Role of CRP in the Evaluation and Diagnostic use in Assessment of Neonatal Sepsis in Tertiary care Hospital

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Abstract

Introduction: Neonatal sepsis, a clinical syndrome of bacteremia with systemic signs and symptoms of infection in the first 4 weeks of life is a major cause of morbidity and mortality in newborn. Early diagnosis is critical, as sepsis can progress more rapidly in neonates than in adults. In a developing country like Nepal, where proper medical care and obstetrical facilities are still scarce in the rural areas. With limited resources, early diagnosis and treatment if crucial, may not be achieved. C-reactive protein (CRP) is the most common diagnostic marker of infection. The aims of this study is to evaluate the role of CRP in the diagnosis of neonatal sepsis. Material and Methods: This prospective observational study were done in the department of paediatrics, UCMS-TH, Bhairahawa from May 2019 to March 2020. Clinically suspected neonatal sepsis cases were enrolled in the study. Venous blood were collected before giving any intravenous fluid or antibiotics for complete blood counts, CRP levels, blood sugar and blood culture and send to laboratory within half hour of collection. All patients included in this study were treated accordingly and followed up strictly. Results: A total of 440 patients clinically diagnosed as neonatal sepsis were studied. The mean age was found 10.1±8.5 days with range from 1 to 28 days and more than half (51.6%) babies belonged to age ≤7 days. More than two third 71.4% babies were male and 28.6% were female. 236(53.6%) babies were found CRP positive and 204(46.4%) were CRP negative. 25.5% babies were blood culture positive and 74.5% babies were culture negative.364 (82.73%) babies were cured and 76 (17.27%) were died. Conclusion: CRP is a simple, relatively cheap and sensitive study in prediction of neonatal sepsis. It can be useful test in settings with limited resources. : Serum CRP is the most sensitive marker of sepsis.

Keywords: C-reactive protein, Antibiotic, Neonate, Sepsis.

INTRODUCTION

Sepsis is the commonest cause of neonatal mortality; it is responsible for about 30-50% of the total neonatal deaths in developing countries [1]. It is estimated that up to 20% of neonates develop sepsis and approximately 1% die of sepsis related causes [2]. All neonates suspected to have sepsis should have a septic screen to corroborate the diagnosis. However, the decision to start antibiotics need not be conditional to the sepsis screen result, if there is a strong clinical suspicion of sepsis. The clinical diagnosis of neonatal sepsis is difficult because the signs and symptoms are not always specific. There is no laboratory test with 100% sensitivity and specificity [3]. Blood culture has been considered the gold standard for confirmation of diagnosis but the results are available only after 48-72 hours. Moreover, in many cases blood culture fails to detect the offending organism/bacteria. So, the search for a reliable test continues, especially one that is useful in culture-negative cases [4]. It encompasses various systemic infections of the newborn such as septicaemia, meningitis, pneumonia, arthritis, osteomyelitis and urinary tract infections [5]. Neonatal mortality rate of Nepal is 33 per 1,000 live births and sepsis is one of the leading cause. In B.S year 2070/71, of the total neonates presenting to government health facilities, 13.9% had possible bacterial infection and 42.1% had local bacterial infection. The major causes of neonatal deaths in Nepal are infection, birth asphyxia, preterm birth, and hypothermia [6].

Despite advances in maternal and neonatal care, infection remains a frequent and important cause of neonatal and infant mortality and morbidity [7]. Neonatal mortality is associated with about 41% of all death among under-five children [8]. Newborn infants, especially sick or preterm infants, are at risk of developing severe infections (such as bloodstream
Infections) during their stay on neonatal units. Infections are often difficult to diagnose early with certainty, and quick tests such as measuring the blood level of a protein that responds to infection (called C-reactive protein) are sometimes used to help make an earlier diagnosis. Neonatal sepsis evaluation, a common occurrence in the Neonatal Intensive Care Unit (NICU), is usually comprised a complete blood count (CBC), blood culture and possibly cerebrospinal fluid (CSF), and urine cultures. When an infant appears ill the clinical findings such as temperature instability, respiratory distress, apnea, bradycardia, and hypotension are nonspecific [10, 11]. Sometimes, especially if noted in isolation, these findings are transient, and no infection is confirmed. If an organism is isolated from a sterile site such as blood or CSF, the treatment course is straightforward. For the majority of remaining infants, the nonspecific symptoms resolve at some point after sepsis evaluation, but antibiotics have already been started. Often an organism is not isolated from the various specimens collected, bringing uncertainty of when to discontinue antibiotic treatment. The use of biomarkers, such as C-reactive protein (CRP), can assist in sepsis management decision-making [12]. The use of rapid blood testing for CRP has become common in pediatrics to help evaluate infection in children and increasingly more common during the evaluation of sepsis in neonates [13].

CLASSIFICATION OF NEONATAL SEPSIS:
Neonatal sepsis can be classified into two major categories depending up on the onset of symptoms, EARLY ONSET SEPSIS (EOS): It presents within first 72 hours of life. In severe cases, the neonate may be symptomatic at birth. Infants with EOS usually present with respiratory distress and pneumonia. The source of infection is generally the maternal genital tract. LATE ONSET SEPSIS (LOS): It usually presents after 72 hours of age. The source of infection in LOS is either nosocomial (hospital acquired) or community acquired and neonates usually present with septicemia, pneumonia or meningitis.

MATERIAL AND METHODS
It is a prospective observational study, conducted during May 2019 to March 2020 in the Department of Paediatrics, UCMS. A detailed history and thorough physical examination were done in each neonate on admission. History included age of newborn, sex, gestational age, I/o prolonged rupture of membrane (PROM), intrapartum fever or fever 3 days before delivery, per vaginal foul smelling discharge, prolonged labour and features of sepsis. Physical examination included respiratory rate, heart rate, temperature, chest indrawing, grunting, cyanosis, convulsion, breath sound, added sound, weight, jaundice, bleeding manifestation, status of fontanelles, umbilicus and capillary refill time. Venous blood was collected before giving any intravenous fluid or antibiotics for blood sugar, complete blood counts, CRP levels and blood culture and sensitivity. Sample was sent to laboratory with in half hour of collection. CRP levels > 6mg/L considered as positive and < 6mg/L considered as negative. All patients included in this study were treated accordingly and followed up strictly. The outcome and relevant data from history, physical examination and investigations were recorded in predesigned questionnaire.

Inclusion Criteria
Neonates were enrolled on the basis of signs and symptoms of clinical sepsis after through clinical examination and proper history taking. The clinical criteria considered were – poor feeding, irritability / excessive cry, lethargy poor cry and reflexes, fever, hypothermia, jaundice, vomiting, abdominal distension, tachypnoea and grunting, convulsions, diarrhea, pusules, cyanosis, bulged fontanelle, bleeding, poor perfusion / shock, apnea. Also significant predisposing factors for presumed early onset sepsis was taken into consideration during inclusion of cases.

Exclusion Criteria
Neonates who received antibiotics before Admission, Neonates who died before work up was complete, Neonates who underwent surgery, Congenital anomalies e.g. tracheoesophageal fistula, lobar agenesis, malrotation of the gut, complex heart diseases, neural tube defects etc. Inborn errors of metabolism.

Investigations Done – Sepsis Screening
- Absolute Neutrophil Count (ANC) :< 1800/cu mm
- Immature to Total Neutrophil (I/T) RATIO :> 0.2(immature neutrophils/ANC), highlysensitive of NNS,
- Total Leukocyte Count (TLC) :< 5000/cu mmr>20,000/cu mm
- CRP :> 6mg/dL
- Micro ESR (µ-ESR): >15 mm/1” hr, specifictbut moderate sensitivity
- Platelets<150000/cu mm

Infections
- Blood Culture

Thenumber of neonates with positive blood culture was recorded and analyzed. The blood culture positivity was considered when both gram stain and culture was positive. All the positive cultures were sent to the department of microbiology for a detailed report including the causative pathogen and the antibiotic sensitivity test.

Ethical Clearance
- The approval of Institutional Review Committee of Universal College of Medical Sciences, Bhairahawa, Nepal was taken before the initiation of experiment. Registration No. UCMS/IRC/117/19. All the protocols and experiments were conducted in compliance with the ethical principles and guidelines.

Statistical Analysis
- Data were processed manually and analyzed with the help of SPSS (Statistical package for social

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RESULTS

A total of 440 patients clinically diagnosed as neonatal sepsis were studied. The mean age was found 10.1±8.5 days with range from 1 to 28 days and more than half (51.6%) babies belonged to age ≤7 days. More than two third (71.4%) babies were male and 28.6% were female. 236(53.6%) babies were CRP positive and 204(46.4%) were negative CRP. 25.5% babies were blood culture positive and 74.5% babies were culture negative.

Table 1 showed 282 (64.1%) Hindu, 110(25.0%) Muslim, and others were 48(10.9%). 314(71.4%) male and 126(28.6%) female neonate with male to female ratio of 2.5:1. However 232(52.7%) were delivered by normal vaginal delivery and 208(47.3%) were delivered by caesarian section. maternal risk factor for sepsis were 132(30.0%) and without risk factor for sepsis 308(70.0%). PV leakage were present in 52(11.8%) cases. 82(18.6%) neonate were born low birth weight and 358(81.4%) were normal weight (>2.5kg). 82(18.6%) were preterm, 342(77.7%) were term baby and 16(3.6%) cases were post-term baby.

Among the patients with suspected neonatal sepsis, the most common presenting clinical feature was respiratory distress followed by feeding problems and lethargic.

Table-3: CRP valve in suspected neonatal sepsis

| CRP    | No. of patients | No (%) |
|--------|----------------|--------|
| Positive | 236            | 53.6%  |
| Negative | 204            | 46.4%  |
| Total    | 440            | 100%   |

Out of total 440 suspected neonatal sepsis, 236(53.6%) were CRP positive and 204(46.4%) were CRP negative.

Table-4: Distribution of isolated organism

| Organisms            | Number | Percentage |
|----------------------|--------|------------|
| Coagulase negative   | 44     | 43.13%     |
| Klebsiella           | 24     | 23.52%     |
| Actinobacter species | 12     | 11.76%     |
| Staphylocus epidermis| 4      | 3.92%      |
| Enterococcus species | 4      | 3.92%      |
| MR staphylococci species | 12 | 11.76% |
| Grampositivecoccus CONS | 2     | 1.96%      |
| Total                | 102    | 100.0%     |
Of the 440 cases were evaluated for sepsis, the blood culture was positive in 102 (24.9%) cases. In which Coagulase negative Staphylococci 44 (43.3%) and Klebsiella species 24 (23.52%) were the most common Gram positive and Gram negative organisms.

Table-5: Relationship between CRP and glucose level

| CRP     | <45 | 45-145 | >145 | Total |
|---------|-----|--------|------|-------|
| Positive| 204 | 236    | 236  | 676   |
| Negative| 204 | 236    | 236  | 676   |
| Total   | 408 | 472    | 472  | 1352  |

Majority (51.7%) of CRP positive patients were found normoglycemic, 33.9% were found hypoglycemic and only 14.4% were found hyperglycemic.

**DISCUSSION**

Out of 440 suspected neonatal sepsis patients, 314 (71.4%) were males and 126 (28.6%) were females. Among the patients with suspected neonatal sepsis, the most common presenting clinical feature was respiratory distress followed by feeding problems and lethargic. In this study, Coagulase negative staphylococcus (CONS) (43.13%), Klebsiella species (23.52%), Acinetobacter species (11.76%), Pseudomonas aeruginosa (9.32%), Enterococcus species (3.2%) and Methacillin resistant Staphylococcus aureus (11.76%) were the common organisms associated with sepsis. In a similar study done in India most prevalent organisms were coagulase negative staphylococcus (CONS) followed by Coagulase positive staphylococcus, streptococcus faecalis, alpha- hemolytic Streptococcus, klebsiella, proteus, E.coli and Candida albicans [14]. The causative organisms in neonatal sepsis vary from place to place and the frequency of the causative organisms is different in different hospitals and even in the same hospital at different time. The other published data in Nepal on the subject shows E.coli as the most common isolate [15, 16].

Definitive diagnosis rests upon a positive blood culture, to identify the pathogen and determine its antibiotic susceptibility pattern, but for better survival and outcome, simple and rapid diagnostics tests are required as adjuncts to the blood culture for early and effective initiation of treatment to the septicemia in neonates.

Majority (51.7%) of CRP positive patients were found normoglycemic, 33.9% were found hypoglycemic and only 14.4% were found hyperglycemic. It was observed that 82.73% patients were cured and 17.27% were died.

C-reactive protein ≥6mg/dl was considered as positive result for sepsis screen. Predictive accuracy of CRP of this study is compared with other studies. In present study, CRP positive 236 (53.6%) and negative 204 (46.4) were found in suspected sepsis. CRP proved to be the most efficient of all the markers of sepsis. The principal ligand to CRP is phosphocholine, which is found in lipopolysaccharide, bacterial cell walls, as well as in most biological membranes [17]. CRP is part of the acute-phase response which aims to neutralize the inflammatory agent and to promote the healing of the injured tissue [18].

Thus male babies were more affected by suspected neonatal sepsis than female babies. In the part to biological difference it is proposed to be due to genetic origin for the origin of the sex different in vulnerability to infection. The special source of vulnerability open to females by virtue of her possession of to X-chromosomes in contrast to the single ‘X’ of the male. Hence it is genetic locus on the ‘X’ Chromosomes involve with synthesis of immunoglobulins responsible for the sex difference as per Thomas C. Washburn et al., [19].

Bacterial infection stimulates the hepatocytes to produce CRP: a nonspecific immune response, which is a useful clinical marker for the individual host-pathogen interaction. Since the half-life of CRP is less than 3 days, a rapid fall is seen with successful therapy. Early diagnosis of neonatal sepsis is very difficult when it is based only on clinical signs. The clinical profile is neither uniform nor specific and could mislead the health care professionals. Keeping in mind the mortality caused by neonatal sepsis, empirical treatment should not be delayed but again this can result in unnecessary and prolonged exposure to antibiotics in this early age group. Cost effective and rapid diagnostic tool is one way to tackle this issue. CRP and leukocyte count should be considered to decrease neonatal mortality rate in the country.

**CONCLUSION**

CRP is a simple cheap and relatively sensitive, specific with good negative predictive value in the prediction of neonatal sepsis and can be useful test in the settings with limited resources.

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