HIGHLIGHTS OF PRESCRIBING INFORMATION These highlights do not include all the information needed to use STRENSIQ safely and effectively. See full prescribing information for STRENSIO.

STRENSIQ[®] (asfotase alfa) injection, for subcutaneous use Initial U.S. Approval: 2015

RECENT MAJOR CHANGES	
KECENT MAJUK UNANGES	
Dosing and Administration (2.3, 2.4)	6/2020
Warnings and Precautions (5.4)	6/2020

-INDICATIONS AND USAGE-

STRENSIQ is a tissue nonspecific alkaline phosphatase indicated for the treatment of patients with perinatal/infantile- and juvenile-onset hypophosphatasia (HPP). (1)

DOSAGE AND ADMINISTRATION-

Perinatal/Infantile-Onset HPP (2.1)

- Recommended dosage regimen is 2 mg/kg administered subcutaneously three times per week, or 1 mg/kg administered six times per week. Injection site reactions may limit the tolerability of the six times per week regimen.
- The dose may be increased to 3 mg/kg three times per week for insufficient efficacy.

Juvenile-Onset HPP (2.2)

Recommended dosage regimen is 2 mg/kg administered subcutaneously three times per week, or 1 mg/kg administered six times per week. Injection site reactions may limit the tolerability of the six times per week regimen.

Preparation and Weight-Based Dosing (2.3):

- Caution: Do not use the 80 mg/0.8 mL vial in pediatric patients weighing less than 40 kg because the systemic asfotase alfa exposure achieved with the 80 mg/0.8 mL vial (higher concentration) is lower than that achieved with the other strength vials (lower concentration). A lower exposure may not be adequate for this subgroup of patients.
- See full prescribing information for tables of weight-based dosing by treatment regimen.

Administration (2.4):

- For subcutaneous injection only.
- Rotate injection sites. Do not administer to areas that are reddened, inflamed or swollen.

-DOSAGE FORMS AND STRENGTHS-

Injection: 18 mg/0.45 mL, 28 mg/0.7 mL, 40 mg/mL, or 80 mg/0.8 mL solution in single-dose vials. (3)

- CONTRAINDICATIONS -

None. (4)

- WARNINGS AND PRECAUTIONS -

- <u>Hypersensitivity Reactions:</u> Monitor and if a severe reaction occurs, discontinue treatment and initiate appropriate medical treatment. (5.1)
- Lipodystrophy: Localized reactions were reported after several months of treatment; follow proper injection technique and rotate injection sites. (5.2)
- Ectopic Calcifications (eye and kidneys): Monitor using ٠ ophthalmologic examinations and renal ultrasounds at baseline and periodically during treatment. (5.3)
- Possible Immune-Mediated Clinical Effects: Evaluate patients for antibody formation if clinically indicated. (5.4)

-ADVERSE REACTIONS-

Most common adverse reactions ($\geq 10\%$) are injection site reactions, lipodystrophy, ectopic calcifications and hypersensitivity reactions. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact Alexion Pharmaceuticals, Inc. at 1-844-259-6783 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

-DRUG INTERACTIONS-

- Drug Interference with Laboratory Tests: Alkaline Phosphatase (ALP) is used as a detection reagent in many laboratory tests and the presence of asfotase alfa in clinical laboratory samples could result in erroneous test results. Inform laboratory personnel and discuss use of an alternative testing platform for patients on treatment. (7.1)
- Serum Alkaline Phosphatase: Serum ALP measurements are expected to be elevated during treatment and may be unreliable for clinical decision making. (7.1)

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling.

Revised: 6/2020

FULL PRESCRIBING INFORMATION: CONTENTS*

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FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

STRENSIQ[®] is indicated for the treatment of patients with perinatal/infantile- and juvenile-onset hypophosphatasia (HPP).

2 DOSAGE AND ADMINISTRATION

2.1 Dosage for Perinatal/Infantile-Onset HPP

The recommended dosage regimen of STRENSIQ for the treatment of perinatal/infantileonset HPP is 6 mg/kg per week administered subcutaneously as either:

- 2 mg/kg three times per week, or
- 1 mg/kg six times per week. Injection site reactions may limit the tolerability of the six times per week regimen [see Adverse Reactions (6.1)].

The dose of STRENSIQ may be increased for lack of efficacy (e.g., no improvement in respiratory status, growth, or radiographic findings) up to 9 mg/kg per week administered subcutaneously as 3 mg/kg three times per week.

2.2 Dosage for Juvenile-Onset HPP

The recommended dosage regimen of STRENSIQ for the treatment of juvenile-onset HPP is 6 mg/kg per week administered subcutaneously as either:

- 2 mg/kg three times per week, or
- 1 mg/kg six times per week. Injection site reactions may limit the tolerability of the six times per week regimen [see Adverse Reactions (6.1)].

2.3 Preparation and Weight-Based Dosing Tables

Caution: Do not use the 80 mg/0.8 mL vial of STRENSIQ in pediatric patients weighing less than 40 kg because the systemic exposure of asfotase alfa achieved with the 80 mg/0.8 mL vial (higher concentration) is lower than that achieved with the other strength vials (lower concentration). A lower exposure may not be adequate for this subgroup of patients [see Dosage Forms and Strengths (3), Clinical Pharmacology (12.3)].

1. Determine the total weekly volume needed for the prescribed dosage based on the patient's weight and recommended dosage. Follow these steps to determine the patient dose.

Total weekly dose (mg) = patient's weight (kg) x prescribed dose (mg/kg/week)

Total injection volume (mL) per week = Total dose (mg/week) divided by the STRENSIQ concentration (40 mg/mL or 100 mg/mL). Note product concentrations are: 40 mg/mL (vial strengths 18 mg/0.45 mL, 28 mg/0.7 mL, 40 mg/mL) or 100 mg/mL (vial strength 80 mg/0.8 mL).

Round total injection volume to the nearest hundredth of a mL

Total number of vials per week = Total injection volume divided by vial volume (mL)

2. Determine the number of injection days per week (three or six per week).

3. Determine dose per injection day. Patient weights should be rounded to the nearest kilogram when determining dose. Use the following tables for guidance, for patients administering 2 mg/kg three times per week (Table 1), 1 mg/kg six times per week (Table 2) and for dose escalations to 3 mg/kg three times per week, recommended only for patients with perinatal/infantile-onset HPP [see Dosage and Administration (2.1)] (Table 3).

Table 1:Weight-Based Dosing for Administration of 2 mg/kg Three Times per
Week

Body Weight (kg)*	Dose to Inject	Volume to Inject	Vial Configuration
3	6 mg	0.15 mL	18 mg/0.45 mL
4	8 mg	0.2 mL	18 mg/0.45 mL
5	10 mg	0.25 mL	18 mg/0.45 mL
6	12 mg	0.3 mL	18 mg/0.45 mL
7	14 mg	0.35 mL	18 mg/0.45 mL
8	16 mg	0.4 mL	18 mg/0.45 mL
9	18 mg	0.45 mL	18 mg/0.45 mL
10	20 mg	0.5 mL	28 mg/0.7 mL
15	30 mg	0.75 mL	40 mg/1 mL
20	40 mg	1 mL	40 mg/1 mL
25	50 mg	1.25 mL	Two 28 mg/0.7 mL vials
30	60 mg	1.5 mL	Two 40 mg/1 mL vials
35	70 mg	1.75 mL	Two 40 mg/1 mL vials
40	80 mg	0.8 mL	80 mg/0.8 mL
50	100 mg	1 mL	Two 80 mg/0.8 mL vials
60	120 mg	1.2 mL**	Two 80 mg/0.8 mL vials
70	140 mg	1.4 mL**	Two 80 mg/0.8 mL vials
80	160 mg	1.6 mL**	Two 80 mg/0.8 mL vials

* Do not use the 80 mg/0.8 mL vial of STRENSIQ in pediatric patients weighing less than 40 kg [see Clinical Pharmacology (12.3)].

^{**} When preparing a volume for injection greater than 1 mL, split the volume equally between two syringes, and administer two injections. When administering the two injections, use two separate injection sites.

Body Weight (kg)*	Dose to Inject	Volume to Inject	Vial Configuration
3	3 mg	0.08 mL	18 mg/0.45 mL
4	4 mg	0.1 mL	18 mg/0.45 mL
5	5 mg	0.13 mL	18 mg/0.45 mL
6	6 mg	0.15 mL	18 mg/0.45 mL
7	7 mg	0.18 mL	18 mg/0.45 mL
8	8 mg	0.2 mL	18 mg/0.45 mL
9	9 mg	0.23 mL	18 mg/0.45 mL
10	10 mg	0.25 mL	18 mg/0.45 mL
15	15 mg	0.38 mL	18 mg/0.45 mL
20	20 mg	0.5 mL	28 mg/0.7 mL
25	25 mg	0.63 mL	28 mg/0.7 mL
30	30 mg	0.75 mL	40 mg/1 mL
35	35 mg	0.88 mL	40 mg/1 mL
40	40 mg	1 mL	40 mg/1 mL
50	50 mg	0.5 mL	80 mg/0.8 mL
60	60 mg	0.6 mL	80 mg/0.8 mL
70	70 mg	0.7 mL	80 mg/0.8 mL
80	80 mg	0.8 mL	80 mg/0.8 mL
90	90 mg	0.9 mL	Two 80 mg/0.8 mL vial
100	100 mg	1 mL	Two 80 mg/0.8 mL vial

Table 2:Weight-Based Dosing for Administration of 1 mg/kg Six Times per
Week

* Do not use the 80 mg/0.8 mL vial of STRENSIQ in pediatric patients weighing less than 40 kg [see Clinical Pharmacology (12.3)].

Table 3:Weight-Based Dosing for Administration of 3 mg/kg Three Times per
Week – Only for Perinatal/Infantile-Onset HPP*

Body Weight (kg)**	Dose to Inject	Volume to Inject	Vial Configuration
3	9 mg	0.23 mL	18 mg/0.45 mL
4	12 mg	0.3 mL	18 mg/0.45 mL
5	15 mg	0.38 mL	18 mg/0.45 mL
6	18 mg	0.45 mL	18 mg/0.45 mL
7	21 mg	0.53 mL	28 mg/0.7 mL
8	24 mg	0.6 mL	28 mg/0.7 mL
9	27 mg	0.68 mL	28 mg/0.7 mL
10	30 mg	0.75 mL	40 mg/1 mL
15	45 mg	1.13 mL***	Two 28 mg/0.7 mL vials
20	60 mg	1.5 mL***	Two 40 mg/1 mL vials
25	75 mg	1.88 mL***	Two 40 mg/1 mL vials

* A regimen of 3 mg/kg three times per week is recommended only for patients with perinatal/infantile-onset HPP [see Dosage and Administration (2.1)]

** Do not use the 80 mg/0.8 mL vial of STRENSIQ in pediatric patients weighing less than 40 kg [see Clinical Pharmacology (12.3)].

**** When preparing a volume for injection greater than 1 mL, split the volume equally between two syringes, and administer two injections. When administering the two injections, use two separate injection sites.

4. Take the unopened STRENSIQ vial(s) out of the refrigerator 15 to 30 minutes before injecting to allow the liquid to reach room temperature.

Do not warm STRENSIQ in any other way (for example, do not warm it in a microwave or in hot water).

- 5. Inspect the solution in the vial(s) for particulate matter and discoloration. STRENSIQ is supplied as a clear, slightly opalescent or opalescent, colorless to slightly yellow aqueous solution; a few small translucent or white particles may be present. Discard any vial(s) not consistent with this appearance.
- 6. Assemble injection supplies. Administer STRENSIQ using sterile disposable 1 mL syringes and ½ inch injection needles, between 25 to 29 gauge are recommended. The use of two different gauge needles is recommended, a larger bore needle (e.g. 25 gauge) for withdrawal of the medication, and a smaller bore needle (e.g. 29 gauge) for the injection. For doses greater than 1 mL, the injection volume should be split equally between two 1 mL syringes. Always use a new syringe and needle for each injection.
- 7. Remove vial cap, aseptically prepare the vial and insert the syringe into the vial to withdraw the prescribed dose for administration. Do not shake.
- 8. Remove any air bubbles in the syringe and verify the correct dose.
- 9. STRENSIQ vials are for one time use only. Discard any unused product.

2.4 Administration

STRENSIQ is for subcutaneous injection only.

- When administering two injections, use two separate injection sites.
- Administer STRENSIQ within 3 hours upon removal of the vial(s) from refrigeration.
- Rotate the injection from among the following sites to reduce the risk of lipodystrophy: abdominal area, thigh, deltoid, or buttocks [see Warnings and *Precautions (5.2), Adverse Reactions (6.1)*].
- Do NOT administer injections in areas that are reddened, inflamed, or swollen.
- Inject STRENSIQ subcutaneously into the determined site and properly dispose of the syringe and the needle.

3 DOSAGE FORMS AND STRENGTHS

STRENSIQ is a clear, slightly opalescent or opalescent, colorless to slightly yellow aqueous solution; few small translucent or white particles may be present. The product is available as:

• Injection: 18 mg/0.45 mL, 28 mg/0.7 mL, 40 mg/mL, or 80 mg/0.8 mL solution in single-dose vials

4 CONTRAINDICATIONS

None.

5 WARNINGS AND PRECAUTIONS

5.1 Hypersensitivity Reactions

Hypersensitivity reactions, including anaphylaxis, have been reported in STRENSIQtreated patients. Signs and symptoms consistent with anaphylaxis included difficulty breathing, choking sensation, nausea, periorbital edema, and dizziness. These reactions have occurred within minutes after subcutaneous administration of STRENSIQ and have been observed more than 1 year after treatment initiation. Other hypersensitivity reactions have also been reported in STRENSIQ-treated patients, including vomiting, fever, headache, flushing, irritability, chills, erythema, rash, pruritus and oral hypoesthesia [see Adverse Reactions (6.1)].

If a severe hypersensitivity reaction occurs, discontinue STRENSIQ treatment and initiate appropriate medical treatment. Consider the risks and benefits of re-administering STRENSIQ to individual patients following a severe reaction. If the decision is made to re-administer the product, monitor patients for a reoccurrence of signs and symptoms of a severe hypersensitivity reaction.

5.2 Lipodystrophy

Localized lipodystrophy, including lipoatrophy and lipohypertrophy, has been reported at injection sites after several months in patients treated with STRENSIQ in clinical trials *[see Adverse Reactions (6.1)]*. Advise patients to follow proper injection technique and to rotate injection sites *[see Dosage and Administration (2.4)]*.

5.3 Ectopic Calcifications

Patients with HPP are at increased risk for developing ectopic calcifications. Events of ectopic calcification, including ophthalmic (conjunctival and corneal) and renal (nephrocalcinosis, nephrolithiasis), have been reported in the clinical trial experience with STRENSIQ. There was insufficient information to determine whether or not the reported events were consistent with the disease or due to STRENSIQ. No visual changes or changes in renal function were reported resulting from the occurrence of ectopic calcifications.

Ophthalmology examinations and renal ultrasounds are recommended at baseline and periodically during treatment with STRENSIQ to monitor for signs and symptoms of ophthalmic and renal ectopic calcifications and for changes in vision or renal function.

5.4 Possible Immune-Mediated Clinical Effects

In clinical trials, most STRENSIQ-treated patients developed anti-asfotase alfa antibodies and neutralizing antibodies which resulted in reduced systemic exposure of asfotase alfa *[see Immunogenicity (6.2)]*. In postmarketing reports, some STRENSIQ-treated patients

with initial therapeutic response subsequently developed recurrence and worsening in disease-associated laboratory and radiographic biomarkers (some in association with neutralizing antibodies) suggesting possible immune-mediated effects on STRENSIQ's pharmacologic action resulting in disease progression *[see Adverse Reactions (6.3)]*. The effect of anti-asfotase alfa antibody formation on the long-term efficacy of STRENSIQ is unknown. There are no marketed anti-asfotase alfa antibody tests. If patients experience progression of HPP symptoms or worsening of disease-associated laboratory and imaging biomarkers after a period of initial therapeutic response to STRENSIQ, consider obtaining anti-asfotase alfa antibody testing by contacting STRENSIQ Medical Information at Alexion at 1-888-765-4747 or by email at medinfo@alexion.com. Close clinical follow up is recommended.

6 ADVERSE REACTIONS

The following adverse reactions are described below and elsewhere in the labeling:

- Hypersensitivity Reactions [see Warnings and Precautions (5.1)]
- Lipodystrophy [see Warnings and Precautions (5.2)]
- Ectopic Calcifications [see Warnings and Precautions (5.3)]
- Possible Immune-Mediated Clinical Effects [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 rates in the clinical trials of another drug and may not reflect the rates observed in practice.

The data described below reflect exposure to STRENSIQ in 99 patients with perinatal/infantile- or juvenile onset HPP (age 1 day to 58 years) treated with STRENSIQ, most for more than 2 years (range 1 day to 312 weeks [78 months]): 51 patients received at least 96 weeks (24 months) of treatment and 39 patients received 168 weeks (42 months) or more of treatment *[see Clinical Studies (14)]*.

Common Adverse Reactions

Overall, the most common adverse reactions reported were injection site reactions (63%). Other common adverse reactions included lipodystrophy (28%), ectopic calcifications (14%), and hypersensitivity reactions (12%).

Table 4 summarizes the adverse reactions that occurred at a rate of at least 10% in clinical trials following subcutaneous injection of STRENSIQ by patient population and STRENSIQ dosage regimen.

The frequency of injection site reactions, lipodystrophy and ectopic calcification were higher in patients with juvenile-onset HPP as compared to perinatal/infantile-onset HPP patients.

The majority of injection site reactions resolved within a week. Two patients experienced injection site reactions that led to reductions of their STRENSIQ dose. One patient switched from six times per week dosing to 3 times per week dosing as a result of injection site reactions. One other patient experienced a severe injection site reaction of injection site discoloration and withdrew from the trial.

		Perinatal/		1
	In	Juvenile- onset HPP		
Adverse Reaction Category or Term	6 mg/kg ner 6 mg/kg/weg		Total (N=79) n (%)	STRENSIQ (N=20) n (%)
Injection site reactions	38 (58)	6 (46)	44 (56)	18 (90)
Erythema	29 (44)	3 (23)	32 (41)	15 (75)
Discoloration/ Hypopigmentation	11 (17)	1 (8)	12 (15)	8 (40)
Pain/ Tenderness	10 (15)	1 (8)	11 (14)	8 (40)
Pruritus/ Itching	10 (15)	0 (0)	10 (13)	7 (35)
Swelling	8 (12)	0 (0)	8 (10)	6 (30)
Induration	9 (14)	1 (8)	10 (13)	3 (15)
Macule	4 (6)	0 (0)	4 (5)	7 (35)
Reaction, not otherwise specified	6 (9)	1 (8)	7 (9)	4 (20)
Bruising	6 (9)	0 (0)	6 (8)	4 (20)
Nodule	2 (3)	0 (0)	2 (3)	2 (10)
Other injection site reactions ^b	10 (15)	3 (23)	13 (17)	4 (20)
Ectopic calcifications	3 (5)	0 (0)	3 (4)	11 (55)
Lipodystrophy	12 (18)	2 (15)	14 (18)	14 (70)
Injection site atrophy	4 (6)	2 (15)	6 (8)	8 (40)
Injection site	5 (8)	0 (0)	5 (6)	6 (30)
hypertrophy		0.423		
Other lipodystrophy ^c	4 (6)	0 (0)	4 (5)	1 (5)
Hypersensitivity reactions	7 (11)	3 (23)	10 (13)	2 (10)
Vomiting/emesis	2 (3)	2 (15)	4 (5)	2 (10)
Other hypersensitivity reactions ^d	6 (9)	2 (15)	8 (10)	2 (10)

Table 4:Adverse Reactions Reported in at Least 10% of Patients with
Perinatal/Infantile- or Juvenile-onset HPP Enrolled in STRENSIQ
Clinical Trials

^a Adverse reactions are from the combined period of 6 mg/kg and above (i.e. total drug exposure regardless of starting dose and intermediary doses as long as the patient reached doses > 6 mg/kg)

^b Other injection site reactions include injection site rash, inflammation, papule, hemorrhage, hematoma, urticaria, warmth, calcification, mass, scar and cellulitis.

^c Other lipodystrophy includes lipohypertrophy.

^d Other hypersensitivity reactions include erythema/redness, pyrexia/fever, irritability, nausea, pain, rigor/chills, hypoesthesia oral, headache, flushing, and anaphylaxis.

Less Common Adverse Reactions

Adverse reactions that occurred at rates less than 1% included:

- Hypocalcemia
- Renal Stones
- Chronic hepatitis
- Decreased vitamin B6

Long-Term Safety

In long-term extension trials reflecting a median exposure to STRENSIQ of 142 weeks (range 0.1 weeks to 392 weeks) in 112 patients with perinatal/infantile- (n = 89), juvenile- (n = 22), and adult-onset (n = 1) HPP (age at enrolment = 1 day to 66.5 years), the most common adverse reactions were similar to those reported in Table 4.

6.2 Immunogenicity

As with all therapeutic proteins, there is potential for 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 the antibodies in the studies described below with the incidence of antibodies in other studies or to other asfotase alpha products may be misleading.

During clinical trials, anti-asfotase alfa antibodies have been detected in patients receiving treatment with STRENSIQ using an electrochemiluminescent (ECL) immunoassay. Antibody positive samples were tested to determine the presence of neutralizing antibodies based on in vitro inhibition of the catalytic activity of STRENSIQ.

Among STRENSIQ-treated patients with hypophosphatasia (HPP) in clinical studies who had post-baseline antibody data available, 97/109 (89%) tested positive for anti-asfotase alfa antibodies at some time point during STRENSIQ treatment. Among those 97 patients, 55 (57%) also tested positive for neutralizing antibodies at some time point during STRENSIQ treatment. No correlation was observed between the anti-asfotase alfa antibody titers and the neutralizing antibody (% inhibition) values. Formation of anti-asfotase alfa antibody resulted in a reduced systemic exposure of asfotase alfa *[see Clinical Pharmacology (12.3)]*.

6.3 **Postmarketing Experience**

The following adverse reactions have been identified during post-approval use of STRENSIQ. 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.

Some STRENSIQ-treated patients with initial therapeutic response to STRENSIQ subsequently developed worsening in disease-associated laboratory and radiographic biomarkers (some in association with neutralizing antibodies) suggesting possible immune-mediated effects on STRENSIQ's pharmacologic action resulting in disease progression [see Warnings and Precautions (5.4)].

7 DRUG INTERACTIONS

7.1 Drug Interference with Laboratory Tests

Laboratory Tests Utilizing Alkaline Phosphatase as a Detection Reagent

Studies have shown that there is analytical interference between asfotase alfa and laboratory tests that utilize an alkaline phosphatase (ALP)-conjugated test system, rendering erroneous test results in patients treated with STRENSIQ. ALP-conjugated test systems are utilized to measure substances such as hormones, bacterial antigens and antibodies. Therefore, it is recommended that laboratory assays which do not have ALP-conjugate technology be used when testing samples from patients who are receiving STRENSIQ.

To avoid erroneous test results for patients treated with STRENSIQ, inform laboratory personnel that the patient is being treated with STRENSIQ and discuss the use of a testing platform which does not utilize an ALP-conjugated test system.

Serum Alkaline Phosphatase

High serum ALP measurements detected through clinical laboratory testing are expected in patients receiving STRENSIQ and reflect circulating concentrations of asfotase alfa.

Do not rely on serum ALP measurements for clinical decision making in patients treated with STRENSIQ.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

There are no available data on STRENSIQ use in pregnant women to inform a drug associated risk. In animal reproduction studies, asfotase alfa administered intravenously to pregnant rats and rabbits during the period of organogenesis showed no evidence of fetotoxicity, embryolethality or teratogenicity at doses causing plasma exposures up to 21 and 24 times, respectively, the exposure at the recommended human dose (*see Data*).

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

Data

Animal Data

Asfotase alfa administered during the period of organogenesis to rats (from gestation Day 6 to Day 19 post-partum) and rabbits (on gestation days 7 to 19) at intravenous doses up to 50 mg/kg/day, approximately 21 and 24 times the human AUC of 65486 ng.h/mL at 2 mg/kg dose administered three times weekly for a 50 kg individual, respectively did not cause any adverse effects on embryofetal development. A pre- and post-natal development study in pregnant rats showed no evidence of adverse effects on pre- and post-natal development at intravenous doses (from Day 6 of gestation to Day 19 postpartum) of asfotase alfa up to 50 mg/kg/day approximately 21 times the human AUC of 65486 ng.h/mL at 2 mg/kg dose administered three times weekly for a 50 kg individual.

8.2 Lactation

Risk Summary

There are no data on the presence of asfotase alfa 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 STRENSIQ and any potential adverse effects on the breastfed infant from asfotase alfa or from the underlying maternal condition.

8.4 Pediatric Use

The safety and effectiveness of STRENSIQ for the treatment of perinatal/infantile- and juvenile-onset HPP have been established in pediatric patients. Use of STRENSIQ for this indication is based on 4 prospective, open-label clinical trials conducted in 89 pediatric patients with perinatal/infantile-onset or juvenile-onset HPP [see Clinical Studies (14)].

8.5 Geriatric Use

No patients with perinatal/infantile- or juvenile-onset HPP aged 65 years were enrolled in clinical trials of STRENSIQ. Therefore, there is no information available to determine whether patients aged 65 years and over respond differently from younger patients.

11 DESCRIPTION

Asfotase alfa is a tissue nonspecific alkaline phosphatase (TNSALP) produced by recombinant DNA technology in a Chinese hamster ovary cell line. Asfotase alfa is a soluble glycoprotein composed of two identical polypeptide chains. Each chain contains 726 amino acids with a theoretical mass of 161 kDa. Each chain consists of the catalytic domain of human TNSALP, the human immunoglobulin G₁ Fc domain and a decaaspartate peptide used as a bone targeting domain. The two polypeptide chains are covalently linked by two disulfide bonds.

STRENSIQ (asfotase alfa) injection is a sterile, preservative-free, nonpyrogenic, clear, slightly opalescent or opalescent, colorless to slightly yellow, with few small translucent or white particles, aqueous solution for subcutaneous administration. STRENSIQ is supplied in glass single-dose vials containing asfotase alfa; dibasic sodium phosphate, heptahydrate; monobasic sodium phosphate, monohydrate; and sodium chloride at a pH between 7.2 and 7.6. Table 5 describes the content of STRENSIQ vial presentations.

Ingredient	Quantity per Vial					
Asfotase Alfa	18 mg/0.45 mL	28 mg/0.7 mL	40 mg/mL	80 mg/0.8 mL		
Dibasic sodium phosphate, heptahydrate	2.48 mg	3.85 mg	5.5 mg	4.4 mg		
Monobasic sodium phosphate, monohydrate	0.28 mg	0.43 mg	0.62 mg	0.5 mg		
Sodium chloride	3.94 mg	6.13 mg	8.76 mg	7.01 mg		

Table 5: Content of STRENSIQ Vial Presentations

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

HPP is caused by a deficiency in TNSALP enzyme activity, which leads to elevations in several TNSALP substrates, including inorganic pyrophosphate (PPi). TNSALP is a metallo-enzyme that catalyzes the hydrolysis of phosphomonoesters with release of inorganic phosphate and alcohol. Elevated extracellular levels of PPi block hydroxyapatite crystal growth which inhibits bone mineralization and causes an accumulation of unmineralized bone matrix which manifests as rickets and bone deformation in infants and children and as osteomalacia (softening of bones) once growth plates close, along with muscle weakness. Replacement of the TNSALP enzyme upon STRENSIQ treatment reduces the enzyme substrate levels.

12.2 Pharmacodynamics

Perinatal/infantile- and juvenile-onset HPP patients treated with STRENSIQ had reductions in plasma TNSALP substrates, PPi and pyridoxal 5'-phosphate (PLP) within 6 to 12 weeks of treatment. Reductions in plasma PPi and PLP levels did not correlate with clinical outcomes.

Bone biopsy data from perinatal/infantile-onset and juvenile-onset HPP patients treated with STRENSIQ demonstrated decreases in osteoid volume and thickness indicating improved bone mineralization.

12.3 Pharmacokinetics

Based on data in 38 HPP patients, the pharmacokinetics of asfotase alfa exhibit dose proportionality across the dose range of 0.3 mg/kg to 3 mg/kg once every other day (three times a week) and appear to be time-independent. Steady state exposure was achieved as

early as three weeks after the administration of the first dose. The elimination half-life following subcutaneous administration was approximately 5 days.

Table 6 summarizes the pharmacokinetic parameters following multiple doses in 20 HPP patients after subcutaneous administration of STRENSIQ at 2 mg/kg three times per week in Study 2 (age of less than or equal to 5 years) and Study 3 (age of greater than 5 to 12 years), indicating the pharmacokinetics were similar between patients in the two age groups.

	Study 2	Study 3		
Ν	14	6		
Age (year)	3.4 ± 2.1 (0.2, 6.2)	8.6 ± 2.2 (6.1, 12.6)		
Weight at baseline (kg)	$11.2 \pm 5.0 \\ (2.9, 17.1)$	$21.2 \pm 7.9 \\ (11.4, 35.4)$		
t _{last} (h)	$48.1 \pm 0.1 \\ (47.9, 48.3)$	$\begin{array}{c} 48.0 \pm 0.1 \\ (48.0, 48.1) \end{array}$		
t _{max} (h)	$14.9 \pm 10.4 \\ (0, 32.2)$	$20.8 \pm 10.0 \\ (11.9, 32.2)$		
C _{max} (ng/mL)	$1794 \pm 690 \\ (856, 3510)$	$\begin{array}{c} 2108 \pm 788 \\ (905, 3390) \end{array}$		
AUC _t (h*ng/mL)	66042 ± 25758 (27770, 119122)	89877 ± 33248 (37364, 142265)		
Accumulation Ratio ^a	1.5	3.9		

Table 6:Summary of Pharmacokinetic Parameters Following Multiple
Subcutaneous Administration of STRENSIQ 2 mg/kg Three Times
per Week

 a Ratio values reflect the fold increase of AUCt from Week 1 based on mean AUCt, values.

Data are presented as mean \pm standard deviation (range). Study 3 includes patients with perinatal/infantile- or juvenile-onset of disease. $t_{last},$ time of last

concentration; t_{max} , time of maximal concentration; C_{max} , maximal concentration;

AUCt, area under the concentration-time curves over a dosing interval of 48 hours

Population PK analysis of asfotase alfa concentrations supports weight-based dosing because body weight is a major covariate of asfotase alfa clearance. The formulation concentration had an impact on the systemic exposure of asfotase alfa in HPP patients. The higher concentration formulation (80 mg/0.8 mL vial) achieved an approximately 25% lower systemic asfotase alfa exposure (i.e., concentrations and AUC) compared to the lower concentration formulations (18 mg/0.45 mL, 28 mg/0.7 mL or 40 mg/mL vials) at the same dose of STRENSIQ [see Dosage and Administration (2.1)].

Anti-Drug Antibody Effects on Pharmacokinetics

Formation of anti-drug antibodies resulted in reduced systemic exposure of asfotase alfa.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Long-term studies in animals to evaluate carcinogenic potential or studies to evaluate mutagenic potential have not been performed with asfotase alfa. Asfotase alfa at intravenous doses up to 50 mg/kg/day administered daily in pregnant rats (approximately 21 times the human AUC of 65486 ng.h/mL at 2 mg/kg dose administered three times weekly for a 50 kg individual) was found to have no adverse effect on fertility and reproductive performance of male and female rats.

14 CLINICAL STUDIES

14.1 Perinatal/Infantile-Onset HPP

Study 1 was a 24-week prospective single-arm trial in 11 patients with severe perinatal/infantile-onset HPP. In this study, 7/11 (64%) were female and 10/11 (91%) patients were white, and age ranged from 3 weeks to 39.5 months. Severe perinatal/infantile-onset HPP was defined as biochemical, medical history and radiographic evidence of HPP as well as the presence of any of the following: rachitic chest deformity, vitamin B6-dependent seizures, or failure to thrive. Ten of 11 patients completed the 24-week trial and continued treatment in the extension phase. Nine patients have been treated for at least 216 weeks (54 months) and 4 patients have been treated for over 240 weeks (60 months). Patients received subcutaneous STRENSIQ 3 mg/kg per week for the first month; subsequently, dose increases up to 9 mg/kg per week were allowed for changes in weight and/or for lack of efficacy. All 10 patients required dose increases up to 6 mg/kg per week or higher; 9 patients increased between 4 and 24 weeks after starting treatment and 1 patient increased after 70 weeks due to suboptimal clinical response. One patient's dose was decreased from 9 mg/kg per week to 6 mg/kg per week based on PK data.

Study 2 was a prospective open-label study in 59 patients with perinatal/infantile-onset HPP. In this study, 32/59 (54%) were female, 46/59 (78%) were white, and age ranged from 1 day to 78 months. Patients received subcutaneous STRENSIQ at 6 mg/kg per week for the first 4 weeks. Ten patients received dose increases higher than 6 mg/kg per week due to suboptimal clinical response, with dose increases occurring between 8 and 24 weeks after starting treatment. The recommended dosage regimen of STRENSIQ for the treatment of perinatal/infantile-onset HPP is up to 9 mg/kg per week administered subcutaneously as 3 mg/kg three times per week [*see Dosage and Administration (2.1)*].

Forty-one patients were treated for at least 24 weeks (6 months) and 15 patients were treated for at least 96 weeks (24 months).

Survival and Ventilation-Free Survival in Perinatal/Infantile-Onset HPP

Survival and invasive ventilation-free survival were compared in STRENSIQ-treated patients (Studies 1 and 2) with a historical cohort of untreated patients with similar clinical characteristics (Table 7 and Figure 1).

Table 7:Survival and Invasive Ventilation-Free Survival in STRENSIQ-
Treated versus Historical Control Patients with Perinatal/ Infantile-
Onset HPP (Pooled Studies 1 and 2)

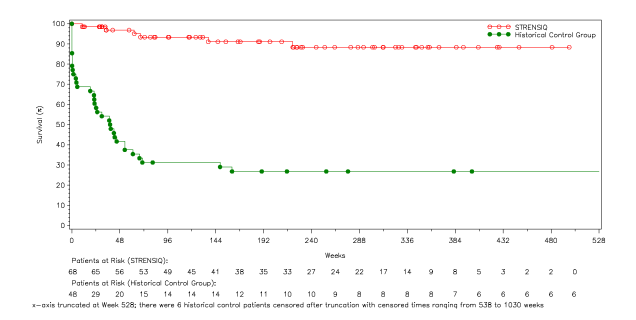
	STRENSIQ- Treated	Historical Controls
Survival	n = 68	n = 48
Alive at Point of Last Contact (%)	91	27
Hazard Ratio (STRENSIQ/Historical Control), 95% Confidence Interval*	0.1 (0.05,	
Kaplan-Meier Estimate and Alive at Age 1 Year (Week 48) (%)	97	42
Invasive Ventilation-Free Survival**	n = 54	n = 48
Alive and Not on Ventilation at Point of Last Contact (%)	85	25
Hazard Ratio (STRENSIQ/Historical Control), 95% Confidence Interval*	0.2 (0.09,	
Kaplan-Meier Estimate of Alive and Not on Ventilation at Age 1 Year (Week 48) (%)	96	31

* Adjusted for year of diagnosis.

** Alive and not initiating invasive ventilation after start of STRENSIQ treatment. STRENSIQ-treated patients on invasive ventilation at baseline were excluded from this analysis.

In patients who required any form of respiratory support, 21 of 26 (81%) of the treated patients survived through their last assessment (median age at last assessment was 3.2 years of age), versus 1 of 20 (5%) of historical controls.

Figure 1: Overall Survival in STRENSIQ-Treated versus Historical Control Patients with Perinatal/ Infantile-Onset HPP (Pooled Studies 1 and 2)



Skeletal Manifestations in Perinatal/Infantile-Onset HPP

Radiographs from 68 STRENSIQ-treated perinatal/infantile-onset HPP patients, including 64 patients in Studies 1 and 2, and 4 patients in Study 3 *[see Clinical Studies (14.2)]*, were examined to assess HPP-related rickets using the 7-point Radiographic Global Impression of Change (RGI-C) scale. Patients with a minimum RGI-C score of +2 were defined as "responders". Radiologic improvements could be seen by Month 24; at last assessment, 50/68 [74%] treated patients were rated as RGI-C responders. No comparative data were available from historical controls. The mean time interval between the baseline and last RGI-C assessment was 24 months (range was 1 month to 67 months).

Eighteen perinatal/infantile-onset HPP patients experienced fractures during the course of treatment. There were insufficient data to determine the effect of STRENSIQ on fractures.

Growth in Perinatal/Infantile-Onset HPP

Height and weight measurements (as measured by Z-scores) were available posttreatment for 72 perinatal/infantile-onset HPP patients, including 68 patients enrolled in Studies 1 and 2, and 4 patients enrolled in Study 3 (Table 8).

Table 8:Perinatal/Infantile-Onset Height and Weight Measurements as
Measured by Z-Score (Studies 1 and 2)

	Height Z-score					Weight	Z-score	
	Ba	aseline	Last Assessment		ment Baseline		Last Assessment	
	Mean	Min, Max	Mean	Min, Max	Mean	Min, Max	Mean	Min, Max
Studies 1 and 2* (N=68)	-3.3	-10.1, 0.9	-2.9	-10.6, 0.4	-3.2	-23.8, 0	-2.4	-20.9. 1.1
Study 3 (N=4)**	-2.6	-6.6, -0.7	-1.5	-5.8, 0.4	-2.5	-8.2, -1.0	-1.5	-5.4, 0.5

*The mean time interval between baseline and last assessment was 21 months (range was 1 month to 72 months).

**The mean time between baseline and last assessment was 56 months (range was 53 months to 60 months).

Long-Term Extension Trials in Perinatal/Infantile-Onset HPP

Long-term data were collected in 68 STRENSIQ-treated patients with perinatal/infantile onset HPP in Studies 1 and 2 and an additional 10 patients enrolled in Study 2. The longest duration of follow-up in the 78 patients was 7 years (84 months). At point of last contact, 69/78 (88%) STRENSIQ-treated patients had survived.

14.2 Juvenile-Onset HPP

Study 3 was a prospective open-label 24-week trial that included 8 juvenile-onset HPP patients and 5 perinatal/ infantile-onset HPP patients; 11/13 (85%) were male and 12/13 (92%) were white *[see Clinical Studies (14.1)]*. On study entry, patients were 6 to 12 years of age. All 8 juvenile-onset patients entered the extension study and were treated for at least 48 months. At trial entry, patients were randomized to receive subcutaneous STRENSIQ 6 mg/kg per week or 9 mg/kg per week. Two patients received dose reductions during the primary treatment period, including one patient who experienced a decrease in vitamin B6 levels and one patient who experienced recurrent injection site reactions. During the extension phase, the dosing regimen for all patients was initially changed to 3 mg/kg per week. Dosing was subsequently increased to 6 mg/kg per week, with no patients requiring doses higher than 6 mg/kg per week. The recommended dosage regimen of STRENSIQ for the treatment of juvenile-onset HPP is 6 mg/kg per week *[see Dosage and Administration (2.1)]*.

Growth in Juvenile-Onset HPP

Height and weight measurements (as measured by Z-scores) in 8 STRENSIQ-treated juvenile-onset HPP patients were compared with a historical cohort of 32 untreated patients with similar clinical characteristics (Table 9). Height and weight data for historical patients were collected from medical records.

Table 9:Juvenile-Onset Height and Weight Measurements as Measured by Z-
Score (Study 3)

	Height Z-score					Weight	Z-score	
Baseline		Baseline Last Assessment		Ba	aseline	Last A	ssessment	
	Mean	Min, Max	Mean	Min, Max	Mean	Min, Max	Mean	Min, Max
STRENSIQ (N=8)*	-1.5	-3.8, 0	-0.9	-2, 0	-1.1	-3.5, 2.3	0	-1.3, 2.2
Control (N=32)**	-1.1	-4.9, 2.6	-1.1	-4.9, 1.8	-1.2	-5, 2.1	-1	-5.7, 2.1

* The mean time interval between baseline and last assessment was 55 months (range was 53 months to 60 months).

** The mean time interval between baseline and last assessment was 61 months (range was 19 months to 109 months).

Skeletal Manifestations in Juvenile-Onset HPP

Radiographs from 8 STRENSIQ-treated juvenile-onset HPP patients and 32 historical controls were compared to assess HPP-related rickets using the 7-point RGI-C (Radiographic Global Impression of Change) scale. Patients who achieved a RGI-C score of 2 or higher (corresponding to substantial healing of rickets) were classified as being responders to treatment. All 8 treated patients were rated as responders by Month 54 of treatment. The mean duration between the baseline and last RGI-C assessments for control patients was 56 months (range was 8 to 95 months). At last assessment, 2/32 (6%) of control patients were rated as responders.

Eight of 20 (40%) patients with juvenile-onset HPP experienced new fractures during the course of treatment. There were insufficient data to assess the effect of STRENSIQ on fractures.

Gait/Mobility in Juvenile-Onset HPP

Gait was assessed using a modified Performance Oriented Mobility Assessment-Gait (MPOMA-G) scale in 8 STRENSIQ-treated juvenile-onset HPP patients at 6-month intervals out to 36 months. Mobility was also assessed using the 6 Minute Walk Test (6MWT) in 7 of the 8 patients. Step length improved by at least 1 point in either foot in 6/8 patients compared to 1/6 (17%) control patients. The proportion of patients who had 6MWT percent predicted values within the normal range for age, sex, and height-matched peers increased from 0/8 patients at baseline to 6/6 patients (100%) by Month 48 and all 6 were also able to walk longer distances at this time point compared to baseline.

Long-Term Extension Trials in Juvenile-Onset HPP

Long-term data were collected in 8 patients with juvenile-onset HPP treated with STRENSIQ for at least 6 years (72 months). At last assessment, 7 patients with available 6MWT results had maintained improvements in gait/mobility.

16 HOW SUPPLIED/STORAGE AND HANDLING

STRENSIQ is supplied as a sterile, nonpyrogenic, preservative-free, clear, slightly opalescent or opalescent, colorless to slightly yellow aqueous solution; a few small translucent or white particles may be present. The product is available as single-dose vials in a carton of one (1) or twelve (12) vials at the following strengths:

Strength	National Drug Code (NDC)	Quantity of Vials in Carton			
18 mg/0.45 mL	NDC 25682-010-01	1			
	NDC 25682-010-12	12			
28 mg/0.7 mL	NDC 25682-013-01	1			
	NDC 25682-013-12	12			
40 mg/mL	NDC-25682-016-01	1			
	NDC-25682-016-12	12			
For pediatric patients 40 kg and greater					
80 mg/0.8 mL	NDC 25682-019-01	1			

Table 10: STRENSIQ Vial Presentations

Strength	National Drug Code (NDC)	Quantity of Vials in Carton
	NDC 25682-019-12	12

STRENSIQ vials must be stored in the original carton until the time of use under refrigerated conditions at 2° C to 8° C (36° F to 46° F) and protected from light.

Once removed from refrigeration, STRENSIQ should be administered within 3 hours.

Do not use beyond the expiration date stamped on the carton.

DO NOT FREEZE OR SHAKE.

Vials are for one time use only. Discard any unused product.

17 PATIENT COUNSELING INFORMATION

Advise the patient or caregiver to read the FDA-approved patient labeling (Patient Information and Instructions for Use).

Advise patients or caregivers of the following:

Preparation

- When preparing a volume for injection greater than 1 mL, split the volume equally between two syringes, and administer two injections. When administering the two injections, use two separate injection sites.
- Take the unopened STRENSIQ vial(s) out of the refrigerator 15 to 30 minutes before injecting to allow the liquid to reach room temperature.
- Inspect the solution in the vial(s) for particulate matter and discoloration.
- Assemble injection supplies. Administer STRENSIQ using sterile disposable syringes and injection needles. The syringes should be of small enough volume that the prescribed dose can be withdrawn from the vial with reasonable accuracy. Always use a new syringe and needle.
- Remove vial cap, aseptically prepare the vial and insert the syringe into the vial to withdraw the prescribed dose for administration.
- Remove any air bubbles in the syringe and verify the correct dose.
- STRENSIQ vials are for one time use only. Discard any unused product [see Dosage and Administration (2.3)].

Administration

- Administer STRENSIQ within 3 hours upon removal of the vial(s) from refrigeration.
- Rotate the injection site to reduce the risk of lipohypertrophy and injection site atrophy.
- Do NOT administer injections in areas that are reddened, inflamed, or swollen.

• Inject STRENSIQ subcutaneously into the determined site and properly dispose of the syringe and needle [see Dosage and Administration (2.4)].

Hypersensitivity Reactions

• Reactions related to administration and injection may occur during and after STRENSIQ treatment. Inform patients and/or caregivers of the signs and symptoms of hypersensitivity reactions and have them seek immediate medical care should signs and symptoms occur [see Warnings and Precautions (5.1)].

Lipodystrophy

• Lipohypertrophy (enlargement or thickening of tissue) and localized atrophy (depression in the skin) have been reported at injection sites after several months. Follow proper injection technique and rotate injection sites [see Warnings and Precautions (5.2)].

Possible Immune-Mediated Clinical Effects

• Anti-drug antibodies may develop during treatment which may interfere with STRENSIQ's pharmacologic action. Inform patients or their caregivers to contact their healthcare provider if they experience worsening symptoms of HPP (e.g., increased respiratory support, increased difficulty walking, new fractures) [see Warnings and Precautions (5.4)].

Hypophosphatasia (HPP) Registry

• A registry has been established in order to better understand HPP in the population, and to monitor and evaluate long-term treatment effects of STRENSIQ. Patients and their caregivers should be encouraged to participate and advised that their participation is voluntary and may involve long-term follow-up. For more information, visit <u>www.hppregistry.com</u>

STRENSIQ is manufactured by:

Alexion Pharmaceuticals, Inc.

121 Seaport Boulevard

Boston, MA 02210

U.S. License Number: 1743

This label may not be the			
For current labeling information, pleas			
Patient Information	the nearest hospital emergency room right away if you have any of the sizes and sumptoms of a serieus allergic reaction including.		
STDENCIO® [strop! sik]	signs and symptoms of a serious allergic reaction including:		
STRENSIQ [®] [stren' sik]	 difficulty breathing nausea or vomiting 		
(asfotase alfa) injection, for subcutaneous use	 swelling of your eyes, lips dizziness 		
What is STRENSIQ?	or tongue o itching of lips, tongue or		
STRENSIQ is a prescription medicine used to treat people with perinatal, infantile,	 hives throat 		
and juvenile onset hypophosphatasia (HPP).	 feeling faint choking sensation 		
Before using STRENSIQ, tell your healthcare provider about all of your medical			
conditions, including if you:	• skin thickening or pits at the injection site (lipodystrophy). Lipodystrophy		
	at the injection site has happened several months after using STRENSIQ.		
are allergic to asfotase alfa or any of the ingredients in STRENSIQ. See the end	• calcium build up in your eyes and kidneys. You healthcare provider should		
of this leaflet for a complete list of ingredients in STRENSIQ.	check your eyes and kidneys while you use STRENSIQ.		
are pregnant or planning to become pregnant. It is not known if STRENSIQ will			
harm your unborn baby.	The most common side effects of STRENSIQ include local skin injection site		
are breastfeeding or plan to breastfeed. It is not known if STRENSIQ passes			
into your breast milk. Talk to your healthcare provider about the best way to feed your baby if you use STRENSIQ.	thinning, swelling, pits, and bumps) and calcium build up in your eyes and kidneys.		
 experience decreased efficacy with STRENSIQ. 	Nulleys.		
• experience decreased enreacy with strensig.	These are not all the possible side effects of STRENSIQ. For more information, ask		
Tell your healthcare provider about all the medications you take, including	your healthcare provider or pharmacist. Call your healthcare provider for medical		
prescription and over-the counter medicines, vitamins, and herbal supplements.	advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.		
р р	General information about the safe and effective use of STRENSIQ		
There is a registry for people who use STRENSIQ. The purpose of this registry is to	Medicines are sometimes prescribed for purposes other than those listed in a		
collect information about HPP and about what happens when you use STRENSIQ for	Patient Information leaflet. Do not use STRENSIQ for a condition for which it was		
a long time. For more information about this registry, talk with your healthcare	not prescribed. Do not give STRENSIQ to other people, even if they have the		
provider or go to www.hppregistry.com.	same symptoms that you have. It may harm them. You can ask your pharmacist		
How should I use STRENSIQ?	or healthcare provider for information about STRENSIQ that is written for		
See the "Instructions for Use" that come with STRENSIQ for detailed	healthcare professionals.		
instructions about the right way to use STRENSIQ.	What are the ingredients in STRENSIQ?		
Use STRENSIQ exactly as your healthcare provider tells you to.	Active ingredient: asfotase alfa.		
 Your healthcare provider will tell you how much STRENSIQ to use and when to use it. 	 Inactive ingredients: dibasic sodium phosphate, heptahydrate; monobasic sodium phosphate, monohydrate and sodium chloride. 		
Your healthcare provider may change your dose if needed.	Manufactured by: Alexion Pharmaceuticals, Inc., Boston, MA 02210 USA		
• Change (rotate) your injection site with each injection. Do not use the same	U.S. License Number: 1743		
injection site for each injection.	For more information, go to <u>www.strensiq.com</u> or call 888-765-4747.		
What are the possible side effects of STRENSIQ?	This Patient Information has been approved by the U.S. Food and Drug Administration		
STRENSIQ may cause serious side effects, including:	Approved: 10/2016		

 serious allergic (hypersensitivity) reactions. Serious allergic reactions have happened in some people who use STRENSIQ. Stop using STRENSIQ and

INSTRUCTIONS FOR USE

STRENSIQ[®] [stren' sik]

(asfotase alfa) injection, for subcutaneous use vial

Read this "Instructions for Use" before you start using STRENSIQ and each time you get a refill. There may be new information. This information does not take the place of talking to your healthcare provider about your medical condition or your treatment.

Do not share your syringes or needles with anyone else. You may give an infection to them or get an infection from them.

Supplies needed to give your STRENSIQ injection (See Figure A):

- 1 or 2 STRENSIQ vial(s).
- 1 or 2 sterile disposable 1 mL syringes for injection with 25 to 29 gauge (G), ½ inch needles.
 - The use of two different gauge needles is recommended, a larger bore needle (e.g. 25 gauge) for withdrawal of the medication, and a smaller bore needle (e.g. 29 gauge) for the injection.
 Always use a new syringe and needle for each injection.
- 2 alcohol wipes
- 1 gauze or cotton ball
- a clean flat surface, like a table
- 1 sharps container for throwing away used needles and syringes. See
 "Disposing of used needles and syringes" at the end of these instructions.

Figure A



Storing your STRENSIQ dose:

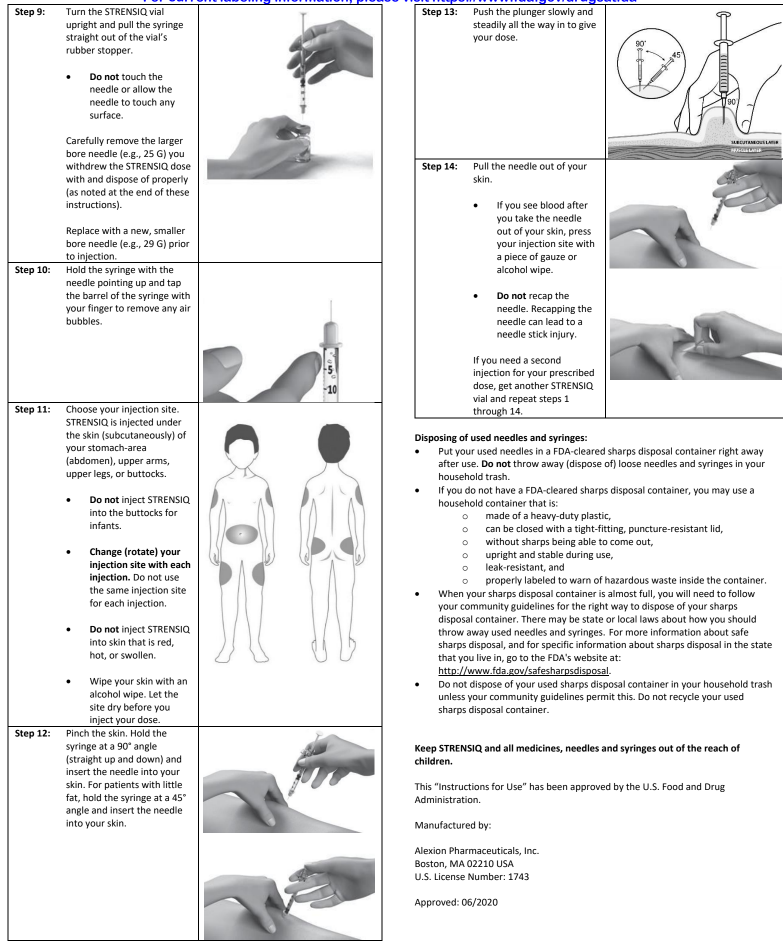
- Store STRENSIQ in the original carton in the refrigerator between 36°F to 46°F (2°C to 8°C) until you are ready to use it.
- Do not freeze your STRENSIQ vials. Do not use STRENSIQ if it has been frozen.
- Do not shake your STRENSIQ vials.
- Protect STRENSIQ from light until you are ready to use it.
- Do not use STRENSIQ after the expiration date printed on the carton.
- STRENSIQ vials are for 1 time use only. Throw away any unused STRENSIQ left in the vial.

Preparing your STRENSIQ dose and giving your STRENSIQ injection:

- Prepare a clean flat surface, like a table or counter top.
- Remove the unopened STRENSIQ vial(s) out of the refrigerator and allow it to sit at room temperature for at least 15 to 30 minutes. Injecting STRENSIQ when cold can make the injection feel uncomfortable.
 - Do not warm STRENSIQ in any way other than letting it sit at room temperature (for example, do not warm it in a microwave or in hot water).
- Gather all the supplies you will need to give your STRENSIQ injection.
- Wash your hands with soap and water.
- Utilize STRENSIQ within 3 hours after removing it from the refrigerator.
- Inject STRENSIQ exactly as your healthcare provider has shown you.

Step 1:	Check the liquid in the	
	STRENSIQ vial. STRENSIQ	
	liquid should look clear or	
	slightly yellow and may have	
	a few small white particles in	

	in the test of the second s	
	it. Do not use it if the liquid is	
	discolored or contains any	
	•	
	lumps or large particles in it.	
	Throw it away and get a new	
	vial.	
Step 2:	Using your thumb, flip the	
•	plastic cap off the STRENSIQ	and the second
	vial.	
		Rele
		in the second
Step 3:	Remove the larger bore	
Step 5.	-	
	needle (e.g., 25 G) from the	
	package. Pick up the syringe	
	and place the needle on the	
	tip of the syringe. Push down	
	and twist the needle onto	
	the syringe until it is tight.	
Step 4:	Hold the syringe with the	
	needle pointing up and pull	
	back the plunger until the	
	top of the plunger reaches	
	the line for your prescribed	
	dose.	
Step 5:	Pull the cap straight off of	_
	the needle.	
	-	-1
	• Do not touch the	
	needle or let the	The
	needle touch any	
	surface.	
Char: C		
Step 6:	Hold the STRENSIQ vial	Т
	firmly on a flat surface, then	
	push the needle through the	
	rubber stopper of the	
	STRENSIQ vial.	
		00 ⁵
		. 4
		45
		A
Step 7:	Keeping the needle in the	
	STRENSIQ vial, lift the vial	
	- /	
	and turn it upside down with	
	the needle pointing toward	
	the ceiling. Slowly push the	
	plunger all the way in.	
Stor 0.	With the needle tip still in	
Step 8:	•	- here
	the liquid, slowly pull back	
	the plunger until the top of	
	the plunger reaches the line	
		8
	slightly past the line for your	
	slightly past the line for your prescribed dose. Do not pull	
	slightly past the line for your prescribed dose. Do not pull the needle out of the	
	slightly past the line for your prescribed dose. Do not pull	
	slightly past the line for your prescribed dose. Do not pull the needle out of the STRENSIQ vial. Slowly push	
	slightly past the line for your prescribed dose. Do not pull the needle out of the STRENSIQ vial. Slowly push the plunger up until the top	
	slightly past the line for your prescribed dose. Do not pull the needle out of the STRENSIQ vial. Slowly push the plunger up until the top of the plunger reaches the	
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	slightly past the line for your prescribed dose. Do not pull the needle out of the STRENSIO vial. Slowly push the plunger up until the top of the plunger reaches the line for your prescribed dose.	



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