Bacteriological Analytical Manual (BAM)

Chapter 5: *Salmonella*

**Authors:** Wallace H. Andrews (ret.), Hua Wang, Andrew Jacobson (ret.), Beilei Ge, Guodong Zhang, and Thomas Hammack

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Revision History

- September 2023: Section D.7 updated by requiring the use of both typical and atypical \textit{Salmonella} culture controls.
- April 2023: Chapter 5 was converted to PDF and labeled as BAM Chapter 5: \textit{Salmonella} (April 2023 Edition). Section C.28 updated to remove standing at room temperature and pH adjustment, which are steps not performed at the time of SLV and MLV.
- November 2022: Sections C.28 and E.9 updated to include screenshots for AB 7500 Fast setup and representative Genie instrument outputs, recommended GspSSD2.0 as the preferred LAMP master mix, and updated template preparation method to include a final centrifugation step. Updated section C.7a to include oysters.
- March 2022: Section C.28 updated to include dairy feed as a validated animal food matrix, AB 7500 Fast as an alternative LAMP assay platform, two additional reagent choices for LAMP, and removed duplicate testing recommendation for LAMP screening of \textit{Salmonella} in animal food protocol; Section E.9.d updated to include AB 7500 Fast as an alternative LAMP assay platform, two additional reagent choices for LAMP, and removed duplicate testing recommendation for LAMP confirmation of \textit{Salmonella} isolates protocol; Section E.11 was updated by linking SeqSero to Protocols.IO.
- October 2021: Section C.2.c updated to correct a typo; Section E.11 serotyping submission information and shipping point of contact updated.
- September 2021: Section D.7 updated to include serological-formulas of ATCC control cultures 12325 and 29934; Section E.9.c updated to remove SmartCycler instructions from the Real-time PCR confirmation test.
- June 2021: Section A.27 updated to include ultraviolet light source; Section C.7 updated by changing nut/nut meat preparation procedure from blending to soaking; Section C.23 updated by removing erroneous requirement to adjust the pH of preenrichment before incubation; Section C29 added method for spent sprout irrigation water from alfalfa, mung bean, and broccoli varieties; Section D.7 updated by adding Green Fluorescent Protein control strains.
- May 2021: Section C.28 updated to include an alternative positive control, a video demonstration, and software updates for LAMP screening for \textit{Salmonella} in animal food protocol; Section E.9.d updated to include a video demonstration and software updates for LAMP confirmation of \textit{Salmonella} isolates protocol; Section C.7.a updated to include mushrooms.
- February 2021: Removed text from section E.9.b.
- July 2020: Section c.10. Added sample set up for non-powder forms of allspice, cinnamon, cloves, and oregano. Added reference 17.
December 2019: Section C7. Limited the foods covered by Section C7 to those not listed elsewhere in Section C.

November 2019: Section C.10.c Added detailed procedure to prepare sample composites for oregano, cinnamon, allspice, and cloves.

November 2019: Section E9 updated to include additional options for identification of Salmonella.

July 2018: Section C7 revised to include vegetables not included in C23 or C27.

March 2018: Added real-time quantitative PCR for confirmation of Salmonella isolates protocol and validated preenrichment changes for leafy produce, herbs and sprouts; Vegetables removed from revised Section C7; Validated Preenrichment broth change in section C23.

August 2016: Added the Salmonella Flipbook, a pictorial general guide to aid analysts in the detection and identification of Salmonella growing on the plating media and screening tubes used in the BAM Chapter 5 Salmonella method. See Section E8. (Prepared By: Matthew J. Forstner, Laboratory Services, Minnesota Department of Agriculture). (PDF, 13Mb)

December 2015: A section for the Statens Serum Institute Procedure was added to Section E: Identification of Salmonella.

May 2014: The VITEK 2 method of Presumptive generic identification of Salmonella was updated.

February 2014: Section on Detection and isolation of Salmonella from shell eggs was replaced, and validation data and additional references were added in an Appendix.


November 2011: Addition to Section C: Preparation of foods for isolation of Salmonella: Leafy green vegetables and herbs.

February 2011: Removed link to Appendix 1: Rapid Methods for Detecting Foodborne Pathogens (now archived).

December 2007: Mamey pulp method added, and Section D revised.

June 2006: Eggs method revised for shell eggs and liquid whole eggs.

April 2003: Frog legs method, Lactic casein, Rennet casein, Sodium caseinate and Rabbit carcass methods revised, top ears and other dog chew toys added. Removed section A.25, Mechanical shaker.

October 2001: Extension of the applicability of the orange juice method in section C.19 to apple juice and apple cider.


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Introduction

Several changes are being introduced in this edition of BAM (8th Edition). The first change involves the expanded use of Rappaport-Vassiliadis (RV) medium for foods with both high and low levels of competitive microflora. In the previous edition, RV medium was recommended only for the analysis of shrimp. Based on the completion of AOAC precollaborative (5, 6) and collaborative (7, 8) studies, RV medium is now being recommended for the analysis of all foods, except guar gum. In addition, RV medium replaces selenite cystine (SC) broth for the analysis of high microbial load foods. RV medium replaces lauryl tryptose broth for use with dry active yeast. Tetrathionate (TT) broth continues to be used as the second selective enrichment broth. However, TT broth is to be incubated at 43°C for the analysis of high microbial load foods and at 35°C for the analysis of low microbial load foods, including guar gum.

The second change involves the option of refrigerating incubated preenrichments and selective enrichments of low-moisture foods for up to 72 h. With this option, sample analyses can be initiated as late as Wednesday or Thursday without weekend work being involved.

The third change involves reducing the period of incubation of the lysine iron agar (LIA) slants. In the former edition (BAM-7), triple sugar iron (TSI) agar and LIA slants were incubated at 35°C for 24 ± 2 h and 48 ± 2 h, respectively. Unpublished data have demonstrated that the 48 h reading of LIA slants is without diagnostic value. Of 193 LIA slants examined, all gave definitive results within 24 ± 2 h of incubation. No significant changes altered the final test result when the slants were incubated an additional 24 h. Thus, both the TSI and LIA slants are now incubated for 24 ± 2 h.

The fourth change involves the procedure for surface disinfection of shell eggs. In the previous edition (BAM-7), shell eggs were surface-disinfected by soaking in 0.1% mercuric chloride solution for 1 h followed by soaking in 70% ethanol for 30 min. Mercuric chloride is classified as a hazardous waste and is expensive to dispose of according to Environmental Protection Agency guidelines. In this edition (BAM-8) shell eggs are now surface-disinfected by soaking for at least 10 sec in a 3:1 solution consisting of 3 parts of 70% alcohol (ethyl or isopropyl) to 1 part of iodine/potassium iodide solution.

The fifth change involves the sample preparation of eggs. Egg contents (yolk and albumen) are thoroughly mixed before analysis. After mixing the egg contents, 25 g (ml) are added to 225 ml trypticase (tryptic) soy broth supplemented with ferrous sulfate.

A method for the analysis of guar gum has been included. When guar gum is preenriched at a 1:9 sample/broth ratio, a highly viscous, non-pipettable mixture results. Addition of the enzyme cellulase to the preenrichment medium, however, results in a readily pipettable mixture.

A method for orange juice (pasteurized and unpasteurized) has been included due to recent orange juice-related outbreaks.
The directions for picking colonies from the selective plating agars have been made more explicit to reflect the intent of the method. In the absence of typical or suspect colonies on the selective plating agars, it is recommended that atypical colonies be picked to TSI and LIA slants. This recommendation is based on the fact that up to 4% of all *Salmonella* cultures isolated by FDA analysts from certain foods, especially seafoods, during the past several years have been atypical.

Finally, since the publication of BAM-7, a 6-way comparison was conducted of the relative effectiveness of the three selective plating agars recommended in the BAM (*bismuth sulfite, Hektoen enteric, and xylose lysine desoxycholate agars*) and three relatively new agars (EF-18, xylose lysine Tergitol 4, and Rambach agars). Our results (9) indicated no advantage in replacing any of the BAM-recommended agars with one or more of the newer agars. Thus, the combination of selective plating agars recommended in BAM-7 remains unchanged.

### A. Equipment and Materials

1. Blender and sterile blender jars *(see BAM Chapter 1)*

2. Sterile, 16 oz (500 ml) wide-mouth, screw-cap jars, sterile 500 ml Erlenmeyer flasks, sterile 250 ml beakers, sterile glass or paper funnels of appropriate size, and, optionally, containers of appropriate capacity to accommodate composited samples

3. Sterile, bent glass or plastic spreader rods

4. Balance, with weights; 2000 g capacity, sensitivity of 0.1 g

5. Balance, with weights; 120 g capacity, sensitivity of 5 mg

6. Incubator, 35 ± 2 °C

7. Refrigerated incubator or laboratory refrigerator, 4 ± 2°C

8. Water bath, 49 ± 1°C

9. Water bath, circulating, thermostatically-controlled, 43 ± 0.2°C

10. Water bath, circulating, thermostatically-controlled, 42 ± 0.2°C

11. Sterile spoons or other appropriate instruments for transferring food samples

12. Sterile culture dishes, 15 × 100 mm, glass or plastic

13. Sterile pipets, 1 ml, with 0.01 ml graduations; 5 and 10 ml, with 0.1 ml graduations

14. Inoculating needle and inoculating loop (about 3 mm id or 10 5l), nichrome, platinum-iridium, chromel wire, or sterile plastic

15. Sterile test or culture tubes, 16 × 150 mm and 20 × 150 mm; serological tubes, 10 × 75 mm or 13 × 100 mm

16. Test or culture tube racks
17. Vortex mixer
18. Sterile shears, large scissors, scalpel, and forceps
19. Lamp (for observing serological reactions)
20. Fisher or Bunsen burner
21. pH test paper (pH range 6-8) with maximum graduations of 0.4 pH units per color change
22. pH meter
23. Plastic bags, 28 × 37 cm, sterile, with resealable tape. (Items 23-24 are needed in the analysis of frog legs and rabbit carcasses.)
24. Plastic beakers, 4 liter, autoclavable, for holding plastic bag during shaking and incubation
25. Sponges, non-bactericidal (Nasco cat # B01299WA), or equivalent
26. Swabs, non-bactericidal, cotton-tipped.
27. Ultraviolet (UV) light source with an emission wavelength between 360 to 400 nm when Green Fluorescent Protein (GFP)-UV control strains are used

B. Media and Reagents

For preparation of media and reagents, refer to Methods 967.25-967.28 in *Official Methods of Analysis* (1).

1. Lactose broth (M74)
2. Nonfat dry milk (reconstituted) (M111)
3. Selenite cystine (SC) broth (M134)
4. Tetrathionate (TT) broth (M145)
5. Rappaport-Vassiliadis (RV) medium (M132). NOTE: RV medium must be made from its individual ingredients. Commercial formulations are not acceptable.
6. Xylose lysine desoxycholate (XLD) agar (M179)
7. Hektoen enteric (HE) agar (M61)
8. Bismuth sulfite (BS) agar (M19)
9. Triple sugar iron agar (TSI) (M149)
10. Tryptone (tryptophane) broth (M164)
11. Trypticase (tryptic) soy broth (M154)
12. Trypticase soy-tryptose broth (M160)
13. MR-VP broth (M104)
14. Simmons citrate agar (M138)
15. Urea broth (M171)
16. Urea broth (rapid) (M172)
17. Malonate broth (M92)
18. Lysine iron agar (LIA) (Edwards and Fife) (M89)
19. Lysine decarboxylase broth (M87)
20. Motility test medium (semisolid) (M103)
21. Potassium cyanide (KCN) broth (M126)
22. Phenol red carbohydrate broth (M121)
23. Purple carbohydrate broth (M130)
24. MacConkey agar (M91)
25. Nutrient broth (M114)
26. Brain heart infusion (BHI) broth (M24)
27. Papain solution, 5% (M56a)
28. Cellulase solution, 1% (M187)
29. Tryptose blood agar base (M166)
30. Universal preenrichment broth (M188)
31. Universal preenrichment broth (without ferric ammonium citrate) (M188a)
32. Buffered peptone water (M192)
33. Dey-Engley broth (M193)
34. Potassium sulfite powder, anhydrous
35. Chlorine solution, 200 ppm, containing 0.1% sodium dodecyl sulfate (R12a)
36. Ethanol, 70% (R23)
37. Kovacs' reagent (R38)
38. Voges-Proskauer (VP) test reagents (R89)
39. Creatine phosphate crystals
40. Potassium hydroxide solution, 40% (R65)
41. 1 N Sodium hydroxide solution (R73)
42. 1 N Hydrochloric acid (R36)
43. Brilliant green dye solution, 1% (R8)
44. Brom cresol purple dye solution, 0.2% (R9)
45. Methyl red indicator (R44)
46. Sterile distilled water
47. Tergitol anionic 7 (R78)
48. Triton X-100 (R86)
49. Physiological saline solution, 0.85% (sterile) (R63)
50. Formalinized physiological saline solution (R27)
51. *Salmonella* polyvalent somatic (O) antiserum
52. *Salmonella* polyvalent flagellar (H) antiserum
53. *Salmonella* somatic group (O) antisera: A, B, C1, C2, C3, D1, D2, E1, E2, E3, E4, F, G, H, I, Vi, and other groups, as appropriate
54. *Salmonella* Spicer-Edwards flagellar (H) antisera
55. Modified Buffered Peptone Water (M192b)

C. Preparation of foods for isolation of *Salmonella*

The following methods are based on the analysis of a 25 g analytical unit at a 1:9 sample/broth ratio. Depending on the extent of compositing, add enough broth to maintain this 1:9 ratio unless otherwise indicated. For samples not analyzed on an exact weight basis, e.g., frog legs, refer to the specific method for instructions.

1. **Dried egg yolk, dried egg whites, dried whole eggs, liquid milk (skim milk, 2% fat milk, whole, and buttermilk), and prepared powdered mixes (cake, cookie, doughnut, biscuit, and bread), infant formula, and oral or tube feedings containing egg.** Preferably, do not thaw frozen samples before analysis. If frozen sample must be tempered to obtain analytical portion, thaw suitable portion as rapidly as possible to minimize increase in number of competing organisms or to reduce potential of injuring *Salmonella* organisms. Thaw below 45°C for 15 min with continuous agitation in thermostatically controlled water bath or thaw within 18 h at 2-5°C. Aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. For nonpowdered samples, add 225 ml sterile lactose broth. If product is powdered, add about 15 ml sterile lactose broth and stir with sterile glass rod, spoon, or tongue depressor.
to smooth suspension. Add 3 additional portions of lactose broth, 10, 10, and 190 ml, for total of 225 ml. Stir thoroughly until sample is suspended without lumps. Cap jar securely and let stand 60 ± 5 min at room temperature. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2 with sterile 1 N NaOH or 1 N HCl. Cap jar securely and mix well before determining final pH. Loosen jar cap about 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

2. **Eggs**

   a. **Shell eggs** [14,15]. Eggs with chipped, cracked, or broken shells are not included in the sample. Remove any adherent material from the shell eggs surface. Disinfect egg surface with a solution consisting of 3 parts of 70% alcohol (ethyl or isopropyl) to 1 part iodine/potassium iodide solution. Prepare 70% alcohol solution either by diluting 700 ml 100% alcohol with sterile distilled water for a final volume of 1,000 ml or by diluting 700 ml 95% alcohol with sterile distilled water for a final volume of 950 ml. Prepare iodine/potassium iodide solution by dissolving 100 g potassium iodide in 200-300 ml sterile distilled water. Add 50 g iodine and heat gently with constant mixing until the iodine is dissolved. Dilute the iodine/potassium iodide solution to 1,000 ml with sterile distilled water. Store iodine/potassium iodide solution in an amber glass-stoppered bottle in the dark if not used immediately. Prepare the disinfection solution by adding 250 ml iodine/potassium iodide solution to 750 ml 70% alcohol solution and mix well. Submerge eggs in disinfection solution for 10 seconds (make sure not less than 10 seconds). Remove eggs from the solution and allow to air dry. Each sample shall consist of twenty (20) eggs, for a total of fifty (50) samples per poultry house. Eggs are cracked aseptically into a 4L sterile beaker or other suitable container by gloved hands, with a change of gloves between samples. Mix samples thoroughly with a sterile tool by gloved hands until yolks are completely mixed with the albumen, with a change of gloves between samples. Preenrich the 20-egg sample by adding 2 L sterile trypticase soy broth (TSB; room temperature) and mix well with a sterile tool. Cover securely and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

   See *Salmonella Appendix* for validation data

   b. **Liquid whole eggs (homogenized).** Combine fifteen (15) 25 ml test portions into a 375 ml composite contained in a 6-liter Erlenmeyer flask. Composites are held at room temperature (20-24°C) for 96 ± 2 h. After 96 ± 2 h, add 3,375 ml sterile trypticase soy broth (TSB) supplemented with ferrous sulfate, as described above, and mix well by swirling. Let stand 60 ± 5 min at room temperature. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

   c. **Hard-boiled eggs (chicken, duck, and others).** If the shell eggs are still intact, disinfect the shells as described above and aseptically separate the shells from the
eggs. Pulverize the eggs (egg yolk solids and egg white solids) aseptically and weigh 25 g into a sterile 500 ml Erlenmeyer flask or other appropriate container. Add 225 ml trypticase soy broth (TSB) (without ferrous sulfate) and mix well by swirling. Continue as in D.1-11.

3. **Nonfat dry milk**

   a. **Instant.** Aseptically weigh 25 g sample into sterile beaker (250 ml) or other appropriate container. Using sterile glass or paper funnel (made with tape to withstand autoclaving), pour 25 g analytical unit gently and slowly over surface of 225 ml brilliant green water contained in sterile 500 ml Erlenmeyer flask or other appropriate container. Alternatively, 25 g analytical units may be composited and poured over the surface of proportionately larger volumes of brilliant green water. Prepare brilliant green water by adding 2 ml 1% **brilliant green dye solution** per 1000 ml sterile distilled water. Let container stand undisturbed for 60 ± 5 min. Incubate loosely capped container, without mixing or pH adjustment, for 24 ± 2 h at 35°C. Continue as in D.1-11, below.

   b. **Non-Instant.** Examine as described for instant nonfat dry milk, except that the 25 g analytical units may not be composited.

4. **Dry whole milk.** Examine as described for instant nonfat dry milk, except that the 25 g analytical units may not be composited.

5. **Casein**

   a. **Lactic casein.** Aseptically weigh 25 g sample into sterile beaker (250 ml) or other appropriate container. Using sterile glass or paper funnel (made with tape to withstand autoclaving), pour 25 g analytical unit gently and slowly over the surface of 225 ml **universal preenrichment broth (UPB)** contained in sterile 500 ml Erlenmeyer flask or other appropriate container. Analytical units (25 g) may be composited. Let container stand undisturbed 60 ± 5 min. Incubate loosely capped container, without mixing or pH adjustment, for 24 ± 2 h at 35°C. Continue as in D.1-11, below.

   b. **Rennet casein.** Aseptically weigh 25 g sample into sterile beaker (250 ml) or other appropriate container. Using sterile glass or paper funnel (made with tape to withstand autoclaving), pour 25 g analytical unit gently and slowly over the surface of 225 ml **lactose broth** contained in sterile 500 ml Erlenmeyer flask or other appropriate container. Analytical units (25 g) may be composited. Let container stand undisturbed 60 ± 5 min. Incubate loosely capped container, without mixing or pH adjustment, for 24 ± 2 h at 35°C. Continue as in D.1-11, below.

   c. **Sodium caseinate.** Aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Add 225 ml sterile lactose broth and mix well. Analytical units may be composited. Let stand 60 min at
room temperature with jar securely capped. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Loosen jar about 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

6. **Soy flour.** Examine as described for rennet casein, except 25 g analytical units (25 g) may not be composited.

7. **Fresh, frozen, or dried products.** Preferably, do not thaw frozen samples before analysis. If frozen sample must be tempered to obtain analytical portion, thaw below 45°C for <15 min with continuous agitation in thermostatically controlled water bath or thaw within 18 h at 2-5°C.
   
a. **Egg-containing products (noodles, egg rolls, macaroni, spaghetti), cheese, dough, prepared salads (ham, egg, chicken, tuna, turkey), fruits, crustaceans (shrimp, crab, crayfish, langostinos, lobster), mushrooms, fish and oysters.** Aseptically weigh 25 g sample into sterile blending container. Add 225 ml sterile lactose broth and blend 2 min. Aseptically transfer homogenized mixture to sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container and let stand 60 ± 5 min at room temperature with jar securely capped. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Mix well and loosen jar cap about 1/4 turn. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

b. **Nuts and nut meats.** Aseptically weigh 25 g sample into a sterile wide mouth Erlenmeyer flask or other appropriate container. Add 225 mL universal preenrichment broth (UPB) and mix well by swirling. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

c. **Nut butter.** Aseptically weigh 25 g sample into a sterile wide mouth Erlenmeyer flask or other appropriate container. Add 225 mL universal preenrichment broth (UPB) and blend 2 min. Aseptically transfer homogenized mixture to sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Mix well and loosen jar cap about 1/4 turn. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

d. **Vegetables.** Aseptically weigh 25 g sample into a sterile wide mouth Erlenmeyer flask or other appropriate container. Add 225 mL universal preenrichment broth (UPB) and mix well by swirling. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

8. **Dried yeast (active and inactive yeast).** Aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Add 225 ml sterile trypticase soy broth (TSB). Mix well to form smooth suspension. Let stand 60 ± 5 min at room temperature with jar securely capped. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2, mixing well before determining
final pH. Loosen jar cap 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

9. **Frosting and topping mixes.** Aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Add 225 ml nutrient broth and mix well. Cap jar securely and let stand 60 ± 5 min at room temperature. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Loosen jar cap about 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

10. **Spices**

   a. **Black pepper, white pepper, celery seed or flakes, chili powder, cumin, paprika, parsley flakes, rosemary, sesame seed, thyme, and vegetable flakes.** Aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Add 225 ml sterile trypticase soy broth (TSB) and mix well. Cap jar securely and let stand 60 ± 5 min at room temperature. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Loosen jar cap about 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

   b. **Onion flakes, onion powder, garlic flakes.** Aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Pre-enrich sample in trypticase soy broth (TSB) with added K$_2$SO$_3$ (5 g K$_2$SO$_3$ per 1000 ml TSB, resulting in final 0.5% K$_2$SO$_3$ concentration). Add K$_2$SO$_3$ to broth before autoclaving 225 ml volumes in 500 ml Erlenmeyer flasks at 121°C for 15 min. After autoclaving, aseptically determine and, if necessary, adjust final volume to 225 ml. Add 225 ml sterile TSB with added K$_2$SO$_3$ to sample and mix well. Continue as in C-10a.

   c. **Allspice, cinnamon, cloves, and oregano.** There are no known methods for neutralizing the toxicity of these 4 spices (powder or non-powder) at this time. Dilute spices beyond their toxic levels to examine them for the presence of *Salmonella*. Examine allspice, cinnamon, and oregano at 1:100 sample/broth ratio, and cloves at 1:1000 sample/broth ratio.

   In more detail, for oregano, cinnamon, and allspice, mix each sample thoroughly; take 25.0 g from each of 15 subsamples for a total of 375.0 g. Thoroughly mix this composite. Then from the 375.0 g composite, measure 37.5 g, and add to 3712.5 ml of sterile pre-enrichment broth (TSB). Shake and mix well for pre-enrichment. For food categories requiring 30 subsamples, create 2 sets of composites, each consisting of 15 subsamples; for those requiring 60 subsamples, create 4 sets of composites, each consisting of 15 subsamples. Then follow the procedures for 15 subsamples described above. For cloves, create a 375.0 g composite from 15 subsamples according to the procedures described above. Then from the 375.0 g composite, measure 3.75 g, and add to 3746.25 ml of
sterile pre-enrichment broth (TSB). Shake and mix well for pre-enrichment. A minimum of 37.5 g sample size is required, 10 tests should be conducted from the same 375.0 g composite. For food categories requiring 30 subsamples, create 2 sets of composites, each consisting of 15 subsamples; for those requiring 60 subsamples, create 4 sets of composites, each consisting of 15 subsamples. Then follow the procedures for 15 subsamples described above.

After the pre-enrichment procedures described above, examine these spices as described in C-10a, above, maintaining recommended sample/broth ratios.

Non-powder forms of allspice, cinnamon, cloves, and oregano (whole, chunks, pieces, leaves). Aseptically weigh 25 g sample into a sterile Whirl-Pak® filter bag or equivalent. Add 225 ml TSB and vigorously shake for 60 sec manually. Transfer the rinsate immediately into a fresh sterile bag. It is crucial to add TSB into the sample immediately before shaking and to transfer the rinsate out immediately after shaking. If the samples are very absorptive of broth (such as oregano leaves), 475 ml of TSB should be added to 25 g of samples. Continue to pre-enrich the rinsate as in C-10a.

11. Candy and candy coating (including chocolate). Aseptically weigh 25 g sample into sterile blending container. Add 225 ml sterile, reconstituted nonfat dry milk and blend 2 min. Aseptically transfer homogenized mixture to sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container and let stand 60 ± 5 min at room temperature with jar securely capped. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Add 0.45 ml 1% aqueous brilliant green dye solution and mix well. Loosen jar caps 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

12. Coconut. Aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Add 225 ml sterile lactose broth, shake well, and let stand 60 ± 5 min at room temperature with jar securely capped. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Add up to 2.25 ml steamed (15 min) Triton X-100. Limit use of these surfactants to minimum quantity needed to initiate foaming. For Triton X-100 this quantity may be as little as 2 or 3 drops. Loosen jar cap about 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

13. Food dyes and food coloring substances. For dyes with pH 6.0 or above (10% aqueous suspension), use method described for dried whole eggs (C.1, above). For laked dyes or dyes with pH below 6.0, aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Add 225 ml tetrathionate broth without brilliant green dye. Mix well and let stand 60 ± 5 min at room temperature with jar securely capped. Using pH meter, adjust pH to 6.8 ± 0.2. Add 2.25 ml 0.1% brilliant
**green dye solution** and mix thoroughly by swirling. Loosen jar cap about 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

14. **Gelatin.** Aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Add 225 ml sterile lactose broth and 5 ml 5% aqueous *papain solution* and mix well. Cap jar securely and incubate at 35°C for 60 ± 5 min. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Loosen jar cap about 1/4 turn and incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

15. **Meats, meat substitutes, meat by-products, animal substances, glandular products, and meals (fish, meat, bone).** Aseptically weigh 25 g sample into sterile blending container. Add 225 ml sterile lactose broth and blend 2 min. Aseptically transfer homogenized mixture to sterile wide-mouth, screw-cap jar (500 ml) or other appropriate container and let stand 60 ± 5 min at room temperature with jar securely capped. If mixture is powder or is ground or comminuted, blending may be omitted. For samples that do not require blending, add lactose broth and mix thoroughly; let stand for 60 ± 5 min at room temperature with jar securely capped.

Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Add up to 2.25 ml steamed (15 min) *Tergitol anionic 7* and mix well. Alternatively, use steamed (15 min) *Triton X-100*. Limit use of these surfactants to minimum quantity needed to initiate foaming. Actual quantity will depend on composition of test material. Surfactants will not be needed in analysis of powdered glandular products. Loosen jar caps 1/4 turn and incubate sample mixtures 24 ± 2 h at 35°C. Continue as in D.1-11, below.

16. **Frog legs.** (This method is used for all domestic and imported frog legs.) Place 15 pairs of frog legs into sterile plastic bag and cover with sterile lactose broth at a 1:9 sample-to-broth (g/ml) ratio (see A, 23-24, above). If single legs are estimated to average 25 g or more, examine only one leg of each of 15 pairs. Place bag in large plastic beaker or other suitable container. Mix well and let stand 60 ± 5 min at room temperature. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Place plastic bag containing the frog legs and lactose broth into plastic beaker or other suitable container. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

17. **Rabbit carcasses.** (This method is used for all domestic and imported rabbit carcasses.) Place rabbit carcass into sterile plastic bag. Place bag in beaker or other suitable container. Add sterile lactose broth at a 1:9 sample-to-broth (g/ml) ratio to cover carcass (see A, 23-24, above). Mix well by swirling and let stand 60 ± 5 min at room temperature. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

18. **Guar gum.** Aseptically weigh 25 g sample into sterile beaker (250 ml) or other appropriate container. Prepare a 1.0% cellulase solution (add 1 g cellulase to 99 ml sterile

15
distilled water). Dispense into 150 ml bottles. (Cellulase solution may be stored at 2-5°C for up to 2 weeks). Add 225 ml sterile lactose broth and 2.25 ml sterile 1% cellulase solution to sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. While vigorously stirring the cellulase/lactose broth with magnetic stirrer, pour 25 g analytical unit quickly through sterile glass funnel into the cellulase/lactose broth. Cap jar securely and let stand 60 ± 5 min at room temperature. Incubate loosely capped container without pH adjustment, for 24 ± 2 h at 35°C. Continue as in D.1-11, below.

19. **Orange juice (pasteurized and unpasteurized), apple cider (pasteurized and unpasteurized), and apple juice (pasteurized).** Aseptically add 25 ml sample to 225 ml universal preenrichment broth (UPB) in a sterile, wide mouth, screw-capped jar (500 ml) or other appropriate container. Swirl the flask contents thoroughly. Cap jar securely and let stand 60 ± 5 min at room temperature. Do not adjust pH. Incubate loosely capped container for 24 ± 2 h at 35°C. Continue as in D.1-11, below (treat as a low microbial load food).

20. **Pig ears and other types of dog chew pieces.** Place 1 piece (or 2-3 pieces if smaller sizes) from each sample unit into sterile plastic bag. Place bag into large beaker or other suitable container. Add sterile lactose broth at a 1:9 sample-to-broth (g/ml) ratio to cover pieces (see A, 23-24, above). Mix well by swirling and let stand 60 ± 5 min at room temperature. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Add either steamed (15 min) Tergitol anionic 7 or steamed (15 min) Triton X-100 up to a 1% concentration. For example, if 225 ml lactose broth is added, the maximum volume of added surfactant is 2.25 ml. Limit use of these surfactants to minimum quantity to initiate foaming. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

21. **Cantaloupes.** Preferably, do not thaw frozen samples before analysis. If frozen sample must be tempered to obtain analytical portion, thaw below 45°C for <15 min with continuous agitation in thermostatically controlled water bath or thaw within 18 h at 2-5°C.

For comminuted or cut fruit, aseptically weigh 25 g sample into sterile blending container. Add 225 ml sterile universal preenrichment broth (UPB) and blend 2 min. Aseptically transfer homogenized mixture to sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container and let stand 60 ± 5 min at room temperature with jar securely capped. Do not adjust pH. Mix well and loosen jar cap about 1/4 turn. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

For whole cantaloupes, do not rinse even if there is visible dirt. Examine the cantaloupes "as is".

Place the cantaloupe into a sterile plastic bag. Add enough UPB broth to allow the cantaloupe to float. The volume of UPB broth may be 1.5 times the weight of the
cantaloupes. For example, cantaloupes weighing 1500 g will probably need a volume of approximately 2250 ml UPB broth to float. Add more broth, if necessary. Place the plastic bag, with cantaloupes and UPB broth, into a 5 liter beaker, or other appropriate container, for support during incubation. Allow the open-end flap of the plastic bag to "fold over" so as to form a secure, but not air-tight, closure during incubation.

Let stand for 60 ± 5 min at room temperature. Do not adjust pH. Incubate slightly opened bag, containing cantaloupe, for 24 ± 2 h at 35°C. Continue as in **D.1-11**, below.

22. **Mangoes.** Preferably, do not thaw frozen samples before analysis. If frozen sample must be tempered to obtain analytical portion, thaw below 45°C for <15 min with continuous agitation in thermostatically controlled water bath or thaw within 18 h at 2-5°C.

For comminuted or cut fruit, aseptically weigh 25 g sample into sterile blending container. Add 225 ml sterile buffered peptone water (BPW) and blend 2 min. Aseptically transfer homogenized mixture to sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container and let stand 60 ± 5 min at room temperature with jar securely capped. Mix well by swirling and loosen jar cap about 1/4 turn. Incubate 24 ± 2 h at 35°C. Continue as in **D.1-11**, below.

For whole mangoes, do not rinse even if there is visible dirt. Examine the mangoes "as is".

Place the mango into a sterile plastic bag. Add enough BPW to allow the mango to float. The volume of BPW may be 1.0 times the weight of the mangoes. For example, mangoes weighing 500 g will probably need a volume of approximately 500 ml BPW broth to float. Add more broth, if necessary. Place the plastic bag, with mangoes and BPW broth, into a 5 liter beaker, or other appropriate container, for support during incubation.

Let stand for 60 ± 5 min at room temperature. Incubate slightly opened bag for 24 ± 2 h at 35°C. Continue as in **D.1-11**, below.

23. **Tomatoes.** For comminuted or cut fruit, aseptically weigh 25 g sample into sterile blending container. Add 225 ml universal preenrichment broth (UPB) and blend 2 min. Aseptically transfer homogenized mixture to sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container and let stand 60 ± 5 min at room temperature with jar securely capped. Do not adjust pH. Mix well and loosen jar cap about 1/4 turn. Incubate 24 ± 2 h at 35°C. Continue as in **D.1-11**, below.

For whole tomatoes, do not rinse even if there is visible dirt. Examine the tomatoes "as is".

Place the tomato into a sterile plastic bag or other suitable container (sterile foil covered beaker can be used). Add enough UPB broth to allow the tomato to float. The volume
of UPB broth may be 1.0 times the weight of the tomato. For example, tomatoes weighing 300 g will probably need a volume of approximately 300 ml UPB broth to float. Add more, if necessary. Place the plastic bag (if used), with tomato and UPB broth, into a sterile beaker (beaker size is dependent on the size of the tomato), or other appropriate container, for support during incubation. Allow the open-end flap of the plastic bag to "fold over" so as to form a secure, but not air-tight, closure during incubation.

Let stand for 60 ± 5 min at room temperature. Do not adjust pH. Incubate slightly opened bag for 24 ± 2 h at 35°C. Continue as in D.1-11, below.

24. **Environmental testing.** Sample environmental surfaces with sterile swabs or sponges. Place the swab/sponge in a sterile Whirl-Pak® bag, or equivalent, that contains enough Dey-Engley (DE) broth to cover the swab/sponge.

Transport swabs/sponges in an insulated transport container with frozen gel packs to keep the samples cold, but not frozen. If samples cannot be processed immediately, refrigerate at 4 ± 2°C. Start sample analysis within 48 ± 2 h of collection.

Add swab/sponge to 225 ml lactose broth in a sterile, wide mouth, screw-capped jar (500 ml) or other appropriate container. Swirl the flask contents thoroughly. Cap jar securely and let stand 60 ± 5 min at room temperature. Mix well by swirling and determine pH with test paper. Adjust pH, if necessary, to 6.8 ± 0.2. Incubate 24 ± 2 h at 35°C. Continue examination as in D.1-11, below.

25. **Alfalfa seeds and mung beans.** Aseptically weigh 25 g alfalfa seeds or mung beans into a sterile 500 mL Erlenmeyer flask. Aseptically add 225 mL lactose broth to the test portion and swirl the Erlenmeyer flask. Cover the mouth of the Erlenmeyer flask with sterile aluminum foil and allow contents to stand at room temperature for 60 ± 5 min. Adjust the pH of the culture to 6.8 ± 0.2, if necessary. Incubate for 24 ± 2h at 35 ± 2°C. Continue as in D.1-11, below (treat as high microbial load food).

26. **Mamey pulp.** If frozen, sample must be tempered to obtain analytical portion. Thaw below 45°C for <15 min with continuous agitation in thermostatically controlled water bath or thaw within 18 h at 2-5°C.

For mamey pulp, suspected to be contaminated with *S. Typhi*, aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container. Add 225 ml sterile universal preenrichment broth (UPB) without ferric ammonium citrate, mix by swirling, and let stand 60 ± 5 min at room temperature with jar securely capped. Do not adjust pH. Mix well and loosen jar cap about 1/4 turn. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below. Treat as a low microbial load food.

For mamey pulp, NOT suspected to be contaminated with *S. Typhi*, aseptically weigh 25 g sample into sterile, wide-mouth, screw-cap jar (500 ml) or other appropriate container.
Add 225 ml sterile universal preenrichment broth (UPB), mix by swirling, and let stand 60 ± 5 min at room temperature with jar securely capped. Do not adjust pH. Mix well and loosen jar cap about 1/4 turn. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

27. **Fresh leafy green vegetables, herbs and sprouts (baby spinach, cabbage, iceberg lettuce, Romaine lettuce, Spring mix, basil, cilantro, dill, curly parsley, culantro, Italian parsley, watercress, alfalfa, mung bean, clover, radish and broccoli sprouts)**. Aseptically weigh 25 g into a sterile wide mouth Erlenmeyer flask or other appropriate container. Add 225 mL universal preenrichment broth (UPB) (for cabbage, adding 225 ml modified buffered peptone water (M192b)) and completely soak contents without any homogenization. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

28. **Animal food (dry cat food, dry dog food, cattle feed, dairy feed, horse feed, poultry feed, and swine feed)**. Aseptically weigh 25 g sample into a sterile Whirl-pak filter bag, or equivalent. Place bag into large container or rack for support during incubation. Add 225 ml sterile buffered peptone water (BPW). Mix well by swirling and brief hand-massage. Incubate 24 ± 2 h at 35°C. Continue as in D.1-11, below.

Animal food can be screened for *Salmonella* using loop-mediated isothermal amplification (LAMP). If screening with LAMP, only samples positive by the LAMP method should be confirmed by culture below (D.1-11).

Protocol: [Screening of *Salmonella* in animal food by LAMP](https://www.fda.gov/downloads/ForConsumers/ForHealthProfessionals/ConsumerUpdates/UCM566961.pdf) (PDF 1.08 MB). Contact: Beilei Ge, FDA-CVM (beilei.ge@fda.hhs.gov).

29. **Spent sprout irrigation water from alfalfa, mung bean, and broccoli varieties**. Aseptically add 375 ml sample to 1,125 ml universal preenrichment broth (UPB) in a sterile Whirl-Pak filter bag, or equivalent. Place bag into large container or rack for support during incubation. Mix well by swirling. Incubate 24 ± 2 h at 35 ± 2°C. Continue examination as in D.1-11, below (treat as a high microbial load food).

### D. Isolation of *Salmonella*

1. Tighten lid and gently shake incubated sample.

   **Guar gum and foods suspected to be contaminated with *S. Typhi***. Transfer 1 ml mixture to 10 ml selenite cystine (SC) broth and another 1 ml mixture to 10 ml tetrathionate (TT) broth. Vortex.

   **All other foods**. Transfer 0.1 ml mixture to 10 ml Rappaport-Vassiliadis (RV) medium and another 1 ml mixture to 10 ml TT. Vortex.

2. Incubate selective enrichment media as follows:
Foods with a high microbial load. Incubate RV medium 24 ± 2 h at 42 ± 0.2°C (circulating, thermostatically-controlled, water bath). Incubate TT broth 24 ± 2 h at 43 ± 0.2°C (circulating, thermostatically-controlled, water bath).

Foods with a low microbial load (except guar gum and foods suspected to be contaminated with *S. Typhi*). Incubate RV medium 24 ± 2 h at 42 ± 0.2°C (circulating, thermostatically controlled, water bath). Incubate TT broth 24 ± 2 h at 35 ± 2.0°C.

Guar gum and foods suspected to be contaminated with *S. Typhi*. Incubate SC and TT broths 24 ± 2 h at 35°C.

3. Mix (vortex, if tube) and streak 3 mm loopful (10 µl) incubated TT broth on bismuth sulfite (BS) agar, xylose lysine desoxycholate (XLD) agar, and Hektoen enteric (HE) agar. Prepare BS plates the day before streaking and store in dark at room temperature until streaked.

4. Repeat with 3 mm loopful (10 µl) of RV medium (for samples of high and low microbial load foods) and of SC broth (for guar gum).

5. Refer to 994.04 in *Official Methods of Analysis* (1) for option of refrigerating incubated sample preenrichments and incubated sample selective enrichments (SC and TT broths only) of low moisture foods. This option allows sample analyses to be initiated as late as Thursday while still avoiding weekend work.

6. Incubate plates 24 ± 2 h at 35°C.

7. Examine plates for presence of colonies that may be *Salmonella*.

**TYPICAL *Salmonella* COLONY MORPHOLOGY**

Pick 2 or more colonies of *Salmonella* from each selective agar plate after 24 ± 2 h incubation. Typical *Salmonella* colonies are as follows:

If typical colonies are present on the BS agar after 24 ± 2 h incubation, then pick 2 or more colonies. Irrespective of whether or not BS agar plates are picked at 24 ± 2 h, reincubate BS agar plates an additional 24 ± 2 h. After 48 ± 2 h incubation, pick 2 or more typical colonies, if present, from the BS agar plates, only if colonies picked from the BS agar plates incubated for 24 ± 2 h give atypical reactions in triple sugar iron agar (TSI) and lysine iron agar (LIA) that result in culture being discarded as not being *Salmonella*. See sections D.9 and D.10, below, for details in interpreting TSI and LIA reactions.

**ATYPICAL *Salmonella* COLONY MORPHOLOGY**

In the absence of typical or suspicious *Salmonella* colonies, search for atypical *Salmonella* colonies as follows:
SUGGESTED CONTROL CULTURES

Please see control cultures below. Analysts must use both typical and atypical *Salmonella* controls for sample analysis, since it may be more difficult to confirm the presence of an atypical strain in a sample without an atypical control strain.

Green fluorescent protein (GFP) control strains which fluoresce under UVA light, more specifically at wavelengths between 360 to 400 nm, have been developed by FDA and have been licensed by FDA to Microbiologics for distribution (https://www.microbiologics.com; 200 Cooper Avenue North St. Cloud, MN 56303, 1-800-599-2847). The following cultures may be purchased from Microbiologics:

*Salmonella* enterica subsp. *enterica* serovar Gaminara Sal57 / FDA SAL5695 (typical, lactose negative, H₂S positive), Catalog No. 01278UV.

*Salmonella* enterica subsp. *enterica* serovar Senftenberg Sal59 / FDA Sal5697 (atypical, lactose negative, H₂S negative), Catalog No. 01226UV.

*Salmonella* enterica subsp. *enterica* serovar Mbandaka Sal58 / FDA SAL5696 (atypical, sucrose positive, H₂S positive), Catalog No. 01230UV.

Non-GFP fluorescent *Salmonella* control strains may also be used for sample analysis. In addition to the positive control cultures (typical *Salmonella*; lactose negative, H₂S positive), 3 additional *Salmonella* cultures are recommended to assist in the selection of atypical *Salmonella* colony morphology on selective agars. These cultures are a lactose-positive, H₂S -positive *S. diarizonae* (47:i:z53:z57, ATCC 12325) and a lactose-negative, H₂S -negative *S. Abortusequi* (ATCC 9842); OR a lactose-positive, H₂S -negative *S. diarizonae* (60:r:e,n,x,z15, ATCC 29934). These cultures may be obtained from the American Type Culture Collection (http://www.atcc.org/), 10801 University Boulevard, Manassas, VA 20110-2209.

a. **Hektoen enteric (HE) agar.** Blue-green to blue colonies with or without black centers. Many cultures of *Salmonella* may produce colonies with large, glossy black centers or may appear as almost completely black colonies.

b. **Xylose lysine desoxycholate (XLD) agar.** Pink colonies with or without black centers. Many cultures of *Salmonella* may produce colonies with large, glossy black centers or may appear as almost completely black colonies.

c. **Bismuth sulfite (BS) agar.** Brown, gray, or black colonies; sometimes they have a metallic sheen. Surrounding medium is usually brown at first, but may turn black in time with increased incubation, producing the so-called halo effect.

d. **HE and XLD agars.** Atypically a few *Salmonella* cultures produce yellow colonies with or without black centers on HE and XLD agars. In the absence of typical *Salmonella* colonies on HE or XLD agars after 24 ± 2 h incubation, then pick 2 or more atypical *Salmonella* colonies.
e. **BS agar.** Atypically some strains produce green colonies with little or no
darkening of the surrounding medium. If typical or suspicious colonies are not
present on BS agar after 24 ± 2 h, then do not pick any colonies but reincubate an
additional 24 ± 2 h. If typical or suspicious colonies are not present after 48 ± 2 h
incubation, then pick 2 or more atypical colonies.

8. Lightly touch the very center of the colony to be picked with sterile inoculating needle
and inoculate TSI slant by streaking slant and stabbing butt. Without flaming, inoculate
LIA slant by stabbing butt twice and then streaking slant. Since lysine decarboxylation
reaction is strictly anaerobic, the LIA slants must have deep butt (4 cm). Store picked
selective agar plates at 5-8°C.

9. Incubate TSI and LIA slants at 35°C for 24 ± 2 h. Cap tubes loosely to maintain aerobic
conditions while incubating slants to prevent excessive H2S production. *Salmonella* in
culture typically produces alkaline (red) slant and acid (yellow) butt, with or without
production of H2S (blackening of agar) in TSI. In LIA, *Salmonella* typically produces
alkaline (purple) reaction in butt of tube. Consider only distinct yellow in butt of tube as
acidic (negative) reaction. Do not eliminate cultures that produce discoloration in butt of
tube solely on this basis. Most *Salmonella* cultures produce H2S in LIA. Some
non- *Salmonella* cultures produce a brick-red reaction in LIA slants.

10. Cultures that give an alkaline butt in LIA should be retained as
potential *Salmonella* isolates and submitted for biochemical and serological tests if the
TSI slant is either alkaline over acid or acid over acid. Cultures that give an acid butt in
LIA and an alkaline slant and acid butt in TSI should also be considered
potential *Salmonella* isolates and should be submitted for biochemical and serological
tests. Cultures that give an alkaline butt in LIA, but where there is no change in the TSI
for both the slant and the butt should be discarded. Cultures that give an acid butt in LIA
and an acid slant and acid butt in TSI may be discarded as not being *Salmonella*. Test
retained, presumed-positive TSI cultures as directed in **D.11**, below, to determine if they
are *Salmonella* species, including *S. arizonae*. If TSI cultures fail to give typical reactions
for *Salmonella* (alkaline slant and acid butt) pick additional suspicious colonies from
selective medium plate not giving presumed-positive culture and inoculate TSI and LIA
slants as described in **D.8**, above.

11. Apply biochemical and serological identification tests to:

   a. Three presumptive TSI cultures recovered from set of plates streaked from RV
   medium (or SC broth for guar gum), if present, and 3 presumptive TSI agar
   cultures recovered from plates streaked from TT broth, if present.

   b. If 3 presumptive-positive TSI cultures are not isolated from one set of agar plates,
   test other presumptive-positive TSI agar cultures, if isolated, by biochemical and
   serological tests. Examine a minimum of 6 TSI cultures for each 25 g analytical
   unit or each 375 g composite.
E. Identification of *Salmonella*

1. **Mixed cultures.** Streak TSI agar cultures that appear to be mixed on [MacConkey agar](#), [HE agar](#), or [XLD agar](#). Incubate plates 24 ± 2 h at 35°C. Examine plates for presence of colonies suspected to be *Salmonella*.
   
a. **MacConkey agar.** Typical colonies appear transparent and colorless, sometimes with dark center. Colonies of *Salmonella* will clear areas of precipitated bile caused by other organisms sometimes present.
   
b. **Hektoen enteric (HE) agar.** See [D.7a](#), above.
   
c. **Xylose lysine desoxycholate (XLD) agar.** See [D.7b](#), above. Transfer at least 2 colonies suspected to be *Salmonella* to TSI and LIA slants as described in [D.7](#), above, and continue as in [D.9](#), above.

2. **Pure cultures**

   a. **Urease test (conventional).** With sterile needle, inoculate growth from each presumed-positive TSI slant culture into tubes of [urea broth](#). Since occasional, uninoculated tubes of urea broth turn purple-red (positive test) on standing, include uninoculated tube of this broth as control. Incubate 24 ± 2 h at 35°C.
   
b. **Optional urease test (rapid).** Transfer two 3-mm loopfuls of growth from each presumed-positive TSI slant culture into tubes of [rapid urea broth](#). Incubate 2 h in 37 ± 0.5°C water bath. Discard all cultures giving positive test. Retain for further study all cultures that give negative test (no change in color of medium).

3. **Serological polyvalent flagellar (H) test**

   a. Perform the polyvalent flagellar (H) test at this point, or later, as described in [E.4](#), below. Inoculate growth from each urease-negative TSI agar slant into either 1) [BHI broth](#) and incubate 4-6 h at 35°C until visible growth occurs (to test on same day); or 2) [trypsicase soy-tryptose broth](#) and incubate 24 ± 2 h at 35°C (to test on following day). Add 2.5 ml formalinized physiological saline solution to 5 ml of either broth culture.
   
b. **BD DIFCO™ Procedure.** Select 2 formalinized broth cultures and test with *Salmonella* polyvalent flagellar (H) antisera per manufacturer's instructions. Place 0.5 ml of appropriately diluted *Salmonella* polyvalent flagellar (H) antiserum in 10 × 75 mm or 13 × 100 mm serological test tube. Add 0.5 ml antigen to be tested. Prepare saline control by mixing 0.5 ml physiological saline solution with 0.5 ml formalinized antigen. Incubate mixtures in 48-50°C water bath. Observe at 15 min intervals and read final results in 1 h.

   **Positive** — agglutination in test mixture and no agglutination in control.

   **Negative** — no agglutination in test mixture and no agglutination in control.
**Nonspecific** — agglutination in both test mixture and control. Test the cultures giving such results with Spicer-Edwards antisera.

c. Statens Serum Institute Procedure. Perform the polyvalent flagellar (H) test using Statens Serum Institute *Salmonella* polyvalent flagellar (H) antisera. The *Salmonella* is grown over night on a non-selective agar medium. Swarm agar is the best suited medium for growing cultures for H typing, but H antigens can be serotyped from a non-selective agar medium if the H antigens are well expressed. Add a small drop of antiserum (approx. 20 µL) on a glass slide or plastic petri dish (15 × 100 mm). Transfer culture using an inoculating loop from several colonies to the drop of antiserum and mix well. The amount of culture should be sufficient to give a distinct milky turbidity. Tilt the slide or petri dish for 5-10 seconds. A positive reaction is seen as visible agglutination, whereas a negative reaction is seen as homogeneous milky turbidity. A late or weak agglutination should be considered negative. Physiological saline (0.85%, pH 7.4) is used as a negative control and must be negative.

4. **Spicer-Edwards serological test.** Use this test as an alternative to the polyvalent flagellar (H) test. It may also be used with cultures giving nonspecific agglutination in polyvalent flagellar (H) test. Perform Spicer-Edwards flagellar (H) antisera test as described in E.3b, above. Perform additional biochemical tests (E.5a-c, below) on cultures giving positive flagellar test results. If both formalinized broth cultures are negative, perform serological tests on 4 additional broth cultures (E.3a, above). If possible, obtain 2 positive cultures for additional biochemical testing (E.5a-c, below). If all urease-negative TSI cultures from sample give negative serological flagellar (H) test results, perform additional biochemical tests E. 5a-c, below).

5. **Testing of urease-negative cultures**

   a. **Lysine decarboxylase broth.** If LIA test was satisfactory, it need not be repeated. Use lysine decarboxylase broth for final determination of lysine decarboxylase if culture gives doubtful LIA reaction. Inoculate broth with small amount of growth from TSI slant suspicious for *Salmonella*. Replace cap tightly and incubate 48 ± 2 h at 35°C but examine at 24 h intervals. *Salmonella* species cause alkaline reaction indicated by purple color throughout medium. Negative test is indicated by yellow color throughout medium. If medium appears discolored (neither purple nor yellow) add a few drops of 0.2% bromcresol purple dye and re-read tube reactions.

   b. **Phenol red dulcitol broth or purple broth base with 0.5% dulcitol.** Inoculate broth with small amount of growth from TSI culture. Replace cap loosely and incubate 48 ± 2 h at 35°C, but examine after 24 h. Most *Salmonella* species give positive test, indicated by gas formation in inner fermentation vial and acid pH (yellow) of medium. Production of acid should be interpreted as a positive reaction. Negative test is indicated by no gas formation in inner fermentation vial.
and red (with phenol red as indicator) or purple (with bromcresol purple as indicator) color throughout medium.

c. **Tryptone (or tryptophane) broth.** Inoculate broth with small growth from TSI agar culture. Incubate 24 ± 2 h at 35°C and proceed as follows:

1. **Potassium cyanide (KCN) broth.** Transfer 3 mm loopful of 24 h tryptophane broth culture to KCN broth. Heat rim of tube so that good seal is formed when tube is stoppered with wax-coated cork. Incubate 48 ± 2 h at 35°C but examine after 24 h. Interpret growth (indicated by turbidity) as positive. Most *Salmonella* species do not grow in this medium, as indicated by lack of turbidity.

2. **Malonate broth.** Transfer 3 mm loopful of 24 h tryptone broth culture to malonate broth. Since occasional uninoculated tubes of malonate broth turn blue (positive test) on standing, include uninoculated tube of this broth as control. Incubate 48 ± 2 h at 35°C, but examine after 24 h. Most *Salmonella* species cultures give negative test (green or unchanged color) in this broth.

3. **Indole test.** Transfer 5 ml of 24 h tryptophane broth culture to empty test tube. Add 0.2-0.3 ml Kovacs' reagent. Most *Salmonella* cultures give negative test (lack of deep red color at surface of broth). Record intermediate shades of orange and pink as ±.

4. **Serological flagellar (H) tests for Salmonella.** If either polyvalent flagellar (H) test (E-3, above) or the Spicer-Edwards flagellar (H) test tube test (E-4, above) has not already been performed, either test may be performed here.

5. Discard as not *Salmonella* any culture that shows either positive indole test and negative serological flagellar (H) test, or positive KCN test and negative lysine decarboxylase test.

6. **Serological somatic (O) tests for Salmonella.** (Pre-test all antisera to *Salmonella* with known cultures.)

   a. **Polyvalent somatic (O) test.** Using wax pencil, mark off 2 sections about 1 × 2 cm each on inside of glass or plastic petri dish (15 × 100 mm). Commercially available sectioned slides may be used. Emulsify 3 mm loopful of culture from 24-48 h TSI slant or, preferably, tryptose blood agar base (without blood) with 2 ml 0.85% saline. Add 1 drop of culture suspension to upper portion of each rectangular crayon-marked section. Add 1 drop of saline solution to lower part of one section only. Add 1 drop of *Salmonella* polyvalent somatic (O) antiserum to other section only. With clean sterile transfer loop or needle, mix culture suspension with saline solution for one section and repeat for other section containing antiserum. Tilt mixtures in back-and-forth motion for 1 min and
observe against dark background in good illumination. Consider any degree of agglutination a positive reaction. Classify polyvalent somatic (O) test results as follows:

**Positive** — agglutination in test mixture; no agglutination in saline control.

**Negative** — no agglutination in test mixture; no agglutination in saline control.

**Nonspecific** — agglutination in test and in control mixtures. Perform further biochemical and serological tests as described in Edwards and Ewing's *Identification of Enterobacteriaceae* (2).

b. **Somatic (O) group tests.** Test as in E.6a, above, using individual group somatic (O) antisera including Vi, if available, in place of *Salmonella* polyvalent somatic (O) antiserum. For special treatment of cultures giving positive Vi agglutination reaction, refer to sec. 967.28B in *Official Methods of Analysis* (1). Record cultures that give positive agglutination with individual somatic (O) antiserum as positive for that group. Record cultures that do not react with individual somatic (O) antiserum as negative for that group.

7. **Additional biochemical tests.** Classify as *Salmonella* those cultures which exhibit typical *Salmonella* reactions for tests 1-11, shown in Table 1. If one TSI culture from 25 g analytical unit is classified as *Salmonella*, further testing of other TSI cultures from the same 25 g analytical unit is unnecessary. Cultures that contain demonstrable *Salmonella* antigens as shown by positive *Salmonella* flagellar (H) test but do not have biochemical characteristics of *Salmonella* should be purified (E.1, above) and retested, beginning with E.2, above.

Perform the following additional tests on cultures that do not give typical *Salmonella* reactions for tests 1-11 in Table 1 and that consequently do not classify as *Salmonella*.

a. **Phenol red lactose broth or purple lactose broth.**

1. Inoculate broth with small amount of growth from unclassified 24-48 h TSI slant. Incubate 48 ± 2 h at 35°C, but examine after 24 h.

   **Positive** — acid production (yellow) and gas production in inner fermentation vial. Consider production of acid only as positive reaction. Most cultures of *Salmonella* give negative test result, indicated by no gas formation in inner fermentation vial and red (with phenol red as indicator) or purple (with brom cresol purple as indicator) throughout medium.

2. Discard as not *Salmonella*, cultures that give positive lactose tests, except cultures that give acid slants in TSI and positive reactions in LIA, or cultures that give positive malonate broth reactions. Perform further tests on these cultures to determine if they are *S. arizonae*. 

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b. **Phenol red sucrose broth** or **purple sucrose broth**. Follow procedure described in E.7a-1, above. Discard as not *Salmonella*, cultures that give positive sucrose tests, except those that give acid slants in TSI and positive reactions in LIA.

c. **MR-VP broth**. Inoculate medium with small amount of growth from each unclassified TSI slant suspected to contain *Salmonella*. Incubate 48 ± 2 h at 35°C.

   1. Perform Voges-Proskauer (VP) test at room temperature as follows: Transfer 1 ml 48 h culture to test tube and incubate remainder of MR-VP broth an additional 48 h at 35°C. Add 0.6 ml α-naphthol and shake well. Add 0.2 ml 40% KOH solution and shake. To intensify and speed reaction, add a few crystals of creatine. Read results after 4 h; development of pink-to-ruby red color throughout medium is positive test. Most cultures of *Salmonella* are VP-negative, indicated by absence of development of pink-to-red color throughout broth.

   2. Perform methyl red test as follows: To 5 ml of 96 h MR-VP broth, add 5-6 drops of methyl red indicator. Read results immediately. Most *Salmonella* cultures give positive test, indicated by diffuse red color in medium. A distinct yellow color is negative test. Discard, as not *Salmonella*, cultures that give positive KCN and VP tests and negative methyl red test.

d. **Simmons citrate agar**. Inoculate this agar, using needle containing growth from unclassified TSI agar slant. Inoculate by streaking slant and stabbing butt. Incubate 96 ± 2 h at 35°C. Read results as follows:  

   **Positive** — presence of growth, usually accompanied by color change from green to blue. Most cultures of *Salmonella* are citrate-positive.

   **Negative** — no growth or very little growth and no color change.

e. **Classification of cultures**. Classify, as *Salmonella*, cultures that have reaction patterns of Table 1. Discard, as not *Salmonella*, cultures that give results listed in any subdivision of Table 2. Perform additional tests described in *Edwards and Ewing's Identification of Enterobacteriaceae* (2) to classify any culture that is not clearly identified as *Salmonella* by classification scheme in Table 1 or not eliminated as not being *Salmonella* by test reactions in Table 2. If neither of 2 TSI cultures carried through biochemical tests confirms the isolate as *Salmonella*, perform biochemical tests, beginning with E.5, on remaining urease-negative TSI cultures from same 25 g analytical unit.

8. **Additional Resources:**  
   
   **Salmonella Flipbook**, a pictorial general guide to aid analysts in the detection and identification of *Salmonella* growing on the plating media and screening tubes used in the BAM Chapter 5 *Salmonella* method. (Prepared By: Matthew J. Forstner, Laboratory Services, Minnesota Department of Agriculture).
Table 1. Biochemical and serological reactions of *Salmonella*

<table>
<thead>
<tr>
<th>#</th>
<th>Test or substrate</th>
<th>Positive Result</th>
<th>Negative Result</th>
<th>Salmonella species reaction(a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Glucose (TSI)</td>
<td>yellow butt</td>
<td>red butt</td>
<td>+</td>
</tr>
<tr>
<td>2.</td>
<td>Lysine decarboxylase (LIA)</td>
<td>purple butt</td>
<td>yellow butt</td>
<td>+</td>
</tr>
<tr>
<td>3.</td>
<td>H₂S (TSI and LIA)</td>
<td>blackening</td>
<td>no blackening</td>
<td>+</td>
</tr>
<tr>
<td>4.</td>
<td>Urease</td>
<td>purple-red color</td>
<td>no color change</td>
<td>−</td>
</tr>
<tr>
<td>5.</td>
<td>Lysine decarboxylase broth</td>
<td>purple color</td>
<td>yellow color</td>
<td>+</td>
</tr>
<tr>
<td>6.</td>
<td>Phenol red dulcitol broth</td>
<td>yellow color and/or gas</td>
<td>no gas; no color change</td>
<td>+(b)</td>
</tr>
<tr>
<td>7.</td>
<td>KCN broth</td>
<td>growth</td>
<td>no growth</td>
<td>−</td>
</tr>
<tr>
<td>8.</td>
<td>Malonate broth</td>
<td>blue color</td>
<td>no color change</td>
<td>−(c)</td>
</tr>
<tr>
<td>9.</td>
<td>Indole test</td>
<td>red color at surface</td>
<td>yellow color at surface</td>
<td>−</td>
</tr>
<tr>
<td>10.</td>
<td>Polyvalent flagellar test</td>
<td>agglutination</td>
<td>no agglutination</td>
<td>+</td>
</tr>
<tr>
<td>11.</td>
<td>Polyvalent somatic test</td>
<td>agglutination</td>
<td>no agglutination</td>
<td>+</td>
</tr>
<tr>
<td>12.</td>
<td>Phenol red lactose broth</td>
<td>yellow color and/or gas</td>
<td>no gas; no color change</td>
<td>−(c)</td>
</tr>
<tr>
<td>13.</td>
<td>Phenol red sucrose broth</td>
<td>yellow color and/or gas</td>
<td>no gas; no color change</td>
<td>−</td>
</tr>
<tr>
<td>14.</td>
<td>Voges-Proskauer test</td>
<td>pink-to-red color</td>
<td>no color change</td>
<td>−</td>
</tr>
<tr>
<td>15.</td>
<td>Methyl red test</td>
<td>diffuse red color</td>
<td>diffuse yellow color</td>
<td>+</td>
</tr>
<tr>
<td>16.</td>
<td>Simmons citrate</td>
<td>growth; blue color</td>
<td>no growth; no color change</td>
<td>v</td>
</tr>
</tbody>
</table>

a: +: 90% or more positive in 1 or 2 days; −: 90% or more negative in 1 or 2 days; v: variable.
b: Majority of *S. arizonae* cultures are negative.
c: Majority of *S. arizonae* cultures are positive.
Table 2. Criteria for discarding non-Salmonella cultures

<table>
<thead>
<tr>
<th>#</th>
<th>Test or substrate</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Urease</td>
<td>positive (purple-red color)</td>
</tr>
<tr>
<td>2.</td>
<td>Indole test and Polyvalent flagellar (H) test; or Indole test and Spicer-Edwards flagellar test</td>
<td>positive (red color at surface) negative (no agglutination) positive (red color at surface) negative (no agglutination)</td>
</tr>
<tr>
<td>3.</td>
<td>Lysine decarboxylase and KCN broth</td>
<td>negative (yellow color) positive (growth)</td>
</tr>
<tr>
<td>4.</td>
<td>Phenol red lactose broth</td>
<td>positive (yellow color and/or gas)^(a),(b)</td>
</tr>
<tr>
<td>5.</td>
<td>Phenol red sucrose broth</td>
<td>positive (yellow color and/or gas)^(b)</td>
</tr>
<tr>
<td>6.</td>
<td>KCN broth, Voges-Proskauer test, and Methyl red test</td>
<td>positive (growth) positive (pink-to-red color) negative (diffuse yellow color)</td>
</tr>
</tbody>
</table>

a Test malonate broth positive cultures further to determine if they are *S. arizonae*.
b Do not discard positive broth cultures if corresponding LIA cultures give typical *Salmonella* reactions; test further to determine if they are *Salmonella* species.


   a. **Serological confirmation:** perform the *Salmonella* serological somatic (O) test (E.6, above) and the *Salmonella* serological flagellar (H) test (E.3, above) or the Spicer-Edwards flagellar (H) test (E.4, above). Serological confirmation must always be paired with biochemical confirmation.

   b. **Biochemical confirmation:** Instead of the conventional biochemical tube system above, any of 5 commercial biochemical systems (API 20E, Enterotube II, Enterobacteriaceae II (AOAC OMA 978.24), MICRO-ID (AOAC OMA 989.12), or Vitek 2 GN (AOAC OMA 2011.17) may be used. Commercial biochemical kits should not be used as a substitute for serological tests (I).

   c. **Real-time PCR confirmation test:** Confirmation of *Salmonella* isolates by real-time qPCR (PDF, 1.3 MB).

   d. **Loop-mediated isothermal amplification (LAMP) confirmation test:** Confirmation of *Salmonella* isolates by LAMP (PDF, 1.08 MB). Contact: Beilei Ge, FDA-CVM (beilei.ge@fda.hhs.gov).
e. **ANSR® Salmonella confirmation test (AOAC OMA method 2013.14):** using isothermal nucleic acid amplification assay based on the nicking enzyme amplification reaction (NEAR) technology for the identification and confirmation of *Salmonella*.

g. **Bruker MALDI Biotyper Method (AOAC OMA method 2017.09):** using matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometry (MS) for the identification and confirmation of bacteria.

h. **Other instrumental confirmation methods validated per FDA’s Microbiological Methods Validation Guidelines, AOAC’s Appendix J, or ISO 16140-6:2019** are acceptable if approved for use by FDA’s Microbiology Methods Validation Subcommittee.

i. **Classification of cultures:**
   
   1. Report as *Salmonella* those cultures classified as *Salmonella* by a and b, or by one of c, d, e, f or h methods.
   
   2. Discard cultures not confirmed as *Salmonella* by a, and one of b, c, d, e, f or h methods.
   
   3. For cultures having conflicted confirmation by above methods, classify according to additional tests specified in E, 2-7, above, or additional tests as specified by Ewing (2), or send to reference typing laboratory for definitive serotyping and identification.

10. **Treatment of cultures giving negative flagellar (H) test.** If biochemical reactions of certain flagellar (H)-negative culture strongly suggest that it is *Salmonella*, the negative flagellar agglutination may be the result of nonmotile organisms or insufficient development of flagellar antigen. Proceed as follows: Inoculate motility test medium in petri dish, using small amount of growth from TSI slant. Inoculate by stabbing medium once about 10 mm from edge of plate to depth of 2-3 mm. Do not stab to bottom of plate or inoculate any other portion. Incubate 24 h at 35°C. If organisms have migrated 40 mm or more, retest as follows: Transfer 3 mm loopful of growth that migrated farthest to Trypticase Soy-Tryptose Broth. Repeat either polyvalent flagellar (H) (E.3, above) or Spicer-Edwards (E.4, above) serological tests. If cultures are not motile after the first 24 h, incubate an additional 24 h at 35°C; if still not motile, incubate up to 5 days at 25°C. Classify culture as nonmotile if above tests are still negative. If flagellar (H)-negative culture is suspected of being a species of *Salmonella* on the basis of its biochemical reactions, FDA laboratories should submit the culture to
for further identification and/or serotyping. Laboratories other than FDA should make arrangements with a reference laboratory for the serotyping of *Salmonella* cultures.

11. **Submission of cultures for serotyping.** Submit 1 isolate of each somatic group recovered from each analytical unit, unless otherwise instructed. Submit cultures on BHI agar slants in screw-cap tubes (13 × 100 mm or 16 × 125 mm) with caps secured tightly. Label each tube with sample number, subsample (analytical unit) number, and code, if applicable. Isolate(s) should be submitted to your local serotyping group for WGS and SeqSero analysis.

If isolate(s) require additional serotyping, submit culture(s) on BHI agar slants in screw-cap tubes with caps secured tightly. Label each tube with sample number, subsample (analytical unit) number, and code, if applicable. Place cultures in culture container with official FDA seal. Submit memo or cover letter for each sample number to expedite reporting of results. Prepare cultures for shipment according to requirements for shipment of etiological agents (3). Label secondary shipping container according to ref. 4. Send container by most rapid mail service available. Maintain duplicate cultures of those submitted for serotyping only on those samples under consideration for legal action.

Microbiology Field laboratories should send the isolates to Denver Laboratory for traditional serotyping:

Denver Laboratory
Food and Drug Administration
1 Denver Federal Center
6th Avenue & Kipling Street
Building 20, Entrance W- 10
Denver, CO 80225-0087
Attention: Shauna Madson
Tel # 303-236-9631
Fax # 303-236-9675
References


