

Aug 14, 2019

Version 3

# O DASH Protocol V.3

DOI

dx.doi.org/10.17504/protocols.io.6ivhce6

Amy Lyden<sup>1</sup>, Emily Crawford<sup>1</sup>, Jenai Quan<sup>1</sup>, Saharai Caldera<sup>2</sup>, David Dynerman<sup>1</sup>

<sup>1</sup>CZ Biohub; <sup>2</sup>UCSF

Chan Zuckerberg Biohub



Amy Lyden

# Create & collaborate more with a free account

Edit and publish protocols, collaborate in communities, share insights through comments, and track progress with run records.

Create free account





DOI: https://dx.doi.org/10.17504/protocols.io.6ivhce6

Protocol Citation: Amy Lyden, Emily Crawford, Jenai Quan, Saharai Caldera, David Dynerman 2019. DASH Protocol. protocols.io <a href="https://dx.doi.org/10.17504/protocols.io.6ivhce6">https://dx.doi.org/10.17504/protocols.io.6ivhce6</a>

**Manuscript citation:** 



**License:** This is an open access protocol distributed under the terms of the **Creative Commons Attribution License**, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

Protocol status: Working

We use this protocol and it's working

Created: August 14, 2019

Last Modified: August 14, 2019

Protocol Integer ID: 26933

**Keywords:** CRISPR, sequencing, cas9, depletion, ribosomal, sequencing library, useful for rna, guide rna, human mitochondrial ribosomal rna, unique barcoding of the rna, abundant species such as human mitochondrial ribosomal rna, set of guide rna, seq of human metagenomic sample, human metagenomic sample, rna, crispr, dash treatment, sequencing space, rrna, ligation of adapter, dash protocol, hybridization, ligation, dash, seq library

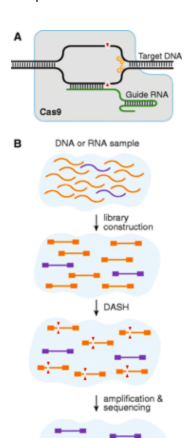


### **Abstract**

This protocol is for performing Depletion of Abundant Sequences by Hybridization (DASH) after preparing sequencing libraries and pooling together.

DASH is most useful for RNA-seq of human metagenomic samples, where abundant species such as human mitochondrial ribosomal RNAs (rRNAs) occupy a majority of the sequencing space available, leaving a minor fraction for regions of interest.

DASH treatment is performed after ligation of adapters and unique barcoding of the RNA-seq library. It employs CRISPR-Cas9 complexed to a set of guide RNAs (gRNAs) targeted to the abundant regions to be depleted in a given library. These abundant regions in the library are then cleaved, leaving only the fragments with intact adapters on both ends to be further amplified and sequenced.





# Guidelines

This protocol describes DASH for an RNA library prepared using the NEBNext Ultra II RNA Library Prep Kit. Other standard library preparation kits may be used instead, as long as DASH is applied after ligation of adapters.



### **Materials**

#### **MATERIALS**

- ₩ USER Enzyme 250 units New England Biolabs Catalog #M5505L
- X NEBNext Adaptor for Illumina New England Biolabs Catalog #E7337 in Kits E7335, E7500, E771
- **X** Thermocycler
- 🔯 NEBNext Ultra II RNA Library Prep Kit for Illumina 96 rxns New England Biolabs Catalog #E7770L
- Nuclease-free water Ambion Catalog #AM9932
- Qubit dsDNA HS kit Life Technologies Catalog #Q32851
- 🔯 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- TruSeq i7/i5 Indexing Primers Custom (or NEBNext® Multiplex Oligos for Illumina) **New England Biolabs Catalog #**E7500L
- **X** Cas9 40μM
- Dual guide RNAs (40μM targeted to genes or regions to be depleted crisprRNA and tracr RNA quantified by RNA Qubit)
- Proteinase K New England Biolabs Catalog #P8107S
- SPRI beads (homemade) or Ampure XP beads
- 🔯 Kapa HiFi Real-Time Amplification Kit Kapa Biosystems Catalog #KK2702
- 🔯 Zymo DNA Clean & Concentrator 5 **Zymo Research Catalog** #D4014
- Magnetic rack for PCR strips
- X gPCR machine
- Illumina P5 and P7 primers 5uM combined; P5: 5' AATGATACGGCGACCACCGAGATCT P7: 5' CAAGCAGAAGACGCATACGAGAT

### STEP MATERIALS

- X NEBNext Ultra II RNA Library Prep Kit for Illumina 96 rxns New England Biolabs Catalog #E7770L
- 🔀 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- Proteinase K New England Biolabs Catalog #P8107S
- SPRI beads (homemade) or Ampure XP beads
- 🔯 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies



- X Kapa HiFi Real-Time Amplification Kit Kapa Biosystems Catalog #KK2702
- Illumina P5 and P7 primers 5uM combined; P5: 5' AATGATACGGCGACCACCGAGATCT P7: 5' CAAGCAGAAGACGGCATACGAGAT
- SPRI beads (homemade) or Ampure XP beads
- Qubit dsDNA HS Assay kit Thermo Fisher Scientific Catalog #Q32854
- 🔀 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- **X** Cas9 40μM
- 🔯 10X Cas9 Activity Buffer (500mM Tris pH 8.0 1M NaCI 100mM MgCL2 10mM TCEP)
- Dual guide RNAs (40μM targeted to genes or regions to be depleted crisprRNA and tracr RNA quantified by RNA Qubit)



## **Protocol materials**

- Proteinase K New England Biolabs Catalog #P8107S
- TruSeq i7/i5 Indexing Primers Custom (or NEBNext® Multiplex Oligos for Illumina) New England
- X Kapa HiFi Real-Time Amplification Kit Kapa Biosystems Catalog #KK2702
- 🔯 Zymo DNA Clean & Concentrator 5 **Zymo Research Catalog #**D4014
- Illumina P5 and P7 primers 5uM combined; P5: 5' AATGATACGGCGACCACCGAGATCT P7: 5' CAAGCAGAAGACGGCATACGAGAT
- X Zymo DNA Clean & Concentrator 5 **Zymo Research Catalog** #D4014
- 🔯 USER Enzyme 250 units New England Biolabs Catalog #M5505L
- 🔀 NEBNext Ultra II RNA Library Prep Kit for Illumina 96 rxns New England Biolabs Catalog #E7770L
- 🔯 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- Dual guide RNAs (40µM targeted to genes or regions to be depleted crisprRNA and tracr RNA quantified by RNA Qubit)
- Dual guide RNAs (40μM targeted to genes or regions to be depleted crisprRNA and tracr RNA quantified by RNA Qubit)
- Magnetic rack for PCR strips
- Marmocycler 1
- SPRI beads (homemade) or Ampure XP beads
- 🔯 NEBNext Ultra II RNA Library Prep Kit for Illumina 96 rxns New England Biolabs Catalog #E7770L
- Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- 🔯 10X Cas9 Activity Buffer (500mM Tris pH 8.0 1M NaCl 100mM MgCL2 10mM TCEP)
- SPRI beads (homemade) or Ampure XP beads
- SPRI beads (homemade) or Ampure XP beads
- Qubit dsDNA HS kit Life Technologies Catalog #Q32851
- 🔀 NEBNext Adaptor for Illumina New England Biolabs Catalog #E7337 in Kits E7335, E7500, E771
- Qubit dsDNA HS Assay Kit **Thermo Fisher Scientific Catalog** #Q32851
- Illumina P5 and P7 primers 5uM combined; P5: 5' AATGATACGGCGACCACCGAGATCT P7: 5' CAAGCAGAAGACGGCATACGAGAT
- **X** Cas9 40μM
- **X** qPCR machine
- 🔯 Qubit dsDNA HS Assay kit Thermo Fisher Scientific Catalog #Q32854



- Nuclease-free water **Ambion Catalog #**AM9932
- Qubit dsDNA HS kit Life Technologies Catalog #Q32851
- Kapa HiFi Real-Time Amplification Kit Kapa Biosystems Catalog #KK2702
- **X** Cas9 40μM
- Proteinase K New England Biolabs Catalog #P8107S
- 🔯 10X Cas9 Activity Buffer (500mM Tris pH 8.0 1M NaCI 100mM MgCL2 10mM TCEP)
- 🔯 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- 🔯 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- 🔀 NEBNext Ultra II RNA Library Prep Kit for Illumina 96 rxns New England Biolabs Catalog #E7770L
- Qubit dsDNA HS Assay Kit **Thermo Fisher Scientific Catalog** #Q32851
- 🔯 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- **X** Cas9 40μM
- 🔯 10X Cas9 Activity Buffer (500mM Tris pH 8.0 1M NaCl 100mM MqCL2 10mM TCEP)
- Dual guide RNAs (40μM targeted to genes or regions to be depleted crisprRNA and tracr RNA quantified by RNA Qubit)
- X Zymo DNA Clean & Concentrator 5 **Zymo Research Catalog #**D4014
- Proteinase K New England Biolabs Catalog #P8107S
- SPRI beads (homemade) or Ampure XP beads
- 🔯 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- Qubit dsDNA HS kit Life Technologies Catalog #Q32851
- Illumina P5 and P7 primers 5uM combined; P5: 5' AATGATACGGCGACCACCGAGATCT P7: 5' CAAGCAGAAGACGGCATACGAGAT
- Kapa HiFi Real-Time Amplification Kit Kapa Biosystems Catalog #KK2702
- SPRI beads (homemade) or Ampure XP beads
- Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies
- Qubit dsDNA HS Assay kit Thermo Fisher Scientific Catalog #Q32854

# Troubleshooting



## Before start

Prepare your sequencing libraries, and make sure you have gRNAs for DASH. We used the DASHit software to generate an optimized set of guide RNA sequences based on a dataset of unwanted genes. (For DASHit software, visit: <a href="https://github.com/czbiohub/guide\_design\_tools/blob/master/dashit/dashit-reads/dashit-reads.org">https://github.com/czbiohub/guide\_design\_tools/blob/master/dashit/dashit-reads.org</a>. You can buy the gRNAs or buy DNA templates and transcribe them. For an IVT protocol for crispr RNA and tracr RNA, please contact <a href="mailto:emily.crawford@czbiohub.org">emily.crawford@czbiohub.org</a> or <a href="mailto:amy.lyden@czbiohub.org">amy.lyden@czbiohub.org</a>)



# Prepare indexed RNA-seq library

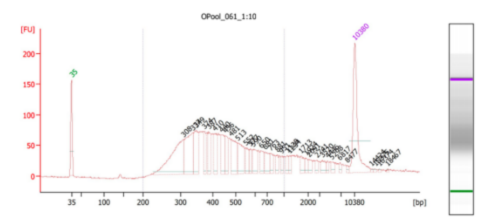
- 1 Follow all steps using NEB Ultra II RNA library preparation kit. Use an input RNA volume of 25ng if available, or less if not available. Perform 12-18 cycles of indexing PCR.
  - NEBNext Ultra II RNA Library Prep Kit for Illumina 96 rxns **New England Biolabs Catalog #**E7770L

## 2 Choose one option:

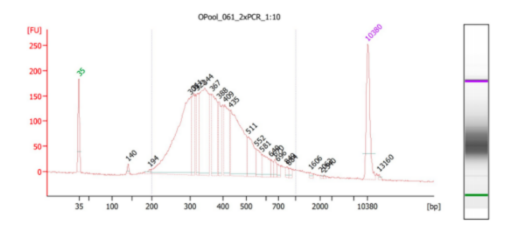
- **a. Pooled DASH:** If there are multiple samples, you may pool them at normalized concentrations of DNA between 200 1000 bp by Qubit or BioA/Tapestation/Fragment Analyzer, or using a preliminary low-depth sequencing run (such as an iSeq or MiSeq) of an equivolume pool to determine pooling ratios. Alternatively, if you suspect that your samples may vary drastically in DASHable material, you can pool equivolumes of your samples and DASH, then run an iSeq or MiSeq sequencing. Then pool normalized concentrations based on these values. This pooled library will go into a single DASH reaction.
- **b. Single Sample DASH:** Additionally if you suspect your samples may vary in DASHable material or have a high amount of DASHable material, you may DASH each sample individually and pool them after. In this case, each sample will have its own DASH reaction, and the following steps will be performed on each individual library.
- 3 Quantify by HS DNA Qubit.
  - 🔯 Qubit dsDNA HS Assay Kit Thermo Fisher Scientific Catalog #Q32851
- 4 Perform a HS DNA Bioanalyzer (or other fragment analysis) and ensure for your library(ies):



- 1) that adapter dimers have been removed. Several SPRI bead clean-ups at a sample:bead volume ratio of 1:1 may be necessary.
- 2) there is sufficient sample in between the 200 1000 bp range.
- 3) There is no 'PCR bubble' (appears as secondary bump in fragment analysis trace at higher bp regions, see trace below). PCR bubble will be eliminated with 1-2 extra PCR cycles. We recommend.
- 100ng of DNA in 23µL H20
- 25μL Kapa HiFi Real-Time Amplification Master Mix
- 2μL Illumina primers at 5μM.



Library pre-PCR cycles with DNA in 2000-10000bp region. This library may be over-amplified and cannot be used in DASH.



Library post 2xPCR cycles with no DNA in 2000-10000bp region. This confirms library previously had PCR bubble. This library can be used in DASH.

🔀 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies



5 Each DASH reaction requires at least 10 µL of 2.8 nM DNA library (approximately 0.83 ng/μL for an average 450 bp library). You may concentrate your sample by SPRI bead clean-up if necessary.

### DASH

- 6 Thaw necessary reagents and let come to room temperature before use. If you will not be using for more than 10 minutes, put on ice and take off 5 minutes prior to mixing.
  - **X** Cas9 40μM
  - ₩ TCEP) 10X Cas9 Activity Buffer (500mM Tris pH 8.0 - 1M NaCl - 100mM MgCL2 - 10mM
  - Dual guide RNAs (40μM targeted to genes or regions to be depleted crisprRNA and tracr RNA - quantified by RNA Qubit)
- 7 Prepare master mix (MM) of Cas9 and gRNAs:

#### Note

Ensure that the dual-guide RNAs have been annealed at 95°C for 30 seconds and then cooled to room temperature. No additional heating is recommended if pairs were annealed prior to storage.

#### Note

Cas9-gRNAs MM will be replenished once in the protocol. Ensure you have enough reagents for 2X MM for each sample.

Reagent	Stoc k Con cent ratio n	Fina I Con cent ratio n	1X cas 9- gRN A MM
10X cas9 buffer	10X	1.25 X	2.5μ L
cas9	40μ Μ	5μΜ	2.5μ L



dual-guide RNAs	40μ Μ	10μ Μ	5μL
			10μL total

**DASH Master Mix** 

- 8 Mix well by gently pipetting and tapping and incubate this mixture at \$\mathbb{\mathbb ♦ 00:05:00 .
- 9 Add  $\perp$  10  $\mu$ L of a 2.8 nM barcoded DNA library (pooled or not) to every  $\perp$  10  $\mu$ L of the above Cas9-gRNA mix.
  - o This reaction may be scaled up from the original 1X (  $\perp$  20  $\mu$ L DASH reaction)
  - o Calculator for µg moles: <a href="https://www.promega.com/a/apps/biomath/">https://www.promega.com/a/apps/biomath/</a>

Rea gent	Stoc k Con cent ratio n	Fina I Con cent ratio n	1X cas 9- gRN A MM
10X cas9 buff er	10X	1.25 X	2.5μ L
cas9	40μ Μ	5μΜ	2.5μ L
dual - guid e RNA s	40μ Μ	10μ Μ	5μL
DNA libra ry	2.8n M	1.4n M	10μL
			20µ L total

Final DASH reaction (1X, 20µL)



- 10 Incubate the DASH reaction at \$\mathbb{8}\$ 37 °C for \( \blue{\cdot} \) 00:30:00 .
- Before your incubation is up, prepare a second batch of 1X Cas9-gRNA MM for all your samples and incubate mixture at \$\mathbb{s}\$ 37 °C for \$\infty\$ 00:05:00 .

X Zymo DNA Clean & Concentrator - 5 Zymo Research Catalog #D4014

- 14 Incubate the DASH reaction at \$\mathbb{8}\$ 37 °C for \( \mathbb{O} \) 01:30:00 .

#### Note

Total incubation of DNA library in Cas9-gRNA MM should be 02:00:00 . Column clean-up timing has not been optimized, and could be split to be 37 °C for 01:00:00 , column clean, and then 37 °C for 01:00:00 .



- Add  $\Delta$  1  $\mu$ L proteinase K to each  $\Delta$  20  $\mu$ L reaction and mix well by gently tapping or pipetting.
  - Proteinase K New England Biolabs Catalog #P8107S
- Incubate at \$50 °C for 00:15:00 for cas9 inactivation.

# Clean up DASH reaction

17 Prepare to do a SPRI selection of sample:bead 1:0.9 (equivalent to Ampure 0.7X)

### Note

This goal of this step is to purify your target library away from the DASH reagents (buffers, Cas9, gRNAs) and the DASH-digested fragments. Depending on DASH efficiency and the size of the expected digested fragments, this ratio may be altered to best remove the fragments.

#### Note

Other magnetic beads such as AmpureXP may be used instead of SPRI. However, please take note of the different size cut-offs for sample:bead ratios, as they may vary from the homebrew SPRI beads used in this protocol.

- 18 Equilibrate SPRI beads to room temperature and vortex well to mix.
  - 🔀 SPRI beads (homemade) or Ampure XP beads
- Add beads equivalent to 0.9X the sample volume to each sample tube (for  $\Delta$  21  $\mu$ L of sample, add  $\Delta$  18.9  $\mu$ L beads).
- 20 Mix well by pipetting or tapping the tubes and spin down briefly.

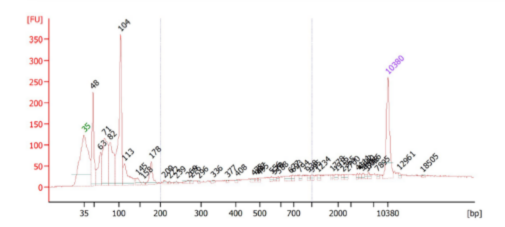


- 21 Incubate for 00:05:00 at room temperature, then put the tubes on the magnetic rack. Allow beads to separate on the magnet for 3-5 minutes, or until the supernatant is clear.
- 22 Keeping the tubes on the magnet, carefully remove and discard the supernatant.
- Add Δ 200 μL 80% ethanol (prepared fresh). Incubate beads for 00:01:00 and then remove the ethanol.
- 24 Repeat the above ethanol wash step.
- Allow the beads to air dry for approximately 00:05:00. Do not over-dry. Dry beads should appear matte (rather than glossy) but should not have a cracked appearance. Overdried beads may not resuspend or elute well.
- Remove tubes from magnet and add Δ 27 μL nuclease-free H2O.
- 27 Resuspend well by vortexing, tapping the tubes, and spinning down briefly.
- Allow 00:02:00 for DNA to elute from beads, then transfer tubes back to magnet
- Allow the beads to separate on magnet.
- 30 Collect  $\[ \underline{A} \]$  25  $\mu L$  of supernatant from each sample and transfer it to a clean PCR tube.
- Run a fragment analysis, such as with the HS dsDNA Bioanalyzer kit or the Agilent HS D5000 Tapestation kit.
  - Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies



### **Expected result**

If DASH is successful, the trace may be characterized by the presence of small peaks between 30 – 100 bp. The size of these peaks will depend on the proportion of expected depleted product over target product, and how closely spaced together the target guide RNAs were designed.



DASHed sample showing the digested fragments and gRNAs (30-140), an adapter dimer (178), and low concentration of target library (sloping from 300 – 1000+). Bioanalyzer internal markers in green (35) and purple (10380). This sample needs additional cleanup and amplification.

32 If desired, also quantify by HS DNA Qubit.

**☒** Qubit dsDNA HS kit **Life Technologies Catalog** #Q32851

As needed, perform additional clean-up steps to remove the digested fragments and adapter dimers in your sample. Proceed to amplification of target library if your concentration in the 200-1000bp region is very low, or looks nonexistent.

# **Amplification of Target Library**

Perform this Real-Time PCR amplification step if after cleaning out the digested fragments and adapter dimers your library concentration is too low for loading onto a sequencer.



#### Note

This step could be optional, if your DASHable material is low and your concentration after DASH is high. However, we have found it to be almost always necessary.

- Thaw Kapa HiFi Real-Time Amplification Master Mix and Standard 2 on ice, plus Illumina P5 and P7 primers
  - XX Kapa HiFi Real-Time Amplification Kit Kapa Biosystems Catalog #KK2702
  - Illumina P5 and P7 primers 5uM combined; P5: 5'

    AATGATACGGCGACCACCGAGATCT P7: 5' CAAGCAGAAGACGGCATACGAGAT
- 36 Prepare the following reaction for each library:

Reagent	Volu me (1X)
2X Kapa HiFi Real-time Amplification Master Mix	25μL
5μM Illumina P5 and P7 primers	2μL
DNA (bring up volume with H20 if needed)	23μ L
	50μ L

- Place each sample in a tube or strip of tubes that is physically separated from the others, so that they can be removed one at a time.
- 38 Put  $\stackrel{\text{\@textbf{A}}}{=}$  50  $\mu$ L of Standard 2 in a PCR tube
- Cycle with the following conditions in an RT-PCR machine. Make sure that if your machine does a baseline correction, that you remove it.

Tem pera ture	Tim e	Cycl es
---------------------	----------	------------



98C	45 sec	1
98C	15 sec	25
63C	30 sec	
72C	1m 45 sec	
Plate read		
72C		

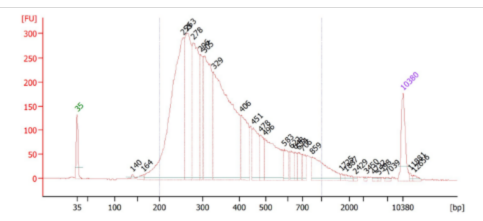
Cycling conditions for PCR

- Watch the qPCR graph during the plate read. When a sample reaches the fluorescence threshold given by standard 2, remove that sample **DURING the 72°C for 20 second timeframe** after the plate read.
- Perform a SPRI cleanup as described in `Clean up DASH reaction` section above at a sample:bead ratio of 1:0.9 to remove amplification reagents, and elute in a suitable volume of water. 

  go to step #17
  - SPRI beads (homemade) or Ampure XP beads
- 42 Quantify and analyze your library by HS DNA Qubit and HS dsDNA Bioanalyzer.
  - **Q** Qubit dsDNA HS Assay kit Thermo Fisher Scientific Catalog #Q32854
  - 🔀 Bioanalyzer chips and reagents (DNA High Sensitivity kit) Agilent Technologies

### **Expected result**

This trace should resemble a regular library with a majority of the library sized between 200 – 1000bp:



Same DASHed sample from above.SPRI cleaned at sample:bead ratio 1:0.7, 2 amplification cycles. Small primer dimer at 140bp.

- 43 Perform more SPRI clean-ups or amplification if necessary.
- 44 Submit your library for sequencing!