Nonautomated Blood Cultures in a Low-Resource Setting: Optimizing the Timing of Blind Subculture

Sien Ombelet,1,2* Marjan Peeters,1 Chhundy Phe,3 Achilleas Tsoumanis,1 Chun Kham,3 Syna Teav,3 Erika Vlieghe,4 Thong Phe,3 and Jan Jacobs1,2

1Department of Clinical Sciences, Institute of Tropical Medicine, Antwerp, Belgium; 2Department of Immunology, Microbiology and Transplantation, KULeuven, Leuven, Belgium; 3Sihanouk Hospital Center of Hope, Phnom Penh, Cambodia; 4Department of General Internal Medicine, Infectious and Tropical Diseases, University Hospital Antwerp, Antwerp, Belgium

Abstract. Laboratory procedures for blood cultures in a hospital in Phnom Penh were adapted to optimize detection of *Burkholderia pseudomallei*, an important pathogen in this setting. The effects of these changes are analyzed in this study. Blood cultures consisted of two BacT/ALERT bottles (bioMérieux, Marcy-l’Etoile, France). Growth was detected visually by daily inspection of the bottles. In 2016, the aerobic–anaerobic pair (FA/FN FAN) was substituted by an aerobic pair of BacT/ALERT FA Plus bottles. Blind subculture (BS) (subculture in the absence of visual growth) was advanced from day 3 to day 2 of incubation in July 2016. In July 2018, it was further advanced to day 1 of incubation. From July 2016 to October 2019, 9,760 blood cultures were sampled. The proportion of cultures showing pathogen growth decreased from 9.6% to 6.8% after the implementation of the laboratory changes (*P* < 0.001). Advancing the BS from day 3 to day 2 led to an increased proportion of pathogens detected by day 3 (92.8% versus 82.3%; *P* < 0.001); for *B. pseudomallei*, this increase was even more remarkable (92.0% versus 18.2%). Blind subculture on day 1 similarly increased the proportion of pathogens detected by day 2 (82.9% versus 69.0% overall, 66.7% versus 10.0% for *B. pseudomallei*; both *P* < 0.001). However, after implementation of day 1 subculture, a decrease in recovery of *B. pseudomallei* was observed (12.4% of all pathogens versus 4.3%; *P* < 0.001). In conclusion, earlier subculture significantly shortens time to detection and time to actionable results. Some organisms may be missed by performing an early subculture, especially those that grow more slowly.

INTRODUCTION

Bloodstream infections (BSIs) lead to considerable morbidity and mortality worldwide.1–3 Blood cultures are currently the most important tool for diagnosis of BSIs.4 Furthermore, they enable performing antibiotic susceptibility testing of the causative pathogens, which is instrumental in patient management and antibiotic stewardship. However, in many low-resource settings, blood culture surveillance is lacking.5–7 Automated equipment for blood culture incubation is too expensive and its maintenance too exigent for these settings.

Since 2007, blood cultures are collected in Sihanouk Hospital Center of Hope (SHCH), a nongovernmental organization 30-bed hospital in Phnom Penh, Cambodia. A pair of BacT/ALERT blood culture bottles (bioMérieux) are sampled from separate venipunctures and incubated in a static incubator, as opposed to an automated incubator. The bottles are daily inspected visually to detect changes in the color indicator on the bottles. These procedures were outlined in a previous publication, describing results of the blood culture surveillance for the period 2010–2015 in SHCH.8

One of the conclusions of the previous study was that detection of *Burkholderia pseudomallei*, a key pathogen in this setting and the causative agent of melioidosis, is tardive compared with other pathogens, despite results from another study, showing fast growth of *B. pseudomallei* in automated blood culture systems.9 Another finding from our surveillance was the increased value of aerobic bottles compared with anaerobic bottles, especially for the detection of non–glucose-fermenting Gram-negative organisms such as *B. pseudomallei*. Some procedural changes were therefore implemented in the laboratory, to increase and speed up detection of this pathogen (Figure 1).8

First, instead of performing a blind subculture (BS) (i.e., a subculture of all aerobic bottles in the absence of visual signs of growth) on day 3 of incubation, it was decided to advance the BS to day 2 of incubation. Second, the aerobic–anaerobic pair of bottles was replaced by an aerobic–aerobic pair. Moreover, the BS medium was changed from chocolate agar to McConkey agar because of the high number of contaminants found on chocolate agar. Incidentally, the bottle types used from 2010 to 2015 (the charcoal-containing BacT/ALERT FA FAN and FN FAN) were withdrawn from the market by bioMérieux and replaced by the resin-containing BacT/ALERT FA Plus bottles. These changes were consecutively implemented from April to June 2016.

In this study, we describe how these procedural changes led to differences in time to detection (time between incubation and first sign of growth) and in “time to colonies” (time between incubation and first recovery of colonies on agar) of key pathogens. An interim analysis 2 years after implementation led to additional changes in the procedures (July 2018): BS was further advanced to day 1 of incubation and was performed on blood agar instead of McConkey agar, to demonstrate earlier growth of *Staphylococcus aureus* BSI. Figure 1 summarizes the procedural changes over the different surveillance periods.

METHODS AND MATERIALS

Laboratory materials and procedures. BacT/ALERT FA Plus bottles (reference number 410851) were used for blood culture sampling. For adults, a set of blood culture bottles (2 × 10 mL in two FA Plus bottles) was sampled. For children 15 years of age or younger, guidelines recommend sampling of only one (aerobic) bottle. However, for 66% of cultures in
children in SHCH (n = 128), two bottles had been sampled. On arrival in the laboratory, bottles were weighed and incubated in a static incubator at 35°C ± 1°C for 7 days. Bottles were inspected twice daily for signs of growth (such as color change of the indicator). Bottles with signs of growth were further processed by performing Gram stain of the bottle and subculture on solid media appropriate according to the Gram stain result. A BS was performed on McConkey agar for each bottle on day 2 of incubation from July 2016 to July 2018. From July 2018 to November 2019, BS was performed on blood agar for each bottle on day 1 of incubation.

**Laboratory methods for identification and antibiotic susceptibility testing.** After growth of colonies on subculture, bacterial isolates were identified using conventional microbiological techniques and Analytical Profile Index (API®) tests (bioMérieux) when necessary. Antibiotic susceptibility was tested with disk diffusion. Antibiotic disks used were Neo-Sensitabs™ (Rosco Diagnostica, Taastrup, Denmark).

**Inclusion and exclusion criteria.** Indications for blood culture sampling were based on the systemic inflammatory response syndrome criteria: presence of tachycardia, tachypnea, fever, or hypothermia (temperature > 38°C or < 36°C); altered mental status; and presence of inflammatory parameters such as leukocytosis or leukopenia. Homemade glass bottles and solitary bottles from adults (or children aged > 15 years) were excluded as they do not reflect optimal blood culture methods and to facilitate comparison with the period of 2010–2015, when these bottles were also excluded from the analysis. Cultures are sampled free of charge, excluding any economical bias to the patient population included in the study.

**Data collection and extraction.** Patient demographic data, clinical information (such as use of antibiotics 14 days before blood culture sampling) as well as detailed microbiological data were extracted from the laboratory information system (structured query language [SQL]) into Microsoft® Excel® for Office 365 (Microsoft Corp., Redmond, WA) for the period July 2016 up to November 2019. Doubtful results and likely errors were verified with the laboratory notebooks. Only paired bottles were considered for analysis, except for children younger than 15 years of age, for whom also single bottles were included. Homemade bottles were excluded from analysis.

**Blood volume measurement and calculation.** Blood culture bottles were measured on arrival in the laboratory. The weight was noted in the laboratory logbooks and entered in the SQL database (Oracle Corporation, Redwood City, CA). The volume of blood was calculated by subtracting the mean empty weight of a BacT/ALERT FA Plus bottle (measured mean weight = 61.9 g, SD = 0.12 g) and next dividing the result by the density of blood (=1.06 g mL−1). Bottles from children aged ≤15 years were not included in the analysis of the blood volume, as guidelines for optimal volume in children differ. The volume of blood was calculated by subtracting the mean empty weight of a BacT/ALERT FA Plus bottle (measured mean weight = 61.9 g, SD = 0.12 g) and next dividing the result by the density of blood (=1.06 g mL−1). Bottles from children aged ≤15 years were not included in the analysis of the blood volume, as guidelines for optimal volume in children differ.

**Incubation delay.** Day of blood culture request and day of reception in the laboratory were compared to assess possible delay in incubation (needle-to-incubator time).

**Statistical analysis.** Export of the data from SQL to Microsoft Excel was cleaned and imported into R (R Foundation for Statistical Computing, Vienna, Austria). Statistical tests were performed using R version 3.6.1 (July 5, 2019). Univariate differences in proportions, means, and median values were assessed using chi-square test, Student’s t-test, and Wilcoxon rank-sum test, respectively. A segmented logistic regression, modeling the effect of one (July 2016) or two interventions (July 2016 and July 2018), on pathogen growth rate was performed. To assess for the impact of other variables on pathogen growth, such as age, gender, hospitalization status, prior antibiotic use, volume sampled, and delay in incubation, multivariate logistic regression models were calculated for each surveillance period separately and for periods 2 and 3 combined. Presence of a trend over time in the total number of cultures sampled and absolute number of grown pathogens was assessed using Poisson regression. Trends over time in pathogen growth rate and recovery on BS were assessed using logistic regression.

**Definitions.** Definitions of blood culture-related terms used in this study are given in Table 1. For further reference, the surveillance period of 2010–2015 is called “period 1,” the surveillance period of July 2016 to June 2018 is called “period 2,” and the surveillance period of July 2018 to October 2019 is called “Period 3.” The proportion of blood cultures showing growth of a pathogen will be referred to as “pathogen growth rate.”

**RESULTS**

**Blood culture results from July 2016 to October 2019.** From July 2016 to October 2019, 20,894 bottles were sampled (Figure 2). After exclusion of homemade and solitary bottles, 19,523 bottles remained for analysis. These correspond to 9,760 cultures from 9,314 suspected episodes of BSI. They were obtained from 8,276 patients with median age of 52 years (2–98 years), including 247 children (age ≤18 years). Slightly more than half of patients were female (59.1%). Most of the
suspected BSI episodes (96.5%) were community acquired (Table 2).

In 7.1% of suspected BSI episodes, a pathogen was recovered from blood culture. In total, 664 pathogens were recovered. There were five polymicrobial infections, that is, episodes for which two different pathogens were recovered. The most frequent pathogens were *Escherichia coli*, *Salmonella* Paratyphi A, and *B. pseudomallei* (Figure 2), which accounted together for 62.8% of all recovered pathogens. The contamination rate expressed per bottle and per culture was 2.7% and 5.1%, respectively. In 26 of 496 contaminated cultures, an identical contaminant was found in both bottles (5.2%).

Suspected episodes of patients under antibiotic treatment (antibiotic administered 2 weeks or less ago) showed growth more often (8.0%) than episodes of patients not under antibiotic treatment (6.5%) (P = 0.017; Table 2). Suspected healthcare-associated infections were also associated with higher growth rates (P = 0.017; Table 2). The mean blood volume sampled per bottle was 9.51 mL (SD 1.90); this was significantly higher than the mean volume of the period 2010–2015, when it was 7.99 mL (P < 0.0001). Correct filling rate (between 8 mL and 12 mL) increased from 59.5% of all bottles to 79.4% (P < 0.001).

Blind subculture yielded 184 pathogens, representing 27.7% of all recovered pathogens. All contaminants found on BSI (n = 81) were recovered in the period July 2018 to October 2019, when the medium for BS was changed to blood agar: this allowed the growth of Gram-positive organisms. The most commonly isolated pathogen on BS was *Salmonella* Paratyphi A (n = 71; 38.6%), followed by *B. pseudomallei* (n = 40; 21.5%) and *E. coli* (n = 25; 13.6%). Proportionally, BS contributed most to detection of *B. pseudomallei*, yielding 66.7% of all recovered *B. pseudomallei*.

**Comparison of yield between the different surveillance periods.** Pathogen growth rate per suspected episode was 10.2% in period 1. In period 2, pathogen growth rate decreased to 7.1% (P < 0.001). In period 3, growth rate remained at 7.0%. The absolute number of cultures sampled per month increased during this period, from 177 per month to 239 per month in period 2 and to 252 cultures per month in period 3, as shown in Table 3. As also shown in this table, the recovery of pathogens in absolute numbers remained stable over the whole surveillance period (P = 0.17).

Table 4 shows the output of the multivariate and segmented regression models for the different surveillance periods. Pathogen growth rate per culture was stable over time within each surveillance period, but a significant decline in the

---

**TABLE 1**

Definitions used in this study

| Term | Definition |
|------|------------|
| Blood culture | Adults: a collection of blood culture bottles (2 or more) sampled at the same time. Children aged < 15 years: one blood culture bottle. |
| Blood culture set | Two aerobic bottles sampled from the same patient at the same time. |
| Solitary bottle | One bottle sampled from the same (adult) patient at the same time. |
| Cancelled blood cultures | Blood culture bottles that arrived in the laboratory but were not further worked up because of cancellation by the treating physician. |
| Suspected BSI episode | A suspected BSI episode was defined as all blood cultures sampled within a 14-day interval from the first sample, unless growth (see in the following text). |
| Culture-confirmed BSI episode | A BSI episode was defined as 1) the initial recovery of a pathogen from a suspected BSI episode, 2) the recovery of a pathogen different from the initial pathogen ≥ 48 hours after the recovery of the initial pathogen, or 3) the recovery of the same pathogen after at least a 14-day interval since the previous grown culture with this pathogen. |
| BS | A subculture performed in the absence of any visual signs of growth (in this case, change in color of the growth indicator). |
| Contamination rate | Skin and environmental bacteria (coagulase-negative staphylococci, *Corynebacterium* species, and *Bacillus* species) were categorized as blood culture contaminants. The rate of contaminants was defined as the number of bottles grown with contaminants divided by the total number of bottles collected (as each bottle was sampled by a separate venipuncture) and expressed as a percentage. |
| Pathogen growth rate | The number of blood cultures demonstrating growth of a pathogen, divided by the total number of blood cultures and expressed as a percentage. |
| Period 1 | Period of blood culture surveillance from 2010 to 2015; BS of aerobic bottle done on day 3 of incubation. |
| Period 2 | Period of blood culture surveillance from July 2016 to June 2018; BS done on day 2 of incubation. |
| Period 3 | Period of blood culture surveillance from July 2018 to October 2019; BS done on day 1 of incubation. |
| Day of incubation | Days of incubation were mentioned to indicate for instance the time-to-detection. They were defined as follows: Day 0 = reception in the laboratory. Day 1 = after 1 night of incubation. Day 2 = after 2 nights of incubation. Day 3 = after 3 nights of incubation. |
| Time to detection | The time between reception of the bottle in the laboratory and the moment that growth of a pathogen is detected and can be reported. For bottles showing growth first on BS, the time to detection is the day after the BS, that is, the day colonies are witnessed on BS. |
| Time to colonies | The time between reception of the culture in the laboratory and the moment that colonies are available on solid medium (agar) for further testing. |
| Community-acquired and healthcare-associated BSI episodes | Community-acquired and healthcare-associated BSI were defined according to the day of sampling, that is, at ≤ 2 days and > 2 days of hospital admission, respectively. |

*BS = Blind subculture; BSI = bloodstream infection.*
growth rate was seen after the first intervention (July 2016). Prior antibiotic treatment had a negative effect on growth rates in period 1, but a positive, albeit not statistically significant, effect in periods 2 and 3. Contrary to expectations, higher sampled volume was not associated with better recovery of pathogens, nor was increased incubation delay associated with worse recovery of pathogens.

Another important observation was the declining recovery of *B. pseudomallei* in period 3, when the BS was advanced to day 1 (Table 3). This followed a relative increase during period 2, when proportion rose from 8.3% to 12.4% (*P* = 0.02). The proportion of *B. pseudomallei* among all culture-confirmed BSI went down from 12.4% in period 2 to 4.3% (*P* < 0.001). Simultaneously, the proportion of *Salmonella Paratyphi A* further increased to 39.7% of all pathogens, demonstrating an obvious increasing trend since 2010 (Table 3).

**Comparison of time to detection and time to colonies between the different surveillance periods.** The yield of BS increased significantly when the subculture was advanced; it rose from 6.7% of all retrieved pathogens to 9.6% by advancing the subculture to day 2; it further increased to 45.1% by advancing subculture to day 1 (Table 3).

By advancing the BS to day 2 of incubation, the time to detection of growth in general, and specifically of *B. pseudomallei*, shortened significantly (Figures 3 and 4). The cumulative proportion of all pathogens (excluding *B. pseudomallei*) retrieved by day 3 (the day following the BS) increased from 86.6% to 92.9% (*P* = 0.002). For *B. pseudomallei*, cumulative proportion retrieved by day 3 increased rather spectacularly from 18.2% to 92% (*P* < 0.001).

Blind subculture on day 1 further shortened the time to detection. The proportion of total pathogens retrieved by day 2 increased from 69.0% to 82.9% (*P* < 0.001). Time to detection shortened for *B. pseudomallei*; cumulative growth on day 2 increased to 66.7% for all *B. pseudomallei* cultures, from only 10.0% when day 2 BS was performed (*P* < 0.001)(Figure 3).

Figures 3 and 4 also show the cumulative time to colonies; this is the interval between the moment of incubation to the moment colonies are available for identification and antibiotic susceptibility testing. A shorter time to colonies therefore leads to faster actionable results for the clinician. The impact of advancing the BS on time to colonies is even greater than that of advancing the BS on time to detection. A day 2 BS led to 80.1% of all pathogens (including *B. pseudomallei*) showing colony growth by day 3, versus 67.2% with a day 3 BS (*P* < 0.001). Similarly, advancing the subculture to day 1 led to a recovery of colonies on day 2 of 77.1%, versus only 39.3% when a day 2 BS was performed (*P* < 0.001).

**Figure 2.** Breakdown of blood cultures sampled in Sihanouk Hospital Center of Hope from July 2016 to October 2019 (periods 2 and 3). * In five episodes, two different pathogens were retrieved from the same culture.

---

**Table 2**

Demographic and clinical data of patients with suspected and culture-confirmed BSI episodes from July 2016 to October 2019 (periods 2 and 3)

|                          | All suspected BSI episodes | Non-confirmed BSI episodes | Culture-confirmed BSI episodes | P-value |
|--------------------------|----------------------------|---------------------------|-------------------------------|---------|
| No. of patients          | –                          | 8,276                     | 7,641                         | 635     |
| Female (% of number of patients) | –                          | 4,892 (59.1)             | 4,539 (59.4)                 | 353 (55.6) | 0.13 |
| Age, median (range) (years) | –                          | 52 (2–98)                 | 52 (2–95)                    | 48.5 (9–98) | < 0.001 |
| Total no. of BSI episodes | –                          | 9,314                     | 8,653                         | 659     |
| Healthcare vs. community | Community- acquired        | 8,408 (95.7)             | 7,831 (95.8)                 | 577 (93.5) | 0.017 |
| (% of number of episodes) (n = 8,790)* | Healthcare- associated     | 382 (4.3)                 | 342 (4.2)                    | 40 (6.5) |
| Antibiotic treatment† (% of number of episodes) (n = 9,113)* | Recent                     | 3,737 (40.1)             | 3,439 (40.6)                 | 298 (45.2) | 0.017 |

*BSI = bloodstream infection. *P*-values were calculated for differences in parameters between confirmed and non-confirmed episodes, using a multivariate logistic regression model of (suspected) episodes during periods 2 and 3.

* Episodes for which no data were available were not included in this analysis.

† Recent antibiotic treatment was defined as administration of antibiotics in the last 2 weeks.
showed an increasing trend, whereas the total number of pathogens retrieved increased in period 2 but decreased significantly in period 3.

| Pathogen growth rate | 10.7% | 7.1%** | 7.0% | P < 0.001 |
|----------------------|-------|--------|------|------------|
| Total number of pathogens retrieved | 1,087 | 387 | 277 | -- |
| Mean per month | 16.5 | 16.1 | 17.3 | P = 0.25 |
| Mean per month | 1.4 | 2.0* | 0.8** | P = 0.53 |
| Mean per month | 2.2 | 3.1* | 6.9** | P < 0.001 |

**BS = blind subculture; B. pseudomallei = Burkholderia pseudomallei. Pathogen growth rate was defined as proportion of blood cultures showing growth of a pathogen. Statistically significant differences with the previous surveillance period are indicated with an asterisk (* = P < 0.05; ** = P < 0.001). Logistic and Poisson regressions were used to detect a linear trend over time. This table shows that pathogen growth rate significantly declined in periods 2 and 3, but that the total number of pathogens retrieved remained stable. The total number of Salmonella Paratyphi A retrieved showed an increasing trend, whereas the total number of B. pseudomallei retrieved increased in period 2 but decreased significantly in period 3.**

DISCUSSION

This analysis of blood culture data from SHCH studied the effect of implementation of procedural changes in the laboratory on important quality indicators such as time to detection of pathogens and blood culture yield.

Advancing the BS to day 2 led to shorter time to detection overall and higher relative recovery of B. pseudomallei. This higher recovery may also have been caused by the change of an aerobic–anaerobic pair to an aerobic–aerobic pair; a previous analysis had shown that B. pseudomallei grew mainly in the aerobic bottle of a blood culture set (70/99 B. pseudomallei cultures grew only in the aerobic bottle). When an aerobic–anaerobic pair was used, 10 of 62 cultures growing B. pseudomallei showed growth in just one bottle (16.1%). Had an aerobic–anaerobic pair been used, half of these infections would theoretically have been missed. However, B. pseudomallei incidence may also have increased in the region during that period; epidemiological data on B. pseudomallei are scarce, but Calmette Hospital reported their incidence of B. pseudomallei to the Melioidosis Research Coordination Network, and observed a steady increase in identified cases from 2013 to 2018.

The overall pathogen growth rate per suspected episode decreased from 10.2% in period 1 to 7.1% for periods 2 and 3. This is still within the recommendations of blood culture guidelines, which recommend a pathogen growth rate between 6% and 12%, but the decrease was nonetheless significant and a cause of concern. Reasons for a declining growth rate can be related to efficiency of recovery of pathogens or to sampling indications. The indications for blood culture sampling have not changed between 2010 and the present, but we observed a sharp increase in the number of blood cultures sampled when comparing periods 2 and 3 with period 1 (Table 3). Because of hospital budget problems, the number of patients was limited in the period before 2016. The hospital went to full function of both outpatient departments and hospital beds since 2016. The number of pathogens detected per month, however, remained stable. These results suggest the drop in pathogen growth rate may have been caused by increased sampling, rather than suboptimal recovery of pathogens.

The replacement of an anaerobic bottle by another aerobic bottle led to an expected decrease in recovery of strictly anaerobic pathogens; it is however unlikely this would have had a significant effect on pathogen growth rate, as only 34 strictly anaerobic pathogens were recovered in period 1 (3.0% of all grown cultures; 0.3% of all sampled blood cultures). The omission of anaerobic bottles is contested, as many authors point out that anaerobic infections may not always be correctly predicted by clinical signs; antibiotic resistance among anaerobic bacteria is increasing, and some facultative pathogens show faster growth in anaerobic than aerobic bottles. However, our own evaluation of the use of anaerobic bottles in SHCH showed slower and less efficient overall pathogen growth in anaerobic bottles. Given the importance of the obligate aerobic pathogen B. pseudomallei in this setting and the inability of the laboratory to perform anaerobic antibiotic susceptibility testing due to technical limitations, we believe anaerobic bottles are of limited value in SHCH, which is in line with findings of other studies.

Further advancing the BS to day 1 increased the overall yield of BS and detection of B. pseudomallei by day 2 of incubation significantly (Table 3, Figure 3). The effect of the advanced BS on time to detection for pathogens other than B. pseudomallei was not statistically significant (Figure 4). Overall pathogen growth rate remained stable (despite further increase in numbers of cultures sampled as shown in Table 3), but relative frequency of B. pseudomallei decreased markedly. This relative decrease may have been partly due to a resurgence of the previously described Salmonella Paratyphi A outbreak in...
Cambodia that was observed simultaneously (Table 3). However, the absolute number of*B. pseudomallei* retrieved per month decreased significantly as well. Unfortunately, no official data about epidemiology of*B. pseudomallei* in this region are available after 2018, but personal communication from the Diagnostic Microbiology Development Program, working in Takeo Hospital, did not reveal a similar decrease in*B. pseudomallei* incidence in Takeo Province. Given the importance of BS in the detection of*B. pseudomallei*, its overall slow growth, as described previously, and scarce visual signs of growth of non-fermenting Gram-negative organisms, it is possible that growth of*B. pseudomallei* was missed by advancing the subculture.

![Cumulative time-to-detection *Burkholderia pseudomallei*](image1)

![Cumulative time-to-colonies *Burkholderia pseudomallei*](image2)

**Figure 3.** Cumulative time to detection (first sign of growth) and time to colonies for all pathogens except *Burkholderia pseudomallei*, comparing the three periods of surveillance. Statistically significant differences between periods of surveillance are indicated with asterisks (* = P < 0.05; ** = P < 0.001). This figure appears in color at www.ajtmh.org.
Optimal timing of BS largely depends on the goal of BS. An early BS (within 12–24 hours of incubation) decreases the time to detection, whereas a late BS serves as a final check for growth, to ensure no pathogens are missed. The “Clinical Microbiology Procedures Handbook” recommends a late subculture after 72 hours of incubation, whereas Clinical & Laboratory Standards Institute recommends performing subculture after 24–48 hours of incubation, and Cumitech advises an even earlier subculture after 12–18 hours of incubation.

A study looking at the optimal timing for subculture specifically for the detection of *B. pseudomallei* found that an early day 1 subculture confirmed growth of 52.3% of all *B. pseudomallei* cultures. An additional BS on day 2 yielded an extra 28.5% of grown cultures. However, the results of our study suggest that performing only a day 1 BS may decrease the overall recovery of *B. pseudomallei*. For settings where *B. pseudomallei* is an important pathogen, we therefore recommend BS on day 2 of incubation as the best trade-off between time to detection and pathogen growth rate.

The potential benefits of early BS are not limited to earlier detection of pathogens; it also leads to earlier recovery of colonies, which can be immediately used for supplementary testing. Fast results of antibiotic susceptibility testing, for example, have a large impact on patient management. These tests depend on colony growth; therefore, shorter time to colonies leads directly to faster actionable results, clinical decision-making, and directed treatment. This is particularly important in a context of increased multidrug-resistant pathogens and endemic *B. pseudomallei* infection, as empirical treatment may not adequately cover these pathogens. As shown in Figures 3 and 4, the effect of BS was more pronounced on time to colonies than on time to detection of pathogens. Performing a BS on all bottles adds a substantial workload, and thus costs to a blood culture system. The replacement of an anaerobic by an aerobic bottle doubled the workload for BS, as this practice is only recommended for aerobic bottles. Pathogen growth rates of blood cultures are low (recommended pathogen growth rate is 6–12%, implying the large majority of sampled cultures do not show growth); therefore, most of the work associated with performing BSs does not result in faster recovery of pathogens. Faster pathogen detection must therefore be balanced against increased labor costs. Labor costs are generally lower in most low-resource than in high-resource settings; therefore, BS is probably more efficient in these settings. The added value of increased and faster recovery of pathogens with BS should be balanced against this increased workload and an increased risk of contamination of the blood culture bottles.

![Cumulative time-to-detection for all pathogens excluding B. pseudomallei](image)

**Figure 4.** Cumulative time to detection (first sign of growth) and time to colonies for *Burkholderia pseudomallei*, comparing the three periods of surveillance. Statistically significant differences between periods of surveillance are indicated with asterisks (* = P < 0.05; ** = P < 0.001). This figure appears in color at www.ajtmh.org.
The change in blood culture bottles (FA FAN to FA Plus Bact/ALERT bottles) could have contributed to faster growth, although there was no significant change in visual signs of positivity on day 1 (the only day not affected by changes in BS timing); cumulative first growth on day 1 increased only slightly from 33.1% to 36.7% after implementation of the new bottles ($P = 0.13$). However, cumulative time to detection by day 2 increased significantly for pathogens not including B. pseudomallei in period 2, indicating a possible bottle effect as BS could not have impacted growth on day 2 yet (Figure 4). Moreover, the presence of resins in the FA Plus blood culture medium may have had an impact on growth rates of blood cultures taken under antimicrobial therapy, as resins have been shown to successfully inhibit the effect of antibiotics present in serum. $^{32-34}$ This is indeed suggested by the fact that prior antibiotic therapy had a negative effect on the growth rate in period 1 (odds ratio [OR] of 0.78, $P < 0.001$), whereas cultures under antibiotic therapy were more likely to show growth in periods 2 and 3, after the implementation of the resin-containing bottles (OR: 1.17 and 1.13, respectively, $P > 0.059$) (Table 4).

Another effect of the change in bottles could be related to visual detection of growth. The bottles are currently used off-label, as they are specifically constructed for use in an automated incubator. Other studies have used a similar approach, but also with bottles not containing resins. $^{35}$ Visual growth characteristics such as turbidity and puff balls can be less clearly visible in resin-containing bottles. $^{36}$ The observed decrease in pathogen growth rate could be related to less efficient detection of growth, especially of organisms that cause little visible change in the color indicator.

As compared with surveillance period 1, volume of blood culture bottles was higher, and significantly more bottles were correctly filled in periods 2 and 3. A monthly feedback system was implemented in 2014, whereby quality indicators such as bottle weight and contamination rate were presented to nursing teams, creating increased awareness of correct blood culture sampling procedures. The success of this feedback system is further highlighted by the observation that these filling volumes compare favorably with those reported from high-income countries, despite the limitations inherent to low-resource settings. $^{37,38}$ Volume did however not appear to have an impact on growth rates. This could be explained by the assumption that sampling high volumes of blood from severely ill patients is more difficult and less easily obtained; these patients are however more likely to suffer from BSI. No information on disease severity was available for this study.

Even with early BS, time to detection of many pathogens, most notably non-fermenters such as B. pseudomallei, remains behind that of automated systems. $^{9,39-41}$ A possible explanation for this observation is faster detection of growth by the algorithms in the automate software than visual assessment of the indicator. However, improved time to detection may also be related to faster growth. Blood culture automates agitate the blood cultures continuously during incubation, which has been shown to increase recovery of pathogens, probably due to increased oxygenation. $^{42-46}$ Moreover, temperature in automated incubators is presumably more stable than that in a conventional incubator, which has to be opened in its entirety each time a bottle is added or removed.

The limitations of this study are its observational nature and the simultaneous implementation of multiple laboratory changes. Causal effects of one factor on outcomes such as yield and speed of detection are therefore hard to pinpoint. Moreover, other changes in hospital policy and in underlying population trends could have influenced our results. In the case of time to detection and time to colonies, however, there is strong evidence that advancing the BS led to faster recovery of pathogens. The shifting proportions of different key pathogens could have been reflections of changing underlying epidemiology, although there are indications that the laboratory changes affected detection of B. pseudomallei. Another limitation was the lack of information on time delays (in hours) between the sampling of blood cultures and the time of incubation, which could have provided an alternative hypothesis for poor detection rates of B. pseudomallei. We found no relation between incubation delay and growth; however, delay was only measured in days, precluding demonstration of the effect of more subtle time delays. It has been shown that non-fermenting Gram-negative organisms can go into a bacteriostatic phase when pre-incubated, inhibiting their detection in automated systems. $^{47}$ However, we believe this problem is less important in this setting, as no automated detection was used, and pre-incubation is unlikely to impact yield of BS or visual detection. Moreover, the lack of terminal subculture (additional subculture at the end of the incubation period) during this study precludes conclusions on how many pathogens were missed by advancing the BS. It was therefore decided to temporarily introduce a terminal subculture in SHCH to revalidate the BS procedure.

In conclusion, this study demonstrates that rigorous follow-up of blood cultures and quality indicators is possible in low-resource settings and can lead to significant improvements in laboratory and sampling procedures. A faster detection of important pathogens is possible by relatively inexpensive methods such as introducing or advancing a BS early in the period of incubation. This study also demonstrates the possible risks associated to such changes, and the need for continuous monitoring. Based on these study results, it was decided that BS on day 2 was the preferred subculture timing for SHCH, optimizing the balance between time to detection and pathogen recovery. Despite the best efforts, time to detection of these manual blood cultures systems still lags behind that of automated blood culture systems. Robust, affordable detection systems to further shorten time to detection for low-resource settings are warranted.

Received April 6, 2020. Accepted for publication September 27, 2020.

Acknowledgments: We are grateful to all study participants, laboratory staff, nursing and clinical staff, data manager, and management staff at SHCH, without whom our research would not have been possible.

Financial support: This work was funded by the Belgian Directorate of Development Cooperation (DGD) through the Fourth Framework Agreement between the Belgian DGD and the Institute of Tropical Medicine, Belgium.

Data availability: The database for this manuscript will be made open access. Access requests for ITM research data can be made to ITM’s central point for research data access by means of submitting the completed Data Access Request Form. These requests will be reviewed for approval by ITMs Data Access Committee (https://www.itg.be/E/data-sharing-open-access).

Disclosure: Ethical approval was granted as part of the study: “Surveillance of antimicrobial resistance among consecutive blood culture isolates in tropical settings, V4.0” (IRB ITM 613/08, EC UZA 8/20/96,
This is an open-access article distributed under the terms of the Creative Commons Attribution (CC-BY) License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

REFERENCES
1. Fleischmann-Struzek C, Goldfarb DM, Schlattmann P, Schlapbach LJ, Reinhardt K, Kissoon N, 2018. The global burden of paediatric and neonatal sepsis: a systematic review. Lancet Respir Med 6: 168–170.
2. Vincent JL et al., 2014. Assessment of the worldwide burden of critical illness: the Intensive Care over Nations (ICON) audit. Lancet Respir Med 2: 380–386.
3. Fleischmann C, Scherag A, Adhikari NKJ, Hartog CS, Tsaganos T, Schlattmann P, Angus DC, Reinhardt K, International Forum of Acute Care Trials, 2016. Assessment of global incidence and mortality of hospital-treated sepsis current estimates and limitations. Am J Respir Crit Care Med 193: 259–272.
4. Peker N, Couto N, Sinha B, Rossen JW, 2018. Diagnosis of bloodstream infections from positive blood culture samples and directly from blood samples: recent developments in molecular approaches. Clin Microbiol Infect 24: 944–955.
5. Ombrelet S et al., 2019. Clinical bacteriology in low-resource settings: today’s solutions. Lancet Infect Dis 18: e248–e258.
6. Petti CA, Polage CR, Quinn TC, Ronald AR, Sander MA, 2006. Laboratory medicine in Africa: a barrier to effective health care. Clin Infect Dis 42: 377–382.
7. Otu A, Elston J, Nsutebu E, 2015. Sepsis in Africa: practical steps to stem the tide. Pan Afr Med J 21: 323.
8. Peeters M et al., 2019. Slow growth of Burkholderia pseudomallei compared to other pathogens in an adapted blood culture system. J Med Microbiol 68: 1195–1199.
9. Tiangpitayakorn C, Songsivilai S, Piyasangthong N, Dharakul T, Wattanagoon Y, White N, 1990. Blood culture techniques for detecting typhoidal serovar Paratyphi A infections, Phnom Penh, Cambodia. J Clin Pathol 53: 233–238.
10. Mahidol Oxford Tropical Medicine Research Unit, 2016. Melioidosis Info: Databases. Available at: https://www.melioidosis.info INFO.aspx?pageID=107. Accessed October 20, 2020.
11. Baron E, Dunne W, Weinstein M, Weich D, Wilson D, Yagupsy P, 2005. Cumitech 1C, blood culture IV. Baron E, ed. Cumitech 1C. Washington, DC: American Society for Microbiology.
12. Ven a A, Muñoz P, Alcalá L, Fernandez-Cruz A, Sanchez C, Valero M, Bouza E, 2015. Are incidence and epidemiology of anaerobic bacteremia really changing? Eur J Clin Microbiol Infect Dis 34: 1621–1629.
13. Barthlett JG, Dick J, 2000. The controversy regarding routine anaerobic blood cultures. Am J Med 108: 505–506.
14. Grohs P, Mainardi JL, Podgielan I, Hanraux X, Eckert C, Buu-Hoi A, Varon E, Gutmann L, 2007. Relevance of routine use of the anaerobic blood culture bottle. J Clin Microbiol 45: 2711–2715.
15. Gajdács M, Urbán E, 2020. Relevance of anaerobic bacteremia in adult patients: a never-ending story? Eur J Microbiol Immunol 10: 64–75.
16. Urbán E, 2012. Five-year retrospective epidemiological survey of anaerobic bacteremia in a university hospital and review of the literature. Eur J Microbiol Immunol (Bp) 2: 140–147.
17. Gajdács M, Urbán E, 2020. Relevance of anaerobic bacteremia in adult patients: a never-ending story? Eur J Microbiol Immunol 10: 64–75.
18. Leber AL, 2016. New observations on the biology of Burkholderia pseudomallei. Clin Infect Dis 62(5): 612–618.
19. Cutnell J, Johnson K, 1998. Physics, Fourth Edition. Hoboken, NJ: Wiley.
20. Bardin JD, Tekippe M, 2016. Diagnosis of bloodstream infections in children. J Clin Microbiol 54: 1418–1424.
21. Mahidol Oxford Tropical Medicine Research Unit, 2016. Melioidosis Info: Databases. Available at: https://www.melioidosis.info INFO.aspx?pageID=107. Accessed October 20, 2020.
22. Baron E, Dunne W, Weinstein M, Weich D, Wilson D, Yagupsy P, 2005. Cumitech 1C, blood culture IV. Baron E, ed. Cumitech 1C. Washington, DC: American Society for Microbiology.
23. Ven a A, Muñoz P, Alcalá L, Fernandez-Cruz A, Sanchez C, Valero M, Bouza E, 2015. Are incidence and epidemiology of anaerobic bacteremia really changing? Eur J Clin Microbiol Infect Dis 34: 1621–1629.
24. Bartlett JG, Dick J, 2000. The controversy regarding routine anaerobic blood cultures. Am J Med 108: 505–506.
25. Grohs P, Mainardi JL, Podgielan I, Hanraux X, Eckert C, Buu-Hoi A, Varon E, Gutmann L, 2007. Relevance of routine use of the anaerobic blood culture bottle. J Clin Microbiol 45: 2711–2715.
26. Gajdács M, Urbán E, 2020. Relevance of anaerobic bacteremia in adult patients: a never-ending story? Eur J Microbiol Immunol 10: 64–75.
27. Leber AL, 2016. New observations on the biology of Burkholderia pseudomallei. Clin Infect Dis 62(5): 612–618.
28. Gajdács M, Urbán E, 2020. Relevance of anaerobic bacteremia in adult patients: a never-ending story? Eur J Microbiol Immunol 10: 64–75.
29. Cutnell J, Johnson K, 1998. Physics, Fourth Edition. Hoboken, NJ: Wiley.
30. Bardin JD, Tekippe M, 2016. Diagnosis of bloodstream infections in children. J Clin Microbiol 54: 1418–1424.
cultures using BacT/ALERT 3D (automated) blood cult 20: 530–533.
40. Biondi EA, Mischler M, Jerardi KE, 2014. Blood culture time to positivity in febrile infants with bacteremia. JAMA Pediatr 168: 844–849.
41. Lambregts MMC, Bernards AT, Van Der Beek MT, Visser G, De Boer MG, 2019. Time to positivity of blood cultures supports early re-evaluation of empiric broad-spectrum antimicrobial therapy. PLoS One 14: e0208819.
42. Hawkins BL, Peterson EM, de la Maza LM, 1986. Improvement of positive blood culture detection by agitation. Diagn Microbiol Infect Dis 5: 207–213.
43. Arpi M, Larsen SO, 1992. Comparison of detection speed and yield in agitated and non-agitated aerobic blood culture bottles. APMIS 100: 1061–1065.
44. Prag J, Nir M, Jensen J, Arpi M, 1991. Should aerobic blood cultures be shaken intermittently or continuously? Appl Microbiol 99: 1078–1082.
45. Ellner PD, Kiehn TE, Beebe JL, McCarthy LR, 1976. Critical analysis of hypertonic medium and agitation in detection of bacteremia. J Clin Microbiol 4: 216–224.
46. Weinstein MP, Mirrett S, Reimer LG, Reller LB, 1989. Effect of agitation and terminal subcultures on yield and speed of detection of the oxoid signal blood culture system versus the BACTEC radiometric system. J Clin Microbiol 27: 427–430.
47. Klaerner HG, Eschenbach U, Kamereck K, Lehn N, Wagner H, Miethke T, 2000. Failure of an automated blood culture system to detect nonfermentative gram-negative bacteria. J Clin Microbiol 38: 1036–1041.