



DepleteX® Mito DNA Depletion Kit

KIT2002

For post-library depletion of Illumina NGS libraries

Contents

Product Overview	
Workflow	3
Kit Contents and Storage	2
Required Materials and Equipment	5
Best Practices	6
Revision Log	θ
Input Material	_
Input Material	
Protocol	8
	_
Step A: Depletion of Mitochondrial DNA	
Step B: PCR Amplification	10
Subsequent Steps	11

Contact Us

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Product Overview

The DepleteX® Mito DNA Depletion Kit targets DNA molecules derived from the human mitochondrial chromosome for removal from next generation sequencing (NGS) libraries. The kit leverages the CRISPR Cas9 endonuclease and its associated guide RNA to target and cleave human mitochondrial sequences. In samples with high mitochondrial content, such as ATAC-Seq libraries, use of the DepleteX Mito DNA Depletion Kit maximizes the re-assignment of sequencing reads to informative nuclear content and increases sequencing coverage of DNA derived from the nuclear fraction.

Description	
Total Assay Time	~3 hours
Hands-on Time	~1 hour
Samples Per Kit	24 samples
Sample Type	Human DNA NGS Library
DNA Library Input	10 ng
DNA Library Size	≥ 200 bp
Designed to Deplete	Human mitochondrial DNA
Method	CRISPR-Cas9 mediated depletion
Validated Library Prep	NEBNext® Ultra II FS DNA Library Prep Kit



Workflow

2

1 RNA Library Prep

DepleteX Mito DNA Depletion Kit (Post-Library Depletion)

Step A: Depletion of Mitochondrial DNA

- ☐ Hands-on time: ~30 minutes | Total time: ~2 hours
- 1. RNP Complex Formation & CRISPR Digestion
- 2. Bead Size Selection
- Safe Stopping Point

Step B: PCR Amplification

- Hands-on time: ~30 minutes | Total time: ~1 hour
- 1. PCR Amplification
- 2. Bead Cleanup
- Safe Stopping Point

3 Sequencing





Kit Contents and Storage

The DepleteX Mito DNA Depletion Kit contains enough material to deplete 24 NGS libraries. The kit contents and storage temperatures are indicated in the tables below.

ASY1064: Depletion Reagents (8 depletion reactions per box)

3 boxes

Storage at -20°C

Kit contents	Part number	Quantity per box
Cas9	REA1039	1 tube
10X Cas9 Buffer	REA1040	1 tube
RNase Inhibitor	REA1041	1 tube
Nuclease-Free Water	REA1042	1 tube
Tris Buffer	REA1043	1 tube
P5 Primer	REA1044	1 tube
P7 Primer	REA1045	1 tube
2X PCR Mix	REA1046	1 tube

ASY1069: Guide RNA for Mito DNA Depletion

1 Bag

Storage at -80°C

Kit contents	Part number	Quantity per bag
Guide RNA (Mito DNA)	REA1052	1 tube





Required Materials and Equipment

Туре	Item	Supplier
	0.5 mL, 1.5 mL DNA LoBind Tubes	Eppendorf (Cat# 022431021)
Plastic	0.2 mL thin wall PCR tubes	General Lab Supplier
Consumables	Low-Retention, Filtered, Sterile Tips (10 μL, 20 μL, 200 μL and 1000 μL)	General Lab Supplier
	AMPure XP Beads	Beckman Coulter (Cat# A63881)
Reagents	Absolute Ethanol, 200 Proof	General Lab Supplier
	Qubit dsDNA HS Assay Kit	ThermoFisher Scientific (Cat# Q32854)
	Single Channel Pipettes (10 μL, 20 μL, 200 μL, and 1000 μL)	General Lab Supplier
	Multichannel Pipettes (10 μL, 20 μL, and 200 μL)	General Lab Supplier
	Vortex Mixer	General Lab Supplier
	Microcentrifuge	General Lab Supplier
.	PCR Magnetic Rack or Stand	General Lab Supplier
Equipment	Ice Bucket	General Lab Supplier
	PCR Thermal Cycler	General Lab Supplier
	Qubit Fluorometer	ThermoFisher Scientific (Cat# Q33238)
	Automated electrophoresis Instrument such as Agilent TapeStation or 2100 BioAnalyzer	General Lab Supplier





Best Practices

General

- Periodically, optimizations and revisions are made to the kit components and user manual. Therefore, following the protocol included with or appropriate for the kit in question is important. This can be done by comparing the name and version number of the product to the name and version number of the protocol. If you need further assistance in this regard, please contact support@jumpcodegenomics.com
- This protocol describes the reagents, best practices, workflow, and method details for DepleteX Rare Transcript Boost kit.
- Use properly calibrated pipettes as library preparation is sensitive to pipetting error.
- Follow good laboratory practices throughout the protocol, including RNA-safe procedures when handling RNA. Ensure all pipette tips, microcentrifuge tubes, and other consumables are DNase- and RNase-free.
- When undertaking the protocol, always proceed immediately to the next step. If a stop is necessary, safe stopping points are available. Refer to the workflow schematic on page 3.

Reagent Handling

- Do not remove Cas9 and RNase Inhibitor from storage until before use. Maintain on ice during reaction setup. Return to -20°C immediately after use.
- Do not remove the guide RNA from storage until immediately before use. Maintain on ice during reaction setup. Return to -80°C immediately after use.
- We recommend a maximum of three freeze-thaw cycles for the Guide RNA. It is strongly recommended that multiple smaller aliquots of the Guide RNA be prepared when the reagent is first thawed if more than three freeze-thaw cycles are expected.
- Do not freeze AMPure® XP beads.
- Allow AMPure XP beads to come to room temperature before use. A 30-minute incubation on the laboratory bench is usually sufficient.
- Vortex AMPure XP beads immediately before use. Ensure that the beads are in a uniform suspension before use.
- Use magnetic stands appropriate for PCR tubes.

Equipment Handling

• Thermal cycling should be performed with a heated lid, except where specified otherwise. Thermal cycler(s) must support uniform heating up to 100 μL sample volume.

Revision Log

Version	Date	Description
V1.0	April 2024	Launch





Input Material

Please review and follow the guidelines below for preparing RNA-Seq libraries that are optimal for depletion:

The DepleteX Mito DNA Depletion Kit is optimized for 10 ng of NGS library material. Efficient depletion rates and high-quality NGS data have been validated with the use of the NEBNext® Ultra II FS DNA Library Prep Kit (Cat# E7805).

Accurate DNA quality assessment is recommended to maximize depletion efficiency and the efficiency of downstream steps. Validate input DNA using a fluorometric based method, such as a Qubit Fluorometer. Evaluate quality using an automated electrophoresis method, such as the Agilent Bioanalyzer System with a High Sensitivity DNA reagent kit.

Before Proceeding to Step A.

Please follow the guidelines below:

- Review the library preparation method(s). The libraries should have been prepared and stored under nuclease-free conditions.
- Ensure that the libraries are uniquely indexed.



USER MANUAL v1.0

Protocol

Step A: Depletion of Mitochondrial DNA

☐ Hands-on time: ~ 30 min | Total time: ~2 hours

Reagent Preparation

Item	Storage	Handling
DNA Library ≥ 200bp (10 ng)		
Cas9	-20°C	If frozen, thaw on ice. Finger-tap mix and spin down. Keep on ice. Return to freezer immediately
RNase Inhibitor		after use.
Guide RNA (Mito DNA)	-80°C	
Nuclease-Free Water		Thaw at room temperature.
10X Cas9 Buffer	-20°C	Vortex briefly and spin down. Keep on ice.
AMPure XP Beads	4°C	Bring to room temperature. Vortex and invert mix.
80% Ethanol	Room Temperature	Prepare fresh.

A1: Ribonucleoprotein (RNP) Complex Formation & CRISPR Digestion

1. Dilute the NGS libraries intended for depletion to a final quantity of 10 ng in 9 μL using Nuclease-Free Water. Set aside for CRISPR digestion (Step A1.5).

2. At room temperature, combine the following reagents in the order listed below in a 0.2 mL PCR tube:

RNP Complex Formation Reaction	Volume
Nuclease-Free Water	2.5 μL
10X Cas9 Buffer	1.0 μL
Cas9	2.0 μL
RNase Inhibitor	1.0 μL
Guide RNA (Mito DNA)	3.5 µL
Total Volume	10 μL

- 3. Mix gently by flicking the tube or pipetting up and down. Collect the contents at the bottom of the tube by briefly spinning the tube in a microcentrifuge.
- 4. Incubate the reaction mix at room temperature for 10 minutes. This constitutes the "Ribonucleoprotein (RNP) Complex."
- 5. To the tube containing the RNP complex, add the following reagents in the order listed below at room temperature:

CRISPR Digestion Reaction	Volume
RNP Complex (Previous Step A1.4)	10 μL
DNA Library (10 ng)	9 μL
10X Cas9 Buffer	1 μL
Total Volume	20 μL

6. Mix gently by flicking the tube or pipetting up and down slowly. Collect the contents at the bottom of the tube by briefly spinning the tube in a microcentrifuge.





7. Place the tube in a thermal cycler with a heated lid set to ≥80°C. Run the following program:

Temperature	Cycle Time
37°C	16 hours
4°C	HOLD

8. Following incubation, proceed to the next step (Bead Size Selection).

A2: Bead Size Selection

- 1. Add 30 μL of Nuclease-Free Water to the CRISPR digestion reaction from Step A1.7.
- 2. Add 40 μL (0.8X) of resuspended AMPure XP beads to the reaction. Mix well by pipetting up and down. Collect the contents at the bottom of the tube by briefly spinning the tube in a microcentrifuge.
- 3. Incubate at room temperature for 10 minutes. Mix the sample halfway through the incubation by pipetting up and down.
- 4. Place the tube on a magnetic stand until the solution is clear (3-5 minutes). Discard the supernatant without disturbing the beads.
- 5. Add 200 μL of freshly prepared 80% ethanol to the tube while still on the magnetic stand. Wait 30 seconds, then remove and discard the ethanol. Do not remove the tube from the magnetic stand while performing the ethanol wash.
- 6. Repeat the previous wash step.
- 7. Briefly spin the tube, place it back on the magnetic stand. Ensure that all residual ethanol is removed at this step.
- 8. Open the cap of the tube while it is on the magnetic stand and allow the beads to air dry for up to 5 minutes.
- 9. Remove the tube from the magnetic stand and add 40 μL of Nuclease-Free Water to the beads. Mix well to resuspend the beads by pipetting up and down. Collect the contents at the bottom of the tube by briefly spinning the tube in a microcentrifuge.
- 10. Incubate at room temperature for 10 minutes. Mix the sample halfway through the incubation by pipetting up and down. Collect the contents at the bottom of the tube by briefly spinning the tube in a microcentrifuge.
- 11. Place the tube on the magnetic stand. Allow the solution to clear and transfer the supernatant containing the eluted DNA to a new 0.2 mL PCR tube.
- 12. Proceed to the next step (PCR Amplification).
- SAFE STOPPING POINT: If stopping at this point in the protocol, store the sample at -20°C.





Step B: PCR Amplification

Hands-on time: ~30 min | Total time: ~1 hour

Reagent Preparation

ltem	Storage	Handling
2X PCR Mix	-20°C	If frozen, thaw on ice. Finger-tap mix and spin down. Keep on ice.
P7 Primer	20%	Thaw at room temperature. Vortex briefly and spin down. Keep on ice.
P5 Primer	-20°C	Vortex briefly and spin down. Keep on ice.
Tris Buffer	-20°C	Bring to room temperature.
AMPure XP Beads	4°C	Vortex and invert mix.
80% Ethanol	Room Temperature	Prepare fresh.

B1: PCR Amplification

1. On ice, combine the following reagents in the order listed below in a 0.2 mL PCR tube:

Amplification Reaction	Volume
Depleted DNA product (Step A2.11)	40 μL
2X PCR Mix	50 μL
P7 Primer	5 μL
P5 Primer	5 μL
Total Volume	100 μL

2. Place the reaction in a thermal cycler with a heated lid set to 105°C. Run the following program:

Temperature	Cycle Time	Number of cycles
95°C	2 min	1 cycle
98°C	20 sec	
55°C	30 sec	7 cycles*
72°C	30 sec	
72°C	2 min	
4°C	HOLD	1 cycle

^{*}The number of PCR cycles after depletion correlates inversely with DNA input. With the recommended input of 10 ng and 10 PCR cycles, a final total library yield of 100-300 ng is expected.

3. Proceed immediately to the next step (Bead Cleanup).

B2: Bead Cleanup

- 1. Add 80 μ L (0.8X) of resuspended AMPure XP beads to the reaction. Mix well by pipetting up and down. Collect the contents at the bottom of the tube by briefly spinning the tube in a microcentrifuge.
- 2. Incubate the sample at room temperature for 10 minutes. Mix the sample halfway through the incubation by pipetting up and down.
- 3. Place sample tube on a magnetic stand until the solution is clear (3-5 minutes). Discard the supernatant without disturbing the beads.
- 4. Add 200 μL of freshly prepared 80% ethanol to the tube while still on the magnetic stand. Wait 30 seconds, then remove and discard the ethanol. Do not remove the tube from the magnetic stand while performing the ethanol wash.
- 5. Repeat the previous wash step.





- 6. Briefly spin the tube, place it back on the magnetic stand. Ensure that all residual ethanol is removed at this step.
- 7. Open the cap of the tube while it is on the magnetic stand and allow the beads to air dry for up to 5 minutes.
- 8. Remove the tube from the magnetic stand and add 30 μL of Tris Buffer to the beads. Mix well to resuspend the beads by pipetting up and down. Collect the contents at the bottom of the tube by briefly spinning the tube in a microcentrifuge.
- 9. Incubate the sample at room temperature for 10 minutes. Mix the sample halfway through the incubation by pipetting up and down. Collect the contents at the bottom of the tube by briefly spinning the tube in a microcentrifuge.
- 10. Place the tube on the magnetic stand. Allow the solution to clear and transfer the supernatant containing the eluted DNA to a new DNA LoBind tube. This constitutes the depleted DNA library.



SAFE STOPPING POINT: If stopping, store the library at -20°C.

Subsequent Steps

Jumpcode Genomics recommends assessing the library yield using a dsDNA-specific fluorescence-based method (such as a Qubit fluorometer) and library size on an Agilent Bioanalyzer 2100 or equivalent instrument before sequencing for best results.

After the second (and final) bead cleanup, most DNA fragments in the depleted library should be larger than 200 bp. If so, the library can be loaded directly on an Illumina sequencer. If a large proportion of fragments are less than 200 bp, it is recommended that a gel-based size selection step be performed to isolate DNA between 200 bp and 500 bp in size. The DNA can be loaded on a sequencing instrument after gel purification.

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