HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use PROLASTIN®-C LIQUID safely and effectively. See full prescribing information for PROLASTIN-C LIQUID.

PROLASTIN®-C LIQUID (Alpha1-Proteinase Inhibitor [Human]) Solution for Intravenous Injection
Initial U.S. Approval: 1987

INDICATIONS AND USAGE

PROLASTIN-C LIQUID is an Alpha1-Proteinase Inhibitor (Human) (Alpha1-PI) indicated for chronic augmentation and maintenance therapy in adults with clinical evidence of emphysema due to severe hereditary deficiency of Alpha1-PI (alpha1-antitrypsin deficiency).

Limitations of Use:
- The effect of augmentation therapy with any Alpha1-PI, including PROLASTIN-C LIQUID, on pulmonary exacerbations and on the progression of emphysema in Alpha1-PI deficiency has not been conclusively demonstrated in randomized, controlled clinical trials.
- Clinical data demonstrating the long-term effects of chronic augmentation or maintenance therapy with PROLASTIN-C LIQUID are not available.
- PROLASTIN-C LIQUID is not indicated as therapy for lung disease in patients in whom severe Alpha1-PI deficiency has not been established.

DOSAGE AND ADMINISTRATION

For intravenous use only.

Dose: 60 mg/kg body weight intravenously once per week.
Dose ranging studies using efficacy endpoints have not been performed with any Alpha1-PI product, including PROLASTIN-C LIQUID.
Use a sterile 15 micron in-line filter when administering the product (not supplied).
Administration: 0.08 mL/kg/min as determined by patient response and comfort.

DOSAGE FORMS AND STRENGTHS

For injection: approximately 1,000 mg in a single-use vial containing 20 mL of solution for injection.
CONTRAINDICATIONS

- Immunoglobulin A (IgA) deficient patients with antibodies against IgA. (4)
- History of anaphylaxis or other severe systemic reaction to Alpha1-PI. (4)

WARNINGS AND PRECAUTIONS

- Severe hypersensitivity and anaphylactic reactions may occur in IgA deficient patients with antibodies against IgA. Discontinue administration of the product and initiate appropriate emergency treatment if hypersensitivity reactions occur. (5.1)

- Because PROLASTIN-C LIQUID is made from human plasma, it may carry a risk of transmitting infectious agents, e.g., viruses, the variant Creutzfeldt-Jakob disease (vCJD) agent, and, theoretically, the Creutzfeldt-Jakob disease (CJD) agent. (5.2)

ADVERSE REACTIONS

The most common adverse reactions during PROLASTIN-C LIQUID clinical trials in > 5% of subjects were diarrhea and fatigue, each of which occurred in 2 subjects (6%). (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact Grifols Therapeutics Inc. at 1-800-520-2807 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

See 17 for PATIENT COUNSELING INFORMATION. Revised: 9/2017
13 NON-CLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

13.2 Animal Toxicology and/or Pharmacology

14 CLINICAL STUDIES

15 REFERENCES

16 HOW SUPPLIED/STORAGE AND HANDLING

17 PATIENT COUNSELING INFORMATION

* Sections or subsections omitted from the full prescribing information are not listed.
FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

PROLASTIN-C LIQUID is an Alpha1-Proteinase Inhibitor (Human) (Alpha1-PI) indicated for chronic augmentation and maintenance therapy in adults with clinical evidence of emphysema due to severe hereditary deficiency of Alpha1-PI (alpha1-antitrypsin deficiency).

Limitations of Use

- The effect of augmentation therapy with any Alpha1-PI, including PROLASTIN-C LIQUID, on pulmonary exacerbations and on the progression of emphysema in Alpha1-PI deficiency has not been conclusively demonstrated in randomized, controlled clinical trials.
- Clinical data demonstrating the long-term effects of chronic augmentation or maintenance therapy with PROLASTIN-C LIQUID are not available.
- PROLASTIN-C LIQUID is not indicated as therapy for lung disease in patients in whom severe Alpha1-PI deficiency has not been established.

2 DOSAGE AND ADMINISTRATION

For intravenous use only.

2.1 Dose

- The recommended dose of PROLASTIN-C LIQUID is 60 mg/kg body weight administered intravenously once weekly.
- Dose ranging studies using efficacy endpoints have not been performed with any Alpha1-PI product.
- The carton and the label on each vial of PROLASTIN-C LIQUID show the actual amount of functionally active Alpha1-PI in milligrams (as determined by the capacity to neutralize porcine pancreatic elastase).

2.2 Preparation and Handling

1. Allow unopened PROLASTIN-C LIQUID to warm up to room temperature before administration.
2. Remove the plastic flip top from the vial.
3. Swab the exposed stopper surface with alcohol and allow to dry.
4. Inspect the PROLASTIN-C LIQUID visually for particulate matter and discoloration prior to pooling. The product may contain a few protein particles. The solution is clear, colorless or pale yellow or pale green. Do not use if the product is discolored or cloudy.
5. Pool PROLASTIN-C LIQUID from several vials to achieve the intended mg/kg body weight dose into an empty, sterile intravenous solution container using aseptic technique.

6. Keep pooled solution at room temperature for administration within three hours.

2.3 Administration

- Visually inspect parenteral drug products for particulate matter and discoloration prior to administration, whenever solution and container permit. The product may contain a few protein particles. Do not use if discolored or cloudy.
- Use a sterile 15 micron in-line filter when administering the product.
- Infuse PROLASTIN-C LIQUID separately, without mixing with other agents or diluting solutions.
- Infuse PROLASTIN-C LIQUID intravenously at 0.08 mL/kg/min as determined by patient response and comfort. The recommended dosage of 60 mg/kg takes approximately 15 minutes to infuse.

3 DOSAGE FORMS AND STRENGTHS

PROLASTIN-C LIQUID is supplied in a 1,000 mg (approximate) single-use vial containing 20 mL of solution for injection. The actual amount of functionally active Alpha1-PI in milligrams is printed on the vial label and carton.

4 CONTRAINDICATIONS

PROLASTIN-C LIQUID is contraindicated in:

- IgA deficient patients with antibodies against IgA, due to the risk of severe hypersensitivity.
- Patients with a history of anaphylaxis or other severe systemic reaction to Alpha1-PI.

5 WARNINGS AND PRECAUTIONS

5.1 Hypersensitivity Reactions

Hypersensitivity reactions, including anaphylaxis, may occur. Monitor vital signs and observe the patient carefully throughout the infusion. Early signs and symptoms of hypersensitivity reactions may include pruritus; generalized urticarial; flushing; swollen lips, tongue, or uvula; wheezing; tightness of the chest; dyspnea; hypotension; and syncope. If hypersensitivity symptoms occur, promptly stop PROLASTIN-C LIQUID infusion and begin appropriate therapy. Have epinephrine and other appropriate therapy available for the treatment of any acute anaphylactic or anaphylactoid reaction. [see Patient Counseling Information (17)]
PROLASTIN-C LIQUID may contain trace amounts of IgA. Patients with known antibodies to IgA, which can be present in patients with selective or severe IgA deficiency, have a greater risk of developing potentially severe hypersensitivity and anaphylactic reactions.

5.2 Transmissible Infectious Agents

Because PROLASTIN-C LIQUID is made from human plasma, it may carry a risk of transmitting infectious agents, e.g., viruses, the variant Creutzfeldt-Jakob disease (vCJD) agent, and, theoretically, the Creutzfeldt-Jakob disease (CJD) agent. This also applies to unknown or emerging viruses and other pathogens. The risk of transmission of infectious agents has been reduced by screening plasma donors for prior exposure to certain infectious agents, by testing for the presence of certain virus infections, and by including steps in the manufacturing process with the demonstrated capacity to inactivate and/or remove certain infectious agents. Despite these measures, this product may still potentially transmit disease.

Report all infections thought by a physician possibly to have been transmitted by this product to Grifols Therapeutics Inc. (1-800-520-2807).

6 ADVERSE REACTIONS

The most serious adverse reaction observed during clinical trials with PROLASTIN-C was an abdominal and extremity rash in one subject. [see Warnings and Precautions (5.1)] The most common adverse reactions observed at a rate of > 5% in subjects receiving PROLASTIN-C LIQUID were diarrhea and fatigue, each of which occurred at a rate of 6% (two subjects each).

6.1 Clinical Trials Experience

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

One clinical trial was conducted with PROLASTIN-C LIQUID: a 16 week, multicenter, randomized, double-blind crossover study to assess the safety, immunogenicity, and pharmacokinetic comparability of PROLASTIN-C LIQUID to PROLASTIN-C in 32 subjects.

Adverse reactions (as defined in the footnote to Table 1) occurring in >5% of subjects during the 16 week double-blind crossover treatment period are shown in Table 1.
Table 1: Adverse Reactions Occurring in >5% of Subjects during the Double-Blinded Crossover Treatment

<table>
<thead>
<tr>
<th>Adverse Reaction*,†</th>
<th>PROLASTIN-C LIQUID (N=32)</th>
<th>PROLASTIN-C (N=31)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of Subjects with Adverse Reaction (percentage of all subjects)</td>
<td>No. of Subjects with Adverse Reaction (percentage of all subjects)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>2 (6)</td>
<td>0</td>
</tr>
<tr>
<td>Fatigue</td>
<td>2 (6)</td>
<td>0</td>
</tr>
</tbody>
</table>

* An adverse reaction is defined as any adverse event that occurred where either a) the event was not considered “unrelated” to administration of the product, or b) the occurrence was during or within 72 hours of the end of the previous infusion of the product, or c) the investigator's causality assessment of the event was missing or indeterminate, or d) the incidence during treatment with 1 investigational product was 130% or more of the incidence during treatment with the other investigational product.

† Source: the randomized double-blinded comparator trial of PROLASTIN-C LIQUID vs PROLASTIN-C.

Table 2 below displays the adverse reaction (defined as per Table 1) rate as a percentage of infusions received during the 16 week double-blinded treatment period.

Table 2: Adverse Reaction Frequency as a Percent of All Infusions and Occurring More than Once in the PROLASTIN-C LIQUID Group during the 16 Week Double Blinded Treatment Period

<table>
<thead>
<tr>
<th>Adverse Reaction*</th>
<th>PROLASTIN-C LIQUID No. of infusions: 252</th>
<th>PROLASTIN-C No. of infusions: 245</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of Adverse Reactions (percentage of all infusions)</td>
<td>No. of Adverse Reactions (percentage of all infusions)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>3 (1.2)</td>
<td>0</td>
</tr>
<tr>
<td>Fatigue</td>
<td>2 (0.8)</td>
<td>0</td>
</tr>
</tbody>
</table>

* Source: the randomized double-blinded comparator trial of PROLASTIN-C LIQUID vs PROLASTIN-C.

A total of 23 COPD exacerbations were reported for a total of 18 individual subjects. Twelve subjects (12/32, 38%) during PROLASTIN-C LIQUID treatment experienced 13 COPD exacerbations, and 9 subjects (9/31, 29%) during PROLASTIN-C treatment had 10 COPD exacerbations. Three COPD exacerbations occurred during the Follow-Up Period after PROLASTIN-C LIQUID treatment and 1 COPD exacerbation occurred in the Follow-up Period.
period after PROLASTIN-C treatment. The overall rate of pulmonary exacerbations during
treatment with either product was 1.9 exacerbations per subject-year. No exacerbation was
considered to be serious, except for one event after PROLASTIN-C treatment during the
Follow-Up period (due to hospitalization).

Two separate prior clinical trials were conducted with PROLASTIN-C: 1.) a 20 week, open-
label, single arm safety study in 38 subjects (single-arm open-label trial), and 2.) a 16 week,
randomized, double-blind, crossover pharmacokinetic comparability study vs. PROLASTIN
in 24 subjects, followed by an 8 week open-label treatment with PROLASTIN-C
(randomized double-blinded comparator trial). Thus, a total of 93 subjects were exposed to
PROLASTIN-C in clinical trials.

The most serious adverse reaction observed during clinical trials with PROLASTIN-C was
an abdominal and extremity rash in one subject. The rash resolved subsequent to outpatient
treatment with antihistamines and steroids. Two instances of a less severe, pruritic abdominal
rash were observed upon rechallenge despite continued antihistamine and steroid treatment,
which led to withdrawal of the subject from the trial.

Grifols assessed the randomized double-blinded comparator trial of PROLASTIN and
PROLASTIN-C for adverse reactions (as defined in the footnote to Table 3) occurring during
each 8 week double-blind crossover treatment period, as shown in Table 3.
### Table 3: Adverse Reactions Occurring during the First 8 Weeks of Each Double-Blinded Treatment

<table>
<thead>
<tr>
<th>Adverse Reaction *†</th>
<th>PROLASTIN®-C</th>
<th>PROLASTIN®</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of subjects: 24</td>
<td>No. of subjects: 24</td>
<td></td>
</tr>
<tr>
<td></td>
<td>No. of Subjects with Adverse Reaction (percentage of all subjects)</td>
<td>No. of Subjects with Adverse Reaction (percentage of all subjects)</td>
<td></td>
</tr>
<tr>
<td>Upper respiratory tract infection</td>
<td>3 (12.5%)</td>
<td>1 (4.2%)</td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>1 (4.2%)</td>
<td>2 (8.3%)</td>
<td></td>
</tr>
<tr>
<td>Pruritus</td>
<td>1 (4.2%)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Urticaria</td>
<td>1 (4.2%)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Nausea</td>
<td>1 (4.2%)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Peripheral edema</td>
<td>1 (4.2%)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Pyrexia</td>
<td>1 (4.2%)</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

* An adverse reaction is defined as any adverse event where either a) the incidence with PROLASTIN-C was greater than with PROLASTIN, or b) the occurrence was within 72 hours of treatment, or c) the event was otherwise considered related or possibly related to the drug.

† Source: the randomized double-blinded comparator trial.

Table 4 below displays the adverse reaction (defined as per Table 3) rate as a percentage of infusions received during the 8 weeks of each double-blinded treatment.
### Table 4: Adverse Reaction Frequency as a Percent of All Infusions during the First 8 Weeks of Each Double-Blinded Infusion Treatment

<table>
<thead>
<tr>
<th>Adverse Reaction*</th>
<th>PROLASTIN®-C</th>
<th>PROLASTIN®</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of infusions: 188</td>
<td>No. of infusions: 192</td>
</tr>
<tr>
<td>No. of Adverse Reactions</td>
<td>No. of Adverse Reactions</td>
<td></td>
</tr>
<tr>
<td>(percentage of all infusions)</td>
<td>(percentage of all infusions)</td>
<td></td>
</tr>
<tr>
<td>Upper respiratory tract infection</td>
<td>3 (1.6%)</td>
<td>1 (0.5%)</td>
</tr>
<tr>
<td>Headache</td>
<td>1 (0.5%)</td>
<td>3 (1.6%)</td>
</tr>
<tr>
<td>Pruritus</td>
<td>1 (0.5%)</td>
<td>0</td>
</tr>
<tr>
<td>Urticaria</td>
<td>1 (0.5%)</td>
<td>0</td>
</tr>
<tr>
<td>Nausea</td>
<td>1 (0.5%)</td>
<td>0</td>
</tr>
<tr>
<td>Peripheral edema</td>
<td>1 (0.5%)</td>
<td>0</td>
</tr>
<tr>
<td>Pyrexia</td>
<td>1 (0.5%)</td>
<td>0</td>
</tr>
</tbody>
</table>

* Source: the randomized double-blinded comparator trial.

Table 5 below displays the adverse reactions occurring in two or more subjects during the single-arm open-label trial.
Table 5: Adverse Reactions Occurring in Two or More Subjects (>5%) during the 20 Week Single-Arm Open-Label Trial

<table>
<thead>
<tr>
<th>Adverse Reaction</th>
<th>PROLASTIN®-C No. of subjects: 38</th>
<th>No. of Subjects with Adverse Reaction (percentage of all subjects)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upper respiratory tract infection</td>
<td></td>
<td>6 (15.8%)</td>
</tr>
<tr>
<td>Urinary tract infection</td>
<td></td>
<td>5 (13.2%)</td>
</tr>
<tr>
<td>Nausea</td>
<td></td>
<td>4 (10.5%)</td>
</tr>
<tr>
<td>Chest pain</td>
<td></td>
<td>3 (7.9%)</td>
</tr>
<tr>
<td>Back pain</td>
<td></td>
<td>2 (5.3%)</td>
</tr>
<tr>
<td>Chills</td>
<td></td>
<td>2 (5.3%)</td>
</tr>
<tr>
<td>Cough</td>
<td></td>
<td>2 (5.3%)</td>
</tr>
<tr>
<td>Dizziness</td>
<td></td>
<td>2 (5.3%)</td>
</tr>
<tr>
<td>Dyspnea</td>
<td></td>
<td>2 (5.3%)</td>
</tr>
<tr>
<td>Headache</td>
<td></td>
<td>2 (5.3%)</td>
</tr>
<tr>
<td>Hot flush</td>
<td></td>
<td>2 (5.3%)</td>
</tr>
<tr>
<td>Oral candidiasis</td>
<td></td>
<td>2 (5.3%)</td>
</tr>
</tbody>
</table>

* An adverse reaction is defined as any adverse event that occurred where either a) the occurrence was within 72 hours of treatment, or b) the event was otherwise considered related or possibly related to the drug.
† Source: the single-arm, open-label trial.

Ten exacerbations of chronic obstructive pulmonary disease were reported by 8 subjects in the 24 week crossover pharmacokinetic study. During the 16 week double-blind crossover phase, 4 subjects (17%) had a total of 4 exacerbations during PROLASTIN-C treatment and 4 subjects (17%) had a total of 4 exacerbations during PROLASTIN treatment. Two additional exacerbations in 2 subjects (8%) occurred during the 8 week open-label treatment period with PROLASTIN-C. The overall rate of pulmonary exacerbations during treatment with either product was 0.9 exacerbations per subject-year.
Immunogenicity

The detection of antibody formation is highly dependent on the sensitivity and specificity of the assay. Additionally, the observed incidence of antibody (including neutralizing antibody) positivity in an assay may be influenced by several factors including assay methodology, sample handling, timing of sample collection, concomitant medications, and underlying disease. For these reasons, comparison of the incidence of antibodies to PROLASTIN-C LIQUID with the incidence of antibodies to other products may be misleading.

In the randomized, crossover pharmacokinetic clinical trial, no immunogenicity response was observed in subjects dosed with PROLASTIN-C LIQUID or PROLASTIN-C.

In the single-arm, open-label safety clinical trial, three treatment naïve subjects out of 36 subjects evaluated developed antibody to Alpha1-PI at week 24 after receiving PROLASTIN-C. A fourth subject (non-naïve) was positive prior to and after receiving PROLASTIN-C, but levels were unchanged during the study. None of the four antibody specimens was able to neutralize the protease inhibitor capacity of PROLASTIN-C. In the randomized, crossover pharmacokinetic clinical trial comparing PROLASTIN-C and PROLASTIN, none of 24 subjects developed antibodies to PROLASTIN-C.

6.2 Postmarketing Experience

Because postmarketing reporting of adverse reactions is voluntary and from a population of uncertain size, it is not always possible to reliably estimate the frequency of these reactions or establish a causal relationship to product exposure.

Expected postmarketing experience for PROLASTIN-C LIQUID is based on the reactions reported for PROLASTIN-C. The reactions which have been chosen for inclusion due to their seriousness, frequency of reporting, possible causal connection to PROLASTIN®–C, or a combination of these factors, are:

- **General/Body as a Whole:** Fatigue, malaise, influenza-like illness, pain, asthenia
- **Immune system:** Hypersensitivitiy including anaphylactoid/anaphylactic reactions
- **Cardiovascular:** Tachycardia
- **Musculoskeletal:** Arthralgia, myalgia
- **Gastrointestinal:** Vomiting, diarrhea
- **Investigation:** Blood pressure increased
8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

There are no data with PROLASTIN-C LIQUID use in pregnant women to inform a drug-associated risk. Animal reproduction studies have not been conducted with PROLASTIN-C LIQUID. It is not known whether PROLASTIN-C LIQUID can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. PROLASTIN-C LIQUID should be given to a pregnant woman only if clearly needed.

In the U.S. general population, the estimated background risk of major birth defect and miscarriage in clinically recognized pregnancies is 2-4% and 15-20%, respectively.

8.2 Lactation

Risk Summary

There is no information regarding the presence of PROLASTIN-C LIQUID in human milk, the effects on the breastfed infant, or the effects on milk production. The developmental and health benefits of breastfeeding should be considered along with the mother’s clinical need for PROLASTIN-C LIQUID and any potential adverse effects on the breastfed infant from PROLASTIN-C LIQUID or from the underlying maternal condition.

8.4 Pediatric Use

Safety and effectiveness in the pediatric population have not been established.

8.5 Geriatric Use

Clinical studies of PROLASTIN-C LIQUID did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. As for all patients, dosing for geriatric patients should be appropriate to their overall situation.

11 DESCRIPTION

PROLASTIN-C LIQUID is a sterile, concentrate of Alpha1-PI for intravenous infusion. The solution is clear, colorless or pale yellow or pale green. Each vial of PROLASTIN-C LIQUID contains approximately 1,000 mg of functionally active Alpha1-PI as determined by capacity to neutralize porcine pancreatic elastase. The specific activity of PROLASTIN-C LIQUID is ≥ 0.7 mg functional Alpha1-PI per mg of total protein. PROLASTIN-C LIQUID has a purity of ≥ 90% Alpha1-PI (Alpha1-PI protein/total protein). PROLASTIN-C LIQUID has a pH of 6.6–7.4, a sodium phosphate content of 0.013–0.025 M, and is stabilized with 0.20-0.30 M of alanine. The total sodium concentration is ≤ 100 mEq/L. PROLASTIN-C LIQUID contains no preservative.
PROLASTIN-C LIQUID is produced from pooled human plasma through modifications of
the PROLASTIN process using purification by polyethylene glycol (PEG) precipitation,
anion exchange chromatography, and cation exchange chromatography. All Source Plasma
used in the manufacture of PROLASTIN-C LIQUID is non-reactive (negative) by FDA-
licensed serological test methods for hepatitis B surface antigen (HBsAg) and antibodies to
hepatitis C virus (HCV) and human immunodeficiency virus types 1 and 2 and negative by
FDA-licensed Nucleic Acid Technologies (NAT) for HCV and human immunodeficiency
virus type 1 (HIV-1). In addition, all Source Plasma is negative for hepatitis B virus (HBV)
by either an FDA-licensed or investigational NAT assay. The goal of the investigational
HBV NAT test is to detect low levels of viral nucleic acid; however, the significance of a
negative result for the investigational HBV NAT test has not been established. By in-process
NAT, all Source Plasma is negative for hepatitis A virus (HAV). As a final plasma safety
step, all plasma manufacturing pools are tested by serological test methods and NAT.

To evaluate further the virus safety profile of PROLASTIN-C LIQUID, *in vitro* studies have
been conducted to validate the capacity of the manufacturing process to reduce the infectious
titer of a wide range of viruses with diverse physicochemical properties. These studies
evaluated the inactivation/removal of clinically relevant viruses, including human
immunodeficiency virus type 1 (HIV-1) and hepatitis A virus (HAV), as well as the
following model viruses: bovine viral diarrhea virus (BVDV), a surrogate for hepatitis C
virus; pseudorabies virus (PRV), a surrogate for large enveloped DNA viruses (e.g., herpes
viruses); vesicular stomatitis virus (VSV), a model for enveloped viruses; reovirus type 3
(Reo3), a non-specific model for non-enveloped viruses; and porcine parvovirus (PPV), a
model for human parvovirus B19.

The PROLASTIN-C LIQUID manufacturing process has several steps (Cold Ethanol
Fractionation, PEG Precipitation, and Depth Filtration) that are important for purifying
Alpha1-PI as well as removing potential virus contaminants. Two additional steps,
Solvent/Detergent Treatment and 15 nm Virus Removal Nanofiltration, are included in the
process as dedicated pathogen reduction steps. The Solvent/Detergent Treatment step
effectively inactivates enveloped viruses (such as HIV-1, VSV, HBV, and HCV). The 15 nm
Virus Removal Nanofiltration step has been implemented to reduce the risk of transmission
of enveloped and non-enveloped viruses as small as 18 nm. *Table 6* presents the virus
reduction capacity of each process step and the accumulated virus reduction for the process
as determined in viral validation studies in which virus was deliberately added to a process
model in order to study virus reduction. In addition, the Solvent/Detergent Treatment step
inactivates ≥ 5.4 log_{10} of West Nile virus, a clinically relevant enveloped virus.
Table 6: Virus Reduction (Log_{10}) for the PROLASTIN®-C LIQUID Manufacturing Process

<table>
<thead>
<tr>
<th>Process Step</th>
<th>Enveloped Viruses</th>
<th>Non-enveloped Viruses</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HIV-1</td>
<td>BVDV</td>
</tr>
<tr>
<td>Cold Ethanol Fractionation</td>
<td>1.5</td>
<td>1.7</td>
</tr>
<tr>
<td>PEG Precipitation</td>
<td>4.3</td>
<td>2.8</td>
</tr>
<tr>
<td>Depth Filtration</td>
<td>≥ 4.7</td>
<td>4.0</td>
</tr>
<tr>
<td>Solvent/Detergent Treatment</td>
<td>≥ 6.2</td>
<td>≥ 4.6</td>
</tr>
<tr>
<td>15 nm Virus Removal Nanofiltration</td>
<td>≥ 6.9</td>
<td>≥ 4.7</td>
</tr>
<tr>
<td>Accumulated Virus Reduction</td>
<td>≥ 23.6</td>
<td>≥ 17.8</td>
</tr>
</tbody>
</table>

* Not determined. VSV inactivation and/or removal was only determined for the Solvent/Detergent Treatment and 15 nm Virus Removal Nanofiltration steps.
† Not applicable. This step is only effective against enveloped viruses.

Additionally, the manufacturing process was investigated for its capacity to decrease the infectivity of an experimental agent of transmissible spongiform encephalopathy (TSE), considered as a model for the variant Creutzfeldt-Jakob disease (vCJD) and Creutzfeldt-Jakob disease (CJD) agents. Studies of the PROLASTIN-C LIQUID manufacturing process demonstrate that a minimum of 6 log_{10} reduction of TSE infectivity is achieved. These studies provide reasonable assurance that low levels of vCJD/CJD agent infectivity, if present in the starting material, would be removed.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Alpha_{1}-PI deficiency is an autosomal, co-dominant, hereditary disorder characterized by low serum and lung levels of Alpha_{1}-PI. Smoking is an important risk factor for the development of emphysema in patients with Alpha_{1}-PI deficiency. Because emphysema affects many, but not all individuals with the more severe genetic variants of Alpha_{1}-PI deficiency, augmentation therapy with Alpha_{1}-PI is indicated only in patients with severe Alpha_{1}-PI deficiency who have clinically evident emphysema.

Only some Alpha_{1}-PI alleles are associated with clinically apparent Alpha_{1}-PI deficiency. Approximately 95% of all severely deficient patients are homozygous for the PiZ allele. Individuals with the PiZZ variant typically have serum Alpha_{1}-PI levels less than 35% of the average normal level. Individuals with the Pi(null)(null) variant have undetectable Alpha_{1}-PI protein in their serum. Individuals with these low serum Alpha_{1}-PI levels, i.e., less than 11 \mu M, have a markedly increased risk for developing emphysema over their lifetimes. In addition, PiSZ individuals, whose serum Alpha_{1}-PI levels range from approximately 9 to 23
µM, are considered to have moderately increased risk for developing emphysema, regardless of whether their serum Alpha1-PI levels are above or below 11 µM.

Augmenting the levels of functional protease inhibitor by intravenous infusion is an approach to therapy for patients with Alpha1-PI deficiency. The intended theoretical goal is to provide protection to the lower respiratory tract by correcting the imbalance between neutrophil elastase and protease inhibitors. Whether augmentation therapy with any Alpha1-PI product actually protects the lower respiratory tract from progressive emphysematous changes has not been demonstrated in adequately powered, randomized controlled, clinical trials. Although the maintenance of blood serum levels of Alpha1-PI (antigenically measured) above 11 µM has been historically postulated to provide therapeutically relevant anti-neutrophil elastase protection, this has not been proven. Individuals with severe Alpha1-PI deficiency have been shown to have increased neutrophil and neutrophil elastase concentrations in lung epithelial lining fluid compared to normal PiMM individuals, and some PiSZ individuals with Alpha1-PI above 11 µM have emphysema attributed to Alpha1-PI deficiency. These observations underscore the uncertainty regarding the appropriate therapeutic target serum level of Alpha1-PI during augmentation therapy.

The pathogenesis of emphysema is understood as described in the “protease-antiprotease imbalance” model. Alpha1-PI is understood to be the primary antiprotease in the lower respiratory tract, where it inhibits neutrophil elastase (NE). Normal healthy individuals produce sufficient Alpha1-PI to control the NE produced by activated neutrophils and are thus able to prevent inappropriate proteolysis of the lung tissue by NE. Conditions that increase neutrophil accumulation and activation in the lung, such as respiratory infection and smoking, will in turn increase levels of NE. However, individuals who are severely deficient in endogenous Alpha1-PI are unable to maintain an appropriate antiprotease defense, and, in addition, they have been shown to have increased lung epithelial lining fluid neutrophil and NE concentrations. Because of these factors, many (but not all) individuals who are severely deficient in endogenous Alpha1-PI are subject to more rapid proteolysis of the alveolar walls leading to chronic lung disease. PROLASTIN-C LIQUID serves as Alpha1-PI augmentation therapy in the patient population with severe Alpha1-PI deficiency and emphysema, acting to increase and maintain serum and lung epithelial lining fluid levels of Alpha1-PI.

12.2 Pharmacodynamics

Chronic augmentation therapy with the predecessor product, PROLASTIN® (Alpha1-Proteinase Inhibitor [Human]), administered weekly at a dose of 60 mg/kg body weight, results in increased levels of Alpha1-PI and functional anti-neutrophil elastase capacity in the epithelial lining fluid of the lower respiratory tract of the lung, as compared to levels prior to commencing therapy with PROLASTIN. However, the clinical benefit of the increased levels at the recommended dose has not been demonstrated in adequately powered, randomized, controlled clinical trials for any Alpha1-PI product.

PROLASTIN-C LIQUID increases antigenic and functional (anti-neutrophil elastase capacity, ANEC) serum levels.
12.3 Pharmacokinetics

The pharmacokinetic (PK) study was a randomized, double-blind, crossover trial comparing PROLASTIN-C LIQUID to PROLASTIN-C conducted in 32 adult subjects age 44 to 71 years with severe Alpha1-PI deficiency. Eighteen subjects were male and 14 subjects were female. Sixteen subjects were randomized to each treatment sequence. All but one subject had the PiZZ genotype and the remaining subject was PiSZ. Twenty-eight subjects had received prior Alpha1-PI augmentation therapy and 4 subjects were naïve to Alpha1-PI augmentation therapy. Study subjects were randomly assigned to receive either 60 mg/kg body weight of functional PROLASTIN-C LIQUID or PROLASTIN-C weekly by intravenous infusion during the first 8-week treatment period. Following the last dose in the first 8-week treatment period, subjects underwent serial blood sampling for PK analysis and then crossed over to the alternate treatment for the second 8-week treatment period.

Following the last treatment in the second 8-week treatment period, subjects underwent serial blood sampling for PK analysis. In addition, blood samples were drawn for trough levels before infusion at Weeks 6, 7, 8, and 9, as well as before infusion at Weeks 14, 15, 16, and 17. A final PK sample was drawn at Week 20 (4 weeks after the last dose) to correct for endogenous Alpha1-PI levels.

The key pharmacokinetic parameter was the area under the plasma concentration-time curve (AUC0-7days) following 8 weeks of treatment with PROLASTIN-C LIQUID or PROLASTIN-C. The 90% confidence interval (1.03-1.08) for the ratio of AUC0-7days for PROLASTIN-C LIQUID and PROLASTIN-C indicated that the 2 products are bioequivalent, i.e. the entire range falls within the 80 – 125% interval. Figure 1 shows the concentration (functional activity) vs. time curves of Alpha1-PI after intravenous administration of PROLASTIN-C LIQUID and PROLASTIN-C.

<table>
<thead>
<tr>
<th>Table 7: Pharmacokinetic Parameters of Alpha1-PI in Plasma</th>
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<tbody>
<tr>
<td>Treatment</td>
</tr>
<tr>
<td>PROLASTIN-C LIQUID n=30</td>
</tr>
<tr>
<td>PROLASTIN-C n=28</td>
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Figure 1: Mean Plasma Alpha$_1$-PI Concentration (functional activity) vs. Time Curves Following Treatment with PROLASTIN-C LIQUID or PROLASTIN-C

Trough levels measured at steady state during the PK study using an antigenic content assay showed PROLASTIN-C LIQUID resulted in a mean trough of 17.72 µM and PROLASTIN-C resulted in a mean trough of 16.88 µM.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenesis, mutagenesis, and impairment of fertility studies were not performed; PROLASTIN-C LIQUID is a biologic purified from human plasma.
13.2 Animal Toxicology and/or Pharmacology

Intravenous administration of five daily doses of PROLASTIN-C LIQUID to rabbits at a dose up to 600 mg/kg per day (10-fold higher dose than the recommended human dose of 60 mg/kg administered weekly), did not result in any signs of toxicity. Further, there were no differences in safety and tolerability of PROLASTIN-C and PROLASTIN-C LIQUID in nonclinical testing.

14 CLINICAL STUDIES

The clinical efficacy of PROLASTIN-C LIQUID in influencing the course of pulmonary emphysema or pulmonary exacerbations has not been demonstrated in adequately powered, randomized, controlled clinical trials.

A total of 23 subjects with the PiZZ variant and documented emphysema were studied in a single-arm, open label clinical trial with PROLASTIN, the predecessor product. Nineteen of the subjects received PROLASTIN, 60 mg/kg, once weekly for up to 26 weeks (average 24 weeks). Blood levels of Alpha1-PI were maintained above 11 µM. Bronchoalveolar lavage studies demonstrated statistically significant increased levels of Alpha1-PI and functional ANEC in the epithelial lining fluid of the lower respiratory tract of the lung, as compared to levels prior to dosing.

In addition to the PROLASTIN-C LIQUID/PROLASTIN-C crossover trial described above, in which 31 subjects received PROLASTIN-C, PROLASTIN-C has been studied in 62 individual subjects in 2 separate clinical trials. The first study was a crossover pharmacokinetic study. [see Clinical Pharmacology (12.3)] The second PROLASTIN-C clinical trial was a multi-center, open-label single arm safety study conducted to evaluate the safety and tolerability of PROLASTIN-C. In this study, 38 subjects were treated with weekly intravenous infusions of 60 mg/kg body weight of PROLASTIN-C for 20 weeks. Half the subjects were naïve to previous Alpha1-PI augmentation prior to study entry and the other half were receiving augmentation with PROLASTIN prior to entering the study. A diagnosis of severe Alpha1-PI deficiency was confirmed by the demonstration of the PiZZ genotype in 32 of 38 (84.2%) subjects, and 6 of 38 (15.8%) subjects presented with other alleles known to result in severe Alpha1-PI deficiency. These groups were distributed evenly between the naïve and non-naïve cohorts.

15 REFERENCES


16 HOW SUPPLIED/STORAGE AND HANDLING

- PROLASTIN-C LIQUID is supplied in a single-use vial with the total Alpha1-PI functional activity, in milligrams, stated on the vial label and carton.

- Components of the packaging do not contain natural rubber latex.

<table>
<thead>
<tr>
<th>NDC Number Carton</th>
<th>Approximate Alpha1–PI Functional Activity</th>
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<tr>
<td>13533-705-01</td>
<td>1,000 mg</td>
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</table>

- Store refrigerated at 2-8°C (36-46°F) for the period indicated by the expiration date on its label.
  - Product may be stored at room temperatures not exceeding 25°C (77°F) for up to one month, after which the product must be used or immediately discarded.

- Do not freeze.

17 PATIENT COUNSELING INFORMATION

- Inform patients of the signs of hypersensitivity reactions including pruritus; generalized urticarial; flushing; swollen lips, tongue, or uvula; wheezing; tightness of the chest; dyspnea; hypotension; and syncope. Advise patients to discontinue use of the product and contact their physician and/or seek immediate emergency care, depending on the severity of the reaction, if these symptoms occur. [see Warnings and Precautions (5.1)]

- Inform patients that PROLASTIN-C LIQUID is made from human plasma and may carry a risk of transmitting infectious agents that can cause disease (e.g., viruses, the vCJD
agent and, theoretically, the CJD agent). Explain that the risk of PROLASTIN-C LIQUID transmitting an infectious agent has been reduced by screening plasma donors for prior exposure to certain infectious agents, by testing the donated plasma for certain current virus infections, and by inactivating and/or removing infectious agents during manufacturing. [see Warnings and Precautions (5.2)]

- Inform patients that administration of PROLASTIN-C LIQUID has been demonstrated to raise the plasma level of Alpha1-PI, but that the effect of this augmentation on pulmonary exacerbations and on the rate of progression of emphysema has not been demonstrated in adequately powered, randomized, controlled clinical trials for any Alpha1-PI product. [see Clinical Studies (14)]

Manufactured by:

**GRIFOLS**

**Grifols Therapeutics Inc.**

Research Triangle Park, NC 27709 USA

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