



OUR EXPERIENCE. YOUR SUCCESS.

COMPREHENSIVE SERVICES & SUPPORT

Greenleaf's comprehensive services and wealth of experience ensure that clients can count on expert direction as they encounter complex regulatory challenges. Greenleaf professionals work as teams specializing in product quality, manufacturing, and compliance; medical devices and combination products; and drug and biological products.



Product Quality, Manufacturing & Compliance

Greenleaf's Product Quality, Manufacturing, and Compliance Team provides assistance and support to companies, trade associations, and other stakeholders regulated by the U.S. Food and Drug Administration (FDA). Greenleaf experts identify and promote practices that will align a client's approach with the FDA's quality, safety, and compliance expectations.



Medical Device & Combination Products

Greenleaf's Medical Device and Combination Products Team guides clients through the complex regulatory process, from the earliest stages of product development, through the FDA review process, to marketing authorization and compliance with postmarket requirements and quality systems.



Drug & Biological Products

Greenleaf serves as a trusted partner when maneuvering the sophisticated process of bringing new therapeutics to market. The Drug and Biological Products Team specializes in providing strategic and technical guidance on medical product development, regulatory review, and postmarket requirements.

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

GREENLEAF'S APPROACH

Greenleaf is committed to serving our clients' needs with extensive expertise, unwavering integrity, and strategic insight in a manner that supports availability of safe, effective, and high-quality drugs, biologics, and devices.

TEAM OF EXPERTS

Greenleaf's team of advisors offers a rare blend of perspectives developed as leaders in both the public and private sectors. This wealth of experience informs Greenleaf's understanding of the broad life sciences industry and enables us to deliver valuable insight throughout the product lifecycle.

REGULATORY SERVICES

The firm's targeted regulatory capabilities include:

Strategic and technical guidance for medical product development and regulatory review

Product quality, manufacturing, and compliance

FDA meeting preparation and communication

Advisory Services

Medical product labeling and promotion

Regulatory policy consultation, strategic planning, and communications

Compliance assessments, remediation, and inspection readiness

Greenleaf is a regulatory consulting firm and does not provide legal advice or legal services.

UNMATCHED REGULATORY EXPERTISE

Greenleaf's decades of hands-on experience in the regulatory space is unmatched. The firm's team of experts includes former leaders and regulatory professionals from the FDA, Capitol Hill, top global pharmaceutical and medical device companies, leading law firms, and the top U.S. biotechnology trade organization.

DISTINGUISHED EXPERIENCE & LEADERSHIP

Greenleaf's team is comprised of experts with a combined total of more than 300 years of FDA experience.



JOHN TAYLOR

President and Principal, Compliance & Regulatory Affairs

John Taylor joined Greenleaf following a distinguished FDA career of more than 20 years. Taylor served in many high-profile positions at the FDA, as well as in senior leadership roles within industry.



KATHLEEN SONNTAG

Chief Operating Officer

Kathleen Sonntag is a veteran in the financial services field, with a broad career emphasizing information technology, data analysis, and the use of technology to enable business processes.



JOHN JENKINS, M.D.

Principal, Drug & Biological Products

With a 25-year career at the FDA, including 15 years in senior leadership positions within the Center for Drugs, Dr. Jenkins is an expert in the statutes and regulations that guide drug development.



BOB MEYER, M.D.

Principal, Drug & Biological Products

Dr. Meyer brings more than 25 years of regulatory, industry, and academic leadership to Greenleaf, including prominent roles at the FDA, Merck, and the University of Virginia.



KAREN MIDTHUN, M.D.

Principal, Drug & Biological Products

An infectious disease physician by training, Dr. Midthun's 28-year career in public service includes her role as Director of the FDA's Center for Biologics Evaluation and Research (CBER).



DANIEL SCHULTZ, M.D.

Principal, Medical Devices & Combination Products

Dr. Schultz's distinguished 35-year public service career includes his role as a member of the U.S. Public Health Service and as Director of the FDA's Center for Devices (CDRH).



KATE COOK

Principal, Regulatory Policy

Kate Cook joined Greenleaf following more than two decades with the FDA leading regulatory policy development and serving as legal counsel on biological, medical device, and drug issues.



DAVID ELDER

Principal, Regulatory Compliance

A 23-year veteran of the FDA, David Elder served as a senior FDA official with prominent roles in domestic and foreign inspections, recalls and emergencies, and compliance actions.



CYNTHIA SCHNEDAR

Principal, Regulatory Compliance

Cynthia Schnedar's 25-year compliance career includes her role as Director of the Office of Compliance for the FDA's Center for Drug Evaluation and Research (CDER).

COMPREHENSIVE SERVICES

Members of the Greenleaf team work cross-functionally to provide a full-service engagement that ensures clients can count on expert direction as they encounter regulatory challenges. Greenleaf's collaborative services include:

Litigation Support Services

Greenleaf and our network of experts work closely with litigators representing FDA-regulated clients in disputes related to medical devices, drug and biological products, and product quality, manufacturing, and compliance.

Advisory Services

Greenleaf understands the complex environment within which life sciences transactions take place and advises investors on potential issues and regulatory risks that may be identified during such transactions.

Cell & Gene Therapy

Greenleaf assists sponsors of cell and gene therapies by optimizing FDA interactions and submissions to support development, manufacturing, quality, and regulatory review.