

2. Perform PCR amplification as follows:

Step	Temperature	Time	Cycle
Initial Denaturation	95°C	5 min	1
Denaturation	95°C	15 sec	35
Annealing ²	55°C for controls	15 sec	
Extension ²	72°C	15 sec (for controls) 2-3 kb/min	
Final Extension	72°C	5 min	1

² Adjust Annealing temperature and Extension time for your Target-Specific primers.

3. Run 5 µl of each product on a 2% agarose gel. Confirm correct size of Target-Specific amplicon (variable) and each Control amplicon (expected size is 503 bp). Keep the remaining PCR products on ice or at -20°C until Part D. If non-specific amplifications appear, repeat the PCR using touchdown method.

Part C – sgRNA Synthesis

1. Prepare the following reactions on ice:

Component	Control	Target-Specific
sgRNA Control Oligo	1 µl	-
Target-Specific Oligo (10 µM)	-	1 µl
Scaffold Template and Primer Mix	1 µl	1 µl
MegaFi Pro Fidelity 2X PCR MasterMix	12.5 µl	12.5 µl
Nuclease-free Water	10.5 µl	10.5 µl

2. Perform PCR Amplification as follows:

Step	Temperature	Time	Cycle
Initial Denaturation	95°C	5 min	1
Denaturation	95°C	15 sec	35
Annealing	55°C	15 sec	
Extension	72°C	15 sec	
Final Extension	72°C	5 min	1

3. Run 5 µl of each product on a 2% agarose gel and expect 130 bp amplicons.

4. After correct bands are confirmed, prepare the following mix on ice:

Component	Control sgRNA	Target-Specific sgRNA
sgRNA Control Oligo PCR Product	4 µl	-
Target-Specific Oligo PCR Product	-	4 µl
sgRNA Synthesis Enzyme Mix	4 µl	4 µl
2X sgRNA Synthesis Buffer	10 µl	10 µl
Nuclease-free Water	2 µl	2 µl

5. Mix and centrifuge briefly. Incubate reactions at 37°C for 30 minutes to transcribe sgRNA.

6. Products can be used directly or stored at -80°C for long-term storage.

Part D – In vitro Cas9 Cleavage

1. Prepare the following reactions on ice:

Component	Control RNP Complex (Volume x 4) ³	Target-Specific RNP Complex ⁴ (Volume x 1)
Control sgRNA	4 µl	-
Target-Specific sgRNA	-	1 µl
spCas9 Nuclease Protein	8 µl	2 µl
Nuclease-free Water	28 µl	7 µl

³ Control RNP complex is multiplied by 4 (3 controls + 1 to account for pipetting error)

⁴ Target-Specific RNP complex mix recipe is for 1 reaction only. Adjust volumes accordingly depending on the number of pellets being tested.

2. Mix and centrifuge briefly. Incubate reactions at 37°C for 10 minutes to assemble the RNP complexes.

3. Prepare the following reactions on ice:

Component	Wild-Type Control	Monoallelic Control	Biallelic Control	Target-Specific
Wild-Type Control PCR product from Part B	4 µl	-	-	-
Monoallelic Control PCR product from Part B	-	4 µl	-	-
Biallelic Control PCR product from Part B	-	-	4 µl	-
Target-Specific PCR product from Part B	-	-	-	4 µl
10X Cas9 Reaction Buffer	2 µl	2 µl	2 µl	2 µl
Control RNP Complex from Part D	10 µl	10 µl	10 µl	-
Target-Specific RNP Complex from Part D	-	-	-	10 µl
Nuclease-free Water	4 µl	4 µl	4 µl	4 µl

4. Mix and centrifuge briefly. Incubate at 37°C for 1 hour.

5. Inactivate reactions by adding 1 µl of RNP Degradator into each tube. Incubate at 68°C for 15 minutes.

6. Run the assay products on a 2% agarose gel and assess the cleavage patterns.

Part E – Analysis of Cleavage Products

Analyze the cleavage of the controls. Controls should show the following cleavage patterns:

Sample	Expected Bands
Wild-Type Control	294 bp + 209 bp
Monoallelic Control	503 bp + 294 bp + 209 bp
Biallelic Control	503 bp

For Target-Specific samples, assess cleavage patterns based on PCR product length and sgRNA placement. spCas9 cleaves DNA 3 bases upstream of the PAM sequence. Wild-Type cell samples should fully cleave showing 2 bands. For CRISPR-edited cells:

- 1 band - **Biallelic editing**
- 2 bands - **Unedited WildType cells**
- 3 bands - **Monoallelic editing**

For a more detailed product manual, please follow the QR Code below:

