

# Interactive Web Response Systems



Optimize Inventory Management



Manage Subject Randomization



Simplify Clinical Supply Management



Eliminate Bias

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!





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# **INTERACTIVE WEB RESPONSE SYSTEMS** BENEFITS

#### Recruitment

- 24/7 availability of web interface to quickly register and randomize subjects into your study
- Recruitment controls with study level, stratification level, and country level and site level enrollment caps

#### Randomization

- Configurable, simple to complex randomization codes
- Block-based and centrally or locally managed randomization
  sequences with multiple stratifications
- Manage adaptive study designs and mid-study changes

#### Supply Management

- Automated & manual shipments to suit your operational needs
- Real-time trial supply inventory tracking with predictive reordering algorithms
- Multi-depot modules with depot-to-depot & site-to-site transfers to help sponsors balance IMP distribution, manage stock-outs, utilize near expiry supplies and facilitate ad hoc supplies, thus reducing overall cost, wastage and manual efforts

Octalsoft's CTMS, eTMF and IWRS applications are well-designed and well-integrated systems customized with all contributors and participant inputs and interactions. The team members are hard-working (24X7 available), highly talented who are always thinking out of the box, and the team brings together exceptional skills and invaluable experience.

> Dr Rajiv Yadav Manager-Clinical Research Alkem Laboratories Ltd.

#### Dispensation

- Integrated complex dispensation logic
- Auto-dose calculations including up & down titrations

#### Masking & Unblinding

- User role based controlled access to maintain masking of unblinded information
- Integrated emergent & non-emergent code breaking negates the redundancy of paper envelopes

#### • Real-time & Critical Insights

- Customized automated alerts for critical updates like low level supply trigger, near-to-expiry date trigger, IP not acknowledged by a site beyond a defined period, etc
- Extensive list of reports for blinded and unblinded users with 3D/stack/unstack graphical representation of data with filtration options
- Complete IMP reconciliation

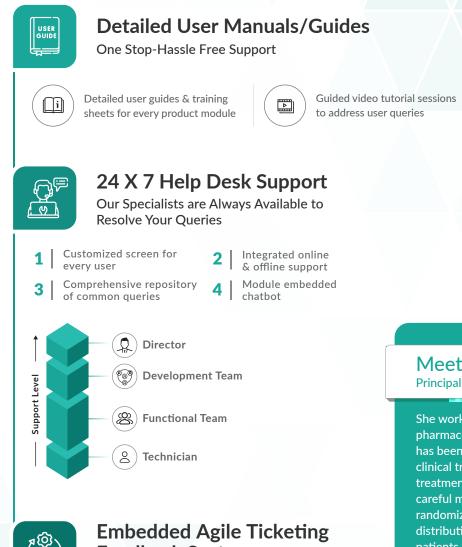
#### Application Programming Interfaces (APIs)

- End-to-end IWRS system integration with EDC and CTMS
- API with third-party applications/systems

#### Rapid Deployment & Quick Support

- Configurated and pre-validated system with abbreviated UAT for faster deployment
- Proactive focus on remaining available and customer-focused throughout the study execution
- Integrated support model with online feedback ticketing system, email & support team contact numbers specific for each deployment

# OUR UNIQUE CUSTOMER SUPPORT STRUCTURE



# Feedback System

Your Query is Our Priority

Issue based (bug, enhancement, data rectification & others) ticket resolution system to address urgent queries on priority



#### **AI Powered** Module Specific Chatbot Personalized Just for You!

Specific phone key number allocation to every client 1

- 2 Local & international support
- 3 Integrated online & offline support

#### Meet Dr. Selena. **Principal Investigator**

?

She works at a large-scale pharmaceutical company. She has been working on a new clinical trial for a cancer treatment drug, which requires careful management of patient randomization and drug distribution. With hundreds of



patients to track and various drugs to distribute, Selena realized that she needed a more efficient way to manage the process.

Exhaustive FAQ sheets per

product module

Selena was introduced to the Octalsoft IWRS by a colleague, and it turned out to be the perfect solution for her needs. With the IWRS, Selena was able to easily manage the entire process of patient randomization and drug distribution in one central location. The system also provided real-time data on patient enrollment, drug inventory, and treatment compliance, which allowed Selena to make informed decisions and adjustments to the trial as needed.

The IWRS also helped Selena save time and resources. Before using the system, Selena and her team had to manually manage patient randomization and drug distribution, which was time-consuming and prone to errors. With the IWRS, these processes were automated, allowing Selena and her team to focus on other aspects of the trial, such as patient recruitment and data analysis.

Thanks to the IWRS, Selena was able to achieve her goal of managing the clinical trial more efficiently, with greater accuracy and transparency.

### WHY CONSIDER OUR eCLINICAL SOLUTION?



#### Single Unified Mobile-Enabled Platform

End-to-end physically & logically secured cloud solution that effortlessly manages data collection, IP supplies, randomization, study compliance, and documents





Patient Centric

Integrated ePRO & eConsent enables you to conduct patient-centric virtual trials to access diverse population all over the world for smarter and faster trials



Audit Ready Database

End-to-end physically & logically secured cloud solution that effortlessly manages data collection, IP supplies, randomization, study compliance, and documents



Global Mega Trials in Multiple Languages

Experience conducting global trials for all types of studies and trial phases with US FDA Submission | EMA Submission | DCGI Submission



Shortest build times with over 90% of studies deployed within 2-4 weeks. Easy-to-use trial builder allows quicker configuration of eCRFs



Configurable, Customizable, Cost Effective & Easy to Use

The platform can easily integrate with your existing system and can also get designed as per your requirements for increased flexibility, mobility and scalability



Unparalleled Global Customer Support

End-to-end physically & logically secured cloud solution that effortlessly manages data collection, IP supplies, randomization, study compliance, and documents

**Our Values** 

For 15+ years, we've mastered the art of Simplifying Technology so that greater discoveries can happen sooner. As a leading eClinical Solution provider, we've always invested in the right expertise, experts, and technology that keeps us, and you at the forefront of science and medicine. As a trusted tech partner for some of the world's leading clinical trial initiatives, our values of innovation, reliability, and client satisfaction guide everything we do.



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