

DRUG & BIOLOGICAL PRODUCTS

To ensure success in today's rapidly evolving biopharmaceutical industry, companies must recognize and be prepared for a dynamic regulatory landscape. Guided by decades of experience, Greenleaf's team of experts provides unmatched knowledge of the life sciences regulatory process and serves as a trusted partner for companies navigating the complexities of product lifecycle management.

GREENLEAF'S APPROACH

- Greenleaf's in-depth knowledge and understanding of the U.S. Food and Drug Administration (FDA) as a regulator provides clients with a trusted partner when navigating the complex process of bringing new therapeutics to market. The team's approach, firmly grounded in established principles of public health practices, is guided by decades of regulatory experience in drug and biological products development, spanning all therapeutic areas.
- The Drug and Biological Products Team delivers a variety of services from the earliest stages of product development through postapproval commitments. Services include monitoring and assessing the regulatory environment for emerging trends, analyzing the impact of agency actions on current development programs, and reviewing the competitive landscape for specific therapeutic areas.
- Greenleaf experts specialize in clinical trial design, FDA submissions and the review process, as well as postmarket requirements, including safety monitoring.
- Experts from Greenleaf's Drug and Biological Products Team also provide advisory services that include extensive research and due diligence for firms engaged in potential mergers and acquisitions that require regulatory risk assessments before and after a life sciences transaction.

ABOUT GREENLEAF

Greenleaf Health is a leading FDA regulatory consulting firm that provides strategic and technical guidance to pharmaceutical and medical device companies researching, developing and manufacturing innovative solutions to pressing global public health challenges.

COMPREHENSIVE SERVICES

Greenleaf's Drug and Biological Products Team has a robust blend of regulatory expertise and FDA institutional knowledge, providing strategic and technical guidance in an array of areas, including clinical trial design, FDA filings and review process, and postmarket requirements such as safety monitoring.

