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APPENDICES

APPENDIX I

Media and Identification Procedures.

Media

1. Trypticase soy agar (Mearck, Germany)

Peptone from casein	15.0	g
Peptone from soymeal	5.0	g
Sodium chloride	5.0	g
Agar	15.0	g
Distilled water	1,000.0	ml

pH: 7.3 +/- 0.2 at 25°C

Media preparation:

All of ingredients were dissolved in distilled water, heat to boiling and then sterilized by autoclaving at 121°C, 15 pounds/inch² pressure, for 15 minutes. The sterile medium was cooled to 45°C to 50°C, and dispensed into sterile plates.

2. Swarm agar

Bacto beef extract	3.0	g
Bacto peptone	5.0	g
Bacto agar	5.0	g
Natriumdesoxycholol (C ₂₄ H ₃₉ NaO ₄)	0.3	g
Distilled water	1,000.0	g

pH: 7.3 +/- 0.2 at 25°C

Media preparation:

All of ingredients were dissolved in distilled water, heat to boiling and then sterilized by autoclaving at 121°C, 15 pounds/inch² pressure, for 15 minutes. The sterile medium was cooled to 45°C to 50°C, and dispensed into sterile plates.

Identification procedures

1. Biochemical tests

1.1 Triple sugar iron agar

Purpose: Triple sugar iron (TSI) agar is a screening medium used to identify gram-negative bacilli based on ability to ferment the carbohydrates glucose, sucrose, and lactose to produce H₂S gas.

Principle and interpretation: TSI agar contain protein, NaCl, Lactose, sucrose, dextrose, a sulfur source, an H₂S indicator, a pH indicator, and agar. The medium includes ten times as much lactose and sucrose as glucose. Bacteria that ferment glucose produce a variety of acids, turning the colour of the medium from red to yellow. Larger amounts of acid are produced in the butt of the tube (fermentation) than in the slant of the tube (respiration). Organisms growing on TSI also form alkaline products from the oxidative decarboxylation of peptone. These alkaline products neutralize the small amounts of acids present in the slant but are unable to neutralize the large amounts of acid present in the butt. Thus, the appearance of an alkaline (red) slant and an acid (yellow) butt after 24 hours incubation indicates that the organisms is a glucose fermenter but is unable to ferment lactose and sucrose.

Bacteria that ferment lactose or sucrose (or both), in addition to glucose, reduce such large amounts of acid that the oxidative deamination of protein that may occur in the slant does not yield enough alkaline products to cause a reversion of pH in that region. Thus, these bacteria produce an acid slant and acid butt. It is impossible to determine from the TSI reaction whether both lactose and sucrose are being fermented or only one of these

carbohydrates is being fermented; individual carbohydrate fermentation tests are required to make this assessment.

Gas production (CO_2 and hydrogen) is detected by the presence of cracks or bubbles in the medium. These are formed when the accumulated gas escapes.

H_2S gas is produced as a result of the reduction of thiosulfate. H_2S is a colourless gas and can be detected only in the presence of an indicator, in this case ferric ammonium sulfate. H_2S combines with the ferric ions of ferric ammonium sulfate to produce the insoluble black precipitate ferrous sulfide. Reduction of thiosulfate proceeds only in an acid environment, and blackening usually occurs in the butt of the tube. Although the black precipitate may frequently obscure the colour of the butt, it can be assumed that the organism is a glucose fermenter because of the requirement for an acid environment. The reactions can be summarized as follows:

Alkaline slant/acid butt: glucose only fermented

Acid slant/acid butt: glucose and sucrose fermented or glucose and lactose fermented or glucose, lactose, and sucrose fermented

Bubbles or cracks present: gas produced

Black precipitate present: H_2S gas produced

Ingredients and preparation: Mix the ingredients, heat to boiling, dispense into tubes, and sterilize at 121°C for 15 minutes, and allow tubes of medium to cool in a slanted position.

Pancreatic digest of casein, USP	10 g
Peptic digest of animal tissue, USP	10 g
NaCl	1 g
Lactose	10 g
Sucrose	10 g
D-Glucose	1 g

Ferric ammonium sulfate	0.2 g
Sodium thiosulfate	0.2 g
Phenol red	25 g
Agar	13 g
Distilled water	1 L

Final pH 7.3-7.4

Procedure: Inoculate test cultures to TSI agar by first touching a sterile bacteriologic needle to a colony and then stabbing the needle into the deep agar region of the medium. Hence withdrawing the needle, move it from side to side over the surface of the medium. Incubate cultures at 37°C for 18 to 24 hours. Examine cultures for colour of the slant, butt, gas cracks, and blackening caused by H₂S.

1.2 Indole test

Purpose: Indole broth is used for distinguishing between bacteria based on ability to produce indole from tryptophan.

Principle and interpretation: Indole broth contains tryptophan-rich peptone and NaCl. The tryptophan present in peptone is oxidized by certain bacteria to indole, skatole, and indoleacetic acid. The intracellular enzymes that are responsible for metabolizing tryptophan to these compounds are collectively termed tryptophanase. Indole is detected in broth cultures of bacteria with an alcoholic *p*-dimethylaminobenzaldehyde reagent. Indole reacts with the aldehyde to give a red product in the alcoholic layer of the broth-reagent mixture.

Two reagents were used to detect indole: Kovac's and Ehrlich. Ehrlich reagent is believed to be more sensitive than Kovac's and is recommended for detection of indole production by anaerobic bacteria and nonfermentative gram negative organisms. Kovac's

reagent was used initially to classify members of the family *Enterobacteriaceae* and should be used with these organisms.

Ingredients and preparation: Mix the ingredients, heat to boiling, dispense into tubes, and sterilize at 121°C for 15 minutes.

Indole broth:

Pancreatic digest of casein, USP	20 g
NaCl	5 g
Distilled water	1 L

Final pH 7.2

Reagents:

Kovac's indole reagent. Dissolve the aldehyde in the alcohol and slowly add acid to the mixture.

Alcohol, amyl or isoamyl	150 ml
p-Dimethylaminobenzaldehyde	10 g
Hydrochloric acid, concentrated	50 ml

Procedure: Inoculate the test organism into indole broth, incubate at 35°C for 18 to 24 hours, and test as follows.

Indole test: Add 2 to 3 drops of Kovac's reagent directly to the broth culture, shake gently, and observe for development of a red colour in the upper alcohol layer.

1.3 Urea agar

Purpose: Urease agar are used for distinguishing between species of aerobic bacteria based on ability to hydrolyze urea.

Principle and interpretation: A variety of media are used to test for ability to hydrolyze urea. The hydrolysis of urea by urease to ammonia is accompanied by a rise in

pH of the medium and a concomitant change in the color of the indicator from yellow to pink-red.

Ingredients and preparation: Mix urea basal ingredients, sterilize by filtration, and add sterile agar solution (50°C). Mix and dispense into tubes, and allow tubes of medium to cool in a slanted position.

Urea base:

Pancreatic digest of gelatin, USP	1 g
NaCl	5 g
Monopotassium phosphate	2 g
D-Glucose	1 g
Urea	20 g
Phenol red	12 mg
Distilled water	100 ml

Final pH 6.8

Agar solution:

Agar	15 g
Distilled water	900 ml

Urea agar:

Urea base	100 ml
Agar solution	900 ml

Procedure: Inoculate the organism to the urea agar, incubate for 24 to 48 hours at 35°C, and observe for a red color change in the medium.

1.4 Citrate agar, Simmons

Purpose: Simmons citrate agar is used to distinguish gram-negative bacteria based on their ability to utilize as a sole source of carbon.

Principle and interpretation: Several theories have been proposed to explain the mechanism of citrate agar. Only one is presented here. Organisms that metabolize citrate as a sole source of carbon cleave citrate to oxaloacetate and acetate via the citrate enzyme. Another enzyme, oxaloacetate decarboxylase, then converts oxaloacetate to pyruvate and CO₂. CO₂ combines with sodium and water to form Na₂CO₃, an alkaline compound. As a result, the pH of the medium rises and the indicator (bromthymol blue) changes from green to Prussian blue. Presence of the blue color constitutes a positive finding for citrate utilization.

Ingredients and Preparation: Mix the following ingredients, heat to boiling, dispense into test tubes, and sterilize at 121°C for 15 minutes. Cool each tube of medium in a slanted position.

Sodium citrate	2 g
NaCl	5 g
MgSO ₄	0.2 g
Ammonium dihydrogen phosphate	1 g
Dipotassium phosphate	1 g
Bromthymol blue	80 mg
Agar	15 g
Distilled water	1 L

Final pH 6.9

Procedure: Lightly inoculate the test organism to the surface of citrate medium, incubate at 35°C for 24 to 48 hours, and observe for a Prussian blue color change.

1.5 Malonate broth

Purpose: Malonate broth is used for differentiation of members of the family *Enterobacteriaceae*, especially *Salmonella* species.

Principle and interpretation: Malonate broth tests for utilization of sodium malonate as a sole source of carbon. The medium contains buffer, pH indicator, sodium malonate, required salts, and a small amount of yeast extract and glucose. The pH indicator, bromthymol blue, is a deep Prussian blue at its alkaline end point (pH 7.6), yellow at its acidic end point (pH 6.0), and green when uninoculated (pH 6.7). Bacteria that are capable of using malonate as a source of energy and carbon produce alkaline by products that change the color of the medium to blue. Bacteria that are unable to use malonate as a carbon source usually do not grow and the pH of the medium does not change; the indicator remains green. Some malonate-negative strains may produce a yellow color owing to fermentation of glucose.

Ingredients and preparation: Mix the ingredients, heat to boiling, dispense into tubes, and sterilize at 121°C for 15 minutes.

Yeast extract	1 g
Ammonium sulfate	2 g
Dipotassium phosphate	0.6 g
Monopotassium phosphate	0.4 g
NaCl	2 g
Sodium malonate	3 g
D-Glucose	0.25 g
Bromthymol blue	0.025 g
Distilled water	1 L

Final pH 6.7

Procedure: Inoculate the test organism into malonate broth and incubate at 35°C for 18 to 24 hours.

2. Anti serum to *Salmonella* species

The antiserum using in Thailand are following :

1. ***Salmonella* polyvalent A – 67 antiserum** : is composed of group A, group B, group C, group D and every group to group 67 antisera.
2. ***Salmonella* polyvalent A –I antiserum** : is composed of group A, group B, group C, group D group E, group F, group G, group H, group I antiserum
3. ***Salmonella* polyvalent O : 17 – O : 67 antiserum** : is composed of group J (O :17) , group K (O : 18) to group O : 67
4. ***Salmonella* polyvalent H : H antiserum** : is composed of all of flagella both phase I and II which can described as H:a,H: b,H:c,H: d, H:eh toH: z₆₁
5. ***Salmonella* polyvalent H : L antiserum** : is composed of flagella as described as : H : l,v , H:v, H:w, H:z₁₃ , H:z₂₈
6. ***Salmonella* polyvalent H : G antiserum** : is composed of flagella as described as : H:f, H:g, H:s, H:t, H:m, H:p, H:q
7. ***Salmonella* polyvalent H : unspecific** : is composed of all phase 2 flagella as described as : H:1,2 , H:2, H:5, H:6, H:7 and H: z₆

The strains which will be tested must had been tested for biochemical characteristic already, the two important tests are TSI, Urea agar, LDC, ONPG, Indole, and VP. It should be subcultured the tested strain to NA or TSA plate and incubate at 37°C for 24 hr. before test for serological test. Detection of the O-antigen is performed by slide agglutination method.

Procedure for O – antigen test. Put one drop of 0.85% of NSS (control) on glass slide. By using loop, put one loop full cultures from TSA onto the drop of NSS and mix together, if show agglutination in 30 seconds it's indicate that it's rough strains, it can't be used to test with antiserum. If no agglutination with NSS (control) it'll be right to

test with antiserum. Put one drop of *Salmonella* polyvalent A-67 antiserum and *Salmonella* polyvalent A-I antiserum on each test area, put one loop full of cultures from NA to each antiserum, carefully mixed the culture and O-serum well. Rock the glass slide gently for one minute. If agglutination is found with which antisera indicated that the strain has that type antigen, but in this step we can only screen because the antisera in composed of variant group, can occurred in 2 antisera, agglutinating both with A-67 and A-I indicated that the strain will be a group of A-I group, agglutinating with A-67 but not agglutinating with A-I determined that the strain will be a group of 17-67. Then test with each single group antisera, if positive with in group, report the strain is that group and further test with O antigen of that group flower Antigenic Formulas of The *Salmonella* serovars 1997, edition 7th and 2001, edition 8th, for example, if positive with group B, must test with O:1, O:4, O:5, O:12, O:27 antisera, if positive with group C, must test with O:6, O:7, O:8, O:14, O:20 antisera, if positive with group D, must test with O:1, O:9, O:12, O:46 antisera, if positive with group E, must test with O:1, O:3, O:10, O:15, O:19, O:34 antisera.

Procedure for Vi antigen test. Some serovars of *Salmonella* Typhi, and *Salmonella* Paratyphi C don't agglutinate with O – antigen for the first step, because these strains almost have Vi – antigen, then must test with Vi-antiserum for the first step. In some case it's necessary to suspend the strain in NSS and boiled at 100^oC for 15 minutes to destroy Vi – antigen. Then take the sediment to test with O – antigen, will be positive.

Procedure for H – antigen test (Sven Gard Technique). Transfer the culture from TSA to swarm agar by spot-inoculate at the center, incubate at 37^oC for 18-24 hrs. Motility strains will spread on the surface of swarm agar. Use these strains to test for H – antigen, add one drop of H-antiserum and one drop 0.85% NSS (control) separately to each test area on the glass slide. Using loop, pick culture from the edge of swarm agar, carefully mix with H-antiserum and 0.85% NSS (control) separately. Rock

the glass slide gently for 30 seconds to 1 minutes. Observe the reaction, not agglutinate with 0.85%NSS (control). Start from polyvalent H antiserum for the first step to reduce your work time, so that always use polyvalent H:H antiserum, polyvalent H:L antiserum, polyvalent H:G antiserum, H:unspecific antiserum, if positive with polyvalent H:H antiserum and polyvalent H:L antiserum further test for specific antisera such as : H : l, v , H : v , H : w , H : z₁₃ , H : z₂₈ . If positive with polyvalent H : H antiserum and polyvalent H : G antiserum further test for specific antisera such as H : f , H : g , H : s , H : t , H : m , H : p , H : q. If positive with polyvalent H : H antiserum and polyvalent H : unspecific antiserum, test for specific antisera such as H:1,2 , H:2, H:5, H:6, H:7 and z₆. If positive with only polyvalent H: H antiserum, test for specific antisera H : a , H : b , H : c , H : d , H : eh etc. By choosing the specific antisera which specified to often found serovars. Absorbing for H – antigen, by drop concentrated antiserum, should be 1 : 800 or 1 : 1,600 titer, which specified to be found for 0.09 ml. in small petridish, pour melted swarm agar, shake well to mix homogeneous. Allow swarm agar to cool, inoculate culture from first plate swarm agar and incubate at 37^oC for 18 – 24 hrs. Specific antibody in swarm agar will be combined with antigen, colony which isn't specified will spread, further test for another phase antigen. And test for another phase H – antigen, using the same procedure. Absorbing for both 2 phase H-antigen, by drop specified concentrated antiserum another one petridish and pour method swarm agar, shake vigorously, and allow agar to cool. Transfer culture from the second petridish to the last petridish, incubate at 37^oC for 18 hrs. Observe the result if there isn't any spread strain in swarm agar, report 2 phase antigen which found. (Popoff and Minor, 1997; 2001)

APPENDIX II**Reagents for molecular analysis****Stock solution****1. 5 M NaCl (pH 8.0)**

NaCl	292.2 g
Deionized water	1,000 ml

This stock reagent was prepared by dissolved by 292.2 g of NaCl in 750 ml of deionized water, then the pH was adjusted to 8.0 with conc. HCl. The final volume was bought up to 1,000 ml with deionized water. The stock reagent steriled by autoclaving at 121°C, 15 pounds/inch² pressure, for 15 minutes. The stock reagent was stored at room temperature.

2. 0.5 M EDTA (pH 8.0)

Ethylene diaminetetraacetic acid	186.12 g
Deionized water	1,000 ml

This stock reagent was prepared by dissolved by 186.12 g of ethylene diaminetetraacetic acid in 750 ml of deionized water, then the pH was adjusted to 8.0 with conc. HCl. The final volume was bought up to 1,000 ml with deionized water. The stock reagent steriled by autoclaving at 121°C, 15 pounds/inch² pressure, for 15 minutes. The stock reagent was stored at room temperature.

3. 1 M Tris (pH 8.0)

Tris base	121.14 g
Deionized water	1,000 ml

This stock reagent was prepared by dissolved by 121.14 g of Tris base in 750 ml of deionized water, then the pH was adjusted to 8.0 with conc. HCl. The final volume was bought up to 1,000 ml with deionized water. The stock reagent

steriled by autoclaving at 121°C, 15 pounds/inch² pressure, for 15 minutes. The stock reagent was stored at room temperature.

4. 10 X TBE

Tris base	108	g
Boric acid	55	g
EDTA	7.44	g
Deionized water	1,000	ml

This stock reagent was prepared by dissolved all of ingredients in 1,000 ml of deionized water. The stock reagent steriled by autoclaving at 121°C, 15 pounds/inch² pressure, for 15 minutes. The stock reagent was stored at room temperature.

Buffer

1. SE buffer (75 mM NaCl pH 8.0, 25 mM EDTA pH 8.0)

5 M NaCl	15	ml
0.5 M EDTA	50	ml

This buffer was prepared by mixing these stock solution in 1,000 ml of sterile deionized water. The SE buffer was stored at room temperature.

2. Lysis buffer (50 mM Tris pH 8.0, 50 mM EDTA pH 8.0, 1% Sarcosyl)

1 M Tris	50	ml
0.5 M EDTA	100	ml
Sarcosyl	10	g

This buffer was prepared by mixing these stock solution in 1,000 ml of sterile deionized water. The lysis buffer was stored at room temperature.

3. TE buffer (10 mM Tris pH 8.0, 1mM EDTA pH 8.0)

1 M Tris	10	ml
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0.5 M EDTA	2	ml
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This buffer was prepared by mixing these stock solution in 1,000 ml of sterile deionized water. The TE buffer was stored at room temperature.

4. 0.5 X TBE buffer

10 X TBE	125	ml
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Sterile deionized water	2,375	ml
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This buffer was prepared by mixing, and then was stored at room temperature.

Enzymes

1. Proteinase K (10 mg/ml)

Proteinase K	100	mg
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Sterile deionized water	10	ml
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To prepared this stock reagent, 100 mg of Proteinase K was dissolved in 10 ml of sterile deionized water. The stock reagent was stored at -20°C

2. Restriction endonuclease enzyme *Xba*I

2.1 Pre-incubation (per reaction)

10X restriction enzyme <i>Xba</i> I buffer	10	μl
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Sterile deionized water	90	μl
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This buffer was freshly preped by mixing before use.

2.2 Restriction enzyme digestion (per reaction)

Restriction enzyme <i>Xba</i> I (15 units/μl)	1.33	μl
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10X restriction enzyme <i>Xba</i> I buffer	5	μl
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0.1% bovine serum albumin	5	μl
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Sterile deionized water	38.67	μl
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This restriction enzyme *Xba*I solution was freshly

prepared by mixing before use.

Agarose gel

1. 1% Low melting point agarose gel (per sample)

Low melting point agarose	0.1	g
TE buffer	10	ml

This 1% low melting point agarose gel was prepared by suspending 0.1 g of low melting point agarose in 10 ml of TE buffer. The agarose was melted by microwave oven.

2. 1% Ultrapure high melting temperature agarose gel

Ultrapure high melting temperature agarose	0.9	g
0.5X TBE	90	ml

This 1% ultrapure high melting temperature agarose gel was prepared by suspending 0.9 g of ultrapure high melting temperature agarose in 90 ml of 0.5X TBE buffer. The agarose was melted by microwave oven.

Reagent

Ethidium bromide solution

One pellet (11 mg) of ethidium bromide was dissolved in 11 ml of ultrapure water.

Working solution: The 40 μ l ethidium bromide stock solution was mixed with 300 ml ultrapure water before use.

APPENDIX III

Chemical agents, Enzyme, Molecular marker, Materials, and Instruments

1. Chemical agents

- Low melting point agarose (Bio-Rad, USA)
- Ultrapure high melting temperature agarose (USB, USA)
- Tris base (USB, USA)
- EDTA (USB, USA)
- Sodium chloride (Merch, USA)
- Sodium lauroyl sarcosine (USB, USA)
- Boric acid (Bio-Rad, USA)
- Ethidium bromide solution (USB, USA)
- Anti-serum for *Salmonella* species (SAP, Thailand)
- Mc Farland No.4 (Bio Merieux)

2. Enzyme and Molecular Marker

- Proteinase K (USB, USA)
- Restriction endonuclease enzyme *Xba*I (USB, USA)
- Restriction endonuclease enzyme *Xba*I buffer (USB, USA)
- Bovine Serum Albumin (USB, USA)
- Lambda ladder marker (USB, USA)

3. Materials

- Eppendorf tube
- Test tube (Pyrex, USA)
- Plastic tube bottom size 50 ml (Falcon,)

Tip
Cylinder
Beaker
Flask
Glass bottle (Duran, Germany)

4. Instruments

Incubator 37°C (Mettler, Germany)
Shaking waterbath (Mettler, Germany)
Vortex mixer (Scientific, USA)
Eppendorf microcentrifuge (Tomy Seiko, Japan)
Freezer (-20°C) (Sanyo, Japan)
Refrigerator (Sanyo, Japan)
Automatic pipette, p10/p20/p200/p1000/p5000 (Gilson Medical Electronic, France)
pH meter (Beckman, USA)
Pulsed-Field Gel Box (Bio-Rad, USA)
Pump, Gel Molds (Bio-Rad, USA)
Cooling system (Bio-Rad, USA)
Power supply, Pulsed wave switcher (Bio-Rad, USA)
Contour-clamped homogenous electric field apparatus (Bio-Rad, USA)
UV transilluminator (Bio-Rad, USA)
Polaroid camera (Bio-Rad, USA)
Biological safety cabinet (Yamato, Japan)
Autoclave (Yamato, Japan)
Microwave oven (Sanyo, Japan)

BIOGRAPHY

Mr. Thanarat Kaewsawang was born on October 21, 1974 in Bangkok, Thailand. I graduated with the Bachelor degree of Science in Medical Technology from the Faculty of Medical Technology, Rangsit University in 1996. I am a medical technologist of Department of Pathology, Nopparat Rajathanee General Hospital, Bangkok, Thailand.