 datasets, respectively, with the optimal *p*'s (in the range of [0.05, 0.25]) (Fig. 9b, Supplementary Fig. S9b, and Supplementary Fig. S11b).



Fig. 9. Impacts of the autoencoder regularization strategy on DE gene identification accuracy measured by recall. **a**, Weight decay regularization. **b**, Dropout regularization. The dash lines show the recall without imputation. Each point is the average of the results obtained from five random seeds used for autoencoder training.

Moreover, weight decay with the optimal λ 's improves the recall, precision, and TNR from the baseline values (before imputation) on every dataset, while dropout with the optimal *p*'s fails to improve the recall and TNR from the baseline values on 6 and 20 datasets, respectively. The preference of weight decay over dropout is similar to our evaluation of cell clustering but different from our evaluation of imputation accuracy. A possible reason is that we use manual masking (random or double exponential) in the evaluation of imputation accuracy but not in the evaluation of cell clustering or DE gene identification. Our results reveal the importance of evaluation metrics in autoencoder design. However, despite the discrepancy in preferring weight decay or dropout, our evaluation results consistently suggest that regularization is crucial to the imputation performance of autoencoders.

3. Discussion

Sparsity is one of the major hurdles in scRNA-seq data analysis [36]. To address the sparsity issue, more than 70 computational imputation methods have been developed, many of which are autoencoder-based [37], motivated by the success of neural networks in computer vision and natural language processing [38]. Compared with traditional statistical and machine learning methods, autoencoder-based imputation methods have unique characteristics. First, autoencoder-based imputation methods require no assumptions on the underlying distribution of scRNA-seq data. This data-driven characteristic avoids the model specification bias in traditional methods. Second, autoencoder-based imputation methods can effectively handle large-scale scRNA-seq data by innovative hardware. especially the using GPU. Third. autoencoder-based imputation methods have high flexibility granted by their neural network design. With a carefully-designed loss function, they have incorporated multiple functionalities such as imputation, dimension reduction, and batch effect normalization [39].

However, how to optimize the design of autoencoders for scRNA-seq data remains a challenge. Successful applications of autoencoders in other fields, especially computer vision, rely on empirical studies that optimize the autoencoder design on massive datasets. Existing autoencoder-based imputation methods for scRNA-seq data follow certain guidelines (e.g., using the ReLU activation function) but ignore other guidelines (e.g., using a deep and narrow neural network) without comprehensive investigations. Our study partially fills this gap, and our results confirm the deep learning field's common wisdom that deeper and narrower autoencoders have better imputation performance. Meanwhile, we have an unexpected finding that sigmoid and tanh outperform ReLU and ReLU-modified forms as activation functions. This finding is consistent with another study, in which a neural network with the tanh activation function outperforms the ReLU counterpart in the cell-type classification task based on scRNA-seq data [40], reflecting the fact that scRNA-seq data are distinct from image data and thus need specific investigations.

Although our results favor deep and narrow neural networks, a previous study found that deep neural networks do not improve celltype classification on scRNA-seq data [41]. We hypothesize that this discrepancy is due to the two tasks' different natures: our imputation task is a regression problem whose predictive targets are continuous gene expression values (after preprocessing), while the previous study's task is a classification problem whose predictive targets are discrete cell-type labels. We hypothesize that, compared to classification, imputation requires deeper neural networks with a higher predictive capacity. Moreover, even though deeper autoencoders exhibit advantages in our study, the benefit of depth saturates when the number of layers surpasses 10, which is a much shallower architecture compared to the state-of-the-art deep neural networks with hundreds of layers used in computer vision. However, this discrepancy is consistent with the relationship between data complexity and model capacity. One image is typically a three-dimensional tensor (RGB channels \times width \times length) [42] and much more complex than one cell encoded in a one-dimensional vector in scRNA-seq data. Hence, it is unsurprising that scRNA-seq data need a shallower neural work than image data do.

Regarding the regularization strategy, we find that dropout excels in improving the overall imputation accuracy, and weight decay (using the L_2 penalty) excels in improving the downstream cell clustering and DE gene analysis. Although dropout does not penalize all weight parameters, it randomly sets a certain proportion of weight parameters to zeros and thus can be interpreted as a stochastic L_1 penalization. From this point of view, the L_1 and L_2 penalizations have complementary advantages in improving the overall imputation accuracy and the downstream cell clustering and DE gene analysis, respectively. We should also note that the three masking schemes are simplified approximations of, but are not themselves, the true missing mechanism, and the ultimate goal of imputation is to enhance downstream analyses. Therefore, weight decay may provide stronger benefits in real-world applications.

We observe that a similar autoencoder design exhibits distinctive imputation performance on scRNA-seq datasets with different characteristics. Datasets generated by 10X Genomics, e.g., jurkat, monocyte, and 293 t, show higher overall imputation accuracies (Fig. 2). Those datasets share some common characteristics, including high throughput (i.e., large numbers of cells) and large zero proportions (Supplementary Table S1). On the other hand, datasets generated by Smart-seq-total, Smart-seq2, and Fluidigm C1 show lower overall imputation accuracies (Fig. 2). Those datasets have lower throughput with smaller zero proportions (Supplementary Table S1). When it comes to enhancing downstream cell clustering, datasets that contain more ground-truth cell types, such as manno_human, chen, and lake, pose a greater challenge for autoencoders to enhance the separation of cell types. This is evident from Fig. 6 and Supplementary Table S2. Even when the correct number of clusters is set in K-means clustering, it is more difficult for autoencoders to improve the cell clustering in those datasets. We also observe that autoencoders are less effective at improving DE gene identification on smaller datasets, such as Astrocytes, Macrophages, and Tanycytes, which only have hundreds of cells (Supplementary Table S3). Since the number of ground-truth DE genes is the same for all datasets, we suspect that the number of cells plays a critical role in the effectiveness of autoencoder-based imputation methods.

Our findings indicate that the performance of autoencoder-based imputation methods is contingent upon the data characteristics and the downstream tasks involved. While deep autoencoders have the ability to capture non-linear and complex relationships in scRNA-seq data, their effectiveness may be limited in datasets with low complexity. As a result, we observe performance saturation as the number of layers increases to a point (Figs. 2, 6, and 8). Considering the higher computational resources required for deeper autoencoders, it becomes crucial to balance the tradeoff between imputation performance and computational time for autoencoder design. Researchers must weigh the potential gains in imputation performance against the increased computational demands associated with deeper autoencoders.

Our study is conducted based on the original autoencoder. Some imputation methods use variants extended from the original autoencoder. For example, scVI is based on a variational autoencoder to learn a probabilistic latent space of the input data [39]; scScope adopts an iterative autoencoder to impute data using many iterations [43]; DCA uses the negative log-likelihood of a zero-inflated negative binomial model as the loss function to estimate the parameters of an autoencoder [16]. Despite their differences, these autoencoder variants all have the encoder-decoder neural network structure and a non-linear activation function, with some including a regularization strategy. Therefore, our study can be easily generalizable to these autoencoder variants.

A major limitation of our study is that we adopt a greedy search strategy, instead of a global search strategy, to optimize the autoencoder design, due to the high computation complexity of autoencoder training. Hence, the autoencoder designs we find as optimal in this study are not guaranteed to be globally optimal. One potential solution is to use advanced experimental design strategies, such as the space-filling design [44,45] and the fractional factorial design [46,47], to expand the search space so that the optimized design is closer to the global optimum. Another limitation is that we set all hidden layers to the same width (32, 64, 128, or 256) in each autoencoder, aiming to reduce the complexity of varying widths. Although there is no consensus on how the width heterogeneity across layers would affect an autoencoder's learning capacity [48], it is possible that an autoencoder with hidden layers of different widths might have better imputation performance. We will leave these two improvements for future research.

In summary, the performance of autoencoder-based imputation methods relies on key aspects of the autoencoder design, including the neural network architecture, activation function, and regularization strategy. Borrowing guidelines from practices in other fields cannot guarantee good performance on scRNA-seq data. Future methodological development should pay more attention to optimizing the autoencoder design and allow users to adjust the design for application needs.

4. Methods

4.1. Autoencoder for imputing scRNA-seq data

An autoencoder is a multi-layer neural network that aims to reconstruct the input data [49]; it includes an encoder that embeds high-dimensional data into a low-dimensional latent space and a decoder that reconstructs the high-dimensional data from the low-dimensional embeddings (Fig. 1a). Let *X* be a sparse scRNA-seq input matrix after appropriate preprocessing and normalization (see Methods section "Data preprocessing and normalization"). Let *Y* be the dense matrix output by an autoencoder. Both *X* and *Y* have *n* rows (cells) and *m* columns (genes). Denote H_k , an *n*-by- l_k matrix, as the output of the *k*th hidden layer (with width l_k) of the autoencoder, where k = 1, 2, ..., h, with *h* as the total number of hidden layers. Then the output of the first hidden layer is

$$H_1 = f(XW_1 + B_1)$$

where W_1 is an *m*-by- l_1 weight matrix, and B_1 is an *n*-by- l_1 bias matrix with *n* identical rows. Here, *f* is an element-wise nonlinear activation function. Similarly, the output of the (k+1)th hidden layer is

$$H_{k+1} = f (H_k W_{k+1} + B_{k+1}),$$

where W_{k+1} is an l_k -by- l_{k+1} weight matrix, and B_{k+1} is an *n*-by- l_{k+1} bias matrix with *n* identical rows. Finally, the output of the autoencoder is

 $Y = H_h W_{h+1} + B_{h+1},$

where W_{h+1} is an l_h -by-*m* weight matrix and B_h is an *n*-by-*m* bias matrix with *n* identical rows. Training the autoencoder means estimating (learning) the parameters in the weight matrices $W_1, W_2, ..., W_{h+1}$ and the bias matrices $B_1, B_2, ..., B_{h+1}$ by minimizing the mean squared error (MSE) between the input *X* and the output *Y* on the nonzero values of *X*. Let \mathcal{W} be the set of weight matrices $W_1, W_2, ..., W_{h+1}$ and \mathcal{B} be the set of bias matrices $B_1, B_2, ..., B_{h+1}$. Then the loss function is.

$$MSE(\mathscr{W},\mathscr{B}) = \frac{\sum_{i}^{n} \sum_{j}^{m} (Y_{ij} - X_{ij})^{2} I(X_{ij} \neq 0)}{\sum_{i}^{n} \sum_{j}^{m} I(X_{ij} \neq 0)}, \text{where } I(\cdot) \text{ is an indicator func-}$$

tion. The sets of weight and bias matrices $\{\widehat{\mathscr{W}}, \widehat{\mathscr{B}}\}$ are estimated by

 $\{\widehat{\mathscr{W}},\widehat{\mathscr{B}}\} = \mathrm{argmin}_{(\mathscr{W},\mathscr{B})}\mathrm{MSE}(\mathscr{W},\mathscr{B}).$

Since the minimization of MSE is a non-convex optimization problem [48], the backpropagation algorithm [50] is utilized to train the autoencoder (see Methods section "Autoencoder training and imputation"). In the imputation step, the zero entries in the input matrix X are replaced by their nonzero counterparts in the output matrix Y. That is, the imputed scRNA-seq data matrix \hat{X} is calculated as

$$\widehat{X} = X + Y \circ I(X = 0)$$

where $^{\circ}$ is the element-wise product.

Several modifications to the original autoencoder have been made since the debut of autoencoder-based imputation methods. For example, DCA [16] models the scRNA-seq data by a negative binomial (NB) distribution with or without zero-inflation (ZI) (that is, ZINB or NB distribution) and learns the autoencoder by maximizing the likelihood of ZINB or NB calculated on the output *Y*; scVI [39] learns a variational autoencoder [51] by forcing each hidden unit in the hidden layer to follow a ZINB distribution; DeepImpute [31] learns an autoencoder by minimizing the weighted MSE, in which a gene's weight is its expression level; LATE [52] considers cells or genes as observations, learns an autoencoder for each consideration, and selects the autoencoder with a smaller MSE; scScope [43] learns an autoencoder by using the imputed data as input iteratively. Despite these modifications, all autoencoder-based imputation methods require the specification of the autoencoder design aspects we examine in this work.

4.2. Regularization in autoencoder

Regularization is a technique for constraining the complexity of a machine learning model so that the model would have better generalizability [53]. There are two regularization strategies, among others, commonly used to improve the imputation accuracy of autoencoders—weight decay [48] and dropout [54]. Weight decay incorporates the L_2 norm of weight parameters into the loss function to penalize large weights in an autoencoder. The weight and bias parameters under the weight decay $\{\widehat{\mathscr{W}}, \widehat{\mathscr{B}}\}_{\text{weight decay}}$ are given by

 $\{\widehat{\mathscr{W}}, \widehat{\mathscr{B}}\}_{\text{weight decay}} = \operatorname{argmin}_{(\mathscr{W}, \mathscr{B})} \operatorname{MSE}(\mathscr{W}, \mathscr{B}) + \lambda ||W||_2^2$, where $||W||_2$ is the L_2 norm of weight parameters, and λ is a tuning parameter that controls the degree of penalization.

Rather than penalizing the magnitudes of weight parameters, dropout regularization randomly sets a proportion of hidden units to zero in the training of an autoencoder. Dropout forces the autoencoder not to rely on particular hidden units and thus reduces overfitting [54]. Specifically, suppose that z_k is a random vector with l_k elements, where l_k is the number of hidden units in the hidden layer H_k . Each random variable in z_k independently follows a Bernoulli distribution with a parameter $p_k \in (0, 1)$. A matrix Z_k is constructed to have the same dimensions as those of the hidden layer matrix H_k , and every row in Z_k is set to z_k . Then in the training, the calculation of hidden layer H_{k+1} under dropout regularization is

$$H_{k+1} = f [(H_k \circ Z_k) W_{k+1} + B_{k+1}]$$

where $^{\circ}$ is the element-wise product. Note that with a trained autoencoder, the calculation of hidden layers in the imputation step does not involve the dropout operation. In our analysis, we set $p_1 = p_2 = ... = p_h = p$, where *h* is the total number of hidden layers. We call *p* the dropout rate in the following text.

4.3. Three masking schemes for introducing artificial zeros

In scRNA-seq data, distinguishing biological zeros and nonbiological zeros (i.e., missing values) is challenging without external information [17,18]. Hence, to evaluate the overall imputation accuracy, we design three masking schemes to introduce artificial zeros into scRNA-seq data and measure the differences between the artificial zeros' imputed values and original values (Fig. 1b-d). The three masking schemes represent three typical assumptions of missing mechanisms in scRNA-seq data [9].

Random masking: we randomly mask (by setting values to zeros) 50% of nonzero entries in the scRNA-seq data matrix. Random masking means that the missing mechanism is independent of the gene expression levels, and it has been widely used in previous work to evaluate the imputation accuracy [18].

Median masking: we mask the nonzero entries less than or equal to the overall median in the scRNA-seq data matrix. Median masking assumes a complete dependence of the missing mechanism on the gene expression levels.

Double exponential masking: we assume that a gene's probability of having missing values depends on the gene's mean expression level across all cells. Hence, lowly expressed genes are more likely to have missing values than highly expressed genes. Specifically, for gene *j*, let \overline{X}_j be the mean expression level (natural-log-plus-one-transformed read count) of nonzero values across all cells and p_j be the probability of missing values. Then gene *j*'s probability of having missing values is modeled by a double exponential function [9].

$$p_j = \exp \Big(-\lambda \overline{X}_j^2 \Big).$$

Let Z_{ii} be a random variable that indicates whether to mask the

nonzero expression of gene *j* in cell *i*. Then $Z_{ij} \sim Bernoulli(p_j)$. If $Z_{ij} = 1$, then the nonzero expression of gene *j* in cell *i* will be masked. The value of λ is determined such that 50% nonzero entries in the scRNA-seq data matrix are masked.

A previous study showed that artificial zeros introduced by the three masking schemes exhibited different impacts on scRNA-seq data analysis [9]. Although random masking has been widely used to evaluate imputation accuracy, its assumption is unrealistic. In contrast, median masking is an extreme way of modeling the observation that lowly expressed genes have more zeros than highly expressed genes have [15, 55]. Between the two masking schemes, double exponential masking uses a probabilistic model to reflect this observation at the individual gene level. Together, the three masking schemes provide a comprehensive evaluation of imputation accuracy from different perspectives.

4.4. Data preprocessing and normalization

All real and synthetic scRNA-seq datasets used in this study are cellby-gene count matrices. They are preprocessed and normalized in three steps. First, we remove the genes expressed in fewer than three cells and the cells having fewer than 200 genes expressed. Second, we divide each count by its cell library size (i.e., the cell's total count) and then multiply it by 10000 (library size normalization). Then we add one to the normalized count and apply the natural-log transformation. Third, we select 2000 highly variable genes using the vst method implemented in the FindVariableFeatures function of the Seurat package [34] (v4.0). After preprocessing and normalization, the dimensions of all scRNA-seq data matrices are cell number \times 2000. Note that the preprocessing and normalization steps apply to the pre-imputed datasets only. The imputed datasets, when used as the input of cell clustering and DE gene analysis, will not go through the preprocessing and normalization steps.

4.5. Training of autoencoders and imputation

The training of autoencoders is implemented using the Pytorch deep learning library [56] (v 1.8.1) on a server with two Intel Xeon E5-2687 W v4 CPUs, 256 GB memory, an Nvidia GeForce RTX 2080 Ti GPU, and Ubuntu 18.04 system. After preprocessing, normalization, and masking (masking is only necessary for the evaluation of overall imputation accuracy), we split each dataset's cells into a training set (80% of cells) and a validation set (20% of cells). We utilize the Adam optimization algorithm [21] to train the autoencoder with a 0.001 learning rate and a 64 batch size. After every epoch of training on the training set, we impute the validation set using the current autoencoder and calculate the MSE between the original nonzero values in the validation set and their corresponding imputed values. We continue the training until either the MSE does not decrease over 20 epochs or the total number of epochs surpasses 10000. In the imputation step, the trained autoencoder accepts the preprocessed and normalized scRNA-seq data matrix as input (with the dimensions as cell number \times 2000) and outputs a data matrix of the same dimensions. The final imputed data matrix is constructed by replacing the zero values in the input matrix with their counterparts in the output matrix. The nonzero values in the input matrix remain in the final imputed data matrix.

4.6. Calculation of imputation normalized root mean squared error (NRMSE)

Denote by X the sparse scRNA-seq input matrix after preprocessing and normalization; \hat{X} is the imputed scRNA-seq data matrix; M is the set of masked entries in the scRNA-seq data matrix. The MSE between the imputed values and the original values of the masked entries, MSE_{mask}, is calculated as

$$MSE_{mask} = \frac{\sum_{i}^{n} \sum_{j}^{m} (X_{ij} - \widehat{X}_{ij})^{2} I(X_{ij} \in M)}{\sum_{i}^{n} \sum_{j}^{m} I(X_{ij} \in M)}.$$

Denote the mean masked values \overline{X}_{mask} as

$$\overline{X}_{\text{mask}} = \frac{\sum_{i} \sum_{j} X_{ij} I(X_{ij} \in M)}{\sum_{i} \sum_{j} \sum_{i} I(X_{ij} \in M)}$$

Then the imputation NRMSE is calculated as NRMSE_{imputation} = $\frac{\sqrt{MSE_{mask}}}{\overline{X}_{mask}}$.

4.7. Activation functions

We evaluate seven activation functions in this study, including logistic function (sigmoid), hyperbolic tangent function (tanh), rectified linear unit (ReLU), leaky ReLU (with two hyperparameters), exponential linear units (ELU), and scaled exponential linear units (SELU). An activation function accepts a linear combination of the outputs from the last layer as input and applies a nonlinear transformation to the linear combination. All activation functions take a real-valued scalar as input and output a real-valued scalar.

The sigmoid activation function is a bounded differentiable function with positive and continuous derivatives. Its value ranges from 0 to 1. The function form of sigmoid is

$$f(x) = \frac{1}{1 + e^{-x}}.$$

The tanh activation function is also a bounded differentiable function with positive and continuous derivatives. Its value ranges from -1 to 1. The functional form of tanh is

$$f(x) = \frac{e^x - e^{-x}}{e^x + e^{-x}}.$$

The ReLU activation function conducts a threshold operation that outputs zero for a negative input and acts as an identity function for a positive input. Its value ranges from 0 to $+\infty$, and it has discrete derivatives. The functional form of ReLU is

$$f(x) = \begin{cases} x & \text{if } x \ge 0\\ 0 & \text{if } x < 0 \end{cases}$$

The leaky ReLU activation function modifies ReLU by introducing a small negative slope when the input is negative. Its value ranges from $-\infty$ to $+\infty$, and it has discrete derivatives. The functional form of leaky ReLU is

$$f(x) = \begin{cases} x & \text{if } x \ge 0\\ ax & \text{if } x < 0 \end{cases}$$

where $\alpha > 0$ is a hyperparameter. In our analysis, we set α to 0.01 and 0.2 — the default values in two popular deep learning libraries Pytorch [56] and TensorFlow [57].

The ELU activation function replaces the linear negative part of leaky ReLU with an exponential function. Its value ranges from $-\infty$ to $+\infty$, and it has discrete derivatives. The functional form of ELU is

$$f(x) = \begin{cases} x & \text{if } x \ge 0\\ a e^x - 1 & \text{if } x < 0 \end{cases},$$

where $\alpha > 0$ is a hyperparameter. In our analysis, we set α to 1 — the default value in Pytorch.

The SELU activation function further adds a scale factor to ELU and changes the constant in the negative part. Its value ranges from $-\infty$ to $+\infty$, and it has discrete derivatives. The functional form of ELU is

$$f(x) = \tau \{ \begin{array}{cc} x & \text{if } x \ge 0\\ ae^x - a & \text{if } x < 0 \end{array} \}$$

where $\tau > 0$ and $\alpha > 0$ are predefined parameters with $\tau = 1.05$ and $\alpha = 1.67$.

4.8. Weight decay and dropout parameter settings

When evaluating the impact of weight decay regularization on the imputation accuracy, we set the hyperparameter λ to nine values, including 0, 1e-7, 5e-7, 1e-6, 5e-6, 1e-5, 5e-5, 1e-4, and 5e-4. Larger λ values indicate stronger regularization imposed on the autoencoder, and $\lambda = 0$ means no regularization. When evaluating the impact of weight decay regularization on the cell clustering and DE gene analysis, we set the hyperparameter λ to 17 values, including 0, 1e-7, 5e-7, 1e-6, 5e-6, 1e-5, 5e-5, 1e-4, 5e-4, 5e-3, 0.01, 0.05, 0.1, 0.5, 1, and 5. When evaluating the impact of dropout regularization on the imputation accuracy, cell clustering, and DE gene analysis, we set the dropout rate *p* to 11 values, including 0, 0.01, 0.02, 0.05, 0.1, 0.15, 0.2, 0.25, 0.3, 0.35, and 0.4. Larger *p* values indicate stronger regularization imposed on the autoencoder, and *p* = 0 means no regularization.

4.9. Cell clustering analysis

We use the function kmeans in R programming language to conduct K-means clustering on the pre-imputed and imputed scRNA-seq datasets (Supplementary Table S2). We set the parameter centers (the number of clusters k in the K-means clustering) to the correct number of cell types in each dataset. We set the parameter nstart to 25, which repeats the clustering 25 times by randomly selecting 25 sets of initial cluster centers and returns the result with the minimum sum of pairwise distances within clusters [58]. The dimensions of input data matrices for K-means clustering is cell number \times 2000 without further dimension reduction (that is, each cell is a 2000-dimensional vector). Note that before clustering, pre-imputed datasets are preprocessed by following the procedure described in the section "Data preprocessing and normalization." The imputed datasets are directly clustered by K-means clustering.

We use adjusted Rand index (ARI) and adjusted mutual information (AMI) to measure the performance of cell clustering. Let $U = \{u_1, u_2, ..., u_c\}$ be the true partition of *c* classes and $V = \{v_1, v_2, ..., v_c\}$ be the partition obtained by K-means clustering. Let n_i and m_j be the numbers of observations in class u_i and cluster v_j , respectively. Let r_{ij} be the number of observations in both class u_i and cluster v_j . The ARI is calculated as

$$\frac{\sum_{i=1}^{c} \sum_{j=1}^{c} {\binom{r_{ij}}{2}} - \left[\sum_{i=1}^{c} {\binom{n_i}{2}} \sum_{j=1}^{c} {\binom{m_j}{2}}\right] / {\binom{n}{2}}}{\left[\sum_{i=1}^{c} {\binom{n_i}{2}} + \sum_{j=1}^{c} {\binom{m_j}{2}}\right] / 2 - \left[\sum_{i=1}^{c} {\binom{n_j}{2}} \sum_{j=1}^{c} {\binom{m_j}{2}}\right] / {\binom{n}{2}}},$$

where *n* is the total number of observations and $n = \sum_{i=1}^{c} n_i = \sum_{j=1}^{c} m_j$. The AMI is calculated as

$$\frac{2I(U,V)}{H(U) + H(V)}$$

where I(U, V) is the mutual information of U and V, and H(U) and H(V) are the entropies of U and V respectively [59]. We utilize the functions ARI and AMI in R package aricode to calculate ARI and AMI, respectively.

4.10. Simulation of synthetic scRNA-seq data

We utilize simulator scDesign [19] to generate 20 synthetic scRNA-seq data with ground-truth DE genes. The 20 real datasets for training scDesign (Supplementary Table S3) are preprocessed by following the procedure described in the section "Data preprocessing and normalization." For each real dataset, we execute the function design_data in R package scDesign to simulate one synthetic dataset. Each synthetic dataset contains two cell types with 1000 cells per type, and 10% of genes are differentially expressed between the two cell types. The sequencing depth of each synthetic dataset is equal to the

median cell library size of the corresponding real dataset multiplied by the cell number (2000). Other parameters of the function design_data are set to their default values. All synthetic datasets are count matrices with dimensions as cell number \times 2000.

4.11. DE gene analysis

We conduct DE gene analysis on the aforementioned 20 synthetic datasets and their imputed counterparts. For pre-imputed synthetic datasets, the gene expression counts of each cell are divided by the total counts of that cell (library size) and then multiplied by 10000 (library size normalization). The results are then added by one before being natural-log transformed. We utilize the function FindMarkers in R package Seurat to identify the DE genes between the two cell types. We set the parameter test.use to "MAST" and identify genes with Bonferroni-corrected p-values under 0.05 as DE genes. Based on the ground-truth DE genes, we calculate the precision, recall, and TNR for each pre-imputed synthetic dataset and the corresponding imputed dataset.

4.12. Sensitivity analysis with varying numbers of highly variable genes

We perform a sensitivity analysis by varying the number of highly variable genes to assess the robustness of our major findings. Specifically, we examine the impacts of autoencoder architecture, activation function, and regularization on imputation accuracy using random masking. Two scenarios are considered, where the highly variable genes are set to 1000 and 3000, respectively. The results of this analysis are summarized in Supplementary Figs. S12–S19. Notably, our investigation confirms that our major findings remain consistent across the three different numbers of highly variable genes (1000, 2000, and 3000).

Declaration of Competing Interest

The authors declare no conflict of interests.

Data and code availability

The source code is available at the GitHub repository: https://github. com/xnnba1984/Benchmarking-the-Autoencoder-Design-for-Imputing-Single-Cell-RNA-Sequencing-Data. The datasets used in this study can be found at the Zenodo repository: https://zenodo.org/record/7504311#. Y7ei0naZMuK.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.csbj.2023.07.041.

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