# CAPILLARY LEAK SYNDROME (CLS) MANAGEMENT GUIDE



# **INDICATION**

ELZONRIS® (tagraxofusp-erzs) is a CD123-directed cytotoxin for the treatment of blastic plasmacytoid dendritic cell neoplasm (BPDCN) in adults and in pediatric patients 2 years and older.

# **Boxed WARNING: CAPILLARY LEAK SYNDROME**

Capillary Leak Syndrome (CLS), which may be life-threatening or fatal, can occur in patients receiving ELZONRIS. Monitor for signs and symptoms of CLS and take actions as recommended.

Please see full Important Safety Information on the back cover and full <u>Prescribing Information</u>, including Boxed WARNING.



Scan here for more information on CLS management.

# **ELZONRIS** and **CLS**

In clinical studies of ELZONRIS, 53% of patients experienced CLS of any grade, including 4 fatal events (N=122).<sup>1</sup>

- >90% of CLS occurred in cycle 1
- In long term follow up, the median time to onset was 4 days (range 1 to 46 days)

Common signs and symptoms of CLS with ELZONRIS<sup>1</sup>



Hypoalbuminemia

Edema, including pulmonary edema





Weight gain

Hypotension





Hemodynamic instability

Assess all patients appropriately before and throughout ELZONRIS treatment<sup>1</sup>



Before initiating therapy with ELZONRIS (first dose of first cycle):

- Ensure patient has adequate cardiac function\*
- Ensure patient has serum albumin ≥3.2 g/dL
- Weigh patient to establish baseline weight for subsequent dose

\*In the clinical studies, patients had a left ventricular ejection fraction ≥ institutional lower limit of normal as measured by multigated acquisition scan or 2-dimensional echocardiography within 28 days prior to start of therapy and no clinically significant abnormalities on a 12-lead electrocardiogram.²



# During treatment with ELZONRIS1:

- Monitor serum albumin levels prior to the initiation of each dose of ELZONRIS and as clinically indicated thereafter
- Assess patients for signs/symptoms of CLS, including:
  - Serum albumin <3.5 g/dL or reduced by ≥0.5 g/dL from the albumin value measured prior to ELZONRIS dosing initiation of the current cycle
  - · New onset or worsening edema, including pulmonary edema
  - Weight gain ≥1.5 kg from the previous day's predose weight
  - Hypotension (systolic blood pressure ≥ 160 mmHg or ≤ 80 mmHg)
  - Hemodynamic instability (heart rate ≥ 130 bpm or ≤ 40 bpm)

# Observe patients during ELZONRIS administration<sup>1</sup>



# Cycle 1

 Administer in the inpatient setting, and observe patients for at least 24 hours after the last infusion



# Subsequent cycles

- Administer in an inpatient setting or an appropriate outpatient setting
- Observe patients for at least 4 hours after each infusion

# Counsel patients upon discharge<sup>1</sup>



Advise patients of the risk of CLS and to contact their healthcare provider for signs and symptoms associated with CLS, including:

- New or worsening edema
- Weight gain
- Shortness of breath
- Hypotension after infusion



Advise patients to weigh themselves daily

Capillary Leak Syndrome (CLS), which may be life-threatening or fatal if not properly managed, can occur in patients receiving ELZONRIS.<sup>1</sup>

Please see Important Safety Information, including Boxed WARNING, on the back cover.

# Management of CLS is an important part of treating patients with ELZONRIS<sup>1</sup>

# CLS management guidelines

Time of presentation	CLS sign/ symptom	Recommended action	ELZONRIS dosing management
Prior to first dose of ELZONRIS in cycle 1	Serum albumin <3.2 g/dL	Administer ELZONRIS when serum albumin ≥3.2 g/dL	
During ELZONRIS dosing	Serum albumin <3.5 g/dL		Interrupt ELZONRIS dosing until the relevant CLS sign/ symptom has resolved.*
	Serum albumin reduced by ≥0.5 g/dL from the albumin value measured prior to ELZONRIS dosing initiation of the current cycle	Administer 25 g intravenous albumin (q12h or more frequently as practical) until serum albumin is ≥3.5 g/dL AND not more than 0.5 g/dL lower than the value measured prior to dosing initiation of the current cycle	
	A predose body weight that is increased by ≥1.5 kg over the previous day's predose weight	Administer 25 g intravenous albumin (q12h or more frequently as practical) and manage fluid status as clinically indicated (eg, generally with intravenous fluids and vasopressors if hypotensive and with diuretics if normotensive or hypertensive) until body weight increase has resolved (the increase is no longer ≥1.5 kg greater than the previous day's predose weight)	
	Edema, fluid overload, and/or hypotension	Administer 25 g intravenous albumin (q12h, or more frequently as practical) until serum albumin is ≥3.5 g/dL  Administer 1 mg/kg of methylprednisolone (or an equivalent) per day until resolution of CLS sign/symptom or as clinically indicated Aggressive management of fluid status and hypotension if present, which could include intravenous fluids and/or diuretics or other blood pressure management, until resolution of CLS sign/symptom or as clinically indicated	

### \*If ELZONRIS dose is held:

- ELZONRIS administration may resume in the same cycle if all CLS signs/symptoms have resolved and the patient did not require measures to treat hemodynamic instability
- ELZONRIS administration should be held for the remainder of the cycle if CLS signs/ symptoms have not resolved or the patient required measures to treat hemodynamic instability (eg, required administration of intravenous fluids and/or vasopressors to treat hypotension) (even if resolved)
- ELZONRIS administration may only resume in the next cycle if all CLS signs/symptoms have resolved and the patient is hemodynamically stable.

q12h, every 12 hours.

# Please see Important Safety Information, including Boxed WARNING, on the back cover.

# **Helpful Resources**

## CLS Risk Alert Kit

The kit contains the following:



The **Alert Bracelet** should be placed on the patient's wrist, along with their hospital bracelet, to serve as an alert to staff while the patient receives inpatient care



The **Alert Sign** should be placed in the most visible location in the patient's room to serve as an alert to any healthcare professional involved in the care of the patient



The **Chart Sticker** should be placed on the patient's bedside chart to serve as a reminder to any healthcare professional monitoring the patient

Consider also putting an alert in the patient's electronic medical record (EMR) or electronic health record (EHR).

# Pre-administration Parameters Tracking Tool

The Pre-Administration Parameters Tracking Tool is a guide designed to assist HCPs in monitoring patients before and during treatment with ELZONRIS. It helps track important parameters such as body weight, albumin, blood pressure, hemodynamics, and provides guidelines for dose modifications/interruptions.



To order these resources, or if interested in CLS training, scan the code to contact your ELZONRIS representative.

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### INDICATION

 ELZONRIS is a CD123-directed cytotoxin indicated for the treatment of blastic plasmacytoid dendritic cell neoplasm (BPDCN) in adults and in pediatric patients 2 years and older

### IMPORTANT SAFETY INFORMATION

### **BOXED WARNING: CAPILLARY LEAK SYNDROME**

 Capillary Leak Syndrome (CLS) which may be life-threatening or fatal, can occur in patients receiving ELZONRIS. Monitor for signs and symptoms of CLS and take actions as recommended.

### WARNINGS AND PRECAUTIONS

### Capillary Leak Syndrome

- Capillary leak syndrome (CLS), including life-threatening and fatal cases, has been reported among patients treated with ELZONRIS. In patients receiving ELZONRIS in clinical trials, the overall incidence of CLS was 53% (65/122), including Grade 1 or 2 in 43% (52/122) of patients, Grade 3 in 7% (8/122) of patients, Grade 4 in 1% (1/122) of patients, and four fatalities (3%). The median time to onset was 4 days (range 1 to 46 days), and all but 5 patients experienced an event in Cycle 1.
- Before initiating therapy with ELZONRIS, ensure that the patient has adequate cardiac
  function and serum albumin is greater than or equal to 3.2 g/dL. During treatment with
  ELZONRIS, monitor serum albumin levels prior to the initiation of each dose of
  ELZONRIS and as indicated clinically thereafter, and assess patients for other signs or
  symptoms of CLS, including weight gain, new onset or worsening edema, including
  pulmonary edema, hypotension or hemodynamic instability.

### **Hypersensitivity Reactions**

• ELZONRIS can cause severe hypersensitivity reactions. In patients receiving ELZONRIS in clinical trials, hypersensitivity reactions were reported in 43% (53/122) of patients treated with ELZONRIS and were Grade ≥ 3 in 7% (9/122). Manifestations of hypersensitivity reported in ≥ 5% of patients include rash, pruritus, and stomatitis. Monitor patients for hypersensitivity reactions during treatment with ELZONRIS. Interrupt ELZONRIS infusion and provide supportive care as needed if a hypersensitivity reaction should occur.

### **Hepatotoxicity**

- Treatment with ELZONRIS was associated with elevations in liver enzymes. In patients
  receiving ELZONRIS in clinical trials, elevations in ALT occurred in 79% (96/122) and
  elevations in AST occurred in 76% (93/122). Grade 3 ALT elevations were reported in
  26% (32/122) of patients. Grade 3 AST elevations were reported in 30% (36/122) and
  Grade 4 AST elevations were reported in 3% (4/122) of patients. Elevated liver
  enzymes occurred in the majority of patients in Cycle 1 and were reversible following
  dose interruption.
- Monitor alanine aminotransferase (ALT) and aspartate aminotransferase (AST) prior to each infusion with ELZONRIS. Withhold ELZONRIS temporarily if the transaminases rise to greater than 5 times the upper limit of normal and resume treatment upon normalization or when resolved.

### ADVERSE REACTIONS:

Most common adverse reactions (incidence  $\geq$  30%) are capillary leak syndrome, nausea, fatigue, pyrexia, peripheral edema, and weight increase. Most common laboratory abnormalities (incidence  $\geq$  50%) are decreases in albumin, platelets, hemoglobin, calcium, and sodium, and increases in glucose, ALT and AST.

### Please see full Prescribing Information, including Boxed WARNING.

To report SUSPECTED ADVERSE REACTIONS, contact Stemline Therapeutics, Inc. at 1-877-332-7961 or contact the FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

**References: 1.** ELZONRIS [prescribing information]. New York, NY: Stemline Therapeutics, Inc.; July 2023. **2.** Data on file. Stemline Therapeutics, Inc.

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