T-Pro Protein Free Blocking Buffer



In PBS (JK92-W003) 500 ml In TBS (JK92-W004) 500 ml



This product is for laboratory research ONLY and not for diagnostic use.

Description

The T-Pro Protein Free Blocking Buffers contain a proprietary compound for blocking excess binding sites in ELISA, Western blotting, arrays and other immunochemical applications. This blocking buffer reduces or eliminates many of the problems encountered with traditional protein-blocking reagents, such as cross-reactivity and interference from glycosylation. Additionally, T-Pro Protein Free Blocking Buffers are compatible with antibodies and avidin/biotin systems.

Important Product Information

- The usage as described in these instructions may differ from other blocking solutions.
- Use the protein-free blocking buffers at the supplied concentration; do not dilute blocking buffer.
- A final concentration of 0.05% Tween-20 Detergent in the blocking buffer often improves blocking; however, it is not required nor recommended for all systems.
- The protein-free blocking buffers may be used as a protein stabilizer for drying antigen- or antibody-coated microplates. Dry plate completely before sealing in a plastic bag with desiccant. Store plate at 2^8 °C.

Storage

T-Pro Protein Free Blocking Buffer is stable for 2~8°C

Instructions

Procedure for Blocking ELISA Plates

- 1 Coat the ELISA plate with antigen or antibody according to standard procedures.
- Add 300 μ L of the blocking buffer to each well and incubate for 30 minutes at room temperature. Alternatively, add 300 μ L of blocking buffer to each well and immediately invert plate to empty contents. Repeat this process two more times.
- Proceed with assay or invert plate, and allow it to completely dry for ~2 hours. Place dry plate in a plastic bag or other container with desiccant and store at 2~8°C.

Procedure for Blocking Membranes

Note: Typically, adding a final concentration of 0.05% Tween-20 to the blocking buffer produces the best results.

- 1 Add sufficient Protein-Free Blocking Buffer to cover the entire surface of the membrane.
- 2 Incubate for 15~30 minutes at room temperature on a rocking platform.
- Continue the blotting procedure do not using the Protein-Free Blocking Buffer to dilute primary and secondary antibodies.
- * Continue the blotting procedure do not using the T-Pro Protein Free Blocking Buffer to dilute primary and secondary antibodies.