

DNA Normalization Magnetic Beads (50-300 ng Input)

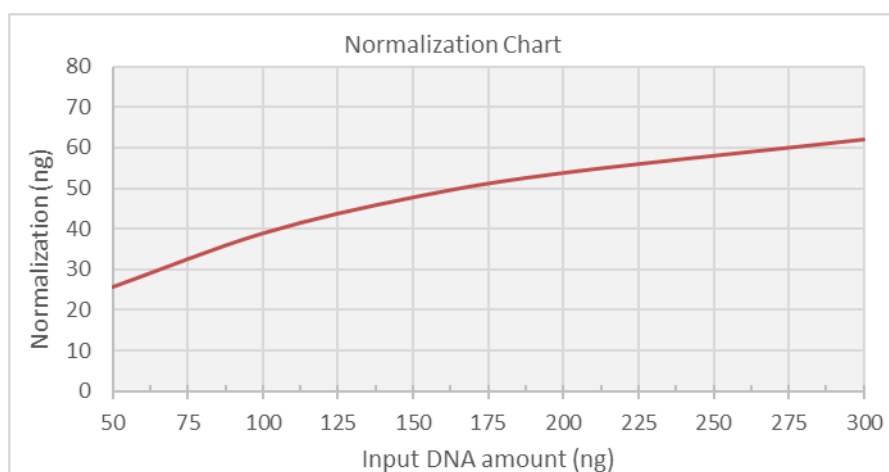
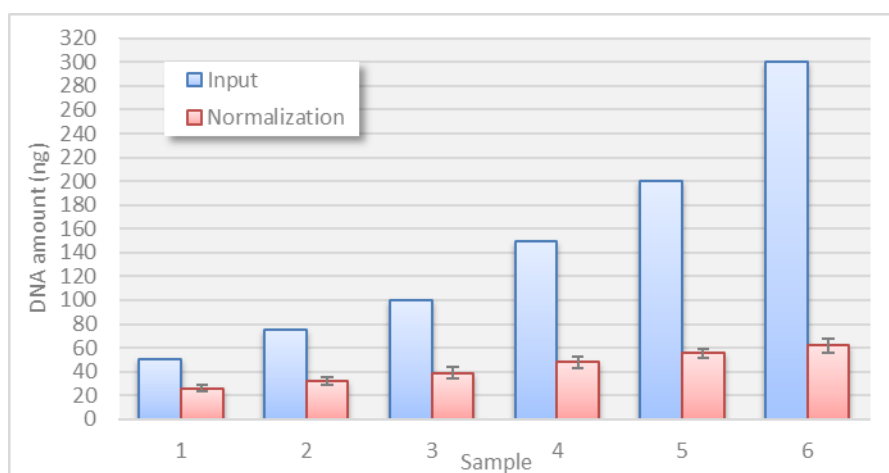
Catalog No.	40075S	40075L
Rxns*	48	96

*Based on 20 µl of sample volume

Description

The **DNA Normalization Magnetic Beads (50-300 ng Input)** was developed for normalization of NGS libraries, PCR fragments, and sheared genomic DNA based on our magnetic bead technology. Magnetic beads has DNA-binding capacity. With the limitation of the DNA-binding capacity, normalized amount of DNA is recovered, and excess unbound DNA is separated.

The normalized DNA concentration can be obtained from DNA samples of various concentrations. Traditional fluorescent methods of DNA quantification require the setup of a standard curve from known DNA concentrations, compare with DNA samples from various sources, which would need extensive efforts and operating reagents. The protocol is simple, fast, centrifugation free, and filtration free. Additional DNA quantification and dilution is not necessary after DNA normalization with magnetic beads. Labor, time, and reagent are saved with the normalization beads.



Specification

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DNA input range	50-300 ng
DNA input type	NGS library, PCR fragment, sheared genomic DNA
Normalized range	25-60 ng
Elution volume	> 10 µl
Technology	Magnetic beads

Features

- Consistent normalization of various sources
 - NGS libraries
 - PCR fragments
 - Sheared genomic DNA
- Excellent dynamic input range: 6 folds from 50 ng to 300 ng
- Magnetic beads technology
 - No centrifugation
 - No column
 - No filtration
- Elution: as low as 10 µl without yield loss
- Removal of unwanted components and impurities

Component

Catalog No.	40075S	40075L
PG8 Buffer	700 µl	1400 µl
MK1 Buffer	96 µl	192 µl
B3M Buffer	96 µl	192 µl
SS7 Beads	96 µl	192 µl
Elution Buffer	960 µl	1920 µl

Storage Condition

- Store at 4°C, stable up to 6 months.

Reagent & Equipment Needed (not provided in this reagent)

- Magnetic particle concentrator
- 96-well plates
- 100% ethanol and 80% ethanol (prepare before use)
- Thermal cycler

Protocol

Note:

- Store DNA samples at 4°C before use.
 - DNA samples without buffer are preferred. PCR buffer may be fine depending on the specific buffer composition.
- 1) Prepare **Normalization Beads** (20 µl per sample) by adding the following to a tube/bottle, and mix by vortexing the Normalization Beads thoroughly. **Attention:** vortex or shake the **SS7 Beads** tube to resuspend the beads thoroughly just before use.

PG8 Buffer	14 µl
MK1 Buffer	2 µl
B3M Buffer	2 µl
<u>SS7 Beads</u>	<u>2 µl</u>
Total	20 µl

- 2) Transfer 20 µl of the Normalization Beads to the wells of a 96-well plate, incubate the plate on ice for 3 min. Slow pipetting of the viscous Normalization Beads is needed for precise aliquot.
- 3) Add 20 µl of DNA samples, mix by pipetting gently and thoroughly with a multichannel pipettor. Remove the plate from the ice, load the plate on a thermal cycler, and incubate at 22°C for 10 min.
- 4) Load the sample on a magnet, incubate for 5 min.
- 5) Add 150 µl of **100% ethanol**, incubate for 2 min, and **slowly** remove the supernatant without disturbing the pellet.
- 6) **Slowly** add 200 µl of **80% ethanol** without disturbing the beads. Incubate for 2 min and remove the supernatant **slowly and carefully** with a multichannel pipettor. Remove residual ethanol **carefully** with fine tips.
- 7) Remove the plate from the magnet and resuspend the beads in at least 10 µl of **Elution Buffer** (10 mM Tris-HCl). Make sure all beads are suspended in the Elution Buffer. DNA can also be eluted in water (pH>6.0), low TE buffer, or TE Buffer. Resuspension of the beads in less than 10 µl may reduce the yield and generate more yield variation.
- 8) Load the plate on the magnet, incubate for 1 min, and transfer the supernatant (containing normalized DNA samples) to a new plate or tubes without disturbing the beads.

Quality Control

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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