An evaluation of entecavir treatment among nucleos(t)ide-naïve Moroccan patients with chronic hepatitis B

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ABSTRACT

Objective: To analyse the efficacy and safety of entecavir (ETV) treatment in nucleos(t)ide (NUC)-naïve Moroccan patients with chronic hepatitis B.

Methods: We retrospectively analysed 41 NUT-naïve Moroccan patients with chronic hepatitis B who received ETV 0.5 mg/day monotherapy for at least 3 months, of whom 3 were HBV envelope antigen (HbeAg) positive and 38 were HBeAg negative. The primary end point was the proportion of patients achieving virological response. Secondary end points included biochemical response (alanine transaminase [ALT] normalisation), serological response (HbeAg and HBV surface antigen [HBsAg] loss or seroconversion) and safety.

Results: The median follow-up duration was 74 weeks (48–144 weeks) and mean age was 43.8 years. Of 41 patients, 6 were primary non-responders and 2 achieved partial virological response at week 48, whereas 35 achieved undetectable hepatitis B virus (HBV) DNA at month 12. Viral suppression was maintained in 97.6% of patients after 3 years of ETV treatment. One patient experienced a virological breakthrough at month 12 of treatment. ALT normalisation occurred in 100% of the patients after 1 year of treatment. Only three patients in our study were HbeAg positive, of whom one has experienced seroconversion at month 12 of treatment. However, HBsAg loss or seroconversion was not achieved during the period of the study. No serious adverse event was reported.

Conclusions: These preliminary results showed that ETV is a safe and potent inhibitor of HBV in NUC-naïve Moroccan patients, but we need to observe more patients for a longer period of time, in order to assess the long-term effectiveness, safety, resistance profile and predictive factors for virological and serological response of ETV.

BACKGROUND

Chronic hepatitis B (CHB) is a major global health problem with more than 400 million people chronically infected in the world.1 Owing to its 25–40% associated lifetime risk of development of cirrhosis-related complications and hepatocellular carcinoma (HCC),2 it causes more than 600 000 deaths every year3 and currently represents 5–10% of liver transplantation indications.4 Morocco is considered a low endemic area thanks to the success of the vaccination programme; nevertheless, CHB still remains an important national disease burden.5

Antiviral therapy is an effective way of improving quality of life and survival by preventing disease progression and even reversing liver fibrosis. The end point of treatment is to ensure a degree of virological suppression that will then lead to biochemical remission, histological improvement and prevention of complications.6 CHB management has witnessed a paradigm shift in the past decade since the introduction of oral nucleotide/nucleoside analogues. Entecavir (ETV) is currently recommended as a first-line choice in current hepatitis B virus (HBV) treatment guidelines because of its potent viral suppression and large genetic barrier to resistance.7
To the best of our knowledge, there are no previous published studies or data regarding ETV treatment in Morocco. In this study, we assessed the efficacy and safety of ETV on patients with CHB receiving ETV treatment in our unit.

METHODS

Study population

This retrospective longitudinal study collected consecutive patients from our outpatient clinic between January 2011 and December 2015. Patients were eligible if they were aged 18–70 years and had CHB defined by hepatitis B surface antigens (AGHBs) carriage beyond a 6-month period. As recommended by European Association for the study of the liver (EASL) practice guidelines, treatment indications were based mainly on the combination of three criteria: Serum HBV DNA levels ≥2000 IU/mL, serum ALT levels above the upper limit of normal and severity of liver disease assessed by liver biopsy showing moderate to severe active necroinflammation and/or at least moderate fibrosis.

Patients were excluded from the study if they had hepatitis D virus, hepatitis C virus and HIV, hepatocellular carcinoma or comorbidities including alcoholic, autoimmune, metabolic liver disease with steatosis or steatohepatitis.

Among 65 patients treated with ETV daily for at least 3 months, 24 were excluded because of co-infection with Hepatitis C virus (n=11) and unavailable results for baseline and/or month 3 sera HBV DNA (n=13). A total of 41 patients were included in this analysis and took ETV monotherapy 0.5 mg/day (Bristol-Myers Squibb, Shanghai) for at least 3 months.

Study design

Pretherapeutic biological evaluation was performed systematically for each patient including: Routine haematological analysis, hepatobiliary enzymes, serological analysis, hepatic synthetic function, blood urea nitrogen, creatinine, thyroid-stimulating hormone ultrasensitive (TSHus) and blood glucose level. Serum ALT level was 34 (26–255) IU/L and the serum HBV DNA level was 3.47 (2.3–5.1) log10 IU/mL. The median treatment duration was 74 weeks (range, 48 to 144 weeks).

RESULTS

Baseline characteristics of study patients

The mean age of the 41 patients included in the study was 43.8±10.9 years. Only 26/41 of the patients (63.4%) with CHB were diagnosed through a screening test (Table 1). There were 31 men (75, 6%) and 5 patients (12.2%) with cirrhosis. The median baseline serum ALT level was 34 (26–255) IU/L and the serum HBV DNA level was 3.47 (2.3–5.1) log10 IU/mL. The median treatment duration was 74 weeks (range, 48 to 144 weeks). Of 41 patients, 3 (7.3%) were HBeAg-positive and 38 (92.7%) were HBeAg-negative.

End points and definitions

The primary end point was the proportion of patients achieving virological response defined as undetectable HBV DNA (<100 IU/mL) by a sensitive PCR assay evaluated every 3–6 months during therapy. Primary non-response is defined as less than 1 log10 IU/mL decrease in HBV DNA level from baseline at 3 months of therapy. Partial virological response is defined as a decrease in HBV DNA of more than 1 log10 IU/mL but detectable HBV DNA after at least 6 months of therapy in compliant patients. Virological breakthrough is defined as a confirmed increase in HBV DNA level of more than 1 log10 IU/mL compared to the nadir (lowest value) HBV DNA level on therapy. Secondary end points included biochemical response (ALT normalisation), serological response (HBV envelope antigen (HbeAg) and HBV surface antigen (HbsAg) loss or seroconversion) and treatment-related adverse events.

Table 1 Characteristics of patients at baseline

<table>
<thead>
<tr>
<th>Baseline demographics</th>
<th>HBeAg-positive</th>
<th>HBeAg-negative</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>n (%)</td>
<td>3 (7.3)</td>
<td>38 (92.7)</td>
<td>41 (100)</td>
</tr>
<tr>
<td>Male (%)</td>
<td>3 (100)</td>
<td>28 (73.7)</td>
<td>31 (75.6)</td>
</tr>
<tr>
<td>Age (years, mean)</td>
<td>41.7±14.4</td>
<td>44±10.8</td>
<td>43.8±10.9</td>
</tr>
<tr>
<td>Presence of cirrhosis (%)</td>
<td>1 (33)</td>
<td>4 (10.5)</td>
<td>5 (12.2)</td>
</tr>
<tr>
<td>Interferon experienced (n, %)</td>
<td>3 (100)</td>
<td>24 (58.5)</td>
<td>27 (65.8)</td>
</tr>
<tr>
<td>Naïve patients (n, %)</td>
<td>0 (0)</td>
<td>14 (36.8)</td>
<td>14 (34.1)</td>
</tr>
<tr>
<td>ALT (U/L, median)</td>
<td>40 (30–42)</td>
<td>31 (26–255)</td>
<td>34 (26–255)</td>
</tr>
<tr>
<td>HBV DNA (log10 IU/mL)</td>
<td>5 (2.3–5.1)</td>
<td>3 (2.5–4.7)</td>
<td>3.47 (2.3–5.1)</td>
</tr>
</tbody>
</table>

ALT, alanine transaminase; HbeAg, HBV envelope antigen; HBV, hepatitis B virus.
TREATMENT OUTCOMES

Virological response
Six patients (14.6%) were primary non-responders and 2 achieved partial virological response (4.8%) at week 48, whereas 35 (85.3%) achieved undetectable HBV DNA at month 12. Viral suppression was maintained in 40/41 patients (97.6%) after 3 years of ETV treatment (figure 1). One patient (2.4%) experienced a virological breakthrough at month 12 of treatment.

Biochemical and serological response
ALT normalisation occurred in 100% of the patients after 1 year of treatment (figure 1). Only 3/41 patients in our study were HbeAg positive, of whom one experienced seroconversion at month 12 of treatment. However, HBsAg loss or seroconversion was not achieved during the period of the study.

Safety and side effects
During this study, there were no significant clinical side effects reported, especially no induced renal impairment and no lactic acidosis. Patients with liver cirrhosis did not develop any clinical decompensation or HCC during ETV therapy.

DISCUSSION
The main end point of chronic HBV treatment with nucleos(t)ide (NUC) is complete viral suppression. Once HBV DNA is cleared, continuing treatment may achieve HBeAg clearance and antiHbe seroconversion in HBeAg-positive patients and less frequently HbsAg clearance and antiHBs seroconversion in HBeAg-positive and negative patients.9 In the present study, the results of the 3-year analysis of 41 NUC-naïve Moroccan patients with CHB were presented. Precore mutant CHB was found in 38 (92.7%) of 41 patients. This finding agrees well with data reported in studies, confirming that it is the predominant type of CHB in the Mediterranean basin10 and thus in Morocco.11 HBeAg-negative CHB is a potentially severe disease that rapidly progresses to cirrhosis and related complications.12 Therefore, appropriate use of effective therapy is an important issue in the management of this group of patients.

There was obvious serum HBV DNA reduction and increased virological response at month 3 and month 6, which confirm ETV to be a highly potent antiviral agent. The rate of ALT normalisation was increased over time too, which is consistent with previous results. The virological response was found to be 85, 3% at the end of 12 months and more than 95% patients after 36 months of ETV treatment. In Spain, treatment for 12 months was associated with 82% HBV DNA negativisation, 26% HBeAg and 2% HBsAg clearance rates in 190 NUC-naïve patients.13 In Italy, treatment of 100 consecutive NUC-naïve patients reported 94% HBV DNA negativisation, 33% HBeAg and 15% HBsAg clearances rates after 36 months of continuous ETV treatment.14 Results from Hong Kong showed that 222 treatment NUC-naïve patients achieved 92% HBV DNA negativisation, 44% HBeAg and 0, 45% HBsAg clearance rates after 36 months of ETV treatment.15 All these studies showed a very low incidence of ETV resistance and virological breakthrough: <1%. In our study, one patient (2, 4%) had experienced a virological breakthrough at month 12 of treatment and was treated with ETV plus Adéfovir ADV (Ténofovir unavailable in Morocco).

Those present results are consistent with other experiences16–18 showing increasing virological and biochemical response rates with continuous treatment. Controversially, they showed lower HBeAg clearance and seroconversion rates which emphasise the importance of long-term treatment. In addition, our study confirms that ETV treatment is safe as reported in clinical trials.19 There were no serious side effects and no patient discontinued ETV due to intolerance.

CONCLUSION
In summary, we acknowledge that this study has its limitations. Even though patients were prospectively followed, it is a retrospective study with a small sample size. However, the findings are consistent with other studies, confirming the efficacy and safety of ETV in the management of CHB in Morocco.
size. Also, we did not measure HbsAg levels at baseline and on-treatment response due to economic reasons (non-covered test by health insurance). In conclusion, we were able to demonstrate that ETV treatment is a highly effective anti-HBV treatment for Moroccan patients, with a favourable tolerability profile. This study has shown that ETV is a potent inhibitor of HBV, with greater efficacy in NUC-naïve Moroccan patients, but we need to observe more patients for a longer period, in order to assess the long-term effectiveness, safety, resistance profile and predictive factors for the virological and serological responses of ETV.

Competing interests None declared.

Ethics approval Faculty of Medicine and Pharmacy, Ethics Committee, Rabat, Morocco.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement No additional data are available.

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REFERENCES