



**PHARMA** 

Industry



CANADA Location



>**\$2B**Investment in R&D



>100 Clinical Trial Projects

## **Client Requirement**

- Manage and monitor multiple clinical trials across the globe with the highest level of accuracy and precision.
  - Ensure patient safety, accurately assess and report outcomes, and comply with global regulatory
- requirements.
- Organize and store documents in a CDISC-compliant eTMF structure.
- Customize country-specific finance budget sheets.
  - Convert labor-intensive clinical operations
- management into an **effortless and seamless** process by deploying CTMS.



## **Our Solution**

Octalsoft provided a customized & centralized system for capturing trial data, monitoring trial progress, generating reports, and addressing open action items. We currently host a database of 1000+ trials, which help maintain clinical operations and subject-specific data in compliance with the highest regulatory standards. With Octalsoft CTMS, the client:

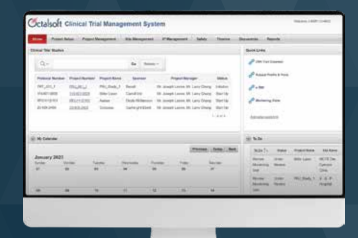
- Made real-time, data-driven decisions, increased the efficiency of the business processes, and contained costs, resulting in timely trial completion and license filings.
- Increased productivity of the CRAS by automating tasks, allowing real-time information sharing, and effective time management.
- Generated an intensive database of verified Clinical Trial Sites and Principal Investigators, giving it a sustainable competitive edge
- Enabled integration of IWRS & EDC for subject-specific critical data points. The data points included but were not limited to subject statuses & visit progression, recruitment efforts, and subject safety.
- Accessed flexible exports, smart reports, dashboards, and multiple reporting tools as per the management requirement.
- ✓ Improved document management and quality processes through Octalsoft eTMF.
- CTMS Finance Module helped with online tracking of pass-through and non-pass-through study costs and investigator milestone payments, and claim management in case of SAES

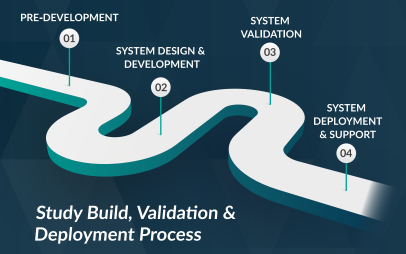




## **Module Features**

- Clinical Site Visit Reports
- Milestones Planning and Tracking
- **Resource Planning and Tracking**
- **Action Items & Deviations**
- Finance Management for Trial Budgeting and **Payments**
- Site Budget and Payments





- Site Monitoring (Site Qualification, Selection, Initiation, and Closure)
- Travel Expenses & Timesheet Tracking
- Regulatory Documents & Compliance Tracking
- Training Management
- Adverse Event Tracking
- Integration with Other eClinical Solutions

## **Our Key Platform Benefits**

Modularized & Integrated Platform With Single Sign In

Unparalleled **Customer Support** 





**Cost Effective** 



**Compliant With Highest Regulatory Standards** 



Simple & Intutive **User Interface** 



To know more, email us at info@octalsoft.com inquiry@octalsoft.com +1 (240) 547-4400 +91 (760) 098-3010 Schedule a **Demo** with our Product Specialist today!

