

MYALEPT® REMS Program

Program Requirements | Training & Enrollment

A Risk Evaluation and Mitigation Strategy (REMS) is a strategy to manage known or potential serious risks associated with a drug product and is required by the Food and Drug Administration (FDA) to ensure that the benefits of a drug outweigh its risks.

The MYALEPT REMS Program was developed with the FDA:

To educate prescribers about:

- the development of anti-metreleptin antibodies that neutralize endogenous leptin and/or MYALEPT and the serious adverse events that may result from these antibodies,
- the risk of lymphoma, and
- appropriate patient selection

Program Requirements

MYALEPT is available only through the MYALEPT REMS Program. The MYALEPT REMS Program requirements include:

For Prescribers:

- **Certification** of prescribers of MYALEPT
 - Certification consists of completion of **training, and enrollment** in the MYALEPT REMS Program
- **Completion** of a Prescription Authorization form for **each** new prescription.

Find out more about [Training & Enrollment](#).

For Pharmacies:

- **Restricted distribution** of MYALEPT to patients with completed Prescription Authorization Forms from prescribers who are certified in the MYALEPT REMS Program.

Training & Enrollment

Healthcare providers who prescribe MYALEPT must review the prescriber training materials to enroll in the MYALEPT REMS Program.

Steps to Prescriber Certification

1. Review the Prescriber Education Materials

- [MYALEPT Prescribing Information](#)
- [Prescriber Training Module](#)

2. Complete and submit the MYALEPT REMS Program Prescriber Enrollment Form

- Print and sign the [Prescriber Enrollment Form](#) or request a copy by calling 1-855-669-2537
- Submit the form via Fax to 1-877-328-9682

By completing the Prescriber Enrollment Form, the prescriber agrees to comply with the MYALEPT REMS Program requirements. A confirmation of your certification in the MYALEPT REMS program will be sent to you so you can begin to prescribe MYALEPT.

Reporting Adverse Reactions

Healthcare providers should report all suspected adverse events.

Please contact the company at 1-855-669-2537 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.com.

REMS Materials

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-  [Prescribing Information](#)
-  [Myalept REMS Letter for HCPs \(Infections\)](#)

MYALEPT REMS Program Contact Us

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Fax: 1-877-328-9682

Hours of Operation:
Monday - Friday,
8:00am-8:00pm
Eastern Time

MYALEPT[®] REMS Program

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