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**Current evidence on the management of hepatitis B in pregnancy**

Maraolo AE *et al.* HBV in pregnancy

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**Abstract**

Hepatitis B virus (HBV) infection is one the main public health problem across the globe, since almost one third of the world population presents serological markers of contact with the virus. A profound impact on the epidemiology has been exerted by universal vaccination programmes in many countries, nevertheless the infection is still widespread also in its active form. In the areas of high endemicity (prevalence of hepatitis B surface antigen positivity > 7%), mother-to-child transmission represents the main modality of infection spread. That makes the correct management of HBV in pregnancy a matter of utmost importance. Furthermore, the infection in pregnancy needs to be carefully assessed and handled not only with respect to the risk of vertical transmission, but also with respect to gravid women health. Each therapeutic or preventive choice deserves to be weighed up attentively. On many aspects evidence is scarce or controversial. This review will highlight the latest insights into the paramount steps in managing HBV in pregnancy, with particular attention to recommendations from recent guidelines and data from up-do-date research syntheses.

**Key words**: Hepatitis B; Pregnancy; Therapy; Antiviral prophylaxis; Immunoprophylaxis; Hepatitis B immunoglobulin

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**Core tip:** Hepatitis B is still a matter of concern worldwide. Particularly challenging is the correct management of infection during pregnancy. Two aspects have to be taken into account: The potential need to treat the mothers and, at once, the necessity to prevent the vertical transmission of the virus to the infants. This review will discuss the most up-to-date evidence upon therapeutic and preventive interventions in the several scenarios characterizing the course of hepatitis B in pregnancy.

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**INTRODUCTION**

Despite the availability of effective preventive measures, particularly active immunization through vaccination[1], hepatitis B virus (HBV) infection is still today a major public health issue worldwide, being the 3.61% of the global population chronically infected, as expressed by the prevalence of hepatitis B surface antigen (HBsAg)-positivity, according to the most recent and robust estimates[2]. Of course, there is a relevant heterogeneity both across and within continents and states as far as endemicity is concerned: Western Pacific Region and Africa are the world areas with the highest prevalence[2].

Even larger is the number of people having serologic markers of previous contact with HBV: About 2 billion subjects worldwide[3]. These individuals, showing markers of resolved/occult HBV disease, deserve special attention in case they undergo immunosuppressive treatment[4]. Nevertheless, and not surprisingly, the major disease burden is related to chronic infection: In 2013 HBV was responsible of nearly 686000 deaths globally, a figure that places the virus in the top 20 causes of mortality among humans[5]. Of note, differently from other major communicable diseases, the burden of viral hepatitis, mainly driven by HBV and hepatitis C virus (HCV), has increased in terms of morbidity and mortality between 1990 and 2013[6].

Chronic HBV infection may evolve to cirrhosis, a condition characterized by profound alteration of liver architecture and function, in about 20% of subjects and may result in hepatocellular carcinoma (HCC), as a consequence of cirrhosis itself or of viral pro-oncogenic properties[7]. In turn, cirrhosis may be responsible for a vast array of complications: Infections[8], mainly spontaneous bacterial peritonitis[9] and bloodstream infections[10], ascites[11], hepatorenal syndrome[12], variceal bleeding[13].

Despite the remarkable efforts during the last decades aimed at implementing effective vaccination strategies worldwide[14], 50 million new cases of hepatitis B are still diagnosed each year, most due to mother-to-child transmission (MTCT)[15].

As matter of fact, the transmission routes differ according to the entity of HBV endemicity: In areas of high prevalence (> 7%), the vertical transmission prevails, whereas in low endemic regions (prevalence < 2%) sexual transmission is the major culprit[16]. The way and the timing of transmission are crucial factors influencing the probability of developing chronic HBV infection: Indeed, this likelihood is higher in subjects infected perinatally (up to 90%) when compared with rate of chronicity in adults after the acute phase (< 1%)[17]. Around 15%-40% of individuals suffering from chronic hepatitis develops cirrhosis[18].

All these figures underpin the necessity of correctly managing pregnant women with HBV infection, in order to reduce the burden of disease. Attention must be paid not only in developing countries, but also in regions such as Australia, United States, Western Europe, where immigration from areas of high HBV endemicity may represent a challenge for physicians not accustomed to manage HBV infection in particular settings[15]. Of course, HBV infection in pregnancy not only is a problem for infants, but also for women’s health (Figure 1). In this review, the current state of the art regarding the best management of HBV in pregnancy, both for the mother and child, will be discussed.

**HBV INFECTION IN PREGNANCY: FROM THE PERSPECTIVE OF GRAVID WOMEN**

***How HBV impacts on the health of pregnant subjects***

The relationship between liver diseases and pregnancy is proteiform, and three categories of pathological conditions can be described: The ones representing underlying status, pre-existing to the moment of conception; the ones coincidental with maternity; eventually, the ones specific of pregnancy (for example, pre-eclampsia)[19]. Viral hepatitis can fall into the first two categories[19].

As far as acute hepatitis B is concerned, its occurrence during pregnancy is not associated with higher mortality, and also the related clinical picture is not distinguishable from that in the general population[20]. This notion was further confirmed by a case-control study run in China, comparing 22 pregnant patients and 87 matched non-gravid women, all suffering from acute hepatitis B: No difference with regard to mortality and incidence of fulminant hepatitis was detected[21]. Of note, the HBsAg loss and seroconversion rates were lower in the first group, suggesting that pregnancy might act as a risk factor for chronicity[21].

An interesting and recent systematic review has assessed the impact of inactive HBV carriage on gravid women health, showing that this condition is not associated with complications in pregnancy, so this condition does not need any particular therapeutic measure[22]. When it comes to chronic (active) HBV infection already established before conception, the immunological modifications that occur in pregnancy may raise the level of HBV viremia, whereas alanine aminotransferase (ALT) levels are normal or just above the upper limit of normal (ULN)[23].

A more relevant exacerbation of chronic hepatitis might happen after delivery in a notable percentage of women (up to 45%), as observed in a small retrospective cohort involving 38 pregnancies in 31 subjects, in which the flare was defined as three times increase in ALT levels within 6 mo post-partum[24]. Authors suggested that this phenomenon, also found in the subgroup of women (8/13, 62%) who had undergone a course of lamivudine (LAM) during the third trimester, was attributable to the restoration after delivery of the immune system, whose functions are previously altered to prevent foetus rejection[24]. The topic has been further elucidated by a subsequent and prospective study recruiting 126 women: Post-partum flares, defined as ALT levels twice the ULN or the baseline, were described in 27 (25%) individuals, usually asymptomatic and with spontaneous resolution[25]. At multivariate analysis, HBeAg positivity turned out to be the most relevant predictor of post-partum flares, although just barely not reaching the statistical significance (*P* = 0.051)[25]. Even a further study, a multicentre retrospective cohort involving 101 women and 113 pregnancies, did not identify clear risk factors for exacerbation of chronic hepatitis B after delivery[26].

With regard to maternal complications, chronic HBV infection does not seem a risk factor for many of them, as derived by research syntheses. A meta-analysis collecting data on 9088 placenta previa cases and 15571 placental abruption cases failed to demonstrate an association with HBV, implicated as driver of an inflammation state able to induce dysfunction of trophoblasts: Odds ratio (OR) equal to 0.98 with 95%CI equal to 0.60-1.62 and OR = 1.42 with 95%CI: 0.93-2.15, respectively[27]. A further meta-analysis involving 439514 subjects showed that HBV was not associated with increased risk of gestational diabetes mellitus (adjusted OR = 1.11, 95%CI: 0.96-1.28), a link suggested by the potential role played by the virus in inducing insulin resistance[28]. Another research synthesis of observational studies did not detect a statistical significance between chronic HBV infection and preterm labor (OR = 1.12, 95%CI: 0.94-1.33)[29]. More recently, a meta-analysis including 11566 women has, quite surprisingly, highlighted a negative association between chronic HBV infection and preeclampsia was observed (OR = 0.77, 95%CI: 0.65-0.90, *P* = 0.002): Actually the protective effect, probably due to impaired immune response and/or increased immune-tolerance caused by the virus (preeclampsia is linked with exaggerated activation of immune system), was apparent only in Asian population, as derived from the subgroup analysis[30].

Another aspect to be considered is how the most advanced stage of chronic liver disease, cirrhosis, impact on the health of pregnant women, in the particular setting of HBV infection. Cirrhosis is, fortunately, an infrequent occurrence in pregnancy due to two factors: The development of end-stage liver disease requires time and more often takes place when women have gone beyond their reproductive age; moreover, hypothalamic-pituitary dysfunction related to cirrhosis[31] may ensue in anovulation and amenorrhea[32]. However, when present, cirrhosis is a relevant health issue for pregnant women. In a large population-based retrospective study in the United States on gravid women, comparing 339 cirrhotic cases with 6625 matched-controls, maternal mortality and complications of pregnancy (*e.g.*, uterovaginal haemorrhage, pre-eclampsia, peri-partum infections) were higher among individuals suffering from liver disease: For example, the maternal death rate was 1.8% *vs* 0% (*P* < 0.0001)[33]. Mortality among cirrhotic pregnant women was higher in case of viral aetiology (HBV as well as HCV)[34]. The high burden of liver cirrhosis in pregnancy has been confirmed in a more recent prospective study, matching 176 cirrhotic gravid women with 2179 pregnant non-cirrhotic women and 1034 cirrhotic but not pregnant female subjects[34]. Maternal mortality rate was superior in the study group (7.8%) than in the first (0.2%) and in the second control group (2.5%; *P* = 0.001); variceal haemorrhage during vaginal delivery was the most frequent reason of maternal death[34]. Indeed, the rupture of oesophageal varices represents probably the most important complications among the ones directly related to cirrhosis in pregnant women, especially in the advanced phase of pregnancy or during labor[35]. An important predictor of liver-related complications during pregnancy is a model for end-stage liver disease (MELD) score ≥ 10[36].

***Treatment criteria for acute and chronic hepatitis B in pregnancy***

The paramount issue is: Which pregnant women with HBV infection should be treated[37]?

In case of acute hepatitis B, the main goal of the treatment should be the prevention of acute liver failure[38]. The quality of current evidence upon pharmacological interventions in this setting is unfortunately very low[39]. Nevertheless, rarely antiviral therapy is necessary, since the large majority of adult patients (> 95%) have a full and spontaneous recovery[38], and, as mentioned above, the clinical course of this entity does not differ between pregnant and non-pregnant women. The problem is the management of cases suffering from severe acute HBV infection[40]. First, in case of serious hepatitis affecting gravid women, differential diagnosis is essential, to rule out, for example, diseases unique to pregnancy such as acute fatty liver of pregnancy (AFLP) as well as haemolysis, elevated liver enzymes and low platelets (HELLP) syndrome, representing two hepatic emergencies in the third trimester[40].Once the viral aetiology is established, borrowing the recommendations applying to general population, the treatment should rely on nucleos(t)ide analogue (NA) agents and the patients should be considered, as extrema ratio, for liver transplantation[38]. Therapy with NAs can prevent acute liver failure and the related mortality[38], but needs to be started early in the course of severe acute hepatitis, otherwise the protective effect does not display[41]. Lacking high-quality data[39], there is high uncertainty regarding the best therapeutic options; recommendations for general population support the use of tenofovir disoproxil fumarate (TDF), entecavir (ETV), lamivudine (LAM)[38]. To date, there is no information about the use in severe acute hepatitis B of tenofovir alafenamide (TAF), the prodrug of tenofovir developed to ameliorate the safety and tolerability profile of TDF[42]. Among the above-mentioned NAs, LAM and TDF are preferable in pregnancy, in particular the second one, owing to its high resistance barrier that is fundamental in case of prolongation of therapy for chronicity[43]. In extreme circumstances, liver transplantation might be a therapeutic option even during pregnancy, if hepatic decompensation exceeds the point of no return: The difficulty of the task is doubled, being involved two organisms (the mother and the foetus), but successful cases are described, also related to fulminant hepatitis B[44].

At any rate, the management of severe acute hepatitis B in pregnancy would need more robust data to draw firm conclusions. Today, a case-by-case approach is needed. Chronic HBV infection is surely a more common scenario in pregnancy than fulminant hepatitis B. In general population, according to the most authoritative guidelines endorsed by societies from all over the world, the main criteria for treatment are based on serum HBV DNA levels, serum ALT levels and severity of liver disease[38,45,46]. Despite some discrepancies (for example, the locution “inactive carriers”, discouraged by Asian and European guidelines but kept in the American ones), there is a substantial consensus upon the following items, concerning situations wherein antiviral therapy is recommended: Cirrhosis; absence of cirrhosis, but viraemia > 20000 International Units (IU)/mL and ALT levels > 2×ULN; no cirrhosis, viraemia > 2000 IU/mL, ALT 1-2 × ULN but at least moderate to severe inflammation on liver biopsy[38,45,46]. In general population, there are two therapeutic options: Interferon-alfa (IFNα) and NAs[16]. The rationale underpinning the choice of one strategy over another one is different: IFNα is administered to provide a long-term immunological control through a finite duration treatment, attempting to achieve the so-called “functional cure” by HBsAg loss, but it is burdened by several side effects; NAs have a definitely better safety and tolerability profile, provide a very good virological control (persistent inhibition of HBV replication), but the duration of therapy is indefinite[47].

At any rate, in pregnant women IFNα is contraindicated, therefore the therapeutic armamentarium is limited to NAs[48]. Among this category, currently TDF (alternatively TAF) and ETV are considered the first-line drugs when starting a new therapy for chronic HBV infection, combining excellent safety profile with a genetic barrier of resistance higher than earlier available agents such as LAM and telbivudine (LdT)[38,45]. Nevertheless, according to the 5-class labelling used by the Food and Drug Administration (FDA) until 2015 as for safety in pregnancy, ETV has been classified as “category C” agent, differently from LdT and TDF, labelled as “category B” drugs: That means the absence of teratogenic effects in animal studies[49]. As matter of fact, TDF is the drug of choice for pregnant females with chronic HBV infection requiring antiviral therapy (*i.e.*, advanced fibrosis and cirrhosis), also in the light of the huge amount of data from the setting of gravid women under treatment against the human immunodeficiency virus (HIV), in which TDF is administered safely in combination with other antiretroviral agents throughout the pregnancy, since the first trimester[50].

A delicate issue is how to handle cases of women who become pregnant when already on anti-HBV treatment[51]. As a rule, appropriateness of therapy should be re-evaluated, striking a balance between benefits for the mother and the safety of the foetus[50]. Whichever the diseases severity, IFNα must be immediately stopped; therapy with a NA can be continued in case of advanced fibrosis or cirrhosis, switching to TDF if therapy was started before conception with another drug (for instance, ETV)[51]. In case of mild disease (*e.g.*, no advanced fibrosis, normal ALT levels, viraemia between 2000 UI/mL and 20000 IU/mL) discontinuation of therapy until delivery might be a viable option, as long as an adequate monitoring is carried out to immediate re-start treatment if necessary[51]. Eventually, another matter of concern is the management of HBV resistance cases: In pregnant women there is limited experience, nonetheless, in gravid subjects experiencing treatment failure (HBV DNA rebound) under LAM or LdT, switching to TDF appears a safe and effective option[52]. In Figure 2 the current knowledge regarding the treatment of HBV infection in pregnant women is summarized.

**HBV INFECTION IN PREGNANCY: FROM THE PERSPECTIVE OF FOETUSES AND NEWBORNS**

***Does HBV damage the product of conception?***

Besides the “long-term” risks of HBV MTCT such as chronic hepatitis, cirrhosis and HCC[53], physicians caring pregnant women with HBV infection have to take into account the “short-term” potential consequences for the product of conception: For example, small for gestational age (SGA), foetal distress, preterm birth (PTB), low birth weight (LBW), congenital anomalies and neonatal jaundice[54].

PTB, defined as a birth occurring earlier than 37 completed weeks of gestation, represents one of the most feared complications, being worldwide the main cause of death in children under 5 years of age[55]. A very large population-based cohort study run in China, involving 489965 women who had singleton livebirths (of whom 20827, the 4.3%, with HBV infection diagnosed before pregnancy), showed that, adjusting for several covariates, in comparison with gravid subjects without HBV infection, HBsAg positive and HBeAg negative pregnant women had a 26% higher risk of PTB, whereas in women who were both HBsAg and HBeAg positive this percentage was equal to 20%[56]. Higher risk were observed also as for early PTB (before 34 wk of gestation); unfortunately, data on viral load were not available, at any rate the results of this recent study advocates proper medical intervention against HBV in pregnancy to improve neonatal outcomes[56].

Another large population-based study (sample size over 2000000 people), conducted in the United States and focused on neurological complications at birth, demonstrated that women with HBV, compared with gravid subjects without HBV, had a higher likelihood to generate infants who suffered from brachial plexus injury, even after adjusting for several confounders (OR = 2.04, 95%CI: 1.15-3.60)[57]. In a prospective cohort study in China, investigating 21004 pregnant women, of which were 513 HBV-positive and 20491 HBV-negative, no differences between the two groups were detected with regard to the rate of stillbirth, SGA and LBW, but the proportion of miscarriage was higher among gravid subjects with HBV (adjusted OR = 1.71, 95%CI: 1.23-2.38), but also in this study data on viraemia were lacking[58].

***Criteria and options for antiviral prophylaxis against HBV MTCT***

The viral load is just the key factor to determine the risk of HBV MTCT[59], that, in absence of any preventive measure, ranges from 10% to 40% when mothers are HBeAg-negative and from 70% to 90% in HBeAg-positive mothers[60], much higher rates in comparison with the other hepatotropic virus, HCV (0%-30%)[61].

The modalities of vertical transmission are: Intrauterine, peripartum and post-natal infection[59]. The transmission in utero is the most insidious route, since it is represents the most important cause of passive-active immunoprophylaxis failure[62]. There is no consensus to correctly define this occurrence (many criteria have been proposed, such as, among the many, persistent serum anti-HBc IgM positive after birth or Pre-S1 protein positivity in umbilical blood); supposed mechanisms are the passage of serum/body fluid through damaged placenta, the transmission of infected germ cells and the transfer of infected placenta or peripheral blood mononuclear cells[62]. HBeAg-positivity is a notable driver of intrauterine transmission: The antigen can cross the placenta barrier through leakages or through infected cells, and it is linked with higher levels of HBV replication[35]. Natal transmission during delivery represents the most impactful modality, being offspring exposed to blood or other maternal body fluids while passing the genital tract[62]. Finally, postnatal infection, encompassing all the cases in which the transmission occurs after delivery, because of contacts with maternal fluids such breast milk or blood[62].

All guidelines, in line with the recommendations of the World Health Organization, support the administration, within 12 h of birth, of active (first dose of anti-HBV vaccine) and passive immunization through hepatitis B immunoglobulin (HBIG) to the offspring born to HBsAg-positive mothers, a measure able to abate the rate of HBV MTC to > 90% to < 10%[38,45,46] and supported by high-quality evidence[63]. The paramount issue is how to avoid the failure of passive-active immunoprophylaxis, mainly ascribable to intrauterine transmission[64]. The most recent international guidelines recommend, for gravid women not already on NAs treatment, the use of antiviral prophylaxis from week 24-28 of gestation (third trimester)if viral load > 200000 IU/mL and/or serum HBsAg levels > 4 log10 IU/mL[38,45]. The viraemia threshold was first set based on a retrospective study involving 869 mother-newborns pairs who had received proper immunoprophylaxis: Failures occurred only in infants born to HBeAg-positive women with viral load > 200000 IU/mL (maternal viraemia levels, along with detectable HBV DNA in the cord blood, was the main risk factor at multivariate analysis)[65]. Subsequently, serum HBsAg levels emerged as surrogate marker for viral load as well as predictive variable of HBV MTCT[66,67].

The benefit of antiviral prophylaxis as additional measure to HBIG and vaccination was clear in a meta-analysis of 26 studies involving 3622 pregnant women, with a risk ratio equal to 0.3; the use of LdT, LAM, and TDF turned out to be safe[68]. There is no randomized controlled trial (RCT) directly comparing NAs as for HBV MTCT prevention: A Bayesian network meta-analysis (NMA) in 2016 demonstrated greater efficacy of LdT over LAM, but TDF was not taken into account[69]. A more recent NMA failed to demonstrate a superiority of TDF *vs* LAM[70]. At any rate, TDF is the favourite choice because of his superior barrier to resistance[38,45]. Indeed, TDF was the drug of choice for an non-randomized trial and two RTCs *vs* placebo published during the last three years[71-73]. The first two studies demonstrated that TDF (administered throughout the last trimester of pregnancy until 1 mo post-partum) decreased significantly the rate of HBV vertical transmission in comparison with placebo in women HBeAg-positive having high viral load[71,72]. On the contrary, the last and more recent RCT, not considered by guidelines due to publishing timing reasons, failed to detect a significative difference between the TDF group and the placebo arm (no events out of 147 newborns *vs* 3 infection out of 147 infants, respectively, *P* = 0.29)[73]. The study protocol contemplated the administration of TDF or placebo from 28 wk of gestation to 2 mo post-delivery in HBeAg-positive women, the large majority of them having viraemia > 20000 UI/mL, and its sample size was as large as the ones of the previous studies taken altogether[73]. Therefore, the results of this negative trial brings into question the usefulness of NAs, specifically TDF, as additional preventive measure during last period of pregnancy and will need to be considered and put in the right perspective by the next research syntheses and guidelines; one of the possible explanation is the very early administration of HBV vaccination (the median time was just 1.2 h after birth)[74].

Pending new compelling evidence, the combination of HBIG and vaccine at birth is the mainstay of HBV MTCT prevention in newborns; antiviral prophylaxis in late pregnancy may be considered for HBeAg-positive gravid women with high viral load[75]. Furthermore, the absence of harm, weighing risks and benefits, might tip the scale in favour of NAs administration during the last trimester: The alarm raised by a case-control study (74 TDF-exposed and 69 TDF-unexposed infants) about the risk of lower neonatal bone mineral content (difference equal to 12% at 1 mo of birth) because of TDF during late pregnancy[76] has been refuted by a subsequent work (conducted by the same study group) on 509 children: At 2 years of age TDF was not linked with lower length or head circumference[77]. This is in accordance with evidence from research synthesis confirming the safety, both for mothers and their offspring, of TDF use in pregnancy[78]. If administered, there is no consensus about when to stop prophylaxis: Some guidelines support its prolongation until 12 wk after delivery[38], others until 4 wk post-partum[45]: The protocol of main trials about TDF provided for the use of drug for 4[71,72] or 8[73] wk after delivery. The point is to strike a balance between the potential risk of interfering with breastfeeding and the benefit on possible post-partum hepatitis flares[35]. More conservative recommendations[45] rely on a prospective study recruiting 91 women (101 pregnancies), showing no advantages in terms of hepatitis flare rate for gravid subjects who extended antiviral prophylaxis with TDF beyond 4 wk after delivery[79]. Nevertheless, prolongation of antiviral prophylaxis[38] might be useful at least for women with elevated ALT during pregnancy, since they present a higher risk of post-partum hepatitis flare, as showed by a Chinese study wherein mothers were administered LdT[80]. With regard to other preventive strategies, unfortunately there is high uncertainty, also due to very low available evidence, upon the potential benefits of the antenatal administration of HBIG, to exploit the materno-fetal diffusion through the placenta, that reaches its peak during the third trimester[81].

***Prevention of HBV MTCT: Beyond pharmacological options***

The last issues involve the following topics: Delivery modalities, invasive procedures during pregnancy and breastfeeding[38,45]. There is a huge debate about the efficacy of caesarean section (C-section) as preventive measure. Guidelines do not back its elective implementation[45], although meta-analyses reveal that C-section, compared with vaginal delivery, significantly decrease the risk of HBV vertical transmission[82,83]. The problem is the high heterogeneity of the studies whose results have been retrieved and analysed by these research syntheses, one collecting data from 10 studies[82] and the most recent from only Chinese datasets[83]: These relevant limitations advocate well designed studies to be performed in order to shed light on this matter.

Unfortunately, there is scarce evidence regarding the best practice when invasive procedures are carried out. As far as amniocentesis is concerned, a quite recent matched case-control study (63 infants whose HBsAg-positive mothers had underwent the procedure and 198 newborns whose HBsAg-positive mothers had not underwent amniocentesis) found that HBV MTCT was more frequent among cases (6.35% *vs* 2.53%; *P* = 0.226); notably, the difference was apparent when maternal viral load was taken into account, especially above the threshold of 200000 IU/mL (50% *vs* 4.5%, *P* = 0.006)[84]. Neither cases nor controls were born to mothers who were administered antiviral prophylaxis during pregnancy[84]. No strong recommendations can be drawn on this basis, therefore, waiting for studies that will investigate the potential role of antiviral prophylaxis in women with high viraemia undergoing amniocentesis, guidelines suggest that a careful assessment of harms and benefits of the invasive procedure is necessary[45].

The last topic is breastfeeding. On one hand, lactating is allowed as long as the standard measures of passive/active prophylaxis are taken[38,45,46]. On the other hand, there are some concerns about the safety of NAs, particularly TDF, during breastfeeding[38,45,46]. Experiences in HIV field indicate that antivirals are well tolerated[85] and in particular TDF appears to be safe as to infant outcomes[86]. In Figure 3 a summary of the current knowledge regarding the HBV MTCT prevention is depicted.

**CONCLUSION**

When facing HBV in pregnancy, there are two different problems to address: The first is represented by the maternal liver disease, the second by the risk of MTCT. The two issues are actually strictly inter-connected, but choices regarding potential antiviral use can profoundly differ, especially as for timing. Unfortunately, to date many questions present answers backed up by low-quality evidence, a not rare occurrence when pregnancy is involved: For instance, it is not simple to set up large and multicentre RCTs in this setting. Moreover, there is constant need to take carefully into account benefits and harms of each intervention, potentially impacting not on one but on two lives. Regarding the first issue, in essence the indications for treatment of general population also apply to pregnant women. The drug of choice is represented by TDF; in case the gravid subjects are already on treatment with another NA, a switch is advised. IFN is absolutely contraindicated.

As to the second issue, the only mandatory measure, underpinned by incontrovertible evidence, is represented by providing passive and active immunoprophylaxis to the newborns, starting the schedule as early as possible at birth. The use of antivirals as preventive weapon, for women not falling in the categories that require treatment, is recommended during the last trimester (until 4-12 wk after delivery) just in case of high viraemia (> 200000 UI/mL), but evidence collected so far is not solid. The drug of choice also in this case is TDF, although there is not direct or indirect proof of superiority over other NAs allowed in pregnancy; nevertheless, among them it shows the highest barrier to resistance. To date there are no data regarding the use of TAF in pregnancy, which could represent an important option, combining the same efficacy with a better safety profile.

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**Vertical transmission**

Intrauterine, perinatal and post-natal infection

Potential damage to the product of conception

**Maternal liver disease**

Acute or chronic hepatitis

Cirrhosis

**Hepatitis B in pregnancy**

**Figure 1 Hepatitis B in pregnancy: the two side of the problem.**

|  |  |
| --- | --- |
|  | Use of IFN in any case |
|  | Discontinuation, until delivery, of NA agent commenced before pregnancy in case of mild hepatitis |
|  | Use of LAM or TDF for severe acute hepatitis Use of TDF or for chronic hepatitis/cirrhosisSwitching to TDF if the women was before pregnancy on treatment with other drugs |

**Figure 2 Treatment of acute and chronic hepatitis B in pregnancy.** The column marked by the red circle refers to interventions that are not allowed. The column marked by the yellow circle refers to interventions that are backed up by low-quality or conflicting evidence. The column marked by the green circle refers to the best practice according to current evidence. IFN: Interferon; NA: Nucleos(t)ide analogue; LAM: Lamivudine; TDF: Tenofovir disoproxil fumarate.

|  |  |
| --- | --- |
|  | Avoiding lactation |
|  | Antiviral prohylaxis with TDF during the third trimester (until 4-12 wk after delivery) in case of maternal viral load > 200000 IU/mL C-section |
|  | HBIG and vaccination at birth as early as possible (in newborns) |

**Figure 3 Prevention of hepatitis B virus vertical transmission.** The column marked by the red circle refers to interventions that are not allowed. The column marked by the yellow circle refers to interventions that are backed up by low-quality or conflicting evidence. The column marked by the green circle refers to the best practice according to current evidence. HBV: Hepatitis B virus; TDF: Tenofovir disoproxil fumarate; IU: International units; C-section: Caesarean section; HBIG: Hepatitis B immunoglobulin.