

Double Antibody Sandwich ELISA (DAS-ELISA)

Our ELISA reagents are optimized using greiner bio-one microplates, medium binding. Before opening the tubes containing coating antibody (IgG) and IgG-AP- Conjugate please spin down all the liquid by a short centrifugation (approx. 3000rpm for a few seconds).



- 1. Dilute specific antibody in coating buffer (recommended dilution see delivery note and tube); i.e. 20µl in 20 ml buffer at a recommended dilution of 1:1000 or 40µl in 20 ml buffer at a recommended dilution of 1:500. Add 200µl to each well of the microtiter plate.
- 2. Cover the plates and incubate at 37 °C for 2- 4 h.
- 3. Wash plate with PBS-Tween using wash bottle, soak for a few minutes and repeat washing two times. Blot plates by tapping upside down on tissue paper.



- Extract samples 1:20 (w/v) in extraction buffer. Add 200 μl aliquots of the test sample to duplicate wells.
- 5. Cover the plates and incubate overnight at 4 °C.
- 6. Wash three times as in step 3.



- 7. Add 200 μ l enzyme conjugate, recommended dilution is given in the delivery note, in conjugate buffer.
- 8. Cover the plates and incubate at 37 °C for 2- 4 hours.
- 9. Wash three times as in step 3.



- 10. Add 200 μ l aliquots of freshly prepared substrate (1 mg /ml para- nitrophenyl- phosphate in substrate buffer) to each well.
- 11. Cover the plate and incubate at 37°C for 30-60 min, or as long as necessary to obtain clear reactions.
- 12. Assess results by:
 - a) Visual observation
 - b) Spectrophotometric measurement of absorbance at 405 nm

Reference

Clark, M. F. and Adams. A. N. 1977. Characteristics of the microplate method of enzyme-linked immunosorbent assay for the detection of plant viruses. *Journal of General Virology* 34: 475-483

QC-SOP-0087 Anl. 002 Double Antibody Sandwich ELISA (DAS-ELISA), Version 2.0

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Buffers used in ELISA

1. Coating buffer (pH 9.6)

1.59 g sodium carbonate (Na₂CO₃)
2.93 g sodium bicarbonate (NaHCO₃)
0.20 g sodium azide (NaN₃)
Dissolve in 900 ml H₂O, adjust pH to 9.6 with HCl and make up to 1 l.

2. PBS (pH 7.4) phosphate buffered saline

8.0 g sodium chloride (NaCl) 0.2 g monobasic potassium phosphate (KH_2PO_4) 1.15 g dibasic sodium phosphate (Na_2HPO_4) 0.2 g potassium chloride (Na_3) 0.2 g sodium azide (NaN_3) Dissolve in 900 ml N_2O_3 adjust pH to 7.4 with NaOH or HCl and make up to 1 l.

3. PBS-Tween (PBST)

PBS + 0.5 ml Tween 20 per liter

4. Sample extraction buffer (pH 7.4)

PBST + 2% PVP (e.g. Serva PVP-15 polyvinyl pyrrolidone)

5. Sample extraction buffer (pH 8.5) for Begomoviruses

0.05 M Tris containing 0.06 M sodium sulfite, pH 8.5

6. Conjugate buffer

PBST + 2% PVP + 0.2% egg albumin (e.g. Sigma A-5253)

7. Substrate buffer

97 ml diethanolamine 600 ml H_2O 0.2 g sodium azide (NaN₃) Adjust to pH 9.8 with HCl and make up to 1 liter with H_2O

Buffers can be stored at 4 ° C for at least 2 months. Warm to room temperature before use.

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ELISA Troubleshooting

1. No color development

- a) Did you omit any steps?
- b) Did you use the correct buffer for each step?
- c) Is your enzyme OK? Serum OK?
- d) Is your positive control homologous to antiserum (IgG)?

Recommendations - Do a titration plate. Use a reliable positive control in each plate. Pretest enzyme conjugate on substrate.

2. Nonspecific color development

- a) If in edge wells only:
 - Make sure the humidity in the incubator is sufficiently high.
- If this does not help, don't use edge or border wells, fill with buffer only.
- b) If in whole plate:
 - incomplete washing
 - old substrate
 - use recommended ELISA plate (greiner medium binding)
 - error in loading sequence

Recommendations - Use reliable negative control in each plate.

Use fresh substrate and check for spontaneous color change. Cover plates while incubating. Check pH of the buffers used.

- c) Some wells with inconsistent or unexpected reactions
 - incomplete washing
 - error in loading test antigens
 - spillage between wells

Recommendations - Use extra wash step, handle plates carefully with lids on, use predetermined loading pattern before loading. Blot top of plate after rinsing.

3. Color development very rapid; some color in healthy samples

- a) Enzyme conjugate concentration too high
- b) Substrate concentration too high

Recommendations - Use enzyme conjugate and substrate concentrations that will give OD_{405 nm} of about 1.0 in 30 to 60 min with good antigen source.

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