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# **DamID-seq protocol**

This protocol is based on the following published research:

Marshall OJ, Southall TD, Cheetham SW, Brand AH (2016) Nat Protocols 11:1586–1598

Marshall OJ, Brand AH (2015) Bioinformatics 31:3371-3

Southall TD, Gold KS, Egger B, Davidson CM, Caygill EE, Marshall OJ, Brand AH (2013) *Dev Cell* 26:101-12

Vogel MJ, Peric-Hupkes D, van Steensel B (2007) Nat Protoc 2:1467-1478.

# **Tissue preparation**

Aim to process enough material to give >10,000 driven cells

- 1. Take embryos, dissected tissue or heads from -80°C freezer (stored without buffer)
  - Add 75 μL sterile H<sub>2</sub>O + 20μL 500mM EDTA NB: Addition of EDTA is optional, but recommended when working with tissue containing the gut (or any tissue with high concentrations of nucleases/proteases). We have not tried lower concentrations than 100mM; lower concentrations may also work.
  - 3. **If working with embryos or heads**, use sterile (wash in 100% ethanol) pestle to homogenise samples (smush heads by rotating pestle between fingers and pressing down to bottom of tube)

# DNA extraction (Zymo Quick-DNA Miniprep plus kit) (2hrs) [Day 1]

The following assumes the use of a vacuum manifold. If using without a manifold, add in centrifugation steps as per the manufacturer's protocol.

- 1. Mix together 10 μL **Proteinase K** (Zymo Quick-DNA kit) + 95μL **Solid Tissue Buffer** (Zymo Quick-DNA kit; blue) per sample and vortex to make a **Prot K master mix**.
- Add 105µL of the Prot K master mix to each sample and flick gently to mix; digest at 56°C for 1-3 hours (heat block)
- 3. Cool to RT, add 400µL **Genomic Binding Buffer** (Zymo Quick-DNA kit) and mix by gentle inversion
- 4. Add all solution to a spin column on vacuum manifold and draw through
- 5. Add 400µL **DNA Pre-Wash Buffer** (Zymo kit) and draw through
- 6. Add 700 $\mu$ L g-DNA Wash Buffer (Zymo kit) and draw through
- 7. Add 200 $\mu L$  g-DNA Wash Buffer and draw through
- 8. Transfer column to a collection tube and spin at maximum speed in benchtop centrifuge for 2 mins to dry the column
- Transfer spin column to a new 1.5 ml tube, add 50 μL DNA Elution Buffer and leave at RT for at least 30 min. Spin at >6000 x g for 1 min and keep the flow-through
- 10. Optional: run 1 $\mu$ l on a 1% gel to check quality (should be a single band on the top of gel and not a smear)

# **DpnI digestion** (overnight + 30 mins)

- 1. Transfer 43.5µl of elution to a new 1.5ml tube
- Add 5µL CutSmart Buffer and 1.5µl DpnI, mix very gently and digest at 37°C overnight. (*This digestion step can be reduced to two hours if required.*) DpnI can be optionally heat-inactivated for 20 mins at 80°C. [Day 2]
- Clean-up with Machery-Nagel PCR Purification kit (100μL Buffer NTI; 2x 700μL washes); elute in 32μL DEPC H<sub>2</sub>O [Label tube "A"]

4. Transfer 15µL of sample into an 8-well PCR strip; store the remainder at -20°C as a backup. There is generally very little/undetectable amounts of DNA at this point, as uncut genomic DNA (which should be the majority of DNA in most use cases) will not pass through the spin columns – the purified DNA should only come from induced cells.

(If working with large amounts of tissue and/or if concerned about DNA shearing, the concentration of samples can be measured and a maximum of 750ng should be used going forward. However, such a high yield is highly unusual. Very low or undetectable yields at this stage produce excellent data – do not throw samples away!)

```
Adaptor ligation (2.5 hrs)
```

- 1. Add 4µL Adaptor Ligation Buffer (AL)
- 2. Add 1µL T4 DNA ligase and mix well
- 3. Ligate for 2 hours (16°C for 2 hours  $\rightarrow$  65°C for 10 mins in PCR machine)

# **DpnII digestion (2.5 hrs)**

- 1. Add 19µL DpnII Digestion Buffer (DIIB)
- 2. Add 1µL **DpnII enzyme** and mix well
- 3. Digest at 37°C for 2 hours
- 4. Heat inactivate at 65°C for 20 mins

PCR amplification (2.5 hrs + 30 mins)

- 1. Add 118µL PCR Buffer (PCR)
- 2. Add 2µL MyTaq polymerase and mix well
- 3. Split into 4 x 40µL reactions in a PCR strip using multichannel
- 4. Run PCR (use DamID 17 programme in PCR machine):

```
68°C 10 min
94°C 30 sec
65°C 5 min
68°C 15 min
Repeat 3x
94°C 30 sec
65°C 1 min
68°C 10 min
Repeat 17x
94°C 30 sec
65°C 1 min
68°C 2 min
68°C 5 min
4°C hold
```

This PCR protocol is from Vogel et al., 2007. Both the initial long extension for 10 mins and the subsequent long cycles appear to be required. A 3 min extension time, as present in the final 17 cycles, should amplify all fragments below 5kb; these represent 99.97% of all GATC fragments in the D. melanogaster genome.

(An argument could be made that fragments larger than 5kb will not represent genuine binding at that fragment, given Dam-fusion proteins typically only methylate within +/-2kb of their binding site in a seemingly normal distribution. On that basis, longer amplification times should probably be discouraged.)

- 5. PCR purify (160μL sample + 320μL Buffer NTI).
   [Day 3]

   If using MyTaq, wash column three times with kit's EtOH wash buffer before elution.
   Use

   32μL DEPC H<sub>2</sub>O to elute (leave water on column for at least 5 min) [Label tubes "C"]
- 6. Run 1 $\mu$ L on a gel to check quality (should be a smear between ~400bp and 2kb)
- 7. Spec DNA conc on nanodrop

# QC and sonication (>30 mins + overnight)

- 1. Dilute samples: 2µg DNA in 90µL DEPC H<sub>2</sub>O in 1.5mL TPX sonication tubes *Using dedicated sonication tubes is important to obtain consistent results*
- 2. Add 10µL CutSmart Buffer and mix well; cool on ice
- 3. Sonicate in Diagenode Bioruptor Nextgen:
  - a. Add 500mL ice to chamber, top up to the line with water and mix
  - b. Cool tube holder cassette on ice before using
  - c. Sonicate for 5 cycles (27 secs on, 27 secs off) on high power The aim is to achieve a ~300bp average fragment size. All sonicators are different; if using a different sonication device you will need to optimise the conditions first.
- 4. Check fragment size on Tapestation (genomic tape) ensure that average size is ~300bp Slight variations in fragment size are acceptable. Very large fragments (>600bp) may impede clustering efficiency and sequencing yields
- 5. Add 1µL AlwI enzyme and digest overnight at 37°C AlwI removes the DamID adaptors and initial GATC sequence (cuts at GGATCNNNN|N). Digestion can be carried out either before or after sonication; it makes no difference to the end result. AlwI cannot be heat inactivated, but will not cause issues if carried over in subsequent steps.
- 6. Transfer 70µL (or 90µL) of each sample into PCR 8-well strips for library prep

# Library preparation (~6 hrs)

[Day 4]

The following protocol uses homebrew reagents and is not compatible with Illumina kits. If you wish to use a commercial Illumina kit, we recommend the TruSeq Nano kit – follow the manufacturer's protocol except for the PCR step, in which case still use a reduced number of cycles (6 cycles seems optimum for most cases – see notes below). Commercial AmpureXP beads are a (vastly more expensive) alternative to homemade Seramag beads.

# Purify with Seramag beads (40 mins)

- Add 105μL of Seramag beads to 70μL sample (or 135μL to 90μL) and mix well Pipette mix 20x and then flick mix the strip and pulse down. Rapid mixing of beads and sample is important. Use 90μL of samples if concentration prior to sonication was low.
- 2. Incubate at RT, 10'
- 3. Place on magnetic stand 10' (or until clear)
- 4. Remove supernatant
- 5. Wash twice in 190µL 80% EtOH (30 secs wash time)
- 6. Let plate stand for 5' to air dry
- 7. Resuspend in 25µL Resuspension Buffer and remove from magnetic stand
- 8. Mix well and incubate 2' RT
- 9. Place on magnetic stand for 5' (or until clear)
- 10. Put 22.5 $\mu$ L of the supernatant into a new, clean tube for the next step

#### Adjust concentrations (30 mins)

- 1. Measure library conc on Qubit
- 2. Dilute samples to (no more than) 500ng of DNA in 20µL Resuspension Buffer

#### End repair (1 hr)

- 1. Add 7.5µL End Repair Buffer (ERB)
- 2. Add 2.5µL End Repair enzyme and mix well
- 3. Run PCR program "Truseq 1 2 ER Hi":
  - a. Incubate for 30 mins at 30°C
    - b. Heat inactivate enzymes for 20 mins at 75°C

#### Adenylate 3' Ends (40 mins)

- 1. Add 0.75µL Klenow 3' to 5' exo- enzyme and mix well
- 2. Run PCR program "Truseq 2 AT":
  - a. Incubate for 30 mins at 37°C
- 3. Proceed immediately to adaptor ligation

#### Adaptor ligation (20 mins)

- 1. Add 2.5µL NEB Quick Ligase enzyme
- 2. Add  $2.5\mu L$  of relevant adaptor
- 3. Run PCR program "Truseq 3 lig":
  - a. Incubate for 10 mins at 30°C
- 4. Add 5µL Stop Ligation Buffer

#### Clean-up with Seramag beads (35 mins)

- 1. Add 40µL Seramag beads and mix well
- 2. Incubate at RT, 10'
- 3. Place on magnetic stand 5' (or until clear)
- 4. Remove supernatant
- 5. Wash twice in 190µL 80% EtOH (30 secs wash time)
- 6. Let plate stand for 5' to air dry
- 7. Resuspend in 52.5µL Resuspension Buffer and remove from magnetic stand
- 8. Mix well and incubate 2' RT
- 9. Place on magnetic stand for 5' (or until clear)
- 10. Put  $50\mu L$  of the supernatant into a new, clean tube for the next step

#### Clean-up with Seramag beads again (35 mins)

- 1. Add 50µL Seramag beads and mix well
- 2. incubate at RT, 10'
- 3. Place on magnetic stand 5' (or until clear)
- 4. Remove supernatant
- 5. Wash twice in 190 $\mu$ L 80% EtOH (30 secs wash time)
- 6. Let plate stand for 5' to air dry
- 7. Resuspend in 22.5µL Resuspension Buffer and remove from magnetic stand
- 8. Mix well and incubate 2' RT
- 9. Place on magnetic stand for 5' (or until clear)
- 10. Put 20μL of the supernatant into a new, clean tube for the next step *This second clean-up is required to ensure complete removal of sequencing adaptor dimers* – *should these be present in the final library they will vastly out-compete genuine library sequences when hybridising to the flow cell.*

# Enrich DNA fragments (20 mins)

- 1. Add 5µL PCR Primer Cocktail
- 2. Add 25µL PCR Master Mix
- 3. Run PCR program "Truseq 4 PCR 6 cycles":

```
98°C for 30 secs
6 cycles of:
98°C for 10 secs
60°C for 30 secs
72°C for 30 secs
72°C for 5 mins
```

Note: aim to avoid concatamer formation (see troubleshooting section below)

# Clean-up with Seramag beads (final time) (35 mins)

- 1. Add  $50\mu L$  Seramag beads and mix well
- 2. incubate at RT, 10'
- 3. Place on magnetic stand 5' (or until clear)
- 4. Remove supernatant
- 5. Wash twice in 190µL 80% EtOH (30 secs wash time)
- 6. Let plate stand for 5' to air dry
- 7. Resuspend in 32.5µL Resuspension Buffer and remove from magnetic stand
- 8. Mix well and incubate 2' RT
- 9. Place on magnetic stand for 5' (or until clear)
- 10. Put  $30\mu L$  of the supernatant into a new clean tube

#### Library QC and multiplexing (1 hr)

- 1. QC sample with tapestation (use genomic tape) ensure that concatemers are not present (see "Troubleshooting" below) and that adapter dimers are not present (small peak around 240bp) in the sample. Record the average fragment length for each sample.
- 2. Measure DNA conc with Qubit (see attached protocol)
- 3. Calculate molarity of each sample from fragment length and concentration use the spreadsheet template (ask Owen for this), or see the "Using the Qbit" section for details as to how to calculate molarity manually
- 4. Pool samples to give 20nM final DNA conc (with all libraries at equal concentration) use the spreadsheet template to calculate this automatically
- 5. Sequence as single-end 50nt (SE50) on Illumina sequencer.

# **Bioinformatic processing of samples**

- Log<sub>2</sub>(Dam-fusion/Dam) binding profiles can be obtained from either raw reads in FASTQ format or aligned BAM files using the damidseq\_pipeline software, freely available from <a href="https://owenjm.github.io/damidseq\_pipeline">https://owenjm.github.io/damidseq\_pipeline</a>. Detailed usage instructions and examples are available from the website.
- 2. Profiles can be viewed using browser software such as IGV, as detailed on the website.
- 3. Calling expressed genes from RNA pol II ratio files generated by the damidseq\_pipeline software can be carried out using the polii.gene.call.r script, freely available from <a href="https://github.com/owenjm/polii.gene.call">https://github.com/owenjm/polii.gene.call</a>.

#### Notes

- Wear gloves, use filter tips and clean, autoclaved H<sub>2</sub>O to avoid contamination *Exceptions to filter tips: DNeasy/Qiagen kits before final elution Always use filter tips when preparing sequencing libraries The use of DEPC H<sub>2</sub>O throughout this protocol, while technically unnecessary, reflects the need to use a clean, contamination-free source of H<sub>2</sub>O*
- 2. Be very gentle with samples prior to DpnI digestion. Any genomic DNA shearing will result in broken ends that can potentially ligate DamID adapters. This will lead to random signals. Do not vortex, but mix samples gently by inversion and/or slow pipetting with a blue tip (and/or *very* gentle flicking). Always check the quality of the DNeasy-prepped DNA on a gel before continuing with the protocol.
- 3. For forward planning, each sample uses 278.5µL beads during the sequencing library prep

#### Citations

If you find this protocol useful, please cite <u>http://www.ncbi.nlm.nih.gov/pubmed/23792147</u> (original TaDa publication) and <u>http://www.ncbi.nlm.nih.gov/pubmed/26112292</u> (bioinformatic processing and NGS method); or <u>http://www.ncbi.nlm.nih.gov/pubmed/27490632</u> (Nature Protocols)

# **Reagents and equipment**

MyTaq HS DNA polymerase (Bioline, Cat# BIO-21112 (200µl) or BIO-21113 (500µl))

Note: it's been hard to find a replacement for the old Clontech Advantage cDNA polymerase. MyTaq HS seems the best so far; Clontech's Advantage 2 will work, but at reduced yield and is not recommended. Another, as yet untested, alternative would be Cesium Klentaq AC LA from DNA Polymerase Technology.

Seramag SpeedBead, 3 EDAC/PA5, 1µm, 15mL (Fisher Scientific, Cat# 12326433)

Note: Seramag beads need to be diluted in PEG before use as described in the dedicated section below. AmpureXP beads (which are ready to use and do not need further preparation) may be substituted for Seramag beads.

#### NEBNext® High-Fidelity 2X PCR Master Mix 1.25mL (NEB, Cat# M0541S)

*Alternative to NEBNext PCR MM (works equally well, but may be more expensive):* Kapa HiFi HS ReadyMix 1.25mL (Anachem, Cat# KK2601)

#### Quick Ligase kit (NEB, Cat# M2200S)

We're just using the Quick ligase enzyme, which is simply NEB T4 ligase at 2000 U/ $\mu$ L. However, at \$97 per 30 $\mu$ L, this is cheaper than high conc. NEB T4 ligase (#M0202T) at \$64 per 10 $\mu$ L.

#### NEB enzymes:

T4 ligase (NEB, Cat# M0202S) T4 DNA polymerase (NEB, Cat# M0203S) T4 polynucleotide kinase (NEB, Cat# M0201S) Klenow fragment (NEB, Cat# M0210S) Klenow 3' to 5' exo- (NEB, Cat# M0212L)

NEB restriction enzymes:

DpnI (Cat# R0176L) DpnII (Cat# R0543L) AlwI (Cat# R0513L)

dNTPs (NEB, Cat# N0447S)

# <u>Kits</u>

Zymo Quick-DNA miniprep plus (Cat# D4069)

Macherey-Nagel NucleoSpin Gel and PCR Clean-up (Cat# 740609.250)

Other equivalent kits also work; we find these to be the most effective in terms of price, performance and ease of use. Note that for genomic DNA extraction, we recommend a kit that allows efficient elution in  $50\mu L$ .

#### DamID buffers

Ligation buffer (AL)		x100	Aliquot
10x T4 ligase buffer	2	200	40
0.8uL ds ADR	0.8	80	
H2O	1.2	120	
DpnII digestion buffer (DIIB)		x100	Aliquot
Dpn II Buffer	4	400	190
H2O	15	1500	
PCR		x50	Aliquot
5X MyTaq HS Buffer	32	1600	1180
DamID-PCR primer (50 µM)	2.5	125	
DEPC H2O	83.5	4175	

#### Homebrew sequencing buffers

The following buffers and components are mostly adapted from Ethan Ford's protocol

End Repair Buffer (ERB)	1x (7.5μL)	Master mix x50
NEB T4 ligase buffer 10x	3.0	150
10mM dNTPs	1.2	60
DEPC H <sub>2</sub> O	3.3	165
Total	7.5	375
End Repair enzymes	1x (2.5μL)	Master mix x50
T4 DNA polymerase (NEB, 3U/µL)	1.14	56.82
Klenow Fragment (NEB, 5 U/µL)	0.23	11.36
T4 polynucleotide kinase (NEB, 10 U/µL)	1.14	56.82
Total	2.5	125.00

#### PCR Master Mix (PCR MM)

NEBNext 2x HiFi Hotstart Ready-mix (aliquot into 105µL aliquots and use as supplied)

**PCR Primer Cocktail (PPC)** 25 μL PCR1 primer (100 μM) 25 μL PCR2 primer (100 μM) 50 μL DEPC H<sub>2</sub>O

**Resuspension Buffer (RSB)** 10mM Tris-HCl, pH 8.0 0.1 mM EDTA

**Ligation Stop Solution** 0.5M EDTA

# Ligation of DamID adaptors

Making ds AdR stock (50uM):

- 1. Take 50  $\mu$ L AdRt (100uM) and 50  $\mu$ L AdRb (100 uM)
- 2. Incubate in removable metal heating block, 95°C, 2 mins
- 3. Remove heating block and allow to cool to room temperature (should take > 45mins)

# DamID oligos

AdRt CTAATACGACTCACTATAGGGCAGCGTGGTCGCGGCCGAGGA

AdRb TCCTCGGCCG

# Annealed structure:

5' CTAATACGACTCACTATAGGGCAGCGTGGTCGCGGGCCGAGGA 3'

# 3' GCCGGCTCCT 5'

# DamID\_PCR

GGTCGCGGCCGAGGATC

#### Sequencing adaptors and primer sequences

(adaptor barcodes highlighted in red; additional bases for higher indices in blue)

Universal	AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATC*T
Index 1	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACAGATCTCGGATCTCGGATGCCGTCTTCTGCTT*G
Index 2	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>CGATGT</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 3	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>TTAGGC</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 4	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>TGACCA</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 5	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>ACAGTG</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 6	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>GCCAAT</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 7	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>CAGATC</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 8	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>ACTTGA</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 9	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACGATCAGATCTCGTATGCCGTCTTCTGCTT*G
Index 10	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>TAGCTT</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 11	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACGGCTACATCTCGTATGCCGTCTTCTGCTT*G
Index 12	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>CTTGTA</b> ATCTCGTATGCCGTCTTCTGCTT*G
Index 13	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACAGTCAACAATCTCGTATGCCGTCTTCTGCTT*G
Index 14	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACAGTTCCGTATCTCGTATGCCGTCTTCTGCTT*G
Index 15	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACA <b>TGTCA</b> GAATCTCGTATGCCGTCTTCTGCTT*G
Index 16	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACCGGTCCCGGATCTCGTATGCCGTCTTCTGCTT*G
Index 18	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACGTCGCACATCTCGTATGCCGTCTTCTGCTT*G
Index 19	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>GTGAAA</b> CGATCTCGTATGCCGTCTTCTGCTT*G
Index 20	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>GTGGCC</b> TTATCTCGTATGCCGTCTTCTGCTT*G
Index 21	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACGTTTCGGAATCTCGTATGCCGTCTTCTGCTT*G
Index 22	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>CGTACG</b> TAATCTCGTATGCCGTCTTCTGCTT*G
Index 23	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACGAGTGGATATCTCGTATGCCGTCTTCTGCTT*G
Index 25	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCAC <b>ACTGAT</b> ATATCTCGTATGCCGTCTTCTGCTT*G
Index 27	/5Phos/GATCGGAAGAGCACACGTCTGAACTCCAGTCACATTCCTTTATCTCGTATGCCGTCTTCTGCTT*G
PCR1	AATGATACGGCGACCACCGA*G
PCR2	CAAGCAGAAGACGGCATACGA*G

/5Phos/ = 5' Phosphorylation; \* = Phosphorothioate linkages
We order adaptors from IDT @ 100 nmol synthesis + standard desalting

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#### **Annealing adaptors**

- 1. Resuspend adaptor oligos at 100µM in TE + 50mM NaCl
- 2. Mix  $25\mu$ L of relevant index +  $25\mu$ L Universal primer in PCR tube
- 3. Incubate in removable metal heating block, 95°C, 2 mins
- 4. Remove heating block and allow to cool to room temperature (should take > 45mins)

#### Notes re. adaptors

1. If multiplexing four or fewer libraries, selecting adaptors with barcodes that are too similar may result in a reduced number of reads passing the filter. In this case, the preferred indexes to use (in order) are 4, 7, 6 and 8.

# **Preparation of Seramag beads**

(*modified from* Rohland N and Reich D. (2012) Cost-effective, high-throughput DNA sequencing libraries for multiplexed target capture. *Genome Res* 22:939–46. doi:10.1101/gr.128124.111)

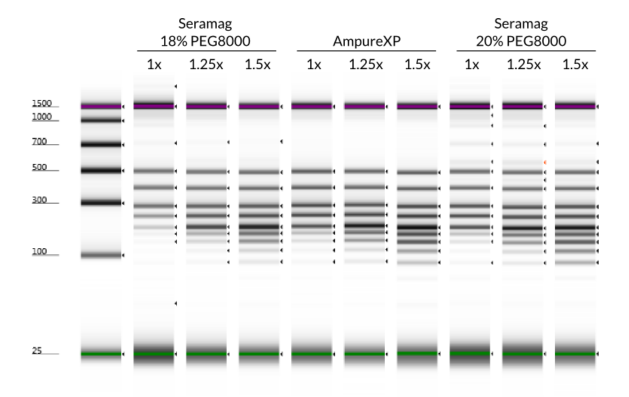
1. In 15mL Falcon tube add:

3g PEG-8000 (20% w/v final conc) 3mL 5M NaCl (1M final) 150μL 1M Tris pH 8.0 (10mM final) 30μL 0.5M EDTA (1mM final)

- 2. add DEPC  $H_2O$  to 14mL
- 3. mix by inversion until PEG is disolved (~5mins)
- 4. mix Speedbeads container well to ensure beads are in suspension
- 5. add  $300\mu L$  Seramag Speedbeads and mix by inversion
- 6. add DEPC H20 to 15mL volume and store at  $4^{\circ}$ C

#### Test beads by purifying Bioline Hyperladder V

- 1. Make mastermix of diluted ladder:  $10\mu$ L ladder +  $20\mu$ L DEPC H<sub>2</sub>O per sample
- 2. Purify ladder with 1x, 1.25x, 1.5x amounts of beads
- 3. Elute in  $20\mu$ L DEPC H<sub>2</sub>0
- 4. Run on tapestation using D1K tape (should look like 20% PEG-8000 lanes):



#### Notes

- 1. Rohland and Reich used 18% PEG-8000; however, this does not appear to perform as well as 20% in ladder purification (compared with AmpureXP)
- 2. Based on current prices:

Ampure beads are £689.67 for 60mL working solution Seramag beads are £317.62 for 15mL, which makes 750mL working solution Seramag beads are therefore more than 27x more cost-effective (!)

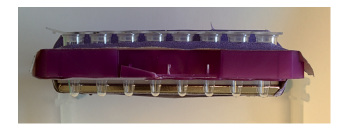
#### Optional: homemade magnetic rack for bead purification

Thermoscientific provide a 96-well magnetic plate for £763, which works reasonably well. However, the total retail cost of the neodymium magnets in this plate is  $\sim$ £6 (!)

Whilst it would be possible to buy individual ring magnets and glue them to a 96-well rack with epoxy resin, I've found it easier to make a simple 8-well (i.e. one PCR strip) magnetic plate using:

1x N42 Neodymium magnet (75x10x3mm) (e.g. <u>these magnets</u>)
1x 20μL filter tip box insert (acting as the 96-well support)
1x 75x10x2.5mm plastic spacer (supplied with a pair of the strip magnets)

The magnet together with spacer is taped to the tip box insert as shown below.



**Important:** as the magnet needs to remain firmly in contact with the edge of the strip tubes at all times, use a second PCR strip and/or tape on the side of the strip to hold the magnet in place.

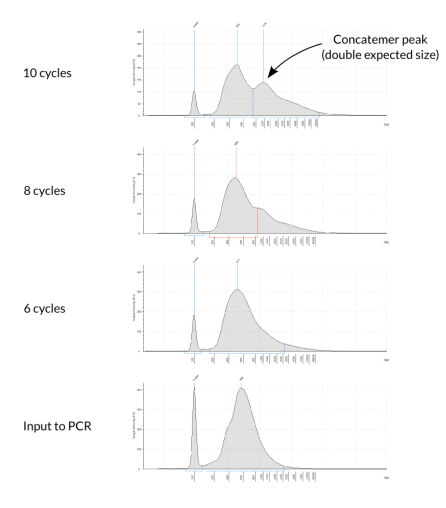
The advantage of using a strip magnet rather than a circular magnet is that the beads are pulled to one side of the tube rather than sitting in a ring, greatly facilitating liquid removal. (The disadvantage is that such a configuration is not compatible with 96-well high-throughput solutions).

Strength N42 seems to work well, but there should be no harm in using more powerful magnets if you can obtain them (the only result will be a faster purification process).

#### Troubleshooting

#### Secondary peak after PCR

This generally only happens using 8 or 10 (or more) cycles as PCR reaction is exhausted – a secondary peak twice the size of the original peak is seen on the tapestation genomic plots. This peak results from concatemers of the amplified product. If you see this, reduce number of PCR cycles and try again (6 cycles seems to work well for TaDa).



# Using the tapestation

- Prepare ladder and samples in 8-well PCR strips: 2μL pre-made ladder dilution in lane 1 3μL sample buffer + 1μL sample in subsequent lanes (Pre-made ladder dilution: ladder:loading-dye 1:10 or 1:15. Using 1:15 will save significant reagent costs but will obviously affect quantitation.)
- 2. Cap strip and mix well by vortexing (pulse down and mix several times)
- 3. Turn on tapestation and laptop
- 4. Launch Tapestation software and wait until prompted to insert the tape
- 5. Open tapestation, place tape in slot in correct orientation (barcode towards you on the right) We recommend using genomic tapes rather than D1K HS tapes – the genomic tapes give a better picture of adaptor-concatemer formation if present. D5K tapes may present an even better solution here, but we haven't tried these yet.
- 6. Insert strip with ladder and samples and close the machine
- 7. Select the wells you want to run on the software, and add labels in the table that appears
- 8. Click "Start" to begin the run (will take ~10 min to complete)
- 9. Once run is complete and analysis software is open, use the region settings to determine average fragment sizes and save a report

# Using the Qbit

- 1. Prepare samples as per the included protocol.
- 2. NB use  $1\mu$ L sample + 199 $\mu$ L working solution for each sample
- 3. With each sample, press the button to obtain the DNA concentration
- 4. DNA molarity  $\cong \frac{1,500}{fragment \ size \ in \ bp} \times [concentration \ in \ ng/\mu l]$
- 5. Use a spreadsheet template to calculate final pooling of samples

QUICK

CARD

REFERENCE

# Qubit<sup>™</sup> Assays

# www.invitrogen.com/qubit

**NOTE:** For best results, store the dye and the buffer at room temperature. Store the DNA, RNA, and protein standards at 4°C. Ensure that all assay reagents are at **room temperature** before you begin.

- 1. Set up two Assay Tubes for the standards (three for the protein assay) and one tube for each user sample.
- Prepare the Qubit<sup>™</sup> Working Solution by diluting the Qubit<sup>™</sup> reagent 1:200 in Qubit<sup>™</sup> buffer. Prepare 200 µL of Working Solution for each standard and sample.
- 3. Prepare the Assay Tubes\* according to the table below.

	<b>Standard</b> Assay Tubes	User Sample Assay Tubes
Volume of <b>Working Solution</b> (from step 2) to add	190 μL	180–199 μL
Volume of <b>Standard</b> (from kit) to add	10 μL	_
Volume of <b>User Sample</b> to add	_	1–20 μL
Total Volume in each Assay Tube	200 µL	200 µL

\* Use only thin-wall, clear 0.5 mL PCR tubes. Acceptable tubes include Qubit<sup>®</sup> assay tubes (set of 500, Cat. no. Q32856) or Axygen PCR-05-C tubes (VWR, part no. 10011-830).

- 4. Vortex all tubes for 2–3 seconds.
- Incubate the tubes for 2 minutes at room temperature (15 minutes for the Qubit<sup>™</sup> protein assay).
- 6. Insert the tubes in the Qubit<sup>®</sup> 2.0 Fluorometer and take readings. For detailed instructions, refer to the Qubit<sup>®</sup> 2.0 Fluorometer manual.
- 7. *Optional:* Using the Dilution Calculator feature of the Qubit<sup>®</sup> 2.0 Fluorometer, determine the stock concentration of your original sample.

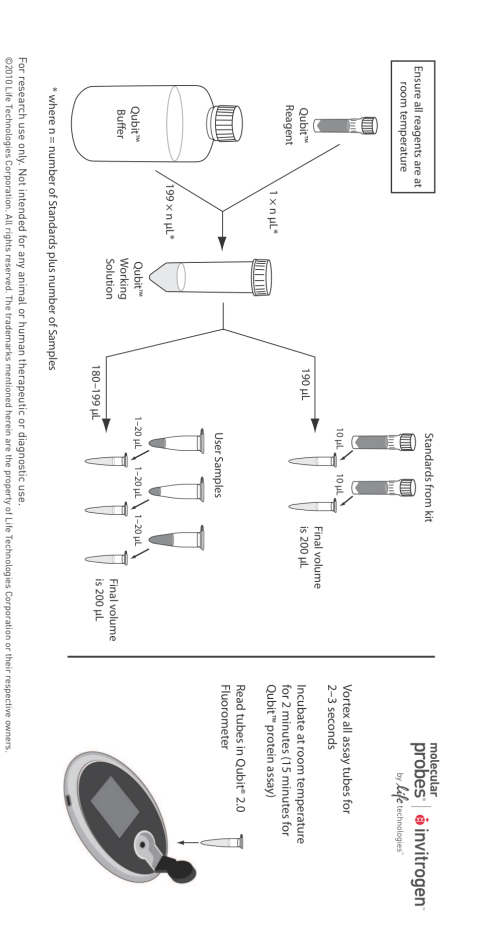
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Marshall Lab DamID-seq protocol