



PathoSEEK®

Salmonella & STEC Multiplex Assay

with SenSATIVax® DNA Purification

QUICK GUIDE

DNA Purification:

Flower:

1. Weigh 1g sample into one side of the mesh lining of a Whirl-Pak bag then transfer 9 mL TSB into the bag. Hand homogenize for 1 minute.
2. Enrich at 37°C 16-24 hours.
3. Remove from the incubator and hand homogenize for 1 minute.
4. Transfer 1mL from the bag (non sample side of mesh lining) into a 1.5 mL tube.
5. Add 50 µL MGC Cell Lysis Buffer.
6. Vortex for 10 seconds. Let sit for 2 minutes then centrifuge at high speed for 5 minutes.
7. Transfer 200 µL supernatant to a well of extraction plate. Avoid pellet at bottom of tube.
8. Add 200 µL MGC DNA Binding Buffer and tip mix. Let sit at room temp 5 minutes.
9. Move the extraction plate to the magnet. Let the beads settle for 5 minutes.
10. Remove and discard supernatant (~400 µL) while avoiding beads.
11. Add 400 µL 70% Ethanol. Wait 30 seconds. Remove and repeat for a total of 2 washes.
12. Let the beads dry for 5-15 minutes.
13. Remove plate from magnet, add 50 µL MGC Elution buffer and mix to resuspend beads. Let sit for one minute.
14. Move the sample plate to the magnet.
15. Let beads settle for 1 minute, then transfer the eluent to a new well or plate.

Non-Flower:

1. Weigh 1g sample into 15 or 50 mL tube.
2. Add 2.4 mL TSB, vortex.
3. Enrich at 37°C for 16-24 hours.
4. Add 4.6 SenSATIVAx MIP Solution A.
5. Vortex to adequately homogenize.
6. Transfer 1 mL to 1.5 mL tube.
7. Add 10 µL of a Fresh 1:5k dilution of SCCG Internal Control.
8. Vortex and spin 10 minutes in high-speed centrifuge.
9. Transfer 600 µL to a new tube.
10. Add 600 µL chloroform.
11. Vortex well and centrifuge 5 minutes in high speed centrifuge.
12. Transfer 100 µL of supernatant to a well of extraction plate.
13. Add 100 µL SenSATIVAx MIP Solution B and tip mix.
14. Add 200 µL MGC DNA Binding Buffer and tip mix, let sit at room temp 5 minutes.
15. Move plate to magnet, let beads settle for 5 minutes.
16. Remove and discard supernatant (~400 µL) while avoiding beads.
17. Add 400 µL 70% Ethanol, wait 30 seconds, remove, repeat for a total of 2 washes.
18. Let the beads dry for 5-15 minutes.
19. Remove plate from magnet, add 50 µL MGC Elution buffer and mix to resuspend beads.
20. Let sit for 1 minute, move the plate to the magnet.
21. Let beads settle for 1 minute, transfer eluent to a new well.

qPCR Setup:

1. Prepare Master Mix and Positive Control Dilution

PCR Reagent Volumes

| Reagents | 1 Reaction |
|----------------------|---------------|
| qPCR Master Mix | 3.75 μ L |
| Assay Probe Mix | 1 μ L |
| Reaction Buffer | 0.8 μ L |
| Water | 8.2 μ L |
| Total Assay Probe MM | 13.75 μ L |

2. Prepare enough master mix for your samples plus two controls (positive and NTC). Add 10% overage to the master mix components to account for pipetting and dead volumes.
3. Dilute the stock assay positive control 1:10 with nuclease free water. 9 μ L water, 1 μ L positive control, vortex and spin down.
4. Transfer 5 μ L of each sample, 5 μ L of diluted assay positive control and 5 μ L of water to separate wells of a qPCR plate.
5. Transfer 13.75 μ L of freshly prepared qPCR Master Mix to each well and slowly tip mix.
6. Seal plate, spin in plate centrifuge and load on qPCR instrument.
7. Set up MGC qPCR cycling parameters:
 - a. 95°C, 5 minutes
 - b. 40 cycles of:
 - i. 95°C, 15 seconds
 - ii. 65°C, 90 seconds
8. Start run - Run takes approximately 1 hour and 40 minutes.

Data Analysis:

| PathoSEEK™ Assay | Cq Value | Fluor | Negative Control (Cq) | CFU threshold (CFU/g) |
|------------------------|----------|---------|---|-----------------------|
| <i>STEC</i> | ≤ 40 | FAM | No Cq | Presence/Absence |
| <i>Salmonella sps.</i> | ≤ 40 | ROX | No Cq | Presence/Absence |
| Internal Control* | ≤35 | HEX | *Internal control verifies the presence or absence of plant DNA obtained through the DNA Purification Process | |
| Assay Positive Control | ≤35 | FAM/ROX | | |

1. Confirm Assay Positive control well and assay NTC well to ensure the results are as expected.
 - a. Assay positive control should have a Cq value ≤ 35 for FAM and ROX.
 - i. No HEX signal should be observed in the control wells
 1. If HEX signal is observed a Cq of >35 is acceptable.
 - b. Assay NTC should have NO Cq value for FAM and ROX.
 - i. No HEX signal should be observed in the control wells
 1. If HEX signal is observed a Cq of >35 is acceptable.
 - c. Confirm Cq values against amplification plots

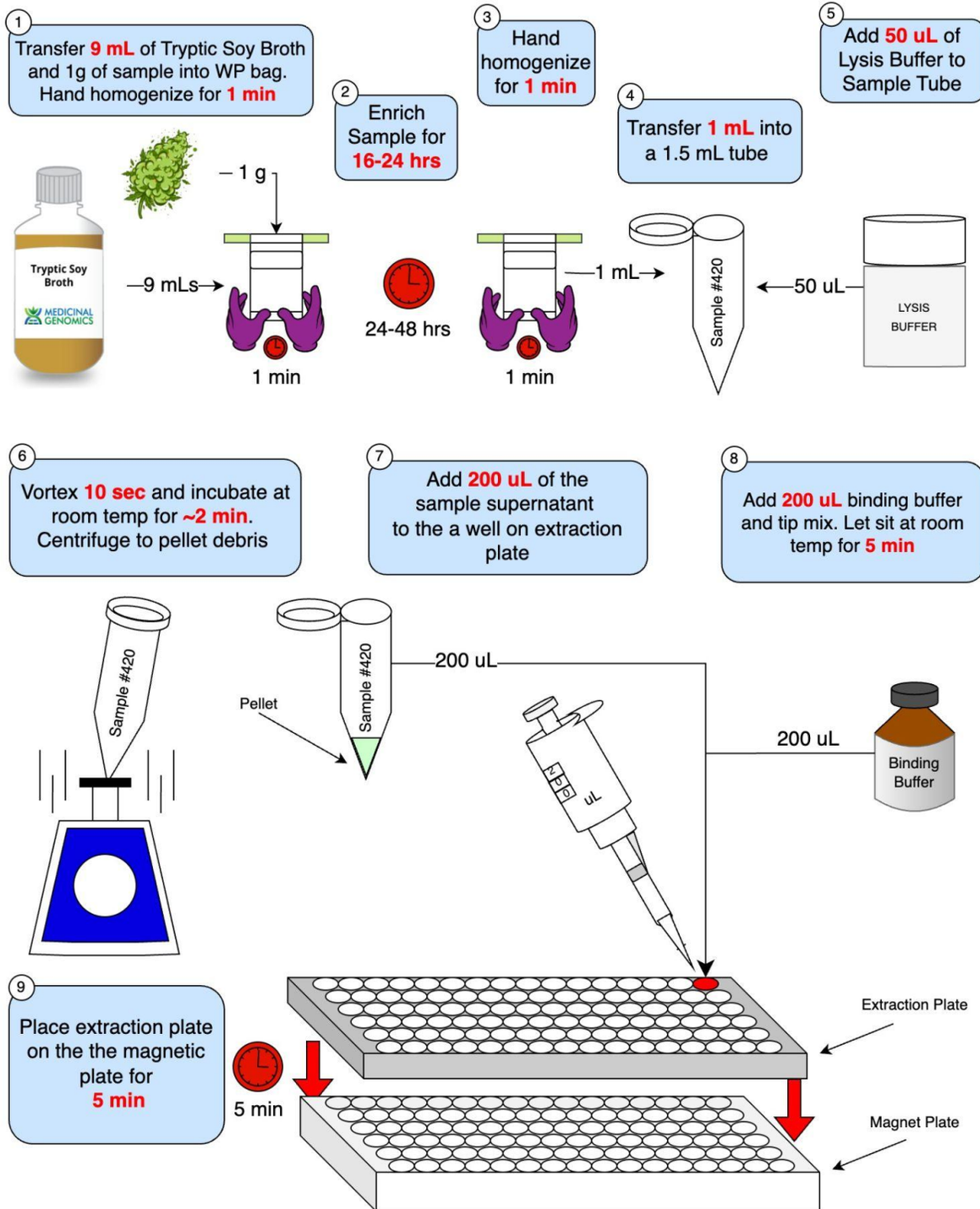
2. Unknown Samples
 - a. Internal DNA Purification Control (HEX)
 - i. HEX signals in sample wells should be ≤ 35 for flower ≤ 40 for non flower
 - b. Samples positive for STEC or Salmonella will show amplification which results in a Cq value ≤ 40
 - i. See table above to determine whether STEC or Salmonella is present
 - c. Confirm Cq values against amplification plots

For detailed User Guide please visit the Medicinal Genomics website

<https://medicinalgenomics.com/product-literature/>

Workflow Diagram: SenSATIVAx DNA Purification from Flower

SenSATIVax[®]
DNA Purification from FLOWER
 Aspergillus Assay



**STEPS 10-12
KEEP ON
MAGNETIC
PLATE**

10 Pipette to the bottom center of the plate well **WITHOUT** disturbing the ring of binding beads
Remove and discard all of the supernatant

Side view of well

Top view of well

Binding beads will form a **ring** at the bottom of the well

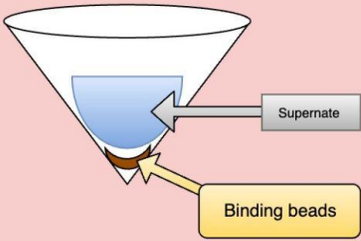
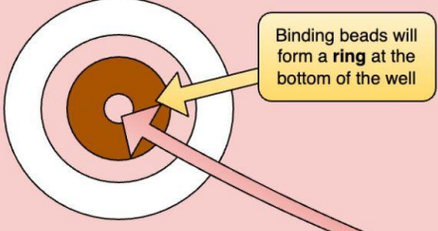
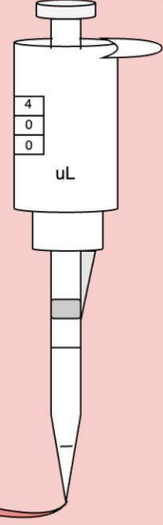
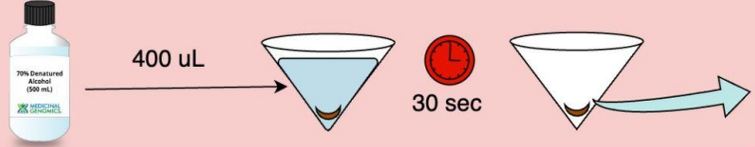
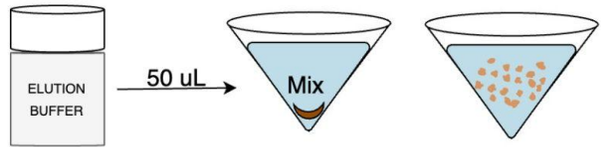
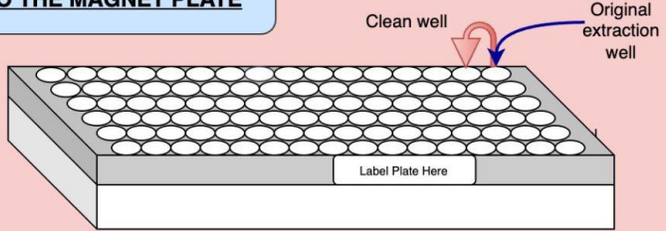
11 Pipette **400 uL** of 70% Ethanol into well (**DO NOT MIX**).
Let it sit for **30 seconds**, then remove all ethanol **WITHOUT** disturbing the ring of binding beads. **Repeat this step! (2 washes)**

12 Let binding beads dry for **5-15 minutes**
Make sure they do not over dry!

13 **REMOVE FROM MAGNET PLATE**
Add **50 uL** of Elution Buffer to each well and mix until the binding beads are fully resuspended.
Let sit for **1 minute**

14 **PLACE EXTRACTION PLATE ONTO THE MAGNET PLATE**

15 Let beads settle for **1 minute**, then transfer **50 uL** the eluent to a new clean well

**For detailed User Guide please visit the Medicinal Genomics website
<https://medicinalgenomics.com/product-literature/>**

DISCLAIMER

This test was developed, and its performance characteristics determined by Medicinal Genomics Company, for laboratory use. Any deviations from this protocol are not supported by MGC

The results may vary based on laboratory conditions. Altitude and humidity are among factors known to affect the growth of bacterial and fungal species. All thresholds were determined based on the results using the Agilent AriaMX or BIO-RAD CFX96 Touch® Real-Time PCR Detection System.

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