Early Warning Signs of High Dose Methotrexate (HDMTX) Induced Acute Kidney Injury (AKI)

Recommendations from the Consensus Guidelines for the Use of Glucarpidase in Patients with HDMTX-Induced AKI and Delayed MTX Clearance


These recommendations were adapted from published consensus guidelines in which the authors provided guidance for many of the time points that are routinely monitored in clinical practice. Consulting with an experienced oncologist, nephrologist, or clinical pharmacist who is familiar with managing patients treated with HDMTX infusions is recommended if the patient has prior HDMTX-induced AKI or a GFR < 75 mL/min/1.73 m².

Early warning signs

Treatment with glucarpidase is recommended in the case of excessively high MTX concentrations and rising creatinine.

• However, serum creatinine is a suboptimal biomarker of AKI, as creatinine rise may lag significantly from the time of the renal insult.

• Elevated plasma MTX concentration may indicate HDMTX-induced AKI prior to a significant change in creatinine.

• Thus, clinicians administering HDMTX should be familiar with the expected plasma MTX concentration at the various time points after infusion.

Considerations based on MTX dose and infusion duration

1 – 8 g/m² MTX Infused over 24 – 42 Hours

• For a 24-hour infusion, a plasma MTX concentration >120µM (54.5µg/mL) at the end of the infusion or a creatinine increase 50% over baseline warrants additional plasma MTX monitoring at 36 hours. If the 36-hour MTX concentration is above 30µM (13.6µg/mL), the 42-hour MTX concentration is above 10µM (4.54µg/mL), or the 48-hour concentration is above 5µM (2.27µg/mL) and the serum creatinine is rising, glucarpidase is recommended.

• For 36 – 42 hour infusions, if the 48-hour MTX concentration is above 5µM (2.27µg/mL) and serum creatinine is rising, glucarpidase is recommended.

8 – 12 g/m² MTX Infused over ≤ 6 Hours

• A plasma MTX concentration > 1500 µM (681µg/mL) at the end of the infusion warrants additional plasma MTX monitoring at 24 hours. If the 24-hour MTX concentration is above 50µM (22.7µg/mL), the 36-hour concentration is above 30µM (13.6µg/mL), the 42-hour concentration is above 10µM (4.54µg/mL), or the 48-hour concentration is above 5µM (2.27µg/mL), and the serum creatinine is rising, glucarpidase is recommended.

For methotrexate doses of ≤ 1g/m² that are infused over 36 – 42 Hours

• If the 48 hour concentration is above 5µM (2.27µg/mL) and the serum creatinine is rising, glucarpidase is recommended.

Administration of glucarpidase should optimally occur within 48 – 60 hours from the start of the HDMTX infusion, because life-threatening toxicities may not be preventable beyond this time point.


Indication and Limitations of Use

Voraxaze® (glucarpidase) is indicated for the treatment of toxic plasma methotrexate concentrations (>1μmol/L) in patients with delayed methotrexate clearance due to impaired renal function.

Voraxaze® is not indicated for use in patients who exhibit the expected clearance of methotrexate (plasma methotrexate concentrations within 2 standard deviations of the mean methotrexate excretion curve specific for the dose of methotrexate administered) or those with normal or mildly impaired renal function because of the potential risk of subtherapeutic exposure to methotrexate.

Please see Important Safety Information on reverse side and accompanying full Prescribing Information.
**Important Safety Information for Voraxaze® (glucarpidase)**

**Warnings and Precautions:**

**Serious Allergic Reactions:**
Serious allergic reactions, including anaphylactic reactions, may occur.

**Monitoring Methotrexate Concentration/Interference With Assay:**
Methotrexate concentrations within 48 hours following Voraxaze® administration can only be reliably measured by a chromatographic method due to interference from metabolites. Measurement of methotrexate concentrations within 48 hours of Voraxaze® administration using immunoassays can overestimate the methotrexate concentration.

**Continuation and Timing of Leucovorin Rescue:**
Leucovorin should not be administered within 2 hours before or after Voraxaze® dose because leucovorin is a substrate for Voraxaze®.

For the first 48 hours after Voraxaze®, administer the same leucovorin dose as given prior to Voraxaze®. Beyond 48 hours after Voraxaze®, administer the same leucovorin dose as given prior to Voraxaze®.

Therapy with leucovorin should be continued until the methotrexate concentration has been maintained below the leucovorin threshold. Continue hydration and alkalinization of the urine as indicated.

**Adverse Reactions:**
In clinical trials, the common related adverse events (occurring in >1% of patients) were:

- Paresthesias
- Flushing
- Nausea and/or vomiting
- Hypotension
- Headache

**MTX Monitoring Guideline and Glucarpidase Treatment Algorithm**

<table>
<thead>
<tr>
<th>MTX Dose</th>
<th>MTX Monitoring Strategy</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤1 g/m²</td>
<td>Over 36 hours</td>
</tr>
<tr>
<td>≤8 g/m²</td>
<td>Over 24 hours</td>
</tr>
<tr>
<td>≤12 g/m²</td>
<td>Over 12 hours</td>
</tr>
</tbody>
</table>

Glucarpidase strongly recommended in the context of a rising creatinine.

Administration of glucarpidase should optimally occur within 48 – 60 hours after the start of the HDMTX infusion.

Glucarpidase Monitoring Guideline and Glucarpidase Treatment Algorithm

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[Abbreviations: Bl., baseline; Cr, serum creatinine; LV, leucovorin (folinic acid, citrovorum factor, 5-methyltetrahydrofolate); MTX, methotrexate; [MTX], plasma methotrexate concentration]

Adapted from Ramsey, et. al p. 7 fig 5
VORAXAZE® (glucarpidase)
For Injection, for intravenous use
Initial U.S. Approval: 2012

--- RECENT MAJOR CHANGES ---
Immunogenicity (6.2) 03/2013

--- INDICATIONS AND USAGE ---
- VORAXAZE® (glucarpidase) is a carboxypeptidase enzyme indicated for the treatment of toxic plasma methotrexate concentrations (>1 micromole per liter) in patients with delayed methotrexate clearance due to impaired renal function. (1.1)
- Limitation of use: VORAXAZE is not indicated for use in patients who exhibit the expected clearance of methotrexate (plasma methotrexate concentrations within 2 standard deviations of the mean methotrexate excretion curve specific for the dose of methotrexate administered) or those with normal or mildly impaired renal function because of the potential risk of subtherapeutic exposure to methotrexate. (1.2)

--- DOSAGE AND ADMINISTRATION ---
Administer VORAXAZE as a single intravenous injection of 50 Units per kg.

--- DOSAGE FORMS AND STRENGTHS ---
Lyophilized powder 1,000 Units per vial (3)

--- CONTRAINDICATIONS ---
None

--- WARNINGS AND PRECAUTIONS ---
- Serious allergic reactions, including anaphylactic reactions, may occur. (5.1)
- Measurement of methotrexate using immunoassays is unreliable for samples collected within 48 hours following VORAXAZE administration. (5.2)
- Continue therapy with leucovorin until the methotrexate concentration has been maintained below the leucovorin treatment threshold for a minimum of 3 days. (5.3)
- Do not administer leucovorin within 2 hours before or after a dose of VORAXAZE. (5.3)
- For 48 hours after VORAXAZE administration, determine the leucovorin dose based on the patient’s pre-VORAXAZE methotrexate concentration. (5.3)
- Continue hydration and alkalization of the urine as indicated. (5.3)

--- ADVERSE REACTIONS ---
In clinical trials, the most common related adverse events (occurring in >1% of patients) were paraesthesia, flushing, nausea and/or vomiting, hypotension and headache. (6)

To report SUSPECTED ADVERSE REACTIONS, call 877-377-3784 or contact FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

--- DRUG INTERACTIONS ---
- Leucovorin is a substrate for VORAXAZE. Do not administer leucovorin within 2 hours before or after a dose of VORAXAZE. (7.1)
- Other potential exogenous substrates of VORAXAZE include reduced folates and folate antimetabolites. (7.2)

--- USE IN SPECIFIC POPULATIONS ---
Pregnancy: No human or animal data. Use only if clearly needed. (8.1)
Renal Impairment: No dose adjustment is recommended in patients with renal impairment. (8.6)

See Section 17 for PATIENT COUNSELING INFORMATION.

Revised: 03/2013

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

1 INDICATIONS AND USAGE

1.1 Indication
VORAXAZE (glucarpidase) is indicated for the treatment of toxic plasma methotrexate concentrations (>1 micromole per liter) in patients with delayed methotrexate clearance due to impaired renal function.

1.2 Limitation of Use
VORAXAZE is not indicated for use in patients who exhibit the expected clearance of methotrexate (plasma methotrexate concentrations within 2 standard deviations of the mean methotrexate excretion curve specific for the dose of methotrexate administered) or those with normal or mildly impaired renal function because of the potential risk of subtherapeutic exposure to methotrexate.

2 DOSAGE AND ADMINISTRATION

2.1 Recommended Dose
Administer VORAXAZE as a single intravenous injection of 50 Units per kg.

2.2 Administration
Administer VORAXAZE intravenously as a bolus injection over 5 minutes. Flush intravenous line before and after administration of VORAXAZE.

2.3 Preparation
1. Reconstitute the contents of the vial with 1 mL of sterile saline for injection, USP.
2. Roll and tilt the vial gently to mix. Do not shake.
3. Inspect the vial and discard VORAXAZE if the solution is not clear, colorless, and free of particulate matter.
4. Use reconstituted VORAXAZE immediately or store under refrigeration at 36°F to 46°F (2°C to 8°C) for up to 4 hours if not used immediately. VORAXAZE contains no preservative and is supplied as a single-use vial. Discard any unused product [see How Supplied/Storage and Handling (16)].

3 DOSAGE FORMS AND STRENGTHS
Lyophilized powder 1,000 Units per vial

4 CONTRAINDICATIONS
None

5 WARNINGS AND PRECAUTIONS

5.1 Serious Allergic Reactions
Serious allergic reactions occurred in less than 1% of patients [see Adverse Reactions (6.1)].

5.2 Monitoring Methotrexate Concentration/Interference with Assay
Methotrexate concentrations within 48 hours following administration of VORAXAZE can only be reliably measured by a chromatographic method. DAMPA (4-deoxy-4-amino-N10-methylpteroic acid) is an inactive metabolite of methotrexate resulting from treatment with VORAXAZE. DAMPA interferes with the measurement of methotrexate concentration using immunoassays resulting in an erroneous measurement which overestimates the methotrexate concentration. Due to the long half-life of DAMPA (t1/2 of approximately 9 hours), measurement of methotrexate using immunoassays is unreliable for samples collected within 48 hours following VORAXAZE administration [see Clinical Pharmacology (12.1)].

5.3 Continuation and Timing of Leucovorin Rescue
Continue to administer leucovorin after VORAXAZE. Do not administer leucovorin within 2 hours before or after a dose of VORAXAZE because leucovorin is a substrate for VORAXAZE [see Drug Interactions (7.1)].

For the first 48 hours after VORAXAZE, administer the same leucovorin dose as given prior to VORAXAZE [see Warnings and Precautions (5.2)]. Beyond 48 hours after VORAXAZE, administer leucovorin based on the measured methotrexate concentration. Do not discontinue therapy with leucovorin based on the determination of a single methotrexate concentration below the leucovorin treatment threshold. Therapy with leucovorin should be continued until the methotrexate concentration has been maintained below the leucovorin treatment threshold for a minimum of 3 days.

Continue hydration and alkalinization of the urine as indicated.

6 ADVERSE REACTIONS
Serious allergic reactions, including anaphylactic reactions, may occur. The most common adverse reactions (incidence >1%) with VORAXAZE are paraesthesias, flushing, nausea and/or vomiting, hypotension, and headache.

6.1 Clinical Trials Experience
Because clinical trials are conducted under controlled but widely varying conditions, adverse reaction rates observed in clinical trials of VORAXAZE cannot be directly compared to rates in the clinical trials of other drugs and may not reflect the rates observed in practice.

The evaluation of adverse reactions in patients treated with VORAXAZE is confounded by the population in which it was studied, patients with toxic plasma methotrexate levels due to impaired renal function. Adverse reactions related to toxic methotrexate levels due to prolonged methotrexate clearance include myelosuppression, mucositis, acute hepatitis, and renal dysfunction and failure.

The safety of VORAXAZE is based on data from 290 patients who were treated in 2 single-arm, open-label, multicenter trials enrolling patients who had markedly delayed methotrexate clearance secondary to renal dysfunction. Patients with osteosarcoma were eligible for these studies if the plasma methotrexate concentration was greater than 50 µmol/L at 24 hours, greater than 5 µmol/L at 48 hours, or greater than 2 standard deviations above the mean methotrexate elimination curve at least 12 hours after methotrexate administration and there was a 2-fold or greater increase in serum creatinine above baseline. All other patients were eligible for these studies if the plasma methotrexate level was greater than 10 µmol/L more than 42 hours after the start of the methotrexate or the plasma level was greater than 2 standard deviations above the mean methotrexate excretion curve at least 12 hours following methotrexate and the serum creatinine was greater than 1.5 times the upper limit of normal or the creatinine clearance was less than 60 mL/min at least 12 hours following methotrexate administration.

Study 1, conducted by the National Cancer Institute (NCI), enrolled 184 patients; safety information is available for 149 patients. VORAXAZE was given at a dose of 50 Units/kg as an intravenous injection over 5 minutes. Patients with pre-
VORAXAZE methotrexate concentrations >100 μmol/L were to receive a second dose of VORAXAZE 48 hours after the first dose. The protocol specified that patients continue receiving intravenous hydration, urinary alkalization and leucovorin, and that leucovorin administration be adjusted to ensure that it was not administered within two hours before or after VORAXAZE.

In Study 1, VORAXAZE-related adverse reactions were collected on a flow sheet with a daily log of adverse reactions characterized as “glucarpidase toxicity.” Additional safety information was collected from clinical records submitted by treating physicians. This information was abstracted and categorized using the National Cancer Institute (NCI) “Common Terminology Criteria for Adverse Events” (CTCAE) version 3 scale.

The Study 1 population enrolled patients with a median age of 18 years (1 month to 85 years); 63% were male, and the underlying malignancies were osteosarcoma/sarcomas in 32%, and leukemia or lymphoma in 63% of patients. One (n=106) or 2 (n= 30) doses of VORAXAZE were administered intravenously; the number of doses was not specified in 13 patients. Doses ranged from 18 to 98 Units/kg, with a median dose of 49 Units/kg.

Study 2 is an ongoing expanded access program. At the time of data cut-off, 243 patients were enrolled and safety data was available for 141 patients. VORAXAZE was given at a dose of 50 Units/kg as an intravenous injection over 5 minutes. The criterion for allowing patients to receive a second glucarpidase dose was not specified in the protocol. The protocol specified that patients continue receiving intravenous hydration, urinary alkalization and leucovorin, and that leucovorin administration be adjusted to ensure that it was not administered within two hours before or after VORAXAZE.

Study 2 enrolled patients with a median age of 17 years (6 months to 85 years); 64% were male, and the underlying malignancies were osteogenic sarcoma in 32%, and leukemia or lymphoma in 62% of patients. One (n=122) or 2 (n= 18) doses of VORAXAZE were administered intravenously; the number of doses was not specified for 1 patient. Doses ranged from 6 to 189 Units/kg, with a median dose of 50 Units/kg.

In Study 2 only VORAXAZE-related adverse reactions were collected and severity was graded according to NCI CTCAE version 3.

Among the 290 patients included in the safety evaluation of VORAXAZE, there were 8 deaths within 30 days of VORAXAZE exposure that were not related to progressive disease. Twenty-one of 290 patients (7%) experienced adverse reactions that were assessed as related to VORAXAZE. Most were Grade 1 or 2 events. One patient experienced related Grade 3 flushing. The most common related adverse reactions that were not hematologic, hepatic or renal events were paresthesia, flushing, and nausea and/or vomiting, which each occurred in 2% of patients (Table 1).

<table>
<thead>
<tr>
<th>Adverse Reaction</th>
<th>N= 290 n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paresthesias</td>
<td>7 (2%)</td>
</tr>
<tr>
<td>Flushing&lt;sup&gt;1,2&lt;/sup&gt;</td>
<td>5 (2%)</td>
</tr>
<tr>
<td>Nausea/Vomiting</td>
<td>5 (2%)</td>
</tr>
<tr>
<td>Headache</td>
<td>2 (1%)</td>
</tr>
<tr>
<td>Hypotension</td>
<td>2 (1%)</td>
</tr>
<tr>
<td>Blurred Vision</td>
<td>1 (&lt;1%)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>1 (&lt;1%)</td>
</tr>
<tr>
<td>Hypersensitivity</td>
<td>1 (&lt;1%)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>1 (&lt;1%)</td>
</tr>
<tr>
<td>Rash</td>
<td>1 (&lt;1%)</td>
</tr>
<tr>
<td>Throat irritation/Throat tightness</td>
<td>1 (&lt;1%)</td>
</tr>
<tr>
<td>Tremor</td>
<td>1 (&lt;1%)</td>
</tr>
</tbody>
</table>

<sup>1</sup> This incidence includes the following terms: flushing, feeling hot, burning sensation.

<sup>2</sup> One of these reactions was classified as Grade 3 in severity.

### 6.2 Immunogenicity

As with all therapeutic proteins, there is potential for immunogenicity. In clinical trials, 121 patients who received one (n=99), two (n=21), or three (n=1) doses of VORAXAZE were evaluated for anti-glucarpidase antibodies. Twenty-five of these 121 patients (21%) had detectable anti-glucarpidase antibodies following VORAXAZE administration, of which 19 received a single dose of VORAXAZE and 6 received two doses of VORAXAZE. Antibody titers were determined using a bridging enzyme-linked immunosorbent assay (ELISA) for anti-glucarpidase antibodies.

Neutralizing antibodies were detected in 11 of the 25 patients who tested positive for anti-glucarpidase binding antibodies. Eight of these 11 patients had received a single dose of VORAXAZE. However, the development of neutralizing antibodies may be underreported due to lack of assay sensitivity.

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 incidence of antibodies to VORAXAZE with the incidence of antibodies to other products may be misleading.
7 DRUG INTERACTIONS

7.1 Use of VORAXAZE with Leucovorin
Leucovorin is a substrate for VORAXAZE. Do not administer leucovorin within 2 hours before or after a dose of VORAXAZE. No dose adjustment is recommended for the continuing leucovorin regimen because the leucovorin dose is based on the patient’s pre-VORAXAZE methotrexate concentration [see Warnings and Precautions (5.3) and Clinical Pharmacology (12.3)].

7.2 Other Substrate Interference
Other potential exogenous substrates of VORAXAZE may include reduced folates and folate antimetabolites.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy
Pregnancy category C.

There are no adequate and well controlled studies with VORAXAZE in pregnant women and animal reproduction studies have not been conducted with VORAXAZE. Therefore, it is not known whether VORAXAZE can cause fetal harm when administered to a pregnant woman. VORAXAZE should be given to a pregnant woman only if clearly needed.

8.3 Nursing Mothers
It is not known if VORAXAZE is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised when VORAXAZE is administered to a nursing woman.

8.4 Pediatric Use
The effectiveness of VORAXAZE in pediatric patients was established in Study 1. Of the 22 patients in the efficacy dataset in Study 1, 12 were pediatric patients with ages ranging from 5 to 16 years. Three of the six pediatric patients with a pre-VORAXAZE methotrexate concentration of 1-50 µmol/L achieved a rapid and sustained clinically important reduction (RSCIR) in plasma methotrexate concentration, while none of the six pediatric patients with a pre-VORAXAZE methotrexate concentration >50 µmol/L achieved a RSCIR [see Clinical Studies (14)].

The pooled clinical safety database for VORAXAZE included data for 147 patients from 1 month up to 17 years of age. No overall differences in safety were observed between these patients and adult patients.

8.5 Geriatric Use
Of the total number of 290 patients in clinical studies of VORAXAZE, 15% were 65 and over, while 4% were 75 and over. No overall differences in safety or effectiveness were observed between these patients and younger patients.

8.6 Renal Impairment
No dose adjustment of VORAXAZE is recommended for patients with renal impairment [see Clinical Pharmacology (12.3)].

8.7 Hepatic Impairment
No specific studies of VORAXAZE in patients with hepatic impairment have been conducted.

10 OVERDOSAGE
There are no known cases of overdose with VORAXAZE.

11 DESCRIPTION
VORAXAZE (glucarpidase) is a carboxypeptidase produced by recombinant DNA technology in genetically modified Escherichia coli. Glucarpidase is a 390-amino acid homodimer protein with a molecular weight of 83 kDa. Each potency Unit corresponds to the enzymatic cleavage of 1 µmol/L of methotrexate per minute at 37°C.

VORAXAZE is supplied as a sterile, preservative-free, white lyophilized powder in single-use vials. Each vial contains 1,000 Units of glucarpidase, lactose monohydrate (10 mg), Tris-HCl (0.6 mg) and zinc acetate dihydrate (0.002 mg).

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action
VORAXAZE (glucarpidase) is a recombinant bacterial enzyme that hydrolyzes the carboxyl-terminal glutamate residue from folic acid and classical antifolates such as methotrexate. VORAXAZE converts methotrexate to its inactive metabolites 4-deoxy-4-amino-N10-methylpteroyl acid (DAMPA) and glutamate. VORAXAZE provides an alternate non-renal pathway for methotrexate elimination in patients with renal dysfunction during high-dose methotrexate treatment.

12.2 Pharmacodynamics
Plasma methotrexate concentrations within 48 hours following administration of VORAXAZE can only be reliably measured by a chromatographic method because DAMPA interferes with the immunoassays [see Warnings and Precautions (5.2)]. Following administration of VORAXAZE 50 Units/kg to patients in Study 1, methotrexate concentration measured by a chromatographic method was reduced by ≥97% within 15 minutes in all 22 treatment-evaluable patients, and was maintained at a >95% reduction up to 8 days in 20 of the 22 patients [see Clinical Studies (14)].

12.3 Pharmacokinetics
The pharmacokinetics of glucarpidase in the absence of methotrexate were studied in eight healthy subjects following an intravenous injection of VORAXAZE 50 Units/kg over 5 minutes. Serum glucarpidase activity levels were measured by an enzymatic assay and serum total glucarpidase concentrations were measured by ELISA.

Serum glucarpidase activity levels declined with a mean elimination half-life (t1/2) of 5.6 hours. The mean Cmax was 3.3 µg/mL and the mean area under the curve (AUC0-Inf) was 23.3 µg h/mL. The mean systemic clearance (CL) was 7.5 mL/min. The mean volume of distribution (Vd) was 3.6 L, suggesting that glucarpidase distribution is restricted to plasma volume. The pharmacokinetic parameters derived from the serum total glucarpidase concentrations were similar to those generated by serum glucarpidase activity levels except for a longer t1/2 of 9 hours.

Renal Impairment
The pharmacokinetics of glucarpidase in the absence of methotrexate were studied in four subjects with severe renal impairment (creatinine clearance <30 mL/min). Following an intravenous dose of 50 Units/kg of VORAXAZE, the mean pharmacokinetic parameters were similar to those observed in healthy subjects except for a longer t1/2 of 8.2 hours as compared to 5.6 hours in healthy subjects by the enzymatic assay.
Drug Interactions
In a study of cancer patients receiving a high-dose methotrexate (≥1 g/m²) and leucovorin rescue regimen, intravenous administration of 50 Units/kg VORAXAZE 2 hours before leucovorin reduced (6S)-leucovorin AUC0-3h by 33% and Cmax by 52%, and also reduced its active metabolite, (6S)-5-methyltetrahydrofolate, AUC0-3h by 92% and Cmax by 93% [see Drug Interactions (7.1)].

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
VORAXAZE has not been evaluated in animals for carcinogenic or mutagenic potential or for impairment of fertility.

14 CLINICAL STUDIES
The efficacy of VORAXAZE was evaluated in a subset consisting of 22 treatment-evaluable patients enrolled in Study 1. Study 1 was a single-arm, open-label study in patients who had markedly delayed methotrexate clearance (defined as more than 2 standard deviations greater than the mean excretion curve for methotrexate) secondary to renal dysfunction. All patients received VORAXAZE 50 Units/kg as an intravenous injection over 5 minutes; those patients with pre-VORAXAZE methotrexate concentrations >100 μmol/L were to receive a second dose of VORAXAZE 48 hours after the first dose. The protocol specified that patients continue receiving intravenous hydration, urinary alkalinization and leucovorin, and that leucovorin administration be adjusted to ensure that it was not administered within two hours before or after VORAXAZE.

Efficacy was evaluated in a subset of patients enrolled in Study 1 who met the inclusion criteria for the study, had a pre-VORAXAZE methotrexate concentration >1 μmol/L, and had both pre- and post-treatment plasma samples available for determination of methotrexate concentration by a chromatographic method analysis. The main outcome measure was the proportion of patients who achieved a rapid and sustained clinically important reduction (RSCIR) in plasma methotrexate concentration, defined as an attainment of plasma methotrexate concentration ≤1 μmol/L at 15 minutes that was sustained for up to 8 days following the initial injection.

Of the 22 patients in the efficacy dataset, the median age was 15.5 years (5 to 84 years); 59% were male, and the most common underlying cancers were osteogenic sarcoma (50%) and leukemia or lymphoma (45%).

Ten of the 22 patients achieved RSCIR (45% [95% CI 27, 65%]). Of the 12 patients who failed to achieve RSCIR, 5 patients (23%) attained a transient plasma methotrexate concentration of ≤1 μmol/L. In these 5 patients, the median increase of plasma methotrexate concentration from their nadir was 1.4 μmol/L (0.3 to 2.5 μmol/L).

Table 2 summarizes the results of RSCIR and exploratory analyses following the first dose administration of VORAXAZE. An exploratory analysis in subgroups determined by pre-VORAXAZE methotrexate concentration suggests that the likelihood of attaining a RSCIR following the first VORAXAZE injection correlates with the pre-VORAXAZE methotrexate concentration (Table 2). In an additional exploratory analysis, all 9 patients with pre-VORAXAZE methotrexate concentrations >50 μmol/L achieved greater than a 95% reduction in methotrexate concentrations for up to 8 days following the initial injection of VORAXAZE although none of them achieved a RSCIR.

### Table 2: Results of RSCIR and Exploratory Analyses Following the First Dose of VORAXAZE

<table>
<thead>
<tr>
<th>Pre-VORAXAZE Methotrexate Concentration (μmol/L)</th>
<th>Number of Patients</th>
<th>Patients Achieving RSCIR (%)</th>
<th>Patients with &gt;95% Rapid Reduction in Methotrexate Concentration and Maintained up to 8 Days (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;1</td>
<td>22</td>
<td>10 (45%)</td>
<td>20 (91%)</td>
</tr>
<tr>
<td>&gt;1 to ≤50</td>
<td>13</td>
<td>10 (77%)</td>
<td>11 (85%)</td>
</tr>
<tr>
<td>&gt;50 to ≤100</td>
<td>2</td>
<td>0</td>
<td>2 (100%)</td>
</tr>
<tr>
<td>&gt;100</td>
<td>7</td>
<td>0</td>
<td>7 (100%)</td>
</tr>
</tbody>
</table>

RSCIR: rapid and sustained clinically important reduction in methotrexate concentration.

Lack of Efficacy with a Second Dose of VORAXAZE
Six of the seven patients with pre-first dose VORAXAZE methotrexate concentrations >100 μmol/L received a second 50 Units/kg dose of VORAXAZE administered 48 hours after the first dose. Among them, none of the four patients with pre-second dose VORAXAZE methotrexate concentrations >1 μmol/L achieved a RSCIR. The remaining two patients achieved a RSCIR but their pre-second dose VORAXAZE methotrexate concentrations were already ≤1 μmol/L.

Deaths Attributable to Methotrexate Toxicity
There are no controlled trials comparing VORAXAZE plus supportive care to supportive care measures alone in patients with toxic plasma methotrexate concentrations due to impaired renal function; therefore there are no data regarding the effect of VORAXAZE on survival or toxic deaths due to methotrexate. VORAXAZE did not prevent fatal methotrexate toxicity in 3% of patients in the safety population.

16 HOW SUPPLIED/STORAGE AND HANDLING
VORAXAZE is supplied as a sterile, preservative-free white lyophilized powder in an individually packaged glass vial closed with a bromo butyl elastomeric stopper and blue flip-off seal.

1,000 Units of glucarpidase per vial (1 vial per carton) NDC 50633-210-11

Store VORAXAZE at 36°F to 46°F (2°C to 8°C). Do not freeze. Do not use VORAXAZE after the expiration date on the vial.

17 PATIENT COUNSELING INFORMATION
- Inform patients that allergic reactions, including potentially serious reactions, may occur during VORAXAZE treatment.
- Advise patients to immediately report any signs and symptoms of infusion reactions such as fever, chills, flushing, feeling hot, rash, hives, itching, throat tightness or breathing problems, tingling, numbness, or headache.
- Inform patients of the importance of continued monitoring of methotrexate blood concentrations and renal status at the appropriate times after discharge from the hospital.