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In vitro transcription of crRNA and tracrRNA from DNA oligos for cas9 enrichment and nanopore sequencing (for Bac - PULCE)

Forked from In Vitro Transcription for dgRNA

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SUBMIT TO PLOS ONE

ABSTRACT

For CRISPR-cas9 protocols including BAC-PULCE, T7 polymerase is used to transcribe the crRNA and tracrRNA to make dgRNA for cas9. The two components of the dual guides are the crRNA (containing your variable 20 nt target plus a 22 nt constant region) and the tracrRNA (a 72 nt constant region).

We used a combination of these protocols:

https://www.protocols.io/view/in-vitro-transcription-for-dgrna-3bpgimn

/dx.doi.org/10.17504/protocols.io.3bpgimn

https://international.neb.com/protocols/2013/04/02/standard-rna-synthesis-e2050

Nanopore protocol Cas-mediated PCR-free enrichment - please refer to this protocol in particular for further important detail around sequencing the DNA library.

Using modified "In Vitro Transcription for dgRNA V.2" (Lyden et al 2019):

In this protocol, we are designing crRNAs against which Cas9 will target. Using this sequence, we will add a T7 RNA polymerase binding site at the 5' end, a 3' tracrRNA binding site, and use the reverse complement of this to order our DNA oligos (below more details):

Designing crRNAs using CHOPCHOPv3 (Labun et al 2019), insert your gene target and CHOPCHOP will find a sequence of 20nt's that will end in an 'NGG-3'

5'----NNNNNNNNNNNNNNNNNNNNNNNNNGG----3'

where 20Ns are your target site. Cas9 cuts between the 17th and 18th N of your target.

The sequence of each crRNA should be as follows, with the Ns replaced by your 20 nt target:

The underline portion is the T7 RNA polymerase binding site. The addition at the 3' end represents the region to which an 18mer will bind, in order to allow T7 a double stranded binding site.

18mer T7:

TAATACGACTCACTATAG

DOI

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PROTOCOL CITATION Olin Silander 2021. In vitro transcription of crRNA and tracrRNA from DNA oligos for cas9 enrichment and nanopore sequencing (for Bac - PULCE) . protocols.io https://dx.doi.org/10.17504/protocols.io.brbnm2me 6 FORK NOTE FORK FROM Forked from In Vitro Transcription for dgRNA, Amy Lyden **KEYWORDS** DASH, FLASH, cas9, dgRNA, IVT, transcription, Bac PULCE 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 CREATED Jan 10, 2021 LAST MODIFIED Feb 23, 2021 PROTOCOL INTEGER ID 46158 **GUIDELINES** Work in an RNAse free space! we used RNAse ZAP on all work surfaces prior to carrying out method. use nuclease free tubes lobind eppendorf tubes. MATERIALS TEXT **MATERIALS** Nuclease-free water Ambion Catalog #AM9932 **⊠** Qubit RNA HS Assay Kit **Thermo Fisher** Scientific Catalog #Q32852 SPRI beads (homemade) or Ampure XP beads Contributed by users ★ tracrRNA template (90nt) IDT
 Step 4 8 10X T7 Buffer (400 mM Tris pH 7.9 - 200 mM MgCl2 - 50 mM DTT - 20 mM spermidine (Sigma 85558)) store at -80C Contributed by users Scientific Catalog #R0481

 Magnetic Tube Rack for 1.5mL or 15mL tubes Contributed by users Catalog #12321D

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STEP MATERIALS

Citation: Olin Silander (02/23/2021). In vitro transcription of crRNA and tracrRNA from DNA oligos for cas9 enrichment and nanopore sequencing (for Bac - PULCE) ÃÂ . https://dx.doi.org/10.17504/protocols.io.brbnm2me

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ScrRNA template (60nt) IDT

    ★ tracrRNA template (90nt) IDT

Scientific Catalog #ND-1000
Scientific Catalog #R0481
8 10X T7 Buffer (400 mM Tris pH 7.9 - 200 mM MgCl2 - 50 mM DTT - 20 mM spermidine (Sigma 85558)) store at -80C Contributed by users ■
Step 2

    ⋈ Nuclease-free water

Ambion Catalog #AM9932

    Nuclease-free water

Ambion Catalog #AM9932
SPRI beads (homemade) or Ampure XP beads Contributed by users

    ⊠ Ethanol 100% Contributed by users
    Step 11
Qubit RNA HS Assay Kit Thermo Fisher
Scientific Catalog #Q32852
users Catalog #5067-1548
■ NTP quality varies from one vendor to another. We have had consistent success with Thermo cat # r0481 and Life Tech
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- NTP quality varies from one vendor to another. We have had consistent success with Thermo cat # r0481 and Life Tech
 AM81110G, -20G, -30G, and -40G, used at a final concentration of 1 mM each
- We purify our own T7, and experiments should be optimized for each batch of T7, or for a commercial T7.

BEFORE STARTING

In this protocol, we are designing crRNAs against which Cas9 system will target. Using this sequence, we will add a T7 RNA polymerase binding site at the 5' end, a 3' tracrRNA binding site, and use the reverse complement of this to order our DNA oligos, with these dgRNAs we are able to cut and sequence regions of interest of a desired sample (below more details):

Designing crRNAs using CHOPCHOPv3 (Labun et al 2019, https://chopchop.cbu.uib.no), insert your gene target and CHOPCHOP will find a sequence of 20nt's that will end in an 'NGG-3' (PAM site)

5'---NNNNNNNNNNNNNNNNNNNGG----3'

where 20Ns are your target site. Cas9 cuts between the 17th and 18th N of your target.

The sequence of each crRNA should be as follows, with the Ns replaced by your 20 nt target:

The underline portion is the T7 RNA polymerase binding site. The addition at the 3' end represents the region to which an 18mer will bind, in order to allow T7 a double stranded binding site.

18mer T7:

TAATACGACTCACTATAG

further details from the FLASH protocol to help in designing your sequences -

Designing the crRNA(s): (contains your target sequence) S. pyogenes cas9 requires a 20-nt target directly 5' to a PAM motif "NGG" (where N is any nucleotide). The NGG is not present in the guide RNA itself. So when choosing a target you are looking for a sequence that matches the following pattern (and don't forget that you can target either strand): where the 20 Ns in bold are your target site. Cas9 will cut between the 17th and 18th nt of the target, yielding the following products: 5'----NNNNNNNNNNNNNNNN3' 5'NNNNGG----3' ☐ The sequence of each crRNA should be as follows, with the Ns replaced by your 20 nt target: The underlined portion is the T7 transcription site. T7 only requires its own 18 nt binding site to be doublestranded; the rest of the template can be single stranded. Thus the template can be constructed by purchasing two oligos from IDT: the reverse complement of the 60 nt sequence listed above, plus an 18 nt oligo to make the T7 site double stranded: 60mer reverse complement: 18mer T7: **TAATACGACTCACTATAG** The tracrRNA: (constant for all dgRNA) ☐ The sequence of the tracrRNA template should be as follows: <u>TAATACGACTCACTATAG</u>GACAGCATAGCAAGTTAAAATAAGGCTAGTCCGTTATCAACTTGAAAAAGTGGCACCGAG **TCGGTGCTTTTT** Just as with the crRNA, only the T7 binding site needs to be double stranded, so the following two oligos can be purchased from IDT: 90mer reverse complement: AAAAAGCACCGACTCGGTGCCACTTTTTCAAGTTGATAACGGACTAGCCTTATTTTAACTTGCTATGCT **GTCCTATAGTGAGTCGTATTA** 18mer T7: **TAATACGACTCACTATAG** RNAse Zap equipment, gloves and bench prior to starting protocol. Turn on heat block to reach a temperature of 8 95 °C . Annealing T7 to crRNA and tracrRNA template Pool your crRNA DNA oligo sequences in equimolar amounts. We order our crRNA templates from IDT with oligos diluted in nuclease free water at a concentration of 10µM in 1.5ml Eppendorf tubes. ScrRNA template (60nt) IDT **⊠** Nuclease-free water

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Preparation

Ambion Catalog #AM9932

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3 Add an equimolar amount of T7 primer to your crRNA pool. For example, reconstitute T7 primer to 10μM, and pool 125μL of your 10μM crRNA pool with 125μL of your T7 primer at 10μM.

4 Add an equimolar amount of T7 primer to your tracrRNA. For example, if you have reconstituted your tracrRNA to 100μM, pool 125μL of your tracrRNA at 100μM to 125μL of your T7 primer at 100μM.

StracrRNA template (90nt) IDT ■

Anneal tracrRNA template + T7 primer and crRNA template + T7 primer by heating to § 95 °C on a heat block or thermocycler for © 00:02:00 and allowing them to cool to room temperature slowly on the bench in a tube rack.

In Vitro Transcription

6 For IVT we used the T7 E2050S kit from NEB and followed the standard RNA synthesis method. Thaw and keep on ice the necessary components - NTP buffer mix.

₩ HiScribe T7 Quick High Yield RNA Synthesis Kit - 50 rxns New England

Biolabs Catalog #E2050S

7 Assemble the reactions in the following order in nuclease free 1.5ml lobbind tubes for both the crRNA & tracrRNA . total volume can be scaled accordingly. We usually had a total reaction volume of 90μL.

Α	В	С
1x	CrRNA	TracrRNA
Nuclease free water	ΧμL	XμL
NTP buffer mix	30µ	30µ
Oligo RNA template	(1µg)	(1µg)
T7 Polymerase Mix	бμL	6µL
Total Volume Mix	90µL	90µL

^{*} To find the volume of oligo RNA template required we used NEBio calculator ss: moles to mass (https://nebiocalculator.neb.com/#/ssdnaamt)

8 Incubate at § 37 °C overnight. We usually incubate for 16 hours using a heat block.

RNA Purification with SPRI beads

9 Proceed to purification. We used Ampure beads to purify gRNAs after transcription.

SPRI beads (homemade) or Ampure XP beads Contributed by users

10 Equilibrate Ampure XP beads to room temperature.

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11	For every 200 μ L of IVT reaction, add 300 μ L of 100% ethanol (1:1.5 ratio i.e 90 μ l IVT reaction mixture then add 135 μ l of ethanol). The solution should turn a cloudy white (precipitation of RNA) upon addition of ethanol. This step helps the short RNAs bind to the beads. This can be done in several 1.5mL tubes.
	⊠Ethanol 100% Contributed by users
12	For every $500\mu L$ of IVT reaction + 100% ethanol, add $500\mu L$ of Ampure XP beads to the solution (1:1 ratio i.e 225μ l) of IVT reaction and ethanol and mix well by inverting or pipetting with a P1000.
13	Incubate at room temperature for $ \odot 00:05:00 $.
14	Place the tubes on a 1.5 mL magnetic separation rack.
15	Wait © 00:05:00 to allow the beads to separate if using a 1.5mL rack. Allow a longer time if necessary to ensure proper separation of beads liquid before proceeding. while waiting make up a batch of 80% ethanol for next step.
16	Make up a batch of 80% ethanol for following wash steps. Usually 5ml is sufficient for washing both crRNA & tracrRNA tubes.
17	Remove and discard the supernatant while tube remains in magnetic rack.
18	Rinse the beads with 1mL of 80% ethanol if using a 1.5mL tube . It is not necessary to resuspend the beads.
19	Wait © 00:01:00 then remove and discard the ethanol.
20	Repeat the wash step as described above. (Add the same amount of 80% ethanol, wait \bigcirc 00:01:00 , then discard the ethanol.)
21	Remove residual ethanol that collects at the bottom of the tube by using a P200 or P20.
22	Air dry the beads for © 00:05:00 in a 1.5mL tube or until the beads lose their glossy appearance. Sufficiently dry beads will appear matte. Be careful not to let the beads get too dry (appearing cracked or dusty).
23	Elute the RNA by resuspending the beads with an appropriate amount of nuclease-free H2O depending on the desired volume and concentration. For the proceeding nanopore sequencing we usually elute with 30-50µL nuclease free H2O.

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⊠ Nuclease-free Water Contributed by users

- $24 \quad \text{Flick tubes to combine and allow the RNA to elute off the beads by incubating at room temperature for } \\ \text{$00:10:00} \ .$
- 25 If necessary, pulse-spin the tubes to collect any liquid along the sides of the tubes.
- Place the tubes on the magnetic rack and allow them to separate until water is clear. This will take 5-10 minutes for a 1.5mL tube.
- 27 Collect the eluted RNA, being careful not to take up beads.

Quantify, anneal and aliquot dgRNA

28 Using the XR RNA Qubit kit, quantify 1μL of the eluted tracrRNA and 1μL of the eluted crRNA. Follow standard XR RNA Qubit protocol. if concentration is too large to read initially make a 1 in 10 dilution.

⊠ Qubit™ RNA XR Assay Kit **Thermo Fisher**

Scientific Catalog #Q33223

29 Store RNA at δ-80 °C in small aliquots (we usually store 3μl-6μl aliquots in PCR tubes) in order to avoid freezethaws. The crRNA & tracrRNA should be freezed separately. Annealing to make the dgRNAs should be done only prior to sequencing.

Preparing input DNA *From Oxford nanopore protocol - please refer to their protocol for further detail

- 30 Prepare $5\mu g$ of high molecular weight genomic DNA (a range of 1-10 μg can be used but we found that higher DNA input works better).
- 31 Prepare DNA in nuclear free water to adjust to a 24µl total volume in a PCR tube. Flick and spin down to incorporate.

Preparing cas9 ribonucleoprotein complexes (RNPs)

- 32 Preheat thermal cycler to § 95 °C . Thaw an aliquot of NEB CutSmart Buffer, mix by vortexing and place on ice.
- 33 $\,$ In a PCR tube add in the following order (ensure crRNA & tracrRNA are equimolar 100 $\mu M)$:

Α	В
Duplex Buffer	8µL
crRNA pool	1µL
(100µM)	
tracrRNA	1µL
(100µM)	

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*To help calculate the volume required from the amount of crRNA & tracrRNA previously made we used the biotin calculator and the value was then divided by 100 but pipetting error was also considered. (https://www.bioline.com/media/calculator/01_07.html - Calculation of the molar concentration)

Pipette mixture up and down. Place in thermal cycler for 5 mins at 8 95 °C

34 Cool to room temperature and spin down to pool liquids - the crRNA: tracrRNA complexes will now be annealed.

Forming RNPs

35 In a 1.5ml Eppendorf tube

Α	В
Annealed tracrRNA : crRNA	10μL
complex	
10x NEB cutsmart buffer	10μL
Nuclease free H2O	79.2µL
Pyrogenes Hifi Cas9 nuclease	0.8µL
Total Volume	100µL

^{*} further information can be found in the Oxford protocol around scaling this reaction down accordingly.

Mix by flicking tube and incubate at room temperature for 30 mins then place on ice, the RNP complexes are now formed.

Phosphorylate genomic DNA

36 In a PCR tube add:

Α	В
10x CutSmart Buffer	3µL
Your Genomic DNA	24µL
Total reaction Volume	27µL

Mix gently by flicking tube and spin down.

⊠ CutSmart Buffer - 5.0 ml **New England**

Biolabs Catalog #B7204S

37 Add $3\mu I$ of CIP to the tube

12m

Using a Thermocycler, incubate at $\$ 37 °C for \odot 00:10:00 , $\$ 80 °C for \odot 00:02:00 then hold at $\$ 20 °C .

⊠ Alkaline Phosphatase, Calf Intest (CIP) - 1,000 units New England

Biolabs Catalog #M0290S

Cleave and dA-tail the target DNA

38 Thaw dATP tube, vortex to mix and place on ice.

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39 In a 1.5ml Eppendorf lobind tube make up dATP mix to 10mM stock. Vortex to mix and spin down (1μl of dATP to 9μl nuclease free water), then place on ice.

20m

⋈ Nuclease-free Water Contributed by users

40 To the PCR tubes containing the 30μl of desphophorylated DNA add:

Α	В
Desphosphorylated genomic DNA sample	30µl
Cas9 RNPs	10µl
10 mM dATP	1µl
NEB Taq polymerase	1µl
Total reaction volume	42µl

Carefully mix the contents by inversion of tube, spin down and place in thermal cycler.

Incubate sample for the following cycle: § 37 °C for © 00:15:00 § 72 °C for © 00:05:00 and hold at § 4 °C

Adaptor Ligation

41 Thaw Ligation Buffer (LNB) at room temperature, mix via pipetting due to viscosity. Place on ice.

Thaw the adaptor mix (AMX), mix by flicking tube and pulse spin. Return to ice.

Place Ampure XP beads at room temperature

- 42 Using a wide-bore tip transfer contents from the previous step from the PCR tube into a 1.5ml lobind eppendorf tube.
- 43 In a seperate 1.5mL Eppendorf Tube assemble:

Α	В
Ligation Buffer (LNB)	20µl
Nuclease free water	ЗμΙ
NEBNext Quick T4 DNA ligase	10µl
Adapter Mix (AMX)	5µl
Total reaction volume	38µl

 $\operatorname{\mathsf{Mix}}$ by pipetting up and down to ensure viscous LNB is incorporated

44 Add 20µl of this adapter ligation mix to the cleaved and dA-tailed sample in the other 1.5ml eppendorf tube. Mix gently by flicking up and down. immediately after mixing, add the remainder of the adapter ligation mixture to the cleaved and dA-tailed sample, which will yield an 80µl ligation mix.

Mix by flicking the tube and pulse spin to collect droplets.

Incubate mixture for © 00:10:00 at room temperature.

Purify adapter ligated DNA - AMpure XP bead purification

45 Resuspend the AMpure XP beads by vortexing.

Thaw short fragment buffer (SFB) and EB from SQK-LSK109

46 Add 1x volume (80μl) of TE pH8.0 to the ligation mix and mix by gentle flicking of the tube. Then add 0.3x volume (48uL) of AMPure XP beads to the ligation sample.

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	Gently invert tube then spin down
	Incubate sample at room temperature for $© 00:10:00$ - do not pipette or agitate sample.
47	Spin down sample and pellet on a magnetic rack, then pipette off the supernatant.
48	Using 250µl of Short Fragment Buffer (SFB) resuspend the beads and flick to incorporate. Return tube to magnetic rack. Allow beads to pellet.
	Remove supernatant via pipette and discard.
	Repeat this SFB wash step.
49	After removal of supernatant, let sample dry for ③ 00:00:15 . Add 13uL of Elution Buffer (EB). Incubate for ⑤ 00:10:00 at room temperature.
	Pellet beads on a magnet and remove 12uL of eluate to a clean 1.5mL Eppendorf DNA LoBind Tube.
	This prepared library will be used for loading into the flow cell. Store on ice until ready to load.
Priming	and loading the flow cell 5m
50	Thaw sequencing buffer (SQB), loading beads (LB), Flush tether (FLT) and one tube of flush buffer (FB). After thawed place on ice.
51	Mix sequencing buffer (SQB) and flush buffer (FB) tubes by vortexing, spin down and return to ice.
	Mix the flush tether (FLT) by pipetting, then return to ice.
52	Open the lid of the nanowire sequencing device and slide the flow cell's priming port cover clockwise so that the priming port is visible.
	With a P1000 μ l pipette set to 200 μ l insert tip into the priming port and turn the wheel until the dial shows 220-230 μ l whereby a small volume of buffer will be seen in the tip.
53	Prepare the flow cell priming mix: add 30µl of flush tether (FLT) directly to the tube of flush buffer (FB) and mix by pipetting up and down.
54	Load 800µl of the priming mix into the flow cell via the priming port, avoiding the introduction of air bubbles.
	Wait © 00:05:00
55	Thoroughly mix the contents of the Loading beads (LB) via pipetting.
56	In a new 1.5ml lobind eppendorf tube prepare the library for loading:

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Α	В
Sequencing Buffer (SQB)	25μΙ
Loading Beads (LB)	13µl
DNA library	12µl
Total volume	50μΙ

57 Gently lift the sample port cover to make the sample port accessible.

Load 200µl of the priming mix into the flow cell via the priming port - avoiding the introduction of air bubbles.

- 58 Mix the prepared library by gently pipetting up and down immediately prior to loading.
- Add 50µl of sample to the flow cell via the sample port in a slow drop wise fashion. Ensure each drop flows into the port before adding the next.
- $60 \quad \text{Gently replace the sample port cover, close the priming port and replace the MinION lid.}$

Starting the sequencing run 5m

61 Follow the steps as outlined by Nanopore to start the sequencing run.