A novel approach for fast detection of sepsis with Gram-negative bacterial infection

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Summary

Sepsis, a life-threatening systemic infection, requires quick treatment. Gram-negative bacteria (GNB) are the major causative pathogens and their endotoxin can be a surrogate biomarker for diagnosis. We explored a fast identification of GNB by first culturing blood to increase endotoxin levels and then detecting endotoxin by Tachypleus amebocyte lysate (TAL) with kinetic turbidimetric assay (KT-TAL). Heating samples could significantly increase the endotoxin released from GNB; speed and time of centrifugation, and sample dilution could affect the endotoxin results. At a high GNB load, endotoxin was detected 3 h after culture, 6.5 h earlier than the BD BACTEC blood culture system detecting GNB. At a low GNB load, endotoxin was detected at 9 h after culture, 13 h earlier than the BD BACTEC system. In a sepsis patient with Acinetobacter baumannii, we detected endotoxin at 12 h after culture, while the BD BACTEC system needed 28.5 h for detection, allowing physicians an earlier decision on appropriate treatment.

Introduction

Sepsis requires immediately appropriative treatment, since each 1-h delay in diagnosis will increase mortality rates by 5–10% (Pfafflin and Schleicher, 2009; Schuts et al., 2016). A fast detection of a blood-borne pathogen is thus critical for rescuing life (Kibe et al., 2011; Turner et al., 2017).

At present, blood culture is the gold standard for the diagnosis of sepsis, but it needs blood culture for about 16–28 h (Sarkar et al., 2015), hampering early treatment decision. In addition, not all GNB in blood could be detected by the BD BACTEC system (Thomas et al., 1984).

Amebocyte Lysate (AL) is the most sensitive reagent to detect the endotoxin of GNB (Cooper et al., 1971; Levin et al., 1972). The enzymatic clotting cascade can be kinetically recorded with a turbidimetric reader (Jorgensen et al., 1973). This sensitive assay has been widely used in detecting the endotoxin in different biosamples (Cooper et al., 1972; Jorgensen and Jones, 1975). Compared to the conventional culture assay for GNB, the endotoxin AL assay is more sensitive and reliable (McCormey et al., 1983). However, components of plasma/serum can interfere with endotoxin detection at the onset of sepsis. (Gnauck et al., 2015; Bottiroli et al., 2017). Hence, it is urgently needed to develop a better approach for detecting endotoxin.

We propose the TAL-based approach shown in Fig. 1 for an early diagnosis and treatment monitoring of GNB sepsis.

BD BACTEC blood culture system detecting GNB. At a low GNB load, endotoxin was detected at 9 h after culture, 13 h earlier than by the BD BACTEC system. In a sepsis patient with Acinetobacter baumannii, we detected endotoxin at 12 h after culture, while the BD BACTEC system needed 28.5 h for detection, allowing physicians an earlier decision on appropriate treatment.
New procedure for LPS detection in blood suspected with GNB infection

**A**  
Add 3–5 ml blood to culture bottle  
↓  
17°C, shaking at 250rpm  
Take 0.5 ml sample every hour starting on 2 h after culture  
↓  
Process the samples:  
1. Dilute 1:4–10  
2. Heat at 100°C for 10 min  
3. Spin at 450 g for 3 min  
4. Dilute supernatant at 1:10  
5. Analyze with KT-TAL system  
↓  
Compare with LPS standard curve to determine if GNB in suspected blood grow with times

**B**  
Fig. 1. Flow chart for new procedure of LPS detection in blood suspected with GNB infection and its reference standard: (A) 3–5 ml blood was added into culture bottle. 2 h later, 0.5 ml sample was taken every hour for dynamic test of LPS. The samples were diluted, heated, spun and diluted again before LPS quantitative measurement with KT-TAL system. (B) The results were compared with LPS standard curve to determine if LPS increased as GNB in suspected blood grew with times.

Endotoxin exists as free endotoxin and endotoxin associated with intact cell walls (Jorgensen et al., 1973). Heating not only promotes the release of endotoxin, but also denatures and precipitates interfering material (Hurley, 1995). Indeed, heating increased the amount of detectable endotoxin (Table 1), thereby shortening the time for endotoxin detection. No increase over time was seen for ET with a culture from a Gram-positive bacterium (Table 1).

Since the KT-TAL assay measures gel formation with endotoxin, it requires a clean and transparent sample. To remove cloudy materials, samples were centrifuged. Centrifugation at 5000 g for 3 min leads to a 15% loss of endotoxin, while at 450 g for 3 min, only 9% was lost. If the sample is clear and transparent, no centrifugation is necessary.

To define an optimal dilution factor, we took blood samples cultured in BD BACTEC aerobic bottles spiked with 1 EU ml⁻¹ of endotoxin. At 1:10 and 1:20 dilution, endotoxin was not or only inefficiently detected, while at 1:40 and 1:100 dilution, endotoxin level close to the spiked 1 EU ml⁻¹ endotoxin was recovered (>97%) (Figure S1).

To test if the combination of blood culture with KT-TAL assay could detect ET of GNB earlier than the conventional BD BACTEC system, the same amounts (0.5 McF) of *E. coli* or *Klebsiella pneumoniae* were diluted at 1:10⁻⁸ and resulted in a positive reaction after 3–4 h in the KT-TAL assay compared to 9.5 and 12.5 h in the BD BACTEC system. At the 1:10⁻⁹ dilutions, the detection time was for *E. coli* 9 and 22 h for KT-TAL assay and BD BACTEC respectively.

Finally, a patient with suspected and later proven *VITEK 2, bioMérieux* Acinetobacter baumannii sepsis was tested. The BD BACTEC system reported pathogen

![Fig. 1](image-url)

**Table 1.** Effect of heating on ET release from GNB samples and assay specificity.a

<table>
<thead>
<tr>
<th>Bacterial samples</th>
<th>Sampled after hours of culture</th>
<th>ET detected (EU ml⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Unheated</td>
<td>Heated</td>
</tr>
<tr>
<td><em>E. coli</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>0.144</td>
<td>6.689*</td>
</tr>
<tr>
<td>5</td>
<td>0.326</td>
<td>10.854*</td>
</tr>
<tr>
<td>6</td>
<td>0.999</td>
<td>&gt;14.125*</td>
</tr>
<tr>
<td>7</td>
<td>4.237</td>
<td>&gt;14.125*</td>
</tr>
<tr>
<td>8</td>
<td>10.706</td>
<td>&gt;14.125*</td>
</tr>
<tr>
<td>9</td>
<td>&gt;14.125</td>
<td>&gt;14.125</td>
</tr>
<tr>
<td><em>Staphylococcus aureus</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4–9</td>
<td>&lt;0.007</td>
<td>&lt;0.007</td>
</tr>
</tbody>
</table>

a. Sample diluted at 1:4 with ET-free water, heated at 100°C for 10 min, spin at 450 g for 3 min, and then the supernatant was further diluted at 1:10 for ET measurement with KT-TAL system. The experiments were repeated three times with the same tendency, that is in the GNB *E. coli* group the ET of heated samples was much higher than the unheated samples (*P < 0.05*), while in Gram-positive *Staphylococcus aureus* group, there was no difference of ET between the heated and unheated samples.
positive with GNB at 28.5 h after culture, while KT-TAL system detected endotoxin at 12 h, 16.5 h earlier than the conventional BD BACTEC system did.

The TAL-based turbidimetric assay provides quantitative information, allowing a follow-up of GNB infection before and after treatment within 1 h assay time that only needs a 37°C shaker for bacteria culture, a microplate reader and TAL reagent kit. However, sepsis with Gram-positive pathogens will be missed.

Conflict of interest

None declared. All the blood sample collection was approved by the institutional review board committee of the First Affiliated Hospital at Fujian Medical University (#2013-29). The operation processes were followed the institutional safety guideline for laboratory involved in handling of pathogen bacteria.

References


Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Table S1 Optimizations of sample dilution by recovery test of spiked samples.