



You are cordially invited to attend a non-CME educational program

A New 2L+ Combination Therapy for Adults With RRMM After a PI and an Immunomodulatory Agent



Presented by

Subrena Powell, Nurse Practitioner

Oncology Clinical Educator, an employee of Johnson & Johnson

Date

**Tuesday, October 20, 2026
at 6:00 PM Eastern Time**

Location

**Orsay
3630 Park Street,
Jacksonville, Florida 32205
(904) 381-0909**

We look forward to your participation in this informative discussion. Reserve your spot now by contacting your Johnson & Johnson Representative, Subrena Powell at spowell12@its.jnj.com (813) 955-4842, or visit <https://janssenspeakerprograms.my.site.com/Registration/s?l=a12vt00000Fxabv>

Please RSVP one (1) week prior to the program date

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INDICATIONS AND USAGE

TECVAYLI® (teclistamab-cqyv) is a bispecific B-cell maturation antigen (BCMA)-directed CD3 T-cell engager indicated for the treatment of adult patients with relapsed or refractory multiple myeloma:

- in combination with daratumumab and hyaluronidase-fig in patients who have received at least one prior line of therapy, including a proteasome inhibitor and an immunomodulatory agent.
- as monotherapy, in patients who have received at least four prior lines of therapy, including a proteasome inhibitor, an immunomodulatory agent and an anti-CD38 monoclonal antibody.

IMPORTANT SAFETY INFORMATION

WARNING: CYTOKINE RELEASE SYNDROME and NEUROLOGIC TOXICITY including IMMUNE EFFECTOR CELL-ASSOCIATED NEUROTOXICITY SYNDROME. Cytokine release syndrome (CRS), including life-threatening or fatal reactions, can occur in patients receiving TECVAYLI. Initiate treatment with TECVAYLI step-up dosing schedule to reduce risk of CRS. Withhold TECVAYLI until CRS resolves or permanently discontinue based on severity. Neurologic toxicity, including Immune Effector Cell-Associated Neurotoxicity Syndrome (ICANS) and serious, life-threatening, or fatal reactions, can occur in patients receiving TECVAYLI. Monitor patients for signs or symptoms of neurologic toxicity, including ICANS, during treatment. Withhold TECVAYLI until neurologic toxicity resolves or permanently discontinue based on severity.

TECVAYLI is available only through a restricted program called the TECVAYLI and TALVEY® Risk Evaluation and Mitigation Strategy (REMS).

Please read additional Important Safety Information on page 2 and accompanying full Prescribing Information, including Boxed WARNING, for TECVAYLI.