

# Glyphosate in Green Coffee Bean Sample Extraction

#### 1. Intended Use

For the detection of Glyphosate in green coffee beans.

### 2. Sensitivity

10 ppb in matrix

## 3. Materials and Reagents Required

Glass vials – 4 mL and 20 mL with Teflon-lined caps Conical Centrifuge Tubes – 15 mL or 50 mL Deionized water Serological pipettes, 5 mL or 10 mL Microcentrifuge tubes, 2.0 mL Microcentrifuge capable of 10,000 rpm, Analytical balance Weighing boats or equivalent Spatula Rotator and/or shaker Vortex mixer Hammer and zip-lock bags (4 MIL heavy duty) Coffee grinder, blender/food processor, or equivalent Sieve or mesh filter, 500 microns (PN 500103) or equivalent ABRAXIS® Glyphosate Sample Diluent (PN 500082) ABRAXIS® Glyphosate ELISA kit (PN 500205)

#### 4. Notes and Precautions

This procedure is intended for use with green coffee bean samples. Other matrices should be thoroughly validated before use with this procedure.

- It is highly recommended to add the appropriate amount of green coffee beans into a 4 MIL heavy-duty bag and homogenized/pulverized with hammer, and then finely ground with a coffee grinder, blender, or equivalent. The ground coffee beans should then be passed through a sieve or mesh filter of 500-micron pore size before extraction to produce accurate results.
- To minimize the potential for sample contamination due to carryover from the preparation of highly contaminated samples, the blender jar should be thoroughly washed and dried between samples.
- Due to the viscous nature of the sample extracts, if possible, the microtiter plate should be placed on a plate shaker or vortex mixer fitted with a micro-well plate holder adapter for the incubations with the antibody and conjugate solutions (Steps F.2 and F.3 in the ABRAXIS® Glyphosate ELISA user's guide). This will allow for the appropriate mixing of all reagents in the microtiter wells.
- Assay should be performed with ABRAXIS® ELISA kit as soon after extraction as possible. Samples should not sit more than one day in plastic microcentrifuge tubes before being run with the ELISA kit.
- This procedure is for research use only. It is not intended for diagnostic procedures.

#### 5. Extraction Procedure

- 5.1 Grind sample using a grinder, blender or equivalent.
- 5.2 Pass the ground sample through a sieve or mesh filter (500-micron pore size) and collect in a disposable weighing boat. If more sample size is needed, repeat steps 5.1 and 5.2.
- 5.3 Weigh 0.5 g of finely ground sample into a 15 or 50 mL conical centrifuge tube, or 20 mL glass vial.

- 5.4 Add 10 mL of deionized water to sample (20-fold dilution).
- 5.5 Vortex vigorously for 10 15 seconds. Place sample on rotator or shaker at 40 rpm for 10 minutes.
- 5.6 After mixing, let sample settle for at least 2 minutes. Transfer 1.5 mL of extracted sample to a clean, appropriately labeled 2.0 mL microcentrifuge tube.
- 5.7 Centrifuge tube at 8100 x g for 5 minutes. Make sure the centrifuge is properly balanced.
- 5.8 Add 850 μL of ABRAXIS® Glyphosate Sample Diluent to an appropriately labeled 4 mL glass vial. Add 150 μLof the supernatant (from 5.8) to the ABRAXIS® Glyphosate Diluent in the vial (1:6.67 dilution) and vortex or mixfor 15 seconds.
- 5.9 Derivatize the sample according to Section D Test Preparation in step 7 of *Derivatization of Standards, Control and Samples* instructions of the ABRAXIS® ELISA kit.
- 5.10 Perform assay as noted in Section F Assay Procedure instructions provided in the kit.

#### 6. Evaluation of Results

The ELISA results must be multiplied by a factor of 133.3 to account for the necessary dilution. Samples showing a concentration lower than Standard 1 (0.075 ppb) should be reported as < 10 ppb of Glyphosate. Highly contaminated samples (those outside of the calibration range of the assay) must be diluted and re-analyzed to obtain an accurate quantitative result.

# 7. For ordering or technical assistance contact:

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