



## 7 **▼**Jun 01, 2022

# © 2-step PCR mixture and conditions (Barcoded-head primers for seqs pooling) V.7

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protocol.



#### PCR mixture and condition (2X SUPERGREEN PCR MASTER MIX)

Yin-Tse Huang 2022. 2-step PCR mixture and conditions (Barcoded-head primers for seqs pooling) **protocols.io** 

 $\label{lem:https://protocols.io/view/2-step-pcr-mixture-and-conditions-barcoded-head-pr-cadgsa3w Yin-Tse Huang$ 

protocol

Herbold CW, Pelikan C, Kuzyk O, Hausmann B, Angel R, Berry D, Loy A. 2015. A flexible and economical barcoding approach for highly multiplexed amplicon sequencing of diverse target genes. Front. Microbiol. [Internet] 6:731. Available from: http://dx.doi.org/10.3389/fmicb.2015.00731

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1 Wear glove, clean up the working bench w. 1% bleach

## For 1' PCR head-primers

2 Prepare 1' PCR master mixutre for head-primers (prepare 1.2X of solutions for pipetting error if needed)

PCR mixture for head-primers for each reaction

A	В	С	D		
Component	Volume	Volume	Final conc.		
		(1.2X)			
Forward Primer (10 µM)	1.6 µl	1.9 μΙ	1 μΜ		
Reverse Primer (10 µM)	1.6 μΙ	1.9 μΙ	1 μΜ		
2X Supergreen PCR Master Mix	7.8 µl	9.4 μΙ	-		
ddH20	4.1 μΙ	4.9 µl	-		
Total volume	15 μΙ	18 μΙ	-		

Negative control ALWAYS NEEDED! For example, if you have 5 PCR reactions to run, prepare master mixture for 6 reactions (5 DNA template + 1 negative control).

- 3 Mix the 1' PCR master mixture gently by pippeting. Quick spin the tube.
- 4 Transfer **□15 μL** 1' PCR master mixutre in 8-strip PCR tubes.
- 5



Add **30.6** µL DNA template in 8-strip PCR tubes, resulting in a **15.6** µL reaction mixture for 1' PCR.

**Negative control** contains only  $\Box 15 \mu L$  master mixture but not DNA template

- Mix the reaction mixture gently by tapping the tubes. Quick spin the tubes.
- Carry out PCR using the following condition:

#### 1' PCR condition for head-primers

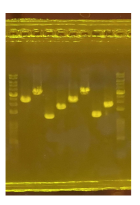
Α	В	С	D
Step	Temp	Sec	Cycle
Initial denaturation	95 °C	30-180 (a)	
Denaturation	98 °C	15	20-25 cycles
Annealing	64-68 °C varied (b)	15	
Extension	72 °C	60-180 (c)	
Final extension	72 °C	210	
Preservation	Preservation	4 °C	∞

a. Varied depend on template complexity
b. Annealing varied, 62-65C is working based on test on 220530; Refer to 1' PCR primers for annealing temperature
c. 1kb ~ 1min extension; enough time allow full extension of sequence

#### 7.1 1' hear-primers used in Huang lab

Α	В	С	D	
Name	Sequence	Tm°C	CG%	
NS1B1ngs_H1	GCTATGCGCGAGCTGCcctngttgatyctgccagt	71.7	60	
ITS4ngs_H1	GCTATGCGCGAGCTGCtcctscgcttattgatatgc	69	55.6	
LR5_H1	GCTATGCGCGAGCTGCtcctgagggaaacttcg	70.2	60.6	
EF1-526F_H1	GCTATGCGCGAGCTGCgtcgtygtyatygghcaygt	71	59.3	
EF1-1567R_H1	GCTATGCGCGAGCTGCachgtrccrataccaccratctt	70.6	56	
EF1-2218R_H1	GCTATGCGCGAGCTGCatgacaccracrgcracrgtytg	72.2	60.3	
Ben2f_H1	GCTATGCGCGAGCTGCtccagactggtcagtgtgtaa	70.5	56.8	
Bt2b_H1	GCTATGCGCGAGCTGCaccctcagtgtagtgacccttggc	74.5	62.5	
T22_H1	GCTATGCGCGAGCTGCtctggatgttgttgggaatcc	70.3	56.8	
RPB2-3bF_H1	GCTATGCGCGAGCTGCggwggwtayttyatyatyaatgg	65.6	48.7	
RPB2-7cR_H1	GCTATGCGCGAGCTGCcccatrgcttgyttrcccat	72.3	59.7	
fRPB2-11aR_H1	GCTATGCGCGAGCTGCgcrtggatcttrtcrtcsacc	71.7	60.8	

## Carry out **electrophoresis** for inspection of DNA products



Gel before markdown

Markdown wells and upload the pictures to the Lab Google drive



Marked gel picture go to the Lab Google drive

#### For 2' PCR barcoded-head primers

10 Prepare 2' PCR master mixutre for barcoded-primers (prepare 1.2X of solutions for pipetting error if needed)

PCR mixture for barcoded-primers for each reaction (NO PRIMERs at this point!!)

A	В	С	D	
Component	Volume	Volume (1.2X)	Final conc.	
2X Supergreen PCR Master Mix	10.75 μL	12.9 µL	-	
ddH20	10.75 μL	12.9 µL	-	
Total volume	21.5 μL	25.8 μL	-	

Negative control ALWAYS NEEDED! For example, if you have 5 PCR reactions to run, prepare master mixture for 6 reactions (5 DNA template + 1 negative control).

- 11 Mix the 2' PCR master mixture gently by pippeting. Quick spin the tube.
- 12 Transfer  $\blacksquare$ 21.5  $\mu$ L of the 2' PCR master mixture to 8-strip PCR tubes.
- 13 Add **■2.5 µL pre-mixed barcoded-head primers** (Forward + Reverse) to each PCR tubes.
- 14 🛕

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**Negative control** contains only  $\square$ 24  $\mu$ L master mixture and premixed barcoded-head primers but not DNA template

- 15 Mix gently by tapping the tubes. Quick spin the tubes.
- 16 Carry out 2' PCR using the following condition:
  - 2' PCR condition for barcoded-head primers

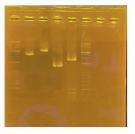
Α	В	С	D		
Step	Temp	Sec	Cycle		
Initial denaturation	98 °C	30			
Denaturation	98 °C	15	10-15 cycles		
Annealing	64-68 °C varied (a)	15	-		
Extension	72 °C	60 (b)			
Final extension	72 °C	210			
Preservation	Preservation	4 °C	∞		

a. Annealing varied,  $\bf 65\,C$  is working based on test on 220531; Refer 2' PCR primers for annealing temperature b. 1kb ~ 1min extension; enough time allow full extension of sequence

## 16.1 2' barcoded-head primers used in Huang lab

Α	В	С	D	E	F	G	Н	ı	J
Name (1st)	Sequence	Tm°C	CG%	Name (2nd)	Sequence	Tm°C	CG%	Name (3rd)	Sequence
F1-1	aaatctcattttctggtaggccggctatgcgcgagctgc	71.7	53.8	F2-1	ggccattcgtagtaagtagttcagctatgcgcgagctgc	70	53.8	F3-1	gggctagagtttaaccttaattag
F1-2	aacccgcacgtctattttaagaagctatgcgcgagctgc	70.1	51.3	F2-2	cagatactaattctcgtcacatggctatgcgcgagctgc	68.5	51.3	F3-2	taatttatggtctacagacgttag
F1-3	aactagtttatttcaaattaatagctatgcgcgagctgc	61.3	35.9	F2-3	agagattgagcttatccgtttttgctatgcgcgagctgc	68.7	48.7	F3-3	gtaagcgctcatacagcaaacta
F1-4	aatatgaageteegacatatetggetatgegegagetge	69.5	51.3	F2-4	ccattgattccagattatcatgtgctatgcgcgagctgc	67.7	48.7	F3-4	atccgcatatctaactcgaaagc
F1-5	acagtacaaacacggctcattaagctatgcgcgagctgc	70	51.3	F2-5	ccgtaggacttcgtagtttaaacgctatgcgcgagctgc	70.1	53.8	F3-5	gtgcgtgcaaatcgttttgtatcgc
F1-6	acagttgaccatccgacattatcgctatgcgcgagctgc	71.2	53.8	F2-6	gtcatgaccgctacatattctcagctatgcgcgagctgc	70	53.8	F3-6	gtggggataagcttgacattttag
F1-7	caccataaatgagattgctggaggctatgcgcgagctgc	70.7	53.8	F2-7	cgataggacaagcaatgtactcagctatgcgcgagctgc	70.1	53.8	F3-7	gttggataggaggtcaaacaga
F1-8	accgtgtatgtattcgtgttacggctatgcgcgagctgc	71	53.8	F2-8	cgatataatcgatccgccataccgctatgcgcgagctgc	71.4	56.4	F3-8	gtttaacccataagtgcgaccate
R1-1	acgtcaatgctattcccagtcaagctatgcgcgagctgc	71.4	53.8	R2-1	cagcgtttccaaagacattattggctatgcgcgagctgc	69.7	51.3	R3-1	taaaaattcctggaactccacag
R1-2	acgtcgaggtatcataaatacttgctatgcgcgagctgc	67.9	48.7	R2-2	cgctttatatgcttaaagtacccgctatgcgcgagctgc	69.1	51.3	R3-2	taatccatgtgtctattcttagagc
R1-3	cgcaagtatcgtctttcatagtcgctatgcgcgagctgc	70.1	53.8	R2-3	ctgctgagttataccacagtgacgctatgcgcgagctgc	71.5	56.4	R3-3	gtaaatctaggtgtaaaatgagt
R1-4	agtcaatgctttgggtacataaggctatgcgcgagctgc	70.1	51.3	R2-4	tacgttatattaactctagccgagctatgcgcgagctgc	67.4	48.7	R3-4	ctttcttataatatccgggctaagc
R1-5	atatcttacacaaaagtatcgttgctatgcgcgagctgc	65.4	43.6	R2-5	tagaatgtcaacacaagtaggacgctatgcgcgagctgc	69.5	51.3	R3-5	cttttgtaatggtgttgtccgttgct
R1-6	cgactattttaacttccgcaacagctatgcgcgagctgc	69.3	51.3	R2-6	gaaatcgaacaaattctgccttcgctatgcgcgagctgc	69.7	51.3	R3-6	taggtatcattctcatcctatcggc
R1-7	atcgttggttctatgttcaggtagctatgcgcgagctgc	69.5	51.3	R2-7	gaatcatcaagaagggaacaacagctatgcgcgagctgc	69.3	51.3	R3-7	tatagagacgggtttcggtaaaa
R1-8	atgagatctatctagtacccgttgctatgcgcgagctgc	68.8	51.3	R2-8	gactatagtgaaaaatcacatacgctatgcgcgagctgc	65.8	46.2	R3-8	tattagtatagatagacactcggg
R1-9	gaggagccatagagtatcaatgagctatgcgcgagctgc	69.7	53.8	R2-9	atgatcctaccggagatttacctgctatgcgcgagctgc	70.7	53.8	R3-9	tcaggggtagaagactagttgta
R1-10	atttgtagcattgaataggagcagctatgcgcgagctgc	68.7	48.7	R2-10	gagttctggatatctatgggcttgctatgcgcgagctgc	70	53.8	R3-10	tcgggttaaatgctaagcgtaatq
R1-11	caaattcctatcgtacgtgatccgctatgcgcgagctgc	70.2	53.8	R2-11	gcaacgaaacattcgttaagtatgctatgcgcgagctgc	68.1	48.7	R3-11	tgctgaaaacaggaagtctcact
R1-12	caagaactagacgctgctcttaagctatgcgcgagctgc	70.2	53.8	R2-12	tgcccaatatgttagcaccctaagctatgcgcgagctgc	71.4	53.8	R3-12	accattcctaatagccaataggg

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Marked gel picture go to the Lab Google drive

