

# Anti-Myc Magnetic Particles, 200 nm PRODUCT DATA SHEET

# Anti-Myc Magnetic Particles, 200 nm

Cat No: AM-Anti-Myc-20

## Description

Anti-Myc magnetic beads are 200 nm nano magnetic beads that covalently bind a large number of high-quality murine anti-Myc monoclonal antibodies on the surface. Nano-sized magnetic beads provide a large specific surface area, have more binding sites, use less magnetic beads, and have a low non-specific adsorption rate. This product is widely used in the immunoprecipitation (IP) or immunoco-precipitation (Co-IP) with Myc label fusion protein in cell lysate, cell secreted supernatant, serum, ascites and other samples. Due to the magnetic separation, it can save up to 40% of the time per IP and Co-IP.

For custom sizes, formulations or bulk quantities please contact our customer service department.

Website: www.abvigen.com Phone: +1 929-202-3014 Email: info@abvigenus.com

#### Characteristics

Matrix	Silicon based magnetic particles
Ligand	Murine monoclonal antibody against Myc
Particle size	200 nm
Concentration	10 mg/mL
Binding ability	≥ 0.5 mg Myc tag fusion protein/mL magnetic particles
Scope of application	IP, Co-IP, etc
Shelf life	Stable storage at 2 ~ 8°C for 2 years

#### **Operation process**

Note: Prepare your own buffer or purchase our IP kit directly.

# Adherent cell sample:

1. Remove the medium and wash the cells twice with PBS.



- 2. Cells were collected into 1.5 mL EP tube and IP Lysis/Wash Buffer was added proportioned, and add corresponding inhibitor such as PMSF. After mixing, the cells were placed on ice for  $5 \sim 20$  min (mixed several times).
- 3. At 4°C,  $12000 \sim 16000 \text{ xg}$ , centrifuge for 10 min to collect the supernatant and place it on ice for subsequent experiments (or place it at -80°C for long-term storage).

# Petri dish IP Lysis/Wash Buffer Recommended volume:

Petri dish size/surface area	Volume of IP Lysis/Wash Buffer
100 mm x 100 mm	500 ~ 1000 μL
100 mm x 60 mm	100 ~ 300 μL
6-well plate	100 ~ 200 μL

## Suspended cell sample:

- 1. Cells were collected at 4°C, 500 ~ 1000 xg, 10 min, and the supernatant was discarded.
- 2. Wash the cells with PBS once, that is, suspend the cell mass with PBS at  $4^{\circ}$ C,  $500 \sim 1000$  xg, 5 min, collect the cells, discard the supernatant.
- 3. Resuspend cells with pre-cooled IP Lysis/Wash Buffer. The 500  $\mu$ L IP Lysis/Wash Buffer was used for every 50 mg cells. At the same time, add the corresponding inhibitors such as PMSF, mix well and put on the ice for 5  $^{\sim}$  20 min (mix several times during the period).
- 4. At  $4^{\circ}$ C,12000  $\sim$  16000 xg, centrifuge for 10 min to collect the supernatant and place it on ice for subsequent experiments (or place it at -80°C for long-term storage).

# Serum sample:

It is recommended to dilute a serum sample with IP Lysis/Wash Buffer to a final concentration of target protein of 50 to 150  $\mu$ g/mL and place it on ice for future use (or place it at -20°C for long-term storage).

#### Immunoprecipitation:

Note: In order to ensure the uniform distribution of magnetic beads, mix the magnetic beads in the bottle by repeatedly reversing or slightly vortexing before use.

- 1. Add 20 to 50 μL of Anti-Myc magnetic bead into 1.5 mL centrifuge tube.
- 2. Add 500  $\mu$ L pre-cooled PBS to the magnetic bead and mix gently.
- 3. Put the centrifugal tube into the magnetic rack and collect magnetic beads to one side of the centrifugal tube. Remove the supernatant.



- 4. Add 200  $^{\sim}$  500  $\mu$ L IP Lysis/Wash Buffer to the centrifuge tube. Reverse the centrifugal tube several times or mix with a slight vortex for 1 min. Use a magnetic rack to collect magnetic beads. Remove the supernatant.
- 5. Add the protein sample containing the Myc label into the centrifuge tube fitted with magnetic beads and incubate at room temperature for  $1 \sim 2$  h, or  $4 \circ C 2 \sim 4$  h.
- 6. Collect magnetic beads with a magnetic rack, remove uncombined samples, and store them for analysis.
- 7. Add 1000  $\mu$ L IP Lysis/Wash Buffer to the centrifuge tube and gently mix the magnetic ball for 5  $^{\sim}$  10 min. Collect the beads and discard the supernatant. Repeat twice.
- 8. **Denaturing elution:** Add 80  $^{\sim}$  100  $\mu$ L SDS-PAGE Sample Loading Buffer (1×) to the centrifuge tube and heat the sample in 100°C water bath or metal bath for 10 min. Magnetic beads are separated by magnetic rack to retain the supernatant containing the target antigen.

**Note:** The following elution methods can also be used to maintain protein activity.

Low pH Elution: Add 100  $\mu$ L Elution Buffer to centrifuge tube. Keep mixed and incubate centrifuge tube at room temperature for 5  $^{\sim}$  10 min. Magnetic beads are magnetically separated to retain the supernatant containing the target antigen. Low pH is neutralized by adding 20  $\mu$ L Neutralization buffers per 100  $\mu$ L of wash out solution.

#### Notes

- 1. Before performing the experiment, please read this operation manual carefully.
- 2. Do not centrifuge, dry or freeze magnetic beads at high speed, these operations will cause magnetic beads to gather and reduce the binding ability.
- 3. The affinities between different types of antibodies and antigen Lysis are different in IP experiments. The binding of antibodies to antigens is also affected by IP Lysis/Wash Buffer, so you can do your own experiments by optimizing the operation details or screening and formulating buffers.
- 4. Magnetic beads should be fully oscillated and uniform before use. Magnetic beads should be kept in storage solution to prevent drying.
- 5. This product is for scientific research only.

#### **Problem solving**

1. No immunoprecipitation of antigen

1 The sample contains too little antigen to be detected



Recommendation: To verify the expression and/or cleavage efficiency of proteins in the lysate by SDS-

PAGE or protein immunoblotting; If necessary, increase the sample size

(2) The ingredients in IP Lysis/Wash Buffer interfere with the binding of antigens to antibodies

Recommendation: Immunoprecipitation and rinsing with other buffers (e.g., TBS containing 0.5% CHAPS).

2. Low protein intake

(1) The protein is degraded

Recommendation: Add protease inhibitor

(2) The amount of magnetic beads used is not enough

Recommendation: Increase the amount of magnetic beads used to capture immune complexes

(3) The amount of target protein in the sample is not enough

**Recommendation:** Increase the amount of antigen sample

3. Multiple non-specific bands

There are non-specific proteins that bind to the magnetic beads

Recommendation: Add 50 ~ 350 mM NaCl to IP Lysis/Wash Buffer. Increase elution intensity and

frequency.

4. Magnetic beads gather

The aggregation of magnetic beads in the low pH Elution Buffer is a normal phenomenon and does

not affect the normal use of magnetic beads.

Recommendation: The magnetic beads can be restored to uniform state by using IP Lysis/Wash Buffer

to neutral and then oscillating with Tris buffer (pH 7.5) containing 0.1% (v/v) Tweene-20 and treated

with ultrasonic water bath. None of the above treatments affect the antibody binding efficiency of the

magnetic beads. A non-ionic detergent with a final concentration of 0.1% (v/v), such as Tween-20 or

Triton X-100, can also be added to effectively prevent the accumulation of magnetic beads.

Note: Ultrasonic treatment will also remove antibodies captured by magnetic beads in the sample

solution, so this method should not be used before elution of magnetic beads after sample addition.

**Ordering Information** 

Website: www.abvigen.com

Phone: +1 929-202-3014

Email: info@abvigenus.com