

**CENTER FOR DRUG EVALUATION AND  
RESEARCH**

***APPLICATION NUMBER:***

**761052Orig1s000**

**LABELING**

## HIGHLIGHTS OF PRESCRIBING INFORMATION

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

**BRINEURA (cerliponase alfa) injection, for intraventricular use**  
**Initial U.S. Approval: 2017**

### INDICATIONS AND USAGE

Brineura is a hydrolytic lysosomal N-terminal tripeptidyl peptidase indicated to slow the loss of ambulation in symptomatic pediatric patients 3 years of age and older with late infantile neuronal ceroid lipofuscinosis type 2 (CLN2), also known as tripeptidyl peptidase 1 (TPP1) deficiency. (1)

### DOSAGE AND ADMINISTRATION

- Aseptic technique must be strictly observed during preparation and administration. Brineura should be administered by, or under the direction of a physician knowledgeable in intraventricular administration. Brineura is administered to the cerebrospinal fluid (CSF) by infusion via a surgically implanted reservoir and catheter. (2.1)
- Pre-treatment of patients with antihistamines with or without antipyretics or corticosteroids is recommended 30 to 60 minutes prior to the start of infusion. (2.2)
- The recommended dosage is 300 mg administered once every other week as an intraventricular infusion followed by infusion of Intraventricular Electrolytes over approximately 4.5 hours. (2.2)
- For complete information on preparation, specific intraventricular access device for use, and administration, see the full prescribing information. (2.1, 2.3, 2.4, 2.5)

### DOSAGE FORMS AND STRENGTHS

Injection: Brineura 150 mg/5 mL (30 mg/mL) solution, two single-dose vials per carton co-packaged with Intraventricular Electrolytes Injection 5 mL in a single-dose vial. (3)

### CONTRAINDICATIONS

- Acute intraventricular access device-related complications (e.g., leakage, device failure, or device-related infection). (4)
- Patients with ventriculoperitoneal shunts. (4)

### WARNINGS AND PRECAUTIONS

- Intraventricular Access Device-Related Complications: Inspect the scalp for skin integrity and for signs of intraventricular access device leakage. Do not administer if there are signs of device leakage or infection. Routinely send CSF samples for testing to detect subclinical device-related infections. (2.5, 5.1)
- Cardiovascular Adverse Reactions: Monitor vital signs before, during, and post-infusion. Monitor Electrocardiogram (ECG) in patients with a history of bradycardia, conduction disorder, or with structural heart disease, during the infusion. In patients without cardiac abnormalities, perform regular 12-lead ECG evaluations every 6 months. (2.5, 5.2)
- Hypersensitivity Reactions: Observe patients during and after the infusion. If a severe hypersensitivity reaction occurs, immediately stop the infusion and initiate appropriate treatment. (5.3)

### ADVERSE REACTIONS

Most common adverse reactions ( $\geq 8\%$ ) are: pyrexia, ECG abnormalities, decreased CSF protein, vomiting, seizures, hypersensitivity, increased CSF protein, hematoma, headache, irritability, pleocytosis, device-related infection, bradycardia, feeling jittery, and hypotension. (6.1)

**To report SUSPECTED ADVERSE REACTIONS, contact BioMarin at 1-866-906-6100 or FDA at 1-800-FDA-1088 or [www.fda.gov/medwatch](http://www.fda.gov/medwatch).**

**See 17 for PATIENT COUNSELING INFORMATION**

Revised: 04/2017

## FULL PRESCRIBING INFORMATION: CONTENTS\*

### 1 INDICATIONS AND USAGE

### 2 DOSAGE AND ADMINISTRATION

- 2.1 Important Preparation and Administration Information
- 2.2 Dosage
- 2.3 Method of Administration
- 2.4 Preparation for Infusion
- 2.5 Intraventricular Infusion Procedure

### 3 DOSAGE FORMS AND STRENGTHS

### 4 CONTRAINDICATIONS

### 5 WARNINGS AND PRECAUTIONS

- 5.1 Intraventricular Access Device-Related Complications
- 5.2 Cardiovascular Adverse Reactions
- 5.3 Hypersensitivity Reactions

### 6 ADVERSE REACTIONS

- 6.1 Clinical Trials Experience
- 6.2 Immunogenicity

### 8 USE IN SPECIFIC POPULATIONS

- 8.1 Pregnancy
- 8.2 Lactation
- 8.4 Pediatric Use

### 11 DESCRIPTION

### 12 CLINICAL PHARMACOLOGY

- 12.1 Mechanism of Action
- 12.3 Pharmacokinetics

### 13 NONCLINICAL TOXICOLOGY

- 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

### 14 CLINICAL STUDIES

### 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

Brineura is indicated to slow the loss of ambulation in symptomatic pediatric patients 3 years of age and older with late infantile neuronal ceroid lipofuscinosis type 2 (CLN2), also known as tripeptidyl peptidase 1 (TPP1) deficiency.

### 2 DOSAGE AND ADMINISTRATION

#### 2.1 Important Preparation and Administration Information

- Aseptic technique must be strictly observed during preparation and administration.
- Brineura should be administered by, or under the direction of a physician knowledgeable in intraventricular administration.
- Brineura is administered into the cerebrospinal fluid (CSF) by infusion via a surgically implanted reservoir and catheter (intraventricular access device). Brineura is intended to be administered via the Codman<sup>®</sup> HOLTER RICKHAM Reservoirs (Part Numbers: 82-1625, 82-1621, 82-1616) with the Codman<sup>®</sup> Ventricular Catheter (Part Number: 82-1650). The intraventricular access device must be implanted prior to the first infusion. It is recommended that the first dose be administered at least 5 to 7 days after device implantation.
- Brineura is intended to be administered with the B Braun Perfusor<sup>®</sup> Space Infusion Pump System. The essential performance requirements for this syringe pump used to deliver Brineura are as follows:
  - Delivery rate of 2.5 mL/hr with delivery accuracy of +/- 1 mL/hr
  - Compatible with 20 mL syringes provided in the Administration Kit for use with Brineura
  - Occlusion alarm setting to  $\leq$  281 mm Hg
- Administer Brineura and the Intraventricular Electrolytes using the provided Administration Kit for use with Brineura components [*see How Supplied/Storage and Handling (16)*].

#### 2.2 Dosage

The recommended dosage of Brineura in pediatric patients 3 years of age and older is 300 mg administered once every other week by intraventricular infusion. Administer Brineura first followed by infusion of the Intraventricular Electrolytes each at an infusion rate of 2.5 mL/hr. The complete Brineura infusion, including the required infusion of Intraventricular Electrolytes, is approximately 4.5 hours.

Pre-treatment of patients with antihistamines with or without antipyretics or corticosteroids is recommended 30 to 60 minutes prior to the start of infusion.

#### 2.3 Method of Administration

Brineura and the Intraventricular Electrolytes must only be administered by the intraventricular route, using the provided Administration Kit for use with Brineura. Each vial of Brineura and Intraventricular Electrolytes is intended for a single dose only.

Each infusion consists of 10 mL of Brineura followed by 2 mL of Intraventricular Electrolytes. The complete infusion must be administered using an infusion set with a 0.2 micron inline filter. The Intraventricular Electrolytes are used to flush the infusion line, port needle, and intraventricular access device in order to fully administer Brineura and to maintain patency of the intraventricular access device.

## 2.4 Preparation for Infusion

*Gather supplies:*

- Brineura and Intraventricular Electrolytes Injection vials (package 1 of 2) [*see How Supplied/Storage and Handling (16)*]
- Administration Kit for use with Brineura (package 2 of 2) [*see How Supplied/Storage and Handling (16)*]
- Syringe pump (not supplied)

Inspect the Administration Kit infusion components to ensure the components are in the individual packages and have not been compromised.

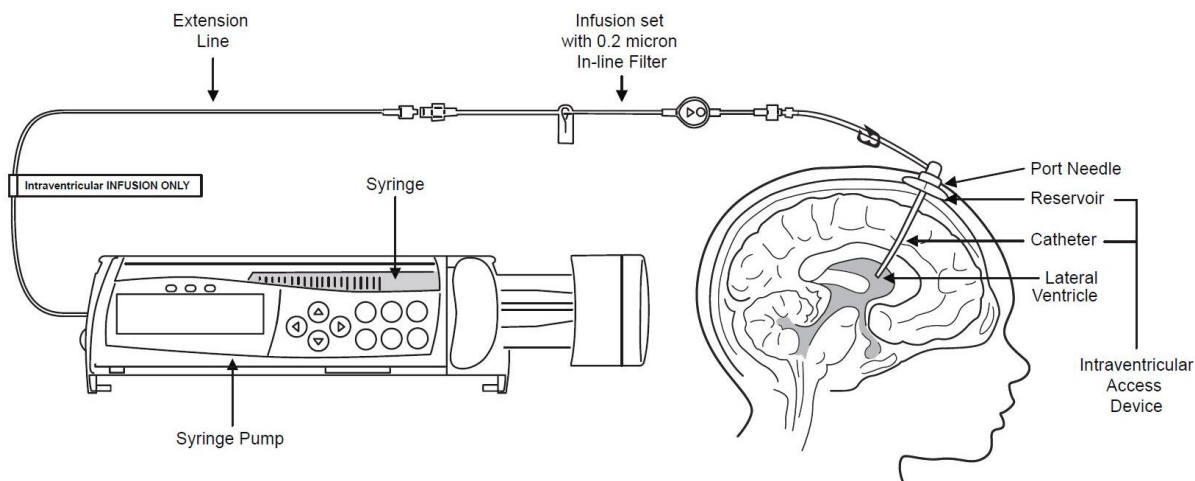
Thaw Brineura and Intraventricular Electrolytes Injection vials at room temperature for approximately 60 minutes. **Do not** thaw or warm vials any other way. **Do not** shake vials. Condensation will occur during thawing period. **Do not** re-freeze vials or freeze syringes containing Brineura or Intraventricular Electrolytes.

Inspect fully thawed Brineura and Intraventricular Electrolytes Injection vials. Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit. Brineura is a clear to slightly opalescent and colorless to pale yellow solution. Intraventricular Electrolytes is a clear to colorless solution. **Do not use** if the solutions are discolored or if there is other foreign particulate matter in the solutions. Brineura vials may occasionally contain thin translucent fibers or opaque particles. These naturally occurring particles are cerliponase alfa. These particles are removed via the 0.2 micron inline filter without having a detectable effect on the purity or strength of Brineura. Intraventricular Electrolytes may contain particles, which appear during the thaw period; however, these dissolve when the solution reaches room temperature.

## 2.5 Intraventricular Infusion Procedure

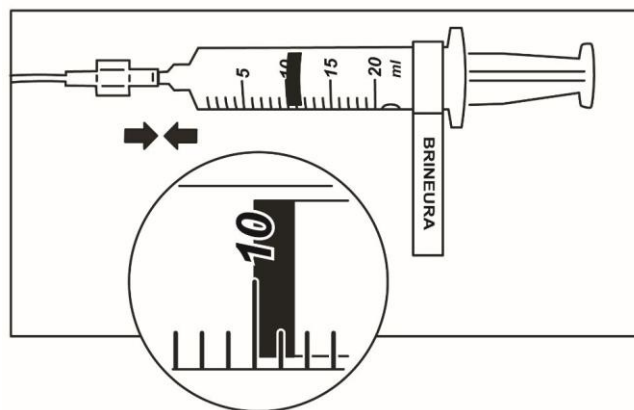
### Intraventricular Infusion of Brineura

Figure 1 represents the intraventricular infusion system set up. Use aseptic technique during the infusion. Follow the steps below to proceed with the intraventricular infusion.



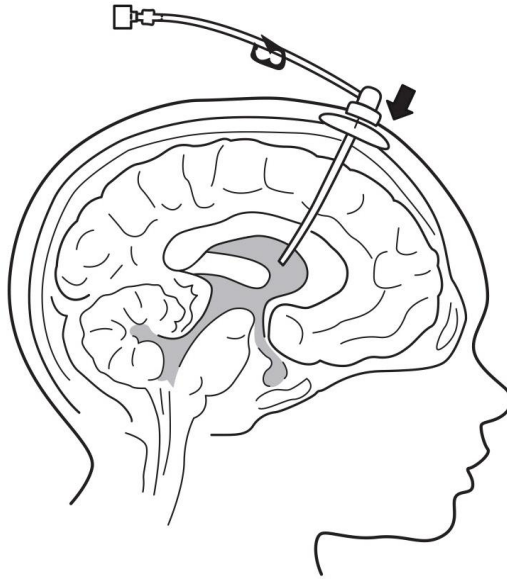
**Figure 1**

1. Use aseptic technique when preparing the Brineura syringe for infusion. Label one-sterile syringe “Brineura” and attach the syringe needle. Remove the green flip-off caps from the two Brineura vials. Use the “Brineura” labeled syringe to withdraw a total of 10 mL from the Brineura vials. **Do not** dilute Brineura. **Do not** mix Brineura with any other drug.
2. Label the infusion line “intraventricular infusion only” (see Figure 1).
3. Attach the syringe containing Brineura to the extension line (see Figure 2). Then connect the extension line to the infusion set with a 0.2 micron inline filter (see Figure 1).



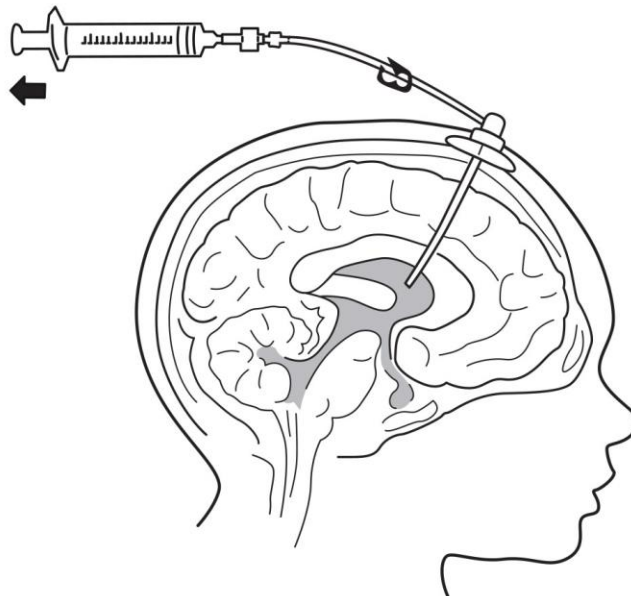
**Figure 2**

4. Prime the infusion components with Brineura.
5. Inspect scalp for signs of intraventricular access device leakage or failure and for potential infections [see *Warnings and Precautions* (5.1)].
6. Prepare the scalp for intraventricular infusion per institution standard of care.
7. Insert port needle into intraventricular access device (see Figure 3).



**Figure 3**

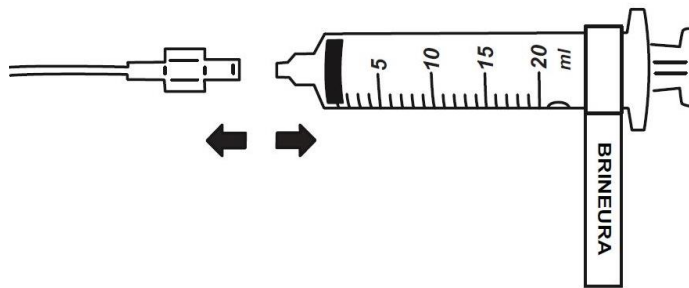
8. Connect a separate empty sterile single-use luer lock syringe, no larger than 3 mL (not provided) to the port needle. Withdraw 0.5 mL to 1 mL of CSF to check patency of intraventricular access device (see Figure 4) and send specimen for culture.
  - **Do not** return CSF to intraventricular access device.
  - Routinely send CSF samples for infection monitoring [see *Warnings and Precautions* (5.1)].



**Figure 4**

9. Attach the infusion set with 0.2 micron inline filter to the port needle (see Figure 1).
  - Secure the components per institution standard of care.

10. Place the syringe containing Brineura into the syringe pump and program pump to deliver at an infusion rate of 2.5 mL per hour. Set the occlusion alarm setting to alert at pressure  $\leq 281$  mm Hg. See syringe pump operating manual for details. **Do not** deliver as a bolus or manually.
11. Administer pre-medication 30 to 60 minutes prior to the start of infusion [see *Dosage and Administration* (2.2)].
12. Monitor vital signs (blood pressure, heart rate) prior to the start of infusion, periodically during infusion, and post-infusion [see *Warnings and Precautions* (5.2)].
13. Initiate infusion of Brineura at a rate of 2.5 mL per hour.
14. Periodically inspect the infusion system during the infusion for signs of leakage or delivery failure [see *Warnings and Precautions* (5.1)].
15. When the Brineura infusion is complete, detach and remove the empty syringe from the pump and disconnect from the tubing (see Figure 5). Proceed to Step 14 for Intraventricular Electrolytes infusion.

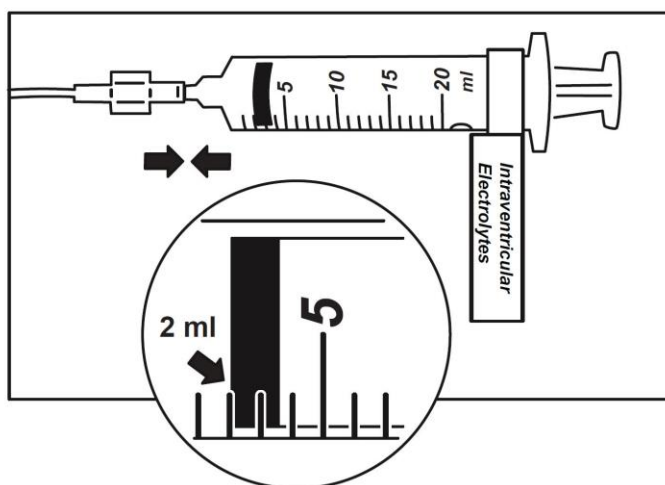


**Figure 5**

#### Intraventricular Infusion of Intraventricular Electrolytes

Administer the Intraventricular Electrolytes provided **after** Brineura infusion is complete.

16. Use aseptic technique when preparing the Intraventricular Electrolytes syringe for infusion. Label one sterile syringe "Intraventricular Electrolytes" and attach the syringe needle. Remove the yellow flip-off cap from the Intraventricular Electrolytes Injection vial. Withdraw 2 mL of Intraventricular Electrolytes. Discard the remaining unused portion.
17. Attach the syringe to the extension line (see Figure 6).



**Figure 6**

18. Place the syringe containing Intraventricular Electrolytes into the syringe pump and program pump to deliver at an infusion rate of 2.5 mL per hour. Set the occlusion alarm setting to alert at pressure  $\leq 281$  mm Hg. See syringe pump operating manual for details. **Do not** deliver as a bolus or manually.
19. Initiate infusion of Intraventricular Electrolytes at a rate of 2.5 mL per hour.
20. Periodically inspect the infusion system during the infusion for signs of leakage or delivery failure.
21. When the Intraventricular Electrolytes infusion is complete, detach and remove the empty syringe from the pump and disconnect from the infusion line.
22. Remove the port needle. Apply gentle pressure and bandage the infusion site per institution standard of care.

Dispose of the infusion components, needles, unused solutions and other waste materials in accordance with local requirements.

#### Storage of Thawed Product

Use thawed Brineura and Intraventricular Electrolytes immediately. If not used immediately, store unopened vials in the refrigerator at 2°C to 8°C and use within 24 hours.

#### Storage of Product in Syringes

Use product held in labeled syringes immediately. If not used immediately, store product held in labeled syringes in the refrigerator at 2°C to 8°C up to 4 hours prior to infusion.

### **3 DOSAGE FORMS AND STRENGTHS**

Injection: Brineura 150 mg/5 mL (30 mg/mL) solution, two single-dose vials per carton co-packaged with Intraventricular Electrolytes Injection 5 mL in a single-dose vial. Brineura is a clear to slightly opalescent and colorless to pale yellow solution. Intraventricular Electrolytes is a clear to colorless solution [see *How Supplied/Storage and Handling* (16)].

## 4 CONTRAINDICATIONS

Brineura is contraindicated in patients with:

- acute intraventricular access device-related complications (e.g., leakage, device failure, or device-related infection) [see *Warnings and Precautions (5.1)*].
- ventriculoperitoneal shunts.

## 5 WARNINGS AND PRECAUTIONS

### 5.1 Intraventricular Access Device-Related Complications

Brineura must be administered using aseptic technique to reduce the risk of infection. Healthcare professionals should inspect the scalp for skin integrity to ensure the intraventricular access device is not compromised prior to each infusion [see *Dosage and Administration (2.5)*].

Brineura is contraindicated if there are signs of acute intraventricular access device-related complications (e.g., leakage, device failure or signs of device-related infection such as swelling, erythema of the scalp, extravasation of fluid, or bulging of the scalp around or above the intraventricular access device) [see *Contraindications (4)*]. In case of intraventricular access device complications, discontinue the Brineura infusion and refer to the device manufacturer's labeling for further instructions.

The signs and symptoms of device-related infections may not be apparent, therefore, CSF samples should routinely be sent for testing to detect subclinical device infections [see *Dosage and Administration (2.5)*].

In clinical studies with Brineura, intraventricular access device-related infections were observed in two patients. In each case, antibiotics were administered, the intraventricular access device was replaced, and the patient continued on Brineura treatment.

Material degradation of the intraventricular access device reservoir may occur after approximately 105 perforations of the intraventricular access device. The intraventricular access device may require replacement as soon as, or prior to, 105 administrations of Brineura, equating to approximately 4.3 years of regular administrations.

### 5.2 Cardiovascular Adverse Reactions

Monitor vital signs (blood pressure, heart rate) before infusion starts, periodically during infusion, and post-infusion in a healthcare setting [see *Dosage and Administration (2.5)*]. Upon completion of the infusion, clinically assess the patient status. Continued observation may be necessary if clinically indicated.

Perform electrocardiogram (ECG) monitoring during infusion in patients with a history of bradycardia, conduction disorder, or with structural heart disease, as some patients with CLN2 disease may develop conduction disorders or heart disease. In patients without cardiac abnormalities, regular 12-lead ECG evaluations should be performed every 6 months.

In the clinical studies, hypotension was reported in 2 (8%) patients, which occurred during or up to eight hours after Brineura infusion. Patients did not require alteration in treatment, and reactions resolved spontaneously or after intravenous fluid administration [see *Adverse Reactions (6.1)*].

### 5.3 Hypersensitivity Reactions

Hypersensitivity reactions have been reported in Brineura-treated patients during the clinical studies. A total of 11 (46%) patients experienced hypersensitivity reactions during the infusion or within 24 hours of

completion of the infusion. The signs and symptoms observed concomitantly with hypersensitivity reactions included pyrexia, vomiting, pleocytosis or irritability. Patients were routinely pre-medicated with antihistamines with or without antipyretics or corticosteroids, prior to infusion of Brineura.

Due to the potential for anaphylaxis, appropriate medical support should be readily available when Brineura is administered. If anaphylaxis occurs, immediately discontinue the infusion and initiate appropriate medical treatment. Observe patients closely during and after the infusion. Inform patients/caregivers of the signs and symptoms of anaphylaxis, and instruct them to seek immediate medical care should signs and symptoms occur.

The management of hypersensitivity reactions should be based on the severity of the reaction and may include temporarily interrupting the infusion, and/or treatment with antihistamines, antipyretics, and/or corticosteroids. If a severe hypersensitivity reaction occurs, immediately discontinue the infusion and initiate appropriate medical treatment.

## **6 ADVERSE REACTIONS**

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

- Intraventricular Access Device-Related Complications *[see Warnings and Precautions (5.1)]*
- Cardiovascular Adverse Reactions *[see Warnings and Precautions (5.2)]*
- Hypersensitivity Reactions *[see Warnings and Precautions (5.3)]*

### **6.1 Clinical Trials Experience**

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

The safety of Brineura was evaluated in 24 patients with CLN2 disease who received at least one dose of Brineura in a clinical study with extension of up to 161 weeks *[see Clinical Studies (14)]*. Table 1 summarizes the most common adverse reactions that occurred in Brineura-treated patients through 96 weeks.

**Table 1: Adverse Reactions Reported in  $\geq 8\%$  of Symptomatic Pediatric Patients with CLN2 Disease in the Brineura Single-Arm Clinical Study with Extension at Week 96**

| Adverse Reaction          | Patients Treated with Brineura<br>n=24 (%) |
|---------------------------|--------------------------------------------|
| Pyrexia*                  | 17 (71)                                    |
| ECG abnormalities†        | 17 (71)                                    |
| Decreased CSF protein     | 17 (71)                                    |
| Vomiting                  | 15 (63)                                    |
| Seizures‡                 | 12 (50)                                    |
| Hypersensitivity§         | 11 (46)                                    |
| Increased CSF protein     | 5 (21)                                     |
| Hematoma                  | 5 (21)                                     |
| Headache                  | 4 (17)                                     |
| Irritability              | 4 (17)                                     |
| Pleocytosis               | 4 (17)                                     |
| Device-related infection¶ | 2 (8)                                      |
| Bradycardia               | 2 (8)                                      |
| Feeling jittery           | 2 (8)                                      |
| Hypotension               | 2 (8)                                      |

\* Pyrexia includes: pyrexia and increased body temperature

† ECG abnormalities include: non-specific repolarization abnormality, notched QRS, ST segment elevation, biphasic T wave abnormality, supraventricular extrasystoles, bradycardia, sinus tachycardia, and intraventricular conduction delay

‡ Seizures include: atonic, generalized tonic-clonic, focal, and absence.

§ Hypersensitivity includes: immune reactions and signs and symptoms observed concomitantly with hypersensitivity reactions including pyrexia, vomiting, pleocytosis or irritability

¶ Device-related infections include: *Propionibacterium acnes* and *Staphylococcus epidermidis*

### Description of Selected Adverse Reactions

#### *Seizures*

Seizures were reported in 12 of 24 (50%) patients. The seizure types reported include atonic, generalized tonic-clonic, focal, and absence. Seizures were managed with standard anti-convulsive therapies and did not result in discontinuation of Brineura treatment.

#### *Device-Related Complications*

Adverse reactions related to the device were observed in 12 of 24 (50%) of patients. Device-related adverse reactions include infection, delivery system-related complications, and pleocytosis. Nine of these patients (38%) experienced adverse reactions, which involved complications of the non-implanted delivery system components. Four patients (16%) had device-related adverse reactions, which required medical intervention, including two patients (8%) with intraventricular access device-related CNS infections, and one patient (4%) each with leakage of the intraventricular access device and pleocytosis. Device-related infections were diagnosed by increased CSF pleocytosis and microbiology culture and organism identification, without accompanying signs and symptoms of meningitis. Intraventricular access devices were replaced and infections were treated with antibiotics. Device-related complications did not result in discontinuation of Brineura treatment [see *Warnings and Precautions (5.1)*].

#### *Hematoma*

Hematoma adverse reactions were reported in 5 (21%) patients treated with Brineura and presented as hematoma, post procedural hematoma, traumatic hematoma and subdural hematoma. Hematomas did not require treatment and did not interfere with Brineura infusion.

### *Hypersensitivity*

Hypersensitivity reactions were reported in 11 out of 24 patients (46%) treated with Brineura during or within 24 hours after completion of the Brineura infusion, despite pre-medication with antihistamines with or without antipyretics or corticosteroids [see *Warnings and Precautions (5.1)*]. The most common manifestations observed concomitantly with hypersensitivity included pyrexia with vomiting, pleocytosis, or irritability, which are not consistent with classic immune mediated hypersensitivity. Symptoms resolved over time or with administration of antipyretics, antihistamines and/or corticosteroids and no patient discontinued treatment with Brineura.

One patient experienced hypoxia (decreased oxygen saturation less than 88% by pulse oximeter), 8 hours after Brineura infusion, followed by a low mean arterial pressure at 15 hours post infusion. Symptoms resolved after oxygen administration, airway repositioning and normal saline infusion. One patient reported decreased oxygen saturation (90% by pulse oximeter), 45 minutes after starting Brineura with associated low diastolic blood pressures. Hypoxia resolved after oxygen administration. No treatment was administered for the low diastolic blood pressure, which returned to normal while the patient continued to receive Brineura infusion without change to the infusion rate or dose.

## **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 antibodies to cerliponase alfa in the studies described below with the incidence of antibodies in other studies or to other products may be misleading.

Anti-drug antibodies (ADAs) to cerliponase alfa were detected in both serum and CSF in 79% and 33%, respectively, of patients treated with Brineura for up to 161 weeks. Patients who experienced hypersensitivity adverse reactions were tested for drug-specific IgE and found to be negative, including three patients for whom grade 3 (severe) hypersensitivity adverse reactions were reported. No association was found between serum or CSF ADA titers and incidence or severity of hypersensitivity. Drug-specific neutralizing antibodies (NAb) have not been evaluated.

## **8 USE IN SPECIFIC POPULATIONS**

### **8.1 Pregnancy**

#### Risk Summary

There are no available data on Brineura use in pregnant women to inform a drug-associated risk of pregnancy-related outcomes. Animal reproduction studies have not been conducted using cerliponase alfa.

The estimated background risk of major birth defects and miscarriage for the indicated population is unknown. All pregnancies have a background risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2 to 4% and 15 to 20%, respectively.

## 8.2 Lactation

### Risk Summary

There are no data on the presence of cerliponase alfa in human milk, the effects on the breastfed child, or the effects on milk production. The lack of clinical data during lactation precludes a clear determination of the risk of Brineura to an infant during lactation; therefore, the development and health benefits of breastfeeding should be considered along with the mother's clinical need for Brineura and any potential adverse effects on the breastfed infant from Brineura or from the underlying maternal condition.

## 8.4 Pediatric Use

Safety and effectiveness of Brineura have been established in pediatric patients 3 years of age and older. Pediatric use of Brineura to slow the loss of ambulation in symptomatic pediatric patients 3 years of age and older with late infantile neuronal ceroid lipofuscinosis type 2 (CLN2), also known as tripeptidyl peptidase 1 (TPP1) deficiency, is supported by a non-randomized single-arm dose escalation clinical study with extension in patients with CLN2 disease and compared to untreated patients with CLN2 disease from an independent natural history cohort [see *Clinical Studies (14)*]. Safety and effectiveness in patients less than 3 years of age have not been established.

## 11 DESCRIPTION

Cerliponase alfa is a purified human enzyme produced by recombinant DNA technology in a Chinese hamster ovary cell line. The active substance is a recombinant human tripeptidyl peptidase-1 (rhTPP1), a lysosomal exopeptidase. The primary activity of the mature enzyme is the cleavage of N-terminal tripeptides from a broad range of protein substrates.

Cerliponase alfa contains 544 amino acids with an average molecular mass of 59 kDa. The mature enzyme is 368 amino acids in length. There are 5 consensus N-glycosylation sites on rhTPP1 that contain high mannose, phosphorylated high mannose and complex glycosylation structures.

Brineura (cerliponase alfa) Injection and Intraventricular Electrolytes Injection are administered by intraventricular infusion. The solutions are sterile, nonpyrogenic, and free of foreign particulates. Brineura is a clear to slightly opalescent and colorless to pale yellow solution. Intraventricular Electrolytes is a clear to colorless solution.

Brineura and Intraventricular Electrolytes Injection are packaged in 10 mL clear Type 1 single-dose glass vials [see *How Supplied/Storage and Handling (16)*]. Each vial of Brineura provides 5 mL of solution containing 150 mg cerliponase alfa. Each vial of Intraventricular Electrolytes Injection provides 5 mL of solution. Both Brineura and Intraventricular Electrolytes Injection are formulated with the following excipients: calcium chloride dihydrate (1.05 mg); magnesium chloride hexahydrate (0.8 mg); potassium chloride (1.1 mg); sodium chloride (43.85 mg); sodium phosphate, dibasic, heptahydrate (0.55 mg); sodium phosphate, monobasic, monohydrate (0.4 mg); and Water for Injection, USP. The pH of the solution is between 6.2 to 6.8 for Brineura, and between 6.0 to 7.0 for Intraventricular Electrolytes Injection.

Each vial contains: sodium: 0.76 mEq, and potassium: 0.015 mEq.

## 12 CLINICAL PHARMACOLOGY

### 12.1 Mechanism of Action

CLN2 disease is a neurodegenerative disease caused by deficiency of the lysosomal enzyme tripeptidyl peptidase-1 (TPP1), which catabolizes polypeptides in the CNS. TPP1 has no known substrate specificity.

Deficiency in TPP1 activity results in the accumulation of lysosomal storage materials normally metabolized by this enzyme in the central nervous system (CNS), leading to progressive decline in motor function.

Cerliponase alfa (rhTTP1), a proenzyme, is taken up by target cells in the CNS and is translocated to the lysosomes through the Cation Independent Mannose-6-Phosphate Receptor (CI-MPR, also known as M6P/IGF2 receptor). Cerliponase alfa is activated in the lysosome and the activated proteolytic form of rhTTP1 cleaves tripeptides from the N-terminus of proteins.

### 12.3 Pharmacokinetics

The pharmacokinetics of cerliponase alfa were evaluated in patients with CLN2 disease who received intraventricular infusions of 30 mg (0.1 times the approved recommended dosage), 100 mg (approximately 0.3 times the approved recommended dosage), and 300 mg over approximately 4.5 hours once every other week.

Cerliponase alfa CSF exposure following the initial single dose administration of Brineura increased less than proportionally across doses of 30 mg, 100 mg, and 300 mg. There was no apparent accumulation of cerliponase alfa in CSF or plasma when Brineura was administered at a dose of 300 mg once every other week.

Cerliponase alfa pharmacokinetics have high inter-subject and intra-subject variability. Following intraventricular infusions of 300 mg of Brineura at Day 1, Week 5, and Week 13, the pharmacokinetic parameters in CSF and plasma were assessed in 14 patients and are summarized in Table 2.

**Table 2: Pharmacokinetic Parameters of Cerliponase Alfa Following Intraventricular Infusion (approximately 4.5 hours in duration) of Brineura 300 mg Every Two Weeks**

|                     | Parameter                          | Median [Min, Max]  |                     |                     |
|---------------------|------------------------------------|--------------------|---------------------|---------------------|
|                     |                                    | Day 1              | Week 5              | Week 13             |
| CSF                 | N                                  | 13                 | 14                  | 13                  |
|                     | T <sub>max</sub> <sup>*</sup> , hr | 4.5 [4.3, 5.8]     | 4.3 [3.8, 4.5]      | 4.3 [4.0, 4.5]      |
|                     | C <sub>max</sub> , mcg/mL          | 1260 [359, 4380]   | 1630 [376, 4670]    | 1390 [1110, 2340]   |
|                     | AUC <sub>0-t</sub> , mcg-hr/mL     | 9290 [3660, 19000] | 12400 [4620, 26200] | 10500 [7000, 18200] |
|                     | V <sub>ss</sub> , mL               | 245 [78.4, 909]    | 196 [85.4, 665]     | 186 [131, 257]      |
|                     | CL, mL/hr                          | 32.3 [15.8, 81.9]  | 24.2 [11.4, 64.9]   | 28.7 [16.5, 42.9]   |
|                     | t <sub>1/2</sub> , hr              | 6.2 [5.5, 16.3]    | 7.4 [3.3, 9.5]      | 7.7 [5.1, 9.4]      |
| Plasma <sup>#</sup> | N                                  | 12                 | 12                  | 9                   |
|                     | T <sub>max</sub> <sup>*</sup> , hr | 12.0 [4.3, 24.5]   | 12.0 [7.5, 24.2]    | 12.3 [4.3, 75.9]    |
|                     | C <sub>max</sub> , mcg/mL          | 1.3 [0.2, 3.9]     | 1.9 [0.2, 4.3]      | 1.0 [0.03, 2.6]     |
|                     | AUC <sub>0-t</sub> , mcg-hr/mL     | 16.2 [1.1, 69.9]   | 40.1 [11.1, 78.9]   | 9.5 [0.2, 51.6]     |
| CSF/Plasma Ratio    | N                                  | 11                 | 12                  | 9                   |
|                     | C <sub>max</sub>                   | 1200 [305, 4530]   | 809 [202, 9370]     | 1320 [541, 51200]   |
|                     | AUC <sub>0-t</sub>                 | 393 [115, 1910]    | 340 [126, 1780]     | 1330 [167, 38900]   |

<sup>\*</sup>T<sub>max</sub> expressed as time since start of ~4.5 hour infusion.

<sup>#</sup>V<sub>ss</sub>, CL, and t<sub>1/2</sub> were not estimated due to insufficient plasma pharmacokinetic data

The estimated CSF volume of distribution of cerliponase alfa following intraventricular infusion of 300 mg of Brineura (median  $V_{ss}$  = 245 mL) exceeds the typical CSF volume (100 mL).

Cerliponase alfa is a protein and is expected to be degraded through peptide hydrolysis.

## **13 NONCLINICAL TOXICOLOGY**

### **13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility**

Carcinogenicity, genotoxicity, and fertility studies have not been performed with cerliponase alfa. Based on the mechanism of action, cerliponase alfa is not expected to be tumorigenic.

## **14 CLINICAL STUDIES**

The efficacy of Brineura was assessed over 96 weeks in a non-randomized single-arm dose escalation clinical study with extension in symptomatic pediatric patients with late infantile neuronal ceroid lipofuscinosis type 2 (CLN2) disease, confirmed by TPP1 deficiency. Brineura-treated patients were compared to untreated patients from a natural history cohort. The Motor domain of a CLN2 Clinical Rating Scale was used to assess disease progression. Scores ranged from 3 (grossly normal) to 0 (profoundly impaired) with unit decrements representing milestone events in the loss of motor function (ability to walk or crawl). Due to the inability to establish comparability for the CLN2 Language domain ratings between the clinical study with extension and the natural history cohort, efficacy of Brineura for the Language domain cannot be established.

Twenty-four patients, aged 3 to 8 years were enrolled in the Brineura single-arm clinical study. Sixty-three percent of patients were female and 37% were male. Ninety-six percent of patients were Caucasian and 4% were Asian. One patient withdrew after week 1 due to inability to continue with study procedures; 23 patients were treated with Brineura 300 mg every other week for 48 weeks, and continued treatment during the extension period.

In the clinical study with extension, patients were assessed for decline in the Motor domain of the CLN2 Clinical Rating Scale at 48, 72 and 96 weeks. Decline was defined as having an unreversed (sustained) 2-category decline or an unreversed score of 0 in the Motor domain of the CLN2 Clinical Rating Scale. Patients' responses to Brineura treatment were evaluated if at screening a combined Motor plus Language CLN2 score of less than 6 was recorded. Two patients with a combined Motor plus Language CLN2 score of 6 were excluded from the analyses; they maintained that score throughout the study period. The patient who terminated early was analyzed as having a decline at the time of termination. Data used in the analyses from the natural history cohort began at 36 months of age or greater and at the first time a Motor plus Language CLN2 score less than 6 was recorded.

Motor scores of the 22 Brineura-treated patients in the clinical study with extension were compared to scores of the independent natural history cohort that included 42 untreated patients who satisfied inclusion criteria for the clinical study. The results of logistic modeling with covariates (screening age, screening motor score, genotype: 0 key mutations (yes/no)), demonstrated the odds of Brineura-treated patients not having a decline by 96 weeks were 13 times the odds of natural history cohort patients not having a decline (Odds Ratio (95% CI): 13.1 (1.2, 146.9)).

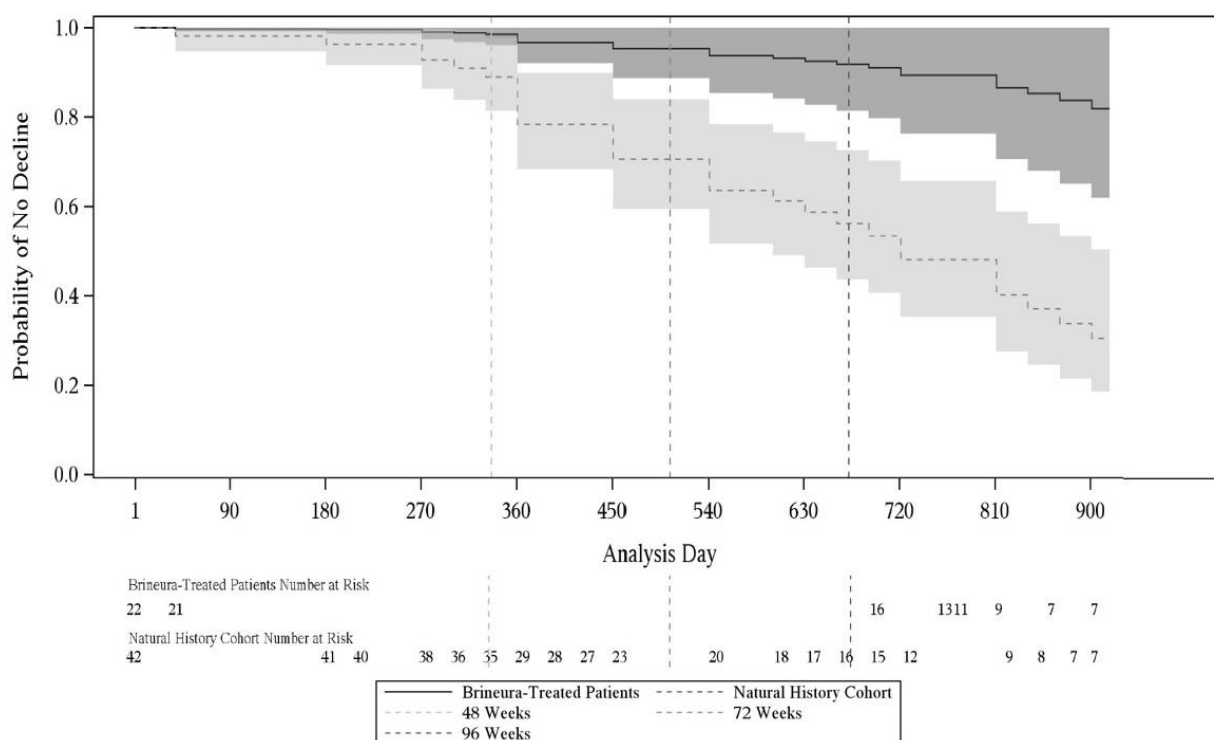
#### Descriptive non-randomized comparison

In an unadjusted non-randomized comparison, of the 22 patients treated with Brineura and evaluated for efficacy at week 96, 21 (95%) did not decline, and only the patient who terminated early was deemed to have a decline in the Motor domain of the CLN2 Clinical Rating Scale. Results from the natural history cohort demonstrated progressive decline in motor function; of the 42 patients in the natural history cohort,

21 (50%) experienced an unreversed (sustained) 2-category decline or unreversed score of 0 in the Motor domain of the CLN2 Clinical Rating Scale over 96 weeks.

Given the non-randomized study design, a Cox Proportional Hazards Model adjusted for age, initial motor score, and genotype was used to evaluate time to unreversed 2-category decline or unreversed score of 0 in the Motor domain. This model showed a lesser decrease in motor function in the Brineura-treated patients when compared to the natural history cohort (see Figure 7).

**Figure 7. Estimated Time to Unreversed (Sustained) 2-Category Decline or Unreversed Score of Zero in Motor Domain for Symptomatic Pediatric Patients in the Brineura Single-Arm Clinical Study with Extension and for Patients in a Natural History Cohort (Based on the Cox Proportional Hazards Model Adjusting for Covariates)**



Shading represents 95% confidence intervals.

Follow-up for the natural history cohort begins at 36 months of age or greater and at the first time a Motor plus Language CLN2 score less than 6 was recorded.

The Brineura-treated population is the full population (N=24) minus two patients with baseline Motor plus Language CLN2 score = 6. Covariates: screening age, screening Motor score, genotype: 0 key mutations (yes/no). "Screening age" was defined in the natural history cohort as the age at the first time a Motor plus Language CLN2 score less than 6 was recorded, and no earlier than 36 months of age. The "screening Motor score" of the natural history cohort was defined as the Motor score at the screening age.

Decline is defined as an unreversed (sustained) 2-category decline or unreversed score of 0 in the Motor domain of the CLN2 Clinical Rating Scale.

#### Motor Domain Scores: Matched Patients Only

To further assess efficacy, the 22 patients from the Brineura clinical study with a baseline combined Motor plus Language CLN2 score less than 6 were matched to 42 patients in the natural history cohort. Patients were matched based on the following covariates: baseline age at time of screening within 3 months, genotype (0, 1, or 2 key mutations), and baseline Motor domain CLN2 score at time of screening.

Using the Motor domain of the CLN2 Clinical Rating Scale, decline was defined as having an unreversed 2-category decline or an unreversed score of 0. At 96 weeks, the matched analysis based on 17 pairs

demonstrated fewer declines in the Motor domain for Brineura-treated patients compared to untreated patients in the natural history cohort (see Table 3).

**Table 3: Proportion of Matched Symptomatic Pediatric Patients with CLN2 Disease without Decline\* in the Brineura Single-Arm Clinical Study with Extension and in the Natural History Cohort assessed at Weeks 48, 72, and 96**

| Time Point/Period         | Natural History Cohort (N=17) | Brineura-Treated (N=17) | Difference    | Odds Ratio***  |
|---------------------------|-------------------------------|-------------------------|---------------|----------------|
|                           | n (%)                         | n (%)                   | % (95% CI**)  | OR (95% CI)    |
| Follow-up through Week 48 | 13 (76)                       | 16 (94)                 | 18% (-19, 51) | 4 (0.4, 200)   |
| Follow-up through Week 72 | 11 (65)                       | 16 (94)                 | 29% (-7, 61)  | 5.9 (0.7, 250) |
| Follow-up through Week 96 | 6 (35)                        | 16 (94)                 | 59% (24, 83)  | 11 (1.6, 500)  |

\*Decline is defined as an unreversed (sustained) 2-category decline or unreversed score of 0 in the Motor domain of the CLN2 Clinical Rating Scale.

\*\* Exact confidence interval for risk difference (Santner and Snell)

\*\*\*Based on McNemar's Exact test

Matched on baseline age at time of screening within 3 months, genotype (0, 1, or 2 key mutations), and baseline Motor domain CLN2 score at time of screening.

The Brineura-treated population is based on the full population minus two patients with baseline Motor plus Language CLN2 score = 6.

## 16 HOW SUPPLIED/STORAGE AND HANDLING

Brineura is supplied as a sterile, clear to slightly opalescent and colorless to pale yellow solution for intraventricular infusion and Intraventricular Electrolytes Injection is supplied as a clear to colorless solution for intraventricular infusion; both are included in package 1 of 2. The Administration Kit for use with Brineura is supplied separately as package 2 of 2 [see *Dosage and Administration (2.2)*].

### Package 1 of 2

Each Brineura (cerliponase alfa) Injection vial has a green flip-off cap (plastic), and contains 150 mg cerliponase alfa per 5 mL (30 mg/mL).

Each Intraventricular Electrolytes Injection vial has a yellow flip-off cap (plastic), and contains 5 mL of solution.

| Contents of Package 1                                                                                                    | NDC Number   |
|--------------------------------------------------------------------------------------------------------------------------|--------------|
| Brineura (cerliponase alfa) Injection (2 vials of 150 mg/5 mL)<br>Intraventricular Electrolytes Injection (1 vial, 5 mL) | 68135-811-02 |

### Package 2 of 2

The Administration Kit for use with Brineura is supplied separately and contains the following single-use, sterile infusion components:

- Two 20-mL syringes
- Two syringe needles (21 G, 25.4 mm)
- One extension line
- One infusion set with 0.2 micron inline filter
- One port needle (22 G, 16 mm)

### Storage

Brineura (cerliponase alfa) Injection and Intraventricular Electrolytes Injection:  
Store upright in a freezer (-25°C to -15°C) in original carton to protect from light.

Administration Kit for use with Brineura:

Store in original carton separately from Brineura. Do not freeze.

## **17 PATIENT COUNSELING INFORMATION**

- **Intraventricular Access Device-Related Complications**  
Advise patients and caregivers of the risk of device-related infections. If any signs of infection are present, instruct patients to immediately contact their healthcare provider [*see Warnings and Precautions (5.1)*].
- **Cardiovascular Adverse Reactions**  
Advise patients and caregivers that hypotension and/or bradycardia may occur during and following the infusion of Brineura. Instruct patients immediately to contact their healthcare provider if these reactions occur [*see Warnings and Precautions (5.2)*].
- **Hypersensitivity Reactions**  
Advise patients and caregivers that hypersensitivity reactions related to Brineura treatment, including fever, vomiting, and irritability may occur. Due to the potential for anaphylaxis, inform patients and caregivers of the signs and symptoms of anaphylaxis, and instruct them to seek immediate medical care should signs and symptoms occur [*see Warnings and Precautions (5.3)*].

Manufactured by:

BioMarin Pharmaceutical Inc.

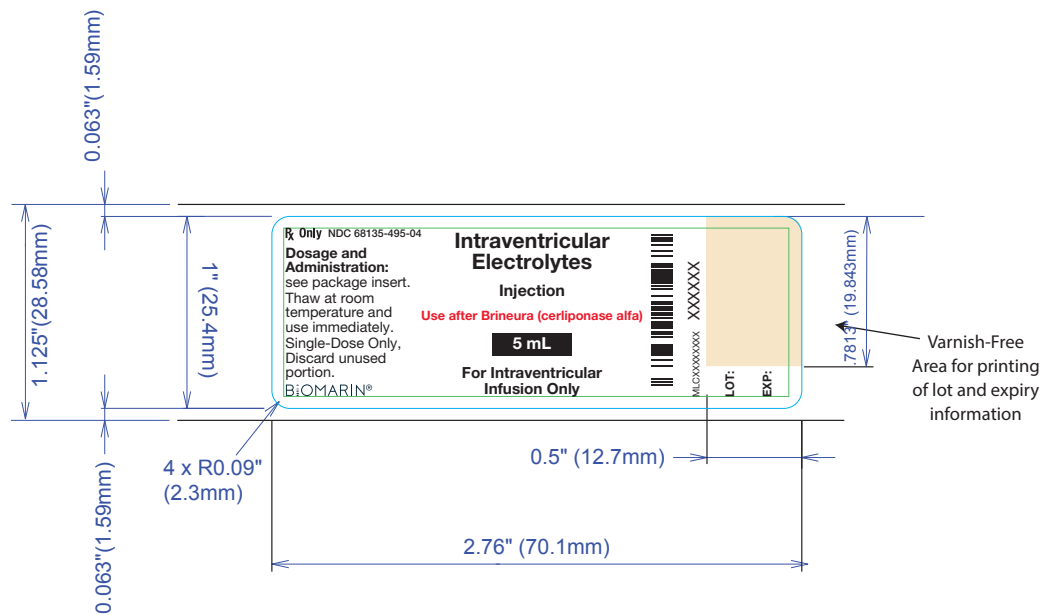
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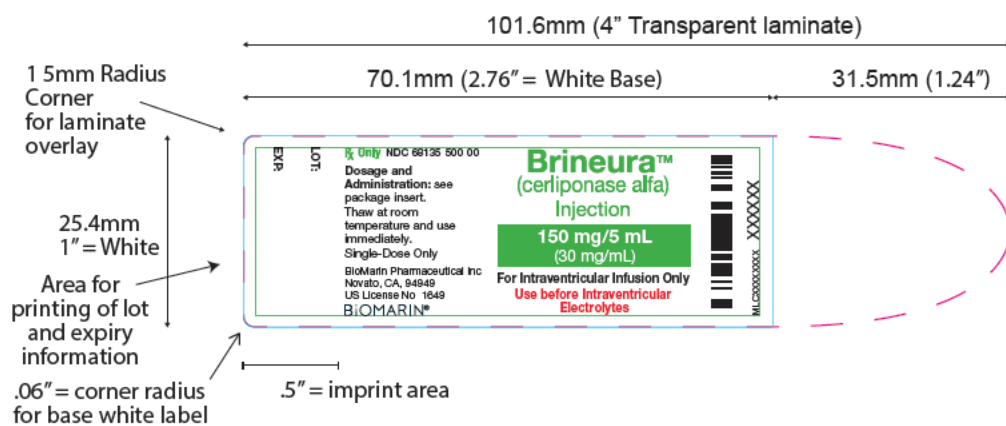
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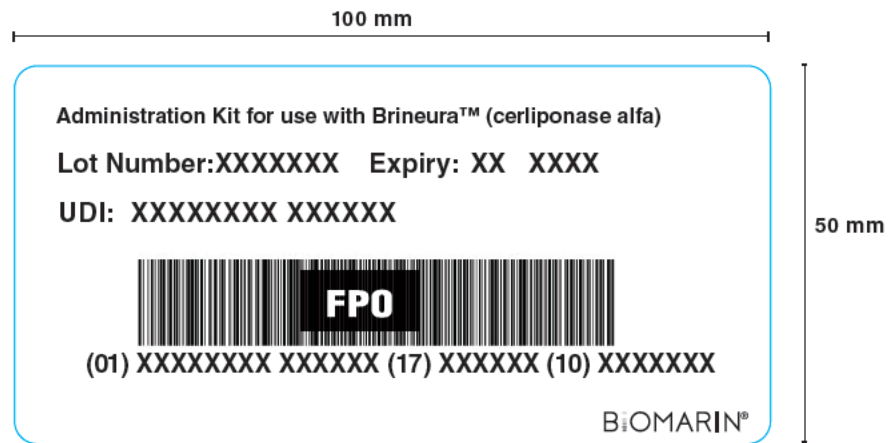
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JULIE G BEITZ  
04/27/2017