

Bases / Media Supplements

# **Technical Information**

### Casman MiVeg Broth Base

### Product Code : VM1766

Application:- Casman MiVeg Broth Base with blood is recommended for isolation of fastidious microorganisms such as Haemophilus influenzae & Neisseria gonorrhoeaefrom clinical specimens, under reduced

Composition		
Ingredients	Gms / Litre	
MiVeg peptone No.3	10.00	
MiVeg hydrolysate No.1	10.00	
MiVeg extract	3.00	
Dextrose	0.50	
Corn starch	1.00	
Sodium chloride	5.00	
Nicotinamide	0.05	
p-Amino benzoic acid (PABA)	0.05	
Final pH ( at 25°C)	7.2±0.2	
** Formula adjusted standardized to suit perfor	mance personators	

\*\* Formula adjusted, standardized to suit performance parameters.

#### Principle & Interpretation

Casman MiVeg Media is prepared by using vegetable peptones instead of animal based peptones thereby making this media free from BSE/TSE risks. Members of the genus *Haemophilus* and *Neisseria* are fastidious microorganisms that require addition of growth factors. This media is the modifications of the media described by Casman (1, 2, 3) for cultivatior of *Haemophilus* and *Gonococci*, which replaced previous formulations requiring fresh meat infusion, fresh and heated blood etc.

It contains MiVeg peptone No.3, MiVeg hydrolysate No.1 and MiVeg extract supplies amino acids and other complex nitrogenous nutrients. Dextrose enhances growth of pathogenic cocci. Corn starch inhibits the growth of *Neisseria gonorrhoeae* 8 prevents fatty acid without interfering with the haemolytic reaction and it also neutralizes the inhibitory action of dextrose. Addition of blood aids the growth factors required for *Haemophilus influenzae* i.e. hemin or X factor and Nicotinamide Adenine Dinucleotide (NAD) or V factor. Horse and rabbit blood are preferred as they are relatively free of NADase, an enzyme that destroys V factor (4). Nicotinamide is added to the medium to inhibit nucleotidase of erythrocytes that destroys V factor. PABA serves as a growth factor.

The medium should be inoculate as soon as the specimen arrives at the laboratory. After incubation Haemophilus influenzae produces colourless to grey colonies with a characteristic 'mousy' odour while *Neisseria gonorrhoeae* produces small colourless to greyish-white colonies on this medium.

### Methodology

Suspend 29.6 grams of powder media in 1000 ml distilled water. Mix thoroughly. Heat to boiling to dissolve the medium completely. Sterilize by autoclaving at 15 lbs pressure (121°C) for 15 minutes. Cool to 45°-50°C and add 0.5ml of sterile defibrinated rabbit blood to each tube. Mix well and dispense as desired.

### **Quality Control**

#### Physical Appearance

Light yellow coloured, may have slightly greenish tinge, homogeneous, free flowing powder.

#### Colour and Clarity of prepared medium

Basal medium is yellow coloured. With addition of blood, cherry red coloured opalescent solution in tubes.

#### Reaction

Reaction of 2.96 % w/v aqueous solution pH: 7.2 ±0.2 at 25°C





Dehydrated Culture Media Bases / Media Supplements

#### pH range

#### 7.0-7.4 Cultural Response/Characteristics

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Cultural characteristics observed after an incubation at 35-37°C for 40-48 hours		
Organisms (ATCC)	Inoculum (CFU)	Growth
Haemophilus influenzae (35056)	102-103	good-luxuriant
Neisseria meningitidis (13090)	102-103	good-luxuriant
Streptococcus pneumoniae (6303)	102-103	good-luxuriant
Streptococcus pyogenes (19615)	102-103	good-luxuriant
Streptococcus mitis (9895)	102-103	good-luxuriant

## 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., 17:281.
- 2. Casman, 1942, J. Bact., 43:33.
- 3. Casman, 1947, J. Bact., 53:561.
- 4. 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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