

Specialty pharmacy and distributors list



The LUMAKRAS™ network gives providers and patients multiple dispensing options



Specialty Pharmacies (SPs)



In-office Dispensing Pharmacies

Specialty pharmacy network

LUMAKRAS is available for order from these specialty pharmacies who also provide support to help patients with their prescribed treatments:

Specialty pharmacy	Phone number	Fax number
Biologics	800-850-4306	800-823-4506
Humana	800-720-7258	877-405-7940
AcariaHealth	407-903-1308	833-762-0871
Onco 360	877-662-6633	877-662-6355
Optum Specialty Pharmacy	877-445-6874	877-342-4596
Accredo Health Group	For patients: 877-732-3431 MDO ONLY: 877-783-2262	866-352-3973
CVS Specialty	888-280-1193	855-296-0210
US Bioservices	877-757-0667	888-899-0067

- Check with your patient's insurance for specific coverage requirements and specialty pharmacy mandates.
- Payer and PBM specialty pharmacy mandates may apply regardless of the manufacturer-defined specialty pharmacy network.

Specialty distributors network

These authorized specialty distributors stock and sell LUMAKRAS:

Specialty distributor	Phone number	Website
ASD Healthcare	800-837-5403	www.asdhealthcare.com
Oncology Supply	800-633-7555	www.oncologysupply.com
Cardinal Health SPD – Hospital & SP's	855-855-0708	www.cardinalhealth.com
Cardinal Health SPD - Clinics	877-453-3972	www.cardinalhealth.com
McKesson Plasma and Biologics	877-625-2566	connect.mckesson.com
McKesson Specialty Care Distribution	855-477-9800	mscs.mckesson.com/CustomerCenter

This table is current as of 06/30/2021.

The specialty pharmacies and distributors identified can fill and distribute LUMAKRAS and are provided herein for informational purposes only and is not an endorsement or recommendation to use any particular pharmacy or distributor over another. This list may not be exhaustive and is subject to change. It represents no statement, promise, or guarantee by Amgen Inc. concerning coverage and/or availability of the product for any particular patient and it is the responsibility of the health care provider to determine the appropriate services provided to their patients.

Please see page 2 for **Indication and LUMAKRAS™ Important Safety Information**. [Click here for LUMAKRAS™ full Prescribing Information](#).

INDICATION AND IMPORTANT SAFETY INFORMATION

INDICATION

LUMAKRAS is indicated for the treatment of adult patients with *KRAS G12C*-mutated locally advanced or metastatic non-small cell lung cancer (NSCLC), as determined by an FDA-approved test, who have received at least one prior systemic therapy.

This indication is approved under accelerated approval based on overall response rate (ORR) and duration of response (DOR). Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial(s).

IMPORTANT SAFETY INFORMATION

Hepatotoxicity

- LUMAKRAS™ can cause hepatotoxicity, which may lead to drug-induced liver injury and hepatitis.
- Among 357 patients who received LUMAKRAS™ in CodeBreak 100, hepatotoxicity occurred in 1.7% (all grades) and 1.4% (grade 3). A total of 18% of patients who received LUMAKRAS™ had increased alanine aminotransferase (ALT)/increased aspartate aminotransferase (AST); 6% were grade 3 and 0.6% were grade 4. In addition to dose interruption or reduction, 5% of patients received corticosteroids for the treatment of hepatotoxicity.
- Monitor liver function tests (ALT, AST, and total bilirubin) prior to the start of LUMAKRAS™, every 3 weeks for the first 3 months of treatment, then once a month or as clinically indicated, with more frequent testing in patients who develop transaminase and/or bilirubin elevations.
- Withhold, dose reduce, or permanently discontinue LUMAKRAS™ based on severity of adverse reaction.

Interstitial Lung Disease (ILD)/Pneumonitis

- LUMAKRAS™ can cause ILD/pneumonitis that can be fatal. Among 357 patients who received LUMAKRAS™ in CodeBreak 100 ILD/pneumonitis occurred in 0.8% of patients, all cases were grade 3 or 4 at onset, and 1 case was fatal. LUMAKRAS™ was discontinued due to ILD/pneumonitis in 0.6% of patients.
- Monitor patients for new or worsening pulmonary symptoms indicative of ILD/pneumonitis (eg, dyspnea, cough, fever). Immediately withhold LUMAKRAS™ in patients with suspected ILD/pneumonitis and permanently discontinue LUMAKRAS™ if no other potential causes of ILD/pneumonitis are identified.

Most Common Adverse Reactions

- The most common adverse reactions $\geq 20\%$ were diarrhea, musculoskeletal pain, nausea, fatigue, hepatotoxicity, and cough.

Drug Interactions

- Advise patients to inform their healthcare provider of all concomitant medications, including prescription medicines, over-the-counter drugs, vitamins, dietary and herbal products.
- Inform patients to avoid proton pump inhibitors and H₂ receptor antagonists while taking LUMAKRAS™.
- If coadministration with an acid-reducing agent cannot be avoided, inform patients to take LUMAKRAS™ 4 hours before or 10 hours after a locally acting antacid.

Reference: 1. LUMAKRAS™ (sotorasib) prescribing information, Amgen.

Please see LUMAKRAS™ full [Prescribing Information](#).

For more information, visit [LUMAKRASHCP.com](https://www.lumakrashcp.com)

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LUMAKRAS[™]
(sotorasib) 120 mg tablets

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