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9 **Clinical utility of quantifying hepatitis B surface antigen in African patients with chronic**  
10 **hepatitis B**11 Gerrit Post<sup>1</sup>, Jess Howell<sup>2</sup>, Amina Sow<sup>3</sup>, Gibril Ndow<sup>4</sup>, Isabelle Chemin<sup>5</sup>, Gora Lo<sup>3</sup>, Amie  
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69 **Abstract (n=250 words)**

70 The clinical utility of quantifying hepatitis B surface antigen (qHBsAg) levels in African subjects  
71 with chronic hepatitis B virus (HBV) infection has been poorly documented. From a multicenter  
72 cohort of 944 HBV-infected African patients we aimed to assess whether qHBsAg alone can  
73 accurately identify i) those in a HBeAg-negative chronic HBV infection phase at low risk of liver  
74 disease progression and ii) those in need of antiviral therapy according to the 2017 EASL  
75 guidelines. We analyzed 770 HBV mono-infected treatment-naïve patients, mainly males (61%)  
76 from West Africa (92%), median age 35 years (IQR: 30-44), median HBV DNA: 95.6 IU/ml  
77 (10.0-1,300.0), median qHBsAg 5,498 IU/ml (1,171-13,000), HBeAg-pos 38 (5%). A total of  
78 464/770 (60.2%) patients were classified as HBeAg-negative chronic infection (median age 36  
79 years (31-46), median ALT 23 IU/l (18-28), median HBV-DNA 33.5 IU/ml (3.8-154.1), median  
80 LSM 4.8 kPa (4.1-5.8)) and qHBsAg levels had poor accuracy to identify these subjects with  
81 an AUROC at 0.58 (95%CI: 0.54-0.62), sensitivity 55.0%, specificity 55.6%; 118/770 (15.3%)  
82 patients were eligible for treatment according to the 2017 EASL criteria. qHBsAg correlated  
83 poorly with HBV DNA and had poor accuracy to select patients for antiviral therapy with an  
84 AUROC at 0.54 (0.49-0.60), sensitivity 46.6%, specificity, 46.9%. In African treatment-naïve  
85 HBV-infected subjects, the clinical utility of qHBsAg to identify subjects in HBeAg-negative  
86 infection phase or subjects eligible for antiviral therapy seems futile. Whether qHBsAg levels  
87 can be used as a predictor of long-term liver complications in Africa needs to be further  
88 investigated.

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93 Infection with hepatitis B virus (HBV) is a major public health issue worldwide. Sub-Saharan  
94 Africa accounts for an estimated 60 million people chronically infected with HBV (1). However,  
95 access to HBV diagnosis and treatment is limited in most African countries with only 2% of  
96 HBV-infected people diagnosed and less than 1% of those eligible for antiviral therapy  
97 receiving treatment in 2016 in sub-Saharan Africa (2). In patients with chronic HBV infection,  
98 the assessment of liver disease and initiation of antiviral therapy mainly rely on three

99 measurements: alanine aminotransferase (ALT) levels, HBV viral load, and liver fibrosis  
100 estimation using liver histology or liver stiffness measurement (e.g. by vibration-controlled  
101 transient elastography [Fibroscan®]) (3). Except for ALT levels, these tests are difficult to  
102 perform in clinical practice in Africa and are therefore major barriers to scale-up HBV screen-  
103 and-treat interventions in this region. Indeed, measurement of HBV viral load currently relies  
104 on nucleic acid testing using quantitative real-time polymerase chain reaction (qRT-PCR),  
105 which is expensive (up to €150 in Africa) and requires high quality laboratories and well-trained  
106 technicians. In addition, Fibroscan® devices remain expensive and are often only accessible in  
107 capital cities in private sectors (4). To overcome these limitations, simplified diagnostic tools  
108 and algorithms are needed. For liver fibrosis assessment, biochemical scores (e.g aspartate-  
109 aminotransferase (AST)-to-platelet ratio index (APRI) or gamma glutamyl-transpeptidase to  
110 platelet ratio (GPR)) have been proposed but their diagnostic accuracy is debated in African  
111 populations (5, 6). To quantify HBV viral replication, new molecular technologies (e.g  
112 GeneXpert) (7) or serological biomarkers (e.g Hepatitis B core-related antigen (HBcrAg)) (8)  
113 have been validated as accurate alternatives to the conventional qPCR, but they are still  
114 difficult to implement at large scale in resource-limited countries.

115 HBV-infected subjects in the HBeAg-negative chronic infection phase (previously known as  
116 chronic inactive carriers) should be at low risk of liver disease progression (9). Therefore, in  
117 Africa, where a large proportion of HBV-infected people are in this phase (10) a simple  
118 biomarker to easily identify these subjects and those in need of antiviral therapy would be  
119 extremely useful to scale-up HBV screen-and-treat interventions.

120 In the last decade the quantification of serum HBsAg (qHBsAg) levels, which reflects the  
121 transcriptional activity of covalently closed circular DNA (cccDNA), has been used routinely in  
122 high-income countries, to define the phase of the infection and predict liver-related  
123 complications in patients with chronic HBV infection (11, 12). Levels of qHBsAg have been  
124 also identified as a useful marker to monitor response to treatment and predict relapse after  
125 stopping nucleos(t)ide analogue therapy (13). Although qHBsAg levels is not recommended to  
126 select patients for antiviral therapy, its clinical utility to predict or monitor response to treatment  
127 is mentioned in international HBV guidelines (3). HBsAg quantification is a simple and more  
128 affordable test (less than €2 per test) than qPCR.

129 Data on qHBsAg levels in HBV-infected patients has been mainly collected in Western (13-15)  
130 and Asian countries (16-19). In African HBV-infected subjects, qHBsAg levels and its clinical  
131 utility have been very inadequately documented (20, 21) with conflicting results. In particular,

132 whether it could simplify the identification of subjects in an HBeAg-negative chronic infection  
133 phase and of patients in need of antiviral treatment in Africa is unknown.

134 We hypothesized that qHBsAg levels could be used in clinical routine to easily distinguish  
135 patients in need of antiviral therapy from those in an HBeAg-negative chronic infection phase  
136 considered at low risk of liver disease complications.

137 This study aimed to assess whether HBsAg levels alone is accurate to 1) identify subjects in a  
138 HBeAg-negative chronic HBV infection phase and 2) select subjects for antiviral therapy  
139 amongst treatment-naïve HBV-infected people in Africa.

140

## 141 **Methods**

### 142 **Study population**

143 Data from treatment-naïve, HBV-infected African patients, enrolled in African (Burkina  
144 Faso/The Gambia/Senegal) and European (France/Germany) cohorts, were retrospectively  
145 analyzed. Data included basic demographic information (age, gender, excessive alcohol  
146 intake, country of birth), as well as laboratory data (HBV viral load, qHBsAg levels, Hepatitis B  
147 e antigen (HBeAg) serology (ETI-EBK Plus, Diasorin, Italy or Abbott Diagnostics, Chicago, IL,  
148 USA), liver enzyme levels (ALT, AST, GGT, platelet count, and co-infection sero-status (HIV),  
149 hepatitis C virus (HCV), hepatitis Delta virus (HDV)). Patients with excessive alcohol intake as  
150 defined by intake >20g/day or HIV, HCV or HDV co-infections were excluded from the final  
151 analysis.

### 152 **Laboratory investigations**

#### 153 **HBV DNA measurement**

154 HBV DNA levels were measured by qPCR using commercialized assay (Abbott Diagnostics,  
155 Chicago, IL, USA) in all countries except The Gambia where a validated in-house qPCR was  
156 used (22).

#### 157 **HBs antigen quantification**

158 qHBsAg levels were measured in all patients using the Abbott Architect chemiluminescent  
159 microparticle assay (Abbott Diagnostics, Chicago, IL, USA). Samples were tested at dilution  
160 1/500 as recommended with a range of quantification of 25-125,000 IU/ml after dilution. We  
161 also calculated the qHBsAg/HBV DNA ratio that reflects the relationship between HBsAg  
162 production and HBV replication and might be a better marker of viral activity than HBV viral  
163 load or qHBsAg levels alone (23).

#### 164 **HBV genotyping**

165 HBV genotyping was determined in a subgroup of patients. PCR and nested PCR products  
166 were sent to Genome Express (Grenoble, France) for sequencing. Edited sequences were  
167 submitted to BLAST analysis, and the highest scoring complete HBV genome was retained. A  
168 phylogenetic tree was built from >2000 complete HBV genome sequences contained in  
169 GenBank, and HBV genotype/subgenotype clades were identified based on information in the  
170 GenBank entries and/or in original publications.

#### 171 **Liver fibrosis assessment**

172 The severity of liver fibrosis was assessed using fasting LSM in all patients. As recommended,  
173 a valid Fibroscan® (Echosens, France) value was defined as at least 10 valid measurements,  
174 a success rate of at least 60% and an IQR/median-ratio of less than 30% (24). To stage the  
175 degree of liver fibrosis, we used previously validated LSM cut-offs in African patients with  
176 chronic HBV infection:  $\geq 7.9$  kPa for clinically significant fibrosis ( $\geq F2$ ) and  $\geq 9.5$  kPa for  
177 cirrhosis (6). In 88 patients liver biopsies were carried out and fibrosis grading was reported in  
178 METAVIR score.

#### 179 **HBV infection phases**

180 We determined the phase of HBV infection according to the EASL 2017 guidelines using a  
181 single time point analysis (supplemental table 1). Subjects in the HBeAg-negative HBV  
182 infection phase were defined as having a negative HBeAg serology and HBV DNA  $< 2.000$   
183 IU/ml and ALT  $< 40$  IU/L and none or mild liver fibrosis based on LSM ( $< 7.9$  kPa) or liver  
184 histology analysis if available (F0-F1) and no or mild activity (A0-A1).

#### 185 **Hepatitis B treatment eligibility**

186 We applied the 2017 EASL treatment criteria based on a single time point as usually done in  
187 resource-limited areas. We applied the upper limit of normal for ALT as 40 IU/L irrespective of  
188 gender as recommended by EASL (3) (Suppl. Table 2).

#### 189 **Statistical analysis**

190 Statistical analyses were performed using IBM SPSS Statistics version 24. We report means  
191 and standard deviation for all metric and normally distributed variables. If normal distribution  
192 could not be assumed, median and interquartile ranges (IQR) are presented. Since none of  
193 our variables shows a normal distribution, we used Mann-Whitney-U and Kruskal Wallis tests  
194 as well as Spearman's correlation coefficient to compare or correlate different variables. A two-  
195 sided P value of less than 0.05 was considered statistically significant.

196 The capability of qHBsAg levels to correctly identify patients with HBeAg-negative chronic  
197 infection, as well as to select patients for antiviral treatment, were evaluated by the receiver

198 operating characteristic (ROC) curve. The optimal cut-offs for HBsAg levels were selected to  
199 minimize the absolute difference between the sensitivity and specificity.

## 200 **Results**

### 201 **Study population**

202 We extracted data from 944 treatment-naïve African patients with chronic HBV infection and  
203 available qHBsAg levels. Most of them were recruited in West Africa (n=882) through the  
204 PROLIFICA research program (10, 25) in The Gambia and Senegal (n=689). A minority of  
205 African patients (n=149) was enrolled in European cohorts. Finally, complete data from 770  
206 treatment-naïve, HBV mono-infected African patients were analyzed (Figure 1). Table 1  
207 summarizes the characteristics of the study population according to the different phase of HBV  
208 infection.

209 All patients were born in Africa and were mainly from West Africa (n=711, 92.3%), a minority of  
210 patients was from North Africa (n=14, 1.8%) and Central or East Africa (n=45, 5.8%). Most of  
211 them were males (n=468, 61%), median age of 35 years (IQR: 30-44), 38 (4.9%) subjects  
212 were tested positive for HBeAg, median LSM was 5.3 kPa (4.4 – 7.0) with 133 patients having  
213 significant liver fibrosis (7.9%) or cirrhosis (9.4%), median ALT level was 25 IU/L (20 - 36) and  
214 median HBV DNA level was 95.6 IU/ml (10.0–1,300.00) with a median qHBsAg levels of  
215 5,497.6 IU/ml (1,170.8 – 13,000.0). HBV genotype was determined in a subgroup of 202  
216 patients: Genotype E was predominant (178 (88.1%)) whilst a minority of patients were  
217 infected with genotype A (24 (11.9%)).

218

219 The HBsAg levels were higher in HBeAg-positive patients compared to HBeAg-negative  
220 patients (10,407.5 IU/ml (2,582.5 – 33,320.0) vs 5,343.4 IU/ml (1,142.1 – 13,000.0), p=0.003).  
221 HBsAg levels were higher in patients classified with HBeAg-negative chronic hepatitis than in  
222 those with HBeAg-negative chronic infection (7,025.39 IU/ml (2,305.71 – 14,694.00) vs 4,526.1  
223 IU/ml (460.2 – 13,000.0), p<0.001) (Table 1).

224 qHBsAg to HBV DNA ratio also differed according to HBeAg sero-status (0.01 (0.00 – 0.18) vs.  
225 17.3 (0.6 – 268.0, p<0.001). HBV DNA ratio was the highest in patients with HBeAg-negative  
226 chronic infection 60.8 (4.1 – 561.3) and higher than that observed in HBeAg-negative chronic  
227 hepatitis subjects 1.37 (0.14 - 24.72), p<0.001.

### 228 **Correlation between HBsAg levels and HBV viral load**

229 In the whole study population, including both HBeAg-positive and HBeAg-negative subjects,  
230 there was a poor correlation between qHBsAg levels and HBV DNA levels (r=0.270, p<0.001).

231 In a subgroup of HBeAg-positive patients the correlation was higher as compared to the

232 correlation observed in HBeAg-negative patients ( $r=0.565$  and  $r=0.252$ ,  $p<0.001$  respectively)  
233 (Figure 2). In a subset of patients with available HBV genotype determination ( $n=202$ ), the  
234 correlation of qHBsAg with HBV DNA was poor even after stratifying by the genotypes  
235 ( $r=0.090$ ,  $p=0.234$  in genotype E and  $r=-0.043$ ,  $p=0.843$  in genotype A).

#### 236 **Performance of qHBsAg to identify patients with HBeAg-negative chronic infection.**

237 In the entire study population 464 (60.3%) subjects were identified as having HBeAg-negative  
238 chronic infection (Table 1). The performance of qHBsAg levels was poor to identify these  
239 subjects with an AUROC at 0.58 (0.54 – 0.62), sensitivity 55.0%, specificity 55.6% for a best-  
240 identified threshold of 5,660 IU/ml (Table 2).

#### 241 **Performance of qHBsAg levels to predict HBV treatment eligibility**

242 Applying the 2017 EASL guidelines, 118 (15.3%) were eligible for antiviral therapy. Using the  
243 2017 EASL criteria as a reference, qHBsAg levels failed to correctly identify patients eligible for  
244 treatment with an AUROC at 0.54 (0.49 – 0.60) and low sensitivity and specificity at 46.6% and  
245 46.9% respectively, with a best identified cut-off of 6,000 IU/ml (Table 2).

246 Our results did not differ between patients living in Africa and those living in Europe and were  
247 similar irrespective of the use of commercial or in-house HBV PCR (data not shown).

#### 248 **Discussion**

249 In a large cohort of African treatment-naïve patients with chronic HBV infection, we found that  
250 qHBsAg levels had a poor clinical utility to identify HBV-infected subjects with HBeAg-negative  
251 chronic HBV infection or to select HBV-infected subjects for antiviral therapy in Africa.

252 In most African countries, hepatitis B screening interventions are usually based on only the  
253 detection of HBsAg using a rapid POC test after finger prick, and people have a significantly  
254 limited access to HBV DNA measurement. We could not confirm our hypothesis that qHBsAg  
255 levels could facilitate a rapid triage of HBV-infected patients.

256 To the best of our knowledge the utility of qHBsAg has never been examined in HBV-infected  
257 African patients, so far. To date, only two studies analyzed the levels of qHBsAg in African  
258 subjects; one study from Senegal assessed the fluctuation of qHBsAg levels in 87 HBV-  
259 infected patients with normal ALT level (20) but did not examine whether qHBsAg levels could  
260 help classifying the phase of natural history of chronic HBV infection; another study conducted  
261 in the UK assessed the qHBsAg levels in 259 Genotype E patients born in West Africa, but  
262 only included patients with a viral load above 2,000 IU/ml and raised ALT level (21).

263 As previously shown in Asian (19) and European (15) patients, our study found that qHBsAg  
264 levels varied according to the positivity of HBeAg in African HBV-infected patients with the  
265 lowest levels observed in HBeAg-negative patients and the highest levels measured in HBeAg-

266 positive patients. We also found that qHBsAg to HBV DNA ratio was the highest in subjects  
267 with HBeAg-negative chronic infection as previously suggested (14, 26). In this phase of the  
268 infection, HBsAg is transcribed from integrated DNA while cccDNA transcription is inhibited in  
269 this phase (26-29).

270 As observed in other studies (19-21, 29), we found a poor correlation between qHBsAg levels  
271 and HBV viral load, especially in HBeAg-negative subjects. In West African HBV-infected  
272 patients with viral load  $\geq 2,000$  IU/ml and raised ALT level, Chakrabarty et al. found lower  
273 qHBsAg levels in patients with advanced liver fibrosis (21). We also found low level of HBsAg  
274 in subjects with cirrhosis but no correlation was observed between LSM and qHBsAg levels in  
275 our study (data not shown).

276 We assessed the accuracy of qHBsAg levels for the identification of HBeAg-negative subjects  
277 with chronic infection. From a public health perspective, the validation of a simple marker to  
278 easily identify patients in this phase is highly needed in Africa. Indeed, the vast majority of  
279 HBV-infected subjects in Africa are classified in this phase (10) and might be at low risk of liver  
280 disease progression (9) suggesting a possible cost-effective one-step liver assessment with  
281 major cost savings for resource-limited African countries. Unfortunately, in contrast to previous  
282 findings (14, 30), qHBsAg levels failed in our study to accurately identify subjects in this phase.  
283 This could be explained by relatively high levels of qHBsAg levels of subjects with HBeAg-  
284 negative chronic infection in our study (median 4526.1 IU/ml (460.2 – 13,000) despite very low  
285 median HBV viral load 33.50 IU/ml (3.79 - 154.07)) as reported in previous African studies (20,  
286 21). Asian and Western studies (14, 16) reported much lower qHBsAg levels in their population  
287 of inactive chronic carriers. Indeed, qHBsAg levels do not reflect the production of virions but  
288 rather the presence of defective HBsAg particles (spheres and filaments) that can dramatically  
289 exceed the amount of competent infectious virions by  $10^2$  to  $10^5$ . Whether the high levels of  
290 qHBsAg observed in African subjects in the HBeAg-negative chronic infection phase is  
291 attributable to genotypes or genetic variability of HBV, or clinical outcomes in Africa, remains to  
292 be confirmed.

293 Using the 2017 EASL treatment criteria, we also confirmed the poor performance of qHBsAg  
294 levels for identifying subjects in need of antiviral therapy. From a public health perspective the  
295 identification of patients in immediate need of antiviral therapy based on a one-step screening  
296 intervention would be very useful in resource-constrained areas in Africa.

297 Our study has some limitations. First, patients were assessed on a single time point; however,  
298 most patients with HBV infection living in resource-limited countries have a single assessment;  
299 we were unable to provide longitudinal data at this stage. Therefore, we did not assess the

	Full study population n=770	HBeAg- CI n=464	HBeAg- CH N=206	HBeAg+ CI n=13	HBeAg+ CH n=15	Cirrhosis n=72	P value*
Age, years	35 (30 - 44)	36 (31 - 46)	35 (29 - 40)	32 (28 - 38)	32 (28 - 37)	36 (30 - 42)	P=0.004
Males, n	468 (60.8%)	238 (51.3 %)	148 (71.8 %)	8 (61.5 %)	12 (80.0 %)	62 (86.1 %)	p<0.001
West African, n	711 (92.3%)	448 (96.6 %)	169 (82 %)	12 (92.3 %)	10 (66.7 %)	72 (100 %)	p<0.001
East African, n	45 (5.8 %)	14 (3.0 %)	29 (14.1 %)	1 (7.7 %)	1 (6.7 %)	0	p<0.001
North African, n	14 (1.8 %)	2 (0.4 %)	8 (3.9 %)	0	4 (26.7 %)	0	p<0.001

300 utility of qHBsAg levels to predict the development of liver disease complications over time. We  
301 will however address this question in the future. Secondly, our study population was mainly  
302 from West Africa and therefore the proportion of non-E genotype was small. Thirdly, we used  
303 two different methods for HBV DNA quantification; however, our findings were similar  
304 irrespective of the use of a commercial or in-house PCR. Fourthly, we did not analyze the  
305 impact of pre-core and pre-S mutations on the qHBsAg levels although these mutations may  
306 influence the level of HBsAg and contribute to liver disease complications (17, 31)

307 In conclusion, although the quantification of HBsAg levels is a simple and inexpensive test its  
308 clinical utility to identify subjects at low risk of liver disease progression (HBeAg-negative  
309 chronic infection phase) or subjects eligible for immediate antiviral treatment in Africa is poor.  
310 Additional markers and strategies are needed to simplify the stratification of HBV-infected  
311 subjects in order to scale up screen-and-treat interventions in Africa.

312

313 **Authors' contribution:** ML,PI,GP,YS,JH,MT designed the study. GP,ML,YS were in charge of  
314 the statistical analysis. AS,JH,AC,GL,IC,CTK were in charge of the virological analysis.  
315 ML,GP,PI,YS drafted the first manuscript. All authors contributed to patient recruitment and  
316 approved the manuscript.

317 **Conflicts of interest:** ML, PI, YS, MT received research funding from Gilead US.

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321 and their families.

322 **Data availability statement:** The data that support the findings of this study are available from  
323 the corresponding author upon reasonable request.

324

325

326 **Tables:**

327

LSM, kPa	5.3 (4.4 – 7.0)	4.8 (4.1 – 5.8)	5.9 (4.8 – 8.0)	5.3 (3.3 - 6.5)	6.6 (5.9 - 7.8)	12.45 (10.7 - 17.6)	p<0.001
HBV DNA (IU/ml)	95.64 (9.98 – 1300.00)	33.50 (3.79 - 154.07)	2781.11 (110.06 - 20804.00)	7801425.18 (330451.31 – 129130845.8)	632.4 (4.29 - 584370.55)	2124.03 (164.57 – 69870.67)	p<0.001
qHBsAg (IU/ml)	5497.62 (1170.77 – 13000.0)	4526.1 (460.2 – 13000)	7025.39 (2305.71 – 14694.00)	33320.00 (13196.00 – 52000.00)	7771.00 (574.30 – 27267.28)	4720.8 (1204.94 – 7839.58)	p<0.001
qHBsAg to HBV DNA ratio	13.26 (0.36 – 238.81)	60.8 (4.1 – 561.3)	1.37 (0.14 - 24.72)	0 (0 - 0.01)	0.21 (0.01 - 22.36)	1.09 (0.04 – 10.67)	p<0.001
HBV Genotype, n, m=202							
E	178 (88.1 %)	106 (89.8 %)	41 (85.4 %)	5 (83.3 %)	3 (100 %)	23 (85.2 %)	p=0.186
A	24 (11.9 %)	12 (10.2 %)	7 (14.6%)	1 (16.7 %)	0	4 (14.8 %)	p=0.511
ALT, IU/l	25 (20 – 36)	23 (18 - 28)	43 (25 - 58)	32 (24 – 33)	46 (38 – 67)	35 (25 - 57)	p<0.001
AST, IU/l, m=765	30 (25 - 37)	27 (23 - 32)	37 (29 - 48)	34 (28 – 39)	40 (34 – 55)	37 (31 – 65)	p<0.001
GGT, IU/l, m=723	26 (20 - 38)	24 (18 - 31)	33 (24 - 49)	34 (20 – 46)	28 (19 – 56)	38 (26 – 72)	p<0.001
Platelet count, 10 <sup>9</sup> /L , m=731	205(163 – 251)	212 (171 - 257)	194 (158 -246)	243 (195 – 273)	190 (137 – 235)	147 (123 – 209)	p<0.001
2017 EASL treatment eligibility, n	118 (15.3 %)	5 (1.1 %)†	29 (14.1 %)	8 (61.5 %)	4 (26.7 %)	72 (100%)	p<0.001

328 **Table 1.** Characteristics of the study population according to the HBV infection phases. \*based  
329 on either Kruskal Wallis or Chi-squared test. Data are presented as median (IQR) or n (%). †  
330 all 5 patients had a family history of HCC.

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332 HBeAg- CI = HBeAg negative chronic infection, HBeAg- CH = HBeAg negative chronic  
333 hepatitis, HBeAg+ CI = HBeAg positive chronic infection, HBeAg+ CH = HBeAg positive  
334 chronic hepatitis, LSM = liver stiffness measurement, HBV DNA = Hepatitis B virus  
335 deoxyribonucleic acid, qHBsAg = quantification of serum HBsAg, ALT = alanine  
336 aminotransferase, AST= aspartate aminotransferase, GGT = gamma-glutamyl transferase.

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	Entire study population n=770	
	qHBsAg level (IU/ml) to identify HBeAg-neg CI subjects	qHBsAg level (IU/ml) to select patients to antiviral therapy
<b>AUROC (95% CI)</b>	0.58 (0.54 - 0.62)	0.54 (0.49 - 0.60)
<b>Optimal cut-off</b>	5,660	6,000
<b>Sensitivity (%)</b>	55.0	46.6
<b>Specificity (%)</b>	55.6	46.9
<b>PLR</b>	1.24	0.88
<b>NLR</b>	0.81	1.14

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340 **Table 2:** Performance of HBsAg levels for the identification of HBeAg-neg chronic infection  
341 (CI) patients (left column) and treatment eligibility (right column)

342 *Offs: Abbreviations: AUROC: area under the receiver operating characteristics, CI: chronic*  
343 *infection, PLR: positive likelihood ratio, NLR: negative likelihood ratio*

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456 **Figure legends**

457

458 **Figure 1:** Study flow chart

459

460 **Figure 2:** Correlation between HBsAg levels [IU/ml] and HBV viral load [IU/ml] in the whole  
461 study population (A), in HBeAg-negative patients (B) and HBeAg-positive patients (C). Axis are  
462 on logarithmic scale.

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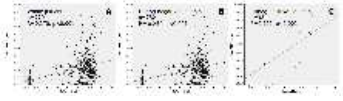
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