PARTNERING IN PROGRESS

14th Annual AACI CRI Meeting

JULY 12–14, 2022

Loews Chicago O’Hare Hotel
The Association of American Cancer Institutes (AACI) Clinical Research Innovation (CRI) program serves as a network for research leaders to develop and share best practices for the efficient operation of clinical trials offices (CTO) at AACI cancer centers.

The programming of the 14th Annual AACI CRI Meeting, “Partnering in Progress,” aligns with CRI’s strategic goal of stimulating cancer center interactions to maximize resources by creating opportunities for peer-to-peer networking and collaboration.

2022 Strategic Plan & Goals

1. Increase AACI cancer center participation in CRI
2. Collect and disseminate benchmarking data to develop cancer center clinical trial best practices
3. Integrate CRI into AACI programs and initiatives
4. Assist cancer centers in improving community outreach and engagement to increase enrollment in clinical trials
5. Promote diversity, equity, and inclusion for training and career development
6. Develop clinical research education and resources for professional development
7. Increase engagement with industry and other stakeholders to support CRI
8. Develop outcomes to drive change and advance cancer center clinical research programs
9. Create a network for leadership to continue fostering communication and mentoring opportunities

AACI CRI Welcome

Meeting App and Social Media

The agenda, presentations, exhibitor information, and attendees list are available on the meeting app.

To access the app:
- Download “Cvent Events” from your mobile app store.
- Log in with the name and email used during registration.
- You will receive a 6-digit verification code. Enter your code and log in.
- Within the Cvent Events app, search for “14th Annual AACI Clinical Research Innovation Meeting” and download the meeting.

The meeting app can also be accessed on your desktop at: https://cvent.me/NQV48v

Twitter: @AACI_Cancer
Facebook: AACICancer
Hashtag: #CRI2022
Wireless Network: AACI CRI Wi-Fi
Password: 2022cri

AACI CRI 2022 Steering Committee

Chair: Tara L. Lin, MD
The University of Kansas Cancer Center

Leslie P. Byatt, CCRC, ACRP-PM
University of New Mexico Comprehensive Cancer Center

Tiffany Colvin, CCRC
University of Colorado Cancer Center

Thomas J. George, Jr., MD, FACP
University of Florida Health Cancer Center

Janie Hofacker, RN, BSN, MS
Association of American Cancer Institutes

Christopher Loertscher, MA
USC Norris Comprehensive Cancer Center

Patricia M. LoRusso, DO, PhD
Yale Cancer Center; Yale School of Medicine

Eneida Nemecek, MD, MS, MBA
OHSU Knight Cancer Institute

Bhanu Pappu, PhD, MHA
Simmons Comprehensive Cancer Center
UT Southwestern Medical Center

Anne Schnatterly, MBA, BSN, RN, CCRP
WVU Cancer Institute

Susanna Sellmann, MRT, BSc., CCRP
Princess Margaret Cancer Centre
University Health Network

Rachna Shroff, MD, MS
The University of Arizona Cancer Center

Kate Shumate, MPA, CCRP
UCSF Helen Diller Family Comprehensive Cancer Center

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AACI CRI Meeting Objectives

- Offer an industry perspective on diversity related to the workplace and clinical trial recruitment
- Address how cancer centers utilize their institutional support to broaden a culture of diversity, equity, and inclusion (DEI)
- Share strategies that CTOs are using to engage research team collaboration with community outreach and engagement (COE) “champions” and the benefits of working with community liaisons or navigators to increase trial enrollment
- Discuss ways to provide community patients access to research drugs and improve the research treatment experience using community-based navigators, home health care, and telehealth
- Share successful recruitment models for rapid onboarding of staff to address the unprecedented employee shortages resulting from the “Great Resignation”
- Discuss the expanding use of innovative technology to enhance staff efficiency and patient experience in clinical trials
- Review the advantages of using a clinical trial feasibility process to ensure a realistic assessment of trial site capabilities for conducting trials
- Learn the benefits of rightsizing clinical trial portfolios for streamlining CTO operations while addressing staffing gaps to improve patient outcomes
- Review the practical implications of preparing a National Cancer Institute (NCI) Cancer Center Support Grant competing renewal, with a focus on the Clinical Trials Reporting Program, revisions to the Funding Opportunity Announcement, and the funding of investigator-initiated trials
- Understand cancer center catchment areas, the NCI’s new diversity and inclusion reporting requirements, and prospects for continued virtual site visits

Who Attends This Meeting?

- Individuals from AACI member cancer centers, including:
  - CTO administrative directors, medical directors, managers, and supervisors
  - Deputy/associate directors of clinical research administration
  - Cancer center administrators
  - Research regulatory management and staff
  - Clinical research finance directors, managers, and supervisors
  - Biostatisticians and informatics specialists
- Employees of U.S. Department of Health and Human Services agencies and offices, including the NCI and the U.S. Food and Drug Administration
- AACI sustaining members
- AACI Corporate Roundtable members
- Representatives from industry, including drug development companies, clinical research organizations, and consultants
- Information technology companies that support cancer center clinical research management
- Like-minded organizations promoting patient access to clinical trials
Management and Program Management Warhol
In this breakout session, managers will discuss how to effectively motivate and communicate with staff to balance workloads, increase productivity and efficiency, and maintain a positive environment.

Regulatory Management Avedon A
Breakout session participants will share best practices for creating and maintaining the clinical trial regulatory file and electronic systems to reduce paper and increase efficiency. Participants will also discuss working with regulatory bodies, the U.S. Food and Drug Administration, and local and central institutional review boards; continuing renewals; and managing trial closures.

Trial Coordination Avedon B
Participants in this breakout session will describe best practices for coordination of clinical trials to include methods for screening and consenting patients, confirming trial eligibility, coordinating protocol procedures and investigational drug ordering, reporting serious adverse events, and documenting sources in the electronic medical record.

Trial Start-up Louvre 2
Trial activation involves teamwork from people in various roles, working both inside and outside the clinical trials office. Breakout participants will discuss challenges and methods for shrinking the trial activation timeline while working with the cancer center’s protocol and review monitoring system to prioritize trials. Other topics include coordinating workflows for developing trial budgets, negotiating contracts, and developing Medicare coverage analysis.

2:30 PM Networking Break Artist Foyer
Supported by Essex Management

3:00 PM Clarity and Perspective: The NCI Cancer Center Support Grant and New Reporting Guidelines
Cassatt Ballroom
National Cancer Institute (NCI) officials will provide an overview of recent and proposed changes to NCI’s Cancer Center Support Grant (CCSG), including new guidelines for reporting data for pragmatic trials. Panelists will also review the practical implications of preparing a CCSG competing renewal, with a focus on the Clinical Trials Reporting Program, revisions to the Funding Opportunity Announcement—including funding of investigator-initiated trials and catchment area considerations—new diversity and inclusion reporting requirements, and prospects for continued virtual site visits.

Moderator: Kate Shumate, MPA, CCRP
UCSF Helen Diller Family Comprehensive Cancer Center
Krzysztof Ptak, PhD
National Cancer Institute

4:00 PM Poster Session Avedon CD

5:30 PM Welcome Reception Artist Foyer
Supported by Florence

Wednesday, July 13

7:00 AM General Breakfast Guggenheim Ballroom – Museum Wing

7:00 AM CTO Administrative Directors’ Breakfast (invitation only) Prado 1

7:00 AM CTO Medical Directors’ Breakfast (invitation only) Prado 2

8:00 AM Exhibits Open Artist Foyer

8:15 AM Welcome Cassatt Ballroom
Tara L. Lin, MD
The University of Kansas Cancer Center

8:30 AM Keynote Presentation: Optimizing Diversity Efforts Cassatt Ballroom
Bristol Myers Squibb People and Business Group Lead Monique Phillips, PMP, will offer an industry perspective on diversity—both as it relates to workplace culture and to clinical trial recruitment. She will be joined in discussion by ovarian cancer survivor and patient advocate Kimberly Richardson, of Black Cancer Collaborative. Topics will include patient navigation; implicit bias; outreach and education strategies; and diversity, equity, and inclusion programs.

Monique Phillips, PMP
Bristol Myers Squibb

Kimberly Richardson
Black Cancer Collaborative

9:30 AM Meeting Our Patients on Their Terms Cassatt Ballroom
Panelists will discuss ways to improve the treatment experience for patients, including the use of community-based navigators and drug shipments to affiliate sites. Additionally, a patient advocate will share their unique perspective.

Moderator: Bhanu Pappu, PhD, MHA
Simmons Comprehensive Cancer Center, UT Southwestern Medical Center

Sean DeFrates, PharmD
Robert H. Lurie Comprehensive Cancer Center of Northwestern University

Annie Ellis
Memorial Sloan Kettering Cancer Center

Fabian Robles, MSc-HLM
Simmons Comprehensive Cancer Center, UT Southwestern Medical Center
Using Technology to Simplify CTO Administrative Responsibilities  Cassatt Ballroom
Supported by Veeva

Artificial intelligence innovations such as automated trial-patient matching and other technology solutions (e.g., electronic informed consent and electronic regulatory management systems) are enhancing clinical trials office efforts to reduce administrative burden on staff that are sometimes overwhelmed by the shifting workplace demands caused by the ongoing COVID-19 pandemic. This panel discussion will explore the expanding use of technology to enhance both staff efficiency and patient experience in clinical trials.

Moderator: Thomas J. George, Jr., MD, FACP
University of Florida Health Cancer Center

Tricia Bentz, MHA, CCRP
Hollings Cancer Center, Medical University of South Carolina

Jocelyn Brown, CCRP
Stephenson Cancer Center, University of Oklahoma

Ashley Drawz, MSRC, CCRC
Robert H. Lurie Comprehensive Cancer Center of Northwestern University

Ensuring a Good Study: The Clinical Trial Feasibility Process  Cassatt Ballroom

A rigorous clinical trial feasibility process helps to ensure a realistic assessment of your ability to conduct the trial. Panelists in this session will examine steps in the process such as assessing internal and environmental capacity, aligning the clinical trial with study design, and evaluating the timing and other logistics of opening a trial at an affiliate site. Pharmacy considerations like investigational drug availability and dosage will also be discussed along with approaches to interfacing with a cancer center's protocol review and monitoring committee.

Moderator: Leslie P. Byatt, CCRC, ACRP-PM
University of New Mexico Comprehensive Cancer Center

Frances Brogan, MSN, RN, OCN, CCRP
Herbert Irving Comprehensive Cancer Center
Columbia University Irving Medical Center

Donna Katz, MSN, APRN, FNP-BC, OCN
UCLA Jonsson Comprehensive Cancer Center

Brian Miller, PharmD
UPMC Hillman Cancer Center

Trial Activation Outside the Clinical Trials Office (CTO) Warhol
Trial activation requires the teamwork of members both within and outside the CTO. Breakout session participants will discuss their methods for working with their finance and contracts teams and sponsors to activate trials within required timelines and meet trial budget requirements.

Managing Trials From Beginning to End Louvre 3
Session participants will discuss rightsizing your clinical trial portfolio in the current environment to address your cancer patients' needs and maintain CTO efficiency.

Shared Investigator Platform (SIP) Avedon B
Breakout session participants will describe the challenges with SIP adoption and trial activation. Topics include site facility setup, establishing investigator profiles, and staffing the site administrator role. Representatives from Cognizant will be available to answer questions.

Reviewing Investigator-Initiated Trials With Your Data Safety Monitoring Committee Pollock
Participants in this breakout session will discuss how they work with their Data Safety Monitoring Committee to review safety reports and monitor investigator-initiated trials.

Managing Multisite Investigator-Initiated Trials (IIT) Louvre 1
Participants will discuss the management of multisite IITs and best practices for providing investigational drugs and managing the regulatory file.

Managing Your Community Trial Sites Avedon A
Participants will discuss trial site selection, trial monitoring, and ensuring timely and accurate data collection. They will also address financial considerations for sites working with trial site affiliates.

Diversity, Equity, and Inclusion: Working With Your COE Louvre 2
Participants will continue the conversation on methods for engaging the community. Discussion may include working with community navigators and patient advocates; determining how to diversify and retain staff; and enrolling more patients in clinical trials.

Networking and Dessert Break  Artist Foyer
Supported by Advarra
3:15 PM  **2022 CRI Abstract Presentations  Cassatt Ballroom**

Abstracts received from AACI cancer center members focus on oncology research that illuminates clinical research challenges and solutions, accelerating cancer drug development. The CRI steering committee has selected three abstracts for presentation at this year’s meeting. The abstract presentations will be followed by a Q&A session.

**Moderator: Tara L. Lin, MD**  
*The University of Kansas Cancer Center*

**Angela Fritsche, MPA**  
*Mayo Clinic Comprehensive Cancer Center*

**Erin Hastings Monari, PhD, CCRP**  
*University of Florida Health Cancer Center*

**Josefina Sanchez, CCRC**  
*Sylvester Comprehensive Cancer Center*  
*University of Miami Health System*

4:15 PM  **Honorable Mention Poster Session  Cassatt Ballroom**

Abstract authors will share challenges they experienced within their clinical trials office and the innovative solutions implemented to improve cancer center operations.

**Moderator: Tara L. Lin, MD**  
*The University of Kansas Cancer Center*

**Christina Crabtree-Ide, PhD, MPH**  
*Roswell Park Comprehensive Cancer Center*

**Theresa Cummings, RN, DBA, CCRP**  
*UNC Lineberger Comprehensive Cancer Center*  
*University of North Carolina at Chapel Hill*

**Emily Hautamaki, RN, MPH, CCRP**  
*Cedars-Sinai Cancer*

**Ashley McCauley, MA, CCRP**  
*Fred Hutchinson Cancer Center*

**Justin Miller**  
*The Tisch Cancer Institute at Mount Sinai*

**Lillian Neal, MSc, CCRP**  
*Hollings Cancer Center, Medical University of South Carolina*

**Renata Panchal, MS**  
*Memorial Sloan Kettering Cancer Center*

**Beth Scanlan, MAP, CCRP**  
*UAMS Winthrop P. Rockefeller Cancer Institute*

5:15 PM  **Vendor Presentation: Advarra**

**Site-Sponsor Consortium: Connecting Research Sites and Sponsors Through an Integrated Technology Ecosystem**

In this session, Laura Hilty, vice president, strategy at Advarra, will outline the challenges and present solutions for cancer centers overburdened by sponsor-driven technology. Hilty will share metrics from a recent survey conducted by Advarra and Society for Clinical Research Sites and will highlight progress made by a new consortium of sites, sponsors, and clinical research organizations. The consortium is working together to connect existing site technology to sponsor systems seamlessly, addressing the duplicative efforts, longer timelines, and reduced patient focus often associated with sponsor-provided technology. Hilty will showcase progress made by Advarra as a result of the consortium’s efforts and provide actionable insight for attendees to take advantage of this connected ecosystem, empowering site stakeholders to drive efficiencies across the clinical trial lifecycle.

8:00 – 10:00 PM  **Hospitality  Ice Bar, Lobby Level**  
*Supported by WellSky*
9:15 AM Rightsizing the Clinical Trial Portfolio  Cassatt Ballroom

Much as a quarterly review of financial investments helps to generate income and minimize losses, a clear-eyed analysis of a clinical trial portfolio can streamline CTO operations and improve research and patient outcomes. Helping disease teams prioritize staffing, closing trials, increasing trial efficiency numbers, and focusing on the National Cancer Institute’s Cancer Center Support Grant requirements, especially as they relate to investigator-initiated trials, are some of the steps toward rightsizing a clinical trial portfolio that will be examined in this session.

Moderator: Anne Schnatterly, MBA, BSN, RN, CCRP  
WVU Cancer Institute

Michael Pearl, MD  
Stony Brook Cancer Center

Theresa Sberna, MPH  
Vanderbilt-Ingram Cancer Center

10:15 AM Break

10:30 AM Successful Staff Retention and Training  Cassatt Ballroom

Amid the “Great Resignation” and unprecedented staffing shortages, recruiting, training, and retaining employees have become top priorities among cancer centers. Speakers will share successful recruitment strategies and models for rapid onboarding. They will also address rightsizing trial portfolios and maintaining trial accruals despite staffing gaps.

Moderator: Christopher Loertscher, MA  
USC Norris Comprehensive Cancer Center

Chloe Fournier, MBA, CCRP  
Duke Cancer Institute, Duke University Medical Center

Stacey Lewis, RN, BSN, OCN, BBA  
Atrium Health Wake Forest Baptist Comprehensive Cancer Center

Andrea Skafel, MSc, CCRP  
UCSF Helen Diller Family Comprehensive Cancer Center

11:30 AM Closing Remarks  Cassatt Ballroom

Janie Hofacker, RN, BSN, MS  
Association of American Cancer Institutes

11:45 AM Adjourn
AACI CRI Meeting 2022 Abstracts

FIRST PLACE:
5. Successful Methods of Addressing Clinical Research Staff Turnover
N. Nahmias, J. Sanchez, A. Olier-Pino, A. Allred, K. Aviles, L. Corrales
Sylvester Comprehensive Cancer Center, University of Miami Health System

SECOND PLACE:
32. Starting Off on the Right Foot: Elevating the Voice of Community Stakeholders During the IIT Development Process
E. Monari, S. Szurek, A. Ivey, T. George, A. Anderson, E. Shenkman, C. Evans, A. Lawson-Ross
University of Florida Health Cancer Center

THIRD PLACE:
8. Clinical Research Coordinator Workload Estimation and Tracking
M. Repede, D. Beighley, K. Putz, A. Fritsche, G. Nowakowski
Mayo Clinic Comprehensive Cancer Center

Abstracts are organized by category and completion status, then in alphabetical order by cancer center.

Clinical Trial Operations

1. Clinical Trial Office Response to COVID-19 at an Academic Comprehensive Cancer Center
E. Bentlyewski, F. Brogan, R. Shelton, J. Jurcic, A. Lassman
Herbert Irving Comprehensive Cancer Center, Columbia University Irving Medical Center

2. CTO Benchmarking Study
M. Contreraz, T. Lautenschlaeger, K. Lee, K. Miller, B. Hicks, A. Sands, F. Bhimani, M. Gee, T. Hanson, B. Richardson
Indiana University Melvin and Bren Simon Comprehensive Cancer Center

3. Single Institution Experience of Integrating Radiation Oncology Clinical Research Into Comprehensive Cancer Center CTO
Indiana University Melvin and Bren Simon Comprehensive Cancer Center

4. Solutions for Clinical Research Continuity During the COVID-19 Pandemic
N. Nahmias, J. Sanchez, A. Olier-Pino, A. Allred, K. Aviles, L. Corrales
Sylvester Comprehensive Cancer Center, University of Miami Health System

5. Successful Methods of Addressing Clinical Research Staff Turnover
N. Nahmias, J. Sanchez, A. Olier-Pino, A. Allred, K. Aviles, L. Corrales
Sylvester Comprehensive Cancer Center, University of Miami Health System

6. eRegulatory Process and Software Implementation in Times of Crisis
J. Stern, R. Jones, C. Henrichs
Vanderbilt-Ingram Cancer Center

7. Automated Reporting for Clinical Trial Operations
M. Rump, M. Kilbane, K. McCaffrey, B. Matia, J. Lindberg
Cleveland Clinic Cancer Center

8. Clinical Research Coordinator Workload Estimation and Tracking
M. Repede, D. Beighley, K. Putz, A. Fritsche, G. Nowakowski
Mayo Clinic Comprehensive Cancer Center

9. Data Automation to CIBMTR
C. Thomas, R. Panchal, J. Konecny, T. Casali
Memorial Sloan Kettering Cancer Center

10. Memorial Sloan Kettering Cancer Alliance’s Cancer Control and Population Science Subcommittee
S. Yoon
Memorial Sloan Kettering Cancer Center

11. Data Talks: Using Non-Compliance Tracking to Improve Patient Safety
E. Harms, N. Borror, E. Menne
Siteman Cancer Center

12. Digital Transformation of Sylvester Comprehensive Cancer Center Research Lab: A Work in Progress
P. Seo, A. Blivin-Martinez, J. Trent
Sylvester Comprehensive Cancer Center, University of Miami Health System

13. Implementing “Protected Time” to Increase Clinical Research Coordinator Data Entry Efficiency
J. Miller, S. Scheiner, C. Varnadoe-Rothman, S. Lopiano, N. Taylor, P. Garcia
The Tisch Cancer Institute at Mount Sinai

14. Study Start-up Dashboard
D. Mudaranhakam, S. Pepper, A. Tribitt
The University of Kansas Cancer Center

15. Establishing a Pre-Screening Process – Creating the Right Tool
UAMS Winthrop P. Rockefeller Cancer Institute

16. Navigating Oncology Clinical Trials in the Era of COVID-19 – A CRC Perspective
M. Boota, K. Allen, S. Panozzo, M. Kovak, M. Birrer
UAMS Winthrop P. Rockefeller Cancer Institute

17. Microsoft Power Platform: Improving Efficiency, Communication, and Documentation in the Clinical Research Setting
M. Fritz, J. Plassmeyer, M. Horak, T. Cronauer (Horne), S. Perry, J. Griffo, D. Cleary
UPMC Hillman Cancer Center

18. Surveying Staff Satisfaction to Work Toward Improved Employee Retention
J. Binder, D. Ritter, K. Garcea, M. Fritz, M. Horak, D. Cleary
UPMC Hillman Cancer Center

*Honorable Mention
Poster not present at meeting
2022 Abstracts

Finance/CCSG/PRMS

19. Creation of a Consort Diagram to Visualize Participant Enrollment and Allocation at the Memorial Sloan Kettering Data and Safety Monitoring Committee
C. Kolenut, K. Napolitano, X. Lekperic, S. Hanley, K. Tan, E. O'Reilly, S. Slovin
Memorial Sloan Kettering Cancer Center

J. Migliacci, X. Lekperic, B. Seko, K. Kaufman, K. Napolitano, S. Hanley, A. Rodavitch
Memorial Sloan Kettering Cancer Center

21. PRMC Member Workload Survey After Charter Alignment With NCI Requirements
B. Hughes, C. Allen, T. Herzog, C. Vollmer, M. Marcum, N. Kurtzweil
University of Cincinnati Cancer Center

22. Impact of the SRMC Zero Tolerance Policy on DSG Trial Portfolios
J. Walsh, T. Guinn, Jr., T. George, A. Anderson, A. Ivey
University of Florida Health Cancer Center

23.*Taking a Closer Look: Standardizing Disease Focus Groups to Strengthen Trial Portfolios
L. Neal
Hollings Cancer Center, Medical University of South Carolina

24. Automating and Streamlining the 2-Stage Scientific Review Process
Mayo Clinic Comprehensive Cancer Center

25. Clinical Research: Following the Money Phase III
R. Geary, P. Eggleton, M. Kovak, M. Birrer, A. Smith, Z. Feng, N. Pruss
UAMS Winthrop P. Rockefeller Cancer Institute
Information Technology Research Systems, University of Arkansas for Medical Sciences

26. Monitoring Study Enrollment Demographics: PRMS-COE Collaboration at University of Colorado Cancer Center (UCCC)
D. McCollister, D. Pacheco, A. Henningham, E. Borrayo, C. Cost
University of Colorado Cancer Center

Investigator-Initiated Trials

27.*Development of a Multisite Investigator-Initiated Trial Coordinating Center at Cedars-Sinai Cancer
E. Hautamaki, D. Ngo, A. Tan, P. Chang
Cedars-Sinai Cancer

E. Hautamaki
Cedars-Sinai Cancer

29. Building IND Infrastructure to Ensure Compliance and Enable Growth
UNC Lineberger Comprehensive Cancer Center, University of North Carolina at Chapel Hill

30. Development of an Investigator-Initiated Trial Intake Process at Cedars-Sinai Cancer
E. Hautamaki, P. Chang, D. Ngo, A. Tan
Cedars-Sinai Cancer

31. Streamlining Data Collection: Implementation of an EDC FHIR Lab Interface
E. Crecelius, M. O’Dwyer, L. Logan, S. Balu, J. Frank, R. Johnson, R. Church, C. Lee, J. Morrison
UNC Lineberger Comprehensive Cancer Center, University of North Carolina at Chapel Hill

32. Starting Off on the Right Foot: Elevating the Voice of Community Stakeholders During the IIT Development Process
E. Monari, S. Szurek, A. Ivey, T. George, A. Anderson, E. Shenkman, C. Evans, A. Lawson-Ross
University of Florida Health Cancer Center

33. UF’s IIT Think Tank Experiment
E. Monari, A. Ivey, T. George, A. Anderson
University of Florida Health Cancer Center

Quality Assurance, Remote Monitoring, and Auditing

34. Proactive Quality Assurance Through Dual Review of Eligibility and Consent
K. Thorne
Huntsman Cancer Institute, University of Utah

35. Transforming Risk Management: Technological Evolution of MSK’s Clinical Research Quality Assurance Program
A. Granobles, M. Satter, S. Puleio, F. Puma, N. Brosnan, K. Yataghene
Memorial Sloan Kettering Cancer Center

36.*Virtual Monitoring and Auditing Digitization in Decentralized Clinical Trials: Source Document Verification, System Scheduling, and Real Time Protocol Performance Feedback
M. Buckley, J. Lengfellner, M. Latif, K. Yataghene, C. Houston, S. Terzulli, N. Cimaglia, P. Sabbatini
Memorial Sloan Kettering Cancer Center

*Honorable Mention

*Honorable Mention
2022 Abstracts

37. Automating Data Safety Monitoring Committee (DSMC) Progress Reports
T. McSpadden, S. Grotnic
University of Colorado Cancer Center

38. Introducing a Quality Management System Into the Mayo Clinic Cancer Center Clinical Research Office
K. Alexander, K. Croghan, A. Fritsche, J. Summer Bolster, J. Welte
Mayo Clinic Comprehensive Cancer Center

39. Preparing and Sharing Subject Cases for Remote NCTN Audit
K. Rygalski, M. Russell, D. Kitterman
University of Illinois Cancer Center

40. Standardized Quality Metrics in Cancer Clinical Trials: A Qualitative Study
Vanderbilt-Ingram Cancer Center

41. Digitalizing and Automating Clinical Research Protocol Regulatory Binders for Greater Efficiencies
M. Buckley, R. Lehrman, J. Lengfellner, M. Latif, K. Yapaghenie, C. Houston, S. Terzulli, P. Sabbatini
Memorial Sloan Kettering Cancer Center

42. Delegation of Authority – A Simplified Process
B. Scanlan, A. Holley, M. Kovak, B. Lehman, P. Newman, R. Perry, D. Wade, M. Birrer
UAMS Winthrop P. Rockefeller Cancer Institute

43. Optimization of a Regulatory eBinder Platform
S. Rebar, K. Lopez, D. Cervantes
Fred Hutchinson Cancer Center

44. Simplifying and Improving Training and Delegation Documentation
R. Kingsford, L. Hayes, L. Lujan
Huntsman Cancer Institute, University of Utah

45. Supporting Virtual Clinical Trials: How the Generation of DOAs in PIMS Has Enabled Clinical Trial Compliance in a Remote World
P. Lim
Memorial Sloan Kettering Cancer Center

Regulatory

47. The Effectiveness of an Innovative Competency-Based Education and Training Program on Decreasing Audit Findings
E. Dawkins, S. Cole, N. Nahimas, P. Seo, and J. Brown
Sylvester Comprehensive Cancer Center, University of Miami Health System

E. Cunningham, L. Dunham, B. Olsen
Karmanos Cancer Institute, Wayne State University

49. Implementation of Small Group Trainings to Expedite Initial Onboarding for Clinical Research Staff and Increase Connection Between New Employees
D. Kreitner, M. Wanchoo, D. Castro, C. Burgin
OHSU Knight Cancer Institute

40. Standardized Quality Metrics in Cancer Clinical Trials: A Qualitative Study
Vanderbilt-Ingram Cancer Center

Training and Career Development

46. Investing in Investigator Training: Developing Tools to Close the Gap
UNC Lineberger Comprehensive Cancer Center, University of North Carolina at Chapel Hill

47. The Effectiveness of an Innovative Competency-Based Education and Training Program on Decreasing Audit Findings
E. Dawkins, S. Cole, N. Nahimas, P. Seo, and J. Brown
Sylvester Comprehensive Cancer Center, University of Miami Health System

E. Cunningham, L. Dunham, B. Olsen
Karmanos Cancer Institute, Wayne State University

49. Implementation of Small Group Trainings to Expedite Initial Onboarding for Clinical Research Staff and Increase Connection Between New Employees
D. Kreitner, M. Wanchoo, D. Castro, C. Burgin
OHSU Knight Cancer Institute

50. Staffing Pipeline Creation: Clinical Research Internship for Undergraduate BIPOC Students
T. Cummings, A. Walens, A. Leak-Bryant, V. Carlisle, M. Haines, C. Lee
UNC Lineberger Comprehensive Cancer Center, University of North Carolina at Chapel Hill

51. Using Surveys to Evaluate Staff Onboarding Experiences: Pandemic to Present
C.L. Allen, P. Rose, M. Marcum, N. Kurtzweil
University of Cincinnati Cancer Center

52. Piloting a New Investigator E-Learning Onboarding Program
J. Thomas, M. Murphy, T. George, A. Anderson, E. Monari, A. Ivey
University of Florida Health Cancer Center

Trial Recruitment & Community Outreach and Engagement

53. Community Outreach, Community of Color Accrual: CS-Link Registration and OnCore Enhancements to Capture Ethnic Subgroups, Gender, and Sexual Orientation
J. Gomez, B. Rimel
Cedars-Sinai Cancer

54. Implementing Structured Assessments to Determine Research Readiness and Capacity Among Community-Based Clinical Oncology Network Sites
C. Crabtree-Ide, R. Evans, E. Bouchard, K. Noyes, M. Reid, L. Smith, K. Glaser
Roswell Park Comprehensive Cancer Center

55. Use of Epic My Reports to Increase Trial Accruals While Decreasing Pre-Screening Time Spent
A. Gatta, A. Kavadas, J. Davenport, K. McCaffrey, M. Kilbane
Taussig Cancer Institute, Cleveland Clinic Cancer Center

*Honorable Mention
Poster not present at meeting
2022 Abstracts

56. Improving Gender Diversity and Representation in Clinical Trials
S. Ward, C. Evans, D. Castro, C. Burgin
OHSU Knight Cancer Institute

57. Machine Evaluation of Catchment Area Relevance Through Text Mining
P. Arlen, J. Chakko, B. Mahal, G. DeGennaro
Sylvester Comprehensive Cancer Center, University of Miami Health System

58. Process Improvement of the Precision Medicine Program at Sylvester Comprehensive Cancer Center: An Exploration of Different Models to Increase Awareness and Clinical Trial Enrollment
P. Seo, J. Figueredo, J. Trent
Sylvester Comprehensive Cancer Center, University of Miami Health System

59. Prescreening GI Cancer Clinic Schedules for Clinical Trial Recruitment – Plan to Make a Difference in Study Accruals
J. Siddiqui, A. Loechtenfeldt, J. Parker
University of Cincinnati Cancer Center

60. Study Consent Rates and Decline Reasons at the University of Illinois Cancer Center
D. Kitterman, M. Russell, Y. Molina, O. Danciu
University of Illinois Cancer Center

61. Reduction of Days From Referral to Phase I Consultations
J. Bourgeois, T. Goodale, S. Mackoon, D. Arnett, E. Judson, C. Lewis
Winship Cancer Institute of Emory University

Trial Start-up and Activation

62. Collaboration to Develop Recommendations to Improve Trial Activation Timelines
T. Werner, T. Lin, C. Houston, D. Otap, M. Nashawati, L. Ashmore, E. Buell, A. Zafirovski, K. Much
Huntsman Cancer Institute, University of Utah; The University of Kansas Cancer Center; Memorial Sloan Kettering Cancer Center; Mays Cancer Center at UT Health San Antonio MD Anderson; Robert H. Lurie Comprehensive Cancer Center of Northwestern University; AbbVie; Genentech; Janssen Oncology; Merck

63. Practical Benefits of Defining and Implementing Structured Intake and New Study Assignment in a Centralized Start-up Model
A. McCauley, M. Winkler, M. Poduri, M. Hibbert
Fred Hutchinson Cancer Center

64. Four Years and Beyond: Progress With the Committee on Radiation
Memorial Sloan Kettering Cancer Center

65. It’s About Time: A Simplified Approach to NCI Trial Activation
Memorial Sloan Kettering Cancer Center

66. Strategies to Expedite Activation of Expanded Access Protocols at Memorial Sloan Kettering Cancer Center
X. Lekperic, E. Valentino, S. Hanley, A. Rodavitch
Memorial Sloan Kettering Cancer Center

67. Enhancing 1st Stage Protocol Review – A Quantitative Approach
L. Wall, A. Spratt, R. Szmulowitz
The University of Chicago Medicine Comprehensive Cancer Center

68. Evaluation of a Prioritization Matrix for Electronic Order Build in an Investigational Drug Service
A. Smith, K. Bottenberg, J. Rudolph, K. Redic
University of Michigan Rogel Cancer Center

69. Clinical Trial Research Group (CTRG) Guidelines for Trial Portfolio Management
Huntsman Cancer Institute, University of Utah

70. Technology and Centralization in Early Study Start-up Activities
E. Lebleu, L. Lujan, J. Moehle, T. Werner
Huntsman Cancer Institute, University of Utah

71. Enhancing Transparency and Interoperability: Developing an Enterprise-Level Portal to Streamline Trial Activation Processes
P. Arlen, M. Santiago, K. Williams, L. Thyssen, G. DeGennaro, A. Ward, N. Reyes, C. Valdivia
Sylvester Comprehensive Cancer Center, University of Miami Health System

72. Improving Trial Activation Timelines: A Comprehensive Process Improvement Project
P. Arlen, L. Thyssen, K. Williams
Sylvester Comprehensive Cancer Center, University of Miami Health System

73. Value Stream Mapping: Maximizing Value, Minimizing Waste, and Improving Flow Across the Clinical Trial Activation Process
P. Arlen, L. Thyssen, K. Williams
Sylvester Comprehensive Cancer Center, University of Miami Health System

74. Implementation of a Feasibility Committee – University of Cincinnati Cancer Center (UCCC) Study Operations & Administrative Review (SOAR)
A. Kastl, M. Marcum
University of Cincinnati Cancer Center

*Honorable Mention

*Poster not present at meeting
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