Product Registration and Submission Management, an important aspect of Regulatory Information Management (RIM), touches almost every step of the life science product lifecycle.

Working with health and regulatory authorities worldwide can present efficiency challenges from the earliest stages of drug or device conception, all the way through product end of life. Issues include data handling and regulatory-related decision making processes. Segmented systems that don’t work together can drive disconnects and decreased visibility between regulatory portfolio planning, milestone tracking, and submission processes.

All this can put companies at risk when it comes to getting products into the hands of patients who need them, while making sure those products are safe for consumption.

How can life sciences organizations assure proper product registration and submission, process transparency, and regulatory compliance?

**MEET THE CHALLENGE**

With Appian, life sciences organizations can gain efficiency and control over their RIM process. Appian can provide the capabilities to manage and establish transparency for multiple levels of registration across all levels, global and regional, and across multiple regulatory jurisdictions.

Built on the Appian low-code application platform, Product Registration and Submission Management can:

- Cut down on repeat data entries with a simplified employee workflow
- Provide real-time visibility into processes
- Reduce global risk with increased visibility and greater control
- Provide flexibility for worldwide regulations
- Speed delivery of new products to market by streamlining processes
FOCUS
The Product Registration and Submission Management application enables life sciences organizations to focus on:

• Submission quality with easier resource planning
• Continuous improvement of regulatory operations
• Addressing new and changing regulations
• Faster, more informed decision making

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

• Regulatory Intelligence Database
• Labeling Process & Materials Management
• 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.

Information management at your fingertips for regulatory associates located all over the world and designed for all associates involved in the regulatory business process.

— HEAD OF GLOBAL REGULATORY OPERATIONS, TOP MULTINATIONAL PHARMA COMPANY