STUDY PROTOCOL

An observational study protocol to explore loneliness and systemic inflammation in an older adult population with chronic venous leg ulcers

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Abstract

Background: Chronic venous leg ulcers (CVLUs) are the most common type of lower extremity wound. Even when treated with evidenced-based care, 30–50% of CVLUs fail to heal. A specific gap exists about the association between psychosocial stressors, particularly loneliness, and biomarkers of inflammation and immunity. Loneliness is highly prevalent in persons with CVLUs, has damaging effects on health, and contributes to the development of multiple chronic conditions, promotes aberrant inflammation, and diminishes healing. However, the confluence of loneliness, inflammation and the wound healing trajectory has not been elucidated; specifically whether loneliness substantially mediates systemic inflammation and alters healing over time. This study seeks to address whether there is a specific biomarker profile associated with loneliness, CVLUs, and wound healing that is different from non-lonely persons with CVLUs.

Methods: An observational prospective study will identify, characterize and explore associations among psychosocial stressors, symptoms and biomarkers between 2 CVLU groups, with loneliness+ (n = 28) and without loneliness- (n = 28) during 4 weeks of wound treatment, measured at 3 time points. We will examine psychosocial stressors and symptoms using psychometrically-sound measures include PROMIS® and other questionnaires for loneliness, social isolation, depression, anxiety, stigma, sleep, fatigue, pain, quality of life, cognition, and function. Demographics data including health history, sex, age, wound type and size, wound age, and treatment will be recorded from the electronic health record. We will characterize a biomarker panel of inflammatory genes including chemotaxic and growth factors, vascular damage, and immune regulators that express in response to loneliness to loneliness and CVLUs using well-established RNA sequence and PCR methods for whole blood samples. In an exploratory aim we will explore whether age and sex/psychological stressors and symptoms indicate potential moderation/mediation of the effect of loneliness on the biomarker profile over the study period.

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Background

It is well established that individuals with chronic wounds, particularly those with chronic venous leg ulcers (CVLUs), have substantially reduced quality of life [1–3]. A wound is defined as chronic when it fails to reduce in size by 50% for at least 6 weeks after onset [4]. Up to 150 million individuals worldwide live with slow or non-healing ulcers. This is problematic because CVLUs account for ~80% of all chronic ulcers [5] that take months to years to heal [6]; for those that do heal, as many as 70% recur within 3 months [7].

Most CVLUs are associated with physical, psychological, and social symptoms. Physically, people with CVLUs experience symptoms such as odor, drainage, and leg swelling, which affect functional ability and physical appearance [8]. Self-management of CVLUs has been linked to fatigue and pain [9], and physical exhaustion [10] from the intense care required. Psychological symptoms reported by people with CVLUs include fear that the ulcer will never heal, will get infected, or that an amputation may occur [1, 10]. In the literature, cognitive dysfunction, poor self-image and worry [1], depression and anxiety have been reported [9]. These physical and psychological symptoms are known to negatively influence mobility, sleep, and mood, and limit professional, familial and social relationships, especially contact with friends [11]. Frequent and prolonged care is reported to lead to feelings of anger and resentment [12].

It is possible that sex, psychosocial stressors, and symptoms play a predominant role in perpetuating a chronic, non-healing state yet these factors are under-studied, rarely assessed, and not consistently managed during clinical encounters. It is concerning that 30–50% of CVLUs fail to heal, even when treated based on best clinical practices which include: compression and leg elevation to improve venous blood flow, wound dressings to control necrotic tissue and wound exudate, and advanced topical biological therapies such as skin substitutes and growth factors to alter the local wound environment [13]. In the past 10 years, two studies have discovered that males with CVLUs report higher scores for both depression and anxiety [14, 15]. The findings regarding sex and associated symptoms underscore the critical need to explore the psychosocial and biological mechanisms that influence wound healing [16].

A specific gap exists when seeking knowledge about the association between psychosocial stressors, particularly loneliness, and biomarkers of inflammation and immunity. Loneliness has damaging effects on health, and contributes to the development of multiple chronic conditions [17], and diminished healing. Population-based longitudinal research indicates that loneliness predicts functional decline, morbidity and mortality, independent of objective social isolation, depression and health behavior [18, 19]. Loneliness is defined as a distressing feeling that accompanies the perception that one’s social needs are not being met by the quantity (social isolation) or especially the quality of one’s social relationships with individuals and/or the community [17, 20–23]. Loneliness affects ~50% of older adults in the U.S. [24], 68% of older adults with chronic wounds [25], and is associated with cardiovascular disease, hypertension, stroke, cancer, functional disability, and cognitive decline [26–28]. The negative health effects of loneliness exceed those of cigarette smoking, physical inactivity, and obesity [17, 29–31].

A growing body of evidence indicates wound healing is substantially altered by loneliness and social isolation in both human and animal models [32–34]. Loneliness is associated with altered molecular mechanisms such as inflammation and increased levels of pro-inflammatory cytokines [35], that may impede wound healing [36–38]. In addition, exposure to psychosocial adversity has been shown to elicit vascular, neuroendocrine physiological responses including increased peripheral resistance and dysregulation of the hypothalamic-pituitary-adrenocortical (HPA) axis [38].

A growing literature in human social genomics suggests that psychosocial adversity and a negative social environment are associated with differential expression of hundreds of gene transcripts involved with inflammation and antiviral response, known as the conserved transcriptional response to adversity (CTRA) pathway. Loneliness has been associated with up-regulated inflammation by enhancing the transcription of genes such as interleukin-1 alpha and beta, interleukin-6, interleukin-8, cyclooxygenase 2, and tumor necrosis factor [39, 40] and the potent pro-inflammatory NF-kappa B gene [41]. The CTRA pathway is characterized by decreased expression of Type I interferon-related antiviral genes and increased...
expression of proinflammatory genes. However there is a paucity of literature to determine whether these same genes are also expressed in individuals with loneliness and chronic wounds [42]. The premise of this study is that a ~50-gene composite score previously used to assess the CTRA profile will be up-regulated in lonely (L+) individuals with CVLUs supporting the link between chronic wounds and inflammatory genes [43, 44]. The overexpression of these wound-related genes, in particular inflammatory cytokines may be considered candidates for the prediction of treatment response among patients with L+ and CVLUs. Experimental studies in animal and human models show a causal impact of adverse social conditions on CTRA gene expression [45]. Specific to chronic wounds, systemic inflammation is a well-established biological pathway linked to poor wound healing outcomes and is the focus of our study.

Recognizing that inflammatory biomarkers such as elevated cytokines and altered gene expression exist in CVLUs, we hypothesize that substantially heightened inflammation is a common molecular mechanism with a distinct profile that underlies both loneliness and poor wound healing in a chronic wound population compared to a wound population without loneliness. However the confluence of loneliness, inflammation and the wound healing trajectory has not been elucidated; specifically whether loneliness substantially mediates systemic inflammation and alters healing over time. The major question this study seeks to address towards advancing wound and symptoms sciences is whether there is a specific biomarker profile associated with loneliness, CVLUs, and wound healing that is different from non-lonely patients with CVLUs by addressing the following aims:

**Aim 1**
Examine whether psychosocial stressors (i.e., social isolation, social support) and symptoms (i.e., fatigue, pain, depression, anxiety, sleep disturbance, reduced QOL) differ between (L+) and (L-) patients with CVLUs using well-validated questionnaires.

**Aim 2**
Characterize a biomarker (chemotaxic factors, growth factors, vascular damage and immune regulators) profile common to loneliness and CLVU using well-established RNA sequencing and PCR methods for whole blood samples. We will start the analysis using a 1.8-fold change in gene expression, either positive or negative, to generate a composition score to assess the effect of loneliness and stressors on biomarkers.

**Exploratory aim 3**
Explore whether age and sex/psychological stressors and symptoms indicate potential moderation/mediation of the effect of loneliness on the biomarker profile over the study period.

**Methods/design**

**Overall strategy**
The overarching aims of this observational prospective study are to identify, characterize and explore associations among psychosocial stressors, symptoms and biomarkers (panel of inflammatory genes that express in response to loneliness) between 2 CVLU groups, L+ (n=28) and L- (n=28) during wound treatment at 3 time points – baseline (V1), end of month 1 (V2) and end of month 2 (V3) - over a 2-month period. Eligibility criteria include: female and male; ≥ 60 years of age; CVLU of ≥6 weeks duration, and English speaking. Patients will be excluded if they are undergoing chemotherapy (affects immune factors), or taking steroids (may decrease inflammation and affect cytokine levels). We do not want to exclude participants based on wound treatment because the point of the proposal is to better understand loneliness in people with wounds. We are collecting treatment data but it will not determine inclusion. We also want to characterize participants by loneliness based on treatment but in order to do that, we will need to do a subsequent adequately powered study to determine the influence of treatments. Demographic and health data will be obtained from participants and the electronic health record. All participants will be recruited from the local wound communities/wound clinics in the southeastern region of the U.S.

Participant compensation will be provided. In return for their time and effort, each participant will receive a check for $25.00 for each completed study visit. Checks will be mailed to the address that the participant provides to the researchers. Additionally, we will offer a “fish bowl” prize drawing at each visit in which patients can “win” an additional $1 to up to $25 at V2 and V3 – we have used this contingency management strategy in our previous wound prevention studies with great success (7% attrition rate in a recently completed randomized controlled trial). The odds for the prize drawing are devised so that participants will not in all likelihood receive disproportionate amount of study compensation. A CONSORT diagram of participant flow through the study is provided below in Fig. 1.

**Demographics**
Age, race, ethnicity, sex, marital status, education, disability status, income (socioeconomic status), and employment information will be collected by study personnel and recorded on the investigator-generated study specific demographics form developed for use in our previous studies. We will enter all data for the 3 visits into our study database in REDCap, a secure electronic...
documentation data storage management system. There are known sex differences in the neuroendocrine stress response and in the prevalence, symptoms, and correlates of loneliness in adults [46]. In our prior studies we reported that, in older adults, female sex was predictive of loneliness yet in rural areas such as Appalachia, mid-life men had higher mean loneliness scores [47]. Therefore, it is likely that loneliness impacts the health of males and females differently, making it critical to include sex as a biological variable in the study design. We will attempt to recruit equal numbers of males and females per loneliness group (L+ vs L-). Though no sex-specific hypotheses will be tested in the proposed study, we plan to explore possible differences of the effect of loneliness by sex (moderation) in the analyses, for example, through inclusion of sex-by-interaction terms. As this study is not powered to confirm hypotheses regarding moderating effects, these analyses will be considered hypothesis generating and descriptive rather than hypothesis testing. Per NIH guidelines, we will also report outcomes separately by sex.

**Instruments**

The SPIRIT diagram (Table 1) below summarizes the schedule of enrollment, assessments, and visits across the study.

**Aim 1. Stressors and symptoms measures**

In our previous trials, we tested the feasibility of the battery of instruments which took approximately 45 min to complete and found this was acceptable to our patients. However we are mindful of response burden and the time commitment for completing questionnaires and having blood specimens obtained. At the 3 data collection periods during regular wound clinic visits, participants can opt to answer questionnaires via assistance from study personnel in paper form or electronically on a tablet. Participants in both groups will receive the same battery of instruments. Psychometrically-sound measures include PROMIS® and other measures of loneliness, social isolation, depression, anxiety, stigma, cognition, and function. Data on wound type and size, wound age, and treatment will be recorded from the electronic health record. In the event that a participant expresses psychological distress when responding to psychological questionnaires (e.g., depression scale), our study wound physician will be contacted. We developed a plan for referral of depressed or suicidal participants.

**Aim 2: blood draws for genomics analyses**

Patient blood samples will be drawn by study personnel via venipuncture directly into PAXgene RNA tubes which inhibit RNA degradation and allow for purification of
RNA for subsequent sequencing analyses. All blood samples will be drawn during the clinic visit, placed in the transport bag, immediately delivered directly to the MUSC laboratory at room temperature, and slowly frozen to −20 °C prior to processing, shipping, and analysis. RNA from patient samples will be sequenced in a blinded fashion in batches. We will attempt to schedule patients for clinic visits between 9 and 11 am to reduce the influence of diurnal variations.

### Feasibility and process monitoring

We will observe 10% of data collection procedures including questionnaires and specimen collection at least monthly. The team will track other study procedures through review of consenting logs, noting any problems encountered during interviewing and specimen analysis, participant burden of completing questionnaires/time taken, and document other barriers such as transportation that arise during data collection.

### Data analyses

Univariate descriptive statistics and frequency distributions will be calculated as appropriate for all variables. Inflammatory profiles, loneliness, psychosocial variables, and physical and functional variables will be obtained at baseline (V1), month 1 (V2) and month 2 (V3). Preliminary analyses will explore patterns of missing data for all outcome variables. We will also explore differences in demographic and other variables such as age (60–74, ≥75 years), sex, chronic illnesses, cognition, health status, functional activity, nutritional status, length of time to heal (healing trajectory), and wound treatment type across the 3 time points. All statistical analyses will be conducted using SAS Statistical Software Version 9.4 (Copyright © 2016 by SAS Institute Inc., Cary, NC, USA).

### Analyses for aim 1

The primary variables for aim 1 are social isolation and social support (psychosocial stressors) and fatigue, pain,

**Table 1** Study SPIRIT diagram

| TIMEPOINT | STUDY PERIOD |
|-----------|--------------|
|           | ENROLLMENT   | VISIT SCHEDULE |
|           | Day 0        | Week 4         | Week 8         |
| ENROLLMENT| Eligibility screening checklist | X         | X         | X |
|           | Informed consent | X         | X         | X |
|           | Allocation (L+/L-) | X         | X         | X |
| ASSESSMENTS AND MEASURES | Demographics and characteristics | X         | X         | X |
|           | Wound assessment and medical health history | X         | X         | X |
| VEINES-QOL/Sym - 26 item | PROMIS Cognitive Function 6a | X         | X         | X |
|           | PROMIS Global Health Scale - 10 item | X         | X         | X |
|           | PROMIS Anxiety 6a | X         | X         | X |
|           | PROMIS Fatigue 6a | X         | X         | X |
|           | PROMIS Pain Interference 6b | X         | X         | X |
|           | PROMIS Sleep Disturbance 6a | X         | X         | X |
|           | Katz Activities of Daily Living - 6 item | X         | X         | X |
|           | Mini nutritional status - 7 item | X         | X         | X |
|           | UCLA Loneliness Scale - 20 item | X         | X         | X |
|           | Medical Outcomes Social Support Survey - 19 item | X         | X         | X |
|           | Patient Health Questionnaire - 9 item | X         | X         | X |
|           | The Stigma Scale: Body Image and the Skin - 8 item | X         | X         | X |
|           | Blood Sample: RNA for deep sequencing analysis* | X         | X         | X |

*Chemotactic factors: CCL2/JE/MCP-, CCL3/MIP-1 alpha, CCL4/MIP-1 beta, Granzyme B, CCL5/RANTES, CCL11/Eotaxin, CCL19/MIP-3 Beta, CCL20/MIP-3 alpha, CX3CL1/Fractalkine, PD-1/L1/B7-H1, CXCL1/GRO alpha/IL-12/PLA, CXCL2/GRO, and CXCL10

*Immune regulators: IL-3, IL-4, IL-5, IL-6, IL-7, IL-8/CXCL8, IL-10, IL-12 p70, IFN-alpha IL-13, IFN-beta IL-15, IFN-gamma IL-17/IL-17A, IL-1 alpha/IL-1F1, IL-1B/IL-25, CD40 Ligand/TNFSF5, IL-1 beta/IL-1F2, IL-13, IL-1ra/IL-1F3, and IL-2

*Vascular damage: VCAM, ICAM, CCRP, MMP-1, MMP-3, and MMP-

*Growth factors: PDGF-AB/BB, TGF-alpha, EGF, TNF-alpha, FGF basic/FGF2 Flt-3 Ligand TRAIL, G-CSF, GM-CSF, VEGF, and PDGF-AA
depression, anxiety, sleep disturbance, reduced QOL (symptoms). To examine differences in and variability of psychosocial stressors and symptoms over time between the L+ and L- groups, we will compare the longitudinal profile of the psychosocial stressors and symptoms (in individual models) between the groups over the 3-month study period using mixed effects models (MEM) analyses. These analyses will estimate the average change in the dependent variable within each group (L+ vs. L-) and individual change in variables for participants. MEM analyses allow for missing data, measurement of study subjects at different time points during the study, and time varying covariates. MEM can also take into account the effect of clustering, e.g., correlation of repeated measurements within one subject. The psychosocial stressor or symptom will be used as the dependent variable with group (L+ vs. L-), time, and time-by-group interaction as primary independent variables. The GLMM analysis approach allows for time-varying variables; we will also report separately whether and how many patients switch from L+ to L- or vice versa. Nutritional status, functional activity, and wound treatment will be investigated during secondary analyses through inclusion as covariates in the longitudinal models (individually). Further, frequency distributions of adverse events and serious adverse events will be obtained. Proportions within categories of adverse events and serious adverse events for the L+/L- groups will be compared via chi-square analyses.

Analyses for aim 2

All RNA sequencing will be performed at the UCLA Social Genomics Core Lab. RNA will be extracted from PAXgene tubes (Qiagen RNeasy), tested for suitable mass and integrity, and converted to cDNA libraries using a high-efficiency enzyme system (Lexogen QantSeq 3’ FWD). Normalized cDNA libraries will be sequenced using an Illumin HiSeq 4000 instrument, targeting > 10 million reads per sample. Reads will be aligned to the reference human transcriptome and quantified as gene transcripts per million total mapped reads using the STAR aligner. Transcript abundance data will be log2 transformed for analysis by linear statistical models quantifying differential gene expression as a function of key predictors (e.g., loneliness wound healing) while controlling for potential confounders, particularly age, sex, race/ethnicity, BMI, smoking history, and heavy alcohol consumption history. The CTRA will be measured by a pre-specified 53-gene composite score used in previous social genomics studies, [48, 49] including 19 gene transcripts involved in inflammation (i.e., IL1B, IL6, IL8, TNF), 34 gene transcript involved in innate antiviral responses (IFNB, IFI+, MX+, and OAS-family genes). Scores will be composed by z-score standardizing expression of each gene and summing the resulting values after sign-reversing the antiviral genes to reflect their inverse contribution to the CTRA profile.

Analyses for aim 3

The goal of aim 3 is to explore whether age and sex/psychological stressors and symptoms indicate potential moderation/mediation of the effect of loneliness on the biomarker profile over the study period. The composite score of the biomarker expression values will be used as the dependent variable for this analysis. Subsequently, to compensate for the limited sample size of this exploratory analysis, a bootstrapping procedure that resamples the data with replacement will be employed to provide more accurate estimates of variability (95% CIs). The analysis will involve a comparison of the composite score between L+ and L- using MEM as described above. Moderators will be explored through inclusion of an interaction term in the model, for example, age-by-loneliness. Potential mediators will be explored individually using MEM. Exploratory analyses will follow the modified procedure suggested by MacKinnon, Fairchild & Fritz [48] of the causal steps approach developed by Baron & Kenny [50].

Wound healing trajectory definitions

A normal healing trajectory is considered a reduction in wound surface area of 10–15% per week, or at least 30% over a 2-week period. Wounds that fail to decrease in size by 30% over the first 2 weeks of treatment have a 68% probability of failing to heal within 24 weeks [51]. Ulcer area reduction at 2 weeks predicts failure to heal by 24 weeks in the venous leg ulcers of patients living alone. Overall time to closure depends on wound size and characteristics, patient age, nutritional status, comorbid conditions, adherence to treatment, and additional factors to be explored in this study. While there is no wound severity score for venous leg ulcers (unfortunately there is no staging or classification score for these types of wounds similar to staging criteria for pressure injuries), consensus on severity among wound care experts is emerging and is based on wound clinical characteristics as follows:

Grade 3 = size > 10 cm², infection/biofilm/osteomyelitis, ulcer > 4 weeks without change in size, high exudate, pain > 7 numeric rating scale (NRS), necrotic debris/fibrin covering > 50% of wound bed, odor, periwound inflammation/maceration, large lower extremity edema based on calf circumference - would require Level 1 treatment approaches (see below);
Grade 2 = size 5–10 cm², critically colonized, moderate exudate, pain 4–6 NRS, necrotic debris/fibrin 25–50%, moderate lower extremity edema - would require Level 2 treatment; and,
Grade 1 = size < 5 cm², light exudate, pain < 4 NRS, necrotic debris/fibrin < 25% or > 75% pink granulation tissue, mild lower extremity edema – would require Level 3 treatment.

Healing is determined by consistent reduction in size and symptoms and presence of epidermis covering 100% of wound bed. There are treatment differences depending on wound characteristics such as drainage, necrotic debris, leg edema, and infection. We will account for changes in grade and treatment by treatment types (levels) in our analyses as follows:

Level 1: inelastic, multi-component, high compression [> 40 mmHg], mechanical/sharps debridement, antibiotics, advanced healing technologies such as skin substitutes, hyperbaric oxygenation;
Level 2: moderate compression [30–40 mmHg], exudate management, enzymatic debridement; and,
Level 3: moderate compression, moist healing environment.

Sample size considerations
With 28 participants per group we will have 80% power to detect an effect size of 0.6 standardized units effect size between the 2 groups, assuming 2 post-baseline time points; intra-class correlation no greater than 0.4; level of significance \[\alpha\] = 0.05, two-tailed. Assuming standard deviation (sd) = 2, this translates into a raw effect size (difference in change scores) of 1.2. If the pooled sd is 5 units, the standardized effect size is equivalent to a raw detectable difference in change scores between the groups of 4.5 raw scale units. We will enroll 10 males and 10 females prior in each group so that we can complete thorough data analysis for outcome comparisons by sex. If necessary we will over enroll to achieve 28 participant per group at V3. The distribution of patients at one of our main wound centers is 60.0% male, 40.0% female; 60.0% African American, and 40.0% White with mean age 70 years.

Data and safety monitoring
Considering the study design and specific procedures to be performed, the overall risk level for enrolled participants is considered to be minimal. Accordingly, this study will employ the use of an Independent Safety Monitor (Board certified D.O.) who will be provided with real-time electronic access to participant data. This physician will be responsible for reviewing and grading adverse events, monitoring the study safety profile, and making recommendations to the Principal Investigator regarding study modification, termination, and continuance.

Discussion
There is a critical need to understand the trajectory of wound healing and interrelated psychosocial factors in lonely persons with CVLUs and to differentiate molecular mechanisms to advance future treatment approaches aimed to improve chronic wound healing and quality of life. We have demonstrated considerable expertise on the phenomenon of loneliness and its relation to health for chronically ill adults in the U.S. in analyses of Health and Retirement Study (HRS) data that identified predictors and outcomes of loneliness [52, 53] incorporated into the psychoneuroimmunological (PNI) conceptual model (Fig. 2).

Descriptive studies of loneliness in adults in Appalachia with chronic illness have reported loneliness correlated with depression \(r = .388, p < .05\), lower quality of life scores \(r = .272, p < .05\), and diminished social support \(r = .374, p < .05\), particularly diminished emotional support \(r = .459, p < .01\) [54]. Studies of stroke survivors discovered that loneliness was correlated \(p < .05\) with poorer quality of life on 13 domains from the Neuro-Qol [55, 56] and in systematic reviews, relationships were explicated between loneliness and multiple chronic conditions, providing evidence that loneliness may have higher prevalence in rural-residing individuals and that interventions are needed for loneliness in this population [17]. Further, recent qualitative work has discovered that the emotions of worry, anger, and fear are present for lonely older women and that lonely stroke survivors have unmet connection needs [47, 57].

A funded study led to the development of LISTEN, a novel intervention for loneliness that has been deemed feasible, acceptable, and initially effective for diminishing loneliness, and concurrently, diminishing measures of inflammation as persons became less lonely [58]. Finally, in a study of gene expression in lonely adults in a pilot randomized controlled trial of LISTEN, data suggest gene expression for inflammation and immunity changed in a favorable hypothesized direction [59]. The team spent extensive time identifying which variables to include, guided by the PNI model so that we could logically determine a gene expression profile in people with wounds, building on the current evidence base. Note also that previous research on gene expression in the HRS sample has also verified the association of loneliness with increased CTRA gene expression in older adults [60].

Previous research in social genomics has linked adverse social conditions to increased expression of pro-inflammatory genes in correlational human studies, in experimental animal and human models showing causal effects, and in cellular model systems dissecting the specific molecular signaling pathways involved [39, 43]. Loneliness in particular has been repeatedly linked to
up-regulated expression of pro-inflammatory genes (i.e., *IL1B*, *IL6*, *IL8*, *TNF*), and down-regulated expression of genes involved in innate antiviral responses (*IFNB*, *IFIs*, *MX*, *OAS*) [18, 60–66]. This recurring pattern appears to reflect the activation of CTRA in myeloid-lineage immune cells [39, 67]. Our in-depth review of the literature guided the selection of these genes and is based on the best available data. We believe the novelty is in the application of the gene analysis to a population that suffers with wounds. However, the impact of this biomarker profile on chronic wound healing remains unstudied.

We have identified several challenges to our study including relevance of selected biomarkers, models to assess healing, sample size, generalizability of results, and racial/ethnic composition. Increased expression of inflammatory markers is associated with a variety of factors such as older age, multiple chronic conditions (obesity, diabetes), pain, smoking, lower socioeconomic status, and poor functional status [68–70]. We recognize the majority of individuals with CVLUs is generally older (≥60 years of age), has high functional impairment, poor quality of life, and multiple chronic conditions. However it is not known whether there is a distinct profile in patients with chronic wounds and in particular those with loneliness. We will account for these co-variates in our analyses.

Currently there is no assessment mechanism available to predict healing that includes an inflammatory biomarker or psychosocial variables such as loneliness and social isolation. If relationships are indicated among our study variables (and confirmed subsequently), these factors could be added to currently available diseasespecific risk factor models to predict non-healing in wound populations. The M.A.I.D. score, predominantly used in wound care, was created out of 4 clinically-defined parameters to estimate long-term clinical outcomes [71]. However, there are no psychosocial, physical or biological components in this score, thus it lacks full scope risk factors for accurate prediction.

We recognize the small sample size may not allow for detecting statistically significant differences; however, as the focus of the exploratory aim is to generate hypotheses rather than to confirm, we will be able to establish preliminary relationship profiles among the psychosocial stressors and symptom variables. In addition, we will investigate whether there is an indication for clinically relevant differences that support further focused exploration in future studies.

We acknowledge generalizability of the anticipated results is limited in this small exploratory study and anticipate, in a future trial, we will have an adequately powered study and a greater diversity (Latinx) of participants. To this end, we will develop study materials, for example, in Spanish to reflect the diversity of the population.

The projected racial and ethnic composition of the participant sample is a concern given the established differences in gene expression among and between these

![Conceptual model of Key Study Variables Mapped to Major Concepts of the Psychoneuroimmunological (PNI) Paradigm](image-url)
groups; it is unlikely that the least frequent groups (e.g., Hispanic) can be properly evaluated in the proposed analyses, impacting on rigor. We agree and acknowledge the concern about the differences in gene expression among various ethnic/racial groups; for this small exploratory trial, we are focused on fidelity to our PNI model. Though it is not the primary aim of this observational study to determine ethnic differences, we have included age, sex, and race and ethnicity as individual person factors based soundly in the PNI model and these variables will be included and accounted for in preliminary analyses.

Conclusion

Our study addresses a highly prevalent clinical problem - chronic venous leg ulcers (CVLUs) that affect millions of individuals worldwide, causing considerable suffering, disability and poor quality of life. The objective of this exploratory project is to assess stressors, symptoms and biomarkers associated with lonely and non-lonely individuals with CVLUs. Findings are expected to improve understanding of molecular mechanisms common to loneliness and inflammation towards development of a biopsychosocial prognostic indicator of healing potential in persons with chronic wounds. If differences in the biomarker profile of L+ individuals and CVLUs are found, further exploration of co-variables could form the bases of a new type of assessment that could potentially predict nonhealing.

Abbreviations

CVLUs: Chronic venous leg ulcers; CTRA: Conserved transcriptional response to adversity; HPA: Hypothalamic-pituitary-adrenocortical; HRS: Health and Retirement Study; L-: Without loneliness; L+ : With loneliness; LST EN: Loneliness Intervention using Story Theory to Enhance Nursing sensitive outcomes; M:A.ID.: Palpable pedal pulses (II); Wound area (A); Ulcer duration (D), and Presence of multiple ulcerations (M); M:EM: Mixed effects models; NRS: Numeric rating scale; PNI: Psychoneuroimmunological; sd: Standard deviation

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Authors’ contributions

TK, LT, SC, and RMH developed the protocol. MMueller, RMH and SC will analyze and interpret the patient data regarding the study variables including biological data. MMadisetti and MP established the protocols for data collection and management activities. All authors read and approved the final manuscript.

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Availability of data and materials

The datasets used and/or analyzed during the current study will be housed and stored at the Medical University of South Carolina in accordance with the academic institution’s Records Retention policy. Data will be made available from the corresponding author on reasonable request after publication of the results on the main research questions.

Ethics approval and consent to participate

This protocol underwent scientific peer review at the National Institute of Nursing Research. The study was approved by the Medical University of South Carolina Institutional Review Board (IRB #00102546), conforms to the Declaration of Helsinki, and will be conducted in compliance with Good Clinical Practices (GCP) of the International Conference of Harmonization (ICH). Written informed consent will be gathered from all study participants.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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References

1. Kouris A, Armyra K, Christodoulou C, et al. Quality of life psychosocial characteristics in Greek patients with leg ulcers: a case control study. Int Wound J. 2016;13(6):744–7.
2. Kapp S, Miller C, Santamaria N. The quality of life of people who have chronic wounds and who self-treat. J Clin Nurs. 2018;27(1–2):182–92.
3. Santos V, Oliveira ADS, Amaral A, Nishi ET, Junquaia JB. Quality of life in patients with chronic wounds: magnitude of changes and predictive factors. Rev Esc Enferm USP. 2017;51:e03250.
4. Zenliman J, Valle MF, Malas MB, et al. Chronic Venous Ulcers: A Comparative Effectiveness Review of Treatment Modalities. Rockville (MD); Agency for Healthcare Research and Quality (US); 2013 Dec. Report No.: 13(14)-EHC121-EF.
5. Collins L, Seraj S. Diagnosis and treatment of venous ulcers. Am Fam Physician. 2010;81(9):989–96.
6. Abbade LP, Lastoria S. Venous ulcer: epidemiology, physiopathology, diagnosis and treatment. Int J Dermatol. 2005;44(6):449–56.
7. Langer V. Preventing leg ulcer recurrence. Indian Dermatol Online J. 2014; 5(4):534–5.
8. Phillips P, Lumley E, Duncan R, et al. A systematic review of qualitative research into people’s experiences of living with venous leg ulcers. J Adv Nurs. 2018;74(3):550–63.
9. Stechmiller JK, Lyon D, Schultz G, et al. Biobehavioral mechanisms associated with nonhealing wounds and Psychoneurologic symptoms (pain, cognitive dysfunction, fatigue, depression, and anxiety) in older individuals with chronic venous leg ulcers. Biol Res Nurs. 2019;21(4):407–19.
10. Szewczyk MT, Moscicka P, Jawien A, et al. Quality of life in patients with leg ulcers or skin lesions - a pilot study. Postepy Dermatol Alergol. 2015;32(6):465–9.
11. Farah E, Blanes L, Hochman B, Mesquita Filho M, Ferreira L. Health-related quality of life, self-esteem, and functional status of patients with leg ulcers. Wounds. 2011;23(1):4–10.
12. Platsidaki E, Kouris A, Christodoulou C. Psychosocial aspects in patients with chronic leg ulcers. Wounds. 2017;29(10):306–10.
13. BroszczaK DA, Sydes ER, Wallace D, Parker TJ. Molecular aspects of wound healing and the rise of venous leg ulceration: Omics approaches to enhance knowledge and aid diagnostic discovery. Clin Biochem Rev. 2017; 38(1):35–55.
14. da Silva MH, de Jesus MC, Merighi MA, de Oliveira DM, Biscotto PR, Silva GP. The daily life of men who lives with chronic venous ulcer: phenomenological study. Rev Gaucha Enferm. 2013;34(3):95–101.
15. Kelechi TJ, Mueller M, Dooley M. Sex differences in symptom severity and clusters in patients with stage C4 and stage C5 chronic venous disease. Eur J Cardiovasc Nurs. 2017;16(1):28–36.

16. Aguilar AC, Sadigursky D, Martins LA, Menezes TM, Santos AL, Reis LA. Social repercussions experienced by elderly with venous ulcer. Rev Gaucha Enferm. 2016;37(3):353-302.

17. Pettite T, Mallow J, Barnes E, Petrone A, Barr T, Theeke L. A systematic review of loneliness and common chronic physical conditions in adults. Open Psychol J. 2015;8(Suppl 2):113–32.

18. Cacioppo JT, Cacioppo SP, Capitanio JP, Cole SW. The neuroendocrinology of social isolation. Annu Rev Psychol. 2015;66:733–67.

19. O’Suilleabhain PS, Gallagher S, Stephane S, Zelley S, Theeke L. Sex differences in symptom severity and social isolation in the elderly during 19 years of follow-up. Psychosom Med. 2019;81(6):521–6.

20. Hawkley LC, Cacioppo JT. Loneliness. In: Sprecher S, editor. Encyclopedia of human relationships. Thousand Oaks: Sage; 2009. p. 985–90.

21. Hawkley LC, Cacioppo JT. Loneliness matters: a theoretical and empirical review of consequences and mechanisms. Ann Behav Med. 2010;40(2):218–27.

22. Peplau LA, Perlman D. Loneliness: a sourcebook of current theory, research and therapy. New York: Chichester: Wiley; 1982.

23. Leigh-Hunt N, Bagguley D, Bash K, et al. An overview of systematic reviews of loneliness and cardiovascular health. Antioxid Redox Signal. 2018;29(9):837–51.

24. Holt-Lunstad J, Smith TB, Layton JB. Social relationships and mortality risk: a meta-analytic review. PLoS Med. 2010;7(7):e1000316.

25. Wang J, Mann F, Lloyd-Evans B, Ma R, Johnson S. Associations between loneliness and perceived social support and outcomes of mental health problems: a systematic review. BMC Psychiatry. 2018;18(1):156.

26. Cohen-Mansfield J, Hazan H, Lerman Y, Shalom V. Correlates and predictors of loneliness in older adults: a review of quantitative results informed by qualitative insights. Int Psychogeriatr. 2016;28(4):557–76.

27. Dahlberg L, McKee KJ. Correlates of social and emotional loneliness in older people: evidence from an English community study. Aging Ment Health. 2014;18(4):504–14.

28. Bolton L. Exploring social isolation of leg ulcer patients. Wounds. 2017; 29(4):122–4.

29. Pyer LM, Yang L, da Rocha JM, Engeland CG. The effects of social isolation on wound healing mechanisms in female mice. Physiol Behav. 2014;127:64–70.

30. Pyer LM, Yang L, Mckenzie C, et al. Contrasting mechanisms by which social isolation and restraint impair healing in male mice. Stress. 2014;17(3):256–65.

31. Meyer T, Witz Rh. Mechanisms of mitochondrial redox signaling in psychosocial stress-responsive systems: new insights into an old story. Antioxid Redox Signal. 2018;28(9):760–72.

32. Smith KJ, Gavey S, Ne RJ, Kontai P, Victor C. The association between loneliness, social isolation and inflammation: a systematic review and meta-analysis. Neurosci Bull. 2020;36(2):1152-59.

33. Vingilis E, Hiyoshi A, Lintjes M, Fall K, Montgomery S. Longitudinal analysis of loneliness and inflammation at older ages. English longitudinal study of ageing. Psychoneuroendocrinology. 2019;110:104421.

34. Shivowitz-Etz S, Sarag O. Does loneliness ‘get under the skin’? Associations of loneliness with subsequent change in inflammatory and metabolic markers. Aging Ment Health. 2019;23(10):1588–66.

35. Cole SW. Social regulation of human gene expression: mechanisms and implications for public health. Am J Public Health. 2013;103(Suppl 1):S58–92.

36. Irwin MR, Cole SW. Reciprocal regulation of the neural and innate immune systems. Nat Rev Immunol. 2011;11(9):625–32.

37. Black DS, Irwin MR, Olmstead R, Ji E, Crabb Breen E, Motivala SJ. Tai chi meditation effects on nuclear factor-kappaB signaling in lonely older adults: a randomized controlled trial. Psychoso Psychosom. 2014;83(5):315–7.

38. Ganapathy N, Venkatasaran SS, Daniel R, Aravind RJ, Kumararukshavan VB. Molecular biology of wound healing. J Pharm Biomed Anal. 2012; 45(Suppl 2):S534–7.

39. Baud A, Mulligan MK, Casale FN, et al. Genetic variation in the social environment contributes to health and disease. PLoS Genet. 2017;13(1):e1006498.

40. Wiegand C, Schonfelder U, Abel M, Ruth P, Kaatz M, Hipler UC. Pro tease and pro-inflammatory cytokine concentrations are elevated in chronic compared to acute wounds and can be modulated by collagen type I in vitro. Arch Dermatol Res. 2010;302(6):419–28.

41. Powell ND, Sloan EK, Bailey MT, et al. Social stress up-regulates inflammatory gene expression in the leukocyte transcriptome via beta-adrenergic induction of myelopoesis. Proc Natl Acad Sci U S A. 2013;110(41):16574–9.

42. Oyola MG, Handa RJ. Hypothalamic-pituitary-adrenal and hypothalamic- pituitary-gonadal axes: sex differences in regulation of stress responsivity. Stress. 2017;20(5):476–94.

43. Theeke LA, Mallow J, Gianni C, Legg K, Glass C. The experience of older women living with loneliness and chronic conditions in Appalachia. Rural Ment Health. 2015;39(2):61–72.

44. Fredrickson BL, Gre wen KM, Aloe SB, et al. Psychological well-being and the human conserved transcriptional response to adversity. PLoS One. 2015; 10(3):e0121839.

45. Fredrickson BL, Gre wen KM, Coffey KA, et al. A functional genomic perspective on human well-being. Proc Natl Acad Sci U S A. 2013;110(33):13684–9.

46. Borton RM, Kenny DA. The moderator-mediator variable distinction in social psychological research: conceptual, strategic, and statistical considerations. J Pers Soc Psychol. 1986;51(6):1173–82.

47. Parker CN, Finlayson JK, Edwards HE. Ulcer area reduction at 2 weeks predicts failure to heal by 24 weeks in the venous leg ulcers of patients living alone. J Wound Care. 2016;25(11):626–34.

48. Theeke LA. Predictors of loneliness in U.S. adults over age sixty-five. Arch Psychiatr Nurs. 2007;21(3):387–96.

49. Theeke LA. Sociodemographic and health-related risks for loneliness and outcome differences by loneliness status in a sample of U.S. older adults. Res Gerontol Nurs. 2010;3(2):113–25.

50. Theeke LA, Goins RT, Moore J, Campbell H. Loneliness, depression, social support, and quality of life in older chronically ill Appalachians. Aust J Psychol. 2012;146(1):155–71.

51. Gershon RC, Lai JS, Bode R, et al. Neuro-QOL: quality of life items for adults with neurological disorders: item development and calibrations based upon clinical and general population testing. Qual Life Res. 2012; 21(3):475–86.

52. Theeke L, Horstman P, Mallow J, et al. Quality of life and loneliness in stroke survivors living in Appalachia. J Neurosci Nurs. 2014;46(6):E3–15.

53. Theeke L, Lucke-Wold AN, Mallow J, Horstman P. Life after stroke in Appalachia. Int J Nurs Sci. 2017;4(2):105–11.

54. Theeke LA, Mallow J, McBurney A, Rellick S, Van Gilder R. Effectiveness of LISTEN on loneliness, neuroimmunological stress response, psychosocial functioning, quality of life, and physical health measures of chronic illness. Int J Nurs Sci. 2016;3(3):242–51.

55. Theeke LA, Mallow J, Moore J, et al. Using gene expression analysis to examine changes in loneliness, depression and systemic inflammation in lonely chronically ill adults. Open J Nurs. 2016;6(8):620–31.

56. Cole SW, Levine ME, Arevalo JM, Ma J, Weir DR, Cirmmns EM. Loneliness, eudaimonia, and the human conserved transcriptional response to adversity. Psychoneuroendocrinology. 2015;21:1–7.

57. Cole SW, Hawkley LC, Arevalo JM, Sury C, Rose RM, Cacioppo JT. Social regulation of gene expression in human leukocytes. Genome Biol. 2007;8(9):R189.

58. Snodgrass JS, Dengah II, Huang LY, Mcg T, et al. Social genomics of healthy and disordered internet gaming. Am J Hum Biol. 2018;30(S5):e23146.

59. Cole SW, Capitanio JP, Chun K, Arevalo JM, Ma J, Cacioppo JT. Myeloid differentiation architecture of leukocyte transcriptome dynamics in perceived social isolation. Proc Natl Acad Sci U S A. 2015;112(49):15142–7.

60. Mocks M, Irwin MR, Jevtic I, et al. Trait sensitivity to social disconnection enhances pro-inflammatory responses to a randomized controlled trial of endoxin. Psychoneuroendocrinology. 2015;62:336–42.

61. Cacioppo JT, Cacioppo S, Cole SW, Capitanio JP, Goossens L, Boomser MA. Loneliness across phylogeny and a call for comparative studies and animal models. Perspect Psychol Sci. 2015;10(2):202–12.
66. Capitano JP, Hawkley LC, Cole SW, Cacioppo JT. A behavioral taxonomy of loneliness in humans and rhesus monkeys (Macaca mulatta). PLoS One. 2014;9(10):e110307.

67. Cole SW. Human social genomics. PLoS Genet. 2014;10(8):e1004601.

68. Hopman WM, Buchanan M, VanDenKerkhof EG, Harrison MB. Pain and health-related quality of life in people with chronic leg ulcers. Chronic Dis Inj Can. 2013;3(3):167–74.

69. McRobert J. Smoking and its effects on the healing process of chronic wounds. Br J Community Nurs. 2013;18(Sup3):S18–23.

70. Gouin JP, Kiecolt-Glaser JK. The impact of psychological stress on wound healing: methods and mechanisms. Crit Care Nurs Clin North Am. 2012;24(2):201–13.

71. Becket S, Pietsch AM, Kuper M, et al. M.a.I.D.: a prognostic score estimating probability of healing in chronic lower extremity wounds. Ann Surg. 2009;249(4):677–81.

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