Missed opportunities for prevention of perinatal transmission of hepatitis B: A retrospective cohort study

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BACKGROUND: Perinatal transmission of hepatitis B virus (HBV) can occur despite postexposure prophylaxis (PEP). Recent literature suggests that antiviral treatment during pregnancy when maternal HBV DNA levels are elevated can further decrease vertical transmission. However, HBV DNA screening is not routinely performed antenatally.

OBJECTIVE: To determine the rates of HBV prevalence and perinatal transmission in an antenatal cohort.

METHODS: A retrospective review of public health records (December 2008 to December 2010) was performed for both mothers and newborns.

RESULTS: A total of 725 mother-infant pairs were included. Of these, 574 of 715 (80%) women had antenatal hepatitis B e antigen (HBeAg) testing performed, and 127 of 574 (22%) were HBeAg positive (HBeAg+). Of babies born to hepatitis B surface antigen-positive (HBsAg+) mothers, only 573 of 725 (79%) received complete PEP. In addition, 172 of 725 (24%) infants did not receive post-PEP blood testing or were lost to follow-up. Of the 552 infants with results available, seven cases (1.3%) of mother-to-child HBV transmission were observed, six of which involved infants born to HBeAg+ women.

CONCLUSIONS: Our findings suggest that routine HBeAg screening could identify a subset of mother-infant pairs among HBsAg+ pregnant women who are at higher risk for vertical HBV transmission. Determination of viral load in expectant HBeAg+ mothers may provide more precise insight into HBV transmission to their infants.

Key Words: Hepatitis B; Pregnancy; Vertical transmission

Globally, 350 million people are chronically infected with hepatitis B virus (HBV) (1). The risk of developing chronic infection correlates inversely with the age at which HBV infection is acquired because the majority of those infected perinatally develop chronic infection. Chronic HBV infection is associated with liver-related diseases such as hepatocellular carcinoma (2). Over the past three decades, perinatal transmission of HBV has been dramatically reduced by the use of postexposure prophylaxis (PEP) (3). In British Columbia, PEP consists of hepatitis B immunoglobulin (HBIG) and vaccine administration to the newborn at birth, followed by three additional doses at two, four and six months of age (4).

With PEP, the rate of vertical transmission is reported to be 1% to 2% for quiescent chronic carrier mothers (hepatitis B surface antigen positive [HBsAg+]) but may be as high as 32% for mothers with active HBV infection (hepatitis B e antigen positive [HBeAg+]) (5).

The strongest predictor of vertical transmission is high levels of maternal serum HBV DNA. The presence of HBeAg generally correlates with high HBV DNA levels; however, HBeAg alone is not indicative of the magnitude of risk. For every log increase in HBV DNA level above 106 copies/mL, the risk of PEP failure increases (6). Missed opportunities exist worldwide to prevent unnecessary HBV transmission such as use of full implementation of PEP as well as consideration of antiviral therapy in pregnancy to reduce the level of HBV DNA.

Trials conducted in high HBV-endemic regions show that the use of antiviral medication in the last trimester of pregnancy, when maternal HBV DNA levels are >106 copies/mL, may reduce the rate of vertical transmission (7-10).

In the current study, we had three aims: to determine the prevalence of HBsAg and HBeAg among pregnant women in British Columbia; to determine the rate of perinatal HBV transmission in this
TABLE 1
Infant (n=725) hepatitis B surface antigen (HBsAg) status at eight to 12 months of age according to maternal (n=715) hepatitis B e antigen (HBeAg) status

<table>
<thead>
<tr>
<th>HBsAg</th>
<th>Maternal HBeAg status*</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBsAg-</td>
<td>337</td>
<td>100</td>
</tr>
<tr>
<td>HBsAg+</td>
<td>0</td>
<td>6</td>
</tr>
<tr>
<td>Unknown</td>
<td>110</td>
<td>21</td>
</tr>
<tr>
<td>Total</td>
<td>447</td>
<td>127</td>
</tr>
</tbody>
</table>

Data presented as n. *10 sets of twins within this sample. There were 725 mother-infant pairs, consisting of 715 women and 725 infants. Maternal data points for women with twins were counted twice. – Negative; + Positive

METHODS

Study design
A retrospective review of the treatment and follow-up of infants born to HBsAg+ women between December 2008 and December 2010 was undertaken. All pregnant women in two health authorities within British Columbia who tested and were reported to Public Health Services as HBsAg+ were approached to participate. Inclusion criteria included a live birth during the study period to 12 months postenrollment (to account for deliveries that occurred >40 weeks after enrollment).

Data collection
Data regarding HBV infection status were compiled from two public health data registries: the Primary Access Regional Information System, used by the Vancouver Coastal Health Authority, and the integrated Public Health Information System, used by the Fraser Health Authority. Data were cross-matched between public health records and the British Columbia Public Health Microbiology and Reference Laboratory at the British Columbia Centre for Disease Control, which perform reflex HBeAg testing on HBsAg+ prenatal specimens. The data extraction identified a cohort of women with confirmed HBV infection who were pregnant or became pregnant during the study period.

HBsAg+ pregnant women identified through the data registries were contacted antenatally by public health nurses and informed of their diagnosis, provided with recommendations for treatment of their infants and follow-up testing. The women’s care providers were also contacted by mail and informed of the risk of perinatal HBV transmission to the infant.

HBsAg+ infants born before February 2009, four doses at zero, two, four and six months of age were administered. Some infants (11.7%) did not complete their vaccination series, while another portion of approximately 20% did not receive the postbirth vaccination doses as per the provincial public health recommended schedule. Some infants (11.7%) did not complete their vaccination series, while another portion of infants (6.8%) received their immunoprophylaxis off schedule.

Data presentation
Vaccine(s) | Completion of vaccine schedule
--- | ---
HBsAg | Yes | No | Off-schedule* | Unknown
--- | --- | --- | --- | ---
HBV vaccine | 681 (93.9) | 31 (4.3) | – | 13 (1.8)
Complete | 698 (96.3) | 1 (0.1) | – | 26 (3.6)
series† | 573 (79.0) | 85 (11.7) | 49 (6.8) | 18 (2.5)

Data presented as n (%). *Defined as any dose that was administered at least ≥1 month(s) off the recommended schedule. †Defined as hepatitis B immunoglobulin (HBIG) and hepatitis B virus (HBV) vaccine administered at birth, followed by additional doses of HBV vaccine at two, four and six months of age. For at-risk infants born before February 2009, four doses at zero, two, four and six months age were administered.

The proportion of infants receiving HBV serological testing post-prophylaxis (eight to 12 months of age), and the rate of immunoprophylaxis failure, as indicated by infants’ HBsAg positivity despite PEP, were determined. Outcomes were stratified according to maternal HBsAg status.

RESULTS

Maternal HBV carrier rate
In total, 725 mother-infant pairs (10 sets of twins) were identified in whom maternal HBsAg screening was positive between December 2008 and December 2010. The total number of deliveries within the corresponding health regions in British Columbia during the same time frame was 55,734 (British Columbia Vital Statistics Agency, 2008, 2009), corresponding to an overall maternal HBV carrier rate of 1.3%. Of the 725 mother-infant pairs identified, 574 (79%) underwent maternal antenatal HBsAg testing. Of these, 127 (22.1%) women were HBsAg+. Follow-up infant testing results, performed to determine whether perinatal transmission had occurred, were not performed or not available in the public health databases for 173 (23.8%) infants.

Perinatal HBV transmission rate
Of the 552 infants who had available follow-up HBV testing results, perinatal transmission was observed in seven (1.3%) cases. In the groups of women who were HBsAg+ and HBsAg negative, vertical transmission rates were 5.5% (n=6) and 0% (n=0), respectively (Table 1). One transmission occurred in a mother who did not undergo HBsAg testing.

Infant vaccine history
Most infants received the first dose of HBV vaccination (96.3%) and HBsAg (93.9%) shortly after birth (Tables 2 and 3). However, approximately 20% did not receive the postbirth vaccination doses as per the provincial public health recommended schedule. Some infants (11.7%) did not complete their vaccination series, while another portion of infants (6.8%) received their immunoprophylaxis off schedule.

Clinical characteristics of HBV-positive infants
The clinical characteristics of the seven HBV-positive infants are described in Table 4. When the association of infant HBsAg positivity and PEP completion rates were examined, all seven infants who acquired HBV infection received vaccine, albeit two off-schedule. Of the seven infants who tested positive for HBsAg, one was documented to have anti-HBs levels that would have been considered immune in the absence of HBsAg. Of the seven HBsAg+ infants, breastfeeding data were available for five, all of whom were breastfed. Five were delivered vaginally and two by Caesarean section. Of the 545 HBsAg-negative infants, breastfeeding data were available for 304: 271 (89.1%) were breastfed, 31 (10%) were not breastfed and two (0.66%) received donor milk.
DISCUSSION

Main findings
Approximately 1.3% of the pregnant cohort studied was HBsAg+, of whom one in five were also HBeAg+. Of the babies born to the HBV-positive mothers, 20% did not receive PEP as per the recommended provincial public health schedule. In addition, almost one-quarter of infants did not undergo the recommended post-PEP serological testing at eight to 12 months of age to confirm PEP effectiveness.

Of the 552 infants who underwent post-PEP testing, seven cases of mother-to-child transmission (MTCT) were documented, corresponding to a 1.3% vertical transmission rate. This rate is lower than the perinatal HBV transmission rate of 2.1% that was reported in an adjacent Canadian province (Alberta) among a cohort of 980 exposed infants (11). Significantly, the maternal HBeAg was positive in six of the seven cases of perinatal transmission in the study, all of whom had completed PEP, suggesting that high maternal serum levels of HBV DNA was a likely significant factor in PEP failure. Because HBV viral loads were not captured in the databases and may not have been routinely ordered during the study period, these values were not available for analysis. The seventh case of transmission occurred from a mother with unknown HBeAg status.

Strengths and limitations
The present study represents an important evaluation of the efficacy of standardized HBV perinatal PEP in a jurisdiction with standardized guidelines and full medical coverage under these circumstances. It represents the two largest health authorities in British Columbia, which include a representative sample of provincial births because 63% occur within these regions (12). In addition, the ethnic composition within these regions is highly diverse, with 27% of the population identifying as Asian and 10% identifying as South Asian (13).

These data have several limitations. First, the present study was retrospective in design; however, the study reveals the current practice of maternal HBV testing, PEP receipt and follow-up testing in British Columbia, in the absence of observational bias. The data sets were incomplete and there were insufficient data available to fully assess the maternal and neonatal factors that may influence transmission. For example, we are unaware of whether any women were treated with antenatal antivirals in these pregnancies because this information was not collected. In addition, other potential risk factors, such as the duration of breastfeeding and whether it was supplemented with formula, the duration of labour, rupture of membranes or invasive fetal testing information was unavailable. Recent evidence suggests, however, that breastfeeding is not associated with a significant, incrementally increased risk for HBV vertical transmission among immunized infants (14).

Interpretation
Despite the introduction of PEP programs with proven efficacy for decreasing HBV transmission (15), almost 20% of infants in the present study did not complete their vaccination series, even though their parents and health care providers received specific PEP instructions as part of the public health program. Issues that may contribute to incomplete vaccine receipt include language barriers and family relocation, resulting in challenges for public health staff to assess vaccine uptake and subsequent infant testing results. Health care provider recommendation is associated with vaccine uptake (16). Because vaccination is the single most effective way of preventing HBV infection, programs designed to target patients with vaccine adherence challenges are paramount.

Canadian hepatology guidelines published in 2012 on the management of chronic hepatitis B recommend that all pregnant women who are HBsAg+ have HBV DNA levels determined (17). Higher rates of PEP failure occur with increasing HBV DNA levels (6). These guidelines also recommend that antiviral therapy be considered for pregnant women with HBV DNA viral loads >2x10^6 IU/mL, in an effort to prevent PEP failure and vertical transmission.

In the present study, six of seven transmissions that occurred were in women who tested positive for HBeAg. Had these women been treated with antenatal antiviral therapy, as suggested in several treatment trials (7-10), these neonatal infections may have been prevented.

Mutations in the HBsAg region can occur in vertical maternal transmission of HBV under the immunological selection pressure of PEP (18,19). One such transmission was observed in an Alberta cohort, in which maternal HBeAg and HBV DNA levels were undetectable in the first trimester (20). Of interest, and of potential concern, is the infant who was infected with HBV (as indicated by HBsAg seropositivity) despite a serum anti-HBs titre that was in the protective range (Table 4). Although molecular sequencing was not

## TABLE 3

<table>
<thead>
<tr>
<th>Infant HBsAg status</th>
<th>HBsAg</th>
<th>HBV vaccine at birth</th>
<th>Complete vaccine series</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Negative</td>
<td>513</td>
<td>25</td>
<td>527</td>
</tr>
<tr>
<td>Positive</td>
<td>7</td>
<td>0</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>520</td>
<td>25</td>
<td>543</td>
</tr>
</tbody>
</table>

Data presented as n. *Defined as any dose that was administered at least ≥1 month(s) off the recommended schedule; †Total are variable because not all infants had data available for all points of hepatitis B virus (HBV) immunoglobulin (HBsAg)/vaccine receipt in addition to infant hepatitis B surface antigen (HBsAg) testing.

## TABLE 4

<table>
<thead>
<tr>
<th>Case number</th>
<th>Year of birth</th>
<th>Maternal HBeAg status</th>
<th>HBsAg status</th>
<th>Anti-HBs</th>
<th>Adequate immunity</th>
<th>PEP receipt</th>
<th>Mode of delivery</th>
<th>Breastfeeding*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2008</td>
<td>Positive</td>
<td>Positive</td>
<td>Negative</td>
<td>Not immune</td>
<td>Off-schedule</td>
<td>Vaginal (forceps)</td>
<td>Partial</td>
</tr>
<tr>
<td>2</td>
<td>2008</td>
<td>Positive</td>
<td>Positive</td>
<td>Negative</td>
<td>Not immune</td>
<td>Complete</td>
<td>Vaginal (vacuum)</td>
<td>Exclusive</td>
</tr>
<tr>
<td>3</td>
<td>2008</td>
<td>Positive</td>
<td>Positive</td>
<td>Negative</td>
<td>Not immune</td>
<td>Complete</td>
<td>Vaginal (SVD)</td>
<td>Exclusive</td>
</tr>
<tr>
<td>4</td>
<td>2009</td>
<td>Positive</td>
<td>Positive</td>
<td>Positive</td>
<td>Immune</td>
<td>Off-schedule</td>
<td>Caesarean section</td>
<td>Unknown</td>
</tr>
<tr>
<td>5</td>
<td>2009</td>
<td>Positive</td>
<td>Positive</td>
<td>Negative</td>
<td>Not immune</td>
<td>Complete</td>
<td>Vaginal (SVD)</td>
<td>Unknown</td>
</tr>
<tr>
<td>6</td>
<td>2009</td>
<td>Unknown</td>
<td>Positive</td>
<td>Negative</td>
<td>Not immune</td>
<td>Complete</td>
<td>Vaginal (SVD)</td>
<td>Exclusive</td>
</tr>
<tr>
<td>7</td>
<td>2010</td>
<td>Positive</td>
<td>Positive</td>
<td>Negative</td>
<td>Not immune</td>
<td>Complete</td>
<td>Caesarean section (elective)</td>
<td>Partial</td>
</tr>
</tbody>
</table>

*Defined as any dose that was administered at least ≥1 month(s) off the recommended schedule. HBeAg Hepatitis B e antigen; HBs Hepatitis B surface antigen; PEP Postexposure prophylaxis; SVD Spontaneous vaginal delivery.
performed in this particular case, it is possible that a similar phenomenon may have contributed to a possible explanation for PEP prophylaxis failure in this instance. This phenomenon, although uncommonly documented, provides additional rationale for identifying HBV-infected mothers at higher risk for vertical HBV transmission and administering selective antiviral treatment in those cases, in addition to the currently recommended PEP and post-PEP serological assessment.

CONCLUSION
A significant proportion of infants in British Columbia born to HBsAg+ mothers are not receiving a full course of recommended PEP and/or have incomplete follow-up. Moreover, MTCT of HBV still occurs despite complete, timely HBV PEP, which particularly affects infants whose mothers are HBeAg+. The present study does not add any new data to what is known about HBV transmission; however, our findings highlight that adherence to PEP clinical guidelines in British Columbia is suboptimal and needs improvement, and may be relevant to other Canadian jurisdictions facing similar challenges.

While a growing body of evidence indicates that maternal antiviral treatment can abrogate HBV vertical transmission in maternal HBeAg+ cases, investigations on maternal and fetal antiviral toxicities, rebound hepatitis after treatment cessation, safety of breastfeeding during treatment and the treatment-associated emergence of viral drug resistance are still needed. The optimal timing for HBeAg or HBV DNA testing in pregnancy in relation to risk determination for MTCT remains uncertain. Future studies may aim to identify modifiable perinatal risk factors in HBsAg+ women to further reduce the risk of HBV vertical transmission and the global burden of HBV disease.

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