

Dehydrated Culture Media Bases / Media Supplements

# **Technical Information**

# **Triple Sugar Iron Agar**

### Product Code: DM 1021

**Application:** Triple Sugar Iron Agar is used for the identification of gram-negative enteric bacilli on the basis of dextrose, lactose and sucrose fermentation and hydrogen sulphide production.

Composition**						
Ingredients	Gms / Litre					
Peptic digest of animal tissue	10.000					
Casein enzymic hydrolysate	10.000					
Yeast extract	3.000					
Beef extract	3.000					
Lactose	10.000					
Sucrose	10.000					
Dextrose	1.000					
Sodium chloride	5.000					
Ferrous sulphate	0.200					
Sodium thiosulphate	0.300					
Phenol red	0.024					
Agar	12.000					
Final pH ( at 25°C)	7.4±0.2					
**Formula adjusted, standardized to suit performance parameters						

### Principle & Interpretation

Sulkin and Willett originally proposed Triple Sugar Iron Agar<sup>(1)</sup> Which was modified by Hajna<sup>(2)</sup> for identifying members of Enterobacteriaceae. This medium complies with the recommendation of APHA, for the examination of meat and food products<sup>(3),</sup> Including milk and dairy products<sup>(4)</sup> microbial limit test for confirming the presence of Salmonellae<sup>(5,6)</sup> and in the identification of gram-negative bacilli<sup>(6,7).</sup>

Casein enzymic hydrolysate, peptic digest of animal tissue, yeast extract and beef extract provide nitrogenous compounds, sulphur, trace elements and vitamin B complex etc. Sodium chloride maintains osmotic equilibrium. Lactose, sucrose and dextrose are the fermentable carbohydrates. Sodium thiosulphate and ferrous ions make H<sub>2</sub>S indicator system. Phenol red is the pH indicator. Organisms that ferment glucose produce acids, turning the colour of the medium from red to yellow.

More amount of acids are liberated in butt (fermentation) than in the slant (respiration). Growing bacteria also form alkaline products from the oxidative decarboxylation of peptone and these alkaline products neutralize the large amounts of acid present in the butt. Thus the appearance of an alkaline (red) slant and an acid (yellow) butt after incubation indicates that the organism is a glucose fermenter but is unable to ferment lactose and/or sucrose. Bacteria that ferment lactose or sucrose (or both), in addition to glucose, produce large amounts of acid enables no reversion of pH in that region and thus bacteria exhibit an acid slant and acid butt. Gas production (CO<sub>2</sub>) is detected by the presence of cracks or bubbles in the medium, when the accumulated gas escapes. Thiosulphate is reduced to hydrogen sulphide by several species of bacteria and H<sub>2</sub>S combines with ferric ions of ferric salts to produce the insoluble black precipitate of ferrous sulphide. Reduction of thiosulphate proceeds only in an acid environment and blackening usually occurs in the butt of the tube. Triple Sugar Iron Agar should be used in parallel with Urea Agar / Broth (DM1112/DM1111) to distinguish between *Salmonella* and *Proteus* species. The reactions can be summarized as follows Alkaline slant / acid butt-only glucose fermented Acid slant / acid butt-glucose and sucrose fermented or glucose and lactose fermented or all the three sugars, glucose, lactose and sucrose fermented.Bubbles or cracks present-gas production Black precipitate present-H<sub>2</sub>S gas production Some members of the *Enterobacteriaceae* and H<sub>2</sub>S producing *Salmonella* may not be H<sub>2</sub>S positive on TSI Agar. Some bacteria may show H<sub>2</sub>S production on Kligler Iron Agar but not on TSI Agar. This can happen because utilization of sucrose in TSI Agar suppresses the enzymic pathway that result in H<sub>2</sub>S production.





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### Methodology

Suspend 64.52 grams of powder media in 1000 ml distilled water. Shake well & heat to dissolve the medium completely. Mix well and distribute into test tubes. Sterilize by autoclaving at 15 lbs pressure (121°C) for 15 minutes. Allow the medium to set in sloped form with a butt about <sup>1</sup> inch long.

Note: For better results, the medium can be sterilized by autoclaving at 10 lbs pressure (115°C) for 15 minutes.

### **Quality Control**

### Physical Appearance

Light yellow to pink homogeneous free flowing powder

#### Gelling

Firm, comparable with 1.2% Agar gel.

#### Colour and Clarity of prepared medium

Pinkish red coloured clear to slightly opalescent gel forms in tubes as slants.

#### Reaction

Reaction of 6.45% w/v aqueous solution at 25°C. pH : 7.4±0.2

pH Range:-7.20-7.60

#### Cultural Response/Characteristics

DM1021: Cultural characteristics observed after an incubation at 35-37°C for 18-24 hours.

Organism	Inoculum (CFU)	Growth	Slant	Butt	Gas	H₂S
Citrobacter freundii ATCC 8090	50-100		acidic reaction, yellowing ofthe medium	acidic reaction, yellowing of the medium	•	positive,blackening of medium
Enterobacter aerogene. ATCC 13048	s 50-100		acidic reaction, yellowing ofthe medium	acidic reaction,yellowing of the medium	positive reaction	negative, noblackening of medium
Escherichia coli ATCC 25922	50-100		acidic reaction, yellowing ofthe medium	acidic reaction,yellowing of the medium	reaction	negative, noblackening of medium
Klebsiella pneumonia ATCC 13883	50-100		acidic reaction, yellowing ofthe medium	acidic reaction,yellowing of the medium	positive reaction	negative, noblackening of medium
Proteus vulgaris ATCC 13315	50-100		alkalinereaction, redcolour of themedium	acidic reaction,yellowing of the medium	-	positive,blackening of medium
Salmonella Paratyphi A ATCC 9150	50-100		alkalinereaction, redcolour of themedium	acidic reaction,yellowing of the medium	positive reaction	negative, noblackening of medium
Salmonella Typhi ATCC 6539	50-100		alkaline reaction, redcolour of themedium	acidic reaction, yellowing of the medium	-	positive,blackening of medium
Salmonella Typhimurium ATCC 14028	50-100		alkalinereaction, redcolour of themedium	acidic reaction,yellowing of the medium	•	positive,blackening of medium
Shigella flexneri ATCC 12022	50-100		alkalinereaction, redcolour of themedium	acidic reaction,yellowing of the medium	negative reaction	negative, noblackening of medium
Escherichia coli ATCC 8739	50-100		acidic reaction, yellowing ofthe medium	acidic reaction,yellowing of the medium	positive reaction	negative, noblackening
Escherichia coli NCTC 9002	50-100		acidic reactionyellowing ofthe medium	acidic reaction,yellowing of the medium	positive reaction	negative, noblackening of





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Klebsiella pneumoniae 50-100 luxuriantacidic reaction, yellowing ATCC 10031

ofthe medium

acidic reaction, yellowing of the medium

medium positive negative, noblackening reaction of medium

# Storage and Shelf Life

Dried Media: Store below 30°C in tightly closed container and use before expiry date as mentioned on the label. **Prepared Media:** 2-8<sup>0</sup> in sealable plastic bags for 2-5 days.

### **Further Reading**

1. Sulkin E.S. and Willett J.C., 1940, J. Lab. Clin. Med., 25:649.

2. Hajna A.A., 1945, J. Bacteriol, 49:516.

3. Downes F. P. and Ito K., (Eds.), 2001, Compendium of Methods for the Microbiological Examination of Foods, 4th Ed., APHA, Washington, D.C.

4. Wehr H. M. and Frank J. H., 2004, Standard Methods for the Microbiological Examination of Dairy Products, 17th Ed., APHA Inc., Washington, D.C.

5. Finegold S. M. and Baron E. J., 1986, Bailey and Scotts Diagnostic Microbiology, 7th Ed., The C.V. Mosby Co., St. Louis. 6. Eaton A. D., Clesceri L. S. and Greenberg A. W., (Eds.), 2005, Standard Methods for the Examination of Water and Wastewater, 21st Ed., APHA, Washington, D.C.

7. MacFaddin J., 1985, Media for Isolation-Cultivation-Identification-Maintenance of Medical Bacteria, Vol. 1, Williams and Wilkins, Baltimore.

### Disclaimer :

- User must ensure suitability of the product(s) in their application prior to use.
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