

**Catalog No.:** D300-1000Mix

**Product Name:** Tissue-Direct™ PCR Genotyping Kit (MIX ONLY)

**Description:** Tissue-Direct™ PCR Genotyping Kit contains all the reagents needed for quick preparation of animal genomic DNA and genotyping by direct PCR amplification. Tissues used for this kit can be mouse tail, ear pieces, nail tips and cultured cells, or any other animal tissues such as muscle, liver, hair shaft, skin, etc. The PCR product can be directly loaded onto agarose gel without addition of loading dye for visualization of the experimental results.

| Kit Contents:      | D300-100 | D300-1000 |
|--------------------|----------|-----------|
| Size:              | 100 rxns | 1000 rxns |
| DNA Prep Solution: | 10ml     | 2x50ml    |
| 2X Direct-PCR Mix: | 1ml      | 10x 1ml   |

**Storage:**  
or at -20°C for long-term.

The whole kit can be stored at 4°C for up to three months

### General Protocol

#### **I. DNA Sample Preparation:**

1. Place the sample into a PCR tube:
  - For mouse tissues from tail, ear or nail: 1-3 mm in length or diameter.
  - Animal tissues: 1-3mg.
  - Cultured cells: 10µl of cell culture.
  - Other samples: similar amount or volume as above.
2. Add 100µl of the **DNA Prep Solution** into the tube containing the sample.
3. Heat the sample for 10 minutes at 95°C in a PCR machine.
4. Take out the sample and mix a few times. The sample is now ready for PCR. **Optional:** the sample can be centrifuged briefly and use the supernatant for PCR. Avoid any undigested tissue or debris.

## II. PCR Amplification:

- To set up the PCR reaction, add the following reagents to a PCR tube or PCR plate, for example:

|                      |              |
|----------------------|--------------|
| 2X Direct-PCR Mix:   | 10µl         |
| Primers:             | 1µl          |
| Sample:              | 1µl          |
| Water:               | 8µl          |
| <b>Total volume:</b> | <b>20 µl</b> |

### Note:

- If needed, scale up your PCR according to your specific case, such as using 50 µl as PCR reaction volume. **However, always use half of your PCR volume for the 2X Direct-PCR Mix.**
  - When large numbers of samples are processed with same primers, the 2X Direct-PCR Mix, water and primers can be premixed and aliquoted.
- Mix the PCR reaction gently avoid creating foam or bubbles. A brief centrifugation may be needed to collect the reaction mix to the bottom of the PCR tube or plate.
  - Perform the PCR thermal cycling. The following table is a typical example of a PCR.

| Step             | Temperature | Time      | Cycles |
|------------------|-------------|-----------|--------|
| Initial Denature | 95°C        | 3 min     | 1      |
| Denature         | 95°C        | 0.5-1 min | 30-35  |
| Annealing        | 50-65°C     | 0.5-1 min |        |
| Extension        | 72°C        | 1 min/kb  |        |
| Final Extension  | 72°C        | 7 min     | 1      |
| Hold             | 4°C         | ∞         |        |

- After PCR, The amplified products can be directly loaded onto an agarose gel for checking the results.

## Troubleshooting: Problems and Solutions

**Q1.** Samples are not completely digested or dissolved.

**A1.** Samples are not expected to be digested or dissolved completely. Do not worry. Sufficient DNA will be released for PCR without complete digestion of the samples.

**Q2.** Little or no PCR product is detected.

### **A2.** General Solution:

- Make sure that there are no PCR components missed.
- More PCR cycles may be needed.
- Primers may not be designed optimally.

- d) Try different annealing temperature and extension time or use a touchdown PCR program.
- e) Too much sample may have been used, in that case the samples can be easily diluted 10 times with H<sub>2</sub>O or 10mM Tris-HCl buffer, pH 8.5.
- f) If you have tried all above and it is still not working, read the Specific Solution below.

**A2. Specific Solution:** Some templates and primers are difficult for PCR. If you have tried changing all the possible PCR parameters and your PCR is still not working well, we recommend that you try our **Conquest™ Genotyping PCR Optimizing Kit (Cat. No. D911)**. This kit has successfully performed PCR on many tough PCR templates and primers. The Master Mixes within the **Conquest™ Genotyping PCR Optimizing Kit** are formulated with multiple thermal DNA polymerases and pre-optimized PCR buffers and enhancers. It is in four different ready-to-use 2X format combinations. Using this **Conquest™ Genotyping PCR Optimizing Kit** simplifies the process to determine optimal conditions for each unique combination of template and primers. You will easily find a specific Master Mix that works well with your templates and primers.

- Q3.** High background or multiple PCR products.
- A3.** Adjust your annealing temperature or use touchdown PCR program.
- Q4.** Negative control shows PCR product or false positive result.
- A4.** Reagents or your samples may be contaminated.

END