TYZEKA® (telbivudine) tablets, for oral use
TYZEKA® (telbivudine) solution, for oral use

Initial U.S. Approval: 2006

TYZEKA® (telbivudine) solution, for oral use

TYZEKA Dose

<table>
<thead>
<tr>
<th>Creatinine Clearance (mL/min)</th>
<th>Oral Solution (100 mg/5 mL)</th>
<th>Tablet (600 mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>greater than or equal to 50</td>
<td>30 mL once daily 1 tab every 24 hrs</td>
<td>1 tab every 24 hrs</td>
</tr>
<tr>
<td>30 – 49</td>
<td>20 mL once daily 1 tab every 48 hrs</td>
<td>1 tab every 72 hrs</td>
</tr>
<tr>
<td>less than 30 (not requiring dialysis)</td>
<td>10 mL once daily 1 tab every 72 hrs</td>
<td></td>
</tr>
<tr>
<td>End stage renal disease (ESRD)</td>
<td>6 mL once daily 1 tab every 96 hrs</td>
<td></td>
</tr>
</tbody>
</table>

When administered on hemodialysis days, administer TYZEKA after hemodialysis (2.2).

WARNING: SEVERE ACUTE EXACERBATIONS OF HEPATITIS B

Severe acute exacerbations of hepatitis B have been reported in patients who discontinued anti-hepatitis B therapy, including TYZEKA. Hepatic function should be monitored closely in patients who discontinue therapy. Resumption of anti-hepatitis B therapy may be warranted (5.1).

INDICATIONS AND USAGE

TYZEKA® (telbivudine) solution, for oral use

WARNINGS AND PRECAUTIONS

Lactic Acidosis: If suspected, treatment should be suspended (5.2).

Myopathy and Rhabdomyolysis: TYZEKA should be interrupted if myopathy is suspected; and discontinued if confirmed. It is unknown whether risk of myopathy is increased with concomitant use of other medications associated with myopathy (5.3).

Peripheral Neuropathy: Risk increased when TYZEKA used in combination with alfa interferons, avoid concomitant use. TYZEKA should be interrupted if peripheral neuropathy is suspected; and discontinued if confirmed (4, 5, 4, 7).

ADVERSE REACTIONS

In clinical trials, the most common adverse reactions (greater than or equal to 3%), of any severity, were: fatigue, increased creatine kinase (CK), headache, cough, diarrhea, abdominal pain, nausea, pharyngolaryngeal pain, arthralgia, pyrexia, rash, back pain, dizziness, myalgia, ALT increased, dyspepsia, insomnia, and abdominal distension (6.1). The most common adverse events resulting in TYZEKA discontinuation included increased CK, nausea, diarrhea, fatigue, myalgia, and myopathy (6.1).

OVERDOSAGE

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide.

Revised: 12/2018

CONTRAINDICATIONS

Combination of TYZEKA with pegylated interferon alfa-2a: Increased risk of peripheral neuropathy (4).

DRUG INTERACTIONS

Coadministration of TYZEKA with drugs that affect renal function may alter plasma concentrations of TYZEKA and/or coadministered drug (7).

Coadministration of TYZEKA with pegylated interferon alfa-2a increases the risk and severity of peripheral neuropathy (7).

To report SUSPECTED ADVERSE REACTIONS, contact Novartis Pharmaceuticals Corporation at 1-888-669-6682 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

CONTRAINdications

Tablet: 600 mg tablets.
Oral Solution: 100 mg per 5 mL solution.

INDICATIONS AND USAGE

See full prescribing information for complete boxed warning.

WARNINGS AND PRECAUTIONS

Boxed Warning

3/2018

WARNINGS AND PRECAUTIONS

Lactic Acidosis (5.2)      3/2018

Myopathy and Rhabdomyolysis (5.3)     3/2018

PATIENT COUNSELING INFORMATION

*Sections or subsections omitted from the full prescribing information are not listed
WARNING: SEVERE ACUTE EXACERBATIONS OF HEPATITIS B

Severe acute exacerbations of hepatitis B have been reported in patients who have discontinued anti-hepatitis B therapy, including TYZEKA. Hepatic function should be monitored closely with both clinical and laboratory follow-up for at least several months in patients who discontinue anti-hepatitis B therapy. If appropriate, resumption of anti-hepatitis B therapy may be warranted [see Warnings and Precautions (5.1)].

1 INDICATIONS AND USAGE

1.1 Chronic Hepatitis B

TYZEKA is indicated for the treatment of chronic hepatitis B in adult patients with evidence of viral replication and either evidence of persistent elevations in serum aminotransferases [alanine aminotransferase (ALT) or alanine aminotransferase (AST)] or histologically active disease.

The following points should be considered when initiating therapy with TYZEKA:

- This indication is based on virologic, serologic, biochemical and histologic responses in nucleoside treatment naïve adult patients with HBeAg positive and HBeAg negative chronic hepatitis B with compensated liver disease [see Clinical Studies (14)].
- For HBeAg-positive patients, TYZEKA should only be initiated in patients with hepatitis B virus (HBV) DNA less than 9 log10 copies per mL and ALT greater than or equal to 2x Upper Limit of Normal (ULN) prior to treatment.
- For HBeAg-negative patients, TYZEKA should only be initiated in patients with HBV DNA less than 7 log10 copies per mL prior to treatment.
- On-treatment response should guide continued therapy [see Dosage and Administration (2.1) and Microbiology (12.4)].
- TYZEKA has not been evaluated in patients co-infected with human immunodeficiency virus (HIV), hepatitis C virus (HCV), or hepatitis D virus (HDV).
- TYZEKA has not been evaluated in liver transplant recipients or in patients with decompensated liver disease.
- TYZEKA has not been studied in well-controlled trials for the treatment of patients with established nucleoside analog reverse transcriptase inhibitor-resistant HBV infection, but is expected to be cross-resistant to lamivudine [see Microbiology (12.4)].
- The safety and efficacy of TYZEKA have not been evaluated in Black/African American or Hispanic patients [see Use in Specific Populations (8.9)].

2 DOSAGE AND ADMINISTRATION

2.1 Adults and Adolescents (16 years of age and older)

Due to higher rates of resistance that may develop with longer term treatment among patients with incomplete viral suppression, treatment should only be initiated, if pre-treatment HBV DNA and ALT measurements are known, in the following patient populations:

For HBeAg-positive patients, HBV DNA should be less than 9 log10 copies per mL and ALT should be greater than or equal to 2x ULN prior to treatment with TYZEKA.

For HBeAg-negative patients, HBV DNA should be less than 7 log10 copies per mL prior to treatment with TYZEKA.
HBV DNA levels should be monitored at 24 weeks of treatment to ensure complete viral suppression (HBV DNA less than 300 copies per mL). Alternative therapy should be initiated for patients who have detectable HBV DNA after 24 weeks of treatment. Optimal therapy should be guided by further resistance testing.

The recommended dose of TYZEKA for the treatment of chronic hepatitis B is 600 mg once daily, taken orally, with or without food.

TYZEKA oral solution (30 mL) may be considered for patients who have difficulty with swallowing tablets.

### 2.2 Renal Impairment

TYZEKA may be used for the treatment of chronic hepatitis B in patients with impaired renal function. No adjustment to the recommended dose of TYZEKA is necessary in patients whose creatinine clearance (CrCl) is greater than or equal to 50 mL per min. Adjustment of the total daily dose of TYZEKA oral solution or of the interval for administration of TYZEKA tablets is required in patients with CrCl less than 50 mL per min including those with end stage renal disease (ESRD) on hemodialysis (Table 1).

<table>
<thead>
<tr>
<th>Creatinine Clearance (mL/min)</th>
<th>TYZEKA Oral Solution Dose (5 mL = 100 mg)</th>
<th>TYZEKA Tablet Dose (1 tablet = 600 mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>greater than or equal to 50</td>
<td>30 mL once daily</td>
<td>1 tablet every 24 hrs</td>
</tr>
<tr>
<td>30-49</td>
<td>20 mL once daily</td>
<td>1 tablet every 48 hrs</td>
</tr>
<tr>
<td>less than 30 (not requiring dialysis)</td>
<td>10 mL once daily</td>
<td>1 tablet every 72 hrs</td>
</tr>
<tr>
<td>ESRD</td>
<td>6 mL once daily</td>
<td>1 tablet every 96 hrs(^1)</td>
</tr>
</tbody>
</table>

\(^1\) When administered on hemodialysis days, TYZEKA should be administered after hemodialysis.

### 2.3 Hepatic Impairment

No adjustment to the recommended dose of TYZEKA is necessary in patients with hepatic impairment.

### 2.4 Duration of Therapy

For patients with incomplete viral suppression (HBV DNA greater than or equal to 300 copies per mL) after 24 weeks of treatment, alternate therapy should be instituted. HBV DNA should be monitored every 6 months to ensure continued response. If patients test positive for HBV DNA at any time after their initial response, alternative treatment should be instituted. Optimal therapy should be guided by resistance testing.

The optimal duration of therapy with TYZEKA for patients with chronic hepatitis B is unknown.

### 3 DOSAGE FORMS AND STRENGTHS

**Tablets**

TYZEKA (telbivudine) 600 mg tablets are white to slightly yellowish film-coated, ovaloid-shaped tablets, imprinted with “LDT” on one side.

**Oral Solution**

TYZEKA (telbivudine) oral solution is a clear, colorless to pale yellow, passion fruit flavored liquid. TYZEKA oral solution contains 100 mg of telbivudine per 5 milliliters. Therefore, 30 mL of the oral solution provides a 600 mg dose of telbivudine.

### 4 CONTRAINDICATIONS

TYZEKA is contraindicated in combination with pegylated interferon alfa-2a because of increased risk of peripheral neuropathy [see Warnings and Precautions (5.4) and Drug Interactions (7)].
5 WARNINGS AND PRECAUTIONS

5.1 Exacerbations of Hepatitis B after Discontinuation of Treatment

Severe acute exacerbations of hepatitis B have been reported in patients who have discontinued anti-hepatitis B therapy, including TYZEKA. Hepatic function should be monitored closely with both clinical and laboratory follow-up for at least several months in patients who discontinue anti-hepatitis B therapy. If appropriate, resumption of anti-hepatitis B therapy may be warranted [see Adverse Reactions (6.1)].

5.2 Lactic Acidosis

Lactic acidosis and severe hepatomegaly with steatosis, including fatal cases, have been reported with the use of nucleoside analogues. Postmarketing cases of lactic acidosis have been reported with TYZEKA. Cases were often associated with other serious conditions (e.g. rhabdomyolysis) and/or associated with muscle-related events (e.g., myopathy, myositis) [see Warnings and Precautions (5.3)]. Some cases were also associated with pancreatitis, liver failure/hepatic steatosis and renal failure. Treatment with TYZEKA 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).

5.3 Myopathy and Rhabdomyolysis

Cases of myopathy/myositis have been reported with TYZEKA use, several weeks to months after starting therapy. Myopathy has also been reported with some other drugs in this class. Rhabdomyolysis, including fatal cases, has been reported during postmarketing use of TYZEKA. Some of the muscle related events (e.g., myopathy, myositis, and rhabdomyolysis) reported with TYZEKA were associated with lactic acidosis [see Warnings and Precautions (5.2) and Adverse Reactions (6.2)].

Uncomplicated myalgia has been reported in TYZEKA-treated patients [see Adverse Reactions (6.1)]. Myopathy, defined as persistent unexplained muscle aches and/or muscle weakness in conjunction with increases in creatine kinase (CK) values, should be considered in any patient with diffuse myalgias, muscle tenderness, or muscle weakness. Among patients with TYZEKA-associated myopathy, no pattern with regard to the degree or timing of CK elevations has been observed. In addition, the predisposing factors for the development of myopathy among TYZEKA recipients are unknown. Patients should be advised to report promptly unexplained muscle aches, pain, tenderness, or weakness. TYZEKA therapy should be interrupted if myopathy is suspected, and discontinued if myopathy is confirmed. It is unknown whether the risk of myopathy during treatment with drugs in this class is increased with coadministration of other drugs associated with myopathy, including but not limited to: corticosteroids, chloroquine, hydroxychloroquine, certain HMGCoA reductase inhibitors, fibric acid derivatives, penicillamine, zidovudine, cyclosporine, erythromycin, niacin, and certain azole antifungals. Physicians initiating concomitant treatment with any drug associated with myopathy should monitor patients closely for any signs or symptoms of unexplained muscle pain, tenderness, or weakness.

5.4 Peripheral Neuropathy

Peripheral neuropathy has been reported with TYZEKA alone or in combination with pegylated interferon alfa-2a and other interferons. In one clinical trial, an increased risk and severity of peripheral neuropathy was observed with the combined use of TYZEKA 600mg daily and pegylated interferon alfa-2a 180 micrograms once weekly compared to TYZEKA or pegylated interferon alfa-2a alone [see Contraindications (4) and Drug Interactions (7)]. Such risk cannot be excluded for other dose regimens of pegylated interferon alfa-2a, or other alfa interferons (pegylated or standard). The safety and efficacy of TYZEKA in combination with pegylated interferons or other interferons for the treatment of chronic hepatitis B have not been demonstrated. Patients should be advised to report any numbness, tingling, and/or burning sensations in the arms and/or legs, with or without gait disturbance. TYZEKA therapy should be interrupted if peripheral neuropathy is suspected, and discontinued if peripheral neuropathy is confirmed [see Adverse Reactions (6.1)].
6 ADVERSE REACTIONS

The following adverse reactions are discussed in other sections of the labeling:

- Severe acute exacerbations of hepatitis after discontinuation of treatment [see Boxed Warning and Warnings and Precautions (5.1)]
- Lactic Acidosis [see Warnings and Precautions (5.2)]
- Myopathy and Rhabdomyolysis [see Warnings and Precautions (5.3)]
- Peripheral Neuropathy [see Warnings and Precautions (5.4)]

6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to the rates in the clinical trials of another drug and may not reflect the rates observed in clinical practice.

Assessment of adverse reactions is primarily based on two trials (007 GLOBE and NV-02B-015) in which 1,699 subjects with chronic hepatitis B received double-blind treatment with TYZEKA 600 mg per day (n = 847 subjects) or lamivudine (n = 852 subjects) for 104 weeks. The median duration of therapy was 104 weeks for both treatment groups.

In the 104 week clinical trials, most adverse experiences reported with TYZEKA were classified as mild or moderate in severity and were not attributed to TYZEKA. Selected adverse events of any severity which were reported in greater than or equal to 3% of TYZEKA and lamivudine recipients are shown in Table 2. With the exception of increased CK, which was reported more frequently among TYZEKA recipients, the adverse event profile was similar for the two drugs.

<table>
<thead>
<tr>
<th>Adverse Event (Preferred Term)</th>
<th>TYZEKA N = 847</th>
<th>Lamivudine N = 852</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatigue</td>
<td>106 (13)</td>
<td>95 (11)</td>
</tr>
<tr>
<td>CK increased</td>
<td>90 (11)</td>
<td>52 (6)</td>
</tr>
<tr>
<td>Headache</td>
<td>83 (10)</td>
<td>95 (11)</td>
</tr>
<tr>
<td>Cough</td>
<td>52 (6)</td>
<td>45 (5)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>50 (6)</td>
<td>46 (5)</td>
</tr>
<tr>
<td>Abdominal pain, upper</td>
<td>49 (6)</td>
<td>52 (6)</td>
</tr>
<tr>
<td>Nausea</td>
<td>45 (5)</td>
<td>40 (5)</td>
</tr>
<tr>
<td>Pharyngolaryngeal pain</td>
<td>38 (5)</td>
<td>31 (4)</td>
</tr>
<tr>
<td>Arthralgia</td>
<td>37 (4)</td>
<td>38 (5)</td>
</tr>
<tr>
<td>Pyrexia</td>
<td>34 (4)</td>
<td>27 (3)</td>
</tr>
<tr>
<td>Rash</td>
<td>33 (4)</td>
<td>21 (3)</td>
</tr>
<tr>
<td>Back pain</td>
<td>33 (4)</td>
<td>32 (4)</td>
</tr>
<tr>
<td>Dizziness</td>
<td>32 (4)</td>
<td>43 (5)</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>29 (3)</td>
<td>31 (4)</td>
</tr>
<tr>
<td>Myalgia</td>
<td>27 (3)</td>
<td>17 (2)</td>
</tr>
<tr>
<td>Condition</td>
<td>TYZEKA 600 mg (n = 847)</td>
<td>Lamivudine 100 mg (n = 852)</td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>-------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>ALT increased</td>
<td>27 (3)</td>
<td>31 (4)</td>
</tr>
<tr>
<td>Dyspepsia</td>
<td>24 (3)</td>
<td>39 (5)</td>
</tr>
<tr>
<td>Insomnia</td>
<td>24 (3)</td>
<td>22 (3)</td>
</tr>
<tr>
<td>Abdominal distension</td>
<td>22 (3)</td>
<td>19 (2)</td>
</tr>
<tr>
<td>Pruritus</td>
<td>18 (2)</td>
<td>23 (3)</td>
</tr>
<tr>
<td>Hepatitis B exacerbation</td>
<td>17 (2)</td>
<td>36 (4)</td>
</tr>
</tbody>
</table>

Adverse events reported in greater than or equal to 3% subjects in either treatment group.

Moderate to severe (Grade 2-4) adverse events were reported in 239/847 (28%) of TYZEKA recipients and 229/852 (27%) of lamivudine recipients. The profile of adverse events of moderate to severe intensity was similar in both treatment groups and no individual adverse event was reported in greater than 2% of subjects in either treatment group.

Discontinuations due to adverse events were reported in 4% of TYZEKA recipients and 4% of lamivudine recipients. The most common adverse events resulting in TYZEKA discontinuation included increased CK, nausea, diarrhea, fatigue, myalgia, and myopathy.

Peripheral neuropathy was reported as an adverse event in less than 1% (2/847) of subjects receiving TYZEKA monotherapy [see Warnings and Precautions (5.4)]. Of TYZEKA-treated subjects less than 1% (5/847) were diagnosed with myopathy/myositis (presenting with muscular weakness) [see Warnings and Precautions (5.3)].

**Laboratory Abnormalities**

Frequencies of selected treatment-emergent laboratory abnormalities in the 007 GLOBE and NV-02B-015 trials are listed in Table 3.

**Table 3**  
Selected Treatment-Emergent Grade 3-4 Laboratory Abnormalities\(^a\) in Patients with Chronic Hepatitis B in the 104-Week Pooled 007 GLOBE and NV-02B-015 Trials

<table>
<thead>
<tr>
<th>Test</th>
<th>TYZEKA 600 mg (n = 847)</th>
<th>Lamivudine 100 mg (n = 852)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CK greater than 7.0 x ULN</td>
<td>13%</td>
<td>4%</td>
</tr>
<tr>
<td>ALT greater than 10.0 x ULN and 2.0 x baseline(^b)</td>
<td>5%</td>
<td>8%</td>
</tr>
<tr>
<td>ALT greater than 3 x baseline</td>
<td>7%</td>
<td>13%</td>
</tr>
<tr>
<td>AST (SGOT) greater than 3.0 x baseline</td>
<td>6%</td>
<td>10%</td>
</tr>
<tr>
<td>Lipase greater than 2.5 x ULN</td>
<td>2%</td>
<td>4%</td>
</tr>
<tr>
<td>Amylase greater than 3.0 x ULN</td>
<td>less than 1%</td>
<td>less than 1%</td>
</tr>
<tr>
<td>Total Bilirubin greater than 5.0 x ULN</td>
<td>less than 1%</td>
<td>less than 1%</td>
</tr>
<tr>
<td>Neutropenia (ANC less than or equal to 749/mm(^3))</td>
<td>2%</td>
<td>2%</td>
</tr>
<tr>
<td>Thrombocytopenia (Platelets less than or equal to 49,999/mm(^3))</td>
<td>less than 1%</td>
<td>less than 1%</td>
</tr>
</tbody>
</table>

Abbreviations: CK, creatine kinase; ANC, absolute neutrophil count.

On-treatment value worsened from baseline to Grade 3 or Grade 4 during therapy

American Association for the Study of Liver Diseases (AASLD) definition of acute hepatitis flare

**CK Elevations**

CK elevations were more frequent among subjects on TYZEKA treatment. By 104 weeks of treatment, Grade 1-4 CK elevations occurred in 79% of TYZEKA-treated subjects and 47% of lamivudine-treated subjects. Grade 3 or 4 CK elevations occurred in 13% of TYZEKA-treated subjects and 4% of lamivudine-treated subjects.
subjects. Most CK elevations were asymptomatic, but the mean recovery time was longer for subjects on TYZEKA than subjects on lamivudine.

Among TYZEKA-treated subjects with Grade 1-4 CK elevations, 10% developed a musculoskeletal adverse event compared to 5% of lamivudine-treated subjects. A total of 2% (13/847) TYZEKA-treated subjects interrupted or discontinued trial drug due to CK elevation or musculoskeletal adverse events.\(^1\)

\(^1\)Includes the Preferred Terms: back pain, chest wall pain, non-cardiac chest pain, chest discomfort, flank pain, muscle cramp, muscular weakness, musculoskeletal pain, musculoskeletal chest pain, musculoskeletal discomfort, musculoskeletal stiffness, myalgia, myofascial pain syndrome, myopathy, myositis, neck pain, and pain in extremity.

**ALT Flares During Treatment**

The incidence of ALT flares, defined as ALT greater than 10 x ULN and greater than 2 x baseline, was similar in the two treatment arms (3%) in the first six months. After Week 24, ALT flares were reported less frequently in the TYZEKA arm (2%) compared to the lamivudine arm (5%). Periodic monitoring of hepatic function is recommended during chronic hepatitis B treatment.

**Exacerbations of Hepatitis after Discontinuation of Treatment**

In the subset of subjects who discontinued treatment prematurely for reasons other than efficacy, or who elected not to continue TYZEKA in another clinical trial, 9/154 (6%) TYZEKA-treated and 10/180 (6%) lamivudine-treated subjects experienced an exacerbation of hepatitis (ALT elevation greater than 2 x baseline and greater than 10 x ULN) in the 4-month post-treatment period.

**Results at 208 Weeks**

After 104 weeks of blinded therapy in trials 007 GLOBE and NV-02B-015, 667 subjects received TYZEKA in an open-label extension trial, CLDT600A2303. Of those initially randomized to TYZEKA therapy, 78% of subjects (530/680) from trial 007 GLOBE and 82% (137/167) of subjects from trial NV-02B-015 enrolled into the extension trial and continued TYZEKA treatment for up to 208 weeks. The long-term TYZEKA safety population in trial CLDT600A2303 consisted of 655 subjects, including 518 subjects from trial 007 GLOBE and 137 subjects from trial NV-02B-015.

The overall safety profile from the pooled analysis up to 104 and 208 weeks was similar. Grade 3/4 CK elevations occurred in 16% of subjects (104/655) treated with TYZEKA in trial CLDT600A2303. Most Grade 3/4 CK elevations were asymptomatic (74% of subjects without any muscle related adverse reaction) and transient (98% of episodes lasted one or two visits (visit interval 2 - 12 weeks) and 87% of subjects had one or two episodes). Most Grade 3/4 CK elevations (93%) resolved spontaneously or returned to baseline levels. Two cases of myopathy and two cases of myositis were reported in the 655 TYZEKA-treated subjects.

Among the cohort of 655 subjects continuing TYZEKA for up to 208 weeks in trial CLDT600A2303, including the subgroup of patients (n = 223) with mild renal impairment [estimated glomerular filtration rate (eGFR) 60-90 mL per min)] at baseline, mean estimated GFR assessed by modification of diet in renal disease (MDRD) did not decline.

**6.2 Postmarketing Experience**

The following adverse reactions have been reported during post approval use of TYZEKA. Because these reactions were reported voluntarily from a population of unknown size, it is not possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

**Musculoskeletal, and Connective Tissue Disorders:** Rhabdomyolysis

**Nervous System Disorders:** Paraesthesia, hypoaesthesia

**Metabolism and Nutrition Disorders:** Lactic acidosis
7 DRUG INTERACTIONS

TYZEKA is excreted mainly by passive diffusion so the potential for interactions between TYZEKA and other drugs eliminated by renal excretion is low. However, because TYZEKA is eliminated primarily by renal excretion, concomitant administration of TYZEKA with drugs that alter renal function may alter plasma concentrations of TYZEKA [see Clinical Pharmacology (12.3)].

A clinical trial investigating the combination of TYZEKA, 600 mg daily, with pegylated interferon alfa-2a, 180 micrograms once weekly by subcutaneous administration, indicates that this combination is associated with an increased risk of peripheral neuropathy occurrence and severity, in comparison to TYZEKA or pegylated interferon alfa-2a alone [see Contraindications (4) and Warnings and Precautions (5.4)].

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Pregnancy Exposure Registry

There is a pregnancy exposure registry that monitors pregnancy outcomes in women exposed to TYZEKA during pregnancy. Healthcare providers are encouraged to register patients by calling the Antiretroviral Pregnancy Registry (APR) at 1-800-258-4263.

Risk Summary

Prospective pregnancy data from the APR are insufficient to adequately assess the risk of birth defects, miscarriage or adverse maternal or fetal outcomes. Available data from the APR show no increase in the overall risk of major birth defects of telbivudine compared with the background rate for major birth defects of 2.7% in a U.S. reference population of the Metropolitan Atlanta Congenital Defects Program (MACDP). The rate of miscarriage is not reported in the APR. All pregnancies have a background risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of miscarriage is 15%-20% of clinically recognized pregnancies.

In animal reproduction studies, no adverse developmental effects were observed in rats and rabbits at exposures 6- and 37-times higher, respectively, than those observed at the recommended human dose (RHD) of 600 mg per day (see Data).

Data

Human Data

Based on prospective reports to the APR of live births following exposure to telbivudine-containing regimens during pregnancy (including 245 exposed in the first trimester and 13 exposed in the second/third trimester), the prevalence of birth defects in live births was 1.2% (95% confidence interval [CI]: 0.3% to 3.5%) for telbivudine with first trimester exposures; the 13 second/third trimester exposures reported to the APR are insufficient to make a risk assessment. Prospective reports from the APR of overall major birth defects in pregnancies exposed TYZEKA is compared with a U.S. background major birth defect rate. Methodological limitations of the APR include the use of MACDP as the external comparator group. Limitations of using an external comparator include differences in methodology and populations, as well as confounding due to the underlying disease.

Animal Data

In an embryo-fetal development study, pregnant rabbits received telbivudine at oral doses of 50, 250 or 1000 mg/kg/day during the period of organogenesis (gestation Day 6 to 18). No adverse embryo-fetal effects were observed at doses up to 1000 mg/kg/day. At this dose level, an increased incidence of abortions associated with maternal toxicity was observed at an exposure approximately 37 times the systemic exposure at the RHD of 600 mg per day.

In a prenatal and postnatal development study, pregnant rats received oral doses of telbivudine at 100, 250, or 1000 mg/kg/day from organogenesis, through late gestation, delivery, and lactation (gestation Day 7 to

Reference ID: 4368857
lactation/postpartum Day 20). No adverse effects were observed at doses up to 1000 mg/kg/day, at exposures approximately 6 times the systemic exposure at the RHD.

8.2 Lactation

Risk Summary

It is not known whether TYZEKA is present in human breast milk, affects human milk production, or has effects on the breastfed infant. Telbivudine is present in the milk of rats (see Data). The developmental and health benefits of breastfeeding should be considered along with the mother’s clinical need for TYZEKA and any potential adverse effects on the breastfed infant from TYZEKA or from the underlying maternal condition.

Data

Telbivudine was the primary drug-related component excreted into the milk of lactating rats following a single oral dose of 10 mg per kg on lactation Day 12, with milk concentrations of up to approximately 2.8 times that of maternal plasma concentrations observed 3 hours post-dose.

8.4 Pediatric Use

Safety and effectiveness of TYZEKA in pediatric patients have not been established.

Juvenile Animal Toxicity Data

In a juvenile animal study, reduced fertility associated with oral telbivudine doses \( \geq 1000 \) mg/kg/day was observed when administered to sexually immature rats (from postpartum Day 14 to 70). No fertility effects were observed in sexually mature animals [see Nonclinical Toxicology (13.1)].

8.5 Geriatric Use

Clinical trials of TYZEKA did not include sufficient numbers of subjects aged 65 and older to determine whether they respond differently from younger subjects. In general, caution should be exercised when prescribing TYZEKA to elderly patients, considering the greater frequency of decreased renal function due to concurrent disease or other drug therapy. Renal function should be monitored in elderly patients, and dosage adjustments should be made accordingly [see Clinical Pharmacology (12.3) and Dosage and Administration (2.2)].

8.6 Renal Impairment

TYZEKA is eliminated primarily by renal excretion, therefore dose regimen adjustment is recommended in patients with CrCl less than 50 mL per min, including patients with ESRD requiring hemodialysis [see Dosage and Administration (2.2) and Clinical Pharmacology (12.3)].

8.7 Liver Transplant Recipients

The safety and efficacy of TYZEKA in liver transplant recipients have not been evaluated. The steady-state pharmacokinetics of TYZEKA was not altered following multiple dose administration in combination with cyclosporine. If TYZEKA treatment is determined to be necessary for a liver transplant recipient who has received or is receiving an immunosuppressant that may affect renal function, such as cyclosporine or tacrolimus, renal function should be monitored both before and during treatment with TYZEKA [see Clinical Pharmacology (12.3) and Dosage and Administration (2.2)].

8.8 Co-infected Patients

TYZEKA has not been investigated in co-infected hepatitis B patients (e.g., patients co-infected with HIV, HCV, or HDV).

8.9 Racial/Ethnic Minorities

The safety and efficacy of TYZEKA have not been evaluated in Black/African American or Hispanic patients. It is not known if safety and efficacy can be extrapolated from studied populations.
10 OVERDOSAGE

There is no information on intentional overdose of TYZEKA, but one subject experienced an unintentional and asymptomatic overdose. Healthy subjects who received TYZEKA doses up to 1800 mg per day for 4 days had no increase in or unexpected adverse events. A maximum tolerated dose for TYZEKA has not been determined. In the event of an overdose, TYZEKA should be discontinued, the patient must be monitored for evidence of toxicity, and appropriate general supportive treatment applied as necessary.

In case of overdosage, hemodialysis may be considered. Within 2 hours, following a single 200 mg dose of telbivudine, a 4-hour hemodialysis session removed approximately 23% of the telbivudine dose.

11 DESCRIPTION

TYZEKA is the trade name for telbivudine, a synthetic thymidine nucleoside analogue with activity against hepatitis B virus (HBV). The chemical name for telbivudine is 1-((2S,4R,5S)-4-hydroxy-5-hydroxymethyltetrahydrofuran-2-y1)-5-methyl-1H-pyrimidine-2,4-dione, or 1-(2-deoxy-β-L-ribofuranosyl)-5-methyluracil. Telbivudine is the unmodified β-L enantiomer of the naturally occurring nucleoside, thymidine. Its molecular formula is C_{10}H_{14}N_{2}O_{5}, which corresponds to a molecular weight of 242.23 g/mol. Telbivudine has the following structural formula:

![Telbivudine Structural Formula](image)

Telbivudine is a white to slightly yellowish powder. Telbivudine is sparingly soluble in water (greater than 20 mg per mL), and very slightly soluble in absolute ethanol (0.7 mg per mL) and n-octanol (0.1 mg per mL).

TYZEKA film-coated tablets are available for oral administration in 600 mg strength. TYZEKA 600 mg film-coated tablets contain the following inactive ingredients: colloidal silicon dioxide, magnesium stearate, microcrystalline cellulose, povidone, and sodium starch glycolate. The tablet coating contains titanium dioxide, polyethylene glycol, talc, and hypromellose.

TYZEKA oral solution is available for oral administration in 100 mg per 5 mL strength. TYZEKA oral solution contains the following inactive ingredients: citric acid anhydrous, benzoic acid, passion fruit flavor, sodium saccharin, sodium hydroxide, and purified water. A 600 mg dose (30 mL) of TYZEKA oral solution contains approximately 47 mg of sodium.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

TYZEKA is an antiviral drug against the hepatitis B virus [see Microbiology (12.4)].

12.2 Pharmacodynamics

In a randomized, partially single-blinded, placebo and active-controlled, four-period crossover trial, 53 healthy subjects were administered TYZEKA 600 mg, a supratherapeutic TYZEKA 1800 mg dose, placebo, and moxifloxacin 400 mg. After 7 days of dosing, TYZEKA did not prolong the QT interval. The maximum placebo-adjusted mean (upper 1-side 95% CI) change from baseline in QTcF on Day 7 were 3.4 msec (5.9 msec) for the 600 mg and 4.4 msec (6.9 msec) for the 1800 mg dosing regimens.
12.3 Pharmacokinetics in Adults

The single- and multiple-dose pharmacokinetics of TYZEKA were evaluated in healthy subjects and in patients with chronic hepatitis B. TYZEKA pharmacokinetics are similar between both populations.

Absorption and Bioavailability

Following oral administration of TYZEKA 600 mg once-daily in healthy subjects (n = 12), steady state peak plasma concentration (C_max) was 3.69 ± 1.25 microgram per mL (mean ± SD) which occurred between 1 and 4 hours (median 2 hours), area under the curve (AUC) was 26.1 ± 7.2 microgram * hour per mL (mean ± SD), and trough plasma concentrations (C_trough) were approximately 0.2-0.3 microgram per mL. Steady state was achieved after approximately 5 to 7 days of once-daily administration with ~1.5-fold accumulation, suggesting an effective half-life of ~15 hours.

Effects of Food on Oral Absorption

TYZEKA absorption and exposure were unaffected when a single 600 mg dose was administered with a high-fat (~55 g), high-calorie (~950 kcal) meal. TYZEKA may be taken with or without food.

Distribution

*In vitro* binding of telbivudine to human plasma proteins is low (3.3%). After oral dosing, the estimated apparent volume of distribution is in excess of total body water, suggesting that telbivudine is widely distributed into tissues. Telbivudine was equally partitioned between plasma and blood cells.

Metabolism and Elimination

No metabolites of telbivudine were detected following administration of [14C]-telbivudine in humans. Telbivudine is not a substrate, or inhibitor of the cytochrome P450 (CYP450) enzyme system.

After reaching the peak concentration, plasma concentrations of TYZEKA declined in a biexponential manner with a terminal elimination half-life (T_1/2) of 40-49 hours. TYZEKA is eliminated primarily by urinary excretion of unchanged drug. The renal clearance of TYZEKA approaches normal glomerular filtration rate suggesting that passive diffusion is the main mechanism of excretion. Approximately 42% of the dose is recovered in the urine over 7 days following a single 600 mg oral dose of TYZEKA. Because renal excretion is the predominant route of elimination, patients with moderate to severe renal dysfunction and those undergoing hemodialysis require a dose regimen adjustment [see Dosage and Administration (2.2)].

Special Populations

Gender: There are no significant gender-related differences in TYZEKA pharmacokinetics.

Race: There are no significant race-related differences in TYZEKA pharmacokinetics.

Pediatrics and Geriatrics: Pharmacokinetic studies have not been conducted in children or elderly subjects.

Renal Impairment: Single-dose pharmacokinetics of TYZEKA have been evaluated in subjects (without chronic hepatitis B) with various degrees of renal impairment (as assessed by creatinine clearance). Based on the results shown in Table 4, adjustment of the dose regimen for TYZEKA is recommended in patients with creatinine clearance of less than 50 mL per min [see Dosage and Administration (2.2)].

<table>
<thead>
<tr>
<th>Renal Function (CrCl in mL/min)</th>
<th>C_max (µg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal (greater than 80) (n = 8) 600 mg</td>
<td>3.4 ± 0.9</td>
</tr>
<tr>
<td>Mild (50-80) (n = 8) 600 mg</td>
<td>3.2 ± 0.9</td>
</tr>
<tr>
<td>Moderate (30-49) (n = 8) 400 mg</td>
<td>2.8 ± 1.3</td>
</tr>
<tr>
<td>Severe (less than 30) (n = 6) 200 mg</td>
<td>1.6 ± 0.8</td>
</tr>
<tr>
<td>ESRD/Hemodialysis (n = 6) 200 mg</td>
<td>2.1 ± 0.9</td>
</tr>
</tbody>
</table>

Reference ID: 4368857
### Pharmacokinetic Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value (µg•hr/mL)</th>
<th>Value (L/hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AUC_{0-INF}</td>
<td>28.5 ± 9.6</td>
<td>7.6 ± 2.9</td>
</tr>
<tr>
<td>CL_{RENAL}</td>
<td>32.5 ± 10.1</td>
<td>5.0 ± 1.2</td>
</tr>
<tr>
<td></td>
<td>36.0 ± 13.2</td>
<td>2.6 ± 1.2</td>
</tr>
<tr>
<td></td>
<td>32.5 ± 13.2</td>
<td>0.7 ± 0.4</td>
</tr>
<tr>
<td></td>
<td>67.4 ± 36.9</td>
<td></td>
</tr>
</tbody>
</table>

### Renally Impaired Subjects on Hemodialysis

Hemodialysis (up to 4 hours) reduces systemic TYZEKA exposure by approximately 23%. Following dose regimen adjustment for creatinine clearance [see Dosage and Administration (2.2)], no additional dose modification is necessary during routine hemodialysis. When administered on hemodialysis days, TYZEKA should be administered after hemodialysis.

### Hepatic Impairment

The pharmacokinetics of TYZEKA following a single 600 mg dose have been studied in subjects (without chronic hepatitis B) with various degrees of hepatic impairment. There were no changes in TYZEKA pharmacokinetics in hepatically impaired subjects compared to unimpaired subjects. Results of these studies indicate that no dosage adjustment is necessary for patients with hepatic impairment.

### Drug Interactions

Drug-drug interaction studies show that lamivudine, adefovir dipivoxil, cyclosporine, pegylated interferon alfa-2a, and tenofovir disoproxil fumarate do not alter TYZEKA pharmacokinetics. In addition, TYZEKA does not alter the pharmacokinetics of lamivudine, adefovir dipivoxil, cyclosporine, or tenofovir disoproxil fumarate. No definitive conclusion could be drawn regarding the effects of TYZEKA on the pharmacokinetics of pegylated interferon alfa-2a due to the high inter-individual variability of pegylated interferon alfa-2a concentrations. At concentrations up to 12 times that in humans, telbivudine did not inhibit \textit{in vitro} metabolism mediated by any of the following human hepatic microsomal cytochrome P450 (CYP) isoenzymes known to be involved in human medicinal product metabolism: 1A2, 2C9, 2C19, 2D26, 2E1, and 3A4. Based on the above results and the known elimination pathway of telbivudine, the potential for CYP450-mediated interactions involving telbivudine with other medicinal products is low.

### 12.4 Microbiology

#### Mechanism of Action

Telbivudine is a synthetic thymidine nucleoside analogue with activity against HBV DNA polymerase. It is phosphorylated by cellular kinases to the active triphosphate form, which has an intracellular half-life of 14 hours. Telbivudine 5'-triphosphate inhibits HBV DNA polymerase (reverse transcriptase) by competing with the natural substrate, thymidine 5'-triphosphate. Incorporation of telbivudine 5'-triphosphate into viral DNA causes DNA chain termination. Telbivudine is an inhibitor of both HBV first strand (EC_{50} value = 1.3 ± 1.6 micromolar) and second strand synthesis (EC_{50} value = 0.2 ± 0.2 micromolar). Telbivudine 5'-triphosphate at concentrations up to 100 micromolar did not inhibit human cellular DNA polymerases α, β, or γ. No appreciable mitochondrial toxicity was observed in HepG2 cells treated with telbivudine at concentrations up to 10 micromolar.

#### Antiviral Activity

The antiviral activity of telbivudine was assessed in the HBV-expressing human hepatoma cell line 2.2.15, as well as in primary duck hepatocytes infected with duck hepatitis B virus. The concentration of telbivudine that effectively inhibited 50% of viral DNA synthesis (EC_{50}) in both systems was approximately 0.2 micromolar. The anti-HBV activity of telbivudine was additive with adefovir in cell culture, and was not antagonized by the HIV NRTIs didanosine and stavudine. Telbivudine was not antagonistic to the anti-HIV activity of abacavir, didanosine, emtricitabine, lamivudine, stavudine, tenofovir, or zidovudine. Transient reductions in HIV-1 RNA have been seen in some patients after administration of telbivudine in the absence of antiretroviral therapy. The clinical significance of these reductions has not been determined.

### Resistance

**Trial NV-02B-007 (007 GLOBE):** In an as-treated analysis of the Phase III global registration trial, 59% (251/429) of treatment-naïve HBeAg-positive and 89% (202/227) of treatment-naïve HBeAg-negative subjects...
receiving TYZEKA 600 mg once daily achieved undetectable serum HBV DNA levels (less than 300 copies per mL) by Week 52. Of those who continued treatment beyond Week 52, 58% (243/418) and 85% (190/224) of HBeAg-positive and HBeAg-negative TYZEKA recipients, respectively, had undetectable HBV DNA at Week 104 (or at the end of dosing in treatment Year 2).

Genotypic analysis of paired baseline and treatment failure isolates from 181 evaluable subjects with amplifiable HBV DNA and greater than or equal to 16 weeks of TYZEKA treatment showed that the rtM204I/V substitution was associated with virologic failure (HBV DNA greater than or equal to 1,000 copies per mL) and virologic rebound (HBV DNA greater than or equal to 1 log₁₀ increase above nadir). The rtM204I/V substitution was detectable in isolates from 78% (142/181) of evaluable subjects, and was frequently found with substitutions rtL80I/V and rtL180M. The rtM204I/V substitution was found infrequently with rtV27A, rtL82M, rtV173L, rtT184I/S, rtA200V, rtL229F/V/W, and rtR289K substitutions. The HBV of 16 subjects developed rtA181S/T amino acid substitutions while receiving TYZEKA. Eight of these 16 subjects had outgrowth of HBV expressing an rtM204I/V substitution without the rtA181S/T substitutions and 1 subject’s HBV had both the rtM204I and rtA181T substitutions.

Trial CLDT600A2303: After 2 years of TYZEKA monotherapy in the 007 GLOBE trial, 77% (505/656) of subjects entered the open-label CLDT600A2303 extension trial to continue TYZEKA for up to 2 additional years, including 349 subjects who had undetectable levels of HBV DNA and 156 subjects who were viremic at entry. The rtM204I/V substitution was detectable in the virus from 83% (39/47) of the subjects losing viral suppression and having evaluable genotypic data. Of evaluable viremic subjects entering the extension, 25/33 (76%) developed rtM204I/V substitutions. Overall, 64 subjects developed genotypic resistance to TYZEKA with evidence of emerging rtM204I/V substitutions during the 2 years of TYZEKA treatment in this extension trial.

Subjects with higher baseline viral load had higher rates of genotypic resistance to TYZEKA, while subjects who achieved HBV DNA levels less than 300 copies per mL at Week 24 had lower rates of genotypic resistance to TYZEKA. The cumulative frequency of genotypic resistance (emergence of the rtM204I/V substitution) to TYZEKA in nucleos(t)ide treatment-naïve subjects was 7% and 22% at Weeks 52 and 104 of the controlled 007 GLOBE trial, and 30% and 35% at Weeks 156 and 208 of the open-label extension trial (CLDT600A2303), respectively (Table 5).

One-hundred-sixty-seven subjects (25% of those in the 007 GLOBE trial) were treated with TYZEKA according to current dosing recommendations [see Indications and Usage (1.1)]. Eighty-four percent (140/167) of these subjects qualified at 24 weeks for continued TYZEKA treatment (HBV DNA less than 300 copies per mL). Retrospective calculation of the cumulative rate of genotypic resistance to TYZEKA for this subgroup of subjects was 0%, 3%, 12%, and 16% at Weeks 52, 104, 156, and 208, respectively (Table 5).

<table>
<thead>
<tr>
<th>Table 5</th>
<th>Cumulative Rates of Genotypic Resistance to TYZEKA through Week 208</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trial</td>
<td>Overall Study Population</td>
</tr>
<tr>
<td></td>
<td>Cumulative genotypic resistance rate¹</td>
</tr>
<tr>
<td>NV-02B-007 (007 GLOBE trial)</td>
<td>Week 52</td>
</tr>
<tr>
<td></td>
<td>Week 104</td>
</tr>
<tr>
<td>CLDT600A2303 (104-week extension trial)</td>
<td>Week 156</td>
</tr>
<tr>
<td></td>
<td>Week 208</td>
</tr>
</tbody>
</table>

¹The cumulative rates of genotypic resistance to TYZEKA were calculated using the formula previously described by Pawlotsky et al. (2008).

²TYZEKA dosing recommendations are provided in this Package Insert [see Indications and Usage (1.1) and Dosage and Administration (2.1)].
Cross-Resistance

Cross-resistance has been observed among HBV nucleos(t)ide analogues. In cell-based assays, lamivudine-resistant HBV strains expressing either the rtM204I substitution or the rtL180M/rtM204V double substitution had greater than or equal to 1,000-fold reduced susceptibility to telbivudine. Telbivudine retained wild-type phenotypic activity (1.2-fold reduction) against HBV expressing rtM204V alone. Most subjects (92%, 155/169) whose virus developed lamivudine resistance-associated substitutions (rtM204I/V) during 2 years of lamivudine treatment in the 007 GLOBE trial remained viremic (HBV DNA greater than 300 copies per mL) after up to 2 years of TYZEKA monotherapy in the CLDT600A2303 extension trial, including 91% (50/55) of the subjects with the rtM204V substitution.

HBV encoding the adefovir resistance-associated substitution rtA181V showed 3- to 5-fold reduced susceptibility to telbivudine in cell culture. The rtA181S and rtA181T substitutions conferred 2.7- and 3.5-fold reductions in susceptibility to telbivudine, respectively. The rtA181T substitution is associated with decreased clinical response in subjects with HBV treated with adefovir and entecavir. HBV encoding the adefovir resistance-associated substitution rtN236T remained susceptible to telbivudine.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Telbivudine has shown no carcinogenic potential. Long term oral carcinogenicity studies with telbivudine were negative in mice and rats at exposures up to 14 times those observed in humans at the therapeutic dose of 600 mg per day.

There was no evidence of genotoxicity based on in vitro or in vivo tests. Telbivudine was not mutagenic in the Ames bacterial reverse mutation assay using S. typhimurium and E. coli strains with or without metabolic activation. Telbivudine was not clastogenic in mammalian-cell gene mutation assays, including human lymphocyte cultures and an assay with Chinese hamster ovary cells with or without metabolic activation. Furthermore, telbivudine showed no effect in an in vivo micronucleus study in mice.

Effects on fertility were studied in rats administered telbivudine as juveniles or adults. Juvenile rats were treated with telbivudine at doses of 0, 250, 1000, and 2000 mg per kg per day from post natal days 14 to 70. These rats were mated following a 5 week drug-free recovery period. Up to 50% reduction of fertility was associated with doses 1000 mg per kg per day and higher, which was equivalent to a systemic exposure approximately 7.5 times that achieved in humans at the therapeutic dose. The no observed adverse effect level (NOAEL) for effects on fertility or mating parameters was 250 mg per kg per day, which was equivalent to systemic exposure levels 2.5 to 2.8 times that achieved in humans at the therapeutic dose. In contrast, such reduction of fertility was absent in adult rats, when either male or female rats were treated with telbivudine at doses up to 2000 mg per kg per day, equivalent to a systemic exposure approximately 14 times that achieved in humans at the therapeutic dose.

14 CLINICAL STUDIES

14.1 Clinical Experience in Nucleoside-Naïve Adults

The safety and efficacy of long-term (104-week) TYZEKA treatment were evaluated in one active-controlled, clinical trial (NV-02B-007 GLOBE Trial) that included 1,367 subjects with chronic hepatitis B and a smaller supportive trial (NV-02B-015) that included 332 subjects. Subjects were 16 years of age or older, with chronic hepatitis B, evidence of HBV infection with viral replication (HBsAg-positive, HBeAg-positive or HBeAg-negative, HBV DNA detectable by a PCR assay), and elevated ALT levels greater than or equal to 1.3 x ULN, no evidence of hepatic decompensation, and chronic inflammation on liver biopsy compatible with chronic viral hepatitis.

NV-02B-007 GLOBE Trial

The Week 52 and Week 104 results of the 007 GLOBE trial are summarized below.
The 007 GLOBE trial was a Phase III, randomized, double-blind, multinational trial of TYZEKA 600 mg once daily compared to lamivudine 100 mg once daily for a treatment period of 104 weeks in 1,367 (n = 680 TYZEKA; n = 687 lamivudine) nucleoside-naïve chronic hepatitis B HBeAg-positive and HBeAg-negative subjects. The primary data analysis was conducted after all subjects had reached Week 52.

**HBeAg-positive Subjects** (n = 458 TYZEKA; n = 463 lamivudine) The mean age of subjects was 32 years, 74% were male, 82% were Asian, 12% were Caucasian, and 6% had previously received alfa-interferon therapy. At baseline, subjects had a mean Knodell Necroinflammatory Score greater than or equal to 7; mean serum HBV DNA as measured by Roche COBAS Amplicor® PCR assay was 9.52 log10 copies per mL; and mean serum ALT was 153 IU per L. Pre- and post-liver biopsy samples were adequate for 86% of subjects.

**HBeAg-negative Subjects** (n = 222 TYZEKA; n = 224 lamivudine) The mean age of subjects was 43 years, 77% were male, 65% were Asian, 23% were Caucasian, and 11% had previously received alfa-interferon therapy. At baseline, subjects had a mean Knodell Necroinflammatory Score greater than or equal to 7; mean serum HBV DNA as measured by Roche COBAS Amplicor® PCR assay was 7.54 log10 copies per mL; and mean serum ALT was 140 IU per L. Pre- and post-liver biopsy samples were adequate for 92% of subjects.

**Clinical Results**

Clinical and virologic efficacy endpoints were evaluated separately in the HBeAg-positive and HBeAg-negative subject populations.

The primary endpoint of Therapeutic Response at Week 52 was a composite endpoint requiring suppression of HBV DNA to less than 5 log10 copies per mL in conjunction with either loss of serum HBeAg or ALT normalization. Key secondary endpoints included histologic response, ALT normalization, and measures of virologic response.

At Week 52, in HBeAg-positive subjects, 75% of TYZEKA subjects and 67% of lamivudine subjects had a Therapeutic Response; in HBeAg-negative subjects, 75% of TYZEKA subjects and 77% of lamivudine subjects had a Therapeutic Response.

Analysis of the histological response at Week 52 is shown in Table 6.

<table>
<thead>
<tr>
<th>Table 6</th>
<th>Histological Improvement and Change in Ishak Fibrosis Score at Week 52 (007 GLOBE Trial)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HBeAg-positive (n = 797)</td>
</tr>
<tr>
<td></td>
<td>TYZEKA 600 mg (n = 399)</td>
</tr>
<tr>
<td><strong>Histologic Response</strong>&lt;sup&gt;2&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>Improvement</td>
<td>69%</td>
</tr>
<tr>
<td>No Improvement</td>
<td>19%</td>
</tr>
<tr>
<td>Missing Week 52 Biopsy</td>
<td>12%</td>
</tr>
<tr>
<td><strong>Ishak Fibrosis Score</strong>&lt;sup&gt;3&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>Improvement</td>
<td>41%</td>
</tr>
<tr>
<td>No Change</td>
<td>39%</td>
</tr>
<tr>
<td>Worsening</td>
<td>9%</td>
</tr>
<tr>
<td>Missing Week 52 Biopsy</td>
<td>12%</td>
</tr>
</tbody>
</table>

<sup>1</sup>Subjects with greater than or equal to one dose of trial drug with evaluable baseline liver biopsies and baseline Knodell Necroinflammatory Score greater than or equal to 2.

<sup>2</sup>Histologic Response defined as greater than or equal to 2 point decrease in Knodell Necroinflammatory Score from baseline with no worsening of the Knodell Fibrosis Score.

<sup>3</sup>For Ishak Fibrosis Score, improvement defined as greater than or equal to a 1-point reduction in Ishak Fibrosis Score from baseline to
Subjects were eligible to continue blinded treatment to Week 104. In the ITT population, 624/680 (92%) TYZEKA recipients and 599/687 (87%) lamivudine recipients completed trial treatment to Week 104. At Week 104, in HBeAg-positive subjects, 63% of TYZEKA subjects and 48% of lamivudine subjects had a Therapeutic Response, while in HBeAg-negative subjects 78% of TYZEKA subjects and 66% of lamivudine subjects had a Therapeutic Response.

Selected virologic, biochemical, and serologic outcome measures at Weeks 52 and 104 are shown in Table 7.

Table 7 Virological, Biochemical and Serologic Endpoints at Weeks 52 and 104 (007 GLOBE Trial)

<table>
<thead>
<tr>
<th>Response Parameter</th>
<th>HBeAg-positive (n = 921)</th>
<th>HBeAg-negative (n = 446)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>TYZEKA 52</td>
<td>Lamivudine 52</td>
</tr>
<tr>
<td></td>
<td>600 mg (n = 458)</td>
<td>100 mg (n = 463)</td>
</tr>
<tr>
<td>Mean HBV DNA Reduction from Baseline (log_{10} copies/mL) ± SEM</td>
<td>-6.45 (0.11)</td>
<td>-5.74 (0.15)</td>
</tr>
<tr>
<td>% Subjects HBV DNA undetectable by PCR</td>
<td>60%</td>
<td>56%</td>
</tr>
<tr>
<td>ALT Normalization</td>
<td>77%</td>
<td>70%</td>
</tr>
<tr>
<td>HBeAg Seroconversion</td>
<td>23%</td>
<td>30%</td>
</tr>
<tr>
<td>HBeAg Loss</td>
<td>26%</td>
<td>35%</td>
</tr>
</tbody>
</table>

1Roche COBAS Amplicor® Assay (LLOQ less than or equal to 300 copies/mL).
2ALT normalization assessed only in subjects with ALT greater than ULN at baseline.
3HBeAg seroconversion and loss assessed only in subjects with detectable HBeAg at baseline.

Subjects who achieved non-detectable HBV DNA levels at 24 weeks were more likely to undergo e-antigen seroconversion, achieve undetectable levels of HBV DNA, normalize ALT, and were less likely to develop resistance at one and two years.

NV-02B-015 Trial

The efficacy results of the 007 GLOBE trial were supported by results of trial NV-02B-015. This was a Phase III, randomized, double-blind, trial of TYZEKA 600 mg once daily compared to lamivudine 100 mg once daily for a treatment period of 104 weeks in 332 (n = 167 TYZEKA; n = 165 lamivudine) nucleoside-naïve chronic hepatitis B HBeAg-positive and HBeAg-negative Chinese subjects. The primary efficacy endpoint was serum HBV DNA reduction from baseline. In this trial, the composite endpoint Therapeutic Response was a key secondary endpoint. Histological response was not assessed as an outcome measure in this trial.

Clinical Results

Among HBeAg-positive subjects (n = 147 TYZEKA; n = 143 lamivudine) results for key endpoints at Week 104 included Therapeutic Response (66% vs. 41%), mean HBV DNA reduction (-5.47 vs. -3.97 log_{10} copies per mL), HBV DNA PCR negativity (58% vs. 34%), ALT normalization (73% vs. 59%), HBeAg loss (40% vs. 28%) and HBeAg seroconversion (29% vs. 20%), for TYZEKA and lamivudine, respectively. Because the number of HBeAg-negative subjects in this trial was small (n = 42), definitive conclusions could not be drawn regarding efficacy outcomes in this subpopulation.
16 HOW SUPPLIED/STORAGE AND HANDLING

Tablets
TYZEKA 600 mg tablets are white to slightly yellowish film-coated, ovaloid-shaped tablets, imprinted with “LDT” on one side.
Bottle of 30 tablets (NDC 0078-0538-15) with child-resistant closure.
Store TYZEKA tablets in original container at 25°C (77°F), excursions permitted to 15°C-30°C (59°F-86°F) [see USP Controlled Room Temperature].

Oral Solution
TYZEKA (telbivudine) oral solution is a clear, colorless to pale yellow, passion fruit flavored liquid. TYZEKA oral solution contains 100 mg of telbivudine per 5 milliliters.
Bottle containing 300 mL oral solution (NDC 0078-0539-85) with child-resistant closure and embossed dosing cup. The dosing cup is intended for measurement of TYZEKA oral solution only.
Store TYZEKA oral solution in original container at 25°C (77°F), excursions permitted to 15°C-30°C (59°F-86°F) [see USP Controlled Room Temperature]. Use within two months after opening the bottle. Do not freeze.

For all medical inquiries call: 1-877-8-TYZEKA (1-877-889-9352).
Keep this and all drugs out of the reach of children.

17 PATIENT COUNSELING INFORMATION
See FDA-approved patient labeling (Medication Guide and Instructions for Use)

Severe Acute Exacerbation of Hepatitis after Discontinuation of Treatment
Inform patients that discontinuation of anti-hepatitis B therapy, including TYZEKA, may result in severe acute exacerbations of hepatitis B. Advise the patient to not discontinue TYZEKA without first informing their healthcare provider [see Warnings and Precautions (5.1)].

Lactic Acidosis and Severe Hepatomegaly
Lactic acidosis and severe hepatomegaly with steatosis, including fatal cases, have been reported with use of drugs similar to TYZEKA. Advise patients to contact their healthcare provider immediately and stop TYZEKA if they develop clinical symptoms suggestive of lactic acidosis or pronounced hepatotoxicity [see Warnings and Precautions (5.2)].

Myopathy and Rhabdomyolysis
Advise patients that cases of myopathy/myositis have been reported with TYZEKA and to report promptly unexplained muscle weakness, tenderness, or pain [see Warnings and Precautions (5.3)].

Peripheral Neuropathy
Inform patients that peripheral neuropathy has been reported with TYZEKA alone or in combination with pegylated interferon alfa-2a and other interferons. Advise patients to report promptly any numbness, tingling, and/or burning sensations in the arms and/or legs, with or without difficulty walking [see Warnings and Precautions (5.4)].

Important Administration Instructions
Inform patients that it is important to take TYZEKA on a regular dosing schedule with or without food and to avoid missing doses as it can result in the development of resistance [see Dosage and Administration (2.1)]. Inform patients to discuss with their physician any changes in their regimen or discontinuation as deterioration
of liver disease may occur in some cases with treatment discontinuation. Advise patients on a low sodium diet that TYZEKA oral solution contains approximately 47 mg of sodium per 600 mg dose (30 mL).

Treatment Duration

Advise patients that in the treatment of chronic hepatitis B, the optimal duration of treatment is unknown. The relationship between response and long-term prevention of outcomes such as hepatocellular carcinoma is not known.

Pregnancy Registry

Advise patients that there is a pregnancy exposure registry that monitors pregnancy outcomes in women exposed to TYZEKA during pregnancy [see Use in Specific Populations (8.1)].

Distributed by:
Novartis Pharmaceuticals Corporation
East Hanover, New Jersey 07936
T201X-XX
TYZEKA® (Tie-zee′-ka) (telbivudine) tablets

TYZEKA® (Tie-zee′-ka) (telbivudine) oral solution

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

TYZEKA can cause serious side effects, including:

Worsening of your hepatitis B virus (HBV) infection after discontinuation. Your HBV infection may get worse or become very serious if you stop taking TYZEKA.
- Take your TYZEKA exactly as your healthcare provider tells you to.
- Do not let your TYZEKA run out. Refill your medicine or talk to your healthcare provider before your TYZEKA is all gone.
- Do not stop taking your TYZEKA without talking to your healthcare provider.

Your healthcare provider will monitor your health and do regular blood tests to check your liver if you stop taking TYZEKA. Tell your healthcare provider right away about any new or unusual symptoms that happen after you stop taking TYZEKA.

Too much lactic acid in your blood (lactic acidosis). Too much lactic acid is a serious but rare medical emergency that can lead to death. Tell your healthcare provider right away if you get these symptoms: weakness or being more tired than usual, unusual muscle pain, being short of breath or fast breathing, stomach pain with nausea and vomiting, cold or blue hands and feet, feel dizzy or lightheaded, or a fast or abnormal heartbeat.

Severe liver problems. In rare cases, severe liver problems that can lead to death have happened in some people who take medicines like TYZEKA. Tell your healthcare provider right away if you get these symptoms: skin or the white part of your eyes turns yellow, dark “tea-colored” urine, light-colored stools, loss of appetite for several days or longer, nausea, or stomach-area pain.

Muscle problems (myopathy). TYZEKA can cause muscle problems, including unexplained muscle pain, muscle weakness or muscle tenderness. Serious muscle problems can happen, including muscle breakdown (rhabdomyolysis). Muscle breakdown can cause kidney damage that can lead to death. Tell your healthcare provider right away if you have unexplained muscle aches, pain, tenderness, or weakness.

Nerve problems. People who take TYZEKA alone or with any type of injectable interferon product can have nerve problems such as numbness, tingling, burning sensations in the arms or legs (peripheral neuropathy), or problems walking. Call your healthcare provider right away if you have any of these symptoms. Your healthcare provider may delay or stop your treatment with TYZEKA if you have any signs or symptoms of nerve problems.

What is TYZEKA?

TYZEKA is a prescription medicine used for the treatment of long-term (chronic) infection with hepatitis B virus (HBV) in people 16 years of age and older who also have active liver damage.

TYZEKA will not cure HBV. TYZEKA may lower the amount of HBV in the body. TYZEKA may improve the condition of your liver.

TYZEKA may lower the ability of HBV to multiply and infect new liver cells. It is not known if TYZEKA is safe and effective in children younger than age 16.
Who should not take TYZEKA?

Do not take TYZEKA if you also receive Pegasys® (peginterferon alfa-2a). Taking TYZEKA with peginterferon alfa-2a may increase your risk of nerve problems.

Before taking TYZEKA, tell your healthcare provider about all of your medical conditions, including if you:

- have HIV-1 (human immunodeficiency virus type 1), hepatitis C or hepatitis D (delta) virus infection
- have had a liver transplant
- have kidney problems
- are receiving hemodialysis
- are pregnant or plan to become pregnant. It is not known if TYZEKA will harm your unborn baby. It is not known if TYZEKA helps prevent a pregnant mother from passing hepatitis B virus (HBV) to her baby. Tell your healthcare provider if you become pregnant during treatment with TYZEKA.
  - Pregnancy Registry: There is a pregnancy registry for women who take antiviral medicines during pregnancy. The purpose of the registry is to collect information about the health of you and your baby. Talk to your healthcare provider about how you can take part in this registry.
- are breastfeeding or plan to breastfeed. It is not known if TYZEKA can pass into your breast milk or if it can harm your baby.

Tell your healthcare provider about all the medicines you take including prescription and over-the-counter medicines, vitamins, and herbal supplements.

TYZEKA may interact with other medicines. Especially tell your healthcare provider or pharmacist if you are also being treated with Pegasys® (pegylated interferon alfa-2a), or any type of injectable interferon product for chronic hepatitis B or C. See "Who should not take TYZEKA?"

TYZEKA oral solution contains salt (sodium). Tell your healthcare provider if you are on a low sodium diet.

How should I take TYZEKA?

- See the detailed Instructions for Use for information about how to take a dose of TYZEKA oral solution. Take TYZEKA exactly as your healthcare provider tells you to.
- Take TYZEKA 1 time each day with or without food.
- Do not change your dose or stop taking TYZEKA without talking to your healthcare provider first.
- Tell your healthcare provider if you have trouble swallowing tablets. TYZEKA also comes as a liquid (oral solution).
- If you miss a dose of TYZEKA, take it as soon as you remember and then take your next dose at the regular time. If it is almost time for your next dose, skip the missed dose.
- If you take more than your prescribed dose of TYZEKA, call your healthcare provider right away.
- It is important to stay under your healthcare provider’s care during treatment with TYZEKA. Your healthcare provider will regularly test the level of the hepatitis B virus in your blood.

What are the possible side effects of TYZEKA?

TYZEKA can cause serious side effects. See "What is the most important information I should know about TYZEKA?"

The most common side effects of TYZEKA include:

- back pain
- dizziness
- muscle aches
- upset stomach
- trouble sleeping
- stomach area (abdominal) swelling
- certain abnormal blood tests

These are not all the possible side effects of TYZEKA.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

How should I store TYZEKA?

- Store TYZEKA tablets and TYZEKA oral solution in their original container at room temperature between...
68°F to 77°F (20°C to 25°C).
- Do not freeze TYZEKA oral solution.
- Use TYZEKA oral solution within 2 months after opening the bottle.
- Keep the bottle closed tightly.
- Throw away TYZEKA when it is outdated or no longer needed by taking TYZEKA to a community take-back disposal program, if available, or by placing TYZEKA in a closed container (such as a sealed bag) in the household trash. Remove all identifying information from the original TYZEKA container before throwing it out.

Keep TYZEKA and all medicines out of the reach of children.

<table>
<thead>
<tr>
<th>General information about the safe and effective use of TYZEKA.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use TYZEKA for a condition for which it was not prescribed. Do not give TYZEKA to other people, even if they have the same symptoms you have. It may harm them. You can ask your healthcare provider or pharmacist for information about TYZEKA that is written for health professionals.</td>
</tr>
</tbody>
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<table>
<thead>
<tr>
<th>What are the ingredients in TYZEKA tablets and TYZEKA oral solution?</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Active ingredient:</strong> telbivudine</td>
</tr>
<tr>
<td><strong>Inactive ingredients in TYZEKA tablets:</strong> colloidal silicon dioxide, magnesium stearate, microcrystalline cellulose, povidone and sodium starch glycolate. The tablet coating contains titanium dioxide, polyethylene glycol, talc, and hypromellose.</td>
</tr>
<tr>
<td><strong>Inactive ingredients in TYZEKA oral solution:</strong> citric acid anhydrous, benzoic acid, passion fruit flavor, sodium saccharin, sodium hydroxide, and purified water. A 600 mg dose (30 mL) of TYZEKA oral solution contains about 47 mg of sodium.</td>
</tr>
</tbody>
</table>

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For more information call 1-877-889-9352 or go to www.TYZEKA.com.

This Medication Guide has been approved by the U.S. Food and Drug Administration

Revised: December 2018
Instructions for Use

Make sure that you read, understand, and follow these instructions carefully so that you take TYZEKA oral solution the right way.

1. Supplies needed to take a dose of TYZEKA oral solution

To take your dose of TYZEKA you will need:
1. Bottle containing the medicine.
2. Child-resistant cap.
3. Oral dosing cup with 6, 10, 20, and 30 mL markings.

See Figure 1.

2. Important Information

- Do not pour any extra medicine back into the bottle. This will help prevent contamination.
- Close the bottle right away after use.
- Clean the dosing cup with clean water after each use.
- Keep the dosing cup dry between uses.
- The dosing cup is intended for measurement of TYZEKA oral solution only.

3. Prepare a dose of TYZEKA oral solution using the dosing cup

1. Remove the dosing cup from the cap.
2. Press down and turn the child-resistant cap at the same time, to open the bottle. See Figure 2a and 2b.
3. Before you pour the solution, check for the marking on the dosing cup that matches the amount of medicine that has been prescribed.

4. Hold the dosing cup at eye level. This will help to make sure that you measure the right amount of medicine. Carefully and slowly pour the prescribed amount of solution from the bottle into the dosing cup, until the solution reaches the line that matches the correct line on the dosing cup. See Figure 3.

**If you pour too much medicine into the cup, pour the extra medicine into the sink. Do not pour it back into the bottle.**

5. Take the medicine right away. See Figure 4.

6. Close the bottle tightly.

7. Rinse the dosing cup with clean water.

8. Dry the dosing cup using a dry, clean tissue before you put it back onto the cap. See Figure 5.