



LoopSeq™ 16S Microbiome 24-Plex LoopSeq™ 16S & 18S Microbiome 24-Plex Quick Start Guide

Version 3.3

Sample Dilution

- Quantify template DNA by Qubit
- Dilute DNA stock to **2 ng/ul** using Buffer EB

Barcode Assignment PCR

- Spin down PCR plate
- Add **5ul of diluted DNA** sample to their respective wells (one sample per well) and **mix by pipetting** (do not vortex)
- Seal the plate with fresh caps or sealing film, and **spin down the PCR plate**
- Initiate the following PCR program:

Temperature	Duration	STEP
95°C	3 min	1 cycle
98°C	15 seconds	
48°C	20 seconds	2 cycles
72°C	2 min	
4°C	HOLD	

Set the heated lid at **100°C**
Ramp rate = **2 to 3°C/second**

Barcode Distribution

- Pool **5ul** from each of the 24 reactions into a single tube (ideally 1.5 mL) and mix by pipet (do not vortex)
- Transfer **95ul** of this pool to a new PCR tube
- Add **15ul of Inactivation Enzyme A** to the reaction
- Incubate at 37°C for 10 minutes, then 80°C for 5 minutes, then HOLD at 4°C
- Perform a cleanup reaction using **66ul** SPRI reagent
- Wash twice in 200ul 80% ethanol, do **not** dry beads
- Elute in **20ul** Buffer EB

Pause Point. Store sample at 4°C for up to 24 hours if needed



Calibration

- Add 18ul EB buffer to a 1.5 mL tube
- To this tube, add 2ul of the eluted sample (after 24-plex pooling and cleanup)
- This is the “diluted sample pool”

- Thaw on ice a chosen 2x qPCR Master Mix. Only use the following qPCR Master Mix reagent validated for this kit:

KAPA SYBR FAST qPCR Kit Master Mix, Catalog No. KK4600
BioRad iQ SYBR Green Supermix, Catalog No. 1708880

- Using a 96-well plate suitable for real-time SYBR-based qPCR, the following samples will be set up in the following reactions as duplicates:
 - 4 calibration standards (provided)
 - 1 sample (your samples, inactivated)
 - 1 NTC (no template control, your Buffer EB)

- Prepare a Master Mix for the above 6 reactions in duplicate with overage:
 - To a 1.5mL tube, add **140ul** of a chosen 2x qPCR Master Mix
 - Add **70ul of Calibration Reaction**

- Mix by vortex and briefly centrifuge
- Aliquot **15ul** of the Master Mix into 12 wells on a 96-well plate

- A. Column 1: Fill two wells with 5ul of **Calibration Standard 1**
- B. Column 2: Fill two wells with 5ul of **Calibration Standard 2**
- C. Column 3: Fill two wells with 5ul of **Calibration Standard 3**
- D. Column 4: Fill two wells with 5ul of **Calibration Standard 4**
- E. Column 5: Fill two wells with 5ul of Buffer EB (this is the negative control)
- F. Column 6: Fill two wells with 5ul of “diluted sample pool”

- Seal the plate using optically clear film or caps
- Briefly centrifuge the sealed plate for at least 20 seconds

- Initiate the following real-time qPCR program:

Temperature	Duration	STEP
95°C	3 min	1 cycle
98°C	15 seconds	
60°C	20 seconds	35 cycles
72°C	2 min	
4°C	HOLD	

Set the heated lid at **100°C**

SYBR signal is collected during the 2 min elongation

- Analyze the data on the Loop Genomics website at:

www.loopgenomics.com/start

- Calculate the volume of your sample pool to be carried forward into the next step of Barcode Distribution. Choose the volume appropriate for **HiSeq/NextSeq/NovaSeq**

Barcode Distribution

- Using a new PCR tube, combine the following to a final volume of 50 μ l:

Amplification Mix B	30μl
Nuclease-free water	20 - μl
DNA from previous step	μl*

* Volume may vary based on result of the calibration

- Initiate the following PCR program:

Temperature	Duration	STEP
95°C	3 min	1 cycle
98°C	15 seconds	
60°C	20 seconds	20 cycles
72°C	2 min	
4°C	HOLD	

Set the heated lid at 100°C
Ramp rate = 2 to 3°C/second

- Perform a cleanup reaction using **30 μ l** SPRI reagent
- Wash twice in 200 μ l 80% ethanol, do **not** dry beads
- Elute in **15 μ l** Buffer EB

- To the eluted sample, add **5 μ l** of **Distribution Mix**
- Add **2 μ l** of **Distribution Enzyme** the reaction
- Incubate at 20°C for 15 minutes, then 75°C for 5 minutes, then HOLD at 4°C

- Add **75.5 μ l** of **Activation Mix** to the reaction
- Add **2.5 μ l** of **Activation Enzyme** to the reaction
- Incubate at 20°C for 2 hours (heated lid off), then HOLD at 4°C

- Add **6 μ l** of **Neutralization Enzyme** to the reaction
- Incubate at 37°C for 15 minutes, then HOLD at 4°C

- Perform a cleanup reaction using **80 μ l** SPRI reagent
- Wash twice in 200 μ l 80% ethanol, do **not** dry beads
- Elute in **20 μ l** Buffer EB

Pause Point. Store sample at 4°C for up to 24 hours if needed



Library Preparation

- Using a new PCR tube, combine **on ice** the following:

Fragmentation Mix	20μl
Fragmentation Enzyme	10μl
DNA from previous step	20μl



- Finger-flick** to mix (do not vortex)
- Centrifuge for 2 seconds and immediately put back on ice
- Set up the following PCR program:

Temperature	Duration	STEP
4°C	1 min	Pause
32°C	12 min	1 cycle
65°C	30 min	1 cycle
4°C	HOLD	

Set the heated lid at 100°C

- Initiate the empty PCR machine (no tubes yet) then **pause** during the first 4°C step

- Place the assembled reaction in the PCR tube into the paused PCR machine and **resume** the program

- Take out the PCR tube once the program is completed
- Add **40 μ l** of **Ligation Mix** to the reaction
- Add **10 μ l** of **Ligation Enzyme** to the reaction
- Incubate at 20°C for 15 minutes (heated lid off), then HOLD at 4°C

- Perform a cleanup reaction using **80 μ l** SPRI reagent
- Wash twice in 200 μ l 80% ethanol, do **not** dry beads
- Elute in **20 μ l** Buffer EB

Pause Point. Store sample at 4°C for up to 24 hours if needed



- Using a new PCR tube, combine the following:

Index Master Mix	25μl
Index Primer P1 to P4	5μl (choose one Index Primer)
DNA from most previous elution	20μl

- Initiate the following PCR program:

Temperature	Duration	STEP
95°C	3 min	1 cycle
98°C	15 seconds	
60°C	30 seconds	12 cycles
72°C	30 seconds	
72°C	1 min	1 cycle
4°C	HOLD	

Set the heated lid at 100°C

- Perform a cleanup reaction using **40 μ l** SPRI reagent
- Wash twice in 200 μ l 80% ethanol, do **not** dry beads
- Elute in **20 μ l** Buffer EB

Process Complete: Ready for sample QC and sequencing