Clinical Researcher
May 2019
HOME STUDY TEST
Mapping the Way to Site Success

Earn 2.0 Continuing Education Credits
Two articles from the May 2019 issue of Clinical Researcher have been selected as the basis for a Home Study test that contains 20 questions. For your convenience, the selected articles and test questions are combined and posted in the form of this printable PDF at https://www.acrpn.org/professional-development/training/home-study/, where the test may be purchased. The test will be active until May 31, 2020. This activity is anticipated to take two hours. Answers must be submitted using the electronic answer form online (members $30; non-members $50). Those who answer 80% or more of the questions correctly will receive an electronic statement of credit by e-mail within 24 hours. Those who do not pass can retake the test for no additional fee.

The Clinical Researcher archive is at https://www.acrpn.org/resources/clinical-researcher/.

CONTINUING EDUCATION INFORMATION
The Association of Clinical Research Professionals (ACRP) is an approved provider of clinical research continuing education credits.

Contact Hours
The Association of Clinical Research Professionals (ACRP) provides 2.0 contact hours for the completion of this educational activity. These contact hours can be used to meet the maintenance requirements for certification programs of the Academy of Clinical Research Professionals. [ACRP-2019-HMS-005]

ACRP DISCLOSURE STATEMENT
The Association of Clinical Research Professionals (ACRP) requires everyone who is in a position to control the planning of content of an education activity to disclose all relevant financial relationships with any commercial interest. Financial relationships in any amount, occurring within the past 12 months of the activity, including financial relationships of a spouse or life partner, that could create a conflict of interest are requested for disclosure. The intent of this policy is not to prevent individuals with relevant financial relationships from participating; it is intended that such relationships be identified openly so that the audience may form their own judgments about the presentation and the presence of commercial bias with full disclosure of the facts. It remains for the audience to determine whether an individual’s outside interests may reflect a possible bias in either the exposition or the conclusions presented.

ACRP EDITORIAL ADVISORS
Suheila Abdul-Karrim, CCRA, CCRT, FACRP
Victor Chen, MSc
Fraser Gibson, CCRC, CCRA, CCRP
Stefanie La Manna, PhD, MPH, ARNP, FNP-C
Christina Nance, PhD, CPI
Paula Smailes, DNP, RN, MSN, CCRP, CCRC
Jerry Stein, PhD, ACRP-CP
Shirley Trainor-Thomas, MHSA
Heather Wright, CCRC
Nothing to Disclose

ACRP STAFF/VOLUNTEERS
James Michael Causey (Editor-in-Chief)
Gary W. Cramer (Managing Editor)
Jan Kiszko, MD
Deepti Patki, MS, CCRC
Barbara van der Schalie
Nothing to Disclose
Imagine seeing a product you want to buy, but when you go to the website, it asks you to print an order form and mail or fax it in. Would you trust that provider or system with your credit card information? Probably not. And yet, that’s what we are asking patients to do with something far more important— their health—when it comes to clinical trials. People are used to technology, apps, and accessibility in their daily lives, so it is no surprise that a 2016 survey showed 31% of 137 responding patients reported themselves as more likely to participate in a clinical trial if it has a mobile app. [1]
In fact, Deloitte reports there are more than 260,000 health apps worldwide and the members of 70% of patient groups are using at least one app to manage their condition. By utilizing digital solutions, pharma companies can tap into these progressively health-aware consumers. However, those working in the industry, specifically within clinical trials, often feel that the patients in their studies receive a mixed experience of technology, and many simply feel overwhelmed by too many systems.

So why isn’t the pharmaceutical industry moving more quickly to adopt solutions that patients—and sites and study teams—want? Barriers to digitally focused patient centricity include uncertainty over regulators’ expectations and requirements, and data safety and privacy concerns with the rise of medical apps and other digital technologies. Such trust issues can result in patients being unwilling to engage with pharma, while low levels of health and digital literacy also affect the ability to engage with patients effectively. Attracting the right talent to support a patient-centric ecosystem can also be a stumbling block, with a traditional product-based culture causing barriers to an agile, patient-centric approach.

Increased willingness to take a patient-centric approach to technology adoption to improve data capture, patient understanding, and patient engagement is needed to drive real return on investment for retention, compliance, protocol adherence, and overall study timelines. The first important step is improving adoption, but equally important is seamless integration and a single, unified user experience for the patient and sites. It is not enough to give them technology—it must blend into their daily lives so that it doesn’t get in their way. In other words, we seek easy-to-use solutions built with patients (and how they will actually use the tech) in mind. This article will therefore outline how electronic clinical outcome assessment (eCOA), electronic informed consent (eConsent), and patient engagement apps can work together in one solution to combine to transform the patient experience in clinical trials.

As an aside for readers who are concerned that their patients may lack the technologies necessary for the purposes described in this article, eCOA and eConsent are commonly offered using provisioned devices, so that all patients can utilize them, without regard to their personal mobile phone usage. Patient engagement is typically offered to patients on their personal devices, with either mobile app (for smartphone users) or SMS (text messages) for non-smartphone users.
SMS works on any mobile phone, so this ensures that patient engagement can be provided to virtually any patient.

Creating a Seamless Patient Experience

In clinical studies, it is important to employ approaches that enable the optimal assessment of the study concepts of interest. Where this involves use of a technology solution, the aim is to simplify processes, make participation easier, improve quality, facilitate decision making, and collect reliable, honest data.

Patients and their families are understandably impacted by technology as they are going through a trial. In a typical study, for example, one patient may experience four devices, such as a wearable, a reminder app, eConsent, and an e-Diary. It is not uncommon for a trial participant to also use a payment and reimbursement website, all accessed via different systems and devices. As new technologies emerge, they appear via specialist vendors, so for each software and device, patients have a learning curve, different access points, and compliance requirements. As a result, the patient is faced with a dizzying array of disparate communications, devices, and systems, and if the process is too complex, the patient is more likely to quit the study and deviate off protocol. Seamless integration and a single, unified user experience are again the key to making participation as easy as possible without overburdening the patient. It might seem obvious, but they only need the tools that they need.

Sites today are leaning on technology help desks more than ever in their struggles to provide patients all the training they require. Fortunately, by combining these technologies into a single project delivery team, sites receive streamlined and coherent training, which makes them more effective educators to patients.

However, it is not just the patient who needs to be considered, as technology has an impact from multiple perspectives. Improving the clinical trial experience for all involved is a priority. For study teams, enabling them to see real-time study progress and improve efficiencies is the top priority for managing the complexities of a trial and not adding to it. Transparent, real-time insight is the key here. Meanwhile for management, any solution should improve oversight, efficiencies, and successful outcomes for return on investment.
Less Vendor Focused, More Solution Focused

As mentioned, seamless patient experience requires successful collaboration across many different areas of a trial—outreach, recruitment websites, screening, verification, eConsent, registration, data collection, data return, and incentives. How to bring all these elements together needs to be considered carefully, especially in a virtual trial where there may be no physical site. Figure 1 demonstrates what successful collaboration could look like.

**Figure 1: Coordination Among Multiple Systems Sets a Higher Bar for a Seamless Patient Experience**

![Diagram showing coordination among multiple systems]

eConsent, eCOA, and patient engagement solutions are the cornerstones of a dependable, patient-centric technology solution to help improve the speed, reliability, and insight of clinical research. Combined into an integrated solution, it makes for a powerful approach.
Patients need to be informed and retention starts on the right foot through the informed consent process. eConsent sets the patient up to fully understand the study well, and the engagement aspect carries them forward. eConsent incorporates patient-friendly features such as familiar and convenient mobile devices, multimedia video education, pop-up glossary terms, digestible electronic consent form sections, clearer assessments, and enhanced accessibility (i.e., audio narration, large fonts, etc.). There is a growing body of evidence to support the idea that eConsent improves patient comprehension and study retention. One study{3} demonstrated an increase in assessment understanding with eConsent education; meanwhile, CenterWatch{4} highlighted that it is possible to enroll 25% fewer patients to reach the same completion goals as paper-based studies. In addition, consent-related deviations are the second most common audit findings (e.g., people consenting on the wrong version, not re-consenting to new protocol amendments, etc.). eConsent practically eliminates these deviations and improves compliance, thus delivering major benefits in time and cost at the end of a study.

The benefits also extend to study teams, offering advantages such as real-time consent monitoring, bring-your-own-device (BYOD) implementation, SaaS content system creation (with customers configuring their own programs rather than relying on the platform provider for implementation services), eSignature, print-to-sign functionality, and a fully validated solution that provides value anywhere in the world.

*eCOA/ePRO*

With patient engagement–focused eCOA and electronic patient-reported outcome (ePRO, in which patient surveys are collected to help assess responses to the therapy that can’t be measured by lab tests) technologies, the patient has a better experience within the study, which underpins the ultimate goal of capturing the most reliable data and achieving higher retention rates and improved protocol compliance. Whilst a paper-based option could seem easier for patients to use to record information whenever needed, eCOA forces them to respond *correctly*, with guided
responses, and only during the times specified by the protocol, thus preventing “junk data” from being entered at whim or convenience and misleading sponsors.

Access to reminders about personalized medications, appointments, and other study-specific details, benefiting from a single, mobile touchpoint that integrates study commitments into their daily life, are extras that increase engagement among patients. Content libraries presented in digital form, including video, PDF, audio, images, and text, can also facilitate comprehension and help keep the patient fully informed about the study. Bringing these together is an important way to guide patients through their study experiences—giving them the information they need all in one place. This is crucial in a virtual study, which offers less “hand holding” to patients from the site, making the reliance on technology more important.

Patient Engagement

Engaging a patient on a mobile device may drive better patient outcomes, compliance, change behaviors, etc. In the arena of mobile health, it is obvious that technology is vital, but it is important to look at the tools in terms of the value they deliver. Effective patient engagement solutions should eliminate the need for patients to access multiple disparate systems or rely on manual paper practices to keep up with study requirements. For study teams, mobile programs should be configured to match specific study protocols, work across all smartphones and tablets, and be deployable in a BYOD model.

Such a single, streamlined experience has numerous benefits for all stakeholders, and is a significant aid to making evidence-based decisions. Successful patient engagement apps preferred by patients are ideally integrated into a seamless solution providing a much-simplified experience. They integrate visit reminders, documents, site contact info, critical updates, reimbursements, and courier services (e.g., Uber) into a single app. So even when multiple third-party providers are delivering the services, it is invisible to patients because it is all behind the scenes of the user-friendly app. The aim of technology should be to provide a better patient experience. Increasingly complex protocols are challenging enough without overwhelming technologies, plus a single patient experience inspires confidence in overall study conduct.
A streamlined experience is also simpler for sites, requiring less effort but greater impact. Single sign-on and data sharing across systems fosters simplicity and efficiency for site staff, while centralized data entry reduces the possibility of erroneous data and can reduce operational time and cost. Patient engagement apps also offer advantages for sponsors, as trials run smoothly, with faster recruitment and improved retention. Apps can increase patient comprehension of the study and reduce patient burden, resulting in higher completion rates. Efficiencies for sites translate to more active sites, and faster enrollment and less manual data entry provides better data accuracy and integrity.

In a recent vaccine study, patients were offered the option to receive patient engagement messages on their mobile device. Those who chose to receive SMS reminders, including visit reminders and engagement messages, had a 50% lower drop-out rate than those who did not. This also correlated with higher completion rates in the study, demonstrating a significant opportunity to guide the patient. The added benefit of this technology is that it can be used on both a patient’s own device and a provisioned device (BYOD). In a virtual study model, a personal device is more common, making it easy for the patient to download an app.

A First for Patients—Combining ePRO, eConsent, and Engagement

Patient engagement is fundamental not only within a traditional clinical trial setting, but also within a virtual trial, offering a new method of collecting safety and efficacy data from clinical trial participants. Virtual trials (also referred to as direct-to-patient, siteless, or remote trials) are decentralized trials that are less site-centric. The benefits of such trials are that they increase access to include hard-to-reach patient populations (i.e., new geographies) and allow the patient to perform study requirements independently away from the site. Challenges include drug supply, identity verification, consent, and more. Virtual studies are poised to take full advantage of the technologies that are out there, and this is where the value of collaboration and integration really comes into its own.

Case Study—An Integrated Solution

A recently created study sponsored by a top 10 pharmaceutical company looked at developing a deeper understanding and body of evidence around quality of life (QOL) metrics for a specific
indication. There is a lot known about patient experience measured by biomarkers; however, the sponsor believed that the therapy has benefits to the patient that extend beyond optimizing the biomarker measure response. The sponsor wanted to conduct a health-related, QOL virtualized, observational study to attract a large number of patients with this indication and engage with them, with the goal of learning more about their QOL with this indication. Recruitment began in January 2018 with the goal of recruiting up to 1,500 patients. The sponsor was looking for an integrated technology solution to improve patient retention and identity verification. They wanted a solution that increased verification accuracy and reduced patient effort while increasing their comprehension of the study.

The integrated, end-to-end solution for patients was to be a second-generation initiative which presented opportunities and objectives to address challenges experienced in the previous trial, including:

**The Challenges**

- Patients were quick to join the program, but many did not persist (very high drop-off rates)
- A lack of patient engagement and support through the program
- Sponsor needed a reasonable assurance of each patient’s identity/medical condition

**Key Requirements**

- Take consent to the next level by increasing patients’ comprehension of the study
- Do more than collect data
- Improve patient retention
- Be patient friendly
- Follow a similar model to the app store that patients are familiar with to download an app
- Take specific care verifying the patient’s medical condition (via disease-specific screening questionnaire)
- Keep the patient’s interest through to completion
The Solution

The gold standard approach encompasses digital outreach, patient discovery via a landing page and a study overview section, screening, and eConsent through to an engagement app via the patient’s own device, making the process as easy as possible for the patient.

Such a solution offers a much-improved experience to patients and study teams. For patients, it offers a seamless experience, with less work and fewer systems and interfaces, whilst also building trust and increased understanding of the study. Patients are seamlessly guided through the process of learning about a study, understanding their eligibility, consenting, and beginning their engagement through their mobile device. The BYOD diary/patient engagement tool was quickly set up via an automatically initiated activation message sent directly to the patient’s phone. An effortless patient activation process takes the burden off sites and patients. Ease of use and rapid patient setup process instills confidence in users, as the system is intuitive even to first-time users. For study teams, this results in faster recruitment and better retention, higher conversion rates, and better diary completion rates. Full launch documentation was provided, including ethics committee/institutional review board (EC/IRB) submissions, data protection statements, and testing documentation, making deployment virtually effortless for the client team.

Despite targeting a niche patient population, the study reached its enrollment goal well before the target “last person in” date and well under the original outreach budget. Completion rates of the consent form suggested a good balance of user experience and rigor of consent form.

Automated study reminders kept patients on track with the protocol and reduced the burden on clinical research sites who would ordinarily remind study subjects to fulfill their commitments to the program. The solution delivered protocol-specific reminders about study medications, appointments, and PRO diary windows, as well as other need-to-know details. These were sent via text message, push notification, e-mail, or voice.
Completion rates (for both the diaries and the overall study experience) were approximately 10 times higher than the predecessor study, significantly exceeding the expectations of the sponsor.

**Some Thoughts About Privacy**

Our industry is faced with a complex and ever-changing privacy and data security landscape. Each provider of patient-facing technologies approaches these requirements differently, which creates a challenge for all parties involved. Sponsors expend significant effort auditing/qualifying each provider. Sites (and their EC/IRB groups) need to assess and verify that patient privacy rights are upheld by each provider. Patients need to reach a level of comfort with the idea that the multiple technologies with which they are interacting are all safe and secure.

By bringing all of these technologies together, this landscape is greatly simplified for all parties. Additionally, given the scale of providing multiple solutions, a larger organization can more easily stay fully up to speed on changing regulatory frameworks and legal requirements across the many geographic territories in which research is conducted.

**A Future-Proof Gold Standard Solution**

There is huge potential for thinking differently about how existing technologies can be utilized to enable novel measurements for health outcomes and health status in patients, while making the process as easy as possible for patients to participate in a trial. The goal is to move the industry beyond paper, for its obvious limitations and risks, while navigating the current landscape of decentralized, fragmented systems that do not talk to each other. First and foremost, we need to increase adoption, but just as important is the need to utilize technology in a seamlessly integrated way that fits into patient lives to collect life-changing insights without the trial hindering the process or the patient.

The vision of an integrated system with eConsent, eDiaries, reminder apps, and ePRO in one mobile solution that offers a unified solution—for improved patient experience with increased virtual capabilities—is here, but that is not the end. Novel use of technology encompassing wearable devices, smartphone sensors, and motion-based gaming platforms to collect new endpoints is coming along quickly. A future-proof gold standard solution is available through
best-of-breed collaboration and pioneering solution integration intrinsic to this vision of transforming trials and the patient experience.

References


Jeff Lee is Product Lead for Patient Engagement and eConsent with CRF Bracket.
Assessing the capacity of a clinical research coordinator (CRC) is something that many medical research centers struggle with. Not all studies have the same needs or require the same level of support. What is an appropriate balance of minimal risk versus greater than minimal risk or complex studies? How can the complexity of a clinical trial be assessed in a consistent manner across varying disease types? Research leaders struggle with these questions and how to adequately staff their research units to prevent burnout or decreased quality of services.

Most research positions across institutions are extramurally funded, which presents challenges for not only immediate staffing needs, but makes getting to a state of predictive staffing nearly impossible. Over the last several years, Mayo Clinic Florida has gone through an iterative process to develop and refine a tool with the dual purpose of assessing the complexity of a clinical trial and, by extension, using that complexity assessment to determine the trial capacity of a research coordinator.
Background

The tool is based upon the widely available National Cancer Institute (NCI) complexity assessment,\(^1\) which focuses on the key elements of hematology/oncology trials as they relate to number of study arms, complexity of treatment, data collection complexity, and ancillary studies. These elements were categorized as standard, moderate, or highly complex. The NCI focused primarily on community-based programs, and scored on tasks related to direct patient care interactions.\(^2\)

In our local development process, we wanted a tool with a broad scope so that it could be applied to studies of all diseases and even to non-treatment trials. We used the NCI tool as the foundation, but modified it so that any clinical trial at Mayo Clinic Florida would be assessed for complexity in a consistent manner. Through this process, we were able to score biobanks, registries, expanded access situations, drug trials, and device trials in a uniform fashion.

We considered all stages of running a study to determine overall study complexity. The first iteration of the tool was comprised of 21 unique elements each with a possible score of 0 to 3 points. For example, we included items scored on values such as recruitment strategies, principal investigator (PI) experience, screening procedures, number of visits, numbers of departments involved, frequency of monitoring, and activities at follow-up.

One of the elements assessed is the amount of data that must be collected by the local site. Data collection has a significant impact on how much effort will be required for the trial. A score of 3 would indicate that the data are complex and might need to be entered within three days.

Another example is the amount of time needed to complete a baseline/randomization visit. Minimal effort—a score of 1—would relate to any tasks in the baseline/randomization visit that occurred in less than four hours of total time. Baseline/randomization visits requiring greater than eight hours total time received a maximum score of 3.

Guidelines included in the tool provide instructions for what to do in cases such as when baseline and screening visits occur on the same day (only one value is scored 0 to 3 while the other is marked N/A). Figure 1 details how the scores can be applied to the categories.
The highest possible score when adding up all 21 items is 63 points. The elements of the complexity tool relate to the overall study design, team engagement, target accrual, consenting processes, length of study, monitoring elements, billing requirements, and whether there are any associated ancillary studies.

The clinic’s Research Leadership team held several brainstorming meetings to refine the complexity assessment criteria upon which any study would be evaluated for content validity across cancer, neurology/neuroscience, and general non-cancer trials. These meetings occurred until we could ensure the tool could be applied consistently and comprehensively across all clinical trials. We then tested our tool on current protocols in all specialty disease areas, across Phase I through IV and pilot studies, observational registries or biobanks, and device trials by having multiple team members score a study to determine if they resulted in the same or similar
scores based upon their interpretation of the protocol. Once content validity was established, we were ready to develop a standard for using the Complexity Tool as a predictive work load indicator for trial capacity of a research coordinator.

**Developing a Standard**

We began validating the tool in early 2017, with the Research Leadership team on the Florida campus scoring all active studies to date through December 2016 (n=430). Each study received a complexity score. These scores were added to the overall portfolio management tracking tool in use at that time. The portfolio tracker lists studies by PI, Disease Type, and Coordinator, and was used to view the research activity within the disease or investigator’s study portfolio at any given time, including studies assigned to a specific research coordinator.

Using the Complexity Tool, the team was then able to review these portfolios to see what the cumulative complexity scores were for a designated group of staff or disease type (see Figure 2). In this versatile method, clinical trial portfolios could be evaluated for overall complexity through various views such as by PI, study coordinator, disease type, non-cancer vs. cancer trial, or clinical department. The scores varied depending on how the data were viewed, which led to further discussions on greater development and use of the tool.

<table>
<thead>
<tr>
<th>Study</th>
<th>Enrollment Status</th>
<th>Lead Coordinator</th>
<th>Complexity Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trial 1</td>
<td>Open</td>
<td>CRC 1</td>
<td>42</td>
</tr>
<tr>
<td>Trial 2</td>
<td>Open</td>
<td>CRC 1</td>
<td>39</td>
</tr>
<tr>
<td>Trial 3</td>
<td>Closed to Enrollment - No Patients on Tx</td>
<td>CRC 1</td>
<td>4</td>
</tr>
<tr>
<td>Trial 4</td>
<td>Open</td>
<td>CRC 1</td>
<td>42</td>
</tr>
<tr>
<td>Trial 5</td>
<td>Open</td>
<td>CRC 1</td>
<td>48</td>
</tr>
<tr>
<td>Trial 6</td>
<td>Open</td>
<td>CRC 1</td>
<td>54</td>
</tr>
<tr>
<td>Trial 7</td>
<td>Closed to Enrollment - Has Active Patients</td>
<td>CRC 1</td>
<td>44</td>
</tr>
<tr>
<td>Trial 8</td>
<td>Open</td>
<td>CRC 1</td>
<td>50</td>
</tr>
<tr>
<td>Trial</td>
<td>Status</td>
<td>CRC</td>
<td>Score</td>
</tr>
<tr>
<td>---------</td>
<td>-------------------------------</td>
<td>------</td>
<td>-------</td>
</tr>
<tr>
<td>9</td>
<td>Open</td>
<td>CRC 1</td>
<td>30</td>
</tr>
<tr>
<td>10</td>
<td>Open</td>
<td>CRC 2</td>
<td>43</td>
</tr>
<tr>
<td>11</td>
<td>Closed to Enrollment - Has Active Patients</td>
<td>CRC 2</td>
<td>43</td>
</tr>
<tr>
<td>12</td>
<td>Open</td>
<td>CRC 2</td>
<td>48</td>
</tr>
<tr>
<td>13</td>
<td>Open</td>
<td>CRC 2</td>
<td>50</td>
</tr>
<tr>
<td>14</td>
<td>Closed to Enrollment - No Patients on Tx</td>
<td>CRC 2</td>
<td>10</td>
</tr>
<tr>
<td>15</td>
<td>Closed to Enrollment - Has Active Patients</td>
<td>CRC 2</td>
<td>42</td>
</tr>
<tr>
<td>16</td>
<td>Closed to Enrollment - Has Active Patients</td>
<td>CRC 2</td>
<td>48</td>
</tr>
<tr>
<td>17</td>
<td>Closed to Enrollment - Has Active Patients</td>
<td>CRC 2</td>
<td>39</td>
</tr>
</tbody>
</table>

| Total Complexity Score for this Team: | 676 |
| Total Complexity for CRC 1:          | 353 |
| Total Complexity for CRC 2:          | 323 |

While at first glance there did not appear to be general logic or break points in the scoring, once the studies were grouped by disease type, there were obvious natural breaks in the scoring that could be used to delineate what could be considered a high-, moderate-, or low-complexity trial. For example, a trial score of 45 or higher on the tool was considered highly complex; scores of 30 to 44 moderately complex; and scores below 30 low in complexity.

**Addition of a Step-Down Score**

From the initial review and feedback, it was recognized that there are varying stages of work throughout the life cycle of a clinical trial. To accommodate the fluctuating needs or varying effort required, a step-down scoring process was incorporated. The initial score calculated is considered the trial’s overall complexity, and its use assumes the study is active and enrolling patients.
The first step-down score is to accurately assess the study needs once a trial closes to accrual. This step accounts for trials that may still have active patients being followed, but are no longer accruing new patients. To determine this first step-down score, the six elements that relate to enrollment and baseline assessments are subtracted from the overall complexity score.

The second step-down score is for trials that have closed to accrual and have no patients on active treatment. This score applies to trials that are in long-term follow-up only—for example, those assessing survivorship. To determine this score, only the last four elements in the complexity tool are applied for a total maximum score of 12 points (see Figure 3).

<table>
<thead>
<tr>
<th>Figure 3: Step-Down Categories</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Follow-up (Post End of Treatment) Scoring Elements</strong></td>
</tr>
<tr>
<td>Post-treatment visits in 1st year (follow-up visits after drug administration)</td>
</tr>
<tr>
<td>Post-treatment visits after 1st year</td>
</tr>
<tr>
<td>Frequency of required telephone and non-clinic visits</td>
</tr>
<tr>
<td>Complexity of the follow-up visits (Additional testing at follow-up)</td>
</tr>
</tbody>
</table>

Through these three possible steps, a consistent and scalable method for assessing the complexity of any type of trial in a uniform way that accounts for the various stages of the trial was made possible, replicable, and scalable. This step-down score takes into consideration the complexity of each CRC, regardless of whether the study is accruing or only following patients.

**Conclusion**

In summary, the Research Leadership team was able to develop, test, and implement a complexity assessment that allowed for a uniform review of any clinical trial. Through its implementation, we were able to standardize what is considered highly or moderately complex.

After implementation, it was identified that further analysis could be completed to correlate the complexity score of a study with the workload capacity of a CRC. Further development of the
tool was approved, and results of the next iteration of the Complexity Tool will be discussed in a subsequent publication.

References


Alexa Richie, DHSc, is a Research Operations Manager at Mayo Clinic Florida.

Dale Gamble, MHSc, is a Program Manager at Mayo Clinic Florida.

Andrea Tavlarides, PhD, is a Research Supervisor at Mayo Clinic Florida.

Carol Griffin is a Research Operations Administrator at Mayo Clinic Florida.
HOME STUDY

Mapping the Way to Site Success

Three Times the Charm—Transforming Patient Centricity with eConsent, eCOA, and Patient Engagement

LEARNING OBJECTIVE

After reading this article, the participant will be able to describe the importance of unified eConsent, eCOA, and patient engagement technologies to patient-centric research.

DISCLOSURE

Jeff Lee: Employee of CRF Bracket (now Signant Health)

1. A 2016 survey showed what percentage of responding patients reported themselves as more likely to participate in trials featuring mobile apps?
   A. 12%
   B. 19%
   C. 31%
   D. 48%

2. The article mentions barriers to digitally focused patient centricity, including which of the following?
   1. Patients are largely unwilling to pay more to participate in clinical trials using apps.
   2. The pharmaceutical industry has data safety and privacy concerns regarding apps.
   3. Most regulatory authorities have been vocally resistant to the adoption of patient centricity.
   4. It is difficult to attract the right talent to support a patient-centric ecosystem.
   A. 1 and 3 only
   B. 1 and 4 only
   C. 2 and 3 only
   D. 2 and 4 only
3. What “first step” does the author cite as important for taking a patient-centric approach to technology adoption?
   A. Protesting federal restrictions on the wider use of digital technologies in research.
   B. Improving adoption with seamless integration and a unified user experience.
   C. Convincing more study sponsors to allow rapid introduction of apps to clinical trials.
   D. Finding ways to pass more of the technology-related expenses on to the consumer.

4. Which of the following is cited as a set of problems patients have when new technologies appear via specialist vendors?
   A. Legal requirements, accessibility of purchase points, poor packaging
   B. Limited functionality, lack of portability to study sites, disposability requirements
   C. Learning curve, different access points, compliance requirements
   D. Lookalike products, power requirements in different countries, complicated informed consent

5. What solutions does the article list as cornerstones for improving the speed, reliability, and insight of clinical research?
   A. eConsent, eCOA, and patient engagement
   B. eVerify, EDC, and patient recruitment
   C. eDiaries, IVRS, and patient screening
   D. eSignatures, CTMS, and patient accountability

6. The article cites which of the following goals that can be improved by eConsent?
   A. Patient-reported outcomes and study endpoints
   B. Patient comprehension and study retention
   C. Patient compliance and study closeout
   D. Patient advocacy and study protocol development

7. As opposed to a paper-based option for clinical outcome assessments, which of the following does eCOA force patients to do?
   A. Respond immediately
   B. Respond positively
   C. Respond anonymously
   D. Respond correctly

8. According to the article, which of the following patient needs should effective patient engagement solutions eliminate?
   A. Accessing multiple systems to keep up with study requirements
   B. Accounting for time zone changes when moving from site to site
   C. Accepting enrollment into studies without informed consent discussions
   D. Allowing study coordinators to contact them at any time of day or night
9. In a vaccine study cited in the article, how did the drop-out rate among patients who received SMS reminders compare to those who did not?
A. Patients receiving reminders dropped out twice as much as those who did not.
B. Patients receiving reminders had a 25% drop-out rate versus 65% among those who did not.
C. Patients receiving reminders dropped out half as much as those who did not.
D. Patients receiving reminders had a 100% drop-out rate versus 0% among those who did not.

10. How did completion rates for diaries and the overall study experience in a new study combining ePRO, eConsent, and patient engagement compare to those in the sponsor’s previous study?
A. The rates were approximately half as high in the new study.
B. The rates were approximately 10 times higher in the previous study.
C. The rates were approximately twice as high in the previous study.
D. The rates were approximately 10 times higher in the new study.

---

**Trial Complexity and Coordinator Capacity: The Development of a Complexity Tool**

**LEARNING OBJECTIVE**

After reading this article, the participant should be able to describe the notable complicating factors of assessing the capacity of clinical research coordinators to handle trial duties and the elements of a complexity tool for addressing them.

**DISCLOSURE**

Alexa Richie, DHSc; Dale Gamble, MHSc; Andrea Tavlarides, PhD; Carol Griffin: Nothing to disclose

11. Which of the following are mentioned in the article’s introduction as being challenges to the ability of a medical research center when assessing clinical research coordinator (CRC) capacity?
1. Studies have different levels of risk and complexity.
2. Studies have different needs and require support at different levels.
3. Consistent billing of study procedures across different study types.
4. Consistent assessment of trial complexity across different disease types.

A. 1, 2, and 3 only
B. 1, 2, and 4 only
C. 1, 3, and 4 only
D. 2, 3, and 4 only
12. Which of the following is noted as a challenge of extramural funding of research positions?
   A. Budgeting for adequate staff training and development.
   B. Complying with Good Clinical Practice.
   C. Reaching a state of predictive staffing.
   D. Completing study enrollment on schedule.

13. What kind of programs did the National Cancer Institute complexity assessment primarily focus on?
   A. Community-based
   B. Outside the United States
   C. Public-private partnership
   D. Academic medical center

14. What did the Mayo Clinic Florida team consider to determine overall study complexity?
   A. All sources of financial support.
   B. All stages of running a study.
   C. All study staff qualifications.
   D. All regulatory expectations.

15. Which of the following are mentioned as factors related to elements of the complexity tool?
   1. Staff certifications
   2. Team engagement
   3. Billing requirements
   4. Consenting processes
   A. 1, 2, and 3 only
   B. 1, 2, and 4 only
   C. 1, 3, and 4 only
   D. 2, 3, and 4 only

16. What was the next step after content validity of the tool was established?
   A. Developing a standard to use it to predict achievability of enrollment goals.
   B. Developing a standard to use it to predict principal investigator (PI) involvement.
   C. Developing a standard to use it to predict workload for a CRC’s trial capacity.
   D. Developing a standard to use it to predict patients’ placebo responses.

17. The tool allows clinical trial portfolios to be evaluated for complexity through which of the following views?
   A. PI, study coordinator, disease type, non-cancer vs. cancer trial, clinical department
   B. PI, monitor, disease severity, non-blinded vs. blinded trial, regulatory oversight
   C. PI, project manager, rare disease, multi-arm vs. single-arm trial, funding source
   D. PI, institutional review board, pre-marketing vs. post-marketing trial, duration
18. Which of the following is true about natural breaks in the complexity tool’s scoring?
   A. They can be used to identify when trial complexity will lead to treatment futility.
   B. They can be used to identify trial complexity as exceeding or not exceeding authorized levels.
   C. They can be used to identify trial complexity as low, moderate, or high.
   D. They can be used to identify when trial complexity will lead to high enrollment.

19. How did the Mayo Florida Clinic team accommodate the fluctuating needs/varying effort required during a trial’s life cycle within the complexity tool?
   A. By incorporating a rapid staff training regimen.
   B. By incorporating a step-down scoring process.
   C. By incorporating a standard operating procedures policy.
   D. By incorporating a “one and done” trial restriction on PIs.

20. What feature of the tool is used when trials have closed to accrual and have no patients on active treatment?
   A. A second step-down score for long-term trials.
   B. A green light to pursue publication of trial results.
   C. A non-disclosure agreement binding all study staff.
   D. A binding contract for verification of all study data.