

Instructions for Use

TRYPTIC SOY AGAR (TSA) MEDIA, USP, IRRADIATED

Cat. no. P520	Tryptic Soy Agar (TSA), with Lecithin and Tween® 80, USP, Irradiated, Triple Bagged, Contact Plate, Lok-Tight™, 15ml*	10 plates/bag
Cat. no. P520R	Tryptic Soy Agar (TSA), with Lecithin and Tween® 80, USP, Irradiated, Triple Bagged, Red-Tinted Contact Plate, Lok-Tight™, 15ml*	10 plates/bag
Cat. no. W520	Tryptic Soy Agar (TSA) with Lecithin and Tween® 80, USP, SterEM™, Irradiated, Triple Bagged, 15x100mm Plate, 26ml*	10 plates/bag
Cat. no. W540	Tryptic Soy Agar (TSA), USP, SterEM™, Irradiated, Triple Bagged, 15x100mm Plate, 34ml*	10 plates/bag
Cat. no. W540R	Tryptic Soy Agar, USP, SterEM™, Irradiated, Triple Bagged, Red-Tinted 15x100mm Plate, 34ml*	10 plates/bag
Cat. no. W570	Tryptic Soy Agar (TSA), USP, SterEM™, Irradiated, Triple Bagged, 15x100mm Plate, 26ml*	10 plates/bag
Cat. no. W570R	Tryptic Soy Agar (TSA), USP, SterEM™, Irradiated, Triple Bagged, Red-Tinted 15x100mm Plate, 26ml*	10 plates/bag
* A fourth sterile sample bag is included for packaging after the sample is collected.		

INTENDED USE

Hardy Diagnostics Tryptic Soy Agar (TSA), USP, Irradiated is recommended for use as a general growth medium for the detection and enumeration of microorganisms in the microbiological examination of non-sterile products. Tryptic Soy Agar (TSA) with Lecithin and Tween®, USP, Irradiated is recommended for use as a general growth medium for establishing microbiological trends, alerts, and action levels in biologically controlled environments.

The media are formulated according to the United States Pharmacopoeia (USP) <62> and meet the harmonized USP/EP/JP standards for the microbial examination of non-sterile products.

This product is not intended to be used for the diagnosis of human disease.

SUMMARY

The formulation of Tryptic Soy Agar, USP is prepared according to the *United States Pharmacopoeia* standard formula for Soybean-Casein Digest Agar Medium.⁽²⁾

Hardy Diagnostics Tryptic Soy Agar (TSA) contains digests of soybean meal and casein which provide amino acids and other nitrogenous compounds to promote the growth of a variety of microorganisms. Sodium chloride is added to maintain the osmotic equilibrium. Agar is the solidifying agent. Lecithin and Tween® 80 are added to neutralize the antimicrobial effects of disinfectants or cleaning solutions used on environmental surfaces.

The plates are triple bagged and sterilized by irradiation to promote a higher sterility assurance level.

FORMULA

Ingredients per liter of deionized water:*

TSA contains:	
Pancreatic Digest of Casein	15.0gm
Peptic Digest of Soybean Meal	5.0gm
Sodium Chloride	5.0gm
Agar	15.0gm
TSA with Lecithin & Tween [®] 80 also contains:	
Lecithin	0.7gm
Tween [®] (polysorbate) 80	5.0ml

Final pH 7.3 +/- 0.2 at 25°C.

* Adjusted and/or supplemented as required to meet performance criteria.

STORAGE AND SHELF LIFE

Storage: Store at 15-30°C., and store all media away from direct light. Media should not be used if there are any signs of contamination, deterioration, discoloration, or if the expiration date has passed. Product is light and temperature sensitive. Protect from freezing.

Do not use irradiated media if there is any damage to the packaging prior to use.

For irradiated media: Inspect each bag prior to opening and using the product.

The expiration date on the product label applies to the product in its intact packaging when stored as directed. The product may be used and tested up to the expiration date on the product label and incubated for the recommended incubation times as stated below.

Refer to the document "[Storage](#)" for more information.

PRECAUTIONS

This product may contain components of animal origin. Certified knowledge of the origin and/or sanitary state of the animals does not guarantee the absence of transmissible pathogenic agents. Therefore, it is recommended that these products be treated as potentially infectious, and handle observing the usual universal blood precautions. Do not ingest, inhale, or allow to come into contact with skin.

This product is for laboratory use only. It is to be used only by adequately trained and qualified laboratory personnel. Observe approved biohazard precautions and aseptic techniques. All laboratory specimens should be considered infectious and handled according to "standard precautions." Refer to the document "[Guidelines for Isolation Precautions](#)" from the Centers for Disease Control and Prevention.

For additional information regarding specific precautions for the prevention of the transmission of all infectious agents from laboratory instruments and materials, and for recommendations for the management of exposure to infectious disease, refer to CLSI document M29: *Protection of Laboratory Workers from Occupationally Acquired Infections*.

Sterilize all biohazard waste before disposal.

Refer to the document "[Precautions When Using Media](#)" for more information.

PROCEDURE

Before Use: For irradiated media, it is possible that variation in temperature and pressure during shipping and storage may cause condensation on the innermost bag surrounding the plates. If condensation of the packaging or plates is observed, remove the plates from the innermost packaging in a sterile environment and allow them to dry for 10-15 minutes before use.

The medium should be warmed to room temperature and the surface should be dry prior to use. To reduce the potential for cross-contamination, it is strongly suggested that appropriate gowning and glove procedures, designated aseptic processing areas, appropriate sporicidal disinfectants, and environmental monitoring procedures be strictly followed to reduce the likelihood of accidental contamination.⁽¹⁾

Method of Use: For environmental monitoring procedures, consult USP <1116> *Microbiological Evaluation of Clean Rooms and Other Controlled Environments*.⁽¹⁾ Incubate media using appropriate atmospheric, temperature, and duration conditions as outlined by the test method.⁽¹⁾ Count the number of colonies and report as the number of colony forming units (CFU).

Sedimentation Plate Method: Place the plate on a clean piece of paper and expose the agar by removing the lid. Do not invert the lid while removed to avoid exposure to falling sediment. Expose the agar for 15 minutes or longer, depending upon established procedures, and replace the lid. Incubate according to laboratory protocol.

Impact Air Sampling Method: Use the plate size specified for the impact air sampling unit. Remove the sampler head and place the plate, lid up, into the slot. Aseptically remove the lid and expose the agar; do not invert the lid while removed to avoid exposure to falling sediment. Place the sampler head back on the unit and turn the unit on; sample a specific volume of air according to laboratory procedure. After sampling, remove the sampler head, aseptically return the lid of the plate and remove the plate from the sampling unit; incubate per laboratory protocol.

INTERPRETATION OF RESULTS

Consult the appropriate reference for information regarding further testing and identification of microbial cultures.⁽¹⁻²⁾

Because of the inherent variability of environmental sampling methods, it is more useful to trend contamination recovery results rather than focus on the number of colonies recovered from a single sample. Action should be required when the contamination recovery rate trends above the recommended action levels for a significant time.

If action levels have been identified, a thorough investigation into the adequacy of personnel work practices, operational procedures, cleaning procedures and solutions, and air filtration efficiency within the processing area must be made. Once changes have been made, monitoring procedures must be repeated to determine if the changes made were effective. Documentation of all monitoring results, remedial action and follow-up monitoring must be maintained. Consult listed reference for more detailed information concerning plate count methods.⁽¹⁾

LIMITATIONS

It is recommended that biochemical, immunological, molecular, or mass spectrometry testing be performed on colonies from pure culture for complete identification of bacteria and/or fungi.

Avoid repeated and/or extreme variations in temperatures during storage, as this can cause the release of excessive moisture from the media in the bags and plates.

Accurate counting may be difficult with molds or spreading colonies.

Sampling challenges may occur with irregular, porous, rough or textured media surfaces.

Contact plate media are not recommended for sampling crevices or irregular surfaces.

Storage at 2-8°C may result in moisture build-up inside sealed packs.

Refer to the document "[Limitations of Procedures and Warranty](#)" for more information.

MATERIALS REQUIRED BUT NOT PROVIDED

Standard microbiological supplies and equipment such as loops, other culture media, swabs, applicator sticks, incinerators, and incubators, etc., as well as serological and biochemical reagents, are not provided.

QUALITY CONTROL

Hardy Diagnostics tests each lot of commercially manufactured media using appropriate quality control microorganisms and quality specifications as outlined on the Certificate of Analysis (CofA) and the CLSI document M22-A3 *Quality Assurance for Commercially Prepared Microbiological Culture Media*. The following microorganisms are routinely used for testing at Hardy Diagnostics:

Test Organisms**	Inoculation Method*	Incubation			Results
		Time	Temperature	Atmosphere	
<i>Staphylococcus aureus</i> ATCC® 6538	J	1-3 days	30-35°C	Aerobic	Growth
<i>Candida albicans</i> ATCC® 10231	J	1-5 days	30-35°C	Aerobic	Growth
<i>Bacillus spizizenii</i> ATCC® 6633	J	1-3 days	30-35°C	Aerobic	Growth
<i>Aspergillus brasiliensis</i> ATCC® 16404	J	1-5 days	30-35°C	Aerobic	Growth
<i>Pseudomonas aeruginosa</i> ATCC® 9027	J	1-3 days	30-35°C	Aerobic	Growth
Also included for Cat. nos. P520 and P520R:					
<i>Staphylococcus epidermidis</i> ATCC® 12228	A	1-3 days	30-35°C	Aerobic	Growth
<i>Clostridium sporogenes</i> ATCC® 19404	A	1-3 days	30-35°C	Anaerobic	Growth

Representative samples from each lot of irradiated media are held for seven days to confirm the media meet the validated sterilization process sterility assurance level (SAL) of 10⁻⁶ following ANSI/AAMI/ISO 11137.

* Refer to the document "[Inoculation Procedures for Media QC](#)" for more information.

** Tested in accordance with USP <61> and <62>. ⁽²⁾

USER QUALITY CONTROL

End users of commercially prepared culture media should perform QC testing in accordance with applicable government regulatory agencies, and in compliance with accreditation requirements. Hardy Diagnostics recommends end users check for signs of contamination and deterioration and, if dictated by laboratory quality control procedures or regulation, perform quality control testing to demonstrate growth or a positive reaction and to demonstrate inhibition or a negative reaction, if applicable. Hardy Diagnostics quality control testing is documented on the certificate of analysis (CofA) available from Hardy Diagnostics [Certificate of Analysis](#) website. Also refer to the document "[Finished Product Quality Control Procedures](#)," and the CLSI document M22-A3 *Quality Assurance for Commercially Prepared*

Microbiological Culture Media for more information on the appropriate QC procedures. See the references below.

PHYSICAL APPEARANCE

Tryptic Soy Agar (TSA) Agar should appear translucent, and light amber in color.

Tryptic Soy Agar (TSA) Agar with Lecithin and Tween® 80 should appear clear to slightly hazy and light amber in color.



Uninoculated plate of Tryptic Soy Agar, Irradiated.

REFERENCES

1. American Public Health Association. 2012. *Standard Methods for the Examination of Water and Wastewater*, 22nd ed. APHA, Washington, D.C.
2. The Official Compendia of Standards. *USP-NF*. United States Pharmacopeial Convention, Rockville, MD.
3. *Quality Assurance for Commercially Prepared Microbiological Culture Media*, M22. Clinical and Laboratory Standards Institute (CLSI - formerly NCCLS), Wayne, PA.

ATCC is a registered trademark of the American Type Culture Collection.

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[Ordering Information](#)

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management system is certified to ISO 13485.

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