#### RUV-III-NB: A robust scRNA-seq normalization methods

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MSPGH and School of Mathematics and Statistics
The University of Melbourne
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# Single-Cell Sequencing

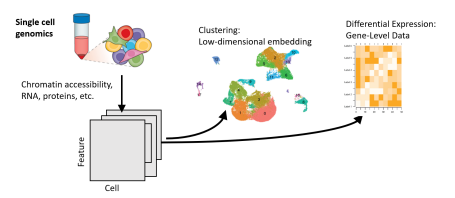


Figure adapted from Longo, Guo, Ji and Khavari (2021, Nat. Rev. Genetics)

Clustering is used to identify cell states; DE is used to identify marker genes that differentiate states

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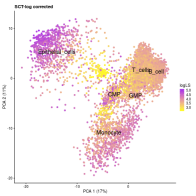
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- Motivation I: Current normalization methods remove biology when unwanted variation (UV) are associated with biology.
- Motivation II: Most methods only return dimensional reduction (cell embedding) unsuitable for downstream analyses.
- ullet RUV-III-NB takes into account biology imes UV association and return adjusted data for all genes.

## **NSCLC Study**

Non-small cell lung carcinoma ( $\sim$  6,000 cells) study using 10x platform (from one batch)

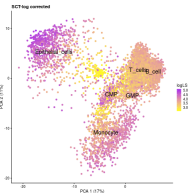
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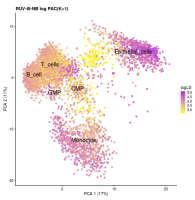


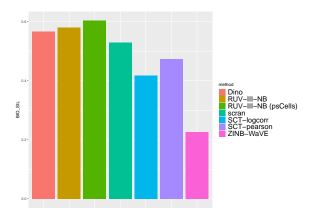
 Biology (cell-type) is associated with library size (UV), with the larger Epithelial cells and Monocytes have higher LS.

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 RUV-III-NB separates Monocytes better and makes Epithelial cells cluster tighter.



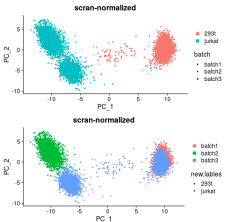


• Only RUV-III-NB and Dino improve biological silhouette score relative to scran.

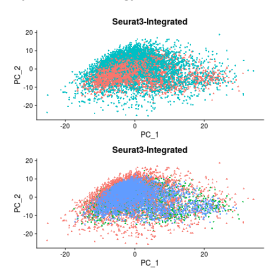
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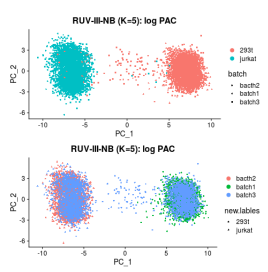
 There's a strong batch effects for Jurkat cells and biology (cell-type) is associated with batch (UV).



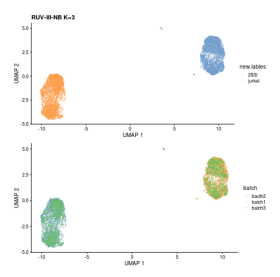
#### Seurat completely removes biology



RUV-III-NB removes batch effects without removing biology



### RUV-III-NB removes batch effects without removing biology



• Let  $\mu_{\mathbf{g}} = (\mu_{\mathbf{g1}}, \mu_{\mathbf{g2}}, \dots, \mu_{\mathbf{gN}})^{\mathsf{T}}$  be the vector of NB mean parameter for gene g across N cells, we assume  $\mathbf{y_g} \sim \mathbf{NB}(\mu_{\mathbf{g}}, \psi_{\mathbf{g}})$ , with

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- We assume that we have cell state information for  $n_a \le 3,000$  cells. This cell state information can come from:
  - For cell types: highly-confident annotation after initial LS normalization
  - For other factors, e.g. treatment, we have this information from experimental design.

For cells with annotation,

$$\log \mu_{\mathrm{g}}^{\mathtt{a}} = \zeta_{\mathrm{g}} + \mathsf{M} eta_{\mathrm{g}} + \mathcal{W}_{\mathtt{a}} lpha_{\mathrm{g}},$$

 $\mathbf{M}(\mathbf{n_a}\times\mathbf{m})$  matrix that contains dummy variables for cell states,  $\mathbf{W_a}(\mathbf{n_a}\times\mathbf{K})$  is rows subset of a K-dimensional *unknown* unwanted factors  $\boldsymbol{W}$  associated with annotated cells,  $\beta_{\mathbf{g}}\sim\mathbf{N}(\mathbf{0},\lambda_{\beta}^{-1}\mathbf{I_m}), \alpha_{\mathbf{g}}\sim\mathbf{N}(\mathbf{0},\lambda_{\alpha}^{-1}\mathbf{I_k})$ 

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For cells without annotation,

$$\log \boldsymbol{\mu}_{g}^{u} = \boldsymbol{\zeta}_{g} + \beta_{gc} + \boldsymbol{W}_{u}\boldsymbol{\alpha}_{g},$$

 $\mathbf{W_u}$  is rows subset of W associated with the un-annotated cells and  $\beta_{gc} \sim N(0, \lambda_{\beta}^{-1})$ 

 We also assume there is a negative control gene set (C) so that for any genes in this set,

$$\log \mu_g = \zeta_g + W \alpha_g,$$

 $\mathbf{W}(\mathbf{N} \times \mathbf{k})$  is a K-dimensional *unknown* unwanted factors for all cells

#### Adjusted data: log percentile-invariant adjusted count (PAC)

**Q** Calculate percentile under full fitted model:  $r_{gc} = rac{a_{cg} + b_{cg}}{2}$ , where

$$\begin{array}{lcl} \mathbf{a}_{gc} & = & F_{NB}(y_{gc}; \mu_{gc} = \mathbf{e}^{\hat{\zeta}_g + \hat{\beta}_{gc} + \hat{\mathbf{w}_c}^T \hat{\alpha}_g}, \hat{\psi}_g) \\ b_{gc} & = & F_{NB}(y_{gc} + 1; \mu_{gc} = \mathbf{e}^{\hat{\zeta}_g + \hat{\beta}_{gc} + \hat{\mathbf{w}_c}^T \hat{\alpha}_g}, \hat{\psi}_g) \end{array}$$

and  $\hat{w}_c$  the c<sup>th</sup> row of the matrix  $\hat{W}$ 

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② Invert the percentile under NB distribution where the mean is shifted to have average unwanted variations, where  $\overline{w}$  is vector of entries equal to the average level  $N^{-1} \sum_{c=1}^{N} \hat{w}_c$  of unwanted variation.

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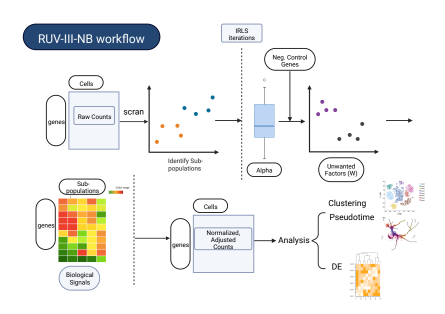
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- lacktriangledisplays Add 1 and take  $\log o \log(\mathsf{PAC}_{gc} + 1)$



### Parameter Estimation

- Iterative reweighted least squares (IRLS)-based
- $\bullet$  Parameters  $\zeta_{\it g}$  ,  ${\it W}_{\it g}$  ,  ${\it W}_{\it a}$  and  $\alpha_{\it g}$  are estimated using annotated cells
- ullet Parameters  $eta_{gc}$  and  $oldsymbol{W}_u$  are estimated using un-annotated cells.

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 Cell states information (M matrix): some cells need to have known cell states.

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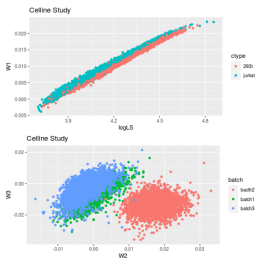
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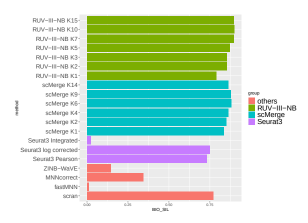
- Cell states information (M matrix): some cells need to have known cell states.
- Negative control gene sets: RUV-III-NB is a robust against a degree of miss-specification
- The number of unwanted factors (K): slight overestimation does not remove biological signals.

## Cell line Study: W estimates

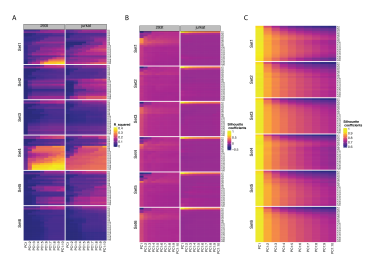
RUV-III-NB correctly identifies logLS and batch as the unwanted factors.



RUV-III-NB's performance is quite robust for a range of assumed unwanted factors (K)

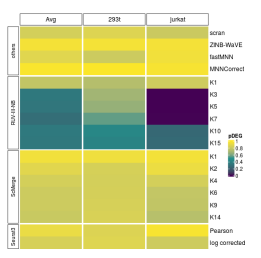


Robust performance with different sets of negative control genes



## Cell line Study: DEG

DEG of the same cell types located in different batches. RUV-III-NB adjusted data has the smallest amount of batch effects



### ZINB extension

 UMI data dominates in scRNA-seq world but there are still platforms without UMI

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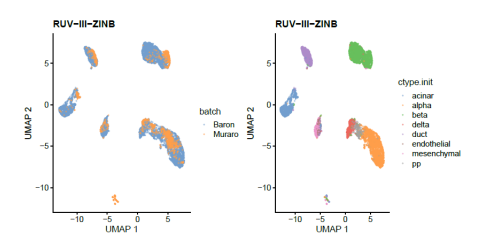
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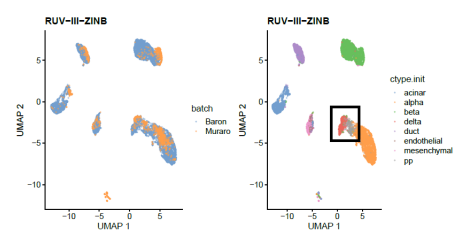
### **RUV-III-ZINB**



This is achieved with only 5% of the cells having known annotations.

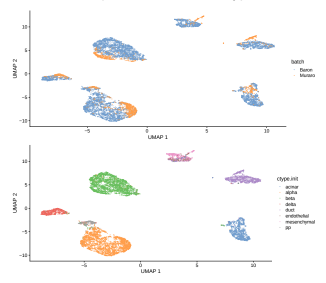
# Robustness against incorrect annotation?

We rerun RUV-III-ZINB assuming that the delta and PP cells are of the same cell-type



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#### RUV-III-ZINB can still separate the two cell-types



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- Future works: extensions to scMultiOmics and spatial transcriptomics.

## Acknowledgments

- Terry Speed, Ramyar Molania, Jianan Wang (WEHI)
- Alysha de Livera (La Trobe)
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