In vitro transcription of guide RNAs

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Abstract

In vitro T7 template synthesis and transcription sgRNA protocol with SPRI beads

Citation: Mark Dewitt & Julia Wong In vitro transcription of guide RNAs. protocols.io

dx.doi.org/10.17504/protocols.io.dm749m

Published: 13 Aug 2015

Guidelines

Assembly Oligos:

T7FwdVar:

GGATCCTAATACGACTCACTATAG---guide-sequence---GTTTTAGAGCTAGAA

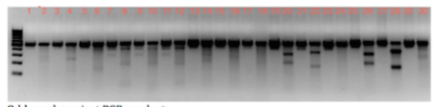
T7RevLong:

AAAAAAGCACCGACTCGGTGCCACTTTTTCAAGTTGATAACGGACTAGCCTTATTTTAACTTGCTATTTCTAGCTC
TAAAAC

Amplification Primers:

T7FwdAmp: GGATCCTAATACGACTCACTATAG

T7RevAmp: AAAAAAGCACCGACTCGG



Odd numbers: just PCR product Even numbers: with T7E1 treatment

Here's a gel that shows some good guides and some that didn't work examples of more promising looking guides would be #'s 20, 22, 26, and 28.

Protocol

Making the Template (for 1 sgRNA)

Step 1.

see guidelines for the primers

Set up the following 50 µl reaction



50 μl Additional info:

₽ PROTOCOL

. T7 PCR Template for 1sgRNA Mixture

CONTACT: Jacob Corn

Step 1.1. 35.5 μl H2O

Step 1.2.

10 μl 5x Phusion HF Buffer

Step 1.3.

 $1 \, \mu l \, 10 \, mM \, dNTPs$

Step 1.4.

1 μ l T7FwdVar (1 μ M) \leftarrow THIS IS THE PART WE DESIGN

Step 1.5.

1 μl T7RevLong (1 μM)

Step 1.6.

0.5 μl T7FwdAmp (100 μM)

Step 1.7.

0.5 μl T7RevAmp (100 μM)

Step 1.8.

0.5 μl Phusion HF DNA polymerase (2u/μl)

PCR

Step 2.

Run PCR:

95° 30 sec

95° 10 sec

57° 10 sec

72° 10 sec

30x steps 2-4

72° 2 min

4° hold

No PCR cleanup necessary at this point

Making template

Step 3.

Mix: Final conc.

2 μl 10x Buffer 1x 2 μl ATP (100 mM) 10 mM 2 μl GTP (100 mM) 10 mM 2 μl CTP (100 mM) 10 mM 2 μl UTP (100 mM) 10 mM

8 μl DNA template (85 ng/μl) ←PCR PDTS 25 ng/μ

12 μl T7 RNA polymerase mix

20 µl total

Incubated this mix for 18 hours at 37° in a thermalcycler. Added 1 µl of RNase-free DNase; incubated

20 min, room T.

O DURATION

00:18:00

Run PCR

Step 4.

Run PCR

95C 30s

95C 10s

57C 10s

72C 10s

steps 2-4, 30 cycles

72 2min

4C hold

No PCR cleanup necessary at this point

T7 transcription

Step 5.

Make 20 µl total T7 transcription mix

■ AMOUNT

20 µl Additional info:

₽ PROTOCOL

. T7 transcription mix

CONTACT: Jacob Corn

T7 transcription

Step 6.

Incubate transcription mix for 18 hours at 37° in a thermalcycler

© DURATION

18:00:00

T7 transcription

Step 7.

Add 1 µl of RNase-free DNase; incubated 20 min, room T

© DURATION

00:20:00

SPRI Beads clean-up of sgRNAS (96-well plate):

Step 8.

IVT sqRNAs to 20uL (or 10uL)

SPRI Beads clean-up of sgRNAS (96-well plate):

Step 9.

Bring volume to 150 uL with 100% EtOH (this helps binding of small fragments)

SPRI Beads clean-up of sgRNAS (96-well plate):

Step 10.

Add 5X SPRI (we use homemade SeraPure beads for RNA binding) 5*10 (IVT sqRNA)= 50 uL of SPRI Beads 5*20 (IVT sgRNA)= 100 uL SPRI Beads SPRI Beads clean-up of sgRNAS (96-well plate): **Step 11.** Pipette to mix 10 times SPRI Beads clean-up of sgRNAS (96-well plate): **Step 12.** Incubate 5 minutes at room temperature © DURATION 00:05:00 SPRI Beads clean-up of sgRNAS (96-well plate): **Step 13.** Place on magnetic stand, 5 min **O DURATION** 00:05:00 SPRI Beads clean-up of sgRNAS (96-well plate): **Step 14.** Discard supernatant SPRI Beads clean-up of sgRNAS (96-well plate): **Step 15.** Wash#1 Add 200 uL, 80% EtOH. Wait 2 min. Remove EtOH. © DURATION 00:02:00 **Step 16.** 8b)Wash #2: Add 200 uL, 80% EtOH. Wait 2 min. Remove EtOH. © DURATION 00:02:00 SPRI Beads clean-up of sqRNAS (96-well plate): Step 17. Air dry 5-10 min (pellet will change from a glossy/wet to matte/dry.) **O DURATION** 00:10:00 SPRI Beads clean-up of sgRNAS (96-well plate): **Step 18.** Elute 20 uL of water or TE. Pipette mix 10 times. SPRI Beads clean-up of sqRNAS (96-well plate): **Step 19.** Incubate 2 minutes at room temperature © DURATION 00:02:00 SPRI Beads clean-up of sgRNAS (96-well plate):

Step 20. Place on r

Place on magnetic stand, 5 min

© DURATION 00:05:00

SPRI Beads clean-up of sgRNAS (96-well plate):

Step 21.

Keep Supernatant. Transfer to a new plate / tubes.