Factors influencing psychological, social and health outcomes after major burn injuries in adults: cohort study protocol

Martha Druery, Peter A Newcombe, Cate M Cameron, Jeffrey Lipman

ABSTRACT

Introduction The goal of burn care is that ‘the quality of the outcome must be worth the pain of survival’. More research is needed to understand how best to deliver care for patients with burns to achieve this aim. Loss of independence, function as well as loss of income for patients with burns and carers cause a significant burden at both individual and societal levels. Much is being done to advance knowledge in the clinical care field; however, there has been a paucity of research exploring psychosocial outcomes. This paper describes the study background and methods, as implemented in an Australian cohort study of psychosocial outcomes after major burn injuries.

Methods and analysis In this inception cohort study, a target sample of 230 participants, aged 18 years or over, admitted to a single statewide burns centre with a burn injury are identified by hospital staff for inclusion. Baseline survey data are collected either in person or by telephone within 28 days of the injury and participants then followed up with telephone interviews at 3, 6 and 12 months postburn. Injury and burns treatment information is collected from medical records. Social support is measured as a predictor variable using the Multidimensional Scale of Perceived Social Support. Outcome data are collected via standardised measures in the domains of Quality of Life (SF-12, EQ-5D, BSHS-B), depression (PHQ-9), post-traumatic stress disorder (PCL-C, PAS), community integration (CIQ-R) and Quality-Adjusted Life Years (EQ-5D). Additional survey questions measure life satisfaction, return to work and public services utilisation at 12 months postinjury. Data analysis methods will include analysis of variance, Pearson correlation and hierarchical multiple regression analyses.

Ethics and dissemination Hospital-based and University of Queensland Human Research Ethics Committees have approved the protocol. Results from the study will be disseminated at national and international conferences, in peer-reviewed journals and in a doctoral thesis.

Trial registration number Australia New Zealand Clinical Trials Registry (ACTRN12616000828426). Retrospectively registered on 23 June 2016; pre-results.

INTRODUCTION

Identifying determinants and barriers to acceptable and optimal burns outcomes is central to developing targeted interventions, programs and models of care. Cost-effectiveness of clinical treatment is also an important consideration due to the significant health expenditure on resource intensive burn care. It is widely agreed in the burns literature that outcome is multidimensional for burns survivors given the spectrum of impacts this type of injury has on people, physically, emotionally, psychologically and socially.

While a number of studies consistently report that a large proportion of burns survivors make a satisfactory functional recovery, there remain difficulties with other aspects of injury adjustment and a percentage of patients demonstrating poorer outcomes. These findings led to concentrated investigations into more global outcomes research.

Broader outcomes studies have investigated Quality of Life (QoL). Health-Related QoL

Strengths and limitations of this study

- This is a longitudinal, prospective study that incorporates injury, treatment, personal and social factors to determine predictors of long-term psychological, social and health outcomes of hospitalised adults postburn in an Australian context. As such, it will address a number of the methodological limitations that have constrained previous burns outcomes research.
- The methodology of researcher-led follow-up is intended to limit attrition, resulting in high continuing participation rates.
- The target of a larger sample size than previously reported in similar studies, in combination with multiple validated measures, will enhance the rigour of this research.
- Participants of the proposed study are patients of one Australian burns unit and therefore the results may not be generalisable to other Australian burns survivors or those from other countries.
- A further limitation is the observational nature of the study. However, observational data are sufficient to address the key research objective of reporting the psychological, social and health outcomes of the study group at 12 months postburn.

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There exists a broad spectrum of factors which have been investigated in terms of their predictive or causal relationship with burns outcomes such as QoL and HRQoL. The overall consensus is that while injury severity is strongly predictive of poorer outcome in terms of functional independence and QoL, the role of psychological and environmental factors has been empirically acknowledged. The few studies that have applied a multifactorial theoretical model to investigate the hierarchical association and direction of causality between these separate factor groupings have been significantly constrained by methodological limitations such as retrospective, cross-sectional designs, small sample size and one time point of data collection nearly 10 years postinjury.

The present research takes a multidimensional view. It investigates a potential range of factors, hypothesised to predict physical, psychological and health outcomes in functional, mental health, social and life satisfaction domains. The broader societal impacts of return to work, public services utilisation and ‘Quality Adjusted Life Years’ (QALYs) health utility weights will also be investigated.

The aim of this study is to identify factors that are predictive of psychosocial outcomes 12 months after a burn injury in order to determine which hospitalised adults are likely to experience poor long-term outcomes postburn.

METHODS AND ANALYSIS

Study design

This study uses a single-centre prospective longitudinal cohort design to capture the reported dynamic nature of burn recovery over time. Baseline data are collected as soon as feasible after the burn injury to gather preinjury information. Intermediate data collection occurs at 3 months postinjury (when the majority of patients will have experienced the exposure and impacts of returning to the community) and 6 months postinjury (when the acute period of care is usually completed and a rehabilitation routine has been established). Outcome data are collected at 12 months postburn.

Study setting

Participants are recruited from the Professor Stuart Pegg Adult Burn Centre at the Royal Brisbane and Women’s Hospital in Australia, the sole statewide specialist adult burns referral centre for Queensland, northern New South Wales, the Northern Territory and the Pacific Islands. An average of 390 patients with a new burn injury are admitted to this burns unit annually.

Study population

Inclusion criteria

Eligible participants are those who have been admitted for at least 24 hours as an initial hospitalisation to the Professor Stuart Pegg Adult Burns Unit at the Royal Brisbane and Women’s Hospital. They have either sustained a major burn injury defined as affecting ≥10% of the total body surface area (unless affecting the hands/forearms, feet/lower legs, head/face or perineum), the presence of airway burn or an inhalation injury, a length of stay >7 days or any burn injury requiring specialist multidisciplinary care in a burn centre, such as an operative procedure. To be eligible, participants are aged 18 years or over in order to legally provide consent for their own participation. Participants who are non-English speaking are also invited to participate, as this potentially marginalised group has previously been excluded from burns outcomes research in Australia.

Exclusion criteria

The exclusion criteria includes patients who lack the requisite medicolegal capacity to provide consent to participate, those who are admitted for subsequent treatment of a previously sustained injury, not domiciled in Australia, not expected to remain so for 12 months postinjury or those who are currently incarcerated in a correctional facility. Patients with burns who are identified by the burns treating team to be experiencing significant physical or psychological symptoms are excluded as are patients who are identified to be exhibiting aggressive or violent behaviour. Patients with burns who were experiencing significant functional impairment prior to their burn injury, resulting in a high level of dependence on residential care services due to advanced age or disability are also ineligible.

Sampling, recruitment and consent

Eligible hospitalised participants in an adult burns unit are identified by nursing staff who introduce the study, advise patients of their eligibility, provide a Participant Information and Consent Form and seek agreement for a researcher to attend the unit for further discussion. Recruitment commenced in October 2016 and is ongoing. Eligibility of agreeing participants is confirmed by a member of the clinical team in relation to cognitive and psychological capacity and the researcher attends the burns unit at a clinically appropriate time. The researcher then verbally explains the study, discusses the Participant Information and Consent Form, answers questions and seeks consent. Patients who are physically able to sign the form, do so with a witness present to also sign.

Data collection

Researchers conduct an 8-minute baseline interview with consenting participants and access injury data from medical records and the hospital Burns Database. Researchers then conduct follow-up telephone interviews at 3, 6 and 12 months postdate of injury. The duration of interviews are approximately 20 minutes for time points 2 and 3 and 30 minutes for time point 4.

Data regarding injury, treatment, personal and social factors are recorded at baseline and outcomes at the follow-up time points as reported in table 1.

Table 1  Data collected at baseline and follow-up

<table>
<thead>
<tr>
<th>Injury data</th>
<th>% total body surface area burned, % full thickness burn, circumstances, mechanism, bodily location, respiratory injury, multitrauma</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Baseline</strong></td>
<td></td>
</tr>
<tr>
<td>Preburn personal and social factors</td>
<td>Age, gender, relationship status, employment status, income, education, remoteness of residence, housing, history of trauma, prior substance use, material resources, physical and psychological comorbidities, ethnicity, primary language</td>
</tr>
<tr>
<td>Previous health status—EQ-5D⁵⁵</td>
<td>Mobility, self-care, usual activities, pain/discomfort, anxiety/depression</td>
</tr>
<tr>
<td>Social support—MSPSS¹⁶</td>
<td>Family, friends, significant other, global support</td>
</tr>
<tr>
<td><strong>3, 6 and 12 months postdate of injury</strong></td>
<td></td>
</tr>
<tr>
<td>Psychological well-being</td>
<td>Depression risk: PHQ-9²⁴ PTSD risk: PCL-C²⁶ PTSD and depression risk postinjury: PAS³⁶</td>
</tr>
<tr>
<td>Composite score</td>
<td>Total symptom severity score PAS_P (PTSD) PAS_D (depression)</td>
</tr>
<tr>
<td>QoL outcomes</td>
<td>Burns specific outcomes: BSHS-B²⁹ Generic outcomes: SF-12²⁴</td>
</tr>
<tr>
<td>Affect and relations Function Skin involvement Mental Component Summary Physical Component Summary</td>
<td></td>
</tr>
<tr>
<td>Community integration</td>
<td>CIQ-R⁴³ Home Integration Subscale Social Integration Subscale Productivity Subscale Electronic Social Networking Subscale</td>
</tr>
<tr>
<td>12 months postdate of injury</td>
<td></td>
</tr>
<tr>
<td>Life satisfaction</td>
<td>Single-item Likert scale—question developed for the purposes of this study</td>
</tr>
<tr>
<td>Societal outcomes</td>
<td>QALY/Health utility: EQ-5D²⁵ Return to work and self-reported public service utilisation questions developed for the purposes of this study</td>
</tr>
<tr>
<td>Treatment factors</td>
<td>Total length of hospital stay, number of operative procedures, intubation, ICU length of stay</td>
</tr>
<tr>
<td>Complications</td>
<td>Heterotopic ossification, deceased</td>
</tr>
</tbody>
</table>

BSHS-B, Burn Specific Health Scale—Brief version; CIQ-R, revised Community Integration Questionnaire—Revised; EQ-5D, EuroQol Group Standardised Measure of Health Status; ICU, intensive care unit; MSPSS, Multidimensional Scale of Perceived Social Support; PAS, Post-traumatic Adjustment Scale; PCL-C, PTSD Symptom Checklist—Civilian version; PHQ-9, Patient Health Questionnaire; PTSD, post-traumatic stress disorder; QALY, Quality Adjusted Life Years; QoL, Quality of Life; SF-12, 12-Item Short Form Health Survey.

Research measures

Independent predictor variables

Predisposing risk factors

Premorbid individual factor data including age, gender and ethnicity are sourced from the Hospital Based Clinical Information System (HBCIS). Questions relating to premorbid use of tobacco, alcohol and recreational drugs and comorbidities have been replicated from New Zealand injury-outcome research, based on census questions.¹⁵ Premorbid social factor data such as marital status and place of residence are also sourced from HBCIS. Remoteness of residence is determined from the Australian Bureau of Statistics postcode classifications.

Questions relating to income, employment, education, housing tenure, material resources, trauma and violence history are derived from previous research.¹⁵

Social support

Due to the significance of social support as a predictor of postburn functional outcomes,⁶ it is important to investigate in relation to psychosocial outcomes. The Multidimensional Scale of Perceived Social Support (MSPSS) is a 12-item self-report instrument which measures interpersonal functioning and social support from family, friends and significant other on a 7-point Likert scale from ‘Very strongly agree’ to ‘Very strongly
disagree. A global support score can be derived from all three subscales. Good internal reliability and validity have been reported and although not validated for a burns population, it has been used extensively among mental health, injury and general populations.

**Burns factors**

Injury factors including percentage of total body surface area burned, percentage of full thickness burn, circumstances of injury, mechanism of injury, visible location of injury, presence of inhalation injury and of heterotopic ossification are sourced from the Royal Brisbane and Women’s Hospital burns unit database with assistance from the database manager. Treatment factors including length of stay, number of operative procedures, ventilation and intensive care unit length of stay are also sourced from the hospital burns unit database as this information is routinely collected for all admitted patients with burns.

**Outcome variables**

**Quality of Life and life satisfaction**

There is no consensus in the literature in relation to a definition or consistent measurement tool for QoL. There is also confusion whether the many component variables of QoL should be considered determinants or indicators. QoL has been defined by the WHO as ‘a person’s perception of his/her position in life within the context of the culture and value systems in which he/she lives and in relation to his/her goals, expectations, standards, and concerns’ (WHOQOL Group, p28). This term has been adopted in injury outcomes literature to conceptualise the multidimensional nature of recovery across physical, functional, psychological and social domains from the patient’s perspective.

There is consensus in the general injury literature to include both generic QoL and condition-specific outcome measures in study designs. The benefit of generic outcome measurement is that it allows comparisons with normative, other injury and disease groups. Outcome measurement in the present study was informed by the most frequently used generic QoL tools in burns research; the 36-Item Short Form Health Survey by the most commonly used generic QoL tools in burns research; the 36-Item Short Form Health Survey—Brief version (SF-36) and EuroQol Group Standardised Measure of Health Status (EQ-5D).

While the SF-36 has been validated for use in a Australian burns population, the SF-12 was preferred for use in the current study due to its comparable psychometric properties combined with its brevity, thereby reducing participant burden for this repeat measures design. This instrument includes 12 items from the original SF-36, scored to provide two summary scores: a Mental Component Summary and a Physical Component Summary.

The EQ-5D was developed by the EuroQol Group as a generic self-report measure of health status and for economic appraisal. It comprises two components. The first is a descriptive system which includes responses of ‘No problems’, ‘Some problems’ or ‘Extreme problems’ to the five dimensions of mobility, self-care, usual activities, pain/discomfort and anxiety/depression. The second component is a Visual Analogue Scale where the participants rate their health on a given day on a vertical scale ranging from 0=‘Best imaginable health state’ to 100=‘Worst imaginable health state’. The EQ-5D is recommended for use by the Cologne Group in relation to QoL outcome measurement for traumatic injuries and has been chosen for its sound psychometric properties and extensive use throughout the world, including burns outcomes research.

Condition-specific QoL measurement is particularly informative due to the individual consequences of burn injury that would not otherwise be captured by generic tools in isolation. The 40-item Burn Specific Health Scale—Brief version (BSHS-B) has been the most frequently used measure of burns outcome in recent decades. The BSHS-B has been validated to measure burns outcomes in the domains of simple abilities, hand function, heat sensitivity, treatment regimens, body image, affect, interpersonal relationships, sexuality and work with excellent internal consistency reported as ranging between 0.75 and 0.93. Willebrand and Kildal validated a simplification of these nine domains into three clusters: affect and relations (affect, interpersonal relationships and sexuality), function (simple abilities, hand function) and skin involvement (heat sensitivity, treatment regimens and body image). These authors treated the work subscale as a separate domain due to an identified issue of double loadings.

In addition to these, concept analyses have found that subjective satisfaction with life is also of merit. Therefore, a single question referring to overall life satisfaction has been included with a 5-point Likert scale response, ranging from 1 (Extremely dissatisfied with my life in general at the moment) to 5 (Extremely satisfied with my life in general at the moment).

**Psychological well-being**

The domains of mental health that are recommended for routine outcome measurement by the American Burn Association (ABA) are depression and post-traumatic stress disorder. They recommend the Patient Health Questionnaire (PHQ-9) for screening of depression in the inpatient or outpatient adult burns population. The PHQ-9 is a 9-item self-report measure which has been adapted for assessing depression severity from the Primary Care Evaluation of Mental Disorders instrument (PRIME-MD). The items are scored from 0 (Not at all) to 3 (Nearly every day), providing a composite score from 0 to 27 and lower-bound cut-off points for depression severity levels of none (0), mild (5), moderate (10), moderately severe (15) and severe (20). The internal reliability of the PHQ-9 has been reported to be excellent (α=0.89) with similarly sound sensitivity, specificity and construct validity.

The ABA recommends the Post-Traumatic Stress Disorder (PTSD) Symptom Checklist—Civilian version (PCL-C) for assessing PTSD if assessment by clinical interview is not feasible. These authors report that the
Networking' (ESN) subscale, scoring improvements and revised in 2014 (CIQ-R) to include an 'evaluating social with burns.' It has not been validated for use with the burns Centrelink income support and corrective services.

Public service utilisation
Questions with binary 'Yes/No' answers will be included in the survey booklet at 12 months postburn in relation to usage of non-burns related health services, mental health services, alcohol and drug services, public rental housing, Centrelink income support and corrective services.

Free-text fields will be used to collect comments in relation to the nature of these services.

Health utility
The EQ-5D, which is used as a generic QoL outcome measure also provides a health utility outcome/QALY measure. Each participant’s health state is derived from scoring the five descriptive system domains into a 5-digit code. This code can then be converted into a single summary index with a formula that results in a QALY for each participant.

Sample size
Online calculation using Epi Info online software (https://www.cdc.gov/epiinfo/7/index.htm) yielded a target sample size of 138 completions at 12 months postburn. Input values for the sample size calculation were based on a of 5%, a standard power of 90%, with a large effect size and incidence or exposure estimates as follows. The expected incidence of poor psychosocial outcomes in the burns population with good social resources is 20% and for those with poor social resources is 50%, based on reported figures in the OECD Better Life Index (https://www.oecdbetterlifeindex.org) and conventional calculations from injury and disease epidemiology literature, with a ratio of exposed to unexposed participants estimated at 2.5. Allowing for 40% attrition, which approximates that reported in the burns literature, a final sample size of 230 is required for statistical significance.

Data analysis
Summary statistics will be used to describe the study cohort. Categorical data will be expressed as frequencies and percentages. A missing values analysis will be conducted to determine the randomness of the missing data. The outcome of these analyses will guide the appropriate missing value replacement method (eg, multiple imputation, linear interpolation) as required to ensure statistical power is maintained at 90%. The statistical methods will include analysis of variance to explore changes across time points and Pearson correlation to explore relationships between risk factors, burns factors and social support with psychological, social and health outcomes. Hierarchical multiple regression analyses will be used to test the predictability of injury, treatment, social and environmental factors on burns outcomes, adjusted for potentially confounding covariates (eg, premorbid individual factors). All statistical analyses will be two-tailed and considered significant at p<0.05. Free-text data collected in relation to public service utilisation will be subject to thematic analysis for descriptive reporting of response categories.

DISCUSSION
The primary focus of this study is to examine associations and predictors for long-term psychological, social and health outcomes for hospitalised patients with burns encompassing life satisfaction, psychological well-being,
functional outcomes, health status, return to work and community integration. It is hypothesised that preburn social factors such as low income, unemployment, remoteness of residence, poor housing tenure, limited material resources and restricted social support will be significant predictors of poorer psychosocial outcomes in adult survivors with burns. It is also hypothesised that limited preburn social resources will be significantly associated with poorer health utility weights and greater public service utilisation at 12 months postinjury.

Burns studies have used various time points for prospective longitudinal data collection but few have provided a rationale for the choice of schedule. The US National Institute on Disability and Rehabilitation Research Burn Model System Database is a large repository of burn injury incidence, treatment and outcome data which has designated follow-up outcome data to be collected at the time of discharge then at 6, 12 and 24 months after injury. These time points are similar to those propounded by the Trauma Registry of the American College of Surgeons at time of discharge, 1 month, 6 months and 12 months postburn. The Cologne Group, a multinational consensus forum regarding QoL outcome measurement of trauma survivors, also recommends data collection at T0 (preinjury), T1 (acute phase—3 months postinjury), T2 (rehab phase—6 months) and T3 (long-term outcome phase—12 months). These time points represent phases of recovery calculated from date of injury as opposed to date of discharge, which was abandoned by the European Consumer Safety Association working group on injury-related outcomes measurement guidelines due to variability. It has been shown that there is little change after 12 months postburn, a finding echoed in the general injury literature, so this has been chosen as an end-point for outcome data collection in the present study. Although the practice of retrospectively collecting patient-reported preinjury data is prone to recall bias, it is important to determine the preburn situation and circumstances of participants. Data are not collected at the time of discharge as this is a variable time point dependent on injury severity, complications and other factors. The schedule will remain static for all participants for consistency of analysis.

An economic analysis of costs associated with injury also supports this methodology given that the major costs associated with long-term, static sequelae eclipse the high cost of medical and rehabilitative treatment in the acute and initial improvement phases. There is a risk that survivors with burns who consent to participate in the study will differ from those who decline, which will introduce bias. Comparison of participants and non-participants will be possible in relation to routinely collected hospital data for all admitted patients with burns. Attrition is an added risk for any longitudinal study design, introducing potential constraints in generalisability of findings from the study cohort to the burns population. However, it will be possible to assess any differences in those who become lost to follow-up over the course of the project.

Recall bias is limited due to the prospective design of the proposed study; however, this will be a potential source of error in relation to baseline measurement of premorbid individual and social factors as well as health status. An attempt to minimise this is made by collecting baseline data as soon as possible after admission to hospital or after resumption of cognitive capacity. The measures invite participants to recall varying periods of time from ‘at the moment’ to ‘in the last month’. While this is also a potential source of bias, it has been reported not to be a significant difficulty for respondents.

Participants of the proposed study are patients of one Australian burns unit and therefore the results may not be generalisable to other Australian burns survivors or those from other countries. However, the Queensland population is largely representative of the nation, so the relationships between factors and outcomes as well as direction of effects are likely to be similar in other regions. As cognitively impaired burns survivors are excluded due to the limitations inherent in proxy reporting, results will not be generalisable to those with cognitive deficits. The study will also only recruit survivors with burns who were admitted for their burn injury. Those who did not require treatment in the statewide burns centre or hospitalisation may also be significantly affected by their burn injury but not be represented by this sampling method.

An added limitation is due to the decision not to include a measure of cognitive functioning. Although other researchers have extended the EQ-5D to measure this, the amendment arguably undermines the tool’s psychometric properties. Measurement of cognitive functioning is beyond the scope of the present study which is focused on the potential relationship between preinjury social factors and psychosocial outcomes.

A further limitation is the risk of response bias or Hawthorne Effect due to the interview modality of administering the surveys, which is inherent in self-report measures. It would be optimal to integrate self-report responses with clinician assessments or observations; however, feasibility of the study precluded this option.

This study addresses a number of the methodological limitations that has constrained previous research as it is a longitudinal, prospective study that incorporates injury, treatment, personal and social factors to determine predictors of long-term psychological, social and health outcomes of hospitalised adults postburn in an Australian context. The methodology of researcher-led follow-up is intended to limit attrition and the anticipated high participation rate will optimise recruitment of a representative sample for generalisable findings. The target of a larger sample size than previously reported in similar studies, in combination with multiple validated measures, contribute to the rigour of this research.

The findings will provide detailed information about the multiple impacts experienced over time by survivors of major burn injuries in the study population. It will reveal factors that are associated with the highest risk of adversity and the greatest societal costs. The results are...
anticipated to assist in identifying those patients with burns who are more likely to experience poorer psychosocial outcomes, in order that evidence-based intervention, services and programs can be delivered to target these more vulnerable people. Identifying factors influencing injury outcomes is intended to lead to future research on modifiable factors for better outcomes in the population after serious burn-injury and cost-effective improvements.

**Ethics and dissemination**

The study protocol has been approved by the Royal Brisbane and Women’s Hospital Human Research Ethics Committee (HREC/15/QRBW/145) and the Behavioural Social Sciences Ethics Review Committee, University of Queensland (2015001089). The appropriate permissions have been granted for access and usage of all specified databases, clinical information systems, patient records and standardised measures. Results from this study will be disseminated at national and international conferences, in peer-reviewed journals and in a doctoral thesis.

**Correction notice**

This paper has been amended since it was published Online First. Owing to a scripting error, some of the publisher names in the references were replaced with ‘BMJ Publishing Group’. This only affected the full text version, not the PDF. We have since corrected these errors and the correct publishers have been inserted into the references.

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**Contributors**

MD conceived of the study and its design, drafted the grant proposal, manages its coordination as the principal investigator of the study and drafted the manuscript. PAN and CMC participated in the design of the study and all authors, including JL, critically revised the manuscript and approved the final version of the manuscript.

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**Competing interests**

None declared.

**Ethics approval**

The study protocol has been approved by the Royal Brisbane and Women’s Hospital Human Research Ethics Committee (HREC/15/QRBW/145) and the Behavioural Social Sciences Ethics Review Committee, University of Queensland (2015001089).

**Provenance and peer review**

Not commissioned; peer reviewed.

**Data sharing statement**

This article does not contain any data. No datasets have been analysed during the current phase of the study; therefore, data sharing is not applicable. Study materials and future datasets may be available from the corresponding author on reasonable request.

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