

Dehydrated Culture Media Bases / Media Supplements

Technical Information

Casman Agar

Product Code: DM 1201

Application: - Casman Agar with blood is used for isolation of fastidious microorganisms from clinical specimens under reduced oxygen tension.

Composition**		
Ingredients	Gms / Litre	
Proteose peptone	10.000	
Tryptose	10.000	
Beef extract	3.000	
Dextrose	0.500	
Corn starch	1.000	
Sodium chloride	5.000	
Nicotinamide	0.050	
p-Amino benzoic acid (PABA)	0.050	
Agar	14.000	
Final pH (at 25°C)	7.3±0.2	

**Formula adjusted, standardized to suit performance parameters

Principle & Interpretation

Being fastidious in nature *Haemophilus* and *Neisseria* require the addition of X and V- growth factors for in vitro cultivation ⁽¹⁾. Blood-enriched medium for cultivation of *Haemophilus* and gonococci has been describe else where ⁽¹⁻³⁾. The medium was developed to replace the previously described media whose preparation is time consuming using fresh and heated blood and meat infusion to supply the essential nutrients for growth of these fastidious organisms⁽²⁻³⁾. Blood supplies both factor-X (hemin) and factor-V (Nicotinamide Adenine Dinucleotide), which is required for growth of *Haemophilus influenzae*. Sheep blood lacks factor-V due to NADase, an enzyme that destroys factor-V (4). Horse and rabbit blood supplies both X&V factor and are relatively free of NADase activity, Nicotinamide is added to medium to inhibit nucleotidase of erythrocytes that may destroy factor V.

Proteose peptone, tryptose and beef extract provide amino acids and other complex nitrogenous nutrients. Dextrose improves growth of pathogenic cocci. Corn starch prevents fatty acids from inhibiting the growth of *Neisseria gonorrhoeae,* without interfering with haemolytic reaction and also neutralizes the inhibitory action of dextrose. Inoculate the medium as soon as the specimen arrives at the laboratory. After incubation *H. influenzae* produces colourless to grey colonies with a characteristic mousy odour while *N. gonorrhoeae* produces small colourless to greyish-white colonies.

Methodology

Suspend 43.6 grams of poder media in 1000 ml distilled water. Shake well & heat to boil to dissolve the medium completely. Sterilize by autoclaving at 15 lbs pressure (12 1°C) for 15 minutes. Cool to 50°C and aseptically add 0.15% v/v sterile water lysed blood (water: blood:: 3:1) of 5% sterile blood. Mix well and dispense as desired.





Bases / Media Supplements

Quality Control

Physical Appearance

Cream to yellow homogeneous free flowing powder

Gelling

Firm, comparable with 1.4% Agar gel.

Colour and Clarity of prepared medium

Basal medium: Yellow coloured clear to slightly opalescent gel. After addition of 5%w/v sterile defibrinated blood : Cherry red coloured After addition of 5%w/v sterile defibrinated blood : Cherry red coloured After addition of 5%w/v sterile defibrinated blood: opaque gel forms in Petri plates.

Reaction

Reaction of 4.36% w/v aqueous solution at 25°C. pH : 7.3±0.2

pH range 7.10-7.50

Cultural Response/ characteristices

DM 1201: Cultural characteristics observed with added water-lysed blood, after an incubation at 35-37°C for 40-48 hours.

Organism	Inoculum (CFU)	Growth	Recovery	Haemolysis
Haemophilus influenzae ATCC 35056	50-100	good	50-70%	none
Neisseria meningitidis ATCC 13090	50-100	luxuriant	>=70%	none
Streptococcus mitis ATCC 9811	50-100	luxuriant	>=70%	beta
Streptococcus pneumoniae ATCC 6303	50-100	luxuriant	>=70%	alpha
Streptococcus pyogenes ATCC 19615	50-100	luxuriant	>=70%	beta

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. Casman, 1947, Am. J. Clin. Pathol., 17:28 1.

- 2. Casman, 1942, J. Bact., 43:33.
- 3. Casman, 1947, J. Bact., 53:561.

Krunveide and Kuttner, 1938, J. Exp. Med., 67:429

Disclaimer:

- User must ensure suitability of the product(s) in their application prior to use.
- The product conform solely to the technical information provided in this booklet and to the best of knowledge research and development work carried at **CDH** is true and accurate
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