An Implementation Strategy for the Use of Chromogenic Media in the Rapid, Presumptive Identification of Candida Species

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Abstract: The purpose of this study was to devise an algorithm utilizing chromogenic media for the detection and presumptive identification of Candida species in a clinical microbiology laboratory. In order to select one chromogenic medium, a limited evaluation of 3 commercially available yeast chromogenic media: BBL CHROMagar Candida, Bio-Rad CandiSelect 4, and Oxoid OCCA, was conducted with 12 C. albicans, 8 C. glabrata, 10 C. tropicalis, and 9 C. krusei. The presumptive species identification after 48 hours of incubation on BBL and Bio-Rad media were comparable, with more than 80% of the Candida species correctly identified, and both were superior to Oxoid. An algorithm based on use of chromogenic media for the rapid presumptive identification of C. albicans, C. tropicalis, C. krusei, and C. glabrata was developed for routine use in the microbiology laboratory. A presumptive identification is available within 24 to 48 hours utilizing this algorithm.

Keywords: Chromagar, chromogenic media, candida.

INTRODUCTION

Yeasts are a common cause of nosocomial infection in hospitalized patients. Candida species are reported as the fourth most commonly isolated pathogen from blood cultures in hospitalized patients [1, 2]. The most common Candida species causing bloodstream infections include C. albicans, C. glabrata, C. tropicalis, and C. parapsilosis [3]. Rapid identification of these non-albicans species is critical to patient management as C. krusei and C. glabrata exhibit decreased susceptibility to fluconazole [4], which is frequently selected for empiric therapy.

Conventional yeast identification includes microscopic morphological identification and biochemical studies, which require technical expertise and may take three to four days. Chromogenic media have recently been developed to facilitate simple and rapid yeast identification for the more commonly isolated species. Current chromogenic yeast media isolate and differentiate C. albicans, C. glabrata, C. krusei, and C. tropicalis based on colonial morphology and color production [5-10]. The identification is considered presumptive, particularly for C. glabrata, C. krusei, and C. tropicalis, and requires confirmatory testing by standard yeast identification methods. The use of chromogenic media as a primary medium for Candida species detection from clinical specimens is supported by the high sensitivity and specificity reported in the current literature [5, 9, 10]. Chromogenic media are advantageous as they also enable the detection of mixed yeast cultures, which may not be detected on standard media due to the similar colonial morphology of different yeast species [11].

Currently, there are only 3 commercially available chromogenic yeast media: BBL CHROMagar Candida (CHROMagar; Becton Dickinson Canada Inc.), Bio-Rad CandiSelect 4 (CandiSelect; Bio-Rad), and Oxoid OCCA (OCCA; Oxoid Ltd.). Comparative studies have been conducted with CHROMagar and OCCA [12] as well as CHROMagar and CandiSelect [10, 13]; however, a comparison of all three media has not been reported previously. In order to select the best chromogenic medium for use in the routine identification of yeast in a clinical microbiology laboratory, a limited evaluation of the 3 commercially available chromogenic yeast media was conducted.

The purpose of this study was to devise an algorithm to optimize the use of chromogenic media for the detection and presumptive identification of Candida species in a clinical microbiology laboratory. New microbiological products are frequently developed and made available to laboratories; however, the best way to incorporate these products into current laboratory processes is not always evident. This study was designed to address this question for yeast chromogenic media.

MATERIALS AND METHODOLOGY

Candida Strains

Thirty-nine Candida strains including 12 C. albicans, 8 C. glabrata, 10 C. tropicalis, and 9 C. krusei were evaluated in this study. All strains were selected from clinical bloodstream Candida isolates from patients in Toronto,
which are stored in the microbiology laboratory at the Hospital for Sick Children, Toronto, Canada. Isolates were identified by conventional means, i.e. germ tube test, morphology on cornmeal tween 80 agar and API 20C AUX.

Candida Identification on Chromogenic Media

Each Candida strain was subcultured twice onto blood agar from the stock culture prior to inoculation of the 3 chromogenic media (CHROMagar, CandiSelect, and OCCA). Standard inocula were prepared such that 30 – 100 colonies were inoculated onto each of the 3 chromogenic media. In accordance with the manufacturers’ instructions, the CHROMagar and CandiSelect were incubated aerobically at 35°C and the OCCA was incubated aerobically at 30°C. Color development and colonial morphology were evaluated daily for 2 days and a presumptive identification of each isolate was recorded. The color and colonial morphology used to identify the organisms is detailed in Table 1. The selection of isolates and the evaluation of colony color and morphology on the chromogenic media were conducted by separate individuals to complete the study in a blinded manner.

The identification of 3 chlamydospore negative C. albicans was confirmed by DNA sequencing. Sequencing was conducted using ITS3 and ITS4 fungal primers previously described by Chen et al. [14].

A retrospective review of all blood cultures and sterile site specimens from 2007 and 2008 that included a yeast workup was conducted to validate the developed algorithm.

RESULTS AND DISCUSSION

Accuracy of the Chromogenic Media

All 3 media are able to identify C. albicans, C. krusei, and C. tropicalis. CandiSelect is marketed with an additional indication for the presumptive identification of C. glabrata. Although CHROMagar and OCCA are not marketed with an indication for C. glabrata, C. glabrata strains have a characteristic appearance on both media that is different from the three species for which they have been validated (see Table 1). As such, we evaluated their utility in making a presumptive diagnosis of C. glabrata infection. After 24 hours, CHROMagar, CandiSelect, and OCCA correctly identified 36%, 49%, and 3% of the Candida species, respectively. By 48 hours, 85%, 85%, and 62% of the Candida species were correctly identified by CHROMagar, CandiSelect, and OCCA, respectively. The overall percent identification of Candida species provided by each media after 24 and 48 hours of incubation is presented in Fig. (1). As demonstrated in Fig. (2), CHROMagar and CandiSelect correctly identified all C. albicans and C. krusei by 48 hours whereas OCCA only identified 92% of the C. albicans and 56% of the C. krusei. At 48 hours, CHROMagar, CandiSelect, and OCCA identified 50%, 60%, and 40% of the C. tropicalis. CHROMagar, CandiSelect, and OCCA correctly identified 88%, 75%, and 50% of C. glabrata strains by 48 hours, respectively.

Three chlamydospore negative C. albicans were correctly identified as C. albicans on all three chromogenic media after 24 to 48 hours of incubation. Sequencing of the rRNA genes confirmed the C. albicans identification.

A small number of discordant identifications were observed on the three media. The discordant identifications were as follows: CHROMagar, 1 C. tropicalis identified as C. glabrata; CandiSelect, 1 C. glabrata identified as C. tropicalis; OCCA, 1 C. albicans identified as C. tropicalis, 1 C. tropicalis identified as C. albicans, and 1 C. tropicalis identified as C. glabrata.

Selection of a Chromogenic Medium for Routine Use

The amount of time required for identification and the accuracy of the species identification were comparable for CHROMagar and CandiSelect. As shown in Fig. (1), CHROMagar (98%) provided a greater percentage of correct or possible identifications at 48 hours than either CandiSelect (85%) or OCCA (80%). However, CHROMagar and CandiSelect were comparable in their ability to identify the 4 Candida species: C. albicans, C. glabrata, C. krusei, and C. tropicalis (Fig. 2). Similarly, Sendid et al. demonstrated that an equivalent proportion of C. albicans, C. glabrata, C. krusei, and C. tropicalis isolates was identified by CHROMagar and CandiSelect [13]. OCCA was found to be inferior to CHROMagar and CandiSelect at 24 and 48 hours with respect to the speed of the presumptive identification and the proportion of correct identifications, as previously reported [12]. A 72 hour incubation is recommended by the manufacturer of OCCA; however, we required a presumptive identification within 48 hours for the development of a rapid identification schema. Accordingly, the results after 72 hours of incubation were not considered in our study. The observation that CHROMagar was superior to OCCA supports the current literature [12].

Although CandiSelect is advantageous as it provides a presumptive identification for C. glabrata, the distinct color development of the four Candida species on CHROMagar was easier for the technologists to learn and may result in

<table>
<thead>
<tr>
<th>Media</th>
<th>C. albicans</th>
<th>C. glabrata</th>
<th>C. tropicalis</th>
<th>C. krusei</th>
</tr>
</thead>
<tbody>
<tr>
<td>BBL CHROMagar Candida</td>
<td>Light to medium green</td>
<td>Small pink</td>
<td>Dark blue to metallic blue</td>
<td>Light mauve to mauve with whitish border, rough</td>
</tr>
<tr>
<td>Bio-Rad CandiSelect 4</td>
<td>Pink to purple</td>
<td>Pale turquoise centre with white periphery</td>
<td>Intense turquoise</td>
<td>Turquoise-blue, rough</td>
</tr>
<tr>
<td>Oxoid OCCA</td>
<td>Green</td>
<td>Beige/yellow-brown</td>
<td>Dark blue</td>
<td>Pink-brown, dry</td>
</tr>
</tbody>
</table>

Table 1. Colonial Morphology and Color Development on BBL CHROMagar Candida, Bio-Rad CandiSelect 4, and Oxoid OCCA for the Presumptive Identification of C. albicans, C. glabrata, C. tropicalis, and C. krusei
Implementation of Candida Chromogenic Media

The rationale for the ultimate selection of CHROMagar in our laboratory included its demonstrated accuracy in this study, the wide acceptance of CHROMagar as the medium of choice based on favorable reports in the literature [5, 6, 8, 9, 11, 12], and the preference by the laboratory technologists due to its distinct color profile.

greater consistency of yeast identification. The rationale for the ultimate selection of CHROMagar in our laboratory included its demonstrated accuracy in this study, the wide acceptance of CHROMagar as the medium of choice based on favorable reports in the literature [5, 6, 8, 9, 11, 12], and the preference by the laboratory technologists due to its distinct color profile.

Fig. (1). Overall Percent Identification of Candida species Provided by BBL CHROMagar Candida, Bio-Rad CandiSelect 4, and Oxoid OCCA after 24 and 48 Hours of Incubation.

Correct identification, identification matches database; Possible identification, color development indicative, but inconclusive for identification (e.g. Mauve colony on CHROMagar indicative of C. krusei, but inconclusive without the white border); discordant identification, identification discordant with database; and no identification, no species-specific color development.

Fig. (2). Percent of C. albicans, C. glabrata, C. krusei, and C. tropicalis Correctly Identified by BBL CHROMagar Candida, Bio-Rad CandiSelect 4, and Oxoid OCCA after 24 and 48 Hours of Incubation.
Targeted Implementation of BBL CHROMagar Candida

In order to utilize the CHROMagar medium in a cost-effective manner in our laboratory, the medium is included in the routine investigation of specimens for which complete yeast identification is warranted. In particular, the medium is utilized as a primary isolation medium for sterile site cultures, including blood, CSF, sterile fluids (abdominal, joint, pericardial, peritoneal), and biopsies.

The identification schema developed for the implementation of CHROMagar for routine yeast identification from sterile sites is presented in Fig. (3). This schema enables the rapid presumptive identification of C. albicans, C. tropicalis, and C. krusei. As CHROMagar is not marketed for the presumptive identification of C. glabrata, a rapid confirmatory test is suggested rather than a presumptive identification based on colony colour. In our schema, a presumptive identification is made based on the color and colonial morphology observed on the chromogenic media as soon as it is possible, following 24 or 48 hours of incubation. At that point, an interim report is provided to the clinician with the presumptive yeast identification and a comment that confirmation will be forthcoming.

A final identification requires confirmatory testing. As per our schema, the additional tests differ based on the presumptive identification and include morphology on direct microscopic examination, morphology on cornmeal tween 80 agar, detection of urease, API 20C AUX results, and RAT (rapid assimilation of trehalose) test (REMEL Inc., Lenexa, KS). The identification of C. glabrata from CHROMagar with our schema relies on the presence of typical macroscopic morphology (small pink), typical microscopic morphology (small, circular yeast cells) [15], and a positive RAT test. The RAT test has been shown to be highly sensitive and specific in the confirmation of C. glabrata identification from chromogenic media [8]. The confirmatory tests for C. glabrata can be completed on the same day as the pink colonies are observed, which permits a rapid final identification on the same day that a presumptive identification would be available. Isolates with discordant presumptive identifications are subjected to additional testing: Cornmeal Tween 80 agar from presumptive C. tropicalis and C. krusei and Cornmeal Tween 80 agar and API20C AUX for presumptive C. glabrata.

C. dubliniensis cannot be definitively distinguished from C. albicans on chromogenic media [6]; thus, green colonies...
are reported as *C. albicans/C. dubliniensis*. Most *Candida* yeasts other than *C. albicans, C. glabrata, C. krusei, and C. tropicalis* appear to be shades of pink, lavender or ivory, which are distinguishable from the four yeasts for which a presumptive identification is possible. However, several uncommon strains of *Candida*, including *C. inconspicua, C. firmeata*, and some strains of *C. norvegensis* and *C. lipolytica*, may have the rough colonial morphology, pink/purple hue and pale border characteristic of *C. krusei*, making them indistinguishable morphologically [6]. Thus, confirmation of the identify of *C. krusei* is required, in this case with an API 20 AUX.

The accuracy of the chromogenic media also enables it to be relied upon for the identification of repeat positive specimens. *C. albicans/C. dubliniensis, C. glabrata, C. krusei or C. tropicalis* that are re-isolated from the same site within two weeks of a complete identification are referred to the previous identification based on the color and colonial morphology observed on the chromogenic medium without further testing.

**POST-IMPLEMENTATION REVIEW OF CANDIDA SPP. IDENTIFICATIONS WITH THE RAPID, PRESUMPTIVE IDENTIFICATION ALGORITHM UTILIZING BBL CHROMOMAG CANDIDA MEDIA**

Since 2007, 102 blood culture and 26 sterile site specimens were processed with the yeast chromogenic media following the algorithm detailed in Fig. (3). There were 78 blood cultures from 21 patients and 21 sterile site isolates from 6 patients available for full analysis upon retrospective review. Seventy-one *C. albicans, 7 C. glabrata, 1 C. krusei, and 19 C. parapsilosis* were recovered from these specimens. The presumptive identifications provided by the yeast chromogenic media were confirmed for all isolates of *C. glabrata, C. krusei* and 98.6% of the *C. albicans*. One culture with *C. albicans* grew as 2 morphotypes; one morphotype originally grew as a white colony on the yeast chromogenic media. The average time to presumptive identification for *C. albicans, C. glabrata, and C. krusei* was 1.5 days (range: 1-4 days), 3 days (range: 2-4 days), and 1 day, respectively. The average time to final identification for *C. albicans, C. glabrata, C. krusei, and C. parapsilosis* was 2.3 days (range: 1-9 days), 3.7 days (range: 3-4 days), 5 days, and 4.4 days (range: 3-6 days), respectively.

**CONCLUSION**

Significant work evaluating the sensitivity and specificity of yeast chromogenic media has been reported previously [5, 6, 8-10, 12, 13]; however, none of them has suggested implementation schemes for routine use in hospital microbiology laboratories. This study is unique as it details a simple and practical method for incorporating chromogenic media into routine yeast identification from specimens for which complete yeast identification is warranted. In the retrospective review following the implementation of the algorithm, it was shown to provide accurate identifications to clinicians by at least one day earlier than the final identifications completed by traditional methods.

The ability to refer specimen identifications to previous specimen results on the basis of a single test improves laboratory turn-around-time and cost-efficiency. Another advantage of the chromogenic media is its ability to identify mixed cultures [11]. The identification of mixed cultures with *C. glabrata or C. krusei* may impact treatment decisions due to their intrinsic fluconazole resistance. The potential to identify mixed *Candida* cultures, the rapid presumptive species identification, and the improved turn-around-time of repeat positive specimens will significantly improve laboratory processes and ultimately patient care.

**ACKNOWLEDGEMENTS**

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**REFERENCES**


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