

實作作業說明

報告時間：(待定)

報告地點：線上會議

本實作請下載「線上公開的皮膚相關 scRNA-seq 資料」進行完整分析。

課程設計 9+1 項 *Checkpoints*，其中 **第 0, 5-8 項為必答**，其餘可視需求選答。請於分析與撰寫報告時，依序回應下列問題並提出具體佐證（圖表、統計量或文獻）：

| # | 檢核主題 | 檢查點與關鍵問題 |
|---|-------------------|--|
| 0 | 資料來源與預處理記錄 | · 請詳細紀錄資料來源與樣本說明、分析流程、R 及套件版本與參數設定。 |
| 1 | 品質管制 (QC) 設定 | · 閾值與過濾策略是否合理？ · 是否可能過度過濾而排除關鍵細胞？ |
| 2 | 主成分數量 (PCs) 選擇 | · 依據哪些指標或概念決定 PC 數？ |
| 3 | 批次效應校正 | · 整合後是否改善 batch effect 且保留生物訊號？ · 若為跨平台／實驗室／物種整合，需注意哪些問題？ |
| 4 | 群集解析度調整 | · 最終採用的解析度是否合理？如何判斷過高或過低？ · 是否使用量化指標進行解析度優化？ |
| 5 | 細胞類型標註 (必答) | · 標註結果是否符合已知生物學知識？ · 是否有值得關注的細胞族群變化或新穎細胞族群？ |
| 6 | 差異表達基因 (DEG) (必答) | · 目前使用的篩選門檻與統計方法是否恰當？為什麼？ · 這些 DEG 是否與既有的疾病機轉、治療靶點、或細胞功能相關？是否提供新見解？ |
| 7 | 功能富集與細胞通訊 (必答) | · 富集分析與通訊網絡揭示何種機制假設？ · 與臨床或病理的連結為何？ |
| 8 | 結果整合 (必答) | · 各分析結果間是否具有的一致性？是否可整合為一個生物機制假說或模型？ |
| 9 | 軌跡分析 (若適用) | · 如何界定起點 (root)？ · 起點選擇對軌跡推論結果造成何種影響？ |

提交格式

- 書面報告：Word 或 PDF，附完整程式碼、主要圖表與關鍵結果說明
- 口頭報告：10 分鐘簡報 + 5 分鐘 Q&A

Practical Assignment Instructions

Oral Presentation

Date: TBD

Venue: Online Meeting

Assignment Description:

Participants are required to download a **publicly available skin-related scRNA-seq dataset** and conduct a complete analysis.

The assignment includes **9+1 checkpoints**, where **Checkpoints 0 and 5–8 are mandatory**, and the rest are optional based on your analysis needs.

Please respond to the following checkpoints in order, with **supporting evidence** such as plots, statistical metrics, or references.

| # | Topic | Checkpoints & Key Questions |
|---|---|---|
| 0 | Data Source and Preprocessing | <ul style="list-style-type: none">Record dataset source, sample information, analysis workflow, R and package versions, and parameter settings in detail. |
| 1 | Quality Control (QC) Settings | <ul style="list-style-type: none">Are the filtering thresholds and strategies reasonable?Could essential cell populations be excluded due to over-filtering? |
| 2 | Selection of Principal Components (PCs) | <ul style="list-style-type: none">What indicators or concepts were used to determine the number of PCs? |
| 3 | Batch Effect Correction | <ul style="list-style-type: none">Does the integration effectively reduce batch effects while preserving biological signals?For cross-platform/lab/species integration, what issues arise? |
| 4 | Clustering Resolution Adjustment | <ul style="list-style-type: none">Is the chosen resolution appropriate? How do you determine if it is too high or low?Did you use quantitative metrics to optimize resolution? |
| 5 | Cell Type Annotation (Required) | <ul style="list-style-type: none">Do the annotations match known biological knowledge?Are there notable population shifts or novel cell types? |

| # | Topic | Checkpoints & Key Questions |
|---|--|---|
| 6 | Differential Gene Expression (Required) | <ul style="list-style-type: none"> • Are the current thresholds and statistical methods appropriate? Why? • Do the DEGs relate to disease mechanisms, therapeutic targets, or cell functions? |
| 7 | Functional Enrichment & Cell-Cell Communication (Required) | <ul style="list-style-type: none"> • What mechanistic hypotheses do enrichment and communication analyses suggest? • How do these findings connect to clinical or pathological features? |
| 8 | Integration of Results (Required) | <ul style="list-style-type: none"> • Are the findings consistent across analyses? • Can the results be synthesized into a coherent biological model or hypothesis? |
| 9 | Trajectory Analysis (if applicable) | <ul style="list-style-type: none"> • How is the root state defined? • How does the root selection affect the inferred trajectory? |

Submission Format

- **Written Report:**

Submit as a Word or PDF file, including the full analysis code, key plots, and explanations of major findings.

- **Oral Presentation:**

10-minute presentation followed by 5 minutes of Q&A.