HEPSERA™

(adefovir dipivoxil)

Tablets

Rx Only

WARNINGS

1. SEVERE ACUTE EXACERBATIONS OF HEPATITIS HAVE BEEN REPORTED IN PATIENTS WHO HAVE DISCONTINUED ANTI-HEPATITIS B THERAPY, INCLUDING THERAPY WITH HEPSERA. HEPATIC FUNCTION SHOULD BE MONITORED CLOSELY IN PATIENTS WHO DISCONTINUE ANTI-HEPATITIS B THERAPY. IF APPROPRIATE, RESUMPTION OF ANTI-HEPATITIS B THERAPY MAY BE WARRANTED (SEE WARNINGS).

2. IN PATIENTS AT RISK OF OR HAVING UNDERLYING RENAL DYSFUNCTION, CHRONIC ADMINISTRATION OF HEPSERA MAY RESULT IN NEPHROTOXICITY. THESE PATIENTS SHOULD BE MONITORED CLOSELY FOR RENAL FUNCTION AND MAY REQUIRE DOSE ADJUSTMENT (SEE WARNINGS AND DOSAGE AND ADMINISTRATION).

3. HIV RESISTANCE MAY EMERGE IN CHRONIC HEPATITIS B PATIENTS WITH UNRECOGNIZED OR UNTREATED HUMAN IMMUNODEFICIENCY VIRUS (HIV) INFECTION TREATED WITH ANTI-HEPATITIS B THERAPIES, SUCH AS THERAPY WITH HEPSERA, THAT MAY HAVE ACTIVITY AGAINST HIV (SEE WARNINGS).

4. LACTIC ACIDOSIS AND SEVERE HEPATOMEGALY WITH STEATOSIS, INCLUDING FATAL CASES, HAVE BEEN REPORTED WITH THE USE OF NUCLEOSIDE ANALOGS ALONE OR IN COMBINATION WITH OTHER ANTIRETROVIRALS (SEE WARNINGS).

DESCRIPTION

HEPSERA™ is the tradename for adefovir dipivoxil, a diester prodrug of adefovir. Adefovir is an acyclic nucleotide analog with activity against human hepatitis B virus (HBV).

The chemical name of adefovir dipivoxil is 9-[2-{bis[(pivaloyloxy)methoxy]phosphinyl}-methoxy]ethyl]adenine. It has a molecular formula of C_{20}H_{32}N_{5}O_{8}P, a molecular weight of 501.48 and the following structural formula:
Adefovir dipivoxil is a white to off-white crystalline powder with an aqueous solubility of 19 mg/mL at pH 2.0 and 0.4 mg/mL at pH 7.2. It has an octanol/aqueous phosphate buffer (pH 7) partition coefficient (log p) of 1.91.

HEPSERA tablets are for oral administration. Each tablet contains 10 mg of adefovir dipivoxil and the following inactive ingredients: croscarmellose sodium, lactose monohydrate, magnesium stearate, pregelatinized starch, and talc.

**Microbiology**

*Mechanism of Action:*

Adefovir is an acyclic nucleotide analog of adenosine monophosphate. Adefovir is phosphorylated to the active metabolite, adefovir diphosphate, by cellular kinases. Adefovir diphosphate inhibits HBV DNA polymerase (reverse transcriptase) by competing with the natural substrate deoxyadenosine triphosphate and by causing DNA chain termination after its incorporation into viral DNA. The inhibition constant (Kᵢ) for adefovir diphosphate for HBV DNA polymerase was 0.1 µM. Adefovir diphosphate is a weak inhibitor of human DNA polymerases α and γ with Kᵢ values of 1.18 µM and 0.97 µM, respectively.

*Antiviral Activity:*

The *in vitro* antiviral activity of adefovir was determined in HBV transfected human hepatoma cell lines. The concentration of adefovir that inhibited 50% of viral DNA synthesis (IC₅₀) varied from 0.2 to 2.5 µM.

*Drug Resistance:*

**Clinical Studies 437 & 438**

Genotypic and phenotypic analyses of serum HBV DNA from adefovir dipivoxil (10 mg or 30 mg) treated HBeAg-positive patients (n = 215; study 437) and HBAg-negative patients (n=56; study 438) at baseline and week 48 did not identify mutations in the HBV DNA polymerase gene that may
confer reduced susceptibility to adefovir. An unconfirmed increase of $\geq 1 \log_{10}$ copies/mL in serum HBV DNA was observed in some patients. The molecular basis and/or the clinical significance for the observed unconfirmed increases are not known.

**Cross-resistance:**

Recombinant HBV variants containing lamivudine-resistance-associated mutations (L528M, M552I, M552V, L528M + M552V) in the HBV DNA polymerase gene were susceptible to adefovir *in vitro*. Adefovir has also demonstrated anti-HBV activity (median reduction in serum HBV DNA of 4.3 $\log_{10}$ copies/mL) against clinical isolates of HBV containing lamivudine-resistance-associated mutations (study 435). HBV variants with DNA polymerase mutations T476N and R or W501Q associated with resistance to hepatitis B immunoglobulin were susceptible to adefovir *in vitro*.

**CLINICAL PHARMACOLOGY**

**Pharmacokinetics**

The pharmacokinetics of adefovir have been evaluated in healthy volunteers and patients with chronic hepatitis B. Adefovir pharmacokinetics are similar between these populations.

**Absorption:**

Adefovir dipivoxil is a diester prodrug of the active moiety adefovir. Based on a cross study comparison, the approximate oral bioavailability of adefovir from HEPSERA is 59%.

Following oral administration of a 10 mg single dose of HEPSERA to chronic hepatitis B patients (n=14), the peak adefovir plasma concentration ($C_{\text{max}}$) was $18.4 \pm 6.26$ ng/mL (mean $\pm$ SD) and occurred between 0.58 and 4.00 hours (median = 1.75 hours) post dose. The adefovir area under the plasma concentration-time curve (AUC$_{0-\infty}$) was $220 \pm 70.0$ ng•h/mL. Plasma adefovir concentrations declined in a biexponential manner with a terminal elimination half-life of $7.48 \pm 1.65$ hours.

The pharmacokinetics of adefovir in subjects with adequate renal function were not affected by once daily dosing of 10 mg HEPSERA over seven days. The impact of long-term once daily administration of 10 mg HEPSERA on adefovir pharmacokinetics has not been evaluated.

**Effects of Food on Oral Absorption:**

Adefovir exposure was unaffected when a 10 mg single dose of HEPSERA was administered with food (an approximately 1000 kcal high-fat meal). HEPSERA may be taken without regard to food.

**Distribution:**

*In vitro* binding of adefovir to human plasma or human serum proteins is $\leq 4\%$ over the adefovir concentration range of 0.1 to 25 µg/mL. The volume of distribution at steady-state following intravenous administration of 1.0 or 3.0 mg/kg/day is $392 \pm 75$ and $352 \pm 9$ mL/kg, respectively.
PROPOSED FINAL LABELING FOR HEPSERA TABLETS

As of September 20, 2002

Metabolism and Elimination:

Following oral administration, adefovir dipivoxil is rapidly converted to adefovir. Forty-five percent of the dose is recovered as adefovir in the urine over 24 hours at steady state following 10 mg oral doses of HEPSERA. Adefovir is renally excreted by a combination of glomerular filtration and active tubular secretion (See DRUG INTERACTIONS).

Special Populations:

Gender

The pharmacokinetics of adefovir were similar in male and female patients.

Race

Insufficient data are available to determine the effect of race on the pharmacokinetics of adefovir.

Pediatric and Geriatric Patients

Pharmacokinetic studies have not been conducted in children or in the elderly.

Renal Impairment

In subjects with moderately or severely impaired renal function or with end-stage renal disease (ESRD) requiring hemodialysis, C<sub>max</sub>, AUC, and half-life (T<sub>1/2</sub>) were increased compared to subjects with normal renal function. It is recommended that the dosing interval of HEPSERA be modified in these patients (See DOSAGE AND ADMINISTRATION).

The pharmacokinetics of adefovir in non-chronic hepatitis B patients with varying degrees of renal impairment are described in Table 1. In this study, subjects received a 10 mg single dose of HEPSERA.
Table 1

Pharmacokinetic Parameters (Mean ± SD) of Adefovir in Patients with Varying Degrees of Renal Function

<table>
<thead>
<tr>
<th>Renal Function Group</th>
<th>Unimpaired</th>
<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline Creatinine Clearance (mL/min)</td>
<td>&gt;80 (n=7)</td>
<td>50 - 80 (n=8)</td>
<td>30 - 49 (n=7)</td>
<td>10 - 29 (n=10)</td>
</tr>
<tr>
<td>Cmax (ng/mL)</td>
<td>17.8±3.22</td>
<td>22.4±4.04</td>
<td>28.5±8.57</td>
<td>51.6±10.3</td>
</tr>
<tr>
<td>AUC 0-∞ (ng•h/mL)</td>
<td>201±40.8</td>
<td>266±55.7</td>
<td>455±176</td>
<td>1240±629</td>
</tr>
<tr>
<td>CL/F (mL/min)</td>
<td>469±99.0</td>
<td>356±85.6</td>
<td>237±118</td>
<td>91.7±51.3</td>
</tr>
<tr>
<td>CLrenal (mL/min)</td>
<td>231±48.9</td>
<td>148±39.3</td>
<td>83.9±27.5</td>
<td>37.0±18.4</td>
</tr>
</tbody>
</table>

A four-hour period of hemodialysis removed approximately 35% of the adefovir dose. The effect of peritoneal dialysis on adefovir removal has not been evaluated.

Hepatic Impairment

The pharmacokinetics of adefovir following a 10 mg single dose of HEPSERA have been studied in non-chronic hepatitis B patients with hepatic impairment. There were no substantial alterations in adefovir pharmacokinetics in patients with moderate and severe hepatic impairment compared to unimpaired patients. No change in HEPSERA dosing is required in patients with hepatic impairment.

Drug Interactions:

Adefovir dipivoxil is rapidly converted to adefovir in vivo. At concentrations substantially higher (> 4000 fold) than those observed in vivo, adefovir did not inhibit any of the common human CYP450 enzymes, CYP1A2, CYP2C9, CYP2C19, CYP2D6, and CYP3A4. Adefovir is not a substrate for these enzymes. However, the potential for adefovir to induce CYP450 enzymes is unknown. Based on the results of these in vitro experiments and the renal elimination pathway of adefovir, the potential for CYP450 mediated interactions involving adefovir as an inhibitor or substrate with other medicinal products is low.

The pharmacokinetics of adefovir have been evaluated following multiple dose administration of HEPSERA (10 mg once daily) in combination with lamivudine (100 mg once daily), trimethoprim/sulfamethoxazole (160/800 mg twice daily), acetaminophen (1000 mg four times daily) and ibuprofen (800 mg three times daily) in healthy volunteers (n = 18 per study).

Adefovir did not alter the pharmacokinetics of lamivudine, trimethoprim/sulfamethoxazole,
acetaminophen and ibuprofen.

The pharmacokinetics of adefovir were unchanged when HEPSERA was co-administered with lamivudine, trimethoprim/sulfamethoxazole and acetaminophen. When HEPSERA was co-administered with ibuprofen (800 mg three times daily) increases in adefovir $C_{\text{max}}$ (33%), AUC (23%) and urinary recovery were observed. This increase appears to be due to higher oral bioavailability, not a reduction in renal clearance of adefovir.

**INDICATIONS AND USAGE**

HEPSERA is indicated for the treatment of chronic hepatitis B in adults with evidence of active viral replication and either evidence of persistent elevations in serum aminotransferases (ALT or AST) or histologically active disease.

This indication is based on histological, virological, biochemical, and serological responses in adult patients with HBeAg+ and HBeAg- chronic hepatitis B with compensated liver function, and in adult patients with clinical evidence of lamivudine-resistant hepatitis B virus with either compensated or decompensated liver function.

**Description of Clinical Studies**

**HBeAg-positive Chronic Hepatitis B:**

Study 437 was a randomized, double-blind, placebo-controlled, three-arm study in patients with HBeAg-positive chronic hepatitis B that allowed for a comparison between placebo and HEPSERA. The median age of patients was 33 years. Seventy-four percent were male, 59% were Asian, 36% were Caucasian, and 24% had prior interferon-α treatment. At baseline, patients had a median total Knodell Histology Activity Index (HAI) score of 10, a median serum HBV DNA level as measured by an experimental polymerase chain reaction assay of 8.36 log$_{10}$ copies/mL and a median ALT level of 2.3 times the upper limit of normal.

**HBeAg-negative (anti-HBe positive / HBV DNA positive) Chronic Hepatitis B:**

Study 438 was a randomized, double-blind, placebo-controlled study in patients who were HBeAg-negative at screening, and anti-HBeAg positive. The median age of patients was 46 years. Eighty-three percent were male, 66% were Caucasian, 30% were Asian, and 41% had prior interferon-α treatment. At baseline, the median total Knodell HAI score was 10, the median serum HBV DNA level as measured by an experimental polymerase chain reaction assay was 7.08 log$_{10}$ copies/mL, and the median ALT was 2.3 times the upper limit of normal.

The primary efficacy endpoint in both studies was histological improvement at week 48; results of
which are shown in Table 2.

### Table 2

**Histological Response at Week 48***

<table>
<thead>
<tr>
<th></th>
<th>Study 437</th>
<th>Study 438</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HEPSERA 10 mg</td>
<td>Placebo</td>
</tr>
<tr>
<td>(n=168)</td>
<td>(n=161)</td>
<td>(n=121)</td>
</tr>
<tr>
<td>Improvement**</td>
<td>53%</td>
<td>25%</td>
</tr>
<tr>
<td>No Improvement</td>
<td>37%</td>
<td>67%</td>
</tr>
<tr>
<td>Missing/ Unassessable Data</td>
<td>10%</td>
<td>7%</td>
</tr>
</tbody>
</table>

*Intent-to-Treat population (patients with ≥ 1 dose of study drug) with assessable baseline biopsies.

**Histological improvement defined as ≥ 2 point decrease in the Knodell necro-inflammatory score with no worsening of the Knodell fibrosis score.

Table 3 illustrates the changes in Ishak Fibrosis Score by treatment group.

### Table 3

**Changes in Ishak Fibrosis Score at Week 48**

<table>
<thead>
<tr>
<th></th>
<th>Study 437</th>
<th>Study 438</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HEPSERA 10 mg</td>
<td>Placebo</td>
</tr>
<tr>
<td>(n=150)</td>
<td>(n=146)</td>
<td>(n=112)</td>
</tr>
<tr>
<td>Number of adequate biopsy pairs</td>
<td>Ishak Fibrosis Score</td>
<td>Improved*</td>
</tr>
<tr>
<td>Unchanged</td>
<td>55%</td>
<td>60%</td>
</tr>
<tr>
<td>Worsened</td>
<td>11%</td>
<td>21%</td>
</tr>
</tbody>
</table>

*Change of 1 point or more in Ishak Fibrosis Score.
At week 48, improvement was seen in respect to mean change in serum HBV DNA ($\log_{10}$ copies/mL), normalization of ALT, and HBeAg seroconversion as compared to placebo in patients receiving HEPSERA (Table 4).

### Table 4

<table>
<thead>
<tr>
<th>Study 437</th>
<th>Study 438</th>
</tr>
</thead>
<tbody>
<tr>
<td>HEPSERA 10 mg</td>
<td>Placebo HEPSERA 10 mg</td>
</tr>
<tr>
<td>(n=167)</td>
<td>(n=171)</td>
</tr>
<tr>
<td>Mean change ± SD in serum HBV DNA from baseline ($\log_{10}$ copies/mL)</td>
<td>-3.57±1.64</td>
</tr>
<tr>
<td>ALT Normalization</td>
<td>48%</td>
</tr>
<tr>
<td>HBeAg Seroconversion</td>
<td>12%</td>
</tr>
</tbody>
</table>

* Patients with HBeAg-negative disease cannot undergo HBeAg seroconversion

In studies 437 and 438, continued treatment with HEPSERA to 72 weeks resulted in continued maintenance of mean reductions in serum HBV DNA observed at week 48. An increase in the proportion of patients with ALT normalization was also observed in study 437. The effect of continued treatment with HEPSERA on seroconversion is unknown.

**Pre- and Post-Liver Transplantation Patients:**

HEPSERA was also evaluated in an open-label, uncontrolled study of 324 chronic hepatitis B patients pre- (n=128) and post- (n=196) liver transplantation with clinical evidence of lamivudine-resistant hepatitis B virus (study 435). The median baseline HBV DNA as measured by an experimental polymerase chain reaction assay was 7.4 and 8.2 $\log_{10}$ copies/mL, and the median baseline ALT was 1.8 and 2.1 times the upper limit of normal in pre- and post-liver transplantation patients, respectively. Results of this study are displayed in Table 5. Treatment with HEPSERA resulted in a similar reduction in serum HBV DNA regardless of the patterns of lamivudine-resistant HBV DNA polymerase mutations at baseline. The clinical significance of these findings as they relate to histological improvement is not known.
PROPOSED FINAL LABELING FOR HEPSERA TABLETS
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Table 5

<table>
<thead>
<tr>
<th>Efficacy Parameter</th>
<th>Pre-Liver Transplantation</th>
<th>Post-Liver Transplantation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(n=128)</td>
<td>(n=196)</td>
</tr>
<tr>
<td>Mean change ± SD in HBV DNA from baseline (log_{10} copies/mL)</td>
<td>-3.8±1.4</td>
<td>-4.1±1.6</td>
</tr>
<tr>
<td>Stable or improved Child-Pugh-Turcotte score</td>
<td>92%*</td>
<td>96%</td>
</tr>
<tr>
<td>Normalization of: **</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALT</td>
<td>76%</td>
<td>49%</td>
</tr>
<tr>
<td>Albumin</td>
<td>81%</td>
<td>76%</td>
</tr>
<tr>
<td>Bilirubin</td>
<td>50%</td>
<td>75%</td>
</tr>
<tr>
<td>Prothrombin time</td>
<td>83%</td>
<td>20%</td>
</tr>
</tbody>
</table>

* 24 week data
** Denominator is patients with abnormal values at baseline.

Clinical Evidence of Lamivudine Resistance:

In study 461, an ongoing double-blind, active controlled study in 59 chronic hepatitis B patients with clinical evidence of lamivudine-resistant hepatitis B virus, patients were randomized to receive either HEPSERA monotherapy or HEPSERA in combination with lamivudine 100 mg or lamivudine 100 mg alone. At week 16, the mean ± SD decrease in serum HBV DNA as measured by an experimental polymerase chain reaction assay was 3.11 ± 0.94 log_{10} copies/mL for patients treated with HEPSERA and 2.95 ± 0.64 log_{10} copies/mL for patients treated with HEPSERA in combination with lamivudine. There was a mean decrease in serum HBV DNA of 0.00 ± 0.28 log_{10} copies/mL in patients receiving lamivudine alone. The clinical significance of these observed changes in serum HBV DNA has not yet been established.

CONTRAINDICATIONS

HEPSERA is contraindicated in patients with previously demonstrated hypersensitivity to any of the components of the product.

WARNINGS
Exacerbations of Hepatitis after Discontinuation of Treatment

Severe acute exacerbation of hepatitis has been reported in patients who have discontinued anti-hepatitis B therapy, including therapy with HEPSEERA. Patients who discontinue HEPSEERA should be monitored at repeated intervals over a period of time for hepatic function. If appropriate, resumption of anti-hepatitis B therapy may be warranted.

In clinical trials of HEPSEERA, exacerbations of hepatitis (ALT elevations 10 times the upper limit of normal or greater) occurred in up to 25% of patients after discontinuation of HEPSEERA. Most of these events occurred within 12 weeks of drug discontinuation. These exacerbations generally occurred in the absence of HBeAg seroconversion, and presented as serum ALT elevations in addition to re-emergence of viral replication. In the HBeAg-positive and HBeAg-negative studies in patients with compensated liver function, the exacerbations were not generally accompanied by hepatic decompensation. However, patients with advanced liver disease or cirrhosis may be at higher risk for hepatic decompensation. Although most events appear to have been self-limited or resolved with re-initiation of treatment, severe hepatitis exacerbations, including fatalities, have been reported. Therefore, patients should be closely monitored after stopping treatment.

Nephrotoxicity

Nephrotoxicity characterized by a delayed onset of gradual increases in serum creatinine and decreases in serum phosphorus was historically shown to be the treatment-limiting toxicity of adefovir dipivoxil therapy at substantially higher doses in HIV-infected patients (60 and 120 mg daily) and in chronic hepatitis B patients (30 mg daily). Chronic administration of HEPSEERA (10 mg once daily) may result in nephrotoxicity. The overall risk of nephrotoxicity in patients with adequate renal function is low. However, this is of special importance in patients at risk of or having underlying renal dysfunction and patients taking concomitant nephrotoxic agents such as cyclosporine, tacrolimus, aminoglycosides, vancomycin and non-steroidal anti-inflammatory drugs (See ADVERSE REACTIONS).

It is important to monitor renal function for all patients during treatment with HEPSEERA, particularly for those with pre-existing or other risks for renal impairment. Patients with renal insufficiency at baseline or during treatment may require dose adjustment (See DOSAGE AND ADMINISTRATION). The risks and benefits of HEPSEERA treatment should be carefully evaluated prior to discontinuing HEPSEERA in a patient with treatment-emergent nephrotoxicity.

HIV Resistance

Prior to initiating HEPSEERA therapy, HIV antibody testing should be offered to all patients. Treatment with anti-hepatitis B therapies, such as HEPSEERA, that have activity against HIV in a chronic hepatitis B patient with unrecognized or untreated HIV infection may result in emergence of HIV resistance. HEPSEERA has not been shown to suppress HIV RNA in patients; however, there are limited data on the use of HEPSEERA to treat patients with chronic hepatitis B co-infected with HIV.
Lactic Acidosis/Severe Hepatomegaly with Steatosis

Lactic acidosis and severe hepatomegaly with steatosis, including fatal cases, have been reported with the use of nucleoside analogs alone or in combination with antiretrovirals.

A majority of these cases have been in women. Obesity and prolonged nucleoside exposure may be risk factors. Particular caution should be exercised when administering nucleoside analogs to any patient with known risk factors for liver disease; however, cases have also been reported in patients with no known risk factors. Treatment with HEPSERA should be suspended in any patient who develops clinical or laboratory findings suggestive of lactic acidosis or pronounced hepatotoxicity (which may include hepatomegaly and steatosis even in the absence of marked transaminase elevations).

PRECAUTIONS

Drug Interactions

Since adefovir is eliminated by the kidney, co-administration of HEPSERA with drugs that reduce renal function or compete for active tubular secretion may increase serum concentrations of either adefovir and/or these co-administered drugs.

Apart from lamivudine, trimethoprim/sulfamethoxazole and acetaminophen, the effects of co-administration of HEPSERA with drugs that are excreted renally, or other drugs known to affect renal function have not been evaluated (See CLINICAL PHARMACOLOGY).

Patients should be monitored closely for adverse events when HEPSERA is co-administered with drugs that are excreted renally or with other drugs known to affect renal function.

Ibuprofen 800 mg three times daily increased adefovir exposure by approximately 23%. The clinical significance of this increase in adefovir exposure is unknown (See CLINICAL PHARMACOLOGY).

While adefovir does not inhibit common CYP450 enzymes, the potential for adefovir to induce CYP450 enzymes is not known.

The effect of adefovir on cyclosporine and tacrolimus concentrations is not known.

Duration of Treatment

The optimal duration of HEPSERA treatment and the relationship between treatment response and long-term outcomes such as hepatocellular carcinoma or decompensated cirrhosis are not known.

Animal Toxicology

Renal tubular nephropathy characterized by histological alterations and/or increases in BUN and serum creatinine was the primary dose-limiting toxicity associated with administration of adefovir dipivoxil in animals. Nephrotoxicity was observed in animals at systemic exposures approximately 3-10 times higher than those in humans at the recommended therapeutic dose of 10 mg/day.

Carcinogenesis, Mutagenesis, Impairment of Fertility
Carcinogenicity studies in mice and rats receiving adefovir have been conducted. In mice, at dose levels of 1, 3, or 10 mg/kg/day, no treatment-related increases in tumor incidence were found at 10 mg/kg/day (systemic exposure was 10 times that achieved in humans at a therapeutic dose of 10 mg/day). In rats dosed at levels of 0.5, 1.5, or 5 mg/kg/day, no drug-related increase in tumor incidence was observed. The exposure at the high dose was four times that at the human therapeutic dose. Adefovir dipivoxil was mutagenic in the \textit{in vitro} mouse lymphoma cell assay (with or without metabolic activation). Adefovir induced chromosomal aberrations in the \textit{in vitro} human peripheral blood lymphocyte assay without metabolic activation. Adefovir was not clastogenic in the \textit{in vivo} mouse micronucleus assay at doses up to 2,000 mg/kg and it was not mutagenic in the Ames bacterial reverse mutation assay using \textit{S. typhimurium} and \textit{E. coli} strains in the presence and absence of metabolic activation. In reproductive toxicology studies, no evidence of impaired fertility was seen in male or female rats at doses up to 30 mg/kg/day (systemic exposure 19 times that achieved in humans at the therapeutic dose).

**Pregnancy**

Pregnancy Category C:

Reproduction studies conducted with adefovir dipivoxil administered orally have shown no embryotoxicity or teratogenicity in rats at doses up to 35 mg/kg/day (systemic exposure approximately 23 times that achieved in humans at the therapeutic dose of 10 mg/day), or in rabbits at 20 mg/kg/day (systemic exposure 40 times human).

When adefovir was administered intravenously to pregnant rats at doses associated with notable maternal toxicity (20 mg/kg/day, systemic exposure 38 times human), embryotoxicity and an increased incidence of fetal malformations (anasarca, depressed eye bulge, umbilical hernia and kinked tail) were observed. No adverse effects on development were seen with adefovir administered intravenously to pregnant rats at 2.5 mg/kg/day (systemic exposure 12 times human).

There are no adequate and well-controlled studies in pregnant women. Because animal reproduction studies are not always predictive of human response, HEPSERA should be used during pregnancy only if clearly needed and after careful consideration of the risks and benefits.

**Pregnancy Registry**

To monitor fetal outcomes of pregnant women exposed to HEPSERA, a pregnancy registry has been established. Healthcare providers are encouraged to register patients by calling 1-800-258-4263.

**Labor and Delivery**

There are no studies in pregnant women and no data on the effect of HEPSERA on transmission of HBV from mother to infant. Therefore, appropriate infant immunizations should be used to prevent neonatal acquisition of hepatitis B virus.

**Lactating Women**

It is not known whether adefovir is excreted in human milk. Mothers should be instructed not to breast-feed if they are taking HEPSERA.

**Pediatric Use**
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Safety and effectiveness in pediatric patients have not been established.

Geriatric Use

Clinical studies of HEPSERA did not include sufficient numbers of patients aged 65 and over to determine whether they respond differently from younger patients. In general, caution should be exercised when prescribing to elderly patients since they have greater frequency of decreased renal or cardiac function due to concomitant disease or other drug therapy.

ADVERSE REACTIONS

Assessment of adverse reactions is based on two studies (437 and 438) in which 522 patients with chronic hepatitis B received double-blind treatment with HEPSERA (n=294) or placebo (n=228) for 48 weeks. With extended therapy in the second 48 week treatment period, 492 patients were treated for up to 109 weeks, with a median time on treatment of 49 weeks.

In addition to specific adverse events described under the WARNINGS section, all treatment-related clinical adverse events that occurred in 3% or greater of HEPSERA-treated patients compared with placebo are listed in Table 6. A summary of grade 3 and 4 laboratory abnormalities during therapy with HEPSERA compared with placebo is listed in Table 7.

Table 6

<table>
<thead>
<tr>
<th></th>
<th>HEPSERA 10 mg (n=294)</th>
<th>Placebo (n=228)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthenia</td>
<td>13%</td>
<td>14%</td>
</tr>
<tr>
<td>Headache</td>
<td>9%</td>
<td>10%</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>9%</td>
<td>11%</td>
</tr>
<tr>
<td>Nausea</td>
<td>5%</td>
<td>8%</td>
</tr>
<tr>
<td>Flatulence</td>
<td>4%</td>
<td>4%</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>3%</td>
<td>4%</td>
</tr>
<tr>
<td>Dyspepsia</td>
<td>3%</td>
<td>2%</td>
</tr>
</tbody>
</table>
Laboratory Abnormalities

Table 7

<table>
<thead>
<tr>
<th>Grade 3-4 Laboratory Abnormalities</th>
<th>HEPSERA 10 mg</th>
<th>Placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reported in ≥ 1% of All HEPSERA-Treated Patients in the Pooled 437-438 Studies (0-48 weeks)</td>
<td>(n=294)</td>
<td>(n=228)</td>
</tr>
<tr>
<td>ALT (≥ 5 x ULN)</td>
<td>20%</td>
<td>41%</td>
</tr>
<tr>
<td>Hematuria (≥ 3+)</td>
<td>11%</td>
<td>10%</td>
</tr>
<tr>
<td>AST (≥ 5 x ULN)</td>
<td>8%</td>
<td>23%</td>
</tr>
<tr>
<td>Creatine Kinase (≥ 4 x ULN)</td>
<td>7%</td>
<td>7%</td>
</tr>
<tr>
<td>Amylase (≥ 2 x ULN)</td>
<td>4%</td>
<td>4%</td>
</tr>
<tr>
<td>Glycosuria (≥ 3+)</td>
<td>1%</td>
<td>3%</td>
</tr>
</tbody>
</table>

In patients with adequate renal function, increases in serum creatinine ≥ 0.3 mg/dL from baseline were observed in 4% of patients treated with HEPSERA 10 mg daily compared with 2% of patients in the placebo group at week 48. No patients developed a serum creatinine increase ≥ 0.5 mg/dL from baseline by week 48. By week 96, 10% and 2% of HEPSERA-treated patients, by Kaplan-Meier estimate, had increases in serum creatinine ≥ 0.3 mg/dL and ≥ 0.5 mg/dL from baseline, respectively (no placebo-controlled results were available for comparison beyond week 48). Of the 29 of 492 patients with elevations in serum creatinine ≥ 0.3 mg/dL from baseline, 20 out of 29 resolved on continued treatment (≤ 0.2 mg/dL from baseline), 8 of 29 remained unchanged and 1 of 29 resolved on discontinuing treatment (see Special Risk Patients section below for changes in serum creatinine in patients with underlying renal insufficiency at baseline).

Special Risk Patients

Pre- (n=128) and post-liver transplantation patients (n=196) with chronic hepatitis B and clinical evidence of lamivudine-resistant hepatitis B virus were treated in an open-label study with HEPSERA for up to 129 weeks, with a median time on treatment of 19 and 56 weeks, respectively. The majority of these patients had some degree of underlying renal insufficiency at baseline or other risk factors for renal dysfunction during treatment. Increases in serum creatinine ≥ 0.3 mg/dL from baseline were observed in 26% of these patients by week 48 and 37% by week 96 by
Kaplan-Meier estimates. Increases in serum creatinine $\geq 0.5$ mg/dL from baseline were observed in 16% of these patients by week 48 and 31% by week 96. Of the 41 of 324 patients with elevations in serum creatinine $\geq 0.5$ mg/dL from baseline, 7 of 41 resolved on continued treatment ($\leq 0.3$ mg/dL from baseline), 18 of 41 remained unchanged and 16 of 41 had not resolved. Additionally, decreases in serum phosphorus were observed in 4% of these patients by week 48, and 6% by week 96 by Kaplan-Meier estimates. One percent (3 of 324) of pre- and post-liver transplantation patients discontinued HEPSERA due to renal events.

Due to the presence of multiple concomitant risk factors for renal dysfunction in these patients, the contributory role of HEPSERA to these changes in serum creatinine and serum phosphorus is difficult to assess.

The most common treatment-related adverse events reported in pre- and post-liver transplantation patients treated with HEPSERA with a 2% frequency or higher include:

- **Body as a whole:** asthenia, abdominal pain, headache, fever
- **Gastrointestinal:** nausea, vomiting, diarrhea, flatulence, hepatic failure
- **Metabolic and Nutritional:** increases in ALT and AST, abnormal liver function
- **Respiratory:** increased cough, pharyngitis, sinusitis
- **Skin and Appendages:** pruritus, rash
- **Urogenital:** increases in creatinine, renal failure, renal insufficiency

**OVERDOSAGE**

Doses of adefovir dipivoxil 500 mg daily for 2 weeks and 250 mg daily for 12 weeks have been associated with gastrointestinal side effects. If overdose occurs the patient must be monitored for evidence of toxicity, and standard supportive treatment applied as necessary.

Following a 10 mg single dose of HEPSERA, a four-hour hemodialysis session removed approximately 35% of the adefovir dose.

**DOSAGE AND ADMINISTRATION**

The recommended dose of HEPSERA in chronic hepatitis B patients with adequate renal function is 10 mg, once daily, taken orally, without regard to food. The optimal duration of treatment is unknown.

**Dose Adjustment in Renal Impairment:**

Significantly increased drug exposures were seen when HEPSERA was administered to patients with renal impairment (See Pharmacokinetics). Therefore, the dosing interval of HEPSERA should be adjusted in patients with baseline creatinine clearance $< 50$ mL/min using the following suggested guidelines (See Table 8). The safety and effectiveness of these dosing interval adjustment guidelines have not been clinically evaluated. Additionally, it is important to note that
these guidelines were derived from data in patients with pre-existing renal impairment at baseline. They may not be appropriate for patients in whom renal insufficiency evolves during treatment with HEPSERA. Therefore, clinical response to treatment and renal function should be closely monitored in these patients.

<table>
<thead>
<tr>
<th>Creatinine Clearance (mL/min)*</th>
<th>≥50</th>
<th>20-49</th>
<th>10-19</th>
<th>Hemodialysis Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recommended Dose and Dosing Interval</td>
<td>10 mg every 24 hours</td>
<td>10 mg every 48 hours</td>
<td>10 mg every 72 hours</td>
<td>10 mg every 7 days following dialysis</td>
</tr>
</tbody>
</table>

*Creatinine clearance calculated by Cockcroft-Gault method using lean or ideal body weight.

The pharmacokinetics of adefovir have not been evaluated in non-hemodialysis patients with creatinine clearance < 10 mL/min; therefore, no dosing recommendation is available for these patients.

**HOW SUPPLIED**

HEPSERA is available as tablets. Each tablet contains 10 mg of adefovir dipivoxil. The tablets are white and debossed with “10” and “GILEAD” on one side and the stylized figure of a liver on the other side. They are packaged as follows: Bottles of 30 tablets (NDC 61958-0501-1) containing desiccant (silica gel) and closed with a child-resistant closure.

Store in original container at 25 °C (77 °F), excursions permitted to 15-30 °C (59-86 °F) (See USP Controlled Room Temperature).
PROPOSED FINAL LABELING FOR HEPSERA TABLETS

As of September 20, 2002

PATIENT INFORMATION

HEPSERA™ (hep-SER-rah)

Generic Name: (adefovir dipivoxil) tablets

Read this information carefully before you start taking HEPSERA. Read and check for new information each time you get more HEPSERA. This information does not take the place of talking with your doctor about your medical condition or your treatment.

What is the most important information I should know about HEPSERA?

1. Some people who take HEPSERA get a worse or very serious hepatitis when they stop taking it. This usually happens within 12 weeks after stopping. You will need to have regular blood tests to check for liver function and hepatitis B virus levels if you stop taking HEPSERA.

2. HEPSERA may cause a severe kidney problem called nephrotoxicity. It usually happens in people that already have a kidney problem, but it can happen to anyone that uses HEPSERA. You will need to have regular blood tests to check for kidney function while you are taking HEPSERA.

3. If you get or have HIV that isn’t being treated with medicines, HEPSERA may increase the chances your HIV infection cannot be helped with usual HIV medicines. This can happen if you get or have HIV and don’t know it, or if your HIV is not being treated while you are taking HEPSERA. You should get an HIV test before you start taking HEPSERA and anytime after that when there’s a chance you were exposed to HIV.

4. Some people who have taken nucleoside analog medicines, like HEPSERA, have developed a serious condition called lactic acidosis (build up of an acid in the blood). Lactic acidosis is a medical emergency and must be treated in the hospital. Call your doctor right away if you get any of the following signs of lactic acidosis:
   - You feel very weak or tired.
   - You have unusual (not normal) muscle pain.
   - You have trouble breathing.
   - You have stomach pain with nausea and vomiting.
   - You feel cold, especially in your arms and legs.
   - You feel dizzy or lightheaded.
   - You have a fast or irregular heartbeat.

Some people who have taken medicines like HEPSERA have developed serious liver problems
called hepatotoxicity, with liver enlargement (hepatomegaly) and fat in the liver (steatosis). Call your doctor right away if you get any of the following signs of liver problems.

- Your skin or the white part of your eyes turns yellow (jaundice).
- Your urine turns dark.
- Your bowel movements (stools) turn light in color.
- You don’t feel like eating food for several days or longer.
- You feel sick to your stomach (nausea).
- You have lower stomach pain.

You may be more likely to get lactic acidosis or serious liver problems if you are very overweight (obese) or have been taking nucleoside analog medicines [Combivir (zidovudine plus lamivudine), Epivir-HIV, Epivir-HBV (lamivudine), Hivid (zalcitabine), Retrovir (zidovudine), Trizivir (zidovudine plus lamivudine plus abacavir), Videx (didanosine), Viread (tenofovir disoproxil fumarate), Zerit (stavudine), Ziagen (abacavir)], like HEPSERA, for a long time.

What is HEPSERA?

HEPSERA is a medicine used to treat adults with continuing (chronic) infections with active hepatitis B virus. HEPSERA has not been studied in adults over the age of 65 or in children.

- HEPSERA will not cure your chronic hepatitis B.
- HEPSERA may help lower the amount of hepatitis B virus in your body.
- HEPSERA may lower the ability of the virus to multiply and infect new liver cells.
- We do not know if HEPSERA will reduce your chances of getting liver cancer or liver damage (cirrhosis) from chronic hepatitis B.
- We do not know how long HEPSERA may help your hepatitis. Sometimes viruses change in your body and medicines no longer work. This is called drug resistance.
- HEPSERA does not stop you from spreading hepatitis B to others by sex or sharing needles. So practice safe sex and needle use.

Who should not take HEPSERA?

- Do not take HEPSERA if you are allergic to any of the ingredients in HEPSERA. The active ingredient in HEPSERA is adefovir dipivoxil. See the end of this leaflet for a complete list of all the ingredients in HEPSERA.

Tell your doctor if:
• You are pregnant. We do not know if HEPSERA can harm your unborn child. You and your
doctor will need to decide if HEPSERA is right for you. If you take HEPSERA and you are
pregnant, talk to your doctor about how you can be on the HEPSERA pregnancy registry.
• You are breast-feeding. We do not know if HEPSERA can pass through your milk and if it can
harm your baby. You will need to choose either to breast feed or take HEPSERA, but not both.
• You have kidney problems now or had them before. Your dose and schedule of HEPSERA
may be reduced. Blood tests will need to be done regularly to see how your kidneys are working.

Tell your doctor about all the medicines you take, including prescription and non-prescription
medicines, vitamins, and herbal supplements. Some medicines may affect how HEPSERA works,
especially medicines that affect how your kidneys work. HEPSERA can affect how your other
medicines work. Your dose of HEPSERA and the other medicines may be changed. Do not take
any other medicines while you are taking HEPSERA, unless your doctor has told you it is
okay.

How should I take HEPSERA?

• Your doctor will tell you how much HEPSERA to take.
• Your doctor will tell you when and how often to take HEPSERA.
• Take HEPSERA the same time each day that your doctor tells you. If you forget to take
HEPSERA, take it as soon as you remember that day. Do not take more than 1 dose of
HEPSERA in a day. Do not take 2 doses at the same time. Call your doctor or pharmacist if you
are not sure what to do.
• Do not change your dose of HEPSERA or stop HEPSERA without talking to your doctor. Your
hepatitis may get worse if you change doses or stop.
• You may take HEPSERA with or without food.
• When your HEPSERA supply gets low, call your doctor or pharmacy for a refill. Do not run out
of HEPSERA.
• If you take too much HEPSERA, call your local poison control center or emergency room right
away.

Some patients get worse or very serious hepatitis B symptoms when they stop taking HEPSERA
(see, “What is the most important information I should know about HEPSERA?”). We don’t know
how long you should use HEPSERA. You and your doctor will need to decide when it is best for
you to stop taking HEPSERA. After you stop taking HEPSERA, your doctor will still need to check
your health and take blood tests to check your liver for a few months.

What should I avoid while taking HEPSERA?

Avoid doing things that can spread hepatitis B since HEPSERA doesn’t stop you from passing the
infection to others.

- Do not share needles or other injection equipment.
- Do not share personal items that can have blood or body fluids on them, like toothbrushes or razor blades.
- Do not have any kind of sex without protection. Practice “safe sex” using condoms and dental dams.

What are the possible side effects of HEPSERA?

HEPSERA can cause the following serious side effects: (See, “What is the most important information I should know about HEPSERA?”)

1. a worse or very serious hepatitis if you stop taking it
2. a severe kidney problem called nephrotoxicity
3. increase your chance of developing a form of HIV that cannot be treated with usual HIV medicines
4. lactic acidosis and liver problems

The most common side effects of HEPSERA are weakness, headache, stomach pain and nausea. The most common side effects in patients with liver transplants and chronic hepatitis B are weakness, headache, stomach pain, and itching. Some patients with liver transplants also had changes in the way their kidneys worked.

These are not all of the possible side effects of HEPSERA. For more information, ask your doctor or pharmacist.

General information about the safe and effective use of HEPSERA:

Medicines are sometimes prescribed for conditions not mentioned in patient information leaflets. Do not use HEPSERA for a condition for which it was not prescribed. Do not give HEPSERA to other people, even if they have the same symptoms that you have.

This leaflet summarizes the most important information about HEPSERA. If you would like more information, talk with your doctor. You can ask your doctor or pharmacist for information about HEPSERA that is written for health professionals.

HEPSERA tablets should be stored at room temperature and should be stored in their original container.

What are the Ingredients of HEPSEERA?
PROPOSED FINAL LABELING FOR HEPSERA TABLETS

As of September 20, 2002

Active Ingredient: Adefovir dipivoxil

Inactive Ingredients: croscarmellose sodium, lactose monohydrate, magnesium stearate, pregelatinized starch and talc
This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/
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Mark Goldberger
9/20/02 05:41:06 PM