Real life turnaround time of blood cultures in the clinical microbiology laboratory: results of the first Italian survey, May 2015

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Summary

Background and aims: Blood culture (BC) results are essential to guide antimicrobial chemotherapy for patients with sepsis. However, BC is a time-consuming exam, which can take several days. Reducing BCs turn around time (TAT) could impact on multiple outcome parameters and TAT monitoring is an important tool for measurement of microbiology laboratory performance. The aim of this study was to provide an overview of BC TATs among Italian microbiology laboratories.

Materials and methods: Five laboratories collected and recorded, for a month period, date and time of the BC processing events. Cumulative TATs were analysed using the GraphPad software.

Results: Participating laboratories reported data from 302 sepsis episodes. The median time from when the BC system produced a positive signal until Gram-stain results were reported was 7.6 hours. A rapid molecular identification and antimicrobial susceptibility testing (AST) was performed in 26.5% of BCs. Mean TAT for identification report was significantly lower when a molecular approach was adopted (12 vs. 28.7 hours, P<0.001). Similarly, results of the molecular AST were obtained more than 24 hours in advance compared with phenotypic AST (mean 13.2 vs. 47.6, P<0.001). TATs from BC positivity of laboratories opened 7 days/week were not significantly lower than those of laboratories opened 6 days/week.

Conclusions: BC is a time-consuming exam, however, molecular identification and AST methods can drastically reduce time to results. The lack of difference between TATs observed for laboratories working 7 days/week and 6 days/week, coupled with a high rate of BCs turning positive during the night enable to conclude that the most urgent measure to reduce TATs is the expansion of laboratory regular duty hours.

Introduction

Sepsis is a severe disease associated with high morbidity and mortality (7). Although conflicting data exist, early administration of appropriate empirical antimicrobial therapy, coupled with supportive treatment, have been shown to have an important impact on outcome (6,9,10,16).
Blood culture (BC) is the cornerstone for aetiological diagnosis of septicemia and results are important to shift from empiric to definitive antimicrobial chemotherapy. However, BC is a time-consuming exam, which can take longer than 48 hours.

Although robust data showing a significant correlation between timeliness of reporting blood culture results and mortality are lacking, it seems likely that the two outcomes are associated. However, well structured studies demonstrated that the more rapid is the laboratory data reporting, the lower is the patients length of stay (both in Intensive Care Unit and hospital) and the antimicrobial consumption (4).

Newly introduced technologies such as PCR-based methods, fluorescence in situ hybridization and MALDI-TOF have expedited the causative pathogen identification (1,5,8,19). Also on the antimicrobial susceptibility testing (AST) scenario there are several chances to reduce the TAT: i) direct inoculation of automatic susceptibility testing instruments or use of young culture on agar media (12,13,17,20), and ii) use of liquid culture or time lapse microscopy based technologies (1,11,14).

Molecular methods performed directly on positive blood cultures (broths) are able to provide preliminary information on antimicrobial resistance genes presence and pathogens identification in a turn-around time (TAT) of approximately three hours, hence resulting in a dramatic change of the Clinical Microbiology Laboratory contribution to the management of septic patients (2,3,15,18).

TAT is a key indicator of clinical laboratory performance and is used by many clinicians to judge the quality of the laboratory service. For this reason we implemented a Clinical Microbiology Laboratory Network (CMLN), including laboratories adopting new approaches for expediting ID and AST results, aimed to provide an updated overview of real-life blood culture TATs among Italian microbiology laboratories.

Materials and Methods

A CMLN, consisting of five large Italian laboratories referring to members of the Gruppo di Lavoro Infezioni Paziente Critico working group (belonging to Associazione Microbiologi CLinici Italiani) was created in April 2015. Three out of five laboratories were located within Hospitals with ≥1000 beds, while two in Hospitals with 400-700 beds.

CMLN Laboratories were asked to collect and record in a dedicated database, for a month period (1 to 30 May 2015), the date and time of the following BC processing events: i) laboratory check-in (the time when the BC was taken in charge by the laboratory staff); ii) BC positivity signal (instrument alarm); iii) Gram-stain microscopic examination completed and results successfully reported to clinical staff; iv) bacterial/fungal identification, obtained with rapid molecular methods other than MALDI-TOF or biochemical ID report; v) MALDI-TOF or biochemical bacterial/fungal ID report (including results obtained with lysis-filtration protocols, young culture processing and conventional overnight approach); vi) molecular detection of antimicrobial resistance markers, directly from BC liquid, report; vii) phenotypic AST final report (including results obtained with automated AST instruments or broth microdilution method using young or overnight culture).

Analysis was restricted to data regarding the first positive blood culture bottle from a set, for all consecutive, non-replicated, sepsis episodes occurred within the study period. BCs positive for contaminant (positivity of one bottle/set only for skin colonizing bacteria) were excluded from the analysis.

TATs were expressed in hours and the minimum, maximum, median, 25th percentile and 75th percentile values were calculated using the GraphPad software (GraphPad Software, Inc.) TATs were compared using the two-tailed t test.

The participating laboratories were also asked to provide full information on systems used for: BCs monitoring; conventional ID; molecular pathogen ID from BC liquid and detection of resistance determinants markers; phenotypic AST; software used for results reporting and storage (laboratory information system). Moreover, the laboratories were asked to provide the number of BC exams performed in the previous year (2014) as indicator of their workload, together with details on opening days/week and hours/day.

Results

Among the participating laboratories, two out of five, processed positive BCs 7 days/week while the other three processed positive BCs 6 days/week. All participants used continuously monitoring blood culture systems.

During opening days, participating laboratories processed BCs over a timespan ranging from 9 to 15 hours with an intermittent processing schedule during opening hours. The majority of IDs were obtained by MALDI-TOF technology and phenotypic AST was performed mainly with automated AST systems (Table 1 and data not shown).

Overall, during the surveillance period, CMLN laboratories reported data from 302 sepsis episodes (range 41-91 episodes per Laboratory). The molecular ID and AST were performed for 80 BCs (26.5%) and the phenotypic AST was performed in 277 cases (92%). The mean and median times to positivity from check-in (Check-Pos) were 15.1 and 20.4 hours, respectively.

Among all laboratories, the median time from when the BC system produced a positive signal until Gram-stain results were reported (Pos-gram) was 7.6 hours, ranging from 0.2 to 86.4 hours (mean 10.3 hours). TAT analysis is summarised in Table 2 and Figure 1. Mean TAT for ID report was significantly lower when a molecular approach was adopted (12 vs. 28.7 hours, P<0.001). Similarly, results of the molecular AST were obtained more than 24 hours in advantage comparing with phenotypic AST (mean 13.2 vs. 47.6, P<0.001).

Cumulatively, TATs from BC positivity of Laboratories opened 7 days/week were not significantly lower than those of Laboratories opened 6 days/week with mean Pos-Gram: 11.5 vs. 9.1, mean Pos-ID: 30 vs. 27.9 and mean Pos-AST: 48 vs. 47.2 hours.

Regarding time to positivity, 49.3% of blood cultures turned positive during the night (from 20:00 to 8:00) and therefore were processed the following morning.

Interestingly, the TATs distribution for identification and molecular AST were very similar to that of Gram-stain report (Figure 1).

Discussion and Conclusions

Overall, the analysis of data from CMLN showed that the determination of phenotypic conventional AST is still a time-consuming process that requires a mean of 47.6 hours from BC positivity. However, the adoption of AST protocols based on the use of young culture or rapid broth enrichment protocols can lead to the obtainment of AST results within 12 hours.

The relevant advantage of use of MALDI-TOF ID is confirmed by the fact that mean TAT of ID anticipates by more than 20 hours the phenotypic AST result (mean TAT 28.7 vs. 47.6 hours).

Our analysis confirms that molecular ID and AST methods able to give rapid results from BC broth are breakthrough technologies that can dramatically reduce time ID and AST results (TAT 12 and 13.2 hours from BC positivity, respectively). However, due to the high cost of these technologies in comparison to conventional ID and AST, they can be adopted in a subset of selected cases only (in our surveillance, molecular ID and AST were performed in approximately a quarter of
Table 1. Results obtained form laboratory tests.

<table>
<thead>
<tr>
<th>Centre</th>
<th>Beds</th>
<th>BCs in 2014</th>
<th>N. positive BCs, May 2015</th>
<th>N. sepsis episodes, May 2015</th>
<th>N. molecular AST</th>
<th>System used for molecular AST</th>
<th>Laboratory opening hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>607</td>
<td>14,356</td>
<td>97</td>
<td>42</td>
<td>8</td>
<td>GeneXpert (Cepheid Sunnyvale, CA, USA)</td>
<td>8:00 to 20:00, 7 days/week</td>
</tr>
<tr>
<td>2</td>
<td>1644</td>
<td>22,000</td>
<td>400</td>
<td>91</td>
<td>45</td>
<td>Easyplex (Amplex BioSystems Giessen, Germany), GeneXpert</td>
<td>7:00 to 19:00, 6 days/week</td>
</tr>
<tr>
<td>3</td>
<td>460</td>
<td>25,047</td>
<td>248</td>
<td>41</td>
<td>19</td>
<td>Film Array (bioMérieux Marcy l’Etoile, France), home brew PCR</td>
<td>7:30 to 20:00, 7 days/week</td>
</tr>
<tr>
<td>4</td>
<td>1000</td>
<td>10,151</td>
<td>139</td>
<td>62</td>
<td>4</td>
<td>GeneXpert</td>
<td>7:00 to 16:00, 6 days/week</td>
</tr>
<tr>
<td>5</td>
<td>1297</td>
<td>19,482</td>
<td>277</td>
<td>66</td>
<td>4</td>
<td>Verigene (Nanosphere, Northbrook, IL, USA)</td>
<td>7:00 to 22:00, 6 days/week</td>
</tr>
</tbody>
</table>

BC: blood cultures; AST: antimicrobial susceptibility testing.

Table 2. Results of turn around time analysis.

<table>
<thead>
<tr>
<th></th>
<th>Check-Pos</th>
<th>Pos-Gram</th>
<th>Pos-ID</th>
<th>Pos/IDM</th>
<th>Pos-ATBM</th>
<th>Pos-ATB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of values</td>
<td>302</td>
<td>302</td>
<td>222</td>
<td>80</td>
<td>80</td>
<td>277</td>
</tr>
<tr>
<td>Range</td>
<td>1.7-109.4</td>
<td>0.2-86.4</td>
<td>1-132</td>
<td>1-38.4</td>
<td>1-41.5</td>
<td>9.6-138.4</td>
</tr>
<tr>
<td>Median</td>
<td>15.1</td>
<td>7.6</td>
<td>23.5</td>
<td>7.9</td>
<td>8.6</td>
<td>41.2</td>
</tr>
<tr>
<td>Mean</td>
<td>20.4</td>
<td>10.3</td>
<td>28.7</td>
<td>12</td>
<td>13.2</td>
<td>47.6</td>
</tr>
</tbody>
</table>

Check Pos, Check-in to positivity; Pos-Gram, positivity to Gram-stain; Pos-IDM, positivity to molecular pathogen identification; Pos-ID, positivity to pathogen identification; Pos-ASTM, positivity to molecular detection of resistance markers; Pos-AST, positivity to phenotypic antimicrobial susceptibility testing.

Figure 1. Time intervals used for the turnaround time analysis (in days): from check-in to positivity (Check-Pos); from positivity to Gram-stain reporting (Pos-Gram); from positivity to molecular pathogen identification reporting (Pos-IDM); from positivity to pathogen identification reporting (Pos-ID); from positivity to molecular detection of resistance markers reporting (Pos-ASTM); from positivity to phenotypic antimicrobial susceptibility testing results (Pos-AST). Boxes indicate the range from 25 to 75%.

sepsis episodes). Overall, participating laboratories declared to process with molecular ID and AST the positive BCs obtained from patients admitted to Intensive Care Units, Neonatal Intensive Care Units and Haematology wards.

In all participating laboratories, even for fast molecular methods and Gram-stain, we found a significant delay of observed versus theoretical TATs. This phenomenon is probably related to the high proportion (49.6%) of BC that turned positive outside of the regular duty hours, causing several hours delays in the analysis of positive BCs.

The lack of difference between TAT observed for laboratories working 7 days/week and 6 days/week enabled us to conclude that the most urgent measure to reduce TATs in microbiology is the progressive expansion of regular duty hours up to 24 hours/day.

One possible limitation of our study is that information on the preanalytical phase were not available for all laboratories, since storage and transport times are generally out of the laboratory’s control. Therefore this phase was excluded from the analysis. However, even if the preanalytical phase could influence the time elapsed from check-in to BC positivity, the time elapsed from BC positivity to the results of the various analytical steps is not expected to be biased by the conditions of the preanalytical phase and, in our opinion, these data can be reliably used as indicators of the performance of the laboratory workflow.

References

2. Banerjee R, Teng CB, Cunningham SA, et al. Randomized trial of


