

## LETTERS TO THE EDITOR

# Validation of Whole Blood Rapid Diagnosis Test for Hepatitis B

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We recently published a report validating point-of-care rapid diagnostic tests (RDT) for the diagnosis of Hepatitis B virus (HBV) infection in serum. In the current report, we validated a whole-blood RDT for HBsAg in the form of a test-strip. The test was validated in 55 HBV positive individuals across all genotypes other than F, and in 20 HBV negative individuals in the Netherlands. The RDT showed 100% sensitivity and specificity. The low cost and use in whole blood allows this RDT to be useful in resource-limited locations, further validation in such settings will be of importance.

To the editor:

We recently published a report validating point-of-care rapid diagnostic tests (RDT) for the diagnosis of Hepatitis B virus (HBV) infection [1]. We believe that expedited and cost-effective diagnosis of HBV infection via RDT is critical to understand the epidemiology of HBV in resource-limited settings and implement prevention and treatment strategies [2]. In that report, a whole-blood rapid test for HBV surface antigen (HBsAg), the universally accepted test to diagnose chronic HBV infection, in the form of “cassette” did not produce reliable results, with a sensitivity of only 56% (albeit a specificity of 100%). In the current report, we validated a whole-blood RDT for HBsAg in the form of a test-strip (PRECHECK Bio Inc, Korea). The test was validated in 55 HBV positive (HBV-pos) and 20 HBV negative (HBV-neg) individuals in the Netherlands. The median age was 38 years (IQR 3–47) in the HBV-pos group, with 60% being male and 33 years (IQR 30–37) in the HBV-neg group, with 85% being male. The RDT showed 100% sensitivity and specificity, with complete correlation for HBsAg positive and negative results with the local gold standard at Erasmus University, Rotterdam (LIAISON XL, Diaorin, Italy). All HBV-pos individuals were negative for hepatitis C and D virus, as well as human immunodeficiency virus. The samples included individuals on treatment and inactive carriers, all HBV genotypes other than F, with the most common genotypes being D (23%), C (20%), and A (15%). Median HBV DNA was 20 IU/ml (IQR 20–285), median HBsAg level 510 IU/ml (IQR 150–2900), and 16 subjects (29%) were HBeAg positive. Seventy percent of HBV+ individuals were on treatment (47% on entecavir and 21% on tenofovir).

We believe this addition to our previous study is important as it provides validation of a RDT for HBsAg using

whole blood, in a rapid and cost-effective manner, therefore allowing the test to be used without the need of any laboratory resources. Our test was validated in one institution (Erasmus MC, the Netherlands) in a variety of genotypes. However, we believe further validation in resource-constrained settings will be of importance.

### Data Accessibility Statement

Study data will be made available upon request of the corresponding author.

### Ethics and Consent

Ethical approval was given by the ethical committee of the Erasmus MC, Rotterdam. MEC-2017\_1140.

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### Competing Interests

The authors have no competing interests to declare.

### Author Contributions

Jose D. Debes: study design, data analysis, writing; Gertine van Oord: data collection, writing; Andre Boonstra: study design, data analysis, writing.

### References

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2. **Amini A, Varsaneux O, Kelly H, et al.** Diagnostic accuracy of tests to detect hepatitis B surface antigen: A systematic review of the literature and meta-analysis. *BMC Infect Dis*. 2017; 17: 698. DOI: <https://doi.org/10.1186/s12879-017-2772-3>

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