Early Diagnostic Markers for Neonatal Sepsis - Haematological Scoring System, C-Reactive Protein and Procalcitonin

Dr Malvika Gaur¹, Dr Arathi C.A.²

Abstract: Background: Neonatal sepsis is associated with high mortality and morbidity rate, as the clinical manifestations are nonspecific. Therefore, the need arises for early diagnostic markers of neonatal sepsis like Haematological scoring system (HSS), C-Reactive protein (CRP), Procalcitonin (PCT). Aim: To evaluate the role of diagnostic parameters – HSS, CRP and PCT in the early detection of neonatal sepsis. Methods: 50 cases of neonatal sepsis were studied, HSS was calculated as per the 7 point Rodwell et al scoring system. PCT was calculated by the semi-quantitative kit. Statistical analysis: The results were compared with each other (comparative study design), with the gold standard (blood culture). Specificity, sensitivity, PPV & NPV were calculated. Result: Blood culture was Positive - 46 % cases, negative - 54% cases, most common isolate obtained Coagulase negative staphylococcus aureus (CONS) 47.3% cases, Staphylococcus aureus (34.78% cases). HSS obtained Sepsis very likely - 56% cases, Sepsis possible - 40% cases, Sepsis unlikely - 4% cases. CRP results obtained positive results - 60% cases, negative - 40% cases. PCT results were positive - 86% cases and negative - 14% cases. Conclusion: Among the various screening tests, PCT has higher sensitivity of 91.3% as compared to CRP (73.9%), HSS (65.2%). Thus, PCT is a better marker for screening neonatal sepsis. However it is not very specific, can be raised in various other conditions.

Keywords: Blood culture, HSS, Neonatal sepsis, PCT

1. Introduction

Neonatal sepsis is associated with high mortality and morbidity rate, as the clinical manifestations are nonspecific.¹ therefore, it is very essential to diagnose the sepsis in early phase and also it is equally important to rule out neonatal sepsis.² Therefore, the need arises for early diagnostic markers of neonatal sepsis like Haematological scoring system (HSS), C-Reactive protein (CRP) and Procalcitonin (PCT). An early diagnosis using a sensitive marker can reduce the mortality and improve the outcome. According to the National Neonatal Perinatal Database (NNPD) report 2002-2003, the incidence of neonatal septicemia in tertiary care institutions has been reported to be 14.5 per 1000 live births (2.3%) and contributes to 16% morbidity rate, as the clinical manifestations are nonspecific. Therefore, it is very essential to diagnose the sepsis in early phase and also it is equally important to rule out neonatal sepsis.² Therefore, the need arises for early diagnostic markers of neonatal sepsis like Haematological scoring system (HSS), C-Reactive protein (CRP) and Procalcitonin (PCT). An early diagnosis using a sensitive marker can reduce the mortality and improve the outcome. According to the National Neonatal Perinatal Database (NNPD) report 2002-2003, the incidence of neonatal septicemia in tertiary care institutions has been reported to be 14.5 per 1000 live births (2.3%) and contributes to 16% of all mortalities among the hospital born neonates. (National Neonatal Perinatal Database: Report for the year 2002-03. National Neonatology Forum, India).³

1.1 Aim

To evaluate the role of diagnostic parameters – HSS, CRP and PCT in the early detection of neonatal sepsis.

1.2 Objective

1) To evaluate the role of HSS, CRP and PCT in the early detection of neonatal sepsis.
2) To compare between HSS, CRP and PCT, as a better marker in the early detection of neonatal sepsis.
3) To derive the sensitivity and specificity of PCT as an early diagnostic marker.

2. Materials and Methods

In the present study 50 cases of clinically suspicious cases of neonatal sepsis were studied.

Inclusion criteria: Clinically suspected cases of neonatal sepsis admitted in the neonatal intensive care (NICU) and neonates who developed signs and symptoms of sepsis while they were admitted in NICU.

Exclusion criteria: 1) Suspected cases of septicaemia, where antibiotics have already been administered
2) Inborn errors of metabolism
3) Congenital anomalies

An approval from the ethical committee was obtained and ethical practices were observed during the study. Written valid consent was taken from the parents. A detailed clinical history and findings were recorded. The blood samples were collected and processed for HSS, CRP and PCT. Blood culture – 1 ml of blood was drawn aseptically and inoculated into blood culture bottles containing 10 ml of brain heart infusion broth.

HSS was calculated as per the 7 point Rodwell et al scoring system in the present study.⁴

<table>
<thead>
<tr>
<th>SN</th>
<th>Criteria</th>
<th>Abnormality</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Total WBC count</td>
<td>≤5,000/µl</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>Total PMN count</td>
<td>≤1,800/µl &amp; ≥5,400/µl (at birth)</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>Immature PMN count</td>
<td>≤600/µl</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>I:T PMN ratio</td>
<td>≤ 0.3</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>I:M PMN ratio</td>
<td>≤ 0.3</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>Degenerative PMN count</td>
<td>≥ 0.3</td>
<td>1</td>
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<tr>
<td></td>
<td>Toxic granules (picture 1)</td>
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Abnormal total PMN count is assigned score of 2, instead of 1, if no mature polymorphs are seen on peripheral smear examination to compensate for low immature to mature ratio. Thus a score of 0 to 8 was obtained and interpreted as – score ≤2 - Sepsis unlikely, 3-4 - Sepsis possible, ≥5 - Sepsis very likely.4

CRP was assessed using the rapid slide latex agglutination qualitative method (LAB-CARE DIAGNOSTICS (INDIA) PVT LTD). PCT was assessed using B.R.A.H.M.S PCT-Q KIT (Picture 3) manufactured by Thermo SCIENTIFIC, an immunochromatographic test for the semi-quantitative detection of PCT concentrations in serum/plasma. PCT level of <0.5 ng/ml - Sepsis not likely, ≥0.5 ng/ml to <2ng/ml - Sepsis possible, ≥2 ng/ml to 10 ng/ml - Sepsis likely and ≥10 ng/ml - Sepsis very likely.

Statistical analysis : The results were compared with each other and with the gold standard (blood culture). Specificity, sensitivity, PPV & NPV were then calculated.

3. Result

Blood culture was Positive in 46 % cases and negative in 54% cases. Blood culture most common isolate obtained Coagulase negative staphylococcus aureus (CONS) 47.3% cases and Staphylococcus aureus (34.78% cases) (Picture 4.5).

HSS obtained Sepsis very likely (56% cases), Sepsis possible (40% cases) and Sepsis unlikely (4% cases). CRP results obtained positive results in 60% cases and negative in 40% cases. PCT results were positive in 86% cases and negative in 14% cases.

Table 1: Correlation of HSS with gold standard (blood culture)

<table>
<thead>
<tr>
<th>HSS</th>
<th>Blood culture</th>
<th>Total</th>
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<tbody>
<tr>
<td></td>
<td>Positive</td>
<td>Negative</td>
</tr>
<tr>
<td>Positive</td>
<td>15</td>
<td>13</td>
</tr>
<tr>
<td>Negative</td>
<td>8</td>
<td>14</td>
</tr>
<tr>
<td>Total</td>
<td>23</td>
<td>27</td>
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</table>

Table 2: Correlation of CRP with gold standard (blood culture)

<table>
<thead>
<tr>
<th>CRP</th>
<th>Blood culture</th>
<th>Total</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Positive</td>
<td>Negative</td>
</tr>
<tr>
<td>Positive</td>
<td>17</td>
<td>15</td>
</tr>
<tr>
<td>Negative</td>
<td>6</td>
<td>14</td>
</tr>
<tr>
<td>Total</td>
<td>23</td>
<td>27</td>
</tr>
</tbody>
</table>

Table 3: Correlation of PCT with gold standard (blood culture)

<table>
<thead>
<tr>
<th>PCT</th>
<th>Blood culture</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Positive</td>
<td>Negative</td>
</tr>
<tr>
<td>Positive</td>
<td>21</td>
<td>22</td>
</tr>
<tr>
<td>Negative</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
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</tbody>
</table>

Among the various screening tests, PCT has a higher sensitivity of 91.3% as compared to CRP (73.9%) and HSS (65.2%). Thus, PCT is a better marker for screening neonatal sepsis. However it is not very specific and can be raised in various other conditions. 94% cases were healthy when discharged, 2% were discharged against medical advice and 4% died.

It was noted that neonatal risk factors associated with sepsis were low birth weight (<2.5 kg) (78% cases), preterm (<37 weeks of gestational age) (80% cases), lower socioeconomic status (82% cases) and born to multigravida mothers (72% cases). Early onset neonatal sepsis (presented within 7 days of birth) was seen in (78% cases). Respiratory distress syndrome was seen in 51.28% cases and Meconium aspiration syndrome in 20.51% cases. Maternal risk factors associated with neonatal sepsis were Anaemia during pregnancy (32.69% cases), Pregnancy induced hypertension (30.77% cases) and Premature rupture of membranes (26.92% cases).

4. Discussion

Blood culture is gold standard for definitive diagnosis of neonatal sepsis, but it has its own limitations. The yield of a positive blood culture ranges from 8-73%. A negative blood culture does not exclude sepsis and about 26% of all neonatal sepsis could be due to anaerobes. Maternal antibiotics given in preterm deliveries may suppress the growth of bacteria in culture.

HSS is a simple cost, quick and effective tool in the early diagnosis of neonatal sepsis, but its sensitivity is unsatisfactory. Therefore it cannot provide a guideline to decisions regarding antibiotic therapy. It can be applied to even those neonates who have received antibiotic therapy. WBC varied widely across studies, with sensitivity & specificity ranging from 17% to 90% and 31% to 100%. WBC maybe helpful in diagnosing sepsis, however normal WBC counts maybe observed in as many as 50% of culture proven sepsis cases, and neonates who are not affected may also have abnormally high WBC counts as a result of the stress of delivery.

The associated band count and a leftward shift of the myeloid immaturity measurements may improve the diagnostic yield, but their subjective measurement is problematic. Neutrophilia itself is not a reliable or sensitive indicator of infection. Thrombocytopenia was frequently associated with sepsis and indicated poor prognosis. It is an important parameter in supporting the diagnosis of sepsis, although it appears to be a late finding and nonspecific. 21 cases of low platelet count were seen in the present study.

Table 4: Sensitivity, specificity, NPV and PPV of PCT, CRP and HSS

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>PPV</th>
<th>NPV</th>
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<tbody>
<tr>
<td>PCT</td>
<td>91.3%</td>
<td>18.5%</td>
<td>48.8%</td>
<td>71.4%</td>
</tr>
<tr>
<td>CRP</td>
<td>73.9%</td>
<td>51.9%</td>
<td>56.7%</td>
<td>70.0%</td>
</tr>
<tr>
<td>HSS</td>
<td>65.2%</td>
<td>51.9%</td>
<td>53.6%</td>
<td>63.3%</td>
</tr>
</tbody>
</table>

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CRP is commonly used for detection of sepsis in neonates, but it is not useful as an early phase infection marker and it lacks specificity. Long duration between invasion by infectious agent and the rise in serum CRP concentrations. CRP can be considered as a specific but late marker of neonatal infections.

PCT rapidly increases in 6 to 8 hours, and then a plateau in 12 to 48 hours. Plasma elimination is approx 25 to 30 hours. Therefore it has value in early detection of neonatal sepsis, and shows quick reduction in its level post antibiotic therapy. PCT is more sensitive than CRP in the diagnosis of septicaemia, meningitis and UTI. PCT used together with CRP, a negative PCT test may help in “ruling out” while a raised CRP result helps in “ruling in”, the possibility of sepsis, particularly of the late onset type. The early response to appropriate antibiotic therapy can be evaluated by PCT in the septic neonates, but not by CRP. Late response to treatment can be evaluated by both CRP and PCT. It is not the sole marker of neonatal sepsis, and is relatively expensive.

This study correlated with a study by Sucilathangam G in 2012 studied PCT and CRP in neonates admitted to NICU. They concluded that PCT was more sensitive than CRP in the detection of neonatal sepsis. A negative PCT test result may help to “rule out”, while a raised CRP result helps to “rule in”, the possibility of sepsis. Monsef A and Eghbalian F in 2012 concluded that PCT has a high sensitivity, specificity, PPV and NPV for the diagnosis of neonatal sepsis. Mamdouh M. Esmat in 2012 concluded that the serum levels of PCT is more reliable marker than the serum levels of CRP in the early diagnosis of neonatal sepsis and in the evaluation of the response of the disease to the antibiotic therapy. H Altunhun in 2011 studied PCT, CRP and blood cultures of neonates admitted to NICU for neonatal sepsis and concluded that PCT measurement at birth may initially be normal, a serial PCT measurement at 24 hours of age may be more helpful for an early diagnosis. During the first 24 hours of life PCT is a more sensitive marker of infection that CRP. Ibeh Isaiah Nnanna in 2011 concluded that PCT monitoring could be helpful in the early diagnosis of neonatal septicemic infection in the intensive care unit. Both absolute values and variations should be considered and evaluated in further studies. A combination of 3 or all of 4 tests was highly specific 95%–100%.

Limitations in this study were - small sample size, lack of follow up repeat PCT levels and blood culture negative results, even in clinically proven sepsis, this reduced the sensitivity and specificity of CRP and PCT.

5. Conclusion

HSS and CRP are simple, quick and cost effective tool in the early diagnosis of neonatal sepsis, but its sensitivity in detection of neonatal sepsis is unsatisfactory. PCT had a higher sensitivity (91.3%) as compared to CRP & HSS. PCT is a better marker for screening neonatal sepsis. However is not very specific and can be raised in various other conditions.

6. Acknowledgements

1. Dr Arathi CA – my guide for her constant support and guidance. She is MD Pathology, working in Sri Siddhartha medical college, Tumkur during the study.

2. Dr Sharda Devi MY, Department of microbiology, Shri Siddhartha medical college, Tumkur, Karnataka for carrying out the blood cultures and providing reports and photography, and cooperation.

Funding: Dr Malvika Gaur

Competing Interests: NA

References


Table legends

Tables are already added in the article
Table 1 : Correlation of HSS with gold standard (blood culture)
Table 2 : Correlation of CRP with gold standard (blood culture)
Table 3 : Correlation of PCT with gold standard (blood culture)
Table 4 : Sensitivity, specificity, NPV and PPV of PCT, CRP and HSS.

Picture legends

Picture 1 – Peripheral smear showing band forms and toxic granules (Leishman stain, 100 x)
Picture 2 – Peripheral smear showing cytoplasmic vacuolations (Leishman stain, 100 x)
Picture 3 – Commercially available PCT – Q Kit : positive PCT (≥10 ng/ml)
Picture 4 – MacConkey agar showing growth for Staphylococcus aureus
Picture 5: Blood agar showing growth for Coagulase negative Staphylococcus aureus