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© Cell Interaction by Multiplet sequencing (CIM-seq) V.2

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dx.doi.org/10.17504/protocols.io.b2byqapw



Single cell sequencing methods facilitate the study of tissues at high resolution, revealing rare cell types with varying transcriptomes or genomes, but so far have been lacking the capacity to investigate cell-cell interactions. Here, we introduce CIM-seq, an unsupervised and high-throughput method to analyze direct physical cell-cell interactions between every cell type in a given tissue. CIM-seq is based on RNA sequencing of incompletely dissociated cells, followed by computational deconvolution of these into their constituent cell types using machine learning. CIM-seq is broadly applicable to studies that aim to simultaneously investigate the constituent cell types and the global interaction profile in a specific tissue.

DOI

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https://www.nature.com/articles/s41592-021-01196-2

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cell interaction, rnaseq, single cell

_____ protocol,

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55384
MATERIALS
Biolabs Catalog #M0262L

⋈ HotStart ReadyMix (KAPA HiFi PCR kit) Kapa

Biosystems Catalog #KK2601
⊗ psfTn5 addgene Catalog #79107
⊠ 10% SDS
solution Teknova Catalog #S0287
SMARTScribe Reverse
Transcriptase Takarabio Catalog #634888
Aldrich Catalog #M1028
XTriton X-100
Sigma Catalog #93426
XKAPA HiFi PCR kit with dNTPs Fisher
Scientific Catalog #NC0142652

    □ Recombinant RNAse
    □

Inhibitor Takarabio Catalog #2313A
Betaine 5M Sigma
Aldrich Catalog #B0300
Fisher Catalog #4456740
⊠ USB Dithiothreitol (DTT), 0.1M Solution Thermo
Fisher Catalog #707265ML
⊠ UltraPure™ DEPC-Treated Water Thermo
Fisher Catalog #750023
⊠ dNTP Mix (10 mM each) Thermo
Fisher Catalog #R0192
Sera-Mag Speed Beads Ge
Healthcare Catalog #65152105050250
```



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₩ Hard-Shell® 384-Well PCR Plates thin wall skirted BioRad

Sciences Catalog #HSP3801

℧ Qubit 1X dsDNA HS Assay Kit **Thermo Fisher**

Scientific Catalog #Q33231

□ PEG 8000 - (Polyethylene glycol) Sigma

Aldrich Catalog #P2139

XTAPS Sigma

Aldrich Catalog #T5130

Oligonucleotides (all ordered from IDT using Standard desalting, except barcodes ordered in solution/plates)

Oligo-dT:

(N2:25252525)

IS_PCR: 5'-AAGCAGTGGTATCAACGCAGAGT-3'

TSO: 5'-AAGCAGTGGTATCAACGCAGAGTACATrGrG+G-3'
ME-A: 5'-TCGTCGGCAGCGTCAGATGTGTATAAGAGACAG-3'
ME-B: 5'-GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAG-3'

ME-Rev: 5'-/5Phos/CTGTCTCTTATACACATCT-3'

Illumina-compatible barcodes used (Sxxx/Nxxx series, n=784) are available as a supplementary table in the manuscript.

Before preparing cell lysis plates it is recommended to thoroughly clean all equipment with 70% EtOH and RNAse away to prevent contamination and avoid RNA degradation.

Prepare Lysis buffer

- 1 CIM-seq is compatible with both plate based and droplet based single cell RNA-seq methods. The following protocol is for Smart-seq2 based CIM-seq. For droplet based (10x), see the methods section of the manuscript and the vignettes in the Enge lab Github repository.
- 2 Prepare Lysis Buffer:

NOTE: Reagents are prepared on ice, working quickly. ERCC is stored in single-use aliquots at \$ -80 °C , thawed on ice and added last.

Reagent:	Reagent	μl per
	concentration:	reaction:
H20	-	1.31
Inhibitor	1 U/μΙ	0.05
ERCC	-	0.05
(1:600		
000)		
Triton-	0.2%	0.04
X100		
(10%		
solution)		
10mM	2.5mM/each	0.5
dNTP		
100uM	2.5μΜ	0.05
dT		
Total	-	2

Add **2 µL lysis buffer mix** to each well. Cover with appropriate lids. Spin down.

Snap freeze on dry ice. Store until use at 8 -80 °C

Sort cells

3 Sort single cells and multiplets (aggregates of multiple cells) into **2 μL lysis buffer mix.**Multiplets can be discerned from singlets by gating on the basis of FSC-W (Forward scatter - Width) and FSC-H (Forward scatter - Height) (see **Figure 1**).

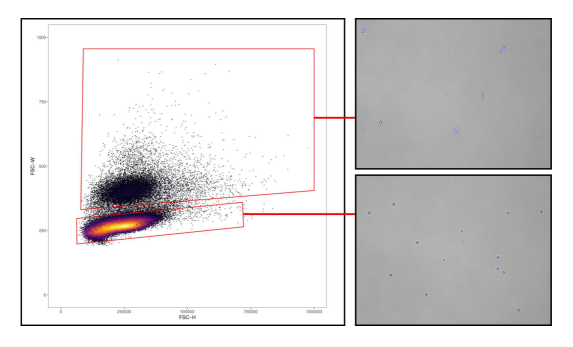


Figure 1. Gating scheme and result of FSC-W and FSC-H based sort of multiplets (top) and singlets (bottom) using HCT116 cells .

Following sort, immediately seal with appropriate seals (approved for -80C > 100C) and centrifuge

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at **32000 x g, 4°C, 00:05:00**.

Reverse transcription and cDNA amplification

4 Primer annealing

5 Prepare RT master-mix

Made fresh.

Reagent:	Reaction	Reagent
	concentration:	volume:
SmartScribe	15u/µl	0.475
RNase	1.66u/µl	0.125
Inhibitor		
5x First	1X	1
Strand		
buffer		
DTT	8.33mM	0.25
(100mM)		
Betaine	1.66M	1
(5M) [fridge]		
MgCl2 (1M)	10mM	0.03
[bench]		
TS0	1.66µM	0.05
(100uM)		
H20	-	0.07
Total	-	3

Dispense **□3 µL** per well.

Cover plate with new film and spin down.

6 Incubate in thermocycler

8 42 °C © 01:30:00

84°C hold



7 cDNA preamplification

Made fresh.

Reagents	Reaction	Reagent
	concentration:	volume:
H20	-	1.0688
Kapa HiFi HotStart ReadyMix	1X	6.25
(2x)		
IS_PCR primer (10uM)	0.1μΜ	0.125
Lambda Exonuclease	0,045u/µl	0.05625
Total	-	7.5000

Dispense \blacksquare 7.5 μ L per well . Total reaction volume will be 12.5 μ l. Spin down. Cover with new lid.

8 Incubate in thermocycler with the following program:

Step	Temperature	Time	Cycles
Lambda	37°C	30 min	1x
exonuclease			
Initial	95°C	3 min	1x
denaturation			
Denaturation	98°C	20 sec	18-24x
Annealing	67°C	15 sec	
Elongation	72°C	4 min	
Final	72°C	5 min	1x
elongation			
	4C	Hold	

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cDNA cleanup

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We prepare SPRI-beads in 20% PEG-8000 solution as in: https://openwetware.org/wiki/SPRI_bead_mix#Ingredients_for_50_mL_2

Using 20% SPRI-bead solution:

- 1. Add 0.7x the reaction volume of SPRI beads per well. Mix well by pipetting. (i.e \blacksquare 8.75 μ L
- SPRI-bead solution for $\blacksquare 12.5 \, \mu L$ reaction volume)
- 2. Incubate © 00:05:00 & Room temperature
- 3. Place on magnetic stand for © 00:03:00
- 4. Carefully remove supernatant
- 5. Add 40 μl 80% EtOH and incubate **© 00:00:30**
- 6. Remove EtOH (without disturbing the beads)
- 7. Wash again with EtOH. Make sure to remove well.
- 8. Allow beads to air-dry for **© 00:10:00 © 00:15:00**
- 9. Remove plate from magnetic stand
- 10. Elute beads in $\blacksquare 15 \,\mu L$ EB or TE buffer. Mix well by pipetting
- 11. Incubate **⋄ 00:05:00 ♦ Room temperature**
- 12. Place on magnetic plate for © 00:03:00
- 13. Optional: Carefully remove supernatant to the elution plate

10 cDNA quantification

We measure concentration of random wells using Qubit HS dsDNA, adapted to a 96-well plate reader.

- 1. Add **97 μL** of 1X Qubit HS dsDNA solution to a flat-bottom, black plate
- 2. Add 3 µL of cDNA sample
- 3. Add Standards (NOTE: We make a 8-step ladder from $0 ng/\mu l \rightarrow 10 ng/\mu l$ Qubit Standard DNA in TE buffer)
- 3. Read in plate reader using 485nM excitation/528nm emission
- 4. Calculate cDNA concentration

11 (optional) cDNA quality control

Using Agilent HS 5000 DNA chips (or equivalent)

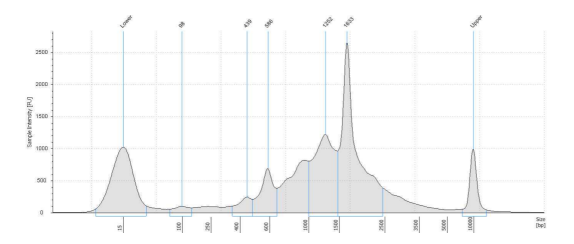


Figure 2. cDNA profile of single cell run on HS D5000 Agilent tapestation

12 Make cDNA dilution plate

Dilute cDNA in water based on average concentration from Qubit measurements. Target concentration 150pg per μ l in 15μ L.

cDNA tagmentation

13 Tn5 digestion

Tn5 is produced from psfTn5 (Addgene #79107), purified to ~3mg/ml and assembled with Illumina Tn5 adapters (see oligos) as in Picelli et al, 2014.

Picelli S, Björklund AK, Reinius B, Sagasser S, Winberg G, Sandberg R (2014). Tn5 transposase and tagmentation procedures for massively scaled sequencing projects.. Genome research.

https://doi.org/10.1101/gr.177881.114

14 Prepare Tn5 master mix

NOTE: TAPS-PEG Buffer contains PEG, which is viscous. Buffer should equilibrate to room temperature before use to allow proper mixing.



Reagent	Reaction	µl per
	conc.	reaction
Nuclease	-	1.05
free H2O		
TAPS-	10mM	0.5
PEG	TAPS,	
(50mM	5mM	
TAPS,	MgCl2,	
25mM	8% PEG-	
MgCl2,	8000	
40%		
PEG-		
8000)		
psfTn5,		0.25
loaded		
with		
50µM		
MEDS-		
A/B		
Total		1.8

Dispense 1.8 µL per well in a new plate(tagmentation plate)

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Add $\mathbf{D0.7} \, \mu \mathbf{L} \, \text{cDNA}$ (normalized to 150pg/ μ l)

Mix well by vortexing plate. Cover with new lid and spin down.

16 Incubate in thermocycler at § 55 °C © 00:10:00

Remove immediately and stop reaction by adding $\Box 1 \mu L$ per well of 0.1% SDS.

Vortex, spin down and incubate **© 00:07:00** at **§ Room temperature**

cDNA library PCR and barcoding

17 Make PCR master-mix

Reagents	µl per	
	reaction	
H20	13.25	
5x buffer	5	
dNTPs	0.75	
KAPA	0.5	
Total	19.50	

Dispense $\blacksquare 19.5 \,\mu L$ per well to tagmentation plate (containing $\blacksquare 3.5 \,\mu L$ sample after step 14)

18 Add primers/barcodes

 $\mathbf{2}\mu\mathbf{L}$ per well (from 384-well index plates, with 3.75 μ M/each forward/reverse primers; **see** oligos in materials).

Total reaction volume is $\blacksquare 25~\mu L$ ($\blacksquare 3.5~\mu L$ sample + $\blacksquare 19.5~\mu L$ PCR mix and $\blacksquare 2~\mu L$ primers).

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Vortex. Spin down and cover. Incubate in thermocycler as following:

Step	Temperature	Time	Cycles
Gap fill	72°C	3 min	1x
First	95°C	30 sec	1x
denature			
Denature	95°C	15 sec	12x
Denature	67°C	30 sec	
Denature	72°C	45 sec	
Final	72°C	4 min	1x
extension			
	4-10°C	hold	

20 Pool $\mathbf{2.5} \, \mu \mathbf{L}$ from each well to an 1.5ml Eppendorf tube.

21 Library cleanup

- 1. Add 0.9x pooled library volume of SPRI-bead solution. Incubate for © 00:05:00 at
- **8** Room temperature .
- 2. Place on magnetic rack for **© 00:03:00**.



- 3. Remove supernatant without disturbing magnetic beads.
- 4. Add at least 1 mL 80% EtOH (fresh). Incubate for 00:00:30.
- 5. Remove supernatant.
- 6. Repeat EtOH wash.
- 7. Air dry for **© 00:10:00 © 00:15:00** .
- 8. Re-suspend beads thoroughly in 100 µl EB or TE buffer.
- 9. Place eppendorf on magnetic rack for **© 00:03:00**.
- 10. Transfer supernatant to new 1.5ml Eppendorf tube.
- 11. Repeat cleanup (from step 1-7) and elute in 30 µl EB or TE buffer.
- 12.(Optional) Place eppendorf on magnetic rack for © 00:03:00 and transfer supernatant to new tube.

22 Pooled library QC

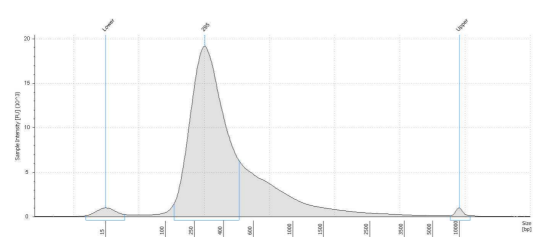


Figure 3. cDNA profile of a library of 784 cells (both single cells and multiplets) on HS D5000 Agilent tapestation.

Data pre-processing

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A series of **pre-processing steps** must be performed in order to generate a counts file:

1. Trim reads, remove adapter sequences and align RNAseq data to reference genome using STAR:

Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, Batut P, Chaisson M, Gingeras TR (2013). STAR: ultrafast universal RNA-seq aligner.. Bioinformatics (Oxford, England).

https://doi.org/10.1093/bioinformatics/bts635

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2. Remove duplicate reads using Picard:

McKenna A, Hanna M, Banks E, Sivachenko A, Cibulskis K, Kernytsky A, Garimella K, Altshuler D, Gabriel S, Daly M, DePristo MA (2010). The Genome Analysis Toolkit: a MapReduce framework for analyzing next-generation DNA sequencing data.. Genome research. https://doi.org/10.1101/gr.107524.110

3. Generate transcript counts file using HTSeq:

Anders S, Pyl PT, Huber W (2015). HTSeq--a Python framework to work with high-throughput sequencing data.. Bioinformatics (Oxford, England).

https://doi.org/10.1093/bioinformatics/btu638

Dimensionality reduction and classification

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Once a counts file has been generated the data can be analyzed. CIM-seq requires four arguments in order to run:

- 1. The raw counts data with gene IDs as rownames and sample IDs as colnames.
- 2. The ERCC spike-in counts data with gene IDs as rownames and sample IDs as colnames.
- 3. The dimensionality reduced representation of the data.
- 4. A class for each of the individual singlets.

In order to generate the last two of these we recommend using the Seurat package in R, as CIMseq is implemented in R as well. A number of tutorials for Seurat can be found on the Satijalab website:

https://satijalab.org/seurat/vignettes.html

CIM-seq

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CIM-seq can be downloaded from: https://github.com/EngeLab/CIMseq

Or installed directly in R using the devtools package:

devtools::install_github("EngeLab/CIMseq")

The CIM-seq vignette can be found at:

https://github.com/EngeLab/CIMseq/tree/master/vignettes

