

[Cat. No.] TA-1016-1

Introduction

Bioneer AccuNanoBead Biotin Magnetic Nanobeads are uniform, silica-based paramagnetic nanoparticles are coupled with a biomolecule, such as Biotin, and are utilized in the magnetic separation and isolation of avidin and streptavidin-labeled components. The particles have a large surface area with high capture efficiencies.

Features & Benefits

AccuNanoBead Biotin Magnetic Nanobeads are densely coated with Biotin. Nanobeads are utilized in the magnetic separation of avidin and streptavidin-labeled molecules. Biotin magnetic beads are stable, pre-blocked beads with high binding capacity that provide rapid and efficient biomolecule purification from complex samples. Design of Bioneer AccuNanoBead Biotin Magnetic Nanobeads enables faster binding kinetics, High yield, purity, and quality in many biomedical and research applications.

Components

Components	Amount
AccuNanoBead™ Biotin Magnetic NanoBeads	0.5g/25ml In DW

* **Note:** For research use only. Not for use in diagnostic or therapeutic procedures.

Materials to be Prepared by User

Magnetic Separator	
Binding/Wash Buffer	TBS – 0.05% Tween 20 detergent
Elution buffer	8M guanidine-HCL, Ph 1.5
Deionized Water	

* **Note:** Buffer could be changed depending on user's needs.

Specifications

AccuNanoBead™ Biotin Magnetic NanoBeads	
Composition	Biotin based magnetic nanobeads
Binding capacity	HABA/Avidin Loading : $\geq 7\mu\text{mol/g}$ of beads
Size	Average 400 nm
Concentration	0.5g/25ml

Storage

Store at 2~6°C. This product can be stable for 3 years at 2~6°C

Expired date

Indicated on the label.

Precautions

Do not vigorously vortex AccuNanoBead™ Biotin Magnetic NanoBeads. An exact protocol may need to be optimized by the user

Online Resources



Korean



English

Visit our product page for additional information and protocols

Ordering Information

Description	Cat. No.
AccuNanoBead™ Biotin Magnetic NanoBeads	TA-1016-1

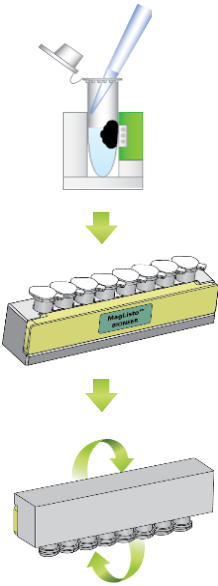
Notice

BIONEER corporation reserves the right to make corrections, modifications, improvements and other changes to its products, services, specifications or product descriptions at any time without notice.

Explanation of Symbols

Batch Code	Catalog Number	Caution	Consult Instructions For Use
Contains Sufficient for <n> tests	Do not Re-use	Manufacturer	Research Use Only
Temperature Limitation	Use-by Date		

Experimental Procedures (The protocols are scalable and can be optimized)

	Steps	Procedure Details
1	Procedures 	1. Add 100 μL (0.5 mg) of beads to 1 mL of binding buffer in each tube to wash particles. 2. Magnetically separate using a magnetic separator for 2 minutes or until the supernatant is clear. 3. Remove the supernatant and wash once more by adding 1 mL of binding buffer. 4. Repeat step 2 and remove the supernatant. 5. Resuspend beads by adding 450 μL of binding buffer. 6. Add 50 μL of serum or cell culture supernatant to the beads. 7. Gently mix using vortex or rotator for 30 minutes. 8. Magnetically separate using a magnetic separator for 2 minutes or until the supernatant is clear. 9. Remove supernatant and wash with 0.5 mL Binding/Wash buffer to remove unbound proteins. 10. Repeat steps 8 and 9 once more. Remove supernatant. 11. Add 100 μL of elution buffer to beads and mix well. 12. Incubate at room temperature for 10 minutes with occasional gentle mixing or vortex. 13. Desalt or dialyze the eluted sample to put them into a suitable buffer.