PANCREAZE™ (pancrelipase) delayed-release capsules

These highlights do not include all the information needed to use PANCREAZE™ safely and effectively. See full prescribing information for PANCREAZE™.

PANCREAZE™ is a combination of porcine-derived lipases, proteases, and amylases indicated for the treatment of exocrine pancreatic insufficiency due to cystic fibrosis or other conditions (1).

INDICATIONS AND USAGE

Initial U.S. Approval – XXXX

Dosage

PANCREAZE™ is not interchangeable with any other pancrelipase product.

Children Older than 12 Months and Younger than 4 Years

• Enzyme dosing should begin with 1,000 lipase units/kg of body weight per meal to a maximum of 2,500 lipase units/kg of body weight per meal (or less than or equal to 10,000 lipase units/kg of body weight per day), or less than 4,000 lipase units/g fat ingested per day. (2.1)

Children 4 Years and Older and Adults

• Enzyme dosing should begin with 500 lipase units/kg of body weight per meal to a maximum of 2,500 lipase units/kg of body weight per meal (or less than or equal to 10,000 lipase units/kg of body weight per day), or less than 4,000 lipase units/g fat ingested per day. (2.1)

Limitations on Dosing

• Dosing should not exceed the recommended maximum dosage set forth by the Cystic Fibrosis Foundation Consensus Conferences Guidelines. (2.1)

Administration

• PANCREAZE should be swallowed whole. For infants or patients unable to swallow intact capsules, the contents may be sprinkled on soft acidic food with a pH of 4.5 or less, e.g., applesauce. (2.2)

Dosage Forms and Strengths

• Capsules: 4,200 USP units of lipase; 10,000 USP units of protease; 17,500 USP units of amylase. Capsules have a yellow opaque body and clear cap, printed with “McNEIL” and “MT 4” (3)

• Capsules: 10,500 USP units of lipase; 25,000 USP units of protease; 43,750 USP units of amylase. Capsules have a pink opaque body and clear cap, printed with “McNEIL” and “MT 10” (3)

• Capsules: 16,800 USP units of lipase; 40,000 USP units of protease; 70,000 USP units of amylase. Capsules have a salmon opaque body and clear cap, printed with “McNEIL” and “MT 16” (3)

• Capsules: 21,000 USP units of lipase; 37,000 USP units of protease; 61,000 USP units of amylase. Capsules have a white opaque body and cap, printed with “McNEIL” and “MT 20” (3)

WARNINGS AND PRECAUTIONS

None (4)

ADVERSE REACTIONS

• Fibrosing colonopathy is associated with high-dose use of pancreatic enzyme replacement. Exercise caution when doses of PANCREAZE exceed 2,500 lipase units/kg of body weight per meal (or greater than 10,000 lipase units/kg of body weight per day). (5.1)

• To avoid irritation of oral mucosa, do not chew PANCREAZE or retain in the mouth. (5.2)

• Exercise caution when prescribing PANCREAZE to patients with gout, renal impairment, or hyperuricemia. (5.3)

• There is theoretical risk of viral transmission with all pancreatic enzyme products including PANCREAZE. (5.4)

• Exercise caution when administering pancrelipase to a patient with a known allergy to proteins of porcine origin. (5.5)

ADVERSE REACTIONS

• Treatment emergent adverse events occurring in at least 2 patients (greater than or equal to 10%) receiving PANCREAZE or placebo are abdominal pain, abdominal pain upper, flatulence, diarrhea, abnormal feces, and fatigue. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact McNeil Pediatrics at 1-800-525-7736 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

USE IN SPECIFIC POPULATIONS

Pediatric Patients

• The safety and effectiveness of PANCREAZE were assessed in pediatric patients, aged 6 to 30 months old and aged 8 to 17 years old. (8.4)

• The safety and efficacy of pancreatic enzyme products with different formulations of pancrelipase in pediatric patients have been described in the medical literature and through clinical experience. (8.4)

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide.

Revised: April 2010

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FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE
PANCREAZE (pancrelipase) is indicated for the treatment of exocrine pancreatic insufficiency due to cystic fibrosis or other conditions.

2 DOSAGE AND ADMINISTRATION

2.1 Dosage
PANCREAZE is not interchangeable with other pancrelipase products.

PANCREAZE is orally administered. Therapy should be initiated at the lowest recommended dose and gradually increased. The dosage of PANCREAZE should be individualized based on clinical symptoms, the degree of steatorrhea present, and the fat content of the diet (see Limitations on Dosing below).

Dosage recommendations for pancreatic enzyme replacement therapy were published following the Cystic Fibrosis Foundation Consensus Conferences.\(^1\)\(^2\)\(^3\) PANCREAZE should be administered in a manner consistent with the recommendations of the Conferences provided in the following paragraphs. Patients may be dosed on a fat ingestion-based or actual body weight-based dosing scheme.

Infants (up to 12 months)
Infants may be given 2,000 to 4,000 lipase units per 120 mL of formula or per breast-feeding. Do not mix PANCREAZE capsule contents directly into formula or breast milk prior to administration [see Dosage and Administration (2.2)].

Children Older than 12 Months and Younger than 4 Years
Enzyme dosing should begin with 1,000 lipase units/kg of body weight per meal for children less than age 4 years to a maximum of 2,500 lipase units/kg of body weight per meal (or less than or equal to 10,000 lipase units/kg of body weight per day), or less than 4,000 lipase units/g fat ingested per day.

Children 4 Years and Older and Adults
Enzyme dosing should begin with 500 lipase units/kg of body weight per meal for those older than age 4 years to a maximum of 2,500 lipase units/kg of body weight per meal (or less than or equal to 10,000 lipase units/kg of body weight per day), or less than 4,000 lipase units/g fat ingested per day.
Usually, half of the prescribed PANCREAZE dose for an individualized full meal should be given with each snack. The total daily dose should reflect approximately three meals plus two or three snacks per day.

Enzyme doses expressed as lipase units/kg of body weight per meal should be decreased in older patients because they weigh more but tend to ingest less fat per kilogram of body weight.

**Limitations on Dosing**

Dosing should not exceed the recommended maximum dosage set forth by the Cystic Fibrosis Foundation Consensus Conferences Guidelines.1,2,3 If symptoms and signs of steatorrhea persist, the dosage may be increased by a healthcare professional. Patients should be instructed not to increase the dosage on their own. There is great inter-individual variation in response to enzymes; thus, a range of doses is recommended. Changes in dosage may require an adjustment period of several days. If doses are to exceed 2,500 lipase units/kg of body weight per meal, further investigation is warranted.

Doses greater than 2,500 lipase units/kg of body weight per meal (or greater than 10,000 lipase units/kg of body weight per day) should be used with caution and only if they are documented to be effective by 3-day fecal fat measures that indicate a significantly improved coefficient of fat absorption. Doses greater than 6,000 lipase units/kg of body weight per meal have been associated with colonic strictures, indicative of fibrosing colonopathy, in children with cystic fibrosis less than 12 years of age [see Warnings and Precautions (5.1)]. Patients currently receiving higher doses than 6,000 lipase units/kg of body weight per meal should be examined and the dosage either immediately decreased or titrated downward to a lower range.

### 2.2 Administration

**PANCREAZE** should always be taken as prescribed by a healthcare professional.

**Infants (up to 12 months)**

PANCREAZE should be administered to infants immediately prior to each feeding, using a dosage of 2,000 to 4,000 lipase units per 120 mL of formula or per breast-feeding. Contents of the capsule may be sprinkled on small amounts of acidic soft food with a pH of 4.5 or less (e.g., applesauce) and given to the infant within 15 minutes. Contents of the capsule may also be administered directly to the mouth. Administration should be followed by breast milk or formula. Contents of the capsule **should not** be mixed directly
into formula or breast milk as this may diminish efficacy. Care should be taken to ensure that PANCREAZE is not crushed or chewed or retained in the mouth, to avoid irritation of the oral mucosa.

*Children and Adults*

PANCREAZE should be taken during meals or snacks, with sufficient fluid. **PANCREAZE capsules and capsule contents should not be crushed or chewed.** Capsules should be swallowed whole.

For patients who are unable to swallow intact capsules, the capsules may be carefully opened and the contents sprinkled on small amounts of acidic soft food with a pH of 4.5 or less (e.g., applesauce). The PANCREAZE-soft food mixture should be swallowed immediately without crushing or chewing, and followed with water or juice to ensure complete ingestion. Care should be taken to ensure that no drug is retained in the mouth.

### 3 DOSAGE FORMS AND STRENGTHS

The active ingredient in PANCREAZE evaluated in clinical trials is lipase. PANCREAZE is dosed by lipase units.

PANCREAZE is available in 4 color coded capsule strengths.

Other active ingredients include protease and amylase. Each PANCREAZE capsule strength contains the specified amounts of lipase, protease, and amylase as follows:

- 4,200 USP units of lipase; 10,000 USP units of protease; 17,500 USP units of amylase capsules have a yellow opaque body and clear cap, printed with “McNEIL” and “MT 4”
- 10,500 USP units of lipase; 25,000 USP units of protease; 43,750 USP units of amylase capsules have a pink opaque body and clear cap, printed with “McNEIL” and “MT 10”
- 16,800 USP units of lipase; 40,000 USP units of protease; 70,000 USP units of amylase capsules have a salmon opaque body and clear cap, printed with “McNEIL” and “MT 16”
- 21,000 USP units of lipase; 37,000 USP units of protease; 61,000 USP units of amylase capsules have a white opaque body and cap, printed with “McNEIL” and “MT 20”

### 4 CONTRAINDICATIONS

None.
5 WARNINGS AND PRECAUTIONS

5.1 Fibrosing Colonopathy

Fibrosing colonopathy has been reported following treatment with different pancreatic enzyme products. Fibrosing colonopathy is a rare serious adverse reaction initially described in association with high-dose pancreatic enzyme use, usually with use over a prolonged period of time and most commonly reported in pediatric patients with cystic fibrosis. The underlying mechanism of fibrosing colonopathy remains unknown. Doses of pancreatic enzyme products exceeding 6,000 lipase units/kg of body weight per meal have been associated with colonic strictures in children less than 12 years of age. Patients with fibrosing colonopathy should be closely monitored because some patients may be at risk of progressing to stricture formation. It is uncertain whether regression of fibrosing colonopathy occurs. It is generally recommended, unless clinically indicated, that enzyme doses should be less than 2,500 lipase units/kg of body weight per meal (or less than 10,000 lipase units/kg of body weight per day) or less than 4,000 lipase units/g fat ingested per day [see Dosage and Administration (2.1)].

Doses greater than 2,500 lipase units/kg of body weight per meal (or greater than 10,000 lipase units/kg of body weight per day) should be used with caution and only if they are documented to be effective by 3-day fecal fat measures that indicate a significantly improved coefficient of fat absorption. Patients receiving higher doses than 6,000 lipase units/kg of body weight per meal should be examined and the dosage either immediately decreased or titrated downward to a lower range.

5.2 Potential for Irritation to Oral Mucosa

Care should be taken to ensure that no drug is retained in the mouth. PANCREAZE should not be crushed or chewed or mixed in foods having a pH greater than 4.5. These actions can disrupt the protective enteric coating resulting in early release of enzymes, irritation of oral mucosa, and/or loss of enzyme activity [see Dosage and Administration (2.2) and Patient Counseling Information (17.4)]. For patients who are unable to swallow intact capsules, the capsules may be carefully opened and the contents sprinkled to a small amount of acidic soft food with a pH of 4.5 or less, such as applesauce. The PANCREAZE-soft food mixture should be swallowed immediately and followed with water or juice to ensure complete ingestion.

5.3 Potential for Risk of Hyperuricemia

Caution should be exercised when prescribing PANCREAZE to patients with gout, renal impairment, or hyperuricemia. Porcine-derived pancreatic enzyme products contain purines that may increase blood uric acid levels.
5.4 Potential Viral Exposure from the Product Source

PANCREAZE is sourced from pancreatic tissue from swine used for food consumption. Although the risk that PANCREAZE will transmit an infectious agent to humans has been reduced by testing for certain viruses during manufacturing and by inactivating certain viruses during manufacturing, there is a theoretical risk for transmission of viral disease, including diseases caused by novel or unidentified viruses. Thus, the presence of porcine viruses that might infect humans cannot be definitely excluded. However, no cases of transmission of an infectious illness associated with the use of porcine pancreatic extracts have been reported.

5.5 Allergic Reactions

Caution should be exercised when administering pancrelipase to a patient with a known allergy to proteins of porcine origin. Rarely, severe allergic reactions including anaphylaxis, asthma, hives, and pruritus, have been reported with other pancreatic enzyme products with different formulations of the same active ingredient (pancrelipase). The risks and benefits of continued PANCREAZE treatment in patients with severe allergy should be taken into consideration with the overall clinical needs of the patient.

6 ADVERSE REACTIONS

The most serious adverse reactions reported with different pancreatic enzyme products of the same active ingredient (pancrelipase) include fibrosing colonopathy, hyperuricemia and allergic reactions [see Warnings and Precautions, (5)]

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.

The short-term safety of PANCREAZE was assessed in two clinical trials conducted in 57 patients with exocrine pancreatic insufficiency (EPI) due to CF. Study 1 was conducted in 40 patients, ages 8 years to 57 years; Study 2 was conducted in 17 patients, ages 6 to 30 months. In Study 1, PANCREAZE was administered in a dose of approximately 6,300 lipase units per kilogram per day for lengths of treatment ranging from 8 to 26 days; in Study 2, PANCREAZE was administered in four treatment arms (doses of 1,375, 2,875, 4,735, and 5,938 lipase units per kilogram per day) for lengths of treatment ranging from 6 to 11 days. The population was nearly evenly distributed in gender, and approximately 96% of patients were Caucasian.
Study 1 was a randomized, double-blind, placebo-controlled, study of 40 patients, ages 8 to 57 years, with EPI due to CF. In this study, patients received PANCREAZE at individually titrated doses (not to exceed 2,500 lipase units per kilogram per meal) for 14 days, followed by randomization to PANCREAZE or matching placebo for 7 days of treatment. The mean exposure to PANCREAZE during this study, including titration period and randomized withdrawal period, was 18 days.

The incidence of adverse events (regardless of causality) was higher during placebo treatment (60%) than during PANCREAZE treatment (40%). The most common adverse events reported during the study were gastrointestinal complaints, which were reported more commonly during placebo treatment (55%) than during PANCREAZE treatment (30%). The type and incidence of adverse events were similar in children (8 to 11 years), adolescents (12 to 17 years), and adults (greater than 18 years).

Table 1 enumerates treatment-emergent adverse events that occurred in at least 2 patients (greater than or equal to 10%) treated with either PANCREAZE or placebo in Study 1. Adverse events were classified by Medical Dictionary for Regulatory Activities (MedDRA) terminology.

Table 1: Treatment-Emergent Adverse Events Occurring in at least 2 Patients (greater than or equal to 10%) in Either Treatment Group of the Placebo-Controlled, Clinical Study of PANCREAZE

<table>
<thead>
<tr>
<th>MedDRA Primary System Organ Class Preferred Term</th>
<th>PANCREAZE (N=20) n (%)</th>
<th>Placebo (N=20) n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gastrointestinal Disorders</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>2 (10%)</td>
<td>3 (15%)</td>
</tr>
<tr>
<td>Abdominal pain upper</td>
<td>1 (5%)</td>
<td>3 (15%)</td>
</tr>
<tr>
<td>Flatulence</td>
<td>1 (5%)</td>
<td>3 (15%)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>0 (0%)</td>
<td>4 (20%)</td>
</tr>
<tr>
<td>Abnormal feces</td>
<td>0 (0%)</td>
<td>3 (15%)</td>
</tr>
<tr>
<td><strong>General Disorders And Administration Site Conditions</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatigue</td>
<td>0 (0%)</td>
<td>2 (10%)</td>
</tr>
</tbody>
</table>

Study 2 was a randomized, investigator-blinded, dose-ranging study of 17 patients, ages 6 to 30 months, with EPI due to CF. All patients were transitioned from their usual PEP treatment to PANCREAZE at 375 lipase units per kilogram body weight per meal for a 6 day run-in period. Patients were then randomized to receive PANCREAZE at one of four doses (375, 750, 1,125, and 1,500 lipase units per kilogram body weight per meal) for 5 days. Adverse events were collected on patient diary entries and at each study visit.
The most commonly reported adverse events were gastrointestinal, including diarrhea and vomiting, and were similar in type and frequency across treatment arms and to those reported in the double-blind, placebo-controlled trial (Study 1).

6.2 Postmarketing Experience
Post-marketing data for PANCREAZE has been available since 1988. The safety data is similar to that described below.

Delayed- and immediate-release pancreatic enzyme products with different formulations of the same active ingredient (pancrelipase) have been used for the treatment of patients with exocrine pancreatic insufficiency due to cystic fibrosis and other conditions, such as chronic pancreatitis. The long-term safety profile of these products has been described in the medical literature. The most serious adverse events included fibrosing colonopathy, distal intestinal obstruction syndrome (DIOS), recurrence of pre-existing carcinoma, and severe allergic reactions including anaphylaxis, asthma, hives, and pruritus. The most commonly reported adverse events were gastrointestinal disorders, including abdominal pain, diarrhea, flatulence, constipation and nausea, and skin disorders including pruritus, urticaria and rash. In general, these products have a well defined and favorable risk-benefit profile in exocrine pancreatic insufficiency.

Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

7 DRUG INTERACTIONS
No drug interactions have been identified. No formal interaction studies have been conducted.

8 USE IN SPECIFIC POPULATIONS
  8.1 Pregnancy
Teratogenic effects

Pregnancy Category C: Animal reproduction studies have not been conducted with pancrelipase. It is not known whether pancrelipase can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. PANCREAZE should be given to a pregnant woman only if clearly needed. The risk and benefit of pancrelipase should be considered in the context of the need to provide adequate nutritional support to a pregnant woman with exocrine pancreatic insufficiency. Adequate caloric intake during pregnancy is important for normal maternal weight gain and fetal
growth. Reduced maternal weight gain and malnutrition can be associated with adverse pregnancy outcomes.

8.3 Nursing Mothers
It is not known whether this drug is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised when PANCREAZE is administered to a nursing woman. The risk and benefit of pancrelipase should be considered in the context of the need to provide adequate nutritional support to a nursing mother with exocrine pancreatic insufficiency.

8.4 Pediatric Use
The short-term safety and effectiveness of PANCREAZE were assessed in two clinical studies in pediatric patients with EPI due to CF; one study included patients ages 6 to 30 months, and the other included patients ages 8 years to 17 years.

Study 1 was a randomized, double-blind, placebo-controlled study in 40 patients, 14 of whom were pediatric patients, including 7 children aged 8 to 11 years, and 7 adolescents aged 12 to 17 years. The safety and efficacy in pediatric patients in this study were similar to adult patients [see Adverse Reactions (6.1) and Clinical Studies (14)].

Study 2 was a randomized, investigator-blinded, dose-ranging study in 17 pediatric patients aged 6 to 30 months. When patient regimen was switched from their usual PEP regimen to PANCREAZE, patients showed similar control of their fat malabsorption [see Adverse Reactions (6.1) and Clinical Studies (14)].

The safety and efficacy of pancreatic enzyme products with different formulations of pancrelipase consisting of the same active ingredient (lipases, proteases, and amylases) for treatment of children with exocrine pancreatic insufficiency due to cystic fibrosis has been described in the medical literature and through clinical experience.

Dosing of pediatric patients should be in accordance with recommended guidance from the Cystic Fibrosis Foundation Consensus Conferences [see Dosage and Administration (2.1)]. Doses of other pancreatic enzyme products exceeding 6,000 lipase units/kg of body weight per meal have been associated with fibrosing colonopathy and colonic strictures in children less than 12 years of age [see Warnings and Precautions (5.1)].

10 OVERDOSAGE
In Study 1, a 10 year-old patient was administered a PANCREAZE dose of 12,399 lipase units per kilogram per day for the duration of the open-label and randomized withdrawal periods. The patient experienced mild abdominal pain throughout both study periods.
Abnormal chemistry data at the end of the study included mild elevations of aspartate aminotransferase (AST), alanine aminotransferase (ALT), and serum phosphate. Abnormal hematology data at the end of the study included mild elevations of hematocrit. No abnormalities from analyses of urinalysis or uric acid were noted.

Chronic high doses of pancreatic enzyme products have been associated with fibrosing colonopathy and colonic strictures [see Dosage and Administration (2.1) and Warnings and Precautions (5.1)]. High doses of pancreatic enzyme products have been associated with hyperuricosuria and hyperuricemia, and should be used with caution in patients with a history of hyperuricemia, gout, or renal impairment [see Warnings and Precautions (5.3)].

11 DESCRIPTION
PANCREAZE is a pancreatic enzyme preparation consisting of pancrelipase, an extract derived from porcine pancreatic glands. Pancrelipase contains multiple enzyme classes, including porcine-derived lipases, proteases, and amylases.

Each capsule for oral administration contains enteric-coated microtablets that are each approximately 2 mm in diameter.

The active ingredient evaluated in clinical trials is lipase. PANCREAZE is dosed by lipase units. Other active ingredients include protease and amylase.

Inactive ingredients in PANCREAZE include cellulose, colloidal anhydrous silica, crospovidone, magnesium stearate, methacrylic acid ethyl acrylate copolymer, montan glycol wax, simethicone emulsion, talc and triethyl citrate.

PANCREAZE is available in four color coded strengths. Each PANCREAZE capsule strength contains the specified amounts of lipase, protease, and amylase as follows:

**4,200 USP units of lipase;** 10,000 USP units of protease; 17,500 USP units of amylase. The hard gelatin capsules have a yellow opaque body and clear cap imprinted with “McNEIL” and “MT 4”. The capsule shell contains gelatin, titanium dioxide, sodium lauryl sulfate, sorbitan monolaurate, iron oxide, and gelatin capsule imprint ink.

**10,500 USP units of lipase;** 25,000 USP units of protease; 43,750 USP units of amylase. The hard gelatin capsules have a pink opaque body and clear cap imprinted with “McNEIL” and “MT 10”. The capsule shell contains gelatin, titanium dioxide, sodium lauryl sulfate, sorbitan monolaurate, iron oxide, and gelatin capsule imprint ink.
16,800 USP units of lipase; 40,000 units of protease; 70,000 USP units of amylase. The hard gelatin capsules have a salmon opaque body and clear cap imprinted with “McNEIL” and “MT 16”. The capsule shell contains gelatin, titanium dioxide, sodium lauryl sulfate, sorbitan monolaurate, iron oxide, and gelatin capsule imprint ink.

21,000 USP units of lipase; 37,000 units of protease; 61,000 USP units of amylase. The hard gelatin capsules have a white opaque body and cap imprinted with “McNEIL” and “MT 20”. The capsule shell contains gelatin, titanium dioxide, sodium lauryl sulfate, sorbitan monolaurate, and gelatin capsule imprint ink.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action
The pancreatic enzymes in PANCREAZE catalyze the hydrolysis of fats to monoglyceride, glycerol and free fatty acids, proteins into peptides and amino acids, and starches into dextrins and short chain sugars such as maltose and maltroiose in the duodenum and proximal small intestine, thereby acting like digestive enzymes physiologically secreted by the pancreas.

12.3 Pharmacokinetics
The pancreatic enzymes in PANCREAZE are enteric-coated to minimize destruction or inactivation in gastric acid. PANCREAZE is expected to release most of the enzymes in vivo at pH greater than 5.5. Pancreatic enzymes are not absorbed from the gastrointestinal tract in appreciable amounts.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
Carcinogenicity, genetic toxicology, and animal fertility studies have not been performed with pancrelipase.

14 CLINICAL STUDIES
The short-term safety and efficacy of PANCREAZE were evaluated in two studies conducted in 57 patients with exocrine pancreatic insufficiency (EPI) associated with cystic fibrosis (CF).

Study 1 was a randomized, double-blind, placebo-controlled study of 40 patients, ages 8 to 57 years, with EPI due to CF. In this study, patients received PANCREAZE at individually titrated doses (not to exceed 2,500 lipase units per kilogram per meal) for 14 days (open label period) followed by randomization to PANCREAZE or matching placebo for 7 days of treatment (double-blind withdrawal period). Only patients with coefficient of fat absorption (CFA) ≥80% in the open label period were randomized to the
double-blind withdrawal period. The mean dose during the controlled treatment period was 6,400 lipase units per kilogram per day. All patients consumed a high-fat diet (greater than or equal to 100 grams of fat per day) during the treatment period.

The primary efficacy endpoint was the change in CFA from the open label period to the end of the double-blind withdrawal period. The CFA was determined by a 72-hour stool collection period during both treatment periods, when both fat excretion and fat ingestion were measured (Table 2).

### Table 2. Change in CFA in Study 1 (Open Label Period to End of Double-Blind Withdrawal Period)

<table>
<thead>
<tr>
<th></th>
<th>PANCREAZE n=20</th>
<th>Placebo n=20</th>
</tr>
</thead>
<tbody>
<tr>
<td>CFA [%]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Open Label Period* (Mean, SD)</td>
<td>88 (5)</td>
<td>91 (5)</td>
</tr>
<tr>
<td>End of Double-Blind Withdrawal Period† (Mean, SD)</td>
<td>87 (8)</td>
<td>56 (25)</td>
</tr>
<tr>
<td>Change in CFA† [%]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Open Label Period to End of Double-Blind Withdrawal Period (Mean, SD)</td>
<td>-2 (6)</td>
<td>-34 (23)</td>
</tr>
<tr>
<td>Treatment Difference Point Estimate (95% CI)</td>
<td>33 (25, 40)</td>
<td></td>
</tr>
</tbody>
</table>

*Minimum of 72 hours from start of open label period.
†Double-blind withdrawal period ranged from 4 to 7 days.
†p<0.001

At the end of the double-blind withdrawal period, the mean change in CFA from the open label period to the end of the double-blind withdrawal period was -2% with PANCREAZE treatment compared to -34% with placebo treatment. There were similar responses to PANCREAZE by age and gender.

Study 2 was a randomized, investigator-blinded, dose-ranging study of 17 patients, ages 6 months to 30 months (mean 18 months) with EPI due to CF. The final analysis population was limited to 16 patients; 1 patient was excluded due to withdrawal of consent. All patients were transitioned from their usual PEP treatment to PANCREAZE at 375 lipase units per kilogram body weight per meal for a 6 day run-in period. Patients were then randomized to receive PANCREAZE at one of four doses (375, 750, 1,125, and 1,500 lipase units per kilogram body weight per meal) for 5 days. The CFA was measured at the end of the run-in period and at the end of the randomized period (Table 3).
Table 3. Change in CFA in Study 2 (End of Run-in Period to End of Study)

<table>
<thead>
<tr>
<th>CFA (%)</th>
<th>375 units lipase/kg/meal n=4</th>
<th>750 units lipase/kg/meal n=4</th>
<th>1,125 units lipase/kg/meal n=4</th>
<th>1,500 units lipase/kg/meal n=4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 6* (Mean, SD)</td>
<td>93 (2)</td>
<td>90 (5)</td>
<td>81 (11)</td>
<td>93 (3)</td>
</tr>
<tr>
<td>Day 11# (Mean, SD)</td>
<td>92 (3)</td>
<td>91 (4)</td>
<td>80 (13)</td>
<td>91 (2)</td>
</tr>
<tr>
<td>Change in CFA (%) Day 6 to Day 11 (Mean, SD)</td>
<td>-2 (3)</td>
<td>1 (3)</td>
<td>-1 (3)</td>
<td>-2 (3)</td>
</tr>
</tbody>
</table>

*End of Run-in Period; #End of Study

Overall, patients showed similar CFA at the end of the run-in period (mean PANCREAZE dose of 1,600 lipase units per kilogram body weight per day) as at the end of the study across the four treatment arms.

15 REFERENCES


16 HOW SUPPLIED/STORAGE AND HANDLING

PANCREAZE (pancrelipase) Delayed-Release Capsules

4,200 USP units of lipase; 10,000 USP units of protease); 17,500 USP units of amylase.

PANCREAZE (pancrelipase) is supplied as hard gelatin capsules with a yellow opaque body and clear cap imprinted with “McNEIL” and “MT 4” and packaged in bottles of 100–(NDC 50458-341-60).

PANCREAZE (pancrelipase) Delayed-Release Capsules
10,500 USP units of lipase; 25,000 USP units of protease; 43,750 USP units of amylase.

PANCREAZE (pancrelipase) is supplied as hard gelatin capsules with a pink opaque body and clear cap imprinted with “McNEIL” and “MT 10” and packaged in bottles of 100–(NDC 50458-342-60).

PANCREAZE (pancrelipase) Delayed-Release Capsules
16,800 USP units of lipase; 40,000 USP units of protease; 70,000 USP units of amylase.

PANCREAZE (pancrelipase) is supplied as hard gelatin capsules with a salmon opaque body and clear cap imprinted with “McNEIL” and “MT 16” and packaged in bottles of 100–(NDC 50458-343-60).

PANCREAZE (pancrelipase) Delayed-Release Capsules
21,000 USP units of lipase; 37,000 USP units of protease; 61,000 USP units of amylase.

PANCREAZE (pancrelipase) is supplied as hard gelatin capsules with a white opaque body and cap imprinted with “McNEIL” and “MT 20” and packaged in bottles of 100–(NDC 50458-346-60).

Storage and Handling
Avoid heat. PANCREAZE hard gelatin capsules should be stored in a dry place in the original container. After opening, KEEP THE CONTAINER TIGHTLY CLOSED between uses to PROTECT FROM MOISTURE. Do not store above 25°C (77°F).

The PANCREAZE 4200 USP Units of lipase bottle contains a desiccant packet. DO NOT eat or throw away the packet (desiccant) in your medicine bottle. This packet will protect your medicine from moisture.

Keep out of reach of children.

DO NOT CRUSH PANCREAZE delayed-release capsules or the capsule contents.

17 PATIENT COUNSELING INFORMATION
See Medication Guide

17.1 Dosing and Administration
- Instruct patients and caregivers that PANCREAZE should only be taken as directed by their healthcare professional. Patients should be advised that the total daily dose should not exceed 10,000 lipase units/kg body weight/day unless clinically indicated. This needs to be especially emphasized for patients eating multiple snacks and meals per day. Patients should be informed that if a dose is
missed, the next dose should be taken with the next meal or snack as directed. Doses should not be doubled [see Dosage and Administration (2)].

- Instruct patients and caregivers that PANCREAZE should always be taken with food. Patients should be advised that PANCREAZE delayed-release capsules and the capsule contents must not be crushed or chewed as doing so could cause early release of enzymes and/or loss of enzymatic activity. Patients should swallow the intact capsules with adequate amounts of liquid at mealtimes. If necessary, the capsules contents can also be sprinkled on soft acidic foods. [see Dosage and Administration (2)].

- Instruct patients to notify their healthcare professional if they are pregnant or are thinking of becoming pregnant during treatment with PANCREAZE [see Use in Specific Populations (8.1)].

- Instruct patients to notify their healthcare professional if they are breast feeding or are thinking of breast feeding during treatment with PANCREAZE [see Use in Specific Populations (8.3)].

### 17.2 Fibrosing Colonopathy
Advise patients and caregivers to follow dosing instructions carefully, as doses of pancreatic enzyme products exceeding 6,000 lipase units/kg of body weight per meal (10,000 lipase units/kg of body weight/day) have been associated with colonic strictures in children below the age of 12 years [see Dosage and Administration (2)].

### 17.3 Allergic Reactions
Advise patients and caregivers to contact their healthcare professional immediately if allergic reactions to PANCREAZE develop [see Warnings and Precautions (5.5)].

**Manufactured by:**
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25436 Uetersen, Germany.

**Manufactured for:**
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