AWARD NUMBER: DAMD17-02-2-0017

TITLE: Population Health Trial for Smokeless Tobacco Cessation with Military Personnel

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Eugene, Oregon 97403

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While smoking cessation has received considerable attention within the military, the use of smokeless tobacco (chewing tobacco and snuff) has not been a focus of medical services or research. Epidemiological data suggest that while smoking has continued to decline both in the general population and within the military, the use of smokeless tobacco products has increased. The primary objective of this research is to develop and evaluate an intervention for smokeless tobacco cessation comprised of proactive recruitment, targeted written and video materials mailed to the participant, and phone call support. At the end of this third year of the study, several accomplishments have been achieved. Accomplishments include visits for orientation and training at dental clinics at 14 additional military installations: 6 AF sites were added for enrollment purposes; 5 Army sites; Camp Pendleton, CA, for Marine enrollment; and 2 Navy clinics in San Diego for enrollment of Navy personnel. A total of 785 participants have now enrolled in the study. Mailed follow-up assessments at three and six months post enrollment were continued, with 587 three-month surveys and 571 six-month surveys completed to date.
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INTRODUCTION

While smoking cessation has received considerable attention within the military, the use of smokeless tobacco (chewing tobacco and snuff) has not been a focus of medical services or research. Epidemiological data suggest that while smoking has continued to decline both in the general population and within the military, the use of smokeless tobacco products has increased. The primary objective of this research is to develop and evaluate an intervention for smokeless tobacco (ST) cessation comprised of proactive recruitment, targeted written and video materials mailed to the participant, and phone call support. The primary hypothesis to be tested is that participants randomized to receive the intervention will quit their tobacco use at a significantly higher rate than participants receiving usual care. Active duty U.S. Armed Forces personnel stationed at military locations that were identified as current ST users when completing their annual preventive oral health assessment have been recruited to participate in a randomized two-group design that compares a brief contact intervention with the usual preventive health care. Follow up assessments were completed at 3- and 6-months by mail or phone to assess the impact of the program.

BODY

In the fourth year of our grant, we finished enrollment and follow-up. In total our study recruited participants at 21 military bases as listed in the Table 1 below. Across all sites we enrolled 785 participants. 587 participants completed the 3-month follow-up and 571 completed the 6-month follow-up.

Table 1. Recruitment Intervention Sites

<table>
<thead>
<tr>
<th>Base</th>
<th>POC</th>
<th>Start Date</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Air Force</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lackland AFB, TX (3 clinics)</td>
<td>Lt Col (Dr.) Alan Peterson</td>
<td>09/02/2003</td>
</tr>
<tr>
<td>Randolph AFB, TX</td>
<td>Col Carlos Esquivel</td>
<td>09/08/2003</td>
</tr>
<tr>
<td>Wright Patterson AFB, OH</td>
<td>Lt Col (Dr.) Jeff Cigrang</td>
<td>09/02/2003</td>
</tr>
<tr>
<td>Dyess AFB, TX</td>
<td>Lt Col Randall Griffin</td>
<td>11/17/2003</td>
</tr>
<tr>
<td>Sheppard AFB, TX</td>
<td>Capt (Dr.) Bruce Abe</td>
<td>11/18/2003</td>
</tr>
<tr>
<td>Brooks City Base, TX</td>
<td>Maj Jacob Palma</td>
<td>01/07/2004</td>
</tr>
<tr>
<td>Laughlin AFB, TX</td>
<td>Capt (Dr.) Mark Halverson</td>
<td>01/13/2004</td>
</tr>
<tr>
<td>Robins AFB, GA</td>
<td>Maj Elizabeth Tandy</td>
<td>06/06/2004</td>
</tr>
<tr>
<td>Mt Home AFB, ID</td>
<td>Capt. William Quinn</td>
<td>08/03/2004</td>
</tr>
<tr>
<td>Little Rock AFB, AR</td>
<td>Lt Col Robert Abbott</td>
<td>08/24/2004</td>
</tr>
<tr>
<td>Langley AFB, VA</td>
<td>Col Robert Sabatini</td>
<td>09/01/2004</td>
</tr>
<tr>
<td>Nellis AFB, NV</td>
<td>Lt Col Jeff Thompson</td>
<td>09/28/2004</td>
</tr>
<tr>
<td>Eglin AFB, FL</td>
<td>Col Mike Garrett</td>
<td>11/09/2004</td>
</tr>
<tr>
<td><strong>Army</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ft. Sam Houston, TX</td>
<td>COL Ronald Lambert</td>
<td>09/18/2003</td>
</tr>
<tr>
<td>Ft. Sill, OK</td>
<td>LTC Charles Sabadell</td>
<td>07/15/2004</td>
</tr>
<tr>
<td>Ft. Polk, LA</td>
<td>COL Thomas MacKenzie</td>
<td>08/16/2004</td>
</tr>
<tr>
<td>Ft. Leavenworth, KS</td>
<td>LTC Robert Windom</td>
<td>08/03/2004</td>
</tr>
<tr>
<td>Ft. Drum, NY</td>
<td>COL Robert Rock</td>
<td>09/04/2004</td>
</tr>
<tr>
<td>Ft. Knox, KY</td>
<td>COL Stephen Rouse</td>
<td>04/05/2005</td>
</tr>
<tr>
<td><strong>Navy</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>North Island Dental Clinic</td>
<td>CAPT Richard A. Joralmon</td>
<td>06/17/2004</td>
</tr>
<tr>
<td><strong>USMC</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Camp Pendleton, Clinic 13</td>
<td>CAPT Wayne Osborne</td>
<td>06/18/2005</td>
</tr>
</tbody>
</table>

Of the 785 participants, 392 were assigned to the Treatment Group (TX). 370 (94%) of TX participants were successfully contacted by phone and offered cessation materials and MI calls, 351 were mailed cessation materials. Figure 1 below illustrates the number of subjects at each
stage of the intervention. Six percent of TX participants either asked to be dropped or were unreachable due to lack of contact information, deployment or did not answer their phone or return messages. 282 (72%) of TX participants completed the three follow-up assessment and 274 (70%) completed the six-month follow-up assessment. 17 TX participants actively dropped during MI calls and 20 TX participants refused additional MI calls but were willing to complete follow-up assessments.

Figure 1. DoD Protocol Flowchart

Enrollment (21 Dental Clinics)

Randomization (785 enrolled participants)

Brief Contact Intervention (392 TX Participants)

Usual Care (393 Control Participants)

Call 1
370 participants completed Call 1

Cessation Materials
351 participants received materials

Call 2
328 participants completed Call 2

Call 3
287 participants completed Call 3

3-month follow-up
282 participants completed 3-month follow-up

6-month follow-up
274 participants completed 6-month follow-up

3-month follow-up
305 participants completed 3-month follow-up

6-month follow-up
297 participants completed 6-month follow-up

Cessation materials offered to participants
174 participants requested cessation materials

Of the 785 participants 393 were assigned to the Usual Care (UC). The UC condition consisted of the standard practice used at the study sites during the annual dental examination of active duty military personnel. In most cases, this included asking about smokeless tobacco use, advising individuals to quit, and providing a referral to the local tobacco cessation program for those interested in quitting. 305 (78%) of UC participants completed the three follow-up assessment and 297 (76%) completed the six-month follow-up assessment. 174 (59%) of UC participants who completed the 6-month survey requested to be mailed cessation materials upon completion of their participation (see Figure 1 above).
At the 3-month follow-up 100 participants were either lost due to lack of contact information (39) or deployed (61). We continued to search the military data base as well as other commonly used search methods for participants lost due to lack of contact information. At the 6-month follow-up 110 participants were lost (35) or deployed (75).

47 total participants dropped out of the study during 3- and 6-month follow-up assessments; bringing the total numbers of drops for this study to 64.

**Participant Characteristics:**

Tables 2 provides descriptive statistics on the sample of participants in the study. As expected the vast majority were male and 89% were white. Most were married (72%) and most had some college (82%). The vast majority were enlisted personnel (87%) and the majority of the sample came from the Air Force (66%).

Table 2. Demographics

<table>
<thead>
<tr>
<th>Demographics</th>
<th>Branch of Military (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>Air Force 65.6%</td>
</tr>
<tr>
<td>White</td>
<td>Army 30.2%</td>
</tr>
<tr>
<td>Hispanic</td>
<td>Marines 3.15</td>
</tr>
<tr>
<td>Married</td>
<td>Navy 1.15</td>
</tr>
<tr>
<td>Some college or higher</td>
<td>Enlisted vs. Officer</td>
</tr>
<tr>
<td>Age</td>
<td>Enlisted 86.8% (n = 681)</td>
</tr>
<tr>
<td>BMI</td>
<td>Officer 13.2% (n = 104)</td>
</tr>
</tbody>
</table>

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<td>Some college or higher</td>
<td>Enlisted vs. Officer</td>
</tr>
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<tr>
<td>BMI</td>
<td>Officer 13.2% (n = 104)</td>
</tr>
</tbody>
</table>

Table 3 reports indicators of tobacco dependence and describes the current tobacco use of the sample. One indication of tobacco dependence is the co-use of cigarettes and smokeless tobacco and 20% of the sample reported concurrently using both tobacco products. Most subjects reported using smokeless daily and using ST products for about 13 years. Two other items of dependence are using ST within 30 minutes of waking in the morning and 24% reported this behavior, while almost 50% reported sometimes or always swallowing the tobacco juice while using snuff or chew.

Table 3. Current Tobacco Use of DoD Smokeless Tobacco Participants – Indicators of Dependence

<table>
<thead>
<tr>
<th>Current Smoker</th>
<th>Current Tobacco Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes 20%</td>
<td># of days chew per week 6.21 (sd = 1.47)</td>
</tr>
<tr>
<td>No/missing data 81.1%</td>
<td>Days tin lasts 3.7 (sd = 8.39)</td>
</tr>
<tr>
<td>Age 30.40 (sd = 7.63)</td>
<td>Age of initiation 17.70 (sd = 5.11)</td>
</tr>
<tr>
<td>BMI 26.7 (sd = 3.23)</td>
<td>Length of use in years 12.80 (sd = 8.39)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Tobacco use by spouse or friends</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spouse smokes 23%</td>
</tr>
<tr>
<td>% of closest friends who chew 81.7%</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>First Chew in the Morning</th>
<th>Swallow the Juice</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;30 min 23.9%</td>
<td>Never 50.1%</td>
</tr>
<tr>
<td>30-60 min 34.5%</td>
<td>Sometimes 37.4%</td>
</tr>
<tr>
<td>&gt;60 min 41.6%</td>
<td>Almost always 12.5%</td>
</tr>
</tbody>
</table>
Recruitment of Intervention Sites

Described below are the Air Force, Army, Navy, and Marine sites (Table 1 above) that participated in our study. The point of contact (POC) is the person located at the dental clinic who oversaw the project at that site.

Air Force

All of the Air Force sites listed above honored the Wilford Hall Medical Center (WHMC) IRB approval of our protocol, with the exception of Wright-Patterson AFB, OH, Eglin AFB, FL, and Brooks City Base, TX, which have their own IRBs. Our protocol had been previously approved by the Wright-Patterson Medical Center IRB in 2003 and by the Brooks City IRB in Jan 04. Our protocol was expeditiously cleared through both the Eglin IRB later in 2004 to allow us to begin recruiting participants there in November 2004.

Army

Four of the participating Army sites are in the Great Plains Medical Region and thus fall under the authority of the BAMC IRB. They were: Ft Sam Houston, TX; Ft Sill, OK; Ft Polk, LA; and Ft Leavenworth, KS. The BAMC IRB approved our protocol for these sites in expeditious fashion, allowing for recruitment to continue (at Ft Sam Houston) and to begin at Ft Sill, Ft Polk, and Ft Leavenworth in the Jul-Aug 2004 timeframe. The BAMC IRB allowed us to use the short, 4-page enrollment packet (see attached Appendix A) that had been approved for the Air Force by WHMC. As a consequence, enrollment at sites in the Great Plains Medical Region was very good and on a par with AF sites where the shorter enrollment packet was used.

North Atlantic Medical Region Army Sites

The remaining Army sites, Ft Drum, NY, and Ft Knox, KY, are in the North Atlantic Medical Region and thus fall under the authority of the Walter Reed Army Medical Center IRB (WRAMC). The process of gaining IRB approval from WRAMC for enrollment at Ft Drum was quite protracted, lasting at least 12-months. Once the approval was obtained, we were required to use a much longer enrollment packet (see attached Appendix B), i.e., 10 pages long vs. the 4-page enrollment packet approved for AFBs (under the WHMC, Wright-Patterson, and Eglin IRBs) and Army sites in the Great Plains Region (BAMC IRB). Initially, we did not realize that we needed to obtain IRB approval from WRAMC to recruit participants at Ft Drum. The AF policy for installations that do not have their own IRB is that a protocol must be reviewed and approved at one of the AF medical center IRBs (e.g., Wilford Hall). To obtain the IRB approval a signed letter must be obtained from the proposed recruitment site installation medical commander authorizing participation in the study and acknowledging that the medical center IRB approval will be accepted because there is not a local IRB. This is the process that we used to obtain IRB approval to recruit participants at 8 AF sites that did not have a local IRB. In reference to Ft Drum, we had received Army IRB approval from both BAMC and Ft Detrick and obtained a signed letter from the Ft Drum medical commander indicating that he would accept the BAMC/Ft Detrick IRB approvals. Using our 4-page enrollment package that had been approved by BAMC and Ft Detrick, we initiated recruitment at Ft Drum and obtained 14 volunteer participants in the first week of recruitment. However, we were then notified that the BAMC/Ft Detrick IRB approvals would not cover Ft Drum and that we needed to stop recruitment and obtain Walter Reed IRB approval. No randomization or treatment had started for any of the initial Ft Drum participants and they were notified that we would need to delay treatment until we obtained WRAMC IRB approval. As mentioned previously, the WRAMC IRB approval took approximately 12 months to obtain and the length of the enrollment package was increased from 4 to 10 pages. We then re-initiated recruitment at Ft Drum and the longer enrollment packet adversely impacted enrollment. Over the subsequent year, we were only
able to recruit 5 participants at Ft Drum using this 10-page enrollment package. It should be noted that this was a minimal risk study to help volunteer participants to quit tobacco and no medications were used.

**Navy and Marines**

The process of gaining IRB approval from the Navy Hospital IRB for enrollment at identified Navy and Marine bases was quite protracted as well, lasting 9-months. Again we were required to use a lengthy enrollment packet (see attached Appendix C). Recruitment at these sites was also negatively impacted by the longer enrollment forms. A total of 33 participants were enrolled at these two clinics. Table 4 summarizes the number of clinics and the number of subjects by branch of service.

**Utilization of Telephone Counseling Guidelines based on Principles of Motivational Interviewing**

A crucial part of the intervention in this study is the phone call support given by project phone counselors. We continued to implement, refine, and monitor the quality of telephone counseling calls that incorporated Motivational Interviewing (MI) techniques to reinforce participants’ own motivation for quitting smokeless tobacco. We continued to conduct regular supervision sessions with phone counselors to ensure quality and consistency of counseling calls across counselors and across time.

**Utilization of Data Entry/Management System**

In this study, data were collected at various points in time at both research sites – Oregon Research Institute (ORI) in Eugene, OR, and Wilford Hall Medical Center (WHMC) in San Antonio, TX. We collected baseline tobacco use data along with consent information from participants at the various dental clinics at the time of participant enrollment. Those data were forwarded by the clinics to ORI for data entry. We collected various data points from participants in the Treatment Group at the time of telephone counseling conducted out of the WHMC research site. We conducted follow-up survey assessments by mail with all participants, in both Treatment and Control Groups, at 3- and 6-months post enrollment and those assessments were both sent and processed at ORI by project staff. If the participant did not respond to requests to complete the mailed survey, we called them to conduct a telephone survey, using the same questions.

In order to centralize and effectively manage the myriad data collected at both sites, we continued in Year-04 to utilize, and in some cases refine, the Electronic Data Management System developed in Year-02. The system was accessible by research staff at WHMC through a virtual private network (VPN) that completely protected participant confidentiality and allowed WHMC research staff in Texas to accomplish data entry and updates as needed. In addition to being a repository for collected data, the system also served as a scheduling function. It scheduled the dates for the three counseling calls to Treatment participants, as well as dates for follow-up assessments. The evolving database provided the basis for all data analysis procedures to be conducted at the conclusion of the data collection phase.

The database development, expansion, and shared input by staff at WHMC and ORI were key activities in the project. The data entry was all done at ORI where the enrollment data and all follow-up data are stored in secure files by participant number. All MI phone contacts were done by phone counselors and the information they collected was input into the database for that participant. The Virtual Private Network connection, which was developed and implemented in Year-02, allowed the phone counselor to enter key data for the participant into the database. The counselor also accessed the database prior to the call to determine the degree of the participant’s readiness to quit, amount of smokeless tobacco (ST) used, and other
relevant information to use in their motivational interview phone calls. This same system was used in the scheduling and tracking of follow-up assessments at 3- and 6-months.

KEY RESEARCH ACCOMPLISHMENTS IN YEAR 04

- Enrolled 147 participants in Year 04.
- Ended recruitment at all 21 military sites.
- Completed 284 three-month follow-up assessments and 351 six-month assessments during Year 04.
- 242 Counseling Calls made in Year 04.
- Continued to utilize and refine an electronic database accessible to project staff at both the ORI and WHMC research sites for tracking participants and data entry.

PRESENTATIONS:


REPORTABLE OUTCOMES

The 3- and 6-month follow-up data are very promising and we obtained very large effect sizes. At the 3-month follow-up, our ST Cessation Program resulted in a 47% quit rate (7-day point prevalence for all respondents) as compared to only 14% with usual care (p < .001). Using an intent-to-treat analysis in which non-respondents are assumed to have not quit, the cessation rate was still three times higher in the active intervention group (33% versus 11%; p < .001). The quit rates were slightly decreased at the 6-month follow-up, but were still very robust with the active intervention group being twice as likely to have quit (45% versus 22% for all respondents (p < .001); 30% versus 15% using an intent to treat analysis (p < .001)). These results are illustrated in Figure 2 below.

Figure 2. Comparison of Quit Rates
CONCLUSIONS

These results provide strong support for the efficacy of a ST cessation program for active duty military personnel. However, the use of trained phone counselors—which we believe was a key factor in the success of our program—makes the dissemination of this type of program a bit complicated and costly. It may be that similar results could be obtained if this program was implemented without the phone counselor component and if it included only the Enough Snuff Manual and the Tough Enough to Quit DVD. This abbreviated intervention (Book + DVD) could also be strengthened by using a population health approach where all ST users are identified during the annual dental exam and then offered a brief intervention in the dental clinic. The potential population health impact might be even greater if we were to provide training to the dental staff in the use of brief (i.e., < 5 minutes) motivational interviewing and motivational enhancement strategies that could be incorporated into their annual exam. The military healthcare setting is one of the few settings were a true population health intervention study could be successfully employed and evaluated. We plan to seek additional funding from the PRMRRP or the National Cancer Institute to conduct a follow-on study to evaluate brief versus enhanced versions of our smokeless tobacco cessation program and to determine the best method for dissemination of the program(s) throughout the DoD dental community. We have completed the preliminary review of the data from the study and we will now be focused on writing the reports of the study outcomes. We have one paper in progress in which we report the baseline data and characteristics of the sample for the study and we will now be working on analyzing the results for the primary and secondary outcomes of the study. We expect to use the no cost extension of the project to conduct a full data analysis and submit several manuscripts for publication.

REFERENCES

None.
APPENDICES

Enrollment Packets

A. Oregon Research Institute, Eugene OR, Wilford Hall Medical Center, TX, Wright-Patterson AFB, OH, Eglin AFB, FL, and Brooks City Base, TX

B. Water Reed Army Medical Center

C. Naval Medical Center San Diego, CA
APPENDICE A
Enrollment Forms

IRB approved enrollment packet (4-pages) used by:
- Oregon Research Institute;
- Wilford Hall Medical Center;
- Wright-Patterson AFB, OH;
- Eglin AFB, FL; and
- Brooks City Base, TX

1. Consent Form (1-page)
2. HIPAA (1-page)
3. Survey (2-Pages)
Statement of Informed Consent: Smokeless Tobacco Use in Military Personnel

This clinic is taking part in a smokeless tobacco research study. **If you chew tobacco or use snuff, we would like you to participate in this study.** We would like to get as close as possible to 100% of smokeless tobacco users to participate. The purpose of this study is to assess the effectiveness of a brief intervention on smokeless tobacco cessation.

**Your participation involves the following:**

1) **Filling out the attached survey.** You do not need to be ready to quit smokeless tobacco to participate. All information collected in this study will be kept confidential. Only research staff will have access to your information. All data will be stored by Oregon Research Institute, and this information will not be available to your dentist, other health care providers, or to anyone in the military.

2) **Willingness to be contacted by phone at home or work and be offered a smokeless tobacco cessation program.** If you agree to participate, you will be randomly assigned (like the toss of a coin) to the treatment group or the control group. If you are assigned to the treatment group, we will call you to discuss your tobacco use and offer you a cessation program that would help you quit using smokeless tobacco on your own at home. If you are assigned to the control group, you will not receive any phone calls or the cessation program. By agreeing to participate in the study you are not obligated to make a quit attempt even if you are assigned to the treatment group.

3) **You will receive two more surveys by mail similar to the one attached, the first in 3 months and the second in 6 months.**

**Risks**  The risks involved in participating in this study may include:

1) **Loss of confidentiality.** We will be getting personal information from you. Your social security number will be used if needed to locate you for follow-up surveys and to document your phone conversations with counselors if you are assigned to the treatment group. There is always the slight possibility that someone who is not authorized might see the personal information that is requested from you. However, it is extremely unlikely that this will occur, and we will take every precaution to assure that your data remain anonymous.

2) **Discomfort in discussing your use of tobacco.** You may be unaccustomed to talking with someone about your tobacco use. However, our counselors have been extensively trained in this area and will be helpful, courteous, and respectful of your needs.

3) **Withdrawal Symptoms.** If you quit using smokeless tobacco, you may experience withdrawal symptoms from nicotine cravings such as hunger, anxiety, restlessness, or sleep disturbance. These symptoms are common for persons quitting their addiction to tobacco products. Our counselors have been extensively trained in this area and will be helpful, courteous, and respectful of your needs.

**Benefits**  The benefits of participating in this study are:

1) You could receive a free smokeless tobacco quitting program that may enable you to quit without attending classes or medical appointments. Quitting tobacco may be the most important lifestyle change you can make to improve your health.

2) The information you give us may help other military personnel in the future.

? **YES, I am interested in participating in this study.** (You do not need to be ready to quit to participate.)

By signing below I give my consent for the information I provide on the attached survey to be used by scientists at Oregon Research Institute (Eugene, Oregon). I understand that completing this survey is voluntary, I may choose to skip any question, and that this information will be kept confidential.

I also give my consent to being contacted in the future, possibly offered a smokeless tobacco cessation program, and being sent follow-up surveys in the mail.

<table>
<thead>
<tr>
<th>Printed name/with rank</th>
<th>Signature</th>
<th>Date</th>
<th>Social Security Number (Optional)</th>
</tr>
</thead>
<tbody>
<tr>
<td>______________________</td>
<td>__________________</td>
<td>______</td>
<td>____________________________</td>
</tr>
</tbody>
</table>

Home address: ____________________________________________  Home phone: (____)_________________  prefer daytime  prefer after 1700

(Street, City, State and Zip Code)

Email address: ____________________________________________  Work phone: (____)_____________  best time is __morning __ later  best time is after 1600

Put an * next to the number where you prefer to be contacted

? **NO, I am not interested in participating in this study.** AGE _____ years old  SEX: ? Male ? Female
HIPAA AUTHORIZATION TO USE AND DISCLOSE
INDIVIDUAL HEALTH INFORMATION FOR RESEARCH PURPOSES

What is an authorization?
Federal privacy regulations provide safeguards for privacy, security, and authorized access to your personal health information. The Federal Health Insurance & Portability Act (HIPAA) protects the privacy of your personal health information that [INSERT Dental Clinic NAME HERE] and Oregon Research Institute collect in the course of this research study. The only health information that we will access is what you provide in the attached questionnaire. We WILL NOT obtain any information from your dental or medical records. Health information that identifies you may not be used or disclosed to others for research purposes unless you give written permission in advance.

1. Authorization. As a research participant, you authorize Dr. Herbert Severson and Lt Col. Alan Peterson and their research staff to use and disclose your individual health information provided in the attached questionnaire for the purpose of conducting the research project entitled “Population Health Trial for Smokeless Tobacco Cessation with Military Personnel”.

2. Information to be Used or Disclosed. Your individual health information that may be used or disclosed to conduct the study includes: tobacco use history and demographic information such as age, gender, race, rank, education level, marital status, height, and weight.

3. Parties Authorized to Disclose Information. The researcher and the researcher's staff may obtain your individual health information from the attached questionnaire.

4. Parties Who May Receive or Use Information. Your individual health information disclosed in the attached questionnaire and information disclosed by you or discovered about you during the course of the research may be disclosed to (1) the study investigators (Dr. Herbert H. Severson and Lt Col Alan Peterson), (2) the sponsor of this research (the Department of Defense U.S. Army Medical Research and Materiel Command), and (3) the Institutional Review Boards that review this research to make sure that it is ethical (Oregon Research Institute, Wilford Hall Medical Center, and Wright-Patterson Medical Center).

5. Right to Refuse to Sign this Authorization. You understand that you do not have to sign this Authorization. However, if you do not sign this authorization, you will not be allowed to participate in this research study. Your decision not to sign this Authorization will not affect any other treatment, payment, or enrollment in health plans or eligibility for benefits.

6. Right to Withdraw Authorization. You understand you can change your mind and withdraw this Authorization at any time by sending a written notice to Dr. Herbert Severson, 1715 Franklin Blvd, Eugene, OR 97403 or Lt Col Alan Peterson, 59MDOS/MMCP, Wilford Hall Medical Center, 2200 Bergquist Drive, Suite 1, Lackland Air Force Base, TX 78236 to inform the researcher of your decision. If you withdraw this Authorization, the researcher may only use and disclose individual health information already collected for the study. No additional health information about you will be collected by or disclosed to the researcher for the study.

7. Potential Re-disclosure. Only as spelled out in #4 above, or in emergencies and/or as required by state law (in the case of child/elder abuse or intent to harm self or others) will information disclosed to others include information that could be used to identify you. If your information is re-disclosed by them, it may no longer be protected by HIPAA.

8. Expiration of Authorization. This authorization does not have an expiration date.

I am the research participant. I have read this information, and I understand I will receive a copy of this Authorization when it has been signed.

_________________________________________
Name of Participant (type or print)

_________________________________________   ___________________
Signature of Participant       Date
Smokeless Tobacco Study

This clinic is taking part in a smokeless tobacco research project. **If you chew tobacco or use snuff,** we would like you to fill out this survey. Filling it out is voluntary and you may choose to skip any question. If you choose not to complete this survey, it will not affect your health care in any way.

1. Today's date: _/___/___
2. In a typical week, how many days do you use chew/snuff? ○ 1 ○ 2 ○ 3 ○ 4 ○ 5 ○ 6 ○ 7
3. How many days does a can/pouch last you? ○ 1 ○ 2 ○ 3 ○ 4 ○ 5 ○ 6 ○ 7 or more
4. How soon after you wake up do you use chew/snuff? ○ Less than 30 min. ○ 30 to 60 min. ○ More than 60 min.
5. How old were you when you began using chew/snuff? __ years old
6. Do you swallow tobacco juice on purpose? ○ Never ○ Sometimes ○ Almost always
7. Have you tried to quit using chew/snuff in the last year? ○ Yes ○ No
8. These statements show how some chew/snuff users feel about quitting. Mark the number that shows how you feel:

<table>
<thead>
<tr>
<th>Not ready to quit</th>
<th>Should consider quitting some day</th>
<th>Should quit but not quite ready</th>
<th>Thinking about cutting down or quitting</th>
<th>Have cut down and seriously considering quitting</th>
<th>Ready to quit now</th>
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<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
</tbody>
</table>

9. How many of your five best friends use chew or snuff? ○ None ○ 1 ○ 2 ○ 3 ○ 4 ○ 5

Please complete the questions on the back of this page. Thank you for your help!
10. Do you currently smoke?  ○ Yes  ○ No

11. On a typical day, how many cigarettes do you smoke?
   ○ None  ○ 1 to 5  ○ 6 to 10  ○ 11 to 15  ○ 16 to 20  ○ 21 to 25  ○ 26 to 30  ○ 31 or more

12. Does your spouse/partner smoke?  ○ Yes  ○ No  ○ Does not apply

13. Have you had two or more years in your life when you felt depressed or sad most days, even if you felt okay sometimes?
   ○ Yes  ○ No

14. In the past year, have you had two weeks or more during which you felt sad, blue, or depressed, or when you lost all interest or pleasure in things that you usually cared about or enjoyed?
   ○ Yes  ○ No

15. In the past seven days, how many drinks of alcohol did you have? (one drink equals a 12-ounce glass of beer or 6-ounce glass of wine or one shot of liquor)
   ○ None  ○ 1-3  ○ 4-6  ○ 7-9  ○ 10-12  ○ 13-15  ○ 16-18  ○ 19 or more

16. How tall are you?  □ □ feet  □ □ inches

17. How much do you weigh?  □ □ □ pounds

18. Your age:  □ □ years old

19. Sex:  ○ Male  ○ Female

20. Do you consider yourself to be Hispanic or Latino?  ○ Yes  ○ No

21. What race do you consider yourself to be?  Select one of the following:
   ○ American Indian or Alaska Native  ○ Asian  ○ Black or African American  ○ Native Hawaiian or other Pacific Islander  ○ White

22. Marital Status:  ○ Single  ○ Married or living with partner

23. Education finished:
   ○ Less than high school degree  ○ High school graduate or equivalent  ○ Some college  ○ College graduate  ○ Post college

You are finished. Thank you for your help!
APPENDICES B
Enrollment Forms

IRB approved enrollment packet (10-pages) – Water Reed Army Medical Center

1. Consent Form (6-pages)
2. HIPAA (2-pages)
3. Survey (2-Pages)
VOLUNTEER AGREEMENT AFFIDAVIT

For use of this form, see AR 70-25 or AR 40-38; the proponent agency is OTSG

PRIVACY ACT OF 1974

Authority: 10 USC 3013, 44 USC 3101, and 10 USC 1071-1087.

Principle Purpose: To document voluntary participation in the Clinical Investigation and Research Program. SSN and home address will be used for identification and locating purposes.

Routine Uses: The SSN and home address will be used for identification and locating purposes. Information derived from the study will be used to document the study; implementation of medical programs; adjudication of claims; and for the mandatory reporting of medical conditions as required by law. Information may be furnished to Federal, State and local agencies.

Disclosure: The furnishing of your SSN and home address is mandatory and necessary to provide identification and to contact you. If future information indicates that your health may be adversely affected. Failure to provide the information may preclude your voluntary participation in this investigational study.

PART A(1) - VOLUNTEER AFFIDAVIT

Volunteers under the provisions of AR 40-38 and AR 70-25 are authorized all necessary medical care for injury or disease which is the proximate result of their participation in such studies.

I, _______________________________ SSN _______________________________, having full capacity to consent and having attained my _______________________________ birthday, do hereby volunteer/give consent as legal representative for _______________________________ to participate in _______________________________.

Population Health Trial for Smokeless Tobacco Cessation Among Military Personnel

under the direction of Jane Bowers, Community Dental Health Hygienist, Fort Drum, New York

conducted at USA DENTAL ACTIVITY, FORT DRUM, NEW YORK

(Name of institution)

The implications of my voluntary participation/consent as legal representative; duration and purpose of the research study; the methods and means by which it is to be conducted; and the inconveniences and hazards that may reasonably be expected have been explained to me by:

Jane Bowers, or her designee, Community Dental Health Hygienist, Fort Drum, New York, phone 315-772-7841JS7.

I have been given an opportunity to ask questions concerning this investigational study. Any such questions were answered to my full and complete satisfaction. Should any further questions arise concerning my rights/the rights of the person I represent on study-related injury, I may contact:

JUDGE ADVOCATE GENERAL (JAG) OFFICE, PHONE 315-772-5261

at FORT DRUM, NEW YORK

(Name, Address and Phone Number of Hospital Include Area Code)

I understand that I may at any time during the course of this study revoke my consent and withdraw/have the person I represent withdrawn from the study without further penalty or loss of benefits; however, if the person I represent may be required (military volunteer) or requested (civilian volunteer) to undergo certain examination if, in the opinion of the attending physician, such examinations are necessary for my/the person I represent's health and well-being. My/the person I represent's refusal to participate will involve no penalty or loss of benefits to which the person I represent is otherwise entitled.

LIMITATIONS TO MEDICAL CARE ARE DESCRIBED IN PART B

PART A (2) - ASSENT VOLUNTEER AFFIDAVIT (MINOR CHILD)

I, _______________________________ SSN _______________________________, having full capacity to consent and having attained my _______________________________ birthday, do hereby volunteer for _______________________________ to participate in _______________________________.

under the direction of _______________________________

Conducted at

(Name of institution)

(Continue on Reverse)

DA FORM 5303-R, MAY 89

PREVIOUS EDITIONS ARE OBSOLETE
PART A(2) - ASSENT VOLUNTEER AFFIDAVIT (MINOR CHILD) (Cont'd)

The implications of my voluntary participation: the nature, duration, and purpose of the research study, the methods and means by which it is to be conducted, and the inconveniences and hazards that may reasonably be expected have been explained to me by

I have been given an opportunity to ask questions concerning this investigational study. Any such questions were answered to my full and complete satisfaction. Should any further questions arise concerning my rights I may contact

at

[Name, Address, and Phone Number of Hospital (Include Area Code)]

I understand that I may at any time during the course of this study revoke my assent and withdraw from the study without further penalty or loss of benefits, however, I may be requested to undergo certain examinations if, in the opinion of the attending physician, such examinations are necessary for my health and well-being. My refusal to participate will involve no penalty or loss of benefits to which I am otherwise entitled.

LIMITATIONS TO MEDICAL CARE ARE DESCRIBED IN PART B

PART B - TO BE COMPLETED BY INVESTIGATOR

INSTRUCTIONS FOR ELEMENTS OF INFORMED CONSENT: (Provide a detailed explanation in accordance with Appendix C, AR 40-28 of AR 70-25)

DESCRIPTION OF THIS STUDY

You are being asked to be in this research study because you answered that you currently use smokeless tobacco at your annual dental exam. Your participation is entirely voluntary. Refusal to participate will not result in any penalty or loss of benefits to which you are otherwise entitled.

The purpose of this study is to assess the effectiveness of a brief intervention on smokeless tobacco cessation with military personnel.

Other studies have shown that telephone support calls from a trained counselor, together with self-help cessation materials sent by mail, is very effective in helping smokeless tobacco users to quit. This study is based upon a published study with military personnel that used the same procedures. A study has reported that these procedures have also been effective with civilian smokeless tobacco users.

If you agree to be in this study, you will be asked: (1) to fill out the attached survey. This survey includes questions regarding your tobacco and alcohol use as well as your demographics (information such as your age, race, education level, and marital status). You do not need to be ready to quit smokeless tobacco to fill out the attached initial survey; (2) to be contacted by phone at home or work and be offered a smokeless tobacco cessation program. If you agree to participate by signing this form, you will be randomly assigned (similar to the flip of a coin) to one of two groups, the Treatment group or the Control group. Your chances of being assigned to each group are equal.

I do □ do not □ (check one & initial) consent to the inclusion of this form in my outpatient medical treatment record.

SIGNATURE OF VOLUNTEER DATE SIGNATURE OF LEGAL GUARDIAN (If volunteer is a minor)

PERMANENT ADDRESS OF VOLUNTEER

TYPOGRAPHICAL NAME OF WITNESS

SIGNATURE OF WITNESS DATE

REVERSE OF DA FORM 5303-R, MAY 89
If you are assigned to the Treatment group, you will receive up to three phone calls. The first call attempt will be made within 1-month upon Oregon Research Institute (ORI, the non-profit research organization conducting this study) receiving the completed enrollment packet (this consent form, the HIPAA authorization form, and the initial survey). Phone counselors, hired and trained by ORI, who have been extensively trained in the area of smokeless tobacco cessation will be making these calls. In the first call you will be offered a smokeless tobacco cessation program that will be mailed to your home. The program includes a 25-minute video tape and a self-help cessation manual. Two additional supportive phone calls will be made to discuss the cessation program, your current tobacco use and how you might apply the program to your own use. The second phone call will be attempted beginning 3-weeks following the date materials are mailed. The third phone call will be attempted either 2-weeks after the second phone call or 2-days following your quit date, should you choose to set a quit date during the second phone call. Each phone call will last up to 20-minutes and all three phone calls will be completed within 3-months of enrolling in this study. If you are assigned to the Control group, you will not receive any phone calls or the cessation program. **By agreeing to participate in the study and completing the initial survey, you are not obligated to make a quit attempt even if you are assigned to the Treatment group.**

Both Treatment and Control groups will be asked to complete and return two more surveys, by mail, the first in 3-months following enrollment and the second in 6-months. These surveys will be mailed to you and will each take 20-minutes to complete. You will be asked to return the completed survey in the provided stamped, addressed envelope. These surveys include questions regarding your health and tobacco use. If you are assigned to the Treatment group and have received the video and manual, each of these surveys will also ask for your feedback about the materials as well as your reaction to the phone counselor.

Both Treatment and Control group participants in the study who do not return the 3-month survey by mail will be contacted by phone and asked to complete the survey over the phone.

**AMOUNT OF TIME FOR YOU TO COMPLETE THIS STUDY**

You will be part of this study for a total of 6-months. The three surveys will take approximately 15-minutes each to complete. If you are assigned to the Treatment group and choose to receive the intervention, you will receive three phone calls, lasting up to 20-minutes each, a video (25-minutes long) and a manual for viewing.

**APPROXIMATE NUMBER OF PEOPLE TAKING PART IN THIS STUDY**

This study is called a multi-service study because participants will be recruited from dental clinics located at all four US military branches. There will be up to 600 people taking part in this study at Fort Drum. A total of 1,600 people will be in the study from all of the dental clinics involved.

**POSSIBLE RISKS OR DISCOMFORTS FROM BEING IN THIS STUDY**

The risks involved in participating in this study may include:

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<td>PERMANENT ADDRESS OF VOLUNTEER</td>
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<td>SIGNATURE OF WITNESS</td>
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REVERSE OF DA FORM 5303-R, MAY 89
(1) **Loss of confidentiality.** We will be getting personal information from you. Your social security number will be used if needed to locate you for follow-up surveys and to document your phone conversations with counselors if you are assigned to the Treatment group. There is always the slight possibility that someone who is not authorized might see the personal information that is requested from you. However, it is extremely unlikely that this will occur, and we will take every precaution to assure that your information remains confidential.

(2) **Discomfort in discussing your use of tobacco.** You may be unaccustomed to talking with someone about your tobacco use. However, our counselors have been extensively trained in this area and will be helpful, courteous, and respectful of your needs.

There are no other expected risks or discomforts from being in this study.

**POSSIBLE BENEFITS OF BEING IN THIS STUDY**

The possible benefits to you from being in this study are (a) you could receive a free smokeless tobacco quitting program that may enable you to quit without attending classes or medical appointments; and (b) quitting tobacco may be the most important lifestyle change you can make to improve your health. However, no benefit can be guaranteed. The information you give us, whether you are assigned to the Treatment or Control group, will help us compare this program to the options already provided to military personnel to assist them in quitting smokeless tobacco use.

**CONFIDENTIALITY (PRIVACY) OF YOUR IDENTITY AND YOUR RESEARCH RECORDS**

The principal investigator will keep records of your being in this study. All information collected in this study will be kept confidential. All datafiles will be maintained by Oregon Research Institute in Eugene, Oregon. This information will not be available to your dentist, other health care providers, or to anyone in the military. These records may be looked at by people from the Walter Reed Department of Clinical Investigation, the Walter Reed Human Use Committee, the Army Clinical Investigation Regulatory Office (CIRO), the United States Army Medical Research and Materiel Command (USAMRMC, the sponsor of this study), the Institutional Review Boards (committees that review this research to make sure that it is ethical) at Wilford Hall Medical Center (a study site), Wright-Patterson Medical Center (a study site), and the Oregon Research Institute, and other government agencies as part of their duties. These duties include making sure that research subjects are protected. Confidentiality of your records will be protected to the extent possible under existing regulations and laws. Complete confidentiality cannot be promised, particularly for military personnel, because information bearing on your health may be required to be reported to appropriate medical or command authorities. Your name will not appear in any published paper or presentation related to this study.

Every precaution will be taken to assure that the data files remain confidential. Upon completing this consent form and the initial survey, you will be assigned a personal identification number. The assigned personal identification number will not contain any part of your social security number. The consent and survey will

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REVERSE OF DA FORM 5303-R, MAY 89
be mailed separately from the dental clinic to the Oregon Research Institute (ORI) in Eugene, Oregon where the data will be entered. The data will be stripped of personal identifiers and will be labeled with only your assigned personal identification number. The master list linking your personal identifying information with your assigned personal identification number will be kept in a separate, password encrypted data file to which only the Principal Investigator at ORI, Dr. Severson, and selected professional staff will have access. This master list will be maintained at ORI. Data files will also be password encrypted. All computer data are protected from unauthorized access by industry standard encryption and firewall techniques. All hard copies of data will be kept in locked files. Reports describing the results of this study will in no way reveal your identity. All personnel who have access to your information are instructed not to talk about the study participants publicly by name, and refer to them either by number or first name only. Such procedures are made part of the training and are overemphasized to ensure that these safeguards are maintained. All employees of ORI are asked to sign a Certificate of Confidentiality before being employed with the organization.

This research study meets the confidentiality requirements of the Health Insurance Portability and Accountability Act (HIPAA). A HIPAA authorization form for this study will be provided to you separately, and you will be asked to sign that form.

CONDITIONS UNDER WHICH YOUR TAKING PART IN THIS STUDY MAY BE STOPPED WITHOUT YOUR CONSENT

Your taking part in this study may be stopped without your consent if remaining in the study might be dangerous or harmful to you. Your taking part in this study may also be stopped without your consent if the military mission requires it, or if you become ineligible for medical care at military hospitals.

ELIGIBILITY AND PAYMENT FOR BEING IN THIS STUDY

Any active duty military personnel who indicate they are a current smokeless tobacco user is eligible for this study. You will not receive any payment for being in this study.

COMPENSATION TO YOU IF INJURED AND LIMITS TO YOUR MEDICAL CARE

Should you be injured as a direct result of being in this study, you will be provided medical care for that injury at no cost to you. You will not receive any compensation (payment) for injury. You should also understand that this is not a waiver or release of your legal rights. You should discuss this issue thoroughly with the principal investigator before you enroll in this study.

Medical care is limited to the care normally allowed for Department of Defense health care beneficiaries (patients eligible for care at military hospitals and clinics).

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REVERSE OF DA FORM 5303-R. MAY 89
WHAT WILL HAPPEN IF YOU DECIDE TO STOP TAKING PART IN THIS STUDY AND INSTRUCTIONS FOR STOPPING EARLY

You have the right to withdraw from this study at any time. If you decide to stop taking part in this research study, send a written notice to Dr. Herbert Severson, 1715 Franklin Blvd, Eugene, OR 97403 to inform the researchers of your decision to withdraw from this study. If you withdraw from this study, you will no longer be contacted. Once you withdraw from the study, the researchers may only use and disclose individual health information already collected for the study. No additional health information about you will be collected by or disclosed to the researchers for the study. By leaving this study at any time, you in no way risk losing your right to medical care.

☐ YES, I am interested in participating in this study. (You do not need to be ready to quit to participate.)

Home address: _______________________________ Rank: _______________

Home phone: _____________________ □ prefer daytime □ prefer after 1700

Work phone: ________________________ □ best time is ___A.M. ___P.M. □ best time is after 1600

Put an * next to the number where you prefer to be contacted

Email address: ____________________________

☐ No, I am not interested in participating in this study. AGE _____ years old Sex: □ Male □ Female

Please feel free to ask any questions that will allow you to clearly understand this study.

A copy of this consent form will be provided to you.

SIGNATURE OF VOLUNTEER DATE SIGNATURE OF LEGAL GUARDIAN (if volunteer is a minor)

PERMANENT ADDRESS OF VOLUNTEER TYPED NAME OF WITNESS

SIGNATURE OF WITNESS DATE

REVERSE OF DA FORM 5303-R, MAY 89
Protocol Title: Population Health Trial for Smokeless Tobacco Cessation Among Military Personnel

Principal Investigator: Jane Bowers

Work Unit #: 04-82001

The Federal Health Insurance Portability and Accountability Act (HIPAA) includes a Privacy Rule that gives special safeguards to Protected Health Information (PHI) that is identifiable, in other words, can be directly linked to you (for example, by your name, Social Security Number, birth date, etc.). We are required to advise you how your PHI will be used.

1. What information will be collected?

For this research study, we will only be collecting personal health information from participants. This information includes: tobacco use history and demographic information such as age, gender, race, rank, education level, marital status, height, and weight.

2. Who may use my PHI within the Military Healthcare System?

Your PHI disclosed in the attached questionnaire and information disclosed by you or discovered about you during the course of the research may be disclosed to (1) the study investigators, (2) the sponsor of this research (the Department of Defense U.S. Army Medical Research and Materiel Command), and (3) the Institutional Review Boards that review this research to make sure that it is ethical (Walter Reed Army Medical Center Department of Clinical Investigation and Human Use Committee, Wilford Hall Medical Center, and Wright-Patterson Medical Center).

3. What persons outside of the Military Healthcare System who are under the HIPAA requirements will receive my PHI?

Your PHI disclosed in the attached questionnaire and information disclosed by you or discovered about you during the course of the research may be disclosed to (1) the study investigator, Dr. Herbert H. Severson of the Oregon Research Institute, (ORI) in Eugene, Oregon, (2) the research project staff, and (3) the ORI Institutional Review Board that reviews this research to make sure that it is ethical.

4. What is the purpose for using or disclosing my Protected Health Information (PHI)?

The investigators of this research project need to use your PHI in order to analyze the information to assess the effectiveness of a brief intervention on smokeless tobacco cessation.

5. How long will the researchers keep my Protected Health Information?

Data will be maintained by ORI for five years after the completion of this research study at which time all documents and data files will be destroyed.

6. Can I review my own research information?

You may look at your personal research information at any time.

7. Can I cancel this Authorization?

Yes. If you cancel this Authorization and stop taking part in this research study, you will no longer be contacted. You can change your mind and withdraw this Authorization at any time by sending a written notice to Dr. Herbert Severson, 1715 Franklin Blvd, Eugene, OR 97403 to inform the researchers of your decision to withdraw from this study. Once you withdraw from the study, the researchers may only use and disclose individual health information already collected for the study. No additional health information about you will be collected by or disclosed to the researchers for the study.
8. What will happen if I decide not to sign this Authorization?

You understand that you do not have to sign this Authorization. However, if you do not sign this authorization, you will not be allowed to participate in this research study. Your decision not to sign this Authorization will not affect any other treatment, payment, or enrollment in health plans or eligibility for benefits.

9. Can my Protected Health Information be disclosed to parties not included in this Authorization who are not under the HIPAA requirements?

There is a potential that your research information will be shared with another party not listed in this Authorization in order to meet legal or regulatory requirements. Examples of persons who may access your PHI include representatives of the Army Clinical Investigation Regulatory Office, the Food and Drug Administration, the DHHS Office for Human Research Protections, and the DHHS Office for Civil Rights. This disclosure is unlikely to occur, but in that case, your health information would no longer be protected by the HIPAA Privacy Rule.

10. Who should I contact if I have any complaints?

If you believe your privacy rights have been violated, you may file a written complaint with the WRAMC Privacy Officer, 6900 Georgia Ave., NW, Washington, DC 20307. Telephone: 202-782-3501.

The signature below acknowledges receipt of this Authorization:

Signature: _____________________________ Date: ________________

If you are a parent, court-appointed representative, or acting as power of attorney, indicate your authority to act for the participant: _____________________

Print Name: _____________________________

A copy of this signed Authorization will be provided to you. 7/21/03
Smokeless Tobacco Study

This clinic is taking part in a smokeless tobacco research project. If you chew tobacco or use snuff, we would like you to fill out this survey. Filling it out is voluntary and you may choose to skip any question. If you choose not to complete this survey, it will not affect your health care in any way.

IMPORTANT: USE BLACK OR BLUE INK PEN
Shade Circles Like This
Not Like This

For optimum accuracy, please print carefully and avoid contact with the edges of the box.
The following will serve as an example:

1. Today's date:  / /
m m d d y y

2. In a typical week, how many days do you use chew/snuff? ○ 1 ○ 2 ○ 3 ○ 4 ○ 5 ○ 6 ○ 7

3. How many days does a can/pouch last you? ○ 1 ○ 2 ○ 3 ○ 4 ○ 5 ○ 6 ○ 7 or more

4. How soon after you wake up do you use chew/snuff? ○ Less than 30 min. ○ 30 to 60 min. ○ More than 60 min.

5. How old were you when you began using chew/snuff?   years old

6. Do you swallow tobacco juice on purpose? ○ Never ○ Sometimes ○ Almost always

7. Have you tried to quit using chew/snuff in the last year? ○ Yes ○ No

8. These statements show how some chew/snuff users feel about quitting. Mark the number that shows how you feel:

<table>
<thead>
<tr>
<th>Not ready to quit</th>
<th>Should consider quitting some day</th>
<th>Should quit but not quite ready</th>
<th>Thinking about cutting down or quitting</th>
<th>Have cut down and seriously considering quitting</th>
<th>Ready to quit now</th>
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</tr>
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</table>

9. How many of your five best friends use chew or snuff? ○ None ○ 1 ○ 2 ○ 3 ○ 4 ○ 5

Please complete the questions on the back of this page. Thank you for your help!
10. Do you currently smoke?  ○ Yes  ○ No

11. On a typical day, how many cigarettes do you smoke?
   ○ None  ○ 1 to 5  ○ 6 to 10  ○ 11 to 15  ○ 16 to 20  ○ 21 to 25  ○ 26 to 30  ○ 31 or more

12. Does your spouse/partner smoke?  ○ Yes  ○ No  ○ Does not apply

13. Have you had two or more years in your life when you felt depressed or sad most days, even if you felt okay sometimes?
   ○ Yes  ○ No

14. In the past year, have you had two weeks or more during which you felt sad, blue, or depressed, or when you lost all interest or pleasure in things that you usually cared about or enjoyed?
   ○ Yes  ○ No

15. In the past seven days, how many drinks of alcohol did you have? (one drink equals a 12-ounce glass of beer or 6-ounce glass of wine or one shot of liquor)
   ○ None  ○ 1-3  ○ 4-6  ○ 7-9  ○ 10-12  ○ 13-15  ○ 16-18  ○ 19 or more


18. Your age: □ □ years old  19. Sex:  ○ Male  ○ Female

20. Do you consider yourself to be Hispanic or Latino?  ○ Yes  ○ No

21. What race do you consider yourself to be? Select one of the following:
   ○ American Indian  ○ Asian  ○ Black or African  ○ Native Hawaiian or
   or Alaska Native  African American  other Pacific Islander  ○ White

22. Marital Status:  ○ Single  ○ Married or living with partner

23. Education finished:
   ○ Less than high  ○ High school graduate  ○ Some college  ○ College graduate  ○ Post college
   school degree  or equivalent

You are finished. Thank you for your help!
APPENDICE C

Enrollment Forms

IRB approved enrollment packet (10-pages) – Naval Medical Center San Diego, CA

1. Consent Form (4-pages)
2. California Experimental Subjects Bill of Rights (1-page)
3. Privacy Act Statement (1-page)
4. HIPAA (2 pages)
5. Survey (2-Pages)
1st DENTAL BATTALION NAVAL DENTAL CENTER  
CAMP PENDLETON  
CAMP PENDLETON, CA 92055-5221

CONSENT BY A SUBJECT FOR VOLUNTARY PARTICIPATION IN A CLINICAL INVESTIGATION (RESEARCH) STUDY

1. You, ______________________, have been asked to voluntarily participate in a research project entitled, "Population Health Trial for Smokeless Tobacco Cessation Among Military Personnel" being conducted at the Naval Medical Center, San Diego by medical researchers from the 1st Dental Battalion Naval Dental Center Camp Pendleton, in collaboration with (1) Oregon Research Institute, (2) the sponsor of this research (the Department of Defense U.S. Army Medical Research and Materiel Command).

2. WHY IS THE STUDY BEING DONE?  
The purpose of this research project is to measure the effectiveness of a brief smokeless tobacco cessation intervention program (designed specifically for use in the Military) compared to the usual preventative health care provided.

3. HOW LONG WILL YOU BE PARTICIPATING IN THE STUDY?  
Once you agree to participate in the study you will begin by completing the initial survey. This starts the timer for your involvement. All participants will receive two follow up surveys by mail, the first in three months and the second at six months.

4. WHAT IS INVOLVED IN THE STUDY?  
Once you have signed this consent form and have completed the attached baseline survey (2-pages) you will be randomly assigned by chance (like the toss of a coin) to the treatment group or the control group. Neither your doctor, the researcher for this study, nor you will be able to choose the group to which you will be assigned. If you are assigned to the treatment group, we will call you to discuss your tobacco use and offer you a cessation program that would help you quit using smokeless tobacco on your own at home. The program involves receiving a smokeless tobacco cessation guidebook, video and three counseling phone calls. By agreeing to participate in the study and completing the initial survey, you are not obligated to make a quit attempt even if you are assigned to the treatment group. If you are assigned to the control group, you will not receive any phone calls or the cessation program. Again, all participants will be mailed a three month and six month follow up survey to complete and return.

5. WHAT IS THE EXPERIMENTAL PART OF THE STUDY?  
The experimental part of this study is to test the intervention program design as well as the effectiveness of the intervention materials designed specifically for use with military personnel in both the short (three month) and long term (six month).

Subject's Initials: ______

IRB Approval Stamp Required  
(no alterations should be made to this document w/out prior approval.)

Page 1 of 4       June 17, 2004
6. **HOW MANY PEOPLE WILL TAKE PART IN THE STUDY?**
A total of 1,600 subjects are expected to participate in this study, of whom up to 200 participants may be from the Naval Medical Center, San Diego.

7. **WHAT ARE THE RISKS OF THE STUDY?**
The risks involved in participating in this study may include:

   1) **Loss of confidentiality.** We will be getting personal information from you. Your social security number will be used if needed to locate you for follow-up surveys and to document your phone conversations with counselors if you are assigned to the treatment group. There is always the slight possibility that someone who is not authorized might see the personal information that is requested from you. However, it is extremely unlikely that this will occur, and we will take every precaution to assure that your data remain anonymous.

   2) **Discomfort in discussing your use of tobacco.** You may be unaccustomed to talking with someone about your tobacco use. However, our counselors have been extensively trained in this area and will be helpful, courteous, and respectful of your needs.

   3) **Withdrawal Symptoms.** If you quit using smokeless tobacco, you may experience withdrawal symptoms from nicotine cravings such as hunger, anxiety, restlessness, or sleep disturbance. These symptoms are common for persons quitting their addiction to tobacco products. Our counselors have been extensively trained in this area and will be helpful, courteous, and respectful of your needs.

8. **ARE THERE BENEFITS TO TAKING PART IN THE STUDY?**
If assigned to the treatment group, the benefits of participating in this study are: (a) you would receive a free smokeless tobacco quitting program that may enable you to quit without attending classes or medical appointments and quitting tobacco may be the most important lifestyle change you can make to improve your health; (b) the information you give us may help other military personnel in the future.

   If assigned to the control group, your participation in this research project may not be of direct benefit to you personally. However, the results of this study may help the investigators gain important knowledge about the effectiveness of this smokeless tobacco quitting program as well as aid in determining the availability of such programs for smokeless tobacco users in the future.

9. **WHAT OTHER OPTIONS ARE THERE?**
This research study is not designed to treat any medical condition that you may have. Therefore, there are no alternative procedure(s) or course of treatment that would be advantageous to you.

10. **WILL I BE PAID TO PARTICIPATE?**
You will not be financially compensated for your participation in this study.

Subject’s Initials: _____

**IRB Approval Stamp Required**
(no alterations should be made to this document w/out prior approval)

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11. WHAT IF I AM INJURED AS A RESULT OF PARTICIPATION IN THIS STUDY?
If you suffer any injury directly related to your participation in this research study, immediate
medical attention is available at the Naval Medical Center, San Diego, or at another closer
medical treatment facility, if applicable. Any injury resulting from your participation in this study
will be evaluated and treated in keeping with the benefits or care to which you are entitled under
applicable Navy, other Department of Defense, and other state or Federal regulations.

12. WHAT ABOUT CONFIDENTIALITY?
In all publications and presentations resulting from this research study, information about you or
your participation in this project will be kept in the strictest confidence and will not be released
in any form identifiable to you personally. However, authorized personnel from the Navy
Medical Department and from the Food and Drug Administration (FDA), where applicable, may
have access to your research file in order to verify that your rights have been adequately
protected.

13. WHOM DO I CALL IF I HAVE QUESTIONS OR PROBLEMS?
If you have any questions regarding this research study, you may contact CAPT Wayne J.
Osborne, at 760-725-8994.

If you have any questions about your rights as an individual while participating in a research
study at the Naval Medical Center, San Diego, you may contact CAPT George Ulrich, MC,
USN, Chairman, Institutional Review Board at (619) 532-8125, or Dr. Warren Lockette,
Head, Clinical Investigation Department at (619) 532-8127.

If you believe that you have been injured as a result of your participation in this research study,
you may contact CDR Steve Bannow, JAGC, USN, Naval Medical Center, San Diego, Legal
Department, at (619) 532-6475.

14. WHAT ARE MY RIGHTS AS A PARTICIPANT?
Your participation in this project is entirely voluntary and your decision not to participate will
involve no penalty or loss of benefits to which you are entitled under applicable regulations. If
you choose to participate, you are free to ask questions or to withdraw from the study at any
time. If you should decide to withdraw from the research project, you will notify CAPT Wayne
J. Osborne, 1st Dental Battalion Naval Dental Center Camp Pendleton, Camp Pendleton,
CA 92055-5221, (760) 725-8994 by sending a written notice to inform the researchers of
your decision. You may also contact them by phone to ensure your timely removal from the
study. If you withdraw this Authorization, the researcher may only use and disclose individual
health information already collected for the study. No additional health information about you
will be collected by or disclosed to the researcher for the study. Your withdrawal will involve no
prejudice to your future health care or any loss of rights or benefits to which you are otherwise
entitled. Any new significant finding developed during the course of this study, which might
affect your willingness to continue participation will be communicated to you.
15. CAN I BE TERMINATED FROM THE STUDY?
The investigator may terminate your participation in this study for the following reasons:
Participant is deployed for an undisclosed period of time during any phase of the study.

16. SIGNATURE

You are making a decision whether or not to participate in the research project above. Your
signature indicates that you have had this information presented to you, have had the
opportunity to ask questions about the research and your participation, and agree to participate
in the study. Further, your signature indicates that you have been provided with a copy of this
consent document and a copy of a document entitled, "California Experimental Subject's Bill of
Rights."

☐ YES, I am interested in participating in this study. (You do not need to be ready to quit to
participate.)

SIGNATURES AND DATE SIGNED:        PRINTED OR TYPED IDENTIFICATION:

Patient / Subject   (Date)          Name / Status / Sponsor's SSN
Witness            (Date)          Name / Grade or Rank
Researcher/Investigator   (Date)  Name / Grade or Rank

Home address:________________________________________________________

House/apt number, street______________________________________________

City, State, Zip Code________________________________________________

Email address:________________________________________________________

Home phone:________________________prefer daytime
prefer after 1700
best time is ___ morning ___ later

Work phone:________________________ best time is after 1800

Put an * next to the number where you prefer to be contacted

☐ No, I am not interested in participating in this study. AGE____ years old SEX: ☐ Male ☐ Female

Subject's Initials:________

IRB Approval Stamp Required
(no alterations should be made to this document w/out prior approval)

Page 4 of 4       June 17, 2004
PATIENT AUTHORIZATION TO USE AND/OR DISCLOSE PROTECTED HEALTH INFORMATION FOR RESEARCH (HIPAA)
(In Keeping with the Health Insurance Portability and Accountability Protection Act)

What is Confidentiality of records all about?

The Naval Medical Center San Diego makes every effort to maintain the confidentiality of protected health information we obtain about you. However, we cannot absolutely guarantee confidentiality because other people may need to see your information in the course of this research study. Most people and organizations will protect the privacy of your information, but may not be required to do so by the law. Also, if the results of this research study are presented at meetings or are published, your name will not be used.

What is HIPAA all about?

The Health Insurance Portability and Accountability Act (HIPAA) requires that we get your permission to use protected health information about you that is either created by or used in connection with this research study. This permission is called an Authorization. The information we use includes your entire research record and supporting information from your medical records, results of laboratory test, X-rays, MRIs, CT scans and observations made by a physician or nurse which are both clinical and research in nature.

What will we do with this information?

Your protected health information will be collected and used during the course of the research study, to monitor your health status, to measure the effects of drugs or devices or procedures, to determine research results, and to possibly develop new tests, procedures, and commercial products.

Your research doctor will use this information to report the results of research to sponsors and federal agencies, like the Food and Drug Administration (FDA). The information may also be reviewed when the research study is audited for compliance. When the study is over, you have the right to see the information and copy it for your records.

Who will we share your information with?

Your information may be shared with any of the following:

- The sponsor of the study, or its agents, such as data repositories
- Other medical centers, institutions, or research investigators outside of the Naval Medical Center San Diego, participating in this research study
- State and Federal agencies which have authority over the research, the Naval Medical Center San Diego or patients. Good examples are: the Department of Health and Human Services (DHHS), the Food and Drug Administration (FDA), the National Institute of Health (NIH), the Office of Human Research Protections (OHRP), and the Department of Social Services (DSS) or other.
- This hospital or clinic.
- Accrediting agencies, such as JCAHO.
- A data safety monitoring board, if applicable
- Clinical staff who may not be involved directly in the research study, but who may become involved in your care, if it is possibly related to treatment
For this research study, the study investigator may share this authorization form and records which identify you to comply with regulatory requirements or for purposes related to this research to:

All documented Principal, Associate, and Sub-investigators, and the Medical Monitor (if one is assigned).

What if you want to revoke or cancel away your Authorization?

If you decide to participate in this research study, your Authorization for this study will not expire unless you revoke or cancel it in writing to the research doctor. If you revoke your Authorization, you will also be removed from the study, but standard medical care and any other benefit to which you are entitled will not be affected in any way.

Revoking your Authorization only affects the use and disclosure (sharing) of information after your written request has been received. Federal law requires sending study information to the FDA for studies it regulates, like studies of drugs and devices. In a case like this, your information may need to be reported to them and cannot be removed from the research records once it is collected.

Do you have to sign this form?

You have the right to refuse to sign this Authorization form and not be a part of this study. You can also tell your study doctor you want to withdraw from the study at any time without revoking the Authorization to use your health information. By signing this research Authorization form, you authorize the use and/or disclosure of your protected health information described above.

**SIGNATURE AND DATE SIGNED:**

Patient/Subject ____________________________  (Date)

Witness ____________________________________ (Date)

Researcher/investigator ______________________  (Date)

**PRINTED OR TYPED IDENTIFICATION:**

Name/Status/Sponsor’s SSN

Name/Grade or Rank

Name/Grade or Rank
PRIVACY ACT STATEMENT

1. Authority. 5 USC 301

2. Purpose. Medical research information will be collected to enhance basic medical knowledge or to develop tests, procedures, and equipment to improve the diagnosis, treatment, or prevention of illness, injury, or functional impairment.

3. Use. Medical research information will be used for statistical analysis and reports by the Department of the Navy, the Department of Defense, and other U.S. Government agencies, provided this use is compatible with the purpose for which the information was collected. Use of the information may be granted to non-Government agencies or individuals by the Chief, Bureau of Medicine and Surgery in accordance with the provisions of the Freedom of Information Act.

4. Disclosure. I understand that all information contained in this Consent Statement or derived from the medical research study described herein will be retained permanently at Naval Hospital Camp Pendleton and salient portions thereof may be entered into my health record. I voluntarily agree to its disclosure to agencies or individuals identified in the preceding paragraph. I have been informed that failure to agree to such disclosure may negate the purposes for which the research study was conducted.

SIGNATURES AND DATE SIGNED:  PRINTED OR TYPED IDENTIFICATION:

Patient / Subject (Date)  Name / Status / Sponsor's SSN
(if Applicable)

Parent / Guardian (Date)  Name / Status
(if Applicable)

Witness (Date)  Name / Grade or Rank
CALIFORNIA EXPERIMENTAL SUBJECTS BILL OF RIGHTS

Any person who is requested to consent to participate as a subject in a research study involving a medical experiment, or who is requested to consent on behalf of another, has the right to:

1. Be informed of the nature and purpose the experiment;

2. Be given an explanation of the procedures to be followed in the medical experiment, and any drug or device to be used;

3. Be given a description of any attendant discomforts and risks reasonably to be expected from the experiment;

4. Be given an explanation of any benefits to the subject reasonably to be expected from the experiment, if applicable;

5. Be given a disclosure of appropriate alternative procedures, drugs, or devices that might be advantageous to the subject with their relative risks and benefits;

6. Be informed of the avenues of medical treatment, if any, available to the subject after the experiment if complications should arise;

7. Be given an opportunity to ask any questions concerning the experiment or the procedures involved;

8. Be instructed that consent to participate in the medical experiment may be withdrawn at any time, and the subject may discontinue participation in the medical experiment without prejudice;

9. Be given a copy of a signed and dated written consent form when one is required;

10. Be given the opportunity to decide to consent or not to consent to a medical experiment without the intervention of any element of force, fraud, deceit, duress, coercion, or undue influence on the subject’s decision; and

11. Be assured that the subject’s confidentiality will be preserved and his/her name will not be released without his/her permission.

Any questions regarding this research study should be directed to the principal investigator or associate investigators. Information is available from the Chairman, Institutional Review Board, established for the protection of volunteers in research projects at this facility by calling (619) 532-8125 or writing the Chairman, Institutional Review Board at Naval Medical Center, Clinical Investigation Department (Code KCA), San Diego, CA 92134-5000.
Smokeless Tobacco Study

This clinic is taking part in a smokeless tobacco research project. If you chew tobacco or use snuff, we would like you to fill out this survey. Filling it out is voluntary and you may choose to skip any question. If you choose not to complete this survey, it will not affect your health care in any way.

For optimum accuracy, please print carefully and avoid contact with the edges of the box. The following will serve as an example:

1. Today's date: [ ]/ [ ]/ [ ]
2. In a typical week, how many days do you use chew/snuff?
   ○ 1 ○ 2 ○ 3 ○ 4 ○ 5 ○ 6 ○ 7
3. How many days does a can/pouch last you?
   ○ 1 ○ 2 ○ 3 ○ 4 ○ 5 ○ 6 ○ 7 or more
4. How soon after you wake up do you use chew/snuff?
   ○ Less than 30 min. ○ 30 to 60 min. ○ More than 60 min.
5. How old were you when you began using chew/snuff?
   [ ] years old
6. Do you swallow tobacco juice on purpose?
   ○ Never ○ Sometimes ○ Almost always
7. Have you tried to quit using chew/snuff in the last year?
   ○ Yes ○ No
8. These statements show how some chew/snuff users feel about quitting. Mark the number that shows how you feel:
   - Not ready to quit
   - Should consider quitting some day
   - Should quit but not quite ready
   - Thinking about cutting down or quitting
   - Have cut down and seriously considering quitting
   - Ready to quit now
   0 1 2 3 4 5 6 7 8 9 10
9. How many of your five best friends use chew or snuff?
   ○ None ○ 1 ○ 2 ○ 3 ○ 4 ○ 5

Please complete the questions on the back of this page. Thank you for your help!
10. Do you currently smoke?  ○ Yes  ○ No

11. On a typical day, how many cigarettes do you smoke?
   ○ None  ○ 1 to 5  ○ 6 to 10  ○ 11 to 15  ○ 16 to 20  ○ 21 to 25  ○ 26 to 30  ○ 31 or more

12. Does your spouse/partner smoke?  ○ Yes  ○ No  ○ Does not apply

13. Have you had two or more years in your life when you felt depressed or sad most days, even if you felt okay sometimes?
   ○ Yes  ○ No

14. In the past year, have you had two weeks or more during which you felt sad, blue, or depressed, or when you lost all interest or pleasure in things that you usually cared about or enjoyed?
   ○ Yes  ○ No

15. In the past seven days, how many drinks of alcohol did you have?  (one drink equals a 12-ounce glass of beer or 6-ounce glass of wine or one shot of liquor)
   ○ None  ○ 1-3  ○ 4-6  ○ 7-9  ○ 10-12  ○ 13-15  ○ 16-18  ○ 19 or more

16. How tall are you?  [ ] feet  [ ] inches  17. How much do you weigh?  [ ] [ ] pounds

18. Your age:  [ ] [ ] years old  19. Sex:  ○ Male  ○ Female

20. Do you consider yourself to be Hispanic or Latino?  ○ Yes  ○ No

21. What race do you consider yourself to be?  Select one of the following:
   ○ American Indian or Alaska Native  ○ Asian  ○ Black or African American  ○ Native Hawaiian or other Pacific Islander  ○ White

22. Marital Status:  ○ Single  ○ Married or living with partner

23. Education finished:
   ○ Less than high school degree  ○ High school graduate or equivalent  ○ Some college  ○ College graduate  ○ Post college

You are finished. Thank you for your help!