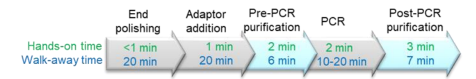


# ChIP-Seq Library Prep Kit (MGI Platform)

| Catalog No. | 34034S       | 34034L       |
|-------------|--------------|--------------|
| Index type  | index        | index        |
| Reactions   | 24 reactions | 96 reactions |

# **Description**

The **ChIP-Seq Library Prep Kit** (MGI Platform) was developed for the construction of high-quality libraries using 5 ng to 400 ng of ChIP DNA as input. The kit is compatible with ChIP DNA fragments generated from both enzymatic methods and mechanical methods (sonication, nebulization etc.).



#### Features

- Fast
  - Total time: 1.5 hrs
  - Hands-on time: 10 min
- Easy procedure
  - Ready-to-use master mix
  - Less reaction components
- Less magnetic beads required: Reduced more than 50%
- Guaranteed quality: Higher library conversion efficiency
- Input ChIP DNA: From 5 ng to 400 ng

### Component

| Catalog No.               | 34034S   | 34034L   |
|---------------------------|----------|----------|
| CS1 Buffer                | 72 ul    | 288 ul   |
| CS1 Enzyme                | 48 ul    | 192 ul   |
| CS2M Buffer               | 336 ul   | 1344 ul  |
| CS2 Enzyme                | 24 ul    | 96 ul    |
| Sodium Chloride (1.67 M)* | 432 ul   | 1728 ul  |
| Primers                   | 5 ul X24 | 5 ul X96 |
| PCR mix                   | 600 ul   | 2400 ul  |

\*Sometimes the tube of Sodium Chloride (1.67 M) may crack during dry ice shipping. Customers need to prepare Sodium Chloride (1.67 M) in this case.

### **Storage Condition**

• Store kit at -20°C, stable up to 12 months.

### Reagent & Equipment Needed (not provided in this kit)

- Magnetic particle concentrator
- PCR thermal cycler
- 96-well PCR plate
- 80% ethanol (prepare before use)
- Sodium Chloride Solution (1.67 M)\*
- Magnetic Beads (BioDynami Cat.# 40051) or equivalent



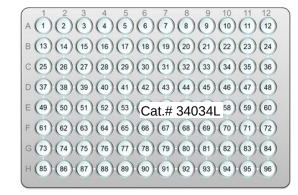
# Library and Index Information

Index (Cat.# 34034S and 34034L)

| Index ID | Index Sequence | Index ID  | Index Sequence | Index ID  | Index Sequence |
|----------|----------------|-----------|----------------|-----------|----------------|
| Index #1 | GCTTGTTCAG     | Index #9  | CCGATGACGT     | Index #17 | GACGCGGTAT     |
| Index #2 | AACAAGCACT     | Index #10 | TTATCTCGAG     | Index #18 | ACGAGACGTC     |
| Index #3 | TTGCCAGTGA     | Index #11 | AGCCGATACC     | Index #19 | CTACTCAAGA     |
| Index #4 | CGAGTCAGTC     | Index #12 | GATGACGTTA     | Index #20 | TGTTATTCCG     |
| Index #5 | GATAGTAACG     | Index #13 | TCGCGATGTC     | Index #21 | CCGTCACTGA     |
| Index #6 | TGAGTGGCTA     | Index #14 | AGTGACACCA     | Index #22 | TGACGCAACT     |
| Index #7 | CCGTCATTAC     | Index #15 | GACATTCAAG     | Index #23 | GTTGTTGCTC     |
| Index #8 | ATCCACCGGT     | Index #16 | CTATCGGTGT     | Index #24 | AACAAGTGAG     |

For Cat.# 34034S, primers will be shipped in 8-stripe PCR tubes with index labels at both ends as shown below. For Cat.# 34034L, primers will be shipped in 96-well plates. Below is the index layout.







# Protocol

# Step 1: End polishing

1) Add the following to one well of a 96-well PCR plate:

| ChIP DNA   | 10 ul (5 ng~400 ng) |  |  |
|------------|---------------------|--|--|
| CS1 Buffer | 3 ul                |  |  |
| CS1 Enzyme | <u>2 ul</u>         |  |  |
| Total      | 15 ul               |  |  |

- 2) Mix by pipetting ten times.
- 3) Incubate at 20°C for 15 min, 70°C for 5 min. Proceed immediately to step 2.

# Step 2: Adaptor addition

1) Add the following to Step 1 reaction mixture. Slow pipetting of the viscous CS2M Buffer is needed for precise aliquot.

| CS2M Buffer | 14 ul |
|-------------|-------|
| CS2 Enzyme  | 1 ul  |
| Total       | 15 ul |

- 2) Mix by pipetting ten times.
- 3) Incubate at 20°C for 20 min.
- 4) Add Sodium Chloride (1.67 M) 18 ul to the reaction mixture. Proceed immediately to step 3.

# Step 3: Pre-PCR purification

- 1) Resuspend Magnetic Beads and transfer 40 ul to the above reaction mixture, mix by pipetting and incubate for 3 min.
- 2) Load the plate on a magnet, incubate for 2 min, and discard the supernatant carefully. Remove all residual supernatant without disturbing the beads.
- 3) Add 180 ul of 80% ethanol, incubate for 30 sec, and discard the supernatant carefully. Remove all residual ethanol without disturbing the beads.
- 4) Remove the plate from the magnet, resuspend the beads in 22 ul of water.
- 5) Load the plate on the magnet, incubate for 1 min, and transfer 20 ul supernatant (containing library) to a new tube without disturbing the beads.

### Step 4: PCR

1) Mix the following in a PCR plate:

| Library | 20 ul |
|---------|-------|
| Primers | 5 ul  |
| PCR mix | 25 ul |
| Total   | 50 ul |

2) Put PCR plate on a thermal cycler, start PCR with the following condition:

| Step                 | Temperature | Time       | Cycles    |   |
|----------------------|-------------|------------|-----------|---|
| Initial Denaturation | 98°C        | 30 seconds | 1         | <ul> <li>*** As a reference:<br/>10-13 cycles for 5-10 ng input;<br/>8-11 cycles for 10-20 ng input;<br/>7-10 cycles for 20-50 ng input;<br/>6-9 cycles for 50-400 ng input.</li> </ul> |
| Denaturation         | 98°C        | 10 seconds | 6-13      |   |
| Annealing/extension  | 65°C        | 70 seconds | cycles*** |   |
| Final Extension      | 65°C        | 2 minutes  | 1         |   |
| Hold                 | 4°C         |            |           |   |

# Step 5: Post-PCR purification

- 1) Resuspend Magnetic Beads and transfer 40 ul to the above reaction mixture, mix by pipetting and incubate for 3 min.
- 2) Load the plate on a magnet, incubate for 2 min, and discard the supernatant carefully.
- 3) Add 180 ul of 80% ethanol, incubate for 30 sec, and discard the supernatant carefully. Remove all residual ethanol without disturbing the beads.
- 4) Remove the plate from the magnet, resuspend the beads in 22 ul of water or Tris-HCl (10 mM).
- 5) Load the plate on the magnet, incubate for 1 min, and transfer 20 ul supernatant (containing library) to a new tube without disturbing the beads.

# **Quality Control**

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Kit components passed stringent functional quality test.

#### **Product Use Limitation**

This product is developed and sold for research purposes and *in vitro* use only. Please refer to BioDynami.com for Material Safety Data Sheet of the product.

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#### About PCR master mix:

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