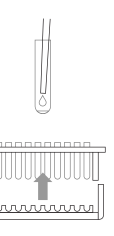


ABRAXIS® Atrazine Magnetic Particle 500001

1.



Remove upper rack from magnetic base. Label test tubes for Standards, Control, and Samples.

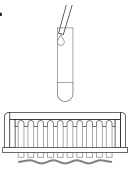
Tube # 1,2	Content Dilutent/Zero, 0 ppb
3.4	Standard 1, 0.1 ppb
5.6	Standard 2, 1.0 ppb
7.8	Standard 3, 5.0 ppb
9.10	Control
11,12	Sample 1
13,14	Sample 2
15,16	Sample 3

Add 200 or 250 µL of either Standards, Control or Samples to the bottom of each test tube by inserting the pipette tip all the way into the bottom of the tube without touching the sides of the tube.



<u>Do not</u> separate upper rack from lower base. Using a smooth motion, *invert* the combined rack assembly over a sink and pour out the tube contents; keep inverted and <u>gently blot</u> the test tube rims on several layers of paper toweling.

7.



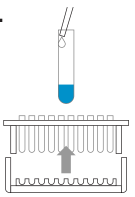
Add 1 mL of Washing Solution down the inside wall of each tube by using the technique described in Box 2. Wait 2 minutes. Using a smooth motion, invert the combined rack assembly over a sink and pour out the tube contents: keep inverted and gently blot the test tube rims on several layers of paper toweling. Repeat this step.

2.



Add 250 µL of Atrazine Enzyme Conjugate down the inside wall of each tube by using the technique described in Box 2. Vortex for 1 to 2 seconds (at low speed to minimize foaming).

8.



Lift the upper rack (with its tubes) off the magnetic base; add 500 μ L of Color Reagent down the inside wall of each tube by using the technique described in Box 2. Vortex for 1 to 2 seconds (at low speed to minimize foaming).

3.



Add 500 µL of thoroughly mixed Atrazine Antibody Coupled Magnetic Particles down the inside wall of each tube by using the technique described in Box 2. Vortex for 1 to 2 seconds (at low speed to minimize foaming).

9.



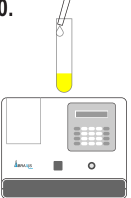
Incubate for 20 minutes at room temperature (15°-30° C). During this period, add 1 mL of Washing Solution into a clean tube for use as an instrument blank in Step 10.

4.



Incubate 15 minutes at room temperature (1 5 °- 30°C).

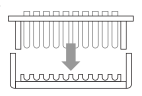
10.



Add 500 μL of Stopping Solution down the inside wall of each tube by using the technique previously described. Read results at 450 nm within 15 minutes after adding the Stopping Solution. Multiply results of samples by the appropriate dilution factor (if any).

[Safety Caution: Stopping Solution contains diluted sulfuric acid.]

5.



Combine the upper rack with the magnetic base; press all tubes into base; allow 2 minutes for the particles to separate.

Gold Standard Diagnostics 124 Railroad Drive Warminster, PA 18974 WEB: www.abraxiskits.com T (215) 357 3911 F (215) 357 5232

Ordering: info.abraxis@us.goldstandarddiagnostics.com Technical Support: support.abraxis@us.goldstandarddiagnostics.com

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