AUTHOR CORRECTION

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Author Correction: The shaky foundations of simulating single-cell RNA sequencing data



The original article can be found online at https://doi.org/10.1186/ s13059-023-02904-1.

*Correspondence: mark.robinson@imls.uzh.ch

 Department of Molecular Life Sciences, University of Zurich, Zurich, Switzerland
SIB Swiss Institute of Bioinformatics, University of Zurich, Zurich, Switzerland
TH Zurich, Zurich, Switzerland
Present address: Friedrich Miescher Institute for Biomedical Research and SIB Swiss Institute of Bioinformatics, Basel, Switzerland Correction: Genome Biol 24, 62 (2023) https://doi.org/10.1186/s13059-023-02904-1

Following publication of the original article [1], it was pointed out that the legend to Table 1 did not match the content of the table.



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The incorrect Table 1 is as follows:

Table 1 Overview of scRNA-seq simulators compared in this study. Methods are ordered alphabetically and annotated according to their (in)ability to accommodate multiple batches and/ or clusters, support for parallelization (parameter estimation and data simulation, respectively), software availability, and publication year. The right-most column catalogues neutral benchmark studies where each simulator was used. (\checkmark = yes, \bigstar = no, (\checkmark) = yes, but based on user input parameters, i.e., no support for parameter estimation, *requires random splitting of cells into two groups, \dagger/\ddagger = internal/prior resampling from empirical parameter distribution, •= no separate estimation step)

	Batches	Clusters	Type(s)	Cell #	Parallelization	Availability	Year	Model
BASiCS [<u>37</u>]	1	×	b	×	√ ×	<u>R/Bioc</u>	2015	NB
ESCO [<u>38</u>]	1	1	n,b,k	1	<i>」</i>	<u>R/GitHub</u>	2020	Gamma-Poisson
hierarchicell [<u>39]</u>	1	×	n,b	1	**	<u>R/GitHub</u>	2021	NB
muscat [<u>40</u>]	1	1	n,b,k	(✔)†	**	<u>R/Bioc</u>	2020	NB
POWSC [<u>41</u>]	×	1	n,k	(✔)†	**	<u>R/Bioc</u>	2020	zero-inflated, log- normal Poisson mixture
powsimR [<u>42]</u>	×	(√)	n*	(✔)†	<i>\</i>	<u>R/GitHub</u>	2017	NB
scDD [<u>43</u>]	×	×	n*	1	11	<u>R/Bioc</u>	2016	Bayesian NB mixture model
scDesign [<u>44</u>]	×	(✓)	n	1	o√	<u>R/GitHub</u>	2019	Gamma-Normal mixture model
scDesign2 [<u>45]</u>	×	1	n,k	1	√ <i>¥</i>	<u>R/GitHub</u>	2020	(zero-inflated) Poisson or NB + Gaussian copula for gene- gene correlations
SCRIP [<u>46</u>]	1	1	n,b,k	1	**	<u>R/GitHub</u>	2020	(Beta-)Gamma- Poisson
SPARSim [<u>47</u>]	1	×	n,b	(✔)‡	**	<u>R/GitLab</u>	2020	Gamma-multivar- iate hypergeo- metric
splatter [<u>15]</u> (Splat model)	(√)	(✓)	n	1	**	<u>R/Bioc</u>	2017	Gamma-Poisson
SPsimSeq [<u>16</u>]	1	×	n,b	1	•*	<u>R/Bioc</u>	2020	log-linear model-based density estima- tion + Gaussian copula for gene- gene correlations
SymSim [<u>48]</u>	1	×	n,b	1	**	<u>R/GitHub</u>	2019	kinetic model using MCMC
ZINB-WaVE [<u>49</u>]	1	1	n,b,k	×	**	<u>R/Bioc</u>	2018	zero-inflated NB
zingeR [<u>50</u>]	×	×	n	(✔)†‡	××	<u>R/GitHub</u>	2017	zero-inflated NB

Table 1 Overview of scRNA-seq simulators compared in this study. Methods are ordered alphabetically and annotated according to their (in)ability to accommodate multiple batches and/or clusters, support for parallelization (parameter estimation and data simulation, respectively), software availability, and publication year. Type(s)'column specifies which type of simulations can be produced (n: "singular" references: single batch or cluster; b: multiple batches; k: multiple clusters). 'Cell #' refers to whether the number of cells can be varied. Symbols: $\checkmark =$ yes, $\bigstar =$ no, (\checkmark) = yes, but based on user input parameters, i.e., no support for parameter estimation, *requires random splitting of cells into two groups, $\dagger/\ddagger =$ internal/prior resampling from empirical parameter distribution, $\bullet =$ no separate estimation step)

	Batches	Clusters	Type(s)	Cell #	Parallelization	Availability	Year	Model
BASICS [<u>37</u>]	1	×	b	×	/ ×	<u>R/Bioc</u>	2015	NB
ESCO [<u>38</u>]	1	1	n,b,k	1	<i>」」</i>	<u>R/GitHub</u>	2020	Gamma-Poisson
hierarchicell [<u>39]</u>	1	×	n,b	1	**	<u>R/GitHub</u>	2021	NB
muscat [<u>40</u>]	1	1	n,b,k	(√)†	**	<u>R/Bioc</u>	2020	NB
POWSC [<u>41</u>]	×	1	n,k	(✔)†	**	<u>R/Bioc</u>	2020	zero-inflated, log- normal Poisson mixture
powsimR [<u>42]</u>	×	(√)	n*	(√)†	\checkmark	<u>R/GitHub</u>	2017	NB
scDD [<u>43</u>]	×	×	n*	1	J J	<u>R/Bioc</u>	2016	Bayesian NB mixture model
scDesign [<u>44</u>]	×	(✓)	n	1	o√	<u>R/GitHub</u>	2019	Gamma-Normal mixture model
scDesign2 [<u>45]</u>	×	1	n,k	1	√ <i>¥</i>	<u>R/GitHub</u>	2020	(zero-inflated) Poisson or NB + Gaussian copula for gene- gene correlations
SCRIP [<u>46</u>]	1	1	n,b,k	1	**	<u>R/GitHub</u>	2020	(Beta-)Gamma- Poisson
SPARSim [<u>47</u>]	1	×	n,b	(✔)‡	**	<u>R/GitLab</u>	2020	Gamma-multivar- iate hypergeo- metric
splatter [<u>15]</u> (Splat model)	(✓)	(✓)	n	1	**	<u>R/Bioc</u>	2017	Gamma-Poisson
SPsimSeq [<u>16</u>]	1	×	n,b	1	•*	<u>R/Bioc</u>	2020	log-linear model-based density estima- tion + Gaussian copula for gene- gene correlations
SymSim [<u>48]</u>	1	×	n,b	1	**	<u>R/GitHub</u>	2019	kinetic model using MCMC
ZINB-WaVE [<u>49</u>]	1	1	n,b,k	×	**	<u>R/Bioc</u>	2018	zero-inflated NB
zingeR [<u>50</u>]	×	×	n	(✔)†‡	**	<u>R/GitHub</u>	2017	zero-inflated NB

The linked citations and the hyperlinks to the availability of data (in table 1) can be found in the original article. The original article [1] has been corrected.

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Reference

1. Crowell HL, Morillo Leonardo SX, Soneson C, et al. The shaky foundations of simulating single-cell RNA sequencing data. Genome Biol. 2023;24:62. https://doi.org/10.1186/s13059-023-02904-1.