February 1, 2017

TO: PARTICIPATING INSTITUTIONS LISTED ON THE TITLE PAGE OF THE PROTOCOL

FROM: Dawne Wenzel, M.A., Protocol Coordinator (E-mail: dwenzel@swog.org)


STATUS NOTICE

Study Chair: Dawn Hershman, M.D., M.S.
Phone number: 212/305-1945
E-mail: dlh23@columbia.edu

IRB Review Requirements
( ) Full board review required
( ) Expedited review allowed
( ) No review required

Status Change
( ) IRB Review only
( ) Activation
( ) Closure
( ) Reactivation

Protocol changes
( ) Eligibility changes
( ) Treatment / Dose Modification / Study Calendar changes
( ) Informed Consent changes
( ) Patient notification not required
( ) Patient notification required
( ) Scientific / Statistical Consideration changes
( ) Specimen Submission changes
( ) Data Submission / Forms changes
( ) Editorial / Administrative changes
( ) Other:

PERMANENT CLOSURE

The above-referenced study has met its accrual goal and will permanently close to accrual effective February 15, 2017 at 11:59 p.m. Pacific.

This memorandum serves to notify the NCI and the SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
December 15, 2016

TO: PARTICIPATING INSTITUTIONS LISTED ON THE TITLE PAGE OF THE PROTOCOL

FROM: Dawne Wenzel, M.A., Protocol Coordinator (E-mail: dwenzel@swog.org)


MEMORANDUM

Study Chair: Dawn Hershman, M.D., M.S.
Phone number: 212/305-1945
E-mail: dlh23@columbia.edu

IRB Review Requirements
( ) Full board review required
( ) Expedited review allowed
( √ ) No review required

Status Change
( ) IRB Review only
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( ) Data Submission / Forms changes
( ) Editorial / Administrative changes
( ) Other:

MEMORANDUM

The purpose of this memorandum is to inform sites of an update to the SWOG Biospecimen Bank kit ordering instructions.

The SWOG Biospecimens Bank’s Kit Management System Version 2 is being released on Monday December 19, 2016. This system will be used to order Specimen Collection Kits when the SWOG Biospecimen Bank is listed in the protocol as the kit supplier for specimen submissions.
The new Kit Management URL will be provided to all active users prior to December 19th. If you try to access Kit Management via the old URL on or after December 19th, you will automatically be redirected to the new Kit Management site.

The first time you access the site, you will be asked to register, even if you have an account in the current Kit Management system.

- Your registration does not need to be approved in order for you to access Kit Management!
- You will receive an email confirmation that you have registered for an account.
- While you may register for an account beginning on December 19th, we ask that you wait to order a kit until you actually need one, so that we can assure that anyone who needs a kit for a patient who is being screened will receive one in a timely manner.
- Remember, we provide kits for many different groups!

The ‘Help’ tab on the site includes a new user guide. We recommend that you review the user guide prior to ordering kits. In addition, requests for assistance can be sent to the Bank using the Request for Assistance entry form under the Help tab on the Kit Management website.

Online ordering with the NEW Kit Management System version offers several benefits to you:

- The new version is much quicker than the previous iteration.
- Users do not need to wait for new accounts to be approved.
- The system includes a helpful description of each kit.
- Users can edit their address, phone number and password without contacting the BPC.
- Users may have more than one shipping address connected to their account.
- Order history can be viewed online so each user can see the orders they have placed, even those not yet shipped by the SWOG Biospecimen Bank.
- The order history will also display the FedEx tracking number, so that users can track their shipments!

**Users will be able to print shipping labels via the Kit Management System instead of contacting the [Bank] to obtain labels.**

This memorandum serves to notify the NCI and the SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
TO: PARTICIPATING INSTITUTIONS LISTED ON THE TITLE PAGE OF THE PROTOCOL

FROM: Kimberly F. Kaberle, Protocol Coordinator (E-mail: kkaberle@swog.org)


REVISION #7

Study Chair: Dawn Hershman, M.D., M.S.
Phone: 212/305-1945
E-mail: dlh23@columbia.edu

IRB Review Requirements

( ) Full board review required. Reason:

( ) Initial activation (should your institution choose to participate)

( ) Increased risk to patient

( ) Complete study redesign

( ) Addition of tissue banking requirements

( ) Study closure due to new risk information

( ) Expedited review allowed

( √ ) No review required

REVISION #7

1. Title Page, Page 1: The version date of the protocol and the model consent form has been updated.

2. Participant Table, Page 2: MI047 and MI132 have been removed as these sites are participating under CRCWM.

This memorandum serves to notify the NCI and the SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
Danika Lew, M.A.
William Barlow, Ph.D.
Dona Marrah
Joseph Unger, Ph.D.
Monica Yee
March 15, 2015

TO: PARTICIPATING INSTITUTIONS LISTED ON THE TITLE PAGE OF
THE PROTOCOL

FROM: Kimberly F. Kaberle, Protocol Coordinator (E-mail: kkaberle@swog.org)

RE: S1200, “Randomized Blinded Sham- and Waitlist-Controlled Trial of
Acupuncture for Joint Symptoms Related to Aromatase Inhibitors in

MEMORANDUM

Study Chair: Dawn Hershman, M.D., M.S.
Phone: 212/305-1945
E-mail: dlh23@columbia.edu

IRB Review Requirements

(   ) Full board review required. Reason:
(   ) Initial activation (should your institution choose to participate)
(   ) Increased risk to patient
(   ) Complete study redesign
(   ) Addition of tissue banking requirements
(   ) Study closure due to new risk information
(   ) Expedited review allowed
(√) No review required

MEMORANDUM

The purpose of this memorandum is to inform sites of a change to the Master Forms Set
(found on the SWOG website at www.swog.org) for the above-noted study. The S1200
Registration Worksheet has been revised under “Stratification Question” to include:

Institution name changes
• “(2) PSOC” has been changed to “(2) FHRC”
• “(3) Grand Rapids CCOP” has been changed to “(3) West Michigan NCORP”
• “(4) Kaiser Permanente NoCa” has been changed to “(4) Kaiser NCORP”
• “(5) Good Samaritan” has been changed to “(5) Good Samaritan/Oregon HSU”
• “(8) Greenville CCOP” has been changed to “(8) Greenville NCORP”

New institution
• (11) PCRC NCORP

This memorandum serves to notify the NCI and the SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
    William Barlow, Ph.D.
    Joseph Unger, Ph.D.
    Danika Lew, M.A.
    Dona Marrah
    Monica Yee
Distribution Date: February 15, 2015  
DCP Submission Date: January 22, 2015

TO: PARTICIPATING INSTITUTIONS LISTED ON THE TITLE PAGE OF THE PROTOCOL

FROM: Kimberly F. Kaberle, Protocol Coordinator (E-mail: kkaberle@swog.org)


REVISION #6

Study Chair: Dawn Hershman, M.D., M.S.  
Phone: 212/305-1945  
E-mail: dlh23@columbia.edu

IRB Review Requirements

- ( ) Full board review required. Reason:  
  - ( ) Initial activation (should your institution choose to participate)  
  - ( ) Increased risk to patient  
  - ( ) Complete study redesign  
  - ( ) Addition of tissue banking requirements  
  - ( ) Study closure due to new risk information

- ( ) Expedited review allowed  
- ( ) No review required

REVISION #6

1. Title Page, Page 1: The version date of the protocol and the model consent form has been updated.

2. Participant Table, Page 2: The participant list has been moved from Page 1 to Page 2. Subsequent pages have been renumbered accordingly. Due to NCORP transition changes, the following institutions have been added to the list:

   a. CA156/Kaiser, Walnut Creek  
   b. CA172/Kaiser, Santa Clara  
   c. CA175/Kaiser, San Jose
The following institutions have been revised:

a. CA154/Kaiser Permanente Northern California has been changed to CA154/Kaiser NCORP.
b. MI021/Grand Rapids CCOP has been changed to CRCWM/West Michigan NCORP.
c. NY024/Columbia University MBCCOP has been changed to NY024/Columbia University NCORP.
d. SC036/Greenville CCOP has been changed to SC036/Greenville Health System Cancer Institute-Eastside.
e. WA011/Puget Sound has been changed to WA007/Swedish Medical Center.

3. Table of Contents, Pages 3-4: The Table of Contents has been updated with revised page numbers.

4. Section 5.1a, Page 20: The following sentence has been removed: “Patients must have undergone breast cancer surgery.” The following text has been added to the last sentence: “If patient has undergone breast cancer surgery, she...”. This change has been made because not all eligible patients undergo breast cancer surgery. We have removed the limitation that they must have undergone surgery and kept in place the criteria that they must have made a complete surgical recovery. Having breast cancer surgery has no impact on the use of aromatase inhibitors in women with early stage breast cancer nor on the primary endpoint of the study, which is to determine whether true acupuncture and waitlist control causes a significant reduction in joint pain related to aromatase inhibitors (AIs) in women with early stage breast cancer as measured by the Brief Pain Inventory-Short Form (BPI-SF) worst pain score at 6 weeks.

5. Section 7.6a, Page 26: “Breaks of up to 30 consecutive days are allowed.” has been removed from the last sentence of the last paragraph of this section for clarification. The acupuncture sessions and follow-up assessments should be collected as close to schedule as possible. In Section 7.6a, flexibility of the acupuncture schedule is described (“The schedule will be flexible to allow for 8-12 sessions during the first 6 weeks, which may be more reflective of what may happen in the “real world” setting.”) and Section 7.7 specifies the wish for follow-up under intent to treat.

6. Section 9.0, Page 30: In the “#” footnote, the following sentence was added for clarification: “The Week 6 and Week 12 study visits are preferably performed after acupuncture Session 12 and Session 18, respectively.”
Institutions **should** update their local consent forms to include the change to the Model Consent Form. SWOG considers that the Model Consent Form change **does not** represent an alteration in risk/benefit ratio. Therefore, local accrual does not need to be suspended pending implementation of this change. Patients need not be informed of the following change unless required by the local IRB.

7. Model Consent Form, Page 8: Under “What are the costs…”, the URL in the fourth paragraph has been updated.

This memorandum serves to notify the NCI and the SWOG Statistical Center.

**cc:** DCP PROTOCOL & INFORMATION OFFICE
William Barlow, Ph.D.
Joseph Unger, Ph.D.
Danika Lew, M.A.
Dona Marrah
Monica Yee
Distribution Date: May 15, 2014
DCP Submission Date: April 28, 2014

TO: Columbia University MBCCOP, Good Samaritan Hospital, Grand Rapids CCOP, Greenville CCOP, Kaiser Permanente Northern California, Lahey Hospital and Medical Center, Puget Sound, St. Luke's Mountain States, University of Southern California, University of Utah

FROM: Kimberly F. Kaberle, Protocol Coordinator (E-mail: kkaberle@swog.org)


REVISION #5

Study Chair: Dawn Hershman, M.D., M.S.
Phone: 212/305-1945
E-mail: dlh23@columbia.edu

IRB Review Requirements

( ) Full board review required. Reason:
( ) Initial activation (should your institution choose to participate)
( ) Increased risk to patient
( ) Complete study redesign
( ) Addition of tissue banking requirements
( ) Study closure due to new risk information
(√) Expedited review allowed
( ) No review required

REVISON #5

1. The Version Date of the protocol and Model Consent Form have been updated. There have been no additional changes to the Model Consent Form.

2. Title Page, Page 1: The NCT number has been added below the title. Under Biostatisticians, Joseph Unger’s degree has been changed from "M.S." to "Ph.D.". The participants list has been reformatted into the new NCI approved table format and moved to the bottom of the page.

3. Table of Contents, Pages 2-3: The Table of Contents has been updated.
4. Section 5.1a, Page 19: The second sentence has been revised from: “Patients must have undergone modified radical mastectomy or breast sparing surgery.” to “Patients must have undergone breast cancer surgery.” in order to allow more eligible patients to participate.

5. Section 5.3k, Page 20: NSAIDs have been added to the list of allowed analgesics. It had been erroneously removed in a previous revision.

6. Section 5.3o, Page 20: Prestudy history and physical within 180 days prior to registration has been added. This addition has caused information from Page 20 to be displaced to Page 21.

7. Section 7.5, Page 24: The third sentence of the second paragraph has been added to allow for seats in the range of 40-50 cm. This was added in order to accommodate sites that did not have seats exactly 42 cm high.

8. Section 7.6, Page 25: The following italicized text was added to the first sentence of the paragraph below the bulleted list in order to allow greater flexibility in scheduling and is a reflection of the real world setting, “...it may be made up in the following weeks for up to 3 sessions...”.

9. Section 8.5, Page 27: The reference to “AdEERS” has been replaced with “CTEP-AERS”. An occurrence of “Study Coordinator” has been updated to “Study Chair”.

10. Section 9.0, Page 28: The S1200 Assessment of Blinding form was added in the Questionnaires section as it was inadvertently missing in previous versions.

11. Section 13.3b, Page 34: Information regarding the affirmation of eligibility was added to the first sentence of the first bullet.

12. Section 15.1c.4, Page 39: The negative sign (-) was removed before -4°C as the correct temperature is 4°C.

13. Section 16.0, Page 42: The SWOG standard confidentiality statement has been added below the “Monitoring” section. This addition caused information from Page 42 to be displaced to Page 43.

14. Section 16.1b, Page 42: The references to the “NCI’s Adverse Event Expedited Reporting System (AdEERS)” have been changed to the “CTEP Adverse Event Reporting System (CTEP-AERS)” throughout this section. The last sentence regarding paper AdEERS forms has been deleted.

15. Section 16.1e, Page 43: The references to “AdEERS” have been changed to “CTEP-AERS” throughout this section.

Please attach this memorandum to your copy of the protocol. Replacement pages are attached for the revised pages referenced above. This memorandum serves to notify the NCI and SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
   William Barlow, Ph.D.
   Joseph Unger, Ph.D.
   Danika Lew, M.A.
   Dona Marrah
   Jo Ann Hartline, M.P.H., M.S.W.
TO: Columbia University MBCCOP, Good Samaritan Hospital, Grand Rapids CCOP, Greenville CCOP, Kaiser Permanente Northern California, Lahey Hospital and Medical Center, Puget Sound, St. Luke's Mountain States, University of Southern California, University of Utah

FROM: Kimberly F. Kaberle, Protocol Coordinator


REVISION #4

Study Chair: Dawn Hershman, M.D., M.S.
Phone: 212/305-1945
E-mail: dlh23@columbia.edu

IRB Review Requirements

( ) Full board review required. Reason:
( ) Initial activation (should your institution choose to participate)
( ) Increased risk to patient
( ) Complete study redesign
( ) Addition of tissue banking requirements
( ) Study closure due to new risk information

(√) Expedited review allowed
( ) No review required

In addition to the protocol revisions below, the form packet, located on the protocol abstract page on the SWOG website (www.swog.org), has been revised with the following changes:

- The following patient questionnaires have been added in Spanish:
  - S1200 Brief Pain Inventory Short Form (BPI-SF) – S1200 Cuestionario Breve Para la Evaluación Del Dolor (Edición Corta) (Form #33424)
  - S1200 Western Ontario and McMaster Universities Osteoarthritis (WOMAC) Index (Version 3.1) – S1200 WOMAC VA3.1 Cuestionario (Form #51571)
• **S1200** Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands (M-SACRAH) – **S1200** Puntuación-Modificada Para la Evaluación Y Cuantificación de Afecciones Reumatoideas Crónicas de Las Manos (Form #60281)

• **S1200** PROMIS Pain Impact-Short Form (PROMIS PI-SF) – **S1200** PROMIS Efectos Del Dolor – Cuestionario Abreviado 6B (Form #33617)

• **S1200** FACT-ES Trial Outcome Index (Version 4) – **S1200** FACT-ES (4ª Versión) (Form #34481)

• **S1200** Aromatase Inhibitor Usage Form – **S1200** Cuestionario de Uso de Los Inhibidores de la Aromatasa (Form #19842)

• **S1200** Assessment of Blinding – **S1200** Evaluación de Cegamiento (Form #449)

**S1200** Registration Form – Lahey Hospital and University of Utah have been added to Page 1 as stratification factors. The form number has been updated from #2385 to #38433.

1. Title Page: The version date, table of contents, and format of the title page have been updated. Columbia University has been changed to Columbia University MBCCOP. Lahey Hospital and Medical Center and University of Utah have been added to the Participants List in order to increase accrual.

2. Schema, Page 4: In the first paragraph, worst pain score of at least “5”, has been changed to “3” to be consistent with revised eligibility criteria in Section 5.3a.

3. Section 4.0, Page 18: Stage 0 Stage grouping has been added as it includes DCIS which is being added as an inclusion eligibility criteria.

4. Section 5.1a, Page 19: Patients with histologically confirmed DCIS are now eligible for the study. This eligibility criterion has been expanded to include these patients because they may also be experiencing joint pain that started or increased since starting an aromatase inhibitor.

5. Sections 5.2-5.5, Pages 19-21: The subsections under each of these secondary heading sections have been renumbered to begin at “a” to follow standard SWOG formatting.

6. Section 5.3a, Page 19: The worst pain score has been changed from “at least 5” to “at least 3”. This has been changed in order to broaden eligibility. The study’s statistics were based on a smaller randomized-controlled trial of true vs. sham acupuncture completed at Columbia University, which used a worst pain score on the BPI item #2 of at least 3 out of 10. This study found a significant difference in mean BPI-SF worst pain scores at 6 weeks, which was lower for the true acupuncture group compared with the sham acupuncture group (3.0 v 5.5; P < .001). The 6 week time point is also the primary endpoint for this current study. (http://jco.ascopubs.org/content/28/7/1154.short)

7. Section 5.3j, Page 20: The following italicized text was added for clarification, “…or intra-articular steroids for joint symptoms within 28 days…”.

8. Section 5.3k, Page 20: The following italicized text was added for clarification as combination NSAIDs are commonly taken by this population and are not opioid analgesics, “…with the exception of combination NSAIDs and acetaminophen)…”
9. Section 5.3m, Page 20: “or Spanish” was added to the end of this eligibility criterion to allow Spanish-speaking patients to participate as translated questionnaires are now available for their use. These questionnaires were added to increase study accrual since two of the study sites (Columbia University MBCCOP and University of Southern California) see a large Spanish-speaking population.

10. Section 7.3a, Page 22: “by review of the web-based study calendar” and the second sentence, “The web-based calendar will be maintained by Columbia University.” have been removed from the first paragraph as sites did not find this additional web-based calendar helpful to document the information required on the S1200 Adherence to Acupuncture Form.

11. Section 7.3b, Page 23: The following sentence has been added as the last sentence in the fifth paragraph in order for the study team to see durability of the free sessions until the patient’s study completion, “Use of the 10 free true acupuncture sessions will be reported using the Columbia University CRF until the patient reaches their 52 week visit.”

12. Section 7.5, Page 24: In the second sentence of the first paragraph, “Detecto DHS 88 Digital Grip Strength Dynamometer in kilograms (kg)” has been changed to “FEI Baseline Digital Smedley Spring Dynamometer (200 pound)” as the original dynamometer is no longer being made by the company.

13. Section 7.6a, Page 25: In the last bullet of this section, “for up to three months after receipt of the voucher” has been removed and replaced with “until the patient reaches their 52 week visit.” This was changed to see the durability of the free sessions until the patient’s study completion.

14. Section 9.0, Pages 28-29: The formatting of the weeks has been corrected. Week “25” has been corrected to “52” and a “^” footnote has been added to this week to allow a +/- 14 day window for patient assessment. The “∑” footnote has been added to allow a +/- 7 day window for patient assessment for Week 2. The “≦” footnote has been added to allow 180 days prior to registration for patient’s history, height, weight, and performance status. Collection of height, weight, and performance status has been removed at the Week 6, 12, 24, and 52 timepoints as this information does not add necessary data to the study and was never electronically captured.

15. Section 11.1, Page 31: In the third sentence of the first paragraph, worst pain score of “5” has been changed to “3” to be consistent with revised eligibility criterion in Section 5.3a.

Please attach this memorandum to your copy of the protocol. Replacement pages are attached for the revised pages referenced above.

This memorandum serves to notify the NCI and SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
William Barlow, Ph.D.
Joseph Unger, M.S.
Danika Lew, M.A.
Dona Marrah
Jo Ann Hartline, M.P.H., M.S.W.
Distribution Date: April 1, 2013
DCP Submission Date: March 7, 2013

TO: Columbia University, Good Samaritan Hospital, Grand Rapids CCOP, Greenville CCOP, Kaiser Permanente Northern California, Puget Sound, St. Luke's Mountain States, University of Southern California

FROM: Kimberly F. Kaberle, Protocol Coordinator


REVISION #3

Study Coordinator: Dawn Hershman, M.D., M.S.
Phone: 212/305-1945
E-mail: dlh23@columbia.edu

IRB Review Requirements

( ) Full board review required. Reason:
( ) Initial activation (should your institution choose to participate)
( ) Increased risk to patient
( ) Complete study redesign
( ) Addition of tissue banking requirements
( ) Study closure due to new risk information

( √ ) Expedited review allowed
( ) No review required

REVISION #3

In addition to the protocol revisions below, the form packet, located on the protocol abstract page on the SWOG website (www.swog.org), has been revised with the following changes:

- **S1200** Registration Worksheet (Form #2384): On Page 1 under Stratification Question, Greenville CCOP has been added. The form # has been changed from #52546 to #2384.

- **S1200** Functional Testing (Form #3766): “If these readings are not within < 10% of each other, then do further contractions, with 1 minute between each, until they are repeatable.” has been removed from the note section.

- **S1200** Aromatase Inhibitor Usage Form (Form #22213): On Page 5, the Medication Adherence Questionnaire has been replaced with the Morisky Medication-Taking Compliance and the references have been updated. The form # has been changed from #39879 to #2213.
1. **Title Page**: The version date and Table of Contents have been updated. Greenville CCOP was added to the participant’s list.

2. **Section 2.0, Page 11**: A paragraph to address safety concerns regarding acupuncture in breast cancer patients has been added prior to the “Mechanism of Action of Acupuncture” section.

3. **Section 5.2d, Page 19**: The minimum required time a patient must be on an aromatase inhibitor has been changed from 90 days to 30 days. Patients must be on a stable dose for at least 30 days.

4. **Section 5.3g, Page 20**: This eligibility criterion has been updated from “must not have had prior acupuncture treatment within the past 12 months or for AI-induced joint symptom at any time” to “must have had two or fewer acupuncture treatments within the past 12 months for any reason except for joint symptoms. Patients must not have had prior acupuncture treatment for joint symptoms at any time.”

5. **Section 5.3i, Page 20**: This eligibility criterion has been added as latex is a component of the ear seeds adhesive. Subsequent sections have been renumbered accordingly.

6. **Section 7.5, Page 24**: The following has been removed from the first paragraph: “If these readings are not within < 10% of each other, further contractions will be undertaken, with 1 minute between each, until they are repeatable. Once repeatable...”.

7. **Section 8.2, Page 26**: The fifth and sixth sentences regarding guidelines for the use of acupuncture in patients with lymphoedema have been added.

8. **Section 9.0, Pages 28 and 29**: “Physical Exam” has been removed from under the “Physical” section, as only height, weight and PS are required pre-study. The location of the forms has been updated in the note below the calendar. The # footnote has changed from +/- 3 days to +/- 10 days. The ^ footnote has been changed from +/- 7 days to +/- 14 days. The ¥ footnote has been added to clarify who can perform the physical examinations.

9. **Section 14.2, Page 35**: This section has been updated as forms are now located on the S1200 abstract page on the SWOG website.

10. **Section 15.1c.4, Page 39**: This section was added to include shipping instructions for specimens collected on Friday.

11. **Section 17.0, Page 55**: References 160 and 161 have been added.

12. **Appendix 18.1, Pages 58 and 60**: In Tables 1 and 2 the heading for the “Auricular acupuncture points” has been updated to state “given in right ear on odd numbered visits and left ear on even numbered visits”. On Table 2 on Page 60, the “shoulder” point has been updated from “…above the elbow, 3 cun above the olecranon, 0.5 cun toward the anterior of the small intestine meridian” to “…5 cun below the anterior axillary fold, and 1 cun anterior to the lung channel.”
The protocol has been reformatted and repaginated to meet the current requirements for electronic protocol submission. This includes addition of second level headings in instances where they were previously absent, reformatting the title page to include all second level headings, reformatting the protocol calendar into M.S. Word, removal of form numbers and removal of the consent form as Section 18.0. References and page numbers affected by these changes have been updated throughout the protocol.

Please attach this memorandum to your copy of the protocol. Replacement pages are attached for the revised pages referenced above.

This memorandum serves to notify the NCI and SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
   William Barlow, Ph.D.
   Joseph Unger, M.S.
   Danika Lew, M.A.
   Dona Marrah
   Jo Ann Hartline, M.P.H., M.S.W.
January 1, 2013

TO: ALL SWOG MEMBER, CCOP AND AFFILIATE MEDICAL INVESTIGATORS AND CLINICAL RESEARCH ASSOCIATES

FROM: SWOG Operations Office

RE: Eligibility Affirmation

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MEMORANDUM

By signing the FDA 1572, every SWOG investigator has agreed to conduct studies in compliance with the protocol, and to personally conduct or supervise the investigation. A critical step in this process is verification of patient eligibility.

Effective January 1st, 2013, every registering investigator or another SWOG investigator designate is required to sign a statement on the Registration Worksheet that the eligibility criteria have been confirmed. This worksheet will not be submitted to Data Operations Office but must be maintained at the local institution for review during audits.

As part of this transition, forms and the forms list (Section 18.2) are being removed from active studies and will be posted separately on the individual protocol abstract page for each study. Subsequent pages have been renumbered accordingly. No other form, protocol, or consent form changes have been made as part of the transition.

If you have any questions, please contact the SWOG Operations Office at 210/614-8808.
REVISION #2

1. Title Page: The version date has been updated. The participants list has been updated to include Good Samaritan Hospital (of Oregon Health and Sciences University), Puget Sound, and St. Luke’s Mountain States. Fred Hutchinson Cancer Center and Oregon Health and Sciences University have been removed.

2. Section 18.2a, Page 50: The form number has been changed from "39748" to "53962".

3. S1200 Registration Worksheet (Form #53962): Under "Stratification Question", the study sites have been updated to be consistent with the participants list on the title page.

This memorandum serves to notify the NCI and SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
    Danika Lew, M.A.
    William Barlow, Ph.D.
    Joseph Unger, M.S.
    Jeri Jardine
REVISION #1

1. Title Page: The version date has been updated.

2. Section 7.3a, Page 19: For clarification, the first sentence has been updated to: Acupuncture adherence will be ascertained by the CRA by review of the web-based calendar and documented on the S1200 Adherence to Acupuncture Form (Form #45337). The second sentence has been added to clarify that Columbia University will maintain the web-based calendar.

3. Section 7.6a, Page 22: The following has been added to the end of the last sentence of the second bullet, “for up to three months after receipt of the voucher.”

4. Section 14.7j, Page 32: “[WEEK 6 AND 12]” has been added.
5. Section 15.1c.3, Page 33: The urine collection tube has been changed from 20 mL to 15 mL.

6. Section 15.2c.4, Page 35: In the third sentence, "Tools of the Trade" has been changed to "Training" and "Program" has been changed to "Module".

7. Section 15.3b, Page 36: The e-mail address for Ramona Jayasena has been corrected to rj2002@columbia.edu.

8. **S1200** Prestudy Form (Form #16333): Under Patient and Disease Description, information regarding lymph nodes has been added. Under Prior Treatment Related to This Cancer, the Date Started, Date Stopped and Currently receiving fields for each Aromatase Inhibitor have been removed. The form number has been updated from 41314 to 16333.

9. **S1200** Functional Testing (Form #3766): The measurement of grip strength has been changed from kPa to kg. The form number has been updated from 37709 to 3766.

10. **S1200** Adherence to Acupuncture (Form #45337): Data collection for Weeks 1 and 3 has been changed to 2 and 4. The form number has been updated from 6044 to 45337.

11. **S1200** Adverse Event Summary Form (Form #37441): Data collection for Weeks 1 and 3 has been changed to 2 and 4 and collection during Weeks 24 and 52 have been removed. The form number has been updated from 5183 to 37441.

12. **S1200** Supplemental Agents Reporting Form (Form #52616): Data collection for Weeks 1 and 3 has been changed to 2 and 4. The following question has been added to Page 2: “Has the patient used therapeutic massage since the last assessment?” The form number has been updated from 9413 to 52616.

13. Section 19.1, Page 96: In the second sentence of the second paragraph, the 3 inch needles have been updated from 34-gauge to 30-gauge. In the ninth sentence, “up to 2 licensed acupuncturists” has been changed to “at least 1 other licensed acupuncturist”.

14. Section 19.1, Page 99: The following has been added after the first sentence: The auricular sham product used will be Sakamura Migrain Ear Pellet, silver, with pellets removed (manufactured by Sakamura Lab & Co., Kyoto, Japan).

15. Section 19.4, Page 102: In the first sentence “a web-based training for” was added. In the first sentence of the second paragraph “acupuncture tables” has been deleted.

16. New form numbers have been added in the following Sections: 5.0 (Page 17), 7.2 (Page 19), 7.3a (Pages 19 and 20), 7.3b (Page 20), 7.4 (Page 21), 14.5 and 14.6 (Page 31), 14.7 (Page 32), 14.9 and 14.10 (Page 32), and 18.2 (Page 50).

This memorandum serves to notify the NCI and SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
    William Barlow, Ph.D.
    Joseph Unger, M.S.
    Danika Lew, M.A.
    Jeri Jardine
Distribution Date: April 1, 2012
E-mailed Date: March 27, 2012

TO: Columbia University, Fred Hutchinson Cancer Research Center, Grand Rapids CCOP, Kaiser Permanente Northern California, Oregon Health and Sciences University, University of Southern California

FROM: Kimberly F. Kaberle, Protocol Coordinator


STATUS NOTICE

Study Coordinator: Dawn Hershman, M.D., M.S.
Phone: 212/305-1945
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IRB Review Requirements

(√) Full board review required. Reason:
   ( √) Initial activation (should your institution choose to participate)
   ( ) Increased risk to patient
   ( ) Complete study redesign
   ( ) Addition of tissue banking requirements
   ( ) Study closure due to new risk information

( ) Expedited review allowed
( ) No review required

ACTIVATION

The study will be activated as of March 27, 2012 at 2:00 P.M. EDT. However, please note that (in additional to requiring IRB approval from your own site) patients cannot be registered to the study until at least one acupuncturist from your site has undergone the acupuncture training that will take place at the Spring 2012 SWOG Group Meeting. The entire protocol is attached for your use.

This memorandum serves to notify the NCI and SWOG Statistical Center.

cc: DCP PROTOCOL & INFORMATION OFFICE
William Barlow, Ph.D.
Joseph Unger, M.S.
Danika Lew, M.A.
Jeri Jardine
SWOG

RANDOMIZED BLINDED SHAM- AND WAITLIST-CONTROLLED TRIAL OF ACUPUNCTURE
FOR JOINT SYMPTOMS RELATED TO AROMATASE INHIBITORS IN WOMEN WITH
EARLY STAGE BREAST CANCER

NCT #01535066

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National Cancer Institute, Office of Research on Women's Health

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OR013/Good Samaritan Hospital
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SC053/Greenville Health System Cancer Institute-Butternut
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WA008/Fred Hutchinson Cancer Research Center
WA007/Swedish Medical Center
WA102/Seattle Cancer Care Alliance
WA179/Salmon Creek Hospital
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SCHEMA

Women with stage I-III hormone receptor-positive breast cancer who are receiving adjuvant aromatase inhibitors (AIs) and report worst pain of at least 3 (out of 10) that has started or increased since initiation of aromatase inhibitor treatment

At baseline:
- Questionnaires
- Functional testing
- Blood/urine collection

RANDOMIZE
2:1:1

Intervention*:
- True Acupuncture twice weekly x 6 weeks (12 sessions) then weekly x 6 weeks (6 sessions)
  vs.
- Sham Acupuncture twice weekly x 6 weeks (12 sessions) then weekly x 6 weeks (6 sessions)
  vs.
- Waitlist control x 12 weeks

At 2 and 4 weeks:
- Telephone contact

At 6, 12, and 24 weeks:
- Questionnaires
- Functional testing
- Blood/urine collection

At 16 and 20 weeks:
- Telephone contact and BPI-SF Questionnaire

At 52 weeks:
- Questionnaires
- Functional testing
- Urine collection

*At 24 weeks, all 3 arms will receive vouchers for 10 true acupuncture sessions.
1.0 OBJECTIVES

1.1 Primary Objective

a. To determine whether true acupuncture administered twice weekly for 6 weeks (8-12 sessions) compared to sham acupuncture and waitlist control causes a significant reduction in joint pain related to aromatase inhibitors (AIs) in women with early stage breast cancer as measured by the Brief Pain Inventory-Short Form (BPI-SF) worst pain score at 6 weeks.

1.2 Secondary Objectives

Secondary objectives are to investigate the effects of true acupuncture administered twice weekly for 6 weeks (8-12 sessions) followed by 6 weekly treatments (4-6 sessions) of maintenance (12-18 sessions total over 12 weeks) compared to sham acupuncture and waitlist control in this study population. The evaluations at 12 and 24 weeks are to determine the benefit of additional 6 weekly acupuncture treatments for maintenance and to determine the durability of response after stopping acupuncture, respectively. The evaluation at 52 weeks is to determine the long-term effects of acupuncture and adherence to AIs.

b. To evaluate the effects of acupuncture on the Brief Pain Inventory-Short Form (BPI-SF) worst pain, worst stiffness, pain severity, and pain-related interference scores at 6, 12, 16, 20, 24, and 52 weeks.

c. To evaluate the effects of acupuncture on Western Ontario and McMaster Universities Osteoarthritis (WOMAC) index (pain, stiffness, and function) for the hips and knees at 6, 12, 24, and 52 weeks.

d. To evaluate the effects of acupuncture on Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands (M-SACRAH) (pain, stiffness, and function) at 6, 12, 24, and 52 weeks.

e. To evaluate the effects of acupuncture on the PROMIS Pain Impact-Short Form (PROMIS PI-SF) at 6, 12, 24, and 52 weeks.

f. To evaluate the effects of acupuncture on quality of life (QOL) as assessed by the Functional Assessment of Cancer Therapy-Endocrine Subscales (FACT-ES) at 6, 12, 24, and 52 weeks.

g. To evaluate the effects of acupuncture on functional testing with grip strength and “Timed Get Up and Go” (TGUG) test at 6, 12, 24, and 52 weeks.

h. To evaluate the effects of acupuncture on analgesic and opioid use at 2, 4, 6, 12, 16, 20, 24, and 52 weeks.

i. To evaluate the effects of acupuncture on self-reported AI adherence at 12, 24, and 52 weeks.

j. To assess AI adherence via urine AI metabolites at baseline, 24, and 52 weeks.

k. To evaluate the effects of acupuncture on serum hormones (estradiol, FSH, LH) and inflammatory biomarkers (serum TNFα, IL-6, IL-12, CRP and urine CTX-II) at 6, 12, and 24 weeks.

l. To evaluate whether polymorphisms in CYP19A1 aromatase gene predict severity of AI-related joint symptoms.

m. To assess the safety and tolerability of acupuncture in this study population.
Use of Aromatase Inhibitors for Breast Cancer

Endocrine treatments, including tamoxifen and aromatase inhibitors, are widely prescribed for all stages of hormone-responsive breast cancer. These agents are primarily directed at inducing estrogen deprivation through blocking estrogen at the receptor level (tamoxifen), or by inhibiting estrogen biosynthesis (aromatase inhibitors). Tamoxifen, a selective estrogen receptor modulator (SERM), is both an antagonist and a partial agonist of the estrogen receptor. (1) Tamoxifen may cause hot flashes, vaginal bleeding, and serious long-term side effects, including endometrial cancer and thromboembolism. (2-4)

Third-generation aromatase inhibitors (AIs), namely anastrozole, letrozole and exemestane, markedly suppress plasma estrogen levels in postmenopausal women by inhibiting the enzyme responsible for the conversion of androgens to estrogens in peripheral tissues (skin, muscle, fat, benign and malignant breast tissue). (5-7) Treatment of hormone receptor-positive breast cancer in postmenopausal women with third-generation AIs has been shown to be superior to tamoxifen with respect to disease-free survival, distant and local recurrence rates, and incidence of contralateral breast cancer. (8-13) The Arimidex® (anastrozole), Tamoxifen, Alone or in Combination (ATAC) trial compared five years of anastrozole to 5 years of tamoxifen in 9,366 postmenopausal women with localized breast cancer. After a median follow-up of 68 months, anastrozole significantly prolonged disease-free survival (HR 0.87, 95% CI 0.78-0.97), time-to-recurrence (HR 0.79, 95% CI 0.70-0.90), and contralateral breast cancers (42% reduction, 95% CI 12-62%). (10) As a result of these studies, they are currently the first-line hormonal therapy in postmenopausal women with early stage breast cancer and standard of care for women who have received 2½ years to 5 years of tamoxifen in the adjuvant settings. (11-13) They are also undergoing evaluation as chemopreventive agents in several large clinical trials for the primary prevention of breast cancer.

Side Effects Associated with Aromatase Inhibitors

Despite the well-proven efficacy of AIs for the treatment of hormone-sensitive breast cancer, some patients suffer from side effects or even stop treatment early due to undesirable toxicities. The most common side effects of AIs are hot flashes, vaginal dryness, musculoskeletal pain and headache, and possibly alterations in serum lipid profiles. (8) In addition, all third-generation AIs increase bone resorption and may predispose to osteoporosis and fractures. (14, 15) In one study, 16% of metastatic breast cancer patients complained of joint pain within 2 months of starting anastrozole and 5% had to discontinue therapy because of severe arthralgia. (16) Discontinuation of anastrozole resulted in resolution of pain-related symptoms. In large adjuvant trials involving AIs, the incidence of musculoskeletal disorders was 20% to 30% and nearly 5% of patients discontinued therapy in the AI group because of toxic effects. (8, 9) Investigators have recently shown that at the end of 4.5 years of therapy, 30% of women have discontinued hormonal treatment, and of those who continue, only 60% are taking the medication 80% of the time or more. (17)

A survey was conducted to assess the prevalence of joint symptoms in postmenopausal women on adjuvant AIs for early stage breast cancer. (18) In the clinical breast oncology academic practice at Columbia University Medical Center (CUMC), 253 consecutive breast cancer patients receiving adjuvant AIs were screened over a 4-month period. Of these women, 200 completed a 25-item self-administered survey asking about the presence of joint pain or stiffness in the preceding week, if symptoms started or worsened after initiating AIs, and the location and severity of symptoms using a 0-10 scale. Among all participants, 50% reported joint pain and 47% joint stiffness within the past week which started or worsened after initiating AIs. Among those with AI-related arthralgias, over 50% reported joint symptoms in the hands and/or knees. The score for severity of joint symptoms on a scale of 0-10 was ≥ 5 in 68% of the patients who -
experienced AI related arthralgia. Subsequently, 53 patients who were initiating AI therapy were prospectively evaluated. Compared to baseline, there was a 49% increase in joint pain severity at 6 months and an 86% increase in pain-related interference at 6 months as assessed by the Brief Pain Inventory-Short Form (BPI-SF). At 3 months, 46% of subjects had at least a 2-point increase in the BPI-SF worst pain score (0-10 scale) from baseline, and at 6 months, a significant decrease in pinch grip strength was detected (p=0.05). The results are consistent with findings from other studies. (16, 20)

In a study evaluating menopausal symptoms in breast cancer patients receiving endocrine therapy, there was a significant change in musculoskeletal pain in women receiving AIs. (21) Fifty percent of women who were asymptomatic at baseline reported variable degrees of pain after one month of treatment. There was a marked increase in the number of patients reporting severe to intolerable symptoms after three months of therapy, which led to treatment interruption in 11% of patients. Patients usually present with polyarthralgia affecting the hands, knees, hips, lower back or shoulders, which is often refractory to conventional pharmacological interventions. (16) The musculoskeletal pain appears to be specific for this class of compounds, regardless of the AI prescribed.

Recent studies have found that AI-induced arthralgia is accompanied by physical changes in the affected joints seen on MRI, EMG, and ultrasound. Women on AIs for six months had decreased grip strength and increased tenosynovial changes seen on MRI. (22) In a study conducted by Dizdar et al., women taking AIs have increased tendon thickness and higher rates of effusions in hand joints/tendons on musculoskeletal sonography compared to women who never received AIs. (23) AI use is also associated with a greater incidence of carpal tunnel syndrome of moderate intensity and short duration. (24)

The risk factors for developing AI-associated arthralgia are unclear. In the ATAC trial, high body mass index (BMI), prior chemotherapy, history of hormone replacement therapy were major risk factors for developing joint symptoms. (25) In a cross-sectional survey conducted at CUMC, prior taxane chemotherapy was associated with a 4-fold increased risk of AI-related joint symptoms, and BMI of 25-30 kg/m² and prior tamoxifen use were inversely associated with AI-associated arthralgias. (18) A recent study found a significant association between greater joint symptoms and lower bone mineral density in women taking AIs. (26) Recent studies have reported an association between a polymorphism in the 3' untranslated region of the CYP19A1 aromatase gene and improved AI treatment efficacy. Less is known about its relationship to toxicity. (27, 26)

AIs are better tolerated than tamoxifen with respect to hot flushes, vaginal bleeding and discharge, endometrial cancer and thromboembolic events. (8) However, patients receiving AIs have a higher incidence of osteoporosis, fractures and musculoskeletal symptoms (arthralgia and other joint disorders) compared to tamoxifen. (8, 9, 13) Although joint pain is not considered a serious side effect, increasing intensity decreases patient compliance and adversely affects quality of life. The risk of important long-term toxicities, including arthritis and osteoporosis, may increase with the use of AIs and the increase in life expectancy among breast cancer survivors. While bisphosphonates such as alendronate are available for the prevention of bone loss, safe and effective treatments are needed to alleviate AI-induced musculoskeletal pain.

**Mechanism of Aromatase Inhibitor-Induced Musculoskeletal Pain**

Estrogen deficiency after menopause has been linked to an increase in several chronic inflammatory conditions, including osteoporosis and osteoarthritis. (29, 30) In terms of preclinical evidence, conjugated equine estrogen demonstrated significant anti-inflammatory activity in a rat model. (31) Several investigators have reported the presence of estrogen receptors in cartilage. (32-34) When adult monkeys were treated with conjugated equine estrogens, chondrocytes incorporated higher levels of sulfate in proteoglycans compared to baseline. (35) Decreased estrogen results in increased release of proinflammatory cytokines from monocytes and
macrophages. (36) The proinflammatory cytokines, interleukin-1 (IL-1) and tumor necrosis factor (TNFα) promote cartilage reabsorption, inhibit synthesis of proteoglycans and cause inflammation. (37-39) From these preclinical results, it seems clear that close interactions exist between estrogen and inflammation as well as cartilage metabolism.

The identification of the two estrogen receptors α and β in human articular chondrocytes provided further evidence that cartilage is sensitive to estrogens. (32, 40-42) Various animal and in vitro studies suggest that estrogen may play a role in the regulation of cartilage turnover and development of joint disease. In an experimental model of postmenopausal osteoarthritis with ovariectomized rats, estrogen deficiency accelerated cartilage turnover and increased cartilage surface erosion, whereas administration of estrogen or SERMs significantly suppressed cartilage degradation. (43) In ovariectomized rats, estrogen prevented the cartilage breakdown caused by IL-6 (44), a proinflammatory cytokine that plays a critical role in the pathogenesis of osteoarthritis (OA). (45) In a sheep model, estrogen replacement therapy reduced the loss of proteoglycans from cartilage adjacent to OA lesions, providing evidence for a chondroprotective effect of estrogen. (46)

Estrogen can influence chondrocyte formation on multiple levels by interacting with cellular growth factors, adhesion molecules, and cytokines. In vitro studies showed a dose-dependent change in matrix protein turnover when cultured chondrocytes were exposed to estradiol. (33, 34, 47, 48) Production of IL-6 and type II collagen in articular chondrocytes was affected by estradiol, suggesting possible mechanisms whereby it may affect cartilage metabolism. (49, 50) In addition, estrogens may decrease the acceleration in subchondral bone remodeling, which is a key factor in the pathophysiology of osteoarthritis. (51)

Several small studies have also evaluated the influence Al therapy has on inflammatory serum markers. Results evaluating CRP, IL-6, and TNFα have been inconsistent. (52-54) Even in studies showing Al-related changes to inflammatory markers, such as CRP, these changes did not correlate with symptoms of Al-induced arthralgia. (53) One study involving 57 women on AIs found an increase in ANA and anti-DNA positivity rates: 40% vs 19% for ANA and 14% vs 0% for anti-DNA. (54) A case-control study evaluating 24 women with Al-related arthralgia and 30 without, were evaluated for a wide range of inflammatory markers. While many of the assayed factors including interferons, interleukins, matrix metalloproteinases (MMPs), and chemokines were different at baseline, these differences did not persist at 6 months. This suggests a possible inflammatory phenotype that predicts the development of this toxicity. (55) These studies were limited by small samples sizes.

The exact mechanism of Al-related arthralgia is unclear, but is thought to be related to estrogen deprivation. The effects of estrogen on bone, endometrium, and breast tissue have been extensively studied. Cartilage is not generally viewed as an estrogen responsive tissue, however, several epidemiological studies and a few intervention studies support that estrogen may have a role in osteoarthritis. OA is a disease of joints, involving both cartilage and bone. Progressive and permanent articular cartilage degeneration is the hallmark of osteoarthritis. (56) Two biochemical markers, serum cartilage oligomeric matrix protein (COMP) and urine crosslinked C-telopeptides of Type II collagen (CTX-II) were shown to correlate with symptoms and x-ray findings of knee OA. (57) Urinary CTX-II, a biochemical marker of type II collagen breakdown, has been negatively correlated with joint surface area and was an important predictor of joint damage assessed by radiographs. (58) Factors influencing the incidence of radiological osteoarthritis include obesity and female gender, especially after entering the menopause. (59, 60) Over 80 years ago, Cecil and Archer first described “arthritis of the menopause” as the rapid development of hand and knee osteoarthritis coinciding with cessation of menses. (61) Various studies report larger increases in women than in men in the incidence and prevalence of hip, knee, and hand osteoarthritis after 50 years of age. (62) Sowers et al. reported significant associations of lower serum estradiol and urinary metabolites with the development of knee osteoarthritis in women. (63) However, other studies have not found a consistent link between osteoarthritis and sex hormone levels. (64-66)
Findings in epidemiologic studies regarding a correlation between estrogen and osteoarthritis are inconsistent. However, observational studies of the incidence and prevalence of osteoarthritis in postmenopausal women with and without hormone replacement therapy (HRT) have supported a protective effect of estrogens in osteoarthritis. In a cross-sectional study by Nevitt et al., women taking HRT had a lower relative risk of developing hip osteoarthritis (OR 0.62; 95% CI 0.49-0.86) compared to women not taking HRT. The protective effect was greater in women on HRT for more than ten years with OR 0.57 (95% CI 0.40-0.82). Two studies found that HRT tended to reduce the incidence of radiological knee osteoarthritis. In a study of women without osteoarthritis, HRT taken for longer than 5 years was associated with a larger volume of tibial cartilage as compared to women who were not taking HRT. The number of years since menopause was more strongly related to diminished knee cartilage volume than age per se, suggesting an effect of hormonal status on knee cartilage. In a randomized controlled trial of HRT, the Heart and Estrogen/progestin Replacement Study (HERS) found no difference in the prevalence of knee pain with or without HRT (24.1% and 26.1%, respectively). However, the Women's Health Initiative intervention trial of HRT recently reported a difference in the incidence of any joint pain or swelling in postmenopausal women on estrogen compared to those who were not on estrogen (70.6% versus 77.2%, p=0.01). In elderly postmenopausal women, estrogen may be an important regulator of osteoarthritis. The long-term effects of profound estrogen suppression in breast cancer patients taking AIs are unknown. The acute and long-term side effects of AIs are becoming an increasingly important issue as more and more women are being treated with these agents. Musculoskeletal pain may be associated with a deterioration of quality of life due to physical disability, sleep disturbance, impaired cognitive function, depression and anxiety, and increased utilization of health care resources. Therefore, targeted interventions that relieve AI-induced musculoskeletal pain are needed.

**Current Treatment for Aromatase Inhibitor-Induced Joint Pain**

Non-steroidal anti-inflammatory drugs (NSAIDs), the most common medications used to treat pain associated with arthritis, may lead to serious renal and gastrointestinal toxicity, particularly in older adults. Selective cyclooxygenase-2 (COX-2) inhibitors, such as rofecoxib and celecoxib, are effective analgesics with fewer gastrointestinal side effects but have recently been shown to elevate the risk of cardiovascular events. Non-toxic treatments that effectively relieve AI-induced musculoskeletal pain are needed. Ongoing studies are evaluating the potential benefits of physical activity, vitamin D, glucosamine/chondroitin and omega-3 fatty acids for treating this toxicity. Since research in this area is limited, a small pilot study evaluating the use of acupuncture to relieve symptoms of AI-associated arthralgias was initially conducted. In this study of 21 women treated with a 6-week course of total body and auricular acupuncture, improvements were reported in pain severity, pain-related functional outcomes, and physical well-being, and no significant adverse events were reported. This study was limited due to its small sample size and lack of an adequate control group. Investigators then went on to conduct a randomized, blinded, sham-controlled study in 38 breast cancer patients with AI-associated arthralgias. True acupuncture for 6 weeks was associated with about a 50% decrease in mean BPI-SF scores compared to sham acupuncture with superficial needling at non-acupuncture points. Similar findings were seen for the WOMAC and M-SACRAH scores. Results found that acupuncture is an effective and well-tolerated strategy for managing this common treatment-related side effect.

**Acupuncture for the Treatment of Joint Pain**

Approximately 25% of the population over the age of 55 years are affected by joint pain and about half of them will have some restriction of normal daily activities. Virtually all individuals over the age of 65 have radiographic evidence of osteoarthritis. With conventional
approaches such as analgesics and exercise, controlling pain and minimizing loss of function are the principal aims of treatment. (63, 84) About two-thirds of individuals who suffer from joint pain have used complementary and alternative treatments to control their symptoms. (85) However, due to the dearth of rigorously derived experimental evidence, most third party payers do not reimburse patients for the cost of many of these procedures.

Acupuncture is a traditional Chinese method of medical treatment involving the insertion of fine, single-use, sterile needles in acupoints according to a system of channels and meridians that was developed by early practitioners of Traditional Chinese Medicine (TCM) over 2000 years ago. The needles are stimulated by either manual manipulation, electrical stimulation, or heat. (86) Patients in pain have imbalance in their Qi or vital energy. The Qi travels through 12 main meridians in the body, connecting the body in a weblike matrix. (87) Within the traditional framework, meridians are thought to represent conduits for electrical signals. (88) During traditional acupuncture, needles are inserted into a meridian point and then manipulated to elicit “de qi.” De qi has been described as a sensation of aching, soreness, or dull pain. According to TCM, elicitation of de qi re-establishes normal flow of Qi and is essential for clinical efficacy. More recently, it has been postulated that sensation of de qi represents activation of nerve receptors during insertion of acupuncture needles at the appropriate location and depth of traditional acupuncture points. (89) Currently, acupuncture is often used with TCM and it is a recognized health profession with strict licensure and regulatory status in 40 states. (90)

Acupuncture is a popular non-pharmacological modality used for treating a variety of conditions, including musculoskeletal pain. Acupuncture has been shown to have short-term analgesic effect in musculoskeletal pain. (91, 92) Clinical trials have found a benefit to patients with knee osteoarthritis when acupuncture is used as an adjunct to conventional management strategies. (93) In a randomized study of 97 patients with osteoarthritis of the knee, acupuncture as a complementary therapy to pharmacological treatments is more effective than pharmacological treatment alone, in terms of reducing pain, improving physical function and health-related quality of life. (94) A trial conducted in Germany concluded that true acupuncture has a better effect than sham acupuncture in the treatment of knee and back pain. (95) However, in two large multicenter trials of acupuncture for osteoarthritis of the knee, one found a benefit in joint pain and function compared to sham acupuncture and the other found no additional improvement in pain scores. (96, 97)

Acupuncture and TCM vary slightly from Western biomedicine in the methods of physical examination, evaluation and diagnostic criteria. For example, the major diagnostic criteria of TCM practitioners include inquiry, inspection of the body, acupuncture channels or meridians, and the tongue, as well as palpation of the channels, points and pulse. Pulse palpation in TCM occurs at the radial artery in the wrist and is assessed for rate, rhythm, force and shape, which all correspond to internal physiological processes according to TCM theory. Using these criteria, a differential diagnosis is made and an appropriate point prescription is applied as treatment.

Acupuncture points are located on the body using anatomical locations and the TCM standard unit of body measurement termed the “cun.” As everyone's body has different dimensions, a cun is defined according to the person whose body is to be treated. For example, 1 cun is equivalent to the width of the thumb, in the middle, at the crease; 3 cun is equivalent to the combined breadth of the 4 fingers, at the level of the pinky finger's first joint above the palm of the hand; and 12 cun is equivalent to the distance from the elbow crease to the wrist crease.

Auricular acupuncture therapy has long been used in China to treat conditions by stimulating certain points on the ear with needles. The history of its use has been documented in various historical texts and auricular therapy is maintained in common use in the present day system of acupuncture. The National Acupuncture Detoxification Association (NADA) protocol has been extensively studied for its effect on addiction and stress relief. This protocol is endorsed by the National Institute on Drug Abuse (NIDA), the governing body of the treatment of chemical dependency, and is being used in some states such as Florida and California within their drug rehabilitation and court systems. (98-100) The function of the NADA protocol with full body acupuncture treatment is to relieve pain and decrease stress.
The most common presentation of musculoskeletal pain according to TCM is “Bi Syndrome.” “Bi” means obstruction or blockage. “Bi Syndrome” describes a set of patterns in which obstruction of qi and blood through the channels causes pain. Qi and blood are defined as the basic elements of all physiological activity according to TCM. The TCM differential diagnosis of “Bi syndrome” encompasses several disorders known in Western biomedicine, such as osteoarthritis, rheumatoid arthritis, rheumatic fever, fibromyalgia, gout and sciatica. (101-103) Of interest to this study, AI-induced musculoskeletal pain follows the pattern diagnosis of “Bi syndrome.”

The use of complementary and alternative medicine (CAM) is growing in the U.S. According to a Center for Disease Control (CDC) report of the 2002 National Health Interview Study, 75% of respondents reported using some form of CAM. (104) Breast cancer patients are the cancer group most likely to use CAM therapies in conjunction with their conventional treatment. (105-107) Therapies used include nutritional and herbal supplements, massage, dietary changes, prayer, yoga and acupuncture. (108-110) An increasing amount of evidence suggests that acupuncture has a role in the supportive care of cancer patients for treating chemotherapy-induced nausea and vomiting, xerostomia, leukopenia, and other chemoradiotherapy-induced symptoms. (111-116) A recent study found that acupuncture treatments given twice weekly for 4 weeks; then weekly for 8 weeks (12 weeks total) was as effective as the antidepressant, venlafaxine, for managing vasomotor symptoms related to hormonal therapy in breast cancer patients. (117) In fact, acupuncture had a more durable effect with fewer side effects compared to venlafaxine and had additional benefits, including increased sex drive, improved energy and sense of well-being.

Acupuncture has also been demonstrated to be safe to administer to breast cancer patients as symptom management for a wide variety of conditions including hot flashes and nausea. Acupuncture has been demonstrated to be safe in two previous studies assessing acupuncture feasibility and safety in treating upper extremity lymphoedema. (160,161)

Mechanism of Action of Acupuncture

Ideally, relief from AI-associated arthralgia should be through non-hormonal mechanisms, so as not to interfere with the efficacy of the drug. Acupuncture is under current investigation for its mechanistic effect within the biomedical research paradigm and its influence on particular conditions or symptom sets. Although the precise mechanism of action is unknown, acupuncture analgesia is thought to be mediated by central mechanisms of pain control through the release of specific neurotransmitters, such as endorphin. (118-120) A recent study suggested that acupuncture modulates pain signals in the central nervous system at the level of the spinal cord, midbrain and cortex via release of endogenous opiates and neurotransmitters. (121, 122) Early studies show that naloxone, an antagonist to opiates, attenuated the analgesic effects of acupuncture in humans and mice. (120, 123) In humans after receiving acupuncture, elevated levels of β endorphin in the cerebrospinal fluid have been observed. (119) Animal studies with electro-acupuncture (EA) have shown that EA at low frequency of 2 Hz stimulates the release of β endorphins, enkephalin and endomorphin, which activates μ and δ opiate receptors. (124) In addition, EA at higher frequency stimulates release of dynorphin which activates the κ opiate receptors. (125)

Another proposed mechanism of acupuncture analgesia involves the serotonergic pathway. Acupuncture has been shown to activate the release of serotonin from the serotonergic raphe spinal neurons in the nucleus raphemagnus, which is associated with the descending pain inhibitory pathway. (126) Serotonin then binds serotonin receptors on inhibitory interneurons in the spinal cord to release enkephalin, which inhibits pain sensation signals. (127) A more recent study in rats found that acupuncture activate neurons in the periductal gray matter of the midbrain, a region in the brain that contributes to the descending pain inhibitory pathway. EA also had an anti-nociceptive effect by abolishing histamine and dopamine release induced by pain stimuli. (128) More recently, researchers are looking at the anti-hyperalgesic effect of acupuncture in inflammatory mouse models. In one in vivo study employing a persistent inflammatory pain rat model, administration of EA increased paw withdrawal latency suggestive of anti-hyperalgesic effect. In addition, EA was able to reduce inflammatory paw edema in the rats. (129, 130)
Sham Acupuncture Interventions

The optimal control group for studies of acupuncture is controversial. Sham acupuncture methods include acupuncture for an unrelated condition, needle insertion at nonacupoint locations, noninsertive simulated acupuncture, or use of retractable needles. However, most acupuncture controls are not completely inert. Potential benefits may be conferred from the physiologic effect of needling even when not performed according to established principles. (131-134) Detrimental effects from sham needling are also possible, and other studies have found no benefit in the sham group. (134)

In the prior sham-controlled study of acupuncture for AI-related joint symptoms, the sham intervention used superficial needle insertion at body locations not recognized as true acupoints. (80) Results showed that 22% of subjects who received sham acupuncture reported at least a 2-point decrease in BPI-SF worst pain scores, consistent with the literature that reports a 25-55% placebo effect for nonacupoint and superficial insertion sham methods. (136) In this study, 57% of patients in the sham group believed they were receiving true acupuncture, suggesting that blinding was effective. For the auricular acupuncture, the sham intervention will involve adding adhesives to the selected points on the helix of the auricle. While a sham auricular intervention was not applied in the prior study, investigators feel that application of the sham auricular intervention may increase the effectiveness of blinding. While future studies may attempt to look at other point protocols and acupuncture interventions oriented in other traditional theories, investigators feel that it is important to remain consistent in this study design. Moreover, investigators will control for a placebo effect based on previous research as reflected in the statistical methods.

Patient Reported Outcomes for Joint Symptoms

This study will rely on a patient-reported outcome (PRO) for the primary endpoint. Traditionally, the Food and Drug Administration (FDA) requires substantial evidence for reaching a conclusion that a drug or intervention will have an effect on a given condition. Evidence that shows not only a change in symptoms, but how that change translates to other specific endpoints, such as the patients' ability to perform activities, is sometimes required by the FDA for approval. (136) The PRO of symptomatic and functional improvement may benefit from the use of objective measures, such as magnetic resonance imaging (MRI) of joints to assess for structural changes or functional assessments, including grip strength. (22)

Studies assessing AI-induced arthralgias have shown a correlation between PROs and objective findings. Morales et al. demonstrated that the subjective symptoms of AI-induced arthralgias in the hands are associated with physiologic changes to the joint and functional impairments. (22) The patients in this study were assessed with serial MRIs of the hands, measurements of grip strength, and symptoms self-assessments. In a 6-month period, women taking AIs were more likely than those on tamoxifen to have an increase in tenosynovial changes as seen on MRI, a decrease in grip strength as measured by a sphygmomanometer, as well as increased pain and stiffness as measured by self-administered questionnaire. (22)

Since there are no well-validated measures specifically for AI-induced arthralgias, scales that captured joint pain, stiffness, and functional status in the hips and knees (WOMAC) and hands (M-SACRAH) were selected, and a general pain scale used in cancer patients (BPI-SF). The BPI-SF (137) is a 14-item questionnaire which asks patients to rate pain over the prior week and the degree to which it interferes with activities on a 0 to 10 scale. Numeric rating scales such as the BPI are among the most common, valid, and reliable measures used to assess cancer pain. (138) The Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC) Version 3.1 is a validated measure for assessing osteoarthritis of the knees or hips and consists of 24 questions related to three subscales: pain, stiffness, and physical function. (139) The Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands (M-SACRAH) consists of three domains assessing pain, stiffness, and functional status in patients suffering from hand osteoarthritis and rheumatoid arthritis. (140) Well-validated quantitative
functional testing with grip strength to assess hand function and the “Timed Get Up and Go” (TGUG) test to assess functional status in the lower extremities have also been included. (141,142)

**Acupuncture for Aromatase Inhibitor-Related Arthralgias**

Researchers at CUMC have extensive experience evaluating interventions for the treatment of AI-induced arthralgias. Since research in this area is limited, a small pilot study evaluating the use of acupuncture to relieve symptoms of AI-associated arthralgias was initially conducted. In this study of 21 women treated with full body and auricular acupuncture twice weekly for six weeks, improvements were reported in pain severity, pain-related functional outcomes, and physical well-being, and no significant adverse events were reported. (79) Another pilot study used electro-acupuncture administered twice weekly for 2 weeks followed by 6 weekly treatments in 12 breast cancer survivors with AI-related arthralgia. (143) Similarly, the patients reported significant reductions in pain severity, stiffness, and joint symptom interference according to the BPI. Subjects also reported a significant decrease in fatigue and anxiety. These studies were limited due to small sample sizes, unblinded interventions, and lack of an adequate control group.

A randomized, blinded, sham-controlled study in 38 women with early stage breast cancer and AI-induced arthralgias to assess the efficacy of acupuncture was then conducted. As is shown in Figure 1, a 6-week course of true acupuncture was associated with about a 50% decrease in mean BPI-SF scores. No change from baseline was observed for the sham arm. At 6 weeks, the mean BPI-SF worst pain scores (range, 0-10) were lower for true acupuncture compared to sham acupuncture (3.0 vs. 5.5, p=0.001), as well as pain severity (2.6 vs. 4.5, p=0.003) and pain-related interference (2.5 vs. 4.5, p=0.002). Similar findings were seen for the WOMAC and M-SACRAH scores which assess knee/hip and hand joint symptoms, respectively. (144) These measures were very effective at capturing the symptoms of AI-related arthralgia.

**Summary**

Due to early detection and improved treatments, women with breast cancer are living longer. The increase in breast cancer survival is largely due to the benefits of hormonal therapy, such as tamoxifen and aromatase inhibitors (AIs), for the treatment of hormone-sensitive breast cancer. Recent clinical trials have demonstrated that AIs are more effective than tamoxifen at reducing breast cancer recurrences. (8-13) However, breast cancer patients receiving AIs have a higher incidence of osteoporosis, bone fractures and musculoskeletal symptoms, particularly joint pain and stiffness. Musculoskeletal pain, which occurs in up to 50% of patients treated with AIs (18), may lead to noncompliance, deterioration of quality of life due to physical disability, sleep disturbance, impaired cognitive function, depression and anxiety, and increased utilization of health care resources. (73-75) Since women with hormone receptor-positive breast cancer require long-term hormonal therapy, an important objective is minimizing long-term side effects to enhance patient compliance and improve quality of life. Therefore, targeted interventions that relieve AI-induced musculoskeletal pain are needed.
There are currently no proven treatments for AI-induced arthralgias. Acupuncture is a popular non-pharmacologic modality that has been shown to have analgesic effects in many chronic pain syndromes, including osteoarthritis. A recent evaluation of acupuncture by the National Institutes of Health concluded that it is a useful adjunct in a range of painful conditions, including musculoskeletal pain. (145) While the use of complementary and alternative medicine for the treatment of persistent musculoskeletal pain continues to increase, rigorous clinical trials examining their efficacy are needed before definitive recommendations can be made. Investigators recently conducted a randomized, blinded, sham-controlled trial to assess the effect of acupuncture on musculoskeletal pain induced by AIs. The true acupuncture group had about a 50% decrease in mean BPI-SF worst pain score compared to the sham acupuncture group (3.0 vs 5.5, p<0.001). (80) There were also differences in pain severity and pain related interference. Acupuncture was well tolerated with no complications. However, despite these promising results, this single institution study had a relatively small sample size, had a single acupuncturist, and was only a 6-week intervention. (80) For the community, cost and lack of insurance coverage limit access to this therapy. A larger, confirmatory, multicenter trial is necessary to truly control for confounding factors, and may provide the data critical to extend coverage for this indication. This approach is innovative because it is a new application of a well recognized pain therapy for breast cancer supportive care. Improved adherence is critical for this class of medications, given the large discontinuation rate due to toxicity, which has been shown to affect survival. (17)

For this current proposal, SWOG will conduct a multicenter randomized, blinded, sham- and waitlist-controlled clinical trial to test the effect of acupuncture on AI-associated joint pain in 228 postmenopausal women with early stage hormone receptor-positive breast cancer who report joint pain that has increased or started since initiation of AI therapy. The primary endpoints are to compare the mean BPI-SF worst pain scores at 6 weeks in the true acupuncture compared to sham acupuncture and waitlist groups in order to confirm initial results of a 6-week acupuncture intervention given twice weekly in a larger multicenter study. Secondary exploratory endpoints include evaluations at 12, 24, and 52 weeks to determine the benefit of additional 6 weekly acupuncture treatments for maintenance and to determine the durability of response 12 and 40 weeks after stopping acupuncture, respectively. This study will be the first large multicenter trial to investigate the effect of acupuncture in treating AI-induced joint symptoms in breast cancer patients. This is a common complaint in this patient population which can lead to the discontinuation of a life-saving therapy. Research has shown that even taking this medication less than 80% of the time can have a negative impact on survival. (17) Results from this study will inform clinicians, insurance companies and patients on whether or not this is an effective therapy, and may help predict who is most likely to benefit from this therapy.

Inclusion of Minorities

This clinical trial was designed to include minorities, but was not designed to measure differences of intervention in minority subgroups. It is expected that the patients accrued to this trial will reflect the minority representation in the local population of the participating sites. Men with breast cancer are not eligible for this study because it is not standard for them to receive aromatase inhibitors. Anticipated accrual to this study by race and ethnicity, based on previous Group trials in this disease type, follows:
### Ethnic Category

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### Racial Category

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<td><strong>Racial Category: Total of all Subjects</strong></td>
<td><strong>228</strong></td>
<td><strong>228</strong></td>
</tr>
</tbody>
</table>

3.0 **DRUG INFORMATION**

There are no drugs used in this study.

4.0 **STAGING CRITERIA**


**DEFINITION OF TNM**

**Primary Tumor (T)**

Definitions for classifying the primary tumor (T) are the same for clinical and for pathologic classification. If the measurement is made by physical examination, the examiner will use the major headings (T1, T2 or T3). If other measurements, such as mammographic or pathologic measurements are used, the subsets of T1 can be used. Tumors should be measured to the nearest 0.1 cm increment.

- **TX** Primary tumor cannot be assessed
- **T0** No evidence of primary tumor
- **Tis** Carcinoma in situ
- **Tis (DCIS)** Ductal carcinoma in situ
- **Tis (LCIS)** Lobular carcinoma in situ
- **Tis (Paget's)** Paget's disease of the nipple with no tumor

**NOTE:** Paget's disease associated with a tumor is classified according to the size of the tumor.
T1  Tumor 2 cm or less in greatest dimension
T1mic Microinvasion 0.1 cm or less in greatest dimension
T1a Tumor more than 0.1 cm but no more than 0.5 cm in greatest dimension
T1b Tumor more than 0.5 cm but not more than 1 cm in greatest dimension
T1c Tumor more than 1 cm but not more than 2 cm in greatest dimension
T2 Tumor more than 2 cm but not more than 5 cm in greatest dimension
T3 Tumor more than 5 cm in greatest dimension
T4 Tumor of any size with direct extension to (a) chest wall or (b) skin, only as described below:
T4a Extension to chest wall, not including pectoralis muscle
T4b Edema (including peau d'orange) or ulceration of the skin of the breast, or satellite skin nodules confined to the same breast
T4c Both T4a and T4b
T4d Inflammatory carcinoma

Regional Lymph Nodes (N)

Clinical

NX Regional lymph nodes cannot be assessed
N0 No regional lymph node metastasis
N1 Metastasis to movable ipsilateral axillary lymph node(s)
N2 Metastasis in ipsilateral axillary lymph nodes fixed or matted, or in clinically apparent* ipsilateral internal mammary nodes in the absence of clinically evident axillary lymph node metastasis
N2a Metastasis in ipsilateral axillary lymph nodes fixed to one another (matted) or to other structures
N2b Metastasis only in clinically detected* ipsilateral internal mammary nodes and in the absence of clinically evident axillary lymph node metastasis
N3 Metastasis in ipsilateral infraclavicular lymph node(s) with or without axillary lymph node involvement, or in clinically detected* ipsilateral internal mammary lymph node(s) and in the presence of clinically evident axillary lymph node metastasis; or metastasis in ipsilateral supraclavicular lymph node(s) with or without axillary or internal mammary lymph node involvement
N3a Metastasis in ipsilateral infraclavicular lymph node(s)
N3b Metastasis in ipsilateral internal mammary lymph node(s) and axillary lymph node(s)
N3c Metastasis in ipsilateral supraclavicular lymph node(s)

*Clinically detected is defined as detected by imaging studies (excluding lymphoscintigraphy) or by clinical examination or grossly visible pathologically.

Pathologic (pN)*

pNX Regional lymph nodes cannot be assessed
pN0 No regional lymph node metastasis identified histologically

NOTE: Isolated tumor cells (ITC) are defined as single tumor cells or small cell clusters not greater than 0.2 mm, usually detected only by immunohistochemical (IHC) or molecular methods but which may be verified on H&E stains. ITCs do not usually show evidence of malignant activity e.g., proliferation or stromal reaction.
pN0 (i-) No regional lymph node metastasis histologically, negative IHC

pN0 (i+) Malignant cells in regional lymph node(s) no greater than 0.2mm (detected by H&E or IHC including ITC)

pN0(mol-) No regional lymph node metastasis histologically, negative molecular findings (RT-PCR)

pN0(mol+) No regional lymph node metastasis histologically, positive molecular findings (RT-PCR)

a Classification is based on axillary lymph node dissection with or without sentinel lymph node dissection. Classification based solely on sentinel lymph node dissection without subsequent axillary lymph node dissection is designated (sn) for "sentinel node," e.g., pN0 (i+) (sn).

b RT-PCR: reverse transcriptase/polymerase chain reaction.

pN0 Metastasis in 1 to 3 axillary lymph nodes, and/or in internal mammary nodes with microscopic disease detected by sentinel lymph node dissection but not clinically detected**

pN1 Metastasis in 1 to 3 axillary lymph nodes, at least one metastasis is greater than 2.0mm

pN1a Micrometastasis (greater than 0.2 mm, none greater than 2.0 mm)

pN1b Metastasis in internal mammary nodes with microscopic disease detected by sentinel lymph node dissection but not clinically detected.**

pN1c Metastasis in 1 to 3 axillary lymph nodes and in internal mammary lymph nodes with microscopic disease detected by sentinel lymph node dissection but not clinically detected.** (If associated with greater than 3 positive axillary lymph nodes, the internal mammary nodes are classified as pN3b to reflect increased tumor burden)

pN2 Metastasis in 4 to 9 axillary lymph nodes or in clinically detected* internal mammary lymph nodes in the absence of axillary lymph node metastasis

pN2a Metastasis in 4 to 9 axillary lymph nodes (at least one tumor deposit greater than 2.0 mm)

pN2b Metastasis in clinically detected* internal mammary lymph nodes in the absence of axillary lymph node metastasis

pN3 Metastasis in 10 or more axillary lymph nodes, or in infraclavicular lymph nodes, or in clinically detected* ipsilateral internal mammary lymph nodes in the presence of 1 or more positive axillary lymph nodes; or in more than 3 axillary lymph nodes with clinically negative microscopic metastasis in internal mammary lymph nodes; or in ipsilateral supraclavicular lymph nodes

pN3a Metastasis in 10 or more axillary lymph nodes (at least one tumor deposit greater than 2.0 mm), or metastasis to the infraclavicular lymph nodes

pN3b Metastasis in clinically detected* ipsilateral internal mammary lymph nodes in the presence of 1 or more positive axillary lymph nodes; or in more than 3 axillary lymph nodes and in internal mammary lymph nodes with microscopic disease detected by sentinel lymph node dissection but not clinically detected**

pN3c Metastasis in ipsilateral supraclavicular lymph nodes

* Clinically detected is defined as detected by imaging studies (excluding lymphoscintigraphy) or by clinical examination.

** Not clinically detected is defined as not detected by imaging studies (excluding lympho-seintigraphy) or by clinical examination.
### Distant Metastasis (M)

M0  No distant metastasis

#### STAGE GROUPING

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<td>T4</td>
<td>N2</td>
<td>M0</td>
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<tr>
<td>Stage IIIC</td>
<td>Any T</td>
<td>N3</td>
<td>M0</td>
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</tbody>
</table>

*T1 includes T1mic

** T0 and T1 tumors with nodal micrometastases only are excluded from Stage IIA and are classified Stage IB.

**NOTE**: Stage designation may be changed if post-surgical imaging studies reveal the presence of distant metastases, provided the studies are carried out within 4 months of diagnosis in the absence of disease progression and provided that the patient has not received neoadjuvant therapy.
5.0 ELIGIBILITY CRITERIA

Each of the criteria in the following section must be met in order for a patient to be considered eligible for registration. Use the spaces provided to confirm a patient's eligibility. For each criterion requiring test results and dates, please record this information on the Prestudy Form and submit to the Data Operations Center in Seattle (see Section 14.0). Any potential eligibility issues should be addressed to the Data Operations Center in Seattle at 206/652-2267 prior to registration.

In calculating days of tests and measurements, the day a test or measurement is done is considered Day 0. Therefore, if a test is done on a Monday, the Monday four weeks later would be considered Day 28. This allows for efficient patient scheduling without exceeding the guidelines. If Day 28 falls on a weekend or holiday, the limit may be extended to the next working day.

SWOG Patient No. __________________________

Patient's Initials (L, F, M) __________________________

5.1 Disease Related Criteria

____ a. Patients must be women with histologically confirmed primary invasive carcinoma of the breast (Stage I, II, or III) with no evidence of metastatic disease (M0) (see Section 4.0) or with histologically confirmed DCIS. If patient has undergone breast cancer surgery, she must have recovered from all side-effects of surgery.

____ b. Patients must be postmenopausal, as defined by at least one of the following:

1. ≥ 12 months since the last menstrual period OR
2. prior bilateral oophorectomy OR
3. current use of a GnRH agonist OR
4. previous hysterectomy with one or both ovaries left in place (or previous hysterectomy in which documentation of bilateral oophorectomy is unavailable) AND FSH values consistent with the institutional normal values for the postmenopausal state. If patient is under the age of 55, FSH levels must be obtained within 28 days prior to registration.

____ c. Patients must be positive for either estrogen receptor (ER) and/or progesterone receptor (PgR) as determined by institutional standard.

5.2 Prior/Current Therapy Criteria

____ a. Patients must currently be taking a third-generation aromatase inhibitor (AI) – anastrozole, letrozole, or exemestane for at least the previous 30 days prior to registration with plans to continue for at least an additional 1 year after registration. Patients may have switched AIs provided that they have been on a stable dose for at least 30 days. Concurrent trastuzumab (herceptin) is allowed.

5.3 Clinical/Laboratory Criteria

____ a. Patients must have completed the S1200 Brief Pain Inventory-Short Form (BPI-SF) within 14 days prior to registration. Patients must have a worst pain score of at least 3 on the Brief Pain Inventory (item #2) that has started or increased since starting AI therapy.
b. Patients must have a Zubrod performance status of 0 to 1 (see Section 10.3).

c. Patients must have had two or fewer prior acupuncture treatments within the past 12 months for any reason except for joint symptoms. Patients must not have had prior acupuncture treatment for joint symptoms at any time.

d. Patients must not have a severe bleeding disorder.

e. Patients must not have an allergy to latex.

f. Patients must not have concurrent medical/arthritic disease that could confound or interfere with evaluation of pain or efficacy including: inflammatory arthritis (e.g., rheumatoid arthritis, systemic lupus, spondyloarthropathy, psoriatic arthritis, polymyalgia rheumatica), gout, episodes of acute monarticular arthritis clinically consistent with pseudogout, Paget’s disease affecting the study joint (knees/hands), a history of septic arthritis or avascular necrosis or intra-articular fracture of the study joint, Wilson’s disease, hemochromatosis, alkaptonuria, or primary osteochondromatosis.

g. Patients must not have a history of bone fracture or surgery of the afflicted knees and/or hands within 6 months prior to registration.

h. Patients must not have a history of illness that, in the opinion of the investigator, might confound the results of the study or pose additional risk to the patient.

i. Patients must not be on narcotics within 14 days of registration.

j. Patients must not have received oral corticosteroids, intramuscular corticosteroids, or intra-articular steroids for joint symptoms within 28 days prior to registration.

k. Patients must not have received topical analgesics (e.g., capsaicin preparations) or any other analgesics (e.g., opiates, tramadol, with the exception of NSAIDS, combination NSAIDs, and acetaminophen) within 14 days prior to registration.

l. Patients must not have received or implemented any other medical therapy, alternative therapy or physical therapy for the treatment of joint pain/stiffness within 28 days prior to registration. Therapeutic massage is allowed.

m. Patients must be able to complete study questionnaires in English or Spanish.

n. No other prior malignancy is allowed except for adequately treated basal cell or squamous cell skin cancer, in situ cervical cancer, DCIS, adequately treated Stage I or II cancer from which the patient is currently in complete remission, or any other cancer for which the patient has been disease-free for > 5 years.

o. Prestudy history and physical must be obtained within 180 days prior to registration.
5.4 Specimen Submission Criteria

- a. Patients must be willing to submit blood and urine samples for serum hormones (estradiol, FSH, LH), inflammatory biomarkers (serum TNFα, IL-6, IL-12, CRP and urine CTX-II), urine AI metabolites, and DNA analysis (CYP19A1), and must be given the option to consent to use of remaining specimens for future translational medicine studies as outlined in Section 15.0. Baseline samples must be obtained prior to beginning intervention.

5.5 Regulatory Criteria

- a. All patients must be informed of the investigational nature of this study and must sign and give written informed consent in accordance with institutional and federal guidelines.

- b. At the time of patient registration, the treating institution’s name and ID number must be provided to the Data Operations Center in Seattle in order to ensure that the current (within 365 days) date of institutional review board approval for this study has been entered into the database.
6.0 STRATIFICATION FACTORS

Patients will be randomized using a 2:1:1 ratio to true acupuncture vs sham acupuncture vs waitlist control. Patient randomization will be dynamically balanced according to study site at time of registration.

7.0 PROCEDURES AND STUDY PLAN

For study-related questions, please contact Dr. Dawn Hershman at 212/305-1945 or Dr. Katherine Crew at 212/305-1732.

7.1 General Considerations

All patients, whether receiving acupuncture treatments (either true or sham) or assigned to the waitlist control group, will have the same clinic visit and telephone contact schedule, including patient-completed questionnaires and functional testing. See Appendix 18.1 for a description of true acupuncture, sham acupuncture, waitlist control.

7.2 Initial Visit

Before beginning the intervention, patients will have completed the following self-administered questionnaires:

- **S1200 Brief Pain Inventory-Short Form (BPI-SF)**
- **S1200** Western Ontario and McMaster Universities Osteoarthritis (WOMAC) Index (Version 3.1)
- **S1200** Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands (M-SACRAH)
- **S1200** PROMIS Pain Impact-Short Form (PROMIS PI-SF)
- **S1200** FACT-ES Trial Outcome Index (Version 4)
- **S1200** Aromatase Inhibitor Usage Form

It should take the patient about 30 minutes to complete the assessments. See Section 15.2 for instructions for administration of the questionnaires. The nurse or CRA must complete the **S1200** Cover Sheet for Patient-Completed Questionnaires. Use of pain medications will be recorded on the **S1200** Supplemental Agents Reporting Form.

Functional testing (grip strength, ‘Timed Get Up and Go’) will be conducted and recorded on the **S1200** Functional testing. See Section 7.5 for instructions for functional testing.

Additionally, blood will be drawn for serum hormones (estradiol, FSH, LH) and inflammatory markers (TNFα, IL-6, IL-12, CRP) and for DNA analysis (CYP19A1) (see Section 15.1). Urine will be collected for urine CTX-II and urine Al metabolites.

7.3 Follow-Up Visits

a. Follow-up visits will occur at 6 and 12 weeks after registration.

Acupuncture adherence will be ascertained by the CRA and documented on the **S1200** Adherence to Acupuncture Form. Adverse events will be assessed and recorded on the **S1200** Adverse Event Summary Form. Use of pain medications will be recorded on the **S1200** Supplemental Agents Reporting Form. At 6 weeks and 12 weeks, to assess the adequacy of blinding, patients in the true and sham acupuncture groups will give an opinion about their treatment assignment. All of this data will be recorded on the **S1200** Assessment of Blinding.
Patients will be instructed to complete the following self-administered questionnaires. It takes about 30 minutes to complete the assessments. See Section 15.2 for instructions for administration of the questionnaires.

- **S1200** Brief Pain Inventory-Short Form (BPI-SF)
- **S1200** Western Ontario and McMaster Universities Osteoarthritis (WOMAC) Index (Version 3.1)
- **S1200** Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands (M-SACRAH)
- **S1200** PROMIS Pain Impact-Short Form (PROMIS PI-SF)
- **S1200** FACT-ES Outcome Trial Outcome Index (Version 4)
- **S1200** Aromatase Inhibitor Usage Form

The nurse or CRA will also complete the **S1200** Cover Sheet for Patient-Completed Questionnaires.

Functional testing (grip strength, ‘Timed Get Up and Go’) will be conducted and recorded on the **S1200** Functional testing. See Section 7.5 for instructions for functional testing.

Additionally, blood will be drawn for serum hormones (estradiol, FSH, LH) and inflammatory markers (TNFα, IL-6, IL-12, CRP) (see Section 15.2). Urine will be collected for urine CTX-II.

b. Follow-up visits after the 12-week study intervention will occur at 24 and 52 weeks after registration.

Patients will be instructed to complete the same self-administered questionnaires as listed in Section 7.3a.

The nurse or CRA will also complete the **S1200** Cover Sheet for Patient-Completed Questionnaires.

Functional testing (grip strength, ‘Timed Get Up and Go’) will be conducted and recorded on the **S1200** Functional testing. See Section 7.5 for instructions for functional testing.

At 24 and 52 weeks, use of pain medications will be recorded on the **S1200** Supplemental Agents Reporting Form. To assess the adequacy of blinding, patients in the true and sham acupuncture groups will give an opinion about their treatment assignment and all three groups will receive a voucher for 10 free true acupuncture sessions to be administered by the study acupuncturist(s) and used off-study at her convenience. Information on additional acupuncture received off-study will be collected. All of this data will also be recorded on the **S1200** Assessment of Blinding. Use of the 10 free true acupuncture sessions will be reported using the Columbia University CRF until the patient reaches their 52 week visit.

Additionally, at 24 weeks, blood will be drawn for serum hormones (estradiol, FSH, LH) and inflammatory markers (TNFα, IL-6, IL-12, CRP) and urine will be collected for CTX-II. At 24 and 52 weeks, urine for AI metabolites will be collected (see Section 15.1).
7.4 Telephone Contact

During telephone contact by the CRA at 2 and 4 weeks, acupuncture adherence will be ascertained and documented on the S1200 Adherence to Acupuncture Form for those patients assigned to receive acupuncture treatment. Adverse events will be assessed and recorded on the S1200 Adverse Event Summary Form. Use of pain medications will be recorded on the S1200 Supplemental Agents Reporting Form.

During telephone contact by the CRA at 16 and 20 weeks, the BPI-SF will be administered and documented on the S1200 Brief Pain Inventory-Short Form (BPI-SF). Use of pain medications will be recorded on the S1200 Supplemental Agents Reporting Form. The nurse or CRA will also complete the S1200 Cover Sheet for Patient-Completed Questionnaires.

7.5 Functional Testing

At baseline, Week 6, Week 12, Week 24, and Week 52, functional testing of the hands and legs will be conducted. Grip strength will be measured with a FEI Baseline Digital Smedley Spring Dynamometer (200 pound). The dominant hand will be recorded and both hands will be tested. Patients will be asked to make 3 maximal voluntary contractions (MVC), with 1 minute between each. Verbal encouragement will be given and the read out of each contraction, in kg, will allow visual feedback. Only the maximum contraction will then be taken to represent the MVC strength.

The “Timed Get Up and Go” (TGUG) test is a physical function assessment tool that measures balance and gait. The test uses a straight-backed chair with arms and a seat height of 42 cm. If there is not a seat that has a height of 42 cm, any seat with a height ranging from 40-50 cm can be used. Lower extremity strength is measured when the patient is asked to stand up from the chair without using her arms, walk 3 meters (10 feet), turn around, walk back to the chair, and be seated. The subject is timed with a stopwatch from the command ‘go’ until she returns to a seated position in the chair. Many recently developed measures of physical activity use timing of tasks to improve objectivity of the scoring. Timing of task performance has been found to provide greater discrimination of differences in ability, especially in elderly subjects with higher functional abilities.

7.6 Acupuncture Treatment

a. Acupuncture Schedule

Patients will be randomized (2:1:1) to true acupuncture vs. sham acupuncture vs. waitlist control. Patients will be blinded to true acupuncture vs. sham acupuncture. Patients will not be blinded to the waitlist control. The study intervention is to begin within 5 working days of registration.

- True acupuncture administered twice weekly for 6 weeks followed by weekly treatments for 6 weeks (12-18 sessions total over 12 weeks).
- Sham acupuncture administered twice weekly for 6 weeks followed by weekly treatments for 6 weeks (12-18 sessions total over 12 weeks).
- Duration of intervention – Study intervention will be continued for 12 weeks unless unacceptable toxicity occurs, the subject has concurrent illness that necessitates withdrawal or subject decides to withdraw from participation for any reason. If the subject’s medical condition is influenced by the knowledge of which study intervention she was assigned, there will be unblinding upon the Investigator’s request.
• A licensed Study Acupuncturist(s) designated at each site will administer the true and sham acupuncture treatments.
• After the 24-week visit, all patients will receive a voucher for 10 free true acupuncture sessions administered by the Study Acupuncturist(s) and to be used at the patient’s convenience and documented in the web-based calendar until the patient reaches their 52 week visit.

During the first 6 weeks of acupuncture, the twice weekly acupuncture sessions should be at least 24 hours apart and if a session is missed, it may be made up in the following weeks for up to 3 sessions per week. The schedule will be flexible to allow for 8-12 sessions during the first 6 weeks, which may be more reflective of what may happen in the “real world” setting. During the second 6 weeks of weekly maintenance acupuncture, there will be no make-up of missed sessions, so patients may have only 1 session per week during the second 6 weeks of the study intervention (Week 7 to 12).

b. Documentation of Acupuncture Visits

The Study Acupuncturist(s) must maintain a careful record of the acupuncture visits and the true or sham point prescription administered at each visit, which will be captured on the proper electronic case report forms (e-CRFs). To maintain blinding, the data on the true or sham point prescription administered at each visit will be entered into a separate locked database, which will only be accessible to the Study Acupuncturist(s) and an unblinded Site Administrator at CUMC who will not have contact with patients.

7.7 Criteria for Removal from Protocol Intervention*

a. Evidence of new cancer or cancer recurrence at any time.

b. Unacceptable toxicity (see Section 8.1).

c. Delay of 30 consecutive days due to any reason after study registration.

d. The patient may withdraw from the study at any time for any reason.

e. Completion of 12 weeks of study intervention (acupuncture). (NOTE: Patients randomized to Waitlist Control are considered off protocol intervention after 12 weeks.)

* Under intent-to-treat, all follow-up assessments will continue, if possible, whether the patient completes intervention or has early removal from intervention.

7.8 Additional Instructions for Patient-Completed Questionnaires

If a patient goes off protocol intervention before the schedule for patient-completed questionnaires has been completed, please administer all patient-completed questionnaires at the scheduled assessment times, even if the patient has begun another treatment. The 6, 12, 24, and 52 week assessment times as well should be defined from registration.

The **S1200** Cover Sheet for Patient-Completed Questionnaires is required with each set of patient-completed questionnaires indicating whether or not the assessment occurred, if assistance was required, and the location of the assessment. If one or all of the questionnaires were not administered, an overall reason must be indicated on the **S1200** Cover Sheet for Patient-Completed Questionnaires. Please see Section 15.2 for more detailed instructions for the questionnaires.
7.9 Discontinuation of Treatment

All reasons for study discontinuation must be documented on the S1200 Off Protocol Notice.

7.10 Follow-Up Period

All patients will be followed for one year from registration. No further follow-up will be required once the patient completes the 52-week visit.

8.0 SAFETY MONITORING

8.1 NCI Common Terminology Criteria for Adverse Events

This study will utilize the CTCAE (NCI Common Terminology Criteria for Adverse Events) Version 4.0 for toxicity and Reportable Adverse Event reporting. A copy of the CTCAE Version 4.0 can be downloaded from the CTEP home page (http://ctep.cancer.gov). All appropriate treatment areas should have access to a copy of the CTCAE Version 4.0.

8.2 Acupuncture Schedule Adjustments

The main toxic effects of acupuncture include a slightly increased risk of minor bruising and bleeding and rarely infection. Needle shock, a vaso-vagal response to needle insertion, occurs about 5% of the time usually during the first or second treatment. (147) Patients may become pre-syncopal, diaphoretic, nauseated and occasionally lose consciousness. These symptoms are easily reversible by removing the acupuncture needles and laying the patient supine. There are no universal guidelines in patients with lymphoedema however there may be hospital, academic medical center, or department specific guidelines. If these exist, the administration of unilateral true acupuncture or sham acupuncture points can be tracked on the S1200 Acupuncture Visit Form. Toxicity will be assessed during telephone contact at 2 and 4 weeks and during study visits at 6 and 12 weeks.

8.3 Reporting adverse events and referrals for patient care

During the study, all study patients will be under the care of either a medical oncologist or a surgical oncologist. If any abnormalities or problems, including possible adverse events (AE) or reportable adverse events, are detected during screening, subsequent measurements, or study activities, the patient will be referred back to their medical provider for appropriate care. The site CRAs and site investigators will be responsible for recording and reporting any AEs and reportable adverse events. The possible routes of referral are as follows:

a. Abnormality/problem detected during screening and subsequent clinic visits: The CRA will notify and discuss with the responsible site investigator. The patient will be referred to their medical provider by the CRA and the responsible site investigator as appropriate. The CRA and responsible site investigator will record and report AEs and reportable adverse events as appropriate.

b. Abnormality/problem detected during telephone contact: The CRA will refer the patient to their medical provider and the responsible site investigator as appropriate. The CRA and responsible site investigator will record and report AEs and reportable adverse events as appropriate.
c. **Abnormality/problem detected during acupuncture session:** The Study Acupuncturist will notify and discuss with the responsible site investigator. The CRA will be notified and the patient will be referred to their medical provider by the CRA and the responsible site investigator as appropriate. The CRA and responsible site investigator will record and report AEs and reportable adverse events as appropriate.

### 8.4 Dose Modification Contacts

For study related questions, please contact Dr. Dawn Hershman at 212/305-1495 or Dr. Katherine Crew at 212/305-1732.

### 8.5 Adverse Event Reporting

Toxicities (including suspected reactions) that meet the expedited reporting criteria as outlined in Section 16.0 of the protocol must be reported to the Operations Office, Study Chair and NCI via CTEP-AERS, and to the IRB per local IRB requirements.
## STUDY CALENDAR

### REQUIRED STUDIES

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<tr>
<th>PRE STUDY</th>
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### LABORATORY

| FSH (if necessary) | X | | | | | | | | | | | | | |
| ER/PgR | X | | | | | | | | | | | | | |

### QUESTIONNAIRES

| S1200 Cover Sheet for Patient-Completed Questionnaires | X | X | | | | | | | | | | | | |
| S1200 BPI-SF | X | | X | X | X | X | X | X | X | X | X | X | X | X | X |
| S1200 WOMAC | X | | X | X | X | | | | | | | | | | |
| S1200 M-SACRAH | X | | X | X | X | | | | | | | | | | |
| S1200 PROMIS PI-SF | X | | X | X | X | | | | | | | | | | |
| S1200 FACT-ES | X | | X | X | X | | | | | | | | | | |
| S1200 AI Usage Form | X | | X | X | X | | | | | | | | | | |
| S1200 Assessment of Blinding | X | | X | X | X | | | | | | | | | | |

Calendar continued on next page. Click here for footnotes.
### PROCEDURES

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### TREATMENT

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**NOTE:** Forms are found on the protocol abstract page on the SWOG website (www.swog.org). Forms submission guidelines are found in Section 14.0. 

√ Blood will be collected to measure serum hormones (estadiol, FSH, LH) and inflammatory biomarkers (TNF, IL-6, IL-12, CRP) at baseline, Weeks 6, 12, and 24. Plasma and buffy coat will be collected at baseline only for DNA extraction and genotyping (CYP19A1). 

β See Section 15.1 and order kit from SWOG Specimen Repository. 

α If the study visit occurs on the same day as an acupuncture visit, then the evaluations should be conducted prior to acupuncture treatment. Study visit to include assessment of adherence to acupuncture use of pain medication, adverse events, and adequacy of blinding. 

Σ +/- 7 days to allow flexibility in scheduling of assessments.  

# +/- 10 days to allow flexibility in scheduling of assessments. The Week 6 and Week 12 study visits are preferably performed after acupuncture Session 12 and Session 18, respectively.  

^ +/- 14 days to allow flexibility in scheduling of assessments.  

π After the Week 24 visit, all patients will receive a voucher for 10 free true acupuncture sessions to be administered by the study acupuncturist(s) and used off-study at her convenience. 

* True/Sham acupuncture will be administered twice weekly for 6 weeks (12 sessions) followed by 6 weekly treatments (6 sessions) of maintenance weeks (18 sessions total over 12 weeks). 

% Patients on the Waitlist Control arm will not receive any acupuncture (true or sham) until given the vouchers at Week 24. All forms must be submitted as required in Section 14.0.  

† Telephone contact at Weeks 2 and 4 will include assessment at adherence to acupuncture, use of pain medications, and adverse events.  

~ Telephone contact at Weeks 16 and 20 will include phone administration of the S1200 Brief Pain Inventory - Short Form and assessment of use of pain medications.  

¥ Pre-study and study examinations can be conducted by a licensed healthcare professional including but not limited to MD, NP, PA, RN. 

◇ May be performed within 180 days prior to registration.
10.0 CRITERIA FOR EVALUATION AND ENDPOINT DEFINITIONS

10.1 Primary Endpoint

Reduction in worst joint pain at 6 weeks between the true acupuncture compared to sham acupuncture and waitlist control groups: A difference of two points in the modified Brief Pain Inventory worst pain score (Item #2) has been identified as a clinically meaningful difference. (148) This item has a scale of 0 to 10 with 0 indicating “No pain” and 10 indicating “Pain as bad as you can imagine”.

10.2 Secondary Endpoints

To investigate the effects of true acupuncture compared to sham acupuncture and waitlist control on the following outcome measures compared to baseline:

a. Brief Pain Inventory-Short Form (BPI-SF) worst pain, worst stiffness (items 15 & 16), pain severity, and pain-related interference scores at 6, 12, 16, 20, 24, and 52 weeks.

b. Western Ontario and McMaster Universities Osteoarthritis (WOMAC) index (pain, stiffness, and function) for the hips and knees at 6, 12, 24, and 52 weeks.

c. Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands (M-SACRAH) (pain, stiffness, and function) at 6, 12, 24, and 52 weeks.

d. PROMIS Pain Impact-Short Form (PROMIS PI-SF) at 6, 12, 24, and 52 weeks.

e. Quality of life as assessed by the FACT-ES Trial Outcome Index (Version 4) at 6, 12, 24, and 52 weeks.

f. Functional testing of the hands (grip strength) and legs (‘Timed Get Up and Go’) at 6, 12, 24, and 52 weeks.

g. Analgesic and opioid use at 2, 4, 6, 12, 16, 20, 24, and 52 weeks.

h. AI adherence at 12, 24, and 52 weeks.

i. Urine Al metabolites at 24 and 52 weeks.

j. Serum hormone biomarkers (estradiol, FSH, LH) and inflammatory biomarkers (serum TNFα, IL-6, IL-12, CRP; urine CTX-II) at 6, 12, and 24 weeks.


l. Safety and tolerability of acupuncture.

This study will also describe adverse events, assess the association between CYP19A1 genotype and severity of AI-related joint symptoms, and assess the adequacy of blinding and use of additional off-study acupuncture treatments.
10.3 Performance Status

Patients will be graded according to the Zubrod performance status scale.

<table>
<thead>
<tr>
<th>POINT</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Fully active, able to carry on all pre-disease performance without restriction.</td>
</tr>
<tr>
<td>1</td>
<td>Restricted in physically strenuous activity but ambulatory and able to carry out work of a light or sedentary nature, e.g., light housework, office work.</td>
</tr>
<tr>
<td>2</td>
<td>Ambulatory and capable of self-care but unable to carry out any work activities; up and about more than 50% of waking hours.</td>
</tr>
<tr>
<td>3</td>
<td>Capable of limited self-care, confined to bed or chair more than 50% of waking hours.</td>
</tr>
<tr>
<td>4</td>
<td>Completely disabled; cannot carry on any self-care; totally confined to bed or chair.</td>
</tr>
</tbody>
</table>

11.0 STATISTICAL CONSIDERATIONS

11.1 Primary Hypothesis

The primary hypothesis of the study is that true acupuncture will decrease joint pain associated with the use of AIs in breast cancer patients compared to sham acupuncture or waitlist control at 6 weeks. Joint pain will be assessed using “worst pain” according to the BPI. Enrollees must currently be taking AIs and must exhibit joint pain with a minimum BPI worst pain score of 3; scores of 3bb to 10 are considered to reflect moderate to severe pain. (149) Patients will be stratified at randomization by study site to account for potential site-specific variation in the administration of acupuncture. The primary endpoint of 6 weeks was chosen in order to confirm initial results of a 6-week acupuncture intervention given twice weekly in a larger multicenter study. (80)

The study team stipulates an alpha=.025 two-sided test (to account for two comparisons; i.e., true acupuncture vs. sham acupuncture and true acupuncture vs. waitlist control), with an estimated 5% non-adherence (reducing the nominal effect size) and 10% dropout rate (increasing the total required sample size) at the primary endpoint evaluation time of 6 weeks after randomization. In addition, the design will incorporate a 10% contamination rate (which also reduces the nominal effect size) based on the assumption that a portion of the patients will have joint pain with a different etiology (that is, not biologically associated with AI treatment and therefore not amenable to the specified intervention). The subjects will be randomized 2:1:1 to: 1) true acupuncture, 2) sham acupuncture, or 3) waitlist control. Power for the design will be a function of the difference detected and the standard deviation (SD) at 6 weeks after randomization. Data on the use of the BPI to assess joint pain from AIs is limited. A difference of 2 points in the BPI worst pain score has been identified as a clinically meaningful difference. (148) In the pilot single institution randomized sham acupuncture-controlled trial for AI-induced joint pain, the standard deviation for BPI “worst pain” at 6 weeks on the control arm was 2.31. (88) For design purposes for this multicenter acupuncture study, a more conservative estimate of standard deviation of 3.0 points will be assumed. Given the number controls in the prior study was small (n=18), assumption of a standard deviation (SD) of 3.0 points for this study is likely conservative.
For a 2-point difference and a 3.0 point SD at 6 weeks, with other parameters as specified above, 208 total eligible patients (104 true acupuncture, 52 sham acupuncture, and 52 waitlist control) would be required for 82% power for the two comparisons, based on two separate t-tests. [Note that in power calculations, the combination of the 5% non-adherence rate and the 10% contamination rate, stipulated above, reduces the nominal effect size of a 2 point difference to 1.7 points; while the 10% dropout rate inflates the estimated sample size by a factor of 1.9 or about 11%.] This approach allows two fully-powered independent comparisons between true acupuncture and the two alternatives. In addition, unlike an ANOVA approach, it does not assume equal variances across the three arms, which may be more appropriate in this setting given the inclusion of a waitlist control. Power will be higher with lower observed standard deviation.

A multiple linear regression analysis of the primary endpoint will include the baseline score as well as the pre-specified stratification factor. To allow for an ineligibility of 8% (based on a recent study, S0715), a total of 228 patients will be enrolled to achieve 208 eligible patients.

The potential for different dropout by arm, especially with respect to the waitlist control, will be mitigated by monthly conferences among study investigators to encourage proper assessment and submission of forms at every required timepoint for all patients. Dropout patterns will be monitored on an ongoing basis.

11.2 Secondary endpoints

a. Secondary hypotheses (Section 10.2) will include assessment of intervention differences at each timepoint individually (weeks 6, 12, 16, 20, 24, and 52), as well as longitudinal modeling of the outcome measures over time. For longitudinal modeling, linear mixed models will be used; if there is evidence of non-random dropout, pattern-mixture models will be utilized as sensitivity analysis. Covariates for longitudinal modeling will include intervention assignment, assessment time, their interaction, baseline score, and other potential confounding variables (i.e. age).

b. Exploratory analyses will also be done to evaluate the relationship between aromatase gene polymorphisms (CYP19A1) and inflammatory biomarkers with severity of joint symptoms and response to acupuncture treatment.

c. A further exploratory analysis at both 6 and 12 weeks according to adherence to intervention assignment will be conducted, with receipt of ≥ 80% of planned treatment sessions defined as “adherent.” (In particular, patients will be divided into the adherent true (≥ 80%), non-adherent true (< 80%), adherent sham (≥ 80%), and non-adherent sham (< 80%) acupuncture groups.) The interaction between treatment adherence category and treatment category will be tested. Adherence to acupuncture sessions as a continuous measure will also be evaluated.

d. The success of blinding will be evaluated by using chi-square tests to compare the percentage of patients in each group who believed that they had received true acupuncture at Weeks 6, 12, 24, and 52.

11.3 Sample Size/Accrual Rate

Based upon a previous cross-sectional survey at Columbia University, 253 consecutive breast cancer patients on adjuvant AI therapy were screened over a 4-month period; 47% reported joint pain which started or worsened after initiation of AI therapy and among those with AI-related arthralgias, about two-thirds reported a worst pain score ≥ 5 points on a 0-10 scale. Therefore, it is anticipated that up to 20 women screened in the clinic
each month may be eligible for this trial. In addition, the Research Recruitment and Minority Outreach Core at Columbia University will be utilized to target women with breast cancer in the community. Based on this strategy, investigators were able to successfully recruit to other intervention studies for AI-related arthralgias, including acupuncture and supplements such as glucosamine/chondroitin and omega-3 fatty acids. Investigators will also advertise through breast cancer support groups and reach out to colleagues in the community who see breast cancer patients.

Accrual is expected to be rapid given the common use of AIs for adjuvant therapy in postmenopausal women with breast cancer. Allowing for 6 months ramp-up and IRB approval time, accrual of 1.5 patients per month for each site would allow completion of the study accrual in approximately 2.5 years. Accrual will be assessed every 6 months after study activation. If monthly average accrual is < 50% 1 year after study activation revision will be considered. For an individual site, if there is no accrual in a 6 month period, the study team will evaluate barriers to accrual. If these changes do not improve accrual, the site will be closed.

11.4 Data and Safety Monitoring Committee

A Data and Safety Monitoring Committee will oversee the conduct of the study. The Committee consists of four members from outside of SWOG, 3 SWOG members, 3 non-voting representatives from the National Cancer Institute (NCI), and the Group Statistician (non-voting). The members of this Committee will receive confidential reports every 6 months from the SWOG Statistical Center, and will meet at the Group’s bi-annual meetings as necessary. The Committee will be responsible for decisions regarding possible termination and/or early reporting of the study.

12.0 DISCIPLINE REVIEW

There will be no formal discipline review for this study.

13.0 REGISTRATION GUIDELINES

13.1 Registration Timing

Patients must be registered prior to initiation of study intervention (no more than 5 working days prior to planned start of study intervention).

13.2 Investigator/Site Registration

The individual registering the patient must have completed the appropriate SWOG Registration Worksheet. The completed form must be referred to during the registration but should not be submitted as part of the patient data. OPEN will also ask additional questions that are not present on the SWOG Registration Worksheet. The individual registering the patient must be prepared to provide answers to the following questions:

- Institution CTEP ID
- Protocol Number
- Registration Step
- Treating Investigator
e. Credit Investigator
f. Patient Initials
g. Patient’s Date of Birth
h. Patient SSN (SSN is desired, but optional. Do not enter invalid numbers.)
i. Country of Residence
j. ZIP Code
k. Gender (select one):
   - Female Gender
   - Male Gender
l. Ethnicity (select one):
   - Hispanic or Latino
   - Not Hispanic or Latino
   - Unknown
m. Method of Payment (select one):
   - Private Insurance
   - Medicare
   - Medicare and Private Insurance
   - Medicaid
   - Medicaid and Medicare
   - Military or Veterans Sponsored NOS
   - Military Sponsored (Including Champus & Tricare)
   - Veterans Sponsored
   - Self Pay (No Insurance)
   - No Means of Payment (No Insurance)
   - Other
   - Unknown
n. Race (select all that apply):
   - American Indian or Alaska Native
   - Asian
   - Black or African American
   - Native Hawaiian or other Pacific Islander
   - White
   - Unknown

13.3 Registration procedures

a. All site staff will use OPEN to enroll patients to this study. OPEN is a web-based application and can be accessed at https://open.ctsu.org, or from the OPEN tab on the CTSU members’ side of the website at https://www.ctsu.org, or from the OPEN Patient Registration link on the SWOG CRA Workbench.

b. Prior to accessing OPEN site staff should verify the following:
   - All eligibility criteria have been met within the protocol stated timeframes and the affirmation of eligibility on the Registration Worksheet has been signed by the registering investigator or another investigator designate. Site staff should refer to Section 5.0 to verify eligibility.
   - All patients have signed an appropriate consent form and HIPAA authorization form (if applicable).
c. Access requirements for OPEN:

- Site staff will need to be registered with CTEP and have a valid and active CTEP-IAM account. This is the same account (user ID and password) used for the CTSU members' web site.
- To perform registrations on SWOG protocols you must have an equivalent 'Registrar' role on the SWOG roster. Role assignments are handled through SWOG.

Note: The OPEN system will provide the site with a printable confirmation of registration and treatment information. Please print this confirmation for your records.

d. Further instructional information is provided on the OPEN tab on the CTSU members’ side of the website at https://www.ctsu.org or at https://open.ctsu.org. For any additional questions contact the CTSU Help Desk at 1-888-823-5923 or ctsucontact@westat.com.

13.4 Exceptions to SWOG registration policies will not be permitted

a. Patients must meet all eligibility requirements.

b. Institutions must be identified as approved for registration.

c. Registrations may not be cancelled.

d. Late registrations (after initiation of treatment) will not be accepted.

14.0 DATA SUBMISSION SCHEDULE

14.1 Data Submission Requirement

Data must be submitted according to the protocol requirements for ALL patients registered, whether or not assigned treatment is administered, including patients deemed to be ineligible. Patients for whom documentation is inadequate to determine eligibility will generally be deemed ineligible.

14.2 Master Forms

Master forms can be found on the protocol abstract page on the SWOG website (www.swog.org) and (with the exception of the sample consent form and the Registration Worksheet) must be submitted on-line via the Web; see Section 14.3a for details.

14.3 Data Submission Procedures

a. SWOG institutions must submit data electronically via the Web by using the SWOG CRA Workbench. To access the CRA Workbench, go to the SWOG Web site (http://swog.org) and logon to the Members Area. After you have logged on, click on the CRA Workbench link to access the home page for CRA Workbench website. Next, click on the Data Submission link and follow the instructions. For new users, the link to a "Starter Kit" of help files may be found by clicking on the Starter Kit link at the Members' logon page.
To submit data via the web the following must be done (in order):

1. You are entered into the SWOG Roster and issued a SWOG Roster ID Number,
2. You are associated as an investigator or CRA/RN at the institution where
   the patient is being treated or followed, and
3. Your Web User Administrator has added you as a web user and has given
   you the appropriate system permissions to submit data for that institution.

For assistance with points 1 and 2 call the Operations Office at 210/614-8808. For
point 3, contact your local Web User Administrator (refer to the "Who is my Web
User Administrator?" function on the swog.org Members logon page). For other
difficulties with the CRA Workbench, please email technicalquestion@crab.org.

b. If you need to submit data that are not available for online data submission, the
   only alternative is via facsimile. Should the need for this occur, institutions may
   submit data via facsimile to 800/892-4007 or 206/342-1680 locally. Please do not
   use cover sheet for faxed data. Please make sure that each page of all faxed
   data include the SWOG patient number, study ID and patient initials.

14.4 Data Submission Overview and Timepoints

a. AFTER REGISTRATION BUT PRIOR TO BEGINNING STUDY INTERVENTION
   (WITHIN 24 HOURS AFTER OBTAINING SPECIMENS):

Submit baseline blood and samples as described in Section 15.0. The SWOG
Specimen Tracking System must be used for sample submission.

b. WITHIN 7 DAYS AFTER REGISTRATION:

Submit the following:

S1200 Prestudy Form
S1200 Functional Testing
S1200 Supplemental Agents Reporting Form

Institutional surgical pathology report to confirm staging.

S1200 Cover Sheet for Patient-Completed Questionnaires
S1200 Brief Pain Inventory-Short Form (BPI-SF)
S1200 Western Ontario and McMaster Universities Osteoarthritis (WOMAC)
   Index (Version 3.1)
S1200 Modified-Score for the Assessment and Quantification of Chronic
   Rheumatoid Affections of the Hands (M-SACRAH)
S1200 PROMIS Pain Impact-Short Form (PROMIS PI-SF)
S1200 FACT-ES Trial Outcome Index (Version 4)
S1200 Aromatase Inhibitor Usage Form
c. **WITHIN 14 DAYS OF THE WEEK 2 AND 4 STUDY ASSESSMENTS:**

Submit the following:
- **S1200** Adherence to Acupuncture Form
- **S1200** Adverse Event Summary Form
- **S1200** Supplemental Agents Reporting Form

d. **WITHIN 14 DAYS OF THE WEEK 6, 12, 24 AND 52 STUDY ASSESSMENTS:**

Submit the following:
- **S1200** Cover Sheet for Patient-Completed Questionnaires
- **S1200** Brief Pain Inventory-Short Form (BPI-SF)
- **S1200** Western Ontario and McMaster Universities Osteoarthritis (WOMAC) Index (Version 3.1)
- **S1200** Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands (M-SACRAH)
- **S1200** PROMIS Pain Impact-Short Form (PROMIS PI-SF)
- **S1200** FACT-ES Trial Outcome Index (Version 4)
- **S1200** Aromatase Inhibitor Usage Form
- **S1200** Functional Testing
- **S1200** Adherence to Acupuncture Form [WEEK 6 AND 12]
- **S1200** Adverse Event Summary Form [WEEK 6 AND 12]
- **S1200** Supplemental Agents Reporting Form
- **S1200** Assessment of Blinding

e. **WITHIN 24 HOURS OF OBTAINING SPECIMENS AT THE WEEK 6, 12, 24 AND 52 STUDY ASSESSMENTS:**

Submit blood and urine samples as described in Section 15.0.

f. **WITHIN 14 DAYS OF THE WEEK 16 AND 20 STUDY ASSESSMENTS:**

Submit the following:
- **S1200** Brief Pain Inventory-Short Form (BPI-SF)
- **S1200** Supplemental Agents Reporting Form
- **S1200** Cover Sheet for Patient-Completed Questionnaires
g. **WITHIN 14 DAYS OF REMOVAL FROM PROTOCOL TREATMENT:**

Submit a copy of the Off Protocol Notice along with the final copy of the S1200 Adverse Event Summary Form.

h. **WITHIN 14 DAYS OF A DIAGNOSIS OF NEW OR RECURRENT CANCER OCCURRING UP TO 52 WEEKS AFTER REGISTRATION:**

Submit copies of:

- Pathology report documenting cancer
- Follow-Up Form

i. **WITHIN 4 WEEKS OF KNOWLEDGE OF DEATH OCCURRING UP TO 52 WEEKS AFTER REGISTRATION:**

Submit the Notice of Death documenting death information.

### 15.0 SPECIAL INSTRUCTIONS

#### 15.1 Collection and Submission of Blood and Urine Specimens

Patients are required to submit blood samples for serum hormones (estradiol, FSH, LH), serum inflammatory markers (TNFα, IL-6, IL-12, CRP), blood for DNA analysis of CYP19A1, urine for CTX-II, and urine for AI metabolites as listed below.

<table>
<thead>
<tr>
<th>Specimen Type</th>
<th>Baseline</th>
<th>Week 6</th>
<th>Week 12</th>
<th>Week 24</th>
<th>Week 52</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood for Serum Hormones</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Blood for Inflammatory Markers</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Blood for DNA</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urine for CTX-II</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Urine for AI metabolites</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

If the patient agrees, left over blood and urine will be banked for future translational medicine studies. **Institutions are required to seek additional patient consent to bank blood and urine for future translational medicine studies.**

a. Specimen collection and submission instructions can be accessed on the SWOG Specimen Submission webpage (https://swog.org/Members/ClinicalTrials/Specimens/STSpecimens.asp), or via the link on the S1200 protocol abstract page on the SWOG website (www.swog.org). Additional collection instructions are located in Section 15.1c.

b. Specimen collection kits may be ordered by the SWOG Specimen Repository Management Application at http://ricapps.nationwidechildrens.org/BPCKitManagement/.
c. Additional Collection Instructions

1. Blood for serum hormones (estradiol, FSH, LDH) and inflammatory markers (TNFα, IL-6, IL-12, CRP) will be collected in two 10 mL red top tubes. Store samples at 4°C for up to 24 hours prior to shipping.

2. Blood for DNA (CYP19A1) will be collected in a 10 mL purple top EDTA tube. Store sample at 4°C for up to 24 hours prior to shipping.

3. Untreated urine will be collected in a 15 mL tube and stored at 4°C for up to 24 hours prior to shipping.

4. Blood samples collected on Friday should be shipped on the same day to the repository. Urine samples collected on Friday should be held over the weekend at 4°C and shipped on Monday.

15.2 Patient Questionnaires: Instructions for Administration

a. Time frame for questionnaires

It is important to note that the time frame for providing ratings differs depending on the scale the patient is completing. The **S1200 Brief Pain Inventory Short Form** (BPI-SF), **S1200 Western Ontario and McMaster Universities Osteoarthritis (WOMAC) Index**, **S1200 Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands** (M-SACRAH), **S1200 PROMIS Pain Impact-Short Form** (PROMIS PI-SF), and **S1200 FACT-ES Trial Outcome Index** (Version 4) should be rated with respect to the past 7 days. (See Section 14.0 for additional questionnaire submission requirements.)

If treatment is delayed due to toxicity, the assessment schedule for the Week 2, 4, 6, 12, 24 and 52 assessments should be defined from the date of registration.

b. Administration of Questionnaires

1. The first time the patient completes the questionnaires: Please read to the patient the instructions attached to each patient questionnaire. Explain the specific administration times for this protocol. Patients should be directed to report all symptoms and limitations whether or not they are related to the cancer or its treatment.

2. It is permissible to assist patients with completing the questionnaires being careful not to influence the patient's response. Note on the cover sheet what assistance was required and indicate reason (e.g., elderly, too sick, etc.). Discourage family members from: 1) being present while the patient completes the questionnaire and/or 2) influencing patient responses to the questions.

3. It is very important to review the questionnaires after the patient has completed them to be sure all of the questions have been answered and that only one answer is marked. a) If the patient has marked more than one answer per question, ask the patient which answer reflects how she is feeling. b) If the patient has skipped a question, tell the patient that a question was not answered and ask if she would like to answer the question. Always give the patient the option to refuse. Indicate on the form by the question that the patient did not want to answer this question.

4. If a patient refuses or cannot complete the questionnaire for some reason, then this must be documented on the cover sheet and faxed to the Data Operations Center in Seattle (see Section 14.3b). This form will also be available for online submission.
5. If a patient misses an appointment or is too sick to complete the questionnaires on the scheduled date, the questionnaire can be mailed to the patient or sent home with her. A telephone interview must be scheduled and completed within one week of the originally scheduled time. Patient responses to questionnaire items are to be obtained during the telephone interview while the patient is looking at her copy of the questionnaire. The date of the telephone interview is to be noted on the cover sheet. In this study, the patient will also complete scheduled telephone assessments at Weeks 16 and 20 only. As noted above, ask the patient to have a copy of the BPI in front of her and then read each item and its possible response options. Do not discuss the patient's pain or help her perception of the pain for the study questions. Reinforce the idea that only the patient perception of pain is relevant for the study outcomes.

c. Additional quality control procedures:

1. When a patient is registered on S1200, a calendar should be made with dates of upcoming patient-completed questionnaires noted. A copy of this calendar can be given to the patient with the notation that the questionnaires should be completed before receiving treatment. You may wish to photocopy the Study Calendar, Section 9.0, and include the patient's name and specific dates. A copy of this should be kept in the patient file.

2. If a patient goes off study intervention prior to the protocol-defined end of intervention at 12 weeks, administer the patient-completed questionnaires according to the protocol-defined assessment schedule (timed from registration date).

3. If a patient refuses or cannot complete the patient questionnaires at one time point, she should be asked to do so at the next scheduled administration time.

4. Anyone involved in the collection of quality of life data in SWOG trials should review the training program available on the SWOG website (www.swog.org). Please log on as a member and go to the CRA Workbench. Inside the Workbench, click on Training and select the training program: Patient-Reported Outcome Questionnaires Training Module. This program is a narrated set of slides designed to standardize the way quality of life data is collected from patients; it takes the place of the Quality of Life Training Video. Questions regarding the quality of life assessments can be addressed to Dr. Carol Moinpour at the Fred Hutchinson Cancer Research Center (206/677-4604).

d. S1200 Cover Sheet for Patient-Completed Questionnaires

For each time point, the nurse or CRA completes the S1200 Cover Sheet for Patient-Completed Questionnaires. The Cover Sheet is submitted with the set of patient-completed forms at each scheduled assessment. The Cover Sheet is very important for tracking how and when the patient forms were completed. When a patient-completed form is not administered at a scheduled time point, it is important to know why the assessment did not occur; the form includes potential reasons for a patient not completing a form. See Section 14.0 for data submission guidelines.
15.3 Acupuncture Quality Control: Instructions to Acupuncturist

An acupuncturist from each site will be required to attend an initiation training session at the SWOG group meeting, watch the online training module, read and follow the training manual and train the other acupuncturists at the site. The other acupuncturists will be required to watch the online training module and follow the training manual. During the initiation, the Columbia training acupuncturist will observe each site acupuncturist perform true and sham acupuncture on a volunteer. All acupuncturists will undergo quality assurance checks throughout the study period.

a. The first time the patient arrives for acupuncture: The acupuncturist will read to the patient the instructions. Explain the specific administration times for this protocol. Patients should be directed to report all symptoms and limitations.

b. The acupuncturist will be the only participant on the study team that is unblinded to sham and true acupuncture assignment. The acupuncturist will log into the CRAB website to retrieve the assignment after the patient has been registered. They will maintain all blinding to both the patient and the investigators. All eCRF’s should be filled out at the time of each visit and sent electronically to the Columbia Coordinating center. An unblinded staff member at CUMC will review everything for accuracy and completeness and will be available for questions by the study acupuncturists regarding data submission of acupuncture visits: Ramona Jayasena, phone 212/304-5579 or e-mail rj2002@columbia.edu.

c. It is very important to review the CRFs after the patient has completed treatment to be sure all of the questions have been answered.

d. If a patient refuses, does not show up or cannot complete the acupuncture for some reason, then this must be documented on the cover sheet and faxed to the Data Operations Center in Seattle as well as the Columbia Coordinating Center (see Section 14.3b).

e. The acupuncturists will participate in a monthly conference call with the Columbia Coordinating Center along with the SWOG Protocol Coordinator to maintain quality control.

16.0 ETHICAL AND REGULATORY CONSIDERATIONS

The following must be observed to comply with Food and Drug Administration regulations for the conduct and monitoring of clinical investigations; they also represent sound research practice:

Informed Consent

The principles of informed consent are described by Federal Regulatory Guidelines (Federal Register Vol. 46, No. 17, January 27, 1981, part 50) and the Office for Protection from Research Risks Reports: Protection of Human Subjects (Code of Federal Regulations 45 CFR 46). They must be followed to comply with FDA regulations for the conduct and monitoring of clinical investigations.

Institutional Review

This study must be approved by an appropriate institutional review committee as defined by Federal Regulatory Guidelines (Ref. Federal Register Vol. 46, No. 17, January 27, 1981, part 56) and the Office for Protection from Research Risks Reports: Protection of Human Subjects (Code of Federal Regulations 45 CFR 46).
Monitoring

This study will be monitored by the Clinical Data Update System (CDUS) Version 3.0. Cumulative CDUS data will be submitted quarterly to CTEP by electronic means. Reports are due January 31, April 30, July 31 and October 31.

Confidentiality

Please note that the information contained in this protocol is considered confidential and should not be used or shared beyond the purposes of completing protocol requirements until or unless additional permission is obtained.

Adverse Experiences

There are no agents in this study.

16.1 Adverse Event Reporting Requirements

   a. Purpose

   Adverse event data collection and reporting, which are required as part of every clinical trial, are done to ensure the safety of patients enrolled in the studies as well as those who will enroll in future studies using similar agents. Adverse events are reported in a routine manner at scheduled times during a trial. (Directions for routine reporting are provided in Section 14.0.) Additionally, certain adverse events must be reported in an expedited manner to allow for more timely monitoring of patient safety and care. The following guidelines prescribe expedited adverse event reporting for this protocol. See also Appendix 18.1 for general and background information about expedited reporting.

   b. Reporting methods

   This study requires that expedited adverse event reporting use CTEP’s Adverse Event Reporting System (CTEP-AERS). The NCI’s guidelines for CTEP-AERS can be found at http://ctep.cancer.gov. An CTEP-AERS report must be sent to SWOG Operations Office by electronically submitting the report via the CTEP-AERS Web-based application located at http://ctep.cancer.gov.

   In the rare event when internet connectivity is disrupted a 24-hour notification is made to NCI by telephone at 301-897-7497. An electronic report MUST be submitted immediately upon re-establishment of internet connection.

   c. When to report an event in an expedited manner

   When the adverse event requires expedited reporting, submit the report within 10 calendar days of learning of the event.

   You may be asked to submit supporting clinical data to the Operations Offices in order to complete the evaluation of the event. If requested, the supporting data should be sent within 5 calendar days by fax to 210-614-0006.
d. Other recipients of adverse event reports

The Operations Office will forward reports and documentation to the appropriate regulatory agencies and drug companies as required. Adverse events determined to be reportable to the Institutional Review Board responsible for oversight of the patient must be reported according to local policy and procedures.

e. Expedited reporting for acupuncture/sham interventions

Reporting requirements for acupuncture intervention are provided in Table 16.1. If there is any question about the reportability of an adverse event or if on-line CTEP-AERS cannot be used, please telephone or email the SAE Program at the Operations Office, 210/614-8808 or adr@swog.org, before preparing the report.

Table 16.1. Expedited reporting requirements for adverse events experienced by patients who have received intervention on this study.

<table>
<thead>
<tr>
<th>Attribution</th>
<th>Grade 4</th>
<th>Grade 5a</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Unexpected</td>
<td>Expected</td>
</tr>
<tr>
<td>Unrelated or Unlikely</td>
<td>CTEP-AERS</td>
<td>CTEP-AERS</td>
</tr>
<tr>
<td>Possible, Probable, Definite</td>
<td>CTEP-AERS</td>
<td>CTEP-AERS</td>
</tr>
</tbody>
</table>

CTEP-AERS: Indicates an expedited report is to be submitted via CTEP-AERS within 10 calendar days of learning of the eventb.

a This includes all deaths within 30 days of the last acupuncture/sham intervention, regardless of attribution. Any death that occurs more than 30 days after the last acupuncture/sham intervention and is attributed (possibly, probably, or definitely) to the intervention and is not due to cancer recurrence must be reported according to the instructions above.

b Submission of the on-line CTEP-AERS report plus any necessary amendments generally completes the reporting requirements. You may, however, be asked to submit supporting clinical data to the Operations Office in order to complete the evaluation of the event. If requested, the specified data should be sent by fax to 210-614-0006.
17.0 BIBLIOGRAPHY


18.0 APPENDIX

18.1 Acupuncture Description and Side Effects
18.2 Patient-Completed Questionnaires
18.4 Measurement of Blood and Urine Biomarkers
18.5 Training of Study Acupuncturists to the Study Protocol
18.5 CRA Functional Testing Training
18.6 Emergency Unblinding Guidelines
18.1 Acupuncture Description and Side Effects

Description of True Acupuncture

The true acupuncture protocol and procedures employed will be devised with adherence to the Standards for Reporting of Controlled Trials in Acupuncture (STRICTA) recommendations. The acupuncture protocol was selected based on a standard Traditional Chinese Medicine (TCM) point prescription and informal practitioner query. A manualized protocol (i.e., a protocol that is standardized but allows the Study Acupuncturist to modify treatments based on the patient’s current joint symptoms) has been developed. The manualized protocol was selected as opposed to a fixed acupuncture point protocol to facilitate the systematic delivery of standardized acupuncture treatments and because it was shown to be effective in other acupuncture trials. (79,80,96,143,150) Patients will be scheduled for twice weekly 45-minute acupuncture sessions (30-minute acupuncture, 15-minute set-up) for 6 weeks (8-12 sessions) followed by 6 weekly treatments (4-6 sessions) for maintenance (12-18 sessions total over 12 weeks). The true acupuncture protocol includes standardized body and auricular treatments (Table 1). This point prescription is based on a standard TCM point prescription to treat “Bi Syndrome” and the National Acupuncture Detoxification Association (NADA) protocol applied to one ear to relieve pain and decrease stress. At every acupuncture visit, the patients assigned to true acupuncture will receive the full body acupuncture prescription and the auricular acupuncture-NADA protocol in one ear (to be alternated at each visit). (151) In addition, each session will include a joint-specific point prescription tailored to up to four of the patient’s most painful joints, including knees, fingers, lumbar area, shoulders, hips, and wrists. (79, 144) The anatomic site selection may vary between visits depending on the patient’s current symptoms.

Acupuncture needles used will be Mac™ single-use, sterile, and disposable. The full body acupuncture needles will be 1 inch, 1.5 inches and 34-gauge or 3 inches and 30-gauge (manufactured by Tian Jin Haing Lin Sou Won Medical Instrument Co., Ltd; distributed by Mac Co., Roslyn Heights, New York, USA) and auricular needles will be 15mm and 38-gauge (Seirin-America, Inc., MA). The needling protocol will consist of first swabbing all points with alcohol and needling auricular points, then needling full-body points. The needles will be inserted to the proper needling depth as determined by standard point locations and to elicit a de qi sensation at all standardized full body acupuncture points. (152) The needles will remain in situ for 20-25 minutes during which the Study Acupuncturist will return to stimulate the needles once, utilizing even needle technique in order to re-elicit the de qi sensation. If needed, to access joint-specific points that are not accessible in the supine position, needles may be inserted after the first set, at the proper depth and angle, eliciting a de qi sensation, and retained for 10 minutes. No electrical stimulation will be used. One licensed Study Acupuncturist will perform all treatments. In addition, at least 1 other licensed acupuncturist will be trained in administering the protocol in the event of schedule conflicts. At 6, 12, 24, and 52 weeks, to assess the adequacy of blinding, patients will give an opinion about their treatment assignment. After the 24-week visit, patients will receive a voucher for 10 free acupuncture sessions to be administered by the study acupuncturist(s) used off-study at her convenience. Any additional acupuncture treatments received off-study will be recorded at 24 and 52 weeks. All of this information will be recorded on the S1200 Assessment of Blinding Form.
### Table 1. True Acupuncture Points

<table>
<thead>
<tr>
<th>Point Name</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Full body acupuncture points (given at each visit)</strong></td>
<td></td>
</tr>
<tr>
<td>SJ 5-wai guan (TB 5)</td>
<td>Posterior to the fourth metatarsalphalangeal joint, in the depression lateral to the tendon of m. extensor digiti minimi of the foot</td>
</tr>
<tr>
<td>GB 41-zulin qi</td>
<td>In the depression anterior and inferior to the head of the fibula</td>
</tr>
<tr>
<td>GB 34-yang ling quan</td>
<td>On the dorsum of the hand, between the 1st and 2nd metacarpal bones, approximately in the middle of the 2nd metacarpal bone on the radial side</td>
</tr>
<tr>
<td>LI 4-he gu</td>
<td>On the dorsum of the foot, at the midpoint of the transverse crease of the ankle joint, in the depression between the tendons of m. extensor digitorum longus and hallucis longus, approximately at the level of the tip of the external maleolus</td>
</tr>
<tr>
<td>ST-41-jie xi</td>
<td>In the depression between the tip of the external maleolus and the anterior side of the tendon achilles</td>
</tr>
<tr>
<td>KD 3-tai xi</td>
<td></td>
</tr>
<tr>
<td><strong>Auricular acupuncture points (given in right ear on odd numbered visits and left ear on even numbered visits)</strong></td>
<td></td>
</tr>
<tr>
<td>Shen men</td>
<td>At bifurcating point between superior and inferior antihelix crux, and lateral 1/3 of the triangular fossa</td>
</tr>
<tr>
<td>Kidney</td>
<td>On the lower border of the inferior antihelix crux, directly above the small intestine point</td>
</tr>
<tr>
<td>Liver</td>
<td>At the posterior aspect of the stomach and duodenum points</td>
</tr>
<tr>
<td>Upper lung</td>
<td>Directly above the heart point in the central depression of the cavum conchae</td>
</tr>
<tr>
<td>Sympathetic</td>
<td>At the end of the inferior antihelix crux</td>
</tr>
<tr>
<td><strong>Joint-specific point protocols (tailored to up to 3 of the patient’s most painful areas with each treatment)</strong></td>
<td></td>
</tr>
<tr>
<td>Knee:</td>
<td></td>
</tr>
<tr>
<td>SP-9</td>
<td>On the lower border of the medial condyle of the tibia in the depression posterior and inferior to the medial condyle of the tibia</td>
</tr>
<tr>
<td>SP-10</td>
<td>With knee flexed, 2 cun above the superior medial border of the patella on the bulge of the medial portion of quadriceps femoris (vastus medialis)</td>
</tr>
<tr>
<td>ST-34</td>
<td>With knee flexed, 2 cun above the superior lateral border of the patella on the line connecting with the anterior superior iliac spine (ASIS)</td>
</tr>
<tr>
<td>Fingers:</td>
<td></td>
</tr>
<tr>
<td>SI-3</td>
<td>When a loose fist is made, the point on the ulnar end of the crease in front of the metacarpalphalangeal joint of the little finger, at the junction of the two skins</td>
</tr>
<tr>
<td>Ba xie</td>
<td>On the dorsum of the hand, at the junction of the two skins, eight in all, making a loose fist to locate the points</td>
</tr>
<tr>
<td>LI-3</td>
<td>When a loose fist is made, the point on the radial side of the index finger, in the depression proximal to the head of the 2nd metacarpal bone</td>
</tr>
</tbody>
</table>
Joint-specific point protocols (tailored to up to 3 of the patient’s most painful areas with each treatment) (contd.)

| Lumbar: Du-3 | Below the spinous process of the fourth lumbar vertebrae, at the level with the crista iliaca |
| Lumbar: Du-8 | Below the spinous process of the ninth thoracic vertebrae |
| Lumbar: UB-23 | 1.5 cun lateral to Du-4 (below the spinous process of the second lumbar vertebrae) at the level of the lower border of the spinous process of the second lumbar vertebrae |
| Shoulder: LI-15 | Antero-inferior to the acromion, on the upper portion of m. deltoideus |
| Shoulder: SJ-14 | On the shoulder, posterior to LI-15, in the depression inferior and posterior to the acromion |
| Shoulder: SI-10 | Directly above the posterior end of the axillary fold, in the depression inferior to the scapular spine |
| Hip: GB-30 | At the junction of the lateral 1/3 and medial 2/3 of the distance between the prominence of the greater trochanter and the hiatus of the sacrum |
| Hip: GB-39 | 3 cun above the tip of the external malleolus on the posterior border of the fibula |
| Wrist: SI-5 | At the ulnar end of the transverse crease on the dorsal aspect of the wrist, in the depression between the styloid process and the ulnar and triquetral bone |
| Wrist: SJ-4 | On the transverse crease of the dorsum of the wrist, in the depression lateral to the tendon of fm. extensor digitorum communis |
| Wrist: LI-5 | When the thumb is tilted upward, the point in the depression between the tendons of m. extensor pollicis longus and brevis on the radial side of the wrist |

* A cun is a measurement used in finding acupuncture points. As each person has different body dimensions, a cun is defined according to the person whose body is to be treated. It is equivalent to the width of the distal phalanx of the person’s thumb, in the middle, at the crease.

Description of Sham Acupuncture

The sham acupuncture intervention, which is a control for acupoint specificity, will use superficial needle insertion at body locations not recognized as true acupoints. Similar to the true acupuncture arm, patients will be scheduled for twice weekly 45-minute acupuncture sessions (30-minute sham acupuncture, 15-minute set-up) for 6 weeks (8-12 sessions) followed by 6 weekly treatments (4-6 sessions) for maintenance (12-18 sessions total over 12 weeks). The sham acupuncture protocol includes full body and joint-specific treatments and an auricular sham intervention (Table 2). At every acupuncture visit, the patients assigned to sham acupuncture will receive the full body sham and auricular sham acupuncture prescriptions. In addition, each session will include a joint-specific sham point prescription tailored to up to four of the patient’s most painful joints, including knees, fingers, lumbar area, shoulders, hips, and wrists. The anatomic site selection may vary between visits depending on the patient’s current symptoms.

Acupuncture needles used will be Mac™ single-use, sterile, and disposable with plastic guide tube. The sham acupuncture needles will be 0.5 inch 34-gauge (manufactured by Tian Jinhaing Lin Sou Won Medical Instrument Co., Ltd; distributed by Mac Co., Roslyn
Heights, New York, USA). The needling protocol will consist of first swabbing all points with alcohol and needling full body and joint-specific sham points, inserting the needle by tapping in just to the upper level of the guide tube, at a transverse angle. The auricular sham procedure will include application of 3 adhesives to selected points on the ear (Table 2). The auricular sham product used will be Sakamura Magrain Ear Pellet, silver, with pellets removed (manufactured by Sakamura Lab & Co., Kyoto, Japan). The auricular adhesives are latex-free and will be loosely attached and pressed to the sham points on the helix of the auricle. The needles and auricular adhesives will remain in situ for 20-25 minutes during which the Study Acupuncturist will return to check the patient and lightly touch each needle handle and auricular adhesive in order to mimic the true intervention. No electrical stimulation will be used. The same Study Acupuncturists will administer both the true and sham acupuncture interventions. At 24 weeks, to assess the adequacy of blinding, patients will give an opinion about their treatment assignment. After the 24-week visit, patients will receive a voucher for 10 free true acupuncture sessions to be administered by the study acupuncturist(s) and used off-study at her convenience. Any additional acupuncture treatments received off study will be recorded at 24 and 52 weeks. All of this information will be recorded on the S1200 Assessment of Blinding and Additional Acupuncture Form.

Table 2. Sham Acupuncture Points

<table>
<thead>
<tr>
<th>Point Name</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Full body sham acupuncture points (given at each visit)</strong></td>
<td></td>
</tr>
<tr>
<td>Sham 1</td>
<td>On the lateral side of the left forearm, near the elbow, 3 cun below the olecranon, 0.5 cun toward the anterior of the small intestine meridian</td>
</tr>
<tr>
<td>Sham 2</td>
<td>On the lateral side of the right forearm, near the elbow, 3 cun below the olecranon, 0.5 cun toward the anterior of the small intestine meridian</td>
</tr>
<tr>
<td>Sham 3</td>
<td>At the lower border of the medial condyle of the left tibia, 1 cun anterior and superior to xi guan (Liv 7) of the liver meridian</td>
</tr>
<tr>
<td>Sham 4</td>
<td>At the lower border of the medial condyle of the right tibia, 1 cun anterior and superior to xi guan (Liv 7) point of the liver meridian</td>
</tr>
<tr>
<td><strong>Auricular (A) sham acupuncture points (given at in right ear on odd numbered visits and left ear on even numbered visits)</strong></td>
<td></td>
</tr>
<tr>
<td>A Sham 1</td>
<td>On the helix of the auricle between helix point #5 and #6</td>
</tr>
<tr>
<td>A Sham 2</td>
<td>On the helix of the auricle between helix point #4 and #3</td>
</tr>
<tr>
<td>A Sham 3</td>
<td>On the helix of the auricle between the ear apex and helix point #1</td>
</tr>
<tr>
<td><strong>Joint-specific sham point protocols (tailored to up to 3 of the patient’s most painful areas with each treatment)</strong></td>
<td></td>
</tr>
<tr>
<td>Knee</td>
<td>2 cun above sham 3 or 4, respectively</td>
</tr>
<tr>
<td>Fingers/ Wrist</td>
<td>On the lateral side of the left and right forearm, near the elbow, 5 cun below the olecranon, 0.5 cun toward the anterior of the small intestine meridian</td>
</tr>
<tr>
<td>Lumbar</td>
<td>On the back at the level of thoracic vertebra 8, 5 cun from the center of the spine, 2 cun from the outer channel of the urinary bladder meridian (needled bilaterally)</td>
</tr>
<tr>
<td>Shoulder</td>
<td>On the lateral side of the left and right arm, 5 cun below the anterior axillary fold, and 1 cun anterior to the lung channel</td>
</tr>
<tr>
<td>Hip</td>
<td>On the thigh, approximately 4 cun above the patella, 1 cun away from the anterior of the gall bladder meridian</td>
</tr>
</tbody>
</table>

Description of Waitlist Control

Patients randomized to the waitlist control group will undergo the scheduled evaluations at baseline, Week 2, Week 4, Week 6, Week 12, Week 24, and Week 52. Patients will be allowed to take acetaminophen and NSAIDs as needed for their joint symptoms, which will be documented as analgesic use at scheduled time points and captured on the S1200 Supplemental Agents Reporting Form. They must agree not to join any other clinical trial for the treatment of AI-related arthralgias while participating in this study.
After the 24 week visit, patients will receive a voucher for 10 free acupuncture sessions to be administered by the study acupuncturist(s) and used off-study at her convenience. Any additional acupuncture treatments received off-study will be recorded at 24 and 52 weeks. All of this information will be recorded on the S1200 Assessment of Blinding and Additional Acupuncture Form.
18.2 Patient-Completed Questionnaires

Joint symptoms (Brief Pain Inventory-Short Form [BPI-SF], Western Ontario and McMaster Universities Osteoarthritis [WOMAC] index for the hips and knees, Modified-Score for the Assessment and Quantification of Chronic Rheumatoid Affections of the Hands [M-SACRAH], and PROMIS Pain Impact-Short Form [PROMIS PI-SF]) and quality of life (Functional Assessment of Cancer Therapy-Endocrine Subscale [FACT-ES]) will be collected from self-administered questionnaires given at baseline, 6 weeks, 12 weeks, 24 weeks, and 52 weeks. (137,139, 140,155)

The BPI-SF is a 14-item questionnaire which asks patients to rate pain over the prior week and the degree to which it interferes with activities on a 0 to 10 scale. (137) Severity is measured as average pain, pain right now, worst pain, and least pain. The BPI-SF responds to both behavioral and pharmacological pain interventions and has been well-validated. Note that we consider a reduction of two or more points on the BPI-SF worst pain item (#2) to correspond to a clinically meaningful decrease in pain.

The WOMAC index Version 3.1 is a validated questionnaire for assessing osteoarthritis of the knees or hips and consists of 24 questions related to three subscales: pain (0-50), stiffness (0-20), and physical function (0-170). (139) Each question is answered by the patients along a standardized 10-point scale with terminal descriptors. The WOMAC (V3.1) osteoarthritis index has been tested extensively for validity, reliability, and responsiveness for measuring changes in symptoms after different interventions. Although originally developed as a standardized assessment to evaluate osteoarthritis, the WOMAC has been used as a generalized symptom scale to evaluate musculoskeletal pain from conditions such as rheumatoid arthritis, total knee replacements, femoral shaft fracture, and osteonecrosis of the femoral head. (155-159) The M-SACRAH consists of three domains assessing pain, stiffness, and functional status in patients suffering from hand osteoarthritis and rheumatoid arthritis, answered on 10-point scale. (140)

The Functional Assessment of Cancer Therapy-General (FACT-G) consists of four subscales including Physical, Social/Family, Emotional, and Functional Well-Being. We will focus on just Physical and Functional Well-Being for this trial. We will also use the Endocrine Subscale (ES) to monitor endocrine side effects caused by AIs, such as hot flashes and vaginal symptoms.(155) ACT scales have five response levels (“not at all” to “very much”), where higher scores reflect better well-being and fewer symptom problems.
18.3 Measurement of Blood and Urine Biomarkers

a. Blood Analyses

Patients will provide blood specimens at baseline, 6 weeks, 12 weeks, and 24 weeks for the measurement of serum hormone biomarkers (estradiol, FSH, LH) and inflammatory biomarkers (IL-6, IL-12, TNFα, CRP). Blood (~20 cc in two 10-ml red-top tubes) will be collected by standard venipuncture technique. Blood in the red top tubes is allowed to clot for 30 minutes to 1 hour (maximum) at room temperature, and centrifuged at 3,000 x g for 20 minutes immediately after clotting, or refrigerated or placed on ice for no more than 6 hours, and centrifuged at 3,000 x g for 20 minutes. Using a pipette, serum is then transferred to eight 1.8-ml Cryovials (1 ml each) and any extra serum should be aliquoted in a separate Cryovial.

Radioimmunoassays (RIAs) will be used to measure serum estradiol, FSH, and LH. Serum IL-6, IL-12, and TNFα will be determined by enzyme-linked immunosorbent assay (ELISA) using a commercially available kit (R&D Systems Inc., Minneapolis, MN). Serum hsCRP will be measured on the COBAS INTEGRA 400 plus system (Roche Diagnostic, Indianapolis, IN). All serum assays will be performed according to the manufacturer’s instructions and conducted at the Irving Center for Clinical Research (ICCR) at CUMC by Dr. S. Cremers.

b. Urine Analysis

At baseline, 6 weeks, 12 weeks, 24 weeks, and 52 weeks, approximately 10 ml of urine will be collected. Using a pipette, urine is then transferred to five 1.8-ml Cryovials (1.5 ml each) and any extra urine should be aliquoted in a separate Cryovial. The urine specimens will be sent in bulk shipment at the end of study on dry ice to the Sports Medicine Research and Testing Laboratory in Salt Lake City. The urine assays will be performed using high performance liquid chromatography (HPLC) tandem mass spectrometry.

c. Genotyping Analysis

At baseline, patients will provide additional blood specimens for plasma/buffy coat to be used for genotyping analyses. Approximately 10 ml of blood will be drawn in a 10-ml EDTA purple-top tube (after the serum sample so that the plasma can be mixed without delay). Just prior to collection of the specimen, it is important to rotate the 10-ml purple-top tube to allow the EDTA solution to coat the inside of the tube. Following collection of the blood, the sample must be inverted gently 10-15 times to mix well.

Genomic DNA will be extracted from the buffy coat (gray layer of WBC) using standard RNase/proteinase K technique. All samples will be genotyped using the ABI PRISM 7900HT Sequence Detection System (Applied Biosystems, Foster City, CA) in a 96-well format. The fluorogenic 5’ nuclease or TaqMan assay will be used to determine genotype in the CYP19A1 aromatase gene. Genotyping will be done by laboratory personnel blinded to treatment assignment. All genotyping studies will be conducted in the laboratory of Dr. R. Santella (CUMC).
18.4 Training of Study Acupuncturists to the Study Protocol

A web-based site initiation visit and formal training of the Study Acupuncturists and a web-based training for Clinical Research Associates (CRAs) administering the functional testing at each site will be performed at the time of study activation. At least two acupuncturists licensed with the appropriate certifications for the state at each site will be trained and certified annually in administering the protocol and will perform all treatments. Each site already has experience in conducting acupuncture clinical trials.

CUMC-supplied study materials, including single-use, sterile, and disposable acupuncture needles, a training manual and online training module of the point protocols for both the true and sham acupuncture interventions and a demonstration of the grip strength and ‘Timed Get Up and Go’ (TGUG) testing, will be provided to each participating site. In addition, sites will be supplied with standardized dynamometers for grip strength testing and stop watches for the ‘Timed Get Up and Go’ test.

The acupuncture training manual will consist of the following:

- Written point protocols and anatomic renderings of all true and sham acupuncture points
- Written description about basic clean needle technique and administration of true and sham acupuncture interventions
- Written explanation for filling out case report forms done by the Study Acupuncturist

The acupuncture online training module will consist of the following:

- Video recording of the true and sham acupuncture points on a human volunteer
- Video recording of true acupuncture needling intervention in 1-2 selected body points and the auricular points
- Video recording of the sham acupuncture needling intervention in 1-2 selected body points and auricular points (taping)
18.5 CRA Functional Testing Training

The functional testing online training module will consist of the following:

- Video recording of how to perform the dynamometer measurements for grip strength testing
- Video recording of the components of the ‘Timed Get Up and Go’ test including how to use the stopwatch and a recording of a human volunteer enacting the test
18.6 Emergency Unblinding Guidelines

a. General Considerations

In this intervention trial, S1200 participants AND study staff are blinded to two of the three treatment assignments, True Acupuncture or Sham Acupuncture (double blind conditions). Guidelines and Procedures for unblinding in emergency situations are given below (see Sections 18.6c and d).

In general, participants SHOULD NOT be unblinded to the treatment assignment unless a condition exists where the knowledge of the participant’s treatment assignment would directly influence or affect her immediate care, or if an emergency situation arises.

Emergency unblinding of all S1200 treatment assignments will be done by the Washington Poison Center (WPC) with medical advice from a panel of two resource physicians appointed for this study. Calls for information about the treatment assignment, or the study in general, should be directed to the SWOG Statistical Center.

b. Guidelines for Discontinuation of Study Intervention

(Unblinding is NOT appropriate when acupuncture is discontinued.) The following events MAY require PERMANENT discontinuation of Coded acupuncture:

- Evidence of new cancer or cancer recurrence at any time.
- Unacceptable toxicity (as defined in Section 8.0)
- A participant who becomes pregnant while on treatment will be removed from protocol treatment
- Cumulative delay ≥ 30 consecutive days after study registration
- Completion of 12 weeks of study intervention (acupuncture).

Permanent discontinuation of the Coded intervention for any reason must be documented on the S1200 Off Protocol Form.

The following events MAY require TEMPORARY discontinuation of Coded Intervention:

- Treatment of and/or hospitalization for a medical problem;
- Assessment of symptoms or side effects potentially related to Coded Intervention.

Temporary discontinuation of the Coded Intervention for any reason must be documented on the S1200 Off Protocol Form.

Refer to Section 7.0 (Procedures and Study Plan) and Section 8.0 (Safety Monitoring) of this protocol for details on discontinuation of study intervention.
c. Guidelines for Emergency Unblinding of Treatment Assignment

The following event MAY require emergency unblinding of treatment assignment:

- A compelling medical need as determined by a physician, e.g., occurrence of a severe or life-threatening reaction, inclusive of an adverse drug reaction, which may have been attributable to treatment assignment, or existence of a condition where the knowledge of the participant's treatment assignment would directly influence or affect her immediate care;

d. Procedure for Emergency Unblinding

The procedure for unblinding the treatment assignment for a participant is as follows:

- All unblinding must be done by the registering physician or designee.
- Call the Washington Poison Control (WPC) collect at 206/526-2121 or at 800/732-6985 if calling from within Washington State. The WPC is accessible 24 hours per day, 365 days per year for unblinding calls. Informational calls should be directed to the Data Operations Center in Seattle during standard business hours.
- Provide the WPC with the following information:
  - Study number: S1200
  - SWOG patient number
  - Participant name
  - Name and telephone number of the caller
  - Reason unblinding is required
- Unblinding for a "compelling medical need" must be authorized by a physician designated as a resource physician for this protocol.

The treating physician (or designee) would provide the WPC with the information needed to determine if unblinding is required for the participant. The WPC would contact the resource physician, provide the required information, and obtain the authorization to unblind, if necessary. Based on the decision of the resource physician, the WPC would call the treating physician with either the unblinded treatment assignment or a treatment recommendation from the resource physician.

If a resource physician cannot be reached by the WPC, treatment of the participant should proceed as if the patient received True Acupuncture.

Unblinding of treatment assignment for any reason must be documented on the S1200 Off Protocol Form. All unblinded patients are taken off treatment and followed per the requirements of SWOG.

Any questions regarding unblinding may be directed to one of the following resource physicians:
Informed Consent Model for S1200

*NOTES FOR LOCAL INSTITUTION INFORMED CONSENT AUTHORS:

This model informed consent form has been reviewed by the DCT/NCI and is the official consent document for this study. Local IRB changes to this document are allowed. (Institutions should attempt to use sections of this document that are in bold type in their entirety.) Editorial changes to these sections may be made as long as they do not change information or intent. If the institutional IRB insists on making deletions or more substantive modifications to the risks or alternatives sections, they may be justified in writing by the investigator and approved by the IRB. Under these circumstances, the revised language, justification and a copy of the IRB minutes must be forwarded to the SWOG Operations Office for approval before a patient may be registered to this study.

Please particularly note that the questions related to banking of specimens for future study are in bolded type and may not be changed in any way without prior approval from the SWOG Operations Office.

Instructions and examples for informed consent authors are in [italics].
A blank line, __________, indicates that the local investigator should provide the appropriate information before the document is reviewed with the prospective research patient.
The term "study doctor" has been used throughout the model because the local investigator for a cancer treatment trial is a physician. If this model is used for a trial in which the local investigator is not a physician, another appropriate term should be used instead of "study doctor".
The dates of protocol updates in the header and in the text of the consent is for reference to this model only and should not be included in the informed consent form given to the prospective research patient.
The local informed consent must state which parties may inspect the research records. This includes the NCI, NCCAM, Office of Research on Women's Health (ORWH), any companies or grantors that are providing study support (these will be listed in the protocol's model informed consent form) and SWOG.

"SWOG" must be listed as one of the parties that may inspect the research records in all protocol consent forms for which patient registration is being credited to SWOG. This includes consent forms for studies where all patients are registered directly through the SWOG Data Operations Office, all intergroup studies for which the registration is being credited to SWOG (whether the registration is through the SWOG Data Operations Office or directly through the other group), as well as consent forms for studies where patients are registered via CTSU and the registration is credited to SWOG.

Readability Statistics:

<table>
<thead>
<tr>
<th>Measure</th>
<th>Score</th>
<th>Target</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flesch Reading Ease</td>
<td>62</td>
<td>&gt;55</td>
</tr>
<tr>
<td>Flesch-Kincaid Grade Level</td>
<td>8.5</td>
<td>&lt;8.5</td>
</tr>
</tbody>
</table>

• Instructions and examples for informed consent authors are in [italics].
• A blank line, __________, indicates that the local investigator should provide the appropriate information before the document is reviewed with the prospective research patient.
• The term "study doctor" has been used throughout the model because the local investigator for a cancer treatment trial is a physician. If this model is used for a trial in which the local investigator is not a physician, another appropriate term should be used instead of "study doctor".
• The dates of protocol updates in the header and in the text of the consent is for reference to this model only and should not be included in the informed consent form given to the prospective research patient.
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"SWOG" must be listed as one of the parties that may inspect the research records in all protocol consent forms for which patient registration is being credited to SWOG. This includes consent forms for studies where all patients are registered directly through the SWOG Data Operations Office, all intergroup studies for which the registration is being credited to SWOG (whether the registration is through the SWOG Data Operations Office or directly through the other group), as well as consent forms for studies where patients are registered via CTSU and the registration is credited to SWOG.
• When changes to the protocol require revision of the informed consent document, the IRB should have a system that identifies the revised consent document, in order to preclude continued use of the older version and to identify file copies. An appropriate method to identify the current version of the consent is for the IRB to stamp the final copy of the consent document with the approval date. The stamped consent document is then photocopied for use. Other systems of identifying the current version of the consent such as adding a version or approval date are allowed as long as it is possible to determine during an audit that the patient signed the most current version of the consent form.

*NOTES FOR LOCAL INVESTIGATORS:

• The goal of the informed consent process is to provide people with sufficient information for making informed choices. The informed consent form provides a summary of the clinical study and the individual’s rights as a research patient. It serves as a starting point for the necessary exchange of information between the investigator and potential research patient. This model for the informed consent form is only one part of the larger process of informed consent. For more information about informed consent, review the "Recommendations for the Development of Informed Consent Documents for Cancer Clinical Trials" prepared by the Comprehensive Working Group on Informed Consent in Cancer Clinical Trials for the National Cancer Institute. The Web site address for this document is http://cancer.gov/clinicaltrials/understanding/simplification-of-informed-consent-docs/

• A blank line, _________, indicates that the local investigator should provide the appropriate information before the document is reviewed with the prospective research patient.

• Suggestion for Local Investigators: An NCI pamphlet explaining clinical trials is available for your patients. The pamphlet is titled: "If You Have Cancer…What You Should Know about Clinical Trials". This pamphlet may be ordered on the NCI Web site at https://cissecure.nci.nih.gov/ncipubs or call 1-800-4- CANCER (1-800-422-6237) to request a free copy.

• Optional feature for Local Investigators: Reference and attach drug sheets, pharmaceutical information for the public, or other material on risks. Check with your local IRB regarding review of additional materials.

*These notes for authors and investigators are instructional and should not be included in the informed consent form given to the prospective research patient.
"A Randomized Blinded Sham- and Waitlist-Controlled Trial of Acupuncture for Joint Symptoms Related to Aromatase Inhibitors in Women with Early Stage Breast Cancer"

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 are a postmenopausal woman currently taking an aromatase inhibitor for the treatment of early stage breast cancer and experiencing joint pain or stiffness. Aromatase inhibitors are a type of hormonal therapy for breast cancer, including anastrozole (Arimidex), letrozole (Femara), and exemestane (Aromasin).

Who is doing this study?

SWOG is sponsoring this trial. SWOG is an adult cancer clinical trials organization. SWOG is funded through the National Cancer Institute, and its network consists of almost four thousand physicians at almost three hundred institutions throughout the United States. Your study doctor has met all requirements to be a member of SWOG and to perform National Cancer Institute-funded research through this Group. This particular study is also being supported by the National Center for Complementary and Alternative Medicine and by the Office for Research on Women's Health.

Why is this study being done?

The purpose of this study is to test whether 12-week use of acupuncture can be used to ease joint pain or stiffness related to aromatase inhibitors for breast cancer treatment. We want to find out what effects, good and/or bad, it has on you and your joint symptoms.

Acupuncture is a traditional Chinese medical treatment which involves the use of very small, thin needles to stimulate specific points in the body. Previous studies have found a benefit of acupuncture for the treatment of knee and back pain. This study will test whether acupuncture can reduce joint pain associated with aromatase inhibitor therapy for breast cancer.

How many people will take part in the study?

About 228 people will take part in this study.
What will happen if I take part in this research study?

Before you begin the study …

You will need to have the following exams, tests or procedures to find out if you can be in the study. These exams, tests or procedures are part of regular health care for postmenopausal women and may be done even if you do not join the study. If you have had some of them recently, they may not need to be repeated. This will be up to your study doctor.

- Medical history and physical exam
- Blood work to check your menopausal status (if necessary).
- You will be asked to take one questionnaire that will ask you to rate your joint pain and stiffness.

During the study …

If the exams, tests and procedures show that you can be in the study, and you choose to take part, then you will need the following procedure.

Randomization

You will be "randomized" into one of the study groups described below. Randomization means that you are put into a group by chance. A computer program will place you in one of the study groups. Neither you nor your doctor can choose the group you will be in.

You will have a 50% chance of receiving true acupuncture, a 25% chance of receiving sham acupuncture (light acupuncture at non-acupuncture points), and a 25% chance of being in the waitlist control group (you will wait 24 weeks before having the choice to receive true acupuncture). You will not be told whether you are receiving either true or sham acupuncture. Only the acupuncturist and staff members at SWOG and Columbia University will be aware of the acupuncture assignment. These are described below.

If you are in the true or sham acupuncture groups, you will be scheduled for 45-minute acupuncture sessions given twice a week for six weeks, then weekly for another six weeks for a total of twelve weeks. If you are in the waitlist control group, you will not receive true acupuncture until after the 24-week evaluation, if you choose. No matter what group you are assigned to after 24 weeks you will receive 10 free vouchers for true acupuncture. You will be allowed to take certain pain medications as necessary, such as acetaminophen (Tylenol) and anti-inflammatory drugs (such as aspirin, ibuprofen, Celebrex). For the duration of the study, you will be asked not to take steroids or narcotics (opiates) for pain relief or start new treatments for pain relief, if possible. The study will take fifty-two weeks to complete. Follow-up visits will be scheduled at six weeks, twelve weeks, twenty-four weeks, and fifty-two weeks.
You will complete the following questionnaires and procedures after randomization.

- **Questionnaires** - Six self-administered questionnaires (which will take about 30 minutes to complete) will be given at the beginning of the study to collect information about how you are feeling physically during your treatment and how you are performing your daily activities. You will be asked to rate joint pain and stiffness, and will be asked to respond to questions regarding aromatase inhibitors and pain medications. If any questions make you feel uncomfortable, you may skip those questions and not give an answer. We will do our best to make sure that your personal information will be kept private.

- **Hand grip strength and activity** - You will be tested for hand grip strength and an activity called “Timed Get Up and Go”. The interviewer will ask you to take a short walk of 10 feet then measure the time it takes you to do so (~15 minutes to complete).

- **Pain medications** - Assess the amount and how often you have taken pain medications.

- **Blood and urine samples** - Blood (about 2 tablespoons) and urine (about 1/3 of a cup) will be taken before starting the study intervention. The blood and urine samples are not part of the usual medical care. The blood and urine samples will be submitted to look at hormone levels, specific biomarker (biological molecule found in blood that is a sign of a normal or abnormal process, or of a condition or disease), inflammatory markers, break down products from the aromatase inhibitors, and DNA (genetic) studies. These evaluations will be done at the end of the entire study and will not be part of your medical record. You will be asked if remaining blood and urine from the samples can be kept for future research purposes. Please see separate questions at the end of this document.

During telephone interviews at 2 week and 4 weeks …

The medical team or staff will record:

- Your acupuncture schedule (if applicable)
- Side effects you may be having
- Pain treatments you are taking

During study visits at 6, 12, 24 and 52 weeks …

You will have the following procedures:

- **Questionnaires** - Six self-administered questionnaires (which will take about 30 minutes to complete) will be given at six and twelve weeks to collect information about how you are feeling physically during your treatment and how you are performing your daily activities. You will be asked to rate joint pain and stiffness. You will also give your opinion about whether you were receiving true acupuncture or sham acupuncture (if applicable). If any questions make you feel uncomfortable, you may skip those questions and not give an answer. We will do our best to make sure that your personal information will be kept private.
• Functional testing for hand grip strength and an activity called “Timed Get Up and Go” during which the interviewer will ask you to take a short walk of 10 feet then measure the time it takes you to do so (~15 minutes to complete)
• Blood and urine samples – Blood (about 1.5 tablespoons) and urine (about 1/3 of a cup) will be taken at six, twelve and twenty-four weeks. Urine samples only will be taken at 52 weeks. The blood and urine samples are not part of the usual medical care. The blood and urine samples will be submitted to look at hormone levels and inflammatory markers. These results will not be made available to you or your study doctor. You will be asked if remaining blood and urine from the samples can be kept for future research purposes. Please see separate consent form at the end of this document.

The medical team or staff will record:

• Your acupuncture schedule (if applicable)
• Side effects you may be having (at Weeks 6 and 12)
• Pain treatments you are receiving
• Additional acupuncture treatments received outside of the study (at Weeks 24 and 52)

During telephone interviews at 16 weeks and 20 weeks …

The medical team or staff will record:

• Your rating of joint pain and stiffness
• Pain treatments you are receiving

At 24 weeks, all 3 groups will receive vouchers for 10 true acupuncture sessions.

How long will I be in the study?

You will be asked to take part in the study for about 52 weeks.

Can I stop being in the study?

Yes. You can decide to stop at any time. Tell the study doctor if you are thinking about stopping or decide to stop.

The study doctor may stop you from taking part in this study at any time if he/she believes it is in your best interest or if the study is stopped.
What side effects or risks can I expect from being in the study?

You may have side effects while on the study. Everyone taking part in the study will be watched carefully for any side effects. However, doctors don’t know all the side effects that may happen. Side effects may be mild or very serious. Your health care team may give you medicines to help lessen side effects. Many side effects go away soon after you stop the acupuncture intervention. In some cases, side effects can be serious, long lasting, or may never go away.

You should talk to your study doctor and acupuncturist about any side effects that you have while taking part in the study.

Risks and side effects related to acupuncture include those which are:

- Likely (> 20%)
  - Aching, warm or tingling sensation from needling

- Less Likely (~5%)
  - Needle shock or feeling ‘faint’ from needle insertion. If this should occur, then the acupuncture needles will be removed and you will be laid flat on the examination table for about 15 minutes or until the symptoms subside.
  - Bruising and minor bleeding

- Rarely (< 1%)
  - Infection

Are there benefits to taking part in the study?

If you agree to take part in this study, there may or may not be direct medical benefit to you. We hope the information learned from this study will benefit other patients with joint symptoms associated with aromatase inhibitor therapy for breast cancer treatment in the future.

What other choices do I have if I do not take part in this study?

Your other choices may include:
- Getting treatment without being in a study
- Taking part in another study
- Getting no treatment

Talk to your doctor about your choices before you decide if you will take part in this study.
Will my medical information be kept private?

We will do our best to make sure that the personal information in your medical record will be kept private. However, we cannot guarantee total privacy. Your personal information may be given out if required by law. If information from this study is published or presented at scientific meetings, your name and other personal information will not be used.

Organizations that may look at and/or copy your medical records for research, quality assurance, and data analysis include:

- SWOG
- The National Cancer Institute (NCI), National Center for Complementary and Alternative Medicine (NCCAM), Office for Research on Women's Health (ORWH) and other government agencies, like the Food and Drug Administration (FDA), involved in keeping research safe for people

What are the costs of taking part in this study?

Taking part in this study may lead to added costs that you or your health insurance plan may be responsible for. Please ask about any expected added costs or insurance problems.

All tests and procedures which are not part of routine clinical care, including the blood and urine tests, will be covered by the study. The acupuncture sessions will be provided free of charge.

Regardless of study arm you are assigned, you will be offered a voucher for ten free true acupuncture sessions to be used off-study at your convenience after the 24-week visit.

For more information on clinical trials and insurance coverage, you can visit the National Cancer Institute’s Web site at http://m.cancer.gov/topics/clinicaltrials/learningabout/payingfor/howinsurance-companies-decide. (updated 1/22/15) You can print a copy of the "Clinical Trials and Insurance Coverage" information from this Web site.

Another way to get the information is to call 1-800-4-CANCER (1-800-422-6237) and ask them to send you a free copy.

What happens if I am injured because I took part in this study?

It is important that you tell your study doctor, ____________________ [investigator’s name(s)], if you feel that you have been injured because of taking part in this study. You can tell the doctor in person or call him/her at ____________________ [telephone number].

You will get medical treatment if you are injured as a result of taking part in this study. You and/or your health plan will be charged for this treatment. The study will not pay for medical treatment.
What are my rights if I take part in this study?

Taking part in this study is your choice. You may choose either to take part or not to take part in the study. If you decide to take part in this study, you may leave the study at any time. No matter what decision you make, there will be no penalty to you and you will not lose any of your regular benefits. Leaving the study will not affect your medical care. You can still get your medical care from our institution.

In the case of injury resulting from this study, you do not lose any of your legal rights to seek payment by signing this form.

Who can answer my questions about the study?

You can talk to your study doctor about any questions or concerns you have about this study. Contact your study doctor __________________ [name(s)] at __________________ [telephone number].

For questions about your rights while taking part in this study, call the __________________ [name of center] Institutional Review Board (a group of people who review the research to protect your rights) at __________________ (telephone number). [Note to Local Investigator: Contact information for patient representatives or other individuals in a local institution who are not on the IRB or research team but take calls regarding clinical trial questions can be listed here.]

Where can I get more information?

You may call the National Cancer Institute's Cancer Information Service at:

1-800-4-CANCER (1-800-422-6237) or TTY: 1-800-332-8615

You may also visit the NCI Web site at http://cancer.gov/

- A description of this clinical trial will be available on http://www.ClinicalTrials.gov. This Web site will not include information that can identify you. At most, the Web site will include a summary of study results. You can search this Web site at any time.

[Note to Informed Consent Authors: the above paragraph complies with the new FDA regulation found at 21 CFR 50.25(c) and must be included verbatim in all informed consent documents. The text in this paragraph cannot be revised.]

- For NCI’s clinical trials information, go to: http://cancer.gov/clinicaltrials/
- For NCI’s general information about cancer, go to: http://cancer.gov/cancerinfo/
• For the National Center for Complementary and Alternative Medicine (NCCAM), go to: http://nccam.nih.gov/.

• For the Office of Research on Women's Health (ORWH), go to http://orwh.od.nih.gov/.

You will get a copy of this form. If you want more information about this study, ask your study doctor.

Future Contact
Occasionally, researchers working with SWOG may have another research idea that relates to people who were on a SWOG study. In some cases, to carry out the new research, we would need to contact patients in a particular study. You can agree or not agree to future contact by circling “yes” or “no”.

I agree to allow my study doctor, or someone approved by my study doctor, to contact me regarding future research involving my participation in this study.

Yes    No

Consent Form for Use of Specimens for Research

We would like to keep some of the blood and urine specimens that are left over for future research. If you agree, these specimens will be kept and may be used in research to learn more about cancer and other diseases. Please read the information sheet called "How Are Specimens Used for Research" to learn more about tissue research.

The research that may be done with your specimens are not designed specifically to help you. It might help people who have cancer and other diseases in the future.

Reports about research done with your specimens will not be given to you or your doctor. These reports will not be put in your health record. The research will not have an effect on your care.

Things to Think About

The choice to let us keep the left over specimens for future research is up to you. No matter what you decide to do, it will not affect your care.

If you decide now that your specimens can be kept for research, you can change your mind at any time. Just contact us and let us know that you do not want us to use your specimens. Then any specimens that remain will no longer be used for research.

In the future, people who do research may need to know more about your health. While SWOG may give them reports about your health, it will not give them your name, address, phone number, or any other information that will let the researchers know who you are.
Sometimes specimens are used for genetic research (about diseases that are passed on in families). Even if your specimens are used for this kind of research, the results will not be put in your health records.

If your confidential genetic information is discovered, you may suffer from genetic discrimination. Genetic discrimination occurs if people are treated unfairly because of differences in their genes that increase their chances of getting a certain disease. In the past, this could have resulted in the loss of health insurance or employment. Because of this, The Genetic Information Nondiscrimination Act of 2008, also referred to as GINA, was passed by Congress to protect Americans from such discrimination. The new law prevents discrimination from health insurers and employers. This act was signed into federal law on May 21, 2008, and went into effect May 2009. This law does not cover life insurance, disability insurance and long-term care insurance.

While this study has safeguards in place to protect your confidential genetic information and to make it extremely unlikely that your identity would be connected with any special studies that are performed on your tissue, it is possible that this information could be discovered by someone who is unauthorized to have access to it.

Your specimens will be used only for research and will not be sold. The research done with your specimens may help to develop new products in the future.

Benefits

The benefits of research using specimens include learning more about what causes cancer and other diseases, how to prevent them, and how to treat them.

Risks

The greatest risk to you is the release of information from your health records. We will do our best to make sure that your personal information will be kept private. The chance that this information will be given to someone else is very small.

Making Your Choice

Please read each sentence below and think about your choice. After reading each sentence, circle "Yes" or "No." If you have any questions, please talk to your doctor or nurse, or call our research review board at IRB's phone number.

No matter what you decide to do, it will not affect your care.
Future Use of Specimens

1. My specimens may be kept for use in research to learn about, prevent, treat or cure cancer.

   Yes  No

2. My specimens may be kept for use in research about other health problems (for example: diabetes, Alzheimer's disease, or heart disease).

   Yes  No

3. Someone may contact me in the future to ask me to allow other uses of my specimens.

   Yes  No

If you decide to withdraw your specimens from the SWOG Repository in the future, a written withdrawal of consent should be submitted by your study doctor to the SWOG Operations Office. Please designate in the written withdrawal whether you would prefer to have the specimens destroyed or returned to the study doctor.

Signature

I have been given a copy of all [insert total of number of pages] pages of this form. I have read it or it has been read to me. I understand the information and have had my questions answered. I agree to take part in this study.

Patient __________________________________________

Date ____________________________________________
Specimen Consent Supplemental Sheets

How are Specimens Used for Research?

Where do specimens come from?

A specimen may be from a blood sample or from bone marrow, tissue from a biopsy, skin, toenails or other body materials. People who are trained to handle specimens and protect donors’ rights make sure that the highest standards of quality control are followed by SWOG. Your doctor does not work for SWOG, but has agreed to help collect specimens from many patients. Many doctors across the country are helping in the same way.

Why do people do research with specimens?

Research with specimens can help to find out more about what causes cancer, how to prevent it, how to treat it, and how to cure it. Research using specimens can also answer other health questions. Some of these include finding the causes of diabetes and heart disease, or finding genetic links to Alzheimer’s.

What type of research will be done with my specimen?

Many different kinds of studies use specimens. Some researchers may develop new tests to find diseases. Others may develop new ways to treat or even cure diseases. In the future, some of the research may help to develop new products, such as tests and drugs. Some research looks at diseases that are passed on in families (called genetic research). Research done with your specimen may look for genetic causes and signs of disease.

How do researchers get the specimen?

Researchers from universities, hospitals, and other health organizations conduct research using specimens. They contact SWOG and request samples for their studies. SWOG reviews the way that these studies will be done, and decides if any of the samples can be used. SWOG gets the specimen and information about you from your hospital, and sends the specimen samples and some information about you to the researcher. SWOG will not send your name, address, phone number, social security number or any other identifying information to the researcher.

Will I find out the results of the research using my specimen?

You will not receive the results of research done with your specimen. This is because research can take a long time and must use specimen samples from many people before results are known. Results from research using your specimen may not be ready for many years and will not affect your care right now, but they may be helpful to people like you in the future.

Why do you need information from my health records?

In order to do research with your specimen, researchers may need to know some things about you. (For example: Are you male or female? What is your race or ethnic group? How old are you? Have you ever smoked?) This helps researchers answer questions about diseases. The information that will be given to the researcher may include your age, sex, race, diagnosis, treatments and family history. This information is collected by your hospital from your health record and sent to SWOG. If more information is needed, SWOG will send it to the researcher.

Will my name be attached to the records that are given to the researcher?

No. Your name, address, phone number and anything else that could identify you will be removed before they go the researcher. The researcher will not know who you are.
**How could the records be used in ways that might be harmful to me?**

Sometimes, health records have been used against patients and their families. For example, insurance companies may deny a patient insurance or employers may not hire someone with a certain illness (such as AIDS or cancer). The results of genetic research may not apply only to you, but to your family members too. For disease caused by gene changes, the information in one person's health record could be used against family members.

**How am I protected?**

SWOG is in charge of making sure that information about you is kept private. SWOG will take careful steps to prevent misuse of records. Your name, address, phone number and any other identifying information will be taken off anything associated with your specimen before it is given to the researcher. This would make it very difficult for any research results to be linked to you or your family. Also, people outside the research process will not have access to results about any one person which will help to protect your privacy.

**What if I have more questions?**

If you have any questions, please talk to your doctor or nurse, or call our research review board at (Insert IRB's Phone Number).