

# Salmonella Quantification (SalQuant™) with the BAX® System for Poultry Primary Production Boot Swab Samples

#### Introduction

Poultry are natural reservoirs for *Salmonella* and during the primary production stage, *Salmonella* can easily increase and spread throughout the flock and facility. Since *Salmonella* thrives in the GI-tract of poultry, managing the internal levels through vaccines, water and food treatments, and cleaning becomes a necessity to help reduce the load of *Salmonella* that enters the processing environment. Poultry houses are routinely sampled by utilizing a boot sock that can be worn on a work boot to physically walk the house to collect a sample that contains litter, liquids, solids, and fecal material. Hygiena's rapid and affordable method to quantify *Salmonella* provides a tool to make on-farm decisions and forewarn the processing facility of an incoming "hotter" flock. This helps the processing facility schedule and prepare to combat increased levels of *Salmonella* to provide safe and wholesome final products.

The objectives of these studies were to develop and verify rapid methods for PCR quantification of Salmonella (SalQuant<sup>TM</sup>) in poultry primary production boot swab samples.

# **Equipment, Supplies and Reagents**

- BAX® System Q7 instrument and supplies
- BAX® System Real-Time PCR Assay for Salmonella KIT2006
- Incubators For maintaining temperatures at 37°C and 42°C
- Brain heart infusion (BHI) broth
- BAX® System MP media MED2003/2016
- BAX® System BPW media MED2010/2011
- Antibiotic

# **Sample Preparation and Enrichment**

#### **Pure Culture Preparation:**

A culture of *Salmonella* Typhimurium strain ATCC 14028 was grown overnight in BPW broth at 37°C in preparation to inoculate boot socks. The culture was serially diluted in BPW broth to obtain a target concentration. Dilutions were plated in triplicate onto BHI agar and incubated at 37°C for 18-24 hours. The culture and dilutions were stored at 4°C until enumeration was complete.

### **Pre-screening of Matrix:**

Mock sampled boot swabs were procured from a commercial poultry producer for this study to represent normal background flora and matrix debris. Boot swabs were screened for naturally occurring *Salmonella* prior to inoculation by adding 100 mL of BPW to each sample, then homogenized by hand for 30 s to create a "slurry". One milliliter of the slurry was transferred into 9 mL BPW, then incubated at 37°C for 18 – 24 h and analyzed using the BAX® System Real-Time Assay for *Salmonella*. Only negative screened boot swabs were utilized to develop the quantitative method.

#### **Inoculation of Matrix:**

Following screening, negative boot swabs slurries were inoculated with an aliquot of the diluted

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Salmonella culture to create 3 biological replications of 5 inoculation levels (1, 10, 100, 1,000, 10,000 CFU/mL of solution). Inoculation unit of mL of solution was utilized to represent inoculation of 1 boot swab since prescreening was required.

#### **Enrichment Procedures for SalQuant™ Development (Figure 1):**

Following inoculation, boot swab slurry <u>primary enrichments</u> were homogenized by hand for 30 s. The <u>secondary enrichment</u> was created by transferring 30 mL of the primary enrichment to a new container and combined with 30 mL of pre-warmed (42°C) BAX MP media with antibiotic. The 60 mL solution was incubated at 42°C for 10 – 12 h. Samples aliquots were removed at 10, 11, and 12 hours for quantification and tested in quintuplet by the BAX® System method described below.

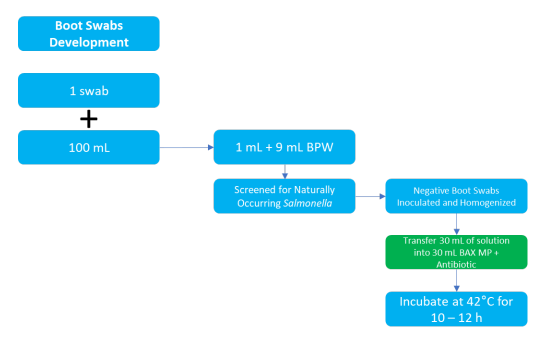


Figure 1. Enrichment procedures for SalQuant™ development.

#### **Verification Methods**

Presumptive negative boot swabs were collected from a commercial grower facility. Boot swabs were inoculated with an aliquot of the diluted *Salmonella* culture to create 3 biological replications of 5 inoculation levels (1, 10, 100, 1,000, 10,000 CFU/mL of slurry. Inoculated boot swabs were held at refrigeration for 48 h as a stress period. Following the stress period, 100 mL of BPW was added to the boot swabs and homogenized by hand for 30 s to create a primary enrichment slurry. Processing of slurries to secondary enrichment occurred 30 − 60 min following homogenization. The 30 mL aliquot of the slurry was transferred to a new container and combined with pre-warmed (42°C) BAX MP media with antibiotic to create the secondary enrichment and incubated at 42°C for 10 h. In parallel, 1 MPN was performed from the primary enrichment of each inoculation level as the reference method. After incubation, each sample was analyzed using the BAX® System Real-Time Assay for *Salmonella*, CT values collected, and estimations performed via SalQuant™ equations.

#### **PCR Method**

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BAX® System Method – For each sample, 5  $\mu$ L of enrichment was added to 200  $\mu$ L prepared lysis reagent (150  $\mu$ L of protease to one 12 mL bottle of lysis buffer) in cluster tubes. Lysis was performed by heating cluster tubes at 37°C for 20 minutes and 95°C for 10 minutes, and then cooling tubes at 4°C. Real-Time *Salmonella* PCR tubes were hydrated with 30  $\mu$ L of lysate and held for 10 minutes on a chilled (4°C) PCR cooling block. All PCR tubes were loaded into the BAX® System Q7 instrument and a full process was run according to the procedure described in the BAX® System User Guide.

#### Reference Method for SalQuant™

- Modified MLG MPN utilizing BAX prevalence testing for rapid confirmation of MPN results after incubation.

#### **Results**

#### **SalQuant™ Curve Development:**

- Boot Swab Development: 10 hours at 42°C produced a linear fit equation with an R<sup>2</sup> of 0.87 and Log RMSE of 0.43

## SalQuant™ MPN Verification (Figure 2):

One 3 x 5 Modified MLG MPN was performed at each level of inoculation to verify the efficacy of SalQuant<sup>M</sup> estimations. There was no statistical difference between MPN and SalQuant<sup>M</sup> estimations for boot swab.

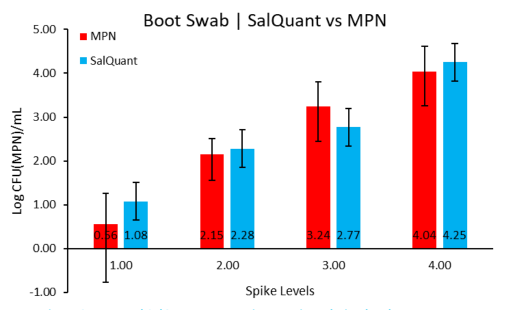


Figure 2. MPN and SalQuant™ comparison per inoculation level.

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## **Conclusions**

Overall, the results of this study demonstrate the ability of BAX® System Real-Time Salmonella to be used by enriching boot swabs to 10 h, complete quantification from 1 - 10,000 CFU/mL of slurry can be achieved simply and effectively. Using SalQuant<sup>TM</sup> approach depicted below (Figure 3) for boot swabs, poultry processors will be able to identify and manage flocks and farms that contain higher levels of Salmonella and take action to reduce exposure during processing and improve food safety.

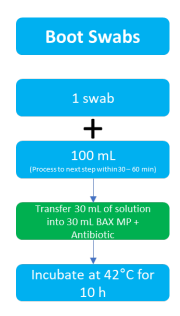


Figure 3. Enrichment protocol for Salmonella quantification (SalQuant™) with the BAX® System from poultry primary production boot swabs.

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