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Datasheet for ABIN1721114 MagSi-S Hydrazide 600 beads

2 Images



Overview		
Quantity:	2 mL	
Application:	Separation (Sep)	
Product Details		
Purpose:	Superparamagnetic silica particles with hydrazide modified surface. Intended for immobilization of antibodies and glycoproteins using aldehyde coupling chemistry with specific orientation.	
Characteristics:	Coating chemistry: Hydrazide Bead concentration: 8 - 20 x 10^9 beads/mL	
Components:	Magnetic silica beads with activated surface	
Material not included:	Buffers and Materials (depending on the application) Magnetic separator for bead separation/collecting Mixer/vortex to homogenize samples and resuspend beads (depending on the application)	
Bead Ligand:	Hydrazide modified surface	
Bead Matrix:	Magnetic Silica particles	
Bead Size:	600 nm	

Application Details

Comment:	MagSi-Tools are surface activated magnetic particles, intended for covalent immobilization of
	proteins (e.g. antibodies, enzymes), peptides, nucleic acids or other molecules of interest.
	Different surface modifications and bead sizes allow for choosing the optimal product for the
	right molecule to be coupled, and for the intended application. Please take into consideration

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Application Details	
	 which groups are available on the ligand for coupling, and try to prevent inactivation or hiding the active or exposed site of the ligand. After coupling the molecule of interest (ligand) is coupled to the magnetic particles, the resulting beads can be used in downstream applications such as: Isolating specific target proteins, antibodies, nucleic acids, cells, viruses, etc. (preparative applications) Detecting specific target proteins, nucleic acids, cells, viruses, etc. (diagnostic applications) Immobilizing enzymes, thereby enhancing stability and minimizing auto-catalysis. Magnetic collection of the particle/enzyme complex allows to remove the enzyme from the reaction, and to reuse it in a new reaction.
Protocol:	Magnetic beads are an ideal tool for immobilizing molecules (proteins, enzymes, antibodies, peptides, nucleic acids, etc.) on a solid phase, to be used for e.g. detecting, enriching, or cleaving specific target molecules. The easy and efficient collection of beads in magnetic fields allows for easy rinsing and removal of excess reagents and ligand after coupling the ligand molecule, as well as easy use in downstream applications. The use of magnetic beads does not require columns or centrifugation steps, and are therefore ideal in high-throughput and automated applications
	Our magnetic beads come in three sizes, 600 nm, 1 μ m and 3 μ m. 600 nm beads have the advantage of having a larger surface area and the sedimentation time of 600 nm MagSi beads is approximately 4 times slower than that of 1.0 μ m beads. This allows longer incubation times without shaking/mixing, and may be important in automated and other high-throughput applications in which shaking/mixing options are often lacking. MagSi beads with a diameter of 3 μ m have stronger magnetic properties and will separate approximately 4x faster than 600 nm beads under same conditions: approximate separation time is \geq 1 minute using a suitable magnet.
Restrictions:	For Research Use only
Handling	
Format:	Liquid
Concentration:	10 mg/mL
Buffer:	Stored in PBS, 0.05% sodium azide

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Preservative:	Sodium azide
Precaution of Use:	This product contains sodium azide: a POISONOUS AND HAZARDOUS SUBSTANCE which should be handled by trained staff only.
Handling Advice:	Store beads in well closed vial and in upright position to prevent drying of the beads since this makes them more difficult to re-suspend. Do not freeze the product! Vortex bead suspension well before use. If you expect iron interference in downstream applications, we strongly advise you to rinse the beads before usage. Before using the beads it is important to rinse with water or PBS to remove any components that could interfere with your test.
Storage:	4 °C
Expiry Date:	12 months

Images



nging nextine socialisme ease which reacts with amme groups concernes in proben. Gluberaldehyde glives more stable probein binding than the carbodienide reagents used with arbscylate beads.

3 Reduction of disulfides with 0.1 M DTE (dithioerythrei); coupling of protein at pM below isoelectric point; deactivate excess thiel with 20 mM PDEA (2-(2-pyridinyidithie) ethane-amine)/ 1M NOCI ref4.3 Image 1.

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Surface activation	Formula	Example Applications
Silica (stored in 0.05% sodium azide)	SI-OH	 End-users' own application (e.g. functionalization of the MagSi beads)
Carboxyl (stored in PBS, 0.05% sodium azide)	R-COOH	 Protein and peptide immobilization Antibody immobilization
Aldehyde (stored in PBS, 0.05% sodium azide)	R-CHO*	- Protein immobilization
Amine (stored in 0.05% sodium azide)	R-NH ₂	- Protein immobilization
Sulfydryl (stored in PBS, 0.05% sodium azide)	R-SH*	 Immobilization via target cysteine groups, coupling to gold surfaces
Tosyl (stored in DSM0:THF 1:1)	R-S O O CH ₃	 Antibody immobilization Protein and peptide immobilization
Hydrazide (stored in PBS, 0.05% sodium azide)	R-CO-N ₂ H ₂	- Glycoprotein immobilization - Protein and peptide immobilization
Epoxy (stored in DSM0:THF 1:1)	R-CH-CH ₂	 Enzyme immobilization Protein and peptide immobilization

coupling of other organic molecules, such as nucleic acids or carbohydrates, is also possible. CHO- and SH-beads have a limited stability, and must be used for coupling ligand within 2-3 weeks after production. Image 2. Active surfaces and example applications of

MagSi-tools