The Site Monitor Reborn as Consultant, Trainer and Manager
By Michael Rosenberg and Norman M. Goldfarb

Historically, the primary role of the clinical research site monitor has been to verify that study data and regulatory documents are correct at the site. Much of the site monitor’s activities are thus tedious routine – essentially checking boxes according to standard operating procedures. For most site monitors, the rest of the job is a craft. As with other crafts such as carpentry and blacksmithing, the site monitor learns much of his/her trade through apprenticeship and on-the-job training. As a result, each site monitor has his/her own way of doing things. When a site monitor leaves a study, his/her replacement often has different ways of doing things, creating a period of confusion and inefficiency while the site adjusts to the new rules. This non-interchangeability of personnel is typical of crafts.

Clinical Research data collection and validation is evolving through the three stages in Table 1:

<table>
<thead>
<tr>
<th>Activity</th>
<th>Stage 1</th>
<th>Stage 2</th>
<th>Stage 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Source document data entry</td>
<td>Paper</td>
<td>Paper</td>
<td>Computer</td>
</tr>
<tr>
<td>Case report form data entry</td>
<td>Paper</td>
<td>Computer</td>
<td>n.a.</td>
</tr>
<tr>
<td>CRF data validation</td>
<td>Manual</td>
<td>Computer</td>
<td>n.a.</td>
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In stage 1, the entire process is manual and based on paper. The data does not enter the computer until it has been verified. In stage 2, case report forms (CRFs) and their verification are automated (eCRF). In stage 3, source documents and verification are automated (eSource), and there is no need for the intermediate CRF steps.

Stage 2 is well-underway and we are seeing the beginnings of Stage 3. A variety of products are now available to capture source data without a paper intermediary.¹,²

In Stage 3, the primary task of site monitors – CRF and source document verification – largely disappears. Most of any remaining manual review processes can be performed in the study sponsor’s offices.

The role of the site monitor will evolve through three additional stages, as described below.

Evolution of the Site Monitor’s Role

Most of the tedious work may disappear, but there are still important functions for site monitors to perform at the site. To start with, the sponsor has a regulatory obligation and business imperative to ensure that the site is conducting the trial properly. The site monitor will continue to review the regulatory binder and other essential documents, but perhaps Stage 4 will see many of these documents standardize and move online with electronic signatures.³

With the time freed up from data verification, the site monitor can look at the bigger picture: What is going well and what isn’t? What are the root causes of any problems? Are they random or systemic? Are they short- or long-term? Provided the site monitor knows
how to recruit subjects, obtain informed consent, conduct study activities, enter data, complete regulatory forms, etc., he/she can then provide advice and train site personnel.

For example, sponsors typically train investigators and study coordinators at investigator meetings and then repeat the material at site initiation visits. Sponsor personnel talk, site personnel listen, and there is some Q&A interchange. If the site monitor knows how to conduct a study visit, the site initiation meeting could instead focus on practicing (simulating) study visits, so the initial few subjects do not suffer from the site’s lack of hands-on experience.

These diagnostic, training and consulting activities occur now to a limited extent. Today, they are performed as a craft, not as a highly-organized industrial process. However, in Stage 5, they can be systematized with proven diagnostic tools, knowledgebases and training materials, including online eLearning modules. Operational modeling and metrics will enable site monitors to identify problem areas and measure progress in a systematic manner.

Two important concepts become important in Stage 5:

- **Risk-adjusted monitoring.** With the availability of operational modeling, site monitors will know which operational data potentially reflect important problems, and which are random noise that will have minimal impact on the study’s outcome. Site monitors can then focus their energy where it matters the most.

- **Adaptive monitoring.** Site monitors will schedule less time at high-performing sites and more at sites that can benefit from their help. As site performance changes over time, e.g., a nurse joins a study team or a competitive study completes enrollment, site monitors can shift their attention to the current problems and opportunities. The data that drives adaptive monitoring can also be used to increase resources to productive sites and shut down unproductive sites.

As routine training and consulting become more standardized and automated, the role of the site monitor evolves further, in Stage 6, into site management. There are, of course, site managers today that work for sites. This new type of site manager will resemble today’s outsourcing managers – the people who manage a sponsor’s relationships with, for example, contract research organizations (CROs).

**Implications for Careers**

Only a small minority of today’s site monitors have experience as study coordinators. Because of the different content of the jobs, it can be difficult for a study coordinator to make a successful transition to site monitoring (assuming the job interests them). As the role of the site monitor evolves to consultant, trainer and manager, study-coordinator experience will become important and the skills will fit better. It is much easier to diagnose problems and provide useful advice if you have personal hands-on experience. This evolution will create problems for research sites that lose experienced personnel. On the other hand, study coordinators will be more motivated to develop their expertise so they can move into a rewarding new career path. Study coordinator losses will create more need for training and consulting at the sites. However, the drain on study sites will be limited because site monitor job satisfaction will increase, turn-over will decrease, and the number of site monitors will diminish.

**Implications for Efficiency and Cost**

As the industry moves from Stage 1 to Stage 6, clinical trial efficiency increases and costs decline:
• Technology and lower-cost labor replace expensive site monitor labor.
• Site monitors decline in numbers and their time is employed on high-value activities at high-impact sites.
• Errors are corrected at inception, minimizing rework.
• More accurate data means that study power increases, reducing sample sizes and statistical gymnastics.
• Research sites become more competent and therefore more efficient.
• Lessons from one site, e.g., with respect to subject recruiting, can be quickly propagated to other sites.
• Because training and consulting will create closer personal bonds and sponsors will want to capitalize on the investment they make training and consulting with sites, sponsors will form long-term relationships with their good sites.

Implications for Data Quality

Automating data collection and verification not only reduces costs. It also improves quality:
• If a study coordinator makes an error on a paper CRF, it may be months before the error is identified. By that time, nobody may remember the circumstances and the same error may have been made numerous times. By catching the error immediately with eCRF, or better yet, eSource, the circumstances are fresh and corrective measures can prevent recurrence.
• Eliminating transcription steps eliminates opportunities for error. Eliminating errors eliminates error correction, which introduces its own errors.
• With the noise of all these errors out of the system, study personnel and site monitors can identify systemic quality trends more accurately and keep the site on course.
• Higher-quality, timely data facilitate adaptive trials, completely eliminating errors in data that is not collected at all.

Conclusion

This discussion is not just theoretical. Studies that have applied the methods described above have reduced costs of site monitoring by as much as 80% and total study costs by as much as 32%.4 By focusing site monitors on higher-value tasks, they make more significant contributions to the success of studies and become more valuable to their employers. By working with research sites as consultants, trainers and managers, they improve site performance and strengthen the relationship between sponsor and site. It is time to move away from utilizing site monitors like Medieval craftsmen and leveraging their talents in a manner suitable for the 21st century.

References


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