



LEMTRADA REMS PATIENT AUTHORIZATION AND BASELINE LAB FORM

**Please fax this completed form to the LEMTRADA REMS at 1-855-557-2478
or submit online at www.LemtradaREMS.com**

This form must be completed within 30 days prior to the first infusion date of each LEMTRADA® (alemtuzumab) patient's treatment course.

*Indicates a mandatory field.

PRESCRIBER INFORMATION (PLEASE PRINT)

Name (Last, First)*		Office Phone Number*
Address*		
City*	State*	ZIP Code*
Prescriber LEMTRADA REMS Identification Number*		

PATIENT INFORMATION (PLEASE PRINT)

Name (Last, First)*
Patient LEMTRADA REMS Identification Number*
Date of Birth (MM/DD/YYYY)*

AUTHORIZATION AND BASELINE LABS

Do you authorize LEMTRADA treatment for the above-referenced patient? Yes No

Do you attest that required baseline laboratory testing has been completed prior to LEMTRADA treatment and within 30 days of the patient's first infusion? Yes No

PRESCRIPTION INFORMATION

Check one* Initial course (1 vial [12 mg/day]) X 5 consecutive days Total number of vials: _____

Subsequent course (1 vial [12 mg/day]) X 3 consecutive days Total number of vials: _____

SIGNATURE

Prescriber Signature* Date*

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If you have any questions regarding the LEMTRADA REMS, call 1-855-676-6326