CONSENT FORM TO PARTICIPATE IN THE
WOMEN’S HEALTH INITIATIVE EXTENSION STUDY

WHI Clinical Coordinating Center
The Fred Hutchinson Cancer Research Center
Seattle, Washington

[Clinical Center]

[Principal Investigator]

[Other Investigators, as appropriate]

[24-Hour Contact]

This form is to tell you about the Women’s Health Initiative (WHI) Extension Study. You are invited to join the WHI Extension Study because you were a participant in either the WHI Clinical Trial or the WHI Observational Study. The WHI Extension Study will extend your participation in the WHI study through 2010. All participants in the WHI Diet Modification, Hormone Program, and Calcium and Vitamin D Clinical Trials, regardless of their treatment assignments to control or placebo or active interventions, and those in the WHI Observational Study are invited to participate in the WHI Extension Study. We expect up to 150,000 participants across the United States to join and continue to contribute important information to women’s health.

Purpose of the WHI Extension Study

The main purpose of the WHI Extension Study is to continue to learn about the health of postmenopausal women over an additional five years of follow-up. We hope to learn more about risks for heart disease, stroke, blood clots, breast cancer, colon cancer, other cancers, and osteoporotic fractures (broken bones), as well as quality of life. For women who were in the WHI Clinical Trials, we hope to learn if the risks or benefits for these health conditions continue or disappear after women have stopped taking their WHI study pills, are no longer in contact with the study nutritionists, or are continuing to follow their usual diet. We also hope to learn if these changes are different for women who were originally assigned to take active or placebo study pills or were in the Dietary Change or Comparison group. For women in the Clinical Trials and the Observational Study, we expect that five additional years of follow-up will help us to learn more about both common and more rare diseases of aging.

Reasons for the WHI Extension Study

Results from the WHI Clinical Trials and Observational Study have given health care providers important information about the effects of diet, hormones, and calcium/vitamin D supplements and initial health characteristics, including family and medical history and lifestyle factors, such as activity levels, on diseases of aging in postmenopausal women. As more and more women are re-evaluating their diet, use of hormone therapy, and other factors, it is important to understand how
how maintaining or changing these factors affects disease over time. These questions have been asked by health care providers and the millions of women who are moving into the next decade of their lives. The WHI Extension Study can find answers by comparing changes in health status over the course of the initial WHI study period with those in the five-year follow-up period.

**What Will You Be Doing?**

If you decide to join the WHI Extension study, you will be asked to fill out health forms each year. You will not be asked to come into the WHI Clinical Center for these activities. The health forms will be similar to the medical history updates you were asked to complete regularly while you were in the WHI Clinical Trial or Observational Study. For the Extension Study, you will only be asked to complete the forms once a year through 2010. The form will be mailed to you, with a pre-paid mailing envelope, and you will be asked to return your completed form by mail. If the WHI staff has any questions after receiving your forms, they will contact you by telephone. We may also ask you to sign a Medical Release form to get more detailed information about health issues that you report on the form.

The amount of time asked of you to participate in the WHI Extension Study will be:

<table>
<thead>
<tr>
<th>Activity</th>
<th>Total Time</th>
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<tr>
<td>Completing health forms</td>
<td>About 30 minutes each year through 2010</td>
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**Benefits and Risks**

We cannot and do not guarantee or promise that you will receive any personal benefits from this study; however, there is also little risk. By taking part in the Extension Study, you will help advance scientific knowledge about postmenopausal women’s health.

There are no risks to completing the health update forms for this study.

**Costs**

You will not be paid for being in the study. The WHI has not set aside funds to pay for any health conditions that you may develop, and will not pay for any health problems or conditions that might occur during the course of this study. These might be covered by your usual insurance, Medicare or Medicaid. You will not be paid back for any wages lost from taking part in this study. If you have questions about your costs, financial responsibilities, and/or medical insurance coverage for this activity, please contact [CC financial office or other budgetary person and institutional department] at [phone number].

**Confidentiality**

All of your study records will be kept confidential to the extent required by law. Your personal identity will not be revealed in any publication or release of results. Only WHI staff at the [Name] Clinical Center and the Clinical Coordinating Center at the Fred Hutchinson Cancer Research Center in Seattle, Washington, will have access to your study number, name, social security number and address for the purpose of study wide mailings, as well as maintaining and updating your study records. The Food and Drug Administration may be allowed to examine your study records for safety reasons. Study records will be kept indefinitely for analysis and follow-up. This study is authorized by Privacy Act 42 United States Code 241.
ACCESS TO YOUR INFORMATION
You have the right to limit the use and sharing of your health information, and you have the right to see your medical records and know who else is seeing them. Everyone involved in this study respects your privacy. Any and all information about you obtained for this research study will be kept confidential and will not be released for any reason without your written permission unless compelled by law. In addition:
- staff are trained to protect your privacy and sign an agreement to do so;
- paper records are stored in locked files;
- computer records are password protected;
- your name or other personal information will not be included with any data shared with anyone who is not part of the WHI Study Research Staff.

USE AND DISCLOSURE OF YOUR MEDICAL INFORMATION
By signing this consent form, you are authorizing the use and disclosure of your health information collected in connection with your participation in this research study. Your information will only be used as we have described in this consent form and any applicable laws. If you decide to stop participating in the study, you may revoke your authorization, except to the extent that the law allows us to continue using your information.

What Information Will Be Used or Disclosed?
Health information related to this study may include, but is not limited to: demographic information, the results of physical exams, blood tests and related records, x-rays, and other diagnostic and medical procedures. This information may be used or disclosed in connection with this research study. We may request from your physician a copy of your medical record related to your diagnosis and treatment of cancer or other diseases if we have obtained from you a signed Medical Release form.

Who May Use or Disclose the Information?
The following parties are authorized to use and/or disclose your health information in connection with this research study:
- The [CC] WHI Research Staff
- The [CC] Administrative Panel on Human Subjects in Medical Research or Institutional Review Board (IRB) and any other unit of the [CC], as necessary
- The WHI Clinical Coordinating Center Staff at the Fred Hutchinson Cancer Research Center in Seattle, Washington
- The Institutional Review Board at the Fred Hutchinson Cancer Research Center

Who May Receive / Use the Information?
The parties listed in the preceding paragraph may disclose your health information to the following persons and organizations for their use in connection with this research study:
- The Office for Human Research Protections in the U.S. Department of Health and Human Services
- The Food and Drug Administration (FDA)
Your information may be redisclosed if the recipients described above are not required by law to protect the privacy of the information.

Expiration
Your authorization for the use and disclosure of your health information will continue indefinitely.

Right to Withdraw
Your decision to be in the study is voluntary. You may quit at any time, for any reason, without notice. We hope that you will take part for the entire time of the study because we need all of this information to draw correct conclusions. If you decide to leave the study, it will not affect your regular medical care.

Other Information
Your joining is important to the success of this study. You may join the WHI Extension Study whether or not you were in the active or placebo group of the WHI Hormone or Calcium and Vitamin D Clinical Trials, in the Dietary Change or Comparison group, or in the WHI Observational Study. You may also join even if you have started taking hormones prescribed by your health care provider or have made any changes to your diet or use of calcium supplements. We have tried to make joining the study as easy as possible for you. Please let us know if there are ways you think it would be easier for you to join this study.

At the beginning of this study, the WHI staff will ask for your permission to contact your spouse, close relative, or friend for updated information about your health in case you are unable to complete the health update forms. If you move, and we are not able to find you, we may try to locate you through nationally available records, such as social security. If you choose not to join the WHI Extension Study, it will not affect your personal medical care or your medical insurance coverage. The study does not replace your usual medical care.

Voluntary Consent
If you have any questions about any part of the study or your rights as a volunteer, a WHI staff person will answer them before you sign this consent form. Also, if you are not satisfied with the manner in which this study is being conducted, or if you have any questions about your rights as a study participant, please call [IRB Official’s Name] in the Institutional Review Board Office of [Clinical Center’s Institution] at [phone number]. If you have any questions at any time, you may call: [Clinical Center name and phone number] or any of the investigators listed at the beginning of this form. Before you sign this form, please ask any questions on any part of this study that is not clear to you.

Investigator's Statement
I have provided an explanation of the above research program. The participant was given an opportunity to discuss the procedures, including risks, and to ask any additional questions. A signed and dated copy of the consent form has been given to the participant.

__________________________ ____________
Signature of Principal Investigator or Designee Date
PARTICIPANT’S STATEMENT

I certify that I have read, or had read to me the description of the WHI Extension Study. I understand this information and voluntarily consent to join. I understand that I may quit the study at any time. I have had a chance to ask questions about the study and my participation and about the need for access to my medical records. They have been answered to my satisfaction. I understand that future questions I may have about the research will be answered by one of the investigators listed above and that any questions I have about my rights as a research subject will be answered by the person identified above. I acknowledge that I will receive a signed copy of this consent form for my records.

___________________________________ ____________
Signature of Participant Date

___________________________________ ____________
Signature of Witness Date