ROLE OF THE CONCENTRATION PROCESS IN THE RECOVERY OF CANDIDA ALBICANS FROM BLOOD

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ABSTRACT

The lysis-centrifugation system (Isolator™) is recognized as the standard method for recovery of Candida spp. from blood. In this study, the effect of the concentration process of this system was compared with conventional methods of blood culture using liquid and biphasic media. The tests were performed in vitro using Candida albicans in counts of 100, 10, 1, and 0.5 cells/ml of blood. Cultures onto chocolate agar were performed with the sediment obtained after centrifugation of Isolator™ tube and with liquid and biphasic media after their incubation for 24 h at 35°C. Gram stain prepared from the conventional methods were also evaluated in the first 24 h. It was possible to detect Candida albicans in blood, regardless both the number of cells or methodology. As blastoconidia were observed in Gram stains at the same time that growth was noted, time for diagnosis was also not different for the compared methods. Therefore, we suggest that the process of concentration is not the single important factor responsible for the recovery rates of Candida albicans from blood by the Isolator™ system.

Key words: candidemia, blood culture, lysis-centrifugation system

INTRODUCTION

The incidence of fungemia has increased considerably in hospitalized patients (1, 21, 28). Candida spp. are the most commonly fungi recovered from blood, and more than 50% of the episodes of candidemia are due to Candida albicans (25, 31). In fact, deep fungal infection due to Candida spp. is responsible for an excess of hospital stay and for an overall mortality rate that may range from 38% to 75% (32, 33). Patients using inadequate antimicrobial treatment had a mortality rate higher than those receiving appropriate antifungal therapy (24, 30). Considering that patients with fungemia rarely present characteristic clinical manifestations, the diagnosis relies upon the findings of the microbiology laboratory, in particular, observation and isolation of fungi from blood. Although an increase in life-threatening invasive yeast infection has been observed, many of these infections either remain undetected or are detected too late to beneficiate patient management (4, 21, 22, 23). The number of fungal cells in blood is generally small, and this poses the major problem to obtain a positive result (27, 4).

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Recovery of *Candida albicans* from blood

The conventional, visually monitored, broth-based blood culture consists of culturing blood in broth (liquid) or biphasic (liquid and solid) media. Their advantages include relative inexpensive supplies, the convenience of collecting specimens and most importantly, the benefits that can derive from the use of a technology that existed for so many years (12, 14). Therefore it is still largely used, specially in small laboratories.

The critical importance of sensitivity and speed of processing blood cultures for proper diagnosis and management of patients stimulated continuous efforts to develop more sensitive blood culture methods. A great deal of work has been done to define optimal media, blood-to-medium ratios, types of anticoagulants and other routine procedures (12).

The development of a lysis-centrifugation system (Isolator; Wampole Laboratories, Cranbury, NJ) improved the recovery rates of fungi from blood. In this system blood cells are lysed to liberate microorganisms from phagocytes and blood is concentrated by centrifugation to prepare an inoculum that is plated onto agar plates. Most clinical comparisons of Isolator™ with broth-based blood culture systems showed higher sensitivity (3, 5, 11, 13, 18, 22), and earlier results using the Isolator™ system (8, 13). In a quantitative study, Kiehn et al. (15) reported advantage of Isolator™ versus broth culture method only if the number of fungal cells present in blood was less than 10 /ml.

Although lysis-centrifugation system has advantages, processing of specimens is labor-intensive and expensive (18), mainly due to the centrifugation process rather than the lysis. Another disadvantage of Isolator™ is the high propensity for contamination due to manipulation of samples (9, 12).

Reports of Murray et al. (19) and Zierdt (35), comparing the of Isolator™ system with conventional broth method plus lytic agent, indicate that the higher performance Isolator™ is due to the release of intracellular microorganisms, so that broth supplemented with lytic agent saponin has been introduced in automated systems (13, 17).

The aim of this work is to evaluate the influence of blood concentration, using the Isolator™ system, in the recovery of *Candida albicans*.

**MATERIALS AND METHODS**

**Methods of blood culture and culture media**

The concentration process was done using the Isolator™ system. Blood was concentrated by centrifugation using a fixed angle rotor, and the remaining pellet was directly plated onto solid media. Tubes of Isolator™ contain saponin to liberate microorganisms from white blood cells and sodium polyanetholsulfonate (SPS) to prevent clotting. Although *in vitro* phagocytosis does not occur easily (16), in this study blood was mixed with the chemical components of the tube before inoculation of *Candida albicans*, in order to allow the action of SPS and saponin, avoiding “in vitro” phagocytosis (12). Therefore, the lysis interference was eliminated in all samples, assenting to determine the effect of the concentration alone.

The conventional methods were represented by biphasic bottles and by 45 ml broth bottles, containing brain heart infusion (BHI - DIFCO Laboratories, Detroit, Michigan), supplemented with 0.05% SPS.

Chocolate agar plates used to subculture *Candida albicans* were prepared with BHI agar (DIFCO Laboratories, Detroit, Michigan).

**Blood**

A total of 30 ml of blood was collected from healthy volunteers after appropriate desinfection of the venipuncture site. From each sample of blood a volume of 9 ml was immediately inoculated into one Isolator™ tube, one biphasic bottle and one broth bottle.

**Candida albicans**

*Candida albicans* isolated from a patient with candidemia, identified by API 20C (BIOMÉRIEUX VITEK, Missouri, USA), was maintained on agar slant at 22°C (20). The inoculation sample was prepared as described elsewhere (2, 6, 16). Briefly, the sample was inoculated onto a chocolate agar plate and incubated for 24 h at 35°C. Five to ten colonies were suspended in sterile saline and centrifuged for 10 minutes at 1,500 rpm. The pellet was washed twice and a suspension of turbidity equivalent to 1000 cells/ml was prepared. The number of cells in this suspension was confirmed by counting the cells in Neubauer’s chamber (7).

**Inoculation and culturing procedures**

The original suspension of 1000 cells of *Candida albicans/ml* was serially diluted to obtain an inoculum ten fold more concentrated than the final tests, i.e., 100, 10 and 5 cells/ml. One ml of these suspensions was inoculated into each of the bottles or tubes, which
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had been previously inoculated with 9 ml of blood. Accordingly, to perform tests with 100 cells of Candida/ml of blood, 1 ml of the suspension containing 1000 cells/ml was used; to obtain tests containing 10 cells of Candida/ml of blood the inocula with 100 cells/ml was used, and so on.

After the inoculation of blood and suspensions of Candida albicans, the samples of conventional methods were incubated at 35°C for 24 h. Subcultures were made as recommended (12): an amount was drain to prepare a Gram smear and a volume of 0.1 ml was plated onto chocolate agar plates.

The Isolator™ tubes were processed as described in the instructions of the manufacturer (29). Finally, 0.1 ml of the pellet was plated onto chocolate agar plates.

Chocolate agar plates were incubated at 35°C for 24 h. In view of the excessive number of colonies, it was impossible to perform individual colony count (Fig. 1). Therefore, magnitude of growth was estimated as degrees of confluent growth.

All tests were performed three times, using fresh suspensions of Candida albicans prepared immediately before each inoculation.

RESULTS AND DISCUSSION

Reports of blood culture using clinical specimens are sometimes difficult to interpret due to problems in standardization of blood samples. Another concern is that clinical specimens usually render a small number of positive cultures for a specific organism such as Candida albicans, and this has statistical implications. Furthermore, in the majority of the studies, the quantification of microorganisms in the sample was not performed (3, 19, 26).

The methodology applied in this study allowed the use of equal volumes of blood in each method as well as inoculation of a previously established number of cells. The number of cells tested was related to those more frequently associated with candidemia (14, 26). Counts of 100 and 10 cells/ml of blood were used to correspond to the counts observed in patients with infections related to catheters. Suspensions of 1 and 0.5 cells/ml were used to resemble patients with candidemia related to other sources of infection (27).

Our results showed that it was possible to detect Candida albicans in blood, regardless both the number of cells and the methodology. In fact, the amount of colonies on agar chocolate plates in each dilution was comparable, despite the method employed (Table 1).

![Figure 1](image.png)

**Figure 1** - Confluent growth of Candida albicans onto chocolate agar plates incubated for 24 h at 35°C.

<table>
<thead>
<tr>
<th>Number of cells of Candida albicans</th>
<th>Method</th>
<th>Growth</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 cells/ml of blood</td>
<td>Concentration</td>
<td>P +++</td>
</tr>
<tr>
<td>100 cells/ml of blood</td>
<td>Conventional</td>
<td>P +++</td>
</tr>
<tr>
<td>10 cells/ml of blood</td>
<td>Concentration</td>
<td>P +++</td>
</tr>
<tr>
<td>10 cells/ml of blood</td>
<td>Conventional</td>
<td>P +++</td>
</tr>
<tr>
<td>1 cell/ml of blood</td>
<td>Concentration</td>
<td>P ++</td>
</tr>
<tr>
<td>1 cell/ml of blood</td>
<td>Conventional</td>
<td>P ++</td>
</tr>
<tr>
<td>0.5 cells/ml of blood</td>
<td>Concentration</td>
<td>P +</td>
</tr>
<tr>
<td>0.5 cells/ml of blood</td>
<td>Conventional</td>
<td>P +</td>
</tr>
</tbody>
</table>

P = positive results; +++ / ++ = confluent growth; + = uncountable colonies.

In order to determine the time required by each method to obtain positive results, we compared observations made in the first 24 h of test, i.e., presence of blastoconidia in Gram stain prepared from the conventional method with growth on solid media from the Isolator™. Blastocconidia were observed in all Gram stains and the number of yeast seen by microscopy was directly related to the original number of cells in the sample (Fig. 2).

According to our results, it appears reasonable to conclude that the concentration method alone did not
reduced the time required for diagnosis, since positive results were obtained 24 h after inoculation of *Candida albicans* in both conventional and concentration methods. The mentioned advantage of Isolator™ in clinical reports may, therefore, be explained by the process of lysis which liberates microorganisms from phagocytes, increasing the number of viable yeasts (19, 35).

We suggest that the conventional method of growth in BHI for 24 h, either in liquid or biphasic media, and the concentration method by Isolator™ display comparable ability for the recovery of yeasts from blood.

Although the process of concentration appears not to improve the recovery of *Candida albicans* from blood, other advantages of the Isolator™ system deserve consideration. Quantitative results, supplied by Isolator™, although unable to determine clinical importance of the microorganism, may be important to determine the source of the infection and to follow therapy (26, 34). It has to be considered, however, that quantitative results are difficult to interpret because the method is not reproducible, and routine variation in the concentration of yeasts in blood during the course of infection is not well delimited (34).

Another advantage of Isolator™ is that direct inoculation of sediment onto agar plates would allow earlier isolation of colonies and a quicker identification (14). This, however, seems not to be so relevant as one can also obtain diagnosis of candidemia in the first 24 h through the visualization of blastoconidia directly in the broth of the conventional method, as we have demonstrated in this study.

Furthermore, identification and sensitivity tests can be done directly from broth (10).

Since concentration alone seems not improve the recovery of *Candida albicans*, the reported superiority of Isolator™ may be explained by lysis. Studies using conventional broth blood culture, supplemented with a lytic agent are therefore warranted. This will certainly be an enormous improvement for laboratories that use conventional methods.

**RESUMO**

**Processo de concentração do sangue na detecção de *Candida albicans***

O sistema de lise-centrifugação (Isolator™) tem sido considerado como método padrão para aumentar as taxas de diagnóstico de candidemia através de hemocultura. Neste estudo, o processo de concentração, segundo este sistema, foi comparado com métodos convencionais de cultivo em meio líquido e bifásico. Foram realizados testes “in vitro” utilizando *Candida albicans* em contagens de 100, 10, 1 e 0,5 células/ml de sangue. Culturas em ágar-chocolate foram realizadas a partir do sedimento obtido pela centrifugação do tubo de Isolator™ e do caldo das culturas convencionais após sua incubação por 24 horas a 35°C. Esfregaços corados pelo Gram, preparados a partir dos métodos convencionais, também foram observados em 24 horas. Foi possível detectar *Candida albicans* independentemente do número de células ou da metodologia utilizada. O tempo para diagnóstico também não foi diferente para os métodos comparados, já que blastoconídios e crescimento foram observados no mesmo prazo de tempo. Assim, sugerimos que o processo de concentração não é o maior fator responsável pelas taxas de recuperação de *Candida albicans* a partir do sangue obtidas pelo sistema Isolator™.

**Palavras-chave:** candidemia, hemocultura, sistema de lise-centrifugação.

**REFERENCES**