

DNA Size Selection Kit (100-200 bp, Magnetic Beads)

Catalog No.	20102S	20102L
Runs*	24	96

*Based on 50 µl of sample volume

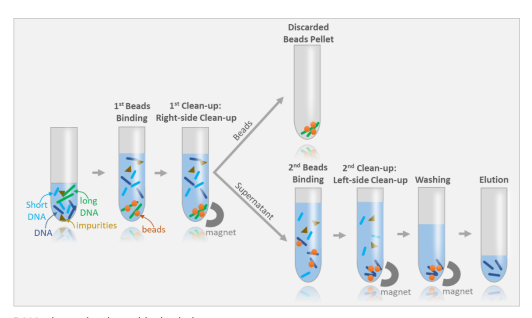
Description

The **DNA Size Selection Kits (100-200 bp, Magnetic Beads)** were developed for DNA size selection using magnetic beads with a selection range spanning from 100 bp to 200 bp. The kits provide a simple and quick approach for the enrichment of a specific range of DNA fragments. The kit workflow allows double-sided size selection for the specific size cutoffs.

The magnetic beads technology uses paramagnetic particles, also known as SPRI (Solid Phase Reversible Immobilization) beads, to bind DNA reversibly and selectively. DNA fragments can be size-selected and purified by changing the properties of the magnetic beads or SPRI beads. The magnetic beads can easily separate the beads-binding DNA from the contaminants and unwanted components in the samples.

DNA size selection is a selective capture of DNA fragments of a specific range of size for next-generation sequencing (NGS) library preparations, PCR, ChIP assay, DNA ligations, endonuclease digestions, adapter removal, and other genomics and molecular biology applications.

Specific DNA fragments at a certain length range can be purified simply using magnetic separation with different beads components, avoiding tedious and time-consuming gel extraction and column-based purification. The first beads-binding step, referred to as the right-side clean-up, removes large DNA fragments. The large DNA fragments are bound to the beads and are discarded. The desired DNA fragments in the supernatant are transferred to a new well, and new beads are added to the supernatant for the second beads-binding, referred to as the left-side clean-up. The double-size selected DNA fragments are eluted after ethanol rinsing.



DNA size selection with dual clean-ups.



Features

- High specificity of size selection: 100-200 bp
- Fast and simple
 - o 20-min protocol
 - o No gels required
 - o No columns required
 - o No centrifugation required
- High recovery of selected DNA fragments
- · Consistent performance: rapid size selection with high reproducibility

Component

Catalog No.	20102S	20102L
D2 Beads	0.96 mL	3.84 mL
F2 Beads	0.96 mL	3.84 mL
Elution Buffer	0.96 mL	3.84 mL

Storage Condition

Store at 4°C, stable up to 12 months.

Reagent & Equipment Needed (not provided in this reagent)

- Magnetic particle concentrator
- 96-well plates or microfuge tubes
- 80% ethanol (prepare before use)

Protocol

Note:

- Bring the beads to room temperature before using.
- Invert or shake the beads bottles thoroughly to resuspend the beads before using.
- Accurate pipetting is needed for precious size selection.
 - 1) Add 40 μl of the D2 Beads to the wells containing 50 μl of samples in a 96-well plate. Slow pipetting of the viscous beads is needed for precise aliquot. Mix by pipetting gently and thoroughly with a multichannel pipettor. Incubate for 5 min. **Note**: Small sample volumes tend to generate more variations. If the sample volume is less than 50 μl, increase the volume to 50 μl by adding nuclease-free water.
 - 2) Load the sample on a magnet, incubate for 3 min, and transfer the supernatant (containing the sample) to a new 96-well plate. Discard the beads pellet.
 - 3) Add 40 µl of the F2 Beads to the wells containing samples. Slow pipetting of the viscous beads is needed for precise aliquot. Mix by pipetting gently and thoroughly with a multichannel pipettor. Incubate for 5 min.
 - 4) Load the sample plate on a magnet, incubate for 3 min, and discard the supernatant carefully.
 - 5) Add 200 µl of 80% ethanol without disturbing the beads. Incubate for 1 min and discard the supernatant carefully. Remove all residual ethanol without disturbing the beads.
 - 6) Remove the plate from the magnet and resuspend the beads in at least 20 µl of water, Elution Buffer (10 mM Tris-HCl), or low TE buffer to elute DNA from the beads. **Note**: Resuspension of the beads in less than 20 µl will reduce the yield. A brief centrifugation step may improve bringing eluates to the bottom of the wells before placing on the magnet.
 - 7) Load the plate on the magnet, incubate for 1 min, and transfer the supernatant (containing the size-selected sample) to a new plate or tubes without disturbing the beads.



Quality Control

Magnetic beads components passed stringent functional quality test.

Product Use Limitation

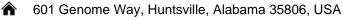
This product is developed and sold for research purposes and *in vitro* use only. Please refer to biodynami.com for Material Safety Data Sheet of the product.

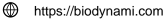
Limited Label License

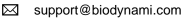
The product is developed and sold exclusively for research purposes and *in vitro* use only. The product or its any individual component has not been tested for use in diagnostics or drug development, and is not suitable for administration to human or animal.

The purchaser of this product is granted a limited, non-transferable right to use the purchased amount of the product only for internal, research purposes for the sole benefit of the purchaser. The buyer cannot sell or otherwise transfer (i) this product (ii) its components or (iii) materials made using this product or its components to a third party or otherwise use this product or its components or materials made using this product or its components for commercial purposes. This product is for internal research purposes only and is not for use in commercial purposes of any kind. "Commercial purposes" includes any activity for which a party receives consideration and may include, but is not limited to, (1) use of the product or its components or derivatives in manufacturing, (2) transfer or sale of vectors made with the product or components or derivatives of the product, (3) use of this product or components or derivatives of the product made therefrom to provide a service, information, or data to a third party in return for a fee or other consideration, or (4) resale of the product or its components or derivatives, whether or not such product or its components or derivatives are resold for use in research. If the purchaser is not willing to accept the limitations of this limited use statement, BioDynami is willing to accept return of the products with a full refund. For information on obtaining additional rights, please contact support@biodynami.com

BioDynami









Version 1.0 (Jul. 2024)