# Introduction

#### Intended Use

 $InTray^{TM}$  SAB-FungID $^{TM}$  w/ CC contains Sabouraud's dextrose agar with chloramphenical and cycloheximide, a selective medium used to aid in the detection of dermatophyte fungi from clinical specimens with mixed microbiota.

# Description and Principle

InTray SAB-FungID w/ CC is a dynamic system with built-in components and features that are designed for user compatibility and ease of dermatophyte fungi detection (i.e., 10x objective microscopy direct from the tray). The medium allows for the growth and observation of distinct colony morphology and color (i.e., pigments) of dermatophyte fungi, while inhibiting most grampositive bacteria, gram-negative bacteria, yeast and saprophytic fungi. Dermatophytes are fungi in the genera Microsporum, Trichophyton and Epidermophyton. They are capable of metabolizing keratin found in skin, hair and nails of living hosts. The fungi characteristically may invade the cutaneous tissue of the living host but rarely penetrate the subcutaneous tissue. Tinea and ringworm are two terms commonly used to describe dermatophytes.

# Reagents and Appearance

SAB-FungID w/ CC appears transparent with a light amber hue and contains peptic/casein digest, dextrose, chloramphenicol (0.050 g/L) and cycloheximide (0.40 g/L) with a final pH of 5.6 ± 0.2 at 25°C

# Precautions, Safety and Disposal

For In Vitro Diagnostic Use. For professional use only.

Read the Safety Data Sheets (SDSs) and follow the handling instructions. Wear appropriate protective eyewear, clothing and aloves.

Once the tray has been inoculated and resealed, re-open only in a biological safety cabinet. Because of the potential for containing infectious materials, the tray must be destroyed by autoclaving at 121°C for 20 minutes.

# Storage

On receipt, store trays at 2-25°C in the dark. Avoid freezing or prolonged storage at temperatures greater than 40°C. Do not use trays if the medium shows signs of deterioration or contamination.

# Shelf Life

InTray SAB-FungID w/ CC expires 12 months from date of manufacture.

# Procedure

# Key Notes Regarding Specimen Collection

Specimen collection poses a major uncertainty in using this device.

NAILS - Often collecting viable material from infected nails is difficult because the living organisms are well under the nail itself. For best results, cut nails into small pieces.

HAIR - Samples should be grasped at the uninfected end and several (3-6) small pieces, about 2 cm long, should be cut from the infected portion for inoculation onto the medium.

**SKIN** - Scrapings should be taken with an inoculation tool that has been moistened with the medium or a sharp blade from the outer ridge of an active lesion. Vesicular fluid is unacceptable for dermatophyte culture. If the infected area is vesiculated, skin scrapings should be taken from the surface.

#### **Prepare Sample:**

Use aseptic technique during specimen collection and handling. Remove any soap residue from the sampling area. Clean the area with 70% alcohol and permit to air dry.

# **Collect Sample:**

InTray SAB-FungID w/ CC is designed for culturing hair, skin and nail samples (i.e., cuttings/scrapings). All specimens should be handled according to CDC infectious materials isolation guidelines; cdc.gov/infectioncontrol/guidelines/isolation

#### Materials Provided

InTray SAB-FungID w/ CC test(s)

#### Materials Required but Not Provided

- Sterile inoculation tool (e.g., cotton swab/forceps/scalpel blade)
- Laboratory incubator capable of incubation at 25-30°C

#### 1 Prepare InTray



Allow tray(s) to warm to 18-25°C before use. Immediately label the tray with patient/sample information and date. Pull back the lower right corner adjacent to the clear window of the tray label until the protective seal is completely visible.

Remove the seal by pulling the tab. Discard the seal.

DO NOT REMOVE OR ALTER THE WHITE FILTER STRIP OVER THE VENT HOLE!

#### 2 Inoculate Sample



Inoculate the specimen on the center surface of the medium. A sterile inoculating loop that has been moistened by touching the surface of the medium may be used for inoculation of solids or scrapings.

Re-seal all around the tray to ensure a complete seal by pressing the edges of the label against the plastic tray. DO NOT COVER THE VIEWING WINDOW. Complete re-seal prevents dehydration!

#### Incubation

Incubate inoculated trays in a dark environment for up to 21 days at 25-30°C. Observe the trays daily through the clear viewing window.

# **Quality Control**

This product has been tested and meets the CLSI (formerly NCCLS) Approved Standard for commercially prepared media (M22-A3). At the time of manufacture, quality control testing is performed on each lot of the InTray SAB-FungID w/ CC. The ability of the media to support growth and demonstrate expected biochemical reactions and morphology is verified by lot.

# Strains for QC Testing SAB-FungID w/ CC

Organism	ATCC <sup>®</sup>	Expected Result
T. mentagrophytes	9533	Growth
T. rubrum	28188	Growth
C. albicans	60193	Partial Inhibition
A brasiliensis	16404	Significant Inhibition
S. aureus	25923	Significant Inhibition
E. coli	25922	Significant Inhibition

# Reading the Results

#### Evaluation

Observe the medium for fungal colony growth, texture, morphology and pigmentation characteristic of dermatophytes and saprophytes. Without opening the InTray SAB-FunalD w/ CC, place the tray under a 10x objective microscope lens to view distinct funaal structures (i.e., hyphae, micro/macro-conidia). See the Quick Identification Chart below for reference. Trays to be used with 10x objective ONLY!

Positives: If, within 1-21 days, the medium shows characteristic dermatophyte fungi colony growth, texture, morphology, pigmentation and microscopic structure, the SAB-FungID w/ CC is presumptive positive.

Negatives: Trays that show no colony growth after 21 days from the date of inoculation are presumptive negative.

#### Dermatophyte Identification

This is a selection of commonly encountered organisms. Please consult the references listed below, as well as other standard mycology and microbiology references.



#### Trichophyton rubrum

Colony appearance: white, pink Reverse appearance: yellow to wine red

Septate hyphae. Macroconidia: (4-6 x 15-30 µm) abundant, rare or absent, however can be long, narrow, thin-walled, parallel sides, 2-8 cells, may form on the ends singly or in groups. Microconidia: (2-3 x 3-5 µm) lateral, teardrop shaped, form on macroconidia.



#### Trichophyton mentagrophytes

Colony appearance: white to yellow Reverse appearance: reddish brown

Septate hyphae. Macroconidia: (4-8 x 20-50 µm) occasionally present, cigar-shaped, thin-walled, narrow attachments to septate hyphae, 1-6 cells, found in young cultures 5-10 days old. Microconidia: usually present in powdery cultures, very round, clustered on branched conidiophores; in fluffy cultures, smaller, fewer, teardrop shaped and easily confused with those of T. rubrum.



#### **Epidermophyton floccosum**

Colony appearance: green, yellow edges

Reverse appearance: vellowish brown

Septate hyphae. Macroconidia: (7-12 x 20-40 µm) smooth, thick and thinwalled, club shaped with rounded ends, two to six cells, singly or clusters. Microconidia: none.

# Saprophyte (Contaminants) Identification



Alternaria sp. Hyphae are septate and dark. Conidiophores are septate, variable in length and sometimes branched. Macroconidia are large (7-10 by 23-24 µm), brown, have both transverse and longitudinal stations, and found singly or in chains. They are usually round at the end nearest the conidiophore, producing a club-like shape.



Aspergillus sp. Microsco pic morphology – Septate hypae (2.5-8 µm in diameter); unbranched condidiophore arises from a specialized foot cell. The conidiophore is enlarged at the tip, forming a swollen vesicle completely or partially covered with flask-shaped phialides. The pialides produce chains of mostly round, sometimes rough conida (2-5 µm in diameter)



Penicillium sp. Microscopic morphology – Septate hyphae (1.5-5 μm in diameter) with branched conidiophores that have secondary branches known at metulae. On the metulae, are flask-shaped phialides that bear unbranched chains of smooth or rough conidia (2.5-5 µm in diameter). The entire structure forms the characteristic "penicillus" or "brush" appearance.

#### Limitations

If a fungal infection is strongly suspected and the culture test result is negative, it may be appropriate to retest giving more care to specimen collection.

- Optimally, simultaneously inoculate samples on both InTray DM-FungID (Cat. Nos. 11-593-001, 11-593-002) and InTray SAB-FungID (Cat. Nos. 11-263-001, 11-263-002) without antimicrobials (e.g., for some pathogenic fungi inhibited by antimicrobials)
- Transfer of fungal growth to slide examination: Stained slide preparations can be made as appropriate based on 10x objective examination of SAB-Fun-gID w/ CC. Potato Dextrose Agar can be used to induce sporulation.
- Final culture identification is to be made by laboratory professionals only and may require further biochemical/culture testing.

InTray SAB-FungID w/ CC is an agar medium that is susceptible to condensation collection within the inner seal, especially when stored at low temperatures and/or having been exposed to extreme temperature fluctuations. If moisture is visible on the surface of the InTrays, dry them (with the seal removed and InTray label in a position allowing for air flow) under a BSL-2 cabinet just prior to inoculation. There should be no visible droplets of moisture on the surface of the agar when they are inoculated. The surface of the dried medium should be smooth and should not show signs (webbed ribbing pattern on the agar surface) of desiccation.

#### References

- Sabouraud R (1892), Ann Dermatol, Syphil, 3:1061,
- Tille, et al. (2014) Bailey & Scott's Diagnostic Microbiology, Elsevier: p. 450.
- Kwon-Chung, KJ and Bennett, JE (1992) Medical Mycology, Lea and Febiger:
- Murray, PR, Baron, ET, Pfaller, MA, Tenover, FC, Yolkem, RH, (1995) Manual of Clinical Micro-biology 6th ed., American Society for Microbiology: Washington, D.C., pp. 709-722.
- Larone, DH (1995) Medically Important Fungi: A Guide to Identification, 2nd ed., American Society for Microbiology: Washington, D.C.

Symbol glossary: biomeddiagnostics.com/l/symbol-glossary

# Document Revision History

Rev. C, Aug 2025

Removed QR Code, updated manufactured by, logo, and company address. Replaced R with TM. DM-FungID catalog numbers were corrected to 11-593-001, 11-593-002, PDA-FungID was discontinued and removed from the Limitations section.



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# InTray™ SAB-FungID™ w/ CC

Sabouraud's Dextrose Agar with Cycloheximide and Chloramphenicol



11-283-001

11-283-002



Not available in all countries; please inquire.

A SELECTIVE CULTURE SYSTEM TO AID DETECTION OF Dermatophyte fungi

For In Vitro Diagnostic Use



