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Bacterial DNA Barcoding PCR Kit (KIT-1110-50)





Bacterial DNA Barcoding PCR Kit

Product No: KIT-1110-50

This kit contains PCR reagents to amplify the 16s rRNA full length gene for Sanger sequencing. The optimized PCR protocol generate end products for Sequencing+ PLUS Services from 1st BASE sequencing services. Customer can align and BLAST the obtained sequencing results to their choice of database for barcoding purposes. If customer choose to send their PCR products for other sequencing provider, the PCR products need to be purified before sequencing. It is an ideal kit to perform DNA barcoding of various bacterial samples readily.

For Research Use Only. Not for use in Diagnostic Procedures.

Kit Contents

No	Product	KIT-1110-50	Storage
1	16S Primer Mix	600 μ L	-20°C
2	16S Enzyme	30 μ L	
3	16S PCR Buffer	750 μ L	
4	Bacterial Plasmid Positive Control	25 μ L	
5	Sequencing Forward Primer, 785F (10 μ M)	150 μ L	
6	Sequencing Reverse Primer, 907R (10 μ M)	150 μ L	



Product Specification

	KIT-1110-50
Sample	Purified Genomic DNA from bacteria
Duration	PCR amplification ~ 45 minutes
Storage	-20 °C

Materials Supplied by Users

- ✓ Thermocycler
- ✓ Electrophoresis reagents and system
- ✓ Sterile nuclease-free 0.2 mL PCR tubes or 96-well plate
- ✓ Sterile nuclease-free pipette and pipette tips

Precautions for Users

- ✓ Always wear a lab coat, disposable gloves and apply proper aseptic techniques to conduct molecular biology experiments.



Protocol

PCR	<ol style="list-style-type: none"> 1. Prepare the PCR Mix according to Table 1. 2. Add 2 μL of purified genomic DNA (gDNA) at the concentration of 15 – 25 ng/μL as DNA Template with each 23 μL of PCR Mix into 0.2mL tube or 96-well plate. 3. Run the PCR Cycle Protocol on Thermocycler according to Table 2. 4. After the PCR cycle is completed, check the present of \sim1.5 kb PCR products on 1% agarose gel according to Figure 1.
Sequencing	<ol style="list-style-type: none"> 5. For the unpurified PCR products that shows single band at \sim1.5 kb on agarose gel electrophoresis, they are ready to send for 1st BASE Sequencing+ PLUS Services using the provided 785F and 907R sequencing primers. <p><i>Tips:</i></p> <ul style="list-style-type: none"> ✓ <i>If there is no amplification or no PCR products generated from your bacteria sample, please refer Troubleshooting Guidelines.</i> ✓ <i>1st BASE Sequencing+ PLUS Services has included PCR clean-up before sequencing. The turnaround time is \sim3 – 4 working days from the day of the unpurified PCR products received by 1st BASE.</i>
Barcoding	<ol style="list-style-type: none"> 6. After the sequencing results are ready, trim off the reads with Quality Value (QV) < 20, align the Forward and Reverse sequencing results. 7. BLAST the aligned sequence against your preferred database, e.g. NCBI, Greengenes or others. 8. The identification of the bacteria is reliable up to genus level and it typically appears within the top-10 of nucleotide BLAST results.



Table 1: Preparation of PCR Mix

Number of Reactions	16S Primer Mix (μL)	16S Enzyme (μL)	16S PCR Buffer (μL)	Total PCR Mix (μL)
2	20	1.0	25.0	46
3	30	1.5	37.5	69
4	40	2.0	50.0	92
5	50	2.5	62.5	115
6	60	3.0	75.0	138
7	70	3.5	87.5	161
8	80	4.0	100.0	184
9	90	4.5	112.5	207
10	100	5.0	125.0	230

Note 1:

- ✓ The PCR Mix must be freshly prepared.
- ✓ The recommended DNA template amount in each PCR is 30 – 50 ng.
- ✓ Each PCR consists of 23 μL of PCR Mix and 2 μL of diluted DNA Template.
- ✓ Both NTC (No Template Control) and positive control reactions are recommended to be included into each round of PCR preparation.
- ✓ For each positive control reaction, use 1 μL of the provided plasmid positive control (5 ng/μL) as DNA Template.
- ✓ For each NTC reaction, use 1 μL of the provided TE Buffer as DNA Template.

Table 2: PCR Cycle Protocol

Step	PCR Process	Time	Temp. (°C)	Number of Cycle
1	Initial Denaturation	2 min	94	1
2	Denaturation	10 sec	98	25 cycles
3	Annealing	30 sec	53	
4	Extension	60 sec	68	

Note 2:

- ✓ Always check the present of PCR end products, which is ~1.5 kb of size on agarose gel electrophoresis before send for 1st BASE Sequencing+ PLUS Services.
- ✓ If you have alternative sequencing service provider, please purify the PCR products (PCR purification reagents not provided with this kit) before sequencing.

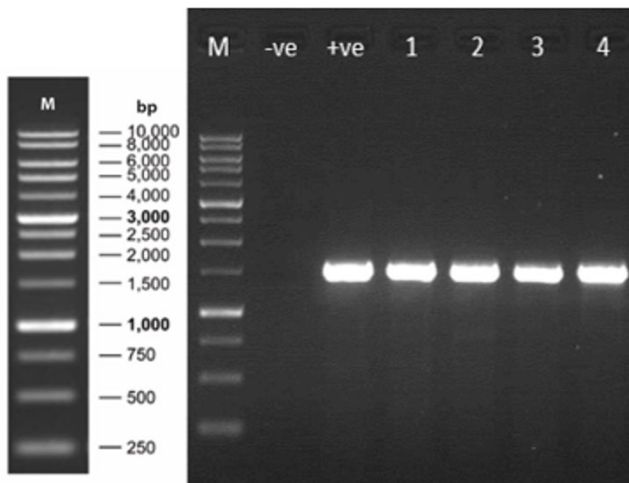


Figure 1. 1uL of un-purified PCR products on 1% TAE Agarose Gel. PCR was performed in 25uL according to Table 2.

-ve: NTC (No Template Control)

+ve: Bacterial Plasmid Positive Control, 1uL per reaction

1 to 4: Bacterial Nucleic Acid, 30 to 50ng per reaction



Troubleshooting Guidelines

Problems	Reasons
No PCR amplification	<ul style="list-style-type: none"> ▪ Presence of PCR inhibitors: Perform 5x - 20x dilution of DNA Template and repeat PCR using the diluted DNA Template.
NTC shown amplification	<ul style="list-style-type: none"> ▪ PCR mix in Table 1 must be prepared freshly. Do not use the premix that was prepared overnight.
Sanger sequencing results shown mixture of signal	<ul style="list-style-type: none"> ▪ More than 1 different copy of 16s gene were detected due to presence of more than 1 type of bacteria. <ol style="list-style-type: none"> a) Re-isolate single colony from the bacteria culture. Repeat the process of extraction and sequencing using the new PCR products amplified from pure culture; or b) If the mixture of bacteria is less than 5, by using the remaining PCR products, you may order cloning prior sequencing services using product code: MBS-3006. If the mixture of bacteria is a lot more complex, you may consider our next-generation sequencing services using product code: NGS-7008. New gDNA is required.



Product Ordering Information

Product Name	Packaging Size	Product Number
Bacterial DNA Barcoding PCR Kit	50 preps	KIT-1110-50
Bacterial DNA Barcoding Kit, including DNA Extraction	50 preps	KIT-1100-50
1st BASE Sequencing+ PLUS Services	2 sequencing reactions for each sample	SS1201
PCR Product Cloning Service PLUS (up to 1.5kb) + Cloning of unpurified PCR product into pJET1.2/ Blunt vector + Colony PCR screening and pick 5 positive colony PCR products for bi-directional sequencing.	1 sample	MBS-3006
Amplicon Sequencing Lite, partial gene (16s v3-v4 or 16s v4) with Basic Bioinformatics Analysis.	1 sample	NGS-7008
Bacterial DNA Barcoding Services from pure isolates, full gene of 16s rRNA + Extraction of gDNA + PCR amplification and Purification + Bidirectional PCR product sequencing + Data analysis (BLAST to show the top 10 matches from database)	1 sample	MBS-5002
Bacterial DNA Barcoding Services from mixed isolates, full gene of 16s rRNA + Extraction of gDNA, PCR Amplification and Purification + Cloning and 8 positive clones of plasmid DNA to send for Bidirectional Sequencing + Data analysis (Alignment & BLAST to show the top 10 matches from database)	1 sample	MBS-5102
<Optional> Custom DNA Barcoding Services	Minimum 5 samples	MBS-5007

Customization of DNA Barcoding kit for your choice of organism is available. Please contact us at <http://www.base-asia.com/find-us> for more information.



Note

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