1. PROTOCOL TITLE: Heated Ultrasound Gel Satisfaction (HUGS): A Randomized Control Trial

2. ABSTRACT: Ultrasound has become one of the fastest growing diagnostic tools for Emergency Physicians, and given the increasing focus on patient satisfaction in the Emergency Department, there may be opportunities to enhance both patient satisfaction and diagnostic accuracy. The purpose of this single-blind, randomized control trial will be to investigate whether simply having warm gel, as compared to room-temperature gel, during a bedside ultrasound significantly improves patient satisfaction scores. If satisfaction scores are significantly higher, then medical directors may find it beneficial to invest in ultrasound gel warmers. We will also investigate two secondary hypotheses: 1) patient perceptions of their medical providers and 2) the overall quality of ultrasound images obtained.

3. OBJECTIVES/SPECIFIC AIMS/RESEARCH QUESTIONS.
Hypothesis: The utilization of warmed ultrasound gel significantly increases patient satisfaction with regards to their care during a visit to the Emergency Department.
Null: The utilization of warmed ultrasound gel does not change patient satisfaction with regards to their care during a visit to the Emergency Department.

Hypothesis 2: Utilization of warmed ultrasound gel significantly improves the quality of images obtained during bedside ultrasound.
Null: The utilization of warmed ultrasound gel does not change the quality of images obtained during bedside ultrasound.

Hypothesis 3: The utilization of warmed ultrasound gel significantly increases patient perceptions of their physician’s professionalism.
Null: The utilization of warmed ultrasound gel does not change patient perceptions of physician professionalism.

4. MILITARY RELEVANCE. Patient satisfaction as an indicator of the quality of healthcare has evolved as an outcome measure within both the federal and civilian sectors. In fact, the DOD military health system incorporated patient satisfaction surveys to serve as a measure for how its beneficiaries perceive their care. In order to further review specific factors that may help to improve patient satisfaction scores, this study will explore the use of warmed ultrasound gel and perceived patient satisfaction. For most soldiers and dependents, warmed ultrasound gel is generally more comfortable than room temperature gel because warmed gel appears to enhance patient comfort and improve overall experience with the procedure. There is a considerable gap in the literature regarding factors that improve enhanced patient comfort in military beneficiary populations. Therefore, one of the contributions of this study will be to clarify precisely what that level of satisfaction is, if use of warmed gel is truly a satisfaction factor, and if the quality of images improve with the use of warmed gel. Generally, we aim to contribute to the generalizable knowledge of medical science, particularly in regards to patient satisfaction and the impact of warmed gel on ultrasound imaging, which would be applicable to both military and civilian settings.

5. BACKGROUND AND SIGNIFICANCE. Patient satisfaction is a growing center focus of Emergency Departments across the country. One study evaluated over 2.4 million ED visits over the course of four
years in eight different states. Their data showed that patients who responded with the lowest quarter of satisfaction scores were twice as likely to file a complaint compared to the top quarter of patients surveyed. They are 93% of physicians in Pennsylvania from six of the most litigious medical services reported that they practice “defensive medicine and within this group, ED physicians had the highest rates of ordering more tests than medically necessary."

In a 2005 study in JAMA, 93% of physicians in Pennsylvania from six of the most litigious medical services reported that they practice “defensive medicine and within this group, ED physicians had the highest rates of ordering more tests than medically necessary.”

In line with the concept of more tests than medically necessary, CT orders in the ED are becoming standard practice as a method of evaluating a wide variety of complaints. Levin, Parker, and Fragos (2014) reported that for every 1000 Medicare ED visits between 2002-2012, CT scanner use increased remarkably across the country. The doses of radiation that come with CT scans, however, can increase the risk to the patient in the long run. For example, one abdominal CT delivers 8mSv, which is roughly equivalent to the radiation from 400 chest x-rays. To counter this increase in CT scans, ultrasound has become one of the fastest growing and safer diagnostic tools for ED physicians over the past decade. Non-cardiac ultrasound use increased by 121% from 2002-2012, with 21 patients per 1000 being scanned. With real-time diagnostic capabilities at the bedside and no harmful radiation, ultrasounds provide clinicians with decision tools to better care for their patients. Class I medical devices, such as the Thermasonic Gel Warmer, heat ultrasound gel to-and-just-above normal body temperatures are frequently seen in Cardiology and OB-GYN clinics, where echocardiograms and pelvic ultrasounds are performed routinely. Given the increase in use of ultrasound and the increased focus on patient satisfaction, our aim is to identify whether warmed gel can truly improve overall patient satisfaction when performing point-of-care ultrasounds in the ED.

6. RESEARCH DESIGN

This is a single-blinded, randomized study will recruit a sample of 124 subjects from military and civilian populations who are treated in the Emergency Department (ED) at SAMMC and require standard of care bedside ultrasound for diagnosis.

7. RESEARCH PLAN

7.1 Selection of Subjects

7.1.1 Subject Population.

7.1.2 Inclusion Criteria:

- Adults 18-89 who are seen in the SAMMC Emergency Department that require standard of care bedside ultrasound

7.1.3 Exclusion criteria:

- Minors under age 18
- Pregnant females
- Adults over age 90
- Vulnerable adults to include those with altered mental status, incarcerated, military basic trainees, or those with a primary language other than English
- Patients with open or broken skin
7.1.2. Source of Research Material.

<table>
<thead>
<tr>
<th>Source of Research Material</th>
<th>Clinical Purposes (Y/N)</th>
<th>Research Purposes (Y/N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Survey</td>
<td>N</td>
<td>Y</td>
</tr>
<tr>
<td>Provider Survey</td>
<td>N</td>
<td>Y</td>
</tr>
<tr>
<td>Bedside Ultrasound</td>
<td>Y (standard of care)</td>
<td>Y</td>
</tr>
</tbody>
</table>

7.1.3. Description of the Recruitment and Prescreening Process. Patients being treated in the ED who require a standard of care bedside ultrasound for diagnosis will be approached by the AI (ultrasonographer). The AI will advise patient that they are there to perform the ultrasound as ordered by his or her doctor and subsequently ask if the patient would be interested in participating in a patient satisfaction survey surrounding the ultrasound. If the patient says they are interested and meets inclusion/exclusion criteria, the AI will provide the patient with the information sheet (see Appendix C) and explain the study to subject.

7.1.4. Subject Screening Procedures. N/A

7.1.5. Consent Process. In a private exam area in the ED, the AI will inquire whether the patient, who required bedside ultrasound and who meets inclusion/exclusion criteria, would be interested in learning about a research study involving how ultrasounds are performed. The AI will provide an information sheet about the study to interested patients. If the patient is interested, the AI will review the information sheet (Appendix C) and answer any questions the patient may have. We are requesting a waiver of documentation of consent for this project as the study procedures are minimal risk and involves no procedures for which written consent is normally required outside the context of research.

Because we are not using or collecting any personally identifiable information for research purposes, a HIPAA authorization does not apply to this research.

7.1.6. Compensation for participation. None

7.2 Drugs, Dietary Supplements, Biologics, or Devices.

7.2.1 N/A

7.2.2 The Thermasonic Gel Warmer (Model 82-03 LED, 120V) is a Class I medical device that warms up to three (3) gel bottles to settings of “Low” (97 Fahrenheit), “Medium” (102 Fahrenheit), or “High” (109 Fahrenheit) (Appendix E). For this study, we will set all gel warmers to the “Medium” setting. For a full listing of the device instruction manual, please visit [http://www.parkerlabs.com/thermasonic](http://www.parkerlabs.com/thermasonic).

A heat-resistant glove, ULine Terry Cloth Glove (Appendix F), will be used by the AI to handle the ultrasound gels. This glove is manufactured to resist temperatures up to 250 degrees Fahrenheit. In line with SAMMC’s infection control practices and patient safety, ultrasound operators will wear a sterile glove over the terry cloth glove to prevent the spread of microbiota from subject to subject. For more information about the glove, please visit their website at [http://uline.com/BL_111/terry-gloves](http://uline.com/BL_111/terry-gloves).

7.3. Study Procedures/Research Interventions. The principal investigator will brief ED physicians on details
of this study by email (Appendix D). ED physicians who identify that a patient will require an ultrasound will page the ultrasound team as part of standard of care. At that time, the AI (ultrasonographer) will ask subject if he/she would like to participate in a research study involving patient satisfaction and diagnostic bedside ultrasound. If the patient agrees, the AI will provide an information sheet (Appendix C) and explain study details/answer any questions. The patient will then be randomized and assigned a HUGS-X number that will be placed in the comments section of the ultrasound image and given a 3-question patient survey (Appendix A) related to patient satisfaction and perceptions about their provider which will be completed at the end of the ultrasound procedure. The patient will be instructed not to provide any PII/PHI when he/she completes the survey after the diagnostic bedside ultrasound and to hand it back to the AI when complete.

Ultrasound and Warm or Room Temp Gel Procedures:

The AIs will use either the warm or cold gel to complete their standard of care diagnostic ultrasound scans. Subjects will be randomized to receive either warm or cold gel depending on the study packet number. The study packet number will contain specific randomized instructions for the AI that provides the appropriate gel numbered 1-6 to utilize and the 3-question patient survey and 1 question provider survey. Patients and providers will be blinded to the intervention meaning both the AI and the patient will be unaware of different gel temperatures being utilized until the moment it is used on the patient’s ultrasound area.

The provider (AI) will be blinded to the intervention used in two ways. The first being that six ultrasound gel warmers, labeled #1-6, will be set up at a station, and the warmers will have been randomly set-up as far which are actually turned on or not by the PI. The AI will not know which are turned on because the indicator lights will be covered with black tape. Additionally, the AI will use heat-resistant gloves to hold the ultrasound gel bottle as described in section 7.2.2. At the end of the subject’s bedside ultrasound, the physician completing the ultrasound will hand the patient a satisfaction survey utilizing visual analog scales. As an additional measure to test our blinding measure, the AI will complete the one question survey to test whether he or she was aware of the warm or room-temperature gel temperature. Both surveys will be collected and stored in a locked box in C-Pod of the Emergency Department until results can be entered in to the electronic spreadsheet (HUGS DataTracker Data Collection Spreadsheet, Appendix G)

The Ultrasound Image Rating:

Ultrasound image rating is conducted weekly by the SAMMC ED/Ultrasound Quality Assurance Team on a weekly basis. These meetings are already a standard practice in our department to review every ultrasound obtained in the department as a way to obtain feedback from all clinicians about the quality of their ultrasound studies. Their standard metric for evaluation is a 5-point Likert-type scale. During this QA review, scores related to images that have a HUGS ID in the comments area will be written down and provided to the PI. Their standard metric for evaluation is a 5-point Likert-type scale.
Table I: Summary of Study Activity/Interventions

<table>
<thead>
<tr>
<th>Assessment:</th>
<th>During ED Visit:</th>
<th>Weekly Ultrasound QA Session:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screening</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Randomization</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Bedside Ultrasound</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Anonymous Patient Survey Appendix A)</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Provider Survey (Appendix B)</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Review of Ultrasound Images for Overall Quality</td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>

7.3.1 Collection of Human Biological Specimens. N/A

7.3.1.1 Laboratory evaluations and special precautions. N/A

7.3.1.2 Specimen storage. N/A

7.3.2 Data Collection. Subject specific data (patient age, gender, and their satisfaction scores) will all be written on the anonymous survey (Appendix A). The primary outcome will be a visual analogue scale, i.e. continuous variable. By collecting a variable with greater data density than a conventional Likert scale (e.g., ordinal data), this will enable us to use more powerful statistical techniques in analyzing our data so minimizing the possibility of a type II (beta) error. We will further evaluate our secondary outcomes with Likert-type scales to minimize millimeter differences between patient scores. Finally, we will use binary data to evaluate whether patients were overall satisfied with their visit.

Primary Outcome #1a: Overall, how satisfied are you with the experience of having a bedside ultrasound today? Ranges from “Least” to “Most” on 10cm VAS (continuous data)

Primary Outcome #1b: Overall, are you satisfied with the care you received today in the Emergency Department? “Yes” or “No” (binary data)

Secondary Outcome #1: Overall, how professional was the physician who did your bedside ultrasound? Ranges from “Least” (1) to “Most” (5) on 5-point Likert-type scale (ordinal data)

The subject survey form will not contain any PHI/PII. Completed packets will be taken to the locked bin in C-Pod 2 of the Emergency Department. We will also store the completed provider surveys in the same container to allow for further evaluation of provider answers regarding which type of gel they believe they used and if the blinding method imposed by the PI prevented the provider from knowing which gel they utilized.

New study packets will be kept in a separate designated bin within the Emergency Department, next to the ultrasound gel warmers, so that associate investigators may access all required materials while working in the department. The PI or available AI will pick up completed packets weekly and will either log the data immediately or store them in the designated locked bin in the ultrasound team room until they can be entered into the HUGS DataTracker spreadsheet.
Subject bedside ultrasounds, like any clinical ultrasound done in our hospital, are uploaded automatically to the IMPAX system. Each week, the SAMMC ED/Ultrasound Quality Assurance Team reviews all bedside ultrasounds for quality assurance purposes. Their standard metric for evaluation is a 5-point Likert-type scale. This type of data is ordinal (and is the basis for secondary outcome #2). These scores will be logged by the PI or available AI into an Excel Spreadsheet found on a password-protected computer behind a locked door in the ultrasound team room. As all investigators are blinded to which ultrasounds were conducted with warm or cold gel, there is no concern for a bias regarding the rating of quality for each ultrasound or during entering survey data to the HUGS DataTracker. Upon transfer of all data to the spreadsheet, the paper surveys and hand-written scores for each ultrasound will be shredded.

7.4 Statistical Consideration

7.4.1 Sample Size Estimation. This study will be developed to assess our primary outcome with a two-tailed alpha of 0.05 and a beta of 0.2 (power of 0.8), with equal numbers in the warm and cold gel groups. Based on previous studies on visual analog scales and patient satisfaction in the Emergency Department, we will be using an effect size of 0.11 (indicating 11 mm on the VAS) with a standard deviation of 0.21 (indicating 21 mm on the VAS).5

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<table>
<thead>
<tr>
<th>Estimate Required Sample Size</th>
<th>114</th>
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<tbody>
<tr>
<td>Estimate Subject Drop Out / Withdrawal</td>
<td>10</td>
</tr>
<tr>
<td>Total Enrollment Requirement</td>
<td>124</td>
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</tbody>
</table>

<table>
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<tr>
<th>Enrollment at Each Site</th>
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<tr>
<td>BAMC</td>
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7.4.2 Primary (i.e., primary outcome variables) and secondary endpoints. As discussed in 7.4.1, this study has been powered to detect an effect size of 11 mm on the VAS for a statistically significant difference when investigating our primary outcome (1a).

#1a: Overall, how satisfied are you with the experience of having a bedside ultrasound today?
- Ranges from “Least” to “Most” on 10 cm on VAS

#1b: Overall, are you satisfied with the care you received today in the Emergency Department?
- “Yes” or “no”

#2: Overall quality of images
- Ranges from lowest quality (1) to highest quality (5) on a Likert-type scale

#3: Overall, how professional was the physician who did your bedside ultrasound?
- Ranges from “Least” (1) to “Most” (5) on Likert-type scale

7.4.3 Data analysis. The patient satisfaction scores will be compared (warm vs. cold groups) using independent sample t-tests. If the data are ordinal or not normally distributed with equal variances, the equivalent non-parametric test will be used. Independent sample t-tests will also be used to compare the quality ratings of the ultrasound images.

7.5 Confidentiality. The PI or available AI will pick up completed packets weekly and will either log the data immediately or store them in the designated locked bin in the ultrasound team room until they can be entered into the HUGS DataTracker spreadsheet. It will not be possible to associate survey forms with patient charts. Data from the anonymous surveys will be entered into an Excel Spreadsheet (HUGS DataTracker, Appendix G) which will be stored on a password protected computer behind a locked door. The completed paper surveys will be immediately shredded once entered electronically. Once the data has been analyzed, the anonymous data...
file will be deleted.

The bedside ultrasounds, which are part of routine clinical care in our department, are stored within the electronic IMPAX database. All completed ultrasounds for the past week within our emergency department are automatically queued for review by our ultrasonography team. This process is inherent in our system and is not specially designed for this research. They will know that an ultrasound study is part of the research study as the “HUGS” identifier will be seen in the comments box of the ultrasound images. Only medical providers within the military healthcare system have access to these scans.

7.5.1 Certificate of Confidentiality. N/A

8.0 RISKS/BENEFITS ASSESSMENT

8.1 Risks. There are minimal risks for this study. Although this survey is anonymous and cannot be tracked to specific individuals, there is a small risk of inadvertent breach of confidentiality. Participants will not be identified and will be reminded not to add identifiers to their survey documents. For the ultrasound images on IMPAX, only military providers within the military healthcare system have access. Risk of a confidentiality risk is similar to the normal, daily risk of breach in our healthcare system.

There is also the risk of brief discomfort to subjects related to the gel temperature. Medical devices heat the gel to a maximum temperature of 104 degrees. Injuries from gel at this temperature or at room temperature are very unlikely to occur.

Although there is a minimal risk of gel being contaminated with microbiota, we will further mitigate the risk by excluding patients with broken skin in the area of where an ultrasound probe would be placed onto the skin.

8.2 Potential Benefits. There may be direct benefits to participants. Subjects enrolled in the warm gel group may benefit from a more comfortable ultrasound.

9.0 ADVERSE EVENTS, UNANTICIPATED PROBLEMS, AND DEVIATIONS

9.1 An adverse event in this study would be either a burn to the patient, infection, or a breach of confidentiality. These are considered rare but serious, as described above. Adverse events during the study would be documented by the ultrasonographer or another member of the research team and reported by the PI to the IRB as required.

9.2 Reporting Unanticipated Problems Involving Risks to Subjects or Others, Serious Adverse Events and Deaths to the Office of the IRB, BAMC.

All unanticipated problems involving risk to subjects or others, serious adverse events, and all subject deaths related to the study will be reported within 48 hours of the research team’s knowledge of the event by phone (210-916-0606), by e-mail (BAMC_IRB_AE@amedd.army.mil), by facsimile (210-916-1650) or via letter addressed to Human Protections Administrator, Office of the Institutional Review Board, Brooke Army Medical Center, Attn: MCHE-CI, 3698 Chambers Pass, Fort Sam Houston, TX 78234-6315. A complete written report will follow the initial notification within 10 working days.

9.3 Research Monitor. N/A

10.0 WITHDRAWAL FROM STUDY PARTICIPATION. Subjects may withdraw from the study at any time prior to turning in his/her survey. Their survey packet, if they began to write on it, would be shredded if they withdraw from the study prior to turning it in. Regardless of participating in the study, the patient would still...
receive the bedside ultrasound as it is part of their clinical care. Because this is an anonymous survey, once
the survey is turned in, data will be used and transferred to the Excel Spreadsheet (HUGS DATA
TRACKER). There would be no adverse effects for the patient if he or she decided to withdraw from the
study.

11.0 USAMRMC Volunteer Registry Database. N/A

12.0 REFERENCES.

1. Cydulka RK, Tamayo-Sarver J, Gage A, and Bagnoli D. Association of patient satisfaction with complaints and risk
3. Levin DC, Rao VM, Parker L, and Frangos AJ. Continued growth in emergency department imaging is bucking the
4. Mettler FA Jr, Huda W, Yoshizumi TT, and Mahesh M. Effective doses in radiology and diagnostic nuclear medicine:
5. Singer AJ, and Thode HC. Determination of the minimal clinically significant difference on a patient visual analog
and Assessment of Department of Defense Effects

13.0 TIME REQUIRED TO COMPLETE THE RESEARCH (including data analysis). Up to 18 months: 12
months for recruitment and up to 6 months for analysis and manuscript preparation.

14.0 STUDY CLOSURE PROCEDURES: All paper surveys will be shredded and any computer data with
subject numbers will be deleted upon entry into the data collection spreadsheet. After study completion, a final
closure report will be submitted to the IRB. Ultimately, a manuscript will be generated with the intent of
submitting to a peer-reviewed medical journal for publication.

Appendices:

Appendix A: Anonymous Subject Survey
Appendix B: Provider Survey
Appendix C: Subject Information Sheet
Appendix D: Email to Emergency Medicine Staff
Appendix E: Thermasonic Product Information Sheet
Appendix F: ULine Terry Cloth Information Webpage
Appendix G: HUGS DataTracker Data Collection Spreadsheet