ANNEX C

Initial Contamination of *Salmonella* Enteritidis in Shell Eggs
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INTRODUCTION

This annex discusses the methods used to estimate the levels of *Salmonella* Enteritidis (SE) in contaminated eggs at lay. These estimates are based on a conceptual model of transovarian contamination (vertical transmission) of SE from hen to egg. SE infection of the hen reproductive system can lead to contamination of the internal contents of shell eggs with SE. It is generally believed that the initial levels of SE are low, though high initial levels of SE contamination can occur. SE contaminating the hen ovary and/or oviduct can be deposited at different locations within the egg, i.e. yolk and/or albumen. Depending where SE are deposited, their ability to grow, and therefore their likelihood of being detected will vary. This annex estimates the initial levels of SE in egg albumen and egg yolk.

To estimate the levels of SE in eggs at lay, studies enumerating the levels of SE in eggs from hens experimentally inoculated with SE were evaluated. A primary concern in using these data was the effect of the mechanism used to contaminate the egg (migration or direct deposition), the age of the egg at collection, the time-temperature history of the egg, and the location of contamination. These variables make it difficult to determine whether the enumerated levels of SE are due to initial contamination levels or due to subsequent SE growth. Assumptions concerning how much growth might have occurred if enumeration was not immediate are included. These assumptions define the model and enable estimates to be made of values for model parameters.

This annex is divided into two main sections: one that discusses distribution of the levels of SE in albumen (*Ea*)-contaminated eggs; and one that discusses distribution of levels of SE in the yolk (*Ey*) and vitelline membrane\(^a\) (*Ev*). These sections provide an estimation of the distribution of initial levels of SE in *Ea* and *Ey*-contaminated eggs, accompanied by the standard deviation and uncertainty surrounding the distribution. The distribution of SE in *Es* eggs and *Salmonella* spp. in *Ep* eggs are assumed the same as that in *Ea* eggs. This assumption was a state of knowledge necessitated by absence of data.

**DISTRIBUTION OF LEVELS OF SE IN *Ea*-CONTAMINATED EGGS AT LAY**

**Estimating the Distribution of the Number of SE Cells in Albumen**

Contamination of the albumen (*Ea*) is thought to be the primary site of SE contamination within eggs, and SE from *Ea* contaminations can migrate to other egg compartments. Therefore, determining the initial levels of SE within the albumen is important to determine risk. To estimate the initial levels of SE in egg albumen, we used raw data provided by Gast\(^1\) and Cogan.\(^2\) These data were from studies originally published by Gast and Beard\(^3\) and Cogan et al.,\(^4\) respectively.

The study of Gast and Beard\(^3\) enumerated SE from contaminated eggs from experimentally inoculated hens. These eggs were collected daily between 4 and 14 days post-inoculation. Three

\(^a\)Vitelline membrane is the surrounding yolk membrane. This membrane can be contaminated with SE without yolk contamination.
handling procedures were used for eggs produced by these treated hens: Group 1 contents of the eggs were analyzed for SE the same day eggs were collected; Group 2 eggs were stored at 7.2°C for 7 days before analysis; and Group 3 eggs were stored at 25°C for 7 days before analysis. Following these handling procedures, albumen samples were collected and frozen. Later, samples were thawed and analyzed for the presence of SE. The raw data from this study[1] are presented in Tables C1 and C2. Analysis of these data is presented in Table C3.

<table>
<thead>
<tr>
<th>TABLE C1 Group 1(^a) and 2(^b) combined raw data.(^1)</th>
<th>Measured SE cfu/ml</th>
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<tr>
<td>Frequency of Observations</td>
<td>2  3  4  6  7  8  9</td>
</tr>
<tr>
<td>(^a) Group 1 sampled on day of lay</td>
<td>2  1  2  1  1  1  1</td>
</tr>
<tr>
<td>(^b) Group 2 stored at 7.2°C and sampled seven days after lay</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TABLE C2 GROUP 3(^a) RAW DATA.(^1)</th>
<th>Measured SE cfu/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency of observations</td>
<td>1  2  3  4  5  7  8  9 12 35 38 85 106</td>
</tr>
<tr>
<td>(^a) Group 3 stored at 25°C and sampled 7 days after lay</td>
<td></td>
</tr>
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As the age of the eggs was unknown, the enumerated values for Groups 1 and 2 (Table C1) might represent initial contamination levels or subsequent SE growth from unknown levels. Therefore, to estimate the initial levels of SE from this study, relative growth that might have occurred during the time prior to analysis of the eggs was considered. From that, the possible amount of growth within the albumen was determined. The Group 3 data (Table C2) were not explicitly used here, as there was a strong possibility that SE growth occurred in these eggs due to holding at 25°C for 7 days. However, these data were used implicitly to determine the percentage of SE-positive eggs (see below; Table C3).

Assigning a Distribution to Initial Contamination in Albumen

For estimating the distribution of the number of SE cells, \(k\), in an egg at lay, it was assumed that the distribution could be written as:

\[
\begin{align*}
  f(0) &= p & \text{if } x = 0 \\
  f(k) &= (1-p)g(k|\psi) & \text{otherwise.}
\end{align*}
\]

where \(g(k)\) is the probability density function of \(k\) when \(k\) is positive, depending upon unknown parameters \(\psi\). It was assumed further that the number of SE cells in an egg at the time of measurement, \(y\), is the product of the initial number of SE cells in the egg, \(k\), and factor, \(r(k)\), describing the relative growth, so that \(E(y|r, k) = r(k)k\), and the distribution of \(y\) given \(r\) and \(k\) was assumed to be distributed as \(b(y|r, k)\). The distribution of \(m(y)\), for \(k > 0\), can be written as:

\[
m(y|\psi, r) = \int b(y|r, k)g(k|\psi) \, dk
\]

(C2)
where the integral represents the summation when the distribution is assumed discrete. Thus, the likelihood of an observation, \( z \), depends upon \( r \) and \( k \),

\[
\text{Lik}(z | \psi, r) = p \delta(z = 0) + (1 - p) \int\int m(y | \psi, r) h(z | y) \mathrm{d}y \mathrm{d}r
\]

(C3)

where \( \delta \) (true expression) = 1 and \( \delta \) (false expression) = 0, and where \( h(z | y) \) is the density of the Poisson distribution with an expected value = \( \lambda y \).

Gast and Beard\(^3\) froze egg samples prior to enumeration. The freeze-thaw process killed about 25% of the SE within the sample.\(^1\) Therefore, the recovery of SE was approximately 75%. To enumerate samples, 5 samples of 0.2 ml albumen were analyzed.\(^3\) Thus it is assumed 1 ml of albumen in total was analyzed. It was also reported that the average volume of the eggs was 40 ml. Thus, \( \lambda = 0.75/40 = 1/53.33 \).

Finally, it was assumed that \( r \) is a random variable with cumulative density function (cdf) \( V(r) \), which may depend upon \( x \) so that the unconditional likelihood of an observation \( z \), given \( x \) and unknown parameters is:

\[
\text{Lik}(z | \psi) = p \delta(z = 0) + (1 - p) \int\int m(y | \psi, r) h(z | y) \mathrm{d}y \mathrm{d}V(r).
\]

(C4)

The distribution \( V \) represents the variation of relative growth among SE-contaminated eggs.

**Estimating the Relative Growth of SE in Albumen**

To use the data of Gast and Beard\(^3\) to determine initial levels of SE in albumen, the level of relative SE growth needed to be determined to account for possible growth within a contaminated egg prior to collection. To do so, assumptions were made about the physiological state of SE cells and the age of eggs. Both factors are predicted to affect the relative growth of SE within albumen.

To determine the amount of growth of SE within albumen, data for SE growth in isolated egg albumen are available.\(^2\) Albumen was isolated, inoculated with 25 ml samples with varying levels of SE, and incubated for 20 and 30°C for 8 days. Levels of SE/mL were then enumerated and, for each egg sample, an exponential growth rate could be estimated from these data when assuming a fixed lag to generation time ratio, \( R_{at} \). However, as indicated by Cogan et al.,\(^4\) the SE cells used in this study were in stationary phase. SE within naturally contaminated eggs would not likely be in stationary phase. Though the amount of growth that would be expected for stationary phase cells is less than that for naturally contaminating SE cells, we are aware of no data to describe the growth phase of naturally contaminating SE, or to estimate the effect of this growth phase on SE growth in albumen. Therefore, it was assumed that the exponential growth rates would be the same for both initial phases of SE cells.
Eggs were collected in the morning for the study by Gast and Beard. Thus, it is possible that some of the collected eggs on a given day were up to 24 hours old. Alternatively, some of the collected eggs could have been laid shortly before they were collected. Therefore, it was assumed that the ages of the eggs in the study were uniformly distributed between 0 and 1 day old when sampled.

Through simulation of data from Cogan, selecting a random time between 0 and 1 for each sample, and computing the amount of SE growth for the sample assuming \( \lambda_t = 1 \), it was computed that \( E(r) = 1.5 \) and the standard deviation is 0.65 (so that \( E(r^2) = 2.7 \)). Thus, from this model, the amount of SE growth that samples with more than a few SE cells would experience is not large, generally less than a 3-fold increase (0.5 log10).

**SE Density and Growth in Albumen**

In a population of eggs inoculated with SE in the albumen, some of the eggs will experience SE growth; yet some will not. Given favorable time and temperature conditions, this could be due to location of the inoculum relative to the yolk, physiological egg variation, pH, and inoculum size. Concerning inoculum size, smaller inoculums of SE are less likely to grow within the albumen as compared to larger sizes. To determine the initial levels of SE deposited within the albumen, consideration for those percentages of eggs that did not show SE growth is needed.

Cogan in inoculated whole eggs in the albumen close to the yolk (Eac) with varying levels of SE and held them at 20 or 30°C for 8 days. For many inoculated eggs, SE growth was not observed, suggesting the distribution of relative SE growth among contaminated eggs, \( V \), of the function \( r \), be assumed such that \( V(1)(k) = v_0(k) > 0 \). The likelihood of \( z \) can then be written as:

\[
E(k, z | \varphi) = p \delta(z = 0) + (1 - p) \int_{k=0}^{\infty} \int_{r=0}^{\infty} h(z | k) v_0(k) d\varphi(k | z) + \int_{r=0}^{\infty} \int_{k=0}^{\infty} h(z | kr^2)(1 - v_0(k)) d\varphi(k | z) d\varphi(r) \]  
(C5)

Equation C7 expresses that the likelihood of SE growth depends upon the number of SE cells, \( k \), at time \( t \) in the initial contamination. For inoculation levels of 2, 25, 250, and 2,500 SE cells per egg, the percentages of samples for which the relative growth was less than 1.25 were 41, 19, 12, and 12%, respectively, nearly independent of the levels \( k \), for sufficiently large \( k \). Though there is a trend toward growth dependence and concentration, this result was unexpected based on the above discussion. However, these percentages include eggs in which SE appear to have had access to yolk nutrients due to yolk membrane breakdown (YMB). To identify the percentages of samples for which the relative growth was less than 1.25 in the albumen only, eggs for which there were indications of YMB were excluded. Upon so doing, the percentages of samples for which relative growth were less than 1.25 increased to 50, 25, 17 and 24% for inoculation levels of 2, 25, 250, and 2,500 SE cells per egg, respectively. Once again, for the samples with inoculation of 2 SE cells per egg, the percentage of eggs showing no growth was large, as expected; however, for sufficiently large \( k \), the levels appeared independent of inoculum
size. Further, for samples with a target inoculum of 2 cells per egg, a number of eggs did not contain SE. Therefore, errors in the number of SE cells that were inoculated could be large.

Most of the derived distributions of the number of SE cells for a target inoculum of 2 and 25 cells per egg is below 15 cells, so that it could be reasonably assumed that 20% or so of the eggs would not experience SE growth. This number, as a rough approximation, was assumed for all albumen-contaminated (Ea) eggs, so that, for the estimation of the distribution g, it was assumed that \( \nu_0 \) and \( V \) are not dependent on \( k \). With this simplification, from the above equations, the first and second non-central moments, respectively, of \( z \) are:

\[
E(z) = (1 - p)(\bar{E}(k))(\nu_0 + (1 - \nu_0)E(r))
\]

and:

\[
E(z^2) = E(z) + (1 - p)\nu_0^2E(k^2) + (1 - \nu_0)E(k^2)E(r^2)
\]

To complete this analysis, \( 1-p \), the percentage of eggs contaminated with SE in the albumen, was calculated from data of Gast and Beard.\(^3\) Hens were experimentally inoculated with SE and collected daily between 4 and 14 days post-inoculation. Three handling procedures were used for eggs produced by these treated hens: Group 1 contents of the eggs were analyzed for SE the same day eggs were collected; Group 2 eggs were stored at 7.2°C for 7 days before analyzing; Group 3 eggs were stored at 25°C for 7 days before analyzing. Following these handling procedures, albumen samples were taken from eggs and frozen. Upon enumeration, samples were thawed and analyzed for the presence of SE (Table C3).

### Table C3 Percent SE-positive samples and mean SE levels (CFU/ML) for three groups of eggs.

<table>
<thead>
<tr>
<th>Number of Positive Samples (%)</th>
<th>Group 1(^a) 4/132 (3)</th>
<th>Group 2(^b) 5/134 (3.7)</th>
<th>Group 1 and 2 9/266 (3.4)</th>
<th>Group 3(^c) 22/138 (15.9)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean SE CFU/ml</td>
<td>5.5</td>
<td>4.6</td>
<td>5.0</td>
<td>15.6</td>
</tr>
<tr>
<td>Geometric Mean</td>
<td>5.1</td>
<td>3.9</td>
<td>4.4</td>
<td>5.6</td>
</tr>
<tr>
<td>Median Value</td>
<td>5.5</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>25(^{th}) Percentile(^d)</td>
<td>3.25</td>
<td>2</td>
<td>2.5</td>
<td>2</td>
</tr>
<tr>
<td>High Value</td>
<td>8</td>
<td>9</td>
<td>9</td>
<td>106</td>
</tr>
</tbody>
</table>

\(^a\)Group 1 sampled on day of lay.
\(^b\)Group 2 stored at 7.2°C and sampled 7 days after lay.
\(^c\)Group 3 stored at 25°C and sampled 7 days after lay.
\(^d\)Computed as \( y = (1-f)x + f x_{i-1} \), where \( x_i \) is the \( i \)th ordered result, and the number of observations can be expressed as \( 4(i+f) - 1 \).

Table C3 shows a striking difference in the percentages of positive samples between those of the first two groups and that of the third group, i.e. 3.4% vs. 15.9%, respectively. A comparison between the first two groups and group 3 suggests that for SE deposited in many of the eggs from groups 1 and 2, initial levels of SE were below the detection limit of the methodology used in the study. In addition, a comparison between group 2 stored at 7.2°C and group 3 stored at
25°C suggests SE deposited in group 3 grew to levels above the detection limit and were therefore recovered more frequently than SE in group 2 eggs, which were held below optimal growth temperatures. For group 3, approximately 16% (22/138) of the egg samples were positive for SE. The actual percentage of SE-positive eggs in this experiment, however, could be greater than reported due to false-negatives. From the data of Cogan,² 22% of the egg samples (after an 8 day storage at 20 or 30°C), for which it was determined that there was no YMB event, showed less than a 25% increase in the number of SE cells (excluding the results from the lowest inoculation group of 2 cells). Because eggs from the study by Cogan² were assumed to represent Eac eggs, it is possible that there would be a larger percentage of no growth for Eaf eggs. For example, if 22% of the Eac eggs and 30% of the Eaf eggs did not experience growth, and 70% of the Ea eggs are Eaf, then the percentage of no growth eggs would be about 28%. Hence for group 3 it is possible that 22% (16%/(1-0.28)) of the eggs were positive for SE. That is, it is possible, that 20% or more of the eggs were positive in groups 1 and 2, of which, only 3-4% of them were detected positive, for approximately an 80 to 85% false-negative rate.

Equations C8 and C9 (described below) were examined for groups 1 and 2 data, assuming that 20% of the eggs were contaminated, so that \( p = 0.80 \), and that 20% of the samples that were positive for SE would not experience growth, so that \( v_0 = 0.2 \). In actuality, eggs with small number of SE cells or exceptionally young eggs may not experience growth of SE, and for those eggs that did experience growth of SE, the expected value would be larger (based on the condition of there being growth). However, to simplify the calculations, this adjustment was ignored, with the expectation that doing so would not cause a large error. With these assumptions, Equations C8 and C9 were used to solve for \( E(k) = 32.22 \) and the standard deviation of \( k \), \( \text{std}(k) = 65.15 \). The parameters for this distribution are given in the following section.

**Distribution of SE Cells Initially Contaminating Albumen**

To describe variation of SE levels within albumen, a lognormal distribution was applied to the data as described above. Historically, a negative binomial distribution has been used to describe microbial count and density data. For this analysis, however, a lognormal distribution was used, as it was necessary to assure positive contamination levels. These issues are discussed below.

To approximate \( g \), a lognormal distribution, it was assumed that \( \ln(k-1) \) is normally distributed with parameters \( (\omega, \varsigma) \), where \( \omega \) is the mean, and \( \varsigma \) is the standard deviation. This distribution assures values of \( k \) greater than 1. An attempt was made to fit a negative binomial distribution without zeros. While values of parameters were derived which resulted in a skewed distribution, in the simulations to determine the confidence intervals for the values of the parameters, the estimated values were negative approximately 45% of the time. This indicated unstable estimates and thus a lognormal distribution was assumed. This distribution depends upon estimates of the mean and standard deviation that are usually stable, even with small numbers of samples. From the values of \( E(k) \) and \( \text{std}(k) \) given above, the estimates of the parameters are \( \omega = 2.60221 \) and \( \varsigma = 1.29535 \). For this distribution, the 95\(^{th} \) percentile was 115 cells and the 99\(^{th} \) percentile was 276 cells. It should be noted also, that different values of \( v_0 \) had a small effect on the estimated values: for \( v_0 = 0.5 \), \( \omega = 2.71 \) and \( \varsigma = 1.30 \), and the 99\(^{th} \) percentile of \( k \) was estimated to be 311 cells.
Assigning Uncertainty for Initial SE Contamination in Albumen

Uncertainty of parameter estimates $\omega$ and $\zeta$ were determined by a bootstrap with 5,030 simulations. First, independent random variables $n$, from a binomial distribution with parameters 9 and 9/266 (Group 1 and 2) and $m$, from a binomial distribution with parameters 22 and 22/138 (Group 3), were generated. Then $n$ random selections from the 9 results, with replacement, were made. Using the $n$ results, estimates of $E(k)$, var$(k)$, $\omega$, and $\zeta$ were derived, as described above. The results for $n$ less than 3 were deleted, because, if 2 positive results were seen, it is probable that another procedure for estimating the distribution would have been developed.

The distribution of the estimates of $\zeta$ was negatively skewed due to a small percentage (0.2%) of exceptionally low results. When eliminating these results (which were 3.5 standard deviation units below the mean of all the results), the distribution was nearly symmetrical (skewness $=-0.14$, from -0.27 with all the results) and kurtosis of 2.65. The distribution of the simulated estimates of $\omega$ was negatively skewed with 5 negative results. Eliminating these five results, leaving 4,998 remaining bootstrap values, the distribution of $\omega$ and $\zeta$ were estimated: the distribution for $\zeta$ was nearly symmetric with skewness of -0.16 and a kurtosis of 2.64; and the distribution for $\omega$ was negatively skewed with skewness of -0.62 and a kurtosis of 4.5.

To derive a symmetric distribution to use for $\omega$, the quantities, $\phi(c) = (5-\omega)^{c}$ were considered for various values of $c$. When $c = 0.6$, the skewness of $\phi(0.6)$ was 0.03 and the kurtosis was 2.19. Thus, for determining the distribution of $\omega$, the distribution of $\phi(0.6)$ was used. The mean of $\phi(0.6)$ is 1.70, which transforms to a value of $\omega = 2.58$, which is close to the estimated value of $\omega$ of 2.60 given above. The standard deviation of the simulated values of $\phi(0.6)$ is 0.29390; the standard deviation of the simulated values of $\zeta$ is 0.1425; and the correlation between these two variables is 0.94310. To determine percentiles for these variables that account for the non-zero kurtosis, an Edgeworth expansion term, $(z^{3}-3z)\kappa_{4}/24$, where $z$ is a quantile of the standard normal distribution and $\kappa_{4}$ is the kurtosis, was used.

Results and Assumptions for Estimating Initial SE Contamination of Albumen

The results of this section estimate the distribution for the number of SE cells within the albumen ($Ea$) of a contaminated egg at the time of lay. This distribution was dependent on the parameter estimates of the mean and the standard deviation. Further, uncertainty for these estimates was determined. These estimates and the uncertainty are used in the exposure assessment and risk characterization to determine the risk to the consumer of consuming an SE-contaminated egg. The assumptions are: 1) at lay, the level, $k$, of SE cells in $Ea$-contaminated eggs is distributed such that $\ln(k-1)$ is a normal distribution with mean, $\omega$, equal to 2.6022 and standard deviation, $\zeta$, equal to 1.2953; and 2) the uncertainties of $\omega$ and $\zeta$ are determined by assuming $\phi = (5-\omega)^{0.6}$ and $\zeta$ to be nearly normally distributed with means of 1.6997 and 1.2953, and standard deviations of 0.2939 and 0.1425, respectively, with correlation coefficient of 0.9431. Generated standardized values from a bivariate normal distribution with zero means, unit standard deviations, and correlation of 0.9431, say $z_{1}$ and $z_{2}$, respectively, are adjusted by computing

$$z_{j} = \phi_{j} + \frac{\kappa_{4}(z_{j}^{2} - 2z_{j})}{24}, \quad j = 1, 2$$

(C8)
where $\kappa_4$ is the kurtosis. For $\phi$, $\kappa_4 = 2.19$ and for $\zeta$, $\kappa_4 = 2.64$. These adjusted values, $z_j$, are multiplied by the corresponding standard deviation (0.2939 for $\phi$ and 0.1425 for $\zeta$) and then added to the corresponding mean values (1.6997 for $\phi$ and 1.2953 for $\zeta$) to calculate the simulated values of $\phi$ and $\zeta$. The simulated value of $\omega$ is $5^{-\phi^{1/0.6}}$.

**DISTRIBUTION OF LEVELS OF SE IN YOLK (EY) AND VITELLINE MEMBRANE (EV)-CONTAMINATED EGGS AT TIME OF LAY**

The model used to describe the initial levels of SE in the yolk or on the vitelline membrane was guided by a conceptual model describing contamination of an egg by an infected hen. SE are capable of contaminating the ovary and oviduct of a hen. Infection of these sites can lead to vertical contamination of the inner yolk contents (Ey) or the vitelline membrane (Ev). Ey and Ev events can occur either directly, as indicated above, or by migration of SE from other locations within an SE-contaminated egg. Depending on the mechanism, some level of SE can be deposited within the yolk or the vitelline membrane. This section provides an estimate of the level of SE for Ey and Ev events at the time of lay. In arriving at these estimates, we used data from experimentally inoculated hens. Issues such as separating the contamination of the vitelline membrane vs. internal yolk contents, SE levels below enumeration, and false-negatives are considered in interpreting these data. These issues are discussed below.

**Estimating the Distribution of the Number of SE Cells in Yolk and on the Vitelline Membrane**

**Conceptual model**

Direct contamination of the yolk or vitelline membrane could occur early in the formation of the egg by contamination of the ovary and/or contamination of the opening of the oviduct, the infundibulum. This could be a large or small contamination event. Given the time the egg is within the hen, growth of SE could occur. Therefore, enumeration of SE within egg albumen at the time of lay does not necessarily indicate the initial contamination level. Indirect contamination of the yolk or vitelline membrane could occur by SE migration from the vitelline membrane or the albumen, respectively. As SE are motile and can move within the egg, after some time SE contaminating the albumen could reside on the vitelline membrane. Further, SE contaminating the vitelline membrane could contaminate the internal yolk contents. Contamination of these compartments would take time relative to the above route of contamination and therefore could demonstrate a lower likelihood of growth. These mechanisms suggest SE contamination levels of the yolk or vitelline membrane observed experimentally might demonstrate a multi-modal distribution due to the two routes of contamination and their potential for growth.
Modeling the Distribution of the Number of SE Cells in Yolk and on the Vitelline Membrane

The study used for determining the initial level of SE contamination in yolk was that of Gast and Holt. Hens were inoculated with SE and eggs collected daily for analysis. Yolk was separated from albumen and the vitelline membrane and yolk contents homogenized. To screen for SE-positive yolks, 10 ml of yolk/egg from among 874 eggs were enriched in broth. This screen identified 21 yolk SE-positive eggs. From here, a 1 ml aliquot from a 3 ml refrigerated companion sample was analyzed for numbers of SE. Among the 21 SE-positive yolk samples, 18 were reported non-enumerable; that is, the initial SE level within these samples was below the limit of detection. Of the remaining 3 samples that could be enumerated, the levels of SE were 4, 27 and 67 cfu/ml. If the volume of the yolk and vitelline membrane was assumed approximately 15 ml, and it was assumed that the distribution of SE cells within the yolk region was uniform, then it could be estimated that there were 1,005 SE cells within the yolk of the 67 cfu/mL egg. This assumption was reasonable because, before the samples were formed, the yolk material was homogenized.

The eggs were collected daily, so that the age of the eggs when the samples were prepared and the contents analyzed are not specifically known. Given that eggs may be contaminated through a contamination of the albumen with subsequent migration to the yolk, the age of the egg is likely an important factor in determining the transition from an Ev to an Ey event. Results presented by Gast and Holt suggest that migration can occur in a relatively short time (10% of Ev contaminated eggs can become Ey in 6 hrs), and thus it was presumed that the event could happen while the egg was still being formed in the oviduct or prior to collection. Once in the yolk, rapid growth of SE could commence following a lag period, upon which the contamination would spread throughout the yolk. Consequently, the distribution of the number of SE cells in the yolk among eggs not more than 1 day old could be highly heterogeneous, possibly, or most likely, multi-modal in nature, due to the different avenues of contamination and growth potential.

This expectation seems to be borne out by the results of Gast and Holt. The fact that 86% (18/21) of the known SE-positive samples could not be enumerated and that 14% (3/21) had high results, suggests a highly skewed, or multi-modal, distribution of SE cell counts within the contaminated yolks of eggs. For the purposes of analysis and modeling, it can be imagined that the yolk is divided into three regions: 1) the 10-mL volume (= v₁) initial screening sample that was analyzed to determine the presence of SE cells; 2) the 1-mL volume (= v₂) comprising 5 0.2 ml portions taken from a 3 ml sample for enumeration; and 3) the remaining volume (= v₃).

Let v be the total volume, assumed to be 15 ml, and pⱼ = the probability that a specified cell would be in the subsample of volume vⱼ, j = 1, 2, 3. Thus, p₁ = 2/3, p₂ = 1/15, and p₃ = 4/15. To compute the probability of SE within each of these regions, let w be the proportion of eggs that are not contaminated, and, for the other eggs, let the cumulative distribution of the number of cells in the egg, x, be F(x|θ), where θ are parameters. For a given x > 0, the probability of the number of cells in the sub-sample, vⱼ, xⱼ, j = 1, 2, 3, such that the sum, x, is a multinomial distribution. Furthermore, it was assumed that there was not 100% recovery, but the probability of a specific cell not being recovered is τ. This can be visualized as further dividing the volumes v₁ and v₂ into four subsections: v₁₁, v₁₂, v₂₁, v₂₂, such that v₂ = τv₁ represents the volume for which the cells are not recovered. Thus, the actual number of SE cells recovered is x₁₁ and x₂₁ for the screening and enumeration samples, respectively.

The probability of a positive result on the screening sample is, Qₙ(x) = 1-(1-p₁)(1-τ). The multinomial structure permits a simple derivation of the probability of detecting y cells in a 1 ml
sample, \( Q_y(x) \), given that there are \( x \) cells in the egg. This event entails the event, \( A \), of \( y \) cells in volume \( v_{21} \), which has the binomial probability with parameters \( x \) and \( p_2(1-\tau) \). However, \( A \) includes the disqualifying event \( B \) of \( x_{11} = 0 \) and \( x_{21} = y \); thus, the probability of this event must be subtracted from the probability of \( A \). The probability of \( B \) is the multinomial event with parameters \( x \), \( p_2(1-\tau) \) and \( 1-(p_1+p_2)(1-\tau) \) for the occurrence of \( x_{11} = 0 \), \( x_{21} = y \), and the remaining cells \( x-y \). Thus, performing the subtraction, it is derived that:

\[
Q_y(x) = \sum_{k=0}^{x} \binom{x}{k} (p_2(l-\tau))^k (1-p_2(l-\tau))^{x-k} \left[ 1 - \left( p_2(l-\tau) \right)^{x-y} \right]
\]

(C9)

where \( y = 0, 1, ..., x \). The likelihood of obtaining \( y \) cells in a 1 ml sample is:

\[
Q_y = \int_{\theta}^{\infty} \left( l - \theta \right) \sum_{y} Q_y(x) dF(x | \theta)
\]

(C10)

and the likelihood of the screening test being negative is:

\[
Q_y = \int_{\theta}^{\infty} \left( l - \theta \right) \sum_{x} Q_x(x) dF(x | \theta)
\]

(C11)

For the purposes of this analysis, it was assumed that the distribution of SE cells in egg yolks can be described as bi-modal: (i) a low-valued population: the 18 enumeration-negative results represent one subpopulation of the distribution for which there was no growth. This subpopulation represents \( Ev \) contaminations; and (ii) high-valued population: the 3 enumeration-positive results represent the other subpopulation for which either there was growth or an initial high level. This subpopulation represents \( Ey \) contaminations. Below, the distribution and uncertainty for initial contamination of each subpopulation is determined.

**Low-Valued Population**

**Modeling the distribution for the low-valued population**

For the low-valued population representing no growth and \( Ev \) contaminations, assume that \( F \) is a non-zero Poisson distribution with parameter \( \rho \), with probability increment, \( dP(k|\rho) \) equal to:

\[
p(x=k) = \frac{\rho^k e^{-\rho}}{k!}
\]

(C12)

where \( k = 1, 2, \) etc. This distribution was selected for convenience, although other distributions such as the logarithmic series distribution could be considered.
**Modeling the likelihood of a negative enumeration sample**

The likelihood of a negative enumeration sample, i.e., a sample that did not experience growth, was computed by first noting that:

\[
\int_0^\infty Q_\rho(k) dP(k | \rho) = (e^\rho - 1)^{-1} \sum_{k=0}^{\infty} \rho^k \left( \frac{(1 - p_{HI})^k - (1 - p_{HI} - p_{HI})^k}{k!} \right)
\]

where \( p_{j1} = p_j(1-\tau) \), so that:

\[
Q_\rho = (e^\rho - 1)^{-1} \left[ (e^\rho - 1)^{-1} - \rho (1 - p_{HI}) - e^\rho (1 - p_{HI} - p_{HI}) \right] \tag{C14}
\]

where \( 1 - w \) is the percentage of eggs that are contaminated, given that the eggs are from the low-value population or negative.

**Modeling the likelihood of a negative screening sample**

Similarly, the likelihood that the screening test is negative is:

\[
Q_{\rho 0} = (e^\rho - 1)^{-1} \left[ (e^\rho - 1)^{-1} - \rho (1 - p_{HI}) - e^\rho (1 - p_{HI} - p_{HI}) \right] \tag{C15}
\]

For example, if \( \rho = 2, \tau = 0.2, \) representing an 80% recovery, and \( w = 0.95, \) then \( Q_0 = 3.4\% \).

**Modeling the probability of observing 1 or more cells for an enumeration sample**

Of further interest was the probability of observing 1 or more cells for an enumeration sample, \( Q_{>1} \). From Equation C2, assuming the non-zero Poisson distribution, or from \( 1 - Q_{>1}Q_0 \), \( Q_{>1} \) was determined to be:

\[
Q_{>1} = (1 - w)(e^\rho - 1)^{-1} \left[ (e^\rho - 1)^{-1} - \rho (1 - p_{HI}) - e^\rho (1 - p_{HI} - p_{HI}) \right] \tag{C16}
\]

In the above example, \( Q_{>1} = 0.384\% \), which from 871 samples would imply an expected 3.345 samples for which enumeration would be 1 cell or more? The measure of fit statistic is -2 log-likelihood ratio:
where \( n_\) is the number of samples that were not positive by the screening test and \( n_0 \) is the number of samples with enumeration results of non-detect. The measure of fit statistic has a value of 12, which indicates a lack of fit, when compared to the percentiles of a chi-square with 1 degree of freedom.

The parameter, \( \rho \), was determined by assuming \( \tau = 0.2 \); setting the parameter \( Q_{1} \) so that the probability of no enumeration sample with a positive result from the \( n + n_0 = 871 \) screening samples is 25%; and minimizing \( \mathcal{L} \). A solution for \( \rho \) and \( w \) was obtained by constructing the Lagrangean equations (with one multiplier), setting the derivatives equal to zero, and solving in Mathcad® 7. For these constraints, \( \rho \) was determined to be 1.3922, \( w = 0.9681 \) (or about 3% of the eggs were contaminated in the yolk or vitelline membrane) and \( \mathcal{L} = 2.773 \), which is significant with \( P\)-value = 0.1, indicating a moderate degree of fit. Uncertainties of these values are determined by considering \( n_\) and \( n_0 \) as random variables arising from a multinomial distribution with the number parameter equal to 874 samples.

### Possibility of Egg Contamination Before or After Lay

Eggs that are members of the low-valued population are assumed to have experienced little growth. This suggests the contamination is not in the yolk, as SE cells in the yolk would likely leave lag phase quickly and be detected easily. This is important as yolk contaminations can lead to substantial numbers of cells within an egg over a short period. Therefore, the need to distinguish between yolk and vitelline membrane contaminations is necessary to establish a realistic estimate of risk. The low-valued eggs might represent small level contaminations in the vitelline membrane due to a migration event from the albumen after the egg was laid. This suggests the contamination is within a location less likely to exhibit rapid growth and could only be within this location from the time of lay to collection (1 day maximum). Eggs with these conditions would experience little growth. However, it is possible the contamination took place before the egg was laid, i.e. direct contamination of the yolk or vitelline membrane and the cells were still in the lag phase. This latter possibility could suggest the contamination was in the vitelline membrane or the yolk for up to 24 hrs within the hen plus the time before collection. Under these conditions, growth would be more likely and risk would be greater. To evaluate these possibilities and therefore establish risk for the low-valued population, an evaluation of possible growth, or actually, no growth, needs to be examined.

To determine the possibility of no growth for the low-valued population, the probability of no growth for an egg a particular age was first calculated. An initial contamination of 1 to 15 cells was assumed. This was followed by estimating the probability of no growth for a randomly selected egg from a population of eggs. Therefore, assuming the contamination to be uniformly distributed over a time for potential contamination, the probability for no growth before or after lay was determined. To determine the probability of no growth for an egg of a particular age, equations were developed that describe stochastically the growth of SE cells. The temperature of the egg, as a function of time, was assumed to be:

\[
T(t) = \min(T_a, T_i) + (T_i - T_a)e^{-x/2} \]

(C18)
where $T_a$ is the ambient temperature, $T_i$ is the initial egg temperature, $t_{lay}$ is the time that the egg was laid, relative to the time that the contamination entered the yolk or vitelline membrane, and $k$ is the exponential cooling rate (natural logarithm units per day). If the contamination enters the yolk or vitelline membrane before the egg is laid, then there are values of $t$ such that $t-t_{lay}$ is negative. Assumptions are: 1) $T_i = 41.1^\circ C$ - the body temperature of the hen; 2) $T_a = 24.4^\circ C$ - room temperatures where the hens reside is maintained at about $24.4^\circ C$; 3) $k = 0.3(24) = 7.2/day$ based on study of cooling rates of eggs in open stacks; and 4) as temperature changes, the ratio of the lag to the generation time, $Rat$, remains constant. For these calculations, it was assumed that $Rat = 5$.

For these calculations, it was assumed that the cells are found in the vitelline membrane, and that the egg yolks contain anti-SE antibodies. With these assumptions, the probabilities of no growth by time $t$ for assumed initial number of SE cells, ranging from 1-15, at the beginning of their lag phase at the time of lay ($t_{lay} = 0$) are given in Table C4. The expected value of the probability of no growth is given in the last row, computed assuming a Poisson distribution with parameter 1.392. The probability of greater than 15 SE cells is very small.

### Table C4 Probability of No Growth by Age (Days) When There Are an Assumed Initial Number of Cells That Contaminate the Vitelline Membrane (EV Eggs) at Lay. The Expected Value was Computed Assuming a Poisson Distribution with Parameter 1.392.

<table>
<thead>
<tr>
<th>Initial Number Cells</th>
<th>Time (days)</th>
<th>0.05</th>
<th>0.10</th>
<th>0.25</th>
<th>0.50</th>
<th>0.80</th>
<th>1.00</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>p0n</td>
<td>0.987</td>
<td>0.966</td>
<td>0.912</td>
<td>0.842</td>
<td>0.780</td>
<td>0.733</td>
</tr>
<tr>
<td>2</td>
<td>p0n</td>
<td>0.974</td>
<td>0.931</td>
<td>0.829</td>
<td>0.713</td>
<td>0.605</td>
<td>0.543</td>
</tr>
<tr>
<td>3</td>
<td>p0n</td>
<td>0.961</td>
<td>0.896</td>
<td>0.755</td>
<td>0.602</td>
<td>0.474</td>
<td>0.410</td>
</tr>
<tr>
<td>4</td>
<td>p0n</td>
<td>0.948</td>
<td>0.864</td>
<td>0.687</td>
<td>0.598</td>
<td>0.367</td>
<td>0.295</td>
</tr>
<tr>
<td>5</td>
<td>p0n</td>
<td>0.932</td>
<td>0.837</td>
<td>0.624</td>
<td>0.432</td>
<td>0.287</td>
<td>0.221</td>
</tr>
<tr>
<td>6</td>
<td>p0n</td>
<td>0.927</td>
<td>0.807</td>
<td>0.574</td>
<td>0.370</td>
<td>0.265</td>
<td>0.168</td>
</tr>
<tr>
<td>7</td>
<td>p0n</td>
<td>0.916</td>
<td>0.777</td>
<td>0.517</td>
<td>0.306</td>
<td>0.174</td>
<td>0.118</td>
</tr>
<tr>
<td>8</td>
<td>p0n</td>
<td>0.897</td>
<td>0.751</td>
<td>0.470</td>
<td>0.258</td>
<td>0.134</td>
<td>0.088</td>
</tr>
<tr>
<td>9</td>
<td>p0n</td>
<td>0.885</td>
<td>0.723</td>
<td>0.428</td>
<td>0.218</td>
<td>0.104</td>
<td>0.064</td>
</tr>
<tr>
<td>10</td>
<td>p0n</td>
<td>0.875</td>
<td>0.699</td>
<td>0.390</td>
<td>0.184</td>
<td>0.081</td>
<td>0.045</td>
</tr>
<tr>
<td>11</td>
<td>p0n</td>
<td>0.869</td>
<td>0.674</td>
<td>0.352</td>
<td>0.155</td>
<td>0.064</td>
<td>0.035</td>
</tr>
<tr>
<td>12</td>
<td>p0n</td>
<td>0.851</td>
<td>0.651</td>
<td>0.322</td>
<td>0.131</td>
<td>0.048</td>
<td>0.027</td>
</tr>
<tr>
<td>13</td>
<td>p0n</td>
<td>0.841</td>
<td>0.627</td>
<td>0.293</td>
<td>0.110</td>
<td>0.038</td>
<td>0.019</td>
</tr>
<tr>
<td>14</td>
<td>p0n</td>
<td>0.829</td>
<td>0.606</td>
<td>0.267</td>
<td>0.093</td>
<td>0.030</td>
<td>0.014</td>
</tr>
<tr>
<td>15</td>
<td>p0n</td>
<td>0.818</td>
<td>0.584</td>
<td>0.241</td>
<td>0.079</td>
<td>0.023</td>
<td>0.010</td>
</tr>
<tr>
<td>Expected Value For No Growth</td>
<td></td>
<td>0.976</td>
<td>0.936</td>
<td>0.843</td>
<td>0.741</td>
<td>0.646</td>
<td>0.592</td>
</tr>
</tbody>
</table>

Results from Table C4 indicate, for instance, that if there were 10 SE cells in the beginning of lag phase at lay, in a full day, there was a 4.7% chance that no growth would take place. Within half a day, there was an 18.4% chance that no growth would take place. To determine the probability of no growth for a randomly drawn egg from a population of eggs, let $Eprob_0(t|t_b)$ be the expected value of no growth for an egg of age $t$, given contamination occurred at time $t_b$. From the results in the last row of Table C4 and others for different times not shown, an
approximation of $E_{prob0}(t|t_b)$ of the form $d/(1+b t^c)$, where $b$, $c$ and $d$ are parameters with values depending on $t_b$ and were estimated from nonlinear regressions. (For the case of Table C4 ($t_b = 0$), $d = 1.01$, $c = 0.92$ and $b = 0.70$). If it is assumed ages of these low valued population eggs when sampled were uniformly distributed over 0 to 1 day, the expected value that there would be no SE growth, $E_{prob}(t_b)$ in a randomly drawn egg from this population would be the integral of the $d/(1+b t^c)$ from $t_b$ to 1. Results for selected times SE of contamination before the egg is laid are given in Table C5. Figure C1 is a graph of the natural logarithm of the computed probability of no growth versus the time of contamination and a smoothed fourth degree polynomial fit.

**Table C5 Probability there would be no growth of SE in a randomly selected egg, given the contamination entered the vitelline membrane at $t$ days before the egg was laid.**

<table>
<thead>
<tr>
<th>Fraction of Day After Lay</th>
<th>-1</th>
<th>-0.9</th>
<th>-0.75</th>
<th>-0.5</th>
<th>-0.25</th>
<th>0</th>
<th>0.15</th>
<th>0.25</th>
<th>0.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Probability of No Growth</td>
<td>29.3%</td>
<td>31.9%</td>
<td>36.4%</td>
<td>45.9%</td>
<td>58.6%</td>
<td>75.9%</td>
<td>83.6%</td>
<td>86.6%</td>
<td>92.6%</td>
</tr>
</tbody>
</table>

![Figure C1](image)

Figure C1 Computed natural logarithm of the probability of no growth versus given times of contamination relative to when the egg is laid. The smoothed line is fourth degree polynomial least squares regression.

The above calculation of the probability of no SE growth assumes that the egg is sampled at an age greater than the time of contamination; that is, it assumes that the eggs are sampled at a random time between $\max(0, t_b)$ and 1. Eggs sampled before $t_b$ of course would show no growth. Including this latter possibility, the probability of a randomly selected egg showing no growth is:
Using the derived polynomial regression, $PN_g$ was derived to be 71%. This value can be visualized as an average of the conditional probabilities of no growth given that the egg became contaminated before and after lay. The former probability is 48% and the latter is 94%. If the beginning of the contamination were assumed to be uniformly distributed over the interval (-1, 1), then 75% of the contaminations would be sampled. Of the positive samples, 67% would be from contaminations that occurred before lay. Consequently, even with a draw of 8 such eggs, the probability of at least 1 egg experiencing growth would be high.

This analysis shows that the assumption that a significant percentage of these eggs are contaminated in the yolk before lay is not plausible. Consequently, for these eggs, it was assumed that the contamination occurred after lay and that the contamination resides within the vitelline membrane.

**RESULTS AND ASSUMPTIONS USED FOR LOW-VALUED POPULATIONS**

1) For the low value population, it was assumed that the contamination was located within the vitelline membrane and occurred post-lay.

2) For the low value population, it was assumed that the number of SE cells at the beginning of a contamination was distributed as a Poisson, without zeros (Equation C6), with parameter, $\rho$, equal to 1.3922.

3) The age of the eggs when the contamination begins in vitelline membrane was distributed uniformly in the interval (0, 1). The assumption that the ages of the eggs when sampled were uniform implies that an expected 50% of the contaminated eggs of this type would not have been sampled.

4) The fraction of low valued positive eggs was estimated as $2(1-w)$, where $w = 0.96813$.

5) Uncertainty of $\rho$ and $w$ was determined by assuming the vector: $n_n, n_0, and n_+$, is distributed as a multinomial distribution with probability parameters: $z_+, z_0, and z_+ = 1- z_+ - z_0$, and number parameter 874. The values from the trinomial distribution were generated by a sequence of two binomial distributions, where the first one generated a value of $n_+$, labeled $m_+$ for this discussion, from the binomial (874, 3/874), and then the second one generated a value of $n_+$ from the binomial(874- $m_+$, 871/874). The strategy is based on the probability law: $P(A, B) = P(A|B)P(B)$, and noting that the conditional probability, $P(A|B)$, and the unconditional probability, $P(B)$, are distributed as binomial distributions. A bootstrap of 1,000 simulations was performed, and various transformations of the results were examined to find ones that were nearly normally distributed. For $\rho$, the transformation $g(\rho) = \ldots$

---

This time interval indicates 24 hrs prior to egg lay through 24 hrs post-lay. It represents a contamination event within the hen that could have taken place before egg lay through the maximum time the egg could have remained before collection and analysis.
\( \ln(1+\ln(\rho)) \) was nearly normally distributed with the mean equal to 0.27671, for which the inverse transformation is 1.3755, and the standard deviation equal to 0.17325 (the skewness = 0.062 and the kurtosis = 0.1873). To adjust for the non-zero kurtosis, Equation C10 was used. For \( w \), the transformation, \( h(w) = (-\ln(w))^{0.75} \) was nearly normally distributed with the mean equal to 0.077364, for which the inverse transformation is 0.96754, and the standard deviation equal to 0.015473 (the skewness = 0.022 and the kurtosis = 0.007). The correlation of \( g(\rho) \) and \( h(w) \) is 0.99961.

**High-Valued Population**

For the high-valued population, the age of the sampled egg is critical as the SE levels observed by Gast and Holt\(^5\) could quickly lead to substantial levels within an egg. For these three eggs (4, 27 and 67 cfu/mL of yolk), it was assumed that there was SE growth taking place when sampled, so that the time that the growth of SE begins is important. Therefore, to identify the initial contamination levels of this subpopulation, the amount of SE growth that might have occurred for these three eggs needed to be calculated. For this, it was assumed 1) that the SE cells are in the vitelline membrane or yolk; and 2) that at least one of the cells is out of its lag phase.

To begin to identify the initial level of the high-valued results (4, 27 and 67 cfu/mL of yolk), it needs to be determined if these three levels could be explained using the Poisson assumption derived above for the initial distribution of the low-valued population. Since the predicted exponential growth rate per day for SE cells in the vitelline membrane at 24.4°C was approximately 3.8 log\(_{10}\)/day, it is possible to explain the three results by assuming a single cell was out of its lag phase sometime before the egg was sampled at less than or equal to 1 day old. However, using the Poisson distribution would not account for the uncertainty that these three results present and therefore was not used in the analysis of this subpopulation.

In addition, it is unclear if by the time the eggs were sampled, growth can be assumed to have taken place. It cannot be dismissed that the ages of some of these eggs when sampled could be less than just a few hours and that these eggs represent serious contaminations with high numbers of SE cells, or that at least a few of them were in the exponential growth phase. Alternatively, it is possible initial contaminations of the yolk or vitelline membrane occurred early in an egg’s development, close to 24 hrs before the egg was laid. This could be due to colonization of the ovary and/or the upper oviduct by SE. By the time the egg was laid, there could be a large and growing contamination, even if the initial number of SE cells in the original contamination was not large. Therefore, the assumption is possible. The complexities are infinite, and the information is infinitesimal.

Nonetheless, the three high results suggest the SE within these yolks could have been growing at the time of collection. If this were true, the time and temperature conditions of the egg would likely influence the expected value of SE. That is, if the eggs were stored differently from the conditions of Gast and Holt\(^5\), then it would be expected that the measured SE levels would differ from those reported. Consequently, it was necessary to make some adjustments accounting for time and temperature storage conditions.

To make this possible, the high-valued population was modeled in such a way to account for the time and temperature differences potentially experience by an individual egg. The strategy was to back-calculate, to a time, \( t(1) \), when it is imagined one cell would have existed out of lag phase, from which the expected subsequent growth would provide a level equal to that at the age that the eggs were sampled under the same conditions.
An Example to Demonstrate Difficulty an Initial Set of Conditions

It is difficult to determine the possible initial states that could give rise to a specific set of results. An example is given that demonstrates the difficulty of determining an initial set of conditions by back-calculating possible growth scenarios, and to show the type of calculations that were done.

In general, there could be \( N \) SE cells initially in the contamination, where only some of them actually begin to grow (divide), while the others do not until later. Suppose the age of an egg when sampled was \( z \), and the obtained result was \( x \). It was assumed that, at the time of sampling, SE cells were growing. Let it be assumed that there was a time, \( t \), less than \( z \), when a single SE cell left its lag phase, and the measured value of \( x \) represents the number of that cell’s progeny.

The actual distribution of the increase of the number of cells is a geometric distribution, with parameter \( p \), where \( p \) is the expected value of \( X \); specifically, the cumulative distribution of the random variable \( X \), representing the increase number of cells, is:

\[
G(X) = 1 - (1 - p)^X + 1
\]  

so that the \( q \)th percentile, \( X_q \), is:

\[
X_q = \frac{\ln(1 - q)}{\ln(1 - p)}
\]

If the result of 67 cfu/mL represented the 99th percentile of the distribution given above, then \( p = 1/261.3 \), or that the expected value would be 261.3, or, in log base 10 units, 2.42 \( \log_{10} \) cfu/ml. If the result represented the 1st percentile, then the expected value would be 5.07 \( \log_{10} \) cfu/ml. This latter possibility could happen for the assumed model if a single cell left its lag phase 45 minutes before the egg was laid and the egg was 1 day old when sampled. However, another possibility is that the SE cell left its lag phase 12 hours before the egg was laid and the age of the egg when sampled was about 1.7 hours. If this were to happen, employing the growth model developed here, it would be expected that the egg at 1 day old would have about 9 \( \log_{10} \) SE cells.

One can go farther and assume that the distribution of SE levels is lognormal for the population sampled and that the results represent a simple random sample from this distribution. The \( p \)th percentile of predicted individual values was determined using:

\[
\hat{x} + k_p (\bar{x} + \frac{1}{\hat{k}})^{\frac{1}{2}}
\]

C-17
where $k_p$ is $p^{th}$ percentile of the $t$-distribution with 2 degrees of freedom. Assuming that results represent cfu/mL in 15 ml of the yolk, and the recovery is 80%, then the upper 99.5$^{th}$ percentile of the individual values is approximately $9.5 \log_{10}$. However, as the ages of the eggs could be between 0 and 1 day old, these results could represent eggs with levels of SE above $9 \log_{10}$ at 1 day. There are neither data nor scientific theory known that would eliminate this possibility.

**ESTIMATING THE INITIAL CONTAMINATION OF THE HIGH-VALUED POPULATION**

To estimate the initial contamination of this subpopulation, it was assumed that the eggs, when sampled, were less than 1 day old. The ages of the eggs when sampled are unknown, so it was assumed that the ages are uniformly distributed over the interval $(0, 1)$. Further, calculations are performed when it was assumed that SE growth takes place in the yolk ($E_y$ contaminations), and that the eggs contain antibodies to SE. To distinguish if the three high-valued results represent $E_v$ or $E_y$ contamination, data from the study of Gast and Holt$^7$ were used. In this study, 29 eggs were found to be contaminated in the yolk or vitelline membrane. Analysis of the internal yolk contents found that only 3 of the 29 were actually yolk contaminations, excluding the membrane. It was assumed that 3 eggs corresponding to the high values from the study by Gast and Holt$^5$ represent $E_y$ contaminations.

The exponential growth rates, $\mu(t)$, as a function of time through the temperature, are described in annex E, and the temperature profile is given in Equation C20. Thus, the expected relative growth between two times, $a$ and $b$, is:

$$r(b, a) = \int_a^b \mu(t) dt$$

(C23)

The strategy for estimating the initial contamination of the high-valued population is as follows: The value $t(1)$ represents the time that a single SE cell left its lag phase and entered into the exponential phase of growth. If $t(1) < 0$ then, in the risk assessments, the calculation of the number of progeny cells when the egg is laid is made assuming a temperature of $41.1^\circ$C. The likelihood of observing $x$ (ignoring measurement error), given the age of the egg, $z$, and the time $t = t(1)$ is:

$$L(x | z, t(1)) = p(z, t(1)) (1 - p(z, t(1)))^{x-1}$$

(C24)

where $p(z, t) = r(z, t)^{-1}$.
Let $h(t|\alpha, \beta)$ be the beta distribution of $t(1)$ within the interval $[-1,1]$, representing the possible times that a SE cell could contaminate the yolk (lower bound, -1, is based on the assumption that it takes approximately 24 hours for an egg to be formed and laid), where $\alpha$ and $\beta$ are the parameters, whose values are to be estimated. Because it is assumed that the ages of the eggs when sampled are uniform and observed values of $x$ only occur when $t(1) < z$, the unconditional likelihood value, $Lik(x|\alpha, \beta)$ is:

$$Lik(x|\alpha, \beta) = \int_{-1}^{1} h(t|\alpha, \beta) \int_{0}^{t(1)} p(z, \tau) (1-p(z, \tau))^{x-1} dz d\tau$$

(C25)

Working with this equation to determine $\alpha$ and $\beta$ is difficult. Thus, a simplified approach for determining values for $\alpha$ and $\beta$ is used as follows: assume that the distribution of ages of the eggs when sampled is the uniform distribution; then, for a given $t(1)$, the expected value of $\ln(x)$, $TE_x(t(1))$, would be:

$$TE_x(t(1)) = \int_{0}^{t(1)} h(t|\alpha, \beta) dz$$

(C26)

Using Equation C27, a value of $t(1)$ was estimated, assuming that $TE_x(t(1)) = \ln(x)$. For $x = 1,256$, corresponding to the sample with measured value of 67 cfu/ml, the value of $t(1)$ is 0.101; for the sample with value 27 cfu/ml, the value of $t(1)$ is 0.140; and for the value of 4 cfu/ml, $t(1)$ = 0.253. These values can be considered the observed values of a sample from a population of times, with density $h(t|\alpha, \beta)$. However, the probabilities of samples being observed as growth samples are not the same because, for times $t>0$, the probability of the egg being sampled at an age less than $t$ is 1-$t$. Thus, the likelihood, $Lik1(t)$, of an observation is:

$$Lik1(t) = h(t|\alpha, \beta)\min(1,1-t)$$

(C27)

reflecting the lower likelihood of being observed, if $t$ is positive. Stirling’s formula, $(x/e)^x(2\pi x)^{0.5}$, for approximating the factorial of $x$ was used for large values of $\alpha$ and $\beta$. Maximum likelihood estimators (MLE) of $\alpha$ and $\beta$ were derived using the logarithmic transformations, $\ln(\alpha)$, and $\ln(\beta)$, and the error covariance matrix derived applies to these transformed values.

Calculations were performed on Mathcad® 7. The MLE estimates of $\ln(\alpha)$ and $\ln(\beta)$ are 4.9065 and 4.6066, with standard errors of 0.8177 and 0.8172, respectively, and a correlation of 0.9957, computed from the Hessian matrix of second derivatives. Figure C2 depicts the curves of the log-likelihood function, fixing $\ln(\alpha)$ and varying $\ln(\beta)$, for values of $\ln(\alpha)$, including the MLE, 4.5, 4.8, 5.0, and 5.3. The curve with the MLE is bolded. As is evident, the log-likelihood function is flat near the MLE estimates, contributing to the relatively large standard errors. The above approach does not account for the uncertainty that would exist for the values of the parameters due to the uncertainty of the values of the exponential growth rates. The affect of
different assumed exponential growth rates would be to translate the times by amounts so that the expected value of the distribution would be most affected.

![Plot of loglikelihood curves for fixed values of ln(α), where the x-axis is a range of ln(β) values. The middle curve (bolded) was generated with the MLE of ln(α); the maximum occurs for the MLE estimate of ln(α) = 4.907 and ln(β) = 4.607.](image)

**RESULTS AND ASSUMPTIONS USED HIGH-VALUED POPULATIONS**

1) It was assumed that the time, \( t(1) \), that one SE cell in the interior of the yolk \( (E_y \) contamination) enters the exponential growth phase was distributed as a beta distribution over the interval \([-1, 1]\) with parameters \( \alpha = e^{4.9065} \) and \( \beta = e^{4.6066} \).

2) The uncertainty was determined by assuming that \( \ln(\alpha) \) and \( \ln(\beta) \) were distributed as a \( t \)-distribution with means equal to 4.9065 and 4.6066, respectively, with standard errors of 0.8177 and 0.8172, respectively, and a correlation of 0.9957. Because there are three observations, there are 2 degrees of freedom for determining the standard deviation from the mean. Thus, 2 degrees of freedom are associated with these estimates. Formally, this would
imply that values of $\ln(\alpha)$ or $\ln(\beta)$ could be negative, or that values of $\alpha$ or $\beta$ are less than 1, a result that seems counter-intuitive. Consequently, generated values of $\alpha$ or $\beta$ that are less than 1 are set equal to 1.

3) The age of the eggs when sampled was assumed uniform over the interval [0, 1], so that any egg that was contaminated before detectable SE levels would have been sampled, while if contaminated after lay would have been sampled with a probability of $\frac{1}{2}$. Thus, there is a 75% chance that the contaminated eggs would have been sampled. Because 3 contaminations were detected, it is possible that 4 could have existed in the population. Consequently, the percentage of eggs that are members of the high value population is $(1.33)^{3/874}$. To determine the uncertainty of this estimate, the number of positive eggs was assumed distributed as 1.33 times a binomial distribution with parameters $3/874$ and 874.
REFERENCES


