Research and Applications

Consent Builder: an innovative tool for creating research informed consent documents

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ABSTRACT

Objective: To describe process innovations related to research informed consent documents, and development and formative evaluation of Consent Builder, a platform for generating consent documents for multicenter studies.

Materials and Methods: Analysis of Institutional Review Board workflows and documents, followed by process redesign, document redesign, and software development. Locally developed software leverages REDCap and LaTeX. A small-scale usability study was conducted.

Results: Process innovations were combining document types, and conceptualizing 2-part informed consent documents: part 1 standardizing the study description and part 2 with local site verbiage. Consent Builder was implemented in the Trial Innovation Network. User survey scores were acceptable; but areas for improvement were noted. LaTeX coding was the biggest challenge for users.

Discussion: The process changes were generally well accepted. The software implementation uncovered unaccounted for assumptions, and variability in IRB review workflow across centers. Technical modifications may be needed before widespread implementation.

Conclusion: We demonstrated proof-of-concept of an approach to generate research consent documents that are consistent across sites in study description, but which allow for customization of local site verbiage. The Consent Builder tool is an example of an operational innovation, helping meet a need that arose in part due to regulations around use of Single IRB for multicenter trials.

Key words: informed consent, user-centered design, IRB (ethics committees, research), multicenter studies

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LAY SUMMARY
This article focuses on research informed consent documents for multicenter trials. Research sites need to review consent documents, even after approved by the designated single institutional review board (IRB), to make sure that local requirements are met. Coordinating centers that manage multicenter trials need to manage a large number of documents. Combined forms may help reduce the number of documents. It is common to find document inconsistencies across sites for the same study. Splitting the consent documents into 2 parts could reduce inconsistencies. Part 1 describes the study itself, and should be identical for all sites. Part 2 provides information that is unique to the site, such as contact information for the local study team. A software tool, Consent Builder, was developed by the Utah Trial Innovation Center to support informed consent document creation, formatting and management. This article describes the software components and workflow, and reports findings from early tool evaluations. Possible areas for improvement in the tool, and in the culture surrounding informed consent for multicenter trials, are discussed.

INTRODUCTION
When the National Institutes of Health (NIH) issued the Single Institutional Review Board IRB (SIRB) policy in 2016, few investigators understood how informed consent documents in multicenter trials would be reviewed. The policy, updated in 2018, requires NIH-supported collaborative studies to use SIRB review. Collaborative, or multicenter studies, are conducted at multiple sites following the same study protocol.

SIRB use does not obviate the need for local review of the study and study documents. While the goal is to streamline Institutional Review Board (IRB) review, sites still need to comply with state and local policies, ensure that the research is feasible at their site, verify qualifications of local study staff, and conduct other Human Research Protection Program (HRPP) reviews. Uncertainty about how to implement SIRB is common and sometimes leads to duplicative full IRB review at study sites, with local edits subsequently introduced into the informed consent form (ICF) inconsistencies between sites.

The informed consent process is intended to help potential participants understand a study, including risks and benefits of the research and their rights as a participant. Researchers usually use a prewritten ICF, and participants sign and date the document to confirm that they have been provided the information and understand it. Informed consent documents are reviewed and approved by the Institutional Review Board(s) providing oversight for the study. Research nurses tend to rely on the ICFs typically reading the document verbatim to participants. Limitations to ICF documents are well known and although IRBs may provide templates or example text the informed consent documents can still fall short of desired readability leading to less than ideal participant understanding. Evolving regulatory and institutional requirements contribute to documents that are long and complex further contributing to understandability challenges.

Guidelines were created for simplification of ICF language and the Revised Common Rule includes requirements that ICF documents begin with a concise presentation of key information although the best format for presenting this information still need to be studied. Research is beginning to demonstrate the value of augmenting standard ICF documents with multimedia, video, fact sheet, or other methods.

There are situations where it may be important to modify consent documents from standard verbiage and there is evidence that document formatting can facilitate understanding; however, in many cases sites request changes to consent documents that are merely preference and not substantive, or changes that focus on institutional interests (eg, liability management) rather than human research protections. Clear guidance is needed, including tools and templates, on how to comply with the SIRB policy. Platforms such as SMART IRB were designed to assist with regulatory aspects but tools and templates are needed to support creation and management of the informed consent documents.

There were multiple ways institutions deal with ICFs for multicenter research. Managing consent forms at the site is burdensome but generic documents may not meet the needs of participants. As SIRB regulation became active, document creation often shifted from local study teams to central coordinating centers but this created new challenges. The sheer number of documents can be overwhelming to coordinate and manage. Adult participants sign the ICF for their own participation. In pediatric studies parents sign a Parental Permission (PP) form for their child’s participation, and children who are old enough sign an Assent document. A multicenter trial at 20 sites enrolling children and young adults (a fairly typical scenario) would require 20 ICF, 20 PP, and 20 Assent documents. If the study is enrolling both English and Spanish speaking participants, the number of documents doubles. These documents must be kept in sync across sites where there are revisions or amendments.

The second challenge was how to support rigor and reproducibility for the consent process. Each site needs to operationalize the same study, yet has site-specific language as well. Local document creation causes multiple back-and-forth revisions between the coordinating center and participating sites. Standardized verbiage (“boilerplate language”) developed by local IRBs may provide portions of the site-specific language but each study still required customization.

OBJECTIVE
This article describes process innovations for informed consent documents, and the development and formative evaluation of software that implements the changes—a document building tool, Consent Builder. REDCap was already being used in some sites to create consent documents. The output was plain-text and additional formatting options were desired including lists and tables, and decorative elements such as dividing lines, logos, or precisely placed images.

MATERIALS AND METHODS
The Utah Trial Innovation Center (TIC) is part of the Trial Network (TIN), formed in 2016 as an initiative within the national Clinical and Translational Science (CTSA) program. The TIN consists of the CTSA hub sites, 3 TICs, and a Recruitment Innovation Center (RIC). The TIN brings together decades of experience in conducting large-scale, multicenter clinical trials that recruit and engage challenging populations. IRB experts and experienced project
managers at the TICs and RIC compared regulatory requirements, IRB checklists, and extant consent document templates to define the initial requirements and process innovations. IRB document review workflows and checklists guided initial software recommendations.

Process innovations

Combined documents

Combining consent for research with HIPAA Authorization language is common practice at some centers. The adult ICF and Parent Permission (PP) forms use nearly identical verbiage. Two types of assent documents were common—a teen assent with verbiage similar to the ICF and PP forms, and a child assent with simpler language. Combining the ICF, PP, and teen assent into a single document was starting to be accepted at some sites.

Two-part consent documents

A breakthrough occurred with the conceptualization of consent forms as 2-part documents. Part 1 or General Language encompasses the study protocol and should be identical across all sites. This could be provided once by a central authority (the study PI). Part 2 (Local Language) includes local contact information and other site-specific verbiage.

Technical innovation: Consent Builder

The Utah TIC developed a tool that implements the document process innovations. The goal was for project managers to be able to produce a richly formatted document that combined general study language (part 1) with site-specific language (part 2), formatted according to site-specific requirements and SIRB reliance agreements. Formatting preferences such as site logo or participant initials on each page needed to be able to be turned on or off separately for each study.

Figure 1 illustrates how a user interacts with the Consent Builder tools. The tool host site was the University of Utah. The host site

sets up the study in the Consent Builder interface, and initiates REDCap surveys for the study. Multicenter studies designate a Lead Site (typically the site affiliated with the Contact PI of the study proposal). All other sites are considered Participating Sites.

1. General language (the science): The Lead Site Principal Investigator or designee enters study protocol language into the Part 1 survey. The survey prompts for background information, study procedures, risks and benefits, and other content. Part 1 language may be separately reviewed/approved by the SIRB along with the initial study application if desired.

2. Local language: Participating sites are provided a copy of the formatted (and SIRB approved, if applicable) Part 1 document along with a link to the part 2 (Site) Survey. Study staff enter their site-specific language, and designate formatting preferences.

3. Generate full consent document: The host (coordinating center) staff use the Consent Builder tool to combine content from parts 1 and 2 and generate the formatted consent document, which is provided back to sites for review and submission to local IRBs.

Architecture

Consent Builder is a suite of tools developed using a tiered web application/server architecture. The locally developed web application leverages REDCap for content management, LaTeX to encode formatting, and generates ICF in Adobe PDF format. The Consent Builder tool accesses survey content via the REDCap application programming interface (API), and merges this content with a LaTeX template stored in the Consent Builder application. Figure 2 depicts the logical division of Consent Builder components and functionality. The web application and service processor use the following components:

- Windows: Windows Server 2016 Datacenter
- Web Server: Apache Tomcat 8.5

Figure 1. Consent Builder workflow. CB: Consent Builder; ICF: Informed Consent Form.
• Java: AdoptOpenJDK
• TeX/LaTeX System: MiKTeX 2.9 or higher
• Directory Service: Microsoft Active Directory (AD)
• Database: MS SQL Server

The database server uses the following components:

• Windows: Windows Server 2016 Datacenter
• Directory Service: Microsoft Active Directory (AD)
• Database: MS SQL Server 2016

REDCap

Consent document content was collected and managed using the REDCap platform hosted at the University of Utah.24,25 REDCap (Research Electronic Data Capture) is a secure, web-based data capture platform, providing an intuitive interface; audit trails; automated export procedures; and procedures and an API for interoperability with external tools. The 3 TICs and RIC all utilized REDCap so it was a logical choice for initial content management. REDCap questionnaires are managed by the Consent Builder host site (University of Utah). An existing REDCap consent form template was modified for Consent Builder to support a 2-part document. Staff at the Utah TIC compared the part 1 and part 2 surveys with IRB review checklists and verified that required components were represented, and located in the appropriate survey part.

LaTeX

Prior experience suggested LaTeX27 as a document formatting platform that could accommodate all the formatting desired by investigators and IRBs. We selected LaTeX in part because Utah staff had used the platform for many years to format study protocols. LaTeX is an open-source typesetting system designed for producing technical and scientific documents. It is widely used in fields such as physics or mathematics.27 Proponents embrace LaTeX because of the control it offers for complex document layouts, customizability, and portability, and it is readily compatible with many external programs. In Consent Builder, document components (headers) are defined in a LaTeX template, along with the ability to switch on or off optional features based on site survey responses. LaTeX provides this versatility via logic commands. Specific signature blocks can be created, and header verbiage adjusted. Images and logos can be added and their placement and dimensions dynamically controlled.

User evaluation

Formative evaluation during development was via informal feedback. IRB experts at the TICs verified that the documents produced by the tool correspond to regulatory requirements. The tool and documents were reviewed at joint TIC/RIC meetings and individually. Regular meetings were held between the software developers, project managers, and IRB staff. During initial use, sites provided feedback directly to the developer. Quality Assurance included standard software code tests and user review. Project managers entered content from manually created documents into the surveys, generated new consent documents, and compared the original document to the newly generated document. IRB staff and experienced IRB reviewers evaluated the generated documents against IRB review checklists.

Perceptions from initial users about using the Consent Builder were formally evaluated in a survey that was administered online via REDCap. The University of Utah IRB determined the user study to be non-human subjects research (IRB number 00149404). We invited participation from project managers who had used the software to build consent documents for at least one actual study (not just testing). Usability and perceived utility were evaluated using the System Usability Scale (SUS).30 We selected the SUS as a short, validated instrument that is an industry standard30–32 and widely recognized benchmark that has been used to evaluate eConsent platforms.33 Limitations of the SUS are that it is minimalistic, originally designed for static interfaces, and may not capture complexities. We measured perceived ease of use and perceived usefulness using 2 questions from the Technology Acceptance Model (TAM) questionnaire34,35 and added custom questions specific to LaTeX.
We continued to collect informal feedback, problems, and comments throughout the initial implementation.

RESULTS

To date, 9 project managers have used Consent Builder to generate consent forms for 13 studies across 119 sites; generating more than 1238 documents. Document changes were largely acceptable to site IRBs although a few were unwilling to allow certain changes such as including HIPAA authorization language within the ICF document. There were site differences in whether part 1 and part 2 could be combined or needed to be reviewed separately. We received survey responses from 7 of the 9 project managers. Experience with the tool varied from use with 1–15 studies. SUS scores range from 0 to 100; and are interpreted as normalized scores (not a percentage). The overall SUS score was 74.58; a score that is considered acceptable but with room for improvement. Table 1 shows the responses to other questions. LaTeX coding problems were the most frequent cause of errors in generating PDF documents.

DISCUSSION

Lessons learned/next steps: software

Although the Consent Builder tool had acceptable usability metrics, and we expect that the principle of 2-part ICF documents will likely gain acceptance, refinement of the Consent Builder tool is needed before hosting could be done at other sites.

Sites responding to surveys do not need to have REDCap installed, because the surveys are accessed in a web browser. However, the choice of REDCap as a content database constrains hosting to sites that have local installations of REDCap. Other databases could manage content but would require modifications to the API calls. In addition, we had manually compared site templates to develop the content questionnaires. A formal data model and standardized representation of for informed consent document content could manage content but would require modifications to the API to sites that have local installations of REDcap. Other databases ever, the choice of REDCap as a content database constrains hosting could be done at other sites.

Lessons learned/next steps: research culture

User preconceptions were an unexpected challenge, including lack of recognition that survey responses would be copied verbatim into the consent document. For example, some users simply typed a name and phone number, assuming the system would automatically convert this to a full paragraph of local contact information text. More detailed user documentation and a user training video were developed to help manage expectations about how the system works, and exemplar response verbiage was added to the REDcap surveys.

The Consent Builder documents aligned well with University of Utah SIRB workflow. The University of Utah IRB embraced the principle of a 2-part consent document, and will accept PDF documents for review. This is not the case at all sites. Efficiencies can be gained by standardizing documents, but users continue to request changes to documents. Many of those requests are preference-based and not substantive. The culture of every researcher wanting a bespoke consent form needs to change along with the processes we use.

Consent Builder addresses a different problem than electronic document signature (eConsent) platforms. Electronic platforms can support participant understanding, at least for some aspects of a study by integrating avatars, on demand glossary, videos, or other features. We recognize that as eConsent tools mature the needs that drove Consent Builder development might be met through other tools. At present, however, centers that coordinate multicenter trials continue to express interest in a tool like Consent Builder that can aid in managing informed consent documents.

CONCLUSION

Innovations are new products, ideas, methods, or processes. The TICs and RIC focus on operational innovations which are improvements in how studies are designed or conducted. Combined ICF reduce the number of documents that need to be managed. Two-part consent keeps the study description consistent across sites. We demonstrated initial proof-of-concept of a Consent Builder tool, which meets a need to support centralized management of the ICF documents in a multicenter trial.

Tools like Consent Builder, or templates within eConsent systems, primarily standardize document structure and format. There continues to be a need for research regarding the content of those documents. Improving informed consent will require evolution in the culture around informed consent documents. Improved understanding of the regulations may support reduced document complexity. Further research may provide evidence that changes from a culture of customized documents based on personal preference, to customizations that are evidence based.

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AUTHOR CONTRIBUTIONS
All listed authors meet the criteria for authorship. JB and JMD focused on tool conception and process improvements. JO, MR, and CE analyzed IRB workflows, evaluation of document changes, and tool output fit with IRB processes. RE was the initial architect and software developer, and collected data for the evaluations. KAS contributed to software design, conception and design of the user study, IRB approval of the user study. ER, JB, and KAS provided the initial paper content. All authors critically revised content, provided final approval of the version to be published, and agree to be accountable for all aspects of the work.

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CONFLICT OF INTERESTS STATEMENT
None declared.

DATA AVAILABILITY
Data from the user survey may be obtained upon request from the corresponding author. Consent Builder software intellectual property is managed by the University of Utah Partners for Innovation, Ventures, Outreach, and Technology (PIVOT) center.

REFERENCES
1. National Institutes of Health. Final NIH Policy on the Use of a Single Institutional Review Board for Multi-Site Research [NOT-OD-16-094]. 2016. https://grants.nih.gov/grants/guide/notice-files/NOT-OD-16-094.html. Accessed November 24, 2021.
2. Office for Human Research Protections (OHRP). Revised Common Rule. 2018. https://www.hhs.gov/ohrp/regulations-and-policy/regulations/finalized-revisions-common-rule/index.html. Accessed May 24, 2021.
3. Corneli A, Dombeck CB, McKenna K, et al. Stakeholder experiences with the single IRB review process and recommendations for Food and Drug Administration guidance. Ethics Hum Res 2021; 43 (3): 26–36.
4. Burr JS, Johnson AR, Vaseinia V, et al. Implementing a central IRB model in a multicenter research network. Ethics Hum Res 2019; 41 (3): 23–8.
5. Hahn C, Kaufmann P, Bang S, et al. Tool conception and process improvements. J Gen Intern Med 2006; 21 (8): 888–9.
6. Nusbaum L, Douglas B, Estrella-Luna N, et al. Survey of risks and benefits communication strategies by research nurses. Nurs Ethics 2019; 26 (3): 937–50.
7. Sugarman J, Paasche-Orlow M. Confirming comprehension of informed consent as a protection of human subjects. J Gen Intern Med 2006; 21 (8): 888–9.
8. Nusbaum L, Douglas B, Damus K, et al. Communicating risks and benefits in informed consent for research: a qualitative study. Glob Qual Nurs Res 2017; 4: 233339617732017.
9. Paasche-Orlow MK, Taylor HA, Brancati FL. Readability standards for informed-consent forms as compared with actual readability. N Engl J Med 2003; 348 (8): 721–6.
10. Hlubocky FJ, Kass NE, Roter D, et al. Investigator disclosure and advanced cancer patient understanding of informed consent and prognosis in Phase I Clinical Trials. JOP 2018; 14 (6): e357–67.
11. Abdel-Rahman SM. Facilitating informed permission/assent/consent in pediatric clinical trials. Paediatr Drugs 2019; 21 (4): 205–14.
12. Corneli A, Namey E, Mueller MP, et al. Evidence-based strategies for shortening informed consent forms in clinical research. J Empir Res Hum Eth 2012; 17 (1): 14–25.
13. O’Rourke PP. The Final Rule: when the rubber meets the road. Am J Bioeth 2017; 17 (7): 27–33.
14. Bazzano LA, Djeddour D, Brantley PR. A modern history of informed consent and the role of key information. Ochsner J 2021; 21 (1): 81–5.
15. Solomon ED, Mouzeyry J, Wrobleski MP, et al. Understanding the use of optimal formatting and plain language when presenting key information in clinical trials. J Empir Res Hum Eth 2022; 17 (1–2): 177–92.
16. Sugarman J. Examining provisions related to consent in the Revised Common Rule. Am J Bioeth 2017; 17 (7): 22–6.
17. Kass NE, Sugarman J, Medley AM, et al. An intervention to improve cancer patients’ understanding of early-phase clinical trials. IRB 2009; 31 (3): 1–10.
18. Taylor HA, Washington D, Wang NY, et al. Randomized comparison of two interventions to enhance understanding during the informed consent process for research. Clin Trials 2021; 18 (4): 466–76.
19. Rothwell E, Johnson E, Wong B, et al. Comparison of video, app, and standard consent processes on decision-making for biospecimen research: a randomized controlled trial. J Empir Res Hum Eth 2020; 15 (4): 252–60.
20. Dickert N, Kass N, Pasche-Orlow M, et al. Excluding particular information from consent forms. Account Res 2005; 12 (1): 33–45.
21. Suver CM, Hamann JK, Chin EM, et al. Informed consent in two Alzheimer’s disease research centers: Insights from research coordinators. AJOB Empir Bioeth 2020; 11 (2): 114–24.
22. Wilfond BS, Zabrowski J, Johnson LM. The limitations of “boilerplate” language in informed consent: single IRB review of multisite genetic research in military personnel. Am J Bioeth 2019; 19 (4): 81–2.
23. Bernard GR, Harris PA, Pulley JM, et al. A collaborative, academic approach to optimizing the national clinical research infrastructure: the first year of the Trial Innovation Network. J Clin Transl Sci 2018; 2 (4): 187–92.
24. Harris PA, Taylor R, Minor BL, et al.; REDCap Consortium. The REDCap consortium: building an international community of software platform partners. J Biomed Informat 2019; 95: 103208.
25. Harris PA, Taylor R, Thielke R, et al. Research electronic data capture (REDCap)—a metadata-driven methodology and workflow process for providing translational research informatics support. J Biomed Informat 2009; 42 (2): 377–81.
26. Lind CE, Pytovarova E, Appelbaum P, et al. RELIANCE agreements and single IRB review of multisite research: Concerns of IRB members and staff. AJOB Empir Bioeth 2018; 9 (3): 164–72.
27. LaTeX Project. https://www.latex-project.org/. Accessed November 24, 2021.
28. International Standards Organization (ISO). Document Management—How To Do Things With PDF Documents—Part 2: PDF 2.0. ISO 32000-2:2020 2020. https://www.iso.org/standard/75839.html. Accessed November 24, 2021.
29. Matthews D. Craft beautiful equations in Word with LaTeX. J Biomed Inform 2021; 124: 101320.
30. Finkelstein J, Robins D, Liu J. Usability inspection of multipurpose scalable informed consent platform. Stud Health Technol Inform 2019; 262: 198–201.
34. Davis FD. Perceived usefulness, perceived ease of use, and user acceptance of information technology. *MIS Quarterly* 1989; 13 (3): 319–40.

35. Chuttur MY. Overview of the Technology Acceptance Model: Origins, developments and future directions. *Indiana University, USA. Sprouts: Working Papers on Information Systems* 2009; 9 (37). http://sprouts.aisnet.org/9-37. Accessed November 24, 2021.

36. Knauff M, Nejasmic J. An efficiency comparison of document preparation systems used in academic research and development. *PLoS One* 2014; 9 (12): e115069.

37. University of Utah Trial Innovation Center (TIC). Consent Builder Tool. YouTube video 2018. https://www.youtube.com/watch?v=W0K2pwbQAAc. Accessed July 1, 2022.

38. Rothwell E, Wong B, Rose NC, et al. A randomized controlled trial of an electronic informed consent process. *J Empir Res Hum Res Ethics* 2014; 9 (5): 1–7.

39. Lawrence CE, Dunkel L, McEver M, et al. A REDCap-based model for electronic consent (eConsent): Moving toward a more personalized consent. *J Clin Transl Sci* 2020; 4 (4): 345–53.

40. Merriam-Webster Dictionary. Online Version. https://www.merriam-webster.com/dictionary/innovation. Accessed August 30, 2021.

41. Shah MR, Culp MA, Gersing KR, et al. Early vision for the CTSA Program Trial Innovation Network: a perspective from the National Center for Advancing Translational Sciences. *Clin Transl Sci* 2017; 10 (5): 311–3.

42. Hammer M. Deep change. How operational innovation can transform your company. *Harv Bus Rev* 2004; 82 (4): 84–141.

43. England A, Wade K, Smith PB, et al.; Best Pharmaceuticals for Children Act—Pediatric Trials Network Administrative Core Committee. Optimizing operational efficiencies in early phase trials: The pediatric Trials Network experience. *Contemp Clin Trials* 2016; 47: 376–82.