

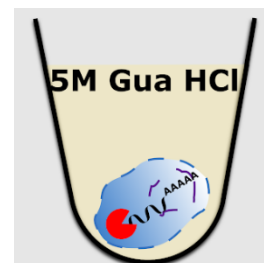
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Version 1

# gmcSCRB-seq protocol V.1

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**We use this protocol and it's working**

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## Abstract

gmcSCRB-seq is an alternative lysis protocol to mcSCRB-seq ([Publication](#) and [Protocol](#)). gmcSCRB-seq uses Guanidine Hydrochloride in the lysis buffer and requires an additional clean up step. We recommend using this alternative lysis protocol in cases where cells are difficult to lyse or where the DNA appears degraded following the pre-amplification step. However, we have observed reduced sensitivity with gmcSCRB-seq compared to mcSCRB-seq and would therefore only recommend using it in cases where the latter does not produce high quality libraries.

## Attachments



[mcSCRBseq\\_oligodT.tx..](#)

44KB

## Guidelines

- For troubleshooting help, feel free to join our [mcSCRB-seq Slack channel](#), leave your question in the comments section, or message us directly.
- The complete list of reagents and plastic ware with order numbers can be found in the 'Materials' section.
- Make sure all steps involving single-cell lysate and RNA before reverse transcription are carried out swiftly.
- Size selection of libraries is optional, but has in our experience improved reliability in cluster densities when sequencing.
- All primer sequences are listed below:

|  | Oligo                             | Vendor | Purification | Concentration | Sequence                                                                   |
|--|-----------------------------------|--------|--------------|---------------|----------------------------------------------------------------------------|
|  | barcoded oligo-dT (E3V6NEXT)      | IDT    | TruGrade     | 2 µM          | Biotin-<br>ACACTCTTTCCCTACACGA<br>CGCTCTTCCGATCT[BC6]<br>[UMI10][T30]VN    |
|  | TSO unblocked (E5V6NEXT)          | IDT    | HPLC         | 100 µM        | ACACTCTTTCCCTACACGA<br>CGCrGrGrG                                           |
|  | PreAmp (SINGV6)                   | IDT    | Desalted     | 10 µM         | Biotin-<br>ACACTCTTTCCCTACACGA<br>CGC                                      |
|  | 3' enrichment primer (P5NEXTPT 5) | IDT    | HPLC         | 5 µM          | AATGATACGGCGACCACC<br>GAGATCTACACTCTTTCCC<br>TACACGACGCTCTTCCG*A<br>*T*C*T |
|  | i7 Index Primer (N7XX)            | IDT    | TruGrade     | 5 µM          | CAAGCAGAAGACGGCATA<br>CGAGAT[i7]GTCTCGTGGG<br>CTCGG                        |

Find the cell barcode sequences in the attached text file (Abstract tab).



## Materials

|  | NAME                                                        | CATALOG # | VENDOR                         |
|--|-------------------------------------------------------------|-----------|--------------------------------|
|  | EDTA<br>0.5<br>M                                            | E7889     | Sigma<br>Aldrich               |
|  | Ethanol,<br>absolute                                        | 9065.4    | Carl<br>Roth                   |
|  | Exonuclease I<br>(20<br>U/ $\mu$ l)                         | ENO582    | Thermo<br>Fisher<br>Scientific |
|  | Exonuclease I<br>Reaction<br>Buffer<br>(10x)                | ENO582    | Thermo<br>Fisher<br>Scientific |
|  | IGEPAL<br>CA-630                                            | I8896     | Sigma<br>Aldrich               |
|  | Maxima<br>H-Reverse<br>Transcriptase<br>(200<br>U/ $\mu$ l) | EP0753    | Thermo<br>Fisher<br>Scientific |
|  | Maxima<br>RT<br>Buffer<br>(5x)                              | EP0753    | Thermo<br>Fisher<br>Scientific |
|  | Polyethylene<br>glycol                                      | 89510     | Sigma<br>Aldrich               |



|  |                                                             |                            |                                  |
|--|-------------------------------------------------------------|----------------------------|----------------------------------|
|  | 800<br>0                                                    |                            |                                  |
|  | PBS<br>7.4                                                  | 1001<br>0-23               | Gibco -<br>Thermo<br>Fischer     |
|  | Phusion<br>HF<br>Buffer                                     | B051<br>8                  | New<br>England<br>and<br>Biolabs |
|  | Proteinase<br>K                                             | 903<br>4                   | Takara                           |
|  | Sera-<br>Mag<br>Speed<br>Beads                              | 6515<br>2105<br>050<br>250 | Thermo<br>Fisher<br>Scientific   |
|  | Terra<br>PCR<br>Direct<br>Polymerase<br>Mix                 | 639<br>271                 | Takara                           |
|  | Ultra<br>Pure<br>DNAse/RNase-<br>Free<br>Distilled<br>Water | 1097<br>7-<br>049          | Invitrogen -<br>Thermo<br>Fisher |
|  | Sodium<br>Azide<br>99.5<br>%                                | S20<br>02-<br>100<br>G     | Sigma<br>Aldrich                 |
|  | Sodium<br>Chloride<br>5M                                    | S515<br>0-1L               | Sigma<br>Aldrich                 |



|  |                                                |             |                            |
|--|------------------------------------------------|-------------|----------------------------|
|  | Trizma hydrochloride solution 1M pH 8.0        | T2694       | Sigma Aldrich              |
|  | Bioanalyzer High Sensitivity DNA Analysis Kits | 5067-4626   | Agilent Technologies       |
|  | MinElute Gel Extraction Kit                    | 28606       | Qiagen                     |
|  | Nextera XT DNA Library Preparation Kit         | FC-131-1096 | illumina                   |
|  | Quant-iT PicoGreen dsDNA Assay Kit             | P7589       | Invitrogen - Thermo Fisher |
|  | dNTPs (25 mM each)                             | R0182       | Thermo Fisher Scientific   |
|  | Aluminum                                       | 391-1275    |                            |



|  |                                     |            |                                         |
|--|-------------------------------------|------------|-----------------------------------------|
|  | seals for cold storage              |            |                                         |
|  | Adhesive PCR film seals             | AB0558     | <a href="#">ThermoFisher Scientific</a> |
|  | twin.tec 96-well DNA LoBind Plates  | 0030129504 | <a href="#">Eppendorf</a>               |
|  | twin.tec 384-well DNA LoBind Plates | 0030129547 | <a href="#">Eppendorf</a>               |
|  | 0.5 ml PCR clean tube DNA LoBind    | 0030108035 | <a href="#">Eppendorf</a>               |
|  | 1.5 ml PCR clean tube DNA LoBind    | 0030108051 | <a href="#">Eppendorf</a>               |



|  |                                                            |                    |                                                  |
|--|------------------------------------------------------------|--------------------|--------------------------------------------------|
|  | 5.0<br>ml<br>PCR<br>clea<br>n<br>tube<br>DNA<br>LoBi<br>nd | 003<br>0108<br>310 | Epp<br>end<br>orf                                |
|  | 15<br>ml<br>PCR<br>clea<br>n<br>tube<br>DNA<br>LoBi<br>nd  | 003<br>0122<br>208 | Epp<br>end<br>orf                                |
|  | E-<br>Gel<br>EX<br>Agar<br>ose<br>Gels<br>, 2%             | G40<br>200<br>2    | Invitr<br>oge<br>n -<br>Ther<br>mo<br>Fish<br>er |

## Before start

Wipe bench surfaces with RNase Away and keep working environment clean.

## Preparation of lysis plates

- 1 Prepare **Lysis Buffer** for the number of plates needed.

### Note

Note that 96-well plates are the preferred setup for gmcSCRB-seq, as the well cleanup (Step 9-15) would be more difficult with 384-well plates.

|  | Reagent                    | 96-well plate |
|--|----------------------------|---------------|
|  | 8M Guanadine Hydrochloride | 343.75        |
|  | 2-mercaptoethanol          | 5.5           |
|  | NEB HF Phusion buffer (5x) | 1.1           |
|  | H2O                        | 199.65        |
|  | <b>Total</b>               | 550           |

- 2 Prepare 96 well plate(s) containing 5  $\mu$ L **Lysis Buffer** per well.

### Note

Lysis plates should be prepared shortly before use, but can be stored for up to 1 week at room temperature. Prior to use, double check to make sure salts have not fallen out of solution or that the lysis buffer has not evaporated.

 5  $\mu$ L Lysis Buffer

## Sample Collection

- 3 Sort 1 cell to each well of a 96 well plate containing 5  $\mu$ L **Lysis Buffer**.
- 4 Immediately seal the plate with an aluminium cold storage seal.



- 5 In a cooled centrifuge, spin down the plate for 30 sec @ max. speed and place immediately on dry ice.

🧊 -80 °C Store plates containing single-cell lysates in a -80 °C freezer for up to 6 months.

## Proteinase K Digest

- 6 Thaw plates briefly (up to 1 min) at room temperature
- 7 Spin down (30 sec @ 1000 rcf) in a centrifuge pre-cooled to 4 °C.

🧊 4 °C

- 8 Prepare **Clean-up Beads**:

|  | Reagent                    | Amount       |
|--|----------------------------|--------------|
|  | PEG 8000                   | 11 g         |
|  | NaCl, 5M                   | 10 mL        |
|  | Tris-HCL, 1M, pH 8.0       | 500 µL       |
|  | EDTA, 0.5M                 | 100 µL       |
|  | IGEPAL, 10% solution       | 50 µL        |
|  | Sodium Azide, 10% solution | 250 µL       |
|  | UltraPure Water            | up to 49 mL  |
|  | <b>Total</b>               | <b>49 mL</b> |

-Add all ingredients into a 50 mL falcon tube, but do not add the total amount of water until after PEG is completely solubilized

- Incubate at 40°C and vortex regularly until PEG is completely dissolved
- Resuspend bead stock carefully (Sera-Mag Speed Beads)
- Pipette 1000 µL of bead suspension into a 1.5 mL tube
- Place on magnet stand
- Remove supernatant



- Add 1000  $\mu$ L 10 mM Tris-HCl, pH 8.0, 1 mM EDTA (TE), and resuspend beads
- Place on magnet stand
- Remove supernatant
- Repeat wash one more time
- Add 900  $\mu$ L 10 mM Tris-HCl, pH 8.0, 1 mM EDTA (TE), and resuspend beads
- Add to PEG solution above and mix well.

**Note**

Beads should be prepared ahead of time and can be stored at 4 °C or room temperature.

- 9 Add 10  $\mu$ L of **Clean-up Beads** to each well (1:2 ratio).

 10  $\mu$ L Clean-up Beads

- 10 Seal the plate and vortex to mix the bead-lysate mixture. Briefly spin down the plate (<200 rcf, <10 sec) so all of the mixture is at the bottom of the well but the beads are still in solution.

- 11 Incubate for 5 minutes at RT.

 00:05:00 binding of the cDNA onto the beads

 20 °C Room temperature

- 12 Place on magnet stand until clear

- 13 Discard supernatant

- 14 Wash twice with 50  $\mu$ L 80% ethanol (while on magnet) and discard supernatant

 50  $\mu$ L 80% ethanol (freshly prepared)

- 15  00:05:00 air dry beads

**Note**

Depending on lab temperature and humidity, drying times can vary.



## Reverse Transcription

16 Prepare **Reverse Transcription Mix** as follows:

| Reagent                              | 96-well plate                |
|--------------------------------------|------------------------------|
| UltraPure Water                      | 88 $\mu$ L                   |
| PEG 8000 (50 % solution)             | 165 $\mu$ L                  |
| Maxima RT Buffer (5x)                | 220 $\mu$ L                  |
| dNTPs (25 mM each)                   | 44 $\mu$ L                   |
| TSO E5V6NEXT unblocked (100 $\mu$ M) | 22 $\mu$ L                   |
| Maxima H Minus RT (200 U/ $\mu$ l)   | 11 $\mu$ L                   |
| <b>Total</b>                         | <b>550 <math>\mu</math>L</b> |

### Note

- RT MM can be prepared while plate is incubating or while drying on magnet.
- If ERCCs will be used, decrease the amount of H<sub>2</sub>O and add appropriate amount of ERCCs.
- Caution: Reverse Transcription Mix with PEG needs to be mixed carefully!

 4 °C Keep Reverse Transcription Mix on ice

17 Once drying is complete, add 4  $\mu$ L H<sub>2</sub>O to each well.

 4  $\mu$ L H<sub>2</sub>O

18 Add 5  $\mu$ L Reverse Transcription Mix to each well.

 5  $\mu$ L Reverse Transcription Mix

**Note**

If a robot (eg. Formulatrix Mantis) is used, make sure to calibrate correctly to the viscous solution.

- 19 Add 1  $\mu$ L of barcoded oligo-dT primer [2  $\mu$ M] (E3V6NEXT adapter) to each well.

 1  $\mu$ L barcoded oligo-dT primer [2  $\mu$ M] (E3V6NEXT adapter)

- 20 Seal plate with a PCR seal, vortex briefly and spin down (30 sec @ 1000 rcf) in a centrifuge pre-cooled to 4 °C.

- 21 In a thermocycler with heated lid, incubate:

 42 °C 90 min

 8 °C  $\infty$

**cDNA Pooling & Purification**

- 22 Vortex to mix the bead-cDNA mixture. Briefly spin down the plate (<200 rcf, <10 sec) so all of the mixture is at the bottom of the well but the beads are still in solution.

- 23 Prepare **Bead Binding Buffer**

|  | <b>Rea<br/>gent</b>               | <b>Amo<br/>unt</b> |
|--|-----------------------------------|--------------------|
|  | PEG<br>800<br>0                   | 11 g               |
|  | NaCl<br>, 5M                      | 10<br>mL           |
|  | Tris-<br>HCL,<br>1M,<br>pH<br>8.0 | 500<br>$\mu$ L     |
|  | EDT<br>A,<br>0.5<br>M             | 100<br>$\mu$ L     |
|  | IGEP<br>AL,<br>10%                | 50 $\mu$<br>L      |

|  |                                                 |                    |
|--|-------------------------------------------------|--------------------|
|  | solut<br>ion                                    |                    |
|  | Sodi<br>um<br>Azid<br>e,<br>10%<br>solut<br>ion | 250<br>μL          |
|  | Ultra<br>Pure<br>Wat<br>er                      | up<br>to 5<br>0 mL |
|  | <b>Tota<br/>l</b>                               | <b>50<br/>mL</b>   |

-Add all ingredients into a 50 mL falcon tube, but do not add the total amount of water until after PEG is completely solubilized

-Incubate at 40°C and vortex regularly until PEG is completely dissolved

- 24 Pool all wells (including beads) of one plate into a 2 mL tube and add 960 μL (ratio 1/1)  
**Bead Binding Buffer**

 960 μL Bead Binding Buffer

#### Note

Vortex to mix the sample and bead binding buffer.

25

 00:05:00 binding of the cDNA onto the beads

 20 °C Room temperature

- 26 Place on magnet stand until clear

- 27 Discard supernatant

- 28 Wash twice with 2 mL 80% ethanol (while on magnet) and discard supernatant



🧪 2  $\mu$ L 80% ethanol (freshly prepared)

29 ⌚ 00:05:00 air dry beads

## Exonuclease I Treatment

30 Elute cDNA in 17  $\mu$ L UltraPure Water & transfer to new tube

🧪 17  $\mu$ L UltraPure Water

### Note

Avoid transferring beads as these can inhibit the downstream PCR.

31 To the 17  $\mu$ L cDNA, add:

🧪 2  $\mu$ L Exonuclease I Buffer (10x)

🧪 1  $\mu$ L Exonuclease I (20 U/ $\mu$ L)

## Full length cDNA amplification

32 In a thermocycler with heated lid, incubate:

🔥 37 °C 20 min (ExoI digest)

🔥 80 °C 10 min (Heat inactivation)

🔥 8 °C  $\infty$  (Store)

33 Prepare **PreAmplification Mix** as follows:

| Reagent                            | 1x                          |
|------------------------------------|-----------------------------|
| Terra direct Buffer (2x)           | 25 $\mu$ L                  |
| SINGV6 Primer (10 $\mu$ M)         | 1 $\mu$ L                   |
| Terra polymerase (1.25 U/ $\mu$ L) | 1 $\mu$ L                   |
| UltraPure Water                    | 3 $\mu$ L                   |
| <b>Total</b>                       | <b>30 <math>\mu</math>L</b> |

34 Add 30  $\mu$ L of **PreAmplification Mix** directly to the Exonuclease I digested sample.

🧪 30  $\mu$ L PreAmplification Mix

## cDNA purification & quantification

35 In a thermocycler with heated lid, incubate as follows:

| Step                 | Temperature | Time   | Cycles |
|----------------------|-------------|--------|--------|
| Initial Denaturation | 98 °C       | 3 min  | 1x     |
| Denaturation         | 98 °C       | 15 sec | 13-21x |
| Annealing            | 65 °C       | 30 sec |        |
| Elongation           | 68 °C       | 4 min  |        |
| Final Elongation     | 72 °C       | 10 min | 1x     |
| Store                | 8 °C        | ∞      |        |

### Note

Cycle number highly depends on the input amount and should be optimized depending on the specific cell type used in the experiment. For ES cells, 13-15 cycles are sufficient.

36 Mix PreAmplification PCR with 40 µL **Clean-up Beads** (1/0.8 ratio)

 40 µL Clean-up Beads

37  00:05:00 binding of the cDNA onto the beads

38 Place on magnet until clear and discard supernatant

39 Wash twice with 150 µL 80% ethanol (while on magnet) and discard supernatant

 150 µL 80% ethanol (freshly prepared)

40  00:05:00 air dry beads

41 Elute cDNA in 15 µL UltraPure Water & transfer to new tube

 15 µL UltraPure Water

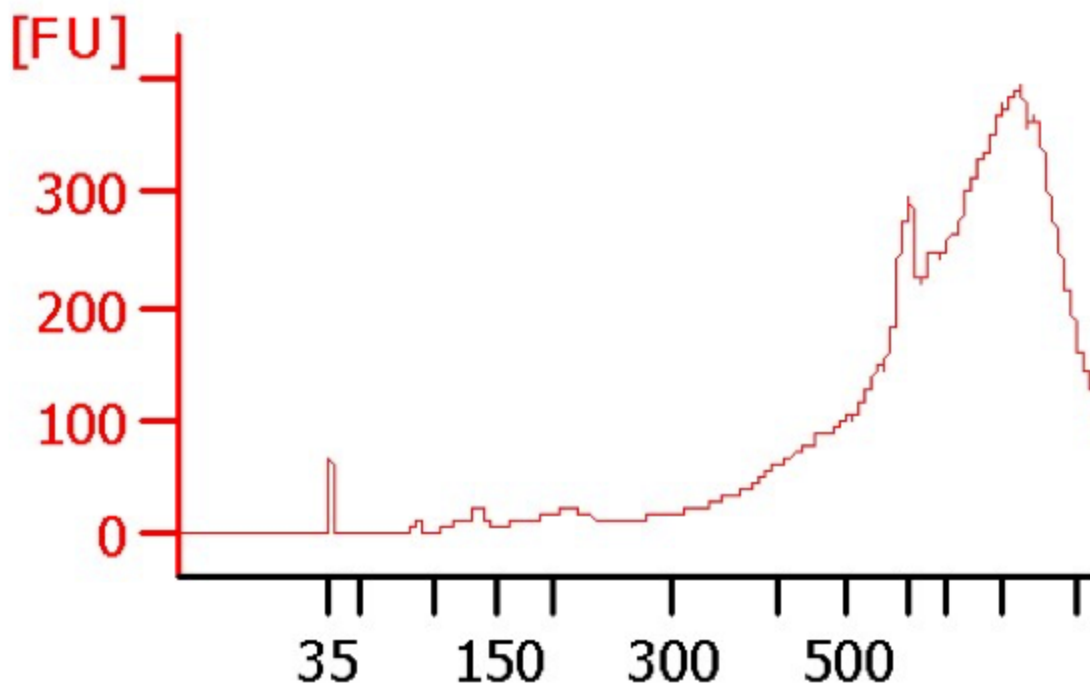
- 42 Quantify the cDNA using the Quant-iT PicoGreen dsDNA assay kit or equivalent Qubit following the manufacturer's protocol. Use 1  $\mu$ l of clean cDNA for quantification.

#### Expected result

cDNA concentration should be > 1 ng/ $\mu$ l, depending on cell type and cycle number

- 43 Optional: Quality check the cDNA using the Agilent 2100 Bioanalyzer with High Sensitivity DNA Analysis Kits.

#### Expected result



## Tagmentation, Library PCR & Indexing

- 44 Prepare **Tagmentation Mix** and dispense 19  $\mu\text{L}$  to a new 96-well plate.

| Reagent                    | 1x                                 |
|----------------------------|------------------------------------|
| Tagment DNA Buffer (2x)    | 10 $\mu\text{L}$                   |
| Amplicon Tagment Mix (Tn5) | 5 $\mu\text{L}$                    |
| UltraPure Water            | 4 $\mu\text{L}$                    |
| <b>Total</b>               | <b>19 <math>\mu\text{L}</math></b> |

 19  $\mu\text{L}$  Tagmentation Mix

- 45 Dilute cDNA to 0.8 ng/ $\mu\text{L}$  and add 1  $\mu\text{L}$  to each reaction.

 1  $\mu\text{L}$  cDNA (0.8 ng/ $\mu\text{L}$ )

- 46 In a thermocycler with heated lid, incubate as follows:

 55  $^{\circ}\text{C}$  Tagmentation

 00:10:00 Tagmentation

- 47 To stop the reaction, add 5  $\mu\text{L}$  NT buffer to each reaction and mix by pipetting up and down.

 5  $\mu\text{L}$  NT Buffer

 00:05:00 Incubation at room temperature

- 48 Prepare **3' Enrichment PCR Mix** as follows and add 24.5  $\mu\text{L}$  to each tagmentation reaction.

| Reagent                      | 1x                                   |
|------------------------------|--------------------------------------|
| NPM PCR Mix                  | 15 $\mu\text{L}$                     |
| P5NEXTPT5 (5 $\mu\text{M}$ ) | 0.5 $\mu\text{L}$                    |
| UltraPure Water              | 9 $\mu\text{L}$                      |
| <b>Total</b>                 | <b>24.5 <math>\mu\text{L}</math></b> |

 24.5  $\mu\text{L}$  3' Enrichment PCR Mix

- 49 Add 0.5  $\mu\text{L}$  of i7 index primer (5  $\mu\text{M}$ )

 0.5  $\mu\text{L}$  i7 index primer (5  $\mu\text{M}$ )

50 In a thermocycler with heated lid, incubate as follows:

| Step                 | Temperature | Time   | Cycles |
|----------------------|-------------|--------|--------|
| Gap-fill             | 72 °C       | 3 min  | 1x     |
| Initial Denaturation | 95 °C       | 30 sec |        |
| Denaturation         | 95 °C       | 10 sec | 13x    |
| Annealing            | 55 °C       | 30 sec |        |
| Elongation           | 72 °C       | 1 min  |        |
| Final Elongation     | 72 °C       | 5 min  | 1x     |
| Store                | 8 °C        | ∞      |        |

51 Mix Index PCR with 50 µL **Clean-up Beads** (1/1 ratio)

 50 µL Clean-up Beads

52  00:05:00 binding of DNA onto the beads

53 Place on magnet until clear and discard supernatant

54 Wash twice with 150 µL 80% ethanol (while on magnet) and discard supernatant

 150 µL 80% ethanol (freshly prepared)

55  00:05:00 air dry beads

56 Elute cDNA in 20 µL UltraPure Water & transfer to new tube

 20 µL UltraPure Water

## Size selection

57 Load complete library onto an 2% Agarose E-Gel EX and run for 10 minutes.

 00:10:00

- 58 As soon as the Gel run has finished open the Gel framing using the Gel opening tool
- 59 Excise the Library from 300bp to 900bp using a clean scalpel
- 60 Gel purify the slice using the Qiagen MinElute Kit following manufacturer's guidelines:
- Add 450  $\mu$ L Buffer QG
  - Dissolve the gel slice in QG for 10 min @ 42 °C
  - Add 150  $\mu$ L Isopropanol to the sample and mix by inverting
  - Transfer sample to spin column and centrifuge at 16 000 x g for 1 min
  - Discard flow through and add 500  $\mu$ L Buffer QG
  - Centrifuge at 16 000 x g for 1 min and discard flow through
  - Add 700  $\mu$ L Buffer PE
  - Centrifuge at 16 000 x g for 1 min and discard flow through
  - Centrifuge again at 16 000 x g for 1 min to remove residual ethanol
  - Transfer column to a new 1.5 mL microcentrifuge tube
  - Add 20  $\mu$ L H<sub>2</sub>O to column and incubate for 1 min
  - Centrifuge at 16 000 x g for 1 min to elute and discard the spin column

#### Note

The Monarch DNA Gel Extraction Kit (NEB T1020L) can also be used.

## Library Quantification

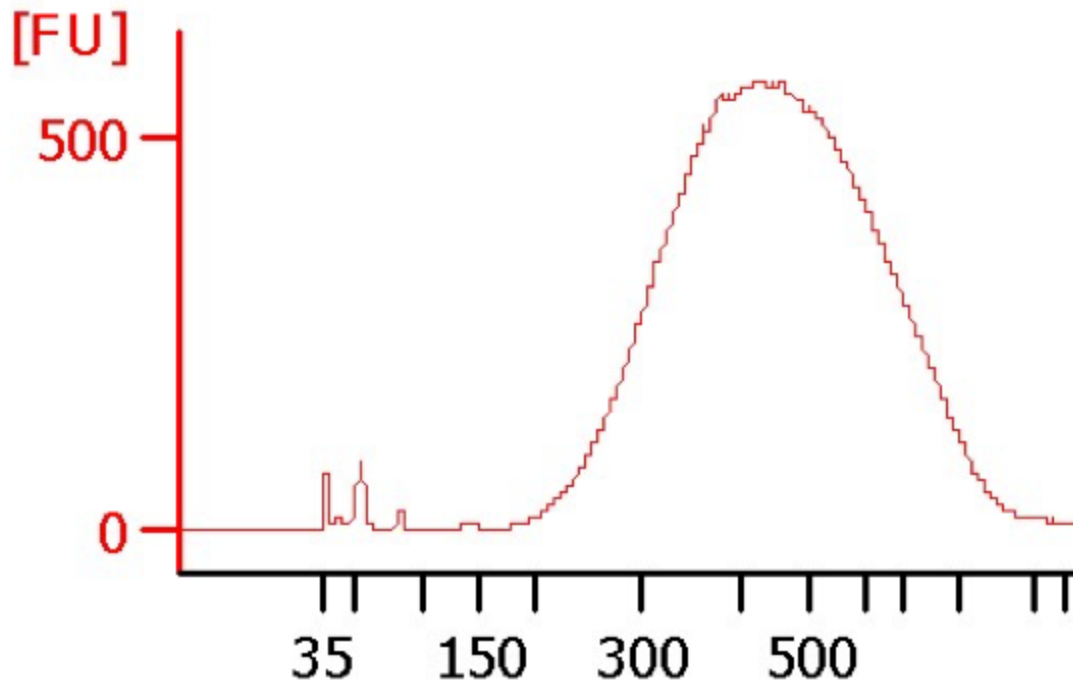
- 61 Quantify and quality control the library using the Agilent 2100 Bioanalyzer with High Sensitivity DNA Analysis Kits.
- Load both the library and a 1/10 dilution on two different lanes of the chip.

#### Note

If cDNA was not quality controlled previously, it is strongly suggested to run cDNA on the same chip.

### Expected result

Successful libraries will typically exceed 3-5 ng/ $\mu$ l concentration.



- 62 If pooling several libraries, combine equal molar amounts.

## Sequencing

- 63 Sequence your library on any compatible Illumina sequencer. Dilute libraries to recommended molarity according to Illumina's recommendations (eg. 2 nM). Select the following paired-end read-length settings:



|  | Read    | Cycles | Content            |
|--|---------|--------|--------------------|
|  | Read 1  | 16     | Cell barcode & UMI |
|  | Index 1 | 8      | i7 Index           |
|  | Index 2 | 0      |                    |
|  | Read 2  | 50     | cDNA fragment      |

## Primary data processing using zUMIs

64 Download and install zUMIs including all dependencies.

### Software

zUMIs

NAME

Linux

OS

<https://github.com/sdparekh/zUMIs>

REPOSITORY

<https://www.biorxiv.org/content/early/2017/10/18/153940>

SOURCE LINK

65 Copy the sequencing data from the sequencer and run bcl2fastq without demultiplexing.

### Command

```
bcl2fastq --use-bases-mask Y16,I8,Y50 --create-fastq-for-index-reads
```

66 Run zUMIs by specifying your parameters in the associated yaml file or using the **YAML config Rshiny application**. Below is an example yaml file for a typical gmcSCRB-seq run.

## Command

## Example zUMIs yamI

```
project: Example_project_name
sequence_files:
  file1:
    name: /data/R2.fq.gz
    base_definition: BC(1-8)
  file2:
    name: /data/R1.fq.gz
    base_definition:
      - BC(1-6)
      - UMI(7-16)
  file3:
    name: /data/R3.fq.gz
    base_definition: cDNA(1-50)
reference:
  STAR_index: /data/STAR5idx
  GTF_file: /data/Species.gtf
  additional_STAR_params: ''
  additional_files: /data/ERCC92.fa
out_dir: /data/zUMIs
num_threads: 15
mem_limit: 0
filter_cutoffs:
  BC_filter:
    num_bases: 1
    phred: 20
  UMI_filter:
    num_bases: 1
    phred: 20
barcodes:
  barcode_num: ~
  barcode_file: /data/bc.txt
  automatic: no
  BarcodeBinning: 0
  nReadsperCell: 100
counting_opts:
  introns: yes
  downsampling: '0'
  strand: 0
  Ham_Dist: 0
```



```
velocityto: no
primaryHit: yes
twoPass: yes
make_stats: yes
which_Stage: Filtering
Rscript_exec: Rscript
STAR_exec: STAR
pigz_exec: pigz
samtools_exec: samtools
zUMIs_directory: /data/zUMIs2/zUMIs
read_layout: SE
```