Is occult Hepatitis B virus infection with detectable anti-HBs infectious or not?

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In the recent issue of Middle East Journal of Digestive Diseases, we read with great interest the article by Bahari et al.1, “Significance of response to hepatitis B vaccine in subjects with isolated antibody to hepatitis B core antigen”. We wanted to specify some important matters related to the article.

Initially, definition of occult hepatitis B virus infection is the existence of hepatitis B virus (HBV) deoxyribonucleic acid (DNA) in the liver of individuals negative for hepatitis B surface antigen (HBsAg) by currently available assays.2 Serum HBV DNA titres may be detectable (usually very low, <200 IU/ml) or undetectable, and may show fluctuation in occult hepatitis B virus infection patients. Also, occult hepatitis B virus infection may occur not only seropositive for antibodies to hepatitis B core antigen (anti-HBc) and/or anti-bodies to HBsAg (anti-HBs), but also seronegative.2

Isolated anti-HBc positivity or anti-HBc alone, most commonly associated with occult hepatitis B virus infection, develops particularly because of the reduction of corresponding antibodies to undetectable levels after the resolution of infection.3 In this case, with HBV vaccination as implemented in the article, anti-HBs levels increase in a short time.1 However, isolated anti-HBc positivity can be seen in the beginning of the convalescence period of current infection or with the reduction of HBsAg to undetectable levels after long-term inactive HBV carriage. The reduction of HBsAg levels can be shown by only high sensitive HBsAg assays and may be with low HBV DNA titres. Moreover, false isolated anti-HBc positivity has been reported up to 35% via repetitive tests.4,5

The more important topic is whether occult hepatitis B virus infection with detectable anti-HBs is infectious or not. Theoretically, high anti-HBs titres can neutralize the infectivity of virus particles, on the other hand, infection can emerge if the product to be transfused is positive for HBV DNA. HBV transmission from patients with occult hepatitis B virus infection with anti-HBs positivity to immunocompetent recipients has been shown.6 In addition, it has also been reported that anti-HBs titres lower than 100 IU/ml have insufficient protectivity in the presence of HBV DNA and products containing HBV DNA may be infectious and may cause serious clinical status in patients with severe immun deficiency even though anti-HBs levels are high.6 Anti-HBc positive donations are accepted only if HBV DNA is negative and anti-HBs titres are at least 100 IU/ml in European Union countries and at least 200 IU/ml in Japan.7,8

Consequently, serological tests and vaccination of solely anti-HBc donors

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might be an option or tool for donor screening, but a combination of serological tests with nucleic acid amplification tests (individuate or at least pooled) seems more reasonable and reliable for the reasons mentioned above (e.g., seronegative occult hepatitis B virus infection, isolated anti-HBc positivity due to the reduction of HBsAg titres with low HBV DNA levels, donor loss because of the false isolated anti-HBc positivity and infectivity of occult hepatitis B virus infection with detectable anti-HBs). However, the costs, the need for equipment, and experienced staff or accessibility may be the problems as shared by Bahari et al.1

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CONFLICT OF INTEREST
The author declares no conflict of interest related to this work.

Answer to Dr. Murat Afyon
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We thank the author of the letter gratefully for his/her attention and the valuable comments. Surely, we agree with the comments that had been mentioned in the letter. However, the design of our study pointed to somewhat a different issue. We had aimed at investigating the pattern of response to hepatitis B vaccine in those who had isolated HBC antibodies and to correlate or compare the results with PCR test in order to identify patients with occult HBV infection.

As the author of the letter has mentioned, in occult HBV infection, serum HBV DNA titers may be detectable or even undetectable. Also, occult HBV infection may occur not only seropositive for antibodies to hepatitis B core antigen (anti-HBc) and/or antibodies to HBsAg (anti-HBs), but also seronegative. Therefore, the diagnosis of occult HBV infection would be difficult in patients with both negative HBV DNA titers and negative HBV antigens/antibodies. As we have shown in our article, 19 (21.1%) of the cases had no sero-conversion (anti-HBs titers <10 mIU/mL) 30 days after the third dose, although they had negative PCR results. Accordingly, using vaccination may help to distinguish patients with occult HBV infection among those who have both negative HBV DNA titers and negative HBV antigens and antibodies.

On the other hand, studies have shown that even in the presence of high anti-HBs titers, hepatitis B reactivation occurred after immunosuppressive therapy in anti-HBc positive patients. Therefore, anti-HBs titers with high titer may not be reliable enough to donate blood safely.

All together, combination of hepatitis B vaccination and HBV DNA PCR may help to exclude patients with occult HBV infection, but the risk of hepatitis B transmission will not be zero.

REFERENCES


