Conducting clinical trials is a complex, challenging series of tasks that involves generating and consolidating data, managing and organizing people, and making informed decisions. PAREXEL’s IMPACT® Express offers a quick-to-deploy, easy-to-learn, cost effective clinical trials management solution.

IMPACT® Express provides complete trial oversight, tracks sites and progress, and delivers clinical cost tracking functionality in a secure hosted environment. Built with ultimate speed and ease of use in mind, IMPACT® Express can be deployed in four weeks and includes an extensive training program supported by the Perceptive Institute.

IMPACT® Express draws upon PAREXEL Informatics’ 20+ years of Clinical Trials Management Systems innovation and the experiences of more than 26,000 users in more than 70 countries. The world’s leading biopharmaceutical, medical device and research organizations have used PAREXEL CTMS solutions to manage more than 25,000 trials involving almost one million subjects and nearly 400,000 sites. Today, 100 percent of the Top 15 pharmaceutical companies use PAREXEL technologies.
With IMPACT® Express you can start managing your clinical trial in as little as four weeks. IMPACT® Express’ process-driven approach is focused on affordability and flexibility. Its simple, out-of-the-box deployment means no need for technical infrastructure or resources. And since it includes comprehensive training, your users will be running the system confidently from trial start-up to trial completion.

**KEY FEATURES:**
- Process-driven approach focused on affordability and flexibility
- Solution deployment in four weeks
- Complete trial tracking and oversight from a single, easy-to-implement, hosted solution
- Investigator and management selection tools to accelerate study start-up
- Site management and monitoring for both office and field-based monitors
- Real-time study recruitment reporting
- Intelligent assessment of workload to help site monitors plan site visits
- Flexible online end user training

**IMPACT® EXPRESS DELIVERABLES INCLUDE:**
- Hosted, standard set-up supported by a validation certification
- Centralized clinical trial management database
- High-level standard clinical processes
- Customer-specific reference data
- Performance analytics and reporting
- Administrator access to IMPACT® Express
- Internal and external data sharing
- Training material covering how to perform all activities

With IMPACT Express you can start managing your clinical trial in as little as four weeks.