## Absence of mutations in the HBsAg "a" determinant during REP 2139 therapy validates serum HBsAg reductions observed in the REP 102 protocol

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#### Figure 1 (A): MHR haplotype complexity during treatment in responders **BACKGROUND & AIMS METHODS** with hepatitis B virus (HBV) may lead to acute or chronic hepatitis. , 240 million people are chronically infected with HBV, and hately 1 million people die annually from HBV-related diseases. Serum samples of 12 patients obtained throughout therapy and during follow-up were used for analysis. Sample were selected up to the lowest possible HBsAg result (highlighted in green in Table 1) containing sufficient HBV DNA for sequencing analysis (see Table division the second sec 16 14 12 10 8 6 4 2 2 Number of haplotypes ◆ P2 ● P3 △ P4 × P6 × P7 ○ P8 appro Nucleic acid polymers (NAPa) clear HBsAg from the Mood by blocking its release from infected hepatocytes. In the REP 102 protocol (NCT02646189) achieved 2-7 log reductions of serum HBsAg accompanied by 3-9 log reductions in serum HBV DNA and the appearance of ant-HBs (A-Mahaliba et al., PCR products for direct and deep sequencing were prepared by single or semi nested PCR of a HBsAg fragment of HBV DNA. T2 Sampling Time points HBsAg is the primary diagnostic target of serological diagnostic in HBV infection. The detection of HBsAg depends on the "a" determinant region (124 - 147aa). Mutations within this region may impair detection of HBsAg and allow HBV to escape vaccine induced immunity or passive immunoglobulin therapy. 4 / 9 responder patients increases decline at end of the treatment. Haplotyp ndergo any clear changes in abundance Deep sequencing targeted the major hydrophilic region (MHR) (including the "a" determinant) of HBsAg. treatment, folic 5/9 responder wed by a sharp de Direct sequencing was performed by Supremerun® and analysed with Geneious® software. Figure 1 (B): MHR haplotype complexity in non-responders . Sequence analysis of a 376 bp fragment of HBsAg was performed determine haplotypes in patients at different phases of REP 2139 there REP 102 protocol to better understand its outcome. 12 10 8 6 4 Number of haplotypes NGS analysis was performed on Illumina® data from all 12 REP 102 patients. Investigate possible sequence differences among the haplotypes betw responder and non-responder patients, which could help understand me of the drug action. -0-P10 The NGS pipeline involved read filtering (Cutadapt), assembly (BWA), variant calling (GATK) and haplotype reconstruction (QuasiRecomb) to detect variants within Investigate to what extent the accumulation of mutations in HBsAg region be responsible for the kinetic of response to NAPs. Т3 T1 T2 Sampling Time points the samples. Explore and identify the mutations within the "a"- determinant region that m result in reduced detection of HBsAg by standard diagnostic assays. roughout the treatment for two of the non-ent 5) shows a dramatic increase following

RESULTS

- Of 12 patients treated (1 gtA, 4 gtD, 7 gtC), 9 responders (with HBsAg reduction) and 3 non responders (1 log or less HBsAg reduction) were identified (Al-Mahtab et al., 2016, see Table 1).
- No mutations evolved in the "a" determinant region during REP 2139 therapy in all 12 patients (Figure 2). Haplotypes either decreased or remained unchanged during treatment (Figure 1).

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- Among the 3 non-responders (Patient 1, 5 & 10), a total of 4 mutations were observed occurring outside the "a" determinant region include G76C, L88P, V118G and P120T (Figure 3).
- In the 9 responder patients, 24 different mutated positions were observed occurring outside the "a" determinant region which include G76C, F85C, L88P, L98V, D99G, Q101R, Q101K, M103I, V106F, L109P, L109Q, G112R, T115N, V118G, G119R, P120T, R122K, P153T, A159V, F161Y, V168A, F170S, Q181R V184A, P188L, P188H, and T189I (Figure 2, Table 2).
- Within the "a"-determinant, mutations were observed in some haplotypes I126S (Patient 3) and 1126T (Patient 12) with frequencies of 18% and 13% respectively, which disappeared as REP 2139 therapy continued (Figure 2).
- Additionally G130R (Patient 8), G145R (Patient 6) and D144E (Patient 9) mutations were reported with a very low frequencies (3.4%, 6% and 2.89% respectively), which also disappeared as REP 2139 therapy continued (Figure 2)



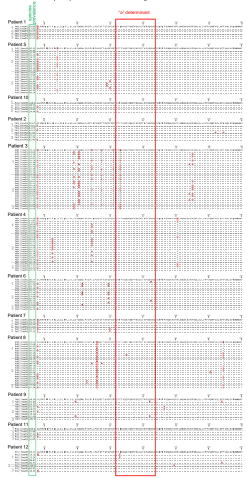
U / mL) hed in Al-Mahtab et al., 2016. ient due to unrelated infectious gastroenteritis for 2 weeks at this sampling point Table 2 Common MHR mutations shared across the responders

Amino acid position	Amino acid substitution	Responder Patients
76	G -> C	2, 3, 4, 6, 7, 8, 9, 11, 12
109	P > L	3
	L > Q	4
	L -> Q	8
115	N -> T	3
	T -> N	8
126	1-> S	3
	1.> T	12
188	P.>L	9

## **CONCLUSIONS & PERSPECTIVE**

- Mutation of the "a"-determinant region of HBsAg or alteration in MHR quasispecies does not occur with REP 2139 therapy.
- Evolution of HBsAg variants escaping detection by HBsAg assays does not occur with chronic exposure to REP 2139.
- No relationship between haplotypes present within the MHR of HBsAg and response to REP 2139 therapy were observed.
- Treatment with REP 2139 does not appear to induce any selection pressure on the MHR.
- Circulating virus species persisting at later stages of REP 2139 treatment may not be recognized by an impaired host immune response.

Figure 2. Deep sequencing analysis. HBsAg quasispecies prevalence in the major hydrophilic region around the "a"-determinant during exposure to REP 2139 is presented. Quasispecies with > 1% prevalence are presented. The location of the "a"-determinant within the sequence is identified by the red box. The points for quasispecies listings are identified on the left. Individual species prevalence is identified inside the green box.



## REFERENCES

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