

Your Journey to Adaptive Site Monitoring

CluePoints' Site Profile & Oversight Tool (SPOT) enables adaptive site monitoring so teams can swiftly pinpoint anomalies and translate insights into actionable strategies. Sponsors and CROs can evaluate site performance more efficiently, adjusting site visits to better account for risk and resource workload.

SPOT integrates seamlessly into any RBQM environment. You can either deploy it as a standalone solution or enhance CluePoints' Central Monitoring Platform with SPOT's advanced features.

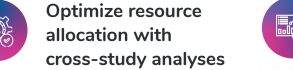
See How SPOT Can Benefit You



Decrease site visits to increase cost savings and efficiency



Better prepare for site visits and integrate multiple systems





Guide critical thinking for site monitoring visits

Make informed decisions regarding future site selection

CluePoints' Central **Monitoring Platform** (CMP) is a customizable Risk-Based Quality Management (RBQM) solution that enables you to detect potential trial risks using comprehensive clinical and operational data. It offers powerful visualization, management, and documentation tools for all resulting actions, powered by a sophisticated

statistical engine.

Enhance Your Technology-Driven Clinical Trial Strategy

SPOT's specialized focus and integration capabilities meet both the fundamental and complex needs of clinical research teams distinguishing it from generic tools in the market. SPOT transforms site monitoring, equipping teams with adaptive, Al-powered tools designed to streamline site visit scheduling and decrease manual data processing.

- O Data-Driven Site Scoring
- User-Triggered Actions
- Automated Actions
- Risk Mitigation
- Data Visualization

- Accessibility Controls
- Study-Level Configurations

Pick Your Path

Yes, I'm a Current CMP User.

- 1. Explore how SPOT integrates with CluePoints' RBQM technology and review features that enhance existing functionality
- 2. Request a demo to see SPOT in action alongside RBQM and understand the combined workflow
- 3. Assess the benefits of integrating SPOT with our platform and compare against other tools on the market*
- 4. Finalize integration, onboarding, and training plans

No, I'm Not a Current CMP User.

- 1. Explore how SPOT integrates with your current infrastructure and review features that enhance existing functionality
- 2. Request a demo to see SPOT in action in a non-CluePoints' environment
- 3. Evaluate how SPOT's standalone capabilities fit your organization, compare against other tools on the market*, and consider the Central Monitoring Platform in addition
- 4. Finalize integration, onboarding, and training plans

*The Comparison

SPOT is a comprehensive solution tailored specifically for site monitoring in clinical trials. Unlike generic data visualization tools that can be adapted for various industries, SPOT is purpose-built to support the unique requirements of clinical research. More specifically, CluePoints created SPOT to address the challenges of fixed monitoring schedules, a lack of visualization to enable actionable insights, and inefficient resource allocation.

30% of clinical trial expenses are linked to site monitoring.

Another major differentiator is SPOT's flexible configuration. Consolidate multiple data inputs and integrate document decision-making capabilities directly into third-party systems, such as Clinical Trial Management Systems (CTMS), where you can comprehensively visualize operational data and make efficient, informed decisions.

Connect with a CluePoints technology expert to explore how SPOT can be tailored to fit your organization's unique needs, complex studies, and existing workflows.



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