

TECHNICAL DATA SHEET

Article No. 9166

MYP Agar, prepared plates

SYNONYMS

Mannitol Egg Yolk Polymyxin Agar, Bacillus Cereus Mossel Agar

SPECIFICATION

Prepared plates. Selective solid medium, according to Mossel, for the isolation and identification of Bacillus cereus from food samples. FIL-IDF 181, ISO 7932, ISO 21871.

Color: Orange

pH: 7.2 ± 0.2 at 25 °C

COMPOSITION IN G/L

Casein peptone 10.000 Mannitol 10.000 Sodium chloride 10.000 Meat extract 1.000 Phenol red 0.025 12.000 Agar Polymixin B Sulphate 100,000IU Egg Yolk 100 ml

PACKAGING DETAILS

9166-20PLATES

20 prepared plates 90 mm

Content: $21 \pm 2 \text{ ml}$

Packaging unit: 1 box with 2 packs of 10 plates/pack. Single cellophane.





GUIDELINES

Description:

Mossel's formulation has been developed to detect and enumerate *B. cereus* in any food stuff. It is both selective and differential for this microorganism. Polymyxin addition inhibits most of the accompanying bacteria, but does not affect the growth of *B. cereus*. This bacterium does not ferment mannitol and thus there is no change in the colour of the indicator around the colonies. The lecithinase activity of *B. cereus* produces a halo or zone of white precipitate around the colonies.

A count of *B. cereus* of more than 100.000 cells/g of food sample is considered hazardous, since the accumulated phosphorilcholine may cause toxic symptoms in children. For this reason a viable enumeration has to be performed to evaluate the actual population of cells.

Technique:

For plate inoculation follow the laboratories' standard methods or the applicable norms (spiral plating method, econometric methods, streak plating, dilution banks, spread plating with drigralsky rod, among them). According to the authors, dehydrated or dry samples must be treated in the following way: 20 g of sample is mixed with 90 ml of Tryptone Water for a minimum period of 1 hour, at room temperature. Afterwards, add an additional 90 mL of Tryptone Water and homogenize. If necessary dilute 1:10. Proceed to a 1/10 serial dilution bank using Tryptone water as the diluent if necessary. With a Drigalsky loop, spread aliquots of 0,1 mL over the surface of the agar plates and let the agar medium absorb the aliquots. Incubate the plates at 30 °C for 18-24 hours to allow spore germination before giving definite results.

Suspected colonies have the following appearance: irregular borders, pink colour becoming purple in the centre, with a halo of white precipitate (mannitol +). Colonies with yellow halos must be discounted (mannitol -). Confusion with other colonies of Gram positive bacilli is possible, and hence, confirmation tests must be carried out i.e. glucose fermentation, gelatine degradation and nitrate reduction.

MICROBIOLOGICAL CONTROL

Inoculate: Practical range 100 ± 20 CFU. min. 50 CFU (productivity)/ 10₄-10₆ (selectivity).

Microbiological control according to ISO 11133:2014/A1:2018.

Aerobiosis. Incubation at 30 ± 1 °C, read after 24 ±3h - 44 ±4h.

Microorganism	Growth
Bacillus cereus ATCC® 11778, WDCM 00001	Good-pink colonies with halo of precipitation
Escherichia coli ATCC® 25922, WDCM 00013	Inhibited
Bacillus subtilis ATCC® 6633, WDCM 00003	Yellow colonies without halo

Sterility control:

Incubation 48 hours at 30-35 $^{\circ}$ C and 48 hours at 20-25 $^{\circ}$ C: NO GROWTH. Check at 7 days after incubation in same conditions.





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STORAGE

2-14 °C

SHELF LIFE

3 months unopened from date of manufacture

updated: 30.08.2022

