Effectiveness and Safety of Tenofovir Disoproxil Fumarate in Chronic Hepatitis B: A 3-Year, Prospective, Real-World Study in France

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Abstract
Background and Aims Tenofovir disoproxil fumarate (TDF) demonstrated potent and sustainable antiviral efficacy and a good safety profile in patients with chronic hepatitis B (CHB) in controlled clinical trials. Real-world data are important to confirm effectiveness and safety data in patient populations encountered in routine clinical practice.
Methods This non-interventional, prospective, 36-month study included treatment-naïve and treatment-experienced patients with CHB initiating their first TDF regimen (monotherapy or combination therapy) in routine clinical practice in France. Clinical, virologic, biochemical, compliance, and safety data were collected.
Results Data from 440 consecutive patients from 58 centers were analyzed. The majority of the cohort was male
(71 %), hepatitis B “e” antigen-negative (HBeAg–) (74 %), and treatment-experienced (56 %); 11 % were aged ≥65 years; and comorbidities were reported in 39 %. After 12 months, 92 % of the overall cohort achieved virologic response (HBV DNA <69 IU/mL) which was maintained to 36 months (96 %); virologic response was achieved by >90 % of patients irrespective of HBeAg status, age, or prior treatment history. At 36 months, 77 % of patients had normal alanine aminotransferase levels. Fourteen patients lost hepatitis B surface (HBs) antigen, and seven seroconverted to anti-HBs. TDF was well tolerated over the 36-month study, including in 14 women who became pregnant during the study. Median estimated glomerular filtration rate did not change markedly from baseline irrespective of prior treatment history.

Conclusions TDF demonstrated potent virologic and biochemical responses across a broad range of patients reflective of routine clinical practice. The safety profile was consistent with results from pivotal trials.

Keywords HBV · Tenofovir · Routine practice · Real-world

Introduction

Chronic infection with hepatitis B virus (HBV) represents a major global health problem, responsible for around 780,000 deaths worldwide, largely through serious liver-related sequelae [1]. Chronic HBV infection (CHB) prevalence varies widely, being highest in sub-Saharan Africa and East Asia, where 5–10 % of the adult population is chronically infected. HBV infection is becoming a public health concern in countries where prevalence was previously reduced due to factors including increased migration from countries with intermediate/high HBV prevalence [2]. Although recent data on HBV prevalence in France are lacking, a 2004 national survey estimated hepatitis B surface antigen (HBsAg) prevalence was 0.68 %, corresponding to 280,821 chronic carriers [3]. A 2001 French survey of patient mortality related to hepatitis B and C estimated that 1507 deaths were attributed to HBV infection, corresponding to 2.5 deaths per 100,000 persons [4].

Many studies have demonstrated the correlation between viral load and the risk of developing cirrhosis and hepatocellular carcinoma (HCC) [5–8]. As it is not possible to eradicate HBV [9], CHB treatment aims to provide sustained suppression of HBV replication, thus preventing disease progression and death. Tenofovir disoproxil fumarate (TDF) is a potent nucleotide analog recommended as a first-line therapy for CHB in international and national guidelines [9–11]. Pivotal randomized controlled trials have reported high efficacy and a favorable safety profile with TDF treatment for up to 8 years in treatment-naive and treatment-experienced patients [12–15], with associated fibrosis regression and cirrhosis [14]. No drug resistance to TDF has been observed.

Published data from real-world studies with TDF, which include the highly diverse patients encountered in routine practice, are lacking, and are mostly retrospective, but are required in order to confirm and expand upon the results of clinical trials [16, 17]. The prospective, non-interventional, multicenter VIREAL study investigated the use of TDF in routine clinical management of CHB in France, and evaluated its effectiveness and safety in a “real-world” cohort of patients.

Patients and Methods

VIREAL is a multicenter, non-interventional, prospective study to assess the long-term effectiveness and safety of TDF treatment in a real-life cohort of patients with CHB managed in routine clinical settings in France. Participating sites included teaching hospitals, general hospitals, and private practices specializing in CHB to provide a representative cross section of CHB management. Patients were enrolled at 58 centers across France. Recruitment took place between June 2009 and April 2010, and the database was locked in October 2013.

The study was conducted in accordance with Good Epidemiological Practice guidelines, was approved by the French Medical Board, and was authorized by the National Computers and Privacy Commission. Patients gave written consent for the collection of anonymized medical data from their medical files.

Study Population

The study enrolled treatment-naive or treatment-experienced patients ≥18 years of age with hepatitis B “e” antigen (HBeAg)-negative (HBeAg–) or HBeAg-positive
(HBeAg+) CHB, initiating a first prescription of TDF at enrollment. TDF treatment initiation was at the treating physician’s discretion. Key exclusion criteria included prior treatment with TDF, evidence of HCC, or co-infection with human immunodeficiency virus (HIV), hepatitis C virus, or hepatitis D virus.

Study Assessments and Data Collection

Data on safety and on the clinical, laboratory, and virologic course of TDF-treated patients from the routine testing/evaluations conducted by the treating physician were captured. Study-specific laboratory tests, resistance monitoring, or prespecified visits were not required. As visit times varied between practices, patient assessments were assigned a nominal visit month of 3, 6, 12, 18, 24, 30, and 36 months based on the number of days since the baseline visit within a visit window of ± 3 months of the nominal visit month. Data were collected via an electronic case report form and included the following: patient demographics; anti-HBV treatment history; viral genotype; serum HBV DNA levels; HBeAg and HBsAg; alanine aminotransferase (ALT); other routine laboratory parameters; renal function tests (serum creatinine, creatinine clearance [CrCl], estimated glomerular filtration rate [eGFR] calculated by Cockcroft–Gault formula at the beginning of the study which was replaced by the Chronic Kidney Disease Epidemiology Collaboration formula); liver histology and progression of liver-related complications; resistance to anti-HBV therapy; adverse events (AEs) and serious AEs; pregnancy; adherence to therapy; and therapy interruptions.

Virologic response was defined as HBV DNA <69 IU/mL. Adherence was assessed using a six-question questionnaire completed by the patient at each visit, which was first applied in the field of hypertension [18]. The total number of “yes” answers defined the level of adherence: 0 = good; 1–2 = minor problems; and >3 = poor adherence.

Statistical Analysis

Mean, standard deviation (SD), median, range, or first and third quartile (Q1, Q3) were calculated for continuous variables, together with the total number of observations and the number of non-missing and missing values. For categorical variables, number and percent of patients are reported. Missing data were excluded from the efficacy analysis. The safety analysis included all treated patients.

Comparative analyses were performed using Pearson’s Chi-square test for qualitative variables, and Student’s t test for analyses of variance for quantitative variables. Non-parametric tests were used in the case of a non-normal distribution. Statistical analysis was performed using SAS® (Version 9.1).

Results

In total, 440 patients were prospectively enrolled. Of these, 328 completed the study, and 112 discontinued the study [49 (46.7 %) patients were lost to follow-up; 27 (24.1 %) were discontinued at the investigator’s discretion; 12 (10.7 %) patients made the decision to discontinue; 10 patients died (8.9 %); seven (6.3 %) patients discontinued due to an adverse drug reaction; and seven (6.3 %) were withdrawn as they did not fit the inclusion/exclusion criteria].

The overall population was predominantly male and around three-quarters were HBeAg– (Table 1). The largest proportion of patients was of European decent (38.6 %). Mode of HBV infection was unknown in almost half of patients (46 %). Where this was known (n = 237), the most commonly cited routes were the following: at birth or during early childhood (54.4 %); family exposure (18.1 %); nosocomial infection (10.5 %); sexual exposure (6.3 %); and drug use (2.5 %).

Just over half of the patients (n = 258) were treatment-experienced, with around two-thirds having previously received an adefovir (ADV)-containing regimen: monotherapy or in combination with lamivudine (LAM) (Table 2). The majority (179/258) of the treatment-experienced patients were switched directly to TDF. Seventy-nine patients were not receiving treatment at baseline and the washout period of previous treatment was variable. Of the treatment-experienced patients, 64 (24.8 %) terminated previous therapy and 38 (14.7 %) initiated TDF due to persistent viremia/suboptimal response and viral relapse (defined as a confirmed increase in HBV DNA level of more than 1 log_{10} IU/mL compared with the nadir [lowest value] HBV DNA level on therapy) [9]. Mean baseline HBV DNA levels were lower in treatment-experienced patients than treatment-naïve patients (Table 2). A higher proportion of treatment-experienced patients had undetectable viral load (HBV DNA <69 IU/mL) at baseline compared with treatment-naïve patients (61.1 vs. 5.5 %, respectively, Table 2). Viral genotype was known for only 7 % of patients so is not included in the analysis (Table 1).

Baseline ALT levels varied widely [median (range) 35 (7–828) IU/mL]. At baseline, more than half of patients had normal ALT (ALT < upper limit of normal) (Table 2). The proportion of patients with normal ALT was higher in treatment-experienced compared with treatment-naïve patients (71.3 vs. 35.3 %, respectively), and in HBeAg– compared with HBeAg+ patients (59.5 vs. 42.9 %, respectively).
Around two-thirds of patients had a report of liver biopsy at baseline. The majority had no/minimal fibrosis (METAVIR score F0–F1) or moderate fibrosis (F2); 14.4% had cirrhosis (F4). Clinical signs of cirrhosis were reported in 17 patients (co-existing signs were possible): ascites (n = 4), esophageal varices (n = 13), and splenomegaly (n = 10).

The most commonly reported comorbidities were hypertension and diabetes mellitus; prior renal insufficiency was reported in 14 patients, and 14 patients had a history of liver (n = 11) or renal (n = 3) transplantation. In all, 122 patients (27.7%) had baseline eGFR <90 mL/min/1.73 m², with 24 of these patients having a baseline eGFR <60 mL/min/1.73 m²; no patient had baseline eGFR <30 mL/min/1.73 m².

### TDF Treatment

At study initiation, most treatment-naïve patients (91.2%) received TDF as monotherapy, with the remainder receiving TDF in combination with LAM. The majority of treatment-experienced patients (63.4%) received initial treatment with TDF as monotherapy, with the remainder receiving TDF in combination with entecavir (ETV) (19.8%) or LAM (21.3%). Most patients remained on TDF monotherapy for the entire study duration, with 70 receiving add-on medication, predominantly ETV (n = 15) or LAM (n = 45). Overall 17 patients switched to an alternative treatment (ETV n = 14; LAM n = 3).

### Virologic Response

By month 12, 92% of the overall population with available data (n = 308) had HBV DNA <69 IU/mL, rising to 96% at 36 months (n = 259). A greater proportion of HBeAg– patients compared with HBeAg+ patients achieved HBV DNA <69 IU/mL by month 12 (90.0 vs. 80.3%, respectively). Irrespective of HBeAg status, treatment history, or whether patients received TDF as monotherapy or in combination, >90% of patients with available data achieved HBV DNA <69 IU/mL by month 36. A total of 27 patients had one or more resistance-associated variants (RAVs) at baseline, including those associated with LAM and ADV. Three of these patients had HBV DNA <69 IU/mL at baseline and final visit; all remaining patients achieved a virologic response during treatment, the majority by month 3, and maintained this response to their last visit. Of the 18 patients with resistance-associated mutations at baseline and HBV DNA levels available at baseline and final visit, 13 had undetectable HBV DNA at the final visit. No correlation was observed between adherence score and virologic response. Older age did not appear to affect response; of the 26 patients aged ≥65 years at baseline who completed 36 months of treatment, all (100%) achieved HBV DNA

| Characteristic | N = 440 |
|----------------|--------|
| Age, years, mean (SD) | 45.3 (14.3) |
| ≥65 years, n (%) | 48 (10.9) |
| Male, n (%) | 312 (70.9) |
| Geographic descent, n (%) | |
| Europe | 170 (38.6) |
| Middle East | 21 (4.8) |
| Asia Pacific | 101 (23.0) |
| North Africa | 44 (10.0) |
| Sub-Saharan Africa | 87 (19.8) |
| North and South America | 11 (2.5) |
| Unknown | 6 (1.4) |
| Genotype, n (%) | |
| A | 4 (0.9) |
| B | 5 (1.1) |
| C | 7 (1.6) |
| D | 9 (2.0) |
| E | 5 (1.1) |
| Missing | 410 (93.2) |
| Patients with cirrhosis, n (%)<sup>a</sup> | 68 (15.5) |
| Liver biopsy performed, n (%) | 298 (67.7) |
| Fibrosis stage (METAVIR), n (%) | |
| F0–F1 | 105 (36.5) |
| F2 | 88 (30.6) |
| F3 | 53 (18.4) |
| F4 | 42 (14.6) |
| Missing | 10 |
| Comorbidities, n (%)<sup>b</sup> | |
| Hypertension | 48 (10.9) |
| Diabetes mellitus | 40 (9.1) |
| Cardiovascular disease | 16 (3.6) |
| Rheumatic disease | 15 (3.4) |
| Renal insufficiency/dysfunction | 14 (3.2) |
| Malignant disease | 10 (2.3) |
| Neuropsychiatric disease | 9 (2.1) |
| Other | 104 (23.6) |

<sup>a</sup> History of cirrhosis as defined by the investigator; method not further specified

<sup>b</sup> Where reported (n = 170). Multiple comorbidities were possible
Among 167 patients with HBV DNA <69 IU/mL at baseline, 13 patients experienced virologic breakthrough (defined as patients with baseline HBV DNA <69 IU/mL and an HBV DNA of ≥69 IU/mL for at least one post-baseline visit) over the course of the study. The annual rate of virologic breakthrough was 3.6, 3.6, and 0.6 % at 12, 24, and 36 months, respectively. For 11 of these 13 patients, breakthrough was transient. All patients with virologic breakthrough continued TDF and completed the study. No baseline RAVs were reported for the 13 patients with virologic breakthrough. A minor problem of compliance was reported for four patients, and the treatment adherence questionnaire was not completed for eight patients.

### Biochemical Response

Mean (SD) ALT improved from 67.2 (95.07) IU/mL at baseline to 31.3 (18.01) IU/mL at month 36. The proportion of patients with normal ALT levels increased steadily to 76.8 % (74.6 % treatment-naïve; 77.7 % treatment-experienced) by 12 months, and this proportion was maintained to 36 months (78.0 % overall; 75.0 % treatment-naïve; 79.4 % treatment-experienced).
Serologic Response

Of the 80 patients who were HBeAg+ at baseline with ≥1 additional HBeAg status test at some stage during the study, 36 (45 %) lost HBeAg and 20 achieved HBeAg seroconversion during the study (five additional patients were anti-HBe+ at baseline). Three patients who lost HBeAg and seroconverted to anti-HBe+ during the study reconverted and were HBeAg+, anti-HBe− at their final study visit.

During the study, 14 patients (3 %) lost HBsAg. (Three HBeAg+ at baseline, 10 HBeAg−, and one with unknown HBeAg status at baseline). Of the 14 patients with HBsAg loss, five were treatment-naïve and nine were treatment-experienced (ADV + LAM, n = 4; ADV + LAM + -PEG-IFN, n = 2; ADV + PEG-IFN, n = 2; ADV + ETV, n = 1). The cumulative rate of HBsAg loss was 0.96 % in the first year, 2.89 % in year 2, and 3.37 % in year 3. Seven patients (Five HBeAg−) achieved HBsAg seroconversion during the course of the study. The
cumulative rate of seroconversion was 0.72 % in the first year, 1.20 % in year 2, and 1.68 % in year 3.

Safety

TDF was well tolerated over the 36-month study. In total, 41 patients (9.3 %) reported 68 AEs considered by the physician to be related to TDF (Table 4). There was no difference in the proportions of treatment-naive and treatment-experienced patients reporting adverse events (8.2 vs. 10.1 %, $P = 0.711$). Of the 16 serious AEs reported in 16 patients, nine were considered to be related to TDF (visual impairment, nausea, asthenia, gait disturbance, weight decrease, muscular weakness, musculoskeletal pain, depression, psoriasis, all $n = 1$).

Table 4 Adverse events considered by the physician to be related to TDF

| Patients (n) | Number of adverse events/patient |
|-------------|----------------------------------|
|             | 1                                | 27 |
|             | ≤2                               | 36 |
|             | ≤3                               | 39 |
|             | ≤8                               | 41 |
| Serious adverse events$^a$ | 9 | |
| Adverse events occurring in >1 patient | | |
| Abdominal pain$^b$ | 8 |
| Asthenia | 7 |
| Nausea | 6 |
| Vomiting | 5 |
| Diarrhea | 5 |
| Hypophosphatemia | 4 |
| Headache/migraine | 2 |
| Renal related | | |
| Abnormal renal function tests$^c$ | 4 |
| Renal failure | 3 |
| Renal impairment | 2 |
| Renal tubular disorder | 2 |
| Muscle spasms | 2 |
| Discontinuation of TDF due to adverse events$^d$ | 23 |

$^a$ Serious adverse events comprised visual impairment, nausea, asthenia, gait disturbance, weight decrease, muscular weakness, musculoskeletal pain, depression, psoriasis (all $n = 1$)

$^b$ Abdominal pain includes the MedRA preferred terms abdominal pain upper ($n = 1$) and abdominal pain ($n = 7$)

$^c$ Abnormal results included creatinine renal clearance decreased ($n = 2$) and blood creatinine increased ($n = 2$)

$^d$ Reasons for discontinuation in >1 patient (multiple reasons per patient were possible): nausea ($n = 5$) vomiting ($n = 4$); asthenia ($n = 3$); renal failure ($n = 3$); diarrhea ($n = 2$); abdominal pain ($n = 2$); renal impairment ($n = 2$); renal tubular disorder ($n = 2$)

There were 10 deaths during the study [HCC $n = 1$; leukemia $n = 2$; breast cancer $n = 1$; suicide post-liver transplantation $n = 1$; stroke $n = 1$; post-liver transplantation complications $n = 1$; cholangiocarcinoma $n = 1$; and end-stage liver disease (hepatic encephalopathy $n = 1$; spontaneous bacterial peritonitis $n = 1$)]. No deaths were considered to be related to TDF.

Sixteen pregnancies were reported during the study. The majority of these patients had become pregnant while taking TDF with one case where TDF was administered prophylactically. Follow-up data were available for 14 cases. Of these, the mean age of the mothers was 29 years, and 57 % ($n = 8$) were HBeAg+; META VIR fibrosis scores (where available) were F0 in one patient, F1 in three patients, and F2 in six patients. Median duration of TDF exposure during pregnancy was 35 weeks (range 5–39 weeks). Therapy was interrupted in four patients (obstetricians’ decision $n = 2$; patient’s decision $n = 1$; HBsAg loss $n = 1$) at weeks 12, 16, 23, and 25 of pregnancy. One patient discontinued treatment at week 5 (obstetrician’s decision). Two HBeAg+ mothers had HBV DNA $>10^6$ IU/mL at delivery (one TDF discontinuation; one non-adherence). Vaccination and immunoprophylaxis according to the French guidelines were recommended for all newborns. Median gestational age at delivery was 39 weeks (range 34–40 weeks). No AEs related to TDF were observed, no neonatal complications arose, and no birth defects were reported. Complications during pregnancy, labor, or postpartum were: $n = 1$; premature labor controlled by medication and $n = 2$; postpartum hemorrhage. Five patients reported breastfeeding, with three breastfeeding while receiving TDF without any adverse consequences for the babies up to 1 year. No infant was reported to be HBsAg+ at 9 months of age. Of five infants with anti-HBs testing, all were anti-HBs+ (84–308 IU/mL).

During the course of the study, median change in CrCl and serum creatinine remained relatively stable over 36 months of treatment, regardless of prior treatment (Table 5). Serum phosphorus levels were also stable over the study period.

Mean eGFR declined slightly over the treatment period (Table 6). Older patients ($\geq 65$ years) and patients previously treated with ADV-containing regimens had lower eGFR at baseline, which declined slightly or remained stable over the treatment period (Fig. 2a). When subdivided by baseline eGFR, mean eGFR declined slightly over time in patients with baseline CrCl $\geq 90$ mL/min/1.73 m$^2$, and increased slightly in patients with mild or moderate renal impairment at baseline (Fig. 2b). Overall, 65 (15 %) had a decline in eGFR of $\geq 20$ % versus baseline and 26 patients (6 %) had a decline in eGFR of $\geq 30$ % versus baseline (Table 7). Prior treatment did not have a
significant effect on the proportion of patients with at least a 20 or 30% decline in eGFR. However, significantly fewer patients in the treatment-naive versus treatment-experienced group and in the group with no prior ADV therapy versus prior ADV therapy had an eGFR value of \(<60\text{ mL/min/1.73 m}^2\) for at least one post-baseline visit (\(P < 0.001\) for both).

Renal endpoints were observed in approximately 3% of patients: five patients had a serum creatinine increase of at least 0.5 mg/dL from baseline and two patients had a decreased creatinine clearance. Three cases of renal failure, two cases of renal impairment, and two cases of renal tubular dysfunction were reported as mild or moderate AEs. None of these patients had a reported history of renal impairment. One patient had been treated previously with ADV and LAM, one had been treated previously with ADV, two had been treated previously with ADV and LAM with LAM therapy continuing during the study. One patient had diabetes, two had hypercholesterolemia, and two were of older age (67 years and 72 years). TDF was discontinued in all patients. No renal-related serious AEs were reported.

**Major Liver-Associated Events**

Two cases of confirmed HCC and three cases of suspected HCC arose during the course of the study (months 3 and 12, and months 24 and 36, respectively). All patients were reported to have cirrhosis at baseline; both patients with confirmed HCC also had clinical signs or symptoms of advanced liver disease at baseline (esophageal varices in one patient; esophageal varices and ascites in the second). Two patients died due to complications of end-stage liver disease (hepatic encephalopathy \(n = 1\); spontaneous bacterial peritonitis \(n = 1\)). Neither death was considered to be related to TDF.

**Discussion**

In this study of CHB patients managed in routine clinical practice in France, treatment with TDF resulted in sustained virologic suppression. Virologic suppression was maintained over 36 months, irrespective of HBeAg status
or treatment history. In this diverse patient population, treatment was well tolerated with a safety profile similar to that reported in clinical trials. In addition, efficacy data from this study compare favorably to those reported in the pivotal clinical trials of TDF in CHB.

### Table 6 Change in estimated glomerular filtration rate from baseline according to prior treatment status

| Treatment group | Baseline, \( (n) \) | Change in eGFR (CKD-EPI) from baseline to: |
|-----------------|----------------------|------------------------------------------|
|                 | Month 12, \( n \), median (range) (mL/min/1.73 m\(^2\)) | Month 24, \( n \), median (range) (mL/min/1.73 m\(^2\)) | Month 36, \( n \), median (range) (mL/min/1.73 m\(^2\)) |
| Overall         | 360                  | 249 (−41.5, 38.54) \( P = 0.044 \) | 217 (−61.03, 38.76) \( P = 0.681 \) | 204 (−48.78, 34.91) \( P = 0.033 \) |
| Treatment-naive | 143                  | 91 (−37.91, 22.00) \( P = 0.015 \) | 79 (−36.02, 29.86) \( P = 0.240 \) | 75 (−48.78, 34.54) \( P = 0.009 \) |
| Treatment-experienced | 217                  | 158 (−41.05, 38.54) \( P = 0.48 \) | 138 (−61.03, 37.86) \( P = 0.700 \) | 129 (−40.78, 34.91) \( P = 0.536 \) |
| Prior ADV       | 157                  | 117 (−41.05, 38.54) \( P = 0.534 \) | 101 (−61.03, 37.86) \( P = 0.762 \) | 101 (−38.50, 33.87) \( P = 0.888 \) |
| No prior ADV    | 203                  | 132 (−37.91, 33.18) \( P = 0.032 \) | 116 (−50.07, 29.86) \( P = 0.327 \) | 103 (−48.78, 34.91) \( P = 0.008 \) |

ADV adefovir, CKD-EPI chronic kidney disease epidemiology collaboration, eGFR estimated glomerular filtration rate

\( P \) values calculated using Fisher’s exact test (two-sided)

### Fig. 2 Mean estimated glomerular filtration rate. a By age and treatment history; b by baseline creatinine clearance (CrCl) category. ADV adefovir, CKD-EPI chronic kidney disease epidemiology collaboration, CrCl creatinine clearance, eGFR estimated glomerular filtration rate

The diversity of the patient population in the VIREAL cohort compared with that in clinical trials is reflected in the wide variation of baseline parameters including ALT and viral load, the varied geographical origin of the patients, and the high proportion of patients with reported...
comorbidities. Despite this, the efficacy data from the current study compare favorably to those reported in the pivotal clinical trials of TDF in CHB. Following the initial randomized study period of 48 weeks, patients included in the pivotal clinical trials of TDF were enrolled in an open-label study and treated for 7 additional years. After 26 months of treatment with TDF, 99% of HBeAg– patients and 93% of HBeAg+ patients achieved HBV DNA ≤69 IU/mL [13] in the on-treatment population, which is similar to that included in VIREAL. Further data have demonstrated that virologic response is maintained to 5 years, with 99% of HBeAg– patients achieving HBV DNA ≤29 IU/mL [14]. Recently presented 8-year data from this study show that this response is maintained, with a good safety profile and no evidence of virologic resistance [15]. Our data are similar to those reported in a prospective/retrospective real-world European study including 374 treatment-naïve patients, where 97% of patients had undetectable HBV DNA (PCR negative) after 4 years of TDF treatment [16]. Patients in this European cohort and the majority of patients in the pivotal clinical trials were treatment-naïve whereas around half of the patients in the VIREAL cohort were treatment-experienced.

By month 36 of the current study, 94% of treatment-experienced patients achieved HBV DNA ≤69 IU/mL, including all 18 patients with resistance-associated variants at baseline. These results are consistent with data from a randomized trial of TDF or TDF/emtricitabine in patients who previously had an incomplete response to ADV [19]. This study demonstrated that response was not influenced by the presence of ADV- or LAM-associated mutations. In contrast, retrospective analysis of patients treated in Germany and The Netherlands found that although LAM resistance did not influence antiviral response, the presence of ADV resistance impaired TDF efficacy (100% vs. 52% probability of HBV DNA ≤69 IU/mL, respectively) [20]. The variability in these results demonstrates that further research is required before the relationship between ADV resistance and TDF response is fully understood.

Numerous studies have shown an association between HBV DNA levels and disease progression [5, 8] and that long-term HBV DNA suppression can reduce serious liver-related sequelae [14, 21] reinforcing the importance of HBV DNA suppression in the treatment of CHB. Overall, 78% of VIREAL patients had normal ALT levels after 36 months of TDF treatment, again similar to that seen in pivotal trials [3, 12, 15]. Serologic response in VIREAL was also similar to that seen in clinical trials, with 45% of HBeAg+ patients losing HBeAg in VIREAL compared with 34% in the clinical trial population after 3 years [13]. In all, 14 patients in VIREAL lost HBsAg during the course of the study, and seven of these patients also seroconverted to anti-HBs. Interestingly, these patients were predominantly HBeAg– at baseline, while all except one patient who showed HBsAg loss/seroconversion in the clinical trial population were HBeAg+. This may partly result from the proportion of HBeAg– patients in this cohort compared with the clinical trial population.

TDF was well tolerated over 36 months in VIREAL, with a safety profile similar to that reported in clinical trials. Treatment-related serious AEs were low, and the majority of patients stayed on TDF treatment until their last visit. There was no statistical difference in the number of AEs related to TDF reported in treatment-naïve and treatment-experienced patients. Although there was a slight decline in mean serum creatinine, CrCl, and eGFR, statistical comparison to baseline should be viewed with caution due to the high proportion of missing data. Furthermore, a decrease in eGFR is expected in patients over time due to aging. We observed

| Table 7 | Proportion of patients with a decline in estimated glomerular filtration rate of ≥20% compared with baseline, ≥30% compared with baseline and eGFR values ≤60 mL/min/1.73 m² |
|------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
| Proportion of patients with eGFR decline ≥20% versus baseline, n (%) | Overall | Treatment-naïve n = 182 | Treatment-experienced n = 258 | Prior ADV n = 179 | No Prior ADV n = 261 |
| cGFR ≤60 mL/min/1.73 m², n (%) | 65 (15) | 22 (12) | 43 (17) | 31 (17) | 34 (13) |
| P | 0.220 | 0.308 | 0.001 | 0.221 | 1.00 |

ADV adefovir, eGFR estimated glomerular filtration rate
P values calculated using Fisher’s exact test (two-sided)
that significantly fewer treatment-naïve patients had an eGFR value of <60 mL/min/1.73 m² compared with treatment-experienced patients. This could be due to the fact that treatment-experienced patients have received treatment for a longer duration than patients who were receiving TDF as their first treatment. While serum creatinine levels are commonly used markers of renal function, other urinary and serum proteins may be earlier, more accurate markers of renal toxicity [22]. However, testing of such biomarkers is not routinely used in clinical practice, and creatinine clearance was therefore used in this real-life study to monitor renal function. In all, only seven cases of renal AEs were reported, all of which were of mild-to-moderate intensity, and were managed by TDF discontinuation.

Our study adds to the increasing data on the use of TDF during pregnancy. In VIREAL, treatment with TDF during pregnancy, including during the first trimester, was not associated with any AEs during pregnancy, complications during labor, neonatal complications, or birth defects. TDF has been assigned pregnancy category B by the Food and Drug Administration, compared with category C for LAM, ADV, and ETV. A recent review examining data on the use of TDF in pregnancy in both HIV and HBV infection [23] together with that from the 1800 pregnancies included in the Antiretroviral Pregnancy Registry, concluded that the safety data for TDF during pregnancy are generally reassuring for pregnancy outcomes and for lack of congenital or other severe anomalies in exposed infants.

The limitations of this study are those generally related to non-interventional, uncontrolled, observational studies. There was a variable amount of missing data which was particularly notable for some parameters. The reasons for this loss may include the treating physician not performing certain tests routinely, patients being lost to follow-up, or database entry errors. In addition, sites utilized their local laboratory for testing which means that inter-laboratory variation is possible. The data presented here is reflective of the practice of the clinicians included in the study and their individual monitoring procedures. As a result, some parameters of interest, such as bone mineral density and bone biomarkers, which are known to be associated with antiretroviral therapy [24], were not specifically assessed in this study. However, the main strength of the study is the diversity of the patient population meaning the results are relevant to real-world patients and to other countries with similar management strategies and HBV demographics.

In conclusion, TDF shows significant and sustained antiviral activity against HBV and a favorable safety and tolerability profile in a diverse population of patients managed in routine clinical practice, including treatment-experienced patients, older patients, and patients with advanced liver disease or renal insufficiency. The effectiveness and safety results of TDF in clinical practice are similar to those reported in clinical trials.

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Author contributions All authors recruited patients and collected data or were involved in data interpretation. All authors were involved in drafting the manuscript and critical revision of the manuscript for important intellectual content, and all authors approved the final version of the manuscript.

Compliance with ethical standards

Conflict of interest P.M. is a speaker and investigator for BMS, Boehringer-Ingelheim, Tibotec, Janssen, Gilead, Roche, and Merck. F.Z. has served as an investigator for Gilead and a consultant/speaker for Gilead and Bristol-Myers Squibb. C.H. has served as a consultant/speaker for Roche, Bristol-Myers Squibb, Merck, Janssen, AbbVie, and Gilead Sciences. X.C. has served as a consultant/speaker for Gilead, EMS, and Janssen-Cilag. M.B. has received research grants from Bayer, Gilead Sciences, Janssen and Merck. He has served as an advisory board member and speaker for AbbVie, Bristol-Myers Squibb, Gilead Sciences, GlaxoSmithKline, Idenix, Janssen, Merck, Roche and Vertex and is a consultant/speaker for Gilead. J.P.Z. consults and is on the speakers’ bureau for Gilead, Bristol-Myers Squibb, Janssen, Siemens, and MSD and he consults for AbbVie. S.M.C. has received fees for participation in advisory boards for Novo Nordisk and Gilead, and for lectures from AstraZeneca, Bayer, Boehringer Ingelheim, Bristol-Myers Squibb, Eli Lilly, Euthérapié, GlaxoSmithKline, Gilead, Janssen, Landbeck, MSD, Merck Sharp and Dohme, Novartis, Novo Nordisk, Otsuka Pharmaceutical France, Pfizer, Sanofi, Servier, and Wyeth. J.F.C. is a board member of Gilead. P.P. is a Gilead employee and has Stock-Options and RSU. O.L. is a Gilead employee. V.T. and C.S. were Gilead employees at the time of submission of this manuscript but are no longer working for Gilead. B.R., A.P., R.T., J.D., D.O., G.P.P., G.R., and D.L., have no conflicts of interest to declare.

Ethical approval For this type of study formal consent is not required. The study was conducted in accordance with Good Epidemiological Practice guidelines, was approved by the French Medical Board, and was authorized by the National Computers and Privacy Commission.

Informed consent Patients gave written consent for access to their medical records.

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