Prevention of mother to child transmission of hepatitis B virus infection in Nigeria: A call to action

Abstract: Background: Sub-Saharan Africa including Nigeria has the second largest global burden of chronic carriers of hepatitis B virus (HBV) infection after Asia. Mother-to-child transmission (MTCT) of HBV is the most common route of transmission in high endemic areas. MTCT of hepatitis B virus infection continues to occur despite the interventions of hepatitis B vaccinations and immunoglobulins in settings where it is practiced. Infants most at risk are those whose mothers have high HBV DNA viral loads and produce the protein HBeAg. Various Nigerian studies have reported high HBV infection rates as well as HBeAg positivity among pregnant women. These HBV infections usually occur intrapartum and rarely intrauterine. Mothers with HBeAg positivity known to be associated with higher HBV DNA viral loads have been linked with higher chances of MTCT as HBeAg is the only structural HBV protein that can cross into the placental circulation.

In the absence of post exposure prophylaxis about 40 percent of infants delivered by HBV infected mothers could develop HBV infections, and about 25% of them may come down with chronic hepatitis and resulting possible complications including liver cirrhosis and hepatocellular cancer later in life.

The prevention of transmission of retroviral infection from mother-to-child has been a success story of the 21st century and such feat could be replicated for HBV infection. The standard PMTCT of HBV currently will comprise: timely prenatal screening, starting anti-viral therapy for pregnant women with HBeAg positivity and high viral load, infant post-exposure prophylaxis and follow-up of infants of HBsAg positive mothers.

There is no co-ordinated PMTCT of HBV programme in place in our setting despite the huge burden of the disease in Nigeria. Hence the need therefore to develop a home grown PMTCT programme of HBV to help tackle the burden of the disease in our country. An evidence based review of current best practice guidelines for the prevention of mother-to-child transmission of HBV for use in low and medium resource income settings with hepatitis B hyperendemicity will be quiet apt in this circumstance.

This document therefore will be useful as a quick guide to Paediatricians, Obstetricians, Family Physicians, General Practice Doctors and other allied health workers charged with the care of pregnant mothers and their young children.

Methods: Relevant literatures published in English language or translated into English were searched manually and electronically in PUBMED and SCOPUS for the period between 1990 and 2016 on the subject. Keywords searched included: epidemiology of HBV infection, MTCT of HBV, and its preventive strategies including prenatal screening, anti-viral agents in pregnancy, infant post exposure prophylaxis and follow-up of infected children.

Results: Over 35 scholarly articles on HBV epidemiology, MTCT, and preventive measures as well as follow-up models were retrieved and analyzed.

Conclusion: Universal screening of all pregnant women for HBV infection is the most effective
strategy for the prevention of MTCT of HBV, as effective preventive measures could be applied starting from pregnancy to delivery while infants of HBsAg positive mothers should receive timely post exposure prophylaxis and followed up for possible development of chronic hepatitis B infection. Key words: Prevention of mother to child transmission; Hepatitis B virus infection; Resource limited settings.

Introduction

Hepatitis B virus (HBV) infection is a potentially life threatening liver infection. It is a major global public health problem and known to cause chronic hepatitis with high risk of mortality resulting from hepatic cirrhosis and cancer.1 Sub-Saharan Africa has the second largest global burden of chronic carriers of HBV infection after Asian continent.2 Estimates of hepatitis B antigenaemia seroprevalence of 6-20% have been reported, making Sub-Saharan Africa a hyper-endemic region.3,4 Nigeria is also a hyper-endemic country for HBV infection, with varying reported rates ranging from 0.5 to 44.7 percent in children.5-9

In highly endemic regions HBV infection is commonly spread from mother to child prenatally, during the period of delivery (perinatal transmission), or through horizontal transmission (exposure to infected blood/body or other fluids) particularly from an infected child to an uninfected child during the first five years of life.1,10 In Nigeria varying reports of horizontal as well as vertical transmission as the most common route of HBV transmission in children exist.11,12 MTCT has also been shown to occur during pregnancy and/or delivery.13 Mother-to-child transmission (MTCT) of HBV infection has been reported to be responsible for over a third of cases of chronic HBV infections worldwide.14 Hepatitis B infection in pregnancy poses a serious threat to the infant at birth. In the absence of post exposure prophylaxis, about 40 percent of infants delivered by HBV infected mothers could develop HBV infections, and about 25% of whom will die resulting from complications of chronic liver disease.14 Several Nigerian studies in pregnant women have reported HBV rates between 8.3 to 12.8 percent.11,15,16,17

MTCT of hepatitis B virus infection continues to occur despite the interventions of hepatitis B vaccinations and immunoglobulins in settings where it is practiced.14 It is pertinent to understand that the most significant risk factor in transmission of HBV is high maternal viral load.18 Nigerian studies have shown high levels of HBeAg positivity in pregnant women.19,20 Despite presence of hepatitis B surface antigen positivity, mothers who are HBeAg with HBV DNA levels greater than or equal to 10^6 copies per millilitre (greater than 200,000 in/ml) are at greatest risk of transmitting HBV to their infants.21

Screening pregnant women for HBsAg (HBV infection), starting anti-viral therapy for those with HBeAg positivity and high viral load, providing infant post-exposure prophylaxis (PEP) using the administration of hepatitis B vaccine within 24 hours of birth followed by completion of the HBV vaccine series; are recognized strategies for reducing MTCT transmission rates and the global burden of a new chronic HBV infection.10,22

The World Health Organization has actively initiated responses to the control of HBV pandemic through promoting the prevention of its transmission.13

Of note also is that even where women have access to birth dose vaccine of HBV and hepatitis B immunoglobulin there remains a 5-10 percent failure rate. This occurs in women with high HBV viral loads. In this group of mothers, antiviral therapy during pregnancy has shown to significantly reduce the risk of MTCT.23 In circumstances where mothers do not need the anti-viral therapy for their own (HBV infection) health; the antiviral therapy could be used during pregnancy with the aim of reducing the risk of MTCT of HBV.

Considering that the prevention of transmission of retroviral infection from mother-to-child has been a success story in this 21st Century.22 Such feat could be replicated for HBV infection particularly in hyperendemic settings like ours. However gaps still exist on the current recommendations for managing HBV infected mothers as well as their exposed infants particularly in our setting. Also failure in follow-up of HBV exposed infants abound despite the potential complications associated with chronic HBV infections particularly liver cirrhosis and hepatoma.

It therefore becomes pertinent to review the literature on the current best practice guidelines on prevention of mother-to-child transmission (PMTCT) of HBV infection in other to further empower our clinicians particularly Obstetricians and Paediatricians alike as well as other allied health professionals in addressing this problem adequately in order to reduce the burden of HBV through vertical transmissions in high endemic settings like ours.

Nigeria has introduced the birth dose of hepatitis B vaccine as well other primary series hepatitis B vaccination since 2004.9 The birth dose of HBV vaccine has been shown to be strategic towards PMTCT of HBV, however ensuring universal HBsAg screening in pregnancy, starting anti-viral therapy for pregnant women with HBeAg positivity and high viral load, timely administration of the birth dose of HBV vaccination and follow-up of exposed infants will remarkably help to realize this objective of PMTCT of HBV. Also efforts should be made to educate carrier mothers (after testing) and their families on the prevalent risk factors and essence of pre-
ventive measures.

Current implications are that there is urgent need for Nigeria and other countries in the Sub-Saharan Africa as well as Asia where hyper-endemicity of HBV exist to articulate evidence – based, home adaptable and sustainable programmes on prevention of mother-to-child-transmission of HBV geared towards the eradication of the HBV infection.

In Nigeria the current national programme on the PMTCT of human immune deficiency virus (HIV) infection could be leveraged upon in the realization of HBV prevention. All that is needed is the political will, which basically implies political strategic financial investment in the programmes to make the PMTCT of HBV in Nigeria - a dream come true.

Hepatitis B Virus Pathology

The HBV is a member of the hepadnaviridae family (a hepatotropic group of DNA viruses). The mature HBV virion is a spherical double layered “Dane particle” that has 4 different genes including the outer surface envelope of glycoprotein (hepatitis B surface antigen; HBsAg); an inner portion of the virion designated as hepatitis B core antigen (HBcAg), the nucleocapsid which encodes the viral DNA and a non-structural antigen known as hepatitis B epsilon (e) antigen (HBeAg). The HBeAg is a non-particulate soluble antigen from HBcAg by proteolytic self clearance.

The HBeAg is a marker of active viral replication and correlates favourably with HBV DNA levels. The HBV virion also contains a DNA polymerase which exhibits reverse transcriptase activity and genomic replication occurs through an intermediate RNA template as well as protein from the x-region (HBX) which is needed for the viral replication and acts as a transcriptional transactivator of the viral genes and a wide range of host gene parameters.

The replication of HBV occurs usually in the liver; however it could occur in the lymphocytes, spleen, kidneys, and pancreas.

Mother-to-child transmission of hepatitis B virus infection

Prior to the era of the introduction of universal hepatitis B vaccinations the possible risk of a child becoming a chronic carrier of hepatitis B virus infection was about 10 – 30 percent if born to mother who is HBsAg positive but HBeAg negative; and up to 90% if mother is also HBeAg positive.

The most important risk factor for the transmission of hepatitis B in pregnancy is the maternal HBV DNA levels. It has also been proven that failures in MTCT infant post exposure prophylaxis (PEP) usually occur in mothers with HBV DNA viral threshold of ≥10⁶ to ≥10⁹ copies per millilitre. Maternal acquisition of HBV infection particularly in third trimester of pregnancy or intrapartum carries the highest risk of transmission of HBV infection.

The risk of maternal transmission of HBV infection in pregnancy and peri-natal period is dependent on maternal HBeAg positivity. Mothers with positive HBeAg have a transmission rate of 70-90 percent whereas those with a negative HBeAg have a rate probably less than 10%.

However the precise mechanism of HBV transmission remains largely uncertain but it appears that infection may occur intrapartum, or rarely, in-utero. Hepatitis B viral DNA and HBsAg have been detected in amniotic fluid, placental cells and virginal secretions of HBsAg positive women during pregnancy and in cord blood of their neonates.

Prenatal transmission

The possibility of prenatal transmission of HBV has been speculated to be low. However, mothers with HBeAg positivity known to be associated with higher HBV DNA viral loads have been linked with higher chances of MTCT as HBeAg is the only structural HBV protein that can cross into the placental circulation.

Intra-partum transmission

The intra-partum period has been shown to be associated with higher chances of MTCT of HBV. This exposure occurs via micro-transfusion or haematologic leaks of maternal blood to the fetus during contractions or through inoculations of mucosal membranes or breaks in the skin (e.g. during some obstetric procedures including insertion of fetal scalp electrodes in settings where it is practised).

The mode of delivery (vaginal or caesarean section) does not seem to increase or decrease the risk of perinatal HBV infection. Most studies found no difference in MTCT among babies delivered by caesarean or spontaneous versus operative vaginal deliveries among infants who received post exposure prophylaxis (PEP). However there is no consensus recommendation as per the preferred mode of delivery in high risk mothers.

Breastfeeding

Though possible markers of HBV have been demonstrated in breast milk and colostrum from HBsAg – positive women; transmission of HBV through breast milk is not a significant source of infection, as reported in some pre-HBV routine neonatal prophylaxis studies.

Reported prevalence of HBV infection in breastfed and non-breastfed infants appear same, however, some of the studies did not take maternal HBV DNA viral load into cognizance. Also, mothers with cracked or bleeding nipples may momentarily stop breastfeeding as meta-
analysis of studies among mothers with absence of such symptoms did not identify an increase in MTCT rates of HBV when breastfed babies received post exposure prophylaxis (PEP).22

It is recommended that HBV positive mothers should breastfeed their infants provided that the infants receive timely and adequate PEP at birth.

**Clinical features**

Most cases of acute HBV infection usually manifest with non-specific presentations including nausea, vomiting, loss or poor appetite, fever, myalgia and weakness.22

One percent of cases of acute hepatitis including those occurring in neonatal period may develop fulminant hepatitis. Course of cases of acute, fulminant hepatitis B virus infection could be irreversible and result in liver failure with possible mortality.33

Majority of cases of HBV infections including acute and chronic hepatitis are asymptomatic. Chronic HBV infection however has three immunologic phases: immune tolerant (absent or minimal liver inflammation, high viraemia); immune active (alanine amino-transferase rises, high viraemia, liver inflammation rises; high viraemia, liver inflammation, and fibrosis improve, anti-HBe is present); and the inactive chronic hepatitis (the non-replicative or inactive immune-control phase) which follows successful sero-conversion from an HBeAg – positive to anti-HBe state.34

The immune tolerant phase usually appears in individuals who had perinatal infection from HBeAg positive mothers.

**Preventive measures**

**Prenatal hepatitis b virus infection screening**

Chronic HBV infection is characterized by the presence of HBsAg for at least 6 months (with or without concurrent HBeAg positivity). The persistence of HBsAg is the principal marker of risk of developing chronic liver disease and hepatocellular carcinoma later in life.9

It is recommended that all pregnant women should be tested routinely for HBsAg during an early prenatal visit (preferably in the first trimester) in every pregnancy irrespective of if they had been previously vaccinated for HBV or tested.1

Un-booked pregnant mothers or women who were not screened prenatally particularly those who engage in behaviours that put them at high risk for infection especially intravenous drug users, those with more than one sexual partners in the previous six months or an HBsAg-positive sex partner, evaluation or treatment for a sexually transmitted disease and those with clinical hepatitis should be tested at the time of admission to the hospital for delivery.35

The prenatal HBsAg screening is key to successful HBV PMTCT programme as the maternal status is obtained early enough and appropriate measures applied.36

Rapid immunochromatographic hepatitis B surface antigen tests with acceptable performance are available to identify HBV infected pregnant women.21

Cost-effective point-of-care tests for combined HBsAg and HBeAg detection and determination of HBV viral load are currently being developed37 and further resources should be channeled towards making it a success story.

**Prenatal prevention for mothers**

The maternal HBeAg positivity as well as elevated HBV DNA viral load are directly related to rate of intrauterine transmission.38 Therefore reduction of maternal DNA viral load of HBV is an effective method of decreasing the rate of HBV infection in infants particularly in endemic areas.39,40

**Use of anti-viral agents in pregnancy**

The probable reasons for use of anti-viral (anti-retroviral) therapy in chronic hepatitis B in pregnancy are based on maternal viral load, liver enzyme levels (liver function test), HBeAg status, liver histology, and HIV co-infection status.41

The benefits of anti-retroviral (ARV) prophylaxis during pregnancy is to decrease viraemia so as to limit breakthrough HBV infection.42 The administration of anti-retroviral therapy (ART) starting from late pregnancy will reduce maternal HBV viral load and potentially reduce the possible risk of MTCT.42

The use of antiviral agents during pregnancy mainly for prophylaxis of perinatal HBV transmission would entail diligent evaluation of potential risks and benefits among infants and their pregnant mothers alike.43 Some studies in animal specimens have shown severe growth restriction and reduced bone mineral density in exposed fetuses to ART.

However, recent efficacy data on prophylaxis of PMTCT of HBV have shown acceptable safety profile in pregnant women. Equally commencement of the ART in the third trimester of pregnancy among mothers with high viral load (e.g. ≥200U/mL) to prevent breakthrough perinatal HBV transmission has also been successful without concerns. Tenofovir, lamivudine and telbivudine are nucleos(t)ide inhibitors which act as chain termination in DNA elongation and can be administered from 28 weeks of pregnancy.23

Initially lamivudine (3TC) was commonly used with benefit of reduction in transmission of HBV transmission; however, it was shown to be associated with development of resistance owing to its low genetic barrier. On the other hand, tenofovir, has a high barrier to resistance and has been used extensively in the setting of HIV in pregnancy. Use of tenofovir in pregnancy has resulted in
significant reductions among mothers, without any remarkable adverse effects.44,45

Based on safety data from the ARV in Pregnancy Registry in pregnant HIV positive women who have received tenofovir and/or lamivudine or emtricitabine; tenofovir appears to be the preferred ARV, owing to its better resistance profile, and more extensive safety data in pregnant HBV positive mothers.23,43

As ongoing caution is being advised on the use of ART in pregnancy; the absence of adverse drug events in mothers and their infant pairs who received ART in pregnancy up to date offers promise in their use.44 Some guidelines recommend treatment for non-cirrhotic patients (including pregnant woman) with serum HBV DNA levels greater than 20,000 IU/mL (> 10^5 copies/mL) and evidence of liver diseases.55

Intra-partum/Neonatal transmission

Use of monitors should be avoided as much as possible. In centres where it is obtainable the use of fetal scalp electrode monitoring as well as fetal blood sampling should be avoided particularly if mother is HBsAg positive. Also standard precautions should be implemented for care of all women in labour. Appropriate cleaning of the newborn should equally be practiced upon delivery the neonate’s eyes as well as the non-intact skin should be cleaned properly using water as soon as possible after birth.46

Immunoprophylaxis

The double usage of active and passive immunoprophylaxis is the ideal measure to prevent MTCT of HBV infection.47 PEP with hepatitis B vaccine and hepatitis B immune-globulin (HBIG) administered within 12 to 24 hours after birth followed by completion of a three dose vaccination series, has been shown to protect 85 to 95% of infants whose mothers were positive for both HBsAg and HBeAg.45

Birth dose hepatitis B vaccine

The recombinant DNA – derived hepatitis B vaccines have been available and licensed for use since 1982. The World Health Organization (WHO) as well as the Centers for Disease Control (CDC) recommends universal HBV vaccination for all infants, and that the first dose should be given as soon as possible after birth preferably within the first 12 hours of birth.55

In Nigeria the universal HBV vaccination was initiated by the Federal Government in 2004; despite that huge gap still exists in meeting the targets as majority of the children receive the birth dose vaccine outside the recommended time owing to maternal as well as programme – related factors.49,50

The birth dose of HBV vaccine is very important as that serves as a safety net as non-testing, errors in testing, reporting and documentation of maternal HBsAg status could occur.51

In the United States, the CDC in 2005 recommended that the Advisory Committee on Immunization Practices (ACIP) and all health professionals should administer the HBV vaccine to all newborns before hospital discharge in order to protect them against the HBV infection.52

However, in Nigeria it has been noted that many babies whom were delivered in health facilities do not receive the birth dose of recommended vaccinations before the hospital discharge.53 Nigeria recommends that the birth dose of HBV should be given preferably within the first 24 hours of birth but up to first 2 weeks of life. This birth dose is usually given as a monovalent vaccine. It is of note that giving the birth dose after the neonatal period reduces the efficacy of PEP which decreases with increasing time as exposure.54

Nigerian HBV Vaccination Schedule

The birth dose of HBV vaccine followed by two more doses to complete the primary series is the conventional. However, WHO recommends that four doses may be given in line with the programmatic protocol of National Schedules.55

The pentavalent vaccine which replaced the trivalent DPT was introduced in Nigeria in 2012, and contains HBV vaccine among others as a combination vaccine to be commenced at age 6 weeks through ten and 14 weeks respectively. However, Nigeria currently gives the birth dose in addition to three combined doses in form of pentavalent vaccine doses.

The complete vaccine series induces protective antibody levels in more than 95% of infants, children and young adults. Protection lasts at least 20 years and is probably life long. The vaccine has excellent record of safety and effectiveness and the immunogenicity of the combined hepatitis vaccines is equivalent to that of their individual antigens administered separately.1

Immunization escape mutants in the S – gene of HBV have been detected, and concerns have been expressed that these variants could replicate in the presence of vaccine induced anti-HBS or anti-HBS contained in HBIG. However no evidence exist that s-gene immunization escape mutants posing a risk to currently existing programmes using hepatitis B vaccine.55

Further studies including surveillance are needed to track any possible emergence of these variants in future.

Vaccinations in preterm/Low Birth Weights

Previously, the American Academy of Pediatrics (AAP) routinely recommended that the birth dose of HBV vaccine for preterm and low birth weight infants born to HBsAg negative mothers be delayed up till the infant
weighed 2 kilograms or is two months of chronological age,\textsuperscript{56} owing to reduced immunogenicity in preterm less than two kilograms (<2kg).\textsuperscript{56} However recent guidelines by the AAP recommends that such children should receive the first dose of hepatitis B vaccine by 30 days of chronological age irrespective of the gestation age. Also if such preterm/LBW shows consistent weight gain necessitating hospital discharge before 30 days of age, the first dose of hepatitis B vaccine should be given at the time of hospital discharge. The remaining three doses of hepatitis B vaccine series should be completed on the same schedule as applicable to full term infants, ensuring that the second dose is given at least 30 days after the initial dose.\textsuperscript{56}

Also of note is that currently the ACIP does not recommend routine booster doses of hepatitis B vaccine for persons who completed the primary HBV immunization series in childhood. \textsuperscript{57}

The immunogenicity of hepatitis B vaccine is low in individuals with severe immunosuppressive states including AIDS, and timely administration of the HBV vaccine commencing at birth should be the rule rather than the exception as the HIV immunosuppression worsens overtime among infected children.\textsuperscript{58}

Hepatitis B vaccination is recommended in HBsAg negative women in pregnancy with high likelihood of acquisition of the infection particularly in those having multiple sexual partners or intravenous drug abusers.\textsuperscript{58}

**Passive immunization**

It provides passively acquired anti-HBS and temporary protection when administered in standard doses. Hepatitis B immunoglobulin (HBIG) is typically used as an adjuvant to the hepatitis B vaccine in the PMTCT of hepatitis B infection in exposed infants, providing temporarily protection for 3 to 6 months.

Administration of hepatitis B immunoglobulin and hepatitis B vaccine within 24 hours of birth, followed by the completion of the vaccine series is 85 – 95% efficacious in PMTCT of HBV.\textsuperscript{14}

HBIG administered alone is the primary means of protection after an HBV exposure for non-responders to hepatitis B vaccination.

**Follow up**

MTCT of hepatitis B virus continues to occur despite the use of adequate preventive measures including vaccinations and HBIG administration.\textsuperscript{15} The ACIP recommends post-HBV vaccination testing for all infants born to HBsAg positive women. The testing consists of HBsAg for infection and anti-HBS for response to vaccination. The optimal timing for detecting protective antibodies is one to 2 months after the final Hepatitis B vaccine dose at ≥ 9 months of age. The testing for follow-up should commence at 9 to 18 months of age, at least 1 month after the last dose of vaccine.\textsuperscript{46}

The anti-HBS levels done earlier than 9 months of age may reflect passive immunization with HBIG.\textsuperscript{28} Anti–HBS levels of more than 10m/u/L indicate adequate protection whereas babies with anti-HBs levels of less than 10m/u/L need to be re-vaccinated with the entire 3 dose schedule.

Babies who are HBsAg positive are infected and need further evaluation and follow-up by a paediatric gastroenterologist.

Currently Nigerian Society for Paediatric Gastroenterology Hepatology and Nutrition (NISPGHAN) and Society for Gastroenterology and Hepatology in Nigeria (SOGHIN) are in the forefront of spreading the good news of HBV prevention in Nigeria and should be encouraged by the authorities by doing the needful.

**Conclusion**

Universal screening of all pregnant women for HBV infection is the most effective strategy for the prevention of MTCT of HBV, as effective preventive measures could be applied starting from pregnancy to delivery while infants of HBsAg positive mothers should receive timely and adequate PEP as well as follow up for possible development of chronic hepatitis B infection.

**Author’s contributions**

CBE: Study conceptualization, literature search, manuscript draft, critical editing of the manuscript for important intellectual content.

NBO, OFA: Critical editing of the manuscript for important intellectual content.

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