Mpx

Guidance for Tecovirimat Use

Updated June 7, 2023

What You Need to Know

- Tecovirimat (also known as TPOXX or ST-246) is FDA-approved for the treatment of human smallpox disease caused by variola virus in adults and children. The use of tecovirimat for the treatment of other orthopoxvirus infections, including mpx, remains unapproved at this time.

- However, the National Institute of Allergy and Infectious Diseases, part of the National Institutes of Health, is now enrolling patients in the Study of Tecovirimat for Human Mpx Virus (STOMP), which is designed to assess whether tecovirimat is safe and effective for the treatment of mpx in people with the disease.

- Providers are encouraged to inform patients with mpx about STOMP and to recommend they consider enrollment in STOMP.

- Access to oral tecovirimat is also available for patients with mpx who meet eligibility criteria (e.g., have severe disease or involvement of anatomic areas that might result in serious sequelae or are at high risk for severe disease) under CDC’s expanded access Investigational New Drug (EA-IND) protocol.

Treatment Considerations

The ongoing mpx outbreak in the United States is caused by Clade Iib of the virus that causes mpx. Patients with mpx benefit from supportive care and pain control that is implemented early in the illness (Clinical Considerations for Pain Management of Mpx). Illness depends on a person’s immune response. For most patients with intact immune systems, supportive care and pain control may be enough. However, because prognosis depends on multiple factors, such as initial health status, concurrent illnesses, previous vaccination history, and comorbidities, supportive care and pain control may not be enough for some patients (for example, those with weakened immune systems).

Data on the effectiveness of tecovirimat in treating people with mpx are not available but studies using a variety of animal species have shown that tecovirimat is effective in treating disease caused by orthopoxviruses. In animal studies, tecovirimat has been shown to decrease the chance of dying from infections with orthopoxviruses when given early in the disease course. A clinical trial that focused on safety in healthy people without mpx virus infection showed the drug had an acceptable safety profile; the effectiveness of tecovirimat was not studied in this trial.

Data from the published literature and additional recently released data from the U.S. Food and Drug Administration suggest that there may be a low barrier to virus developing resistance to tecovirimat; indiscriminate use could promote resistance and render tecovirimat, first line treatment for orthopoxviruses, ineffective for patients. Alternate therapeutics have more concerning safety profiles than tecovirimat.

When considering the use of tecovirimat, clinicians and patients should understand 1) the lack of tecovirimat effectiveness data to date in people with mpx, 2) the lack of data indicating which patients might benefit the most from tecovirimat, and 3) the concern for the development of resistance to tecovirimat, which could render the drug ineffective for any treated patients.

Tecovirimat should be considered for use in people who have the following clinical manifestations:

- Severe disease—consider severe disease when a patient has conditions such as hemorrhagic disease; large number of lesions such that they are confluent; sepsis; encephalitis; ocular or periorbital infections; or other conditions requiring hospitalization
Involvement of anatomic areas which might result in serious sequelae that include scarring or strictures—these include lesions directly involving the pharynx causing dysphagia, inability to control secretions, or need for parenteral feeding; penile foreskin, vulva, vagina, urethra, or rectum with the potential for causing strictures or requiring catheterization; anal lesions interfering with bowel movements (for example, severe pain); and severe infections (including secondary bacterial skin infections), especially those that require surgical intervention such as debridement.

Tecovirimat should also be considered for use in people who are at high risk for severe disease:

- People currently experiencing severe immunocompromise due to conditions such as advanced or poorly controlled human immunodeficiency virus (HIV), leukemia, lymphoma, generalized malignancy, solid organ transplantation, therapy with alkylating agents, antimetabolites, radiation, tumor necrosis factor inhibitors, or high-dose corticosteroids, being a recipient of a hematopoietic stem cell transplant <24 months post-transplant or ≥24 months but with graft-versus-host disease or disease relapse, or having autoimmune disease with immunodeficiency as a clinical component.
- Pediatric populations, particularly patients younger than 1 year of age.
- Pregnant or breastfeeding people.
- People with a condition affecting skin integrity—conditions such as atopic dermatitis, eczema, burns, impetigo, varicella zoster virus infection, herpes simplex virus infection, severe acne, severe diaper dermatitis with extensive areas of denuded skin, psoriasis, or Darier disease (keratosis follicularis).

For patients at high risk for progression to severe disease, tecovirimat should be administered early in the course of illness along with supportive care and pain control.

More Details: Treatment Information for Healthcare Professionals

People who are ineligible for tecovirimat treatment under the EA-IND include:

- Patients (or their legally authorized representatives) who are unwilling to sign an informed consent and refuse tecovirimat treatment.
- Patients with known allergy to tecovirimat and/or inactive ingredients of tecovirimat formulation.

Available Formulations

Tecovirimat is available as an oral capsule (200 mg) and injection for intravenous (IV) administration. Drug absorption of oral formulation is dependent on adequate concurrent intake of a full, fatty meal.

IV tecovirimat should not be administered to patients with severe renal impairment (CrCl <30 mL/min). Oral formulation remains an option for this population. IV tecovirimat should be used with caution in patients with moderate (CrCl 30-49 mL/min) or mild (CrCl 50-80 mL/min) renal impairment as well as in pediatric patients < 2 years of age given immature renal tubular function.

Adverse Reactions

Oral
- Headache (12%)
- Nausea (5%)
- Abdominal pain (2%)
- Vomiting (2%)
- Neutropenia was found in one study participant.

IV
- Infusion site pain (73%)
- Infusion site swelling (39%)
- Infusion site erythema (23%)
- Infusion site extravasation (19%)
- Headache (15%)
Drug-Drug Interactions

Significant interactions have been reported in healthy adults with co-administration of repaglinide (hypoglycemia) and midazolam (decreased effectiveness of midazolam).

Special Populations

Pregnancy/Lactation

Although tecovirimat has not been studied in pregnant and nursing women, they are not excluded from treatment if deemed appropriate following careful clinical assessment and discussion of risks/benefits with patient using a shared decision-making model. There are no human data to establish the presence or absence of tecovirimat-associated risk of fetotoxicity, effect on milk production, the presence of drug in human milk, and/or effects on breastfed children. No fetotoxicity was found in animal studies, though tecovirimat was detected in trace amounts in milk.

Pediatrics

Tecovirimat has been used in a 28-month-old child with no adverse effects attributed to the drug, but no clinical studies have been done in pediatric populations. Monitoring of renal function is recommended in pediatric patients <2 years of age, given theoretical concerns that renal immaturity in young pediatric patients may result in higher exposure of hydroxypropyl-β-cyclodextrin, an ingredient in IV tecovirimat. Animal studies have shown potential for nephrotoxicity at very high exposure levels of hydroxypropyl-β-cyclodextrin.

References


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