

Ecologiena
Nonionic Surfactant
APE ELISA KIT
(Microplate)
User's Guide

Measuring Principle (Competitive ELISA)

1. Competitive Reaction

The test is based on the recognition of APE by specific monoclonal antibodies. APE present in the sample and an APE-enzyme conjugate (i.e. APE labeled with a coloring enzyme) are premixed and added into each well of a microplate, and allowed to compete for limited number of binding sites of specific antibodies immobilized on the surface of the wells. When the APE concentration is higher relative to the enzyme conjugate, the APE will predominantly bind the antibody and vice versa.

2. Chromogenic Reaction

Unbound APE and excess antigen-enzyme conjugates are washed out. The chromogenic substrate is then added to develop color in conjunction with the enzyme conjugate. The amount of APE-enzyme conjugate remaining bound to the antibody will determine the color intensity. The higher APE concentration in sample, for example, leads to less antigen-enzyme conjugate bound to the antibody binding sites in a microplate well, generating a lighter color, i.e. lower absorbance.

3. Quantitative Analysis

The standard curve, a dose-response curve obtained from known concentrations of APE standards, is determined from the absorbance at 450nm. The APE concentration in each sample is accurately calculated by interpolation using the absorbance intensity obtained from the standard curve.

Kit Content

	Contents	Volume	Quantity	Storage
1	MoAb-Coated Microplate	96 Wells	1 Plate	2-8°C
2	APE Standard Concentrate (NP10EO, 4mg/L 20%MeOH)	4mL	1 Vial	2-8°C
3	Antigen-enzyme Conjugate	for 7mL	2 Vials	2-8°C
4	Buffer Solution- <i>white cap</i>	8mL	2 Vials	2-8°C
5	Uncoated Microplate	96 Wells	1 Plate	---
6	Wash Solution (6-fold concentration)	50mL	1 Vial	2-8°C
7	Chromogen Solution	250µL	1 Vial	2-8°C
8	Substrate Solution- <i>red marker</i>	15mL	1 Vial	2-8°C
9	Stop Solution- <i>black cap</i>	15mL	1 Vial	2-8°C
10	Instruction Booklet	---	1	---

Other Essential or Recommended Reagents, Materials

Essential

1. Disposable test tubes (e.g. IWAKI, item No. 9831-1207)
2. Glass fiber filters (e.g. ADVANTEC Co., item No. 36481047 ϕ 47mm) and filtering equipment
3. Micropipettes (20µL -100µL and 100µL -1000µL) and tips
4. Multichannel pipettes (50µL -300µL) and tips
5. Microplate reader (450nm wavelength) (e.g. TECAN, SPECTRA Classic)
6. Stop watch

3. Standard Solution

Dilute 4mg/L APE concentrate solution (20% methanol) with methanol and/or distilled water and prepare 10% methanol solution containing APE from 0.02mg/L to 1mg/L. Following is an example.

Standard solution (mg/L)	1.0	0.5	0.2	0.1	0.05	0.02	0
4mg/L APE concentrate (μL)	250	125	50	25	25	20	0
Methanol (μL)	50	75	90	95	195	396	100
Distilled water (μL)	700	800	860	880	1780	3584	900
Total (mL)	1.0	1.0	1.0	1.0	1.0	1.0	1.0

- Pour concentrate first into a disposable tube, then add methanol and finally distilled water for dilution to avoid non-specific adsorption onto the tube surface.
- Prepare the standard APE solution just before the test. Standard solution, once diluted from the concentrate, is NOT reusable at a later date. Prepare new standard solution for every test session.
- Be sure the standard concentrate is tightly capped after use and store it in a refrigerator. The standard solution must also be sealed or capped tightly to avoid methanol evaporation.
- Keep the methanol concentration of standard solutions at 10%. Higher methanol content in the sample may damage antibody and lower content may result in inaccurate readings.
- Dilute the concentrate in a single step to minimize adsorption onto the tube surface.
- Mix by filling the tip and expelling the contents with a pipette. Do not stir vigorously, with a Vortex mixer for example. Otherwise foaming may generate incorrect optical density.

4. Antigen-enzyme Conjugate Solution

Reconstitute antigen-enzyme conjugate powder with 7mL of buffer solution to prepare antigen-enzyme conjugate solution.

- Store the conjugate solution at 2-8°C; it will be stable for approximately 2 weeks.
- Mix by filling the tip and expelling the contents with a pipette.
- Mix a pair of reconstituted solutions when you use them altogether.

5. Mixture of Standard/Sample and Conjugate Solution

Transfer 100μL of APE standard (or sample) and 100μL of conjugate solution into each well of the uncoated microplate and mix by filling the tip and expelling the contents with a pipette.

- Dispense standard solution first, and then add conjugate solution to avoid non-specific adsorption on the inner surface of the well.
- Use 10% methanol solution as a blank.

6. Competitive Reaction

Dispense 100μL aliquots of the above mixture into each coated well of the microplate. Incubate the microplate for 60 minutes at room temperature (18-25°C).

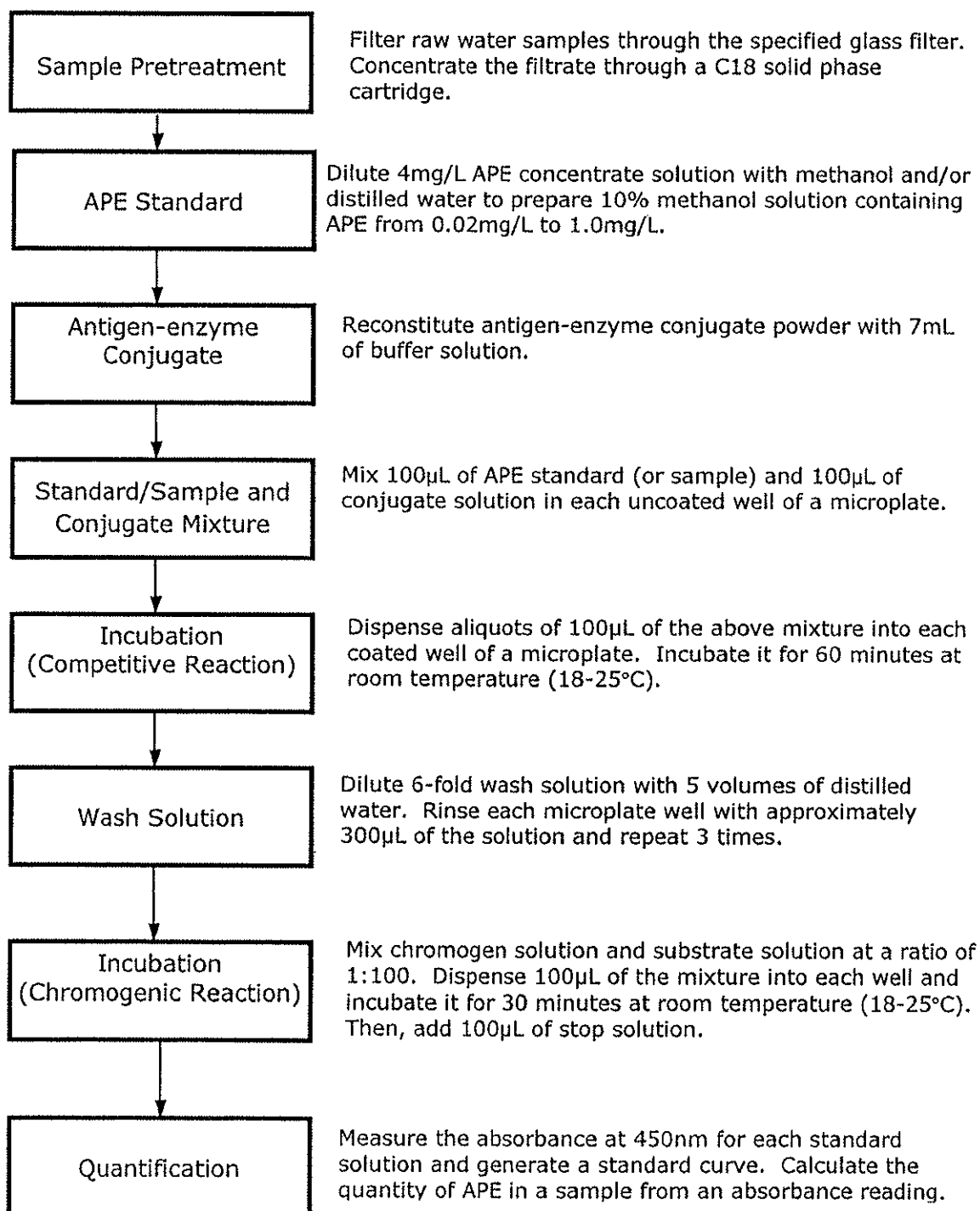
Split the microplate, with a strip ejector for example, to use necessary number of wells. This microplate is breakable into 12 strips, each of which consists of 8 wells. Place back the unused plate strips in the packet, seal and store them at 2-8°C.

APPENDIX

1. Flowchart for APE Measurement

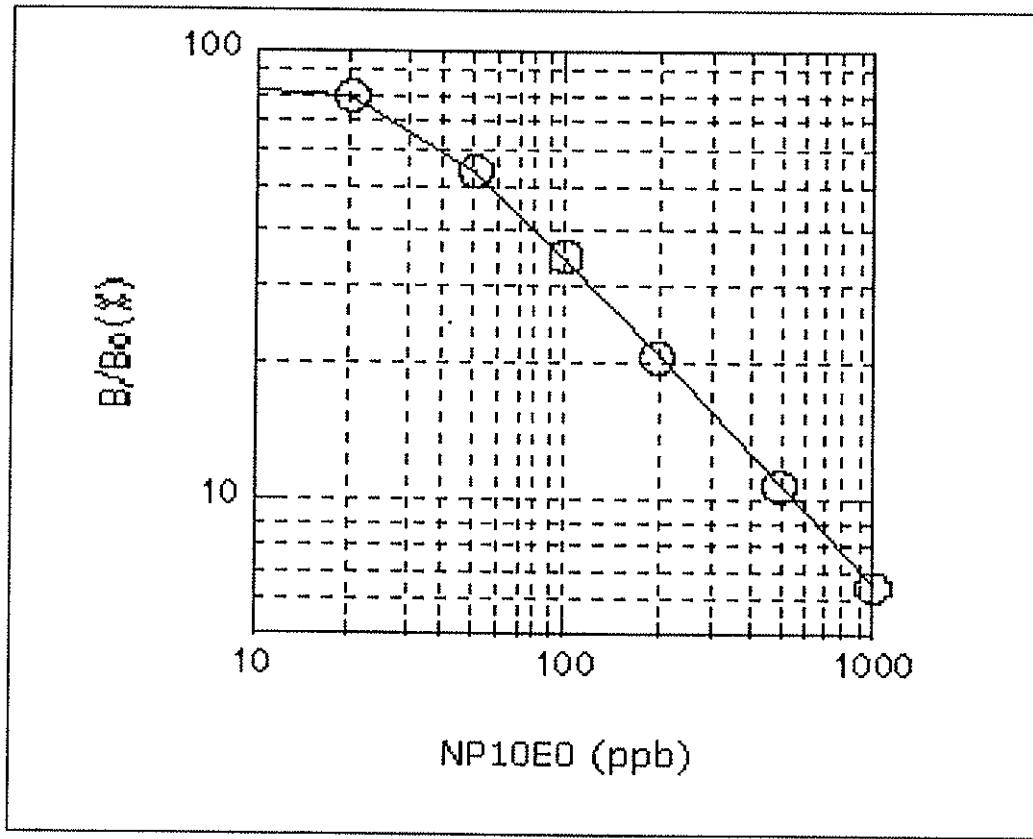
Note: Take out all the kit contents from a refrigerator and let them reach room temperature (18-25°C) prior to the assay.

<Please follow the steps describing in the text: Test Protocol>



4. APE Standard Curve

This test kit has a wide detection range between 0.02 mg/L and 1.0mg/L. Samples within this range can be directly applied to the assay only after filtration. Samples outside of the range must be either diluted with 10% methanol or extracted with solid phase concentration prior to analysis.



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