Obtain visibility throughout the entire labeling process
Meet regulations and requirements with a compliant solution
Automate workflows to improve process efficiency
Manage processes more effectively with unified data and interfaces

Labeling Process and Material Management, a critical element of Regulatory Operations, is a complex activity that demands thorough attention to detail throughout the product development and commercialization lifecycle. The labeling and material management process requires specialists’ input and careful coordination across organizations and multiple functions to ensure accuracy and promote the quality and safety of the delivered product.

Coupled with the need to meet the diverse and shifting regulatory requirements across global markets, it is no surprise that achieving effective label management remains a major challenge for many life sciences organizations.

How can life sciences organizations more efficiently and successfully manage the labeling and material management process?

MEET THE CHALLENGE
Appian offers a solution that can provide a more comprehensive end-to-end review and management of the labeling process throughout the lifecycle of a pharmaceutical or medical device, from early product development through the end of product life. The system affords all stakeholders clear visibility into the overall process orchestration, with automated workflows that provide timely, transparent views across processes in R&D, Quality, Pharmaco- and Medical Device Vigilance, Regulatory Affairs, Medical Affairs, and Manufacturing & Supply Chain.

Appian’s solution can help organizations:

• Tie together safety findings, health authority citations, manufacturing changes, and new research and development findings.

• Simplify extensive processes by offering situational awareness and clear notifications when there is a threat for bottlenecks, along with opportunities for more efficient solutions.

• Manage the creation and maintenance of the Core Company Data Sheets (CCDS), label text creation, Product Information Leaflets (PILs), and all product related artwork and updates throughout the product life cycle.

• Access process monitoring capabilities for distribution on local, regional, and global levels.

• Manage the creation, splitting, cloning, and merging of submission packages across various jurisdictions at the global, regional and local levels.
FOCUS
The Labeling Process and Material Management application enables life sciences organizations to focus on:

- Ensuring quality information is provided for labeling input
- Keeping up with industry trends and changes
- Easy collaboration for experts from various teams throughout the organization

TAKE CONTROL
Using Appian, you can quickly build, deploy, and scale new regulatory operations enterprise applications, including:

- Product Registration & Submission Management (RIM)
- Regulatory Intelligence Database
- Regulatory Start Up
- Identification of Drug and Medicinal Products (IDMP)
- Medical Devices National Registry

PREPARE FOR THE FUTURE
The future of the life sciences industry depends on its ability to bring the highest quality products to market quickly and cost-effectively.

It takes speed and power to transform the life science product lifecycle. The Appian low-code application platform provides both.

With Appian, organizations can build web and mobile apps faster, run them on the Appian cloud, and manage complex processes, end-to-end, without limitations.

Appian provides a leading low-code software development platform that enables organizations to rapidly develop powerful and unique applications. The applications created on Appian’s platform help companies drive digital transformation and competitive differentiation.

For more information, visit www.appian.com