

Alliance Audit Regulatory Review

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November 5,2015

Presentation Objectives

- Identify the Regulatory Review Process and the Documents that will be reviewed
 - Prior to the audit
 - Protocols approved through the utilization of a local IRB
 - Protocols approved through the utilization of the CIRB
 - During the audit
 - Protocols approved through the utilization of local IRB
 - Protocols approved through the utilization of the CIRB
- Describe Common Regulatory Deficiencies found during an audit



Regulatory Review

- Regulatory review is the review of the foundation documents for conducting a particular study at your site.
 - Two Step Process
 - Pre-review
 - At time of audit
 - Two Parts
 - IRB review
 - Informed Consent Content Review



 Per CTMB guidelines section 4.1, the list of protocols and patient cases selected will be supplied to the site at least 2 weeks (no more than 4) prior to the audit



• For each protocol selected for audit the site will forward the following regulatory documents to the Chicago Office prior to the audit date





- For each protocol selected for audit the site will forward the following regulatory documents to the Chicago Office prior to the audit date
 - Initial (Final) IRB Protocol Approval
 - Continuing / Annual Renewal Approvals
 - Required Amendment / Update Approvals
 - Selected Locally Utilized Informed Consent Form
 - Applicable Corresponding Model Consent



- For each protocol selected for audit the site will forward the following regulatory documents to the Chicago Office prior to the audit date
 - Trials reviewed under the CIRB
 - Approval letter from CIRB noting local IRB acceptance
 - Study specific worksheet with local context
 - Selected locally utilized informed consent form
 - Applicable corresponding model consent
 - All other CIRB approval documents will be reviewed at the time of audit



Step I: Regulatory Pre-Review Part I: IRB Review





Initial IRB Protocol Approval





Initial IRB Protocol Approval

July 1, 2014

Project Number:

Project Title:

What are we looking for?

 Approval date and signature by the Chair (or designee) Alliance A011106 ALTernate Approaches for Clinical Stage II or III Estrogen Receptor Positive Breast Cancer NeoAdjuvant TrEatment (ALTERNATE) in Postmenopausal Women: A Phase III Study Alliance for Clinical Trials in Oncology

Sponsor: Primary Investigator: Meeting Date: IRB Approval Date: IRB Expiration Date: Type of Approval:

5/22/2014 6/26/2014 5/21/2015 Full Committee Review

Dear Investigator:

This is to certify that your research proposal involving human subject participants has been reviewed and **approved** by the IRB. This approval is based upon the assurance that you will protect the rights and welfare of the research participants, employ approved methods of securing informed consent from these individuals, and not involve undue risk to the human subjects in light of potential benefits that can be derived from participation.

Approval of this research is contingent upon your agreement to:

- Adhere to all Policies and Procedures Relating to Human Subjects, as written in accordance with the Code of Federal Regulations (45 CFR 46).
- (2) Maintain copies of all pertinent information related to the research study including, but not limited to, video and audio tapes, instruments, copies of written informed consent agreements, and any other supportive documents in accordance with the Research Records Retention Policy.
- (3) Report potentially serious events to IRB by completing the "Adverse Event Report".

Submit deviations from previously approved research activities which were immediately necessary to eliminate apparent and immediate dangers to the subjects.

- (5) Submit Amendments to the IRB for any proposed changes from the previously approved project. Changes may not be initiated without prior IRB review and approval.
- (6) Submit an Application for Continuing Review to the regulations and policies require continuing review of research at intervals appropriate to the degree on risk, but not less than once per year.

If you have any questions regarding the human subject protection process, please do not hesitate to contact

our office.

Very truly yours,





Initial IRB Protocol Approval

July 1, 2014

Project Number:

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- Approval date and signature by the Chair (or designee)
- Full Board Review -

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IRB Coordinator



Initial IRB Protocol Approval

July 1, 2014

Project Number:

Project Title:

What are we looking for?

- Approval date and signature by the Chair (or designee)
- Full Board Review -
- Approval was received prior to patient enrollment

Alliance A011106 ALTernate Approaches for Clinical Stage II or III Estrogen Receptor Positive Breast Cancer NeoAdjuvant TrEatment (ALTERNATE) in Postmenopausal Women: A Phase III Study Alliance for Clinical Trials in Oncology

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Very truly yours,



IRB Coordinator

Part I: IRB Review Continuing / Annual Reviews





Part I: IRB Review Continuing / Annual Reviews

What are we looking for?

 Approval is <a> 365 days from last review/initial
 approval

DATE:	April 20, 2015
TO:	M.D.
FROM:	Institutional Review Board
STUDY TITLE:	[564237-3] A041202 - A Randomized Phase III Study of Bendamustine Plus Rituximab Versus Ibrutinib Plus Rituximab Versus Ibrutinib Alone in Untreated Older Patients (>65 Years of Age) With Chronic Lymphocytic Leukemia (CLL)
IRB REFERENCE #:	· · · · · · · · · · · · · · · · · · ·
SUBMISSION TYPE:	Continuing Review/Progress Report
ACTION:	APPROVED
APPROVAL DATE:	April 2, 2015
EXPIRATION DATE:	April 1, 2016
REVIEW TYPE	Full Board

- · Continuing Review IRB Submission Form
- Consent Form, Update 1, Dated 6/1/14 (Addition of Physician Name Submitting consent for renewal stamp)
- HIPAA Consent/Authorization (Only submitting for renewal stamp)
- Alliance Data and Safety Monitoring Board (DSMB) Report Dated 12/23/14

Thank you for your submission of the items as listed for the above research study. Institutional Review Board has approved the items as submitted at its

regularly scheduled meeting.

Please remember that informed consent is a process beginning with a description of the study and insurance of participant understanding followed by a signed consent form. Informed consent must continue throughout the study via a dialogue between the researcher and research participant. Federal regulations require each participant receive a copy of the signed consent document.

Please note that any revision to previously approved materials must be approved by this office prior to initiation. Please use the appropriate revision forms for this procedure.

All SERIOUS and UNEXPECTED adverse events must be reported to this office. Please use the appropriate adverse event forms for this procedure. All FDA and sponsor reporting requirements should also be followed.

Please report all NON-COMPLIANCE issues or COMPLAINTS regarding this study to this office.

Please note that all research records must be retained for a minimum of three years.

Based on the risks, this project requires Continuing Review by this office on an annual basis. Please use the appropriate renewal forms for this procedure.

If you have any questions, please contact or Please include your study title and reference number in all correspondence

with this office.

Chairperson



Part I: IRB Review Continuing / Annual Reviews

What are we looking for?

- Approval is
 365 days from last review/initial
 approval
- Full board reviewed for protocols w/ active
 recruitment or subjects on active treatment

DATE:	April 20, 2015
TO:	M.D.
FROM:	Institutional Review Board
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Chairperson







What are we looking for?

- Approvals are obtained within 90 days of the group's notification date
 - Alliance broadcasts occur on the 1st and 15th of the month
 - CTSU broadcasts occur on the 8th and 22nd of the month
- The IRB review is appropriate to the requirement (i.e. full board vs. expedited)



Protocol Update #06 03/25/2015

ALLIANCE FOR CLINICAL TRIALS IN ONCOLOGY

PROTOCOL UPDATE TO ALLIANCE A021202

PROSPECTIVE RANDOMIZED PHASE II TRIAL OF PAZOPANIB (NSC # 737754, IND 75648) VERSUS PLACEBO IN PATIENTS WITH PROGRESSIVE CARCINOID TUMORS

Pazopanib and matching placebo will be supplied by GlaxoSmithKline and distributed by CTEP

X Update:	Status Change:
Eligibility changes	Activation
Therapy / Dose Modifications / Study Calendar changes	Closure
X Informed Consent changes	Suspension / temporary closure
Scientific / Statistical Considerations changes	Reactivation
Data Submission / Forms changes	
Editorial / Administrative changes	
X Other : Updated CAEPR	

The changes included in this update to A021202 have been made in response to the NCI Action Letter from Dr. Pamela Harris dated March 24, 2015. This Action Letter is posted on the A021202 Study Page on the Alliance web site. A revised CAEPR with the new risk has been added to the protocol. Therefore, the model consent form has been revised to incorporate this new risk consistent with the new NCI Model Template Instructions.

IRB approval (or disapproval) of this update is required within 90 days. Full Board review is recommended. Please follow your IRB of record's policies.

No new patients may be consented onto this protocol until IRB approval for this amendment has been obtained. Patients consented on or before March 25, 2015 may be enrolled onto this trial before local IRB approval of the revised protocol and informed consent form has been obtained.



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Common IRB Major Deficiencies





Common IRB Major Deficiencies

•Amendment approvals obtained greater than 90 days post group's notification

•Continuing review approved by expedited review when full board review is needed

•Expired continuing reviews greater than 30 days late



Step I: Regulatory Pre-Review Part II: Informed Consent Content









- A minimum of 3 consents will be selected for review
- For each consent selected the site will forward the following to the Chicago Office prior to the audit date (including CIRB reviewed studies)
 - Current approved locally utilized informed consent form
 - Applicable model consent



ALLIANCE MODEL CONSENT FORM:

RANDOMIZED PHASE II STUDY COMPARING CABOZANTINIB (NSC #761968 and IND #116059) WITH COMMERCIALLY SUPPLIED SUNITINIB IN PATIENTS WITH PREVIOUSLY UNTREATED LOCALLY ADVANCED OR METASTATIC RENAL CELL CARCINOMA

This is a clinical trial, a type of research study. Your study doctor will explain the clinical trial to you. Clinical trials include only people who choose to take part. Please take your time to make your decision about taking part. You may discuss your decision with your friends and family. You can also discuss it with your health care team. If you have any questions, you can ask your study doctor for more explanation.

You are being asked to take part in this study because you have advanced or metastatic kidney cancer

Why is this study being done?

The purpose of this study is to find out what effects, good and/or bad, two study drugs called sunifinib and cabozantinib have on you and on advanced or metastatic kidney cancer. Sunifinib has been approved by the FDA and cabozantinib is an investigational drug. Both medications target special proteins that are on the surface of the kidney cancer cell and both drugs are taken by mouth.

How many people will take part in the study?

About 150 people will take part in this study.



Consent Form

Title of Protocol

Randomized Phase II Study Comparing Cabozantinib (NSC#761968 and IND#116059) with Commercially Supplied Sunitinib in Patients with Previously Untreated Locally Advanced or Metastatic Renal Cell Carcinoma.

Who is conducting this study?

Principal Investigator:

Sub-Investigators:

Sponsor:

Alliance for Clinical Trials in Oncology (Alliance)

Alliance for Clinical Trials in Or

Protocol No: A031203 Update #5: 01Oct2014 UPB # : Pro00004221

This is a clinical trial, a type of research study. Your study doctor will explain the clinical trial to you. Clinical trials include only people who choose to take part. Please take your time to make your decision about taking part. You may discuss your decision with your friends and family. You can also discuss it with your health care team. If you have any questions, you can ask your study doctor for more explanation.

Why have I been asked to take part in this research study?

You have been asked to take part in this study because you have advanced or metastatic kidney cancer.

Why is this study being conducted?

The purpose of this study is to find out what effects, good and/or bad, two study drugs called sunitinib and cabozantinib have on you and on advanced or metastatic kidney cancer. Sunitinib has been approved by the FDA and cabozantinib is an investigational drug. Both medications

Page

Version Date: 09/16/14

1



Version II Dec. 2002 CD, 06/03 Application packet Version 6/03 Revised 01/12 PD 1 of 18 Pt. Initials____

- Informed Consent Forms are reviewed for the 8 basic required elements of a consent (21CFR50.25)
 - Study involves research
 - Description of foreseeable risks
 - Description of benefits
 - Disclosure of alternatives
 - Description describing confidentiality maintenance
 - Compensation / treatment in the case of injury
 - Contact information for questions regarding research/rights
 - Participation is voluntary



- Informed Consent Forms are reviewed for additional elements (21CFR50.25)
 - Treatment may involve risks
 - Anticipated circumstances in which subject's participation may be terminated
 - Additional costs to the subject
 - Consequences for subject's decision to withdraw
 - Subject will be informed of significant new findings
 - Approximate number of subjects
 - A copy of this form will be given to the subject
 - http://www.ClinicalTrials.gov website listed per U.S. law



CIRB Trials

Yes the informed consent form is reviewed!

Because **CIRB** is the **IRB of record** your locally utilized consent must be a <u>word for word</u> match with the model consent with the exception of what is approved **by the CIRB** on the study specific worksheet with local context



Informed Consent Content Common Major Deficiencies





Informed Consent Content Common Major Deficiencies

- Omission of one or more risks
- Omission of one or more of the required informed consent elements
- Changes to the following without Alliance approval
 - Additions to the risks
 - Additions / Omissions to the list of alternative options
 - Changes to the translational research section (including the questions)



Step II: Regulatory Review at Time of Audit





Step II: Regulatory Review at Time of Audit

- Review of regulatory approval documents for any unannounced protocols
- Review submission of unanticipated / IND reports per your IRB policy
- CIRB reviewed trials
 - Ensure all CIRB regulatory approvals are on file at your site
 - Continuing / Annual review approvals
 - All required amendment / update approvals
 - Review of any approvals from the local IRB prior to CIRB review acceptance (if applicable)



Step II: Regulatory Review at Time of Audit

- Resolve any regulatory and consent discrepancies found during the pre-review / time of audit
- Assess any regulatory findings



Conclusion: Regulatory Review





Conclusion: Regulatory Review

- Regulatory review is a two step process
 - Pre-review (prior to the audit date)
 - Items reviewed at the time of the audit
- Regulatory review occurs in two parts
 - IRB review
 - Informed Consent Content review
- Common Major Deficiencies
 - IRB review
 - Informed Consent Content review



THANK YOU!

Ouestions from Audience Answers from Presenter

