Policies and Procedures

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1 Introduction

The Alliance for Clinical Trials in Oncology (Alliance) was created in July 2011 by the merger of three National Cancer Institute (NCI) funded cancer cooperative groups: American College of Surgeons Oncology Group (ACOSOG), Cancer and Leukemia Group B (CALGB), and North Central Cancer Treatment Group (NCCTG). Per the Alliance Constitution, the missions of the three component cooperative groups were joined in the new vision statement:

“To reduce the impact of cancer on people by uniting a broad community of scientists and clinicians from many disciplines, committed to discovering, validating and disseminating effective strategies for the prevention and treatment of cancer.”

The Alliance receives grant funding from the National Cancer Institute (NCI). The Alliance is one of the Network Groups for the NCI National Clinical Trials Network (NCTN) and serves as a research base for the NCI Community Oncology Research Program (NCORP). The Alliance complies with the NCTN and NCORP Program Guidelines, related NCI policies and procedures and the Code of Federal Regulations (CFR). As an NCTN and NCORP Group, the Alliance utilizes centralized NCI systems for the management of clinical trials.

1.1 Specific aims

The Alliance is founded upon more than 60 years of cooperative group experience, but re-designed to meet the current challenges of cancer clinical and translational research. The Alliance is an experienced multi-institutional cancer clinical trials group that provides a comprehensive and highly efficient clinical trials infrastructure, access to experienced collaborators across all disciplines of oncology clinical trials research, and a diverse portfolio of trials for patients with breast, gastrointestinal, genitourinary, respiratory, central nervous system, hematological malignancies, and selected rare tumors.

As an integral component of the National Clinical Trials Network (NCTN), the Alliance has the leadership, experience, infrastructure, and member commitment required to achieve the scientific, operational, and collaborative aims outlined below.

1.1.1 Scientific aims

Alliance scientific programs conduct trials of highest possible clinical and translational impact that define new standards of care for patients with cancer. Programs have the following specific aims:

1. To conduct multimodality studies of adult cancers that include novel approaches to treatment and evaluation of patient outcomes based upon improved understanding of the molecular pathogenesis of these diseases
2. To develop treatments specific for molecularly defined disease subsets

3. To develop and implement novel clinical trial designs that facilitate evaluation of target-directed therapies

4. To introduce imaging response as a biomarker to direct therapy

5. To improve treatment outcomes by studying psychosocial adaptation to cancer, symptom management, and cancer survivorship

6. To study the unique therapeutic, psychosocial, economic, functional, and biological features of cancer in special populations including those with rare tumors, the elderly, underrepresented minorities, and those who are economically disadvantaged

1.1.2 Operational aims

The infrastructure of the three component cooperative groups has been merged into a single, fully integrated system that is optimally designed to serve the NCTN and NCORP research community. The Alliance operations units have the following specific aims:

1. To support a broadly based institutional member research network that includes a balance of academic and community researchers of all disciplines who are committed to conducting high impact cancer clinical trials

2. To provide operational capabilities for clinical and translational trials that are efficient, innovative, and make maximal use of available resources to achieve accurate and timely clinical trials results

3. To maintain responsible stewardship of important public resources, including clinical trials data and outcome-linked biospecimens, so that these can be used to conduct the best possible cancer treatment discovery and biomarker validation research

4. To train the next generation of investigators to meet the continuing challenges of cancer clinical and translational research

1.1.3 Collaborative aims

The Alliance is committed to collaborating with the NCI and all NCTN members to achieve the overall goals of the NCTN. Specific aims for collaboration include the following:
1. To participate to the fullest possible extent in clinical trials planning and management committees convened by the NCI including the Disease/Modality Specific Steering Committees, the Group Banking Committee, and other planning groups.

2. To collaborate with other network groups, cancer centers, Specialized Programs of Research Excellence (SPOREs), and selected organizations outside of the NCTN to optimally leverage available resources to achieve NCTN scientific objectives.

3. To promote accrual to all NCTN trials among its institutional members.

4. To practice responsible resource sharing in order to achieve the goals of the NCTN as a whole.
1.2 Overview of program structure

As outlined in the Alliance Constitution and Bylaws, the primary governance body of the Alliance is the Board of Directors, which represents the group’s institutional members. The Alliance is led by the group chair with assistance from the group vice chair. The Alliance is also supported by five program directors/principal investigators, each responsible for a specific program integrating discipline-related science and operational functions across all disease committees. The Executive Committee represents the Board of Directors and assists the group chair in planning and coordinating group activities.

The Alliance structure is disease-centered, with multi-modality involvement and significant input from both academic- and community-based researchers, full involvement of patient advocates, and routine participation of mentored junior investigators (see table 1-1). The group chair is responsible for the conduct and quality of scientific activities and efficient operation of the Alliance, represents the Alliance in its business with the NCI and other parties, and serves as the spokesperson for Alliance. The group chair directs eight multidisciplinary disease committees and six modality committees, and is responsible for central administration, finance, quality assurance and membership services.

Table 1-1. Alliance program structure

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**Alliance Policies and Procedures — Introduction 1-4**
In addition to the traditional modalities of surgery, radiation oncology, medical oncology, pathology and biostatistics, Alliance research is enriched by involvement of patient advocates, oncology nurses, community oncologists, and specialists in imaging, laboratory medicine, information technology, bioinformatics, outcomes and comparative effectiveness research, research ethics, and disparities research. In order to productively manage this deep scientific scope, Alliance uses a program approach that provides a structure to support researcher involvement and innovation in these many fields, yet maintains operational efficiency.

The Alliance includes five programs, each led by an Alliance program director who operates under the direction of the Group Chair to manage scientific, administrative, and operational activities of the group (see figure 1-1). Specifically, each program includes an operational unit and one or more scientific or administrative committees that report directly to the program director. Each program effectively interacts with the disease committees, and the program director is responsible for ensuring optimal integration of their program’s activities into study development and execution. For example, each disease committee requires the involvement of biostatistics (Statistics and Data Management Program), protocol development and study concept review (Central Protocol Operations Program), biomarker development and biorepository support (Translational Research Program), and cancer control research and community oncology participation (Cancer Control Program). The fifth program, the American College of Surgeons Clinical Research Program, provides an interface between the Alliance and the American College of Surgeons (ACS), a >75,000 member...
A professional organization that has led cancer care and research programs since its inception in 1913. The program-based structure of the Alliance is an innovative approach to management of cooperative group research. The Alliance programs create an interactive environment that fosters integration across disciplines and operational units, and has proved to be a highly effective structure for maximizing efficiency.

### 1.2.1 Office of the Group Chair

The Office of the Group Chair is responsible for administrative and fiscal affairs. This includes support for scientific leadership, administrative committees, membership services, regulatory compliance/audits, travel, meetings, financial services and grants administration. The Office of the Group Chair coordinates a per-case payment program, using funds provided by the NCI and other federal agencies, to defray the costs incurred by institutions in treating and following patients on the group’s clinical trials. The Office of the Group Chair is the communications hub of the Alliance, providing regular distribution of information essential for the conduct of group business to participating members, NCI, regulatory agencies (e.g., Food and Drug Administration (FDA), Office of Human Research Protections (OHRP), Institutional Review Board (IRB)), other NCTN groups and the general public. The office organizes all group meetings, coordinates communications, education and training, maintains the Alliance website, and produces a variety of publications, including a monthly newsletter.

In addition to the group chair, three senior leaders provide support to Alliance members through this office (see figure 1-1). The group vice chair stands in for the group chair for any responsibility within the Office of the Group Chair. The associate group chair for Cancer Center Collaborations ensures that

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**Figure 1-2. Office of the Group Chair**

*Alliance Policies and Procedures — Introduction 1-6*
Alliance research is optimally linked to cancer center clinical and translational programs. The associate group chair for Advocacy promotes patient advocacy initiatives.

Staff working within the Office of the Group Chair are located at Brigham and Women’s Hospital in Boston, MA, University of Chicago in Chicago, IL, and Duke University in Durham, NC. The CAO, CFO and the group vice chair share responsibility for overseeing the operational functions of the Office of the Group Chair. The CAO oversees administrative operations including grant preparation, communications, education and training, member roster tracking, audit and regulatory compliance monitoring, and other member services activities. Figure 1-2 illustrates the administrative, financial, and regulatory operations staff and their areas of responsibility.

1.2.2 Statistics and Data Management Program

The Alliance Statistics and Data Management Program, also referred to as the Statistics and Data Center (SDC), supports the activities of the group by achieving the highest standards for the conduct of clinical trials in terms of study design, statistical methodology, data management, protection of patients and their data, and regulatory compliance. The SDC also strives to continually improve the efficiency of Alliance systems and processes.

Two scientific committees, Biostatistics and Bioinformatics, serve as core resources for Alliance investigators at all stages of the study process from design to analysis and reporting. These scientific teams are also responsible for developing innovative statistical designs to improve the efficiency and reliability of Alliance trials.

In addition to its scientific committees, the Alliance SDC houses key operational functions for data management, study monitoring, and information systems. The SDC is primarily located at the Mayo Clinic in Rochester, MN with additional statistical staff at other locations. The leadership at each site works collaboratively to ensure optimal collaboration with Alliance investigators. Figure 1-3 illustrates the SDC senior leadership roles.
1.2.3 Central Protocol Operations Program

The Central Protocol Operations Program oversees the development and maintenance of all study protocols generated by Alliance scientific committees. Staff members manage the complex process of study protocol development, including scientific team coordination, study concept review and prioritization, including collaboration with the Translational Research Program (TRP) for their scientific review, NCI submission, protocol document development and maintenance, study budgeting, and coordination of study-specific logistics such as biomarker study funding (including collaboration with TRP to prepare BIQSFP applications), pharmaceutical and regulatory affairs (e.g., drug distribution and IND reporting), other regulatory logistics (e.g., NCI CIRB submission and credentialing) and implementation of accrual management plans. This process adheres to NCI Operational Efficiency Working Group (OEWG) timelines, which are carefully monitored by Protocol Operations Office staff. The Protocol Office also distributes protocols to sites and serves as focal point of communication for both study chairs and investigators throughout Alliance member institutions.

Once a protocol is activated, Protocol Office staff implement protocol amendments and other protocol communications, maintain all regulatory
support documentation, and serve on numerous NCI committees that work to improve the cooperative group process. The Central Protocol Operations Program represents the interests of the group in many study-specific negotiations with the NCI, pharmaceutical firms, other network groups, international collaborators, and the public. Figure 1-4 illustrates the Protocol Office staff and their areas of responsibility.

1.2.4 Translational Research Program (TRP)

With the advent of molecularly driven oncology, the Translational Research Program is essential for the development and execution of trials performed by each Alliance disease and modality committee. The TRP facilitates the scientific agenda by supporting the basic and translational researchers who work within Alliance committees. The TRP director, in collaboration with the chairs of disease committees, names a translational research leader for each disease. These individuals work within the TRP to ensure optimal integration of translational endpoints into Alliance trials. These researchers also promote successful collaboration between Alliance committees and researchers within Specialized Programs of Research Excellence (SPORES), cancer centers, and other research groups. In addition, discipline committees within the TRP, such as Pharmacogenomics and Population Pharmacology, Imaging, and Pathology provide both scientific input and operational support for Alliance translational research. Figure 1-5 illustrates the TRP senior leadership roles.

A key TRP operational component involves management of tissue resources collected during Alliance clinical trials. The TRP coordinates the Alliance Integrated Biorepositories, an operations unit with locations at The Ohio State University, Washington University Medical Center, the Mayo Clinic, and Brigham and Women’s Hospital. The TRP also manages a network of Molecular Reference Laboratories that provide specialty biospecimen services that are required for Alliance research protocols.

Figure 1-5. Translational Research Program

Alliance Policies and Procedures — Introduction 1-9
1.2.5 Cancer Control Program

Research in cancer control is integrated throughout the scientific programs and operations of the Alliance. The Cancer Control Program serves as the research base for the NCI Community Research Oncology Program (NCORP), as well as non-NCORP community oncology members. There are five scientific domains of the Cancer Control Program (CCP): Cancer Prevention, Symptom Intervention, Health Outcomes, Cancer in the Elderly, and Health Disparities. The Office of Director for Cancer Control oversees administrative components of the Cancer Control Program, including Leadership, Community Oncology Membership Services (including the Community Oncology Committee), Administrative/ Operations, and Pilot Projects/Consulting. Research conducted by the scientific committees of the Cancer Control Program is integrated with the Alliance disease committees and TRP so that each Alliance treatment study can be leveraged as appropriate to include cancer control endpoints. This integration occurs by placement of cancer control researchers and community oncology members in disease and modality committees. In addition, a leadership team reviews each Alliance trial concept for opportunities to contribute to cancer control research.

The activities of the Cancer Control Program are central to the work of the Alliance. In particular, the Cancer in the Elderly, Health Disparities, and Health Outcomes Committees are essential for achieving the goals of the cancer treatment trials program. Figure 1.6 illustrates CCP leadership roles. In addition, the Community Oncology Committee is responsible for ensuring participation by community oncology leaders in treatment trial and translational research study design and execution. A community oncology co-chair is required for every protocol.

Figure 1.6. CCP Leadership

1.2.5.1 NCORP Specific Aims
The overarching aim of the Alliance National Cancer Institute Community Oncology Research Program (NCORP) Research Base is to reduce the burden of cancer by conducting high-quality multidisciplinary, multi-site interventional and observational clinical trials, as well as database analyses. The NCORP places special emphasis upon issues affecting minority, underserved and elderly patient groups, and upon building strong collegial relationships with NCORP Community sites and Minority/Underserved Community sites. There are three specific aims within the overall research base:

1. To reduce the incidence and prevalence of clinically significant cancers

In support of aim 1, NCORP will reduce the incidence and prevalence of clinically significant cancers by a) identifying patients at greatest risk for developing specific cancers, b) screening those patients to detect early stage disease amenable to curative therapies, c) employing effective pharmaceutical interventions in at risk patients, and d) developing effective strategies to reduce individual behaviors that increase cancer risk. NCORP will also develop strategies to identify pre-symptomatic recurrent cancers in patients with a prior diagnosis of cancer, and who were previously treated with curative intent, in order to intervene with potentially curative therapies.

2. To alleviate the symptoms of cancer and the toxicities of cancer treatment, and

In support of aim 2, NCORP will alleviate the symptoms of cancer and the toxicities of cancer treatment by a) understanding the pathophysiology and natural history of untoward symptoms associated with cancer and/or cancer therapy, b) identifying factors that increase patient risk for these symptoms, and c) finding effective strategies for the prevention and treatment of such symptoms.

3. To improve the delivery of cancer care in community and academic practices.

In support of aim 3, to improve the delivery of cancer care in community and academic practices, NCORP will focus upon three strategies; a) patient-centered outcomes and
comparative effectiveness research, b) cancer economics, and c) systems redesign and organizational change.

To support each of these three primary aims, NCORP will conduct health outcomes research to improve understanding of the patient experience with disease, treatment, and survivorship. To achieve this mission, four research priorities were identified: a) to embed patient-reported outcomes (PROs) in Alliance clinical trials; b) to conduct primary PRO methodology research; c) to study relationships of genetic/biological mechanisms with PROs; and d) to evaluate the use of PROs to improve care delivery and quality.

NCORP will also identify and intervene to eliminate disparities in cancer incidence, morbidity, mortality, and clinical trial participation among underserved and minority populations. Strategies to reduce disparities are: a) to conduct stand alone and companion trials to assess and/or intervene to improve health disparities; b) to examine existing data in the Alliance to assess and/or monitor disparities among populations that experience disparities; c) to provide education, strategies on, and monitoring of accrual of underserved and minority populations to Alliance studies; and d) to integrate relevant community members and providers into the Alliance to facilitate identification of eligible populations, health disparities, and solutions to address disparities in cancer outcomes.

Finally, in support of each specific aim, NCORP will a) address treatment issues including efficacy among older cancer patients, b) improve quality of life and maintain and/or improve function among older cancer patients; and c) assess the role and potential value of geriatric assessment tools in cooperative group trials and to develop models for predicting toxicity and functional decline.

1.2.6 American College of Surgeons Clinical Research Program

The American College of Surgeons (ACS) has long-standing programs that define and improve the quality of cancer care. ACS is the parent organization of the Commission on Cancer (CoC), a consortium of professional societies that improves survival and quality of life for cancer patients through standard-setting, prevention, research, education, and the monitoring of quality care. ACS is also the sponsor of the National Cancer Database (NCDB), a joint program of CoC and the American Cancer Society. CoC develops and disseminates cancer care standards and tracks quality metrics to improve patient outcomes. One of the most important CoC quality metrics is participation in clinical trials, and each of the more than 1500 CoC sites across the United States has a clinical trials participation target of at least 5% of its cancer patients.
The mission of the Alliance/American College of Surgeons Clinical Research Program (ACS CRP) is to reduce the impact of cancer by increasing knowledge and awareness of new evidence and practice standards; increase the participation of community oncology surgeons in cancer research and cancer care activities; develop and implement evidence-based practices in surgical oncology; and create opportunities for meaningful health services research. The program has four committees that have unique goals and activities and that work together to reach the program's overall research goals. These committees include the Education Committee, Dissemination & Implementation Committee, Cancer Care Standards Development Committee, and Cancer Care Delivery Research Committee. The ACS CRP shares responsibility with the Alliance, ACS and the CoC for developing surgical standards for use in Alliance clinical trial protocols and CoC accreditation as well as for disseminating new evidence-based knowledge.

1.2.7 Member institutions

Membership in a network group is required for enrollment of patients on group protocols. Alliance member networks may be Lead Academic Participating Sites (LAPS) or NCORP networks. LAPS and NCORP institutional networks receive grants from the NCI to support their infrastructure and participation in NCI-funded clinical trials. Non-LAPS and non-NCORP institutions receive per-case payments from the Alliance NCI-grants to support their clinical trial participation.

A principal investigator and a co-principal investigator, who are responsible for managing the site according to all Alliance and NCTN policies, lead Alliance member institutions. Membership evaluation involves assessment of each site’s past clinical trials accrual and audit history, and requires that each potential member agree to adhere to the policies and procedures of the Alliance.
1.3 Committees

Alliance organizational structure, as defined in its Constitution and Bylaws, calls for its research agenda to be driven by a number of scientific committees, whose activities are supported by administrative committees with research infrastructure needs executed by operations units. Alliance is a large and diverse organization, working across many institutions. To permit optimal leadership and accountability, the group is structured into programs (see section 1.2). The assignment of Alliance committees and operations units to the group chair and to the programs is shown in table 1-1.

1.3.1 Scientific committees

Alliance trials are conducted by scientific committees of two types: disease committees and modality/discipline committees. Disease committees serve as the primary site of study concept generation. Modality/discipline committees foster cross-disease participation of a modality or discipline in Alliance research. Scientific committee chairs are either proposed by the group chair or, for those committees within Alliance programs, are nominated by the appropriate program director. The Alliance Executive Committee approves all chair appointments. Each scientific committee chair names several vice chairs, who are also approved by the Executive Committee. Committee members are appointed by the committee chair with input from the vice chairs and from modality/discipline committee leaders.

Diversity of leadership and membership is built into the scientific committee structure. The committee leadership (chair plus several vice chairs) must include representatives from medical oncology, surgical oncology (for solid tumor committees), radiation oncology, translational research, and transplantation (leukemia and myeloma committees). Alliance disease committees include, at a minimum, two representatives each from the disciplines of medical oncology, surgical oncology (solid tumor committees), translational research, radiation therapy, community oncology practices and young investigators (those within five years of fellowship completion), as well as patient advocates and liaisons from Cancer Control committees, as applicable.

Alliance research is supported by a number of committees that ensure participation of essential modalities and disciplines in trial design and execution. As cancer research has become more complex and specialized, the number and variety of these committees has increased. Several committees play important roles in designing and executing Alliance trials. Some modality/discipline committees, however, are not sites of study concept development, but instead provide focal points for member involvement,
enable collaboration and increase the impact of Alliance trials by promoting interactions with key member groups.

1.3.2 Administrative committees

Administrative committees conduct business as required to ensure the effective and ethical operation of the Alliance. Administrative committees reporting directly to the Board of Directors include the following: Membership, Institutional Performance Evaluation, Audit, and Constitution and Bylaws. The chairs of each of these committees are proposed by the group chair, and approved by the Board of Directors. Administrative committees reporting directly to the Executive Committee include: Data and Safety Monitoring Board, Conflict of Interest, and Publications.
2 Institutional membership

Members of the Alliance will be institutions meeting all requirements for membership, which include accrual, data quality and timeliness, adherence to Alliance policies and procedures, and participation in Alliance scientific activities. See the Alliance Bylaws for additional details. Institutional member networks consist of a main member with or without affiliates or components.

2.1 Membership criteria

Refer to the Alliance Bylaws sections 1-4 for qualifications for prospective members.

The Membership Committee considers the following aspects in their evaluation of prospective members:

- Multi-disciplinary institutional resources for clinical trials
- Scientific interests
- Prior clinical research experience
- Level of participation in cancer research cooperative group trials
- Patient population
- Prior institutional performance evaluation metrics
- Satisfactory audit results
- Other regulatory considerations
2.2 Applying for membership

The Alliance reviews institutional membership applications monthly or as needed. The institutional membership application is available on the Alliance public website under the Membership tab (http://www.allianceforclinicaltrialsinoncology.org). An application will be reviewed by the Membership Committee only if the institution has an active NCI ID and FWA. The Membership Committee evaluates the completed applications for appropriateness of facilities, institutional resources and past performance in clinical research. Following a decision by the Membership Committee, applicants will be notified of approval status. If the Membership Committee approves the application, it then submits its recommendation for approval to the Board of Directors for vote. Refer to the Alliance Bylaws section 5 for additional details regarding the membership evaluation procedure.

Affiliate applications can be approved by the Membership Committee without Board approval.
2.3 Membership activation

If the Board of Directors approves the Membership Committee’s recommendation for approval, applicants will receive a notification of approval status with additional information. Alliance staff will activate the member on the Alliance roster in the Cancer Trials Support Unit (CTSU) Regulatory Support System (RSS) and the Clinical Trials Monitoring Branch (CTMB)-Audit Information System. Alliance staff will manage the PI and Lead CRP roles in the institution roster(s). The Lead CRP will add persons and person roles to the institution roster via the NCORP Management System (NCORP SYS) or CTSU Roster Update Management System (RUMS). Upon activation of Alliance membership, the institutional network will be granted access to the Alliance website and Alliance Web applications. Alliance members will have access to clinical trials on the CTSU menu.

2.3.1 Roster

2.3.1.1 A site must be included on the roster if it meets the following definition of engagement in research as defined by OHRP (45 CFR part 46). An institution is engaged in a particular non-exempt human subjects research project when its employees or agents for purposes of the research project obtain:

1. Data about the subjects of the research through intervention or interaction with them
2. Identifiable private information about the subjects of the research, or
3. The informed consent of the human subjects for the research

2.3.1.2 NCI Tiers

The Alliance adheres to the institution membership structure as mandated by the NCI. There are four types of member networks that are structured based on their funding mechanism. The member networks can have up to 3 levels (tiers) of member types:

- Tier 1 members of Lead Academic Performance Site (LAPS) and NCI Community Oncology Research Program (NCORP) represent the administrative offices of the member network. Tier 1 of the Main member networks (non-LAPS, non-NCORP) can either be an administrative office of a health system (if approved by CTSU) or an accruing institution.
• Tier 2 members include affiliates of Main members, NCORP components, and LAPS Main member, LAPS affiliates, LAPS Integrated components as identified in the LAPS grant. LAPS can also have aligned affiliates. Aligned affiliates are institutions/performance sites that are affiliated with the LAPS network but are not included in the LAPS grant. The Alliance Operations Center grant provides the per case payments for aligned affiliates.
• Tier 3 members are sub-component/sub-affiliates. A sub-component or sub-affiliate is an institution or practice site that shares the same FWA, IRB, governance structure, employees of either a Tier I or Tier II member. An example of a sub-affiliate is a physician practice that has a primary clinical site and has additional office locations where the same physicians treat patients. The primary clinic site is the parent and the additional locations are sub-affiliates.

2.3.2 Regulatory documentation

Regulatory documentation includes: documentation that the institution has a current Federalwide assurance (FWA) with the Office for Human Research Protections (OHRP); current Food and Drug Administration (FDA) 1572 forms and financial disclosure forms (FDFs) for all investigators; and certification that all investigators have received training in Human Subjects Protection (HSP) and Good Clinical Practice (GCP).

NCI policy requires all persons participating in any NCI-sponsored clinical trial to register and renew their registration annually. Registration is accomplished via the NCI Registration and Credential Repository (RCR). Additional details can be found on the NCI/CTEP website.

2.3.3 Financial documentation for institutions

Financial documentation for institutions includes a services agreement signed by the principal investigator and institutional official and W-9 form confirming correct legal name and tax-ID of the institution.
2.4 Responsibilities of a main member

The principal investigator will be required to sign a membership agreement that includes a summary of key policies and procedures, including conflict of interest, scientific misconduct, membership accrual requirements, confidentiality, audit requirements, institutional performance and publications.

The main member institution is responsible for all aspects of conducting Alliance clinical trials within its network. The main member is responsible for monitoring the conduct of a study both at the main member and all sites within its network.

Responsibilities are listed below. Affiliate and sub-affiliate institution have their own unique characteristics but each main institution must be sure that mechanisms are in place so that these responsibilities are met.

2.4.1 Communications

The main member institution must confirm that all research staff have access to the Alliance electronic distribution of information. This information includes new protocols, addenda, memos, letters, and miscellaneous items from the Alliance. The Alliance clinical research office at the institution is frequently located in the oncology or hematology department of a hospital or medical school and it is vitally important that a good communications network is established so that Alliance members from other modalities (e.g., pathology, radiation oncology, surgery, transplant, imaging, correlative sciences) receive information on a timely basis regarding Alliance protocols, meetings, and other relevant topics. It is the responsibility of the main member to assure that their network institutions have the same type of communications network established to distribute information to all disciplines within the affiliate.

2.4.2 Electronic communication

The Alliance makes use of electronic mail and the website to provide information to its members. It is the responsibility of the main member to confirm that participants are able to access this information. The Alliance requires all members to have a unique e-mail address. All network and site PIs, Co-PIs, Lead CRPs and Secondary Lead CRPs are required to receive broadcast emails.
2.4.3 Management of network data

Data forms should be submitted according to specifications in the protocol. The main member is responsible for the data quality and timeliness of their network sites.

If an affiliate institution changes main member networks, the new main member becomes responsible for the timely submission of data for all Alliance patients at the affiliate institution, including patients registered through the previous main member.

A main member institution is responsible for collection of data for patients at an affiliate institution even if that affiliate is dropped from the network. The Institutional Performance Evaluation Committee (IPEC) includes, in its evaluation of a main network, patients from dropped affiliates who are still in the evaluation window.

2.4.4 Investigational drug handling

All affiliates order drugs directly from either the NCI or from a private source as specified in the protocol. However, the main member is responsible for ensuring that all federal regulations regarding investigational drugs are adhered to by the main member and the affiliates. Annually, each Alliance investigator must sign a FDA Form 1572 stating that the investigator will adhere to the federal regulations and each main member should confirm that its investigators are in compliance and have a current FDA Form 1572 on file with the Pharmaceutical Management Branch. Each institution that orders drugs is responsible for any protocol specific requirements related to drug ordering and shipping. Refer to the Investigator’s Handbook on the NCI/CTEP website for more specific investigational drug information. See also Section 12, Investigational Product.

2.4.5 Human subjects protection

The main member is responsible for ensuring that all federal regulations are adhered to regarding protection of human subjects. No patient may be entered on a study until the protocol has been reviewed and approved by the IRB of record for the institution where the patient is being treated. Alliance protocols also require a patient to sign an informed consent and the registering institution must confirm that the informed consent has been signed before the patient can be registered to the study. Protocol-specified research interventions, including interventions conducted at a facility external to the registering institution, must be covered under an IRB approval.
2.4.6 Training

The main member serves as a resource for institutional personnel to further their understanding of clinical studies and to expand and encourage participation in the studies. Training programs should be provided for all personnel. The Alliance conducts education and training sessions during the Alliance Group meetings and posts educational resources on its website. All Alliance members are encouraged to participate in these training opportunities.
2.5 Institutional roles and responsibilities

2.5.1 Main member principal investigator

2.5.1.1 Network responsibilities

The main member principal investigator (PI) is responsible for the conduct of Alliance activities at a main member institution and for the integrity of all data submitted from the institution’s affiliate network. The PI is ultimately responsible for the conduct of research and regulatory compliance at affiliate institutions. The PI is responsible for managing the funds to support the work of the Alliance at their institution, and receive other funds from the Alliance in support of Alliance activities.

The obligations of institutional membership are set forth elsewhere in these policies. It is the job of the PI to ensure that these are met by all institutions in the network or to correct deficiencies in institutional performance that are documented by Alliance mechanisms, set forth elsewhere in these policies.

Each main member institution shall also have a co-principal investigator, who shall assume responsibility in place of the principal investigator if for any reason the principal investigator is unable to perform duties required for Alliance institutional membership.

Each affiliated institution in a network must name a responsible principal investigator. This PI may be the main member PI or another investigator responsible for clinical trial conduct at the affiliate institution with oversight from the main member PI.

2.5.1.2 Institutional responsibilities

Membership in Alliance is granted to an institution not an individual. It is the institution's responsibility to ensure that the Alliance research program is vigorously and competently administered at that institution, and to recommend to the group chair and Membership Committee, as appropriate, changes in the institutional PI. Although the Membership Committee considers the qualifications of PIs when approving institutions for membership in the Alliance, and must acknowledge changes in PI when proposed by the institution, the Alliance is not involved in
the nomination or selection process which occurs at the institutional level.

The PI receives Alliance communications concerning activities at his/her institution, or appoints individuals to act on behalf of the PI for these purposes. The PIs name individuals from their institutions as authors on Alliance publications, according to Alliance guidelines on publication. The PI takes responsibility for the performance of their institution's interdisciplinary team of Alliance participants, and for the introduction of new scientists to Alliance activities. The PI ensures that specialists from relevant oncology disciplines are available within the institution to support the activities of Alliance; makes certain that the institution meets minimum accrual standards required to maintain Alliance membership; and oversees all aspects of data and specimen management for Alliance studies within the institution. The PI also ensures that Alliance studies are conducted with appropriate attention to the protection of human subjects in research, all applicable regulations and that the physicians who oversee the conduct of Alliance studies disclose potential conflicts of interest. The PI ensures that the delegation of authority and tasks is documented and that research personnel are adequately trained.

2.5.2 Affiliate member principal investigator

The principal investigator (PI) for an affiliate institution is responsible for the conduct of Alliance activities at an Alliance institution, human subjects protection and the integrity of all data submitted from the institution.

These responsibilities are similar to the responsibilities of the principal investigator at the main member institution.

2.5.3 Clinical research professionals

Clinical research professionals (CRPs) at an Alliance institution may include clinical research associates (CRAs), surgical CRAs, oncology research nurses, and others. In general, responsibilities for CRPs at an Alliance institution include the following:

- Obtain IRB approval for Alliance protocols, consent forms, annual continuing review, and any protocol amendments that require IRB approval
• Obtain patient consent (and re-consents, when appropriate) for participation in Alliance studies
• Maintain study-specific regulatory and training files
• When authorized, register consented eligible patients to Alliance studies.
• Submit accurate protocol-required data, specimens and supporting documents according to protocol requirements
• Maintain a research record of supporting documents for each Alliance patient
• Participate in Alliance audits at the institution
• Maintain a patient notification policy

2.5.3.1  **Lead CRP**

Each Alliance institutional network must designate a lead CRP to receive and distribute communications from the Group and be the primary clinical research professional contact for the network. A secondary CRP should be designated to serve as a backup to the lead CRP. Institutional responsibilities of the lead CRP vary by network.

2.5.4  **Withdrawn or terminated institutions**

If an institution is withdrawn from the Alliance or terminated by the Alliance, the institution will remain responsible for data submission until such time that there are no longer patients in treatment or follow up, or the patient(s) are transferred to another Alliance member. The main member remains responsible for data from withdrawn affiliates.
2.6 Office for Human Research Protections assurances

2.6.1 Assurances

The regulations require that each institution engaged in the conduct of research involving human subjects provide a written assurance of compliance that it will comply with the requirements set forth in these regulations. The document is referred to as an assurance. Each assurance sets forth the commitment of the institution to employ the basic ethical principles of the Belmont Report and to comply with the regulations. There are several kinds of assurance documents. Where an independent investigator is to provide an assurance of compliance to OHRP the document is called an agreement.

Under the Department of Health and Human Services (HHS) human subjects protection regulations at 45 C.F.R. 46.103, every institution engaged in human subjects research supported or conducted by DHHS must obtain an assurance of compliance approved by the Office for Human Research Protections (OHRP).

All institutions applying for membership in the Alliance that do not currently have an assurance must obtain a Federalwide Assurance (FWA). The institution is responsible for ensuring that all institutions and investigators engaged in its U.S. federally supported human subject research operate under an appropriate OHRP or other federally approved assurance for the protection of human subjects.

2.6.2 Reporting institutional assurance compliance

The institution’s FWA must be included with the member’s roster information and remain current. Alliance must have documentation that there has been prospective review, at least annual continuing review, and review of significant protocol updates.
2.7 Institutional Review Boards

Each Alliance member institution must have an approved institutional review board (IRB) under the HHS Regulations for the Protection of Human Subjects (45 CFR 46) in order to enter patients on Alliance protocols. The IRB must follow the federal regulations regarding IRBs. The IRB must also be registered with the Food and Drug Administration (FDA). If the NCI Central Institutional Review Board (CIRB) is utilized by the local IRB through facilitated review, the CIRB is considered the IRB of record.

At the time of institutional audit, the performance of the IRB with respect to review of Alliance protocols and protocol amendments is evaluated. In addition, consent forms used within the institution are examined in order to determine whether they meet the standards required by OHRP. For institutions using CIRB, documentation of CIRB approvals including the CIRB Facilitated Review Acceptance Form will be reviewed, as well as the local informed consent form.

The Alliance may take various actions including suspension of accrual by an institution when it receives information from any source alleging that an IRB fails to comply with federal regulations. In such instances, Alliance informs the CTMB and an audit team may be assembled by staff at the CTMB, in conjunction with OHRP and the Office of Research Integrity (ORI).

2.7.1 Reporting and review requirements

As noted above, the Alliance must have documentation that there has been prospective review, at least annual continuing review and the review of significant protocol updates. IRB approval documentation is submitted to the CTSU. This information is entered into the CTSU/RSS database and is referred to when a patient is being registered. Documentation must state the type of review, list the protocol number (and if it is a review of a protocol update, it must list the protocol update number) and an IRB member or administrator must sign it. The protocol number and the update number, if applicable, must be clearly documented. Initial and continuing review documents must be submitted to the Cancer Trials Support Unit (CTSU) and Alliance staff will access the information in the CTSU database.

Annual continuing review must continue as long as patient data are being submitted. However, if no patients are currently receiving treatment and only data are being submitted, the Alliance accepts expedited review. Institutions must continue to submit studies that are not yet terminated to their IRB for continuing review. The Alliance audit team confirms that informed consent was obtained after initial review and that appropriate continuing review and significant protocol updates have taken place.
2.7.2 Federal record-keeping requirements for IRBs

The institutional review board that reviewed the study must keep records and minutes of the review per the federal guidelines. Institutions retain their discretion to organize and store IRB records in any manner that is consistent with the requirements of HHS regulations at 45 CFR 46.115. Electronic storage is acceptable as long as all records are accessible for inspection and copying by the Alliance, OHRP, FDA and other regulatory agencies, as applicable.
2.8 Institutional audits

2.8.1 History

As the world's largest sponsor of clinical trials of investigational antineoplastic agents and cancer clinical trials, the National Cancer Institute (NCI) must ensure that research data generated under its sponsorship are of high quality, reliable, and verifiable. The NCI quality assurance and monitoring policies for clinical trials have been in evolution since the start of the National Clinical Trials Network (formerly the Clinical Trials Cooperative Group) Program in 1955. One important aspect of the quality assurance program is that investigators in the NCTN undergo peer review as part of the funding process. As the NCI clinical research program has increased in size and complexity, the systems for quality control became more formal and systematic.

In 1982, the NCI made on-site monitoring a requirement for the Clinical Trials Cooperative Group Program, cancer centers, and any other investigators conducting clinical trials under its IND sponsorship. Because quality control and assurance programs were in place in many cooperative groups, the NCI delegated much of its responsibility for on-site monitoring of investigational agent studies and clinical trials to the cooperative groups. The guidelines were later expanded to include monitoring of Community Clinical Oncology Programs (CCOPs) components by cancer centers that serve as their research bases.

In 2014, the Cooperative Group Program was replaced by the NCI National Clinical Trials Network (NCTN) program. In addition, the Community Clinical Oncology Program (CCOP) combined with the NCI Community Cancer Center Program (NNCORP) to create the NCI Community Oncology Research Program (NCORP).

2.8.2 Quality assurance

Since the multicenter nature of group trials presents obvious questions about variability, the groups long ago recognized the need for formal quality control and monitoring. Procedures were developed to monitor the overall progress of studies and for ensuring adherence to protocol and procedural requirements.

The groups perform two distinct kinds of monitoring. The first is periodic review of the overall progress of each study to assure that the projected accrual goals are met on a timely basis, that over accrual is avoided, that eligibility and evaluability rates do not fall below minimum acceptable
standards, and that risks are not excessive. The groups perform this function at least semiannually prior to their group meetings.

The second type of monitoring is a systematic and independent audit of trial related activities and documents to assure the quality of trial execution at the level of the investigator. The audit process enhances the delivery of accurate and reliable clinical trials data and results according to the protocol, sponsor’s standard operating procedures, applicable regulatory requirements, and good clinical practices (GCP). This is commonly an on-site process, and consists of reviewing a subset of patients on a trial. The audit program assures that the data used to analyze the trials are an accurate reflection of the primary data. The program requires an on-site comparison of the submitted data with the primary medical record for a sample of patient cases. At the same time, compliance with regulatory requirements for the protection of human subjects and investigational drug accountability are checked. The audit also provides educational support to the clinical trials sites regarding issues related to data quality, data management, and other aspects of clinical research quality assurance.

Also included in these central quality assurance measures is the assessment of protocol compliance. This is done in an increasingly systematic way and on an ongoing basis. For example, most groups conduct central pathology review for selected studies to reduce variability in diagnosis. To ensure adherence to protocol-specified treatment, radiotherapy films and surgery reports are also monitored centrally. Checks of submitted data sheets for protocol compliance ensure that treatment is delivered according to protocol stipulations and that appropriate study tests have been obtained. The study chair and/or the statistical center are responsible for confirming each case's eligibility and evaluability, based on the information gathered through these quality control mechanisms.

2.8.3 NCI audit participation

The Clinical Trials Monitoring Branch (CTMB) of the NCI maintains oversight responsibility for the network group auditing programs. The most recent CTMB Audit Guidelines for the establishment of auditing programs have been incorporated into the Alliance policies. The complete federal document can be found on the NCI/CTEP website (NCI Guidelines for Auditing Clinical Trials). The CTMB Guidelines may be referenced for any policies and procedures that are not specified within the Institutional Audits Policy.

CTMB staff reviews all audit schedules and all reports of audit findings. To assure consistency of auditing across the group/cancer center research bases,
a CTMB representative may attend on-site audits. Staff from the CTMB may make specific recommendations for action if they do not believe the action taken by the network group or site has been adequate.

The CTMB, as part of their clinical trials auditing service, contracts review of some audits. The role of the NCI representative is to monitor the audit process and to ensure that the requirements of the CTMB for auditing are being met. They review the audit case reports prepared by the auditors, assess the audit exit interview, participate in the pharmacy audit, etc. and provide the CTMB with a detailed report on the conduct and outcome of the audit.

2.8.4 **Overview of Alliance auditing policies and procedures**

The Alliance Audit Committee was developed to provide assurance that the data reported on Alliance research records, of all types, accurately reflect the data as reported in the primary patient record.

To ensure that data management practices in each Alliance institution adhere to protocol guidelines, submitted information is accurate and complete, and all Federal Human Subjects regulations and NCI guidelines for investigational drugs have been followed, the audits conducted of member institutions examine a meaningful and random sample of the following:

- Clinical records and abstracts
- Imaging reports and techniques
- Pathology, cytochemistry and RT submission compliance, if applicable
- Operative reports
- Laboratory data
- IRB reviews and consents
- Investigational drug compliance documents

2.8.5 **Scheduling of audits**

2.8.5.1 **Selection of main member and affiliate member institutions for audit**

All institutions are audited at least once every 36 months, but all are at risk for audit during any one year. New main member institutions are audited no longer than 18 months after entry of the first patient to assure performance standards are being met and as an educational experience for the new investigators and their staff. The initial audit may be sooner based on accrual. Initial audits are conducted on-site. Routine audits will be scheduled within 36
months after the previous audit. For high accruing main member institutions, it may be appropriate to audit these institutions on a more frequent interval given the high number of cases for review.

The Alliance Audit Program may request main members to conduct on-site pharmacy audits of their affiliates, utilizing the same on-site audit procedures used by the Alliance. If requested, each main member must appoint a pharmacy audit liaison to manage the affiliate pharmacy audits. The audit liaison should be a member of Alliance who is versed in the Alliance’s audit policies. All pharmacy audit liaisons should have previous auditing experience and/or are required to participate in training sessions and/or modules. Physicians and staff from affiliates may not audit another affiliate.

Alternatively, these affiliates may be audited when the Alliance conducts the on-site audit of the main member institution.

Affiliate institutions must provide all required documents to conduct the audit at the main member institution the day of the audit or earlier if determined by the Alliance. It is strongly recommended that a representative from the affiliate be present at the main member institution during the audit. A separate Preliminary Report of Audit Findings and Final Audit Report are required for the main member institution and each affiliate institution audited.

An effort will be made to audit a pharmacy on-site at least every other audit, including a re-audit, if the deficiencies are related to drug inventory and the institution has registered patients on one or more studies with IND agents since the previous audit.

### 2.8.5.2 Scheduling audits for NCORPs and NCORP components

One audit will usually be conducted for the NCORP as a whole. Protocols and patient cases must be selected for review from each component where accrual has occurred. If the NCORP is audited as one entry, only one preliminary report and final audit report is required. This is the preferred method for auditing NCORPs and their components. Alternatively, the NCORP components may be audited as a separate entity.
2.8.5.3 **Scheduling of audits for inactive sites**

Institutions remain at risk for audit even if their membership in the Group is no longer Active, since they have made a commitment to long-term follow-up of patients with provision of good quality data according to the study schedule.

2.8.5.4 **Single-Site Audit Initiative (Multi-Group Audits [MGA])**

Certain sites/organizations may be subject to audit by more than one Network Group at the same time. This CTMB and CTSU initiative is intended to promote more efficient auditing practices, and are conducted according to these audit guidelines. These audits are coordinated by the CTSU.

2.8.5.5 **Case/protocol selection**

A minimum of four protocols representing studies conducted at the site should be selected when applicable. Emphasis should be given to registration trials, IND, multi-modality, advanced imaging studies, and prevention/cancer control trials, as well as those with high accrual.

A **minimum** number of cases equivalent to 10% of patients accrued since the last audit will be reviewed. The 10% of cases reviewed apply to each participating site being audited. For selection purposes, the 10% of chosen cases will always be rounded up. For selection of patient cases the following apply where appropriate:

1. **10% Group/NCORP cases**
2. **10% from protocols with advanced imaging studies/imaging studies embedded in treatment protocols**
3. **10% of DCP cancer control/prevention cases**
4. **A patient case from every registration trial must be selected for audit. This includes every NCI site Code being audited.**

While most cases will be selected from patients accrued since the previous audit, any patient case may be at risk for selection for
audit. In addition, at least one or more unannounced cases will be reviewed, if the total accruals warrant selection of unannounced cases. These cases may have a limited or full audit review. A limited review may include reviewing the patient informed consent document, patient eligibility and general data quality. However, if the unannounced cases only receive a limited review, these cases do not count towards the minimum of 10%.

Random selection of patient cases is used as often as possible balanced with the need to consider other factors such as date of enrollment, case complexity, treatment arm, etc.

2.8.5.6 Notification of audit

Institutions are notified of the date of the audit at least three months prior to the audit, although in some special circumstances the interval may be shorter. A list of the cases selected is sent to the institution 14-28 days prior to the audit to allow adequate time to prepare.

2.8.5.7 Audit team

Audit team members include Alliance audit staff and members of the Audit Committee. Principal investigators and clinical research professionals from any Alliance institution may also be asked to serve as ad hoc auditors. The auditors must be knowledgeable about the protocols to be reviewed, Alliance audit procedures, clinical trials methodology, NCI policies, and Federal regulations. All auditors must complete Alliance auditor training prior to their first audit and must maintain a signed confidentiality agreement on file at the Chicago office of the Alliance.

Alliance auditors will not complete site-specific training, such as EMR, HIPAA, etc, but will maintain a current human subjects training certification.

Each main member or NCORP principal investigator is responsible for recommending physicians who are able to serve as physician auditors.
2.8.6 Audit preparation by the institution

Principal investigators and institutional clinical research professionals are responsible for preparing for an audit.

The institution is responsible for ensuring that all relevant materials are available for review. If an institution is audited off-site at the Network Main Member, NCORP, or LAPs main member, the following records must be available:

1. IRB approvals, continuing reviews, amendment approvals, and safety reports.
2. Current versions of requested protocols.
3. Current locally utilized informed consent forms along with applicable model consent forms.

Note: The regulatory items above may be requested prior to the audit. At least three local consent forms will be audited.

4. NCI Drug Accountability Record Forms (DARFs) for control and satellite pharmacies, agent receipts, returns/destruction logs, transfer records, and/or logs for imaging/radiopharmaceutical agents.

Note: The pharmacy should be alerted that the auditors may conduct an on-site inspection of storage, security, and temperature monitoring logs. The pharmacy items above may be requested prior to the audit.

5. Complete medical records.

Note: De-identified source documentation is not acceptable. When imaging is used for disease response, physician auditors may request to review images.

6. Other relevant source documents or information, e.g. reports from the Imaging Core Laboratories, Central Laboratory/Pathology reports, etc.

7. For imaging studies: source documents/worksheets used for imaging acquisition, processing, quality assurance documentation, reader’s interpretation, record of imaging
administration, patient/study participant monitoring (vital signs, monitoring of contrast reactions, etc.) and log of staff signatures and imaging responsibilities.

For comprehensive instructions on preparing for an audit, please see the information posted on the Alliance website.

### 2.8.7 Conduct of an Alliance audit

The auditors review specific data relating to regulatory requirements and research.

#### 2.8.7.1 Regulatory requirements

An audit consists of reviewing and evaluating (1) conformance to IRB, informed consent content requirements, and maintenance of delegation of tasks log (if applicable) (2) drug accountability and pharmacy compliance including the use of NCI DARFs, or NCI approved drug accountability forms, and (3) individual patient cases. During the audit, each of these three components are independently assigned an assessment of either Acceptable, Acceptable Needs Follow-up, or Unacceptable, based on findings at the time of the audit. Assessment is based on evaluation of critical, major and lesser deficiencies.

For each component rated as Acceptable Needs Follow-up or Unacceptable, the institution is required to electronically submit a written response and/or Corrective and Preventive Action (CAPA) plan to Audit@AllianceNCTN.org. Once approved by the Alliance, the CAPA plan will be forwarded to the CTMB. The approval of CAPA plans does not constitute approval of site-specific policies and procedures. Each audit report indicates the date the Alliance must receive the response/CAPA plan. If the plan is not received and approved by the date indicated in the audit report, patient registration may be suspended at that institution.

A re-audit is mandatory for any component rated as Unacceptable. Depending on the individual circumstances a re-audit may also be scheduled when the result is designated Acceptable, Needs Follow-up.
2.8.7.1.1 Critical, Major and lesser deficiencies

Deficiencies are categorized as either “critical”, "major" or "lesser"; examples are provided in the appropriate sections. An exhaustive list of examples is not given, but the examples are intended to guide the reviewers in their assessment and categorization of specific deficiencies. Deficiencies too trivial to warrant comment are not included in the report.

Critical deficiency: any condition, practice, process or pattern that adversely affect the rights, safety or well-being of the patient/study participant and/or the quality and integrity of the data; includes serious violation of safeguards in place to ensure safety of a patient/study participant and/or manipulation and intentional misrepresentation of data.

Major deficiency: a protocol variance that makes the resulting data questionable.

Lesser deficiency: a deficiency that does not affect the outcome or interpretation of the study and is not described as a major deficiency. An unacceptable frequency of lesser deficiencies is treated as a major deficiency.

2.8.7.2 Review of IRB documentation and informed consent content

See section 5.2 of the CTMB Audit Guidelines for complete details concerning IRB documentation and informed consent content.

2.8.7.2.1 IRB documentation

Before a patient enters a study, all federal requirements for the protection of human subjects must be met. Every institution must have documentation of IRB approval.

Maintaining a separate chronologic file for correspondence regarding IRB information for each protocol is recommended so that information regarding annual renewals and changes in protocols is readily available for audit review.
Documentation of initial IRB approvals with the IRB chair's signature and date, annual re-approvals for each audited protocol and approval for amendments should be available at the site visit for review by the audit team. The same is true for IRB review of safety reports. If an institution being audited is covered by another institution's IRB, the written agreement should be available for review.

For institutions that use the NCI Central Institutional Review Board (CIRB) as their IRB of record for particular trials, the following items must be provided for auditing:

1. Initial approval letter from CIRB to the Principal Investigator (PI) for study activation
2. CIRB Approval of the Annual Signatory Institution Worksheet About Local Context
3. Documentation that IRB approval was obtained prior to patient registration
4. Reporting of any unanticipated problems, serious non-compliance and/or continuing non-compliance problems per OHRP/FDA policy
5. Other correspondence with CIRB such as annual re-approvals, protocol amendments, etc.

Critical IRB deficiency:
- Any finding identified before or during an audit that is suspected to be fraudulent activity

Major IRB deficiencies may include but are not limited to:
- Initial approval by expedited review for protocols requiring full board review per OHRP guidelines.
- Expedited re-approval for situations other than approved exceptions.
- Registration and/or treatment of patient prior to full IRB approval.
- Re-approval delayed more than thirty days, but less than one year.
- Registration of patient on protocol during a period of delayed re-approval or during a temporary suspension (i.e., Request for Rapid Amendment).
- Missing re-approval.
• Expired re-approval.
• Internal reportable adverse events reported late or not reported to the IRB.
• Failure to submit or submitted after 90 days, any reportable external safety report to the IRB that is considered an unanticipated problem as defined by OHRP, unless there is a IRB policy that does not mandate reporting of external safety reports.
• Lack of documentation of IRB approval of a protocol amendment or action letter that affects more than minimal risk or IRB approval is greater than 90 days after the Network Group’s notification; this includes a Request for Rapid Amendment (RRA) resulting from an action letter indicating temporary suspension of accrual with expedited review permitted.

Lesser IRB deficiencies may include but are not limited to:
• Protocol annual re-approval delayed less than 30 days.
• Delayed re-approval for protocol closed to accrual for which all patients/study participants have completed therapy.

2.8.7.2.2 Informed consent content (ICC)

The audit team verifies that the most recent IRB-approved local informed consent document for at least three protocols (if the number of protocols allows) contains the elements required by federal regulations. In addition, each of the three informed consent documents should be checked to ensure they contain the risks and alternatives listed in the model informed consent document approved by the NCI. If CTSU case(s) are reviewed, at least one local informed consent document should be reviewed for content.

Risks, opt in/opt out Alliance-specific translational research questions and alternatives to study treatment may not be added or deleted from the model informed consent document.

If the site identifies a significant error in risk (e.g. missing risks, or risks erroneously attributed to the drug), the responsible investigator must send an email
to the protocol coordinator listed on the study cover page and the Alliance regulatory group providing written justification for correction of the identified error. The Alliance will determine if a protocol amendment is required to address the issue.

Institutions using the NCI Central Institutional Review Board (CIRB) as their IRB of record must follow the NCI-CIRB policy regarding acceptable and prohibited ICD modifications.

**Critical ICC Deficiency:**

- Any finding identified before or during an audit that is suspected to be fraudulent activity

**Major ICC deficiencies** related to informed consent content (does not represent an all-inclusive list of the major deficiencies that may be found):

- Omissions of one or more risks/side effects as listed in the model informed consent document.
- Omission of one or more revisions to the informed consent per protocol amendment or failure to revise an informed consent in response to an NCI action letter regarding risks that require a change to the informed consent.
- Omission of one or more required informed consent elements required by federal regulations.
- Changes made to the informed consent document not approved by the IRB of record.
- Multiple cumulative effects of minor problems for a given informed consent.

**Lesser ICC Deficiencies:**

- When the CIRB is the IRB of record, failure to have the informed consent document locally implemented within 30 days of notification (posted on the CTSU website)
- IRB approved informed consent document with incorrect version date
2.8.7.2.3 Review of the Delegation of Task Log (if applicable)

The Clinical Investigator (CI) is held responsible for the conduct of a clinical trial and may delegate activities/duties associated with the clinical trial to his/her staff. In such a case, a Delegation of Task Log (DTL) must be maintained and include anyone who contributes significant trial-related duties. This log is generated and maintained by institution and protocol by the CI via the DTL link on the CTSU website.

Auditors will request the DTLs for appropriate protocols and review for implementation and maintenance.

Critical DTL Deficiency:

- Any finding identified before or during an audit that is suspected to be fraudulent activity

Major DTL Deficiency:

- Performing tasks not assigned to individual
- Failure to keep DTL current
- Individual not listed on DTL

2.8.7.2.4 Assessing the IRB, ICC and DTL

The following categories outlined in table 2-1 should be used in assigning a final assessment to the IRB/ICC component of the audit.
Table 2-1. IRB/ICC/DTL audit assessment categories

| Acceptable          | • No deficiencies identified.  
|                     | • Few lesser deficiencies identified.  
|                     | • Any major deficiencies identified during the audit that were addressed and/or corrected prior to the audit for which a written and dated CAPA plan exists, and no further action is required by the Alliance, or NCORP, the institution, or the principal investigator because no similar deficiency has occurred since the CAPA plan was implemented. However, this approach may not be applicable if a deficiency is associated with a safety concern and determined that further action is necessary. In any case, the Alliance will provide the CTMB with a copy of the CAPA plan at the time the final audit report is submitted or by the date follow up is due. |
| Acceptable Needs Follow-up | • Multiple lesser deficiencies identified.  
|                         | • Major deficiencies identified during the audit but not corrected and/or addressed prior to the audit.  |
| Unacceptable          | • A single critical deficiency identified.  
|                         | • Multiple major deficiencies identified.  
|                         | • Excessive number of lesser deficiencies identified.  |

Alliance uses an algorithm as a guideline to determine the final assessment for the IRB/ICC component of an audit. The Alliance tallies the total number of items that are reviewed for a particular IRB/ICC review. IRB records for each protocol that are reviewed and each individual consent reviewed are considered separate items. If a single critical deficiency is identified or if the total number of major deficiencies cited is 20% or greater of the total items that are reviewed for this segment of the audit, the IRB/ICC component of the audit is rated Unacceptable.

While this algorithm is used to assess the majority of IRB/ICC audit ratings, exceptions may be made by the Audit Steering Committee in consultation with the chair of the Audit Committee and the Chief Administrative Officer.

2.8.7.3 Review of accountability of investigational agents and pharmacy operations

An effort will be made to audit a pharmacy on-site at least every other audit, including a re-audit if the deficiencies are related to drug inventory and/or security and the institution has registered patients on one or more studies with IND agents since the previous audit.

Agent accountability and storage procedures described in this section are required under federal regulations and NCI policy for
NCI-supplied study agents (by PMB/CTEP or designated company/Group for DCP and imaging agents). See NCI/CTEP policies under the Agent Management section of the CTEP/PMB website.

An Oral NCI Investigational Agent (Drug) Accountability Record Form (Oral DARF) has been created and all transactions with oral agents must be recorded on this DARF. Agent transactions for formulations other than oral must be recorded on the NCI Investigational Agent (Drug) Accountability Record Form (DARF).

A waiver statement allowing use of electronic DARFs (eDARFs) has not been issued by the NCI and the NCI does not endorse any eDARF pharmacy package. Institutions that choose to use an electronic accountability system must ensure the database is capable of producing a paper printout that is identical to the NCI DARF. Electronic accountability system database limitations are not valid reasons for improper accountability documentation according to NCI policy.

All protocols that use investigational drugs, or commercially available drugs for an investigational purpose when designated by the protocol, must have a specific drug supply for use with that protocol only. This means there may be several supplies of the same drug, each designated for use for only one protocol. Separate NCI DARFs for each study listed by study number must be kept. Multi-agent protocols require a separate NCI DARF for each agent. Each different strength or dosage of a particular agent must also have a separate NCI DARF. For open-label studies, multiple patients may be treated with one drug and each drug receipt and dispensing date is to be recorded on that NCI DARF. DARFs cannot be patient-specific, except in the instance where the drug is being compared with a placebo in double-blind fashion and is supplied per patient by NCI. Refer to the NCI/CTEP Investigator's Handbook for information on drug accountability and the NCI regulations for accountability of investigational agents.

Auditors are required to inspect the drug logs and tour the area where the investigational drugs are stored (on-site audits). The pharmacy (if one participates in the handling of protocol drugs) must also be visited to evaluate storage and security compliance. Arrangements should be made with the staff pharmacist for the audit
team to visit the pharmacy area. If no pharmacy is used, drug-handling procedures in the clinic/office must be audited.

The investigator ordering and/or dispensing agents (or co-signing for others) must be currently registered with PMB, DCTD, NCI. Procedures must be in place in the pharmacy and followed to ensure that the person prescribing the DCTD-agent is an investigator currently registered with PMB and/or the prescription is co-signed by the registered investigator.

2.8.7.3.1 Guidelines for conducting the review

Because of the difficulty categorizing critical, major and lesser deficiencies related to investigational drug accountability and storage, auditors will determine the rating of this component based on the findings of compliance to the required procedures for drug accountability and storage.

The following table lists compliant and non-compliance issues for the review of accountability of investigational agents and pharmacy operations.

<table>
<thead>
<tr>
<th>Compliance</th>
<th>Non-Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Maintain complete, accurate and timely records of agent disposition of all study-supplied agents using NCI Investigational Agent (Drug) Accountability Record Forms (DARFs)</td>
<td>• NCI DARF not maintained or not maintained completely, accurately or on a timely basis</td>
</tr>
<tr>
<td>• Oral study-supplied agents are documented on the Oral DARF</td>
<td>• Oral NCI DARF not maintained for oral study-supplied agents, not maintained completely, accurately or on a timely basis</td>
</tr>
<tr>
<td>• NCI DARFs are utilized to track cancer control/imaging study-supplied agents, or other accountability log captures the same information as NCI DARF</td>
<td>• Lack of a DARF(s) to verify cancer control/imaging study supplied agents are administered to patients/study participants</td>
</tr>
<tr>
<td>• Paper and/or electronic DARFs (eDARFs) contains all required information; paper printout of eDARF is identical to NCI DARF</td>
<td>• Paper and/or electronic DARFs (eDARFs) do not contain all information or are not completed as required; paper printout of eDARF is not identical to the NCI DARF</td>
</tr>
<tr>
<td>• Corrections on DARFs are lined out, initialed and dated with no erasures and whiteouts; corrections on eDARFs are documented</td>
<td>• Erasures or “whiteouts” on paper DARF</td>
</tr>
<tr>
<td>• Agent was dispensed to a registered patient/study participant and documented on the appropriate DARF</td>
<td>• Corrections are not lined out, initialed and dated on paper DARF</td>
</tr>
<tr>
<td>• Appropriate documentation of multi-dose vial agent dispensing to multiple patients/study participants on separate lines of the DARF</td>
<td>• Corrections are not appropriately documented on eDARF in electronic inventory system</td>
</tr>
<tr>
<td>• Patient/study participant returns of oral study-</td>
<td>• Study-supplied agent dispensed to a registered patient/study participant and not recorded on the</td>
</tr>
</tbody>
</table>
supplied agents are documented on the oral DARF

- Patient/study participant returns of non-oral, non-patient-specific agent supplies are not documented on the DARF
- Patient/study participant returns of non-oral, patient-specific agent supplies are documented on the DARF
- [For NCI-sponsored Study] An institution or centralized pharmacy service (Control) may receive NCI-supplied study agent directly from NCI and is permitted to deliver (transport, not re-ship or repackage) NCI-supplied study agent to the institution’s Satellite Dispensing Areas
- [For NCI-sponsored Study] Study Agent has been transferred to an authorized investigator and/or protocol with CTEP approval

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appropriate DARF

- Multiple dose vials not used for more than one patient/study participant and/or doses not documented correctly on separate lines of the DARF
- Dispensing of study-supplied agent to a non-registered patient/study participant recorded on the DARF
- Patient/study participant returns of oral study-supplied study agents not documented on the Oral DARF
- Patient/study participant returns of non-oral, non-patient-specific agent supplies are documented on the DARF
- Patient/study participant returns of non-oral, patient-specific agent supplies are not documented on the DARF
- [For NCI-sponsored Study] NCI-supplied study agents are repackaged and/or reshipped to other investigators, patients, or locations by mail or express carrier
- [For NCI-sponsored Study] Study agent has been transferred to an unauthorized investigator or protocol without CTEP approval

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### Table 2-3. Assessing compliance for DARFs protocol and study agent specific

<table>
<thead>
<tr>
<th>Compliance</th>
<th>Non-Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Only study-supplied agents used to treat patients/study participants and study-supplied agents not used for other purposes</td>
<td>- Substitution of any study-supplied agent, with non-study supplied study agent, including commercial agents</td>
</tr>
<tr>
<td>- Protocol using multiple study-supplied agents have a separate DARF for each agent</td>
<td>- DARF maintained by lot #</td>
</tr>
<tr>
<td>- Separate DARFs are maintained by protocol, study agent, strength, ‘dosage form’ (e.g., oral, injectable), and by ordering investigator</td>
<td>- One DARF used for more than one protocol</td>
</tr>
<tr>
<td>- A separate patient-specific DARF is maintained for each patient/study participant on a patient-specific supply study, as directed by the protocol</td>
<td>- One DARF used for a protocol using multiple study agents</td>
</tr>
<tr>
<td></td>
<td>- One DARF used for multiple agent strengths, dosage forms, or ordering investigators</td>
</tr>
<tr>
<td></td>
<td>- Single DARF used for multiple patients/study participants on study when patient-specific DARF should be maintained</td>
</tr>
<tr>
<td></td>
<td>- Study-supplied agent used for pre-clinical or laboratory studies without written approval by NCI</td>
</tr>
</tbody>
</table>
### Table 2-4. Assessing compliance for satellite records

<table>
<thead>
<tr>
<th>Compliance</th>
<th>Non-Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Satellite Dispensing Area DARF is used at each location where study-supplied agent is received from the Control dispensing area and is stored more than a day</td>
<td>• No satellite DARFs in use when required</td>
</tr>
<tr>
<td>• Satellite Dispensing Area records are available the day of the audit</td>
<td>• Satellite DARFs not available at the time of the audit</td>
</tr>
<tr>
<td>• Satellite Dispensing Area and Control records match and are accurately maintained</td>
<td>• Satellite and Control records do not match or are not accurately maintained</td>
</tr>
<tr>
<td>• Unused and un-dispensed study-supplied agent is documented on Satellite Dispensing Area DARF as returned to Control for disposition (i.e., transfer, return and/or to be locally destroyed)</td>
<td>• Unused and un-dispensed study-supplied agent is not documented as returned to Control dispensing area; Satellite Dispensing Area is inappropriately transferring and/or locally destroying study-supplied agent</td>
</tr>
</tbody>
</table>

### Table 2-5. Assessing compliance for NCI DARFs kept as primary transaction record

<table>
<thead>
<tr>
<th>Compliance</th>
<th>Non-Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Study-supplied agent order receipts/documentation (paper or electronic) are retained and available for review</td>
<td>• Study-supplied agent order receipts/documentation are not retained or not available for review</td>
</tr>
<tr>
<td>• Documentation on Control DARF of study-supplied agent transactions such as agent returns, authorized agent transfers or authorized agent local destruction</td>
<td>• Lack of documentation on Control DARF of study-supplied agent transactions and local destruction</td>
</tr>
<tr>
<td>• Balance on DARF matches physical inventory</td>
<td>• Quantities not accounted for in physical inventory; quantity does not match DARF</td>
</tr>
<tr>
<td>• [For NCI-sponsored Study] Written documentation of NCI authorization for transfer of study-supplied agent between investigators, protocols or institutions or for local destruction of unused/un-dispensed NCI-supplied study agent is maintained (paper or electronic)</td>
<td>• [For NCI-sponsored Study] No written documentation of NCI authorization of transfer or local destruction of NCI-supplied study agent maintained</td>
</tr>
</tbody>
</table>

### Table 2-6. Assessing compliance for return of drug to NCI

<table>
<thead>
<tr>
<th>Compliance</th>
<th>Non-Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Return of unused/un-dispensed NCI-supplied study agent to NCI or locally destroyed with NCI authorization when notified study agent is no longer suitable for clinical use; Return Form or local destruction authorization is maintained</td>
<td>• Unused/un-dispensed NCI-supplied study agent is not returned, not transferred to an appropriate NCI protocol or not destroyed within 90 days of notification from NCI; NCI- supplied study agent is locally destroyed without NCI authorization or not locally destroyed per local institution’s destruction policy</td>
</tr>
<tr>
<td>• Return of unused/un-dispensed NCI-supplied study agent to NCI or locally destroyed with NCI authorization or transferred to another NCI protocol (with NCI approval), when studies are complete or discontinued. Return Form</td>
<td>• Agent returned to PMB that should have been destroyed on-site or agent returned to PMB that</td>
</tr>
<tr>
<td>or local destruction authorization is maintained</td>
<td>was not supplied by PMB</td>
</tr>
<tr>
<td>------------------------------------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td>• NCI-supplied study agent is returned, transferred or locally destroyed within 90 days of study completion, when requested by the NCI, or when patients/study participants are in follow-up and NCI-supplied agent is not being administered</td>
<td>• Failure to maintain Return Form or documentation of authorized local destruction; no written NCI authorization for transfer or local destruction</td>
</tr>
<tr>
<td>• [For Non-NCI sponsored Study] Study agent final disposition of inventory is documented on DARF</td>
<td>• Unused/un-dispensed NCI-supplied study agents not returned, transferred or locally destroyed within 90 days when patients/study participants are in follow-up and no NCI-supplied study agent is being administered</td>
</tr>
<tr>
<td></td>
<td>[For Non-NCI sponsored Study] Study agent final disposition of inventory is not documented on DARF</td>
</tr>
</tbody>
</table>
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**Policy Number:** 2.8  
**Section:** Institutions – 2  
**Date Revised:** January 1, 2018

### Table 2-7. Assessing compliance for agent storage

<table>
<thead>
<tr>
<th>Compliance</th>
<th>Non-Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Each study-supplied agent is stored separately by protocol, strength, ‘dosage form’ (e.g., oral, injectable) and by ordering investigator</td>
<td>• Study-supplied agent is not stored separately by protocol, strength, ‘dosage form’ (e.g., oral, injectable) and/or by ordering investigator</td>
</tr>
<tr>
<td>• Study-supplied agent is stored under proper conditions (i.e., refrigeration, freezer or room temperature) with appropriate documentation and maintenance of temperature monitoring</td>
<td>• Study-supplied agent not stored under proper temperature conditions; temperature monitoring documentation not maintained</td>
</tr>
</tbody>
</table>

### Table 2-8. Assessing compliance for adequate security

<table>
<thead>
<tr>
<th>Compliance</th>
<th>Non-Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Study-supplied agent is stored in a secure area that can be locked</td>
<td>• Study-supplied agent is stored in an unsecured area</td>
</tr>
<tr>
<td>• Storage areas shall be accessible only to authorized individuals; unauthorized individuals are supervised by an authorized individual</td>
<td>• Unauthorized individuals have access to a secure area without supervision</td>
</tr>
</tbody>
</table>

### Table 2-9. Assessing compliance for authorized prescription(s)

<table>
<thead>
<tr>
<th>Compliance</th>
<th>Non-Compliance</th>
</tr>
</thead>
<tbody>
<tr>
<td>• [For NCI sponsored Study] Investigator prescribing or co-signing a prescription for study-supplied agent has an active investigator registration with CTEP and is an authorized prescriber for the protocol</td>
<td>• [For NCI sponsored Study] Investigator prescribing or co-signing an order for study supplied agent does not have an active investigator registration with CTEP or is not an authorized prescriber for the protocol</td>
</tr>
<tr>
<td>• [For NCI sponsored Study] An order for a study- supplied agent is signed or co-signed by an active, authorized registered CTEP investigator prior to study agent dispensing and administration</td>
<td>• [For NCI sponsored Study] An order for a study-supplied agent is not signed or co-signed by an authorized and registered investigator prior to study agent dispensing and administration</td>
</tr>
<tr>
<td>• Procedures are in place in the pharmacy and followed to ensure that the person prescribing or co-signing prescriptions for study-supplied agent is an authorized prescriber</td>
<td>• Pharmacy does not have procedures in place to ensure person prescribing or co-signing prescriptions for study-supplied agent is an authorized prescriber</td>
</tr>
</tbody>
</table>
2.8.7.3.2 Assessing the accountability of investigational agents and pharmacy operations

The following categories in table 2-10 should be used in assigning a final assessment to this component of the on-site audit. CTMB strongly recommends an “on-site” audit be conducted every other 3-year cycle. The main member, NCORP, or the Alliance may conduct an on-site pharmacy inspection.

Table 2-10. Pharmacy audit assessment categories

| Acceptable | • Compliance found for all categories.  
| • Any non-compliant item identified during the audit that was addressed and/or corrected prior to audit for which a written and dated Corrective and Preventive Action (CAPA) plan exists and no further action is required by the Network Group, NCORP Research Base, CTSU, the institution, or the principal investigator. No further action is necessary because no similar non-compliance issues have occurred since the CAPA was implemented. However, this approach may not be applicable if the non-compliance is associated with a safety concern and determined that further action is necessary. |
| Acceptable Needs Follow-up | • Category found non-compliant during the audit, which was not corrected and/or addressed prior to the conduct of the on-site audit. |
| Unacceptable | • A single Critical Non-compliance finding  
| • Multiple non-compliant categories identified.  
| • Inability to track the disposition of NCI-supplied study drug |
| No Assessment Required | • No IND or NCI-supplied study drug is in stock or in use during the audit period.  
| • This designation applies under the following two conditions:  
| • The review of the pharmacy consists of only security, storage and review of pharmacy procedures to ensure investigator has an active PMB registration.  
| • Review of security, storage and pharmacy procedures were found to be compliant. |
| Limited Review Needs Follow-up | • Non-compliance identified under Pharmacy and audit was limited to review of storage, security and/or pharmacy procedures; and CAPA plan or follow-up response is requested. |
2.8.7.4 Review of patient case records

Alliance patient data submitted by the institution to the Statistics and Data Center (SDC) are compared to patient source documents so that the submitted data will be verified against the primary medical record.

Assessment of patient cases should include:

1. Properly signed and dated consent documents (using the original consent documents when possible), including documentation of the consent process

2. All eligibility criteria

3. Correct treatment and treatment sequence

4. Evaluation of disease outcome/tumor response

5. Reporting of adverse events related to treatment

6. General quality of the data submitted, supporting documents uploaded and required/optional specimens submitted

Data that could likely affect every major study endpoint described in the protocol objectives and statistical sections are reviewed using primary documents either by the audit team or as part of central data review.

Auditing Patient Cases for Studies in Medidata RAVE

Targeted Source Data Verification is a system utilized by auditors reviewing patient records to electronically record audit activity directly in iMedidata Rave (Rave) for those studies using Rave to manage patient clinical data.

Source documents should be independently verifiable. Copies of Group study forms generally are not considered to be primary source documents. The use of flow sheets as primary source documentation is strongly discouraged, except for flow sheets that are signed, dated and accepted as part of the official institutional medical record. Primary laboratory reports, progress notes, etc., are considered adequate. Documentation of oral drug administration should be included in the patient's primary record independent of
the flow sheet (e.g., notation in progress notes or photocopy of prescription, as well as documentation in the NCI Drug Accountability Record Form where appropriate).

Per GCP requirements, corrections to paper source documents are to be done by a single line through the error, initials of the person making the corrections, and the date of correction. The correction on CRFs should be supported by the source data. For unusual changes, a brief explanation should be given. If there is conflicting information in the source documents, the PI should indicate in a study note which information was used and why those data were chosen.

Auditor review of source documentation through electronic medical records and electronic imaging is allowable. A staff member must be present to assist with navigating through the system.

Per FDA regulations, the medical record should contain documentation in the case history for each study volunteer that the study consent document was explained to the patient, questions were answered, and informed consent was obtained. This documentation should be included in a progress note, nurse’s note, or elsewhere in the medical record to verify informed consent was obtained.

The CTMB Guidelines section 5.4 allows for missing documentation in the patient case review at the time of the audit to be submitted to the audit team after the audit. The audit team leader will provide the site with a list of unconfirmed items at the exit interview. The missing documentation must be submitted in one submission to the audit team leader within one week following the audit.

A critical deficiency is defined as any finding identified before or during an audit that is suspected to be fraudulent activity.

A major deficiency is defined as a variance from protocol-specified procedures that makes the resulting data questionable.

A lesser deficiency is a deficiency that is judged to not have a significant impact on the outcome or interpretation of the study and is not described above as a major deficiency. An unacceptable frequency/quantity of lesser deficiencies should be treated as a
major deficiency in determining the final assessment of a component.

2.8.7.4.1 Examples of critical, major and lesser deficiencies

**Informed Consent-Critical Deficiencies**
- Any finding identified before or during an audit that is suspected to be fraudulent activity
- Consent form document not signed and dated by the patient/study participant (or parent/legally authorized representative, if applicable)
- Patient/study participant signature cannot be corroborated
- Consent form not protocol specific

**Informed Consent-Major Deficiencies**
- Failure to document the informed consent process with the study participant
- Patient/study participant signs consent form document containing changes not approved by the CIRB/IRB
- Consent form document missing
- Translated consent, short form or other form of translation not available or signed/dated by a non-English speaking patient/study participant
- Consent form not signed by patient prior to study registration/enrollment
- Consent form does not contain all required signatures
- Consent form used was not the most current IRB-approved version at the time of patient registration
- Consent form does not include updates or information required by IRB
- Re-consent not obtained as required
<table>
<thead>
<tr>
<th>Policy Name: Institutional Audits</th>
<th>Policy Number: 2.8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Section: Institutions – 2</td>
<td>Date Revised: January 1, 2018</td>
</tr>
</tbody>
</table>

- Consent of ancillary/advanced imaging studies not executed properly

**Eligibility – Critical Deficiency**
- Any finding identified before or during an audit that is suspected to be fraudulent activity

**Eligibility – Major Deficiencies**
- Review of documentation available at the time of the audit confirms patient/study participant did not meet all eligibility criteria and/or eligibility requirements were not obtained within the timeframe as specified by the protocol
- Documentation missing; unable to confirm eligibility [Exception: Patients deemed ineligible based on laboratory/pathology reports following registration and changes based on central review of material.]

**Treatment – Critical Deficiencies**
- Any finding identified before or during an audit that is suspected to be fraudulent activity
- Incorrect agent/treatment/intervention used

**Treatment – Major Deficiencies**
- Additional agent/treatment/intervention used which is not permitted by protocol
- Dose deviations or incorrect calculations (error greater than +/- 10%)
- Dose modification/treatment/intervention not per protocol; incorrectly calculated
- Treatment/intervention incorrect, not administered correctly, or not adequately
documented

- Timing and sequencing of treatment/intervention not per protocol
- Unjustified delays in treatment/intervention

**Disease Outcome/Response – Critical Deficiency**
- Any finding identified before or during an audit that is suspected to be fraudulent activity

**Disease Outcome/Response – Major Deficiencies**
- Inaccurate documentation of initial sites of involvement
- Tumor measurements/evaluation of status or disease not performed, not reported, or not documented per protocol
- Protocol-directed response criteria not followed
- Claimed response (i.e., partial response, complete response, stable) cannot be verified or auditor could not verify the reported response
- Failure to detect cancer (as in a prevention study) or failure to identify cancer progression

**Adverse Events – Critical Deficiency**
- Any finding identified before or during an audit that is suspected to be fraudulent activity

**Adverse Events – Major Deficiencies**
- Failure to report or delayed reporting of an
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- Adverse event that would require filing an expedited Adverse Event (AE) report or reporting to the Group
- Adverse events not assessed by the investigator in a timely manner (per protocol)
- Grades, types, or dates/duration of serious adverse events inaccurately recorded
- Adverse events cannot be substantiated
- Follow-up studies necessary to assess adverse events not performed
- Recurrent under- or over-reporting of adverse events

#### General Data Management Quality – Critical Deficiency
- Any finding identified before or during an audit that is suspected to be fraudulent activity

#### General Data Management Quality – Major Deficiencies
- Recurrent missing documentation in the patient/study participant records
- Protocol-specified laboratory tests not done, not reported or not documented
- Protocol-specified diagnostic studies including baseline assessments not done, not reported or not documented
- Protocol-specified research/advanced imaging studies not done or submitted appropriately
- Frequent data inaccuracies
- Errors in submitted data
- Delinquent data submission (> 6 months delinquent is considered a major deficiency; a 3-6 month delinquency is considered a lesser deficiency)
2.8.7.4.2 Assessing the findings from patient case records

The following categories in table 2-11 should be used in assigning a final assessment to this component of the audit.

Table 2-11. Patient case records audit assessment categories

<table>
<thead>
<tr>
<th>Acceptable</th>
</tr>
</thead>
<tbody>
<tr>
<td>• No deficiencies identified.</td>
</tr>
<tr>
<td>• Few lesser deficiencies identified and no follow-up is requested</td>
</tr>
<tr>
<td>• Any major deficiency identified during the audit that was addressed and/or corrected prior to the audit for which a written and dated Corrective and Preventive Action (CAPA) plan exists and no further action is required by the Alliance, NCORP Research Base, the institution, or the principal investigator because no further deficiency has occurred since the CAPA plan was implemented. However, this approach may not be applicable if a deficiency is associated with a safety concern and determined that further action is necessary (to be discussed with CTMB liaison). In either case, CTMB must receive a copy of the CAPA at the time the final report is submitted.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Acceptable Needs Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Multiple lesser deficiencies identified.</td>
</tr>
<tr>
<td>• Major deficiencies identified during the audit but not corrected and/or addressed prior to the audit.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Unacceptable</th>
</tr>
</thead>
<tbody>
<tr>
<td>• A single critical deficiency identified.</td>
</tr>
<tr>
<td>• Multiple major deficiencies identified.</td>
</tr>
<tr>
<td>• Multiple lesser deficiencies of a recurring nature found in a majority of the patient cases reviewed.</td>
</tr>
</tbody>
</table>

The Alliance uses an algorithm (table 2-12) as a guideline in assessing the final rating for the patient case review. The number of patients reviewed is multiplied by six (there are six categories in the patient case review; informed consent, eligibility, treatment, disease outcome/response, adverse events, and general data quality). The number 100 is divided by the product. The result is the point value assigned to each lesser deficiency. Each major deficiency is worth double the point value that is assigned to a lesser deficiency. The point value for all major deficiencies and lesser deficiencies should then be added. This sum is then subtracted from 100 in order to determine the final rating score.

<p>| |</p>
<table>
<thead>
<tr>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A final rating score of less than 70 is considered an unacceptable assessment for the patient case review segment of the audit.</td>
</tr>
<tr>
<td>A final rating score of less than 77 is considered unacceptable for a re-audit.</td>
</tr>
</tbody>
</table>
Table 2-12. Final rating for the patient case review

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Line</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients.</td>
<td>1.</td>
</tr>
<tr>
<td>Number of lesser deficiencies.</td>
<td>2.</td>
</tr>
<tr>
<td>Number of major deficiencies.</td>
<td>3.</td>
</tr>
<tr>
<td>Multiply line 1 by 6, which is the number of categories. This is the number of items.</td>
<td>4.</td>
</tr>
<tr>
<td>Divide 100 by line 4.</td>
<td>5.</td>
</tr>
<tr>
<td>This is the point value for each lesser deficiency.</td>
<td></td>
</tr>
<tr>
<td>Multiple line 5 by 2.</td>
<td>6.</td>
</tr>
<tr>
<td>This is the point value for each major deficiency.</td>
<td></td>
</tr>
<tr>
<td>Multiple line 2 by line 5.</td>
<td>7.</td>
</tr>
<tr>
<td>This is the score for lesser deficiencies.</td>
<td></td>
</tr>
<tr>
<td>Multiple line 3 by line 6.</td>
<td>8.</td>
</tr>
<tr>
<td>This is the score for major deficiencies.</td>
<td></td>
</tr>
<tr>
<td>Add lines 7 and 8.</td>
<td>9.</td>
</tr>
<tr>
<td>This is the total deficiency score.</td>
<td></td>
</tr>
<tr>
<td>Subtract line 9 from 100.</td>
<td>10.</td>
</tr>
<tr>
<td>This is the final rating score.</td>
<td></td>
</tr>
</tbody>
</table>

While this algorithm is used to assess the ratings of the majority of patient case review audits, the group chair or designee, in consultation with the Chair of the Audit Committee, Audit Program Director, and Chief Administrative Officer, may make exceptions.

A minimum number of four patient cases are required for utilization of the algorithm.

The audit ratings for audits with less than four patient cases will be assessed on a case-by-case basis.

2.8.7.5 Exit interview

At the conclusion of the visit, the audit team conducts an exit interview. It is expected that the Principal Investigator or designee and designated staff be present at the exit interview. Additional personnel may be present at the discretion of the principal investigator. An appropriate amount of time should be set aside for the audit team to review with the institution the preliminary
findings, items reviewed “off-site”, and recommendations from the audit team.

The exit interview should provide an opportunity for immediate dialogue, feedback, clarification, and most importantly, education.

During this interview, specific problems or questions are discussed. The list of unconfirmed items should be reviewed and provided to the PI and/or lead CRP by the audit team leader. General issues of concern and the major deficiencies should be brought to the attention of the institution staff. It is very important to discuss these issues and to allow the principal investigator to provide clarifications or explanations that could have a direct influence on the final report submitted to the NCI.

2.8.8 Re-audits

A re-audit is mandatory for any component rated as Unacceptable if the institution continues to participate in the Alliance or NCORP Research Base. It is not necessary that the re-audit be conducted on-site. Depending on the nature of the deficiencies that resulted in the Unacceptable rating, the re-audit may be conducted as an off-site review. A re-audit should be done no later than one year after an Unacceptable audit or when sufficient patients have been accrued.

If only the IRB or pharmacy component is rated Unacceptable, an off-site re-audit of that component may be conducted depending on the nature of the deficiencies. Unacceptable pharmacy audits for security or shelf balance issues will be conducted on-site.

If the patient case review component is rated Unacceptable, re-audits must be conducted on-site. In such cases, the IRB/ICC and pharmacy components will also be audited. On a case-by-case basis, complete re-audits (three components) may be conducted after an Unacceptable rating in only the IRB/ICC or pharmacy component.

2.8.9 Audit review

2.8.9.1 Audit evidence of scientific misconduct

The audit team leader must notify the Alliance Chief Administrative Officer, or in his/her absence another designated person within the Office of the Group Chair, immediately if the audit team uncovers any evidence of systematic or apparently
<table>
<thead>
<tr>
<th>Policy Name: Institutional Audits</th>
<th>Policy Number: 2.8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Section: Institutions – 2</td>
<td>Date Revised: January 1, 2018</td>
</tr>
</tbody>
</table>

deliberate submission or intent to submit false data to the Alliance. The Chief Administrative Officer immediately notifies the Group Chair, the Chair of the Audit Committee, and CTMB of this occurrence. See also section 3.4, Individual Scientific Misconduct Policy.

If still on site and it is practical to do so, the audit team will immediately takes steps to preserve the evidence of false data submission and undertake expansion of the audit to gather additional information. A re-audit with an augmented team which may include NCI, Office of Research Integrity (ORI), and FDA representatives will be scheduled by Alliance in cooperation with the appropriate federal agencies.

Any data irregularities identified through quality control procedures or through the audit program that raise any suspicion of intentional misrepresentation of data must be immediately reported to the Alliance Chief Administrative Officer who will report suspicions or findings to the Group Chair, the Chair of the Audit Committee, and the NCI. The CTMB must be notified immediately by telephone of any findings suspicious and/or suggestive of intentional misrepresentation of data and/or disregard for regulatory safeguards for any of the three (IRB/ICC, pharmacy, and patient case) components of an audit. It should be emphasized the irregularity/misrepresentation does not need to be proven and a reasonable level of suspicion suffices for CTEP notification. It is also essential that involved individual(s) and/or institutions follow their own institutional misconduct procedures in these matters.

### 2.8.9.2 Action taken based on audit results

For audits where the findings indicate poor data quality or noncompliance with regulatory requirements, Alliance may take a variety of actions depending on the scope and severity of the problem.

- The PI and institution's staff is advised of the problems encountered during the audit and advised of ways to improve performance.

- If the Alliance is not satisfied that the problems are correctable, it may choose to terminate the membership or affiliate status of the institution.
• Audit reports are reviewed by Alliance audit staff and then forwarded to the principal investigator, outlining the assessment of the audit and any recommendation for action to be taken. If an institution has received an Unacceptable rating in any of the three components (IRB/ICC, pharmacy, patient case), or Acceptable Needs Follow-up (ANFU) with a re-audit requirement, the Audit Committee will also receive an electronic copy of the report.

• The principal investigator and the lead clinical research professional receive final audit reports a maximum of 70 days after an audit takes place. Included with the Final Audit Report is a cover memo that states the audit ratings, explains which deficiencies must be addressed with a written corrective and prevention plan and gives a due date.

• The CAPA plan must include measures for prevention of deficiencies in the future. A response confirming correction of a specific deficiency (e.g., submission of a data form or adverse event report) is insufficient without an overall corrective plan. In many cases, corrective action may entail a review of policies and procedures, additional training of clinical research staff and/or communication with the IRB regarding procedures and timelines. In addition, preventative plans need to be included to ensure the issues do not re-occur and double-check systems are in place.

• If a CAPA plan is determined to be unsatisfactory, and/or if additional information or documentation is required, the Audit Program Director will contact the principal investigator and the lead clinical research professional to obtain an additional response. If the request(s) for an additional response are not answered in a timely fashion, patient registration privileges at the institution may be suspended.

• The CAPA plan is due 15 business days from the date the report was distributed.

• An unacceptable rating in the IRB/ICC, patient case review, or pharmacy sections of the audit is evaluated on a case-by-case basis by the Chief Administrative Officer and/or Group Chair and may also warrant immediate suspension of registration privileges depending upon the evaluation. Registration privileges are reinstated upon receipt of a CAPA plan and
approval of the plan by the Audit Program Director, in consultation with the Chief Administrative Officer.

- If an institution fails to provide an acceptable CAPA plan for one or more audit components rated as Acceptable Needs Follow-Up or Unacceptable within 45 days of when the Final Audit Report was initially distributed, written notice will be provided to the principal investigator that the corrective action is overdue, and a five day working grace period will be granted for the submission of the CAPA plan. If a CAPA plan is not received within this five-day grace period, patient registration privileges may remain suspended. If the institution is an affiliate, patient registration privileges for the main member may also be suspended at this time.

- If the CAPA plan is not submitted within the five-day grace period, it must include a written explanation from the PI that explains the reason for the delay. The suspension of patient registration privileges will not be lifted until an acceptable CAPA plan is submitted and approved by the Audit Program Director, in consultation with the Chief Administrative Officer, and is forwarded and reviewed by the CTMB.

2.8.9.3 Report submission to CTMB

Report of preliminary audit findings must be submitted to the CTMB within one working day of completing the audit. Critical and Major deficiencies should be described. This report is not intended to be a complete or exhaustive list of all deficiencies contained in the final audit report.

The Alliance audit program staff is responsible for submitting all audit reports and related correspondence to the CTMB. If the CTMB has any comments or questions, the audit staff is notified. The audit staff forwards CTMB comments, if appropriate, to the principal investigator and the lead clinical research professional.

2.8.9.4 Changes to the Alliance database subsequent to audit

The Statistics and Data Center staff receive copies of audit reports. The SDC staff is responsible for determining if data changes may be required based on audit findings.
2.9. Continuing Alliance membership

The Alliance Bylaws outline procedurally how Alliance membership status is evaluated. Each institutional member is re-evaluated for performance in Alliance activities by the Membership Committee semi-annually. The Alliance Institutional Performance Evaluation Committee (IPEC) reviews institutional performance semi-annually. All Alliance institutions are subject to periodic audits. The Membership Committee receives reports from the IPEC, the Audit Committee, and other committee reports as needed to evaluate institutional status. Based on the information received from the various sources, the Membership Committee recommends:

- Continue institutional membership
- Suspend patient registration privileges until specific deficiency is corrected
- Change to probationary status
- Mandated change in membership type or expulsion
- Expulsion from the Alliance

Institutions must annually achieve the required number of patient registrations per year (15 for main member networks, and five for affiliates) based on a rolling three-year average.

2.9.1. Main members

Main members that do not fulfill the accrual requirement of 15 patient registrations per year, based on a three-year rolling average, for two consecutive calendar years will be subject to having their membership type changed to an affiliate in the year following the second year that the three-year rolling average was below 15 patient registrations. They would be allowed four months to find a main member with which to affiliate. It is understood that any affiliates of the main member would also need to find a new main member. If the affiliation agreements cannot be executed in this time frame, the main member (and their affiliates/sub affiliates) will be dropped from participation in Alliance.

At the spring Alliance meeting, the main members likely to be affected by this policy will receive a warning letter from the Membership Committee. Prior to the fall Alliance meeting, main members will be informed of the recommendation for a change in membership type and be given the opportunity to appeal at the fall Board of Directors meeting.

The Membership Committee may recommend exceptions to the Board of Directors for approval. If an exception is granted or an appeal is approved, the affected institution will be granted a grace period of one year. If the network does not meet their accrual requirement at the end of the grace period, the network will be subject
to having their membership type changed to an affiliate, without an opportunity to appeal. If the main member and/or their affiliates do not find another main member with which to affiliate by the end of the grace period, their Alliance membership will be terminated, as of January 1st in the year following the grace period.

2.9.2. Affiliates

Affiliates must achieve at least five patient registrations per year based on a three-year rolling average. Affiliates that do not fulfill their accrual requirement for two consecutive calendar years, will be subject to having their Alliance membership terminated, as of January 1st of the year following the three-year period. At the spring Alliance meeting, the affiliate members likely to be affected by this policy will receive a warning letter. Prior to the fall Alliance meeting, main members will be informed of the recommendation for a change in membership type and be given the opportunity to appeal at the fall Board of Directors meeting. The Membership Committee will include a list of at-risk affiliates to the Board of Directors for approval.
2.10 Institutional Network Performance Evaluation

The Alliance membership networks will be evaluated twice yearly coinciding with the Alliance Meetings in three primary areas: quality, timeliness, and group participation. Points will be assigned based on multiple parameters, as shown below. The points will be added to derive an overall score. An overall score can range from -15 to +16.

A network with an overall score below 0 in any evaluation period requires review by the Institutional Performance Evaluation Committee (IPEC) for potential action, including warning or probation. As stated in the Institutional Probation Policy (section 2.11), a network with an overall score of -1 to -5 will receive a warning for substandard performance. The IPEC may recommend probation if a network meets one of the following criteria:

- Two successive evaluation periods with substandard overall scores of -3 or less.
- One evaluation period with substandard overall score of -6 or less.
- Three successive evaluation periods with substandard scores of -2 for timeliness.

2.10.1 Institutional Network Performance Evaluation Scoring System

Below tables 2-20 through 2-22 outline the parameters for each primary area (quality, timeliness, and group participation).

Table 2-20. IPEC scoring for quality

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Values</th>
<th>Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ineligibility (% of patients with eligibility review completed that were deemed ineligible)</td>
<td>&gt;3%</td>
<td>-1</td>
</tr>
<tr>
<td></td>
<td>1-3%</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>&lt;1%</td>
<td>1</td>
</tr>
<tr>
<td>Main member audit (for each component—IRB/ICC, pharmacy, patient case—the most current audit results of acceptable, acceptable needs follow-up [ANFU] or unacceptable will be evaluated)</td>
<td>Unacceptable</td>
<td>-2</td>
</tr>
<tr>
<td></td>
<td>ANFU</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Acceptable</td>
<td>2</td>
</tr>
<tr>
<td>Specimen condition (% of samples intact out of all samples received)</td>
<td>&lt;97%</td>
<td>-1</td>
</tr>
<tr>
<td></td>
<td>97-99%</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>&gt;99%</td>
<td>1</td>
</tr>
</tbody>
</table>

NOTE: This includes all patients evaluated who were accrued by the membership on RAVE trials (patients are not filtered by date of registration to the trial).
Early termination of follow-up (% of patients deemed lost to follow-up, withdrew consent for follow-up or deemed canceled, i.e., protocol treatment not received)  
i.e.: # patients that terminated follow-up early / # patients that were accrued by the membership  
NOTE: This includes all patients accrued by the membership on RAVE trials (patients are not filtered by date of registration to the trial).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Values</th>
<th>Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;3%</td>
<td></td>
<td>-1</td>
</tr>
<tr>
<td>1-3%</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>&lt;1%</td>
<td></td>
<td>1</td>
</tr>
</tbody>
</table>

Table 2-21. IPEC scoring for timeliness

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Values</th>
<th>Points</th>
</tr>
</thead>
</table>
| Data submission (% of eCRFs submitted on time)  
i.e. # forms received on time during report period / total of # forms that were due during the time period plus # forms due before the time period that are still outstanding  
Baseline and treatment forms are given a 15-day grace period after the target date.  
Follow up forms are given a 30-day grace period after the target date.* | <75% | -2 |
| 75%-80% | -1 |
| 80%-85% | 0 |
| 85%-90% | 1 |
| >90%   | 2 |

Response to Queries (% of issued queries that were resolved on time)  
i.e. # query responses received on time during report period / total of # query responses that were due during the time period plus # query responses due before the time period that are still outstanding  
Queries are given a 15 day grace period after the target date.* | <75% | -2 |
| 75%-80% | -1 |
| 80%-85% | 0 |
| 85%-90% | 1 |
| >90%   | 2 |

Specimen Submission (% of baseline samples received on time)  
i.e. # specimens received on time during report period / # specimens that were due during the time period | <75% | -2 |
| 75%-80% | -1 |
| 80%-85% | 0 |
| 85%-90% | 1 |
| >90%   | 2 |

* The grace period for timeliness is based on standards developed by an NCI working group.

Table 2-13. IPEC scoring for group participation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Values</th>
<th>Points</th>
</tr>
</thead>
<tbody>
<tr>
<td>Audit participation by physicians and clinical research professionals (CRPs) in the past two years</td>
<td>No participation</td>
<td>0</td>
</tr>
<tr>
<td>MD or CRP participation</td>
<td>1</td>
<td></td>
</tr>
</tbody>
</table>
2.11 Institutional probation

The Alliance is committed to ensuring that Alliance member institutions meet high quality standards in the conduct of clinical research and the protection of human subjects. Alliance monitors compliance to federal regulations and Alliance guidelines through various mechanisms, including on-site audits and institutional performance evaluations. The criteria for institutional probation set forth below allow Alliance to identify and monitor institutions that have demonstrated substandard performance, with the goal of improving performance at institutions on probation.

2.11.1 Probation based on institutional network performance evaluation

The Institutional Performance Evaluation Committee (IPEC) reviews the performance of main member networks according to the Institutional Network Performance Evaluation Scoring System. The main member networks will be evaluated twice yearly in three primary areas: quality, timeliness, and group participation. Please see the Institutional Network Performance Evaluation Policy (section 2.10) for additional information.

2.11.1.1 Criteria for warnings of substandard institutional network performance

Prior to a recommendation for probationary status, the IPEC may issue warnings to networks with substandard overall scores of -1 to -5 during one evaluation period.

2.11.1.2 Criteria for IPEC recommendation of probation of main member networks

The IPEC may recommend probation to the Membership Committee if a network meets one of the criteria below.

- Two successive evaluation periods with substandard overall scores of -3 or less
- One evaluation period with substandard overall score of -6 or less
- Three successive evaluation periods with substandard scores of -2 for timeliness
2.11.2 Recommendation of probation for an affiliate member

In rare circumstances, IPEC may recommend probation of an affiliate, if it is determined that the substandard overall score for two consecutive evaluation periods is attributable to a particular affiliate.

If the network is underperforming in more than one area, IPEC considers the entire network to be underperforming and recommends probation for the entire network.

2.11.3 Probationary process

The intent of the probationary process is to provide a network the opportunity to improve its Alliance clinical research program, and regain status as an Alliance member in good standing.

The Institutional Performance Evaluation Committee reviews the performance of main members and affiliates using established criteria. The chair of IPEC notifies the principal investigator (PI) in writing of the conclusions of the IPEC.

The IPEC may recommend to the Membership Committee that an institutional network be placed on probation based on substandard performance. Following review and discussion, the Membership Committee votes to determine whether to recommend to the Board of Directors that an institutional network be placed on probation.

The Membership Committee shall communicate the recommendation of probation to the PI of the main member network so evaluated, at a date no later than 30 days prior to the scheduled Board of Directors meeting. The network PI may appeal the recommendation to the Board of Directors before a final decision is rendered. The Board of Directors shall make the final decision and a simple majority shall indicate final approval of recommendations.

After the Board of Directors votes to place a network or individual network sites on probation, the group chair or designee (e.g., chief administrative officer) notifies in writing the main member principal investigator of probationary status, the deficiencies cited, and the penalties associated with probationary status. The group chair or designee copies an institutional official (e.g., dean, executive vice president, cancer center director, hospital director) who is responsible for oversight of the Alliance program.
The principal investigator is required to submit a response and a detailed site improvement plan to the Office of the Group Chair within 30 days of the notice. The Office of the Group Chair may be involved in the development of the site improvement plan in conjunction with the institution. The institutional site improvement plan should address key infrastructural issues contributing to poor performance. The Group Chair or designee may suspend patient registration privileges, if a satisfactory site improvement plan is not received.

During the probationary period, accrual will be closely monitored by the Alliance with increased utilization of quality control procedures at the time of patient registration and timely review of data submission. The member institution may also be assigned a mentor by the Alliance.

Until the probationary status is lifted, the Alliance does not recognize the institution(s) as a member in good standing. Institutions that do not resolve issues responsible for probationary status within one year following an extension of probationary status, and who cannot successfully resolve such issues by changing to another membership level, will be expelled from Alliance. The Membership Committee shall communicate the recommendation to the PI of the main member network so evaluated, at a date no later than 30 days prior to the scheduled Board of Directors meeting. The network PI may appeal the recommendation to the Board of Directors before a final decision is rendered. The Board of Directors shall make the final decision and a simple majority shall indicate final approval of recommendations for lifting of probationary status or one-year extension of probationary status. A two-thirds vote is required for a change in institutional membership level or expulsion of a member from the Alliance. Institutions who are expelled from Alliance may re-apply for membership no sooner than three years after the date of expulsion. See section 8 of the Alliance Bylaws.

All correspondence regarding probationary status of affiliates is addressed to the main member network PI. It is the responsibility of the network PI to inform the individual network institution of probationary status and to work with the institution to develop an appropriate corrective action plan.

The IPEC, Membership Committee, and Board of Directors are scheduled to review probationary status semi-annually. The Audit Committee will report unacceptable audit results to the IPEC and the Membership Committee, as appropriate.
2.11.4 Probation based on unacceptable audits

In compliance with the CTMB Guidelines, if a participating institution (main or affiliate) is deemed unacceptable for the same audit component(s) on two consecutive audits, the institution will be placed on probation. Probationary status may be conferred by the Office of the Group Chair, in conjunction with the Audit Committee. This may occur prior to and separate from the IPEC, Membership Committee, and Board of Directors deliberations. The group chair and chair of the Audit Committee will notify the Membership Committee when probationary action has been taken as a result of unacceptable audits and request an affirmative vote as appropriate. Audit ratings are included in the IPEC criteria for institutional evaluation.

Following a second unacceptable audit for the same audit component, the group chair or designee (e.g., chief administrative officer) notifies in writing the main member principal investigator of probationary status, the deficiencies cited and the penalties associated with probationary status. The group chair or designee copies an institutional official (e.g., dean, executive vice president, cancer center director, hospital director) who is responsible for oversight of the Alliance program.

The principal investigator is required to submit a response and a detailed site improvement plan to the group chair or designee, within 30 days of the notice. The Office of the Group Chair and audit personnel may be involved in the development of the site improvement plan in conjunction with the institution. The institutional site improvement plan should address key infrastructural issues contributing to poor performance. The group chair or designee may suspend patient registration privileges, if a satisfactory site improvement plan is not received.

During the probationary period, accrual will be closely monitored by the Alliance with increased utilization of quality control procedures at the time of patient registration and timely review of data submission. The member institution may also be assigned a mentor by the Alliance.

2.11.4.1 Implications of probationary status

The implications of probationary status for Alliance participation and membership depend on the level of membership and duration of the probationary status. At each anniversary of a network or network institution probation, the IPEC, Membership Committee, and Board of Directors review the status of the cited institution and
votes by majority on the progression of the sanctions according to the following schedule.

**Immediate**

If the network is placed on probation and the institution has a voting seat on the Board of Directors, the PI does not vote at the Board of Directors meetings. If a network institution is placed on probation, the PI retains the privilege to vote at the Board of Directors meetings.

The Alliance operations staff will work closely with the institution to assist in resolving the issues that resulted in a probationary status.

**Year 1 Anniversary**

The network’s accrual privileges are limited according to the following guidelines.

- A main member network is limited to registering 15 patients per calendar year, or 50% of the rolling three-year annual average (up to 100 patient registrations), based on calendar years, whichever is greater. The accrual limitation will be in effect until probation is lifted.
- If the cause for probation is data driven, network accrual privileges may temporarily be limited to 15 patient registrations until the data issues are resolved. Upon resolution of data issues the probationary accrual limitations (15 patient registrations or 50% of annual average whichever is greater) are in effect until probation is officially lifted.
- An affiliate that is placed on probation is not permitted to register more than five patients per year.

**Year 2 Anniversary**

Expulsion. The Board of Directors may vote to terminate membership of the network or affiliate in the Alliance. See section 8 of the Alliance Bylaws regarding conditions for expulsion.
2.12 **Institutional retention of study records**

The following definitions apply in this policy:

- **Research records** are usually maintained by the investigator or research staff, may be separate from the hospital records, and may contain the original signed informed consent form and copies of key protocol parameters.
- **Source documents** include original patient medical records, hospital charts, lab printouts, radiological reports, correspondence, scans, X-rays, patient-completed forms, etc.
- **Flow sheets and case report forms** are created by the Alliance, completed by the institution, and submitted from the participating sites to the Alliance Statistics and Data Center.

The registering institution identified at registration, or, in the case of a transfer, the institution that accepts the responsibility for the patient, is responsible for maintaining and keeping all regulatory and original source documentation.

If the study treatment does not include investigational agents, then the research records (except for signed informed consent) and Alliance case report forms and flow sheets may be discarded after the study has been terminated. The institutional review board that reviewed the study must keep records and minutes of the review per federal guidelines and their own institutional policies.

If the study includes investigational agents, then in addition to the above requirements, records may only be destroyed two years after the New Drug Application (NDA) or Biologic License Application (BLA) has been approved or withdrawn, or the Investigation New Drug (IND) has been withdrawn/closed. The pharmacy at the institution must keep the ordering records for each agent per the federal requirements and the disposition of the investigational agent must be documented in the drug accountability form.

Source documentation, including the informed consent forms, should be retained indefinitely at the registering institution. In many instances, the signed informed consent form is included in the research records and not in the medical records. The Alliance does not collect signed informed consent forms. If the original signed informed consent form is not charted to hospital source documentation and is maintained in the research records, the signed informed consent form must be removed before the research record is destroyed and retained as would be done for source documents.
2.13 Non-member Collaborators

Non-member collaborators (NMCs) are institutions or networks that participate on Clinical Therapy Evaluation Program (CTEP) and Division of Cancer Prevention (DCP) sponsored protocols but are not full member institutions of the Alliance or a participating organization. Most non-member collaborators with the Alliance are international organizations.

In addition to their own country’s regulations, International groups must comply with US federal regulations such as:

- Obtaining Federalwide assurance (FWA) with the Office for Human Research Protections (OHRP); and

- Obtaining State Department Clearance. The Alliance will submit State Department Clearance to the NCI on behalf of the international collaborator.

NCI policy also requires all persons participating in any NCI-sponsored clinical trial to register and renew their registration annually. Registration is accomplished via the NCI Registration and Credential Repository (RCR). Additional details can be found on the NCI/CTEP website.
3 Participants

Individual members of the Alliance fall into three categories: institutional, staff (Alliance operations, statistics and data), and special member.

3.1 Participant Categories

Institutional members belong to an Alliance member institution and are involved with Alliance studies. This category includes the following:

- Principal investigators
- Investigators in all modalities and disciplines
- Pharmacists
- Clinical research professionals and oncology nurses
- Coordinators (e.g., pharmacy, radiation oncology, imaging, surgery, pathology)
- Cytogeneticists
- Administrative staff
- Laboratory researchers
- Fellows in oncology-related disciplines

Alliance staff may be located at an Alliance institution, but are responsible for group functions, including network group management, protocol development, regulatory affairs, statistical support and management of group data. This category includes Alliance operations and program staff as follows:

- Statistics and Data Center
- Office of the Group Chair
- Central Protocol Operations Program
- Cancer Control Program
- American College of Surgeons Clinical Research Program
- Translational Research Program
- Biorepositories

Special members are not located at an Alliance institution but interact with other Alliance participants in group activities. This category includes the following:

- Laboratory personnel handling Alliance samples at a non-Alliance institution
- Imaging/RT personnel evaluating data from Alliance studies
- Active participants relocated to non-Alliance member institutions (e.g., a study chair who has moved to a non-Alliance institution but is continuing to serve as chair)
• Patient advocates
• Investigators who participate in Alliance committees or studies but are not located at Alliance institutions
• Consultants who provide advice to Alliance leadership/committees within their area of expertise but do not actively participate in the research of the research programs of the group
• Data and Safety Monitoring Board (DSMB) members
• Representatives from federal agencies (FDA, NIH, etc)
3.2 Membership and participant registration

3.2.1 Applying for membership and registration

The institutional membership application is available on the Alliance public website (http://www.allianceforclinicaltrialsinoncology.org).

The lead Clinical Research Professional (CRP) or Secondary Lead CRP is responsible for adding and withdrawing all institutional members via CTSU Roster Update Management System (RUMS) or NCORP-SYS.

NCI policy requires all persons participating in any NCI sponsored clinical trial to register and renew their registration annually. Registration is accomplished via the NCI Registration and Credential Repository (RCR). RCR utilizes five person registration types:

- **Investigator (IVR)** — MD, DO, or international equivalent
- **Non-Physician Investigator (NPIVR)** — advanced practice providers (e.g., NP or PA) or graduate level researchers (e.g., PhD)
- **Associate Plus (AP)** — clinical site staff (e.g., RN or CRA) with data entry access to CTSU applications (e.g., RUMS, OPEN, RAVE, TRIAD)
- **Associate (A)** — other clinical site staff involved in the conduct of NCI-sponsored trials
- **Associate Basic (AB)** — individuals (e.g., pharmaceutical company employees) with limited access to NCI-supported systems

All Investigators (IVRs), Non-Physician Investigators (NPIVRs), and Associate Plus (APs) are required to obtain Human Subjects Protocol and Good Clinical Practice (GCP) Training to be in compliance with the NIH. The training provider, course title, completion date, and expiration date, if applicable, and the provider's training certificate must be uploaded in the NCI Required Training subsection of the NCI Biosketch.

All persons applying for Alliance membership must obtain an NCI/CTEP-IAM account, access the RCR system, and complete an annual NCI person registration.

Additional details are available on the CTEP website https://ctep.cancer.gov/investigatorResources/default.htm. Alliance leaders and committee chairs may request special membership for an individual. The request is sent to the Office of the Group Chair.
3.2.2 Alliance person database

The CTSU maintains a database of all Alliance individual members in the Regulatory Support System (RSS).

The institutional principal investigator and the lead CRP are responsible for ensuring that the roster of institutional members is accurate and up-to-date, utilizing the CTSU Roster Update Management System (RUMS) and providing timely notification to Alliance of changes to PIs and lead CRPs.

Alliance staff claim individual members as “persons” in the Alliance roster in and ensures the accuracy of the Alliance person roster.

The Alliance may release portions of the roster to persons who are not Alliance members upon approval by the Alliance group chair or designee. Individuals who wish to request the roster should send a request and justification to the chief administrative officer.
3.3 Traveling on official Alliance business

Alliance members whose travel expenses are paid by an Alliance grant must follow federal guidelines regarding reimbursement of travel expenses. Each institutional grants and contracts office that reimburses travel has its own policy regarding how federal travel funds are to be reimbursed. Please refer to the specific grants and contracts office of the institution that is funding travel expenses for instructions on how to file expense reports.

For information on travel support available from the Alliance, see the Alliance Travel Policy (refer to the Alliance website under the ‘Meetings’ heading). In addition to support for travel to group and committee meetings, the Alliance also provides travel support for the institutional audit program.
3.4 Individual scientific misconduct

The integrity of Alliance data is dependent upon the work of many individuals at all levels of the group. No event is more damaging to the reputation of the clinical research that Alliance and the other network groups perform than the discovery of submission of false or fraudulent data. Inclusion of such data in our analyses may invalidate the scientific conclusions reached. These invalid conclusions may result in the setting of inappropriate medical practice standards consigning large groups of patients to inferior therapy. Moreover, the violation of the trust between the patient and the healthcare team by such an event will erode the relationships required for conduct of clinical trials and harm the public's perception of all medical investigations. As such, evidence of any systematic or intentional attempt to submit false data of any sort to the Alliance will be dealt with in the most rapid and vigorous manner possible. In addition to withdrawing Alliance membership from those affected, and suspending accrual from the institution(s), the Alliance will assist appropriate governmental bodies in the prosecution of the individuals involved.

The Alliance publicizes its policies concerning scientific misconduct in a variety of forums, including the group meeting sessions, the group newsletter, and other means. Specific training sessions in ethics for investigators, clinical research professionals, statisticians, and other personnel are offered.

This training includes instructions on means whereby Alliance members can bring possible instances of scientific misconduct to the attention of those required to investigate it, how to deal with improper data that may have been recorded, and how to correct, if necessary, the scientific record based upon data that are inaccurate.

3.4.1 Receipt of allegations of scientific misconduct

Individuals who have been asked to falsify data or who believe they have knowledge that others are falsifying data must inform the chief administrative officer (CAO) at the Alliance as soon as possible via whatever means (phone, letter, fax, e-mail, personal contact) is practical. The CAO completes a detailed accounting of the notification. If this notification occurs by phone, the CAO asks the party making the call if a witness to the call is desired. The policies of Alliance and NCI require a thorough investigation of any allegation of scientific misconduct while at the same time taking whatever actions are reasonable and proper to preserve the confidentiality of the informant and, until misconduct is proven, to protect the reputation of those accused. Although anonymous calls for the purpose of notification are discouraged since they may lead to less effective resolution of the matter, they are, nevertheless, accepted. This notification does not supersede or replace any notification also required by the institution from which the report originates. Alliance participants should contact the grants and contracts
**Policy Name:** Individual Scientific Misconduct  
**Policy Number:** 3.4  
**Section:** Participants – 3  
**Date Revised:** January 1, 2018

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offices of their institutions to ascertain the correct procedures for reporting such matters at their institution.

### 3.4.2 Processing of allegation within Alliance

Upon receipt of an allegation of scientific misconduct, the CAO immediately brings the matter to the attention of the group chair or, in the absence of the group chair, the group vice chair.

When notification is complete, the group chair, group vice chair, or CAO immediately contacts the Cancer Therapy Evaluation Program (CTEP) Clinical Trials Monitoring Branch to report the incident. Subsequent to this notification, other actions may be required. These may include the immediate suspension of accrual to protocols in the involved institution and further investigation (see below).

### 3.4.3 Investigation of the allegation

In concert with NCI or other agencies, Alliance develops and implements a plan to investigate the allegation. This investigation usually consists of a thorough audit (see section 2.8).

The terms to be used by various committees and officers in connection with the investigation of possible episodes of scientific misconduct have been deliberately chosen to remove any restriction or impediment to whatever action Alliance committees, Executive Committee and Board of Directors may eventually choose to take in a given case. The Alliance may take action against a participant or institution independently whether or not the individual is found guilty in civil or criminal proceedings by others.

The terms used in the audit section of these policies to define institutional performance are used to describe adherence to protocol as well as the quality of data and other submitted materials. In this section we distinguish between erroneous data that result from unintentional mistakes and omissions, and data that are systematically erroneous or untrue.

It is acknowledged that in any process as complex as clinical research occasional errors of many sorts may occur. These may include typographical mistakes, miscalculations of numeric data, omissions of tests, doses, or procedures, delays of treatments, etc. These events when encountered are characterized by the terms used in the audit section and may generate actions concerning the institution as specified elsewhere in these policies.
Falsification of information is to be distinguished from inaccuracies arising from sources noted in the preceding paragraph. Examples include an ineligible patient falsely made eligible, a non-responding patient said to have responded, an abnormal laboratory result made normal, omitted doses of treatment said to have been given, etc. When wrong information is provided systematically, intent to deceive may be inferred. Occasional divergences of opinion among investigators are to be expected in any clinical trial, and data arising from such divergences are to be distinguished from those that are systematic attempts to deceive. When necessary, the Alliance Audit Committee, Institutional Performance Evaluation Committee, Membership Committee, Executive Committee, and Board of Directors render judgment as to whether a given problem represents scientific misconduct and take appropriate actions as defined elsewhere in these policies.

Notwithstanding procedures for revoking membership, halting institutional accrual, or taking other action as defined in these policies or in the Alliance Constitution and Bylaws, the Alliance group chair takes immediate action as defined here when allegations or proof of scientific misconduct occurs within Alliance.

### 3.4.4 Actions to be taken if allegation of scientific misconduct is proved

If false data have been submitted to the Alliance Statistics and Data Center, the data are segregated and reviewed. The SDC staff is responsible for determining what data changes may be required (see also section 2.8).

### 3.4.5 Publication and retractions

If the data have been used in any analyses in preparation of an abstract, the abstract will be revised, if possible, based on a new analysis without the suspect data, or a disclaimer will be offered during the presentation of the revised data. If such data have been used for preparation of a manuscript, the paper will be withdrawn until a new analysis can be conducted. If the manuscript with the false data has been published, the journal will be asked to publish a retraction and re-analysis at the earliest possible time.

It is understood that correction of published information derived from flawed data is of great importance to the public and the scientific community. The Alliance will issue such corrections to relevant journals within 30 days of the time that false data are discovered, or with CTEP consent, whenever a re-analysis can be completed. In addition Alliance has agreed to make its computer data and documentation available to CTEP for analysis when necessary in a national health emergency.
3.4.6 Actions against individuals

An allegation of scientific misconduct may result in immediate action on the part of the group chair to suspend patient registrations by a participant or a member institution. Subsequently, possible actions relevant to institutions occur through usual committee processes described elsewhere in these policies.

Allegations of scientific misconduct by individuals are brought by Alliance staff, the Audit Committee, or others to the Alliance Executive Committee for investigation. Those accused may be asked to appear before the Committee. In such matters, because of the possibility of injury to patients or the public health, time is of the essence. The Executive Committee sets the schedule for the appearance and testimony of the accused. On the basis of the investigation, the Committee may either take no action or may make recommendations to the Alliance Board of Directors. Recommendations to the Board may include severing the membership of the accused, removing the accused from study chairmanship or authorship, censure, or any other action the Executive Committee feels is appropriate.

The accused is provided with the written recommendation of the Executive Committee to the Board. At the meeting of the Board, or in writing prior to the meeting, the accused may offer a rebuttal of the Executive Committee recommendations, but may not offer evidence not previously considered by the Executive Committee. The Board acts on the recommendation of the Executive Committee, accepting it, rejecting it, or changing it, as the Board deems appropriate.

3.4.7 Confidentiality

The action of the Board is final and is a matter of record. It is documented in the minutes of the Board and communicated to the relevant Alliance institution. The deliberations of the Board, the Executive Committee, evidence and audits collected by the committees of the group, and the statements of the accused are held confidential by the Alliance. However, any and all evidence of misconduct is shared with the NCI and/or other appropriate governmental bodies.
3.5 Conflict of interest

3.5.1 Disclosure

3.5.1.1 Introduction

A financial conflict of interest (FCOI) in research means significant financial interest that could directly and significantly affect the design, conduct, analysis or reporting of research. For Alliance for Clinical Trials in Oncology, each person proposed to hold a leadership or staff role that impacts the design, conduct, analysis or reporting of research results must comply with financial disclosure requirements.

The Alliance study chairs/co-chairs, committee chairs, group leaders, Data and Safety Monitoring Board, institutional investigators and Alliance operations staff members are required to disclose financial arrangements >$5,000 per year, as defined in this policy.

FCOI training, review of the Alliance Conflict of Interest (COI) Policy and submission of the Alliance COI form must be completed prior to research participation and at least annually. Updated COI forms are required to be submitted within 30 days of a change in financial arrangements. Study specific COI forms must be submitted until study results are published. Alliance training on the COI Policy and other educational materials will be provided during the annual Alliance Group Meeting.

3.5.1.2 Study chairs/co-chairs

Prior to concept submission, proposed study chairs/co-chairs complete the Alliance Conflict of Interest Disclosure Form (see Alliance website).

The Conflict of Interest Disclosure Form is updated annually or more until the study is published. The Conflict of Interest (COI) Committee, appointed by the group chair, reviews the information on the disclosure form and makes a recommendation to the Alliance Executive Committee concerning possible conflict of interest. The Executive Committee considers this recommendation and, if necessary, additional information, and decides whether a conflict of interest exists that would prevent the individual from serving as a study chair/co-chair. The recommendation of the
Executive Committee is sent to the group chair for action. Disclosure is required for all financial arrangements >$5,000/year.

### 3.5.1.3 Committee chairs/group leaders/institutional investigators/Alliance staff

The Alliance disease, discipline and modality committee chairs and vice chairs complete the Alliance Conflict of Interest Form annually. Institutional principal investigators of Alliance main members, members of the Executive Committee, and staff (defined as all employees) of the Alliance Statistics and Data Center, Alliance Operations/Program Offices also complete the disclosure statement.

This statement is updated annually or more frequently when it is deemed necessary by the individual involved. The COI Committee reviews all disclosure statements. The COI Committee makes recommendations to the Executive Committee concerning possible conflicts of interest. The Executive Committee considers the recommendation(s) and, if necessary, additional information, and decides whether action is needed to manage or prevent a potential conflict of interest. The recommendation of the Executive Committee is sent to the group chair for action. Disclosure is required for all financial arrangements >$5,000/year.

In addition to main member principal investigators, institutional investigators participating in an Alliance study may be required, on a study-specific basis, to disclose financial arrangements as defined in this policy.

### 3.5.1.4 Data and Safety Monitoring Board

The Data and Safety Monitoring Board (DSMB) members submit the Alliance Conflict of Interest Disclosure Form. At the time of each DSMB meeting each member verbally discloses any conflicts pertinent to studies under review and/or recuses themselves from participation in the deliberations of the DSMB. Disclosure is required for financial arrangements >$5,000/year.
3.5.2 Decisions on matters of conflict of interest

The ramifications of the procedures described in this policy preclude preparing guidelines for every possible situation that could give rise to conflict of interest or the perception thereof. For this reason, the COI Committee is broadly charged with using the guidance of the definitions offered below in arriving at a recommendation that is in the best interest of the public, the patients, and the advancement of science. In this activity, the COI Committee members understand that the committee has considerable latitude and flexibility with respect to rendering its decisions. In arriving at a recommendation as to the presence of actual or perceived conflict of interest the COI Committee uses the following definitions as guidance.

3.5.3 Definitions of potential conflict of interest

- **Research Product.** A research product includes a drug, technique, or technology or medical device.

- **Investigator.** Any person who is responsible for the design, conduct, analysis or reporting of research. The investigator must disclose potential conflicts of interest and/or related financial arrangements of any individual with whom the investigator directly shares income (e.g., spouse, children or domestic partner).

- **Conflict of Interest.** A professional, proprietary and/or financial arrangement on the part of the individual, or any individual with whom the individual directly shares income, that may directly and significantly affect the design, conduct, analysis, or reporting of research.

3.5.3.1 Professional interest

The investigator or sponsoring committee chair or vice chair has played a substantial role in the prior development of the product or technology being studied by the Alliance. A professional interest may exist not only where the entity’s products or services are the subject of Alliance-related activity or otherwise under consideration by the Alliance, but also where the entity’s products or services are in competition with those under consideration.

Financial relationships that exist between an individual and a commercial entity in circumstances such as those described below for which compensation is provided in amounts that exceed those listed. A financial arrangement may also exist if the investigator has had a substantial ongoing affiliation with an organization.
having a role in the development or sale of a product or technology including organizations holding patents to or licenses for the development or sale of research products including instances in which the investigator serves as an officer, director, trustee, general partner, employee, or on a scientific advisory board or in a similar capacity for such an organization. Such organizations also include those with which the investigator is negotiating for or has an arrangement concerning prospective employment or affiliation, or those from which the investigator receives or expects to receive compensation exceeding $5,000 annually for honoraria, consultative services, paid authorship or from their fiscal intermediaries such as medical services or continuing medical education companies. All non-government or non-academic travel reimbursement from a for-profit entity must be disclosed including the purpose, sponsor/organizer, destination, duration and additional information as needed. Conflict of interest may also exist if an individual receives $100,000 or more over a three-year period for research funding that is not designated for a particular study or contracted product through their employing institution (i.e., “unrestricted educational grants”). The significance of the conflict will depend, to some degree, on whether reimbursement for professional activities involves compensation limited to that normally required to support the scientific process, or is substantially larger, leading to actual or potential personal financial gain to the investigators or any individuals with whom they directly share income.

An investigator with financial relationships >$25,000/year in a privately held business, equity interest in a publicly traded company sponsor that exceeds $50,000/year, or ≥5% ownership interest (including common stock) in either a privately held or publicly traded business, will generally be prohibited from assuming chairmanship of a study. An investigator with financial relationships >$25,000/year or equity interest in a publicly traded company sponsor that exceeds $50,000/year, or ≥5% ownership interest (including common stock) in either a privately held or publicly traded business, who also serves on the Executive Committee must recuse themselves from participation in the deliberations of the Executive Committee where a conflict or appearance of conflict of interest may exist.
3.5.3.2 Proprietary interest

The investigator has financial interest in the research product being evaluated because the investigator or any individuals with whom they directly share income has a material interest in the product or technology that may result in financial gain, e.g., the investigator is receiving compensation that could be affected by study outcome such as compensation that is explicitly greater for a favorable result or the investigator is receiving annual royalties or other compensation at a value exceeding $5,000/year following the commercial sale of the product or technology. Such royalties may be in the form of personal compensation to the investigator or may be used in support of the investigator's research.

The investigator has financial interest in the research product being evaluated because the investigator or any individuals with whom they directly share income has an equity interest (including common stock) exceeding $5,000/year, or ≥5% ownership interests (including stock options) in a start-up company, the stock of which is not publicly traded, or options exceeding $5,000/year in a commercial enterprise that will benefit from the sale of the product or technology.

3.5.3.3 Miscellaneous and multiple financial interests

There may be other instances in which an investigator or any individuals with whom they directly share income has an affiliation or relationship such that objective impartiality could be questioned. In such instances, the investigator should disclose the nature and extent of such affiliation or relationship on the disclosure form.

Alliance leaders may have individual financial interests related to industry partnerships or other affiliations that do not exceed the threshold of $25,000. Multiple disclosures of >$5,000 are subject to review by the Alliance COI Committee. The Committee may request a management plan including oversight by co-leaders.

Committee chairs with financial interests in products, actively under investigation or proposed in committee sponsored studies, may be required to publicly disclose potential conflicts and/or recuse themselves from relevant discussions.
Committee chairs with financial interests exceeding thresholds defined in this policy may be subject to management plans and restrictions, per section 3.5.4 below.

3.5.4 Management plan for conflicts of interest

Prior to concept submission, study activation, as financial arrangement change and at least annually, all members of the study leadership team are required to complete a Conflict of Interest Form as described above. When the Alliance staff identifies potential conflicts of interest, these issues are referred to the Conflict of Interest Committee for review. If the COI Committee determines that a conflict of interest exists, then the management plan for study leaders with financial arrangements between $5,000/year and $25,000/year or equity interest in a publicly traded company sponsor ≤$50,000/year, outlined below will be enacted to ensure accurate and unbiased data collection and reporting for studies undertaken by the Alliance.

- The study chair and the study statistician jointly oversee all trials. No aggregate outcome data are shared with the study chair until it is released from the DSMB oversight for DSMB monitored studies or deemed ready for sharing based on the trial statistician for non-DSMB monitored studies. For all phase III trials, the study chair does not have access to the raw data except for what is provided in the eligibility and case evaluation summary. When a potential conflict of interest exists for the study chair, the study co-chair or their designees will be required to take a significant role in reviewing data and preparing study results for publication or presentation. These steps will be taken in addition to the existing policy of distributing drafts of all manuscripts to the relevant disease committee members and the Alliance principal investigators for review prior to external submission.

- Independent review by NCI: CTEP will be informed of the COI Committee determination that the potential for a conflict of interest exists on the part of the Study Chair.

- Independent review by a Data and Safety Monitoring Board will continue to be provided for all phase 3 trials. In cases where a potential conflict of interest exists for the study chair, copies of relevant COI findings will be forwarded to the DSMB during their review of the relevant study. A representative from CTEP participates in DSMB meetings and will have access to this information at that time.
• The study statistician, the study co-chair, or his or her designee, and the professional staff of the Alliance Statistical Center will undertake management of data independent of the study chair.

• Financial conflict disclosures of institutional investigators are subject to institutional conflict of interest policies. The Alliance may request a mitigation plan from investigators exceeding thresholds, including documented institutional management plans in compliance with institutional requirements. Independent review of studies by network group leadership beyond the sponsoring committee will be undertaken.

In the event of conflicts exceeding the $25,000 annual threshold or equity interest in a publicly traded company sponsor of $50,000 annual threshold, or ≥5% ownership interest (including common stock), or direct employment with an industry partner, the following policies will be enacted.

• The individual in question may not serve as study chair or co-chair or serve in an oversight capacity as chair of the committee sponsoring the trial if such a conflict is deemed to exist while the study is actively accruing patients and until the primary study analysis has been completed. In this circumstance, the group chair will appoint a new study chair without such a conflict, or when a conflict exists for the committee chair, then the committee vice-chair or their designee will assume responsibility for study oversight.

• The new study chair and the study statistician will assume primary responsibility for data management, analysis, and presentation and publication of study results.

• The individual with a conflict of interest may retain rights of authorship on publications derived from the study in accordance with the requirements for disclosure of conflicts of interest established by the relevant publishing authorities. Any individuals with a significant conflict of interest such that they are ineligible for a study chair or co-chair role cannot serve as either first, corresponding, or senior (last) author of an Alliance publication. When a conflict exists for the committee chair or vice-chair the committee leader may not serve as either first, corresponding or senior author. If all of the key individuals of a study show a significant conflict of interest such that they are ineligible, then the disclosures are sent to COI committee for review.
• The Alliance may disapprove study participation of institutional investigators exceeding maximum thresholds, upon review of the institutional plan to mitigate bias.

3.5.5 Review of disclosure statements

The Conflict of Interest Committee meets no less frequently than once per year and reviews disclosure statements and makes recommendations concerning possible conflicts of interest to the Alliance Executive Committee.

3.5.6 Actions on conflict of interest

The Executive Committee recommends to the group chair actions to be taken with respect to significant conflict of interest.

3.5.7 Penalties for failure to observe conflict of interest policies

Lack of compliance with these policies is referred to the Alliance Executive Committee. The Executive Committee will conduct and complete a retrospective review within 120 days of identified noncompliance and document findings. The Executive Committee recommends whatever action it deems appropriate to the Alliance Board of Directors. The Board of Directors receives this recommendation and takes whatever action it deems appropriate, accepting the recommendation or applying a greater or lesser penalty than that recommended. Failure to submit conflict of interest forms or to comply with COI management plans by individuals subject to the COI policy may result in suspension or termination of Alliance membership privileges including study or committee chairpersonship. Public disclosure

3.5.8 Public disclosure

Financial conflicts of interest must be disclosed in each public presentation of research results. Financial conflicts of interest must be disclosed during Alliance committee meetings, including study development discussions. In addition, the Alliance will make FCOI information publicly available within five days of a written request.

3.5.9 Record keeping

The Alliance Staff maintains records of all financial disclosures and all actions taken by the Alliance with respect to each conflicting interest for a minimum of three years after the grant period within which the forms were collected has ended. Summary recommendations of the Conflict of Interest
Committee are reported to the Executive Committee and become part of the minutes of that committee.

### 3.5.10 Reporting Financial Conflicts of Interest (FCOI)

The Alliance reports Financial Conflicts of Interest (FCOI) that could directly and significantly affect the design, conduct or reporting of NIH funded research.

The Alliance submits COI disclosures to the Cancer Therapy Evaluation Program (CTEP), as a part of the CIRB Submission. A management plan is provided as appropriate.

The Alliance provides an FCOI report to the awardee Institution receiving Alliance grants (e.g., Brigham and Women’s Hospital) according to the requirements of the Institution.

### 3.5.11 Alliance Conflict of Interest Committee

The Alliance Conflict of Interest Committee is a volunteer committee comprised of Alliance investigators. The committee reviews financial conflict of interest disclosures related to trials supported by the Alliance and Alliance for Clinical Trials in Oncology Foundation.
4 Committees

4.1 Committees and their function in Alliance

The Alliance has scientific (disease, modality, and discipline) and administrative committees. These committees are responsible for the scientific, administrative oversight and quality assurance activities of the Alliance.

4.1.1 Disease committees

The Alliance disease committees are responsible for developing and conducting the scientific agenda of the Alliance. They collaborate closely with the modality and discipline committees and many of the Alliance studies are multimodality studies that address more than one research hypothesis.

4.1.2 Discipline committees

The Alliance discipline committees, working in conjunction with the disease committees or on their own, are responsible for studies that focus on new methodologies for treating cancer or minimizing the burden of cancer for individuals and their family members. These committees also develop studies addressing the fundamental biology of cancer, cancer risk assessment and prevention.

4.1.3 Modality committees

The Alliance modality committees develop educational programs and/or provide quality control services and serve as a scientific resource for other committees.

4.1.4 Administrative committees

The Alliance administrative committees are responsible for the administrative and quality assurance activities of the Alliance.
4.2 How to form a committee

The proposal to form a new scientific committee is brought before the Executive Committee for review and approval. The approval and formation of a new committee are also brought to the attention of the chair of the Constitution Committee, since changes to the Alliance Constitution or Bylaws may be required.

The group chair may form administrative, ad hoc committees, and working groups; the Executive Committee must approve these committees and the chairs of newly formed committees.
4.3 **Committee membership**

The group chair, with the approval of the Executive Committee, names the committee chair and the committee chair selects the committee members. Members are encouraged to bring their ideas to the committee chair for consideration and let the committee chair know of their interest in being on the committee. Committee membership is rotated at appropriate intervals.
4.4 Roles and responsibilities in committees

4.4.1 Committee chair nomination and approval

Committee chairs are either proposed by the group chair or, for those committees within Alliance programs, are nominated by the appropriate program director. All chair appointments are approved by the Alliance Executive Committee. The chair is chosen and their performance is evaluated on the basis of the leadership they can provide to the area of committee responsibility.

4.4.2 Committee chair responsibilities

It is the responsibility of the committee chair to coordinate the activities of the committee and to ensure that the work of the committee is performed in a timely manner.

4.4.2.1 Administration

Committee membership: The committee chair names the members of the committee. The number of members may not exceed the number designated by the group chair. The committee chair is responsible for rotating members off of the committee and adding new ones as needed.

The committee chair nominates the vice chair of the committee to the group chair for the group chair’s review and approval. The committee chair also nominates subcommittee chairs, if applicable, to the group chair for the group chair’s review and approval.

Committee liaisons: The committee chair names liaisons to the committee from other committees after discussion with the other committee chair.

Group and committee meetings: The committee chair or designee prepares agendas for committee and group meetings.

The committee chair also identifies invitees to the meetings and requests support for travel, as appropriate.

The committee chair may schedule conference calls as often as needed to accomplish the work of the committee.
Conflict of interest: The committee chair, vice chair, study chair and study co-chair (of studies that have not been published, current and pending studies) are required to complete a conflict of interest disclosure form at least annually (see Alliance Policies and Procedures, section 3.5 Conflict of Interest).

Scientific misconduct: Each participant in the Alliance is expected to review and comply with the section on individual scientific misconduct in the Alliance Policies and Procedures (see section 3.4 Individual Scientific Misconduct).

The committee chair and all investigators are expected to comply with federal guidelines regarding human subjects training requirements.

Annual progress reports: The committee chair prepares an annual progress report for the committee for inclusion in the annual grant progress report that the Alliance must submit to NCI.

Competing renewal report: The committee chair prepares a committee report whenever the Alliance submits a competing renewal application.

4.4.2.2 Protocol development and management

The committee chair assigns study chairs and evaluates study chair performance on an ongoing basis.

The committee chair supervises the protocol process from concept development through publication of results. This includes overseeing the development and review of concepts, submitting concepts to the Study Concept Review Committee, providing input throughout the protocol development process, and reviewing publications including interim agenda reports, abstracts and manuscripts. Committee chairs are responsible for mentoring study chairs and guiding them through the protocol development process, including forms development. The committee chair acts as a mediator if other members of the study team cannot reach resolution on significant issues that arise during the life cycle of a protocol. Committee chairs also participate in the development of protocol amendments as required.
The committee chair reviews study accrual on an ongoing basis and consults with the study team to develop appropriate action plans for studies that are accruing at a slower pace than anticipated.

The committee chair attends DSMB meetings for studies within the committee.

The committee chair participates in study team conference calls as appropriate.

The committee chairs may be contacted by investigators, oncology nurses and clinical research associates when the study chair is unavailable, with questions pertaining to a specific study (e.g., clarification of eligibility, treatment issues). Nobody, including the committee chair, may grant waivers of eligibility criteria.

The committee chair regularly communicates with the data managers. The committee chair may be called on to answer protocol questions in the absence of the study chair.

The committee chair regularly communicates with the statisticians responsible for the committee regarding protocol development, monitoring of ongoing studies and analysis/publication of results.

### 4.4.2.3 Publications

The committee chair, along with the committee’s primary statistician, works with study chairs to complete manuscripts in a timely manner. If the study chair is not able to write a manuscript in a timely manner, then the committee chair discusses with the group chair reassignment of that study to another individual who will be able to write the manuscript.

The committee chair reviews the committee’s statistical study summaries before they are distributed to the Alliance.

The committee chair reviews the committee’s abstracts/manuscripts/presentations prior to public distribution.

The committee chair works with his/her counterpart in other network groups to ensure that intergroup studies have a representative from each group that participated in the study.
4.4.2.4 Intergroup collaborations

If appropriate, the committee chair discusses collaboration with his/her counterpart in other national groups.

The committee chair may also explore collaborations with international groups in conjunction with the CPOP office. International groups must comply with the federal regulations of the United States in addition to their own country’s regulations.

4.4.2.5 Finances

Travel to Alliance meetings: Committees that do not have a separate grant have travel funds available in the Office of the Group Chair from various grants and other sources to support travel to the committee meetings. Travel expense reimbursement policies and frequently asked questions may be found on the Alliance website in the “Meetings” section.

Funding to support research projects: The committee chair works with the committee members and interested participants in the Alliance demonstrating interest in applying for additional funding. If the project appears feasible, the committee chair asks the person who is responsible for the project to discuss the project with the executive officer and the primary statistician so that appropriate budgets, supporting the efforts of the Alliance operations offices, will be included in the application.

If funding is requested to support a research project, the group chair, appropriate committee chair(s), executive officer, and group statistician are copied on correspondence regarding this project.

Details concerning the proposed funding are included with the concept when it is submitted to the SCRC for review. Some concepts are not approved if no new funds are brought in to finance them. Information about the potential sources of support—federal or non-federal—should appear on the cover sheet that accompanies the concept.

The proposal is submitted to the CPOP office for review of the scientific plan and budgetary requests prior to submitting an application to the NCI or other granting agencies. If the proposal is submitted to a non-federal source, the Alliance Foundation reviews
All negotiations with industry collaborators are handled by the CPOP office staff, not by the investigator or committee chair who proposed the project. The Alliance and the party arrange the details of the drug/device provision directly to group members.

4.4.3 Committee vice chairs

The committee vice chairs assists the committee chair in carrying out the responsibilities of the committee and assumes responsibility for the committee when the committee chair is absent. The committee chair nominates candidates for committee vice chairs, who are approved by the Executive Committee.

4.4.4 Subcommittee chairs/cadre leaders

The cadre leader is appointed by the committee chair with the approval of the group chair. The cadre leader coordinates the activities of a subcommittee.

4.4.5 Committee members

Committee members, based on their expertise and interest in that particular area, are appointed by the committee chair with input from the vice chairs and from modality/discipline committee leaders. Principal investigators may nominate Alliance members to the committee for consideration by the committee chair, but only the committee chair appoints the committee members.

Patient advocates are assigned to other committees (besides the Patient Advocate Committee) and advise committees on various aspects of clinical research, providing the patient perspective.
4.5 Electing Executive Committee members

The group chair appoints a nominating committee consisting of at least three individuals, no more than one of whom may be from any single member institution. The nominating committee, after consultation with the chairs of the appropriate modality committee and the cancer control committee, proposes a candidate(s) for vacant positions on the Executive Committee. In addition, individual institutional members may make nominations at the time of the election by the Board of Directors. Each position on the Executive Committee is filled by a separate election. Each election is conducted by closed ballot. In the event of a plurality, only the top two candidates are entered into a runoff election.

Each elected representative to the Executive Committee serves a three-year term and may only be elected for three consecutive terms, but is eligible for re-election following a term out of office. The terms of office of the elected members of the Executive Committee overlap to provide continuity of committee activities. See the Alliance Constitution for additional details.
5 Meetings

The purpose of meetings of the Alliance for Clinical Trials in Oncology is to provide a forum to plan, conduct, and share the results of clinical trials research with the membership of Alliance. Meetings are a necessary and vital communication function of the Alliance. Continuing medical education credits may be offered for physicians, clinical research professionals, oncology nurses, and pharmacists, when appropriate.

5.1 Group meetings

Group meetings are held on a biannual basis and are open to all Alliance members. Most disease, modality, discipline, and administrative committees meet during the three- or four-day meeting. The disease, modality, and discipline meetings are open to all attendees. Disease, modality, and discipline chairs have the option to have a closed session for their committee members in addition to the open session. This request must be communicated to the meetings manager before a meeting schedule is published. All administrative committee meetings are closed sessions and only open to committee members or invited guests. In addition, several committees may sponsor workshops and educational forums as time and space allows. The group chair sets the agenda for the plenary session.

Outside speakers may be invited to address committees, but are subject to approval by the group chair. Funding may not be available and should be verified with the Office of the Group Chair prior to committing financial support for travel and honoraria.

Alliance members receive information regarding group meetings in newsletters and on the Alliance website (http://www.allianceforclinicaltrialsinoncology.org). The travel policies and reimbursement forms are included on the member’s side of the Alliance website under the ‘Member Services’ tab and the heading ‘Meetings’.

5.1.1 Attendance

All members who plan to attend a group meeting should complete an online registration form. A registration fee is required for non-member attendees. Attendance records will be maintained at the Office of the Group Chair.

Attendees are welcome to go to any open session at the meeting. Attendance at closed sessions is only open to committee members or with approval from the committee chair.

5.1.2 Travel funding for group meetings

Travel funding for attendees is based on the committee rosters. A subset of committee members is selected by the committee chairs to receive funded
travel spots and necessary staff are also funded to attend. Travel expenses for all other meeting participants are the responsibility of the participant or their institution. Non-members that have full/partial registration fee covered by the Alliance (e.g., guest speakers, other select attendees) will be given a code for use in the online registration.

5.1.3 Study accrual reports and publications

Reports summarizing the progress of active studies are generated by the Alliance Statistics and Data Center and distributed at the group meeting (at least annually). The summary also includes a listing of published manuscripts and abstracts. The primary purpose of these reports is to inform Alliance meeting attendees as well as the National Cancer Institute of the current status of Alliance research.
5.2 **Identification of funded travelers and expense reports**

All committee members on a roster are eligible to be funded to attend group meetings. It will be the responsibility of the committee chairs to identify a subset of their committee roster to be funded for group meetings.

5.2.1 **Committee member funding and roster updates**

Committee chairs must submit the “Committee Funding Form” to the Office of the Group Chair at least three months prior to each meeting. The form includes the current committee roster as well as past funding and attendance information, where applicable. Committee chairs must submit the form with roster updates and indicate their funded travelers. Committee chairs with supplemental funding from the Alliance for Clinical Trials in Oncology Foundation should submit the names of travelers utilizing these funds at the same time.

If the form is not received by the deadline, the list of funded travelers will default to the last funded list for that committee. There will be no exceptions or changes after this date.

5.2.2 **Travel funding notification**

Travelers who are funded by the Alliance or Alliance for Clinical Trials in Oncology Foundation will be notified via invitation email. The invitation email will include information on arranging travel, travel policy, and reimbursement of allowable expenses.

5.2.3 **Expense reports**

Funded travelers must submit for reimbursement of out of pocket expenses to the Alliance within 180 days of travel. All expenses must comply with the Alliance travel policy. Expenses that do not comply with policy cannot be reimbursed and will be removed from the expense report or adjusted as needed to comply. Given the volume of expense reports, it is not always possible to notify the traveler when this occurs. The attendee will be contacted if the expense report is not completed correctly or if receipts are missing or inadequate.
5.3 Continuing Education (CE) Credit

The Alliance for Clinical Trials in Oncology offers a variety of Continuing Education (CE) credit opportunities. Through sessions occurring at the Alliance’s semi-annual Group Meetings, members can receive Society of Clinical Research Associates (SOCRA) and/or Nursing CE credit. Sessions eligible for SOCRA credits may also be eligible for CE credits from the Association of Clinical Research Professionals (ACRP). As a research based organization, the Alliance prioritizes these CE opportunities for the membership and values members’ need to develop increased skillsets to further aid in clinical research.

5.3.1 Continuing Education (CE) Requirements

Members interested in receiving CE credits must fill out the registration application and indicate credit requests prior to the Group Meeting.

CE credits are tracked through the Group Meeting registration system. CE credits and certificates may not be issued if they are not designated at the time of registration.

For sessions to count for CE credits members must ensure timely attendance at these sessions. It is the responsibility of the individual member to ensure that attendance is recorded at each session for CE credit.

In order to receive both SOCRA and Nursing credit, an additional required post Group Meeting feedback survey must be completed within 30 days of the survey’s release.

All individuals interested in CE credit must complete all required materials and/or inquire about their CE credit within 6 months of all CE credit events.

5.3.2 Continuing Education (CE) Credit Certificates

Once all the requirements for CE credits are completed, the Alliance provides members with a completion certificate stating the amount of hours/credits individuals have received.

All credit certificates will be sent out after the 30-day deadline and once participants have completed the required survey.

While the Alliance will provide a certificate citing an individual’s total CE hours/credits, it is the responsibility of the member to keep track of
### Policy Name: Continuing Education (CE) Credit

<table>
<thead>
<tr>
<th>Policy Number: 5.3</th>
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</thead>
<tbody>
<tr>
<td><strong>Section:</strong> Meetings – 5</td>
</tr>
<tr>
<td><strong>Date Revised:</strong> January 1, 2018</td>
</tr>
</tbody>
</table>

...individual hours/credits and the submission process necessary for each desired CE program.
6 Study protocol

This section of the Policies and Procedures describes Alliance clinical trial characteristics and conduct, including definitions of study types, study team roles, development of a study protocol, and policies relating to study conduct.

6.1 Study types

Each Alliance study is characterized either as a “treatment” study or as a “non-treatment” study.

6.1.1 Treatment studies

Treatment refers to therapy for diagnosed cancer including chemotherapy, surgery, radiotherapy, or other therapy, including adjuvant therapy, as long as it is directed against the cancer.

6.1.2 Non-treatment studies

All other studies are classified as non-treatment, even those for which there is therapy for some secondary condition. Non-treatment studies can stand alone or can be a companion to one or more treatment studies.

6.1.2.1 Companion studies

A companion study is conducted in conjunction with one or more treatment or other intervention studies. Companion studies may investigate pharmacology, tumor biology, quality of life, symptom management, economic outcomes, or other areas of interest to the group.

A companion study may be embedded within another study to reduce administrative and IRB work for participating institutions, decrease the number of consent forms a trial participant must sign, or facilitate translational research. In order to receive a separate study number, the study component should be an objective (or more than one objective) of the main trial, as listed in the protocol document. Companion studies with separate study numbers do not necessarily have to be published at the same time as the parent study, and may be published as a distinct manuscript. The component should also have a separate study chair who is not the parent study chair listed on the protocol cover page.
The Alliance Executive Committee has determined that an embedded companion study may be assigned 0.25 membership accrual credits. Companion membership accrual credits will be separated from any accrual-based NCI credits or payment amounts, as described on study funding sheets.
6.2 Study participation

Unless otherwise indicated, Alliance studies are open to all members of the group. In accordance with U.S. Department of Health and Human Services (HHS) policy, member institutions must receive IRB approval prior to registering trial participants on an Alliance study. Some studies may require limited access or establish individual credentialing requirements (see section 7).

6.2.1 Limited access studies

Limited access studies restrict trial participant registration to a specific list of institutions indicated on the protocol cover page. Affiliates or networked institutions may not participate unless specifically stated on the protocol cover page. Main member institution participation does not guarantee affiliate institution participation. An affiliate institution may participate, if listed on the protocol cover page, regardless of whether its corresponding main member institution also participates. The study chair, in consultation with the committee chair, determines the list of limited access institutions.

As per NCI requirements, limited access studies may not include members outside of the Lead Participating Organization. Permission for the addition of institutions outside of the Lead Participating Organization to limited access studies must be obtained from the NCI.

6.2.2 Credentialing

Studies may require credentialing, an authorization before investigators and/or institutions can participate. Credentialing is often conducted at the level of an individual investigator, e.g., a surgeon is credentialed to perform a particular surgical procedure. Institutions may also need to be authorized to participate in a particular study, e.g., an approved transplant institution. Authorizations may be study-specific, and may require fulfillment of additional regulatory requirements. Requirements for credentialing and/or authorization are included within the protocol document.

6.2.3 Non-Alliance members

Members of other network groups may participate in certain Alliance studies via the CTSU and the Oncology Patient Enrollment Network (OPEN). Requirements for submission of study data and materials are the same as for Alliance members.
6.3 Study team roles and responsibilities

6.3.1 Study chair

The study chair is responsible for proposing the research idea to, and obtaining approval from, the sponsoring committee chair. The study chair works with the committee chair, committee statisticians, appropriate committee members, committee liaisons, and other study team members to refine the concept and, upon review by the Alliance Study Concept Review Committee (SCRC) and approval by the Cancer Therapy Evaluation Program (CTEP) or the Division of Cancer Prevention (DCP), to develop the trial. Trial development includes writing and revising sections of the protocol, participating in conference calls with the study team and CTEP or DCP, and working with statisticians and the data management staff to define the required data elements that must be captured on the case report forms.

While the trial is active, the study chair responds to requests for clarification of protocol details, participates in the development of trial amendments, and, when appropriate, participates in case reviews. For phase 1 trials, the study chair is required to convene regularly scheduled conference calls with the primary statistician, representatives from each participating institution, and other staff as appropriate to evaluate toxicities encountered and to make decisions concerning dose escalation, modification of cohort size, etc.

Upon completion of the primary endpoint, and in conjunction with the primary statistician, the study chair is responsible for ensuring that the results of the study are published or reported to the scientific community in a timely manner.

6.3.1.1 Moving study chair to a non-Alliance institution

If the study chair moves to a non-Alliance institution, the committee chair appoints an Alliance-based study co-chair, if one has not already been named for the study. The study chair may continue to serve in the full capacity of study chair with the agreement of the appropriate committee chair and if no conflicts of interest have arisen because of the move of the study chair.

6.3.1.2 Replacing study chair

Study chairs will have their performance carefully evaluated and will be replaced if performance is not satisfactory. If a study chair is forced to relinquish responsibility for a study, the group chair (or
6.3.2 Study co-chair

It is expected that study co-chairs contribute in a meaningful way to the study conduct, for example, by answering questions from institutions related to their role on the study. Study co-chairs are responsible for the section of the protocol specific to their modality or discipline, such as surgery, imaging, radiation, community involvement, etc. Identification as a study co-chair on the protocol face page does not assume authorship.

At least one member of the study leadership team in the role of chair or co-chair shall be a community oncologist (see section 13 of Alliance Bylaws).

6.3.2.1 Moving study co-chair to a non-Alliance institution

If the study co-chair moves to a non-Alliance institution, the study co-chair may continue to serve as study co-chair with the agreement of the appropriate committee chair and if no conflicts of interest have arisen because of the move of the study co-chair.

6.3.2.2 Replacing study co-chair

Study co-chairs will have their performance carefully evaluated and will be replaced if performance is not satisfactory. If a study co-chair is forced to relinquish responsibility for a study, the group chair (or designee) and committee chair will appoint a new study co-chair and re-assign authorship responsibility.

6.3.3 Committee chair

The committee chair is responsible for the scientific portfolio and priorities of his/her committee, including protocol development, conduct and analysis and publication of results. As delegated by the Alliance Executive Committee, the committee chair approves concepts for further development and may select or assign study chairs or co-chairs. The committee chair is responsible for submitting study concepts that emerge from his/her committee to the SCRC. For more information see section 4.

6.3.4 Primary statistician

6.3.4.1 Primary statistician
The primary statistician has primary responsibility for all statistical aspects of the protocol, including description of the study design, calculation of the sample size necessary to meet the primary objective of the study, and description of the interim and final analyses that will be used to investigate the primary and secondary hypotheses of the study. The primary statistician oversees the development of case report forms and the forms schedule.

For studies monitored by the Data and Safety Monitoring Board (DSMB), the primary statistician is responsible for preparing the monitoring reports presented to the DSMB (see section 16). After the study is closed, the primary statistician directs the final data analysis of the data and assists the study chair in preparation of a manuscript.

6.3.4.2 Secondary statistician

The secondary statistician assists the primary statistician. During the development of the protocol, the secondary statistician works in collaboration with data management staff, the primary statistician, and the study chair to develop case report forms.

6.3.5 Data managers

Data managers review protocols, create data submission schedules, and work with the study chair, statisticians, protocol coordinators, clinical research professional liaisons, oncology nurse liaisons, and information systems personnel to create new case report forms (paper or electronic). The data managers are responsible for the data management of assigned protocols.

6.3.6 Protocol coordinator

Protocol development occurs under the direction of the protocol coordinator. Protocol coordinators will establish timelines for protocol development, and work with study team members to draft, review and revise the protocol. They serve as the liaison for all protocol related correspondence with CTEP, DCP and CIRB, and are responsible for communicating official CTEP, DCP or CIRB communications to study team members.

Post-study activation, the protocol coordinator fields questions from sites, coordinates answers from study team members to sites, and works with members of the study team or other functional areas to address study issues. The protocol coordinator is responsible for managing any protocol
amendments, working with members of the study team or other functional areas as appropriate.

6.3.7 Executive officer

The executive officer, monitors protocol development and assists the protocol coordinator with issues requiring physician input, for example reviewing SCRC meeting minutes or evaluating the appropriateness of eligibility criteria or dose modifications. The executive officer assists with reviews of serious adverse events (SAEs) and CTEP Adverse Event Reporting System (CTEP-AERS) reports, provides guidance on study-specific emergency actions, reviews correspondence with NCI, and responds to queries when the study chair is unavailable. The executive officer also participates in logistical activities of protocol development, for example assessing study budget needs or study feasibility. Additionally, the executive officer assists in the coordination of industry interactions.
6.4 Protocol development

6.4.1 Protocol numbering

A concept submitted for review by the Study Concept Review Committee (SCRC) or the Translational Research Program (TRP) Executive Committee, or concepts containing data-only requests, has a study number assigned by the Alliance database (table 6-1). The study number will be assigned prior to concept review.

The first character of the study number is an A, followed by two digits that indicate the committee associated with the protocol. The next two digits indicate the year the concept was introduced. The final two digits are assigned consecutively for that committee as concepts are submitted to the SCRC. For example, the Breast Committee is A01, so A011204 would refer to the fourth breast cancer concept submitted in 2012.

Table 6-1. Alliance protocol numbering system

<table>
<thead>
<tr>
<th>Alliance Committee</th>
<th>Committee Number</th>
<th>Sample Study Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breast</td>
<td>A01</td>
<td>A011101</td>
</tr>
<tr>
<td>Gastrointestinal</td>
<td>A02</td>
<td>A021101</td>
</tr>
<tr>
<td>Genitourinary</td>
<td>A03</td>
<td>A031101</td>
</tr>
<tr>
<td>Leukemia</td>
<td>A04</td>
<td>A041101</td>
</tr>
<tr>
<td>Lymphoma</td>
<td>A05</td>
<td>A051101</td>
</tr>
<tr>
<td>Myeloma</td>
<td>A06</td>
<td>A061101</td>
</tr>
<tr>
<td>Neuro-Oncology</td>
<td>A07</td>
<td>A071101</td>
</tr>
<tr>
<td>Respiratory</td>
<td>A08</td>
<td>A081101</td>
</tr>
<tr>
<td><strong>Alliance Scientific Discipline Committee</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Experimental Therapeutics</td>
<td>A09</td>
<td>A091101</td>
</tr>
<tr>
<td>Imaging</td>
<td>A10</td>
<td>A101101</td>
</tr>
<tr>
<td>Leukemia Correlative Sciences</td>
<td>A11</td>
<td>A111101</td>
</tr>
<tr>
<td>Pathology</td>
<td>A12</td>
<td>A121101</td>
</tr>
<tr>
<td>Pharmacogenomics and Population Pharmacology</td>
<td></td>
<td>A131101</td>
</tr>
<tr>
<td>Radiation Oncology</td>
<td>A14</td>
<td>A141101</td>
</tr>
<tr>
<td>Solid Tumor Correlative Sciences</td>
<td>A15</td>
<td>A151101</td>
</tr>
<tr>
<td>Transplant</td>
<td>A16</td>
<td>A161101</td>
</tr>
<tr>
<td><strong>Alliance Cancer Control Program</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cancer in the Elderly</td>
<td>A17</td>
<td>A171101</td>
</tr>
<tr>
<td>Health Disparities</td>
<td>A19</td>
<td>A191101</td>
</tr>
<tr>
<td>Health Outcomes</td>
<td>A20</td>
<td>A201101</td>
</tr>
<tr>
<td>Prevention</td>
<td>A21</td>
<td>A211101</td>
</tr>
<tr>
<td>Symptom Intervention</td>
<td>A22</td>
<td>A221101</td>
</tr>
<tr>
<td>Cancer Care Delivery Research</td>
<td>A23</td>
<td>A231101</td>
</tr>
</tbody>
</table>
To more easily connect any embedded companion trial with a treatment study, a two-letter and number extension is added (table 6-2). For example, “A021101-ST1” is a solid tumor correlative sciences embedded companion study that appears in study A021101. If more than one type of embedded companion is included in the treatment or intervention study for the same type of companion, then sequential numbers are assigned (e.g., A021101-ST2, A021101-ST3, etc.).

Table 6-2. Alliance protocol numbering system - embedded studies

<table>
<thead>
<tr>
<th>Committee</th>
<th>Embedded Study Suffix</th>
<th>Sample Study Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cancer in the Elderly</td>
<td>EL</td>
<td>A021101-EL1</td>
</tr>
<tr>
<td>Comparative Effectiveness Research *</td>
<td>ER *</td>
<td>A021101-ER1 *</td>
</tr>
<tr>
<td>Health Disparities</td>
<td>HD</td>
<td>A021101-HD1</td>
</tr>
<tr>
<td>Health Outcomes</td>
<td>HO</td>
<td>A021101-HO1</td>
</tr>
<tr>
<td>Prevention</td>
<td>PR</td>
<td>A021101-PR1</td>
</tr>
<tr>
<td>Symptom Intervention</td>
<td>SI</td>
<td>A021101-SI1</td>
</tr>
<tr>
<td>Imaging</td>
<td>IM</td>
<td>A021101-IM1</td>
</tr>
<tr>
<td>Leukemia Correlative Sciences</td>
<td>LC</td>
<td>A041101-LC1</td>
</tr>
<tr>
<td>Pathology</td>
<td>PA</td>
<td>A021101-PA1</td>
</tr>
<tr>
<td>Pharmacogenomics and Population Pharmacology</td>
<td>PP</td>
<td>A041101-PP1</td>
</tr>
<tr>
<td>Solid Tumor Correlative Sciences</td>
<td>ST</td>
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</tr>
<tr>
<td>Cancer Care Delivery Research</td>
<td>CD</td>
<td>A021101-CD1</td>
</tr>
</tbody>
</table>

* not in use

6.4.2 Concept

6.4.2.1 Concepts other than translational research and data-only requests

Concepts are discussed at Alliance disease/modality/discipline committee meetings. If the concept includes various committee components, each relevant committee must approve the concept before it can be submitted for review.

The Alliance requires treatment studies to be submitted to the SCRC on an appropriate NCI/CTEP Letter of Intent (LOI) or Concept submission form. Cancer control studies (e.g., non-treatment studies) do not have an NCI-specific concept submission form, and are to be submitted to the Alliance SCRC in the same format as required for concept submission to NCI DCP. If applicable, concept submission should occur after NCI Task Force review. An Alliance Conflict of Interest Form (completed by the study chair) and an Alliance Concept Submission Form must
accompany the concept submission to the SCRC. Details concerning the proposed funding must be included with the concept submission.

The committee chair must submit concepts to the SCRC. If the concept is submitted by a designate, the committee chair must indicate his/her approval of the concept in writing.

Concepts submitted by investigators external to the Alliance will be reviewed by the SCRC.

6.4.2.2 Concepts containing data-only requests

Studies that only require data that are already available in the Alliance Statistics and Data Center (data-only studies), and are not part of the original objectives of the parent Alliance study, will be considered for approval once the primary study analyses are published. If the proposed study requires data from a trial that is under active monitoring by the DSMB, the DSMB must review and approve the release of the data (see section 16).

The proposed data-only study may include data generated by a correlative study. Requests for use of biospecimens are covered by a separate review procedure, as noted in the translational research section.

Requests for a data set that will be analyzed outside of the Alliance Statistics and Data Center (SDC) fall under the Data Sharing policies (see sections 6.11 and 15). Typically, these requests will originate outside of the Alliance.

The Alliance requires that Alliance-led data-only studies be submitted for review. Data-only study proposals should be submitted on the Alliance Data Sharing Request Form located on the Alliance website, under ‘concept submission.’

Prior to submission for Alliance review and approval, the request will be reviewed by the committee chair and committee statistician. The statistician will generate an Alliance SDC workload estimate. If the proposal is generated from a committee other than the committee that sponsored the original clinical/translational study, approval from that original committee chair and statistician is also required. In most cases, the original study chair will be involved in these discussions.
It will sometimes be the case that the data requested for analyses are not in the electronic database but will need to be abstracted from charts and reports. Data abstractions can only be performed if there is adequate funding and staff available.

For requests expected to require ≤25 hours of effort from the SDC, review and approval will be by the associate directors of SDC. For such requests, the investigator will be notified of the decision within three weeks of submitting all requisite items. Proposals expected to require >25 hours of effort will be reviewed by the Alliance Executive Committee.

As specified in section 6.14, proposals requiring collection of additional data from Alliance institutions are discouraged and must be reviewed by the SCRC.

### 6.4.3 Developing the protocol

#### 6.4.3.1 Communications post-SCRC and NCI concept approval

Upon approval by the appropriate concept review body Alliance SCRC, all subsequent communications with NCI CTEP must occur through members of the Central Protocol Operations Program (CPOP). CPOP submits the approved NCI LOI or Concept Submission Form to CTEP for approval. The Alliance Cancer Control Program Manager submits concepts to DCP for approval.

Once CTEP or DCP approves the concept, the study team may begin developing the protocol. The protocol coordinator maintains the official, master version of the protocol document. Upon DCP concept approval, all subsequent communications with NCI DCP must occur through CPOP.

#### 6.4.3.2 Protocol authoring

Following concept approval by the SCRC CTEP or DCP, the protocol coordinator seeds the Alliance Model Protocol template with information from the NCI approved concept/LOI. The study chair, study co-chair(s) and primary statistician(s) are responsible for authoring the first full draft of the protocol. The protocol coordinator edits the draft to Alliance standards and circulates it for initial review by the study chair, study co-chair(s), committee chair and vice chair, primary statisticians, data manager, the responsible executive officer, the, and the director of translational research.
Based on the comments received, a revised draft is constructed by the protocol coordinator and the study chair. This draft is then circulated for expanded review to the above reviewers, plus the following additional internal reviewers: director of protocol operations, group chair, IT systems management unit, and other members of data operations as appropriate. External reviewers include liaisons from Pharmacy, CRP, Oncology Nursing, and Patient Advocates Committees, as well as representatives from IROC, and specimen repositories, as appropriate.

After internal reviews are completed, the protocol is submitted by the protocol coordinator to CTEP, DCP or other appropriate review agency. The Alliance will adhere to all NCI-mandated protocol development timelines.

6.4.3.3 Determining the trial participant eligibility criteria

In general, there should be as few eligibility requirements as possible, with the requirements only excluding those for whom the study is clearly inappropriate.

Alliance studies typically require trial participants to be at least 18 years old. In certain diseases, younger patient populations may be considered.

6.4.3.4 Inclusion of women and minorities

It is the policy of the National Institutes of Health (NIH) that women and members of minority groups and their subpopulations must be included in all NIH supported biomedical and behavioral research projects involving human subjects, unless a clear and compelling rationale and justification is provided that explains why inclusion is inappropriate with respect to the health of the subjects or the purpose of the research. The inclusion of women and minorities in Alliance protocols is a standard item of CTEP review. All protocols submitted to NCI include appropriate sections on women and minorities.

6.4.3.5 Determining the trial participant follow-up period

Each protocol must explicitly state the required follow-up time, and the maximum time period for which data are required for each trial participant. The requirement is based on study objectives and statistical design considerations, including those of companion
studies. Disease committees may also specify disease-specific rules.

### 6.4.3.6 External protocol review

When ready, protocols are submitted to CTEP or DCP for review. Phase III, select phase II, and select cancer control trials are also reviewed by an NCI Central Institutional Review Board (CIRB). Changes mandated by the NCI, CIRB or FDA do not need to be reviewed by the SCRC. In other cases, significant changes to the protocol, e.g., change in trial design or a significant change to sample size, must be re-reviewed by the SCRC.

Once all necessary external and internal approvals have been secured, the protocol is activated, generally in the next scheduled protocol posting.

### 6.4.4 Developing case report forms

The following policy describes the process of assembling the forms necessary to collect the scientific data required to meet the protocol objectives. The policy covers scientific and supplemental data form development and revision. **Note: When the term "form" is used in this section, it refers to the data collection form and the form instructions, whether paper or electronic.** Scientific forms are defined as those forms that are used for study data collection. Supplemental forms are those forms providing reference information necessary for completion of scientific forms.

#### 6.4.4.1 Determining data to be collected

Decisions about the amount and type of data collected are made jointly by the study chair, committee chair, primary statistician, and executive officer, if one is assigned to the study. As a general principle, Alliance studies attempt to collect the minimum amount of data required to meet the scientific objectives of the study.

#### 6.4.4.2 Making use of standard Alliance forms

The Alliance Global Library of supplemental forms should be used for all studies. Whenever possible, the study chair and primary statistician should agree to make use of the Alliance’s existing scientific forms.
6.4.4.3 Using Translated Patient-Reported Questionnaires

The most commonly used patient-reported questionnaires for Alliance protocols will be made available in the North American primary languages, i.e., English, Spanish, and French Canadian. If a translated questionnaire is not readily available, the study chair must choose between: 1) restricting participation to English speakers only or 2) allowing accrual of patients with other non-English primary languages. If option 2, then the study chair must decide whether to: 1) pursue formal translation of the questionnaire or 2) allow on the spot translation by either professional translators at the institution or the patient’s family/friends.

The Alliance preference is to design all Alliance studies to allow accrual of patients with other non-English primary languages using on the spot translation by either professional translators at the institution or the patient’s family/friends. The Alliance Model Protocol Template includes the appropriate information for this option.

However, if a formal translation is requested, the investigator must send an email request to QOL@alliancenctn.org. All translation requests will need to be reviewed and approved by the Cancer Control Program (CCP) leadership.

6.4.4.4 Using copyrighted forms

Any use of copyrighted forms should be coordinated through the Alliance. A copyrighted form is used as-is within the Alliance form shell. NO MODIFICATIONS MAY BE MADE TO THE FORM BY ANY ALLIANCE PARTICIPANTS. Only the copyright holder may make changes.

When the use of a copyrighted form requires a fee, and there is no specific grant funding the use of the copyrighted form, approval to disburse any Alliance funds must be granted by the group chair or the principal investigator for the Cancer Control Program as appropriate.

6.4.4.5 Forms design
All Alliance forms contain basic identifying features and adhere to a common format. Appropriate data management and IT staff ensure adherence to standard Alliance case report form formats.

### 6.4.4.6 Forms review and approval

All forms and instructions go through two review stages (initial and final review) before they can be used in a study or for administrative purposes.

The following individuals provide the final forms approval:

- Primary statistician
- Clinical trials manager
- Quality review specialist
- Protocol coordinator and executive officer (as applicable) (for information only)
- Study chair
- Modality/discipline co-chairs, as applicable

Other approvals may be obtained as deemed necessary by the development team. Upon receipt of all final approvals, further changes may not be made unless required by NCI review. The Alliance will not activate a study until all form approvals have been received.

### 6.4.4.7 Forms revision

When a form requires changes after study activation, the study developer will revise the form following either an expedited change pathway in the case of urgently needed changes or the bundled changes pathway. Changes will be bundled if the change request is not related to patient safety or primary endpoint analysis. Bundled changes will be pushed to production per a regular schedule. Forms distribution system

Most Alliance forms are available on the [Alliance website](http://alliance.org). Forms not available on the website may be obtained by contacting the appropriate Alliance data manager.

### 6.4.5 Participation in intergroup studies
With few exceptions, all studies are to be available to all members of the NCTN. Exceptions may include certain DCP sponsored studies and selected phase I or early phase II studies. Studies may have co-chairs from other groups who were involved in the study design added to the protocol. These individuals should be included in protocol development when possible and must be adequately informed about progress and problems with the protocols for which they are responsible. Substantive amendments, e.g., those changing the study design or requiring a significant change in sample size, must always be discussed with representatives of the other groups.
6.5 Activating a study

After receiving final protocol approval from CTEP or DCP, the Alliance Protocol Office activates the study, in coordination with Alliance IT, registration, and data management staff. A notice indicating that a study is officially open for accrual is issued by the responsible protocol coordinator in the protocol posting on the Alliance website.
6.6 Waivers

6.6.1 Eligibility waivers

No eligibility waivers will be granted.

6.6.2 Other waivers

The Alliance adheres to CTEP’s policy not issue or approve any waivers for protocol deviations, including eligibility criteria, treatment schedules, dose modifications, toxicity assessment, response criteria, and statistical aspects.
6.7 Updating a study

6.7.1 Revisions and amendments

Protocol updates containing revisions and amendments may be generated in response to decisions by the study chair to change some aspect of the study design or conduct. All amendments that are not merely editorial in nature will be reviewed by the following: study chair executive officer (if applicable), committee chair (if applicable), and primary statistician, the executive officer in charge of drug distribution (if applicable), the, director of translational research operations, and data management personnel.

Updates may also be generated in response to information or requests from external agencies, such as safety letters or action letters distributed by CTEP.

For any studies monitored by the DSMB, approval of substantive updates by the DSMB is required prior to submission to NCI. If the update includes changes in the trial design, these changes must first be discussed with NCI before submission to the DSMB, unless the DSMB has requested the change in trial design based on safety or outcome data available only to the DSMB.
6.8 Suspending a study

A suspension is a temporary cessation of accrual to a protocol, either planned or unplanned. Suspension may also result in a temporary cessation or modification of treatment of patients already registered to a study. An unplanned decision to suspend a study may be made by the study team based upon the recommendation of the NCI CTEP/DCP or industry partner, study chair, the primary statistician, relevant committee chair(s), or the DSMB.
6.9 Unblinding trial participants

The Alliance conducts clinical trials that mask, or blind, the identity of treatments given to trial participants and, sometimes, investigators. The DSMB, CTEP, or DCP may recommend that study accrual be stopped and treatment assignments be unblinded for all trial participants because of toxicity or safety concerns.

There are three scenarios, described below, where treatment assignments may be unblinded for individual trial participants.

Intentional unblinding of a treatment assignment, other than by the methods described below, is a serious breach of scientific ethics. The Alliance policies concerning scientific misconduct will be employed to investigate and report such incidents (see section 3.4).

6.9.1 Emergency unblinding

A trial participant’s treatment assignment can be unblinded in emergent situations with approval of the appropriate Alliance executive officer (or designee) only if unblinding would influence management of the situation, e.g., if a child has swallowed a vial of pills. Study chairs, primary statisticians, and other Alliance staff are not permitted to approve emergency unblinding requests. Emergency unblinding requests should be directed to the executive officer on call, 24 hours a day, 365 days a year. If an Alliance executive officer determines unblinding is warranted, they will contact the registration office staff. The executive officers and the Group chair are the only personnel who can unblind a study patient.

6.9.2 Protocol Specific unblinding

The protocol may specify that a trial participant’s treatment assignment can or should be unblinded based on certain criteria as specified in the protocol, such as for the purpose of crossover from placebo to active drug at disease progression. Protocol-specified unblinding may be performed by the Registration Office during regular business hours, with confirmation from the primary statistician (or designee) that the protocol-specified criteria have been reached. No executive officer (or designee) approval is required.

6.9.3 Elective unblinding

If allowed per-protocol, a trial participant, family member, or treating physician may request unblinding of the treatment assignment in non-emergent situations in order to inform subsequent disease management decisions. Elective unblinding is only permitted if the trial participant has met
<table>
<thead>
<tr>
<th>Policy Name: Unblinding Trial Participants</th>
<th>Policy Number: 6.9</th>
</tr>
</thead>
<tbody>
<tr>
<td>Section: Study Protocol – 6</td>
<td>Date Revised: January 1, 2018</td>
</tr>
</tbody>
</table>

the trial's primary endpoint. Elective unblinding will be performed by the Registration Office during regular business hours, with confirmation from the primary statistician (or designee) that the appropriate criteria have been met. If the patient has not met the primary endpoint, or if the appropriate criteria have not been met, the Registration Office will refer the caller to the appropriate executive officer (or designee) to discuss the situation. The protocol and Model Consent Form must specify whether elective unblinding will be permitted and, if permitted, that requestors should contact the Registration Office.
6.10 Closing a study

Closing a study means that accrual to the study is permanently stopped. It is possible to close only a portion of a study.

6.10.1 Procedures for closing a study

The decision to close a study is made by the primary statistician, in consultation with the study chair and committee chair (and the DSMB for phase 3 studies or other studies monitored by the DSMB). If unexpected adverse events occur, members of the study team may initiate the process. For phase 3 studies (or other studies monitored by the DSMB), the DSMB may recommend early closure of a study for reasons of patient safety or of differential treatment effectiveness.

For routine study closures, in order to allow sites to register patients who are already in the process of being worked up for the study, the Alliance routinely sets a future closing date, usually two weeks, once adequate accrual has been achieved. This may result in modest over-accrual to the study. Exceptions to this policy are phase 1 studies, for which over-accrual is not allowed, and certain phase 2 studies. These studies require tighter control of the number of patients registered and treated. More rapid study closures may be necessary for patient safety reasons.

6.10.2 Notifying patients about early closure of clinical trials

Disclosure to individual participants of study results often follows a recommendation that accrual be terminated early and/or that protocol specified treatment be discontinued or significantly modified. However, disclosure must not violate any state or federal laws regarding breaking the code on anonymized data.

The trial participant who provided the original consent to participate in the research is informed of the results of the clinical trial by his/her treating physician or designee. Participants are informed in a manner that will ensure that they receive the results with a minimum of disruption to the patient-physician relationship.
6.11 Release of data

6.11.1 Studies monitored by the DSMB

If a trial is being monitored by the DSMB (see section 16), requests for release of data (immature and mature) to the study team must be submitted to the DSMB. If the request is approved, the data can be released to the study team and can only be used within the scope specified by the DSMB in their approval, see section 16.2.6.

6.11.2 Studies not monitored by the DSMB

6.11.2.1 Adverse event/toxicity data

If adverse event/toxicity data are not the primary endpoint for a trial or a key secondary endpoint, these data should be freely available to the internal study team for analysis throughout the trial, even if they are a secondary endpoint. Note that if the trial is a blinded trial, the assessment of the data must adhere to the NCI policy for reporting adverse event data for blinded studies.

6.11.2.2 Mature endpoint data

When the primary statistician has ascertained that the study endpoint data have met the criteria as described in the protocol for final analysis, the data can be released to the internal study team for analysis. Results of the analysis can be made public through abstracts, presentations, and publications.

6.11.2.3 Immature endpoint data

Immature endpoint data are data that have not met the criteria as described in the study protocol for final analysis.

6.11.2.3.1 Study is closed to accrual

If a study is closed to accrual but the endpoint has not yet met the criteria as described in the protocol for final analysis, the internal study team must submit a written request for access to the data to the Alliance committee (co-) lead statistician(s). The request should specify the following:
- The purpose of accessing the immature endpoint data (e.g., for planning a new study, for potential modification of the existing study)
- The endpoint data being requested
- The data analysis plan for the requested endpoint data
- The individuals who will have access to the analysis results
- How confidentiality will be ensured
- The potential impact on the study

If approved by the Alliance committee (co-) lead statistician(s), the data will be released to the study statisticians for analysis. The results of the analysis can only be shared with the individuals specified in the request, can only be used for the purpose stated in the request, and must be kept confidential.

6.11.2.3.2 Study is open to accrual

Requests for access to endpoint data while a study is still accruing patients will be granted only in extraordinary circumstances. If a study is open to accrual, the internal study team must submit a written request for access to the data to the program director and associate chair of the Alliance Statistics and Data Management Program. The request should specify the following:

- The purpose of accessing the immature endpoint data (e.g., for planning a new study, for potential modification of the existing study)
- The endpoint data being requested
- The data analysis plan for the requested endpoint data
- The individuals who will have access to the analysis results
- How confidentiality will be ensured
- The potential impact on the completion of the study
If approved by the leadership of the Alliance Statistical Units and Data Management Program, these data will be released to the study statisticians for analysis. The results of the analysis can only be shared with the individuals specified in the request, can only be used for the purpose stated in the request, and must be kept confidential.

6.11.2.4 Appeal process

If the internal study team disagrees with a denial for early access to the study data, they can appeal. For closed trials, the appeal should be made to the program director/co-director of the Alliance Statistics and Data Management. Unit. For open trials, the appeal should be made to the associate chair of the Alliance Statistics and Data Management.
6.12 Completing a study

A study is declared completed by the study chair, the primary statistician and the relevant committee chair(s). Ordinarily, this occurs when the study has met all of its objectives, a definitive analysis has been performed, and an article has been published. Rarely, a study may be declared completed when the study chair and statistician agree that no analysis or publication of the study will be done. This latter category is considered “completed-administratively.”

The classification of a study as “completed” has operational consequences, indicated below.

6.12.1 Archiving paper records

CALGB Legacy studies - Paper files of patient data are kept at the CALGB Statistical and Data Center for three years after study closure to be available for institutional audits. Three years after closure, paper records are archived in Duke off-site storage if a study is completed. These records can be retrieved within 24 hours by contacting the staff assistant at the Data Operations Office, who is responsible for requesting delivery from the storage facility.

ACOSOG & NCCTG Legacy studies – As applicable, paper files of patient data are stored electronically at the Alliance Statistics and Data Center in a document imaging system. Upon receipt of records they are scanned and stored electronically. The system is web-based and records can be viewed once authorization access has been approved. The stored electronic data are available for audit by requesting them from the Data Operations Office.

6.12.2 Archiving study database

The data for a completed study remain in the Alliance database.

The Alliance Statistics and Data Center maintains a library of data sets used in monitoring reports, interim analyses and manuscripts.

The data sets used in monitoring reports, interim analyses and manuscripts are stored as SAS data sets or ASCII files with attached data dictionary. The statistician who prepares the reports or analyses is responsible for copying the necessary data files. The statistician uses naming conventions to index the data files by the study number, the type of report and the date the report was prepared. All data sets are archived on a designated archive server. At the discretion of the statistician, additional files may also be archived.
6.12.3 Study chair access to additional data

Copies of data received by the Alliance Statistics and Data Center for completed studies are not automatically sent to the study chair unless explicitly requested by the study chair. All requests for study data should be sent to the study statistician.
6.13 Terminating a study

Studies may have all follow-up terminated for all trial participants either because all trial participants have been followed for the protocol-specified period or because it is decided that no further follow-up is needed. Upon termination, no further follow-up data, including new queries, are collected from participating sites. All studies are reviewed annually by the primary statisticians to determine if continued follow-up is required. A list of all studies with terminated follow-up is publicized on the Alliance website.

Study team members wishing to extend patient follow-up beyond the protocol-specified interval must obtain permission from the group statistician. A protocol amendment must also be generated.
6.14 Retrospective data collection from closed or completed studies

Generally, proposals that require the collection of additional material from Alliance sites will not be approved. Retrospective collection of data is expensive and time-consuming. These requests usually require IRB review at each participating site and may require obtaining additional patient consent and/or authorization. The Alliance may consider such requests in special circumstances provided adequate funding is available for both the Alliance Statistics and Data Center effort and for participating institutions. Studies that require the collection of additional material will be reviewed by the Alliance Study Concept Review Committee.
### 7 Patient registration

All Alliance institutions are allowed to register patients to Alliance and other trials posted on the Cancer Trials Support Unit (CTSU) menu.

#### 7.1 Authorization of institutions to register patients

Institutions intending to register patients must have IRB approval for the study. IRB approval documentation must be submitted to CTSU for entry into RSS, prior to enrollment of the first patient. Submission instructions are available on the Regulatory page of the CTSU website.

Compassionate (expedited, emergency) approval, in which an institution wants immediate approval to put a patient on a treatment study not yet approved by its IRB, is not allowed. The IRB must give full-board approval before patients may be registered on a treatment study. Select non-treatment studies, such as laboratory or survey studies that present minimal risk to participants, may qualify for expedited review, which is noted at the time of protocol activation.

Institutions may have their accrual privileges suspended by the Alliance leadership.

#### 7.1.1 Limited access studies

Some studies may limit access to a subset of institutions, for quality assurance or other reasons (e.g., phase 1 studies). Participating sites in limited access studies will be identified in CTSU RSS.
7.2 Authorization of participants to register patients

All Alliance studies will use the Oncology Patient Enrollment Network (OPEN) online registration system (https://open.ctsu.org) maintained by the CTSU except where otherwise indicated in the protocol. Site participants who register patients on NCTN trials must be registered with the Cancer Therapy Evaluation Program (CTEP) and have a valid and active CTEP Identity and Access Management (IAM) account. This is the same account (username and password) used to access the member portion of the CTSU website (https://www.ctsu.org). Registration is accomplished via the NCI Registration and Credential Repository (RCR). Refer to the CTEP website for additional details on registration types and required documentation.

To perform patient registrations (including pre-registrations), the site user must be assigned the ‘Registrar’ role in the CTSU’s Regulatory Support System (RSS), found under the ‘Regulatory’ tab in the member portion of the CTSU website. The principal investigator (PI) of each main member must approve all personnel authorized to register patients.
7.3 **Credentialing**

If a protocol requires credentialing for the registering physician (e.g., to demonstrate proficiency in performing a particular type of surgery) or the registering institution (e.g., to administer radiation therapy), then the credentialing requirements listed in the protocol must be met before patient registration may proceed.
7.4 Confirming patient eligibility

The institution confirms eligibility before registration or randomization by verifying the eligibility criteria listed in the protocol. Institutions should refer to all relevant protocol sections in order to ensure that all conditions for appropriate entry of a patient on study are met.

Exceptions to eligibility criteria or other protocol requirements will not be granted.

Treatment should begin no later than one week after registration, unless otherwise specified in the protocol.

As a general rule, sites should not register a patient to more than one interventional study when it is expected that one protocol’s intervention might impact the other study’s endpoints (e.g., registering a patient for two studies, where both protocols’ treatments are expected to have an impact upon response, overall survival, etc.), although exceptions may be allowed by the study chair in collaboration with the executive officer. The study team should carefully consider the scientific and practical implications before allowing such exceptions.
7.5 Procedures to register patients to Alliance studies

Registration to Alliance studies is available 24 hours a day via OPEN. All participating sites (Alliance and non-Alliance sites) will use OPEN to enroll patients. OPEN can be accessed from the member portion of the CTSU website.

A study-specific Registration Worksheet/Eligibility Checklist is available for each study. Information required at registration includes:

- Registering institution and investigator names and CTEP ID numbers
- Patient demographic information
- Pre-study and eligibility information
- Stratification factors
- Companion study participation information if applicable

The OPEN system will provide the site with a printable confirmation of registration and treatment information.

7.5.1 Pre-registration

For select studies it is necessary to obtain a patient ID for study screening and eligibility using a pre-registration procedure. Patients will be pre-registered using OPEN.
7.6 Registration on weekends or after business hours

Patients must be consented and registered to a treatment study before protocol treatment begins, with the following exception.

Treatment prior to registration is allowed if all of the following criteria are met:

- Patient is to be registered to an acute leukemia or high-grade Burkitt’s-like lymphoma non-randomized study, or to a study in which the induction arm is standard chemotherapy.
- Immediate treatment is necessary (i.e., patient is in medical crisis).
- This treatment policy exception is stated in the protocol.

Patient must be registered on the next business day. The institution must document in writing the reason why treatment was started before registration and submit documentation to the Alliance Statistics and Data Center.
7.7 Registration to companion studies

Patients may be registered to companion studies at the same time as they are registered to the treatment study. It is important for the registering institution to check protocol requirements for companion studies (e.g., whether patient participation is mandatory or optional) before registering the patient. The majority of companion studies are “embedded” within the treatment study, that is, the description of the companion study, registration and data collection procedures, and consent are included within the treatment protocol and consent form. Some companions are “freestanding”, that is, described in a separate protocol document with a separate consent form. Freestanding companion studies may be optional for the institution as well (i.e., they do not need to be offered to the patient).

A patient who has been registered to a treatment study may later be registered to a companion study, if allowed by the protocol. This may happen even though the registering institution for the treatment study is no longer a member of the Alliance, provided the institution has accepted responsibility for the patient via transfer, including patient registration and submission of the patient's data.
7.8 Procedure to register patients to intergroup studies

For registration of patients to intergroup studies not coordinated by the Alliance that are available on the CTSU menu, the Alliance institution must use OPEN. The institution must indicate its network group affiliation with the Alliance in order to receive enrollment credit for the Alliance.
8 Data management

8.1 Data submission

8.1.1 Completing forms

8.1.1.1 Alliance general instructions: all forms (electronic CRFs and paper forms)

All data forms and supporting documentation as required by the study are submitted to the Alliance Statistics and Data Center (SDC) using either Rave (for Alliance studies) or the legacy CALGB, and NCCTG systems. Access to Rave requires that the site has IRB approval of the study and that site staff have an iMedidata Rave account and have completed eLearning for their Rave role.

Use forms specified in the study data submission schedule and available on the Alliance website (http://www.allianceforclinicaltrialsinoncology.org), CTSU website or in the electronic data capture system used for the study. Do not store electronic copies of the form on your computer; always download the most recent copy from the Alliance website or CTSU site. Forms for intergroup studies are distributed by the coordinating group and may be obtained from their website, the CTSU website (https://www.ctsu.org), or iMedidata. If you are unable to locate an intergroup form, contact the responsible coordinating group.

When submitting copies of hospital records (path reports, lab results, etc.) make ONE-SIDED COPIES ONLY. Remove all patient identifiers and write the Alliance study and patient number on each page. For Rave, the supporting documents can be uploaded to the eCRF.

8.1.1.2 Instructions for forms submitted during treatment and follow-up

Many Alliance forms are study- or disease-specific, but these general instructions may be used for all forms described below.

1. Each form must be accompanied by required documentation as specified in the protocol for the same time period. Check the data submission schedule in the forms package for required
data. The information recorded on each form should reflect only those events occurring during the time period covered.

2. The time period covered by each form is specified in the data submission schedule. The coding convention for the covered time period is as follows:

If the data submission schedule states that forms are required for each phase/cycle of treatment, the time period covered by the forms should be from day one of each treatment phase/cycle up until the administration of the subsequent treatment. This allows for capture of responses and adverse events attributable to the entire phase/cycle but not fully assessed until the patient returns for the next treatment phase/cycle.

**Follow-up and response forms**

1. Record the dates of objective status, e.g., response or progression, only during the time period in which the event occurred.

2. The criteria for assessing response are specified in the protocol. For example in many solid tumor studies, overall objective status is determined per RECIST criteria.

3. Supporting documentation of response, relapse or progression must be submitted as required by the protocol.

**Adverse event forms**

General instructions for all adverse event forms are as follows:

- All studies use the NCI’s Common Terminology Criteria for Adverse Events (CTCAE) that is available on the Alliance and the Cancer Therapy Evaluation Program (CTEP) websites (http://ctep.cancer.gov/protocolDevelopment/electronic_applications/etc.htm). Use only these criteria to identify events and determine grade severity. The version of the CTCAE is specified within the protocol.

- The forms used with the CTCAE are study-specific. Each form provides a list of solicited events for which grade must be coded. Additional fields are provided for specifying other events that occur.
• Code grade “5” if the event caused the death of the patient. Code only one grade 5 event for a patient. Code contributing events that are not the primary cause of death per CTCAE grade criteria.

• Note that for some events certain grades are not defined and are not allowed (e.g., grade 3 or 4 alopecia).

Adverse Event Expedited Reporting System (CTEP-AERS)

Expedited adverse events are reported using the NCI’s Cancer Therapy Evaluation Program Adverse Event Reporting System (CTEP AERs, located at https://ctepcore.nci.nih.gov/ctepaers. Guidelines for CTEP-AERS reporting are included in each protocol.

Only file one CTEP-AERS report per course/cycle. Amend the previous report for the cycle if the adverse event data needs to be corrected, the adverse events worsen, or new adverse events occur that require expedited reporting.

• Don’t assume that all hospitalization require CTEP-AERS reporting— check the protocol.

• The “Surgical Intervention” section is to be used ONLY for the protocol related surgery.

8.1.2 Submission of data forms

The Alliance requires capture of data per protocol for all patients on Alliance treatment studies. Data continue to be submitted per protocol until the patient reaches the endpoints defined in the protocol (e.g., relapse/progression), or until follow-up is discontinued per protocol instructions.

8.1.2.1 General data submission instructions

For patients on phase 1, 2, and 3 studies, data submission is required as indicated by the general rules in table 8-1. However, data submission requirements specified in the protocol take precedence over those indicated in the table. For example, if a study includes treatment with a drug that may cause chronic toxicity, the protocol may require collection of adverse event data after study endpoints have been reached.
For all patients registered to phase 1, 2, and 3 treatment studies, survival information must be provided as specified in the protocol. Survival data must continue to be submitted until indicated otherwise by the protocol, that is until the patient reaches a follow-up truncation point stated in the protocol or until follow-up is discontinued for the entire protocol. Survival dates and dates of death must contain the day, month and year the patient was last known to be alive.

Death of a patient is reported on the forms specified in the study-specific data submission schedule.

**Overdue data**

The current expectations for form submission before being considered delinquent are: Baseline and treatment forms: within 30 days of target date, Follow-up Forms: within 60 days of target date.

For studies using Medidata Rave, study-specific delinquency lists are available in real time via the Rave task summary. To assist with site performance, delinquent data reports are provided by the Alliance SDC and are available on the Alliance website to main members rostered lead and secondary lead CRPs.

**New malignancies**

All new malignancies that occur following treatment and fall within the protocol specified time period must be reported.

Table 8-1. Alliance data submission guidelines

<table>
<thead>
<tr>
<th>Status</th>
<th>Example</th>
<th>Baseline/On-Study</th>
<th>Off Treatment</th>
<th>Adverse Event +</th>
<th>Endpoint Data Relapse/or Progression</th>
<th>New Malignancy +</th>
<th>Survival</th>
</tr>
</thead>
<tbody>
<tr>
<td>Typical Patient Pathway</td>
<td>Patient on or has completed RX, but has not reached follow-up completion as defined in protocol</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Ineligible</td>
<td>Patient registered to study and deemed ineligible after SDC review</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>
8.1.2.2 Registered patients who never receive treatment (canceled patients)

Patient eligibility and willingness to participate in the protocol must be carefully assessed prior to registration to ensure the patient’s ability to comply with protocol requirements. A patient may not be removed from an Alliance protocol after being registered. Patients will be given a status of “canceled” if no protocol treatment is ever given.

For canceled patients, the institution provides the SDC with sufficient paperwork to document the reason why treatment was never given. Data submission requirements for canceled patients are provided in table 8.1.
8.1.2.3 Transfer of patient to another institution

A patient on an Alliance study may transfer their study related care to another institution. It is the responsibility of the institution transferring the patient to ensure that all transfer procedures are followed. The institution accepting the patient transfer must have IRB approval for the protocol. A transferring patient must sign a new informed consent form with the accepting institution.

Prior to the transfer, the site clinical research professional (CRP) ensures that all data are up-to-date and all queries have been addressed and resolved. This will be confirmed by the Alliance Data Manager prior to the patient being officially transferred. Copies of all data required by the protocol and subject records must be submitted to the accepting institution. Once the data are updated the site is required to call the Alliance Registration Office for official documentation of the transfer and transfer of responsibilities.

Both sites will be responsible for their data. The transferring institution is eligible for audit of all patient data submitted up to the date of transfer. The accepting institution is responsible for submitting all subsequent data required by the protocol after the informed consent is signed.

For patients registered via one of the Alliance legacy registration systems, both the treating investigator at the transferring institution and the treating investigator at the accepting institution must complete the Alliance Patient Transfer Form, which can be found on the Alliance website. The completed Alliance Patient Transfer Form must be sent to the Alliance SDC, per the instructions on the form.

The sites should follow the CTSU guidelines for patients registered via OPEN. Both the treating investigator at the transferring institution and the treating investigator at the accepting institution must complete the CTSU’s patient transfer form, which can be found on the CTSU website. The completed form must be sent to the CTSU Operations Center.

The Alliance database does not reflect the transfer until the completed transfer form has been signed by both institutional treating investigators and has been received at the Alliance SDC. For patients registered through OPEN, the CTSU will forward the
transfer information to the Alliance SDC. The Alliance database continues to reflect accrual from the institution that registered the patient.

8.1.2.4 Withdrawn consent to treat or follow

If a patient refuses further protocol treatment after therapy has begun, the institution continues to submit all data required by the protocol unless the patient specifically withdraws consent to be followed. A research participant’s discontinuation or refusal of research treatment or intervention is not a withdrawal of consent to participate in the research study. This participant is still considered to be part of the study and should be followed per protocol/group policy.

A patient may, on rare occasions, withdraw consent for continued protocol participation. A verbal or written withdrawal of consent by the patient must be documented in the patient’s research record. A patient’s refusal to comply with follow-up visits or requirements is not considered to be an implied withdrawal of consent. Institutions must follow the Confirmation of Lost to Follow-up procedure (section 8.1.2.5) for a noncompliant patient who has not specifically withdrawn consent.

The institution must have written documentation that clearly states the level of withdrawal for follow-up. Written documentation can include one or more of the following:

- A signed and dated letter from the study participant documenting the withdrawal of consent (preferred).
- A clinic note from the research record documenting the date of phone conversation with study participant and the withdrawal of consent.
- A signed and dated letter from the Principal Investigator or treating physician on institution letterhead documenting the withdrawal of consent.
- A signed and dated letter from the study participant’s power of attorney or guardian documenting the withdrawal of consent on the participant’s behalf.

The statement indicates whether the patient is withdrawing consent solely for clinical follow-up or if both clinical and survival follow-up are refused. The following is the suggested wording for the refusal statement:
“[Patient’s initials, Alliance ID #] withdrew consent to be followed with respect to (clinical status/clinical and survival status) on Alliance study (study #). Treatment Physician’s signature_____________ Date signed________________”

A copy is kept in the patient’s record at the follow-up institution. Based on information provided in the statement, the patient is removed from requirements for further follow-up of the appropriate type. All required study data up to and until the date consent withdrawal declared is expected to be submitted to the lead group/sponsor. Data generated after the date consent withdrawal declared should not be submitted. Patients that have withdrawn consent are removed from calculations of institutional performance related to timeliness. However, the percentage of patients that have withdrawn consent is included in the metrics for institutional performance related to data quality (see section 2.10).

A study participant may rescind their consent withdrawal. Upon rescission of the consent withdrawal:

- Study participant’s status is re-activated.
- Documentation may be provided in same fashion as for consent withdrawal designation.
- Reminders and data expectancy are reactivated.

8.1.2.5 Confirmation of lost to follow-up status

Institutions may confirm that a patient is lost to follow-up using specific procedures.

Note: Study participants who refuse aspects of participation or withdraw consent from further participation should not be designated as lost to follow-up. The guidelines for patient withdrawal/consent withdrawal should be followed in these situations.

8.1.2.5.1 Procedure for confirming a patient is lost to follow-up

After a period of two years in which the institution has tried with unsuccessful results to contact a patient, the patient may then be declared lost to follow-up. Institutions may confirm that a patient is lost to follow-up. Recommended contact strategies include:
• Contact the patient by phone (e.g., residence, work, cell). Search the patient’s medical record.
• Contact patient’s primary care physician (e.g., family doctor) if permitted. Check appropriate registries for the region for information about the patient’s death.
• Contact people listed for the patient (e.g., family members) if permitted.
• Send a letter or letters to the patient at the last known address. A diligent effort to contact the patient is required and should be documented.

For the patient to be confirmed lost, the institution must provide the Alliance SDC with the Alliance Confirmation of Lost to Follow-up form.

The Alliance SDC does not require submission of additional details of the attempts to contact the patient, but documentation of the attempts made during the 2 years should be retained in the patient’s institutional research record for purposes of audits.

### 8.1.2.5.2 Retrospective data submission

If a patient is confirmed lost, the institution continues to be responsible for submitting protocol-required data (e.g., on-study, treatment, follow-up information) for the period from patient registration through the date the patient is deemed lost to follow-up. For the period of time between the last contact with the patient and the date they are deemed lost to follow-up, the site must record in Rave that no contact occurred including the date of the attempt to contact the patient.

### If a lost patient is found

If a patient is re-contacted or additional data are received that change the patient’s survival or clinical status (from “lost to follow-up”), the institution must contact the data manager for the study. The data manager will inactivate the Lost to Follow up form and advise the site as to the appropriate forms for completion and submission.
8.1.3 Submission of samples, specimens, and modality materials

Specimens and modality materials (e.g., karyotypes, images) are to be submitted to the modality office or repository as specified in the Alliance protocol. Procurement, processing, submission schedules, and shipment instructions are provided in each protocol, as well as in the Alliance Biospecimen Management System (BioMS). Alliance patient ID number, study number, institution, and specimen ID should appear on submitted materials, unless otherwise specified in the protocol.

If a registered patient refuses further protocol treatment but agrees to be followed, samples may be submitted as required by the protocol. If a registered patient withdraws consent for participation in the study or consent for follow-up, samples may not be submitted. At any point in the trial, study participants can withdraw consent to (1) further specimen collection, and/or (2) change their permissions for future use of previously collected specimens. If samples have already been submitted but not distributed to investigators, when the patient withdraws consent, those samples will be withdrawn from the biorepository and will be disposed of appropriately – either destroyed or, in the case of paraffin blocks, returned to the submitting institution. Attempts will be made by the repository staff to retrieve any samples that have been sent from the repository to investigators. However, processed samples and the research data generated from them will not be rescinded, and may be used in study analyses. See sections 11.2 and 11.3 for additional information.

8.1.4 Submission of samples for intergroup studies

Samples, specimens, and modality materials are submitted per protocol-specific instructions.
8.2 Receipt and distribution of data forms by SDC

Refer to the data submission section of the protocol for instructions on how to submit data to the Alliance Statistics and Data Center.

Data for studies coordinated by other network groups are submitted directly to the coordinating group via the instructions outlined in their data submission section of the protocol.
8.3 Quality assurance performed by Data Management Unit

Data submitted for Alliance-coordinated studies are reviewed by the data manager responsible for the study. Quality assurance checks are performed to verify the completeness and accuracy of reporting, as well as intra- and interform consistency. A careful review of the data also is conducted to evaluate protocol compliance, e.g., patient eligibility, stratification, safety reconciliation, treatment and endpoints. When discrepancies are found or data are missing, data personnel query the institution.

8.3.1 Quality checks of on-study and eligibility data

Quality checks of on-study data include a detailed review of eligibility criteria and supporting documentation requested in the protocol. The first eligibility review is performed via the OPEN registration system. Upon receipt of the eligibility material and supporting documentation the DM performs a second quality check.

If a patient is found to be ineligible or of questionable eligibility, the data manager will request review by the study chair. If the study chair and data manager do not agree on the eligibility of a patient, the study statistician attempts to resolve the problem. If the statistician cannot resolve the problem, the statistician will contact an executive officer or the group chair for determination. The data manager will notify the institution of any patients deemed ineligible.
8.4 Alliance case evaluation process

Within a large clinical trials network, it is essential that patient information is collected and quantified in a standard manner across institutions and in particular that adverse events and outcome measures (response, relapse, etc.) are properly assessed. A case evaluation is a formal, centralized, clinical review by the study chair on the accuracy and consistency of key adverse event and outcome data reported by the treating institution for an individual patient (case) entered on a treatment study. The evaluation, which is required by the NCI, provides a centralized review of the data forms and other supporting documents by a medical expert, and ensures accurate data.

The patient (case) on a specific treatment study, not the institution, is the unit of evaluation. It is not the intent of the case evaluation process to evaluate the institution. Institutional evaluation is an independent process and is described in section 2.10.

8.4.1 Objectives

The objectives of the case evaluation process are to provide an assessment by the study chair of the following:

- Treatment compliance
- Study endpoint(s)
- Adverse events

8.4.2 Studies requiring case evaluation

Only studies that contain an intervention component, whether for cancer treatment or control, require case evaluation. Case evaluations may be performed on other studies upon request of the study team and joint approval of the director of statistics and director of data management. Similarly, if a study team wishes to have their study excused from these requirements joint approval is necessary.

The study chair has the final responsibility for the case evaluation. While study chairs and other study team members are involved in ongoing monitoring and review of all patient data, a case evaluation is usually performed only once per patient. A patient summary report is created when a patient reaches an endpoint defined in the study. The study chair is notified when a report is generated. The study chair can perform the review in real time or in small batches. If the reviews are performed in batches, the schedule for the review is determined by the study team. The entire case evaluation process for the study must be completed prior to the final statistical analysis to be used for publication of results.
For studies with fewer than 100 patients, all cases must be evaluated by the study chair. For large studies with 100 or more patients, the first 100 consecutive patients enrolled, and then 10 percent of the remaining target up to a maximum of 300 cases must be evaluated. Patients who were enrolled but never treated may be omitted. Additional cases may be evaluated as deemed necessary by the study statistician. All exceptions must be approved by directors of data management and statistics. In particular for some trials, review of all protocol specified events may be required.

Abstracts submitted to professional society meetings are exempt from these requirements. For most internal purposes (e.g., routine progress reports, interim analyses), it is not essential to have completed the number of case evaluations required for an external publication.

### 8.4.3 Case evaluation form

The study chair completes the case evaluation form to record his/her evaluation of the case based on an assessment of the patient summary report of data coded on the case report forms, and any other supporting documentation. The case evaluation form solicits the study chair's opinion regarding adverse events, response, relapse or disease progression, and survival as recorded in the database. The study chair provides specific comments about treatment violations or inadequate reporting.

#### 8.4.3.1 Patient summary report

Provided by the SDC, the patient summary report is a computer-generated review of a patient’s major clinical events. Reports will be based on a core set of items for all studies; additional items are determined by study phase and type (cancer/non-cancer treatment, QOL, etc.). Table 8-2 outlines the data topics included in the report.

<table>
<thead>
<tr>
<th>Treatment Compliance</th>
<th>Study Endpoint(s)</th>
<th>Study-Specific For Other Studies</th>
<th>Adverse Events (when applicable)</th>
<th>Additional Items</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date treatment or intervention started</td>
<td>Required for all studies:</td>
<td>Examples: skeletal-related event, lymphedema, submission of final questionnaire</td>
<td>Phase 1 studies:</td>
<td>Determined by study team</td>
</tr>
<tr>
<td>Date treatment or intervention ended</td>
<td>Primary and critical secondary endpoints</td>
<td>Date(s) of endpoint(s)</td>
<td>Phase 2 studies:</td>
<td>Approved by directors (Data Management, Statistics)</td>
</tr>
<tr>
<td>Number of cycles or interventions given</td>
<td>Examples: clinical tumor response, pathologic tumor response, disease recurrence or progression, death</td>
<td></td>
<td>Phase 3 studies:</td>
<td>Case by case basis</td>
</tr>
<tr>
<td>Reason treatment or intervention ended</td>
<td>Date(s) of endpoint(s)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dosing compliance (for treatment trials)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
8.4.4 Procedures

The data manager, who is assigned to the specific study, monitors and coordinates all case evaluation procedures. A patient summary report is generated at the time of case evaluation. The study chair reviews the data in the patient summary report to complete the case evaluation form. The study chair may also review the case report forms as part of the review process. A study chair’s access to additional case report form data is based on study phase. Study chairs will have full access to data for phase 1 and 2 studies. Study chairs will not have access to case report forms for phase 3 studies.

The data manager reviews discrepancies and other problems noted by the study chair, and queries the site if necessary.

The study statistician will be notified if the study chair has not completed their review according to the agreed upon timeline. Serious delinquency of the study chair will be reported to the committee chair. Possible consequences for serious delinquency are prevention from serving as future study chair and loss of authorship on the primary manuscript.
9 Information systems

The objective of Alliance Systems Management Unit and Information Systems Unit (SMU/ISU) policies is to ensure the confidentiality, integrity, and availability of Alliance systems and data. ISU staff is located in Durham, NC (Duke University); Rochester, MN (Mayo Clinic); St. Louis, MO (Washington University); and Chicago, IL (University of Chicago). SMU staff is located in Rochester, MN, and Durham, NC. ISU and SMU personnel work collaboratively, maintaining controls at all levels to ensure that all necessary standards are met.

This policy and procedures document contains two parts. Member Information describes policies and procedures for Alliance members who require access to the ISU applications, databases and equipment. SMU/ISU Operations shows policies and procedures used by the SMU/ISU staff to establish and maintain the applications, databases, and equipment for which they are responsible.

9.1 Member information

SMU/ISU develops and maintains the Alliance Information Systems (IS) that institutional and internal Alliance members use to enter and manage patient and study data. SMU/ISU also manages the Alliance website (http://www.allianceforclinicaltrialsinoncology.org), including the member site, and all Alliance databases. The website provides access to Alliance Web applications and other information useful to members, and is updated regularly as additional Alliance applications and reports are made available. The databases are the repository for member, patient, and study data.

In general, users of Alliance information systems are registered members – persons who assist with Alliance studies or other Alliance mission-related tasks. A primary objective of SMU/ISU is to provide efficient and reliable systems that enable the members to perform their assigned tasks, while safeguarding Sensitive Electronic Information (SEI) and Protected Health Information (PHI), and meeting the requirements defined by regulatory bodies including Health Insurance Portability and Accountability Act (HIPAA) and Health Information Technology for Economic and Clinical Health (HITECH).

9.1.1 Member account request and setup

All Alliance members must have a Cancer Therapy Evaluation Program (CTEP) ID and a CTEP Identity and Access Management (IAM) account in order to log into the member portion of the Alliance website. Refer to the CTEP website (http://ctep.cancer.gov) or to the Cancer Trial Support Unit (CTSU) website (http://www.ctsu.org) for additional information.
<table>
<thead>
<tr>
<th><strong>Policy Name:</strong></th>
<th>Member Information</th>
<th><strong>Policy Number:</strong></th>
<th>9.1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Section:</strong></td>
<td>Information Systems – 9</td>
<td><strong>Date Revised:</strong></td>
<td>January 1, 2018</td>
</tr>
</tbody>
</table>

Alliance member accounts give access to the Alliance member site (a restricted area of the Alliance website), and to Alliance IS Web applications. Prior to using the applications, Alliance members must be working with an institution that has IRB approval for an Alliance-based clinical trial, be authorized (as appropriate) to work with a given study’s clinical trial data, and receive required Web application training. Alliance Web applications are available to registered Alliance members only. However, users from other research groups may be given access to the Alliance website.

Additional member account setup is also required for user access to the individual legacy ACOSOG, CALGB, and NCCTG websites. Access to the ACOSOG, CALGB, and NCCTG websites is available using links on the Alliance website until the full transition of those functions and content is migrated wholly to the Alliance website. Access to non Web-based applications for staff members is provided by the ACOSOG, CALGB, and NCCTG organizations.

### 9.1.1.1 Individual institution members

To register a new member and request access to Alliance Web applications, an authorized institution representative must follow the application procedure specified on the Alliance website. During the application process, the prospective member’s role assignment(s) is specified. When the application is approved, appropriate accounts are created in the Alliance Information Systems. The member’s CTEP username and password is used to access the Alliance member site and SMU/ISU Web applications.

### 9.1.2 Institution registration

Alliance Institutional membership gives an institution the ability to participate in Alliance clinical trials. Institutional membership requirements and application instructions are available on the Alliance website under the ‘Membership’ heading.

### 9.1.3 Alliance application accounts

The Alliance uses Web-based and non Web-based applications for the capture, management, and reporting of clinical data for most Alliance-sponsored studies. As needed, users (who meet the above requirements) from other research groups may be given access to the Alliance website. For all Alliance applications access, an application must be completed and submitted to the administrator who issues access credentials.
9.1.4 User names and passwords

The Alliance requires each user to have a unique user name and password prior to accessing Alliance information systems that contain identifiable patient information. Sharing of accounts or passwords is prohibited. If the Alliance becomes aware of violations, it may be required to report the non-compliance to the offending institution’s security officers.

In addition to employing unique user names and passwords, each user must adhere to access restrictions for their accounts, guard their passwords, and change passwords regularly.

9.1.5 Roles and permissions

During Alliance registration, members are assigned roles and permissions that determine the specific Alliance data they may access, and which tasks they may perform.

A member may hold one role or many roles. Roles are defined as group roles, institution roles, committee roles, or study roles. A member holding a role is granted all of the data access privileges defined for the role. When a member holds more than one role, any necessary operation must be defined for access with at least one of the roles held by the member.

Typically, institutional members may access data from their own institutions only. Members from main member institutions can access data from their own institutions and their affiliate institutions. Alliance staff members may access only data necessary to fulfill their job responsibilities.

Members are further granted permissions, which are actions (e.g., read, update) that may be performed on the data they access.

Beyond assigned privileges and permissions, any privilege may be granted, with proper approval, to a specific member. Institutional members who need access to additional data should contact the Alliance Help Desk and request the additional privilege. Help Desk staff will forward the information to Alliance management for approval. Refer to the Alliance website under the ‘Contact’ heading for Help Desk contact information.

9.1.6 System availability

All Alliance systems are available 24 hours a day, seven days a week, with exceptions for system maintenance. Whenever possible, system maintenance will occur on a planned basis, with one week notice provided to Alliance
members. Unscheduled maintenance may occur as needed to resolve critical security vulnerabilities or to resolve other critical systems issues.

In the event of an unscheduled outage, an SMU/ISU employee will send a message to the established contact lists of users of Alliance systems. If network or internet connectivity problems occur such that users cannot access Alliance systems or send email, an SMU/ISU employee posts a message on the Alliance website under the ‘News’ heading.

9.1.7 User support

Alliance members require information systems that support all activities related to the conduct of clinical trials. To help meet these requirements, the Alliance provides technical support by trained Alliance Service Center employees to assist users with system, database, Web application, Internet, or study-related problems. Alliance Service Center employees may also create trouble tickets to document user issues prior to assignment to appropriate technical staff.

9.1.7.1 Alliance Service Center

For systems support, the Alliance Service Center is available Monday through Friday from 9 AM to 5:30 PM Eastern Time (8 AM to 4:30 PM Central Time). Refer to the Alliance website under the ‘Contact’ heading for the Alliance Service Center contact information.

For non-business hour emergency support, refer to phone numbers published in study protocols or memoranda, the Alliance website, and in the Alliance Service Center recorded off-hours message.
9.2 SMU/ISU operations

The remainder of this section contains policies the Alliance SMU/ISU uses to ensure the efficient and effective operation of its computing environment. The policies are divided into the following topics:

- Software development
- Documentation
- Technology selection and change management
- Usage of computing resources
- Security
- Backups and data retention
- Disaster recovery

Alliance IS staff locations adhere to the site-specific institutional IS policies of Mayo Clinic. In many cases SMU/ISU policies and procedures are more restrictive than institutional policies because of the national scope of the Alliance. However, all SMU/ISU policies and procedures serve the best interests of patients and members by providing the highest level of safety and security regarding data collection, maintenance, and reporting. Beyond safety and security criteria, the policies reflect the most efficient and effective means for meeting the goals of the Alliance and industry best practices.

9.2.1 Software development

SMU/ISU develops software applications and interfaces that generate, collect, maintain, and transmit data for clinical trials conducted by the Alliance. The applications are developed using a variety of development tools, technologies, and databases.

ISU uses a tiered software development environment to ensure proper testing and migration from the development to production environments. Software is first deployed to a development environment for initial testing by the software development staff. Software is subsequently deployed to an integration environment for software quality assurance and user acceptance testing, prior to being released into the production environment. New software is deployed during scheduled downtimes unless they are deemed urgent or critical, in which case the software release is migrated as soon as possible.

Completed deployment plans are required prior to implementing upgrades or other software changes in the production environment. Each release is planned to allow thorough testing prior to its deployment to the production environment.
Software developers are required to use development tools that have been carefully reviewed and established as standard within the ISU. Developers use online software development tools to capture project notes, requirements, technical specifications, screenshots, and links to appropriate source code repositories. Developers check all new and modified code into a source code control system that supports team development on various projects, prevents accidental file loss, allows backtracking to previous versions, and manages releases.

Security and confidentiality of study data are maintained at all times. To protect sensitive and confidential information, applications incorporate sound security practices and comply with HIPAA guidelines. All Alliance applications safeguard protected health information (PHI) by requiring secure logins, limiting access to authorized users, and implementing encryption schemes for data transmission.

9.2.2 Documentation policies

Documentation may include but is not limited to user manuals, job aids, training manuals, development documentation, policies, and standard operating procedures. All SMU/ISU documentation – whether in-process or released – is housed in a common server location.

During documentation development, writers follow consistent documentation templates. Documents intended for external (non-SMU/ISU) audiences are reviewed by the Training Team before they are finalized for publication. The reviewer list is dependent upon the document content.

9.2.3 Technology selection and change management

As resources permit, the Alliance works to maintain a state-of-the-art computing environment. Alliance developers use open-source software customized to Alliance needs, or software developed in-house. In some cases, commercial software solutions are a better choice, and are used if the vendor places a high priority on integration capabilities. The Alliance does not use commercial systems provided by a single vendor that would create a closed environment.

SMU/ISU projects are prioritized by the SDC Directors and monitored through Program Operations.
9.2.4 Usage of computing resources

9.2.4.1 Alliance staff and members

As part of its mission, the SMU/ISU acquires, develops, and maintains computers, databases, and networks. These computing resources are intended for Alliance-related purposes, including direct and indirect support of the Alliance mission.

Use of Alliance computing resources is not completely private. While the Alliance does not routinely monitor individual usage, normal operation and maintenance requires the backup of data and communications, the logging of activity, the monitoring of general usage patterns, and other activities necessary for the provision of service. Under prescribed circumstances, the Alliance may also specifically monitor the activity and accounts of individual Alliance computing resource users, including individual login sessions and the content of individual communications.

The Alliance does not permit use of its computing resources for personal, financial, or other gain.

9.2.4.2 Alliance staff

Alliance staff housed at institution locations must adhere to locally established usage requirements.

9.2.5 Security

SMU/ISU uses industry best practices to protect information against unauthorized access, use, or destruction. Access is controlled in order to limit the exposure of sensitive patient data.

Four categories of access control are implemented:

- Facilities
- Network and servers
- Database
- Application (includes Alliance website(s) and Web applications)

For all users, the SMU/ISU completes an Alliance authorization and account creation process before physical or electronic access is granted.
When a user no longer requires access, and authorization to terminate an account is received, SMU/ISU Help Desk employees terminate the user’s role(s) and disable accounts that provide access to the Alliance software applications.

### 9.2.5.1 Alliance Statistical Center facilities security

Permission to access to the Alliance Statistical Center facilities is determined by local institution guidelines.

The Alliance Data Center is a high security environmentally controlled and monitored computer room within the Statistical Center. Access to Alliance Data Center facilities is controlled through use of an Access Identification Card provided to permanent staff or vendors who have management authorization to be at the Data Center and have received training. Vendors may be approved for temporary or long-term access. Visitors that require access to the Data Center must receive management pre-approval and must sign in at the wall-mounted computer near the entrance to the facilities. In addition, visitors must be escorted during their visit by a cardholder with Data Center access authorization.

The Alliance monitors for and protects its computer resources against environmental hazards. Systems are centrally monitored, kept in temperature-controlled conditions, and are protected against electrical power surges and short-term outages. Backup generators are available onsite to ensure the continuous operation of the Data Center in case of long-term utility power failures. To comply with local building and fire codes, computer resources are protected by automatic smoke detection and fire suppression equipment.

### 9.2.5.2 Network and server security

SMU/ISU passwords and network/server security upgrades must be managed to conform to local institution practice.

Alliance systems housed at the Statistical Center are protected by enterprise firewalls and network security, which provides continuous monitoring to identify and prevent malicious access.

Server login passwords are encrypted and stored in their encrypted form in protected files.
An administrative user account is a specific account type that allows access for system administration purposes, including setup of user accounts. Administrative users only are authorized to manage accounts and servers. An Alliance computing manager responsible for specific work units designates Alliance SMU/ISU staff administrative users and assigns them a unique user ID and password.

### 9.2.5.3 Database security

Because of the highly sensitive nature of data collected by the Alliance, and the right to privacy of patients entered on clinical trials, only authorized members with a need to know will be given access to data in the Alliance databases. SMU/ISU ensures the security and integrity of the databases through password controls, logging, monitoring, and auditing.

SMU/ISU implements database auditing for relevant features related to data definition, security administration, and logon failures. The SMU/ISU implements both database and application level auditing for relevant features related to data manipulation, security administration, and logon failures. A database audit trail is used to record date, time, and user for various levels of standard and suspicious activity. A correction history is available to record date, time, and user for all data manipulation activity.

SMU/ISU monitors each database product software lifecycle, and ensures that appropriate updates are applied. Each new release and version of the database software is identified, considered for installation, and installed after rigorous validation.

For database patches, SMU/ISU follows industry best practices. Database security patches are installed only after they are validated against the SMU/ISU computing environment. If validation is successful, installation will occur as soon as possible after the date of release.

Database user accounts are set up after authorization by the appropriate manager. Database passwords expire at preset intervals per institution standards and must be changed when required. SMU/ISU employees inactivate or remove user accounts immediately upon notification of termination of employment or Alliance membership.
9.2.5.4 Application security

SMU/ISU develops and/or supports software applications for use by Alliance members. These applications enable such functions as patient registration, specimen tracking, reporting, and data entry and review. ISU applications encrypt all data transmission to ensure security and confidentiality of data as it is entered and viewed.

Authentication and authorization services ensure consistent security for applications. Users are provided accounts and roles that determine their access, at a granular level, to data and functions. These roles are periodically reviewed by SMU/ISU directors, with final approval for changes given by the Alliance IT Committee.

9.2.6 Backups and data retention

All system and database backup and recovery procedures adhere to industry best practices. Alliance data is safeguarded against loss via industry standard backup and retention schedules. Backups are performed on a daily basis. Using the backup scheduler and policy engine, data backups are targeted to a tape library located in a remote data center physically separated from the primary infrastructure hosting Alliance application and data services. Data are retained for a period not less than 30 days. Alliance employee workstations are managed by local desktop support and fall under the backup policies of the support environment.

9.2.6.1 System and database backups

Controlled and monitored backup rotations protect all servers, file storage devices, and server security information.

9.2.6.2 Servers

Servers are designated by institution policies as critical or non-critical. All servers receive a weekly full backup, and incremental backups. In the event of file loss, file corruption, or total equipment loss, SMU/ISU is able to recover from the previous full and incremental backups. Maximum file loss would be 24 hours.

9.2.6.3 Retention and storage

Backup tapes are retained for two months. Longer data retention is additionally determined by the study. Security access files for all
machines supported by the SMU/ISU are backed up and retained as required by HIPAA.

Backups are stored offsite from the main data center. Statistical archives are stored in SAS data sets rather than in the Alliance database and are housed on a separate server.

9.2.7 Disaster recovery

The Alliance has a formal disaster recovery plan to be used in the event of a significant failure of regular computing services. The plan identifies the primary and backup members of the disaster recovery assessment team and the functional systems area for which each person is responsible. If an event occurs that requires the attention of the team, all members assemble to begin an assessment of the situation for their respective area and prepare an estimate of the time and level of effort required to restore operations. Restoration efforts are directed by the ISU leadership. Recovery time will depend on the nature of the disaster.
10 Publications Committee charter and mission guidelines

“The Publications Committee shall review existing policies and best practices concerning authorship of scientific publications, and shall recommend to the Executive Committee for its approval a set of requirements for authorship of Alliance publications. These requirements shall be in the form of a guidance policy for Alliance publications and shall address rules governing authorship and disclosure of conflict of interest for Alliance publications. The chair and vice chair of the Publications Committee shall include one individual who is a scientific leader and one who is a community oncology leader. The Publication Committee shall include representatives from the Central Protocol Operations Program and the Statistics and Data Management Program, as well as other members as deemed appropriate. The Publications Committee shall meet at a frequency of not less than once yearly. The Publications Committee shall also adjudicate in a timely manner any issues related to publication of Alliance manuscripts, and make recommendations concerning these matters to be acted upon by the Executive Committee.”

— Statement from the Alliance Constitutions and Bylaws

10.1 Data ownership

Data generated by Alliance Group activity, using Alliance resources, or associated with the Alliance belong to the Alliance. Therefore, the Alliance, through its publication policy, has oversight over the use and publication of any and all Group data. All planned abstracts or manuscripts reporting results of Alliance studies to a meeting or journal for publication are to undergo pre-submission review and approval, based on this Policy and Procedures document.
10.2 Committee members

Members of the Alliance Publications Committee are nominated by the committee chair to serve 3-year terms (renewable one time), and are expected to attend a minimum of 75 per cent of committee meetings.
### 10.3 Group Review members

<table>
<thead>
<tr>
<th>Reviewer’s Group Role</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>All co-authors of publication</td>
<td></td>
</tr>
<tr>
<td>Chair, Publications Committee*</td>
<td></td>
</tr>
<tr>
<td>Chief Financial Officer</td>
<td></td>
</tr>
<tr>
<td>Committee Chair</td>
<td>Applicable studies only</td>
</tr>
<tr>
<td>Director, Biospecimens and Correlative Science Operations*</td>
<td>Translational studies only</td>
</tr>
<tr>
<td>Director, Central Operations*</td>
<td></td>
</tr>
<tr>
<td>Director, Regulatory Affairs</td>
<td></td>
</tr>
<tr>
<td>Executive Officer</td>
<td>Applicable studies only</td>
</tr>
<tr>
<td>Group Administrator</td>
<td></td>
</tr>
<tr>
<td>Group Chair*</td>
<td></td>
</tr>
<tr>
<td>Group Statistician*</td>
<td></td>
</tr>
<tr>
<td>Manager, Publications Operations</td>
<td></td>
</tr>
<tr>
<td>NCI representative</td>
<td></td>
</tr>
<tr>
<td>Industry representative, according to study agreement</td>
<td>Applicable studies only†</td>
</tr>
<tr>
<td>Executive Committee members</td>
<td>Half of the EC membership (excluding those asterisked in this table) is selected to review publications in 6-month rotations</td>
</tr>
</tbody>
</table>

*Member of the Executive Committee who reviews publications in all rotations.
†Determined by Director of Regulatory Affairs
10.4 Abstract and manuscript preparation

10.4.1 General principles


The study chair is responsible for providing leadership and writing manuscripts/abstracts for publications that describe an Alliance study. The document entitled “CHECKLIST – Recommended Content for Alliance Manuscripts and Meeting Abstracts” provides guidance related to title page, authorship, acknowledgements, scientific content for different sections, as well as template wording for support, monitoring, informed consent, locations of data collection and statistical analyses, randomization scheme, quality assurance, meta- or pooled analysis, and data lock. All authors are expected to review and follow this checklist.

The study chair sends the initial draft manuscript/abstract to all the co-authors for review, including the faculty and staff statisticians. All authors, including those assigned authorship based on accrual, are responsible for careful and meaningful review. The first author takes into account all comments and suggestions by co-authors and incorporates them into the revised draft, as appropriate. After initial co-author review, the study chair sends the revised draft to the publications coordinator (publications@AllianceNCTN.org) as an MS Word file; this way the Alliance files are properly up to date. This revised draft is sent for Group Review (see sections 10.5.3 and 10.5.4).

It is the responsibility of the corresponding author to collect and send to the journal all journal-specific conflict of interest forms prior to manuscript submission for publication. Any individual with a conflict of interest that is sufficient to make them ineligible for a study chair role cannot serve as either first or senior (last) author of an Alliance publication.

10.4.2 Cover page

It is important for the study number(s) to appear early in the manuscript/abstract for ease of retrieval in literature searches. The title section of the cover page of the manuscript should indicate the Alliance or legacy study number(s) about which the manuscript is written. As example: “Phase III Alliance A1K study of drug A vs. drug B for treatment of X”. For abstracts and manuscripts generated from the ACOSOG, CALGB, and NCCTG legacy groups, recommendation is to add “Alliance” after the study number. As example: “Phase III ACOSOG A1K (Alliance) study of drug A vs. drug B for treatment of X”.

Alliance Policies and Procedures - 10-4
If it is not possible to include all study numbers in the title, the author should insert wording such as “A combined analysis of Alliance studies” in the title; include the study numbers within the abstract or introduction section.

Each cover page of a manuscript also indicates the supporting grant numbers for all authors listed. This is done by use of a footnote after each author's name, with the footnote itself containing the name and location of the main member institution in which the author was affiliated when the study was activated, followed by the National Institutes of Health (NIH) grant number. Appropriate acknowledgment of other funding sources should be included as well (e.g., the Breast Cancer Research Foundation or company XYZ).

10.4.3 Authorship

Alliance authorship guidelines follow those of the publicly available International Committee of Medical Journal Editors (ICMJE) recommendations for authorship:

“The ICMJE recommends that authorship be based on the following 4 criteria:

• Substantial contributions to the conception or design of the work; or the acquisition, analysis, or interpretation of data for the work; AND

• Drafting the work or revising it critically for important intellectual content; AND

• Final approval of the version to be published; AND

• Agreement to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

All those designated as authors should meet all four criteria for authorship, and all who meet the four criteria should be identified as authors. Those who do not meet all four criteria should be acknowledged.”

If there are questions or discrepancies related to author order based on the study chair’s decision and the publications guidelines, as seen below, arbitration is required by the Alliance Publications Committee chair and the Alliance Group chair, with input from the other Group Review members.

10.4.3.1 Publication on the primary study endpoint

The listing and order of authorship for a manuscript/abstract for a primary study endpoint is determined by overall workload contribution, intellectual...
contribution, and participant accrual. Each author is responsible for obtaining any required clearances from his/her own institution (or network).

The first author of the manuscript/abstract is usually the study chair or co-chair. A study chair who moves to a non-Alliance institution may continue to serve in the full capacity of study chair with the agreement of the appropriate committee chair and if no conflicts of interest have arisen because of the move of the study chair. The original study chair therefore retains authorship rights by virtue of serving in the full capacity of the study chair role.

The first author is generally followed by the study’s primary statistician. Authorship should be granted to the responsible executive officer. The study community co-chair should be included as an author if appropriate by ICMJE recommendations stated above. If the modality co-chair participated in the design of the study and wrote the modality section of the protocol, they should be an author on primary endpoint publications. Pathologists, radiologists and other specialists who perform quality assurance (QA) for a study should be included in the authorship of any publications that result from the study, unless the publication is independent of QA results of their findings. The decision for inclusion of an Alliance quality assurance specialist/data manager, clinical research professional or nurse as a co-author is to be made by the study chair in consultation with the primary statistician and disease/modality committee chair, and must be made according to ICMJE recommendations.

Other individuals making significant contributions according to ICMJE recommendations may be listed.

Institutional authorship based on accrual is separate from (and in addition to) study chair, committee chair or other contributors. Institutional authorship representation on primary study publications is awarded to an institutional network whose participant accrual contribution fulfills the following guidelines:

<table>
<thead>
<tr>
<th>Total number of participants in the study</th>
<th>Number of participants at a network, based on total study accrual</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fewer than 100 total study accrual</td>
<td>25% of the total or 8 participants, whichever is less</td>
</tr>
</tbody>
</table>

*Alliance Policies and Procedures* - 10-6
The principal investigator of a network makes the assignment of authorship after being informed by the publications operations manager or publications coordinator of institutional merit. The network principal investigator is best suited to determine the assignment of authorship and may assign himself/herself, another physician in the same or another specialty, or an individual from the main member or an affiliate. In most cases, authorship is assigned to the highest accruing investigator in the institutional network. Institutional nurses or clinical research professionals making significant contributions should also be considered for authorship. Generally, the individual given the authorship assignment should be someone who was working at the institution during the period of accrual and who made substantive contributions to accrual at the institution. All authors should be included in manuscript preparation and approval.

For manuscripts/abstracts involving other National Clinical Trial Network (NCTN) group studies, it is not necessary to include all other NCTN group institutions, but it is expected that groups that endorsed the study and enrolled >10% of patients should have at least one author included in the report of treatment studies.

All primary manuscripts (excluding those for multi-group studies) also acknowledge each institution that enrolled participants on the study, as an appendix. The relevant local principal investigator, their institution, and grant numbers are listed in that appendix.

When the study is a limited access pilot of fewer than 30 patients, involving only a few institutions, the study chair, primary statistician and committee
10.0 Policy Name: Authorship

10.4 Section: Publications – 10.0

Policy Number: 10.4

Date Revised: January 1, 2018

chairs should discuss authorship. Ideally, all institutions participating will be represented.

10.4.3.2 Publication on a secondary (correlative) study

A secondary (correlative) study may include observations utilizing existing datasets or compilation of results from several studies. The secondary study may have been approved as a sub-study in an original protocol document, or may be a new study that was proposed by an Alliance or non-Alliance investigator. The work may involve biospecimens, quality of life, symptom analyses, and economic analyses, among others. The intention of the Alliance authorship policy is to be appropriately inclusive, consistent with authorship guidelines from major journals and the ICMJE.

Information related to the Alliance and its grant numbers should be in the face page of secondary manuscripts.

1. **Authorship on publications of a secondary study included in the original Alliance or legacy protocol**

All of the following are invited to participate in review of abstract/manuscript data, publication development and approval and should receive authorship if appropriate by ICMJE recommendations:

- Study chair, study co-chair, executive officer, and community co-chair of the original study. Authorship by a modality co-chair on secondary endpoint publications should be a function of their involvement in the secondary analysis.
- Study chairs from other cooperative groups that accrued patients or samples to the secondary study
- Correlative study statistician and primary statistician of the original study if different
- Pathologists, radiologists and other specialists who perform quality assurance (QA) for the study, unless the publication is independent of QA results of their findings.
- Accrual authors

   For accrual authors on CALGB and NCCTG publications, the principal investigator of the highest accruing network selects the network author based on investigator accrual or other study contribution. No minimum accrual threshold is required for the network or selected author.
2. **Authorship on publications of a secondary study not in an original Alliance or legacy protocol; study proposed by Alliance investigator**

New secondary studies include observations utilizing existing datasets or specimens, or a compilation of results thereof from several studies that were not part of the original objectives of the primary study or studies.

a. When manuscripts/abstracts are prepared for new secondary studies, potential authorship should be extended to the following, but final authorship determination should be based on ICMJE recommendations:

- Study chair(s) of original Alliance study or studies, correlative study statistician, primary statistician of original Alliance study or studies. Co-chairs from other cooperative groups that accrued any patients or specimens may be included if Alliance author or Alliance committee chair requests.
- Pathologists, radiologists and other specialists who perform quality assurance (QA) for a study, unless the publication is independent of QA results of their findings.
- Researchers performing the secondary study

After primary study chair(s), primary statistician(s), QA specialists and researchers, other investigators who were involved in the primary study or studies may not necessarily be included in secondary study publications; instead, authorship is determined by an individual’s contribution specific to the secondary study and by ICMJE recommendations. Order of authorship should reflect the magnitude and effort contributed by each author to the secondary analyses, which may be independent of the primary studies’ analyses or accrual.

b. Authorship based solely on accrual is not a criterion for this category of abstract or manuscript. Accrual investigators are recognized in an acknowledgement section rather than with authorship, unless they are among the investigators.
conducting the secondary use study, in which case authorship depends upon contribution. It is expected that all investigators who contributed data to the secondary analyses will also

- be involved in interpretation of those data
- be given the opportunity to participate fully in preparation of resultant manuscripts/abstracts
- be acknowledged as co-authors on those manuscripts/abstracts.

This may also apply to non-tissue secondary abstracts/manuscripts if the data collected by the investigators from the collaborative groups will be utilized.

3. **Authorship on publications of a secondary study not in an original Alliance or legacy protocol; study proposed by non-Alliance investigator**

This category includes abstracts and manuscripts led by outside investigators who have been granted access to Alliance data or biospecimens.

Authorship decisions regarding the non-Alliance correlative study chair and statistician and non-Alliance researchers performing the secondary study are made by the non-Alliance investigator and team.

NCI rules do not mandate that the Alliance investigators be considered for authorship. We suggest that outside investigators consider including the following Alliance leadership team in the preparation and formal approval of the manuscript:

- Alliance study chair(s), of original Alliance study or studies
- Alliance primary statistician(s) of original Alliance study or studies
- Investigators who contributed annotated tumor specimens
10.5 Abstract and manuscript timelines

10.5.1 Timelines for abstract and manuscript preparation

The process of abstract and manuscript generation for phase III studies begins promptly after the Alliance Data and Safety Monitoring Board (DSMB) has determined that the study results may be released and the study chair has completed case evaluations. For phase II studies, the process begins when the study chair has received the study summary from the study’s primary statistician. Of note, the statistician may need to conduct additional analyses in collaboration with the study team. Once the statistical analyses are completed, the statistician sends a copy of the analyses to the study chair and notifies the disease/modality chair (refer to the Statistical Summary Report Timelines Document).

The first abstract/manuscript is expected to be based on the mature primary endpoint of the study. Submission of abstracts before data on the primary endpoint are completed is not generally endorsed, but may be considered on individual cases. Some examples are description of unexpected toxicities, enrollment procedures or data, and companion studies that are not dependent on the primary endpoint. This decision to submit an abstract before primary endpoint data are mature is made as a collaborative effort between the study chair, study primary statistician, committee chair, Group chair, and Publications Committee.

Almost all abstracts submitted to a meeting must be followed by a full manuscript (except in special situations that should be discussed with the Alliance Publications coordinator prior to the abstract submission); the manuscript should be sent to the Alliance publications coordinator (publications@AllianceNCTN.org) for Group Review no later than 6 months after the meeting. We suggest that the abstract author create a draft manuscript by the time of meeting presentation using the statistical analysis that is prepared for the meeting abstract to optimize time and effort. This initial draft can be used as a guide from which to develop a final version that is sent to potential co-authors, etc., prior to submission to the Alliance publications coordinator.

For publications in which an abstract is not prepared prior to developing a draft manuscript, the draft manuscript should be sent to the publications coordinator within 2 months from completion of the statistical summary report.

10.5.2 Delinquency in manuscript preparation

As stated above, it is expected that a draft manuscript is completed at the time of data presentation at a medical meeting. When a study chair has not completed a draft manuscript according to this timeline, the disease or modality committee chair
initiates a discussion with the study chair, as a warning (cc to publications@AllianceNCTN.org). After receiving a warning notice from the committee chair, the study chair has 30 days to submit a first draft of the manuscript to the protocol office.

If the study chair is unable to complete the manuscript in the expected time period, 2 actions by the disease and modality committee chairs may follow: (1) reassignment of first authorship and (2) prevention of the delinquent author from chairing a future Alliance concept or study for at least one year. The appropriate disease and modality committee chairs then request from the Group chair (and Publications Committee chair) permission to reassign the manuscript to an investigator responsible for a large percentage of accrual or with a substantial intellectual contribution to the study. The reassignment of authorship of a paper rests with the appropriate disease or modality chairs, who should in turn notify both the new author and the study’s executive officer of the reassignment. The disease or modality chair should clarify to the new author that the first draft of the manuscript should be ready within 30 days after reassignment.

10.5.3 Timelines for review and revision of abstracts submitted to the Alliance publications coordinator

A meeting abstract must be submitted by the first or corresponding author to the publications coordinator (publications@AllianceNCTN.org) as a Word document at least 2 weeks prior to the meeting abstract submission deadline. The author receives scientific comments from Group reviewers typically within 2 days. Comments concerning authorship may also be sent to the corresponding author. After revising the abstract based on Group Review, the first author must send the revised abstract to co-authors for their approval. When the abstract is accepted, the author must send the acceptance email and the final submitted abstract to all co-authors and to the publications coordinator within 1 week after acceptance.

10.5.4 Timelines for review and revision of manuscripts submitted to the Alliance publications coordinator

The publications coordinator (publications@AllianceNCTN.org) reviews authorship within 2 working days and submits the authorship to the study chair within those 2 working days. Barring any discrepancies or concerns between the study chair and publications coordinator’s list and order, the publications coordinator submits the manuscript for Group Review within 2 working days. The Alliance manuscript review (aka Group Review) members are described in the Group Review section above.
Reviewers are expected to provide written input to the publications coordinator within 7 working days. All abstracts and manuscripts (except those resulting from data sharing) must be reviewed by an independent Alliance faculty statistician.

All comments from the Group Review should be sent to the manuscript’s first author, the corresponding author, the chair of the Publications Committee, and the publications operations manager. The first author is expected to discuss suggestions with the study statistician, review comments, and complete a second version of the manuscript within 4 weeks. Inability to meet this timeline should be discussed with the modality/disease committee chair. Based on the situation, further discussion with the Publications Committee chair may be required, to better assist the author.

10.5.5 Approval of abstracts and manuscripts

All comments received from reviewers during Group Review are sent to the chair(s) of the Alliance Publications Committee. The Publications Committee chair(s) are responsible for approving abstracts and manuscripts, or requesting revisions followed by re-review.
### 10.6 Abstract or manuscript submission to meeting or journal

The study chair revises the manuscript/abstract based on internal and external reviews outlined above and sends the co-authors the revised publication for their approval. The author submits the approved manuscript/abstract to the journal or association for review, complying with all submission requirements. The study chair also sends a copy of the submitted manuscript/abstract to the publications coordinator for inclusion in the Alliance publication files within 1 week after submission.
### 10.7 Publication of abstract or manuscript

The study chair/corresponding author advises the publications coordinator (publications@AllianceNCTN.org) of the status of all abstracts and manuscripts submitted to a meeting or journal for publication. Letters of acceptance and a PDF file of the published abstract or printed manuscript must be sent by the study chair/corresponding author to the publications coordinator within 14 days of availability. This is necessary for the Alliance publication files to be accurate and complete (including the full citation). This material is reviewed every 3 months by the Publications Committee. To facilitate access to Group study results, Alliance publication citations are posted in the publications section on the Alliance Web site.
10.8 Publicizing Research Information

All communication related to the dissemination of Alliance research to external audiences is handled by the Alliance communications specialist. This includes all written or recorded communication (i.e., press releases, news releases, press statements, video releases) directed to members of the news media, stakeholders, and the public, regarding the activation, progress, results and findings of Alliance research. This also relates to all communication generated by an institution or industry partner based on Alliance research. Such communication must be submitted the communication specialist (communications@AllianceNCTN.org) for review at least one week prior to its release. Also refer to Section 14.3, Dissemination of Information to the General Public.
10.9 Summary of study results for the public

The lead author must submit the completed plain language study results summary template to the publications coordinator (publications@AllianceNCTN.org) when the manuscript is sent for Alliance Group review. If a manuscript is not accompanied by a completed template, Group review will be delayed until its receipt.

For a phase III or randomized phase II study, a public study result summary of the trial design, goals and results is created by the Publications Committee, with input from the lead author of the manuscript, Patient Advocate Committee and Oncology Nursing Committee, using the plain language template for consistent and understandable information. The primary audience for public study result summaries includes study participants.

The Alliance web content administrator posts the public summary to the Alliance website at a time that coincides with publication of the manuscript.
10.10 NIH Public Access Policy compliance

The U.S. government provides full-text content of scientific journal literature to the public through PubMed Central. All peer-reviewed journal articles resulting from Alliance NIH funding that are accepted for publication on or after April 7, 2008 must appear in PubMed Central no later than 12 months after the official publication date, according to NIH Public Access Policy NOT-OD-08-033. In PubMed Central, they may appear as either accepted final peer-reviewed manuscripts or final published articles (see definitions in table 1 below). Failure to comply may result in withholding of federal funds to the Alliance.

The level of author involvement in compliance depends upon the journal in which the manuscript is published. Most journals assist the author in submitting a journal article for use by PubMed Central (formatted as either the accepted peer-reviewed manuscript or final published article). In rare situations, the author may be entirely responsible for completing this process (e.g., manuscripts published in Leukemia & Lymphoma). Some journals provide assistance options that must be selected by the author at the time of manuscript submission. Therefore, at the time of manuscript submission, the first author should consult with the journal or visit the journal Web site to determine the journal’s method to assure compliance with the NIH Public Access Policy.

The table below provides a summary of the document submission methods (methods A, B, C and D, as described by NIH) and document approval steps required by NIH for compliance. It also indicates, by method, the responsible parties and journals that frequently published Alliance manuscripts. The table is based on NIH training, and was developed by the Alliance to consolidate instructions for authors. In summary, the author need not submit or approve the document when publishing in a journal that uses Method A or B, although most publishers that offer Method B charge an extra fee. The author need only provide approvals when a journal uses Method D. The author is responsible for both manuscript submission and approvals when a journal uses Method C. Under Alliance policy, the author must ensure that all steps are taken to comply with NIH requirements.

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
<th>Responsible Parties</th>
<th>Journals</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Automatically deposited by publisher</td>
<td>Author</td>
<td>Leukemia &amp; Lymphoma</td>
</tr>
<tr>
<td>B</td>
<td>Author provides final document</td>
<td>Author</td>
<td>Leukemia &amp; Lymphoma</td>
</tr>
<tr>
<td>C</td>
<td>Author provides final document and approves</td>
<td>Author</td>
<td>Leukemia &amp; Lymphoma</td>
</tr>
<tr>
<td>D</td>
<td>Author provides final document, approves, and explicitly agrees to NIH policy</td>
<td>Author</td>
<td>Leukemia &amp; Lymphoma</td>
</tr>
</tbody>
</table>
Submission methods, process steps and responsible parties for compliance with NIH Public Access Policy*

<table>
<thead>
<tr>
<th>Steps in Process</th>
<th>Submission Method Used by Journals and Responsible Party</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Step 1:</strong> Submitting the file ‡</td>
<td>If journal uses <strong>Submission Method A:</strong> Publisher posts published article to PMC no later than 12 months after publication.</td>
</tr>
<tr>
<td></td>
<td>If journal uses <strong>Submission Method B:</strong> Same as Method A, except author must opt for publisher assistance (usually for a fee).</td>
</tr>
<tr>
<td></td>
<td>If journal uses <strong>Submission Method C:</strong> Upon manuscript acceptance, author submits manuscript via NIHMS system. ‡</td>
</tr>
<tr>
<td></td>
<td>If journal uses <strong>Submission Method D:</strong> Upon manuscript acceptance, publisher submits manuscript via NIHMS system. ‡</td>
</tr>
<tr>
<td><strong>Step 2:</strong> Approving submitted materials (Required step after submitting the file) ‡</td>
<td>Publisher approves</td>
</tr>
<tr>
<td></td>
<td>Publisher approves</td>
</tr>
<tr>
<td></td>
<td>Author approves via NIHMS, after notification from NIHMS of action required. ‡</td>
</tr>
<tr>
<td><strong>Step 3:</strong> Approving PMC web version (Required step after NIHMS or PMC creates web version)</td>
<td>Publisher approves</td>
</tr>
<tr>
<td></td>
<td>Publisher approves</td>
</tr>
<tr>
<td></td>
<td>Author approves via NIHMS, after notification from NIHMS of action required.</td>
</tr>
<tr>
<td></td>
<td>Author approves via NIHMS, after notification from NIHMS of action required.</td>
</tr>
</tbody>
</table>

* Based on information available at https://publicaccess.nih.gov as of July 1, 2014.
† **Final published article:** journal’s authoritative copy of the paper, including all modifications from publishing peer review process, copy editing/style edits, formatting. **Final accepted peer-reviewed manuscript:** author’s final manuscript of peer-reviewed paper accepted for publication, including all modifications from the peer review process. Only one version of paper must be submitted.
‡ **NIHMS:** the National Institutes of Medicine Manuscript Submission System.
€ For Methods C and D, steps 1 and 2 must be completed within 90 days after article’s official date of publication in order to be compliant with NIH public access policy. This is to allow completion of processing steps and PMC posting by 12 months after publication.

Training on an author's responsibilities in complying with the NIH Public Access Policy is provided at http://publicaccess.nih.gov/communications.htm and http://www.nihms.nih.gov/help/#slideshow. Answers to frequently asked questions are available at NIHMS FAQ. To ask questions about the process of compliance with the NIH Public Access Policy, authors should contact the NIHMS or PubMed Central help desks using the following URLs:

- NIH Public Access: PublicAccess@nih.gov
- NIHMS: https://nihms.nih.gov/db/sub.cgi?page=email&from=grant_suggest&mid=

Alliance Policies and Procedures - 10-19
## 10.11 Quick view of Alliance publication timelines

<table>
<thead>
<tr>
<th>Type of publication</th>
<th>Timelines</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Meeting abstract</strong></td>
<td>Send to publications coordinator:</td>
</tr>
<tr>
<td></td>
<td>2 weeks prior to meeting submission deadline or per online schedule</td>
</tr>
<tr>
<td></td>
<td>2 days for scientific review</td>
</tr>
<tr>
<td></td>
<td>7 days for operations review during high volume</td>
</tr>
<tr>
<td></td>
<td>Send to publications coordinator:</td>
</tr>
<tr>
<td></td>
<td>1. Copy of submitted abstract within 1 week after submission</td>
</tr>
<tr>
<td></td>
<td>2. Acceptance email and PDF of published abstract no later than 2 weeks after available</td>
</tr>
<tr>
<td><strong>Manuscript with no prior meeting abstract</strong></td>
<td>Send to publications coordinator:</td>
</tr>
<tr>
<td></td>
<td>2 months after completion of the statistical summary report along with completed public study summary template, if applicable</td>
</tr>
<tr>
<td></td>
<td>7 days for scientific review</td>
</tr>
<tr>
<td><strong>Manuscript that follows a meeting abstract</strong></td>
<td>Send to publications coordinator:</td>
</tr>
<tr>
<td></td>
<td>6 months after presentation at meeting along with completed public study summary template, if applicable</td>
</tr>
<tr>
<td></td>
<td>7 days for scientific review</td>
</tr>
<tr>
<td><strong>Alliance-approved manuscript submitted to journal</strong></td>
<td>Journal submission:</td>
</tr>
<tr>
<td></td>
<td>Determine the journal’s NIH Public Access Policy method to assure compliance with government policy if manuscript is accepted</td>
</tr>
<tr>
<td></td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td>NA</td>
</tr>
<tr>
<td><strong>Accepted manuscript</strong></td>
<td>Manuscript acceptance:</td>
</tr>
<tr>
<td></td>
<td>Submit to NIHMS for use by PubMed Central if journal does not assist; respond to NIHMS requests for approval</td>
</tr>
<tr>
<td></td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td>If submission Method C or D was used, provide in NIHMS:</td>
</tr>
<tr>
<td></td>
<td>Approval of submitted or posted materials</td>
</tr>
<tr>
<td></td>
<td>Approval of PMC web version</td>
</tr>
<tr>
<td><strong>External study communications, if applicable</strong></td>
<td>Send to publications coordinator and communications specialist:</td>
</tr>
<tr>
<td></td>
<td>1 week</td>
</tr>
<tr>
<td></td>
<td>NA</td>
</tr>
</tbody>
</table>

### Contact information
- Alliance publications coordinator: [publications@AllianceNCTN.org](mailto:publications@AllianceNCTN.org)
- Alliance communications specialist: [communications@AllianceNCTN.org](mailto:communications@AllianceNCTN.org)
- NIHMS: [https://nihms.nih.gov/db/sub.cgi?page=email&from=grant_suggest&mid=](https://nihms.nih.gov/db/sub.cgi?page=email&from=grant_suggest&mid=)
11 Alliance Biorepositories and Biospecimen Resource (ABBR) and Translational Research

11.1 ABBR Infrastructure and Oversight

11.1.1 The Alliance ABBR is comprised of five federated biorepository facilities located at four academic medical centers.

11.1.1.1 Alliance Biorepository at the Ohio State University (OSU). Formerly known as the “CALGB PCO”, this facility stores primarily fixed tissue and biofluids from legacy, CALGB solid tumor and lymphoma studies, as well as solid tumor and biofluid biospecimens from newer Alliance studies.

11.1.1.2 Alliance Hematological Malignancy Biorepository (HEME). Formerly known as the “CALGB Leukemia Bank”, this facility also resides at The Ohio State University and stores specimens from patients with acute or chronic leukemia, myelodysplastic syndrome, or multiple myeloma who are enrolled on an Alliance protocol. HEME primarily receives blood and bone marrow specimens, and, in some cases, buccal smears.

11.1.1.3 Alliance Lung Cancer Tissue Bank (LCTB). The Alliance Lung Cancer Tissue Bank (LCTB) is located at the Brigham and Women’s Hospital in Boston, MA. The purpose of the LCTB is to collect, catalog and store frozen samples of lung carcinoma and when possible, portions of involved lymph nodes and adjacent uninvolved lung tissue obtained from previously untreated patients. In addition to tissue specimens, blood samples are also collected pre- and post-resection from the patients to provide a source of quality DNA, RNA and protein for molecular studies.

11.1.1.4 Alliance Biorepository at Washington University in St. Louis (WUSTL). Formerly known as the “ACOSOG Specimen Bank” this CAP-accredited facility collects and stores frozen and fixed tissue, and biofluids from breast, lung, GI, and other solid tumor Alliance trials.

11.1.1.5 Alliance Biorepository at Mayo Clinic (MAYO). Formerly known as the “NCCTG Biospecimen Resource”, this second CAP-accredited facility processes and stores biospecimens associated with neuro-oncology studies, and is also the designated repository.
11.1.2 Biospecimen tracking, reporting, and inventory management is integrated across all biorepository sites and centrally coordinated at the WUSTL biorepository, through the use of the Alliance BioMS biospecimen management tool.

11.1.3 Although each biorepository site maintains its own local set of policies and standard operating procedures to comply with institutional requirements, those individual site policies specifically pertaining to Alliance trial biospecimen integrity and management are harmonious and meet the minimal standards set forth in this document.

11.1.4 The ABBR is supported by a National Cancer Institute (NCI) U24 funding mechanism. Each of the Alliance biorepository leaders at the four academic institutions serve as a co-Principal Investigator (PI) on the U24 grant, with the WUSTL bank director currently serving as contact PI.

11.1.5 One or more of the ABBR U24 grant PIs also serves on the Alliance Translational Research Program (TRP) Executive Committee and the Alliance Executive Committee. These appointees are charged with ensuring that the ABBR serves the needs of the NCTN Alliance network.

11.1.6 Three of the ABBR U24 grant PIs (or their designees) also serve on the NCTN Group Banking Steering Committee (GBC). The GBC is charged with developing and adopting harmonized policies and practices across all NCTN biospecimen resources.

11.1.7 The TRP Executive Committee is primarily responsible for oversight of compliance of the Alliance repositories with Alliance and NCI policies regarding specimen collection and distribution. In addition, this committee is responsible for ensuring that the repositories follow the NCI guidance document “Best Practices for Biospecimen Resources” that was published and updated in 2011. Each Alliance biorepository site will undergo periodic audits to ensure compliance with the NCI Best Practices (http://biospecimens.cancer.gov/practices) and oversight for the audits will be a function of the TRP Executive Committee.

11.1.8 The Alliance Translational Research Program (TRP), the TRP biorepository sub-committee, study chairs and correlative science co-chairs, individual disease/modality/discipline committees (usually the vice-chair of the
disease/modality/discipline in charge of translational research) are jointly responsible for: (1) determining biospecimens that should be collected on each Alliance trial and the appropriate methods for collection and processing of those biospecimens and (2) ensuring that the ABBR sites have the appropriate quality control and quality assurance procedures in place for biospecimen handling, processing, storage and distribution.

11.1.9 As the Alliance steward of biospecimens, each biorepository director agrees to procure, store, process and distribute the specimens according to Alliance and NCI policy. In addition, if the biorepository does not comply with Alliance policy, the Alliance can move the biospecimens to another approved Alliance location.
11.2 Biorepository Functions

The ABBR serves a number of important functions in the context of NCTN Alliance clinical trials. These roles include, but are not limited to:

11.2.1 Biospecimen Collection. The ABBR may design, construct, and distribute supplies and ‘kits’ to facilitate biospecimen collection from remote sites. It is the responsibility of the ABBR to ensure that the design of such materials maintain biospecimen integrity during collection and transport while minimizing cost and logistical complications at the clinical site. The ABBR is also responsible for prospectively tracking and reporting on biospecimen collection activities for all Alliance clinical trials and when necessary, work with other Alliance team members to resolve systematic hindrances with biospecimen collection.

11.2.2 Storage. The ABBR is responsible for storing all biospecimens collected on NCTN and NCORP Alliance trials using methods that optimally preserve biological integrity and ensure biospecimen security.

11.2.3 Processing. The ABBR may be responsible for initial processing of tissue and biofluid specimens to a stable state for long-term storage. This may include centrifugation and/or separation of blood components and processing or embedding of tissue samples. At the discretion of each ABBR biorepository PI, the trial-associated biorepository site may develop and validate specialized processing methods to support specific trial procedures. An ABBR site may also perform secondary processing procedures, such as nucleic acid extraction, tissue sectioning, or tissue microarray (TMA) construction in order to create ‘assay ready’ materials that may be distributed for correlative science studies.

11.2.4 Quality Assurance. The ABBR is responsible for conducting or facilitating the conduct of quality assurance procedures for all collected biospecimens. This includes documenting physical quality of all specimens received, ensuring that proper biospecimen identification is preserved, facilitating histopathology review of tissue specimens when necessary, and ensuring that all material leaving the biorepository is fit for purpose and of suitable quality for all studies planned with those biospecimens.

11.2.5 Regulatory Compliance. The ABBR is the custodian and ‘honest broker’ of all biospecimens collected from patients enrolled on Alliance clinical trials. The ABBR ensures that biospecimens are appropriately de-identified and utilized for scientific studies that are commensurate with the corresponding patient informed consent.
11.2.6 Distribution. The ABBR works with other components of the Alliance to facilitate the review and distribution of biospecimens for correlative science studies.

11.2.7 Direct submission. For all Alliance trials, the ABBR should be the primary resource for all biospecimen collection, processing, and storage activities. In some cases, however, it may not be desirable or feasible to have biospecimens sent or processed by the ABBR. In these cases, with permission from the Director of Translational Research Operations and/or the Principal Investigator of the Alliance Translational Research Program, biospecimens may be sent directly to an investigator or commercial laboratory. However, even in such cases the investigator or commercial laboratory must follow all policies and procedures related to Alliance biospecimen tracking and handling (as outlined in this document). Furthermore, all biospecimens still remain under the custodianship of the ABBR and any remnant specimens must be returned to the ABBR at the completion of the assay. Examples include:

11.2.7.1 Assay requires rapid processing of fresh biospecimens using a technology or platform that is not available at the ABBR.

11.2.7.2 Assay is consumptive of the entire biospecimen and no material would remain for banking or future use anyway.

11.2.7.3 Assay is an integral biomarker assay that must be performed in a clinically accredited clinical laboratory and/or with rapid turnaround time, following clinical standards of biospecimen identity management and chain of custody.

11.2.8 Depending upon the specific trial design, the ABBR may support biospecimen activities for three different study types as defined by the NCI, the NCTN, and the Alliance. Each activity may be supported by a different funding mechanism, as explained below.

11.2.8.1 Integral Biomarker Studies. Studies in which biospecimens are mandatory and collected to perform an assay (or pathology review) in ‘real-time’ for the purposes of determining patient eligibility, arm assignment, or stratification. As noted above, biospecimens collected for integral biomarker studies may be sent directly to the relevant assay lab. However, once the biomarker assay is complete, unused biospecimens must be sent to the ABBR for other embedded or secondary use studies, unless determined otherwise by the ABBR director.
11.2.8.2 **Integrated (Embedded) Correlative Studies.** Studies in which biospecimens are collected to perform a well described, pre-defined correlative biomarker study that may be a secondary or tertiary end point of the trial itself. Collection may or may not be mandatory. With appropriate consent, remnant biospecimens from integrated correlative studies may be stored and used for stand-alone secondary correlative science studies.

11.2.8.3 **Biobanking for Stand-alone Secondary Correlative Studies.** Collection of biospecimens in the absence of a specific study that is described in the trial protocol itself, but that may be stored and made available for future studies proposed by investigators within or outside of the Alliance or the NCTN groups.

11.2.9 In addition to facilitating biospecimen collection for Alliance clinical trials, the ABBR may serve as a biorepository site for any NCTN intergroup trial, even if the Alliance is not the ‘lead group’ for that trial. As described below, support for intergroup trial biobanking activities must be pre-arranged prior to trial activation.
11.3 Biospecimen Collection Funding

A number of different funding mechanisms support ABBR activities. Funding is dependent upon the trial and the nature of the activity.

11.3.1 NCI U24 Biorepository Funding. The Alliance U24 biorepository grant is designed to support the staff and resources necessary for basic biorepository operations that include routine biospecimen processing, biospecimen storage, biospecimen information management, and administrative functions. Activities that are NOT supported by U24 funding include:

11.3.1.1 Design, manufacturing, and shipping of specialized biospecimen procurement kits.

11.3.1.2 Procedures related to biospecimen procurement at the site.

11.3.1.3 Biospecimen shipping.

11.3.1.4 Specialized biospecimen processing.

11.3.1.5 Pathologist time for central pathology review to confirm diagnosis.

11.3.1.6 Extraction of nucleic acids, or other secondary biospecimen processing.

11.3.2 Clinical Trial Budget. For some trials where biospecimen collection, processing, or pathology review is integral to the trial itself, these expenses may be primarily part of the trial budget and supplemented by U24 biorepository funding where appropriate. Otherwise, funding may be obtained from other sources noted below.

11.3.3 BIQSFP. The NCI BIQSFP mechanism may be used to support the conduct of integral and/or integrated biomarker studies as well as the expense of biospecimen procurement, shipping, and processing to conduct those studies. This funding mechanism will not support collection of biospecimens for other correlative studies or biobanking purposes.

11.3.4 Non-NCI Funding. Funding from other non-NCI sources (e.g. Komen Foundation, DOD, Breast Cancer Research Foundation), if obtained, may be used to support the construction and distribution of specialized biospecimen collection kits, reimbursement for research biospecimen procurement procedures, and specialized processing at the ABBR, when needed.
11.3.5 **Research Grants (Federal and Non-federal).** Investigators requesting biospecimens for either embedded / integrated correlative science studies or secondary use studies should anticipate that there will be nominal costs associated with the preparation (i.e. TMA slides, nucleic acid extraction, tissue quality assurance review) and distribution of biospecimens for funded research projects. These should be supported by research grant budgets, with expenses returned to the appropriate ABBR site to help support operations.

11.3.5.1 Costs for secondary processing of biospecimens for research studies will be charged by each ABBR site. Charges will be dictated by individual ABBR site policies.

11.3.5.2 Additionally, for secondary use studies, a standardized ‘application’ and/or ‘processing fee’ may be charged, in keeping with NCI NCTN policies.

11.3.6 Prior to any trial activation, an appropriate and sufficient funding source(s) should be identified to support all aspects of required biospecimen-related activities, from procurement to distribution. Funding resource(s) for integral biomarker must be secured prior to study activation.
11.4 Correlative Science and Biospecimen Collection Protocol Development

11.4.1 Proposals to utilize the specimens collected in a prospective trial ideally should be included in the clinical trial protocol concept at the time it is submitted to the Alliance Study Concept Review Committee. An appropriately powered, foreseeably funded, biospecimen-based correlative science study with a strong biological and/or clinical rationale may be included as a secondary end-point of the trial itself and will not need further review or approval once it has been approved in the context of the trial itself.

11.4.2 Once an NCI-approved trial concept moves to protocol development phase, stakeholders from the TRP Pathology Committee and/or TRP Biorepository Committee should begin immediate work with the Alliance Offices, disease/modality/discipline committee, Alliance Statistics and Data Center, Trial study chair(s), and Correlative Science co-chair(s) to develop the integral /integrated / biobanking study plan and biospecimen collection logistics.

11.4.3 Investigators performing laboratory studies may serve as study chairs of Alliance correlative science companion trials.

11.4.4 All embedded correlative science (CS) research requires review and approval by the disease/modality/discipline committee and TRP prior to submission of the main study to the NCI for final protocol approval. Subsequent review of the embedded CS research may be also required by the Alliance biorepository and disease/modality/discipline committee CS vice chairs during the protocol development process. Additional review of other relevant Translational Research Program sub-committees, such as Pathology Committee, Imaging Committee, Pharmacogenomics and Population Pharmacology Committee, Sequencing Committee may also be required for some studies.

11.4.5 Collection time points and biospecimens to be collected at each time point will be defined in a biospecimen collection calendar. Considerations in developing the correlative science and biospecimen collection plan include:

   11.4.5.1 Biospecimens that are required for planned integral / integrated biomarker studies.

   11.4.5.2 Low cost, minimally invasive collection opportunities (ideally synchronized with collections required for integral / integrated biomarker studies or standard of care) that can be leveraged to create a trial-based biospecimen resource for future correlative science studies.
11.4.5.3 Biospecimens, collection methods, and collection time points that minimize cost and simplify the logistics of collection, shipping, and processing.

11.4.6 Protocols that include a “research use only” biopsy must specify eligible biopsy location(s), methods and number of cores must be defined, along with other protocol specific requirements. Source(s) of funding for research tissue collection must be identified (see section 11.3).

11.4.7 Protocols that require extensive specimen sampling or processing, non-standard specimen collection time point, or the use of “kits” must be reviewed and approved by the TRP Operation Director and the ABBR director. Source of funding for any “kits” or special collection materials must be identified (see section 11.3).

11.4.8 Protocols that require central pathology review require approval by the TRP Operation Director and Pathology Committee. Source(s) of funding for real time central pathology review must be identified (see section 11.3).

11.4.9 Protocols that require central imaging review require approval by the TRP Operation Director, Imaging Committee and Imaging and Radiation Oncology Core lab (IROC). Source(s) of funding for real time central pathology review must be identified (see section 11.3).

11.4.10 Protocols that require international specimen shipping must be reviewed and approved by the TRP Operation Director and the ABBR director. Sources of funding for international specimen delivery must be identified (see section 11.3).

11.4.11 Although not required, it is strongly recommended that the study chair contact the TRP Operation Director, TRP Executive Officer, the ABBR director, and the appropriate disease committee CS co-chairs, the disease pathology cadre leaders, the disease Imaging Committee liaison, or other relevant TRP subcommittees, if applicable, prior to study concept submission to the SCRC.

11.4.12 Amendments to the main study wherein the embedded CS research is modified require review and approval by the main Study Chair, Correlative Science Study co-Chair, study statistician, and TRP Operation Director. If these changes involve modification to the standard protocols for biospecimen collection, processing, or shipping, then review and approval is also needed from the ABBR director. If these changes involve modification to the standard protocols
for imaging collection or processing, then review and approval is needed from the Alliance Imaging Committee and/or IROC.

11.4.13 Once a biospecimen collection schedule is created and approved by all stakeholders, a budget will be created. Based upon the cost and the parameters discussed in section 11.3 Biospecimen Collection Funding, appropriate funding must be identified.

11.4.14 The ABBR site that will support biospecimen collection for a trial will be determined by the ABBR director, with approval from the corresponding ABBR site director. Considerations for choosing the ABBR site include:

11.4.14.1 Existing site capacity and resources to manage a new collection.

11.4.14.2 Need for central pathology review or other correlative science support. To minimize shipping costs and logistical complications, trials where pathology support or correlative study assays will be provided by an institution that is also an ABBR site should also use that site for biobanking.

11.4.14.3 Biospecimens from neuro-oncology and Alliance cancer control program trials will be preferentially banked at the MAYO site.

11.4.14.4 Biospecimens from hematological malignancy trials will be preferentially banked at the HEME site.

11.4.15 Sites with logistical inability (e.g., sites outside the continental U.S.) to collect, process and ship specimens according to the protocol must apply for a waiver for exemption with the Alliance Protocol Operations Office, the Alliance TRP, the Alliance Statistics and Data Center (SDC), the study chair and the disease/modality/discipline committees. An administrative memorandum stating the shipping issue(s) and any protocol violation(s) from the site must be approved and filed with Protocol Operations through the assigned Protocol Coordinator prior to study activation. Collection, processing and shipping instructions for these sites will be provided on a study-by-study basis by the assigned biorepository.
11.5 Biospecimen Collection Policies

11.5.1 Each trial protocol document or associated Correlative Science Manual (CSM) must specify how to collect, prepare and ship specimens to the appropriate ABBR site. Questions regarding the collection and/or shipment of the materials should be directed to the assigned biorepository site where the specimen is being sent.

11.5.2 Each trial protocol or CSM document will follow a standard set of protocols (SOPs) for biospecimen collection, shipping, and processing.

11.5.3 All sites are required to send protocol-mandated biospecimens to the appropriate ABBR site, providing that appropriate patient consent is obtained and it is physically possible to send such biospecimens.

11.5.4 In cases where institutional policy prohibits the release of clinical pathology tissue blocks, an enrolling site may receive permission to submit a tissue block alternative (such as unstained slides or a tissue punch from the block) provided that permission is granted by the TRP Operations Director and trial study chair(s) and study co-chair(s). Note that for some protocols, submission of a tissue block may be absolutely required for participant enrollment.

11.5.5 ABBR biorepository sites themselves are not clinically-accredited medical laboratories. Therefore, any biospecimen processing that must be performed by a clinically accredited analytical laboratory (e.g. for integral biomarker testing or return of individual patient results) should not be performed by the Alliance biorepository. The Alliance biorepository is allowed to receive and store slides for retrospective histopathology review. All local diagnostic slides submitted for histopathology review can be returned to submitting sites upon request.

11.5.6 All specimens shipped to Alliance repositories must have patient consent and be accompanied by the appropriate paperwork as outlined in the protocol (e.g. forms, pathology report, etc.).
### 11.6 Biospecimen Processing and Storage Policies

**11.6.1** For diagnostic clinical pathology tissue specimens that have been submitted to any Alliance repository, the appropriate representative sections and/or cores will be prepared and the block will remain on file and will be available to the submitting institution for any medical-legal need.
11.7 Biospecimen Reporting and Tracking

11.7.1 All biospecimens submitted by sites are tracked by a database system (the Biospecimen Management System—BIOMS). Any exception must be granted by the ABBR director.

11.7.2 Each specimen submitted must be accompanied by the appropriate paperwork, as required by the protocol. Local records are kept in addition to the database. Local records will be secured in a locked cabinet/office at all times and database security will follow that recommended by the Alliance Statistics and Data Center (SDC).
11.8 Patient Consent, Confidentiality, and Regulatory Compliance

11.8.1 Patient consent for studies must be obtained prospectively. Consent forms must include adequate information to assess risks.

11.8.2 For trials that involve integral biomarker assessment to determine eligibility or treatment stratification, biospecimen submission for the integral biomarker assay is mandatory from all sites and all patients. All non-integral embedded correlative science requiring specimen submission must be offered to all patients enrolled on the study, although patients may opt not to participate. Therefore, specimen submission for non-integral correlatives, in general, is optional for the patient but not optional for the site. Exceptions to site participation in specific embedded correlative science studies may be granted by the study chair(s), in consultation with the corresponding correlative sciences co-chair(s) and the Translational Research Program Principle Investigator, in circumstances when the requisite resources or other infrastructure are not available at that site. In some rare instances, non-integral specimens can be mandatory for patients to participate after the group chair and/or the principal investigator of the Translational Research Program grant permission.

11.8.3 In the case of future (secondary use) studies that will use biospecimens collected for an Alliance clinical trial, including germ line susceptibility studies (studies of heritable genes), participants are asked to grant broad permission (i.e., it is unknown exactly what tests might be appropriate or performed in the future at the time the specimen is banked). Participants will NOT be re-contacted for each individual study.

11.8.4 Previously banked material that was not originally intended for extensive DNA studies (e.g., whole genome sequencing, whole exome sequencing, and genome-wide association studies) and for which informed consent was not originally obtained may be used for such research, but in these cases whether a re-consent must be obtained from the participant at the institutional level or not will be determined by the Alliance Ethics committee. For deceased patients, where re-consent is not practicable, whether a waiver of consent must be obtained at the institutional level or not will also be determined by the Alliance Ethics committee.

11.8.5 A unique Alliance biospecimen identification number will be assigned to each biospecimen submitted to Alliance Biorepositories. At the Alliance biorepositories, biospecimens must be stored and distributed with this number only. Investigators may not receive any patient identifiers, only the unique
biorepository specimen number. However, this may not apply to biospecimens sent directly to an investigator or commercial laboratory (see section 11.2.7).

11.8.6 Only authorized biorepository personnel may have access to match the unique sample ID with the Alliance patient ID number and only authorized Alliance statisticians may have the ability to link the unique specimen ID number, patient information, and clinical outcome. Exceptions must be approved by the principal investigator of the Translational Research Program and the group statistician for the Alliance.

11.8.7 If a registered patient withdraws consent from treatment but agrees to be followed on protocol, biospecimens may be submitted as required by the protocol.

11.8.8 If a registered patient withdraws consent for participation in the study or consent for follow-up, biospecimens may not be submitted.

11.8.9 If biospecimens have already been submitted but not distributed to investigators at the time when the patient withdraws consent, those biospecimens will be withdrawn from the repository and will be disposed of appropriately – either destroyed or, in the case of tissues, returned to the submitting institution upon request. Attempts will be made to retrieve any specimens that have been sent from the repository to investigators. However, processed specimens and the research data generated from them will not be rescinded, and may be used in study analyses.

11.8.10 Biospecimens are not released from the repository to investigators until the Alliance statistician assigned to the study or designee confirms the record of patient consent in the Alliance database. If a specimen is present in the repository but is later found to not have the appropriate patient consent, the specimen will be withdrawn from the repository and will be disposed of appropriately – either destroyed or, in the case of a diagnostic clinical pathology tissue blocks, returned to the submitting institution.

11.8.11 It is the Alliance policy that the Alliance biorepository shall not release clinical, pathology reports submitted by sites to correlative science investigators. Requests for data elements collected from local pathology reports should be submitted to the Alliance Data Center. This rule does not apply to study pathologists performing retrospective central reviews.
11.8.12 Disagreement between investigators and statisticians with respect to consent language for specific analyses will be adjudicated and decided by Alliance ethics leadership, statisticians, and the translational research program.

11.8.13 Reports (including manuscripts, abstracts, and progress reports) may never list any patient by name or initials. If needed, only unique identification codes may be used.

11.8.14 Unless indicated in the protocol and performed in a CLIA-certified laboratory, results from correlative science studies may not be provided to the patient or physician. Upon request, information may be made available as aggregate data in the form of abstracts or manuscripts.

11.8.15 The Alliance maintains Certificates of Confidentiality for each of its repositories from the US Department of Health and Human Services (HHS), which protects against the involuntary release of information collected during the course of the study. The researchers involved in a project may not be forced to identify a patient in any legal proceedings (criminal, civil, administrative, or legislative) at the federal, state, or local level. However, some information may be required by the Federal Food, Drug, and Cosmetic Act, the HSS, or for purposes of program review or audit.

11.8.16 For biospecimens sent directly to an investigator or commercial laboratory, certain Protected Health Information (PHI), such as patient initials and collection dates, may be sent to the investigator/commercial laboratory along with the biospecimens. In those cases, the trial protocol and informed patient consent form will inform sites and patients of any potential regulatory considerations.
11.9 Biospecimen Pathology Review

11.9.1 In no cases will the Alliance or an Alliance study pathologist render a clinical diagnosis. It is assumed that the submitting institution and the appropriate institutional pathologist will have rendered a clinical diagnosis in a way that is most appropriate for standard of care for the patient, prior to submission. In particular, fresh, ‘research only’ biopsy specimens will not receive a clinical diagnosis from an Alliance pathologist. If it is deemed necessary to make a histopathologic diagnosis of a biospecimen collected from a patient with an uncertain diagnosis (e.g., a metastatic lesion of a presumptive but unconfirmed primary origin), then it is incumbent upon the institution to perform any necessary diagnostic evaluation prior to submitting the biospecimen to the Alliance, even if the trial will perform a central pathology review.

11.9.2 In any case involving an apparent significant discrepancy between an observation made by an Alliance study pathologist and a diagnosis rendered at the submitting institution, the Alliance pathologist takes the following steps to determine the nature of the problem:

11.9.2.1 The study pathologist will verify the case identifiers. If the case was submitted to the Alliance biorepository for retrospective central diagnosis confirmation, the study pathologist will notify the biospecimen repository regarding the potential diagnostic discrepancy in the case. If the problem is clerical (e.g., incorrect specimen submitted to or distributed from the biorepository), the study pathologist and/or repository rectifies the problem directly with the submitting institution through Alliance institutional personnel (e.g., the institutional clinical research professional).

11.9.2.2 If it is determined that all case identifiers are correct, the Alliance study pathologist will contact the institutional clinical research professional (CRP) and, if necessary, will arrange to contact the submitting pathologist. The Alliance study pathologist will discuss the case with the submitting pathologist and detail the findings and the need for a re-review by the submitting institution. The Alliance study pathologist will discuss with the responsible institutional CRP and/or submitting pathologist whether other/additional pathologic materials from that case exist that might explain a discrepancy. Any problems related to case identification, specimen selection, or additional diagnostic information or materials will be discussed and resolved, if possible, by this direct communication,
and the nature of the resolution will be communicated to the repository by the study pathologist.

11.9.2.3 If an apparent discrepancy still exists, the appropriate Pathology Committee leader and at least one other committee member will review the case to confirm the diagnostic discrepancy. It is highly recommended that the study pathologist, the pathology committee leader and the submitting pathologist discuss the case directly before the final confirmation of discrepancy.

11.9.2.4 If the discrepancy is confirmed, the study pathologist or the chair of the Pathology Committee will immediately report the correct diagnosis to the responsible data coordinator. The data coordinator will report the correct diagnosis to the clinical research professional at the submitting institution. It is the responsibility of the clinical research professional to notify the submitting pathologist and the physician who registered the patient that there is a difference in diagnosis. The Alliance SDC will consider the discrepancy in the final analysis of the study.
11.10 Accessing Banked Biospecimens Overview

11.10.1 An Alliance membership is not required to request Alliance specimens.

11.10.2 Samples are furnished to the investigator by the appropriate Alliance specimen repository for the purpose of the project as approved. Research must be limited to that described in the approved protocol. Investigators may not share any portion of specimen or derivative specimen with another investigator or lab without permission of the NCI and the Alliance.

11.10.3 Investigators must discuss return of all unused specimens to the Alliance specimen repository prior to the completion of their correlative study. This includes RNA, DNA, urine, plasma, serum, tissue, slides, unstained sections, etc.

11.10.4 When investigators request specimens for nucleic-acid based (RNA / DNA) studies, it is the policy of the ABBR that whenever possible, only nucleic acid derivatives aliquots prepared by the ABBR will be distributed to the investigators. Exceptions can only be made with approval from the ABBR director.

11.10.5 No diagnostic, clinical pathology tissue blocks shall be released to research investigators. In general, no research blocks shall be released to research investigators either. However, exceptions for research block release may be granted by the ABBR director.

11.10.6 Once the project is approved, the investigator will be responsible for ensuring that his/her research is conducted under regulatory policies (human subjects, intellectual property, material transfer) governing their individual institution, as well as those set forth by the Alliance/NCI.

11.10.7 Correlative science investigators are required to have funding for their projects prior to receiving specimens.

11.10.7.1 In order to facilitate the successful application for funding, the Alliance will review concepts without established funding. For this review, investigators must provide the information requested for a preliminary concept review. In order to receive a letter of support from the Alliance, interested investigators must provide a preliminary concept at least six weeks prior to the grant deadline. Exceptions to this rule have to be approved by the principal investigator of the Translational Research Program.
Approved preliminary concepts, must include a description of the collaboration with the Alliance in their proposal submission and they must comply with the Alliance guidelines, which have been written to ensure scientific integrity, patient confidentiality, specimen protection, and support of the Alliance infrastructure resources.

Any collaboration with the Alliance that impacts Alliance resources, including protocol development, data management, statistical analysis, and specimen banking may require additional funding support. In addition to funds to support laboratory science (supplies, equipment, personnel, etc.), investigators may also be required to establish contracts and agreements with the Alliance, and/or subcontracts with the different resource offices of the Alliance being used, including the following:

11.10.7.3.1 Alliance Group Chair’s Office
11.10.7.3.2 Statistics and Data Center (for data management and statistical support)
11.10.7.3.3 Any relevant biorepository (for sample preparation and distribution, etc.)

Subcontract arrangements must be performed in accordance with Alliance policy and submitted in advance to ensure appropriate time is given for review and sign-off. A final copy of the grant must be submitted to and approved by the Alliance before submission to the granting agency.
Policy Name: Stand-alone Secondary Biospecimen Use Studies  
Policy Number: 11.11

Section: Biospecimen Repositories and Translational Research – 11  
Date Revised: January 1, 2018

11.11 Stand-alone Secondary Biospecimen Use Studies

11.11.1 Any proposal to utilize biospecimens from an Alliance NCTN study will be reviewed by the NCI NCTN Core Correlative Science Committee (CCSC) through a process managed by NCI. NCI NCTN-CCSC is charged with scientific review & prioritization of proposals requesting use of banked, non-reserved biospecimens collected from NCTN trials for use in correlative science studies. NCTN-CCSC prioritization ensures optimal use of these irreplaceable clinical trial biospecimens.

11.11.2 All correlative science investigators must agree to use the specimens for only the NCI NCTN-CCSC-approved research project and to follow Alliance and NCI policies and procedures. Investigators will be charged for all services that the biorepository provides (see 11.3)

11.11.3 Submission to the Alliance Translational Research Program for approval is strongly encouraged, but not required to obtain specimens from Alliance NCTN studies. A letter of support will be provided for the proposals endorsed by the Alliance.

11.11.4 A correlative science proposal should be based on an innovative idea, built around a strong biologic hypothesis including preliminary data supporting the hypotheses and/or feasibility, be scientifically valid and have significant clinical relevance. The investigator must demonstrate expertise, both technical and scientific, relevant to the work proposed. Therefore, previous publications in the area and/or preliminary data are required. Preliminary data are also required to evaluate the scientific rationale and logistics of the concept, the performance characteristics of the assay(s) to be employed (including accuracy compared to a gold standard, reproducibility, variability, and/or other available analytic validation), and to demonstrate clinical relevance.
11.12 Data Generation, Ownership, and Publications

11.12.1 Data from all laboratory tests performed on samples from any Alliance repository will be submitted to the Alliance SDC, usually via electronic means. The analysis of the data will be conducted by the responsible Alliance statistician or designee with the necessary expertise. The designee must be approved by the principal investigator of the TRP and the group statistician. The group statistician must approve any exception to this rule.

11.12.2 All publications must be reviewed and approved by the Alliance, following guidelines in the Alliance Policies and Procedures. The grant support of the appropriate Alliance repository will be acknowledged in publications.
12 Investigational agents

In Alliance studies, any agent that is provided to institutions is considered “investigational” for purposes of this policy. For investigational agents used under an IND, the IND holder is either the NCI or Alliance. Investigational agents may be provided by NCI/CTEP or directly by the industry partner. Investigational agents may be distributed to the institutions by NCI, industry, the Mayo Cancer Center Pharmacy or a third party distributor. The Alliance generally follows PMB policies (https://ctep.cancer.gov/branches/pmb/default.htm) and CTEP investigator guidelines (https://ctep.cancer.gov/investigatorresources/investigators_handbook.htm) for all IND investigational agents, irrespective of the IND holder.

12.1 Agent Accountability and Procurement

12.1.1 National Cancer Institute (NCI) Investigational Agents

Investigators must have current investigator registration documents (FDA Form 1572, Financial Disclosure Form, HSP/GCP training, biosketches, CV) on file with the NCI in order to receive investigational agents. These registrations must be renewed annually. Registration must be completed via the NCI Registration and Credential Repository (RCR). Additional information regarding registration types and required documentation is available at the Cancer Therapy Evaluation Program (CTEP) website (https://ctep.cancer.gov/investigatorresources/).

Investigational agents provided or distributed by the NCI are ordered through the Pharmaceutical Management Branch (PMB) Online Agent Order Processing (OAOP) application. Access to the OAOP system requires a CTEP Identity and Access Management (IAM) account and the maintenance of an active account status and a current password.

NCI distributes investigational agents for which it holds the IND and may also distribute NCI investigational agents, either Alliance-held IND or IND exempt, provided by industry.

12.1.2 Investigational Agents distributed by the Alliance

Instructions for ordering agents distributed by the Alliance or third party distributors vary from study to study, and can be found in the Drug Formulation, Availability and Preparation section of the protocol. The specific form required to ship drug to an institution is described in the protocol.
12.1.3 Shipment of investigational agents

Investigators have a single 1572 form on file with the PMB. Multiple pharmacy addresses may be listed on the 1572 form. By providing accurate shipping information this will assure that the FDA regulations are being followed, along with decreasing investigational agent shipping delays and expense and ensures accountability.

Investigational agent(s) will only be shipped to the designated pharmacy on the 1572 form of the investigator who is ordering the agent and must not be shipped to any other addresses after receipt at the pharmacy of record. Alternatively, investigators at affiliate institutions may order agents directly from PMB and not through their main member institution.

PMB policy allows centralized pharmacies to receive investigational agents for re-distribution to local satellite institutions and affiliated investigators who are registered with PMB and have designated a “central pharmacy” as their shipping address. If investigational agent is ordered through the main member institution, then the agent can be couriered to the satellite location if necessary. When agents are transported between control and satellite locations, care must be taken to ensure all appropriate storage conditions are maintained.

In the instance of investigators who staff more than 1 location, investigational agent(s) should be ordered to the central pharmacy where the patient will be receiving the investigational agent.

PMB policy also forbids secondary distribution of investigational agents to physicians who are not listed on the Delegation of Tasks Log (DTL) or transfer of investigational agents between institutions or other sites. Shipment of agents directly to patients is not allowed.

12.1.4 Use of Investigational Agents

Investigational agents must be used only in accordance with the protocol and only for patients registered on the study. Investigators must not charge for or seek reimbursement for investigational agents.

Commercial agents may not be substituted for an investigational agent nor can an investigational agent be used to “pay back” or “replace” commercial supplies.
The Alliance audits the pharmacy according to the NCI Guidelines for Auditing Clinical Trials (CTMB Guidelines) section 5.3 (https://ctep.cancer.gov/branches/ctmb/clinicalTrials-monitoring.htm).

Compliance with investigational drug use and accountability procedures is reviewed at the time of Alliance audits and will result in the pharmacy audit being rated as critical non-compliant, non-compliant, compliant or not reviewed. A rating of critical non-compliant will automatically result in an Unacceptable audit rating for Drug Accountability and Pharmacy. Any Unacceptable rating will require a re-audit within 12 months.

Auditors review investigational agents provided by industry partners according to the same procedures used for agents provided by NCI.

12.1.5 Storage and Accountability of Investigational Agents

A pharmacist or designated individual is responsible for investigational drug ordering, storage, dispensing and accountability. All study site personnel responsible for investigational agent(s) accountability must be listed on the Delegation of Tasks Log (DTL). The appropriate NCI Drug Accountability Record Form (DARF) should be used to record the receipt and disposition of all drugs supplied (by the NCI or pharmaceutical companies) for Alliance protocols. Specific procedures for completing DARFs and policies for storage and accountability are available on the PMB website. Guidelines are also available in the CTEP Investigator’s Handbook.

12.1.6 Deviation from Study Protocol

The appropriate Alliance protocol resource must be contacted if the handling or dispensing of the investigational agent(s) deviate from the protocol instructions. The deviation must be reported to the IRB of record, documented in a note-to-file, and retained in the records of the site.
12.2 Investigational New Drug Applications

Alliance reviews each study in development to determine if an IND/IDE application is required for a trial.

12.2.1 Investigational New Drug (IND)

12.2.1.1 IND Required

IND applications for Alliance-held INDs are submitted to the FDA by the Alliance Chicago Office. The Alliance group chair is the Responsible Investigator on all Alliance IND applications. The FDA will provide documentation of IND approval.

12.2.1.2 IND Exemption

If the FDA determines that an IND is not required, the FDA will provide documentation of IND exempt status.

12.2.2 Investigational Device Exemption (IDE)

Alliance studies may include the use of investigational devices. As in the case of INDs, the Alliance will submit an application for Investigational Device Exemption. If a study includes an investigational device, the protocol provides instructions on how to obtain the device as well as information regarding any special handling requirements that must be followed.

12.2.3 FDA Reporting

For Alliance-held INDs, annual reports, correspondence, amendments, and all other reporting requirements are submitted by the Alliance Chicago Office. All adverse events (AE) whose causality may be both serious and/or unexpected (SUSAR) are reported to the FDA by the Alliance.
13 Industry relations

The primary sponsor of Alliance studies is the National Cancer Institute, through research grants supporting these studies and the necessary infrastructure. However, Alliance also works with pharmaceutical companies and other health-related industry concerns to allow access to new investigational drugs or products that are relevant to Alliance research interests and to acquire financial support for unfunded or under-funded activities of the Alliance. Financial support, if acquired, helps to defray the costs of protocol development, implementation, data management, monitoring, patient tests that are not covered by insurance, laboratory studies, auditing, and statistical analysis.

Negotiations with industry are managed through the Alliance Chicago Office. Study chairs and committee chairs are not authorized to negotiate or sign agreements on behalf of Alliance.

The Federal principles governing industry collaborators in oncology trials are well established. The relationship is described in a document entitled “NCI – Cooperative Group – Industry Relationship Guidelines” (http://ctep.cancer.gov/industryCollaborations2/guidelines.htm) that focuses on manufacturer-NCI drug development agreements called Cooperative Research and Development Agreements (CRADAs) and Clinical Trials Agreements (CTAs). In addition, the NCI may distribute drugs for a network group trial under a Clinical Supply Agreement (CSA), independent of a CRADA or CTA. Many network group trials, however, involve drugs that are not the subject of such agreements, but the basic tenets of these agreements still apply to Alliance-industry collaborations.

13.1 Industry documents

Studies supported by pharmaceutical companies require a legal agreement in order for funding to be provided to the Alliance for Clinical Trials in Oncology Foundation, a tax-exempt nonprofit organization with the mission of supporting the research and educational programs of the Alliance. Other documentation, e.g., inclusion of standard language in the protocol document, and/or a letter of understanding from Alliance regarding drug or device/services provision, may be necessary for selected studies.

Examples of information included in each document are provided below.

13.1.1 Legal agreement for provision of financial support

Description of funding to be provided, payment schedule, reporting requirements, data (if any) to be provided, advertising, termination, scope of work, and responsibilities of the parties. The parties to the legal agreement are the industry collaborator and the Foundation.
### 13.1.2 Protocol document

Standard language in the protocol document provided by NCI when drug is provided under a CRADA, CSA or CTA between the NCI and industry collaborator. The NCI language can be found in the document entitled “NCI Standard Protocol Language for Collaborative Agreements” ([http://ctep.cancer.gov/protocolDevelopment](http://ctep.cancer.gov/protocolDevelopment)).

Specific language in the protocol document may be required when drug is provided directly to the Alliance.

### 13.1.3 Letter of understanding regarding drug or device/services provision

Reports summarizing the progress of active studies are generated by the Alliance Statistics and Data Center and distributed at the group meeting (at least annually). The summary also includes a listing of published manuscripts and abstracts. The primary purpose of these reports is to inform Alliance meeting attendees as well as the National Cancer Institute of the current status of Alliance research.

This letter, from the Alliance group chair, may discuss:

- Information regarding the structure and function of the Alliance
- Information regarding how the drug/device/service will be used/applied
- Information regarding the specific study for which the drug/device/service is provided
- Reference to the Alliance Policies and Procedures, as appropriate
13.2 Confidential and proprietary information

In studies involving collaboration with the pharmaceutical industry, it is the responsibility of all institutional participants to maintain confidentiality with regard to proprietary, trade secret, or other confidential information. Confidential and proprietary industry information is strongly discouraged from inclusion in study protocol document.

All study chairs must abide by group policy that requires strict confidentiality of study information (see section 6). The Alliance statistician carries out all interim analyses in a confidential manner. No one other than those explicitly authorized to be part of the monitoring process has access to the results. All such persons must keep all aspects of their deliberations in strict confidence until the release of results is approved. Any violation of confidentiality is considered a serious offense.
13.3 Data ownership in the context of industry collaboration

Pursuant to NCI policy, the Alliance owns all data resulting from its trials. It is willing to provide accrual updates, copies of adverse event reports, regulatory documents, and study summaries to industry collaborators. If a collaborator wishes trial results or regulatory information to use for internal or regulatory purposes, the appropriate terms can be negotiated. Typically this requires a legal agreement. If a trial involves two or more investigational drugs, each company must normally consent to all data being provided to the other company/ies.
13.4 Release of data

Trial data may be made available to industry collaborators pursuant to executed agreements for data transfer. The data resulting from Alliance studies (except for adverse events reports and other data as mentioned in section 13.3) should be available to industry collaborators within six months of data maturity for primary endpoint, as long as funding is available for such endeavor. Confidential data under active monitoring by the DSMB are not released without the approval of the Alliance DSMB. Industry collaborator requests for data are managed by the Alliance Chicago office.
13.5 Indemnification

The Alliance and its Foundation are not liable for any acts or omissions of the industry collaborator with respect to the conduct of Alliance studies. The Alliance requests that the collaborator indemnify all investigators against loss under customary product liability principles, including responsibility for drug information in the Investigator’s Brochure. Because it lacks the legal power to do so (i.e., because it is not a legal entity), the Alliance is not in a position to indemnify collaborators or manufacturers against claims due to negligence of its members.
13.6 Intellectual property and patent rights

An invention resulting from work performed by an Alliance investigator generally is the property of either the investigator or the Alliance institution with which he or she is affiliated. It is the policy of the Alliance that investigators shall disclose to the Alliance group chair any inventions or discoveries, whether patentable or not, resulting from Alliance studies (aka “Intellectual Property”) in writing within ninety (90) days of discovery thereof. Upon receipt of the notification, the group chair, along with the investigator(s), will consider the appropriate institutional officials to participate in a meeting to discuss matters of recognition and remuneration related to the patenting, licensing, exploitation or commercialization of the Intellectual Property. It is not the intent of this policy to interfere with the publication of research results.

The Alliance has a contractual relationship with each of its member institutions that requires adherence to this policy and all policies described in the Alliance Policies and Procedures. In addition, the provisions in the NCI “Intellectual Property Option to Collaborator” (http://ctep.cancer.gov/industryCollaborations2/intellectual_property.htm) terms of award modifications apply to Alliance studies using investigational agents.
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13.7 Publication of study results

Consistent with the traditional principle of academic freedom, the right of the public to know about government-funded trial results, and the policy of major medical journals, the Alliance requires that its investigators have the absolute right to publish all study results. At least thirty days before submission of a manuscript for publication, a copy is provided to the industry collaborator for advisory review and comment, so that the manufacturer can protect patent opportunities and review for disclosure of proprietary information. See NCI Standard Protocol Language for Collaborative Agreements ([http://ctep.cancer.gov/industryCollaborations](http://ctep.cancer.gov/industryCollaborations)). If patent-related action is necessary, an additional sixty-day delay is provided.
13.8 Use of agent/devise provided by industry collaborator

Any agent/device provided by an industry collaborator may not be used by participating institutions and investigators for any purpose outside the scope of the protocol. Neither the institution nor the investigator may charge any third party payer or patient enrolled in the study for the agent/device, and the institution or investigator may not include the cost of such agent/device in any cost report to third party payers.
14 Public relations

14.1 Authorized group representation

No one other than the group chair, or the authorized representative of the chair, may represent the Alliance in any manner.
14.2 Public service

The Alliance receives major support from the National Cancer Institute (NCI), Division of Cancer Treatment and Diagnosis (DCTD), Cancer Therapy Evaluation Program (CTEP). The goal of this national program is to seek improved methods of cancer therapy, a goal shared by the Alliance. The Alliance represents a bridge between the NCI and cancer patients throughout the country who receive new methods of treatment devised by Alliance investigators and approved by the NCI. In addition, the Alliance depends upon the scientific and financial resources provided by academic medical institutions and community sites throughout the country. The group is committed to conduct its science in a spirit of open inquiry, to critically evaluate promising new ideas and technology, and to take measures to minimize the risks of these new treatments to study participants. The institutions that participate in Alliance studies must agree to furnish study-related data concerning participants who have consented to be enrolled in Alliance studies, regardless of the level of institutional funding, and to undergo audits that evaluate and help to insure the integrity of the data collected. In turn, the Alliance provides these medical centers with the opportunity to see their ideas evaluated in definitive national trials and to provide novel therapies to their patients.

The Alliance also receives support from the NCI’s Division of Cancer Prevention (DCP) and Division of Cancer Control and Population Sciences (DCCPS). Using these resources, the Alliance pursues studies to reduce the incidence and prevalence of clinically significant cancers, to alleviate the symptoms of cancer and the toxicities of cancer treatment, and to improve the delivery of cancer care in community and academic practices, with special emphasis on issues affecting minority, underserved, and elderly patient groups.

Thus, the Alliance serves three constituencies:

1. The public whose taxes support Alliance, in part

2. The research participants who agree to take part in Alliance-sponsored clinical and cancer control research

3. The academic institutions and community sites that support our many scientists, physicians, and staff
14.3 Dissemination of information to the general public

It is the responsibility of the Alliance Office of the Group Chair and each member of the Alliance to furnish accurate information concerning the Alliance and its research programs to the general public.

Questions from the public fall into several categories and are answered according to category.

- **Questions about new treatments:** These questions are usually referred to an executive officer or a protocol coordinator. If the Alliance has a relevant protocol that is open to accrual, it is appropriate to describe it and refer patients to an appropriate Alliance institution. If the question comes from a geographic location not served by Alliance, it should be indicated that there are other network groups that may also have studies that are appropriate, and that information concerning all NCI-sponsored clinical trials in cancer may be obtained from several websites (see the Alliance website [http://www.allianceforclinicaltrialsinoncology.org](http://www.allianceforclinicaltrialsinoncology.org) under the ‘Resources’ heading).

- **Questions of a medical nature about a specific patient:** The Alliance personnel do not furnish medical advice. For answers to questions of this nature, individuals are referred to the patient's physician.

- **Requests from patients or physicians for copies of Alliance protocols and forms:** Alliance protocols are considered confidential documents and are generally not provided to the public. A non-Alliance physician may receive a copy of an Alliance protocol upon request and after approval by the principal investigator of the Alliance Central Protocol Operations Program (CPOP) or the group chair. The request must be made in writing and the intended use of the protocol must be clearly stated.

- **Inquiries concerning gifts to support cancer research:** Questions should be referred to the Alliance for Clinical Trials in Oncology Foundation.

- **Questions about the Alliance history, structure, and membership:** Refer to the Alliance communications specialist, chief administrative officer (CAO), or refer to the Contact us section of the Alliance website.

- **Questions about Alliance research results:** The Alliance works closely with the NCI, industry partners, member institutions and patient advocacy groups to disseminate information regarding the activation, progress, results and findings of its research. All requests should be referred to the Alliance communications specialist.
Requests for access to the Alliance website: The Alliance website contains sections available to the general public as well as sections that are accessible only to Alliance members or others who have been granted access. The Alliance periodically receives requests for access to the password-protected section of the website. Such requests should be submitted in writing to the CAO and should explain the purpose of the request in detail. The CAO authorizes access if the request is deemed appropriate. In most cases, access is time-limited.

- All other questions: Refer to the Alliance communications specialist.
14.4 Confidentiality of patient information

The Alliance has instituted procedures designed to protect the privacy of its clinical trial participants. Although there are some limits to non-disclosure of information to federal regulatory agencies, the Alliance intends to protect the privacy of its clinical trial participants to the limit allowed by the law. The Alliance consent form describes the steps taken in this regard. Alliance information systems are HIPAA compliant and Alliance has received a Certificate of Confidentiality from the NIH to protect information about specimens or data obtained from participants in Alliance studies.

Information about Alliance clinical trials may also be provided to pharmaceutical companies, foundations and others that support the work of the group. In all instances the Alliance takes steps to protect the privacy of the study participant. Patient identifiers (including but not limited to patient name, social security number, address and phone number) are not released. Alliance reports and publications do not present information that would allow the identification of its study participants.
15 Data sharing

Each Alliance study has a formal protocol document, which includes a statement of the objectives of the study. Patient consent and authorization are obtained to collect the individual patient data required for addressing the study objectives. These data are transmitted from the treating or enrolling institution to the Alliance Statistics and Data Center (SDC), where the data are reviewed, processed and stored in the Alliance database. Not all information submitted becomes part of the electronic database; for example, only some information on supporting documents such as operative and pathology reports may be entered into the database. The electronic database is used as the basis for the analysis of Alliance studies, with the analyses performed by the staff at the Alliance SDC.

The procedures described here do not cover requests – from the National Cancer Institute (NCI), the Food and Drug Administration (FDA), or other federal agencies – for information required by federal regulations or by the terms of the grant awards from federal agencies (e.g., Cancer Therapy Evaluation Program [CTEP], NCI, and National Institutes of Health [NIH]) to the Alliance. Such requests will be honored as expeditiously as possible.

This policy covers requests for existing data, not requests for collection of additional data. Requests for individual-level genomic or other high-dimensional data not used in the primary publication (see section 15.4) may be subject to other NCI and NIH regulations.

The data requested by an investigator can include data generated from Alliance laboratory correlative studies. However, requests for use of biospecimens are covered by a separate evaluation and review procedure described in section 11.

The sharing of data with industry is further described in section 13.

15.1 Guidelines for availability of data sets

For phase 3 studies it is anticipated that individual-level de-identified data sets, that would be sufficient to reproduce results provided in a publication (i.e., published manuscript) containing the primary study analysis, will be available to individuals via the requesting procedures described in section 15.2 generally within six months of publication of the manuscript. It is anticipated that data sets containing patient-level entry data of all baseline variables summarized in the publication will be available within 12 to 15 months after the publication of the primary analysis.

For non-phase 3 studies, a patient data set containing the variables analyzed in the primary results paper will be available upon request (subject to restrictions in sections 15.3 and 15.4). This process could take several months, based on the type of request and workload amount/priorities of the SDC. Since these studies could be quite small, the release of data may also be constrained by the ability to de-identify data.
For publications that are not presenting the primary analysis of the trial, patient data sets containing the variables analyzed in the manuscript will be available upon request (subject to restrictions in sections 15.3 and 15.4). This process could take several months depending on workload and prioritization within the SDC.

Release of data collected in a clinical trial conducted under a binding collaborative agreement between CTEP and a pharmaceutical/biotechnology company must be in compliance with the terms of the binding collaborative agreement and must be approved by CTEP and the company. Release of data is also subject to the terms of any contracts between the Alliance and other entities, which cover any of the requested data. These two considerations could, in some instances, delay the release of data to requesting investigators.
15.2 Request procedures

While most analyses of Alliance studies are performed at the Alliance SDC, the Alliance also makes research data available to other investigators, as required by the policies of the NIH. An investigator who wishes to use individual patient data from one or more of the Alliance studies must make a formal request to the Alliance Chicago Office.

The Alliance requires documentation, which includes a brief description of the project, as well as documentation of Institutional Review Board (IRB) approval or exemption from the institution of the requesting investigator (see section 15.3). The Alliance also requires the investigator to sign a data use agreement specifying who will have access to the individual patient data and specifying that it will not be shared with other outside this specified set of individuals unless first approved by the Alliance.

There will be no scientific review of requests for data. If the Alliance is unable to fulfill a request, the Alliance will inform the investigator(s) of the reason the request cannot be fulfilled. In most cases it is likely the investigator(s) will be able to amend the request to comply with the procedures. If the Alliance believes the request will not be amendable, the Alliance will inform the investigator of the appeals process outlined in section 15.6, and also notify the lead chief of the Clinical Investigations Branch (CIB) of CTEP in the Division of Cancer Treatment and Diagnosis (DCTD) at the NCI and the lead NCTN program director. Release of the data is subject to the disclaimer in section 15.5.
15.3 Regulatory considerations

All research use of data collected on human subjects from network group studies led by the Alliance Central Protocol Operations Program and Alliance SDC is subject to applicable Office of Human Research Protections (OHRP) regulations and to applicable regulations of the Privacy Rule of the Health Insurance Portability and Accountability Act (HIPAA). Generally, patients have only consented to have their health information used for the objectives of the clinical trial in which they participated. Use of the data for other research projects is allowed only if an IRB has determined that use of the data in the project meets the minimal risk criteria for conducting the research without the patient's consent, if the use of the data in the project is exempt from consent requirements, or if the project does not constitute human subjects research. The required level of review or approval will generally depend on the degree to which the data have been rendered fully anonymous, de-identified, or coded.

Guidance on these matters can be found in the OHRP document “Guidance on Research Involving Coded Private Information or Biological Specimens” located at http://www.hhs.gov/ohrp/policy/cdebiol.html. Information is also available on the NIH website (http://privacyruleandresearch.nih.gov/clin_research.asp) for Clinical Research and the HIPAA Privacy Rule. The criteria for de-identification of data under HIPAA are given in the Code of Federal Regulations, Part 46, Section 164.514. It is possible to conduct most projects using coded data (as described in the OHRP Guidance) that meet the criteria for a limited data set that can be released under a data use agreement (as described in Part 46 of the CFR, Section 164.512 and in the NIH HIPAA guidance documents), without obtaining additional patient consent or authorization.
15.4 Genomic data sharing

15.4.1 NIH data sharing policies

In accordance with NIH data sharing policies, genomics data generated from Alliance studies are deposited into the database on Genotypes and Phenotypes (dbGaP). The Alliance Bioinformatics Unit is responsible for this process.

NIH data sharing policies are evolving as a consequence of the cost of investments in large-scale data generation in high-throughput genotyping and sequencing of samples collected in NIH-funded research studies. There is strong consensus that making data and complete results of studies broadly available to the scientific community helps to insure that the investments made in data collection and genomic (and other “-omic” studies requiring substantial investment of resources and generating large-scale data) studies provides the greatest benefit to stakeholders including NIH, taxpayers and the scientific community.

At the same time, there is appropriate concern for maintaining the privacy of patients participating in such studies, and respecting the consent procedures within existing studies. Unique challenges to full compliance with data sharing policies arise in the context of clinical trials because some of the outcomes measured within clinical trials are available relatively early in the trial, while some of the key outcomes are not available until the primary endpoint(s) of the trial are met. Because high-throughput genomic data can be generated very rapidly and may be appropriately applied to outcomes available early in the trial, the desire to facilitate data sharing and fully comply with NIH data sharing policies will inevitably collide with long-standing practices in clinical trials research that have traditionally precluded sharing of data from a study until it is completed.

15.4.2 Alliance genomics studies

Alliance genomics studies are typically conducted as companions to Alliance clinical trials. To ensure that the data sharing process addresses the concerns of all parties involved, a steering committee consisting of the relevant committee and study chairs and statisticians will be formed for each genomic study. Any steps to be taken with respect to data sharing will be reviewed by the relevant steering committee and proceed only upon approval.

Alliance Policies and Procedures — Data Sharing 15-5
It should be noted that as NIH policies regarding high-throughput genomic data sharing continue to evolve, it is expected that the corresponding Alliance policies will of necessity evolve as well.

15.4.2.1 Genotype data

De-identified (coded) high throughput genotype data (including intermediate files and/or information useful for copy number variation analysis) will be made available to public repositories (such as dbGaP) upon completion of quality control studies. The Alliance statistician associated with analysis of the trial and genotype data will determine when quality control studies have been completed, and will prepare data for submission. Publications by others that make use of only Alliance genotype data (for example, as control data for other studies) may be published at any time after submission.

15.4.2.2 Phenotype data

Phenotype data will be submitted at the completion of the trial once all data have been subject to quality and integrity checks. All phenotype data that are part of the Alliance electronic database, have been checked for quality and integrity, and are used in genetic studies will be deposited. The Alliance statistician associated with analysis of the trial and genotype data will determine when the standard Statistics and Data Center quality control processes have been completed and will prepare data for submission. Publications by others making use of Alliance phenotype data (with or without genotype data) will be embargoed until after publication of the primary paper reporting the primary endpoint results of the clinical trial. As in the case of any Alliance data sharing request, no phenotype data on a DSMB monitored study, will be released without a formal approval from the DSMB.

15.4.2.3 Results databases

As considerable time may elapse between submission of genotype and submission of phenotype data, the Alliance will develop results databases (see example at [http://www.pgscore.org](http://www.pgscore.org)) to serve results of genotype - phenotype association studies for phenotypes that have not yet been deposited. For example, genome-wide association studies conducted on intermediate phenotypes (e.g., pharmacogenetic phenotypes, or surrogate...
outcomes) may be available through the web site before final phenotype data are deposited in dbGaP because of the length of time required to obtain and quality check full outcome information. The use of results databases permitting extensive queries will improve access of the scientific community to results of the studies and serve as the necessary intermediate between completion of initial genetic studies (which may involve intermediate data) and completion of the clinical trial. Results databases may be made publicly available upon completion of Alliance-approved analyses that have undergone review of the steering committee and, in the case of studies that have not yet met the primary endpoint, been approved by the DSMB.
15.5 Release conditions and disclaimer

A simple, formal data use agreement specifying who will have access to the individual patient data (and specifying that it will not be shared with others outside this specified set of individuals) as well as covering the release conditions described below and the regulatory considerations described in sections 15.3 and 15.4 above is required.

It is anticipated that most data requests can be provided as non-complex data sets in electronic form.

Sometimes the data requested for analysis will not all be coded in the Alliance database, but will be available from supplementary material that was submitted as part of the trial. In this case, the data would need to be abstracted from the supplementary material. Data abstractions can only be performed if adequate funding to support the abstraction is available. Even if funding is available, the Alliance may not have staff available to perform the abstraction. In this situation, Alliance may consider inviting the investigator(s) to the Alliance SDC to perform the abstraction. Some funding for clerical support may still be required. Likewise, when data requested require data sets not available in easily obtained electronic format, especially for older trials, the Alliance may require funding for support to create the data set in an electronic format.

In releasing the data, the Alliance makes no representations and extends no warranties of any kind, either expressed or implied. There are no expressed or implied warranties of merchantability or fitness for a particular purpose, or that the use of the data will not infringe any patent, copyright, trademark, or other proprietary rights. No indemnification for any loss, claim, damage, or liability is intended or provided.

Copies of any manuscript arising from the project associated with the data request must be sent to the Alliance; however, approval of the manuscript is not a condition for use of the data.
15.6 Appeals process

If a request for data is denied, the applicant may appeal the decision. The appeal is reviewed by the Alliance group chair, the lead NCTN program director, CTEP associate director or his/her designee, and an outside statistician (i.e., a statistician who is not a member of the Alliance). The outside statistician is named jointly by the Alliance group chair and the lead NCTN program director.
15.7 Fees

Routine costs associated with preparing standard data sets are viewed by NCI as covered by grants for the Alliance Operations Center and Alliance SDC funded under the NCTN Program. Fees will not be charged for the release of non-complex electronic data sets. For complex data sets where substantial work is involved, fees may be charged for preparing and documenting the data set. Any fees will be limited to the actual time, effort, and materials required for preparing and documenting the data set.
16 Study monitoring and interim analyses

The primary purpose of monitoring a clinical trial is to ensure the safety and well-being of the specific participants entered on the trial. All treatment protocols must include a formal monitoring plan. All randomized phase 2 all phase 3 trials, and some specially-designated trials are formally monitored by a standing Alliance Data and Safety Monitoring Board (DSMB). The monitoring functions for other treatment trials (e.g., phase 1 and non-randomized phase 2), including accrual monitoring, are carried out by the study chair, the primary statistician, and the executive officer along with other members of the study team and Alliance staff. Non-treatment trials do not usually require formal monitoring procedures, however the DSMB does monitor selected Cancer Control studies.

16.1 Study monitoring by the DSMB

16.1.1 Studies requiring DSMB monitoring

All Alliance-led phase 3 and randomized phase 2 CTEP or DCP sponsored trials are monitored by the Alliance DSMB. Other studies may be monitored by the DSMB if deemed appropriate by the group chair and DSMB chair.

16.1.2 Function of the DSMB

The responsibilities of the DSMB are as follows:

1. The primary responsibility of the DSMB is to review adverse event data in conjunction with interim analyses of outcome efficacy data (prepared by the study statistician) and to recommend whether the study needs to continue per protocol or be changed or terminated based on these analyses. For phase 3, phase 2/3, and randomized phase 2 trials, the committee also determines whether and to whom outcome results should be released prior to the reporting of study results at the time specified in the protocol.

2. The DSMB reviews reports of related studies performed by the network groups or other organizations to determine, considering information and recommendations supplied by the study team, whether the group study needs to be changed or terminated.

3. The DSMB oversees the safety and accrual data; however it is also the responsibility of the study team to review the safety and accrual information on a regular basis.
4. All clinical trial data release requests (e.g., baseline data for correlative studies or assay methodology evaluation) on DSMB monitored studies have to be submitted to the DSMB for review and approval.

5. The DSMB reviews major modifications to the study proposed by the study committee (e.g., termination, dropping an arm based on toxicity results or other trials reported, increasing target sample size).
16.2 Overview of DSMB procedures

Each study to be monitored requires periodic (at least every 6 months) confidential reports to be prepared by the primary statistician. These reports are submitted to the DSMB, a single standing committee established for the purpose of reviewing all of the individual reports. No individuals other than members of the DSMB receive a copy; specifically, the study chair does not receive a copy.

16.2.1 Confidentiality

All interim analyses are carried out in a confidential manner. No one other than those explicitly authorized to be part of the monitoring process have access to the results. All such persons must keep all aspects of their deliberations in strict confidence. Any violation of confidentiality is considered a serious offense.

All members of the DSMB are required to sign a written confidentiality pledge. The Alliance SDC maintains confidential files of all reports and actions taken on each study. No communication of the deliberations of the committee, either written or oral, may be made except as provided for in these DSMB policies and procedures. Any violation of confidentiality must be reported to the group chair.

16.2.2 Membership

The DSMB chair is nominated for a five-year term by the group chair and confirmed by CTEP. The group statistician is a non-voting member of the DSMB. All other members are appointed by the group chair for three-year terms, and include individuals primarily from outside of the Alliance. The majority of the voting DSMB members cannot be affiliated with the Alliance, and voting quorums for a DSMB meeting require that the majority of voting members not belong to the Alliance. Individuals are selected based on their breadth of experience, reputation for objectivity, absence of the actual conflicts of interest or the appearance of conflicts of interest, and knowledge of good clinical trial methodology. There is at least one lay member and a voting statistician from outside the group. One or more CTEP physician(s) and a CTEP statistician, selected by CTEP, are non-voting ex officio members, as are one or more DCP physician(s), selected by the DCP. Members of the DSMB who chair or co-chair studies being monitored by the committee excuse themselves from all DSMB discussions concerning that study and do not receive DSMB reports concerning that study. Members of the DSMB who are leaders (chair or vice chair) of disease or modality committees excuse themselves from all DSMB discussions concerning...
studies being conducted by their committee and do not receive DSMB reports concerning those studies.

16.2.3 Meetings

The DSMB meets at least twice yearly, ordinarily in conjunction with scheduled group meetings (see section 5). Additional DSMB meetings may be held at any time or in any form as decided by the DSMB chair. At a minimum, an in person (face to face) meeting must be held at least every 18 months.

The DSMB meeting itself consists of open (optional, per discretion of DSMB chair) and closed (required) sessions for each study under consideration. During the open session, the study chair, primary statistician, and committee chair are available to answer questions posed by DSMB members. During the closed session, the DSMB decides what action, if any, is required. The study chair, primary statistician, and committee chair must absent themselves from the closed session even if they are members of the DSMB.

16.2.4 Recommendations

The results of each DSMB meeting are summarized in a formal report by the Alliance Group Statistician and sent by the DSMB chair to the group chair within one week of the meeting (urgent matters are addressed immediately). The DSMB report contains recommendations on whether to modify or close each study reviewed, whether to release and report the results, and whether to continue accrual or follow-up. A primary recommendation (e.g., continue with no change; recommended or required modification; release study results and stop further DSMB monitoring) must be included in the document. The group chair must approve these recommendations before any action is taken.

In the unlikely situation that the Alliance group chair does not concur with the DSMB recommendation, the Alliance group chair must discuss his/her reasons for not accepting the DSMB recommendation with the chief, Clinical Investigations Branch (CIB). The chief, CIB, will then inform the CTEP associate director of the recommendation of the DSMB and of the Alliance group chair’s reasons for disagreeing with the recommendation. The CTEP associate director, chief, CIB, and the Alliance group chair, in consultation with the DSMB chair, will be responsible for reaching a mutually acceptable decision about the study. Confidentiality will be maintained during these discussions, but relevant data will be shared with the Alliance group chair, chief, CIB, CTEP associate director, and other parties whom they wish to involve in reaching a decision. In the exceptional circumstance that a mutually acceptable decision cannot be reached, final responsibility for a decision will
rest with the CTEP associate director in consultation with the director of the Division of Cancer Treatment and Diagnosis.

The group chair, or designee, is responsible for notifying the study chair, primary statistician, and committee chair before the recommendations of the DSMB are carried out. An edited version of the recommendations is distributed to Alliance membership. The Alliance Central Office keeps an archive of DSMB minutes and recommendations.

**16.2.4.1 Study change for patient safety reasons**

In the event that the DSMB recommends a study change for patient safety reasons (including early stopping for inferior therapy), the Alliance group chair will act to implement the change as expeditiously as possible. For studies that are being closed based on a DSMB recommendation, although CTEP/DCP pre-approval is not required, the Alliance group chair (or his/her designee) must inform and discuss the closure of the study with the chief, CIB, (chief COPTRG if a DCP study) or his/her designee before disclosing the study closure to anyone. If the DSMB recommends closure of a study, the NCI/DCTD physician member of the DSMB will provide the current 24/7 contact information for the chief, CIB, or his/her designee.

**16.2.4.2 Study closure due to slow accrual**

In the event that the DSMB recommends a study be closed early due to slow accrual, then the recommendation of the DSMB would be processed as described in 16.2.4.1 above. Note: NCI/DCTD/CTEP may have additional closure policies that apply to studies with slow accrual that have not yet had formal interim efficacy analyses presented to the DSMB.

**16.2.4.3 Study change for non patient safety reasons**

In the event that the DSMB recommends a change in a study for reasons other than either patient safety (e.g., to extend accrual because of an event rate lower than expected) or study closure due to slow accrual, the DSMB will provide to the Alliance group chair an adequate rationale. In the absence of disagreement, the Alliance group chair (working with the study chair) will be responsible for having an amendment prepared and submitted to CTEP’s Protocol and Information Office reflecting the recommendations of the DSMB and providing the rationale for the changes. (This is
required even if NCI/DCTD/CTEP approval has been obtained prior to the amendment being presented to the DSMB.) NCI/DCTD/CTEP approval of the amendment will be required prior to implementation of the change, although it is anticipated that a decision to override the recommendation of the DSMB will be made only in the most exceptional circumstances. In the event that the Alliance group chair disagrees with the DSMB recommendation, the recommendation would be processed as described in 16.2.4.1 above.

For DSMB recommendations specific to cancer prevention and control trials funded by a NCORP Research Base grant, the appropriate NCI staff to include and report to are the DCP/Community Oncology and Prevention Trials Research Group (COPTRG) program director (instead of the NCI/DCTD physician member of the DSMB), the chief of COPTRG (instead of the chief, CIB) and the associate director for clinical research in DCP (instead of the CTEP associate director), and the director of the Division of Cancer Prevention (instead of the director of the Division of Cancer Treatment and Diagnosis).

### 16.2.5 Study modifications

Major modifications to the study design by the study team not motivated by confidential outcome data or patient safety/toxicity data (e.g., increasing the sample size because of more rapid than expected accrual) must be discussed with NCI/DCTD/CTEP/DCP before being presented to the DSMB for consideration. If NCI/DCTD/CTEP/DCP is willing to approve the modifications, the network group informs the DSMB at the next scheduled DSMB meeting.

### 16.2.6 Release of results

For phase 3, phase 2/3, and randomized phase 2 trials, any release of outcome data (either internal to the network group, to NCI personnel not members of the DSMB, or external [e.g., a paper presented at professional society meetings, seminars, papers, etc.]) prior to the final approval of general dissemination of results must be reviewed and recommended for approval by the DSMB to the Alliance group chair. In general, outcome data from phase 3, phase 2/3, and randomized phase 2 trials would not be routinely made available to individuals outside of the DSMB until accrual has ceased and all patients have concluded their randomized treatment, and completed study follow-up and/or reached a protocol-specified endpoint. After this time point, the DSMB may recommend the release of outcome data on a confidential
basis to the study chair for planning the preparation of manuscripts, and/or to a small group of individuals for purposes of planning future trials. The DSMB will consider special requests for information from the disease committee chair prior to that time point. The DSMB should be made aware of any communication of analysis results from phase 3, phase 2/3, and randomized phase 2 trials outside of the statistical center at any time. The Alliance group chair may not be able to accept the recommendation of the DSMB to release data for a specific trial if the Alliance and/or NCI/DCTD/CTEP has a binding agreement with a company collaborator (or other entity) that specifies data exclusivity for the trial without discussing the release with CTEP (for Alliance trials with a CTEP binding agreement) and/or the company or other collaborator (for Alliance studies that are under other binding agreements).

16.2.7 Presentation of results by treatment group

The DSMB assesses relative efficacy according to the protocol specified interim analyses; therefore results by treatment are presented and discussed. No treatment-specific results, coded or not, are released to anyone not on the DSMB.

16.2.8 Phase 2/3 trials

With respect to implementation of phase 2 decision rules in phase 2/3 designs of clinical trials, any protocol-specified phase 2 decision-rule analysis must be performed within six weeks from the date the required number of events are observed and reported in the database. If the trial follows the decision rule (i.e., continues or stops depending on whether the continuation threshold is met), then the Alliance notifies the DSMB and chief, CIB of the status of the trial (i.e., continuing or stopping) based on the protocol-specified phase 2 decision rule. In the unlikely event that the study statistician wishes to request permission not to follow the protocol pre-specified decision rule, such a request must first be discussed with NCI/DCTD/CTEP by conference call within two weeks of the required number of events being observed / reported in the database. This request (change in the design of the trial) needs to be approved by the CTEP associate director or his/her designee in consultation with the chief, CIB who will notify the Alliance Chicago Office in writing of NCI decision regarding the request. If NCI/DCTD/CTEP is willing to approve the request, the Alliance must then seek DSMB approval within three weeks on receiving NCI/DCTD/CTEP approval before submitting an official amendment to CTEP’s Protocol and Information Office to change the design of the trial regarding the phase 2 decision rule.
16.2.9 Industry-supported studies

Studies supported by industry are also covered by these policies and procedures. Industry representatives may not serve on the Alliance DSMB.

16.2.10 Conflict of interest

Individuals invited to serve on the DSMB are subject to the Alliance Conflict of Interest policy (see section 3.5).
16.3 Monitoring phase 1 and 2 studies

16.3.1 Phase 1 studies

Phase 1 studies are ordinarily limited access studies. Routine monitoring is carried out via conference call among representatives of each participating institution, the primary statistician, the study chair and members of the study team. A representative from each institution must participate whenever the institution has any participants currently receiving protocol therapy. Institutions that fail to submit toxicity data as required or that do not participate in the conference calls will be prohibited from continuing to enroll participants on the study.

16.3.2 Phase 2 studies

Non-randomized phase 2 studies are routinely monitored by the study team (study chair, primary statistician, executive officer, protocol coordinator, data management personnel) and other Central Protocol Operations Program staff (e.g. director of regulatory affairs) as applicable. Each phase 2 protocol must specify the efficacy and adverse event monitoring plan to be used.
Chapter 1 Introduction

Summary of Changes

Section 1 Introduction Committee membership

Added new 3rd paragraph “The Alliance receives grant funding from the National Cancer Institute (NCI). The Alliance is one of the Network Groups for the NCI National Clinical Trials Network (NCTN) and serves as a research base for the NCI Community Oncology Research Program (NCORP). The Alliance complies with the NCTN and NCORP Program Guidelines, related NCI policies and procedures and the Code of Federal Regulations (CFR). As an NCTN and NCORP Group, the Alliance utilizes centralized NCI systems for the management of clinical trials.”

Section 1.1 Specific aims
Changed 55 years to “60 years” in the first sentence

Section 1.1.2 Operational aims
Added “and NCORP” to the first sentence: “The infrastructure of the three component cooperative groups has been merged into a single, fully integrated system that is optimally designed to serve the NCTN and NCORP research community”

Section 1.2 Overview of program structure Conflict of Interest
Added “principal investigators” to the 3rd sentence of the first paragraph “The Alliance is also supported by five program directors/principal investigators, each responsible for a specific program integrating discipline-related science and operational functions across all disease committees.”

Minor updates to table 1-1 Alliance program structure

Figure 1-1 Alliance Leadership – Remove “Patient” from position title “Associate Group Chair Patient Advocacy”

Section 1.2.1 Office of the Group Chair

Added “communications” as task of the office of the group chair in the 1st paragraph
Deleted the following sentence from the 2nd paragraph “The group vice chair is the chair of the Publications Committee, and has primary responsibility for Alliance manuscript review and for coordinating manuscript authorship and timely submission.”

Added sentence to 2nd paragraph “The associate group chair for Advocacy promotes patient advocacy initiatives.”

Deleted “Mayo Clinic in Rochester, MN” Figure 1-2 Office of the Group Chair
• Moved Publications Manager, and Publications Coordinator position under Group Chair,
• Added Regulatory Compliance Manager
• Deleted Regulatory Affairs Manager, Roster Supervisor, Roster Staff, and Audit

Alliance Policies and Procedures – Appendices A1
Managers
Deleted the following text “As mentioned above, the group vice chair oversees publications processes and compliance with Alliance publications policies and timelines.” “Finally, the associate group chair for Patient Advocacy provides an important representative from the Office of the Group Chair to patient advocate organizations outside of the Alliance.”

Section 1.2.2 Statistics and Data Management Program
Removed office locations, as we do not need that much detail in the P&P. Deleted specific locations of Ohio State University and MD Anderson.

Section 1.2.3 Central Protocol Operations Program
Changed title of Figure 1-4 Protocol Office to “Central Protocol Operations Program”
Added “Executive Officer” “Pharmaceutical Affairs Manager” and “Associate Director” to Figure 1-4

Section 1.2.4 Translational Research Program (TRP)
Figure 1-5 Translational Research Program – Deleted ‘Director Solid Tumor’ and ‘Director Leukemia Correlative Science.’ Added ‘Operations’ to ‘Director Translational Research,’ Changed ‘Director’ to ‘Chair Pharmacogenetics and Population Research.’ Added ‘Executive Officer’

Section 1.2.6 American College of Surgeons Clinical Research Program
Updated language to make current. Deleted the following text “The ACS Clinical Research Program is an innovative program that meets three important Alliance needs. First, it provides partnerships and infrastructure that enable comparative effectiveness research, a burgeoning field in an era of health care reform and a particularly useful approach to addressing surgical questions in cancer treatment. Second, this program allows Alliance investigators to use the NCDB to inform the design of both treatment and comparative effectiveness studies. This resource is particularly effective as a tool to assess accrual potential for studies of rare tumors, or those with a challenging randomization plan for which knowledge of regional practice patterns is essential to study planning. Finally, the program’s Member Services and Education Committees provide support that integrates surgical researchers into the Alliance. This outreach is particularly important to maintaining involvement of community surgeons in Alliance work, as they are less likely than medical or radiation oncologists to have in-house support for clinical research.”

Section 1.2.7 Member institutions
Added the following text to the 1st paragraph “Alliance member networks may be Lead Academic Participating Sites (LAPS) or NCORP networks. LAPS and NCORP institutional networks receive grants from the NCI to support their infrastructure and participation in NCI-funded clinical trials. Non-LAPS and non-NCORP institutions receive per-case payments from the Alliance NCI-grants to support their clinical trial participation.”

Section 1.3.1 Scientific committees
Editorial changes

Section 1.3.2 Administrative committees
Deleted “Study Concept Review”
Chapter 2 Institutions

Summary of Changes

Section 2 Institutional Membership
Minor change deleted “Institutional” from first sentence

Section 2.2 Applying for membership
Updated frequency of application review to “monthly or as needed” was previously “twice monthly.”
Added location of membership application on the Alliance website. Added language stating applications will be reviewed by Membership Committee ‘only if the institution has an active NCI ID and FWA”

Added the following text “Affiliate applications can be approved by the Membership Committee without Board approval.”

Section 2.3 Membership activation

Additional text regarding PI and Lead CRP added to the first paragraph “Alliance staff will add the PI and Lead CRP to the institution roster(s). The Lead CRP will add persons and person roles to the institution roster via the NCORP Management System (NCORP SYS) or CTSU Roster Update Management System (RUMS).”

Section 2.3.1 Roster

Deleted the first sentence “Alliance complies with the NCIs Unified Site Code Policy”

Added additional language regarding criteria for including site on the Alliance roster “A site must be included on the roster if it meets the following definition of engagement in research as defined by OHRP (45 CFR part 46). An institution is engaged in a particular non-exempt human subjects research project when its employees or agents for purposes of the research project obtain: 1. Data about the subjects of the research through intervention or interaction with them 2. Identifiable private information about the subjects of the research. or 3. The informed consent of the human subjects for the research.”

Deleted 5 bullet points at end of paragraph “•Direct receipt of CTEP agent • Enrollment of patients/research participants • Institution’s whose employees, representatives, and/or agents are authorized to obtain informed consent from patients • Direct receipt of federal funds • Directly responsible for submission of data to the study sponsor or their designee”

Section 2.3.2 Regulatory Documentation

Deleted last sentence of the 1st paragraph regarding institutions needing documentation that a procedure in place to notify patients of new information regarding toxicities and outcomes, and all members acknowledging the Individual Scientific Misconduct Policy”

Added new 3rd paragraph detailing Registration and Credential Repository (RCR) Requirements
“NCI policy requires all persons participating in any NCI-sponsored clinical trial to register and renew their registration annually. Registration is accomplished via the NCI Registration and Credential Repository (RCR). Additional details can be found on the NCI/CTEP website.”

Section 2.3.3 Added “for institutions” to section title

Section 2.4 Responsibilities of the main member

Added as first paragraph “The principal investigator will be required to sign a membership agreement that includes a summary of key policies and procedures, including conflict of interest, scientific misconduct, membership accrual requirements, confidentiality, audit requirements, institutional performance and publications.”

Section 2.4.2 Electronic communication

Added sentence to end of paragraph “All network and site PIs, Co-PIs, Lead CRPs and Secondary Lead CRPs are required to receive broadcast emails.”

Section 2.4.5 Human subjects protection

Added new sentence at end of paragraph “Protocol-specified research interventions, including interventions conducted at a facility external to the registering institution, must be covered under an IRB approval.”

Section 2.4.6 Training

Added new sentence at end of paragraph “The Alliance conducts education and training sessions during the Alliance Group meetings and posts educational resources on its website. All Alliance members are encouraged to participate in these training opportunities.”

Section 2.5.1.1 Network responsibilities

Added new 4th paragraph “Each affiliated institution in a network must name a responsible principal investigator. This PI may be the main member PI or another investigator responsible for clinical trial conduct at the affiliate institution with oversight from the main member PI.”

Section 2.5.1.2 Institutional responsibilities

Added new sentence to the last paragraph “The PI ensures that the delegation of authority and tasks is documented and that research personnel are adequately trained.”

Section 2.5.2 Affiliate member principal investigator

Added “human subjects protection” to first sentence of first paragraph “The principal investigator (PI) for an affiliate institution is responsible for the conduct of Alliance activities at an Alliance institution, human
subjects protection and the integrity of all data submitted from the institution.”

Section 2.5.3 Clinical research professionals

Added new 3rd bullet “Maintain study-specific regulatory and training files

Section 2.5.4 Cytogeneticist

Deleted entire section

Added new section 2.5.4 Withdrawn or terminated institutions

“If an institution is withdrawn from the Alliance or terminated by the Alliance, the institution will remain responsible for data submission until such time that there are no longer patients in treatment or follow up, or the patient(s) are transferred to another Alliance member. The main member remains responsible for data from withdrawn affiliates.”

Section 2.6.2 Reporting institutional assurance compliance

Added new sentence to the 1st paragraph “The institution’s FWA must be included with the member’s roster information and remain current”

Deleted “This information is entered into the CTSU/RSS database and is referred to when a patient is being registered. Documentation must state the type of review, list the protocol number (and if it is a review of a protocol update, it must list the protocol update number) and an IRB member or administrator must sign it. The protocol number and the update number, if applicable, must be clearly documented. Initial and continuing review documents must be submitted to the Cancer Trials Support Unit (CTSU) and Alliance staff will access the information in the CTSU database.” (moved to section 2.7.1)

Section 2.7.1 Reporting requirements – changed section name to “Reporting and review requirements”

Deleted first paragraph “Any substantive changes of information concerning risks or alternative procedures and/or translational research contained in the model informed consent document must be justified in writing by the investigator. Investigators must forward copies of such changes, with their justifications, to the Alliance for review.”

Added the following text after the first sentence “IRB approval documentation is submitted to the CTSU. This information is entered into the CTSU/RSS database and is referred to when a patient is being registered. Documentation must state the type of review, list the protocol number (and if it is a review of a protocol update, it must list the protocol update number) and an IRB member or administrator must sign it. The protocol number and the update number, if applicable, must be clearly documented. Initial and continuing review documents must be submitted to the Cancer Trials Support Unit (CTSU) and Alliance staff will access the information in the CTSU database.”

Section 2.8 Institutional Audits

Alliance Policies and Procedures – Appendices A5
Extensive changes made throughout the entire section to reflect the new CTMB Audit Guidelines. Please refer to tracked changed version of the Policy and Procedures

Section 2.9.2 Affiliates

Minor changed moved “…five patient registrations per year based on a three-year rolling average.” to the first paragraph

Section 2.10.1 Institutional Network Performance Scoring

Updates to Table 2.20 IPEC Scoring Quality, 2.21 IPEC scoring for timeliness

Section 2.11.3 Probationary process

Minor changed “affiliated” to “individual network sites” throughout section

Section 2.11.4.1 Implication of probationary status

Minor changed “affiliated” to “network institution” throughout section

Added new Section 2.13 Non-member Collaborators

“Non-member collaborators (NMCs) are institutions or networks that participate on Clinical Therapy Evaluation Program (CTEP) and Division of Cancer Prevention (DCP) sponsored protocols but are not full member institutions of the Alliance or a participating organization. Most non-member collaborators with the Alliance are international organizations.

In addition to their own country’s regulations, International groups must comply with US federal regulations such as:

- Obtaining Federalwide assurance (FWA) with the Office for Human Research Protections (OHRP); and

- Obtaining State Department Clearance. The Alliance will submit State Department Clearance to the NCI on behalf of the international collaborator.

NCI policy also requires all persons participating in any NCI-sponsored clinical trial to register and renew their registration annually. Registration is accomplished via the NCI Registration and Credential Repository (RCR). Additional details can be found on the NCI/CTEP website.”
Chapter 3 Participants

Summary of Changes

Section 3 Participants

Reorganized sections 3.1-3.2

Added Section 3.1 titled Participant Categories. Text has not changed

Added Section 3.2 “Membership and participant registration”

Section 3.2.1 “Applying for membership and registration”(formerly section 3.1, Applying for membership in Alliance)

Deleted the first sentence “Individuals apply for membership as Alliance institutional members via their principal investigator”

Deleted 3rd sentence “The principal investigator submits a request for membership for all prospective physician members, along with a copy of their curriculum vitae, 1572 and a completed Roster Update Form.”

Modified the last sentence in the first paragraph to “The lead Clinical Research Professional (CRP) or Secondary Lead CRP is responsible for adding and withdrawing all institutional members via CTSU RUMS or NCORPSys and include the Human Subject Protection Training type and date completed”

Added new paragraph describing Registration and Credentialing Repository Requirements

“NCI policy requires all persons participating in any NCI sponsored clinical trial to register and renew their registration annually. Registration is accomplished via the NCI Registration and Credential Repository (RCR). RCR utilizes FIVE person registration types:

- **Investigator (IVR)** — MD, DO, or international equivalent
- **Non-Physician Investigator (NPIVR)** — advanced practice providers (e.g., NP or PA) or graduate level researchers (e.g., PhD)
- **Associate Plus (AP)** — clinical site staff (e.g., RN or CRA) with data entry access to CTSU applications (e.g., RUMS, OPEN, RAVE, TRIAD)
- **Associate (A)** — other clinical site staff involved in the conduct of NCI-sponsored trials
- **Associate Basic (AB)** — individuals (e.g., pharmaceutical company employees) with limited access to NCI-supported systems”

All persons applying for Alliance membership must obtain an NCI/CTEP-IAM account, access the RCR system, and complete an annual NCI person registration.

Additional details are available on the CTEP website [https://ctep.cancer.gov/investigatorResources/default.htm](https://ctep.cancer.gov/investigatorResources/default.htm). Alliance leaders and committee chairs may
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request special membership for an individual. The request is sent to the Office of the Group Chair.”

Section 3.2.2 “Alliance person database” (formerly section 3.2)

Modified second sentence: The institutional principal investigator and the lead CRP are responsible for ensuring that the roster of institutional members is accurate and up-to-date, utilizing the CTSU Roster Update Management System (RUMS) and providing timely notification to Alliance of changes to PIs and lead CRPs.

Section 3.5 Conflict of Interest

The Conflict of Interest policy was updated to include collection and review of study-specific COI disclosures from institutional investigators, in addition to the main member PI. Note: Study-specific COI disclosures may be collected on select studies, not all studies.

The COI policy was updated to implement additional review considerations determined by the Alliance COI committee related to disclosures from Alliance leadership. These considerations include multiple disclosures that individually do not meet thresholds set forth in the current policy. Multiple disclosures of >$5,000 are subject to review by the Alliance COI Committee. The Committee may request a management plan including oversight by co-leaders. The policy was also updated to strengthen the language regarding public disclosure of potential COI.

Section 3.5.1.1 Introduction (COI)

In the 2nd paragraph deleted “Study Concept Review Committee” added “institutional investigators”

Section 3.5.1.3 Committee chairs/group leaders/Alliance staff

Changed section name to “Committee chairs/group leaders/institutional investigators/Alliance staff”

Added new paragraph at end of section “In addition to main member principal investigators, institutional investigators participating in an Alliance study may be required, on a study-specific basis, to disclose financial arrangements as defined in this policy.”

Section 3.5.1.4 Data and Safety Monitoring Board and Study Concept Review Committee members

Changed section name to “Data and Safety Monitoring Board”

Deleted “and Study Concept Review Committee (SCRC) from the 1st paragraph

Section 3.5.3.2 Proprietary interest

Added to first sentence “the investigator is receiving compensation that could be affected by study outcome such as compensation that is explicitly greater for a favorable result”

Deleted section “An investigator with financial arrangements >$25,000/year in a privately held business

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or equity interest in a publicly traded company sponsor that exceeds $50,000/year, or ≥5% ownership interest (including common stock) in either a privately held or publicly traded business would be prohibited from assuming chairmanship of a study. An investigator with financial relationships >$25,000/year or equity interest in a publicly traded company sponsor that exceeds $50,000/year, or ≥5% ownership interest (including common stock) in either a privately held or publicly traded business, who also serves on the Executive Committee must recuse themselves from participation in the deliberations of the Executive Committee where a conflict or appearance of conflict of interest may exist.”

Section 3.5.3.3 Miscellaneous

Changed section name to “Miscellaneous and multiple financial interest”

Added the following

“Alliance leaders may have individual financial interests related to industry partnerships or other affiliations that do not exceed the threshold of $25,000. Multiple disclosures of >$5,000 are subject to review by the Alliance COI Committee. The Committee may request a management plan including oversight by co-leaders.

Committee chairs with financial interests in products, actively under investigation or proposed in committee sponsored studies, may be required to publicly disclose potential conflicts and/or recuse themselves from relevant discussions.

Committee chairs with financial interests exceeding thresholds defined in this policy may be subject to management plans and restrictions, per section 3.5.4 below.”

Section 3.5.4 Management plan for conflicts of interest

Modified first paragraph to the following: “Prior to concept submission, study activation, as financial arrangement change and at least annually, all members of the study leadership team are required to complete a Conflict of Interest Form as described above.”

Deleted first sentence of the 1st bullet “Independent review of studies by network group leadership beyond the sponsoring committee will be undertaken.”

Added 5th bullet with the following language “Financial conflict disclosures of institutional investigators are subject to institutional conflict of interest policies. The Alliance may request a mitigation plan from investigators exceeding thresholds, including documented institutional management plans in compliance with institutional requirements. Independent review of studies by network group leadership beyond the sponsoring committee will be undertaken.”

Added “or direct employment with an industry partner” to the following paragraph “In the event of conflicts exceeding the $25,000 annual threshold or equity interest in a publicly traded company sponsor of $50,000 annual threshold, or ≥5% ownership interest (including common stock), or direct employment with an industry partner, the following policies will be enacted.”

Added text to 8th bullet “When a conflict exists for the committee chair or vice-chair the committee

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leader may not serve as either first, corresponding or senior author. If all of the key individuals of a study show a significant conflict of interest such that they are ineligible, then the disclosures are sent to COI committee for review.”

Added 9th bullet “The Alliance may disapprove study participation of institutional investigators exceeding maximum thresholds, upon review of the institutional plan to mitigate bias.”

Section 3.5.8 Public disclosure

Added the following text “Financial conflicts of interest must be disclosed during Alliance committee meetings, including study development discussions.”

Added new section 3.5.11 Alliance Conflict of Interest Committee

The Alliance Conflict of Interest Committee is a volunteer committee comprised of Alliance investigators. The committee reviews financial conflict of interest disclosures related to trials supported by the Alliance and Alliance for Clinical Trials in Oncology Foundation.
Chapter 4 Committees

Summary of Changes

Section 4.3 Committee membership
Removed the following text “There is no application for committee membership”

Section 4.4.2.1 Conflict of Interest
Added frequency language to state conflict of interest disclosure form is required at least annually

Section 4.4.2.1 Training
Deleted the following text “Training: The committee chair completes the Alliance Study Chair Workshop training modules, upon appointment and upon request of the Central Protocol Operations Program (CPOP) Director and informs the study chairs (past study chairs who have not published a manuscript, current study chairs and pending study chairs) within the committee that each one is also expected to complete the modules. The CPOP office notifies study chairs and committee chairs as to who is expected to attend.”

Section 4.4.2.4 Intergroup collaborations
Deleted the following text “Refer to the National Clinical Trials Network Program’s Guidelines for Development, Conduct and Analysis of Clinical Trials with International Collaborating Institutions.”

Section 4.4.2.5 Finance
Deleted the title “chief financial officer” from the Funding to support research project paragraph, as discussion of project budgets with the CFO is not required in all situations.

Changed text in 3rd paragraph “Executive Committee” to “SCRC” in the following sentence “Details concerning the proposed funding are included with the concept when it is submitted to the Executive Committee for review.”

Deleted “…when the Executive Committee reviews it.” From the 3rd paragraph

Section 4.4.4 Subcommittee chairs
Changed subcommittee chair to ‘cadre leader’ in text.

Section 4.4.5 Committee members
Deleted the following text “Committee members are appointed for three-year renewable terms, with no limit to the number of terms. The committee chair may also appoint leaders to coordinate the activities of a subsection of the committee.”

Section 4.5 Electing Executive Committee members
In 1st paragraph changed “CCOP” to “cancer control”
2nd paragraph updated language regarding term limits to indicate representatives on Executive Committee may serve three year terms, and only be elected for three consecutive term, per the Alliance Constitution.

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Chapter 5 Meetings

Summary of Changes

Section 5.1 Group meetings
Editorial changes, and added clarifying language regarding location of travel information

Section 5.1.2 Travel funding for group meetings
Added clarifying language regarding travel funding

Section 5.2 Committee meetings
Deleted section on committee meetings and renumbered subsequent sections

Section 5.2 Identification of funded travelers and expense reports
Editorial changes to remove references to committee meetings and clarify expense report requirements

Section 5.3.2 (previous section number) Discretionary funding
Section deleted

Section 5.3 (new) Continuing Education Credit
Added section 5.3 “Continuing Education (CE) Credit”
Chapter 6 Study Protocol

Summary of Changes

Section 6.1.2.1 Companion studies
Added clarifying language regarding companion studies “Companion studies with separate study numbers do not necessarily have to be published at the same time as the parent study, and may be published as a distinct manuscript.”

Added clarifying language regarding necessity of the component having a separate study chair “who is not the parent study chair”

Section 6.2.1 Limited access studies
Added the following paragraph “As per NCI requirements, limited access studies may not include members outside of the Lead Participating Organization. Permission for the addition of institutions outside of the Lead Participating Organization to limited access studies must be obtain from the NCI.”

Section 6.3.2 Study co-chair
Added clarifying language to existing section regarding role of study co-chair in protocol development “It is expected that study co-chairs contribute in a meaningful way to the study conduct, for example, by answering questions from institutions related to their role on the study. Study co-chairs are responsible for the section of the protocol specific to their modality, such as surgery, imaging, radiation, etc. Identification on the protocol face page of a study co-chair will not assume to lead to authorship.”

Section 6.3.5 Data managers
Changed data personnel to data managers

Section 6.3.6 Protocol Coordinator
Modified language in first paragraph to the following “Protocol development occurs under the direction of the protocol coordinator. Protocol coordinators will establish timelines for protocol development, work with study team members to draft, review and revise the protocol. They serve as the liaison for all protocol related correspondence with CTEP, DCP and CIRB, and are responsible for communicating official CTEP, DCP or CIRB communications to study team members.”

In the 2nd paragraph, clarified post study-activation activities of the protocol coordinator

Section 6.3.7 Executive officer
Editorial changes to clarify role of executive officer and update AdEERS to CTEP-AERS

Section 6.4.1 Table 6-1 Alliance protocol numbering system
Deleted “Comparative Effectiveness Research, A18, A181101” from the table

Section 6.4.2.1 Concepts other than translational research and data-only requests
Deleted the following language “In addition, the concept must have been reviewed by TRP prior to submission to the SCRC even if no translational research has been planned”
Added language regarding Cancer Control to the 2nd paragraph “Cancer control studies (e.g., non-treatment studies) do not have an NCI-specific concept submission form, and are to be submitted to the Alliance SCRC in the same format as required for concept submission to NCI DCP.”

Section 6.4.2.2 Translational Research Concepts
Deleted section since translational research is captured in chapter 11 and renumbered subsequent sections

Section 6.4.2.2 (new section number) Concepts containing data-only requests
Clarified language

Section 6.4.3.1 Communications post-concept approval
Added clarifying language regarding process for DCP “The Alliance Cancer Control Program Manager submits concepts to DCP for approval.”

Section 6.4.3.2 Protocol authoring
Editorial changes and deletion of 4th paragraph regarding translational research

Section 6.4.3.3 Determining the trial participant eligibility criteria
Added language from 6.4.3.4 “Alliance studies typically require trial participants to be at least 18 years old. In certain diseases, younger patient populations may be considered.”

Section 6.4.3.4
Text deleted “Alliance studies typically require trial participants to be at least 18 years old. In certain diseases, younger patient populations may be considered.” and moved to section 6.4.3.3

Section 6.4.3.6 External protocol review
Clarified the types of protocols submitted to CTEP or DCP “Phase III, select phase II, and select cancer control trials…”

Section 6.4.4.3 Using Translated Patient-Reported Outcomes
New section added “The most commonly used patient-reported questionnaires for Alliance protocols will be made available in the North American primary languages, i.e., English, Spanish, and French Canadian. If a translated questionnaire is not readily available, the study chair must choose between: 1) restricting participation to English speakers only or 2) allowing accrual of patients with other non-English primary languages. If option 2, then the study chair must decide whether to: 1) pursue formal translation of the questionnaire or 2) allow on the spot translation by either professional translators at the institution or the patient’s family/friends. The Alliance preference is to design all Alliance studies to allow accrual of patients with other non-English primary languages using on the spot translation by either professional translators at the institution or the patient’s family/friends. The Alliance Model Protocol Template includes the appropriate information for this option.

However, if a formal translation is requested, the investigator must send an email request to QOL@alliancenctn.org. All translation requests will need to be reviewed and approved by the Cancer Control Program (CCP) leadership.”

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Section 6.4.5 Participation in intergroup studies
Deleted text “There are two mechanisms for intergroup participation: either by placement of a study on the CTSU menu (most phase 3 studies), or by agreement between one or more network groups to participate in a protocol. Each intergroup study coordinated by Alliance has co-chairs designated by the other groups. These individuals must be adequately informed about progress and problems with the protocols for which they are responsible.”

Added the following paragraph “With few exceptions, all studies are to be available to all members of the NCTN. Exceptions may include certain DCP sponsored studies and selection phase I or early phase II studies. Studies may have co-chairs from other groups who were involved in the study design added to the protocol. These individuals must be adequately informed about progress and problems with the protocols for which they are responsible.”

Section 6.5 Activating a study
Deleted last sentence in section “All Alliance treatment and intervention trials will be accessible from the member portion of the CTSU website”

Section 6.7.1 Revisions and amendments
Editorial changes and deletion of requirement for SCRC review

Section 6.8 Suspending a study
Deleted “surrogate” from the last sentence

Section 6.9.1 Emergency unblinding
Clarified language regarding emergent unblinding requests: “Emergency unblinding requests should be directed to the executive officer on call, 24 hours a day, 365 days a year. If an Alliance executive officer determines unblinding is warranted, they will contact the registration office staff. The executive officers are the only personnel who can unblind a study patient.”

Sections 6.10 – 6.14
Repaginated
Chapter 7 Patient Registration

Summary of Changes

Section 7.1 Authorization to register patients
Moved section regarding authorization of institutions to register patients from 7.3 to 7.1

7.2 (new section number) Authorization of participants to register patients

Added the following sentence to the 2nd paragraph “Registration is accomplished via the NCI Registration and Credential Repository (RCR). Refer to the CTEP website for additional details on registration types and required documentation.

Section 7.3 (new section number) Credentialing
Renumbered from 7.2 to section 7.3
Chapter 8 Data Management

Summary of Changes

Section 8.1.1.1 Alliance general instructions: all forms (electronic CRFs and paper forms)
Deleted “ACOSOG,” and “or faxed to the CTSU for appending to the form in Rave,” editorial changes

Section 8.1.1.2 Instructions for forms submitted during treatment and follow-up
Under Follow up and response forms
- Deleted “by combining target lesions, non-target lesions, target lymph node lesions, non-target lymph node lesions and any new lesions or sites of disease.”
- Deleted “Once a patient achieves a “PR” or “CR” they remain a “PR or “CR” until the criteria for progression are met.”

Under Adverse event forms deleted:
- Deleted “forms, and data entry screens from the last sentence of the first bullet
- Deleted the 3rd bullet “Code the grade that reflects the most severe adverse event or most abnormal lab value occurring during the reporting period’ You will never have more than one AdEERS report per cycle”
- Deleted the 6th bullet Each event included in the CTCAE is linked to a MedDRA code. Refer to the protocol for the MedDRA code version. Don’t assume that all hospitalization require AdEERS reporting—the guidelines are set by whoever hold the IND,”
- Deleted 7th bullet “The “Surgical Intervention” section is to be used ONLY for the protocol related surgery.”

Under Adverse Event Reporting System
- Changed “AdEERS” to “CTEP-AERS” throughout section
- Added bullet “Don’t assume that all hospitalization require CTEP-AERS reporting— check the protocol.
- Added bullet “The “Surgical Intervention” section is to be used ONLY for the protocol related surgery.”

Section 8.1.2.1 General data submission instructions
Overdue data section
- Added new paragraph “The current expectations for form submission before being considered delinquent are: Baseline and treatment forms: within 30 days of target date, Follow-up Forms: within 60 days of target date.”

- Deleted text “Delinquent data submission of greater that 3 months but less than 6 months is a Lessor deficiency in an audit. Greater than 6 months is a Major deficiency.”

- Clarified last paragraph to describe study specific delinquency lists, added more detail to describe tools available on the Alliance website

New malignancies section
• Clarified first sentence to indicate all new malignancies that “fall within the protocol specified time period” must be reported and deleted end of sentence which described where it should be reported
• Deleted “For legacy ACOSOG, CALGB, and NCCTG studies, submit new malignancy data as specified in the protocol”

Updates to table 8-1: Addition of clarifying language

Section 8.1.2.2 Registered patients who never receive treatment (canceled patients)
Deleted “Upon receipt of the documentation, a data manager reviews it and determines the status of the patient” from 2nd paragraph

Section 8.1.2.3 Transfer of patients to another institution
3rd paragraph changed to “Prior to the transfer, the site clinical research professional (CRP) ensures that all data are up-to-date and all queries have been addressed and resolved. This will be confirmed by the Alliance Data Manager prior to the patient being officially transferred. Copies of all data required by the protocol and subject records must be submitted to the accepting institution”

Added the following paragraph “Once the data are updated the site is required to call the Alliance Registration Office for official documentation of the transfer and transfer of responsibilities.”

Last two paragraphs changed “via the CTSU” to “OPEN”

Section 8.1.2.5 Confirming of lost to follow-up status

Deleted the following text from the 1st paragraph “Patients that are confirmed lost to follow-up are removed from calculations of institutional performance related to timeliness. However, the percentage of patients deemed lost to follow-up is included in the metrics for institutional performance related to data quality.”

Section 8.1.2.5.1 Procedures for confirming a patient is loss to follow-up

Under the second bullet deleted text “Check the Social Security Death Index (http://www.stevemorse.org/ssdi)”

Fourth bullet deleted text regarding requirement to send certified mail “by certified mail (https://www.usps.com/ship/insurance-and-extra-services.htm) with request for return receipt to the patient at the last known domestic address, or by registered priority international mail with request for return receipt.” Added text “A diligent effort to contact the patient is required and should be documented.”

Deleted text regarding the Alliance Confirmation of Lost to Follow up form requirements for patients lost to follow up “…which asks for limited details about the procedures used by the institution. The form must indicate both of the following items: 1) No date of death can be found in the Social Security Death Index. 2) The certified or registered international letter has been returned unclaimed or marked addressee unknown, or it has been received (as documented by return receipt) but has resulted in no
response from the patient over a one month period after it was received.”

Deleted the following text: “For an audit, evidence of diligent effort to contact the patient may include: A copy of the Web page from the Social Security Death Index indicating that no information was found for the patient, and A certified or registered international letter returned to the institution unclaimed or marked addressee unknown. Alternatively, if the letter is delivered, but a response from the patient is not obtained by the institution within one month, the return receipt acknowledging the letter was received is evidence of due diligence. Because certified and registered international mail do not require the signature of a specific individual at the destination address, date of receipt of the letter should not be used to update the survival date of the patient. Even if the signed name on the return receipt for the certified or registered international mail appears to be that of the patient, it is not appropriate to update survival based on the signature on the returned receipt because of the level of uncertainty in signature verification. A telephone log or other documentation of attempts to contact the patient via telephone.”

Section 8.1.2.5.2

Continuing responsibilities for data submission - Deleted section “Patients that are confirmed lost to follow-up are removed from calculations of institutional performance related to timeliness. However, the percentage of patients deemed lost to follow-up is included in the metrics for institutional performance related to data quality (see section 2.10).”

Retrospective data submission - Added the following text “the patient is deemed lost to follow-up. For the period of time between the last contact with the patient and the date they are deemed lost to follow-up, the site must record in Rave that no contact occurred including the date of the attempt to contact the patient” Deleted “…of last contact with the patient”

Section 8.2 Receipt and distribution of data forms by SDC

Added clarifying language to 2nd paragraph: “Data for studies coordinated by other network groups are submitted directly to the coordinating group via the instructions outlined in their data submission section of the protocol.”

Section 8.3 Quality assurance performed by Data Management

Added clarifying language to 3rd sentence of first paragraph

Section 8.3.1 Quality checks of on study and eligibility data

Modified first paragraph

New Language: Quality checks of on-study data include a detailed review of eligibility criteria and supporting documentation requested in the protocol. The first eligibility review is performed via the OPEN registration system. Upon receipt of the eligibility material and supporting documentation the DM performs a second quality check.

Section 8.4 Alliance Case Evaluation Process

Repaginated
Chapter 9 Information Systems

Summary of Changes
Section 9.1 Member information
Minor changes

Section 9.1.1.1 Individual institution members
Clarified language regarding registering new members. Added and deleted text in the last sentence (new text in bold below):

“When the application is approved, appropriate accounts are created in the Alliance Information Systems. The member’s CTEP username and password is used to access the Alliance member site and SMU/ISU Web applications.”

Deleted the following text from the last sentence “and the member is given a username and password for accessing the Alliance member site and SMU/ISU IS Web Applications.”

Section 9.1.6 System Availability
Added text to the first sentence to clarify availability of Alliance systems (new text in bold below): “All Alliance systems are available 24 hours a day, seven days a week, with exceptions for system maintenance. Whenever possible, system maintenance will occur on a planned basis, with one week notice provided to Alliance members. Unscheduled maintenance may occur as needed to resolve critical security vulnerabilities or to resolve other critical systems issues.”

Deleted the following text from the first sentence: “the exception of scheduled maintenance, which occur on a planned basis with a week prior notice provided to Alliance members”

Minor edits to 2nd paragraph

Section 9.1.7.2 Emergency calls for systems support
Deleted entire section “During business hours, contact the Alliance Service Center. For emergencies outside of business hours, call the Alliance on-call telephone number shown on the Alliance website under the ‘Contact’ heading. Trained SMU/ISU employees respond to application support emergencies as soon as possible.”

Section 9.2.1 Software development
Minor changes to first paragraph to clarify SMU/ISU activities. Added text after the first sentence in the 2nd paragraph to describe software development (new text in bold below):

“ISU uses a tiered software development environment to ensure proper testing and migration from the development to production environments. Software is first deployed to a development environment for initial testing by the software development staff. Software is subsequently deployed to an integration environment for software quality assurance and user acceptance testing, prior to being released into the production environment. New software is deployed during scheduled downtimes unless they are deemed urgent or critical, in which case the software release is migrated as soon as possible.”

Section 9.2.3 Technology selection and change management
Deleted Committees no longer active: Alliance IT Committee, Alliance SMU/ISU Leadership Committee
Chapter 10 Publications

Summary of Changes

Section 10.4.3.1 Publication on the primary study endpoint

Added the following sentence to the 3rd paragraph “If the modality co-chair participated in the design of the study and wrote the modality section of the protocol, they should be an author on primary endpoint publications.”

Section 10.4.3.2 Publication on a secondary (correlative) study

Authorship on publications of a secondary study included in the original Alliance or legacy protocol

Added text to the first bullet “Authorship by a modality co-chair on secondary endpoint publications should be a function of their involvement in the secondary analysis.”

Added 4th bullet “Pathologists, radiologists and other specialists who perform quality assurance (QA) for the study, unless the publication is independent of QA results of their findings.”

Authorship on publications of a secondary study not in an original Alliance or legacy protocol; study proposed by Alliance investigator

Added 2nd bullet “Pathologists, radiologists and other specialists who perform quality assurance (QA) for a study, unless the publication is independent of QA results of their findings.”

Added “QA specialist” to paragraph “After primary study chair(s), primary statistician(s), QA specialists and researchers, other investigators who were involved in the primary study or studies may not necessarily be included in secondary study publications; instead, authorship is determined by an individual’s contribution specific to the secondary study and by ICMJE”

Section 10.5.4 Timelines for review and revisions of manuscripts submitted to the Alliance publications coordinator

Added text to 2nd paragraph “All abstracts and manuscripts (except those resulting from data sharing) must be reviewed by an independent Alliance faculty statistician.”

Added new Section 10.5.5 Approval of abstracts and manuscripts

All comments received from reviewers during Group Review are sent to the chair(s) of the Alliance Publications Committee. The Publications Committee chair(s) are responsible for approving abstracts and manuscripts, or requesting revisions followed by re-review.

Section 10.8 Press Release

Changed name of the section to “Publicizing Research Information“

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Modified section as follows: “All communication related to the dissemination of Alliance research to external audiences is handled by the Alliance communications specialist. This includes all written or recorded communication (i.e., press releases, news releases, press statements, video releases) directed to members of the news media, stakeholders, and the public, regarding the activation, progress, results and findings of Alliance research. This also relates to all communication generated by an institution or industry partner based on Alliance research. Such communication must be submitted the communication specialist (communications@AllianceNCTN.org) for review at least one week prior to its release. Also refer to Section 14.3, Dissemination of Information to the General Public.”

Section 10.10 NIH Public Access Policy compliance

Deleted the “Method used by journals” section of the table

10.11 Quick view of Alliance publication timelines

Under Type of publication in the table changed “Press release” to “External study communications”
Changed group review period for External study communications from 2 days to “1 week”
Chapter 11 Biospecimen Repositories and Translational Research

**Summary of Changes**

Chapter 11 was completely rewritten to reflect the current infrastructure and policies for the Alliance Biospecimen Repositories and Translational Research. Please refer to updated chapter for updated content.
Chapter 12 Investigational Agents

Summary of Changes

Section 12 Investigational Agents
The introductory paragraph was modified to read as follows:

“In Alliance studies, any agent that is provided to institutions is considered “investigational” for purposes of this policy. For investigational agents used under an IND, the IND holder is either the NCI or Alliance. Investigational agents may be provided by NCI/CTEP or directly by the industry partner. Investigational agents may be distributed to the institutions by NCI, industry, the Mayo Cancer Center Pharmacy or a third party distributor. The Alliance generally follows PMB policies (https://ctep.cancer.gov/branches/pmb/default.htm) and CTEP investigator guidelines (https://ctep.cancer.gov/investigatorresources/investigators_handbook.htm) for all IND investigational agents, irrespective of the IND holder.”

Section 12.1.1 National Cancer Institute Investigational Agents
Deleted the first paragraph “NCI distributes investigational agents for which it holds the IND, and on occasion NCI distributes investigational agents that are provided by industry (Alliance-held IND or IND exempt)”

Updated information in second paragraph regarding investigator regulatory requirements, and added text regarding the new NCI Registration and Credential Repository “Investigators must have current investigator registration documents (FDA Form 1572, Financial Disclosure Form, HSP/GCP training, biosketches, CV) on file with the NCI in order to receive investigational agents. These registrations must be renewed annually. Registration must be completed via the NCI Registration and Credential Repository (RCR). Additional information regarding registration types and required documentation is available at the Cancer Therapy Evaluation Program (CTEP) website (https://ctep.cancer.gov/investigatorresources/).”

Deleted the following text “Investigators at affiliate institutions order agents directly from the NCI, not through their main member institution. Satellite locations may order and receive agents through control locations (i.e., the location that orders and receives agents from NCI). When agents are transported between control and satellite locations, care should be taken to ensure appropriate storage conditions (e.g., cold packs).”

Added the following paragraph “NCI distributes investigational agents for which it holds the IND and may also distribute NCI investigational agents, either Alliance-held IND or IND exempt, provided by industry.”

Section 12.1.2 changed title to “Investigational Agents distributed by the Alliance” was previously “Investigational agents supplied by industry”

Modified section to the following: “Instructions for ordering agents distributed by the Alliance or third party distributors vary from study to study, and can be found in the Drug Formulation, Availability and Preparation section of the protocol. The specific form required to ship drug to an institution is described in the protocol.”
Added new Section 12.1.3 Shipment of investigational agents and renumbered subsequent sections

“Investigators have a single 1572 form on file with the PMB. Multiple pharmacy addresses may be listed on the 1572 form. By providing accurate shipping information this will assure that the FDA regulations are being followed, along with decreasing investigational agent shipping delays and expense and ensures accountability.

Investigational agent(s) will only be shipped to the designated pharmacy on the 1572 form of the investigator who is ordering the agent and must not be shipped to any other addresses after receipt at the pharmacy of record. Alternatively, investigators at affiliate institutions may order agents directly from PMB and not through their main member institution.

PMB policy allows centralized pharmacies to receive investigational agents for re-distribution to local satellite institutions and affiliated investigators who are registered with PMB and have designated a “central pharmacy” as their shipping address. If investigational agent is ordered through the main member institution, then the agent can be couriered to the satellite location if necessary. When agents are transported between control and satellite locations, care must be taken to ensure all appropriate storage conditions are maintained.

In the instance of investigators who staff more than one location, investigational agent(s) should be ordered to the central pharmacy where the patient will be receiving the investigational agent.

PMB policy also forbids secondary distribution of investigational agents to physicians who are not listed on the DTL or transfer of investigational agents between institutions or other sites. Shipment of agents directly to patients is not allowed.”

Section 12.1.4 Use of Investigational Agents

Deleted the text “if the agent is being provided by the NCI or industry” from sentence #1 in the 2nd paragraph. New paragraph reads “Commercial agents may not be substituted for an investigational agent if the agent is being provided by the NCI or industry” nor can an investigational agent be used to “pay back” or “replace” commercial supplies.”

Added the following text “The Alliance audits the pharmacy according to the NCI Guidelines for Auditing Clinical Trials (CTMB Guidelines) section 5.3.

Deleted the following text from the 3rd paragraph “A list of supplied agents for Alliance studies can be found on the Alliance website”

Deleted and updated text in the last paragraph to reflect current CTMB Audit Guidelines new paragraph reads “Compliance with investigational drug use and accountability procedures is reviewed at the time of Alliance audits and will result in the pharmacy audit being rated as critical non-compliant, non-compliant, compliant or not reviewed. A rating of critical non-compliant will automatically result in an Unacceptable audit rating for Drug Accountability and Pharmacy. Any Unacceptable rating will require a re-audit within 12 months.”

Section 12.1.5 Storage and accountability of investigational agents

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Appendix Name: Summary of Changes

Section: Appendices

Date Revised: January 1, 2018

Added sentence to first paragraph regarding DTL requirement “All study site personnel responsible for investigational agent(s) accountability must be listed on the Delegation of Tasks Log (DTL).”

Added “appropriate” to the following sentence “The appropriate NCI Drug Accountability Record Form (DARF) should be used to record the receipt and disposition of all drugs supplied (by the NCI or pharmaceutical companies) for Alliance protocols”

Removed CTEP website that linked to forms as they are no longer available on the CTEP website

Added new Section 12.1.6 Deviation from Study Protocol

“The appropriate Alliance protocol resource must be contacted if the handling or dispensing of the investigational agent(s) deviate from the protocol instructions. The deviation must be reported to the IRB of record, documented in a note-to-file, and retained in the records of the site.

Section 12.2 Investigational New Drug Applications

Deleted the following text “IND applications for Alliance-held INDs are submitted to FDA by the Alliance Chicago Office. The Alliance group chair is the Responsible Investigator on Alliance IND applications.”

Added the following text “Alliance reviews each study in development to determine if an IND/IDE application is required for a trial.”

Added the following new sections

“12.2.1 Investigational New Drug (IND) Application

12.2.1.1 IND Required

IND applications for Alliance-held INDs are submitted to the FDA by the Alliance Chicago Office. The Alliance group chair is the Responsible Investigator on all Alliance IND applications. The FDA will provide documentation of IND approval.

12.2.1.2 IND Exemption

If the FDA determines that an IND is not required, the FDA will provide documentation of IND exempt status.”

Section 12.2.2 Investigational Device Exemption (IDE)

Minor changes

Added new section 12.2.3 FDA Reporting

“For Alliance-held INDs, annual reports, correspondence, amendments, and all other reporting requirements are submitted by the Alliance Chicago Office. All adverse events (AE) whose causality may be both serious and/or unexpected (SUSAR) are reported to the FDA by the Alliance

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Chapter 14 Public Relations

Summary of Changes

Section 14.2 Public service
Added clarifying language to the 2\textsuperscript{nd} paragraph regarding the NCI Division of Cancer Prevention and Division of Cancer Control and Population Sciences.

Section 14.3 Dissemination of information to the general public

2\textsuperscript{nd} bullet - Questions of a medical nature about a specific patient
  ▪ Clarified section to note that the section is applicable to all Alliance personnel.

5\textsuperscript{th} bullet - Questions about the Alliance history, structure, and membership
  ▪ Added text to refer readers to the Alliance contact us section of the website.

New 6\textsuperscript{th} bullet added - Questions about Alliance research results
  ▪ Text added, “The Alliance works closely with the NCI, industry partners, member institutions and patient advocacy groups to disseminate information regarding the activation, progress, results and findings of its research. All requests should be referred to the Alliance communications specialist.”

Section 14.4 Confidentiality of patient information
Changed NCI to “NIH” in the 1\textsuperscript{st} paragraph
Chapter 16 Study Monitoring and Interim Analysis

Summary of Changes

Section 16 Study monitoring and interim analysis
Minor changes throughout first paragraph:
Deleted “therapeutic trials are monitored and all” from the 2nd sentence. New sentence now reads: “All treatment protocols must include a formal monitoring plan.”

3rd sentence modified (new text in bold): “All randomized phase 2 all phase 3 trials, and some specially-designated trials are formally monitored by a standing Alliance Data and Safety Monitoring Board (DSMB).”

Section 16.1.2 Function of the DSMB
Added and deleted text to #1 and #3, sections now read:

“#1 The primary responsibility of the DSMB is to review adverse event data in conjunction with interim analyses of outcome efficacy data (prepared by the study statistician) and to recommend whether the study needs to continue per protocol or be changed or terminated based on these analyses. For phase 3, phase 2/3, and randomized phase 2 trials, the committee also determines whether and to whom outcome results should be released prior to the reporting of study results at the time specified in the protocol.”

“#3 The DSMB oversees the safety and accrual data; however it is also the responsibility of the study team to review the safety and accrual information on a regular basis.”

Added additional function:

“#4 All clinical trial data release requests (e.g., baseline data for correlative studies or assay Methodology evaluation) on DSMB monitored studies have to be submitted to the DSMB for review and approval.”

#5 Deleted “prior to their implementation”

Section 16.2 Overview of DSMB procedures
Added frequency of “at least every 6 months” to the first paragraph. Section now reads:
“Each study to be monitored requires periodic (at least every 6 months) confidential reports to be prepared by the primary statistician. These reports are submitted to the DSMB, a single standing committee established for the purpose of reviewing all of the individual reports. No individuals other than members of the DSMB receive a copy; specifically, the study chair does not receive a copy.”

Section 16.2.1 Confidentiality
Modified 2nd paragraph to indicate Alliance SDC maintains confidentiality files; previous text indicated DSMB

Section 16.2.4 Recommendations
Minor edits to the first paragraph (new text is bold):

Alliance Policies and Procedures – Appendices A28
“The results of each DSMB meeting are summarized in a formal report by the Alliance Group Statistician and sent by the DSMB chair to the group chair within one week of the meeting (urgent matters are addressed immediately). The DSMB report contains recommendations on whether to modify or close each study reviewed, whether to release and report the results, and whether to continue accrual or follow-up. A primary recommendation (e.g., continue with no change; recommended or required modification; release study results and stop further DSMB monitoring) must be included in the document. The group chair must approve these recommendations before any action is taken.”

Section 16.2.4.3 Study change for non patient safety reasons

Modified the 2nd sentence (new text in bold):

“In the absence of disagreement, the Alliance group chair (working with the study chair) will be responsible for having an amendment prepared and submitted to CTEP’s Protocol and Information Office reflecting the recommendations of the DSMB and providing the rationale for the changes.”

Section 16.2.5 Study modifications

Modified paragraph (new text in bold)

Old Language: “Major modifications to the study design not motivated by confidential outcome data or patient safety/toxicity data (e.g., increasing the sample size because of more rapid than expected accrual) must be discussed with NCI/DCTD/CTEP before being presented to the DSMB for consideration. If NCI/DCTD/CTEP is willing to approve the modifications, the network group may then seek DSMB approval before submitting an official amendment to CTEP’s Protocol and Information Office.”

NEW Language “Major modifications to the study design by the study team not motivated by confidential outcome data or patient safety/toxicity data (e.g., increasing the sample size because of more rapid than expected accrual) must be discussed with NCI/DCTD/CTEP/DCP before being presented to the DSMB for consideration. If NCI/DCTD/CTEP/DCP is willing to approve the modifications, the network group informs the DSMB at the next scheduled DSMB meeting.”

Section 16.2.6 Release of results

Deleted all instances of “blinded” from section.

Modified the 2nd sentence to include additional details on when outcome data is made available. The sentence now reads “In general, outcome data from phase 3, phase 2/3, and randomized phase 2 trials would not be routinely made available to individuals outside of the DSMB until accrual has ceased and all patients have concluded their randomized treatment, and completed study follow-up and/or reached a protocol-specified endpoint.”

Section 16.2.8 Phase 2/3 trials and

Minor changes throughout section
Appendix Name: Summary of Changes
Appendix: A

Section: Appendices
Date Revised: January 1, 2018

Section 16.2.9 Industry-supported studies
Minor change in section

Section 16.3.2 Phase 2 studies
Added text to last sentence (new text in bold):
“Non-randomized phase 2 studies are routinely monitored by the study team (study chair, primary statistician, executive officer, protocol coordinator, data management personnel) and other Central
Protocol Operations Program staff (e.g. director of regulatory affairs) as applicable. Each phase 2 protocol must specify the *efficacy and adverse event* monitoring plan to be used.”
**Appendix B - Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Term</th>
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<tbody>
<tr>
<td>1572</td>
<td>Statement of Investigator (Form FDA 1572)</td>
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<tr>
<td>ABBR</td>
<td>Alliance Biorepositories and Biospecimen Resource</td>
</tr>
<tr>
<td>ACoS</td>
<td>American College of Surgeons</td>
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<tr>
<td>ACOSOG</td>
<td>American College of Surgeons Oncology Group</td>
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<td>ACS</td>
<td>American Cancer Society</td>
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<td>AER</td>
<td>Adverse Event Report</td>
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<td>AIS</td>
<td>Audit Information System</td>
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<tr>
<td>Alliance</td>
<td>Alliance for Clinical Trials in Oncology</td>
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<tr>
<td>ANFU</td>
<td>Acceptable needs follow-up</td>
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<tr>
<td>ASCII</td>
<td>American Standard Code for Information Interchange</td>
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<tr>
<td>BioMS</td>
<td>Biospecimen Management System</td>
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<td>BIQFSP</td>
<td>Biomarker, Imaging and Quality of Life Studies Funding Program</td>
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<tr>
<td>BLA</td>
<td>Biologic License Application</td>
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<td>CALGB</td>
<td>Cancer and Leukemia Group B</td>
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<tr>
<td>CAO</td>
<td>Chief administrative officer</td>
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<tr>
<td>CAP</td>
<td>College of American Pathologists</td>
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<td>CAPA</td>
<td>Corrective and Preventive Action</td>
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<td>Community Clinical Oncology Program</td>
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<td>CCP</td>
<td>Cancer Control Program</td>
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<td>CCSC</td>
<td>Core Correlative Science Committee</td>
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<td>CE</td>
<td>Continuing Education</td>
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<td>CFO</td>
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<td>CFR</td>
<td>Code of Federal Regulations</td>
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<td>Clinical Laboratory Improvement Amendments</td>
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<td>Commission on Cancer</td>
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<td>COI</td>
<td>Conflict of Interest</td>
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<td>Abbreviation</td>
<td>Full Term</td>
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<td>CPOP</td>
<td>Central Protocol Operations Program</td>
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<td>CR</td>
<td>Complete Response</td>
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<td>Cooperative Research and Development Agreements</td>
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<td>CRFs</td>
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<td>CTCAE</td>
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<td>CV</td>
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<td>DARF</td>
<td>Drug Accountability Record Form</td>
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<tr>
<td>dbGaP</td>
<td>Database of Genotypes and Phenotypes</td>
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<td>DCCPS</td>
<td>Division of Cancer Control and Population Sciences</td>
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<td>DCP</td>
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<td>DCTD</td>
<td>Division of Cancer Treatment and Diagnosis</td>
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<td>DNA</td>
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<td>DSMB</td>
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<td>DTL</td>
<td>Delegation of Tasks Log</td>
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<td>Good Clinical Practice</td>
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<td>Identification</td>
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<td>Investigational New Drug</td>
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<td>Information Technology</td>
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<td>Lung Cancer Tissue Bank</td>
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<td>Leukemia Tissue Bank</td>
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<td>MAYO</td>
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<td>MedDRA</td>
<td>Medical Dictionary for Regulatory Activities</td>
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<td>NCCCP</td>
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<td>NCDB</td>
<td>National Cancer Data Base</td>
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<td>OAOP</td>
<td>Online Agent Order Processing</td>
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<td>OEWG</td>
<td>Operational Efficiency Working Group</td>
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_Alliance Policies and Procedures – Appendices B- 3_
### Abbreviations

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<tr>
<th>Abbreviation</th>
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<tr>
<td>OHRP</td>
<td>Office for Human Research Protections</td>
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<tr>
<td>OPEN</td>
<td>Oncology Patient Enrollment Network</td>
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<td>ORI</td>
<td>Office of Research Integrity</td>
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<td>PCO</td>
<td>Pathology Coordinating Office</td>
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<tr>
<td>PDF</td>
<td>Portable Document Format</td>
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<tr>
<td>PHI</td>
<td>Protected Health Information</td>
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<td>PI</td>
<td>Principal investigator</td>
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<td>PMB</td>
<td>Pharmaceutical Management Branch</td>
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<td>PPP</td>
<td>Pharmacogenomics and Population Pharmacology</td>
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<td>Partial Response</td>
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<td>PRO</td>
<td>Patient Reported Outcomes</td>
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<td>Quality assurance</td>
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<td>Quality Assurance Review Center</td>
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<td>Quality of life</td>
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<td>RECIST</td>
<td>Response Evaluation Criteria In Solid Tumors</td>
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<td>Ribonucleic acid</td>
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<td>RRA</td>
<td>Request for Rapid Amendment</td>
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<td>Serious adverse event</td>
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<td>Statistics and Data Center</td>
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<td>SPOREs</td>
<td>Specialized Programs of Research Excellence</td>
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<td>STL</td>
<td>Washington University Bank</td>
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<td>SUSAR</td>
<td>Suspected Unexpected Serious Adverse Reaction</td>
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<td>TMA</td>
<td>Tissue Microarray</td>
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<td>TRP</td>
<td>Translational Research Program</td>
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<td>Abbreviation</td>
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<td>W-9</td>
<td>Request for Taxpayer Identification Number and Certification (Form W-9)</td>
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