Healing the Past by Nurturing the Future—co-designing perinatal strategies for Aboriginal and Torres Strait Islander parents experiencing complex trauma: framework and protocol for a community-based participatory action research study

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ABSTRACT

Introduction Child maltreatment and other traumatic events can have serious long-term physical, social and emotional effects, including a cluster of distress symptoms recognised as ‘complex trauma’. Aboriginal and Torres Strait Islander (Aboriginal) people are also affected by legacies of historical trauma and loss. Trauma responses may be triggered during the transition to parenting in the perinatal period. Conversely, becoming a parent offers a unique life-course opportunity for healing and prevention of intergenerational transmission of trauma. This paper outlines a conceptual framework and protocol for an Aboriginal-led, community-based participatory action research (action research) project which aims to co-design safe, acceptable and feasible perinatal awareness, recognition and assessment strategies for Aboriginal parents experiencing complex trauma.

Methods and analysis This formative research project is being conducted in three Australian jurisdictions (Northern Territory, South Australia and Victoria) with key stakeholders from all national jurisdictions. Four action research cycles incorporate mixed methods research activities including evidence reviews, parent and service provider discussion groups, development and psychometric evaluation of a recognition and assessment process and drafting proposals for pilot, implementation and evaluation. Reflection and planning stages of four action research cycles will be undertaken in four key stakeholder workshops aligned with the first four Intervention Mapping steps to prepare programme plans. Ethnics and dissemination protocols are consistent with the National Health and Medical Research Council Indigenous Research Excellence criteria of engagement, benefit, transferability and capacity-building. A conceptual framework has been developed to promote the application of core values of safety, trustworthiness, empowerment, collaboration, culture, holism, compassion and reciprocity. These include related principles and accompanying reflective questions to guide research decisions.

INTRODUCTION

Child maltreatment and other adverse childhood experiences (ACEs) are an international health priority,1 contributing to a wide range of long-lasting physical, social and emotional...
Health issues. There is growing international consensus to recognise a cluster of distress symptoms people may experience following childhood exposure to severe threats, called complex post-traumatic stress disorder (PTSD; complex trauma). This classification describes a symptom profile that typically follows traumatic experiences of a prolonged nature or repeated adverse events from which separation is not possible. These symptoms include ‘affect/emotional dysregulation’, ‘negative self-concept’ and ‘relational disturbances’, in addition to previously recognised PTSD symptoms of ‘re-experiencing the events (triggers), avoidance and a sense of threat’. These traumatic experiences often involve interpersonal violation and occur within childhood family or institutional care giving systems (eg. childhood abuse, severe domestic violence, torture or slavery). Broader societal factors can amplify or counteract the impact of potentially traumatic experiences. Aboriginal and Torres Strait Islander (Aboriginal) people in Australia are particularly affected by complex trauma, following a legacy of historical trauma, which includes state-sanctioned systematic removal of Aboriginal children from their families and ongoing discrimination. While community cohesion, access to services and cultural continuity have been shown to have a protective effect for some trauma-related outcomes among Aboriginal people, within the context of colonisation sociocultural risk factors experienced by many Aboriginal communities are likely to amplify rather than counteract the effects of complex trauma originating from childhood experiences.

There are strong associations between child maltreatment and a wide range of physical and psychological morbidity and risk factors, including smoking, eating disorders, unplanned pregnancies and adverse birth outcomes. Critically, these long-lasting relational effects can impede the capacity to nurture and care for children, leading to ‘intergenerational cycles’ of trauma. Experiences of child maltreatment are not equally distributed across general populations and WHO uses a sociocultural framework to highlight the links between higher levels of social adversity and increased rates of child maltreatment experienced in some communities worldwide. These factors also interact and create a ‘compounding intergenerational effect’ on health inequities. As such, this is a crucial issue for improving health equity worldwide. ‘Life course approaches’ are central to understanding complex intergenerational causal pathways and also for identifying critical ‘intervention points’ for prevention and support to improve health equity.

The transition to parenting during the perinatal period (pregnancy to 2 years after birth) is a critical ‘life course’ transition for parents who have experienced complex trauma. Trauma responses may be triggered by the intimate nature of experiences associated with pregnancy, birth and breast feeding, and the attachment needs of the infant. The long-lasting relational effects can impede the capacity of parents to nurture and care for their children, and may contribute to ‘intergenerational cycles’ of trauma.

Conversely, the transition to parenthood offers a unique life-course opportunity for emotional healing and development. A positive strengths-based focus during this often-optimistic period has the potential to transform the ‘vicious cycle’ of intergenerational trauma into a ‘virtuous cycle’ that contains positively reinforcing elements. When parents can manage trauma responses and provide love and nurturing care, this love is returned by children, and trauma responses can be realigned, promoting healing in the parent and optimal development for the infant.

Frequent scheduled contacts with perinatal care providers before and after childbirth and across the first 2 years offer an opportunity for providing comprehensive system-based supports for people experiencing complex trauma during this period. This is particularly important because it may be the first time many of this predominantly young childbearing population have had contact with universal health services since childhood. Despite these clear risks and opportunities, few interventions are available for parents with specific histories of maltreatment, and there are no systematic, culturally informed processes or evidence of effective strategies to identify and support Aboriginal parents experiencing complex trauma.

The benefits of involving communities in co-designing healthcare strategies are increasingly recognised. This is critical in the perinatal period for Aboriginal families experiencing complex trauma for several reasons. First, there is very limited evidence of effective interventions internationally. Australian guidelines for the treatment of complex trauma and trauma-informed care emphasise the need for complex trauma to be understood within relational networks and social environments if it is to be adequately addressed. Aboriginal Australians, despite suffering great disadvantage and adversity, demonstrate strong resistance to those actions that are foreign to Aboriginal culture, including separation from families, discrimination and removal from country. Thus, we will engage in respectful collaborative research with and alongside Aboriginal people and keep Aboriginal people’s strengths and protective factors to the fore. These strengths include rich cultural relationships and kinship networks that foster relatedness and connectedness for children. Collaboration with local Aboriginal leaders and Aboriginal organisations has been shown to be critical in adapting child trauma therapies among other Indigenous communities.

Second, Aboriginal conceptualisations of social and emotional well-being are holistic and incorporate connection to land, culture, spirituality, family and community; all of which are impacted by complex trauma, which is sometimes referred to as ‘relational trauma’. The rich relational understandings of well-being may offer important insights for other Indigenous and non-Indigenous communities.

Third, there are risks associated with identifying parents with complex trauma. Labelling individuals as ‘at risk’ has the potential to undermine parents’ existing resilience and coping skills, and trigger inappropriate notifications.
to a potentially punitive child protection system. These concerns are particularly salient for Aboriginal communities, with the history of colonisation and forced child removals from families, and ongoing high rates of infants being removed from Aboriginal families, which have had devastating ongoing intergenerational impacts. Finally, despite a history of childhood adversity, most parents are able to nurture and care for their children. Evidence suggests that examining these ‘cycles of discontinuity’ are an important place to start to illuminate innovative strategies for support.

Aims and objectives

*Healing the Past by Nurturing the Future* is a formative Aboriginal-led, community-based participatory action research (action research) project, which aims to co-design perinatal strategies to support Aboriginal parents experiencing complex trauma. There is currently insufficient evidence to identify potentially acceptable, feasible and effective strategies to support Aboriginal parents experiencing complex trauma, hence the focus of this project is formative research.

The expected outcomes of the project are to identify strategies that are considered acceptable to Aboriginal parents and feasible for service providers. Piloting, implementation and evaluation of the effectiveness of these perinatal strategies will be the subject of a sequential project following this formative design stage.

The co-design strategies aim to improve four key domains of perinatal care:

- **Awareness** of the impact of trauma on parents or ‘trauma-informed’ perinatal care to minimise the risks of triggering and compounding trauma responses.
- **Safe recognition** of parents who may benefit from assessment and support, with processes to reduce risk of harm.
- **Assessment** of complex trauma symptoms to accurately identify parents experiencing distress.
- **Support** strategies for parents to heal, including psychological/emotional, social, cultural and physical strategies.

The purpose of this protocol paper is to illustrate the processes, frameworks and methods used by an Aboriginal-led research team to generate rigorous context-relevant strategies, while also fostering cultural and emotional safety for participants, partners, research staff and the broader Aboriginal community. This paper includes an outline of the following elements:

- Community involvement in the project.
- Conceptual framework for developing safe research processes.
- Research activities within the four action research cycles and Intervention Mapping (IM) steps.
- Ethical considerations and research dissemination plans.

Due to the evolving nature of action research and co-design research, submissions for Human Resarch Ethics Committee (HREC) approval are planned in three distinct ‘ethics phases’, following key stakeholder co-design workshops one, two and three. At the time of submitting this protocol, phase I and II HREC approval had been granted, and HREC submission is planned for phase III in late 2019. Therefore, this protocol includes a detailed description of ‘phase I and II’ activities, with a brief outline only of anticipated phase III activities (highlighted in text).

**METHODS AND ANALYSIS**

Patient and public (community) involvement

This project involves Aboriginal people at every level, and a detailed description is outlined in the National Health and Medical Research Council (NHMRC) Indigenous Research Excellence Criteria (see online supplementary file 1). In summary, the majority of the investigator team are Aboriginal with extensive expertise in this area. The need for this research has been identified in national Aboriginal conferences and formally supported by three Aboriginal community controlled ‘peak bodies’, who play a leading role in Aboriginal health initiatives, the Aboriginal Medical Services Alliance of Northern Territory; the Aboriginal Health Council of South Australia and the Victorian Aboriginal Community Controlled Health Organisation.

We are using an *action research model* to ensure ongoing community involvement is built into the research plan, including refinement of the research questions. Action research draws on phenomenology and critical theory to generate constructivist grounded theory using mixed methods. It involves a practical community-based focus and collaboration for action. The focus of the first year has been meaningful *community engagement* to enable action research. We have established formal partnerships and recognise the leadership of five partner service organisations with this project, including: Central Australian Aboriginal Congress (Northern Territory); Nunkuwarrin Yunti of South Australia and Women’s and Children’s Health Network (South Australia); the Royal Women’s Hospital (Victoria) and the Bouverie Family Healing Centre (Victoria).

Participants in this study include Aboriginal parents, perinatal service providers, Aboriginal elders and key stakeholders (service providers, researchers, policy-makers and community leaders working to address complex trauma). Participants are required to provide informed consent prior to participating in study activities, and draft findings of each activity are provided to participants for feedback, prior to broader community dissemination. We invite key stakeholders from all Australian jurisdictions to participate in the four co-design workshops to enable broader national collaboration in planning for subsequent programme pilot, implementation and evaluation.

**Conceptual framework: developing safe research processes**

To articulate the values for the project and address risks and contextual complexities, we have developed a conceptual framework (figure 1) drawing on holistic Aboriginal
constructs of social and emotional well-being. Protocols that have been critical for informing this conceptual framework include:

- **Power Threat Meaning Framework (PTMF)**, an overarching structure for identifying patterns in emotional distress, unusual experiences and troubling behaviour, as an alternative to psychiatric diagnosis and classification (p. 5). We will incorporate the PTMF by reframing behaviours related to complex trauma as normal self-protective responses to threatening situations rather than pathological deficits.

- **Principles for population-based screening** to assess the benefits, risks, costs, acceptability, accuracy and potential risk of harms resulting from recognising and assessing parents experiencing complex trauma.

- **Indigenous research methodologies** that involve privileging Aboriginal worldviews, self-determination and Aboriginal community control.

The conceptual framework incorporates two elements:

- Four main domains of awareness, recognition, assessment and support.
- Eight core values with related principles and questions.

**Four main domains of recognition, assessment, awareness and support**

The four main domains were developed during the early community engagement stages of the project which revealed concerns about the use of language such as ‘screening’ and ‘intervention’, which implies ‘something is wrong’ with a person, and is not consistent with PTMF framing of trauma to ask ‘what has happened to you’. There are also sensitivities in the context of Aboriginal communities in Australia, with controversial Government ‘interventions’ imposed on Aboriginal communities. The domains of ‘recognition’ and ‘assessment’ broadly align with ‘screening’ strategies that incorporate a safe and feasible two-tiered process for care providers to recognise parents who may require more in-depth assessment for complex trauma; and ‘intervention’ approaches to improve trauma-informed perinatal care and minimise the risks of re-traumatising parents (awareness), and provide trauma-specific support.

**Eight core values with related principles and questions**

Using online searches and team members’ clinical knowledge, we identified seven frameworks that included trauma-informed values and principles. These values and principles were mapped and consensus was reached by the project team for eight core values: safety, trustworthiness, empowerment, collaboration, culture, holism, compassion and reciprocity. Each contains action-oriented principles that enable the core values to be realised, and are accompanied by questions developed to aid reflection on whether the activity under consideration is consistent with the core value (see online supplementary file 2).

**Setting**

Research activities will be conducted in three of seven Australian jurisdictions selected on the basis of existing research relationships and expressed interest by key stakeholders: Northern Territory, South Australia and Victoria. Approximately 23% of Australian Aboriginal people live in these three jurisdictions across mixed urban, rural and remote demographic contexts.

**Data storage and triangulation**

All data will be securely stored using REDCap software and accessible only to members of the project team. Wherever possible, data will be stored in de-identified form. However, where concerns exist about the health of a participant, the safety plans and responses relating to that participant will be stored to enable appropriate follow-up by healthcare professionals.

Multiple data sources will be triangulated within this project (as described below), which will increase confidence in the findings through the confirmation of proposed ideas from two or more independent sources. Data collection tools are designed to progressively inform the co-design of safe, acceptable and feasible perinatal awareness, recognition, assessment and support strategies.

**Research approaches**

An Intervention Mapping (IM) approach is used in this project to frame the co-design process. IM uses ‘theory and evidence as foundations for taking an ecological approach to assessing and intervening in health problems and engendering community participation’ (p. 7). This formative research project addresses IM steps one to four, which are aligned with four key stakeholder workshops (figure 2). IM steps five and six (implementation and evaluation) will form the basis of a subsequent project.

**Action research processes** will be used to foster an iterative co-design process comprising four ‘plan-act-observe-reflect’ cycles. The ‘reflect’ and ‘plan’ action research stages
will be conducted in four key stakeholder workshops, which align with the first four steps of IM. The ‘act’ and ‘observe’ stages of the action research cycles involve a series of mixed method ‘research activities’ that will be refined in each ‘reflect’ and ‘plan’ stage within the workshops. We outline research activities within each of the IM steps and action research cycles below. We note that HREC approval has been received for ‘phase I and II’, but that activities planned for a ‘phase III’ HREC submission have not been approved and are subject to review (thus briefly outlined here).

1. Action research cycle and IM step 1: developing relationships and understanding the problem

This first action research cycle includes: (1a) evidence reviews, (1b) the first key stakeholder workshop, aligned with IM step 1, (1c) mapping domains included within existing assessment tools and (d) a pilot discussion group with senior Aboriginal women. Each of these activities is described further below:

1a: Evidence reviews: scoping review and evidence map of studies involving parents in the perinatal period with a history of childhood maltreatment; and comprehensive systematic reviews

The purpose of the scoping review and evidence map was to identify preliminary evidence, and enable development of protocols for a series of comprehensive systematic reviews (see online supplementary file 3). The scoping review findings have been incorporated into subsequent research activities, including: presentation at workshop 1; generating ‘cards’ of key issues described by parents elsewhere in discussion groups with senior Aboriginal women and parents and scoping ‘strengths’ to be included in an assessment tool. The scoping review has also been critical to refine the search strategy for a series of comprehensive reviews.

1b: Key stakeholder workshop 1

The purpose of workshop 1, aligned with IM step 1 (understanding the problem and developing a logic model), was to provide a forum for preliminary engagement with key stakeholders to:

► Introduce the rationale for the project and share preliminary evidence from the scoping review to enable informed discussion and clarification of goals (logic model).
► Establish safety protocols for working with parents, service providers, key stakeholders, team members and the wider Aboriginal community.
► Understand the context and issues for key stakeholders regarding identifying and supporting Aboriginal parents experiencing complex trauma.

Recruitment and sample: key stakeholders were identified through consultation and using a snowballing recruitment process of advertising about the project through Aboriginal and academic health networks, professional meetings and conferences. People expressing interest in the project were included in a key stakeholder email list, and received updates about the project and invitations to the workshops which were cost-free to enable attendance. Approximately 40 people participated in workshop 1.
Data collection and analysis: a facilitation guide was developed to address the aims of the workshop (see online supplementary file 4) and promote a culturally and emotionally safe environment. Strategies to support any participants who may experience ‘triggers’ themselves (ie, trauma responses) during the workshop and psychological support were provided.

Data were collected in the form of workshop materials developed by participants (butchers paper notes) and observer notetakers. Data were collated into themes and circulated to workshop participants to check the accuracy of the interpretations. A summary of the workshop is available on the project website. In keeping with the action research process, findings were reflected on and used for planning workshop 2 (2a) and developing the conceptual framework and a detailed safety protocol.

1c: Scoping assessment tools
The purpose of scoping existing assessment tools for complex trauma and/or a parental history of child maltreatment, and for assessing resilience and strengths was to:
► Map the range of areas of distress included within existing assessment tools.
► Enable informed consultation with key stakeholders about each of the main areas of distress and if all important areas were considered.
► Map domains of resilience and strengths.

Data collection and analysis: distress assessment tools were identified through the scoping review and consultation. For each tool, data were extracted on: description of the tool; key references; validation information; symptoms of distress and/or trauma exposures measured. Data were synthesised into summary ‘areas of distress’ (see online supplementary file 5), and further refined by the research team for presentation to key stakeholders at workshop 2.

Strengths domains were mapped from existing resilience tools, mediating/moderating factors and ‘strategies parents use’ in the scoping review, and data generated from a discussion group with senior Aboriginal women and in key stakeholder workshop 2.

1d: Pilot discussion group with senior Aboriginal women
The purpose of this discussion group was to:
► Consult with community leaders about the effects of complex trauma during the perinatal period for Aboriginal parents, and what might help or hinder the parenting transition.
► Pilot qualitative methods proposed for use with parents, and gather feedback on the safety and appropriateness of these approaches and tools.

Recruitment and sample: a convenience sample of six to eight senior Aboriginal members of a community group that had expressed interest in the project.

Data collection and analysis: a facilitation plan was developed that included use of: visual tools and natural materials to facilitate discussions; cards illustrating the main themes from the scoping review to build on existing research; third person scenarios to increase safety and minimise the ‘directness’ of sensitive discussions so they were not intrusive; use of metaphors and symbolism; and a ‘strengths-based’ focus on ‘healing’ rather than ‘trauma’. The discussion group was facilitated by an Aboriginal psychologist (YC) and Aboriginal midwife (CC) with expertise in conducting discussion groups with Aboriginal people. Additional psychological support was available in line with the detailed safety plan.

A detailed discussion group protocol was developed (available on request). Data were collected in the form of visual notes and images provided by group participants, observer notes and a recording of the discussion which was transcribed verbatim. Two Aboriginal researchers (YC, CC) independently coded data into themes (thematic analysis) and these were discussed with participants to check the interpretation of the data accurately reflected both what was said as well as the intent. Themes were shared with key stakeholders at workshop 2 for reflection and planning of subsequent parent discussion groups.

2. Action research cycle and IM step 2: scoping assessment domains with a focus on research evidence and community knowledge, and developing objectives
The second action research cycle includes: (2a) a second key stakeholder workshop, aligned with IM step 2, (2b) refining the assessment tool domains and preliminary questions for parents, (2c) identifying ‘gold standard’ assessment for comparison in psychometric testing, training and cultural adaptation (if required) and (2d) first round of discussion groups with parents who have experienced complex childhood trauma.

2a: Key stakeholder workshop 2
The purpose of workshop 2 was to reflect on the activities from action research cycle 1 and plan for ethics phase II. This is aligned with IM step 2, which involves refining the project objectives and consulting with key stakeholders regarding:
► The areas of distress to be included in an assessment tool.
► Reflection on pilot discussions with senior Aboriginal women regarding areas of strengths and pretesting the proposed approach for working with parents.

Recruitment and sample: key stakeholders were identified as described in 1b, with approximately 60 participants attending.

Data collection and analysis: a facilitation guide was developed to address the aims of the workshop (see online supplementary file 6) and promote a culturally and emotionally safe environment. A traditional healer (Ngangkerre) worked alongside the registered psychologist to cater for different support needs and recognise the equal value of respective expertise.

Data regarding the 12 summary areas of distress were gathered using a modified Delphi approach. Each area of distress was allocated to a table and facilitator.
Participants gathered in groups of six to eight at one table and were given individual forms (non-identified) with a description of the area of distress, with additional information provided by the facilitator. They were asked to indicate the degree of ‘importance’ (1–5) of the area of distress, and discuss and/or document any comments about why, who, where and how questions regarding this area of distress should be asked. These discussions will be central for informing co-design of safe ‘recognition’ strategies in workshop 3. Participants rotated around all 12 tables. Data were transcribed and imported into NVivo for thematic analysis and future triangulation with data to be collected at workshops 3 and 4.

Reflections regarding the discussion group with senior Aboriginal women and pretesting the discussion group ‘tree of life’ approach for use with parents were recorded by participants pictorially using sticky notes on butcher paper. The ‘tree of life’ approach was used as it provides a hopeful and inspiring approach to talking about challenging issues and generates visual images to promote shared understanding, and had been used by effectively by an Investigator in other settings (JA). This positive ‘tree of life’ tool aligned with the ‘strengths-based’ focus on parents hopes and dreams and the support parents need moving forward, rather than dwelling on past experiences. These images were photographed, data were coded into themes and imported into NVivo for thematic analysis and future triangulation with other data sources to inform co-design of awareness and support strategies.

2b: Developing assessment tool areas of distress and strength questions for parents
The purpose of refining the areas of distress and strength questions that may be included in an assessment tool is to enable initial evaluation of ‘face validity’ of the questions with parents and identify any important issues requiring direct discussion with parents.

Data collection and analysis: data collected in key stakeholder workshop 1 (1b), scoping assessment tools (1c) and workshop 2 (2a) will be collated in NVivo for thematic analysis. These themes and issues will be refined in consultation with the research team to propose questions related to ‘areas of distress’ to be included in an assessment tool. Questions for assessing each of these areas of distress will be drafted, based on questions validated in existing tools (International Trauma Questionnaire and a version of the Harvard Trauma Questionnaire adapted for Aboriginal people and cultural resources regarding mental health literacy). The purpose of refining the areas of distress and strength questions will be developed by the research team, with the research team, and ‘pretested’ in a convenience sample of Aboriginal colleagues. The proposed questions will be incorporated into the first round of discussion groups with parents to evaluate preliminary ‘face validity’ of the proposed questions.

2c: Identifying ‘gold standard’ assessment for comparison in psychometric testing, training and cultural adaptation (if required)
The purpose of this activity is to identify the best possible ‘gold standard’ for comparison with our proposed assessment tool.

Data collection and analysis: a preliminary list of suitable tools for use as a ‘gold standard’ was generated by consensus within the research team following a systematic and transparent process of consideration. From this, the trauma section of the WHO World Mental health Composite International Diagnostic Interview has been proposed. Consultation about the proposed ‘gold standard’ will also be conducted with three or four additional key external psychiatric and psychological experts.

Up to six Aboriginal psychologists will train together in the use of the ‘gold standard’ structured clinical interview to enable them to reflect and use their cultural and clinical expertise. They will advise whether any aspects need adaptation for use with Aboriginal parents.

2d: First round of discussion groups with Aboriginal parents
The purpose of the first round of discussion groups with Aboriginal parents is to:

► Understand key perinatal experiences affecting Aboriginal parents and what kinds of awareness (trauma-informed care) and support strategies might help or hinder the transition to parenting for parents experiencing complex trauma.
► Evaluate the ‘face validity’ of draft questions in a preliminary assessment tool.

Recruitment and sample: approximately 24 Aboriginal parents will be invited to participate in discussion groups, one to three groups per participating jurisdiction with up to eight parents in each. The size of the group will be determined by the study coordinator in consultation with service provider staff regarding the most appropriate mix of gender, the level of comfort of participants in group discussion and language. We estimate that this will be sufficient to produce theoretical saturation of thematic categories, particularly when triangulated with data from the pilot discussion group and key stakeholder workshops. However, if saturation of themes is not reached we will consider further discussion groups as needed.

Individual parents will be recruited through the services they attend for perinatal care using direct and indirect methods. Service providers will be given written and verbal information about the study by the research team. Service providers will then ask potentially eligible parents if they give consent to be contacted by the research team to discuss the study in more detail and consider if they would like to consent to participate.
in the discussion group. Parents may be asked if they would like to be contacted by the research team in a private area while waiting to attend for services, after a consultation, or during other community activities. Additionally, flyers will be displayed describing the purpose of the study and providing contact details for parents to contact the research team directly.

Inclusion criteria: participants will be eligible to participate if they identify as Aboriginal and/or Torres Strait Islander, are aged 16 years or older and they or their partner are currently pregnant or have a child <2 years of age.

Exclusion criteria: parents experiencing current serious mental illness (e.g., acute psychoses or other mental health difficulties that may affect their capacity to provide informed consent and/or pose a risk to the safety of the parent and other participants in the discussion group). This will be assessed by service staff prior to asking for consent to be contacted, and by the research team prior to asking for consent to participate in the discussion group.

Data collection and analysis: a facilitation plan has been refined based on feedback from the pilot discussion group (1d) and workshop 2 (2a). The discussion group will be facilitated by an Aboriginal researcher with expertise in conducting discussion groups with Aboriginal people. Psychological support will be provided. The facilitation plan (see supplementary file 7) includes use of: visual tools and natural materials to facilitate discussions; cards illustrating the main themes from the scoping review to build on existing research; third person scenarios to increase safety and minimise the ‘directness’ of sensitive discussions so they are not intrusive; use of metaphors and symbolism to explain complex phenomena and a ‘strengths-based’ focus. Data will be collected using visual notes prepared by participants in a ‘tree of life’ activity to frame discussions about the needs for Aboriginal parents experiencing complex trauma, and transcribed audio recordings of the discussions.

Two researchers will independently conduct thematic analysis and discuss draft themes with participants to check the interpretation of the data. The themes from this discussion group will be triangulated with data from previous project activities and shared with key stakeholders participating in workshop 3 to inform co-design of a preliminary awareness and support strategies.

Additional face-to-face interviews will be conducted with up to nine parents to assess the ‘face validity’ of a preliminary list of distress and strengths questions. These will be further refined in workshop 3.

3. Action research cycle and IM step 3: developing acceptable and feasible perinatal awareness, recognition, assessment and support strategies

The third action research cycle includes: (3a) key stakeholder co-design workshop 3, aligned with IM step 3, (3b) psychometric evaluation of assessment tool, (3c) a second round of discussion groups with parents and (3d) discussion groups with service providers.

3a: Key stakeholder co-design workshop 3

The purpose of workshop 3, aligned with IM step 3, is to co-design the preliminary recognition and assessment process and possible awareness and support strategies.

Recruitment and sample: key stakeholders will be identified as previously described, with up to 60 participants anticipated.

Data collection and analysis: a facilitation guide will be developed to address the aims of the workshop and promote a culturally and emotionally safe environment as per previous workshop. The workshop will incorporate triangulated data from previous action research cycles to foster informed co-design of for preliminary:

- Awareness and support strategies, informed by scoping review, qualitative systematic review of parents views, intervention review and relevant data from discussion groups and key stakeholder workshops. The purpose is to generate an overinclusive range of options, for further refinement in parent discussion groups to rank and assess acceptability, and service provider discussion groups to assess feasibility.
- Recognition and assessment strategies, informed by data from the scoping review, scoping of assessment tools, key stakeholder workshop 2 exercise and the face validity assessments in parent discussion groups. The purpose is to develop processes to foster safe recognition of parents who may benefit from further assessment, to be further refined following parent and service provider discussion groups, and an overinclusive list of assessment tool items for psychometric evaluation and refinement.

Summary of proposed activities to be submitted for ‘phase III’ HREC approval

The detailed methods for the following activities will be refined based on feedback from ‘reflection’ and ‘planning’ from activities described in ‘ethics phase I and II’ in consultation with partner organisation staff, and submitted for ethical approval. A brief outline of main activities, aims and sample size estimates are included below.

3b: Psychometric evaluation of assessment tool

The psychometric evaluation aims to develop a valid assessment tool that enables perinatal care providers to accurately identify strengths, as well as complex trauma symptoms (measurement sensitivity) while minimising the erroneous identification of parents who are not experiencing complex trauma symptoms (measurement specificity).

The sensitivity of a complex trauma assessment will need to be high for the inventory to be effective and appropriate for use in practice, where our priority would be that all parents who could benefit from further assessment and support are recognised. Based on previous estimates of PTSD and complex trauma, we conservatively estimate that 20% of Aboriginal parents will meet...
subthreshold criteria of at least two symptoms. Identifying parents meeting subthreshold criteria will maximise the sensitivity of the instrument to identify PTSD and complex trauma and we estimate that a sensitivity of 90% would be achieved. Thus, a sample size of 173 participants will be required to yield an estimate of the instrument sensitivity with a two-sided 95% CI with a width of 10% of the estimate. This sample size will also enable estimation of the specificity of the instrument to correctly identify participants who had not experienced complex trauma.

3c: Second round of discussion groups with parents, which aims to assess the acceptability of the proposed recognition and assessment process; and awareness and support strategies

Approximately 24 Aboriginal parents will be recruited to participate in discussion groups, one to three groups per participating jurisdiction with up to eight parents in each.

3d: Discussion groups with service providers, which aim to assess the feasibility of the proposed recognition and assessment process; and awareness and support strategies

Approximately 24 service providers will be recruited to participate in discussion groups, one to two groups per participating jurisdiction with up to eight service providers in each.

4. Action research cycle and IM step 4: planning for pilot, implementation and evaluation

The fourth and final action research cycle includes a fourth key stakeholder workshop and drafting plans with perinatal service providers to pilot, implement and evaluate safe acceptable and feasible perinatal awareness, recognition, assessment and support strategies for Aboriginal parents experiencing complex trauma.

4a: Key stakeholder co-design workshop 4

The purpose of workshop 4, aligned with IM step 4 to ‘refine strategies and prepare to pretest’, aims to reflect on the research findings with service providers and develop plans for seeking funding to pilot, implement (IM step 5) and evaluate (IM step 6) perinatal awareness, recognition, assessment and support strategies.

ETHICS AND DISSEMINATION

Ethics

Action research poses unique challenges for seeking HREC approval. While there is an overarching structure and an outline of main activities, the detail required for ethical approval evolves during the action research process. In this project, submissions for HREC approval are being submitted to relevant jurisdictional authorities in three phases, with HREC approval for phases I and II granted at the time of submission. This is particularly important in a project involving sensitive content such as complex trauma, where the HREC need to examine draft tools and resources to consider risks for triggering distress symptoms against potential benefits.

This staged approach also enables piloting and reflection on the ‘safety’ of the research activities and flexibility to refine research processes. For example, in this project, discussions were first held with a predominantly professional group of ‘key stakeholders’ in workshop one, then with a group of senior Aboriginal women in a ‘pilot’ discussion, and then a proposed approach was ‘pretested’ in a second ‘key stakeholder’ workshop, prior to submitting the final plans for discussion groups directly with Aboriginal parents. The intent is to ensure our approach and processes maximise safety and minimise the risk of distress for parents, while also gathering the data needed to inform iterative development of awareness, recognition, assessment and support strategies. At the time of submitting this protocol, HREC approval had been granted for phase I and II (figure 2).

The funding proposal for this project was assessed by an Indigenous research panel using the NHMRC Indigenous Research Excellence criteria (see online supplementary file 1) developed to promote ethical and culturally appropriate research with Aboriginal communities. In addition, we have developed a conceptual framework (figure 1), which outlines the ethical and cultural values for this project. A specific safety framework describes how the primary value of safety will be fostered for parents, service providers, key stakeholders and team members, and the broader Aboriginal community.

Dissemination

We have developed a research dissemination plan (available on request), in line with the NHMRC Indigenous Research Excellence criteria (see online supplementary file 1) and the value of reciprocity.

The research dissemination plan includes:

► Offering two-way information exchange for all community meetings (ie, prior to the meeting asking if there are any presentations about topics people would like us to offer to their staff and community members about complex trauma and parenting).

► Publication of articles in open access journals with links to relevant Aboriginal health websites.

► Face-to-face presentations in national and international conferences.

► Translating all findings into plain language summaries.

► Incorporating art, presentations and other mediums to present information.

► Preparing a video/short YouTube clip with essential information for community members and making this freely available on the project website and sharing at community meetings.

► Ensuring all relevant information is presented on the research website, which is regularly monitored for accuracy, optimised for search engine performance and follows accessibility guidelines.
Contributors CC is the study Principal Investigator and drafted this protocol based on the project proposal and other relevant project planning documents involving many people as outlined in acknowledgements and author contribution statements. GG, SJB, JA, DG, HH, KG, YC, SC, FKM, CA, SEB, HM, TH and JN are study investigators who contributed to development of the project proposal, project planning and drafting the manuscript. FKM conducted sample size estimate calculations for psychometric evaluation of the assessment tool. DD assisted with development of the conceptual framework, study planning and drafting the manuscript. NR, SH and YC are employed on the project and have contributed to development of planning documents, conceptual framework, ethics submissions which involved many considerations outlined in this protocol, and drafting the manuscript. All authors read and approved the final manuscript.

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REFERENCES


52. CATSINAM, ACM. *Birthning on country position statement*. Canberra: College of Aboriginal and Torres Strait Islander Nurses and midwives, Australian College of Midwives, CRANA Plus, 2016.


