AACC Guidance Document on the Clinical Use of Procalcitonin

Authors:

1Allison Chambliss, PhD, DABCC, FAACC
Department of Pathology
University of Southern California
Los Angeles, CA

1Khushbu Patel, PhD, DABCC, FAACC
Department of Pathology and Laboratory Medicine
Children’s Hospital of Philadelphia
Philadelphia, PA

Jessica M Colón-Franco, PhD, DABCC, FAACC
Department of Laboratory Medicine
Cleveland Clinic
Cleveland, OH

Joshua Hayden, PhD, DABCC, FAACC
Department of Laboratories
Norton Healthcare
Louisville, KY

Sophie E, Katz, MD, MPH
Division of Infectious Diseases, Department of Pediatrics
Vanderbilt University Medical Center
Nashville, TN

Emi Minejima, PharmD
Department of Clinical Pharmacy
University of Southern California School of Pharmacy
Los Angeles, CA

Wesley Self, MD
Department of Emergency Medicine
Vanderbilt University Medical Center
Nashville, TN

2Alison Woodworth, PhD, DABCC, FAACC
Department of Pathology and Laboratory Medicine
University of Kentucky Medical Center
Lexington, KY

1These authors contributed equally to this report

2To whom correspondence should be addressed: alison.woodworth@uky.edu
**List of abbreviations**

PCT, procalcitonin  
LRTI, lower respiratory tract infection  
ICU, intensive care unit  
RCT, randomized controlled trial  
CRP, C-reactive protein  
ED, emergency department  
CAP, community-acquired pneumonia  
AECOPD, acute exacerbation of chronic obstructive pulmonary disease  
VAP, ventilator-associated pneumonia  
GOLD, Global Initiative for Chronic Obstructive Lung Disease  
AUC, area under curve  
ARI, acute respiratory infection  
POCT, point-of-care testing  
IFCC, International Federation of Clinical Chemistry and Laboratory Medicine  
PCT-WG, IFCC Working Group on Standardization of Procalcitonin Assays
Table of Contents

Abstract ........................................................................................................................................... 5
Background ....................................................................................................................................... 5
Content ............................................................................................................................................. 5
Summary .......................................................................................................................................... 5

Introduction ..................................................................................................................................... 6

Can PCT results be utilized to inform treatment decisions in both initiation and cessation of antimicrobial therapy in adult patients with sepsis or respiratory infections? ........................................................................ 6
  Antibiotic initiation in the critically ill ............................................................................................ 6
  Antibiotic cessation in the critically ill ............................................................................................ 8
  Other biomarkers evaluated to guide antibiotic initiation or cessation in the critically ill .......... 10
  Antibiotic initiation in respiratory tract infections ........................................................................ 11
  Antibiotic cessation in respiratory tract infections ....................................................................... 13
  Other biomarkers evaluated to guide antibiotic initiation or cessation in respiratory tract infections ..................................................................................................................................................... 15

Is PCT an accurate predictor of outcomes (e.g., mortality, respiratory failure, shock) in adult populations? .......................................................................................................................................... 17
  Outcomes from single PCT measurements .................................................................................. 18
  Outcomes from sequential PCT measurements and PCT clearance ........................................... 19
  Outcomes for patients presenting to the emergency department .................................................. 20
  Outcomes differ by patient diagnoses, sepsis definitions, and study populations ....................... 22
  Outcomes other than 28/30-day mortality .................................................................................... 23
  Other biomarkers evaluated to predict outcomes in patients with sepsis and/or respiratory tract infections .............................................................................................................................................. 24

Can PCT results be utilized to inform treatment decisions in both initiation and cessation of antimicrobial therapy in neonatal and pediatric patients with sepsis or respiratory infections? .......................................................................................................................... 28
  PCT for detecting bacterial infections and initiating antibiotics in pediatric and neonatal patients ......................................................................................................................................................... 29
  Antibiotic cessation in neonatal and pediatric populations ......................................................... 31

Is PCT an accurate predictor of outcomes (e.g., mortality, respiratory failure, shock) in pediatric populations? ...................................................................................................................................... 33
  Outcomes from single PCT measurements .................................................................................. 33
  Outcomes from sequential PCT measurements ............................................................................ 34

When and how often should PCT be measured? Which cut-off(s) should be used? ....................... 35
  Timing and frequency for antibiotic initiation and cessation in adults ......................................... 36
<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Timing and frequency for outcomes prediction in adults</td>
<td>37</td>
</tr>
<tr>
<td>Change in PCT concentrations over time to predict outcomes in adults</td>
<td>38</td>
</tr>
<tr>
<td>Timing and frequency for antibiotics initiation and cessation in pediatric patients</td>
<td>39</td>
</tr>
<tr>
<td>Timing and frequency for outcomes prediction in pediatric patients</td>
<td>41</td>
</tr>
<tr>
<td>How should PCT be incorporated into antimicrobial stewardship efforts?</td>
<td>42</td>
</tr>
<tr>
<td>What pre-analytical factors affect PCT results and/or interpretation?</td>
<td>44</td>
</tr>
<tr>
<td>Acceptable sample types</td>
<td>44</td>
</tr>
<tr>
<td>Stability and storage</td>
<td>45</td>
</tr>
<tr>
<td>What FDA-approved methods are available to measure PCT, and how do they compare?</td>
<td>45</td>
</tr>
<tr>
<td>BRAHMS PCT assays</td>
<td>46</td>
</tr>
<tr>
<td>Diazyme PCT assays</td>
<td>47</td>
</tr>
<tr>
<td>Other PCT assays</td>
<td>48</td>
</tr>
<tr>
<td>Are clinical decision points (cut-offs) comparable across PCT assays?</td>
<td>48</td>
</tr>
<tr>
<td>What are possible confounding factors for the interpretation of PCT results?</td>
<td>50</td>
</tr>
<tr>
<td>Conclusions</td>
<td>51</td>
</tr>
<tr>
<td>References</td>
<td>53</td>
</tr>
<tr>
<td>Figures</td>
<td>65</td>
</tr>
<tr>
<td>Tables</td>
<td>67</td>
</tr>
</tbody>
</table>
Abstract

Background
Procalcitonin (PCT), a peptide precursor of the hormone calcitonin, is a biomarker whose serum concentrations are elevated in response to systemic inflammation caused by bacterial infection and sepsis. Clinical adoption of PCT in the United States has only recently gained traction with an increasing number of FDA-approved assays and expanded indications for use. There is interest in the use of PCT as an outcomes predictor as well as an antibiotic stewardship tool. However, PCT has limitations in specificity, and conclusions surrounding its utility have been mixed. Further, there is a lack of consensus regarding appropriate timing of measurements and interpretation of results. There is also a lack of method harmonization for PCT assays, and questions remain regarding whether the same clinical decision points may be used across different methods.

Content
This guidance document aims to address key questions related to the use of PCT to manage adult, pediatric, and neonatal patients with suspected sepsis and/or bacterial infections, particularly respiratory infections. The document explores the evidence for PCT utility for antimicrobial therapy decisions and outcomes prediction. Additionally, the document discusses analytical and pre-analytical considerations for PCT analysis and confounding factors that may affect the interpretation of PCT results.

Summary
While PCT has been studied widely in various clinical settings, there is considerable variability in study designs and study populations. Evidence to support the use of PCT to guide antibiotic cessation is compelling in the critically ill and in some lower respiratory tract infections but is lacking in other clinical scenarios, and evidence is also limited in the pediatric and neonatal populations. Interpretation of PCT results requires guidance from multidisciplinary care teams of clinicians, pharmacists, and clinical laboratorians.
Introduction

Can PCT results be utilized to inform treatment decisions in both initiation and cessation of antimicrobial therapy in adult patients with sepsis or respiratory infections?

Multiple meta-analyses have evaluated the utility of PCT to guide decisions on antibiotic initiation for sepsis and respiratory tract infections (3–7). Data about the use of PCT to safely reduce antibiotic treatment for patients with sepsis and lower respiratory tract infection (LRTI) is encouraging. In patients with sepsis admitted to an intensive care unit (ICU), there is relatively strong evidence to support the use of PCT to reduce antibiotic duration (8–14). The data regarding use of PCT in patients with LRTI is less straightforward, with more recent studies demonstrating little benefit with addition of PCT (15–18).

This section will focus on studies that included evaluation of outcomes based on using PCT to inform decisions regarding antibiotic initiation and cessation.

Antibiotic initiation in the critically ill

There were limited number of studies that addressed the impact of PCT utilization on rates of antibiotic initiation in critically ill patients in the ICU. Layios et al. implemented a PCT protocol to guide antibiotic initiation and specifically reported the rate of initiation of antibiotics as an outcome (19). This was a randomized controlled trial (RCT) of five ICUs and 509 total patients and evaluated the rate of initiation of antibiotics, which was not significantly decreased by the availability of PCT compared to standard of care (PCT 62.6% vs control 57.7%, p=0.11). Possible explanations for the lack of benefit included a low rate (25%) of PCT results that were <0.25 ng/mL, critical illness of patients that prompted clinicians to overrule the PCT treatment algorithm, and a low rate (57%) of initiation of antibiotics in the control arm compared to prior studies. Jensen et al. also implemented an antibiotic initiation PCT protocol in a multicenter RCT study of critically ill patients in the ICU in Denmark (N=1,200) (20). In the
PCT arm, an “alert procalcitonin” notification was provided when the initial PCT was ≥1 ng/mL to initiate antibiotics or when subsequent PCT levels were not decreasing by 10% from the previous day to intensify the antibiotic course. In this study, they found the PCT group had a longer antibiotic course by a median of 2 days (PCT median 6 days (IQR 3-11) vs standard of care of 4 days (IQR 3-10). Antibiotic initiation rates were only reported for guideline concordant cases with PCT ≥1 ng/mL which was 82.1% in the PCT group and 82.4% in the standard of care group. As 28-day mortality rates were comparable between the two groups (PCT vs standard of care: HR 0.98, 95% CI 0.83-1.16), the authors concluded that PCT was not effective to guide initiation or escalation of antibiotic therapy over standard of care practices and led to increased use of broad-spectrum antibiotics.

Three studies that had a mixed initiation and cessation PCT protocol reported the rate of antibiotic initiation compared to a control group. In the PRORATA trial, a large randomized clinical trial (N=621) by Bouadma et al. that included critical care patients in France, 28/307 (9%) in the PCT group did not receive antibiotics at study inclusion in accordance to the pre-specified algorithm (<0.5 ng/mL) (12). Eight of the PCT group then proceeded to be given antibiotics within 5 days and 7/8 survived. In comparison, 15/314 (5%) of the control group did not receive antibiotics at inclusion and 8/15 proceeded to be given antibiotics within 5 days, and 1/5 survived. Otherwise, the rate of initiation of antibiotics in patients who had PCT levels that fell outside of the pre-specified range included 65/307 (21%) patients that were initiated on antibiotics when PCT was <0.5 ng/mL and 4/307 (1%) that were not given antibiotics although the PCT was ≥0.5 ng/mL. Additional details on antibiotic exposure outcomes related specifically to antibiotic initiation were not teased out from the total use combining the initiation and cessation protocol. Overall algorithm adherence was 53% in the PCT group. In a second randomized controlled trial with mixed initiation and cessation PCT protocol of ICU patients with severe acute exacerbation of COPD (AECOPD) (N=302), although there was no significant difference in the proportion of those who received antibiotics at baseline (PCT group 58% vs control 62%, p= ns), by day 1 there were
significantly fewer patients in the PCT arm who remained on antibiotics (P<0.001) (16). In a subgroup analysis, those who were not on antibiotic therapy initially and used the PCT to guide antibiotic initiation had a significantly higher 3-month mortality compared with the control group (PCT 19/61, 31% vs control 7/58, 12%, p=0.015). In summary, the data on the utility of PCT to guide initiation of antibiotics in critically ill patients is currently limited and does not show benefit in decreasing antibiotic prescriptions.

Antibiotic cessation in the critically ill

We reviewed five RCTs comparing PCT-guided antibiotic duration with standard antibiotic duration (Suppl. Table 1) with sample sizes that ranged from 110 to 1,546 patients (8–14,21). Overall, 3,757 patients were enrolled in these trials. All studies demonstrated a reduction in antibiotic use with PCT guidance compared to standard antibiotic duration, with reductions in days on antibiotics of about 2 days.

Major limitations to study interpretation included the variation in timing of PCT measurements among studies and variation in the PCT concentration thresholds utilized to drive discontinuation of antibiotic therapy among studies. One study by Hochreiter et al. did not report frequency of PCT measurement or turn-around time for PCT result notification (13). The other seven studies each obtained PCT measurements on enrollment and then at variable time intervals ranging from daily to every 5-7 days until ICU or hospital discharge (8–12,14,21). Absolute cut points for antibiotic discontinuation were 0.25 ng/mL (14), 0.5 ng/mL (9–12) or 1 ng/mL (8,13,21). Most studies also allowed for antibiotic discontinuation based on a relative decrease in PCT concentrations over time. The majority of studies used an 80-90% decrease from peak value to direct antibiotic cessation, although one study used a 50% drop compared to the previous value (8), and another used a 25% decrease from peak concentration (13). Only two studies reported turn-around time for PCT result notifications from
lab personnel to study providers (12,14). In both studies, results were available within 2-3 hours of specimen collection (13). The other 4 studies each obtained PCT measurements on enrollment and then at variable time intervals ranging from daily to every 3 days until ICU or hospital discharge (8–12,14,21). Absolute cut points for antibiotic discontinuation were 0.5 ng/mL (9–12) or 1 ng/mL (8,13,21). Most studies also allowed for antibiotic discontinuation based on a relative decrease in PCT concentration over time. The majority of studies used an 80-90% decrease from peak value to direct antibiotic cessation, although one study used a 50% drop compared to the previous value (8), and another used a 25% decrease from peak concentration (13). Only one study reported turn-around time for PCT result notifications from lab personnel to study providers, and results were available within 2-3 hours of specimen collection (12).

Compliance rates with PCT driven antibiotic cessation algorithms also varied greatly among studies, ranging from 28.7 to 97%. Most often, providers chose to continue antibiotics despite a low PCT level. Non-compliance with algorithm guidance may have skewed study results. However, as noted earlier, all studies demonstrated significant decrease in antibiotic use when using PCT guidance, and non-compliance with algorithms would have more often led to similar results in the PCT and standard of care groups.

In a 2018 meta-analysis, Meier et al. describe 523 patients with positive blood cultures from 13 clinical trials that randomly assigned patients to either PCT-guided treatment or a standard of care group (22). The mean duration of antibiotic therapy was 2.86 days shorter in the PCT guided group compared to the control group (95% confidence interval -4.88 to -0.84 days, \(p=0.006\)), and mortality was again similar between groups (16.6% PCT vs. 20% control, \(p=0.263\)). This study highlights that PCT-guidance for antibiotic duration may be a safe and effective way to decrease antibiotic use among patients with bacteremia.
In summary, all studies evaluated demonstrated a reduction in antibiotic use with PCT guidance compared to standard antibiotic duration. Limitations found across studies included differences in compliance rates for the antibiotic cessation algorithms that may have limited the benefit in certain populations.

Other biomarkers evaluated to guide antibiotic initiation or cessation in the critically ill

Like PCT, CRP is an acute phase reactant, upregulated in response to inflammation (Figure 2). Due to its longer half-life, its utility in antibiotic monitoring is limited. Several studies have investigated its potential to guide antibiotic therapy. For example, in a randomized controlled trial in two ICUs in Brazil, investigators compared PCT vs C-reactive protein (CRP) for the primary outcome of antibiotic duration in critically ill patients with severe sepsis or septic shock (23). Like PCT, CRP is an acute phase reactant, upregulated in response to inflammation (Figure 2). Due to its longer half-life, its utility in antibiotic monitoring is limited. In the PCT arm, discontinuation of antibiotic therapy was recommended if PCT fell to <0.1 ng/mL (if the initial PCT was <1 ng/mL), or if there was a decrease in PCT by 90% from peak (if initial PCT was >1ng/mL). In the CRP arm, discontinuation of antibiotic therapy was recommended if CRP fell to <25mg/L (if the initial CRP was <100 mg/L) or if CRP decreased by 50% from peak (if the initial CRP was >100 mg/L). Of the 94 total patients included, they found the median duration of antibiotics to be similar between the groups (PCT median 7 days vs. CRP 6 days, p=0.06), even after adjusting for severity of illness. Mortality was similar in both groups (21 patients died in each group, p=0.86). Thus, authors concluded that CRP, which costs significantly less than PCT per test, was as useful as PCT in guiding antibiotic cessation.

Currently there is limited data available on the utility of CRP compared to PCT in guiding antibiotic initiation or cessation in a critically ill cohort across different ICU settings. In addition, as CRP is typically a less expensive inflammatory marker than PCT and may be more readily available in clinical...
laboratories, additional cost effectiveness studies are needed to address whether PCT is a more clinically advantageous inflammatory marker to guide antibiotic initiation or cessation in the critically ill.

**Antibiotic initiation in respiratory tract infections**

A 2018 meta-analysis by Schuetz et al. included 26 randomized clinical trials of PCT in respiratory tract infections, of which 13 were conducted in the ICU setting, 11 in emergency departments, and 2 in primary care (24). Infection types included community acquired pneumonia (CAP), acute exacerbation of chronic obstructive pulmonary disease (AECOPD), acute bronchitis, hospital acquired pneumonia, and ventilator-associated pneumonia (VAP). Initiation of antibiotics was significantly reduced overall in the PCT group compared to the control (70% vs 85%, adjusted OR 0.27 (0.24 to 0.32), p<0.0001). Lower antibiotic prescription rates were seen in primary care (PCT 23% vs control 63%, adjusted OR 0.13 (0.09 to 0.18), p<0.0001), emergency department (ED) (PCT 69% vs control 83%, adjusted OR 0.49, p<0.0001), and ICU settings (PCT 92% vs control 99%, 0.02 (0.01 to 0.05), p<0.0001). Antibiotic prescription rates were similarly lower when analyzed by infection type for all included infection types except for VAP, where antibiotic initiation was 100% in both the PCT and control arms.

Since the publication of the meta-analysis by Schuetz et al. (24), Huang et al. published the ProACT study, a 14 center, randomized clinical trial (15) evaluating PCT use in acute LRTI of 1656 adult patients in the US. This study excluded severely ill patients requiring endotracheal intubation or intravenous vasopressors. Of the patients evaluated in the ED, there was 72.9% PCT algorithm adherence, with highest algorithm adherence for acute bronchitis (82.4%) and the lowest in CAP (39.4%). There was no difference in the primary outcome of antibiotic exposure during the first 30 days between the PCT group vs. control group (mean antibiotic days 4.2 vs 4.3 days, difference -0.05 days, 95% CI -0.6 to 0.5, p=0.87). The secondary outcome of percentage of patients receiving an antibiotic
prescription in acute bronchitis in the ED was significantly lower in the PCT group (17.3% vs 32.1%; risk difference, -14.8%, 99.86% CI -28.5 to -1.1). The authors noted that a study limitation was that PCT information was not available to all prescribers prior to when decisions about antibiotic initiation could be made, thus the true effect of PCT information to guide decisions on whether to initiate antibiotics may be lacking. Also of note is the lower rate of antibiotic prescription even in the control arm in the ED (38.7%) compared to control arms in the studies in the meta-analysis by Schuetz et al. (85%) (24).

Patients presenting with acute heart failure represent a challenging population to evaluate the need for antibiotic therapy for respiratory tract infections due to diagnostic uncertainty. Mockel et al. evaluated whether PCT-guided initiation of antibiotics in a study population of patients with suspected or confirmed heart failure could provide benefit by decreasing unnecessary starts of antibiotics (25). In heart failure patients presenting to the ED with primary symptom of dyspnea, antibiotic initiation was recommended if PCT was >0.2 ng/mL. In this RCT of 742 total patients, the initiation rate of antibiotic was similar between the study groups (PCT-guided 18% vs. standard of care 14%, p=0.145) and the primary outcome of all-cause 3-month mortality was similar (PCT-guided 14% vs. standard of care 6.6%, 90% CI -0.3 - 13.5%). PCT algorithm adherence was 83% in the PCT group. The overall rate of pneumonia in this study was only 7.5%, which the authors predict was likely due to study clinicians avoiding randomization of patients with a high suspicion of pneumonia, where immediate initiation of antibiotics was warranted without having to wait for the PCT result. Thus, the authors concluded that PCT-guided initiation of antibiotics was not more effective than standard of care in the rates of antibiotic utilization.

AECOPD represents another population that poses a diagnostic challenge in LRTI. A meta-analysis by Ni et al. included 23 studies of PCT usage in patients presenting with severe AECOPD with the primary outcome of length of stay and treatment failure (26). Six RCTs involving a total of 942 patients found no difference in treatment failure rates (RR 0.85, 95% CI 0.66-1.09) or length of stay (weighted mean difference = -0.1, 95% CI –0.98 - 0.79) between the PCT-guided group and the control group. PCT-
prepared treatment significantly reduced the antibiotic prescription rate by 34% (RR 0.66, 95% CI 0.62-0.71).

Bremmer et al. evaluated patients with low PCT (<0.25 ng/mL) in non-critically ill AECOPD in a retrospective study on the effects on 30-day all-cause hospital readmissions (27). This study excluded ICU patients, patients with immunocompromising conditions, patients on mechanical ventilation, and patients with pneumonia. Comparing patients who received ≤24h vs >24h of antimicrobial therapy, there was no difference in all-cause 30-day readmission (15.5% vs. 17.4%, p=0.63) and COPD-related 30-day readmission rates (11.2% vs. 12.3%, p=0.743), concluding that in a non-critically ill cohort of AECOPD admissions, antibiotics may be withheld safely utilizing PCT guidance.

In summary, PCT-guided therapy can significantly decrease antibiotic initiation without compromising safety in LRTI. The clinical benefit was more consistently seen in lower acuity patients such as those in the primary care setting with CAP, AECOPD, or acute bronchitis. More data is needed to support its use in antibiotic initiation for critically ill patients or patients with immunocompromising conditions. Data is currently not available to support the use of PCT in patients with the suspicion of VAP to guide antibiotic initiation decisions.

**Antibiotic cessation in respiratory tract infections**

There were 11 RCTs comparing PCT-guided therapy with standard antibiotic therapy for treatment of patients with LRTI, including CAP, COPD, non-pneumonia LRTI, and VAP (Suppl. Table 2) (15,17,18,28–32). Sample sizes ranged from 45 to 1,656 patients; only one study had fewer than 100 patients. There were ten RCTs comparing PCT-guided therapy with standard antibiotic therapy for treatment of patients with LRTI, including CAP, COPD, non-pneumonia LRTI, and VAP (Suppl. Table 2) (15,18,28–32). Sample sizes ranged from 101 to 1,656 patients. Overall, 3,905 patients were enrolled in these trials. There were an additional three retrospective studies evaluating PCT for antibiotic discontinuation (33,34) and one
prospective cohort study (35). The impact of PCT measurement on antibiotic use varied among studies, with 6/11 demonstrating reduction in antibiotic use with PCT-guidance. None of the four RCTs published since 2016 have demonstrated benefit in using PCT to decrease antibiotic use in patients with RTIs (15–18). Possible reasons for the lack of benefit seen may be due to a different population being investigated (severe AECOPD vs mild to moderate AECOPD) (16), lower algorithm adherence rates (17) compared to prior studies (30), or more contemporary studies including antimicrobial stewardship in the standard of care arm that reduced the antibiotic duration.

Timing of PCT measurements and frequency with which PCT was measured varied among studies. The first measurement was obtained at or within 24 hours of enrollment in all studies. The timing of the next measurement varied in all studies, ranging from 12 h to 5 days after the first. No studies required PCT measurements after day 10, and most stopped testing on day 7. Of the studies that mentioned timing from sample collection to provider notification of results, most were available within one hour. Cut points used to discontinue antibiotics also varied across studies. Ten of the 11 studies used an absolute value of 0.25 ng/mL to recommend antibiotic discontinuation (15–18,29,30,36–38). The remaining study used an absolute value of 0.5 ng/mL (39). Five of the studies utilizing LRTI patients also allowed for antibiotic discontinuation based on a relative decrease in PCT concentration of ≥80% (18,30,39,40) or ≥90% (16) when compared to the peak PCT concentration or the concentration at randomization. Compared to studies using PCT for antibiotic discontinuation in sepsis, compliance rates with PCT algorithm guidance were higher in pneumonia studies, ranging from 61% to 85%. Compliance rates were not reported in two of the studies (16,18). A more detailed discussion of PCT collection timing and frequency is included in a later section of this document.

All included RCTs published prior to 2011 demonstrated reduction in antibiotic use with PCT guidance and did not demonstrate increase risk of mortality (if included as an outcome) (30,37–39). However, RCTs published since 2016 have not demonstrated reduced antibiotic use with PCT guidance.
compared to control (15–17,36,40). Proposed reasons for this decrease in effect size include shorter baseline treatment durations for most patients with LRTI and low algorithm compliance rates. Two recently-published retrospective studies with more than 300 patients in each study demonstrated significantly shorter antibiotic duration in patients with LRTI in whom a PCT-guided antibiotic cessation algorithm was followed (33,34).

Schuetz et al. published in 2017 a Cochrane review of 26 randomized clinical trials on respiratory tract infections with a patient level meta-analysis of 6708 participants (5). All cause 30-day mortality was significantly lower with the PCT-guided therapy (adjusted OR 0.83, 95% CI 0.70 to 0.99, p=0.037).

Reduction in total antibiotic exposure (mean 8.1 days compared to 5.7 days, regression coefficient -2.43 days (95% CI -2.71 to -2.15, p<0.001) was observed, although the rate of initiation of antibiotics was not reported.

In summary, conclusions regarding the impact of PCT measurement on antibiotic cessation in LRTI are mixed among studies. Earlier studies and meta-analysis show overall decrease in antibiotic use with PCT, while the more recent RCTs have shown no difference relative to standard of care without PCT.

Other biomarkers evaluated to guide antibiotic initiation or cessation in respiratory tract infections

There are limited studies available that evaluated the performance of other biomarkers (e.g., CRP) compared to PCT on antibiotic prescribing patterns for LRTI. However, CRP has been evaluated in respiratory tract infections for its utility to guide antibiotic prescribing decisions, most notably in the primary care settings, where it has demonstrated decreased antibiotic prescriptions (41–43). In an outpatient setting, point-of-care CRP testing led to significantly fewer antibiotic prescriptions for acute LRTI and rhinosinusitis as compared to standard of care (43.4% vs 56.6%, relative risk 0.77, 95% CI 0.56-0.98) in the UK (44).
Butler et al. evaluated CRP to guide antibiotic prescription decisions in AECOPD in a multi-center, randomized controlled trial (n=653) (45). A CRP point-of-care test was performed at presentation in the intervention arm, with guidance that for CRP <20 mg/L, antibiotics were unlikely to be beneficial, for CRP 20-40 mg/L, antibiotics may be beneficial in the presence of purulent sputum, and for CRP >40mg/L, antibiotics were likely to be beneficial. The availability of rapid CRP results significantly decreased the number of patients who received an antibiotic prescription (47.7% vs 69.7%, adjusted OR 0.31, 95% CI 0.21-0.45). Similarly, Prins et al. found that using CRP significantly decreased antibiotic prescriptions compared to using Global Initiative for Chronic Obstructive Lung Disease (GOLD) guidelines (based on symptoms of purulent sputum) in their randomized controlled trial of 220 adults in the Netherlands (CRP 31.7% vs GOLD 46.2%, p=0.028) (46). However, neither study directly compared CRP vs. PCT performance.

Additional data is needed to compare CRP vs PCT for antibiotic initiation or cessation in other LRTI, such as pneumonia, and in the inpatient setting. As CRP is typically a less expensive inflammatory marker test than PCT, cost-effectiveness studies comparing the two markers would be helpful to guide laboratories considering implementation of PCT.

**Key summary points**

- In patients admitted to the ICU, PCT should be used to reduce antibiotic duration. Data does not support using PCT to guide initiation of antibiotics in these patients.
- In patients with LRTI, PCT may be used to safely reduce antibiotic exposure and duration, but there is less evidence to support this recommendation, particularly in VAP.
- Studies to date have shown significant variation in PCT testing algorithms in terms of cut points and timing of PCT measurements.
• Other inflammatory markers, such as CRP, have shown benefit in decreasing antibiotic initiation in primary care settings and AECOPD, although conclusions are mixed. Direct comparison of CRP and PCT is still lacking in LRTI. As CRP is typically less expensive and may be a more widely available biomarker than PCT, additional cost-effectiveness studies are also needed.

Is PCT an accurate predictor of outcomes (e.g., mortality, respiratory failure, shock) in adult populations?
PCT is upregulated in response to proinflammatory signals and its concentrations increase with increased disease severity (47). Concentration decreases over time are associated with disease recovery, while consistently elevated or increasing PCT concentrations are a signal of persistent or more severe illnesses. Thus, researchers have studied the ability of PCT to predict outcomes in a variety of disease states. Among the most commonly studied outcomes are mortality and disease progression.

The literature investigating the potential utility of PCT in predicting outcomes is significant. Mortality is an important outcome among critically ill patients given that approximately 270,000 septic patients die each year in the United States alone (48). With recent updates to FDA approved intended uses of these assays, we chose to look more closely at publications since 2010 with at least 100 patients enrolled that investigated the correlation between PCT concentrations and mortality. 14 of these looked at 28/30-day all-cause mortality (Suppl. Table 3). Among the 28-day mortality studies, the patient populations ranged from ED, ICU and/or hospital in-patients with sepsis, respiratory tract infections, and other infections. Two of these studies were separate large meta-analyses of >3,000 patients each looking at PCT in septic and RTI patients (49,50). One of these studies was a randomized control trial, however the investigation of 28-day mortality was a secondary analysis (51).
Outcomes from single PCT measurements

All studies demonstrated an association between initial PCT concentrations measured at presentation and/or evolution over time (discussed below) and 28/30-day mortality (Suppl. Table 3). Most of the studies show that, independent of patient population, PCT is significantly higher in patients that will go on to die within 28/30 days compared to those who survived (8,52,52–56). Areas under the ROC Curve (AUCs) for initial PCT measurements to predict 28 day mortality range from 0.56 – 0.82 among studies where the time of PCT measurement was clearly delineated (8,49–52,55,57–59). In their meta-analysis, Liu et al. demonstrated in a diverse patient population that a single PCT measurement had a moderate ability to predict mortality. The area under the summary ROC curve was 0.73 (0.69 – 0.77) (n=13 studies) (50). Follow up PCT measurements in patients with signs of infection at 72 hours after the initial test were also significantly higher in patients that died (59). However, in patients with autoimmune disease and sepsis, PCT concentrations at 72 hours were not significantly different in survivors compared to non-survivors (60). This group also looked at the correlation between PCT peak concentration and mortality, and found variable results depending on patient population (60). Similarly, in their large multicenter prospective observational study with over 1700 septic shock patients, Ryoo et al. were unable to demonstrate that elevated PCT, measured at presentation to the ED, was significantly predictive of 28 day mortality (61). The authors attribute this apparent discrepancy, compared to similar studies, to PCT being measured too soon after presentation and their patient population (ED versus ICU). In summary, a single PCT measurement at presentation is associated with higher mortality rates, however patient population, time of PCT measurement, and other factors confound study results and limit the utility of a single PCT measurement to predict mortality.
Outcomes from sequential PCT measurements and PCT clearance

Procalcitonin has a distinct expression pattern following an infectious insult (Figure 2). Further, PCT expression persists throughout infection and decreases as the infection is cleared (47). Therefore, many have advocated monitoring PCT kinetics/concentration change over time as a prognostic marker in patients with infections (47). Sometimes referred to as the PCT delta, this change is PCT over time has been associated with clinical outcomes. Specifically, a Procalcitonin decrease over time, referred to as PCT clearance, suggests that a patient is responding to antimicrobial or other therapy. By contrast persistent PCT expression over time, also called PCT non-clearance, is associated with poor outcomes in critically ill and septic patients (50).

Among recent studies, five demonstrated that a lack of PCT clearance over time was a good predictor of 28/30 day mortality (50,51,60,62,63). Of these, one was a large meta-analysis where a subset analysis of 9 studies with 868 septic patients demonstrated that lack of PCT clearance, defined by a decrease in PCT of at least 25% (range 25 – 70%) in 48 hours – 7 days, could predict mortality (28 day and hospital) with a summary AUC of 0.79 (0.75 – 0.83), and pooled relative risk of mortality was 3.05 (2.35 – 3.95) (50). The authors concluded that while the prognostic utility of the initial PCT value is limited, PCT non-clearance better predicts mortality and its performance is superior to following clearance of other biomarkers like lactate. Subsequently, one essential study of 13 US medical centers including 858 patients admitted to the ICU with severe sepsis or septic shock from the ED or other hospital locations, demonstrated that a lack of PCT decrease over time was a good predictor of 28/30 day mortality. The study found that although PCT was higher in non-survivors (mean 5.2 (95% CI 3.9-7.0) vs 3.4 ng/mL (95% CI 2.8-4.0), p<0.02), this baseline value was a poor predictor of 28-day mortality (AUC 0.56; 95% CI 0.51-0.60) (62). In patients in which PCT did not decrease by >80% between baseline and day 4, 28-day mortality was 20.0% (hazard ratio 1.97, multivariate adjustment, p=0.009), twice as high as the group with this decline (p=0.001). At this cut-off, sensitivity was 77% (p5% CI (69-85) and specificity was 39% (95% CI 35-43) with similar performance regardless of the patients being at the ICU.
or not at day 4. Notably, ICU residency by day 4 was a strong independent predictor of mortality (hazard ratio 2.69, multivariate adjustment, p<0.0001), with much higher mortality than among those discharged to the hospital floors (26 vs 9%). In a secondary analysis comparing PCT at baseline and day 1, PCT increased by 30% (95% CI 15-47) and by 0% (95% CI -7-6) for those who died and survived, respectively (p<0.0001), and mortality increased 3-fold in patients with an increase in PCT compared to a decrease in PCT (29 versus 12%, p<0.0001). Mortality was approximately 3 times higher if PCT did not decrease by 80%, regardless of the initial PCT concentration being above or below 2 ng/mL. This study demonstrated that both short and longer serial approach/PCT clearance are stronger prognosticators than an initial single PCT measurement. Finally, in their large RCT of 1089 patients with severe sepsis or septic shock, Elke et al. demonstrated that a PCT decrease < 20% from baseline to day 1 was associated with a significantly higher 28-day mortality rate compared to those in whom PCT declined by > 20%. In this same patient population, there was a significantly lower risk of mortality for those in whom PCT declined by > 50% between days 0 and 4 compared to those with a < 50% PCT decline. Although the parameters of the PCT delta calculation are not standardized, the lack of PCT clearance over time does predict 28-day mortality in diverse patient populations.

Outcomes for patients presenting to the emergency department

Some differences are noted in the utility of PCT among different patient populations. In general, there are far fewer studies in ED patients looking at the ability of initial PCT concentrations to predict mortality. The meta-analysis by Liu et al. only included 3 studies in the ED, which limited their ability to evaluate PCT's ability to predict mortality in ED patients (50). An individual patient meta-analysis including 2,605 ED patients concluded that PCT measurement at presentation predicted mortality (AUC 0.67, p<0.001, OR 1.82), and correlated with treatment failure (i.e. death, ICU admission, re-hospitalization, and complications or recurrent or worsening infection within 28 days) (AUC 0.64,
p<0.01, OR 1.85) (49). In reality, this performance is modest for predicting treatment failure and mortality and awaits confirmation by other studies. In their meta-analysis of septic patients (ED and hospitalized), while pooled mean PCT concentrations measured on days 1 and 3 were both significantly different between survivors and non-survivors and able to predict mortality, the correlation was stronger once ED patients were excluded (64). In their multicenter trial, Saeed et al. investigated several biomarkers’ abilities to predict 28-day mortality when measured at initial assessment in ED patients with suspected infections. PCT measurement at initial ED evaluation showed moderate ability to predict 28-day mortality (AUC = 0.72 – 0.75), which is similar to other studies in diverse patient populations (52). Of note, the mortality rate in this patient population was low at ~7%. In their multicenter, multinational ED cohort study, Sager et al. demonstrated that PCT concentrations, when measured during the ED stay, predicted 30-day mortality (55). In contrast, in a third study looking at ED patients with septic shock (20.7% mortality), Ryoo et al. showed that initial PCT measurement was not an independent predictor of 28-day mortality. The authors suggest that their findings may differ from other studies because all PCT measurements were collected prior to initiation of antimicrobial therapy (61). Unlike Ryoo’s population, most other studies utilized patients having received prior antibiotics (50,51,62). In their large meta-analysis of patients with RTI, initial PCT concentrations predicted mortality at 30 days when measured at ED admission, but not at ICU admission, with AUCs of 0.67 and 0.5, respectively (49). In their study, Yu et al. demonstrated that in patients with a suspected infection in the ED or hospital floor, addition of an initial PCT concentration to qSOFA score ≥ 2 significantly improved prediction of 30-day mortality (57).

In general, most studies comparing the prognostic utility of initial PCT concentrations and PCT clearance showed differences in its ability to predict 28/30-day mortality depending on patient location (i.e., ED versus ICU). This is likely due to differences in disease severity and treatments noted in several studies.
Outcomes differ by patient diagnoses, sepsis definitions, and study populations

Groups have also identified differences in PCT’s prognostic utility across patients’ diagnoses. In septic patients, discrepancies among studies could be due to the sepsis definition utilized. Studies published prior to 2016 likely utilized either the Sepsis-1 or 2 definition to classify patients, whereas the Sepsis-3 definition may have been used in more recent studies (65). In their meta-analysis, Elke and colleagues demonstrated that mortality rates differed among sepsis populations depending on the definition utilized. However, regardless of the sepsis definition (Sepsis-1 vs Sepsis-3), PCT concentrations measured within 24 hours of a severe sepsis or septic shock diagnosis were a poor predictor of 28 day mortality (AUC was 0.56 for both populations) (51). Interestingly, in their meta-analysis of septic patients, while PCT concentrations measured on days 1 and 3 were both significantly different between survivors and non survivors in the total population, in a subgroup analysis of patients with severe sepsis or septic shock, PCT concentrations were not significantly different in patients who died (64).

Among patients with RTI, initial PCT concentrations correlated with 30 day mortality in patients with COPD and CAP, but not in patients with acute bronchitis or VAP (49). Elke et al. demonstrated that PCT, measured at baseline, had marginally improved mortality prediction among patients with pneumological compared to intraabdominal infection (AUC 0.58 versus 0.52). Further, initial PCT concentrations were significantly correlated with 28-day mortality among patients with Gram positive and negative infections, but not among patients with fungal infections(51). Most studies noted differences in the PCT’s ability to predict 28/30-day mortality across different diagnosis, likely due to differences in disease severity, treatments and mortality rates in the individual populations.

One other source of confusion regarding the prognostic value of PCT is that significant heterogeneity exists among data sets in individual studies, making it difficult to draw conclusions. In their meta-analysis, Liu et al. demonstrated significant heterogeneity across data sets in individual
studies, especially their mortality rates, which ranged from 17 – 66.7% (50). In a large meta-analysis of
patients with RTI the mortality rate was only 6%, leading to high negative predictive values for PCT (49).
In another large meta-analysis conducted by Arora and colleagues looking at the utility of PCT to predict
mortality among septic patients, statistical heterogeneity of the patient populations across studies was
high, and mortality rates ranged from 13 – 69% (64). More recently, in their study of patients with
suspected infection, Yu et al. reported a mortality rate of 9% and were also able to demonstrate high
negative predictive values for prediction of 30 day mortality (57). In contrast, in their study of septic
patients, Elke et al. demonstrated that initial PCT concentrations were a poor predictor of 28 day
mortality, and their mortality rate was 27% (51). As expected, heterogeneity in patient populations
among different studies is significantly impacted by mortality rates and thus the performance
characteristics of PCT to predict mortality (50).

Outcomes other than 28/30-day mortality
While most of the above discussion has focused on 28/30-day mortality, studies have
investigated other outcomes as well. For example, several studies addressed PCT and its correlation
with mortality during the patient’s hospitalization (50,51,59,62) and/or at other time points including: 7,
14, or 90 days, 1-year post presentation, or undefined mortality endpoints (44, 46,50,57,59,60).
There has also been significant interest in utilizing PCT to predict response to therapy. In
general, PCT concentrations and/or evolution of PCT concentrations over time are good predictors of
successful or failed treatments in a variety of patient populations (49,68). PCT concentrations at
presentation were also correlated with general disease progression, admission to the ICU and/or length
of hospital stay (59). Zaccone et al. showed that PCT measured within 12 hours of admission among
1063 critically ill patients was an accurate predictor of ICU transfer (69). Similarly, in a large population
of patients with lower acute respiratory infection (ARI), initial PCT concentrations correlated with
treatment failure at 30 days. Further initial PCT concentrations even correlated with treatment failure among patients with certain upper ARIs like the common cold or Rhinosinusitis (49). In contrast, in a population of patients with febrile UTI, PCT concentrations at presentation, day 3 or PCT clearance over time were able to predict treatment failure with AUCs of 0.52, 0.55, and 0.58 respectively (70).

In summary, PCT concentrations increase with disease severity in patients with sepsis and RTIs as well as in other select patient populations. Elevated PCT concentrations measured at ED or hospital admission in patients with sepsis or LRTI are associated with a greater risk for 28–30-day mortality. Similarly, a lack of PCT clearance over time is also associated with a great risk of mortality. Similar trends were observed with other mortality outcomes as well as treatment response and/or disease progression. However, significant heterogeneity in study populations across studies, especially related to mortality rates, limits our ability to formally recommend the use of PCT as a predictor of prognosis.

Other biomarkers evaluated to predict outcomes in patients with sepsis and/or respiratory tract infections

Algorithms which use biomarker results to stratify patients by mortality risk and which provide actionable information for patient management are a promising tool for patient care. PCT is likely the best studied biomarker for this purpose in the context of sepsis and/or respiratory tract infections; however, its performance varies across studies (Suppl. Table 3). Although PCT is FDA-cleared for predicting outcomes and disease progression in patients with sepsis, factors such as optimal cut-offs and recommended frequency of testing vary by clinical context and thus complicate its operationalization (71–75). These variables are further discussed in later sections. Other biomarkers with prognostic roles include standard of care tests such as lactate and CRP, as well as a growing list of candidate biomarkers including Interleukin-6, mid-regional proadrenomedullin (MR-proADM), presepsin and multibiomarker models. Below, we summarize the evidence for the biomarkers currently used clinically and will provide an outlook on promising early findings for the newer biomarkers.
Lactate is a key biomarker routinely used for outcome prediction in patients with sepsis despite the low quality of evidence (76). Although it can be elevated in other contexts, increases in blood lactate indicate tissue hypoxia and its measurement is a surrogate marker of hypoperfusion. Patients with elevated lactate concentrations have poor outcomes while sufficient decreased lactate over time (also referred to as lactate clearance) is associated with decreased mortality. In adults patients admitted to the ICU, the likelihood of mortality decreased by 11% for every 10% increase in lactate clearance (77). In children with septic shock, failure to achieve a lactate clearance of >10% increased the risk of mortality (LR 2.83; 95% CI 1.82-4.41) (78). Surviving Sepsis Campaign guidelines recommend to measure lactate promptly after sepsis is suspected or identified and to re-measure it if elevated >2 mmol/L. (76). A lactate value of ≥4 mmol/L warrants fluid resuscitation and normalization is targeted (76). Randomized Control Trials evaluating lactate clearance for therapy-guidance and outcomes vary in their conclusions. For in-hospital mortality, one RTC found a reduction only when adjusted for risk factors (79) while another did not find an effect (80). Tian et al. reported that 10% and 30% lactate clearance was not associated with a reduction in 7 day mortality rate, but the 28 day mortality was significantly lower in patients with ample lactate clearance(81). A meta-analysis including these studies (547 patients), concluded the use of lactate clearance to guide therapy reduces the risk of mortality (risk ratio of 0.65, 95% CI 0.49 – 0.85) (82). It is important to consider that lactate clearance is only appropriate for use in patients with severe sepsis and/or septic shock as it is not elevated in early and/or mild sepsis.

C-reactive protein (CRP) has been heavily studied as a sepsis biomarker and is frequently utilized to monitor numerous inflammatory disorders. The evidence is controversial regarding its role as an outcomes predictor in patients with sepsis and/or respiratory tract infections. Several studies report that in comparison to PCT, CRP measurements at study enrollment and/or admission are not significantly higher in survivors compared to non-survivors (51,57,59). In two studies comparing
prognostic utility of several biomarkers, including PCT and CRP, in patients presenting to the ED. PCT and CRP were significantly associated with 28 day mortality, but neither was considered an independent predictor of mortality (52,61). In general, CRP lacks significant clinical utility as a prognostic marker.

Other new biomarkers may have promise but have not been adopted into clinical practice. MR-proADM is a product of proADM, generated in a 1:1 ratio with adrenomedulin, a calcitonin peptide family mostly known for its vasodilatory activity. Like PCT, MR-proADM elevations are not specific to infections. Several studies have established a relationship between MR-proADM and outcomes such as disease progression and mortality, and its superiority compared to other biomarkers including PCT, CRP, copeptin and presepsin (71–75). One such study, a prospective multicenter study, reported that MR-proADM outperformed PCT and CRP, and clinical scores such as SOFA/qSOFA and National Early Warning Score (NEWS) for ICU admission and 28-day mortality in ED patients (n=684) at presentation and 3 days after (59). MR-proADM had the strongest association by univariate analysis with requirement for ICU admission (OR 4.1 [2.3 – 7.1] vs. PCT OR 2.2 [1.5-3.4] vs. CRP 2.1 [1.2-3.6]), and 28-day mortality (MR-proADM OR 4.1 [2.6-6.5] vs. PCT OR 1.9 [1.3-2.7] vs. CRP OR 1.0 [0.7-1.5]). Adding MR-proADM and PCT increases the correlation with mortality (HR 5.7 [2.8-11.6]), a combination that could be explored further. The study utilizes a non-commercially available point-of-care analyzer for PCT and MR-proADM. Although the availability of rapid testing could increase access to these biomarkers for decision-making, more data will be needed to demonstrate the required analytical performance characteristics to support such applications. Studies using a laboratory assay available for MR-proADM (83) demonstrated consistent MR-proADM performance. Mearelli, and colleagues showed that MR-proADM improves qSOFA’s outcome prediction ability (84). Of 8 biomarkers evaluated (CRP, lactate, soluble interleukin 2 receptor alpha (sIL2Ralpha), soluble triggering receptor expressed on myeloid cell-1 (sTREM-1), secretory phospholipase A2 group II (sPLA2GIIA), presepsin, and MR-proADM) in a secondary analysis of a study in 5 EDs in Italy, the addition of some clinical parameters, CRP, lactate
and MR-proADM, yielded the highest AUROC of 0.83 (95% CI 0.8-0.87). The study by Elke and colleagues also reported MR-proADM was the strongest mortality predictor at baseline relative to PCT and CRP in patients with sepsis (AUC 0.73 vs 0.56 for PCT and 0.55 for CRP) and septic shock (AUC 0.72 vs 0.50 for PCT and 0.53 for CRP); MR-proADM remained the strongest predictor at days 1, 4, 7, and 10. In summary, there is some data to suggest a potential role for MR-proADM in outcomes prediction in septic patients, however, currently, MR-proADM assays are not widely available or cleared for this use.

Presepsin, a subtype of soluble cluster-of-differentiation marker protein 14 (CD14), is released into the blood after lipopolysaccharides in microorganisms bind CD14 in monocytes and macrophages. Its prognostic role has been extensively described but mainly in observational studies with small sample sizes. Presepsin is significantly higher in non-survivors in ED and ICU settings and with a weighted pooled standardized mean difference of 1.09 (95% CI 0.78-1.41) for 30-day mortality. When compared to PCT, there is no conclusive evidence pointing to the superiority of presepsin for mortality prediction. A meta-analysis of 9 studies and approximately 1,500 patients in ED and ICU settings concluded that presepsin is not superior to PCT for mortality prediction (85). The AUC of PCT was 0.81 (95% CI, 0.78–0.84) with a pooled sensitivity of 0.76 (95% CI, 0.55–0.89) and specificity of 0.74 (95% CI, 0.33–0.94) and the AUC of presepsin was of 0.77 (95% CI, 0.73–0.81) with pooled sensitivity and specificity of 0.83 (95% CI, 0.72–0.90) and 0.69 (95% CI, 0.63–0.74), respectively. These studies utilized a unified POC assay (PATHFAST, LSI Medience Corp, The Netherlands), yet cut-offs and clearance strategy are not standardized.

As discussed above with only a few examples, new biomarkers, alone or in combination, show early promising results for roles in predicting outcome. However, the evidence to support the utility of these biomarkers for outcome prediction strengthened is an active area of research. Of note, these biomarkers have almost exclusively been studied in developed counties and in relatively small sample sizes.
cohorts. Although some have been published as mentioned above, future studies on these candidate biomarkers in randomized controlled studies and across different medical centers and with larger sample size will be needed to demonstrate if these promising early results can be validated. Moreover, these studies should focus on establishing evidence-based cut-offs and interpretative criteria is necessary to draw meaningful conclusions for real-live applications. Until then, PCT is the only assay in the US cleared by the FDA for 28-day mortality prediction in critically ill-patients. Despite the FDA approval, the lack of uniformity in the studies make recommendation of a specific clearance cut off challenging.

**Key summary points**

- PCT concentrations increase with disease severity in patients with sepsis and respiratory tract infections.
- Elevated PCT concentrations measured at ED presentation or hospital admission in patients with sepsis or LRTI are associated with a greater risk of 28–30-day mortality.
- In patients with sepsis and/or RTI, the lack of PCT clearance overtime is associated with a greater risk of 28 – 30-day mortality.
- The heterogeneity of patient populations in the studies makes is difficult to form uniform recommendations for the use of PCT as an outcome predictor.

Can PCT results be utilized to inform treatment decisions in both initiation and cessation of antimicrobial therapy in neonatal and pediatric patients with sepsis or respiratory infections?
There are numerous studies evaluating the role of PCT in critically ill children in the pediatric ICU (PICU). Many of these studies have investigated the use of PCT to help differentiate serious bacterial infections, distinguish bacterial from viral meningitis (86–93). Since the use of PCT in this manner informs decisions on treatment and initiation of antibiotics, there has been much interest in using PCT to identify or rule out sepsis and serious bacterial infections. Unlike the adult studies, the pediatric studies did not directly evaluate the rate of antibiotic initiation and instead focused on evaluating the sensitivity, specificity and negative predictive value of PCT in detecting bacterial infections.

PCT for detecting bacterial infections and initiating antibiotics in pediatric and neonatal patients

A meta-analysis of 12 studies involving over 7000 children (<18 years) reported a sensitivity and specificity of 55% and 85%, respectively, for detecting serious bacterial infections (87). For invasive bacterial infections, sensitivity and specificity was 82% and 86%, respectively, using a PCT cutoff of 0.5 ng/mL. In this analysis, serious bacterial infections included a broad spectrum of infections ranging from bacterial meningitis to urinary tract infections as well as a subgroup of severe invasive bacterial infections including bacterial meningitis, sepsis and bacteremia. The negative predictive value of PCT was approximately 99% for invasive bacterial infections and ranged from 79.5% to 96.7% for serious bacterial infections.

A meta-analysis of 28 studies that included over 2600 neonates with suspected sepsis PCT showed a sensitivity of 85% for detecting sepsis and when combined with CRP, the sensitivity improved to 91% (88). In a prospective trial of 80 children, PCT was better able to detect severe infections among PICU patients when compared to CRP or WBC (89). Another prospective study of 64 PICU patients showed that PCT outperforms CRP alone in detecting bacterial infections; however, PCT was insufficiently sensitive with an AUC of 0.71 (90). A study of 85 PICU patients with suspected sepsis demonstrated a
negative predictive value of 90% when a combination of CRP (<4 mg/dL) and PCT (<1.75 ng/mL) is used to identify critically ill children with a low risk of bacterial infection (91). A retrospective study of over 600 PICU patients, reported a negative likelihood ratio of 0.3 for PCT in ruling out infection (92). There is an increasing amount of evidence that suggests PCT in combination with other biomarkers may be useful in ruling out infection or identifying infants with low risk for serious infections. Kuppermann and colleagues developed a clinical prediction rule [Pediatric Emergency Care Applied Research Network (PECARN)] for identifying <60 day old infants across 26 emergency departments that are at low risk for serious bacterial infections to avoid unnecessary lumbar punctures, antibiotic exposure, and hospitalizations. This algorithm incorporated urinalysis, absolute neutrophil count, and PCT and demonstrated a sensitivity of 97.7% and specificity of 60.0%, with a negative predictive value of 99.6% (93). However, further validation of such multi-biomarker algorithms is necessary. An external validation of the PECARN rule yielded a sensitivity and specificity of 89.8% and 55.5% for serious bacterial infections, with an AUC of 0.726 when tested in a cohort of 1247 infants presenting to a pediatric emergency department in Spain (94). The authors caution the use of this prediction rule in young infants with a short history of fever.

Similar observations regarding the ability of PCT to rule out bacterial infections have been made in children <18 years old with LRTI. Using a PCT cutoff of <0.25 ng/mL a 96% NPV, 85% sensitivity and 45% specificity was observed for ruling out typical bacterial CAP in a study of 532 hospitalized children with radiographically confirmed CAP (95). Another group had evaluated the utility of PCT in guiding antibiotic treatment in 319 children with pneumonia. A PCT guided algorithm using a threshold <0.25 ng/mL for withholding antibiotics in the study group (n=155) resulted in 85.8% children receiving fewer antibiotic prescriptions and a shorter exposure time (3.9% vs 25.2%) compared to the control group without a significant difference in recurrence of respiratory symptoms or new antibiotic prescriptions in the following month (96). However, another study of children (1 month-18 years old) presenting with
LRTI found that using PCT-guided algorithms did not alter antibiotic prescription rates but did reduce duration of antibiotic exposure by ~3 days. In this study a PCT >0.5 ng/mL was used to initiate antibiotics and a PCT 0.25-5 ng/mL was left to clinical discretion (97).

Collectively these studies suggest that PCT has insufficient sensitivity in detecting sepsis, serious bacterial infections or LRTI and should not be used as a stand-alone marker to make decisions on initiation of antibiotic therapy. Measuring PCT alone may miss patients with localized but serious infections (98). It is important to note that PCT normal ranges can vary in healthy neonates within the first days of life (peaking at 1 day of life and decreasing to normal by day 2 and 3) (Table 1). Additionally, PCT is not well studied in immunocompromised critically ill children and it is unknown whether these findings can be extrapolated to this population.

Antibiotic cessation in neonatal and pediatric populations

While there is no strong evidence to support using PCT concentrations to guide antibiotic initiation in pediatric and neonatal populations, there is some evidence that suggests PCT is useful to safely discontinue antibiotics this population. Neonatal and pediatric PCT trials are summarized in Suppl. Table 4. The largest multi-center RCT in neonates that assessed the impact of PCT-guided decision making on duration of antibiotic therapy was the NeoPIns trial (99), which enrolled 1710 neonates of >34 weeks gestational age suspected of early-onset sepsis in the first 72 hours of life. 866 neonates were assigned to the experimental arm and 844 to the control arm. In the experimental arm, PCT was measured at 12, 24, 36-72 and every 24-48 hours thereafter until discontinuation of antibiotics. If two consecutive PCT values fell within the normal range for the age, antibiotics were discontinued. PCT guided decision-making led to a 10-hour reduction in antibiotics exposure. However, the impact on re-infection and death during the first month of life could not be entirely assessed due to low occurrence rates of these adverse events. The cut-offs for PCT in this study ranged from 0.5-10 ng/mL, stratified
based on hours after birth (Table 1). It is worth noting that the PCT discontinuation or continuation recommendation per protocol was overruled in 25% of neonates in the experimental arm. A 22.4-hour reduction in antibiotics exposure was observed in an earlier single center study with 121 neonates (61 in control arm and 60 in experimental arm) (100). The study criteria and design were the same as the NeoPlIns trial; however, a single <10 ng/mL PCT cut-off was used in this study. Additionally, the sample size was not sufficient to assess safety measures. One of the limitations of the NeoPlIns trial is that this study included only neonates >34 weeks of gestational age, and the utility of PCT in pre-term infants is unknown.

A similar reduction of antibiotic exposure has been reported in children <18 years old. A single-center study from Spain examined the impact of implementation of a PCT-guided protocol in antibiotic decision-making (101). 114 patients were examined prior to implementation of the protocol and 112 after implementation. Antibiotics were discontinued if there was a 50% decrease in PCT value or PCT values dropped below <0.5 ng/mL. Implementation of this PCT-guided protocol resulted in a reduction of 1.1 days of antibiotic exposure without adverse outcomes. The compliance of antibiotic de-escalation in the PCT protocol was only 54.8%; however, prior to PCT implementation, de-escalation only occurred in 26% of the patients. Using a PCT cutoff of <0.25 ng/mL in the pediatric LRTI population has also demonstrated a reduction in antibiotic exposure (96,97).

The studies evaluating the role of PCT for antibiotic cessation are limited in the pediatric populations and further investigation is warranted. Overall, the studies to date suggest that there is a role for PCT in guiding discontinuation of antibiotics in both neonatal and pediatric populations. Though the studies did not report any adverse outcomes, there was a low rate of re-infection or death occurrence in the control arms for the impact to be assessed.

Key summary points
- PCT should not be used as a stand-alone test for the diagnosis of sepsis or to guide antibiotic initiation in pediatric patients.
- PCT can guide safe cessation of antibiotics in neonates and pediatric patients with suspected sepsis who show clinical improvement.
- In neonates, PCT concentrations rise and fall rapidly, thus cut-offs need to be stratified by age (hours after birth).
- There is no consensus for PCT cutoffs or clearance rates to guide duration of antibiotic therapy in pediatric patients.

**Is PCT an accurate predictor of outcomes (e.g., mortality, respiratory failure, shock) in pediatric populations?**

A few studies have examined the role of PCT as a prognostic predictor in pediatric populations. As with adult populations, higher PCT concentrations are associated with sepsis severity and increased risk of death (102–105). Conversely, low or normal PCT values have excellent negative predictive values for adverse outcomes. In an observational prospective study of 65 children with meningococcal infections, patients with PCT concentrations <10 ng/ml survived; whereas all patients with PCT ≥ 10 ng/mL developed multiple organ dysfunction syndrome or died (106). Another study looking at PCT kinetics in pediatric patients with systemic inflammatory response syndrome and organ failure after open heart surgery showed similar outcomes, where PCT concentrations <10 ng/mL post-surgery survived (107).

**Outcomes from single PCT measurements**

Single PCT measurements at admission cannot predict the likelihood of a patient developing severe sepsis or septic shock. In a retrospective single center study evaluating 109 critically ill children who had
a PCT measurement within 48 hours of admission, 61 patients with septic shock had a median PCT concentration of 7.16 ng/mL with an interquartile range of 2.21-42.28 ng/mL; whereas, another 48 patients without septic shock had a median PCT concentration of 0.91 ng/mL and a interquartile range of 0.10-10.80 ng/mL(92). Though PCT concentrations were higher in patients that progressed to severe sepsis or septic shock, there was significant overlap in PCT concentrations between the two groups. Therefore, PCT measurements alone are unable to predict probability of developing severe sepsis or septic shock. Similarly, in another study of 64 patients with meningococcal sepsis and septic shock, median PCT levels on admission were higher in children with septic shock compared to children with sepsis (270 ng/mL and 64.4 ng/mL, respectively). However, in this study there was no significant difference in PCT ranges for survivors (n=42, 5.7-672.3ng/mL) and non-survivors (n=13, 55-646.4) with septic shock (108). Similarly, in a study of 75 children with septic shock though non-survivors had higher PCT concentrations (median 273 ng/mL for non-survivors vs. 82 ng/mL for survivors), the range of PCT concentrations among survivors and non-survivors was similar (3.3-759.8 ng/mL and 5.1-736.4 ng/mL, respectively). The maximum length of ICU stay in this study was 32 days (102).

Outcomes from sequential PCT measurements

Hatherill and colleagues also evaluated changes in PCT concentrations after treatment. They found that of 39 children with sequential PCT measurements, 16 (41%) showed no fall in PCT after 24 hours of treatment and the observed mortality in this group was 44% compared to 9% in 23 patients that showed a decline (102). The RESOLVE phase III trial examined the biomarker response in children with severe sepsis after treatment with either placebo or Drotrecogin alfa. In this study, 251 survivors showed a decline in PCT concentrations compared to the 37 non-survivors who showed an increase in PCT concentrations 24 hours post-treatment regardless of whether patients were in the trial or control arm (109). Overall, these studies demonstrated that a decline in PCT values after treatment is associated with better survival rates; however, the percent change in PCT concentrations were not provided in
these studies. Poddar et al. evaluated whether reduction in PCT can predict 28-day mortality in 20266 children admitted to the PICU with severe sepsis or septic shock. Of the 14 children that survived to 28268 days, the percent reduction in PCT was 75.5% compared to a 200% increase in PCT concentrations in the269 6 non-survivors between day of admission and 72-96 hours later(110).

Given the limited number of patients evaluated in these studies and low rate of mortality, larger275 studies are needed to determine extent of PCT reduction required to accurately predict mortality in276 pediatric patients. Additionally, outcomes assessment was only a secondary measure assessed in most277 of these studies and therefore details of PCT kinetics and percent reduction were largely missing.

Key summary points

- An elevated PCT is generally suggestive of a worse outcome in pediatric patients with severe283 sepsis or septic shock.
- A single PCT measurement has limited prognostic value since many studies have shown285 significant overlap in PCT concentrations among survivors and non-survivors.
- Serial PCT measurements may be predictive of mortality during ICU stay; however, additional287 studies are needed to define interpretive criteria in pediatric and neonatal patients.

When and how often should PCT be measured? Which cut-off(s) should be used?

PCT guided algorithms have been investigated to optimize antimicrobial therapy and predict outcomes299 including mortality, disease progression, and length of stay. This strategy has safely reduced antibiotic300 treatment in septic patients in different clinical settings (i.e., ED, ICU) and various etiologies, particularly301 respiratory infections (Suppl. Table 1 and 2). However, recent trials did not confirm these findings (15–303 18,36). Algorithms evaluated across studies differ in positive cut-off(s), timing of serial testing, and PCT304
assays used. Commonly, PCT algorithms consist of recommendations to initiate or discontinue antibiotics, typically using different cut-offs based on acuity, clinical setting (i.e., ED, ICU), and patient population.

### Timing and frequency for antibiotic initiation and cessation in adults

The recommended PCT cut-off tiers for patients in the ED and hospital wards are <0.1, 0.1-0.25, 0.26-0.5 and >0.5 ng/mL ([Suppl. Tables 1 and 2](#)). In patients with uncertain clinical suspicion of infection, PCT concentrations <0.1 and 0.1-0.25 ng/mL indicate that initiation of antibiotics is strongly discouraged and discouraged, respectively. However, if there is no clinical improvement, the studies referenced in Suppl. Tables 1 and 2 support that PCT should be rechecked after 6-24 hrs. PCT concentrations 0.26-0.5 and >0.5 ng/mL indicate that antibiotics are encouraged and strongly encouraged, respectively, and to re-measure PCT every 2-3 days to assess for the opportunity of discontinuation of antibiotics. Upon re-evaluation, PCT concentrations <0.1 and 0.1-0.25 ng/mL strongly encourages and encourages discontinuation of antibiotics, respectively. When PCT drops by >80% from its peak value, antibiotic discontinuation is also recommended. PCT concentrations 0.26-0.5 and >0.5 ng/mL discourage and strongly discourage discontinuing antibiotics, respectively. A simplified approach for patients with moderate illness outside of the ICU consists of empiric antibiotic initiation based on clinical practice guidelines and initial assessment or if PCT is 0.25 ng/mL, measuring PCT at least daily and stopping antimicrobial therapy if PCT decreases to <0.25 ng/mL or by at least 80% from peak concentration combined with improvement of clinical symptoms (111). Not surprisingly, different RCTs attempting to assess the feasibility of using PCT for initiating or discontinuing antibiotics across different patient populations and settings have utilized different algorithms and cut-offs, as discussed above. Less is known about the broader applicability of these algorithms for predicting other outcomes such as mortality.
Timing and frequency for outcomes prediction in adults

In the US, most PCT assays were initially FDA cleared to predict disease progression. More recently, some intended uses have expanded to mortality risk assessment and antibiotic management decision making. A higher 28-day risk of all-cause mortality is predicted in patients with PCT concentrations that increase or decline ≤ 80% from the day severe sepsis or septic shock were first diagnosed (Day 0) or Day 1 to Day 4 (62). In this context, the main overlap with the cut-offs from the antibiotic initiation/cessation algorithms discussed above is the assessment of PCT clearance in patients with high acuity disease. Not all studies that demonstrated an association between PCT concentrations at presentation and 28/30-day mortality reported a concentration cut-off, and those that did used a variety of cut-offs, complicating our ability to recommend an evidence-based cut-off for mortality prediction. Moreover, reported cut-offs differ across patient populations and diseases.

Some studies used a cut-off of 0.25 ng/mL (59,111), while others used cut-offs such as those described above for antimicrobial stewardship (i.e. 0.1, 0.25, 0.5, 2.0 ng/mL)(49,55), and others reported a variety of cut-offs (50,53,54) (Suppl. Table 3). The study by Kutz and colleagues, a meta-analysis of 14 trials, found an association between increasing PCT concentrations at presentation and adverse outcomes such as treatment failure and mortality but only in ED patients and in patients with ARI and CAP, but not ICU or primary care settings. A cut-off of 0.1 ng/mL resulted in sensitivities of 86.1 (95% CI 82.4-89.3) and 92.5% (95% CI 86.2-96.5) in ED patients for treatment failure and mortality, respectively, and >90% for mortality in patients in the ICU, or with ARI or CAP. For these same parameters, specificity approaches 80% at a PCT cut-off of 2.0 ng/mL. Similarly, a multi-national prospective study evaluating several PCT cut-offs (0.05, 0.1, 0.25, 0.5 ng/mL) in samples collected at ED admission from nearly 7,000 patients showed association between 30-day mortality and increasing PCT concentrations (55). The study reported OR of 7.31 (95% CI 3.32-14.75) for patients with an admission PCT >0.5 ng/mL. In comparison, samples with PCT<0.1 ng/mL or <0.05 ng/mL had OR of 1.71 (95% CI 0.87-3.34) and of 1.0,
respectively. A large multicenter prospective study of over 1,700 patients with septic shock and PCT measured in the ED at diagnosis derived an optimal PCT cut-off of 0.17 ng/mL but found that PCT was not an independent predictor of 28-day mortality (61). In their study of severely ill trauma patients, those with PCT of 5 ng/mL or greater were at higher risk of dying (OR 3.65, 95% CI 1.03-12.9) (54). A study in patients with tuberculosis reported that baseline PCT concentrations >0.13 ng/mL predicted mortality with OR of 7.9 (95% CI 3.2-19.7); however, reported cut-offs for mortality ranged from 0.05-0.12 on days 7, 14 and 28, highlighting the need for a simplified approach to integrating PCT into mortality prediction, such as an integrated risk-score.

**Change in PCT concentrations over time to predict outcomes in adults**

The MOSES study found that while baseline PCT was a poor predictor of 28-day mortality (AUC 0.56; 95% CI 0.51-0.60), failed PCT clearance (≤80%) between baseline and day 4 doubled the mortality (62). On Day 1, mortality was 3-fold lower in patients with decreased PCT (29 versus 12%), regardless of the initial PCT concentration. In a study in patients with intra-abdominal sepsis, 5-day 70% PCT clearance predicted mortality while PCT clearance at days 3 and 4 did not differentiate survivors and non-survivors (63). Lower PCT clearance cut-offs have also been reported to effectively predict mortality. Elke et al. demonstrated that patients with PCT clearance by 20% at day 1 or by 50% at day 4 had lower mortalities in the ICU and in-hospital (16.8 and 24.1 vs 28.9 and 30.4%, respectively) (51). In patients with autoimmune disease, PCT peak concentrations did not differ between survivors and non-survivors while PCT clearance on days 5 and 7 was significant (p=0.06 and 0.005, respectively) (60). In this study, clearance was calculated from the PCT peak concentration, an approach difficult to operationalize as it is challenging to predict when the peak PCT will occur prospectively.

The value of both PCT concentrations and clearance for 28/30-day, ICU or in-hospital mortality prediction was evaluated in a meta-analysis, which included 23 studies (up to 2014) with 3,994 patients.
The studies associating absolute values of PCT with mortality included different clinical settings (ED and various ICUs) and PCT cut-offs (0.12-14.27 ng/mL). For the evaluation of PCT clearance, the studies were mostly ICU (1 SICU) and the clearance cut-offs ranged from 25%-70%. In both cases, sample size and mortality varied significantly. The AUCs were 0.77 (95% CI 0.73-0.80) and 0.79 (95% CI 0.75-0.83) for a single PCT concentration vs. PCT clearance, respectively. Non-clearance is associated with a RR for mortality of 3.05 (95% CI 2.35-3.95). The meta-analysis provides helpful insight into the value of PCT for mortality prediction but does not help elucidate the optimal cut-off to use or the definition of clearance and time points used.

Unfortunately, the degree of heterogeneity in these studies is significant (64). With such heterogeneity and inconsistency across studies, it is difficult to make a uniform recommendation regarding the utility of PCT in predicting prognosis. Further, even if a consistent cut-off for the PCT delta calculation were identified to predict 28-day mortality, it is unclear as to whether these predictions have any impact on patient care. Additional studies are needed to establish standardized cut-offs and/or PCT clearance parameters in the prediction of 28-day mortality and to determine the clinical utility of a 28-day mortality prediction in patients with sepsis and LRTI patients.

Timing and frequency for antibiotics initiation and cessation in pediatric patients

The majority of evidence described earlier in the pediatric populations demonstrates that PCT has some utility in identifying patients with sepsis and severe bacterial infections; however, PCT does not have adequate sensitivity to serve as a stand-alone test to guide decisions on antibiotic therapy initiation. A meta-analysis consisting of over 7000 children showed that a PCT cutoff of 0.5ng/mL had a sensitivity of only 55% (specificity 85%) for detecting serious bacterial infections and the negative predictive value ranged from 79.5%-96.7 % (87). A prospective study of 64 PICU patients showed that a PCT cut-off of 2.5 ng/mL at admission had optimal sensitivity and specificity (68% and 74%, respectively),
with a negative predictive value of 78%; whereas a CRP cut-off of 40 mg/L had a sensitivity of 95% and specificity of 42% with a negative predictive value of 94% (90). Another study of 85 PICU patients demonstrated a negative predictive value of 90% for ruling out bacterial infection in patients with systemic inflammatory response syndrome when PCT and CRP were used together, with a CRP cut-off of <4 mg/DL and PCT cut-off of <1.75 ng/mL measured <4 hours after initiation or expansion of antibiotics (91). A retrospective analysis of 600 PICU patients reported a negative likelihood ratio of 0.3 for PCT using a cut-off of <0.1 ng/mL. When used in combination with a CRP < 0.8 mg/dL the negative likelihood ratio was 0.1 for bacterial infection (98).

A meta-analysis of 2600 neonates showed that using PCT in combination had a sensitivity of 91% for detecting neonatal sepsis compared to using CRP and PCT alone, with a sensitivity of 71% and 85%, respectively. The cutoff intervals for PCT proposed by sub-analysis was 0.5-2ng/mL and a cut-off value of >10 mg/L for CRP yielded the highest sensitivity and specificity (88). Due to the rapid changes in PCT concentrations in neonates, caution should be used for assigning an absolute PCT cut-off in neonates <72 hrs of age. As in the pediatric population, the cut-off for PCT used to detect bacterial infections varies by study in neonates and will be dependent on the algorithmic approach. For example, in the prediction rule described by Kuppermann and colleagues to rule out serious bacterial infections in infants, a PCT concentration of ≤1.71 ng/mL was used in conjunction with a negative urinalysis result and absolute neutrophil count of ≤4090/μL (93). Collectively, these studies suggest that PCT in combination with CRP has superior performance in ruling out bacterial infection compared to PCT measurements alone in both pediatric and neonatal populations. However, the timing of PCT measurement and proposed cut-offs vary greatly study to study. Overall, the major limitations for comparing studies examining the value of PCT in detecting sepsis or bacterial infections are a lack of a uniform definition for sepsis, varied inclusion and exclusion criteria, and lack of harmonized criteria for time of sampling and interpretation.
The NeoPIns trial provides the strongest evidence and guidance for cessation of antibiotics in neonates >34 weeks of gestational age and with onset of sepsis within the first 72 hours of life. In the trial when PCT concentrations fell within the normal range for the age (Table 1), antibiotics were discontinued. PCT was measured at 12, 24, 36-72 hrs and every 24-48 hours until discontinuation of antibiotics (99). The NeoPIns trial demonstrated a 22.4-hour reduction in antibiotic exposure. In children <18 years old, one single center study evaluated a PCT reduction criteria of 50% or PCT decrease below 0.5 ng/mL. PCT was measured at baseline, 24, 48 and 72 hrs of antibiotic treatment. Using this criteria for antibiotic cessation, a 1.1 day reduction in antibiotic exposure was observed without adverse outcomes (101). While more studies are needed in pediatric and neonatal populations, the evidence for PCT use in antibiotic cessation is encouraging.

Guidelines from the American Academy of Pediatrics on assessment of febrile well-appearing infants 8 to 60 days old recommend using PCT >0.5 ng/mL for initiating antibiotic treatment citing evidence that PCT is an independent predictor of bacterial infections with better performance characteristics than CRP, ANC and WBC (112).

Timing and frequency for outcomes prediction in pediatric patients

The prognostic ability of single increased PCT measurement is poor in predicting mortality, as the studies mentioned above showed that even though the median PCT concentrations were higher in non-survivors compared to survivors, the ranges of PCT concentrations in both groups overlapped significantly (67). Thus, a PCT cut-off concentration for mortality predication cannot be recommended. As evidenced by the MOSES trial in the adult population, serial PCT measurements are likely more useful than single PCT measurements in predicting the risk of death in pediatric populations. However, large, prospective multicenter studies examining the prognostic accuracy of PCT reduction are lacking for the pediatric population. Additionally, small sample sizes, low death rates and differences in study design hinder the ability to make any conclusions on timing, frequency, cut-offs and reduction for PCT.
measurements needed to make accurate predictions of mortality (102,109). Only one single center study evaluated the ability of serial PCT measurements 4 days apart to predict 28-day morality. The authors found that a 75.5% reduction in PCT concentrations was seen in survivors; however the study size was small consisting of only 20 children (14 survivors and 6 non-survivors) (110). It is also important to note that PCT is usually not the only factor in predicting mortality in septic ICU patients as other clinical information (cause of illness, other underlying conditions and other test results) are considered.

Key summary points

- Decision thresholds for antimicrobial use/discontinuation vary by diagnosis and acuity and should considered in the context of other clinical signs and symptoms.
- No consistent PCT cut-off concentration has been established to predict mortality.
- Insufficient PCT clearance over time is associated with a significantly greater risk of 28–30-day mortality in septic and LRTI patients.
- No consistent PCT clearance calculation parameters have been established to predict 28–30-day mortality.
- Routine measurement of PCT to predict mortality is not recommended due to lack of consistent PCT cut-offs and/or PCT clearance parameters and insufficient evidence demonstrating a benefit to estimating 28-day mortality risk in septic and LRTI patients.

How should PCT be incorporated into antimicrobial stewardship efforts?

As adherence to a pre-defined PCT algorithm has been reported to vary in real-world PCT studies, active antimicrobial stewardship intervention paired with PCT results may be needed. Other rapid diagnostic technology, primarily in microbial identification from cultures, has shown mortality benefit only when an antimicrobial stewardship intervention was paired with the result (113). Effective
antimicrobial stewardship requires a multidisciplinary team effort involving infectious disease physicians, nurses, pharmacists, and laboratorians (114). The CDC’s 2019 “Core Elements of Hospital Antimicrobial Stewardship Programs” guidelines indicate that the use of procalcitonin might help identify patients in whom antibiotics can be stopped because bacterial pneumonia is unlikely (115). These guidelines indicate that laboratory and stewardship personnel can work collaboratively to present lab data in a way that supports optimal antibiotic use and is consistent with the institution’s expected practices. Further, the laboratory can collaborate with stewardship program personnel to develop guidance for clinicians around clinical decision making based on laboratory results.

We reviewed the literature for studies that assessed the effectiveness of incorporating PCT into antimicrobial stewardship efforts. Many of the reviewed studies describe providing extensive education and feedback to the prescribers and the development of a clinical decision pathway or algorithm to provide guidance prior to implementation of PCT at their institution (15,27,33,116). Examples of how education was provided include didactic educational sessions, case reviews, and creation of pocket cards (116). Notification to the prescriber of the PCT result through text page or e-mail with guidance on interpretation of the result or prospective audit and direct feedback to the prescriber may provide benefit (36). Although the specific cut-off of PCT values to recommend antibiotic therapy varied among the studies published, tailoring and developing an institution-specific algorithm is important based on the patient population, type of infections, or level of acuity seen at the institution, as well as the analytical assay used. In the real-world clinical setting, over-ruling of an algorithm was commonly seen in critically ill patients, thus the development of different algorithms based on PCT interpretation in mild, moderate, or severe presentation with differing antibiotic recommendations-based on severity may be warranted (111). As with any other antimicrobial stewardship intervention implemented at an institution, continued prospective audit with provider feedback on appropriate use and interpretation of PCT will likely be needed (117).
Moradi et al. describes the implementation of a clinical decision support tool to alert a
prescriber in the electronic medical record in a pre- vs post-intervention quasi-experimental study (118).

If the patient met three criteria: 1) positive viral PCR result by FilmArray® Respiratory Panel, 2) a PCT
result of <0.25 ng/mL, and 3) had one or more antibiotic ordered, an alert would populate when a
prescriber opened the patient’s chart indicating “the results suggest a viral infection, please reassess
necessity of antibiotics as indicated.” Without real-time intervention made by an infectious diseases
physician or pharmacist, this intervention provided a significant decrease in inpatient antibiotic days of
therapy (post-intervention mean 5.8 days vs 8 days, p<0.001) and rate of discharge antibiotic
prescription (post-intervention 20% vs 47.8%, p<0.001) and duration of outpatient antibiotic therapy
(0.9 days vs 2.4 days, p<0.001). Antimicrobial stewardship programs should similarly implement
strategic use of clinical decision support tools in conjunction with rapid diagnostic tools to optimize the
use of PCT in antibiotic decision making.

**Key summary points**

- Antimicrobial stewardship programs should implement strategic use of clinical decision
  support tools in conjunction with rapid diagnostic tools to optimize the use of PCT in
  antibiotic decision making.

**What pre-analytical factors affect PCT results and/or interpretation?**

**Acceptable sample types**

The predominant sample used for the analysis of PCT is serum or plasma obtained from the collection of
venous whole blood. Most data from regulatory filings of PCT assays suggest no difference between
serum or plasma (EDTA, lithium heparin, sodium heparin) when using the acceptance criteria: slope =
1.0 ± 0.1, $r^2 \geq 0.95$ (119–123). Arterial samples have also been reported to yield comparable PCT values
to venous samples (124). The validity of capillary blood samples is an important consideration for potential point-of-care testing (POCT) applications of PCT, and one study showed that capillary and venous samples were comparable when analyzed on the same platform (slope = 1.01 [95% confidence interval (CI) 1.00 – 1.05], intercept = 0 [95% CI: 0 – 0], $r^2 = 0.98$) (125). However, direct comparison between capillary blood on the investigated device and venous blood on reference methods showed greater bias, as discussed further below.

### Stability and storage

The only known enzymes to process PCT are found intracellularly, thus there is little known contribution of extracellular proteases to PCT degradation in vitro. Published data have shown no significant change in PCT concentration during room temperature storage for up to 24 hours (124,126). PCT stability can be further prolonged by storage at lower temperatures, such as up to five days at 4 °C and more than 2 weeks at ≤-20 °C (124,126,127). Limited studies have demonstrated that up to three freeze-thaw cycles do not significantly impact PCT levels (124,126).

#### Key summary points

- PCT shows excellent agreement between arterial and venous samples in common vacutainer tubes.
- Limited data suggest that capillary blood could be an acceptable PCT sample type, though more studies are required.

### What FDA-cleared methods are available to measure PCT, and how do they compare?

Numerous FDA-cleared immunoassays are available to measure PCT (Table 2). A detailed analytical overview of many of these assays has been provided by Schuetz and colleagues as well as the
International Federation of Clinical Chemistry and Laboratory Medicine (IFCC) Working Group on Standardization of Procalcitonin Assays (128,129). While many PCT assays were initially approved only for indications of risk assessment (e.g., for progression to severe sepsis and septic shock) and/or risk of 28-day all-cause mortality, most assays are now also approved for antibiotic therapy decision making in patients with suspected or confirmed sepsis or LRTI (CAP, acute bronchitis, and AECOPD). We reviewed the literature to compile reported statistics from method correlation experiments performed between the various available PCT assays (Suppl. Table 5). For the purposes of this review, we considered the BRAHMS PCT sensitive Kryptor to be the reference method, as it was an early-available method that was used in most of the initial clinical trials that first established PCT clinical decision points (37,38). Our subsequent review of method correlation studies is divided into assays that use BRAHMS-licensed PCT reagents, Diazyme-licensed PCT reagents, and other PCT reagents.

**BRAHMS PCT assays**

Ten published studies that compared various PCT assays to the BRAHMS PCT sensitive Kryptor were included in our review (125,126,130–137). Nine of these studies included a test method(s) that utilized BRAHMS-licensed reagents, and these studies were performed across several immunoassay analyzers. For the BRAHMS assays, as compared to the Kryptor reference method, reported slopes ranged from 0.795 to 1.40, and correlation coefficients \( r \) ranged from 0.8864 to 0.997 (Suppl. Table 5). The Abbott ARCHITECT method repeatedly showed slight to moderate negative proportional biases across multiple studies (slopes ranged from 0.806 to 0.97), while the bioMérieux VIDAS and MINI VIDAS consistently showed positive proportional biases (slopes ranged from 1.188 to 1.40). Correlation of the Roche Elecsys cobas methods to the Kryptor varied by the analyzer model; the lower-throughput e411 showed a negative proportional bias (slope = 0.795) while the higher-throughput e600 series and e801 exhibited slopes closer to 1. Intercepts that may be indicative of clinically-significant constant biases included the
ADVIA Centaur (intercept = 0.40) and the cobas e 601 (intercept = -0.47). Other cobas analyzers did not
demonstrate significant intercepts.

Several of the studies further characterized method correlation by quantifying the percent
agreement of categorical interpretation of PCT results relative to the Kryptor at commonly used clinical
decision points. Dipalo et al. found optimal agreement ranging from 94-98% for the Centaur, cobas
e601, LIASON, and VIDAS at the following concentrations: 0.5, 2.0, 10.0 ng/mL (137). Similarly, Lippi et
al. found agreement of 96-99% for the BRAHMS methods tested at 0.10, 0.25, 0.5, 2.0, and 10.0 ng/mL
(131). Conversely, Gruzdys et al. found that an overall negative bias of the ARCHITECT method
translated to predicted medical decision concentrations for the ARCHITECT that were significantly lower
than their Kryptor counterparts, particularly at 0.50 and 2.00 ng/mL (126). However, not all ARCHITECT
studies found reduced agreement at clinical decision points. The varied results of agreement studies are
likely due, in part, to the small number of data points included within each concentration range.

A POCT method, BRAHMS PCT direct, showed reasonable correlation to the Kryptor lab-based method,
more so in venous whole blood (slope = 0.98) than in capillary (slope = 0.90) samples (125). It should be
noted that this study used a combination of both the BRAHMS PCT sensitive Kryptor and Elecsys
BRAHMS PCT assays as the reference method and did not distinguish between them for this analysis.

Diazyme PCT assays

Four published studies were included that compared Diazyme PCT reagents on various immunoassay
analyzers to the Kryptor method (130,131,134,137). Reported slopes ranged from 0.6543 to 1.19, and
correlation coefficients (r) ranged from 0.85 to 0.960 (Suppl. Table 5). Interestingly, the two most
extreme slopes were generated from studies on the same analyzer (Roche cobas c 702), though the
negative bias was generated from serum studies and the positive bias was generated from lithium-
heparin plasma studies, each from separate investigators. Overall, correlation coefficients were lower
with the Diazyme reagent methods than with the BRAHMS reagent methods. Analysis of categorial
characterization often showed lower agreement at clinical decision points for the Diazyme assays than
for the BRAHMS assays. For example, Dipalo et al. found agreement ranging from 83-86% at 0.5 ng/mL
for four Diazyme methods, while agreement increased to 90-92% at 2.0 ng/mL and to 98% at 10.0 ng/mL
(137). Similarly, Lippi et al. reported agreement between 83% (at 0.25 ng/mL) and 96% (10 ng/mL) (131).

Other PCT assays
Assays not classified as using BRAHMS or Diazyme-licensed PCT reagents include the Beckman Access
assays and Snibe MAGLUMI. One published study was included which compared each of these methods
to the Kryptor method (131). While the Access methods showed reasonable agreement to the Kryptor
method, the MAGLUMI showed a large positive proportional bias (slope=1.51) that yielded
overestimates from the Kryptor (Suppl. Table 5). Agreement with the Kryptor at clinical decision points
was at least 96% across all concentrations (0.10, 0.25, 0.5, 2.0, and 10.0 ng/mL) for the Access assays,
while agreement for the MAGLUMI ranged from 91-96%.

Key summary points
- BRAHMS-licensed PCT immunoassays have demonstrated good overall correlation with the Kryptor
  method and have shown high categorical agreement around common clinical decision points. Non-
  BRAHMS assays have generally shown reduced correlation to the Kryptor.
- Biases across methods may impact reference intervals and the interpretation of PCT results across
  clinical decision ranges.

Are clinical decision points (cut-offs) comparable across PCT assays?
While there is currently no true reference method for PCT, an assay’s agreement with the BRAHMS PCT
sensitive Kryptor assay provides a basis to determine the feasibility of use of common PCT clinical
decision points. Several published studies, described above, demonstrate that while the various PCT methods generally compare to the BRAHMS PCT sensitive Kryptor with reasonable agreement, there are minor to moderate biases in patient sample materials that suggest that method harmonization has not been achieved. These biases preclude PCT from being trended across multiple methods and require the use of assay-dependent reference intervals. Importantly, these biases also impact the interpretation of PCT results across clinical decision ranges, especially at low PCT concentrations, which have the highest implications for diagnostic purposes and antibiotic decision making. Our review indicates that assays which use BRAHMS-licensed reagents have demonstrated stronger correlation to the BRAHMS PCT sensitive Kryptor method than other assays that have been introduced more recently. The analytical differences observed in these correlation studies could be attributed to many factors, including the various detection methods used by the available immunoassays as well as the PCT antibodies used in the assay reagents. Assay manufacturers typically provide limited to no information regarding the antibody design or specific PCT epitopes targeted in their assays, though BRAHMS assays have been reported to employ two monoclonal antibodies on the calcitonin and katacalcin segments, respectively (128).

Notably, the IFCC has an active Working Group on Standardization of Procalcitonin Assays (PCT-WG) seeking to develop and validate standard reference materials and a reference measurement procedure for PCT by stable isotope dilution mass spectrometry (138). The IFCC PCT-WG has indicated that PCT assays belonging to the BRAHMS family participate in a harmonization program using the BRAHMS PCT sensitive Kryptor as a reference method, which supports our findings that BRAHMS assays correlate better with the Kryptor than non-BRAHMS assays (128). However, there remains a need for higher order reference material such that all PCT assays could be harmonized. In the meantime, laboratories seeking to newly implement PCT methods should perform robust accuracy studies to a reference method, preferably the BRAHMS PCT sensitive Kryptor or a BRAHMS method, and should
carefully consider the implications for clinical interpretation. Further, each laboratory should establish or verify a reference interval specific to the chosen platform. Laboratories should work closely with clinical colleagues, such as those in Infectious Diseases and on antimicrobial stewardship teams, to align PCT interpretive algorithms with the chosen analytical method.

Key summary points

- There is not yet a reference method for PCT. In the meantime, the BRAHMS PCT sensitive Kryptor assay should be considered the gold standard since it was used in most of the initial clinical trials that first established PCT clinical decision points.

- The same PCT method should be used to trend PCT values for the same patient, as there is currently an absence of method harmonization, and biases exist between current methods.

- Labs implementing PCT or changing PCT methods should verify or establish the reference interval of their method and should perform robust correlation studies to the BRAHMS PCT sensitive Kryptor or another BRAHMS method using patient samples across clinically-relevant concentration ranges.

What are possible confounding factors for the interpretation of PCT results?

Proper interpretation of PCT results requires careful consideration of the patient’s clinical condition and the myriad of factors that can influence PCT. While PCT is often used as a surrogate marker for bacterial infections, there are conditions where elevated PCT can be observed in the absence of bacterial infection. These clinical situations include inflammatory events such as severe trauma, various major surgeries, and cardiogenic shock. Patients under these circumstances typically require monitoring for systemic infection. However, PCT should be interpreted with caution in the context of these non-bacterial elevations, particularly if acute inflammatory conditions occur in-between trended PCT measurements. While PCT may show general elevations in burns, some studies have suggested that PCT
may still be able to distinguish septic from non-septic burn patients (67,142). Some non-bacterial infections, such as malaria and some fungal infections, have also shown to non-consistently elevate PCT (143,144). Further, patients with significantly compromised renal function may not be able to clear PCT at a normal rate, and this can potentially cause PCT elevations (139). Finally, there is a natural elevation of PCT in healthy neonates just after birth (145). While there is growing evidence that PCT may have a role for guidance of antibiotic therapy in neonates, neonatal-specific reference ranges should be used. Generally, in cases of PCT elevations due to non-bacterial inflammatory processes, traditional PCT clinical decision points for outcome prediction and antimicrobial therapy decisions will not be valid. Alternatively, the kinetics (increases or decreases over time) of PCT may be monitored and considered in the context of potential background increases.

### Key summary points

- Various clinical scenarios outside of bacterial infection may cause elevated PCT results, including trauma, surgery, shock, and renal dysfunction.
- PCT should be interpreted with caution in clinical settings of non-bacterial inflammatory processes and should not be trended around acute inflammatory events.
- Traditional PCT clinical decision points should not be used in the setting of non-bacterial PCT elevations.
- Age-specific reference ranges should be used to interpret PCT in neonates.

### Conclusions

While the literature has accumulated with RCTs investigating the use of PCT, increasingly in the United States with the more recent availability of FDA-approved assays, there exists considerable variabilities in study designs and study populations that make it difficult to provide specific evidence-based recommendations for PCT protocols. In general, evidence to support the use of PCT to guide antibiotic
cessation is compelling, particularly in the critically ill and in some LRTIs, but is lacking in other clinical scenarios. Current data on the utility of PCT to guide the initiation of antibiotics is limited and does not demonstrate a benefit. In the pediatric and neonatal populations, some studies have established a role for PCT-guided protocols in reducing antibiotic exposure; however, the utility of PCT has not yet been well-studied in pre-term infants. While elevations in PCT are generally correlative with poor outcomes, no consistent PCT concentration(s) have been established to predict mortality. PCT interpretation guidance should consider the analytical method used, ideally its comparison to the BRAHMS Kryptor method, and patients should be trended using the same analytical assay. Improved outcomes from PCT implementation are more likely to be realized when the test is used in conjunction with antimicrobial stewardship programs, institutional interpretive algorithms, and clinical decision support tools. Successful implementation of clinical PCT requires a multidisciplinary effort amongst laboratorians, pharmacists, and infectious disease providers.


to guide duration of antibiotic therapy in intensive care patients: a randomized prospective

treatment duration in septic patients: a randomized trial. Am J Respir Crit Care Med.
2008;177:498–505.

15. Huang DT, Yealy DM, Filbin MR, Brown AM, Chang C-CH, Doi Y, et al. Procalcitonin-
2018;379:236–49.

algorithm to guide initial antibiotic therapy in acute exacerbations of COPD admitted to the

Guideline-Based Clinical Assessment Versus Procalcitonin-Guided Antibiotic Use in


Procalcitonin usefulness for the initiation of antibiotic treatment in intensive care unit

guided interventions against infections to increase early appropriate antibiotics and improve

Procalcitonin Algorithm in Critically Ill Adults with Undifferentiated Infection or Suspected
Sepsis. A Randomized Controlled Trial. Am J Respir Crit Care Med. American Thoracic

guided antibiotic treatment in patients with positive blood cultures: A patient-level meta-

versus C-reactive protein for guiding antibiotic therapy in sepsis: a randomized trial. Crit
Care Med. 2013;41:2336–43.

guided antibiotic treatment on mortality in acute respiratory infections: a patient level meta-

Management of acute heart failure with Procalcitonin in EUnorpe: results of the randomized


2017;318:1241.


80. Jones AE. Lactate Clearance vs Central Venous Oxygen Saturation as Goals of Early Sepsis TherapyA Randomized Clinical Trial. JAMA. 2010;303:739.


Figures

Figure 1. Schematic of cytokine-mediated inflammatory host response pathway leading to adipocyte secretion of procalcitonin (top) compared to normal physiological secretion of calcitonin from thyroidal C-cells (bottom).
Figure 2. Relative kinetic expression pattern of procalcitonin upon inflammatory insult as compared to other inflammatory markers. Procalcitonin increases in the plasma within 2-6 hours, peaks at approximately 12 hours, and has a half-life of about 24 hours. Adapted from reference 2.
Table 1. Normal hourly values of post-birth PCT

<table>
<thead>
<tr>
<th>Time after birth (hours)</th>
<th>PCT (ng/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-6</td>
<td>0.5</td>
</tr>
<tr>
<td>6-12</td>
<td>2</td>
</tr>
<tr>
<td>12-18</td>
<td>5</td>
</tr>
<tr>
<td>18-36</td>
<td>10</td>
</tr>
<tr>
<td>36-48</td>
<td>5</td>
</tr>
<tr>
<td>48-60</td>
<td>2</td>
</tr>
<tr>
<td>60-72</td>
<td>1</td>
</tr>
<tr>
<td>&gt;72</td>
<td>0.5</td>
</tr>
</tbody>
</table>

*PCT cutoffs used in NeoPlns trial (ref 99)
### Table 2. Summary of FDA-approved PCT assays and associated reagent licensing

<table>
<thead>
<tr>
<th>Manufacturer (platform)</th>
<th>BRAHMS PCT a</th>
<th>Diazyme PCT b</th>
<th>Other b</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abbott (Alinity)</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abbott (Architect)</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Beckman Coulter (AU)</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Beckman Coulter (Unicel, Access)</td>
<td></td>
<td></td>
<td>(Access PCT)</td>
</tr>
<tr>
<td>bioMerieux (VIDAS)</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BRAHMS (Kryptor)</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DiaSorin (LIAISON)</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diazyme (DZ-Lite)</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Fujirebio (Lumipulse)</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ortho-Clinical (VITROS)</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Roche (cobas)</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Siemens (Atellica)</td>
<td>X</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**FDA approved indications:**

*a* to aid in the risk assessment of critically ill patients on their first day of ICU admission for progression to severe sepsis and septic shock; to aid in assessing the cumulative 28-day risk of all-cause mortality for patients diagnosed with severe sepsis or septic shock in the ICU or when obtained in the emergency department or other medical wards prior to ICU admission, using a change in PCT level over time; to aid in decision making on antibiotic therapy for patients with suspected or confirmed lower respiratory tract infections (LRTI) defined as community-acquired pneumonia (CAP), acute bronchitis, and acute exacerbation of chronic obstructive pulmonary disease (AECOPD) – in an inpatient setting or an emergency department; to aid in decision making on antibiotic discontinuation for patients with suspected or confirmed sepsis

*b* to aid in the risk assessment of critically ill patients on their first day of ICU admission for progression to severe sepsis and septic shock
**Supplementary Table 1.** Summary of characteristics for 8 trials included in the analysis of procalcitonin-guided antibiotic discontinuation for patients with sepsis in the intensive care unit

<p>| Author, Year, Country | N   | Assay Used                                                                 | Frequency of PCT Measurement and Notifications | PCT Criteria for Stopping Antibiotics | Compliance Rates | Findings – Antibiotic Use                                                                 | Study Strength |
|------------------------|-----|----------------------------------------------------------------------------|-----------------------------------------------|--------------------------------------|------------------|------------------------------------------------------------------------------------------|----------------|----------------------------------|
| Bloos, 2016, Germany(1)| 1089| Not explicitly mentioned – described as time-resolved amplified cryptate emission technology | Days 0, 1, 4, 7, 10 and 14 if patient still in ICU. Frequency of notification not reported. | Only on days 7, 10 or 14: PCT &lt;1 ng/mL or ≥ 50% drop compared with previous value | 79% overall, 40.9% by day 7 | 4.5% reduction in antimicrobial exposure, no difference in 28-day mortality. | 2, RCT         |
| De Jong, 2016, France(2)| 1546| Kryptor or Vidas depending on site                                          | Daily until ICU discharge or until 3 days after systemic antibiotics are stopped. Frequency of notification not reported. | PCT &lt; 0.5 ng/mL or ≥ 80% drop from peak value | 44% for patients with antibiotics stopped within 24h and 53% for those with antibiotics | 2-day reduction in antibiotic therapy. Decrease in 28-day and 1-year mortality. | 2, RCT         |</p>
<table>
<thead>
<tr>
<th>Study</th>
<th>PCT Cut-Off</th>
<th>Test Method</th>
<th>Frequency of Notifications</th>
<th>PCT &lt; 1 ng/mL</th>
<th>PCT &lt; 0.25 ng/mL and Low Clinical Suspicion for Infection or ≥ 90% Drop from Peak Value</th>
<th>9 vs 11 Days (p=0.58), 90-Day Mortality 18% vs 16% (p=0.54)</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shehabi, 2014, Australia(3)</td>
<td>394</td>
<td>Not explicitly mentioned – “automated immunoassay analyzers”</td>
<td>Daily until the first of ICU discharge or 7 days. Frequency of notifications not reported.</td>
<td>PCT &lt; 1 ng/mL, PCT &lt; 0.25 ng/mL and low clinical suspicion for infection or ≥ 90% drop from peak value</td>
<td>97%</td>
<td>2, RCT</td>
<td></td>
</tr>
<tr>
<td>Annane, 2013, France(4)</td>
<td>62</td>
<td>BRAHMS PCT-sensitive Kryptor assay</td>
<td>6h, day 3 and day 5. Frequency of notifications not reported.</td>
<td>PCT &lt; 0.5 ng/mL</td>
<td>81% at 6h, 83% on day 3 and 63% on day 5</td>
<td>14% reduction in # of patients on antibiotics at day 5, non-significant (95% CI 0.6 to 1.14)</td>
<td>3, RCT</td>
</tr>
<tr>
<td>Deliberato, 2013, Brazil(5)</td>
<td>81</td>
<td>VIDAS BRAHMS PCT</td>
<td>Day 0, 5 or 7 then every 48h until hospital discharge, or antibiotics stopped. Frequency of notifications not reported.</td>
<td>PCT &lt; 0.5 ng/mL or ≥ 90% drop from peak value</td>
<td>100%</td>
<td>3, RCT</td>
<td></td>
</tr>
<tr>
<td>Study Authors and Year, Location</td>
<td>Patient ID</td>
<td>Study Design</td>
<td>Initial Treatment Schedule</td>
<td>PCT Criteria for Stopping Antibiotics</td>
<td>Percentage of Patients Stopping Antibiotics</td>
<td>Days of Antibiotic Use Difference</td>
<td>Additional Observations</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>------------</td>
<td>--------------</td>
<td>----------------------------</td>
<td>--------------------------------------</td>
<td>------------------------------------------</td>
<td>-------------------------------</td>
<td>-------------------------</td>
</tr>
<tr>
<td>Bouadma, 2010, France(6)</td>
<td>618</td>
<td>BRAHMS Kryptor Day 0, then daily until treatment was finished. Results available within 2h of blood draw.</td>
<td>PCT &lt; 0.5 ng/mL or &gt; 80% drop from peak value</td>
<td>28.7%</td>
<td>2.7 day difference in antibiotic use (p&lt;0.001)</td>
<td>2, RCT</td>
<td></td>
</tr>
<tr>
<td>Hochreiter, 2009, Germany(7)</td>
<td>110</td>
<td>Brahms PCT LIA Not reported</td>
<td>PCT &lt; 1 ng/mL or &gt; 25-35% drop from peak value over 3 days</td>
<td>Not reported</td>
<td>5.9 vs 7.9 days, p&lt;0.001</td>
<td>3, RCT</td>
<td></td>
</tr>
<tr>
<td>Nobre, 2008, Switzerland(8)</td>
<td>79</td>
<td>Brahms Kryptor PCT Day 0, then Day 3 if initial PCT &lt; 1 ng/mL or day 5 if initial PCT &gt; 1 ng/mL. Results available within 3h of blood draw.</td>
<td>If baseline &gt; 1: stop when (1) PCT &lt; 0.25 ng/mL or (2) &gt; 90% drop from peak value If baseline &lt; 1: stop when PCT &lt;0.1 ng/mL</td>
<td>81%</td>
<td>3.5-day shorter median duration of antibiotic therapy. Similar mortality. 2-day shorter ICU stay in PCT group (p=0.03)</td>
<td>3, RCT</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: RCT = Randomized Control Trial, ICU = intensive care unit
**Supplementary Table 2.** Summary of characteristics for 11 trials included in the analysis of procalcitonin-guided antibiotic discontinuation for patients with lower respiratory tract infections

<table>
<thead>
<tr>
<th>Author, Year, Country</th>
<th>Disease State</th>
<th>N</th>
<th>Assay Used</th>
<th>Frequency of PCT Measurement and Notifications</th>
<th>PCT Criteria for Stopping Antibiotics</th>
<th>Compliance Rates</th>
<th>Findings – Antibiotic use</th>
<th>Study Strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Montassier, 2019, France(9)</td>
<td>CAP</td>
<td>285</td>
<td>BRAHMS Kryptor</td>
<td>On enrollment and days 0, 3, 5 and 7. Results available within 1 hour.</td>
<td>PCT &lt;0.25 ng/mL</td>
<td>76%</td>
<td>Mean antibiotic duration 9 days (PCT) vs. 10 days (control) (p=0.21). Rates of serious adverse outcomes at 30 days were 15% (PCT) vs. 20% (control) (95% CI -4% to 14%)</td>
<td>2, RCT</td>
</tr>
<tr>
<td>Wussler, 2019, Switzerland(10)</td>
<td>CAP</td>
<td>45</td>
<td>VIDAS BRAHMS</td>
<td>On enrollment and day 5. Notification frequency not mentioned.</td>
<td>PCT &lt;0.25 ng/mL or &gt; 80% drop from PCT level at randomization</td>
<td>Not reported</td>
<td>Median antibiotic duration 10.5 days in each group (p=0.387). In-hospital mortality 5 patients (PCT) vs. 4 patients (control) (p=1)</td>
<td>3, RCT</td>
</tr>
<tr>
<td>Author, Year, Location</td>
<td>Disease</td>
<td>Code</td>
<td>Test</td>
<td>Enrollment Strategy</td>
<td>PCT Cut-off</td>
<td>Notification Frequency</td>
<td>Outcomes</td>
<td>Conclusion</td>
</tr>
<tr>
<td>------------------------</td>
<td>---------</td>
<td>------</td>
<td>------</td>
<td>---------------------</td>
<td>-------------</td>
<td>-----------------------</td>
<td>----------</td>
<td>------------</td>
</tr>
<tr>
<td>Daubin, 2018, France(11)</td>
<td>COPD</td>
<td>302</td>
<td>Elecsys BRAHMS PCT</td>
<td>6h after enrollment, then days 1, 3 and 5. Notification frequency not mentioned.</td>
<td>PCT &lt;0.25 ng/mL or &gt; 90% drop from PCT level at randomization</td>
<td>Not reported</td>
<td>Mean antibiotic duration 7.9 days (PCT) vs. 7.7 days (control) (p=0.75). 90-day mortality 20% (PCT) and 14% (control) (95% CI for adjusted difference -0.3 to 13.5%)</td>
<td>3, RCT</td>
</tr>
<tr>
<td>Huang, 2018, USA(12)</td>
<td>LRTI</td>
<td>1656</td>
<td>Vidas BRAHMS</td>
<td>Enrollment, days 1, 3, 5 and 7 if still hospitalized and receiving antibiotics. Notification frequency not mentioned.</td>
<td>PCT &lt;0.25 ng/mL</td>
<td>64.8%</td>
<td>Mean antibiotic duration 4.2 (PCT) vs. 4.3 (control) days (p=0.87). Proportion of patients with adverse outcomes 11.7% (PCT) vs. 13.1% (control) (p=&lt;0.001 for noninferiority)</td>
<td>1, RCT</td>
</tr>
<tr>
<td>Corti, 2016, Denmark(13)</td>
<td>COPD</td>
<td>120</td>
<td>Mini Vidas</td>
<td>Within 24h of admission, and on days 3, 5 and 7.</td>
<td>PCT &lt;0.25 ng/mL or &gt; 80% drop from peak value</td>
<td>61% on day 1</td>
<td>Median antibiotic duration 3.5 (PCT) vs. 8.5 (control) days (p=0.02). Mortality</td>
<td>3, RCT</td>
</tr>
<tr>
<td>Study</td>
<td>Disease</td>
<td>Size</td>
<td>Test 1</td>
<td>Test 2</td>
<td>Test 3</td>
<td>Test 4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------------------------</td>
<td>---------</td>
<td>------</td>
<td>--------</td>
<td>--------</td>
<td>--------</td>
<td>--------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Branche, 2015, USA(14)</td>
<td>LRTI</td>
<td>200</td>
<td>VIDAS</td>
<td>BRAHMS</td>
<td>On enrollment and 12h later. Notification frequency not mentioned</td>
<td>PCT &lt; 0.25 ng/mL</td>
<td>64%</td>
<td>Median antibiotic duration 3 (PCT) vs 4 (control) days (p = 0.71). No deaths occurred during hospitalization.</td>
</tr>
<tr>
<td>Long, 2011, China(15)</td>
<td>CAP</td>
<td>156</td>
<td>BRAHMS</td>
<td>Kryptor</td>
<td>At admission and days 3, 6 and 8. Frequency not mentioned.</td>
<td>PCT &lt; 0.25 ng/mL</td>
<td>84.4%</td>
<td>Mean antibiotic duration 5 (PCT) vs 7 (control) days (p&lt;0.001). No deaths in either group within 28 days.</td>
</tr>
<tr>
<td>Schuetz, 2009, Switzerland(1)</td>
<td>LRTI</td>
<td>233</td>
<td>Kryptor</td>
<td>PCT</td>
<td>At admission and on days 3, 5, 7 and at hospital discharge. Results available within 1 hour.</td>
<td>PCT &lt;0.25 ng/mL or &gt; 80% drop from peak value</td>
<td>71%</td>
<td>Mean duration of antibiotics 5.7 (PCT) vs. 8.7 days (control) (relative change - 34.8%, 95% CI -40.3% to -28.7%).</td>
</tr>
<tr>
<td>Study</td>
<td>Infection Type</td>
<td>Sample Size</td>
<td>Test</td>
<td>Antibody Delivery</td>
<td>Antibody at Admission and</td>
<td>Antibody Duration</td>
<td>Mortality Rate</td>
<td>Mortality Rate Difference</td>
</tr>
<tr>
<td>-------</td>
<td>----------------</td>
<td>-------------</td>
<td>------</td>
<td>-------------------</td>
<td>--------------------------</td>
<td>------------------</td>
<td>---------------</td>
<td>--------------------------</td>
</tr>
<tr>
<td>Stolz, 2009, Switzerland (17)</td>
<td>VAP</td>
<td>101</td>
<td>BRAHMS Kryptor</td>
<td>At admission and on day 2, then daily until day 10. Available within 1 hour.</td>
<td>PCT &lt;0.5 ng/mL or ≥ 80% drop from peak value</td>
<td>84%</td>
<td>27% reduction in antibiotic duration in PCT group compared to control (p=0.038). Mortality rate at 28-days was 16% (PCT) vs. 24% (control) (p=0.327).</td>
<td></td>
</tr>
<tr>
<td>Briel, 2008, Switzerland (18)</td>
<td>LRTI</td>
<td>455</td>
<td>Kryptor</td>
<td>At admission and on day 3. Results available in 2-4 hours.</td>
<td>PCT &lt;0.25 ng/mL</td>
<td>85%</td>
<td>Mean antibiotic duration 6.2 days (PCT) vs. 7.1 days (control), adjusted decrease with PCT 1 day (95% CI 0.4 to 1.7 days). No deaths in PCT group, 1 patient in the control group died.</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Country</td>
<td>Study Design</td>
<td>N</td>
<td>Comparator</td>
<td>Intervention</td>
<td>End Point</td>
<td>Outcome Measures</td>
</tr>
<tr>
<td>-------</td>
<td>------</td>
<td>---------</td>
<td>--------------</td>
<td>---</td>
<td>------------</td>
<td>-------------</td>
<td>-----------</td>
<td>------------------</td>
</tr>
<tr>
<td>Christ-Crain, 2006, Switzerland(19)</td>
<td>CAP</td>
<td>BRAHMS Kryptor</td>
<td>At admission and on days 4, 6 and 8. Results available within 1 hour.</td>
<td>PCT &lt;0.25 ng/mL</td>
<td>Not reported</td>
<td>Median antibiotic duration 5 (PCT) vs. 12 (control) days (p&lt;0.001). Mortality not reported.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Beye, 2019, France(20)</td>
<td>VAP</td>
<td>157</td>
<td>Daily while patient receiving antibiotics. Notification frequency not mentioned.</td>
<td>PCT &lt; 0.5 ng/mL or &gt; 80% drop from peak value if initial PCT &gt; 0.5 ng/mL</td>
<td>48%</td>
<td>Median antibiotic duration 8 (PCT) vs. 9.5 (control) days. (p=0.002). Mortality not reported.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Newton, 2019, USA(21)</td>
<td>Any infection</td>
<td>857</td>
<td>Provider discretion</td>
<td>No formal algorithm</td>
<td>73.7% ASP rec compliance</td>
<td>Reduction of mean length of therapy of 3, retrospective analysis</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Non-RCTs**
<table>
<thead>
<tr>
<th>Study</th>
<th>Setting</th>
<th>Test</th>
<th>Testing Schedule</th>
<th>PCT Threshold</th>
<th>Turnaround Time</th>
<th>Duration of Antibiotics</th>
<th>Mortality Evaluation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Akagi, 2019, Japan(22)</td>
<td>CAP 335</td>
<td>Spherelight Brahms PCT</td>
<td>Days 5, 8 and 11 and every 3 days thereafter if needed. Result turnaround time not reported.</td>
<td>PCT &lt; 0.2 ng/mL</td>
<td>Not reported</td>
<td>Median duration 8 (PCT) vs. 11 (control) days (p&lt;0.001). Mortality not evaluated.</td>
<td>3, historical control study</td>
</tr>
<tr>
<td>Townsend, 2018, USA(23)</td>
<td>LRTI 374</td>
<td>BRAHMS Kryptor</td>
<td>Day 0, then day 2 if admitted to floor or day 1 if admitted to ICU. Testing once daily.</td>
<td>≥ 80% drop from peak value</td>
<td>70%</td>
<td>Median duration of antibiotics 6 (PCT) vs. 7 (control) days (p=0.045). 30-day mortality 4% (PCT) vs. 4.5% (control) (p=0.82).</td>
<td>2, Pre-post study</td>
</tr>
</tbody>
</table>

Abbreviations: RCT = Randomized Control Trial, CAP = Community Acquired Pneumonia, LRTI = Lower Respiratory Tract Infection, VAP = Ventilator-associated Pneumonia, COPD = Chronic Obstructive Pulmonary Disease
**Supplementary Table 3.** Clinical trials investigating a single Procalcitonin measurement or PCT Clearance, defined as a decrease in PCT concentrations over time, as a predictor of all cause 28/30-day mortality

<table>
<thead>
<tr>
<th>Author, Year, Country</th>
<th>Patient Population</th>
<th>Study Type</th>
<th>No. of Subjects</th>
<th>PCT Assay</th>
<th>When was PCT measured?</th>
<th>PCT Cutoff(s) (ng/mL)</th>
<th>PCT Clearance %</th>
<th>Significant findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sager, 2017, multinational (France, Switzerland, USA)* (24)</td>
<td>ED</td>
<td>Multicenter, prospective observational (TRIAGE trial)</td>
<td>6970</td>
<td>Kryptor®, B.R.A.H.M.S.</td>
<td>During ED stay</td>
<td>0.05, 0.1, 0.25, and 0.5</td>
<td>n/a</td>
<td>AUC = 0.75 for entire cohort, 0.85 for metabolic disease and 0.82 for cardiovascular disease; Mortality increased stepwise within higher PCT cut-offs (0.05, 0.1, 0.25, 0.5 ng/mL) from 1%, 3%, 7%, 13% to 15%, respectively</td>
</tr>
<tr>
<td>Shim, 2019, Korea (25)</td>
<td>ED patients with suspected infection</td>
<td>Prospective observational</td>
<td>199</td>
<td>HUBI-PCT, Humasis (POC-WB) Roche BRAHAMS (serum)</td>
<td>At presentation</td>
<td>n/a</td>
<td>n/a</td>
<td>AUC = 0.694 for POC-WB and 0.731 for serum</td>
</tr>
<tr>
<td>Gonzalez del Castillo, 2019, Spain (26)</td>
<td>ED patients with suspected infection</td>
<td>Prospective</td>
<td>684</td>
<td>POC (Samsung LABGEO IB10, Nexus, USA)</td>
<td>At presentation and at 72 hours</td>
<td>0.2 at presentation, 1.78 at 72 hrs</td>
<td>n/a</td>
<td>AUC = 0.68 at baseline; AUC = 0.63 at 72 hours</td>
</tr>
<tr>
<td>Schuetz, 2017, US (27)</td>
<td>ICU patients with severe sepsis and septic shock</td>
<td>Blinded prospective, multicenter trial (MOSES Trial)</td>
<td>858</td>
<td>Kryptor®, B.R.A.H.M.S.</td>
<td>Daily for 5 days</td>
<td>&lt;0.5, 0.5-2.0 and &gt;2.0</td>
<td>Decrease by &gt;80%</td>
<td>AUC = 0.56 at baseline AUC = 0.621 PCT change Day 4 vs day 0; 20.0% mortality in patients in which PCT decreased by ≤ 80% by day 4 (HR 1.97, multivariate, p=0.009) vs 10.4% (p=0.001)</td>
</tr>
<tr>
<td>Elke, 2018, Germany, multisite (28)</td>
<td>ICU patients with sepsis and septic shock</td>
<td>Secondary analysis of an RCT (SISPCT)</td>
<td>1089</td>
<td>PCT Kryptor®, B.R.A.H.M.S.</td>
<td>Enrollment and days 1, 4, 7, 10</td>
<td>1.02 Low severity; 47 in high severity - at baseline - cut off varied by day</td>
<td>Decrease by ≥50%</td>
<td>AUC = 0.56 at baseline (septic patients) AUC = 0.5 at baseline (septic shock patients) 29.5% mortality in patients in which Patients with PCT decreased &lt;50% between baseline and day 4 by &lt;50% between baseline and day 4 vs 17.1% (P&lt;0.001)</td>
</tr>
<tr>
<td>Study</td>
<td>Type</td>
<td>Patients/Setting</td>
<td>Prospective/Retrospective</td>
<td>ED, D (HITEMP)</td>
<td>Retrospective V (TRIAGE)</td>
<td>Validation D (HITEMP)</td>
<td>Validation V (TRIAGE)</td>
<td>PCT Cut-off</td>
</tr>
<tr>
<td>------------------</td>
<td>-----------------------------</td>
<td>----------------------------------------------------------------------------------</td>
<td>----------------------------</td>
<td>---------------</td>
<td>--------------------------</td>
<td>-----------------------</td>
<td>-----------------------</td>
<td>--------------</td>
</tr>
<tr>
<td>Saeed, 2019</td>
<td>ED patients with suspected infection</td>
<td>Derivation cohort (D) - England, France, Italy, Sweden and Spain and Validation cohort (V) - France, Switzerland and the US</td>
<td>Prospective (D)- (HITEMP) and retrospective (V) - (TRIAGE)</td>
<td>1175 (D)</td>
<td>Kryptor®, B.R.A.H.M.S.</td>
<td>During ED stay</td>
<td>0.21 (D)</td>
<td>0.15 (V)</td>
</tr>
<tr>
<td>Osawa, 2020</td>
<td>Patients with TB</td>
<td>Prospective, observational</td>
<td>μTASWako i30 immunoanalyzer (Fujifilm Wako Pure Chemical)</td>
<td>Days 0, 7, 14, and 28</td>
<td>Varied by day: 0.13 (day 0), 0.05 (day 7), 0.12 (day 14), and 0.06 (day 28)</td>
<td>n/a</td>
<td>PCT concentrations above the cut-off predicted mortality on days 0, 7, 14, and 28 days with OR of 7.9, 14.3, 20.0 and 7.3, respectively (p≤0.005).</td>
<td></td>
</tr>
<tr>
<td>Peng, 2020</td>
<td>ICU patients w/ abdominal sepsis</td>
<td>Retrospective</td>
<td>Roche Brahms</td>
<td>ICU admission and for the next 5 days</td>
<td>&gt;70% decrease</td>
<td>AUC = 0.726 PCT Clearance day 1 to day 5 OR 0.16 (P&lt;0.001)– PCT Clearance (day 5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shi, 2015</td>
<td>ICU patients w/ AID</td>
<td>Prospective, observational</td>
<td>Brahms Diagnostica</td>
<td>ICU admission and days 1,3,5, and 7</td>
<td>n/a</td>
<td>≥50% decrease</td>
<td>OR 5.1 (P= 0.01)-PCT clearance &lt; 50% (day 5)</td>
<td></td>
</tr>
<tr>
<td>Yu 2019</td>
<td>ED or admitted patients with systemic infection</td>
<td>Retrospective</td>
<td>Biomerieux Vidas</td>
<td>Within 24 hours of admission</td>
<td>&lt;0.25, 0.25 to 2, and &gt;2</td>
<td>n/a</td>
<td>AUC = 0.73 PCT + qSOFA</td>
<td></td>
</tr>
<tr>
<td>Ryoo, 2019</td>
<td>ED patients with septic shock</td>
<td>Retrospective</td>
<td>N/A</td>
<td>Prior to resuscitation and antibiotic administration</td>
<td>&gt;17.0</td>
<td>n/a</td>
<td>Alone, PCT not associated with mortality but OR= 1.55 (P = 0.001) PCT+ CRP &gt; cutoff vs both not elevated</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Setting</td>
<td>Participants</td>
<td>Design</td>
<td>Main Findings</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>---------</td>
<td>--------------</td>
<td>--------</td>
<td>---------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bloos, 2011, Canada, Europe and US (35)</td>
<td>ICU patients with pneumonia and mechanical ventilation</td>
<td>multicenter, international, observational</td>
<td>Lumitest B.R.A.H.M.S. GmbH</td>
<td>Within 48 hours of enrollment and then daily for 14 days</td>
<td>AUC = 0.7 Baseline AUC = 0.74 Max Concentration</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sakran, 2012 US (36)</td>
<td>Trauma ICU Patients</td>
<td>Prospective Observational</td>
<td>Brahms Kryptor</td>
<td>0, 6, 24 hours, daily until death or 30 days</td>
<td>OR 3.65 (p=0.04)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kutz, A 2015, multinational (37)</td>
<td>Primary Care, ED, and ICU patients with Acute ARI</td>
<td>Secondary analysis of a meta-analysis</td>
<td>Brahms Kryptor</td>
<td>At primary care visit, ED admission and ICU admission</td>
<td>AUC = 0.50 (PC), p= 0.6 AUC = 0.67 (ED), p &lt; 0.001 AUC = 0.71 (ICU – LRI), P&lt;0.001</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: ARI, Acute respiratory tract infection
**Supplementary Table 4. Summary of pediatric PCT trials**

<table>
<thead>
<tr>
<th>Author, Year, Country</th>
<th>N</th>
<th>Age Groups</th>
<th>Assay Used</th>
<th>Cut-off/interoperative Criteria</th>
<th>Frequency of PCT Measurement and Notifications</th>
<th>PCT Criteria for Stopping Antibiotics</th>
<th>Compliance Rates</th>
<th>Findings</th>
<th>Study Strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>NeoPIns 2017, 18 hospitals across EU and Canada. (38)</td>
<td>866 and 844 in experimental and control arm, respectively</td>
<td>neonates gestational age &gt;34 weeks with suspected sepsis in 72 hr of life</td>
<td>Roche Elecsys (LOQ 0.06 ng/mL), Kryptor, Roche Diagnostics immunoassay analyzer (LOQ 0.02 ng/mL)</td>
<td>Cut-offs ranging from 0.5-10 ng/mL stratified based on time after birth</td>
<td>12, 24, 36-72, and 24-48 hrs on indication until discontinuation of antibiotic treatment</td>
<td>2 consecutive PCT values within normal range</td>
<td>recommendations overruled in 25% of neonates</td>
<td>10 hr reduction in antimicrobial exposure. 3 hr reduction in hospital stay. Impact on re-infection or death (during 1st month of life) could not be assessed due to low occurrence</td>
<td>2</td>
</tr>
<tr>
<td>Stocker et al., 2010, Swizterland(39)</td>
<td>121 newborns, 61 in control arm and 60 in experimental arm</td>
<td>neonates gestational age &gt;34 weeks</td>
<td>Roche BRAHMS Kryptor</td>
<td>for 18-34 hr &lt;10 ng/mL defined as normal (50% of the highest PCT concentration observed in neonates with respiratory distress without infection)</td>
<td>blood collected 0, 36-72, and 24-48 hrs until discontinuation of antibiotic treatment. In PCT arm additional sample at 12 hr (+/- 6hr)</td>
<td>2 consecutive PCT values within pre-defined age-adjusted cut-off values</td>
<td>recommendation to discontinue abx overruled in 2/60 patients in PCT arm</td>
<td>22.4 hr reduction in abx exposure. Unable to assess hospital stay</td>
<td>2</td>
</tr>
<tr>
<td>PROSACAB 2019, Spain(40)</td>
<td>114 in pre PCT-guided protocol group and 112 in post PCT-protocol group</td>
<td>&lt;18 years admitted after cardiac surgery; excluded neonates</td>
<td>ATOM SA, Brahms Diagnostica, Mi</td>
<td>decrease in PCT by 50% of maximum PCT value or &lt; 0.5 ng/mL</td>
<td>PCT measured daily during 3 days after surgery and continued if there was suspicion for infection until resolution</td>
<td>decrease in PCT by 50% of maximum PCT value or &lt; 0.5 ng/mL</td>
<td>26% in group 1 and 54.8% in group 2</td>
<td>1.1 day reduction in antibiotics exposure and 2 more antibiotic free days observed in PICU without adverse outcomes</td>
<td>2</td>
</tr>
</tbody>
</table>
**Supplementary Table 5.** Summary of comparison studies of various PCT assays to the BRAHMS PCT sensitive Kryptor

<table>
<thead>
<tr>
<th>Test method</th>
<th>Slope (95% CI, if provided)</th>
<th>Intercept (95% CI, if provided)</th>
<th>Correlation</th>
<th>Regression type</th>
<th>Number of samples</th>
<th>Specimen type</th>
<th>Specimen storage notes</th>
<th>Author, Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRAHMS procalcitonin assays</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ADVIA Centaur</td>
<td>0.80</td>
<td>0.40</td>
<td>r=0.984</td>
<td>Deming</td>
<td>100</td>
<td>serum</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
<td>Dipalo et al., 2015(41)</td>
</tr>
<tr>
<td>ARCHITECT</td>
<td>0.834 (0.808, 0.857)</td>
<td>-0.013 (-0.027, -0.010)</td>
<td>--</td>
<td>Passing-Bablok</td>
<td>192</td>
<td>serum and lithium-heparin plasma</td>
<td>Analysis performed over several days, storage not specified</td>
<td>Gruzdys et al., 2019(42)</td>
</tr>
<tr>
<td>ARCHITECT c16000</td>
<td>0.97</td>
<td>0.01</td>
<td>r=0.995</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
<td>Lippi et al., 2019(43)</td>
</tr>
<tr>
<td>ARCHITECT i2000SR</td>
<td>0.91 (0.87, 0.96)</td>
<td>-0.2 (-0.3, 0.0)</td>
<td>r=0.986 (0.984, 0.988)</td>
<td>Deming</td>
<td>584</td>
<td>plasma and serum</td>
<td>Analyzed either fresh or up to 24 h at 2-8°C</td>
<td>Soh et al., 2018(44)</td>
</tr>
<tr>
<td>ARCHITECT i2000</td>
<td>0.806 (0.722, 0.890)</td>
<td>-0.006 (-0.063, 0.050)</td>
<td>r=0.8864</td>
<td>Passing-Bablok</td>
<td>39</td>
<td>serum</td>
<td>One freeze-thaw cycle (-20°C), multicenter, all analyzed on same day</td>
<td>Chambliss et al., 2019(45)</td>
</tr>
<tr>
<td>Instrument</td>
<td>Lower limit (95% CI)</td>
<td>Upper limit (95% CI)</td>
<td>Correlation coefficient</td>
<td>Method</td>
<td>Samples</td>
<td>Conditions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>---------------------</td>
<td>----------------------</td>
<td>----------------------</td>
<td>-------------------------</td>
<td>-----------------</td>
<td>---------</td>
<td>-----------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BRAHMS PCT direct</td>
<td>0.90 (0.86, 0.96)</td>
<td>0.03 (0.02, 0.04)</td>
<td>r²=0.97</td>
<td>Passing-Bablok</td>
<td>202</td>
<td>capillary (fingertip) whole blood Samples analyzed fresh, methods run consecutively</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BRAHMS PCT direct</td>
<td>0.98 (0.93, 1.02)</td>
<td>0.02 (0.01, 0.03)</td>
<td>r²=0.95</td>
<td>Passing-Bablok</td>
<td>219</td>
<td>venous whole blood (EDTA) Samples analyzed fresh, methods run consecutively</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cobas e 411</td>
<td>0.795 (0.733, 0.857)</td>
<td>0.005 (−0.016, 0.050)</td>
<td>r=0.8974</td>
<td>Passing-Bablok</td>
<td>37</td>
<td>serum One freeze-thaw cycle (-20°C), multicenter, all analyzed on same day</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cobas e 601</td>
<td>1.12</td>
<td>-0.47</td>
<td>r=0.988</td>
<td>Deming</td>
<td>100</td>
<td>serum One freeze-thaw cycle (-70°C), multicenter, all analyzed on same day</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cobas e 602</td>
<td>0.9125</td>
<td>0.021</td>
<td>r=0.99</td>
<td>Passing-Bablok</td>
<td>239</td>
<td>serum Analyzed within 24 h stored at 2-8°C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cobas e 801</td>
<td>0.89</td>
<td>-0.01</td>
<td>r=0.985</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LIAISON</td>
<td>1.02</td>
<td>0.22</td>
<td>r=0.986</td>
<td>Deming</td>
<td>100</td>
<td>serum One freeze-thaw cycle (-70°C), multicenter,</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*a Kutz et al., 2016(46) | Chambliss et al., 2019(45) | Dipalo et al., 2015(41) | Ceriotti et al., 2017(47) | Lippi et al., Med, 2019(43) | Dipalo et al., 2015(41) |
<table>
<thead>
<tr>
<th>Instrument</th>
<th>B</th>
<th>Intercept (95% CI)</th>
<th>Correlation Coefficient</th>
<th>Method</th>
<th>Sample Type</th>
<th>Storage and Analysis Details</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>LIAISON XL</td>
<td>1.15</td>
<td>-0.02</td>
<td>r=0.997</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
</tr>
<tr>
<td>LIAISON XL</td>
<td>1.172143</td>
<td>-0.0102086</td>
<td>r=0.99</td>
<td>Passing-Bablok</td>
<td>171</td>
<td>serum</td>
<td>Remnant samples, storage not specified</td>
</tr>
<tr>
<td>Lumipulse G</td>
<td>1.14</td>
<td>-0.01</td>
<td>r=0.986</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
</tr>
<tr>
<td>Lumipulse G600II</td>
<td>1.001</td>
<td>-0.052</td>
<td>r²=0.99</td>
<td>Passing-Bablok</td>
<td>138</td>
<td>heparinized plasma</td>
<td>Samples analyzed fresh, methods run simultaneously</td>
</tr>
<tr>
<td>VIDAS</td>
<td>1.40</td>
<td>-0.09</td>
<td>r=0.979</td>
<td>Deming</td>
<td>100</td>
<td>serum</td>
<td>One freeze-thaw cycle (-70°C), multicenter, all analyzed on same day</td>
</tr>
<tr>
<td>VIDAS</td>
<td>1.19</td>
<td>-0.06</td>
<td>r=0.994</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
</tr>
<tr>
<td>Method</td>
<td>Slope</td>
<td>Intercept</td>
<td>Corr</td>
<td>Eq</td>
<td>N</td>
<td>Type</td>
<td>Conditions</td>
</tr>
<tr>
<td>----------------</td>
<td>---------</td>
<td>-----------</td>
<td>-------</td>
<td>--------</td>
<td>------</td>
<td>---------------</td>
<td>-------------------------------------</td>
</tr>
<tr>
<td>MINI VIDAS</td>
<td>1.188</td>
<td>-0.132</td>
<td>r=0.9873</td>
<td>Passing-Bablok</td>
<td>39</td>
<td>serum</td>
<td>One freeze-thaw cycle (-20°C), multicenter, all analyzed on same day</td>
</tr>
<tr>
<td><strong>Diazyme procalcitonin methods</strong> (Diazyme PCT reagent used on the analyzer indicated)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ADVIA 2400</td>
<td>1.01</td>
<td>-0.23</td>
<td>r=0.927</td>
<td>Deming</td>
<td>100</td>
<td>serum</td>
<td>One freeze-thaw cycle (-70°C), multicenter, all analyzed on same day</td>
</tr>
<tr>
<td>ARCHITECT c16000</td>
<td>0.95</td>
<td>-0.16</td>
<td>r=0.899</td>
<td>Deming</td>
<td>100</td>
<td>serum</td>
<td>One freeze-thaw cycle (-70°C), multicenter, all analyzed on same day</td>
</tr>
<tr>
<td>AU5800</td>
<td>0.87</td>
<td>-0.04</td>
<td>r=0.899</td>
<td>Deming</td>
<td>100</td>
<td>serum</td>
<td>One freeze-thaw cycle (-70°C), multicenter, all analyzed on same day</td>
</tr>
<tr>
<td>AU5800</td>
<td>1.10</td>
<td>-0.89</td>
<td>r=0.960</td>
<td>Deming</td>
<td>129</td>
<td>serum</td>
<td>Samples analyzed fresh (within 3 h from collection), methods run simultaneously</td>
</tr>
<tr>
<td>cobas c 501</td>
<td>0.85</td>
<td>0.07</td>
<td>r=0.899</td>
<td>Deming</td>
<td>100</td>
<td>serum</td>
<td>One freeze-thaw cycle (-</td>
</tr>
</tbody>
</table>
70°C), multicenter, all analyzed on same day

<table>
<thead>
<tr>
<th>Assay</th>
<th>Lab 1</th>
<th>Lab 2</th>
<th>Study Method</th>
<th>Sample</th>
<th>Storage</th>
<th>Time</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>cobas c 702</td>
<td>0.6543</td>
<td>0.014</td>
<td>Passing-Bablok</td>
<td>239</td>
<td>serum</td>
<td>Analyzed within 24 h stored at 2-8°C</td>
<td>Ceriotti et al., 2017(47)</td>
</tr>
<tr>
<td>cobas c 702</td>
<td>1.19</td>
<td>0.01</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
<td>Lippi et al., 2019(43)</td>
</tr>
</tbody>
</table>

Other procalcitonin assays

<table>
<thead>
<tr>
<th>Assay</th>
<th>Lab 1</th>
<th>Lab 2</th>
<th>Study Method</th>
<th>Sample</th>
<th>Storage</th>
<th>Time</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Access (Access)</td>
<td>1.12</td>
<td>-0.03</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
<td>Lippi et al., 2019(43)</td>
</tr>
<tr>
<td>Access (DXI)</td>
<td>1.19</td>
<td>-0.07</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
<td>Lippi et al., 2019(43)</td>
</tr>
<tr>
<td>MAGLUMI</td>
<td>1.51</td>
<td>-0.10</td>
<td>Passing-Bablok</td>
<td>176</td>
<td>lithium-heparin plasma</td>
<td>One freeze-thaw cycle (-70°C), multicenter, analyzed on same day</td>
<td>Lippi et al., 2019(43)</td>
</tr>
</tbody>
</table>

*This study used a combination of both BRAHMS PCT sensitive Kryptor and Elecsys BRAHMS PCT as the reference methods and did not distinguish between them.*
**Supplementary Table References:**


