Wolters Kluwer provides current, accurate, and authoritative references and professional tools with analysis and insight into virtually every aspect of healthcare law. *Cheetah™ Life Sciences Library* gives you access to practical resources and professional references on everything from maintaining full compliance in all areas of food, drug and cosmetics regulation to the rapidly shifting area of medical device regulation. Given the ever-changing regulatory environment, it’s crucial to have access to current and reliable resources that cover the issues of greatest concern. Using *Cheetah™ Life Sciences Library* helps you make the correct choices, maintain compliance, establish effective policies, and provide reliable guidance using:

- **Expert analysis and authoritative content**—The industry’s leading professionals deliver expert explanations, citations, and annotations covering every key area of life science law, in addition to full text of all laws and regulations.
- **Practice tools**—The FDA Reform Navigator Smart Chart allows users to create a customizable chart of any provision of the FDA Food Safety Modernization Act including summaries of each provision.
- **Current awareness**—Regular updates, reporters, and newsletters keep life science attorneys and professionals completely current on all key developments.

**CCH® Food, Drug and Cosmetics Law Library**

Access analysis and full text of laws and regulations governing food, drugs, biologics, and cosmetics.

The Food, Drug and Cosmetics Law Library is your complete online resource covering the Federal Food, Drug and Cosmetics Act and over 50 related federal laws reported in full text, along with concise explanations and expert analysis, pertinent regulations, proposals, rulings, decisions and other developments.

**Designed for fast research**

The *Library* offers a number of tools for fast information retrieval, including cross-references from headnotes and regulations to related text and from report letter stories to applicable headnotes, source documents and regulations.

**Available in Internet or print format**

Both print and Internet versions provide expert explanations of the proposed and final rules governing FDA-regulated products, citations to relevant laws and regulations, and summary annotations to rulings and interpretations of the law.

Internet subscribers receive the full text of all draft and final guidances published by the FDA and other selected documents. Internet subscribers receive the additional advantage of hyperlinks, which allow rapid point-and-click access to all cross-references.

**The Food, Drug, and Cosmetics Newsletter**—a monthly report on significant changes to laws and regulations, new court cases, and important FDA releases related to food and drugs, is included with your subscription.

**Food, Drug and Cosmetics Law Library** covers the following areas:

- **Administrative**—Federal agency information, rulemaking procedures, and more
- **Enforcement**—Prohibited actions, inspections, packaging, and more
- **Food**—Adulteration, food, safety, and security, labeling, misbranding, and more
- **Human drugs and biological products**—Applications, adulteration, and more
- **OTC drugs**—Labeling, monographs, and more
- **Veterinary drugs**—Applications, medicated animal feeds, and more
- **Controlled substances**—Schedules, quotas, registration, labeling, and more
- **Cosmetics**—Purity, safety, and more
- **Color additives**—Listing, petitions, and more
Cheetah™ Life Sciences Library

CCH® Medical Devices Library
Your resource for comprehensive analysis of law relating to medical, diagnostic and radiological devices.

The Medical Devices Library tracks new developments concerning the regulation of medical device manufacturing and marketing—including classification, standards, inspections, seizures and recalls. The Library provides clear, expert analysis and explanations of device regulation, thoroughly discussing compliance issues and approval for Class I, Class II and Class III devices. Additionally, the Library equips you with the reference tools you need to manage product compliance, including reproductions of required FDA forms.

The Medical Device Newsletter, a monthly report on significant changes to laws and regulations, new court cases, and important FDA releases related to medical devices, is also included in your full subscription.

The Medical Devices Library includes the text of the Federal Food, Drug, and Cosmetics Act, The Public Health Services Act, pertinent portions of other public laws and statutes, as well as FDA regulations.

Current Awareness

Both the Food, Drug and Cosmetics Law Library and Medical Devices Library include newsletters with stories highlighting significant regulations, guidances and other actions by the FDA. Stories in the newsletters are linked to primary source content.

Both libraries also include Tracker—a simple and dynamic way to deliver the latest food, drug, and devices primary source content via email every business day.

Choose from a wide variety of food and drug news topics, from advertising to inspections to misbranding. Then, determine what types of documents you need, whether it’s court cases, federal register notices, or guidance documents. We deliver the news to you via e-mail. It’s fast, easy, and convenient.

Tracker allows you to:

• Set up searches once and get results as often as the database is updated. Receive the results via e-mail.
• View everything on any given topic you’ve chosen by a specific date, or over a select period of time.
• “Select All” categories and get anything e-mailed to you that is added to the database.
• Get short summaries of new documents so you can decide whether they are worth accessing.
• Link directly to an entire document right from the Tracker message.
• Search by topic, by document type, or both.

Food, Drug and Cosmetics Tracker covers:
• Administration of FD&C Act
• Drug Approvals
• Food Safety
• Inspections
• Misbranding
• Packaging and Labeling
• Safety and Effectiveness and many other topics

Medical Devices Tracker covers:
• Device Classification
• Enforcement
• Good Manufacturing Practices
• International Harmonization
• Premarket Notification
• User Fees
• Warnings and many other topics

For more information, please call 800-449-6435 or visit us at: wolterskluwerlr.com/cheetah
Other health law resources available from Wolters Kluwer

**Health Law Daily**

*Wolters Kluwer Health Law Daily*—a daily reporting service created by attorneys, for attorneys—delivers breaking news, court decisions, legislation, and regulatory developments.

**Cheetah™ Healthcare Law**

*Cheetah™ Healthcare Law* gives you access to practical resources and professional references on everything from maintaining full compliance in all areas of healthcare regulation to managing reimbursement efficiently and effectively. Given the ever-changing health care environment, it’s crucial to have access to current and reliable resources that cover the issues of greatest concern. Using Cheetah™ Healthcare Law, you can make the correct choices, maintain compliance, establish effective policies, and provide reliable guidance.

**Medicare and Medicaid Guide**

*Medicare and Medicaid Guide* is the leading resource covering reimbursement for health care services and compliance. Its easy-to-use format provides the primary source information you need, and the expert guidance for using that knowledge to your best advantage.

**State Health Law Library**

*State Healthcare Law Library* (including State Medicaid) is a comprehensive source of state-by-state healthcare law information. This Library contains all of the authoritative material needed to keep on top of the latest changes in today’s fast moving legislative and regulatory arenas. This fully searchable, electronic resource brings full text reporting of health care laws and regulations for each of the 50 states and the District of Columbia right to your desktop. Customize your state health care library by choosing only the state(s) you need.

For more information, please call 800-449-6435 or visit us at: wolterskluwerlr.com/cheetah