

Bases / Media Supplements

Technical Information

Kanamycin Esculin Azide Broth

Product Code: DM 1776

Application: - Kanamycin Esculin Azide Broth is used for isolation of Group D Streptococci in foodstuffs.

Composition**						
Ingredients	Gms / Litre					
Casein enzymic hydrolysate	20.000					
Yeast extract	5.000					
Sodium chloride	5.000					
Sodium citrate	1.000					
Esculin	1.000					
Ferric ammonium citrate	0.500					
Sodium azide	0.150					
Kanamycin sulphate	0.020					
Final pH (25°C)	7.0±0.2					
**Formula adjusted, standardized to suit performance parameters						

Principle & Interpretation

Enterococci may be considered an essential part of the autochthonous microflora of humans and animals. Faecal streptococci with group D Lancefield antigens are grouped as Enterococci. Lancefield Group D- faecal Streptococci are contaminants of various food commodities, especially those of animal origin. Kanamycin Esculin Azide Broth is formulated as per Mossel et al ^(1, 2) to detect Enterococci in foodstuffs and used it for the dip slide technique for bacteriological monitoring of foods ⁽³⁾.

Casein enzymic hydrolysate and yeast extract provides essential nutrients for Enterococci. Kanamycin sulphate and sodium azide are the selective inhibitory components. Esculin and ferric ammonium citrate together forms the indicator system to detect esculin-hydrolyzing Streptococci, which form black zones around the colonies. The black zones are produced from the formation of black iron phenolic compounds derived from esculin-hydrolysis products and ferrous ions. Mossel et al ⁽⁴⁾ described the following procedure - 1gm or 1ml mixed food is added to 9 ml of pre-chilled diluent (Tryptone water DM1463) and ton told dilutions are prepared. The ton told dilutions are inoculated in Kanamycin Esculin Azide Broth and incubated at 35-37°C for 16-24 hours. If blackening of medium occurs, streaking is done on agar (DM1510) and after incubation confirmatory tests are carried out.

There is no universal medium that will isolate all strains of Enterococci⁽⁵⁾. Unless a presumptive count is acceptable all isolates should have their identity confirmed with further tests.

Methodology

Suspend 32.67 grams of powder media in 1000 ml distilled water. Shake well & heat to dissolve the medium completely. Sterilize by autoclaving at 15 lbs pressure (121°C) for 15 minutes. Dispense as desired.

Caution: Sodium azide has a tendency to form explosive metal azides with plumbing materials. It is advisable to use enough water to flush of the disposables.





Dehydrated Culture Media Bases / Media Supplements

Quality Control

Physical Appearance

Cream to yellow w/greenish tinge homogeneous free flowing powder

Colour and Clarity of prepared medium

Medium amber coloured, clear solution in tubes.

Reaction

Reaction of 3.27% w/v aqueous solution at 25°C. pH : 7.0±0.2

pH Range 6.80-7.20

Cultural Response/ characteristices

DM 1776 Cultural characteristics observed after an incubation at 35-37°C for 18-24 hours.

Organism	Inoculum (CFU) Growth		Recovery	Esculin Hydrolysis
Enterococcus bovis ATCC 27960	50-100	50-100	>=50%	positive, blackening of medium around the colony
Enterococcus faecium ATCC 19434	50-100	50-100	>=50%	positive, blackening of medium around the colony
Escherichia coli ATCC 25922	>=10 [°]	>=10°	0%	
Enterococcusfaecalis ATCC 29212	50-100	50-100	>=50%	positive, blackening of medium around the colony
Staphylococcus aureus ATCC 25923	>=10 [°]	>=10 [°]	0%	

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° in sealable plastic bags for 2-5 days.

Further Reading

1. Mossel D. A. A., Bijker P. G. H. and Eelderink I., 1978, Arch. Lebensmittel - hyg., 29: 121.

2. Mossel D. A .A. el al, 1978, In : `Streptococci., Skinner F. A. and Quesnel L. B. (Eds.), SAB Symposium, Series No.7, Academic Press, London.

3. Mossel D. A. A. et al, 1976, Lab. Practice, 25:3 93.

- 4. Mossel D. A. A., Harrenwijn G. A. and Elzebroek B. J. M., 1973, UNICEF, Geneva.
- 5. Reuter G., 1985, Inter. J. Food. Microbiol., 2.103-114.

Disclaimer:

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