

# Single-Cell Analysis of Regeneration-Organizing Cells in Xenopus Tail

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## Abstract

The regeneration-organizing cell (ROC) displays a unique gene expression profile and plays a pivotal role in the regeneration of *Xenopus* tail. In this study, we apply single-cell analyses to systematically investigate the unique gene expression profiles of this newly identified regeneration-organizing cell. Clustering and visualization reveal distinct cellular populations, with the ROC cluster clearly separated, confirming its transcriptional uniqueness. Clustering quality is further validated through multiple evaluation metrics. Using logistic regression and the Wilcoxon rank-sum test, we identify marker genes in ROCs. *Apoc1.like.L* is identified as a key marker by both methods, but it is highly expressed across multiple cell types, whereas *pltp.S* serves as a specific marker for ROCs. We apply multiple denoising and batch integration techniques to improve the robustness of our findings. Notably, MAGIC and Harmony correction significantly improves clustering metrics and the former has a noticeable impact on the top 50 expressed genes, while the previous distinctive ROC marker genes can still be reliably identified, further confirming the robustness of our analysis pipeline. Code is available at [https://github.com/DanJiayi/STAT5243\\_Project1](https://github.com/DanJiayi/STAT5243_Project1).

**Keywords:** Regeneration-organizing cell, Single-cell analyses, *Xenopus* tail regeneration

## 1 Introduction

*Xenopus laevis* tadpoles exhibit remarkable regenerative capacity, particularly in their tails. Recent studies have identified a novel epidermal cell type, the regeneration-organizing cell (ROC), which displays a unique gene expression profile and plays a central role in orchestrating tissue regeneration. Previous work [1] using single-cell transcriptomics has identified the existence of ROCs and provided a mechanistic understanding of the initiation and organization of tail regeneration. However, systematic pipelines for single-cell analysis of ROCs have not yet been fully established. In this study, we aim to develop a ROC-focused single-cell analysis framework that integrates clustering with gene expression profiling, and further evaluate the reliability of our findings through extensive denoising and batch integration techniques.

## 2 Methods

**Data preprocessing.** The data with use is consistent with [1]. Raw count matrices were normalized to the same total counts per cell, and log-transformed to stabilize variance. Genes expressed in fewer than 3 cells and cells with fewer than 200 detected genes were removed. Highly variable genes were then identified (top 2,300 genes ranked by dispersion), and used for downstream analyses.

**Clustering Analysis.** To perform clustering, we first reduced the dimensionality of the data using principal component analysis (PCA). Based on the top 30 PCs, we constructed a k-nearest neighbor graph with  $k = 15$ . Using this graph, we performed community detection with both the Louvain and Leiden algorithms, generating alternative cluster partitions of the data. Parameters was adjusted to yield a number of clusters consistent with the 46 known clusters in the dataset, and the clustering results was visualized with UMAP. We evaluated the clustering quality using multiple metrics: Adjusted Rand Index (ARI), Rand Index (RI), Silhouette Score, and Calinski–Harabasz Score.

**Marker selection and Gene Ontology analysis.** We applied differential expression analysis to identify marker genes for ROC. Two complementary statistical approaches were employed: logistic regression and the Wilcoxon rank-sum test. We extracted the top 50 marker genes ranked by each method, annotated the clusters with key markers to confirm their close association with ROC, and compared their expression between ROC and other clusters to select ROC-specific markers. Functional enrichment of the selected marker genes was performed using g:Profiler.

**Data denoising.** To mitigate technical noise and enhance the biological signal, we applied two complementary denoising approaches. First, we used

scVI, a variational autoencoder-based framework, to learn a latent representation of the data and generate normalized denoised expression values (library size set to 1e4, trained for 100 epochs). Second, we applied MAGIC, a diffusion-based imputation method, to recover gene–gene relationships and smooth expression profiles. Both denoised datasets were saved for downstream analyses. After denoising, we re-performed clustering and marker selection, and compared the results with the original analysis to validate the improvement in clustering performance and the robustness of the identified marker genes. Parameters were tuned to maintain a stable number of clusters; the same applies below.

**Batch integration.** For batch effect correction, we used two complementary approaches. Harmony integrates single-cell data by iteratively adjusting the principal component embeddings to remove batch-specific variation while preserving biological structure. Specifically, Harmony was applied using batch labels as covariates, yielding corrected embeddings. As an alternative, we applied BBKNN (Batch Balanced KNN), which modifies the neighborhood graph construction to ensure balanced representation of cells from each batch. Similarly, We re-performed clustering and tested their effectiveness by comparing with the previous results.

**Code availability.** Code supporting this study is available at [https://github.com/DanJiayi/STAT5243\\_Project1](https://github.com/DanJiayi/STAT5243_Project1).

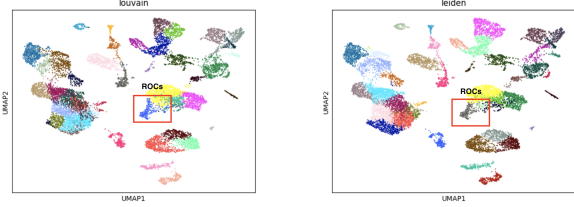
## 3 Results

### 3.1 Clustering Analysis and Visualization

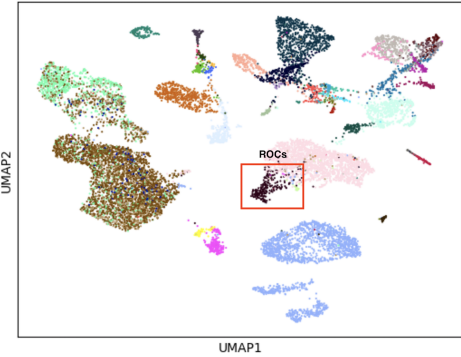
#### 3.1.1 Visualization of clustering results

We applied both the Louvain and Leiden algorithms after PCA-based dimensionality reduction (all clustering in this study was tuned to match the number of clusters in the reference (46)). The UMAP visualization (Fig. 1) shows that both methods separated the cells into distinct clusters. The strong separation among clusters suggests that cell types are characterized by distinct gene expression signatures, with certain lineage-specific markers driving the divergence. For example, immune-related clusters were well distinguished from epithelial and mesenchymal populations, reflecting the underlying functional heterogeneity.

The concordance with the reference cluster annotations (Fig. 2) further supports that these clustering approaches effectively recover biologically relevant cellular identities. Notably, both clustering methods successfully distinguished ROCs (highlighted by the red box in the figure), underscoring the unique gene expression characteristics of these cells.



**Fig. 1:** Clustering results.



**Fig. 2:** Reference cluster annotations.

#### 3.1.2 Clustering evaluation metrics

To quantitatively evaluate clustering performance, we compared the Louvain and Leiden algorithms using four common metrics: Adjusted Rand Index (ARI), Rand Index, Silhouette score, and Calinski–Harabasz score. As shown in Table 1, both methods achieved comparable results. Louvain and Leiden yielded similar ARI values and high Rand Index values (~0.915), indicating a reasonable consistency with the reference labels. Leiden slightly outperformed Louvain in ARI and Silhouette score, suggesting that Leiden provides more coherent clusters. Louvain obtained a slightly higher CH score, reflecting that it may produce clusters with more distinct separation in terms of variance-based criteria. Overall, the results indicate comparable clustering performance between the two methods.

**Table 1:** Clustering metrics.

Method	ARI	Rand Index	Silhouette	CH Score
Louvain	0.3056	0.9140	0.1977	1880
Leiden	0.3230	0.9151	0.2056	1867

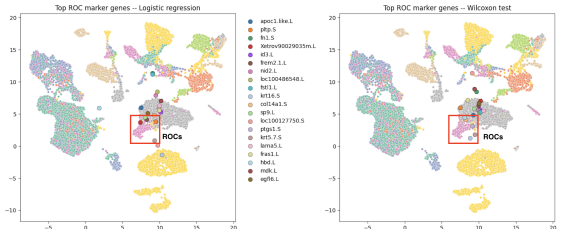
### 3.2 Marker Selection and Gene Analysis

#### 3.2.1 Marker Selection

We applied two complementary strategies: logistic regression and Wilcoxon rank-sum test, to identify marker genes of the ROC cluster. From the top 50 markers identified by each method, we observed a substantial overlap of 29 genes,

highlighting a consistent marker signature across statistical approaches. When compared with the reference list provided in Supplementary Table 3 of [1], 8 of the logistic regression-derived markers and 5 of the Wilcoxon-derived markers were retained.

We annotated the top 20 ROC markers identified by both methods on the UMAP plot (by computing a weighted average of UMAP coordinates based on the expression of these genes). As shown in ,the key markers identified by the two methods were both closely localized to the ROC cluster, confirming the reliability of our results.

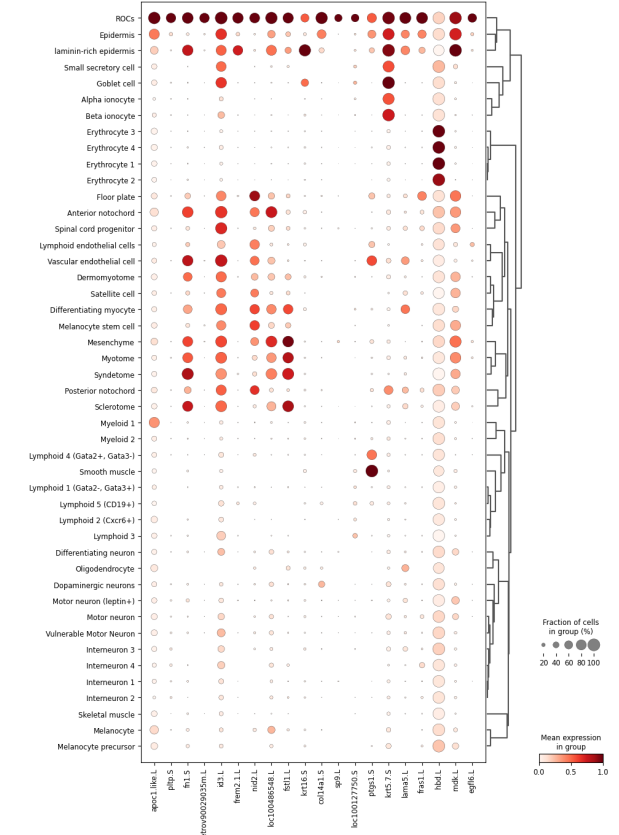


**Fig. 3:** Top ROC markers (annotated on UMAP).

Using logistic-regression-based markers as an illustrative case, the top five ROC marker genes were apoc1.like.L, pltp.S, fn1.S, Xetrov90029035m.L, and id3.L (apoc1.like.L consistently ranked in the top two across both methods). As shown in Fig. 4, apoc1.like.L, fn1.S, and id3.L also showed high expression in the Epidermis or other cell types. In contrast, pltp.S and Xetrov90029035m.L were specifically expressed in the ROC cluster. Both of these ROC-specific genes are consistent with the findings reported in Supplementary Table 3 of [1], with pltp.S also being independently identified by the Wilcoxon-based marker selection. These results highlight pltp.S as specific markers of ROCs.

### 3.2.2 Gene Ontology analysis

GO enrichment analysis revealed that the logistic-regression-based marker genes were strongly associated with extracellular matrix (ECM) organization. Among the most significant terms, “basement membrane” was enriched with 8 marker genes, highlighting a strong link between ROCs and basement membrane formation. Given the basement membrane’s critical role in maintaining tissue architecture, providing structural support, and mediating cell adhesion, this finding suggests that ROCs may contribute specifically to ECM remodeling at epithelial or endothelial interfaces. Other enriched terms, such as “extracellular matrix,” “extracellular region,” and “collagen-containing extracellular matrix,” further reinforce this ECM-associated signature.



**Fig. 4:** Expression of top ROC marker genes.

**Table 2:** Top 10 significantly enriched GO terms from ROC marker genes.

Name	p-value	Intersection
Basement membrane	$4.30 \times 10^{-10}$	8
Extracellular region	$1.67 \times 10^{-8}$	25
Extracellular matrix	$1.79 \times 10^{-7}$	11
External encapsulating structure	$1.83 \times 10^{-7}$	11
Extracellular matrix structural constituent	$1.86 \times 10^{-7}$	8
Extracellular space	$5.58 \times 10^{-7}$	21
Collagen-containing extracellular matrix	$2.36 \times 10^{-6}$	9
Structural molecule activity	$1.06 \times 10^{-5}$	13
Anatomical structure morphogenesis	$1.13 \times 10^{-5}$	19
Cell motility	$1.01 \times 10^{-4}$	15

### 3.3 Data denoising

We assessed the impact of scVI and MAGIC denoising on clustering performance (parameters were

adjusted to maintain a stable number of clusters). As shown in Table 3 and Table 4, while both methods maintained ARI and Rand Index values close to original results, MAGIC substantially increased the Silhouette and Calinski–Harabasz scores. This suggests that MAGIC improves spatial separability and enhance the effectiveness of gene expression signals for downstream analyses, yet it does not guarantee a change in the clustering assignments.

**Table 3:** Clustering metrics - scVI

Method	ARI	Rand Index	Silhouette	CH Score
Louvain	0.3067	0.9152	0.2333	2959
Leiden	0.2594	0.9129	0.2479	2879

**Table 4:** Clustering metrics - MAGIC

Method	ARI	Rand Index	Silhouette	CH Score
Louvain	0.3077	0.9141	0.3801	5076
Leiden	0.2454	0.9110	0.3681	5147

After denoising, we re-identified the marker genes for ROCs. Taking logistic regression-based markers as an example, only 31 and 33 of the top 50 expressed genes overlapped with the previous results after each denoising method (when using the Wilcoxon test, the impact is larger, and the overlap with the reference [1] increases to 19), while the previously top marker genes—apoc1.like.L, pltp.S, and fn1.S—still can be identified. Notably, pltp.S continued to be a ROC-specific marker. These findings confirm the robustness of our earlier conclusions and demonstrate that denoising has a substantial impact on marker identification, highlighting its necessity.

### 3.4 Batch integration

We further adopted two batch correction strategies, Harmony and BBKNN. As shown in Table 5 and Table 6, Harmony correction noticeably improved ARI, Rand index, and Calinski–Harabasz score in both clustering methods, suggesting that it mitigates batch effects and stabilizes gene expression profiles, thereby enhancing clustering performance. However, BBKNN did not yield significant improvements in this task.

**Table 5:** Clustering metrics - Harmony.

Method	ARI	Rand Index	Silhouette	CH Score
Louvain	0.3195	0.9156	0.2005	2030
Leiden	0.3381	0.9170	0.2013	1964

**Table 6:** Clustering metrics - BBKNN.

Method	ARI	Rand Index	Silhouette	CH Score
Louvain	0.2731	0.9112	0.1896	1868
Leiden	0.2939	0.9136	0.2124	1974

Since batch integration methods such as Harmony and BBKNN operate on the latent embedding or neighborhood graph rather than altering the original expression matrix, the identification of marker genes remains unchanged.

## 4 Conclusion

In this work, we applied single-cell analyses to systematically investigate the unique gene expression profiles of regeneration-organizing cells (ROCs). Multiple clustering methods demonstrate that ROC possesses a distinct transcriptional profile, and various evaluation metrics confirm the robustness of our clustering results. Using logistic regression and Wilcoxon test, We identified the top 50 ROC marker genes, mapped them onto the clusters, and profiled their differential expression. The results of the two methods were highly consistent, revealing that genes such as apoc1.like. and pltp.S serve as key markers of ROCs. GO enrichment analysis further indicated that these highly expressed genes are strongly associated with extracellular matrix organization. Finally, Denoising with MAGIC and batch integration with Harmony both significantly improved multiple clustering metrics. While denoising notably affected the top 50 genes, the previously identified key ROC marker genes remained detectable, confirming the reliability and reproducibility of our pipeline and findings.

## References

- [1] Can Aztekin, TW Hiscock, JC Marioni, JB Gurdon, BD Simons, and Jerome Jullien. Identification of a regeneration-organizing cell in the *xenopus* tail. *Science*, 364(6441):653–658, 2019.