Nucleos(t)ide Analogues for Reducing Hepatocellular Carcinoma in Chronic Hepatitis B Patients: A Systematic Review and Meta-Analysis

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Abstract

**Background/Aims:** Studies have shown that nucleos(t)ide analogue (NA) treatment can reduce the risk of hepatocellular carcinoma (HCC) in chronic hepatitis B (CHB) patients, but it is unclear which NA is most effective. We performed a meta-analysis and systematic review comparing the efficacies of NAs in CHB patients. **Methods:** We searched literature databases for randomized controlled trials (RCTs) and observational studies that analyzed the hepatic biochemical response, virological response, seroconversion rate, drug resistance rate, and HCC incidence rate in CHB patients treated with NAs. Meta-analyses were performed with RevMan and Stata/SE software. **Results:** Twelve cohort studies and one RCT were selected, in which entecavir (ETV), lamivudine (LAM), telbivudine (LdT), and/or tenofovir disoproxil fumarate (TDF) were evaluated in CHB patients. The meta-analysis showed that ETV was superior to LAM with regard to the HCC incidence (p=0.0001), biochemical response (p=0.001), virological response (p=0.02), and drug resistance (p=0.00001), and ETV was superior to LdT with regard to the virological response (p=0.0002) and drug resistance (p=0.0002). We found no significant difference between ETV and TDF with regard to the HCC incidence (p=0.08), biochemical response (p=0.39), virological response (p=0.31), serological conversion (p=0.38), or drug resistance (p=0.95). NA-treated patients with pre-existing cirrhosis had a 5.49 times greater incidence of HCC than those without cirrhosis (p=0.00001). **Conclusions:** ETV or TDF should be used for long-term first-line monotherapy in CHB patients according to the current guidelines. Standardized protocols are needed for future studies of ETV and TDF to facilitate conclusive comparisons. Patients with cirrhosis are at significantly elevated risk for HCC, despite the benefits of NA treatment.

**Key Words:** Chronic hepatitis B; Hepatocellular carcinoma; Nucleos(t)ide analogues; Entecavir; Lamivudine; Telbivudine; Tenofovir; Meta-analysis
INTRODUCTION

Liver cancer is the third leading cause of cancer-related mortality in China,¹ the world’s most populous nation, and the second leading cause of cancer-related mortality worldwide.² Hepatocellular carcinoma (HCC) is the most common primary malignancy of the liver, accounting for 70% to 90% of primary liver cancer cases.²³ Hepatitis B virus (HBV) infection is a major risk factor for HCC,⁴ and the incidence of HCC is highest in areas where HBV infection is endemic.⁵⁶ In recent decades, the increased availability of antiviral treatments and HBV vaccines has resulted in reductions in the incidence of HBV infection in various regions.³⁷ However, in 2012, approximately 240 million people had chronic HBV infection (CHB),⁸ and HBV immunisation coverage in areas of endemicity has not increased substantially since then, except in Southeast Asia.⁹ Thus, HBV-related HCC remains a serious threat to public health on a global scale.

Most CHB patients are treated with alpha-interferon and/or a nucleos(t)ide analogue (NA).¹⁰ Interferon induces the expression of hundreds of genes that enhance the innate immune response against HBV-infected hepatocytes,¹¹ whereas currently available NAs act directly to suppress HBV replication by inhibiting viral reverse transcription.¹² The use of interferon in clinical practice is, however, often limited because it has severe side effects, which include thrombocytopenia, neutropenia, lymphopenia, insomnia, and depression.¹³,¹⁴ Side effects of NA treatment are generally mild and infrequent.¹⁴ At present, NAs approved for treating CHB worldwide include entecavir (ETV), tenofovir disoproxil fumarate (TDF), tenofovir alafenamide (TAF), lamivudine (LAM), adefovir dipivoxil (ADV), and telbivudine (LdT), among which ETV and TDF are recommended for first-line treatment because of the lower incidences of resistance observed with ETV and TDF, compared to other NAs.¹⁴,¹⁵ Continuous treatment with NAs can delay clinical progression of CHB in patients with and without cirrhosis.¹⁶-¹⁸ However, while NAs suppress viral replication, they do not completely eliminate HBV in many patients.¹⁹,²⁰ Therefore, many CHB patients need
long-term antiviral treatment.\textsuperscript{14,15,21}

Clinical studies have shown that NA treatment also reduces the risk of HCC in CHB patients to varying degrees.\textsuperscript{16,22-26} However, many studies have not found significant differences between different NAs with regard to reductions in HCC incidence,\textsuperscript{27-32} except in patients with pre-existing cirrhosis.\textsuperscript{24,33} Direct comparisons between studies of different NAs are confounded by differences in study design, selection methods, treatment regimens, and follow-up durations.\textsuperscript{34} Relatively few large-scale studies have been performed that evaluated the efficacies of different NAs for reducing HCC risk, most of which have consisted of retrospective analyses,\textsuperscript{26,27} which have an inherently higher risk of selection bias than a prospective study or randomised control trial (RCT). Furthermore, a number of studies comparing the efficacy of different NAs for reducing HCC incidence, including one RCT,\textsuperscript{28} have analysed data from relatively small samples,\textsuperscript{32,35-37} which might call into question the statistical power of the findings from each.

Given that long-term treatment is suggested for many CHB patients, the choice of which NA to use in such circumstances should, of course, consider the patient’s biochemical response to antiviral treatment, virological response, seroconversion, nephrotoxicity, and NA resistance, properties which have been widely studied. However, the relative efficacies of NAs for long-term reduction of HCC risk and the effects of cirrhosis on HCC outcomes in NA-treated CHB patients are topics that have not been evaluated thoroughly.

To facilitate evidence-based selection of NAs for long-term antiviral treatment in CHB patients, we performed a systematic review and meta-analysis of investigations of NA efficacy that examined HCC incidence as a primary outcome in patients with or without pre-existing cirrhosis. Given that ETV is likely the most widely used first-line antiviral treatment for CHB currently, we performed our analysis using ETV as the reference against which other NAs were compared.
MATERIALS AND METHODS

1. Selection criteria

We included RCTs and observational studies that met the following inclusion criteria: (1) Language: Published in English. (2) Patients: Adults who had received a diagnosis of CHB based on HBsAg-positive lab results or a documented history of HBV infection for ≥6 months, with or without cirrhosis; were either treatment naive or had previously received NAs; and if previously treated with an NA, had no drug resistance at enrolment. (3) Intervention: One treatment group receiving ETV and at least one other group receiving TDF, TAF, LAM, ADV, or LdT; and a minimum treatment duration of 3 months for all groups. (4) Comparator: ETV treatment group. (5) Outcomes: The primary outcome examined was HCC incidence at ≥1 year following initiation of NA treatment. We excluded studies that included patients who developed HCC or died within the first 6 months of treatment and those that included patients who received medications that might affect HCC risk, such as IFN, metformin, statins, or antiplatelet agents.

Secondary outcomes included hepatic biochemical response, virological response, seroconversion, antiviral drug resistance, and development of cirrhosis. Favourable biochemical response was defined as the normalisation of the level of alanine aminotransferase (ALT) as assessed by routine hepatic panel. We defined early cirrhosis as histological findings (Ishak score 5–6) or radiological findings in the absence of decompensated hepatic function or portal hypertension. Favourable virological response was defined as the loss of HBeAg with or without anti-HBe antibodies and undetectable HBV DNA, or undetectable HBV DNA in an HBeAg-negative patient. Antiviral drug resistance was defined as the reappearance of HBV DNA after a period of non-detectable HBV DNA. We excluded articles with previously published data, meta-analyses, review articles, editorials and other types of commentary articles, conference abstracts and presentations, and any publication from which data regarding HCC incidence could not be extracted. Studies
that lacked clear definitions of biochemical response, virological response, drug resistance, and cirrhosis were excluded from the respective subgroup analysis.

2. Data sources and searches

On December 31, 2018, we performed separate searches of Medline, EMBase, and Cochrane Library using the following terms: (1) “[hepatocellular carcinoma OR liver cell carcinoma OR liver cancer OR hepatoma] AND [antiviral OR antiviral drug OR nucleoside analogue OR nucleotide analogue OR nucleos(t)ide analogue]”; (2) “[entecavir OR Baraclude] AND [tenofovir disoproxil fumarate OR tenofovir disoproxil OR tenofovir OR Viread OR tenofovir alafenamide OR Vemlidy] AND [hepatocellular carcinoma OR liver cell carcinoma OR liver cancer OR hepatoma]”; (3) “[entecavir OR Baraclude] AND [lamivudine OR Epivir OR 3TC OR Zeffix] AND [hepatocellular carcinoma OR liver cell carcinoma OR liver cancer OR hepatoma]”; (4) “[entecavir OR Baraclude] AND [adefovir OR adefovir dipivoxil OR bis-POM PMEA OR Preveon OR Hepsera] AND [hepatocellular carcinoma OR liver cell carcinoma OR liver cancer OR hepatoma]”; and (5) “[entecavir OR Baraclude] AND [telbivudine OR Sebivo OR Tyzeka] AND [hepatocellular carcinoma OR liver cell carcinoma OR liver cancer OR hepatoma]”. Two researchers (XW, ZD) independently viewed the titles and abstracts of the articles retrieved, and those that were obviously irrelevant to the selection criteria were discarded.

3. Study selection and data extraction

Study selection and data extraction were performed independently by two researchers (XW, ZD). Titles of the retrieved studies were screen to remove duplicates. Afterward, the abstracts were reviewed to exclude those that did not meet the inclusion criteria. The full-text versions of the remaining articles were then
reviewed to determine inclusion/exclusion. For articles in which they were not in agreement, a third researcher (XL) was consulted to reach a majority decision. Data extraction was performed using a standardised form, which included the following information: (1) Basic study information included the authors’ names, publication date, study location, study dates and duration, sample sizes, and details of the study design, including descriptions of blinding, allocation concealment, and randomisation for RCTs. (2) Baseline characteristics of the study subjects included age, sex, ethnicity, serum biochemical data, CHB diagnostic criteria, HBV DNA level, cirrhosis, decompensation of cirrhosis, previous therapy for CHB, and previous NA treatment. (3) Intervention details included the NA drug used, duration of NA treatment, and follow-up period. (4) Outcome measures included the incidence of HCC, development of cirrhosis or decompensation, incidence of NA resistance, serological conversion, and biochemical and virological responses to NA treatment.

4. Quality assessment

The methodological quality of the data from selected studies was evaluated based on the risk of various biases involved according to the study design. Methodological quality of data from the cohort studies was assessed using the Newcastle-Ottawa Scale (NOS), by which a total score of 0 to 9 stars was assigned to each study. A maximum of 4 stars for selection, 2 stars for comparability, and 3 stars for outcome was combined to calculate the total NOS score, with the maximum score of 9 stars indicating lowest possible risk of bias. Methodological quality in RCTs was assessed using the Jadad Scale, whereby a maximum score of 5 indicated lowest possible risk of bias.

5. Statistical analysis
The Review Manager (RevMan) 5 software, version 5.3, and Stata/SE software, version 15.0, were used for the meta-analysis of the selected studies and the presentation of the results. Comparisons of the outcome incidences are expressed as risk ratio (RR) and 95% confidence interval (CI), which were calculated using random or fixed effects models, and the results are presented as Forest plots. Heterogeneity was evaluated based on the $I^2$ statistic. A fixed-effects model was used when significant heterogeneity existed among the studies, and a subgroup analysis was conducted to identify factors contributing to heterogeneity. A random-effects model was used when significant heterogeneity was not detected. A descriptive analysis was used if the source of heterogeneity between two groups could not be eliminated. Heterogeneity between groups was evaluated using a chi-squared analysis with $\alpha=0.10$. The effect size was estimated based on the Z statistic, which was evaluated using a chi-squared analysis with $\alpha=0.05$. In the case of heterogeneity, a sensitivity analysis was used to apportion the source of heterogeneity by step-wise removal of one of each of the studies in the data set (n-1) over consecutive iterations (n). Funnel plots were used to examine the potential for publication bias, and the Begg’s test and Egger’s test were performed with a level of significance set at $p<0.10$.

RESULTS

1. Search results and literature screening

Study selection is depicted in the flow diagram in Fig. 1. A total of 2,940 articles were retrieved in the electronic searches, from which 740 duplicates were removed. In abstract/title screening of the remaining 2,200 articles, we discarded 2125 articles that did not meet the inclusion criteria. Full-text screening of the remaining 75 articles resulted in the exclusion of 62 articles, leaving a total of 13 articles that were selected for meta-analysis. No studies of TAF met the inclusion criteria.
2. Study characteristics and quality

The basic characteristics of the selected studies are shown in Table 1. A total of 13 studies were selected for meta-analysis, which included 12 cohort studies\textsuperscript{24,27,29-33,35-37,41,42} and 1 RCT\textsuperscript{28}. Only one study\textsuperscript{35} examined the effects of ADV treatment on HCC incidence. The study locations included Korea (n=4), Japan (n=2), Taiwan (n=2), Turkey (n=2), Canada (n=1), Spain (n=1), and Greece (n=1). Most of study participants were of Asian or Caucasian ethnicity. The results of bias risk assessment are also shown in Table 2. Only one cohort study\textsuperscript{24} scored 9 stars on the NOS, whereas four of the cohort studies\textsuperscript{29,35-37} had a total score of 5 stars or less. The single RCT selected for our meta-analysis scored 2 on the Jadad Scale. Data regarding ADV-treated patients\textsuperscript{35} was deemed insufficient for meta-analysis due to small sample size (n=11).

3. HCC incidence in NA-treated CHB patients

In our initial meta-analysis, we compared the incidence of HCC in CHB patients treated with ETV with that of patients treated with other NAs. Treatment with ETV reduced the incidence of HCC by 21\% (RR, 0.79; 95\% CI, 0.49 to 1.27; p=0.34), compared with that of other NAs, but the difference was not statistically significant (Fig. 2A). However, significant heterogeneity in HCC incidence data was detected among the 13 studies (I\textsuperscript{2}=81\%, p<0.00001), so subgroup analyses of the different NAs were performed. In the seven studies in which LAM was used,\textsuperscript{24,27,30,33,35-37} while heterogeneity persisted (I\textsuperscript{2}=43\%, p=0.10), the HCC incidence in CHB patients treated with ETV was 55\% lower than that of those who were treated with LAM (RR, 0.45; 95\% CI, 0.30 to 0.67; p<0.0001). A sensitivity analysis was performed to identify the source of heterogeneity, but did not provide a statistically significant result (data not shown). The incidence of HCC in ETV-treated CHB patients was lower than that in LdT-treated patients\textsuperscript{28,35,36} (RR, 0.72; 95\% CI, 0.24 to 2.14) (Fig. 2C), but was higher than that in TDF-treated CHB patients\textsuperscript{35,36} (RR, 1.52; 95\% CI, 0.94 to 2.44) (Fig.
However, these differences in HCC incidence were not statistically significant (p>0.05 for both). These results were undoubtedly influenced by heterogeneity in the two data sets. While heterogeneity in the ETV versus LdT and ETV versus TDF subgroup analyses was less than that in the overall analysis, the results were not statistically significant (p>0.05 for both). A subgroup analysis of ETV versus ADV was not performed due to the small sample of ADV-treated patients (n=11) in the Coffin et al. study, which was the only study that used ADV among the 13 studies included in our meta-analysis. Funnel plots for the overall (Fig. 2A), ETV versus LAM (Fig. 2B), and ETV versus TDF (Fig. 2D) subgroup analyses did not indicate publication bias (Begg’s and Egger’s tests: p>0.1 for all). No funnel plot was constructed for ETV versus LdT because only three studies were analysed.

4. Biochemical response in NA-treated CHB patients

In the three studies that stringently examined biochemical response to NA treatment, treatment with ETV increased the incidence of favourable biochemical response in CHB patients by 12% (RR, 1.12; 95% CI, 1.01 to 1.24; p=0.04), compared with that observed in patients treated with LAM, TDF, or LdT (Fig. 3A). However, significant heterogeneity was detected among the three studies (I²=53%, p=0.10). In the subgroup analyses (Fig. 3A), ETV treatment significantly increased the incidence of favourable biochemical response in CHB patients (RR, 1.32; 95% CI, 1.11 to 1.56; p=0.001), compared to LAM treatment, whereas no significant difference was observed in the incidence of favourable biochemical response between ETV-treated CHB patients and those treated with TDF (RR, 1.06; 95% CI, 0.93 to 1.20; p=0.39) or those treated with LdT (RR, 1.09; 95% CI, 0.96 to 1.23; p=0.17). Although the Begg’s test of the funnel plot data indicated potential publication bias (p=0.089) (Fig. 3A), it was likely confounded by the small number of studies in the analysis of biochemical response (n=3), whereas the Egger’s test did not indicate publication bias.
bias (p=0.119) (Fig. 3A). Funnel plots were not constructed for the subgroup analyses due to the small number of studies in each.

5. Virological response in NA-treated CHB patients

Six of the selected studies examined virological response in CHB patients treated with NAs. Among these studies, treatment with ETV increased the incidence of favourable virological response by 13% (RR, 1.13; 95% CI, 0.99 to 1.30; p=0.08), compared with that in patients treated with LAM, LdT, or TDF (Fig. 3B), but the difference was not statistically significant. Significant heterogeneity was detected among these studies ($I^2=91\%$, p=0.00001). Subgroup analyses showed that rate of favourable virological response for ETV was 15% greater than that for LAM (RR, 1.15; 95% CI, 1.03 to 1.29; p=0.02), and 37% greater than that for LdT (RR, 1.37; 95% CI, 1.16 to 1.62; p=0.0002). The rate of favourable virological response in ETV-treated patients versus that in TDF-treated patients was statistically similar (RR, 0.95; 95% CI, 0.99 to 1.30; p=0.31). However, both the Begg’s and Egger’s tests of the funnel plot data indicated potential publication bias in the overall analysis (p=0.072 and p=0.010, respectively) (Fig. 3B).

6. Seroconversion in NA-treated CHB patients

Four of the selected studies examined viral seroconversion in CHB patients treated with NAs. In our meta-analysis, we found no significant difference in the rates of seroconversion between patients treated with ETV and those who received LAM, LdT or TDF (RR, 0.98; 95% CI, 0.80 to 1.20; p=0.84) (Fig. 3C). Heterogeneity in the seroconversion data was not detected among the studies analysed ($I^2=0\%$, p=0.54), but the result was not statistically significant. The Begg’s and Egger’s tests of the funnel plot data for the overall analysis did not indicate publication bias (p>0.10 for both) (Fig. 3C).
7. Drug resistance in NA-treated CHB patients

Six of the selected studies\textsuperscript{24,27,28,30,32,37} examined the incidence of NA resistance in CHB patients. Our meta-analysis showed that ETV treatment reduced the development of NA resistance by 95% (RR, 0.05; 95% CI, 0.02 to 0.12, \( p < 0.00001 \)), compared with that observed in patients receiving another NA (Fig. 3D). Subgroup analyses were performed because significant heterogeneity (\( I^2 = 59\% \), \( p = 0.02 \)) was detected, which showed that the incidence of ETV resistance in CHB patients treated was 97% and 96% lower that of LAM (RR, 0.03; 95% CI, 0.02 to 0.04; \( p < 0.00001 \)) or LdT (RR, 0.04; 95% CI, 0.01 to 0.22; \( p = 0.0002 \)), respectively. We found no significant difference in the rates of NA-resistance between CHB patients treated with ETV and those who were treated with TDF (\( p = 0.95 \)). The Begg’s and Egger’s tests of the funnel plot data for the overall analysis did not indicate publication bias (\( p > 0.10 \) for both) (Fig. 3D).

8. HCC incidence in NA-treated CHB patients with cirrhosis

Seven of the selected studies\textsuperscript{24,27,29-32,35} reported the incidence of HCC in CHB patients with pre-existing cirrhosis who underwent NA treatment. Our meta-analysis found that the incidence of HCC in NA-treated patients with pre-existing cirrhosis was 5.49 times higher than that of those who did not have cirrhosis before initiating NA treatment (RR, 5.49; 95% CI, 3.79 to 7.94, \( p < 0.00001 \)) (Fig. 4). Significant heterogeneity in the data regarding HCC in CHB patients with cirrhosis was not detected among the studies analysed (\( I^2 = 22\% \), \( p = 0.26 \)). The Begg’s and Egger’s tests of the funnel plot data did not indicate publication bias (\( p > 0.10 \) for both) (Fig. 4).

**DISCUSSION**
All NAs used in the treatment of CHB competitively bind HBV DNA polymerase, which inhibits viral reverse transcription, thus blocking viral replication. However, NAs have no effect on the covalently closed circular DNA (cccDNA) of HBV that remains stable for long periods in the nucleus of infected hepatocytes. If NA treatment is stopped before the viral DNA polymerase is inactivated or eliminated by cellular protein turnover, HBV levels rebound rapidly as the cccDNA is used as template to produce viral transcripts which in turn serve as templates for viral reverse transcription. Therefore, optimal CHB clinical outcome requires long-term suppression of HBV replication. Stemming the progression of cirrhosis and preventing HCC are additional primary clinical objectives that are intrinsically linked to viral suppression. However, whether different NAs have equivalent effects on HCC development is unclear, and the impact of cirrhosis on the protective effects of NAs has not been adequately characterised.

The current guidelines for CHB treatment recommend ETV, TDF, or TAF as first-line therapy. These recommendations are largely based on the incidence of drug resistance observed for the different NAs. Long-term use of LAM can result in drug resistance and viral breakthrough. Resistance against ADV and LdT monotherapies also occurs. The loss of HBV suppression due to resistance against these NAs requires rescue therapy in which either ETV or TDF is added to the treatment regimen. Although resistance to ETV is rare, certain viral mutations conferring LAM or LdT resistance can increase the risk of developing ETV resistance, and previous exposure to LAM has been shown to increase the risk of ETV resistance even in the absence of detectable LAM resistance. As a consequence, current guidelines recommend switching to TDF or adding ADV for rescue therapy in patients with LAM or LdT resistance, whereas ETV is recommended in patients with ADV resistance. In rare cases of ETV resistance, TDF should be added for antiviral rescue. Resistance to TDF has not been confirmed. With regard to selecting the best NA for reducing the risk of HCC, a low rate of drug resistance should be a prerequisite to ensure long-term

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suppression of viral replication. Our findings of our meta-analysis regarding the incidences of drug resistance support these recommendations.

The results of our meta-analysis showed that the incidence of HCC in CHB patients treated with ETV was 21% lower (p=0.34) than that among patients treated with other NAs, which included TDF, LAM, ADV, and LdT. In the overall analysis, the only studies with RRs that did not favour ETV were those in which patients were treated with TDF (Fig. 2A). In the subgroup analysis, we analysed the data from eight studies, and found no significant difference in HCC incidence between ETV and TDF (p=0.08) (Fig. 2D). However, it is clear from the forest plot in Figure 2A that the studies with the largest confidence intervals in the data set also had the smallest and greatest effect sizes (Coffin et al. and Hsu et al., respectively), which were likely confounding factors given that a sensitivity analysis could not isolate a single study as the source of heterogeneity (data not shown).

Our findings regarding HCC incidence for ETV versus TDF (Fig. 2D) are consistent with those of a study by Tsai and colleagues (2017). Their retrospective cohort study had a relatively large sample size (n=546), but it was not selected our meta-analysis because they included patients also receiving medications that might influence the risk of HCC. Tsai et al. reported that the cumulative incidence of HCC in ETV-treated patients (n=359) was not significantly different than that in TDF-treated patients (n=83). No RCTs of ETV versus TDF in a large sample have been reported, and no RCTs using TDF met our stringent study selection criteria. The results of our meta-analysis of studies comparing HCC incidence for ETV versus TDF highlight an important need for the implementation of standardised protocols for evaluating ETV and TDF efficacy in CHB patients, which has recently been voiced by others in the scientific community.

Only three studies were included in our subgroup analysis of HCC incidence in ETV-treated versus that LdT-treated patients (Fig. 2C), and the combined sample of LdT-treated patients in our analysis was
relatively small (n=63). These may represent potential confounders of our results. Furthermore, we did not perform an analysis of publication bias due to the small number of studies in our ETV versus LdT subgroup analysis. However, our findings regarding HCC incidence for ETV versus LdT are also consistent with those of Tsai et al. They reported that the cumulative incidence of HCC in ETV-treated patients (n=359) was not significantly different than that in LdT-treated patients (n=104). We also found that the risk of HCC in ETV-treated patients was statistically similar to that in LdT-treated patients (p>0.05).

Our subgroup analysis of HCC risk for ETV versus LAM treatments for CHB found that the risk of HCC in patients treated with ETV was 55% lower than that of patients who were treated with LAM (p<0.0001) (Fig 2B). Studies in Asia and Europe have shown that LAM achieves HBV DNA reduction, HBeAg serological conversion, and ALT normalisation, thereby delaying or preventing disease progression and reducing the incidence of HCC. Although the incidence of NA resistance is highest for LAM, the cost of LAM is approximately 80% to 86% lower than that of other NAs, and LAM may still be used as first-line treatment for CHB in some countries in an effort to maintain cost-effective government-provided health care. Although one study found that TDF monotherapy was more cost-effective for treating CHB than other NAs based on average annual and lifetime disease costs per patient, lifetime cost and life expectancy, and quality adjusted life years, a later study reported that cost-effectiveness of LAM plus ADV was similar to that of TDF. However, no combination of different NAs is currently recommended by the current guidelines, except for rescue therapy following the onset of drug resistance.

Antiviral NA treatment is thought to indirectly reduce the risk of HCC by reducing HBV DNA load, improving liver inflammation, and promoting seroconversion of HBeAg. Therefore, we compared these clinical outcomes in ETV-treated CHB patients with those in CHB patients receiving other NAs. The overall analysis of hepatic biochemical response found that the incidence of favourable response was significantly
greater among ETV-treated patients (p=0.04) (Fig. 3A), and the subgroup analysis showed that, while the rate of favourable biochemical response in ETV-treated patients was significantly higher than that of LAM-treated patients (p=0.0001), there was no significant difference between ETV and TDF or LdT. However, only one study was included in each of our analyses of LAM and LdT, which could have biased our results. In the overall and subgroup analyses, the effects of ETV on serological conversion were not significantly different from those of other NAs (Fig. 3C). However, LAM-treated patients were not included in these analyses, and only one study was used in the comparison of ETV and TDF, which might have confounded our analysis.

In our overall analysis, treatment with ETV increased the incidence of favourable virological response by 13% (Fig. 3B), but this result was not statistically significant (p=0.08). While virological response in ETV-treated patients was significantly better than that in patients treated with LAM or LdT (p<0.05 for both), no significant difference in virological response was observed between ETV and TDF (p=0.31). Heterogeneity in the overall analysis of virological response was reduced in the subgroup analyses, but these results were not significant. Furthermore, the funnel plot analysis for virological response indicated a significant probability that publication bias affected our analysis. Therefore, with regard to virological response in CHB patients, our findings are not inconsistent with the recommendations of current CHB treatment guidelines suggesting either these two drugs for first-line monotherapy. However, like that observed in our analysis of HCC incidence, our lack of statistically significant findings of virological response for ETV versus TDF highlights the importance of developing and implementing standardised protocols in future studies in order to facilitate statistically conclusive analyses.

In a previous systematic review and Bayesian analysis of 20 RCTs of NA treatment for CHB, Woo et al. reported that TDF-treated patients had better rates of seroconversion and favourable virological and
biochemical responses than those of patients treated with LAM, ETV, LdT, or ADV. They also reported that ETV was more effective for reducing HBV DNA level and normalising ALT than were LAM, LdT, or ADV, and that ETV was more effective for improving hepatic histological response than all other NAs. We did not include hepatic histological response in our meta-analysis of NA treatments because, in our experience, histological data quite often are not obtained during follow-up for NA treatment if favourable virological, immunological, and hepatic biochemical responses are achieved, except perhaps in RCTs. Furthermore, all of the RCTs included in the Woo et al. meta-analysis were published before 2010, and none of them performed a direct comparison of ETV and TDF. Our meta-analysis included studies published between 2011 and 2018, which should provide a better representation of clinicians increased experiences using NAs and improvements in the guidelines for CHB treatment.

Multiple previous studies have found that pre-existing cirrhosis is an important risk factor for HCC in CHB patients receiving NA therapy. Previous studies show that NAs can halt the progression of liver cirrhosis histologically, prevent the occurrence of hepatic decompensation, and reduce the incidence of HCC by inhibiting virus replication. However, multiple studies have confirmed that HCC can develop in CHB patients despite effective HBV suppression. Our meta-analysis found that the incidence of HCC in NA-treated patients with pre-existing cirrhosis was 5.49 times higher than that of those without cirrhosis (p<0.00001) (Fig. 4). Therefore, future studies should place greater emphasis on subgroup analyses, especially with regard to elucidating the role of cirrhosis on long-term effects of ETV, TDF, and TAF treatments for CHB.

Although our findings show the reduction in HCC risk was statistically similar between ETV and TDF (P > 0.05), ETV may be preferred over TDF, due to the greater potential of TDF for adverse effects and perhaps a wider experience among physicians using ETV since its approval for CHB treatment in 2005, whereas TDF
was approved in 2008. The tenofovir prodrug, TAF, was recently approved for treating CHB in the USA, Europe, and Japan, and has a lower incidence of renal toxicity and bone demineralisation than TDF. However, very little long-term data is available for TAF with regard to HCC risk, so studies of TAF were not included in our analysis. Likewise, only 11 ADV-treated patients were included in our analysis, and we did not subject data for these patients to subgroup analysis.

Our findings are subject to certain additional limitations. Only one randomised controlled trial was included in our meta-analysis, whereas the remaining 12 studies were retrospective cohort studies. This might be viewed as a shortcoming, due to the greater potential for selection bias in retrospective studies, and study design has been shown to be a crucial factor in the analysis of the effects of NAs on HCC risk. A previous systematic review and meta-analysis found that, while RCTs showed no significant benefit of NA treatment on HCC risk, case-control studies showed a significant benefit, and results for cohort studies showed an increased risk of HCC in NA-treated patients. Inconsistencies between the results of this previous meta-analysis and our findings present herein suggest that differences in study design might have contributed to the heterogeneity detected in our analysis of HCC incidence, as our meta-analysis included ten cohort studies and one RCT. Future large-scale RCTs with standardised methods are needed to better clarify the long-term effects of NAs on HCC risk, especially for comparisons of ETV versus TDF.

In the studies subjected to meta-analysis, we found no significant difference in HCC incidence, biochemical response, virological response, serological conversion, or drug resistance between ETV and TDF treatment groups (p>0.05), which highlights the need for standardised protocols for future studies comparing ETV and TDF efficacy for reducing HCC risk and improving secondary outcomes. Our results did, however, show that ETV was superior to LAM with regard to HCC incidence, biochemical response, virological response, and drug resistance, and ETV was superior to LdT with regard to virological response
and drug resistance (p<0.05 for all). We also found that, despite the benefit of NA treatment, the incidence of HCC was higher in CHB patients with cirrhosis than in those without cirrhosis, which highlights the need for early detection and successful intervention using a high genetic barrier NA to avoid resistance and viral breakthrough. Our findings provide clinical guidance by forming a basis for the selection of optimal first-line antiviral therapies for CHB patients, and place an ever-growing burden on clinical researchers to design and implement higher quality studies of the effects of NA treatment on HCC risk.

CONFLICTS OF INTEREST

The authors declare no conflict of interest.

ACKNOWLEDGEMENTS

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Author contributions: Conception and design of the study: Y.J., Z.Y. Acquisition and analysis of data: XW, X.L., Z.D., L.Y. Statistical analysis: X.W. Writing - original draft: X.W., X.L. Writing - review and editing: X.W., Z.Y. Approval of final version of the manuscript: all authors.
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50. Flemming J, Terrault N. Tenofovir vs Entecavir for Hepatocellular Carcinoma Prevention in Patients With Chronic Hepatitis B: One of These Things Is Not Like the Other. JAMA oncology 2018.


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Figures legends

Fig. 1. Flowchart of study selection.

Fig. 2. (A) Forest and funnel plots for the comparison of HCC incidence between ETV-treated CHB patients and those treated with other NA antiviral drugs (Begg’s test: $z = -0.07$, $p = 1.000$; Egger’s test: $t = 0.34$, $p = 0.743$); (B) Forest and funnel plots for the comparison of HCC incidence between ETV and LAM (Begg’s test: $z = -0.30$, $p = 0.764$; Egger’s test: $t = -1.51$, $p = 0.191$); (C) Forest plot for the comparison of HCC incidence between ETV and LdT; (D) Forest and funnel plots for the comparison of HCC incidence between ETV and TDF (Begg’s test: $z = -0.60$, $p = 0.540$; Egger’s test: $t = -1.78$, $p = 0.135$).

Fig. 3. (A) Forest and funnel plots for the comparison of the biochemical response in ETV-treated CHB patients and CHB patients treated with LAM, LdT, or TDF as controls (Begg’s test: $z = 1.70$, $p = 0.089$; Egger’s test: $t = -2.64$, $p = 0.119$); (B) Forest and funnel plots for the comparison of the virological response in CHB patients treated with ETV and those treated with LAM, LdT, or TDF as controls (Begg’s test: $z = 1.00$, $p = 0.072$; Egger’s test: $t = 4.02$, $p = 0.010$); (C) Forest and funnel plots for the comparison of the serological conversion in CHB patients treated with ETV and those treated with LAM, LdT, or TDF as controls (Begg’s test: $z = 0.00$, $p = 1.000$; Egger’s test: $t = -1.23$, $p = 0.436$); (D) Incidence of drug resistance in CHB patients treated with ETV compared with that in those treated with LAM, LdT, or TDF as controls (Begg’s test: $z = 0.75$, $p = 0.452$; Egger’s test: $t = 0.99$, $p = 0.380$).

Fig. 4. Forest and funnel plots for the comparison of HCC incidence in NA-treated patients with CHB and liver cirrhosis (LC) and CHB patients without liver cirrhosis (Begg’s test: $z = 1.13$, $p = 0.260$; Egger’s test: $t = -1.24$, $p = 0.281$).
Fig. 1. 1403 records identified through Medline searching 1537 records identified through other sources

2200 records after duplicates removed

2200 abstracts screened

2125 records excluded:
597 not related to HBV infection
254 not related to Entecavir and HCC
163 basic science
763 reply, commentary, review
348 HCC incidence not reported

75 full-text articles assessed for eligibility

62 full-text articles excluded:
49 related to Entecavir only
13 HCC incidence not reported

13 studies included

Fig. 1.
Fig. 2.
Fig. 3.
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Total events: 366, 85
Heterogeneity: Tau^2 = 0.05; Chi^2 = 7.65, df = 6 (P = 0.26); I^2 = 22%
Test for overall effect: Z = 9.04 (P < 0.00001)

Fig. 4.