**KEY FEATURES**

- Full-featured CTMS providing trial planning, start-up, monitoring, financial administration, supplies tracking and investigator management functionalities
- Modular, flexible packages designed for both on-premise and hosted environments
- Global deployment including international language and currency support
- Full integrated site visit data from MySites to Data-Driven Monitoring allowing for optimal monitoring oversight
- Advanced site selection capabilities including investigator profiling and search functionality based on historical trial performance and customer-defined metrics
- Application Program Interfaces (APIs) to enable automated data import from any clinical system

**KEY BENEFITS**

- Available as both a hosted and on-premise solution based upon the unique needs of your organization
- Provides unparalleled flexibility and scalability to support companies of all size and scope
- Fast solution deployment and process expertise enables accelerated time to value
- Simplifies trial management through a single, centralized system
- Automated integration with ClinPhone® RTSM, DataLabs® EDC, and Data-Driven Monitoring applications eliminates the need for re-keying data and discrepancy resolution
- Highly configurable and adaptable to sponsor workflow and terminology
- Active user community directly influences product direction and promotes knowledge sharing amongst IMPACT® CTMS users

**Part of the Perceptive MyTrials® framework, enabling integration with clinical trial software applications to help users plan, design and conduct clinical trial programs in a single place**

**Value of a Single Centralized System**

Clinical trials constitute the single most expensive component of the drug development process. In particular, clinical trial operations represent one of the most resource-intensive areas within a biopharmaceutical company and often involve tedious, manual processes for collecting, aggregating and rationalizing information from a wide variety of data sources. A CTMS yields immediate efficiencies and cost savings for clinical operations.

By providing a single, centralized system to orchestrate operational and administrative activities, a CTMS allows biopharmaceutical companies and
CROs to intelligently manage the complexities of clinical trials. Instead of relying on disparate “islands” of information, often in direct conflict with each other, CTMS users benefit from sharing consistent trial data across the entire enterprise. This allows the sponsor or CRO to streamline processes, appropriately respond to current statuses and proactively manage their trials. IMPACT CTMS is an enterprise-wide solution used by many top global pharmaceutical companies and CROs to plan, administer and track every aspect of clinical trials. It is available as a hosted solution, enabling customers to enjoy rapid and flexible deployment of our leading CTMS solution without high upfront costs.

Customers can also choose to implement IMPACT CTMS as an on-premise solution and install it within their operating environment. With one of the largest user bases and over 20 years of proven success, you are in the hands of true CTMS experts when you join the IMPACT solution community.

Making It Easier for You

IMPACT CTMS has been developed with flexibility and configurability in mind. The exceptional breadth of functionality has been designed to adapt to sponsor-specific workflows and terminologies. With IMPACT CTMS, clients do not have to modify their working practices and compromise with inflexible or rigid functionality. The system can manage trials of any complexity and support organizations of any size and global reach with complete flexibility to configure different study types for the different needs of each customer. IMPACT CTMS has a large, global user organization—the IMPACT User Group (IUG). The IUG is an independent organization run by users for the benefit of the user community. It is a forum for exchanging knowledge on IMPACT CTMS usage and a direct channel for feeding back product needs and direction to the Perceptive MyTrials® eClinical Suite. The IUG annually holds two formal, in-person meetings—one in Europe and one in North America. In addition, there are special interest groups that meet throughout the year to exchange information on specific subject areas and provide targeted feedback. For us, customers are not just nameless faces; they are part of the close-knit community of biopharmaceutical professionals in a meaningful, long-term partnership.

The IMPACT CTMS solution is a key part of our leading integrated eClinical Suite. The automated integration with the ClinPhone RTSM and DataLabs EDC and Data-Driven Monitoring applications allows users to seamlessly share trial data. Enabled through our unique integration platform, IMPACT CTMS users enjoy accurate and timely data without having to duplicate activities. The integration of IMPACT CTMS with key applications enables your entire organization to benefit from the true power of CTMS. The IMPACT CTMS solution is an integral part of Perceptive MyTrials® through which we are able to converge our integrated suite of clinical trial software applications.