CE IVD



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INSTRUCTIONS FOR USE

TRYPTIC SOY BROTH

Ready-to-use flasks



Tryptic Soy Broth from the left: un-inoculated tube, growth of *B.subtilis*

1 - INTENDED USE

In vitro diagnostic. General purpose medium for the sterility test and for the microbiological examination of pharmaceutical products according to the harmonized methods of EP, USP, JP. For suspension, enrichment and cultivation of microbial strains isolated from clinical specimens on other culture media.

2 - COMPOSITION -TYPICAL FORMULA *

Pancreatic digest of casein	17.0 g
Soy peptone	3.0 g
Sodium chloride	5.0 g
Dipotassium hydrogen phosphate	2.5 g
Glucose	2.5 g
Purified water	1000 mL

^{*}the formula may be adjusted and/or supplemented to meet the required performances criteria.

3 - PRINCIPLE OF THE METHOD AND EXPLANATION OF THE PROCEDURE

Tryptic Soy Broth is a general purpose medium that supports the growth of a wide variety of aerobic and facultative anaerobic bacteria and fungi. Tryptic Soy Broth is used for sterility testing and for the microbiological examination of pharmaceutical products with EP, USP, JP harmonized methods (casein soybean digest broth) and complies with the quality specifications reported therein.²

In clinical microbiology Tryptic Soy Broth is used for suspension, enrichment and cultivation of microbial strains isolated on other culture media and for the preparation of inocula in quality control test procedures.

Supplemented with 20% glycerol, Tryptic Soy Broth may be used for the long-term maintenance of microbial strains; supplemented with 0,1-0,15% of agar it may be used for enhancing growth of anaerobes. Tryptic Soy Broth is used in food bacteriology as the basal medium to which a variety of selective compounds may be added for selective enrichment of pathogens. Tryptic Soy Broth may be used also for blood cultures.

Pancreatic digest of casein and soy peptone are sources of carbon, nitrogen, vitamins and minerals for microbial growth; glucose is a source of energy; sodium chloride maintains osmotic balance, dipotassium hydrogen phosphate is included as a buffer system.

4- METHOD OF PREPARATION

The medium is ready to use. If required by the user's procedures, add the needed sterile supplements and distribute the medium into sterile tubes under aseptic conditions.

5 - PHYSICAL CHARACTERISTICS

 $\begin{array}{ll} \mbox{Medium appearance} & \mbox{yellow, limpid} \\ \mbox{Final pH at 20-25°C} & 7.3 \pm 0.2 \end{array}$

6 - MATERIALS PROVIDED - PACKAGING

Product	Туре	REF	Pack
Tryptic Soy Broth	Ready-to-use flasks	5121552	6 x 100 mL; 6 glass bottles with flat bottom and aluminium screw-cap; packaging: cardboard box.
		5121553	6 x 200 mL; 6 glass bottles with flat bottom and aluminium screw-cap; packaging: cardboard box.

7 - MATERIALS REQUIRED BUT NOT PROVIDED

Incubator and laboratory equipment as required, sterile tubes, sterile loops, needles and swabs, ancillary culture media and reagents for the identification of the colonies.

8 - SPECIMENS

Un-supplemented Tryptic Soy Broth should not be used for the direct inoculation of clinical specimens. In clinical microbiology the specimens consist of microbial colonies grown on other culture media. In pharmaceutical microbiology, samples consist of products on which to perform the sterility test or the detection for specific microorganisms. Refer to the European Pharmacopoeia for sample collection and transport procedures.²

9 - TEST PROCEDURE

With a bacteriological needle or loop inoculate the liquid medium in a test tube or bottle with a colony grown on another isolation medium. Incubate at the temperature and for the time required by laboratory procedures. Usually, an incubation temperature of 35 ± 2° C for 18-24 hours is adequate for cultivation of common anaerobes and facultative anaerobes.

For sterility testing and for use of Tryptic Soy Broth as a pre-enrichment medium for the detection of specific microorganisms in pharmaceutical products, consult the European Pharmacopoeia.²







10 - READING AND INTERPRETATION

The presence of microorganisms is indicated by a varying degree of turbidity, specks and flocculation in the medium. The un-inoculated control remains clear and without turbidity after incubation. The characteristics of the growths are closely related to the type or types of microorganisms grown.

11 - USER QUALITY CONTROL

All manufactured lots of the product are released for sale after the Quality Control has been performed to check the compliance with the specifications. However, the end user can perform its own Quality Control in accordance with the local applicable regulations, in compliance with accreditation requirements and the experience of the Laboratory. Here below are listed some test strains useful for the quality control of un-supplemented medium.³

CONTROL STRAINS INCUBATION T°/ t / ATM EXPECTED RESULTS S.aureus ATCC 25923 35-37°C / 18-24H / A good growth E.coli ATCC 25922 35-37°C / 18-24H / A good growth

User quality control of TSB used for microbiological examination of pharmaceutical products should meet the requirements of EP² A: aerobic incubation; ATCC is a trademark of American Type Culture Collection

12 - PERFORMANCES CHARACTERISTICS

Prior to release for sale, a representative sample of all lots of ready to use tubes of Tryptic Soy Broth and of the raw material used for the production of prepared tubes (dehydrated Tryptic Soy Broth REF 402155), is tested for productivity by comparing the results with a previously approved Reference Batch.

Productivity is tested by dilution to extinction method, by inoculating 1 mL of appropriate decimal dilutions of organisms in test tubes and incubating at 30-35°C or at 20-25°C for 18-24 hours or for 24-72 hours and recording the highest dilution showing growth in Reference Batch (Gr_{RB}) and in Test Batch (Gr_{RB}). Productivity is tested with the following strains: *B.subtilis* ATCC 6633, *C.albicans* ATCC 10231, *A.brasiliensis* ATCC 16404, *S.aureus* ATCC 6538, *P.aeruginosa* ATCC 9027, *E.coli* ATCC 8739, S.Typhimurium ATCC 14028. The productivity index Gr_{RB} - Gr_{TB} for each test strain shall be ≤ 1 .

13 - LIMITATIONS OF THE METHOD

- Tryptic Soy Broth is not suitable for the cultivation of fastidious microorganisms (e.g. *Haemophilus* or *Neisseria* spp.) and for the cultivation of strict anaerobes.
- · Sub-cultures onto suitable solid media are necessary for purification of the culture and to perform identification tests.
- This culture medium is intended as an aid in the diagnosis of infectious diseases; the interpretation of the results must be made considering the patient's clinical history, the origin of the sample and the results of other diagnostic tests.

14 - PRECAUTIONS AND WARNINGS

- This product is a qualitative *in vitro* diagnostic, for professional use only; it is to be used by adequately trained and qualified laboratory personnel, observing approved biohazard precautions and aseptic techniques.
- This product is not classified as dangerous according to current European legislation.
- This culture medium contains raw materials of animal origin. The ante and post mortem controls of the animals and those during the production and distribution cycle of the raw materials, cannot completely guarantee that the product doesn't contain any transmissible pathogen. Therefore, it is recommended that the product be treated as potentially infectious, and handled observing the usual specific precautions: do not ingest, inhale, or allow to come into contact with skin, eyes, mucous membranes. Download the TSE Statement from the website www.biolifeitaliana.it, describing the measures implemented by Biolife Italiana for the risk reduction linked to infectious animal diseases.
- All laboratory specimens should be considered infectious.
- The laboratory area must be controlled to avoid contaminants such as culture medium or microbial agents.
- Be careful when opening screw cap flasks to prevent injury due to breakage of glass.
- Ready-to-use flasks of Tryptic Soy Broth are subject to terminal sterilization by autoclaving.
- Sterilize all biohazard waste before disposal. Dispose the unused medium and the sterilized plates inoculated with samples or microbial strains in accordance with current local legislation.
- Do not use the culture medium as active ingredient for pharmaceutical preparations or as production material intended for human and animal consumption.
- The Certificates of Analysis and the Safety Data Sheet of the product are available on the website www.biolifeitaliana.it.
- The information provided in this document has been defined to the best of our knowledge and ability and represents a guideline for the proper use of the product but without obligation or liability. In all cases existing local laws, regulations and standard procedures must be observed for the examination of samples collected from human and animal organic districts, for environmental samples and for products intended for human or animal consumption. Our information does not relieve our customers from their responsibility for checking the suitability of our product for the intended purpose.

15 - STORAGE CONDITIONS AND SHELF LIFE

Upon receipt, store flasks in their original pack at 2-8°C away from direct light. If properly stored, the flasks may be used up to the expiration date. Do not use the flasks beyond this date. Flasks from opened secondary packages can be used up to the expiration date. Before use, check the closing and the integrity of the screw cap. Opened flasks must be used immediately for the inoculation or for the preparation of tubed medium. If the medium is sub-divided in tubes, the user is responsible of the process. Do not use flasks with signs of deterioration (e.g. microbial contamination, abnormal turbidity, precipitate, atypical colour).

16 - REFERENCES

- 1. MacFaddin JF. Media for Isolation-Cultivation-Identification-Maintenance of Medical Bacteria. Baltimore: Williams & Wilkins; 1985.
- 2. European Pharmacopoeia, current edition
- 3. CLSI (formerly NCCLS) Quality Control of Commercially Prepared Culture Media. Approved Standard, 3rd edition. M22 A3 vol. 24 n° 19, 2004.



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CE IVD

TABLE OF APPLICABLE SYMBOLS

REF or REF Catalogue number	LOT Batch code	IVD In vitro Diagnostic Medical Device	Manufacturer	Use by
Temperature limitation	Contents sufficient for <n> tests</n>	Consult Instructions for Use	For single use only	Fragile, handle with care

REVISION HISTORY

Version	Description of changes	Date
Revision 1	Updated layout and content in compliance with IVDR 2017/746	2021/09
Revision 2	Removal of obsolete classification	2023/04

Note: minor typographical, grammatical, and formatting changes are not included in the revision history.