# Improvements for Seurat

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#### 1 Introduction

As an extension for seurat package, honfleuR has the following chases:

- Workspace before I submit pull request, and hopefully it will be merged into seurat package.
- Incorporate new algorithms to expand its capability of learning data.
- Design and enrich functions and parameters to make seurat compatible with more research of interests.
- Improve running speed while following seurat syntax and reproducing results.

## 2 Summary of changes

| Category                                                       | Seurat          | honfleuR          | What's New                                        |
|----------------------------------------------------------------|-----------------|-------------------|---------------------------------------------------|
| Localization Localization Localization Localization Clustering | addImputedScore | fill_imputed_expr | interface for additional imputation strategies    |
|                                                                | fit.gene.k      | fit_gene_k        | 10X faster; strict control of biological meanings |
|                                                                | initial.mapping | initial_mapping   | 1X faster                                         |
|                                                                | refined.mapping | refined_mapping   | 17X faster                                        |
|                                                                | jackStraw       | jackStraw2        | debug                                             |

eval\_seurat function evaluates Seurat performances on landmark genes and draws ROC curves, reproducing Fig3-G&H.

# 3 Data imputation - fill\_imputed\_expr

Impute expression of each landmak gene (response) based on other genes with variable expressions (predictors).

honfleuR has three imputation strategies:

- 1. Lasso. Reproduce the results of seurat.
- 2. PLSR. Account for potential linear dependencies among predictors.
- 3. Tilling lasso. Learn the data structure and perform imputation.

| Step | Lasso                |   | PLSR            |   | Tilling Lasso                         |
|------|----------------------|---|-----------------|---|---------------------------------------|
| 1    | Focus on landmark ge | 1 | Focus<br>landma | 1 | Focus on specific landmark gene $G$ . |

| Step | Lasso                                                                                                                 | PLSR                                                                                                                | Tilling Lasso                                                                                                                      |
|------|-----------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------|
| 2    | Given the matrix with cells ID on rows and genes on columns, train linear regression model with lasso regularization. | Given the matrix with cells ID on rows and genes on columns, train a PLSR (Partial Least Squares Regression) model. | Given the data matrix with cells ID on rows and genes on columns, shuffle rows randomly.                                           |
| 3    |                                                                                                                       |                                                                                                                     | Set first 20% samples as "unseen" data.                                                                                            |
| 4    |                                                                                                                       |                                                                                                                     | Use the rest 80% samples as training dataset to train a lasso model.                                                               |
| 5    |                                                                                                                       |                                                                                                                     | Apply the model on samples selected on<br>Step 2, impute landmark gene expression<br>in these cells.                               |
| 6    |                                                                                                                       |                                                                                                                     | Set second 20% samples as "unseen" data.<br>Repeat Step 3-5 until the expression of land-<br>mark gene is imputed among all cells. |
| 7    | Apply lasso model on same matrix to impute expression of gene $G$ .                                                   | Apply PLSR model on same matrix to impute expression of gene $G$ .                                                  | Repeat Step 2-6 for 10 times. The average values are the imputed expressions for gene $G$ among all cells.                         |
| 8    | Iterate Step 1-7 for other landmark genes.                                                                            | Iterate Step 1-7 for other landmark genes.                                                                          | Iterate Step 1-7 for other landmark genes.                                                                                         |

#### 3.1 Use Lasso

Run the following codes to test whether honfleuR reproduces results.

The result is (expected to be) TRUE.

#### 3.2 Use PLSR.

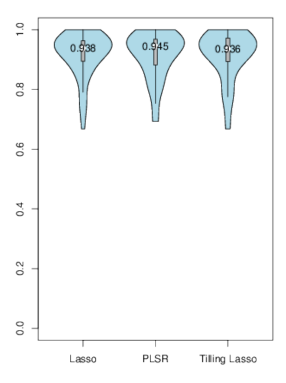
### 3.3 Use Tilling-lasso

#### 3.4 Quick summary

- 1. honfleuR expands seurat capability in addition to linear regression.
- 2. The interface of honfleuR is unified, i.e. setting parameterscheme = c('lasso', 'plsr', 'tlasso').
- 3. honfleuR follows the seurat syntax and does not alter seurat's codes frame.

## 3.5 ROC analysis for imputation scheems

As the way of performing ROC analysis as in Figure 3H of manuscript, the following AUC results are given by using honfleuR's eval\_seurat function.



## 4 Estimate bimodal distribution of landmark gene - fit\_gene\_k

First, honfleuR fixes a bug related with biology. There is a biological issue that original fit.gene.k omits. There are following two scenarios:

```
fit.gene.k(zf, "SOX3", do.k = 3)
```

do.k = 3 means it is assumed that the in situ pattern of gene G has 3 expression levels: low, med, high. In tutorial it was default as 2.

```
fit.gene.k(zf, "SOX3", start.pct=mean(zf@insitu.matrix[,"SOX3"]))
```

start.pct sets the initial percentage of cells in "on" state therefore the dataset is expected to be binary if start.pct is in action.

The above two calls are appropriate. However, current implementation of seurat allows the following extreme case taking place legally:

```
fit.gene.k(zf, "SOX3", do.k = 5, start.pct=mean(zf@insitu.matrix[, "SOX3"]))
```

It is conflict that five (any number greater than 2) different expression levels and "on/off" presumption coexists. Therefore honfleuR comes up with a patch, see fit\_gene\_k.

Furthermore, fit\_gene\_k is 10X faster than fit.gene.k. Run the following codes to see the efficiency boost (on my laptop decrease from 24s down to 2s).

The result is TRUE meaning that honfleuR builds the model more efficiently without losing correctness. See results here estimated by using microbenchmark package.

# 5 Cells mapping - initial\_mapping and refined\_mapping

initial\_mapping and refined\_mapping exactly follows twins functions initial.mapping and refined.mapping respectively.

initial\_mapping is 1X faster (11s down to 5s), and refined\_mapping is 17X times faster (98s down to 5s). See boosting results here estimated by using microbenchmark package.