A Cost-Utility and Cost-Effectiveness Analysis of Different Oral Antiviral Medications in Patients With HBeAg-Negative Chronic Hepatitis B in Iran: An Economic Microsimulation Decision Model

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Abstract

**Background:** Although hepatitis B infection is the major cause of chronic liver disease in Iran, no studies have employed economic evaluations of the medications used to treat Iranian patients with chronic hepatitis B (CHB). Therefore, the cost-effectiveness of the different treatment options for this disease in Iran is unknown.

**Objectives:** The aim of this study was to compare the cost utility and cost-effectiveness of medication strategies tailored to local conditions in patients with HB e antigen (HBeAg)-negative CHB infection in Iran.

**Methods:** An economic evaluation of the cost utility of the following five oral medication strategies was conducted: adefovir (ADV), lamivudine (LAM), ADV + LAM, entecavir (ETV), and tenofovir (TDF). A Markov microsimulation model was used to estimate the clinical and economic outcomes over the course of the patient’s lifetime and based on a societal perspective. Medical and nonmedical direct costs and indirect costs were included in the study and life-years gained (LYG) and quality-adjusted life-years (QALY) were determined as measures of effectiveness. The results are presented in terms of the incremental cost-effectiveness ratio (ICER) per QALY or LYG. The model consisted of nine stages of the disease. The transition probabilities for the movement between the different stages were based on clinical evidence and international expert opinion. A probabilistic sensitivity analysis (PSA) was used to measure the effects of uncertainty in the model parameters.

**Results:** The results revealed that the TDF treatment strategy was more effective and less costly than the other options. In addition, TDF had the highest QALY and LYG in the HBeAg-negative CHB patients, with 13.58 and 21.26 (discounted) in all comparisons. The PSA proved the robustness of the model results. The cost-effectiveness acceptability curves showed that TDF was the most cost-effective treatment in 59% - 78% of the simulations of HBeAg-negative patients, with WTP thresholds less than $14010 (maximum WTP per QALY).

**Conclusions:** The use of TDF in patients with HBeAg-negative CHB seemed to be a highly cost-effective strategy. Compared with the other available medication options, TDF was the most cost-saving strategy. Thus, TDF may be the best option as a first-line medication. Patients can also be switched from other medications to TDF.

**Keywords:** Chronic Hepatitis B, Cost-Utility and Cost-Effectiveness Analysis, Oral Antiviral Therapy, Markov Microsimulation Model, Probabilistic Sensitivity Analysis

1. **Background**

Hepatitis B remains a serious health problem worldwide, as it is the most common and serious form of hepatitis and liver disease (1). People with the disease have an increased risk of mortality and morbidity (2). In addition, treatment costs are high, especially in developing countries (2). Currently, an estimated 350 million people...
worldwide have chronic HB (CHB) infection, which results in about 500,000 to 1.2 million deaths per year (3, 4). In Iran, according to the latest statistical data released by the office for the prevention of viral hepatitis in the ministry of health and medical education, the prevalence of hepatitis B in the country was estimated to be about 1.5% (5, 6). Given that the population of Iran was about 78 million in 2014 (7), the total number of chronic HB virus (HBV) carriers was estimated to be 1,170,000 people (5). The genotype D is the predominant genotype of hepatitis in the country (8, 9), and more than 70% of cases are HB e antigen (HBeAg) negative (10).

Given the mortality rate among untreated CHB patients (20% - 25%) (11), long-term and severe outcomes of CHB infection, and its complicated pattern of management, timely diagnosis and treatment are critical to prevent patients progressing to the severe stages of the disease. Although medication therapy cannot cure CHB, except in rare cases (11), early treatment can significantly prevent the progress of severe liver failure and reduce mortality (12).

Oral treatments (oral nucleos(t)ide analogues) are preferred as the first-line therapy for most genotypes of the disease, as they have fewer side effects compared with interferon (12). The goal of drug therapy for CHB patients is to improve their quality of life and survival by preventing progression of the disease to more severe stages, such as liver cirrhosis and liver cancer. At present, five oral medication strategies are used to treat hepatitis in Iran: adefovir (ADV), lamivudine (LAM), ADV + LAM, entecavir (ETV), and tenofovir (TDF). The use of ETV is less common because of its high cost and the fact that it is not indexed in the Iran Drug List. However, to evaluate the cost-effectiveness of all oral drugs in Iran, ETV was included in the studied medication strategies.

Economic evaluation studies published in recent years showed that TDF was more cost effective than other drugs (12-16). However, most of the published economic evaluation studies on CHB treatment have been carried out in the U.S., Europe, and Southeast Asian countries. In contrast, in the Eastern Mediterranean region, specifically Iran, no studies have used economic evaluation models in the assessment of medication therapy protocols for patients with hepatitis B.

Accordingly, due to the lack of sufficient scientific evidence, the best strategy for the treatment of hepatitis B patients in Iran is unclear. The provision of accurate data on both the economic and clinical impact of HBV infection can help policymakers to make informed decisions to provide better health and medical treatment services for these patients. A number of factors necessitated this study. These included the high prevalence of HBV, which accounts for the majority of hepatitis infections in the country, its role as the major cause of cirrhosis and liver cancer (5, 17, 18), the need to identify optimum hepatitis B strategies tailored to local conditions, and the absence of economic evaluation studies on the disease in Iran and the Eastern Mediterranean region.

2. Objectives

This study aimed to assess the cost utility and cost-effectiveness of various medication strategies for HBeAg-negative CHB from a societal perspective to determine the most appropriate protocol for treating HBV infection in Iran.

3. Methods

3.1. Overview

This study consisted of an economic evaluation of the cost utility of five oral medication strategies: ADV, LAM, ADV + LAM, ETV, and TDF. Unlike other studies in the field of hepatitis, we used a Markov microsimulation model to identify economic and clinical outcomes of HBV infection, which requires a long-term follow up due to the chronic and recurring nature of the infection (12). The outcome measures used in the model included life-years gained (LYG), quality-adjusted life-years (QALY), and the costs of the disease and treatment strategy in each state. For extracting costs, a questionnaire was prepared, and its validity and reliability were tested. As this study adopted a societal perspective, both medical and nonmedical costs (direct and indirect) were evaluated. The time horizon of the study was the patient’s lifetime, with one-year cycles, lasting a total of 35 years. In common with other published economic evaluation studies (12, 13) and in accordance with the opinions of experts, the model was run with one-year cycles to cope with the nature of the changes in the various stages of the disease. As the duration of the model in this study was more than one year, the costs and effects of the medication therapy for patients were calculated for one year in 2014. These were discounted during the patient’s lifetime, with an annual rate of 7.2% (19) and 3% (20), respectively. These discount rates have been applied in previous economic evaluation studies published in Iran (21). Excel and Tree Age software, V. 2011 were used to analyze the data.

3.2. Study Population and Treatment Regimens

The sample size of this study consisted of 155 patients with HBeAg-negative CHB. The sample size was calculated based on the statistical sample of the present study and the
profiles registered in the studied centers. Therefore, the characteristics of the patients matched those of the real population of patients who were referred to the research center of gastroenterology and hepatology in Baghiatollah University and Baghiatollah hospital and Nemazee hospital in Shiraz. The centers were selected using purposive sampling because they are referral centers for hepatitis in the country, and people from different parts of the country are referred to these centers.

The mean age of the patients in the sample was 40 y, and 67% of the study group were males.

At the end, a microsimulation analysis of a hypothetical cohort population of 10000 individuals was carried out.

3.3. Inclusion and Exclusion Criteria

The inclusion criteria were as follows: infected with CHB, a biopsy- or fibroscan-confirmed diagnosis of CHB, received the medications evaluated in the study for at least one year, and had a record. The exclusion criteria were patients with acute hepatitis B, HBeAg-positive CHB and newly diagnosed patients, other disease states, and other types of hepatitis.

The study was approved by the ethics committee of the Pharmaceutical Sciences Research Center of Tehran University of Medical Sciences.

The medication therapy regimens included in the model were based on the guide for the treatment of chronic hepatitis B (22). The regimens were as follows: ADV (10 mg daily), LAM (100 mg daily), TDF (300 mg daily), ETV (0.5 mg daily), and a combination of ADV and LAM. All the medications were taken orally.

3.4. Switching Medications

Based on clinical evidence worldwide (23-25) and in Iran (26) on the development of drug resistance, side effects of drugs, or futility of drugs in some cases, medication regimens may need to be altered to enable the treatment to be continued and prevent disease progression in patients.

This study used Tree Age software to design and evaluate a medication-switching model. The medications were switched in cases of drug resistance, side effects, and no response to the initial treatment. According to published studies, the development of resistance was higher with ADV and LAM than other drugs (27-30). However, as gastroenterology and hepatology specialists in Iran often prescribe a combination of two medications (ADV + LAM and TDF), withdrawal was designed only for these two medications, and the possibilities of resistance were considered only for this state.

Concerning the option to switch to other medications, according to the existing registry of gastroenterology and hepatology Centers, 37% and 63% of patients resistant to LAM, respectively, switched to TDF and ADV + LAM; moreover, 48% and 52% of patients resistant to ADV, respectively, switched to TDF and ADV + LAM.

3.5. Model Structure

Figure 1 presents a schematic diagram of the Markov model of the studied sample and for a hypothetical cohort population of 10000 HBeAg-negative patients aged over 18 y.

As shown in the diagram, CHB patients receiving antiviral drugs can have a recurring transition to different disease states. The specified states and their changes were determined based on evidence from other studies and the opinions of experts. Patients in all stages of the disease can develop hepatocellular carcinoma (HCC) and die. The patients included in the initial distribution of the Markov microsimulation model were selected in proportion to a real sample of patients with CHB.

3.6. Key Modeling Assumptions

Based on the available clinical evidence and the results of a review of the medical records of the patients included in the study, the resistance to ETV and TDF drugs was assumed to be zero.

In the first five years, the drug resistance rates for both LAM and ADV were considered to be variable (31). In subsequent years, they were determined based on the values in the fifth year.

As clinical evidence suggests that combination therapy may not result in more effective therapy (32-34) and due to the lack of published data, it was assumed that the transition probabilities between some of the health states were equal to the most effective component of the combination medication shown in Table 1.

It was assumed that nucleos(t)ides had no effect on mortality in patients with HCC and decompensated cirrhosis (DC).

It was assumed that the side effects associated with nucleos(t)ides had no impact on cost, quality of life, and mortality. Nevertheless, the cost of monitoring renal failure was considered in the cost analysis of the model.

For some transitions, such as hepatitis B surface antigen (HBsAg) to HCC, CHB to HBsAg, and CHB to UnHBV DNA to HCC, data on the individual drugs were not available. In these cases, it was assumed that all treated patients had an equal chance of improvement or disease progression, irrespective of the type of medication used.
3.7. Transitional Probabilities

Tables 1 and 2 present the transitional probabilities for all the parameters included in the model. Due to the absence of data from Iranian studies, especially on the clinical efficacy of different medication protocols used for patients with CHB, most of the transitional probabilities between health states were extracted from international studies and evidence on hepatitis B (12, 13, 31-49).

The mortality rate for the general population was extracted from life tables for 2012 and analyzed separately by age and sex (50). To determine the rate of deaths due to hepatitis B, the standardized mortality ratio reported in published articles was used (51). Some probabilities entered in the model were based on the medical records of patients at the centers included in the study.

As the data on clinical efficacy were reported as rates in some articles, they were converted to probabilities using the following Formula 1 and then entered in the model:

\[ p = 1 - exp(-rt) \]

Where \( p \) is the probability, \( r \) is the instantaneous rate, and \( t \) is the time (52).

3.8. Health Outcomes

Health outcomes were specified in terms of LYG and QALY. Utility scores were obtained using EQ-5D questionnaires administered via face-to-face interviews conducted with 155 HBeAg-negative CHB patients in 2014. Related complications, such as compensated cirrhosis (CC), DC, and HCC, were recorded. The interviewees in the study consisted of outpatients referred to the research center for gastroenterology and hepatology in Baghiatoallh University in Tehran (the capital of Iran and a hepatitis referral center) and inpatients admitted to two major hospitals in Tehran and Shiraz. These hospitals are the main treatment centers for liver transplantation patients.

After completing the EQ-5D questionnaire, for the first time, we tried to consider the national values of Iran that were obtained in a separate study by Goudarzi (53) using time trade-off (TTO), and we changed the 5-digit codes of the questionnaire into numerical utility (Table 3). In addition, in a separate calculation using values reported in a U.K. study, the 5-digit codes were changed again into numerical utility and used in the sensitivity analysis of the model.

3.9. Treatment Costs

The cost analysis method used in a study performed in 2005 was applied (5). The costs of each medication strategy in each state for the treatment of CHB, CC, DC, and HCC were calculated, as shown in Table 2. Data on the direct and indirect costs were extracted from both specialists’ and patients’ views and a review of inpatient and outpatient medical records.
Table 1. Input Parameters Used in the Economic Microsimulation Decision Model: HBeAg-Negative CHB

| Variable | Mean | Dist | Lower Values | Upper Values |
|----------|------|------|--------------|--------------|
| Prob CHB to HCC with LAM | 0.040 | Beta | 0.02 | 0.09 |
| Prob CHB to HCC with TDF | 0.032 | Beta | 0.008 | 0.05 |
| Prob CHB to HCC with ADV + LAM | 0.033 | Beta | 0.02 | 0.09 |
| Prob CHB to HCC with ETV | 0.048 | Beta | 0.03 | 0.09 |
| Prob CC to DC with ADV | 0.027 | Beta | 0.004 | 0.0154 |
| Prob CC to DC with TDF | 0.007 | Beta | 0.00 | 0.0099 |
| Prob CC to DC with ADV + LAM | 0.034 | Beta | 0.0055 | 0.0623 |
| Prob CC to DC with ETV | 0.069 | Beta | 0.044 | 0.054 |
| Prob CC to CC with ADV + LAM | 0.027 | Beta | 0.004 | 0.0154 |
| Prob CC to CC with ADV | 0.039 | Beta | 0.02 | 0.073 |
| Prob CC to CC with LAM | 0.040 | Beta | 0.0045 | 0.0728 |
| Prob CC to CC with ETV | 0.019 | Beta | 0.014 | 0.054 |
| Prob HCC to LT | 0.019 | Beta | 0.0055 | 0.037 |
| Prob DC to LT | 0.010 | Beta | 0.00 | 0.2 |
| Prob CHB to CC with TDF | 0.075 | Beta | 0.036 | 0.08 |
| Prob CHB to CC with ADV | 0.031 | Beta | 0.00 | 0.0463 |
| Prob CHB to CC with ADV + LAM | 0.022 | Beta | 0.0305 | 0.0495 |
| Prob CHB to CC with ETV | 0.017 | Beta | 0.00 | 0.047 |
| Prob CHB to CC with LAM | 0.022 | Beta | 0.0305 | 0.0495 |
| Prob CHB to CC with TDF | 0.006 | Beta | 0.00 | 0.0069 |
| Prob CC to CC with ADV | 0.045 | Beta | 0.022 | 0.054 |
| Prob DC to CC with ADV + LAM | 0.025 | Beta | 0.002 | 0.054 |
| Prob DC to CC with ETV | 0.049 | Beta | 0.02 | 0.011 |
| Prob DC to CC with LAM | 0.046 | Beta | 0.02 | 0.011 |
| Prob DC to CC with TDF | 0.029 | Beta | 0.18 | 0.013 |
| Prob CHB to UnHBV DNA with ADV | 0.489 | Beta | 0.254 | 0.771 |
| Prob CHB to UnHBV DNA with TDF | 0.938 | Beta | 0.794 | 0.993 |
| Prob CHB to UnHBV DNA with ADV + LAM | 0.489 | Beta | 0.254 | 0.771 |
| Prob CHB to UnHBV DNA with ETV | 0.720 | Beta | 0.575 | 0.877 |
| Prob CHB to UnHBV DNA with LAM | 0.318 | Beta | 0.235 | 0.424 |
| Prob CHB to UnHBV DNA with ADV | 0.005 | Beta | 0.00 | 0.013 |
| Prob CHB to UnHBV DNA with CC | 0.100 | Beta | 0.145 | 0.275 |
| Prob CHB to UnHBV DNA with CHB | 0.075 | Beta | 0.036 | 0.08 |
| Prob CHB to UnHBV DNA with TDF | 0.038 | Beta | 0.00 | 0.048 |
| Prob CHB to UnHBV DNA with ADV | 0.028 | Beta | 0.05 | 0.013 |
| Prob CHB to UnHBV DNA with ETV | 0.005 | Beta | 0.052 | 0.015 |
| Prob CHB to UnHBV DNA with UnHBV DNA | 0.052 | Beta | 0.0004 | 0.046 |
| Prob CHB to HBeAg | 0.004 | Beta | 0.052 | 0.0099 |
| Prob CHB to HBeAg | 0.015 | Beta | 0.0099 | 0.074 |
| Prob CHB to UnHBV DNA resistance to LAM or ADV | 0.529 | Beta | 0.075 | 0.967 |

Abbreviations: HBeAg, hepatitis B e antigen; CHB, chronic hepatitis B; Prob, probability; CC, compensated cirrhosis; DC, decompensated cirrhosis; HCC, hepatocellular carcinoma; Sero, seroconversion; LT, liver transplantation; W, withdrawal; LAM, lamivudine (100 mg); TDF, tenofovir (300 mg); ETV, entecavir (0.5 mg); ADV, adefovir (10 mg); HBsAg, hepatitis B surface antigen; Dist, distribution.

The costs (in US$) were based on 2014 tariffs. The exchange rate was 26,668 rials per dollar (54). To enable international comparisons, the costs were changed to international dollars using a purchasing power parity (PPP) $ exchange rate of 8565.41 rials per $ (55).

3.10. Cost-Effectiveness Analysis

Based on the results of the aforementioned stages, a model was designed using Tree Age software, and the extracted data were entered into the model. Considering the time horizon of the study, the values for the costs, effectiveness, and cost-effectiveness were calculated as monetary currencies, QALY, and cost per QALY, respectively, for all the
Table 2. Input Parameters Used in the Economic Evaluation Model: HBeAg Negative

| Variable                  | Mean | Dist  | Lower Values | Upper Values |
|---------------------------|------|-------|--------------|--------------|
| Cost of CHB with ADV      | 4256 | Gamma | 3405         | 5107         |
| Cost of CHB with ADV + LAM| 4334 | Gamma | 3684         | 4970         |
| Cost of CHB with ETV      | 15496| Gamma | 13800        | 18000        |
| Cost of CHB with LAM      | 4206 | Gamma | 3800         | 5000         |
| Cost of CHB with TDF      | 4922 | Gamma | 4100         | 5600         |
| Cost of CC with ADV       | 5280 | Gamma | 4224         | 6337         |
| Cost of CC with ADV + LAM | 5358 | Gamma | 4600         | 6100         |
| Cost of CC with ETV       | 16594| Gamma | 14500        | 18500        |
| Cost of CC with LAM       | 5231 | Gamma | 4500         | 6000         |
| Cost of CC with TDF       | 5944 | Gamma | 5200         | 6700         |
| Cost of DC with ADV       | 22476| Gamma | 17980        | 26970        |
| Cost of DC with ADV + LAM | 22554| Gamma | 18900        | 26600        |
| Cost of DC with ETV       | 33709| Gamma | 28000        | 40000        |
| Cost of DC with LAM       | 22426| Gamma | 19000        | 26000        |
| Cost of DC with TDF       | 23339| Gamma | 19000        | 27000        |
| Cost of HCC with ADV      | 40671| Gamma | 33350        | 47990        |
| Cost of HCC with ADV + LAM| 40749| Gamma | 33400        | 48000        |
| Cost of HCC with ETV      | 50992| Gamma | 45000        | 58000        |
| Cost of HCC with LAM      | 40621| Gamma | 34000        | 47000        |
| Cost of HCC with TDF      | 45909| Gamma | 33000        | 50000        |
| Utility CHB               | 0.83 | Beta  | 0.68         | 0.85         |
| Utility CC                | 0.71 | Beta  | 0.64         | 0.75         |
| Utility DC                | 0.48 | Beta  | 0.40         | 0.53         |
| Utility HCC               | 0.51 | Beta  | 0.27         | 0.59         |
| Utility Sero              | 0.95 | Beta  | 0.90         | 0.95         |
| Utility Un DNA            | 0.83 | Beta  | 0.77         | 0.85         |

Abbreviations: HBeAg, hepatitis B e antigen; ADV, adefovir; LAM, lamivudine; ETV, entecavir; TDF, tenofovir.

Interventions and their incremental cost-effectiveness ratio (ICERs) were estimated and compared using the following formula (56):

$$ICER_{TDF vs. other antivirals} = \frac{Cost_{TDF} - Cost_{other antivirals}}{QALY/LYG_{TDF} - QALY/LYG_{other antivirals}}$$ (2)

3.11. Uncertainty Analysis

In this study, PSA was performed to evaluate the effects of uncertainty on the parameters included in the model. The probability distributions of all the parameters entered into the model are presented in Tables 1 and 2. A second-order Monte Carlo simulation was performed using 5000 trials to analyze the probabilistic sensitivity. The PSA results are presented using cost-effectiveness acceptability curves and a scatter plot of the incremental cost-effectiveness. Due to the lack of an explicit WTP threshold in Iran, based on the proposal of the world health organization for developing countries, a WTP per every QALY of one to three times the gross domestic product per capita was used (57). In Iran, the value in 2014 when its threefold value was about $14010 was $4670 dollars.
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Table 3. Comparison of the Cost-Effectiveness of the Five Drug Strategies in Patients With HBeAg-Negative CHB Based on the Number of QALY Using the Markov Microsimulation Model

| Strategy | Total Cost (USD) | LYG | QALY | Incr. Cost (USD) | Incr. QALY | ICER Per QALY | Subset |
|----------|-----------------|-----|------|-----------------|-----------|---------------|--------|
| TDF      | 47,257          | 21.26 | 11.60 | 0               | 0         | 0             | Dominated |
| LAM      | 64,996          | 17.69 | 10.93 | 16,839          | -2.67     | -6,800        | Dominated |
| ADV      | 69,510          | 18.59 | 11.51 | 22,251          | -2.09     | -10,642       | Dominated |
| Lam + Adv | 73,896         | 18.31 | 11.34 | 26,373          | -2.27     | -10,743       | Dominated |
| ETV      | 139,532         | 18.92 | 11.47 | 92,275          | -0.13     | -18,527       | Dominated |

Abbreviations: HBeAg, hepatitis B e antigen; Incr., incremental, ICER, incremental cost-effectiveness ratio; LYG, life-years gained; QALY, quality-adjusted life-years; TDF, tenofovir; LAM, lamivudine; ADV, adefovir; ETV, entecavir.

4. Results

4.1. Base-Case Analysis

In this study, 155 HBeAg-negative CHB patients who were referred to selected centers were enrolled in the study and classified by age, sex, and duration of disease in the initial model. The mean age of the total sample was 40 y, 67% of the sample were males, and 33% were females.

The results of the cost-effectiveness analysis obtained using the Markov microsimulation model and based on QALY and LYG showed that TDF was less expensive and more effective than the other medication strategies. Hence, TDF was the most cost-effective of the medication strategies (ADV, ETV, and LAM and ADV + LAM) (Figure 2).

Table 3 presents the cost-effectiveness (QALY and LYG), incremental costs, incremental effectiveness, and ICER, as well as the ratings and status of domination of the five drug therapy strategies in patients with HBeAg-negative CHB. As shown in the table, compared with TDF, the other strategies (LAM, ADV, ADV + LAM, and ETV) were associated not only with lower QALY and LYG but also with higher additional costs. During a patient’s lifetime, compared with the alternatives, TDF had the lowest cost and highest expected QALY and LYG. ETV had the highest cost per every unit of QALY and LYG when compared with the other strategies.

4.2. Uncertainty Analysis

In this study, PSA was used to measure the effects of uncertainty in the parameters on the results of the model. Using 5000 trials, a second-order Monte Carlo simulation and PSA were conducted. The uncertainty of the results based on the range of WTP thresholds, cost-effectiveness acceptability curves, and a scatter plot of the incremental cost-effectiveness are presented in Figures 3 and 4.

An acceptability curve is one of the best curves for planning and policymaking. It can help policymakers and health system planners determine the cost-effectiveness of every intervention per payment. The results of the acceptability curves showed that TDF was the most cost-effective treatment strategy in 59% - 78% of simulations for thresholds below $14010 for patients with HBeAg-negative disease (Figure 3).

By showing what percentage of points is located in the accepted area (below the threshold), scatter plots present more detailed information for comparisons between single drugs. The scatter plot results are presented only for TDF compared with the alternatives. As compared with LAM, ADV, ETV, and ADV + LAM, the results demonstrated that TDF was in an acceptable area below the threshold in 85%, 84%, 62%, and 83% of cases in patients with HBeAg-negative disease. Hence, it was the best of the other alternatives and the most cost-effective strategy (Figure 4). Therefore, the results of the sensitivity analysis did not change the status of TDF as the most effective treatment. This finding is an indication of the robustness of the results of the study.

5. Discussion

This is the first comprehensive economic evaluation study of hepatitis B patients in Iran and the Eastern Mediterranean region. The study aimed to evaluate oral nucleos(t)ide analogue strategies for HBeAg-negative CHB and its related complications from a pharmacoeconomic societal perspective.

In the present study, the ICERs based on QALY and LYG in patients with HBeAg-negative CHB showed that of the five therapeutic strategies (ADV, LAM, ADV + LAM, ETV, and TDF) used in Iran, TDF was more effective and less expensive. This finding is in line with the results of He et al.’s study (12) and Tantai et al.’s study (58). Despite some minor differences, the results of this study are also consistent with those of previously published economic evaluation studies of HBV, such as the CUA study by Dakin et al. in the U.K. (13) and Buti et al.’s study (56) in Spain. The results of the present study demonstrated that TDF in patients with HBeAg-negative CHB had the lowest expected cost and highest expected QALY and LYG in a lifetime span. The results of this study are consistent with those of Tantai et al. (58).

The results of the PSA revealed that TDF was less expensive and more effective than the other drug therapy
The cost-effectiveness acceptability curve, which was the output curve of PSA, also showed that TDF was the most cost-effective therapy in 59% - 78% of simulations in HBeAg-negative cases for thresholds below $14010. The results of this study are consistent with those of an earlier study by He et al. (12), who reported that TDF was the most effective strategy in 65% - 78% of cases.

Furthermore, as shown by the results of the incremental cost-effectiveness scatter plot, which was another output of the PSA, in most cases, compared with the other alternatives, TDF was in an acceptable area below the threshold. In patients with HBeAg-negative CHB, compared with the alternatives (LAM, ADV, ETV, and ADV + LAM), TDF was in acceptable area below the threshold in 85%, 84%, 62%, and 83% of cases and was superior to the other strategies. The results of the present study are in line with those of He et al. (12).

In addition, the results obtained from the estimation of QALY and cost savings associated with the use of TDF in HBeAg-negative patients compared with the alternatives showed that it reduced the progress of the disease and serious complications in the future. In addition, it led to cost savings and a very significant increase in the number of QALYs. The results of this study are consistent with those of studies conducted by Jing et al. and Tantai et al. (12, 58). TDF seems to be the optimal drug regimen to stop or slow the progression of liver disease. Therefore, TDF, which was identified as the most cost-saving strategy in this study and

**Figure 2.** Comparison of the Cost-Effectiveness of the Five Drug Strategies in Patients With HBeAg-Negative CHB Based on LYG and QALY

**Figure 3.** Cost-Effectiveness Acceptability Curve Obtained Through a Monte Carlo Simulation

strategies and that it achieved the best results. It also confirmed that TDF was superior to all the other drug treatment strategies. As the results of the PSA did not alter the status of TDF as the most effective treatment, the results of the present study seem to be robust. The findings of the current study are in line with those of studies by He et al. in Canada, Wiens in Brazil, and Tantai et al. in Thailand (12, 16, 58).
some previously published studies (12, 16, 58), should be the first choice in cases of LAM or ADV drug resistance.

Most of the drugs in the present study are manufactured in Iran and therefore do not require high expenditures by either patients or the health system. However, given the better response of TDF compared with the other drugs, it can prevent and reduce high costs associated with complications (i.e., cirrhosis and cancer) that emerge in severe stages of the disease. A regular review of the literature after the introduction of new drugs can ensure that the dominant drugs are excluded and replaced by the dominant drugs. This will benefit all those involved in the health system (i.e., the government, insurance industry, and patients).

The strengths of this study are as follows: The model included both direct medical and various indirect nonmedical costs, and it employed the Monte Carlo Markov method for microsimulation modeling. In addition, the data on the costs and effectiveness were collected locally and based on observations, patients’ self-reports, and experts’ opinions. Furthermore, the values of the country were used for the first time to convert the 5-digit code numbers of the
questionnaire to numerical utility. An additional strength of the model used in the study was the ability to switch to another drug therapy for those patients who gave up drug therapy. This ability distinguishes the present model from models used in previous studies. Finally, for the first time, a microsimulation approach was used to assess the cost-effectiveness of antiviral drugs in patients with chronic HBV infection.

One of the limitations of this study was the absence of data in some cases, especially data on the transitional probabilities of the disease. Hence, a constant rate was used in this study. Although the results showed that TDF was the superior and most cost-effective option for HBeAg-negative patients, to generalize the results to other settings, various factors could influence the findings. These include the epidemiology of the disease, demographics of the study population, resource availability, costs, valuing outcomes by individuals, thresholds, and the use of indicators of effectiveness. However, as this study included a range of important features of the patients, including age, sex, and duration of the disease, and the data were extracted from several main referral centers in the country, many of the results can likely be generalized.

In conclusion, according to the results of this study, among nucleos(t)ide analogues evaluated, TDF was the optimal and most cost-effective strategy for patients with CHB. Hence, it can be used as first-line therapy and for patients switching from other drugs. Furthermore, as TDF can reduce treatment times and manage drug resistance, TDF can be recommended as first-line therapy for patients with CHB.

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Footnote

Authors’ Contribution: Khosro Keshavarz: gathering the data, analyzing the collected data, and contributing to the design of the research method; Abbas Kebriaeizadeh, Seyed Moayed Alavian, and Shekoufeh Nikfar: designing the research method, revising the manuscript, and final approval of the data; Ali Akbari Sari and Farhad Lotfi: designing the research method and revising the manuscript; Maryam Keshvari, Mohsen Rezaei Hemami, Amir Hashemi Meshkini, and Mehdi Javanbakht: data gathering and drafting the manuscript; All the authors read and approved the final manuscript.

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