

ELISA Protocol

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Introduction

ELISA (Enzyme-Linked Immunosorbent Assay) is a sensitive biochemical technique for detecting and quantifying proteins, antibodies, hormones, and other biomolecules in biological samples.

Various types of ELISA exist, including direct, indirect, sandwich, and competitive ELISA procedures. Sandwich ELISA is particularly beneficial for detecting low-abundance proteins in complex samples, as it employs two antibodies for improved specificity and sensitivity.

Reliable results depend heavily on protocol adherence, quality reagents, and optimized conditions. Boston BioProducts offers high-quality reagents specifically designed for consistent, accurate ELISA outcomes.

Browse Our Buffers

Materials & Reagents

Here's everything you need to carry out our ELISA protocol. In addition to Boston BioProducts' highquality buffers and reagents, you'll need the following materials and equipment:

- 96-well microplate
- Capture and detection antibodies
- Plate reader (spectrophotometer)
- Multichannel pipettes and tips
- Plate sealer films

Reagent	Product Name	CAT#
Coating Buffer	Borate Buffer (0.5M, pH 8.5)	BB-66
	<u>Carbonate-Bicarbonate Buffer (1M, pH 9.0)</u>	BB-64
Blocking Buffer	Casein (3%, PBS)	IBB-120
	Bovine Serum Albumin, BSA (5% TBST)	IBB-189
	<u>Gelatin (10%)</u>	IBB-184
	Casein (3%, TBS)	IBB-130
	Non-Fat Powdered Milk (Blotto)	P-1400
Wash Buffer	<u>PBST (10X, 0.5% Tween, pH 7.4)</u>	IBB-171
	<u>PBST (20X, 2% Tween, pH 7.4)</u>	IBB-170X
	<u>TBST (20X, 2% Tween, pH 7.4)</u>	IBB-180X
	<u>TBST (10X, 0.5% Tween, pH 7.6)</u>	IBB-181-6
	<u>PBS (10X, pH 7.4)</u>	BM-220
	<u>Citric Acid Buffer (2M)</u>	BPS-356
Antibody Dilution Buffer	<u>PBST (0.05% Tween-20, pH 7.4)</u>	IBB-171R
	<u>TBST (0.05% Tween-20, pH 7.4)</u>	IBB-181R
	Bovine Serum Albumin (BSA, 5% TBST)	IBB-189
	<u>1M Tris-HCl (pH 8.0)</u>	BBT-80

Reagent	Product Name	CAT#
Stop Solution	Sodium Hydroxide (1 N)	BZ-8041
	<u>Hydrochloric Acid (1N)</u>	BZ-8010
	Sodium Dodecyl Sulfate, SDS (10%)	BM-230A
	Sodium Dodecyl Sulfate, SDS (20%)	BM-230
	<u>EDTA (0.5 M, pH 8.0, RNase Free)</u>	BM-150-DR
	Acetic Acid (2 M)	BPS-211

Boston BioProducts' Protocol

This ELISA protocol outlines key steps, from plate coating to final signal detection, for a reproducible and optimized ELISA workflow. While this protocol primarily details the Sandwich ELISA workflow, it also provides insights into the other ELISA types to help you select the best assay for your research needs.



Sample Preparation

Your sample preparation may vary, but optimizing this step is a critical part of the ELISA protocol. It ensures antigen integrity and reduces background interference, leading to better results.

Our general advice by sample type:

- Serum or plasma: Centrifuge to remove debris and dilute in <u>blocking buffer</u> or sample diluent to fall within the assay's dynamic range.
- **Cell culture supernatant:** Clarify by centrifugation and store on ice or at-20°C until use.
- **Tissue or cell lysates:** Use a <u>lysis buffer</u> compatible with ELISA (avoid detergents or agents that interfere with antibody binding). Centrifuge lysates and use the supernatant.

Ý Avoid repeated freeze-thaw cycles, which can degrade protein targets.

Coating

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Coating the plate with the capture antibody enables specific immobilization of the target antigen, forming the foundation for sensitive and selective detection in Sandwich ELISA. If you're using a kit with pre-coated plates, you can proceed directly to the blocking step.



Add 100 μ l to each well, seal the plate, and incubate overnight at 4°C

To preserve coated antibodies during longer storage, consider adding a protein stabilizer such as <u>trehalose</u> or <u>sucrose</u>.

3) Remove excess solution and wash 3x with wash buffer.

Blocking

Coating the plate with the capture antibody enables specific immobilization of the target antigen, forming the foundation for sensitive and selective detection in Sandwich ELISA. If you're using a kit with pre-coated plates, you can proceed directly to the blocking step.

1) Add 200 μ l of a <u>blocking solution</u> to each well.

Other recommended blocking agents include Casein (3% in <u>PBS</u> or <u>TBS</u>), <u>Gelatin</u> (10%), and <u>milk powder</u>, depending on compatibility with your assay system.

- (2) Incubate for 1 hour at room temperature.
- 3) Aspirate and wash 3x with wash buffer.

Sample Addition

During this step, the antigen from your sample binds specifically to the immobilized capture antibody. Accurate sample dilution and incubation are essential to ensure optimal binding and reliable quantification within the assay's dynamic range.

Not doing a sandwich ELISA?

Here's how direct, indirect, and competitive ELISA procedures differ.

Mini direct ELISA protocol

- $\overbrace{1}$ Coat the plate directly with antigen.
- 2) Block with <u>BSA</u>, <u>Casein</u>, or <u>milk powder</u>.
- 3) Add enzyme-conjugated primary antibody.

- 4) Incubate, <u>wash</u>, and add TMB substrate.
- 5) Stop the reaction and read at 450 nm.
 - ▲ Direct ELISA has fewer steps but lower signal amplification and higher background risk, making it a simple and fast ELISA protocol ideal for detecting a strong, high-purity antigen.

Mini indirect ELISA protocol

- 1 Coat plate with known antigen.
- (2) Block with <u>Casein</u> or <u>BSA</u> in <u>TBST</u>.
- 3) Add a sample containing primary antibody.
- 4) Add enzyme-linked secondary antibody.
- 5) Stop the reaction and read at 450 nm.

The indirect ELISA protocol provides signal amplification but may introduce crossreactivity with the secondary antibody. It is useful for detecting antibodies in a sample, such as in serological testing.

Mini competitive ELISA protocol

- $\widehat{1}$ Coat plate with known antigen.
- 2) Incubate sample antigen with enzyme-conjugated antibody.
- 3 Add mixture to the coated well.
- (4) <u>Wash</u>, develop signal, stop reaction, and read.
 - ▲ Signal intensity is inversely proportional to sample concentration in competitive ELISA, therefore the competitive ELISA procedure is mostly used when the analyte is small or available in limited quantities.
- Prepare antigen standards and samples in a compatible diluent, such as <u>PBST</u> or <u>TBST</u>.
 - Add 100 μ l per well and incubate 1–2 hours at room temperature.
- 3) Wash thoroughly 3x with <u>wash buffer</u>.

Detection Antibody Incubation

The detection antibody binds to a second epitope on the captured antigen, forming the "sandwich" complex. When enzyme-conjugated, this antibody enables signal generation upon substrate addition, allowing quantification of your target analyte.



-) Add 100 μ l per well and incubate 1 hour at room temperature.
- 3) Remove excess solution and wash 3x with <u>wash buffer</u>.

Substrate Incubation

The substrate reacts with the enzyme linked to the detection antibody to produce a measurable signal—typically a color change. Therefore, the enzyme used dictates which kind of substrate to choose.



Add 100 μ l of substrate to each well—including blanks—and incubate the plate at room temperature, protected from light.

- ▲ If using horseradish peroxidase (HRP)-conjugated antibodies, add TMB substrate (preferably cold) and incubate for 15 minutes.
- If using alkaline phosphatase (ALP)-conjugated antibodies, use pNPP substrate and incubate for 30-60 minutes.

Stop Reaction

Stopping the enzymatic reaction preserves the developed signal and prevents further color change. The choice of stop solution depends on the enzyme used and the substrate system, with acidic or denaturing agents commonly employed to halt activity.

1) Add 50 μ l of stop solution to each well.

⚠ If using a TMB reaction, add Sulfuric Acid. The solution will change from blue to yellow.

For ALP-based assays or alternative substrates, other stop solutions such as <u>Sodium</u> <u>Hydroxide</u>, <u>Hydrochloric Acid</u>, <u>EDTA</u>, or <u>Acetic Acid</u> may be appropriate.

For specific applications, <u>SDS</u> may be used to denature the enzyme and stop the reaction without affecting pH-sensitive signals.

2 Immediately measure absorbance at the appropriate wavelength for your substrate using a plate reader (e.g., typically 450 nm for TMB, 405 nm for pNPP).

Incubation time should be optimized to achieve sufficient signal intensity without exceeding the linear range of detection.

Data Analysis

Once the plate has been read, ELISA results are analyzed by comparing sample absorbance values to a standard curve. Accurate curve fitting and replicate consistency are essential for quantifying analyte concentrations with confidence

- 1 Subtract the average blank absorbance (e.g., wells with buffer only) from all sample and standard values to correct for background.
- Plot the standard curve using known antigen concentrations (typically log concentration vs. absorbance).
- 3 Apply curve fitting—commonly linear regression or a 4-parameter logistic (4PL) model—depending on your assay's range and shape.
- (4) Use the curve to interpolate unknown sample concentrations from their absorbance values.
- 5) Check replicates for consistency and exclude any outliers that deviate significantly.
 - A For best results, ensure your sample values fall within the linear portion of the standard curve and re-run samples that fall outside this range.

Troubleshooting & Optimization Tips for ELISA Protocols

Even the most well-planned experiments can encounter unexpected challenges. Common issues such as poor protein transfer, weak signals, and high background noise can significantly impact data quality and reproducibility.

Using high-quality buffers and reagents, combined with a well-optimized protocol, can minimize the risk of these common challenges. Below are some common issues encountered during the SDS page and western blot protocol, alongside some key strategies to help you troubleshoot and optimize your results.

My background signal is too high

The problem:

High background can result from non-specific binding, insufficient washing, or suboptimal blocking.

What to do:

- Increase wash stringency (more washes, higher <u>% Tween-20</u>)
- Try a different blocking agent (e.g., <u>BSA</u> vs. <u>casein</u>)
- Ensure incubation conditions aren't too long or warm

My signals are weak or inconsistent

The problem:

Inadequate antigen capture or low antibody affinity may reduce signal intensity.

What to do:

- Confirm antibody concentrations and storage conditions
- Increase sample incubation time
- Ensure the antigen or sample wasn't degraded during prep

My replicates are highly variable

The problem:

Pipetting error, evaporation, or inconsistent plate handling can affect reproducibility.

What to do:

- Use a multichannel pipette
- Pre-wet tips
- Seal plates during incubation
- Avoid edge effects by not analyzing outer wells unless blocked and filled properly

Why Boston BioProducts?

Consistent buffer composition, ultrapure chemicals, and carefully formulated blocking agents can improve protein separation, enhance target detection, and reduce non-specific binding. Whether it's selecting the right lysis buffer for efficient protein extraction, using high-purity SDS sample buffers for proper denaturation, or employing optimized detection reagents for clear, high-sensitivity results, each component plays a crucial role in data quality.

By integrating Boston BioProducts' high-quality reagents into your SDS-PAGE and Western blotting protocol, you can achieve clearer, more consistent results, increasing the reliability and reproducibility of your protein analysis, and reducing troubleshooting time.



FAQs

What is the difference between Sandwich and Direct ELISA protocols?

The key difference lies in antibody usage and assay design. Sandwich ELISA uses two antibodies—a capture antibody immobilized on the plate and a detection antibody that binds to a different epitope on the same antigen. This format offers high specificity and sensitivity, making it ideal for detecting low-abundance targets in complex samples like serum or cell lysates.

In contrast, a Direct ELISA protocol uses a single enzyme-conjugated primary antibody that binds directly to the immobilized antigen. This approach is faster and simpler, with fewer steps and reagents, but it typically results in lower sensitivity and higher background, especially in unpurified or complex sample matrices.

Can I reuse ELISA plates?

No—ELISA plates are designed for single use and should not be reused. These plates typically have surface treatments that enhance protein binding, and once a plate has been coated or exposed to biological material, it becomes prone to contamination, degradation, or loss of binding efficiency. Reusing plates can lead to inaccurate readings, increased background noise, and unreliable data, compromising the integrity of your assay results.

What's the best way to choose a blocking buffer?

Choosing the right blocking buffer depends on your sample type, antibodies, and assay goals. Common options include <u>BSA</u>, <u>casein</u>, <u>gelatin</u>, and <u>non-fat dry milk</u>, each with different properties. For example, <u>BSA</u> is a widely compatible, low-background choice; <u>casein</u> offers strong blocking in complex matrices; <u>milk powder</u> is inexpensive but may contain biotin or phospho-proteins that interfere with some detection systems. <u>gelatin</u> is a good alternative when avoiding cross-reactivity with serum proteins, especially in anti-mouse or anti-rabbit systems—but it's not ideal for assays involving proteases, as it may be degraded. Testing different blockers during optimization is key to achieving reliable, low-background results.