

UTILIZATION MANAGEMENT MEDICAL POLICY

POLICY: Oncology (Injectable) – Elahere Utilization Management Medical Policy

- Elahere® (mirvetuximab soravtansine-gynx intravenous infusion – ImmunoGen/AbbVie)

REVIEW DATE: 05/06/2026

OVERVIEW

Elahere, a folate receptor alpha (FR α)-directed antibody and microtubule inhibitor conjugate, is indicated for the treatment with FR α positive, platinum-resistant epithelial ovarian, fallopian tube, or primary peritoneal cancer, in adults who have received one to three prior systemic treatment regimens.¹

Dosing Information

The recommended dose of Elahere is 6 mg/kg adjusted ideal body weight (AIBW) administered once every 3 weeks (21-day cycle) as an intravenous infusion until disease progression or unacceptable toxicity.¹ The total dose of Elahere is calculated based on each patient's AIBW using the following formula:

$AIBW = \text{Ideal body weight (IBW [kg])} + 0.4 * (\text{Actual weight [kg]} - \text{IBW [kg]})$

The formula to calculate female IBW is: $\text{Female IBW (kg)} = 0.9 * (\text{height [cm]}) - 92$

Guidelines

The National Comprehensive Cancer Network (NCCN) ovarian cancer (including fallopian tube cancer and primary peritoneal cancer) clinical practice guidelines (version 4.2026 – April 10, 2026) recommend a variety of treatment options as recurrence therapy for platinum-resistant disease.² Single-agent Elahere is listed as a “Preferred” targeted therapy for FR α -expressing tumors ($\geq 75\%$ positive tumor cells) [category 1] for platinum-resistant disease. Elahere + bevacizumab is listed under “Useful in Certain Circumstances” for FR α -expressing tumors ($\geq 25\%$ positive tumor cells) [category 2A]. Elahere is also recommended for platinum-sensitive disease as “Useful in Certain Circumstances” for FR α -expressing tumors ($\geq 75\%$ positive tumor cells) as monotherapy (category 2A) and FR α -expressing tumors ($\geq 50\%$ positive tumor cells) in combination with bevacizumab (category 2B).

POLICY STATEMENT

Prior Authorization is recommended for medical benefit coverage of Elahere. Approval is recommended for those who meet the **Criteria** and **Dosing** for the listed indication. Extended approvals are allowed if the patient continues to meet the Criteria and Dosing. Requests for doses outside of the established dosing documented in this policy will be considered on a case-by-case basis by a clinician (i.e., Medical Director or Pharmacist). All approvals are provided for the duration noted below. Because of the specialized skills required for evaluation and diagnosis of patients treated with Elahere as well as the monitoring required for adverse events and long-term efficacy, approval requires Elahere to be prescribed by or in consultation with a physician who specializes in the condition being treated.

Automation: None.

RECOMMENDED AUTHORIZATION CRITERIA

Coverage of Elahere is recommended in those who meet the following criteria:

FDA-Approved Indication

- Ovarian, Fallopian Tube, or Primary Peritoneal Cancer.** Approve for 1 year if the patient meets ALL of the following (A, B, and C):
 - Patient is ≥ 18 years of age; AND
 - Patient has folate receptor alpha positive disease; AND
 - The medication will be prescribed by or in consultation with an oncologist.

Dosing. Approve up to 6 mg/kg adjusted ideal body weight administered once every 3 weeks (21-day cycle).

Note: To calculate adjusted ideal body weight (AIBW), use the following equation:

$AIBW = \text{Ideal body weight (kg)} + 0.4 * (\text{Actual weight [kg]} - \text{ideal body weight [kg]});$

To calculate female ideal body weight (kg) = $0.9 * (\text{height [cm]}) - 92$

CONDITIONS NOT RECOMMENDED FOR APPROVAL

Coverage of Elahere is not recommended in the following situations:

- Coverage is not recommended for circumstances not listed in the Recommended Authorization Criteria. Criteria will be updated as new published data are available.

REFERENCES

- Elahere® intravenous infusion [prescribing information]. North Chicago, IL: ImmunoGen/AbbVie; July 2025.
- The NCCN Ovarian Cancer Including Fallopian Tube Cancer and Primary Peritoneal Cancer Clinical Practice Guidelines in Oncology (version 4.2026 – April 10, 2026). © 2026 National Comprehensive Cancer Network. Available at: <http://www.nccn.org>. Accessed on April 29, 2026.

HISTORY

Type of Revision	Summary of Changes	Review Date
Early Annual Revision	Ovarian, Fallopian Tube, or Primary Peritoneal Cancer: Additional criteria were added to the requirement of folate receptor positive disease, which are patient has to have either $\geq 75\%$ folate receptor alpha positive tumor cells or patient is using this medication in combination with bevacizumab. The requirement that the patient has tried one systemic regimen and the note of examples of a systemic regimen were removed.	06/05/2024
Annual Revision	Ovarian, Fallopian Tube, or Primary Peritoneal Cancer: The requirement that the patient has either $\geq 75\%$ folate receptor alpha positive tumor cells or is using Elahere in combination with bevacizumab was removed. The requirement that the patient has platinum-resistant disease was removed.	06/04/2025
Annual Revision	No criteria changes	05/06/2026