A RARE PRESENTATION OF HIV IN NEWBORN - OSTEOMYELITIS

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ABSTRACT

BACKGROUND

The Human Immunodeficiency Virus (HIV) is a lentivirus (retrovirus) that causes HIV infection and over time Acquired Immunodeficiency Syndrome (AIDS). AIDS is a condition in humans in which there is progressive failure of immune system, which allows life-threatening opportunistic infections and cancers to thrive. Globally, an estimated 43 million people are infected with HIV. Worldwide, of people living with HIV, the percentage of women with HIV remains at 50%. With maternal diagnostic and prophylaxis during perinatal period, perinatal HIV transmission can be prevented. For this HIV testing and antiretroviral (ARV) prophylaxis and treatment are essential. In our case report, we present a case of preterm baby with HIV seropositivity that was vertically transmitted and accompanied by osteomyelitis. We report this case to emphasise the importance of early detection and early ARV chemotherapy to prevent vertical transmission.

KEYWORDS

Neonatal Osteomyelitis, ARV Chemotherapy, Vertically Transmitted.


BACKGROUND

Neonatal osteomyelitis in HIV positive neonates is a rare and challenging diagnosis, particularly in early onset period. Osteoarticular infections, although uncommon, represent a severe condition in HIV positive neonates. Neonatal osteomyelitis is predominantly caused by staphylococcus aureus with single bone lesion. Lower extremity joints are commonly affected. Early diagnosis and treatment to reduce the sequelae is very important. Diagnosis involves appropriate radiographic studies and culture of blood and material obtained from infected bone by needle aspirate. Treatment requires prolonged intravenous therapy with antibiotics that achieve high bone penetration and are directed against the identified or presumed causative bacteria and Antiretroviral (ARV) chemotherapy. Here, we report a case of neonatal osteomyelitis in a preterm baby with HIV seropositivity that was vertically transmitted.

CASE REPORT

A 32-week-old, male baby with an Apgar score of 8/9 was delivered normally by a 28-year-old unbooked primigravida who presented to obstetric emergency in labour. No antenatal care and investigations were done before. Baby cried immediately after birth, weight was 1.504 gms, vitals were normal, systemic examination unremarkable, no obvious congenital malformation was noted. The neonate was admitted in NICU for preterm care and low birth weight.

At 29 hours of life, the baby developed respiratory distress. Sepsis screen was positive (Micro ESR = 8 mm at 1 hour, CRP = 50 mg/L) Hb = 16.4 g/dL, TLC = 16000/mm3, platelets = 200000/mm3. Chest x-ray was done, which showed bilateral infiltrates. Early onset neonatal sepsis was diagnosed and IV amoxicillin/clavulanic, amikacin, 02 therapy and IV fluids were given. HIV screening of parents were done. Both mother and father were tested positive for HIV-1. Mother CD4 count was 180/mm3. Screening for hepatitis, syphilis and TORCH was negative. Both were referred to ART Centre for HAART.

Baby developed DIC at 94 hrs. of life which manifested as gastrointestinal bleed, (PT/PTT - prolonged, D-dimer = positive, platelet = 40,000/mm3), for which treatment was given (vitamin K injection, FFP, platelet transfusion, total parenteral nutrition – amino acid, intralipid and MVI). Later on at 120 hrs. of life baby developed acute non-alcoholic renal failure (blood urea = 87 mg/dL, serum creatinine = 1.7 mg/dL), which was managed conservatively. Amino acid concentration was reduced and antibiotics was changed to piperacillin-tazobactam and cefotaxime. Investigations were repeated on day 6. Total leucocyte count = 11000/mm3 with CD4 count of 25%. Sepsis screen was still positive. Blood culture sent on day 1 and day 6 was sterile. Apparent improvement was noted after 2 weeks. Activity was better, tolerated feeds well, renal profile normalised. Antibiotics were stopped.

On day 20, swelling of both thighs (left > right) with decreased and painful limb movement of both legs were noted. X-ray revealed gross osteomyelitis with massive periosteal reaction involving both femurs and left tibia. Ultrasonography of limbs revealed marked subcutaneous cellulitis with thickening and increase echogenicity of subcutaneous fat. Irregular cortical bone surface was present along with periosteal reaction. Doppler examination of both lower limbs was normal. Ceftriaxone sodium was started. Blood culture sent on day 22 revealed growth of staphylococcus aureus sensitive to vancomycin, clindamycin and trimethoprim.
sulfamethoxazole. IV vancomycin was started and continued for 3 weeks.

Baby responded well to the treatment and condition of baby improved during hospital stay, which was evident by decrease in the swelling and improvement of movements. Baby was discharged on 50th day. DNA PCR done on day 40 and repeated on day 43 were positive for HIV-1, CD4 count was 25%, classified as HIV clinical disease stage 3 with advanced immunosuppression [WHO 2007]. Antiretroviral therapy (HAART; zidovudine + lamivudine + nevirapine) was started on day 43 along with Pneumocystic jiroveci pneumonia prophylaxis with cotrimoxazole.

On follow up - last follow up was at 15 months, growth and development of baby was satisfactory. There was no limb length discrepancy or angular deformity and the child was able to bear full weight without support.

**DISCUSSION**

HIV infection is a serious condition with high mortality. HIV cases are increasing due to difficulty in diagnosis, long latent period and routes of transmission. We have reported this case to show the importance of universal HIV screening for all pregnant women, importance of antiretroviral therapy, safe delivery practices and modified infant feeding.

The overall incidence rate for bone and joint infections is 0.12 per 1000 live births and 0.67 per 1000 neonatal intensive care (NICU) admissions(1) with a mortality rate of 7.3%.(2) The most common bacterial pathogen causing osteomyelitis in children is staphylococcus aureus in all age groups.(3) Group B streptococcus (streptococcus agalactiae) and gram negative organism (E. coli and Klebsiella pneumoniae) are also important in neonatal period.(4) The most commonly affected joints are the Hip joints (31%), Knee joints (25%) and Ankle joints (18%).(5) If the mother is infected with HIV, the foetus has a 25% chance of being infected by the virus. Upto 50% of infants get HIV from their mother late in pregnancy or during delivery.

Riordan et al has reported osteomyelitis caused by streptococcus pneumoniae 9S, resistant to penicillin in a 6-month-old girl with HIV by PTCT (Parent To Child Transmission).(6) Yao et al have reported osteomyelitis in 0.9% of a retrospective record review of 888 adult patients with HIV.(7)
Mother To Child Transmission (MTCT) is the most important source of Human Immunodeficiency Virus (HIV) in children below the age of 15 years. The rate of perinatal transmission is 15% - 25% in developed countries and 25% - 45% in developing countries.(8) The rate of perinatal transmission without intervention is 19% - 36%(9); 25% - 30% of total transmission occurs during the prenatal period, mainly in late pregnancy(10) while 70% - 75% of total transmission occurs during the intranatal period. Postnatal transmission is via breast milk and accounts of 10% - 16% of all transmission.(11) In 1994 Paediatric AIDS Clinical Trial Group (PACTG) Protocol, 076 demonstrated that the administration of zidovudine during pregnancy and labour and then to newborn decreased the risk of perinatal transmission of HIV by 68% from 25.5% to 8.3%. (12) Children born to seropositive mother should be followed closely during the first several months of life for evidence of HIV infection, so that optimal care can be provided.

Prevention of HIV infection is one of the priorities of all health strategies, especially those related to maternal and child health. Educational programmes should seek to explain the risk behaviour associated with HIV infection, the implications of HIV infection in pregnancy outcome and the ways in which the risk of HIV infection can be reduced. HIV infection should be identified prior to pregnancy or as early as possible. This provides the best opportunity to prevent infant HIV infection and to identify and start therapy as soon as possible in infants who become infected.

CONCLUSION

Neonatal osteomyelitis in HIV positive infant, although a rare presentation remains a diagnostic and therapeutic challenge. Osteomyelitis should be considered in newborn infants presenting with clinical signs of sepsis without any obvious focus. This is essential for early diagnosis and treatment to prevent any long-term morbidity.

REFERENCES