



Gold Nanoparticles-GALG (H+L) PRODUCT DATA SHEET

Gold Nanoparticles-GALG (H+L)

Description

Goat anti-llama secondary antibodies are produced in goat and target llama IgG antibodies, along with their subclasses. This antibody-antibody interaction allows for the development of bio-assays and detection of llama antibody-labelled targets. Abvigen affinity isolated anti-Llama IgG antibody produced in goat and coupled to gold nanoparticles. Suitable for use in applications such as lateral flow, immunoblotting, light microscopy, and electron microscopy applications procedures for secondary detection of Llama antibody labeled samples.

Product List

Cat No	Product Name	Concentration	Size
ABGN-5-GALG (H+L)	Gold Nanoparticles, 5 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-10-GALG (H+L)	Gold Nanoparticles, 10 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-15-GALG (H+L)	Gold Nanoparticles, 15 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-20-GALG (H+L)	Gold Nanoparticles, 20 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-30-GALG (H+L)	Gold Nanoparticles, 30 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-40-GALG (H+L)	Gold Nanoparticles, 40 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-50-GALG (H+L)	Gold Nanoparticles, 50 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-60-GALG (H+L)	Gold Nanoparticles, 60 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-70-GALG (H+L)	Gold Nanoparticles, 70 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-80-GALG (H+L)	Gold Nanoparticles, 80 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-90-GALG (H+L)	Gold Nanoparticles, 90 nm-GALG (H+L)	OD 3	0.5 mL
ABGN-100-GALG (H+L)	Gold Nanoparticles, 100 nm-GALG (H+L)	OD 3	0.5 mL



Characteristics

Core size range: 5 nm ~ 100 nm

Concentration: OD=3

Conjugated antibody: Goat affinity purified anti-Llama IgG (H+L)

Clonality: Polyclonal

Working dilution: 1:10 ~ 1:100 (application dependent, optimization might be required)

Storage buffer: 20 mM Tris (pH 8.0), 150 mM NaCl, 20% glycerol (v/v), 1% BSA

Applications

Gold conjugates are suitable for use in lateral flow immunoblotting, light microscopy, and electron microscopy applications.

Standard Immunogold Dot-Blot Protocol

(Adapted from Moeremans et al.)

1. Spot one microlitre drops of a serial dilution of your protein (1 ug ~ 1 ng) in PBS supplemented with 0.5 µg/mL of BSA on nitrocellulose or PVDF membrane.
2. Let protein drops dry into the membrane.
3. Block Membrane for 30 min using 1% (w/v) dry milk in 1X PBS at room temperature.
4. Incubate with primary antibody for 2 h at room temperature.
5. Wash membrane 3x5 min with blocking solution prepared as above.
6. Incubate for 2 h (or longer for increased sensitivity) with secondary gold conjugate diluted 1:10 (OD=0.3) times with blocking solution (0.2% Blocking Solution).
7. Wash 3x5 min as above.
8. Dry membrane and record data.
9. (OPTIONAL) Proceed with silver enhancement to improve sensitivity.

Storage and Stability

Store undiluted in storage buffer at 2-8°C. Stable for 4 months if stored as specified.

DO NOT FREEZE.

Notes

This product is for R&D use only, not for drug, household, or other uses.



Ordering Information

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