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TITLE: A Translational Approach to Validate in Vivo Anti-tumor Effects of Chloroquine on Breast Cancer Risk

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CONTRACTING ORGANIZATION: Baylor College of Medicine Houston, Texas 77030-3498

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13. SUPPLEMENTARY NOTES

14. ABSTRACT

SUBJECT & PURPOSE: This translational epidemiologic study conducted online aims to confirm preclinical data on the chemopreventive potential of chloroquine (aminoquinoline), a well-characterized anti-malarial drug. BACKGROUND: Exposure to chloroquine, an off-patent anti-malarial drug with a 60-year history of use by millions, reduces the incidence of breast cancer in genetically programmed rats by 37%. METHODS & SCOPE: About 65% of Peace Corps volunteers received chloroquine prophylactically between 1965 and 1990. Therefore, we will collect chloroquine exposure, breast cancer risk, and breast cancer diagnosis data from returned volunteers who served during this period through an online application. We will characterize participants into chloroquine exposed and unexposed groups, based on country of service and self-reported exposure status. The cost and time efficiencies afforded by this study design will allow the translation of preclinical data on breast cancer chemoprevention into public health and potentially promote the repositioning of a well-characterized and inexpensive drug.

15. SUBJECT TERMS

Breast cancer, chloroquine, chemoprophylaxis

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INTRODUCTION

SUBJECT & PURPOSE: This translational epidemiologic study aims to confirm preclinical data on the chemopreventive potential of chloroquine (aminoquinoline), a well-characterized anti-malarial drug. BACKGROUND: Exposure to chloroquine, an off-patent anti-malarial drug with a 60-year history of use by millions, reduces the incidence of breast cancer in genetically programmed rats by 37%. METHODS & SCOPE: About 65% of Peace Corps volunteers received chloroquine prophylactically between 1965 and 1990. Therefore, we will collect chloroquine exposure, breast cancer risk, and breast cancer diagnosis data from returned volunteers who served during this period through an online application. We will characterize participants into chloroquine exposed and unexposed groups, based on country of service and self-reported exposure status. The cost and time efficiencies afforded by this study design will allow the translation of preclinical data on breast cancer chemoprevention into public health and potentially promote the repositioning of a well-characterized and inexpensive drug.

BODY

The following is an update on tasks outlined in the proposed Statement of Work.

Task 1. Obtain human subjects regulatory approval of study (months 1-4)

Completed in Year 1. The human subjects regulatory approval is complete for the study. Baylor College of Medicine's IRB (Appendix A) and Human Research Protection Office (HRPO) Office of Research Protections (ORP) U.S. Army Medical Research and Materiel Command (USAMRMC) have approved the human subjects application.

IRB renewal submission will occur on July 14, 2014. We will request renewal for one year.

Task 2. Create a breast cancer survey instrument (month 1-5)

Completed in Year 1. Participant workflow documents have been developed (Appendix B), and a questionnaire has been written (Appendix C). We completed development of the application to collect the data using a web-based participant interface in Year 2 and launched the application in April 2014.

Task 3. Data Management System (months 1-18)

Year 1 milestones. No existing system of which we are aware has all of the technical features this project requires, nor the ability to meet its statistical sampling needs. During year 1 of funding, we finalized technical documents, architected the data model for the database, and developed CorpsChronicles, a secure, database-backed web application developed to manage large, socially oriented epidemiologic studies. CorpsChronicles will be used to collect information from RPCVs to determine if chloroquine exposure in this cohort associates with reduced breast cancer incidence compared with the general US population.

CorpsChronicles was designed to (i) allow research subjects to securely answer dynamic questionnaires; (ii) manage electronic recruitment and referral workflows; (iii) send email reminders to participants to complete surveys; and (iv) provide feedback to end users on the overall progress of the study in a way that does not compromise the results. Because breast cancer known risk factors include family history, race, and lifestyle, the online survey elicits self-reported data on potential confounding variables which will be accounted for in the statistical analysis to maximize internal validity. Two hundred RPCVs were selected as initial seeds, with a goal of up to 18,000 respondents across all chains.

Because CorpsChronicles is a complex system that must support myriad workflows, be easy-to-use by a diverse body of respondents, be able to accommodate hundreds of simultaneous end users, and support usage outside of the primary intended workflows, we elected to be as cautious as possible in designing and testing the system. This choice was made because (i) any major issues not caught in the planning and testing phases may cause end users to not complete their surveys, (ii) these same issues would likely reduce the number of respondents making the study fall potentially short of its target accrual and (iii) we anticipate that accruals will be extremely rapid owing to the high level of enthusiasm from RPCVs for the project.

Year 2 milestones. During Year 2, we user-tested the CorpsChronicles application in (i) internal testing with Baylor faculty and staff and (ii) in think-aloud testing (Olmsted-Hawala and Murphy, 2013. Protocol and instrument attached in Appendices.) with five Returned Peace Corps Volunteers from different age cohorts from the study population who had different levels of familiarity with online applications, different experiences with breast cancer and, most importantly, no prior knowledge of how the system should function, thus making them excellent real-world test cases. Based on the feedback from this testing, we fixed bugs and revised the system to make it more responsive and user-friendly. Throughout Year 2 and in Year 3, in accordance with iterative software design, our application developer continues to develop iterative revisions of the system to refine existing features, add new features and address issues that arise through production use. This iterative design is informed by a database of participant questions and comments through the study toll-free telephone number and email help line (See Appendix).

Task 4. Recruit a cohort of 14,000 to 18,000 female returned Peace Corps volunteers who served from 1961 through 1990 in both malaria-endemic and non-endemic parts of the world and data collection (months 3-18)

Year 1 milestones. We created a database of eligible potential volunteers who have expressed interest in participating in BCM research from an IRB-approved study "CREATING A DATABASE OF POTENTIAL PARTICIPANTS," in which RPCVs were recruited at the Peace Corps 50th Anniversary conference in Washington DC in September 2012 and a gathering of returned volunteers in June 2013 in Minneapolis, MN.

In Year 2, this database served as the source of "seeds" for Respondent Driven Sampling. In April 2014, we began data collection by emailing all potential volunteers in the database an invitation to participate in the study with link to the online survey instrument with an embedded number to identify each responder. We sent the seeds a reminder email before sending them the link, to help them remember the study and their willingness to play a role in it.

Simultaneously, we launched an informational campaign in concert with the National Peace Corps Association (NPCA). This campaign includes: websites and blogs, including PeaceCorpsConnect.org. PeaceCorpsWorldwide.org, and the websites of independent and NPCA-affiliated RPCV member groups; NPCA's monthly e-newsletter with 35,008 subscribers; mass emails or listsery posts to membership of NPCA, Peace Corps Worldwide, and independent and NPCA-affiliated RPCV member groups; and social media tools including Facebook.

Interim analysis performed at one month post-activation to ensure appropriate accruals with the sampling method, Respondent Driven Sampling (RDS), was encouraging enough to continue with the planned RDS methodology. We will perform a second interim analysis in July 2014. If subject accrual is not meeting our goals at that time, we will implement our back-up plan, traditional snowball sampling.

Enrollment

TABLE 1: ENROLLMENT, FULL STUDY, AS OF MAY 23, 2014

Race/Ethnicity	Female
All ethnicities	71
American Indian or Alaskan Native	0
Asian/Non Vietnamese	0
Black of African American	0
Hispanic or Latino	0
Mixed Race or Ethnicity	0
Native Hawaiian or Pacific Islander	0
Vietnamese	0
White	70
Unknown	1

TABLE 2: ENROLLMENT, THINK-ALOUD PILOT

Race/Ethnicity	Female
All ethnicities	5
American Indian or Alaskan Native	0
Asian/Non Vietnamese	0
Black of African American	0
Hispanic or Latino	1
Mixed Race or Ethnicity	0
Native Hawaiian or Pacific Islander	0
Vietnamese	0
White	4

Task 5. Identify the cause of death of deceased female returned Peace Corps volunteers who served from 1961 through 1990 (months 6-18)

Year 1 milestones. In the original protocol and grant applications, we proposed using the National Death Index (NDI) to match the names of the deceased provided by study participants to those in the NDI to verify causes of death. However, it was determined that relying on participants to provide accurate identifying information on those potential participants who have died would yield an unrepresentative sample of those who died and may be upsetting to participants. Therefore, Network Scale-Up Method (NSUM) will be used to estimate the number of deceased RPCVs, eliminating both the need to ask for identifying information of those who have died from participants and the use of the National Death Index.

NSUM estimates the size of a hidden or hard-to-reach population (i.e., deceased RPCVs) by assuming that, in general, people's social networks are representative of the general population in which we live and move (HR Bernard, 2010). Using the questions already crafted for our primary sampling mechanism of RDS, we will ask the respondents one additional question to determine the number of female RPCVs they know who have died. This number, along with the number of female RPCVs each respondent knew and the total number of female Peace Corps Volunteers who served between 1961 and 1990, will be used to estimate the number of deceased women. Implementation of NSUM eliminates the need for NDI, and thus we request that the funds intended for NDI be re-distributed.

Years 2 and 3. Analysis will begin once data collection is completed.

<u>Task 6</u>. Examine the association between chloroquine exposure and breast cancer in our cohort of RPCVs who served between 1961 and 1990 (months 18-21)

Task 6 will begin in Year 3 of the study (no-cost extension).

Task 7. Disseminate findings (months 21-24)

We have presented posters on this project at two conferences and presented an oral presentation about the project at one conference. A third oral presentation is planned for June 2014. See Reportable Outcomes section for references and Appendices for posters and manuscripts.

KEY RESEARCH ACCOMPLISHMENTS

- Year 1: Development of a Java Enterprise Edition 6 application that uses the model-view-controller architecture with the JBoss Seam framework to manage its various components. The application runs within JBoss 7 middleware and utilizes Oracle 11g for data persistence, though Hibernate makes it possible to swap out Oracle for other database management systems. Security is managed by enhanced Seam security, encrypted passwords, role-based permissions, Captchas, and unique "coupon" codes that serve as keys for respondents. Referrals of respondents are based on a novel social network sampling methodology, Respondent-Driven Sampling (RDS), in which initial respondents ("seeds") identify members of their social network who may want to participate and these members in turn identify members of their network thus creating network "chains" for each seed. Data from links in each chain are compared back to their seed data using RDSAT freeware; demographic and clinical characteristic data is compared between groups using chi-square or Fisher's exact test and t-test or Wilcoxon rank-sums. Multivariate logistic regression is used to assess associations between chloroquine exposure and breast cancer incidence, with the Network Scale-Up Method, a social networking method, applied to estimate the hard-to-count population of RPCVs.
- Year 2: Survey launched.

REPORTABLE OUTCOMES

Publications:

Mayfield P, McOwiti A, Barney J, Qu Shijing, Sexton KR, Harris AM, McArthur K, Hilsenbeck SG, Bondy ML, Conneely O, Spitz M, Dacso CC, Becnel LB. CorpsChronicles: An Application Leveraging Social Networks for Online Epidemiologic Survey Studies. Submitted to American Medical Informatics Annual Symposium Washington, DC. November 16-20, 2013. (Appendix F)

Sexton K, Harris AM, McArthur K, Bondy M, Hilsenbeck S, Becnel L, Mayfield P, Conneely O, Spitz M, Queen C, Dacso C. Designing a translational epidemiologic study: Chloroquine and breast cancer chemoprevention in Returned Peace Corps Volunteers. 140th APHA Annual Meeting, San Francisco, CA. October 2012. (Appendix G)

Annual Report: A Translational Approach to Validate in-Vivo Anti-tumor Effects of Chloroquine on Breast Cancer Risk

Mayfield PK, McOwiti AM, Barney J, Darlington Y, Vanjani G, Becnel LB. C3PR, CODR & Corps Chronicles: case studies of clinical and epidemiological research databases within the DLDCC. Dan L. Duncan Cancer Center 2012 Symposium. Baylor College of Medicine. Houston, TX. (Appendix H)

O'Brien J. Chloroquine and breast cancer chemoprevention in Returned Peace Corps Volunteers: Study Progress and Future Steps. National Peace Corps Association 2104 Annual Gathering. Nashville, TN.

Experience/training supported by this award: One undergraduate student, Adesola Oyewole, University of Houston, received summer training in clinical research, June – August 2012.

CONCLUSION

We have begun to accrue subjects and gather data, but we have not yet analyzed the results. If Respondent Driven Sampling fails to meet our recruitment goals, we will implement our alternative plan: traditional snowball sampling from the National Peace Corps mailing list of over 35,000 people.

REFERENCES

Olmsted-Hawala E, Murphy B. Think-aloud protocols for website usability testing. Wednesday, January 23, 2013. http://www.howto.gov/training/classes/think-aloud-protocols-for-usability-testing.

Bernard HR, Hallett T, Iovita A, et al. Counting hard-to-count populations: the network scale-up method for public health. Sex Trans Inf. 86(Suppl 2): ii11-ii15, 2010.

APPENDICES

- A. Baylor College of Medicine IRB Approval
- B. Participant Workflow Documents
- C. Questionnaire Text
- D. Screenshots of the CorpsChronicles Application
- E. Study Description Developed with the NPCA
- F. Mayfield P, McOwiti A, Barney J, Qu Shijing, Sexton KR, Harris AM, McArthur K, Hilsenbeck SG, Bondy ML, Conneely O, Spitz M, Dacso CC, Becnel LB. CorpsChronicles: An Application Leveraging Social Networks for Online Epidemiologic Survey Studies. Submitted to American Medical Informatics Annual Symposium Washington, DC. November 16-20, 2013.
- G. Sexton K, Harris AM, McArthur K, Bondy M, Hilsenbeck S, Becnel L, Mayfield P, Conneely O, Spitz M, Queen C, Dacso C. Designing a translational epidemiologic study: Chloroquine and breast cancer chemoprevention in Returned Peace Corps Volunteers. 140th APHA Annual Meeting, San Francisco, CA. October 2012.
- H. Mayfield, PK, McOwiti, AM, Barney, J, Darlington, Y, Vanjani, G, and LB Becnel. C3PR, CODR & Corps Chronicles: Case Studies of Clinical and Epidemiological Research Databases within the DLDCC. Dan L. Duncan Cancer Center 2012 Symposium. Baylor College of Medicine. Houston, TX.
- I. Think Aloud Testing Protocol and Instrument
- J. Toll-Free Telephone Number and Email Help Line Statistics



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H-31160 - A TRANSLATIONAL APPROACH TO VALIDATE IN-VIVO ANTI-TUMOR EFFECTS OF CHLOROQUINE ON BREAST CANCER RISK

APPROVAL VALID FROM 1/23/2013 TO 7/2/2013

Dear Dr. DACSO

The Institutional Review Board for Human Subject Research for Baylor College of Medicine and Affiliated Hospitals (BCM IRB) is pleased to inform you that the research protocol named above was approved.

The study may not continue after the approval period without additional IRB review and approval for continuation. You will receive an email renewal reminder notice prior to study expiration; however, it is your responsibility to assure that this study is not conducted beyond the expiration date.

Please be aware that only IRB-approved informed consent forms may be used when written informed consent is required.

Any changes in study or informed consent procedure must receive review and approval prior to implementation unless the change is necessary for the safety of subjects. In addition, you must inform the IRB of adverse events encountered during the study or of any new and significant information that may impact a research participants' safety or willingness to continue in your study.

The BCM IRB is organized, operates, and is registered with the United States Office for Human Research Protections according to the regulations codified in the United States Code of Federal Regulations at 45 CFR 46 and 21 CFR 56. The BCM IRB operates under the BCM Federal Wide Assurance No. 00000286, as well as those of hospitals and institutions affiliated with the College.

Sincerely yours,

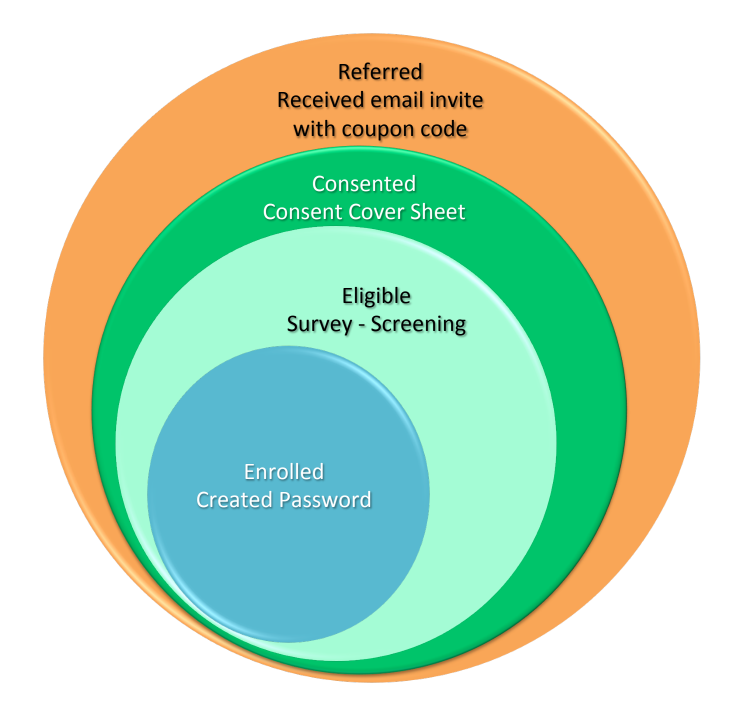
DANIELLE R SORELLE,

Danielle Excelle

Institutional Review Board for Baylor College of Medicine and Affiliated Hospitals



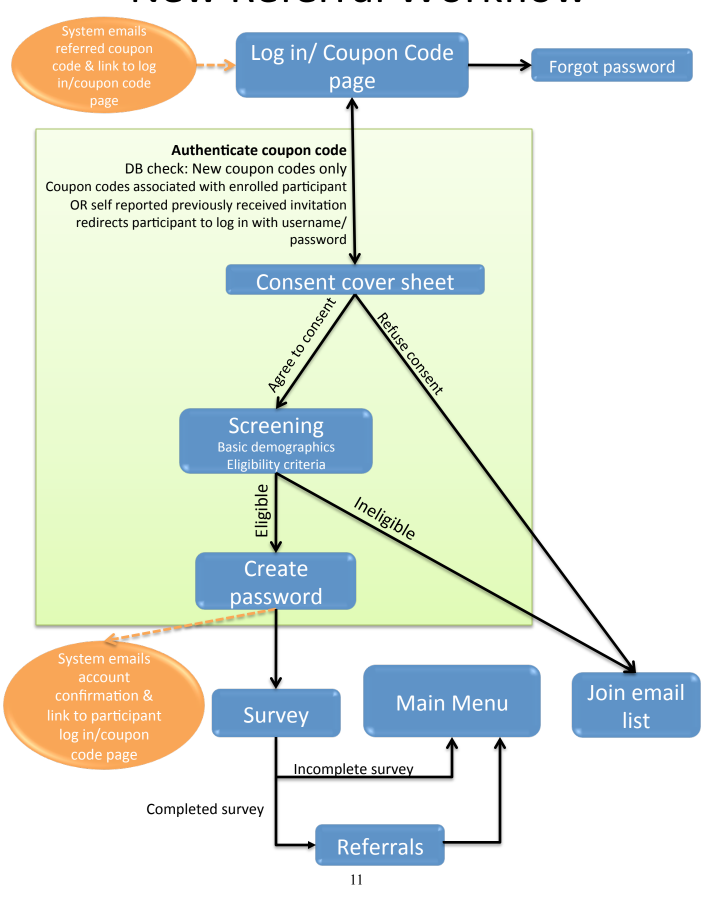
Enrollment Process



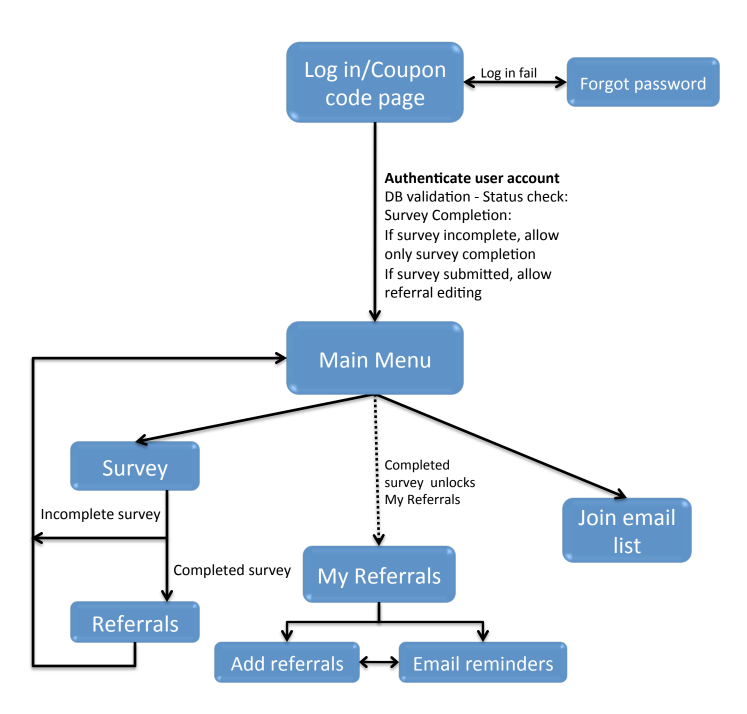
Public Website Workflow



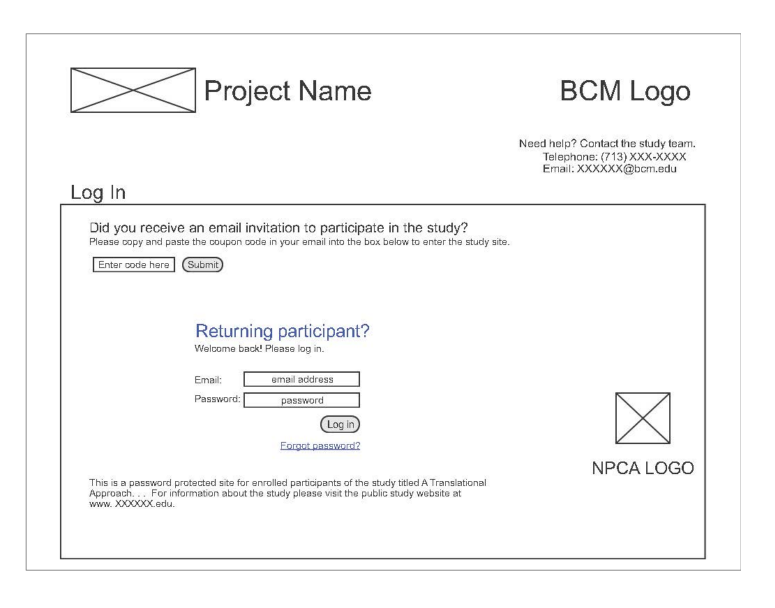
New Referral Workflow



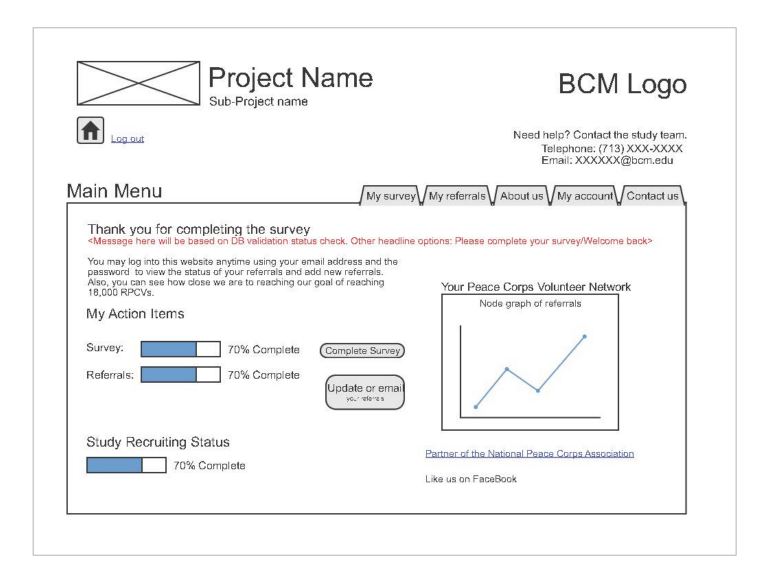
Returning Participants Workflow



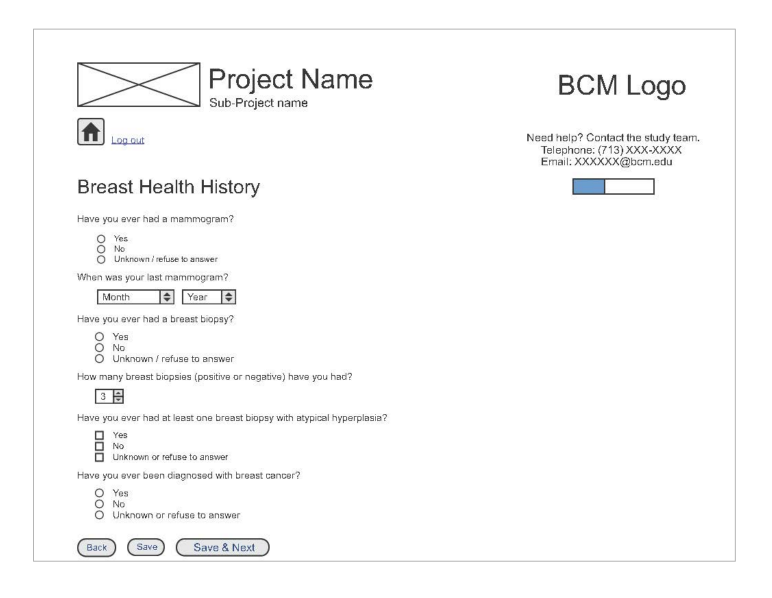
Sample Screen: Log in/Coupon Code page



Sample Screen: Main Menu



Sample Screen: Survey



Email Notifications

Study Site

- 1. Invitation to Participate (Seeds and Referrals)
- 2. Account Creation Confirmation (Seeds and Referrals)
- 3. Reminder to Start Survey (Seeds and Referrals)
- 4. Reminder to Complete Survey (Seeds and Referrals)
- 5. Reminder to Start Referrals (Seeds and Referrals)
- 6. Reminder to Complete Referrals (Seeds and Referrals)

Public Site

1. Mailing List Confirmation

Questionnaire used in CorpsChronicles

About You <*Screening>* 1. Name* ______ MIDDLE 2. Email address* _____ 3. Confirm email address* 4. May we contact you if we have further questions about your responses? ☐ Yes ☐ No 6. Place of birth City _____ State ____ or Check box if not born in the United States If foreign-born, list country of birth: Race (Check all that you MOST CLOSELY identify): ☐ American Indian or Alaska Native ☐ Asian ☐ Black or African-American ☐ Native Hawaiian or Other Pacific Islander ☐ White ☐ Unknown or refuse to answer 8. Ethnicity (Check one that you MOST CLOSELY identify): ☐ Hispanic or Latino ☐ Not Hispanic or Latino ☐ Unknown or refuse to answer 9. Sex* ☐ Male

☐ Female

Your Peace Corps Service

10. Country of Service <drop -="" _<drop="" al="" dec="" down="" down<drop="" jan="" list="" month="" of="" service:="" years="">_/<dropdo< th=""><th>Jan – Dec>_ /<dropdown 1961-2015="" list=""></dropdown></th></dropdo<></drop>	Jan – Dec>_ / <dropdown 1961-2015="" list=""></dropdown>
*Require response on years of service (M	conth of service not required.)
BUTTON: Add another country/ year	of service
*Repeat these questions as many times as	the RPCV has tours of service.
Are you currently active duty military person Graph Yes Graph No	nel?*

Create Password

Please create a password for this survey. With your email address and password, you will be able to log into this website to finish your incomplete survey, add RPCV referrals and see the status of previously submitted referrals, and view study updates.

Your username is the email address at which you received your emailed survey invitation. Please create a password below.

Password Confirm password

Your RPCV Social Network

The following questions are about your fellow volunteers in the Peace Corps. For the purposes of this study, the definition of knowing someone is that you know them and they know you by sight or by name and that you could contact them.

11.	Approximately how many women do you know who served in the Peace Corps at any time between 1961 and 1990?
	women
12.	Of those women you know/knew in the Peace Corps during that time period, about how many of them served at the same time and same country you served?
	women
13.	Of those women you know/knew in the Peace Corps during that time period, about how many of them did you meet after you returned home?
	women
14.	Of those women you know/knew in the Peace Corps during that time period, about how many of them have died since you've returned home?
	women

Your History of Medications to Treat or Prevent Malaria

Chloroquine is a drug that is taken to prevent or treat malaria. When it is taken to prevent malaria it is taken once a week. Other names for chloroquine are: Chloroquine phosphate oral and Aralen phosphate. Malarone, Lariam, Paludrine, and Doxycycline are other drugs that are sometimes used for malaria prevention. They are not the same as Chloroquine.

15.	Have you eve	er taken a drug to prev	vent or treat mal	aria at any tin	ne in your life?	
	☐ Yes ☐ No ☐ Unknown					
		complete the box belo during your Peace Co				
	Drug Name	Peace Corps Use?	Country where Taken	Reason	Approximate Time Taken	How Ofter Drug was Taken
-	Drop Down (include Do Not Know and Other)	Y/N	Drop Down (Include Do Not Know and Other)	Prevention Treatment	Mm/yy- mm/yy	Daily Weekly Bi-Weekly Other
Dio XX	XXX.	A DRUG Tug that is not in this literate taken chloroquine		, and the second	oordinator at XX	XX-XXX-
		o Question 18. Skip to Question 18	<u>3.</u>			
17	'. If you took c	chloroquine for anothe	er reason, please	complete the	following:	
	Reason for U					
<d1< td=""><td></td><td>e Time Taken: MM> <dropdowns fo<="" td=""><td>or YYYY> - <dr< td=""><td>opdowns for N</td><td>MM> <dropdov< td=""><td>vns for</td></dropdov<></td></dr<></td></dropdowns></td></d1<>		e Time Taken: MM> <dropdowns fo<="" td=""><td>or YYYY> - <dr< td=""><td>opdowns for N</td><td>MM> <dropdov< td=""><td>vns for</td></dropdov<></td></dr<></td></dropdowns>	or YYYY> - <dr< td=""><td>opdowns for N</td><td>MM> <dropdov< td=""><td>vns for</td></dropdov<></td></dr<>	opdowns for N	MM> <dropdov< td=""><td>vns for</td></dropdov<>	vns for
ΥY	YYY>Menstrua	al History				

The next set of questions is about your menstrual periods.

18. How old were you when you started your first menstrual period?
years
19. Have you reached menopause, that is your periods have stopped for 12 months or longer for reasons other than pregnancy, or you have no uterus?
 ☐ Yes ☐ No, but periods are irregular Skip to Question 22. ☐ No Skip to Question 22. ☐ Refuse to Answer Skip to Question 22
20. How old were you when your periods stopped?
years or
21. Why did your periods stop?
□ Natural menopause □ Surgical menopause □ Chemotherapy or radiation treatment □ Do not know □ Other

Pregnancy History

babies?

IF YES:

next question.

Select one option. If No, go on to the

How long did you breastfeed?

8 7 7					
Have you ever been pregnant? Please include live births, stillbirths, tubal or ectopic, miscarriage / termination.					
☐ Yes ☐ No Skip to Question 25. ☐ Refuse to answer Skip to Que	estion 25.				
	ve you had, including all live births, still births and tubal or erminations? Include current pregnancy if applicable.				
pregnancies					
24. Are you currently pregnant?					
☐ Yes ☐ No ☐ Unknown or refuse to answer					
	Your pregnancies. Please include all pregnancies (live miscarriages/terminations), and all sons/daughters whether he information for each pregnancy.				
Repeat section for each pregnancy	Pregnancy # (Please indicate pregnancy number)				
How old were you when you became pregnant?	years Don't recall/ refuse to answer				
What was the outcome of this pregnancy? Circle one option	□ Full-term live birth □ Preterm live birth □ Stillbirth (>5 months) □ Miscarriage(<5 months)□ Induced/Elective abortion □ Refuse to answer				

	Oon't recall	

Breast Health History

25. Have you ever had a mammogram?
☐ Yes ☐ No Skip to Question 27. ☐ Unknown or refuse to answer Skip to Question 27.
26. When was your last mammogram?
//
27. Have you ever had a breast biopsy?
☐ Yes ☐ No Skip to Question 30. ☐ Unknown or refuse to answer Skip to Question 30.
28. How many breast biopsies (positive or negative) have you had?
biopsies
29. Have you ever had at least one breast biopsy with atypical hyperplasia?
☐ Yes ☐ No ☐ Unknown or refuse to answer
30. Have you ever been diagnosed with breast cancer?
☐ Yes ☐ No Skip to Question 39. ☐ Unknown or refuse to answer Skip to Question 39.

Breast Cancer Diagnosis <to be completed only if Question 30 is "Yes">

31. Approximate Date of Diagnosis: <dropdown> / <dropdown> MM YYYY</dropdown></dropdown>
32. What imaging technique was used to detect the breast mass? (Check all that apply)
☐ Mammogram ☐ Ultrasound ☐ MRI ☐ Other:
The following questions are regarding markers that are important in your breast cancer treatment
33. ER (estrogen receptor) status:
☐ Positive ☐ Negative ☐ Unknown
34. PR (progesterone receptor) status:
☐ Positive ☐ Negative ☐ Unknown
35. HER2 (human epidermal growth factor receptor 2) status:
☐ Positive ☐ Negative ☐ Unknown
36. How was your breast cancer treated? (Check all that apply)
 □ Mastectomy □ Lumpectomy □ Radiation Therapy □ Chemotherapy □ Hormonal Therapy, such as tamoxifen or aromatase inhibitors (Arimidex/anastrozole, Aromasin/exemestane, Femara/letrozole) □ Targeted therapy, such as Herceptin/trastuzumab, Avastin/bevacizumab □ Other: □ Unknown
37. Approximate Height at Diagnosis: feet inches
38 Approximate Weight at Diagnosis: nounds

Tobacco History

39. Have you smoked at least 100 cigarettes (5 packs) in your lifetime?
☐ Yes ☐ No, never Skip to Question 42. ☐ Unknown or refuse to answer Skip to Question 42.
40. Do you now smoke cigarettes?
☐ Not at all ☐ Some days ☐ Everyday
41. On average, how many packs do/did you smoke per day?
packs
Alcohol History The following questions are about your consumption of alcoholic drinks. These include been wine, wine coolers, and liquor (cocktails, whiskey, tequila, gin, rum, etc).
42. In your entire life, have you had at least a total of 12 drinks of any type of alcoholic beverage?
☐ Yes ☐ No Skip to Question 45. ☐ Unknown or refuse to answer Skip to Question 45.
43. Approximately how many drinks per week do you consume?
☐ 1-3 ☐ More than 3 ☐ Unknown or refuse to answer
44. How many years have you consumed alcohol?
years Check box if you do not know or refuse to answer

Family History of Cancer

The next section is about the cancer history of members of your family. These include living and deceased family members. Please do not include half-siblings.

45. Are you ad	opted?
☐ Yes S☐ No	kip to Question 47
☐ Unknov	wn or refuse to answer Skip to Question 47
46. Have any o with cancer	f your immediate blood relatives (parents, siblings, children) ever been diagnosed?
_	Skip to Question 47. where the state of the

For each of these family members, please tell us which of the above cancers they have/had and the approximate age of their diagnosis. Select "DK" if Don't Know.

Family Member	Age Diagnosed	Type of Cancer
Drop Down	Number/DK	Drop Down

Birth Control

The next section is about the use of birth control pills and hormonal contraceptives. These are taken for various reasons, including preventing pregnancy, irregular periods, etc.

-	-	her hormonal contraceptives for at least one injections, implants, and patches.		
☐ Yes, I am currently usi☐ Yes, I used in the past☐ Unknown Skip to Que☐ Refuse to answer Ski	(more than 6 morestion 50.	nths)□ No Skip to Question 50.		
48. How old were you when you started taking birth control pills/hormonal contraceptives?				
☐ Less than 30 ☐ 30-39				
40-49				
☐ 50-59 ☐ 60 or older				
49. What type of contraceptives did/do you use?				
Туре	Ever Taken?	Total number of years OR total number of months		
Birth control pills	☐ Currently ☐ Formerly			

Туре	Ever Taken?	Total number of years OR total number of months
Birth control pills	☐ Currently ☐ Formerly ☐ No	
Birth control injections (ex. Depo Provera)	☐ Currently ☐ Formerly ☐ No	
Birth control implants (ex. Norplant)	☐ Currently ☐ Formerly ☐ No	
Birth control patch (ex. Ortho Evra)	☐ Currently ☐ Formerly ☐ No	
Vaginal ring (ex. Nuva Ring)	☐ Currently ☐ Formerly ☐ No	

Hormone Replacement Therapy (To be answered only if Question 19 is "Yes")

The next section is about use of female hormones. These are often given to relieve menopausal symptoms such as hot flashes. These hormones may have been in the form of pills, shots, skin patches, creams, or vaginal suppositories.

50. Have you ever used hormone replacement therapy such as estrogen or progesterone? These include pills, shots, skin patches, creams, or vaginal suppositories. Do not include birth control pills or fertility drugs.
☐ Yes, currently use (within past 6 months) ☐ Yes, used in the past (longer than 6 months ago) ☐ No, never used Skip to End. ☐ Unknown Skip to End. ☐ Refuse to answer Skip to End.
51. In total, how long did you use replacement hormones? (years or months)
years or months
52. What type of hormone replacement therapy do/did you use? (Please complete table below)

Туре	Ever taken?	Total number of years <u>OR</u> total number of months
Pills containing estrogen only (Ex. Premarin, Estrace, Estratest, Ogen)	☐ Currently ☐ Formerly ☐ No	
Pills containing progestin only (Ex. Provera, Cycrin, MPA)	☐ Currently ☐ Formerly ☐ No	
Pills containing estrogen plus progestin (Ex. Prempro, Premphase, Femhrt)	☐ Currently ☐ Formerly ☐ No	
Patches containing estrogen only	☐ Currently ☐ Formerly ☐ No	
Patches containing estrogen plus progestin (Ex: Combipatch)	☐ Currently ☐ Formerly ☐ No	— ☐ Years ☐ Months ☐ Don't recall
Type - Unrecalled	☐ Currently ☐ Formerly ☐ No	— ☐ Years ☐ Months ☐ Don't recall
Other:	☐ Currently ☐ Formerly ☐ No	

My Referrals

RPCV Referral #1

In order to reach as many female RPCVs as possible, we are asking that you provide the names and contact information for up to 3 female RPCVs who served between 1961 and 1990.

We will email or mail a letter to each woman you refer to the study with a link to the secure study website and an explanation of the study.

Please fill out as much information as you can on each RPCV so that we may invite them to participate in the study.

Eull Name:				
	FIRST	MIDDLE	LAST	
		a code):		
Served in Peace	Corps <country< td=""><td>drop down> From <ye< td=""><td>ear drop down> to <y< td=""><td>'ear Drop down></td></y<></td></ye<></td></country<>	drop down> From <ye< td=""><td>ear drop down> to <y< td=""><td>'ear Drop down></td></y<></td></ye<>	ear drop down> to <y< td=""><td>'ear Drop down></td></y<>	'ear Drop down>
Sex: Male/Fema	le			
When we contact Yes/no	t this RPCV, may	we please tell her that	you referred her to u	s?
BUTTON: SAV BUTTON: ADI	YE O PERSONAL N	ОТЕ		
RPCV Referral	#2			
Full Name:	FIRST			
		MID		LAST
		a code):		
Served in Peace	Corps <country< td=""><td>drop down> From <ye< td=""><td>ear drop down> to <y< td=""><td>'ear Drop down></td></y<></td></ye<></td></country<>	drop down> From <ye< td=""><td>ear drop down> to <y< td=""><td>'ear Drop down></td></y<></td></ye<>	ear drop down> to <y< td=""><td>'ear Drop down></td></y<>	'ear Drop down>
Sex: Male/Fema	le			
When we contac Yes/no	t this RPCV, may	we please tell her that	you referred her to u	s?
BUTTON: SAV	Æ.			

BUTTON: ADD PERSONAL NOTE

BUTTON: ADD PERSONAL NOTE

RPCV Referral #3		
Full Name:		
FIRST	MIDDLE	LAST
Email Address:		
Telephone Number (including area code	e):	
Served in Peace Corps <country d<="" drop="" td=""><th>down> From <year down="" drop=""> to <</year></th><td>Year Drop down></td></country>	down> From <year down="" drop=""> to <</year>	Year Drop down>
Sex: Male/Female		
When we contact this RPCV, may we pl	lease tell her that you referred her to	us?
Yes/no		
BUTTON: SAVE		

Appendix D: Screenshots of the CorpsChronicles Application

Email Received by Study Participants:

Dear	,

You have been referred by a fellow returned Peace Corps volunteer (RPCV) who thinks you might be interested in participating in an online study designed for RPCVs. This study is being conducted by researchers at Baylor College of Medicine to examine the relationship between certain medications taken by PCVs as part of their service and a reduced risk of health problems, such as cancer, later in life. In order to test this hypothesis, we need up to 18,000 female RPCVs who served between 1961 and 1990 to fill out a secure online questionnaire.

If you are a woman who served in the Peace Corps between 1961 and 1990, you are eligible to participate in this important study. We hope you will consider taking part, and can directly access your survey by clicking here.

Alternatively, you may also copy and paste the survey URL, https://corpschronicles-stg.research.bcm.edu/study/verify, into your web browser then paste in your unique coupon code, 938b4489-8c5c-439b-bf27-f7bad4de43f6, into the survey webpage.

Each coupon code is unique, so forwarding this email will not allow another person to take the survey. You will have an opportunity to refer other RPCVs and automatically create coupon codes for them at the end of your survey through the CorpsChronicles RPCV site.

If you need any assistance with the survey, please call the RPCV study team at (855)-869-0829 or email us at corpsstudyquestion@bcm.edu.

If you do not wish to receive any further emails or calls regarding this study, you may respond to this email with the word REMOVE in the subject line or call us at (855)-869-0829.

Sincerely,

Clifford C. Dacso, MD, MPH

Principal Investigator

'A Translational Approach to Validate in-vivo Anti-tumor Effects of Chloroquine on Breast Cancer Risk'

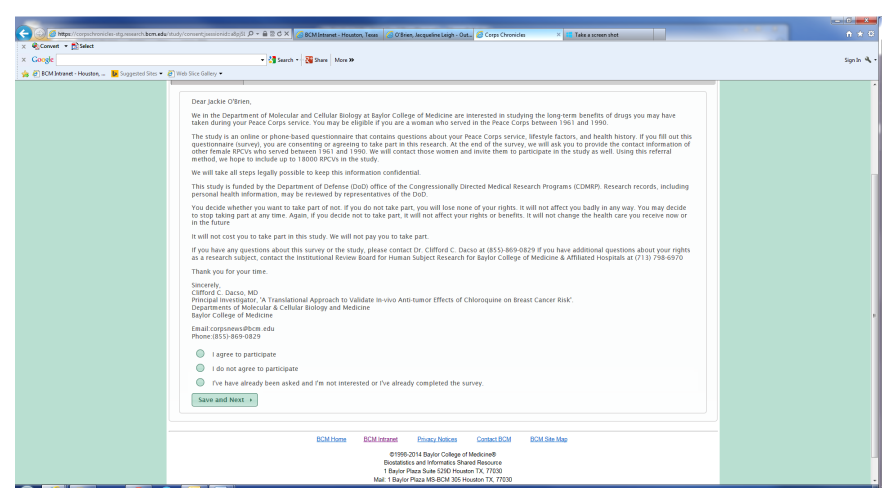
Departments of Molecular & Cellular Biology and Medicine

Baylor College of Medicine

Email: corpsnews@bcm.edu

Phone: (855)-869-0829

First page seen when participant clicks on email:

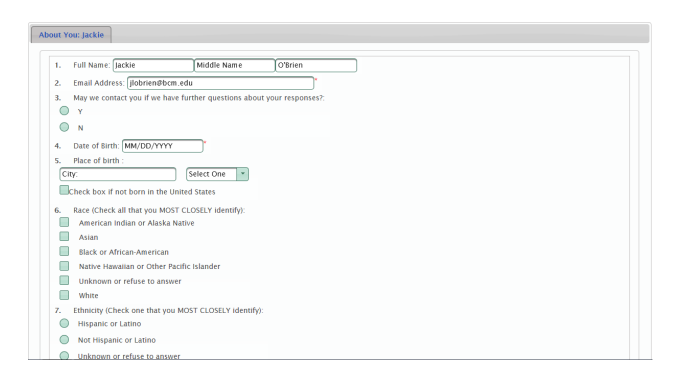


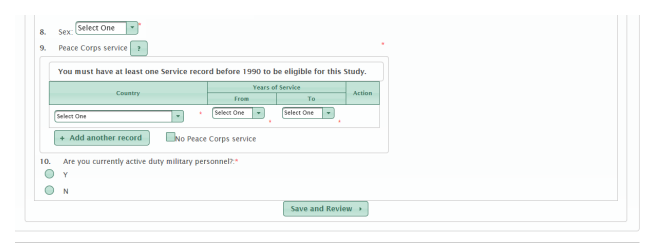
About You:





For Help, get the Quickstart Guide Still need Help? Contact the study team: Call Toll Free: (855)-869-0829 Email: corpsstudyquestion@bcm.edu

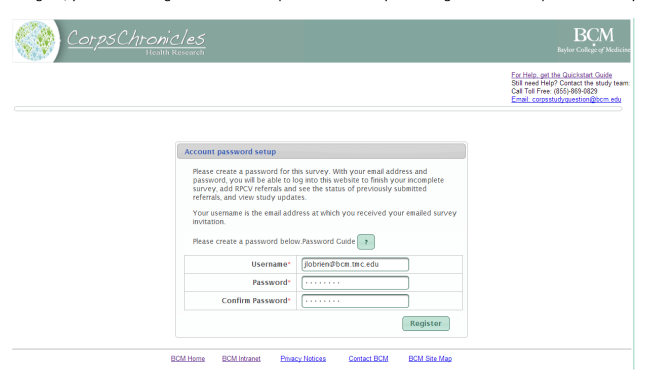




BCM Home BCM Intranet Privacy Notices Contact BCM BCM Site Map

©1998-2014 Baylor College of Medicine®
Biostatistics and Informatics Shared Resource
1 Baylor Plaza Suite 520P Houston TX, 77030
Mail: 1 Baylor Plaza MS-BCM 305 Houston TX, 77030
Phone: 713-798-4155 | Fax: 713-798-6822
E-mail: dldcc-informatics@breastcenter.tmc.edu
Last Modified: 03/19/2014 08:48:15:7800

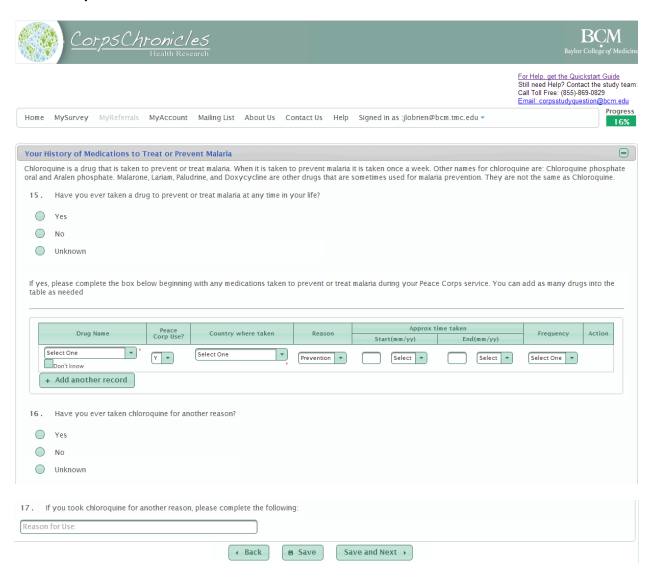
If eligible, you can then register and create a password so that you can log back in to complete the survey and add referrals:



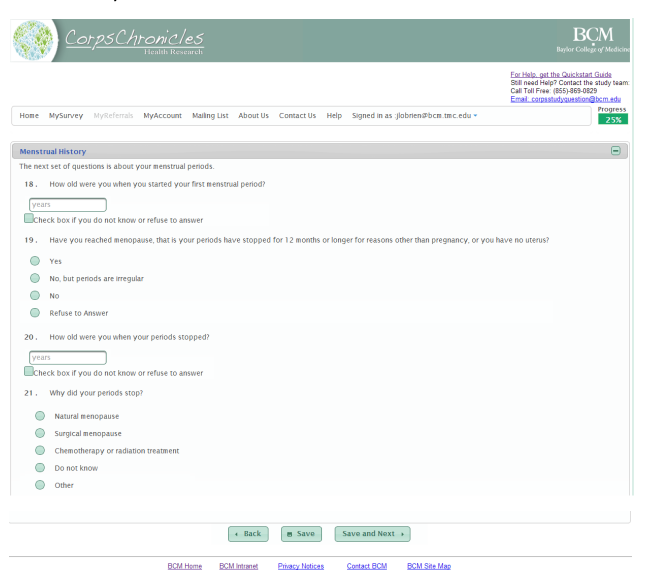
Your RPCV Social Network:



Your History of Medications to Treat or Prevent Malaria:

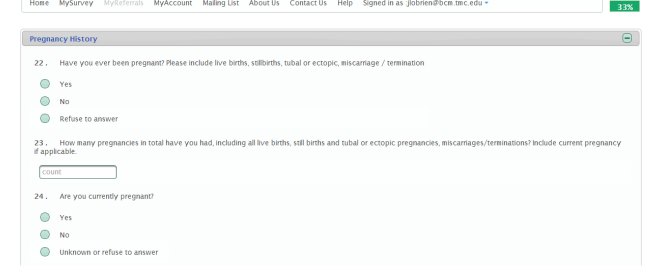


Menstrual History:



Pregnancy History:



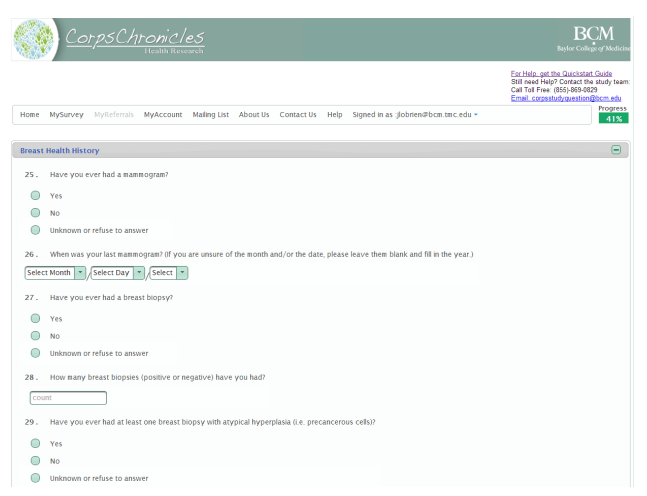


The next set of questions regards all of your pregnancies. Please include all pregnancies (live births, tubal or ectopic, stillbirths, and miscarriages/terminations), and all sons/daughters whether currently living or not. Please fill out the information for each pregnancy.

Pregnancy #	How old were you when you became pregnant?	What was the outcome of this pregnancy? 🦻	Did you breastfeed this or any of these babies?	Action
	Don't recall/ refuse to answer	Full-term live birth Preterm live birth Stillbirth (>5 months) Miscarriage(<5 months) Induced/Elective abortion Refuse to answer	Yes No Don't recall Not applicable	
- Add another r	ecord			

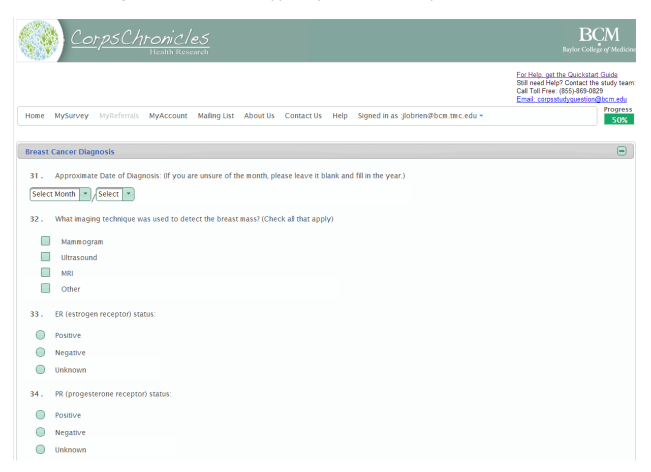
■ Save Save and Next →

Breast Health History:



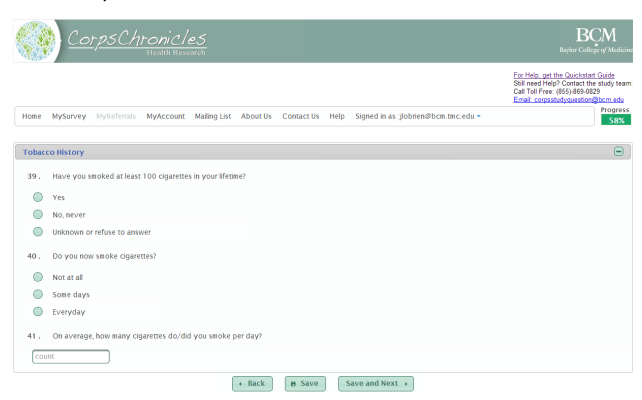


Breast Cancer Diagnosis (This section is skipped if you answer no to question #30):



35.	HER2 (human epidermal growth factor receptor 2) status:
	Positive
	Negative
	Unknown
36 .	How was your breast cancer treated? (Check all that apply)
	Mastectomy
	Lumpectomy
	Radiation Therapy
	Chemotherapy
	Hormonal Therapy, such as tamoxifen or aromatase inhibitors (Arimidex/anastrozole, Aromasin/exemestane, Femara/letrozole)
	Targeted therapy, such as Herceptin/trastuzumab, Avastin/bevacizumab
	Other
	Unknown
37.	Approximate Height at Diagnosis:
feet	
inch	291
38.	Approximate Weight at Diagnosis:
pou	unds
	← Back ← Save and Next →

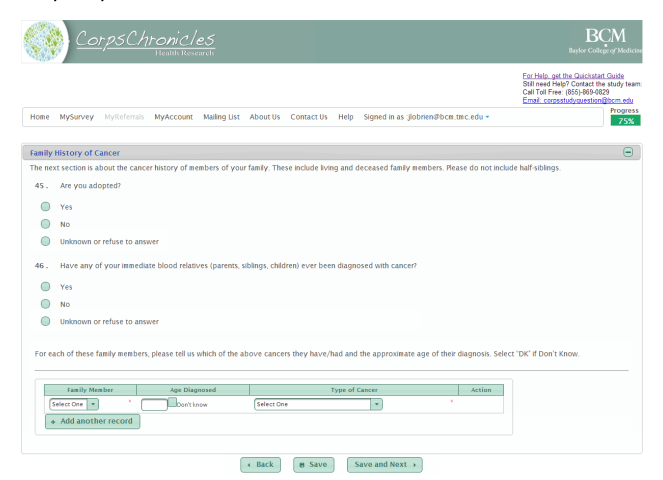
Tobacco History:



Alcohol History:



Family History of Cancer:



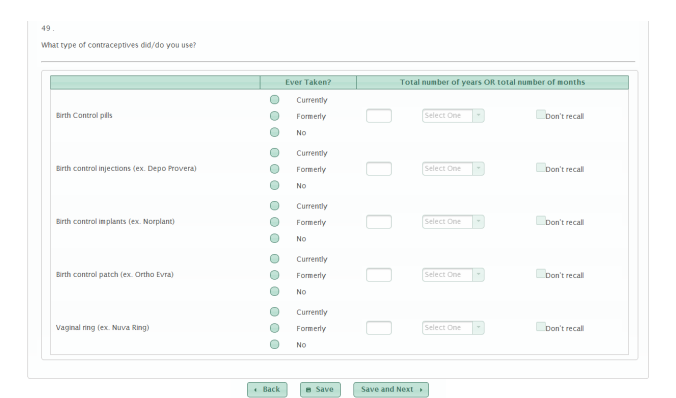
Birth Control:

60 or older



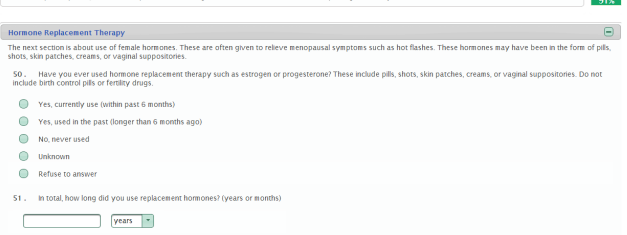
For Help, get the Quickstart Guide
Still need Help? Contact the study team:
Call Toll Free: (855)-869-0829
Email: corpsstudyquestion@bcm.edu

Progress Home MySurvey MyReferrals MyAccount Mailing List About Us Contact Us Help Signed in as :Jlobrien@bcm.tmc.edu • 83% Birth Control The next section is about the use of birth control pills and hormonal contraceptives. These are taken for various reasons, including preventing pregnancy, irregular periods, etc. 47. Have you ever taken birth control pills or other hormonal contraceptives for at least one month for any reason? These include pills, injections, implants, and patches. Yes, I am currently using (within 6 months) Yes, I used in the past (more than 6 months) O No Unknown Refuse to answer 48. How old were you when you started taking birth control pills/hormonal contraceptives? Less than 30 30-39 40-49 50-59



Hormone Replacement Therapy:





What type of hormone replacement therapy do/did you use? (Please complete table below) E Pills containing estrogen only (Ex. Premarin, Estrace, Estratest, Ogen)

	Ever Taken? Total number of years OR total number of month
Pills containing estrogen only (Ex. Premarin, Estrace, Estratest, Ogen)	Currently Formerly Select One Don't recall
Pills containing progestin only(Ex. Provera, Cycrin, MPA)	Currently Formerly No
Pills containing estrogen plus progestin(Ex. Prempro, Premphase, Femhrt)	Currently Formerly Select One Don't recall
atches containing estrogen only	Currently Formerly Select One Don't recall
ratches containing estrogen plus progestin (Ex: Combipatch)	Currently Formerly Select One Don't recall
Type - Unrecalled	Currently Formerly Select One Don't recall
Other	Currently Formerly Select One Don't recall

Back Save and Review

Study Description developed with NPCA

Peace Corps Connect home page sliding graphic headline:

Are you a woman who served in the Peace Corps between 1961 and 1990?

You could be a part of an exciting new study examining breast cancer risk.

(Graphic links to Study Info page)

Study Info Page

Breast Cancer Risk Study for Returned Peace Corps Volunteers

The National Peace Corps Association has partnered with researchers at Baylor College of Medicine to determine if there is a link between the risk of cancers, including breast cancer, and medications taken during Peace Corps service. The investigators have developed an online survey for RPCVs to measure health and health-affecting behaviors.

Our goal is that EVERY woman who served in the Peace Corps between 1961 and 1990 is represented in this study.

Studies in animals suggest that people who took a commonly used medication in the past to prevent or cure malaria may be at lower risk of developing some diseases today, such as cancer or heart disease. Baylor College of Medicine is developing studies to examine this link, beginning with this online survey to compare a large group of women who took the medication to a large group of women who didn't.

Female RPCVs who served between 1961 and 1990 represent an ideal group of people in whom to study this possible link, because about half of RPCVs took medication as part of their service and about half of them did not. Additionally, over 20 years has passed since their service and related medication use, so we can look at health changes over a long period of time.

The survey consists of about 50 questions and can be taken online or over the phone. Question topics include lifestyle risk factors for diseases and a brief medical history.

ACTION: Please share this page with people you served with in the Peace Corps.

PAGE 2 (JUMP LINKS TO THESE HEADERS)

How can I access the survey?

Identifying as many RPCVs who served between 1961 and 1990 as possible is critical to the study's validity; however, no exhaustive list of RPCVs who served during that time is available. To recruit a generalizable sample, the BCM researchers are using an innovative sampling method called Respondent-Driven Sampling. Based on social networks, this new approach works like a chain letter, in which participants answer the survey and invite their friends to answer the survey, adding onto the chain of RPCVs. The longer the chain of participants, the more representative the sample will be of the entire group of RPCVs. You have to be referred to the study by another woman who served between 1961 and 1990 to take the survey.

For more information on Respondent-Driven Sampling methodology, click here. <Link to paper on RDS>

This novel sampling method requires a small group of a few hundred RPCVs to start a movement that will grow to up to 18,000 RPCVs! Researchers have compiled a list of several hundred female RPCVs who will plant the seeds of the survey. Each of these women will invite up to 7 other RPCVs to participate. Those 7 women will each invite 7 more women, and so on.

ACTION: Watch your email for an invitation to fill out the survey and be sure to invite your friends to participate when you get one.

Who can take the survey?

We would like every woman who served between 1961 and 1990 to be represented in this study. No matter where you served, what medications you took or did not take, or if you have even been diagnosed with cancer or not, if you are a woman who served between 1961 and 1990, then you can take the survey.

To get a complete picture of all the women who served between 1961 and 1990, the researchers also need to count those who have died since their service. If you served with women who have died, you can give the researchers their names and service dates and they can be counted as well. To submit the name of a woman who has died, click here.

What about men or those who didn't serve in the Peace Corps?

Although men will not fill out this RPCV survey, there will be studies in the future that include men. If you would like to learn about future studies, you can join our mailing list.

Men or friends of the Peace Corps are encouraged to submit names of women who have died since their service. If you served with women who have died, you can give

the researchers their names and service dates and they can be counted as well. To submit the name of a woman who has died, click here.

Why should I take this survey? Who benefits?

While there are no direct benefits to the RPCVs who help with the survey, this survey may help researchers discover new uses for an off-patent (which means that no pharmaceutical company owns it) and inexpensive drug.

Who is paying for this study?

In the early 1990s, a powerful grassroots advocacy movement campaigned for an increase in breast cancer research funding, and in 1993 the National Breast Cancer Coalition presented President Clinton with a 2.6-million signature petition for "a comprehensive plan to end the breast cancer epidemic." Congress responded by appropriating funds targeted specifically toward winning what came to be known as "The War on Breast Cancer." This appropriation marked the beginning of the Congressionally Directed Medical Research Program (CDMRP). The CDMRP is a unique partnership among the U.S. Congress, the public, and the Department of Defense to reduce bottlenecks and gaps in medical research. The project is funded by a grant from the Congressionally Directed Medical Research Projects administered by the US Department of Defense.

CDMRP is funded through the Department of Defense (DoD), via annual Congressional legislation known as the Defense Appropriations Act. For most programs, the DoD sends a multi-year budget request to Congress in the form of the President's Budget. However, dollars for the CDMRP are not considered part of the DoD's core mission, and are therefore not included in the DoD's requested budget. Rather, the dollars to fund CDMRP are added every year during the budget approval cycle by members of the House or Senate, in response to requests by advocates. This gives the CDMRP a unique ability to respond quickly to changes in science and public need. In addition to breast cancer, CDMRPs support research for diseases such as ALS (Lou Gehrig's disease), MS, and autism.

Are there any risks to participating in the study?

The risks to you are minimal. Participation in research may involve some loss of privacy. However, your records will be handled as confidentially as possible. Access will be limited to the data manager and the researchers organizing the study and will require a password. No information will be used for research without additional permission. Your contact information will not be shared with anyone outside of Baylor College of Medicine.

Are there any financial considerations?

There will be no cost or payment to RPCVs or their loved ones who respond to the survey.

How long will the study take?

The online survey will take a participant up to 40 minutes. You will be able to save the survey at any time to return and log into the study website to finish it later.

What about RPCVs who do not have Internet access?

The study can be completed over the telephone.

What is the National Peace Corps Association's role in this study?

The National Peace Corps Association is assisting Baylor College of Medicine in raising awareness about this path-breaking study. The National Peace Corps Association will not have access to individual survey answers, but will help share the overall study results in a variety of ways. Because of the unique nature of this research and the potential for such a great benefit to society, NPCA is sharing its entire database with Baylor College of Medicine and helping to spread the word so that everyone within the greater Peace Corps community, including friends and family members of Peace Corps Volunteers, are informed about the study and encouraged to participate. (You may control your preferences for such data sharing opportunities by selecting "profile update" accessible at https://secure.peacecorpsconnect.org/npcassa.)

When can I see the final results of this study?

The survey results will be published in academic journals, on PeaceCorpsConnect.org, and will be emailed to people registered to receive information on future studies link to future studies registration>. Because of the tremendous task of collecting surveys from up to 18,000 RPCVs, final survey results will not be collected, analyzed, and published until mid-2014.

What do I do if I have questions, now or later?

Contact the study team directly at XXX@bcm.edu.

CorpsChronicles: An Application Leveraging Social Networks for Online Epidemiologic Survey Studies

¹Mayfield P, ¹McOwiti A, ¹Barney J, ¹Qu Shijing, ¹Sexton KR, ²Harris AM, ²McArthur K, ¹Hilsenbeck SG, ¹Bondy ML, ¹Conneely O, ¹Spitz M, ^{1,2}Dasco CC, ¹Becnel LB

INTRODUCTION: Exposure to chloroquine, an off-patent anti-malarial drug with a 60-year history of use by millions, reduces the incidence of breast cancer in genetically programmed rats by 37%. To study whether such exposure may have a protective effect in humans requires a robust bioinformatics system. CorpsChronicles, is a secure, database-backed web application developed to manage large, socially-oriented epidemiologic studies. The first such study is of Returned Peace Corps Volunteers (RPCVs) who served from 1961-1990 in malaria-endemic areas who received chloroquine prophylactically. CorpsChronicles was used to collect information from RPCVs to determine if chloroquine exposure in this cohort associates with reduced breast cancer incidence compared with the general US population.

METHODS: CorpsChronicles is a Java Enteprise Edition 6 application that uses the model-viewcontroller architecture with the JBoss Seam framework to manage its various components. The application runs within JBoss 7 middleware and utilizes Oracle 11g for data persistence, though Hibernate makes it possible to swap out Oracle for other database management systems. Security is managed by enhanced Seam security, encrypted passwords, role-based permissions, Captchas, and unique "coupon" codes that serve as keys for respondents. Survey skip patterns are managed by Drools rules engine to allow independent, business-driven rules to be implemented for individual studies. "Friend recommendations" of respondents are based on a novel social network sampling methodology, Respondent-Driven Sampling (RDS), in which initial respondents ("seeds") identify members of their social network who may want to participate and these members in turn identify members of their network thus creating network "chains" for each seed. Data from links in each chain are compared back to their seed data using RDSAT freeware; demographic and clinical characteristic data is compared between groups using chi square or Fisher's exact test and t-test or Wilcoxon rank-sums. Multivariate logistic regression is used to assess associations between chloroquine exposure and breast cancer incidence, with the Network Scale-Up Method, a social networking method, applied to estimate the hard-to-count population of RPCVs.

RESULTS: CorpsChronicles (i) allows research subjects to securely answer dynamic questionnaires; (ii) manages electronic recruitment and referral workflows; (iii) sends email reminders to participants to complete surveys; and (iv) provides feedback to end users on the overall progress of the study in a way that does not compromise the results. Because breast cancer known risk factors include family history, race, and lifestyle, the online survey elicits self-reported data on potential confounding variables which will be accounted for in the statistical analysis to maximize internal validity. Two hundred highly-connected RPCVs were selected as initial seeds, with a goal of 17,000 respondents across all chains. Interim analyses were performed at one and two months post-activation to ensure appropriate accruals with RDS.

DISCUSSION: In this talk, sampling mechanisms, technical components of CorpsChronicles and preliminary results of the study will be discussed.

¹Baylor College of Medicine and ²The Methodist Hospital Research Institute, Abramson Center for the Future of Health, Houston, TX, 77030, USA

Designing a Translational Epidemiologic Study: Chloroquine and breast cancer chemoprevention in Returned Peace Corps Volunteers

Krystal Sexton, PhD ¹ Amy M. Harris, MPH ^{2,3} Kara McArthur ^{2,3} Melissa L. Bondy, PhD ¹ Susan Hilsenbeck, PhD ¹ Lauren Becnel, PhD ¹, Pamela Mayfield ¹, Orla Conneely, PhD ¹ Courtney M. Queen, PhD ³ Margaret R. Spitz, MD ¹ Clifford C. Dacso, MD, MPH ^{1,2,3}

BCM
Baylor College of Medicine

(1) Baylor College of Medicine, (2) The Methodist Hospital Research Institute, Abramson Center for the Future of Health, (3) University of Houston

INTRODUCTION

This study represents a multidisciplinary approach for drug repositioning through translational research using novel sampling & analysis methods.

Background

- Drug Repositioning is a promising approach to reducing the cost and timeline of new cancer drug development (routinely >\$1 billion and 10-15 years for FDA approval).
- Agents with established toxicologic, pharmacokinetic, and pharmacodynamic profiles are redeployed for new conditions.
- Following-up laboratory screening of such drugs with well-designed retrospective cohort studies, rather than a traditional, prospective clinical trial, could greatly expedite the identification and testing of promising drues.

Chloroquine, an off-patent malaria prophylaxis with a 60-year history in millions of people, reduces the incidence of breast cancer in rats by 37%. [3-5]



Fig. 1: Chloroquine reduces the incidence of NMU-induced mammary tumors in WT female Wistar-Furth rats (dark line), but shows no breast cancer protective effects in a BALB/c p53-null mammary epithelium model (light line). [5]

The challenge, particularly in the context of the dismal success rate of translational medicine [1], was to design an epidemiological study that quickly and cost-effectively determine chloroquine's effectiveness in humans.

Challenges in designing the study

- Time: Breast cancer can develop several decades after exposure to chloroquine.
- A prospective study would require 10 50 years of follow up
- Study population: This study requires a representative sample of a well-defined study population
 - The study population must include participants with and without breast cancer, including those who have died of breast cancer.
- Risk factors: Breast cancer is a complex disease affected by family history of breast cancer, race, and lifestyle factors.

OBJECTIVE

Design an epidemiologic study to quickly, safely, and cost-effectively evaluate the effect of chloroquine on breast cancer risk in humans.

ACKNOWLEDGEMENTS

Apollo McOwity, Abramson Center for the Future of Health, Funded by the Department of Defense CDMRP program, National Peace Corps Association





METHODS

Study Design: A retrospective cohort study

Exposure of Interest: Use of chloroquine as malaria prophylaxis before the year 1990

- Timeline for exposure ends in the year 1990 because this is when the CDC revised its recommendations for malaria prophylaxis in resistant regions away from chloroquine.
- . Timeline also takes into account the long latency of breast cancer.
- Chloroquine use measured by self-report and cross-referenced by Peace malaria prevention protocols by year and location.

Outcome: Breast cancer diagnosis (self-reported)

Population: Female Returned Peace Corps volunteers (RPCVs) who served between 1961 and 1990 (N=51,828). During that period, ~65% of volunteers were required to take chloroquine for the term of their service.



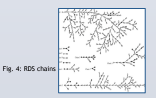
Fig. 2: Peace Corps countries



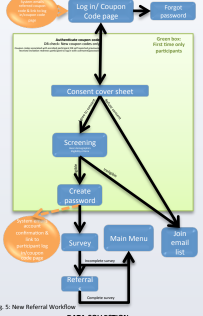
Respondent-Driven Sampling [2,7]

RPCVs are a highly educated group of people who tend to have tight social relationships with other RPCVs.

- Large membership association (National Peace Corps Assoc) and 100s of small membership groups.
- Respondent-Driven Sampling (RDS) is a variation of snowball sampling that builds a broad representative sample.
- Uses a mathematical system for weighting the sample to compensate for its not having been drawn as a simple random
- Allows for valid inferences about the characteristics of the population from which the sample is drawn.
- Starts with a sample of the target group (called "seeds") and expands through successive waves of recruitment, capitalizing on strong social ties among group members.
- RDS has never been done using Internet-based data collection on this scale



New Referral Workflow



DATA COLLECTION

Online Data Collection Tool

Developing a proprietary data collection application, which will:

- Collect survey responses online with piping and skip patterns
- Allow participants to input referrals
- Track RDS chains of referrals
- Follow up with participants via email to increase response rate
- International and national standards compliance (CDISC BRIDG model, NCI EVS controlled vocabularies and data elements)
- Technology stack: Java Enterprise Edition 6, JBoss 7, RHEL 5, Oracle 11g, F5 proxy server



Project Name Sub-right one Sub

Fig 7: Online Data Collection Tool Wireframe

DATA ANALYSIS

- The online RDS data will be analyzed using the Respondent Driven Sampling Analysis Tool (RDSAT) (Ithaca, NY).
- Demographic and clinical characteristics will be compared between groups.
- Categorical variables (e.g., family history) chi square or Fisher's exact test
- Continuous variables (e.g., age at diagnosis) t-tests or Wilcoxon rank-sum.
- RDSAT will calculate RDS weights.

 Multivariate logistic regression model survival will be used to assess the association between chloroquine exposure and breast cancer, adjusted for confounder survival rates from SEER database and CDC.

 Sample size & minimum detectable (PG).

 Sample size & minimum detectable (PG).

This is equivalent to changing "lifetime risk" from 1 in 9 to 1 in 12 with an associated number needed to treat of 36.

EXPECTED RESULTS

As in any retrospective study design, we must work to minimize bias and confounders. However, the gains in cost and time efficiencies allow the translation of preclinical data on breast cancer chemoprevention into the repositioning of a well-characterized and relatively benign drug to reduce breast cancer risk.

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C3PR, CODR & Corps Chronicles: Case Studies of Clinical and Epidemiological Research Databases within the



DLDCC

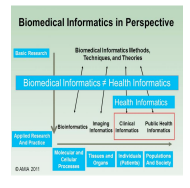
Pamela K. Mayfield, Apollo McOwiti, Jonathan Barney, Yolanda Darlington, Geetu Vanjani and Lauren B. Becnel

Biostatistics and Informatics Shared Resource (BISR), Dan L. Duncan Cancer Center & Lester and Sue Smith Breast Center, Baylor College of Medicine

Abstract

The Biomedical Informatics Group (BIG) of the DLDCC Biostatistics and Informatics Shared Resource provides services to Cancer Center members conducting clinical or epidemiologic research. BIG members were actively involved in establishing national and international data and object standards such as the NIH's standardized case report forms and components of the BRIDG model from the Clinical Data Interchange Standards Consortium (CDISC). Locally, we have implemented these standards in sister clinical research databases, C3PR and CODR. C3PR is a patient registration system created by the National Cancer Institute that BIG hosts to manage core study and patient data for all cancer clinical trials. Certain gaps in C3PR functionality (e.g. streamlined generation of Summary 4 and other reports, data management for PRMS and data review oversight committees) were addressed with the design/creation of a sophisticated, web-based companion application, CODR, which is tightly integrated with C3PR and provides an end-to-end view of clinical trial activity across all DLDCC programs. In addition to the standards-based clinical research support, BIG is currently implementing Corps Chronicles, a system that allows tens of thousands of research subjects to securely answer questionnaires using rules engine-managed set of logic. As a result, this web-based system must be highly scalable, able to handle thousands of simultaneous connections, and incorporate workflow and business logic to manage numerous recruitments, accruals and both internal and external communications (eg, reminders to participants to complete surveys). Although the initial system is being built in support of a breast cancer prevention study involving Peace Corps volunteers, the system design will be generalized to allow addition of survey-based studies for other cancers in the future.

Types of Informatics



BISR Contributions & Collaborations

LOCAL

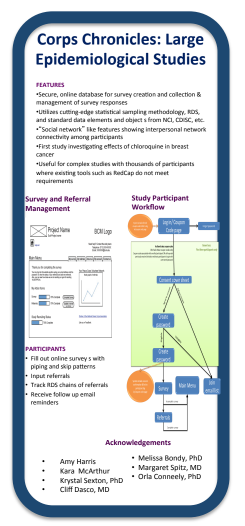
- Developed & hosted international and national standardscompliant online databases
- •Clinical & epidemiological/population science research focus areas

NATIONAL & INTERNATIONAL

- •Participated in the creation of standard case report forms for clinical trials
- •Major co-developers of the NCI Life Science Business Analysis Model (BAM)
- •Contributors to the Clinical Data Interchange Standards Consortium's (CDISC) BRIDG Domain Analysis Model & NCI Enterprise Vocabulary Services terminologies and data elements



CODR: Clinical Trials Oversight Secure, online database for clinical trials oversight committee (IRB, DSM, etc.) management Provides reminders of studies with upcoming & overdue Fully integrated with C3PR Oversight Committee **CODR Data Model Using CDISC** Standards & Reporting **Contributions to National Clinical Trials Management: Data and Object Models** NCI Biomedical Research BAM UML Use Cases Are Used to Create BAM Activity Diagrams from which Components of the BRDIG Object Models Were Developed Acknowledgements, See C3PR Section



Appendix I: Think Aloud Testing Protocol and Instrument

Pre-Test of the Online Survey Document Protocol, Summary, and Instructions

The Pre-Test of the Online Survey will consist of two parts: a Think-Aloud and a Debriefing.

Number of Subjects: 5

Time commitment from subjects: 1 hour

I. Think Aloud

Think Aloud is a method that allows researchers to understand, at least in part, the thought process of a subject as she completes a survey. The researcher listens while the user attempts to complete the survey and "narrates" her experience. Ideally, the observer speaks only to remind the user to "please keep talking" should she lapse into silence.

By thinking aloud while attempting to take the survey, users can explain their thought processes as they read the questions and complete the survey, and illuminate any difficulties they encounter or places they feel uncomfortable.

This Think Aloud will be done on the telephone with one subject and one or two CITI-trained research interviewers. Test subjects will be matched to the ultimate users of the survey on all dimensions, including age, sex, and study exclusion and inclusion criteria. The Think Aloud will take place in the actual environment where a user would need to complete the task; that is, the Think Aloud subject will be at home, taking the survey on her home computer, while the interviewer listens on the telephone.

Because the survey is web-based, the Think Aloud will be supplemented by objective data gathered by the survey program: response latency (how long it took the user to complete the entire survey, in minutes), backups (which questions the user back-tracked to complete), and entry errors (which questions provoked backspaces and changed answers).

Think Aloud Protocol Steps

1. The interviewer will prepare the subject.

- ♦ Describe the goal of the task, "we are pre-testing this online survey instrument to identify any problems, questions, or concerns users might have in completing it."
- ♦ Briefly explain the Think Aloud procedure to your subject. "Please say aloud everything you think while you answer the questions in the survey."
- ◆ Do a practice Think Aloud task to familiarize the subject with the procedure.
- § Ask the subject to open their Web browser, email, and navigate to the emailed survey, making sure that they are comfortable with describing their actions and thoughts in detail.
- ♦ Tell the user that:
- § You are testing the instructions, not the user, and that any difficulties are your fault, not theirs
- § They can stop the task at any time if they become uncomfortable.
- § They may ask questions at any point in the process, but you may not answer them.

§ You will not tell them when they have completed the task; they must determine this on their own.

2. Think Aloud

- ◆ Verify that the subject has no remaining questions about the Think Aloud process.
- ♦ Ask the subject to begin the survey.
- ◆ Throughout, if necessary, prompt the user with "please keep talking."
- ♦ Take extensive notes: everything the user says is relevant.

3. Thank the Subject and Prepare for the Debriefing

♦ When the subject believes she has completed the survey, thank her for participating and ask her to please answer 11 more questions about the survey as a whole.

II. Debriefing Protocol Steps

Ask the subject the following questions. Write down their answers in as complete a form as possible.

- 1. Is there any question that you did not feel comfortable answering? Which one? Why? (repeat as often as necessary)
- 2. Do you have any suggestions for improving the questionnaire?
- 3. Would you feel comfortable forwarding this survey in its current configuration? Why not?
- 4. How did you feel about the length of the survey?
- € Too long
- € An appropriate length
- € Too short
- 5. I get the impression that this study comes from Baylor College of Medicine and is a legitimate academic study.
- € Strongly agree
- € Somewhat agree
- € Not sure
- € Somewhat disagree
- € Strongly disagree
- 6. I felt concerned about my privacy or confidentiality.
- € Strongly agree

€	Not sure				
€	Somewhat disagree				
€	Strongly disagree				
7. I	feel comfortable sending my answers to this survey to the research team over the Internet.				
€	Strongly agree				
€	Somewhat agree				
€	Not sure				
€	Somewhat disagree				
€	Strongly disagree				
	8. I would be willing to submit contact information for up to 7 friends to invite them to participate in this study.				
€	Strongly agree				
€	Somewhat agree				
€	Not sure				
€	Somewhat disagree				
€	Strongly disagree				
9. I would be more likely to submit contact information of friends if					
Thank the subject for participating. Ask if she has any questions. Refer any questions you cannot answer to Dr. Dacso for follow up and response.					

Somewhat agree

Appendix J: Toll-Free Telephone Number and Email Help Line Statistics

Table 1: User Queries and Investigator Responses

Coupon Code	Issue Raised	Date	Person	Status
·			Who Attended to Issue	
1ad296b7-ec5b- 4c15-b9eb- c51cebc0d686	Unable to answer questions because they were greyed out. Went to pick up the phone. Came back and was logged out	3.26.14	K.M.	Escalated to bioinformatics. Determined issue was user error. Participant was able to answer survey on second attempt.
27bb4f5f-10c4- 4289-a777- 932ecd776f77	In filling out the study questions, there is no place to indicate if you breast fed your baby. I cannot move forward unless I answer but there is no place to say yes or no.	3.27.14	K.M.	Escalated to bioinformatics. Viewed subjects responses via participant assistance feature. Determined by bioinformatics to be user error.
52a154b9-2e70- 4457-a916- 3cca1f205907	REMOVE	3.25.14	P.M.	Designated "DO NOT CONTACT"
	REMOVE	3.31.14	C.L.	Designated "DO NOT CONTACT"
491a3d81-6e4f- 4480-aefe- 6ff249978adb	Complained about receiving too many emails after she submitted the survey	4.9.14	J.O'B.	Designated "DO NOT CONTACT"
fdebeeba-d8b0- 4abb-bf5d- 110b8dc2a608	Requested to join study via facebook page; concerned about source of invitation and confused about why info about sign-up not on facebook page	4.12.14	J.O'B.	Replied to email explain study methodology; offered that she could have friends email us if she is uncomfortable with referral. Thanked us for reply, but will not be participating.
c2766a3c-dc1e- 42d5-8f85- 7c78e1a2e8bd	Logged out of survey because the site wouldn't let her answer the breast feeding question and wouldn't let her proceed without answering.	4.16.14	J.O'B.	Replied with email; told her to log out & in again, and if that is not successful, contact us for participant assistance. She replied an hour later; was able to select the breastfeeding question when she tried a second time.
e84bb5ed-e2ed- 45ce-8975- db4e675f4204	Emailed to say she signed up ages ago, and helped us with pilot. Also states she hasn't taken the real survey.	4.16.14	J.O'B.	Sent an email to clarify that this most recent email is her official invitation, and that she should use the link to take the survey and add referrals
e84bb5ed-e2ed- 45ce-8975- db4e675f4204	Emailed with problem with breast feeding question. "it won't let me put in a pregnancy until I say if I breastfed, but there is nothing to	4.17.14	J.O'B.	Sent email to tell her to try and log out, then log in again. Escalated to bioinformatics. Programming error identified, and then fixed.

	click to say yes/no"			
a52ae3c5-ee80- 4db7-b009- a0487cec93f3	Emailed with request to amend her response	4.17.14	J.O'B.	Emailed to tell her we are working on figuring out how to edit responses. Will investigate with bioinformatics. Update: bioinformatics says it is not possible to amend. She emailed again on 4/30 to follow up, and I told her it is not possible to change her answers at this time.
8b199351-ff23- 4f63-a0b2- 7f828f375dd0	Could not get past questions 47-49. Also said, "the diagnostic doesn't make sense."	4.30.14	J.O'B.	Emailed to ask her exactly what the problem was, and what browser she's using. Told her she has to check a response to every row in the table. RESOLVED.
a3ed7cad-c183- 4417-97f2- 7f6c3c497ea2	Emailed because she completed the study, but got a reminder to start the study	5.6.14	J.O'B.	Somehow she has two invitations (two email address in system). Replied to her by email, and shut off email notifications for second account.
246bc734-8f0c- 4b61-8ca8- 3b32177d3f0f	Got kicked off of survey in the middle of taking it	5.7.14	J.O'B.	Emailed link to her; told her to let us know if it happens again
c6f7a58f-5f6f- 4a39-9df0- d4fbd712c2a9	REMOVE	5.13.14	J.O'B.	Designated "DO NOT CONTACT"
f5444138-f3f0- 4c52-b217- 929de921753e	Has MS; says she is unable to complete survey	5.15.14	J.O'B.	Designated "DO NOT CONTACT". Offered to do survey over the phone, but she says she has no interest because she "cannot relate to the questions about how many women were doing what in her group." Commented that 52 years ago was too long ago to remember.
28c77102-135a- 4e4c-b35c- 3bd8e5e7002a	Reported that she could not log on again even with correct username and password	5.15.14	J.O'B.	Advised her to close browser and start again, and if not to reset password.
c384f6dc-adf1- 42ae-9491- 4634d27b3bd3	Requested email address change to her primary account	5.22.14	J.O'B.	Escalated to bioinformatics
8b199351-ff23- 4f63-a0b2- 7f828f375dd0	REMOVE	5.22.14	J.O'B.	Designated "DO NOT CONTACT"
ba6acdbd-cd4d- 48fe-a5b8- 35f2f0b9fce2	Sent blank email with subject, "a balky e-form"	5.22.14	K.M.	Requested clarification from participant of exact problem
979fefe3-b569- 4c82-8a91- 83b0e85e642	Emailed requesting account update, but email was blank	5.22.14	J.O'B.	Requested clarification from participant