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

**Y** Forked from <u>2-step PCR mixture and conditions</u> (Barcoded-head primers for seqs pooling)

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**ABSTRACT** 

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

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protocols.io

b.2015.00731

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#### **MANUSCRIPT CITATION:**

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/fmic

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**Protocol status:** Working We use this protocol and it's working

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#### **PROTOCOL** integer ID:

80784

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 (1.2X)	Final conc.
Forward Primer (10 µM)	1 μΙ	1.2 μΙ	0.4 μΜ
Reverse Primer (10 µM)	1 μΙ	1.2 μΙ	0.4 μΜ
ZEJU PCR Master Mix	12.5 μΙ	15 μΙ	-
ddH20	9.25 μΙ	11.1 μΙ	-
Total volume	23.75 μΙ	28.5 µl	-

#### Note

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 Δ 23.75 μL 1' PCR master mixutre in 8-strip PCR tubes.

Add 🗸 1.25 µL DNA template in 8-strip PCR tubes, resulting in a 🔼 25 µL reaction mixture for

1' PCR.

#### Note

Negative control contains only 🚨 23.75 µL master mixture but not DNA template

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

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

A	В	С	D
Step	Temp	Sec	Cycle
Initial denaturation	95 °C	180	
Denaturation	95 °C	30	
Annealing	60-66 °C varied (b)	30	20-35 cycles
Extension	72 °C	180	
Final extension	72 °C	420	
Preservation	Preservation	4 °C	∞

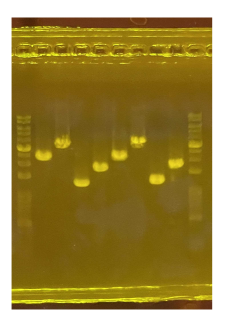
- b. Annealing varied, 60-66C is working; Refer to 1' PCR primers for annealing temperature
- c. 1kb ~ 1min extension; enough time allow full extension of sequence

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

A	В	С	D
Name	Sequence	Tm°C	CG%

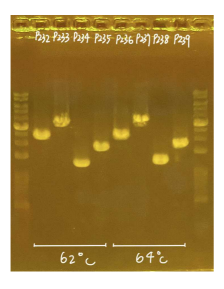
A	В	С	D
NS1B1ngs_H1	GCTATGCGCGAGCTGCcctngttgatyctgccag t	71.7	60
LR5_H1	GCTATGCGCGAGCTGCtcctgagggaaacttcg	70.2	60.6
EF1-526F_H1	GCTATGCGCGAGCTGCgtcgtygtyatygghca ygt	71	59.3
EF1-2218R_H1	GCTATGCGCGAGCTGCatgacaccracrgcracr gtytg	72.2	60.3

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



Gel before markdown

9 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

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.
ZEJU PCR Master Mix	7.5 µL	9 μL	-
ddH20	5.55 µL	6.66 µL	-
Total volume	13.05 µL	15.66 µL	-

#### Note

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.

- Transfer  $\underline{A}$  13.05  $\mu L$  of the 2' PCR master mixture to 8-strip PCR tubes.
- Add Add 1.2 µL pre-mixed barcoded-head primers (Forward + Reverse) to each PCR tubes.
- Add  $\Delta 0.75 \,\mu L$  of 1' PCR product as template, resulting in  $\Delta 15 \,\mu L$  reaction mixture for 2' PCR.

Negative control contains only  $\Delta$  14.25  $\mu L$  master mixture and premixed barcoded-head primers but not DNA template

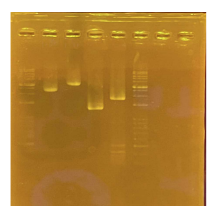
- Mix gently by tapping the tubes. Quick spin the tubes.
- 16 Carry out 2' PCR using the following condition:

## 2' PCR condition for barcoded-head primers

A	В	С	D	
Step	Temp	Sec	Cycle	
Initial denaturation	98 °C	30		
Denaturation	98 °C	15		
Annealing	64-68 °C varied (a)	15	10-15 cycles	
Extension	72 °C	20 (b)		
Final extension	72 °C	210		
Preservation	Preservation	4 °C	∞	

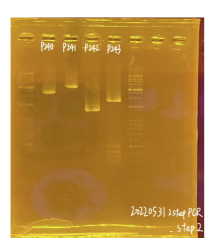
- a. Annealing varied,  $\bf 65~C$  is working based on test on 220531; Refer 2' PCR primers for annealing temperature
- b. 1kb ~ 1min extension; enough time allow full extension of sequence

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



Gel before markdown

# 18 Markdown wells and upload the pictures to the Lab Google drive



Marked gel picture go to the Lab Google drive