

PRODUCT INFORMATION SHEET

Version: 1 Revision Date: 08/08/2020

MagicLink[™] HRP Antibody Labeling Kits

Components

		Catalog Number	Storage
Components		BP-50061	
		1 x 5 mg	
А	MAGIC NHS (MW ~900)	1 vial	-20C
В	LINK activated HRP	1 vial	-20C
С	Reaction Buffer	30 ml	RT
D	Protein concentrator	1	RT

Overview

MagicLink[™] HRP Antibody Labeling Kits are the 3rd generation HRP conjugation technology which can be used to conjugate horseradish peroxidase (HRP) to protein, antibody, amine modified oligo, etc. The labeling kits feature the most stable linkage between HRP antibody on the market. The instant and efficient labeling reaction yield 95 to 100% HRP conjugates.

These kits are specifically optimized to label antibodies at 5 mg. The kit's format is based on instant reaction between functional group MAGIC and LINK at room temperature. By following the protocol provided in the kits, the end users can label their antibody with MAGJIC NHS to get MAGIC-antibody which instantly reacts with LINK activated HRP (provided in the kits) to achieve Ab-HRP conjugates.





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At a Glance

Protocol summary

- 1. Add reaction buffer into antibody.
- 2. Transfer whole antibody solution to MAGIC NHS vial.
- 3. Incubate at room temperature for 60 minutes.
- 4. Remove excess MAGIC by concentrator/filtration device.
- 5. Mix MAGIC-antibody with LINK activated HRP.
- 6. Storage.

Note: Upon receipt, store the kits at 4°C. When stored properly, the kits should be stable for six months. Alternatively, components A and B can be stored at-20°C. Do not freeze reaction buffer (component C). Warm all the components and centrifuge the vials briefly before opening, and immediately prepare the required solutions before starting your conjugation. The following SOP is an example for labeling goat anti-mouse IgG antibody.

Preparation of Working Solution

For labeling 5 mg of purified antibody (assuming the target antibody concentration is 5 mg/ml), final volume would be in 1 ml of reaction buffer (component C). For lyophilized antibody, add 500 μ l of reaction buffer to reconstitute. Antibody, in liquid form and not in 1X PBS, pH 7.2 – 7.5 may require buffer exchange, see note.

Note

- If you have a different concentration, adjust the antibody volume accordingly to make 5 mg antibody available for your labeling reaction.
- The antibody should be dissolved in 1X phosphate buffered saline (PBS), pH 7.5; if the antibody is dissolved in glycine buffer, it must be dialyzed against 1X PBS, pH 7.2-7.5, or use Amicon Ultra-0.5, Ultracel-10 Membrane, 10K MWCO (Cat # UFC501008 from Millipore) to remove free amines or ammonium salts (such as ammonium sulfate and ammonium acetate) that are widely used for antibody precipitation.
- Impure antibodies or antibodies stabilized with bovine serum albumin (BSA) or gelatin will not be labeled well.
- For optimal labeling efficiency the final antibody concentration range of 5 mg/ml is recommended.
- The presence of sodium azide will inhibit HRP activity.

Labeling Protocol

React Antibody with MAGIC NHS reaction:

Add the antibody solution directly into the vial of Magic NHS (component A), and mix them well by repeatedly pipetting for a few times or vortex the vial for a few seconds. Keep the antibody labeling reaction mixture at room temperature for 60 minutes. The antibody-labeling reaction mixture can be rotated or shaken for longer time if needed.

Purify the MAGIC-Antibody Solution

Note: Kit provides 2 concentrators for purification; each gets half the volume, (i.e., for 1 ml total MAGIC-antibody,





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each concentrator receives 500 µl).

- 1. Hydrate concentrator membrane 'filter devices' with 400 to 500 μl of reaction buffer or DI water, and microcentrifuge 14,000 x g, for 3 minutes. Discard, liquid from filter devices and collection tubes.
- Spin down by adding labeled antibody to the concentrators/filter devices up to 500 μl. Microcentrifuge at 14,000 x g, 8 - 10 minutes, or to minimum volume ~ 50 μl left in the filter devices. Discard waste from the collection tubes.
- 3. Desalt by adding reaction buffer to the filter devices up to 500 μ l. Microcentrifuge at 14,000 x g, 8 10 minutes, or to minimum volume ~ 50 μ l left in the filter devices. Discard waste from the collection tubes.
- 4. Repeat step 3, twice.
- 5. Collect labeled antibody from filter devices into a microcentrifuge tube.
- 6. Optional for maximum recovery, add reaction buffer, volume determined by the user, to the filter devices to rinse out residual antibody, microcentrifuge pulse spin, collect antibody/reaction buffer from filter device, add to the microcentrifuge tube from step 5, mix.
- 7. Determine sample concentration, then proceed to the next section.

HRP-Antibody Conjugation

- 1. Make LINK-HRP solution by adding 250 μl ddH₂O into the vial of LINK-HRP (component B), mix well by repeatedly pipetting for a few times or vortex the vial for a few seconds.
- 2. Mix vial of LINK-HRP solution into the purified MAGIC-antibody solution (from step purify the MAGIC-antibody solution, above), at different ratios, mix well and rotating the mixture for 1 hour at room temperature.

HRP to Ab ratios	Volume of LINK-HRP solution to add to ~5 mg MAGIC-Ab
5:1	250 μl
4:1	200 μl
3:1	150 μl
2:1	100 μl
1:1	50 μl

MAGIC-antibody to LINK-HRP labeling ratios:

Note:

- It is recommended the mix Link activated HRP and antibody at 3:1.
- Magic activated protein/antibody should be used right away.
- For a different protein, user need optimize the protein/HRP mix ratio for fit the application accordingly.

The HRP-antibody conjugate is now ready to use. For immediate use, the HRP-antibody conjugate need be diluted with the buffer of your choice. For longer term storage, HRP-antibody conjugate solution need be concentrated or freeze dried.

Storage of HRP-Antibody Conjugate

The antibody conjugate should be stored at > 0.5 mg/ml in the presence of a carrier antibody (e.g., 0.1% bovine serum albumin). For longer storage, the HRP-antibody conjugates could be lyophilized and stored at ≤ -20 °C.



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Troubleshooting

Problem	Possible cause	solution	
Low or no	Buffer containing primary	Buffer exchange the antibody into a non-amine-containing buffer such as the PBS provided	
conjugation with	amine	by desalting columns or dialysis	
MAGIC NHS	MAGIC NHS was hydrolyzed	Use reagent immediately upon reconstitution	
	Carrier protein was present the antibody solution	Remove carrier protein before conjugation by using Protein A, G or A/G resin or an antibody clean-up kit. This will reduce competition for labeling	

Address: 6625 Top Gun Street, Suite 103 San Diego, CA 92121 Phone: +1-858-677-6760; Fax: +1-858-677-6762

Website: https://www.broadpharm.com Email: sales@broadpharm.com