

Bases / Media Supplements

# **Technical Information**

## Modified Duncan Strong (DS) Medium

#### Product Code: DM 2237

**Application:** - Modified Duncan Strong (DS) Medium is used for isolation and differentiation of *Clostridium perfringens* from other clostridia from foods on the basis of raffinose fermentation.

Composition**		
Ingredients	Gms / Litre	
Proteose peptone	15.000	
Yeast extract	4.000	
Sodium thioglycollate	1.000	
Disodium phosphate	10.000	
Raffinose	4.000	
Final pH (25°C)	7.8±0.2	
**Formula adjusted, standardized to suit perform	nance parameters	

#### **Principle & Interpretation**

*Clostridium perfringens*, a gram-positive, rod shaped, anaerobic, spore-forming bacteria, is the major cause of food poisoning in humans. This organism *is* commonly found in raw meats, poultry, dehydrated soups and sauces, raw vegetables and certain other foods or food ingredients.

A heat labile enterotoxin produced only by sporulating cells induces the major symptoms of diarrhea in perfringens poisoning.

Modified Duncan Strong (DS) Medium is formulated as per Duncan and Strong <sup>(1)</sup>, This is also recommended by APHA <sup>(2)</sup> for the isolation and differentiation of *C. perfringens* from other Clostridia from foods on the basis of raffinose fermentation. It is also used for the rapid detection of the *Clostridium perfringens* enterotoxin <sup>(3)</sup>.

Proteose peptone and yeast extract provide nitrogenous compounds and other nutrients for bacterial growth. Sodium thioglycollate helps to create anaerobic conditions suitable for clostridial growth. Disodium phosphate acts as a buffering agent. Raffinose in the medium is fermented only by by *C. perfringens* to produce acid within 72 hours of inoculation which can be tested by transfer 1 ml of culture to a test tube or spot plates and adding 2 drops of 0.04% bromothymol blue. A yellow colour indicates acid production.

Inoculate about 2 gm of the food sample into 15-20 ml of Chopped Liver Broth (DM1606). After an incubation at 35-37°C for 20-24 hours, tubes showing turbidity are streaked on Perfringens Agar Base (DM1837) containing Egg Yolk Emulsion (MS2045) to obtain presumptive *C. perfringens*. These presumptive colonies can be confirmed by inoculating into Motility Nitrate Medium, Buffered (DM1630), Lactose Gelatin Medium (DM1628) and Modified Duncan Strong (DS) Medium (DM2237) (2).

#### Methodology

Suspend 34 grams of powder media in 1000 ml distilled water and mix thoroughly. Sterilize by autoclaving at 15 lbs pressure (121°C) for 15 minutes. Dispense into sterile tubes. Check one or two tubes for measuring the pH.





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### Quality Control

Physical Appearance				
Cream to yellow homogeneous free flowing the second s	ng powder			
Colour and Clarity of prepared medium				
Yellow coloured clear solution without an	y precipitate			
Reaction				
Reaction of 3.4% w/v aqueous solution at	25°C. pH : 7.8±0.2			
pH range 7.60-8.00				
Cultural Response/ characteristices				1.00
DM 2237: Cultural characteristics observ	ved after an incubati	on at 35-37°C for 48-72 ho	burs.	
Organism	Inoculum (CFU)	Growth	Raffinose fermentation	
Clostridium perfringens ATCC 12924	50-100	good-luxuriant	positive reaction	
Clostridium sporogenes ATCC 11437	50-100	good-luxuriant	negative reaction	

### 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. Duncan C. and Strong D., 1969, Appl. Microbiol., 16: 82.

2. Labbe R. G. and Rey D. K., 1979, Appl. Microbiol., 13: 559.

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

#### **Disclaimer**:

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