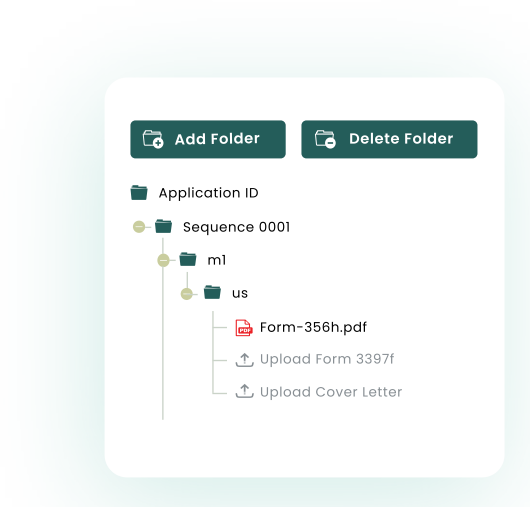


End-to-End Regulatory Affairs Software

What is RIMS?

RIMS, short for Regulatory Information Management System, is a centralized software platform that helps Pharmaceutical and Life Sciences organizations manage regulatory information for their products. It enables end-to-end tracking of Regulatory Activities and reduces efforts to obtain Regulatory Compliance.

The DYAZ RIMS Platform Offers



Template Creation

Pre-built Templates

Build submission plans using fully customizable inbuilt agency complaint eCTD folder structure

Smart, Reusable Content

Create variables in documents for easy and consistent reuse across submissions

Document Management

Collaborative Content Authoring

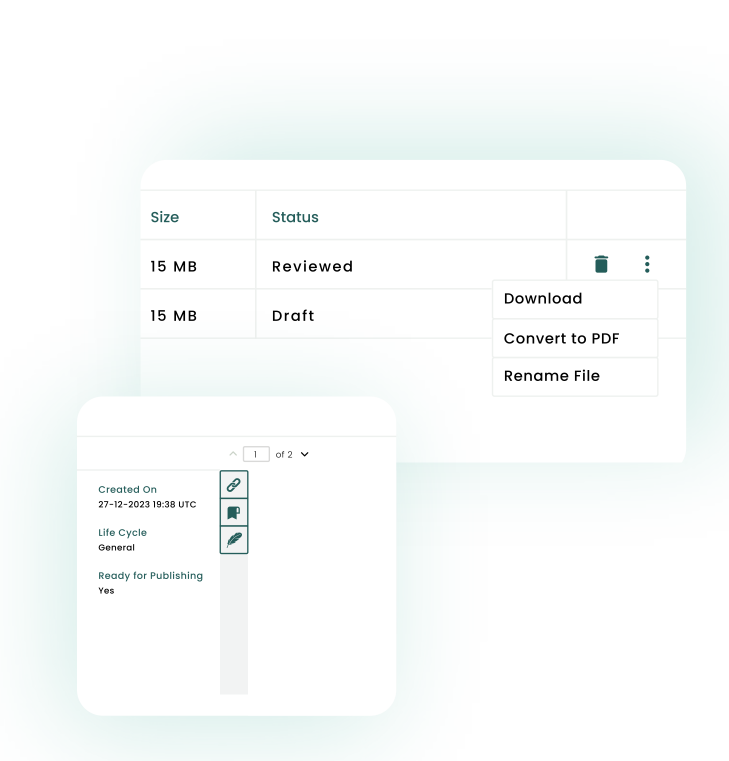
Create, edit, and share structured submission content without external documents or spreadsheets

Onedrive Integration

Open & Edit the document in Onedrive, accompanied by Bookmarking, Hyperlinking and eSignature.

Convert DOC to PDF

Render documents in the correct PDF format with auto-generated TOCs and appendices based on your content plan



Actions Required App 12, Seq 0003 Overdue Revise the target date, contact RA Lead at earliest 01-05-2024 08:25 IST View more **Document Properties:** Created On: 27-12-2023 19:38 UTC Size: Reviewer: **Aaron Grant** Approver: Lifecycle: Replace Link the Document:) Link

eCTD Submissions

Comprehensive Lifecycle Management

Assign tasks, identify document lifecycle operations and navigate among files across all the sequences with the help of our integrated eCTD Viewer

Submission Alerts and Tracking

Receive automated alerts and notifications about your applications and never miss a deadline

Sequence Import & Export

Easy migration of sequences from an external platform and download submission ready sequences

Query Management

Track correspondence with health authorities and manage approval processes

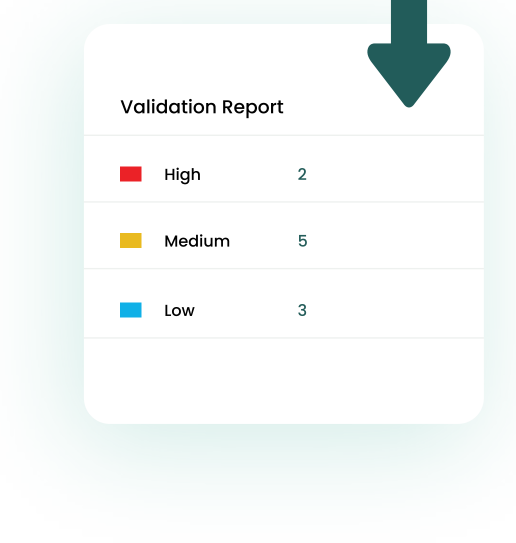
Validations

Reduce Risk of Errors Validate sequences with latest ICH Standards and Regional

Guidelines, preventing technical errors and ensuring on-time approvals

Download Validation Report Review technical errors and handle them within the

system



Benefits of DYAZ RIMS



Manage Roles & Privileges



⊘ Cloud Storage



Flexible Subscriptions

Agency Compliances



Administation

(USFDA)



(EMA)

Economic Community of



West African States (ECOWAS)

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